



Fitness effects of somatic mutations accumulating during vegetative growth

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Abstract

The unique life form of plants promotes the accumulation of somatic mutations that can be passed to offspring in the next generation, because the same meristem cells responsible for vegetative growth also generate gametes for sexual reproduction. However, little is known about the consequences of somatic mutation accumulation for offspring fitness. We evaluate the fitness effects of somatic mutations in *Mimulus guttatus* by comparing progeny from self-pollinations made within the same flower (autogamy) to progeny from self-pollinations made between stems on the same plant (geitonogamy). The effects of somatic mutations are evident from this comparison, as autogamy leads to homozygosity of a proportion of somatic mutations, but progeny from geitonogamy remain heterozygous for mutations unique to each stem. In two different experiments, we find consistent fitness effects of somatic mutations from individual stems. Surprisingly, several progeny groups from autogamous crosses displayed increases in fitness compared to progeny from geitonogamy crosses, likely indicating that beneficial somatic mutations occurred in some stems. These results support the hypothesis that somatic mutations accumulate during vegetative growth, but they are filtered by different forms of selection that occur throughout development, resulting in the culling of expressed deleterious mutations and the retention of beneficial mutations.

Keywords Acquired mutations · Autogamy · Autogamy depression · Cell lineage selection · Dominance · *Erythranthe guttata* · Geitonogamy · *Mimulus guttatus* · Somatic mutations

Introduction

Mutation is the source of variation for evolution and adaptation, but organisms differ in whether mutations originating during gamete formation (meiosis) or somatic growth (mitosis) contribute to heritable variation. For the vast majority of organisms, including viruses, unicellular microbes, and some multicellular eukaryotes, sexual reproduction

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is rare or absent. In these organisms, mutations can occur during mitotic cell replication, and the primary mechanism for adaptation and diversification occurs via selection on cell lineages without recombination. By contrast, acquired mutations occurring during somatic growth of animals are not heritable. This is because in most metazoans (but possibly excepting corals; Barfield et al. 2016; Schweinsberg et al. 2014), the germline is determined early in development and relatively few cell divisions occur before the formation of gametes (the Weismann Barrier; Buss 1983). Consequently, heritable mutations typically occur in animals only during the development of gonads and gametes.

Plants differ from animals and microbes, because mutations contributing to heritable variation can arise both during gamete formation and somatic growth (Antolin and Strobeck 1985; Klekowski and Godfrey 1989). This is due to the fact that plants lack a separate germline and grow from the division of a population of undifferentiated meristem cells within the stem tip that is known as the central zone. These germ cell lineages go on to produce future stem, leaf, and reproductive tissues (e.g., flowers). Therefore, as plants grow, individual ramets of the same genet (separate stems or vegetatively propagated plants) can continue to accumulate mitotic mutations, which can make their way into the gametes and thus be passed to the next generation (Bobiwash et al. 2013; Dubrovina and Kiselev 2016; Klekowski 2003; McKnight et al. 2002; Schmid-Siebert et al. 2017; Schultz and Scofield 2009; Watson et al. 2016; Yu et al. 2020). This aspect of plant biology is well known (Ally et al. 2010; Monro and Poore 2009; Monroe et al. 2022; Reusch and Bostrom 2011), and somatic mutation accumulation has been important in agriculture, where the origin of many clonally-derived varieties of fruits, including citrus, apples, and wine grapes, have been cultivated by grafting from genetically differentiated bud tips (Aradhya et al. 2003; Jarni et al. 2015; McKey et al. 2010; Miller and Gross 2011; Pelsy et al. 2015; Vezzulli et al. 2012). Thus, since somatic mutations can be heritable, they may be an important source of genetic variation for evolution. However, there is disagreement over the extent and evolutionary importance of somatic mutation accumulation in plants (Burian et al. 2016; Hanlon et al. 2019; Kuhlemeier 2017; Plomion et al. 2018; Schmid-Siebert et al. 2017; Schultz and Scofield 2009; Watson et al. 2016).

When we consider the potential for meiotic and somatic mutations to contribute to the total mutational load of plant populations—particularly for long-lived plants—it becomes evident that not all of the mutations occurring during a plant's lifespan are passed to the next generation (Cruzan 2018, pp. 86–98). Indeed, plants have mechanisms of “developmental selection” (Buchholz 1922; Langridge 1958; Williams et al. 1999) that occur during vegetative growth and reproduction to filter the set of mutations that are inherited by progeny (Monroe et al. 2022). New somatic mutations occur as a single copy within the diploid genome, so their fitness effects for the germ cells that carry them will depend on their expression in the heterozygous state. Mathematical models have demonstrated that unexpressed mutations (i.e. neutral and recessive deleterious mutations) are likely to accumulate as germ cells divide, but expressed mutations that reduce cell growth will be eliminated. More rarely, expressed beneficial mutations will arise, such that any mutations that elevate the rate of cell division will tend to increase in frequency until they have replaced the entire germ cell population (clonal selective sweep; Lang et al. 2013; Nowell 1976). As a consequence, the composition of mutations carried by the germ cell population will be altered (Elena and Lenski 2003; Greaves and Maley 2012; Long et al. 2015; Orive 2001; Otto and Hastings 1998; Otto and Orive 1995). This process—referred to as cell lineage selection—will lead to some ramets carrying deleterious somatic mutations, while others could possess beneficial mutations.

In addition to cell lineage selection, some proportion of recessive deleterious mutations may be eliminated during the haploid life stage due to pollen tube attrition and pollen competition (Gametophytic Selection; Armbruster and Rogers 2004; Arunkumar et al. 2013; Cruzan 1989; Harder et al. 2016; Mable and Otto 1998; Mulcahy 1979). Moreover, a portion of deleterious mutations will be homozygous in zygotes, which can lead to higher rates of seed and fruit abortion (Selective Embryo Abortion; Husband and Schemske 1995; Korbecka et al. 2002), thereby increasing the average fitness of surviving offspring (Cruzan and Thomson 1997; Mena-Alí and Rocha 2005). The sum effects of these processes of developmental selection, which include cell lineage selection, gametophytic selection, and selective embryo abortion, may filter the set of mutations that enters the next generation. Therefore, understanding the role of somatic mutations for plant biology is critical to fundamental assumptions concerning the frequencies of deleterious and beneficial mutations in populations, and to the processes of adaptation and diversification.

In this study, we make crosses within individual clones of hermaphroditic plants to assess the fitness effects of inherited mutations that accumulated during somatic growth. Although it is often challenging to track the effects of somatic mutations that accumulate within a single generation, plants with separate stems contain distinct germ cell lineages that are derived from the same zygote. As a consequence, each stem can potentially contain different sets of somatic mutations that have originated during growth. By making crosses either within the same flower (autogamy) or between flowers on separate stems of the same plant (inter-ramet geitonogamy—hereafter referred to as geitonogamy), we can produce progeny segregating for somatic mutations unique to each stem (Fig. 1). These crosses are both self-fertilizations, but the offspring of each cross type will differ in the complement of somatic mutations that they inherit. For a diploid plant, we can assume that somatic mutations ($a \rightarrow a'$) will be in the heterozygous state when they first appear. For progeny generated via autogamy, a somatic mutation will segregate as 25% homozygous ($a'a'$), 50% heterozygous (aa'), and 25% the original (wildtype) homozygote (aa). By contrast, because progeny from geitonogamous crosses will segregate for somatic mutations that are unique

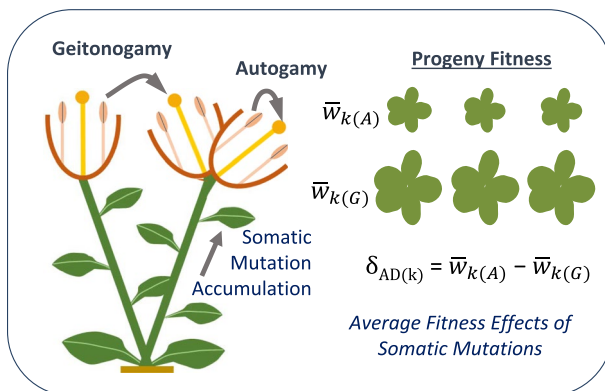


Fig. 1 Experimental design to test for the average fitness effects of somatic mutations accumulating in stems of *Mimulus guttatus* during vegetative growth. A proportion of somatic mutations accumulating during stem growth (dark blue arrows) is made homozygous after within-flower (autogamous) self-pollinations, while all somatic mutations will be heterozygous after between-stem (geitonogamous) self-pollinations. Comparison of the mean fitness of autogamous seedlings ($\bar{w}_{k(A)}$) to geitonogamous seedlings from the same stem ($\bar{w}_{k(G)}$) provides an estimate of the average fitness effects of somatic mutations unique to each stem ($\delta_{AD(k)} = \bar{w}_{k(A)} - \bar{w}_{k(G)}$)

to each stem, 50% of offspring will be carrying mutations in the heterozygous state, and none of the progeny will be homozygous for mutations that arose in a single stem. Thus, the average fitness effects of somatic mutations can be evaluated by comparing the difference in fitness of progeny generated by autogamous and geitonogamous crosses (Bobiwash et al. 2013; Schultz and Scofield 2009).

The effects of deleterious somatic mutations often are apparent as higher rates of embryo abortion after autogamous compared to geitonogamous pollinations, which is referred to as autogamy depression (Schultz and Scofield 2009; Fig. 1). While autogamy depression for seed and fruit abortion has been observed in several species (reviewed in Bobiwash et al. 2013), no previous study has evaluated the fitness effects of somatic mutations inherited by progeny. In this study, we develop and validate methods that use autogamous and geitonogamous self-pollinations to estimate the fitness effects of somatic mutations segregating in offspring, and then use these methods in two separate experiments to estimate the fitness effects of somatic mutations accumulating during vegetative growth in perennial *Mimulus guttatus* DC (*Erythranthe guttata* G.L. Nesom; Phrymaceae).

Methods

Estimating the fitness effects of somatic mutations

As described above, progeny from autogamous self-pollinations will be homozygous for a proportion of somatic mutations, while progeny from geitonogamous crosses will be heterozygous. Consequently, we can estimate the average fitness effects of somatic mutations that are unique to each stem by comparing the average fitness of progeny from autogamous ($\bar{w}_{k(A)}$) and geitonogamous ($\bar{w}_{k(G)}$) crosses to a given stem k as: $\delta_{AD(k)} = \bar{w}_{k(A)} - \bar{w}_{k(G)}$. The parameter $\delta_{AD(k)}$ has been described before in a slightly different format (referred to as Autogamy Depression; Schultz and Scofield 2009; Bobiwash et al. 2013) and summarizes the average magnitude and overall direction of the fitness effects of all expressed somatic mutations that have been transmitted to the next generation. We note that $\delta_{AD(k)}$ is a quantitative genetic estimate of the sum effect of all somatic mutations that accumulated in a stem (Bobiwash et al. 2013; Schultz and Scofield 2009) and is analogous to estimates of inbreeding depression (Charlesworth and Willis 2009). While standing genetic variation can also lead to fitness effects in offspring, there is no a priori reason to expect these effects to differ in progeny from fruits on different stems of the same plant. Hence, differences in the fitness of progeny from autogamous and geitonogamous pollinations are expected to reflect only the effects of recessive and partially dominant somatic mutations that have accumulated during stem elongation.

Validation of fitness estimates

Given that $\delta_{AD(k)}$ is based on fitness differences between offspring from autogamous and geitonogamous crosses, it is possible that variation in the dominance of somatic mutations may reduce the reliability of the parameter. In particular, combining somatic mutations from two separate stems in progeny from geitonogamous crosses may reduce fitness and generate estimates of $\delta_{AD(k)}$ that falsely indicate the presence of beneficial somatic mutations in autogamous progeny, when mutations are actually deleterious (i.e. causing a change in the sign of the estimate). Therefore, to evaluate the reliability of $\delta_{AD(k)}$, we

simulated three different scenarios for variation in the strength of selection and dominance of somatic mutations across multiple stems using randomization functions in Excel (Microsoft Office Professional 2019). First, we assumed that there was only one somatic mutation in each of the two stems, and we calculated expected fitness in the geitonogamous (w_G) and autogamous (w_A) progeny as;

$$w_G = 1 + 0.5(h_1s_1) + 0.5(h_2s_2), \text{ and}$$

$$w_A = 0.25(1) + 0.5(1 + h_1s_1) + 0.25(1 + s_1).$$

where s_1 and s_2 represent the selective effects of mutations, and h_1 and h_2 represent the dominance coefficients of single somatic mutations in stem 1 and stem 2, respectively. Dominance values were chosen assuming that deleterious somatic mutations with high levels of dominance would be eliminated by cell lineage selection during vegetative growth, and beneficial mutations with high dominance would not display strong fitness differences between autogamous and geitonogamous progeny. For simulations, we randomly chose values of s ranging from -0.2 to 0.2 for 200 pairs of stems. Values of h for $s > 0$ were allowed to range from 0 to 0.7 for each stem. Because deleterious alleles tend to be recessive (Dudash and Carr 1998; Peters et al. 2003), values of h were constrained to range between 0 and 0.1 for $s < 0$. To match our experimental design in *Mimulus guttatus* (see below), we calculated the fitness of autogamous progeny (w_A) for one of the stems in each pair. We then calculated w_G using information on s and h for both stems in each pair. Then we calculated the difference in fitness between each estimate as a measure of autogamy depression ($\delta_{AD(k)} = w_A - w_G$), compared the estimate of $\delta_{AD(l)}$ to values of s_l , and evaluated the frequency of estimates that were opposite in sign to the actual selective value. This simulation was repeated 20 times (4,000 pairs of stems total) and average values were calculated.

One limitation of the formulation above is that estimates of w_G will be out of range when large numbers of mutations are involved (i.e., $w_G = 1 + 0.5 \sum h_i s_i$ for a large number of loci segregating for deleterious alleles). To remedy this, we used two sets of fitness calculations for simulations with multiple loci. For the first, we assumed that interactions among somatic mutations would be additive. We estimated the average effect of n loci segregating for deleterious alleles (i.e. $\sum h_i s_i / n$). If we apply this approach for two loci (one mutation per stem) we obtain,

$$w_G = 1 + 0.375(h_1s_1 + h_2s_2), \text{ and}$$

$$w_A = 0.25 + 0.25(1 + h_1s_1) + 0.25(1 + h_2s_2) + 0.25(1 + (h_1s_1 + h_2s_2)/2).$$

For a second set of simulations that examined the effects of alleles at multiple loci, we assumed the interactions were multiplicative. In this case, the fitness estimates of geitonogamous and autogamous progeny become,

$$w_G = 1 + 0.25(h_1s_1 + h_2s_2) + 0.25(h_1s_1h_2s_2), \text{ and}$$

$$w_A = 0.25 + 0.25(1 + h_1s_1) + 0.25(1 + h_2s_2) + 0.25(1 + (h_1s_1h_2s_2)).$$

For both the additive and multiplicative interaction scenarios, we conducted simulations of 200 stems 20 times as described above (a spreadsheet to conduct simulations is available as Appendix 1).

Fitness estimates based on variation among autogamous progeny

To provide additional confirmation of the $\delta_{AD(k)}$ estimates, we developed an independent approach for estimating the fitness effects of somatic mutations that is based entirely on the variance in mean fitness among autogamous progeny from the same fruit. If somatic mutations affect offspring fitness, variation in fitness should be greater for progeny groups from autogamy than from geitonogamy, as long as mutations do not have complete expression in heterozygotes (i.e. $h < 1.0$; Appendix 2). This is because somatic mutations will segregate as homozygotes and heterozygotes in autogamous progeny but will remain heterozygous in the progeny of geitonogamous crosses. Therefore, we can estimate the fitness effects of somatic mutations based on the standard deviation in fitness from autogamous crosses (SD), according to the equation: $w_{SD} = cSD$, where c is the slope of the linear relationship between w_{SD} and SD . For dominance levels of $h = 0.0$ and 1.0 , $c = 2.31$, and the slope reaches a maximum of 2.83 when $h = 0.5$ (Appendix 2; Fig. S1). Thus, using only the variation in fitness among progeny from autogamous crosses, we obtain a good approximation of the magnitude of the fitness effects (w) for somatic mutations accumulating in each stem. This approach is independent of $\delta_{AD(k)}$, which therefore provides an important means of confirming our estimates of the average fitness effects of somatic mutations present in individual stems.

Tests of predictions

From the considerations above, we can identify three predictions for the fitness effects of somatic mutations that occur during stem growth and are passed to offspring. First, on a single stem, the average fitness of progeny deriving from an autogamous cross should be different from the fitness of progeny arising from a geitonogamous cross made using pollen from a different stem. We test for differences in the fitness of progeny from autogamous vs. geitonogamous crosses, and use the mean fitness of each progeny group paired by stem to estimate the sign and magnitude of the average fitness effects of somatic mutations using the formula $\delta_{AD(k)} = \bar{w}_{k(A)} - \bar{w}_{k(G)}$ as described above. The second prediction is that somatic mutations unique to individual stems will result in greater variation in mean fitness among autogamous progeny from the same fruit compared to geitonogamous progeny. We test this prediction by comparing the variance among autogamous and geitonogamous progeny groups. Since we expect each stem to possess unique somatic mutations that will be made homozygous in autogamous progeny, there should be greater variation in mean fitness among autogamous progeny from different stems compared to geitonogamous progeny, which will be heterozygous for somatic mutations or homozygous for the non-mutant allele. Consequently, the third prediction is that somatic mutations will result in significant cross type by stem interactions for mean fitness. Below, we test these three hypotheses using two different experiments with *Mimulus guttatus*.

Study system

Populations of *M. guttatus* display a wide range of life histories—from annuals to herbaceous perennials that outcross to varying degrees (Wu et al. 2008). We use perennial *M. guttatus* plants that produce substantial vegetative growth prior to initiation of flowering. These plants are easily propagated from rosettes, and selfing produces substantial numbers of seeds to allow for statistical comparisons. To increase the chances of detecting somatic mutations that impacted fitness, we exposed plants to novel environments. Furthermore, we exposed parental plants and seedlings to the same controlled environments to evaluate the fitness effects of mutations (Baer et al. 2007; Halligan and Keightley 2009; Shaw et al. 2002).

First experiment

We grew plants of *M. guttatus* in the Research Greenhouse facility at Portland State University from seed collected in July 2013 from three different populations in northern Oregon (Jackson Bottom Wetlands–JB: 45.501794 N, – 122.98776 W; and two from Saddle Mountain–SMB: 45.9861 N, – 123.6859 W, and SMC: 45.9634 N, – 123.6837 W). We assumed that greenhouse conditions were different enough from field environments to provide a novel environment. In August 2013, seeds were cold stratified on moist paper towels at 2 °C for 30 days prior to being sown in soil. Seedlings were transplanted to pots (approximately 10 × 10 × 12 cm) and grown for seven months before the application of pollination treatments. Temperature was maintained between 21 and 26 °C during the day, and 15–21 °C at night. Supplemental HID lights ran for 12 h a day when the seedlings first emerged, and 14 h a day during adult growth.

After plants became established and began producing multiple stems, we conducted autogamous and geitonogamous self-pollinations using flowers on stems 15 to 20 cm in length from two plants from each of four maternal families representing each of the three populations (2 plants × 4 families × 3 populations). Flowers from pairs of stems on individual plants (ramets of the same genet) were reciprocally crossed (geitonogamy), or individual flowers from these same stems were self-pollinated (autogamy). A total of 139 pollinations were conducted across two treatments: limited (pollen was applied to stigmas with one touch from a plastic pipette tip) or excess (where the stigma surface was coated with pollen) in an attempt to manipulate the intensity of gametophytic selection (Cruzan and Barrett 1996). Pollinations were conducted on 12 different days (pollination date) over several weeks in July 2014. Mature fruits were collected and placed individually into paper envelopes, and their contents were examined under a Leica MZ-16 stereoscope. The first 100 ovules from each fruit were categorized as filled seeds (brown, almond-shaped), unfertilized ovules (small, flattened and light-colored), or aborted (larger than unfertilized, dark-colored, shriveled). Ovules that were flattened and appeared to lack endosperm were assumed to be unfertilized or aborted and were not used in germination tests. Seed set and ovule abortion were analyzed using ANOVA models with the GLM procedure of SAS (SAS 2008), with population, maternal plant nested within population, and pollination date as random effects, and cross type (autogamous or geitonogamous) and pollination treatment as fixed effects. Seed set and ovule abortion data were approximately normal so were not transformed prior to analysis.

We assessed the fitness of progeny arising by autogamy and geitonogamy in the same greenhouse environment that was used to grow the parental plants. Seedlings from a subset of ten maternal plants (at least two plants from each of the three populations) that had fruits from both cross types and at least 20 filled seeds were sown in soil and transplanted to 36-cell trays (blocks) in September in a randomized incomplete block design. After 3 months of growth, the progeny were scored for survival, and above ground biomass was measured after drying at 60 °C for at least 24 h. Since all seedlings germinated within a few days of each other, the final biomass is an estimate of growth rate. The fitness of each progeny was estimated as its final biomass, which was log transformed and weighted by the survival frequency of progeny from the same cross. Growth rate is considered to be an appropriate estimate of fitness for perennials (Younginger et al. 2017). Furthermore, we evaluated fitness of seedlings under novel selection regimes in the greenhouse rather than under field conditions, which allowed us to minimize environmental differences between the growth environments of parental plants and seedlings. These estimates were rescaled relative to the maximum value from all crosses, so that \bar{w} ranged from 0 to 1 across progeny from all crosses.

Second experiment

To control for variation among genets and to further assess the fitness effects of mutations that accumulated during vegetative growth, a single plant (genet BV; obtained from Willamette Gardens native plant nursery, Corvallis, OR) was vegetatively propagated to generate 12 plants (ramets) that were exposed to high salinity and control conditions. This genet was originally propagated from wild-collected seed and retains a high level of genetic diversity (J.A. Schwoch, unpub. data). Plants were grown in pea gravel (4–8 mm) in pots placed in four 53 L tubs using a flood and drain hydroponics system (flooding at 15 min intervals). Two tubs had no added salt and were used as controls, and two tubs had high salinity. The initial salt concentration in the high salinity treatment was 5 mM but increased weekly to 25 mM after plants became established. Salt concentrations were monitored using a conductivity meter to ensure stable concentrations. To provide nutrients, 30 ml of hydroponics fertilizer (FloraGrow, Planet Natural, Bozeman, MT) was added per tub. During the course of the experiment, some plants grew substantially. We imposed selection to favor the fastest growing ramets over the next three months by repeatedly removing single rosettes and transplanting them back into the hydroponics system up to four times.

To promote stress recovery, plants were transplanted to soil for six months, which included a two-month vernalization period in a growth chamber (4 °C and 8 h light; Conviron E8, Controlled Environments Ltd., Winnipeg, Manitoba, Canada). After vernalization, plants were returned to the greenhouse to induce flowering. Autogamous and geitonogamous pollinations were made to pairs of flowers at single nodes or consecutive nodes (seven nodes and 14 pollinations total) on the largest ramets in each of the control and high salt treatments. To account for somatic mutation turnover that may occur due to the effects of cell lineage selection during stem growth, we compared progeny from autogamous and geitonogamous pollinations for a subset of four stems that produced fruits from pairs of flowers from the same node. Without a priori knowledge of the expression of somatic mutations in heterozygotes, it is difficult to determine the best pollen donor for geitonogamous crosses. Consequently, we opted to generate progeny from geitonogamy by pollinating flowers with pollen from a potentially more genetically divergent ramet from the other treatment (i.e. salt pollinated with control pollen, and control pollinated with salt pollen;

two stems from each treatment). The fruits were collected, and the total number of aborted and mature seeds and unfertilized ovules were counted under a dissecting microscope. Seeds were planted in soil in trays with three seeds per cell. Seeds were cold stratified in moist soil for three weeks before they germinated in the greenhouse.

To determine whether seedlings from autogamous crosses from ramets exposed to salt stress showed improved performance under the same conditions, all progeny were exposed to high salinity. After germination and establishment in soil, seedlings from autogamous and geitonogamous crosses from control and salt stress ramets were transplanted into pots filled with pea gravel and subjected to high salt in the hydroponics system, as described above. A total of 239 seedlings from 11 fruits (five autogamous and six geitonogamous) were randomly and evenly distributed among 12 hydroponic tubs to ensure equal representation across blocks (tubs). Plant size was measured as the product of the length and width of vegetative spread after two months of growth and was used as a proxy for overall plant performance. Since plants germinated within a few days of each other, plant size represents a good estimate of growth rate. Salt concentration increased from 10 to 37.5 mM over the course of the experiment to induce mortality (~57% across all progeny groups). Fitness was estimated as plant size (log transformed to improve normality) weighted by the survival frequency for progeny from the same cross.

Data analyses

Data were analyzed to compare the means and variance in autogamous and geitonogamous progeny fitness to test the three predictions described above. We first test the prediction that somatic mutations would generate greater variation among autogamous compared to geitonogamous progeny grouped by fruit separately in each experiment. We test this prediction using two-way ANOVAs and the stem \times cross type (i.e. autogamy vs. geitonogamy) interaction for progeny fitness (Prediction 3 above). We then use data pooled from the two experiments (see below) to test the prediction that the fitness of autogamous and geitonogamous progeny paired by stem will differ (Prediction 1), and that, among progeny within fruits, there will be greater variation in fitness among autogamous compared to geitonogamous progeny (Prediction 2).

Results

Validation of fitness estimates

Our simulations revealed that values of $\delta_{AD(k)}$ were accurate across a range of values and assumptions for the effects of somatic mutations. For the simulations assuming one mutation per stem, estimates of $\delta_{AD(k)}$ and s_I were strongly correlated, with R^2 values ranging from 0.76 to 0.82 (see Fig. S4 for example results). For simulations assuming additive interactions among multiple somatic mutations, the relationship between $\delta_{AD(k)}$ and s_I was stronger (R^2 ranging from 0.86 to 0.90), and was stronger still when we assumed multiplicative interactions (R^2 ranging from 0.93 to 0.95).

Overall, these results indicate that our estimate of $\delta_{AD(k)}$ is a valid estimate of s ; however, it does appear that we are underestimating s (i.e., the slope of the line is less than 1; Fig. S4), making $\delta_{AD(k)}$ a conservative estimate of the fitness effects of somatic mutations. It is also notable that our simulation assumes a wider range of h values than are generally

observed for deleterious mutations. Estimates indicate that the product of hs for deleterious alleles is generally around 0.02, and that the relationship between s and h is generally negative (hyperbolic; Lynch et al. 1999). Minor effects of deleterious somatic mutations are expected, because somatic mutations having strong effects would be eliminated by cell lineage selection. Therefore, these simulations confirm that $\delta_{AD(k)}$ provides a valid estimate of the mean fitness of somatic mutations occurring within a stem.

Experimental results

In the first experiment, there was no effect of cross type on seed set (Table S5 in Appendix 3), but there was higher abortion of developing seeds after autogamous compared to geitonogamous crosses (Table S6). For ramets from both experiments, there was significant variation in mean fitness among seedlings from fruits generated from autogamous crosses, but not among seedling groups derived from geitonogamy. Consequently, tests of the cross-type by stem interaction were significant for both experiments (Tables S1 and S2 in Appendix 3), which is consistent with the third prediction described above. Since the results from these two experiments were qualitatively similar, we combined the data for all of the following analyses (10 stems from Experiment 1 and four from Experiment 2, for a total of 14 comparisons).

To test the first prediction described above, that inheritance of somatic mutations will have larger effects on autogamous progeny fitness ($\bar{w}_{k(A)}$), we compared them with the fitness of geitonogamous progeny from the same stem ($\bar{w}_{k(G)}$; Tables S3 in Appendix 3). As validated by our simulations, the difference in mean fitness for the two cross types provides an estimate of the average fitness effects of somatic mutations unique to each stem. The total fitness effects of all mutations ($\delta_{AD(k)}$) occurring in each stem (first experiment) or stem/node combination (second experiment) were calculated as the difference in fitness between progeny from autogamous and geitonogamous crosses, as described above ($\delta_{AD(k)} = \bar{w}_{k(A)} - \bar{w}_{k(G)}$). We observed extensive variation in $\delta_{AD(k)}$ among stems, with nine stems that were significantly different from zero (Fig. 2). Four of the stems had average fitness

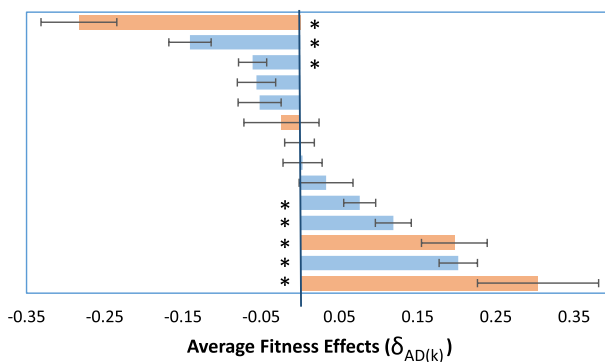


Fig. 2 Estimates of $\delta_{AD(k)}$ for fourteen different stems (ramets) of *Mimulus guttatus* from two separate experiments (Experiment 1–blue bars; Experiment 2–orange bars) based on mean progeny fitness after autogamous $\bar{w}_{k(A)}$ and geitonogamous $\bar{w}_{k(G)}$ self-pollinations ($\delta_{AD(k)} = \bar{w}_{k(A)} - \bar{w}_{k(G)}$). Horizontal lines represent standard errors. Asterisks indicate values of $\delta_{AD(k)}$ that are significantly different from zero based on the t value, calculated as $t = \delta_{AD(k)} / \sqrt{SE}$ with $n-1$ df, where n is the mean of sample sizes for progeny from autogamy and geitonogamy. The relationship between $\bar{w}_{k(A)}$ and $\bar{w}_{k(G)}$ across stems is shown in Fig. S3. Means and sample sizes for progeny groups are available in Table S2 and Fig. S4 in Appendix 3

effects of somatic mutations that were positive, suggesting a net beneficial effect of somatic mutations transmitted to offspring. In addition, the average fitness of progeny from autogamous and geitonogamous crosses on the same stems or nodes was positively correlated (Fig. S2), which indicates that the effects of somatic mutations from the pollen-donor stem were minimal and do not account for the instances of beneficial mutation estimates found in some stems. Note also that the $\delta_{AD(k)}$ estimates that deviated the most from zero generally had the highest $\bar{w}_{k(A)}$ values (Fig. S3). However, this was not always the case, as one stem (stem 9) with a value of $\delta_{AD(k)}$ close to zero produced progeny with relatively high fitness after both autogamous and geitonogamous pollinations.

To make an independent estimate of the average fitness effects for mutations unique to each stem or node, we used the standard deviation in progeny fitness from autogamous crosses based on the relationship $w_{SD} = cSD$, where $c = 2.83$, which assumes dominance close to $h = 0.5$ (qualitatively similar results are obtained if we choose other values of c between 2.83 and 2.31, and when $h = 0$ or 1; Appendix 2). Because the sign of w_{SD} could not be inferred directly from this approach, we used the sign estimated from the $\delta_{AD(k)}$ method (Fig. 3; Appendix 2). For stems with values of $\delta_{AD(k)}$ greater than zero, there was a strong positive relationship between $\delta_{AD(k)}$ and estimates of w_{SD} made from the within-family variation among progeny from autogamy (Fig. 3). In contrast, the relationship for negative values of $\delta_{AD(k)}$ appeared to be driven largely by a single observation. Note that this observation was not supported by a similarly high value of w_{SD} , and the remaining negative fitness effects were more modest based on both estimates. This one highly negative estimate of $\delta_{AD(k)}$ may be due to a large influence of mutations from the second stem on the fitness of geitonogamous progeny (note that this stem had the highest estimate of $\bar{w}_{k(G)}$; Fig. S2). It is also notable that the variation in fitness for progeny from autogamy did not decline to zero for values of $\delta_{AD(k)}$ close to zero, which could be due to the presence of both beneficial and detrimental mutations, and possibly genetic background effects (i.e. epistasis), but it may also reflect uncontrolled environmental variation. Overall, the results from both approaches reveal consistent estimates of the average fitness consequences associated with the accumulation of somatic mutations in individual stems.

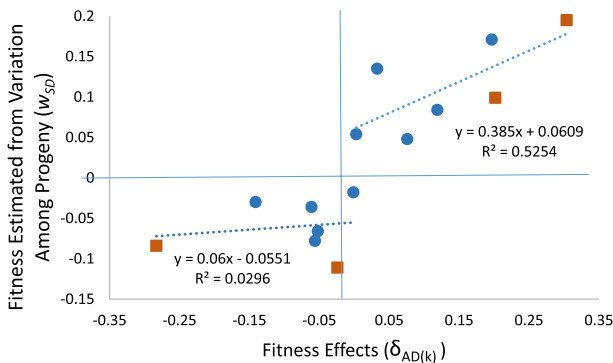


Fig. 3 Relationships between estimates of fitness effects of somatic mutations in *Mimulus guttatus*, based either on the difference in fitness of progeny from autogamy and geitonogamy ($\delta_{AD(k)}$), or the standard deviation in fitness within progeny groups from autogamy for each stem (w_{SD}). Estimates of w_{SD} corresponding to negative values of $\delta_{AD(k)}$ were transformed to negative values. Estimates from Experiment 1 are indicated by blue circles and from Experiment 2 are orange squares. Dashed lines indicate the separate relationships for positive and negative values of fitness estimates

Discussion

In this study, we have provided evidence that somatic mutations accumulating during vegetative growth can affect the fitness of progeny in the next generation. Consistent with previous work (Bobiwash et al. 2013), we observed significant autogamy depression in the form of greater ovule abortion in autogamous relative to geitonogamous crosses. However, we also detected more variation in survival and growth rate for progeny from autogamous compared to geitonogamous self-pollinations, which provides evidence that somatic mutations that accumulated during vegetative growth can have demonstrable effects on the fitness of plants in the next generation. Both the differences in the mean fitness between progeny from autogamy and geitonogamy ($\delta_{AD(k)}$) and variation in fitness of autogamous progeny (w_{SD}) provided consistent estimates of the average fitness effects of somatic mutations segregating in progeny. We found evidence for the effects of beneficial mutations in progeny from some stems, with estimates of $\delta_{AD(k)}$ and w_{SD} exceeding 0.1 in four cases, while estimates for negative fitness effects were more modest (mostly > -0.15). These results imply that, in these plants, many deleterious mutations can be culled by the various types of developmental selection prior to seed dispersal.

Somatic mutations accumulating during vegetative growth had an overall positive effect for four of the stems tested. This result appears to contrast with widely held views that the appearance of beneficial somatic mutations should be exceedingly rare (Charlesworth and Willis 2009; Crow 1993). However, this finding is consistent with expected outcomes of clonal evolution occurring during vegetative growth. We expect that most somatic mutations in germ cells in the central zone of the apical meristem will be neutral and occur at low frequencies, but some could contribute to higher rates of division for some cell lineages over others (Otto and Hastings 1998; Otto and Orive 1995). This hypothesis is supported by the observation that the majority of somatic mutations tend to occur at low frequencies in stem tissue, and a minority occur at high frequencies (Yu et al. 2020; Schwoch et al. unpublished data), suggesting that clonal selective sweeps due to the appearance of beneficial mutations have occurred during stem elongation. While the stem cell population is small (e.g., ~ 35 in *Arabidopsis*; Reddy and Meyerowitz 2005), selection on somatic mutations was apparently strong enough to overcome genetic drift, resulting in higher fitness of autogamous progeny from some stems. Thus, it appears that cell lineage selection has the potential to retain beneficial somatic mutations, as individual cell lineages divide at faster rates resulting in clonal evolution during vegetative growth.

Although there may be few opportunities for beneficial changes to alter basic cellular metabolism, it is becoming apparent from experimental evolution studies with microbes that even basic aspects of cellular metabolism can be sensitive to environmental conditions, which can increase the chances that mutations in clonal populations are beneficial (Böndel et al. 2019; e.g., Lang et al. 2013; Lee and Marx 2013; Maharjan et al. 2015). In the experiments described above, we exposed plants to novel environments either in the greenhouse (experiment 1) or in high salinity in hydroponics (experiment 2). Moreover, genomic evidence suggests that clonal selective sweeps, which may be indicative of the spread of beneficial somatic mutations in the meristem tissue, are more common for stems that have recently been exposed to high salinity (J. A. Schwoch, unpublished data). In this regard, cell lineage selection in a plant meristem represents a potentially powerful forum for the removal of deleterious somatic mutations while favoring the retention of beneficial ones. In addition, gametophytic selection and selective embryo abortion can act as prominent additional filters (Armbruster and Rogers 2004; Arunkumar et al. 2013; Cruzan 1989; Harder

et al. 2016; Mable and Otto 1998; Mulcahy 1979), but they are most likely to have effects by culling deleterious mutations. Regardless, the combined effects of these different forms of developmental selection appear to have had a considerable effect on filtering of somatic mutations under controlled conditions in the greenhouse, such that the distribution of fitness effects among stems has shifted to include more beneficial mutations than expected. Future studies should test whether similar findings are found under field conditions, which could indicate a prominent role for somatic mutation in local adaptation.

An alternative explanation for the observed fitness differences is that exposure to environmental stress has induced heritable epigenetic modifications (Quadrana and Colot 2016). However, phenotypic responses due to epigenetic modifications are expected to be consistent and predictable, as they are hypothesized to represent an adaptive response to historic exposure to similar stressors (Baulcombe and Dean 2014; Itabashi et al. 2018). The results of the experiments described here are unlikely to support a role for epigenetics, because fitness responses in the next generation were inconsistent in direction and magnitude, and they were not predictable based on environmental exposure of the parent stem. Moreover, there is no a priori reason to expect that epigenetic modifications would differ between fruits from autogamous and geitonogamous crosses developing on the same stem. We observed that the mean fitness of progeny from autogamy displayed both increases and decreases compared to the geitonogamy controls, which is consistent with the hypothesis that individual ramets are accumulating unique complements of somatic mutations. Furthermore, these conclusions are supported by the observation of numerous somatic variants in the transcriptomes of meristem tissue from multiple ramets derived from a single genet (Yu et al. 2020; Schwoch et al. unpublished data). Thus, the results of the current study indicate that somatic mutations accumulating during stem growth are most likely to be responsible for the observed fitness effects.

The potential for the acquisition of mutations during vegetative growth is a well-known aspect of plant biology (Bobiwash et al. 2013; Klekowski 2003; Schultz and Scofield 2009), but no previous study has demonstrated the effects of somatic mutations on the fitness of progeny in the next generation. Moreover, most studies have focused on the detrimental effects of somatic mutations; chloroplast mutants have been observed in a number of species (Klekowski 2003), declines in pollen fertility were found in older clones of quaking aspen (Ally et al. 2010), and higher rates of seed and fruit abortion after autogamous pollinations were found in several studies (reviewed in Bobiwash et al. 2013). In contrast, agriculturalists have taken advantage of beneficial somatic mutations to improve economically important plants (Aradhya et al. 2003; Jarni et al. 2015; McKey et al. 2010; Miller and Gross 2011; Pelsy et al. 2015; Vezzulli et al. 2012), and a handful of studies report phenotypic responses to selection in asexual lineages. For example, Breese et al. (1965) succeeded in selecting for increased tillering ability (production of new grass stems) within genets of perennial ryegrass (*Lolium perenne*). Similarly, artificial selection on clonal lineages effectively improved branching in the red seaweed, *Asparagopsis armata* (Monro and Poore 2009). The current study on *M. guttatus* contributes to this literature by highlighting the potential for plants to exhibit significant levels of clonal variation within a single generation.

The transmission of beneficial mutations in autogamous crosses may explain some heretofore difficult to understand results from mutation accumulation studies. Our results suggest that beneficial somatic mutations are likely to be partially dominant (i.e. not completely recessive), because they would have to be expressed in the heterozygous state to be favored by cell lineage selection. While somatic mutations may become homozygous through mitotic recombination, this appears to be rare in non-cancerous somatic growth

(LaRocque et al. 2011). Mutations accumulating during vegetative growth could contribute to standing genetic variation, but autogamy may be more effective for the accumulation of beneficial somatic mutations in populations than geitonogamy or outcrossing, because beneficial mutations could be made homozygous in progeny in a single generation. Depending on the crossing design, outcrossing would take at least two generations for beneficial somatic mutations to be made homozygous, and under geitonogamy they would remain heterozygous. It is striking that high rates of beneficial mutation accumulation have been observed in at least some mutation accumulation studies in the autogamous plant *Arabidopsis thaliana* (Rutter et al. 2012, 2018, 2010; Shaw et al. 2002), but not in outcrossing and partially-selfing species of *Amsinckia* (Schoen 2005). With a few exceptions (e.g., Baer et al. 2005; Denver et al. 2010), nearly all mutation accumulation studies on animals consistently show a prevalence of deleterious mutations (Baer et al. 2007; Halligan and Keightley 2009; but see Bao et al. 2022). Our results suggest that the adaptive potential of autogamous plants may be greater than previously thought, which may help explain the wider geographic ranges of selfing compared to closely-related outcrossing species (Grant and Kalisz 2020; Grossenbacher et al. 2015). Although developmental selection has the potential to contribute to adaptation in all plants, its effects may be enhanced in autogamous lineages, because beneficial mutations arising during vegetative growth have a greater chance of becoming homozygous in offspring and being retained across generations.

As stems grow, mutations can be generated during every mitotic cell division, so the potential for somatic mutation accumulation in plants appears substantial. Thus, understanding how long-lived plants, such as trees, avoid mutational meltdown from the accumulation of deleterious somatic mutations remains a longstanding question. Paradoxically, however, the rate of mutation accumulation observed across generations in plant and animal genomes is similar (Gaut et al. 2011). One hypothesis for this pattern is that, similar to animal germlines, a subset of cell lines in the apical meristem undergoes fewer mitotic divisions, which would protect lineages from the negative effects of mutation accumulation during development of the soma (Plant Germline Hypothesis; Burian et al. 2016; Cruzan 2018; page 90). An alternative hypothesis posits that somatic mutations are generated in apical meristems during plant growth, but these mutations are filtered by developmental selection prior to the establishment of offspring (Somatic Mutation Accumulation Hypothesis; Cruzan 2018, p. 86). Plants have retained the capacity to undergo clonal evolution from their algal ancestors, and thus developmental selection during growth and reproduction has the potential to skew the distribution of fitness effects of transmitted mutations to include a larger proportion of beneficial variants than would be expected through random processes. In addition, this provides a reasonable explanation for why longer-lived plants appear to have slower rates of mutation accumulation across generations (Gaut et al. 2011; Yue et al. 2010). This could be due to the fact that longer generation time leads to more time between recombination events, which can lead to more background selection in non-recombining cell lineages during vegetative growth (Cruzan 2018, pp. 94–95). Recent studies in oaks, spruce, and eel grass (Hanlon et al. 2019; Plomion et al. 2018; Yu et al. 2020), as well as our unpublished data from *M. guttatus*, confirm that multiple somatic variants are likely, even in plants of very different size. Future work that combines information from experiments evaluating the genomic consequences of somatic variation with anatomical estimates of stem cell population dynamics will allow for the development of new models that provide insights into the extent and limitations of somatic evolution in plants.

In conclusion, despite the potential for somatic mutation accumulation to generate novel genetic variation in plant populations, its role in evolution remains almost entirely unexplored. Our estimates of the fitness effects of somatic mutations were consistent across two

different methods and indicate that some stems accumulated primarily deleterious mutations while others produced autogamous progeny with high fitness, which likely indicates the presence of beneficial mutations. Even though high levels of mutation accumulation are often believed to be detrimental, the basic biology of plants suggests that the role of somatic mutations in plant evolution should be considered carefully in the future. Future lines of investigation will improve our understanding of these fundamental aspects of plant development and evolution that may have contributed to the remarkable diversification of plants, and may help to account for some of the variation in mutation rates detected among lineages.

Data archiving

Data and all scripts for data analysis will be submitted to DataDryad upon publication.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s10682-022-10188-3>.

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Author contributions JAS conducted the greenhouse experiments, processed samples, and assisted with data analysis and wrote a portion of the manuscript. MAS assisted with data analysis and data presentation, and wrote a portion of the manuscript. MBC conducted data analyses and simulations and wrote the majority of the manuscript.

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Data availability Data and all scripts for data analysis will be submitted to DataDryad upon publication.

Code availability Included as Appendix 1.

Declarations

Conflict of interest The authors have no conflicts of interest.

Consent for publication All authors have read the manuscript and agree with its contents.

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