Rapid identification of *Meloidogyne chitwoodi*, *M. hapla*, and *M. fallax* using PCR primers to amplify their ribosomal intergenic spacer

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**Summary** - Primers used to amplify the Internal Transcribed Spacer (ITS) were reoriented divergently and used to amplify the genomic rDNA intergenic spacer (IGS) region of *M. chitwoodi* via the polymerase chain reaction. The location of the IGS within the resulting 5.8 kilobase pair DNA fragment was identified by subcloning individual restriction fragments and sequencing their terminal regions. Additional primers were constructed which could amplify the IGS with only minimal flanking sequences of the 28S and 18S rDNA. Agarose gel electrophoresis of amplified fragments obtained from PCR amplification of DNA from *M. chitwoodi*, *M. hapla*, and *M. fallax* demonstrated size polymorphisms. Direct identification of nematodes based on size polymorphisms of an amplified product is more efficient than the usual procedure of amplification followed by digestion with restriction endonucleases.

**Résumé** - Identification rapide de *Meloidogyne chitwoodi*, *M. hapla* et *M. fallax* par utilisation d’amorces d’amplification en chaîne par polymérase en vue d’amplifier leurs espaces intergéniques - Des amorces servant à amplifier la région interne transcrite de l’espacement (ITS) ont été réorientées de manière divergente pour amplifier par réaction en chaîne par polymérase (PCR) l’espacement intergénique (IGS) de l’ADNr génomique de *Meloidogyne chitwoodi*. La position de l’IGS dans le fragment d’ADNr obtenu de 5,8 Kb a bien été identifiée par sous-clonage individuellement des fragments de restriction et en séquençant leurs régions terminales. Des amorces additionnelles ont été construites pour amplifier l’IGS avec un minimum de séquences des gènes 28S et 18S de l’ADNr. Les fragments amplifiés par PCR de l’ADNr de *M. chitwoodi*, *M. hapla* et de *M. fallax* ont montré des tailles différentes par électrophorèse sur gel d’agarose. L’identification directe des nématodes à partir de différences de taille des fragments amplifiés est plus efficace que la méthode usuelle d’amplification suivie de digestion à l’aide d’enzymes de restriction.

**Key-words** : amplified fragment length polymorphism, intergenic ribosomal spacer (IGS), *Meloidogyne*, nematodes, polymerase chain reaction (PCR), taxonomy.

A few species of root-knot nematodes from the genus *Meloidogyne* are the most damaging parasitic nematodes in worldwide agriculture (Sasser, 1980). *Meloidogyne hapla* is common in North America and in Europe. The distribution of *M. chitwoodi*, and its three recently described races (Golden et al., 1980; Santo & Pinkerton, 1985; Mojtabahdi & Santos, 1994), is being verified in some European and other countries. *M. chitwoodi* is also common in several western states of the USA but it has not yet been found in Canada, where a quarantine is in effect against this species. A third species, *M. fallax* has just been described from Dutch populations previously identified as *M. chitwoodi* (Van Meggelen et al., 1994; Karssen et al., 1996).

Nematode identification usually requires morphometric measurements of juveniles and mature females, as well as differential host range studies. Although reliable, these methods remain both specialized and time consuming. However, recently developed molecular techniques are simplifying and expediting species identification in the genus *Meloidogyne*. Castagnone-Sereno et al. (1993) separated *M. arenaria*, *M. incognita*, *M. javanica*, and *M. hapla*, using highly repetitive DNA sequences such as satellite DNA. Other advances rely on polymerase chain reaction (PCR) amplification of a specific sequence and restriction fragment length polymorphisms (RFLPs) in that sequence among the species of interest. The advantages of PCR technology in nematode identification are rapidity, since results are obtained in a few hours, and most importantly, that only one or very few juvenile specimens are required (Vrain & McNamara, 1994). Accordingly, Powers and Harris (1993) could identify *M. incognita*, *M. javanica*, *M. arenaria*, *M. hapla*, and *M. chitwoodi*, from RFLP analysis of a mitochondrial intergenic region, while Zijlstra et al. (1995) separated *M. chitwoodi* from *M. hapla*, *M. fallax*, *M. incognita*, and *M. javanica*, using RFLPs in the ITS of ribosomal DNA (rDNA) repeats.

A further advance of these methods would be to eliminate the need for restriction enzyme digestion, thus directly separating nematode species by the size of the PCR amplified DNA fragment. This would require identifying and amplifying a region of variable size between species. The InterGenic Spacer (IGS) of genomic rDNA is a highly variable region in nematodes and other organisms (Simeone et al., 1982; Gerbi, 1985; Webster et al., 1990; Vahidi & Honda, 1991), often characterized by the presence of multiple short repeated
sequences. We hypothesized that the variable length of rDNA IGS sequences could be used to separate species of root-knot nematodes. Our objective was to design primers that would amplify the IGS regions of *M. hapla* and *M. chitwoodi*, and the newly described species, *M. fallax*, and identify these species using the variable length of the amplified IGS fragment.

**Materials and methods**

**Nematodes**

The populations of *M. chitwoodi*, *M. hapla*, and *M. fallax* used in this study are listed in Table 1. Genomic DNA was purified from nematode cultures as previously described (Vrain et al., 1992; Zijlstra et al., 1995).

**Table 1. Populations of Meloidogyne chitwoodi, M. hapla, and M. fallax used in the study.**

<table>
<thead>
<tr>
<th>Population</th>
<th>Location</th>
<th>Host</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. chitwoodi</em></td>
<td>New Mexico</td>
<td>Cotton</td>
</tr>
<tr>
<td><em>M. hapla</em></td>
<td>Michigan</td>
<td>Cotton</td>
</tr>
<tr>
<td><em>M. fallax</em></td>
<td>California</td>
<td>Cotton</td>
</tr>
</tbody>
</table>

**Polymerase Chain Reaction**

Amplifications of DNA targets greater than 5 kb were performed using 2 μl eLONGase Enzyme Mix (GibcoBRL, Burlington, ON, Canada) in 30 μl PCR reactions containing 60 mM Tris-SO₄ (pH 9.1), 15 mM (NH₄)₂SO₄, 1.6 mM MgSO₄, 200 μM dNTPs, 500 nM each primer, and 100 ng genomic DNA. Cycling parameters consisted of 30 s denaturing for 15 s, 50 °C annealing for 30 s, and 72 °C extension for 8 min. DNA targets of less than 3 kb were amplified using 1 unit of Taq DNA polymerase (Appligene-Oncor, France) in 30 μl PCR reactions containing 10 mM TrisHCl (pH 9.0), 50 mM KCl, 1.5 mM MgCl₂, 0.1 % Triton X100, 0.2 mg/ml BSA, 65 μM dNTPs, 500 nM each primer, and 10 ng genomic DNA. Cycling parameters consisted of 95 °C denaturing for 30 s, 46 °C annealing for 45 s (3 cycles), 52 °C annealing for 45 s, (37 cycles), and 72 °C extension for 2 min. All PCR reactions were performed in a robocycler gradient 96 (Stratagene, La Jolla, CA, USA).

**Oligonucleotide Synthesis**

Primer sequences previously employed for amplification of the rDNA ITS region (Vrain et al., 1992) were used as a basis for designing a new primer set. The inverted primers G: 5'-TCCCTTAGTAACGCGGAGTG-3', and B: 5'-TGTAACAGGCAGGGGACG-3' (Fig. 1) were capable of amplifying a portion of the rDNA cistron (an approximately 5.8 kb DNA fragment) that included the IGS. Additional primers to the 3' end of the 28 S gene (283-N: 5'-TTCGAGTAACGCCGGTTAACAG-3'), and to the 5' end of the 18 S gene (185-0: 5'-CAGTTCAGGCAGGATCACA-3'; 185-1: 5'-GTGAACACACACTCCTCATC-3') were used to amplify the IGS region alone (Fig. 1). All oligonucleotide primers were synthesized by phosphoramidite chemistry using the Oligo1000M (Beckman).

**Cloning**

Amplified DNA from *M. chitwoodi* was separated from the genomic DNA template and primers by electrophoresis using 1 % (w/v) low-melting point agarose.

![Fig. 1. Structural map of the rDNA cistron of Meloidogyne chitwoodi. The hybridization positions of oligonucleotide primers are shown on the map, and the orientation of each primer for amplifying the IGS via PCR is indicated by an arrow.](image-url)

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NTPs, A, C, G, or 15 s, 8 min, using 1 μl Oncor, 10 mM MgCl₂, 0.1% Nonidet P-40, 0.01 M Tris-HCl, pH 9.0, and 100 nM primers. The reaction was carried out at 94 °C for 45 s, 34 cycles, with consequent 45 s annealing at 46 °C for 45 s, and 1 PCR cycle of 10 min at 72 °C. The PCR products were purified using 0.7% agarose gel electrophoresis in Tris-acetate EDTA buffer at 80 V for 1 h. DNA bands were excised from gels with a sterile scalpel and purified from the agarose using a freeze-thaw procedure (Qian & Wilkinson, 1993). The purified fragments were cloned into the Smal site of pSK' (Stratagene, La Jolla, CA, USA) by blunt-end cloning using T4 DNA ligase (GibcoBRL, Burlington, ON, Canada) according to manufacturer's directions. Ligation products were transformed into CaCl₂ competent *Escherichia coli* by a brief heat shock at 42 °C, and cells selected for both resistance to ampicillin and inability to utilize the chromogenic substrate X-gal (lacZ') (Miller, 1972). Alkaline lysis was used to purify recombinant plasmids from single colonies grown overnight in Terrific Broth (Sambrook et al., 1982).

**DNA Sequencing**

Cloned restriction fragments were further purified for sequencing using Qiaquick spin columns (Qiagen Inc., Chatsworth, CA, USA), eluted in 10 mM Tris-HCl, pH 8.0, and stored frozen until further use. Sequencing was accomplished using the PRISM™ DyeTerminator Cycle Sequencing Ready Reaction Kit (Perkin Elmer, Mississauga, ON, Canada) and analyzed by ABI310 capillary electrophoresis system according to the manufacturer’s directions. The sequencing data obtained for the 28S, ITS, and 18S regions were compared with other nucleotide sequences in the GenBank database (http://www.ncbi.nlm.nih.gov) to find homologous regions so that additional primers could be designed to more efficiently amplify the IGS region.

**Results**

Reorientation and modification of the ITS primers allowed successful PCR amplification of a 5.8 kb region of the rDNA cistron of *M. chitwoodii* when a proof-reading *Taq* polymerase was used. Restriction mapping of the cloned fragments obtained from amplifications of both *M. chitwoodii* race 2 and *M. chitwoodii* race 3 allowed further subcloning and DNA sequencing of fragments contained within the 5.8 kb amplicon. Comparisons of nucleotide sequences with other nematodes or related rDNA sequences present within the GenBank database revealed regions to which primers might be designed for amplification of the IGS region from a variety of nema-tode species.

Upon testing of the newly designed primers 283-N and 185-0 (Fig. 1), successful PCR amplification of an approximately 2.6 kb fragment from all three described *M. chitwoodii* races was achieved (Fig. 2). This amplification utilized a more common form of thermostable *Taq* polymerase, and reaction times were reduced by nearly 75%. Additionally, amplified product yields were substantially increased, allowing product from amplification of single juveniles to be visualized by ethidium bromide staining of reactions separated by gel electrophoresis. The rDNA IGS region from several *Melido-gyne* species was amplified and compared using the 283-N/185-0 primer combination. Separation of the amplified products obtained from *M. chitwoodii*, *M. fallax*, and *M. hapla* demonstrated size differences in the amplified product (Fig. 2). The products from *M. chitwoodii* and *M. fallax* differ by approximately 100 bp (2.6 vs 2.7 kb), while the corresponding band from *M. hapla* is much smaller (600 bp). These amplified fragment length polymorphisms were reproducibly obtained from three or more populations of each species tested.

To confirm that each fragment corresponded to the correct amplicon, two further tests were performed as follows. A fifth primer (185-1) was synthesized, hybridizing approximately 200 bp upstream of the 185 5' terminus (Fig. 1), to amplify the IGS region when employed in combination with the 283-N primer. For each species, a band approximately 200 bp smaller than the original 283-N/185-0 product was obtained (Fig. 3). This result was expected if the amplified regions were...
indeed the IGS for which the primers were designed. In addition, we also sequenced the terminal regions of each amplified fragment. The DNA sequences revealed that each band was indeed the product of amplification using both primers contained in the PCR reaction, and that the amplified region exhibited some homology to rDNA sequences of nematodes or other organisms contained within the Genbank database.

**Discussion**

The study of RFLPs in nematode taxonomy routinely involves combining PCR amplification of the rDNA ITS with subsequent restriction enzyme digestion. Our procedure advances upon this technique in three main aspects:  

1. Amplified fragment length polymorphisms can allow single step discrimination of species identity;  
2. Processing time is substantially reduced since restriction enzyme incubation periods are avoided, and only one gel need be run; and  
3. The procedure is less costly since no restriction enzymes are employed, and lower agarose concentrations may be used for electrophoretic detection.

Using the single step protocol, we confirmed the recent reclassification of Baexem-type isolates as a new species (*M. fallax*) distinct from *M. chitwoodii* (Zijlstra et al., 1995). Hence, the sensitivity of molecular methods can identify genetic distinctions even though physiological or host range differences are not yet described. Previously, the potato cyst nematodes *Globodera pallida* and *G. rostochiensis* were directly distinguished by PCR amplification of the spliced leader RNA and SS rRNA spacer region (Stratford & Shields, 1994). Amplified fragment length polymorphisms were also observed within mitochondrial DNA of several *Meloidogyne* species, but did not distinguish *M. chitwoodii* from *M. hapla* (Powers & Harris, 1993). We now routinely use the size differences in the rDNA IGS as a diagnostic tool to differentiate these two species. Sequencing of the rDNA IGS is currently underway in order to produce species-specific probes (Petersen et al., 1995) for detection of quarantined nematodes on imported agricultural commodities.

The presence of a SS gene within the rDNA IGS may contribute to size differences between species (Bloks et al., 1997); however, the small size of the *M. hapla* IGS amplification product suggests a large deletion. Genetic changes in the rDNA IGS are not uncommon; they are readily accommodated by the host since this DNA region is not transcribed (Gerbi, 1985). Indeed, IGS deletions have already been found in *M. arenaria* (Vahidi & Honda, 1991). The presence of such a deletion, found in *M. hapla* isolates from two different continents, is in agreement with morphometric and phylogenetic classification of *M. hapla* as evolving separately from *M. chitwoodii* and *M. fallax* (Golden et al., 1980; Zijlstra et al., 1995).

Nevertheless, our results do not negate the possibility of full length cistrons present within *M. hapla*, since variations within the tandemly repeated rDNA copies can exist within individuals (Williams et al., 1985; Gerbi, 1985; Vahidi & Honda, 1991), and the kinetics of the PCR reaction can favor smaller amplions, resulting in their dominating the final reaction products. Analysis of additional populations and *Meloidogyne* species is necessary to determine whether IGS size polymorphisms are truly species-specific.

The inverse of primers B and G (Fig. 1) that have been used in several published studies of nematode ITS variability were originally designed from homologous sequences between several nematodes and other organisms (Vrain et al., 1992). We have not been able to amplify the IGS region of nematode species from several genera (*Xiphinema*, *Globodera*, *Heterodera*, *Steinernema*, and *Pratylenchus*; data not shown) with primers.
283-N and 185-0. These primers were designed from sequences at the 3' end of the 28S gene and the 5' end of the 18S gene of *M. chitwoodi*, regions lacking the high sequence conservation observed at the other end of each gene. Therefore, we are presently acquiring the nucleotide sequences, at the 3' end of the 28S and the 5' end of the 18S genes of the genera of interest, to design common primers that can amplify the IGS of all species to be compared.

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References


