THE EVOLUTIONARY POTENTIAL OF BAKER’S WEEDINESS
TRAITS IN THE COMMON MORNING GLORY, *Ipomoea purpurea*
(CONVOLVULACEAE)¹

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• *Premise of the study:* Many reports have cited Baker’s list of weediness traits, or those that exemplify the “ideal” weed, yet few have considered the evolutionary potential of such traits as a group. Thus, it is unknown whether constraints on the evolution of increased weediness, such as a lack of genetic variation or genetic correlations between the traits, are present. *Ipomoea purpurea,* the common morning glory, is a problematic weed that exhibits many of Baker’s ideal weed traits.

• *Methods:* We used progeny from a half/full-sib breeding design in a series of three greenhouse experiments to assess the presence of genetic variation, narrow sense heritabilities, and genetic correlations in Baker’s growth, competition, and fitness “weediness” traits in two populations of *I. purpurea.*

• *Key results:* We uncovered genetic variation underlying reproductive fitness traits and competitive ability in at least one population, but no evidence of genetic variation underlying growth rate in either population. Genetic correlations between many of the weediness characters differed significantly from zero; however, their direction and/or magnitude differed between populations.

• *Conclusions:* We found that increased weediness in the common morning glory is more likely to occur through selection on reproductive output and competitive ability rather than through selection on growth rate. Assessing Baker’s traits in a quantitative genetics framework can provide a solid perspective on their evolutionary potential and a unique framework within which to determine how weeds will respond to different environmental stresses and/or scenarios of global climate change.

Key words: Baker’s traits; Convolvulaceae; genetic correlations; genetic variation; heritability; *Ipomoea purpurea*; morning glory; weed.

In 1965, Baker produced a list of weediness traits, or those he thought defined the ideal weed (Table 1) (Baker, 1965, 1974). He compiled this list by comparing traits of weedy and nonweedy congeners; he considered species that exhibit many of these traits to be major weeds, whereas those that exhibit few are minor weeds. Baker’s list is heavily cited in both the invasive plant (e.g., Sutherland, 2004; Richards et al., 2006; Muth and Pigliucci, 2006; Whitney and Gabler, 2008) and weed ecology literature (e.g., Young and Evans, 1976; Holt and Orcutt, 1991; Perrins et al., 1992; Chao et al., 2005) and has been used primarily to predict which weeds will become invasive (Mack, 1996; Daehler, 1998; Goodwin et al., 1999).

Despite the focus on Baker’s list, the processes that govern the evolution of multiple weediness traits have yet to be considered within a species. This omission is especially relevant since Baker himself stated that “weeds are excellent subjects for the study of microevolution” (our italics, Baker, 1974, p. 1).

The evolution of weediness traits may play an important role in the establishment and spread of weedy and invasive species (Sakai et al., 2001; Dlugosch and Parker, 2008). If weediness traits such as competitive ability or relative growth rate lack genetic variation, the potential for the species to evolve into a more troublesome weed might be limited (Maynard Smith et al., 1985; Falconer and Mackay, 1996). While genetic variation has been found in floral and plant growth traits of some weedy plants (Conner and Via, 1993; O’Neil, 1997; Sahli et al., 2008), the evolutionary potential of multiple Baker’s traits has yet to be explicitly considered within a single species. In addition, the possibility that such traits are correlated and thus potentially act as constraints on their respective or dual evolutionary trajectories has not previously been examined (Lande, 1979; Conner and Via, 1993; Stanton and Young, 1994; Agrawal and Stinchcombe, 2009). Assessing Baker’s traits in a quantitative genetics framework can provide perspective on their evolutionary potential; such a view can also allow predictions to be made regarding how agricultural regimes may or may not select for more problematic weeds over time (De Wet and Harlan, 1975).

The common morning glory, *Ipomoea purpurea* (L.) Roth. (Convolvulaceae), a plant that exhibits at least nine of Baker’s 12 ideal weed traits (see boldfaced sections of Table 1), is often found in agricultural fields and is a highly effective competitor that restricts crop productivity by climbing onto and up crop species (Cordes and Bauman, 1984). Plants of this species can grow as rapidly as 20 cm per day (R. S. Baucom, unpublished data) and will produce many flowers in a growing season (daily: 0–80). Flowers are pollinated by bumblebees and generalist species (Epperson and Clegg, 1987; Chang and Rausher, 1998) but are also capable of self-fertilization (Ennos, 1981). Each plant can produce around 10,000 seeds each season (Crowley and Buchanan, 1982). Additionally, *I. purpurea* is tolerant of a variety of stressors including herbivores (Tiffin, 2002), pathogens (Simms, 1993), and herbicide (Baucom and Mauricio, 1993).

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is the mean of the ln-transformed plant biomass at time $$t$$.

Vigorous vegetative reproduction or regeneration from fragments

I. purpurea

Flowers open early in the morning and last a single day. In the United States, the plant senesces or killed by frost (Brown and Clegg, 1984; Uva et al., 1997). Commonly found in disturbed sites in the United States. Germination occurs from other seeds.

TABLE 1. Baker’s ideal weed characteristics. The traits of I. purpurea are in boldface, and traits examined in the present work are underlined. Adapted from Baker (1974).

<table>
<thead>
<tr>
<th>No.</th>
<th>Trait</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Germinates in many environments</td>
</tr>
<tr>
<td>2</td>
<td>Great longevity of seed</td>
</tr>
<tr>
<td>3</td>
<td>Rapid growth through vegetative phase to flowering</td>
</tr>
<tr>
<td>4</td>
<td>Continuous seed production</td>
</tr>
<tr>
<td>5</td>
<td>Self-compatible</td>
</tr>
<tr>
<td>6</td>
<td>When cross-pollinated, unspecialized visitors or wind used</td>
</tr>
<tr>
<td>7</td>
<td>Very high seed output</td>
</tr>
<tr>
<td>8</td>
<td>Tolerant and plastic</td>
</tr>
<tr>
<td>9</td>
<td>Has adaptations for short- and long-distance dispersal</td>
</tr>
<tr>
<td>10</td>
<td>Vigorous vegetative reproduction or regeneration from fragments</td>
</tr>
<tr>
<td>11</td>
<td>Plant not easily drawn from the ground</td>
</tr>
<tr>
<td>12</td>
<td>Ability to compete interspecifically by special means (rosette, choking growth, allelochemicals)</td>
</tr>
</tbody>
</table>

2004). Thus, this species could be classified as one of Baker’s major weeds.

Here, we assessed the evolutionary potential of Baker’s ideal weed traits in I. purpurea using a common garden greenhouse study. Although this species exhibits the majority of Baker’s weediness traits, we focused our efforts on estimates of growth, competition, and reproductive fitness (Table 1, underlined and boldfaced). We asked: (1) Is there evidence of genetic variation underlying Baker’s weediness traits in I. purpurea? (2) Are the traits genetically correlated to one another such that selection on one trait could cause a constraint on the evolution of the other?

MATERIALS AND METHODS

Study species—The common morning glory, Ipomoea purpurea (L.) Roth. (Convolvulaceae), is a climbing annual vine native to central America and commonly found in disturbed sites in the United States. Germination occurs from mid-May to August and flowers are produced about 6 wk after emergence until the plant senesces or killed by frost (Brown and Clegg, 1984; Uva et al., 1997). Flowers open early in the morning and last a single day. In the United States, I. purpurea is often found in maize, soy, and cotton fields and can cause significant yield losses in crop yield (Cordes and Bauman, 1984; Defelice, 2001), which has led to its inclusion on the list of the top 10 troublesome weeds in agriculture in the southern United States (Webster and Coble, 1997).

Crossing design—A partial diallel crossing design was used to generate experimental individuals. Two populations with different collection and breeding histories were assessed so that a broad perspective of the variation within this species could be examined. One population (hereinafter NC), generously provided by M. Rausher, was collected in 1985 in Durham, North Carolina and inbred in the greenhouse for about 15 generations. The other population (hereinafter GA) was originally collected from an agricultural field in Oconee Co., Georgia in 2001 and selfed once in the greenhouse. Crosses were performed for each of the two populations using a full/half-sibling breeding design, with two diallels per population and 10 maternal lines per diallel, yielding 40 maternal lines total. One seed from each of the 40 maternal lines was planted on 5 August 2010 in the University of Cincinnati greenhouse in a 4-inch pot with a general-purpose soil (Fafard 3B Mix, Agawam, Massachusetts, USA), and fertilized with a 20-4-8 NPK fertilizer (Scott’s Osmocote Pro, Marysville, Ohio, USA). Plants were exposed to supplementary sodium halide lights on a 16-h day schedule. Pollinations were performed by touching anthers of the parental plant to the stigma of the maternal parent. To prevent self-pollination, we emasculated all individuals the night before crosses were made. For this partial diallel crossing design (Simms and Rausher, 1987), each individual was crossed as a maternal plant to five paternal plants, and as a paternal plant with five maternal plants (Fig. 1). These crosses produced 200 families total. Replicate seeds from these families were then used for the three common-garden greenhouse experiments described next. The destructive harvest in two of our response variables necessitated a series of three experiments. Due to greenhouse space and seed number limitations, not all 200 families were used in each experiment; however, the highest number possible of common families was used to provide uniformity across experiments.

Experimental procedure—Relative growth rate—in the first experiment, we assessed variation in the relative growth rate of I. purpurea to address Baker’s “rapid growth through vegetative phase to flowering” weediness trait. On 20 December 2010, we scarified and planted eight replicate seeds from 192 families for a total of 1536 individuals. Seeds were planted in supercell containers (164-ml capacity; Stuewe and Sons, Tangent, Oregon, USA) in general-purpose soil (Fafard 3B Mix) in a randomized design in the University of Cincinnati greenhouse and watered daily. After 4 wk of growth, shoots of half of the replicates were harvested. Fourteen days later, shoots of the remaining four replicates were harvested. All plants were in a vegetative phase at the time of both harvests. All shoot mass was dried in a drying oven at 70°C for 3 d and weighed to measure dry biomass. Relative growth rate (RGR) for each family was calculated using the mean natural logarithm-transformed plant biomass ($$W$$) to avoid bias in the formula

$$RGR = \frac{\ln W_f - \ln W_i}{t_f - t_i},$$

where $$\ln W$$ is the mean of the ln-transformed plant biomass at time t (Hoffmann and Poorter, 2002).

Competitive ability—A second experiment was used to examine Baker’s competitive ability (“the ability to compete interspecifically by special means such as choking growth”) in two ways: (1) as the date that plants “grabbed on” to a competitor maize plant or nearest object (such as a bamboo stake); (2) as an estimate of dry biomass of plants grown in competition with maize. A plant was considered to have “grabbed on” when the plant’s vining stem created a C-shaped hook around the maize plant or bamboo stake. Three replicates of 145 families were harvested. All plants were harvested on 30 March and dried in a drying oven at 70°C for an estimate of dry biomass.
Flowering—On 23 February 2011, we planted three scarified replicate seeds of 196 families (588 individuals total) in supercell containers (164-mL capacity; Stuewe and Sons), with general-purpose soil (Fafard 3B Mix), in a completely randomized design in the University of Cincinnati greenhouse. Plants were fertilized and treated with insecticide as needed throughout the duration of the experiment. The length of flowering and total number of flowers was collected once flowering began on 11 April. Every 2 d, the flowers and buds on each plant were counted. Counting the flowers and buds every other day leads to a good approximation of the total number of flowers produced by each individual plant since the flowers of I. purpurea are open for a single day and flower buds are present on the day prior to opening. Flowering data were collected until the end of the experiment on 5 July, 4 mo from the initial planting, which is approximately the length of time I. purpurea plants flower in field conditions. We used total number of flowers and length of flowering as a proxy for Baker’s “very high seed output” and “continuous seed production” weediness characters since our studies were performed in the greenhouse without natural pollinators and since releasing natural pollinators in the greenhouse was not an option. Further, total number of flowers in this species is an excellent predictor of total number of seed (r = 0.924, t = 13.23, N = 32 (maternal lines), P < 0.0001; R. S. Baouco and R. Mauricio, unpublished data); thus, we can be reasonably certain that the length and total number of flowers produced provides a good approximation of reproductive fitness in this annual species.

Data analysis—Data proofing—Data were used only when the plants germinated, survived throughout the duration of the experiment, and in the flowering experiment, flowered. Examination of the data revealed that total number of flowers was nonnormal and exhibited positive skew; thus, we performed a log (y + 1) transformation, which improved normality upon examination of the residuals.

Genetic variation in Baker’s weediness traits—We used a series of univariate mixed model REML analyses of variance to test for the presence of genetic variation in the following Baker’s weediness traits: relative growth rate, total number of flowers, length of flowering, and two measures of competitive ability (see underlined sections of Table 1). The MIXED procedure of the SAS Statistical Software (version 9.2; SAS, Cary, North Carolina, USA) was used to perform separate analyses of each population. We analyzed models for the relative growth rate and flowering experiments with the model (1) trait = µ + diallel + maternal line (diallel) + paternal line (diallel) + + and for the competitive ability experiment with the model: (2) trait = µ + treatment + + where µ is the intercept or mean of the model (fit by default), treatment and replicate diallel as fixed effects and terms involving pedigree (maternal and paternal line) as random effects. The significance of random effects was determined using the likelihood-ratio statistic. The difference between the −2 log likelihood of the full model and the model with the random effect removed was calculated, and the P-value was determined with a χ² test with one degree of freedom. Significant paternal effects would indicate the presence of additive genetic variation, whereas a significant maternal effect indicates the presence of additive and nonadditive genetic variation. The interaction effect of maternal and paternal line was not included in the final model as preliminary analysis indicated that it was not a significant effect, and further, its inclusion does not inform the main conclusions of this study.

To determine whether genetic variation was present for growth rate in the two I. purpurea populations, we used relative growth rate of each family as the dependent variable and diallel and maternal and paternal line nested within diallel as independent variables. Maternal and paternal lines were random effects in the model, and diallel was considered a fixed effect.

Variation in competitive ability was tested using a mixed model with either biomass or the day that the vine grabbed on as the dependent variable, maternal and paternal line nested within diallel, the treatment effect (maize or no maize) and the interaction of maternal and paternal line with treatment as independent variables. Maternal and paternal lines were included as random effects in the model with treatment and diallel as fixed effects, and interaction effects were considered random.

Genetic variation underlying the total number of flowers and length of flowering was likewise tested using a series of mixed model analyses of variance with diallel and the maternal and paternal line nested within diallel as independent variables. Maternal and paternal lines were random effects in the model, and diallel was considered a fixed effect.

For each trait, narrow sense heritability (h²) was estimated as four times the parental variance component divided by the total phenotypic variance as determined in the mixed model ANOVAs (Falconer and Mackay, 1996). Significance tests of individual heritabilities are from the likelihood-ratio tests of the paternal line variance component of the earlier PROC MIXED models (Conner et al., 2003). Population differences in the traits were tested by using the LMEANS function in PROC MIXED with the effects as described already, this time including the term population as a fixed effect (Caruso, 2004).

Genetic correlations and covariances between Baker's weediness traits—We assessed the strength and direction of genetic correlations and covariances among paternal line means of weediness traits by performing the CORR COV procedure in SAS. Paternal line means were calculated by averaging trait values for each paternal line. The significance of genetic correlations and covariances was assessed by resampling the data using the jackknife procedure across each paternal line (Roff and Preziosi, 1994). A t-statistic was used to calculate a P-value for the confidence interval to determine whether the correlations between the two measures differed from zero. P-values were corrected for type I errors using the Bonferroni procedure.

RESULTS

Two of the five measures of Baker’s weediness traits—total number of flowers and length of flowering—differed between the GA and NC populations (F₁,₃₅ = 20.42, P < 0.0001 and F₁,₃₅ = 14.33, P = 0.001, respectively). The NC population flowered 76% longer and had nearly three times more flowers compared to the GA population (GA 6.03 ± 0.76, 18.10 ± 1.95 and NC 17.26 ± 1.67, 31.84 ± 1.81; total number and length of flowering means ± 1 SE, respectively). The estimates of relative growth rate and competitive ability (biomass with maize and day of grabbing on to maize) did not differ significantly between the two populations (F₁,₃₅ = 0.83, P = 0.370; F₁,₃₅ = 1.04, P = 0.314; and F₁,₃₅ = 0.94, P = 0.339; respectively).

We detected additive genetic variation in two of the five traits studied in the NC population (Table 2: i.e., a significant paternal line effect): total number of flowers and length of flowering. One measure of competitive ability, the day that the vine grabbed on to the maize plant or stake, exhibited a trend for paternal line variation (Table 2, P = 0.083). In comparison, the GA population showed no significant additive genetic variance in the weediness traits we investigated; however, this population exhibited a significant maternal line effect of day of grabbing on (Table 2). These results provide evidence of genetic variation in weediness traits underlying the reproductive fitness and competition in at least one population of I. purpurea.

The heritability of these traits ranged from 0 to 0.716 in the NC population (Table 3) and 0 to 0.484 in the GA population (Table 4). Although the heritability for total number of flowers in the GA population was moderate, our mixed model REML analysis did not uncover significant genetic variation in this trait (P = 0.129) in the greenhouse environment.

For both populations, there was a significant treatment effect of maize in terms of competitive ability, i.e., both the day of grabbing on and biomass of plants were significantly impacted by the presence of maize. This competition resulted in a 21% reduction in biomass of I. purpurea plants, and, on average, a lag of 2 d for grabbing on when plants were grown with maize. The interaction effect between maize and the maternal and/or paternal line was not significant in either population, suggesting that the paternal and maternal lines responded similarly in the presence of maize as in the presence of stakes (Table 2).

All of the genetic correlations and covariances among Baker’s weediness traits differed significantly from zero following Bonferroni corrections, with one noted exception in the GA population (Tables 3, 4). In both populations, the length of
Table 2. Results of mixed-model REML analysis demonstrating genetic variation for Baker’s weeding traits in two populations. Models were fit separately for each trait. The variance estimate is given for each random effect, boldfaced if significant, with P-value given in parentheses (calculated from log-likelihood F-statistic). For fixed effects, the F-statistic is given with P-value given in parentheses.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Population: North Carolina</th>
<th>Relative growth rate (P)</th>
<th>Total no. of flowers (P)</th>
<th>Duration of flowering (P)</th>
<th>Biomass (P)</th>
<th>Day of grabbing on to maize (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed effects</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paternal (Diallel)</td>
<td>0.000 (1.000)</td>
<td>0.000 (1.000)</td>
<td>0.000 (1.000)</td>
<td>0.000 (0.752)</td>
<td>0.000 (1.000)</td>
<td></td>
</tr>
<tr>
<td>Maternal (Diallel)</td>
<td>0.000 (1.000)</td>
<td>0.120 (0.048)</td>
<td>80.887 (0.003)</td>
<td>0.001 (0.439)</td>
<td>1.220 (0.083)</td>
<td></td>
</tr>
<tr>
<td>Pat × Maize (Diallel)</td>
<td>0.001 (0.655)</td>
<td>0.000 (1.000)</td>
<td>0.000 (1.000)</td>
<td>0.001 (0.439)</td>
<td>1.220 (0.083)</td>
<td></td>
</tr>
<tr>
<td>Fixed effects</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diallel</td>
<td>0.83 (0.374)</td>
<td>0.36 (0.557)</td>
<td>0.13 (0.724)</td>
<td>4.24 (0.054)</td>
<td>1.16 (0.296)</td>
<td></td>
</tr>
<tr>
<td>Maize</td>
<td>301.7 (&lt;0.0001)</td>
<td>22.51 (0.0002)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Narrow-sense heritabilities (in boldface on the diagonal) and paternal line mean correlations (above diagonal) and covariances (below diagonal) with jackknifed 95% CIs in parentheses, for Baker’s weeding traits in the North Carolina population.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Relative growth rate</th>
<th>Total no. of flowers</th>
<th>Duration of flowering</th>
<th>Biomass with maize</th>
<th>Day of grabbing on to maize</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative growth rate</td>
<td>0.000</td>
<td>−0.321 ± 0.015</td>
<td>−0.188 ± 0.025</td>
<td>−0.247 ± 0.027</td>
<td>0.421 ± 0.020</td>
</tr>
<tr>
<td>Total no. of flowers</td>
<td>−0.007 ± 0.006</td>
<td>0.416*</td>
<td>0.710 ± 0.012</td>
<td>0.048 ± 0.029</td>
<td>0.188 ± 0.030</td>
</tr>
<tr>
<td>Duration of flowering</td>
<td>−0.091 ± 0.012</td>
<td>4.229 ± 2.396</td>
<td>0.716**</td>
<td>0.205 ± 0.018</td>
<td>−0.111 ± 0.029</td>
</tr>
<tr>
<td>Biomass with maize</td>
<td>0.000</td>
<td>0.001 ± 0.008</td>
<td>0.088 ± 0.008</td>
<td>0.000</td>
<td>−0.575 ± 0.019</td>
</tr>
<tr>
<td>Day of grabbing on to maize</td>
<td>0.037 ± 0.002</td>
<td>0.201 ± 0.474</td>
<td>−2.635 ± 0.660</td>
<td>−0.045 ± 0.002</td>
<td>0.171</td>
</tr>
</tbody>
</table>

Notes: All correlations and covariances are significant with Bonferroni corrected P-values. Significance tests of individual heritabilities are from log-likelihood test of the paternal variance component (see Methods) and are denoted with asterisks: * P < 0.05, ** P < 0.01, *** P < 0.001.
To address competitive ability by choking growth, we recorded the day of grabbing on to a stake or plant competitor because we hypothesized that those plants that grab on earliest are likely the ones that also produce the most biomass in the presence of competition. We uncovered a trend for paternal line variation in the day of grabbing on in the NC population (P = 0.083), such that some paternal lines began to vine onto their nearest neighbor earlier than others. Furthermore, the GA population exhibited maternal line variation in this trait (P = 0.026), an effect that includes both additive and nonadditive components of genetic variance.

The presence of variation underlying this trait would be of little consequence if there were no size and/or fitness benefit of producing choking or vining growth early in the growing season. As such, we investigated the relationship between date of grabbing on and biomass while in the presence of maize competition, and we found a significant negative genetic correlation in both populations (GA: r = −0.730, P < 0.001; NC: r = −0.575, P < 0.001), suggesting that plants that vine onto competitors earlier are also larger and thus putatively better competitors. The present series of experiments cannot address whether vining capability is simply due to being larger or to an innately different plant sensory and/or climbing ability. Previous work with Ipomoea hederacea, sister to I. purpurea, found that plants of this species appear to respond to reflectance of nearby objects and to project stems toward neighboring structures such as differently colored bamboo stakes. Further, plants that were successful in climbing weighed more and produced more seeds than morning glories that did not climb (Price and Wilcut, 2007). Thus, it is possible that competitive “vining” capability could be due to more than size in I. purpurea.

The relationship between competitive ability and reproductive fitness differed in the two populations. Ipomoea purpurea plants from the NC population had a negative correlation between date of grabbing on and length of flowering (r = −0.111), suggesting that those that climbed onto their competitor earlier also flowered longer. There was a significant positive correlation between the date of grabbing on to a competitor and the total number of flowers produced (r = 0.188), suggesting that earlier vining would not necessarily produce a greater fitness return. Individuals from the GA population had the opposite relationship: there was a positive correlation between date of grabbing on and duration of flowering (r = 0.132). These results suggest a trade-off between competitive growth and reproductive fitness in this population—those that grab on earlier will flower for a shorter duration, whereas if they grab on later, they flower longer. Despite this potential trade-off, there was no detectable correlation in this population between the total number of flowers produced and day of grabbing on to the competitor. Although it is interesting that the two populations differ regarding presence/absence of genetic variation and the pattern of some genetic correlations, we do not know whether historical, geographic, collection time, or other differences such as breeding history are responsible for the differences. Further studies are required to parse out these differences in populations. We chose to assess these traits in more than one population in an effort to include a wider range of the potential variation in this species.

Our findings of significant genetic variation in reproductive fitness traits—total number of flowers and length of flowering—and a trend for variation in competitive ability suggests that selection on these traits could lead to the evolution of increased/decreased weediness. That we uncovered genetic correlations among such traits suggests that weediness should be considered in a multivariate context (Lande 1979; Agrawal and Stinchcombe, 2009). For example, if, in the NC population, either the total number of flowers or length of flowering were under positive selection, we would predict an increase in the overall reproductive fitness of the plant through the production of more flowers and over a longer period. If, however, selection favored a simultaneous increase in one trait and a decrease in the other, the positive correlation detected here may function as a constraint on the evolution of either trait (Lande, 1979; Lande and Arnold, 1983). Likewise, in the NC population, a trade-off potentially exists between being a grabbing competitor and overall fitness return in that there was a weak positive relationship between the total number of flowers produced and the time in which a plant grabbed on to a nearby object. This relationship might reflect a fitness cost of early grabbing in the absence of intense competition, a highly likely scenario for this species because it is often found vining up agricultural competitor crops. Although most of our genetic correlation values were significant, many of them were low; thus, the extent to which the correlations identified here represent evolutionary constraints is currently unknown. An assessment of these traits in a natural setting would provide insight on the pattern of natural selection on weediness traits.

Although our study explicitly assesses the evolutionary potential of the traits that exemplify Baker’s ideal weed, others have examined the evolutionary potential of some of these traits but in wholly different contexts. For example, additive genetic variation underlies flowering duration and seed number in purple loosestrife (Lythrum salicaria), a noxious semiaquatic perennial weed of North America (O’Neil, 1997). The work of Conner and colleagues finds significant genetic variation in many floral (Conner et al., 2003; Sahl i et al., 2008) and plant growth traits (Conner and Via, 1993) of wild radish (Raphanus raphanistrum). Beyond these two examples in weedy plants, genetic variation underlying floral traits is widely supported (Shore and Barrett, 1990; Maz er and Schick, 1991; Caruso, 2004; Conner, 2006), and across many organisms fitness traits in general are often reported to exhibit higher levels of additive

<table>
<thead>
<tr>
<th>Trait</th>
<th>Relative growth rate</th>
<th>Total no. of flowers</th>
<th>Duration of flowering</th>
<th>Biomass with maize</th>
<th>Day of grabbing on to maize</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative growth rate</td>
<td>0.000</td>
<td>±0.051 ± 0.027</td>
<td>0.271 ± 0.029</td>
<td>−0.523 ± 0.013</td>
<td>0.321 ± 0.020</td>
</tr>
<tr>
<td>Total no. of flowers</td>
<td>0.000 ± 0.000</td>
<td>0.484</td>
<td>0.670 ± 0.017</td>
<td>−0.067 ± 0.028</td>
<td>0.004 ± 0.021 NS</td>
</tr>
<tr>
<td>Duration of flowering</td>
<td>0.050 ± 0.005</td>
<td>±2.802 ± 0.137</td>
<td>0.101</td>
<td>−0.193 ± 0.017</td>
<td>0.132 ± 0.028</td>
</tr>
<tr>
<td>Biomass with maize</td>
<td>0.000 ± 0.000</td>
<td>±0.038 ± 0.003</td>
<td>0.028</td>
<td>−0.046 ± 0.002</td>
<td>0.0000</td>
</tr>
<tr>
<td>Day of grabbing on to maize</td>
<td>0.015 ± 0.001</td>
<td>0.004 ± 0.019 NS</td>
<td>1.797 ± 0.356</td>
<td>−0.750 ± 0.009</td>
<td>0.0000</td>
</tr>
</tbody>
</table>

Notes: All correlations and covariances are significant with Bonferroni corrected P-values, with two noted exceptions (NS). Although reported heritabilities in this population do not differ significantly from zero.
genetic variance than nonfitness traits (Houle, 1992; Merilä and Sheldon, 1999). Furthermore, the relationship between estimates of plant growth and fitness has been considered by others but yet again in different contexts than Baker’s weediness traits. In general, studies show that plants that flower later in the season exhibit greater biomass and/or size (Conner and Via, 1993; Mitchell-Olds, 1996; Latta and Gardner, 2009). While we find a significant negative correlation between relative growth rate and total number of flowers in one population (NC: $r = -0.321$), thus supporting the general finding of a trade off between growth and fitness, we find a significant but weak positive correlation between the total number of flowers and plant biomass in the same population ($r = 0.048$). However in the other population, we find a significant but weak positive correlation between relative growth rate and total number of flowers (GA: $r = 0.051$) and a significant but weak negative correlation between total number of flowers and plant biomass ($r = -0.067$). Since neither of the two investigated populations displayed evidence of genetic variation in relative growth rate and/or biomass, yet displayed evidence of variation in reproductive fitness, it is likely that selection for increased weediness in this species would be apparent through selection on fitness traits rather than growth rate and/or biomass traits. Evidence of genetic variation in both reproductive fitness and competitive ability suggests the relationship between the two traits deserves increased attention in this species.

Baker was the first to compile a list of traits that defined the ideal weed (Baker, 1965, 1974), and since then other lists have followed (De Wet and Harlan, 1975; Patterson, 1985; Roy, 1990). Although the ability to capture weediness in a well-defined and applicable way has appealed to many (Rejmánek, 2000), Baker’s list has been criticized as lacking predictive power (Perrins et al., 1992; Williamson and Fitter, 1996; Mack, 1996; Reichard, 1997). Despite this, surprisingly few reports have explicitly tested for the presence of genetic variation underlying Baker’s weediness traits as a group (Neve et al., 2009), and none have assessed the way in which the traits may be correlated, another oversight as these traits are likely to evolve together (Lande and Arnold, 1983). Addressing the evolutionary potential of traits thought to underlie plant weediness has the potential to help us determine which minor weeds may evolve into major weeds (Baker, 1974; Baker, 1991) and can provide a unique framework within which to determine how major and minor weeds will respond to different environmental stresses (Stanton et al., 2000) and/or scenarios of global climate change (Strauss et al., 2006).

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