Species-specific polymerase chain reaction primer sets for the diagnosis of Tenacibaculum maritimum infection

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ABSTRACT: In this study the specificity and sensitivity of 2 primer pairs, MAR1–MAR2 and Mar1–Mar2, for the detection of Tenacibaculum maritimum were evaluated in parallel using 79 T. maritimum strains isolated from different fish species, as well as 53 representatives of related and unrelated bacterial species. Both primer pairs were species-specific for T. maritimum, since no amplification products were obtained from chromosomal DNA of the non-T. maritimum bacteria tested. However, whereas MAR1–MAR2 identified all the T. maritimum strains studied, producing a unique and clear PCR band of the expected 1088 bp length, the Mar1–Mar2 primer pair failed to amplify the 400 bp specific band in 3 sole isolates. To verify if these strains belonged to T. maritimum species, 2 endonucleases (Pvu I and Sac II) were selected as the most adequate enzymes to confirm the specificity of the MAR1–MAR2 amplified fragment. The digestion patterns obtained with both endonucleases supported the assignation of all the strains to T. maritimum. The sensitivity of both PCR detection methods was also different, showing a reduction of sensitivity in at least one order of magnitude of the Mar1–Mar2 primer pair in comparison with MAR1–MAR2. When the MAR–MAR2 PCR protocol was applied to different seeded turbot tissues, the detection limit was 10^2 to 10^4 T. maritimum cells per reaction. In addition, a nested PCR protocol for detection of this pathogens based on MAR1–MAR2 was developed, which increased the sensitivity by approximately 2 orders of magnitude, ranging from 1 to 250 T. maritimum cells per reaction depending on the tissue employed. The tissues that allowed the most easy detection of T. maritimum were the skin and mucus. Based on the findings reported here, we propose the nested PCR protocol as the most adequate for an accurate detection of T. maritimum in diagnostic pathology as well as in epidemiological studies of gliding bacterial disease of marine fish.

KEY WORDS: Tenacibaculosis · Tenacibaculum maritimum · Nested-PCR · Diagnosis

INTRODUCTION

Diseases caused by bacteria are recognized as major problems affecting the marine fish farming through the world (Austin & Austin 1999, Osorio & Toranzo 2002). Although pathogenic species have been described in most of the existent taxonomic groups, only a small number are responsible for important economic losses in extensive culture worldwide. Tenacibaculum maritimum (formerly Flexibacter maritimus) (Suzuki et al. 2001) has been included among these microorganisms, being the causative agent of gliding bacterial disease (or tenacibaculosis) in a great variety of valuable marine fish species such as turbot Scophthalmus maximus, salmon Salmo salar and Oncorhynchus kisutch, sole Solea senegalensis and Solea solea, gilthead seabream Sparus aurata, red and black seabream Pagrus major and Acanthopagrus schlegeli, and floun-

The disease is preliminarily diagnosed by the clinical signs of the affected fish which usually show eroded and hemorrhagic mouths, ulcerative skin lesions, frayed fins and tail rot. This diagnosis must be supported by the isolation of colonies of this pathogen on appropriate specific media (Pazos et al. 1996), followed by biochemical identification. This traditional culture-based method for the detection of the pathogen requires several days to weeks before results are obtained. In addition, one of the problems in the study of *Tenacibaculum maritimum* is the difficulty of distinguishing it from other phenotypically similar and phylogenetically related species, particularly those of the genera *Flavobacterium* and *Cytobacter* (Suzuki et al. 2001, Bader & Starliper 2002). Another disadvantage in the diagnosis of this infection is the problem of recognizing the colonies of *T. maritimum* among the mixed population that usually overgrow the *T. maritimum* colonies, due to the slow growth characteristic of *T. maritimum*.

All these features make DNA-based methods a very worthwhile approach for diagnosis of tenacibaculosis. Polymerase chain reaction (PCR) has been established as an alternative to traditional detection methods in microbial diagnostics due to the fact that this technique provides a powerful tool for accurate identification of the pathogen from plate cultures, as well as from fish tissues (Cunningham 2002, Osorio & Toranzo 2002). Two PCR primer pairs have been designed for the detection of *Tenacibaculum maritimum* using the 16S ribosomal RNA (rRNA) gene as target. Toyama et al. (1996) selected a pair of primers (MAR1 and MAR2; positions 190 to 206 and 1262 to 1278 respectively in the *Escherichia coli* 16S rRNA numbering system), flanking a 1088 base pair (bp) fragment, which could differentiate *T. maritimum* from the related species *Flavobacterium branchiophilum* and *F. columnare*, as well as from several other fish pathogenic bacteria. More recently, Bader & Shotts (1998) also selected a pair of *T. maritimum* species-specific PCR primers (Mar1 and Mar2; positions 77 to 98 and 456 to 476 respectively) from unique sequence stretches within this gene, delimiting a 400 bp DNA fragment.

However, despite the potential of PCR detection, these primers sets have not been tested simultaneously in order to compare their efficiency using a wide range of strains isolated from different species of marine fish and the environment. In the same way, neither Toyama et al. (1996) or Bader & Shotts (1998) determined the actual detection limits of each primer set with pure and mixed cultures, as well as in fish samples. Therefore, the aim of this study was to examine the specificity and sensitivity of the 2 PCR methods described previously for the identification of the *Tenacibaculum maritimum* strains. In addition, primers described by Toyama et al. (1996) were used to develop a nested PCR method for specific diagnosis of marine tenacibaculosis in fish samples.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** The bacterial strains used in the primer specificity studies are listed in Tables 1 & 2. This collection comprises 76 *Tenacibaculum maritimum* strains isolated from 7 different fish species from 1995 to 2003 (Table 1) and 3 reference strains (NCIMB 2153, 2154T and 2158) from the National Collection of Industrial and Marine Bacteria (Aberdeen, UK) (Table 1). These belong to the main serotypes and clonal lineages described within this pathogen (Avendaño-Herrera et al. 2004a,b). For all experiments, these strains were routinely cultured on *Flexibacter maritimus* Medium (FMM) agar (Pazos et al. 1996) and incubated at 20°C for 72 h. Before the assay, all bacterial strains were confirmed as *T. maritimum* using biochemical tests and serological assays according to Avendaño-Herrera et al. (2004a). In addition, 5 reference strains of other species of the genus *Tenacibaculum*, *T. ovolyticum*, *T. mesophilum* and *T. amylyticum* (NBRC 15947, 15992, 16307T, 16308 and 16310T), obtained from the NITE (National Institute of Technology and Evaluation) Biological Resource Center (Osaka, Japan), 27 strains of related filamentous bacterial species obtained from fish and seaweed and 21 pathogenic fish bacteria were also included as negative controls in the PCR (Table 2). Stock cultures were stored at −70°C in Cryo-bille tubes (AES Laboratory).

**DNA extraction from pure and mixed bacterial culture.** Chromosomal DNA was extracted using InstaGene Matrix (Bio-Rad) for pure bacterial cultures as previously described by Romalde et al. (1999). Briefly, *Tenacibaculum maritimum* and other non-*T. maritimum* bacterial species colonies were collected from the appropriate agar plates, suspended in 1 ml of sterile distilled water and centrifuged at 12 000×g for 1 min. After the supernatants were removed, the pellets were resuspended in 200 µl of InstaGene Matrix and incubated at 56°C for 30 min. Then, the suspensions were mixed at high speed for 10 s and boiled in a water bath for 8 min. The lysates were mixed again at high speed and then were spun at 12 000×g for 3 min. The concentration and quality of each DNA sample was examined spectrophotometrically at 260 nm. DNA from pure
cultures was adjusted to a concentration of 30 ± 2 ng µl⁻¹. All DNA was maintained at −20°C until used for PCR reactions. All the experiments were carried out with DNA obtained in 3 different extractions for each bacterial strain.

**PCR amplification.** All PCR amplifications were performed with the commercial kit Ready-To-Go™ PCR beads (Amersham Pharmacia Biotech), which included all the reagents needed for the PCR reactions (buffer, nucleotides and Taq DNA polymerase), with the exception of the specific primers and DNA template. Two species-specific primer pairs synthesized by Sigma-Genosys were used for the identification of *Tenacibaculum maritimum*: primers MAR1 (5′–AATGCGATCGTAAATA–3′) and MAR2 (5′–CGCTCTTGTGCGAGA–3′) described by Toyama et al. (1996), and primers Mar1 (5′–TGATGCTGCTACAGATGA–3′) and Mar2 (5′–AAATACCTACTCGTAGTACG–3′) designed by Bader & Shotts (1998), which give amplification products of 1088 and 400 bp respectively. One µl of each DNA solution and 2 µmol of each primer were used in the amplification reaction. All samples were subjected to 40 cycles of amplification in 2 different thermal cyclers. Reaction mixtures (25 µl) were simultaneously amplified in T Gradient Thermocycler (Biometra) and Mastercycler personal (Eppendorf) apparatus. The amplification cycles used for denaturation, primer annealing to the template and primer extension were carried out according to each published PCR protocol. Negative controls, consisting of the same reaction mixture but with sterile distilled water instead of template DNA, were included in each batch of PCR reaction. The reproducibility of the results was assessed by repetition of the amplifications in at least 3 independent PCR assays.

**Analysis of PCR products.** Amplified products were detected by horizontal 1% (w/v) agarose gel electrophoresis for 60 min at 100 V in TAE 1× (0.04 M Tris, 0.0001 M EDTA, pH 8.0) electrophoresis buffer, visualized using 0.06 µg ml⁻¹ of ethidium bromide (Bio-Rad) and photographed under UV light and computer digitized (Gel Doc 100, Bio-Rad). A 50 to 2000 bp ladder (Sigma Chemical) was used as a molecular mass marker. The presence of a single product of the appropriate size, identical to the reference strains, was considered as a positive result.

**Restriction enzyme digestion.** After the analysis of all PCR products, the isolates identified as *Tenacibaculum maritimum* with only one of the primer sets were subjected to restriction enzyme digestion. The MapDraw software from the Lasergene Navigator package (DNASTAR) was used to select the most adequate and specific restriction endonucleases for the 1088 bp amplification product. This selection was made on the basis of number, diversity and distribution of restriction fragments produced. Briefly, 2 µl of PCR product from each strain amplified by MAR1 and MAR2 was digested with 30 units of the restriction enzyme PvuII (Amersham Biosciences) according to the instructions of the manufacturer. Ten µl of the digested solution was analysed by horizontal electrophoresis. The restriction fragments of the PCR products from the reference strains were considered as positive controls.

**Determination of PCR sensitivity from pure and mixed cultures.** The detection limits of both species-specific primer sets were evaluated using bacterial suspensions of all reference strains. Pure colonies of *Tenacibaculum maritimum* were picked from FMM agar plates, adjusted to contain 10⁹ cells ml⁻¹ (McFarland Scale 4) and serially diluted in 0.85% sterile saline solution from 10⁹ to 10 cells ml⁻¹. To determine the usefulness of the primer sets to amplify template DNA of *T. maritimum* from mixed cultures, bacterial suspensions of *Listonella anguillarum* (RB2), *Photobacterium damselae* subsp. *piscicida* (TW 398), *Aeromonas salmonicida* (ATCC 14174) and *Lactococcus garvieae* (TW94W) were employed. Thus, bacterial mixtures of *T. maritimum* with 2 other different fish pathogens were prepared for PCR sensitivity testing. To each dilution of the *T. maritimum* strain, 100 µl of each bacterial suspension that contained approximately 10⁷ cells ml⁻¹ was added. Colony forming units (CFU) were estimated by plating each dilution onto appropriate media and counting the bacterial colonies produced.

All the dilutions of pure and mixed cultures were centrifuged at 12 000 × g for 2 min and washed twice with sterile distilled water to extract genomic DNA as before. Conditions for PCR amplification and electrophoresis were the same as described above.

**Determination of PCR sensitivity from seeded fish samples.** On the basis of the results obtained using bacterial cultures, PCR sensitivity was determined employing DNA extracted from in vitro seeded mucus, skin, kidney and blood, using the Toyama et al. (1996) protocol. All the tissue samples were taken from healthy *Tenacibaculum maritimum*-free juvenile turbot (12 g body weight). Mucus samples were obtained with a sterile 10 µl plastic loop or with a sterile scalpel blade by cutting the fish on the lateral line (Evans et al. 2001). The samples were then homogenized with 100 µl in phosphate buffered saline (PBS, pH 7.4) by repeated pipetting. In the case of skin and kidney, samples were collected using aseptic technique and divided into 500 and 30 mg pieces respectively. One hundred µl of blood samples were collected by vein puncture with a syringe and processed immediately. Each fish sample was seeded with 100 µl of the different dilutions of a culture suspension and homogenized
for 60 s. The 3 reference strains of *T. maritimum* were tested in parallel. After incubation for 1 h, DNA extraction was performed with Dynabead® DNA DIRECT™ (Dynal) following the manufacturer’s instruction, with the exception of blood samples, from which DNA was extracted as described above for bacterial culture. Non-inoculated fish samples, employed as negative controls, were processed in the same manner but with PBS instead of bacterial dilution. DNA obtained from the different tissues were maintained frozen at –20°C until used for PCR reactions. Conditions for PCR amplification and electrophoresis were the same as described above. In order to determine the reproducibility of results, all traits were assessed by repetition in at least 3 independent PCR assays, in which DNA obtained in 3 different extractions were employed.

**Nested PCR amplification.** To increase the sensitivity of the Toyama et al. (1996) PCR assay, a nested PCR was conducted on the whole dilution series from pure and mixed cultures, as well as infected tissues and blood with each reference strain. The first round of PCR was performed according to a previously described protocol (Edwards et al. 1989) with minor modifications. We used the same commercial reaction mixture and the universal primer pair pA (5’–AGAGTTGGATCCTGGCTCAG–3’) and pH (5’–AAGGAGGTGATCCAGCCGCA–3’) but the number of cycles was shortened to 25. These primers amplify the region from position 20 to position 1521 (*Escherichia coli* numbering system) of the 16S rRNA gene. After the first amplification by external primer was finished, 1 µl of each PCR product was used as template for the second amplification by the species-specific primer set MAR1 and MAR2. PCR samples were examined by gel electrophoresis and the samples were considered positive if the anticipated 1088 bp product was observed.

**RESULTS**

**PCR amplification**

To test the specificity of each primer pair on the identification of *Tenacibaculum maritimum*, DNA extracted from 79 strains of the target species as well as from 53 strains of other species were used for the PCR with each of the primer pairs. The MAR1–MAR2 primer pair identified all the *T. maritimum* strains studied, producing a unique and clear PCR product of the expected 1088 bp length. In contrast, the Mar1–Mar2 primer pair was capable of amplifying the 400 bp specific band from 76 of 79 strains of *T. maritimum* analyzed. The 3 *T. maritimum* strains (AZ202.1, AZ203.1 and PC438.1) not recognized by the Bader & Shotts (1998) pair of primers were isolated from sole (Table 1). Regardless of the primer set employed, amplification products were not obtained from the other bacterial species tested (Table 2).

When both PCR protocols were compared using 2 different thermal cyclers, no differences were detected in the results of the amplification products. In addition, reproducibility of PCR results was demonstrated since the same results were obtained in at least 3 independent PCR assays for each primer set tested.

**Restriction enzyme digestion**

To verify if the 3 isolates from sole only recognized using the Toyama et al. (1996) primer set belonged to *Tenacibaculum maritimum* species, 2 endonucleases (*Pvu* I and *Sac* II) were selected as the most adequate to confirm the specificity of the amplified fragment. The

<table>
<thead>
<tr>
<th>Type of samples</th>
<th>Isolation source</th>
<th>No. of strains</th>
<th>No. of strains identified with sets MAR1–MAR2</th>
<th>Mar1–Mar2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strains isolated from diseased fish (kidney and/or skin)</td>
<td>Scophthalmus maximus</td>
<td>49</td>
<td>49</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>Solea senegalis</td>
<td>17</td>
<td>17</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>Solea solea</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Sparus aurata</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Salmo salar</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Seriola quinqueradiata</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Dicentrarchus labrax</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Reference strains</td>
<td>Acanthopagrus schlegeli</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>T. maritimum NCIMB2153</td>
<td>Pagrus major</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>T. maritimum NCIMB 2154T</td>
<td>Solea solea</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>T. maritimum NCIMB 2158</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 1. *Tenacibaculum maritimum* strains used in this study and the results from the specific single PCR detection methods employed. NCIMB: National Collection of Marine and Industrial Bacteria (Aberdeen, UK)
restriction profiles of the PCR products obtained with the Toyama et al. (1996) primers, including the isolates not recognized as *T. maritimum* by Mar1 and Mar2 were identical for all the strains tested (Fig. 1), and coincident with the expected sizes calculated using the MapDraw software. Thus, endonuclease *Pvu* I rendered 2 fragments of 400 and 688 bp, while the enzyme *Sac* II produced 2 bands of 336 and 752 bp.

### Detection limit for the primer pairs

The sensitivity and robustness of each species-specific PCR primer set were determined by amplification of DNA extracted from different bacterial suspensions (10⁸ to 10 cells ml⁻¹) of 3 *Tenacibaculum maritimum* reference strains (Table 3). When the MAR1–MAR2 primer pair was tested, the expected 1088 bp PCR product was obtained with samples containing as low as 80 to 500 cells per PCR tube (Fig. 2A). With respect to Mar1–Mar2 primer pair, a fragment of the expected size (400 bp) was also amplified, allowing the detection of 8 x 10² to 5 x 10³ *T. maritimum* cells per PCR tube (Fig. 2B).

When DNA extracted from mixed cultures were used as template in single PCR amplification, the limits of detection of the *Tenacibaculum maritimum* with the 2 primer sets were identical to the level of amplification obtained with DNA from pure culture of reference strains (Table 3). Therefore, the other species of fish pathogens assayed did not interfere in the PCR sensitivity, even when high concentrations of non-*T. maritimum* strains were employed (4.0 ± 1 x 10⁶ CFU ml⁻¹).
With the knowledge that the Toyama et al. (1996) protocol allowed the confirmation of all *Tenacibaculum maritimum* strains examined and due to the best sensitivity achieved with this protocol in pure and mixed cultures, only the primer pair MAR1–MAR2 was tested with DNA extracted from the experimentally seeded fish tissues and blood. Application of the PCR protocol described above showed that *T. maritimum* could be detected in mucus, skin and blood at a detection level on the order of 10² to 10⁴ cells per reaction. However, no amplification product was observed when DNA samples of kidney were tested (Table 3). When the tissues used as negative controls were tested with the same primers, no amplification occurred. Finally, the total time for the procedure, including DNA extraction from samples, amplification and gel electrophoresis, was shorter than 7 h.

**PCR sensitivity from infected fish samples and blood**

Table 3. Sensitivity obtained with the different PCR protocols evaluated in single- and nested-PCR using pure and mixed cultures, as well as fish samples seeded with *Tenacibaculum maritimum*. CFU: colony forming units; NA: no amplification product; NT: not tested

<table>
<thead>
<tr>
<th>Cultures</th>
<th>MAR1–MAR2</th>
<th>Nested-PCR</th>
<th>Single PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure</td>
<td>80–500</td>
<td>1.6 × 10⁴–1.1 × 10⁵</td>
<td>1–5</td>
</tr>
<tr>
<td>Mixed</td>
<td>80–500</td>
<td>1.6 × 10⁴–1.1 × 10⁵</td>
<td>1–5</td>
</tr>
<tr>
<td>Fish samples</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucus</td>
<td>1–2 × 10⁴</td>
<td>5–6 × 10⁶</td>
<td>1–3</td>
</tr>
<tr>
<td>Skin</td>
<td>160–250</td>
<td>1–1.5 × 10⁴</td>
<td>1–3</td>
</tr>
<tr>
<td>Kidney</td>
<td>NA</td>
<td>160–250</td>
<td>1.7–2.3 × 10⁵</td>
</tr>
<tr>
<td>Blood</td>
<td>1–3 × 10⁴</td>
<td>2–4 × 10⁶</td>
<td>5100–200</td>
</tr>
</tbody>
</table>

*The values correspond to the amplification range of the 3 reference strains in triplicate*

**Nested PCR amplification**

When 1 µl of the first round product amplified with the universal primer set was used as a template for species-specific primer in nested PCR assays, the detection limit of all assays decreased by approximately 2 to 3 orders of magnitude compared with conventional PCR assay, regardless of the type of samples employed (Table 3). Then, when pure and mixed cultured *Tenacibaculum maritimum* were used in DNA extraction and nested PCR amplification, the detection limit achieved was about 1 to 5 bacteria per PCR tube (which corresponds to 1.6 × 10² to 1.1 × 10⁴ CFU ml⁻¹). In the case of DNA extracted from mucus and skin samples the nested PCR methods allowed the detection of approximately 1 to 3 *T. maritimum* cell per PCR tube (Fig. 3), and showed a reduction of sensitivity in 1 or 2 log-units for the blood samples. Although the single PCR did not yield amplification products in kidney samples, the nested PCR produced a unique and clear fragment of the expected size (1088 bp) when using this tissue, a sensitivity of approximately 2 ×
One of the most critical steps in the study of bacterial fish diseases is the correct identification of the infectious agent. Rapid diagnosis of outbreaks is essential for an effective control, but current microbiological methods based on culture and biochemical characterization are time-consuming. To overcome these difficulties, DNA-based methods have been developed in recent years for the fast and sensitive diagnosis of the bacterial pathogens in aquatic species. In the case of *Tenacibaculum maritimum*, Toyama et al. (1996) and Bader & Shotts (1998) used the 16S rRNA gene sequence available in public databases for the design of specific primers to be employed in the PCR. However, until now the specificity and sensitivity of these primer sets have not been evaluated in parallel in order to propose the best PCR protocol to detect this pathogen in research laboratories.

Comparing the specificity of the 2 PCR protocols, the sequence of both primer pairs were species-specific for *Tenacibaculum maritimum*, and no amplification products were obtained from chromosomal DNA of other non-*T. maritimum* bacteria tested. These results agree with those reported by the designers of both sets of primers (Toyama et al. 1996, Bader & Shotts 1998). However, when all bacteria listed in Table 1 were screened, the number of strains identified by each primer pair was different. Whereas all the isolates rendered the specific amplicon when the MAR1–MAR2 (Toyama et al. 1996) primer pair was used, the Mar1–Mar2 (Bader & Shotts 1998) primer set failed to identify 3 isolates. However, the digestion patterns of the MAR1–MAR2 amplified products of 1088 bp with the restriction enzymes *Pvu* I and *Sac* II allowed us to confirm the assignation of all the strains to the *T. maritimum* species.

Recently, Cepeda et al. (2003) also employed the Mar1–Mar2 primer pair, making modifications to the PCR programme described by Bader & Shotts (1998) in order to reduce the time to achieve the diagnosis of *Tenacibaculum maritimum*. However, when we used this protocol with pure cultures of *T. maritimum* strains using the same extraction system and identical conditions as described by these authors, none of isolates tested generated an amplification signal (data not shown). This failure should probably be attributed to the extremely short time (2 s) employed in the annealing step, although we can not rule out the influence of other variables which theoretically could alter the test performance, such as source of reagents or disposable supplies.

In the present study, the sensitivity of both PCR detection methods was also different. It has been reported that sensitivity of PCR varies depending on the size of amplicons, and generally primers yielding smaller amplicons will be able to detect lower levels of target organisms (Otta et al. 1999, Hossain et al. 2001). However, our results using either pure or mixed cultures showed a reduction of sensitivity in at least one order of magnitude of the Mar1–Mar2 primer pair in comparison with MAR1–MAR2, although the former primer set yield a shorter amplicon (approximately 700 bp) than the latter one.
Our study has shown that when the single PCR protocol of Toyama et al. (1996) was used with DNA extracted from seeded mucus, skin tissue and blood samples the sensitivity level was 10- to 100-fold lower than that observed with pure and mixed cultures, probably due to presence of host DNA and undefined inhibitors of PCR (Wilson 1997). This level of sensitivity is probably sufficient to detect *Tenacibaculum maritimum* acute infections in fish, but not when the pathogen is present in very low numbers in asymptomatic or carrier fish.

A nested PCR approach greatly increases the sensitivity of detection, and can be useful for the detection of very low copy numbers of template (Osorio & Toranzo 2002). In our case the nested PCR assay supports this fact since an increase in sensitivity of at least 100 times was observed compared to conventional PCR; this sensitivity is in the same range as those achieved for other bacterial fish pathogens (Osorio et al. 1999, Taylor & Winton 2002). Although, we found that all tissues sampled could be used to detect the bacterium in a nested PCR protocol, negative results were obtained when using DNA extracted from kidney samples in a single PCR. Similar problems in detecting other fish pathogens in kidney have been attributed to samples in a single PCR. In our case the nested PCR assay sup-

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In summary, on the basis of the results obtained in this work we propose the Toyama et al. (1996) PCR procedure as the most adequate for an accurate detection of *Tenacibaculum maritimum* in diagnostic pathology as well as in epidemiological studies of gliding bacterial disease of marine fish. In addition, the nested PCR provides at least 100 times more sensitivity than conventional PCR for the detection of *T. maritimum* not only in pure and mixed bacterial cultures, but also in experimentally seeded fish tissues. The accuracy of this method is currently being investigated with natural samples from different fish farms in order to evaluate its efficacy, under field conditions, in detecting the pathogen in diseased and asymptomatic carrier fish.

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