

# Whole-genome duplication increases genetic diversity and load in outcrossing Arabidopsis arenosa

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Genetic variation underpins evolutionary change, but mutation accumulation increases genetic load. Various factors affect the extent of load, such as population size and breeding system, but other important determinants remain unexplored. In particular, whole-genome duplication (WGD)—a pervasive macromutation occurring broadly across Eukaryotes-remains poorly understood in terms of its impact on neutral and selective processes within populations. Using iterative forward simulations and empirical analysis of 632 short- and 16 long-read sequenced individuals of Arabidopsis arenosa (in 23 diploid and 42 natural autotetraploid populations), we measure the effects of WGD on genome-wide diversity and mutation load. Our simulations show how genetic variation gradually rises in autotetraploids due to increased mutational target size. Moreover, mutation load increases due to relaxed purifying selection as ploidies rise, when deleterious mutations are masked by additional chromosome copies. Empirical data confirm these patterns, showing significant increases in nucleotide diversity, ratios of nonsynonymous to synonymous SNPs, and numbers of indels and large structural variants in A. arenosa autotetraploids. However, a rather modest increase in load proxies together with a broad distribution and niche of autotetraploids suggests load accumulation has not yet limited their successful expansion. Overall, we demonstrate a complex interplay between neutral processes and purifying selection in shaping genetic variation following WGD and highlight ploidy as an important determinant of mutation load, genetic diversity, and therefore adaptive potential in natural populations.

Arabidopsis | evolution | genomics | genetic load | natural selection

Genetic diversity is both a result and a determinant of evolutionary change. While some variation mediates adaptations, most mutations are at least mildly deleterious and often disrupt networks that have been fine-tuned over the course of evolution (1, 2). The reduction in fitness of a population, particularly due to recurrent deleterious mutations, is referred to as mutation load (3-6). While DNA mutagenesis increases mutation load, purifying selection reduces it by removing deleterious alleles. Understanding the factors that influence the efficiency of purifying selection—and thus control of mutation load—is foundational for our ability to assess the adaptive potential of natural populations. This has direct relevance to adaptation to environmental challenges (7), agriculture (8), conservation (9-11), and human health (12, 13).

Both theoretical and empirical studies have revealed key determinants of mutation load in natural populations. Population size and the degree of inbreeding are particularly strong factors (6, 14, 15), whose effect in empirical data, however, differ depending on what load component is measured. Realized and masked load reflect whether fitness effects of deleterious alleles are expressed in the current generation or lay hidden in a heterozygous state (as recessive alleles) (6). If deleterious mutations are recessive or partially recessive, masked load increases linearly with population size and realized load drops as deleterious mutations accumulate in a heterozygous state (6). Inbreeding on the other hand unmasks recessive deleterious mutations, increasing realized load. Empirical studies support the notion that various demographic changes modulate selection efficiency and mutation load (15-18). Breeding system is yet another factor that shapes levels of homozygosity and therefore the amount of mutation load (19, 20). Recombination rate shapes mutation load as higher recombination allows more efficient selection (21, 22). And finally, mutation load is determined by the mutation rate itself (23). In sum, recent studies demonstrate an array of population genetic processes affecting mutation load; yet we still lack a comprehensive view of the spectrum of determinants of mutation load in natural populations.

Whole genome duplication (WGD) is a transformative mutation that substantially affects inheritance mode and effective population sizes, which in turn shape genetic

## **Significance**

Our study shows how wholegenome duplication (WGD), a massive and naturally widespread macromutation, increases population genetic variation both through point mutations and via large structural changes in the genome. Specifically, WGD doubles the number of chromosome copies where mutations may occur but also masks their detrimental effects, shielding them by masking via more alleles at each locus. Since genomic variation necessarily underpins evolution, understanding the role of WGD in affecting evolutionary processes gives general insight into mechanisms of adaptation. Given that polyploidy is common in plants, particularly in crops, this research has practical implications for crop breeding and resilience, addressing fundamental questions about the interplay of genome dynamics and limits to adaptability.

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variation and mutation load. WGD has an essential role in the evolution of eukaryotes (24) and is widespread in plants, particularly in crops (25, 26), yet its effect on mutation load has not been fully addressed. Here, we fill in this gap by deconstructing the direct effects of WGD in intraspecific polyploids, autopolyploids, where natural WGD has not been confounded by hybridization on different predictors of mutation load. Theory predicts polysomic inheritance, where all homologous chromosomes recombine without pairing partner preferences, modulates several genetic processes in autopolyploids (27). After one round of WGD, the mutation rate per gene is expected to double (25, 28), while the rate of genetic drift is halved (29). Accordingly, genetic variation in terms of nucleotide diversity ( $\pi$ ) is expected to double (4Ne $\mu$ vs. 8Neµ) (27) if we assume equal population census size. Purifying selection is thought to be weaker in polyploids, predicting an increase in the number and frequency of deleterious alleles (30). This is due to masking of recessive or partly recessive alleles by additional sets of chromosomes.

Despite these clear expectations, empirical data on genetic diversity in autopolyploids are not straightforward. First, genetic diversity has only occasionally been found to be higher (but never doubled) in autotetraploids compared to diploids (31–33). Despite this, increased variation of evolutionarily constrained (genic) regions has been found in autotetraploids for single nucleotide polymorphisms (SNPs) (34, 35), transposable elements (TE) polymorphisms (36) and structural variants (SVs) (37), indicating greater accumulation of mutation load in tetraploid populations compared to diploid counterparts. Nonetheless, estimating load directly from genetic diversity is challenging (6, 38) and we lack a truly integrative analysis of mutation load and its components over many types of genetic markers and load indices in a natural polyploid system.

Here, we provide such an analysis with a mix of mutation load analyses that account for differences in demography between cytotypes and forward-in-time simulations in a wider context of realistic, nonequilibrium evolutionary states. First, we use forward-in-time simulations parameterized by values guided by our Arabidopsis arenosa dataset to generate relevant predictions for the dynamics of genomic variation and mutation load as a function of WGD. Then, we analyze 65 short-read sequenced populations and 16 long-read sequenced individuals of A. arenosa to empirically test these predictions. A. arenosa encompasses diploids and naturally established autotetraploids that were formed by a single WGD event between 19,000 to 30,000 generations ago (34, 39). Its outcrossing breeding system, large and demographically stable populations, and excellent background knowledge of ploidy distribution, broad environmental variation, and evolutionary history allow for testing general hypotheses on the drivers of genomic variation in natural conditions (34, 39-45). In particular, we test the hypothesis that tetraploid populations accumulate more genetic variation and mutation load. By addressing this hypothesis across SNPs, indels, and SVs, we provide robust evidence for the importance of WGD on accumulation of genetic variation, reflecting both an increase in mutational target size and a decrease in selection efficiency.

#### **Results**

Forward-In-Time Simulations Show Pervasive Effects of WGD on Genetic Diversity and Mutation Load. By simulating diploid and autotetraploid (hereafter simply, "tetraploid") populations of the same maximum population census size (carrying capacity  $\sim$ 180,000 individuals, parameterized with our empirical data; see *Methods*), we observed that neutral diversity ( $\pi_{Neutral}$ ) slowly

increases after a diploid population at equilibrium goes through a WGD. However, the increase is gradual (Fig. 1A) and  $\pi_{Neutral}$ in tetraploids does not reach the expected double values of diploids within even half a million generations, i.e., more than an order of magnitude higher than the estimated age of A. arenosa tetraploids (SI Appendix, Fig. S1). This observation corresponds well with analytical predictions on genetic diversity under the coalescent, which generally expect that equilibrium is reached at timescales that are an order of magnitude greater than is the time scale of our simulations (46, 47). Although  $\pi_{Neutral}$  estimates are consistently higher without background selection (reduction of neutral diversity due to purging of linked deleterious variants), the trajectory is very similar when background selection is included (SI Appendix, Fig. S1), suggesting that the overall increase in nucleotide diversity primarily reflects an increase in mutational target size after WGD (doubled number of chromosomes), rather than weakened purifying selection (due to polysomic masking). We did not assume an initial bottleneck following the formation of the tetraploid lineage, as there were no traces of past population size change in A. arenosa, likely due to gradual circa- and post-WGD influx of additional variation from the founding diploid population in the area where the tetraploid originated (34, 39). Yet, even when we assumed such a bottleneck, the overall patterns were similar, although  $\pi_{Neutral}$  of tetraploids drops immediately after the WGD and reaches the value of diploids only after ~20,000 simulated generations (SI Appendix, Fig. S2).

The accumulation of mutation load in tetraploids is also a gradual process. Two indices of mutation load-the ratio of deleterious to neutral nucleotide diversity and the ratio of strongly deleterious to nearly neutral polymorphisms-indicate that tetraploids accumulate a higher mutation load than diploids; this higher load in tetraploids becomes pronounced shortly after WGD (Fig. 1 B and C). This result is robust to demographic effects, as we see similar results if we assume a founding bottleneck at tetraploid formation (SI Appendix, Fig. S2). Unsurprisingly, masking of deleterious mutations after WGD provides immediate fitness benefits for the nascent polyploids (Fig. 1D); however, these benefits are highly transient. As deleterious mutations accumulate, the fitness of the tetraploid population eventually falls below its diploid progenitor (SI Appendix, Fig. S3). These results are robust to different assumptions about allelic dominance (Methods and SI Appendix, Figs. S13–S17)

In summary, our forward simulations show that WGD alters both neutral processes and the impact of purifying selection which should leave detectable footprints on genome-wide diversity in empirical datasets of diploid and autotetraploid populations.

Genetic Diversity and Population Structure in A. arenosa. To decipher how WGD modulates genetic diversity and mutation load in natural populations, we assembled a range-wide sequencing dataset of 632 (222 diploid and 410 tetraploid) A. arenosa individuals. We called genotypes in all accessions with available short-read sequencing data and included 126 newly sequenced individuals covering previously underrepresented geographic regions or genetic lineages, resulting in 65 populations with at least 6 individuals sequenced (Fig. 2A and SI Appendix, Table S1; average depth of coverage 24x). The structure of sampled populations corresponds with previous studies in the species (34, 39, 40, 42, 44): Neighbor-joining tree (Fig. 2D) and PCA (SI Appendix, Fig. S4) analyses show six distinct diploid lineages corresponding to those identified previously in a rangewide sampling (40). Tetraploids form one dominant cluster close to their ancestral diploids from the Western Carpathians (39), and three smaller clusters that show affinities to their sympatric

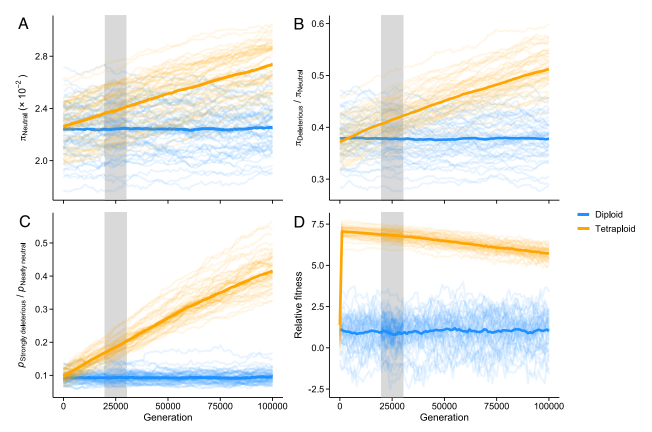


Fig. 1. Forward simulation of genetic diversity and mutation load demonstrates pervasive effects of ploidy. Diploid (blue) and tetraploid (orange) populations during 100,000 generations following WGD. The carrying capacity of both cytotypes was the same and we assumed no founder bottleneck event in tetraploids (for bottleneck scenarios see SI Appendix, Fig. S2). (A) Neutral synonymous nucleotide diversity. (Β) Ratio of nonsynonymous to neutral nucleotide diversity (π<sub>Deleterious</sub>)  $\pi_{\text{Neutral}}$ ). (C) Ratio of strongly deleterious to nearly neutral polymorphisms. (D) Average fitness of a population estimated as the product of fitness effects of all mutations. Fitness estimates were standardized based on values in diploids (mean = 1, SD = 1). In all panels, the solid lines show averages across 100 simulated repeats, while each of the simulation replicates are shown in transparent colors in the background. The gray area highlights the interval of estimated age of the extant A. arenosa tetraploids from Monnahan et al. (34).

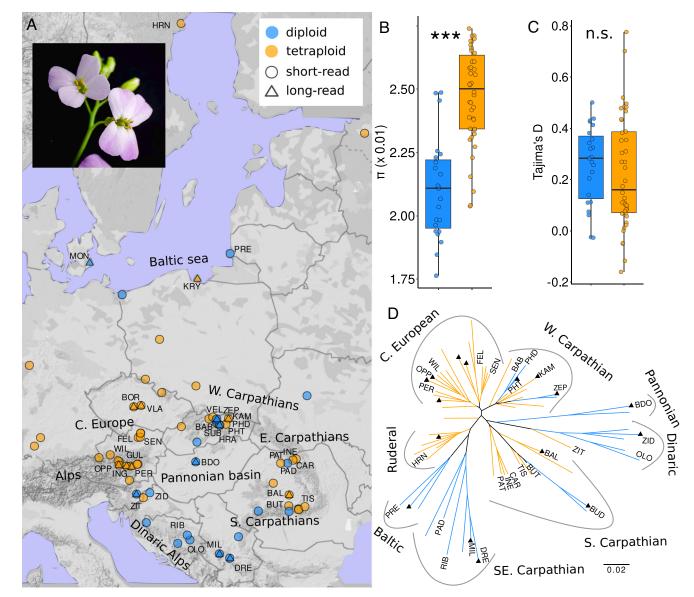
diploid lineages, in line with the documented interploidy gene flow in these areas (34).

Using populations as replicate observations, downsampled to the same number of individuals, we see that nucleotide diversity of putatively neutral fourfold degenerate SNPs ( $\pi_s$ ) is 1.2 times higher in tetraploids (0.025) compared to diploids (0.021) on average (Fig. 2B, DF = 62,  $\beta_{ploidy}$  = 3.9e<sup>-3</sup>,  $p_{ploidy}$  < 0.001, general linear model also accounting for an effect of sequencing depth, SI Appendix, Table S2). The effect of ploidy on diversity is also robust to sampling, remaining statistically significant if we sample the same number of chromosomes (not individuals) per population, or exclude admixed tetraploid populations from ploidy contact zones affected by interploidy gene flow (SI Appendix, Table S2).

The observed difference in neutral diversity can be primarily attributed to the increase of population-scaled mutation rate in tetraploids (8Neµ) compared to diploids (4Neµ). Additional confounding influences of demographic processes within populations seem unlikely in A. arenosa, as we found no difference in Tajima's D between ploidies (Fig. 2C; Wilcoxon rank-sum test of population means, W = 544, P = 0.409). Theoretically, increased mutation rate  $\mu$  in tetraploids could have effects, but estimates of diploid and tetraploid germline mutation rate are not available for A. arenosa, nor is there reason to expect changes in the underlying mutation rate. Based on the insights from our simulations, we speculate that diversity in tetraploids is far less than double that in diploids (only 1.19x), because the relatively recent tetraploid cytotype has not yet reached mutation-drift equilibrium (Fig. 1A).

Signatures of Relaxed Purifying Selection in Tetraploid A. *arenosa* in SNP Data. We next examined whether relaxed purifying selection due to polysomic masking contributes to increased genetic variation upon WGD. To test for a difference in selection efficiency, we compared selectively constrained zerofold degenerate SNPs (where any nucleotide change results in an amino acid replacement), and putatively neutral fourfold degenerate SNPs (where any nucleotide change is permitted without altering the encoded amino acid). To avoid potential effects of sequencing errors, we focused on a subset of 27 deeply sequenced populations, hereafter called core dataset (average sequencing depth of fourfold sites 35× for diploids and 38× for tetraploids (SI Appendix, Figs. S5 and S6); see Methods for details). Site frequency spectra (SFS) for both diploid and tetraploid populations showed an increased proportion of singleton zerofold variants compared to fourfold variants, which indicates that zerofold sites are under purifying selection (Fig. 3 A and B). To test whether purifying selection is relaxed in tetraploids, we used a ratio of overall zero- to fourfold diversity  $(\pi_0/\pi_4)$ , which serves as a proxy for the efficiency of selection relative to drift. This ratio is significantly higher in tetraploid populations than diploids (Fig. 3C, Wilcoxon rank-sum test: W = 24, P = 0.001), in line with the expectation of higher mutation load in tetraploids.

We then asked whether the cytotypes differ in their distribution of fitness effects of new mutations (DFE). The DFE is usually considered to be primarily a species-level trait (48), but WGD leads to major changes in the dominance mode (28), and so may affect DFE via modulation of selection efficiency also within a species. We inferred deleterious DFE from polymorphism in our



**Fig. 2.** Range-wide sampling, genetic diversity, and structure of *A. arenosa* populations. Diploid (blue) and tetraploid (orange) populations sampled throughout the species' native range in Central and Southeastern Europe. (*A*) Locations of the full dataset of 65 populations sequenced for short reads (circles) and 16 individuals sequenced for long reads (triangles). The inset shows flowering *A. arenosa* from W. Carpathians (photo J. Martínek). (*B* and *C*) Within-population pairwise nucleotide diversity  $\pi$  (*B*) and Tajima's D (*C*) calculated based on fourfold degenerate SNPs in all populations (dots) downsampled to the same number of individuals (N = 6). Asterisks indicate significant difference between the ploidies as tested by the Wilcoxon rank sum test. (*D*) Genetic relationships of 65 populations depicted by a neighbor-joining tree based on Nei's distances derived from 92,632 fourfold degenerate SNPs. Deeply sequenced populations from the "core dataset" are labeled with three-letter codes and populations with long read data are indicated by triangles.

core dataset by polyDFE (49). The deleterious DFE statistically differs between diploid and tetraploid populations (PERMANOVA:  $R^2 = 0.174$ , F = 5.29, P = 0.024). Tetraploids showed a lower proportion of novel mutations with a likely highly deleterious effect. They also exhibited higher proportions of novel mutations with inferred moderately deleterious effects (Fig. 3*D*).

DFE inference is based on fitting a model that explains differences between zerofold and putatively neutral fourfold SFS within a population (48, 50, 51). Because the model assumes additivity, it does not account for the fact that, in tetraploids, putatively deleterious zerofold variation may reach higher frequencies due to polysomic masking. This pattern in the SFS likely results in an apparently less deleterious DFE in tetraploids even if the real DFE is the same across ploidies. This interpretation is supported by the fact that DFE rarely changes within an outcrossing species (52, 53); however, at this stage we cannot conclusively differentiate between the effect of polysomic masking on DFE inference and ploidy-related change in actual DFE.

Consistent Signatures of Relaxed Purifying Selection in Tetraploids Across All Variant Types. Finally, we explored whether the relaxed purifying selection suggested for SNPs is also reflected in short indels and larger SVs in our core short-read dataset (indels) and long-read sequenced individuals (SVs). We estimated mutation load based on variant counts, which can be applied consistently across all these data types. Concordant with neutral nucleotide polymorphism values, the total number of derived SNPs, indels, and SVs was significantly higher in tetraploid compared to diploid populations. This held true both when we accounted for the same number of individuals or chromosomes (SI Appendix, Table S3). We thus asked whether it is only due to doubling of theta (4Neµ vs. 8Neµ) or also due to modulation of selection efficiency in coding regions. To test this, we compared putatively neutral and deleterious categories of each variant type. In all cases, we see a positive linear correlation between the number of deleterious variants and putatively neutral variants across populations (Fig. 3).

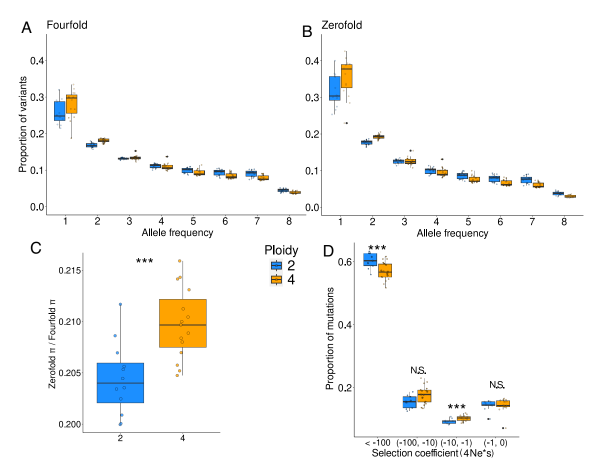


Fig. 3. Allele frequency spectra of both cytotypes and signal of weaker purifying selection efficiency in tetraploid A. arenosa populations. Folded SFS for (A) putatively neutral fourfold and (B) amino acid-changing zerofold SNP categories. The proportions are summarized over values inferred in our core dataset of 27 populations (dots), downsampled to the same number of allele copies. (C) Ratio of zero- to fourfold genetic diversity as a measure of selection efficiency in diploid and tetraploid populations (dots) downsampled to the same number of individuals (N = 6). (D) Discretized distribution of deleterious fitness effects of new mutations (DFE) estimated by polyDFE. Scaled selection coefficient of new mutations (4Ne\*s) is binned into four categories from which "<-100" stands for highly deleterious mutations while "(-1,0)" denotes effectively neutral mutations. The deleterious DFE significantly differs between diploid and tetraploid populations (PERMANOVA: R<sup>2</sup> = 0.174, F = 5.29, P = 0.024). A shift of DFE in tetraploid populations toward moderately deleterious mutations likely reflects the effect of polysomic masking (see text). Boxplots indicate variation among the populations (dots), asterisks indicate statistically significant difference between cytotypes as tested by the Wilcoxon rank-sum test.

Strikingly, however, the slope of this relationship is invariably steeper in tetraploid populations (Fig. 3). This is documented by the significant interaction effect of neutral variation and ploidy level on deleterious variation for the population-level sampled variants (i.e., SNPs and indels, P < 0.001, SI Appendix, Fig. S7 and Table S4). In other words, tetraploids broadly accumulate proportionally greater levels of deleterious variation than diploids do, even when taking neutral variation into account.

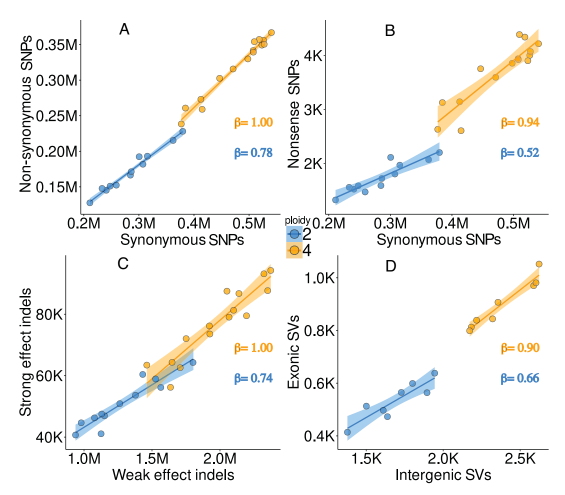
This observation is in line with the expectation of weaker purifying selection in tetraploids due to polysomic masking. In the case of SNPs, we used two different deleterious categories: all nonsynonymous SNPs (Fig. 3A) and a subset of putatively large-effect nonsense SNPs (Fig. 3B). The difference between diploid and tetraploid slopes is greater for nonsense SNPs compared to nonsynonymous SNPs (Fig. 3 A and B), indicating that the effect of polysomic masking is more pronounced for mutations with stronger fitness effects (in the case of nonsense SNPs, likely protein truncation). This corresponds with the fact that more deleterious mutations tend to be also more recessive in Arabidopsis (54) and thus more effectively masked in tetraploids (Fig. 4).

Running these analyses with the same number of chromosomes (as opposed to individuals) per population yielded qualitatively identical results (SI Appendix, Fig. S8 and Table S4), indicating robustness of the result to sampling. Qualitatively identical results

were also obtained when we focused on the total number of alleles per population (instead of sites) which quantifies total mutation load (38) (SI Appendix, Fig. S9). The amount of realized load, quantified as counts of homozygous genotypes, is higher in diploids (SI Appendix, Fig. S10) in line with expected higher homozygosity in diploid populations under equilibrium. Further, we analyzed SNP data from tetraploid populations to assess whether deleterious alleles are fully recessive or exhibit additivity. By examining the proportions of three heterozygous states (simplex 0/0/0/1, duplex 0/0/1/1, and triplex 0/1/1/1) in synonymous and nonsynonymous mutations, we found that nonsynonymous sites are enriched in simplex and depleted in duplex and triplex genotypes. This suggests dosage-dependent purging, with duplex and triplex genotypes removed more frequently (SI Appendix, Fig. S11).

### Discussion

Here, we demonstrated that WGD significantly increases genetic diversity and deleterious variation in natural populations across a range of mutation types in an outcrossing plant species. A key insight is that we demonstrated the combined effect of WGD on both neutral and selective microevolutionary processes across a diverse array of genetic markers and natural populations varying in size and natural environments occupied.



**Fig. 4.** Accumulation of proportionally greater levels of deleterious variation in tetraploids across variant types. The relationship between the number of putatively neutral (horizontal axis) and deleterious (vertical axis) SNPs, insertion-deletion (indels), and SVs in diploid (blue) and tetraploid (orange) *A. arenosa* "core" populations (*A–C*; downsampled to six individuals) or individuals (*D*). Tetraploids show not only consistently more variant sites of both categories, but also proportionally higher accumulation of deleterious variants (slope, β), an indicator of weakened efficiency of purifying selection (the interaction between the number of putatively neutral variants and ploidy is significant for all variant types except for SVs, *SI Appendix*, Table S4). Note that β was calculated for scaled and normalized data to allow direct comparison between site categories. (*A*) Number of nonsynonymous derived (i.e., with minor frequency of a nonreference allele) SNPs as a function of number of synonymous derived SNPs per population. (*B*) Number of nonsense derived SNPs (inserting premature stop codons) as a function of number of synonymous derived SNPs per population. (*C*) Number of strong effect derived indels as a function of number of weak effect derived indels per population. (*D*) Total number of exonic SVs as a function of total number of intergenic SVs per individual. Fully annotated figures with population codes are available in *SI Appendix*, Fig. S7.

Three salient points emerge from these analyses. First, increased genetic diversity following WGD is primarily driven by increase in mutational target size. In one round of WGD, theory predicts mutational target size to double, resulting in an accumulation of twofold higher polymorphism if the number of individuals of pre-WGD and post-WGD population remains constant (28). Forward-in-time simulations support this expectation; however, we show that polymorphism accumulation is gradual and doubled levels are not reached even after half a million generations. In empirical data, we see on average only 1.2-fold increase of neutral genetic polymorphism in tetraploid A. arenosa populations. This is in line with gradual polymorphism accumulation following WGD; however, in empirical data, we cannot directly distinguish the effect of increase in mutational target size and the effect of demography as easily as in simulations where population size was kept constant. Overall, this suggests that differences in population history together with the extensive time required to reach mutation-drift equilibrium set a limit to the expected doubling of genetic diversity after WGD. This may explain why similar diversity has often been reported in empirical comparisons of diploid and autotetraploid natural populations, especially in studies focusing

on recently diverging diploid-autopolyploid systems (e.g., refs. 55–57, but see, e.g., ref. 58).

Second, our analyses suggest that WGD affects selective forces on top of the effect on neutral variation. Specifically, polysomic masking in autotetraploids is expected to relax purifying selection, leading to the accumulation of deleterious mutations. Indeed, we find a relative increase of genetic diversity at constrained sites in tetraploids relative to diploids, both in terms of higher ratios of nonsynonymous to synonymous nucleotide diversity and an altered distribution of fitness effects. Moreover, we show signatures of increased mutation load across different types of genetic variants that well-supplements similar effects in TE insertions, previously reported in a subset of *A. arenosa* populations (36). A consistent, yet nonsignificant trend in SVs calls for further investigations of SV diversity at population levels—an approach that is becoming feasible with the advances in long-read sequencing (59). Altogether, our results suggest a pervasive impact of polysomic masking across the genome.

Third, despite clear evidence for accumulation of deleterious variation, our simulations and empirical data suggest that the recently formed tetraploid *A. arenosa* lineage may still benefit from lower realized load, i.e., the initial fitness effect of masking. This

is consistent with overall successful niche expansion of the autotetraploid A. arenosa (44, 60) and empirical fitness estimates showing the autotetraploids perform as well as their diploid relatives across diverse environments (61). Additionally, the increased diversity may also serve as a pool of potentially adaptive variation especially in periods of environmental turmoil (62). That polyploid A. arenosa populations adapt from a large pool of standing variation, but also occasional novel mutations, has been documented over different extreme environments (63, 64). Nevertheless, our simulations anticipate that these advantages may be transient and the accumulating deleterious variation will likely result in negative fitness consequences and increased mutation load as the tetraploid ages, unless evolutionary "rescue" through rediploidization and purging takes place (65, 66).

It is important to note that our study focuses on empirical results of a single natural WGD event within a diverse and strictly outcrossing species with large and stable populations [(34, 39), Fig. 2]. From studies in diploids, it is known that selfing and/or bottlenecks may affect load (17, 67, 68) but the interaction of these factors with ploidy remain unexplored. In other cases, nascent polyploid lineages may be affected by nonequilibrium demographic histories, as polyploid establishment may be accompanied by initial population bottleneck, range expansions (69), interploidy introgression (65, 70), and strong selection to adapt to the novel polyploid state (71–75). On the other hand, our study also revealed considerable intraspecific variation in load indicators across distinct populations, highlighting the benefit of sampling multiple population replicates when assessing load in a particular lineage. In summary, empirical investigations of diverse mixed-ploidy species spanning a range of breeding systems, polyploid ages, and demographic histories is a rich matter for further study.

Crucially, our simulations indicate that the age of polyploid lineages determines the extent of variation that has newly accumulated following WGD. Recently formed polyploids may thus exhibit a similar amount of deleterious variation as their diploid progenitors, yet higher fitness can still result, due to enhanced masking of recessive deleterious alleles. In turn, we found surprisingly little effect of post-WGD interploidy gene flow on patterns of genetic diversity, despite rampant interploidy admixture (34). This might reflect a broadly shared trans-specific gene pool across outcrossing Arabidopsis species (74, 76). Further, the impact of introgression may be stronger in other mixed-ploidy groups where ancestral and admixing diploid lineages are more diverged (e.g., Betula, (77)). Finally, the pattern of diversity and load may strongly differ in allopolyploids with disomic inheritance, where the inherited load from distinct diploid progenitors seem to play a crucial role (78–81).

Overall, our study highlights the dual role of WGD in increasing genetic diversity and mutation load, that is driven by both neutral processes and relaxed purifying selection. Polyploidy has occurred among a wide range of Eukaryotes; hence these findings situate polyploidy among important determinants of population genetic variation. Understanding how WGD shapes genetic diversity and mutation load in natural populations can inform strategies for managing genetic resources, improving crop resilience, and conserving biodiversity.

#### **Methods**

Simulation Methods. To study factors influencing genetic diversity and mutation load in autotetraploids, we conducted individual-based, forward-in-time simulations using SLiM 4 (82). Polyploids are not directly supported by SLiM, and we used an approach presented in Booker and Schrider (83) to model

autotetraploids. The simulations were conducted as non-Wright-Fisher (nonWF) models, which allow relaxing many assumptions of the standard Wright-Fisher (WF) models. In brief, generations in a nonWF model may overlap and individuals can reproduce multiple times which is the case in short-lived perennial A. arenosa (84). Fitness governs individuals' survival to the next generation, rather than determining the mating success as in a WF model. Census population size (N) is a product of reproduction and survival, resulting in N variation across generations. This dynamic is controlled by the carrying capacity of the environment (K), which is enforced by scaling the fitness of the population based on the relationship of K and N, resulting in exponential growth until K is reached In SLiM 4, nonWF models also provide more flexibility in defining reproduction and recombination than WF models, which we leveraged in simulating tetraploids. This was achieved by simulating two diploid subpopulations and using the SLiM's reproduction callback to allow crossovers to happen between chromosomes housed in the different subpopulations. Selection only acted on a single subpopulation, but dominance was a product of all four chromosomes. More details are provided in the annotated SLiM scripts, available at https:// github.com/thamala/polysim.

Genomic parameters were based on empirical studies conducted on Arabidopsis species; whenever possible, we used parameters inferred for our focal species A. arenosa as well as its close relatives from the genus Arabidopsis. We had not assumed the return of the tetraploid individuals into diploid state as the process of diploidization operates at much longer timescales in flowering plants (millions of generations, (66, 85)) and no signs of diploidization have been observed in autopolyploid A. arenosa (34, 39). However, the nonWF models employed here, especially the model implemented for tetraploids, are computationally more intensive than standard WF models, and therefore we used a rescaling approach to reduce computation times (86). Population size (N), mutation rate  $(\mu)$ , recombination rate (r), and selection coefficients (s) were rescaled by a factor of 20 while retaining the same product of  $N\mu$ , Nr, and Ns as the unscaled data (below we list the rescaled parameters). Note that we used the same scaling parameter for both diploids and tetraploids, resulting in the number of haploid genomes being double in tetraploids for a given N. However, as increased selection coefficients due to scaling could alter evolutionary dynamics associated with genetic load (87), we also conducted a subset of our simulations without scaling. Although the extent of these simulations were limited (no burn-in and maximum of 70,000 generations), we observed that the increase in genetic load in tetraploids was largely unaffected by scaling (SI Appendix, Fig. S12). Furthermore, rather than attempting to simulate whole genomes, we focused on a single chromosome of 100 kb and repeated the simulations 100 times.

We used an empirical mutation rate estimate ( $\mu = 1.39 \times 10^{-7}$ ) from A. thaliana (88), and assumed a ratio of deleterious to neutral  $\mu$  of 2.76, as inferred for A. lyrata coding sequence (89). For crossover rate, we used a genome-wide average estimated for A. arenosa ( $r = 5.6 \times 10^{-7}$ ) (90). Based on our estimates of the DFE, s for deleterious mutations were drawn from a gamma distribution with a mean of -0.048 and a shape parameter alpha of 0.228 (average across the diploid A. arenosa populations). Using data from A. lyrata, Huber et al. (54) coestimated the distribution of selection and dominance coefficients (h) of new mutations. Following their results, we defined dominance coefficients based on the s of each mutation:

$$h = \frac{1}{\frac{1}{\theta_{totages}} - \theta_{rate}},$$
 [1]

where  $\theta_{intercept} = 0.978$  and  $\theta_{rate} = 50328$ , as inferred for A. lyrata (54). This continuous h-s model aims to capture the inverse relationship between selection and dominance often observed in empirical data (91, 92). See SI Appendix, Fig. S13 for a visualization of the h-s model and SI Appendix, Figs. S14 and S15 for showing our main simulation results achieved through purely additive and dominant models, respectively. Following Layman and Busch (93), we defined the fitness effect of each mutation as  $1 + h_x s$ , where  $h_x$  is a ploidy-independent dominance weight:

$$h_x = \frac{1}{1 + \left(\frac{1}{k}\right)^y \left(\frac{1-x}{x}\right)},$$
 [2]

where x is the fraction of mutant copies (e.g., x = 0.5 for Aa and AAaa genotypes). Other components of the function (k and y) were solved as

$$k = 2h, y = \frac{\ln\left(\frac{1}{h} - 1\right)}{\ln\left(\frac{1}{2h}\right)}.$$
 [3]

See *SI Appendix*, Fig. S16 for a visualization of the hX model. We note that although dominance in polyploids is poorly understood (25), increased heterozygosity in polyploids leads to an accumulation of (partially) recessive load even when dominance coefficients in tetraploids are comparable to those in diploids (*SI Appendix*, Fig. S17).

To define population sizes for the simulations, we transformed our empirical estimates of the population mutation rate (4 $Ne\mu$  in diploids and 8 $Ne\mu$  in tetraploids) to Ne by assuming  $\mu = 6.95 \times 10^{-9}$  (88), which gave an average Ne estimate of 183,680 for diploids and 104,680 for tetraploids. However, as we were primarily interested in assessing the effects of WGD (rather than Ne), we assumed the same N for both cytotypes. We started with a diploid population of K = 9,184 (corresponding to unscaled N of 183,680). To define a burn-in period for the simulations, we examined neutral nucleotide diversity ( $\pi_{Neutral}$ ) and the ratio of deleterious to neutral diversities ( $\pi_{Deleterious}$  /  $\pi_{Neutral}$ ) across 50 K generations. Based on the results (*SI Appendix*, Fig. S18), we chose 10 K as a sufficient period for the diploid population to establish before an autotetraploid population was founded by duplicating the genomes of either 10 individuals (bottleneck scenario) or the whole population (nonbottleneck scenario presented in Fig. 1). We acknowledge that switching the whole population to polyploidy is likely unrealistic, but it allows us to distinguish the effects of WGD from the founding bottleneck and it also approaches a realistic situation for A. arenosa (34, 39) when the diversity of nascent autopolyploid population is enriched by post-WGD unidirectional gene flow from sympatric progenitor diploid population via unreduced gametes (72). To evaluate the mutation load of the newly founded autotetraploid population, we estimated nucleotide diversity for deleterious (  $\pi_{\text{Deleterious}}$  ) and neutral (  $\pi_{\text{Neutral}}$  ) variants, calculated the number of strongly deleterious (s < -0.01) and nearly neutral (s > -0.0001) mutations, and estimated fitness for each individual as a product of fitness effects of all mutations. These estimates were then compared to populations that remained diploid throughout the simulation.

**Empirical Data Sampling and Sequencing.** We gathered whole-genome short-read sequencing data of *A. arenosa* from all available published genome-sequencing datasets including populations with at least six and more well-covered individuals (34, 63, 64, 76). We complemented them with additional sequencing (126 individuals, 13 populations) to representatively cover all *A. arenosa* diploid and tetraploid lineages. In total, we gathered short read genome sequences for 634 accessions from 65 populations of *A. arenosa*. We removed reads of low quality and adaptor sequences with trimmomatic-0.36 (94) and we mapped refined reads to the new *A. arenosa* genome (95) by bwa-0.7.15 (96) with default setting. We used picard-2.8.1 to mark duplicate reads and called genotypes with GATK (v.3.7). We used HaplotypeCaller to infer genotypes per individual with respect to its ploidy. Information about ploidy of each individual was adopted from previous studies (published data) or it was detected by flow cytometry following our standard protocol (40) for the newly sequenced individuals.

To filter reliably called SNP genotypes we followed GATK best practices (97) and filtering strategies established in our previous studies involving autopolyploid populations (34, 64). For the SNP-based indices of mutation load and an analysis of population structure we used biallelic SNPs that passed these filtering parameters (FS > 60.0 || SOR > 3 || MQ < 40 || MQRankSum < -12.5 || QD < 2.0 || ReadPosRankSum < -8.0). Further, we masked genes with excessive heterozygosity with at least five fixed heterozygous SNPs in at least two diploid populations as being potentially paralogous genes following Monnahan et al. (34). Also we masked sites with excessive depth of coverage defined as sites with depth that exceeded twice SD of depth of coverage in at least 20 individuals. Retaining only populations with at least six individuals left us with a "full" dataset of 65 populations (632 individuals) with 54,179,280 SNPs of average sequencing depth of 24.2.

For methods that depend on precise inference of allele frequency spectra, we used only populations with mean SNP depth above 23. Applying such criteria left us with 27 sufficiently deeply sequenced core dataset populations (12 diploid and 15 tetraploid) with mean depth of coverage of 36.1 (*SI Appendix*, Fig. S5).

Indel Variant Calling and Filtration. For the indel analysis we focused on the 27 deeply short-read sequenced core populations. We considered only sites that passed GATK best practice filters with the same parameters as for SNPs (FS > 40.0  $\parallel$  SOR > 3  $\parallel$  MQ < 40  $\parallel$  MQRankSum < -12.5  $\parallel$  QD < 2.0  $\parallel$  ReadPosRankSum < -4.0  $\parallel$  INFO/DP > 2\*mean coverage'). On top of that we removed sites with an excess depth coverage calculated as two times the mean coverage of the whole dataset to avoid misassembled paralogous sites. Only indels supported by at least five reads per individual and with absolute length up to 20 bp were considered. We did not remove multiallelic indels due to their ubiquity (48% of sites were multiallelic); we rather split the overlapping indels and counted them as distinct variants. We recovered 14.4 M reliable indels while removing 59.4% of the raw number of indels (35.5 M) that were called by GATK.

SV Detection, Calling, and Filtration. SVs were recovered using Oxford Nanopore long-read sequencing Technology (ONT). We extracted high molecular weight DNA from A. arenosa leaves as described by Russo et al. 2022 (98). The DNA concentration was checked on a Qubit Fluorometer 2.0 (Invitrogen) using the Qubit dsDNA HS Assay kit. Fragment sizes were assessed using the Genomic DNA Tapestation assay (Agilent). Removal of short DNA fragments and final purification to high molecular weight DNA was performed with the Circulomics Short Read Eliminator XS kit. ONT libraries were prepared using the Genomic DNA Ligation kit SQK-LSK109 following the manufacturer's procedure. Libraries were loaded onto R9.4.1 PromethION Flow Cells and run on a PromethION Beta sequencer. Due to the rapid accumulation of blocked flow cell pores or due to apparent read length anomalies on some runs, flow cells used in the runs were treated with a nuclease flush to digest blocking DNA fragments before loading with fresh libraries according to the ONT Nuclease Flush protocol (version NFL\_9076\_v109\_revD\_08Oct2018). FAST5 sequences produced by PromethION sequencer were basecalled using the Guppy6 (https://community.nanoporetech. com) high accuracy basecalling model (dna\_r9.4.1 \_ 450bps\_hac.cfg) and the resulting FASTQ files quality filtered by the basecaller. For read depth and quality of each sample see SI Appendix, Table S5.

Minimap2 (v2.22) (99) was used to map the Nanopore reads against the reference genome of *A. arenosa* (95) with default parameters. We used the following SV calling pipeline developed for autotetraploid *Cochlearia officinalis* and validated using simulated autotetraploid data (37). SVs were identified using Sniffles2 (v2.0.6) (100) run in germline mode, limiting the minimum supporting reads depending on coverage (--minsupport auto). We only kept insertions and deletions between 50 bp and 100 kb in length, as read-alignment-based methods are less accurate at detecting other types of SVs as well as very large SVs. The R package updog based on a genotype-likelihood approach for allele frequency estimation in polyploids (101) was then used to estimate allelic dosage for each individual and SV. The depth of reference-supporting reads and variant-supporting reads for the SVs were retrieved from VCF files generated by Sniffles2. The multidog function was used for SVs with a minimum of 10 supporting reads with "norm model" and setting the ploidy level as 2 and 4 for diploids and tetraploids, respectively.

**Population Structure Inference.** Relationships between all populations (full SNP dataset) have been visualized using PCA and Neighbor joining tree to confirm that our data match the structure observed in the previous studies which focused on the population structure in depth (34, 39, 41, 44). For the inference we used a frequency of 92,632 fourfold degenerate biallelic SNPs with minimum genotype depth 8× and maximum 25% missing genotypes per site. This representation of unlinked sites was retrieved by pruning the fourfold sites with maximum 0.25 linkage disequilibrium coefficient ( $r^2$ ) in 80 kb windows along the genome. We also removed sites where minor allele frequency was lower than 0.1. First, we used principal component analysis (PCA) calculated using gIPCA from the adegenet package (102) to visualize the population structure in multidimensional space. Then we calculated Nei's distances (103) using StAMPP (104) and constructed a neighbor joining tree with the program SplitsTree (105).

**SNP-Based Analyses of Population Diversity.** To test whether genetic diversity differs between diploid and tetraploid populations, we calculated genomewide nucleotide diversity at putatively neutral fourfold degenerate sites and constrained zerofold degenerate sites for each population. We also estimated

Tajima's D based on fourfold degenerate sites serving as a rough proxy of population demographic history. All statistics were computed using the calcwpm function from the program ScanTools (65, github.com/mbohutinská/ScanTools\_ProtEvol), following current best practices in autopolyploid population genomics (106). We included all biallelic SNPs with a minimum genotype depth of 8× and a maximum of 25% missing data per site.

Initially, each population was downsampled to six individuals. However, we were concerned that sampling different numbers of chromosomes per population (12 for diploids and 24 for tetraploids) might bias nucleotide diversity estimates. To equalize the number of sampled alleles across ploidy levels, we instead downsampled tetraploid populations to four individuals and diploid populations to eight individuals (sampling 16 chromosomes in both cases).

We tested for statistical differences in nucleotide diversity, the ratio of zero-to fourfold diversity, and Tajima's D between diploid and tetraploid populations using Wilcoxon's rank sum test. To account for potential technical variation in our diversity estimates, we additionally tested the effect of ploidy on nucleotide diversity using a general linear model, including sequencing depth as a covariate. All statistical analyses were conducted in R version 4.1.2 (107).

**Distribution of Fitness Effects of SNP Mutations.** To test whether distribution of fitness effects (DFE, 2) differs between ploidy levels we first generated carefully polarized unfolded site frequency spectra (SFS) for zero- and fourfold SNPs for each core population using the program est-sfs (108). As this method depends on correct characterization of allele frequency spectra, we focused on a subset of 27 sufficiently deeply sequenced populations (core dataset). Est-sfs program input consists of allele counts of focal species and several outgroups. Allele counts per population of A. arenosa were parsed by an in-house python script sampling 16 chromosomes per population. For the outgroup counts we used A. thaliana and Capsella rubella and we merged the homologous sites of outgroups and our focal A. arenosa by genomic alignment of all species. Discrete DFE was then inferred using the repolarized zero- and fourfold SFS as inputs for each population by the program poly-DFE (49). This program allows inference of DFE with multiple parameters and different prior distributions of fitness effects. In order to explore the parameter space we ran models for all three basic modes of distributions (only deleterious, deleterious + beneficial displaced gamma distribution and deleterious + beneficial exponential distribution). Each distribution model was further refined by including or excluding a nuisance parameter for demography and a parameter for polarization error. We then employed a series of likelihood ratio tests to determine the bestfitting model for each population. For comparison across populations, we finally used a model incorporating a deleterious + beneficial exponential distribution, a nuisance parameter, and a polarization error parameter. This model showed the highest likelihood in model comparisons across the majority of populations. To assess whether the deleterious distribution of fitness effects (DFE) differed between diploid and tetraploid populations, we performed a PERMANOVA (Permutational Multivariate ANOVA) using the adonis function from the vegan R package ver. 2.6-4 (999 permutations, Euclidean distance). Each replicate was represented as a vector of proportions across four discrete scaled selection coefficient bins (4\*Ne\*s) ranging from highly deleterious mutations "<-100", to effectively neutral mutations "(-1,0)". This approach allowed us to test for significant differences in the multivariate composition of the DFE between ploidy levels, while accounting for within-group variation. The difference between ploidy levels in individual selection coefficient bins was tested by Wilcoxon's rank sum test.

Correlational Approach to Determine Mutation Load Across Different Variant Types. We quantified mutation load over SNPs, short indels (both in stringently filtered core short-read dataset of 27 populations), and SVs (all 16 long-read sequenced individuals, one per population) using a measure that allows consistent comparison among these different genetic markers. We calculated correlation between count of putatively neutral variants and selectively constrained variants per population and used these values as an index of mutation load, following studies of human populations (38).

First, we annotated the variants based on their putative phenotypic effect to get the two categories (neutral and constrained by natural selection). For indels and SNPs we used SnpEff (5.1) (109) to annotate the putative phenotypic effect of each variant. For indels we used variants annotated as "HIGH" effect (typically frame-shift mutations) as constrained and "LOW" as neutral (intronic and intergenic variants). In SNPs, we used two constrained categories:

a) nonsynonymous SNPs (i.e., changing the resulting amino acid) b) nonsense SNPs (i.e., causing a premature stop codon) and one neutral category of synonymous SNPs. For SVs we used intergenic SVs (>5 kb away from genes) as neutral category and exonic SVs (i.e., those at least partially covering an exon) as constrained. Then we counted all variable sites with nonreference allele frequency lower than 0.5 (to include only putatively derived variants) per each category and population. The populations were downsampled to six individuals for the SNP and indel dataset to achieve equal sampling effort. Alternatively we also downsampled tetraploid populations to four individuals and diploid populations to eight individuals to explore whether sampling the same number of chromosomes per ploidy yields consistent results. Further we also counted alleles of a given category which represents an alternative summary of total mutation load (38). We also estimated realized mutation load as the number of homozygous genotypes. It shall be noted, however, that comparison of this index between cytotypes is directly affected by overall lower expected homozygosity in tetraploids (q<sup>2</sup> in diploid while q<sup>4</sup> in tetraploid populations).

We tested the effect of ploidy on the number of constrained variants (SNPs, indels, and SVs), while accounting for a "baseline" neutral variation of each population, using generalized linear model from package stats in R version 4.1.2 (106). The model involved the number of constrained variants as response variable that was explained by ploidy and number of neutral variants, their interaction, and the depth of coverage per population as predictors (constrainedN~ploidy\*neutralN+DP). To test for the effect of the interaction between ploidy and number of neutral variants, we compared two hierarchical models using the likelihood ratio test: i) with fixed effect of each predictor vs. ii) with additional ploidy:neutralN interaction. Because the dependent variable is counts we used a Poisson distribution to model residual variance of the model.

Finally, we analyzed SNP data from tetraploid populations to assess whether deleterious alleles are fully recessive or exhibit additive effects. To do this, we compared the proportion of simplex (0/0/0/1), duplex (0/0/1/1), and triplex (0/1/1/1) heterozygous genotypes in synonymous and nonsynonymous mutation categories. Proportion of each heterozygous genotype category from the total number of heterozygous genotypes was calculated per each population downsampled to six individuals. Differences between synonymous and nonsynonymous mutations were tested using the Wilcoxon rank-sum test to evaluate potential dosage-dependent effects.

Data, Materials, and Software Availability. Sequence data that support the findings of this study are deposited in the NCBI (https://www.ncbi.nlm.nih. gov/bioproject/) under BioProjects PRJNA929698 (110), PRJNA284572 (111), PRJNA484107 (112), PRJNA592307 (113), and PRJNA667586 (114) (short read data) and PRJEB83985 (115) (long read data).

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