Research Article

Use of Host-Plant Trait Space by Phytophagous Insects during Host-Associated Differentiation: The Gape-and-Pinch Model

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Ecological speciation via host shifting has contributed to the astonishing diversity of phytophagous insects. The importance for host shifting of trait differences between alternative host plants is well established, but much less is known about trait variation within hosts. I outline a conceptual model, the "gape-and-pin ch" (GAP) model, of insect response to host-plant trait variation during host shifting and host-associated differentiation. I offer four hypotheses about insect use of plant trait variation on two alternative hosts, for insects at different stages of host-associated differentiation. Collectively, these hypotheses suggest that insect responses to plant trait variation can favour or oppose critical steps in herbivore diversification. I provide statistical tools for analysing herbivore trait-space use, demonstrate their application for four herbivores of the goldenrods Solidago altissima and S. gigantea, and discuss their broader potential to advance our understanding of diet breadth and ecological speciation in phytophagous insects.

1. Introduction

The insects have long been held up as providing spectacular examples of rapid diversification and high standing diversity (e.g., [1–3]). Among insects, phytophagous clades often undergo dramatic radiations [4], and phytophagous lineages tend to be more diverse than their nonphytophagous sisters [5, 6]. One likely driver of diversification among phytophagous insects is their tendency to specialize on host-plant species or organs [7–10] and to diversify via host or organ shifts followed by host-associated differentiation (HAD), the evolution of new specialist races or species [9, 11–14]. Because many cases of HAD appear to have proceeded in sympathy [15], a great deal of theoretical and empirical work has focused on understanding ways in which adaptation to different host plants can impose disruptive selection on nascent specialist forms and also reduce gene flow (or permit differentiation in the face of gene flow) between those forms [14–16]. Phytophagous insects, along with parasitoids [17], freshwater fishes [18], seed-eating birds [19], and habitat-specialist plants [20] and lizards [21] have therefore been central to the development of ideas about ecological speciation [22].

A common theme among case studies of ecological speciation is the existence of two alternative niches—microhabitats, resources, reproductive strategies, and so forth—that can be exploited by individuals of a single species, with the potential for disruptive selection to operate between the alternative niches. For phytophagous insects, the alternative niches are a pair of host plant species (or organs). One commonly imagines an evolutionary sequence beginning with an insect exploiting only one of the two alternative hosts. Perhaps via host-choice errors, some individuals occasionally attack individuals of the second host, and if fitness penalties for doing so are not too severe, a host shift occurs and the insect begins to exploit both alternative hosts. (Description of these events as “errors” is standard in the plant-insect literature, but of course this usage is teleological shorthand and can conceal interesting biology. For instance, it might be that genotypes with strong enough host preferences to avoid “errors” would also show costly rejection of some suitable hosts; in this case, the occurrence of host-choice “errors” is simply an adaptive compromise. Nonetheless, for simplicity I retain the standard usage here). Disruptive selection can now begin to favour genotypes better adapted to each alternative host. If reproductive isolation arises between nascent forms,
then ecological speciation can proceed, and a single (perhaps polymorphic) generalist is replaced with a pair of host-specialist races or species. Because reproductive isolation is expected to take some time to evolve, if it can evolve at all, different insects exploiting a pair of alternative host plants are expected to fall on a continuum from generalists to nascent, poorly differentiated host forms to distinct host-specialist sister species [13, 23]. There will be analogous continua for ecological speciation across other kinds of alternative niches, for instance, in parasitoids speciating across hosts or fish across depth niches (e.g., [17, 24]).

The process of HAD in phytophagous insects has been widely discussed, both in general [14] and in the context of a few well-studied model systems (e.g., apple maggot fly [25, 26], goldenrod ball-gall fly [27, 28]). Perhaps unsurprisingly, nearly all studies of HAD have emphasized insect responses to differences in plant traits between the alternative hosts, while downplaying variation in plant traits among individuals within each host. Such an interspecific perspective is obviously appropriate for studies using population-genetic tools to detect host-associated forms and reconstruct their history (e.g., [13, 29–31]), but it is also near universal in studies discussing ecological mechanisms by which host shifts and HAD proceed (e.g., [25, 26, 28, 32–38]). An alternative approach would explicitly recognize within-species variation in host-plant traits and consider possible roles for such variation in favouring or impeding host shifting and HAD. This approach has yet to be applied in earnest to any system, but intriguing hints at its usefulness appear in the literature for the goldenrod ball-gall fly, *Eurosta solidaginis*, and its races on the goldenrods *Solidago altissima* and *S. gigantea*. For example, *Eurosta* of the *S. altissima* race prefer the largest ramets of their host [39], and since *S. gigantea* plants tend to be shorter when *Eurosta* oviposits [40], this preference might discourage host-choice errors by *altissima* flies. In contrast, if *gigantea* flies similarly prefer taller ramets, they could be susceptible to host-choice errors (although *gigantea* flies’ preferences have not been assessed, and neither hypothesis raised here appears to have been tested). Work on the phenology of insect emergence and host-plant growth has similar implications. *Eurosta* adults emerge from *S. gigantea* earlier than from *S. altissima* [41], and this pattern is correlated with availability of rapidly growing ramets of each host to be attacked [40]. Thus, individual *S. altissima* ramets with earlier phenology, or *S. gigantea* ramets with later phenology, might be more likely to be attacked by the "wrong" host race. There is geographic variation in the abundance of such intermediate-phenology ramets, and How et al. [40] suggested that host shifts might be more easily initiated where host phenology overlaps more extensively.

Even for *Eurosta*, however, there are few plant traits for which insect responses have been studied on both alternative hosts, and so we know little about how insect responses to plant trait variation might relate to the ecology of host shifting and HAD. Furthermore, there is no system for which we can compare insect responses to plant trait variation for a set of insects attacking the same plants but differing in stage of host shifting and HAD. I outline here a conceptual model of host trait-space use during host shifting and HAD, along with a statistical approach for investigating trait-space use in phytophagous insects. I suggest hypotheses for temporal changes in host trait-space use over evolutionary time, from initial host-choice errors through to the independent evolution of a pair of well-isolated host-specialist sibling species. I call the overall model the “gape-and-pinch,” or “GAP,” model of trait-space use (the reason for this name will be apparent after the model is described). While I outline the model for plants and phytophagous insects, it will apply to many other systems with some straightforward vocabulary substitutions.

2. Conceptual “GAP” Model of Host Trait-Space Use

All plant species vary intraspecifically for numerous morphological, phenological, and chemical traits, with variation having genetic, epigenetic, and/or environmental causes (e.g., [42–45]). This variation defines a set of phenotypes that are available for attack by a phytophagous insect searching its environment for suitable hosts. This set of phenotypes can be depicted as a cloud of points in a multidimensional trait space, with each point representing an individual plant (or a ramet, for clonal plants; or even a module, when important variation occurs within individuals [46]). It is convenient to consider the two-dimensional case (Figure 1), which can represent either a system in which two plant traits show variation relevant to insect attack or a two-dimensional summary of a higher-dimensional trait space (using principal components to extract two dominant axes of trait variation). I use the term “available trait space” to describe this cloud of points, as combinations of plant traits falling inside it are available to attacking herbivores, while combinations outside are not available (i.e., they do not correspond to real plants which might be attacked). This trait space can be characterized by calculating its centroid (the point whose coordinate on each axis is the mean value of that coordinate for all individuals), its size (average distance from individual plants to the centroid), and its shape.

Consider first an insect interacting with a single host species. Some plant individuals will be attacked, but others will likely escape. The attacked individuals define the “attacked trait space” (filled circles in Figure 1), which must be a subset of the available trait space (and might be expected to be a smaller subset for more specialist herbivores [47, 48]). The relationship between available and attacked trait spaces will depend on active host-selection behavior by insects (insect preference), and also on whether insects can survive on an individual plant after initiating attack (insect performance). Both preference and performance will depend on plant traits—sometimes the same traits, but sometimes not. For simplicity, I use "herbivore attack" to denote the occurrence of feeding herbivores on plants, whether patterns in occurrence arise from preference or performance, and terms like “selective” attack should similarly be taken to include both preference and performance effects. The attacked trait space may represent a common preference by all herbivore individuals, or the sum of herbivore individuals’ distinct preferences in species with strong individual specialization [49, 50].
When herbivore attack is random with respect to plant traits (Figure 1(a)), the attacked trait space will resemble the available trait space in centroid location, shape, and (once corrected for the smaller number of attacked plants) size. For a selective herbivore (one that rejects some available plants), in contrast, the attacked and available trait spaces will differ. Many patterns are possible, but two are particularly likely. First, the herbivore might attack typical plants (those with trait values near the population means) and reject extreme ones, leading to an attacked trait space that is central with respect to the available trait space; the attacked space is smaller than the available space, but the two spaces have similar centroids (Figure 1(b)). Such a pattern might be favoured by selection because (for example) typical plants are most common, and insects preferring them pay lower search costs and experience greater resource availability. Alternatively, herbivore attack might be associated with extreme trait values (e.g., herbivores might perform best on the largest or least-defended individuals), leading to an attacked trait space that is marginal with respect to the available trait space: the attacked trait space is again restricted in size, but in this case the two spaces have different centroids (Figure 1(c)).

Now consider a pair of plant species available for attack (Extension to larger numbers of hosts is possibly but considerably complicating.) Given a common set of measured traits, we should see two clouds of points in trait space (Figure 2) defining a pair of available trait spaces. The distance between available trait spaces defined by different plant species might be large compared to the size of each available trait space [38], but, for closely related pairs of phenotypically variable plants, this need not be so (e.g., for Solidago altissima and S. gigantea, see Figure 3). Two available trait spaces could even be touching or interdigitated, especially for hybrid swarms [51]. One can again consider attacked trait spaces in comparison to available trait spaces on each host, but now there are many more possibilities, as each attacked trait space could be nonselective, central, or marginal (and if marginal, toward or away from the other host). Among possible patterns, I emphasize here a set of trait-space relationships predicted for an insect herbivore moving through a four-step evolutionary sequence: from original specialization on one of the two hosts, through a host shift, to early and late stages of HAD.

2.1. Stage 1: Single-Host Specialists and the Importance of Host-Choice Errors. An insect attacking a single host could show virtually any pattern in the relationship between attacked and available trait spaces, but some patterns are of special interest in the context of possible host-shifting to an evolutionarily novel host. (By a “host shift” I mean the addition of a novel plant to the herbivore species’ diet, which will normally occur without immediate abandonment of the old). Such host shifts are likely to begin when a few individuals attack the “wrong” (novel) host, making it possible for selection to favour the incorporation of the novel host into the insect’s host range. Importantly, the likelihood of host-choice errors is likely to depend on the insect’s use of plant trait space. In particular, imagine an insect showing a marginal attacked trait space on the ancestral host. That marginal attacked trait space could be adjacent to the available trait space defined by the novel host (Figure 2(a)), or could be distant from it. When it is adjacent, host-choice errors are more likely and insects making those errors are more likely to survive on the novel host [52]. In contrast, when the ancestrally attacked trait space is distant from the novel host, host-choice errors (and thus host shifting) should be less likely. I call this the “adjacent errors hypothesis.” The logic mirrors the widespread expectation that host-choice errors and host shifts are more likely between species that resemble each other morphologically, chemically, or phylogenetically [38] but stresses that the distance in trait space that needs to be crossed for a host-choice error depends not only on the distance between available trait spaces but also on how insect preference and performance define the attacked trait space.

2.2. Stage 2: Oligophagous Feeding Following Diet Expansion. Following a host shift that expands diet, our focal insect
species will be oligophagous, feeding on two hosts (ancestral plus novel) rather than one. Because this stage should follow from patterns in attack allowing host-choice errors (adjacent errors hypothesis), attacked trait space is likely to remain marginal on the ancestral host (Figure 2(a)). As host preference and performance evolve to include attack on the novel host, the attacked trait space on that host is likely to be marginal as well, but with the two attacked trait spaces adjacent (Figure 2(b)) because novel plants closer to the ancestrally attacked trait space are more easily colonized. I call this the “adjacent oligophagy hypothesis.”

Note that the adjacency pattern is equivalent to restricted but central use of an available trait space defined by the two hosts in combination (compare Figures 1(b) and 2(b)). If disruptive selection between alternative hosts does not act or is not powerful, oligophagy and the adjacency pattern could be evolutionarily persistent. Alternatively, this stage might be transient, persisting only until disruptive selection has
time to drive HAD of insect subpopulations exploiting the two hosts. The contrast between these possibilities highlights an important fork in the evolutionary road [53], in which disruptive selection favouring HAD is or is not sufficient to overcome gene flow working to homogenize the herbivore population and to maintain an oligophagous diet.

2.3. Stage 3: Nascent Host-Specialist Forms and the Selection-Gene Flow Tension. How might insect trait-space use favour or oppose the ability of disruptive selection to achieve HAD? Craig et al. [52] argued that persistent oligophagy is likely when the two available trait-spaces are very close, with HAD likely when they are more distant. However, their conceptual model assumes that attacked trait spaces on the two hosts remain indefinitely adjacent (their Figure 1). I suggest that there is another important possibility; a critical step in HAD may be the separation of the attacked trait spaces (Figure 2(c)) such that insect subpopulations on the two hosts (now appropriately thought of as nascent host forms) come to attack dissimilar individuals rather than similar ones. I call this the “trait distance-divergence hypothesis”. Separation of the two attacked trait spaces could arise in two different but complementary ways. First, distance between attacked trait spaces could arise simply because disruptive selection for adaptation to the alternative hosts overpowers the homogenizing effect of gene flow between nascent host forms. Under this scenario, the attacked trait spaces could move to opposite ends of the available trait spaces, as in Figure 1(c), or merely further apart than adjacency, depending on the shape of the fitness landscape—that is, fitness optima on the alternative hosts might favour central or marginal trait spaces. Under this scenario, distance between attacked trait spaces is just a symptom by which the progress of HAD can be recognized.

Second, distance could be a product of selection to minimize host-choice errors by each nascent host form or hybridization between them [22]. Host-choice errors could be opposed by selection because they lead to preference-performance mismatches, or because they put larvae in competition with members of the other host form (encouraging divergence by character displacement). Alternatively, host-choice errors coupled with the tendency for phytophagous insects to mate on their host plants could lead to hybrid matings. Hybrid disadvantage is possible given tradeoffs in ability to exploit the alternative hosts, or if hybrids prefer or are best suited for trait-value combinations falling in the gap between the two available trait spaces (this gap is shown narrow in Figure 2 but will often be wider [38]). Selection to reduce hybridization by widening the distance between attacked trait spaces (Figure 1(c)) would be a form of reinforcement [54]. Under the reinforcement scenario, distance between attacked trait spaces is more than a symptom of HAD; once achieved, it serves to reduce gene flow between nascent host forms and permit HAD to progress. (In passing, I note that if selection simply overpowers gene flow, we would expect HAD to involve genetic divergence in genomic islands, whereas if selection opposes hybridization, genome-wide genetic divergence should result via “isolation by adaptation”) [55–57].

In summary, the trait distance-divergence hypothesis holds that attainment of distance between attacked trait spaces (Figure 1(c)) can be both a symptom of HAD and also a factor permitting HAD. The larger the distance between attacked trait spaces, the more likely is the evolution of genetic differentiation between insects on the two hosts. Since genetic differentiation can ease the evolution of distance between attacked trait spaces, this stage of evolution can involve positive feedback [58–60]. In contrast, an insect for which attacked trait spaces remain adjacent (Figure 1(b)) is likely to remain an oligophagous insect with no host-associated structure to its gene pool.

2.4. Stage 4: Pair of Established Host Specialists. As HAD proceeds and gene flow between nascent host forms declines, we would expect the gradual accumulation of more, and more effective, reproductive isolating mechanisms [24, 61, 62]. This should continue until ecological speciation is complete, and the two host-specialist forms attain the status of full biological species. As reproductive isolation becomes enforced by multiple, redundant mechanisms, the importance of separation between attacked trait spaces should decline. Selection will then be free to mould trait-space use independently for each species, and if reinforcement earlier in HAD pushed the attacked trait spaces apart (trait distance-divergence hypothesis), this force can now relax. If selection favours use of central trait space on each host, for example (Figure 2(d); or if it favours nonselective use of trait space on each host), the distance between the attacked trait spaces should decrease. I call this the “distance relaxation hypothesis.”

Note that through the temporal sequence (Figures 2(a)–2(d)), the overall conceptual model suggests a pair of attacked trait spaces that begin close together, move apart, and then move back together like pincers. This movement underlies the terminology “gape-and-pincher” model of trait-space use.

2.5. What about Generalists? The foregoing considered insects that begin as monophagous on one of the two hosts and remain narrowly oligophagous or monophagous at all stages of HAD. However, many herbivores are broader generalists [63] for which we would not expect any of the trait-space patterns shown in Figure 2. In particular, it would be very surprising if a broad generalist showed nonrandom separation between attacked trait spaces (Figures 2(c) and 2(d)). Instead, attacked trait space might be nonselective on both hosts [47, 48], or restricted but marginal along a trait axis orthogonal to the difference between the two hosts (e.g., insects might prefer larger individuals of each host and also attack other, larger species). Such broad generalists are much less likely than host specialists to undergo HAD because (being already adapted to multiple hosts) they are less likely to experience strongly disruptive selection for performance on one host versus another [64].

2.6. Testing the Hypotheses. The four hypotheses that make up the GAP model are logically distinct; finding that one
hypothesis holds (or fails) implies nothing about the others. For example, for a given herbivore, the adjacent errors and adjacent oligophagy hypotheses could hold, but the distance-divergence and distance relaxation hypotheses fail, if HAD proceeds to ecological speciation without any movement of attacked trait spaces away from each other following the host shift.

Each of the four hypotheses can also be posed, and tested, at two levels. First, we can test each hypothesis for a single herbivore. For instance, are attacked trait spaces adjacent on *Solidago altissima* and *S. gigantea* for the narrowly oligophagous *Epiblema scudderiana* (adjacent oligophagy hypothesis)? Of course, such tests focus on patterns, and confirmation of a pattern need not constitute a strong test of underlying mechanism. Second, and more powerfully, we can test each hypothesis for herbivorous insects as a class. For instance, are attacked trait spaces (statistically) further apart for recently divergent and incompletely isolated pairs of host races than they are for more ancient specialist species pairs (distance relaxation hypothesis)? At this level, the hypotheses can hold strongly or weakly (or not at all); that is, the empirical relationship between trait-space distance and extent of reproductive isolation could be stronger or weaker (or non-significant).

Testing the adjacent errors, adjacent oligophagy, trait distance-divergence, and/or distance relaxation hypotheses for individual herbivores will require trait-space use data for large numbers of individuals on the alternative hosts. Some tests will be difficult at the individual-herbivore level (e.g., testing the distance relaxation hypothesis for an individual herbivore would require historical data on past trait-space use, which will only rarely be available). Ultimately, though,

<table>
<thead>
<tr>
<th>Pattern in trait-space use</th>
<th>Hypothesis</th>
<th>Attacked trait spaces marginal?</th>
<th>Attacked trait spaces restricted?</th>
<th>Attacked trait spaces distant?</th>
</tr>
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<tbody>
<tr>
<td>Monophagous, attacked trait space marginal and adjacent to alternative host (Figure 2(a))</td>
<td>Adjacency favours host shifting (adjacent errors hypothesis)</td>
<td>Ancestral host: marginal</td>
<td>Novel host: marginal but rare</td>
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<tr>
<td>Oligophagous, attacked trait spaces marginal and adjacent (Figure 2(b))</td>
<td>Adjacency persists after host shifting (adjacent oligophagy hypothesis)</td>
<td>Ancestral host: marginal</td>
<td>Novel host: marginal</td>
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<tr>
<td>Nascent host races, attacked trait spaces marginal and distant (Figure 2(c))</td>
<td>Distance permits, and is also symptomatic of, trait-space distance</td>
<td>Ancestral host: marginal</td>
<td>Novel host: marginal</td>
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<tr>
<td>Pair of monophagous species, attacked trait spaces central on each host (Figure 2(d))</td>
<td>Other isolating mechanisms reduce importance of trait-space distance</td>
<td>Ancestral host: not marginal</td>
<td>Novel host: not marginal</td>
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1 More strictly, only restricted or marginal trait-space use along the PC axis (or axes) defining the difference between available hosts is directly relevant to the GAP model.

2 But statistical detection is difficult because attack on the novel host is rare.
3. Statistical Methods

I developed statistical methods to test for three patterns in host trait-space use by insect herbivores. These patterns are predicted, in different combinations, by the adjacent errors, adjacent oligophagy, distance-divergence, and distance relaxation hypotheses (Table 1) and thus provide windows on the overall GAP model. The tests share a common framework in that they are based on relationships between attacked and available trait spaces for host plants of two species (Figures 1 and 2). Two tests pertain to the pattern of attack on a single host, and the third to the pattern of attack on each host relative to the other.

First, I test for central versus marginal location of the attacked trait space on each host (“Marginal trait-space test.”). I calculate the centroid of the available trait space (mean PC1 and PC2 scores for all available plants, attacked and unattacked) and that of the attacked trait space (mean PC scores for attacked plants only). I then calculate the distance between available and attacked centroids and compare this to a null distribution of 10,000 such distances calculated following random shuffling of attack status across all plant individuals. The fraction of randomization distances larger than the actual attacked-available distance is a \( P \) value, and when it is small we reject the null hypothesis that attacked and available plants have a common centroid (central or nonselective attack, Figures 1(a) and 1(b)) in favour of the alternative of marginal attack (Figure 1(c)).

When we are unable to detect marginal use of available trait space, we might seek to distinguish between nonselective (Figure 1(a)) and restricted but central (Figure 1(b)) alternatives. To do so, I use the “restricted trait-space test.” I calculate the Euclidean distance from each attacked plant to the centroid of attacked trait space. The size of the attacked trait space is given by the sum of these distances. I then compare this trait-space size to a null distribution the sizes of 10,000 attacked trait spaces generated by randomly shuffling attack status across all plant individuals. Note that shuffling attack status maintains the number of attacked plants, which is critical when calculating the size of a trait space. The fraction of randomization attacked trait spaces smaller than the actual one is a \( P \)-value, and when it is small we reject the null hypothesis that attack is nonselective. Since we are using the restricted trait-space test following a nonsignificant marginal trait-space test, the alternative is that herbivores exploit a restricted but central subset of available trait space.

Finally, I test whether the distance between attacked centroids on the two host plants is smaller or larger than expected at random (“Distant trait-spaces test,” Figure 2(b) versus 2(c)). I first calculate the distance between attacked trait-space centroids on the two alternative hosts. This distance is compared, in a two-tailed test, to a null distribution of 10,000 such distances calculated following random shuffling of attack status across individuals of each plant species (separately). When the actual centroids are farther apart than the mean distance from randomizations, then twice the fraction of randomization distances that are larger than the actual distance is a \( P \)-value, which when small supports rejection of the null hypothesis in favour of the alternative that the two attacked trait spaces are significantly distant. On the other hand, when the actual centroids are closer than the mean distance from randomizations, then twice the fraction of randomization distances that are smaller than the actual distance is a \( P \)-value, which when small supports rejection of the null hypothesis in favour of the alternative that the two attacked trait spaces are significantly adjacent.

The marginal trait-space, restricted trait-space, and distant trait-space tests are implemented in TraitSpaces 1.20, a program written in Microsoft Visual Basic.NET for Windows. The software takes as input a dat file with a row for each individual host plant, and columns for host species identity, presence/absence of each herbivore, and first and second principal components calculated from the host trait matrix. (Principal components may be output from any standard statistical package.) Extension to trait spaces of higher dimensionality, if desired, is straightforward; one could even use an unreduced trait matrix at the cost of some complexity in displaying results. The analytical framework easily accommodates data for other host/attacker systems and could even be applied to cases where consumers use variable microhabitats or food resources. The current version of the TraitSpaces package is available from the author on request.

4. Field Methods

4.1. Study System. The goldenrods Solidago altissima L. and S. gigantea Ait. are clonal perennials codistributed over much of eastern and central North America. Intermixed stands of the two species are common in open habitats such as prairies, old fields, road sides, and forest edges. Individual ramets grow in spring from underground rhizomes, flower in late summer and fall, and die back to ground level before winter. The two species differ most obviously in pubescence [65]: S. altissima stems are sparsely to densely short-hairy, especially basally, while S. gigantea stems are typically glabrous. Both species display extensive intraspecific variation (genetic and plastic) in most traits, including ramet size, pubescence, leaf shape, size, and toothiness, and chemical profiles ([27, 66, 67], S. B. Heard, unpubl. data).

S. altissima and S. gigantea are attacked by a diverse fauna of insect herbivores [68–70], which vary in diet specialization. Some are broad generalists that accept Solidago as part of a taxonomically diverse diet (e.g., the exotic spittlebug Philaenus spumarius [71]), and some are broadly oligophagous, feeding on Solidago among other members of the Asteraceae (e.g., the chrysomelid Exema canadensis [72]). Others are more narrowly oligophagous, attacking only Solidago spp. (e.g., the tortricid stem-galler Epiblema scudderiana [13, 73]). Finally, at least four herbivores have evolved monophagous host races or cryptic species on S. altissima and S. gigantea [13, 30, 74, 75], with divergence ranging from quite recent for the ball–gall fly Eurosta solidaginis (at most 200,000 years, but likely much less) to \( >2 \times 10^6 \) years old for the bunch–gall flies Rhopalomyia solidaginis/R. capitata.

Especially for the better-studied S. altissima, attack by various herbivores is known to vary among clones [68, 76, 77], and with plant traits including ramet size [39, 78],...
growth rate [79], nutritional status [80, 81] and ploidy where this varies locally [82]. These trait-attack relationships involve both plant resistance and insect preference [27] and may be concordant or discordant among different herbivore species [68, 82].

4.2. Field Data. I and my field team gathered data on plant traits and herbivore attack in old-field and trailside Solidago populations in Fredericton, NB, Canada (45° 57′ 30″ N, 66° 37′ 1–20′′ W). Here both S. gigantea and S. altissima are abundant along with S. rugosa, S. juncea, S. canadensis, Euthamia graminifolia, Symphyotrichum spp., and other Asteraceae. S. altissima is exclusively hexaploid in the east, and S. gigantea exclusively diploid, so effects of ploidy on herbivore attack [83] need not be considered here.

It is important to assess the available and attacked trait spaces using traits measured before herbivore attack; otherwise, herbivore responses to plant traits could be confounded with herbivore-driven changes in the same traits. In early June 2004, we marked 104 S. altissima ramets and 186 S. gigantea ramets by setting line transects through well-mixed patches of the two species and marking each ramet touched by the line. At the time of marking, a few ramets had already been attacked by the stem-galler Gnorimoschema gallaecolidaginis (Lepidoptera: Gelechiidae; galls on 4 S. altissima and 3 S. gigantea ramets), but other herbivores

Figure 3: Attacked trait spaces for four goldenrod herbivores on S. altissima (triangles) and S. gigantea (circles). Filled symbols denote attacked plants, and open symbols unattacked ones. Axis labels are shorthand for the first two principal components from a 7-variable morphological dataset; full factor loadings are provided in Table 2.
had yet to attack. We measured 7 morphological traits of our marked ramets, focusing on easily measured traits that were likely to influence herbivore attack, that help distinguish the two study species, or both. We measured stem trichome density by counting, in the field with a hand lens, all trichomes in silhouette along a 10 cm length of stem just below the terminal bud. We measured stem width 5–10 cm above ground using a caliper, and stem height (from ground to the base of the terminal bud) using a measuring tape. For the largest leaf from each ramet, we measured leaf length, leaf width at the widest point, and the number of teeth along one leaf edge. Finally, we weighed each largest leaf before and after drying to constant mass at 45–55°C and calculated percent water content.

We surveyed marked ramets twice weekly until the end of August, identifying herbivores present as specifically as possible without disturbing them on the plant (for some groups, like larval *Trirhabda* beetles, species-level identifications require the removal of the insects to the laboratory, and we wanted to leave plants to experience natural levels of herbivory). When herbivores of the same species were present on consecutive surveys, we were usually unable to determine whether they were the same individuals, so rather than count individuals we classified each ramet as attacked or unattacked by each herbivore over the course of the entire season.

Some marked ramets were lost or damaged during the season, leaving 92 *S. altissima* ramets and 175 *S. gigantea* ramets with comprehensive herbivory and plant-trait data. Four herbivores were identifiable to species and abundant enough to give our analyses reasonable power: the xylem-sucking spittlebug *Philaenus spumarius*, which is broadly polyphagous [71]; the folivorous chrysomelid beetle *Exema canadensis*, which is oligophagous with many hosts in the tribe Astereae [72]; the phloem-sucking aphid *Uroleucon nigrotuberculatum*, which is narrowly oligophagous on *Solidago* spp. [84]; and the gall-making cecidomyiid fly *Rhopalomyia solidaginis/R. capitata*, which is a pair of monophagous specialists (*R. solidaginis* on *S. altissima* and *R. capitata* on *S. gigantea* [13]). All further analyses use this reduced set of 267 ramets and 4 herbivores.

### 5. Field Results and Discussion

#### 5.1. Plant Traits

Among the 7 measured traits stem width, ramet height, and leaf length were strongly intercorrelated (0.72 < r < 0.88), suggesting that all three reflect overall ramet size. The other 18 correlations were weak to moderate (Table 2). The first two principal component axes explained 47% and 28% of the morphological variance (75% total), while no other axis explained more than 9.3%. PC1 largely reflects ramet size (heavy loadings for stem width, ramet height, and leaf length; Table 3), but also leaf toothiness (positively) and water content (negatively). PC2 contrasts trichome counts (strong positive loading; Table 3) with leaf width (strong negative loading) but also includes leaf water content, leaf length, and toothiness (all negative). These two principal components do a good job of capturing both intraspecific and interspecific variation (Figure 3), with *S. altissima* and *S. gigantea* separated primarily along PC2 (the pubescent *S. altissima* with high scores, and the glabrous *S. gigantea* with low scores).

#### 5.2. Herbivore Use of Phenotype Space

Attack rates by *Solidago* herbivores are generally low (often 1–10% or even less), with the exception of some diet generalists and outbreaking species in high-density years (S. Heard, unpubl. data). In our dataset, even though we worked with some of the most common herbivores, only one herbivore on one host had an incidence above 30% (*Exema canadensis* on *S. altissima*, 64% of ramets attacked). Other herbivore/host combinations had lower incidences, with several less than 10% (*Uroleucon nigrotuberculatum* on both hosts and *Rhopalomyia capitata* on *S. gigantea*; Table 4).

The patterns I document in trait-space use could have arisen via herbivore preference, or via performance if poor herbivore growth leads to death or departure of herbivores before surveys can detect them. For most herbivores, repeated surveys allow herbivore detection shortly after attack begins, and so preference is the most likely driver of patterns in attack. However, for gallmakers like *Rhopalomyia* performance at the stage of gall induction could be important.

### Table 2: Correlations among morphological variables measured for *S. altissima* and *S. gigantea*.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Leaf length</th>
<th>Leaf width</th>
<th>Teeth</th>
<th>Water content</th>
<th>Trichomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stem width</td>
<td>0.88</td>
<td>0.18</td>
<td>0.40</td>
<td>-0.42</td>
<td>0.36</td>
</tr>
<tr>
<td>Ramet height</td>
<td>0.72</td>
<td>0.47</td>
<td>0.56</td>
<td>-0.23</td>
<td>0.12</td>
</tr>
<tr>
<td>Leaf length</td>
<td>0.47</td>
<td>0.32</td>
<td>0.56</td>
<td>-0.24</td>
<td>0.57</td>
</tr>
<tr>
<td>Teeth</td>
<td>-0.18</td>
<td>-0.29</td>
<td>-0.24</td>
<td>-0.01</td>
<td>-0.48</td>
</tr>
<tr>
<td>Water content</td>
<td>-0.15</td>
<td>-0.42</td>
<td>-0.36</td>
<td>-0.04</td>
<td>-0.48</td>
</tr>
</tbody>
</table>

### Table 3: Factor loadings for the first two principal components from the morphological data matrix for *S. altissima* and *S. gigantea*.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Loading on PC1</th>
<th>Loading on PC2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stem width</td>
<td>0.63</td>
<td>0.04</td>
</tr>
<tr>
<td>Ramet height</td>
<td>0.51</td>
<td>0.04</td>
</tr>
<tr>
<td>Leaf length</td>
<td>-0.22</td>
<td>0.63</td>
</tr>
<tr>
<td>Teeth</td>
<td>-0.24</td>
<td>0.39</td>
</tr>
<tr>
<td>Water content</td>
<td>-0.39</td>
<td>0.58</td>
</tr>
<tr>
<td>Trichomes</td>
<td>0.34</td>
<td>0.58</td>
</tr>
</tbody>
</table>
Plant genotype effects on gall induction, mismatched with herbivore preference, are known (for example) for *Eurosta solidaginis* on *S. altissima* [85].

The two most generalist herbivores (*Philaenus* and *Exema*) showed similar patterns in trait-space use (Figures 3(a) and 3(b); Table 4). Both showed significant evidence for nonrandom use of available hosts (marginal trait-space test). For *Philaenus* on both hosts, attack was concentrated on larger but less pubescent ramets (higher PC1 and lower PC2), while for *Exema* on both hosts attack was concentrated on larger ramets but did not depend on pubescence (higher PC1). For both species, the distance between attacked trait-spaces on *S. altissima* and *S. gigantea* was slightly but not significantly smaller than expected under the null (distant trait-spaces test).

For the oligophagous *Uroleucon* (Figure 3(c)), attack on both hosts was significantly marginal, being concentrated on larger and less pubescent ramets. The distance between the two attacked trait spaces was slightly, but not significantly, larger than expected under the null. Because this herbivore had the smallest sample size (just 17 attacked ramets total), these tests have much less power than for the more common generalists.

For the monophagous *Rhopalomyia* species pair (Figure 3(d)), there was no evidence on either host for marginal use of trait space, and the restricted trait space test suggests nonselective rather than central use of available trait space (Table 4). The distance between the two attacked trait-spaces was slightly, but not significantly, larger than expected under the null. Sample size, however, was very small for *R. capitata* on *S. gigantea* (8 attacked ramets), so the tests for that species and for the distance between attacked centroids are likely not very powerful.

5.3. Interpretation and Prospects. The clearest pattern in the illustrative dataset is that for three of four herbivores, attack is significantly concentrated on larger ramets (large PC1). The stem gallers *Eurosta* [39] and *Gnorimoschema* [78] also have well-documented associations with larger ramets, something that is common but not universal among phytophagous insects [86, 87]. Such concordance across herbivore species in the use of trait space increases the likelihood of multiple herbivores cooccurring on a single ramet—something very unlikely under the null hypothesis of independent occurrence, since most attack rates are low. Herbivores cooccurring on a plant may compete directly (via resource consumption) or indirectly (via induced resistance) or may even show facilitation [88] although we know little about potential interactions among goldenrod herbivores [89–91]. However, concordance among goldenrod herbivores in use of plant trait space is far from universal [68, 82].

How do the illustrative data fit with the GAP model of trait space use during host shifting and HAD laid out above? The tendency for most herbivores to attack larger ramets (larger PC1) generates pattern in trait-space use. However, this shared tendency means that both attacked trait spaces are offset from the available spaces, in parallel and orthogonally to the contrast between alternative host plants (PC2). The distance between attacked trait spaces is unaffected, and so this ramet-size pattern is not directly relevant to the GAP model. Three of the four herbivores analyzed (*Philaenus*, *Exema*, and *Uroleucon*) have host ranges broader than just the *S. altissima*-S. gigantea pair and might therefore be expected to be rather unselective about traits distinguishing the two hosts. Indeed, *Exema* showed no offsets between attacked and available trait spaces along the principal components axis contrasting *S. altissima* and *S. gigantea* (PC2; Table 4). *Philaenus* and *Uroleucon* did show offsets along this axis, but because they were in the same direction and roughly equal on the two hosts, separation of attacked trait-spaces was not significantly large for either herbivore (Table 4, distant trait spaces test). The fourth herbivore, *Rhopalomyia*, is a pair of relatively old monophagous species [13]. For such a species pair, the distance relaxation hypothesis suggests that the use of distant trait spaces may no longer be an important barrier to gene flow (Figure 2(d)). *Rhopalomyia’s* use of trait space

<table>
<thead>
<tr>
<th>Herbivore</th>
<th>Host</th>
<th># Attacked plants</th>
<th>Distance from available centroid</th>
<th>Marginal trait-space test P</th>
<th>Restricted trait-space test P</th>
<th>Distance between attacked trait-spaces</th>
<th>Distant trait-spaces test P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>PC1</td>
<td>PC2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Philaenus</em></td>
<td><em>S. altissima</em></td>
<td>25</td>
<td>1.00</td>
<td>-0.50</td>
<td>&lt;0.001</td>
<td>— 4</td>
<td>Small</td>
</tr>
<tr>
<td></td>
<td><em>S. gigantea</em></td>
<td>22</td>
<td>1.11</td>
<td>-0.52</td>
<td>0.001</td>
<td>—</td>
<td>Small</td>
</tr>
<tr>
<td><em>Exema</em></td>
<td><em>S. altissima</em></td>
<td>39</td>
<td>0.30</td>
<td>0.05</td>
<td>0.023</td>
<td>—</td>
<td>Small</td>
</tr>
<tr>
<td></td>
<td><em>S. gigantea</em></td>
<td>33</td>
<td>0.65</td>
<td>0.04</td>
<td>0.023</td>
<td>—</td>
<td>Large</td>
</tr>
<tr>
<td><em>Uroleucon</em></td>
<td><em>S. altissima</em></td>
<td>9</td>
<td>1.39</td>
<td>-0.10</td>
<td>&lt;0.001</td>
<td>—</td>
<td>Large</td>
</tr>
<tr>
<td></td>
<td><em>S. gigantea</em></td>
<td>8</td>
<td>2.32</td>
<td>-0.62</td>
<td>&lt;0.001</td>
<td>—</td>
<td>Large</td>
</tr>
<tr>
<td><em>Rhopalomyia</em></td>
<td><em>S. altissima</em></td>
<td>27</td>
<td>0.09</td>
<td>0.35</td>
<td>0.21</td>
<td>0.18</td>
<td>Large</td>
</tr>
<tr>
<td></td>
<td><em>S. gigantea</em></td>
<td>8</td>
<td>0.66</td>
<td>-0.05</td>
<td>0.31</td>
<td>1.0</td>
<td>Large</td>
</tr>
</tbody>
</table>

1 Of 92 available *S. altissima* and 175 available *S. gigantea* ramets.
2 Attacked centroid minus available centroid (PC1 and PC2 components). A positive entry means that ramets with a large PC score are more likely to be attacked.
3 Small if the two attacked trait spaces are adjacent (Figure 2(b)), and large if the two attacked trait spaces are distant (Figure 2(c)).
4 This test is informative only when the marginal trait-space test is not significant.
Table 4: no evidence for marginal attack, and separation between the two attacked trait spaces no larger than expected at random) is consistent with this (Table 1). Overall, none of the illustrative data are inconsistent with the GAP model, but none of the four herbivores analyzed provides a strong test of its hypotheses. I did not find any examples of the patterns hypothesized for a single-host specialist making host-choice errors, for a narrowly oligophagous species immediately following a host shift, or for a pair of nascent forms early in HAD (Table 1, Figures 2(a)–2(c)). This is not surprising, though, because species known to be in early stages of HAD on S. altissima/S. gigantea, such as the ball-gall fly Eurosta solidaginis and the spindle-gall moth Gnorimoschema galleasolidaginis, were insufficiently abundant for analysis. At the broader level of hypothesis testing, four herbivores constitute just a small step towards assessing general patterns in trait-space use through HAD and ecological speciation. It will take many studies like mine, with herbivores on Solidago and other plants, before we can assess the generality of patterns in trait-space use.

Because attack rates for most Solidago herbivores are low, achieving powerful hypothesis tests for any herbivore will entail marking very large numbers of ramets—especially since ramet selection must be done before attack begins to avoid distortions of attack-space measurements if trait values change under herbivore attack. In Solidago, for instance, ramet biomass and height are often reduced by herbivory [91–93]. I am currently expanding on the illustrative study with the goal of securing larger sample sizes for the herbivores studied here and acceptable sample sizes for many more herbivores.

Another obvious limitation of the illustrative dataset is that it includes measurements of only seven plant traits, and conspicuously omits leaf-chemistry traits (and ploidy [83], which varies elsewhere but not in New Brunswick). S. altissima and S. gigantea have complex secondary chemistry, and variation in leaf chemistry is known to influence herbivore attack [66, 67]. Unmeasured morphological traits may also be relevant to insect attack; for instance, Philaenus prefers plant species and individuals with wider leaf axils [94], and this trait varies among S. altissima genotypes (Maddox unpubl. in [77]). Expanding the list of measured traits, and especially incorporating leaf chemistry, is a high priority for future work.

Despite the small numbers of attacked ramets and measured traits that earn the “illustrative” dataset its descriptor, my analysis of trait-space use for four herbivores establishes that the field and analytical approach outlined here is feasible and can detect nonrandom trait-space use. Because the goldenrod system includes such a diverse herbivore community attacking syntopic ramets of the alternative hosts, it offers the potential for great progress in testing hypotheses about host trait-space use during hostshifts and HAD.

6. General Discussion

The literature on how insect preference and performance vary with intraspecific variation in host-plant genotype, morphology, chemistry, and phenology is immense [95, 96]. Similarly, interspecific variation in the same kinds of traits has been widely held up as the key to the macroevolutionary fate of herbivore lineages (host shifting, diversification, specialization, and so forth [38, 63, 64, 97]). What is surprising is that the intersection of these perspectives is so little developed: we know almost nothing about trait-space use in systems where host shifting and HAD are suspected. This gap is clearly illustrated by the two best-studied cases of HAD in phytophagous insects: Eurosta solidaginis on Solidago altissima and S. gigantea and Rhagoletis pomonella on apple and hawthorn. For Eurosta, despite a wealth of information about how preference and performance relate to genetic and trait variation within S. altissima [27], few comparable data are available for flies attacking S. gigantea (except see [40]). For Rhagoletis, much has been written about the importance for HAD of apple-hawthorn differences in ripening phenology [26, 98, 99] and fruit size [100]. However, data on local intraspecific variation in phenology appear to be unavailable (although latitudinal clines have been documented [26]), and data on intraspecific fruit size variation appear limited to confirming significance of interspecific differences in average fruit size [100]. This is not to criticize work on these two model systems, which has pioneered the study of HAD, but rather to draw attention to a significant opportunity for progress.

Of course, the GAP model likely falls short of recognizing the full complexity of trait-space use in nature. While I have focused on snapshots of trait variation and insect use of trait space at a single site and in a single year, both available trait space and its use are likely to vary in space and time. This variation could have interesting and important consequences for HAD. For example, intraspecific variation in Solidago phenology and the difference in average phenology between S. altissima and S. gigantea change in space and time, and phenological differences are involved in host choice for at least two Solidago herbivores undergoing HAD (Eurosta [40]; Gnorimoschema galleasolidaginis, S. B. Heard, unpubl. data). Hawthorn phenologies show latitudinal gradients across space favouring local adaptation rather like that required during Rhagoletis’ host shift to apple [26]. There are thus likely to be places or times that are more conducive to host shifts and HAD than others [40, 52, 53]. Superimposed over this variation in available trait space can be strong geographic variation in insect preference (e.g., [82, 101]) and thus trait-space use. As a consequence, the places or times conducive to host shifting for one insect herbivore might not be so conducive for shifts by another. This is consistent with the evolutionary pattern seen in the Solidago system, in which three gallmakers have made host shifts from S. altissima to S. gigantea but have done so at different times [13].

Thinking about intraspecific variation in plant traits, and patterns of insect attack with respect to that variation, can expand and enhance our view of ecological speciation by phytophagous insects. Testing the hypotheses I frame about trait-space use for herbivores differing in diet breadth and in progress along the evolutionary sequence of HAD (Table 1) could take us a long way towards a predictive understanding of diet evolution and specialization in phytophagous insects. Ultimately, we would like to know for which taxa...
host-shifting and HAD are likely, and for which taxa they are not—and why [13, 38, 64, 102]. While much data collection and analysis lies ahead, the trait-space perspective promises a new and powerful window on the fascinating complexity of insect-plant interactions and herbivore diversification.

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References


