1 RH: LANDERER ET AL.— Intragenomic variation in codon usage

Decomposing mutation and selection to identify mismatched codon usage

- 4 CEDRIC LANDERER^{1,2,*}, RUSSELL ZARETZKI³, AND MICHAEL
- 5 A. GILCHRIST^{1,2}

- ⁶ Department of Ecology & Evolutionary Biology, University of Tennessee, Knoxville, TN 37996-
- 7 1610
- 8 $^{2}\mathrm{National}$ Institute for Mathematical and Biological Synthesis, Knoxville, TN 37996-3410
- ³Department of Business Analytics & Statistics, Knoxville, TN 37996-0532
- *Corresponding author. E-mail: cedric.landerer@gmail.com

Version dated: September 19, 2018

Abstract

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

Here we examine variation in codon usage patterns of endogenous and exogenous genes in the yeast Lachancea kluyveri. Previous studies indicate that the left arm of chromosome C, or $\sim 10\%$ of the L. kluyveri genome, is the result of an large introgression of exogenous genes. Thus, the L. kluyveri genome provides an opportunity to study the adaptation of these exogenous to a novel cellular environment and estimate how the genetic load of these genes changes over time. In order to quantiatively describe L. kluyveri's codon usage environment, we fitted a bayesian, mechanistic model of codon usage bias evolution, ROC-SEMPPR, to L. kluyveri's endogenous gene in order to estimate the strength of mutation bias and selection on codon usage. We then compared these parameter estimates to those we obtained by fitting ROC-SEMPPR to L. kluyveri's exogenous genes, which provides a biased estimate of the ancestral environment of the exogenous genes. Our results indicate the differences in codon usage between L. kluyveri's endogenous and exogenous genes are largely due to differences in mutation bias, rather than selection. Estimating mutation and selection parameters separately for the endogenous and exogenous genes improved our ability to predict empirical estimates of protein synthesis by 17% and avoided errors in identifying L. kluyveri's selectively favored or 'optimal' codons. By comparing our mutation and selection parameters to those estimated for other yeast species, we identified Eremothecium qossypii as the most likely source of L. kluyveri's exogenous genes. Using these parameters and available estimates of mutation rates in yeast, we estimated the age of the introgression to be on the order of 6×10^8 generation. Finally, we estimated the genetic load of the exogeneous genes both at the time of introgression and currently. In summary, our work shows the advantage of using mechanistic models that separate the effects of selection and mutation on codon usage.

36 Introduction

Synonymous codon usage patterns often varies within a genome and between taxa and reflects the effects of 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
complement DNA repair genes. The signature of selection on codon usage is also largely
determined by an organism's cellular environment, such as its translational machinery. In
contrast, the strength of selection on the codon usage of an individual gene is largely determined by its expression level which, in turn, is largely determined by the organism's external
environment. The efficacy of this selection on codon usage relative to drift is a function of
the organism's effective population size N_e which, in turn, is largely determined by its external environment. Thus, the codon usage patterns of a genome provides us with a unique
signature of the organism's internal by decomposing the patterns codon usage bias across a
set of genes allows us to genome provides us with information about the organism's cellular
and external environment.

In general, the strength of selection on codon usage increases with gene expression [Ikemura, 1985, Gouy and Gautier, 1982]. Conversely, the impact of mutation bias on codon
usage declines with gene expression. Thus, we can easily imagine that with increasing gene
expression, codon usage shifts from a process dominated mutation to a process dominated
by selection. Together, the mutation process favoring specific synonymous codons - or mutation bias - and the selection for translation efficiency scaled by gene expression and effective
population size - or selection bias - shape codon usage in a genome. This mutation-selectiondrift balance model allows us to explicitly describe the environment in which genes evolve
with respect to mutation and selection bias. Here we show that estimating the influence
of mutation bias and selection bias on a gene's codon usage allows us to not only predict
protein synthesis rate ϕ , but also, to infer its history and make predictions about its future
with respect to these forces.

Most studies implicitly assume that synonymous codon usage of a genome is reflects

the single mutational and selective cellular environment of the organism. However, any introgressed genes, whether the result of hybridization or horizontal gene transfer, should carry the signature of the exogenous cellular environment whence they came and, in turn, impose a genetic load on the recipient lineage. The magnitude of the exogenous genes' genetic load on the recipient or endogenous lineage should increase as the mutation and selective environments differ between the donor and recipient lineages as well as with the expression level of the genes in the recipient cells. Thus codon usage patterns likely play a critical role in the rates of introgression between lineages and, as a result, can serve as an important source of information about such events.

Here we analyze the synonymous codon usage of the genome of Lachancea kluyveri, the earliest diverging lineage of the Lachancea clade. The Lachancea clade diverged from the Saccharomyces clade prior to the whole genome duplication, about 100 Mya ago. Since its divergence from the other Lachancea, L. kluyveri has experienced a large introgression of exogenous genes. These introgressed exogenous genes replaced XXXX genes, constituting the entire left arm of L. kluyveri's C chromosome, and were identified due to their $\sim 13\%$ elevation in GC content bias relative to the remaining, endogenous genes of the L. kluyveri genome [Payen et al., 2009, Friedrich et al., 2015].

In order to study the effects of introgression and the resulting mismatches in codon usage
in the *L. kluyveri* genome, we use ROC SEMPPR, a mechanistic model of codon usage
bias evolution grounded in population genetics. ROC SEMPPR, which uses a bayesian
MCMC method for model fitting, allows us to quantify the contributions of mutation bias
and selection on to the codon usage patterns of a set of genes. ROC SEMPPR also allows
us to predict a gene's average predicting protein production rate based on its individual
codon usage pattern with a precision comprable to that of more direct empirical methods
[Gilchrist et al., 2015]. By fitting ROC SEMPPR separately to *L. kluyveri*'s endogenous and
exogenous sets of genes we generate a quantiative 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 mostly to differences in mutation bias. In addition, we show that separately fitting ROC SEMPPR to endogenous and exogenous gene sets substantially improves our ability to predict empirical estimates of protein synthesis rates over fitting ROC to a combined dataset of endogenous and exogenous genes.

In order to identify a potential source lineage for L. kluyveri's exogenous gene set we 95 fit ROC SEMPPR to the genomes of 38 yeastspecies. We then compared ROCs parameter estimates of mutation bias and selection of L. kluyveri's exogenous genes to these species 97 and found a strong correlation in only two species, E. qossypii and C. dubliniensis. We also compared the synteny of L. kluyveri's exogenous genes to these lineages. We found strong synteny in a number of cases, most notably in E. qossypii but not C. dubliniensis. 100 As a result, we conclude that of the yeast species we examined, the E. qossypii lineage is 101 most closely related to the donor of L. kluyveri's exogenous genes. Assuming that E. 102 qossypii's mutation bias is similar to the source of the exogenous genes, we estimated the 103 introgression occurred approximately 6×10^8 generations ago using a model of exponential 104 decay to describe the shift in mutation bias of the exogenous genes. Finally, we estimate 105 the selective cost or genetic load of the exogenous genes due to codon usage mismatch using our estimates of the selection parameters from L. kluyveri's endogenous genes and the our estimates of the protein synthesis rate of the exongenous genes. 108

Need to discuss introgression

• Load calculations

109

110

111

114

- Some genes pre-adapted to new environment
- Most genes not
- Load estimate indicates strong selection against introgression sequences alone

• Explaining introgression

- Assuming introduction is continuous as it appears, indicates little recombination during spread
 - Data suggests introgression spread quickly
 - Potential explanations

- * Identified wrong source, though even current load is quite large.
- * Major flaws in our calculation of fitness costs.
- * Failure for positive selection on at amino acid or regulatory sequence at one or more loci, countering the selection on CU
- * Introgression triggered speciation event, thus N_e was very small (< 100) so even if strongly selected against it still had a reasonable probability of fixing.
- * Unlikely event, but introgressions happen frequently. Note here mutation is actually an introgression event, not a nt change. Although pop gen predicts fixation probability is very low. However, pop gen also tells us that if such an unlikely fixation occurs, it is very likely to happen quickly. Thus, continuous nature of introgression also consistent with a rare, maladaptive fixation event.
- * Other adaptive effects of introgression seems most plausible, but since we don't know have reasonable estimates about frequently hybridiations occur nor accurate estimate of how frequent such introgressions fix, the maladaptive explanation is hard to evaluate.
- * Combination of most maladaptive, some adaptive alleles, and speciation could also be a feasible hypothesis.

Terminiology

- Codon families?

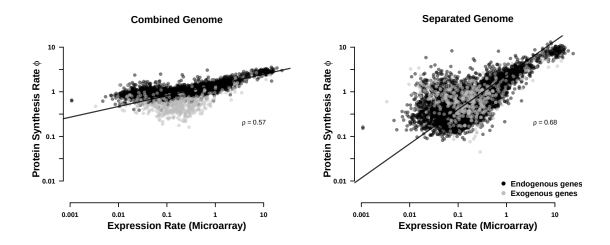


Figure 1: Comparison of predicted protein synthesis rate ϕ to Microarray data from XX for (a) the combined genome and (b) the separated endogenous and exogenous genes. Endogenous genes are displayed in black and exogenous genes in red. Black line indicates type II regression line.

38 Results

$^{_{39}}$ $L.\ kluyveri$'s Genome Contains Signatures from Two Cellular En- $^{_{40}}$ vironments

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 separated into two sets of 4,864 endogenous and 497 exogenous genes. AIC values (Δ AIC = 75,462; Table 1) strongly support the hypothesis that the L.~kluyveri genome consists of genes with two different and distinct patterns of codon usage bias. We found additional support for this hypothesis when we compared our predictions of gene expression to empirically observed values. Specifically, the correlation between our predictions and observed values improved by almost 20%, from $\rho = 0.57$ to 0.68 (Figure 1).

49 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 ΔM and selection $\Delta \eta$ for the two 151 sets of genes. Our estimates of ΔM for the endogenous and exogenous genes were negatively 152 correlated ($\rho = -0.49$), indicating weak concordance of $\sim 5\%$ between the two mutation 153 environments (Figure 2). For example, the endogenous genes show a mutational preference 154 for A and T ending codons in $\sim 95\%$ of the codon families. In contrast, the exogenous genes 155 display an equally consistent mutational preference towards C and G ending codons (Table 156 S1). As a result, only the two codon amino acid Phenylalanine (Phe, F) has the same rank 157 order for the endogenous and exogenous ΔM values.

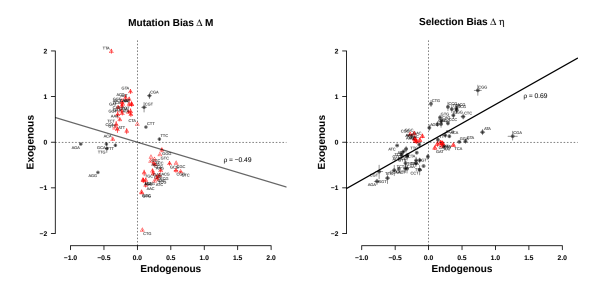


Figure 2: Comparison of (a) mutation bias ΔM and (b) selection bias $\Delta \eta$ parameters for endogenous and exogenous genes. Estimates are relative to the mean for each codon family. Black dots indicate ΔM or $\Delta \eta$ parameters with the same sign for the endogenous and exogenous genes, red dots indicate parameters with different signs. Black line shows the type II regression. Dashed lines mark quadrants.

Our estimates of $\Delta \eta$ for the endogenous and exogenous genes were positively correlated $(\rho = 0.69)$, indicating increased concordance of $\sim 53\%$ between the two selection environments (Figure 2). Nevertheless, the endogenous genes only show a selection preference for

158

159

160

 162 C and G ending codons in $\sim 58\%$ of the codon families. In contrast, the exogenous genes 163 display a strong preference for A and T ending codons in $\sim 89\%$ of the codon families.

The difference in codon preference between endogenous and exogenous genes is striking. 164 Fits to the complete L. kluyveri genome reveal that the relatively small exogenous gene set 165 has a disproportional effect on the model fit. We find that the complete L. kluyveri genome 166 is estimated to share the mutational preference with the exogenous genes in $\sim 78\%$ of codon 167 families with discordance between endogenous and exogenous genes. In two cases, Isoleucine 168 (Ile, I) and Arginine (Arg, R), the strong discrodunce in mutation preference results in a 169 estiamted codon preference in the complete L. kluyveri genome that is not reflected by either 170 endogenous nor exogenous genes. 171

The impact of the small exogenous gene set on the fit to the complete L. kluyveri genome is less prevalent in our estimates of selection bias $\Delta \eta$ but still strong. 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 with discordance between endogenous and exogenous genes. Therefore, it is important to recognize and treat endogenous and exogenous genes as separate sets to avoid the inference of incorrect synonymous codon preferences.

178 Determining Source of Exogenous Genes

We combined our estimates of mutation bias (ΔM) and selection bias $(\Delta \eta)$ with synteny in-179 formation and searched for potential source lineages of the introgressed region. We examined 180 38 yeast lineages of which two (Eremothecium gossypii and Candida dubliniensis) showed a 181 strong positive correlation in codon usage (Figure 3a). The endogenous L. kluyveri genome 182 exhibits codon usage very similar to most yeast lineages examined, indicating little varia-183 tion in codon usage among the examined yeasts (Figure 3b). Four lineages show a positive 184 correlation for ΔM and $\Delta \eta$ with the exogenous genes and have a weak to moderate positive 185 correlation in selection bias with the endogenous genes; but, like the exogenous genes, tend to have a negative correlation in ΔM with the endogenous genes. 187

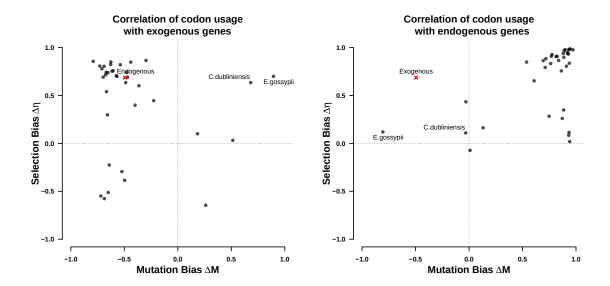


Figure 3: Correlation of ΔM and $\Delta \eta$ of the (a) exogenous and (b) endogenous genes with 38 examined yeast lineages. Dots indicate the correlation of ΔM and $\Delta \eta$ of the lineages with the endogenous and exogenous parameter estimates. All regressions were performed using a type II regression.

Comparing synteny between the exogenous 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 coverage (Figure S1a,b). *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 group(Figure S1b). Given these results, we conclude that the *E. gossypii* lineage is the most likely source of the introgressed exogenous genes.

5 Estimating Introgression Age

We estimated the introgression age using an exponential model of decay for mutation bias, by assuming that E. gossypii is still representative of the mutation bias of its ancestral source lineage at the time of the introgression. We utilize the ΔM estimates for all two codon amino acids and infer the age of the introgression to be on the order of $6.2 \pm 1.2 \times 10^8$ generations. We assume a mutation rate of 3.8×10^{-10} per nucleotide per generation, a value in line with other estimates [Zhu et al., 2014, Lang and Murray, 2008]. *L. kluyveri* experiences between one and eight generations per day, we therefore expect the introgression to have occurred about 205, 000 to 1,600,000 years ago which is longer than previous estimates of Friedrich et al. [2015]. However, our estimates are likely overestimates as they assume a purely neutral decay.

Furthermore, we estimated the persistence of the signal of the foreign cellular environment. Assuming that differences in mutation bias will decay more slowly than differences in selection bias, we predict that the ΔM signal of the source cellular environment will have decayed to be within one percent of the L. kluyveri environment within about $5.4 \pm 0.2 \times 10^9$ generations.

Fitness Burden of the Exogenous Genes

Estimates of selection bias for the exogenous genes show that, while well correlated with the endogenous genes, only nine amino acids share the optimal codon. We therefore expect 213 that the introgressed genes represent a significant reduction in fitness, or genetic load for L. 214 kluyveri, and even more so at the time of introgression. As the introgression occurred before 215 the diversification of L. kluyveri and has fixed since then throughout the various populations, 216 we are left without the original chromosome arm [Friedrich et al., 2015]. However, using our 217 estimates of ΔM and $\Delta \eta$ from the endogenous genes, we can estimate the genetic load of the 218 exogenous genes relative to an expected gene set. We define genetic load as the difference 219 between the fitness of an expected, replaced endogenous gene and the inferred introgressed 220 gene relative to drift $sN_e \propto \phi \Delta \eta$ (See Methods for details). 221

We estimate the genetic load of the exogenous genes at the time of introgression (Figure 4a) and currently (Figure 4b). These estimates are dependent on three key assumptions.

First, we assume again that the current cellular environment of *E. gossypii* is reflective of the ancestral environment. Second, we assume that the current amino acid composition of the exogenous genes is the same as in the replaced endogenous genes. Third, we assume that the

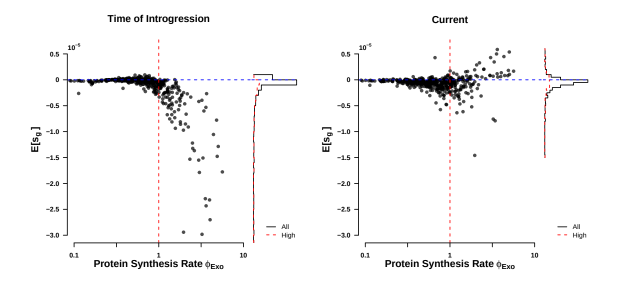


Figure 4: Fitness burden $\Delta s N_e$ (a) at the time of introgression ($\kappa = 5$), and (b) currently ($\kappa = 1$).

difference in the efficacy of selection between E. gossypii and L. kluyveri can be described with a simple scaling term we call κ (Figure S2b). As $\Delta \eta$ is defined as $\Delta \eta = 2N_e q(\eta_i - \eta_j)$, we can not distinguish if κ is a scaling on protein synthesis rate ϕ , effective population size N_e the value of an ATP q[Gilchrist et al., 2015].

At the time of the introgression, we predict that only a few genes were weakly exapted (Figure 4a) with all high expression genes ($\phi > 1$) being maladapted to the novel cellular environment. However, these highly expressed genes show the greatest rate of adaptation to the L. kluyveri cellular environment (Figures 4a, S3).

5 Discussion

Using ROC SEMPPR we show that the L. kluyveri genome contains two distinct signatures of cellular environments, its own endogenous and a foreign exogenous one obtained by an introgression event ($\Delta AIC = 78,000$). Following Payen et al. [2009], who defined the boundary of the anomalous chromosome region based on its elevated GC content, we partitioned the L. kluyveri genome into an endogenous and an exogenous gene set using gene location.

We estimated the codon usage of the entire *L. kluyveri* genome and the separated endogenous and exogenous gene sets (Figure S4). Both, Mutation bias and selection bias differ
between endogenous and exogenous genes. The endogenous genes show a strong mutation
bias towards A/T ending codons, while the exogenous genes show mutation is bias towards
G/C ending codons. We observed the reversed to be true in selection bias, leading to a
strong mismatch in codon usage between the gene sets, supporting our notion of two distinct
signatures of codon usage.

Only half of the codon families share the same optimal codon in the endogenous and 248 exogenous gene sets. However, we find that the strength of selection within a codon family 249 differs between gene sets, causing a change in rank order. Nevertheless, we find a high corre-250 lation for our estimates of selection bias $\Delta \eta$ between the two gene sets. Our estimates of the 251 optimal codon differ in nine cases between endogenous and exogenous genes. Interestingly, 252 when the difference in codon usage is ignored, we find that in seven out of these nine cases 253 the exogenous codon preference is inferred as optimal (Table S2). We find even greater dis-254 coordance in our estimates of ΔM between endogenous and exogenous gene sets (Table S1). 255 Without recognizing this difference in codon preference our estimates would not have been 256 reflective of the actual codon usage of the L. kluyveri genome but of a relatively small introgressed gene set. This shows that a small number of exogenous genes ($\sim 9\%$ of genes) can 258 have a disproportional impact on our estimates of ΔM and $\Delta \eta$ when fitting ROC SEMPPR 259 to the entire L. kluyveri genome. While this is surprising, it highlights the importance to 260 recognize differences in codon usage within a genome. Our results also indicate that we can 261 attribute the higher GC content in the exogenous genes mostly to differences in mutation 262 bias favoring G/C ending codons rather than a novel selective force. 263

Separating the endogenous and exogenous genes improves our estimates of protein synthesis rate ϕ by 17% relative to the full genome estimate ($\rho = 0.59$ vs. $\rho = 0.69$, respectively). Furthermore, we find that the variation in our estimates of ϕ is more consistent with the current understanding of gene expression (compare Figure 1a and b). Small variation in ϕ

estimates may serve as an indicator for the presents of the signature of multiple cellular environments in future work. In the case of the L. kluyveri genome, finding a severe mismatch 269 in ΔM causes ϕ values for low expression genes ($\phi < 1$) to increase towards the inflection 270 point where the dominance of mutation gives way to selection. In the case of the two codon 271 amino acids, the inflection point represents the point at which mutation and selection are 272 contributing equally to the probability of a codons occurrence. We find this inflection point 273 around $\phi = 1$ for most amino acids (Figure S4). However, ROC SEMPPR assumes that 274 estimates of ϕ follow a log-normal distribution with an expected value $E[\phi] = 1$. This as-275 sumption allows us to interpret $\Delta \eta$ as the strength of selection relative to drift (sN_e) for a 276 codon in a gene with the average protein synthesis rate $\phi = 1$. However, tying the mean 277 and standard deviation of the prior distribution together. Therefore, an increase in ϕ for low 278 expression genes has to be meet with a decrease of ϕ for high expression genes, reducing the 279 overall variance in ϕ (see Gilchrist et al. [2015] for details). 280

Having shown that the introgressed exogenous genes reflect a foreign cellular environ-281 ment, we used the quantitative estimates of mutation bias ΔM and selection bias $\Delta \eta$ from 282 ROC SEMPPR to identify potential source lineages. The comparison of the endogenous 283 and exogenous ΔM and $\Delta \eta$ estimates to 38 other yeast lineages revealed that most yeasts examined share similarity in mutation bias (Figure 2). Similar, we find strong similarities in 285 selection bias between examined yeasts, potentially indicating stabilizing selection on codon 286 usage. However, the exogenous genes do not share this commonality (Figure 2a), as their 287 mutation bias strongly deviates from the endogenous genes and most other yeast species 288 examined. This large difference in mutation bias between endogenous and exogenous genes 289 allowed us to limit our candidate list to only two likely lineages, C. dubliniensis and E. 290 gossypii. Interestingly, we did not find Lachancea thermotolerance, a thermophilic lineage 291 closely related to L. kluyveri, as a potential candidate. While L. thermotolerance does have 292 a strong synteny relationship with L. kluyveri, it does not show similarity in codon usage 293 with the exogenous genes and does not share their high GC content. 294

Inference of synteny relationships between the exogenous region and C. dubliniensis and 295 E. gossypii as well as closely related species showed that synteny relationship is limited to the 296 Saccharomycetaceae clade (Figure S1b). E. qossypii showed the highest syntenty coverage 297 and is the only species with similar codon usage. Furthermore, E. qossypii is the only species 298 examined with a GC content > 50\% like it is observed in the exogenous region. The synteny 299 coverage extends along the whole exogenous regions with the exception to the very 3' and 5' 300 end of the region. The lack of synteny at the ends of the region also coincides with a drop 301 in GC content, potentially indicating remains of the original replaced region or increased 302 adaptation. The ancestral introgressed region may have also broken up in E. qossupii as we 303 find non overlapping synteny with chromosomes VI and V as well as have indication that 304 the C chromosome of L. kluyveri very robust to recombination events [Payen et al., 2009, 305 Vakirlis et al., 2016]. 306

With E. gossypii identified as potential source lineage of the introgressed region, we 307 inferred the time past since the introgression occurred using our estimates of mutation bias 308 ΔM . The ΔM estimates are well suited for this task as they are free of the influence of 309 selection and unbiased by N_e and other scaling terms, which is in contrast to our estimates 310 of $\Delta \eta$ [Gilchrist et al., 2015]. We estimated the time since introgression to be on the order of 6×10^8 generations, which is ~ 10 times longer time than a previous estimate by Friedrich 312 et al. [2015] of a minimum of 5.6×10^7 generations . However, our estimate implicitly assumes 313 all mutations are neutral, it is therefore a conservative estimate, potentially overestimating 314 the time since introgression. Our estimate also depend on the assumption that the E. qossypii 315 cellular environment reflects the ancestral environment at the time of the introgression. If the 316 the ancestral mutation environment was more similar to the L. kluyveri environment at the 317 time of the introgression than the E. gossypii environment is today, we would overestimate 318 this time. On the other hand, we would underestimate the time since introgression if the 319 two cellular environments were more dissimilar. We could have attempted to reconstruct 320 the ancestral state of E. qossypii, however, as methods for ancestral state reconstruction are 321

phenomenological, assumptions would be unclear.

The estimates of mutation bias ΔM also allow us the infer the time until the signature of the exogenous cellular environment will have decayed to be indistinguishable at about one percent difference. Our estimate of decay is an order of magnitude greater than our estimate of the time since introgression (5 × 10⁹ and 6 × 10⁸ generations). Estimates of decay based on ΔM are more conservative as we expect differences in $\Delta \eta$ to decay before due to selection favoring the decay.

As we have determined that the introgression event has a long persisting exogenous 329 signature, it is important to understand the fitness consequences of such an event. We 330 estimated the genetic load that the exogenous genes represent assuming that the replaced 331 endogenous genes and the new exogenous genes had the same amino acid composition. This 332 assumption, along with the assumption that the current L. kluyveri cellular environment is 333 reflective of the cellular environment at the time of the introgression is necessary to estimate 334 the expected endogenous sequence that was replaced. Our results show that individual low 335 expression genes contribute little to the genetic load, and show less adaptation to the novel 336 cellular environment (Figure 4, S3). A small number of low expression genes even appear 337 exapted, likely due to the mutation bias in the endogenous genes matching the selection bias in the exogenous genes for G/C ending codons. Highly expressed genes on the other hand have greatly adapted to the L. kluyveri cellular environment. This, however, does not mean that these genes show a higher rate of evolution, but that small changes in their 341 sequence have large impacts on the fitness burden these sequences represent. To this day, the 342 exogenous genes represent a significant fitness burden on L. kluyveri. However, our estimates 343 are conservative as we do not account for potential changes in the codon usage of E. qossypii. 344 While divergent evolution in codon usage between E. qossypii and L. kluyveri would cause 345 us to overestimate the genetic load, convergent evolution, on the other hand, would cause 346 us to underestimate the genetic load. However, as the introgression appears to have reached 347 fixation [Friedrich et al., 2015], the genetic load relative to the replaced chromosome arm is only of theoretical interest.

The large genetic load the exogenous genes represented at the time of the introgression 350 indicates that the fixation of the introgression was a very unlikely event in a population 351 with a large N_e as it is typical for yeasts. It is hard to contextualize the probability of this 352 introgression being fixed as we are not aware of any estimates of the frequency at which 353 such large scale introgressions of genes with very different signatures of codon usage occur. 354 One example is Saccharomyces bayanus, a hybrid of Saccharomyces uvarum, Saccharomyces 355 cerevisiae, and Saccharomyces eubayanus. However, unlike with L. kluyveri and E. gossypii 356 it appears that the donor lineages show similar codon usage. Saccharomyces cerevisiae 357 and Saccharomyces eubayanus show a very strong correlation between selection bias $\Delta \eta$ of 358 $\rho = 0.98$ and a strong correlation between mutation bias ΔM of $\rho = 0.83$ We were unable to 359 identify codon usage for Saccharomyces uvarum. However, L. kluyveri diverged about 85 Mya 360 ago from the rest of the Lachancea clade. This represents between 10^{10} to 10^{11} generations. 361 Assuming for yeasts typical effective population size on the order of 10⁸, we are left with 10^{18} to 10^{19} opportunities for such an event to occur. In addition, the strong mutation bias 363 towards G/C ending codons in the exogenous genes may have contributed to the fixation of this introgression (include figure of ΔM v $\Delta \eta$). It is, on the other hand, also possible that despite their mismatch in codon usage, the exogenous genes have represented a fitness increase due to external environmental factors resulting in the fixation of the introgression. 367 In conclusion, our results show the usefulness of the separation of mutation bias and se-368 lection bias and the importance of recognizing the presence of multiple cellular environments 369 in the study of codon usage. We also illustrate how a mechanistic model like ROC SEMPPR 370 and the quantitative estimates it provides can be used for more sophisticated hypothesis 371 testing in the future. In contrast to other approaches used to study codon usage like CAI 372 [Sharp, 1987] or tAI [dos Reis et al., 2004], ROC SEMPPR is sensitive to differences in mu-373 tation bias. We highlight potential pitfalls when estimating codon preferences, as estimates 374 can be biased by the signature of a second, historical cellular environment. In addition, 375

we show how quantitative estimates of mutation bias and selection relative to drift can be
obtained from codon data and used to infer the fitness cost of an introgression as well as its
history and potential future.

379 Materials and Methods

Separating 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/ (last accessed: 09-27-2014) and the annotation for *L. kluyveri* NRRL Y-12651 (assembly ASM14922v1) from NCBI (last accessed: 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.

390 Model 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). ROC SEMPPR was run from multiple starting values for at least 250,000 iterations, every 50th sample was collected to reduce autocorrelation. After manual inspection to verify that the MCMC had converged, parameter posterior means were estimated from the last 500 samples.

³⁹⁶ Comparing codon specific parameter estimates

Because our ΔM and $\Delta \eta$ are meaningful only for comparisons between synonymous codons, ROC SEMPPR returns mutation bias ΔM and selection bias $\Delta \eta$ parameter values relative to a reference codon. While ROC SEMPPR's choice of the reference codon is largely arbitrary, changes in the reference codon affect [NEED TO COMPLETE] To circumvent this issue, we express our estimates relative to the mean for each codon family.

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.

$$\Delta M_{i,a}^c = \Delta M_{i,a} - \Delta \bar{M}_a \tag{1}$$

$$\Delta \eta_{i,a}^c = \Delta \eta_{i,a} - \bar{\Delta \eta_a} \tag{2}$$

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

406 Synteny

We obtained complete genome sequences from NCBI (last accessed: 02-05-2017). Genomes were aligned and checked for synteny using SyMAP (4.2) with default settings [Soderlund et al., 2006, 2011]. We assessed Synteny as percentage non-overlapping coverage of the exogenous gene region (Figure S1b).

111 Determining introgression timeline

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

$$\frac{dc_1}{dt} = -\mu_{1,2}c_1 - \mu_{2,1}(1 - c_1) \tag{3}$$

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 ΔM_{endo} can be directly related to the steady state of equation 3.

$$\frac{\mu_{2,1}}{\mu_{1,2} + \mu_{2,1}} = \frac{1}{1 + \exp(\Delta M_{endo})} \tag{4}$$

Solving for $\mu_{1,2}$ gives us $\mu_{1,2} = \Delta M_{endo} \exp(\mu_{2,1})$ which allows us to rewrite and solve equation 3 as

$$c_1(t) = \frac{\exp(-t(1 + \Delta M_{endo})\mu_{2,1}) \exp(t(1 + \Delta M_{endo})\mu_{2,1}) + (1 + \Delta M_{endo})K}{1 + \Delta M_{endo}}$$
(5)

where K is

$$K = \frac{-1 + c_1(0) + c_1(0)\Delta M_{endo}}{1 + \Delta M_{endo}}$$
(6)

Equation 5 was solved over time with a mutation rate $m_{2,1}$ of 3.8×10^{-10} per nucleotide per generation [Lang and Murray, 2008]. Initial codon frequencies $c_1(0)$ for each codon family where taken from our estimates of ΔM_{gos} from E. gossypii. Current codon frequencies for each codon family where taken from our estimates of ΔM from the exogenous genes. Mathematica (9.0.1.0) [Inc.] was used to calculate the time t_{exo} it takes for the initial codon frequencies $c_1(0)$ for each codon family to change to the current exogenous codon frequencies. The same equation was used to determine the time t_{endo} at which the signal of the exogenous cellular environment has decayed to within 1% of the endogenous environment.

$_{ ext{420}}$ Estimating fitness burden

To estimate the fitness burden, we made three key assumptions. First, we assumed that the current exogenous amino acid composition of genes is representative of the replaced endogenous genes. 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 of the source lineage and $L.\ kluyveri$ can be expressed as a scaling constant and that protein synthesis rate ϕ has not changed between the replaced endogenous and the introgressed exogenous genes.

We calculated the fitness burden each gene represents assuming additive fitness effects as

$$(sN_e)_g = \sum_{i}^{C} -\kappa \phi_g \Delta \eta_i n_{g,i} \tag{7}$$

where $(sN_e)g$ is the selection against translation inefficiency relative to drift. ϕ_g is the estimated protein synthesis rate for gene g in the exogenous gene set. $n_{g,i}$ is the codon count of each codon i in the codon set C for each gene g. κ is a constant, scaling the efficacy of selection between cellular environments. Like 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 S2b). Thus, we set $\kappa = 1$ if we calculate the $(sN_e)g$ for the endogenous and the current exogenous genes, and $\kappa = 5$ for $(sN_e)g$ 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

$$E[n_{g,i}] = \frac{\exp(-\Delta M_i - \Delta \eta_i \phi_g)}{\sum_{j=1}^{C} \exp(-\Delta M_j - \Delta \eta_j \phi_g)} \times m_{a_i}$$
 (8)

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

We report the fitness burden of the introgression as $\Delta s N_e = (sN_e)_{Intro} - (sN_e)_{Endo}$ where $(sN_e)_{Intro}$ is either the fitness burden at the time of the introgression or presently.

432 Acknowledgments

This work was supported in part by NSF Awards MCB-1120370 (MAG and RZ) and DEB1355033 (BCO, MAG, and RZ) with additional support from The University of Tennessee
Knoxville. CL received support as a Graduate Student Fellow at the National Institute
for Mathematical and Biological Synthesis, an Institute sponsored by the National Science
Foundation through NSF Award DBI-1300426, with additional support from UTK. The
authors would like to thank Brian C. O'Meara and Alexander Cope for their helpful criticisms
and suggestions for this work.

440 References

- T Ikemura. Codon usage and trna content in unicellular and multicellular organisms. *Molec- ular Biology and Evolution*, 2:13–34, 1985.
- M Gouy and C Gautier. Codon usage in bacteria: correlation with gene expressivity. *Nucleic Acids Research*, 10:7055–7074, 1982.
- Clia Payen, Gilles Fischer, Christian Marck, Caroline Proux, David James Sherman, JeanYves Coppe, Mark Johnston, Bernard Dujon, and Ccile Neuvglise. Unusual composition
 of a yeast chromosome arm is associated with its delayed replication. Genome Research,
 19(10):1710–1721, 2009.
- A Friedrich, C Reiser, G Fischer, and J Schacherer. Population genomics reveals chromosome-scale heterogeneous evolution in a protoploid yeast. *Molecular Biology and Evolution*, 32(1):184 192, 2015.

- MA Gilchrist, WC Chen, P Shah, CL Landerer, and R Zaretzki. Estimating gene expression
- and codon-specific translational efficiencies, mutation biases, and selection coefficients from
- genomic data alone. Genome Biology and Evolution, 7:1559–1579, 2015.
- ⁴⁵⁵ Cedric Landerer, Alexander Cope, Russell Zaretzki, and Michael A Gilchrist. Anacoda:
- analyzing codon data with bayesian mixture models. Bioinformatics, 34(14):2496–2498,
- 457 2018.
- 458 Yuan O Zhu, Mark L Siegal, David W Hall, and Dmitri A Petrov. Precise estimates of
- mutation rate and spectrum in yeast. Proceedings of the National Academy of Sciences,
- 460 111(22):E2310–E2318, 2014.
- 461 Gregory I. Lang and Andrew W. Murray. Estimating the per-base-pair mutation rate in the
- yeast saccharomyces cerevisiae. Genetics, 178(1):67 82, 2008. ISSN 0016-6731.
- Nikolaos Vakirlis, Véronique Sarilar, Guénola Drillon, Aubin Fleiss, Nicolas Agier, Jean-
- Philippe Meyniel, Lou Blanpain, Alessandra Carbone, Hugo Devillers, Kenny Dubois,
- Alexandre Gillet-Markowska, Stéphane Graziani, Nguyen Huu-Vang, Marion Poirel,
- 466 Cyrielle Reisser, Jonathan Schott, Joseph Schacherer, Ingrid Lafontaine, Bertrand
- Llorente, Cécile Neuvéglise, and Gilles Fischer. Reconstruction of ancestral chromosome
- architecture and gene repertoire reveals principles of genome evolution in a model yeast
- genus. Genome research, 26(7):918–32, 2016.
- 470 PM Sharp. The codon adaptatoin index a meassure of directional synonymous codon usage
- bias, and its potential applications. Nucleic Acids Research, 15:1281–1295, 1987.
- 472 M dos Reis, R Savva, and L Wernisch. Solving the riddle of codon usage preferences: a test
- for translational selection. Nucleic Acids Research, 32(17):5036–5044, 2004.
- 474 C Soderlund, W Nelson, A Shoemaker, and A Paterson. Symap A system for discovering
- and viewing syntenic regions of fpc maps. Genome Research, 16:1159 1168, 2006.

| Hypothesis | $\log(Likelihood)$ | n | AIC | $\Delta { m AIC}$ |
|------------------------|--------------------|---|-----------|-------------------|
| Endogenous & Exogenous | | | 5,235,598 | 0 |
| Combined | | | 5,311,060 | 75,462 |

Table 1: $\log(Likelihood)$, number of model parameters n, AIC, and Δ AIC.

- C Soderlund, M Bomhoff, and W Nelson. Symap v3.4: a turnkey synteny system with application to plant genomes. *Nucleic Acids Research*, 39(10):e68, 2011.
- Wolfram Research, Inc. Mathematica, Version 9.0. Champaign, IL, 2012.

Figures Figures

Table Table

| 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 | \mathbf{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 |
| 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 S1: Synonymous codons with the greatest mutational preference (i.e. largest ΔM value). Bold face codons indicate synonyms whose ...

Supplementary Material

Supporting Materials for Fitness consequences of mismatched codon usage by Landerer et al..

Tables

| 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 |

Table S2: Synonymous codon preference in the various data sets based on our estimates of $\Delta \eta$

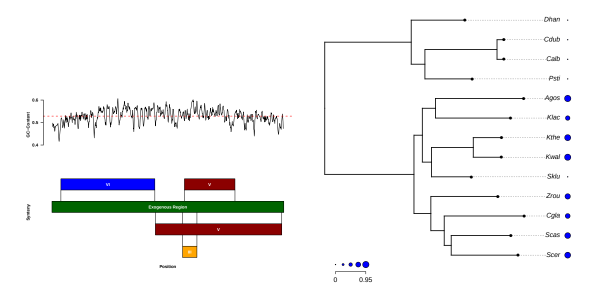


Figure S1: Suppl Fig: Synteny relationship of E. gossypii and the exogenous genes (left), Amount of synteny for each species (Units of std dev) checked for synteny.

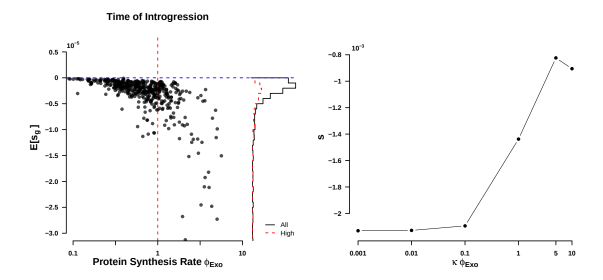
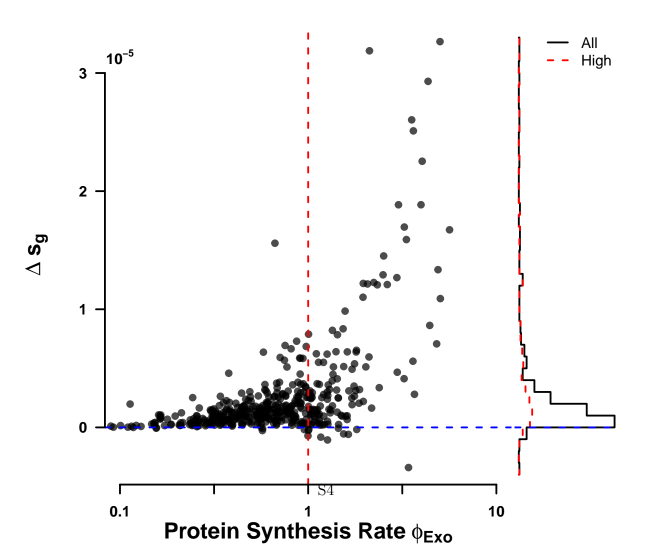


Figure S2: Suppl Fig: Fitness burden (left) without scaling of ϕ , and change of total fitness burden with scaling κ

Change in Selection Coefficient



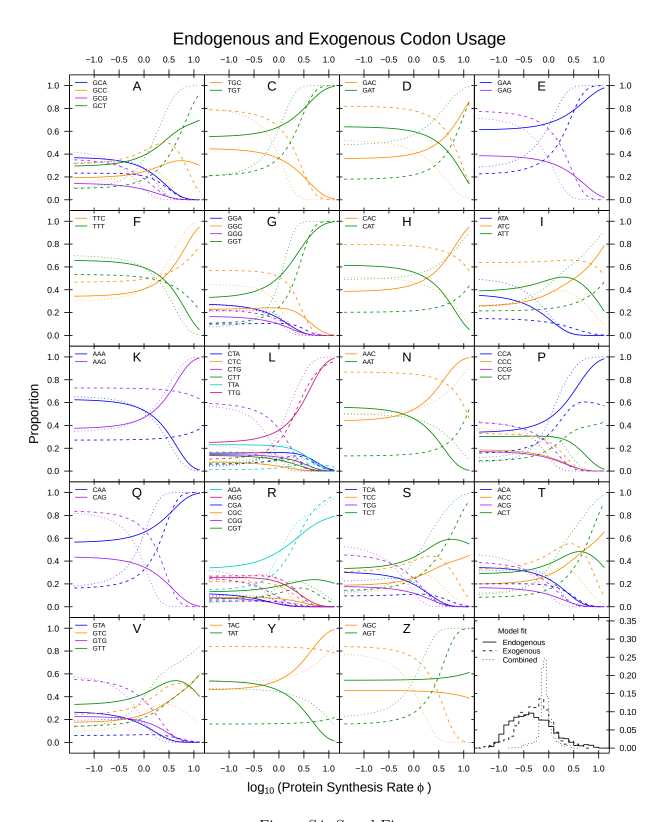


Figure S4: Suppl Fig