**In situ** hybridization on whole larvae: a novel method for monitoring bivalve larvae

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**ABSTRACT:** Accurate and efficient identification to the species level of early larval stages has long been a problematic step in the study of marine invertebrates, due to the extremely small size of the larvae and their lack of diagnostic morphological characters. It is nonetheless, a prerequisite for any ecological study. As a consequence, a number of molecular approaches, mostly based on the PCR technique, have been developed over the last decade. We developed a method relying on specific rRNA-targeted oligonucleotide probes for **in situ** molecular hybridization using whole larvae. A colorimetric reaction following the hybridization allows signal detection with a light microscope. A total of 9 probes have been developed, targeting the species that are major components of benthic communities in several European bays. The method can be applied to roughly sorted wild plankton samples, and some steps could be automated. It is relatively inexpensive to implement and does not require costly equipment or expensive staff training. The overall larval morphology is preserved, allowing visual inspection. The quantitative aspect of the approach is another asset. Field-based studies of larval distribution and behaviour are possible applications of the method, as large numbers of samples can be screened to meet the requirements of adequate spatial and temporal coverage. It is also relevant for routine monitoring of target species, such as species of commercial interest, bio-indicators or invasive species.

**KEY WORDS:** Bivalve larvae · Species identification · **In situ** hybridization · Oligonucleotide probes · 18S rDNA

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**INTRODUCTION**

Pelagic larvae represent a decisive phase in the life cycles of many marine animals, playing a fundamental role in population dynamics and connectivity. In the case of many sessile benthic invertebrates, pelagic larvae represent the exclusive dispersal vector. Precise species identification is prerequisite information for larval ecological studies. The identification of marine invertebrate larvae, especially bivalves, remains nonetheless a notoriously difficult target. Early developmental stages of bivalve larvae can measure <100 µm, and often lack reliable identification characters (e.g. Rees 1950). Morphology can be used for identification at later developmental stages for some species, but this approach requires rare expertise, and long preparation of samples when scanning electron microscopy is used for precise observation (e.g. Fuller & Lutz 1989). Larval identification guides are also restricted to particular geographical areas where such descriptive work has been undertaken for commonly occurring species, and no such comprehensive guide is available for European bivalve species (Larsen et al. 2005).

A variety of molecular approaches, including both DNA-based and immunological methods, have been developed over the last decade to facilitate identification of bivalve larvae (reviewed in Garland & Zimmer 2002). The scope of such methodological developments ranges from one or a few focused species, such as invasive species (Pie et al. 2006) or species of commercial interest — oysters (e.g. Morgan & Rogers 2001),

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scallops (e.g. Frischer et al. 2000, Paugam et al. 2003), surfclams (Bell & Grassle 1998), cockles (André et al. 1999) and mussels (e.g. Wood et al. 2003, Lorenzo-Abalde et al. 2005)—to a wider range of commonly found species (e.g. Hosoi et al. 2004, Larsen et al. 2005). While single-species focused studies have used varied approaches, such as immunology (e.g. Lorenzo-Abalde et al. 2005), RAPDs (André et al. 1999), micro-satellite markers (Morgan & Rogers 2001), RFLPs (e.g. Comtet et al. 2000), dot-blot (e.g. Frischer et al. 2002), PCR amplification and hybridization (Frischer et al. 2000), and multiplex PCR (e.g. Wood et al. 2003), identification methods developed for a range of species mainly rely on derivations of 2 PCR methods, viz. RFLP (e.g. Hosoi et al. 2004) and multiplex PCR (e.g. Larsen et al. 2005).

In their review, Garland & Zimmer (2002) describe the ideal identification technique as (1) involving no direct sub-sampling of organisms from a sample, (2) effective for intact organisms, (3) producing a detectable result through image analysis, (4) relatively inexpensive to develop, (5) relatively inexpensive to use and (6) producing accurate and repeatable results. The methodological development towards bivalve larvae identification presented here was designed to take most of these criteria into account.

In situ hybridization coupled with fluorescent detection (FISH) has been successfully applied as an identification method to a variety of microorganisms, including those from the marine environment, allowing visualization and enumeration of microbial populations (reviewed in Amann et al. 1995), planktonic Archaea and bacteria (e.g. DeLong et al. 1999), small phytoplankton (e.g. Tyrrell et al. 1997, Simon et al. 2000), as well as the identification and localization of bacterial symbionts (e.g. Haygood & Davidson 1997). Application of the methodology to whole invertebrate larvae has been restricted so far to evolution and development research, and has targeted a few marine model species, such as ascidians (e.g. Ueki et al. 1994), sea urchins (e.g. Yaguchi & Katow 2003), gastropods (Nederbragt et al. 2002), and the oyster *Crassostrea gigas* (Fabiox et al. 2004).

We show that in situ hybridization can be used to detect specific bivalve larvae from wild plankton samples. The method relies on specific DNA oligo-probes designed from sequences obtained for a partial, variable region of the 18S rRNA coding gene. Ribosomal RNA-targeted oligonucleotide probes have become a widely used tool for ecological studies in marine microbiology (reviewed in Amann et al. 1995), as their hybridization target sites, the ribosomes, are numerous in single whole cells, allowing efficient hybridization signal detection. The 18S rRNA gene shows sufficient variation in bivalves to be used for designing species-specific probes (Bell & Grassle 1998, Frischer et al. 2002, Larsen et al. 2005). The development of the method involved setting up an in situ hybridization protocol using 18S oligo-probes applicable to whole bivalve larvae, designing species-specific probes for the bivalve species commonly found in European coastal waters, and checking for a specific, easily detectable signal with the designed probes.

**MATERIALS AND METHODS**

The first step of this project was to develop a general protocol for in situ hybridization using whole bivalve larvae collected from wild plankton samples. The aim of this approach was to ensure that any bivalve larva potentially occurring in coastal water samples, whatever its developmental stage or species, could be identified reliably through a detectable hybridization signal. This implied the use of a positive and a negative probe that had already been used for in situ hybridization in previously published studies. This step also led to selection of markers for subsequent specific probe design.

The second step of the project was to build a sequence database to design specific probes. Probe specificity was experimentally assessed through 2 approaches, i.e. detection on dot-blots, using PCR products obtained from adult bivalves as the target, and in situ hybridization using whole larvae collected separately from parents of 4 known species held in hatcheries. In situ hybridization tests were also conducted on samples composed of mixed identified larvae, mimicking simple wild samples.

**Development of a protocol for in situ hybridization on whole larvae.** Plankton sampling and fixation of bivalve larvae: Plankton was sampled in Morlaix Bay off the northern coast of Brittany (France) (48° 40' N 3° 53' W) using a 63 µm mesh size WP2 plankton net. Samples were collected throughout the year; seawater temperatures varied from 8°C to 16°C. Samples were kept in seawater until arrival at the laboratory where veliger-type bivalve larvae were isolated under a dissecting microscope. Larvae were then transferred to a 20% MgCl₂ muscle relaxant solution in order to ensure valves would be open to allow subsequent probe penetration. A 4% paraformaldehyde (PFA) in phosphate-buffered saline solution (PBS) was used as a first fixative for 1 h, before transfer to a microtube containing 96% ethanol at −20°C for long-term storage.

**In situ hybridization:** The reactions were carried out using 50 µm mesh baskets (containing the larvae) placed into cell culture clusters. Samples were first rinsed with gradual series of ethanol/PBT (phosphate-buffered saline containing 0.1% Tween 20) solutions, and then twice in PBT.
Two hybridization temperatures were tested (50 and 55°C), and 2 concentrations of formamide in hybridization buffer (10 and 50%). Pre-hybridization was set for at least 1 h at hybridization temperature, using pre-warmed hybridization buffer (5× saline solution citrate buffer (SSC), 0.2 mg ml⁻¹ salmon sperm DNA, 0.1 mg ml⁻¹ heparin, 2 μL ml⁻¹ Tween 20, 5% dextran sulfate) containing formamide. Larvae underwent hybridization overnight, using a 1 ng μl⁻¹ probe concentration. Two oligonucleotide DNA probes were selected through ProbeBase (Loy et al. 2003). A universal probe, targeting the 18S rRNA of all eukaryotes, EUK516 (Aman et al. 1990), was chosen as a positive control, and a non-specific probe, targeting the 23S rRNA of 2 bacterial strains, Candidatus Brocadia anamoxoxidans and Candidatus Kuenenia stuttgartiensis, Amx1900 (Schmid et al. 2001), was chosen as a negative control. Both were modified with digoxigenin at their 3’ end (MWG Biotech).

Three post hybridization washes were performed using hybridization buffer at hybridization temperature, the first for 1 h and the following 2 for 30 min. Two additional 5 min washes with gentle agitation were carried out with PBT. Larvae were subsequently incubated with a blocking solution (10% goat serum in PBT) for at least 1 h prior to overnight incubation at 4°C with the same solution containing 1‰ Dig-AP conjugate (Roche). Three 10 min washes with gentle agitation were performed with PBT, followed by 2 washes with colour-developing buffer (100 mM Tris-HCl, pH 9.5, 100 mM NaCl, 50 mM MgCl₂, 0.01% Tween 20). Larvae were incubated with BM Purple AP substrate (Roche) at 37°C in the dark until colour development. When a positive signal was detected under the dissecting microscope, the reaction was stopped by PBT washes. Photomicrographs of the larvae were taken using magnifications varying from 100× to 200× and using a Nikon Coolpix 4300 camera. The experiments were repeated 5 times to check consistency of results.

**Design of species-specific probes.** Using the universal primers from Larsen et al. (2005), we sequenced a total of 123 bivalve individuals encompassing 24 species commonly found in European coastal waters. Whenever possible, more than one individual per species was sequenced in order to include intra-specific variability in subsequent analyses.

DNA was extracted using a phenol chloroform protocol, as described in T. Comtet & M. C. Le Goff-Vitry (unpubl.). Each 25 μL PCR sample included 2 mM MgCl₂ (Euroclone), 1.25 μmol of each dNTP (ABgene), 13 pmol of primers Uni 1304F and Uni 1670R (MWG Biotech), 2.5 μl of 10× reaction buffer (Euroclone), 1 unit of Taq Polymerase (Euroclone) and 1.3 μl of 1:100 diluted extracted DNA. PCR was performed with a GeneAmp® PCR System 2700 (Applied Biosystems) thermocycler using an initial denaturation at 94°C for 4 min, followed by 35 cycles of 94°C for 1 min, 57°C for 1 min, 72°C for 1 min, and a final extension at 72°C for 7 min. PCR products were visualised on a 2% agarose gel stained with ethidium bromide and subsequently purified using the Multiscreen® PCR 96 kit (Millipore). Sequencing was performed on an ABI 3100 sequencer using BigDye terminator chemistry (PerkinElmer), following the manufacturer’s protocol. Sequencing was done in both directions. The resulting sequences were checked using Chromas Lite 2.01 (Technelysium, Copyright 1998–2005) and combined together with homologous bivalve sequences obtained from GenBank to build a database total of 197 sequences (Appendix 1, available as Supplementary Material at www.int-res.com/articles/suppl/m343p161_app.pdf).

Winprobe software (Pozhitkov & Tautz 2002) was used to design species-specific oligo probes using this database. When low sequence resolution did not allow the design of species-specific probes, higher taxonomic levels were targeted. Probe specificity was further tested using BLAST (Basic Local Alignment Search Tool: http://130.14.29.110/BLAST/) and selecting the ‘search for short, nearly exact matches’ option.

**Probe specificity tests by dot-blot.** Ideally, all designed probes would be tested through in situ hybridization, using identified larvae from each target species. However, obtaining such larval samples is very demanding in terms of equipment, expertise and time, as it requires conditioning pools of adult bivalves, inducing spawning and rearing larvae at least for a few days. A much more straightforward approach consists of testing probes by dot-blot, using whole 18S PCR products obtained from adult DNA and blotted onto a membrane. This is a common procedure with specific probe development for FISH (fluorescent in situ hybridization; e.g. Edgcomb et al. 1999, Simon et al. 2000). The 9 probes we designed (Table 1) were tested against PCR products obtained from a total of 75 bivalve individuals belonging to 18 different species. Probes were tested individually on membranes containing 8 individuals of Mytilus sp. (Mytilus edulis / Mytilus galloprovincialis complex), 5 of Nucula nucleus, 8 of Ostrea edulis, 6 of Crassostrea gigas, 14 of Veneridae (5 of Venerupispullastra, 3 of Timoecia ovata, 1 of Tapes rhomboides and 5 of Ruditapes decussatus), 7 of Macoma balthica, 7 of Cerastoderma edule, 7 of Glycymeris glycymeris, 8 of Pectinidae (6 of Pecten maximus and 2 of Chlamys varia), and 1 individual each of the following non-target species: Tellina tenuis, Corbula gibba, Laevicardium crassum, Abra alba and Spisula ovalis.
Table 1. List of the 18S rRNA targeted DNA oligoprobes with the taxon specifically targeted

<table>
<thead>
<tr>
<th>Probe name</th>
<th>Probe sequence</th>
<th>Taxon targeted</th>
</tr>
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<tbody>
<tr>
<td>Cra-gig-18S</td>
<td>5'-CGACAGCAATAAGATCGTGTA-DIG</td>
<td>Crassostrea gigas</td>
</tr>
<tr>
<td>Ost-ed-18S</td>
<td>5'-ACAGTTTCGAGATCGTGTA-DIG</td>
<td>Ostrea edulis</td>
</tr>
<tr>
<td>Cer-ed-18S</td>
<td>5'-TTGACGCGGGGACGGATT-DIG</td>
<td>Cerastoderma edule</td>
</tr>
<tr>
<td>Mac-ba-18S</td>
<td>5'-GGCAGCGAATGACGACGTT-DIG</td>
<td>Macoma balthica</td>
</tr>
<tr>
<td>Myt-18S</td>
<td>5'-CCGACGCATAATGGGATCGG-DIG</td>
<td>Mytilus sp.</td>
</tr>
<tr>
<td>Nuc-18S</td>
<td>5'-ACGGAGCGATACCGATCGTG-DIG</td>
<td>Nucula sp.</td>
</tr>
<tr>
<td>Gly-18S</td>
<td>5'-CCCAATGATCGAAGAATAT-DIG</td>
<td>Glycymeris sp.</td>
</tr>
<tr>
<td>Pec-18S</td>
<td>5'-GCAAAGGATAAGCAGGCTT-DIG</td>
<td>Pectinidae</td>
</tr>
<tr>
<td>Ven-18S</td>
<td>5'-CGGGGACTCGATAGAGGATC-DIG</td>
<td>Veneridae</td>
</tr>
</tbody>
</table>

For PCR amplification of the whole 18S gene, we used the universal primers from Moon-van der Staay et al. (2001). Each 25 µl PCR sample comprised 2.6 mM MgCl₂ (Euroclone), 1.25 µl of each dNTP (Abgene), 2.5 µl of 10× reaction buffer (Euroclone), 1.25 unit of Taq Polymerase (Euroclone) and 2 µl of 1:100 diluted extracted DNA. PCR reactions were performed on a GeneAmp® PCR System 2700 (Applied Biosystems) using an initial denaturation at 95°C for 5 min, followed by 34 cycles of 95°C for 1 min, 55°C for 1 min 30 s, 72°C for 2 min, and a final extension at 72°C for 10 min. PCR products were visualised as before.

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Blotting of the PCR products onto a positively charged nylon membrane (Roche) was performed using a Bio-Dot® Microfiltration Apparatus (BioRad) according to the manufacturer’s instructions. The blotted membranes were subsequently incubated with hybridization buffer (5× SSC, 0.1% N-lauryl sarcosine, 0.02% sodium dodecyl sulfate (SDS) and 0.1 mg ml⁻¹ salmon sperm DNA) for 1 h at 42°C in a mini-hybridization oven (Appligene). Hybridization was carried out overnight at 42°C with a probe concentration of 7 pmol ml⁻¹. A total of 6 post-hybridization washes were performed, including 2 washes with 2× SSC 0.1% SDS at room temperature for 5 min, 2 washes with 2× SSC 0.1% SDS at 55°C for 10 min, and 2 final washes with 0.5× SSC 0.1% SDS at 55°C for 20 min. The membranes were then soaked in Buffer 1 (0.1M Tris HCl pH 7.5, 0.1M NaCl, 2 mM MgCl₂, 0.05% Triton 100) for 5 min at room temperature, and then with Buffer 2 (0.1M Tris HCl pH 7.5, 0.1M NaCl, 2 mM MgCl₂, 0.05% Triton 100, 2% Bovine Serum Albumin (BSA)) for 1 h at room temperature. Incubation with Buffer 1 containing Dig-AP conjugate (1/2000 dilution) (Roche) was carried out for 30 min at room temperature. Membranes were then washed (with rocking) 3 times with Buffer 1 for 10 min and then 3 times with Buffer 3 (0.1 M Tris HCl pH 9.5, 0.1 M NaCl, 0.05 M MgCl₂) for 10 min. They were finally incubated with NBT/BCIP (Roche) diluted in Buffer 3 (1/50 dilution) at 37°C in the dark until signal detection. The staining reaction was stopped by rinsing the membranes with Buffer 4 (20 mM Tris HCl pH 7.5, 5 mM ethylenediaminetetraacetic acid (EDTA)). Photographs of the membranes were taken with a Nikon Coolpix 4300 camera. The dot-blot test for each probe was repeated twice in order to check the consistency of the results. In situ test with designed probes. Larvae from 4 species of commercial interest, for which rearing is possible, were obtained from hatcheries. The specific probes for these 4 species were tested by in situ hybridization. Crassostrea gigas larvae were obtained from Station Expérimentale Ifremer d’Argenton (France), Pecten maximus larvae from Ecloserie du Tinduff (France) and Mytilus edulis and Ostrea edulis from Laboratoire Ifremer La Tremblade (France). C. gigas larvae were at the umboned stage, and their mean size was 184 µm. Two batches of P. maximus larvae were available: the youngest were 8 d old, with a mean size of 148 µm; the oldest were 15 d old and 212 µm in mean size. M. edulis larvae were 14 d old, with a mean size of 224 µm. O. edulis larvae were 10 d old, with a mean size of 174 µm. Fixation was carried out as for larvae isolated from plankton samples.

In situ hybridization tests were carried out using a range of hybridization conditions (hybridization temperatures from 46°C to 55°C, formamide concentrations in hybridization buffer from 10% to 50% and probe concentration from 0.25 ng µl⁻¹ to 4 ng µl⁻¹) and 10 individuals per species, for each probe. Cra-gig-18S, Pec-18S, Myt-18S and Ost-ed-18S probes (Table 1) were tested through in situ hybridization using their respective target larvae, but also using non-target larvae, in order to check probe specificity.

These probes were also tested through in situ hybridization using ‘artificial meroplankton samples’ containing a mixture of 10 larvae from each of the 4 species. Morphological characteristics and size differences allowed us to distinguish the different larval species after hybridization. Photographs of the larvae were taken under the microscope using a Nikon Coolpix 4300 camera.

RESULTS

In situ hybridization using rRNA-targeted oligoprobes on whole bivalve larvae from wild samples

A universal protocol for in situ hybridization on whole larvae applicable to wild plankton samples was
successfully developed. We obtained a suitable signal-to-noise ratio with the universal eukaryotic probe at both hybridization temperatures and formamide concentrations. Bivalve larvae isolated from plankton samples stained strongly with the universal probe, while no staining was observed either with the non-specific probe or with no probe (Fig. 1). The signal was detectable under the dissecting microscope after 1 h of incubation with the substrate, whatever the size or morphology of the veliger larvae.

Specific oligo-probes

The lengths of the partial 18S sequences obtained varied between 310 and 440 bp, and were variable enough to allow the design of species-, or genus-specific probes in most cases (Table 1). Out of a total of 9 designed probes, 4 are species-specific and 3 are genus-specific. When it was not possible to obtain species-specific probes, probes were designed to target the genus level (Mytilus, Glycymeris and Nucula), or the family level, in the cases of Pectinidae and Veneridae.

Probe specificity was first assessed through dot-blot tests, using whole 18S PCR products of approximately 1.5 kb obtained from adult bivalve DNA. These tests allowed checking the results of in silico specificity analyses. In 2 cases, probes had to be re-designed because cross-hybridization was encountered. The probe initially designed for Ostrea edulis cross-hybridized with Mytilus edulis and Crassostrea gigas individuals, and the first probe designed for Nucula cross-hybridized with Cerastoderma edule individuals, despite tentative optimizations (data not shown). Two new probes were therefore designed: Ost-ed-18S and Nuc-18S (See Table 1) and tested through dot-blot. At this stage, all tested probes showed a specific signal (Fig. 2). When testing Nuc-18S, all 5 Nucula nucleus individuals showed a strong hybridization signal, and a very faint signal appeared for one of the 7 Glycymeris glycymeris individuals. This result was considered not to compromise the specificity of the probe. The probes targeting the family level (Pec-18S and Ven-18S) showed a positive signal only with the individuals belonging to the species and genera from the targeted family.

A second series of in situ hybridization tests included mixtures of larvae from these 4 separate species to constitute an ‘artificial meroplankton sample’. The overall conservation of larval morphology and size after in situ hybridization allowed visual identification of each larva to species. Hybridization conditions were optimized for each probe, and the consistency of the results under these conditions was validated by repeating the experiment. The conditions giving the best results for each probe are presented in Table 2.
Fig. 2. Probe specificity tests through dot-blot. One probe was tested per membrane. A: Myt-18S (Target genus: *Mytilus*); B: Nuc18S (Target genus: *Nucula*); C: Ost-ed-18S (Target species: *Ostrea edulis*); D: Cra-gig-18S (Target species: *Crassostrea gigas*); E: Ven-18S (Target family: Veneridae); F: Mac-ba-18S (Target species: *Macoma balthica*); G: Cer-ed-18S (Target species: *Cerastoderma edule*); H: Gly-18S (Target genus: *Glycymeris*); I: Pec-18S (Target family: Pectinidae). The table gives the species identities of 18S PCR products blotted onto the membranes, according to their positions. Abbreviations are as follows: *M. sp.*: *Mytilus* sp.; *N. n.*: *Nucula nucleus*; *O. e.*: *Ostrea edulis*; *C. g.*: *Crassostrea gigas*; *V. p.*: *Venerupis pullastra*; *T. o.*: *Timoclea ovata*; *T. r.*: *Tapes rhomboides*; *R. d.*: Ruditapes decussatus); *M. b.*: *Macoma balthica*; *C. e.*: *Cerastoderma edule*; *G. g.*: *Glycymeris glycymeris*; *Pectinidae* (P. m.: *Pecten maximus*; C. v.: *Chlamys varia*); Non target species (T. t.: *Tellina tenuis*; Co. g.: *Corbula gibba*; L. c.: *Laevicardium crassum*; A. a.: *Abra alba*; S. o.: *Spisula ovalis*)
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Fig. 3: Specific probes tested on identified larvae through in situ hybridization. A: Crassostrea gigas; B: Pecten maximus; C: Mytilus edulis; D: Ostrea edulis. A, B, C, D hybridized with 1: Cra-gig-18S probe; 2: Pec-18S probe; 3: M yt-18S; and 4: Ost-ed-18S probe.

Fig. 4: Specific probes tested through in situ hybridization on samples of mixed larvae of known identity. (A) Cra-gig-18S probe, (B) Pec-18S probe, (C) M yt-18S, (D) Ost-ed-18S. Arrows indicate stained larvae.
DISCUSSION

**In situ** hybridization using whole larvae can be used as a reliable identification tool for bivalves. The protocol developed, relying on rRNA-targeted oligonucleotide DNA probes labelled with digoxigenin, has produced consistent results on bivalve larvae isolated from wild plankton samples, as well as on those collected from hatcheries. A partial region of the 18S gene was sufficiently variable to allow design of specific oligo-probes targeting key bivalve species (or higher taxonomic levels) present in North East Atlantic coastal water communities. Probe specificity was experimentally validated. These probes can be used on multi-species assemblages to specifically detect larvae belonging to target species. The value and pitfalls of the method, as well as its potential applications are discussed below.

**Value of in situ hybridization as an identification tool for bivalve larvae**

The approach was successfully applied to veliger-type bivalve larvae of various sizes isolated from plankton samples collected at different times of the year, and to larvae of different ages from hatcheries. It is also potentially applicable to earlier developmental stages, although this was not tested in the present study. This is a considerable asset for an identification method targeting widely diverse larval stages present simultaneously in aquatic environments, in comparison, for example, with morphological or immunological methods. One of the drawbacks of immunological methodology is that the signature molecule on which the reaction relies on is a protein, the expression level of which varies according to exogenous and endogenous factors (Garland & Zimmer 2002).

Specificity is an additional asset of **in situ** hybridization. Probe specificity tested **in silico** was experimentally validated through dot blot tests, and for 4 of them through **in situ** hybridization on whole larvae. The short size of oligonucleotide probes allows high specificity, as a single mismatch in hybridization causes a highly destabilizing effect (Edgcomb et al. 1999). Lack of specificity can be a problem with some identification methods; for instance cross-reaction with non-targeted species has been reported with immunological methods (Paugam et al. 2003).

Through **in situ** hybridization, each larva can be identified individually. This property confers high sensitivity and a quantitative aspect to the method, resulting in high-resolution data for subsequent applications. These are considerable advantages in comparison with molecular approaches relying on DNA extraction and PCR, such as RFLP, multiplex PCR, microsatellites and quantitative PCR. Firstly, PCR failure is common with DNA template extracted from individual larvae, especially after preservation with ethanol or chemical fixatives (Larsen et al. 2005). Moreover, when using as a template DNA extracted from a mixed plankton sample, a bias in estimating numbers of larvae present in a sample through PCR-based methods arises from the fact that various developmental stages are present simultaneously in wild samples, and the quantities of template DNA per larva varies by developmental stage.

**In situ** hybridization is a potentially high throughput approach, allowing simultaneous screening of large quantities of larvae. Larval ecology studies rely on extensive sampling because of the diluted and patchy distribution of larvae in time and space (Garland & Zimmer 2002). Processing such large samples has been a limiting factor so far (Garland & Zimmer 2002). **In situ** hybridization minimizes handling of individual larvae, saving time and limiting potential manipulation error. Throughput could be further enhanced by simultaneous use of differently labelled probes in order to detect several species per sample, as is already the case for phytoplankton and bacterial FISH studies (e.g. Wallner et al. 1993, DeLong et al. 1999). The method could also be coupled advantageously with image-analysis technology, avoiding potential subjectivity for routine screening of samples. Detection and enumeration of stained larvae is a straightforward application of image analysis software (Garland & Zimmer 2002), and coupling of fluorescent detection with a motorised scanning stage and an image analysis system has already been performed in bacterial studies using immunological methods (de Boer & Hall 1998). Most of the experimental procedure could be automated, as the larvae (placed in baskets for the **in situ** protocol) are simply transferred between wells containing different solutions.

An additional advantage of **in situ** hybridization on whole larvae is that it is non-destructive, preserving the overall larval morphology, and hence providing information on the size and shape of labelled larvae. Conservation of larval morphological characters (such as the umbo of *Crassostrea gigas* larvae, or hinge shapes), as well as the relative sizes allowed discrimination of species among larvae collected from multi-species assemblages. Such information can be valuable for crosschecking identification using morphological characters, and for biometric indices used in population dynamic studies. This is an advantage over most molecular approaches requiring the extraction of DNA as a first step.
Potential limitations of in situ hybridization on whole larvae

Tests conducted through in situ hybridization on whole larvae allowed us to monitor signal sensitivity and specificity on different types of samples including samples containing individuals belonging to the same species (Fig. 2) and those containing individuals from 4 different species mixed together (Fig. 3). These experiments allowed testing the effect of different parameters on the hybridization signal, as well as checking for the occurrence of false positives (non-specific staining) and false negatives (non-staining).

While tests carried out using only 1 larval species showed a specific and sensitive hybridization signal over a range of hybridization conditions, fine adjustments were necessary with samples of mixed species in order to achieve the best signal-to-noise ratio. Successful experiments using artificial meroplankton samples were conducted in order to reach optimal hybridization conditions for each individual probe (Table 2). The observations suggest that the application of each specific probe to wild samples may require experimental optimization. This might be explained by the fact that each oligonucleotide probe has a specific dissociation temperature, depending on its length and base composition (Wallner et al. 1993).

Increasing the hybridization temperature enhanced the stringency of the hybridization, until the dissociation temperature of the probe target hybrid was reached, causing hybridization failure (Amann et al. 1995). Formamide was added to the hybridization buffer at various concentrations, as it improves in situ accessibility by weakening the effects of hydrogen bonds responsible for higher-order structure of ribosomes (Amann et al. 1995). However, high formamide concentrations (50%) affect conservation of larval morphology. The optimal probe concentration was determined by testing a range of concentrations from 0.1 to 1 ng µl⁻¹. These correspond to intermediate probe concentrations producing an optimal signal to noise ratio in FISH applications (Wallner et al. 1993). At very low probe concentration (0.01 ng µl⁻¹), the number of probe molecules is not sufficient to bind to all available ribosomes, producing a weak signal, whereas probe concentrations that are too high (20 ng µl⁻¹) increase non-specific staining (Wallner et al. 1993).

Altering hybridization conditions avoided non-specific signals on mixed species samples in most cases. More difficulties were experienced with Ostrea edulis larvae, which showed non-specific staining in a range of conditions when different probes were tested, including the negative probe (Amx1900). However, specific and non-specific staining showed very different patterns and localisations on the larvae. Non-specific staining is commonly encountered with in situ hybridization on whole invertebrates, and might be due to some particular biochemical structure adsorbing the probe, or to some epitope the anti-Dig antibody recognizes (A. Chipman pers. obs.). The fact that non-specific staining was observed with probes for different nucleotide sequences suggests that it is not caused by mispairing of the probe to non-target sequences, but by the binding of the conjugated molecules to other cell components (Wallner et al. 1993). On the other hand, specific signals were either homogeneously distributed over the whole larva, or sometimes focused along the mantle exterior edge and the velum (Fig. 3, image C3 in the matrix). Both organs have key functions in larval development; the edge of the mantle is notably involved in shell secretion, and the velum allows locomotion and acquisition of food (Ruppert & Barnes 2004). These particular areas are thus intensely metabolically active, and are likely to contain numerous ribosomes, targets of the oligonucleotide probes. A number of studies using FISH, especially on single-cell organisms, report that metabolic condition has a strong influence on the intensity of signal from rRNA-targeting oligonucleotide probes (see Amann et al. 1995 for review). This might explain a predominant signal in the most metabolically active cells of the larva.

False negatives were seldom encountered, and were restricted to Mytilus edulis larvae hybridized with the Myt-18S and eukaryotic (EUK516) probes. Problems of target accessibility have been reported in FISH applications on microorganisms; these arise from cell wall prevention of probe penetration into the cytoplasm (Amann et al. 1995). In order to avoid limited target accessibility caused by sealed mollusc shells, we used a solution of MgCl₂ before fixing the larvae, allowing adductor muscles, responsible for closing of the valves, to relax, leaving the shell open for the subsequent hybridization procedure. Meanwhile, shell morphology was kept intact for subsequent visual inspection. The effect of the reagent could be checked on larvae with a protruding velum from between the valves. As false negative cases always corresponded to larvae in which the velum was retracted and the valves closed, a problem might have occurred during larval fixation. An alternative hypothesis is that some of the larvae may have been in very poor metabolic condition, and that their unusually low cellular ribosome content resulted in a signal too weak to be detected.

Relevance of the 18S rRNA-targeted probes for species identification via in situ hybridization

Probes targeting rRNAs have been widely used for identifying and quantifying a range of organisms,
The availability of both conserved and variable regions, allowing the design of universal primers for subsequent amplification of variable regions of the 18S gene, make it a widely used marker for species identification purposes, such as DNA barcoding projects (e.g. Floyd et al. 2002, Bhadury et al. 2006). Finally, the existence of databases of ribosomal sequences (e.g. The European Ribosomal RNA Database: www.psb.ugent.be/rRNA) facilitates the assembly of homologous sequence databases required for the design of specific probes.

Potential applications of the method

Many aspects of the larval stages of marine benthic invertebrates remain poorly understood. The planktonic stage is nevertheless a crucial parameter for consideration at the population and ecosystem level, shaping genetic structures and determining the evolution of populations and eventually communities. Processes governing larval dispersal and recruitment and their relative importance and interactions are not yet fully elucidated. Physical, chemical and ecological processes have been reported as important parameters, together with active larval behaviour at small spatial scales. Studies of larval distributions have so far been hampered by the difficulty of accurately identifying larvae, especially those that are small. Unambiguous, straightforward and high-throughput identification methods are needed to meet the extreme requirements of field-based studies relying on intensive sampling programmes over long periods of time. This is a prerequisite towards the elucidation of larval migration patterns, recruitment processes, and predator/prey population dynamics. In situ hybridization on whole larvae could be used in multidisciplinary studies, providing new insights into population connectivity over a range of scales (e.g. Gilg & Hilbish 2003, Arnold et al. 2005). The method can be easily implemented in any biological laboratory; it does not need costly equipment or consumables and does not require staff to go through advanced specific training. The methodology is flexible. Several steps can be automated, and data acquisition can be made faster through a motorised scanning microscope stage and image analysis technology.

More specific applications include long- or short-term monitoring of multiple or single target species, such as species of commercial interest, bio-indicator, rare or invasive species. Routine screening of plankton samples for biodiversity studies will require the generation of sub-samples to be tested with different probes. Sufficiently large samples would be necessary for subsequent testing of the statistical significance of the results. Alternatively, different probes labelled with different fluorescent dyes could be used simultaneously on wild samples, as is the case for microbial studies (Wallner et al. 1993). Such an approach, increasing the throughput of the method, would, however, require significant protocol optimization. Nested probes, designed for different taxonomic levels and applied to sub-samples of plankton in an ordered top-to-bottom approach, could offer another alternative approach, as previously implemented through FISH with rRNA targeted oligo-probes on microbial samples (Amann et al. 1995). The results obtained with high-level probes allow the selection of probes with narrower specificity designed for the next lowest taxonomic level, providing increasingly refined information on community diversity and composition.

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