Establishment of long-term ostracod epidermal culture

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Abstract
Primary crustacean cell culture was introduced in the 1960s, but to date limited cell lines have been established. *Skogsbergia lerneri* is a myodocopid ostracod, which has a body enclosed within a thin, durable, transparent bivalved carapace, through which the eye can see. The epidermal layer lines the inner surface of the carapace and is responsible for carapace synthesis. The purpose of the present study was to develop an in vitro epidermal tissue and cell culture method for *S. lerneri*. First, an optimal environment for the viability of this epidermal tissue was ascertained, while maintaining its cell proliferative capacity. Next, a microdissection technique to remove the epidermal layer for explant culture was established and finally, a cell dissociation method for epidermal cell culture was determined. Maintenance of sterility, cell viability and proliferation were key throughout these processes. This novel approach for viable *S. lerneri* epidermal tissue and cell culture augments our understanding of crustacean cell biology and the complex biosynthesis of the ostracod carapace. In addition, these techniques have great potential in the fields of biomaterial manufacture, the military and fisheries, for example, in vitro toxicity testing.

Keywords Crustacean · Ostracod · Carapace · Epidermal · Culture

Introduction

Myodocopid ostracods (or ‘seed shrimps’) are a suborder of small marine crustaceans of length between 1 and 32 mm, which pass through a series of six, juvenile growth stages known as instars before reaching the adult stage (McKenzie et al. 1999). Their shrimp-like bodies are suspended within a relatively thin (20–60 μm), bivalved calcified carapace by ‘central adductor muscles’, which run through the body attached to each valve; a hinge joins the two valves enabling them to open and close their carapace while preserving its rigidity (McKenzie et al. 1999). The majority of myodocopid ostracods are benthic; they bury up to 1 cm depth in the sediments of the ocean floor (McKenzie et al. 1999). Most myodocopid ostracods can be distinguished from other ostracods as they possess good vision with well-developed compound eyes (Parker et al. 2019), up to one third of their overall body size (Parker 1998). Some myodocopids in the family Cypridinidae are known to possess transparent carapaces or transparent windows in the parts of the carapace that cover the eyes (Parker et al. 2019). The cypridinid species used in the present study, *Skogsbergia lerneri*, is 2–3 mm in length and found at a depth of 1–130 m along the eastern coast of the American continents from Florida to the top of Brazil and the Caribbean islands (Kornicker 1958) and possess a carapace with uniform transparency.

The carapace structure comprises four layers: a thin outer layer devoid of chitin known as the epicuticle; a dense laminar layering of chitin forming the exocuticle; a less dense laminar layer known as the endocuticle and the epidermal cell layer (Stevenson 1985). This epidermal layer is essential to the formation of the carapace; it shows intense secretory activity and orchestrates secretion of the other layers (Stevenson 1985; Mrak et al. 2015). In addition, the epidermal cells are essential for the reabsorption of calcium from preceding layers just
before ecdysis (or moulting) to store minerals (Glötzner and Ziegler 2000). Ecdysis involves the shedding and formation of old to new exoskeleton and is a fundamental process that is essential for growth of all crustaceans. Understanding the complexity of creation and assembly of the thin, highly transparent, yet presumably strong and durable ostracod carapace would provide huge potential for material manufacture, with numerous applications in the biomaterials industry, as well as facilitating marine in vitro toxicity testing.

To date, crustacean cell culture has proved problematic and is limited (Toullec 1999; Jayesh et al. 2012). The purpose of the present study was to develop an epidermal tissue and cell culture method for *S. lerneri*. To achieve this aim, first an optimal environment for the growth and viability of this cell type was determined, while maintaining sterility and proliferative potential. Next, a microdissection technique to remove the epidermal layer for explant culture was established and finally, a cell dissociation process for cell culture was determined.

**Materials and Methods**

*Source of Tissue* *Skogsbergia lerneri* ostracods used in the present study were collected biannually off the coast of Florida following the acquisition of Special Activity licenses (SAL-16-1796-SR and SAL-19-1796-SR) and Florida Keys National Marine Sanctuary permits (FKNMS-2016-116 and FKNMS-2018-116). The ostracods were then transported to Cardiff University, where, after conditioning to aquaculture conditions for the duration of their lifecycle or until required for experimentation.

For carapace-epithelial culture, the epithelial layer was finally, a cell dissociation process for cell culture would be determined. To isolate the carapace, ostracod valves were opened, under a dissection microscope, by inserting the corner of a single-edged razor blade into the opening at the caudal furca, and the adductor muscle was severed (Fig. 1). While stabilised by forceps at the hinge region, the carapace was opened and internal body parts were removed. The carapace, with or without epidermal layer, was carefully dissected away using microdissection needles.

**X-Ray Microtomography** In order to understand ostracod carapace microstructure, prior to microdissection described above, X-ray microtomography was performed on adult ostracods on Beamline I13 (MT17839-1), at the Diamond Light Source national synchrotron facility (Didcot, UK). This enabled rapid, non-destructive, three-dimensional high spatial and contrast resolution imaging of the *S. lerneri* structure and components. Whole adult *S. lerneri* ostracods (*n* = 3) or single carapace valves (*n* = 3) were mounted onto the metal pin of sample holders (CN Technical Services Ltd, Wisbech, UK) using super glue (Loctite, Westlake, OH).

Raw images were collected using a monochromatic beam and pco.edge 5.5 detector. Samples were scanned through 180°, using a 0.5-s acquisition time at ×4 total magnification and a 12 cm sample-to-detector distance for whole ostracods. For carapace valves, scans were performed at a ×10 total magnification using a 0.2-s acquisition time and a 22.5-cm sample-to-detector distance. Raw data were reconstructed using a series of tomocommands and ImageJ, and then rendered using Avizo 9 software.

**Optimisation of Ostracod Microdissection for Epidermal Culture** Ostracod microdissection was performed, as described above, in ice-cold sterile saline under a dissection microscope. Next, carapace halves were decontaminated prior to placement in culture. Following dissection, carapace halves were incubated in 2.5 μg/ml amphotericin B (Sigma, Gillingham, UK) and/or 2% povidone–iodine in saline (Sigma) for 0–4 min (*n* = 10 per disinfectant tested at each time point). Cell viability, expressed as a percentage, was determined by quantitation of live (green fluorescence) and dead cells (red fluorescence) using a live/dead assay kit (Abcam, Cambridge, UK, ab115347) in images captured using a Leica 6000 fluorescent microscope and Leica Application Suite X (LAS X) software.

For carapace-epithelial culture, the epithelial layer was retained on the carapace. For epidermal explant culture, the epidermal layer was carefully dissected away using microdissection needles (Fig. 1).

**Optimisation of Medium for Epidermal Culture** To determine the optimal combination of culture medium and supplements for epidermal cell culture, carapace-epithelia (i.e. valve with epidermal layer intact, *n* = 15 per culture condition) and/or epidermal explants (*n* = 15 per culture condition) of later-stage lifecycle *S. lerneri* (instar 4–adult) were cultured in...
various media/supplement combinations at ambient temperature (20°C), 25 or 37°C, either in air or 5% CO₂. These temperatures were chosen as a starting point to establish ostracod epidermal cultures based on successful cell and/or tissue culture described for marine fish (ambient room temperature; Nicholson 1989), invertebrates (25–30°C; Hink 1979) and also the seawater temperature of their local habitat at the Florida Keys. In addition, among the culture media tested (see below) were those often used for mammalian cell culture at 37°C. Although this temperature is high for crustacean cells, it has been used to establish fish cell cultures (Shima et al. 1980; Nicholson 1989), so it was included.

Culture media included Leibovitz-15 (L-15) (Sigma), medium 199 (M199) (Sigma), DMEM (high and low glucose) (Thermo Fisher Scientific, Loughborough, UK), Grace’s (Sigma), Schneider’s (Sigma) and Hink’s TNM-FH (Sigma) insect media. Supplements, previously proposed to enhance crustacean cultures (Chen et al. 1986; Luedeman and Lightner 1992; Hsu et al. 1995; Frerichs 1996; Kasornchandra et al. 1999; Toullec 1999; Walton and Smith 1999; Mulford et al. 2001; Shimizu et al. 2001; Lang et al. 2002a; Maeda et al. 2003), in different combinations and/or concentrations, included heat-inactivated foetal bovine serum (FBS, 5, 10 or 20%; Sigma), amphotericin B (2.5 or 0.25 μg/ml), penicillin–streptomycin (100 or 200 μg/ml; Sigma), GlutaMAX (4 or 2 mM; Sigma), MEM non-essential amino acids (1/1000; Gibco, Fisher Scientific, Loughborough, UK), 10 mg/ml copper II sulphate (Sigma), 10 ng/ml epidermal growth factor (EGF; Sigma), 20 ng/ml fibroblast growth factor-basic (FGF; Sigma), 10% salt solution (sodium citrate, Sigma; calcium chloride, Sigma; magnesium chloride, Sigma), 10 mM HEPES (Sigma), 2 mM L-glutamine (Sigma) and sodium chloride (5 or 6 g/l; Sigma).

In addition, to promote epidermal adhesion, epidermal explants were cultured on different substrates (plastic, fibronectin (Sigma), collagen type I (Corning, Corning, New York)) on 12 or 24 multi-well plates (Starlab, Milton Keynes, UK), culture inserts (Corning) and/or 35-mm culture dishes (Corning). Percentage cell viability of epidermal explants cultures (up to 4 wk) and carapace-epithelia cultures (up to 3 wk) was evaluated as described above, with Hoechst 33342 nuclear counterstaining (Molecular Probes, Life Technologies, Loughborough, UK).

**Epidermal Cell Proliferation in Carapace-Epithelia–EdU Incorporation** Cell proliferation in ostracod epidermal cells (n = 5 carapace-epithelial cultures) was assayed using a Baseclick EdU HTS Kit (Sigma). EdU (5-ethynyl-2′-deoxyuridine) was incorporated into the DNA of cycling cells at the beginning of culture by addition of medium containing EdU for 24 and 72 h. The epidermal cells were fixed in 4% paraformaldehyde for 15 min, then permeabilised using 0.1% Triton X-100 in PBS for 10 min, before incubation in the click assay mixture (reaction buffer, catalyst solution, 5-TAMRA-PEG3-Azide dye (excitation = 546 nm, emission = 579 nm) and buffer additive) for 30 min in the dark, at room temperature. Cell nuclei were counterstained with Hoechst 33342 (Molecular Probes, Life Technologies) and visualised using fluorescence microscopy for calculation of percentage proliferating cells.

**Epidermal Cell Proliferation in Carapace-Epithelia–BrdU Incorporation** BrdU immunolocalisation was assayed in cultured intact carapace-epithelia (n = 5) following BrdU incorporation 

in vitro over 48 h in a 10 μM BrdU (ab142567; Abcam) labelling solution. The samples were prepared, fixed and permeabilised as described above for EdU incorporation, then incubated in 1 M HCl for 30 min at room temperature. The samples were then blocked in 5% normal goat serum (ab156046; Abcam) in PBS for 1 h, incubated in rabbit polyclonal anti-BrdU (1/100 dilution, Abcam, ab152095) overnight at 4°C and then rinsed three times in 1× PBS. After an
incubation for 2 h in the dark at room temperature in goat anti-rabbit IgG H&L conjugated to Alexa Fluor 555 (1/1000 dilution, ab150086, Abcam), the samples were mounted in ProLong Gold Anti-fade mountant with DAPI (Thermo Fisher Scientific) and visualised using fluorescent microscopy. Percentage cell proliferation was quantified.

Transmission Electron Microscopy of Cultured Ostracod Epidermal Tissue Cultured epidermal tissue on adult ostracod carapaces at three culture time points (0, 1 and 2 wk, n = 3 at each time point) were removed from culture and incubated at 4°C overnight in Karnovsky fixative (2% paraformaldehyde, 2.5% glutaraldehyde, in 0.1 M cacodylate buffer). Fixed carapace-epidermal valves were transferred to 1.5% potassium ferricyanide/1% osmium tetroxide in 0.1 M cacodylate buffer for 1 h and washed in distilled water, before 1-h immersions in 1% aqueous osmium tetroxide and 1% aqueous uranyl acetate, with thorough washing in distilled water after each. After an additional 1-h incubation in Walton’s lead aspartate (Walton 1979) at 60°C with further washes, samples were dehydrated through serial ethanols to 100%, immersed in propylene oxide, before Durcupan resin infiltration and embedding over 2 d. The resin was hardened at 60°C for 48 h and ultrathin sections (100 nm thick) were cut and viewed on G300 copper grids in a JEM 1010 transmission electron microscope (Jeol, Welwyn Garden City, UK).

Optimisation of Epithelial Cell Dissociation To determine the most effective enzyme for dissociation of the epidermal cells, a number of enzymes were tested (see Table 1) on dissected epidermal layers from later-stage lifecycle S. lerneri ostracods (instar 4–adult, n = 30 ostracods per enzyme protocol). Of all enzymes tested, Dispase I (Sigma) at a concentration from 0.6 to 1.0 U/ml, dissolved in M199 (DPBS) at 25°C was found to be optimal, based on successful cell dissociation of epidermal tissue into discrete cells in a 30-min incubation, while maintaining cell viability.

To further optimise the cell dissociation protocol using Dispase I, epidermal tissue was pooled from 30 ostracods for each protocol tested (triplicate samples for each) which included incubations in Dispase I in M199 (0.6–1.0 U/ml) at 25°C for 5, 10, 20, 30 or 40 min.

The dissociated cells and medium were then passed through an 8-µm filter (MF-Millipore, Sigma) to separate the dissociated cells from any residual tissue remnants. Next, the dissociated cells were thoroughly, but gently, washed in M199 medium, within wells containing 4-µm inserts to fully remove enzyme, while avoiding loss of cells. The cells were then cultured on inserts on coated wells in M199 containing 2 mM GlutaMAX, 10% non-essential amino acids, 200 µg/ml penicillin streptomycin and 0.25 µg/ml amphotericin B for 0, 1, 2 and 3 d. Cell viability, quantified as a percentage, was assessed at each time point, by using a live/dead assay kit (Abcam, ab115347), as described above for epidermal culture.

Determination of Substrate for Epidermal Cell Culture Using optimal conditions for dissociation of the epidermal cells (incubation in Dispase I in M199 (0.6–1.0 U/ml) for 10 min), ostracod epidermal cells were plated on different substrates (i.e. cells were pooled from 30 ostracods per substrate-coated well) to promote adherence during culture in the optimised M199 medium described above. Substrates included collagen type I (48-well; Sigma), collagen type IV (48-well; Sigma), laminin (24-well; Corning), fibronectin (24-well, BioCoat; Corning), Matrigel (48-well; Sigma) and chitosan. For chitosan coatings, 4% (w/v) chitosan (>90% degree deacetylation; Sigma) in 2% (v/v) acetic acid was autoclaved, applied to 48-well plates and allowed to dry for 24 h to form a thin film. The acidity of the films was neutralised with 0.5 M NaOH and washed repeatedly with Hank’s balanced salt solution until film pH returned to a physiological range (pH = 7.4).

Proliferation of Dissociated Epidermal Cells in Culture Epidermal cell proliferation of cultured ostracod epidermal cells was assessed after 7 d of culture in optimised M199 medium, described above, on substrates collagen type I (n = 3 cultures) or IV (n = 3 cultures). The latter were deemed optimal substrates for epidermal cell adherence, viability and spreading (see ‘Results’ section). In brief, following 7 d in culture, cells were incubated in 10 µM BrdU culture medium for 24 h and assayed as described above.

Results

X-Ray Microtomography Reconstructions X-Ray microtomography 3D reconstructions of whole adult S. lerneri ostracods (n = 3) and single carapace valves (n = 3) were rendered to produce high spatial and contrast resolution images that were used to provide information on ostracod microstructure, prior to microdissection (Fig. 2). The 3D images revealed the furca region opening of the carapace, through which the corner of a razor blade could be inserted and the adductor muscle severed, facilitating effective dissection. Image slices (example shown in Fig. 2b) from within a 3D image stack were used to make thickness measurements of the ostracod carapace and epidermal cell layer, in order to get an idea of scale. The carapace and epidermal cell layer thickness in these ostracods ranged from 13.67 to 17.09 µm and 12.48–13.88 µm, respectively.

Epidermal Tissue Culture–Microdissection and Disinfection Protocols Prior to Culture Epidermal cell viability decreased...
with time of exposure to amphotericin B and/or 2% povidone–iodine (Fig. 3) over 4 min. A 13.5–21.4% decrease in viability was observed within the first minute of exposure to either of these anti-microbial agents, decreasing from 66.1% (SEM ± 2.9%), 65.3% (SEM ± 3.4%) and 64.7% (SEM ± 2.4%) to 52.6% (SEM ± 1.7%), 46.3% (SEM ± 1.9%) and 43.3% (SEM ± 1.8%) for 2% povidone–iodine, amphotericin B or a combination of both, respectively. Due to these toxic effects, these disinfectants were eliminated from our method. Instead, a strict protocol was developed using sterile ice-cold 0.9% saline, renewed for each ostracod wash with time restrictions (i.e. 2-min anaesthetisation, a second 20-s wash, before 2 min in ethanol). The ostracod was then subjected to a final wash in 0.9% ice-cold saline water (20 s) before undergoing dissection in 0.9% ice-cold saline water under a dissecting microscope. In addition, autoclaved dissection equipment and immersion of dissection tools in 2% povidone–iodine, followed by 70% ethanol, between each dissection stage was implemented.  

### Table 1  Enzymatic dissociation of ostracod epidermal cells. A number of protocols were tested, with optimal cell dissociation achieved following incubation in 0.6–1.0 U/ml Dispase I in M199 at 25°C for 30 min

| Enzyme/enzyme combination                                      | Temperature (°C) | Duration                  |
|----------------------------------------------------------------|-----------------|---------------------------|
| 0.04% EDTA in DPBS                                             | 25              | 30 min, 2 h, 2.5 h        |
| 0.04% EDTA + Dispase II (2.4 U/ml)                            | 25              | 1 h, 2 h, 3 h             |
| Accutase                                                      | Room temperature, 25, 37 | 30 min, 10 min, 20 min, 1 h |
| 0.25% Trypsin/EDTA                                            | 25              | 20 min, 1 h               |
| 0.05% Trypsin/EDTA                                            | 25, 37          | 20 min, 1 h, 2 h          |
| 0.15% Collagenase I                                           | 25              | 2 h, 2.5 h                |
| Collagenase I (200 U/ml) + Dispase II (2.4 U/ml) + 3 mM CaCl₂ | 25, 30, 37      | 30 min, 2 h               |
| Dispase II in DPBS (2.4 U/ml)                                 | 25              | 1 h, 2 h                  |
| Dispase I in DPBS (0.6–1.0 U/ml)                              | 25              | 30 min                    |
| Dispase I in M199 (0.6–1.0 U/ml)                              | 25              | 1 h                       |
| 0.05% Trypsin/EDTA + Dispase II (2.4 U/ml)                    | 25              | 1 h, 2 h, 2.5 h           |
| 0.25% Trypsin/EDTA + collagenase I (200 U/ml)                 | 25              | 1 h, 2 h                  |
| 0.05% Trypsin/EDTA + collagenase I (200 U/ml)                 | 25              | 1 h                       |
| 0.05% Trypsin/EDTA + collagenase I (200 U/ml) + 3 mM CaCl₂    | 37              | 4 h                       |
| Accutase → 0.05% Trypsin/EDTA + collagenase I (200 U/ml) + 3 mM CaCl₂ | Room temperature → 37 | 20 min → 2 h |
| 0.25% Collagenase IV                                          | 25              | 30 min                    |
| Dispase II (2.4 U/ml) + 0.25% collagenase IV                  | 37              | 1 h                       |
| Dispase II (2.4 U/ml) + 0.04% EDTA + 0.25% collagenase IV + collagenase I (200 U/ml) | 37 | 1 h |

The optimised culture conditions (based on cell viability) for *S. lerneri* carapace-epithelial tissue and epidermal explant cultures included a base medium of M199 with supplements, 10% FBS, 0.25 μg/ml amphotericin B, 200 μg/ml penicillin streptomycin, 10% non-essential amino acids and 2 mM GlutaMAX. The preferential substrate for epidermal explants was fibronectin-coated 35 mm dishes. The preferred culture environment was 25°C within a 5% CO₂ atmosphere. These optimised conditions promoted explant adhesion and cell viability and enabled culture durations up to 4 wk with an average of 60.2% (*n* = 6, SEM ± 5.7%) cell viability for epidermal explants; and 3 wk with an average of 47.4% (*n* = 5, SEM ± 1.6%) cell viability for carapace-epithelia cultures (Fig. 4). Hoechst labelling of cell nuclei confirmed the presence and location of cells in cultured explants at each time point (Fig. 5).  

Cell proliferation assayed following EdU incorporation in the ostracod carapace-epithelia *in vitro* was 17.0% (*n* = 5, SEM ± 1.8%) and 9.0% (*n* = 5, SEM ± 1.9%) of cells at 24 h (Fig. 6a, b) and 72 h (Fig. 6c, d), respectively. Cellular proliferation was confirmed to be 14.0% (*n* = 5, SEM ± 2.8%) in the ostracod carapace-epithelia *in vitro* following BrdU incorporation into carapace halves cultured for 48 h (Fig. 7).
Electron Microscopy of Cultured Ostracod Carapace-Epithelia

Electron microscopy of cultured carapace-epidermal tissue revealed distinct changes with culture duration (0–2 wk—Fig. 8). At 0 wk (Fig. 8a, b), the uncultured epidermal layer cells were tightly packed adjacent cells. After 1 wk in culture, cells appeared less dense (Fig. 8c, d), with gaps present. Cells were no longer in close association with the endocuticle by 2-wk culture (Fig. 8e, f).

**Figure 2.** X-Ray microtomography reconstruction images of an adult ostracod used to understand ostracod carapace microstructure, prior to microdissection. (a) A whole adult ostracod rotated to reveal the inner body parts and appendages through the furca region of the partially open carapace. (b) Microtomography image slice at a depth of 810 μm from within a 3D image stack of an ostracod. (c) A cross-section of the ostracod bivalve displaying the layering of the carapace. C denotes ostracod carapace and E denotes underlying epidermal cell layer.

**Figure 3.** Average percentage cell viability following post-dissection disinfecting steps (n = 10 per disinfectant tested at each time point). Live/dead labelling was used to assess the viability of cells following immersion of carapace halves in 2% povidone–iodine, amphotericin B or a combination of both for 0 to 4 min. Epidermal cell viability decreased with time of exposure to the disinfectants both alone and in combination. Error bars: SEM.

**Figure 4.** Percentage cell viability data for epidermal explant and carapace-epithelia culture (n = 15 per culture condition at 0 wk). Cultures in the optimised medium, which comprised a base medium of M199 with supplements, 10% FBS, 0.25 μg/ml amphotericin B, 200 μg/ml penicillin–streptomycin, 10% non-essential amino acids and 2 mM GlutaMAX, showed a high proportion of live cells at up to 4 wk in epidermal explants and up to 3 wk in carapace-epithelia. Error bars: SEM.
viability (Table 1; Fig. 9a–j). After a 5-min incubation, clumps of cells and tissue remnants were observed (Fig. 9a), while after 10- and 40-min incubations, cell clumps were not identified, and cells appeared fully dissociated into individual cells (Fig. 9b, c) and were viable (Fig. 9e, f, i). However, epidermal cell viability in culture was affected by duration of the Dispase I incubation (Fig. 9), especially after a 40-min incubation with a decline in cell viability at 1 d ($p < 0.05$), 2 d ($p < 0.01$) and 3 d ($p < 0.01$) in culture compared with the 0-d culture, from 65.8% ($n = 3$, SD ± 0.8%) at day 0 to 30.9% ($n = 3$, SD ± 1.7%) on culture day 3. In addition, a significant decrease in cell viability was observed at culture days 1 ($p < 0.05$), 2 ($p < 0.01$) and 3 ($p < 0.01$) in cells subjected to a 5-min Dispase I incubation. The latter may be a result of retention of clumped cells and tissue remnants at this 5-min incubation time. Since complete dissociation of cells (Fig. 9b, e, h) was achieved at 10 min, while maintaining the cell viability over all cultures at all time points, this was deemed the optimal Dispase I incubation time.

**Epidermal Cell Culture - Optimal Dissociation Protocol for Epidermal Cell Culture** Dissociated cells adhered to all substrates tested by culture day 7 (Fig. 10). Cells on laminin (Fig. 10a), chitosan (Fig. 10d), collagen type IV (Fig. 10c) and collagen type I (Fig. 10f) were more easily identified and appeared greatest in number, compared with those on

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**Figure 5.** Confirmation of cells in cultured ostracod epidermal explants. Hoechst 33342 nuclear stain was used to confirm location and presence of cells after (a, e) 1, (b, f) 2, (c, g) 3 and (d, h) 4 wk in culture (a–d ×10 magnification, e–h ×40 magnification). The ostracod epidermal layer is not a flat single layer of cells and, therefore, when visualised the nuclei were in different planes and it was not possible to image all nuclei in focus.

**Figure 6.** Confirmation of epidermal cell proliferation–EdU incorporation. EdU labelling in the epidermal layer of cultured carapace-epithelia following (a) 24 h and (b) 72 h of incubation. Cell proliferation was 17.0% ($n = 5$, SEM ± 1.8%) and 9.0% ($n = 5$, SEM ± 1.9%) of cells at 24 and 72 h, respectively. The ostracod carapace is a curved surface and this surface could not be flattened during sample visualisation. Therefore, when the carapace-epithelia were visualised, the nuclei were in different planes and it was not possible to image all nuclei in focus.
Figure 8. Electron microscopy of cultured epidermal tissue on adult ostracod carapaces at three culture time points (0, 1 and 2 wk, \( n = 3 \) at each time point). (a, b) At 0 wk in culture, the tissue is highly organised in structure compared with (c, d) 1 wk and (e, f) 2 wk. The ‘e’ denotes the epidermal cell layer. Images were captured at \( \times1200 \) (left-hand column) and \( \times2500 \) (right-hand column) magnification. Scale bar: 2 \( \mu m \).

Figure 7. Confirmation of epidermal cell proliferation–BrdU labelling. (a, b) Fluorescent microscopy images of Rabbit anti-BrdU (ab152095) labelling (red fluorescence) in two of the cultured carapace-epithelia samples visualised following in vitro BrdU incorporation into carapace halves cultured for 48 h. Cellular proliferation was confirmed to be 14.0\% (\( n = 5 \), SEM \( \pm \) 2.8\%). Cell nuclei = blue following Hoechst 33342 nuclear staining.
Epidermal Cell Proliferation Percentage epidermal cell proliferation assayed following BrdU incorporation after 7 days of culture on collagen type I (Fig. 11a) and type IV (Fig. 11b) coated plates was 23.7% (n = 3, SD ± 6.0%) and 19.0% (n = 3, SD ± 6.7%), respectively (Fig. 11c).

Discussion

Crustacean cell culture has acquired recognition as a compelling model to aid in the production of diagnostic reagents and probes for use in the shrimp, crayfish and lobster industries (Toullec 1999). Hundreds of cell lines from insects and mammals have been developed; however, despite numerous attempts, current marine crustacean cell lines reported on the Cellosaurus resource database are limited to two SV-40 transformed cell lines of Penaeus stylirostris (Pacific blue shrimp) (OKTr-23 (RRID:CVCL_9U41) and OKTr-1 (RRID:CVCL_9U40); Tapay et al. 1995). To date, no ostracod-derived cell lines have been reported.

The aim of this work was to determine the best methods for sample preparation and the culture of ostracod S. lerneri epidermal tissue (either as carapace-epidermal tissue or explants) and epidermal cells (i.e. cells dissociated from explants). In order to develop a cell culture medium optimised for these cultures, reliable methods of assessing cell viability and proliferation were required. The live/dead labelling assay used provided a simple, fast and quantitative method of evaluating cell viability in order to determine the effectiveness of the medium and/or supplements combination to support cell longevity in culture.

Primary cells introduced into culture are susceptible to contamination, but antibiotics or antimicrobials may limit, but not prevent, the growth of contaminants (Fogh et al. 1971). Yeast and fungal contaminations are especially problematic; therefore, aseptic primary culture techniques are essential. Optimisation of removal of viable epidermal tissue for culture was achieved, despite a number of potentially detrimental processes, including anaesthetisation, decontamination, epidermal dissection and culture protocols, while preventing contamination of cultures. Decontamination of ostracod carapace halves with amphotericin B and/or 2% povidone–iodine had considerable toxic effects on epidermal cell viability even before culture. Therefore, stringent protocols to eliminate contaminant exposure were developed to ensure epidermal cell viability for ostracod cultures of epidermal explants, carapace epidermal tissue cultures and cell cultures.

Thereafter, a variety of media and an extensive range of supplements, including nutrients, growth factors, chemical elements, inorganic compounds and amino acids, were trialled to develop an optimal culture medium. Primary cell cultures from ostracods have yet to be developed; therefore, knowledge of potential media and supplements was sourced from the literature reporting on other attempts of crustacean cell culture. These were predominantly of shrimp primary cell cultures, attained from a number of organ sources, including epidermal cultures (Chen et al. 1986; Luedeman and Lightner 1992; Toullec and Dauphin-Villemant 1994; Hsu et al. 1995; Frerichs 1996; Toullec et al. 1996; Kasornchandra et al. 1999; Owens and Smith 1999; Toullec 1999; Walton and Smith 1999; Mulford et al. 2001; Shimizu et al. 2001; Lang et al. 2002a; Lang et al. 2002b; Maeda et al. 2003). Of the base media tested, M199, which contains a wide variety of vitamins and amino acids, prolonged epidermal tissue survival in culture. This was consistent with M199, along with L-15, being used in the successful culture of crustacean primary cells from various tissues (Brody and Chang 1989; Luedeman and Lightner 1992; Nadala et al. 1993; Ghosh et al. 1995; Hsu et al. 1995; Frerichs 1996; Toullec et al. 1996; Chen and Wang 1999; Itami et al. 1999; Owens and Smith 1999; Shimizu et al. 2001; Lang et al. 2002a, 2002b; Maeda et al. 2003). Inclusion of low concentrations of amphotericin B and penicillin–streptomycin within the media successfully prevented fungal and bacterial contaminations.

Thereafter, an extensive range of supplements, including nutrients, growth factors, chemical elements, inorganic compounds and amino acids, were trialled to develop an optimal culture medium. Of the supplements tested, FBS, non-essential amino acids and GlutaMAX were found to enhance primary epidermal cell survival in both tissue and cell culture. The inclusion of FBS was based on it being a highly complex component used frequently to supplement media with minerals, lipids and hormones (Ma et al. 2017), as well as providing growth and adhesion factors to promote cell attachment and proliferation (Fang et al. 2017). It is unlikely to provide all of the characteristic hormones and growth factors required by crustacean cells (Mulford et al. 2001), although for S. lerneri epidermal cell culture it proved to be adequate for cell survival. Non-essential amino acids were added, as the balance of amino acids in crustacean cultures can impact on cell survival and growth rate (Shimizu et al. 2001).

The inclusion of GlutaMAX was essential as a source of carbon and energy (Butler and Christie 1994), as well as being able to improve cell survival (Pasieka and Morgan 1959). GlutaMAX was used over L-glutamine as it does not degrade spontaneously in culture or generate toxic ammonia which can
limit cell growth (Schneider et al. 1996) and is known to be toxic to life in aquaculture.

In contrast to previous studies of crustacean culture systems in which supplements, such as NE salts (Shimizu et al. 2001), HEPES buffer (Luedeman and Lightner 1992; Hsu et al. 1995; Touleec 1999; Lang et al. 2002a), NaCl (Chen et al. 1986; Hsu et al. 1995; Freirichs 1996; Kasornchandra et al. 1999; Walton and Smith 1999; Shimizu et al. 2001) and bFGF (Hsu et al. 1995), increased cell viability and longevity, these made no contribution to S. lerneri epidermal cultures.

Ostracod epidermal cells proliferated in cultures of carapace-epidermal tissue, explants and cells, demonstrating cell survival following tissue dissection and/or isolation. The decline in proliferation over culture duration may be indicative of initial stress or ‘culture shock’ activation of cell proliferation when placed in culture (Sherr and DePinho 2000; Lang et al. 2004) followed by a period of acclimatisation. The percentage of proliferating cells was similar in all tissue and cell cultures, supporting the use of this optimised medium for epidermal cell survival.

In addition, the proliferative state of cells may be impacted by the ostracod moult stage. Typically, ostracods develop through a series of juvenile instars via a mouling process (ecdysis) which ends when the mature adult instar is reached (Smith et al. 2015). The moult cycle of crustaceans is described in stages: postmoul, intermoul, premoul and ecdysis. In an ostracod lifecycle, the younger instars moult more rapidly than the more mature animals and there is substantial individual variation in instar duration (Turpen and Angell 1971). Some crustaceans (e.g. decapods) spend much of their cycle in the intermoul stage (Passano 1960), where epithelial cells in contact with the cuticle display cellular inactivity (Green and Neff 1972; Chassard-Bouchaud and Hubert 1973; Hubert and Chassard-Bouchaud 1978). It is only during sub-stages of premoul where epithelial mitotic activity becomes apparent (Skinner 1962, 1966; Stevenson 1972; Hubert and Chassard-Bouchaud 1978). It is conceivable that ostracod epithelial cells are only mitotically active for defined and varying time periods during their lifecycle. Further work...
is required to confirm the proliferative state of epithelial cells at different moult stages of *S. lerneri*.

**Conclusions**

The present study is the first to describe the culture of viable ostracod epidermal tissue and cells. Having established a method of maintaining the epidermal layer and cells in vitro, with cell proliferation, there is potential to explore possible ways of mimicking carapace synthesis in three-dimensional culture. The knowledge of this complex construction could then be used to replicate its manufacture at an industrial level to assemble materials where transparency, durability and strength are prerequisites for their functional use. Progress towards establishing marine crustacean cell lines also has positive ramifications for the field of in vitro toxicity testing. Therefore, future research will be focused on extending the survival time of primary ostracod cells and the development of an immortal cell line for sustained and indefinite expansion in vitro.

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