

# Demography and linked selection in wild and domesticated maize

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**Unique selective and demographic forces operate on domesticated plant species.** I don't agree with this statement; characteristics of demography and selection vary across crops and I wouldn't be surprised to find similar demographies and patterns of selection between crop and wild species. These forces interact during and after domestication to generate the patterns of DNA variability that are persistent today. To quantify the interplay between demography and selection, we investigated genetic diversity in maize, one of the most important crops for food, feed, and fuel world-wide. Our sample included whole genome sequence data from 23 maize and 13 teosinte individuals. We obtained a complete estimate of the population size fluctuations and other demographic parameters experienced by maize as it was domesticated from teosinte. Here, we show that maize went through a domestication bottleneck with a population size of approximately 5% that of teosinte before it experienced rapid population size expansion post-domestication. We observe that hard sweeps on genic mutations are not the primary force driving maize evolution. We find that a reduced population size during domestication decreased the efficiency of purifying selection to purge deleterious alleles from maize. However, expansion after domestication has since increased the efficiency of purifying selection to levels exceeding those seen in teosinte. Our results demonstrate that particularly in domesticated species or bottlenecked species right, observations can be generalized to wild species as well; I don't think there's any need to say crops are a special case, demographic and selective history in the ancient and recent past both contribute to genetic variability that is present today, providing substantial implications for the continued improvement of domesticated species. I'd end this more generally, expressing the importance of knowing demography when characterizing selection genome-wide

The genetic properties of populations are determined by a constant interplay between genetic drift and selection. Drift is the consequence of a finite population size, such that random sampling of gametes every generation will incrementally lead to genetic changes. In other words, the demographic history of a population, which includes its population size fluctuations and mating patterns, has a long-lasting impact on the patterns of DNA variability exhibited by that population. Conversely, selection systematically alters a population by favoring particular alleles at the expense of others. In contrast to drift, which results from demographic history, selection is the result of the relationship between genotypes, phenotypes, and ultimately fitness. Importantly, demography and selection do not operate independently to determine genetic variability. Instead, these forces are engaged in a continuous battle by which each masks and obfuscates the outcomes of the other, ultimately meaning that the selective history of a species cannot be properly evaluated without an thorough estimate of demographic history, and demography cannot be evaluated without an understanding of selection.

For instance, supposed neutrally-evolving DNA is often used to estimate historical demographic parameters of a population such as effective population size, structure, and expansion history [1, 2]. However, researchers have called into question whether or not there are reasonable approaches for identifying neutral regions of the genome, since the effects of selection can be wide-ranging [3, 4]. For example, in *Drosophila* it appears that the majority of the genome is impacted by the effects of

selection through linkage [5]. A natural next question, therefore, is how can demographic parameters be estimated independently of selective parameters? One technique utilized to address this problem is limiting analyses to only sites far from genes [6], but as *Drosophila* demonstrates, it is difficult to be certain that sites even in gene-poor regions of the genome are not influenced by linked selection. Ultimately, an understanding of which sites have the potential to be adaptive, and how these affect genome-wide patterns of variability through linkage, is required for reasonable demographic inference.

Maize represents an excellent organism to study these phenomena. Maize is a species of tremendous importance worldwide as both a staple crop [7] and as a model for understanding plant evolution [8]. Broadly speaking, archaeological and genetic studies have established that maize domestication is likely to have taken place in Mexico approximately 9,000 years bp [9, 10]. Teosinte, the most recent wild ancestor to maize, remains extant throughout much of the Americas [11]. Additionally, several large-effect domestication loci [12–14] and putative domestication regions [15] have been identified. But despite all that is known about maize domestication, the param-

## Significance

Both selection and demographic change play important roles in shaping diversity across the genome, but clear empirical examples of the interplay of these two forces are lacking. Here we document the combined effects of demography and linked selection on genome-wide diversity in domesticated maize and its wild ancestor teosinte. We estimate that maize underwent a bottleneck of 5% of the size of the ancestral teosinte population, but that recent expansion has resulted in a maize population perhaps orders of magnitude larger than teosinte. We show that positive selection on new mutations has had relatively little effect on genetic diversity, but that selection against deleterious mutations has dramatically reduced diversity in and immediately around genes in both taxa. We find that the relative effect of selection depend qualitatively on the age of the polymorphisms evaluated: while older polymorphisms in maize show more limited effects of linked selection, new mutations instead reflect its larger current size and more efficient purifying selection. Our results demonstrate that a complete understanding of genome-wide patterns of diversity will require careful assessment of both demographic history and the effects of linked selection.

## Reserved for Publication Footnotes

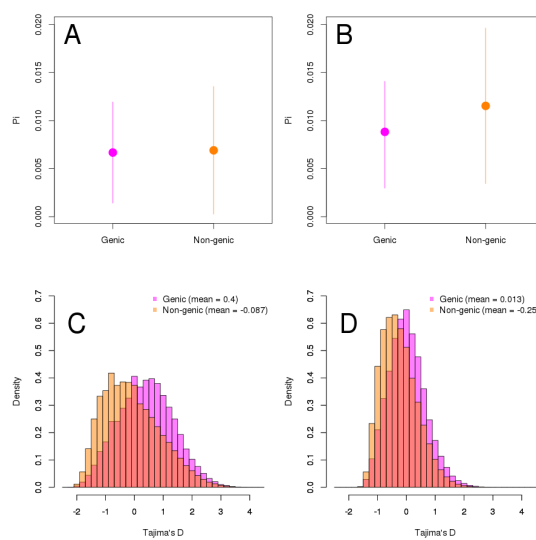
eters of the domestication process remain uncertain. Specifically, the size of the maize domestication bottleneck has not been estimated independently of the bottleneck's duration, nor are there sequence-based estimates of the effective population size of modern maize. Sequence information from maize and teosinte may therefore be utilized to address these questions.

To that end, the objectives of our study were to 1) investigate the relative importance of different forms of selection on whole-genome variability, using maize and teosinte as a model 2) research the impact that the demography history has had on genetic variability in maize, and how this compares to the impact of a different demographic history in teosinte; and 3) precisely estimate the parameters of the maize domestication bottleneck. [16]. *A quick sentence here about why this matters broadly would help...what kind of basic inferences will this research allow us to make about the interplay of demography and selection?*

**D**omesticated plant species evolve in a unique fashion compared to their wild counterparts [17]. This is a result of both the anthropomorphic nature of artificial selection on domestics [18] as well as the demographic characteristics of the domestication bottleneck(s) that they tend to have experienced [19]. However, the complex interplay between selective pressures and demographic limitations, and the impact that this interplay has on identifying selection and understanding demography, is not fully understood. Although a large body of research that involves searching the genomes of domesticated species for evidence of positive selection exists [15, 20–22], these studies tend to focus on identifying or mapping particular genes or regions that play an important role in phenotypic evolution. In contrast, knowledge regarding the impacts that demography and selection have on whole-genome patterns of genetic variability remains limited.

## Results

To investigate how demography and linked selection have shaped patterns of diversity in maize and teosinte, we analyzed data from 23 maize and 13 teosinte genomes from the maize HapMap 2 and HapMap 3 projects [16] *add cite to HM3*. We find broad differences in genic and intergenic diversity consistent with earlier results [15](Figure 1). *pi graph should use greek*



**Fig. 1.** Pairwise diversity  $\pi$  (A,B) and Tajima's D (C,D) in 1kb windows from genic and nongenic regions of maize (A,C) and teosinte (B,D). Shown in A and B are means and one standard deviation.

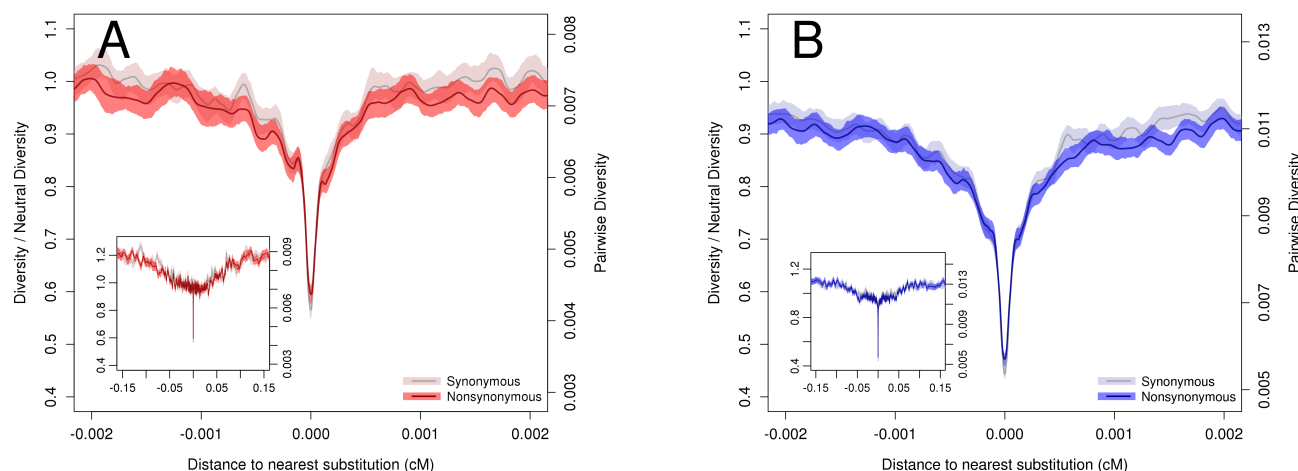
**letter  $\pi$**  In maize, mean pairwise diversity ( $\pi$ ) within genes was significantly lower than at sites at least 5 kb away from genes (0.00668 vs 0.00691,  $p < 2 \times 10^{-44}$ ). Diversity differences in teosinte are even more pronounced (0.0088 vs. 0.0115,  $p \approx 0$ ). Differences were also apparent in the site frequency spectrum, with mean Tajima's D positive in genic regions in both maize (0.4) and teosinte (0.013) but negative outside of genes (-0.087 in maize and -0.25 in teosinte,  $p \approx 0$  for both comparisons). These observations suggest that diversity in genes is not evolving neutrally, but instead is reduced by the impacts of selection on linked sites.

**Hard sweeps do not explain diversity differences.** When selection increases the frequency of a new beneficial mutation, a signature of reduced diversity is left at surrounding linked sites [23]. To evaluate whether patterns of such "hard sweeps" could explain observed differences in diversity between genic and intergenic regions of the genome, we compared diversity around missense and synonymous substitutions between *Trip-sacum* and either maize or teosinte. If a proportion of missense mutations have been fixed due to hard sweeps, diversity around these substitutions should be lower than around synonymous substitutions. We observe this pattern around the causative amino acid substitution in the the domestication locus *tga1* (Figure S1), likely the result of a hard sweep during domestication [14, 24]. *Awesome* Genome-wide, however, we observe no differences in diversity between synonymous and missense substitutions in either maize or teosinte (Figure 2). *I'm curious whether linked selection is affecting the diversity around a sizable proportion of synonymous substitutions. Would it be worthwhile to look at diversity around the subset of synonymous substitutions that are the least linked to missense substitutions just to make sure it is not substantially higher? Or perhaps generate a plot of diversity around synonymous substitutions vs. distance of these subs from missense subs?*

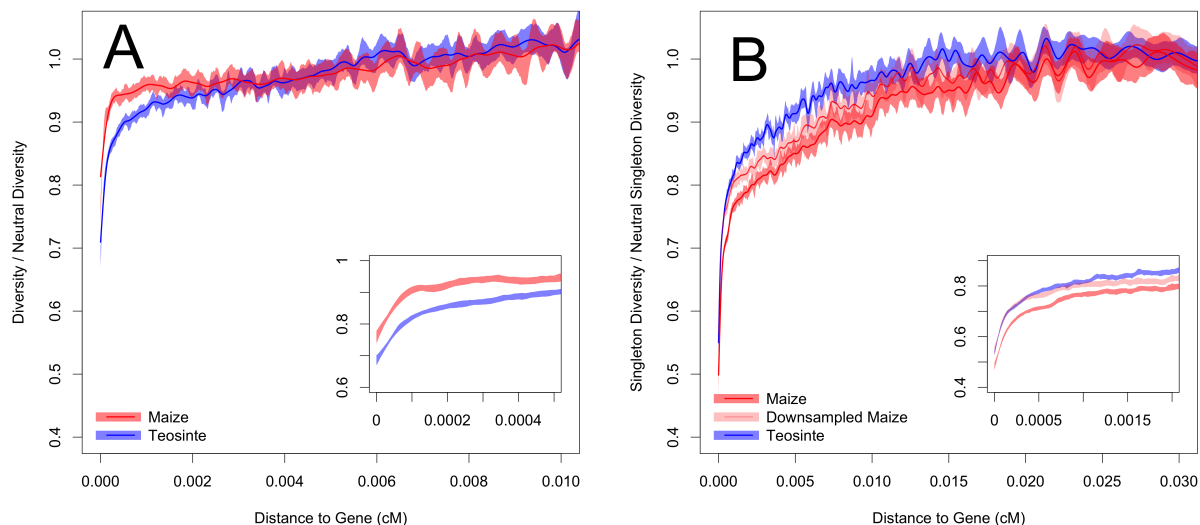
Previous analyses have suggested that this approach may have limited power because a relatively high proportion of missense substitutions will be found in genes that are under weak purifying selection and thus have higher genetic diversity [25]. To address this concern, we took advantage of genome-wide estimates of evolutionary constraint [26], calculated using genomic evolutionary rate profile (GERP) scores [27]. We then evaluated substitutions only in subsets of genes in the highest and lowest 10% quantile of mean GERP score, putatively representing genes under the strongest and weakest purifying selection (Figure S3). As expected, we see higher diversity around substitutions in genes under weak purifying selection *there a table or figure we can reference?*, but we still find no difference between synonymous and missense substitutions in either subset of the data. Taken together, these data suggest hard sweeps do not play a major role in patterning genic diversity in either maize or teosinte.

**Diversity is strongly influenced by purifying selection.** Selection can also reduce diversity in functional regions of the genome via removal of deleterious mutations, a process known as purifying or background selection [28]. We investigated purifying selection in maize and teosinte by evaluating the reduction of diversity around genes. Pairwise diversity is strongly reduced within genes for both maize and teosinte (Figure 3A) but recovers quickly at sites outside of genes, consistent with the low levels of linkage disequilibrium generally observed in these subspecies [16, 29]. The reduction in relative diversity is more pronounced in teosinte, reaching lower levels in genes and occurring over a wider region.

Our comparison of synonymous and missense substitutions has low power to detect the effects of selection acting on multiple mutations or standing genetic variation, because in such



**Fig. 2.** Pairwise diversity surrounding synonymous and missense substitutions in **A** maize and **B** teosinte. Axes show absolute diversity values (right) and values relative to mean nucleotide diversity in windows  $\geq 0.01\text{cM}$  from a substitution (left). Lines depict a loess curve (span of 0.01) and shading represents bootstrap-based 95% confidence intervals. Inset plots depict a larger range on the x-axis.

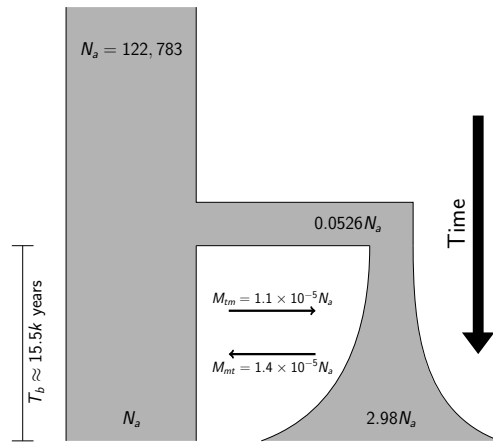


**Fig. 3.** Relative diversity versus distance to nearest gene in maize and teosinte. Shown are **A** pairwise nucleotide diversity and **B** singleton diversity. Relative diversity is calculated compared to the mean diversity in windows  $\geq 0.01\text{cM}$  or  $\geq 0.02\text{cM}$  from the nearest gene for pairwise diversity and singletons, respectively. Lines depict cubic smoothing splines with smoothing parameters chosen via generalized cross validation and shading depicts bootstrap-based 95% confidence intervals. Inset plots depict a smaller range on the x-axis.

cases diversity is not necessarily reduced [30, 31]. Such “soft sweeps” are still expected to occur more frequently in functional regions of the genome and could provide an alternative explanation for the observed reduction of diversity in genes. To test this possibility, we performed a genome-wide scan for selection using the H12 statistic, a method expected to be sensitive to both hard and soft sweeps [32]. Qualitative differences between maize and teosinte remained unchanged even after removing genes in the top 20% of H12 (Figure S6A). *This makes me wonder about the top 20%. Is the pattern of diversity reduction different in this fraction of the genome? If so, that seems like something that should be mentioned.* We interpret these combined results as suggesting that

purifying selection has left a more pronounced signature in the teosinte genome due to the increased efficacy of selection resulting from differences in effective population size.

**Demography of maize domestication.** To explore whether differences in the efficacy of purifying selection between maize and teosinte can be explained by demographic processes, we estimated the parameters of a simple domestication bottleneck model (Figure 4). The most likely model estimates an ancestral population mutation rate of  $\theta = 0.0147$  per bp, which translates to an effective population size of  $N_a \approx 123,000$  individuals. We estimate that the maize population split from teosinte  $\approx 15,000$  generations in the past, with an initial



**Fig. 4.** Parameter estimates for a simple bottleneck model of maize domestication. See methods for details.

size of only  $\approx 5\%$  of the ancestral  $N_a$ . After its split from teosinte, our model posits exponential population growth in maize, estimating a final modern effective population size of  $N_m \approx 370,000$ . Maize and teosinte continue to exchange migrants after the population split, with gene flow between the populations estimated at  $M_{tm} = 1.1 \times 10^{-5} \times N_a$  migrants per generation from teosinte to maize and  $M_{mt} = 1.4 \times 10^{-5} \times N_a$  migrants from maize to teosinte.

In addition to our simple bottleneck model, we investigated two alternative approaches for demographic inference. First, we took advantage of genotyping data from more than 4,000 maize landraces [33] to estimate the modern maize effective population size using low frequency variants informative of population expansion. This analysis yields a much higher estimate of the modern maize effective population size at  $N_m \approx 993,000$ . Finally, we applied a model-free coalescent approach [34] using a subset of our samples. Though this analysis suggests non-equilibrium dynamics for teosinte not included in our initial model, it is nonetheless broadly consistent, identifying a clear domestication bottleneck followed by rapid population expansion in maize to an extremely large extant size of  $\approx 10^9$  (Figure S2).

**Population expansion leads to stronger purifying selection in modern maize.** Motivated by the rapid post-domestication expansion of maize evident in our demographic analysis, we reasoned that low-frequency — and thus younger — polymorphisms might show patterns distinct from pairwise diversity. Singleton diversity around missense and synonymous substitutions (Figure S4) appears nearly identical to results from pairwise diversity (Figure 2), providing little support for a substantial increase in the number or strength of hard sweeps occurring in maize. In contrast, we observe a qualitative shift in the effects of purifying selection: singleton polymorphisms are more strongly reduced in and near genes in maize than in teosinte (Figure 3B), even after downsampling our maize data to account for differences in sample size. This relationship again remained after removing the 20% of genes with the highest H12 values (Figure S6). Finally, while direct comparison of pairwise and singleton diversity within taxa are consistent with at least some non-equilibrium dynamics in teosinte, as indicated by our coalescent-based demographic analysis, these too reveal much stronger differences in maize (Figure S5).

## Discussion

**Hard sweeps do not shape genome-wide diversity in maize.** Our findings demonstrate that classic hard selective sweeps have not contributed substantially to genome-wide patterns of diversity in maize, a result we show is robust to concerns about power due to the effects of background selection [25]. Although our approach ignores the potential for hard sweeps in noncoding regions of the genome, a growing body of evidence argues against hard sweeps as the prevalent mode of selection shaping maize variability. Among well-characterized domestication loci, only the gene *tga1* shows evidence of a hard sweep on a missense mutation [14], while several loci are consistent with “soft sweeps” from standing variation [35, 36] or multiple mutations [13]. Moreover, genome-wide studies of domestication [15], local adaptation [37] and modern breeding [38] all support the importance of standing variation as the primary source of adaptive variation. But Sho's paper and Tanja's paper suggest non-coding variants are important for adaptation...should we mention this caveat? Soft sweeps are expected to be common when  $2N_e\mu_b \geq 1$ , where  $\mu_b$  is the mutation rate of beneficial alleles with selection coefficient  $s_b$  [39]. Assuming a mutation rate of  $3 \times 10^{-8}$  [40] and that on the order of  $\approx 1 - 5\%$  of mutations are beneficial [41], this implies that soft sweeps should be common in both maize and teosinte for mutational targets  $>> 10kb$  — a plausible size for quantitative traits or for regulatory evolution targeting genes with large up- or down-stream control regions [35, e.g.]. Indeed, many adaptive traits in both maize CITE and teosinte CITE are highly quantitative cite Wallace for maize? weber and/or lauter for teosinte? and adaptation in both maize [15] and teosinte [42] has involved selection on regulatory variation. what % of the windows of top H12 score genome-wide occur outside of genes? is that a number to include here? additional consideration of non-genic regions in both the results and discussion would probably be a good idea given Tanja and Sho's results

**Demography of domestication.** Although a number of authors have investigated the demography of maize domestication [43–45], these efforts relied on data only from genic regions of the genome and made a number of limiting assumptions about the demographic model. keep an eye out for Brandon's ninjas We show that diversity within genes has been strongly reduced by the effects of linked selection, such that even synonymous polymorphisms in genes are not representative of diversity at unconstrained sites. weird to follow up a statement about incomplete demographic model with selection result Furthermore, by utilizing the full

joint SFS, we are able to estimate population growth, gene flow, and the strength of the domestication bottleneck without requiring assumptions about its duration.

One surprising result from our model is the estimated timing of domestication at  $\approx 15,000$  years before present. While this appears to conflict with archaeological [46] estimates, we instead argue this estimate reflects the fact that the genetic split between populations likely preceded anatomical changes that can be identified in the archaeological record. We also note that our result may also be inflated due to population structure, as our geographically diverse sample of teosinte may include populations diverged from those that gave rise to maize.

The estimated bottleneck of  $\approx 5\%$  of the ancestral teosinte population seems low given that maize landraces exhibit  $\approx 80\%$  of the diversity of teosinte [15], but our model suggests that the effects of the bottleneck on diversity are likely ameliorated by both gene flow and rapid population growth (Figure 4). Although we estimate that the modern effective size of maize is larger than teosinte, the small size of our sample reduces our power to identify the low frequency alleles most sensitive to rapid population growth [47], and our model is unable to incorporate growth faster than exponential. Both alternative approaches we employ estimate a much larger modern effective size of maize in the range of  $\approx 10^6 - 10^9$ , an order of magnitude or more than the current size of teosinte. Census data suggest these estimates are plausible: there are 47.9 million ha of open-pollinated maize in production [48], likely planted at a density of  $\approx 25,000$  individuals per hectare [49]. Assuming the effective size is only  $\approx 0.4\%$  of the census size (i.e. 1 ear for every 1000 male plants), this still implies a modern effective population size of more than four billion. While these genetic and census estimates are likely inaccurate, all of the evidence points to the fact that the effective size of modern maize is extremely large.

### Demography influences the efficiency of purifying selection.

section needs help – i'm not happy with the writing We find a more pronounced decrease in nucleotide diversity around genes in teosinte compared to domesticated maize (Figure 3A). This difference is likely due to the effects of purifying selection. Purifying selection is more efficient in larger populations [50], and our observation is thus consistent with our demographic model and previous work [43–45, 51] showing a smaller long-term effective population size in maize. To understand how recent population growth has changed the effects of linked selection, we also analyzed diversity of singleton polymorphisms. Under a simple neutral model, lower frequency polymorphisms are expected to be younger, and patterns of diversity at singleton SNPs should thus represent recent evolutionary history. Consistent with its larger current size and thus more efficient purifying selection, we see a stronger decline of singleton diversity around genes in maize (Figure 3B than teosinte).

I would like to see a supplemental figure or maybe just number here on the partial correlation in both teosinte and maize of diversity with recombination after accounting for gene density. do this in 500kb windows and then in 10kb windows. we should do the correlation with both pi and singletons. This let's us compare more explicitly with studies like C-D who evaluate overall patterns of linked selection. will be especially cool if we don't see a big difference in big windows but we do in 10kb windows – suggests C-D looked at wrong scale A consequence of the inefficient purifying selection that maize experienced during its bottleneck is likely that it harbors more weakly deleterious alleles segregating at intermediate frequency than does teosinte. This could be a part of the explanation of why maize inbreds have continued to improve over the past several decades [52]; if deleterious alleles tend to be recessive and are particularly frequent, they will have ample opportunities to display their phenotypes in inbreds. Our results also demonstrate that re-

cent purifying selection in maize has become much more effective, potentially explaining the ongoing improvement of these inbreds as maize lines are continuously selected. Additionally, the large  $N_e$  of modern maize compared to modern teosinte implies that for new mutations, selection will operate much more efficiently in maize. inbreds stuff is OK, but should be shortened. we are aiming at a broader audience that doesn't care about maize. what are more/broader. Chia, Rogers-Melnick, Mezmouk all looked at load. does this have implications for large effect load (i.e. should bneck have purged those?), for heterosis, architecture of quant. traits, etc.

The absence of evidence for a genome-wide impact of hard sweeps differs markedly from observations in *Drosophila* [53] and *Capsella* [54], but is consistent with data from humans [55, 56]. In a related study, [57] showed that selection tends to have an elevated impact on reducing diversity in species with large population size compared to small. This may explain why maize and humans, both bottlenecked species, show similar patterns while drosophila and capsella behave differently. However, we also do not observe abundant hard sweeps in teosinte, which calls this explanation into question.

these results have implications for lots of studies. recent demographic history often ignored (examples). in crops, bneck is considered and loss of diversity, but expansion also likely ubiquitous and consequences not well studied. in humans XYZ, and we see some of that already here. this argues that a full understanding of diversity in both wild and cultivated plants will require recent demography. mention C-D ignored recent demography – here's where rec/pi windows could help.

## Materials and Methods

**BASH, R, and Python scripts.** All scripts used for analysis are available in an online repository at [REPO ADDRESS HERE](#).

**Plant materials.** We made use of published sequences from inbred accessions of teosinte (*Z. mays* ssp. *parviglumis*) and maize landraces from the Maize HapMap3 panel as part of the Panzea project [16, 58, 59]. From these data, we removed 4 teosinte individuals that were not ssp. *parviglumis* or appeared as outliers in an initial principal component analysis conducted with the package adegenet [60] (Figure S7), leaving 13 teosinte and 23 maize that were used for all subsequent analyses (Table S8). We also utilized a single individual of (*Tripsacum dactyloides*) as an outgroup. All bam files are available at [/iplant/home/shared/panzea/hapmap3/bam\\_internal/v3.bams\\_bwamem](#). some clarification here. did we use Lemmon's higher depth teosinte? If so, why does Li say we only had 7 high-depth bam files? shouldn't there be more?

**Physical and genetic maps.** Sequences were mapped to the maize B73 version 3 reference genome [61] ([ftp://ftp.ensemblgenomes.org/pub/plants/release-22/fasta/zea\\_mays/dna/](ftp://ftp.ensemblgenomes.org/pub/plants/release-22/fasta/zea_mays/dna/)) as described by [59]. All analyses made use of uniquely mapping reads with mapping quality score  $\geq 30$  and bases with base quality score  $\geq 20$ ; quality scores around indels were adjusted following Li *et al.* [62]. We converted physical coordinates to genetic coordinates via linear interpolation of the previously published 1cM resolution NAM genetic map [63].

**Estimating the site frequency spectrum.** We estimated both the genome-wide site frequency spectrum (SFS) as well as a separate SFS for genic (within annotated transcript) and intergenic ( $\geq 5\text{kb}$  from a transcript) regions. We used the biomaRt package [64, 65] of R [66] to parse annotations from genebuild version 5b of AGPv3. We estimated single population and joint SFS with the software ANGSD [67], including all positions with at least one aligned read in  $\geq 80\%$  of samples in one or both populations. We assumed individuals were fully inbred and treated each line as a single haplotype. Because ANGSD cannot calculate a folded joint SFS, we first polarized SNPs using the maize reference genome and then folded spectra using  $\delta\alpha\delta i$  [2].

**Demographic inference.** We used the software  $\delta\alpha\delta i$  [2] to estimate parameters of a domestication bottleneck from the joint maize-teosinte SFS, using only sites  $> 5\text{kb}$  from a gene to ameliorate the effects of linked selection. We modeled a teosinte population of constant effective size  $N_a$ , that at time  $T_b$  generations in the past gave rise to a maize population of size  $N_b$  which grew exponentially to size  $N_m$  in the present (Figure 4). The model includes migration of  $M_{mt}$  individuals each generation from maize to teosinte and  $M_{tm}$  individuals from teosinte to maize. We estimated  $N_a$  using  $\delta\alpha\delta i$ 's estimation of  $\theta = 4N_a\mu$  from the data and a

mutation rate of  $\mu = 3 \times 10^{-8}$  [40]. We estimated all other parameters using 1,000  $\delta\alpha\delta i$  optimizations and allowing initial values between runs to be randomly perturbed by a factor of 2. Optimized parameters along with their initial values and upper and lower bounds can be found in table S9. We report parameter estimates from the optimization run with the highest log-likelihood.

We further made use of a large genotyping data set of more than 4,000 maize landraces [33] to estimate the modern maize  $N_e$  from singleton counts. We filtered these data to include only SNPs with data in  $\geq 1,500$  individuals, and then projected the SFS down to a sample of 500 individuals by sampling each marker without replacement 1,000 times according to the observed allele frequencies. We then estimated  $N_e$  from the data assuming  $\mu = 3 \times 10^{-8}$  [40] and the relation  $4N_e\mu = \frac{S}{L}$  [68], where where  $S$  is the total number of singleton SNPs and  $L$  is the total number of SNPs in the dataset.

As a final estimate of demography, we employed MSMC [34] to complement our model-based demographic inference. We used six each of maize and teosinte (BKN022, BKN025, BKN029, BKN030, BKN031, BKN033, TIL01, TIL03, TIL09, TIL10, TIL11 and TIL14), treating each inbred genome as a single haplotype. We called SNPs in ANGSD [67] using a SNP p-value of  $1e-6$  against a reference genome masked using SNPable (.). We then removed heterozygous genotypes and filtered sites with a mapping quality  $< 30$ , a base quality  $< 20$ , or a  $|\log_2(depth)| < 1$ . We ran MSMC with pattern parameters  $20 \times 2 + 20 \times 4 + 10 \times 2$ .

**Diversity.** We made use of the software ANGSD [67] for diversity calculations and genotype calling. We calculated diversity statistics in maize and teosinte in 1 kb non-overlapping windows using filters as described above for the SFS. We used allele counts to estimate the number of singleton polymorphisms in each window, and used binomial sampling to create a second maize data set down-sampled to have the same number of samples as teosinte. We called genotypes in maize, teosinte, and *Tripsacum* at sites with a SNP p-value  $< 10^{-6}$  and when the genotype posterior probability  $> 0.95$ . We identified substitutions in maize and teosinte as all sites with a fixed difference with *Tripsacum* and  $\leq 20\%$  missing data. Substitutions were classified as synonymous, missense, or noncoding using the ensemble variant effects predictor [69]. For each window with  $\geq 100bp$  of data we computed the genetic distance between the window center and the nearest synonymous and missense substitution as well as the genetic distance to the center of the nearest gene transcript.

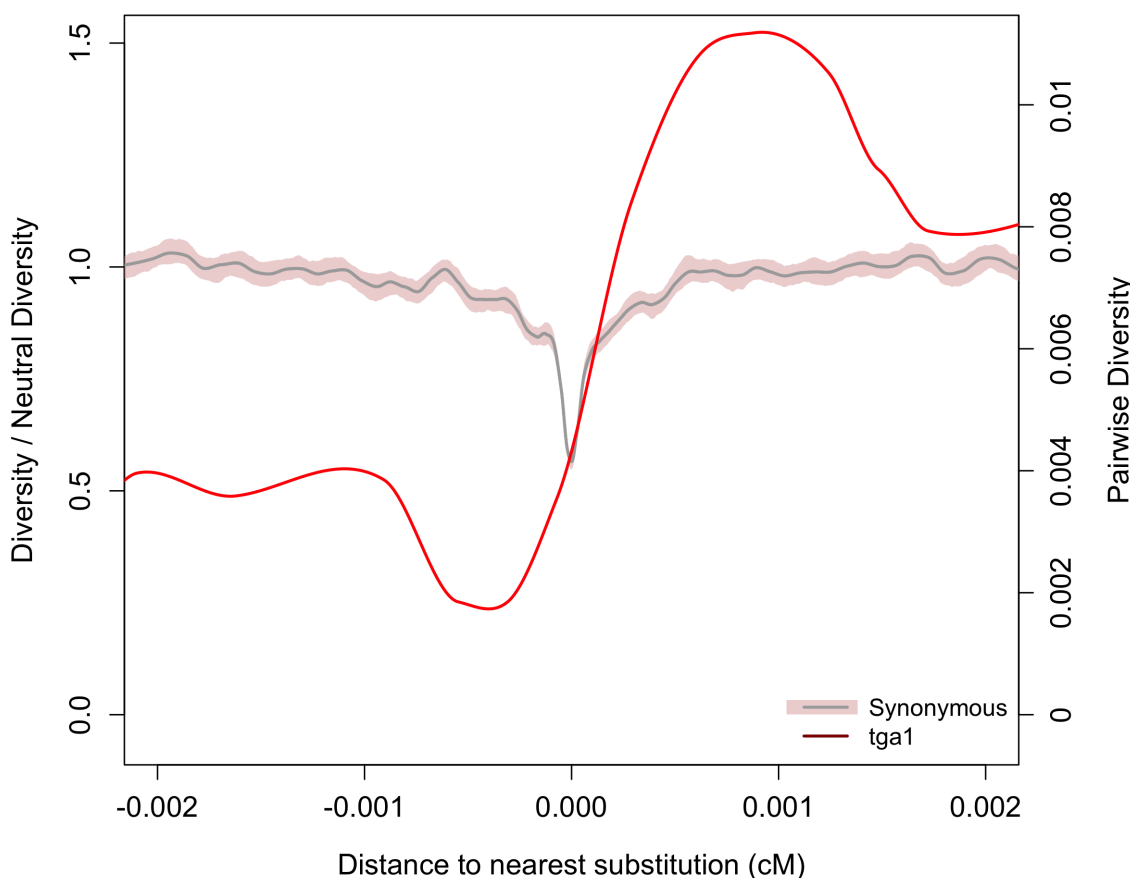
**Selection scan.** We scanned the genome to identify sites that have experienced recent positive selection using the H12 statistic [32] in sliding windows of 200 SNPs with a step of 25 SNPs.

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1. Luikart, G, England, P. R, Tallmon, D, Jordan, S, & Taberlet, P. (2003) *Nature Reviews Genetics* 4, 981–994.
2. Gutenkunst, R. N, Hernandez, R. D, Williamson, S. H, & Bustamante, C. D. (2009) *PLoS genetics* 5, e1000695.
3. Li, J, Li, H, Jakobsson, M, Li, S, Sjödin, P, & Lascoux, M. (2012) *Molecular Ecology* 21, 28–44.
4. Slotte, T. (2014) *Briefings in functional genomics* 13, 268–275.
5. Sella, G, Petrov, D. A, Przeworski, M, & Andolfatto, P. (2009) *PLoS genetics* 5, e1000495.
6. Gazave, E, Ma, L, Chang, D, Coventry, A, Gao, F, Muzny, D, Boerwinkle, E, Gibbs, R. A, Sing, C. F, Clark, A. G, et al. (2014) *Proceedings of the National Academy of Sciences* 111, 757–762.
7. Shifraim, B, Prasanna, B. M, Hellin, J, & Bänziger, M. (2011) *Food Security* 3, 307–327.
8. Strable, J & Scanlon, M. J. (2009) *Cold Spring Harbor Protocols* 2009, pdb-emo132.
9. Smith, B. D. (1995) *The emergence of agriculture*. (Scientific American Library New York).
10. Matsuoka, Y, Vigouroux, Y, Goodman, M. M, Sanchez, J, Buckler, E, & Doebley, J. (2002) *Proceedings of the National Academy of Sciences* 99, 6080–6084.
11. Wilkes, H. G et al. (1967) Teosinte: the closest relative of maize.
12. Doebley, J, Stec, A, & Gustus, C. (1995) *Genetics* 141, 333.
13. Wills, D. M, Whipple, C. J, Takuno, S, Kursel, L. E, Shannon, L. M, Ross-Ibarra, J, & Doebley, J. F. (2013) *PLoS Genet* 9, e1003604.
14. Wang, H, Studer, A. J, Zhao, Q, Meeley, R, & Doebley, J. F. (2015) *Genetics* pp. genetics–115.
15. Hufford, M. B, Xu, X, Van Heerwaarden, J, Pyhäjärvi, T, Chia, J.-M, Cartwright, R. A, Elshire, R. J, Glaubitz, J. C, Guill, K. E, Kaeplier, S. M, et al. (2012) *Nature genetics* 44, 808–811.
16. Chia, J.-M, Song, C, Bradbury, P. J, Costich, D, de Leon, N, Doebley, J, Elshire, R. J, Gaut, B, Geller, L, Glaubitz, J. C, et al. (2012) *Nature genetics* 44, 803–807.
17. Doebley, J. F, Gaut, B. S, & Smith, B. D. (2006) *Cell* 127, 1309–1321.
18. Purugganan, M. D & Fuller, D. Q. (2009) *Nature* 457, 843–848.
19. Ross-Ibarra, J, Morrell, P. L, & Gaut, B. S. (2007) *Proceedings of the National Academy of Sciences* 104, 8641–8648.
20. He, Z, Zhai, W, Wen, H, Tang, T, Wang, Y, Lu, X, Greenberg, A. J, Hudson, R. R, Wu, C.-I, & Shi, S. (2011) *PLoS genetics* 7, e1002100.
21. Vigouroux, Y, McMullen, M, Hittinger, C, Houchins, K, Schulz, L, Kresovich, S, Matsuoka, Y, & Doebley, J. (2002) *Proceedings of the National Academy of Sciences* 99, 9650–9655.
22. Chapman, M. A, Pashley, C. H, Wenzler, J, Hvala, J, Tang, S, Knapp, S. J, & Burke, J. M. (2008) *The Plant Cell* 20, 2931–2945.
23. Smith, J. M & Haigh, J. (1974) *Genetical research* 23, 23–35.
24. Wang, H, Nussbaum-Wagler, T, Li, B, Zhao, Q, Vigouroux, Y, Faller, M, Bomblies, K, Lukens, L, & Doebley, J. F. (2005) *Nature* 436, 714–719.
25. Enard, D, Messer, P. W, & Petrov, D. A. (2014) *Genome research* 24, 885–895.
26. Rodgers-Melnick, E, Bradbury, P. J, Elshire, R. J, Glaubitz, J. C, Acharya, C. B, Mitchell, S. E, Li, C, Li, Y, & Buckler, E. S. (2015) *Proceedings of the National Academy of Sciences* 112, 3823–3828.
27. Davydov, E. V, Goode, D. L, Sirota, M, Cooper, G. M, Sidow, A, & Batzoglou, S. (2010) *PLoS Comput Biol* 6, e1001025.
28. Charlesworth, B, Morgan, M, & Charlesworth, D. (1993) *Genetics* 134, 1289–1303.
29. Tenailion, M. I, Sawkins, M. C, Anderson, L. K, Stack, S. M, Doebley, J, & Gaut, B. S. (2002) *Genetics* 162, 1401–1413.
30. Innan, H & Kim, Y. (2004) *Proceedings of the National Academy of Sciences of the United States of America* 101, 10667–10672.
31. Messer, P. W & Petrov, D. A. (2013) *Trends in ecology & evolution* 28, 659–669.
32. Garud, N. R, Messer, P. W, Buzbas, E. O, & Petrov, D. A. (2015) *PLoS genetics* 11, e1005004.
33. Hearne, S, Chen, C, Buckler, E, & Mitchell, S. (2015) Unimputed gbs derived snps for maize landrace accessions represented in the seed-maize gwas panel (.). Accessed: 2015-02-16.
34. Schiffels, S & Durbin, R. (2014) *Nature genetics*.
35. Studer, A, Zhao, Q, Ross-Ibarra, J, & Doebley, J. (2011) *Nature genetics* 43, 1160–1163.
36. Gallavotti, A, Zhao, Q, Kyozuka, J, Meeley, R. B, Ritter, M. K, Doebley, J. F, Pè, M. E, & Schmidt, R. J. (2004) *Nature* 432, 630–635.
37. Takuno, S, Ralph, P, Swarts, K, Elshire, R. J, Glaubitz, J. C, Buckler, E. S, Hufford, M. B, & Ross-Ibarra, J. (2015) *Genetics*.
38. Beissinger, T. M, Hirsch, C. N, Vaillancourt, B, Deshpande, S, Barry, K, Buell, C. R, Kaeplier, S. M, Gianola, D, & de Leon, N. (2014) *Genetics* 196, 829–840.
39. Messer, P. W & Petrov, D. A. (2013) *Trends in ecology & evolution* 28, 659–669.
40. Clark, R. M, Tavaré, S, & Doebley, J. (2005) *Molecular biology and evolution* 22, 2304–2312.
41. Eyre-Walker, A & Keightley, P. D. (2007) *Nature Reviews Genetics* 8, 610–618.
42. Pyhäjärvi, T, Hufford, M. B, Mezmouk, S, & Ross-Ibarra, J. (2013) *Genome biology and evolution* 5, 1594–1609.
43. Eyre-Walker, A, Gaut, R. L, Hilton, H, Feldman, D. L, & Gaut, B. S. (1998) *Proceedings of the National Academy of Sciences* 95, 4441–4446.
44. Tenailion, M. I, U'Ren, J, Tenailion, O, & Gaut, B. S. (2004) *Molecular Biology and Evolution* 21, 1214–1225.
45. Wright, S. I, Bi, I. V, Schroeder, S. G, Yamasaki, M, Doebley, J. F, McMullen, M. D, & Gaut, B. S. (2005) *Science* 308, 1310–1314.
46. Piperno, D. R, Ranere, A. J, Holst, I, Iriarte, J, & Dickau, R. (2009) *Proceedings of the National Academy of Sciences* 106, 5019–5024.
47. Keinan, A & Clark, A. G. (2012) *science* 336, 740–743.
48. Program, T. M. (1999) Development, maintenance, and seed multiplication of open-pollinated maize varieties. (CIMMYT, Mexico, D.F.), 2 edition.
49. Baden, W. W & Beekman, C. S. (2001) *American Antiquity* pp. 505–515.
50. Kimura, M. (1984) *The neutral theory of molecular evolution*. (Cambridge University Press).
51. Ross-Ibarra, J, Tenailion, M, & Gaut, B. S. (2009) *Genetics* 181, 1399–1413.
52. Meghji, M, Dudley, J, Lambert, R, & Sprague, G. (1984) *Crop Science* 24, 545–549.
53. Sattath, S, Elyashiv, E, Kolodny, O, Rinott, Y, & Sella, G. (2011) *PLoS genetics* 7, e1001302.
54. Williamson, R, Josephs, E, Platts, A, Hazzouri, K, Haudry, A, Blanchette, M, & Wright, S. (2014) *PLoS genetics* 10, e1004622–e1004622.
55. Hernandez, R. D, Kelley, J. L, Elyashiv, E, Melton, S. C, Auton, A, McVean, G, Sella, G, Przeworski, M, et al. (2011) *science* 331, 920–924.

56. Pritchard, J. K., Pickrell, J. K., & Coop, G. (2010) *Current Biology* 20, R208–R215.
57. Corbett-Detig, R. B., Hartl, D. L., & Sackton, T. B. (2015) *PLoS Biol* 13, e1002112.
58. Lemmon, Z. H., Bukowski, R., Sun, Q., & Doebley, J. F. (2014) *PLoS Genet* 10, e1004745.
59. Panzea. (In prep) TBD.
60. Jombart, T & Ahmed, I. (2011) *Bioinformatics* 27, 3070–3071.
61. Schnable, P. S., Ware, D., Fulton, R. S., Stein, J. C., Wei, F., Pasternak, S., Liang, C., Zhang, J., Fulton, L., Graves, T. A., et al. (2009) *science* 326, 1112–1115.
62. Li, H. (2011) *Bioinformatics* 27, 2987–2993.
63. Glaubitz, J. C., Casstevens, T. M., Lu, F., Harriman, J., Elshire, R. J., Sun, Q., & Buckler, E. S. (2014) *PLoS One* 9, E90346.
64. Durinck, S., Spellman, P. T., Birney, E., & Huber, W. (2009) *Nature protocols* 4, 1184–1191.
65. Durinck, S., Moreau, Y., Kasprzyk, A., Davis, S., De Moor, B., Brazma, A., & Huber, W. (2005) *Bioinformatics* 21, 3439–3440.
66. R Core Team. (2014) R: A Language and Environment for Statistical Computing (R Foundation for Statistical Computing, Vienna, Austria).
67. Korneliussen, T. S., Albrechtsen, A., & Nielsen, R. (2014) *BMC bioinformatics* 15, 356.
68. Fu, Y.-X & Li, W.-H. (1993) *Genetics* 133, 693–709.
69. McLaren, W., Pritchard, B., Rios, D., Chen, Y., Flieck, P., & Cunningham, F. (2010) *Bioinformatics* 26, 2069–2070.

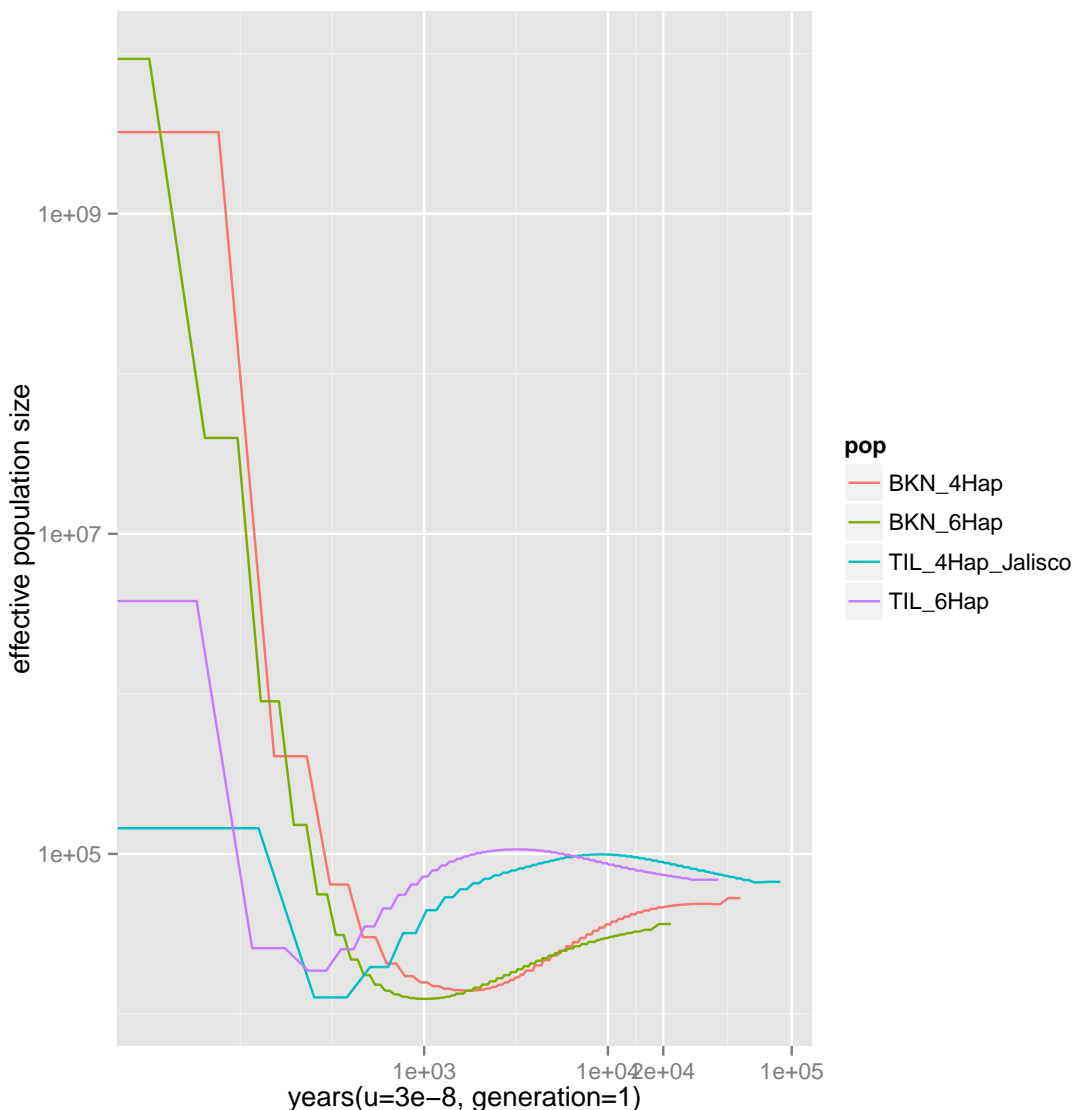




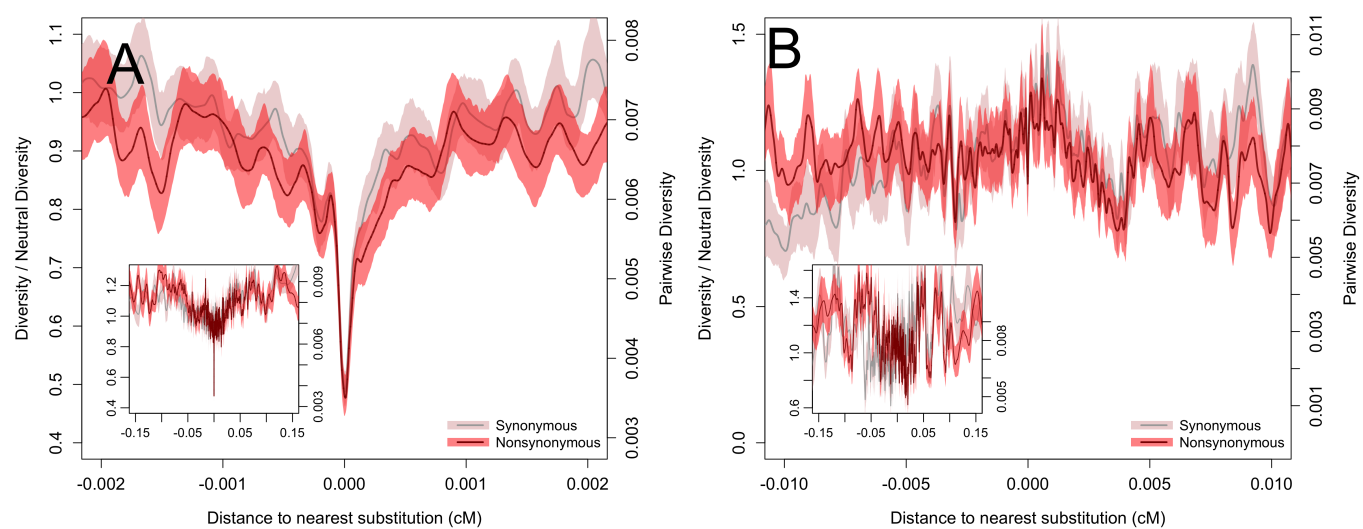
**Fig. S1.** Diversity surrounding the causitive polymorphism at the *tga1* locus is plotted. Since this is only one gene, the large amount of noise compared to our average plots is expected. However, notice that diversity precisely at the causitive polymorphism is reduced and a recovery of diversity is observed away from that site.

#### Supporting Information

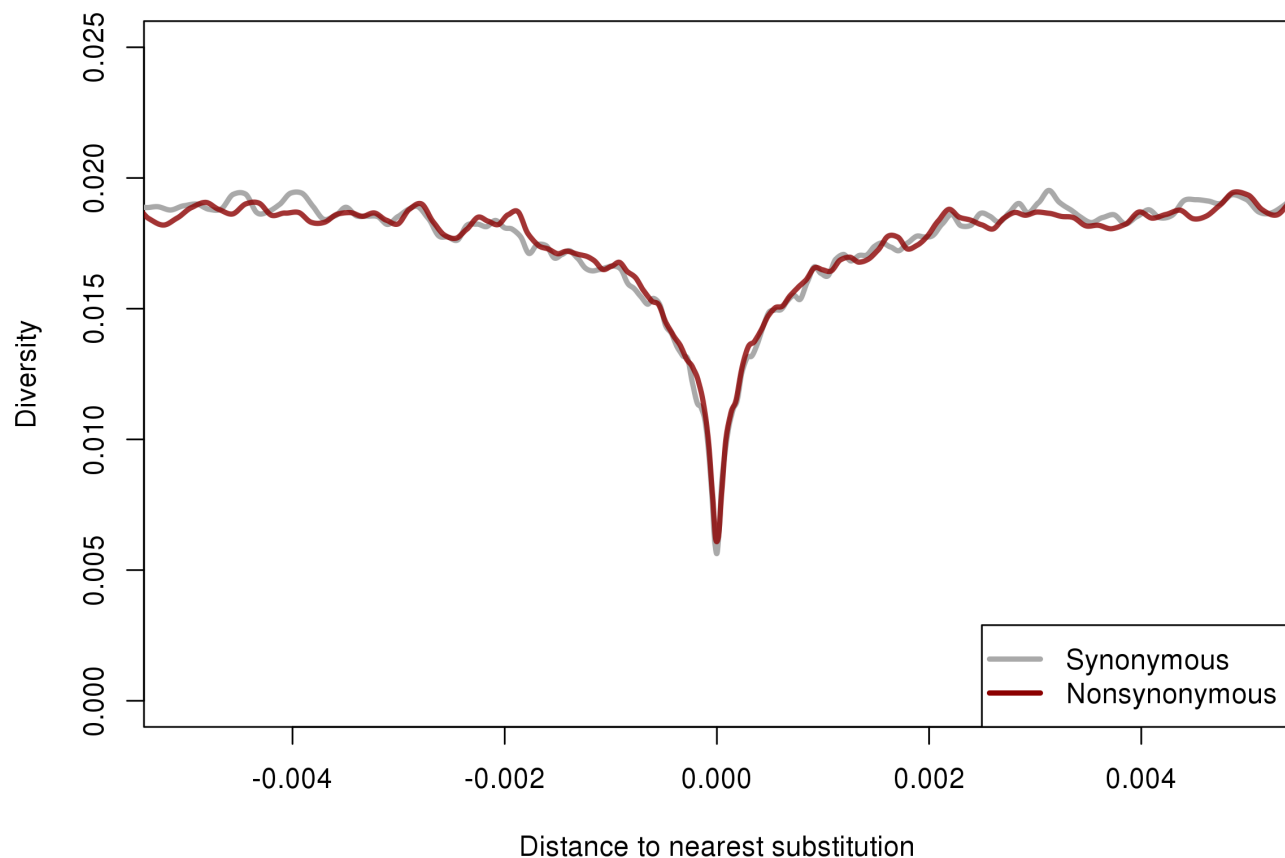




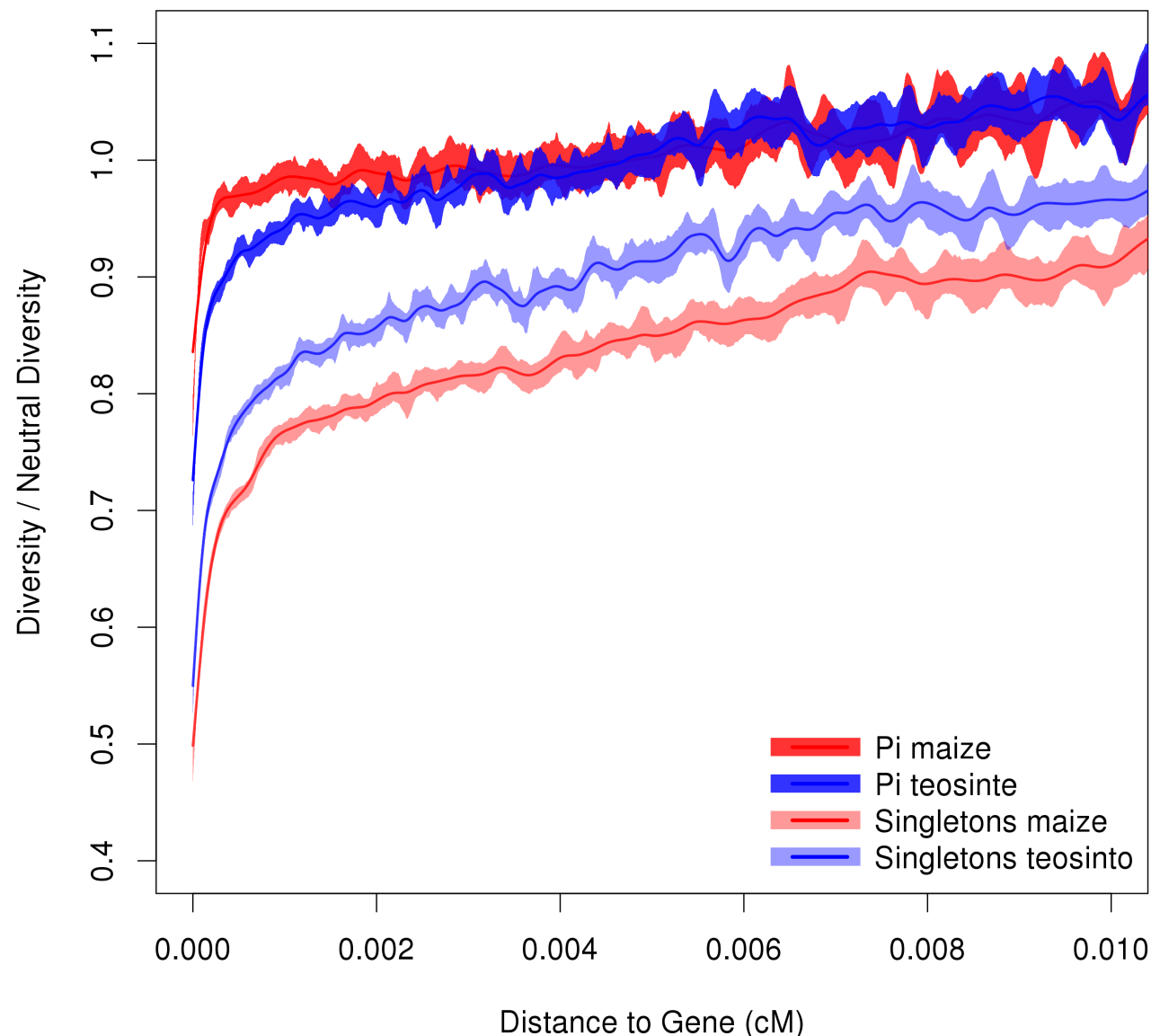
**Fig. S2.** need MSMC caption



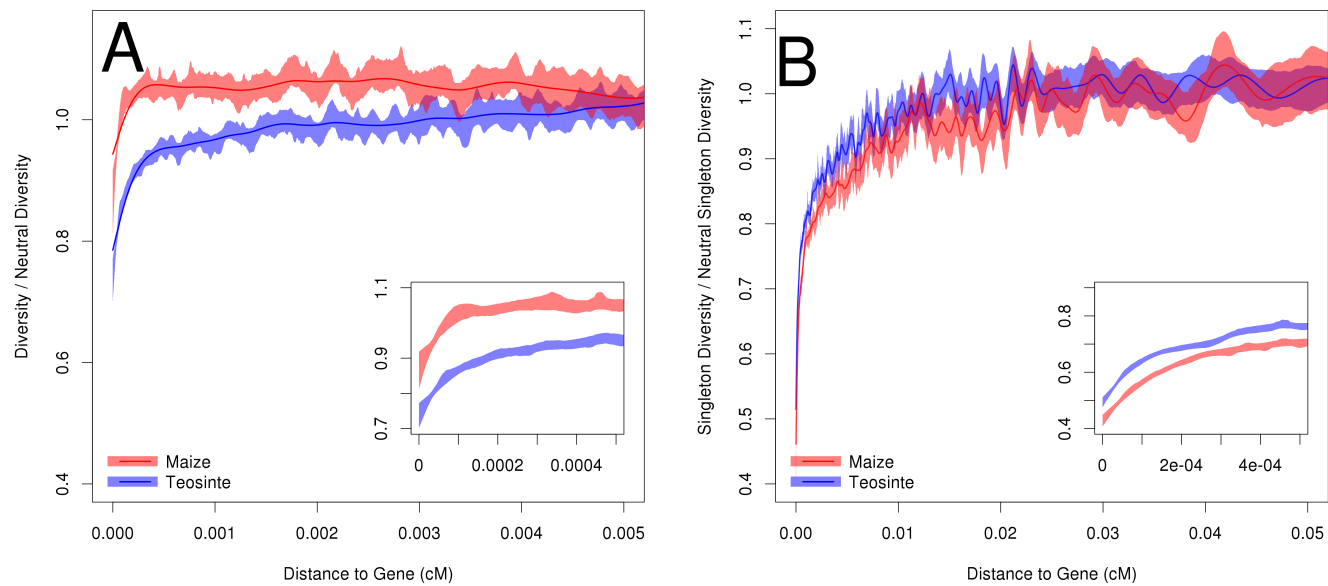
**Fig. S3.** Pairwise diversity surrounding synonymous and nonsynonymous substitutions in maize at highly conserved (A) or unconserved (B) sites. Bootstrap-based 95% confidence intervals are depicted via shading. Inset plots depict a larger range on the x-axis.



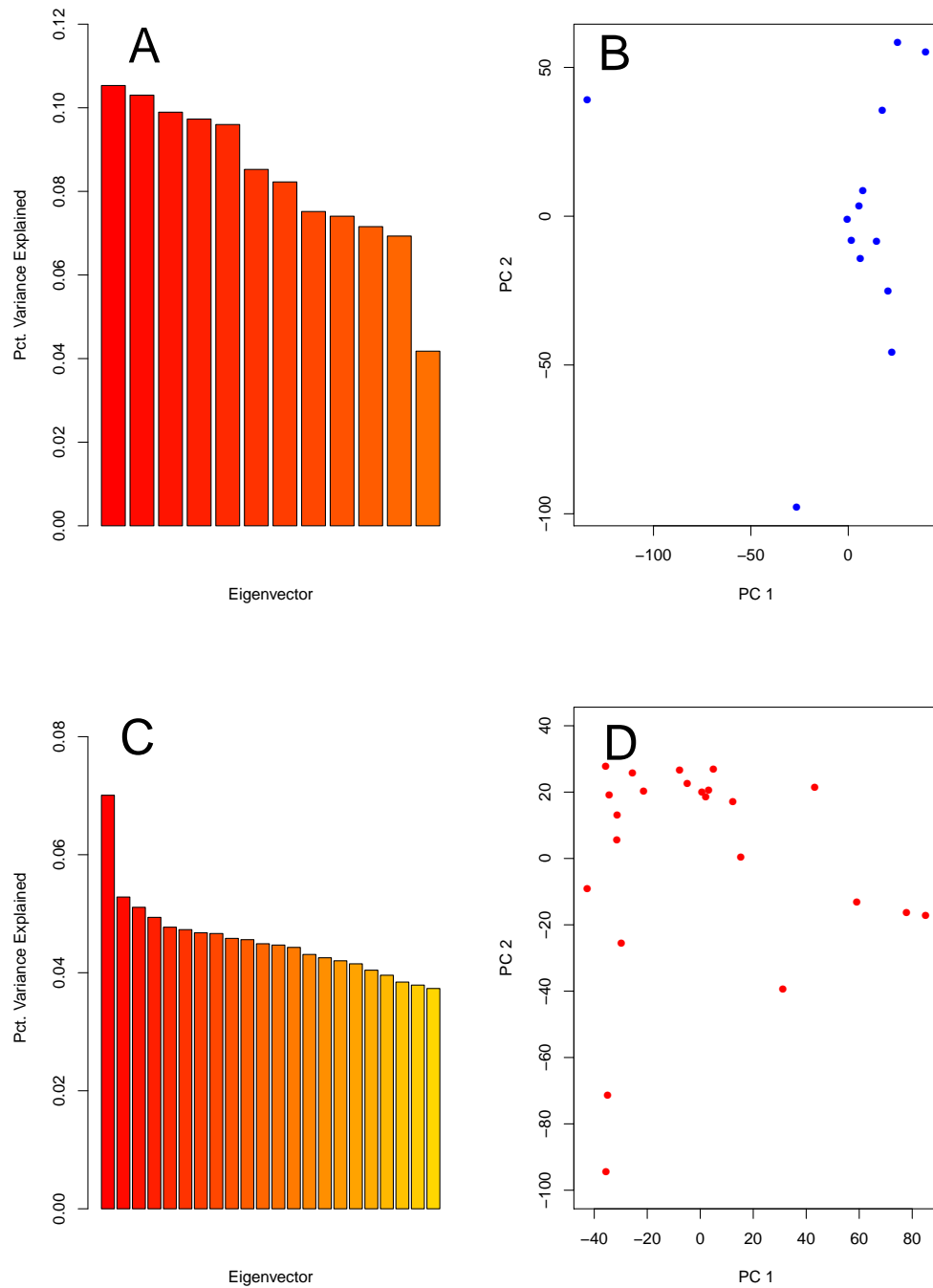
**Fig. S4.** Singleton diversity surrounding synonymous and nonsynonymous substitutions in maize.



**Fig. S5.** Relative diversity versus distance to nearest gene in maize and teosinte. Relative diversity is calculated by comparing to the mean diversity in all windows  $\geq 0.02\text{cM}$  from the nearest gene. Lines depict cubic smoothing splines with smoothing parameters chosen via generalized cross validation and shading depicts bootstrap-based 95% confidence intervals.



**Fig. S6.** Relative level of diversity versus distance to the nearest gene, in maize and teosinte, based on only sites that do not show evidence of hard or soft sweeps according to H12. Two measures of diversity were investigated. **A** displays pairwise diversity, which is most influenced by intermediate frequency alleles and therefore depicts more ancient evolutionary patterns, and **B** depicts singleton diversity, influenced by rare alleles and thus depicting evolutionary patterns in the recent past. Bootstrap-based 95% confidence intervals are depicted via shading. Inset plots depict a smaller range on the x-axis.



**Fig. S7.** Principal component analysis of teosinte and maize individuals to ensure that no close relatives were inadvertently included in our study. Plots are based on a random sample of 10,000 SNPs. **A:** Percentage of total variance explained by each principal component for teosinte. **B:** PC1 vs PC2 for all 13 teosinte individuals. **C:** Percentage of total variance explained by each principal component for maize. **D:** PC1 vs PC2 for all 23 maize individuals.

Maize	Teosinte
BKN009	TIL01
BKN010	TIL02
BKN011	TIL03
BKN014	TIL04-TIP454
BKN015	TIL07
BKN016	TIL09
BKN017	TIL10
BKN018	TIL11
BKN019	TIL12
BKN020	TIL14-TIP498
BKN022	TIL15
BKN023	TIL16
BKN025	TIL17
BKN026	
BKN027	
BKN029	
BKN030	
BKN031	
BKN032	
BKN033	
BKN034	
BKN035	
BKN040	

**Fig. S8.** A list of maize and teosinte individuals included in this study. Sequencing and details were previously described by [cite chia and lemmmon](#)

Parameter	Initial value	Upper bound	Lower bound
$\frac{N_b}{N_a}$	0.02	$1 \times 10^{-7}$	2
$\frac{N_m}{N_a}$	3	$1 \times 10^{-7}$	200
$\frac{T_b}{2N_a}$	0.04	0	1
$\frac{M_{mt}}{N_a}$	$1 \times 10^{-10}$	$1 \times 10^{-7}$	0.001
$\frac{M_{tm}}{N_a}$	$1 \times 10^{-10}$	$1 \times 10^{-7}$	0.001

**Fig. S9.** Parameters, initial values, and boundaries used for model-fitting with  $\delta\alpha\delta i$ . Parameters are shown in the units utilized by  $\delta\alpha\delta i$ , although in the text simplified units are reported.

