

## SPECIAL FEATURE: RESISTANCE EVOLUTION, FROM GENETIC MECHANISM TO ECOLOGICAL CONTEXT

## Editorial

## Resistance evolution, from genetic mechanism to ecological context

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Pesticide use by humans has induced strong selective pressures, reshaping evolutionary trajectories, ecological networks, and even influencing ecosystem dynamics. The evolution of pesticide resistance across weeds, insects, and fungi often leads to negative impacts on both human health and the economy while concomitantly providing excellent systems for studying the process of evolution. In fact, the study of pesticide resistance has been a feature of evolutionary biology since the Evolutionary Synthesis, with Dobzhansky noting in his book *The Genetics and Origins of Species* (1937) that cyanide resistance in the California red scale constituted the “best proof of the effectiveness of natural selection yet obtained”. Following the pioneering work of James Crow and others in the 1950s—which greatly expanded our knowledge of the genetics underlying adaptation—the study of pesticide resistance has shed light on a variety of topics, such as the repeatability of phenotypic evolution across the landscape, “hotspots” of evolution across the genome, and information on the number and type of genetic solutions that populations may employ to strong selection pressures.

Landscape level approaches have come to the forefront over the last 20 years of resistance evolution research, often taking advantage of the fact that replicated populations of the same species are exposed to the same pesticide. Further, the resistance evolution field is turning more attention to the ecological context within which resistance evolution occurs, probably stemming, at least in part, from an historical focus on fitness costs (Baucom, 2019; Cousens & Fournier-Level, 2018). This special feature, “Resistance evolution, from genetic mechanism to ecological context” in *Molecular Ecology* captures the current state of resistance evolution with contributions broadly addressing the question “What has the rapid evolution of

pesticide resistance taught us about genome dynamics and adaptation as well as the ecological context within which resistance evolution occurs?” Below, we contextualize the manuscripts in this special feature that provide insight into the state of the art investigations of resistance evolution across various species of insects, weeds and fungi.

## 1 | GENOMIC BASIS OF ADAPTATION: TARGET SITE AND NONTARGET SITE RESISTANCE MECHANISMS

Pesticides are designed to target particular genes in functional pathways of pest organisms. For example, quinone-outside inhibitor fungicides (QoIs) target the products of the fungal gene cytochrome b, a critical component of aerobic respiration. In plants, the herbicide glyphosate targets the protein product of EPSPS (5-enolpyruvylshikimate-3-phosphate synthase), a central gene in the shikimate acid pathway. One type of pesticide resistance—termed target site resistance—occurs when a mutation in the gene targeted by the pesticide alters the conformation of the protein, reducing or completely eliminating the ability of the pesticide to bind to the protein's active site (Délye et al., 2013). A number of studies have examined target site resistance by sequencing the target locus and assessing the various mutations associated with resistance. Given a priori information about how pesticides work, thus providing obvious genetic candidates to explore, along with the relative simplicity of sequencing a single gene, we currently have a somewhat comprehensive understanding of the type and number of genetic variants associated

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with target site resistance, especially in herbicide resistant weeds (Baucom, 2016; Tranel et al., 2021).

We still have much to learn about target-site resistance evolution, however, especially in the context of broader, genome-scale dynamics. In this special feature, Clarkson et al. (2021) move beyond investigating the dynamics of target-site resistance in isolation by explicitly examining the role of intragenic variation on resistance to pyrethroid, a class of insecticide that is used to control mosquito populations associated with malaria. Their investigation of whole-genome sequence data reveals “a tale of two alleles”: two widespread large-effect target-site resistance alleles within the voltage-gated sodium channel (VGSC) gene appear to be on different evolutionary trajectories. One allele, probably an early ancestral mutation, is associated (i.e., in strong positive linkage disequilibrium) with a subsequent explosion of 13 secondary nonsynonymous mutations, whereas the second allele is associated with fewer mutations. Further, most of these mutations are background-dependent, occurring nearly exclusively on distinct haplotypes—haplotypes that are associated with different signatures of selection despite harbouring the same focal resistance allele, implying important compensatory or enhancing allelic interactions for resistance evolution.

A major contribution to our understanding of the predictability of evolution stems from work examining the repeatability of target site changes that confer pesticide resistance across insect and weed species, respectively (Martin & Orgogozo, 2013). However, whether or not parallel genetic changes lead to resistance among fungal plant pathogens has yet to be succinctly summarized. In this issue, Hawkins and Fraaije (2021) investigate the extent of parallel evolution of individual mutations in target genes among species of fungal pathogens. Focusing on mutations associated with four classes of fungicide, they show that the target-genes vary substantially in the diversity of mutations detected. For two fungicide classes (Qols and MBCs) the same mutations are observed repeatedly across species. In contrast, a greater diversity of resistance mutations was uncovered within genes targeted by azole and SDHI fungicides, providing less evidence for extreme parallelism across species compared to Qols and MBCs.

Another form of target-site resistance is from gene amplification, where increased copy number of the target locus leads to more functional protein and subsequent resistance. In a handful of weeds, an increase in the copy number of the EPSPS locus leads to high glyphosate resistance (reviewed in Gaines et al., 2019); while the underlying mechanism of this copy number increase has been described (Koo et al., 2018), we understand relatively little about the long-term maintenance of copy number variation (CNV) and how gene amplification may influence interactions with other loci. Yakimowski et al. (2021) quantified patterns of variation of target gene copy number and resistance phenotypes within and among populations—the “natural history” of a resistance CNV—to provide insight into the evolution of glyphosate resistance in the agricultural weed *Amaranthus palmeri* in the eastern United States. They detected a steep increase in phenotypic glyphosate resistance at a threshold value of ~15 gene copies, but also found that populations with the highest mean resistance contained some low copy number

individuals (albeit at low-frequency). From 15 to 160 gene copies the level of resistance changed very little; however, the proportion of low-resistance phenotypes gradually decreased in populations with increased copy number, suggesting that dosage of the target gene with increasing copy number might compensate for negative interactions with other loci. Potential positive interactions with other genes were also observed in populations from Georgia. Overall, target gene copy number variation explained a high proportion (~57%) of variation in phenotypic resistance among populations.

In another set of contributions to this special feature, Ravet et al. (2021) and Gaines et al. (2021) take landscape-level approaches to examine EPSPS copy number variation and its association with resistance in kochia (*Bassia scoparia*) and *Amaranthus palmeri*, respectively. In glyphosate resistant kochia, researchers paired an investigation of population structure via SSR markers with qPCR markers designed to examine the EPSPS repeat genotype across western lineages. They found low population structure in this species but evidence for independent evolutionary origins of the EPSPS duplication leading to resistance. In the landscape level assessment of *A. palmeri*, researchers found copy number variation of this target gene—like that found across lineages from the eastern United States by Yakimowski et al. (2021)—present in populations from Brazil and Uruguay, indicating that copy number variation related to resistance is found broadly across the landscape in this species. Interestingly, however, resistance in Argentinian lineages of *A. palmeri* was due in large part to nontarget site resistance mechanisms—that is, resistance mechanisms that do not involve the target site, such as altered translocation or detoxification of the pesticide, among others—rather than elevated copy number of the EPSPS locus. These results show both genomic flexibility in solving the problem of herbicide exposure and the independent, novel evolution of resistance across geography.

Thus, in addition to target site resistance mechanisms, organisms can also evolve resistance through nontarget site mechanisms. Nontarget site resistance mechanisms, which are often thought to be due to polygenic variation, can both confer resistance as well as potentially supplement target-site effects. While both target site and nontarget site mechanisms have previously been uncovered within the same herbicide resistant weed species (as in Gaines et al., 2020), the relative contribution of either type of mechanism has yet to be clearly delineated in any weed species. Using another glyphosate resistant *Amaranthus* species, *A. tuberculatus* (common waterhemp), Kreiner et al. (2021) uncovered the cryptic contribution of genome-wide alleles to glyphosate resistance. On the genomic background of agricultural populations harbouring high frequencies of target-site resistance mechanisms (Kreiner et al., 2019), the authors illustrated a near-equal importance of nontarget and target-site mechanisms. Further, they uncovered hundreds of alleles associated with nontarget site resistance that show not only evidence of recent strong selection from herbicides but a classic trade-off between effect size and allele frequency that implicates pleiotropy as a key constraint to the evolution of herbicide resistance.

As our understanding of the genetic architecture of pesticide resistance and governing selective processes deepens, a key question

will be how consistently such alleles are involved across geographic scales. This question is addressed by Hartmann et al. (2020), who investigated the architecture of azole fungicide in a key wheat pathogen, *Zymoseptoria tritici* across three continents. They uncovered a suite of azole resistance-related loci across the genome including a novel large-effect gene, DHHC palmitoyl transferase. Along with key alleles conferring resistance to three other chemical classes of fungicides, the authors find evidence that the genomic architecture of fungicide resistance is largely distinct across continents, with the exception of large-effect genes that act as hotspots for convergence. Overall, this collection of work characterizing the genetic architecture of pesticide resistance uncovers remarkable complexity in monogenic and polygenic contributions and the processes that govern their assemblage across genomes, from background- and population-specific constraints to the potential for pleiotropic tradeoffs.

## 2 | ECOLOGICAL CONTEXT AND THE INFLUENCE OF INTERACTIONS ON THE EVOLUTION OF RESISTANCE

Pesticide use can have impacts far beyond their intended target organism by influencing the ecology and evolution of organisms with which the target species interact. Contributions in this issue explore the ecological context of resistance evolution by assessing how pesticide application may affect interactions between target and non-target organisms, which may influence downstream ecoevolutionary feedback dynamics.

Iriart et al. (2021) set the stage for our understanding of the ecological context of resistance evolution by reviewing the role of herbicides in driving the ecology and evolution of plants and plant-associates (e.g., pollinators, soil microbes, herbivores, and parasitoids) living in communities at the agroecological interface. They synthesize what is known about how herbicides can alter plant phenotypes from plastic or genetic changes and how plant-associates may be directly or indirectly (via interactions) affected by herbicides. Building off this knowledge, they demonstrate that herbicides can induce sufficiently rapid change in plants and plant-associates to alter both evolution and ecological dynamics over the same timescales, thus producing ecoevolutionary feedbacks. From these insights, they provide suggestions for future research into herbicide catalysed ecoevolutionary dynamics, with the goal of deciphering the effects of herbicides on plant and plant-associates' traits, on species interactions, and on the composition of the broader ecological community.

Herbicide application may alter ecoevolutionary dynamics by selecting for traits that are correlated to resistance, such as earlier flowering time or altered mating patterns (among other changes) thereby potentially modifying mutualistic interactions between plants and their associates. For example, glyphosate resistant populations of the common morning glory (*Ipomoea purpurea*) exhibit higher selfing rates compared to susceptible populations (Kuester et al., 2017),

perhaps due to reproductive assurance associated with being both highly selfing and herbicide resistant. However, an association between the mating system and resistance would not be expected to be maintained over time if the resistant, selfing types exhibited inbreeding depression. In this issue, Van Etten et al. (2021) combined growth chamber and field studies with transcriptome surveys to ask whether genetic lines of *Ipomoea purpurea* selected for increased glyphosate resistance exhibited signs of inbreeding depression (i.e., poorer performance of inbred versus outcross progeny) compared to both nonselected control lines and lines selected for increased susceptibility. Interestingly, they found that while plants from non-selected control lines and susceptible lines exhibited evidence of inbreeding depression, plants from resistant lines provided no evidence for inbreeding depression in most characters. Rather, in the presence of herbicide, resistant lines tended to show outbreeding depression: seeds from resistant lines that were produced via selfing germinated more and grew to be larger plants than those from resistant lines that were produced from outcrossing. Additionally, the authors showed that the expression of genes within the transcriptome mirrored the phenotypic patterns—resistant, inbred plants showed higher expression of genes involved in translation and DNA replication compared to resistant, outcrossed progeny in the presence of glyphosate. Thus, in this case study, continued resistance evolution would support higher self-fertilization and decreased outcrossing. In this way, the maintenance of plant-pollinator interactions could be negatively altered over time in herbicide-exposed populations of *I. purpurea*.

As with herbicides, the evolution of resistance to insecticides has the potential to alter ecological interactions in crop ecosystems, as shown by Paddock et al. (2021). Their study investigated whether the microbial communities differed between herbivorous western corn rootworms (a widespread agricultural pest) that were susceptible or had evolved resistance to the insecticide *Bacillus thuringiensis* (Bt) produced by genetically-modified maize. Their results supported different enteric microbiomes between resistant and susceptible western corn rootworm in that resistant individuals had less rich and diverse bacterial communities. Additionally, western corn rootworm digesting the insecticide caused a severe shift towards more simplified bacterial communities in susceptible western corn rootworm, but not resistant western corn rootworm, suggesting an effect of host-microbial interactions in the evolution of resistance to Bt. Together, these results contribute to our budding understanding of the role that ecology can play in resistance evolution to modern stressors, further perpetuating ecoevolutionary feedbacks and dynamics in natural communities.

## 3 | CONCLUSION

The manuscripts in this special feature cover both a wide conceptual range and a broad group of organisms. The common thread between studies is that they each investigate applied systems to broaden our understanding of either the ecological or genetic factors that

influence adaptation. While many researchers are probably drawn to the study of pesticide resistance evolution for its perceived simplicity—for example, there is a known agent of selection as well as (typically) a known biochemical and often simple genetic target of the pesticide—contributions in this special feature show that both the genetic and ecological context of resistance evolution remains to be deepened in important ways. For example, the population genomic approaches used in these manuscripts provide increased resolution into the number and interactions of alleles involved in resistance, timescale-dependent signatures of selection, heterogeneity among populations in the control of resistance, and gene expression differences associated with both resistance and the mating system. Additionally, the work in this special feature highlights that resistance evolution does not occur in a vacuum; while many examinations of resistance focus on the evolutionary trajectory of a single species exposed to pesticide, the overall adaptation and persistence of a population will concomitantly be influenced by, and likewise influence, other community members such as pollinators, herbivores and the microbial community. Ultimately, deepening the context of resistance evolution by both broadening our genetic toolkits and by assessing community dynamics will allow us to better understand how genetics and ecology are linked and how such linkages can then influence larger-scale ecosystem dynamics.

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