Structural and biological characterization of two freshwater mussel shells (Bivalvia: Unionidae)

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Abstract: The shell of a freshwater mussel (Mollusca: Bivalvia) is a composite biological material that plays an active role in maintaining ecosystem services. Mussel shells have variable structures both within and between species. In this study, the structural and biological characteristics of the shells of Potomida semirugata and Leguminaia wheatleyi were investigated in order to use freshwater mussel shells as a biological material. Close observation of the microstructure of the shells of the two common freshwater bivalves revealed a variation in construction from the outer periostracum to the inner nacreous layer. In P. semirugata, a polygonal arrangement with columnar prisms of different sizes was observed more significantly. In L. wheatleyi the prismatic columns are oval and irregularly polygonal. The nacreous layers of P. semirugata, in which the individual aragonite layers are horizontally overlapped, are more uniformly distributed. The overlapping of the aragonite sheets in L. wheatleyi is more irregular. The presence of calcium carbonate (CaCO₃) in the shells was confirmed by the identification of the characteristic carbonate bands at 701.9, 713.5, and 865.8 cm⁻¹. This information was used to classify and distinguish the different layers of the shells, allowing interspecific comparisons and variations in the different layers. The results clearly showed that shell samples were highly biocompatible and nontoxic compared to the control group. This finding suggests that these materials have promising potential for use in various biomedical applications where biocompatibility is a critical factor. The results indicated that the shells of the freshwater bivalves P. semirugata and L. wheatleyi are biological materials with potential multiple applications for human well-being and environmental quality.

Key words: Calcification, freshwater bivalve, biomineralization, biological material, nacre, scanning electron microscope

1. Introduction

Freshwater mussels have a unique shell structure that is hinged together to protect and support the body (Carter, 1990). Two symmetrical calcareous valves are organomineral composites, and these minerals are generally composed of calcium carbonate (CaCO₃) in the form of calcite and/or aragonite. CaCO₃ is the component that makes up the majority of bivalve shells, but traces of inorganic elements are also present. Unionid shells are characterized by an outer periostracum, an aragonitic prismatic layer, and an inner nacreous layer (Jackson et al., 1988). Due to their different microstructures, mussel shells vary greatly in shape, size, color, and biomass (Lopes-Lima, 2010). The biomineralization process in bivalves is a unique species-specific architecture (Marin et al., 2012; Marie et al., 2017). Mussel shells, with their extraordinary architecture, have attracted attention as a material that can be used in many fields. To this end, many scientists have studied the definition of the microstructures of freshwater mussel shells (Hedegaard, 1997; Moura et al. 2000; Lopes-Lima, 2010; Chakraborty et al., 2020). Indeed, mussels play a major role in natural habitats, both alive and with their perfectly formed shells (Binelli et al., 2014). While freshwater mussel shells are used as seeds in oyster farming, their shells are mostly waste (Anthony and Downing 2001; Haag 2012). Waste shells are a highly valuable resource that is applicable to be used for many purposes (Yao et al., 2014) due to their high CaCO₃ content. Thus far, they have been successfully used for the mining industry (i.e. for sulphate reduction), wastewater treatment, reinforced composite production, environmental remediation (i.e. metal and paint removal, treatment of eutrophic waters), agriculture (i.e. soil improvement), bone regeneration, and cement replacement (Lee et al., 2010; Álvarez et al. 2012; Hamester et al., 2012; González-Chang et al., 2017; Delali et al., 2019). Mussel shells, which preserve the ecology and have a high marketing value, are a true aquaculture biomaterial of significant value (Chakraborty et al., 2020). The characterization of mussel shells, which
are so widely used, is of great importance. The discovery of the cross-layered aragonite structure has contributed to the preference of shells as biological materials (Meyers et al. 2008). The genus Potomida (Bivalvia, Unionida), which has a long history of taxonomic revisions, has a circum-Mediterranean distribution and P. semirugata populations have suffered dramatic declines in most rivers (Lake Gölbaşı) in southern Türkiye (Froufe et al., 2016). L. wheatleyi, the subfamily Gonideinae, is present in Gölbaşı Lake in southeastern Türkiye (Tomilova et al., 2020; Lopes-Lima et al., 2021). P. semirugata and L. wheatleyi are freshwater bivalves commonly found in Lake Gölbaşı. These species, whose habitats are muddy and sandy, migrate to deeper waters when the water temperature drops and live by burying themselves in sediments. In general, they do not prefer areas with aquatic plants as their habitat. They live by burying themselves in the sediment at a depth of 30–40 cm in warm, shallow, stagnant waters. Gölbaşı Lake is an important lake where many people depend on fishing for their livelihood. However, P. semirugata and L. wheatleyi are not sought after for their meat and shells, and their shells are abundant on the lake shore and in the substrate (Şereflişan, 2003). In the present study, the physical characteristics, chemical composition, and various biological properties of freshwater mussel shells were investigated to substantiate their potential as a useful biological material. The size (length and height), shape, and microstructure of the shells were studied to understand their physical characteristics. The chemical and microstructural composition of the shells was analyzed to determine their mineral content and present organic compounds. Various biological properties of the mussel shells were also investigated, such as their biocompatibility, hemocompatibility properties, and potential as a source of biomaterials.

2. Materials and methods

2.1. Study area

The sample collection was carried out at Gölbaşı Lake, located in the Hatay Province of southern Türkiye. Gölbaşı Lake (36°29′E, 36°30′N) covers an area of approximately 400 ha (4,000,000 m²) (Figure 1). It is a natural lake that is fed by groundwater at various places. The sampled lake has a very high fishing pressure, and it is considered to be an important habitat for P. semirugata and L. wheatleyi (Şereflişan, 2003).

2.2. Preparation of the samples

Freshwater mussels P. semirugata and L. wheatleyi (Figure 2) were collected from Gölbaşı Lake in June 2022. Their shells were rinsed with deionized water. Then, the shell length and height of the freshwater mussel species were measured with digital calipers with 0.05-mm accuracy. In the present study, the physical characteristics, chemical composition, and various biological properties of freshwater mussel shells were investigated to substantiate their potential as a useful biological material. The size (length and height), shape, and microstructure of the shells were studied to understand their physical characteristics. The chemical and microstructural composition of the shells was analyzed to determine their mineral content and present organic compounds. Various biological properties of the mussel shells were also investigated, such as their biocompatibility, hemocompatibility properties, and potential as a source of biomaterials.
were mounted on the stubs by gold-plating using a Polaron SC7620 apparatus in a sputter coater before use in the SEM. An electron acceleration potential of 15 kV was used for the microscopic observations. Photographs were taken in a SEM (JEOL JSM-638OLA; Akishima, Tokyo, Japan) at the required magnification. Chemical composition and elemental characterization of the freshwater mussel shells were carried out using the EDS attached to the SEM.

2.4. Fourier-transform infrared (FTIR) spectroscopy analysis

The study was carried out by first mixing 100 mg of KBr with 1% dried shell dust (separately for *P. semirugata* and *L. wheatleyi*). The mixture was then ground to form shell dust KBr pellets for the FTIR spectrum analysis. The infrared spectrum attributable to the shell dust was obtained using an FTIR spectrometer (Jasco FT/IR-6300 type A; serial no. A014461024). For both shell species, the spectra were collected at a resolution of 4 cm⁻¹ with a scan speed of 2 mm/s in the range of 500 to 3500 wavenumbers (cm⁻¹) (Hossain and Aditya, 2013).

2.5. Hemocompatibility test

To assess the hemocompatibility of the freshwater mussel (*P. semirugata* and *L. wheatleyi*) shell powders, their hemolytic activity percentages were determined using the modified method described elsewhere (Wang et al., 2012; Alexandre et al., 2014). Briefly, fresh blood was collected from a healthy sheep at a slaughterhouse and then stabilized with dipotassium ethylenediaminetetraacetic acid (1.5 mg L⁻¹). To obtain healthy red blood cells (HRBCs) for the hemolysis assay, the fresh blood was centrifuged immediately after collection (5000 rpm for 3 min), washed 3 times, and then diluted 10-fold with sterile phosphate-buffered saline (PBS) solution. Then, 2 mL of the diluted HRBCs suspension was transferred to a 5-mL sample vial containing 25 mg of *P. semirugata* and *L. wheatleyi* freshwater mussel shell powder, respectively. Two further sample vials containing 0.4 mL of the diluted erythrocytes and 1.6 mL of sterile distilled water or PBS solution were used as the positive and negative controls, respectively. All of the blood mixtures were then incubated at 37 °C for 2 h, followed by centrifugation (10,000 rpm, 1 min), and the absorbance of the supernatants related to hemoglobin was recorded at 541 nm using a Spectrostar Nano (BMG Labtech, Ortenberg, Germany) equipped with a BMG Labtech ultrafast UV/Vis spectrometer. Hemolytic activity was confirmed twice for each sample.

### Table

<table>
<thead>
<tr>
<th>Species</th>
<th>n</th>
<th>Length ± SD (cm)</th>
<th>Height ± SD (cm)</th>
<th>Distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. semirugata</em></td>
<td>20</td>
<td>6.7 ± 0.38</td>
<td>2.9 ± 0.50</td>
<td>Northwestern Türkiye, Eastern Mediterranean</td>
</tr>
<tr>
<td><em>L. wheatleyi</em></td>
<td>20</td>
<td>8.13 ± 0.33</td>
<td>2.8 ± 0.32</td>
<td>Türkiye, Eastern Mediterranean</td>
</tr>
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and the average of two independent experiments was used to calculate the percentage of hemolytic activity. The percentage of hemolytic activity was calculated using the equation below (1):

\[ HA(\%) = \frac{OD_s - OD_{NC}}{OD_{PC} - OD_{NC}} \times 100. \]  

(1)

Here, OD\(_s\) is the absorbance of the tested sample, OD\(_{NC}\) is the absorbance of the negative control, and OD\(_{PC}\) is the absorbance of the positive control.

2.6. Evaluation of the blood clotting property
To assess the blood clotting index (BCI) of the freshwater mussel (*P. semirugata* and *L. wheatleyi*) shell powders, the method described by Patil et al. (2018) was used with some modifications. For the analysis, sheep blood was collected at the abattoir in an anticoagulant tube and used immediately. The study was briefly performed as follows: Sponge samples were placed in the center of polypropylene petri plates and 100 µL of blood was applied to the surface of the samples. Immediately afterwards, 10 µL of 0.2M CaCl\(_2\) solution was dropped onto the surface and the samples were incubated at 37 °C for 20 min. At the end of the incubation period, 5 mL of sterile distilled water was gently added without breaking up the formed clot. In the next step, the blood-water mixture in the plates was collected and transferred to a polypropylene test tube. After centrifugation at 10,000 rpm, the supernatants were transferred to clean tubes and incubated at 37 °C for 1 h. At the end of the incubation period, the optical density of the samples was recorded at 540 nm using a Spectro star Nano (BMG Labtech) equipped with a BMG Labtech ultra-fast UV/Vis spectrometer. For the optical density measurements, sterile distilled water was used as the target and 5 mL of sterile distilled water with 100 µL of the blood mixture (which was subjected to all of the analysis steps) was used as the positive control. The clotting capacity was confirmed twice for each sample and the average of 2 independent experiments was used to calculate the BCI (%). The BCI was calculated according to the equation of Wang et al. (2012):

\[ BCI(\%) = \frac{OD_s}{OD_{PC}} \times 100 \]  

(2)

Here, OD\(_s\) is the absorbance of the tested sample and OD\(_{PC}\) is the absorbance of the positive control.

2.7. In vitro biocompatibility studies
The in vitro cytotoxicity of the *P. semirugata* and *L. wheatleyi* freshwater mussel shell powders was evaluated using the ISO 10993-5 standard test method. The antiproliferative activities of the samples were measured using the 2, 3-bis-(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide (XTT reagent) colorimetric assay. L929 mouse fibroblast cells were selected as a model cell line to determine the cytotoxicity of the shell powders for the cytocompatibility assay. The cells were cultured in Dulbecco's modified eagle medium (DMEM) supplemented with 1% (v/v) of a preprepared solution of penicillin (100 U/mL), streptomycin (100 U/mL), and 10% (v/v) fetal bovine serum (FBS) and incubated for 24 h at 37 °C with humidity above 90% in an atmosphere of 5% CO\(_2\). They were seeded at a density of \(1 \times 10^4\) cells/ well on a 96-well microplate and incubated at 37 °C for 48 h. After incubation, the cells were treated with different concentrations of compounds (2, 3, 5, and 10 µL) and incubated in an incubator at 37 °C with 5% CO\(_2\) for 48 h. After incubation, the culture medium was discarded, and the wells were rinsed 3 times with PBS. Then, 200 µL of DMEM containing 50 µL of 5 mg/mL XTT reagent was added to each well and incubated for 4 h. The absorbance of the solution was measured spectrophotometrically using a Thermo Scientific Multiskan FC Microplate Photometer reader (Thermo Fisher Scientific Inc., Waltham, MA, USA) at a wavelength of 450 nm. The experiment was repeated in triplicate (Yang et al., 2017; Kucharczyk et al., 2019; Öksüz et al., 2021).

2.8. Statistical analysis
The collected data were presented as the mean ± standard deviation of the mean SD based on at least 6 independent measurements. Data were statistically analyzed using 1-way analysis of variance (ANOVA) followed by the Dunnett and Tukey tests, and significance was achieved at *p ≤ 0.05* using Origin Pro 9.0 software (OriginLab Co., Northampton, USA).

3. Results
3.1. Microstructure observation and characterization with the SEM
The protein-containing sclerotized periostracum layer of the shells of both mussels was very prominent (Figures 3A and 3B). Apparently, the outer surface of the periostracum of the shell of *P. semirugata* and *L. wheatleyi* did not consist of radial, oblique ridges. The prismatic layer showed a typical polygonal arrangement, as shown in the magnified image in Figures 4A and 4B. The nacreous layer of the shells of (A) *P. semirugata* and (B) *L. wheatleyi* is shown in the magnified image in Figures 5A and 5B. The internal shell structure of the 2 bivalves differed in terms of the nacre (Figures 6A and 6B). The shell of *L. wheatleyi* showed an irregular crystalline structure in the nacreous layer (Figure 6B). The lateral view of the prismatic layer shows the columnar convergent prisms of (A) *P. semirugata* and (B) *L. wheatleyi* (Figures 7A and 7B). These columns grow within organic membrane structures. While the columnar prismatic structure of *P. semirugata* (Figure 7A)
Figure 3. The growth front of the outer surface of each shell of (A) *P. semirugata* and (B) *L. wheatleyi*.

Figure 4. The periostracum on the outer surface of each shell of (A) *P. semirugata* and (B) *L. wheatleyi*.

Figure 5. SEM image of the nacreous layer of the shells of (A) *P. semirugata* and (B) *L. wheatleyi*. 
was angular, that of *L. wheatleyi* (Figure 7B) appeared to be more rounded. The horizontally overlapping individual aragonite sheets of *P. semirugata* appeared to be more regular than those of *L. wheatleyi* (Figure 8).

### 3.2. EDS analysis

Figures 9A and 9B show the SEM images and EDS data of the periostracum and nacreous layers in the *L. wheatleyi* shells, respectively. The semiquantitative chemical composition revealed the presence of Ca, Na, Mg, C, and O. In the nacreous layer of *L. wheatleyi*, the chemical composition of the elements ranged from 12.33 to 12.92 atom% for C, 65.63 to 66.49 atom% for O, 0.14 to 0.2 atom% for Na, 0.07 to 0.09 atom% for Mg, 0.06 to 0.08 atom% for Si, 20.06 to 22.6 atom% for Ca, 0.15 to 0.87 atom% for Zr, and 0.57 to 0.75 atom% for Pt (Figure 9B). The chemical composition of the elements in the periostracum layer of *L. wheatleyi* ranged from 7.28 to 12.85 atom% for C, (52.14 to 67.07 atom%) for O, (0.11 to 0.18 atom%) for Na, (0.09 to 0.11 atom%) for Al, (19.3 to 39.35 atom%) for Ca, (0.03 to 0.15 atom %) for Zr, (0.49 to 1.12 atom%) for Pt, and (0.06 to 0.07 atom%) for S (Figure 9A).

Figures 10A and 10B show the SEM-EDS images and EDS data of the periostracum and nacreous layers of the *P. semirugata* shells, respectively. The semiquantitative chemical composition showed that Ca, Na, Mg, C and O were the main elements. In the nacre layer of *P. semirugata*, the chemical composition of elements ranged from 53.04 to 65.80 atom% for C, 29.30 to 38.53 atom% for O, 0.16 to 0.27 atom% for Na, 0.06 to 0.13 atom% for Mg, 0.78 to 1.59 atom% for S, and 1.52 to 4.48 atom% for Ca. The chemical composition of the elements in the periostracum layer of *P. semirugata* ranged from 7.28 to 12.85 atom% for C, 52.14 to 67.07 atom% for O, 0.11 to 0.18 atom% for Na, 0.09 to 0.11 atom% for Al, 0.06 to 0.07 atom% for S, and 19.30 to 39.35 atom% for Ca.
3.3. FTIR analysis

In this experimental study, the functional groups present in the collected mussel shells were analyzed through FTIR spectroscopy. The analysis of the FTIR spectra revealed important information about the functional groups present in the collected mussel shells. Figure 11 shows the FTIR spectra of the two mussel shells, which showed similar characteristics. The peak at 2933.76 cm\(^{-1}\) appeared due to the CH\(_2\) stretching bonds of the aliphatic chains. The spectral peak at 2362.54 cm\(^{-1}\) was identified as the stretching vibration of COO\(^-\) functional groups. The FTIR spectrum showed organic bands of lower intensity; the band at 1786.52 cm\(^{-1}\) was attributed to the carboxylate (carbonyl) groups of the acidic proteins in the organic matrix. Four
bands characteristic of aragonite, corresponding to the CO$_2$ ions, were identified: ν3 at 1467.32 cm$^{-1}$, ν1 at 1081.43 cm$^{-1}$, ν2 at 863.39 cm$^{-1}$, and ν4 at 702.31–713.73 cm$^{-1}$ (Figure 11). The ν4 band corresponded to the planar bending mode of the carbonate vibration and the ν1 band to the symmetric stretching mode. Moreover, the characteristic carbonate ν4 bands of the aragonite were at 713.73 and 702.31 cm$^{-1}$ and the characteristic carbonate ν2 band of the aragonite was at 863.39 cm$^{-1}$, indicating the availability of the aragonite form of CaCO$_3$ that is seen in the shell powders of snails (Anjaneyulu et al., 2015; Hossain et al., 2015).
3.4. Hemostatic characterization

The results of the test, including the hemolytic activity (%) and BCI (%), are shown in Figure 12. Fresh blood was used for the test, and the hemolysis rates were 1.97% and 2.07% for the \textit{P. semirugata} and \textit{L. wheatleyi} shells, respectively (Figure 12A). These results showed that the hemolysis rates slightly exhibited the same trend between the mussel shell groups (Figure 12A).

The BCI results were different between the \textit{P. semirugata} and \textit{L. wheatleyi} shells. The shells of \textit{P. semirugata} had a lower BCI value. The shells of \textit{L. wheatleyi} had a higher BCI for clot formation (Figure 12B).

3.5. In vitro biocompatibility studies

The aim of this study was to evaluate the effect of different concentrations of \textit{P. semirugata} and \textit{L. wheatleyi} shell extracts on the proliferation of L929 mouse fibroblast cells using the XTT assay. The XTT assay measured cell viability as a percentage of the viability observed in the mussel shell samples. Figure 13 shows the cell viability plot using the XTT assay and it was observed that all of the groups supported cell proliferation. The results clearly demonstrated that both of the shell samples were highly biocompatible and nontoxic when compared to the control group.

4. Discussion

The chemical composition of the mussel shells, dominated by CaCO$_3$, indicates that they are a source of CO$_2$ and are therefore important in the carbon cycle (Chakraborty et al., 2020). It involves the calcification of the shell, and the formation of an organic matrix of soluble and insoluble compounds. The presence of CaCO$_3$ is essential for this process (Moura et al. 2000). The bioformed crystals of the shell were quite different in size and shape from those of their nonbiological counterparts (Lopes-Lima et al., 2010). Although the mechanism of biomineralization in molluscs is influenced by environmental factors (such as the physicochemical parameters of the water), it is essentially a cellular process (Moura et al. 2000).

Close observation of the microstructure of the shells of two common freshwater bivalves, \textit{P. semirugata} and \textit{L. wheatleyi}, revealed a variation in their construction from the outer periostracum to the inner nacreous layer. In \textit{P. semirugata}, a polygonal arrangement with columnar prisms of different sizes combined was observed to be more pronounced. In \textit{L. wheatleyi} the prismatic columns are oval and irregularly polygonal. The length and orientation of these fibers play an important role in the growth and remodeling of the shell (Dauphin et al., 2018).

The analysis of the FTIR spectra revealed important information about the functional groups present in the collected mussel shells. As shown in Figure 2, the peak at about 2933.76 cm$^{-1}$ was attributed to the C-H stretching vibration, indicating the presence of methyl and methylene groups in the shells. The spectral peak at 2362.54 cm$^{-1}$ was identified as the stretching vibration of COO$^-$ functional groups. In addition, the band at 1467.32 cm$^{-1}$ corresponded to the C=C bond, which is indicative of unsaturated hydrocarbons. The peak at 1081.43 cm$^{-1}$ was due to C-O stretching, suggesting the presence of proteins in the shell matrix (Marie et al., 2010). The presence of CaCO$_3$ in the mussel shells was confirmed by the identification of the characteristic carbonate bands at 702.31, 713.73, and 863.39 cm$^{-1}$. This information was used to classify...
and distinguish the different layers of the shells, allowing interspecific comparisons and the study of variations in the different layers. As a result of this analysis, it was possible to identify the different layers of the shell, including the outer periostracum, the middle prismatic layer, and the inner nacreous layer. The outer periostracum is the outermost layer of the shell and consists of organic compounds such as chitin and proteins. The middle prismatic layer consists of densely packed CaCO$_3$ crystals, and the inner nacreous layer consists of CaCO$_3$ in a crystalline form arranged in a specific pattern (Sudo et al., 1997; Lopes-Lima et al., 2010; Carter et al., 2012; Hossain et al., 2015; Agbaje et al., 2017).

By distinguishing between the different layers of the shell and examining variation within and between species, a more comprehensive understanding of shell morphology and composition can be obtained. This information can be used to further explore the potential applications of mussel shells in fields such as materials science, biotechnology, and environmental remediation (Carter, 1990).

The results of the test, including the hemolytic activity (%) and coagulation index (%), are shown in Figure 12. Fresh blood was used for the test and the hemolysis rates were 1.97% and 2.07% for the *P. semirugata* and *L. wheatleyi* groups, respectively (Figure 12A). These results showed that the hemolysis rates exhibited slightly the same trend between the shell groups. The number of erythrocytes bound to nanoparticles and the energy of the cell membranes to encapsulate nanoparticles are two factors affecting hemolysis, and both are affected by the external surface area of materials (Yannan et al., 2011). The hemolytic activity of various materials used in healthcare is typically determined and classified according to the American Society for Testing and Materials (ASTM) standards. According to the ASTM F756-00 and ISO 10 993-51, a sample is considered nonhemolytic if the hemolytic index range is less than 2%, mildly hemolytic if it is between 2% and 5%, and hemolytic if it is greater than 5%. In light of the ASTM standards, the hemolysis rates obtained from the mussel shell samples indicated that they are nonhemolytic (Bohorquez-Moreno et al., 2023). Therefore, these results revealed that the studied shells can be used in medical sector applications.

To understand the results of the BCI, it is essential to understand the blood clotting assay, which evaluates the antithrombogenic properties of various biomaterials in contact with whole blood by measuring the amount of free hemoglobin in the medium. Calcium ions play a crucial role in the coagulation reactions that lead to clot formation, making it a critical element of the test (Jang et al., 2010). The BCI results for the *P. semirugata* and *L. wheatleyi* groups were related to their ability to absorb calcium ions. The capacity of the *P. semirugata* shell to absorb calcium ions was limited due to its structural characteristics, resulting in a lower BCI value. On the other hand, the *L. wheatleyi* group had a higher BCI value. This demonstrated that it has a higher tendency toward clot formation since it can

![Figure 13. Cell viability (%) of the freshwater mussel shell powder. *Significant differences compared to the control at p < 0.05.](image-url)
absorb calcium ions more effectively (Singh et al., 2019). The aim of the study was to evaluate the effect of different concentrations of *P. semirugata*, *L. wheatleyi* shell extracts on the proliferation of L929 mouse fibroblast cells using the XTT assay. The XTT assay measured the cell viability as a percentage of the viability observed in the mussel shell samples. Figure 3 shows the cell viability plot using the XTT assay, wherein it can be seen that all of the groups supported cell proliferation with cell viability values above approximately 85%. It was also noted that none of the materials were cytotoxic in the experimental study, as reported in the literature (Latire et al., 2017; El-Bassyouni et al., 2020). Figure 10 shows a significant difference in cell activity for the *P. semirugata* group (107.6 ± 0.8, 3 µL) and the *L. wheatleyi* group (103.4 ± 0.6, 10 µL), indicating their potential to support cell growth. Indeed, mollusc shells contain numerous macromolecules, including soluble proteins, glycoproteins, hydrophobic proteins, chitin, soluble polysaccharides, and lipids (Lowenstam and Weiner 1989; Bédouet et al. 2001; Marie et al. 2011; Marin et al. 2012). *Mytilus edulis* and *Crassostrea gigas* shell extracts have been reported to support cell proliferation, and it has been suggested that shell proteins may play an inducing role in such a finding (Latire et al., 2017). Similarly, a study of the effect of the pearl mussel *Hyriopsis cumingii* Lea on the proliferation of osteoblastic lineage cells showed an increase in the proliferation of these cells (Shen et al. 2006). Although a slight decrease in cell viability was observed in the *P. semirugata* (91.62 ± 0.8, 5 µL) and *L. wheatleyi* (89.25 ± 0.6, 5 µL) groups, the results clearly demonstrated that both shell samples were highly biocompatible and nontoxic compared to the control group. The work presented herein shows that the materials have promising potential for use in various biomedical applications where biocompatibility is a critical factor.

**Ethical approval**

Ethical approval was not needed for this study, as the study dealt with invertebrate organisms.

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**References**


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