

1 **Molecular identification of three co-occurring species of *Acartia* in the Thau**

2 **Lagoon, France.**

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4 P. K. Lindeque¹, S. Boyer² and D. Bonnet².

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6 1. Plymouth Marine Laboratory, Prospect Place, West Hoe, Plymouth, PL1 3DH, UK.

7 pkw@pml.ac.uk, Tel: +44 (0)1752 633100, Fax: +44 (0)1752 633101.

8 2. Laboratoire EcoSym (Ecologie des Systèmes Marins Côtiers), UMR5119, Université

9 Montpellier 2, CC093, Place Eugène Bataillon 34095 Montpellier Cedex 05, France.

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11 **Key Words:** *Acartia*, Thau lagoon, resting eggs, PCR-RFLP

12

13 **Abstract**

14 *Acartia* species, often known to co-occur, can exhibit complex life cycles, such as the
15 production of resting eggs. Studying and understanding their population dynamics is hindered
16 by the inability to identify eggs and early developmental stages using morphological
17 techniques. A simple molecular technique to distinguish between the three species of the
18 *Acartia* genus (*A. clausi*, *A. discaudata* and *Paracartia grani*) that co-occur in the Thau
19 lagoon (40°03 N; 03°41 E) in Southern France has been developed. Direct amplification of a
20 partial region of the mitochondrial cytochrome oxidase I (mtCOI) gene by polymerase chain
21 reaction (PCR) and subsequent restriction fragment length polymorphism (RFLP) results in a
22 unique restriction profile for each species. The technique is capable of determining the
23 identity of individual copepods and eggs, including resting eggs retrieved from sediment
24 samples, illustrating its application in facilitating population dynamic studies of this
25 ubiquitous and important member of the zooplankton community.

26

27 **Introduction**

28 The copepods from the Acartiidae family generally occur in estuaries, harbours and
29 other semi enclosed marine habitats (Alcaraz 1983), in which food is rarely limited (Calbet *et*
30 *al.* 1999). *Acartia* species are commonly known to co-occur in the same environment and in
31 Mediterranean bays, lagoons and harbours, the number of species reported to co-exist varies
32 from two in the ports of Naples (Yamazi 1964) and Marseille (Arfi *et al.* 1981) to six in the
33 small port of Jounieh (Lakkis 1994). In the Thau lagoon, the largest of a string of lagoons that
34 stretch along the Languedoc-Roussillon French coast from the Rhône River to the foothills of
35 the Pyrenees, the genus *Acartia* represents the most abundant taxon in the plankton
36 community (Fatemi 1938; Lam Haoi 1985). To our knowledge, 5 congeneric species have
37 been identified until now in the Thau lagoon: *A. clausi*, *A. bifilosa*, *A. discaudata*, *A.*
38 *margalefi* and *A. latisetosa* (Mathias and Euzet 1962; Lam Haoi 1985). However, unpublished
39 observations (Euzet pers comm; Boyer *et al.* in press) have highlighted that since 1998,
40 *Paracartia grani* has appeared in this lagoon. A two year monitoring program in the Thau
41 lagoon (40°03' N; 03°41' E), has shown that *A. clausi* predominates in winter, while *A.*
42 *discaudata* peaks in spring and *P. grani* in summer (Boyer *et al.* submitted).

43 *Acartia* species are known to have complex life cycles with *A. clausi* and *P. grani*
44 producing resting eggs (i.e. Kasahara *et al.* 1974; Marcus 1990; Guerrero and Rodriguez
45 1998). To study the population dynamics of these species, abundance, stage development and
46 egg production of *Acartia* spp. have been recorded every two weeks during a 2 year survey in
47 the Thau lagoon (Boyer *et al.* submitted). Sediment cores were also collected at two dates
48 (May 2010 and April 2011) to investigate the potential pool of recruits coming from the
49 sediment. In 2010, copepod eggs extracted from the sediment were incubated and a very high
50 hatching success was observed. To establish the species identity of the eggs, they were
51 maintained individually in an attempt to allow them to reach a developmental stage suitable

52 for morphologic identification. Unfortunately, this was never successful as many nauplii and
53 juveniles died before reaching copepodite stages.

54 In situations where morphological characters do not provide sufficient variation to
55 identify to species level, as is the case with the resting eggs and early developmental stages of
56 *Acartia* spp., then taxonomic discrimination can be achieved using genetic characters. Such
57 techniques have been developed for copepods of the genus *Pseudocalanus* (Bucklin *et al.*
58 1998; Grabbert *et al.* 2010), *Calanus* (Lindeque *et al.* 1999; Bucklin *et al.* 1999; Hill *et al.*
59 2001) and *Clausocalanus* (Blanco-Bercial and Alvarez-Marques 2007). However, no such
60 technique has been developed for *Acartia* species. Therefore, during this study we developed
61 a molecular tool to identify individuals, from eggs to adults of the three congeneric species, *A.*
62 *clausi*, *A. discaudata*, and *P. grani*, to facilitate population dynamics studies. These three
63 species were chosen as they are by far the dominant co-occurring species (Lam Haoi 1985;
64 Boyer *et al.* submitted) and because two of them (*A. clausi* and *P. grani*) are known to
65 produce resting eggs, an important life strategy for surviving adverse periods (Kasahara *et al.*
66 1974; Marcus 1990; Guerrero and Rodríguez 1998). The technique therefore, has to be
67 sufficiently robust to work on individual eggs, including resting eggs taken from sediment
68 cores.

69 The gene of choice for our study was the common metazoan DNA barcoding gene,
70 mitochondrial cytochrome oxidase subunit I (mtCOI). This gene has previously been used for
71 the identification and assessment of species diversity and distribution of marine zooplankton
72 (Bucklin *et al.* 1999; Hill *et al.* 2001; Webb *et al.* 2006; Blanco-Bercial and Alvarez-Marques
73 2007; Grabbert *et al.* 2010; Bucklin *et al.* 2010). However, at the beginning of this study, out
74 of the three species *A. clausi*, *A. discaudata* and *P. grani*, only sequence data for the mtCOI
75 gene for *A. clausi* existed on the EMBL DNA sequence database. Therefore, a sequencing
76 effort of adults of all three species collected from the Thau lagoon was necessary. The use of

77 a mitochondrial gene was preferential as such a gene has a high copy number and is relatively
78 accessible to PCR primers allowing PCR amplification to be achieved direct from an
79 individual egg or copepod without prior purification of the DNA. The mitochondrial 16S
80 rRNA gene, which has previously been used for copepod species identification (Bucklin *et al.*
81 1998; Lindeque *et al.* 1999) showed insufficient inter-specific variation between the *Acartia*
82 species (Lindeque unpubl data).

83 A simple and unambiguous method to distinguish between the co-occurring *Acartia*
84 spp. in the Thau lagoon, including early developmental stages and resting eggs, will allow a
85 better description and understanding of the species population dynamics.

86

87

88 **Material and Methods**

89

90 **Sample collection and preservation**

91 Zooplankton samples were collected at a fixed station (40°03 N; 03°41 E) close to
92 Sète channel in the Thau lagoon (south of France). Horizontal hauls were performed in the
93 inner surface water (maximum 1 m depth) with a WP2 plankton net (200- μ m mesh size). The
94 cod-end contents were placed in insulated containers and brought back to the laboratory
95 within 1 h after collection. In the laboratory, healthy females of *A. clausi*, *A. discaudata* and
96 *P. grani* were picked out. Physiological state of the female (ovigerous versus non ovigerous)
97 was determined with a binocular microscope using morphological characteristics (Rose
98 1933). For each species, ovigerous females were incubated for 24 h in a 1 L beaker filled with
99 0.45 μ m filtered sea water, after which the females and the eggs produced were fixed in 95%
100 alcohol.

101 Two sediment cores were sampled on the 27th of May 2010 and the 27th April 2011 at
102 the same monitoring station by a diver. The cores were sliced and each slice preserved at 4°C

103 in darkness before analyses. Resting eggs were extracted from the sediment samples
104 according to the method of Onbé (1978). In 2010, copepod eggs extracted from the sediment
105 were used for hatching experiments. In 2011 no hatching success experiment was run and the
106 eggs collected were directly counted using a binocular microscope and fixed in 95% alcohol.

107

108 **DNA amplification and sequencing**

109 Adult individuals of the three *Acartia* species, *A. clausi*, *A. discaudata* and *P. grani*
110 were prepared for PCR amplification by rehydrating in 22.75 µL DNA grade water in a PCR
111 tube or 96-well plate at room temperature for between 4-6 h. Following this incubation 10 µL
112 of 5x Flexi GoTAQ DNA polymerase buffer (Promega UK) was added to each sample and
113 the sample homogenised using a hyperdermic needle (19G) inserted into a pellet pestle hand
114 held homogeniser (Anachem Ltd) and incubated overnight at 4°C. The remaining PCR
115 reaction components were then added: 5 µL 2 mM dNTPS (Promega UK Ltd), 10 µM of
116 primers LCO-1490 and HCO-2198 (LCO-1490 5'-GGT CAA CAA ATC ATA AAG ATA
117 TTG G-3' and HCO-2198 5'-TAA ACT TCA GGG TGA CCA AAA AAT CA-3', Folmer *et*
118 *al.* 1994), 2 µL 25 mM MgCl₂ and 1.25 U GoTAQ DNA polymerase (Promega UK Ltd.). A
119 positive control of *A. clausi* and *P. grani* extracted genomic DNA (100-200 ng) and a NTC
120 (no template control) were included. Amplifications were carried out in a G Storm or VWR
121 thermocycler. The cycling parameters included an initial denaturation step at 94.0°C (4 min)
122 followed by 40 cycles of 94.0°C (1 min), 50.0°C (1.5 min), 72.0°C (1.5 min). A final
123 extension phase at 72°C (5 min) was followed by storage at 10.0°C. 10 µL aliquots of the
124 amplification reaction were analysed by gel electrophoresis (1%) to check amplification
125 efficiency. 20 µL of successful amplifications were sent away for bidirectional sequencing by
126 LGC Genomics GmbH, Germany. These sequences (n=71) were then used to perform a
127 multiple sequence alignment using ClustalW analysis online at <http://clustalW.genome.jp/> and

128 subsequently a consensus sequence for each species was constructed. The basic local
129 alignment search tool (BLAST) was used to search the EMBL DNA database for sequence
130 similarities. A FastA file of the sequences was converted to a sequence identity matrix using
131 the BioEdit program (Hall 1999). To determine similarity and clustering of the sequences a
132 nonmetric multidimensional scaling plot was constructed using PRIMER (Clarke 1993;
133 Clarke and Warwick 2001).

134

135 **Restriction mapping**

136 The partial consensus sequences of mtCOI DNA for *A. clausi*, *A. discaudata* and *P.*
137 *grani* were restriction mapped using the online NEBcutter V2.0 from New England BioLabs
138 inc. (Vincze *et al.* 2003). From these restriction maps, suitable enzymes were selected to
139 differentially digest each of the three species. To reduce both time and cost, the ability of the
140 restriction enzyme to perform in the same buffer conditions as the amplification reaction were
141 taken in to consideration.

142

143 **Restriction digests**

144 Restriction digests were performed on a 15 µL aliquot of PCR amplification product
145 by the addition of 0.2 µL of BSA (10 µg mL⁻¹) and 5 U of each restriction enzyme (*DdeI* and
146 *PstI* Promega UK Ltd). Incubations were performed at 37°C for 3-4 h. The digestion products
147 were separated by gel electrophoresis on a 2% agarose gel and visualised on a UV
148 transilluminator.

149

150 **Molecular identification of *Acartia* eggs**

151 *Ethanol preserved*

152 Following the development of the RFLP technique based on adults the method was
153 then trialled on *Acartia* eggs collected from egg production experiments and then applied to
154 the copepod eggs collected directly from the sediment. 24 eggs from core 1 (2-2.5 cm depth)
155 collected in April 2011 were used to assess the ability of this technique to identify eggs from
156 the sediment following the method described above.

157 During these studies, it became apparent that *Acartia* eggs, either from egg production
158 experiments or from sediment, are poorly preserved in ethanol for any length of time.
159 Therefore, the method was adapted to ensure successful amplification of DNA following
160 preservation at -20°C. This method is described below.

161

162 *Preserved at -20°C*

163 Eggs were washed into a Petri dish with filtered seawater (fsw). Under a microscope a
164 single egg was aspirated with 0.5 uL fsw and transferred to a new Petri dish containing 5 mL
165 of ultrapure water (MilliQ) to “wash” the egg. The egg was then quickly (to prevent
166 rupturing) aspirated with 23 uL of the ultrapure water and transferred into a well of a 96-well
167 plate. After 4-6 hours, the plate was sealed and frozen at -20°C overnight or longer. The
168 samples were then defrosted and the remaining PCR reaction components added to each well:
169 10 µL of 5x Flexi GoTAQ DNA polymerase buffer (Promega UK), 5 µL 2 mM dNTPS
170 (Promega UK Ltd.), 10 µM of primers LCO-1490 and HCO-2198 (LCO-1490 5'-GGT CAA
171 CAA ATC ATA AAG ATA TTG G-3' and HCO-2198 5'-TAA ACT TCA GGG TGA CCA
172 AAA AAT CA-3', Folmer *et al* 1994), 2 µL 25 mM MgCl₂ and 1.25 U GoTAQ DNA
173 polymerase (Promega UK Ltd). Amplifications and restriction digests were then carried out as
174 described above.

175

176 **Results**

177 A partial region (709 bp) of the mtCOI gene was successfully amplified directly from
178 ethanol preserved individuals of the three *Acartia* spp. *A. clausi*, *A. discaudata* and *P. grani*
179 (Figure 1) collected from Thau lagoon, France. In total 71 sequences were generated for this
180 region of the mtCOI gene; *A. clausi* (n=22; 5 haplotypes; Accession numbers HE 863718-HE
181 863722), *A. discaudata* (n=21; 9 haplotypes; Accession numbers HE 863723-HE 863731) and
182 *P. grani* (n=28; 10 haplotypes; Accession numbers HE 863732-HE 863741). A consensus
183 sequence for each species was constructed (Figure 2). When compared to sequences on the
184 EMBL DNA database, *A. clausi* sequences showed between 96%-100% identity over 85%
185 coverage to existing sequences of *A. clausi* (Accession numbers: HE6647790.1-HE6479.1).
186 No sequence homology greater than 85% was seen for sequences of either *P. grani* or *A.*
187 *discaudata*. Over the amplified partial region of the mtCOI gene, the intraspecific variation
188 ranged from ~2.0% for *A. discaudata*, to ~3.7% for *P. grani* and ~4.0% for *A. clausi*. The
189 interspecific variation between these species was 29% (Figure 3).

190 The sequenced mtCOI regions for each of the three *Acartia* species were mapped and
191 analysed to ascertain suitable restriction enzyme sites such that each species, when digested,
192 would produce a unique restriction profile (Figure 4). The restriction enzymes *DdeI* and *PstI*
193 were chosen as they both produced a unique restriction profile for each species and because
194 the enzymes exhibit optimum activity at the same temperature (37°C) such that restriction
195 digests can be performed in the same buffer conditions as the amplification reaction (Tritle
196 2006). Successful amplifications from individual *Acartia* adults and subsequent RFLP
197 analysis resulted in a characteristic restriction profile or fingerprint for each species (Figure
198 5), thus allowing identification of *Acartia* species to species level.

199 Amplification of the mtCOI partial gene fragment was also successful from eggs
200 collected from *Acartia* egg production experiments preserved in 95% ethanol and from eggs
201 collected from sediment samples preserved in 95% ethanol. Of the 24 eggs processed from

202 sediment samples (Core 1, 2-2.5cm depth, April 2011) 15 amplified successfully. Of these
203 successful amplicons all 15 digested to produce a restriction profile for *A. clausi* (n=5) or *P.*
204 *grani* (n=10). In order to confirm the accuracy of species identification by RFLP on sediment
205 eggs the amplified products were sequenced. In all cases the sequences confirmed the species
206 identity of the individual eggs to be the same as those determined by RFLP. During this study
207 it became apparent that the eggs were not stable stored in ethanol for any length of time. After
208 six months it was no longer possible to successfully amplify the DNA from individual eggs.
209 As such the technique was adapted to provide a protocol for future preservation of individual
210 eggs that is robust and reliable. The technique described was trialled on *A. clausi* eggs laid
211 during egg production experiments and routinely allowed amplification of individual eggs
212 with a success rate of over 80%.

213

214 **Discussion**

215 A simple molecular method involving the direct amplification of a partial region of the
216 gene encoding the mitochondrial protein cytochrome oxidase subunit I (mtCOI) and
217 subsequent restriction fragment length polymorphism (RFLP) has been developed to identify
218 individuals of *A. clausi*, *A. discaudata* and *P. grani*. In order to develop this technique a total
219 of 71 sequences were generated for this region of the mtCOI gene for all three congeneric
220 species collected from the Thau lagoon, France. Over the amplified partial region of the
221 mtCOI gene the intraspecific variation ranged from ~2.0% for *A. discaudata*, to ~3.7% for *P.*
222 *grani* and ~4.0% for *A. clausi*, an accepted level of intra-specific difference for this gene
223 region. Hill *et al.* (2001) found intraspecific variation of between 0.5% and 2.8% for this
224 region of gene in 10 *Calanus* species. For the same region of gene intra-specific variation was
225 less than 2% for *Pseudocalanus* species (Grabbert *et al.* 2010) and between 0.5% and 4.2%
226 for 13 species of *Clausocalanus* (Bucklin *et al.* 2010). When compared against sequences on

227 the EMBL DNA database *A. clausi* showed between 96% and 100% homology with existing
228 *A. clausi* sequences, further helping to confirm the identity of our species. No homology for
229 sequences of either *P. grani* or *A. discaudata* over 85% similarity was seen. In the case of *A.*
230 *discaudata* this is understandable as there are no existing COI sequences for this species on
231 the database. There is however, a recent submission of a putative *P. grani* COI sequence
232 from the Egyptian coast of the Mediterranean Sea (Accession number JQ245071.1) on the
233 database. When aligned with our *P. grani* sequences it showed a variation greater than 30%.
234 This intra specific variation is extremely high for this region of gene, as discussed above the
235 range of intra specific variation is usually between 0% and 4%. It is impossible to explain this
236 high level of variation between the sequences of *P. grani* from the Thau lagoon, France and
237 the *P. grani* sequence from the Egyptian coast of the Mediterranean Sea as the latter is only a
238 single unverified sequence.

239 The inter-specific variation between these congeneric *Acartia* species was 29%. This
240 is also an accepted level of variation for this gene region between congeneric species.
241 Previous studies have shown pair wise sequence differences to be between ~18% for two
242 species of *Pseudocalanus* (Bucklin *et al.* 1998), between 7.4% and 25.7% for 10 congeneric
243 species of *Calanus* (Hill *et al.* 2001) and between 8.6% and 24.9% for 13 species of
244 *Clausocalanus* (Bucklin *et al.* 2010).

245 The developed technique is capable of determining the identity of individual copepods
246 and individual eggs, including resting eggs retrieved from sediment samples. Consideration
247 has been made to produce a reliable, quick and cost effective technique. As such amplification
248 is carried out directly on preserved samples negating the need for prior purification of the
249 DNA. While PCR products can be purified prior to restriction endonuclease digestion, both
250 time and reagents are saved by performing restriction digests in the same buffer conditions as
251 the amplification reaction. Eliminating a purification step after amplification and before

252 restriction digest also eliminates potential pipetting error and decreases assay preparation
253 time.

254 Amplification of adults preserved in 95% ethanol was consistently successful and
255 unproblematic. The success rate of amplification from eggs however, was much lower than
256 for adults. Initially this was thought to be because of the very small size of the eggs
257 ($\sim 74.4 \pm 1.5 \mu\text{m}$ for *A. clausi*; Belmonte 1998) which poses considerable challenges when
258 manipulating them into a PCR tube or 96-well plate. This problem is exasperated when the
259 eggs have been stored in ethanol as they then appear to be ‘static’ and regularly stick to all
260 plastic-ware. Further tests revealed that the eggs ruptured on contact with the ethanol and that
261 the cytoplasm burst through the membrane of the egg. It was possible to successfully amplify
262 DNA from the eggs that had been stored in ethanol if the preservation was under 6 months.
263 However, the eggs appear to be unstable in the ethanol and DNA amplification was
264 impossible after a relatively short period of time (\sim six months). As such a new protocol has
265 been developed whereby eggs, originating from the water column, egg production
266 experiments or sediment, are washed in filtered sea water and then rinsed in ultra-pure MilliQ
267 water. The eggs are then removed with the correct volume of water for the PCR reaction and
268 isolated in a well of a 96-well plate before they rupture. This protocol has many benefits. As
269 the eggs have not been stored in ethanol they are not static, which makes handling much
270 easier. The eggs are isolated in individual wells before rupturing thus preventing any possible
271 cross contamination, the technique significantly reduces the time consuming and difficult
272 manipulation of the eggs compared to storage of multiple eggs in a single vial, and perhaps
273 most importantly, it allows successful amplification of individual eggs.

274 For the first time a technique now exists to identify the 3 congeneric species of co-
275 occurring *Acartia*; *A. clausi*, *A. discaudata* and *P. grani* from the Thau lagoon, France. This
276 technique can identify individuals from eggs to adults including resting eggs found in the

277 sediment. To our knowledge this is the first time that resting eggs in the Thau lagoon have
278 been identified to species level. The developed technique was trialled on a small sample of
279 resting eggs retrieved from the sediment and showed the presence of both *A. clausi* and *P.*
280 *grani* species. These results confer with the literature that both these species are able to
281 produce resting eggs (i.e. Kasahara *et al* 1974; Marcus 1990; Guerrero and Rodríguez 1998).
282 Care must be taken however, as the COI primers used are universal and may therefore
283 amplify COI from other organisms. In the Thau lagoon the resting eggs are easily identifiable
284 as copepod eggs, and although there is a slight chance that other copepods found in the lagoon
285 may produce resting eggs this is firstly unlikely (undocumented for all but *Eurytemora velox*)
286 and secondly these species such as *Eurytemora* and *Centropages* species make up only a very
287 small proportion of the total zooplankton community in the lagoon (<10%) (Lam Haoi 1985).

288 Considering the difficulties of identifying early developmental stages to species using
289 morphological characters and the time constraints of hatching resting eggs and culturing them
290 to maturity in order to allow identification, this new method will allow better population
291 dynamic studies to be performed on these important and dominant members of the
292 zooplankton community in the Thau lagoon, France.

293

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298

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385 **Figure Legends:**

386 **Fig. 1** Amplification of a 709 base pair region of the mtCOI gene from preserved adult
387 *Acartia* M = 100bp DNA ladder, lanes 1-12 = *A. clausi*, lanes 13-24 = *P. grani*, lanes 25-36 =
388 *A. discaudata*. -ve = no DNA template control, +ve = positive control of *A. clausi* and *P.*
389 *grani* extracted genomic DNA (100-200ng)

390

391 **Fig. 2** Consensus sequence of a 709 bp region of the mtCOI gene for the three species *A.*
392 *clausi* (*clausi*), *A. discaudata* (*discaudata*) and *P. grani* (*grani*). * = homology between all
393 three species; **ctnag** = *DdeI* restriction site; **ctgcac** = *PstI* restriction site; primers in italics

394

395 **Fig. 3** Nonmetric multidimensional scaling plot demonstrating similarity and clustering of *A.*
396 *clausi* ▲, *P. grani* ▼ and *A. discaudata* ■. All sequences (n=71) were used to create a
397 sequence identity matrix. Sequences from the same species cluster with >96% similarity
398 ----- *A. clausi* and *A. discaudata* cluster at >80% similarity ----- and all three species
399 cluster at >70% similarity ——

400 .

401 **Fig. 4** Restriction map of a partial region of the mtCOI gene for *A. clausi*, *P. grani* and *A.*
402 *discaudata*. *DdeI* and *PstI* restriction sites are indicated. Resulting fragment lengths are
403 shown in base pairs (bp)

404

405 **Fig. 5** Amplification of a region of mtCOI gene from whole *Acartia* adults and subsequent
406 RFLP analysis with restriction enzymes *DdeI* and *PstI*. M=100bp DNA markers, Lanes 1-3 =
407 restriction profile for *P. grani*; Lanes 4-6 restriction profile for *A. discaudata*; Lanes 7-9 =
408 restriction profile for *A. clausi*; Lane 10 = undigested amplicon

Figure

Fig. 1

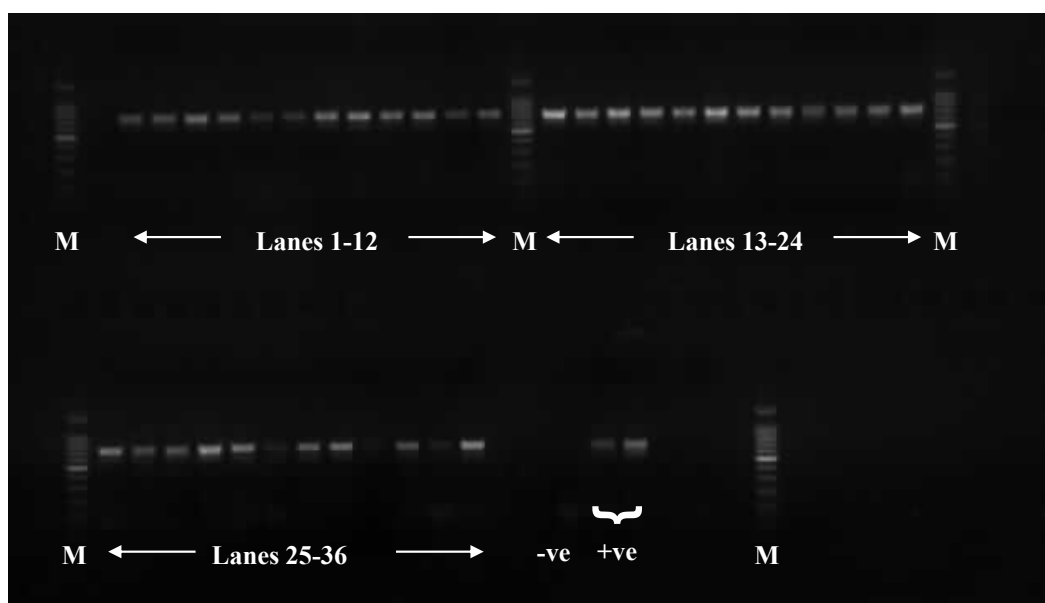


Fig. 1 Amplification of a 709 base pair region of the mtCOI gene from preserved adult *Acartia* M = 100bp DNA ladder, lanes 1-12 = *A. clausi*, lanes 13-24 = *P. grani*, lanes 25-36 = *A. discaudata*. -ve = no DNA template control, +ve = positive control of *A. clausi* and *P. grani* extracted genomic DNA (100-200ng)

Fig. 2

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clausi          GGTCAACAAATCATAAAGATATTGGCACTTTATATTTACTRGCTGGTATRTGGTCGGGGA
discaudata     GGTCAACAAATCATAAAGATATTGGYACTCTYTACCTTTTAGCTGGGATGTGGTCAGGAA
grani          GGTCAACAAATCATAWAGATATTGGGACTTTATATTTATTAGCAGGGSCTTGATCTGGAA
*****
clausi          TAGTRGGCACTGGATTAAGAATRATCATYCGAATRAGCTTGGTCAAGCAGGAAAAATTAA
discaudata     TAGTGGGAACCTGGCTTGAGAATGATCATTCGGATAGAAYTAGCCAAAGCTGGAAAAATTAA
grani          TAGTTGGAACAGGCCTTAGAAATAATTATTCGATTAGARCTGGGTCAAGCTGGAAGATTAA
* * * * *
clausi          TTGGAGATGAYCAAATYTATAACGTAGTTGTAACAGCCCATGCCTTTATTATAATTTTTT
discaudata     TTGGGGATGATCAAATTTATAACGTGGTTCGTTACAGCCACGCGTTTATTATAATTTTCT
grani          TTGGAGATGATCAAATTTACAATGTTGTTGTAACCGCTCATGCYTTTATTATAATTTTTT
****
clausi          TTATAGTTATACCAATTCTAATTGRRGGGTTTGGRAATTGAYTAATYCCCCTAATACTAG
discaudata     TTATAGTAATACCTATTTTAATTGGGGGGTTTGGAAATTGACTAATCCCTCTTATACTWG
grani          TCATAGTYATACCTATTTCTAATTGGAGGATTTGGRAATTGACTAGTGCCTCTTATATTGG
* * * * *
clausi          GAGCTGCAGAYATAGCYTTCCCYCGAATAAATAATAAAGTTTTGRCTCCTACTTCCCTG
discaudata     GGGCTGCTGACATAGCRTTTCCTCCGCATAAATAATAAAGATTTGACTCCTACTCCCCG
grani          GTGCAGCGGATATAGCATTCCCTCGAATAAATAATAAAGATTTGATTCTTAATACCAG
* * * * *
clausi          CTTTAGTYATGCTTTTATCTAGTTCCTTTAGTGGAAAGAGGGGCAGGAACTGGGTGAACAG
discaudata     CCTTGATTATRTTGCTGTCAAGRCTTAGTGGAAAGGGGGGCTGGRACCGGCTGAACAG
grani          CTTTAATTATATTRYTATGTAGTTCATTAGTTGAAAGGGGRGCCGGAACAGGTTGAACAG
* * * * *
clausi          TTTAYCCACCMCTATCTAGAAATATTGCCCATGCAGGRGCTTCAGTTGATTTTCGCAATTT
discaudata     TTTACCCCCCTCTCAAGCAATATTGCCACGCAGGCAGGGCTGTAGATTTTGCATTTT
grani          GTAYCCCCCTTTGTTCGAGGAAYATTGCTCATGCGGGAGCTTCAGTTGATTTTGTCTATTT
* * * * *
clausi          TTTCCCTCCATCTTGCAGGTGCAAGATCTATTTTAGGAGCTGTAACTTTATYCTACTG
discaudata     TTTCACTCCATTTAGCGGGGCCAGTTCATTTCTAGGAGCTGTAAATTTTATTTCTACTA
grani          TTTCTTTACATCTTGGGGGTGAAGTTCATTTTAGGKGTGTTAATTTTATCTCAACTG
****
clausi          TAGGWAACCTTCGATCATTTGGGATAATAGCGGATTTAATACCTTTRTTTAGGTGGGCGG
discaudata     TTGGTAACTTGCATCATTTGGCATGATAGCGGATCTAATACCTTTGTTTAGRTGGGCTG
grani          TTGGAATCTTCGGCATTTTGAATAATTCTTGATCGAATACCATTATTTGCATGAGCAG
* * * * *
clausi          TATTAATTACAGCTGTTTTACTACTYTTATCTTTGCCAGTRTTAGCGGGGGCTATYACTA
discaudata     TGTAATTACAGCGGTTTTGCTTTTATTATCTCTTCCTGTTTTAGCCGGGGCTATTACTA
grani          TATTAATTACAGCAGTTCTATTRCTATTRCTTTACCTGTAYTAGCTGGGGCTATTACTA
* * * * *
clausi          TACTTCTCACTGATCGTAATTTAACTCTTCTTTTTATGACGCCGCGGTGGAGGAGAYC
discaudata     TGCTTTTAAACYGACCGTAATTTGAATTCATCGTTTTACGATGCYGGAGGGGGGGAGAYC
grani          TATTATTRACTGAYCGTAACTTAAATTCATCATCTACGATGCTAGAGGKGGCGCGACC
* * * * *
clausi          CYATCCTTTATCAACATTTATTTTGATTTTTTGGTCACCCTGAAGTTTA
discaudata     CYATCCTGTATCAGCACCTGTTTTGATTTTTTGGTCACCCTGAAGTTTA
grani          CTATCTTRTATCAGCATTTATTTTGATTTTTTGGTCACCCTGAAGTTTA
* * * * *

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Fig. 2 Consensus sequence of a 709 bp region of the mtCOI gene for the three species

A. clausi (*clausi*), *A. discaudata* (*discaudata*) and *P. grani* (*grani*). * = homology

between all three species; **ctnag** = *DdeI* restriction site; **ctgcaac** = *PstI* restriction site;

primers in italics

Fig. 3 Nonmetric multidimensional scaling plot demonstrating similarity and clustering of *A. clausi* ▲
P. grani ▼ and *A. discaudata* ■. All sequences (n=71) were used to create a sequence identity matrix.
Sequences from the same species cluster with >96% similarity ----- *A. clausi* and *A. discaudata*
cluster at >80% similarity ----- and all three species cluster at >70% similarity ———

Fig. 3

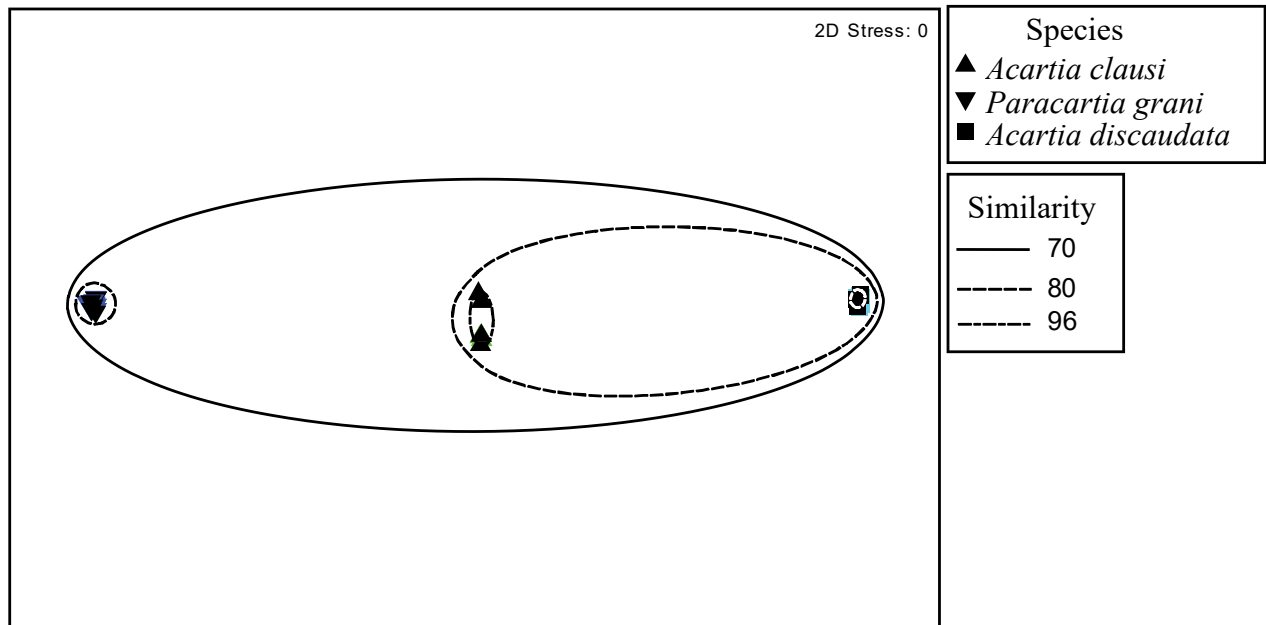


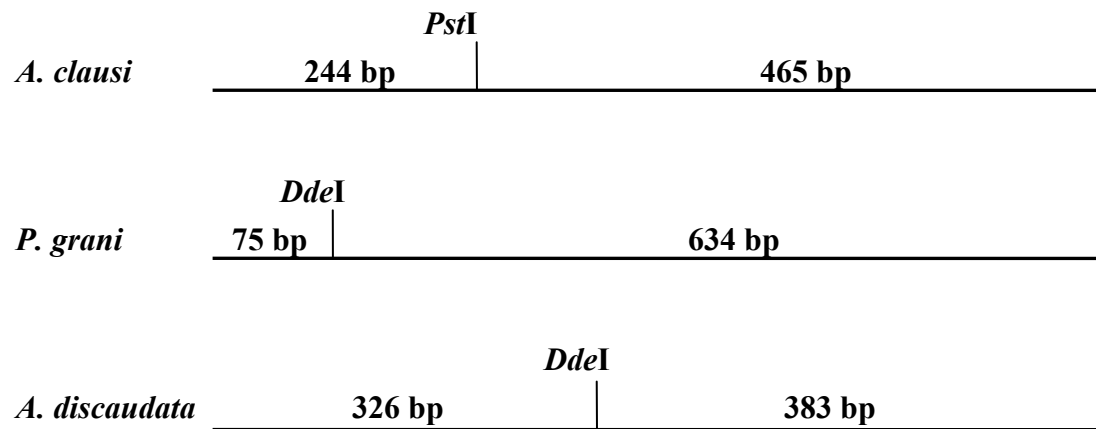
Fig 4

Fig. 4 Restriction map of a partial region of the mtCOI gene for *A. clausi*, *P. grani* and *A. discaudata*. *DdeI* and *PstI* restriction sites are indicated. Resulting fragment lengths are shown in base pairs (bp)

Figure

Fig. 5

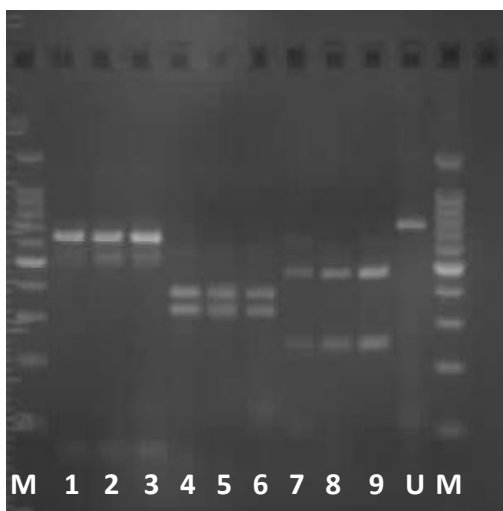


Fig. 5 Amplification of a region of mtCOI gene from whole *Acartia* adults and subsequent RFLP analysis with restriction enzymes *DdeI* and *PstI*. M=100bp DNA markers, Lanes 1-3 = restriction profile for *P. grani*; Lanes 4-6 restriction profile for *A. discaudata*; Lanes 7-9 = restriction profile for *A. clausi*; Lane 10 = undigested amplicon.