SSCP-based identification of members within the *Pseudoterranova decipiens* complex (Nematoda: Ascaridoidea: Anisakidae) using genetic markers in the internal transcribed spacers of ribosomal DNA

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**SUMMARY**

The anisakid nematodes morphologically corresponding with *Pseudoterranova decipiens sensu lato* (s.l.) (Krabbe, 1878) from different seal or sea lion hosts and geographical origins, previously identified as *Pseudoterranova krabbei*, *P. decipiens* (s.s.), *P. bulbosa*, *P. azarasi* and *P. cattani* by multilocus enzyme electrophoresis, were characterized using a DNA approach. Also a population of *P. decipiens* (s.l.) from *Chaenocephalus aceratus*, the blackfin icefish, from Antarctica and another from *Osmerus eperlanus*, the European smelt, from Germany were included in the study. The first (ITS-1) and second (ITS-2) internal transcribed spacers (ITS) of ribosomal DNA (rDNA) were amplified by PCR from individual nematodes and analysed by single-strand conformation polymorphism (SSCP), followed by selective sequencing. While no variation in single-stranded ITS-1 and ITS-2 profiles was detected among samples representing each of the species or populations (with the exception of slight microheterogeneity), SSCP analysis of the ITS-2 amplicons allowed the unequivocal differentiation of all of the 5 sibling species of *P. decipiens* (s.l.) examined, which was supported by sequence differences in ITS rDNA. Samples representing the *P. decipiens* (s.l.) population from *O. eperlanus* had the same SSCP profile as those of *P. decipiens* (s.s.), which was supported by a lack of nucleotide difference in the ITS between them, suggesting that the former represented *P. decipiens* (s.s.). Based on SSCP results and ITS sequence data, *P. decipiens* (s.l.) from *C. aceratus* was genetically most distinct with respect to all other members of *Pseudoterranova* examined, which indicated that it may represent *P. decipiens* E (based on geographical origin) or a distinct species. These findings and the molecular approach taken should have important implications for studying the life-cycles, transmission patterns, epidemiology and population genetics of these anisakid nematodes, and the diagnosis of their infections.

Key words: internal transcribed spacer, polymerase chain reaction, *Pseudoterranova decipiens sensu lato*, ribosomal DNA, single-strand conformation polymorphism, species identification.

**INTRODUCTION**

The morphospecies *Pseudoterranova decipiens sensu lato* (s.l.) (Krabbe, 1878) is a common anisakid nematode of many aquatic hosts and has global geographical distribution (Palm et al. 1994; Palm, 1999; Paggi et al. 2000). Crustaceans, such as copepods, amphipods, shrimps and isopods, serve as intermediate hosts. Fish of various species serve as intermediate or paratenic hosts, and pinnipeds, including seals and sea lions, serve as definitive hosts (Adams, Murrell & Cross, 1997; Palm, 1999; Anderson, 2000). As seals are regarded as the most frequent definitive hosts, and cod among the most frequent intermediate or paratenic hosts, *P. decipiens* (s.l.) is also referred to as the ‘sealworm’ or ‘codworm’. Importantly, humans can also become infected with *P. decipiens* (s.l.) larvae by eating raw or inadequately cooked fish, such as cod (Desowitz, 1986; Kliks, 1986; Oshima, 1987; Adams et al. 1997). The disease is referred to as codworm anisakiasis or pseudoterranoviasis and is recognized as an important fish-borne zoonosis (Oshima, 1987;
Table 1. Adult and larval nematodes representing *Pseudoterranova decipiens* (s.l.), including *Pseudoterranova krabbei* (Pk), *P. decipiens* (s.s.) (Pd), *P. bulbosa* (Pb), *P. azarasi* (Pa), *P. cattani* (Pc) and 2 previously uncharacterized populations (PdCa and PdOe) from different host species

<table>
<thead>
<tr>
<th>Species/population</th>
<th>Sample code</th>
<th>No. of samples</th>
<th>Host species</th>
<th>Geographical origin</th>
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<tr>
<td>Previously characterized by multilocus enzyme electrophoresis</td>
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<td>Pk1-Pk15</td>
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<tr>
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<td>Pb1-Pb10</td>
<td>10</td>
<td><em>Eringnathus barbatus</em> (Bearded seal)</td>
<td>Newfoundland, Canada</td>
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<tr>
<td><em>P. azarasi</em></td>
<td>Pa1-Pa8</td>
<td>8</td>
<td><em>Eumetopias jubata</em> (Steller's sea lion)</td>
<td>Iwanai, Japan</td>
</tr>
<tr>
<td><em>P. cattani</em></td>
<td>Pc1-Pc8</td>
<td>8</td>
<td><em>Otaria byronia</em> (South American sea lion)</td>
<td>Concepción, Chile</td>
</tr>
<tr>
<td>Previously uncharacterized by multilocus enzyme electrophoresis</td>
<td></td>
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<td>PdCa1-PdCa29</td>
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<td><em>Chaenocephalus aceratus</em> (Blackfin icefish)</td>
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<td><em>P. decipiens</em> (s.l.)</td>
<td>PdOe1-PdOe27</td>
<td>27</td>
<td><em>Osmerus eperlanus</em> (European smelt)</td>
<td>Elbe estuary, Germany</td>
</tr>
</tbody>
</table>

Ko, 1995; Pinel *et al.* 1996; Adams *et al.* 1997; Takahashi, 1998). The apparent increase in prevalence and intensity of codworm infection and subsequent condemnation of infected fish can cause substantial economic losses to the fishing industry (Brattey & Davidson, 1996; Rohlwing, Palm & Rosenthal, 1998).

Previous studies employing multilocus enzyme electrophoresis (MEE) have shown that *P. decipiens* (s.l.) comprises at least 6 sibling species (which are morphologically very similar but genetically differentiated), with distinct host preferences and geographical distributions (Paggi *et al.* 1991, 1998; Bullini *et al.* 1994, 1997; George-Nascimento & Llanos, 1995; Mattiucci *et al.* 1998; George-Nascimento & Urrutia, 2000). Three of these sibling species, *P. krabbei*, *P. decipiens* (s.s.) and *P. bulbosa*, have been reported in the North Atlantic, Norwegian and Barents Seas (Paggi *et al.* 1991), while *P. azarasi* has been identified in the Arctic-Boreal and Japanese waters, *P. decipiens* E in Antarctic waters (Bullini *et al.* 1992, 1997; Mattiucci *et al.* 1998), and *P. cattani* in the Southeastern Pacific Ocean (George-Nascimento & Llanos, 1995; George-Nascimento & Urrutia, 2000).

While useful for taxonomic and population genetic studies, the technique of MEE (reviewed by Andrews & Chilton, 1999) has a limitation in that it is reliant on the use of relatively large amounts of fresh or fresh/frozen parasite material for analysis. Polymerase chain reaction (PCR) (Saiki *et al.* 1988) techniques have the advantage over MEE in that their sensitivity allows the analysis of minute amounts of fresh, lyophilized or ethanol-fixed parasite material. Recent PCR-based studies have shown that the first (ITS-1) and/or second (ITS-2) internal transcribed spacers (ITS) of nuclear ribosomal DNA (rDNA) can provide genetic markers for the accurate identification of a number of morphospecies or cryptic species of ascaroid (Jacobs *et al.* 1997; Zhu & Gasser, 1998; Zhu *et al.* 1998a–c, 1999, 2000a, b 2001a, b). Importantly, a study of the *Contracaecum osculatum* complex (see Zhu *et al.* 2000b) indicated that most of its members could be identified and differentiated by single-strand conformation polymorphism (SSCP) analysis of their ITS, suggesting that this rDNA region may also provide specific markers for individual members of the *P. decipiens* complex. In the present study, we employed SSCP to measure the magnitude of sequence variation in the ITS-1 and ITS-2 rDNA within and among 5 recognized members and 2 previously uncharacterized populations of *P. decipiens* (s.l.) from different host species and geographical origins.

**Materials and Methods**

**Parasites**

Adult and larval nematodes corresponding morphologically with *P. decipiens* (s.l.) (see Paggi *et al.* 1991, 2000; Palm *et al.* 1994; Mattiucci *et al.* 1998; Palm, 1999; George-Nascimento & Urrutia, 2000) were used in this study. Their codes, numbers, host species and geographical origins are listed in Table 1. Adult nematodes were collected at necropsy from the stomachs of seal or sea lion definitive hosts, and larval nematodes from the liver, body cavity or musculature of the teleost intermediate or paratenic hosts. Parasites were washed extensively in physiological saline and then frozen at −80 °C or fixed in 50% ethanol. Adult specimens representing the sibling species *P. krabbei*, *P. decipiens* (s.s.), *P. bulbosa*, *P. azarasi* and *P. cattani* were identified previously by MEE analysis (Paggi *et al.* 1991, 1998; George-Nascimento & Llanos, 1995; Bullini *et al.* 1997; Mattiucci *et al.* 1998). Larval specimens representing *P. decipiens* (s.l.) from the teleost hosts *Chaenocephalus aceratus* and *Osmerus eperlanus*, not previously characterized by MEE and thus designated as distinct operational taxonomic units or populations (PdCa and PdOe, respectively), were also included in the study (Table 1).
DNA isolation and enzymatic amplification

Genomic DNA was isolated from individual nematodes by sodium dodecyl-sulphate/proteinase K treatment (Gasser et al. 1993), column-purified (Wizard™ DNA Clean-Up, Promega) and eluted into 40 µl of H2O. Seal DNA was also isolated from liver tissue using the same method. PCR was used to amplify the ITS-1 (plus ~ 80 bp of flanking sequence) with primers NC5 (forward; 5'-GTAGGT-GAACCTTGCGGAAGGATCATT-3') and NC13R (reverse; 5'-GCTGCGTTTCTTTCTCGAT-3'), and the ITS-2 (plus ~ 120 bp of flanking sequence) with primers XZ1 (forward; 5'-ATTGCGCCTC-GGGTTTCTTTTCCTCCGCT-3') and NC2 (reverse; 5'-TTAGTTTCTTTTCTCCGCT-3') (Zhu et al. 2000b). PCR reactions (50 µl) were performed in 10 µl Tris–HCl, pH 8.4, 50 mM KCl, 3 mM MgCl2, 250 µM each of dNTP, 50 pmol of each primer end-labelled with [γ32P]ATP (NEN, DuPont) and 2 U Taq polymerase (Promega) in a thermocycler (Perkin Elmer Cetus) under the following conditions: 94 °C for 5 min (initial denaturation), followed by 30 cycles of 94 °C, 30 sec (denaturation), 55 °C, 30 sec (annealing), 72 °C, 30 sec (extension) and a final extension of 72 °C for 5 min. One microlitre (5–10 ng) of genomic DNA was added to each PCR reaction. Samples with seal DNA or without genomic DNA were included in each amplification run as ‘negative’ controls. An aliquot (4 µl) of each PCR product was examined by agarose gel electrophoresis (Sambrook, Fritsch & Maniatis, 1989).

Single-strand conformation polymorphism (SSCP) analysis

SSCP analysis was carried out as described previously (Zhu & Gasser, 1998). In brief, 10 µl of each PCR product were mixed with an equal volume of loading buffer (10 mM NaOH), 95% formamide, 0.05% bromophenol blue and 0.05% xylene cyanole). After denaturation at 94 °C for 5 min and snap-cooling on a freeze-block (−20 °C), 3 µl of individual samples were loaded into the wells of a 0.4 mm thick, 0.6 × MDE (= mutation detection enhancement, BMA) gel and subjected to electrophoresis in a conventional sequencing rig (Base Runner™, IBI). The conditions for electrophoresis (7 W for 12 h at 20 °C) were standardized for optimal resolution of bands (Zhu & Gasser, 1998). Gels were subsequently dried on to blotting paper and exposed to X-ray film (RP1, Agfa).

DNA sequencing and analyses

Representative amplicons were purified over spin columns (Wizard™ PCR-Preps, Promega) and subjected to manual sequencing using the fmol™ cycle-sequencing kit (Promega), employing the same primers (individually) as used for primary amplification. The 5′ and 3′ ends of the ITS sequences were determined by comparison with those of a range of ascaridoid nematodes, including those within the C. osculatum complex (Jacobs et al. 1997; Zhu et al. 1998a, c, 1999, 2000a, b, 2001a, b), and sequences were aligned manually. Pairwise comparisons were made of the level of sequence difference (D) using the formula D = 1−(M/L) (Chilton, Gasser & Beveridge, 1995), where M is the number of alignment positions at which the 2 sequences have a base in common, and L is the total number of alignment positions over which the 2 sequences are compared. A phenogram was constructed using the unweighted pair group method using arithmetic averages (UPGMA; Sneath & Sokal, 1973).

RESULTS

The ITS-1 and ITS-2 rDNA (including flanking sequences) were amplified individually from genomic DNA prepared from 106 individual worms (Table 1) representing 5 sibling species (P. krabbi, P. decipiens (s.s.), P. bulbosa, P. asarasi and P. cattani) and 2 previously uncharacterized populations of P. decipiens (s.l.). Both the ITS-1 and ITS-2 amplicons were ~ 430 bp in length, and no size variation was detected on agarose gels among any of the samples for both rDNA regions. In no case were amplification products detected in the host or no-DNA control samples (not shown).

The analysis of sequence variation by SSCP has significant technical advantages over other methods, particularly when relatively large numbers of amplicons require analysis in a sequence-dependent manner (Gasser & Zhu, 1999). Hence, this technique was used to screen both the ITS-1 and ITS-2 amplicons for sequence variation within and among the 106 individual nematodes. SSCP analysis of the ITS-1 amplicons resolved 2–3 single-stranded bands per individual, whereas 4–5 bands were displayed for the ITS-2. Except for slight microheterogeneity (Fig. 1, arrowed), no variation in single-stranded ITS-1 and ITS-2 profiles was detected within each of the sibling species or populations, whereas significant differences in SSCP profiles were detected among most of them. For example, P. cattani, P. krabbi, P. decipiens (s.s.), P. bulbosa and P. asarasi each had unique ITS-2 SSCP profiles, which allowed their identification and differentiation from one another (Fig. 1). All individuals of the P. decipiens (s.l.) population PdOe had the same ITS-2 SSCP profile as those representing P. decipiens (s.s.), whereas samples representing the population PdCa all had an ITS-2 SSCP profile which was distinct from all species examined and from population PdOe (Fig. 1).

Of the 212 amplicons scanned by SSCP, the 20 (10 for each rDNA spacer) which represented profiles...
within a species. Microheterogeneity in the SSCP profiles among samples (PdCa). The arrow indicates an example of Osmerus eperlanus (PdOe) or bulbosa (Pb), cattani P the same ITS sequences, those representing those sequences between multiple samples sequenced, length differences varied from 0 to 14 bp in the ITS-1 and 0 to 10 bp in the ITS-2 between samples representing each of the 7 species or populations (Table 2). While no variation in sequence length was detected within any species or population for which multiple samples were sequenced, length differences varied from 0 to 14 bp in the ITS-1 and 0 to 10 bp in the ITS-2 between samples representing each of the 7 species or populations (Table 2).

In accordance with the SSCP results, no substitutional changes were detected in either the ITS-1 or the ITS-2 sequences between multiple samples sequenced for each P. cattani, P. bulbusa and population PdOe, although 2 polymorphic positions were detected in the ITS-1 of P. bulbusa (alignment positions 211 and 233, respectively; Fig. 2) and one in the ITS-2 of both P. cattani and PdOe (alignment positions 62 and 157, respectively; Fig. 2), which was consistent with the microheterogeneity detected in SSCP profiles (Fig. 1). While samples representing P. decipiens (s.s.) and population PbOe had the same ITS sequences, those representing P. cattani, P. krabbei, P. bulbusa, P. azarasi and PdCa each had their unique ITS sequence, allowing their unequivocal molecular identification (Table 3). The ITS-2 representing P. azarasi differed from that of both P. decipiens (s.s.) and PdOe by a single nucleotide, reinforcing the high mutation detection rate of the SSCP method used (Zhu & Gasser, 1998). The G + C content of the sequences varied from 46.4 to 47.9% for ITS-1 and from 43.5 to 45.6% for ITS-2.

The alignment of the ITS-1 and ITS-2 sequences representing the 5 Pseudoterranova species and 2 populations of P. decipiens (s.l.) is shown in Fig. 2. Comparison among all of the sequences (Fig. 2) revealed a total of 65 nucleotide differences (i.e. 27 in the ITS-1 and 38 in the ITS-2, respectively). Of these, 35 (53.8%) were single-base substitutions, representing 25 (38.5%) transitional changes between either purines (A→G; n = 13) or pyrimidines (C→T; n = 12), and 9 (13.8%) transversions (i.e. substitutions between a purine and a pyrimidine). One position (1.5%) represented a multiple-substitution event. The remaining 30 differences (46.2%) represented deletions or insertions.

Pairwise comparisons of the 7 ITS sequences revealed nucleotide differences ranging from 0 to 6.8% (Table 3), which were within the range (0–23%) among members of the C. osculatum complex (Zhu et al. 2000b). With the exception of P. decipiens (s.s.) and population PdOe which had identical ITS sequences, each of the 5 currently recognized sibling species of P. decipiens (s.l.) (based on MEE analysis) could be identified by the presence of at least 1 unequivocal nucleotide difference (Fig. 2). For instance, the G at alignment position 50 in the ITS-2 was unique to P. azarasi, whereas the T’s at both alignment positions 84 and 86 in the ITS-2 allowed the differentiation of P. krabbei from the other taxa (Fig. 2).

The phenogram (Fig. 3) constructed based on the pairwise sequence difference data (Table 3) showed that P. azarasi was genetically more similar to both P. decipiens (s.s.) and population PdOe than it was to any other member of Pseudoterranova, whereas population PdCa was genetically most distinct with respect to all other members of Pseudoterranova examined herein.

**Discussion**

Using MEE analysis, 6 genetically distinct and reproductively isolated species have been detected previously within the morphospecies P. decipiens from the Arctic-Boreal, Boreal and Antarctic waters, and the Southeastern Pacific Ocean (Paggi et al. 1991, 1998; George-Nascimento & Llanos, 1995; Bullini et al. 1997; Mattiucci et al. 1998). A recent study of P. decipiens (s.l.) samples from Atlantic Canada, by restriction fragment length polymorphism analysis of genomic DNA followed by

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**Fig. 1.** Representative SSCP gel displaying sequence variation in the ITS-1 (top) or ITS-2 (bottom) within Pseudoterranova cattani (samples Pc4–Pc8), and among P. cattani, P. krabbei (Pk), P. decipiens (s.s.) (Pd), P. bulbosa (Pb), P. azarasi (Pa), and P. decipiens (s.l.) from Osmerus eperlanus (PdOe) or Chaenocephalus aceratus (PdCa). The arrow indicates an example of microheterogeneity in the SSCP profiles among samples within a species.
Table 2. Length (in bp) and GenBank™ accession numbers of the ITS-1 and ITS-2 sequences representing *Pseudoterranova krabbei* (Pk), *P. decipiens* (s.s.) (Pd), *P. bulbosa* (Pb), *P. azarasi* (Pa), *P. cattani* (Pc) and *P. decipiens* (s.l.) from *Chaenocephalus aceratus* (PdCa) or *Osmerus eperlanus* (PdOe)

(Samples sequenced were selected based on SSCP analysis.)

<table>
<thead>
<tr>
<th>Species/population</th>
<th>Samples sequenced</th>
<th>ITS-1</th>
<th>Accession no.</th>
<th>ITS-2</th>
<th>Accession no.</th>
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<td>296</td>
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<tr>
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</table>

Southern blotting using an ‘heterologous’, cloned rDNA repeat unit as a probe, revealed the presence of *P. decipiens* (s.s.) and *P. bulbosa* in Canadian Atlantic marine fishes and seals (Brattey & Davidson, 1996). This finding was in agreement with results of a previous MEE study (Paggi et al. 1991) which reported the existence of *P. decipiens* (s.s.) and *P. bulbosa* in Atlantic Canada. In the present study, 106 individual nematodes representing 5 currently recognized sibling species within the *P. decipiens* complex (*P. cattani*, *P. krabbei*, *P. decipiens* (s.s.), *P. bulbosa* and *P. azarasi*) and 2 *P. decipiens* (s.l.) populations (PdCa and PdOe) of unknown taxonomic status were characterized by SSCP analysis of their ITS-1 and ITS-2. Each of the 5 sibling species had a unique SSCP profile and ITS sequence, allowing their unequivocal identification and differentiation from one another, which was consistent with previous MEE studies (Paggi et al. 1991, 1998; George-Nascimento & Llanos, 1995; Bullini et al. 1997; Mattiucci et al. 1998). Individuals representing population PdOe from the host *O. eperlanus* had the same SSCP profile and ITS sequence as *P. decipiens* (s.s.), suggesting that this population represented *P. decipiens* (s.s.). This proposal is supported by the geographical origin of PdOe, but independent genetic analysis (e.g. using MEE analysis) is required to verify this. Based on SSCP profile and ITS sequence, population PdCa from the host *C. aceratus* is likely to represent a distinct species. Geographical and host origins as well as its genetic distinctiveness with respect to all other members of the *P. decipiens* complex would support the proposal that it represents *P. decipiens* E. A direct genetic comparison of individuals representing PdCa with those of *P. decipiens* E using the present molecular approaches is warranted when samples representing the latter become available.

*P. decipiens* E is a parasitic nematode occurring in Antarctic waters (Palm, 1999). Its main definitive host is the widely-distributed Weddell seal, *Leptonychotes weddellii* (see Bullini et al. 1992, 1997), and its 3rd-stage larvae occur in a number of fish species in the Weddell Sea and South Shetland waters (Palm et al. 1994, 1998; Palm, 1999). A previous study (Palm et al. 1998) has indicated that most parasite species of the Antarctic fish, *Notothenia coriiceps*, appear to be broadly distributed within Antarctic waters, which would correspond to a low host specificity in higher latitudes. However, to date, it remains unclear whether all *P. decipiens* (s.l.) in the Antarctic belong to a single sibling species (i.e. *P. decipiens* E). Employing the SSCP-sequencing approach, this question could be readily addressed by genetic comparison of *P. decipiens* (s.l.) specimens from a range of intermediate, paratenic and definitive host species and localities within the Antarctic waters.

*P. cattani* is a recently described anisakid of the South American sea lion, *Otaria byronia*, which is distinct with respect to other known *Pseudoterranova* species based on allozymic and morphological characters (George-Nascimento & Llanos, 1995; George-Nascimento & Urrutia, 2000). The adult and larva of this species from the south-eastern Pacific Ocean were recognized previously as *Phoceanema decipiens* (see George-Nascimento & Urrutia, 2000). The results of the present genetic study support the specific taxonomic status of *P. cattani*, given its distinct SSCP profile and ITS sequence with respect to all other members of *Pseudoterranova* examined. Moreover, in addition to *P. decipiens* (s.l.) and *P. cattani*, *P. kogiae* and *P. ceticola* have also been described in the literature (Anderson, 2000). Therefore, the SSCP-sequencing approach employed in this study should be useful for a direct genetic comparison of these *Pseudoterranova* species for which no molecular data are currently available.

Although morphometric keys are available for the identification of adult males of *P. krabbei*, *P. decipiens* (s.s.), *P. bulbosa* and *P. azarasi* (see Di Deco et al. 1994; Mattiucci et al. 1998; Paggi et al. 2000), no
Fig. 2. Alignment of the ITS-1 and ITS-2 sequences representing *Pseudoterranova krabbei* (Pk), *P. decipiens* (s.s.) (Pd), *P. bulbosa* (Pb), *P. azarasi* (Pa) and *P. decipiens* (s.l.) from *Osmerus eperlanus* (PdOe) or *Chaenocephalus aceratus* (PdCa) as well as those of *P. cattani* (Pc). Refer to Tables 1 and 2 for samples sequenced. Sequences have been deposited in the EMBL, GenBank™ and DDBJ databases. Nucleotide differences among the aligned sequences are indicated by asterisks. Polymorphic sites in the sequences are designated with IUPAC codes.
such keys are available for the specific identification of larval stages. A number of crustacean and fish species have been identified as the intermediate and intermediate/paratenic hosts of *P. decipiens* (s.l.), respectively (George-Nascimento & Llanos, 1995; Palm, 1999), but relatively little is currently known about the distributions and role(s) of each host species in the transmission of each sibling species (Paggi et al. 1991). Therefore, the establishment of molecular-genetic tools for the precise identification of sibling species within *P. decipiens* (s.l.) is of significance for further studies of their life-cycles, host preference, transmission patterns and population genetic structures.

The molecular tools should also have significant implications for the diagnosis of pseudoterranoviasis in humans. To date, clinical diagnosis of human gastric and intestinal pseudoterranoviasis has relied mainly on the detection of nematodes upon endoscopy, followed by morphological identification of the larvae recovered (Oshima, 1987; Ko, 1995; Pinel et al. 1996; Adams et al. 1997). However, given the limitations associated with the morphological identification of larval stages or portions thereof, SSCP analysis of ITS-2 can achieve their specific identification from minute amounts of parasite material extracted during endoscopic examination. Indeed, a recent study (D’Amelio et al. 1999) has demonstrated the usefulness of PCR-linked restriction fragment length polymorphism analysis for the specific identification of *Anisakis* larva endoscopically recovered from the stomach of a woman in Southern Italy.

There is a distinct difference between the clinical syndrome of human pseudoterranoviasis in Japan and that in the USA (Oshima, 1987). In Japan, the disease is quite severe, whereas in the USA, it is milder and without clear symptoms attributable to gastro-intestinal invasion (Oshima, 1987). While the reason(s) for this variation in pathogenic effect is currently unclear, Desowitz (1986) suggested that it may be attributable to different ‘strains’ of *P. decipiens* (s.l.) with differing virulence. It would be interesting to genetically compare *P. decipiens* larvae from humans from Japan with those from the USA, employing the SSCP approach. Also, genetic comparison of *P. decipiens* (s.l.) larvae from humans with adult or larval stages from a range of intermediate, paratenic and definitive hosts may establish which species is/are the causative agent(s) for human pseudoterranoviasis and thus pose a public health threat.

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