Separation of Three Species of *Ditylenchus* and Some Host Races of *D. dipsaci* by Restriction Fragment Length Polymorphism

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Abstract: This study examined the ribosomal cistron of Ditylenchus destructor, D. myceliophagus and seven host races of D. dipsaci from different geographic locations. The three species showed restriction fragment length polymorphisms (RFLPs) in the ribosomal cistron, the 18S rDNA gene, and the ribosomal internal transcribed spacer (ITS). Southern blot analysis with a 7.5-kb ribosomal cistron probe differentiated the five host races of D. dipsaci examined. Polymerase chain reaction (PCR) amplification of the ITS, followed by digestion with some restriction endonucleases (but not others), produced restriction fragments diagnostic of the giant race. Because the PCR product from D. myceliophagus and the host races of D. dipsaci was about 900 base pairs and the ITS size in D. destructor populations was 1,200 base pairs, mixtures of populations could be detected by PCR amplification. ITS fragments differentiated between D. dipsaci and Aphelenchoides rhyntium in mixed populations. This study establishes the feasibility of differentiation of the host races of D. dipsaci by probing Southern blots with the whole ribosomal cistron.

Key words: Ditylenchus destructor, D. dipsaci, D. myceliophagus, DNA, host race, internal transcribed spacer, nematode, polymerase chain reaction, restriction fragment length polymorphism, ribosomal DNA, sibling species.

Nematodes belonging to the genus Ditylenchus Filipjev are difficult to identify to species because of highly similar morphology and considerable intraspecific biological variation. Morphological differentiation of Ditylenchus is based mainly on tail shape and size, relative length of the stylet and post-vulval sac, and the number of cuticular lateral lines (10). However, these characters vary according to nematode developmental stage, culture medium, and temperature (2). The conclusion from a recent taxonomic review of the genus Ditylenchus was that identification is difficult and preparation of a workable key is almost impossible (3). Therefore, a sensitive and reliable technique is needed to differentiate species of Ditylenchus.

Ditylenchus dipsaci (Kühn) Filipjev, the stem nematode, is a migratory endoparasite of over 500 species of angiosperms (24). The main method of control of D. dipsaci is crop rotation, but the presence of morphologically indistinguishable races with different host preferences makes rotation difficult. Moreover, the presence of mixtures of nematodes, e.g., Aphelenchoides ritzemabosi and D. dipsaci in alfalfa fields, may confound diagnosis. A practical technique for rapidly and reliably identifying the host races of D. dipsaci is not available. Except for a larger "giant" race on field beans (13), the 30 host races of D. dipsaci (15) are separated on the basis of host preference. The races exhibit varying degrees of reproductive isolation, such as partial or complete reproductive incompatibility (6,21), and a wide range of chromosome numbers, from 2n = 24 to 56 (1).

Although protein electrophoresis separated species of *Ditylenchus* (5), attempts to separate races of *D. dipsaci* by esterase or catalase profiles (7,14) or by polyclonal antibodies developed against the surface antigens (11,28) failed. Palmer et al. (19) generated monoclonal antibodies (MAbs) against the oat race but discovered in an ELISA test that the MAbs were specific to only the original oat race isolate, not to all isolates of that race. Analysis of DNA is a more direct measure of variability than is protein analysis, and the application of

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such molecular analyses recently provided precise identification of three species of *Ditylenchus* but did not differentiate host races of *D. dipsaci* (18).

Ribosomal DNA has proven to be a particularly interesting and useful region for investigating reproductively isolated nematode populations at all taxonomic levels (9,27,29). The multiple copy, tandemly repeated, ribosomal gene cluster contains a wide spectrum of extremely conserved to highly heterogeneous sequence. The DNA sequences of the ribosomal DNA encoding 185, 5.8S, and 26S genes are more conserved than the internal transcribed spacer (ITS) or intergenic spacer (IGS) region. Thus, depending on the organism, regions can be identified that differentiate a range of taxa from the level of genera to that of subspecific populations (12). Although Southern hybridization is a reasonably sensitive method for surveying the fine and gross organization of the ribosomal gene, it has associated disadvantages; for example, it may require a large amount of DNA or radioactive techniques for visualization (4). Polymerase chain reaction (PCR) amplification of specific DNA fragments overcomes these disadvantages (17).

The objective of this study was to determine whether some of the homologous and heterologous regions of the ribosomal gene cluster, namely the whole rDNA cistron, the 18S gene, and the ITS, could separate three species of *Ditylenchus* and nine races of *D. dipsaci*.

MATERIALS AND METHODS

The nematode isolates used in this study originated from North America, Europe, and Africa from cultivated and noncultivated plant hosts (Table 1). Some populations were derived directly from the field and others had been in culture for a number of years. *Aphelenchoides rhyntium* (APH) was maintained on *Botrytis cinerea* culture in the laboratory.

Extraction and restriction of DNA: Nematodes were washed from the lids of fungal plates or extracted from nematodeinfested dried or fresh plant material in a Baermann funnel, and the nematodes were screened from the aqueous suspension with a 30 µm-pore-d sieve. The nematode DNA was extracted (29). Initially, Southern blots of digested Ditylenchus genomic DNA were probed with the whole ribosomal cistron and then reprobed with a portion of the ribosomal gene. Although there was not enough DNA from some D. dipsaci populations to perform Southern hybridizations, all populations were amplified by PCR and subsequently digested with restriction enzymes. Genomic or PCR-amplified DNA samples (0.5 to 1.0 µg) were digested by restriction endonucleases with four, five, or six-base recogni-

TABLE 1. Origin of the Ditylenchus populations used in this study.

	Code	Host	Origin	Source
D. myceliophagus	MYC	Mushroom	Germany	Rhizoctonia cerealis
D. destructor	UK	Solanum tuberosum	Ireland	R. cerealis
	WIS	S. tuberosum	U.S. (Wisconsin)	Excised corn roots
D. dipsaci				
alfalfa	ALF	Medicago sativa	U.S. (Washington)	Greenhouse alfalfa plants
alfalfa	FAL	M. sativa	France	Alfalfa callus
beet	BET	Beta vulgaris	France (Alsace)	Greenhouse faba beans
giant	FGI	Vicia faba	France (Le Rheu)	Greenhouse faba beans
giant	GIA	V. faba	England (Hertfordshire)	Dried zucchini marrows
potato	POT	S. tuberosum	Netherlands	Dried potatoes
red clover	RCL	Trifolium pratense	France (Domagne)	Greenhouse red clover plants
teasel	TEA	Dipsacus fullonum	England (Somerset)	Dried teasel stems
tulip	TUL	Tulipa gesneriana	Netherlands	Dried tulip bulbs

tion sites, according to the manufacturers' recommended procedures (Bethesda Research Laboratories [BRL], Burlington, Ontario; Boehringer Mannheim, Germany; Pharmacia, Baie d'Urfé, Quebec). DNA samples were electrophoresed in horizontal gels (16) of 0.7 to 1.5% agarose, depending on the size of the fragments. Fragment sizes were subsequently estimated with a regression line of size standards and electrophoretic migration. We used a 1-kb ladder (Pharmacia) as size standards.

Southern blots: Two probes (29) were used in Southern blots. The 7.5-kb ribosomal gene fragment from the Bursaphalenchus xylophilus Ibaraki isolate (pBx2) comprises the ribosomal cistron from the 5' end of the 18S coding unit to the 3' end of the 28S coding unit. The 18S probe, from B. mucronatus (pBm3), is a 1.1-kb fragment that contains the entire 18S subunit coding region. The DNA probes used for Southern blots were ³²P-labeled using random oligonucleotide primers (8).

From 1.0 to 2.0 μ g of Rsa I-digested genomic DNA was electrophoresed and Southern blotted to nylon filters (16). Filters were hybridized to denatured rDNA probe labelled with [³²P]dATP at 62 C in 5 × SSPE (1 × SSPE = 0.18 M NaCl, 10 mM (Na_{1.5}) PO₄, 1 mM Na₂EDTA, pH 7.4), 5 × Denhardts (1 × Denhardt's = 0.02% w/v bovine serum albumin, Ficoll 400 and polyvinylpyrrolidone 40) and 2.0% SDS (sodium dodecyl sulfate). The filters were washed four times at 62 C in 2 × SSPE and 0.2% SDS.

For the 18S Southern hybridizations, the nylon filters previously hybridized with the rDNA probe were stripped of the probe by washing twice for 5 minutes in a solution of 1.5 M NaCl and 0.5 M NaOH at room temperature (ca 20 C) and neutralizing for 30– 60 minutes in a solution of 1.0 M ammonium acetate and 0.02 M NaOH, before hybridization with the 18S probe.

Polymerase chain reaction: The ITS region (including ITS 1, ITS 2, and the 5.8S subunit) in 100–200 ng of genomic DNA from each population was amplified with kits, according to the manufacturers' recommendations (Perkin-Elmer Cetus, Eden Prairie, MN; Promega, Madison, WI). The primers used were two universal ITS primers, each 21 base pairs long, developed from *Xiphinema bricolensis* (27); one primer with a sequence starting 171 base pairs from the 3' end of the 18S ribosomal subunit, and the other primer 80 base pairs into the 5' end of the 26S gene.

Negative controls, consisting of the mixture without template DNA but brought to the same volume, were run with each amplification. Reaction profiles were as follows: 1.5 minutes at 96 C, 30 seconds at 50 C, 4 minutes at 72 C; 40 cycles of 45 seconds at 96 C, 30 seconds at 50 C, 4 minutes at 72 C and a final cycle of 45 seconds at 96 C, 30 seconds at 50 C, and 10 minutes at 72 C. The PCR thermocycler used was the Twin Block System EC Cycler (Ericomp). The PCR products from several amplification runs were pooled for digests.

RESULTS

Differentiation of Ditylenchus species: All three nematode species displayed a unique pattern of fragments when Southern hybridizations were probed with the complete ribosomal cistron (Fig. 1). Only a limited amount of DNA was available from D. destructor (WIS), D. dipsaci (BET, FAL, FGI, RCL), and A. rhyntium (APH) populations, and these generated fainter fragment patterns than others. When the described Southern transfer of the digested DNA of D. myceliophagus, D. destructor (UK), and D. dipsaci teasel (TEA) was reprobed with the B. mucronatus fragment containing most of the 18S rDNA gene, D. dipsaci showed a unique pattern of fragments, but identical patterns occurred for D. myceliophagus and D. destructor (data not shown).

Amplification by PCR of DNA from *D.* myceliophagus and all populations of *D.* dipsaci generated a single ITS fragment of about 0.9 kb, whereas the PCR product of *D. destructor* isolates was about 1.2 kb (Fig. 2A). A PCR amplification of a mixed sam-



FIG. 1. Autoradiograph of a Southern transfer of Rsa I-digested genomic DNA from *Ditylenchus myceliophagus, D. destructor,* and five host races of *D. dipsaci* probed with the 7.5-kb rDNA cistron (from *Bursaph elenchus xylophilus*) under moderately stringent conditions. A longer autoradiographic exposure was performed for *D. dipsaci* (12 hours) than for *D. mycelioph agus* and *D. destructor* (8 hours). Populations abbreviated as in Table 1.

ple of *D. dipsaci* (POT) and *D. destructor* (UK) DNA generated respective bands of 0.9 and 1.2 kb, (Fig. 2B). Similarly, PCR amplification of a mixture of *D. dipsaci* (ALF) and *A. rhyntium* (APH) DNA produced two bands corresponding to each genus in the mixture (Fig. 2B).

All enzymes that restricted the ITS fragment provided different patterns for each of the three species of *Ditylenchus* (Figs. 3–6, Table 2). In general, few bands were shared among the species, and the three species were distinguished readily from each other by their RFLPs.

Differentiation of host races of D. dipsaci: Each of five populations of D. dipsaci produced unique fragment patterns when hybridized to a heterologous rDNA cistron (Fig. 1). Three bands at 1.8, 0.6, and 0.5 kb characterized all D. dipsaci populations. Some band sizes (e.g., 2.8 kb, 1.2 kb, and 1.0 kb) were shared between some populations, and others were unique to one D. *dipsaci* isolate examined (e.g., 1.4 kb for the giant race and 0.8 kb for the alfalfa race). When reprobed with the 18S probe, the fragment patterns were identical for four of the *D. dipsaci* host races (ALF, POT, TEA, and POT) and could be distinguished from the giant race pattern by an extra band at 1.4 kb (data not shown).

In order to assess the variability in the PCR-amplified ribosomal spacer region, initially a battery of 14 enzymes was used to digest the ITS fragment of D. dipsaci TEA and ALF. The results of the restriction analysis demonstrated similar banding patterns between these two races (Table 2). The ITS of all the D. dipsaci races including the giant races was digested with seven enzymes, namely, Dde I, Hae III, Hinc II, Hinf I, Hpa II, Pst I, and Rsa I. Digestion of the ITS of each host race, except the two populations of the giant race, produced patterns identical with those of the teasel and alfalfa races. Although not all enzymes, e.g., Pst I and Dde I, showed differences between the giant race and the other D. dipsaci races (Fig. 3), four of the seven enzymes tested (Hae III, Hpa II, Hinf I, and Rsa I) generated patterns unique to this giant race (GIA), and different from those characteristic of all other populations of D. dipsaci, as well as from the other two Ditylenchus species tested (Figs. 4-6). Not all gels are shown, but all fragment sizes generated by the 14 restriction enzymes are tabulated (Table 2).

DISCUSSION

This investigation expands on an earlier study in which Palmer et al. (18) probed Southern hybridizations with random DNA fragments isolated from two libraries, one from *D. myceliophagus* and the other from *D. dipsaci* oat race. Their probes could differentiate between these two species but not define unknown species of *Ditylenchus*.

In the initial Southern blot experiments in the present study, the 7.5-kb rDNA probe derived from *Bursaphelenchus* revealed sequence variation between each species of *Ditylenchus* and between host



FIG. 2. A) Agarose gel of the internal transcribed spacer (ITS) amplified by polymerase chain reaction (PCR) from *Ditylenchus myceliophagus* (MYC), *D. destructor* (WIS, UK), and host races (BET, TUL, GIA, FAL, ALF, TEA) within *D. dipsaci*. B) Two agarose gels showing the ITS fragments resulting from PCR amplification of DNA from *D. myceliophagus* (MYC), *D. dipsaci* (DIP POT), *D. dipsaci* and *D. destructor* (DES UK) DNA mixed together, *D. destructor* (DES UK), water (no DNA), *Aphelenchoides rhyntium* (APH), *D. dipsaci* (DIP ALF), and *A. rhyntium* and *D. dipsaci* mixed. Populations abbreviated as in Table 1. Marker = 1 kb ladder.

races of *D. dipsaci* within the ribosomal cistron. However, closer examination using only the 18S portion of the ribosomal probe failed to distinguish between the three species and between the five host races (except for the giant race). This failure is not unexpected, because this 18S gene is one of the most conserved regions in the ribosomal gene cluster (12).

Each of the three species of Ditylenchus



FIG. 3. Agarose gel of the PCR-amplified internal transcribed spacer (ITS) of *Ditylenchus myceliophagus* (MYC), *D. destructor* (WIS, UK), host races within *D. dipsaci* (abbreviated as in Table 1), and *Aphelenchoides rhyntium* (APH) digested with the restriction enzyme Pst I. Populations of *D. dipsaci* abbreviated as in Table 1. Marker = 1 kb.

were readily separated from each other and A. rhyntium by PCR amplification of the ITS (including the 5.8S gene) and restriction with an appropriate enzyme. Restriction of the ITS fragment produced four groups: D. myceliophagus (MYC), two populations of D. destructor (WIS and UK), the giant race of D. dipsaci, and the other diploid host races of D. dipsaci.

The ITS of the ribosomal cistron in *Ditylenchus* (and *A. rhyntium*) is an appropriate region for species separation but does not differentiate the host races. The host races of *D. dipsaci* were also closely related in another study (18), in which a *D. dipsaci* oat race probe in dot blot experiments did not differentiate the races. In our study, restriction digests of the ITS with 14 different enzymes yielded identical restriction patterns for all but one of the examined races. The polyploid giant race had characteristic patterns of fragments in four (Rsa I, Hpa II, Hae III, Hinf I) of seven enzymic digests.

Sturhan (22,23) suggested that the giant race is not a mere host race but should be considered a sibling species, because it did not produce fertile progeny when crossed with one of the diploid races. However, in another study, fertile F_1 progeny resulting from such crosses were observed (15). Because polyploidy is suspected of playing an important role in the cytogenic evolution of some amphimictic groups of nematodes (26), the hypothesis that the giant race is a sibling species is strengthened. The rDNA sequence heterogeneity presented here also supports this sibling species status.

Mixtures of *D. dipsaci* and *A. ritzemabosi* often coexist in alfalfa (*Medicago sativa*) fields; indeed, the alfalfa samples collected in Wyoming showed typical damage from *D. dipsaci* and contained a mixture of these



FIG. 4. Agarose gel of the PCR-amplified internal transcribed spacer (ITS) of *Ditylenchus myceliophagus* (MYC), *D. destructor* (WIS, UK), host races within *D. dipsaci* (abbreviated as in Table 1), and *Aphelenchoides rhyntium* (APH) digested with the restriction enzyme Hae III. The 0.9-kb fragment in GIA is a consequence of a partial digest. Marker = 1 kb ladder.



FIG. 5. Agarose gel of the PCR-amplified internal transcribed spacer (ITS) of *Ditylenchus myceliophagus* (MYC), *D. destructor* (WIS, UK), host races within *D. dipsaci* (abbreviated as in Table 1), and *Aphelenchoides rhyntium* (APH) digested with the restriction enzyme Hinf I. The faint bands with sizes larger than 350 kb in BET, GIA, FAL, and TEA are the result of a partial digest. Marker = 1 kb ladder.

two species (F. Gray, pers. comm.), which are relatively similar under low magnification. The ITS of A. *rhyntium*, a close relative of A. *ritzemabosi*, could be distinguished in a mixture of this species and D. *dipsaci*. Therefore, this method could be useful to confirm the morphological diagnosis of mixed or pure infestations of nematodes.

The unique patterns of fragments produced from restriction analysis of the ITS



FIG. 6. Agarose gel of the PCR-amplified internal transcribed spacer (ITS) of *Ditylenchus myceliophagus* (MYC), *D. destructor* (WIS, UK), host races within *D. dipsaci* (abbreviated as in Table 1), and *Aphelenchoides rhyntium* (APH) digested with the restriction enzyme Hpa II. The 0.9-kb fragment in GIA is the result of a partial digest. Marker = 1 kb ladder.

TABLE 2. Fragment sizes (kb) resulting from digestion of the internal transcribed spacer of *Ditylenchus myceliophagus*, *D. destructor* (UK and WIS races), and host races and the giant race of *D. dipsaci* with 14 restriction enzymes. TABLE 2. Continued

	D. dipsaci			
Enzyme	Host races	Giant race	D. destructor	D. myceliophagu
Acc I		-†	1200	
4.1 Y	900			900
Alu I	900		870	900
			290	
BamH I		_	1000	
				900
	340			
	220			
Dde I	180		670	
Due I			570	
	310	310	0.0	
				300
	290	290		
	900	900		250
	200	200		130
Dra I		_	1200	100
2.4.2				900
	340			
	250			
Hae III	900	800		
		800	450	450
		200	450	200
			170	
Hinc II			900	900
	800	800		
II. CI			250	
Hinf I			780	630
	440			000
	350	350		
				310
	150	150	180	
Una 11	150	150	1000	
пран			1000	900
		600		000
	320			
	200	200		
X 7 . 7	180		1000	
Nsi I	000		1200	000
Pet I	900		850	900
1311	650	650	000	
				620
	400	400	400	400
Rsa I			6 000	900
		400	600	
		490		

	D. d	ipsaci			
Enzyme	Host races	Giant race	D. destructor	D. myceliophagus	
	450	450			
	250		250		
			170		
	140				
Sau 3A		-	540		
				440	
			400		
	340				
	260				
	200				
			180		
	110				
	100			100	
TaqI		-	640		
	340				
				320	
				260	
	230				
			200	100	
				160	
			150		
	130				

† A dash in the column indicates that this experiment was not performed.

f D. myceliophagus, D. destructor, and D. *ipsaci* are not confused by morphological imilarities between species or geographic opulations, or by host- and temperaturenduced morphological differences within species. Although the host races of D. *ipsaci* examined in this study generally riginated from different geographic locaions and different food sources, Southern vbridizations utilized herein revealed no lifferences, except for the giant race. The wo populations of the giant race from Enland and France generated identical paterns of fragments with three enzymes ested (Pst I, Hinf I, Hpa 2). Similarly, U.S. nd French isolates of the alfalfa race conistently showed the same ITS restriction patterns. In addition, because PCR ampliication has been used successfully on preerved organisms (25), taxonomic features hat are obscured by fixative techniques vill not impede the molecular diagnosis of preserved museum specimens, although uch diagnosis is destructive.

Analysis of the ribosomal cistron by

Southern hybridizations and restriction analysis of fragments amplified by PCR indicates that *Ditylenchus* species and the host races of *D. dipsaci* can be identified using DNA techniques. The use of the ribosomal cistron provides a powerful tool for studying the systematics of unknown species, as well as for the identification of known species in a mixed population of host races in the nematode genus *Ditylenchus*.

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