

RESOURCE ARTICLE

Inference of the distribution of fitness effects of mutations is affected by single nucleotide polymorphism filtering methods, sample size and population structure

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Funding information

Vetenskapsrådet

Handling Editor: Kimberly Gilbert

Abstract

The distribution of fitness effects (DFE) of new mutations has been of interest to evolutionary biologists since the concept of mutations arose. Modern population genomic data enable us to quantify the DFE empirically, but few studies have examined how data processing, sample size and cryptic population structure might affect the accuracy of DFE inference. We used simulated and empirical data (from *Arabidopsis lyrata*) to show the effects of missing data filtering, sample size, number of single nucleotide polymorphisms (SNPs) and population structure on the accuracy and variance of DFE estimates. Our analyses focus on three filtering methods—downsampling, imputation and subsampling—with sample sizes of 4–100 individuals. We show that (1) the choice of missing-data treatment directly affects the estimated DFE, with downsampling performing better than imputation and subsampling; (2) the estimated DFE is less reliable in small samples (<8 individuals), and becomes unpredictable with too few SNPs (<5000, the sum of 0- and 4-fold SNPs); and (3) population structure may skew the inferred DFE towards more strongly deleterious mutations. We suggest that future studies should consider downsampling for small data sets, and use samples larger than 4 (ideally larger than 8) individuals, with more than 5000 SNPs in order to improve the robustness of DFE inference and enable comparative analyses.

KEYWORDS

DFE, missing-data treatment, population structure, sample size, SLiM simulation

1 | INTRODUCTION

The *distribution of fitness effects* (DFE) of new mutations can be described as the probability that a new mutation will have a specific effect on the fitness of an individual. This probability distribution affects the accumulation of genetic variation and can thus

directly impact the evolutionary trajectory of organisms (Bataillon & Bailey, 2014; Keightley & Eyre-Walker, 2007; Ohta, 1992). Understanding the DFE is integral to understanding molecular evolution and remains an important focus in modern evolutionary theory (Chen et al., 2020; Halligan & Keightley, 2009; Kimura, 1968; Ohta, 1973). To date, the arguably most popular

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methods of inferring the DFE are based on contrasting frequencies of putatively neutral and selected polymorphisms presented as a site frequency spectrum (SFS), describing how commonly mutations of different frequencies occur in a population (Gutenkunst et al., 2009; Keightley & Eyre-Walker, 2007; Kim et al., 2017; Tataru & Bataillon, 2019). Since the SFS can be affected by both neutral and selective processes, most methods use the SFS of synonymous mutations to estimate a demographic model representing the effects of population size changes and genetic drift. Meanwhile, the SFS of nonsynonymous mutations are assumed to be shaped by both neutral and selective processes and can therefore be used to estimate the DFE of non-neutral mutations after demography and drift have been accounted for (Boyko et al., 2008; Huang et al., 2021; Keightley & Eyre-Walker, 2007; Kim et al., 2017; Schneider et al., 2011; Tataru & Bataillon, 2019). However, factors other than demography and selection may also affect the shape of the SFS and thus the estimated DFE.

First, SFS-based DFE inferences require that data sets contain no missing sites—all individuals must have complete data for all loci that are to be analysed. Since sequencing techniques are imperfect, such data sets are uncommon (probably nonexistent) in empirical population genomics. As a result, missing-data treatment is an essential first step of data processing. To obtain a complete data set, these data are treated either by filtering out some portion of the data (sub- or downsampling), or filling in the 'gaps' using an algorithm such as imputation (see Section 2.2). Depending on how the treatment is performed, there is a risk of altering the relative allele frequencies in the data set, yielding misleading results (Johri et al., 2021; Larson et al., 2021). Recent studies on DFE have applied different data processing methods; for example, see Hämälä and Tiffin (2020) for imputation, and Gossman et al. (2010) for downsampling. However, it is unknown whether and how the different methods influence DFE estimates.

Second, the sizes of data sets used in published DFE studies vary enormously, from as few as two to several hundred individuals (Chen et al., 2017; Hämälä & Tiffin, 2020). The SFS is highly sensitive to sample size, but the minimum number required to achieve stable DFE estimates remains undetermined (but see Kutschera et al., 2020). Similarly, the number of polymorphic sites necessary for reliable DFE estimation is largely unknown. While some studies of model species use whole genome sequencing with millions of single nucleotide polymorphisms (SNPs) available for analysis (Hämälä & Tiffin, 2020), others may only include a few hundred SNPs (Eyre-Walker & Keightley, 2009; Gossman et al., 2010). Therefore, investigating the impact of sample size (both the number of individuals and sites/SNPs) on DFE estimates is crucial for reliable and accurate DFE estimation.

Finally, most methods of SFS-based DFE estimation first estimate a Wright-Fisher demographic model from the neutral variation in order to control for neutral factors affecting the SFS (Keightley & Eyre-Walker, 2007; Tataru & Bataillon, 2019). Such models assume that mating occurs at random in panmictic populations, even though complete absence of population structure is

likely rare in wild samples. For example, sampling from a large area is preferred for drawing general conclusions about population genetic dynamics, but it increases the likelihood of including genetic structure in the sample (Perez et al., 2018; Zhao et al., 2020). If cryptic genetic clusters are unwittingly included, the demographic model estimated from the data would not fulfil the assumptions underlying the Wright-Fisher model, and subsequent DFE estimates might be biased. However, population stratification has not to our knowledge been examined as a potential factor affecting the accuracy of DFE inference.

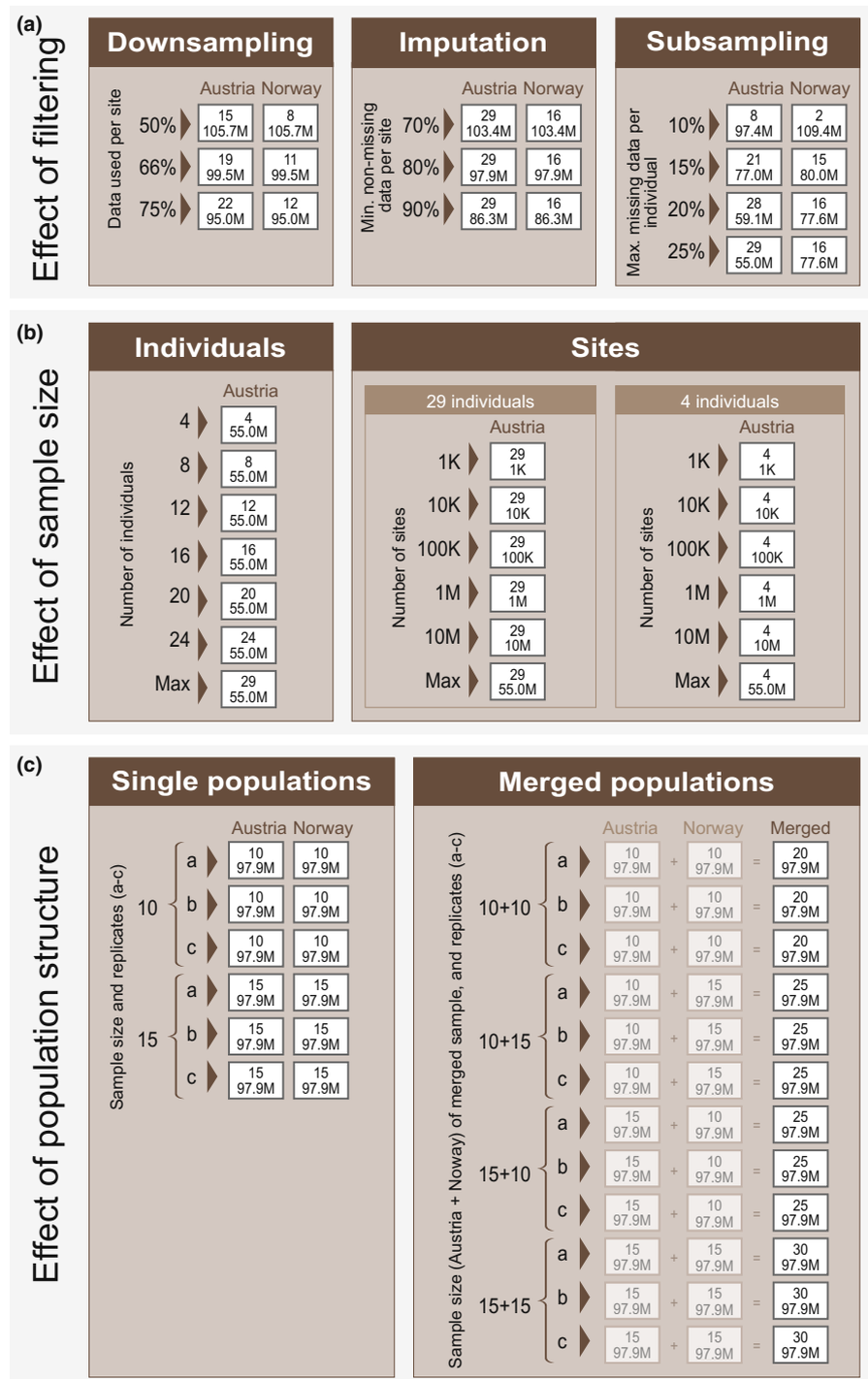
In this study, we test whether and how data processing methods, sample size, SNP number and population structure influence the results of DFE inference, to raise awareness of their potential confounding effects. We used whole genome resequencing data from two populations of *Arabidopsis lyrata* (subsp. *petraea*) to create multiple data sets (Figure 1) with (1) three different methods of missing-data treatment—downsampling, imputation and subsampling—under different filtering thresholds; (2) different numbers of randomly sampled individuals and sites; and (3) samples with induced population stratification, to be contrasted with uniform, single populations. Then, we conducted forward simulation in SLiM 4.0 (Haller & Messer, 2023) to create a population with a known DFE that matches DFEs estimated in *A. lyrata*. Using this known DFE, we evaluate the accuracy of DFE estimates resulting from the different data manipulations. By contrasting the results obtained from the different procedures, we aim to answer the following questions: (1) Do data processing methods and missing-data filtering thresholds affect DFE estimation, and if so, how? (2) How many individuals and SNPs are needed to reach an accurate DFE estimate? and (3) Does population structure affect the DFE, and if so, how? Our results illustrate the importance of careful consideration of all steps in genomic data processing and analysis, both when performing DFE inference and when interpreting its results.

2 | MATERIALS AND METHODS

2.1 | Genomic data set and basic quality control

We downloaded the whole genome resequencing data for two populations of the perennial, diploid obligately outbreeding *Arabidopsis lyrata* subsp. *petraea*, 29 individuals from Austria and 16 individuals from Norway, from the NCBI SRA database (Table S1). The quality of the sequence reads was first assessed with FastQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>). Adapter sequences and low-quality bases were removed using fastp v0.23.0 (Chen et al., 2018) with the parameters '-q 20 -l 36 --cut_front --cut_tail -c'. Clean reads were mapped to the *A. lyrata* v.1.0 genome (<https://plants.ensembl.org/>) using the BWA-MEM algorithm with default parameters (Li, 2013). PCR duplicates were removed using Picard MarkDuplicates (<http://broadinstitute.github.io/picard/>). Reads around putative insertions and

FIGURE 1 Experimental design. We performed three sets of tests to understand the potential influence on the estimated distribution of fitness effects (DFE) of: (a) three methods of missing-data treatment, (b) the number of individuals and sites used, and (c) population structure. Each box represents a derived data set, with the number of individuals shown on top and nucleotide sites below. The study involved two populations of *Arabidopsis lyrata* from Austria and Norway. We created merged populations with subsets of individuals from Austria and Norway as specified on the left of each of the merged boxes (c, greyed out). The estimated DFE of the merged population are compared to that of the contributing populations.



deletions were locally realigned using RealignerTargetCreator and IndelRealigner in the Genome Analysis Toolkit (GATK v.3.7.0; Van der Auwera et al., 2013). Variants were called using the SAMtools and BCFtools pipeline as described previously (Li, 2011). Several filtering steps were performed to minimize genotyping errors: indels and SNPs with mapping quality (MQ) <30 were removed, genotypes with genotype quality (GQ) <20 or read depth (DP) <5 were masked as missing, and all SNPs with a missing rate above 50% or allele number above 2 were removed. After these basic filtering steps, a total of 122,432,856 sites (including invariant sites) were retained in the 45 samples for the following analyses.

2.2 | Missing-data treatment methods

Missing genotypes are common in genomic data sets and should be eliminated before generating an SFS. We tested three methods to treat missing values on the same original data sets—*downsampling*, *imputation* and *subsampling* (Figures 1a and 2), and then compared the DFE inferred from each resulting data set using bootstrapped 95% confidence intervals (CIs).

Downsampling is performed by randomly selecting n genotypes at each site without replacement (Keightley & Eyre-Walker, 2007); sites with fewer than n genotypes available are removed. A 75%

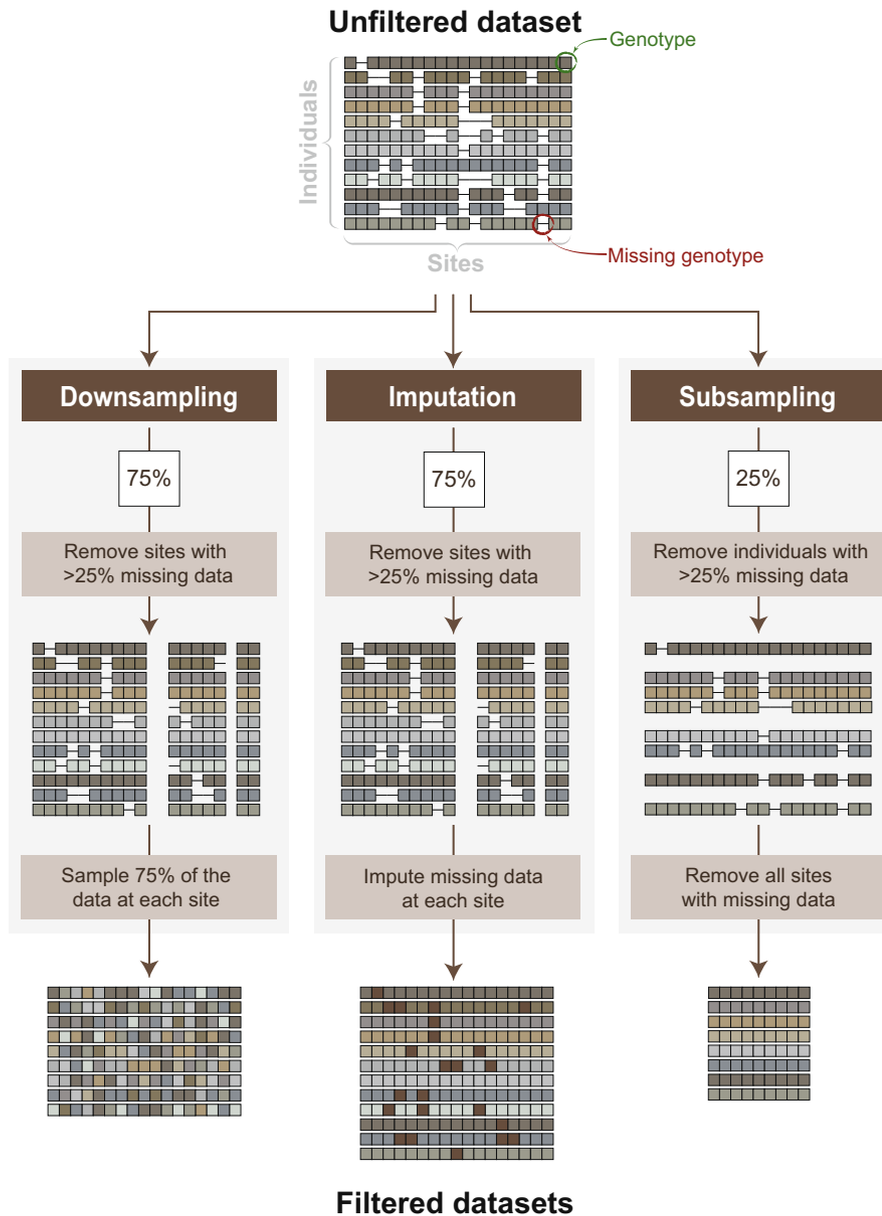


FIGURE 2 Methods of missing-data treatments for site frequency spectrum based analyses. Illustration of the different steps involved in the three missing-data filtering methods examined in this study. Each box corresponds to an individual's genotype at a site, and missing boxes represent missing data for a genotype. In downsampling, step 1 excludes sites at which data is missing in more than a prescribed threshold of individuals (e.g. 25%), while step 2 samples genotypes without replacement from the remaining data at each site. In imputation, as in downsampling, step 1 excludes sites with missing rate more than a prescribed fraction, while step 2 imputes (fills in) missing data. In subsampling, step 1 excludes individuals with missing data in more than a prescribed fraction of sites, while step 2 excludes all sites with missing data.

downsampling threshold in a sample size of 100 individuals means that 75 random genotypes are sampled at each site (Figure 2). Sites that contain <75 genotypes are removed. In this study, we applied downsampling at thresholds 75%, 66% and 50% on both Austrian and Norwegian data sets. The same set of sites were kept and analysed in both populations, making direct comparisons of the DFE between populations possible. Downsampling was performed using a Python script available on Dryad (Papadopoulou & Knowles, 2015) with minor modification (https://github.com/hui-liu/Bioinformatics-Scripts/blob/master/Scripts/Python/sampleDownMSFS_Hui_final.py).

Imputation refers to the statistical inference ('filling in') of missing genotypes using the available linkage information from successfully genotyped samples (Figure 2). We tested thresholds 70%, 80% and 90% on the *A. lyrata* data sets (i.e. excluding sites with less than 70%, 80% and 90% genotype information available), and filled in the missing genotypes at all other sites using Beagle v5.1 (Browning

et al., 2018) with default parameters. We performed imputation using all individuals from both populations, as imputation accuracy tends to increase with sample size, as shown by previous studies (Pook et al., 2020).

Subsampling works in two steps: (1) Individuals who are missing more than a prescribed fraction of their genotype information are excluded, and (2) for the individuals remaining, any site with a missing genotype is removed (Figure 2). This means that the size of a subsampled data set is highly dependent on the individual missing rates and the distribution of missing data across the genome. We first calculated the missingness on a per-individual basis using the parameter '--missing-indv' in VCFtools (Danecek et al., 2011). We then extracted the individuals that had missing rates below the threshold value using '--keep', and finally, we removed all sites containing missing genotypes by setting the parameter '--max-missing 1' in VCFtools. In the *A. lyrata* data set, we tested four maximum missing rates per individual—10%, 15%, 20% and 25% (Note: no individual

had more than 25% missing data). Note that with higher subsampling thresholds, more individuals but fewer sites are retained (Figure 1a).

2.3 | Sample size and SNPs number

To decouple the potential effects of the number of individuals and/or sites on DFE estimation, we randomly sampled 4, 8, 12, 16, 20, 24 or 29 (all) individuals and/or 1K, 10K, 100K, 1M, 10M or 55.0M sites from the Austrian population subsampled at a maximum missing rate of 25% per individual (Figure 1). To investigate the effect of sample size, we kept all sites and compared samples with different numbers of individuals (4–29). Conversely, to investigate the effect of the number of SNPs included in the SFS, all 29 individuals were kept and a randomly chosen subset of 1K to 10M sites were extracted. Finally, the same subsets of 1K to 10M sites were extracted from a data set with only four individuals. By comparing the DFEs from 4 versus 29 individuals for each set of sites, we could see the combined effects of the number of individuals and sites on the estimated DFE and confidence intervals (Figure 3).

2.4 | Manipulating population structure

To gain an overview of the genetic differentiation between the Austrian and Norwegian populations, we performed a principal component analysis (PCA) on the 45 sampled individuals using Eigensoft v.6.1.4 (Price et al., 2006). The data set was filtered at a maximum missing rate of 20% per site and a minor allele frequency (MAF) ≥ 0.05 , retaining 3,921,575 SNPs for the PCA. To investigate whether population structure affects DFE estimates, we randomly selected three different subsets (labelled a, b and c) of 10 and 15 individuals from each of the Austrian and Norwegian populations, imputed at an 80% threshold. Single sets from each population were then combined to form 12 new merged populations with four different configurations (Figure 1c): 10 Austrian+10 Norwegian individuals, 10 Austrian+15 Norwegian individuals, 15 Austrian+10 Norwegian individuals and 15 Austrian+15 Norwegian individuals, each with three replicates. We then estimated the DFE for each subset and all merged samples.

Using the single and merged data sets, we investigated (1) the effect of sample choice within a geographic population on DFE, by comparing the three replicate subsets from a single population (e.g.

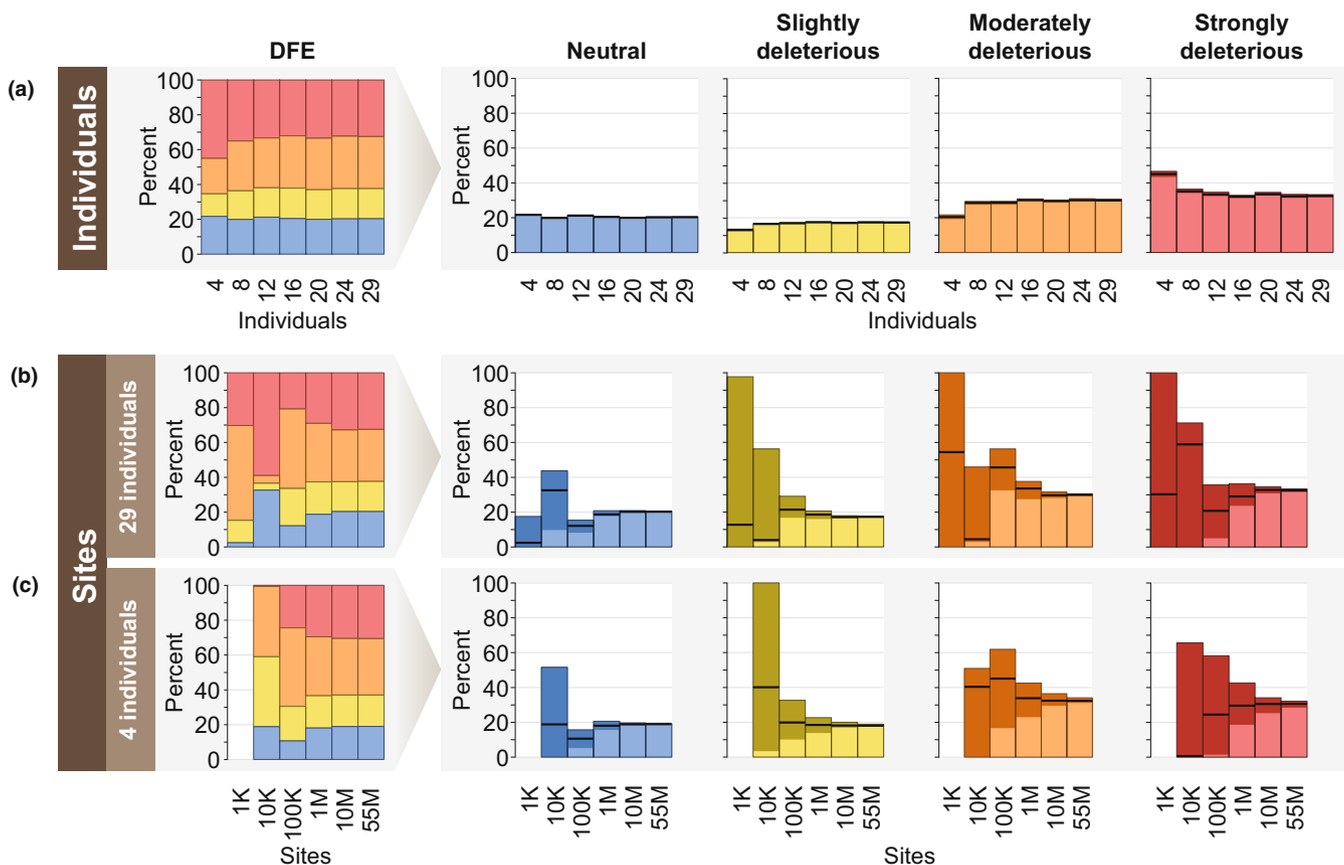


FIGURE 3 Effects of number of individuals and sites on distribution of fitness effects (DFE). DFE estimated from *Arabidopsis lyrata*, (a) random samples of 4, 8, 12, 16, 20 and 24 of the 29 individuals of the Austrian population with 55M sites; (b) all 29 individuals, (c) a random sample of 4 individuals with 1K, 10K, 100K, 1M, 10M and 55M sites. The complete DFE is represented as percentage contribution of each of four categories of mutations: *neutral* (blue), *slightly deleterious* (yellow), *moderately deleterious* (orange) and *strongly deleterious* (red). The DFE for each sample size is represented in two ways: on the left as stacked estimated percentages of the four categories of mutations, and on the right as the estimated percentage of each category of mutations (black bars) together with the 95% confidence intervals (darker coloured areas).

replicates *a* vs. *b* vs. *c* of subset Aus10), (2) the effect of each geographic population on the merged population, by comparing the DFE of the merged population to each of the contributing populations (e.g. replicate *c* of merged population Aus10+Nor15 vs. replicate *c* of subsets Aus10 and Nor15) and (3) the effect of population differentiation (F_{ST}) on DFE in the merged population. The weighted F_{ST} between the two contributing subsets in each merged population was calculated using VCFtools.

2.5 | DFE analyses

We used DFE-alpha (Eyre-Walker & Keightley, 2009), a software that uses a maximum-likelihood approach to determine the shape of the DFE of nonsynonymous mutations. In the simplest model, DFE-alpha assumes that mutations at synonymous sites are selectively neutral and that all nonsynonymous mutations are deleterious. DFE-alpha first estimates a simple demographic model using the SFS of neutral mutations to represent the effect of drift. We modelled the effect of recent demographic change on neutral SFS by assuming one step population size change and inferred the fitness of new deleterious mutations at the selected sites from a gamma distribution while simultaneously fitting the estimated parameters for the demographic model. The estimated fitness effects of new mutations are scaled by effective population size N_e and selection coefficient s as $N_e s$, and divided into four categories: *effectively neutral* ($0 < -N_e s \leq 1$), *slightly deleterious* ($1 < -N_e s \leq 10$), *moderately deleterious* ($10 < -N_e s \leq 100$) and *strongly deleterious* ($-N_e s > 100$). The DFE is presented as the proportion of nonsynonymous mutations that is expected to fall into each of these categories.

We generated a folded SFS for a class of putatively neutral reference sites (4-fold degenerate sites) and a class of selected sites (0-fold degenerate sites) for each data set. We modelled the effects of recent demographic change on the 4-fold sites SFS by assuming a single population size change event and inferred the fitness of new deleterious mutations at the 0-fold sites from a gamma distribution. The 95% CIs for all DFE estimates were calculated by bootstrapping 0-fold and 4-fold sites with replacement for 99 iterations. We performed bootstraps using 999 and 99 iterations in nine samples and found no discernible difference in CI size; all reported CIs are thus based on 99 iterations.

2.6 | Simulations in SLiM

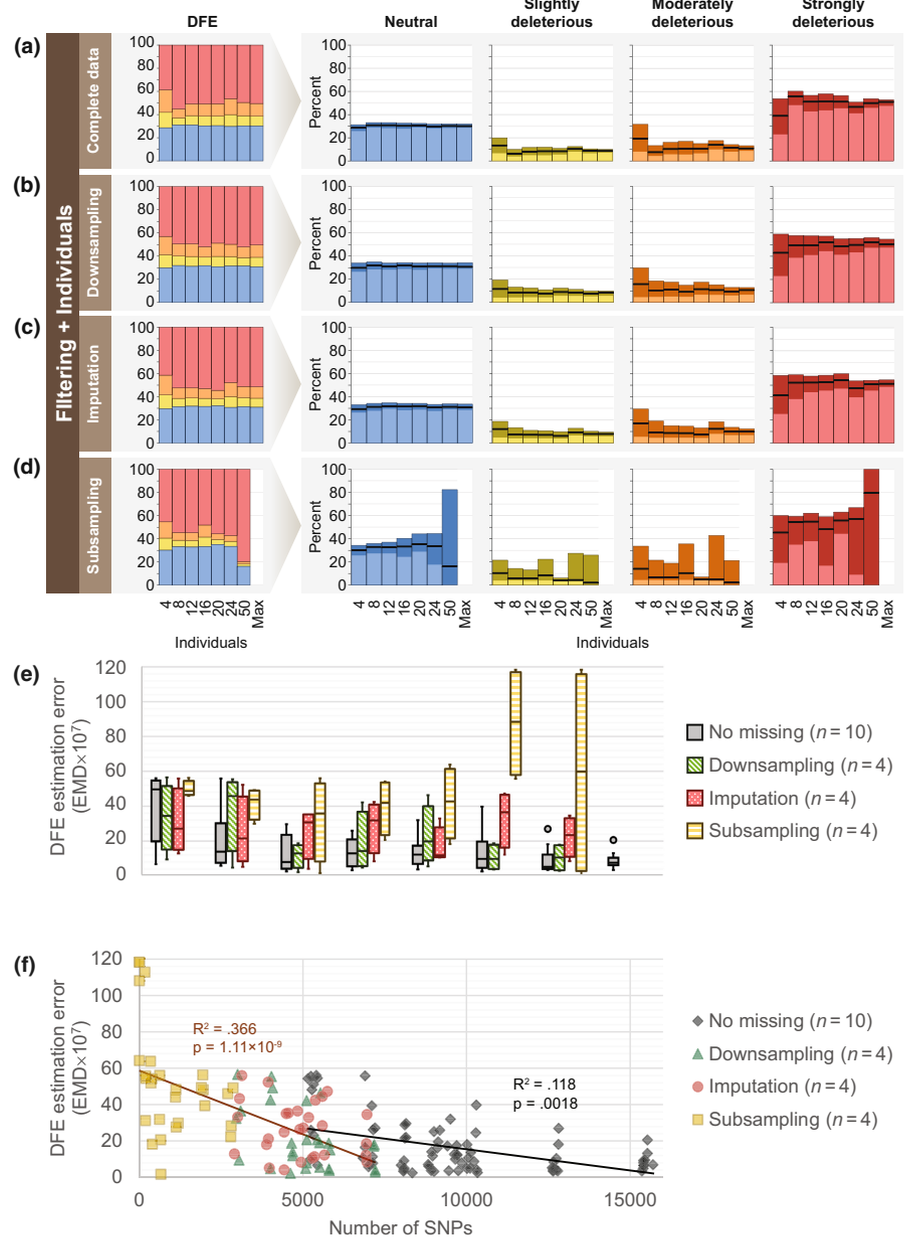
To validate the effects of filtering methods and sample size on DFE estimates, we used SLiM 4.0 to simulate a population with a known DFE, represented by a gamma distribution with shape (β) and mean (Es) parameter values matching the DFE estimated in *A. lyrata*. The simulation consisted of a population of 10,000 outcrossing individuals with a genome size of 5 million sites on one contiguous chromosome, and a uniform recombination rate of 4×10^{-8} (Hämälä & Tiffin, 2020). New mutations occurred at a mutation rate of

5.6×10^{-8} and were drawn from a deleterious DFE with a gamma distribution with $\beta=0.1$ and $Es=-100$. The population state at 60,000 generations was saved as a .trees file, at which point the effective population size N_e had stabilized around 100 individuals with 72,330 segregating deleterious mutations. A neutral burn-in and segregating neutral mutations were then added with recapitation and overlaid mutations based on tree sequence recording, which enables introduction of neutral mutations as though they occurred during the simulation by tracking the genealogy of each genome backwards in time. By overlaying mutations with a mutation rate of 1.4×10^{-8} for neutral mutations, a total of 23,846 segregating neutral mutations were introduced according to SLiM 4.0 (Haller et al., 2019). The resulting data set thus reflects a total mutation rate of 7.0×10^{-8} with a likelihood of selected to neutral mutations occurring at a ratio of 4:1. After adding neutral mutations, the VCF file with 1000 randomly sampled individuals was created and nonsegregating sites (selected or neutral) were added between SNP positions and randomly assigned as either selected (20%) or neutral (5%) to approximate the 0-fold and 4-fold ratios in the empirical *A. lyrata* data set. The resulting VCF file was used in subsequent analyses with DFE-alpha.

To get a baseline accuracy for DFE-alpha, 10 replicates of 100 individuals (the maximum size supported by DFE-alpha) from the simulated data set were analysed, and the estimation error compared with the known DFE was in each case assessed as the Earth Mover's Distance (see below). To investigate the effects of filtering methods, 15% of the sites in each individual in one set of 100 individuals were masked as missing. This data set was filtered with (1) downsampling at a threshold of 85%, (2) imputation at a threshold of 85% or (3) subsampling at a threshold of 15%. However, the subsampled data set retained no 4-fold SNPs in the SFS after filtering, making DFE estimation impossible. We thus instead sampled four replicates of 4, 8, 12, 16, 20, 24 and 50 individuals from the 15% missing data set, and applied subsampling at 100% (i.e. all sites with missing data were excluded). The same sets of sample sizes were then extracted from the downsampled and imputed data sets to compare the accuracy of the different methods while controlling for the effect of sample size. To directly investigate the effect of sample size and SNP number, 10 replicates of 4, 8, 12, 16, 20, 24, 50 and 100 individuals were extracted from the data sets with no missing data and analysed with DFE-alpha (Figure 4a-d).

With the DFE associated with the simulated data sets being known, the accuracy of estimated DFE was assessed by comparing them to the known DFE using Earth Mover's Distance (EMD) implemented in the *transport* package in R (Schuhmacher et al., 2019). Earth Mover's Distance quantifies the dissimilarity between two distributions as the 'work' required to change one distribution to the other, thus taking into account the amount of overlap. In contrast to the widely used Kolmogorov-Smirnov (KS) distance, EMD is not limited by an upper bound, enabling it to more accurately capture substantial differences between distributions. Additionally, EMD is better suited for gauging distances between distributions with long tails. The EMD was evaluated within the range $-10^5 < s < -10^{-3}$ where s represents the selection coefficient for each mutation, in

FIGURE 4 Accuracy of distribution of fitness effects (DFE) estimations by manipulating SLiM simulated data set. DFE estimates and 95% confidence intervals for 4, 8, 12, 16, 20, 24, 50, and a maximum of either 85 (in downsampling) or 100 (in the other cases) individuals, with either (a) no missing data, or 15% missing data and filtered with either (b) downsampling, (c) imputation or (d) subsampling. (e) DFE estimation error, as represented by Earth Mover's Distance (EMD), in different sample sizes without missing data (black, 10 replicates (n) per sample size), or with 15% missing data and filtered with either downsampling (green), imputation (red) or subsampling (yellow), in four replicates each. (f) DFE estimation error in samples plotted against single nucleotide polymorphism (SNP) number, in data sets without missing data (black) as well as with missing-data filtered by downsampling (green), imputation (red) or subsampling (yellow). Linear regression lines for the no missing data (black) and for all of the filtered data sets combined (brown) are displayed to show the trend of EMD over SNP number (0-fold and 4-fold SNPs) in the two groups. Data sets without missing data include 10 replicates of 4–100 individuals, while four replicates of 4–50 individuals are included for the missing-data filtered data sets.



increments of 10^{-3} . Higher EMD values signify a poorer fit between the estimated and true distribution, thus indicating a less accurate result. The EMD values of each data set was plotted against the number of individuals and SNPs with a regression line to illustrate the relationship.

3 | RESULTS

3.1 | The effect of missing-data treatments on DFE in *A. lyrata*

3.1.1 | Downsampling

The data sets downsampled to 50%, 66% and 75% of the genotypes per site retained 105.7M, 99.5M, and 95.0M sites, respectively, for both *A. lyrata* populations (Table 1). The Austrian data

sets contained 15, 19 and 22 'individuals' and 1.39M, 1.46M and 1.47M SNPs (sum of 0- and 4-fold SNPs) for the three thresholds, while the Norwegian population kept 8, 11 and 12 'individuals' and 374K, 366K and 341K SNPs, respectively. The DFE in the Norwegian data sets differed significantly from that of the Austrian population in that neutral mutations were more frequent (31%–33%), while slightly (8%–9%) and moderately (10%–12%) deleterious mutations were less frequent, but the proportion of strongly deleterious mutations was similar (45%–51%) (Table 1). Additionally, the impact of filter thresholds from 50% to 75% on the three deleterious groups of mutations in the two populations showed inverse patterns, for example strongly deleterious mutations increased with the threshold in the Norwegian population but decreased in the Austrian population. While the estimated DFE varied between populations by 1–10 percentage points under the same method and threshold, it also varied by up to 5 percentage points among the downsampling thresholds within each population.

TABLE 1 Estimated DFE using downsampling, imputation and subsampling procedures in Austrian and Norwegian populations of *A. lyrata*.

Filtering method	Individuals	Total sites	O-fold sites	4-fold sites	SNPs in SFS	DFE [95% CI]: % of mutations with $-N_e s$ values of				
						[0, 1]	(1, 10]	(10, 100]	>100	
Austria	50%	105,746,755	20,081,506	4,532,827	1,389,235	24.63 [24.53–24.73]	10.59 [10.47–10.72]	15.11 [14.87–15.34]	49.68 [49.37–49.96]	
	66%	99,518,927	19,796,632	4,465,916	1,455,146	23.26 [23.15–23.35]	11.73 [11.61–11.80]	17.53 [17.30–17.70]	47.48 [47.26–47.80]	
	75%	95,023,967	19,576,647	4,410,244	1,472,536	21.76 [21.69–22.95]	12.91 [11.45–13.00]	20.30 [17.07–20.48]	45.03 [44.75–48.57]	
Imputation	70%	103,436,893	19,988,452	4,509,075	1,686,431	23.08 [22.97–24.04]	12.39 [11.26–12.51]	18.87 [16.47–19.12]	45.66 [45.35–48.25]	
	80%	97,909,623	19,724,874	4,444,971	1,625,688	22.67 [22.57–22.77]	12.12 [11.99–12.22]	18.44 [18.17–18.65]	46.77 [46.50–47.12]	
	90%	86,272,541	19,088,682	4,267,939	1,437,770	20.21 [20.09–20.33]	13.56 [13.46–13.68]	22.24 [21.99–22.52]	43.99 [43.69–44.27]	
Subsampling	10%	97,383,774	19,593,039	4,364,309	843,938	22.96 [22.80–23.28]	11.21 [10.60–11.43]	16.61 [15.39–17.05]	49.22 [48.61–50.79]	
	15%	76,996,896	18,026,562	3,900,662	874,047	19.78 [19.60–19.90]	14.68 [14.50–14.87]	24.81 [24.40–25.28]	40.74 [40.16–41.24]	
	20%	59,099,749	14,532,777	3,066,767	662,641	20.11 [19.41–20.29]	16.87 [16.62–17.62]	29.21 [28.62–31.15]	33.81 [31.81–34.50]	
Norway	25%	54,974,337	13,445,210	2,824,524	609,256	20.31 [20.10–20.56]	17.29 [16.96–17.56]	29.93 [29.16–30.56]	32.47 [31.72–33.38]	
	50%	105,746,755	20,081,506	4,532,827	374,403	33.13 [32.78–33.45]	9.40 [7.96–10.17]	12.06 [9.86–13.30]	45.41 [43.60–48.80]	
	66%	99,518,927	19,796,632	4,465,916	366,254	32.05 [31.58–32.32]	7.91 [7.42–9.95]	9.86 [9.13–13.07]	50.17 [45.39–51.36]	
Imputation	75%	95,023,967	19,576,647	4,410,244	341,308	30.91 [30.70–31.15]	8.11 [7.72–8.67]	10.24 [9.64–11.10]	50.74 [49.38–51.59]	
	70%	103,436,893	19,988,452	4,509,075	399,078	32.83 [32.58–33.04]	6.51 [6.20–6.96]	7.80 [7.37–8.44]	52.86 [51.89–53.62]	
	80%	97,909,623	19,724,874	4,444,971	365,494	31.62 [31.33–31.89]	6.64 [6.26–7.20]	8.04 [7.49–8.85]	53.71 [52.54–54.49]	
Subsampling	90%	86,272,541	19,088,682	4,267,939	268,958	29.30 [29.04–29.68]	8.06 [7.09–8.41]	10.27 [8.79–10.83]	52.38 [51.49–54.50]	
	10%	109,442,991	20,192,673	4,555,968	248,706	7.26 [7.00–7.48]	86.62 [86.37–86.99]	6.12 [5.89–6.30]	0.00 [0.00–0.00]	
	15%	79,983,985	18,525,720	4,136,084	179,342	28.20 [27.89–28.57]	7.97 [7.46–8.23]	10.22 [9.42–10.64]	53.60 [53.01–54.72]	
Norway	20%	77,586,760	18,343,099	4,091,607	171,711	28.36 [27.78–28.69]	7.55 [7.02–8.83]	9.56 [8.76–11.61]	54.53 [51.62–55.79]	
	25%	77,586,760	18,343,099	4,091,607	171,711	28.36 [27.78–28.69]	7.55 [7.02–8.83]	9.56 [8.76–11.61]	54.53 [51.62–55.79]	

Note: For downsampling, thresholds show the percentage of data retained at each locus. Imputation thresholds signify the data quality (inverse of the max missing rate) of the individuals included in the data set prior to imputation. Subsampling thresholds signify the max missing rates per individual. Total sites include all sites in the VCF file. SNPs in the SFS include only SNPs at 0-fold and 4-fold degenerate sites.

Abbreviations: DFE, distribution of fitness effects; SFS, site frequency spectrum; SNPs, single nucleotide polymorphisms.

3.1.2 | Imputation

The imputed data sets retained all individuals (i.e. 29 Austrian and 16 Norwegian individuals), and 103.4M, 97.9M and 86.3M sites at the 70%, 80% and 90% thresholds, respectively. In the Austrian population, 1.69M, 1.63M and 1.44M SNPs were included, while 399K, 365K and 341K SNPs in the Norwegian population, at the three thresholds, respectively. Increasing the threshold from 70% to 90% only caused 2–4 percentage points of variation in each category of mutations (Table 1). Across both populations, the DFE were stable among imputation thresholds, with the Austrian population displaying slightly larger variance.

3.1.3 | Subsampling

In the subsampling trial, we applied four different thresholds, allowing a maximum of 10%, 15%, 20% and 25% missing genotypes per individual. In the Austrian population, a strict threshold of 10% missing data left eight individuals, 97.4M sites and 844K SNPs in the data set, while a relaxed 25% threshold preserved all 29 individuals with 55.0M sites and 609K SNPs (Note: increasing the missing rate from 20% to 25% only added one more individual) (Table 1). Increasing the missing threshold from 10% to 25% decreased the estimated neutral mutations from 23% to 20%, and the strongly deleterious mutations from 49% to 32%, while the slightly and moderately deleterious mutations increased from 11% to 17% and from 17% to 30%, respectively. Overall, change the threshold from 10% to 15% induced the largest difference in the DFE of all stepwise increases (3–8 percentage points of difference in all categories).

In the Norwegian population, the data set filtered with a missing rate of 10% included only two individuals with 109.4M sites and 249K SNPs. At this level, the DFE was estimated to 7% neutral, 86% slightly deleterious, 6% moderately deleterious and no strongly deleterious mutations. Increasing the threshold to 15% increased the number of individuals to 15, retaining 80.0M sites and 172K SNPs, and shifted the DFE to 28% neutral, 8% slightly deleterious, 10% moderately deleterious and 53% strongly deleterious mutation. Further relaxing the missing rate to 20% and 25% included one more individual (16 total) and had little effect on the DFE compared with the data set filtered at 15% (Table 1). Overall, the Austrian population displayed up to 17 percentage points of difference between thresholds, while the Norwegian population displayed up to 79 percentage points of difference when including the data set filtered at 10% missing data.

3.2 | The effect of sample size and sites on DFE

We subsampled the Austrian population of *A.lyrata* into 4, 8, 12, 16, 20 and 24 individual sets, each containing 211K, 320K, 357K, 426K, 512K and 557K SNPs, respectively, from the complete data set of 29 individuals containing 609K SNPs (Figure 3b). We found

that decreasing the sample size from 29 to 4 substantially increased the proportion of strongly deleterious mutations from 32% to 45%, while it decreased the proportion of slightly deleterious mutations from 17% to 13% and moderately deleterious mutations from 30% to 20%. Neutral mutations changed only slightly (from 20% to 22%) (Figure 3a). The partition of DFE remained stable with sample sizes of 8 and upward (≤ 1 percentage point of fluctuation). The 95% CIs remained similar and narrow (0.5%–4%) in all samples.

In the second trial, we randomly sampled 1K, 10K, 100K, 1M and 10M sites in the 29 individuals (with 55.0M sites, 609K SNPs), resulting in 10, 109, 1115, 11.1K and 111K SNPs, in each data set, respectively. We found that the DFE estimates became increasingly unstable with decreasing the number of sites: the data sets with fewer than 1M sites (11.1K SNPs) showed a large variation in DFE values (8–50 percentage points; Figure 3b). Notably, a decrease in the number of sites brought a simultaneous increase of the width of the 95% CIs, in a manner not seen when decreasing the numbers of individuals (Figure 3a vs. b). At 1K sites (10 SNPs), the 95% CIs for the three deleterious categories covered 98%–100% of the entire range of possible values, indicating low confidence in where the true values lie. At 10K sites (109 SNPs), the CIs shrunk but were still large, covering between 34%–71% of the possible values. On average, each 10-fold decrease in the number of sites increased the size of the bootstrapped 95% CIs 2.5 times.

In the third trial, we examined the effect of sites in a small sample of four individuals. The sites chosen were the same as those in the second trial, although the set of 1K sites included too few SNPs to be evaluated and was not shown in Figure 3c. The data sets with 10K, 100K, 1M, 10M and all 55.0M sites had 43, 391, 3821, 38.6K and 211K SNPs, respectively. At 10K sites, the DFE in the 4-individual set was drastically different from the 29 individuals. Furthermore, the 95% CIs of neutral, and slightly and moderately deleterious mutations increased by 18%–81% in the 4-individual relative to the 29-individual data set. The CI for strongly deleterious mutations shrank somewhat in the 4-individual data set but was still large and spanned 66% of the range of possible values. The DFE estimates at 100K sites and above in 4-individual data sets were very similar (≤ 1 percentage point of difference) to the second trial using 29 individuals (Figure 3c vs. b), but the 95% CIs approximately doubled for the three classes of deleterious mutations.

3.3 | Accuracy of DFE-alpha in SLiM simulated data

To determine which missing-data treatment and sample sizes produced the smallest error and thus approximated the true DFE most accurately, we conducted SLiM simulations with a known DFE. The simulation produced a data set with 1000 individuals and 29,944 SNPs. Using 10 replicate samples of 100 individuals, each containing ~15,500 SNPs, the DFE was estimated to 29%–31% neutral, 8%–10% slightly deleterious, 10%–13% moderately deleterious and 48%–52% strongly deleterious mutations; the true DFE should be approximately 30% neutral, 9% slightly deleterious, 11% moderately

deleterious and 50% strongly deleterious mutations, meaning an error of $\pm 1\%$ – 2% can be expected with this data set in optimal conditions. The β and E_s parameters of the gamma distributions were estimated to 0.097–0.128 and -276 – 33 , respectively, yielding error values ($\text{EMD} \times 10^7$) of 3.5–20.5 (Figure 4e). These values are used as reference for the 'maximum' accuracy of DFE-alpha for the simulated data set.

To evaluate the effect of filtering methods, we masked 15% of the genotypes per individual as missing and excluded all missing sites in samples of 4, 8, 12, 16, 20, 24 and 50 individuals (4 replicates of each), which mimics the effect of subsampling at different thresholds. In order to compare these results to downsampling and imputation, the same sample sizes were extracted from the downsampled and imputed data sets created at 85% threshold from the full data set. At a sample size of four individuals, all three methods performed roughly equally well (average EMD was 33.7, 36.4 and 36.5 for downsampling, imputation and subsampling, respectively, Figure 4b–e, Table S2), but subsampling tended to slightly underestimate the proportion of slightly and moderately deleterious mutations (by up to 5 and 7 percentage points, respectively), and overestimate strongly deleterious mutations (by up to 11 percentage points). Downsampling gave the most accurate results based on the average EMD across all sample sizes above eight individuals (Figure 4b,e). Imputation performed slightly worse in all samples except eight individuals (Figure 4c,e). Both downsampling and imputation produced results within 1–3 percentage points of the range of the reference set at all sample sizes above four individuals. Subsampling, however, produced highly variable and noticeably less accurate results even at higher sample sizes (Figure 4d,e). For example, the four replicates of 24 individuals produced EMD values between 3.7–18.8 for downsampling, 12.3–47.4 for imputation and 31.2–112.8 for subsampling (Table S2). We found that subsampling produced the most accurate results at an intermediate sample size (e.g. 16 individuals; EMD from 1.7 to 56.1) and became less accurate at sample sizes where fewer SNPs were retained (e.g. 50 individuals with 5 SNPs remaining; Figure 4b, Table S2).

Our simulated data verified the trends observed in the empirical data, showing that increased sample size correlated with lower error in DFE estimates when the number of SNPs is not a limiting factor. In the data sets of 4, 8, 12, 16, 20, 24 and 50 individuals (10 replicates of each) with no missing genotypes, the EMD values were the largest in samples of 4 and 8 individuals, stabilized around 12–24 individuals, and then decreased further in 50 individuals to a level similar to that in the 100 individuals (Figure 4e). Linear regression in these data sets showed that DFE estimation error (EMD) was negatively correlated with number of individuals ($p = .00179$, $R^2 = .1182$), and even more strongly correlated with the number of SNPs in the data set ($p = 6.38 \times 10^{-6}$, $R^2 = .2311$) (Figure 4f). An even stronger negative correlation between EMD and SNP number was seen when the four replicates of 4–50 individuals from the downsampled, imputed and subsampled data sets were analysed with a joint linear regression ($p = 1.11 \times 10^{-9}$, $R^2 = .3658$) (Figure 4b). Data sets with few SNPs also displayed larger 95% CIs while the number of individuals had a

minor effect on CI size (Figure 4a–d, Table S2), similar to what was observed in the empirical data sets.

In summary, applying different filtering methods and thresholds affected the final data matrix size (number of individuals and SNPs) and subsequent DFE estimates. Imputation and downsampling produced similar and less variable DFE results than subsampling, and downsampling appeared more accurate than imputation for the simulated samples used. Furthermore, higher numbers of individuals and SNPs both increased accuracy of the results, especially at very low sample sizes (4–8 individuals, <5000 SNPs).

3.4 | The effect of population structure on DFE

The PCA of the 45 samples from Austria and Norway showed a distinct separation of the two populations along PC1 (which explained 24.7% of the total genetic variance), and separation of the Austrian population into four visible clusters along PC2 (which explained 7.3% of the total genetic variance) (Figure S1). The weighted F_{ST} between the two populations was 0.228, while the F_{ST} among the four Austrian clusters was relatively small at 0.073. To understand the effect of merging genetically distinct populations on the estimated DFE, we created 12 merged populations with contributions of 10 or 15 individuals from Austria and Norway, with three subsets of each population (Figure 1c). We then calculated the weighted F_{ST} between the contributing subsets to evaluate how the degree of population stratification in a sample affects the joint DFE estimate. We first examined the DFE in the unmerged replicate samples of 10 and 15 individuals from the two populations. Among the replicates of 10 individuals from the Austrian population, a maximum difference of 2, 3, 7 and 6 percentage points were observed in the neutral, slightly, moderately and strongly deleterious mutations. By comparison, no mutation category varied by more than 2 percentage points in the samples of 15 individuals. Comparably stable DFE estimates were observed in the Norwegian samples, with variation in the range of 0, 2, 3 and 4 percentage points for the four categories of mutations in samples of 10 individuals, and less than 1 percentage point of a difference among replicates of 15 individuals (Figure 4a). However, the DFE estimates were markedly different between the two geographical populations, for example neutral mutations shifted up by an average of 9 percentage points while the slight and moderate mutations shifted downwards in Norway compared with Austria. The estimated proportions of strongly deleterious mutations were similar in the two populations.

With this population-specific DFE in mind, we then examined the differences between the merged samples and their respective contributing single population subsets. In most cases, the estimated DFE values for the merged samples were in-between the DFE estimates of the contributing subsets, but not always perfectly intermediate (Figure 5a). The estimated weighted F_{ST} values between the pairs of contributing subsets ranged from 0.218 to 0.263 (mean F_{ST} between 0.085 and 0.131). These estimates are largely in line with previous studies, where mean F_{ST} across European populations

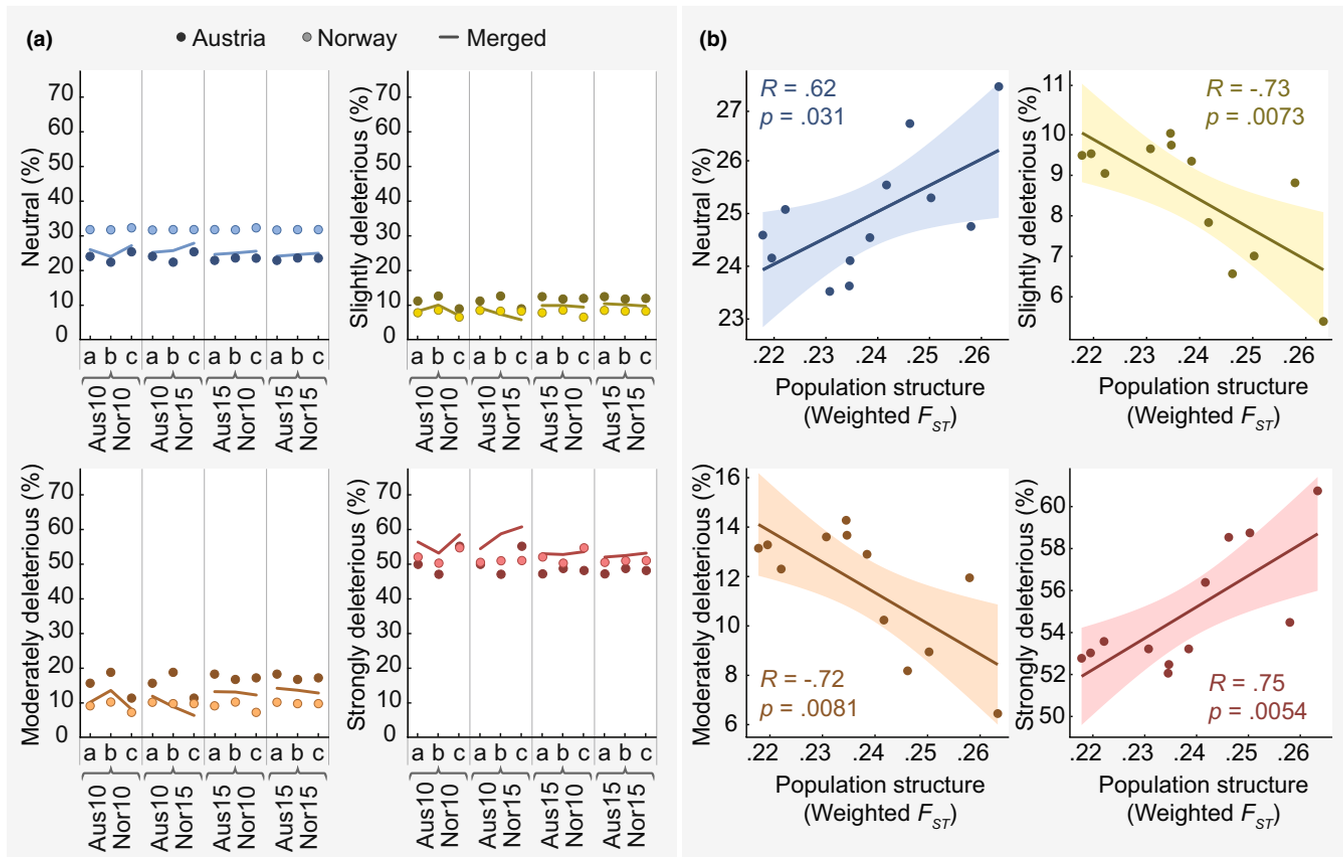


FIGURE 5 Effect of population structure on distribution of fitness effects (DFE). (a) The estimated DFE of the Austrian (dark dots) and Norwegian (light dots) samples of *Arabidopsis lyrata*, compared to merged samples (solid lines) containing both groups in different combinations. The relative sample size from each population is listed along the horizontal axis (bottom), as well the name of each of three replicates (top). (b) Linear regression of the estimated proportion of each of the four mutational categories of the DFE over the F_{ST} between the merged samples, with 95% confidence intervals shown in shaded areas.

of *A. lyrata* ranges between 0.06 and 0.09 (Marburger et al., 2019). Plotting the weighted F_{ST} against the estimated DFE in the merged populations showed an apparent relationship (Figure 5b). Using linear regression, F_{ST} was correlated with the proportion of neutral ($R = .62$, $p = .031$), slightly ($R = -.73$, $p = .0073$), moderately ($R = -.72$, $p = .0081$) and strongly deleterious mutations ($R = .75$, $p = .0054$). These results show that population structure had a significant effect on the DFE, with higher F_{ST} potentially driving up the estimated proportion of neutral and strongly deleterious mutations and reducing the estimates of the less deleterious classes.

4 | DISCUSSION

4.1 | Methods of missing-data treatment affect DFE results

Missing-data treatment is the first step in any genomics analyses. Using simulated data with a known DFE, we were able to evaluate the accuracy of different filtering methods in recovering the true DFE. We found the data set with no missing data produced the most accurate result, followed by downsampling, then imputation, and

then subsampling. The number of SNPs in the downsampled and imputed data sets were similar in all samples, suggesting that any difference in performance between the two methods is likely due to imputation affecting the shape of the SFS in a nonrandom manner. The assumption that deleterious mutations appear as low-frequency alleles in the SFS, in combination with the relatively small sample sizes used in the tests, makes an SFS-based analysis highly reliant on those low-frequency categories, especially singleton SNPs. This could explain our result, as imputation of low-frequency alleles display much higher error rates than higher frequency alleles in imputation procedures (Pook et al., 2020).

Filtering with subsampling produced the least accurate estimates on average. Since increasing the number of individuals in the subsampled data set decreases the number of sites, this filtering method's performance is thus affected by sample size in two ways, both the number of individuals and the number of SNPs available. This effect is expected to be especially strong in data sets where the distribution of missing data is random (as was the case in our simulated data sets), where a highly dissimilar pattern of missing data across individuals excludes a large number of sites by subsampling. This pattern was not as strong in the empirical data sets where the distribution of missing data across individuals was more similar, but still present.

Thus, intermediate sample sizes of individuals are preferable for this method.

The array of tested filtering thresholds on the empirical data sets corroborated the trend and conclusions drawn from the simulated data sets. The empirical data sets proved to be more sensitive to minor changes in filtering thresholds as even slight adjustments resulted in significantly different outcomes in some cases. The DFE estimates in the subsampled data sets were unpredictable, both within and among populations. This is most likely a result of substantial downsizing of the data matrix, since the total number of sites and SNPs were reduced by 50%–90% in the subsampled data sets compared with the other two methods. Downsampling and imputation produced results with similar levels of variation across the different thresholds. With the simulation results in mind, it could be argued that both methods are equally valid in this case as long as sample sizes are satisfactory, and the choice between them might depend on other conditions and computational resources. As a general rule, we recommend filtering data with several thresholds to obtain an overview of the variability produced by each method. This is especially important because the 95% CIs do not provide information about whether the filtered and subsampled data set is representative of the initial population and, as we show in this study, the differences among subsets of samples from the same population can be significant.

A cursory review of recently published DFE estimation studies shows that downsampling is the most frequently used of the three methods tested here (see Castellano et al., 2019; Chen et al., 2020; Gossmann et al., 2010; Liang et al., 2022; Takou et al., 2021). This is not surprising, since downsampling is considerably faster than imputation, yet retains more data than subsampling. Imputation methods require high-quality data sets from the outset to be able to make reliable predictions; data sets with high rates of missing sites and low levels of genome-wide linkage disequilibrium are not ideal for this treatment. With low levels of genome-wide linkage disequilibrium, the presence/absence of any given SNP is mostly uncorrelated with the presence/absence of any other SNP, meaning that there are no patterns of linkage disequilibrium among sites from which imputation can accurately predict the state of a missing site. In such cases, downsampling might be a better choice. With the current rate of improvement in both genome-wide sequence data and computing power, however, we predict an increasing popularity of imputation as a data processing method in DFE estimation and other population genomics analyses. We recommend prefacing any missing-data treatment with an analysis of the prevalence of missing sites and the level of linkage disequilibrium to determine whether imputation is the appropriate method for each data set.

4.2 | Very small sample sizes skew the estimated DFE

A review on DFE estimated in 139 plant and animal species (Chen et al., 2017), each with between 2 and 50 chromosomes sampled,

shows very different DFE distributions. We evaluated the effects of the number of sampled individuals on the estimated DFE when the number of sites was not a limiting factor. We found that DFE estimated from few individuals (<8) were strongly skewed compared with larger sample sizes. In simulated data sets with no missing data, the accuracy of the estimated DFE was highest in the largest sample (100 individuals) and lowest in the smallest samples (4 and 8 individuals), and the samples with >8 individuals displayed markedly improved accuracy of DFE estimates. Similarly, DFE estimates based on four individuals produced the least accurate results using both downsampling and imputation for missing-data treatment.

In the empirical trials, DFE estimates between random sets of four individuals were rather unstable in the Austrian population. In the Norwegian data set subsampled at 10% that kept only two diploid individuals, the proportion of slightly deleterious mutations was greatly overestimated compared to that of the full population size. Results stabilized with a sample size of 8 or more, which is consistent with the findings from the simulated data sets. This suggests that a relatively small number of individuals is needed for reliable DFE estimates when there are many sites available, but that very limited sample sizes increases the risk of producing nonrepresentative results. We thus deem the potential effects of low sample size to be alarming due to their unpredictable and stochastic nature, and caution against using sample sizes below four diploid individuals (eight haploids).

4.3 | Limited sites cause high variability in DFE results

Reducing the number of sites resulted in highly variable and unpredictable DFE estimates even with larger sample sizes. Overall, the negative correlation was observed between the number of SNPs and EMD values in the simulated data sets indicates that the accuracy of SFS-based DFE estimation is limited by the number of SNPs available. This trend was also observed in the empirical data, where estimates based on 1M, 10M and 55M sites in 29 individuals all looked similar, but using 1K–10K sites (59–571 SNPs) produced highly dissimilar results, demonstrating the importance of having a sufficient number of sites and SNPs for reliable SFS-based analyses. The DFE is estimated from SFS, that is the distribution of SNPs of different frequencies in the population. Thus, the number and specific subset of SNPs directly affect the resolution to which we can estimate the shape of the DFE. This would explain why the 95% CIs increased in size as the number of sites decreased. At 1K–10K sites, the confidence intervals spanned the entire range of possible values for several of the mutational categories (Figure 3b). For these data sets, we are therefore left with no confidence that our predicted DFE is close to the true DFE. If the CIs are ignored, the very different DFE estimates from subsets of the same data set could lead to different interpretations of the selection pressures acting on the population. This result illustrates a clear type 1 error; the estimated DFE from our samples of 1K, 10K and 100K sites are not representative of the

full set of sites and produce incorrect inferences that imply differences in the underlying DFE, despite being random subsets of the same data set.

Based on both the empirical and simulated trials, we conclude that DFE estimates of DFE become stochastic and unpredictable with very small number of sites/SNPs, and accuracy is expected to increase significantly with the number of SNPs included; at least 5K SNPs are required to obtain reliable DFE estimates using DFE-alpha.

4.4 | Population structure may skew DFE estimates

By combining samples from the Austrian and Norwegian populations into merged populations, we were able to see how the composition of populations affects DFE estimates. One trend was immediately clear: the estimated proportion of strongly deleterious mutations was higher in the merged populations than in the contributing single population subsets. A high F_{ST} may skew the DFE towards higher estimated proportions of neutral and strongly deleterious mutations and lower proportions of slightly and moderately deleterious mutations. This correlation may not be conclusive, but it indicates that population structure can indeed affect DFE and should be taken into consideration when performing these analyses at a species level. Studies on DFE often include multiple or combined populations to gain a global estimate that characterizes the organism or species (Chen et al., 2017; Hämälä & Tiffin, 2020; Slotte et al., 2010; Zhao et al., 2020). We cannot presently state that pooled samples will always skew the inferred DFE, but it is advisable to estimate the DFE separately in individual populations, as well as from pooled samples to evaluate any deviations caused by pooling that might inform conclusions drawn from the results. A recent study developed a joint DFE approach that enables the analysis of pairs of populations (Huang et al., 2021), which could be practical in examining variance of DFE among populations.

5 | CONCLUSIONS

Accurate estimation of DFE from genomic data hinges on several factors, including the number of sampled individuals, the availability of sites and SNPs, and the approach employed to address missing data. Our study, which utilized both empirical data and forward simulations, explored all these aspects and offers guidance for experimental design of DFE estimation studies. We found that down-sampling is a dependable method of handling missing data, though it may still impact the DFE to some extent. Imputation, while generally accurate, may be less suitable for small samples (≤ 100 individuals, $< 10K$ SNPs) or when genome-wide linkage disequilibrium is very low (as is often the case with highly outbreeding species). We demonstrated that DFE estimates derived from data sets with ≤ 4 diploid individuals or $\leq 5K$ SNPs may be unreliable due to the risk of sampling error and the limited amount of information in the SFS. Furthermore,

strong population structure within samples can potentially skew DFE estimates.

More advanced methods of DFE estimation employ an unfolded SFS, where each SNP is categorized as ancestral or derived based on an outgroup reference genome. While model species can benefit from these sophisticated techniques, most studies must still rely on methods utilizing the folded SFS, and frequently deal with limited sample sizes. Given the extensive body of previously published work employing folded SFS, it is imperative to be able to understand the expected accuracy of DFE estimates in comparative analyses. This study highlights the factors that should be considered when interpreting DFE estimates, thereby enhancing the reliability and relevance of future research.

AUTHOR CONTRIBUTIONS

WZ and XRW designed the empirical study. All authors contributed to designing the simulation study. BH provided support for simulations in SLiM 4.0. BA and WZ performed empirical data analyses. ÅB provided statistical advice. BA, WZ and XRW wrote the manuscript draft. All authors contributed to the revision of the manuscript.

ACKNOWLEDGEMENTS

Genomic data processing and analyses were performed using resources provided by the Swedish National Infrastructure for Computing (SNIC), through the High Performance Computing Centre North (HPC2N). This study was supported by grants from the Swedish Research Council (VR) and T4F program to XRW.

CONFLICT OF INTEREST STATEMENT

The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT

All sequencing data are retrieved from the NCBI SRA database with accession numbers listed in Table S1. Procedures associated with the SLiM simulations are provided to GitHub repository at <https://github.com/beangelica/DFE-filtering>.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Andersson, B. A., Zhao, W., Haller, B. C., Brännström, Å., & Wang, X.-R. (2023). Inference of the distribution of fitness effects of mutations is affected by single nucleotide polymorphism filtering methods, sample size and population structure. *Molecular Ecology Resources*, 23, 1589–1603. <https://doi.org/10.1111/1755-0998.13825>