Taxonomic and Life History Bias in Herbicide Resistant Weeds: Implications for Deployment of Resistant Crops

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Abstract

Evolved herbicide resistance (EHR) is an important agronomic problem and consequently a food security problem, as it jeopardizes herbicide effectiveness and increases the difficulty and cost of weed management. EHR in weeds was first reported in 1970 and the number of cases has accelerated dramatically over the last two decades. Despite 40 years of research on EHR, why some weeds evolve resistance and others do not is poorly understood. Here we ask whether weed species that have EHR are different from weeds in general. Comparing taxonomic and life history traits of weeds with EHR to a control group (“the world’s worst weeds”), we found weeds with EHR significantly over-represented in certain plant families and having certain life history biases. In particular, resistance is overrepresented in Amaranthaceae, Brassicaceae and Poaceae relative to all weeds, and annuality is ca. 1.5 times as frequent in weeds with EHR as in the control group. Also, for perennial EHR weeds, vegetative reproduction is only 60% as frequent as in the control group. We found the same trends for subsets of weeds with EHR to acetolactate synthase (ALS), photosystem II (PSII), and 5-enolpyruvylshikimate-3-phosphate (EPSP) synthase-inhibitor herbicides and with multiple resistance. As herbicide resistant crops (transgenic or not) are increasingly deployed in developing countries, the problems of EHR could increase in those countries as it has in the USA if the selecting herbicides are heavily applied and appropriate management strategies are not employed. Given our analysis, we make some predictions about additional species that might evolve resistance.

Introduction

Evolved herbicide resistance (EHR) has become a threat to agriculture and consequently a food security problem worldwide [1,2]. EHR in weeds was first reported in 1970 [3,4] and widely studied in the 1970s through 1990s [3]. The number of cases has accelerated dramatically over the last two decades [5]. The evolution of resistance to multiple herbicides with different modes of action has also been found within numerous weed species to date [6]. The discovery of resistance to glyphosate (the dominant herbicide worldwide) in the 1990s [7], the introduction of transgenic glyphosate-resistant crops in 1996 [8], and the recent expansion of cases of evolved resistance to glyphosate in weeds, likely due to greater glyphosate usage, have inspired a renewal of interest and resurgence of research into this phenomenon.

Despite 40 years of research on EHR, it is not clear why some weeds evolve resistance faster than others. Baker’s list of characteristics that might be expected in the “ideal weed” is well known [9]; one might expect weeds with EHR to possess a subset of these traits. A cursory review of available data in 2001 revealed that some weeds have a greater propensity to evolve resistance than others [10]. This observation was attributed to opportunity, as many resistant weeds are among the world’s worst weeds [11,12], are widespread, and occur in many cropping systems [10]. Well before EHR was discovered, heritable variability, breeding system, reproductive capacity, annuality, and population size were predicted to correlate with evolution of herbicide resistance [13]. Other plant factors can affect the evolution of resistance, including mutation frequency, generation time, fitness in absence of the herbicide, plasticity, and soil seed reservoir [14], as well as mode of inheritance of resistance, population size, seed dormancy, and gene flow by pollen and seed [15]. While these factors have been tested in models predicting evolution of resistance [14,16], few have been tested empirically.

Given that taxonomic families are relatively cohesive internally but generally vary from one another in ecological traits, we might expect EHR to be represented nonrandomly among plant families. Similarly, based on observations and reports of the propensity for resistance to evolve within certain genera or species, we would expect weeds with HER to be ecologically and taxonomically different than weeds in general. Here we ask whether weed species that have EHR are different from weeds in general and if evolution of resistance to multiple herbicides follows the same patterns.

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Materials and Methods

A list of species with evolved herbicide resistance EHR; (henceforth, the EHR list) was obtained from the database at the website “International Survey of Herbicide Resistant Weeds” http://www.weedscience.org/In.asp created and maintained by Ian Heap with support from the Herbicide Resistance Action Committee, the North American Herbicide Resistance Action Committee, and the Weed Science Society of America [5]. At the time of our study (May 2012), the inventory included 187 species in 31 plant families. For statistical comparison, we chose the two volumes by Holm et al. that inventory the “World’s Worst Weeds” and “World Weeds” [11,12] (henceforth the control list) including 201 species in 49 plant families. Entries from different subspecies of the same species were lumped together. The taxonomy for both lists was updated using the USDA ARS Germplasm Resources Information Network website “GRIN Taxonomy for Plants”, http://www.ars-grin.gov/cgi-bin/npgs/html/index.pl?language=en [17]. All scientific names were then checked against the “The Plant List” website database, http://www.theplantlist.org/, which was also used to obtain current plant family assignments. The two updated lists are presented in Table S1.

Information regarding plant life history was extracted from the Holm et al. [11,12] publications, if available. For species not present in those publications, the information was obtained from major floras (e.g. [18,19]) or citations for individual species entries in the USDA ARS “GRIN Taxonomy for Plants” [17].

All comparisons between the EHR list and the control list were done using a Parson’s chi-square statistic (2-tailed test) with one degree of freedom; \( \alpha = 0.05 \) was used to determine significance. This test was used because it is appropriate for comparisons between theoretical and experimental populations where a data set is large and observations are independent. Calculations were done using R statistical package [20].

We constructed a phylogenetic tree hypothesis for the relationship among plant families with EHR using Phylomatic 2 and the Angiosperm Phylogeny Group 3 derived megatree (http://www.phylodiversity.net/phylomatic/phylomatic.html). Presence and absence of EHR to acetylactate synthase (ALS)- and photosystem II (PSII)-inhibitors was mapped onto the same tree. Character state reconstruction for each EHR group was performed by maximum likelihood using a marginal probability reconstruction with the Asymmetrical parameter Markov-k model of evolution in Mesquite version 2.74 [21]. The results are displayed as likelihood states reported as proportion of the total likelihood and represented as pie charts in each branch node within the tree.

Results and Discussion

Comparisons between the control and EHR lists showed sixty-two species in 19 families are found in common on both lists, representing 33% of the 187 species on the EHR list and 31% of the 201 species on the control list. When ranked by number of species, the same plant families were dominant on both lists (Table 1). Poaceae, Asteraceae, Amaranthaceae, and Brassicaceae were among the top six families on each list, comprising 75% and 57% of species on the EHR and control lists, respectively. This is not surprising since plant families that are overrepresented in the global weed flora would have a greater probability of herbicide exposure and selection of species with EHR [10].

Despite the taxonomic overlap between the EHR and control lists at the family level, we found significant differences with regard to species abundance within families (Table 2). In particular, EHR was overrepresented relative to the control in three of the four aforementioned families (Poaceae, Amaranthaceae, and Brassica-
Table 2. Comparison of plant family and life history traits.

| Family       | All Resistance | ALS Resistance | PSII Resistance | Glycine Resistance | Multiple Resistance |
|--------------|----------------|----------------|-----------------|-------------------|---------------------|
|              | Control list (%) | Complete EHR list (%) | $\chi^2$ statistic | $p$-value | Difference from control list |
|              |                |                |                  |                  |                     |
| Family       |                |                |                  |                  |                     |
| Amaranthaceae| 5              | 10             | 8.55             | 0.004            | +                   |
| Brassicaceae | 4              | 9              | 17.55            | $2.87 \times 10^{-5}$ | + |
| Poaceae      | 23             | 32             | 7.91             | 0.0049           | +                   |
|              |                |                |                  |                  |                     |
| Life-history traits |          |                |                  |                  |                     |
| Annuality 3  | 53             | 86             | 64.26            | $1.08 \times 10^{-15}$ | + |
| Perenniarity | 43             | 11             | 64.32            | $1.06 \times 10^{-15}$ | - |
| Vegetative Reproduction 4 | 68          | 41             | 18.11            | $2.08 \times 10^{-5}$ | - |
|              |                |                |                  |                  |                     |
| Family       |                |                |                  |                  |                     |
| Amaranthaceae| 5              | 11             | 7.88             | 0.005            | +                   |
| Brassicaceae | 4              | 15             | 40.1             | $2.4 \times 10^{-8}$ | + |
| Poaceae      | 23             | 23             | 0.013            | NS                | 0                   |
|              |                |                |                  |                  |                     |
| Life-history traits |          |                |                  |                  |                     |
| Annuality 3  | 53             | 87             | 35.18            | $3.08 \times 10^{-9}$ | + |
| Perenniarity | 43             | 9              | 39               | $4.23 \times 10^{-10}$ | - |
| Vegetative Reproduction 4 | 68          | 48             | 4.91             | 0.03              | -                   |
|              |                |                |                  |                  |                     |
| Family       |                |                |                  |                  |                     |
| Amaranthaceae| 5              | 24             | 51.8             | $6.06 \times 10^{-13}$ | + |
| Brassicaceae | 4              | 6              | 1.3              | NS                | 0                   |
| Poaceae      | 23             | 30             | 2.21             | NS                | 0                   |
|              |                |                |                  |                  |                     |
| Life-history traits |          |                |                  |                  |                     |
| Annuality 3  | 53             | 97             | 35.41            | $2.7 \times 10^{-9}$ | + |
| Perenniarity | 43             | 2              | 43.34            | $4.59 \times 10^{-10}$ | - |
| Vegetative Reproduction 4 | 68          | 25             | 6.6              | 0.01              | -                   |
|              |                |                |                  |                  |                     |
| Family       |                |                |                  |                  |                     |
| Amaranthaceae| 5              | 14             | 3.85             | 0.049            | +                   |
| Brassicaceae | 4              | 0              | None with glycine resistance | | |
| Poaceae      | 23             | 48             | 6.88             | 0.0087           | +                   |
|              |                |                |                  |                  |                     |
| Life-history traits |          |                |                  |                  |                     |
| Annuality 3  | 53             | 176            | 3.73             | 0.05             | +                   |
| Perenniarity | 43             | 17             | 4.95             | 0.026            | -                   |
| Vegetative Reproduction 4 | 68          | 43             | None with glycine resistance | | |
|              |                |                |                  |                  |                     |
| Family       |                |                |                  |                  |                     |
| Amaranthaceae| 5              | 13             | 4.79             | 0.0286           | +                   |
| Brassicaceae | 4              | 8              | 1.92             | 0.1658           | +                   |
| Poaceae      | 23             | 48             | 12.99            | 0.000314         | +                   |
|              |                |                |                  |                  |                     |
| Life-history traits |          |                |                  |                  |                     |
| Annuality 3  | 53             | 94             | 20.38            | $6.35 \times 10^{-6}$ | + |
| Perenniarity | 43             | 3              | 19.66            | $9.26 \times 10^{-6}$ | - |

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**Evolution of Herbicide Resistant Weeds**

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inherited and easily transmitted by seed and pollen [23]. Perennial weeds are only 0.27 times as frequent in the EHR list as found that annual weeds are 1.5 times as frequent, whereas respect to plant habit (annuality, bienniality, perenniality). We understand underlying causes of the phylogenetic patterns number, dominance, and fitness cost of resistance genes) as well as about genetics of resistance to a particular herbicide (frequency, mutation to EHR for any weed species [15]. More information populations, however, little is known about actual rates of variation for resistance appears not to be uncommon in weed becomes increasingly difficult, the taxonomic distribution of EHR phylogenetic patterns in the future.

Panicoideae and Chloridoideae. Dramatic increases in glyphosate found in a single family, Poaceae, and six of the eight genera with variability in glyphosate susceptibility in weeds, EHR to this herbicide was not discovered for two decades following its introduction [8]. The relatively slow appearance of EHR to glyphosate compared to other herbicide groups was initially attributed to genetic and biochemical constraints, restricted use, and lack of soil residual activity resulting in low selection pressure for resistance [8]. Indeed, EHR to glyphosate currently shows a very restrictive phylogenetic distribution with almost half of species found in a single family, Poaceae, and six of the eight genera with EHR in that family are in two closely related subfamilies, Panicoideae and Chloridoideae. Dramatic increases in glyphosate use where transgenic glyphosate-resistant crops are planted have intensified selection pressure for EHR, which might alter these phylogenetic patterns in the future.

Cumulatively, our results for the three herbicide groups tested suggest that as selection of resistance to an herbicide group becomes increasingly difficult, the taxonomic distribution of EHR will be subject to increasing phylogenetic constraint. Genetic variation for resistance appears not to be uncommon in weed populations, however, little is known about actual rates of mutation to EHR for any weed species [15]. More information about genetics of resistance to a particular herbicide (frequency, number, dominance, and fitness cost of resistance genes) as well as about natural mutation frequencies, would improve our ability to understand underlying causes of the phylogenetic patterns detected here.

We also compared the full EHR list and the control list with respect to plant habit (annuality, bienniality, perenniality). We found that annual weeds are 1.5 times as frequent, whereas perennial weeds are only 0.27 times as frequent in the EHR list as in the control list (Table 2). We attribute this difference to the fact that short-lived species are recognized to have a more rapid response to directional selection than long-lived species, leading to greater opportunity for evolution of EHR [14,15]. If length of life cycle is inversely correlated with EHR, then we would predict that because vegetative reproduction effectively extends generation time, perennials with vegetative reproduction should be highly underrepresented among weeds with EHR. As expected, we found the proportion of vegetative reproduction in perennials with EHR to be only 0.6 times that in the control list. The same trends were found in the three herbicide groups evaluated with respect to plant habit and vegetative reproduction (Table 2).

While many traits are predicted to correlate with EHR [13,14,15], some (e.g., fitness, plasticity, soil seed reserve, population size, and gene flow) are strongly influenced by environment and difficult to include in a phylogenetic analysis. Other life history traits, e.g., outcrossing rate, spontaneous hybridization, fecundity, and seed size, might help explain phylogenetic patterns in EHR. Unfortunately, available data are insufficient to allow further quantitative assessment or meta-analysis of these traits for weeds with EHR compared to weeds in general.

A total of 139 species across 44 families on the control list had no reports of EHR. All of these species are found in countries for which reports of EHR occur [5]; thus, the absence of EHR cannot be explained by lack of herbicide exposure or EHR reporting. We note that aquatic and wetland species are significantly more numerous on the control list than the EHR list (χ² = 11.69, df = 2, p = 0.00063) [9,10]. Despite the fact that aquatic weeds are widespread, restricted herbicide use in wetland habitats (with the exception of rice production) apparently precludes selection for EHR [25]. For other species on the control list where no obvious factors account for the lack of EHR, it is likely that habitat, cropping system, or other agronomic factors influence the probability of exposure to herbicides, and thus, selection for EHR.

We also examined patterns between individuals with resistance to multiple herbicides in different functional groups (herbicides with different modes of action). We found that there is a significant increase in multiple resistance in the Amaranthaceae and Poaceae compared with the control list. The abundance of multiple resistance in these families is not surprising given that they are also a subset of the families that have high levels EHR. We also analyzed multiple resistance from the perspective of the herbicides involved. We found that the majority of herbicides have a similar proportion of species with resistance to that herbicide and species with multiple resistance. However, we found that 62% of species with glyphosate resistance have multiple resistance compared to
the approximately 30% among most other herbicide families. It is possible that the cause of increased multiple resistances that include glyphosate is due to the fact that glyphosate is a newer herbicide and is often applied to areas that already have resistance to other herbicides. Therefore, there are increased opportunities for evolution of resistance to glyphosate on a background of resistance to other herbicides.

A major mechanism for multiple herbicide resistance in plants is metabolism by cytochrome P450 monooxygenases \[6,26,27\]. As these enzymes can metabolize herbicides with different modes of action, their existence in weeds with EHR could strongly influence phylogenetic analysis. Unfortunately, while much is known about the role of these enzymes in insecticide resistance \[28\], little is known about their existence and role in weeds \[6,27\]. To date most of the reports of cytochrome P450 monooxygenases have been for grass weeds \[6\], but the limited number of species for which P450-based EHR is known precludes using this information in our analysis.

Figure 1. Phylogenetic tree showing the relationship among 52 plant families. EHR to ALS-inhibitors (left panel) and PS-II inhibitors (right panel) were mapped onto the tree using Mesquite version 2.74. Families with EHR to glyphosate are highlighted in yellow. Relevant plant lineages are labeled within the tree nodes. Lineages that show EHR are depicted in red. The red area within each pie chart indicates the relative support for different ancestor states.

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We expect that herbicide resistance will continue to evolve in regions with the strongest herbicide selection pressure, in particular, in major agricultural countries. For example, considering the case of glyphosate in 2010, the United States, Brazil, and Argentina were #1, #2, and #3 with respect to acreage of crops engineered for glyphosate resistance, collectively comprising more than 80% of the world’s acreage, which comprised 134 million ha in 2009 [29,30]. Presently, there are 112 known instances of EHR to glyphosate [5]; 86 of these were found in the United States, Brazil, and Argentina (more than 75% of the total). While we acknowledge that glyphosate is used for more than weed control in transgenic crops, the strong relationship between their acreage and EHR cannot be ignored.

Our analysis has immediate implications for developing countries that are accelerating or about to accelerate their use of herbicides. As reported by the National Research Council [1], without careful attention to herbicide resistance management (e.g., appropriate herbicide rotations in combination with non-chemical control methods), increased use of herbicides could bring about new instances of EHR [31]. Our results show that new instances of EHR are more likely to be found for annual weeds in the families that we identified as EHR prone. To take a specific example, the recent historic information on glyphosate use is easily available. In 2010, in terms of area planted to glyphosate resistant crops, Paraguay ranked fifth in the world with 2.6 million hectares of glyphosate resistant soybean [29]. That crop was first introduced to Paraguay in 2004 [29]. With six years of intense use, there are now two reports of glyphosate resistance in Paraguay, both forDigitaria insularis (sourgrass, once in 2006 and later in 2008). The situation is not trivial; the latter report indicates an increasing infestation on the order of hundreds of acres at over a dozen locations. We note that this species follows the phylogenetic trends our analysis has identified. Sourgrass is a member of the Poaceae, a family overrepresented for both EHR in general and glyphosate resistance in particular. With continued glyphosate use in Paraguay we would predict some of the following species to rapidly evolve resistance: Echinochloa colona and Eleusine indica, reported to have EHR to other herbicide groups in South America, and even more worrisome, Lolium spp. and Setaria halepense, reported to have glyphosate resistance elsewhere in South America. As herbicide resistant crops (transgenic or not) are increasingly deployed in developing countries, the problems of EHR will likely spread to those countries [32].

Pesticide resistance was first discovered in insects, then fungi, then plants [33], yet this type of analysis of phylogenetic patterns has never been done except at the gene level (e.g. [34,35], but see [22]). We have shown that weeds with EHR are far from a random sample of weedy plant species. Our analysis indicates that certain species in certain families, particularly those species with shorter life cycles, will rapidly evolve herbicide resistance under high selection pressure. Accordingly, local weed managers and scientists can anticipate this evolution and employ recommended herbicide resistance management techniques to delay evolution toward herbicide resistance and preserve the efficacy of a given herbicide [31]. Further analysis of other traits of EHR will refine such strategies.

**Supporting Information**

**Table S1 Complete list of species and life history traits used in this study.** (XLSX)

**Author Contributions**

Conceived and designed the experiments: NCE JSH. Performed the experiments: SRW SMH AMB KTP LCS. Analyzed the data: SRW KS. Contributed reagents/materials/analysis tools: IH. Wrote the paper: JSH NCE.
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