Molecular study supports the position of the New Zealand endemic
genus Lamellomorpha in the family Vulcanellidae
(Porifera, Demospongiae, Tetractinellida),
with the description of three new species

Michelle KELLY¹,*, Paco CÁRDENAS²,*, Nicola RUSH³, Carina SIM-SMITH⁴,
Diana MACPHERSON⁵, Mike PAGE⁶ & Lori J. BELL⁷

¹,³,⁴Coasts and Oceans National Centre, National Institute of Water and Atmospheric Research,
P.O. Box 109–695, Newmarket, Auckland, New Zealand.
²Pharmacognosy, Department of Medicinal Chemistry, BioMedical Centre, Husargatan 3,
Uppsala University, 751 23 Uppsala, Sweden.
³Coasts and Oceans National Centre, National Institute of Water and Atmospheric Research,
Private Bag 14901, Kilbirnie, Wellington, New Zealand.
⁶Coasts and Oceans National Centre, National Institute of Water and Atmospheric Research,
P.O. Box 893, Nelson, New Zealand.
⁷Coral Reef Research Foundation, Box 1765, Koror, 96940 Palau.

*Corresponding authors: michelle.kelly@niwa.co.nz¹, paco.cardenas@ilk.uu.se²
³Email: Nicola.Rush@niwa.co.nz
⁴Email: carina@clearsight.co.nz
⁵Email: Diana.Macpherson@niwa.co.nz
⁶Email: Mike.Page@niwa.co.nz
⁷Email: crrfpalau@gmail.com

Abstract. Due to the possession of huge contort strongyles, and a lack of triaenes in an otherwise
‘astrophorine’ spicule complement, the phylogenetic position of the endemic, monospecific New Zealand
sponge genus, Lamellomorpha Bergquist, 1968, has remained enigmatic. The genus was established
within Jaspidae de Laubenfels, 1968 (in the abandoned order Epipolasida Sollas, 1888), but it was
not until 2002 that the genus was transferred formally to Astrophorina Sollas, 1887, albeit
incertae sedis, by Hooper & Maldonado (2002). In this study, we recognise specimens of Lamellomorpha from
the Subantarctic New Zealand region and Chatham Rise, considered by Bergquist to be conspecific
with the type species, L. strongylata Bergquist, 1968, first described from the Three Kings-Spirits Bay
region of Northland, as the new species, *L. australis* Kelly & Cárdenas sp. nov. These two species of *Lamellomorpha* have differences in external morphology and colour, skeletal architecture and spicules, natural products, geographical distribution, and depth ranges. Sequencing of the COI Folmer barcode/mini-barcode and of 28S (C1–C2 domains) of these two species suggests phylogenetic affinities of *Lamellomorpha* with the tetractinellid suborder Astrophorina and the family Vulcannellidae Cárdenas *et al.*, 2011. Two Subantarctic New Zealand species of the vulcannellid genus *Poecillastra* Sollas, 1888, *P. ducitriaena* Kelly & Cárdenas sp. nov. and *P. macquariensis* Kelly & Cárdenas sp. nov., provide further support for the close relationship of *Lamellomorpha* and *Poecillastra*.

**Keywords.** New species, *Poecillastra*, Porifera, secondary loss, Subantarctic New Zealand.


**Introduction**

*Lamellomorpha strongylata* Bergquist, 1968 (class Demospongiae Sollas, 1885, order Tetractinellida Marshall, 1876, suborder Astrophorina Sollas, 1887 *incertae sedis*) was first described from the Three Kings Islands to the north of New Zealand and recently recollected by the National Institute of Water & Atmospheric Research (NIWA) and the Coral Reef Research Foundation, Republic of Palau (CRRF) from Middlesex Bank to the north of Three Kings Islands and Spirits Bay on the northern tip of the North Island. In the original description, Bergquist (1968) included two specimens from the Campbell Plateau in the Subantarctic New Zealand region (New Zealand Oceanographic Institute (NZOI) Stations B176 and B184, 84 m and 188 m depth, respectively) and considered these to be the same species. While the specimens have not been relocated within the NIWA Invertebrate Collection (NIC), a recent opportunity to examine older material in NIC revealed a surprising number of the ‘southern form’ of *L. strongylata* Bergquist, 1968 from Bounty Platform (NZOI Station A751, 155 m depth), Solander Trough, Campbell Platform, and Macquarie Ridge, all in the Subantarctic New Zealand region, and Mernoo Bank on Chatham Rise, to the east of the South Island. One specimen from the Snares Island Platform was of great interest as it had rare calthrop-like triaenes amongst what appeared to be a spicule complement almost identical to that of *L. strongylata*, suggesting a relationship with calthrop-containing tetractinellid species, such as in families Calthropellidae Lendenfeld, 1907 or Pachastrellidae Carter, 1875.

The systematic position and phylogenetic affinity of *Lamellomorpha* Bergquist, 1968 has been the subject of some debate since it was first described. Bergquist (1968) established the genus within Jaspidae de Laubenfels, 1968 in the abandoned order Epipolasida Sollas, 1888, based on the possession of exclusively monaxon megascleres with asterose microscleres, i.e., lacking triaenes. Bergquist stated that “*L. strongylata* would be a typical *Jaspis* were it not for the completely different microsclere content of microstrongyles and streptasters”, which she likened to those in *Triptolemma simplex* (Sarà, 1959) in the Pachastrellidae.

In 2002, the genus was transferred from Jaspidae to Astrophorina Sollas, 1887 *incertae sedis* by Hooper & Maldonado (2002), who explored a plethora of hypotheses on the relationship of *Lamellomorpha* with Ancorinidae Schmidt, 1870, Pachastrellidae, ‘Lithistid demospongiae’, Hadromerida Topsent, 1894, and Halichondrida Gray, 1867. Bergquist’s initial thoughts were that *Lamellomorpha* was closely comparable to *Coppatias* (*Ecionemia*) *baculifera* (Kirkpatrick, 1903), due to the joint possession of microstrongyles, and to *Jaspis serpentina* Wilson, 1925 on the joint possession of microstrongyles and contort oxeas, and...
thus had affinities with species of *Jaspis* Gray, 1867 and *Coppatias* Sollas, 1888 (*Jaspis*) in the current family Ancorinidae. Hooper & Maldonado (2002) rejected Bergquist’s initial hypotheses on the basis that both species possess euaster microscleres (*Lamellomorpha* has streptasters), a relatively strong synapomorphy for Ancorinidae *sensu stricto*. Hooper & Maldonado (2002) also rejected Bergquist’s suggestion of possible affinity with *Triptolemma simplex*, and thus the family Pachastrellidae, because *Lamellomorpha* lacks the triaene megascleres that are used to judge the affinity with other genera. They also considered any similarity with the form, dimensions and disposition of the megascleres and microscleres to be artificial.

Kelly, in Cryer *et al.* (2000), tentatively assigned specimens of *L. strongylata* and ‘*Lamellomorpha n. sp. 1*’ (Kelly *et al.* 2009) to the desma-bearing, lithistid family Theonellidae Lendenfeld, 1903, after discussions with Professor Murray Munro, University of Canterbury, on the potential origins of the secondary metabolite chemistry of the specimens (Dumdei *et al.* 1997; Li *et al.* 1998; Hickford 2007): Munro’s group identified calyculins, calyculinamides and swinholide H (Dumdei *et al.* 1997), and cyclic peptolide theonellapeptolide IIIe (Li *et al.* 1998), compounds which are related to those found in *Discodermia calyx* Döderlein, 1884 and *Theonella swinhoei* Gray, 1868 suggesting a broad relationship with the lithistid family Theonellidae. Hooper & Maldonado (2002) also put forward the suggestion that *L. strongylata* might be a “lithistid demosponge” that has “lost” its desma megascleres. Indeed, several theonellid lithistids are known with only rudimentary desmas (Kelly-Borges *et al.* 1994: *Discodermia dissoluta* Schmidt, 1880, in the tropical Western Atlantic) or no desmas at all (Australian species *Theonella deliqua* Hall *et al.*, 2014 and *Theonella maricae* Hall *et al.*, 2014). However, the possession of extremely long, contort strongyles, not seen in other lithistids, and a lack of triaenes in *L. strongylata* precludes this assignment.

Cárdenas *et al.* (2011) suggested that *L. strongylata* might be phylogenetically close to *Characella* Sollas, 1886, or to *Pachastrella* Schmidt, 1868, based on the possession of small ectosomal monoaxial spicules. Cárdenas *et al.* (2011) and Cárdenas & Rapp (2012) concluded that Pachastrellidae sensu Maldonado (2002) was polyphyletic and reorganised the genera possessing streptasters into three families (Pachastrellidae; Theneidae Carter, 1883; Vulcanellidae Cárdenas *et al.*, 2011) and three *incertae sedis* groups (*Lamellomorpha incertae sedis, Characella incertae sedis, Neamphius* de Laubenfels, 1953 *incertae sedis*) within the Astrophorina.

Neither Cárdenas *et al.* (2011) nor Cárdenas & Rapp (2012) sampled *L. strongylata* for molecular sequences. Here, we sequence, for the first time, the COI Folmer barcode/mini-barcode and 28S (C1–C2 domains) of *L. strongylata* and the new species *L. australis*, and the second, less common Subantarctic New Zealand species, initially identified as a third species of *Lamellomorpha* with rare, calthrop-like triaenes, but now considered to be a species of *Poecillastra* Sollas, 1888, in the family Vulcanellidae: *Poecillastra ducitriaena* Kelly & Cárdenas sp. nov. The systematic and phylogenetic implications of these species are considered with respect to the broader phylogenetic position of *Lamellomorpha, Poecillastra*, and the wider Tetractinellida.

**Material and methods**

**Collections and morphological systematics**

Specimens were collected by rock and cone dredges as well as by beam and Agassiz medium trawls, from several research vessels between 1962 and 2010. The majority of specimens were collected onboard the National Institute of Water & Atmospheric Research (NIWA) research vessels RV *Tangaroa* and RV *Kaharoa*; numerical voyage identifier and associated stations cited as NIWA Stn TAN(voyage number)/(station number) and NIWA Station KAH(voyage number)/(station number), respectively.
Several specimens from the Three Kings Islands were collected by dredge from RV Kaharoa in 1999, on a voyage chartered by the CRRF.

Specimens were frozen immediately upon collection and then preserved in 70% ethanol or preserved immediately into 70% ethanol (CRRF). Histological sections of the sponges were prepared by embedding a small piece of the sponge in paraffin wax and then sectioning with a microtome at 70 μm. Spicule slides and SEM spicule preparations were made following the methods of Kelly & Sim-Smith (2012). Clean spicules for SEM examination were spread on a plastic disc, air-dried, and coated with platinum for 600 s. Spicules were viewed on a Philips XL30S FEG SEM. Spicule dimensions were measured using a Meji MT5300L compound microscope fitted with a Leica DFC420 microscope camera that was connected to Leica Application Suite imaging software (Leica Microsystems (Switzerland) Ltd.). Spicule measurements in the species descriptions are given as the mean length (range) × mean width (range) of twenty spicule measurements per specimen unless stated otherwise and are based on measurements from the holotype or paratypes and confirmed through examination of all other specimens. Collection information on specimens examined in this study and previous records were gathered in a table made available in the PANGAEA data repository (https://doi.pangaea.de/10.1594/PANGAEA.895370).

Primary and secondary type material of the new species, and additional material, are accessed within the NIWA Invertebrate Collection (NIC) at NIWA, Greta Point, Wellington, using the prefix ‘NIWA-’. Pieces of the holotypes of L. strongylata, L. australis sp. nov., and Poecillastra ducitriaena sp. nov. are also stored at the Zoological Museum in Uppsala, Sweden (prefix ‘UPSZTY-’). Additional abbreviations used in the text include CRRF (Coral Reef Research Foundation, Palau). The taxonomic authority for the new taxa described in this paper is restricted to the authors Michelle Kelly and Paco Cárdenas.

**Systematics**

General classification and the names of class, order, suborder, and family follow the classification proposal by Morrow & Cárdenas (2015). The systematics of the family Vulcanellidae follows Cárdenas et al. (2011). Terminology for the streptaster microscleres follows Sollas (1888), as used in Cárdenas & Rapp (2012): streptasters are categorised as spirasters (small, many actines, twisted, long shaft), metasters (intermediate morphology), plesiasters (large, few actines, short or disappearing shaft), and amphistasters (where actines radiate from both ends of a straight shaft).

**Molecular systematics**

DNA was extracted using a DNeasy Blood and Tissue kit (Qiagen). PCRs were carried out in 25 μl solutions using PuReTaq Ready-To-Go PCR beads (GE Healthcare). Due to poor preservation of the specimens for molecular work (storage in 70% ethanol, instead of the recommended 96% ethanol), the DNA quantities were very low, and it was very degraded (observation on a gel of 1 μl of DNA extract). The complete Folmer fragment could not be sequenced using the standard animal barcoding primer pair LCO1490/HCO2198 (Folmer et al. 1994) so it was sequenced in two parts: the universal minibarcode (130 bp, without primers) was obtained using the primer pair LCO1490/Tetract-minibarR1 (Cárdenas & Moore 2019). Then the second part of the Folmer fragment (539 bp, without primers) was amplified using the primer pair VulcanF2/HCO2198. VulcanCOI-F2 (5'-GGGGATGACCAACTTTATAATG-3') is a new specific primer made to amplify COI in Vulcanellidae species. PCR conditions were (5 min/94 °C; 37 cycles (15 s/94 °C, 15 s/46 ºC, 15 s/72 °C); 7 min/72 °C). The 28S fragment (C1–C2) of 308–369 bp, was obtained using the primer pair C1'/Ep3 (Chombard et al. 1998) and the same PCR program as for COI except that we used 50°C for the annealing temperature. We pruned the comprehensive Tetractinellida COI alignment from Kelly & Cárdenas (2016), to keep only species of Astrophytonra. We added additional sequences of Astrophytonra from the Galapagos (Schuster et al. 2018) along with
the new sequences. The COI data matrix included 115 sequences (with eight Spirophorina Bergquist & Hogg, 1969 outgroups). For 28S, we built an alignment based upon the Astrophorina 28S (C1–D2) alignment from Cárdenas et al. (2011) and added Astrophorina 28S (C1–D2) sequences (Thacker et al. 2013; Schuster et al. 2015, 2018). The 28S data matrix included 126 sequences (with seven Spirophorina outgroups) and was automatically aligned using MAFFT v.7 (Katoh & Standley 2013), L-INS-i option, implemented in AliView 1.18 (Larsson 2014). Phylogenetic analyses were conducted on the CIPRES science gateway v. 3.3 (http://www.phylo.org) (Miller et al. 2010): RAxML 8.2.10 (Stamatakis 2014) for maximum likelihood (ML) and MrBayes v. 3.2.6 (Ronquist et al. 2012) for Bayesian analyses. Bayesian analyses were run with BEAGLE, and consisted of two runs of four chains, each for 5 000 000 generations and sampled every 1000 tree after a 25% burn-in.

**Results**

Class Demospongiae Sollas, 1885  
Order Tetractinellida Marshall, 1876  
Suborder Astrophorina Sollas, 1887

**Family** **Vulcanellidae** Cárdenas, Xavier, Reveillaud, Schander & Rapp, 2011

**Diagnosis** (modified from Cárdenas et al. 2011)
Astrophorina with calthrops, short-shafted triaenes or long-shafted triaenes, in addition to large oxeas and contort or sinuous strongyloxeas. Aster microscleres include several categories of streptasters (spirasters, metasters, amphiasters, and plesiasters). Monaxonic spicules consist of one to three categories of spiny microxeas or microstrongyles.

**Lamellomorpha** Bergquist, 1968


**Diagnosis** (modified from Hooper & Maldonado 2002)
Massive, lamellate stalked-palmate, or paddle-shaped sponges, with a relatively smooth, granular, or fleshy, slightly conulose surface. Ectosomal skeleton a skin-like membrane packed with microstrongyles. Choanosomal skeletal architecture a core of megascleres, which are straight, curved, sinuous, or contort oxeas, frequently modified with one or both ends rounded as in strongyloxeas. These radiate through the stalk and fan. Straight oxeas arise as short subectosomal tracts that emerge oblique to the surface. Roughened microstrongyles or microxeas and streptasters (amphiasters, metasters, and spirasters) scattered throughout the body.

**Type species**

*Lamellomorpha strongylata* Bergquist, 1968 (by monotypy).

*Lamellomorpha strongylata* Bergquist, 1968  
Figs 1–2, 6; Tables 1–2

‘Lamellomorpha n. sp. K & W’ in Cryer et al. 2000: 42 (NIWA 51169 leg.).
Type material

Holotype
NEW ZEALAND • Northeast of Three Kings Islands, NZOI Station B93; 33.983° S, 172.350° E; depth 54–109 m; 22 Oct. 1958; NIWA 356 (NZOI H–33) leg.; beam trawl; UPSZTY 178600 (a piece of the holotype preserved in 70% ethanol, as well as a spicule preparation), NIWA.

Other material examined
NEW ZEALAND – Northeast of Three Kings Islands, NIWA Station Z9678 (KAH9901/27); 34.360° S, 172.712° E; depth 48 m; 26 Jan. 1999; NIWA 51169 and 51172 leg.; UPSZMC 178601 (fragment of NIWA 51172 leg. preserved in 70% ethanol), NIWA • Northeast of Three Kings Islands, NIWA Station Z9686 (KAH9901/43); 34.361° S, 172.686° E; depth 48 m; 27 Jan. 1999; NIWA 51267 leg.; UPSZMC 178603 (fragment preserved in 70% ethanol), NIWA • Northeast of Three Kings Islands, NIWA Station Z9699 (KAH9901/67); 34.360° S, 172.673° E; depth 41 m; 28 Jan. 1999; NIWA 51438 leg.; NIWA • Northeast of Three Kings Islands, NIWA Station Z9710 (KAH9901/85); 34.353° S, 172.765° E; depth 54 m; 28 Jan. 1999; NIWA 51582 leg.; dredge; NIWA • Three Kings Islands, 2.5 nm east of Great Island, NIWA Station Z15944; 34.170° S, 172.210° E; depth 200 m; 16 Apr. 1999; CRRF, NIWA 93474 leg.; dredge; NIWA • Spirits Bay, Northland, NIWA Station KAH0606/D3; 34.36° S, 172.847° E; 15

Fig. 1. Study area showing the collection localities for: *Lamellomorpha strongylata* Bergquist, 1968, around Middlesex Bank, Three Kings Islands, and Spirits Bay (triangles); *L. australis* Kelly & Cárdenas sp. nov., Bounty Platform, Campbell Plateau, Solander Trough, and Macquarie Ridge (Australian EEZ) (circles); *Poecillastra ducitriaena* Kelly & Cárdenas sp. nov., on the eastern edge of the Snares Platform (square); and *Poecillastra macquariensis* Kelly & Cárdenas sp. nov., on Seamount 5, Macquarie Ridge (star).
Fig. 2. Morphology, megascleres and microscleres of *Lamellomorpha strongylata* Bergquist, 1968. A. Deck image showing midnight blue colouration and palmate morphology (NIWA 93474 leg., image Lori J. Bell, CRRF). B. Loss of colouration and radiating skeleton (NIWA 73253 leg.). C. Range of roughened microstrongyles showing centrotylote forms and more typical curved forms (penultimate: NIWA 356 leg., rest: NIWA 93474 leg.). D. Sinuous oxeas (NIWA 93474 leg.). E. Metaster- to amphiaater-like streptasters (left: NIWA 93474 leg., center and right: NIWA 356 leg.); F. Metaster- to amphiaater-like streptasters (left and center: NIWA 356 leg., right: NIWA 93474 leg.).
Table 1. Megascleres and dimensions (μm) of Lamellomorpha strongylata Bergquist, 1968 and L. australis Kelly & Cárdenas sp. nov., given as length [mean (min.–max.)] × width [mean (min.–max.)], n = 10–17 unless stated otherwise.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Straight megascleres</th>
<th>Contort megascleres</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>L. strongylata</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NZOI Holotype 33 (Bergquist 1968)</td>
<td>1980(1000–2808) × 26(14–33)</td>
<td></td>
</tr>
<tr>
<td>NIWA 00356 (NZOI Holotype 33)</td>
<td>1993(1520–2416) × 27(21–31)</td>
<td>1556(1306–1787) × 17(11–24)</td>
</tr>
<tr>
<td>NIWA 93474</td>
<td>1968(1373–2552) × 22(9–30)</td>
<td>1564(1288–1816) × 15(8–21)</td>
</tr>
<tr>
<td><strong>L. australis</strong> sp. nov.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NZOI Station B176 (Bergquist 1968)</td>
<td>1482(1161–1937) × 23(17–28)</td>
<td></td>
</tr>
<tr>
<td>NIWA 89736 leg. (holotype)</td>
<td>1655(1130–1981) × 26(17–41)</td>
<td>not present</td>
</tr>
<tr>
<td>NIWA 93483 leg. (paratype)</td>
<td>1754(1454–2223) × 26(15–36)</td>
<td>2159(1499–3079) × 25(14–31)</td>
</tr>
<tr>
<td>NIWA 93484 leg. (paratype)</td>
<td>1562(1403–1988) × 21(13–27)</td>
<td>2657(2243–3058) × 23(17–33)</td>
</tr>
<tr>
<td>NIWA 93485 leg. (paratype)</td>
<td>1640(1044–2026) × 19(6–25)</td>
<td>3164(2672–3555) × 23(18–27)</td>
</tr>
<tr>
<td>NIWA 93486 leg. (paratype)</td>
<td>1725(1231–2320) × 19(13–28)</td>
<td>2257(1702–3575) × 20(13–32)</td>
</tr>
<tr>
<td>NIWA 93487 leg. (paratype)</td>
<td>1752(1226–2613) × 24(16–31)</td>
<td>2195(1658–3560) × 23(17–34)</td>
</tr>
</tbody>
</table>

May 2005; NIWA 52375 leg.; dredge; NIWA. • Middlesex Bank, Three Kings Rise, NIWA Station TAN1105/43; 33.988° S, 171.751° E; depth 170–174 m; 28 Mar. 2011; NIWA 73243, 73253 leg.; beam trawl; NIWA • Western Continental Slope, Northland, NZOI Station J954 (I808); 34.633° S, 172.225° E; depth 204–192 m; 18 Jun. 1981; collected by rock dredge; specimen now lost, donated by Dame P. R. Bergquist to Dr P. Karuso, Macquarie University, Sydney.

**Description**

The holotype was described by Bergquist (1968) as a “massive, thick, sometimes folded and incurved lamellate sponge”, 130 mm high, 102 mm wide, and 18–22 mm thick, supported by a stout stalk 30 mm in diameter. The surface was described as smooth where the dermal membrane was intact, otherwise ragged due to projecting clumps of oxeas and strongyles. Oscules, 1–2.6 mm in diameter, were found on the convex surface of the lamella and lie flush with the surface (Bergquist 1968). Examination of the numerous preserved specimens in NIC reveal occasional membranous oscules, but it is difficult to tell whether they are restricted to one side of the sponge. However, in the holotype, pores were observed on the opposite side to the oscules, in cribriporal areas, separated by small ridges, or with no boundaries, making a continuous pore surface; each pore is 40–80 μm in diameter. The texture was described as, “firm but compressible, crisp, easily broken”. The colour in life was described as “bright green” and the colour in spirit, “blue green or yellowish green” (Bergquist 1968). The most recent collection was by the Coral Reef Research Foundation in 1999 (NIWA 93474 leg.; Fig. 2A), who described a “dark, royal blue (not navy blue), (palmate) fan sponge with pointed tips, 20 cm high and about 1 cm thick, that tears easily, and which has a fleshy surface”.

**Skeleton**

The description by Bergquist (1968) of the choanosome as “lax and confused with slight traces of radiate construction discernible”, is accurate, but in NIWA 93474 leg. the contort strongyles strongly radiate through the plane of the fan. Bergquist described a “subectosomal region”, in which there were tracts of megascleres, variable in thickness, that curved outward and intersect with the surface at an acute angle; in NIWA 93474 leg. these are predominantly oxeas (Fig. 6A). The ectosome is densely packed with microstrongyles and streptasters, which also occur throughout the sponge, but in much less abundance.
KELLY M. et al., Revision of Lamellomorpha

Table 2. Microscleres and dimensions (μm) of Lamellomorpha strongylata Bergquist, 1968 and L. australis Kelly & Cárdenas sp. nov., given as length [mean (min.–max.)] × width [mean (min.–max.)], n = 10–17 unless stated otherwise.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Microstrongyles</th>
<th>Streptasters</th>
<th>Spirasters</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>L. strongylata</em> Bergquist, 1968</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NZOI Holotype 33 (Bergquist 1968)</td>
<td>25(23–28) × 3(2–4)</td>
<td>10(8–11)</td>
<td>not present</td>
</tr>
<tr>
<td>NIWA 00356 (NZOI Holotype 33)</td>
<td>23(21–28) × 3(2–4)</td>
<td>10(8–15)</td>
<td>not present</td>
</tr>
<tr>
<td>NIWA 93474</td>
<td>27(24–34) × 2(2–3)</td>
<td>9(7–12)</td>
<td>not present</td>
</tr>
<tr>
<td><em>L. australis</em> sp. nov.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NZOI Station B176 (Bergquist 1968)</td>
<td>27(24–30) × 3(2–4)</td>
<td>10(8–11)</td>
<td>not given</td>
</tr>
<tr>
<td>NIWA 89736 leg. (holotype)</td>
<td>37(31–42) × 3(2–4)</td>
<td>12(10–13)</td>
<td>10(8–11)</td>
</tr>
<tr>
<td>NIWA 93483 leg. (paratype)</td>
<td>32(25–41) × 2(2–4)</td>
<td>10(8–12)</td>
<td>10(9–15)</td>
</tr>
<tr>
<td>NIWA 93484 leg. (paratype)</td>
<td>32(29–37) × 2(1–3)</td>
<td>8(6–8)</td>
<td>10(8–12)</td>
</tr>
<tr>
<td>NIWA 93485 leg. (paratype)</td>
<td>28(21–38) × 3(2–5)</td>
<td>10(9–12)</td>
<td>11(9–14)</td>
</tr>
<tr>
<td>NIWA 93486 leg. (paratype)</td>
<td>36(31–40) × 2(2–3)</td>
<td>9(7–11)</td>
<td>11(9–13)</td>
</tr>
<tr>
<td>NIWA 93487 leg. (paratype)</td>
<td>27(19–34) × 3(1–5)</td>
<td>9(8–12)</td>
<td>10(8–14)</td>
</tr>
<tr>
<td>NIWA 93499 leg.</td>
<td>32(26–41) × 2(1–2)</td>
<td>9(8–15)</td>
<td>10(8–12)</td>
</tr>
</tbody>
</table>

Spicules

Megascleres (Table 1; Fig. 2D)

Bergquist (1968) considered the megascleres of *L. strongylata* (Bergquist 1968: 31, 32 (table of spicule dimensions)) to be “strongyles, oxeas and strongyloxeas”, all of similar range in length and width, varying only in relative frequency in the two specimens (presumably the Three Kings holotype and the NZOI Station B176 specimen from Campbell Plateau), with oxeas being dominant in the latter. Re-examination of the holotype megascleres, and those of more recent collections, indicate that there are probably two forms of megascleres: 1) straight to slightly curved oxeas that are common in the subectosomal tracts, ranging from about 1500–1750 μm long and up to 25 μm thick; and 2) massive sinuous or contort oxeas that are usually very thick and frequently modified with one or both ends rounded as in strongyloxeas, rarely as in true strongyles, ranging from about 1600–2375 μm long and up to 40 μm thick. However, it is difficult to distinguish the various megascleres in some specimens, and in some the spicules are much less contort.

Microscleres (Table 2; Fig. 2C–F)

Microstrongyles are “squat, evenly rounded spicules, slightly roughened and occasionally centrotylote” (Bergquist 1968: 31, 32 (table of spicule dimensions)) and range from about 21–34 μm long (Table 2). Bergquist described the streptaster microscleres of *L. strongylata* (Bergquist 1968: 31, 32 (table of spicule dimensions)) as “plesiasters, small spicules with 3–12 smooth, sharply pointed rays”. A re-examination of the holotype (Fig. 2E–F) using scanning electron microscopy has revealed that the streptaster microscleres are metasters and occasionally amphiasters with relatively long microspined rays, all in one size category, following the definition of Sollas (1888), and as used in Cárdenas & Rapp (2012). We describe these spicules as metaster- to amphiaster-like streptasters with heavily spined, relatively long rays in one size category, ranging in length from about 7–15 μm long (Table 2).

Distribution

Northeast of New Zealand.

Substrate, depth range and ecology

Attached to rocky reefs and sediment and rubble-covered rocky platforms, depth 41–200 m.
DNA barcodes

COI. NIWA 51172 leg. (minibarcode, MK033624) and NIWA 51267 leg. (MK033623): no bp differences. 28S (C1-C2). NIWA 51172 leg. (MK033143) and NIWA 51267 leg. (MK033142): no bp differences. We failed to get sequences from the holotype.

Remarks

*Lamellomorpha strongylata* was originally described in considerable detail by Bergquist (1968), and the holotype was redescribed without re-examination more recently by Hooper & Maldonado (2002). No further material was examined. Here, for the first time, we illustrate the sponge as it appears upon collection, showing the beautiful royal blue colouration (Fig. 2A), and illustrate the detail and ornamentation of the microscleres (Fig. 2C–F) using scanning electron microscopy. There is little to add to the original description, consequently the description and skeletal details are provided in comparative prose. *Lamellomorpha strongylata* is restricted to the northernmost tip of New Zealand and beyond to the Three Kings Rise, and is easily recognised in the field by the palmate, tree-like shape and the deep blue to green colouration.

*Lamellomorpha australis* Kelly & Cárdenas sp. nov.

urn:lsid:zoobank.org:act:2A7459B3-0FBA-441B-806C-7C988075843A

Figs 1, 3, 6; Tables 1–2


Etymology

Named for the Chatham Rise and Subantarctic New Zealand distribution of this species (*australis*, south, Latin).

Type material

Holotype

NEW ZEALAND • Subantarctic region of New Zealand, Bounty Platform, NZOI Station A751; 47.743° S, 179.124° E; depth 155 m; 16 Nov. 1962; NIWA 89736 leg.; Agassiz medium trawl; UPSZTY 178605 (a small piece of the holotype preserved in 70% ethanol, as well as a spicule preparation), NIWA.

Paratypes

NEW ZEALAND – same collection data as for the holotype; NIWA 92896 to 92900, 93483, 93484, and 93487 leg.; NIWA • same collection data as for preceding; NIWA 93485 leg.; UPSZTY 178606 (a small piece of the paratype preserved in 70% ethanol), NIWA • same collection data as for preceding; NIWA 93486 leg.; UPSZTY 178607 (a small piece of the paratype preserved in 70% ethanol), NIWA.

Type locality

Subantarctic region of New Zealand, Bounty Platform; depth 155 m.

Other material examined

NEW ZEALAND • Bounty Platform, NZOI Station 1711; 47.833° S, 179.250° E; depth 139 m; 22 Mar. 1979; NIWA 89717 leg.; rock dredge; NIWA • Bounty Platform, NZOI Station A714; 47.725° S, 179.067° E; depth 165 m; 5 Nov. 1962; NIWA 86733 leg.; cone and mesh dredge; NIWA • Bounty Platform, NZOI Station A715; 47.683° S, 179.051° E; depth 121 m; 5 Nov. 1962; NIWA 89720 leg.;
Fig. 3. Morphology, megascleres, and microscleres of *Lamellomorpha australis* Kelly & Cárdenas sp. nov. 

A. Bilamellate form (NIWA 89736 leg., holotype). 

B. Club-shaped morphology showing both sides (NIWA 93483 leg., paratype). 

C. Sinuous oxeas (NIWA 44388 leg.). 

D. Curved, roughened microstrongyles (rounded ends) and microxeas (blunt-pointed ends) (NIWA 44388 leg.). 

E. Metaster-like streptasters with heavily spined, relatively long rays (NIWA 89736 leg. and NIWA 89717 leg.). 

F. Spirasters with dense, short, microspined rays (NIWA 44388 leg. and NIWA 89736 leg.).
cone and mesh dredge; NIWA • Bounty Platform, NZOI Station A757; 47.693° S, 179.058° E; 17 Nov. 1962; NIWA 113894 leg.; NIWA • Solander Trough, NZOI Station D39; 50.967° S, 165.750° E; depth 549 m; 7 May 1963; NIWA 44388 leg.; gear dredge, cone mesh with bag; NIWA • Campbell Platform, NZOI Station B184; 52.615° S, 169.117° E, depth 549 m; 7 May 1963; NIWA 44399 leg.; dredge; NIWA • Macquarie Ridge, NZOI Station TAN0803/69; 52.398° S, 160.657° E; depth 451–438 m; 9 Apr. 2008; NIWA 40328 leg.; epibenthic sled; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W427; 43.077° S, 175.272° E; depth 180–237 m; 20 Feb 1995; NIWA 44240 leg.; Agassiz Trawl; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W446; 43.245° S, 175.458° E; depth 71–76 m; 22 Feb 1995; NIWA 44261 leg.; rock dredge; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W447; 43.245° S, 175.458° E; depth 80–85 m; 22 Feb 1995; NIWA 44263 leg.; rock dredge; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W448; 43.240° S, 175.458° E; depth 74 m; 22 Feb 1995; NIWA 44267 leg.; rock dredge; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W446; 43.247° S, 175.444° E; depth 71–76 m; 22 Feb 1995; NIWA 44272 leg.; rock dredge; NIWA • Chatham Rise, North Mernoo Bank, NIWA Station W435; 43.172° S, 175.340° E; depth 108–113 m; 20 Feb 1995; NIWA 137201 leg.; rock dredge; NIWA • Chatham Rise, South Mernoo Bank, NIWA Station W452; 43.451° S, 175.130° E; depth 126–130 m; 22 Feb 1995; NIWA 44272 leg.; rock dredge; NIWA • Chatham Rise, South Mernoo Bank, NIWA Station W454; 43.451° S, 175.109° E; depth 126–130 m; 22 Feb 1995; NIWA 44262 leg.; rock dredge; NIWA.

Description

Uni- to bilamellate fan sponge, table tennis bat-shaped (Fig. 3A) or club-shaped (Fig. 3B), 130–200 mm high with a short, broad stalk, 2–3 cm thick, and a relatively thick lamella (up to 2 cm thick in places), attenuating towards the margins, which are frequently incised. The specimen from NZOI Station B176 was described by Bergquist (1968) as being 160 mm high, 89 mm wide, and 32–58 mm thick, supported by a stalk that was broken and thus was not measured. Oscules were not visible in the holotype or any other specimen. Pores are inconspicuous and compressed (probably due to the fixation) and were 40–80 μm in diameter (measured on the holotype). Surface relatively smooth with low ridges radiating from the stalk to the fan margins. Texture, relatively soft, compressible. Colour in life and preservative, tan.

Skeleton

Choanosome disorganised, with megascleres orientated more or less parallel with the axis of the fan and stalk (Fig. 6B–D), with single or a couple of megascleres extending beyond the surface from the subectosome. The ectsosome is extremely thick and packed with microstrongyles and streptasters, which also occur in great density throughout the sponge.

Spicules

Megascleres (Table 1; Fig. 3C)

Bergquist (1968) considered the megascleres of the “subantarctic specimen” (presumably the NZOI Station B176 specimen from Campbell Plateau) to have, “predominantly oxeas, some of which are curved, but most are contort”. Our examination of new material reveals that oxeas dominate the megasclere complement; these are rarely to never modified; all have sharp attenuated tips. The majority are straight to slightly curved and contort, but not to the degree seen in L. strongylata. The megascleres reach their greatest length in L. australis sp. nov., up to 3575 μm long in the specimen NIWA 93486 leg. (paratype).

Microscleres (Table 2; Fig. 3D–F)

Bergquist did not differentiate between the microscleres of the holotype of L. strongylata (from the Three Kings) and the subantarctic Campbell Plateau specimens (L. australis sp. nov.), calling them all “plesiasters” in the table of spicule dimensions. However, in pl. 11, figs E2 and F2–3, a clear difference is
obvious between the illustrations of the streptasters: they are metasters in pl. 11, fig. E2 (*L. strongylata*) and larger metasters (pl. 11, fig. F3) and “abnormal microrhabds” in pl. 11, fig. F2 (*L. australis* sp. nov.). The “abnormal microrhabds” of Bergquist (1968: pl. 11, fig. F2) are most likely spirasters (as in our Fig. 3F), the ornamentation of which would not have been visible under light microscopy available at the time.

Thus, *L. australis* sp. nov. has three forms of microsclere: a microxea (Fig. 3D) with attenuating, hastate ends that is usually straight, but may be slightly curved, ranging in length from about 19–40 μm; metaster-like streptasters with heavily spined, relatively long rays (Fig. 3E), ranging in length from about 7–15 μm; spirasters with abundant, short, microspined rays that emanate from a long, spiral axis (Fig. 3F), ranging in length from about 8–14 μm.

**Distribution**

Subantarctic region of New Zealand: Mernoo Bank (depth 71–237 m), north-western Chatham Rise (Dumdei *et al.* 1997; Li *et al.* 1998); Bounty Platform (depth 121–165 m), Solander Trough (depth 549 m), Campbell Platform (depth 344 m), and Macquarie Ridge (depth 451–438 m).

**Substrate, depth range and ecology**

Attached by a thick stalk to sediment covered rocky substrate, depth 71–549 m.

**DNA barcodes**

COI. Holotype (minibarcode, MK033625), no bp differences with the COI minibarcode of *L. strongylata*.

**Remarks**

Bergquist (1968) listed two specimens from Campbell Plateau in the Subantarctic region of New Zealand, from NZOI Station B176 (46 fathoms = 84.12 m) and NZOI Station B184 (103 fathoms = 188.4 m). Unfortunately, neither specimen was found in the NIWA collections and both are presumed lost. However, we did find a specimen from NZOI Station B184 (NIWA 93499 leg.) from a depth of 344 m. Bergquist (1968) considered the two specimens she examined to be conspecific with *L. strongylata*, despite the obvious disjunct distribution, but noted that the subantarctic specimens had predominantly oxeas, an observation we agree with.

Examination of numerous specimens uncovered in NIC allows us to convincingly separate *L. australis* sp. nov. from the type species on geographic distribution, morphology, and skeletal details, despite the COI minibarcodes not differentiating them (Fig. 7). The most obvious difference that separates *L. australis* sp. nov. from *L. strongylata* is the markedly disjunct distribution and depth ranges: *L. strongylata* has only been recorded to the north of New Zealand, 41–200 m depth, while *L. australis* sp. nov. is only found on and south of Mernoo Bank on the Chatham Rise, ranging in depth from 71 m on Mernoo Bank, to 549 m in the Solander Trough. In terms of morphology and colouration in life, *L. strongylata* forms a relatively soft, dark royal blue, palmate sponge, supported by a relatively narrow stalk, while *L. australis* sp. nov. forms a distinctive, tan, paddle-shaped sponge, with thin, incised margins, on a thick, short stalk. In terms of skeletal architecture, the choanosome of *L. australis* sp. nov. is much more densely packed with microscleres than *L. strongylata*, and the former species lacks the relatively distinct subectosomal tracts of the latter. As noted by Bergquist (1968), the megascleres of *L. australis* sp. nov. differ from those of *L. strongylata* in being predominantly oxeas (straight and contort) with no modifications of the tips to strongyloxeas as in *L. strongylata*. In addition, we note that the contort forms are much longer on average, and have a greater size range, than in *L. strongylata*. Finally, microscleres also discriminate *L. australis* sp. nov. from *L. strongylata*. *Lamellomorpha strongylata* has stubby, often centrotylote roughened microstrongyle, while *L. australis* sp. nov. has a relatively fine, curved, slightly
longer roughened microxea. *Lamellomorpha strongylata* has metaster- to amphiaster-like streptasters with heavily spined, relatively long rays in one size category, while *L. australis* sp. nov. has metaster-like streptasters with heavily spined, relatively long rays and spirasters with abundant, short, microspined rays that emanate from a long, spiral axis. Spirasters are absent in *L. strongylata*.

As part of their ongoing investigations into New Zealand marine natural products in sponges, professors Murray Munro and John Blunt and their group in the Department of Chemistry, University of Canterbury, Christchurch, collected what was identified by the late professor Patricia Bergquist as *L. strongylata*, from Mernoo Bank on the Chatham Rise. Vouchers of these sponge specimens were donated to NIC for their preservation and future study and have been re-identified here as *L. australis* sp. nov., extending the known distribution of *L. australis* sp. nov. north to the Chatham Rise. Thus, it is *L. australis* sp. nov., and not *L. strongylata*, from which biologically active secondary metabolites were isolated by the University of Canterbury group, including calyculins (A, B, E, and F), calyculinamides (A and B), swinholide H (Dumdei et al. 1997), and theonellapeptolides (Li et al. 1998; Hickford 2007); identical and related compounds are found in sponges in the genus *Theonella* Gray, 1868 and *Discodermia* du Bocage, 1869 (family Theonellidae).

It has been shown that calyculins and its derivatives (e.g., calyculinamides) could be produced by the filamentous bacteria ‘*Entotheonella*’ spp. in *Discodermia* (Wakimoto et al. 2014). ‘*Entotheonella*’ spp. are especially abundant in *Theonella swinhoei*, as well as in many other demosponges (Wilson et al. 2014). In her PhD thesis, Hickford (2007) noticed that filamentous heterotrophic (Gram positive) bacteria were very abundant in *L. australis* sp. nov. and were associated with several theonellapeptolides. The producer of theonellapeptolides is currently unknown but the results of Hickford (2007) suggest that *L. australis* sp. nov. may be a host for theonellapeptolides-producing ‘*Entotheonella*’-like bacteria. Hickford (2007) also isolated unicellular bacteria from the same specimens and showed these were associated with swinholide H. This result concurs with previous results from Bewley et al. (1996), who identified swinholide A in unicellular bacteria isolates from *Theonella swinhoei* from Palau. However, it is an apparent contradiction with Ueoka et al. (2015) who convincingly show that misakinolide A (swinholide-like compound) from another *Theonella swinhoei* chemotype (chemotype WA from Japan) is produced by ‘*Entotheonella serta*’. Therefore it seems that swinholide-type compounds may be produced by bacteria other than ‘*Entotheonella*’ in *L. australis* sp. nov. and *Theonella swinhoei* (chemotype Palau). Hickford (2007) further states that specimens from the “northern population of *L. strongylata*” (*L. strongylata*) not only had very low quantities of filamentous bacteria (apparently limited to the surface of the sponge), but also missed the biological activity, and therefore may not produce the above mentioned compounds. Thus, to conclude, *L. strongylata* and *L. australis* sp. nov. also clearly differ in terms of natural products and microbial communities.

*Poecillastra* Sollas, 1888

*Poecillastra* Sollas, 1888: 105.

**Diagnosis** (Cárdenas et al. 2011)

Vulcanellidae with spiny microxeas in a single category, triaenes are pseudocaltrops or short-shafted triaenes.

**Type species**

*Poecillastra compressa* (Bowerbank, 1866: 55) (by original designation).
Poecillastra ducitriaena Kelly & Cárdenas sp. nov.  
urn:lsid:zoobank.org:act:E3C570FA-832A-4FB9-97D3-5CDA250A3797  
Figs 1, 4, 6

Etymology
Named for the possession of triaenes in addition to the apparent spiculation of Lamellomorpha, and their guide to the phylogenetic origins of this species (‘duct-’ in the sense of a guide).

Type material
Holotype
NEW ZEALAND • Subantarctic region of New Zealand, east of Snares Island Platform, NIWA Station TRIP3072/8; 48.5° S, 168.0° E; depth 125–213 m; 21 Oct. 2010; NIWA 61944 leg.; fish bottom trawl; UPSZTY 178604 (a small piece of the holotype preserved in 70% ethanol, as well as a spicule preparation), NIWA.

Type locality
Subantarctic region of New Zealand, Snares Island Platform, depth 125–213 m.

Description
Multilamellate, foliose, fan sponge (Fig. 4A), 160 mm high, 104 mm wide, with a short thick stalk about 2 cm thick. Lamella up to 2 cm thick in places, attenuating to curled margins. Oscules were not visible on the holotype. Cribriporal pore areas are widespread between parallel tangential tracts of oxeas; individual pores 80–160 μm in diameter, on both sides of the lamella. Texture firm, compressible, flexible, granular and smooth to the touch. Colour in preservative tan.

Skeleton
Choanosome composed of huge swathes of long straight oxeas, radiating through the lamella, terminating below the surface (Fig. 6E). Contort oxeas are found in the stalk. Ectosome, relatively thick, packed with microxeas and perforated by ostia.

Spicules
Megascleres (Fig. 4B–C)
Oxeas with long fine attenuated tips, 1725(1150–2271) × 16(8–22) μm; contort oxeas in the stalk, 1801(1095–2671) × 13(5–17) μm; short-shafted triaenes, relatively uncommon, rhabd straight, attenuating, 290(260–300) μm, clads curved or acutely bent, occasionally with bifurcating tips, 232(200–250) μm, cladome width 400–500 μm long.

Microscleres (Fig. 4D–F)
Microxeas, heavily microspined, sometimes faintly centrotylote and acutely centrally bent, sharp ended, abundant, 38(28–46) × 3(2–4) μm; metaster- to amphiaister-like streptasters with long, microspined rays, rare, 8(5–12) μm long; spirasters with dense, short rays, rare, 5–7 μm long.

Distribution
East of Snares Island Platform.

Substrate, depth range and ecology
Attached by a thick stalk to sediment covered rocky substrate, depth 125–213 m.
Fig. 4. Morphology, megascleres and microscleres of *Poecillastra ducitriaena* Kelly & Cárdenas sp. nov. (NIWA 61944 leg., holotype). **A.** Preserved holotype showing stalked morphology (both sides). **B.** Short-shafted triaenes with curved clads and short rhabd. **C.** Sinuous and straight oxeas. **D.** Microspined microxeas, sometimes faintly centrotylote, sharp ended. **E.** Metaster- to amphiaster-like streptasters with long, microspined rays. **F.** Spirasters with dense, short rays.
DNA barcodes
COI. Holotype (MK033626); 28 bp difference with the COI of *L. strongylata*; 24 bp difference with the COI of *Poecillastra compressa* (Bowerbank, 1866) (HM592675). 28S (C1-C2). Holotype (MK033144); 5 bp difference with the 28S (C1-C2) of *L. strongylata*; 3 bp difference with the 28S (C1-C2) of *Poecillastra compressa* (HM592757).

Remarks
This remarkable sponge was first identified as a third species of *Lamellomorpha*, as it appeared to have an almost identical form (stalked, multilamellar fan), a megasclere complement of straight and contort oxeas (more or less restricted to the stalk), small centrotylote microxeas, and metasters (albeit rare). Because the short-shafted triaenes were relatively uncommon, it was initially hypothesised that this was a species of *Lamellomorpha* with rudimentary triaenes that ‘showed the way’ to the true affinity of the genus with other triaene-bearing Tetractinellida. However, molecular sequencing consistently linked *Poecillastra ducitriaena* sp. nov. with other *Poecillastra* species (Fig. 7). Despite its consistency with two independent markers, we note that this grouping is not supported (bootstrap of 60 for COI, of 10 with 28S). This may be due to the absence of other subantarctic *Poecillastra* species in our sampling which *Poecillastra ducitriaena* sp. nov. may be closer to (P. Cárdenas, unpublished results).

Although not fully documented (Kelly *et al.* 2009), our knowledge of *Poecillastra* in the New Zealand region is reasonable and includes what we consider to be *Poecillastra laminaris* (Sollas, 1886) (Zeng *et al.* 2016) and *Poecillastra schulzei* (Sollas, 1886). While several undescribed species are known from the New Zealand EEZ, no specimens are known that contain the characteristic contort oxeas of *Poecillastra ducitriaena* sp. nov.

*Poecillastra macquariensis* Kelly & Cárdenas sp. nov.

urn:lsid:zoobank.org:act:78B12A37-93E0-4CA9-9317-81F835B112FE

Fig. 5

Etymology
Named for the type location of this species, the Macquarie Ridge.

Type material

**Holotype**

NEW ZEALAND • Subantarctic region of New Zealand, Seamount 5, Macquarie Ridge, NIWA Station TAN0803/48; 51.096° S, 161.976° E; depth 462–524 m; 4 Apr. 2008; NIWA 52640 leg.; epibenthic sled; NIWA.

Description

Solid stalk of sponge of unknown morphology, 15 mm in diameter, 20 mm high, expanding on the broken, upper surface, sides of stalk sculpted, attachment base contains patches of substrate (Fig. 5A). Surface hispid and scratchy to the touch; texture firm, incompressible. Colour in preservative tan.

**Skeleton**

Stalk composed of huge swathes of contort oxeas and triaenes between which are abundant microscleres.

**Spicules**

*MEGASCLERES* (Fig. 5D–E)

Abundant contort to sinuous oxeas (Fig. 5D) with slightly rounded tips, 3725(2125–5750) × 53(30–70) μm; medium-shafted triaenes (Fig. 5E), rhabd slightly curved, tapering to a sharp tip, 852(550–1225) μm,
Fig. 5. Morphology, megascleres and microscleres of *Poecillastra macquariensis* Kelly & Cárdenas sp. nov. (NIWA 52640 leg., holotype). A. Stalk, showing smooth attachment base with specks of remaining substrate, and broken edge of the upper part of the sponge. B. Microxeas. C. Detail of microxea showing roughened surface. D. Contort oxea of choanosome. E. Range of triaenes with medium to short shafts (calthrops). F. Plesiasters. G. Spiraster to metaster-like streptasters.
Fig. 6. Skeletal arrangement. A. *Lamellomorpha australis* Kelly & Cárdenas sp. nov. showing dense, crustose ectosome packed with microrhabds and other microscleres, a rough megasclere bundle in the subectosome, emerging perpendicular to the surface, and the choanosome densely packed with microscleres (NIWA 89736 leg., holotype). B. *L. australis* sp. nov. showing the subectosome densely packed with microscleres, and megascleres radiating in the plane of the sponge body (NIWA 89717 leg.). C. *L. strongylata* Bergquist, 1968 showing a thin, fleshy ectosome packed with microrhabds and other microscleres, through which projects an ‘extra-axial’ tract of megascleres, traversing the subectosome and projecting through the surface (NIWA 93474 leg.). D. *L. strongylata* showing the ‘axial’ arrangement of contort strongyloxeas in the deep choanosome (NIWA 93474 leg.). E. *Poecillastra ducitriaenea* Kelly & Cárdenas sp. nov. showing a densely packed, crustose ectosome through which subectosomal megascleres project, a relatively cavernous subectosomal region packed with microxeas, and in the deep choanosome, huge swathes of oxeas (NIWA 61944 leg., holotype). Scale bars = 500 μm.
clads of slightly uneven length, slightly curved downwards, 578(450–680) μm, overall cladome width, about 900–1360 μm long, ranging to pseudocalthrops. Broken true oxeas are evident but unmeasurable.

**Microscleres (Fig. 5B–C, F–G)**

Microxeas (Fig. 5B–C), straight to slightly curved, roughened, abundant, 332(260–420) × 7(5–8) μm, n = 20; plesiasters (Fig. 5F), with 3–5 microspined blunt-tipped rays, overall 67(50–100), ray length 37(25–60) μm, n = 10; metaster- to amphistaster- to spiraster-like streptasters (Fig. 5G), with long, microspined rays, abundant, 19(15–20) μm long.

**Distribution**

Macquarie Ridge.

**Substrate, depth range and ecology**

Attached to rock substrate; depth 462–524 m.

**Remarks**

The specimen is the attachment base of a sponge of unknown morphology, but it clearly differs from the holotype of *Poecillastra ducitriaena* sp. nov. in having a very hispid, crisp, scratchy surface, indicating a reduction of the ectosomal crust of microscleres, and the abundance of large megascleres. It is similar to *Poecillastra ducitriaena* sp. nov. in the possession of abundant contort oxeas in the stalk, but differs in the lack of straight oxeas in the stalk and the much larger dimensions of all the spicules; the contort oxeas are up to 2000 μm longer, on average, in *Poecillastra macquariensis* sp. nov., and the triaenes are about double the size of those in *Poecillastra ducitriaena* sp. nov., and much more abundant, the microxeas are about ten times larger, and the sponge contains plesiasters, absent in *Poecillastra ducitriaena* sp. nov.

Because our knowledge of *Poecillastra* in the New Zealand region is reasonable (see above), we have made the decision to record and name this second *Poecillastra* species, despite our lack of information on the body shape, and because surface texture, spicule types and dimensions are so different from those of *Poecillastra ducitriaena* sp. nov.

**Discussion**

We were not successful in obtaining sequences from the holotype of *L. strongylata* (NIWA 356 leg.) but we obtained one Folmer COI sequence from non-type specimen NIWA 51172 leg. and one COI mini-barcode (130 bp) from non-type specimen NIWA 51267 leg., both from the same type locality, the Three Kings Islands. The 28S fragment (C1–C2 domains) was also obtained for the latter two specimens (369 bp each). We obtained one COI mini-barcode from the holotype of *L. australis* sp. nov. (NIWA 89736) but failed to amplify 28S for *L. australis* sp. nov. We obtained the Folmer (661 bp) and the 28S fragments (308 bp) from the holotype of *Poecillastra ducitriaena* sp. nov. (NIWA 61944 leg.).

In the COI tree (Fig. 7), the two species of *Lamellomorpha*, which have identical sequences (at least identical minibarcodes) were a sister group to *Vulcanella* Sollas, 1886; this grouping with *Vulcanella* was poorly supported (bootstrap of 61). *Lamellomorpha + Vulcanella* were sister to a poorly supported *Poecillastra* clade (bootstrap of 60), which included *Poecillastra ducitriaena* sp. nov. Despite these poorly supported nodes, the Vulcanellidae was a very well supported clade suggesting that *Lamellomorpha* is clearly part of this family. The 28S tree confirms the position of *Lamellomorpha* in the Vulcanellidae but with no true support. *Vulcanella* appeared paraphyletic (poorly supported), while the grouping of *Lamellomorpha* was uncertain with respect to *Vulcanella* or *Poecillastra*. This poor resolution may be due to the short sequences we obtained (308–369 bp), which are in fairly conserved domains of 28S and so with few bp differences to differentiate species.
Fig. 7. Astrophorina COI and 28S maximum likelihood (ML) trees reconstructed with RAxML under the generalized time-reversible Gamma (GTRGAMMA) model. ML bootstrap supports (100 bootstrap replicates). GenBank accession numbers are given after each taxon name. In green are the new sequences produced during this study.
The grouping of *Lamellomorpha* with the Vulcanellidae suggested by the molecular markers is in accordance and supported by the morphological revision of this study. Indeed, the lamellate external morphology, the oscule/pore morphologies and distributions, as well as the skeleton organization in *Poecillastra* and *Lamellomorpha* are similar. Some spicules are also clearly homologous, such as the metasters to spirasters. The key discovery in this work is the confirmation that *Lamellomorpha* has integrity as a genus within the family Vulcanellidae, separate from *Vulcanela* and *Poecillastra*. This result generates new hypotheses about the evolution of spicules within the Vulcanellidae. The microrhabds in *Lamellomorpha* which earlier confused taxonomists, can now be considered as reduced *Poecillastra/Vulcanela* microxeas. Secondary losses of spicules have already been shown to be quite common in the Astrophorina (Cárdenas *et al.* 2011). Here, the short-shafted triaenes, which can be scarce (e.g., *Poecillastra compressa*) to quite rare (e.g., *Poecillastra ducitriaena* sp. nov.) in *Poecillastra*, would have been completely lost in *Lamellomorpha*. As for plesiasters (the largest streptaster category), they are absent in *Lamellomorpha*, as in some *Poecillastra* (e.g., *Poecillastra ducitriaena* sp. nov.) or *Vulcanela* (e.g., *Vulcanela horrida* (Schmidt, 1870)).

**Acknowledgements**

Specimens were provided by the NIWA Invertebrate Collection (NIC), and for material collected from the following research projects: Voyage KAH0606 — specimens were collected as part of a Ministry for Primary Industries funded voyage (NIWA project ENV2005–23), titled ‘The Effects of Fishing on the Benthic Community Structure between North Cape and Cape Reinga’; Voyage TAN0803 — specimens were collected during the interdisciplinary New Zealand-Australian MacRidge 2 research voyage (TAN0803), the biological component of which was part of NIWA’s research project, ‘Seamounts: their importance to fisheries and marine ecosystems,’ funded by the New Zealand Foundation for Research, Science and Technology, and CSIRO’s Division of Marine and Atmospheric Research project, ‘Biodiversity Voyages of Discovery,’ funded by the CSIRO Wealth from Oceans Flagship; Voyage TRIP3072 — specimens collected under the Scientific Observer Program funded by the New Zealand Ministry for Primary Industries; Voyage TAN1105 — specimens were collected as part of the Biogenic Habitats on the Continental Shelf project (voyages TAN1105 & TAN1108), funded by New Zealand Ministry of Fisheries (Biogenic Habitats: ZBD200801), New Zealand Foundation for Research, Science and Technology (CCM: CO1X0907), NIWA Capability Fund (CF111358) and Oceans Survey 20/20 RV Tangaroa days funded by Land Information New Zealand; NIWA Stn Z15944, collector CRRF: Coral Reef Research Foundation under contract to the US National Cancer Institute (N02-CM-77249).

We thank emeritus professors Murray Munro and John Blunt, Department of Chemistry, University of Canterbury, Christchurch, for the donation of their sponge vouchers to NIC, for their continued study. We also thank Dr Bruce Marshall, Museum of New Zealand Te Papa Tongarewa, Wellington, for the loan of specimens from their collections. We are grateful to John Rosser MA LTCL for his assistance with suggestions for new species names and advising on the correct form of the new taxon names. Satya Amirapu, Auckland University, carried out our histological work. PC received support from the European Union's Horizon 2020 research and innovation program through the SponGES project (grant agreement No. 679849). This document reflects only the authors’ view and the Executive Agency for Small and Medium-sized Enterprises (EASME) is not responsible for any use that may be made of the information it contains. This research was funded by NIWA under Coasts and Oceans Research Programme Marine Biological Resources: Discovery and definition of the marine biota of New Zealand (2016/2017 to 2018/2019 SCI).
References


**Manuscript received: 23 October 2018**  
**Manuscript accepted: 18 February 2019**  
**Published on: 14 March 2019**  
**Topic editor: Rudy CAM Jocque**  
**Desk editor: Alejandro Quintanar**

Printed versions of all papers are also deposited in the libraries of the institutes that are members of the *EJT* consortium: Muséum national d’Histoire naturelle, Paris, France; Meise Botanic Garden, Belgium; Royal Museum for Central Africa, Tervuren, Belgium; Royal Belgian Institute of Natural Sciences, Brussels, Belgium; Natural History Museum of Denmark, Copenhagen, Denmark; Naturalis Biodiversity Center, Leiden, the Netherlands; Museo Nacional de Ciencias Naturales-CSIC, Madrid, Spain; Real Jardín Botánico de Madrid CSIC, Madrid, Spain; Zoological Research Museum Alexander Koenig, Bonn, Germany.