# Application of mechanistic models to separate the effects of mutation, selection, and drift on protein sequence evolution

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To my mother

# Acknowledgments

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Nothing in Biology Makes Sense Except in the Light of Evolution.

-Theodosius Dobzhansky

Nothing in evolutionary biology makes sense except in the light of population genetics  $\hbox{-Michael Lynch}$ 

## Abstract

Mathematical and statistical models are useful for describing and understanding observations in genetics and genomics. These models have to constantly be updated to reflect current biological understanding. As opposed to descriptive and phenomenological models, mechanistic models allow for the extraction of more biologically relevant information based on underlying principles. Mutation, selection, and genetic drift are the three forces guiding evolution. Mechanistic models rooted in population genetics principles allow us to determine how these forces shape observed data. I demonstrate the usage of mechanistic models to relate protein coding sequences to their fitness landscapes and the evolutionary forces shaping them. Using the yeast *L. kluyveri*, I show the increased cost of protein synthesis due to a large scale introgression with mismatched codon usage. Furthermore, I analyze site-specific selection on amino acids in the beta-lactamase protein TEM, which confers antibiotic resistance in *E. coli* and related species.

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# Chapter 1

## Introduction

Protein synthesis is the most costly metabolic process a cell performs (REEDS et al., 1985; WATERLOW and MILLWARD, 1989; BUTTGEREIT and BRAND, 1995; WARNER, 1999; AKASHI and GOJOBORI, 2002; LINDQVIST et al., 2018) causing selection to maximize the benefit of protein synthesis and performing it as efficiently as possible. Studying the ratio of cost to benefit of protein synthesis is, therefore, important to understand the evolution of protein coding sequences (GILCHRIST et al., 2009; SHAH and GILCHRIST, 2011a; GILCHRIST et al., 2015; BEAULIEU et al., in review). However, the strength of selection varies greatly between genes, from low expression genes with codon usage dominated by mutation bias between nucleotides over highly expressed genes reflecting the dominance of selection for efficient translation of the mRNA, to selection on the amino acid composition required for the function of the protein.

We can formalize the cost and benefit of a protein coding sequence and formulate mathematical models. Mathematical and statistical models have long been used to describe or summarize observations in genetics and genomics. Often without addressing the underlying biological mechanisms - mutation, selection, and genetic drift - shaping DNA sequences, but as phenomelogical descriptions. As researchers learn more about the underlying processes and more genetic and genomic data is available, the mathematical models that allow for the extraction of information from this data have to keep up. For example, after the unraveling of the degenerate genetic code by MATTHAEI and NIERENBERG

(1964); NIERENBERG and MATTHAEI (1961); MAXWELL (1962); LEDER and NIERENBERG (1964), and many others, researchers noticed that synonymous codons are not found in uniform proportions (FITCH, 1976; GRANTHAM et al., 1980; IKEMURA, 1981; GRANTHAM et al., 1981; SHARP et al., 1988). Models of codon usage, however, were long purely descriptive and heuristic (IKEMURA, 1981; BENNETZEN and HALL, 1982; SHARP, 1987; WRIGHT, 1990). Similarly, phylogenetic models have long been phenomelogical (JUKES and CANTOR, 1969; DAYHOFF et al., 1978; KIMURA, 1980; FELSENSTEIN, 1981; ALTSCHUL, 1991), describing the rate of change between states without regards for the forces guiding evolution, mutation, selection, and genetic drift. Zuckerkandl and Pauling (1962) proposed that the evolution of proteins is constant over time and between lineages before the genetic code was fully deciphered and at a time were protein synthesis was barely understood based on their observation that similarity on hemoglobin is correlated with divergence time. This work is therefore focused on the application of mechanistic models rooted in first principles and their application to protein coding sequences

Mechanistic models are used throughout biology (GOLDMAN and YANG, 1994; LAUREAU, 1998; DAVIS and PELSOR, 2001; DORON-FAIGENBOIM and PUPKO, 2007; McGill et al., 2007). By modeling the process underlying the observed data mechanistic models provide insights into the processes and estimates of parameters shaping the data (LIBERLES et al., 2013). A wide variety of information is stored in protein and protein coding sequences, e.g. structure (Anfinsen, 1973), mutation bias (Shah and Gilchrist, 2011a; Gilchrist et al., 2015), protein synthesis rate (Gilchrist, 2007; Gilchrist et al., 2015). Mechanistic models can be used to extract these informations and to study the relative strength of mutation, selection, and genetic drift leading to the observed sequences.

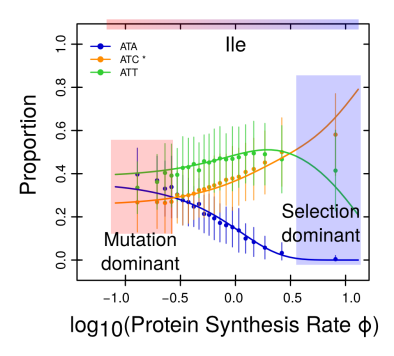


Figure 1.1: ROC SEMPPR model behavior for Isoleucine. The proportion of each codon observed changes with protein synthesis rate. Mutation is dominant when protein synthesis rate is low, mutationally favored codons are observed with the highest frequency. With the increase of protein synthesis rate, the influence of selection increases until the system is dominated by selection. The selectively favored codon is observed with the highest frequency.

# 1.1 Cost: Decomposing Codon Usage

Mutation bias on codon usage is a reflection of the cellular environment while selection on codon usage allows us to make inferences about the cellular and external environment a genome has evolved in. The relative strength of mutation and selection on individual genes varies, allowing us to separate mutation bias and selection, specifically selection against translation overhead cost (GILCHRIST, 2007; SHAH and GILCHRIST, 2011a; GILCHRIST et al., 2015). Genes with low protein synthesis rates are thought to be under weak selection for codon usage and their codon usage is therefore dominated by mutation bias. In contrast, genes with high protein synthesis rates are thought to be under strong selection and their codon usage is therefore dominated by selection. However, mutation bias and selection can differ within the genome.

For example, strand specific mutation bias (LAFAY et al., 1999; ROMERO et al., 2000), differences in the tRNA pool throughout life stages (SAGI et al., 2016), or introgressions and horizontal gene transfer (MDIGUE et al., 1991; LAWRENCE and OCHMAN, 1997) can produce multiple genomic environments. Chapter 2 extends the mechanistic model ROC SEMPPR GILCHRIST et al. (2015) to allow for a mixture distribution of mutation and selection parameters LANDERER et al. (2018) and provides researchers with a software tool to address intra genomic variation in codon usage. However, there is a significant difference to classical mixture approaches. In addition to gene population specific parameters, ROC SEMPPR also estimates a gene specific parameter (protein synthesis rate). Therefore, the protein synthesis rate for each gene has to be estimated assuming that the a gene is in each gene population. This can provide additional insight into the adaptiveness of a gene to alternative genomic environments. Figure 1.1 illustrates how the proportions of synonymous codons change with increasing protein synthesis rate. When the protein synthesis rate is low, mutation bias between codons dominates the proportions of synonymous codons while increasing protein synthesis increases the strength of selection (see GILCHRIST et al. (2015) for details).

In chapter 3, I apply AnaCoDa to analyze the synonymous codon usage of the yeast  $L.\ kluyveri$  which experienced a large scale introgression replacing the whole left arm of chromosome C (FRIEDRICH et al., 2015). I studied the differences in the parameters describing codon usage between the endogenous  $L.\ kluyveri$  genes and the introgressed exogenous genes. Recognizing the differences in codon usage between the endogenous and exogenous genes allowed me to improve prediction of protein synthesis rate, and separate the effects of mutation bias and selection in the endogenous  $L.\ kluyveri$  genes and the introgressed exogenous genes. This information was used to determine a potential donor lineage in  $E.\ gossypii$ , estimate the time since introgression, and estimate the genetic load of the introgression.

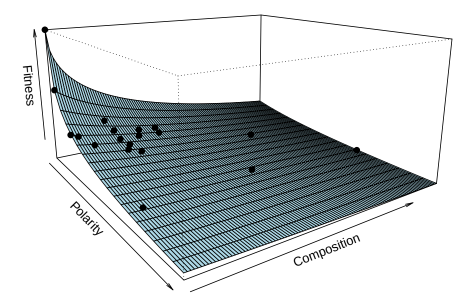


Figure 1.2: Decline in fitness with distance in physicochemical space from the optimal amino acid. Fitness decline of amino acids (black dots) relative to optimal amino acid (Alanine). Weighting of physicochemical properties according to Grantham (1974). The full fitness surface can be described but only 20 discrete amino acid states are available for selection to act on.

# 1.2 Benefit: Selection on Amino acids

Genes are evolving with natural selection favoring proteins that encode their function optimally, with mutations and genetic drift reducing functionality. Amino acid preference and the relative strength of mutation, selection, and genetic drift usually varies between sites along the protein sequence. The number of parameters required to describe protein fitness increases exponentially with the length of the protein if interactions between sites are accounted for. Attempts to incorporate selection into phylogenetic approaches are, therefore, limited to site specific selection. The goal of chapter 4 is to estimate the strength of site specific selection on amino acids from protein coding sequences in a phylogenetic framework.

Ignoring interactions between sites allows to describe the site specific fitness landscape of a protein. Some approaches rely on the description of the full fitness landscape and therefore require  $19 \times L$ , where L is the length of the peptide in amino acids, parameters (Lartillot and Philippe, 2004; Le et al., 2008; Wang et al., 2008; Holder et al.,

2008; Wu et al., 2013; Tamuri et al., 2014). As this is still a large number of parameters the incorporation of experimentally determined site specific selection on amino acids is an attractive alternative (Bloom, 2014; Thyagarajan and Bloom, 2014; Bloom, 2017). Alternatively, assumptions about the nature of selection can reduce the number of parameters required. For example, negative frequency dependent selection (Goldman and Yang, 1994; Muse and Gaut, 1994; Thorne et al., 1996) or stabilizing selection (Beaulieu et al., in review) allow for a reduction in fitness of amino acids with distance in physicochemical space.

SelAC (BEAULIEU et al., in review), a model of stabilizing selection, assesses the fitness of each amino acid relative to the fitness peak (Figure 1.2). Fitness is assumed to decline exponentially with distance in physicochemical space to the optimal amino acid. In chapter 4 I apply SelAC to the  $\beta$ -lactamase TEM and estimate site specific selection on amino acids and compare the inferred fitness landscape to empirical estimates from deep mutation scanning experiments (STIFFLER et al., 2016). I find that experimentally informed amino acid preferences improve model fit but do not accurately reflect the evolution of TEM. Furthermore, I show that the information on site specific selection on amino acids can be extracted from protein coding sequences by models rooted in first principles like SelAC.

# Chapter 2

AnaCoDa: Analyzing Codon Data with Bayesian mixture models

This chapter is a lightly revised version of a paper by the same name published in Bioinformatics and co-authored with Alexander Cope, Russell Zaretzki, and Michael A. Gilchrist.

C. Landerer, A. Cope, R. Zaretzki, M.A. Gilchrist, AnaCoDa: analyzing codon data with Bayesian mixture models, Bioinformatics, 34, 2018, 2496-2498

# 2.1 Abstract

AnaCoDa is an R package for estimating biologically relevant parameters of mixture models, such as selection against translation inefficiency, nonsense error rate, and ribosome pausing time, from genomic and high throughput datasets. AnaCoDa provides an adaptive Bayesian MCMC algorithm, fully implemented in C++ for high performance with an ergonomic R interface to improve usability. AnaCoDa employs a generic object-oriented design to allow users to extend the framework and implement their own models. Current models implemented in AnaCoDa can accurately estimate biologically relevant parameters given either protein coding sequences or ribosome foot-printing data. Optionally, AnaCoDa can utilize additional data sources, such as gene expression measurements, to aid model fitting and parameter estimation. By utilizing a hierarchical object structure, some parameters can vary between sets of genes while others can be shared. Genes may be assigned to clusters or membership may be estimated by AnaCoDa. This flexibility allows users to estimate the same model parameter under different biological conditions and categorize genes into different sets based on shared model properties embedded within the data. AnaCoDa also allows users to generate simulated data which can be used to aid model development and model analysis as well as evaluate model adequacy. Finally, AnaCoDa contains a set of visualization routines and the ability to revisit or re-initiate previous model fitting, providing researchers with a well rounded easy to use framework to analyze genome scale data.

# Availability:

AnaCoDa is freely available under the Mozilla Public License 2.0 on CRAN (https://cran.r-project.org/web/packages/AnaCoDa/).

# 2.2 Introduction

AnaCoDa is an open-source software implemented in R (R CORE TEAM, 2015) that allows researchers to analyze genome-scale data like coding sequences and ribosome footprinting data using evolutionary or analytical models in a Bayesian framework. AnaCoDa was developed to analyze selection on synonymous codon usage in the form of ribosome overhead cost (GILCHRIST et al., 2015; WALLACE et al., 2013; SHAH and GILCHRIST, 2011b). However, other codon metrics like the codon adaptation index (SHARP, 1987) or the effective number of codons (WRIGHT, 1990) are also provided as reference. In addition, three currently unpublished models to analyze coding sequences for evidence of selection against nonsense errors and estimate ribosome pausing times from ribosome footprinting data are included. AnaCoDa implements an adaptive Gibbs sampler within a Metropolis-Hastings Monte Carlo Markov Chain (MCMC). This allows for the incorporation of prior knowledge such as observed gene expression levels and easy sampling from the posterior distribution to estimate parameter values and quantify degree of uncertainty. AnaCoDa provides a mixture distribution option to all implemented models, combining genes into sets by estimating the posterior probabilities of set membership based on gene-set specific parameters shared by all genes assigned to a given set. AnaCoDa provides a generic, mixture distribution option to all implemented models, allowing for the estimation of condition specific parameters or the automatic categorization of data into different sets based on differences in their posterior probabilities of set membership. In addition to the four models provided, AnaCoDa provides a modular infrastructure such that additional genome scale or even phylogenetic models can be integrated.

The **AnaCoDa** framework works with **AnaCoDa** requires gene specific data such as codon frequencies obtained from coding sequences or position specific footprint counts. Conceptually, **AnaCoDa** allows for three different types of parameters. The first type are gene specific parameters such as protein synthesis rate or relative functionality. The second type are gene-set specific parameters, such as mutation bias terms or translation error rates.

These parameters are shared across genes within a set and can be exclusive to a single set or shared with other sets. While the number of gene sets must be pre-defined by the user, set assignment of genes can be pre-defined or estimated as part of the model fitting. Estimation of the set assignment provides the probability of a gene being assigned to a set allowing the user to asses the uncertainty in each assignment. The third type are hyperparameters allowing for the construction and analysis of hierarchical model. Hyperparameters control the prior distribution for gene and gene-set specific parameters such as mutation bias or protein synthesis rate.

## 2.3 Features

**AnaCoDa** provides an interface written in R, a freely available programming language noted for its ease of use for even inexperienced programmers. As a result, **AnaCoDa** is accessible to researchers with minimal computational experience.

The interface of **AnaCoDa** is designed for quick and efficient data analysis. Generally, the only input needed for fitting a model to the data are protein-coding codon sequences in the form of a FASTA file or a flat-file containing codon counts obtained from ribosome foot-printing experiments. **AnaCoDa** also provides visualization functionality, including plotting functions to compare parameter estimates for different mixture distributions and display codon usage patterns. In addition, diagnostic functions such as those for calculating and visualizing the degree of autocorrelation in the parameter traces are provided.

#### Robust and efficient model fitting

AnaCoDa has built-in features designed to improve the robustness and performance of the implemented MCMC approach. For example, the implemented MCMC automatically adapts the proposal width for sampled parameters such that an user defined acceptance range is met, improving sampling efficiency of the MCMC and computational performance. Even though

**AnaCoDa** is written in C++, analysis of large datasets and/or complex models can be very computationally intensive. To protect users from computer failures or aid in the collection of additional MCMC samples, **AnaCoDa** can periodically produce output checkpoint files, which can be used to restart an MCMC chain from a previous time point. In addition, **AnaCoDa** automatically thins all parameter traces - meaning only every  $k^{th}$  sample is kept - increasing the effective number of samples and reducing its memory footprint.

Although AnaCoDa is provided as an R package, the main computational work is implemented in C++. Because R does not provide native C++ support, Rcpp was employed to expose whole C++ classes as modules to R (EDELBUETTEL and FRANCOIS, 2011). Using Rcpp eliminates time consuming data transfers between the R environment and the C++ core during model fitting, resulting in improved computational performance and allows for a fully object-oriented code design (Booch, 1993). As expected, the runtime of AnaCoDa scales linearly with genome size and number of iterations, and scales polynomially with the number of mixture distributions in the data set. The polynomial increase in runtime with the number of mixture distributions is due to the necessity to condition the gene assignment on the estimation of gene specific parameters, such as, protein synthesis rate.

## **Data Simulation**

In addition to fitting the models to datasets, **AnaCoDa** can be used to generate simulated data sets as well. On their own, simulated datasets are useful for model development and analysis. Simulating data under different conditions allows the user to explore model behavior and explore theoretical scenarios. Different conditions can include the addition or elimination of parameters, or simply allowing a set of parameter values to vary. Fitting models to simulated data can provide insight into potential pitfalls or shortcomings when fitting observational data and can serve as the basis for evaluating model adequacy of a model fit to observational data (**MI** et al., 2015). Significant differences between simulated

and observational data suggests the current set of parameters or the model as a whole fail to include or adequately represent biological mechanisms underlying the observed data.

#### Available models

AnaCoDa currently provides codon models for analyzing genome scale data. The ROC model implements and extends the codon usage bias (CUB) models developed by GILCHRIST et al. (2015); Wallace et al. (2013); Shah and GILCHRIST (2011b), which can reliably estimate the strength of selection on ribosome overhead cost, mutation bias and allows for the inference of protein synthesis rates. This model allows for the separation of effects of mutation and selection based on gene ordering by protein synthesis rate, and the addition of a mixture distribution allows for gene clustering based on mutation bias and selection for translation efficiency. In addition to identifying the most efficient codons, ROC provides estimates of mutation bias allowing the approximation of mutation ratios between codons (GILCHRIST et al., 2015; Wallace et al., 2013).

The ability to estimate protein synthesis rates in the absence of empirical data is useful for investigating CUB of non-model organisms for which such data is lacking and enables the usage of protein synthesis rate in comparative frameworks or other analyses requiring protein synthesis rate information (Dunn et al., 2018). Use of the mixture model allows for the investigation of CUB heterogeneity at the genome or gene level. Following the same framework, additional models included in **AnaCoDa** provide estimates of codon-specific nonsense errors rates (FONSE) and ribosome pausing times (PA and PANSE).

Parameters estimated with the evolutionary models ROC and FONSE represent evolutionary averages and do not depend on experimental conditions. In contrast, PA and PANSE estimate the distribution of biologically relevant parameters like ribosome pausing times along a gene from experimental data such as ribosome footprinting data. The distribution can be dependent (PANSE) or independent (PA) of evidence for nonsense errors in the data.

# 2.4 Appendix: Supplementary Material

AnaCoDa allows for the estimation of biologically relevant parameters like mutation bias or ribosome pausing time, depending on the model employed. Bayesian estimation of parameters is performed using an adaptive Metropolis-Hasting within Gibbs sampling approach. Models implemented in AnaCoDa are currently able to handle gene coding sequences and ribosome footprinting data.

#### 2.4.1 The AnaCoDa framework

The AnaCoDa framework works with gene specific data such as codon frequencies or position specific footprint counts. Conceptually, AnaCoDa uses three different types of parameters.

- The first type of parameters are **gene specific parameters** such as gene expression level or functionality. Gene-specific parameters are estimated separately for each gene and can vary between potential gene categories or sets.
- The second type of parameters are **gene-set specific parameters**, such as mutation bias terms or translation error rates. These parameters are shared across genes within a set and can be exclusive to a single set or shared with other sets. While the number of gene sets must be pre-defined by the user, set assignment of genes can be pre-defined or estimated as part of the model fitting. Estimation of the set assignment provides the probability of a gene being assigned to a set allowing the user to asses the uncertainty in each assignment.
- The third type of parameters are **hyperparameters**, such as parameters controlling the prior distribution for mutation bias or error rate. Hyperparameters can be set specific or shared across multiple sets and allow for the construction and analysis of hierarchical models, by controlling prior distributions for gene or gene-set specific parameters.

## Analyzing protein coding gene sequences

AnaCoDa always requires the following four objects:

- **Genome** contains the codon data read from a fasta file as well as empirical protein synthesis rate in the form of a comma separated (.csv) ID/Value pairs.
- Parameter represents the parameter set (including parameter traces) for a given genome. The parameter object also hold the mapping of parameters to specified sets.
- Model allows you to specify which model should be applied to the genome and the parameter object.
- MCMC specifies how many samples from the posterior distribution of the specified model should be stored to obtain parameter estimates.

## 2.4.2 AnaCoDa setup

#### Application of codon model to single genome

In this example we are assuming a genome with only one set of gene-set specific parameters, hence **num.mixtures** = 1. We assign all genes the same gene-set, and provide an initial value for the hyperparameter sphi  $(s_{\phi})$ .  $s_{\phi}$  controls the lognormal prior distribution on the gene specific parameters like the protein synthesis rate  $\phi$ . To ensure identifiability the expected value of the prior distribution is assumed to be 1.

$$E[\phi] = \exp\left(m_{\phi} + \frac{s_{\phi}^2}{2}\right) = 1 \tag{2.1}$$

Therefore the mean  $m_{\phi}$  is set to be  $-\frac{s_{\phi}^2}{2}$ . For more details see GILCHRIST *et al.* (2015).

After choosing the model and specifying the necessary arguments for the MCMC routine, the MCMC is run

```
genome <- initializeGenomeObject(file = "genome.fasta")</pre>
```

runMCMC does not return a value, the results of the MCMC are stored automatically in the mcmc and parameter objects created earlier.

Please note that AnaCoDa utilizes C++ object orientation and therefore employs pointer structures. This means that no return value is necessary for such objects as they are modified within the the runMCMC routine. You will find that after a completed run, the parameter object will contain all necessary information without being directly passed into the MCMC routine. This might be confusing at first as it is not default R behavior.

#### Application of codon model to a mixture of genomes

This case applies if we assume that parts of the genome differ in their gene-set specific parameters. This could be due to introgression events or strand specific mutation difference, horizontal gene transfers or other reasons. We make the assumption that all sets of genes are independent of one another. For two sets of gene-set specific parameter with a random gene assignment we can use:

To accommodate for this mixing we only have to adjust **sphi**, which is now a vector of length 2, **num.mixtures**, and **gene.assignment**, which is chosen at random here.

## Empirical protein synthesis rate values

To use empirical values as prior information one can simply specify an observed expression file when initializing the genome object.

These observed expression or synthesis values  $(\Phi)$  are independent of the number of genesets. The error in the observed  $\Phi$  values is estimated and described by sepsilon  $(s_{\epsilon})$ . The csv file can contain multiple observation sets separated by comma. For each set of observations an initial  $s_{\epsilon}$  has to be specified.

In addition one can choose to keep the noise in the observations  $(s_{\epsilon})$  constant by using the fix.observation.noise flag in the model object.

```
model <- initializeModelObject(parameter = parameter, model = "ROC",
fix.observation.noise = TRUE)</pre>
```

#### Fixing parameter types

It can sometime be advantages to fix certain parameters, like the gene specific parameters. For example in cases where only few sequences are available but gene expression measurements are at hand we can fix the gene specific parameters to increase confidence in our estimates of gene-set specific parameters.

We again initialize the **genome**, **parameter**, and **model** objects.

To fix gene specific parameters we will set the **est.expression** flag to **FALSE**. This will estimate only gene-set specific parameters, hyperparameters, and the assignments of genes to various sets.

If we would like to fix gene-set specific parameters we instead disable the **est.csp** flag.

The same applies to the hyper parameters (est.hyper),

and gene set assignment (est.mix).

We can use these flags to fix parameters in any combination.

## Combining various gene-set specific parameters to a gene-set description.

We distinguish between three simple cases of gene-set descriptions, and the ability to customize the parameter mapping. The specification is done when initializing the parameter object with the **mixture.definition** argument.

We encounter the simplest case when we assume that all gene sets are independent.

The **allUnique** keyword allows each type of gene-set specific parameter to be estimated independent of parameters describing other gene sets.

In case we want to share mutation parameter between gene sets we can use the keyword mutationShared

This will force all gene sets to share the same mutation parameters.

The same can be done with parameters describing selection, using the keyword selectionShared

For more intricate compositions of gene sets, one can specify a custom  $n \times 2$  matrix, where n is the number of gene sets, to describe how gene-set specific parameters should be shared. Instead of using the **mixture.definition** argument one uses the **mixture.definition.matrix** argument.

The matrix representation of **mutationShared** can be obtained by

Columns represent mutation and selection, while each row represents a gene set. In this case we have three gene sets, each sharing the same mutation category and three different selection categories. In the same way one can produce the matrix for three independent gene sets equivalent to the allUnique keyword.

```
# [,1] [,2]
```

```
#[1,] 1 1
#[2,] 2 2
#[3,] 3 3
defMatrix <- matrix(c(1,2,3,1,2,3), ncol=2)</pre>
```

We can also use this matrix to produce more complex gene set compositions.

```
# [,1] [,2]
#[1,] 1 1
#[2,] 2 1
#[3,] 1 2
defMatrix <- matrix(c(1,2,1,1,1,2), ncol=2)</pre>
```

In this case gene set one and three share their mutation parameters, while gene set one and two share their selection parameters.

#### Checkpointing

AnaCoDa does provide checkpointing functionality in case runtime has to be restricted. To enable checkpointing, one can use the function **setRestartSettings**.

```
# writing a restart file every 1000 samples
setRestartSettings(mcmc, "restart_file", 1000, write.multiple=TRUE)
# writing a restart file every 1000 samples
# but overwriting it every time
setRestartSettings(mcmc, "restart_file", 1000, write.multiple=FALSE)
```

To re-initialize a parameter object from a restart file one can simply pass the restart file to the initialization function:

```
initializeParameterObject(init.with.restart.file="restart_file.rst")
```

#### Load and save parameter objects

AnaCoDa is based on C++ objects using the Rcpp (EDELBUETTEL and FRANCOIS, 2011). This comes with the problem that C++ objects are by default not serializable and can therefore not be saved/loaded with the default R save/load functions.

AnaCoDa however, does provide functions to load and save parameter and mcmc objects.

These are the only two objects that store information during a run.

```
#save objects after a run
runMCMC(mcmc = mcmc, genome = genome, model = model)
writeParameterObject(parameter = parameter, file = "parameter.Rda")
writeMCMCObject(mcmc = mcmc, file = "mcmc_out.Rda")
```

As **genome**, and **model** objects are purely storage containers, no save/load function is provided at this point, but will be added in the future.

```
#load objects
parameter <- loadParameterObject(file = "parameter.Rda")
mcmc <- loadMCMCObject(file = "mcmc_out.Rda")</pre>
```

#### 2.4.3 File formats

#### Protein coding sequence

Protein coding sequences are provided by fasta file with the default format. One line containing the sequence id starting with > followed by the id and one or more lines containing the sequence. The sequences are expected to have a length that is a multiple of three. If a codon can not be recognized (e.g AGN) it is ignored.

>YALOO1C

TTGGTTCTGACTCATTAGCCAGACGAACTGGTTCAA
CATGTTTCTGACATTCATTCTAACATTGGCATTCAT

ACTCTGAACCAACTGTAAGACCATTCTGGCATTTAG

>YALOO2W

TTGGAACAAACGGCCTGGACCACGACTCACGCTCT

TCACATGACACTACTCATAACGACACTCAAATTACT

TTCCTGGAATTCCGCTCTTAGACTCAACTGTCAGAA

Empirical gene expression

Empirical expression or gene specific parameters are provided in a csv file format. The first

line is expected to be a header describing each column. The first column is expected be

the gene id, and every additional column is expected to be represent a measurement. Each

row corresponds to one gene and contains all measurements for that gene, including missing

values.

>YALOO1C

ORF, DATA\_1, DATA\_2, ... DATA\_N

YAL001C,0.254,0.489,...,0.156

YAL002W, 1.856, 1.357, ..., 2.014

YAL003W,10.45,NA,...,9.564

YAL005C, 0.556, 0.957, ..., 0.758

Ribosome foot-printing counts

Ribosome foot-printing (RFP) counts are provided in a csv file format. The first line is

expected to be a header describing each column. The columns are expected in the following

order gene id, position, codon, rfpcount. Each row corresponds to a single codon with an

associated number of ribosome footprints.

GeneID, Position, Codon, rfpCount

YBR177C, 0, ATA, 8

23

```
YBR177C, 1, CGG, 1
YBR177C, 2, GTT, 8
YBR177C, 3, CGC, 1
```

## 2.4.4 Analyzing and Visualizing results

#### Parameter estimates

After we have completed the model fitting, we are interested in the results. AnaCoDa provides functions to obtain the posterior estimate for each parameter. For gene-set specific parameters or codon specific parameters we can use the function **getCSPEstimates**. Again we can specify for which mixture we would like the posterior estimate and how many samples should be used. **getCSPEstimates** has an optional argument filename which will cause the routine to write the result as a csv file instead of returning a **data.frame**.

To obtain posterior estimates for the gene specific parameters, we can use the function **getExpressionEstimatesForMixture**. In the case below we ask to get the gene specific parameters for all genes, and under the assumption each gene is assigned to mixture 1.

However we can decide to only obtain certain gene parameters. in the first case we sample 100 random genes.

Furthermore, AnaCoDa allows to calculate the selection coefficients for each codon and each gene. We can use the function **getSelectionCoefficients** to do so. Please note, that this function returns the  $\log(sN_e)$ .

**getSelectionCoefficients** returns a matrix with  $log(sN_e)$  relative to the most efficient synonymous codon.

```
#SAKLOA00154g -0.8494558 -0.045305847 -1.0930388 0 -0.5285367 ...

#SAKLOA00176g -0.4455753 -0.023764823 -0.5733448 0 -0.2772397 ...

#SAKLOA00198g -0.3926068 -0.020939740 -0.5051875 0 -0.2442824 ...

#SAKLOA00220g -0.9746002 -0.051980440 -1.2540685 0 -0.6064022 ...

#SAKLOA00242g -0.3691110 -0.019686586 -0.4749542 0 -0.2296631 ...
```

We can compare these values to the weights from the codon adaptation index (CAI) citepSharp1987 or effective number of codons ( $N_c$ ) (WRIGHT, 1990) by using the functions getCAIweights and getNcAA.

```
caiWeights <- getCAIweights(referenceGenome = genome)
head(caiWeights)

# GCA GCC GCG GCT TGC TGT

#0.7251276 0.6282192 0.2497737 1.0000000 0.6222628 1.0000000
nc.per.aa <- getNcAA(genome = genome)
head(nc.per.aa)

# A C D E F G ...

#SAKLOA00132g 3.611111 1.000000 2.200000 2.142857 1.792453 ...

#SAKLOA00154g 1.843866 2.500000 2.035782 1.942505 1.986595 ...

#SAKLOA00176g 5.142857 NA 1.857143 1.652174 1.551724 3.122449 ...

#SAKLOA00198g 3.800000 NA 1.924779 1.913043 2.129032 4.136364 ...

#SAKLOA00220g 3.198529 1.666667 1.741573 1.756757 2.000000 ...

#SAKLOA00242g 4.500000 NA 2.095890 2.000000 1.408163 3.734043 ...</pre>
```

We can also compare the distribution of selection coefficients to the CAI values estimated from a reference set of genes. Figure 2.1, produced by the code below, shows that selection coefficients for the same codon can vary greatly between the genes.

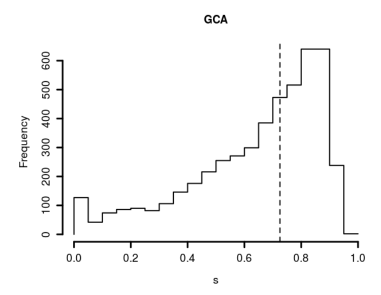


Figure 2.1: Distribution of s for codon GCA for amino acid alanine. Dashed line indicates the CAI weight for GCA. The comparison provides a more nuanced picture as we can see that the selection on GCA varies across the genome.

#### **Diagnostic Plots**

A first step after every run should be to determine if the sampling routine has converged. To do that, AnaCoDa provides plotting routines to visualize all sampled parameter traces from which the posterior sample is obtained (Figure 2.2). First we have to obtain the **trace** object stored within our **parameter** object. Now we can simply plot the **trace** object. The argument **what** specifies which type of parameter should be plotted. Here we plot the selection parameter  $\Delta \eta$  of the ROC model. These parameters are mixture specific and one can decide which mixture set to visualize using the argument **mixture**.

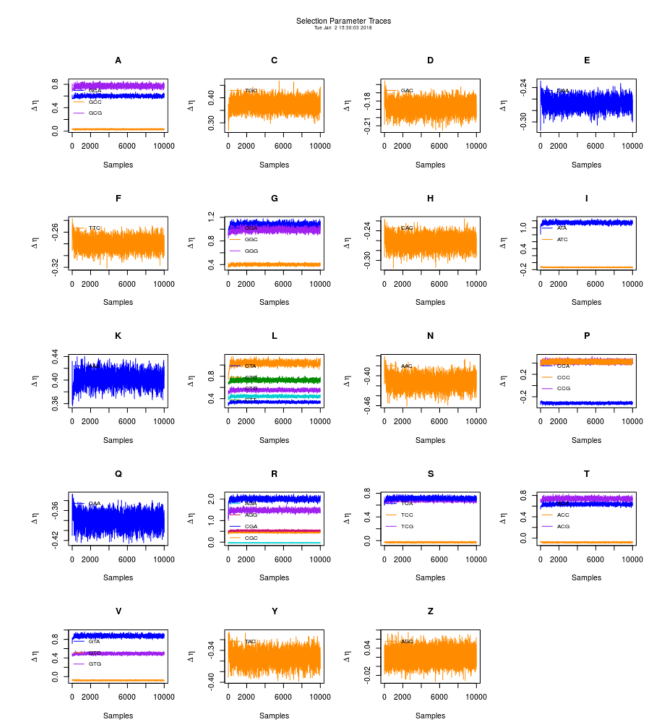


Figure 2.2: Trace plot showing the traces of all 40 codon specific selection parameters  $\Delta \eta$  organized by amino acid.

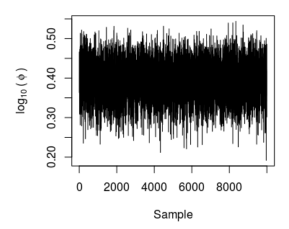


Figure 2.3: Trace plot showing the protein synthesis trace  $\phi$  for gene 669.

```
trace <- getTrace(parameter)
plot(x = trace, what = "Selection", mixture = 1)</pre>
```

A special case is the plotting of traces of the protein synthesis rate  $\phi$  (Figure 2.3). As the number of traces for the different  $\phi$  traces is usually in the thousands, a **geneIndex** has to be passed to determine for which gene the trace should be plotted. This allows to inspect the trace of every gene under every mixture assignment.

```
trace <- parameter$getTraceObject()
plot(x = trace, what = "Expression", mixture = 1, geneIndex = 669)</pre>
```

We find the likelihood and posterior trace of the model fit in the **mcmc object**. The trace can be plotted by just passing the **mcmc** object to the **plot** routine. Again we can switch between  $\log(likelihood)$  and  $\log(posterior)$  using the argument **what**. The argument **zoom.window** is used to inspect a specified window in more detail. It defaults to the last 10 % of the trace. The  $\log(posterior)$  displayed in the figure title is estimated over the **zoom.window** (Figure 2.4).

```
plot(mcmc, what = "LogPosterior", zoom.window = c(9000, 10000))
```

#### log(Posterior Probability): -2605710.60092172

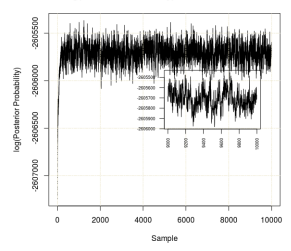


Figure 2.4: Trace plot showing the log(posterior) trace for the current model fit. Window inset shows the last 1.000 samples

#### Model visualization

We can visualize the results of the model fit by plotting the **model** object (Figure 2.5). For this we require the model and the **genome** object. We can adjust which mixture set we would like to visualize and how many samples should be used to obtain the posterior estimate for each parameter. For more details see GILCHRIST *et al.* (2015).

```
# use the last 500 samples from mixture 1 for posterior estimate. plot(x = model, genome = genome, samples = 500, mixture = 1)
```

As AnaCoDa is designed with the idea to allow gene-sets to have independent gene-set specific parameters, AnaCoDa also provides the option to compare different gene-sets by plotting the parameter object. Figure 2.6 allows us to compare the selection parameter estimated by ROC for seven yeast species. The code below illustrates how the figure is plotted.

```
# use the last 500 samples from mixture 1 for posterior estimate.
plot(parameter, what = "Selection", samples = 500)
```



Figure 2.5: Fit of the ROC model for a random yeast. The solid line represent the model fit from the data, showing how synonymous codon frequencies change with gene expression. The points are the observed mean frequencies of a codon in that synthesis rate bin and the whisks indicate the standard deviation within the bin. The codon favored by selection is indicated by a "\*". The bottom right panel shows how many genes are contained in each bin

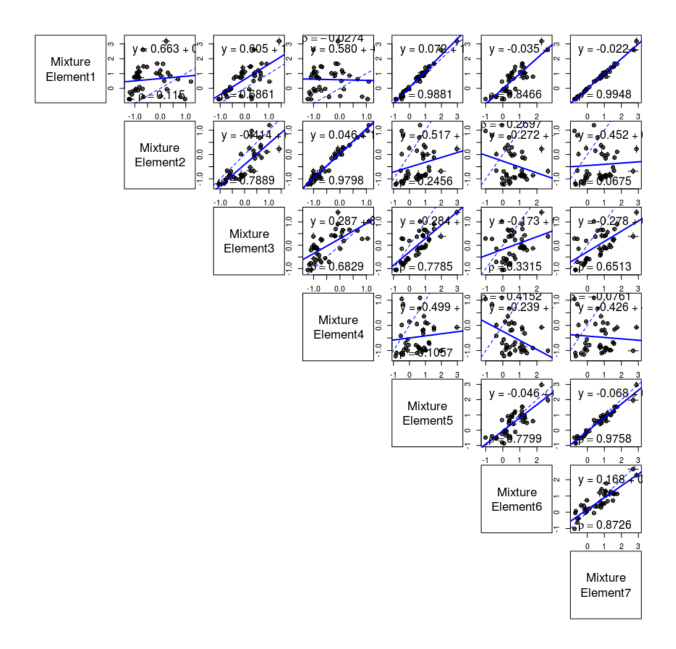


Figure 2.6: Comparison of the selection parameter of seven yeast species estimated with  ${\hbox{ROC-SEMPPR}}.$ 

## Chapter 3

Decomposing mutation and selection to identify mismatched  ${\rm codon\ usage}$ 

This chapter is a lightly revised version of a paper to be submitted to Genome Biology and Evolution and co-authored with Michael A. Gilchrist and Russel Zaretzki.

C. Landerer, R. Zaretzki, M.A. Gilchrist, Decomposing mutation and selection to identify mismatched codon usage

### 3.1 Abstract

For decades, codon usage has been used as a measure of adaptation for translational efficiency of a gene's coding sequence. These patterns of codon usage reflect both the selective and mutational environment in which the coding sequences evolved. Over this same period, gene transfer between lineages has become widely recognized as an important biological pheonmena. Nevertheless, most studies of codon usage implicitly assume that all genes within a genome evolved under the same selective and mutational environment, an assumption violated when introgression occurs. In order to better understand the effects of introgression on codon usage patterns and vice versa, we examine the patterns of codon usage in the yeast which has experienced a large introgression, Lachancea kluyveri. We quantify the effects of mutation bias and selection for translation efficiency on the codon usage pattern of the endogenous and introgressed exogenous genes using a Bayesian mixture model, ROC SEMPPR, which is built on mechanistic assumptions of protein synthesis and grounded in population genetics. We find substantial differences in codon usage between the endogenous and exogenous genes, and show that these differences can be largely attributed to a shift in mutation bias from A/T ending codons in the endogenous genes to C/G ending codons in the exogenous genes. Recognizing the two different signatures of mutation and selection bias improves our ability to predict protein synthesis rate by 17% and allowed us to accurately assess codon preferences. In addition, using our estimates of mutation and selection bias, we to identify  $Eremothecium\ gossypii$  as the most likely source lineage, estimate the introgression occurred  $\sim 6\times 10^8$  generation ago, and estimate its historic and current genetic load. Together, our work illustrates the advantage of mechanistic, population genetic models like ROC SEMPPR and the quantitative estimates they provide when analyzing sequence data.

## 3.2 Introduction

Synonymous codon usage patterns varies within a genome and between taxa, reflecting differences in mutation bias, selection, and genetic drift. The signature of mutation bias is largely determined by the organism's internal or cellular environment, such as their DNA repair genes or UV exposure. While this mutation bias is an omnipresent evolutionary force, its impact can be obscured or even amplified by selection. The signature of selection on codon usage is also largely determined by an organism's cellular environment, such as its tRNA species, their copy number, and post-transcriptional modifications. The strength of selection on the codon usage of an individual gene is largely determined by its expression level which, in turn, is also largely determined by the organism's external environment. In general, the strength of selection on codon usage increases with its expression level (Gouy and GAUTIER, 1982; IKEMURA, 1985; BULMER, 1990), specifically its protein synthesis rate (GILCHRIST, 2007). Thus as gene expression increases, codon usage shifts from a process dominated by mutation to a process dominated by selection. The overall efficacy of selection on codon usage is a function of the organism's effective population size  $N_e$  which, in turn, is largely determined by its external environment. By explicitly modeling the combined forces of mutation, selection, and drift, ROC SEMPPR allows us disentangle the evolutionary forces responsible for the patterns of codon usage bias (CUB) encoded in an species' genome (GILCHRIST, 2007; SHAH and GILCHRIST, 2011a; WALLACE et al., 2013; GILCHRIST et al., 2015), should provide biologically meaningful information about the lineage's historical cellular and external environment.

Most studies implicitly assume that the CUB of a genome is shaped by a single cellular environment. As genes are horizontally transferred, introgress, or combined to form novel hybrid species, one would expect to see the influence of multiple cellular environments on a genomes codon usage pattern (MDIGUE et al., 1991; LAWRENCE and OCHMAN, 1997). Given that transferred genes are likely to be less adapted than endogenous genes to their new cellular environment, we expect a greater fitness burden of transferred genes if donor

and recipient environment differ greatly in their selection bias, making such transfers less likely. More practically, if differences in codon usage of transferred genes are unaccounted for, they may distort parameter estimates. Such distortion could lead to the wrong codon preference for an amino acid, underestimate the variation in protein synthesis rate, or bias mutation estimates when analyzing a genome.

To illustrate these ideas, we analyze the CUB of the genome of Lachancea kluyveri, the earliest diverging lineage of the Lachancea clade. The Lachancea clade diverged from the Saccharomyces clade, prior to its whole genome duplication ~ 100 Mya ago (MARCET-HOUBEN and GABALDN, 2015; BEIMFORDE et al., 2014). Since that time, L. kluyveri has experienced a large introgression of exogenous genes found in all populations (FRIEDRICH et al., 2015). The introgression replaced the left arm of the C chromosome and displays a 13% higher GC content than the endogenous L. kluyveri genome (PAYEN et al., 2009; FRIEDRICH et al., 2015). These characteristics make L. kluyveri an ideal model to study the effects of an introgressed cellular environment and the resulting mismatch in codon usage.

Using ROC SEMPPR, a Bayesian population genetics model based on a mechanistic description of ribosome movement along an mRNA, allows us to quantify the cellular environment in which genes have evolved by separately estimating the effects of mutation bias and selection bias on codon usage. ROC SEMPPR's resulting predictions of protein synthesis rates have been shown to be on par with laboratory measurements (Shah and Gilchrist, 2011a; Gilchrist et al., 2015). We use ROC SEMPPR to independently describe two cellular environments reflected in the *L. kluyveri* genome; the signature of the current environment in the endogenous genes and the decaying signature of the exogenous environment in the introgressed genes. Our results indicate that the difference in GC content between endogenous and exogenous genes is mostly due to the differences in mutation bias of their ancestral environments. Accounting for these different signatures of mutation bias and selection bias of the endogenous and exogenous sets of genes substantially improves our ability to predict present day protein synthesis rates. These endogenous and exogenous gene

set specific estimates of mutation bias and selection bias, in turn allow us to address more refined questions of biological importance. For example, it allows us to identify *E. gossypii* as the most likely source of the introgressed genes out of the 38 yeast lineages with sequenced genomes, estimate the age of the introgression to be on the order of 0.2-1 Mya, estimate the genetic load of these genes, both at the time of introgression and now, as well as make predictions about how the CUB of the introgressed genes will evolve in the future.

### 3.3 Results

# 3.3.1 The Signatures of two Cellular Environments within L. kluyveri's Genome

We used our software package AnaCoDa (Landerer et al., 2018) to compare model fits of ROC SEMPPR to the entire L. kluyveri genome and its genome partitioned into two sets of 4,864 endogenous and 497 exogenous genes. AIC values strongly support the hypothesis that the L. kluyveri genome consists of genes with two different and distinct patterns of codon usage bias ( $\Delta$ AIC = 75,462; Table 4.1). We find additional support for this hypothesis when we compare our predictions of gene expression to empirically observed values. Specifically, the explanatory power between our predictions and observed values improved by  $\sim$  42%, from  $R^2 = 0.33$  to 0.46 (Figure 3.1).

Table 3.1: Model selection of the two competing hypothesis. Reported are the log-likelihood,  $\log(\mathcal{L})$ , the number of parameters estimated n, AIC, and  $\Delta$ AIC values.

| Hypothesis | $\log(\mathcal{L})$ | n     | AIC       | $\Delta AIC$ |
|------------|---------------------|-------|-----------|--------------|
| Separated  | -2,612,397          | 5,402 | 5,235,598 | 0            |
| Combined   | -2,650,047          | 5,483 | 5,311,060 | $75,\!462$   |

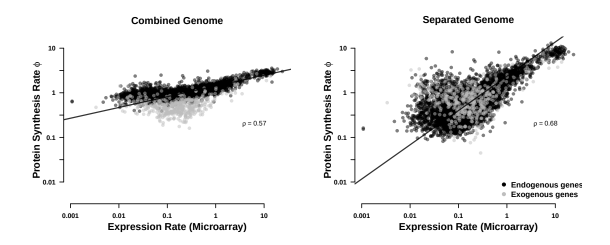


Figure 3.1: Comparison of predicted protein synthesis rate  $\phi$  to microarray data from TSANKOV et al. (2010) for (a) the combined genome and (b) the separated endogenous and exogenous genes. Endogenous genes are displayed in black and exogenous genes in red. Black line indicates type II regression line SOKAL and ROHLF (1981).

# 3.3.2 Comparing Differences in the Endogenous and Exogenous Codon Usage

To better understand the differences in the endogenous and exogenous cellular environments, we compared our parameter estimates of mutation bias  $\Delta M$  and selection  $\Delta \eta$  for the two sets of genes. Our estimates of  $\Delta M$  for the endogenous and exogenous genes were negatively correlated ( $\rho = -0.49$ ), indicating weak concordance of  $\sim 5\%$  between the two mutation environments (Figure 3.2). For example, the endogenous genes show a mutational preference for A and T ending codons in  $\sim 95\%$  of the codon families. In contrast, the exogenous genes display an equally consistent mutational preference towards C and G ending codons (Table 3.2). As a result, only the two codon amino acid Phenylalanine (Phe, F) shares the same rank order across the endogenous and exogenous  $\Delta M$  estimates.

In contrast, our estimates of  $\Delta \eta$  for the endogenous and exogenous genes were positively correlated ( $\rho = 0.69$ ) and showing concordance of  $\sim 53\%$  between the two selection environments (Figure 3.2). We find that the strength of selection within each codon family differs between sets of genes. Overall, the endogenous genes only show a selection preference

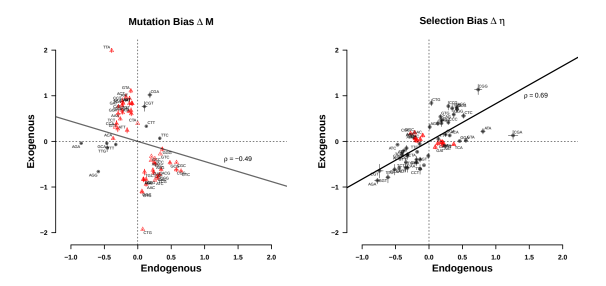


Figure 3.2: Comparison of (a) mutation bias  $\Delta M$  and (b) selection bias  $\Delta \eta$  parameters for endogenous and exogenous genes. Estimates are relative to the mean for each codon family. Black dots indicate  $\Delta M$  or  $\Delta \eta$  parameters with the same sign for the endogenous and exogenous genes, red dots indicate parameters with different signs. Black line shows the type II regression line (Sokal and Rohlef, 1981). Dashed lines mark quadrants.

for C and G ending codons in  $\sim 58\%$  of the codon families. In contrast, the exogenous genes display a strong preference for A and T ending codons in  $\sim 89\%$  of the codon families.

The difference in codon usage between endogenous and exogenous genes is striking. As a result, our estimates of the optimal codon differ in nine cases between endogenous and exogenous genes (Table 3.3). Fits to the complete L. kluyveri genome reveal that the relatively small exogenous gene set ( $\sim 10\%$  of genes) has a disproportional effect on the model fit. We find that the complete L. kluyveri genome is estimated to share the mutational preference with the exogenous genes in  $\sim 78\%$  of the 19 codon families that are discordant between the endogenous and exogenous genes. In two cases, Isoleucine (Ile, I) and Arginine (Arg, R), the strong discordance in mutation preference results in an estimated codon preference in the complete L. kluyveri genome that differs from both the endogenous, and the exogenous genes.

The effect of the small exogenous gene set on the fit to the complete L. kluyveri genome is smaller in our estimates of selection bias  $\Delta \eta$  than  $\Delta M$ , but still large. We find that

the complete L. kluyveri genome is estimated to share the selection preference with the exogenous genes in  $\sim 60\%$  of codon families that show discordance between endogenous and exogenous genes. These results clearly show that it is important to recognize the difference in endogenous and exogenous genes and treat these genes as separate sets to avoid the inference of incorrect synonymous codon preferences and better predict protein synthesis.

#### 3.3.3 Determining Source of Exogenous Genes

We combined our estimates of mutation bias  $\Delta M$  and selection bias  $\Delta \eta$  with synteny information and searched for potential source lineages of the introgressed exogenous region. We examined 38 yeast lineages (Table 3.4) of which two (*Eremothecium gossypii* and *Candida dubliniensis*) showed a strong positive correlation in codon usage (Figure 3.3). The endogenous *L. kluyveri* genome exhibits codon usage very similar to most yeast lineages examined, indicating little variation in codon usage among the examined yeasts (Figure 3.5). Four lineages show a positive correlation for  $\Delta M$  and  $\Delta \eta$  with the exogenous genes and have a weak to moderate positive correlation in selection bias with the endogenous genes; but, like the exogenous genes, tend to have a negative correlation in  $\Delta M$  with the endogenous genes.

Comparing synteny between the exogenous genes, which are restricted to the left arm of chromosome C, and E. gossypii and C. dubliniensis as well as closely related yeast species we find that E. gossypii displays the highest synteny (Figures 3.7 & 3.8). C. dubliniensis, even though it displays similar codon usage does not show synteny with the exogenous region. Furthermore, the synteny relationship between the exogenous region and other yeasts appears to be limited to the Saccharomycetacease clade (Figure 3.8). Given these results, we conclude that of the 38 examined yeast lineages the E. gossypii lineage is the most likely source of the introgressed exogenous genes.

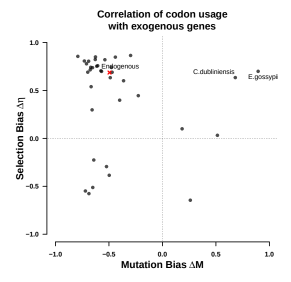


Figure 3.3: Correlation of  $\Delta M$  and  $\Delta \eta$  of the exogenous genes with 38 examined yeast lineages. Dots indicate the correlation of  $\Delta M$  and  $\Delta \eta$  of the lineages with the endogenous and exogenous parameter estimates. All regressions were performed using a type II regression (SOKAL and ROHLF, 1981).

#### 3.3.4 Estimating Introgression Age

We modeled the change in codon frequency as a model of exponential decay, we estimated the age of the introgression assuming that E. gossypii still represents the mutation bias of its ancestral source lineage at the time of the introgression and a constant mutation rate. We infer the age of the introgression to be on the order of  $6.2 \pm 1.2 \times 10^8$  generations. Assuming L. kluyveri experiences between one and eight generations per day, we estimate the introgression to have occurred between 212,000 to 1,700,000 years ago. Our estimate places the time of the introgression earlier than previously assumed (FRIEDRICH et al., 2015).

Using the same approach, we also estimated the persistence of the signal of the exogenous cellular environment. We assume that differences in mutation bias will decay more slowly than differences in selection bias to be able to utilize our bias free estimates of  $\Delta M$ . We predict that the  $\Delta M$  signal of the source cellular environment will have decayed to be within one percent of the L. kluyveri environment in  $\sim 5.4 \pm 0.2 \times 10^9$  generations, or

between 1,800,000 and 15,000,000 years. Together, these results indicate that the mutation signature of the exogenous genes will persist for a very long time.

## 3.3.5 Genetic Load due to Mismatching Codon Usage of the Exogenous Genes

We define genetic load as the difference between the fitness of an expected, replaced endogenous gene and the exogenous gene,  $s \propto \phi \Delta \eta$  due to the mismatch in codon usage parameters (See Methods for details). Estimates of selection bias for the exogenous genes show that, while well correlated with the endogenous genes, only nine amino acids share the same optimal codon. Exogenous genes are, therefore, expected to represent a significant reduction in fitness, or genetic load for L. kluyveri due to this mismatch in codon usage. As the introgression occurred before the diversification of L. kluyveri and has fixed throughout all populations (FRIEDRICH et al., 2015), we can not observe the original endogenous sequences that have been replaced by the introgression. Using our estimates of  $\Delta M$  and  $\Delta \eta$  from the endogenous genes and assuming hat the current exogenous amino acid composition of genes is representative of the replaced endogenous genes, we estimate the genetic load of the exogenous genes at the time of introgression (Figure 3.4a) and currently (Figure 3.4b). We find that the genetic load due to mismatched codon usage was -0.0008 at the time of the introgression and still represents a genetic load of -0.0003 today.

In order to account for differences in the efficacy of selection on codon usage between the donor lineage and L. kluyveri using a linear scaling factor  $\kappa$  (See Methods for details). We predict that a small number of low expression genes ( $\phi < 1$ ) were weakly exapted at the time of the introgression (Figure 3.4a). High expression genes ( $\phi > 1$ ) are predicted to have carried the largest genetic load in the novel cellular environment. These highly expressed genes are inferred to have the greatest degree of adaptation since the time of the introgression to the L. kluyveri cellular environment (Figures 3.4a & 3.10).

Explicitly mention value for genetic load

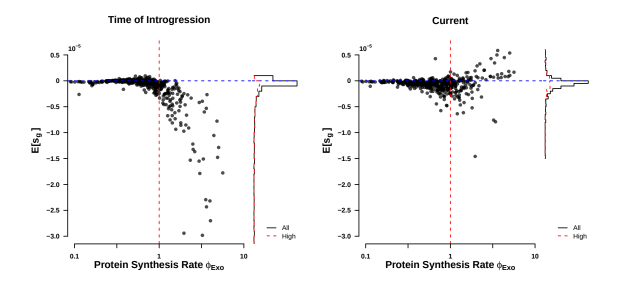


Figure 3.4: Genetic load  $s = \Delta \eta \phi$  (a) at the time of introgression ( $\kappa = 5$ ), and (b) currently ( $\kappa = 1$ ).

## 3.4 Discussion

In order to study the evolutionary effects of an introgression, we used ROC SEMPPR, a mechanistic model of ribosome movement along an mRNA. Our parameter estimates indicate that the L. kluyveri genome contains distinct signatures of mutation and selection bias from both an endogenous and exogenous cellular environment. By fitting ROC SEMPPR separately to L. kluyveri's endogenous and exogenous sets of genes we generate a quantitative description of their signatures of mutation bias and natural selection for efficient protein translation. Our results indicate that the difference in GC content between endogenous and exogenous genes is mostly due to differences in mutation bias, but we also show that the strength and rank order of selection within a codon family differ between endogenous and exogenous cellular environments. Even though the exogenous genes make up only  $\sim 10\%$  of the L. kluyveri genome, when we fail to recognize these differences our estimates of  $\Delta M$  and  $\Delta \eta$  deviate substantial from their actual values (Figure 3.6). While this sensitivity of our parameters to a second cellular environment may be surprising, it highlights the importance of recognizing different cellular environments reflected by a genome. Furthermore, our results

indicate that we can attribute the increased GC content in the exogenous genes mostly to differences in mutation bias favoring G/C ending codons rather than selection.

The separation of the endogenous and exogenous genes improves our estimates of protein synthesis rate  $\phi$  by 42% relative to the full genome estimate ( $R^2 = 0.32$  vs. 0.46, respectively). Furthermore, failing to separately analyze the endogenous and exogenous genes results in an unrealistically small amount of intergenic variation in  $\phi$  (compare Figure 3.1a & b). This behavior is due, in part, to constraining  $E[\phi] = 1$  which allows us to compare the efficacy of selection  $sN_e$  across genomes. Extremely small variances in the  $\phi$  values estimated by ROC SEMPPR could indicate that a genome contains the signature of multiple cellular environments.

The mutation and selection bias parameters  $\Delta M$  and  $\Delta \eta$  of the introgressed exogenous genes contain information, albeit decaying, about its previous cellular environment. We, therefore, utilize  $\Delta M$  and  $\Delta \eta$  to identify potential source lineages. The E. gossypii and C. dubliniensis lineages stand out from the other 36 yeast lineages in that the correlation coefficients between their  $\Delta M$  and  $\Delta \eta$  parameters and those of the exogenous genes are > 0.5 (Figure 3.2). In terms of gene order, we found that synteny with the exogenous genes is limited to the Saccharomycetaceae clade, which C. dubliniensis is outside of. Overall, the synteny coverage extends along the whole exogenous regions with the exception of the 3' and 5' ends of the exogenous region (Figure 3.8b). Further, of the 38 species examined, strength E. qossypii is the only genome with a GC content > 50%, making it most similar to the of exogenous genes. Thus, only the E. qossypii genome displays strong correlations in  $\Delta M$  and selection  $\Delta \eta$ , synteny, and similar GC content with the exogenous genes.

With E. gossypii identified as potential source lineage of the introgressed region, we inferred the time since the introgression occurred using our estimates of mutation bias  $\Delta M$ . Our  $\Delta M$  estimates are well suited for this task as they are free of the influence of selection and unbiased by  $N_e$  and other scaling terms, which is in contrast to our estimates of  $\Delta \eta$  (GILCHRIST et al., 2015). Our estimated age of the introgression of  $6.2 \pm 1.2 \times 10^8$ 

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generations is  $\sim 10$  times longer time than a previous minimum estimate by FRIEDRICH et al. (2015) of  $5.6 \times 10^7$  generations. Our estimate assumes that the current E. gossypii and L. kluyveri cellular environment reflect their ancestral states at the time of the introgression. If the ancestral mutation environments were more similar (dissimilar) at the time of the introgression than now our result is an overestimate (underestimate).

In order to estimate the introgression's genetic load due to codon mismatch, we had to make three key assumptions: 1) at the time of introgression the amino acid sequences of the endogenous genes and exogenous genes where highly similar, 2) the current L. kluyveri cellular environment is reflective of the cellular environment at the time of the introgression, and 3) the E. gossypii cellular environment reflects its ancestral environment at the time of the introgression. In general due to their very nature, low expression genes contribute little to the genetic load. Indeed,  $\sim 30\%$  of low expression exogenous genes ( $\phi < 1$ ) appeared to be exapted at the time of the introgression. These exapted genes are likely due to the mutation bias in the endogenous genes matching the selection bias in the exogenous genes for G/C ending codons. In contrast, highly expressed genes are predicted to have imposed a large genetic load. Many of these genes appear to still represent a significant genetic load. Overall, our estimates of codon mismatch genetic load, therefore, suggest strong selection against the introgression.

It is hard to contextualize the probability of this introgression being fixed as we are not aware of any estimates of the frequency at which such large scale introgressions of genes occur. A related example of a large scale merger of genomic material can be found in S. bayanus, which is currently believed to be a hybrid of S. cerevisiae, S. eubayanus, and S. uvarum lineages. Unlike with L. kluyveri and E. gossypii, the progenitor lineages of S. bayanus have similar codon usage parameters. For example, the correlation between  $\Delta M$  and  $\Delta \eta$  for these two lineages are  $\rho = 0.83$  and 0.98 (data not shown). These similarities in  $\Delta M$  and  $\Delta \eta$  parameters suggest that the genetic load for S. bayanus due to codon usage mismatch is small relative to the exongenous genes considered here. The large genetic load

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of the exogenous genes due to codon mismatch at the time of the introgression would seem to indicate that the fixation of the introgression was either a fluke event or the codon mismatch genetic load was countered by one or more highly advantageous loci within the introgression.

Under the first scenario, our best estimate of the selection coefficient against the introgression based on expected codon mismatch at that time is s=-0.0008 and an effective population size  $N_e$  on the order of  $10^8$  (Wagner, 2005) yields an approximate fixation probability of  $(1-\exp[-s])/(1-\exp[2-sN_e])\approx 10^{-6950}$  (Sella and Hirsh, 2005). Even though L. kluyveri diverged from the rest of the Lachancea clade around 85 Mya (Kensche et al., 2008; Marcet-Houben and Gabaldn, 2015), if we assume 1 to 8 generations/day, which implies  $10^{10}$  to  $10^{11}$  generations since the time of divergence, one round of meiosis for every 1000 rounds of mitosis based on S. paradoxus (?), and  $N_e \approx 10^8$  there were only  $10^{15}$  to  $10^{16}$  opportunities for such an introgression to have occurred and fixed. Clearly, unless there was a severe bottleneck with  $N_e < 1/|s| \approx 1,250$  around the time of introgression, which conceivably could have been triggered by a speciation event, this scenario seems very unlikely.

In the second scenario, where we assume the introgression contained advantageous loci, one may wonder why recombination events did not limit the introgression to only the adaptive loci. Payen et al. (2009) found that the exogenous region has a lower rate of recombination, presumably due to the dissimilarity in GC content and/or a lower than average sequence homology between the exogenous region and the one it replaced. Compatible with this explanation is the possibility of several highly advantageous loci distributed across the region which then drove a rapid selective sweep.

Overall, our results show the usefulness of the separation of mutation bias and selection bias and the importance of recognizing the presence of multiple cellular environments in the study of codon usage. We also illustrate how a mechanistic model like ROC SEMPPR and the quantitative estimates it provides can be used for more sophisticated hypothesis testing in the future. In contrast to other approaches used to study codon usage like CAI (Sharp, 1987)

or tAI (DOS REIS et al., 2003, 2004), ROC SEMPPR accounts for the influence of mutation bias on codon usage. We highlight potential issues when estimating codon preferences, as estimates can be biased by the signature of a second, historical cellular environment. In addition, we show how quantitative estimates of mutation bias and selection relative to drift tAI can be obtained from codon data and used to infer the fitness cost of an introgression as well intro as its history and potential future.

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#### Materials and Methods 3.5

discussion.

#### 3.5.1Separating Endogenous and Exogenous Genes

A GC-rich region was identified by Payen et al. (2009) in the L. kluyveri genome extending from position 1 to 989,693 of chromosome C. This region was later identified as an introgression by FRIEDRICH et al. (2015). We obtained the L. kluyveri genome from SGD Project http://www.yeastgenome.org/download-data/ (on 09-27-2014) and the annotation for L. kluyveri NRRL Y-12651 (assembly ASM14922v1) from NCBI (on 12-09-2014). We assigned 457 genes located on chromosome C with a location within the  $\sim 1Mb$  window to the exogenous gene set. All other 4864 genes of the L. kluyveri genome were assigned to the exogenous genes. All genes could be uniquely assigned to one or the other gene set.

#### 3.5.2Model Fitting with ROC SEMPPR

ROC SEMPPR was fitted to each genome using AnaCoDa (0.1.1) (LANDERER et al., 2018) and R (3.4.1) (R CORE TEAM, 2015). ROC SEMPPR was run from multiple starting values for at least 250,000 iterations, only every 50th step was collected as a sample to reduce autocorrelation. After manual inspection to verify that the MCMC had converged, parameter posterior means were estimated from the last 500 samples.

#### 3.5.3 Comparing Codon Specific Parameter Estimates

Choice of reference codon does reorganize codon families coding for an amino acid relative to each other, therefore all parameter estimates are relative to the mean for each codon family.

$$\Delta M_{i,a}^c = \Delta M_{i,a} - \overline{\Delta M_a} \tag{3.1}$$

$$\Delta \eta_{i,a}^c = \Delta \eta_{i,a} - \overline{\Delta \eta_a} \tag{3.2}$$

Comparison of codon specific parameters ( $\Delta M$  and  $\Delta \eta$ ) was performed using the function lmodel2 in the R package lmodel2 (1.7.3) (Legendre, 2018) and R version 3.4.1 (R Core Team, 2015). Type II regression was performed with re-centered parameter estimates, accounting for noise in dependent and independent variable (Sokal and Rohlf, 1981).

#### 3.5.4 Synteny Comparison

We obtained complete genome sequences from NCBI (on: 02-05-2017). Genomes were aligned and checked for synteny using SyMAP (4.2) with default settings (SODERLUND et al., 2006, 2011). We assess synteny as percentage coverage of the exogenous gene region (Figure 3.8b).

## 3.5.5 Estimating Age of Introgression

We modeled the change in codon frequency over time using an exponential model for all two codon amino acids, and describing the change in codon  $c_1$  as

$$\frac{dc_1}{dt} = -\mu_{1,2}c_1 - \mu_{2,1}(1 - c_1) \tag{3.3}$$

where  $\mu_{i,j}$  is the rate at which codon i mutates to codon j and  $c_1$  is the frequency of the reference codon. Our estimates of  $\Delta M_{\rm endo}$  can be used to calculate the steady state of

equation 3.3.

$$\frac{\mu_{2,1}}{\mu_{1,2} + \mu_{2,1}} = \frac{1}{1 + \exp[\Delta M_{\text{endo}}]}$$
(3.4)

Solving for  $\mu_{1,2}$  gives us  $\mu_{1,2} = \Delta M_{\text{endo}} \exp[\mu_{2,1}]$  which allows us to rewrite and solve equation 3.3 as

$$c_1(t) = \frac{\exp[-t(1 + \Delta M_{\text{endo}})\mu_{2,1}] \exp[t(1 + \Delta M_{\text{endo}})\mu_{2,1}] + (1 + \Delta M_{\text{endo}})K}{1 + \Delta M_{\text{endo}}}$$
(3.5)

where K is

$$K = c_1(0) - \frac{1}{1 + \Delta M_{\text{endo}}} \tag{3.6}$$

Equation 3.5 was solved with a mutation rate  $m_{2,1}$  of  $3.8 \times 10^{-10}$  per nucleotide per generation (Lang and Murray, 2008). Initial codon frequencies  $c_1(0)$  for each codon family where taken from our mutation parameter estimates for E. gossypii  $\Delta M_{gos}$ . Current codon frequencies for each codon family where taken from our estimates of  $\Delta M$  from the exogenous genes. Mathematica (11.3) (Wolfram Research Inc., 2017) was used to calculate the time  $t_{intro}$  it takes for the initial codon frequencies  $c_1(0)$  for each codon family to equal the current exogenous codon frequencies. The same equation was used to determine the time  $t_{decay}$  at which the signal of the exogenous cellular environment has decayed to within 1% of the endogenous environment.

## **Estimating Genetic Load**

To estimate the fitness burden, we made three key assumptions. First, we assumed that the current exogenous amino acid sequence of a gene is representative of its ancestral state and the replaced endogenous gene it replaced. Second, we assume that the currently observed cellular environment of *E. gossypii* reflects the cellular environment that the exogenous genes experienced before transfer to *L. kluyveri*. Lastly, we assume that the difference in the efficacy of selection between the cellular environments due to differences in either effective

population size  $N_e$  or the selective cost of an ATP q of the source lineage and L. kluyverical can be expressed as a scaling constant and that protein synthesis rate  $\phi$  has not changed between the replaced endogenous and the introgressed exogenous genes. Using estimates for  $N_e = 1.36 \times 10^7$  (WAGNER, 2005) for Saccharomyces paradoxus we scale our estimates of  $\Delta \eta$  and define  $\Delta \eta' = \frac{\Delta \eta}{N_e}$ .

We scale the difference in the efficacy of selection on codon usage between the donor lineage and L. kluyveri using a linear scaling factor  $\kappa$ . As  $\Delta \eta$  is defined as  $\Delta \eta = 2N_e q(\eta_i - \eta_j)$ , we can not distinguish if  $\kappa$  is a scaling on protein synthesis rate  $\phi$ , effective population size  $N_e$ , or the selective cost of an ATP q (GILCHRIST, 2007; GILCHRIST et~al., 2015). We calculated the fitness burden each gene represents assuming additive fitness effects as

$$s_g = \sum_{i=1}^{n_g} -\kappa \phi_g \Delta \eta_i' \tag{3.7}$$

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where  $s_g$  is the overall strength of selection for translational efficiency on gene g in the exogenous gene set,  $\kappa$  is a constant, scaling the efficacy of selection between the endogenous and exogenous cellular environments,  $n_g$  is length of the protein,  $\phi_g$  is the estimated protein synthesis rate of the gene in the endogenous environment, and  $\Delta \eta_i'$ , is the  $\Delta \eta$ ' for the codon at position i. As stated previously, our  $\Delta \eta$  are relative to the mean of the codon family. We find that the fitness burden of the introgressed genes is minimized at  $\kappa \sim 5$  (Figure 3.9b). Thus, we set  $\kappa = 1$  if we calculate the  $s_g$  for the endogenous and the current exogenous genes, and  $\kappa = 5$  for  $s_g$  for the fitness burden at the time of introgression. Since we are unable to observe codon counts for the replaced endogenous genes and for the exogenous genes at the time of introgression, we calculate expected codon counts

$$E[n_{g,i}] = \frac{\exp[-\Delta M_i - \Delta \eta_i \phi_g]}{\sum_{j=1}^{C} \exp[-\Delta M_j - \Delta \eta_j \phi_g]} \times m_{a_i}$$
(3.8)

 $m_{a_i}$  is the number of occurrences of amino acid a that codon i codes for.

We report the genetic load of the introgression as  $E[s_g] = s_{\text{intro},g} - s_{\text{endo},g}$  where  $s_{\text{intro},g}$  is the fitness burden of an introgressed gene g either at the time of the introgression or presently.

## 3.6 Acknowledgments

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## 3.7 Appendix: Supplementary Material

Table 3.2: Synonymous codon preference in the various data sets based on our estimates of  $\Delta M$ 

| Amino Acid             | E. gossypii | Endogenous | Exogenous | L. kluyveri |
|------------------------|-------------|------------|-----------|-------------|
| Ala A                  | GCG         | GCA        | GCG       | GCG         |
| Cys C                  | TGC         | TGT        | TGC       | TGC         |
| Asp D                  | GAC         | GAT        | GAC       | GAC         |
| Glu E                  | GAG         | GAA        | GAG       | GAG         |
| Phe F                  | TTC         | TTT        | TTT       | TTT         |
| Gly G                  | GGC         | GGT        | GGC       | GGC         |
| His H                  | CAC         | CAT        | CAC       | CAC         |
| Ile I                  | ATC         | ATT        | ATC       | ATA         |
| Lys K                  | AAG         | AAA        | AAG       | AAA         |
| Leu L                  | CTG         | TTG        | CTG       | CTG         |
| Asn N                  | AAC         | AAT        | AAC       | AAT         |
| Pro P                  | CCG         | CCA        | CCG       | CCG         |
| Gln Q                  | CAG         | CAA        | CAG       | CAG         |
| $\operatorname{Arg} R$ | CGC         | AGA        | AGG       | CGG         |
| $Ser_4 S$              | TCG         | TCT        | TCG       | TCG         |
| Thr T                  | ACG         | ACA        | ACG       | ACG         |
| Val V                  | GTG         | GTT        | GTG       | GTG         |
| Tyr Y                  | TAC         | TAT        | TAC       | TAC         |
| $Ser_2 Z$              | AGC         | AGT        | AGC       | AGC         |

Table 3.3: Synonymous codon preference in the various data sets based on our estimates of  $\Delta\eta$ 

| Amino Acid | E. gossypii | Endogenous | Exogenous | L. kluyveri |
|------------|-------------|------------|-----------|-------------|
| Ala A      | GCT         | GCT        | GCT       | GCT         |
| Cys C      | TGT         | TGT        | TGT       | TGT         |
| Asp D      | GAT         | GAC        | GAT       | GAT         |
| Glu E      | GAA         | GAA        | GAA       | GAA         |
| Phe F      | TTT         | TTC        | TTC       | TTC         |
| Gly G      | GGA         | GGT        | GGT       | GGT         |
| His H      | CAT         | CAC        | CAT       | CAT         |
| Ile I      | ATA         | ATC        | ATT       | ATT         |
| Lys K      | AAA         | AAG        | AAA       | AAG         |
| Leu L      | TTA         | TTG        | TTG       | TTG         |
| Asn N      | AAT         | AAC        | AAT       | AAC         |
| Pro P      | CCA         | CCA        | CCT       | CCA         |
| Gln Q      | CAA         | CAA        | CAA       | CAA         |
| Arg R      | AGA         | AGA        | AGA       | AGA         |
| $Ser_4 S$  | TCA         | TCC        | TCT       | TCT         |
| Thr T      | ACT         | ACC        | ACT       | ACT         |
| Val V      | GTT         | GTC        | GTT       | GTT         |
| Tyr Y      | TAT         | TAC        | TAT       | TAC         |
| $Ser_2 Z$  | AGT         | AGT        | AGT       | AGT         |

| Taxon                      | Abbreviation | NCBI taxonomic ID | Codon Table | % GC  | GC Source    |
|----------------------------|--------------|-------------------|-------------|-------|--------------|
| Candida albicans           | Calb         | 5476              | 12          | 34    | NCBI Geno:   |
| Saccharomyces bayanus      | Sbay         | 4931              | 1           | 40    | NCBI Geno    |
| Trichophyton benhamiae     | Tben         | 63400             | 1           | 49    | NCBI Geno    |
| Tetrapisispora blattae     | Tbla         | 1071379           | 1           | 32    | NCBI Geno    |
| Saccharomyces castellii    | Scas         | 27288             | 1           | 37    | NCBI Geno:   |
| Saccharomyces cerevisiae   | Scer         | 4932              | 1           | 38    | NCBI Geno:   |
| Eremothecium cymbalariae   | Ecym         | 45285             | 1           | 40    | NCBI Geno:   |
| Torulaspora delbrueckii    | Tdel         | 4950              | 1           | 42    | NCBI Geno:   |
| Candida dubliniensis       | Cdub         | 42374             | 12          | 33    | NCBI Geno    |
| Lodderomyces elongisporus  | Lelo         | 36914             | 1           | 37    | NCBI Geno    |
| Saccharomyces eubayanus    | Seub         | 1080349           | 1           | 40    | NCBI Geno:   |
| Debaryomyces fabryi        | Dfab         | 58627             | 1           | 36    | NCBI Geno:   |
| Candida glabrata           | Cgla         | 5478              | 1           | 39    | NCBI Geno:   |
| Eremothecium gossypii      | Egos         | 33169             | 1           | 52    | NCBI Geno:   |
| Meyerozyma guilliermondii  | Mgui         | 4929              | 12          | 44    | NCBI Geno:   |
| Debaryomyces hansenii      | Dhan         | 4959              | 12          | 36    | NCBI Geno:   |
| Lachancea kluyveri         | Lku          | 4934              | 1           | 40/53 | Payen et al. |
| Saccharomyces kudriavzevii | Skud         | 114524            | 1           | 41    | NCBI Geno:   |
| Kluyveromyces lactis       | Klac         | 28985             | 1           | 39    | NCBI Geno:   |
| Lachancea lanzarotensis    | Llan         | 1245769           | 1           | 44    | NCBI Geno:   |
| Yarrowia lipolytica        | Ylip         | 4952              | 1           | 49    | NCBI Geno:   |
| Clavispora lusitaniae      | Clus         | 36911             | 12          | 45    | NCBI Geno:   |
| Kluyveromyces marxianus    | Kmar         | 4911              | 1           | 40    | NCBI Geno:   |
| Saccharomyces mikatae      | Smik         | 114525            | 1           | 38    | NCBI Geno:   |
| Sphaerulina musiva         | Smus         | 85929             | 1           | 51    | NCBI Geno:   |
| Kazachstania naganishii    | Knag         | 588726            | 1           | 46    | NCBI Geno:   |
| Saccharomyces paradoxus    | Spar         | 27291             | 1           | 38    | NCBI Geno:   |
| Candida parapsilosis       | Cpar         | 5480              | 12          | 38    | NCBI Geno:   |
| Spathaspora passalidarum   | Spas         | 340170            | 12          | 38    | NCBI Geno:   |
| Tetrapisispora phaffii     | Tpha         | 113608            | 1           | 34    | NCBI Geno:   |
| Vanderwaltozyma polyspora  | Vpol         | 36033             | 1           | 33    | NCBI Geno:   |
| Lachancea quebecensis      | Lque         | 1654605           | 1           | 47    | Freel et al. |
| Zygosaccharomyces rouxii   | Zrou         | 4956              | 1           | 40    | NCBI Geno:   |
| Scheffersomyces stipitis   | Ssti         | 4924              | 12          | 41    | NCBI Geno:   |
| Lachancea thermotolerans   | Lthe         | 381046            | 1           | 47    | NCBI Geno:   |
| Candida tropicalis         | Ctro         | 5482              | 12          | 33    | NCBI Geno:   |
| Lachancea waltii           | Lwal         | 4914              | 1           | 44    | NCBI Geno:   |
| Cladophialophora yegresii  | Cyeg         | 470704            | 1           | 54    | NCBI Geno:   |

Table 3.4:

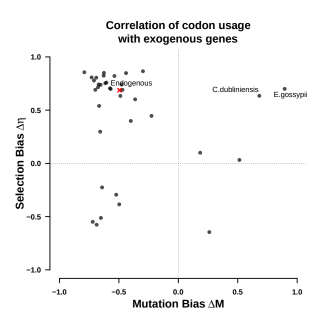


Figure 3.5: Correlation coefficient of  $\Delta M$  and  $\Delta \eta$  of the endogenous genes with 38 examined yeast lineages. Dots indicate the correlation of  $\Delta M$  and  $\Delta \eta$  of the lineages with the endogenous and exogenous parameter estimates. All regressions were performed using a type II regression line (SOKAL and ROHLF, 1981).

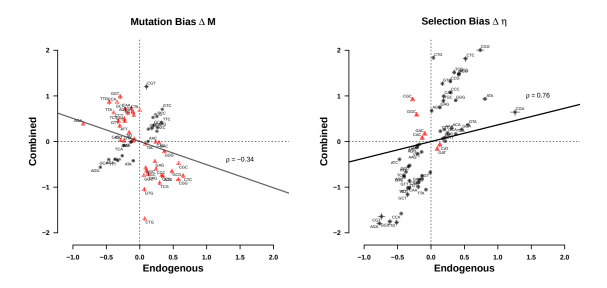


Figure 3.6: Comparison of (a) mutation bias  $\Delta M$  and (b) selection bias  $\Delta \eta$  parameters for endogenous genes and combined gene sets. Estimates are relative to the mean for each codon family. Black dots indicate  $\Delta M$  or  $\Delta \eta$  parameters with the same sign for the endogenous and exogenous genes, red dots indicate parameters with different signs. Black line shows the type II regression line (Sokal and Rohlf, 1981). Dashed lines mark quadrants.

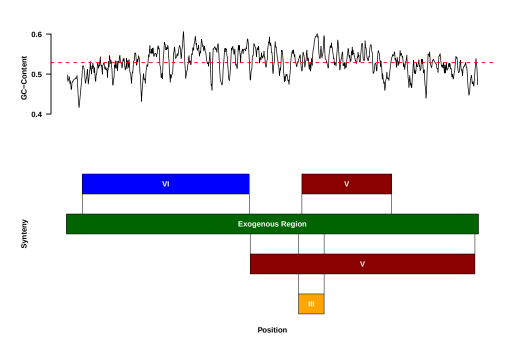


Figure 3.7: Synteny relationship of E. gossypii and the exogenous genes. Indicated is the GC content along the introgression.

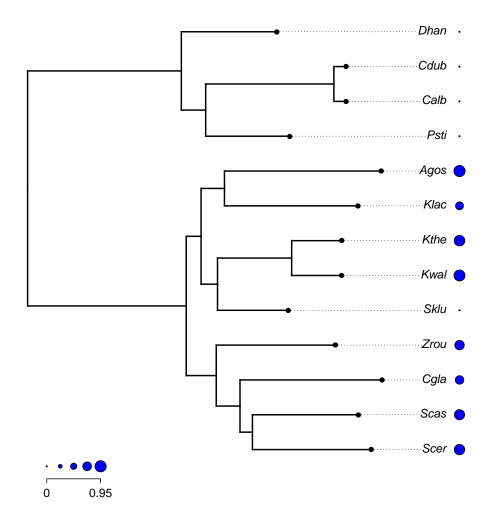


Figure 3.8: Amount of synteny for each species in units of standard deviations for selected species.

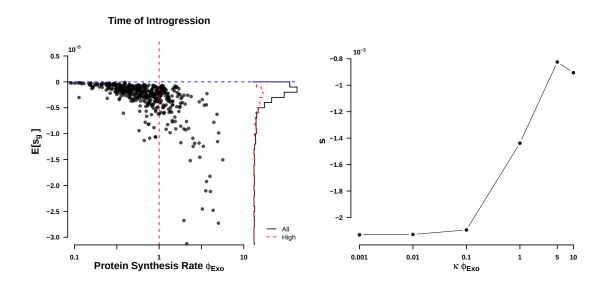


Figure 3.9: Genetic load (left) without scaling of  $\phi$  per gene, and change of total genetic load with scaling  $\kappa$  between E. gossypii and L. kluyveri (right)

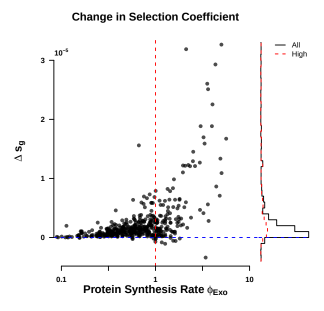


Figure 3.10: Total amount of adaptation estimated to have occured between time of introgression and currently observed per gene.

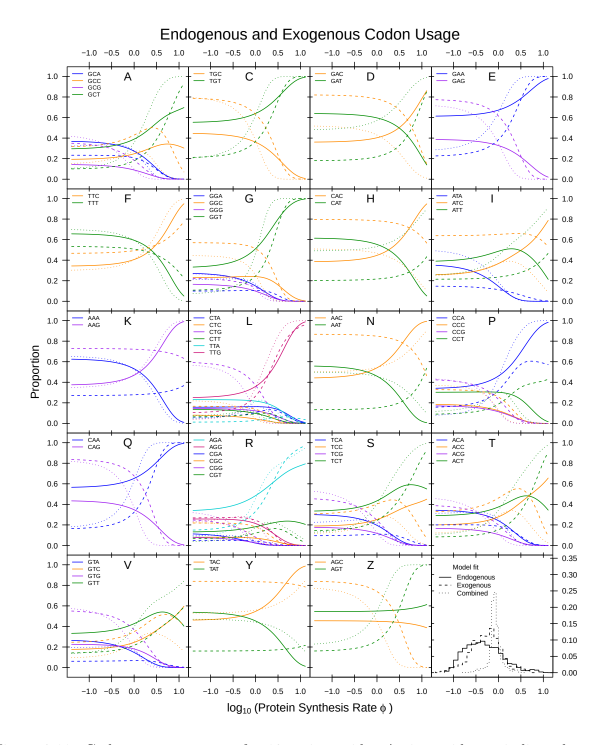


Figure 3.11: Codon usage patterns for 19 amino acids. Amino acids are indicated as one letter code. The amino acids Serine was split into two groups (S and Z) as Serine coded for by two groups of codons that are separated by more than one mutation. Solid line indicates the endogenous codon usage, dashed line indicates the exogenous codon usage, dotted line indicates the combined codon usage.

## Chapter 4

Phylogenetic model of stabilizing selection is more informative about site specific selection than extrapolation from laboratory estimates

This chapter is an early version of a paper to be submitted to Genome Biology and Evolution and co-authored with Michael A. Gilchrist and Brian C. O'Meara.

C. Landerer, B.C. O'Meara, M.A. Gilchrist, Phylogenetic model of stabilizing selection is more informative about site specific selection than extrapolation from laboratory estimates

#### 4.1 Abstract

Here we examine the ability of experimentally inferred, site specific selection for amino acids to improve phylogenetic inferences of sequence evolution. Previous work has shown that laboratory estimates of selection can improve model fit, but did not assess their adequacy. We assess the ability of experimentally inferred site specific selection for antibiotic resistance from deep mutation scanning to inform phylogenetic models. In this study, we use the  $\beta$ -lactamase TEM for which empirical estimates of site specific selection on amino acids are readily available. TEM is an enzyme that catalyzes antibiotics with a  $\beta$ -lactam ring and is found in gram-negative bacteria like *Escherichia coli*. We compare the experimentally inferred site specific selection to our results obtained using SelAC, a new phylogenetic model of stabilizing selection. Using simulations we assess model adequacy, and find that experimentally inferred selection does not adequately reflect evolution in the wild. In contrast, SelAC fits to the data better over models informed by experimentally inferred selection and provides higher model adequacy. We demonstrate the capability of SelAC by estimating the site specific genetic load of the observed TEM variants.

#### 4.2 Introduction

Numerous attempts to incorporate selection into phylogenetic models have been made. Early models only focused on the influence of selection on the substitution rate and fixation probability between a resident and a mutant introduced into a population (GOLDMAN and YANG, 1994; Muse and Gaut, 1994; Thorne et al., 1996). These models however, lack site specific equilibrium codon or amino acid frequencies. The importance of site specific equilibrium frequencies has long been noted (Felsenstein, 1981; Gojobori, 1983). Individual amino acid sites along the protein show differences in evolutionary rates, and wide range of preferences for specific amino acids (Halpern and Bruno, 1998; Ashenberg et al., 2013; Echave et al., 2016). The usage of site specific selection acknowledges the heterogeneity in selection and amino acid preferences along the protein sequence (Hilton et al., 2017).

HALPERN and BRUNO (1998) first introduced a model to incorporate site specific equilibrium frequencies of amino acids. However, they had to concede that their model was too parameter rich and therefore intractable for biological data sets without additional simplifying assumptions. More recent models incorporating site specific equilibrium frequencies still require a large number of parameters to be estimated from the sequence data (Lartillot and Philippe, 2004; Le et al., 2008; Wang et al., 2008; Holder et al., 2008; Wu et al., 2013; Tamuri et al., 2014). Other approaches treat site specific selection as a random effect (Rodrigue et al., 2010; Rodrigue, 2013; Rodrigue and Lartillot, 2014). A full parameterization of site specific equilibrium frequencies for amino acids requires  $19 \times N$  parameters where N is the length of the sequence in amino acids. It is therefore an attractive option to utilize laboratory experiments to empirically estimate site specific strength of selection on amino acids and infer their equilibrium frequencies (Bloom, 2014; Thyagarajan and Bloom, 2014; Bloom, 2017).

Empirical estimates of site specific selection can greatly reduce the number of parameters estimated from phylogenetic data, making it applicable for smaller data sets and allowing

for the fitting of more complex models. Deep mutation scanning (DMS) has recently been used to generate comprehensive site specific estimates of selection (Fowler et al., 2014). The ability to estimate site specific selection allows to estimate site specific amino acid preferences and the fitness consequences a mutation introduces at a particular site (Bloom, 2014; Firnberg et al., 2014; Stiffler et al., 2016). There are, however, also shortcomings. The quality of empirical estimates from DMS, however, depends on many factors including the initial library of mutants and the applied selection (Firnberg and Ostermeier, 2012). Mutation libraries have to be extensive and, therefore, produce a heterogeneous population of competing organisms not usually found in nature. In addition, estimates of selection can only be obtained for fast growing organisms that can be manipulated under laboratory conditions. As many organism can not be cultivated under laboratory conditions or have long generation times, this is a severe limitation to experimentally informed models.

Even in the cases where empirical estimates of site specific selection can be obtained, their utility for phylogenetic reconstruction is questionable. In this study, we assess the ability of experimentally inferred site specific selection to inform phylogenetic models and offer an alternative approach to determine site specific selection. We use site specific estimates of selection for the class A  $\beta$ -lactamase TEM from Stiffler et al. (2016). TEM is an enzyme found in gram-negative bacteria like Escherichia coli that catalyzes antibiotics with a  $\beta$ -lactam ring and provides antibiotic resistance (NEU, 1969). The selection pressure imposed during the DMS experiment was limited to ampicillin and focused solely on TEM-1 (Stiffler et al., 2016). However, TEM variants can also confer resistance to a wide range of other antibiotics (Sougakoff et al., 1988, 1989; Goussard et al., 1991; Mabilat et al., 1992; Chanal et al., 1992; Brun et al., 1994).

In order to do so, we fitted 227 nucleotide and codon models using IQTree and compared their model fits to site specific models of stabilizing selection with (phydms, SelAC+DMS) and without (SelAC) experimentally estimated site specific selection coefficients (NGUYEN et al., 2015; HILTON et al., 2017; BEAULIEU et al., in review). We find that experimentally

inferred selection, while improving model fit, does not adequately reflect observed wild type sequences. In contrast, SelAC (BEAULIEU et al., in review) a mechanistic phylogenetic model of stabilizing selection rooted in first principles with site specific equilibrium frequencies improves model fit, and better predicts sequences found in the wild. In addition, it was previously proposed to extrapolate to the fitness landscape of related proteins using experimentally inferred site specific selection (BLOOM, 2014, 2017). We utilize the TEM homologue SHV, another class A  $\beta$ -lactamase, to demonstrate the problematic with this approach and further highlight the generality of SelAC.

#### 4.3 Results

# 4.3.1 Site Specific Stabilizing Selection on Amino Acids Improves Model Fit

We compared phydms (Hilton et~al., 2017) and SelAC (Beaulieu et~al., in review), models of site specific stabilizing selection on amino acids, to 227 other codon and nucleotide models. We fitted all models to 49 observed sequences of the  $\beta$ -lactamase TEM (Bloom, 2014). The phydms and SelAC models with site specific selection improved model fits by 366 and 934 AICc units, respectively, over the best performing codon or nucleotide models which lacks site specific selection (Table 4.1). In addition, SelAC outperformed the experimentally informed model phydms by 562 to 568 AICc units, depending whether site specific selection was inferred by SelAC or experimentally informed.

SelAC utilizes a hierarchical model and estimates 263 site specific parameters,  $\sim 5\%$  of the  $19 \times N = 4997$  parameters necessary to fully describe site specific selection. In contrast, phydms does not infer any site specific parameters from the phylogenetic data, but utilizes site specific selection estimated from deep mutation scanning experiments. In order to assess the quality of the SelAC fit with experimentally determined site specific selection, we fixed the optimal amino acid at each site to the experimentally determined one in SelAC

Table 4.1: Model selection, shown are the three models of stabilizing site specific amino acid selection (SelAC, SelAC+DMS, phydms) and the best performing codon and nucleotide model (Goldman and Yang, 1994; Zharkikh, 1994). Reported are the log-likelihood  $log(\mathcal{L})$ , the number of parameters estimated n, AIC,  $\Delta$ AIC, AICc, and  $\Delta$ AICc values. See Table 4.3 for results from all models we tested.

| Model         | $\log(\mathcal{L})$ | n   | AIC  | $\Delta { m AIC}$ | AICc | $\Delta { m AICc}$ |
|---------------|---------------------|-----|------|-------------------|------|--------------------|
| SelAC+DMS     | -1768               | 111 | 3758 | 14                | 3760 | 0                  |
| SelAC         | -1498               | 374 | 3744 | 0                 | 3766 | 6                  |
| phydms        | -2061               | 102 | 4326 | 582               | 4328 | 568                |
| SYM+R2        | -2230               | 102 | 4663 | 919               | 4694 | 934                |
| GY94 +F1X4+R2 | -2243               | 102 | 4690 | 946               | 4821 | 1061               |

and refitted the model to the 49 TEM sequences (SelAC+DMS). Incorporating site specific selection estimated from deep mutation scanning experiments into SelAC (SelAC+DMS) yields a similar, but slightly better AICc value to SelAC without that information.

This improvement in AICc of SelAC+DMS over SelAC is solely due to a decrease in the number of parameters estimated In contrast to AICc, the log-likelihood  $log(\mathcal{L})$  of SelAC+DMS is 270  $log(\mathcal{L})$  units worse than the SelAC one (Table 4.1). However, it is statistically speaking unclear if discrete parameters bias ones estimate the Kullback-Leibler divergence in the same way. This is important since 263 of the 374 parameters estimated by SelAC are the discrete optimal amino acid state at each site. Thus, it is possible that we are over penalizing. Therefore, the number of parameter for SelAC we use is conservative. For example, there are only 27 unique site patterns in the TEM alignment, which would yield a total of 138 parameters. This however would likely be an under estimate of the number of parameters estimated. The true number of parameters remains unclear at this point due to the inherent non-independence of the underlying data and the discrete nature of the optimized parameters.

We observe differences in the topology between model fits. The *SelAC* model is currently too slow to estimate the topology, therefore the topology was estimated using the codon model of Kosiol *et al.* (2007). At this point, it is therefore unclear if the difference in topology can be attributes to the experimentally inferred selection. We find that the best

codon model (GY94) (GOLDMAN and YANG, 1994) is outperformed by several nucleotide model e.g. SYM+R2 (ZHARKIKH, 1994). This could be an indication that negative frequency dependent selection like it is modeled in GY94 is not appropriate for TEM (GOLDMAN and YANG, 1994; BEAULIEU et al., in review). Figure 4.1 shows that the estimated phylogenetic trees shift from long terminal branches (SelAC) to longer internal branches (phydms). While the SelAC model fit shows 84% of all evolution happening at the tips, this reduces to 79% in the SelAC+DMS model fit, and 77% in the phydms and GY94 model fits. All models produce polytomies but their location differs between models. Surprisingly, the largest polytomies appear in the experimentally informed phylogeny of phydms. The position of the sequences with the longest branches also differ between SelAC and phydms.

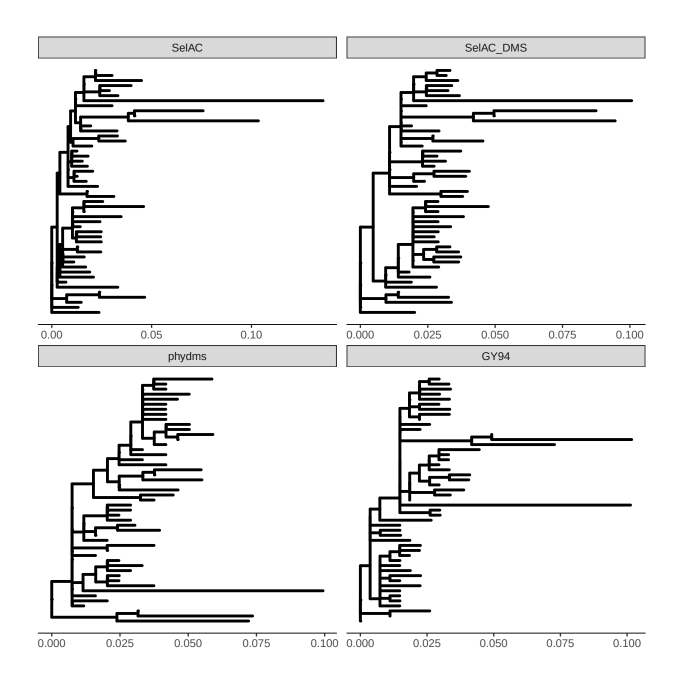


Figure 4.1: Phylogenies resulting from SelAC, SelAC+DMS, phydms, and GY94. As SelAC is currently to slow for the inference of topologies, the topology for the SelAC phylogenies was inferred using the codon model of Kosiol  $et\ al.\ (2007)$ .

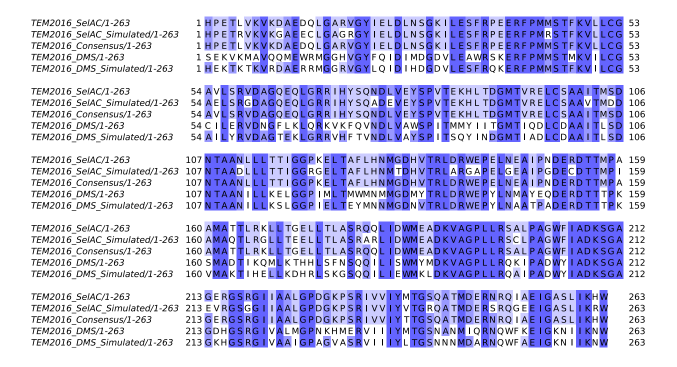


Figure 4.2: Alignment of TEM optimal and simulated sequences. Indicated is the percentage identity at each site.

#### 4.3.2 Laboratory Inferences Inconsistent with Observed Sequences.

The improved model fits of *phydms* relative to more common nucleotide and codon models are, however, deceiving. The site specific selection inferred by DMS is inconsistent with the observed TEM sequences. We find that the sequence of selectively favored amino acids has only 52% sequence similarity with the observed consensus sequence (Figure 4.2). In contrast, the sequence of selectively favored amino acids estimated by *SelAC* shows 99% sequence similarity with the observed consensus sequence. In addition, assuming the site specific selection estimated by DMS, the observed TEM sequences represent an average sequence specific genetic load of 17.88 and an average site specific load of 0.065.

In order to determine if we would expect the observed genetic load under the experimental selection estimates we reconstructed the ancestral TEM sequence and used it as initial conditions for our simulation studies. We simulate under a wide range of effective population

sizes  $N_e$ , and find that the experimentally inferred site specific selection is very strong. The estimated ancestral state is identical to the observed consensus sequence. Simulations of codon sequences under the experimentally inferred site specific selection for amino acids reveal that we would not expect to see the observed TEM sequences. With an effective population size  $N_e$  of  $10^7$ , we find that the simulated sequences show 62% sequence similarity to the observed consensus sequence (Figure 4.3a). Thus, the simulated sequences show a 10% higher similarity to the observed consensus sequence than the sequence of selectively favored amino acids estimated using deep mutation scanning.

In our simulations, only when  $N_e$  is reduced to one individual does drift overwhelm selection (Figure 4.3b). The genetic load of the simulated sequences decrease slowly with increasing  $N_e$ . After simulating until the sequences reached 0.1 expected mutation per site with an effective population size  $N_e = 10^7$  the simulated sequences showed an average sequence specific load of 6.68 and an average site specific genetic load of 0.025 This is less than half of the average sequence and site specific genetic load of the observed sequences. Thus it appears unlikely that the observed sequences have evolved under the DMS inferred site specific selection values.

## 4.3.3 Stabilizing Selection for Optimal Physicochemical Properties Improves Model Adequacy

Model adequacy of SelAC assessed based on sequence similarity and genetic load and shows that SelAC better explains the observed TEM sequences than the experimentally determined site specific selection on amino acids. The observed consensus sequence has 99% sequence similarity with the sequence of selectively favored amino acids estimated by SelAC, this is in contrast to the average sequence similarity of 98% among all 49 observed sequences. In addition, assuming the site specific selection estimated by SelAC, the observed TEM sequences represent an average sequence specific genetic load of  $6.4 \times 10^{-5}$  and an average site specific load of  $2.4 \times 10^{-7}$ .

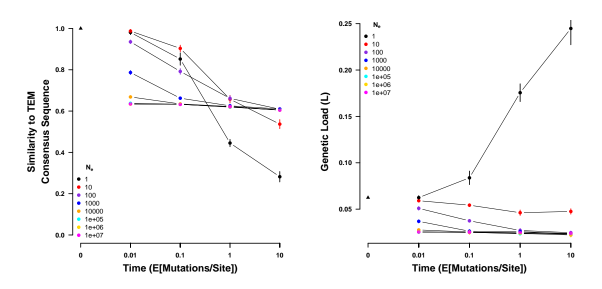


Figure 4.3: Sequences simulated from the ancestral state under the site specific selection on amino acids estimated using deep mutation scanning. (left) Sequence similarity to the observed consensus sequence at various times for a range of values of  $N_e$ . (right) Genetic load of the simulated sequences at various times for a range of values of  $N_e$ . Time is given in number of expected mutations per site, which equals the substitution rate of a neutral mutation. Points indicate sample means and vertical bars indicate standard deviations. Initial sequence is the inferred ancestral state of the TEM variants and indicated by a black triangle.

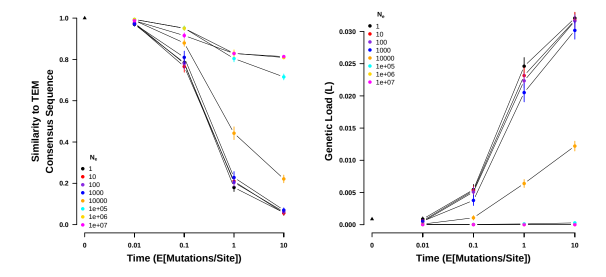


Figure 4.4: Sequences simulated from the ancestral state under the site specific selection on amino acids estimated using SelAC. (left) Sequence similarity to the observed consensus sequence at various times for a range of values of  $N_e$ . (right) Genetic load of the simulated sequences at various times for a range of values of  $N_e$ . Time is given in number of expected mutations per site, which equals the substitution rate of a neutral mutation. Points indicate sample means and vertical bars indicate standard deviations. Initial sequence is the inferred ancestral state of the TEM variants and indicated by a black triangle.

Using the SelAC inferred site specific selection and the reconstructed ancestral TEM sequence as initial conditions we simulated codon sequences forward in time for various time periods and  $N_e$  values. As expected, for small  $N_e$ , simulated sequences drift away from the observed consensus sequence (Figure 4.4a). Because of the high similarity between the optimal amino acid sequence estimated by SelAC and the observed consensus sequence, the genetic load increases drastically as a result. Increasing  $N_e$  to  $10^4$  or above the simulated sequences sequence similarity declines to 83%, indicating that SelAC underestimates selection. After simulating until the sequences reached 0.1 expected mutation per site with an effective population size  $N_e = 10^7$  the simulated sequences showed an average sequence specific load of  $1.3 \times 10^{-5}$  and an average site specific genetic load of  $4.8 \times 10^{-8}$  (Figure 4.4b). Thus, the simulated sequences show a lower genetic load despite the greater divergence from the observed consensus sequence. This be an indication that the selection differs between linages.

To further demonstrate the consistency of SelAC, we simulated codon sequences over the same time periods using 10 sequences where codons where sampled uniform. We find that the sequence similarity increases with effective population size  $N_e$ . The random sequences start of with a similarity of  $\sim 6\%$  and increase  $N_e$  to  $\sim 28\%$  (Figure 4.8a). The same initial sequences simulated under the site specific selection inferred by the deep mutation scanning experiment increase only to  $\sim 18\%$  in sequence similarity over the same period of time.

#### 4.3.4 Estimating Site Specific Selection on Amino Acids

SelAC allows for the estimation of site specific selection on amino acids and the genetic load of an observed amino acid relative to the inferred optimal amino acid. Figures 4.5 and 4.6 illustrate how the genetic load varies along the TEM sequence. The region between residue 80 and 120, where three consecutive helices are located, consists only of selectively favored amino acids and does not show any genetic load. The highest genetic load is found in the unstructured regions and the lowest genetic load is found in  $\beta$ -sheets. However, this difference is not statistically significant (p = 0.17). The largest increase in genetic load is located at the beginning of the last helix. This region strongly contributes to the estimate of similar genetic loads for helices and unstructured regions in the observed TEM sequences (Table 4.2). However, exclusion of this site as outlier does not yield significance (p = 0.09).

SelAC assume that the efficacy of selection G is  $\Gamma$ -distributed with a mean of 1. However, it is possible to estimate site specific values using the parameters estimated by SelAC. We constraint G to a maximum value of 300 in all cases. While this biases our estimate of G, the bias is consistent across all estimates and does not prohibit the comparison of G terms.

The highest efficacy of selection G is estimated in the  $\beta$ -sheet regions which is consistent with the lowest genetic load in these regions. Residues forming the substrate binding site appear to be under the strongest selection, with no accumulated genetic load. However, this is not the case for the two active sites. We find in one sequence (Acinetobacter baumannii, TEM-193) a Lysine, a proton donor, at the proton acceptor site 143 driving the reduced

Table 4.2: Efficacy of selection (G) and genetic load for TEM and SHV, and separated by secondary structure. G was estimated as a truncated variable with an upper bound of 300.

|         |                      |            | G     |      | Genetic               | Load $L_i$            |
|---------|----------------------|------------|-------|------|-----------------------|-----------------------|
| Protein | Secondary Structure  | # Residues | Mean  | SE   | Mean                  | SE                    |
| TEM     |                      | 263        | 219.3 | 7.5  | $15.9 \times 10^{-8}$ | $6.5 \times 10^{-8}$  |
|         | Helix                | 113        | 206.1 | 12.4 | $17.5 \times 10^{-8}$ | $13.1 \times 10^{-8}$ |
|         | $\beta$ -Sheet       | 48         | 238.6 | 15.8 | $6.8 \times 10^{-8}$  | $2.9 \times 10^{-8}$  |
|         | Unstructured         | 102        | 224.8 | 11.4 | $18.6 \times 10^{-8}$ | $8.1 \times 10^{-8}$  |
|         | Active/Binding Sites | 5          | 202.6 | 62.2 | $0.01\times10^{-8}$   | $0.01\times10^{-8}$   |
| SHV     |                      | 263        | 244.9 | 6.8  | $4.0 \times 10^{-8}$  | $1.9 \times 10^{-8}$  |
|         | Helix                | 102        | 234.6 | 11.5 | $7.3 \times 10^{-8}$  | $4.8 \times 10^{-8}$  |
|         | $\beta$ -Sheet       | 66         | 253.1 | 12.8 | $2.1 \times 10^{-8}$  | $1.1 \times 10^{-8}$  |
|         | Unstructured         | 95         | 224.7 | 11.4 | $1.5\times10^{-8}$    | $0.6 \times 10^{-8}$  |
|         | Active/Binding Sites | 5          | 239.9 | 60.0 | $1.5 \times 10^{-8}$  | $1.5 \times 10^{-8}$  |

efficacy of selection G. This is in concordance with the experimental DMS estimates, where proton acceptors are selectively favored. Again, any differences between secondary structure elements are not statistically significant.

It was previously proposed that experimentally inferred site specific selection for amino acids can be used to extrapolate the fitness landscape of related proteins (BLOOM, 2014, 2017). We therefore compared the genetic load, the SelAC selection parameters of our SelAC TEM model fit to a SelAC model fit of SHV, and site specific efficacy of selection G. The genetic load observed in SHV sequences appears to be lower than in TEM with the exception of residues found in  $\beta$ -sheets and the active site (Table 4.2). This is consistent with the elevated efficacy of selection G in SHV. However, only differences in genetic load in the unstructured regions are significantly different between the TEM and SHV sequences, but only at the  $\alpha = 0.05$  significant level (p = 0.04). While the average genetic load across secondary structures is not significantly different, the sites causing increases genetic load differ between SHV and TEM (Figure 4.7). In contrast to TEM, we find the highest genetic load among SHV secondary structure features in the helices (Table 4.2). The highest genetic load in SHV is observed at the end of the first helix. We do find a peak of similar magnitude

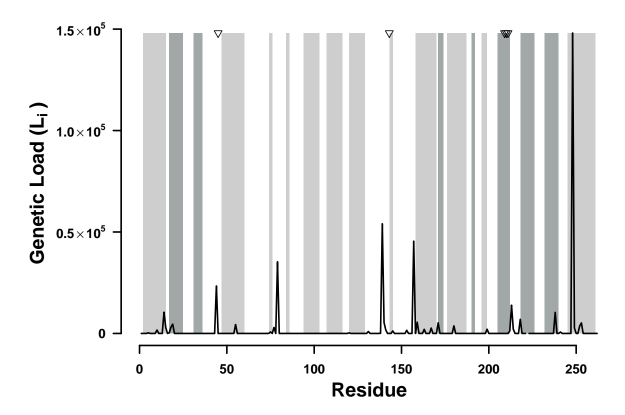


Figure 4.5: Distribution of average site specific genetic load in TEM over all observed TEM variants. Average site specific genetic load is indicated by the black line. Light gray bars indicate where helices are found, and dark gray bars indicate  $\beta$ -sheets. The three residues forming the binding site and the two residues forming the active are indicated by triangles at the top of the plot.

in the TEM sequence at the end of the first helix, but this peak is overshadowed by the increased genetic load at the beginning of the last helix.

We find that the site specific efficacy of selection G differs greatly between SHV and TEM ( $\rho = 0.12$ ), despite a similar estimate of  $\alpha_G$  describing the distribution of G values (Figure 4.10a). Most SelAC selection parameters are very similar between the TEM and the SHV model fit. An exception is the weight for the physicochemical composition property  $\alpha_c$  (Figure 4.10b). Furthermore, we find that the sequences of selectively favored amino acids estimated by SelAC for TEM and SHV only show 68% sequence similarity. These results indicate that the extrapolation from one proteins fitness landscape to another is problematic.

#### 4.4 Discussion

Here we revisited how well site specific estimates of selection from deep mutation scanning experiments inform sequence evolution and compared it to SelAC, a novel phylogenetic model of stabilizing selection. Previous work has shown that laboratory estimates of selection can improve model fit over classical approaches like GY94 (BLOOM, 2014, 2017). While our study confirms this notion, we identify important shortcomings of these laboratory estimates for phylogenetic studies. In contrast, SelAC is a phylogenetic model of stabilizing selection based on physicochemical properties and does not require costly laboratory estimates of selection and is, nevertheless, favored by model selection (Table 4.1). More specifically, it estimates site specific selection on amino acids from the sequence data based on distances between amino acids in physicochemical space (Grantham, 1974; Beaulieu et al., in review). This allows SelAC to be applied to any set of protein coding sequences, eliminating the need to extrapolate from one homologous gene family to the next (e.g. from TEM to SHV). In addition this generality allows for the comparison of model parameters for these proteins.

While previous work showed the advantages of experimentally informed phylogenetics, they did not assess how adequate the estimated selection reflects observed wild-type sequences. The low sequence similarity between the observed consensus sequence and the sequence of selectively favored amino acids estimated by deep mutation scanning experiments is evidence for that. This begs the question how well the experimental selection coefficients represent selection on these sequences in nature. Deep mutation scanning experiments are performed using a comprehensive library of mutants and a strong artificial selection pressure (FIRNBERG and OSTERMEIER, 2012; JAIN and VARADARAJAN, 2014; FOWLER and FIELDS, 2014; FOWLER et al., 2014). This results in very large selection coefficients s and a heterogeneous population of competing individuals unlikely to occur in nature.

The selection pressure imposed during the DMS experiment was limited to ampicillin and focused solely on TEM-1 (STIFFLER et al., 2016). However, TEM variants can also confer resistance to a wide range of other antibiotics, including penicillins, cephalosporins, cefotaxime, ceftazidime, or aztreonam (SOUGAKOFF et al., 1988, 1989; GOUSSARD et al., 1991; MABILAT et al., 1992; CHANAL et al., 1992; BRUN et al., 1994). Thus, the inferred selection is biased towards ampicillin and is inconsistent with the observed TEM sequences (Figure 4.3). This may very well be appropriate to explore the selection on TEM in a hospital environment but is unlikely to be representative of the selection faced by E. coli and other gram-negative bacteria in nature.

If we assume that the DMS selection coefficients underly the evolution of the observed TEM sequences we can think of two possible explanations for the observed sequences. First, the sequences are unable to reach a fitness peak, potentially due to a weak selection pressure or not enough time. Alternatively, the TEM sequences found in nature are highly maladapted, yet with very similar sequences. Both explanations seem unlikely. For example,  $E.\ coli$  has a large effective population size  $N_e$ , estimates are on the order of  $10^8$  to  $10^9$  (OCHMAN and WILSON, 1987; HARTL et al., 1994). We, therefore, expect the observed sequence variants to be near mutation-selection-drift equilibrium. This expectation is supported by our simulations in which we observe a higher sequence similarity with the observed TEM consensus sequence and decreased genetic load even with much smaller  $N_e$ 

(Figure 4.3). Furthermore, previous work showed that the catalytic reaction performed by TEM of penicillin-class antibiotics is close the diffusion limit. As a result some researchers refer to TEM as a perfect enzyme (MATAGNE et al., 1998; STIFFLER et al., 2016). The very large effective population size, however, also raises a concern that the population mutation rate of E.  $coli\ \Theta = 4N_e\mu$  exceeds 0.1 and, thus, violated SelAC's weak mutation assumption (DE KONING and DE SANCTIS, 2018). If the weak mutation assumption is violated evolution is no longer mutation limited and the time between fixation events increases. However, phydms operates under the same weak mutation assumption and the experimentally inferred selection clearly violates the weak mutation assumption.

As experimental selection estimates are not readily available for most organisms and proteins, a possible approach is the extrapolation of empirical estimates to homologous gene families (Bloom, 2014, 2017). When extrapolating the selection estimates from the  $\beta$ lactamase family TEM to the SHV family, the sequence similarity between the observed consensus sequence and the sequence of selectively favored amino acids estimated from deep mutation scanning experiments drops only slightly from 52% to 49%. This may have contributed to the notion that extrapolation to homologous gene families is possible. In contrast, estimates of site specific efficacy of selection G revealed large differences in the site specific selection on amino acids between TEM and SHV. The mismatched physicochemical weights further indicate differences in selection constraints. While the polarity of amino acids is of similar importance in TEM and SHV, amino acid composition appears to play a much greater role in SHV than in TEM. In contrast to the experimental selection estimates, extrapolated from TEM to SHV, the SelAC selection estimates are consistent with the observed sequences, e.g. the selectively favored amino acids estimated by SelAC shows a high sequence similarity with the observed TEM and SHV consensus sequence (99%). Furthermore, SelAC allows to compare parameters between fits to homologous proteins instead of relying on extrapolation.

While SelAC better explains the observed TEM sequences than the experimental estimates of site specific selection on amino acids, it is not without shortcomings itself. SelAC is currently to slow to be used in topology searches, therefore it is unclear if the differences in topology between phydms and SelAC can be attributed to the same inadequacies of experimentally inferred selection. The formulation and implementation of SelAC can and should be improved upon as the simulation of TEM evolution from the ancestral state under the SelAC inferred site specific selection revealed. Starting from the ancestral sequence, the simulated sequences diverge despite stabilizing selection for the optimal amino acid, indicating that SelAC may underestimate selection. While SelAC allows for site heterogeneity in selection for amino acids, it still ignores epistasis. This however, is a shortcoming that is shared with experimental estimates as each mutant typically only carries one mutation (FIRNBERG and OSTERMEIER, 2012; JAIN and VARADARAJAN, 2014). Furthermore, not every protein is under stabilizing selection, however, SelAC is a model stabilizing selection and may therefore not be adequate for every protein. TEM plays a role in chemical warfare with conspecifics and other microbes, therefore some sites may be under negative frequency dependent selection. This potential heterogeneity in selection highlights another shortcoming of SelAC. SelAC assumes the same distribution for the efficacy of selection G and physicochemical sensitivities across the whole protein. However, it possible that residues in different secondary structures or at active sites do not share a common distribution.

As SelAC assumes that the fitness of an amino acid at a site declines with its distance in physicochemical space to the optimal amino acid, the choice of physicochemical properties becomes important. In this study, we used composition, polarity, and molecular volume (Grantham, 1974) for all sites and only estimated their weighting. However, a wide range of additional physicochemical properties of amino acids have been described (Kawashima et al., 2008). A more optimal choice of physicochemical properties may be possible as well as relaxing the assumption that the same properties apply to all sites equally.

In conclusion, experimental estimates of site specific selection on amino acids have to be treated with skepticism and their adequacy should be assessed before using them to inform phylogenetic inferences. This study was initiated to assess the quality of SelAC with the expectation that SelAC could be a faster, cheaper, and more readily available alternative to experimentally inferred selection; specifically in organisms where these experiments are not feasible. Intuitively one would expect that selection coefficients of mutations estimated in living organisms would provide more information on the evolution of proteins than a model relying on many simplifying assumptions. As we show in this study, not only can SelAC estimate site specific selection on amino acids but our approach is a more adequate description of selection on amino acids in nature than experimental estimates.

#### 4.5 Materials and Methods

#### 4.5.1 Phylogenetic Inference and Model selection

TEM and SHV sequences were obtained from Bloom (2017) already aligned. We separated the TEM and SHV sequences into individual alignments. Experimentally fitness values for TEM were taken from Stiffler et al. (2016). We followed (Bloom, 2017) to convert the experimental fitness values into site specific equilibrium frequencies for phydms. phydms (version 2.5.1) was fitted to the site specific selection from Stiffler et al. (2016) using python (version 3.6). SelAC (version 1.6.1) was fitted to the TEM alignment using R (version 3.4.1) (R Core Team, 2015) with and without experimental site specific selection. We assumed the physicochemical properties estimated by Grantham (1974). We choose the constraint free general unrestricted model (Yang, 1994) as mutation model for SelAC. All other models were fitted using IQTree (Nguyen et al., 2015). We report each model's  $\log(\mathcal{L})$ , AIC, and AICc. Models were selected based on the AICc values.

#### 4.5.2 Sequence Simulation

Sequences were simulated by stochastic simulations using a Gillespie algorithm (GILLESPIE, 1976) that was model independent. To calculate fixation probabilities during the simulation we followed Sella and Hirsh (2005). The fitness values were estimated using SelAC or taken from Stiffler et al. (2016). We choose the fitness values resulting from the highest concentration (2500  $\mu g/mL$ ) treatment of ampicillin for our comparison. We rescaled the experimental fitness such that the amino acid with the highest fitness at each site has a value of one. Mutation rates for the simulations were taken from the SelAC or SelAC+DMS fit, respectively. The initial sequences were either a random sequence sampled with uniform codon probabilities or the ancestral sequence reconstructed using FastML (Ashkenazy et al., 2012) (last accessed: 30.09.2018). Each sequence was simulated 10 times and we report average genetic load and sequence similarity and the standard error. The sequences were sampled at times 0.01, 0.1, 1, and 10 expected mutations per site.

#### **4.5.3** Estimating site specific efficacy of selection G

SelAC does not by default estimate site specific values for G but assumes G values follow a  $\Gamma$ -distribution (Felsenstein, 2001). Site specific values for G were optimized by fixing all estimated parameters and performing a maximum likelihood search without the integration over G. In contrast to SelAC that assumes G to be purely positive, we allowed negative values for G but constraint the search to values between -300 and 300 to ensure numerical stability.

#### 4.5.4 Estimating site specific fitness values $w_i$

Following BEAULIEU et al. (in review)  $w_i$  is proportional to

$$w_i \propto \exp(-A_0 \eta \psi) \tag{4.1}$$

where  $A_0$  describes the decline in fitness with each high energy phosphate bond wasted per unit time, and  $\psi$  is the protein's production rate.  $\eta$  is the cost/benefit ratio of a protein (see (BEAULIEU et al., in review) for details). However, SelAC only estimates a composition parameter  $\psi' = A_0 \psi N_e$  thus

$$\psi = \frac{\psi'}{A_0 N_e q} \tag{4.2}$$

SelAC assumes that the effective population size  $N_e = 5 \times 10^6$  and that  $A_0 = 4 \times 10-7$  (GILCHRIST, 2007).

#### 4.5.5 Model Adequacy

Model adequacy was assessed by comparing the observed sequences and simulations under the site specific selection inferred by the deep mutation scanning experiment or SelAC. First, similarity between the sequence of selectively favored amino acids and the observed TEM sequences was assessed. Sequence similarity was measured as the number of differences in the aligned amino acid sequences. Second, the genetic load of the observed and the simulated sequences was calculated using either the site specific selection inferred by the deep mutation scanning experiment or SelAC. The average genetic load for site i in the alignment was calculated as

$$L_i = \frac{w_{max,i} - \overline{w_i}}{w_{max,i}} \tag{4.3}$$

where  $w_{max,i}$  is the fitness of the selectively favored amino acids at position i, either estimated using the site specific selection inferred by DMS or SelAC.  $\overline{w_i}$  represents the average fitness of the residues observed at position i. The average sequence specific genetic load L was calculated as the sum of the site specific genetic loads  $L = \frac{1}{n} \sum_{i=1}^{n} L_i$  where n is the number of amino acid sites.

### 4.6 Acknowledgments

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# 4.7 Appendix: Supplementary Material

Table 4.3: Model selection of 230 models of nucleotide and codon evolution.

| No. | Model        | LnL       | n   | AIC      | $\Delta { m AIC}$ | AICc     | $\Delta AICc$ |
|-----|--------------|-----------|-----|----------|-------------------|----------|---------------|
| 1   | SelAC+DMS+G4 | -1768     | 111 | 3758     | 14                | 3760     | 0             |
| 2   | SelAC + G4   | -1498     | 374 | 3744     | 0                 | 3766     | 6             |
| 3   | phydms       | -2060.85  | 102 | 4326     | 582               | 4328     | 568           |
| 4   | SYM+R2       | -2229.616 | 102 | 4663.232 | 919.232           | 4693.862 | 933.862       |
| 5   | TIMe+R2      | -2232.406 | 100 | 4664.811 | 920.811           | 4694.172 | 934.172       |
| 6   | TVMe+R2      | -2232.838 | 101 | 4667.677 | 923.677           | 4697.668 | 937.668       |
| 7   | TIM3e+R2     | -2234.332 | 100 | 4668.664 | 924.664           | 4698.024 | 938.024       |
| 8   | TIM2e+R2     | -2234.381 | 100 | 4668.763 | 924.763           | 4698.123 | 938.123       |
| 9   | K3P+R2       | -2235.777 | 99  | 4669.553 | 925.553           | 4698.291 | 938.291       |
| 10  | TNe+R2       | -2236.078 | 99  | 4670.155 | 926.155           | 4698.892 | 938.892       |
| 11  | SYM+R3       | -2229.616 | 104 | 4667.232 | 923.232           | 4699.162 | 939.162       |
| 12  | TIM+F+R2     | -2230.958 | 103 | 4667.915 | 923.915           | 4699.191 | 939.191       |
| 13  | TIMe+R3      | -2232.404 | 102 | 4668.808 | 924.808           | 4699.437 | 939.437       |
| 14  | GTR+F+R2     | -2228.537 | 105 | 4667.073 | 923.073           | 4699.665 | 939.665       |
| 15  | K3Pu+F+R2    | -2232.617 | 102 | 4669.234 | 925.234           | 4699.864 | 939.864       |
| 16  | TVM+F+R2     | -2230.105 | 104 | 4668.21  | 924.21            | 4700.14  | 940.14        |
| 17  | TVMe+R3      | -2232.838 | 103 | 4671.676 | 927.676           | 4702.952 | 942.952       |
| 18  | K2P+R2       | -2239.424 | 98  | 4674.847 | 930.847           | 4702.969 | 942.969       |
| 19  | TIM3e+R3     | -2234.332 | 102 | 4672.664 | 928.664           | 4703.293 | 943.293       |
| 20  | TIM2e+R3     | -2234.381 | 102 | 4672.762 | 928.762           | 4703.391 | 943.391       |
| 21  | TIM3+F+R2    | -2233.064 | 103 | 4672.127 | 928.127           | 4703.403 | 943.403       |
| 22  | TIM2+F+R2    | -2233.114 | 103 | 4672.227 | 928.227           | 4703.503 | 943.503       |
| 23  | K3P+R3       | -2235.777 | 101 | 4673.553 | 929.553           | 4703.545 | 943.545       |
| 24  | TN+F+R2      | -2234.624 | 102 | 4673.249 | 929.249           | 4703.878 | 943.878       |
| 25  | TPM3u+F+R2   | -2234.673 | 102 | 4673.347 | 929.347           | 4703.977 | 943.977       |
| 26  | TPM3+F+R2    | -2234.674 | 102 | 4673.348 | 929.348           | 4703.978 | 943.978       |
| 27  | TPM2u+F+R2   | -2234.681 | 102 | 4673.363 | 929.363           | 4703.993 | 943.993       |
| 28  | TPM2+F+R2    | -2234.683 | 102 | 4673.365 | 929.365           | 4703.995 | 943.995       |
| 29  | TNe+R3       | -2236.077 | 101 | 4674.155 | 930.155           | 4704.146 | 944.146       |
| 30  | TIM+F+R3     | -2230.958 | 105 | 4671.915 | 927.915           | 4704.507 | 944.507       |
| 31  | HKY+F+R2     | -2236.266 | 101 | 4674.531 | 930.531           | 4704.522 | 944.522       |
| 32  | GTR+F+R3     | -2228.536 | 107 | 4671.073 | 927.073           | 4705.011 | 945.011       |
| 33  | K3Pu+F+R3    | -2232.617 | 104 | 4673.234 | 929.234           | 4705.163 | 945.163       |
| 34  | TVM+F+R3     | -2230.105 | 106 | 4672.21  | 928.21            | 4705.471 | 945.471       |
| 35  | K2P+R3       | -2239.192 | 100 | 4678.384 | 934.384           | 4707.745 | 947.745       |
| 36  | TIM3+F+R3    | -2233.063 | 105 | 4676.127 | 932.127           | 4708.718 | 948.718       |
| 37  | TIM2+F+R3    | -2233.113 | 105 | 4676.227 | 932.227           | 4708.818 | 948.818       |

Table 4.3 Continued

| No.              | Model        | LnL       | n   | AIC      | $\Delta { m AIC}$ | AICc     | $\Delta$ AICc             |
|------------------|--------------|-----------|-----|----------|-------------------|----------|---------------------------|
| 38               | TN+F+R3      | -2234.624 | 104 | 4677.249 | 933.249           | 4709.178 | $\frac{249.178}{949.178}$ |
| 39               | TPM3u+F+R3   | -2234.673 | 104 | 4677.347 | 933.347           | 4709.277 | 949.277                   |
| 40               | TPM3+F+R3    | -2234.674 | 104 | 4677.348 | 933.348           | 4709.277 | 949.277                   |
| 41               | TPM2u+F+R3   | -2234.681 | 104 | 4677.363 | 933.363           | 4709.293 | 949.293                   |
| 42               | TPM2+F+R3    | -2234.682 | 104 | 4677.364 | 933.364           | 4709.294 | 949.294                   |
| 43               | HKY+F+R3     | -2236.074 | 103 | 4678.148 | 934.148           | 4709.424 | 949.424                   |
| 44               | SYM+I+G4     | -2243.212 | 102 | 4690.424 | 946.424           | 4721.054 | 961.054                   |
| 45               | TVMe+I+G4    | -2244.533 | 101 | 4691.066 | 947.066           | 4721.057 | 961.057                   |
| 46               | TIMe+I+G4    | -2246.457 | 100 | 4692.914 | 948.914           | 4722.275 | 962.275                   |
| 47               | K3P+I+G4     | -2248.166 | 99  | 4694.332 | 950.332           | 4723.069 | 963.069                   |
| 48               | TVM+F+I+G4   | -2241.853 | 104 | 4691.707 | 947.707           | 4723.636 | 963.636                   |
| 49               | TIM3e+I+G4   | -2247.379 | 100 | 4694.758 | 950.758           | 4724.119 | 964.119                   |
| 50               | K3Pu+F+I+G4  | -2245.156 | 102 | 4694.311 | 950.311           | 4724.941 | 964.941                   |
| 51               | GTR+F+I+G4   | -2241.484 | 105 | 4692.968 | 948.968           | 4725.559 | 965.559                   |
| 52               | TIM+F+I+G4   | -2244.418 | 103 | 4694.836 | 950.836           | 4726.112 | 966.112                   |
| 53               | TPM3u+F+I+G4 | -2246.03  | 102 | 4696.06  | 952.06            | 4726.69  | 966.69                    |
| 54               | TPM3+F+I+G4  | -2246.069 | 102 | 4696.138 | 952.138           | 4726.768 | 966.768                   |
| 55               | TIM2e+I+G4   | -2248.934 | 100 | 4697.868 | 953.868           | 4727.228 | 967.228                   |
| 56               | TNe+I+G4     | -2250.587 | 99  | 4699.174 | 955.174           | 4727.911 | 967.911                   |
| 57               | TIM3+F+I+G4  | -2245.534 | 103 | 4697.068 | 953.068           | 4728.344 | 968.344                   |
| 58               | K2P+I+G4     | -2252.181 | 98  | 4700.362 | 956.362           | 4728.484 | 968.484                   |
| 59               | TPM2u+F+I+G4 | -2247.579 | 102 | 4699.158 | 955.158           | 4729.788 | 969.788                   |
| 60               | TPM2+F+I+G4  | -2247.685 | 102 | 4699.371 | 955.371           | 4730     | 970                       |
| 61               | HKY+F+I+G4   | -2249.065 | 101 | 4700.13  | 956.13            | 4730.121 | 970.121                   |
| 62               | TIM2+F+I+G4  | -2247.009 | 103 | 4700.018 | 956.018           | 4731.294 | 971.294                   |
| 63               | TN+F+I+G4    | -2248.511 | 102 | 4701.023 | 957.023           | 4731.652 | 971.652                   |
| 64               | TVMe+I       | -2254.804 | 100 | 4709.608 | 965.608           | 4738.968 | 978.968                   |
| 65               | K3P+I        | -2257.72  | 98  | 4711.439 | 967.439           | 4739.561 | 979.561                   |
| 66               | SYM+I        | -2254.11  | 101 | 4710.221 | 966.220           | 4740.212 | 980.212                   |
| 67               | TIMe+I       | -2257.074 | 99  | 4712.149 | 968.149           | 4740.886 | 980.886                   |
| 68               | TVM+F+I      | -2252.157 | 103 | 4710.315 | 966.315           | 4741.591 | 981.591                   |
| 69               | K3Pu+F+I     | -2254.856 | 101 | 4711.712 | 967.712           | 4741.704 | 981.704                   |
| 70               | TIM3e+I      | -2257.796 | 99  | 4713.592 | 969.592           | 4742.33  | 982.33                    |
| 71               | TPM3+F+I     | -2255.771 | 101 | 4713.543 | 969.543           | 4743.534 | 983.534                   |
| 72               | TPM3u+F+I    | -2255.771 | 101 | 4713.543 | 969.543           | 4743.534 | 983.534                   |
| 73               | K2P+I        | -2261.218 | 97  | 4716.436 | 972.436           | 4743.949 | 983.949                   |
| 74               | GTR+F+I      | -2252.067 | 104 | 4712.133 | 968.133           | 4744.063 | 984.063                   |
| 75<br>           | TIM+F+I      | -2254.783 | 102 | 4713.566 | 969.566           | 4744.195 | 984.195                   |
| 76               | TNe+I        | -2260.579 | 98  | 4717.158 | 973.158           | 4745.28  | 985.28                    |
| 77<br><b>-</b> 0 | TIM3+F+I     | -2255.684 | 102 | 4715.368 | 971.368           | 4745.998 | 985.998                   |
| 78<br><b>7</b> 8 | HKY+F+I      | -2258.352 | 100 | 4716.703 | 972.703           | 4746.064 | 986.064                   |
| 79               | TIM2e+I      | -2259.878 | 99  | 4717.757 | 973.757           | 4746.494 | 986.494                   |

Table 4.3 Continued

| No. | Model            | LnL       | n   | AIC      | $\Delta { m AIC}$ | AICc     | $\Delta$ AICc |
|-----|------------------|-----------|-----|----------|-------------------|----------|---------------|
| 80  | TVMe+G4          | -2258.853 | 100 | 4717.705 | 973.705           | 4747.066 | 987.066       |
| 81  | SYM+G4           | -2257.573 | 101 | 4717.146 | 973.146           | 4747.137 | 987.137       |
| 82  | TPM2+F+I         | -2257.712 | 101 | 4717.423 | 973.423           | 4747.415 | 987.415       |
| 83  | TPM2u+F+I        | -2257.712 | 101 | 4717.423 | 973.423           | 4747.415 | 987.415       |
| 84  | K3P+G4           | -2261.922 | 98  | 4719.844 | 975.844           | 4747.966 | 987.966       |
| 85  | TIMe+G4          | -2260.683 | 99  | 4719.365 | 975.365           | 4748.103 | 988.103       |
| 86  | TN+F+I           | -2258.28  | 101 | 4718.561 | 974.561           | 4748.552 | 988.552       |
| 87  | TIM3e+G4         | -2261.255 | 99  | 4720.51  | 976.51            | 4749.247 | 989.247       |
| 88  | TVM+F+G4         | -2256.108 | 103 | 4718.216 | 974.216           | 4749.492 | 989.492       |
| 89  | TIM2+F+I         | -2257.643 | 102 | 4719.286 | 975.286           | 4749.915 | 989.915       |
| 90  | K3Pu+F+G4        | -2258.971 | 101 | 4719.941 | 975.941           | 4749.933 | 989.933       |
| 91  | TPM3u+F+G4       | -2259.716 | 101 | 4721.433 | 977.433           | 4751.424 | 991.424       |
| 92  | TPM3+F+G4        | -2259.717 | 101 | 4721.434 | 977.434           | 4751.425 | 991.425       |
| 93  | GTR+F+G4         | -2255.75  | 104 | 4719.5   | 975.5             | 4751.43  | 991.43        |
| 94  | TIM+F+G4         | -2258.638 | 102 | 4721.276 | 977.276           | 4751.906 | 991.906       |
| 95  | K2P+G4           | -2265.454 | 97  | 4724.907 | 980.907           | 4752.421 | 992.421       |
| 96  | TNe+G4           | -2264.219 | 98  | 4724.437 | 980.437           | 4752.559 | 992.559       |
| 97  | TIM3+F+G4        | -2259.366 | 102 | 4722.732 | 978.732           | 4753.361 | 993.361       |
| 98  | TIM2e+G4         | -2263.57  | 99  | 4725.141 | 981.141           | 4753.878 | 993.878       |
| 99  | JC+R2            | -2266.233 | 97  | 4726.466 | 982.466           | 4753.98  | 993.98        |
| 100 | F81+F+R2         | -2262.327 | 100 | 4724.654 | 980.654           | 4754.015 | 994.015       |
| 101 | HKY+F+G4         | -2262.499 | 100 | 4724.999 | 980.999           | 4754.359 | 994.359       |
| 102 | TPM2+F+G4        | -2261.915 | 101 | 4725.829 | 981.829           | 4755.82  | 995.82        |
| 103 | TPM2u+F+G4       | -2261.915 | 101 | 4725.829 | 981.829           | 4755.82  | 995.82        |
| 104 | TN+F+G4          | -2262.169 | 101 | 4726.338 | 982.338           | 4756.329 | 996.329       |
| 105 | TIM2+F+G4        | -2261.585 | 102 | 4727.17  | 983.17            | 4757.8   | 997.8         |
| 106 | F81+F+R3         | -2262.028 | 102 | 4728.056 | 984.056           | 4758.685 | 998.685       |
| 107 | JC+R3            | -2265.997 | 99  | 4729.994 | 985.994           | 4758.731 | 998.731       |
| 108 | F81+F+I+G4       | -2274.845 | 100 | 4749.69  | 1005.69           | 4779.05  | 1019.05       |
| 109 | JC+I+G4          | -2279.318 | 97  | 4752.636 | 1008.636          | 4780.149 | 1020.149      |
| 110 | F81+F+I          | -2283.56  | 99  | 4765.119 | 1021.119          | 4793.857 | 1033.857      |
| 111 | JC+I             | -2287.984 | 96  | 4767.968 | 1023.968          | 4794.881 | 1034.881      |
| 112 | F81+F+G4         | -2287.834 | 99  | 4773.669 | 1029.669          | 4802.406 | 1042.406      |
| 113 | JC+G4            | -2292.095 | 96  | 4776.19  | 1032.19           | 4803.103 | 1043.103      |
| 114 | GY94 +F1X4+R2    | -2242.963 | 102 | 4689.926 | 945.926           | 4821.251 | 1061.251      |
| 115 | MGK+F1X4+R2      | -2243.111 | 102 | 4690.221 | 946.221           | 4821.546 | 1061.546      |
| 116 | GY94 + F1X4 + R3 | -2238.022 | 104 | 4684.043 | 940.043           | 4822.271 | 1062.271      |
| 117 | MGK+F3X4+R2      | -2229.923 | 108 | 4675.846 | 931.846           | 4828.729 | 1068.729      |
| 118 | GY94 +F1X4+I+G4  | -2247.179 | 102 | 4698.359 | 954.359           | 4829.684 | 1069.684      |
| 119 | MGK+F1X4+I+G4    | -2247.292 | 102 | 4698.583 | 954.583           | 4829.908 | 1069.908      |
| 120 | MGK+F1X4+R3      | -2241.989 | 104 | 4691.978 | 947.978           | 4830.206 | 1070.206      |
| 121 | MGK+F3X4+R3      | -2224.78  | 110 | 4669.559 | 925.559           | 4830.217 | 1070.217      |

Table 4.3 Continued

|     | 3.6.1.1              | T 7        |     | A T C    | A A T C      | A T C    | A A T C       |
|-----|----------------------|------------|-----|----------|--------------|----------|---------------|
| No. | Model                | <u>LnL</u> | n   | AIC      | $\Delta AIC$ | AICc     | $\Delta AICc$ |
| 122 | GY94 +F1X4+G4        | -2251.144  | 101 | 4704.287 | 960.287      | 4832.263 | 1072.263      |
| 123 | MGK+F1X4+G4          | -2251.472  | 101 | 4704.944 | 960.944      | 4832.919 | 1072.919      |
| 124 | GY94 + F3X4 + R3     | -2227.048  | 110 | 4674.096 | 930.096      | 4834.754 | 1074.754      |
| 125 | GY94 + F3X4 + R2     | -2233.068  | 108 | 4682.136 | 938.136      | 4835.019 | 1075.019      |
| 126 | MGK+F3X4+I+G4        | -2233.539  | 108 | 4683.078 | 939.0781     | 4835.962 | 1075.962      |
| 127 | MGK+F3X4+G4          | -2237.512  | 107 | 4689.024 | 945.024      | 4838.134 | 1078.134      |
| 128 | GY94 + F3X4 + I + G4 | -2238.243  | 108 | 4692.485 | 948.485      | 4845.368 | 1085.368      |
| 129 | GY94 + F3X4 + R4     | -2227.106  | 112 | 4678.213 | 934.213      | 4846.96  | 1086.96       |
| 130 | GY94 + F3X4 + G4     | -2242.394  | 107 | 4698.789 | 954.789      | 4847.899 | 1087.899      |
| 131 | GY94 + F1X4 + I      | -2260.085  | 101 | 4722.169 | 978.169      | 4850.144 | 1090.144      |
| 132 | MGK+F1X4+I           | -2260.345  | 101 | 4722.69  | 978.69       | 4850.665 | 1090.665      |
| 133 | MGK+F3X4+I           | -2246.112  | 107 | 4706.225 | 962.225      | 4855.335 | 1095.335      |
| 134 | MG+F1X4+R2           | -2268.482  | 101 | 4738.963 | 994.963      | 4866.938 | 1106.938      |
| 135 | GY94 + F3X4 + I      | -2252.532  | 107 | 4719.064 | 975.064      | 4868.174 | 1108.174      |
| 136 | MG+F3X4+R2           | -2254.453  | 107 | 4722.906 | 978.906      | 4872.015 | 1112.015      |
| 137 | MG+F1X4+I+G4         | -2272.057  | 101 | 4746.113 | 1002.113     | 4874.089 | 1114.089      |
| 138 | MG+F1X4+R3           | -2267.523  | 103 | 4741.047 | 997.047      | 4875.789 | 1115.789      |
| 139 | MG+F1X4+G4           | -2276.171  | 100 | 4752.342 | 1008.342     | 4877.033 | 1117.033      |
| 140 | MG+F3X4+I+G4         | -2257.945  | 107 | 4729.891 | 985.891      | 4879.001 | 1119.001      |
| 141 | MG+F3X4+G4           | -2261.949  | 106 | 4735.898 | 991.898      | 4881.309 | 1121.309      |
| 142 | MG+F3X4+R3           | -2253.514  | 109 | 4725.027 | 981.027      | 4881.759 | 1121.759      |
| 143 | SYM                  | -2329.878  | 100 | 4859.756 | 1115.756     | 4889.116 | 1129.116      |
| 144 | TIMe                 | -2333.105  | 98  | 4862.21  | 1118.21      | 4890.332 | 1130.332      |
| 145 | TIM3e                | -2333.481  | 98  | 4862.961 | 1118.961     | 4891.083 | 1131.083      |
| 146 | TVMe                 | -2333.164  | 99  | 4864.328 | 1120.328     | 4893.065 | 1133.065      |
| 147 | GTR+F                | -2328.404  | 103 | 4862.809 | 1118.809     | 4894.085 | 1134.085      |
| 148 | K3P                  | -2336.391  | 97  | 4866.783 | 1122.783     | 4894.297 | 1134.297      |
| 149 | MG+F1X4+I            | -2284.946  | 100 | 4769.892 | 1025.892     | 4894.583 | 1134.583      |
| 150 | TVM+F                | -2330.086  | 102 | 4864.172 | 1120.172     | 4894.802 | 1134.802      |
| 151 | TIM+F                | -2331.48   | 101 | 4864.96  | 1120.96      | 4894.952 | 1134.952      |
| 152 | TNe                  | -2336.729  | 97  | 4867.458 | 1123.458     | 4894.972 | 1134.972      |
| 153 | K3Pu+F               | -2333.162  | 100 | 4866.323 | 1122.323     | 4895.684 | 1135.684      |
| 154 | TIM3+F               | -2331.971  | 101 | 4865.942 | 1121.942     | 4895.934 | 1135.934      |
| 155 | TPM3+F               | -2333.648  | 100 | 4867.297 | 1123.297     | 4896.657 | 1136.657      |
| 156 | TPM3u+F              | -2333.648  | 100 | 4867.297 | 1123.297     | 4896.657 | 1136.657      |
| 157 | TIM2e                | -2336.292  | 98  | 4868.584 | 1124.584     | 4896.706 | 1136.706      |
| 158 | MG+F3X4+I            | -2270.442  | 106 | 4752.885 | 1008.885     | 4898.295 | 1138.295      |
| 159 | K2P                  | -2340.015  | 96  | 4872.03  | 1128.03      | 4898.943 | 1138.943      |
| 160 | TN+F                 | -2335.102  | 100 | 4870.204 | 1126.204     | 4899.565 | 1139.565      |
| 161 | HKY+F                | -2336.783  | 99  | 4871.566 | 1127.566     | 4900.303 | 1140.303      |
| 162 | TIM2+F               | -2334.7    | 101 | 4871.401 | 1127.401     | 4901.392 | 1141.392      |
| 163 | TPM2u+F              | -2336.381  | 100 | 4872.761 | 1128.761     | 4902.122 | 1142.122      |
|     |                      |            |     |          |              |          |               |

Table 4.3 Continued

| No. | Model            | LnL       | n   | AIC      | $\Delta { m AIC}$ | AICc     | $\Delta { m AICc}$ |
|-----|------------------|-----------|-----|----------|-------------------|----------|--------------------|
| 164 | TPM2+F           | -2336.381 | 100 | 4872.762 | 1128.762          | 4902.123 | 1142.123           |
| 165 | $_{ m JC}$       | -2366.286 | 95  | 4922.571 | 1178.571          | 4948.892 | 1188.892           |
| 166 | F81+F            | -2362.554 | 98  | 4921.108 | 1177.108          | 4949.229 | 1189.229           |
| 167 | GY94 + F1X4      | -2315.788 | 100 | 4831.575 | 1087.575          | 4956.267 | 1196.267           |
| 168 | KOSI07+FU+R2     | -2325.725 | 97  | 4845.45  | 1101.45           | 4960.675 | 1200.675           |
| 169 | MGK+F1X4         | -2318.048 | 100 | 4836.095 | 1092.095          | 4960.787 | 1200.787           |
| 170 | KOSI07+FU+R3     | -2323.063 | 99  | 4844.126 | 1100.126          | 4965.599 | 1205.599           |
| 171 | MGK+F3X4         | -2304.357 | 106 | 4820.713 | 1076.713          | 4966.124 | 1206.124           |
| 172 | GY94 + F3X4      | -2306.17  | 106 | 4824.339 | 1080.339          | 4969.749 | 1209.749           |
| 173 | KOSI07+FU+I+G4   | -2335.554 | 97  | 4865.108 | 1121.108          | 4980.332 | 1220.332           |
| 174 | KOSI07+FU+G4     | -2339.513 | 96  | 4871.026 | 1127.026          | 4983.218 | 1223.218           |
| 175 | KOSI07+F3X4+R2   | -2315.814 | 106 | 4843.627 | 1099.627          | 4989.038 | 1229.038           |
| 176 | KOSI07+F3X4+R3   | -2310.509 | 108 | 4837.018 | 1093.018          | 4989.901 | 1229.901           |
| 177 | KOSI07+F1X4+R2   | -2333.491 | 100 | 4866.983 | 1122.983          | 4991.674 | 1231.674           |
| 178 | KOSI07+F1X4+R3   | -2328.692 | 102 | 4861.383 | 1117.383          | 4992.708 | 1232.708           |
| 179 | SCHN05+FU+R2     | -2344.705 | 97  | 4883.411 | 1139.411          | 4998.635 | 1238.635           |
| 180 | KOSI07+F1X4+I+G4 | -2337.965 | 100 | 4875.93  | 1131.93           | 5000.621 | 1240.621           |
| 181 | KOSI07+F1X4+G4   | -2341.156 | 99  | 4880.312 | 1136.312          | 5001.784 | 1241.784           |
| 182 | SCHN05+FU+R3     | -2341.179 | 99  | 4880.358 | 1136.358          | 5001.831 | 1241.831           |
| 183 | KOSI07+FU+I      | -2349.617 | 96  | 4891.233 | 1147.233          | 5003.426 | 1243.426           |
| 184 | KOSI07+F3X4+I+G4 | -2323.767 | 106 | 4859.534 | 1115.534          | 5004.944 | 1244.944           |
| 185 | MG+F1X4          | -2342.797 | 99  | 4883.593 | 1139.593          | 5005.065 | 1245.065           |
| 186 | KOSI07+F3X4+G4   | -2327.376 | 105 | 4864.751 | 1120.751          | 5006.534 | 1246.534           |
| 187 | MG+F3X4          | -2328.539 | 105 | 4867.078 | 1123.078          | 5008.861 | 1248.861           |
| 188 | SCHN05+F1X4+R3   | -2340.927 | 102 | 4885.854 | 1141.854          | 5017.179 | 1257.179           |
| 189 | KOSI07+F1X4+I    | -2349.1   | 99  | 4896.2   | 1152.2            | 5017.672 | 1257.672           |
| 190 | SCHN05+F3X4+R3   | -2324.472 | 108 | 4864.944 | 1120.944          | 5017.827 | 1257.827           |
| 191 | SCHN05+FU+I+G4   | -2354.523 | 97  | 4903.046 | 1159.046          | 5018.27  | 1258.27            |
| 192 | SCHN05+F1X4+R2   | -2348.226 | 100 | 4896.452 | 1152.452          | 5021.143 | 1261.143           |
| 193 | SCHN05+F3X4+R2   | -2331.916 | 106 | 4875.833 | 1131.833          | 5021.243 | 1261.243           |
| 194 | SCHN05+FU+G4     | -2358.682 | 96  | 4909.365 | 1165.365          | 5021.558 | 1261.558           |
| 195 | KOSI07+F3X4+I    | -2336.826 | 105 | 4883.653 | 1139.653          | 5025.436 | 1265.436           |
| 196 | SCHN05+F1X4+I+G4 | -2351.096 | 100 | 4902.192 | 1158.192          | 5026.883 | 1266.883           |
| 197 | SCHN05+F1X4+G4   | -2353.895 | 99  | 4905.79  | 1161.79           | 5027.263 | 1267.263           |
| 198 | SCHN05+F1X4+R4   | -2340.593 | 104 | 4889.187 | 1145.187          | 5027.414 | 1267.414           |
| 199 | SCHN05+F3X4+R4   | -2324.102 | 110 | 4868.203 | 1124.203          | 5028.861 | 1268.861           |
| 200 | SCHN05+F3X4+I+G4 | -2338.345 | 106 | 4888.69  | 1144.69           | 5034.101 | 1274.101           |
| 201 | SCHN05+F3X4+G4   | -2341.811 | 105 | 4893.621 | 1149.621          | 5035.404 | 1275.404           |
| 202 | SCHN05+FU+I      | -2370.471 | 96  | 4932.943 | 1188.943          | 5045.135 | 1285.135           |
| 203 | SCHN05+F1X4+I    | -2363.696 | 99  | 4925.391 | 1181.391          | 5046.864 | 1286.864           |
| 204 | SCHN05+F3X4+I    | -2352.81  | 105 | 4915.621 | 1171.621          | 5057.404 | 1297.404           |
| 205 | KOSI07+FU        | -2394.782 | 95  | 4979.563 | 1235.563          | 5088.785 | 1328.785           |

Table 4.3 Continued

| No. | Model         | LnL       | n   | AIC      | $\Delta { m AIC}$ | AICc     | $\Delta { m AICc}$ |
|-----|---------------|-----------|-----|----------|-------------------|----------|--------------------|
| 206 | KOSI07+F1X4   | -2398.44  | 98  | 4992.88  | 1248.88           | 5111.197 | 1351.197           |
| 207 | KOSI07+F3X4   | -2383.159 | 104 | 4974.318 | 1230.318          | 5112.546 | 1352.546           |
| 208 | SCHN05+FU     | -2419.333 | 95  | 5028.665 | 1284.665          | 5137.887 | 1377.887           |
| 209 | SCHN05+F1X4   | -2416.544 | 98  | 5029.088 | 1285.088          | 5147.405 | 1387.405           |
| 210 | SCHN05+F3X4   | -2402.838 | 104 | 5013.675 | 1269.675          | 5151.903 | 1391.903           |
| 211 | GY94 + F + R2 | -2208.59  | 159 | 4735.181 | 991.181           | 5229.161 | 1469.161           |
| 212 | GY94 + F+G4   | -2217.694 | 158 | 4751.388 | 1007.388          | 5234.504 | 1474.504           |
| 213 | GY94 +F+I+G4  | -2213.659 | 159 | 4745.319 | 1001.319          | 5239.299 | 1479.299           |
| 214 | GY94 + F + R3 | -2202.599 | 161 | 4727.198 | 983.198           | 5243.673 | 1483.673           |
| 215 | GY94 + F + I  | -2228.346 | 158 | 4772.691 | 1028.691          | 5255.807 | 1495.807           |
| 216 | GY94 + F + R4 | -2202.61  | 163 | 4731.219 | 987.219           | 5271.26  | 1511.26            |
| 217 | GY94 + F      | -2282.254 | 157 | 4878.509 | 1134.509          | 5351.004 | 1591.004           |
| 218 | KOSI07+F+R2   | -2291.643 | 157 | 4897.286 | 1153.286          | 5369.781 | 1609.781           |
| 219 | KOSI07+F+G4   | -2301.662 | 156 | 4915.325 | 1171.325          | 5377.438 | 1617.438           |
| 220 | KOSI07+F+I+G4 | -2298.418 | 157 | 4910.835 | 1166.835          | 5383.33  | 1623.33            |
| 221 | KOSI07+F+R3   | -2286.723 | 159 | 4891.446 | 1147.446          | 5385.426 | 1625.426           |
| 222 | KOSI07+F+I    | -2311.78  | 156 | 4935.559 | 1191.559          | 5397.672 | 1637.672           |
| 223 | SCHN05+F+R2   | -2310.015 | 157 | 4934.03  | 1190.03           | 5406.525 | 1646.525           |
| 224 | SCHN05+F+G4   | -2316.684 | 156 | 4945.369 | 1201.369          | 5407.482 | 1647.482           |
| 225 | SCHN05+F+I+G4 | -2313.733 | 157 | 4941.467 | 1197.467          | 5413.962 | 1653.962           |
| 226 | SCHN05+F+R3   | -2303.732 | 159 | 4925.463 | 1181.463          | 5419.444 | 1659.444           |
| 227 | SCHN05+F+I    | -2327.127 | 156 | 4966.254 | 1222.254          | 5428.367 | 1668.367           |
| 228 | SCHN05+F+R4   | -2303.45  | 161 | 4928.9   | 1184.9            | 5445.375 | 1685.375           |
| 229 | KOSI07+F      | -2357.579 | 155 | 5025.157 | 1281.157          | 5477.12  | 1717.12            |
| 230 | SCHN05+F      | -2379.264 | 155 | 5068.528 | 1324.528          | 5520.491 | 1760.491           |

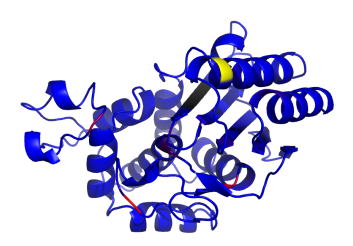


Figure 4.6: Distribution of genetic load in TEM mapped on its structure (1xpb). Average genetic load over all observed TEM variants is indicated by the color, blue low, red medium, yellow high genetic load. Active site is indicated in black.

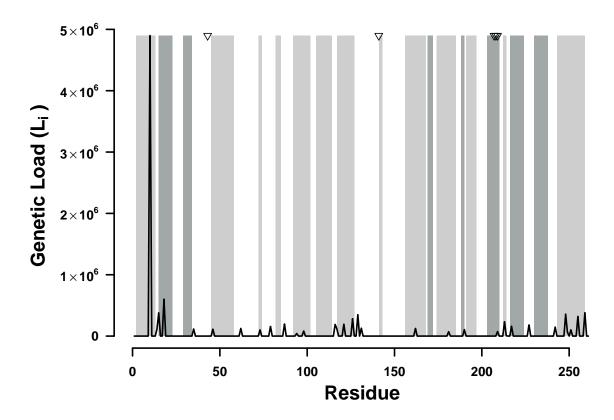


Figure 4.7: Distribution of genetic load in SHV. Average genetic load over all observed SHV variants is indicated by the black line. Light gray bars indicate where helices are found, and dark gray bars indicate  $\beta$ -sheets. The three residues forming the binding site and the two residues forming the active are indicated by triangles at the top of the plot.

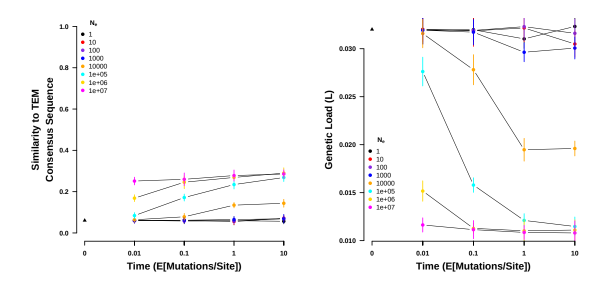


Figure 4.8: Sequences simulated from a random codon sequence under the site specific selection on amino acids estimated using SelAC. (left) Sequence similarity to the observed consensus sequence at various times for a range of values of  $N_e$ . (right) Genetic load of the simulated sequences at various times for a range of values of  $N_e$ . Time is given in number of expected mutations per site, which equals the substitution rate of a neutral mutation. Points indicate sample means and vertical bars indicate standard deviations. Initial sequence is the inferred ancestral state of the TEM variants and indicated by a black triangle.

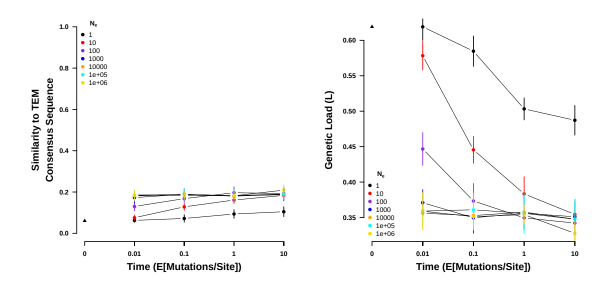


Figure 4.9: Sequences simulated from a random codon sequence under the site specific selection on amino acids estimated using deep mutation scanning. (left) Sequence similarity to the observed consensus sequence at various times for a range of values of  $N_e$ . (right) Genetic load of the simulated sequences at various times for a range of values of  $N_e$ . Time is given in number of expected mutations per site, which equals the substitution rate of a neutral mutation. Points indicate sample means and vertical bars indicate standard deviations. Initial sequence is the inferred ancestral state of the TEM variants and indicated by a black triangle.

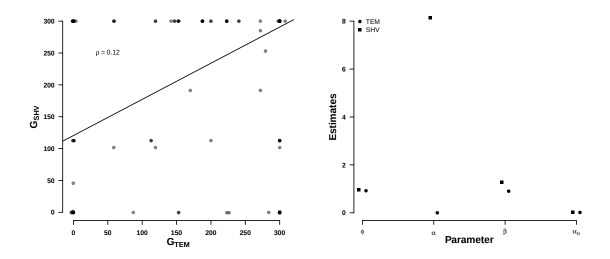


Figure 4.10: Comparison of selection related parameters between TEM and SHV. (left) Estimated site specific efficacy of selection G. (right) Selection related parameter estimates. Protein functionality production rate  $\psi$ , physicochemical weight for amino acid composition  $\alpha_c$ , physicochemical weight for amino acid polarity  $\alpha_p$ , and the parameter describing the distribution of G,  $\alpha_G$  estimated by SelAC.

## Chapter 5

### Conclusion

# 5.1 Summary

Protein synthesis from mRNA is the metabolically most expensive process a cell performs with about 20% of the cells total energy budget (REEDS et al., 1985; WATERLOW and MILLWARD, 1989). The direct cost for the translation of a protein of length L requires 4L+4 high energy phosphate bonds provided by ATP and GTP molecules. Protein synthesis is the results of a complex interplay of many different metabolic and regulatory pathways. Each step of protein synthesis is under selection and prone to errors with consequences for downstream processes. This enormous energy expenditure for the translation of a protein from mRNA leads to strong selection for efficient translation (GILCHRIST, 2007; DRUMMOND and WILKE, 2008; GILCHRIST et al., 2009; SHAH and GILCHRIST, 2011a; GILCHRIST et al., 2015). However, the efficacy of selection varies with the effective population size  $N_e$  between organisms, the rate of protein synthesis, and absolute difference in metabolic expenditure with changes in amino acid and codon usage.

On the other hand, proteins are involved in almost all processes a cell performs. From communication between cells, over the processing of metabolites, to the transport of nutrients. This ratio of cost to benefit is the fundamental concept I applied to understand and separate the effects mutation, selection, and genetic drift have on protein sequence evolution. I approached cost and benefit by applying mechanistic models rooted in first

principles to protein coding sequences. In chapter 3, I focused on the cost of protein synthesis and explored the effects of mismatched codon usage. In chapter 4, I focused on the benefit of protein synthesis and estimated site specific selection on amino acids and assessed their adequacy.

#### 5.1.1 The Value of Mechanistic Models

Mathematical and statistical models exist on a spectrum from descriptive over phenomenological to mechanistic with increasing power to extract information from data. Models allow us to summarize data and identify patterns. They are an essential tool to formalize verbal theory and allow for hypothesis testing. Well formulated models grounded in first principles can provide insights into underlying biological processes. Yet, we still have blackboxes in our models and many phenomena could lead to the model when approximated.

While descriptive and phenomenological models are important contributions to summarize processes, these models lack explanatory power. In contrast, mechanistic models allow researchers to extract information about the processes underlying the data. Mechanistic models, however, require an understanding about the underlying process which may not always be available. Even when this information is available, transition towards mechanistic models can be slow. For example, the most popular models used today to analyze codon usage are still phenomenological (IKEMURA, 1981; BENNETZEN and HALL, 1982; SHARP, 1987; WRIGHT, 1990; DOS REIS et al., 2003, 2004). While these phenomenological models provide good heuristics to explore differences in codon usage or other phenomena, they do not directly account for the evolutionary forces shaping the observed patterns such as selection, mutation, or genetic drift. Accounting for these forces allows for the proposal and testing of more sophisticated hypothesis as I demonstrate in chapter 3 and chapter 4.

## 5.1.2 Mechanistic Models Supplement Experiments

In addition to extracting information about biological processes from data, mechanistic models can help supplement experimental procedures. Empirical estimates of site specific selection are a valuable resource to e.g. identify sites conferring antibiotic resistance (Firnberg et al., 2014; Stiffler et al., 2016). While the unit that selection can act on is the amino acid, amino acids are a complex collection of physicochemical properties. It is, therefore, unclear for which properties amino acids are actually selected and when. Mechanistic models could be used to explore differences in the selection for physicochemical properties within and across proteins. Furthermore hypothesis could be formulated about the differing importance of physicochemical properties between e.g. sites or secondary structure elements

# 5.2 Estimating Protein Functional and Fitness Landscape

The selection on a protein sequence is highly complex. A protein of length L has  $20^L$  possible states it can occupy in a L dimensional fitness landscape. This enormous complexity makes it prohibitively expensive to study protein fitness landscapes without simplifying assumptions. It is therefore important to be aware of potential impacts such assumptions have on the obtained results and how models can be further improved. However, despite such simplifying assumptions, valuable information has been extracted from protein coding sequences.

# 5.2.1 The Importance of Translation Errors

We often think of genes evolving with natural selection favoring proteins that encode their function optimally, with mutations and genetic drift reducing protein functionality. The error rate of protein synthesis is five to six orders of magnitude higher than mutations, Tawfik, 2009; Drummond and Wilke, 2009), creating more erroneous high expression proteins such as ribosomal proteins than error free low expression proteins. Selection on a gene is, therefore, not based solely on the error free protein sequence, but on the average fitness of the population of proteins resulting from a gene by means of error prone protein synthesis. Previous work showed that proteins with functionality essential to an organism can adapt to increased error rates by increasing gene expression and showed increased selection for more stable proteins (Goldsmith and Tawfik, 2009).

Organisms can take two routes to minimize the synthesis of proteins with altered functionality (Drummond and Wilke, 2009). First, organisms can evolve to minimize the rate at which errors during protein synthesis occur, e.g. selecting for codons that minimize translation error rates (Akashi, 2003; Gilchrist and Wagner, 2006). Second, selection could favor proteins with increased robustness to transcriptional and translational errors, e.g. increase protein stability or increase protein synthesis to compensate for non-functional proteins (Goldsmith and Tawfik, 2009).

In chapter 3, I assumed that the translation process is error free, and that each produced protein functions optimal. Thus, I explicitly ignore any selection on the reduction of translation error rates. While selection for the reduction of translation error rates and selection on ribosome overhead cost do not have to be counteracting forces, they could be for some synonymous codon families. The employed ROC SEMPPR framework (GILCHRIST et al., 2015) yields 100% usage of the most efficient codon if proteins synthesis rate is high enough. While individual genes may reach a 100% codon usage of the most efficient codon, we do not observe populations of high expression genes like that in nature. It is therefore unclear if selection for ribosome overhead cost can overpower counteracting selective forces if protein synthesis rate is high enough.

### 5.2.2 Homogeneous Selection

In ROC SEMPPR and SelAC functionality of a protein refers to the ability of a protein to perform its function and the overall need of an organism for the function. The functionality of a protein depends on many factors (DRUMMOND and WILKE, 2009). As a result, we can approximate the functionality of the protein sequence  $\vec{a}$  in a multitude of ways (GIBBS, 1873; GRANTHAM, 1974; COHEN et al., 2009). However, none can capture the full complexity of a folded protein. It is easy to imagine how the strength, or direction of selection can vary between amino acid site, secondary structures and protein domains within the same protein and certainly between proteins. For example, the functionality of a well-adapted protein is unlikely to be increased by an amino acid substitution. However, the effect on the functionality and in turn fitness of a substitution may drastically differ between active sites and structural sites. Similarly, the exchange of an hydrophilic amino acid with a hydrophobic amino acid is likely to have different effects on the surface of a protein than at the core.

The SelAC framework (Beaulieu et al., in review) employed in chapter 4 assumes that the efficacy of selection follows a gamma-distribution. This distribution is applied to all sites. I, therefore, explicitly do not account for potential differences in the distribution of selection between e.g. secondary structure elements. Similarly, selection for physicochemical properties may differ between sites in e.g. the core or at the surface of a protein. Improvements to SelAC with regards to these shortcomings would allow for new hypothesis to be tested and novel information to be gained from the same data.

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