Supplemental Material for: Extreme purifying selection against point mutations in the human genome

Noah Dukler^{1,a}, Mehreen R. Mughal^{1,a}, Ritika Ramani¹, Yi-Fei Huang², and Adam Siepel^{1,*}

¹Simons Center for Quantitative Biology, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY

²Department of Biology and Huck Institutes of the Life Sciences, The Pennsylvania State University, University Park, PA

^aThese authors contributed equally to this work.

*Corresponding author

Supplemental Text

Approximation of the expected number of generations until extinction as a fraction of the neutral expectation

By Kimura and Ohta's formulas [1], the expected number of generations until a new mutant (at initial frequency $\frac{1}{2N}$) is lost from a finite population, as a function of the current population size N and the scaled selection coefficient $S=2N_e s$, where N_e is the effective population size, is approximately given by,

$$t(S) = \frac{2N_e}{N} \left[\ln \left(\frac{N}{S} \right) + 1 - \gamma \right], \qquad \gamma = 0.577...,$$
 (1)

in the case of a semidominant deleterious mutation (with $h=\frac{1}{2}$), and by,

$$t(0) = \frac{2N_e}{N}\ln(2N) \tag{2}$$

in the case of a neutral mutation. In the regime of interest, $\ln(N/S) \gg 1$, so the ratio of these quantities can be roughly approximated as,

$$\frac{t(S)}{t(0)} \approx \frac{\ln(N) - \ln(S)}{\ln(2N)} = \frac{\ln(N) - \ln(S)}{\ln(N) + \ln 2}.$$
 (3)

As discussed in the text, we estimate that ultraselected sites have values of $s_{\rm het}=\frac{1}{2}s$ of about 0.03. Assuming the typical value of $N_e=10^4$ for human populations, $S=2N_es=4N_es_{\rm het}=1200$, meaning that $\ln(S)\approx 7.1$. It is more difficult to know what N should be in this setting, but Weghorn et al. [2] have argued for a plausible range of $0.5-8.0\times10^6$ based on current demographic models for human populations. Thus, $\ln(N)$ ranges from about 13.1 to 15.9, and we obtain values for $\frac{t(S)}{t(0)}$ ranging from 0.43 to 0.53.

References

- [1] Kimura M, Ohta T. The average number of generations until extinction of an individual mutant gene in a finite population. Genetics. 1969;63(3):701–9.
- [2] Weghorn D, Balick DJ, Cassa C, Kosmicki JA, Daly MJ, Beier DR, et al. Applicability of the Mutation-Selection Balance Model to Population Genetics of Heterozygous Protein-Truncating Variants in Humans. Mol Biol Evol. 2019;36(8):1701–1710.
- [3] Yang RY, Quan J, Sodaei R, Aguet F, Segrè AV, Allen JA, et al. A systematic survey of human tissue-specific gene expression and splicing reveals new opportunities for therapeutic target identification and evaluation. bioRxiv. 2018;doi:10.1101/311563.
- [4] Kim BY, Huber CD, Lohmueller KE. Inference of the Distribution of Selection Coefficients for New Nonsynonymous Mutations Using Large Samples. Genetics. 2017;206(1):345–361.

Table S1: Ultraselection across the human genome (less conservative estimates)

Feature	λ_s	\pm (stderr)	no. sites (M)	prop. sites	\exp no. $(M)^a$	exp. prop. b	fold enrich.	\exp . lethal ^c	Shet
CDS	0.149	0.002	33.8	1.18%	4.9	31.6%	26.8	0.12	ı
5' UTR	-0.158	0.002	8.2	0.29%	0.0	0.0%	0.0	0.00	ı
3' UTR	0.023	0.002	36.1	1.26%	0.7	4.6%	3.6	0.02	ı
splice	0.464	0.002	0.8	0.03%	0.4	2.3%	85.0	0.01	2.0%
nonconserved IncRNA ^d	0.008	0.002	453.6	15.78%	1.8	11.8%	0.7	0.04	1
conserved IncRNA ^e	0.055	0.002	23.3	0.81%	1.2	7.7%	9.5	0.03	ı
nonconserved intron ^d	0.008	0.002	972.6	33.83%	4.2	26.8%	0.8	0.10	1
conserved intron ^e	0.057	0.002	44.3	1.54%	2.4	15.3%	6.6	0.06	1
nonconserved intergenic d	0.003	0.002	1255.5	43.67%	0.0	0.0%	0.0	0.00	1
conserved intergenic ^e	0.051	0.002	46.9	1.63%	2.2	14.2%	8.7	0.05	1
Total			2875.1	100.00%	15.6	100.0%		0.43	

"Expected number of ultraselected sites after adjusting for background. In this case, the estimate for nonconserved intergenic regions (0.003) was subtracted from each estimate of λ_s (see **Table 1** for a more conservative correction).

 $[^]b$ Expected proportion of ultraselected sites after adjusting for background. c Expected number of new lethal or near-lethal mutations per diploid individual, assuming a mutation rate of 1.2×10^{-8} per generation per site.

^dSites not classified as conserved by phastCons.

^eSites classified as conserved by phastCons.

Distribution	α^a	θ^a	$\pi_0{}^b$	$\operatorname{mean} g(x)$	mean $f(x)$	mean $h(x)$	λ_s	estimated s_{het}
Kim et al.,	0.1930	0.0168	3.1%	0.0023	0.0032	0.0303	0.0416	-
0d CDS	0.8678	0.0168	3.1%	0.0101	0.0141	0.0275	0.2340	0.0242
miRNA	1.0700	0.0168	0.0%	0.0137	0.0189	0.0312	0.3396	0.0316
TFBS	0.5500	0.0168	70.0%	0.0017	0.0028	0.0277	0.0275	-

^aParameters of assumed Gamma distribution, where α is the shape parameter and θ is the scale parameter ^bWeight of point mass at zero.

Table S2: Means of full simulated DFE (f(x)), DFE associated with remaining rare variants (g(x)), and DFE inferred to be associated with the "missing" rare variants (h(x)) by mixture decomposition (see **Methods**). Also shown are the estimated values of λ_s from simulated data, as well as the corresponding value of s_{het} (for $\lambda_s > 0.18$).

Supplemental Figures

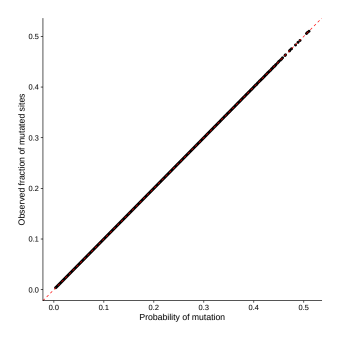


Figure S1: Predicted vs. observed rates of rare variants in designated neutral regions. Each point represents a single 50kb bin. Along the x-axis are the average values of P_i across that bin, as predicted by our mutation model, and along the y axis are the observed rates at which rare variants occur within that bin. The plot shows that the mutation model is well calibrated genome-wide for neutral sites.

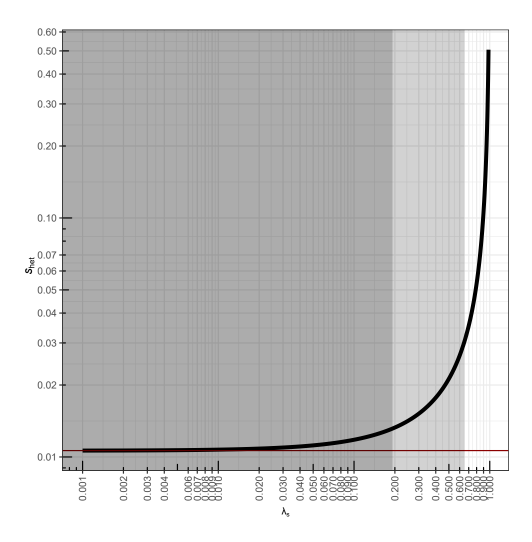


Figure S2: Theoretical relationship between λ_s and the selection coefficient against heterozygous mutations, s_{het} . Curve represents equation 2 with N=71,702 and $c=1.35\times 10^7$ based on our real data set (see **Methods**). The dark shaded region ($\lambda_s<0.18,\,s_{\text{het}}<0.013$) indicates the approximate regime where the relationship no longer yields an accurate estimator for s_{het} with our data, and the lighter shaded region ($0.18<\lambda_s<0.65,\,0.013< s_{\text{het}}<0.03$) indicates the regime where the estimator is slightly inflated but still useful as a guide (see **Supplemental Fig. S3**).

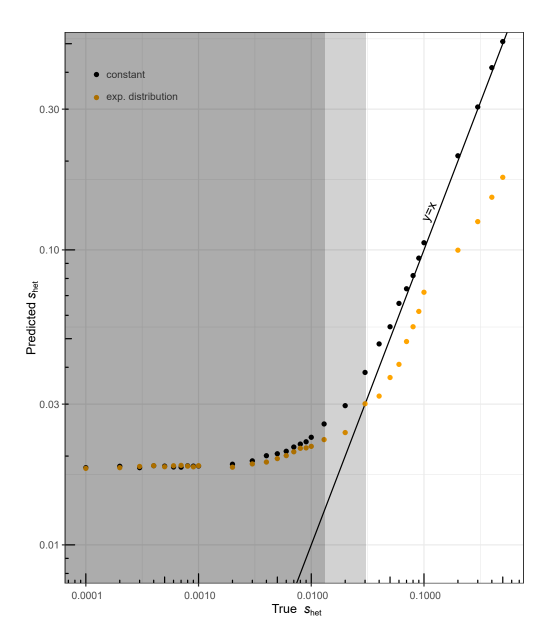


Figure S3: True vs. predicted values of s_{het} in simulation. Data sets of 71,702 diploid individuals and 100,000 sites were simulated using software from ref. [2] with mean s_{het} ranging from 0.0001 to 0.5 (x_{axis}). In one version, all sites were assigned the same "true" value of s_{het} ("constant"; black points) and, in another, sitewise values of s_{het} were drawn from an exponential distribution with the given mean value ("exp. distribution"; orange points). ExtRaINSIGHT was applied to each simulated data set, and then the estimated value of λ_s was converted to a predicted s_{het} (y_{axis}) using equation 2. All simulations assumed a European demographic history (see **Methods**). As in **Supplemental Fig. S2**, the dark and light gray regions respectively indicate the regimes in which the estimator for s_{het} is no longer useful, and is inflated but still approximately useful.

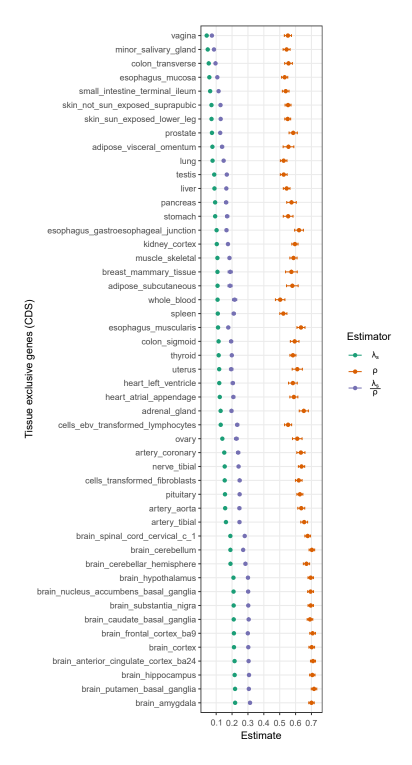


Figure S4: Measures of purifying selection in protein-coding genes exhibiting tissue-specific gene expression. Tissue-specific genes were obtained from ref. [3] as detailed in the Methods section. An estimate for each tissue is shown for both ExtRaINSIGHT (λ_s) and INSIGHT (ρ). Error bars indicate one standard error (see Methods).

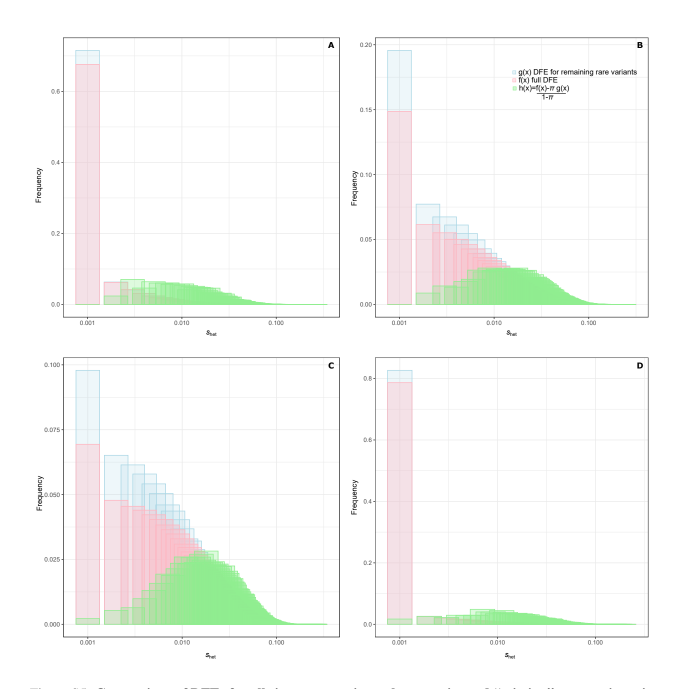


Figure S5: Comparison of DFEs for all sites, rare variants that remain, and "missing" rare variants in simulations. Simulated DFEs (f(x); pink), DFEs for rare variants that remain in the data (g(x); blue), and DFEs inferred by mixture decomposition for the rare variants that are missing (h(x); green). Results are shown for four distinct DFEs: (A) a DFE published by Kim et al. [4] consisting of a mixture of a point-mass at zero (with weight 0.031) and a Gamma distribution with α =0.1930 and θ =0.0168. (B) a modified DFE designed to approximately match our observations at 0d sites in coding regions, consisting of a mixture of a point-mass at zero (weight 0.031) and a Gamma distribution with α =0.8687 and θ =0.0168. (C) a modified DFE designed to approximately match our observations at evolutionarily ancient miRNAs, equal to a Gamma distribution with α =1.07 and θ =0.0168. (D) a modified DFE designed to approximately match our observations at TFBS, consisting of a mixture of a point-mass at zero (with weight 70%) and a Gamma distribution with α =0.55 and θ =0.0168. Means of these distributions along with our λ_s and s_{het} estimates are shown in Supplemental Table S2.