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Project report submitted to the Organic Farming Research Foundation

Project Title:

Genetic population structure of a parasitoid wasp

FINAL PROJECT REPORT

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Summary

Parasitoid insects that use different hosts can have a subdivided population structure that corresponds to host use. A subdivided population structure may favor local adaptation of subpopulations to small-scale environmental differences and may promote their genetic divergence. In this paper, heritable Random Amplified Polymorphic DNA (RAPD) markers visualized by single strand conformational polymorphisms (SSCP) analysis were used to examine the population structure of the parasitoid wasp *Diaeretiella rapae* (Hymenoptera: Braconidae) in an environment where two aphid hosts are available for oviposition. We found 11 codominant and 34 dominant RAPD polymorphisms that conformed to Mendelian segregation patterns. A nested analysis of variance indicated extensive genetic differentiation among six populations ranged from 1.2 - 1.6, indicating a relatively low dispersal rate. Genetic distances were also calculated between populations and the resulting trees indicated that populations less than 1.0 km from each other were genetically differentiated. Our results indicate that *D. rapae* populations are genetically subdivided on a small spatial scale that corresponds to host use patterns.

Introduction

Nearly all species of plants and animals display genetic differentiation at some geographic scale, usually as a result of habitat discontinuity, habitat fidelity or limited dispersal. In some cases, local environmental variation may cause natural selection to operate differently among populations, leading to genetic differentiation (Via 1994). The evolutionary forces responsible for creating genetic differentiation among populations are limited migration, small local population sizes, and restricted matings within a population (Hard & Clark 1989; Rank 1992; Roderick 1996). Dispersal of individuals into other populations can ameliorate these effects by increasing gene flow among populations (Slatkin 1987, 1994). Therefore, the presence of population structuring indicates that one or more of these forces may be operating.

While identifying the forces responsible for genetic differentiation of populations is of interest to evolutionary biologists, it also important to practitioners of biological control. Biological control theory predicts that useful natural enemies will have high dispersal rates compared to their prey or hosts, especially in heterogeneous environments (van Emden 1974; Murdoch 1990; Roderick 1992). This implies that the most successful natural enemies should not form local subpopulations but should remain panmictic. Despite the recognized importance of natural enemy mobility, little is known about their population structure or about the rates of gene flow between local populations of predators and parasitoids (Hopper et al 1993; Roderick 1996).

One of the problems in the analysis of population structure is determining the amount of gene flow between populations. It is usually impossible to detect individual insects moving long distances (but see Antolin and Strong 1987), therefore we must rely on indirect measures such as DNA markers to infer gene flow (Roderick 1996). Historically, the majority of gene flow analyses have examined protein polymorphisms (allozyme markers) to describe allele frequency

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changes in time and space. However, allozymes may not provide adequate resolution for determining migration patterns due to low mutation rates and selection acting to limit variability of protein allozymes (Avise 1994). In addition, Hymenoptera have exceptionally low allozyme variability (Graur 1985). Recent developments of molecular genetic markers that detect DNA-level variability have increased the power of insect population genetic studies (Avise 1994; Haymer 1994; Roderick 1996). We have developed and enhanced the use of Randomly Amplified Polymorphic DNA amplified by the Polymerase Chain Reaction (RAPD-PCR) to generate repeatable and heritable markers useful in molecular taxonomy, genome mapping and population genetics (Hiss et al. 1994; Antolin et al. 1996).

In this study, RAPD-PCR markers were used in conjunction with single strand conformation polymorphism (SSCP) analysis to examine the population structure of the parasitoid wasp *Diaeretiella rapae* M'Intosh (Hymenoptera: Braconidae). *D. rapae* is a parasitoid of many aphid species that feed on various plant types but it is commonly found parasitizing aphids that feed on cruciferous plants. In previous studies, we found that *D. rapae* is primarily attracted to cruciferous plant odors (Vaughn et al. 1996). However, *D. rapae* is used as a biological control agent against the Russian wheat aphid (*Diuraphis noxia*) that feeds on cultivated grasses in the western United States. Thus, *D. rapae* has a choice between two aphid hosts where these plant types are grown together. To assess whether this situation may lead to population differentiation we examined the genetic structure of this parasitoid on a microgeographic scale within managed agricultural systems in northern Colorado.

Methods

Study organisms

D. rapae is a solitary endoparasitoid of many aphid species (Vater 1971; Stary 1976) with

adult females laying a single egg directly into an aphids' body. The parasitized aphid continues to feed for 3-4 d and typically remains in the same location after being attacked. D. *rapae* progresses through four larval instars as it feeds on the internal tissues of the aphid, eventually killing it. When the host dies it becomes a mummy consisting of the hardened exoskeleton of the aphid. The wasp pupates within the aphid host and emerges from the mummy as an adult. Under laboratory conditions $(20^{0} \text{ C}; 60\% \text{ RH})$, egg to adult development requires approximately 9 d.

In northern Colorado, two aphid species are commonly attacked by *D. rapae*, the cabbage aphid (*Brevicoryne brassicae*) and Russian wheat aphid. Both aphid species are serious pests to agricultural crops: *B. brassicae* feeds exclusively on crucifer crops and *D. noxia* feeds exclusively on cultivated cereals. Crucifer crops and cereal crops are grown in close proximity to each other in some regions while in other regions, cereal crops, mainly dryland wheat, are grown in extensive monoculture. This scenario affords *D. rapae* the opportunity to parasitize two aphid hosts which may be separated by many kilometers or as little as several meters.

Field collections

Wasps were collected from crucifer vegetable crops and from cultivated cereal crops in northern Colorado. Individual wasps were removed from aphid colonies on host plants by gently brushing the mummified aphids individually into gelatin capsules. Aphid mummies were collected to ensure that the wasps analyzed were the progeny of adult females that located and chose to parasitize a particular aphid in a particular location. Aphid mummies were returned to the laboratory, were maintained at 23° C until the adults emerged, then were sexed and frozen at 80° C.

A hierarchical sampling design was used to collect *D. rapae* populations from six fields within three regions of northern Colorado (Table 1). Within each region, we located two aphidinfested fields less than 1.0 km from each other. Region I was located at Piedmont Farms, Inc., an organic

vegetable production area. The two wasp populations collected from this area were sampled from Brussels sprouts and winter wheat, separated by 0.25 km. Region II was located 23 km southeast of Region I. The populations sampled in this region were located on 4V Farms, Inc., a cereal production farm in Weld county. Both *D. rapae* populations were collected from winter wheat fields separated by 0.50 km. Region III was located 12 km southeast of Region I and 18 km southwest of Region II. The two populations collected in this region were 1.0 km apart and were collected from a winter wheat field and a cabbage field.

Each of the six populations were sampled three times per year (June, July and August) in 1994 and 1995 to test for seasonal differences in population structure. For each sampling period, 46 individuals were analyzed (23 males and 23 females). The field sampling procedure followed a modified version of Quick Sample, a tool for determining RWA infestations in cereal fields (Legg et al. 1991). Fields were entered at one corner and a plant was sampled for *D. rapae* mummies every 10 -20 paces. Only one wasp mummy was collected from any plant. This procedure ensured that the entire field was sampled and reduced the chance of collecting several progeny from a single female wasp.

DNA extraction, PCR and electrophoresis

DNA was isolated from individual wasps following the protocol of Coen et al (1982). DNA was resuspended in 100 ml TE (10 mM TRIS-HCl, 1mM EDTA, pH 8.0). Details of the RAPD-PCR protocols can be found in Black et al. (1992). One μ l of DNA template was used in each 50 gl PCR reaction. Each set of PCR reactions was checked for contamination by using a negative control (all reagents except template DNA). Samples were stored at 4°C until being electrophoresed.

Complete details of the polyacrylamide gel electrophoresis (PAGE) procedure is described

in Hiss et al (1994) and Black & DeTeau (1996). Modifications to this procedure are in Antolin et al. (1996). Briefly, PCR products (5 p1 of a 50 p1 sample) were mixed with 2 p1 loading buffer at room temperature. The products were electrophoresed on large (40 x 50 cm), thin (0.4 mm) glycerol (5%) polyacrylamide (5%, 2% crosslinking) gels. Samples (5 p1) were loaded into 6 mm sharkstooth combs. Electrophoresis proceeded at constant voltage (350 V) at room temperature for 15 hr. The gels were subsequently stained using silver to detect the mobility of the different DNA markers.

Segregation analysis of RAPD-SSCP markers

The majority of RAPD markers are expressed as phenotypically dominant markers in diploid organisms. However, because of the high resolution offered by the PAGE technique and the haplodiploid sex determination in Hymenopterans, we have also identified numerous codominant markers using a segregation analysis. Inheritance of the bands was tested by generating full-sib families from each of the field populations. Two families were created for each of the two Piedmont populations and one family was created for the remaining four populations. *D. rapae* families were produced by collecting individuals as mummies and rearing them in an incubator (25°C, 60% rH). Upon emergence, mated pairs were formed and the females allowed to oviposit on an aphid infested plant for 24 hr. The mated pair was then stored at -80°C. The progeny were individually collected as mummies, allowed to emerge as adults and also stored at - 80°C. These families were screened with 18 RAPD primers. We chose the following four primers from Operon Technologies, Inc.: A05, COI, CO4 and BAMH1 (5'-ATGGATCCGC, designed by N. M. DeTeau, Colorado State University) with which to test Mendelian inheritance patterns. Repeatability was then tested by amplifying the DNA of each individual in each family twice.

Amplified markers were scored directly from dried gels by measuring band mobility relative to

known size markers (1 Kb ladder, BRL Laboratories). To estimate sizes of the amplified DNA fragments, size standards were fitted to an inverse function that relates fragment size and mobility (Shaffer & Sederoff 1981). Each marker was labeled by the primer used in the PCR reaction and by its mobility on the gel (e.g. A05.275).

Analysis of population subdivision

The RAPD-PCR markers generated from our field collected populations were analyzed in two ways. The most commonly used methods to estimate population subdivision are Wright's F statistics and related estimates (Wright 1978; Cockerham & Weir 1993; Slatkin 1993). These indices measure the correlation between alleles found in subpopulations, relative to the population as a whole. Fst represents the standardized variance in allele frequencies among local populations and ranges between 0, indicating no genetic differentiation, and 1, indicating complete differentiation. However, calculating Fst assumes that populations are in Hardy-Wienberg equilibrium. The frequency of the codominant alleles from the female samples was examined using the computer program BIOSYS-1 (Swofford and Selander 1981) and deviations from Hardy-Wienberg equilibrium were tested using a chi-square analysis.

Estimates of Fst and effective migration rates (Nm) among the females in each population (n=23) were calculated using the computer program RAPDFST provided by W.C. Black IV (Department of Microbiology, Colorado State University, see appendix). The program calculates standard Fst and effective migration rates (Wright 1978) as well as estimates corrected for small sample sizes (Cockerham & Weir 1993) and for both small sample sizes and the effects of dominant loci commonly detected by RAPD-PCR (Lynch & Milligan 1994).

Linkage disequilibrium

Linkage disequilibrium describes the nonrandom association of alleles (Lewontin & Kojima 1960) and can arise in subdivided populations from epistatic natural selection within subpopulations (Lewontin 1974), random genetic drift among subpopulations (Ohta & Kimura 1969) or actual linkage among the markers. Di-locus linkage disequilibrium was estimated using RAPDLD (Black 1995; Apostle et al 1996; see appendix) which estimates linkage disequilibrium among RAPD alleles in the subpopulations.

Genetic distance

The second method of analysis was based upon pair-wise genetic distances between populations, which does not depend upon the assumption of population equilibrium (Slatkin 1993). The number of markers generated with RAPD-PCR allowed us to not only estimate genetic distances but to perform robust statistical analyses of population subdivision at smaller spatial scales. In conjunction with W.C. Black N, a computer program called RAPDDIST (Black 1995) was developed that uses RAPD data to estimate genetic distances for haploid and diploid organisms (see appendix). Nei's unbiased genetic distance (Nei 1978) between pairs of populations were then submitted to Phylip 3.5c (Felsenstien 1995) to create cluster diagrams (trees) based upon UPGMA, an unweighted pair-group method with arithmetic averaging clustering analysis (Avise 1994). RAPDDIST further allows for statistical testing of individual branches on the trees by resampling the data, thus testing for consistency within the RAPD data set in forming the observed groupings. Each sex was analyzed separately for each sampling date and also by combining all sampling dates.

Results

Segregation analysis

The four primers generated a total of 34 dominant (presence/absence) markers and 11 codominant markers in the families, all of which were polymorphic and whose observed frequency was less than 0.92, as recommended by Lynch and Milligan (1994). Primer A05 produced 11 dominant and 4 codominant markers, COI produced 9 dominant and 3 codominant markers, C04 produced 8 dominant and 3 codominant markers and BAMHI produced 6 dominant and 1 codominant marker (Table 2). Repeatability was verified by reamplifying the DNA of each individual in each family with all four primers a second time and identifying the same markers. The inheritance of each marker conformed to Mendelian segregation patterns among the full-sib families. Figure 1 demonstrates common patterns found in the segregation of RAPD-PCR markers: Mendelian inheritance of codominant RAPD markers for haplodiploid families can be determined as follows: if a female having a fast and slow allele is mated to a male with a slow allele, the male offspring will have either a fast or a slow allele and the female offspring will be either heterozygous or homozygous for the slow allele (fig. I a). In the case of a dominant marker, if a female homozygous for the null allele (no band present) is mated to a male that carries a copy of the allele, none of the male offspring will show the marker but all of the female offspring will have a copy because they developed from fertilized eggs (fig. Ib).

Estimation of population subdivision

Deviations from Hardy-Weinberg equilibrium were tested for each of the 11 codominant loci in each of the six populations using the female wasps. No deviations from Hardy-Weinberg equilibrium were found (all)e < 3.33; df = 1; p > 0.10) except for one locus in the population

obtained from BARI (x2 = 3.94; df = 1; p < 0.05). The mean heterozygosity, averaged across sample dates and RAPD loci for each population was calculated and ranged from 0.335 - 0.443 (Table 3).

All three estimates of population subdivision were similar in this study for males and females combined: Fst estimated from Wright (1951) was 0.066 in 1994 and 0.073 in 1995 (p<0.01); Wier & Cockerham's theta (0), which accounts for small sample sizes, was estimated at 0.063 in 1994 and 0.070 in 1995; and Fst estimated from Lynch & Milligan (1994) which accounts for small sample sizes and dominance effects inherent with RAPD loci was estimated at 0.061 in 1994 and 0.069 in 1995. Fst and 0 values for each sampling date are given in Table 4.

The data set was subjected to a nested analysis of variance following Wright (1978) using BIOSYS-1 (Swofford & Selander 1981) to determine the level at which the populations were genetically subdivided. Variance components indicate that most of the variation within the population occurred between fields rather than between the areas containing the fields for all sampling dates in each year (Table 5). The effective migration rates obtained for all populations for 1994 and 1995 were 1.2 and 1.6, respectively.

Linkage disequilibrium

With 34 RAPD loci there were 561 [(34x33)/2] pair-wise comparisons made to estimate linkage disequilibrium. The amount of linkage disequilibrium detected in any population was less than 3% and in no population was this amount significant at a = 0.05. This analysis indicates that the RAPD markers used in this study were randomly distributed throughout the genome.

Cluster analysis

Cluster diagrams were generated for each sex for each sample date and for each sex for all

sample dates combined for each year. There were no differences between the trees generated for males or females at any sampling date and there were no significant differences in topology between sample dates within either sex using strict *consensus* (> 50% grouping). Trees generated using females from all sampling dates within 1994 and 1995 are shown in figure 2. Although there is weak support for population subdivision among areas (e.g.; Piedmont Farms wheat and crucifer populations versus Weld County wheat populations, fig. 2a), complete support is given to each branch that includes the same fields in two different years (fig. 2b). Particularly, crucifer fields (Piedmont farms and Hort Farms) remain genetically distinct from nearby wheat fields (Piedmont Farms and BARI), even less than 1.0 km from each other.

Discussion

("The habitats of most insect species are heterogeneous and discontinuous in time and space and typically contain patches that provide essential resources such as food or reproduction sites (Elton 1949; ,Southwood 1977). In such cases, gene flow between populations can be restricted because individuals tend to move mainly within physically or otherwise separated habitats that contain essential resources. In many invertebrates, a combination of limited dispersal and habitat fidelity results in marked population subdivision, sometimes even over small geographic distances. For example, population substructuring on a microgeographic scale has been demonstrated in a wide variety of organisms, showing that gene flow is often extremely limited, even over relatively short distances (Bilton 1992; Rank 1992; Seppa & Pamilo 1995; Appostol et al. 1996; West 1996). Resource fragmentation is common in agricultural systems especially in parasitoid host interactions where more than one host type is available. Thus, the diverse habitat encountered by parasitoids in agricultural environments may cause population differentiation.

This study of the population genetic structure of D. rapae is one of the first studies to

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demonstrate genetic differentiation in a parasitoid or biological control agent. Because of its use as a biocontrol agent, the biology of this wasp has been extensively studied. However, most of the research conducted on this wasp has concentrated on how it locates aphid hosts (Read et al. 1970; Ayal 1987; Sheehan & Shelton 1989b), especially with reference to chemical cues (Sheehan & Shelton 1989a; Reed et al. 1995; Vaughn et al. 1996). *D. rapae is* attracted to the volatiles emitted from crucifer plants and may possess specialized receptors used in locating habitats with these plant volatiles. Yet, *D. rapae* parasitizes many aphid species not found on crucifer plants. In this study, the population structure of *D. rapae* was examined with particular reference to habitat types which occur in diverse agricultural systems.

One of the difficulties encountered in population genetic studies, especially of Hymenopterans, is the level of resolution offered by indirect estimates of gene flow. With the advent of RAPD-PCR technology, it is. now possible to estimate differentiation of populations on a relatively small spatial scale because of the large number of genetic polymorphisms. In addition, the use of large polyacrylamide gels combined with the greater resolution of silver staining made the analysis of these polymorphisms far easier and more accurate than other methods of detecting genetic differences between populations. By establishing the heritability of RAPD-PCR markers as we have in this study, these markers are as effective as any other genetic marker used in population level studies (Carlson et al. 1991; Hunt & Page 1992; Apostol et al. 1993; Fondrk et a. 1993; Mitchell-Olds 1995; Antolin et al. 1996). The use of genetic markers such as RAPD's also tests whether the entire genome supports the observed groupings rather than just a few loci. This is further supported because none of the RAPD markers were in significant linkage disequilibrium.

The estimates of population subdivision using Wright's F-statistics certainly indicate that the *D. rapae* population is subdivided. The methods of Wright (1951), Wier and Cockerham (1984) and Lynch and Milligan (1994) all produced similar estimates of population differentiation.

Further, the Fst and variance components indicate that most of the variation in northern Colorado occurs between fields rather than between areas containing the fields for all sampling dates within each year. Thus, the individual fields within an area are as different from each other as are fields in different areas, even though pairs of fields within any area are separated by 1 km or less and areas are separated by as much as 23 km. In addition, the comparisons of sampling dates within each year indicates that the subdivision among populations is maintained from year to year, even in small geographical areas, and does not depend on recolonization each year. This result corresponds to *D. rapae* biology because wasps overwinter as fourth instars in aphid hosts, thus remaining in the vacinity(or field) as their parents.

The analysis of the genetic distances between populations also demonstrates that the population is structured in the agricultural environment. These results indicate that once wasps locate habitats with aphids, they remain in those areas and disperse less than 1 km, whether those habitats are wheat fields or crucifer fields. However, the two populations from the Weld County wheat fields appear to be more homogeneous (fig. 1). The two fields group within the same year for both years and the internal node grouping the Weld County Region is high. This has important implications for biological control programs where wasps are released each year. For example, if there are no ameliorating factors involved, *D. rapae will* most likely remain in the specific fields where the releases occur and have low migration rates between fields. Indeed, field observations and previous work with *D. rapae* in flight tunnels (Sheehan & Shelton 1989; Vaughn et al 1996), *D. rapae* was shown to have low a tenacity for sustained flight and spent most of its time at or below the vegetation canopy which likely accounts for its low dispersal rate.

While there are numerous studies of herbivorous insect populations that demonstrate genetic subdivision at small spatial scales (McCauley & Eanes 1987; McCauley et al 1988; McPheron et al 1988; Guttman et al 1989; Guttman & Weigt 1989; Rank 1992), this is the first

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study of a parasitoid wasp to demonstrates genetic subdivision. Predatory lady beetles (Coleoptera: Coccinellidae) are the only natural enemies whose population structure has been studied extensively (Krafsur et al. 1992, 1995; Steiner & Grasela 1993; Coll et al. 1994). These studies employed allozyme electrophoresis to examine gene flow and the main conclusion drawn from each study was that the lady beetle population is not subdivided, even between states in some cases. This suggests that the high mobility of coccinellids contributes to their effectiveness as biological control agents. A recent study, however, has demonstrated genetic subdivision using allozymes between Hawaiian populations of the Coccinellid *Curinus coeruleus* (Follett & Roderick 1996). The levels of gene flow found in this system are similar to the levels of gene flow found in our study.

For insects that inhabit heterogeneous environments, the association with patchily distributed resources has important consequences for the evolution of insects whose dispersal is limited. Because the greatest opportunity for genetic variation in resource use occurs within patchy environments (Via & Lande, 1985; Futuyma & Philippi, 1987), migration between local patches may cause conflicts that produce "tradeoffs" in performance unless the populations can adapt to different environments (Via & Lande, 1985). The possibility of host adaptation is currently being investigated in these *D. rapae* populations.

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Table 1	Sampling locations for <i>D</i> .	rapae	populations in	Colorado in	1995 and	1996.

	and the second			
	Population		Distance	Distance Between Fields
Region	n Location	Field Type	from Region I (Km)	within each Region (Km)
Ι	Piedmont Farms	Wheat	0	
Ι	Piedmont Farms	Crucifer	0	0.25
П	Weld County	Wheat	23	
П	Weld County	Wheat	23	0.05
ш	Hort Farms	Crucifer	13	
Ш	BARI	Wheat	12	1.0

Table 2 Frequency of RAPD markers for all six *D. rapae* populations for females and males for each year. Shown are the frequencies for the band present alleles (scored as '1') for the dominant markers and the 'fast' alleles for the codominant markers (denoted by *).

	1004	1005		1004	1005
) (. 1	1994	1995		1994	1995
Marker	Female Male	Female Male	Marker	Female Male	Female Male
A05.275	0.861 0.842	0.755 0.701	C01.372	0.667 0.690	0.717 0.966
A05.320*	0.694 0.667	0.761 0.696	C01.415	0.283 0.370	0.304 0.396
A05.380	0.619 0.671	0.543 0.630	C01.452	0.333 0.310	0.239 0.283
A05.465	0.521 0.474	0.590 0.523	C01.452	0.543 0.652	0.478 0.578
A05.486	0.385 0.425	0.467 0.511	C01.569	0.696 0.743	0.657 0.701
A05.496	0.306 0.286	0.304 0.413	C01.597	0.667 0.723	0.698 0.723
A05.585	0.717 0.697	0.755 0.722	C01.635*	0.583 0.501	0.555 0.497
A05.620	0.239 0.282	0.360 0.392	C01.791	0.534 0.611	0.567 0.623
A05.682*	0.861 0.810	0.752 0.796	C01.815*	0.457 0.399	0.489 0.409
A05.761*	0.500 0.432	0.564 0.499	C01.1217*	0.472 0.569	0.512 0.577
A05.820	0.417 0.476	0.289 0.334	C01.1615	0.389 0.443	0.355 0.452
A05.855*	0.652 0.718	0.578 0.591	C01.2376	0.826 0.761	0.773 0.697
A05.894	0.804 0.761	0.891 0.800	C04.289	0.306 0.286	0.327 0.291
A05.975	0.572 0.611	0.609 0.701	C04.351	0.778 0.786	0.696 0.775
A05.1254	0.279 0.299	0.317 0.342	C04.415*	0.472 0.515	0.505 0.543
BAMH1.331	0.196 0.174	0.231 0.202	C04.496	0.217 0.330	0.256 0.342
BAMH1.415	0.239 0.196	0.341 0.225	C04.544	0.348 0.477	0.389 0.514
BAMH1.497	0.712 0.784	0.754 0.798	C04.565	0.696 0.772	0.819 0.832
BAMH1.532	0.543 0.652	0.603 0.687	C04.705*	0.286 0.378	0.396 0.403
BAMH1.615	0.457 0.340	0.514 0.448	C04.967	0.222 0.298	0.201 0.279
BAMH1.732*	0.513 0.599	0.609 0.623	C04.1170*	0.592 0.501	0.611 0.577
BAMH1.825	0.876 0.778	0.765 0.698	C04.1852	0.761 0.744	0.679 0.778
			C04.2005	0.577 0.523	0.613 0.549

 Table 3 Mean heterozygosity for the 11 codominant alleles (2 alleles per locus) averaged across

 sample dates and RAPD loci for each population. Numbers in parentheses indicate standard

 errors.

Population	Heterozygosity
Piedmont Wheat	0.335 (0.045)
Piedmont Crucifer	0.416 (0.024)
Weld Wheat I	0.443 (0.030)
Weld Wheat II	0.425 (0.039)
Hort Farms Crucifer	0.392 (0.024)
BARI Wheat	0.412 (0.055)

Table 4 Estimates of Fst and θ for *D. rapae* by date for 1994 and 1995.

	Wright's Fst	θ	Lynch & Milligan's Fst	
Sampling Date	1994 1995	1994 1995	1994 1995	
June	0.070 0.065	0.060 0.068	0.064 0.071	
July	0.059 0.071	0.058 0.062	0.056 0.063	
August	0.066 0.074	0.070 0.073	0.061 0.069	

Table 5 Variance components combined across loci from the nested analysis of variance for D.*rapae* populations collected in Colorado during 1994 and 1995.

Sampling	Variance Components			
Date	Comparison	1994	1995	
····				
JUNE	Fields within Regions	0.619	0.613	
	Fields within the Total	0.424	0.501	
	Regions within the Total	-0.035	-0.059	
JULY	Fields within Regions	0.771	0.690	
	Fields within the Total	0.301	0.405	
	Regions within the Total	-0.070	-0.013	
AUGUST	Fields within Regions	0.809	0.771	
	Fields within the Total	0.299	0.316	
	Regions within the Total	-0.094	-0.073	

Figure 1. Examples of silver-stained PAGE gels demonstrating the inheritance patterns of codominant RAPD-PCR markers (a) and dominant RAPD-PCR markers (b). See text for details of haplodiploid Mendelian inheritance patterns.

Figure 2. Cluster diagrams showing the structure of *D. rapae* female populations in Colorado. Shown are Nei's standard genetic distances among populations of individuals collected during August 1994 and 1995. At each branching point is indicated the bootstrap support (out of 100) for that branch. (A). The pattern indicates that the population of *D. rapae* in northern Colorado is subdivided into many smaller populations, likely a result of limited dispersal, even between fields that are less than 1.0 km from each other (Piedmont wheat fields vs. Piedmont crucifer fields). (B). After collapsing the branching points with low support (less than 75 times out of 100), the pattern still indicates that fields are different from one another.



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