1 RH: BEAULIEU ET AL.— Pop. Gen. Based Phylo.

- Population Genetics Based Phylogenetics Under Stabilizing Selection for an Optimal Amino Acid Sequence: A Nested Modeling Approach
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We present a novel phylogenetic approach based of substitution rates generated from a nested population genetics model. Unlike many simpler codon models, which assume a 17 single substitution matrix for all sites, our model more realistically represents the evolution 18 of protein-coding DNA under the assumption of consistent, stabilizing selection for a gene 19 specific, optimal amino acid sequence. By modeling the cost-benefit function of an amino 20 acid sequence, our model naturally links the strength of stabilizing selection to protein 21 synthesis levels, which, in turn, can be estimated. Our new set of models, which we collectively call SelAC (Selection on Amino acids and Codons), fit phylogenetic data much 23 better than popular models, suggesting strong potential for more accurate inference of phylogenetic trees and branch lengths from existing data. SelAC also demonstrates that a large amount of biologically meaningful information is accessible when using a nested set of 26 mechanistic models. For example, SelAC prediction of gene specific protein synthesis rates 27 correlates well with both empirical (r = 0.34 - 0.48) and other theoretical predictions 28 (r = 0.59 - 0.64) for multiple species. SelAC also provides estimates of which amino acid is 29 optimal for a given site. Finally, because SelAC is a nested approach based on clearly 30 stated biological assumptions, it can be simplified or expanded as needed, such as including 31 shifts in the optimal amino acid sequence within or across lineages.

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Phylogenetic analyses plays a critical role in most aspects of biology, particularly in
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   the fields of ecology, evolution, paleontology, medicine, and conservation. While the scale
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   and impact of phylogenetic studies has increased substantially over the past two decades,
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   the realism of the mathematical models on which these analyses are based has changed
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   relatively little by comparison. The most popular models of DNA substitution used by
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   molecular phylogenetics are simple nucleotide models that are indifferent to the type of
   sequences to which they are applied. For example, when evaluating protein-coding
   sequences these models are inherently agnostic with regards to the different amino acid
   substitutions and their impact on gene function (e.g. F81, F84, HYK85, TN93, and GTR,
   see Yang (2014) for an overview) and, as a result, cannot describe the behavior of natural
   selection at the amino acid or protein level.
          To address this critical shortcoming, Goldman and Yang (1994) and Muse and Gaut
   (1994) independently introduced models which assumed that differences in the
   physicochemical properties between amino acids, or physicochemical distances for short,
   could affect substitution rates. These physicochemical approaches as originally described
   have rarely been used for empirical data; instead these models have served as the basis for
   an array of simpler and, in turn, more popular models that, starting with Yang and Nielsen
   (1998); Nielsen and Yang (1998), typically assume an equal fixation probability for all
   non-synonymous mutations. Thus, these simpler models initially employed a single term \omega
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   to model the differences in fixation probability between nonsynonomous and synonomyous
   changes at all sites. To improve their realism, more complex forms have been developed
   that allow \omega to vary between sites or branches (as cited in Anisimova 2012) and include
   selection on different synonyms for the same amino acid (e.g. Yang and Nielsen 2008)
          In Goldman and Yang (1994); Yang and Nielsen (1998); Nielsen and Yang (1998)
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   and later studies based on their work, \omega is suggested to indicate whether a given site within
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   a protein sequence is under consistent 'stabilizing (\omega < 1) or 'diversifying' (\omega > 1) selection.
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However, the model's actual behavior is inconsistent with how these terms are typically
   defined and understood (e.g. see Pellmyr 2002). Because synonymous substitutions have a
   higher substitution rate than any possible non-synonymous substitutions when \omega < 1, the
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   model behaves as if the resident amino acid i at a given site is favored by natural selection.
   Even when \omega is allowed to vary between sites, the symmetrical nature of the model means
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   that for any given site the strength of selection for the resident amino acid i over its 19
   alternatives is equally strong regardless of their physicochemical properties. Paradoxically,
   selection for amino acid i persists until a substitution for another amino acid, j, occurs. As
   soon as amino acid j fixes, but not before, selection now favors amino acid j equally over
   all other amino acids, including amino acid i. This is now the opposite scenario from when
   i was the resident. Similarly, when \omega > 1, synonymous substitutions have a lower
   substitution rate than any possible non-synonymous substitutions from the resident amino
   acid. Again due to the model's symmetry, the selection against the resident amino acid i is
   equally strong relative to alternative amino acids. The selection against the resident amino
   acid i persists until a substitution occurs at which point selection now favors amino acid i,
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   as well as the 19 other amino acids, to the same degree i was previously disfavored.
          Thus, the simplest and most consistent interpretation of \omega is that it represents the
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   rate at which the selective environment itself changes, and this change in selection
   perfectly coincides with the fixation of a new amino acid. This, in turn, implies that the
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   rate of shifts in the optimal (or pessimal) amino acid is on the time scale as the rate of
   substitution. Contrary to popular belief, \omega does not describe whether a site is evolving
   under a constant regime of stabilizing or diversifying selection, but instead how a very
   particular selective environment changes over time. Given this behavior, \omega based models
   are likely to only reasonably approximate a subset of scenarios such as perfectly
   symmetrical over-/under-dominance or positive/negative frequency dependent selection
   (Hughes and Nei 1988; Nowak 2006).
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Here we propose an approach where selection is based explicitly on seeking to 85 minimize the cost-benefit function  $\eta$  of a protein where protein function is determined solely 86 by the physicochemical properties of the primary amino acid sequence. Our approach, 87 which we call SelAC (Selection on Amino acids and Codons), is developed in the same vein 88 as previous phylogenetic applications of the Wright-Fisher process (e.g. Muse and Gaut 89 1994; Halpern and Bruno 1998; Yang and Nielsen 2008; Rodrigue et al. 2005; Koshi and Goldstein 1997; Koshi et al. 1999; Dimmic et al. 2000; Thorne et al. 2012; Lartillot and 91 Philippe 2004; Rodrigue and Lartillot 2014). Similar to Lartillot's work (Lartillot and Philippe 2004; Rodrigue and Lartillot 2014), we assume there is a finite set of rate matrices describing the substitution process and that each position within a protein is assigned to a particular rate matrix category. Unlike Lartillot's work, we assume a priori there are 20 different families of rate matrices, one family for when a given amino acid is favored at a site. The key parameters underlying these matrices are shared across genes except for gene expression. As a result, SelAC allows us to quantitatively evaluate the support for a particular amino acid being favored at a particular position within the protein sequence.

# MATERIALS & METHODS

Overview

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We model the substitution process as a classic Wright-Fisher process which includes the forces of mutation, selection, and drift (Fisher 1930; Kimura 1962; Wright 1969; Iwasa 1988; Berg and Lässig 2003; Sella and Hirsh 2005; McCandlish and Stoltzfus 2014). For simplicity, we ignore linkage effects and, as a result of this and other assumptions, sequences evolve in a site independent manner.

JMB: At this point, this seems like natural After this it sort of gives awav the punchline of the model andit reads like discussion. suggest cutting this nextbit moving elsewhere. As of now, the intro is way too long BCO: I agree. We're talking about benefits of the model, but people have no idea I'm vet. moving it to

discussion

Because SelAC requires twenty families of  $61 \times 61$  matrices, the number of 107 parameters needed to implement SelAC would, without further assumptions, be extremely 108 large (i.e. on the order of 74,420 parameters). To reduce the number of parameters needed. 109 while still maintaining a high degree of biological realism, we construct our gene and amino 110 acid specific substitution matrices using a submodel nested within our substitution model, 111 similar to approaches in Gilchrist (2007); Shah and Gilchrist (2011); Gilchrist et al. (2015). 112 One advantage of a nested modeling framework is that it requires only a handful of 113 genome-wide parameters such as nucleotide specific mutation rates (scaled by effective 114 population size  $N_e$ ), amino acid side chain physicochemical weighting parameters, and a 115 shape parameter describing the distribution of site sensitivities. In addition to these 116 genome-wide parameters, SelAC requires a gene g specific expression parameter  $\psi_q$  which 117 describes the average rate at which the protein's functionality is produced by the organism 118 or a gene's 'average functionality production rate' for short (for notational simplicity, we 119 will ignore the gene specific indicator q, unless explicitly needed). Currently,  $\psi$  is fixed 120 across the phylogeny, though relaxing this assumption is a goal of future work. The gene 121 specific parameter  $\psi$  is multiplied by additional model terms to make a composite term  $\psi'$ 122 which scales the strength and efficacy of selection for the optimal amino acid sequence 123 relative to drift (see Implementation below). In terms of the functionality of the protein 124 encoded, we assume that for any given gene there exists an optimal amino acid sequence  $\vec{a}^*$ 125 and that, by definition, a complete, error free peptide consisting of  $\vec{a}^*$  provides one unit of 126 the gene's functionality. We also assume that natural selection favors genotypes that are 127 able to synthesize their proteome more efficiently than their competitors and that each 128 savings of an high energy phosphate bond per unit time leads to a constant proportional gain in fitness  $A_0$ . SelAC also requires the specification (as part of parameter optimization) 130

of an optimal amino acid  $a^*$  at each position within a coding sequence. This requirement of

one  $a^*$  per site makes our  $\vec{a}^*$  the largest category of parameters SelAC estimates. Despite

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the need to specify  $a^*$  for each site, because we use a submodel to derive our substitution matrices, SelAC estimates a relatively small number of the parameters when compared to more general approaches where the fitness of each amino acid is allowed to vary freely of any physicochemical properties (Halpern and Bruno 1998; Lartillot and Philippe 2004; Rodrigue and Lartillot 2014).

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As with other phylogenetic methods, SelAC generates estimates of branch lengths and nucleotide specific mutation rates. In addition, the method can also be used to make quantitative inferences on the optimal amino acid sequence of a given protein as well as the realized average synthesis rate of each protein used in the analysis. The mechanistic basis of SelAC also means it can be easily extended to include more biological realism and test more explicit hypotheses about sequence evolution.

#### Mutation Rate Matrix $\mu$

We begin with a 4x4 nucleotide mutation matrix  $\mu$  that describes mutation rates between different bases and, in turn, different codons. For our purposes, we rely on the general unrestricted model (UNREST from Yang 1994) because it imposes no constraints on the instantaneous rate of change between any pair of nucleotides. More constrained models, such as the Jukes-Cantor (JC), Hasegawa-Kishino-Yano (HKY), or the general time-reversible model (GTR), could also be used.

The 12 parameter UNREST model defines the relative rates of change between a pair of nucleotides. Thus, we arbitrarily set the  $G \rightarrow T$  mutation rate to 1, resulting in 11 free mutation rate parameters in the 4x4 mutation nucleotide mutation matrix. The nucleotide mutation matrix is also scaled by a diagonal matrix  $\pi$  whose entries,  $\pi_{i,i}$ , correspond to the equilibrium frequencies of each base. These equilibrium nucleotide frequencies are determined by analytically solving  $\pi \times \mathbf{Q} = 0$ . We use this  $\mathbf{Q}$  to populate a 61 × 61 codon mutation matrix  $\mu$ , whose entries  $\mu_{i,j}$   $i \neq j$  describes the mutation rate from

codon i to j and  $\mu_{i,i} = -\sum_{j} \mu_{i,j}$ . We generate this matrix using a "weak mutation" assumption, such that evolution is mutation limited, codon substitutions only occur one nucleotide at a time. As a result, the rate of change between any pair of codons that differ by more than one nucleotide is zero.

While the overall model does not assume equilibrium, we still need to scale our mutation matrices  $\mu$  by a scaling factor S. As traditionally done, we rescale our time units such that at equilibrium, one unit of branch length represents one expected mutation per site (which equals the substitution rate under neutrality, but would not with selection). More explicitly,  $S = -\left(\sum_{i \in \text{codons}} \mu_{i,i} \pi_{i,i}\right)$  where the final mutation rate matrix is the original mutation rate matrix multiplied by 1/S.

### Protein Synthesis Cost-Benefit Function $\eta$

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SelAC links fitness to the product of the cost-benefit function of a gene  $\eta$  and the 169 organism's average target synthesis rate of the functionality provided by gene  $\psi$ . This is 170 because the average flux energy an organism spends to meet its target functionality 171 provided by the gene is, by definition,  $\eta \times \psi$ . Compensatory changes that allow an 172 organism to maintain functionality even with loss of one or both copies of a gene are 173 widespread (reviewed in 1); here we assume that for finer scale problems than entire loss 174 (for example, a 10% loss of functionality) the compensation is more production of the 175 protein. In order to link genotype to our cost-benefit function  $\eta = \mathbf{C}/\mathbf{B}$ , we begin by defining our benefit function **B**.

MIKE:
Brian, please
provide
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footnote

<sup>&</sup>lt;sup>1</sup>From Cruft: There is evidence of compensation for protein function. Metabolism with gene expression models (ME-models) link those factors to successfully make predictions about response to perturbations in a cell https://www.nature.com/articles/ncomms1928, https://www.sciencedirect.com/science/article/pii/S0958166914002316. For example, an ME-model for \*E. coli\* successfully predicted gene expression levels in vivo http://journals.plos.org/plosone/article?id=10.1371/journal.pone.0045635.

**Benefit:** Our benefit function **B** measures the functionality of the amino acid sequence  $\vec{a}_i$  encoded by a set of codons  $\vec{c}_i$ , i.e.  $a(\vec{c}_i) = \vec{a}_i$  relative to that of an optimal sequence  $\vec{a}^*$ . By definition,  $\mathbf{B}(\vec{a}^*|\vec{a}^*) = 1$  and  $\mathbf{B}(\vec{a}_i|\vec{a}^*) < 1$  for all other sequences. We assume all amino acids within the sequence contribute to protein function and that this contribution declines as an inverse function of physicochemical distance between each amino acid and the optimal one. Formally, we assume that

$$\mathbf{B}(\vec{a}|\vec{a}^*) = \left(\frac{1}{n}\sum_{p=1}^n \left(1 + G_p d(a_p, a_p^*)\right)^{-1}$$
 (1)

where n is the length of the protein,  $d(a_p, a_p^*)$  is a weighted physicochemical distance 178 between the amino acid encoded at a given position p and  $a_p^*$  is the optimal amino acid for 179 that position. For simplicity, we assume all nonsense mutations are lethal by defining the 180 the physicochemical distance between a stop codon and a sense codon as  $\infty$ . The term  $G_p$ 181 describes the sensitivity of the protein's function to physicochemical deviation from the 182 optimimum at site position p. There are many possible measures for physiochemical 183 distance; we use Grantham (1974) distances by default, though others may be chosen. We 184 assume that  $G_p \sim \text{Gamma}$  (shape  $= \alpha_G$ , rate  $= \alpha_G$ ) in order to ensure  $\mathbb{E}(G_p) = 1$ . Given 185 the definition of the Gamma distribution, the variance in  $G_p$  is equal to 186 shape/rate<sup>2</sup> =  $1/\alpha_G$ . Further, at the limit of  $\alpha_G \to \infty$ , the model becomes equivalent to 187 assuming uniform site sensitivity where  $G_p = 1$  for all positions p. Finally, we note that 188  $\mathbf{B}(\vec{a}_i|\vec{a}^*)$  is inversely proportional to the average physicochemical deviation of an amino 189 acid sequence  $\vec{a}_i$  from the optimal sequence  $\vec{a}^*$  weighted by each site's sensitivity to this 190 deviation.  $\mathbf{B}(\vec{a}_i|\vec{a}^*)$  can be generalized to include second and higher order terms of the 191 distance measure d. 192

Cost: Protein synthesis involves both direct and indirect assembly costs. Direct costs consist of the high energy phosphate bonds  $\sim P$  of ATP or GTP's used to assemble the ribosome on the mRNA, charge tRNA's for elongation, move the ribosome forward along the transcript, and terminate protein synthesis. As a result, direct protein assembly costs are the same for all proteins of the same length. Indirect costs of protein assembly are potentially numerous and could include the cost of amino acid synthesis as well the cost and efficiency with which the protein assembly infrastructure such as ribosomes, aminoacyl-tRNA synthetases, tRNAs, and mRNAs are used. When these indirect costs are combined with sequence specific benefits, the probability of a mutant allele fixing is no longer independent of the rest of the sequence (Gilchrist et al. 2015) and, as a result, model fitting becomes substantially more complex. Thus for simplicity, in this study we ignore indirect costs of protein assembly that vary between genotypes and define,

$$\mathbf{C}(\vec{c_i}) = \text{Energetic cost of protein synthesis.}$$
 (2)

$$=A_1 + A_2 n \tag{3}$$

where,  $A_1$  and  $A_2$  represent the direct cost, in high energy phosphate bonds, of ribosome initiation and peptide elongation, respectively, where  $A_1 = A_2 = 4 \sim P$ .

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### Defining Physicochemical Distances

Assuming that functionality declines with an amino acid  $a_i$ 's physicochemical distance from the optimum amino acid  $a^*$  at each site provides a biologically defensible way of mapping genotype to protein function that requires relatively few free parameters. In addition, SelAC naturally lends itself to model selection since we can compare the quality of SelAC fits using different mixtures of physicochemical properties. Following Grantham (1974), we focus on using composition c, polarity p, and molecular volume v of each amino

acid's side chain residue to define our distance function, but the model and its implementation can flexibly handle a variety of properties. We use the Euclidian distance between residue properties where each property c, p, and v has its own weighting term,  $\alpha_c$ ,  $\alpha_p$ ,  $\alpha_v$ , respectively, which we refer to as 'Grantham weights'. Because physicochemical distance is ultimately weighted by a gene's specific average protein synthesis rate  $\psi$ , another parameter we estimate, there is a problem with parameter identifiablity. The scale of gene expression is affected by how we measure physicochemical distances which, in turn, is determined by our choice of Grantham weights. As a result, by default we set  $\alpha_v = 3.990 \times 10^{-4}$ , the value originally estimated by Grantham, and recognize that our estimates of  $\alpha_c$  and  $\alpha_p$  and  $\psi$  are scaled relative to this choice for  $\alpha_v$ . More specifically,

$$d(a_i, a^*) = (\alpha_c [c(a_i) - c(a^*)]^2 + \alpha_p [p(a_i) - p(a^*)]^2 + \alpha_v [v(a_i) - v(a^*)]^2)^{1/2}.$$

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## Linking Protein Synthesis to Allele Substitution

Next we link the protein synthesis cost-benefit function  $\eta$  of an allele with its fixation probability. First, we assume that each protein encoded within a genome provides some 198 beneficial function and that the organism needs that functionality to be produced at a 199 target average rate  $\psi$ . Again, by definition, the optimal amino acid sequence for a given 200 gene,  $\vec{a}^*$ , produces one unit of functionality, i.e.  $\mathbf{B}(\vec{a}^*) = 1$ . Second, we assume that the 201 actual average rate a protein is synthesized  $\phi$  is regulated by the organism to ensure that 202 functionality is produced at rate  $\psi$ . As a result, it follows that  $\phi = \psi/\mathbf{B}(\vec{a}|\vec{a}^*)$  and the cost 203 of a suboptimal amino acid increases the more it decreases the protein's functionality, **B**. In 204 other words, the average production rate of a protein  $\vec{a}$  with relative functionality  $\mathbf{B}(\vec{a}) < 1$ 205 must be  $1/\mathbf{B}(\vec{a}|\vec{a}^*)$  times higher than the production rate needed if the optimal amino acid 206

sequence  $\vec{a}^*$  was encoded since  $\mathbf{B}(\vec{a}^*|\vec{a}^*) = 1$ . For example, a cell with an allele  $\vec{a}$  where  $\mathbf{B}(\vec{a}|\vec{a}^*) = 9/10$  would have to produce the protein at rate  $\phi = 10/9 \times \psi = 1.11\psi$ . Similarly, a cell with an allele  $\vec{a}$  where  $\mathbf{B}(\vec{a}|\vec{a}^*) = 1/2$  will have to produce the protein at  $\phi = 2\psi$ . In contrast, a cell with the optimal allele  $\vec{a}^*$  would have to produce the protein at rate  $\phi = \psi$ .

Third, we assume that every additional high energy phosphate bond,  $\sim P$ , spent per unit time to meet the organism's target function synthesis rate  $\psi$  leads to a slight and proportional decrease in fitness W. This assumption, in turn, implies

$$W_i(\vec{c}) \propto \exp\left[-A_0 \,\eta(\vec{c}_i)\psi\right]. \tag{4}$$

where  $A_0$ , again, describes the proportional decline in fitness with every  $\sim P$  wasted per unit time. Because  $A_0$  shares the same time units as  $\psi$  and  $\phi$  and only occurs in SelAC in conjunction with  $\psi$ , we do not need to explicitly identify our time units. Instead, we recognize that our estimates of  $\psi$  share an unknown scaling term.

Correspondingly, the ratio of fitness between two genotypes is,

$$W_i/W_j = \exp\left[-A_0 \,\eta(\vec{c}_i)\psi\right] / \exp\left[-A_0 \,\eta(\vec{c}_j)\psi\right] \tag{5}$$

$$= \exp\left[-A_0 \left(\eta(\vec{c}_i) - \eta(\vec{c}_j)\right)\psi\right] \tag{6}$$

(7)

Given our formulations of  $\mathbf{C}$  and  $\mathbf{B}$ , the fitness effects between sites are multiplicative and, therefore, the substitution of an amino acid at one site can be modeled independently of the amino acids at the other sites within the coding sequence. As a result, the fitness ratio

for two genotypes differing at a multiple site simplifies to

$$W_i/W_j = \exp\left[-\left(\frac{A_0 (A_1 + A_2 n_g)}{n_g}\right) \sum_{p \in \mathbb{P}} \left[d(a_{i,p}, a_p^*) - d(a_{j,p}, a_p^*)\right] G_p \psi\right]$$

where  $\mathbb{P}$  represents the codon positions in which  $\vec{c_i}$  and  $\vec{c_j}$  differ. Fourth, we make a weak mutation assumption, such that alleles can differ at only one position at any given time, i.e.  $|\mathbb{P}| = 1$ , and that the population is evolving according to a Wright-Fisher process. As a result, the probability a new mutant, j, introduced via mutation into a resident population i with effective size  $N_e$  will go to fixation is,

$$u_{i,j} = \frac{1 - (W_i/W_j)^b}{1 - (W_i/W_j)^{2N_e}}$$

$$= \frac{1 - \exp\left\{-\frac{A_0}{n_g} (A_1 + A_2 n_g) \left[d(a_i, a^*) - d(a_j, a^*)\right] G_p \psi b\right\}}{1 - \exp\left\{-\frac{A_0}{n_g} (A_1 + A_2 n_g) \left[d(a_i, a^*) - d(a_j, a^*)\right] G_p \psi 2N_e\right\}}$$

where b = 1 for a diploid population and 2 for a haploid population (Kimura 1962; Wright 1969; Iwasa 1988; Berg and Lässig 2003; Sella and Hirsh 2005). Finally, assuming a constant mutation rate between alleles i and j,  $\mu_{i,j}$ , the substitution rate from allele i to j can be modeled as,

$$q_{i,j} = \frac{2}{b}\mu_{i,j}N_e u_{i,j}.$$

where, given the substitution model's weak mutation assumption,  $N_e\mu \ll 1$ . In the end, each optimal amino acid has a separate 64 x 64 substitution rate matrix  $\mathbf{Q}_a$ , which incorporates selection for the amino acid (and the fixation rate matrix this creates) as well as the common mutation parameters across optimal amino acids. This results in the creation of 20  $\mathbf{Q}$  matrices, one for each amino acid and each with 3,721 entries which are
based on a relatively small number of model parameters (one to 11 mutation rates, two free
Grantham weights, the cost of protein assembly,  $A_1$  and  $A_2$ , the gene specific target
functionality synthesis rate  $\psi$ , and optimal amino acid at each position p,  $a_p^*$ ). These model
parameters can either be specified a priori and/or estimated from the data.

Given our assumption of independent evolution among sites, it follows that the probability of the whole data set is the product of the probabilities of observing the data at each individual site. Thus, the likelihood  $\mathcal{L}$  of amino acid a being optimal at a given site position p is calculated as

$$\mathcal{L}\left(\mathbf{Q}_{a}|\mathbf{D}_{p},\mathbf{T}\right) \propto \mathbf{P}\left(\mathbf{D}_{p}|\mathbf{Q}_{a},\mathbf{T}\right)$$
 (8)

In this case, the data,  $\mathbf{D}_p$ , are the observed codon states at position p for the tips of the phylogenetic tree with topology T. For our purposes we take T as given but it could be 225 estimated as well. The pruning algorithm of Felsenstein (1981) is used to calculate 226  $\mathcal{L}(\mathbf{Q}_a|\mathbf{D}_p,\mathbf{T})$ . The log of the likelihood is maximized by estimating the genome scale 227 parameters which consist of 11 mutation parameters which are implicitly scaled by  $2N_e/b$ , 228 and two Grantham distance parameters,  $\alpha_c$  and  $\alpha_p$ , and the sensitivity distribution 229 parameter  $\alpha_G$ . Because  $A_0$  and  $\psi_g$  always co-occur and are scaled by  $N_e$ , for each gene g230 we estimate a composite term  $\psi'_g = \psi_g A_0 b N_e$  and the optimal amino acid for each position 231  $a_p^*$  of the protein. When estimating  $\alpha_G$ , the likelihood then becomes the average likelihood 232 which we calculate using the generalized Laguerre quadrature with k=4 points 233 (Felsenstein 2001). 234 Finally, we note that because we infer the ancestral state of the system, our 235 approach does not rely on any assumptions of model stationarity. Nevertheless, as our 236 branch lengths grow the probability of observing a particular amino acid a at a given site 237

approaches a stationary value proportional to  $W(a)^{2N_e-b}$  and any effects of mutation bias (Sella and Hirsh 2005).

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### Implementation

All methods described above are implemented in the new R package, selac available
through GitHub (https://github.com/bomeara/selac) [it will be uploaded to CRAN
once peer review has completed]. Our package requires as input a set of fasta files that each
contain an alignment of coding sequence for a set of taxa, and the phylogeny depicting the
hypothesized relationships among them. In addition to the SelAC models, we implemented
the GY94 codon model of Goldman and Yang (1994), the FMutSel mutation-selection
model of Yang and Nielsen (2008), and the standard general time-reversible nucleotide
model that allows for Γ distributed rates across sites. These likelihood-based models
represent a sample of the types of popular models often fit to codon data.

For the SelAC models, the starting guess for the optimal amino acid at a site comes 250 from 'majority' rule, where the initial optimum is the most frequently observed amino acid 251 at a given site (ties resolved randomly). Our optimization routine utilizes a four stage hill 252 climbing approach. More specifically, within each stage a block of parameters are 253 optimized while the remaining parameters are held constant. The first stage optimizes the 254 block of branch length parameters. The second stage optimizes the block of gene specific 255 composite parameters  $\psi'_g = A_0 \psi_g N_e b$ . The third stage optimizes SelAC's parameters shared 256 across the genome  $\alpha_c$  and  $\alpha_p$ , and the sensitivity distribution parameter  $\alpha_G$ . The fourth 257 stage estimates the optimal amino acid at each site  $a^*$ . This entire four stage cycle is 258 repeated six more times, using the estimates from the previous cycle as the initial 259 conditions for the new one. However, if there is no improvement in the log-likelihood 260 between any successive optimization cycles, we consider the ML solution found and the 261 search is terminated. For optimization of a given set of parameters, we rely on a bounded 262

MIKE: I thought we updated this to be based on the change in LLik function.

JMB: Is this clear now?

subplex routine (Rowan 1990) in the package NLopt (Johnson 2012) to maximize the log-likelihood function. To help the optimization navigate through local peaks, we perform a set of independent analyses with different sets of naive starting points with respect to the gene specific composite  $\psi'$  parameters,  $\alpha_c$ , and  $\alpha_p$ . Confidence in the parameter estimates can be generated by an 'adaptive search' procedure that we implemented to provide an estimate of the parameter space that is some pre-defined likelihood distance (e.g., 2 lnL units) from the maximum likelihood estimate (MLE), which follows Beaulieu and OMeara (2016) and Edwards (1984).

We note that our current implementation of SelAC is painfully slow, and is best suited for data sets with relatively few number of taxa (i.e. < 10). This limitation is largely due to the size and quantity of matrices we create and manipulate to calculate the log-likelihood of an individual site. Ongoing work will address the need for speed, with the eventual goal of implementing SelAC in popular phylogenetic inference toolkits, such as RevBayes (Hhna et al. 2016), PAML (Yang 2007) and RAxML (Stamatakis 2006).

#### Simulations

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We evaluated the performance of our codon model by simulating datasets and estimating the bias of the inferred model parameters from these data. Our 'known' parameters under 279 a given generating model were based on fitting SelAC to the 106 gene data set and 280 phylogeny of Rokas et al. (2003). The tree used in these analyses is outdated with respect 281 to the current hypothesis of relationships within Saccharomyces, but we rely on it simply as 282 a training set that is separate from our empirical analyses (see section below). Bias in the 283 model parameters were assessed under two generating models: one where we assumed a 284 model of SelAC assuming uniform sensitivity across sites (i.e.  $G_p = 1$  for all sites, 285 i.e.  $\alpha_G = \infty$ ), and one where we used the Gamma distribution joint shape and rate 286 parameter  $\alpha_G$  estimated from the empirical data. Under each of these two scenarios, we 287

used parameter estimates from the corresponding empirical analysis and simulated 50 five-gene data sets. For the gene specific composite parameter  $\psi'_g$  the 'known' values used for the simulation were five evenly spaced points along the rank order of the estimates across the 106 genes. The MLE estimate for a given replicate were taken as the fit with the highest log-likelihood after running five independent analyses with different sets of naive starting points with respect to the composite  $\psi'_g$  parameter,  $\alpha_c$ , and  $\alpha_p$ . All analyses were carried out in our selac R package.

## Analysis of yeast genomes & tests of model adequacy

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We focus our empirical analyses on the large yeast data set and phylogeny of Salichos and 296 Rokas (2013). The yeast genome is an ideal system to examine our phylogenetic estimates 297 of gene expression and its connection to real world measurements of these data within 298 individual taxa. The complete data set of Salichos and Rokas (2013) contain 1070 299 orthologs, where we selected 100 at random for our analyses. We also focus our analyses on 300 Saccharomyces sensu stricto and their sister taxon Candida glabrata, and we used the 301 phylogeny depicted in Fig. 1 of Salichos and Rokas (2013) for our fixed tree. We fit the two 302 SelAC models described above (i.e., SelAC and SelAC+ $\Gamma$ ), as well as two codon models, 303 GY94 and FMutSel, and a standard GTR +  $\Gamma$  nucleotide model. The FMutSel model 304 assumes that the amino acid frequencies are determined by functional requirements of the 305 protein while the other models make no assumptions about amino acid frequencies. In all 306 cases, we assumed that the model was partitioned by gene, but with branch lengths linked 307 across genes. 308

For SelAC, we compared our estimates of  $\phi' = \psi'/\mathbf{B}$ , which represents the average protein synthesis rate of a gene, to estimates of gene expression from empirical data.

Specifically, we obtained gene expression data for five of the six species used - four species were measured during log-growth phase, whereas the other was measured at the beginning

someone this last statement about FMutSel? JMB: This seems right based on what understand about this model

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of the stationary phase (S. kudriavzevii) from the Gene Expression Omnibus (GEO). Gene expression in this context corresponds to mRNA abundances which were measured using either microarrays (C. glabrata, S. castellii, and S. kudriavzevii) or RNA-Seq (S. paradoxus, S. mikatae, and S. cerevisiae).

For further comparison, we also predicted the average protein synthesis rate for each gene  $\phi$  by analyzing gene and genome-wide patterns of synonymous codon usage using ROC-SEMPPR (Gilchrist et al. 2015) for each individual genome. While, like SelAC, ROC-SEMPPR uses codon level information, it does not rely on any inter-specific comparisons and, unlike SelAC, uses only the intra- and inter-genic frequencies of synonymous codon usage as its data. Nevertheless, ROC-SEMPPR predictions of gene expression  $\phi$  correlates strongly (Pearson r = 0.53 - 0.74) with a wide range of laboratory measurements of gene expression (Gilchrist et al. 2015).

While one of our main objectives was to determine the improvement of fit that 325 SelAC has with respect to other standard phylogenetic models, we also evaluated the 326 adequacy of SelAC. Model fit, measured with assessments such as the Akaike Information 327 Criterion (AIC), can tell which model is least bad as an approximation for the data, but it 328 does not reveal whether a model is actually doing a good job of representing the data. An 320 adequate model does the latter, one measure of which is that data generated under the 330 model resemble real data (Goldman 1993). For example, Beaulieu et al. (2013) assessed 331 whether parsimony scores and the size of monomorphic clades of empirical data were 332 within the distributions of simulated data under a new model and the best standard model; 333 if the empirical summaries were outside the range for each, it would have suggested that 334 neither model was adequately modeling this part of the biology.

In order to test adequacy for a given gene we first remove a particular taxon from the data set and the phylogeny. A marginal reconstruction of the likeliest sequence across all remaining nodes is conducted under the model, including the node where the pruned

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taxon attached to the tree. The marginal probabilities of each site are used to sample and assemble the starting coding sequence. This sequence is then evolved along the branch, 340 periodically being sampled and its current functionality assessed. We repeat this process 341 100 times and compare the distribution of trajectories against the observed functionality 342 calculated for the gene. For comparison, we also conducted the same test, by simulating 343 the sequence under the standard  $GTR + \Gamma$  nucleotide model, which is often used on these 344 data but does not account for the fact that the sequences are protein coding, and under 345 FMutSel, which includes selection on codons but in a fundamentally different way as our model. 347

## The appropriate estimator of bias for AIC

As part of the model set described above, we also included a reduced form of each of the

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two SelAC models, SelAC and SelAC+ $\Gamma$ . Specifically, rather than optimizing the amino 350 acid at any given site, we assume the the most frequently observed amino acid at each site 351 is the optimal amino acid  $a^*$ . We refer to these 'majority rule' models as SelAC<sub>M</sub> and 352  $SelAC_M + \Gamma$  and the majority rule parameterization accelerates model fitting. 353 Since these majority rule models assume that the optimal amino acids are known 354 prior to fitting of our model, it is tempting to reduce the count of estimated parameters in 355 the model by the number of parameters estimated using majority rule. Despite having 356 become standard behavior in the field of phylogenetics, this reduction is statistically 357 inappropriate unless one uses an additional dataset for this purpose, something we have 358 not seen. Thus, although using majority rule doesn't necessarily give you the most likely 350 parameter estimate, it still uses the data to generate the estimate and, thus, represents a 360 parameter estimated from the data. Because the difference in the number of parameters K361 when counting or not counting the number of nucleotide sites drops out when comparing 362 nucleotide models with AIC, this statistical issue does not apply to nucleotide models. It 363

does, however, matter for AICc, where K and the sample size n combine in the penalty term. This also matters in our case, where the number of estimated parameters for the majority rule estimation differs based on whether one is looking at codons or single nucleotides.

In phylogenetics two variants of AICc are used. In comparative methods

(e.g. Butler and King 2004; O'Meara et al. 2006; Beaulieu et al. 2013) the number of data

points, n, is taken as the number of taxa. More taxa allow the fitting of more complex

models, given more data. However, in DNA evolution, which is effectively the same as a

discrete character model used in comparative methods, the n is taken as the number of

sites. Obviously, both cannot be correct.

The original derivation of AICc by Hurvich and Tsai (1989) assumed a regression 374 model, where the true model was in the set of examined models, as well as approximations 375 in the derivation itself. The appropriateness of this approximation for phylogenetic data, where shared evolutionary history means data points between taxa lack independence is 377 unclear. In any case, we argue that for phylogenetic data, a good estimate of data set size 378 is number of taxa multiplied by number of sites. First, this is what is conventionally seen 379 as the size of the dataset in the field. Second, when considering how likelihood is 380 calculated, the likelihood for a given site is the sum of the probabilities of each observed 381 state at each tip, which is then multiplied across sites. It is arguable that the conventional 382 approach in comparative methods is calculating AICc in the same way. That is, if only one 383 column of data (or "site") is examined, as remains remarkably common in comparative 384 methods, when we refer to sample size, it is technically the number of taxa multiplied by 385 number of sites, even though it is referred to simply as the number of taxa. One notable exception to this appoach to calculating AICc is the program SURFACE implemented by 387 Ingram and Mahler (2013), which uses multiple characters and taxa. 388

Recently, Jhwueng et al. (2014) performed an analysis that investigated what

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MIKE: This sentence doesn't make any sense to me. JMB: Me neither. But I know what it is trying to say. Modified for clarity.

variant of AIC and AICc worked best as an estimator, but the results were inconclusive.

Here, we have adopted and extended the simulation approach of Jhwueng et al. (2014) in

order to examine a large set of different penalty functions and how well they approximate

the remaining portion of the Kullback-Liebler (KL) divergence between two models after

accounting for the deviance (i.e.,  $-2\mathcal{L}$ ) (see Appendix 1 for more details).

RESULTS

By linking transition rates  $q_{i,j}$  to gene expression  $\psi$ , our approach allows use of the same 396 model for genes under varying degrees of stabilizing selection. Specifically, we assume the 397 strength of stabilizing selection for the optimal sequence,  $\vec{a}^*$ , is proportional to the average 398 protein synthesis rate  $\phi$ , which we can estimate for each gene. In regards to model fit, our 399 results clearly indicated that linking the strength of stabilizing selection for the optimal 400 sequence to gene expression substantially improves our model fit. Further, including the 401 shape parameter  $\alpha_G$  for the random effects term  $G \sim \text{Gamma}(\text{shape} = \alpha_G, \text{rate} = \alpha_G)$  to 402 allow for heterogeneity in this selection between sites within a gene improves the  $\Delta AICc$  of 403 SelAC+ $\Gamma$  over the simpler SelAC models by over 22,000 AIC units. Using either  $\Delta$ AICc or 404  ${\rm AIC_w}$  as our measure of model support, the SelAC models fit extraordinarily better than 405  $GTR + \Gamma$ , GY94, or FMutSel (Table 1). This is in spite of the need for estimating the optimal amino acid at each position in each protein, which accounts for 49,881 additional 407 model parameters. Even when compared to the next most parameter rich codon model in 408 our model set, FMutSel, SelAC+Γ model shows over 160,000 AIC unit improvement over 409 FMutSel. 410

With respect to estimates of  $\phi$  within SelAC, they were strongly correlated with both our empirical measurements (Pearson r=0.34-0.48) and theoretical predictions (Pearson r=0.59-0.64) of gene expression (Figure 1 and Figures S1-S2, respectively). In

JMB: I know details of the AIC analyses in appendix. this but seems like a great place the main highlights here. reader may and just say

other words, using only codon sequences, our model can predict which genes have high or low expression levels. The estimate of the  $\alpha_G$  parameter, which describes the site-specific 415 variation in sensitivity of the protein's functionality, indicated a moderate level of variation 416 in gene expression among sites. Our estimate of  $\alpha_G = 1.36$ , produced a distribution of 417 sensitivity terms G ranged from 0.342-7.32, but with more than 90\% of the weight for a 418 given site-likelihood being contributed by the 0.342 and 1.50 rate categories. In simulation, 419 however, of all the parameters in the model, only  $\alpha_G$  showed a consistent bias, in that the 420 MLE were generally lower than their actual values (see Supporting Materials). Other 421 parameters in the model, such as the Grantham weights, provide an indication as to the 422 physicochemical distance between amino acids. Our estimates of these weights only 423 strongly deviate from Grantham's 1974 original estimates in regards to composition weight, 424  $\alpha_c$ , which is the ratio of noncarbon elements in the end groups to the number of side 425 chains. Our estimate of the composition weighting factor of  $\alpha_c$ =0.459 is 1/4th the value 426 estimate by Grantham which suggests that the substitution process is less sensitive to this 427 physicochemical property when shared ancestry and variation in stabilizing selection are 428 taken into account. 429

It is important to note that the nonsynonymous/synonymous mutation ratio, or  $\omega$ , 430 which we estimated for each gene under the FMutSel model strongly correlated with our 431 estimates of  $\phi' = \psi'/\mathbf{B}$  where **B** depends on the sequence of each taxa. In fact,  $\omega$  showed 432 similar, though slightly reduced correlations, with the same empirical estimates of gene 433 expression described above (Figure 2) This would give the impression that the same 434 conclusions could have been gleaned using a much simpler model, both in terms of the 435 number of parameters and the assumptions made. However, as we discussed earlier, not only is this model greatly restricted in terms of its biological feasibility, SelAC clearly 437 performs better in terms of its fit to the data and biological realism. 438

For example, when we simulated the sequence for S. cervisieae, starting from the

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ancestral sequence under both  $GTR + \Gamma$  and FMutSel, the functionality of the simulated sequence moves away from the observed sequence, whereas SelAC remains near the 441 functionality of the observed sequence (Figure 3b). This is somewhat unsurprising, given 442 that both  $GTR + \Gamma$  and FMutSel are agnostic to the functionality of the gene, but it does 443 highlight the improvement in biological realism in amino acid sequence evolution that 444 SelAC provides. We do note that the adequacy of the SelAC model does vary among 445 individual taxa, and does not always match the observed functionality. For instance, our 446 simulations of S. castellii gene function is consistently higher than estimated from the data 447 (Figure 3c). We suspect this is an indication that assuming a single set of optimal amino 448 acid across all taxa may be too simplistic, but we cannot rule out other potential 449 simplifying assumptions in our model, such as a single set of Grantham weights and  $\alpha_G$ 450 values or the simple, inverse relationship between physicochemical distance d and benefit **B**. Finally, we note that our simulation analysis suggested that the best measure of 452 dataset size for estimating KL distance uses a scaled value of the product of number of 453 sites and number of characters. The model comparison approach described above included 454 this assumption. For more details on the simulation approach, see Appendix 1. 455

# DISCUSSION

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Biologically, we know protein-coding DNA sequences largely evolve through the
introduction of new mutations that either become fixed or lost due to selection and/or
drift. Selection on protein coding regions can take many forms, but can generally be
related to the cost of producing the protein and the functional benefit (or potential harm)
caused by the protein product. The gene specific cost of protein synthesis can be affected
by the amino acids used, the direct and indirect costs of peptide assembly by the ribosome,
the use of chaparones to aid in folding, and even the expected lifespan of the protein.

Importantly, these costs can be computed to varying degrees of realism (e.g. Wagner 2005;
Lynch and Marinov 2015). We have previously presented models of protein synthesis costs
that, alternatively, take into account the cost of ribosome pausing (Shah and Gilchrist
2011) or premature termination errors (Gilchrist and Wagner 2006; Gilchrist 2007;
Gilchrist et al. 2009).

Protein benefit or 'function' can be affected by the amino acids at each site and
their interactions. As a result, amino acid substitutions can affect the functionality at key
catalytic sites or, more broadly, the probability of a particular protein fold and, in turn,
the expected functionality of the protein. Linking amino acid sequence to protein function
is a daunting task; thus for simplicity, we assume that for any given desired biological
function to be carried out by a protein, that (a) the biological importance of this protein
function is invariant across the tree, (b) single optimal amino acid sequence that carries out
this function best, and (c) the functionality of alternative amino acid sequences declines
with their physicochemical distance from the optimum on a site by site basis.

We readily acknowledge that sequence space may have more than one local optimum, that physicochemical distance from the optimal primary amino acid sequence is likely a poor model of protein function, and that the biological importance of a function can vary over time. Nevertheless, we believe our cost-benefit approach to be a substantial advance of the more simplistic  $\omega$  models, is complementary to the work of others in the field (e.g. Thorne et al. 2012; Rodrigue and Lartillot 2014), and, in turn, lays the foundation for more realistic work in the future.

For instance, by assuming there is an optimal amino acid for each site, SelAC
naturally leads to a non-symmetrical and, thus, more cogent model of protein sequence
evolution. Because the strength of selection depends on an additive function of amino acid
physicochemical properties, an amino acid more similar to the optimum has a higher
probability of replacing a more dissimilar amino acid than the converse situation. Further,

SelAC does not assume the system is always at the optimum or pessimum point of the fitness landscape, as occurs when  $\omega < 1$  or > 1, respectively.

Importantly, the cost-benefit approach underlying SelAC allows us to link the
strength of selection on a protein sequence to its gene's expression level. Despite its well
recognized importance in determining the rate of protein evolution (e.g. Drummond et al.
2005, 2006), phylogenetic models have ignored the fact that expression levels vary between
genes. In order to link gene expression and the strength of stabilizing selection on protein
sequences, we simply assume that the strength of selection on a gene is proportional to the
average protein synthesis rate of the gene.

One possible mechanism that generates a linear relationship between the strength of 499 selection and gene expression is the assumption of compensatory gene expression. That is, 500 the assumption that any reduction in protein function is compensated for by an increase in the protein's production rate and, in turn, abundance. For example, a mutation which 502 reduces the functionality of the protein to 90% of the optimal protein, would require 503 1/0.9 = 1.11 of these suboptimal proteins to be produced relative to the optimal protein in 504 order to maintain the same amount of that protein's functionality in the cell. Because the 505 energetic cost of an 11% increase in a protein's synthesis rate is proportional to its target 506 synthesis rate, our assumptions naturally link changes in protein functionality and changes 507 in gene expression and its associated costs. Under what circumstances cells actually 508 respond in this manner, remains to be determined. The fact that our method allows us to 500 explain 13-23% of the variation in gene expression measured using RNA-Seq, suggests that 510 this assumption is a reasonable starting point. More importantly, by linking the strength of 511 stabilizing selection for an optimal amino acid sequence to gene expression, we can effectively weight the phylogenetic information encoded in high and low expression genes 513 which tend to evolve at different rates. 514

Because SelAC infers the optimal amino acid for each site, it is substantially more

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parameter rich than more commonly used models such as GTR+Γ, GY94, and FMutSel.

Despite this increase in number of model parameters, SelAC drastically outperforms these
models with AICc values on the order of 10,000s to 100,000s. We predict that SelAC's
performance could be improved even further if we use a hierarchical approach where the
optimal amino acid is not estimated on a per site basis, but rather as a vector of
probability an amino acid is optimal at the gene level.

SelAC makes inferences about the tree, but also about population genetic 522 parameters, and we can validate the assumptions indirectly by comparing our inferences to 523 other empirical data, such as we do with protein synthesis data. More generally, SelAC's 524 assumptions lead to mechanistic and, thus, testable hypothesis about the relationship 525 between mutation, protein function, gene expression, and rates of evolution. More 526 importantly, alternative hypotheses could be used in place of ours and, in turn, phylogenetic and other types of data could be used to evaluate the support of these alternative models. Our hope is that by moving away from the more phenomenological 529 models we can better connect population genetics, molecular biology, and phylogenetics 530 allowing each area inform the others more effectively. 531

A central goal in evolutionary biology is to quantify the nature, strength, and, 532 ultimately, shifts in the forces of natural selection relative to genetic drift and mutation. 533 As data set size and complexity increase, so does the amount of potential information on 534 these forces and their dynamics. As a result, there is a need for more complex and realistic 535 models (Goldman et al. 1996; Thorne et al. 1996; Goldman et al. 1998; Halpern and Bruno 536 1998; Lartillot and Philippe 2004) to accomplish this goal. Although extremely popular 537 due to their elegance and computational efficiency, the utility of  $\omega$  based models in helping us reach this goal is substantially more limited than commonly recognized. Because these 539  $\omega$  models use a single substitution matrix, they are only applicable for situations in which the substitution process and shifts in the selective environment are intrinsic to the

sequence, such as with positive or negative frequency dependent selection; these models do not describe stabilizing or diversifying selection as commonly envisioned (Endler 1986; Pelmyr 2002).

Starting with Halpern and Bruno (1998), a number of researchers have developed 545 methods for linking site-specific selection on protein sequence and phylogenetics 546 (e.g. Koshi et al. 1999; Dimmic et al. 2000; Koshi and Goldstein 2000; Robinson et al. 547 2003; Lartillot and Philippe 2004; Thorne et al. 2012; Rodrigue and Lartillot 2014) Halpern 548 and Bruno (1998) calculated a vector of 19 expected amino acid frequencies for each amino 549 acid site, making it the most general and most parameter rich of these methods. This 550 generality, however, comes at the cost of being purely descriptive; there is no explicit 551 biological mechanism proposed to explain the site specific amino acid frequencies 552 estimated. By grouping together amino sites with similar evolutionary behaviors, Lartillot and colleagues retained the descriptive nature of Halpern and Bruno (1998) work while 554 greatly reduced the number of model parameters needed (Lartillot and Philippe 2004; 555 Rodrigue and Lartillot 2014). SelAC follows in this tradition of using multiple substitution 556 matrices, but includes some key advances. 557

First, by nesting a model of a sequence's cost-benefit function C/B within a 558 broader model, SelAC allows us to formulate and test a hierarchical, mechanistic models of 550 stabilizing selection. More precisely, our nested approach allows us to relax the assumption 560 that physicochemical deviations from the optimal sequence  $\vec{a}^*$  are equally disruptive at all 561 sites within a protein. We found strong support for SelAC's hypothesis that the strength of stabilizing selection against physicochemical deviations from  $\vec{a}^*$  varies between sites  $(\Delta AICc = 20.983)$ . Second, because our substitution matrices are built on a formal description of a sequence's cost-benefit function C/B, we are able to efficiently 565 parameterize 20 different matrices using a relatively small number of genome-wide 566 parameters – e.g. our physicochemical weighting and G distribution shape parameters, and 567

one gene specific gene expression parameter  $\psi$ . While the C/B function on which SelAC currently rests is very simple, nevertheless, it leads to a dramatic increase in our ability to 569 explain the sequence data we analyzed. Importantly, because SelAC uses a formal 570 description of a sequence's C/B, replacing our assumptions with more sophisticated ones 571 in the future is relatively straightforward. Conceptually, our work lies in between that of 572 Lartillot's and Thorne's, where the latter is utilizing even more detailed models of protein 573 structure as a means of linking amino acid substitutions and stabilizing selection. Third, 574 our use of nested models also allows us to make biologically meaningful and testable 575 predictions. By linking a gene's expression level to the strength of purifying selection it 576 experiences, we are able to provide coarse estimates of gene expression. This also suggests 577 that  $\omega$  is best explained as a proxy for gene expression, rather than the nature of selection 578 on a sequence.

One simplifying assumption we make is that the organism can and does compensate 580 for any reduction in protein function by simply increasing the protein's production rate. 581 While this production compensation assumption will clearly not hold in many situations, it 582 does allow us to connect protein function and energetic costs in a simple and biologically 583 plausible manner. Of course, researchers could employ and test other assumptions within 584 our framework, namely, by utilizing more detailed, gene specific knowledge about the 585 relationship between protein function and organism fitness. For example, suppose a protein 586 for a glucose transporter is far less efficient than usual. One possible response and the one 587 envisioned here is that the protein is thus produced at a higher rate to compensate. This 588 would leave the overall ability to transport glucose unchanged. An alternative is that the cell is just less able to transport glucose across membranes. In biology, it is likely that a mixture of such effects exists. However, the production compensation mechanism is likely 591 to have the same costs across proteins, making it a useful first approximation, while the 592 same expression but reduced functionality will have gene specific effects more difficult to 593

model generally (e.g., how does the cost of having glucose transport slow by half compare to the cost of underproducing an anthocyanin for flower color or fewer taste receptor proteins?). Moreover, there is evidence that cells do compensate for lower protein function by increasing gene expression. Nevertheless, by assuming that fitness declines with extraneous energy flux, SelAC explicitly links the variation in the strength of stabilizing selection for the optimal protein sequence among genes, to the variation among genes in their target expression levels  $\psi$ .

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Furthermore, by linking expression and selection, SelAC provides a natural 601 framework for combining information from protein coding genes with very different rates of 602 evolution; from low expression genes providing information on shallow branches to high 603 expression genes providing information on deep branches. This is in contrast to a more 604 traditional approach of concatenating gene sequences together, which is equivalent to assuming the same average protein synthesis rate  $\psi$  for all of the genes, or more recent 606 approaches where different models are fitted to different genes. Our results indicate that 607 including a gene specific  $\psi$  value vastly improves SelAC fits (Table 1). Perhaps more 608 convincingly, we find that the target expression level  $\psi$  and realized average protein 609 synthesis rate  $\phi$  are reasonably well correlated with laboratory measurements and 610 theoretical predictions of gene expression (Pearson r = 0.34 - 0.64; Figures 1, S1, and S2). 611 The idea that quantitative information on gene expression is embedded within 612 intra-genomic patterns of synonymous codon usage is well accepted; our work shows that 613 this information can also be extracted from comparative data at the amino acid level. 614 Of course, given the general nature of SelAC and the complexity of biological 615 systems, other biological forces besides selection for reducing energy flux likely contribute to intergenic variation in the magnitude of stabilizing selection. Similarly, other 617 physicochemical properties besides composition, volume, and charge likely contribute to 618

site specific patterns of amino acid substitution. Thus, a larger and more informative set of

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physicochemical weights might improve our model fit and reduce the noise in our estimates of  $\phi$ . Even if other physicochemical properties are considered, the idea of a consistent, genome wide physicochemical weighting of these terms seems highly unlikely. Since the importance of an amino acid's physicochemical properties likely changes with its position in a folded protein, one way to incorporate such effects is to test whether the data supports multiple sets of physicochemical weights for either subsets of genes or regions within genes, rather than a single set.

Both of these points highlight the advantage of the detailed, mechanistic modeling 627 approach underlying SelAC. Because there is a clear link between protein expression, 628 synthesis cost, and functionality, SelAC can be extended by increasing the realism of the 629 mapping between these terms and the coding sequences being analyzed. For example, 630 SelAC currently assumes the optimal amino acid for any site is fixed along all branches. This assumption can be relaxed by allowing the optimal amino acid to change during the 632 course of evolution along a branch. From a computational standpoint, the additive nature 633 of selection between sites is desirable because it allows us to analyze sites within a gene 634 largely independently of each other. From a biological standpoint, this additivity between 635 sites ignores any non-linear interactions between sites, such as epistasis, or between alleles, 636 such as dominance. Thus, our work can be considered a first step to modeling these more 637 complex scenarios. 638

For example, our current implementation ignores any selection on synonymous

codon usage bias (CUB) (c.f. Yang and Nielsen 2008; Pouyet et al. 2016). Including such

selection is tricky because introducing the site-specific cost effects of CUB, which is

consistent with the hypothesis that codon usage affects the efficiency of protein assembly or

C, into a model where amino acids affect protein function or B, results in a cost-benefit

ratio C/B with epistatic interactions between all sites. These epistatic effects can likely be

ignored under certain conditions or reasonably approximated based on an expectation of

codon specific costs (e.g. Kubatko et al. 2016). Nevertheless, it is difficult to see how one could identify such conditions without modeling the way in which codon and amino acid usage affects C/B.

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This work also points out the potential importance of further investigation into 649 model choice in phylogenetics. For likelihood models, use of AICc has become standard. 650 However, how one determines the appropriate number of parameters estimated in a model 651 is more complicated than generally recognized. Common sense suggests that dataset size is 652 increased by adding taxa and/or sites. In other words, a dataset of 1000 taxa and 100 sites 653 must have more information on substitution models than a dataset of 4 taxa and 100 sites. 654 Our simple analyses agree that the number of observations in a dataset (number of sites  $\times$ 655 number of taxa) should be taken as the sample size for AICc, but this conclusion likely 656 only applies when there is sufficient independence between taxa. For instance, one could imagine a phylogeny where one taxon is sister to a polytomy of 99 taxa that have zero 658 length terminal branches. Absent measurement error or other intraspecific variation, one 659 would have 100 species but only two unique trait values, and the only information about 660 the process of evolution comes from what happens on the path connecting the lone taxon 661 to the polytomy. Although this is a rather extreme example, it seems prudent for 662 researchers to use a simulation based approach similar to the one we take here to determine 663 the appropriate means for calculating the effective number of data points in their data. 664

There are still significant shortcomings in the approach outlined here. Most worrisome are biological oversimplifications in SelAC. For example, at its heart, SelAC assumes that suboptimal proteins can be compensated for, at a cost, simply by producing more of them. However, this is likely only true for proteins reasonably close to the optimal sequence. Different enough proteins will fail to function entirely: the active site will not sufficiently match its substrates, a protein will not properly pass through a membrane, and so forth. Yet, in our model, even random sequences still permit survival, just requiring more

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protein production. Other oversimplifications include the assumption of no selection on
    codon usage, no change of optimal amino acids through time, and no change of the effect of
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   physiochemical properties on fitness through time. However, because we take a mechanistic
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    approach, all of these assumptions can be relaxed through further extension of our model.
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           There are also deficiencies in our implementation. Though reasonable to use for a
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   given topology with a modest number of species, it is currently too slow for practical use
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   for tree search. Our work serves as a proof of concept, or of utility for targeted questions
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    where a more realistic model may be of use (placement of particular taxa, for example).
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    Future work will encode SelAC models into a variety of mature, popular tree-search
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   programs. SelAC also represents a challenging optimization problem: the nested models
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   reduce parameter complexity vastly, but there are still numerous parameters to optimize,
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   including the discrete parameter of the optimal amino acid at each site. A different
   implementation, more parameter-rich, would optimize values of three (or more)
    physiochemical properties per site. This would have the practical advantage of continuous
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    parameter optimization rather than discrete, and biologically would be more realistic (as it
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   is the properties that selection "sees", not the identity of the amino acid itself).
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           In spite of these difficulties, SelAC represents an important step in uniting
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    phylogenetic and population genetic models. While Koshi et al. (1999); Dimmic et al.
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    (2000); Koshi and Goldstein (2000); Robinson et al. (2003); Lartillot and Philippe (2004);
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    Thorne et al. (2012); Rodrigue and Lartillot (2014) are all models of constant, stabilizing
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   selection, SelAC can be generalized further to include diversifying selection. Specifically, by
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   letting SelAC's Grantham weighting term G, which we now assume is \geq 0, to take on
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   negative values, SelAC will behave as if there is a pessimal, rather than optimal, amino
   acid for the given site. In this diversifying selection scenario, amino acids with
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   physicochemical qualities more dissimilar to the pessimal amino acid are increasingly
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   favored, potentially resulting in multiple fitness peaks.
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This ability to extend our model and, in turn, sharpen our thinking about the 698 nature of natural selection on amino acid sequences illustrates the value of moving from 699 descriptive to more mechanistic models in general and phylogenetics in particular. How 700 frequently diversifying selection of this nature occurs is an open, but addressable, question. 701 Regardless of the frequency at which diversifying selection occurs, it leads to the question, 702 "How often does the optimal/pessimal amino sequence change along any given branch?" 703 Due to its mechanistic nature, SelAC can also be extended to include changes in the 704 optimal/pessimal sequence over a phylogeny using a hidden markov modelling approach. 705 Extending SelAC in these ways, will allow researchers to explicitly model shifts in selection 706 on protein sequences and, in turn, quantify their frequency and magnitude. 707

In summary, SelAC allows biologically relevant population genetic parameters to be 708 estimated from phylogenetic information, while also dramatically improving fit and accuracy of phylogenetic models. By explicitly modeling the optimal/pessimal sequence of 710 a gene, SelAC can be extended to include shifts in the optimal/pessimal sequence over 711 evolutionary time. Extending this model in this way will allow researchers to describe not 712 only the dynamic shifts in natural selection, but evaluate how well a given dataset supports 713 such a model. Moreover, it demonstrates that there remains substantially more information 714 in the coding sequences used for phylogenetic analysis than other methods can access. 715 Given the enormous amount of efforts expended to generate sequence datasets, it makes 716 sense for researchers to continue developing more realistic models of sequence evolution in 717 order to extract the biological information embedded in these datasets. The cost-benefit 718 model we develop here is just one of many possible paths of mechanistic model 719 development. 720

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 $_{887}$  Table

		Parameters				Model
Model	logLik	Estimated	AIC	AICc	$\Delta {\rm AICc}$	Weight
$GTR+\Gamma$	-655,166.4	610	1,311,553	1,311,554	284,240	< 0.001
GY94	-612,670.4	111	1,225,563	$1,\!225,\!563$	198,249	< 0.001
FMutSel	-597,140.7	178	1,194,637	1,194,638	167,324	< 0.001
$\mathrm{SelAC}_M$	-478,302.4	50,004	1,056,613	1,076,674	49,360	< 0.001
SelAC	-464,114.8	50,004	1,028,238	1,048,299	20,985	< 0.001
$SelAC_M + \Gamma$	-465,106.9	50,005	1,030,189	1,050,286	22,972	< 0.001
$SelAC+\Gamma$	-453,620.8	50,005	1,007,252	1,027,314	0	> 0.999

Table 1: Comparison of model fits using AIC, AICc, and AICw. Note the subscripts M indicate model fits where the most common or 'majority rule' amino acid was fixed as the optimal amino acid  $a^*$  for each site. As discussed in text, despite the fact that  $a^*$  for each site was not fitted by our algorithm, its value was determined by examining the data and, as a result, represent an additional parameter estimated from the data and are accounted for in our table. Also, the sample size used in the calculation of AICc is assumed to be equal to the size of the matrix (number of taxa x number of sites).

\*\*\* FIGURES

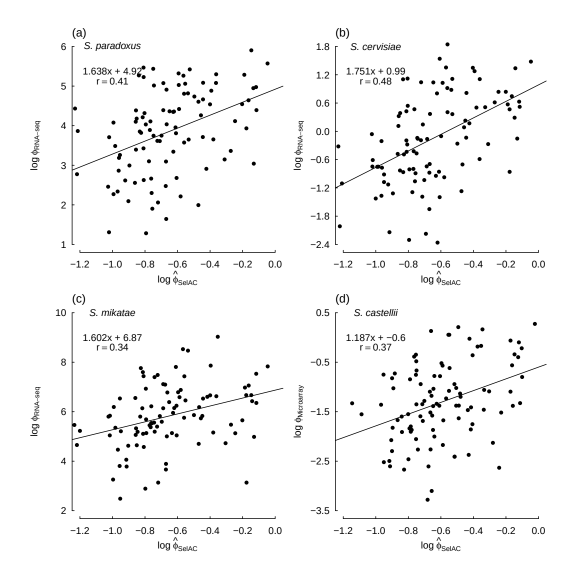


Figure 1: Comparisons between estimates of average protein translation rate  $\hat{\phi}_{SelAC}$  obtained from SelAC+ $\Gamma$  and direct measurements of expression for individual yeast taxa across the 100 selected genes from Salichos and Rokas (2013). Estimates of  $\hat{\phi}_{SelAC}$  were generated by dividing the composite term  $\psi'$  by  $\mathbf{B}(\vec{a}_i|\vec{a}^*)$ . Gene expression was measured using either RNA-Seq (a)-(c) or microarray (d). The equations in the upper left hand corner of each panel represent the regression fit and the Pearson correlation coefficient r.

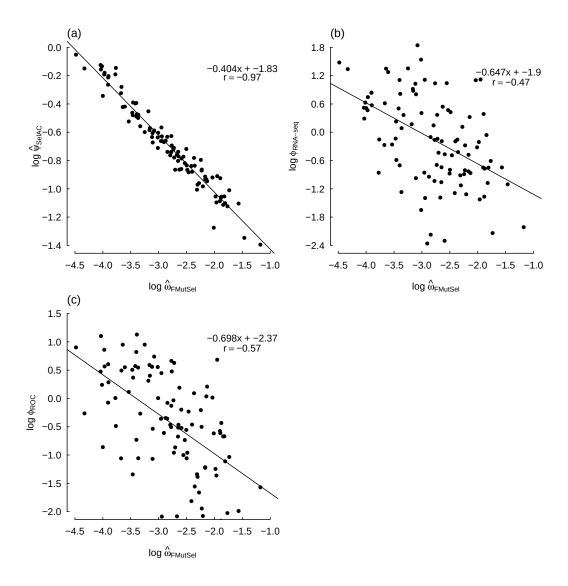


Figure 2: Comparisons between  $\omega_{\rm FMutSel}$ , which is the nonsynonymous/synonymous mutation ratio in FMutSel, SelAC+ $\Gamma$  estimates of protein functionality production rates  $\hat{\psi}_{\rm SelAC}$  (a), RNA-Seq based measurements of mRNA abundance  $\phi_{\rm RNA-seq}$  (b), and ROC-SEMPPER's estimates of protein translation rates  $\phi_{\rm ROC}$ , which are based solely on *S. cerevisiae*'s patterns of codon usage bias (c), for *S. cerevisiae* across the 100 selected genes from Salichos and Rokas (2013). As in Figure 1, the equations in the upper right hand corner of each panel provide the regression fit and correlation coefficient.

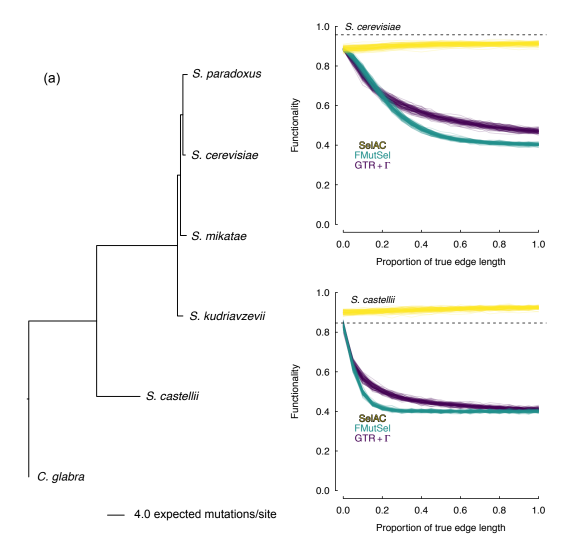


Figure 3: (a) Maximum likelihood estimates of branch lengths under SelAC+ $\Gamma$  for 100 selected genes from Salichos and Rokas (2013). Tests of model adequacy for *S. cerevisiae* (b) and *S. castellii* (c) indicated that, when these taxa are removed from the tree, and their sequences are simulated, the parameters of SelAC+ $\Gamma$  exhibit functionality  $\mathbf{B}(\vec{a}_{\text{obs}}|\vec{a}^*)$  that is far closer to the observed (dashed black line) than data sets produced from parameters of either FMutSel or GTR +  $\Gamma$ .

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Supporting Materials for Population Genetics Based Phylogenetics Under Stabilizing

Selection for an Optimal Amino Acid Sequence: A Nested Modeling Approach by Beaulieu

et al. (In Review).

## Comparisons of SelAC gene expression estimates with empirical measurements

In our model, the parameter  $\phi$  measures the realized average protein synthesis rate of a gene. We compared our estimates of  $\phi$  to two separate measures of gene expression, one empirical (Figure S1), and one model-based prediction that does not account for shared ancestry, for individual yeast taxa across the same set of genes. Our estimates of  $\phi$  are positively correlated with both measures, which are also reasonably well correlated with each other (Figure 1 - S2) On the whole, these comparisons indicate not only a high degree of consistency among all three measures, but also, importantly, that estimates of  $\phi$  obtained from SelAC provide real biological insight into the expression level of a gene.

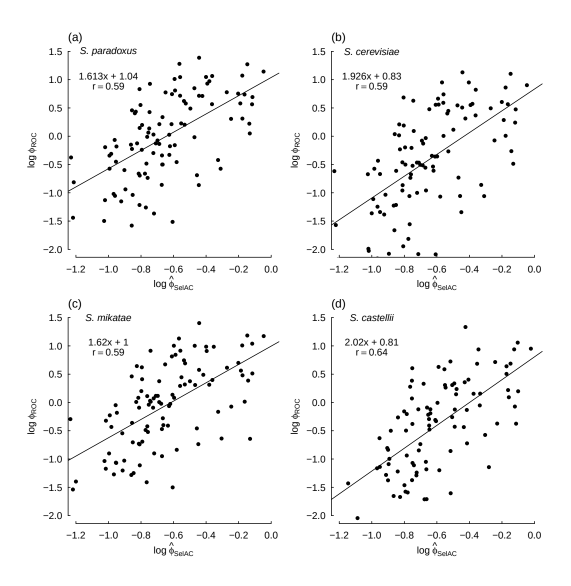


Figure S1: Comparisons between estimates of  $\phi$  obtained from SelAC+ $\Gamma$  and the predicted gene expression from the ROC SEMPER model (Gilchrist et al. (2015)) for individual yeast taxa across the 100 selected genes from Salichos and Rokas (2013). As with figures in the main text, estimates of  $\phi$  were obtained by solving for  $\psi$  based on estimates of  $\psi'$ , and then dividing by  $\mathbf{B}(\vec{a}_i|\vec{a}^*)$ . The equations in the upper left hand corner of each panel represent the regression fit and correlation coefficient.

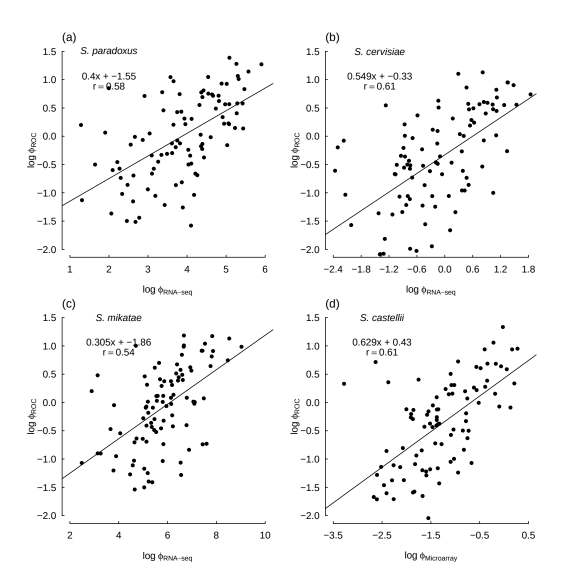


Figure S2: Comparisons of predicted gene expression from the ROC SEMPER model (Gilchrist et al. (2015)) and direct measurements of expression from RNA-Seq or microarray data for individual yeast taxa across the 100 selected genes from Salichos and Rokas (2013). The equations in the upper left hand corner of each panel represent the regression fit and correlation coefficient.

Simulations

Overall, the simulation results indicate that the SelAC model can reasonably recover the

known values of the generating model (Figure S3 - S6). This includes not only the

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parameters in SelAC, but also the optimal amino acids for a given sequence as well as the estimates of the branch lengths. There are a few observations to note. First, the ability to 907 accurately recover the true optimal amino acid sequence will largely depend on the 908 magnitude of the realized average protein synthesis rate of the gene  $\phi$ . This is, of course, 909 intuitive, given that  $\phi$  sets the strength of stabilizing selection towards an optimal amino 910 acid at a site. However, the inclusion of  $\alpha_G$  into SelAC, appears to generally increase 911 values of  $\phi$  and generally improves the ability to recover the optimal amino acids even for 912 the gene with the lowest baseline  $\phi$ . Second, we found a strong downward bias in estimates 913 of  $\alpha_G$ , which actually translates to greater variation among the rate categories. The choice 914 of a gamma distribution to represent site-specific variation in sensitivity was based on 915 mathematical convenience and convention, rather than on biological reality. Nevertheless, 916 we suspect that this bias is in large part due to the difficulty in determining the baseline  $\psi$ 917 for a given gene and the value of  $\alpha_G$  that globally satisfies the site-specific variation in 918 sensitivity across all genes, as indicated by the slight upward bias in estimates of  $\psi$ . A 919 reviewer pointed out that it may also be difficulty for SelAC to account for changing 920 amino-acid, which we agree may also play a role. It has been suggested, in studies of the 921 behavior of the gamma distribution in applications of nucleotide substitution model, that 922 increasing the number of rate categories can often improve accuracy of the shape 923 parameter (Mayrose et al. (2005)). Future work will address this issue. 924



Figure S3: Summary of a 5-gene simulation for a SelAC model where we assume  $\alpha_G = \infty$ , and thus, no site-specific sensitivity in the generating model. The 'known' parameters were based on fitting the same SelAC to the 106 gene data set and phylogeny of Rokas et al. (2003), with gene choice being based on five evenly spaced points along the rank order of the gene specific composite parameter  $\psi'_g$ . The points and associated uncertainty in the estimates of the gene-specific average protein synthesis rate, or  $\psi$  (calculated from  $\psi'$ )(a), nucleotide mutation rates under the UNREST model (b), proportion of correct optimal amino acids for a given gene (c), and estimates of the individual edge lengths are based the mean and 2.5% and 97.5% quantiles across all 50 simulated datasets (d). Gene index on the x-axis refers to the arbitrary number assigned to the simulated gene.



Figure S4: The distribution of estimates of the Grantham weights,  $\alpha_c$  and  $\alpha_p$ , in a SelAC model, where we assume  $\alpha_G = \infty$ , and thus no site-specific sensitivity in the generating model. The dashed line represents the value used in the generating model.



Figure S5: Same figure as in Figure S3, except the generating model includes site-specific sensitivity in the generating model (i.e.,  $\alpha_G$ ).

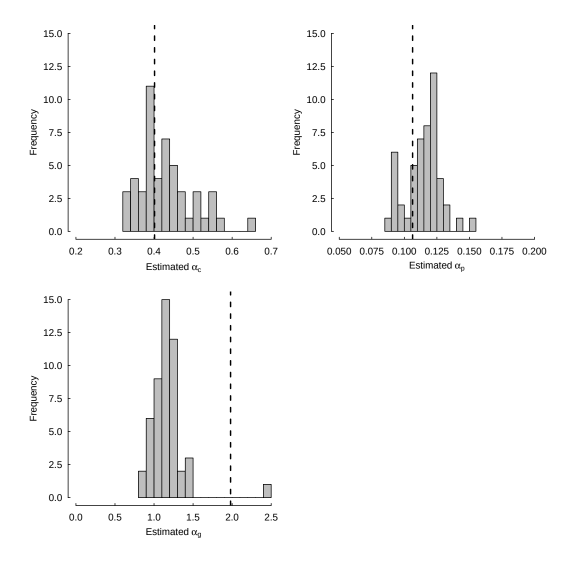


Figure S6: Same figure as in Figure S4, except the generating model includes site-specific sensitivity in the generating model (i.e.,  $\alpha_G$ ). Unlike, Grantham weights, which showed no systematic bias, there is a downward bias in estimates of  $\alpha_G$ .