Mineral phase in shell repair of Manila clam Venerupis philippinarum affected by brown ring disease

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ABSTRACT: The mineral phase of shell repair in the Manila clam Venerupis philippinarum affected by brown ring disease (BRD) was characterised at various scales and at various stages of shell repair by confocal Raman microspectrometry and scanning electron microscopy. Spherulitic and quadrangular aragonite microstructures associated with polyene pigments were clearly observed. Von Kossa staining showed that at the beginning of shell repair, hemocytes are filled with insoluble calcium carbonate salts in all fluids and then are transported toward the extrapallial fluids and the repair sites. Our analyses suggest that after a Vibrio tapetis attack and BRD deposit some clams rapidly cover the deposit, resulting in a modification in the microstructure, which could be produced by the participation of both the mantle and hemocytes.

KEY WORDS: Aragonite · Hemocytes · Shell microstructure · Calcium carbonate · Biomineralization · Mollusc · Raman microspectrometry · Scanning electron microscopy

INTRODUCTION

In marine biota, invertebrates are constantly exposed to external environmental insults. These insults can be of physicochemical nature, such as desiccation, exposure to UV light or accidental cracks, or of biological nature, such as predation or competition for space. During evolution, a majority of sedentary invertebrates have acquired the capacity to develop a mineralised exoskeleton to protect themselves against such occurrences. In addition, their exoskeleton supports their soft bodies (Simkiss & Wilbur 1989). In molluscs, the protective exoskeleton is the shell, which is mainly composed of calcium carbonate. The calcium carbonate layers of molluscan shells can be formed by aragonitic crystals such as in the clam Mercenaria mercenaria (Taylor et al. 1973), by calcitic crystals such as in the oyster Crassostrea virginica (Carriker 1996) or by a mix of the 2 crystal polymorphs such as in the abalone Haliotis rufescens (Nakahara et al. 1982). In exceptional environmental conditions the highly unstable vaterite can also be formed (Spann et al. 2010).

The Manila clam Venerupis philippinarum is a heterodont bivalve of economic importance. Its valves are constructed by the superimposition of 2 aragonitic layers (Taylor et al. 1973). The inner layer is described as ‘homogeneous’ while the outer layer is a ‘composite prismatic’ structure (Boggild 1930, Shimamoto 1986). As in other Veneridae, the attachment of the mantle to the shell along the pallial line separates the extrapallial space into 2 compartments, which contain the central and the peripheral extrapallial fluids (Allam & Paillard 1998, Gosling 2003).

In Europe, this clam is regularly affected by a pathology named brown ring disease (BRD). This disease, which is characterised by the presence of a brown deposit on the inner surface of the valves, is caused by the bacterium Vibrio tapetis (Paillard & Maes 1990). This disease was first recognized in 1987 when mass mortalities of cultured Manila clams were recorded in Landeda, north Finistère (Brittany, France) (Paillard et

Vibrio tapetis colonises the periostracum and inhibits the normal process of shell biomineralisation. The external response of the clam to the bacterial attack consists of a secretion of a brown organic matrix on the inner face of the shell, the well-known ‘brown ring’. Immunological or PCR studies on the different physiological fluids of the clam have shown that, at later stages of the infection, V. tapetis invades the circulatory system (hemolymph) and the extrapallial fluids (Allam 1998, Allam et al. 2002, Paillard et al. 2006). At this advanced stage, clams often die from a hemolymphatic septicaemia. In some cases, clams recover by secreting a white shell layer on the brown organic matrix (Paillard & Maes 1995a,b). The bulk organic matrix associated with the shell repair has been recently characterised using Raman microspectrometry, wavelength-dispersive spectrometry (WDS) microprobe analysis and organic matrix quantification (Trinkler et al. 2009, 2010a). However, the mineral phase and the microstructure of repaired Manila clam shells, including those that exhibit a fully repaired shell, have not yet been studied. In the present study, experimental investigations have been performed on the microstructure and the nature of the calcium carbonate repair layer, using Raman microspectrometry and scanning electron microscopy (SEM). In particular, Raman microspectrometry, a nondestructive and noninvasive technique, provides qualitative and quantitative information on organomineral composites (Urmos et al. 1991, Kontoyannis & Vagenas 2000, Withnall et al. 2003, Hedegaard et al. 2006, Carteret et al. 2009). A complementary characterisation using the Von Kossa stain, which is a stain specific for insoluble calcium salts, was performed on hemolymph and extrapallial fluids, revealing the presence of these salts in hemocytes. These techniques applied to BRD-affected and nonBRD-affected clams provide valuable information on the mineral phase.

MATERIALS AND METHODS

Origin of clams and BRD characterization. Adult Manila clams (25 to 35 mm) used for Raman microspectrometry and Von Kossa stain were collected in field at the Landeda site on 23 April 2007. Most of them exhibited a brown ring (Paillard & Maes 1994). Juvenile clam specimens (10 to 16 mm) used for Raman microspectrometry and SEM were provided by the Satmar nursery, Marennes, France, on 3 May 2007 (Table 1). Shells were systematically opened, emptied of soft tissue by means of a scalpel and rinsed with tap water before determining the stage of BRD on the inner face of the valves using binocular stereomicroscopy. The shell repair stages (SRS) were determined according to previous studies, which identified 5 stages (Paillard & Maes 1994, Paillard 2004). Stage 0 corresponds to a brown deposit without repair. Stage 1 corresponds to the apposition of calcified points. The partial covering of the deposit by wide calcified plates corresponds to Stage 2. Stage 2.5 corresponds to an almost complete repair. In Stage 3, the brown deposit is completely covered by a fully calcified layer (Fig. 1).

Raman microspectrometry. Raman microspectrometer: We used a Raman microspectrometer (T64000, Horiba-Jobin-Yvon) equipped with a confocal system, a motorized x–y stage and a coherent spectrum argon/krypton ion laser emitting at 51435 nm. The laser was focused onto the sample using a microscope equipped with a 100× objective; the resulting spatial resolution was about 1 µm. The scattered light was then analysed using a single monochromator (600 gratings mm⁻¹), coupled to a nitrogen-cooled charge-coupled device (CCD) de-

<table>
<thead>
<tr>
<th>Sample</th>
<th>Origin</th>
<th>Injected with Vibrio tapetis?</th>
<th>BRD-affected?</th>
<th>Size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Raman microspectrometry</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 adults (A1, A2)</td>
<td>Landeda</td>
<td>No</td>
<td>NonBRD-affected</td>
<td>25–35</td>
</tr>
<tr>
<td>5 juveniles (J1, J2, J3)</td>
<td>Satmar Nursery</td>
<td>No</td>
<td>NonBRD-affected</td>
<td>10–16</td>
</tr>
<tr>
<td>12 adults</td>
<td>Landeda</td>
<td>No</td>
<td>BRD-affected: 2 SRS 0, 2 SRS 1, 1 SRS 2, 7 SRS 3</td>
<td>25–35</td>
</tr>
<tr>
<td><strong>SEM</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 juveniles</td>
<td>Satmar Nursery</td>
<td>No</td>
<td>NonBRD-affected</td>
<td>10–16</td>
</tr>
<tr>
<td>10 juveniles</td>
<td>Satmar Nursery</td>
<td>Yes</td>
<td>BRD-affected: 2 SRS 0, 2 SRS 1, 2 SRS 2, 2 SRS 2.5, 2 SRS 3</td>
<td>10–16</td>
</tr>
<tr>
<td><strong>Von Kossa stain</strong></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>9 adults</td>
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<td>No</td>
<td>NonBRD-affected</td>
<td>25–35</td>
</tr>
<tr>
<td>21 adults</td>
<td>Landeda</td>
<td>No</td>
<td>BRD-affected: 1 SRS 0, 1 SRS 1, 14 SRS 2, 2 SRS 2.5, 3 SRS 3</td>
<td>25–35</td>
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</table>
For each sample, parameters were adjusted as follows: laser power (varying from 30 to 50 mW), spectra accumulation (from 2 to 30 times) and exposure time (from 1 to 60 s) (Table 2). Measurements were systematically made at 2 µm depth and sometimes on fresh breaks to avoid surface contamination. Reported Raman spectra were normalised by dividing intensity by acquisition time (s) and laser power (mW). The peak positions and the relative integrated intensities of identifiable vibrational bands and luminescence were determined using the LABSPE software.

**Samples analyzed using Raman microspectrometry:** Raman analysis was first carried out on the internal surface of a nonBRD-affected adult (A1) in 5 regions of the right valve: central zone, peripheral anterior zone, peripheral posterior zone, peripheral ventral zone and siphonal zone. Raman spectra were also acquired on the internal surface of 4 nonBRD-affected juveniles (Table 1). Similar results were obtained with the 4 shells so only 2 of them are presented (see Fig. 4; J1, left valve, peripheral ventral zone; J2, left valve, central zone). In addition, we compared the spectra obtained from the inner and the outer shell layers from 2 nonBRD-affected shells (one adult, A2, and one juvenile, J3) after having cut both of them with cutting pliers to avoid aragonite/calcite thermal transformation (Table 1).

Subsequently, we studied adult clams affected by BRD. For each SRS, analyses were made on the shell surface to characterise the evolution of the Raman signal.

**Table 2.** *Venerupis philippinarum*. Samples and parameters used for Raman microspectrometry analysis. A: adult; J: juvenile

<table>
<thead>
<tr>
<th>Sample</th>
<th>Analysis</th>
<th>Parameter</th>
<th>Acquisition time (s)</th>
<th>Laser power (mW)</th>
<th>Accumulation (times)</th>
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<tbody>
<tr>
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<td>Surface analysis</td>
<td></td>
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<td>2–4</td>
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<tr>
<td>J1</td>
<td></td>
<td></td>
<td>35</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>J2</td>
<td></td>
<td></td>
<td>50</td>
<td>30</td>
<td>2</td>
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<td>10</td>
</tr>
<tr>
<td>2 clams SRS 1</td>
<td></td>
<td></td>
<td>1</td>
<td>30</td>
<td>10–30</td>
</tr>
<tr>
<td>1 clam SRS 2</td>
<td></td>
<td></td>
<td>2</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>2 clams SRS 3</td>
<td></td>
<td></td>
<td>4</td>
<td>30</td>
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<td>A2</td>
<td>Edge analysis</td>
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<td>40</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>J3</td>
<td></td>
<td></td>
<td>30–40</td>
<td>50</td>
<td>2</td>
</tr>
<tr>
<td>1 nonBRD-affected adult</td>
<td>Polyene analysis</td>
<td></td>
<td>30–60</td>
<td>50</td>
<td>2</td>
</tr>
<tr>
<td>5 SRS 3 adults</td>
<td></td>
<td></td>
<td>2–50</td>
<td>30–40</td>
<td>2–10</td>
</tr>
</tbody>
</table>
during the shell repair (2 clams with SRS 0, 2 with SRS 1, 1 with SRS 2 and 2 with SRS 3; Table 1).

**SEM analysis. Inoculation:** *Vibrio tapetis* CECT 4600, isolated from *Venerupis philippinarum* in Landeda (Paillard & Maes 1990), was grown in marine agar (1 l distilled water, 15 g pastag [AES Chemunex], 30 g sea salts [Sigma], 4 g peptone [AES Chemunex], 1 g yeast extract [AES Chemunex] and 0.1 g Fe(PO₄)₂ [Merck]) at 18°C for 72 h before inoculation. Then, *V. tapetis* was re-suspended in sterile sea water (SSW).

Two hundred juvenile clams, obtained from a low susceptibility stock at the Satmar farm (Trinkler et al. 2010b), were injected in the pallial cavity with 0.05 ml of SSW and 200 other clams (same stock) with 0.05 ml of a solution containing *Vibrio tapetis* at 2 × 10⁸ cells ml⁻¹ as previously described (Paillard & Maes 1990). The clams were maintained in marine aquaria at 14°C. A low susceptibility stock of clams was chosen for their capacity to repair their shell (a higher SRS) (Trinkler et al. 2010b). After 6 wk, clams were sampled and their SRS was characterised (Fig. 1).

**Sample preparation for SEM:** Using a QUANTA 200 SEM (FEI), observations were made on the inner valve surface of both nonBRD-affected shells that were not injected and experimentally diseased shells. The outer layer was visible on the edge of the shell. BRD-affected and nonBRD-affected zones were cut out with a scalpel. Fragments from 2 shells of nonBRD-affected clams and from 2 shells of each SRS (0, 1, 2, 2.5 and 3) were soaked in 70% ethanol for 2 h, rinsed with distilled water and dried at 37°C (Table 1). Then, non BRD-affected and repaired fragments were soaked in sodium hypochlorite solution (0.027% active chlorine for 2 h) to remove the organic phase and isolate the mineral phase. As the brown deposit is entirely organic (Trinkler et al. 2010a,b), clams at SRS 0 were not soaked in sodium hypochlorite solution. The fragments were rinsed with distilled water, dried at 37°C and glued onto SEM stubs with conductive silver paint and coated with gold, using a cathodic spray SCD 040 (Blazers Union).

**Von Kossa staining:** The Von Kossa method stains insoluble salts of calcium phosphate and calcium carbonate. The staining was performed according to the following standard procedure (Gabe 1968): 10 min incubation in 5% (w/v) of silver nitrate in water in the dark, 2 × 10 min washes with distilled water to remove the excess silver nitrate, staining with 0.5% (w/v) of hydroquinone in water for 2 min in daylight, a wash with distilled water (10 min), an incubation with sodium thiosulfate (5%, w/v) in water for 30 s, a wash with distilled water (10 min) and a brief immersion (15 s) in aniline blue (3% w/v) in water. Samples were dehydrated (10 min in 70% ethanol, 10 min in 95% ethanol and 2 × 10 min in 100% ethanol) and mounted with a mounting medium (Eukitt). The CEF of a clam at SRS 3 was also soaked in acetic acid (0.1 M, 10 min) to evaluate the specificity of the stain (salts of calcium phosphate and calcium carbonate are dissolved with acetic acid). Then all the slices were observed and photographed with an inverted optical microscope (DM IRB, Leica). The intensity of the staining was visually assessed.

**RESULTS**

**Raman micro-spectrometry results**

NonBRD-affected clams

Raman spectra measured on the inner layer of different valve zones of a nonBRD-affected adult clam were compared (Fig. 2). In the 155 to 1460 cm⁻¹ wave number range, the Raman spectra measured on the surface of the nonBRD-affected adult clam (A1) were similar in all zones of the valve, and the strong bands observed at 155, 205, 271, 704, 1087 and 1460 cm⁻¹ were assigned to the aragonite phase. In the 1018 to 4071 cm⁻¹ wave number range, well-defined bands appeared in the posterior and siphon zones (Table 3). Two particular bands, named R1 (around 1100 to 1140 cm⁻¹) and R4 (around 1500 to 1525 cm⁻¹) were attributed to carbon–carbon single and double bond stretching vibrations, respectively. The observation of these bands revealed the presence of polyene pigments (Table 3, Fig. 3).
Raman spectra of the inner layers of non-BRD-affected adult and juvenile clam shells are shown in Fig. 4. Raman spectra acquired on the surface of non-BRD-affected juvenile clams (J1 and J2) showed the presence of aragonite and polyenes bands and a strong luminescence. In the juvenile spectra (J1, J2), the luminescence intensity was systematically higher than in the adult (A1). In juvenile clam shells, the series of polyene bands were also more difficult to observe because the peaks were of much lower intensity (which is related to the concentration of pigment) and/or were masked by the luminescence of the sample.

Finally, Raman spectra measured on the inner and outer layers of non-BRD-affected adult and juvenile clam shells are compared in Fig. 5. Raman spectra acquired on the edge of a juvenile (J3) and of an adult (A2) specimen, showed well-defined vibrational bands attributed to aragonite and polyenes. Furthermore, the adult outer layer exhibited higher luminescence intensity than did its inner layer. The juvenile outer layer presented lower luminescence intensity than did its inner counterpart (Fig. 5). The luminescence intensity observed in the juvenile inner layer was higher than in the adult inner layer. Finally, the luminescence intensities of the juvenile and adult clam outer layers were similar (J3 and A2, Fig. 5).

BRD-affected clams

Raman spectra acquired on the brown deposits at different SRS are shown in Fig. 6. At SRS 0, before repair, a high-luminescence signal was observed. During the process of shell repair (from SRS 1 to SRS 3), the main aragonite bands (usually observed between 158 and 1087 cm\(^{-1}\)) became more prominent. At the end of the repair process (SRS 3), polyene bands also reappeared in the wave number range 1037 to 4091 cm\(^{-1}\).

Thus, the aragonite bands were always observed in non-BRD-affected adult and juvenile shells as well as in repaired shell and not in organic brown rings. Similarly, the polyene bands were observed in non-BRD-affected and repaired clams but never in the brown rings (Fig. 6).

SEM results

Non-BRD-affected clams

The internal shell surfaces of non-BRD-affected clams were observed by SEM. The inner layer was composed of organised, flattened and imbricate crystals embedded in a thin organic layer (Fig. 7a). After a surface treatment with sodium hypochlorite, the thin organic
layer disappeared (Fig. 7b). In the case of the outer layer (visible at the edge of the shell), the calcium crystals formed mounds, like prisms, in surface view (Fig. 7c).

BRD-affected clams

At the beginning of the BRD process, the brown deposits appeared on the internal shell surface. SEM observations showed that they consisted of a thick organic layer, laid down on the inner shell layer (Fig. 7d). From SRS 1 to SRS 2.5, SEM results showed that calcium carbonate minerals were progressively settled on the brown deposit until the affected zone was almost entirely covered (Fig. 7e–o). At SRS 1, in shells not treated with sodium hypochlorite, minerals seemed to be partially masked under a thick organic matrix, which looked like the BRD organic matrix (Fig. 7e). After sodium hypochlorite treatment, we noticed a great variability of microstructures in SRS 1, from rounded ‘spherulitic-like’ (Fig. 7f) to rectangular (Fig. 7g). In treated fragments, the rectangular minerals were surrounded by a thin organic matrix, and as the sodium hypochlorite treatment degraded this matrix it formed holes (Fig. 7g). During SRS 2, 2.5 and 3, with treatment, spherulitic-like (Fig. 7h,i,l) and rectangular (Fig. 7o) microstructures were also observed. From SRS 1 to SRS 2.5, calcium carbonate minerals were not well organized.

At SRS 2.5, an organic layer appeared and partially covered the calcium carbonate crystal mounds (Fig. 7j,k). At the end of the shell repair (SRS 3), this organic layer entirely covered the crystals (Fig. 7m). It was thick, bright and smooth (Fig. 7j,k,m) and appeared different from the organic matrix of the non-BRD-affected inner layer and from the brown deposit (Fig. 7d). Under the BRD-organic layer, calcium carbonate minerals exhibited an organisation similar to that observed from SRS 1 to SRS 2.5, in particular, rectangular structures were evident (Fig. 7g,o).

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Table 3. *Venerus philippinarum*. Wave numbers of pigment bands on clam shell (cm\(^{-1}\)). R1 designates wave numbers for carbon–carbon single bond stretching vibration and R4 designates wave numbers for carbon–carbon double bond stretching vibration. H: nonBRD-affected; SRS 3: total shell repair

<table>
<thead>
<tr>
<th>Clam status</th>
<th>Example plot</th>
<th>R1</th>
<th>R4</th>
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<tr>
<td><strong>Inner layer</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H Adult</td>
<td>Fig. 2 A1(^a)</td>
<td>1122</td>
<td>1500</td>
</tr>
<tr>
<td>H Juvenile</td>
<td>Fig. 4 J1</td>
<td>1135</td>
<td>1527</td>
</tr>
<tr>
<td>H Juvenile</td>
<td>Fig. 4 J2</td>
<td>1136</td>
<td>1530</td>
</tr>
<tr>
<td>H Adult</td>
<td>Fig. 5 A2</td>
<td>1139</td>
<td>1530</td>
</tr>
<tr>
<td>H Juvenile</td>
<td>Fig. 5 J3</td>
<td>1121</td>
<td>1501</td>
</tr>
<tr>
<td>H Adult</td>
<td></td>
<td>1136</td>
<td>1529</td>
</tr>
<tr>
<td>Adult SRS3</td>
<td>Fig. 6</td>
<td>1126</td>
<td>1505</td>
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<td><strong>Outer layer</strong></td>
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<td>Fig. 5 A2</td>
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<tr>
<td>Juvenile</td>
<td>Fig. 5 J3</td>
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<td>1497</td>
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</table>

\(^a\)Other pigment bands for A1: 1018, 1122, 1166, 1291, 1500, 2130, 2241, 2411, 2502, 2609, 2787, 2975, 3009, 3311, 3681 and 4071 cm\(^{-1}\)

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Fig. 3. *Venerus philippinarum*. R1: wave numbers (cm\(^{-1}\)) (1/wavelength) of carbon–carbon single bond stretching vibration plotted against R4: wave numbers of carbon–carbon double bond stretching vibration. ×: *V. philippinarum* samples reported in Table 3; □: predicted Raman shift of resonantly coupled modes for polynenes with different chain lengths reported by Hedegaard et al. (2006)

Fig. 4. *Venerus philippinarum*. Raman spectra acquired on surface of inner layer of nonBRD-affected clam shells of one adult A1 (ventral zone, dark spectrum) and 2 juveniles J1 (ventral zone, grey spectrum) and J2 (central zone, dark grey spectrum). Baselines are vertically shifted to facilitate comparison. Wavenumber (cm\(^{-1}\)): 1/wavelength; a.u.: arbitrary unit (intensity normalised by dividing intensity by acquisition time [s] and the laser power [mW]). The dome-like curves (juveniles) correspond to the intrinsic luminescence
Fig. 5. *Venerupis philippinarum*. Raman spectra acquired on the edge of nonBRD-affected clam shells of one adult A2 and one juvenile J3 to measure the signatures of inner and outer layers. Baselines are shifted vertically to facilitate comparison. Note that the juveniles exhibit high luminescence. Wavenumber (cm\(^{-1}\)): 1/wavelength; a.u.: arbitrary unit (intensity normalised by dividing intensity by acquisition time [s] and the laser power [mW]).

Fig. 6. *Venerupis philippinarum*. Raman spectra acquired on the surface of 7 samples of Manila clams, with different shell repair stages (SRS). SRS 0: disease without repair; SRS 1: calcified points; SRS 2: calcified plates; SRS 3: total shell repair. Baselines are not shifted vertically. Wavenumber (cm\(^{-1}\)): 1/wavelength; a.u.: arbitrary unit (intensity normalised by dividing intensity by acquisition time [s] and the laser power [mW]).
Fig. 7. *Venerupis philippinarum*. SEM views of the inner surface of Manila clam shells. NonBRD-affected and experimentally diseased shells were observed to describe the development of the microstructure during the shell repair mechanism. (a,b) Non-BRD-affected (N) inner layer. (c) NonBRD-affected outer layer. (d) Brown deposit without shell repair, i.e. shell repair stage (SRS) 0. (e–g) SRS 1. (h,i) SRS 2. (j–l) SRS 2.5. (m–o) SRS 3. Shells in (7b,l,g,i,l,n,o) were soaked in sodium hypochlorite solution (0.027% of active chlorine) for 2 h. Arrows indicate organic matrices.
Von Kossa staining

In all clams, some hemocytes on each glass slide were coloured with Von Kossa stain whatever the conditions and the compartments, but the proportion of stained hemocytes and the intensity of the brown staining varied depending on the nature of the sampled fluid (hemolymph, CEF, PEF) and on the repair stage (Fig. 8, Table 4). The brown intensity of the staining and the proportion of stained hemocytes on each slide were visually assessed. In each case, the localisation of the staining in hemocytes seemed concentrated in vesicles and more diffused in cytoplasm. The results are summarised in Table 4. In nonBRD-affected clams, the proportion of stained hemocytes was lower in hemolymph than in both extrapallial fluids, but the intensity of the staining was lower in the CEF than in the hemolymph and in the PEF (Fig. 8a–c, Table 4). At SRS 1, the proportion of stained hemocytes increased in hemolymph and the brown intensity increased in CEF (Fig. 8d–f, Table 4). At SRS 2, the proportion of stained hemocytes and the brown intensity drastically decreased in hemolymph. At the same time, the proportion of stained hemocytes increased in both extrapallial fluids, similarly to the staining intensity (Fig. 8g–i, Table 4). At SRS 2.5, the proportion of stained hemocytes and the brown intensity increased in the hemolymph and reached a higher level than that recorded for nonBRD-affected hemolymph. At the same time, the brown intensity decreased slightly in both extrapallial fluids (Fig. 8j–l, Table 4). During this stage, the proportion of stained hemocytes and the brown intensity were high or very high in hemolymph and both extrapallial fluids (Fig. 8j–l, Table 4). At the end of the shell repair (SRS 3), the situation was similar to SRS 2.5, with the exception that the proportion of stained hemocytes and the brown intensity decreased slightly in hemolymph, where it was moderate and intermediate, respectively, similar to the level found in nonBRD-affected hemolymph (Fig. 8m–o, Table 4).

At stage SRS 3, a glass slide of CEF rich in brown stains was soaked in acetic acid to induce the disappearance of the brown Von Kossa stain. (Fig. 8p,q). This strongly suggests that the staining was specific for insoluble calcium salts. The remaining localised dark dots are probably aggregates of uric acid or urate.

DISCUSSION

In the present study, we analysed the shell repair process of the Manila clam by using 3 different analytical tools: confocal Raman microspectrometry, SEM and staining of hemocytes for insoluble calcium salts. Raman microspectrometry confirmed that nonBRD-affected shells were fully made of aragonite, the metastable polymorph of calcium carbonate (Kontoyannis & Vagenas 2000, Carteret et al. 2009), which is not surprising: earlier studies revealed that the 2 layers that make up the Manila clam shell, i.e. the outer composite prismatic layer and the inner homogeneous one, are aragonitic (Taylor et al. 1973). More interesting is that aragonite — and only aragonite — was found also in the repair zone, above the brown ring, which is fully organic and does not contain any mineral. Nor did we detect traces of vaterite or calcite. Earlier publications on molluscan shell repair indicate that, in some particular cases, aragonite was sometimes replaced by calcite or by vaterite (Muzii et al. 1966, Saleuddin & Wilbur 1969, Saleuddin 1971, Meenakshi et al. 1975, Blackwelder & Watabe 1977, Suzuki 1979). Another contribution of Raman microspectrometry was the identification of a polyene pigment signature, characterised by 16 bands from 1018 to 4071 cm⁻¹ (Hedegaard et al. 2006). Polyenes were found in both layers of nonBRD-affected shells, as well as in the repaired zone, but they were virtually absent from the brown ring deposit or their presence was masked by the high luminescence of that layer. Polyenes are polyunsaturated long carbon chains of the alkene type, i.e. they exhibit 8 to 13 sequences of alternating double and single carbon—carbon bonds (16 to 26 carbon atoms) with terminal –CH₃ groups (Hedegaard et al. 2006). To our knowledge, this is the first report of the presence of polyenes in the shell of the Manila clam. In earlier work, carotenoids were identified in molluscan shells with Raman spectrometry (Zhang et al. 2001, Withnall et al. 2003, Brink & Van Der Berg 2005), and Hedegaard et al. (2006) mentioned the presence of polyenes in molluscan shells. These polyenes are the pigments responsible for the shell colour such as purple, brown or yellow of the inner and outer layers of clams. Another contribution of the Raman analysis is the detection of an intense luminescence, particularly in the outer layers of adults and juveniles, and the inner layers of juveniles. This luminescence may be due to several organic compounds; those carrying chromophores or fluorescent groups (Bell et al. 2002), organics with amide I groups (Schachar & Solin 1975) or cellulose and lignins in Permian fossil wood (Witke et al. 2004). In bivalves, a high luminescence was recorded in the shell nacre nucleus of Pinctada margaritifera (Rousseau et al. 2005). In the outer layer, the luminescence might be caused by the presence of the pigmentation or by the occurrence of organic components, which are specific to the composite prismatic layer. The last point refers to the high intensity of the luminescence in the inner layer of juvenile specimens in comparison with adults. This may suggest that the ratio of organic matrix to calcium carbonate would be higher in juvenile shells than in those of adults.
**Fig. 8.** *Venerupis philippinarum*. Examples of clam hemocytes coloured with Von Kossa stain, observed by optical microscopy, 400×. The insoluble calcium carbonate salts are dark (brown) coloured. Hemocytes from different fluids of nonBRD-affected and experimentally diseased clams were coloured to show the transport of insoluble calcium carbonate salts during the shell repair mechanism. (a–c) One nonBRD-affected sample. (d–f) One sample at SRS 1. (g–i) One sample at SRS 2. (j–l) One sample at SRS 2.5. (m–o) Two samples at SRS 3. (p,q) Two samples at SRS 3, previously soaked in (p) water or (q) acetic acid. H: hemolymph; CEF: central extrapallial fluid; PEF: peripheral extrapallial fluid.
repairs of the composite prismatic layer, have been observed in shell polygonal units of crystals, such as those in the composite prismatic layer when observed in transverse view. In the repair zone, the microstructures present a greater variability. The formed crystals of the repair layer exhibited 3 shapes: rounded spherulitic, quadrangular and elongated. Spherulits and/or the repair layer formed ‘mounds’. This structure was typical of the outer layer do not have a consistent function during the repair process. Another hypothesis postulates that the mantle responsible for the mineral deposition in the repair zone is moving, i.e. it sometimes withdraws from around the repair zone. A similar phenomenon has been recently observed in the abalone *Haliotis tuberculata* (Fleury et al. 2008). A complex functioning of the mantle can also be inferred from the analyses of the organic matrix. Previous publications (Trinkler et al. 2009, 2010a) showed that the matrix associated with the repair zone was more abundant than the one associated with the homogeneous, nonBRD-affected layer. The ‘repair’ matrix contains luminescent and sulfated components.

The present study supports the suggestion that hemocytes play a key role during the shell repair process. We particularly emphasise their active contribution to the transport of insoluble calcium carbonate salts to the site of mineralisation. Our study corroborates that of Allam (1998), performed on the same species, in which holes were drilled in Manila clam shells and sealed with pieces of sterile coverslip glass. A few days after inoculation of *Vibrio tapetis* in the tested specimens, hemocytes were detected on the coverslips.

If hemocytes are unquestionably involved in cellular responses following shell alteration, they also may have multiple functions, depending on the mollusc model and on the characteristics of the shell injury. For example, in the freshwater snail *Pomacea paludosa*, 3 d after shell removal, hemocytes were detected in the extrapallial space (Watabe & Blackwelder 1980). They eventually aggregated with other cell fragments to form a cellular barrier, which contributed to the deposition of a new mineral phase 7 d.
after shell removal. During this shell repair process, calcium was mostly localised in the intercellular space and hemocytes did not show a higher calcium concentration than that for epithelial cells. The edible mussel _Mytilus edulis_ represents a second example of another role played by hemocytes: after periostracum damage, hemocytes migrated through the epithelium to take part in the production of the regenerated periostracum (Dunachie 1963). In the freshwater bivalve _Anodonta grandis_, after shell damage, hemocytes accumulated in the thickness of the altered mantle, where they were presumably responsible for regenerating the injured part of the mantle (Saleuddin 1967). Finally, in _Crassostrea virginica_ affected by juvenile oyster disease (JOD), a disease characterised by a brown deposit of conchiolin on the inner layer of valves similar to BRD, hemocytes were observed on the conchiolin deposit and in the altered mantle by histology (Bricelj et al. 1992, Ford & Borroco 2001). These 4 examples clearly show that hemocytes, although similarly localised in the subepithelial region of the repairing mantle, play different roles in addition to the transport of the mineral material.

In the present study, we were not able to demonstrate whether the calcium carbonate particles transported by hemocytes were amorphous or crystalline. This question is currently debated. Previous work has found support for the existence of calcium carbonate granules in calcifying molluscan mantle tissues (Robel et al. 1973), either intracellularly (Fournié & Chétail 1982) or extracellularly (Istin & Masoni 1973). In most cases, although no precise studies were done on their structure, these granules were assumed to be amorphous; because they have a greater solubility than their crystalline counterparts, amorphous calcium carbonate minerals represent a source of calcium which is rapidly available and easily remodelled. However, one recent example diverges from this pattern: in the eastern oyster _Crassostrea virginica_, Mount et al. (2004) demonstrated that refractive granulocytes, a subclass of hemocytes, contained calcium carbonate in a crystalline form. A combination of SEM observations and Raman spectrometry measurements showed that these crystals appeared with sharply defined edges and were calcitic. These crystals were directly brought and released at the mineralisation site by granulocytes. So far, it is not known whether the _Crassostrea virginica_ model is the exception or the rule (Mount et al. 2004).

Numerous studies have described how the total hemocyte counts in Manila clam hemolymph increase during the first days of BRD infection, then decrease, whereas the total hemocyte count in extrapallial fluids always increases. This trend was explained as a mobilisation of hemocytes to the extrapallial fluids (Allam et al. 2000a,b). Hemocytes are also observed in the Manila clam mantle, which is altered by _Vibrio tapetis_ during a BRD infection (Allam et al. 2002). However, none of these studies examined the link with shell repair. One study also examined the correlation between hemocytes and resistance to BRD, but did not link it to shell repair: Allam et al. (2001) observed that different populations of _Venerupis philippinarum_ (from France and the USA) and _Ruditapes decussatus_ (from France) did not exhibit the same capability to overcome BRD; greater resistance was associated with a higher concentration of granular hemocytes and greater phagocytic activity.

If these observations are confirmed, we can then seriously envisage the selection of BRD-resistant stocks of Manila clams by sampling their hemocytes. However, additional work on the cell typology of the Manila clam immune system, combined with the identification of molecular markers of hemocyte types using a proteomics approach, appear to be an essential prerequisite.

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