A resurrection experiment finds evidence of both reduced genetic diversity and potential adaptive evolution in the agricultural weed \textit{Ipomoea purpurea}

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\textbf{Abstract}

Despite the negative economic and ecological impact of weeds, relatively little is known about the evolutionary mechanisms that influence their persistence in agricultural fields. Here, we use a resurrection approach to examine the potential for genotypic and phenotypic evolution in \textit{Ipomoea purpurea}, an agricultural weed that is resistant to glyphosate, the most widely used herbicide in current-day agriculture. We found striking reductions in allelic diversity between cohorts sampled nine years apart (2003 vs. 2012), suggesting that populations of this species sampled from agricultural fields have experienced genetic bottleneck events that have led to lower neutral genetic diversity. Heterozygosity excess tests indicate that these bottlenecks may have occurred prior to 2003. A greenhouse assay of individuals sampled from the field as seed found that populations of this species, on average, exhibited modest increases in herbicide resistance over time. However, populations differed significantly between sampling years for resistance: some populations maintained high resistance between the sampling years whereas others exhibited increased or decreased resistance. Our results show that populations of this noxious weed, capable of adapting to strong selection imparted by herbicide application, may lose genetic variation as a result of this or other environmental factors. We probably uncovered only modest increases in resistance on average between sampling cohorts due to a strong and previously identified fitness cost of resistance in this species, along with the potential that nonresistant migrants germinate from the seed bank.

\textbf{Keywords:} agriculture, allelic diversity, genetic drift, selection, temporal evolution

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\section*{Introduction}

The influence of human-mediated selection is perhaps nowhere more prevalent than in the agricultural system. Agricultural weeds, in particular, provide excellent case studies of adaptation to human-mediated selection (Baker 1974). They are exposed to fertilizers, herbicides, irrigation, as well as variable cropping techniques, and these manipulations can impose frequent, strong and highly predictable disturbance regimes (Barrett 1988). Examples of rapid adaptation to these scenarios are present in the literature from early cases of crop mimicry (Baker 1974; Barrett 1983) to the many recent examples of the evolution of herbicide resistance (Barrett 1988). Weedy plants, broadly defined as ‘plants that are growing out of place’ (Kuester \textit{et al.} 2014), are models for understanding rapid evolution and persistence in stressful environments. We currently have a limited understanding, however, of the broad genetic changes that may influence weed populations growing in agricultural landscapes (Vigueira \textit{et al.} 2013; Waselkov & Olsen 2014). These lapses in our knowledge are striking because the population dynamics of agricultural weeds are directly relevant to the global food supply. Agricultural weed infestations reduce worldwide crop
yield by as much as 10% (Oerke 2005), and it has been estimated that crop losses caused by weeds cost the US agricultural economy ~$33B USD per year (Pimentel et al. 2005). Clarifying the evolutionary forces that impact agricultural weeds can provide information on the process of rapid evolution more broadly as well as insight on how weeds survive and persist in agricultural regimes.

Agricultural weeds, which coexist with crops, evolve though unintentional human-mediated selection rather than through direct artificial selection (Stewart & Warwick 2005) and as such they exist in a state that is considered ‘neither wild nor domesticated’ (Vigueira et al. 2013). Weeds are subject to the same forces influencing evolution in nature – notably, genetic drift, selection and gene flow (Jasieniuk et al. 1996) – but they often experience a selection intensity that is much higher than what is usually found in other natural systems. For example, the predominant form of weed control in current farming is through the use of herbicides, which are designed to remove 90% of the weed population (Jasieniuk et al. 1996; Délye et al. 2013). Individuals that survive this high intensity of selection due to either chance or genetic predisposition are founders for the next generation. As the point of selective control – whether through the use of herbicide or another control technique – is to remove a large portion of the population, populations that recolonize are hypothesized to show evidence of genetic bottlenecks (Jasieniuk et al. 1996; Vigueira et al. 2013). As a result, weeds could lose rare alleles important to future adaptation (Nei et al. 1975).

In support of this idea, population genetic surveys have found that weeds tend to exhibit less genetic variation than other groups of plants (Hamrick et al. 1979), and there is some evidence that weed populations from cultivated land exhibit decreased neutral genetic diversity compared to wild populations (Kane & Rieseberg 2008). The majority of the work to date, however, has compared populations across space, that is from cultivated and noncultivated areas (Muller et al. 2010), or ‘wild’ vs. ‘weedy’ populations (Kane & Rieseberg 2008). In contrast, a novel approach that can provide direct evidence for evolutionary change through time is by the use of a resurrection approach in which ancestor and descendant strains of species are compared. In this type of experiment, seeds or propagules sampled from an earlier time point are germinated after remaining dormant for a number of years and compared to descendant populations sampled from the same location (Franks et al. 2007; Orsini et al. 2013). Although resurrection experiments have been used to address key questions about evolutionary constraints in microbial systems (Lenski & Travisano 1994; Lenski 1998), such experiments in eukaryotes have thus far used either a limited number of accessions (Baucom & Mauricio 2010) or a limited number of distinct populations (Franks et al. 2007; Thomann et al. 2015).

Here, we perform a resurrection experiment to determine whether populations of an agricultural weed exhibit evidence of genetic bottlenecks and phenotypic evolution over time. To do so, we use temporally sampled populations of Ipomoea purpurea, an introduced invader of agricultural and disturbed areas in the United States (Defelice 2001). Ipomoea purpurea is native to the central highlands of Mexico (Clegg & Durbin 2000; Defelice 2001), and lineages sampled from natural populations in the USA exhibit low diversity relative to Mexican accessions, suggesting a severe bottleneck occurred following introduction (Fang et al. 2013). Recent work shows a mosaic of glyphosate resistance in populations of I. purpurea across the USA, with some populations exhibiting high resistance (a high proportion of the population that survives glyphosate) and others showing high susceptibility post-herbicide application (Kuester et al. 2015). Previous work has also found that an additive genetic basis underlies glyphosate resistance in I. purpurea (Baucom & Mauricio 2008) and that resistance segregates in genetic lines developed from a single experimental population (Debban et al. 2015).

Although populations of I. purpurea are found primarily within agricultural fields that are treated with glyphosate and other herbicides, the impact of such strong selection and any associated environmental changes on the population genetics of this species remains largely unknown. Given genetic variation underlying resistance, the consistent application of glyphosate should lead to both genotypic and phenotypic evolution, that is evidence of genetic bottlenecks and increased resistance. Here, we test the prediction that agricultural populations, consistently exposed to herbicide over a nine-year period, show both reduced genetic diversity and increased resistance using temporally sampled cohorts of I. purpurea populations. Specifically, we first determine whether the neutral genetic differentiation and diversity of I. purpurea populations have changed between sampling years. We pair this with greenhouse experiments to examine the potential that these populations, sampled from the same fields nine years apart (Fig. 1; Table S1, Supporting information), exhibit increased resistance over time. We find evidence of both genetic bottlenecks and slight increase in the level of resistance, indicating that a noxious weed can adapt to the extreme selection imposed by herbicide applications even as genetic diversity decreases. We further find some indication that highly resistant populations exhibit lower genetic diversity than less resistant
populations, suggesting that herbicide application is responsible for the reduction in neutral genetic diversity. This is the first examination, to our knowledge, of a resurrection experiment that simultaneously identifies both loss of genetic diversity of an agricultural weed over time and potential evidence for adaptive evolution.

Materials and methods

Population sampling

Locations and sampling strategies for 44 Ipomoea purpurea populations were previously described in Kuester et al. (2015). Twenty-six of these populations were sampled in 2003 and resampled in 2012 (see Fig. 1 and Table S1, Supporting information). In 2003, we collected replicate seeds from between 6 and 30 maternal individuals at least 1 m apart from one another along a linear transect. We located the same populations in the fall of 2012 using GPS coordinates, which are accurate to within a few metres. Agricultural fields are highly disturbed by tilling and harvesting each year, and morning glories are predominantly found in areas that have recently experienced soil disturbance via tilling; as a result, this system is not amenable to the maintenance of long-term transects. We are thus making the assumption that adult plants present within the same agricultural field and located within the nearest distance to the GPS coordinates in 2012 are the descendants of the 2003 cohort. Preliminary data from >5000 SNPs generated by genotype-by-sequencing have identified a high number of independent genetic clusters in population structure analyses and a low proportion of recent immigrants into populations (D. Alvarado-Serrano & R. S. Baucom, unpublished data), indicating that our assumptions herein are largely realistic. We estimated population size in the 2012 sampling year by counting the numbers of individuals down a linear transect.

Of the 26 populations that were sampled both years, we randomly chose 10 to examine potential changes in genetic diversity between 2003 and 2012. One seed from an average of 18 maternal lines per population per sampling year (355 individuals total) was germinated, and cotyledons were used for DNA isolation using a CTAB method modified from Culley & Stokes, 2012 [see Kuester et al. (2015)]. The numbers of maternal lines sampled per population were approximately equal between the sampling years and exact numbers are presented in Table S2 (Supporting information).

To assay herbicide resistance among populations and between sampling years, we planted two replicate greenhouse experiments of all 26 populations at the University of Georgia Plant Biology Greenhouses (Athens, GA). One seed from 10 maternal lines per population per sampling year was scarified and planted in pine bark soil in SC10 super conetainers (Stuewe and Sons, Tangent, OR) in six experimental treatments, described below. This design was replicated in its entirety in another greenhouse for a total of 20 seeds per population within each treatment and thus an overall total of 5381 experimental individuals. Plants were randomly assigned to racks that were then randomly assigned to flow trays (4 racks per flow tray). Conetainers were watered daily, and flow trays were filled with water to prevent desiccation. Germination was slightly higher in 2003 compared to 2012 samples (87% and 84% in 2003 and 2012, respectively, \( \chi^2 = 12.27, P < 0.001 \)) and ranged from 50 to 98% across populations.

Plants were sprayed with RoundUp PowerMax® (Monsanto, St Louis, MO, USA) 22 days after planting at rates around the recommended field rate (1.54 kg ai/ha) of 0, 0.21, 0.42, 0.84, 1.70 and 3.40 kg a.i./ha (the 0 kg a.i./ha control treatment was sprayed with water) using a hand-held, CO₂-pressurized sprayer (R & D Sprayers, Opelousas, LA, USA) that delivered 187 L/ha at 206 kPa, 1.5 m above the plants. Three weeks after

![Fig. 1 Map of populations sampled from (A) 2003 and (B) 2012 within the USA. Populations that were genotyped in both 2003 and 2012 are indicated by a triangle (see Table S1, Supporting information for sites used for resistance and growth trait measurements). The per cent survival following 3.4 kg ai/ha of RoundUp® is indicated in colour. Sites were sampled at least 5 km apart.](image-url)
glyphosate application, we scored survival of each plant. Plants were harvested, dried at 72°C for 48 h and measured for total above ground biomass. Biomass values were adjusted to the nonsprayed controls by dividing each individual by the average biomass of its population grown in the nonspray control treatment following standard protocols (Tehranchian et al. 2015). At the time of sampling survival and biomass remaining post-herbicide (3 weeks after herbicide application), none of the plants exhibited signs of regrowth indicating that our measure of resistance does not confound resistance with tolerance (see Baucom & Mauricio 2008).

SSR genotyping and scoring errors
Details on multiplexing SSR markers and scoring procedures can be found in Kuester et al. (2015). Briefly, 15 polymorphic microsatellite loci were used to examine genetic diversity across populations and sampling years, and all individuals were scored by hand. To check accuracy of multilocus genotypes we rescored loci from 200 randomly chosen individuals and found very few scoring errors. We did not find any large allele dropouts or errors due to stutter in any of the loci by population by year combinations. We also examined the influence of null alleles on genetic diversity and found little evidence that potential null alleles altered our estimates or the main conclusions. Details of these analyses are presented in the Supporting Information section.

Temporal genetic differentiation and diversity
We examined the potential that seeds sampled across collection years were genetically differentiated from one another in two ways. First, we estimated genetic differentiation between years (FRT) using hierarchical AMOVA in GENALEX v. 6.5 (Peakall & Smouse 2012). We also performed individual assignment (Paetkau et al. 1995; Cornuet et al. 1999) of individuals to sampling year using GENECLASS2 (Piry et al. 2004). For individual assignment, the inability to assign individuals to a specific sampling year would indicate that individuals sampled in 2012 had not diverged in allelic composition compared to the individuals sampled in 2003. We used the Bayesian method described by Baudouin & Lebrun (2000) as a criterion for computation, and individual assignment was performed using the leave-one-out procedure (Paetkau et al. 2004), where the genotype to be assigned was not included in the population from which it was sampled. We report the –log likelihood of being assigned in each sampled year, by plotting the –log likelihood value of individual assignment to 2003 sample year against the –log likelihood of being assigned to the 2012 sampling year. Lack of temporal change across sampling years would be indicated by overlap of individuals sampled from each year. We calculated expected and observed heterozygosity (H_e and H_o), the number of alleles (Na) and the number of effective alleles (Ne) using GENALEX v. 6.5 (Peakall & Smouse 2012) and allelic richness (AR) using FSTAT v. 2.9.3.2 (Goudet 2005) and determined whether there were reductions in diversity estimates between 2003 and 2012 using Wilcoxon matched-pairs rank-sum tests (Zar 1996). We estimated the inbreeding coefficient (Fis) of each population in each sampling year using GENEPop v 4.5.1 (Rousset 2008) to determine whether there was evidence of inbreeding among populations and whether this significantly differed according to sampling year. Finally, we examined the possibility that populations experienced genetic bottleneck using the program BOTTLENECK (Piry et al. 1999). This program examines the potential for greater expected heterozygosity based on allelic diversity relative to expected heterozygosity estimated under mutation–drift equilibrium (Nei et al. 1975; Cornuet & Luikart 1996). If a significantly high proportion of loci exhibit an allele deficiency relative to expectations based on mutation–drift equilibrium, the population would show signs of a recent reduction in the effective population size and thus a bottleneck (Nei et al. 1975; Cornuet & Luikart 1996). We conditioned analyses on the infinite alleles model (IAM), the stepwise-mutation model (SMM) and the two-phase model (TPM) of microsatellite mutation as we are using microsatellites with a range of repeat motif types – dimeric, trimeric and imperfect motifs – and thus, we have no a priori reason to select one particular mutational model over another [repeat types presented in Table S2, Supporting information of (Kuester et al. 2015)]. All analyses were performed across 1000 iterations assuming mutation–drift equilibrium, and significance was calculated using the Wilcoxon test [appropriate for sample sizes of <30 individuals (Luikart & Cornuet 1998; Luikart et al. 1998)].

Resistance screen
We examined the potential that populations and sampling years varied for resistance using univariate mixed-model analyses of variance. We operationally defined resistance in two ways – first, as a measure of the number of individuals within populations that died as a result of herbicide application, and second, as a measure of the amount of biomass change following herbicide application standardized to controls. Because none of the experimental individuals showed signs of regrowth when survival and biomass postspay were measured, our operational measures of resistance do not conflate resistance with tolerance (which is the ability to regrow following damage). We used the glmer option of the
Ime4 package in R (Bates et al. 2011) to model survival as a binary character; further, we used the lmer option to assess biomass remaining postherbicide. In each model, replicate greenhouse experiment, herbicide treatment, collection year and population were the independent variables with survival or standardized biomass as the dependent variables. We included interactions between population and collection year as well as population, collection year and treatment. Population and its interaction terms were considered random effects in each model whereas all other effects were fixed. We previously identified a significant population effect from the 2012 cohort for survival postherbicide application, which indicated that populations vary in their respective level of resistance (Kuester et al. 2015). Here, we are specifically interested in the year term as well as interaction terms including the year effect, which would indicate that resistance varies between sampling years and/or that populations vary in their level of resistance between years. An F-test was used to determine the significance of fixed effects, and the significance of each random effect in the model was determined using a likelihood ratio test (LRT) in which the full model was compared to a reduced model with the effect of interest removed. The $P$-value was determined using a chi-square test with one degree of freedom. Prior to analysis, we examined the normality of our estimates of biomass with the Shapiro–Wilk test and by visual inspection of quantile–quantile (q–q) plot, and square root transformed this variable to improve normality of the residuals.

Results

Genetic diversity and differentiation

We uncovered reductions in genetic diversity between sampling years among populations (Table 1), with most measures of diversity significantly reduced in 2012 compared to 2003 (Fig. 2). For example, expected heterozygosity was 32% lower in 2012 ($W = 51$, $P = 0.01$), allelic richness was 18% lower ($W = 52$, $P = 0.01$), the effective number of alleles was 43% lower ($W = 51$, $P = 0.01$) and the absolute number of alleles per locus were reduced by 19% in 2012 compared to 2003 ($W = 50$, $P = 0.01$). The observed heterozygosity was 27% higher, on average, in 2012 compared to 2003 ($W = 4$, $P = 0.005$). This difference is probably due to the low observed compared to expected heterozygosity of the 2003 cohort; that is, the inbreeding coefficient ($F_{IS} = 1 - \frac{H_o}{H_e}$) was higher in 2003 vs. 2012 ($F_{2003} = 0.57 \pm 0.05$ (±SE) vs. $F_{2012} = 0.13 \pm 0.04$, respectively; Fig. 2). The difference in average $F_{IS}$ value between 2003 and 2012 was significant ($W = 55$, $P < 0.01$). Although this difference could be due to selection against heterozygotes in 2003, it is more likely indicative of differences in the mating system between sampling years of this mixed-mating, hermaphroditic species. Populations were sampled during a slightly longer window of time in 2003 than in 2012 (10/10–11/3 in 2003 vs. 10/15–10/20 in 2012); however, at least five of the 10 populations were sampled during the same temporal window (10/10–10/20 both years), and these populations exhibit similar differences in $F$ values ($F_{2003} = 0.47 \pm 0.08$ vs. $F_{2012} = 0.12 \pm 0.03$). We do not have information regarding pollinator abundance or any other reason to expect differences in the mating system between years.

Bonferroni-corrected HWE tests, consequently, indicated that more loci were not in HWE equilibrium within populations in 2003 (39 of 150 locus × population combinations), compared to 2012 (1 of 150 locus × population combinations). Processes that lead to heterozygote deficit, such as inbreeding or population substructure, can cause deviations from HWE; alternatively, the presence of null alleles could inflate estimates

<table>
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of homozygosity and lead to deviations from HWE (Kelly et al. 2011). We tested for the potential that null alleles altered our estimates of genetic diversity by removing loci with >25% putative null allele frequencies across populations and re-estimated indices of genetic diversity. We found no evidence that loci with potential null alleles altered our estimates of genetic diversity or the conclusion that diversity is altered between sampling years (Table S3, Supporting information).

We further examined changes in the patterns of allelic diversity by investigating the number of alleles, the number of rare alleles (<10% frequency) and the frequency of rare alleles present in 2003 and 2012. At the species level (i.e. across all populations), we found no evidence for a reduction in the total number of alleles from 2003 to 2012 (42 vs. 44 alleles in each year, respectively) – unexpectedly, we found fewer rare alleles in 2003 than 2012 (10 vs. 17). Only four of the rare alleles present in 2003 were likewise present in 2012, and their frequency was not dramatically increased as would be expected if rare allele frequency changes were responsible for the higher observed heterozygosity in 2012. When examining the number of alleles per population, however, we found that the total number of alleles was reduced in eight of 10 populations, by as much as 12–40% across populations. Two of the ten populations (populations 26 and 28) exhibited gains of low frequency alleles (between 4 and 5 new alleles present in 2012 at frequencies of <10%). Thus, 8 of the 10 populations show reductions in diversity over time likely due to random genetic drift, whereas two of the populations exhibit an increase in the number of alleles, putatively due to migration, drift or mutation.

We tested for a signature of bottleneck events in both the 2003 and 2012 samples using the BOTTLENECK program and found that significance of these tests
dependent on both the specific model employed (IAM, SMM or TPM) and the sampling year. Under the IAM, six populations sampled from 2003 exhibited significant heterozygote excess following corrections for multiple tests, whereas only one – population 32 – exhibited significant heterozygote excess under all three models of microsatellite evolution (Table 2). No populations from 2012 exhibited evidence of heterozygote excess and thus signs of a bottleneck following corrections for multiple tests (Table 2).

We next estimated the effective number of individuals from each sampling year using expected heterozygosity and the equation \( H_e = 4 N_e \mu \) (Nagylaki 1998) with a mutation rate, \( \mu \), of \( 10^{-3} \) (Marriage et al. 2009). We found that the estimated number of individuals from the 2003 populations was significantly higher, on average, compared to that of the 2012 populations (\( N_e \), 2003 = 85, \( N_e \), 2012 = 58; \( W = 67, P = 0.005 \)). Furthermore, we found no significant difference between our census sample size from the 2012 populations and the estimated effective number of individuals from that sampling year (Population size average from census = 70 individuals; \( W = 36.5, P = 0.32 \)). While the difference in estimated number of individuals between sampling years indicated that most populations experienced reductions in size (reductions ranging from 20 to 55 individuals fewer in 2012), populations 26 and 28 both exhibited an estimated gain of 20 individuals.

In line with lower diversity of the majority of populations, we found significant genetic differentiation between individuals sampled from different collection years (AMOVA year effect, \( F_{RT} = 0.218, P = 0.001 \), Table S4, Supporting information), and evidence that individuals sampled as seed in 2003 were more similar to one another than to individuals sampled as seed from the same location in 2012 (Fig. 3); that is, no individual assigned to 2003 was likewise assigned to 2012. We found that the estimate of \( F_{RT} \) was inflated by loci that potentially harboured null alleles; however, after removing these loci from analysis, we found that the \( F_{RT} \) estimate was still significantly different from zero, indicating the presence of between-year genetic differentiation (\( F_{RTo} = 0.133, P = 0.001 \), Table S4, Supporting information). We did not remove loci with potential null alleles from genotypic assignment as these tests are not greatly influenced by their presence (Carlsson 2008).

**Phenotypic evolution**

We examined resistance traits (survival and biomass post-herbicide application) to determine whether there was evidence of changes in resistance between sampling years. Our mixed-effects analyses of variance uncovered a significant year effect for biomass remaining after herbicide application (\( F_{1.3595} = 4.72, P = 0.03 \); Table 3). On average across all populations, the biomass remaining postspray of the 2012 cohort was slightly greater than that of the 2003 cohort (62% vs. 57% in 2012 and 2003, respectively) suggesting moderate

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**Table 2** Tests of heterozygosity excess within populations for each sampling year using the BOTTLENECK program (Cornuet & Luikart’s 1996). Tests were performed using three different models of microsatellite evolution, each of which assumes mutation-drift equilibrium (IAM, infinite alleles model; SMM, stepwise-mutation model; TPM, two-phase model). Probability values from one-tailed Wilcoxon tests are shown, with bolded values indicating statistical significance following corrections for multiple tests (\( P < 0.005 \)).

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**Fig. 3** Scatter plots of log likelihood values from assignment tests of individual *Ipomoea purpurea* plants sampled in 2003 and 2012 based on genotypic data at 15 microsatellite loci. A higher position relative to the y-axis indicates a higher likelihood of being from 2012 pool of individuals, and a higher position relative to the x-axis indicates greater likelihood of being from 2003 pool of individuals.
increases in resistance across populations sampled in 2012 (see Table S5, Supporting information for averages (±SE) among all populations). Likewise, a higher percentage of individuals sampled in 2012 survived herbicide application compared to those sampled from 2003 (49% vs. 42%), but this difference was not significant ($F_{1,5365} = 2.58, P = 0.11$; Table 3).

As in previous work (Kuester et al. 2015), we identified significant population effects for both measures of resistance (Table 3), indicating that populations vary across the landscape for their relative level of herbicide resistance. Here, however, we also find population by year effects in each analysis, indicating that populations differ in their level of resistance across years (Survival, $\chi^2_{1} = 23.74, P < 0.001$; Biomass, $\chi^2_{1} = 7.92, P = 0.005$; Table 3), a result that was significant across all treatment levels of herbicide (Table 3). At the herbicide level closest to the field dose (1.7 kg ai/ha), 16 populations exhibited either the same or increased survival in 2012 compared to 2003 whereas 10 populations exhibited lower survival in 2012 compared to 2003 (Table S5, Supporting information). One population’s survival increased by 79% compared to 2003, indicating that some populations may respond more readily with increased resistance than others. These differences are likewise apparent at the highest dose of herbicide (3.4 kg ai/ha; approximately 2× the field dose), with a significant population by year interaction for both survival and biomass remaining postherbicide application (Survival: Population × Year, $\chi^2_{1} = 16.23, P < 0.001$; Biomass: Population × Year, $\chi^2_{1} = 4.11, P = 0.04$) indicating that the populations differed in resistance level between sampling years. Notably, three populations sampled from TN that were highly resistant in 2012 (Kuester et al. 2015) were similarly resistant in 2003 (Fig. 1A, B, shown at 2× field rate of RoundUp®). The majority of the significant increases identified in the 2012 cohort compared to the 2003 cohort were located in NC and SC (Fig. 1A, B) – while five populations from the 2012 cohort of this region exhibit resistance values significantly greater than the specieswide average (56% survival at 2× field rate of RoundUp®), the 2003 cohorts of these populations exhibited only ~14% survival at 2× field rate. Overall, we identified a slight increase in resistance between sampling years (biomass remaining post-herbicide), and found that the resistance phenotype appears to be dynamic between sampling years, with some populations (central TN) retaining high resistance between sampling cohorts at high levels of herbicide, some populations (Carolinas) showing increased resistance and other populations exhibiting resistance declines.

**Discussion**

Despite the ubiquity and persistence of weedy plant populations, there are few examinations of how their neutral and adaptive genetic diversity may change over time. Here, we use a resurrection experiment to show that populations of weedy Ipomoea purpurea sampled from crop fields concomitantly lose genetic diversity and show signs of potential adaptive evolution in...
herbicide resistance. Our experiments yielded three novel findings. First, we found that seed progenies from populations sampled in 2012 exhibited lower genetic diversity and higher genetic differentiation than seed progenies sampled from the same fields and locations in 2003, suggesting that populations have experienced genetic bottlenecks between sampling periods. Second, heterozygosity excess tests indicated that a significant genetic bottleneck probably also occurred prior to 2003, perhaps due to the dramatic increase in glyphosate use in the late 1990s [see Fig. 1 (Baucom & Mauricio 2004)]. Although we cannot ascribe the loss of neutral genetic variation to the widespread use of herbicide per se, we show that a resistance trait – the amount of biomass maintained following herbicide application – has increased, on average, from 2003 to 2012. We combine these results with a retrospective analysis of a larger and previously published data set showing that highly resistant populations sampled in 2012 exhibit significantly reduced heterozygosity and allelic richness estimates compared to less resistant populations. Below, we discuss each of these major findings.

**Reductions in genetic diversity between sampling years**

We currently have a very limited understanding of how agricultural regimes may influence the population genetics of agricultural weeds. Although processes such as tilling and herbicide use are hypothesized to result in genetic bottlenecks, previous examinations of agricultural weed populations have either failed to uncover substantial reductions in genomewide diversity (Kane & Rieseberg 2008) or have presented largely circumstantial evidence for bottlenecks [i.e. comparisons between species (Hamrick et al. 1979)]. The significant loss in diversity that we uncovered across populations of *I. purpurea* sampled from agricultural fields argues for bottlenecks that were either very strong or occurred more than once, or both. While there are no studies, to our knowledge, that examine the temporal genetics of agricultural weed populations for comparison, it is of note that the average loss of allelic richness that we identified across populations (on average 15% lower between cohorts) is similar in magnitude to that of introduced, colonizing species [18% loss compared to native populations (Dlugosch & Parker 2008)]. Furthermore, using expected heterozygosity estimates from each sampling year, we find that the estimated population sizes have decreased between 2003 and 2012, with the majority of the populations losing reproductive individuals. While we did not take population census data in 2003 for comparison, we find that the estimated population size in 2012 is not significantly different from the census size, suggesting that our estimated population sizes are decent approximations of the true census size. The majority of the populations exhibited loss of alleles between sampling years, however, two populations – #26 and #28 – exhibited gains of low frequency alleles, and the estimated sample size of these two populations likewise increased relative to other populations. The increased diversity of these populations likely was from dormant seeds, migration from another population, or possibly an effect of experimental sampling differences between 2003 and 2012. Emergence of seed stored in the seed bank is incredibly likely – this species can produce a large number of seeds in field conditions (between 3000 and 10 000 per individual), and these heavy, gravity dispersed seeds can remain dormant for ~20 years in the soil (Baskin & Baskin 2000).

Interestingly, while the loss of allelic diversity between 2003 and 2012 suggests a genetic bottleneck has occurred between sampling years, the within-population examination of heterozygosity excess (i.e. the bottleneck test) did not find evidence of genetic bottleneck in the 2012 populations. Instead, our tests of heterozygosity excess uncovered evidence of genetic bottleneck among six populations collected in 2003 under the IAM model, with one population exhibiting evidence of a genetic bottleneck under all three models of microsatellite evolution. Other studies using heterozygosity excess tests have reported limited support of genetic bottleneck in species known to have experienced population declines (Hufbauer et al. 2004; Peery et al. 2012) with the general conclusion that heterozygosity excess tests may be limited to severely bottlenecked populations (Peery et al. 2012). Our results suggest that a genetic bottleneck occurred in some populations prior to the 2003 sampling, possibly following the sharp increase in the widespread use of glyphosate across RoundUp® Ready crops [see Fig. 1 in (Baucom & Mauricio 2004)].

In this scenario, the loss of allelic diversity that we identified in 2012 may simply be a continuation of effects following the initial population bottleneck. Alternatively, it is possible that demographic bottlenecks occurred prior to 2003 and continued between 2003 and 2012, but to a lesser extent between years. While bottleneck tests indicated that a few of the 2012 populations exhibited evidence of genetic bottleneck, these tests were not significant following multiple test corrections. While it is possible that differences in sampling seed could be responsible for the lowered genetic diversity between years, that the majority (8 of 10) populations exhibited significant declines in allele number in 2012 compared to 2003, coupled with the results of bottleneck tests indicating genetic bottleneck prior to 2003 suggest that these populations have experienced demographic declines leading to a reduction in allelic diversity.
Another attribute of the data suggests populations experienced a genetic bottleneck prior to 2003 – we found more locus by population combinations out of HWE compared to 2012. In addition to the evolutionary forces of genetic drift, selection, mutation, migration and nonrandom mating (i.e. inbreeding or nonassortative mating), null alleles can also cause deviations from Hardy–Weinberg equilibrium and will appear as heterozygote deficiency (Dąbrowski et al. 2014). The pattern that we uncovered of low observed heterozygosity relative to expected heterozygosity in our 2003 populations is consistent with the presence of null alleles but also consistent with high levels of inbreeding. To investigate the potential that null alleles influenced our results, we removed loci with putative null alleles from analyses and found that estimates of genetic diversity remained lower in the 2012 populations compared to their 2003 counterparts; further, F indices remained significantly higher in the 2003 sample, implicating widespread inbreeding following a potential bottleneck. *Ipomoea purpurea* is a hermaphroditic species that displays a wide range of outcrossing rates in nature ($T_m$ range: 0.2–0.8, across 20 populations; Kuester et al. unpublished data); thus, it is plausible that inbreeding could follow a large demographic bottleneck in this species. The majority of our loci exhibited $F_{IS} > 0$ in 2003 across populations, further pointing to a scenario of inbreeding following a large demographic change rather than the influence of null alleles. Finally, it is of note that null allele detection methods have been shown to exhibit low reliability when applied to nonequilibrium populations and will overestimate their frequency when populations have recently experienced demographic bottleneck (Dąbrowski et al. 2014).

**Phenotypic evolution**

Recent work provides an interesting contrast between the phenotypic and neutral genetic variations spatially distributed within this system – while neutral genetic differentiation among 44 *I. purpurea* populations is low (i.e. $F_{ST} = 0.127$), populations are significantly differentiated for herbicide resistance across the landscape, with some populations exhibiting high and others very low resistance (Kuester et al. 2015). Our screen of herbicide resistance in 26 of these temporally sampled populations shows that, in addition to a mosaic of resistance across the landscape, the level of resistance has slightly increased, on average, between sampling dates. This finding is interesting in the light of the reduced neutral genetic variation that we identified in eight of our 10 temporally sampled populations, and alternatively, in the light of evidence of potential migrants in two of the 10 populations – reductions in diversity as well as influx of presumably nonadapted variation would either act to impede or to counteract adaptation. These forces, along with recent work showing a severe fitness penalty of herbicide resistance in this species (Van Etten et al. 2015), probably explain why the average increase in resistance that we identified among all populations was modest – perhaps the populations that maintained resistance between sampling years (TN populations) or those that exhibit large increases in resistance (NC/SC populations) were less influenced by susceptible migrants germinating from the seed bank and/or costs of resistance than other populations. The low average increase in resistance across populations could also be due to a range of other factors: it is possible that few populations house additive genetic diversity for resistance, populations may experience different selective regimes, or the response to selection via glyphosate has been constrained by bottleneck events. Interestingly, there was no evidence that plants were different in size between the years (data not shown), indicating that the increased resistance we detected is not due to plants simply being larger in the 2012 cohort and thus better able to withstand herbicide application.

Although we find evidence for a moderate increase in the level of resistance across populations, it is important to note that our phenotypic comparisons were made using field-collected seeds. Thus, our resistance estimates include potential genetic components as well as environmental and maternal effects. This could explain the slight increase in resistance over time: if more 2012 populations had experienced glyphosate application relative to the 2003 populations, we would perhaps be sampling from a subset of the population that experienced herbicide relative to plants that had not, potentially inflating estimates of resistance in the 2012 samples. In 2003, between 80 and 92% of the soya fields in the USA were RoundUp® Ready, and thus sprayed with the herbicide, whereas approximately 20% of corn was RoundUp Ready® in that sampling year. In 2012, 93% of soya planted was RoundUp Ready® and between 73 and 80% of corn was RoundUp Ready® (National Agricultural Statistics Service, 2015, www.nass.usda.gov). Thus, it is possible that our comparison of the temporally sampled phenotypes is influenced by exposure to the herbicide itself in 2012. We considered this using crop type as a proxy for herbicide use, and determined whether biomass remaining post-spray differed according to crop (restricted to soya and corn) separately for both 2003 and 2012. There was no difference in biomass remaining postricide according to crop either year – for example the biomass remaining postricide of morning glories sampled from soya was no different than those sampled from corn – suggesting that the population-level estimates of resistance are not dependent on the crop type (as a proxy for spraying regime) in one particular sampling year.
While we cannot conclude that the phenotypic evolution of resistance identified here is solely due to adaptive evolution, we have previously identified an additive genetic basis underlying glyphosate resistance in one population of this species (Baucom & Mauricio 2008), positive selection for increased resistance in the field (Baucom & Mauricio 2008), and further, have shown that resistance segregates in crosses (Debban et al. 2015) indicating that the genetic potential is present within at least one population. In general, our results suggest that resistance evolution is dynamic in this system, with some populations maintaining high resistance (as measured by survival) between sampling years, other populations exhibiting large increases in resistance and others exhibiting declines in resistance. Continued sampling and assessment of resistance across these populations over time will be necessary to determine whether the populations that exhibited large increases in resistance between 2003 and 2012 maintain high resistance as did populations from central TN. Identification of the genetic basis of resistance across populations, and an assessment of how alleles associated with resistance change over time will decisively test our hypothesis that selection from the use of this herbicide is leading to adaptation in natural populations.

**Has herbicide application caused the genetic bottleneck?**

The populations used in this study were all sampled from crop fields that were farmed prior to and from 2003 onward. While we do not have specific information on herbicide use over this time period, we have historical record for six of 10 years (Table S1, Supporting information) showing that these locations were used for corn and soya crops, both of which make use of herbicides – and largely glyphosate – for weed control. Although other environmental factors (e.g., those associated with climate change) could be responsible for the genetic bottleneck that we report herein, herbicide use is an obvious potential factor. We examined this idea by making use of a larger and previously published data set of 32 populations, sampled in 2012, for which we have estimates of both survival and genetic diversity (Kuester et al. 2015). We performed separate regressions of expected heterozygosity (He) and allelic richness (AR) on estimates of herbicide resistance (proportion survival) at 3.4 kg ai/ha of glyphosate and found a significant negative relationship; that is, more resistant populations exhibit lower genetic diversity (He vs. resistance: $R = -0.375$, $P = 0.04$; AR vs. resistance: $R = -0.345$, $P = 0.05$). There is thus some indication that selection via herbicide application has led to the genetic bottleneck among populations. Interestingly, a population used in both analyses – population 32 – exhibits high survival at 3.4 kg ai/ha of glyphosate, and was the population for which we found evidence of a pre-2003 bottleneck under all models of microsatellite evolution.

**Conclusions**

Weedy plant species found in agricultural fields experience strong selection and thus are hypothesized to be either plastic, capable of adaptation, or saved from extinction through gene flow (Baker 1974; Parker et al. 2003). Using a resurrection approach, we provide evidence that even though genetic variation is lost from the system, some populations show potential signs of adaptation to herbicide application. While previous work indicates that the majority of the gene flow across southeastern populations occurred prior to the widespread adoption and use of glyphosate, suggesting that resistance evolution is due to selection on standing or novel variation within populations, that we identified evidence of potential migrants into the 2012 gene pool (in at least two populations) does not allow us to rule out the hypothesis that resistance can be introduced from outside sources.

Further, while we find evidence of increased resistance, we also show that the absolute change between years was not drastic; large resistance gains were limited to particular populations. These data suggest heightened measures should be taken to reduce the likelihood that seeds are accidentally moved between crop fields through farm machinery or through contaminated seed lots. Finally, we have some evidence that the lower genomewide diversity identified across populations is due to the application of glyphosate; however, we note that we cannot rule out other potential factors, as other herbicides with different mechanisms of action are often applied in crops, other cropping techniques that reduce population sizes might have been employed, and it is also possible that populations have lost diversity due to changes in the climate. The results shown here suggest that this weed, while being a ‘general purpose genotype’ (Baker 1974; Chaney & Baucom 2014), is also capable of adaptive evolution even while losing significant allelic diversity. How likely future adaptation to novel selective forces may be in the future, in the light of reduced variation is unknown.

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**Data accessibility**

SSR Genotyping and morphological data are available on Dryad, doi: http://dx.doi.org/10.5061/dryad.38j37.

**Supporting information**

Additional supporting information may be found in the online version of this article.

**Appendix S1** Materials and methods.

**Table S1** Site and location information for *I. purpurea* populations used in genetic, growth and herbicide resistant assays.

**Table S2** Sampling information for temporally-sampled populations of *Ipomoea purpurea*.

**Table S3** The genetic diversity of populations between sampling years with loci with putative null alleles removed (retaining 8 of the original 15).

**Table S4** Analysis of Molecular Variance (AMOVA) of neutral genetic data estimated using *GENEALEx* v 6.5 (Peakall and Smouse 2012).

**Table S5** The (A) survival and (B) above-ground biomass (standardized to controls) at each treatment level population × year combination.