

Nucleotide Diversity Loss on a Plant Y Chromosome Following Recent Recombination Suppression

Josh Hough^{*,†,1}, Wei Wang[†], Spencer C.H. Barrett[†] and Stephen I. Wright[†]

^{*}Department of Plant Sciences, University of California, Davis, [†]Department of Ecology and Evolutionary Biology, University of Toronto

ABSTRACT X and Y chromosomes differ in effective population size (N_e), rates of recombination, and exposure to natural selection, all of which can affect levels of genetic variability within the genome. On Y chromosomes with suppressed recombination, selection is expected to remove linked neutral variation and reduce the N_e of Y compared to X chromosomes or autosomes. However, non-selective factors including female biased sex ratios and high variance in male reproductive success can also reduce Y-linked N_e , making it difficult to infer the causes of low Y diversity. Here, we investigate the factors affecting X- and Y-linked neutral polymorphism during plant sex chromosome evolution in *Rumex hastatulus* (Polygonaceae), a dioecious annual with recently evolved sex chromosomes. Strikingly, we find that neutral diversity for genes on the Y is on average ~2.1% of the value for their homologues on the X, corresponding to a chromosome-wide reduction of ~93% compared to the neutral expectation. We demonstrate that the magnitude of this diversity loss is inconsistent with a reduced male N_e caused by neutral processes including female-biased sex ratios and high variance in male reproductive success. Rather, using forward population genetic simulations and estimates of the distribution of fitness effects of deleterious mutations, we show that Y diversity can be explained by purifying selection removing linked neutral variation, although selective sweeps may have also contributed. Given the recent origin of *R. hastatulus* sex chromosomes, our study suggests that Y chromosome degeneration in the early stages occurs through strong selective interference among many linked sites rather than primarily through Y-inactivation followed by neutral genetic drift.

KEYWORDS Sex Chromosome Evolution; Nucleotide Diversity; Recombination; Deleterious Mutations

Introduction

Morphologically distinct sex chromosomes have evolved multiple times independently in both plant and animal kingdoms (Westergaard 1958; Ohno 1967; Bull 1983; Charlesworth 1991). Despite clear biological differences between these kingdoms, X and Y chromosomes in both lineages have undergone similar genetic changes. For example, in both groups the loss of recombination between X and Y chromosomes has been associated with an accumulation of deleterious mutations and a gradual loss of genes from the Y chromosome (Hough *et al.* 2014; Bergero *et al.* 2015; Bachtrog 2013), and in some species, this genetic degeneration of the Y has led to the evolution dosage compensation of the X (Charlesworth 1996;

Muyle *et al.* 2012; Mank 2013; Papadopoulos *et al.* 2015). The independent evolution of these phenomena across a broad range of distantly-related species suggest that general evolutionary mechanisms may be involved, but inferring the causes of molecular evolution and patterns of polymorphism in genomic regions that lack recombination is a longstanding challenge for both theoreticians and experimentalists (Charlesworth 1978; Feldman *et al.* 1980; Barton 1995; Charlesworth 1996; Otto and Feldman 1997; Charlesworth and Charlesworth 2000; McVean and Charlesworth 2000).

One fundamental difference between the X and Y chromosomes is that there are 1/3 as many Y-linked gene copies as X-linked ones in a diploid population. Because of this, the effective population size (N_e) of the Y chromosome is expected to be 1/4 that of the autosomes, whereas the N_e for the X chromosome should be 3/4 that of autosomes (assuming an equal number of reproducing females and males). The lowered N_e of the Y chromosome implies that deleterious mutations are

more likely to rise to high frequency because of genetic drift (Nei and Tajima 1981) - an effect that can be exacerbated by selective interference among genetically linked sites (the "Hill-Robertson Effect"). These effects both predict a reduction in the level of neutral polymorphism maintained on the Y chromosome, which at equilibrium is proportional to the product of N_e and the mutation rate, μ (Kimura 1984; Charlesworth *et al.* 1987).

In species with female-biased reproductive sex ratios or extensive male-male competition, a high variance in male reproductive success can also reduce the N_e of genes on the Y chromosome (Caballero 1995; Charlesworth 2001; Laporte and Charlesworth 2002; Pool and Nielsen 2007; Ellegren 2009), suggesting that inferences about the effects of linked selection on the Y need to be distinguished from neutral models that include sex ratio variation as well as male-female differences in reproductive success. High variance in male reproductive success, for example, is predicted to reduce both Y-linked N_e and autosomal N_e (Kimura and Crow 1964; Nomura 2002), and therefore evidence for this effect can be obtained by comparing levels of neutral variability on X and Y chromosomes relative to the autosomes. In particular, high variance in male reproductive success predicts a reduction in the Y/A diversity ratio, but an increase in the X/A ratio. Based on such comparisons, studies in humans have found examples of historical sex-differences in dispersal during human migration (Wilkins 2006), and the inflated X/A ratio has been suggested to reflect a historical excess of breeding females over males (Hammer *et al.* 2008), though there is considerable variation in X/A estimates in human and other demographic scenarios have also been suggested (Bustamante and Ramachandran 2009; Hammer *et al.* 2010).

Despite widespread interest in determining the effects of selection patterns of sex-linked variability diversity (Ellegren 2011; Bachtrog 2013), we still know very little about how linked selection has affected patterns nucleotide polymorphism. In humans and *Drosophila* (McAllister and Charlesworth 1999; Bachtrog and Charlesworth 2000), where large population genomic data sets are readily available, studies examining Y chromosome variability have revealed that Y-linked diversity is considerably lower than predicted under models of neutral evolution (Hellborg and Ellegren 2004; Bachtrog 2013; Wilson Sayres *et al.* 2014). However, it remains unclear whether such inferences about the effects of linked selection on the Y should also apply to systems in which sex chromosomes evolved *de novo* from autosomes much more recently. The effects of diversity loss due to selection might be expected to be lower in such systems because of a shorter history of recombination suppression, but contrary to this, simulations reveal that background selection and selective sweeps both have strongest effects during the early stages of Y-chromosome evolution, before Y-chromosomes have lost the majority of their genes (Bachtrog 2008). Indeed, studies of Y-degeneration in the young plant sex chromosomes *R. hastatulus* and *Silene latifolia* have both revealed signs of rapid Y chromosome degeneration (Hough *et al.* 2014; Papadopoulos *et al.* 2015; Charlesworth 2016). In these cases, if the early stages of Y-chromosome evolution involved widespread Y-linked gene-inactivation (silencing) (Orr and Kim 1998), then it is possible that the strength of linked selection would then be minimized such that gene loss was mostly driven by gene-silencing followed by neutral genetic drift (Bachtrog 2013) rather than selective interference due to widespread linkage disequilibrium.

To investigate the factors affecting nucleotide diversity in the early stages of sex chromosome evolution we analyzed neutral polymorphism levels on X, Y, and autosomal chromosomes in the plant *R. hastatulus* (Polygonaceae). This species is a dioecious annual with highly heteromorphic X and Y chromosomes that are estimated to have evolved approximately 15 MYA (Quesada del Bosque *et al.* 2011; Grabowska-Joachimik *et al.* 2015; Navajas-Pérez *et al.* 2005), making sex chromosomes in this species over 100 million years younger than the well-studied mammalian sex chromosomes (Lahn and Page 1999; Ross *et al.* 2005). *Rumex hastatulus* has also received particular attention because of the unique occurrence of an intraspecific sex chromosome system, in which both XY and XY₁XY₂ males occur in geographically distinct 'chromosomal races' (Smith 1963). The XY₁XY₂ sex chromosome system in this species (the North Carolina race) is thought to have originated through an X-autosome fusion, with the XY system (the Texas race) maintaining the ancestral chromosome complement (Smith 1964). Despite the recent origin of sex chromosomes in both races, a recent study revealed that both the ancestral and neo-Y chromosomes have undergone significant gene loss and functional deterioration (Hough *et al.* 2014). In order to simplify our comparison of polymorphism levels within the genome, we focus on the ancestral Y chromosome that occurs in both sex chromosome races.

Of particular relevance to the present study, *R. hastatulus* populations have been found to consistently exhibit female-biased reproductive sex ratios, with a mean sex ratio of $N_f / (N_m + N_f) = 0.6$ (Pickup and Barrett 2013). Female-biased sex ratios in dioecious plants with sex chromosomes are not uncommon (refs), and their occurrence in *Rumex* provides an excellent opportunity to study both the demographic and selective effects contributing to sex-linked variability. Here, using genome-wide estimates of polymorphism together with forward population simulations, we test whether levels of neutral diversity on sex chromosomes and autosomes in this species are consistent with neutral evolutionary processes, including the joint effects of sex ratio bias and high variance in reproductive success, or whether the removal of linked neutral variation caused by selection on the Y has led to greater loss of diversity than expected under neutrality.

Materials and Methods

Population Samples and Sex-Linked Genes

We analyzed sex-linked and autosomal genes identified from Illumina RNA sequence data from 24 individuals (12 males and 12 females, with 1 male and 1 female from each of 12 populations; 6 populations of each sex chromosome race). Samples were collected in 2010 from throughout the native range of *R. hastatulus* (locations in Table S1), and plants were grown in the glasshouse from seeds collected from open-pollinated females. We extracted RNA from leaf tissue using Spectrum Plant Total RNA kits (Sigma-Aldrich). The isolation of mRNA and cDNA synthesis was conducted according to standard Illumina RNAseq procedures, with sequencing conducted on two Illumina HiSeq lanes with 150-bp end reads at the Genome Quebec Innovation Center. Reads from these 24 samples were mapped to the *R. hastatulus* reference transcriptome (Hough *et al.* 2014), and raw sequences are available from the GenBank Short Read Archive under accession no. SRP041588. Reads were mapped using the BurrowsWheeler Aligner (Li and Durbin 2010), followed by Stampy (Lunter and Goodson 2011).

We used Picard tools (<http://picard.sourceforge.net>) to modify mapping output into the format required for the Genome Analysis Toolkit (McKenna *et al.* 2010) variant calling software, and subsequently removed genes with low coverage (<10x) and Phred Quality Scores <20. The population samples analyzed here were previously reported in Hough *et al.* (2014), where they were used to validate the ascertainment of sex-linked genes identified through segregation analysis. Here, to consider sex linked genes that were identified in each of our sequenced population samples, we focused on the set of 460 X/Y genes for which the Y homolog was found in both the Texas and North Carolina races (i.e., X/Y genes where the Y copy was inferred to be on the Y₁ chromosome).

Autosomal Genes

In evaluating evidence for nucleotide diversity differences between X and Y chromosomes, it is important to distinguish between reduced Y-linked diversity, and the possibility that X-linked diversity is elevated above the level predicted from a neutral model. To do this, we normalized sex-linked diversity estimates by autosomal diversity, and compared empirical X/A and Y/A nucleotide diversity ratios to those predicted from neutral models and from simulations (described below). Because the criteria for identifying autosomal loci in Hough *et al.* (2014) were based on the occurrence of four segregating SNPs per locus, this set of genes is probably higher in diversity than the average autosomal gene. Therefore, for the present analysis we incorporated the larger set of all non-sex linked (putatively autosomal) genes from our transcriptome data. We also filtered this set to remove genes that may have been sex-linked but were not identified as such by Hough *et al.*'s conservative ascertainment criteria. In particular, we removed: (i) any genes in which there was evidence for at least one SNP with a sex-linked segregation pattern, (ii) any genes with SNPs showing fixed heterozygosity in males and fixed homozygosity in females, (iii) any genes with less than 10X coverage or greater than 100X coverage from independently obtained genomic coverage data (to filter out duplicate genes or those with highly repetitive sequences), and (iv) any genes containing SNPs with large (>0.4) allele frequency differences between males and females. Finally, we removed genes with fewer than 100 synonymous sites following this filtering to avoid biasing our results toward genes that may have been particularly short due to assembly problems. This filtering resulted in a set of 12,356 and 11,350 autosomal genes in the Texas and North Carolina races, respectively.

Phasing X and Y alleles

To estimate polymorphism for the X and Y sequences separately, it is necessary to infer the phase of SNPs in sex-linked transcripts in males. In previous work, phasing alleles on *R. hastatulus* sex chromosomes was achieved using segregation analysis from a genetic cross. Here, to phase SNPs from population samples where such segregation data was unavailable, we used HAPCUT (Bansal and Bafna 2008), a maximum-cut based algorithm that reconstructs haplotypes using sequenced fragments (Illumina read data) from the two homologous chromosomes to output a list of phased haplotype blocks containing the SNP variants on each chromosome. Because the resulting haplotype blocks produced by HAPCUT contained SNPs that were phased relative to each other, but not designated to either the X or Y chromosome, we assigned individual variants to X

or Y by independently identifying fixed X-Y differences with each haplotype block (i.e., sites in which all females were homozygous, and all males were heterozygous). Identifying such fixed differences within phased haplotype blocks enabled us to then infer the correct phase (X or Y) of the polymorphisms from HAPCUTs output. This was done by matching the phase of fixed X-Y differences with their neighboring polymorphic sites: when a fixed X-Y difference occurred in the same phased haplotype block as a polymorphic site, the polymorphic variants in that block were assigned to either X or Y based on the known phase of the fixed difference with which they were matched. SNPs that were identified outside of phased blocks, or in blocks without fixed X-Y differences, were recorded as missing data. Finally, we filtered out SNPs with coverage > 60, QUAL score > 60, and those within a distance of 10bp or less from indels. This procedure was conducted using a combination of Perl and Bash scripts, and resulted in fasta-formatted alignments of X and Y sequences for 372 sex-linked genes from the 24 individuals in our study.

We further validated the results of HAPCUTs allele phasing by comparing the accuracy of this method with the phasing-by-segregation method that was conducted in Hough *et al.* (2014). To do this, we first phased the sequence data from parents and their progeny using HAPCUTs algorithm (using the same parameters as for the population data), and then identified cases where SNPs were inferred on the Y chromosome by HAPCUT, but where the true level polymorphism was known to be zero. We identified 7 percent of sex-linked genes with phasing errors of this kind, or which were otherwise determined to have genotyping errors resulting in false SNP calls. This corresponds to a SNP error rate estimate of 1.7×10^{-4} . Note that this rate is very low relative to population-based estimates of polymorphism on the X and autosomes (Table 1), and therefore should have minimal effects on our estimation of X/A polymorphism. However, because the rate is high relative to the expected level of true polymorphism on the Y-chromosome, we further filtered genes in which we found evidence for false-positive Y-polymorphisms arising from: (i) phasing errors caused by gene duplicates (more than two haplotypes), (ii) polymorphisms around indels, and (iii) genotyping errors caused by low Y-expression. This filtering was done by manually inspecting sequences in IGV (Robinson *et al.* 2011) and identifying each individual putative polymorphism on the Y chromosome.

Estimating nucleotide diversity on sex chromosomes and autosomes

For each locus in our analysis, we calculated Wattersons (1975) estimator of the population parameter $\theta = 4N_e\mu$, where N_e is the effective population size, and μ is the mutation rate (Watterson 1975), using a modified version of the Perl program Polymorphura (Bachtrog and Andolfatto 2006). To compare sex-linked and autosomal loci, we calculated the average value of θ , weighted by the number of synonymous sites in each gene (Figure 2; Table 1). We obtained 95 percent confidence intervals for our estimates of the X/A and Y/A diversity ratios by bootstrapping per gene using the BCa method (Efron and Tibshirani 1994) implemented in the Boot package in R (Canty and Ripley 2012), and calculating X/A and Y/A diversity on each iteration for 20000 replicates each. Bootstrapping was conducted on the final filtered set of 173 sex-linked, and 12355 autosomal genes from the Texas race, and separately for the 176 sex-linked and 11349 autosomal genes from the North Carolina

race. Note that the lack of recombination on the Y chromosome implies that assumptions about independence across loci are violated, suggesting that the true uncertainty in our Y/A diversity estimate may be wider than implied by bootstrapping across genes. To address this issue, we also used a maximum likelihood approach, implemented in a modified version of the MLHKA software (Wright and Charlesworth 2004) to independently estimate a credibility interval for the Y/A ratio using coalescent theory (Figure S1). Because of the thousands of genes involved, a likelihood method incorporating divergence to control for heterogeneity in mutation rate is not feasible, since this involves maximizing the likelihood estimate of the mutation rate for each locus independently. Therefore, we assumed no heterogeneity in mutation rate, no recombination between Y-linked genes, and free recombination between autosomal loci. Our model therefore had two parameters; $\theta_{\text{autosomal}}$ and the Y/A ratio. We varied both parameters across a grid of points and evaluated the likelihood, where the Y/A ratio varied from 0.001 to 1, and $\theta_{\text{autosomal}}$ varied from 0.001 to 0.01.

Neutral predictions and the effect of sex ratio bias on diversity

To test whether the levels of diversity we observed on X, Y and autosomal chromosomes could be explained by the occurrence of biased reproductive sex ratios or high variance in reproductive success, we compared our empirical diversity estimates to predictions from a neutral model that jointly considers these two effects.

In particular, the effective population size for genes on each chromosome can be given by (Wright 1931):

$$N_{eA} = \frac{4N_m N_f}{N_m + N_f} \quad (1)$$

$$N_{eX} = \frac{9N_m N_f}{4N_m + 2N_f} \quad (2)$$

$$N_{eY} = \frac{N_m}{2} \quad (3)$$

Using these predictions, we calculated the expected X/A and Y/A ratios of diversity for the estimated *R. hastatus* sex ratio ($N_f/N_m + N_f$) = 0.6 (Pickup and Barrett 2013), assuming that the level of neutral polymorphism is given by the product of the mutation rate and the effective population size, $\theta = 4N_e\mu$ (Watterson 1975). Sex-specific variance in reproductive success can lead to departures from this prediction, and so we also evaluated whether any ratio of male- to female- effective population sizes could explain our observed combination of X/A and Y/A diversity.

Simulations of purifying selection

To test whether our observed level of Y-chromosome diversity could be explained by the effects of linked purifying selection, we followed the approach used in (Wilson Sayres *et al.* 2014) and conducted forward-time simulations of a nonrecombining Y chromosome using the software SFSCODE (Hernandez 2008) and compared our empirical diversity estimates on the Y with those from simulations. This approach was implemented rather than using analytical predictions of background selection primarily because the equilibrium background selection model over-predicts the reduction in diversity when there are many linked sites under selection (Kaiser and Charlesworth 2009), as is expected to be the case for large Y chromosomes that lack crossing over.

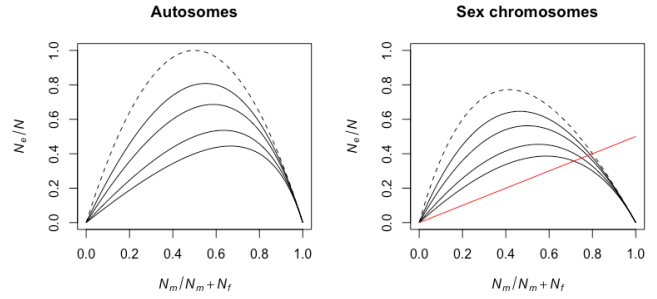


Figure 1 The relation between relative effective population size and sex ratio bias for genes on autosomes (A) and sex chromosomes (B) (Y chromosome in red). The sex ratio is shown as the proportion of males, $N_m/(N_f + N_m)$, where N_m and N_f are the effective number of breeding males and females, respectively, plotted against N_e/N , where $N = N_m + N_f$ and the N_e for sex chromosomes and autosomes is given in equations 1-3. Dotted curves show predictions in the standard neutral model, where both males and females produce Poisson-distributed offspring numbers and the chromosomal N_e 's are given by equations 4-6 (Wright 1931). Solid curves correspond to increasing levels of variance in male reproductive success (Nomura 2002) (see Methods). Assuming $\theta = 4N_e\mu$ and equal neutral mutation rates among genes, the predicted N_e 's are used to generate null predictions for X/A and Y/A ratios of diversity.

To parameterize the forward simulations, we estimated the distribution of fitness effects of deleterious mutations (DFE) using our polymorphism data on the X-chromosome with the program DFE-alpha (Keightley and Eyre-Walker ref). This program fits a gamma distribution of selection coefficients to the observed frequency distribution of nonsynonymous and synonymous polymorphism from the population genetic analyses above. We used this estimated gamma distribution in our forward simulations. To examine the expected reduction in diversity, we varied the number of sites under selection, and obtained an approximate likelihood surface for our observed data based on the proportion of simulations that matched the observed estimate of $\pi_{\text{synonymous}}$. To make the simulation output comparable to our data, we initialized the simulations with our empirically estimated autosomal θ , adjusted for the expected neutral reduction in effective population size on the Y chromosome under a sex ratio of $N_f/(N_f+N_m)=0.6$, and we sampled 6 haploid chromosomes per simulation. Simulated sequences contained 45,331 of linked neutral sequence from which we calculated diversity, matching the number of synonymous sites sampled in our empirical analyses on the Y chromosome (see Supporting Information).

Results and Discussion

Extensive loss of Y-chromosome diversity

Our analysis reveals that diversity on the *R. hastatus* Y chromosome is significantly lower than expected under neutrality, with estimates indicating Y/A=0.02, ..., which is 12.5 fold lower than the standard neutral prediction of Y/A = 0.25 ($P<0.0001$). We also observe that the Y chromosome shows a 40-fold lower than mean diversity on the X chromosome (Table 1). Note that by normalizing X and Y diversity by autosomal diversity, our

Table 1 Estimates of neutral diversity by race on *R. hastatulus* sex chromosomes and autosomes.

chromosome	Texas		South Carolina		Florida	
	θ	θ/θ_A	θ	θ/θ_A	θ	θ/θ_A
A	0.006	1	0.006	1	0.005	1
X	0.0047	0.85	0.0019	0.33	0.0018	0.37
Y	10^{-4}	0.002	10^{-4}	0.002	10^{-4}	0.002

results indicate that the X-Y difference we observed was not due to an elevation of X chromosome diversity, but rather a Y-specific reduction. Conceivably, such low diversity on the Y could arise from a low Y-linked mutation rate, or a lower mutation rate in males compared to females. However, these possibilities are unlikely because the number of synonymous mutations in X and Y lineages, estimated by both parsimony and maximum likelihood, are not significantly different (Hough *et al.* 2014). Similarly, our estimates of weighted average synonymous substitution rate between *R. hastatulus* and the outgroup *R. bucephalophorus* reveal similar levels of synonymous divergence at sex-linked (0.2016) and autosomal genes (0.219) (Hough *et al.* 2014).

In contrast to the X chromosome, however, our data indicate a strong and consistent diversity reduction on the Y chromosome: an approximately 50 fold reduction compared to the mean θ_{Aut} . We next consider several possible models - neutral and selective - that might explain this reduction.

Female biased sex ratio and high variance in male fitness

The occurrence of female-biased sex ratios in *R. hastatulus* has been predicted to lower Y diversity due to its effects on reducing male N_e and therefore the neutrally expected N_e of the Y chromosome (ref). This reduction N_e on the Y is expected to be further attenuated with high variance in male reproductive success (Figure 1), which is not unusual in annual wind-pollinated plants such as *R. hastatulus* that exhibit extensive phenotypic plasticity in plant size and flower production (Harper 1977). Moreover, given that male plants in this species produce large amounts of pollen, and that female flowers are uniovulate, we expected strong competition among males to fertilize females.

In common with most flowering plants we do not have marker-based estimates of the variance in male reproductive success in *R. hastatulus*. However, by comparing our empirical estimates of diversity to predictions from models that jointly predict the effects on diversity of sex ratio bias and male reproductive variance, we evaluated whether these effects could jointly explain the level of Y/A diversity that we observed (see Methods). Conditioning on estimates of sex ratio bias in *R. hastatulus*, ranging from $N_m/(N_m + N_f) = 0.4$ to $N_m/(N_m + N_f) = 0.35$ (Pickup and Barrett 2013), the predicted Y/A diversity ratio is approximately 0.2 (Table1, Figure 3). This is significantly lower than our estimated mean Y/A ratio of 0.02 ($P < 0.0001$), and remains significant even if we consider an upper bound estimate obtained from our likelihood based estimated of the confidence interval. This suggests that the sex ratio effect alone is insufficient to explain our data under the standard sex ratio model.

Assuming that there is extensive variance in male reproduc-

tive success (), the predicted ratios of Y/A diversity (supp) were also significantly higher than our estimates for the empirically estimated sex ratio (table) . We also find, however, that purely neutral models in which the sex ratio was highly female-biased (), with level of variance in male reproductive success on the order of (), predicted a Y/A ratio that could not be rejected (). However, because a highly females biased sex ratio is expected to increase the X/A ratio as well, these models simultaneously predicted a range of X/A ratios that were significantly different from what we observed (Figure). Thus, our results indicate that the combined effects of sex ratio bias and variance in reproductive success cannot jointly explain our observed levels of X, Y, and autosomal diversity.

Background Selection and Selective Sweeps

Conclusions

The observation of widespread degeneration and diversity loss on Y chromosomes illustrates the importance of recombination for maintaining fitness and the genetic variability needed for adaptation (Maynard Smith 1978; Kondrashov 1993; Barton Charlesworth 1998). The extensive loss of diversity on a young plant Y chromosome revealed by our study provides a clear example of the a large genomic regions that lack recombination.

Acknowledgments

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