1	Associational effects and the maintenance of polymorphism in plant
2	defense against herbivores: review and evidence
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#### 25 Abstract

26Many plant species have evolved defense traits against herbivores. Associational effects 27(AE) refer to a kind of apparent interaction where the herbivory risk to a focal plant 28species depends on the composition of other plant species in a neighborhood. Despite 29ample evidence for AE between different plant species, this point of view has rarely 30 been applied to polymorphism in defense traits within a plant species. The purpose of 31 this review is to highlight an overlooked role of conspecific AE in maintaining 32polymorphism in antiherbivore defense. First, I present a general review of AE between 33 plant species and its role in the coexistence of plant species. This viewpoint of AE can 34be applied to genetic polymorphism within a plant species, as it causes frequency- and 35 density-dependent herbivory between multiple plant types. Second, I introduce a case 36 study of conspecific AE in the trichome-producing (hairy) and glabrous plants of 37 Arabidopsis halleri subsp. gemmifera. Laboratory and semi-field experiment illustrated 38 that AE against the brassica leaf beetle *Phaedon brassicae* mediate a minority advantage 39 in defense and fitness between hairy and glabrous plants. Combined with a statistical 40 modeling approach, field observation revealed that conspecific AE can maintain the 41 trichome dimorphism via negative frequency-dependent selection in a plant population. 42Finally, I discuss spatial and temporal scales at which AE contribute to shaping genetic 43variation in antiherbivore defense in a plant metapopulation. Based on the review and 44evidence, I suggest that AE play a key role in the maintenance of genetic variation 45within a plant species.

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47 Keywords: Frequency-dependent selection, Genetic variation, Herbivory, Neighbor
48 effects, Plant defense

#### 49 **1. Background**

50Many plant species have evolved defensive traits, such as spines and toxins, against 51herbivores (Schoonhoven et al. 2005). Natural plant populations exhibit genetic 52variation in regard to chemical (Hughes 1991; van Dam et al. 1999; Chan et al. 2010) 53and morphological defenses (Kivimäki et al. 2007; Wise et al. 2009). It is well known 54that these defense traits are costly for plant growth and/or reproduction (e.g., Mauricio 551998; Elle et al. 1999; Züst et al. 2011). The theory of plant defense assumes that the 56defense-growth tradeoff favors an optimal balance between the cost and benefit of 57defense (Simms 1992), extinguishing variation in defense levels. Thus, it is difficult for 58natural selection to maintain genetic variation in antiherbivore defense traits unless a 59stabilizing mechanism occurs.

60 Plant defense theory has focused on how individual plants allocate their 61 resource to defensive traits (e.g., Rhoades 1979; Simms 1992; Stamp 2003). However, 62 the defensive effects of a particular trait depend not only on a plant's own trait but also 63 on the composition of the other plant types in a neighborhood (Agrawal et al. 2006; 64 Barbosa et al. 2009; Underwood et al. 2014). These phenomena are called associational 65 effects (AEs: also known as *neighbor effects*), which occur when "consumer effects on 66 individuals of one resource organism type, at a given density of that type, are a function 67 of the neighborhood composition of other resource types at particular spatial scales" 68 (Underwood et al. 2014). An increasing number of studies have shown the importance 69 of AE in maintaining plant species diversity (e.g., Hay 1986; Callaway et al. 2005; 70Stastny & Agrawal 2014), but this point of view has rarely been tested in the 71evolutionary ecology of antiherbivore defenses. 72The purpose of the present review is to highlight the overlooked but potentially 73critical role of conspecific AE in the maintenance of genetic variation in antiherbivore 74defense traits. This paper consists of three chapters, in which I first outline the 75ecological roles of AE in maintaining plant species diversity. Second, I introduce a case 76 study on Arabidopsis trichomes to expand the concept of AE to polymorphism in 77defense within a plant species. Finally, I discuss how AE contributes to intraspecific 78variation in plant defense at larger spatial and temporal scales. Based on the review and 79evidence, the present paper suggests that AE plays a key role in maintaining not only 80 species diversity but also genetic variation in antiherbivore defense.

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### 82 1-1. Examples of AE in plant-herbivore interaction

83 The phenomena of associational resistance or susceptibility have been reported across

agricultural (Tahvanainen & Root 1972; Risch 1981; Le Guigo et al. 2012), forest

85 (White & Whitham 2000; Sholes 2008; Castagneyrol et al. 2013), grassland (Agrawal

86 2004; Courant & Fortin 2010; Hahn & Orrock 2016), semi-arid (González-Teuber &

Gianoli 2007), wetland (Rand 2003; Hughes 2012), and aquatic (Hay 1986; Wahl &

Hay 1995) ecosystems (see Appendix for selected examples). Two major patterns have

89 been reported for AE between plant species: Associational resistance or susceptibility

90 refer, respectively, to a situation where a focal plant species is *less* or *more* damaged in

91 the presence of another plant species. Undefended plants may gain "associational

92 resistance" when the defended plants are protected from herbivory by nearby plants

93 (Tahvanainen & Root 1972; Rausher 1981; Hambäck et al. 2000). In the polyculture of

94 the collard green *Brassica oleracea* var. *acephala* (Brassicaceae), Root and colleagues

95 for the first time developed the concept of "associational resistance" and "resource

96 concentration," which refer to the effects of vegetation composition and total density on

97	herbivore populations, respectively (Tahvanainen & Root 1972; Root 1973). Recently, it
98	was also discovered that neighboring plants do not always reduce herbivory damage to
99	other plant species. The term "associational susceptibility" was coined to refer to the
100	situation where defended plants suffer from severe herbivory when they occur nearby
101	undefended ones (White & Whitham 2000; Rand 2003). Barbosa et al. (2009) introduce
102	many more examples of AE and Rautio et al. (2012) outline the terminology regarding
103	AE, comprising the terms "plant defense guild" (Atsatt & O'Dowd 1976), "shared
104	doom" (Wahl & Hay 1995; Emerson et al. 2012), and "associational refuge" (Hjältén et
105	al. 1993; Miller et al. 2009; Emerson et al. 2012; Hughes 2012).
106	The mechanisms underlying AE are involved in various combinations of plant
107	traits and herbivore behaviors. In general, herbivores alter their preferences and/or
108	performance depending on their dietary menu (e.g., Lefcheck et al. 2013; Sato & Kudoh
109	2016a; Wetzel et al. 2016). In particular, theory of optimal foraging suggests that
110	consumers should utilize suboptimal resource as the density of optimal resource
111	becomes low (e.g., Charnov 1976). This behavioral basis of consumers directly alters
112	the consumption rate and thus drives apparent competition between optimal and
113	suboptimal resource (Holt & Kotler 1987). Empirically, mammalian herbivores often
114	avoid spiny or toxic herbs and thereby lead these defended herbs to confer associational
115	resistance to undefended plant species at a patch level (e.g., Hjältén et al. 1993;
116	Callaway et al. 2005; Courant & Fortin 2010; Miller et al. 2009). Visual or chemical
117	crypsis conferred by neighboring plants results in associational resistance against insect
118	herbivores (Tahvanainen & Root 1972; Rausher 1981; Hambäck et al. 2000). Volatile
119	organic chemicals from neighboring plants sometimes induce plant resistance traits and
120	thus drive associational resistance against herbivores (Karban & Maron 2002; Karban

2007; Zakir *et al.* 2013). Direct competition sometimes increases plants' investment in
defense traits and may promote associational resistance (Barton & Brower 2006;
González-Teuber & Gianoli 2007). Polyphagous herbivores move from primary to
secondary host species during their dispersal process, resulting in associational
susceptibility on the part of the former to the latter host plant (White & Whitham 2000;
Agrawal 2004). These examples suggest that the outcomes of AE depend on herbivore
behaviors and plant trait changes.

128While long recognized as apparent competition, the outcomes of AE are more 129 complex than simple competition between two plant species. AE comprise facilitation, 130 exploitation, and other kinds of species interaction (Agrawal 2004; Barbosa et al. 2009). 131 In particular, recent reviews pointed out problems with the terminologies and 132experimental designs used to test AE (Rautio et al. 2012; Underwood et al. 2014), as 133 there has been little consensus regarding the trait of a focal plant type (i.e., defended or 134undefended) and manipulations on the neighborhood composition (presence/absence, 135 relative abundance, or total density of multiple plant types) (see also Appendix). For 136 example, a number of studies compared the magnitude of herbivory on one of two plant 137 species between the presence and absence of another species (Hambäck *et al.* 2000; 138 White & Whitham 2000); however, this setting is unable to determine the likelihood of 139 the coexistence of two plant species (Underwood *et al.* 2014). Many studies focused on 140 the absolute amount of herbivory (Tahvanainen & Root 1972; Hambäck et al. 2000; 141 White & Whitham 2000), while others compared the relative amounts of herbivory 142between multiple plant types (Bergvall et al. 2006; Castagneyrol et al. 2013). 143 Furthermore, some studies focused on the herbivory load to individual plants within a 144 small spatial scale (referred to as a "patch": Rausher 1981; Hambäck et al. 2000;

Karban & Maron 2002) while others compared herbivory among the plant patches at
broader spatial scales (Bergvall *et al.* 2006; Castagneyrol *et al.* 2013). Details of the
experimental design and outcomes are listed for 16 publications (see Appendix).

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149 1-2. Frequency/density-dependent damage caused by AE

150Recently, several authors have begun to discuss how AE enable the coexistence of 151multiple plant species. Underwood et al. (2014) advocated that the focus should be on 152the frequency- or density-dependent damage caused by AE to multiple plant species. If 153AE is more likely to protect rare plant species than abundant species, this rarity 154advantage (i.e., negative frequency dependence) in defense would prevent rare species 155from going extinct due to herbivory, thereby promoting the coexistence of multiple 156plant species. In this refined framework, negative or positive frequency-dependent 157damage indicates associational resistance or susceptibility for rare plant species, 158respectively. Even if multiple plant species can coexist via AE, recovery from low 159density (i.e., negative density dependence) is necessary for the long-term persistence of 160 plant populations. The negative or positive density-dependent damage refers to the 161 "resource concentration or delusion" hypothesis, initially developed by Root (1973) and 162recently reviewed by Underwood et al. (2014) and Hambäck et al. (2014). 163 Several studies suggest the importance of plant frequency and density in 164 determining the outcome of AE, although those showing the importance of both the 165density and frequency dependence are still limited (Table 1). Kim and Underwood 166 (2015) manipulated the density and frequency of the tall goldenrod Solidago altissima 167 (Asteraceae) and the Carolina horsenettle Solanum carolinense (Solanaceae). As a result, 168 S. carolinense received more damage due to the combined effects of the high

169 conspecific density and intermediate frequency of the two species (Kim & Underwood 170 2015). Hahn and Orrock (2016) manipulated both the density and frequency of two 171 related species Solidago nemoralis and S. odora (Asteraceae) in the field and then 172revealed herbivore foraging to be a key mechanism with a behavioral assay. The less 173defended species S. nemoralis gained associational resistance when it was rare due to 174the reduced foraging activity of grasshoppers, with the two *Solidago* species more likely 175attacked at a higher density (Hahn & Orrock 2016). In relation to herbivore foraging, 176Verschut et al. (2016) showed that low-concentration of balsamic vinegar was less 177likely to attract the fruit fly Drosophila melanogaster at a high frequency of high 178 concentration diets, while high-concentration diets were more likely to attract flies at a 179 high frequency of low-concentration diets. Previous studies on the fallow deer Dama 180 *dama* manipulated the tannin concentration as well as the frequencies of two resource 181 types (Bergvall & Leimar 2005; Bergvall et al. 2006). High tannin diets were consumed 182less frequently when low tannin diets were abundant, whereas low tannin diets were 183 consumed more frequently when high tannin diets were abundant (Bergvall & Leimar 184 2005). Furthermore, if two species co-occur in their natural habitats, field surveys help 185 us capture the effects of plant density or frequency in the wild population. Russell and 186 Louda (2005) documented that the native thistle *Cirsium undlatum* (Asteraceae) 187 received less florivory from the exotic weevil Rhinocyllus conicus as the number of 188 another native thistle (*C. canescens*) increased in close vicinity. This long-term survey 189 found that an invasive herbivore creates a novel AE due to the overlap of flowering 190 phenology between two native plant species (Russell & Louda 2004; Russell & Louda 191 2005).

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Although density- or frequency-dependent herbivory have been shown

193 experimentally, we still know little about how AE occurs and contributes to the 194 maintenance of plant species diversity. Thus, it is necessary to link the mechanisms and 195 patterns of AE in a single study system. There is a common set of empirical approaches 196 for understanding the mechanisms and consequences of AE from laboratory to the field 197 study (Fig. 1: see also Appendix for details). The first step is to discover candidate 198 mechanisms of AE (Fig. 1), including the identification of key plant traits and herbivore 199 behaviors, and then to address how they cause non-additive herbivory between multiple 200plant species. Previous studies have done this by conducting laboratory choice 201experiments (e.g., Tahvanainen & Root 1972; White & Whitham 2000; Rand 2003) or 202 by observing herbivore movements between plants (Rausher 1981; Bergvall et al. 2006). 203 The second step includes experimental tests to determine whether plant damage and/or 204 fitness depend on the neighborhood composition. This step is adopted by almost all the 205studies and is the most important for illustrating the relevance of AE. The third step is 206 multi-year field survey to corroborate the experimental evidence with field observations 207 on plant damage and population dynamics across years (Fig. 1). This final step is 208 executable if the focal plant species co-occur within natural populations (Russell & 209 Louda 2004; Callaway et al. 2005). These comprehensive tests from laboratory 210experiment to field observations will reveal the causes and consequences of AE in a 211single study system.

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213 1-3. Plant defense polymorphism via the lens of conspecific AE

214 While the concept of AE was originally developed in interspecific interactions

215 (Tahvanainen & Root 1972; Root 1973), it can be applied to genetic variation within the

same plant species. In agricultural ecosystems, polyculture has been carried out not only

217for different species but also for different cultivars within the same plant species (e.g., 218Cantero & Sanford 1984; Hambäck et al. 2009). The emerging framework of 219"community genetics" may be considered a kind of conspecific AE (Hughes et al. 2008), 220 as these studies have shown that the polyculture of different genotypes alters herbivore 221abundance (e.g., Crutsinger et al. 2006; Johnson et al. 2006; Parker et al. 2010). Studies 222 on community genetics also revealed that plant genetic diversity altered the strength of 223 selection on plant genotypes via changes in plant-herbivore interaction (e.g., Johnson et 224al. 2006; Lankau & Kliebenstein 2009; Parker et al. 2010), although the way in which 225frequency-dependent selection occurs between plant genotypes remains to be explored. 226 Evolutionary game theory provides a logical basis for the roles of AE in 227maintaining polymorphism in defense levels (Sabelis & de Jong 1988; Augner et al. 228 1991; Till-Bottraud & Gouyon 1992; Tuomi et al. 1999). Theoretical studies analyzed 229conditions under which the benefits from AE and the intrinsic cost of defense allow 230defended and undefended plants to coexist. In clonal plants, for example, Till-Bottraud 231and Gouyon (1992) predicted that herbivore preference and the cost of defense lead to 232 the evolutionary stable strategy of producing cyanogenic and acyanogenic clones at an 233intermediate frequency. Empirically, this hypothesis regarding evolutionary stability can 234be tested by asking whether or not rarer plant types have higher fitness compared to 235abundant ones under AE (reviewed by Rautio et al. 2012). If a rarity fitness advantage 236reciprocally occurs among multiple plant genotypes, AE leads to negative 237 frequency-dependent selection that can prevent rare genotypes from going extinct 238 (Clarke 1964; Ayala & Campbell 1974). This criterion would allow empirical 239 researchers to test whether AE can be a mechanism maintaining polymorphism in 240antiherbivore defense.

241Despite its logical basis, only a few studies have focused on conspecific AE 242with regard to the maintenance of polymorphism in antiherbivore defense (Table 1). 243Wise et al. (2009) tested AE in the tall goldenrod Solidago altissima, which possesses a 244genetically based phenotype of ducking stem (called the "candy-cane" morph). This 245candy-cane morph occurs with a rare frequency but across natural populations of S. 246altissima (Wise 2009). Wise et al. (2009) conducted a manipulative experiment with 247two contrasting frequencies of candy-cane and erect-stem morphs (Table 1). The tall 248goldenrod received less damage from the specialist gall fly Eurosta solidaginis when 249the candy-cane morphs were abundant; however, the damage was frequency 250independent, as shown by the lack of a significant morph-by-frequency interaction 251(Wise et al. 2009). Recently, Garrido et al. (2016) tested frequency-dependent selection 252between tolerant (i.e., defense mitigating negative effects of herbivory on plant fitness) 253and resistant (defense directly reducing herbivore attacks) genotypes in the jimson weed 254Datura stramonium (Solanaceae), but found a rarity "disadvantage" against the 255maintenance of defense polymorphism. Thus, it remains unknown whether conspecific 256AE can be a mechanism for the maintenance of defense dimorphism. My collaborators 257and I have investigated AE against a leaf beetle in the trichome dimorphism of 258Arabidopsis halleri subsp. gemmifera (Brassicaceae). In the next chapter, I will 259introduce an example of conspecific AE and its stabilizing effect on a plant defense 260polymorphism.

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### 262 **2. Evidence**

Several species of the genus *Arabidopsis* exhibit a dimorphism of trichome-producing
(hairy) and trichomeless (glabrous) plants. The glabrousness of *Arabidopsis* species is

266 GLABRA1 (GL1 also known as GLABROUS1) (Hauser et al. 2001; Kivimäki et al. 2672007; Kawagoe et al. 2011; Bloomer et al. 2012). Because GL1 is a transcriptional 268factor gene for the initiation of trichome development, the loss of function of this gene 269results in distinct phenotypes of hairy and glabrous plants (Oppenheimer et al. 1991; 270Hülskamp 2004; Ishida et al. 2008). The glabrous phenotype is recessive to the hairy 271ones, and these phenotypes are inherited in the Mendelian fashion (Kärkkäinen & Ågren 2722002). Laboratory studies and field surveys have shown that hairy plants incurred less 273herbivory by insect herbivores compared to glabrous plants (Handley et al. 2005; 274Sletvold et al. 2010; Løe et al. 2007; Kivimäki et al. 2007), while trichomes impose a 275fitness cost on plants (Mauricio 1998; Sletvold et al. 2010; Züst et al. 2011). Owing to 276 the visible and discrete phenotypes with the antiherbivore function, Arabidopsis 277 trichomes provide an excellent system to investigate AE against herbivores within a 278same-plant species. 279Likewise, Arabidopsis halleri subsp. gemmifera (abbreviated hereafter as A.

associated with the loss of function of the key gene of trichome development,

280 halleri) have a dimorphism of hairy and glabrous plants (Fig. 2a). Hairy plants have 281non-glandular trichomes on their leaves and stems, whereas glabrous plants produce no 282trichomes, except on their stem and leaf margin. This presence/absence of leaf 283trichomes is associated with the allelic status of a GL1 orthologue (Kawagoe et al. 2842011). Hairy plants showed a lower fecundity and slower growth than glabrous ones did 285in the absence of herbivores (Kawagoe et al. 2011; Sato & Kudoh 2016b; Sato & Kudoh 2862017b), indicating a fitness cost of the trichome production. Further, trichome 287 production is unlinked with glucosinolate profiles (Sato *et al.* 2014), which may act as a 288chemical defense for Brassicaceae (e.g., Züst et al. 2012).

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289Arabidopsis halleri subsp. gemmifera is a self-incompatible perennial herb 290 distributed across Japan and the Russian Far East. Plants bloom from early April to May 291at the lowland of Japan (Kawagoe & Kudoh 2010). Flowers are pollinated by hoverflies 292and solitary bees. Seeds are dispersed by gravity. After flowering, plants produce clonal 293rosettes on the main and lateral meristems (Sato & Kudoh 2017b), wherein late-spring 294herbivory can directly affect the sexual and asexual reproduction. Arabidopsis halleri 295subsp. gemmifera is attacked by multiple species of herbivorous insects including 296 beetles, butterflies, and sawflies from late-spring to early-summer (Sato & Kudoh 2015; 297Sato & Kudoh 2017a). Among natural populations of A. halleri, the brassica leaf beetle, 298Phaedon brassicae (Fig. 2a), occasionally outbreaks and impacts fruit production by 299 heavily infesting flowering stems (Kawagoe & Kudoh 2010).

300 In this second chapter, I introduce a case study of conspecific AE against the 301 leaf beetle on the trichome dimorphism of A. halleri. This study consists of three 302specific approaches (Fig. 2). The first stage is to identify candidate mechanisms of 303 frequency-dependent damage between hairy and glabrous plants. In the second stage, 304 manipulative experiments are conducted to test fitness consequence and to address 305 whether AE against the leaf beetle results in negative frequency-dependent selection 306 between the two morphs. The final step is to link the experimental findings with field 307 observations by modeling mechanisms that govern plant population dynamics. These 308 three steps will provide comprehensive evidence that AE play a stabilizing role in plant 309 defense polymorphism.

310

311 2-1. Candidate mechanisms of AE

312 Behavioral response of herbivores to multiple resources can be a mechanism of AE (e.g.,

313 Bergvall et al. 2006; Verschut et al. 2016; Hahn & Orrock 2016). Choice assays are 314 often conducted to test the candidate mechanism of AE (Tahvanainen & Root 1972; 315White & Whitham 2000; Rand 2003; Hahn & Orrock 2016). Several studies 316 documented that the feeding preferences of herbivores depend on the presence or 317 relative frequencies of multiple resource types (Chandra & Williams 1983; Cottam 318 1985; Behmer et al. 2001; Bergvall & Leimar 2005; Janz et al. 2005), but evidence 319 regarding such a frequency-dependent herbivory on natural variation in a plant defense 320trait is limited.

321 My collaborators and I have conducted a four-way choice experiment 322 manipulating frequencies of hairy and glabrous leaf discs (Fig. 2b: Sato et al. 2014). 323 Adults of the brassica leaf beetle *P. brassicae* avoided hairy leaves when hairy leaves 324 were rare, whereas the adults did not show a feeding preference when hairy leaves 325 became abundant (Sato et al. 2014). The adult preference for glabrous leaves was also 326 observed in a binary choice experiment using a gl1 mutant of the model plant species 327 Arabidopsis thaliana (Sato 2016). This result from a single-gene mutant provides strong 328 proof for the resistance function of the GL1 gene against P. brassicae. Furthermore, 329 when adult beetles were starved or preconditioned with glabrous diets, they still 330 preferred glabrous plants (Sato & Kudoh 2016b). Contrarily, when adults were 331 preconditioned with hairy diets, this treatment weakened their feeding preference for 332glabrous diets (Sato & Kudoh 2016b). The circumstantial evidence suggests that 333 sequential experience may trigger a non-random response of adult beetles to hairy and 334 glabrous diets. While adult beetles fed on both hairy and glabrous leaves at similar 335 levels under no-choice conditions, larvae fed on hairy diets grew slower than those on 336 glabrous diets. This indicates that hairy diets are suboptimal for P. brassicae.

337 The foraging behaviors of *P. brassicae* were then modeled to reveal the 338 mechanism underlying the pattern of leaf damage to hairy and glabrous plants. Given 339 that *P. brassicae* is flightless and incapable of selecting plants among patches, the 340 optimal diet choice (Charnov 1976) is assumed to be a mechanism by which rare 341 suboptimal diets (i.e., hairy leaves) are less consumed under the small enclosed 342condition. My collaborators and I modified the original model of optimal diet choice to 343 consider imperfect host recognition by P. brassicae, and then fitted the model to the 344 laboratory experiment data of leaf damage under the contrasting frequency of hairy and 345glabrous plants (Sato et al. 2017). As a result of the parameter estimation, it was 346 estimated that decreased energy intake and increased handling time for the leaf beetles 347 accounted for the frequency-dependent damage on hairy plants (Fig. 2c). Hairy plants 348 were less damaged when glabrous plants became abundant, indicating associational 349 resistance for hairy plants. Contrarily, damages to glabrous plants weakly depended on 350the frequency of two plant morphs. These findings indicate that the optimal diet choice 351by P. brassicae can be a determinant of the pattern of AE between hairy and glabrous 352plants.

353

2-2. Manipulative experiment showing a rarity advantage in plant defense and fitness
Even if herbivores feed on hairy and glabrous plants in a frequency-dependent manner,
it is still unknown whether this candidate mechanism leads to frequency-dependent
selection on the trichome dimorphism. Specifically with regard to the brassica leaf
beetle *P. brassicae*, it seems unlikely that herbivore preference alone generates negative
frequency-dependent selection, as they did not avoid rare glabrous diets (Sato *et al.*2014: Fig. 3a). However, game theoretical models suggest that, if plants have defense

traits at the cost of their growth or fecundity, the benefit from associational effects
allows defended and undefended plants to coexist (Augner *et al.* 1991). This hypothesis
was tested using a laboratory and semi-field experiment manipulating the morph
frequency and presence/absence of *P. brassicae* (Fig. 2b; Sato & Kudoh 2016b; Sato &
Kudoh 2017b).

366 Figure 3 shows a schematic explanation of a mechanism by which conspecific 367 AE could be a mechanism for the maintenance of trichome dimorphism. Consistent with 368 the feeding preference assay (Sato et al. 2014), hairy plants were less damaged than 369 glabrous plants only when the hairy ones were rare (Fig. 3a: Sato & Kudoh 2016b; Sato 370 & Kudoh 2017b). This explains the rarity advantage for hairy plants in defense and 371 fitness (Fig. 3a, c). At the end of the experiments, both the hairy and glabrous plants had 372 a rarity advantage in fitness in the presence of beetles (Fig. 3c: Sato & Kudoh 2016b; 373 Sato & Kudoh 2017b), while in the absence of herbivory the glabrous plants had a 374higher growth and fecundity (Fig. 3b: Sato & Kudoh 2016b; Sato & Kudoh 2017b). 375 Because the benefit of associational effects to hairy plants no longer exists under the 376 hairy-abundant condition (Fig. 3a), the cost of defense can be detected when hairy 377 plants become abundant (Fig. 3c). Of note, these fitness consequences were consistently 378 observed for both sexual and asexual reproduction, namely the number of flowers and 379 clones, of A. halleri (Sato & Kudoh 2017b). These experimental findings indicate that, 380 together with the defense-growth tradeoff (Fig. 3c), conspecific AE against the leaf 381 beetle (Fig. 3a) lead to negative frequency-dependent selection between hairy and 382glabrous plants (Fig. 3b) (Sato & Kudoh 2017b).

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384 2-3. Field patterns of herbivory and polymorphism dynamics

385 The set of experiments illustrated conspecific AE that can maintain the trichome 386 dimorphism of A. halleri, but it is still unknown whether these experimental findings 387 represent AE in the wild. Next, field observations on plant damage and demography 388 were conducted in a natural population in which the brassica leaf beetle *P. brassicae* 389 was a major herbivorous insect on A. halleri (Fig. 2b). Because P. brassicae is flightless 390 and the A. halleri population has a fragmented structure, AE is assumed to occur on a 391 small spatial scale. Thus, my collaborators and I tracked the aggregated structure of the 392plants (referred to hereafter as "patch") for four years in the field.

393 Hairy plants received less damage as the frequency of glabrous plants increased 394 in a series of plant patches (Sato et al. 2014; Sato & Kudoh 2017b). In contrast, leaf 395 damage on glabrous plants showed no significant and consistent relationship with the 396 frequency of glabrous plants in a patch (Sato *et al.* 2014; Sato & Kudoh 2017b). This 397 pattern of leaf damage was consistent with the results of a feeding assay. It was also 398 noteworthy that the relative fitness advantage of hairy plants as shown by flower 399 production in the presence of beetles increased with increasing frequencies of glabrous 400 plants (Sato & Kudoh 2017b). This pattern in the flower production agrees with the 401 manipulative experiment detecting negative frequency-dependent selection only in the 402presence of *P. brassicae* (Fig. 3c). These findings support the relevance of 403 frequency-dependent herbivory and selection in the natural population. 404 A multi-year field survey also found that rarer morphs increased in frequency 405in a series of plant patches over time (Sato & Kudoh 2017b). Statistical modeling was 406 used to reveal the mechanisms underlying the temporal dynamics of morph frequency in

407 the field (Sato *et al.* 2017). Based on the aforementioned theory of optimal diet choice,

408 my collaborators and I have modelled the effects of local-scale herbivory and spatial

409 structure on plant demography and then fitted the model to the observed changes in the 410 number of hairy and glabrous plants over four years. Based on a model selection 411 procedure, it was revealed that optimal diet choice by *P. brassicae* contributes to the 412polymorphism dynamics observed in the field (Sato et al. 2017). The estimated 413 magnitude of interpatch dispersal suggested that long-distance seed dispersal was rare 414 and clonal reproduction largely contributed to the plant population growth. According to 415the estimated parameters of antiherbivore defense, rarer morphs had higher fitness and 416 increased in frequency over time when the herbivore diet choice and cost of plant 417 defense were both involved (Fig. 2c). This joint approach using the optimality modeling 418 and field data strongly supports the hypothesis that AE against the leaf beetle promotes 419 the maintenance of the trichome dimorphism via negative frequency-dependent 420 selection.

421In contrast to the constant density in the manipulative experiment (Fig. 3), not 422only frequency but also the number of plants varies among field patches. However, the effects of plant density on damage were much less significant than that of frequency, 423 424 and its magnitude and direction were inconsistent among the study years (Sato et al. 4252014; Sato & Kudoh 2017b). The resource concentration or delusion of A. halleri for P. 426 brassicae was tested in more detail using the field data on the leaf damage and number 427 of A. halleri plants (Sato et al. 2017). If these two effects of plant density occur, there is 428 a non-linear relationship between the total amount of herbivory load and the total 429 number of plants within patches (Hambäck et al. 2014). In A. halleri, however, the total 430 amount of leaf damage increased linearly with the total number of plants in a patch 431 (Sato *et al.* 2017), presumably because the plant density in the field was too low to 432 reach a peak of resource loss by herbivory. Thus, at least for the temporal scales

433 observed, the field survey could not detect a significant role of plant density in AE434 against the leaf beetles.

435 Overall, the multiple lines of evidence have shown that conspecific AE 436 mediated by *P. brassicae* lead to negative frequency-dependent selection and thereby 437 maintain the trichome dimorphism in a natural A. halleri population. If hairy plants are 438 rare, they can escape from leaf beetles and increase in frequency. On the other hand, if 439glabrous plants become rare, they outcompete hairy plants without any costs of 440 trichomes. Consequently, this negative feedback loop prevents rare morphs from going 441 extinct (Fig. 4). To date, negative frequency-dependent selection has been reported for 442plant species interacting with pollinators (Gigord et al. 2001) and pathogens 443 (Antonovics & Ellstrand 1984; Brunet & Munht 2000), but it has not yet been 444 demonstrated in plant defense against herbivores. The case study on A. halleri-P. 445*brassicae* interaction provides the first example to illustrate that conspecific AE drive 446 negative frequency-dependent selection on an antiherbivore defense trait. 447

## 448 **3. Perspective**

449 Genetic polymorphisms are commonly observed in plant defense traits (Hughes 1991;

450 Elle et al. 1999; Wise et al. 2009; Bernhardsson et al. 2013) and not limited to

451 Arabidopsis trichomes. For example, the genetic basis of cyanogenesis in white clover

452 *Trifolium repens* (Fabaceae) is well studied (Hughes 1991). Theory related to

453 herbivore-mediated frequency-dependent selection was applied to geographical

454 variation in a cyanogenesis polymorphism (Till-Bottraud & Gouyon 1992), but the

455 existence of frequency-dependent selection has not been tested experimentally.

456 Dimorphism of the glandular (sticky) and non-glandular (velvety) Datura wrightii

(Solanaceae) is inherited in the Mendelian fashion (van Dam et al. 1999). This chemical 457458defense utilizing glandular exudates is costly for plant fitness (Elle et al. 1999), but it 459remains unknown why these sticky and velvety plants can coexist (Elle & Hare 2000). 460 The genetic basis of glucosinolates, a major secondary metabolite of Brassicales, has 461 been intensively studied. Genetic polymorphisms are observed in physiological 462 pathways in glucosinolate biosynthesis and result in diverse profiles among natural 463 Arabidopsis accessions (Chan et al. 2010). Conspecific AE have not yet been examined 464 in these defense traits but could provide insights into mechanisms for the maintenance 465of antiherbivore defense polymorphism within a plant population.

466 We should note here that not only AE but also other factors may shape genetic 467 variation in antiherbivore defenses. It has been hypothesized that genetic variation may 468 be maintained by spatiotemporal fluctuation in selection pressure and stochastic gene 469 flow among populations (e.g., Levene 1953; Turelli et al. 2001; but see Spichtig & 470Kawecki 2004). The present paper has focused on local-scale AE so far, but to what 471spatial and temporal scales can AE contribute to the maintenance of polymorphism in 472 plant defense? In the final chapter, I discuss a spatial and temporal perspective of 473 conspecific AE and the maintenance of plant defense polymorphism. Concerning the 474spatial scale, several researchers discussed interspecific AE at a landscape scale 475(reviewed by Barbosa et al. 2009). The geographic mosaic of coevolution (Thompson 4762005) proposes a landscape perspective of plant-insect interactions, and this spatial 477 point of view is applied to geographic variation in plant defense (e.g., Berenbaum & 478Zangerl 1998; Zangerl & Berenbaum 2003; Toju et al. 2011; Bernhardsson et al. 2013). 479 In particular to temporal dynamics, the concept of eco-evolutionary dynamics has 480 recently suggested interactions between plant genetic variation and herbivore abundance 481 (Turcotte 2011; Turcotte *et al.* 2013; Agrawal *et al.* 2013). Because not only frequency482 but also density-dependent selection is a key aspect of AE (Table 1), conspecific AE
483 may provide insights into how rapid evolution in plant defenses affects the population
484 dynamics of plants and herbivores.

485

486 3-1. Can AE create a source population for geographical variation in plant defense? 487 Existing theory suggests that negative frequency-dependent selection contributes to 488 geographic cline in a polymorphism, as it makes morph frequency intermediate rather 489 than expected by the results of disruptive selection alone (Clarke 1966; Endler 1977; 490 Till-Bottraud & Gouyon 1992). Till-Bottraud & Gouyon (1992) applied this prediction 491 to explain a cyanogenesis polymorphism along an altitudinal cline in white clover. 492 However, herbivory regimes usually have more complex spatial patterns than a cline, 493 involving differences in herbivore species (Hare & Elle 2002) and the presence of other 494 host plants (Zangerl & Berenbaum 2003). Thus, plant-herbivore interactions are 495 expected to result in complex geographic variation rather than a simple cline in a plant 496 defense polymorphism.

497 The geographic mosaic of coevolution proposes a conceptual framework for 498 investigating the complex metapopulation structure of local adaptation/maladaptation 499 (Thompson 2005). Specifically, the theory of geographic mosaic of coevolving 500polymorphism hypothesizes that disruptive and balancing selection jointly shape 501complex patterns of host defense polymorphism (Thompson 2005). In plant-herbivore 502interactions, Berenbaum and Zangerl (1998) presumed that intense attacks by parsnip 503webworm *Depressaria pastinacella* on abundant chemotypes of a wild parsnip 504 Pastinaca sativa (Apiaceae) led to negative frequency-dependent selection and thus

favored the intermediate chemotype frequency. Given that AE should occur among
individual plants in close vicinity, this may alter spatial patterns of plant defense only at
a local scale. More generally, the metapopulation framework assumes the existence of
source and sink populations of local adaptation (Thompson 2005). When AE favors
polymorphism within a local population, this polymorphic population could be a source
of polymorphism on a metapopulation scale via gene flow (Fig. 5).

511Indeed, herbivore regimes and frequencies of hairy and glabrous plants vary 512among natural populations of Arabidopsis halleri subsp. gemmifera, where AE do not 513 occur against all herbivorous insects on A. halleri. Sato and Kudoh (2015) found that 514AE were detectable when A. halleri interacted with slow-moving herbivores such as P. 515*brassicae*. AE tend to be obscured when plants are attacked by flying herbivores such as 516the green-veined white butterfly *Pieris napi* and cabbage sawfly *Athalia infumata*, 517although trichomes themselves can prevent herbivory by these herbivores (Sato & 518Kudoh 2015). Field surveys revealed that frequencies of glabrous plants declined as 519herbivory pressure became intense (Sato & Kudoh 2017a). Hairy-monomorphic 520 populations were discovered where white butterflies were frequently observed. Despite 521experiencing the severest herbivory pressure, however, a population where *P. brassicae* 522was predominant still had an intermediate morph frequency (Sato & Kudoh 2017a). 523Furthermore, a proximate population was also polymorphic despite the prevalence of 524white butterflies (Sato & Kudoh 2017a). These observations support the hypothesis that 525herbivore-mediated frequency-dependent selection sustains polymorphism, which can 526be a source of genetic variation in proximate populations (Fig. 5b). Further questions 527 remain about how conspecific AE contributes to the maintenance of polymorphism at 528the metapopulation level.

529

530 3-2. Do AE affect population dynamics via the maintenance of defense polymorphism? 531A growing number of studies have shown that herbivores drive the evolution of plant 532defense on ecological time scales (Zangerl & Berenbaum 2008; Agrawal et al. 2012; 533Fukano & Yahara 2012; Sakata et al. 2014). Recent studies have also shown that such 534evolutionary changes in a host plant species alter the population dynamics and 535community structures of herbivores (Crutsinger et al. 2006; Johnson et al. 2006; 536Turcotte et al. 2011; Agrawal et al. 2013), which may in turn affect plant density. Now 537 that conspecific AE is known to maintain polymorphism in plant defense, it is possible 538that such a sustained polymorphism affects eco-evolutionary dynamics in 539plant-herbivore interactions. In the context of AE, Underwood et al. (2014) advocate 540that manipulating both the density and frequency of multiple plant species is important 541for complete understanding of the long-term coexistence of multiple plant species under 542AE (as introduced in Section 1-2.). When this notion is applied to polymorphisms 543within a plant species, AE may play an important role not only through frequency- but 544 also through density-dependent selection in plant defense. 545Once the mechanisms of frequency- and/or density-dependent selection are 546experimentally shown, it is possible to estimate the probability of coexistence from the

experimental data (e.g., Damgaard 1998; Inouye 2001; Miller & Rudgers 2014) or to
predict temporal dynamics in the field (Kim *et al.* 2013; Le Rouzic *et al.* 2015) based on

a given theoretical model. This modeling approach, combined with empirical data, will

be a powerful tool to reveal how AE contribute to eco-evolutionary dynamics in

551 plant-herbivore interactions. For example, Sato et al. (2017) took such a modeling

approach to A. halleri–P. brassicae interactions although the modeling was limited in

predicting long-term interactions between the evolution of plant defenses and herbivore population dynamics. Specifically, the magnitude of herbivory in the current year was not clearly correlated with that in the previous year or with the frequency of hairy plants (Sato *et al.* 2017). If long-term data on herbivory and population dynamics are obtained, the modeling approach might enable us to predict how the existence of polymorphism affects herbivory pressure and in turn alters the morph frequency within a plant species.

559

### 560 3-3. Concluding remarks

561 Associational effects, or neighbor effects, are remarkable phenomena in plant–herbivore

562 interaction, but evidence for conspecific AE is still limited. By focusing on polymorphic

563 defense traits within a plant species, the present review highlights the potential

564 importance of conspecific AE in maintaining polymorphism in an antiherbivore defense

565 trait. The multiple lines of evidence from *P. brassicae–A. halleri* interaction show that

566 conspecific AE maintains antiherbivore defense polymorphism via negative

567 frequency-dependent selection. Future studies should focus on the stabilizing role of AE

568 in genetic variation in plant defense, as diverse plant species possess considerable

569 variation in terms of antiherbivore defense traits.

570



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Factor	Article	Study syst	Neighborhood composition		
		Plant (or Resource)	Herbivore (or Consumer)		
Density	Rand (2003)	Atriplex patula and Salicornia europaea	Polyphagous beetle Erynephala maritima	Presence/absence of the other species with low/high density	
	Russell & Louda (2005)	a (2005) Cirsium canescens and C. undulatum Florivorous weevil Rhinocyllic conicus		Density of the other species in natural populations	
Frequency	Bergvall et al. (2006)	Low- and high-tannin food	Fallow deer Dama dama	Rare or abundant type A against B (= 2 frequency conditions)	
	Wise <i>et al.</i> (2009)	Candy-cane and erect-stem genotype of Solidago altissima	Oligophagous gall-fly Eurosta solidaginis	Rare or abundant type A against B (= 2 frequency conditions)	
	Sato & Kudoh (2016b)	Hairy and glabrous genotype of <i>Arabidopsis halleri</i> subsp. <i>gemmifera</i>	Oligophagous beetle <i>Phaedon</i> brassicae	Rare or abundant type A against B (= 2 frequency conditions)	
	Garrido et al. (2016)	Resistant and tolerent genotype of <i>Datura stramonium</i>	Oligophagous beetle <i>Lema</i> daturaphila	Rare, even, and abundant type A against type B (= 3 frequency conditions)	
Both	Kim & Underwood (2015)	Solidago altissima and Solanum carolinense	(incl. multiple species)	Response surface design of 4 frequency and 4 density conditions	
	Hahn & Orrock (2016)	Solidago nemoralis and S. odora	Grasshoppers	Rare, even, and abundant type A against type B (= 3 frequency conditions) at high/low densities.	
	Verschut et al. (2016)	Balsamic vinegar solutions	Drosophila melanogaster	Different resource concentration (2 levels) with 3 or 5 frequency conditions	

# **Table 1.** Examples of associational effects that cause frequency- and/or density-dependent herbivory between plant species/genotypes.

914	Figure legends
915	Figure 1. Stepwise approach to reveal causes and consequences of associational effects
916	(AE) on plant coexistence. The reference indicates studies comprising a focal approach.
917	Details of the selected references are shown in Appendix.
918	
919	Figure 2. Comprehensive test for the maintenance of the trichome dimorphism of
920	Arabidopsis halleri subsp. gemmifera mediated by associational effects against a leaf
921	beetle Phaedon brassicae. (a) Photographs of plants and insects; (b) Empirical
922	approaches from laboratory experiments to field observation; (c) Modeling herbivory,
923	plant fitness, and polymorphism dynamics. Insets in the panel (c) are modified from
924	Sato <i>et al.</i> (2017).
925	
926	Figure 3. Mechanism through which associational effects and a cost of defense jointly
927	cause negative frequency-dependent selection between hairy and glabrous plants. Panels
928	(a), (b) and (c) represent differences in herbivory, intrinsic fitness, and realized fitness
929	between hair and glabrous plants, respectively. $H > G$ and $H < G$ indicates
930	hairy-abundant and glabrous-abundant condition, respectively.
931	
932	Figure 4. Evolutionary dynamics wherein the trichome dimorphism of Arabidopsis
933	halleri subsp. gemmifera can be maintained by the herbivore-mediated
934	frequency-dependent selection. Frequency-dependent feeding preference by herbivores
935	and plant defense-growth tradeoff results in a negative feedback loop that allows rarer
936	plant morphs to invade into a population.
937	
938	Figure 5. Specific hypothesis for how local-scale associational effects (AE) contribute
939	to geographic variation in plant defense via the maintenance of source population in a
940	polymorphism. (a) Relationship between the herbivory pressure and morph-frequency
941	in the presence (solid line) and absence (dashed line) of AE. (b) Contribution of a
942	source polymorphic population created by AE to the maintenance of polymorphism at a
943	metapopulation scale. Grey and white square represents the region of intense and
944	moderate herbivory, respectively. White arrows represent gene flow from a source to
945	nearby populations.

914

## Mechanism

# Approach 1: Identifying herbivore behaviors and plant traits

- Behavioral assay for herbivores<sup>[1,2,3]</sup> - Measurement of key plant traits<sup>[3,4,5,8]</sup>

# Approach 2: Manipulative experiments demonstrating AE

Manipulating

- Neighborhood composition in plants - Presence/absence of herbivores<sup>[5,6,7]</sup>

# Approach 3: Field pattern of herbivory and plant population dynamics

Multi-year field census on

Patterns of the damage and herbivore abundance on plants<sup>[1,8]</sup>
 Plant fitness<sup>[5,9]</sup> and population growth<sup>[7]</sup>

## Pattern

	[1] Tahvanainen & Root (1972); [2] White & Whitham (2000); [3] Bergvall et al. (2006); [4] Barton & Browers [5] Agrawal (2004); [6] Band (2003); [7] Callaway et al. (2005); [8] Bussell & Loude (2005); [9] Karban & Ma						
947	[5] Agrawar(2004), [6] Rand (2005), [7] Ganaway et al. (2005), [6] Russen & Louda (2005), [9] Randan & Maron (2002)						
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Appendix. Selected examples of associational effects in plant-herbivore interaction. Study system (plants and herbivores), design (experimental design and spatial scales, reciprocal data or not), and results (measurement, outcome, and putative mechanism) are summarized.

Reference	Study system			Study design	Results				
	Herbivore	Plant species	Plant traits	Experimental design	Scale	Reciprocal or not	Measurement	Outcome	Mechanism
Tahvanainen & Root (1972)	A specialist flea beetle ( <i>Phyllotreta</i> cruciferae )	<i>Brassica oleracea</i> and tomato or tobacco	Odor (shown by a choice assay)	Monoculture of <i>B. oleracea</i> and polyculture of three species. The tomato and tobacco was added to the equal density of <i>B. oleracea</i> in the polyculture.	Within-patch t	NA for tomato and tobacco	The number of beetles per plant and plant biomass for <i>B. oleracea</i>	AR: The number of beetles per plant was higher, and plant biomass decreased under the monoculture.	Odor cues for herbivores
Rausher (1981)	A specialist butterfly ( <i>Battus philenor</i> )	Aristolochia reticulata and others species in near vegetation	Unidentified	0.5 m-radius vegetation surrounding an <i>A. reticulata</i> individual was removed. Non-removed patches were assigned to the control.	Within-patch	NA for counterparts	Oviposition per A. <i>reticulata</i> plant	AR: Isolated plants suffer more oviposition, butterflies spend more time in the isolated patch.	Searching effeciency of herbivores
Risch (1981)	Mono- Oligo-, Polyphagous beetles (6 species)	Corn (Zea mays), beans (Phaseolus vulgaris), and squash (Cucurbita maxima)	Unidentified	Intercropping one crops into another; monoculture, diculture triculture for 3 species (= 7 conditions)	Within-patch	Reciprocal	No of beetles per plant	AR: The beetle abundance reduced under polycultic conditions except for a squash- bean diculture	Resource concentration (plant density)
Hay (1986)	Generalist fishes	Multiple species of Marine algae	Unidentified	Solitary or paired culture in a closed cage	Within-patch	Reciprocal	Within-patch: predation risk by fishes	AR for palatable algae to generalist fishes	Physical attachments to unpalatable algae
White & Whitham (2000)	Fall cankerworm (Alsophila pometaria).	Cotton ( <i>Populus</i> angustifolia × P. fremontii), and box elder (Acer negundo)	Unknown, (but a choice test showed that the cankerworm prefer to the box elder).	Transplanting juvenile cottons to three conditions: Nearby box elder, Nearby matured cotton, Stand alone.	Within-patch	NA for the box elder	Larvae density per shoot of the juvenile cotton	AS: Cotton suffered from the higher density nearby a heterospecific counterpart than nearby conspecifics or stand alone.	Herbivore dispersal from palatable to unpalatable plants
Karban & Maron (2002)	Generalist grass- hoppers (incl. multiple species)	Sagebrush (Artemisia tridentata) and tobacco (Nicotiana attenuata)	Volatile chemical	Tobacco was transferred near damaged or intact sagebrush.	Within-patch	NA for Sagebrush	The number of flowers and capsules per tobacco	AR: tobacco produced more flowers and capsules nearby damaged sagebrush.	Volatile communication between plants
Hambäck <i>et al.</i> (2000)	A specialist leaf beetle (Galerucella calmariensis)	An aromatic shrub ( <i>Myrica gale</i> ) and a perrenial herb ( <i>Lythrum salicaria</i> )	Unidentified	Stand-alone or within- <i>M. gale</i> patch condition for <i>L. salicaria</i>	Within-patch	NA for <i>M. gale</i>	Oviposition, leaf damage, seed production per individual <i>L.</i> <i>salicaria</i> .	AR: <i>L. salicaria</i> incurred less oviposition and damage, and produce more seeds nearby <i>M. gale</i>	Odor or vidual cypsis. Predator abundance did not differ between the two conditions.
Rand (2003)	A generalist leaf beetle ( <i>Erynephalam</i> <i>aritima</i> )	Two forbs, <i>Atriplex</i> <i>patula</i> var. <i>hastata</i> and <i>Salicorniae</i> <i>uropaea</i>	Unidentified (but feeding preference was shown by a choice test).	The presence/absence of herbivores for 3 conditions: <i>Atriplex</i> alone with low density, <i>Atriplex</i> alone with high density, <i>Atriplex</i> with <i>Salicornia</i>	Within-patch	NA for Salicornia	Leaf damage, the number of larvae, and survival (%) of plants	AS: <i>Atriplex</i> suffered more leaf damage and less survival nearby <i>Salicornia</i>	Diet-mixing by generalist herbivore
Agrawal (2004)	A specialist beetle (Tetraopes tetraophthalmus)	a milkweed ( <i>Asclepias</i> <i>syriaca</i> ) and a grass ( <i>Liriomyza</i> <i>asclepiadis</i> )	Trichome, Latex, Nitrogen contents	Grasses or beetles were added to a potted milkweed: 4 treatments of Control, Grass+, Beetle+, and Grass&Beetle+. Grass alone, Grass&Milk&Beetle, Grass&Milk was also set.	Within-patch	Grass biomass data: Grass alone = Grass&Milk&Beet le > Grass&Milk	Milkweed biomass, fruit production, fruit mass; and grass biomass	AS: milkweed biomass and fecundity was Control ≒ Grass+ > Beetle+ > Grass&Beetle+.	s Dispersal from primary host plants to secondary ones.

Callaway <i>et al.</i> (2005)	Mammals	Four unpalatable herbs ( <i>Cirsium</i> , <i>Veratrum</i> , <i>Luzula</i> , <i>Alchemilla</i> ) and two palatable herbs ( <i>Anthoxanthum</i> , <i>Phleum</i> )	Spines (for <i>Cirsium</i> ) or toxin (for <i>Veratrum</i> )	Four focal species $\times$ within/outside grazing fence $\times$ exclusion of <i>Cirsium</i> or <i>Veratrum</i> = 24 treatments	Within-patch	NA for <i>Cirsium</i> and <i>Veratrum</i>	Annual growth rates of four plant species.	A P r (
Russell & Louda (2005)	Florivorous weevil, Rhinocyllus conicus	Two congeneric natives, <i>Cirsium</i> <i>canescens</i> and <i>C</i> . <i>undulatum</i>	Plant phenology	> 10-yrs field census in 13 sites where the two plant species co-occur	Within-site	NA for <i>C</i> . canescens	Proportion of weevils and their eggs on <i>C</i> . <i>undulatum</i> .	A v c c
Bergvall et al. (2006)	Fallow deer ( <i>Dama dama</i> )	Unknown. Synthetic food containing different tannin levels	Chemical (Tannin)	Bad-patch and good-patch contains 7:1 and 1:7 of high:low tannin buckets, respectively. Single- and group-foraging of deers were also tested.	Within and between patch	Reciprocal	Log-consumption per bucket (g).	A ti p t
Barton & Bowers (2006)	None (only defense phenotypes were evaluated)	Interspecific: two congeners, <i>Plantago</i> <i>lanceolata</i> and <i>P.</i> <i>major</i>	Chemical (Iridoid glucoside)	2 species $\times$ (conspecific, heterospecific, no neighbor) $\times$ 2 harvesting time points = 12 conditions	Within-patch	Reciprocal	Within-patch: % Dry Weight of Iridoid Glycosides	A s p c c
Le Guigo et al. (2012)	A specialist aphid ( <i>Brevicoryne</i> <i>brassicae</i> ) and generalist aphid ( <i>Myzus persicae</i> )	Three <i>Brassica</i> species and tomato ( <i>Solanum</i> <i>lycopersicum</i> )	Chemical (glucosinolate?)	A <i>Brassicae oleracea</i> individual was surrounded by 6 conspecific or heterospecific individuals (= 7 treatments in total).	Within-patch	Reciprocal	The number of aphids on a focal <i>B</i> . <i>oleracea</i> , compared between conspecific and interspecific conditions	/ v
Castagneyrol et al. (2013)	Leaf-chewers (beetles, caterpillars, and grasshoppers) or Leaf-miners (incl. 5 species)	Betula pendula , Quercus robur , Quercus pyrenaica , Quercus ilex , Pinus pinaster	Plant height, and leaf area, N and P contents, leaf life span	31 possible mono- and poly-culture combinations of 1~5 plant species	Within/Betw een patch	Relative values of two species (for plant height data)	Leaf area loss (by chewers) and leaf- miner abundance at plot and individual scale.	4 1
Kim & Underwood (2015)	3 species of specialists (for <i>S</i> . <i>carolinense</i> ), and 3 species of generalist herbivores	Solanum carolinense, Solidago altissima	Unidentified	Response surface design (1, 6, 12, 18 total plant number with 4 frequencies of <i>S. altissima</i> )	Within-patch	Reciprocal (shown in the supplement)	Leaf damage (%) of individual plants	P

Abbreviation: AR; Associational resistance, AS; Associational susceptibility, NA; No information available

AR: increased growth rate of two palatable species, no increase of growth rate of unpalatable two species nearby *Cirsium* and *Veratrum* 

AR for *C. undulatum* : The proportion of weevils and eggs was negatively correlated with the number of *C. canescens.* 

AS for low-tannin buckets: More consumption on low-tannin buckets in the bad-patch than those in the goodpatch. AR for high-tannin buckets: Less consumption on high-tannin buckets in the good-patch than those in the bad-AS for *P. lanceolata* ?: *P. lanceolata* surrounded by heterospecific plants produced less defensive chemicals compared to those surrounded by conspecifics.

AS when a herbivore is specialist; AR when a herbivore is generalist.

AR against leaf-miners: no AR against leaf-chewers

AS for *S. carolinense and* damage was positively density-dependent.

avoidance by grazers to plant spines and toxins.

Phenological syncronicity of flowering between the two species.

within- and amongpatch foraging behavior of mammalian herbivores.

Plant trait change via direct competition between plants.

Chemical attraction or repulsion

Variation in leaf traits and apparency (plant height)

Unknown (but those depending on the plant density and frequency)