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Attracting pollinators vs escaping herbivores: eco-evolutionary dynamics of plants confronted with an ecological trade-off

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Abstract

Many plant traits are subject to an ecological trade-off between attracting pollinators and escaping herbivores. The interplay of both plant-animal interaction types determines their evolution. As most studies focus on either pollination or herbivory, how they jointly affect the eco-evolutionary dynamics of plant-animal communities is often left unknown. Within a plant-pollinator-herbivore community where interaction strengths depend on trait matching, we consider the evolution of a plant trait involved in both plant-animal interactions. Using adaptive dynamics, we uncover when stabilizing, runaway (i.e. directional) or disruptive selection emerges and its consequences for multispecies coexistence. We find that strong pollination relative to herbivory favors stabilizing selection and coexistence. Strong herbivory relative to pollination fosters runaway selection and threatens coexistence. Importantly, given balanced interactions, joint effects may lead to disruptive selection, allowing the emergence of plant dimorphism. The strength of the ecological trade-off largely explains the occurrence of these contrasting eco-evolutionary dynamics. In particular, plant diversification requires strong trade-offs, with the strongest trade-offs allowing long-term polymorphism. We discuss how our results relate to various empirical cases where the interplay of pollination and herbivory maintains plant polymorphism. Beyond maintenance, our work suggests that it might also have fueled the diversification process itself.

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Introduction

Flowering plants (i.e. angiosperms) are the most diverse and successful plant clade in terrestrial ecosystems, representing almost 90% of the species described (Hernández-Hernández & Wiens, 2020). Most of them rely on animals for pollination (Ollerton et al., 2011). Because animal pollination can favor reproductive isolation, it has been proposed early as an important diversity driver among angiosperms (Grant, 1949). Plant-herbivore interactions have, however, also been identified as potentially fostering diversification due to the ensuing evolutionary arms race between interacting antagonists (Ehrlich & Raven, 1964). These two non-mutually exclusive hypotheses are supported by several phylogenetic investigations. Plant-pollinator coevolution explains higher diversification rates within angiosperms (Hodges & Arnold, 1995; Sargent, 2004; Hernández-Hernández & Wiens, 2020), but plant-herbivore coevolution also spurs plant diversification as a result of defense-counterdefense innovations (Farrell et al., 1991; Becerra et al., 2009). Plant phylogenies reveal that plant-pollinator and plant-herbivore coevolution are often inextricably intertwined (e.g. Armbruster 1997), which advocates for an integrative framework accounting for both interaction types (Fontaine et al., 2011). From an ecological perspective, the positive or negative outcome of many interactions is often context-dependent (Chamberlain et al., 2014), varying for instance with the phenotype of interacting individuals (e.g. larvae vs. adult of nursery pollinators, Hahn and Brühl 2016). This dynamic nature of interactions along a mutualisticantagonistic continuum (Thompson, 1988; Gómez et al., 2023) further highlights the relevance of such an integrative framework.

A large number of plant traits are notably under conflicting selection due to the interplay between pollination and herbivory (Strauss & Irwin, 2004), floral traits in particular (Strauss & Whittall, 2006). Conflicting selection pressures can emerge from several mechanisms acting either in isolation or synergistically. Herbivory-induced changes in plant chemistry potentially reduce pollinator visitations (e.g. Kessler et al. 2011). Many plant traits acting on the plant visibility (size, phenology, floral display, volatile compounds) may increase herbivory (apparency hypothesis, Feeny 1976) while also attracting allies (e.g. Brody 2008). Genetic correlation can also exist between two plant traits, each involved in one plant-animal interaction (Strauss et al. 2004). A decisive consequence is that the selection pressures in the absence of either one animal species would be modified in the presence of both animal groups (Ramos & Schiestl, 2019). Conflicting selection is very often due to shared preferences for plant phenotypes between pollinators and herbivores, a pattern that is largely widespread in nature (Strauss & Whittall, 2006) and that promotes the stable coexistence of the community (Yacine & Loeuille, 2022). This preference pattern indicates that plant species are subject to an ecological trade-off between attracting pollinators and escaping herbivores (Strauss et al., 2002). In other words, an increase in the strength of pollination (e.g. investment in attraction) comes at the cost of an increase in the strength of herbivory, while a decrease in the strength of herbivory (e.g. production of defenses) comes at the cost of a decrease in the strength of pollination. This trade-off has interestingly been shown to support the maintenance of a flower-color polymorphism in the wild radish Raphanus sativus (Irwin et al., 2003), which suggests that the interplay between pollination and herbivory could also lead to disruptive selection and promote diversification.

Conflicting selection arises because pollination and herbivory exert opposite pressures on plant traits (Thompson, 2009), with contrasting implications in terms of diversification potential (Yoder & Nuismer, 2010). These differences are especially prominent under the assumption that interaction strengths increase with the matching between plant and animal phenotypes (i.e. trait-matching). Such an assumption seems reasonable as it has been reported to apply in various instances, including flower color (Irwin et al., 2003) or flower display (Galen & Cuba, 2001) matching animal preferences, chemical volatiles (Theis et al., 2014) matching animal detection abilities, or plant phenology matching that of animals (Brody, 2008). As fitness increases with the strength of a mutualistic interaction, plants are selected to better match their pollinator phenotype and vice versa (e.g. spur length and pollinator tongue, Whittall & Hodges 2007) so that stabilizing selection is

expected (e.g. Kiester et al. 1984). Pollination-induced stabilizing selection has been empirically characterized several times (e.g. Parachnowitsch and Kessler 2010; Sahli and Conner 2011; De Jager and Peakall 2019). It disfavors extreme phenotypes, thus reducing the potential for disruptive selection (Kopp & Gavrilets, 2006; Yoder & Nuismer, 2010; Maliet et al., 2020). Conversely, predation reduces prey fitness so that plants are selected to escape herbivore preferences (e.g. chemical defenses and herbivore tolerance, Becerra et al. 2009). Such arms race dynamics (Ehrlich & Raven, 1964; Dawkins & Krebs, 1979) expose the plant species to runaway selection, as observed in empirical systems (e.g. Mauricio and Rausher 1997; Becerra et al. 2009; Griese et al. 2021). Because apparent competition (Holt, 1977) imposes a cost on phenotype matching between conspecific plants, herbivory also favors increased plant phenotypic divergence thereby enhancing disruptive selection (Kopp & Gavrilets, 2006; Yoder & Nuismer, 2010; Maliet et al., 2020). Note finally that conflicting selection may have far-reaching ecological implications. The relative interaction strength of pollination vs. herbivory has been shown to determine coexistence within plant-pollinator-herbivore communities (Mougi & Kondoh, 2014; Sauve, Fontaine, et al., 2016; Yacine & Loeuille, 2022). Because interaction strengths are the result of plant-animal coevolution, the evolution of plant traits under conflicting selection may have important consequences for multispecies coexistence.

In the present paper, our goals are to determine how the interplay of pollination and herbivory drives the evolution of plant traits involved in both interactions, and how, in turn, such an evolution affects the maintenance of coexistence. We model a community – a plant species interacting simultaneously with a pollinator and a herbivore species - in which plant-animal interactions depend on a plant trait involved in both interactions. Our previous purely ecological investigation of a similar three-species system has shown that a balance between the strength of plant-animal interactions makes the three-species coexistence more likely (Yacine & Loeuille, 2022). Ignoring all evolutionary aspects, this previous study utterly focused on the ecological dynamics and their stability in relationship with plant-animal interaction strengths considered as independently varying parameters. In contrast, pollination and herbivory are here coupled by a focal plant trait whose phenotype determines the strength of both interactions (see Strauss & Whittall (2006) for examples). Each interaction increases in strength with the matching between plant phenotype and animal preferences (or more generally and henceforth, animal phenotypes). Under this trait-matching setting, we study the eco-evolutionary dynamics resulting from the evolution of the plant phenotype using the adaptive dynamics framework (Metz et al., 1992; Dieckmann & Law, 1996).

We are particularly interested in understanding how these dynamics depend on an ecological trade-off to which the plant might be subject. An ecological trade-off is here defined as a positive covariation between the strengths of pollination and herbivory, i.e. stronger pollination (resp. weaker herbivory) comes at the cost of stronger herbivory (resp. weaker pollination). We first show that under our trait-matching assumption, variations in plant phenotype intrinsically entail an ecological trade-off whose strength depends on the dissimilarity between animal phenotypes (details in Methods, Fig. 1). Manipulating animal phenotype dissimilarity, we investigate how the strength of this ecological trade-off affects the type of selection on the plant trait and the maintenance of coexistence. When animal phenotypes are highly dissimilar (weak trade-off), interaction strengths are intrinsically unbalanced making the plant essentially interacting with one animal species so that we expect selection to be stabilizing close to the pollinator phenotype, and runaway close to the herbivore phenotype. Coexistence should then be frequently disrupted as stable coexistence requires the plant-animal interaction strengths to be of similar magnitudes (Yacine & Loeuille, 2022). When animal phenotypes are fairly similar (strong trade-off), the balance between interaction strengths should, in contrast, favor the maintenance of coexistence. The potential for disruptive selection to occur in such instances, and the maintenance of the ensuing polymorphism, is one of the key questions of the present study.

Model & Methods

Model

Population dynamics

We model the dynamics of a plant-pollinator-herbivore module (P, M, H respectively) using a Lotka-Volterra framework:

(1) $\frac{dP}{dt} = P(r_p - c_p P + a_{pm} M - a_{ph} H)$ $\frac{dM}{dt} = M(r_m - c_m M + e_m a_{pm} P)$ $\frac{dH}{dt} = H(r_h - c_h H + e_h a_{ph} P)$

Plants are assumed to have a positive intrinsic growth rate ($r_p > 0$), while both pollinator ($r_m < 0$) and herbivore growth rates ($r_h < 0$) are assumed negative. As in previous models (e.g. Sauve et al. 2014), we thus assume the plant-animal interaction to be obligate for the animals and facultative for the plant. All interacting species undergo negative (quadratic) density-dependence. These terms are here interpreted as intraspecific competition. The plant competition rate (c_p) would notably encompass the competition for space, water or nutrients (Craine & Dybzinski, 2013). Animal competition rates (c_m, c_h) would essentially coincide with interference (e.g. Thébault and Fontaine 2010). These terms might however be more generally considered as any negative regulation not explicitly accounted for in our model (e.g. other predators or resources, diseases....). The per-capita strengths of interspecific interactions are given by a_{pm} for pollination and a_{ph} for herbivory. e_m and e_h are the conversion efficiencies from plants to animals. Table 1 recapitulates model variables and parameters. Our formulation of population dynamics implies an indirect (i.e. plant-mediated) interaction between animal densities, but no direct interaction as would for instance be the case when an animal behaves as both pollinator and herbivore (e.g. nursery pollinators).

Without evolution, stable three-species coexistence occurs when the strengths of pollination (a_{pm}) and herbivory (a_{ph}) are balanced (Yacine & Loeuille, 2022). When pollination is much stronger than herbivory, stability may be lost in the form of unbounded population growth (e.g. Fig. B1, Fig. 5b-d). In this parameter region $(a_{pm} \gg a_{ph})$, our model thus fails at producing biologically realistic dynamics, which indicates that other ignored processes are prominent in such instances (e.g. saturating parameters, wider community context; discussed extensively in Yacine & Loeuille, 2022). One of our main previous results, however, is that the plant-herbivore interaction largely reduces the size of this parameter region, i.e. herbivory often stabilizes unbounded dynamics in plant-insect systems (see Fig. B1.A in appendix B). In the present work, we investigate how the relative strength of pollination vs. herbivory influences plant evolution, and how such evolution, in turn, affects the maintenance of multispecies coexistence. Evolution is characterized in the parameter region over which stable three-species coexistence is obtained (analytical expression of equilibrium (P^* , M^* , H^*) given in Appendix A.I, Supporting Information), as well as in regions of unbounded growth (see Methods).

Plant-animal interactions depend on trait matching

We assume plant-animal interactions to intensify as the matching between plant (t_p) and animal phenotypes – pollinator (t_m) or herbivore (t_h) – increases (equations 2). Interactions are maximal when traits perfectly match (i.e. $|t_p - t_m| = 0$, resp. $|t_p - t_h| = 0$), and weaken when trait-matching is reduced (i.e. $|t_p - t_m| \nearrow$, resp. $|t_p - t_h| \nearrow$). Examples include the color of plant flowers (t_p) and

associated animal preferences (t_m, t_h) (e.g. Irwin *et al.* 2003) or phenological traits (e.g. Brody 2008) such as the date of flowering (t_p) and of animal activity (t_m, t_h) .

(2)
$$a_{pm}(t_p) = \frac{a_{pm0}}{\sqrt{2\pi\sigma_{Pol}^2}} \exp\left[-\frac{\left(t_p - t_m\right)^2}{2\sigma_{Pol}^2}\right]$$
$$a_{ph}(t_p) = \frac{a_{ph0}}{\sqrt{2\pi\sigma_{Her}^2}} \exp\left[-\frac{\left(t_p - t_h\right)^2}{2\sigma_{Her}^2}\right]$$

As σ_{Pol} controls how quickly the strength of pollination decreases with plant-pollinator phenotype dissimilarity, it corresponds to the pollination-niche width, which depends on the generalism of both species. Likewise, σ_{Her} embodies the herbivory-niche width. a_{pm0} and a_{ph0} correspond to basal interaction rates. See table 1 for details.

Variables and parameters **Biological definition** Value Dimension Р Plant biomass density $mass.length^{-2}$ $mass.length^{-2}$ М Pollinator biomass density Variables Η Herbivore biomass density mass.length⁻² Dimensionless Plant phenotype t_p 0 Dimensionless t_m Pollinator phenotype Animal phenotypes Herbivore phenotype [0,3] Dimensionless th Animal phenotype dissimilarity [0,3] Dimensionless $t_h - t_m$ Proxy for the strength of the ecological trade-off (see Methods) Interspecific Basal pollination rate (per capita) [3,9] $(M.L^{-2})^{-1}.t^{-1}$ interaction a_{pm0} parameters $(M.L^{-2})^{-1}.t^{-1}$ Basal herbivory rate (per capita) [3,9] a_{ph0} Pollination niche width [1.5,3] Dimensionless σ_{Pol} Herbivory niche width Dimensionless [1.5,3] σ_{Hei} $time^{-1}$ Plant intrinsic growth rate 10 r_p $time^{-1}$ Pollinator intrinsic growth rate $^{-1}$ r_m Herbivore intrinsic growth rate -4(-1,-1) $time^{-1}$ r_h Other Plant intra-specific competition rate 0.6 (0.6, 1.7) $[(mass.length^{-2}).time]^{-1}$ c_p ecological [(mass.length⁻²).time]⁻¹ Pollinator intra-specific competition rate 0.5 (0.4, 0.7) c_m parameters Herbivore intra-specific competition rate 0.4 $(M.L^{-2})^{-1}.t^{-1}$ C_h Dimensionless Plant to pollinator conversion efficiency 0.2 e_m 0.3 (0.2, 0.3) Dimensionless Plant to herbivore conversion efficiency e_h Mutation probability (per unit of time and plant 2.10^{-7} $[(mass.length^{-2}).time]^{-1}$ μ biomass density) Numerical Mutation amplitude (standard deviation) 0.02 Dimensionless σ simulations Extinction threshold 2.10^{-6} $mass.length^{-2}$ ε

 Table 1 - List of all model parameters and variables with their biological definitions, values and dimensions.

column "Value", "*other ecological parameters*": in parenthesis are provided the values for the robustness ecological parameter sets when they differ from that of our main ecological parameter set (see section Methods/Workflow).

Methods

Definition and emergence of an ecological trade-off

An ecological trade-off between attracting pollinators and escaping herbivory is defined as a positive covariation between the strengths of pollination and herbivory, i.e. both increasing or decreasing. As our focal plant trait is involved in both plant-animal interactions (equations 2), its variation affects both interaction strengths (e.g. Fig. 1b). Because of our trait-matching hypothesis, the induced covariation between the strengths of pollination and herbivory is positive when the plant phenotype is outside the phenotypic interval $[t_m, t_h]$. Variation in plant phenotype there entails an ecological trade-off (Fig. 1b, solid line). Within the interval $[t_m, t_h]$, there is no ecological trade-off (Fig. 1b, dotted line).

We consider the dissimilarity between animal phenotypes $(t_h - t_m)$ to be a proxy for the strength of the ecological trade-off that emerges within our framework. There are two reasons for that. First, as animal phenotype similarity increases (Fig. 1, a vs. b vs. c), the phenotypic region over which plant phenotype variation induces an ecological trade-off increases in size. Second, over this region, in the case of very dissimilar animal phenotypes (i.e. $t_m \ll t_h$, Fig. 1a), plant phenotype variations affect much more one interaction than the other, depending on the closest animal phenotype. In contrast, over the same region but in the case of similar phenotypes (i.e. $t_m \approx t_h$, Fig. 1c), any variation of the plant phenotype has an effect of comparable magnitude on the strength of both interactions. All in all, the ecological trade-off between attracting pollinators and escaping herbivores gets stronger as animal dissimilarity decreases (Fig. 1).





Figure 1 - Animal phenotype dissimilarity determines the strength of the ecological trade-off. Each graph shows the covariation between the strength of plant-animal interactions resulting from varying plant phenotype (t_p varies from -10 to 10 in the direction indicated by the arrows, the orange (resp. red) point indicates when $t_p = t_m$ (resp. $t_p = t_h$). a. b. c. Increasing strength of ecological trade-off with increasing similarity between animal phenotypes (i.e. decreasing dissimilarity). Parameter set: ecological ($r_p = 10, r_m = -1, r_h = -4, c_p = 0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h = 0.3$); interspecific ($t_m = 0, a_{pm0} = a_{ph0} = 9, \sigma_{Pol} = \sigma_{Her} = 2$).

Adaptive dynamics and type of selection

Within a monomorphic plant population with phenotype t_p (resident), we investigate whether a mutant with a new phenotype t_p' can invade. Invasion fitness $w(t_p', t_p)$ is computed as the per capita growth rate of that mutant, when rare and in the environment (P^*, M^*, H^*) set by the resident population (Metz *et al.* 1992, see appendix A.II). When a mutant invades (i.e. $w(t_p', t_p) > 0$), it replaces the resident population, thus becoming the new resident. The sequence of trait

substitutions defines the long-term evolutionary dynamics. Assuming small and rare mutational steps, the variation of phenotype t_p is proportional to the selection gradient (equation 3), i.e. the derivative of invasion fitness with respect to the mutant's trait (Dieckmann & Law, 1996). Evolutionary singularities t_p^* correspond to phenotypes that nullify the selection gradient.

The type of selection - stabilizing, disruptive, or runaway - acting on the plant trait depends on (1) the properties of the evolutionary singularities, and (2) the position of the plant phenotype relative to these singularities. Two independent properties - convergence and invasibility characterize evolutionary singularities (criteria in appendix A.II). Convergence indicates whether the trait evolves toward the singularity in its vicinity. Two types of singularities are convergent: the non-invasible continuously stable strategy (CSS, Eshel, 1983) and the invasible branching point (Geritz et al. 1997). Invasibility specifies whether the singularity may be invaded by nearby mutants (i.e. ESS, Maynard Smith & Price 1973). CSS phenotypes correspond to cases of stabilizing selection (e.g. Fig. 2B.a). Selection is thus considered stabilizing in the basin of attraction of a CSS. Plant-pollinator-herbivore coexistence is notably maintained if a CSS is reached while the three species coexist. We also describe directional selection towards and within a phenotypic region in which our model produces unbounded growth as stabilizing selection (in terms of purely evolutionary dynamics, not ecological). This choice was motivated by both mathematical and biological coherence (details in appendix B.II.1). Unbounded growth regions are notably attractive in terms of evolution (i.e. convergence, note that classical tools of adaptive dynamics (e.g. selective gradient) do not apply within these regions but only in their vicinity). Our choice also preserves the association between stabilizing selection and coexistence maintenance as areas of unbounded growth are regions of phenotypic space in which our model fails to produce realistic dynamics, but in which, from a biological point of view, coexistence should be maintained (notion of "permanent coexistence", Hutson and Law 1985, discussed in Yacine and Loeuille 2022). In contrast to a CSS, a branching point yields the emergence of plant dimorphism due to disruptive selection (e.g. Fig. 2B.d). Accordingly, selection is considered disruptive in the basin of attraction of a branching point. Finally, phenotypes that are not in the basin of attraction of a convergent singularity are under runaway selection. This is possible in the presence of a non-convergent singularity, i.e. a repellor. For the sake of simplicity, we will refer to this set of phenotypes as the basin of attraction associated with runaway selection. Runaway selection should always disrupt plant-pollinator-herbivore coexistence (e.g. Fig. 2B.b). We illustrate how the proportion of phenotypic space under each type of selection is calculated in Appendix B.II.1.

Numerical simulations of community eco-evolutionary dynamics

This framework is completed by numerical simulations starting from a monomorphic plant population (t_p) interacting with a pollinator (t_m) and a herbivore (t_h) population. Mutations are randomly generated following a Poisson process characterized by a mutation probability per unit of time and plant biomass density $\mu = 2.10^{-7}$. Proportionally to phenotype abundances, a parent phenotype is randomly chosen at each mutation event. The mutant phenotype is drawn from a Gaussian distribution centered around the parent phenotype with standard deviation $\sigma = 0.02$. Its initial density is set to a small value ε , taken from the parent population. Symmetrically, populations falling below ε are removed from the system (extinction).

Workflow

Animal phenotypes are fixed parameters (without loss of generality: $t_m = 0, t_h \ge t_m$) while we study the evolution of the plant phenotype (t_p) . Parameters directly affecting plant-animal interactions – i.e. the interspecific parameter set $(t_h - t_m, a_{pm0}, a_{ph0}, \sigma_{Pol}, \sigma_{Her})$ – are at the core of our investigation. Thanks to the model simplicity, several analytical results uncovering various aspects of evolutionary dynamics (e.g. equation 3) are possible. For aspects that cannot be mathematically investigated, we provide numerical resolutions characterizing the variation of eco-evolutionary dynamics along the gradient of trade-off intensity (E3-diagrams, e.g. Fig. 2A). To broaden our understanding of possible evolutionary dynamics, we undertake Monte Carlo experiments (details in appendix B.II). We let interspecific parameters vary independently within

their interval ranges (table 1), which were chosen to explore a wide range of pollination and herbivory intensities (Appendix B.I). Remaining parameters (r_p , r_m , r_h , c_p , c_m , c_h , e_m , e_h) are fixed, but the experiments were conducted for three different sets to assess the robustness of our results: a main ecological parameter set allowing a large range of plant-animal interaction strengths to be compatible with stable coexistence, a second one characterized by symmetrical parameter values for animals, and a third one in which unbounded population growth is made impossible (details in Appendix B.I). We highlight here that our results are essentially robust, as detailed hereinafter. The different Monte Carlo experiments are presented directly in the Result section when needed. Our approach is finally complemented by numerical simulations of the community eco-evolutionary dynamics. Such simulations are used to investigate the emergence of plant dimorphism in the case of disruptive selection.

Results

A typical example of eco-evolutionary dynamics

Eco-evolutionary dynamics qualitatively depend on the strength of the ecological trade-off (Fig. 2). These dynamics are characterized by stabilizing or disruptive selection when trade-offs are strong. They then allow the coexistence of all species. Such a coexistence is, however, often lost in the case of weak trade-offs, where runaway selection is much more common. As developed later on (Fig. 4A), these variations are more general than the specific example we introduce here (Fig. 2A). On Fig. 2A, the plant phenotype always converges towards a CSS phenotype under very strong trade-offs (region I) so that selection is stabilizing. As the trade-off weakens (region II), both stabilizing (Fig. 2B.a) and disruptive selection (Fig. 2B.e) become possible depending on the initial plant phenotype. Stabilizing (resp. disruptive) selection is observed over the basin of attraction of the CSS (black line) (resp. branching point, blue line). These basins of attraction (see arrows) are separated by the repellor (grey line). At weaker trade-offs (region III), the basin of attraction of the CSS covers the whole coexistence area (white) so that selection is always stabilizing (Fig. 2B.b). In region IV, eco-evolutionary dynamics are gualitatively similar to those of region II. For even weaker trade-offs (region V), runaway selection is observed whenever the initial plant phenotype is above the repellor (Fig. 2B.c). Associated eco-evolutionary dynamics lead first to the extinction of pollinators, then of herbivores. Below the repellor, selection stabilizes at the CSS phenotype. For even weaker trade-offs (region VI), the CSS exits the coexistence area so that only runaway selection remains possible. It provokes the extinction of both animal species when starting above the repellor, and that of herbivores when starting below the repellor. The extinction of herbivores then leads to a perfectly matched plant-pollinator community (Fig. 2B.d).

The ecological dynamics induced by each type of selection qualitatively differ (Fig. 2B). Coexistence is always maintained in the case of stabilizing selection. It is always disrupted in the case of runaway selection due to the weakening of one or both plant-animal interactions. Disruptive selection allows the emergence of plant dimorphism. The two plant phenotypes diverge, one leading to stronger plant-animal interactions, the other to weaker plant-animal interactions. Dimorphism can be temporary (Fig. 2B.e) or permanently maintained (Fig. 2B.f). An example of eco-evolutionary landscape associated with permanent dimorphism is given in Fig. S1 (Supporting Information), while the landscape presented in Fig. 2A typically leads to temporary branchings. In addition to trade-off intensity, other interspecific parameters can thus be responsible for qualitative changes in eco-evolutionary dynamics. Most of such (qualitative) variation is covered by the simulations in Fig. 2B.



Figure 2 - Typical examples of eco-evolutionary dynamics with their associated ecoevolutionary landscape. A. Eco-evolutionary landscape. The points labelled by letters indicate the initial conditions of the simulations presented in B. Ecological parameter set: ($r_p = 10, r_m = -1, r_h = -4, c_p = 0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h =$ 0.3). Interspecific parameter set ($a_{pm0} = 5, a_{ph0} = 7, \sigma_{Pol} = 3, \sigma_{Her} = 2.8$) B. Simulated eco-evolutionary dynamics associated with each type of selection. a-b. Stabilizing selection enables the maintenance of coexistence ($t_h = 0.2 \& 1$ respectively). c-d. Runaway selection leads to coexistence loss ($t_h = 2.8 \& 2.2$ respectively). e-f. Disruptive selection allows the emergence of plant dimorphism ($t_h = 0.2 \& 0.1$ respectively). Note that for f, the interspecific parameter set is modified ($a_{pm0} = 5, a_{ph0} = 9, \sigma_{Pol} = 1.7, \sigma_{Her} = 2$). The associated landscape is given in Fig. S1. In B, pictograms depict the community composition, with arrow thickness correlating to interaction strengths.

Opposite effects of pollination vs herbivory on the (local) selection gradient

The selection gradient (equation 3) shows that plant evolution is under two contrasting selective forces (i.e. opposite sign).

(3)
$$\frac{dt_p}{dt} \propto \frac{Selection}{gradient} = -\frac{\underbrace{a_{pm}(t_p)M^*}}{\underbrace{\sigma_{Pol}^2}_{pollination}}(t_p - t_m) + \underbrace{\underbrace{a_{ph}(t_p)H^*}}_{\underbrace{\sigma_{Her}^2}_{herbivory}}(t_p - t_h)$$

Pollination selects for plant phenotypes that are closer to that of pollinators (negative sign). Herbivory selects for phenotypes that are further away from that of herbivores (positive sign). These two selective forces act synergistically when the plant phenotype is within $[t_m, t_h]$ so that evolution enables a simultaneous increase in pollination and decrease in herbivory. In contrast, these two forces become antagonistic when the plant phenotype is outside $[t_m, t_h]$. Pollination selects for stronger plant-animal trait matchings, while herbivory selects for weaker trait matchings. Such conflicting selection captures the ecological trade-off with which plants are confronted.

Pollination and herbivory have also an opposite effect on the evolutionary stability – i.e. noninvasibility – of evolutionary singularities t_p^* . Singularities are invasible if inequality (4) is satisfied (proof and expression of $f(t_p^*)$ in Appendix A.II.2).

(4)
$$Invasibility \Leftrightarrow -\underbrace{a_{pm}(t_p^*)M^*}_{pollination} \left[1 + \frac{f(t_p^*)^2}{\sigma_{Pol}^2}\right] + \underbrace{a_{ph}(t_p^*)H^*}_{herbivory} \left[1 + \frac{f(t_p^*)^2}{\sigma_{Her}^2}\right] > 0$$

Invasibility is thus favored by herbivory and disfavored by pollination. Convergence being however not analytically tractable, we rely on a first Monte Carlo experiment (MC1, details in appendix B.II.2) to investigate the relationship between selection type and plant-animal interactions. We sampled 10 000 interspecific parameter sets and analyzed the nature of all evolutionary singularities allowing ecological coexistence, as well as interaction strengths and animal densities at these singularities. The resulting dataset thereby links the ratio of pollination

to herbivory at the singularity $\left(\frac{a_{pm}M^*}{a_{ph}H^*}\right]_{t_p^*}$ to the type of selection. Stabilizing, runaway and disruptive selections are characterized by contrasting pollination to herbivory ratios (Fig. 3), whose variation explains around two-thirds of the variance in selection type (Kruskal-Wallis).



Figure 3 - Distribution of pollination to herbivory ratios depending on the type of selection. The ratio is measured at the corresponding evolutionary singularity. The medians indicated correspond to absolute values (no log). The effect size of the Kruskal-Wallis test ($p_{value} < 2.2 \ 10^{-16}$) is also indicated. Ecological parameter set: ($r_p = 10, r_m = -1, r_h = -4, c_p = 0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h = 0.3$). Results are from our first Monte Carlo experiment (MC1, appendix B.II.2), with 10 000 interspecific parameter sets sampled.

Effects of interspecific parameters on selective landscapes

Beyond local selection at evolutionary singularities, we now investigate the effect of plantanimal interactions on the selective landscape at large, i.e. the distribution of basins of attraction associated with each type of selection (e.g. Fig. 2A). We build a second Monte Carlo experiment (MC2, details in Appendix B.II.3). For each focal interspecific parameter, we divide the potential range of that parameter (table 1) into six equal intervals to sample from, the sampling of the other interspecific parameters remaining unconstrained. For each sampling (6 x 1000), we calculate the proportion of phenotypic space associated with each type of selection (i.e. basin of attraction, see Appendix B.II.1 for details). We thereby estimate the effect of varying (1) the strength of ecological trade-off (Fig. 4A) and (2) the intensity and generalism of pollination and herbivory (Fig. 4B) on the selective landscape.

Stabilizing selection dominates the selective landscape (Fig. 4), indicating that evolutionary dynamics usually tend to maintain plant-animal coexistence. This result is essentially robust to the modification of the ecological parameter set (Appendix C.II, but see Fig. C3.B).

Disruptive selection is possible at strong trade-offs, but not at weak trade-offs that typically lead to stabilizing or runaway selection (Fig 4A). Disruptive selection indeed requires a balance between pollination and herbivory (Fig. 3). Such a balance has to be achieved over a phenotypic region outside $[t_m, t_h]$ for a branching point to occur. Phenotypes within $[t_m, t_h]$ are continuously selected toward the pollinator phenotype and away from the herbivore one so that no singularity is possible between the two animal traits (see equation 3). When the trade-off is strong, balanced interactions naturally emerge as animal phenotypes are similar. When animals have very dissimilar phenotypes, pollination to herbivory ratios are either large or small outside $[t_m, t_h]$, depending on which animal phenotype is closer to the plant phenotype. As a result, selection is either stabilizing or runaway at weak trade-offs. Note finally that disruptive selection is seldom observed with both our robustness ecological parameter sets (Fig. C2.A & C3.A, Appendix C.II), and that it similarly does not occur at weak trade-offs.

The reported opposite influence of pollination and herbivory on local selection extends in a consistent manner to the selective landscape (Fig. 4B). More precisely, the distribution of basins of attraction associated with each type of selection shifts towards a higher prevalence of stabilizing selection with stronger pollination, and a higher prevalence of runaway selection with stronger herbivory.

Increasing the basal rate of pollination (a_{pm0}) favors stabilizing selection at the expense of disruptive and runaway selection (Fig. 4B.a). On the contrary, higher herbivory rates (a_{ph0}) favor disruptive and runaway selection at the expense of stabilizing selection (Fig. 4B.b). These results suggest that the prevalence of disruptive selection is restricted by strong pollination and fostered by strong herbivory (Fig 4B. a vs. b). A narrower pollination niche width (σ_{Pol}) increases the prevalence of stabilizing selection at the expense of disruptive selection (Fig. 4B.c). In contrast, disruptive and runaway selection become more frequent as the herbivory niche width (σ_{Her}) gets narrower (Fig. 4B.d). Variations are, however, less pronounced for niche widths (σ) than for interaction rates (a_0). Moreover, only in the case of interaction rates are these patterns robust to the variation of the ecological parameter set (Appendix C.II). While basal rate variations have a consistent effect on interaction strength across the phenotypic space, niche width variations increase or decrease interaction strength depending on plant-animal trait matching (i.e. $t_p - t_m$, $t_p - t_h$), likely explaining such a difference.

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Figure 4 - Effect of interspecific parameters on the selective landscape. A. The selective landscape depends on trade-off intensity. B. Opposite effects of pollination and herbivory on selection. Please note that animal phenotype dissimilarity $(t_h - t_m)$ was here further constrained within [0, 1.5]. This was done to better capture the effects of the other interspecific parameters on disruptive selection, the latter being hardly observed for high animal phenotype dissimilarities (i.e. for weak trade-offs, see Fig. 4A and main text). a. b. Variations in basal interaction rates. c. d. Variations in niche widths. Results are from our second Monte Carlo experiment (MC 2, appendix B.II.3), with 1000 interspecific parameter sets sampled at each point. Y-axes indicate the normalized size of the basins of attraction (Mean \pm SE) associated with each type of selection (see appendix B.II.1). Ecological parameter set: $(r_p = 10, r_m = -1, r_h = -4, c_p = 0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h = 0.3)$. See Fig. C2 & C3 (Appendix C.II) for robustness.

Changes in the niches of pollination and herbivory modify community eco-evolutionary dynamics, but these modifications are mainly driven by changes in evolutionary dynamics at strong trade-offs, and changes in ecological dynamics at weak trade-offs (Fig. 5 & S2, Supporting Material).

At weak trade-offs, varying interaction basal rates (Fig. 5) or niche widths (Fig. S2) mainly affects the composition and stability of the ecological communities, i.e. background colors in Fig. 5 & S2. The selective landscape, characterized by stabilizing vs. runaway selection close to the pollinator vs. herbivore phenotype (as in Fig. 2A), remains essentially unaltered. In particular, increasing the basal rate of pollination eventually selects for phenotypes leading to unstable community dynamics (e.g. Fig. 5a vs. 5b, unbounded growth). Increasing the rate of herbivory restores stability (e.g. Fig. 5b vs. 5d). Note also the narrower set of plant phenotypes allowing coexistence when herbivory becomes more specialized (e.g. Fig. S2a vs. S2c). Close to the herbivore phenotype, the resulting increase in herbivory strength (equation 2) reduces plant density so that pollinators no longer survive. Close to the pollinator phenotype, the resulting decrease in herbivory strength impedes herbivore survival. Plant evolution within the three-species community then provokes the extinction of herbivores. More specialized pollination can restore coexistence, but exposes the community to unstable dynamics (Fig. S2c vs. S2d, unbounded growth).

At strong trade-offs, basal rates or niche widths variations primarily modify the selective landscape – i.e. evolutionary singularities in Fig. 5 & S2 – while the ecological context remains essentially unaffected. Evolutionary dynamics may utterly change. For instance, increasing herbivory can shift the selection regime from stabilizing to disruptive (Fig. 5a vs. 5c), due to the modification of the number of singularities (one vs. three), and of their type (a CSS becoming a branching point). The plant-animal community is significantly modified by the emergence of plant dimorphism, maintained in this case. Modifications of the position of singularities, CSSs in particular, can also have far-reaching implications for the maintained community. A stronger herbivory displaces CSS phenotypes closer to animal extinction thresholds (e.g. Fig. 5a vs. 5c), which implies a fragile coexistence. Given small animal densities, coexistence is indeed vulnerable to perturbations or stochasticity. Stronger pollination, in contrast, displaces the CSS phenotypes away from extinction boundaries (e.g. Fig. 5c vs. 5d).

Maintenance of plant dimorphism arising from disruptive selection

Disruptive selection on the plant phenotype arises from the interplay of pollination and herbivory. The resulting dimorphism is often temporary, but stable dimorphism is possible in the case of strong trade-offs. Simulating the eco-evolutionary dynamics following the 676 branching points encountered during our second Monte Carlo experiment when constraining the sampling of trade-off intensity, we indeed find that dimorphism is maintained in only 6% of the cases, all occurring at very strong trade-offs ($t_h - t_m \le 0.5$). At such trade-offs, however, the maintenance of dimorphism is relatively frequent, representing around 27% of the cases (details in appendix D).

Interestingly, we find that the ratio of pollination to herbivory at branching $\left(i.e. \frac{a_{pm}M^*}{a_{ph}H^*}\right]_{t_n^*}$ again

largely explains the maintenance of dimorphism at these strongest trade-offs (Wilcoxon test, $p_{value} < 10^{-10}$ & $effect_{size} \approx 50\%$, see Fig. 6A). Dimorphism maintenance is associated with stronger herbivory than pollination, which suggests a decisive role of herbivory in the process. We finally show that even a slight amount of similarity-dependent competition suffices to maintain dimorphism in the cases where it is lost (Fig. 6B). To do this, we reformulate our model assumptions to make intraspecific plant competition depend on the phenotypes of competing plant individuals (equation 5). Competition between two plant phenotypes (t_{p1}, t_{p2}) now declines as their niche-overlap (i.e. $t_{p1} - t_{p2}$) decreases, α_c and σ_c being the input parameters controlling the proportion and shape of similarity-dependent competition, respectively.

(5)
$$c_p \rightarrow c_p(t_{p1}, t_{p2}) = (1 - \alpha_c)c_p + \alpha_c e^{\frac{-(t_{p1} - t_{p2})^2}{2\sigma_c^2}}c_p$$

Five percent of similarity-dependence can enable the maintenance of dimorphism (Fig. 6B. a vs. b). Ten percent of similarity-dependence can lead to secondary branching events and the emergence of plant polymorphism (Fig. 6B.c), which suggests a potential role of the interplay between pollination, herbivory and associated niche-overlap competition in the diversification of flowering plant species.



Figure 5 - Effect of the basal rates of plant-animal interactions on community ecoevolutionary dynamics. At strong trade-offs, eco-evolutionary dynamics are primarily altered by changes affecting evolutionary singularities. At weak trade-offs, the alteration is mainly mediated by changes in the composition of the ecological community (i.e. background colors). Orange and red dotted lines indicate the pollinator and herbivore phenotype. Arrows indicate evolutionary trajectories and background colors indicate community composition as in Fig. 2A, with light blue figuring regions in which our model fails to produce realistic dynamics (unbounded population growth). <u>Ecological parameter set:</u> ($r_p = 10, r_m = -1, r_h = -4, c_p =$ $0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h = 0.3$). <u>Interaction niche widths:</u> $\sigma_{Pol} = 1.7$, $\sigma_{Her} = 2$.



Figure 6 - The maintenance of plant dimorphism. A. Ratio of pollination and herbivory at branching explains the maintenance of plant dimorphism. The branchings considered here are those occurring at very strong trade-offs ($t_h - t_m \le 0.5, 154/676$), given that dimorphism maintenance is only possible at such strong trade-offs. Wilcoxon $p_{value} < 10^{-10}$. B. Emergence and maintenance of polymorphism are favored by similarity-dependent competition. a. No similarity-dependent competition. The branching corresponds to that presented in Fig. 2B.e. b. Five percent of similarity-dependent competition enables the maintenance of plant dimorphism. c. Ten percent of similarity-dependent competition enables secondary branchings, leading to a polymorphic plant population. Ecological parameter set: ($r_p = 10, r_m = -1, r_h = -4, c_p = 0.6, c_m = 0.5, c_h = 0.4, e_m = 0.2, e_h = 0.3$). Interspecific parameter set: ($a_{pm0} = 5, a_{ph0} = 7, \sigma_{Pol} = 3, \sigma_{Her} = 2.8$). Animal phenotypes are indicated with orange (pollinator) and red (herbivore) dotted lines: ($t_m = 0, t_h = 0.2$).

Discussion

In this article, we investigate how the eco-evolutionary dynamics of plant-pollinator-herbivore communities are driven by the evolution of plant traits under conflicting selection. Conflicting selection arises from the ecological trade-off between attracting pollinators and escaping herbivores which naturally emerges in a trait-matching framework. We find that stronger pollination makes stabilizing selection more prevalent and facilitates coexistence. Stronger herbivory

increases the prevalence of runaway selection and threatens coexistence. Importantly, joint selection may be disruptive, leading to plant diversification. Such a diversification requires strong trade-offs, with the strongest trade-offs allowing its long-term maintenance. At weak trade-offs, coexistence is threatened as runaway dynamics are more frequent. Stabilizing selection can still maintain coexistence, but the intrinsic imbalance of plant-animal interactions makes it vulnerable to perturbations. The strength of the ecological trade-off appears as a structuring determinant of community eco-evolutionary dynamics.

We find that pollination fosters the convergence toward the pollinator phenotype – i.e. stabilizing selection – while herbivory favors the divergence from the herbivore phenotype – i.e. runaway selection. Such opposite selection pressures have notably been found acting on flower color (Irwin et al., 2003; Frey, 2004), shape (Galen & Cuba, 2001) or display (Gómez, 2005), nectar quantity (Adler & Bronstein, 2004) and flowering phenology (Brody, 2008; Ehrlén & Münzbergová, 2009). Conceptually, all traits that make the discovery of a plant species easier for interacting species – i.e. more "apparent" (Feeny, 1976), e.g. high abundance, large size, wide phenology... – may experience ecological trade-offs as they facilitate the discovery by both allies and enemies. A consequence of such trade-offs is that plant phenotypes deviate from optima favored by pollinators (CSSs $\neq t_m$ in Fig. 5), as illustrated by Ramos & Schiestl (2019). Using experimental evolution, the authors show that plants that evolved in the presence of both pollinators and herbivores were less attractive to pollinators than those that evolved with only pollinators, but still more attractive than plants that evolved with hand-pollination, with or without herbivores. The presence of pollinators thus selected toward plant-pollinator trait matching, but the presence of herbivores reduced such matching by adding a runaway component to selection (equation 3).

Recent data have highlighted the occurrence of vast insect declines (Hallmann et al., 2017; Outhwaite et al., 2022). Such declines may be accompanied by changes in the relative frequency of antagonistic vs mutualistic interactions (e.g. herbivores vs pollinators). Given massive pollinator declines (Potts et al., 2010) for instance, plant-animal interactions could become increasingly biased towards antagonism. Given our results, this should favor runaway selection thereby leading to plant phenotypes maladapted to pollinators, further accelerating their declines (see also Weinbach et al. 2022). Herbivore extinctions are also expected as a result of more frequent runaway dynamics. Contrary to our assumption of being fixed, animal phenotypes also evolve to match their resource phenotype in natural settings (e.g. Berenbaum and Zangerl 1998; Muchhala 2006). Runaway dynamics are therefore not expected to systematically provoke the extinction of one or both animals, but their ecological outcome will rather depend on the relative speed of evolution of the different species (e.g. evolutionary rescue, Gomulkiewicz and Holt 1995).

More generally, several assumptions that simplify the evolutionary process are made within our modeling framework. The assumption of fixed animal phenotypes might prove to be relevant for some specific communities in which the plant evolves at a much faster rate than the animals owing to higher mutation rates, standing genetic variation, reproduction rates or lifespans. Animal morphologies, preferences, detection abilities or phenologies may however coevolve with plant phenotypes in natural communities (Thompson, 2009). Results of this coevolution will depend on the relative speeds of evolution of all interactors, which may lead to many different scenarios whose analyses go beyond the scope of this article. Our work aims at developing an in-depth analytical understanding of the dynamics resulting from the sole plant evolution, before more complex coevolutionary scenarios are studied. We acknowledge that it is not straightforward to extend our results to these scenarios as a higher trait dimensionality can substantially alter evolutionary dynamics (Kisdi, 2006; Débarre et al., 2014). Another assumption of our framework is that plant evolution has no other costs than those ensuing from pollination and herbivory. We used this assumption to better highlight the direct implications of ecological trade-offs. This might be a reasonable approximation for specific traits whose selective pressures essentially arise from these plant-animal interactions (e.g. presumably flower color in Raphanus sativus, Irwin et al. 2003), but plant traits are highly multifunctional in general. They are indeed often also involved in other types of interactions (Strauss & Whittall, 2006) or processes such as stress tolerance (Sack & Buckley, 2020). A direct consequence is that phenotypes are usually constrained within an

ecologically viable space, which notably sets a limit to runaway dynamics, mitigating the likelihood of extinctions.

In line with the empirical evidence so far (Strauss et al., 2002; Strauss & Irwin, 2004), our study identifies the strength of ecological trade-off as a key driver of plant-animal eco-evolutionary dynamics. We underline that trade-off intensity affects the maintenance of coexistence, as well as the nature of joint selection, in particular the potential for disruptive selection and plant diversification.

We find that coexistence is fragile or destroyed at weak trade-offs. Runaway dynamics are more frequent (Fig. 4A). Evolutionary-induced unstable dynamics become possible (Fig. 5b&d). In the vicinity of the pollinator phenotype, the plant phenotype is more likely to stabilize after the extinction of herbivores, as indicated by the CSSs leaving the coexistence area (Fig. 2A). When stabilizing selection occurs before herbivore extinction, their low density makes them vulnerable to demographic stochasticity, Allee effects or external disturbances. Because weak ecological trade-offs are associated with fragile plant-pollinator-herbivore coexistence, they should be seldom encountered in natural communities. Accordingly, shared animal preferences for plant phenotype are most often reported in the empirical literature (Adler & Bronstein, 2004; Strauss & Whittall, 2006).

Selection can turn disruptive under stronger ecological trade-offs. We stress that disruptive selection here arises from the joint selection of pollination and herbivory. In the absence of either one interaction, branching is not possible here, as evolutionary dynamics would run away from the herbivore phenotype in the absence of pollination, and would stabilize at the pollinator phenotype in the absence of herbivory. Diversification we uncover here is thus fundamentally different from the one uncovered in previous trait-matching models that consider the two interactions separately (e.g. Yoder and Nuismer 2010; Maliet et al. 2020). While dimorphism can only be maintained when the trade-off is sufficiently strong, long-term maintenance is frequent in such instances. Polymorphism then yields two contrasted phenotypes: one that has many allies and enemies, while the other one has few interactions. Interestingly, such a pattern has been frequently reported in the empirical literature. In Primula farinosa, the fitness advantage of long-scaped phenotypes resulting from higher pollinator visitation rates is balanced by higher levels of fruit predation, allowing the maintenance of dimorphism in scape length (Ehrlén et al., 2002). The maintenance of color dimorphism in Raphanus sativus (Irwin et al., 2003) and Claytonia virginica (Frey, 2004) is similarly supported by animal species preferentially interacting with the same plant phenotype. Field studies of tripartite networks also report a positive correlation between the number (Sauve, Thébault, et al., 2016) or strength (Melián et al., 2009) of plant-pollinator and plant-herbivore interactions of plant species, indicating a rather widespread pattern. This mechanism of diversification under balanced interaction strengths may also apply on longer evolutionary timescales. Using a phylogenetic analysis on 32 Nicotiana species, Adler et al. (2012) show that variations in nicotine defenses among species are largely negatively correlated to pollinator reliance. These tobacco species are therefore either well-defended with few pollinators, or lowly defended and relying on pollination.

We find that disruptive selection requires a local balance between pollination and herbivory (Fig. 3), while its prevalence within the selective landscape increases with the strength of herbivory (Fig. 4B). This is not contradictory as those aspects describe selection at two different scales. To put it simply, disruptive selection can be expected in any three-species community when plant-animal interactions are well-balanced, but communities in which herbivores are more ravenous ($a_{ph0} \nearrow$) will more likely exhibit such evolutionary dynamics. Dimorphism is also more likely to be permanent with stronger herbivory (Fig. 6A), further emphasizing its decisive role in diversification processes. Previous theoretical works that contrast interaction types but one at a time have also highlighted the crucial role of antagonisms for diversification (Kopp & Gavrilets, 2006; Yoder & Nuismer, 2010; Maliet et al., 2020). Given these results, we note that observed declines in the diversity and abundances of herbivore species (Sánchez-Bayo & Wyckhuys, 2019; Atwood et al., 2020) may have far-reaching consequences if they lead to future reductions in diversification rates.

Overall, dimorphism is often temporary within our framework. While we assume constant competition rates in order to focus on how trophic vs mutualistic interactions shape ecoevolutionary dynamics, similarity-dependent competition is likely to occur in natural communities (Macarthur & Levins, 1967). Here, similar plants would share pollinators, which could lead to strong pollination niche-overlap competition for instance due to the dilution of pollen between plant species (Morales & Traveset, 2008; Mitchell et al., 2009). Similarly, shared herbivores would increase apparent competition. Plant traits can also be simultaneously involved in both competitive and plant-animal interactions, either directly (e.g. phenological traits, Schwinning & Kelly 2013) or due to genetic correlations (e.g. plant size, Carmona et al. 2011). Accounting for a small amount of niche-overlap competition suffice to maintain dimorphism in our model, and enhances the potential for further diversification (Fig. 6B). Reproductive isolation may then evolve as the morph that interacts weakly with animal species acquires new pollinators species (Baack et al., 2015), or evolves self-fertilization (Bodbyl Roels & Kelly, 2011). Disruptive selection from the interplay of pollination, herbivory and niche-overlap competition can therefore be construed as a first step toward modeling the evolutionary emergence of plant-pollinator-herbivore networks, which would allow new insights into the eco-evolutionary processes supporting the diversity of natural communities.

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Data, scripts, code and supplementary information availability

The codes used to conduct the Monte Carlo numerical experiments, as well as that used to simulate the community eco-evolutionary dynamics, are available in Zenodo (https://doi.org/10.5281/zenodo.11263481; Yacine & Loeuille, 2024). Supplementary material referenced in text, which notably includes the derivation of mathematical expressions and demonstrations, is available via the same link. It consists of 2 additional figures (S1, S2), one table (S1) and 4 appendices (A, B, C & D).

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