# Integrative taxonomy of calcareous sponges (Porifera: Calcarea) from Réunion Island, Indian Ocean

<sup>1</sup>Universidade Federal do Rio de Janeiro, Instituto de Biologia, Departamento de Zoologia, Av. Carlos Chagas Filho 373, CEP 21941-902, Rio de Janeiro, Brazil

<sup>2</sup>Institut Méditerranéen de Biodiversité et d'Ecologie marine et continentale IMBE, UMR CNRS 7263, IRD 237, Aix Marseille Université, Avignon Université, Station Marine d'Endoume, Rue de la Batterie des Lions 13007 Marseille, France

Received 8 December 2020; revised 19 February 2021; accepted for publication 23 February 2021

The Western Indian Ocean Province is reckoned for its rich marine diversity; however, sponges of the Mascarene Islands ecoregion are still poorly known. In La Réunion, only three species of class Calcarea have been registered. Hence, calcareous sponges were searched in seven sites representing various habitats of the Western coast of La Réunion, but found in only three of them. A total of 23 sponge samples was identified using morphological and molecular taxonomy. This sampling represents 11 species, all new records for the region, and seven of them are new to science: Ascandra mascarenica sp. nov., A. oceanusvitae sp. nov., Janusya indica gen. et sp. nov., Leucascus tenuispinae sp. nov., Lelapiella tertia sp. nov., Soleneiscus intermedius sp. nov. and Leucandra ornata sp. nov.; and a new genus, Janusya gen. nov.. Based on results from this and from previous studies, we propose the synonymization of the order Murrayonida with Clathrinida. A very low sampling effort has thus increased the number of calcareous sponge species from the Mascarenes Islands ecoregion by 69 % and from La Réunion by 367%.

ADDITIONAL KEYWORDS: biodiversity – Indian Ocean – molecular systematics – morphological systematics – phylogenetic systematics – Porifera.

#### INTRODUCTION

Although the Western Indian Ocean Province presents a rich sponge fauna, the Mascarene Islands ecoregion is still poorly known (Van Soest et al., 2012; Van Soest & De Voogd, 2018). The Mascarene Islands ecoregion includes the islands of La Réunion, Mauritius and Rodrigues (Spalding et al., 2007) and, to date, 66 species of sponges have been recorded from this ecoregion (Schuffner, 1877; Vacelet, 1977; Lévi, 1986; Bourmaud, 2003; Van Soest & De Voogd, 2018; Van Soest et al., 2020), 16 belonging to the class Calcarea (Schuffner, 1877; Vacelet, 1977; Bourmaud, 2003; Van Soest & De Voogd, 2018).

Among these three islands, the sponge biodiversity of La Réunion is by far the least known. To our knowledge, the only studies that listed sponges from La Réunion were those of Vacelet (1977), a taxonomic work on sponges from French Polynesia, Madagascar and La Réunion; Bourmaud (2003), an inventory of the reef marine biodiversity of the western coast of La Réunion and Schlever et al. (2016), a work on the benthos growing on submerged lava beds in the eastern coast of La Réunion. Only the first and the second studies recorded calcareous sponges, a total of three species: Vacelet (1977) described two hypercalcified calcareous sponges from La Réunion: the calcaroneans Tulearinia stylifera Vacelet, 1977 and Plectroninia hindei Kirkpatrick, 1900, and Bourmaud (2003) added the calcinean Leucetta microraphis Haeckel, 1872 to the list.

Calcarea represents the only sponge class whose skeleton contains calcite spicules (Borojević *et al.*,

<sup>\*</sup>Corresponding author. E-mail: mklautau@biologia.ufrj.br [Version of record, published online 6 May 2021; http://zoobank.org/urn:lsid:zoobank.org:pub:3E370CBF-7DD1-4E73-BC37-81BF12EDFEED

2002). These sponges are exclusively marine, viviparous and widely distributed at all latitudes and longitudes (Manuel et al., 2002), from intertidal zones down to more than 4000 m deep (Rapp et al., 2011). They are commonly neglected in inventories, possibly because of their small size and preference for cryptic habitats (e.g. Wörheide & Hooper, 1999; Klautau & Borojević, 2001; Van Soest et al., 2012). It is not surprising that only three species were described from Réunion, considering that no specific sampling efforts have been made searching for Calcarea in the waters around this island.

To import skills that did not exist and improve knowledge on the sponge biodiversity, the Vie Océane association and the Marine Natural Reserve of Réunion (RNMR) organized a sponge training course in 2017. Designed on the same model as a previous project in Martinique (Pérez et al., 2017), this course in sponge biology and taxonomy also aimed to provide a first sponge inventory together with first biological explorations of submarine caves. The focus on calcareous sponges resulted in taxonomic identifications and several descriptions presented here, and in a discussion on affinities of the calcareous sponge fauna across the Mascarene Islands, and between the Mascarene ecoregion and other ecoregions of the Western Indian Ocean.

## MATERIAL AND METHODS

## SAMPLE COLLECTION

Seven sites in the western part of La Réunion were investigated, three of them being shallow-water lagoons explored by free diving, the other four localities were reefs or caves explored by scuba diving down to 30 m deep: (1) Cap La Houssaye Reef (CAH2 -21°1'3.2"S, 55°14'13.23"E), cliff and reef bench were explored; (2) Passe de l'Hermitage Reef (PAE2 - 21° 5'11.50"S, 55°13'12.33"E), a fringing reef of the Saint Gilles Ermitage complex; (3) L'Hermitage Lagoon (PAE1 - 21°5'0.83"S, 55°13'24.48"E), a reef complex of La Saline-Ermitage, including the Closerie and the south part of the l'Hermitage Pass; (4) La Saline Lagoon (TRO1 – 21°6'0.73"S, 55°14'26.30"E), a reef complex of La Saline-Hermitage, a hole with water that replaces that of Saint Leu (La Varangue) in heavy swells; (5) Le Sec Jaune Caves  $(SJN2 - 21^{\circ}12'25.63"S,$ 55°16'37.16"E), south area of Pointe au Sel, a cliff and reef banks with numerous overhangs and caves; (6) L'Étang Salé Lagoon (ESB1 – 21°16'5.22"S, 55°19'57.43"E), the southern part of the channel was investigated; (7) Portail Caves (ESB2 – 21°14'31.07"S, 55°17'41.24"E), old lava tunnels.

Each site was visited only once. While the shallow-water sites were explored by four to ten people, most sampling in caves was performed by only one scubadiver. Whenever possible, the studied sponges were photographed *in situ* before collection, and then fixed and preserved in ethanol 93%. A total of 23 calcareous sponges were collected and they were deposited in the Porifera Collection of the Universidade Federal do Rio de Janeiro (UFRJPOR).

#### INSTITUTIONAL ABBREVIATIONS

GW: Gert Wörheide collection, Munich, Germany; MNRJ: Museu Nacional da Universidade Federal do Rio de Janeiro, Brazil; NHMUK: Natural History Museum, London, United Kingdom; PMR: Prirodoslovni Muzej Rijeka, Croatia; QM: Queensland Museum, Brisbane, Australia; RMNH: Naturalis Biodiversity Center, Leiden, The Netherlands; SAM: South Australian Museum, Adelaide, Australia; SMF; Senckenberg Museum, Frankfurt, Germany; SNSB-BSPG: Bayerische Staatssammlung für Paläontologie und historische Geologie, Staatlichen Naturwissenschaftlichen Sammlungen Bayern, Munich, Germany; UFRJPOR: Porifera collection of the Biology Institute of Federal University of Rio de Janeiro, Brazil; WAMZ: Western Australian Museum, Perth, Australia; ZMAPOR: Zoölogisch Museum, Instituut voor Systematiek en Populatiebiologie, Amsterdam, The Netherlands

#### MOLECULAR ANALYSES

Total DNA was extracted using the guanidine/phenol-chloroform protocol (Hillis et al., 1996; modified from Sambrook et al., 1989). To test the taxonomic status of the specimens and their phylogenetic relationships, two molecular markers were used: C-region of the 28S (referred here as C-LSU) for both subclasses Calcinea and Calcaronea, and the region comprising the spacers ITS1 and ITS2 and the 5.8S ribosomal DNA (referred as ITS) for Calcinea only. Polymerase chain reaction (PCR) sequencing conditions and sequence editing were the same used by our group in previous works (e.g. Cóndor-Luján et al., 2018).

In addition to the DNA sequences generated in this work, we also used sequences from GenBank (https://www.ncbi.nlm.nih.gov/genbank). DNA sequences were aligned through the online version of the program MAFFT v.7 (Katoh & Standley, 2013) using the strategy Q-INS-i (Katoh & Toh, 2008), taking the secondary structure of ribosomal DNA into consideration. The final alignment of 49 operational taxonomic units (OTU) ITS sequences' of Calcinea was 1208 bp length, including gaps. A total of 357 conserved sites, 616 variable

sites and 136 singletons were retrieved (Supporting Information, Fig. S1). For the C-LSU, the final alignment comprised 71 OTU and 381 bp length, including gaps and a total of 139 conserved sites, 218 variable sites and 36 singletons (Supporting Information, Fig. S2). For Calcaronea, the C-LSU alignment had 45 OTU and a total length of 430 bp (including gaps), 273 conservative sites, 12 variable sites and 25 singletons (Supporting Information, Fig. S3). The substitution model chosen for the phylogenetic analyses was GTR+G+I for Calcinea (ITS and C-LSU) and TN93+G for Calcaronea (C-LSU only), according to the Model Selection in MEGA 7.0, using an initial automatic (neighbour-joining) tree and considering all sites of the alignment based on the Bayesian information criterion.

A maximum likelihood (ML) tree was generated using the software IQ-Tree 2 (Minh et al., 2020) for each marker. Bootstrap supports were obtained with the ultrafast bootstrap (Hoang et al., 2017) using 1000 pseudo-replicates (Felsenstein, 1985; Russo, 1997). Bayesian inferences (BI) were performed in MrBayes 3.1.2. (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003), using the same evolutionary model parameters as selected for the ML analysis and two runs and four independent chains for 106 generations and a 25% burn-in, resulting in a majority consensus tree (of the sampled trees). Phylogenetic analyses were conducted using the CIPRES Science Gateway (Portal 2) online server (Miller et al., 2012).

Both phylogenies of Calcinea (ITS and C-LSU) and Calcaronea (C-LSU) were midpoint-rooted, as the only appropriate outgroup would be the opposing subclass, and using such distant groups as outgroups would result in loss of information within closely related clades (Hess & Russo, 2007). However, we also built the trees with the opposing subclass as outgroups to confirm that there would be no important change in the topology of the midpoint-rooted trees (Supporting Information, Figs S4, S5).

The uncorrected *p*-distance was calculated in MEGA 7.0 to estimate the intraspecific and interspecific variability, considering the morphological identifications. Two criteria were used for the molecular identification of the specimens: the monophyly of morphologically conspecific specimens, and the genetic dissimilarity (*p*-distance) to the most closely related species with available DNA data.

Species names, location, voucher numbers and GenBank accession numbers of the DNA sequences generated in this work and those obtained from the GenBank database (https://www.ncbi.nlm.nih.gov/genbank) used in the phylogenetic tree are listed in Table 1.

#### MORPHOLOGICAL ANALYSES

Slides of spicules and tangential ('in toto') were made according to standard protocols (Wörheide & Hooper, 1999; Klautau & Valentine, 2003). Skeleton sections were made, including fragments of the sponges, previously stained with a 5% fuchsin alcoholic solution, in paraffin and making thick sections (c. 500 µm) with a microtome. The mounting medium used was Entellan (Merck). Spicule measurements (length and width at the base of the actines) were taken from 20 spicules, when possible, for every category in light microscopy using an ocular micrometer. The biggest and the smallest spicules were searched for, while the other 18 were randomly measured. The results are presented in tabular form presenting minimum (min), mean, standard deviation (SD) and maximum (max) values of both measurements, and number of analysed spicules (N).

Photographs of preserved specimens were taken with a digital camera AxioCam ERc5s coupled to a Zeiss stereomicroscope. Slides photographs were taken with a Canon G12 camera coupled to a Zeiss Axioskop light microscope. Scanning electron microscopy (SEM) was performed in a JSM-6510 microscope at the Instituto de Biologia of the Universidade Federal do Rio de Janeiro. For this purpose, isolated spicules in ethanol 100% were transferred to stubs with double-sided carbon conductive tape and coverslip, dried and sputter coated with gold in a Balzers apparatus at the Instituto de Ciências Biomédicas of the Universidade Federal do Rio de Janeiro.

## RESULTS

Among the 23 calcareous sponges collected, we identified 11 species using both morphology and DNA sequences. All of them are new records for La Réunion. The subclass Calcinea was more diverse than Calcaronea, with eight and three species, respectively. Seven species are new to science, and include a new genus.

The species were distributed among only three sites: Passe de l'Ermitage, Reef and Cave (PAE2), Étang Salé, Portail Caves (ESB2) and Cap Lahoussaye Reef (CAH2; Fig. 1), all of them being caves, PAE2 appearing by far the richest. One single scuba-dive was not enough to cover the large networks of caves found in Le Sec Jaune Caves, what may explain the lack of calcareous specimens after the exploration of this cryptic habitat. On the other hand, whereas a much higher sampling effort was deployed in shallow waters (lagoons), no calcareous sponges could be found in this type of habitat.

 $\textbf{Table 1.} \ \ \textbf{Species names, voucher numbers and GenBank accession numbers used in this study.} \ \ \textbf{*Sequences generated in this study}$ 

| Species   | Locality   | Voucher number               | GenBank acc          | ession               |
|---|--|------------------------------|----------------------|----------------------|
|   |  |                              | ITS                  | C-LSU                |
| Calcinea  |  |                              |                      |                      |
| Arturia alcatraziensis                            | Cape Verde, Africa                               | <b>ZMAPOR 07061</b>          | KC843431             | -                    |
| Arturia spirallata                                | Peru   | MNRJ 11414                   | KC985142             | -                    |
| Arturia sueziana                                  | Red Sea  | SNSB-BSPG GW3120             | KY366410             | KY366370             |
| Arturia sueziana                                  | Maldives   | RMNH10112                    | -                    | MF872755             |
| Ascaltis reticulum                                | Mediterranean Sea                                | UFRJPOR 6258                 | HQ588973             | HQ589014             |
| Ascandra ('Ernstia') chrysops                     | Indonesia  | RMNH 1773                    | -                    | MF686061             |
| Ascandra contorta                                 | Mediterranean Sea                                | UFRJPOR 6327                 | HQ588970             | HQ589011             |
| Ascandra corallicola                              | Norway   | UFRJPOR 6329                 | HQ588994             | HQ589012             |
| Ascandra cf. crewsi                               | French Polynesia                                 | BMOO16290                    | KC843446             | -                    |
| Ascandra falcata                                  | Mediterranean Sea                                | UFRJPOR 5856                 | HQ588962             | -                    |
| Ascandra kakaban                                  | Indonesia  | RMNH1696                     | -                    | MF686077             |
| Ascandra mascarenica sp. nov. *                   | Réunion, Western Indian                          | UFRJPOR 8936                 | -                    | MW350111             |
| Ascandra oceanusvitae sp. nov.*                   | Réunion, Western Indian                          | UFRJPOR 8944                 | -                    | MW350112             |
| Ascandra oceanusvitae sp. nov.*                   | Réunion, Western Indian                          | UFRJPOR 8947                 | MW350104             | -                    |
| Ascandra spalatensis                              | Adriatic, Croatia                                | PMR 17806                    | KP740024             | -                    |
| Bidderia bicolora                                 | Caribbean  | UFRJPOR 8228                 | MG017978             | MG017978             |
| Borojevia brasiliensis                            | Brazil   | UFRJPOR 5406                 | KX548909             | HQ589015             |
| Borojevia cerebrum                                | Mediterranean Sea                                | UFRJPOR 6322                 | HQ588964             | HQ589008             |
| Brattegardia nanseni                              | Greenland  | UFRJPOR 6332                 | HQ588982             | HQ589013             |
| Burtonulla sibogae                                | Indonesia  | RMNH9195                     | -                    | MF686091             |
| Clathrina aurea                                   | Brazil   | MNRJ 8998                    | HQ588968             | HQ589005             |
| Clathrina clathrus                                | Mediterranean Sea                                | UFRJPOR 6315                 | HQ588965             | HQ589009             |
| Clathrina conifera                                | Brazil   | UFRJPOR 8991                 | HQ588959             | HQ589010             |
| Clathrina coriacea                                | Norway   | UFRJPOR 6330                 | HQ588986             | HQ589001             |
| Clathrina curacaoensis                            | Curação  | UFRJPOR 6734                 | MF472607             | MF472612             |
| Clathrina cylindractina                           | Brazil   | UFRJPOR 5206                 | HQ588979             | HQ589007             |
| Clathrina lacunosa                                | Norway   | UFRJPOR 6334                 | HQ588991             | HQ589020             |
| Clathrina lutea                                   | Brazil   | UFRJPOR 5172                 | HQ588961             | HQ589004             |
| Clathrina luteoculcitella                         | Great Barrier Reef (GBR),<br>Australia           | QM G313684                   | HQ588989             | JQ272283             |
| Clathrina ramosa                                  | Chile  | MNRJ 10313                   | HQ588990             | HQ589002             |
| Clathrina rotundata                               | Red Sea  | SMF 11636                    | KY711435             | KY366399             |
| Clathrina rubra                                   | Adriatic Sea                                     | PMR-14306                    | KC479088             | KC479082             |
| Clathrina wistariensis                            | Australia  | QM G313663                   | HQ588987             | JQ272303             |
| Ernstia klautauae                                 | Indonesia  | ZMAPOR 8390                  | KC843451             | -                    |
| Ernstia klautauae                                 | Indonesia  | RMNH9341                     | -                    | MF686062             |
| Ernstia rocasensis                                | Brazil   | UFRJPOR 6664                 | KX548928             | -                    |
| Ernstia tetractina                                | Brazil   | UFRJPOR 5183                 | HQ589000             | HQ589021             |
| Janusya ('Arturia') adusta comb. nov.             | Seychelles, Western Indian                       | ZMAPOR 10612                 | -                    | MF872756             |
| Janusya ('Arturia') angusta comb. nov.            | Indonesia  | ZMAPOR 8221a                 | -                    | MF686060             |
| Janusya ('Arturia') tubuloreticulosa comb. nov.   | Indonesia  | RMNH5547                     | -                    | MF686059             |
| Janusya indica sp. nov.*                          | Réunion, Western Indian                          | UFRJPOR 8932                 | MW350105             | _                    |
| Janusya indica sp. nov.*                          | Réunion, Western Indian                          | UFRJPOR 8945                 | -                    | MW350113             |
| Lelapiella incrustans                             | Vanuatu  | QM G313914                   | _                    | JQ272306             |
| Lelapiella tertia sp. nov.*<br>Leucaltis clathria | Réunion, Western Indian<br>Panama, Caribbean Sea | UFRJPOR 8937<br>UFRJPOR 6942 | MW350106<br>KC843452 | MW350114<br>KC869524 |

Table 1. Continued

| Species                           | Locality                   | Voucher number                         | GenBank acc<br>number | ession   |
|-----------------------------------|----------------------------|--|-----------------------|----------|
|                                   |                            |  | ITS                   | C-LSU    |
| Leucaltis clathria                | Panama, Caribbean Sea      | UFRJPOR 6944                           | KU568493              | -        |
| Leucaltis nodusgordii             | Australia                  | QM G316022                             | AJ633861              | JQ272302 |
| $Leucaltis\ nodusgordii*$         | Réunion, Western Indian    | UFRPOR 8930                            | MW350107              | MW350115 |
| Leucaltis nodusgordii             | Seychelles, Western Indian | <b>ZMAPOR 12436</b>                    | -                     | MF872787 |
| Leucaltis nodusgordii             | Seychelles, Western Indian | <b>ZMAPOR 12443</b>                    | -                     | MF872788 |
| Leucaltis nodusgordii             | Indonesia                  | RMNH1772                               | -                     | MF686078 |
| Leucaltis nodusgordii             | Australia                  | H10                                    | -                     | AY563542 |
| Leucascus flavus                  | Indonesia                  | RMNH 2279                              | -                     | MF686080 |
| Leucascus simplex                 | French Polynesia           | BMOO16283                              | KC843454              | -        |
| Leucascus simplex                 | French Polynesia           | UFRJPOR 6451                           | -                     | MN422254 |
| Leucascus sp.                     | Australia                  | QM G316051                             | -                     | JQ272305 |
| Leucascus tenuispinae sp. nov.*   | Réunion, Western Indian    | UFRJPOR 8927                           | -                     | MW350117 |
| Leucetta antarctica               | Antarctic                  | MNRJ 13798                             | KC849700              | -        |
| $Leucetta\ avocado$               | Mauritius, Western Indian  | NCI 108                                | -                     | KC869542 |
| $Leucetta\ chagosensis^*$         | Réunion                    | UFRJPOR 8939                           | MW350108              | MW350118 |
| Leucetta chagosensis              | Australia                  | -                                      | -                     | AY563543 |
| Leucetta chagosensis              | Indonesia                  | ZMAPOR 13283                           | -                     | MF686085 |
| Leucetta chagosensis              | Rodrigues, Western Indian  | RMNH11657 1                            | -                     | MF872790 |
| Leucetta chagosensis              | Rodrigues, Western Indian  | RMNH11658                              | -                     | MF872791 |
| Leucetta chagosensis              | French Polynesia           | UFRJPOR 6455                           | -                     | MN422257 |
| Leucetta chagosensis              | -                          | -                                      | -                     | AM180994 |
| Leucetta chagosensis              | Australia                  | H10                                    | AF458861              | -        |
| Leucetta chagosensis              | French Polynesia           | BMOO 16210                             | KC843455              | -        |
| Leucetta chagosensis              | Saudi Arabia, Red Sea      | SNSB-BSPG GW3052                       | KY366408              | -        |
| Leucetta delicata                 | Antarctic                  | SMF 11868                              | KC874654              | -        |
| Leucetta floridana                | Brazil                     | UFRJPOR 5630                           | EU781968              | -        |
| Leucetta floridana                | Panama, Caribbean Sea      | P10x2                                  | -                     | KC869538 |
| Leucetta foliata                  | Western Australia          | UFRJPOR 7149                           | -                     | KX499451 |
| Leucetta giribeti                 | Antarctic                  | NHMUK:2017.1.19.1                      | KY670632              | KY670632 |
| Leucetta microraphis              | GBR, Australia             | QM G313659                             | AF479061              | JQ272297 |
| Leucetta potiguar                 | Brazil                     | MNRJ 8474                              | EU781981              | -        |
| Leucetta purpurea                 | Western Australia          | UFRJPOR 7255                           | -                     | KX499450 |
| Leucetta cf. pyriformis           | Antarctic                  | MNRJ 13843                             | KC843457              | -        |
| Leucetta sulcata                  | Rodrigues                  | RMNH 11639_1                           | -                     | MF872798 |
| Leucetta villosa                  | Maldives                   | QM G313662                             | -                     | JQ272295 |
| Leuclathrina translucida          | Maldives                   | SNSB-BSPG GW9394                       | -                     | LT899664 |
| Levinella prolifera               | GBR, Australia             | QM G313818                             | -                     | AM180984 |
| Levinella thalassae               | Rockall Bank, Ireland      | ZMAPOR 18150                           | -                     | MF686093 |
| Murrayona phanolepis              | Australia                  | QM G313992                             | -                     | JQ272304 |
| Murrayona phanolepis              | Australia                  | QM G316290                             | -                     | AM180998 |
| Nicola tetela                     | Curaçao                    | UFRJPOR 6723                           | KU568492              | -        |
| Soleneiscus intermedius sp. nov.* | Réunion, Western Indian    | UFRJPOR 8933                           | -                     | MW350119 |
| Soleneiscus hamatus               | Egypt, Red Sea             | ${\rm SNSB\text{-}BSPG}\;{\rm GW2975}$ | -                     | KY366365 |
| Soleneiscus radovani              | GBR, Australia             | QM G313661                             | -                     | JQ272289 |
| Soleneiscus stolonifer            | GBR, Australia             | QM G313668                             | -                     | JQ272290 |
| Calcaronea                        | M                          | CAM DOOR 40                            |                       | 10070070 |
| Aphroceras sp. 1                  | Tasmania, Australia        | SAM-PS0349                             | -                     | JQ272273 |
| Aphroceras sp. 2                  | Coral Sea, Australia       | QM G316285                             | -                     | AM181001 |
| Aphroceras sp. nov.               | St. Helena, South Atlantic | SA 87                                  | -                     | MH385226 |

Table 1. Continued

| Species                                | Locality  | Voucher number                 | GenBank<br>number | accession            |
|--|---|--------------------------------|-------------------|----------------------|
|  |   |                                | ITS               | C-LSU                |
| Aphroceras sp. nov.                    | St. Helena, South Atlantic                            | SA 89                          | -                 | MH385227             |
| Aphroceras seychellensis comb. nov.*   | Réunion, Western Indian                               | UFRJPOR 8926                   | -                 | MW350109             |
| Aphroceras seychellensis comb. nov.*   | Réunion, Western Indian                               | UFRJPOR 8935                   | -                 | MW350110             |
| Breitfussia schulzei                   | West Greenland  | FB 61                          | -                 | MH385228             |
| Grantia compressa                      | Roscoff, France                                       | SA 80                          | -                 | MH385242             |
| Grantiopsis cylindrica                 | GBR, Australia  | SNSB-BSPG GW973                | -                 | JQ272266             |
| Grantiopsis heroni                     | -   | -                              | -                 | AY563539             |
| Leucandra ananas                       | Trondheimsfjord, Norway                               | SA 136                         | -                 | MH385249             |
| Leucandra aspera                       | -   | -                              | -                 | AY563535             |
| Leucandra falakra                      | Croatia, Adriatic Sea                                 | PMR 13748                      | -                 | KT447560             |
| Leucandra cf. gausapata                | Antarctic   | HT 15                          | -                 | MH385252             |
| Leucandra mozambiquensis               | Mozambique, West Africa                               | ZMA 22408B                     | -                 | MF872766             |
| Leucandra nicolae                      | GBR, Australia  | QM G313672                     | _                 | JQ272278             |
| Leucandra ornata sp. nov.*             | Réunion, Western Indian                               | UFRJPOR 8928                   | _                 | MW350116             |
| Leucandra penicillata                  | West Greenland  | FB 35                          | _                 | MH385256             |
| Leucandra pilula                       | Seychelles, Western Indian                            | ZMA 10528                      | _                 | MF872767             |
| Leucandra spinifera                    | Croatia, Adriatic Sea                                 | IRB SG3                        | _                 | KT447561             |
| Leucandra tahuatae                     | Tahuata, French Polynesia                             | UFRJPOR 6454                   | _                 | MN422253             |
| Leucandrilla sp.                       | Galapagos   | RMNH 11520                     | _                 | MF872768             |
| Leucilla antillana                     | Curação, Caribbean Sea                                | UFRJPOR 6768                   | _                 | MF472615             |
| Leucosolenia botryoides                | Norway  | SA 60                          | _                 | MH385257             |
| Leucosolenia complicata                | Norway  | AA 201                         | _                 | MH385258             |
| Leucosolenia cf. corallorrhiza         | Greenland   | FB 14                          | _                 | MH385270             |
| Paragrantia waguensis                  | Japan   | RMNH 9317                      | _                 | KT277668             |
| Paraleucilla sp. 1                     | St. Helena, South Atlantic                            | SA 83                          | _                 | MH385283             |
| Paraleucilla erpenbecki                | Mozambique, West Africa                               | ZMAPOR 22409C                  | _                 | MF872728             |
| Paraleucilla magna                     | Algarve, Portugal                                     | PT 06                          | _                 | MH385279             |
| Sycon caminatum                        | Madeira Island  | PT 02                          | _                 | MH385301             |
| Sycon villosum                         | Moøyastaken, Norway                                   | SA 24                          |                   | MH385314             |
| Synute pulchella                       | West Australia  | WAMZ 1404                      |                   | JQ272274             |
| Teichonopsis labyrinthica              | South Australia                                       | SAM-PS 0228                    |                   | JQ272264             |
| Ute ampullacea                         | GBR, Australia  | QM G313669                     |                   | AM181002             |
| Ute glabra                             | Portugal  | PT 16                          |                   | MH385315             |
| Ute gladiata                           | Korsfjard, Norway                                     | HT 47                          | -                 | MH385316             |
|  |   | SA 134                         | -                 |                      |
| Ute gladiata                           | Trondheimsfjord, Norway                               | ZMAPOR 11562                   | -                 | MH385317             |
| Ute insulagemmae                       | Seychelles, Western Indian<br>Réunion, Western Indian | UFRJPOR 8925                   | -                 | MF872772<br>MW350120 |
| Ute insulagemmae*                      |   |                                | -                 |                      |
| Ute insulagemmae*                      | Réunion, Western Indian                               | UFRJPOR 8929<br>UFRJPOR 8938   | -                 | MW350121             |
| Ute insulagemmae*<br>Ute insulagemmae* | Réunion, Western Indian                               |                                | -                 | MW350122<br>MW350123 |
| Ute insulagemmae*                      | Réunion, Western Indian                               | UFRJPOR 8942A<br>UFRJPOR 8943A | -                 |                      |
| 9                                      | Réunion, Western Indian                               |                                | -                 | MW350124             |
| Ute insulagemmae*                      | Réunion, Western Indian                               | UFRJPOR 8948                   | -                 | MW350125             |
| Ute aff. syconoides                    | GBR, Australia  | GW975                          | -                 | JQ272270             |
| Ute aff. syconoides                    | GBR, Australia  | QM G313694                     | -                 | JQ272271             |
| Ute aff. syconoides                    | Tasmania, Australia                                   | QM G323233                     | -                 | JQ272269             |

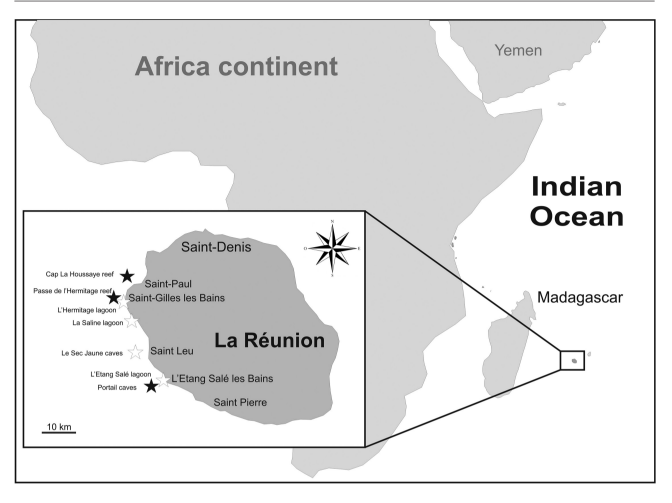


Figure 1. Study area. Sampling localities in La Réunion, Indian Ocean. The black stars indicate the locations where calcareous sponges were found.

## MOLECULAR RESULTS

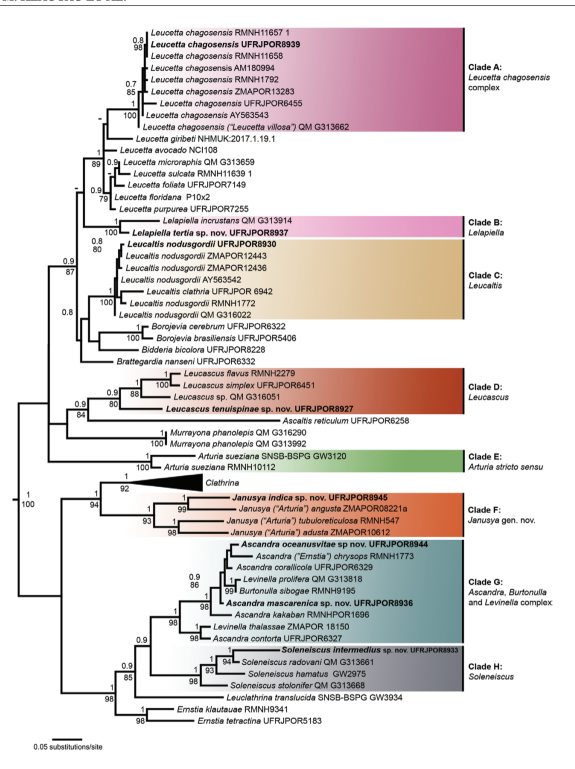
Besides the midpoint-rooted trees (C-LSU of Calcinea and Calcaronea and ITS of Calcinea), we also built C-LSU ML trees rooted with the opposing class (Supporting Information, Figs S4, S5). These trees showed almost the same topology of the midpoint-rooted trees but the midpoint-rooted trees showed better internal resolution. Consequently, we present our molecular results based on the midpoint-rooted phylogenies. Bootstrap and posterior probability values are shown in each ML tree.

## CALCINEA PHYLOGENY (ITS AND C-LSU)

The recovered phylogenetic trees of C-LSU and ITS are shown in Figures 2 and 3, respectively. In both trees, we alphabetically named the clades with specimens from La Réunion (and *Arturia Azevedo et al.*, 2017 s.s.) from A to H. In both C-LSU and ITS trees, the polyphyletic genus *Leucetta* appears monophyletic (bootstrap/posterior

probability: 89%/1 and 73%/0.9, respectively), but only because we did not include Pericharax Poléjaeff, 1883 nor Leucettusa Haeckel, 1872 in the analyses (e.g. Klautau et al., 2013; Lopes et al., 2018). In clade A, the sequence of Leucetta chagosensis Dendy, 1913 from La Réunion grouped with other sequences of that species from the Indo-Pacific and with L. villosa Wörheide & Hooper, 1999 (Table 1; Fig. 2) with a bootstrap/posterior probability of 100%/1 in both trees. The intraspecific p-distance among them was 0.0–2.3% for the C-LSU and 1.5% for the ITS. This relatively high *p*-distance and wide-spread scenario suggest that L. chagosensis may be a complex of cryptic species, as already proposed by Wörheide et al. (2002, 2008), Klautau et al. (2020) and Pasnin et al. (2020). Our results indicate that *L. villosa* is part of this complex (see the 'Taxonomic remarks' of L. chagosensis).

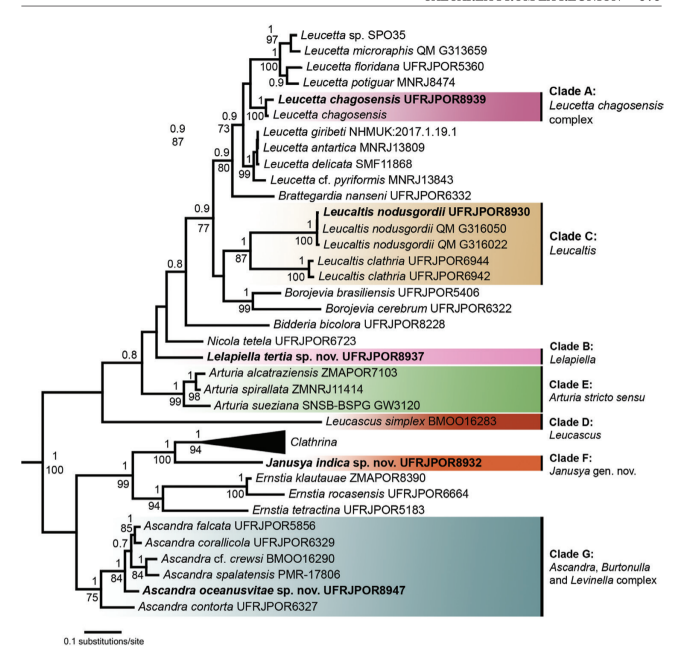
Clade B contains the new species *Lelapiella tertia* (described below), although for the ITS tree we did not have a clade because this species was the first *Lelapiella* Vacelet, 1977 sequence generated. Because



**Figure 2.** Maximum likelihood phylogenetic tree based on C-LSU sequences of the subclass Calcinea. Midpoint-rooted tree. Species sequences generated in this work are in bold. Clades containing species found in this work are highlighted. Support values are shown at the nodes. Support values are shown at the nodes (bootstrap > 70%; posterior probability > 0.7).

we included *Lelapiella incrustans* Vacelet, 1977 in our C-LSU dataset, both species grouped in a clade with a high bootstrap/posterior probability (100%/1)

and *p*-distance of 5.5%. *Lelapiella* did not group with sequences of *Murrayona phanolepis* Kirkpatrick, 1910, another species from the order Murrayonida.



**Figure 3.** Maximum likelihood phylogenetic tree based on ITS sequences of the subclass Calcinea. Midpoint-rooted tree. Species sequences generated in this work are in bold. Clades containing species found in this work are highlighted. Support values are shown at the nodes (bootstrap > 70%; posterior probability > 0.7).

Clade C contains sequences of *Leucaltis clathria* Haeckel, 1872 and *L. nodusgordii* (Poléjaeff, 1883). In the ITS tree, *L. nodusgordii* from La Réunion clustered in a highly supported clade with sequences from Australia (bootstrap/posterior probability: 100%/1) and with low *p*-distance (0.0–0.4%). This clade is closely related to the *L. clathria* clade (bootstrap/posterior probability: 87%/1), but with high genetic divergence of 11.1–11.9% (*p*-distance), confirming that it actually includes two different species and that

the genus is monophyletic. The C-LSU resolution for this clade was not satisfactory, because no lineage differentiation was recovered between *L. clathria* and *L. nodusgordii*. The sequences form a polytomic branch with low resolution and, except for the genus bootstrap/ posterior probability (100%/1), all the bootstraps were low. The *p*-distances range from 0.3 to 3.6%.

For the new species *Leucascus tenuispinae* (described below; clade D), we got only a C-LSU sequence. The genus *Leucascus* Dendy, 1892 was found monophyletic

in our tree (bootstrap/posterior probability: 80%/0.9), *L. tenuispinae* grouping more distantly with *Leucascus* sp. (*p*-distance: 9.3%), *L. flavus* Cavalcanti *et al.*, 2013 (*p*-distance: 10.2%) and *L. simplex* Dendy, 1892 (*p*-distance: 9.9%).

In clade E, both molecular markers recovered the genus Arturia stricto sensu. In the ITS tree, it grouped A. alcatraziensis (Lanna et al., 2007), A. spirallata Azevedo et al., 2015 and A. sueziana (Klautau & Valentine, 2003) (bootstrap/posterior probability: 99%/1). Other sequences of Arturia clustered together with the independent lineage of the new genus Janusya (clade F; bootstrap/ posterior probability: 93%/1 in the C-LSU tree). The new species *Janusya indica* (described below) was the only sequence available of this genus for the ITS marker. Therefore, no intra- or interspecific comments can be made. On the other hand, three other species of Arturia lato sensu were closely related to the new genus Janusya in the C-LSU tree and, consequently, should be transferred. They are Janusya adusta (Wörheide & Hooper, 1999) comb. nov., J. angusta (Van Soest & De Voogd, 2015) comb. nov. and J. tubuloreticulosa (Van Soest & De Voogd, 2015) comb. nov. Janusya angusta grouped with J. indica (bootstrap/posterior probability: 99%/1; p-distance: 9.5%) and Janusya adusta grouped with J. tubuloreticulosa (bootstrap/ posterior probability: 98%/1; p-distance: 10.1%). The variability among these branches ranged from 13.3% (between Janusya indica and J. tubuloreticulosa) to 14.7% (between *J. angusta* and *J. tubuloreticulosa*).

Clade G comprises the genera Ascandra Haeckel, 1872, Burtonulla Borojević & Boury-Esnault, 1986 and Levinella Borojević & Boury-Esnault, 1986. Because no Burtonulla nor Levinella representatives were available in the ITS tree, the genus Ascandra appeared monophyletic (bootstrap/posterior probability: 75%/1), but in the C-LSU, it was found non-monophyletic, with Levinella thalassae Borojević & Boury-Esnault, 1987, L. prolifera (Dendy, 1913) and Burtonulla sibogae Borojević & Boury-Esnault, 1986 being related to *Ascandra* species (bootstrap/posterior probability: 98%/1). Interestingly, one sequence identified as Ernstia chrysops Van Soest & De Voogd, 2018 also grouped with the Ascandra clade. Re-analysing the description of this species, we confirm that it is an *Ascandra*: **A.** chrysops (Van Soest & De Voogd, 2018) comb. nov.

The p-distances for clade Grange from 1.9% [between A. corallicola (Rapp, 2006) and A. falcata Haeckel, 1872] to 11.2% [between A. contorta (Bowerbank, 1866) and the new species A. oceanusvitae] for the ITS, and from 0.6% (between L. prolifera and B. sibogae) to 10.5% (between A. chrysops and A. contorta) for the C-LSU. Our results clearly indicate that Levinella and Burtonulla should be synonymized.

Among clade H, the genus Soleneiscus Borojević et al. 2002 was monophyletic in the C-LSU tree (bootstrap/posterior probability: 98%/1). Our new species, S. intermedius, grouped with S. radovani Wörheide & Hooper, 1999 in a clade with elevated support (bootstrap/posterior probability: 94%/1) and p-distance (5.4%). The other two species are sister to this clade. The p-distances of the genus ranged from 5.4 to 10.9% [between the new species S. intermedius and S. stolonifer (Dendy, 1891)].

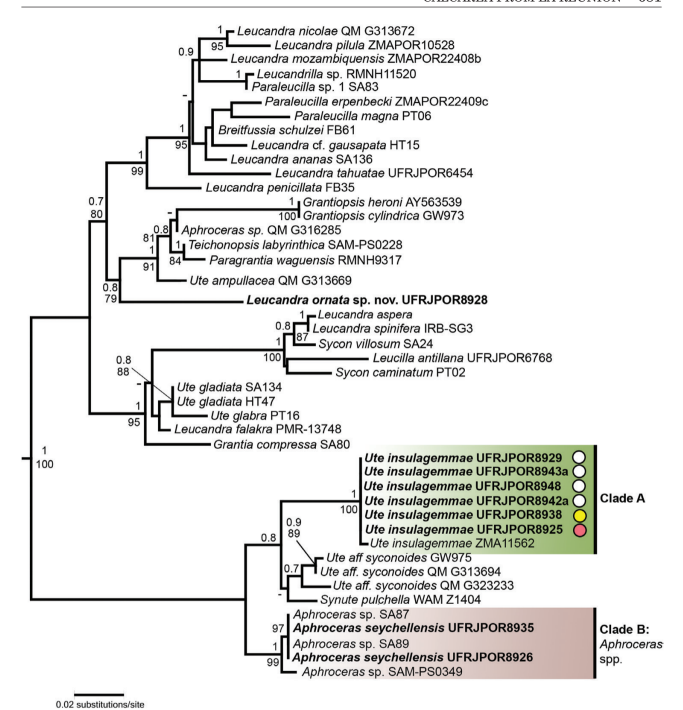
#### CALCARONEA PHYLOGENY (C-LSU)

In this study, we provided sequences for the new species *Leucandra ornata* (described below) and for the species *Aphroceras seychellensis* (Hôzawa, 1940) **comb. nov.** and *Ute insulagemmae* Van Soest & De Voogd, 2018. The obtained ML tree is shown in Figure 4. Corroborating previous results, the family Grantiidae Dendy, 1892 and the three genera were retrieved as non monophyletic. Two separated clades were defined for the cluster of species containing *A. seychellensis* (and other *Aphroceras* spp.) and one for *Ute insulagemmae*.

As a polyphyletic genus, Leucandra Haeckel, 1872 species were distributed in different clades within the subclass Calcaronea (e.g. Alvizu et al., 2018). Our sequence of Leucandra ornata appears as a distinct lineage that groups with a clade formed by Grantiopsis heroni Wörheide & Hooper, 2003, G. cylindrica Dendy, 1892, Ute ampullacea Wörheide & Hooper, 2003, Aphroceras sp., Teichonopsis labyrinthica (Carter, 1878) and Paragrantia waguensis Hôzawa, 1940 (bootstrap/posterior probability: 79%/0.8). These are members of Grantiidae, but also of the families Lelapiidae Dendy & Row, 1913, Jenkinidae Borojević et al., 2000 and Amphoriscidae Dendy, 1892. Therefore, the genus assignment of this species was mostly made according to morphology.

The other sequences of specimens from La Réunion form two well-defined clades. Clade A reunited specimens from La Réunion, morphologically identified by us as *Ute insulagemmae*, with the holotype of this species from the Seychelles (Van Soest & De Voogd, 2018). The clade was supported by high bootstrap and posterior probability values (100%/1) and the genetic *p*-distance did not exceed 0.3%, confirming the morphological identifications. Besides, the specimens included in the phylogenetic tree represented different chromotypes (white, yellow and pink) and these grouped together, showing that for this species, different colours represent intraspecific polymorphism.

Clade B contains the specimens of A. seychellensis from La Réunion and other Aphroceras sequences available from GenBank. Aphroceras seychellensis groups conspecifically with two specimens from



**Figure 4.** Maximum likelihood phylogenetic tree based on C-LSU sequences of the subclass Calcaronea. Midpoint-rooted tree. Species sequences generated in this work are in bold. Clades containing species found in this work are highlighted. Coloured circles represent different chromotypes (yellow, pink or white) of *Ute insulagemmae*. Support values are shown at the nodes (Bootstrap > 70%; posterior probability > 0.7).

St. Helena (South Atlantic; bootstrap/posterior probability: 97%/< 0.7). This clade groups with another specimen of *Aphroceras* from Tasmania (Australia; bootstrap/posterior probability: 99%/1).

The four specimens of *A. seychellensis* presented *p*-distance of 0, and the comparison between this clade and the specimen from Tasmania presented *p*-distance of 0.8%.

Morphological results
Phylum Porifera Grant, 1836
Class Calcarea Bowerbank, 1862
Subclass Calcinea Bidder, 1898
Order Clathrinida Hartman, 1958
Family Clathrinidae Minchin, 1900
Genus Janusya Gen. Nov.

Zoobank registration: urn:lsid:zoobank.org:act: 7F1DD58A-C896-4DF1-9DDC-817B4D30E400.

Type species: Janusya indica sp. nov.

*Etymology:* The new genus is dedicated to Jean Vacelet, who is affectionately known as Janus, in recognition for all his work on Porifera, including calcareous sponges.

Diagnosis: Calcinea in which the cormus comprises a typical clathroid body. A stalk may be present. The skeleton contains regular (equiangular and equiradiate) or subregular triactines and tetractines. Tetractines are less abundant than triactines, sometimes even rare, but they always have needle-like apical actines. Diactines may be added. The aquiferous system is asconoid.

# $\emph{J}$ ANUSYA INDICA SP. NOV.

(FIGS 5, 6; TABLE 2)

Zoobank registration: urn:lsid:zoobank.org:act: 79DA23AC-B3EE-42B0-8BB3-1DA5F592AF9A.

Etymology: Named in reference to the type locality located in the Indian Ocean.

*Diagnosis:* Yellowish to beige *Janusya* in which the cormus is incrusting to globulose, formed by regular and tightly anastomosed tubes with long, apical, water-collecting tube. The skeleton contains fusiform diactines, trichoxeas, triactines and rare tetractines.

Type locality: Passe de l'Hermitage, La Réunion, Indian Ocean.

Type material: Holotype – UFRJPOR 8945 (= 171109-PAE2-TP20). Paratype – UFRJPOR 8932 (= 171109-PAE2-TP6). Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

Additional material examined: UFRJPOR 8931 (= 171109-PAE2-TP5), UFRJPOR 8940 (= 171109-PAE2-TP16). Passe de l'Hermitage Reef,

La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

*Colour:* Sponge yellowish beige in life (Fig. 5A, B) and light brown in ethanol (Fig. 5C, D).

Morphology and anatomy: Sponge incrusting to globulose (Fig. 5). Cormus clathroid, formed by regular and tightly anastomosed tubes (Fig. 5). One or few apical oscula, surrounded by membrane (Fig. 5A). The osculum is a long water-collecting tube, i.e. a larger tube with osculum that receives the water from several other thinner tubes. Underneath the osculum of globulose specimens, there is a cavity that is not lined by a continuous membrane. Cells with yellow granules are present and distributed homogeneously in the mesohyl of the specimens UFRJPOR 8945 and UFRJPOR 8932 (Fig. 6A, B, respectivelly). The aquiferous system is asconoid and the skeleton is composed of diactines, trichoxeas, triactines and rare tetractines (Fig. 6C–F).

Spicules (Table 2)

*Diactines*: Thin, fusiform, but with different tips. They are undulated and slightly curved (Fig. 6C).

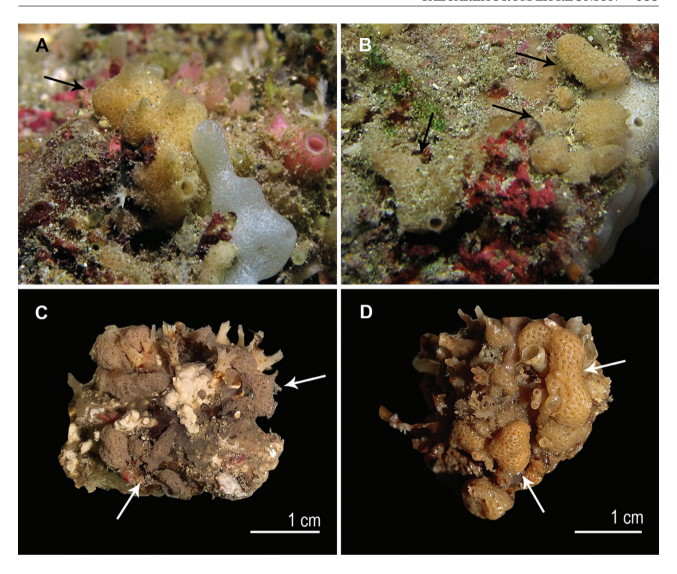
*Triactines:* Equiangular and equiradiate (Fig. 6D), but often the unpaired actine is shorter than the paired ones (Fig. 6E). Actines are conical with blunt to sharp tips.

Tetractines: Equiangular and equiradiate, but often the unpaired actine is shorter than the paired ones in the triactines Actines are conical with blunt to sharp tips (Fig. 6F). The apical actine is thin (needle-like), cylindrical, short, smooth and sometimes curved.

Ecology: Some individuals were associated with Leucaltis nodusgordii (UFRJPOR 8930) and Ute insulagemmae (Fig. 5A).

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: The new genus Janusya is being proposed to reunite the new species described here and other sponges previously assigned to Arturia. The first diagnosis of Arturia was: 'Calcinea in which the cormus comprises a typical clathroid body. A stalk may be present. The skeleton contains regular (equiangular and equiradiate) triactines and tetractines. However, tetractines are more rare. Diactines may be added. Asconoid aquiferous system' (Klautau et al., 2013). But it was modified later: 'Calcinea in which the cormus



**Figure 5.** Janusya indica A, specimen in vivo (UFRJPOR 8931), associated with Leucaltis nodusgordii (white sponge) and near Ute insulagemmae (pink sponge). B, several specimens in vivo (paratype; UFRJPOR 8932). C, fixed specimen (holotype; UFRJPOR 8945). D, fixed specimen (paratype; UFRJPOR 8932). Arrows point to specimens of J. indica gen. et sp. nov.

comprises a typical clathroid body. A stalk may be present. The skeleton contains regular (equiangular and equiradiate) triactines and few tetractines. Diactines may be added. Asconoid aquiferous system' (Fontana *et al.*, 2018).

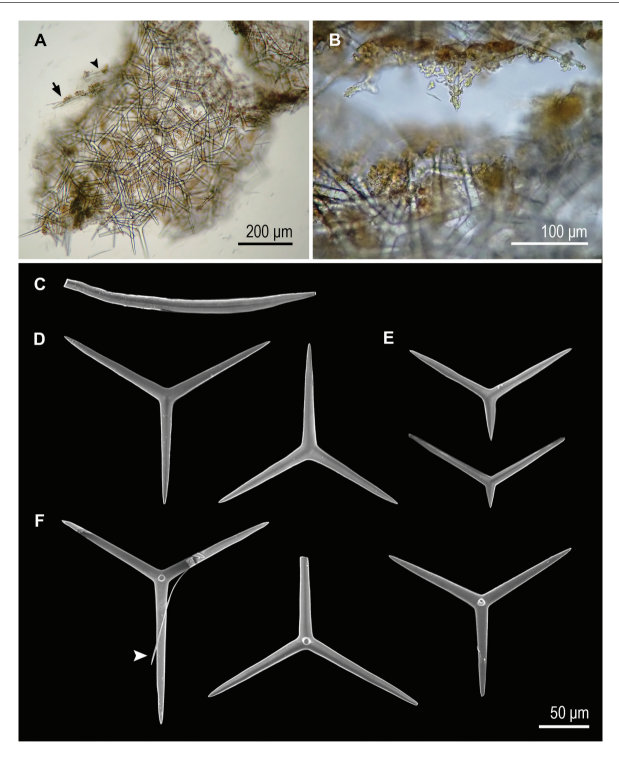
To clarify the difference between *Arturia* and *Janusya* we suggest a new modification in the diagnosis of *Arturia*: Calcinea in which the cormus comprises a typical clathroid body. A stalk may be present. The skeleton contains regular (equiangular and equiradiate) triactines and few tetractines. The apical actine of the tetractines is thinner than the basal ones, but it is conical, never needle-like. Diactines may be added. Asconoid aguiferous system (emend.).

Currently *Arturia* is composed of 15 species: *Arturia* adusta; A. africana (Klautau & Valentine, 2003);

A. alcatraziensis; A. canariensis (Miklucho-Maclay, 1868); A. compacta (Schuffner, 1877); A. darwinii (Haeckel, 1870); A. dubia (Dendy, 1891); A. hirsuta (Klautau & Valentine, 2003) – type species of the genus; A. passionensis (Van Soest et al., 2011); A. spirallata; A. sueziana; A. tenuipilosa (Dendy, 1905); A. trindadensis Azevedo et al., 2017; A. tubuloreticulosa; and A. vansoesti Cóndor-Luján et al., 2018.

Voigt & Wörheide (2016) proposed to include 'Clathrina' adusta, previously transferred to Ernstia by Klautau et al. (2013) in the genus Arturia. However, Voigt et al. (2017) showed in a molecular phylogeny that this species was actually not an Arturia and suggested that it could be a new genus.

Analysing morphologically all species of 'Arturia' presented in our tree, we found that A. alcatraziensis,



**Figure 6.** Janusya indica A, cross-section of the specimen UFRJPOR 8945. B, cross-section of the specimen UFRJPOR 8932 with an apical actine of a tetractine pointing towards the lumen of a tube. C, diactine. D, triactines. E, triactines with shorter unpaired actine. F, tetractines (arrowhead points to a trichoxea).

A. spirallata and A. sueziana have the apical actine of their tetractines only a little thinner than the basal ones, whereas A. adusta, A. angusta, A. tubuloreticulosa and Janusya indica have thin

(needle-like) apical actines. Because the type species of *Arturia* (*A. hirsuta*) has apical actines only a little thinner than the basal ones, we consider the clade of *A. alcatraziensis*, *A. spirallata* and *A. sueziana* as the

| Specimen         | Spicule    | Actine | Length | $(\mu m)$ |      |       | Width | (µm) |     |      | N  |
|------------------|------------|--------|--------|-----------|------|-------|-------|------|-----|------|----|
|                  |            |        | Min    | Mean      | SD   | Max   | Min   | Mean | SD  | Max  |    |
| UFRJPOR 8945 (H) | Diactine   |        | 270.3  | 396.4     | 68.2 | 497.3 | 10.8  | 14.8 | 3.2 | 21.6 | 15 |
| ,                | Triactine  |        | 105.3  | 122.3     | 12.7 | 145.8 | 10.8  | 13.7 | 1.3 | 16.2 | 20 |
|                  | Tetractine | Basal  | 97.2   | 122.3     | 14.6 | 143.1 | 10.8  | 13.6 | 1.3 | 16.2 | 20 |
|                  |            | Apical | 78.3   | 82.4      | 4.1  | 89.1  | 4.1   | 5.2  | 0.6 | 5.4  | 06 |
| UFRJPOR 8932 (P) | Diactine   | _      | 194.6  | 281.1     | 69.2 | 378.4 | 3.2   | 8.8  | 2.7 | 10.8 | 15 |
|                  | Triactine  |        | 121.5  | 145.3     | 16.7 | 180.9 | 13.5  | 17.0 | 1.7 | 20.3 | 20 |
|                  | Tetractine | Basal  | 121.5  | 145.1     | 19.3 | 202.5 | 13.5  | 17.3 | 1.8 | 18.9 | 20 |
|                  |            | Apical | 51.3   | 67.5      | 10.6 | 81.0  | 4.1   | 5.1  | 0.6 | 5.4  | 05 |

**Table 2.** Spicule measurements of *Janusya indica*. H = holotype, P = paratype

true clade of Arturia. Therefore, our new species can be confirmed as a new genus named here Janusya. Moreover, reassessing the descriptions of each species currently assigned to Arturia, we concluded that the new genus Janusya should include J. indica (type species) and five new combinations: J. adusta, J. angusta, J. trindadensis, J. tubuloreticulosa and. possibly, J. darwinii. The last is doubtful, because we did not find the type specimen of this species. Therefore, we used only the drawing published by Haeckel (1872). It is interesting to notice that three of them have a pseudoatrium: J. adusta, J. angusta and J. indica. Finally, Janusya indica can be differentiated from the other species of the genus, because it is the only species with diactines. Besides, the unpaired actine of its triactines and tetractines are usually shorter than the paired ones.

Sponges remaining assigned to Arturia include A. hirsuta (type species), A. africana, A. alcatraziensis, A. canariensis, A. compacta, A. dubia, A. spirallata, A. sueziana, A. tenuipilosa and A. vansoesti.

Family Dendyidae de Laubenfels, 1936 Genus *Soleneiscus* Borojević *et al.*, 2002

Type species: Leucosolenia stolonifer Dendy, 1891.

*Diagnosis:* Dendyidae that grow in the form of an individual ascon tube, with several ascon tubes growing upright from basal, stolon-like tubes, or in the form of creeping, ramified tubes (modified from Borojević *et al.*, 2002).

## SOLENEISCUS INTERMEDIUS SP. NOV.

(FIGS 7, 8; TABLE 3)

Zoobank registration: urn:lsid:zoobank.org:act:C9AE024F-7E3F-4B46-AD5F-A9E6704B59B8.

Diagnosis: Soleneiscus that grows in the form of ramified tubes at the base and whose skeleton is composed of diactines, rare triactines located only at the oscular region and two types of tetractines. The diactines are thin, maximum thickness of  $7.5~\mu m$ .

Etymology: From Latin intermedium, meaning intermediate, in between, for the close morphological similarity with other species.

Type locality: Portail Caves, La Réunion, Indian Ocean.

Type material: Holotype – UFRJPOR 8933 (= 171110ESB2TP6). Portail Caves, La Réunion, Indian Ocean, coll. T. Pérez, 10 November 2017, 18–20 m depth.

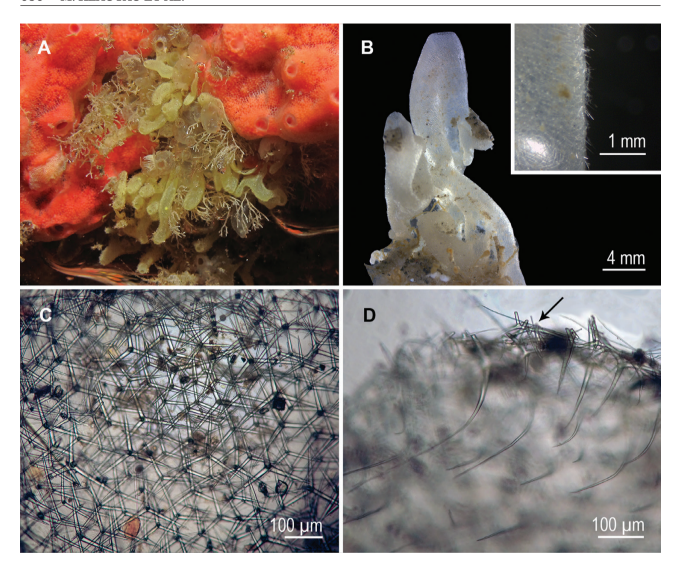
Colour: Sponge light yellow in life and white in ethanol (Fig. 7A, B).

Morphology and anatomy: Cormus formed by ramified and erect tubes, ending in a single osculum without a crown (Fig. 7A, B). Surface hispid due to the presence of diactines lying perpendicular to the surface (Fig. 7B). The lumen of the tubes is also hispid due to the long apical actines of the tetractines (Fig. 7C, D). Aquiferous system asconoid. Skeleton composed of diactines, two size categories of tetractines, and rare triactines (Fig. 8).

Spicules (Table 3)

*Diactines:* Thin, slightly curved and undulated. The tip that penetrates the cormus is thicker, but both of them are sharp (Fig. 8A).

*Triactines:* Regular to subregular, with variable size. They are rare. Actines are conical and sharp. They are present near the oscula (Fig. 8B).



**Figure 7.** Soleneiscus intermedius (holotype; UFRJPOR 8933). A, specimen in vivo living with a red Demospongiae, hydrozoans and ascidians. B, fixed specimen (external hispid surface of a tube in the inset). C, tangential section of a tube. D, lumen of a tube with the apical actine of tetractines. Arrow points to a diactine of the surface.

Tetractines I: Regular or sagittal, large. Actines are conical, sharp and curved (Fig. 8C).

Tetractines II: Regular or sagittal, a little smaller than tetractine I. Actines are slightly conical and sharp. The apical actines are only a little thinner than the basal ones. They are thin, sharp and curved. The apical actines are huge. They are only a little thinner than the basal ones at the base but become thin (Fig. 8D).

*Ecology:* The specimen was growing among a red undetermined sponge, colonies of hydrozoans and ascidians (Fig. 7A).

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: There are nine accepted species of Soleneiscus: S. apicalis (Brøndsted, 1931) from the Antarctic, S. hamatus Voigt et al., 2017 from the Red Sea, S. hispidus (Brøndsted, 1931) from the Antarctic, S. irregularis (Jenkin, 1908) from East Africa, S. japonicus (Haeckel, 1872) from Japan, S. olynthus (Borojević & Boury-Esnault, 1987) from the Bay of Biscay, Atlantic, S. pedicellatus Azevedo et al., 2015 from Peru, S. radovani from the Great Barrier Reef, Australia and S. stolonifer from South Australia. The new species Soleneiscus intermedius is morphologically similar to S. hamatus, S. irregularis and S. stolonifer, because the four species present diactines and two types of tetractines, one with large apical actine and another with thin apical actine. However, the new species has also triactines, but as they are rare and

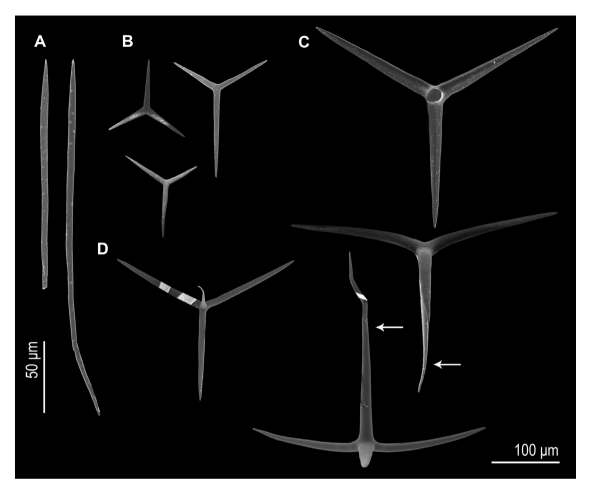


Figure 8. Soleneiscus intermedius (holotype; UFRJPOR 8933). A, diactines. B, triactines. C, tetractines I (arrows point to apical actines). D, tetractines II.

present only near the osculum, they could have been overlooked by the other authors. Nonetheless, even ignoring the presence or absence of triactines, our new species can be differentiated from the others by the size of some spicules (Table 3).

Comparing the new species with S. irregularis and S. stolonifer, the diactines of S. intermedius are much smaller and thinner (S. intermedius:  $250-445/5-7.5~\mu m; S.$  irregularis:  $300-800/16-28~\mu m; S.$  stolonifer:  $700/30~\mu m$ ). The diactines of S. intermedius are most similar to those of S. hamatus ( $136-274/5-8~\mu m$ ). However, these species are genetically distant (Fig. 4), showing that a similar morphology in this group does not, necessarily, represent conspecificity.

This observation also makes us question the identification of some specimens from the Great Barrier Reef (Australia), such as *S. stolonifer* (Wörheide & Hooper, 1999). In the original description of *S. stolonifer*, Dendy (1891) did not separate the tetractines in two different size categories. He mentioned variable

apical actines length of the tetractines (one developed and another short), but he did not separate them in different categories. Besides, the size of the diactines and the external morphology of the specimens from the Great Barrier Reef (Wörheide & Hooper, 1999) were different from those given by Dendy (1891) (see Table 3). These suggest that they are not conspecific, and that the specimens from the Great Barrier Reef probably represent a new species.

FAMILY LEUCALTIDAE DENDY & ROW, 1913 GENUS ASCANDRA HAECKEL, 1872

Type species: Ascandra falcata Haeckel, 1872.

Diagnosis: 'Calcinea with loosely anastomosed tubes. Tubes are free, at least in the apical region. The skeleton contains regular (equiangular and equiradiate) or sagittal triactines and tetractines. The apical actine is thin (needle-like) or thick at the base.

Table 3. Spicule measurements of Soleneiscus intermedius and other Soleneiscus species. H = holotype

| Species              | Spicule       | Actine          | Length (µm) | (mn   |       |        | Width (µm) | ım)     |      |      |    |
|----------------------|---------------|-----------------|-------------|-------|-------|--------|------------|---------|------|------|----|
|                      |               |                 | Min         | Mean  | SD    | Max    | Min        | Mean    | SD   | Max  | N  |
| S. intermedius sp.   | Diactine      |                 | 250.0       | 336.0 | 44.8  | 445.0  | 5.0        | 5.9     | 1.2  | 7.5  | 20 |
| nov. UFRJPOR         | Triactine     |                 | 105.0       | 116.3 | 14.6  | 145.0  | 10.0       | 11.9    | 2.5  | 15.0 | 90 |
| 8933 (H)             | Tetractine I  | Paired          | 190.0       | 227.8 | 21.2  | 260.0  | 20.0       | 23.3    | 2.9  | 30.0 | 20 |
|                      |               | Unpaired        | 135.0       | 221.8 | 39.7  | 285.0  | 15.0       | 23.0    | 4.1  | 30.0 | 20 |
|                      |               | Apical          | 75.0        | 230.5 | 175.8 | >625.0 | 15.0       | 19.8    | 4.4  | 30.0 | 20 |
|                      | Tetractine II | Paired          | 135.0       | 158.5 | 11.0  | 175.0  | 10.0       | 10.0    | 0    | 10.0 | 20 |
|                      |               | Unpaired        | 115.0       | 155.3 | 20.4  | 195.0  | 15.0       | 23.0    | 4.1  | 30.0 | 20 |
|                      |               | Apical          | 25.0        | 50.0  | 18.4  | >70.0  | 5.0        | 5.0     | 0    | 0.0  | 04 |
| S. irregularis after | Diactine      |                 | 300.0       | ı     | 1     | 800.0  | 16.0       | 21 - 24 | ,    | 28   | ı  |
| Jenkin 1908          | Tetractine I  |                 | 150.0       | ı     | 1     | 220.0  |            | 20.0    | ,    |      | ı  |
|                      |               | Apical          | 210.0       | ı     | 1     | 260.0  | 16.0       | 1       | ,    | 20   | ı  |
|                      | Tetractine II |                 | 100.0       | ı     | 1     | 200.0  | 10.0       | 1       |      | 16   | ı  |
|                      |               | Apical          | 120.0       | ı     | 1     | 150.0  |            | 7.0     |      |      | ı  |
| S. stolonifer after  | Diactine      |                 | ı           | 700.0 | 1     | ı      |            | 5.7     |      |      | ,  |
| Dendy 1891           | Tetractine    | Basal           | ı           | 200.0 | 1     | ı      |            | 15.0    |      |      | ,  |
|                      |               | Apical          | ı           | 1     | 1     | 500.0  |            | 1       |      |      | ,  |
| S. stolonifer after  | Diactine      |                 | 150.0       | 207.8 | 1     | 260.0  | 4.0        | 5.7     |      | 8.0  | 25 |
| Wörheide &           | Tetractine I  | Basal           | 80.0        | 129.9 | 1     | 170.0  | 8.0        | 12.2    |      | 21.0 | 25 |
| Hooper 1999          |               | Apical          | ı           | 1     | 1     | 500.0  |            | 1       |      |      | ,  |
|                      | Tetractine II | Paired          | 40.0        | 81.6  | 1     | 110.0  | 8.0        | 9.7     |      | 12.0 | 25 |
|                      |               | Unpaired        | 80.0        | 124.4 | 1     | 160.0  | 8.0        | 9.7     |      | 12.0 | 25 |
| S. hamatus after     | Diactine      |                 | 136.0       | 207.6 | 42.6  | 274.0  | 5.0        | 6.2     | 8.0  | 8.0  | 10 |
| Voigt et al. $2017$  | Tetractine I  | Basal (Regular) | 84.0        | 179.5 | 49.1  | 291.0  | 9.0        | 13.0    | 2.06 | 19.0 | 20 |
|                      |               | Paired          | 84.2        | 156.6 | 32.3  | 272.0  | 10         | 13.5    | 2.8  | 19.0 | 16 |
|                      |               | Unpaired        | 199.0       | 251.1 | 24.1  | 291.0  | 9.0        | 13.5    | 3.1  | 19.0 | 16 |
|                      |               | Apical          | 120.0       | 147.3 | 16.1  | 167.0  | 13.0       | 18.1    | 4.0  | 27.0 | 11 |
|                      | Tetractine II | Basal (Regular) | 0.89        | 134.2 | 37.0  | 235.0  | 5.0        | 8.7     | 2.4  | 15.0 | 20 |
|                      |               | Paired          | 80.0        | 117.7 | 19.0  | 152.0  | 5.0        | 8.9     | 2.4  | 14.0 | 13 |
|                      |               | Unpaired        | 115.0       | 179.6 | 31.9  | 235.0  | 5.0        | 0.6     | 2.3  | 15.0 | 13 |
|                      |               | Apical          | 35.0        | 41.8  | 3.2   | 49.0   | 3.0        | 3.5     | 9.0  | 5.0  | 20 |
|                      |               |                 |             |       |       |        |            |         |      |      |    |

Diactines may be added. Asconoid aquiferous system' (Klautau *et al.*, 2016).

#### ASCANDRA MASCARENICA SP. NOV.

(FIGS 9, 10; TABLE 4)

Zoobank registration: urn:lsid:zoobank.org:act:774A6B27-B4E3-4A85-830F-4BCFB2869574.

*Diagnosis:* Ascandra whose cormus is loosely anastomosed and has apical water-collecting tubes. Skeleton composed of diactines with one lanceolated tip, triactines and two size categories of tetractines. Apical actines are conical.

*Etymology:* The species name refers to ecoregion of the type locality, the Mascarene Islands.

Type locality: Passe de l'Hermitage, La Réunion, Indian Ocean.

Type material: Holotype – UFRJPOR 8936 (= 171109-PAE2-TP12), Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

Colour: White in life and light brown to white in ethanol (Fig. 9A, B).

Morphology and anatomy: Sponge formed by loosely anastomosed tubes (Fig. 9A, B). Large water-collecting tubes are present at the apical region (Fig. 9B). Surface hispid (Fig. 9C). Aquiferous system asconoid. Skeleton composed of tetractines of two sizes, diactines and few triactines (Fig. 9D). The diactines are perpendicular to the tubes.

Spicules (Table 4)

*Diactines:* One tip sharp and the other lanceolated. They are slightly curved and undulated (Fig. 10A).

*Triactines:* Equiangular and equiradiate. They are similar to the tetractines II, but they are less abundant. Actines are slightly conical and sharp (Fig. 10B).

*Tetractines I:* Equiangular and equiradiate. Actines are conical and sharp. The apical actine is conical, sharp, shorter than the basal ones and smooth (Fig. 10C).

Tetractines II: Equiangular and equiradiate. They are smaller than the tetractines I. Actines are slightly conical and sharp (Fig. 10D). The apical actine is conical but much thinner than in the tetractines I, sharp and smooth.

*Ecology:* This species was found protected from the sunlight and associated with a bryozoan, a hydrozoan and other sponges (Fig. 9A).

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: Up to date, seven species of Ascandra with diactines have been described: A. atlantica (Thacker, 1908) from Cape Verde, A. contorta from the Mediterranean Sea, A. densa Haeckel, 1872 from Australia, A. falcata from the Adriatic Sea, A. izuensis (Tanita, 1942) from Japan, A. minchini Borojević, 1966 from the Mediterranean Sea and A. sertularia Haeckel, 1872 from the Indian Ocean (Java).

Considering only the shape of the diactines, the new species can be differentiated from six out of the seven species with diactines. While the diactine of A. mascarenica has one lanceolated tip, A. atlantica, A. contorta and A. izuensis have fusiform diactines with sharp tips, A. densa also has fusiform diactines with one end thicker than the other, A. falcata has sickleshaped diactines with one rounded tip and A. minchini only has microdiactines with a thickening in the midle.

The remaining species, A. sertularia, is probably not an Ascandra. According to the drawings provided by Haeckel (1872), this species has only sagittal spicules, suggesting that it most probably belongs to the genus Leucosolenia. Dendy & Row (1913) already transferred this species to Leucosolenia, but they made this transference because of the asconoid aquiferous system of this species.

## ASCANDRA OCEANUSVITAE SP. NOV.

(FIGS 11, 12; TABLE 5)

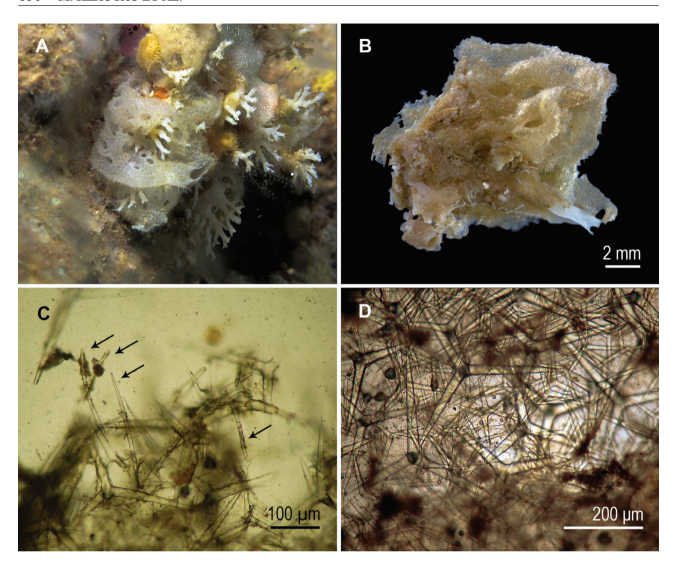
ZooBank registration: urn:lsid:zoobank.org:act:05205855-E843-4E38-B3EF-3DA6548EA5B2.

*Diagnosis:* Ascandra whose cormus is ramified, having only few anastomoses. Water-collecting tubes are not present. Skeleton composed of fusiform diactines, frequently with one tip larger, two size categories of triactines and of tetractines.

*Etymology:* For the Vie Océane Association, one of the institutions responsible for organizing the taxonomy course that allowed the collection of the sponges published here. The name is Latinized to *oceanus* (=ocean) and *vita* (=life).

Type locality: Passe de l'Hermitage, La Réunion, Indian Ocean.

Type material: Holotype - UFRJPOR 8944 (= 171109-PAE2-TP19). Paratype - UFRJPOR 8946



**Figure 9.** Ascandra mascarenica (holotype; UFRJPOR 8936). A, specimen in vivo, living among other sponges, bryozoans, and hydrozoans. B, fixed specimen. C, cross-section of a tube (arrows point to diactines). D, tangential section of the surface.

(= 171109-PAE2-TP21). Other material – UFRJPOR 8947 (= 171109-PAE2-TP22). Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

 ${\it Colour:}$  White in life and light brown or grey in ethanol (Fig. 11A, B).

Morphology and anatomy: Sponge formed by ramified tubes with few anastomoses and no water-collecting tubes (Fig. 11A, B). The surface of the tubes is hispid because of the presence of perpendicular diactines (Fig. 11C). Aquiferous system asconoid. The skeleton is composed of diactines, two size categories of triactines and tetractines. The larger triactines are rare. Diactines lie perpendicular to the tubes. The

small triactines are the most abundant spicules (Fig. 11D).

## Spicules (Table 5)

*Diactines:* Fusiform but with one tip thicker than the other and frequently curved. The thicker tip penetrates the sponge body (Fig. 12A).

*Triactines I:* Equiangular and equiradiate to subregular. It is the largest category of triactine. They are similar to tetractine II. Actines are conical and sharp. They are rare.

*Triactines II:* Equiangular and equiradiate. Actines are slightly conical and sharp. They are the most abundant spicules (Fig. 12B).

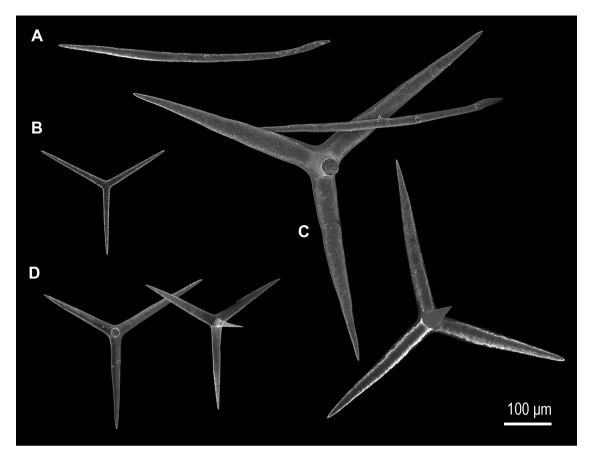


Figure 10. Ascandra mascarenica (holotype; UFRJPOR 8936). A, lanceolated diactines. B, triactine. C, tetractines I. D, tetractines II.

**Table 4.** Spicule measurements of *Ascandra mascarenica*. H = holotype

| Specimen         | Spicule       | Actine | Length | (µm)  | ,    | ,     | Width | (μm) |     | ,    | N  |
|------------------|---------------|--------|--------|-------|------|-------|-------|------|-----|------|----|
|                  |               |        | Min    | Mean  | SD   | Max   | Min   | Mean | SD  | Max  |    |
| UFRJPOR 8936 (H) | Diactine      |        | 200.0  | 328.0 | 91.8 | 530.0 | 10.0  | 13.5 | 3.3 | 20.0 | 20 |
|                  | Triactine     |        | 125.0  | 160.3 | 18.1 | 190.0 | 10.0  | 14.3 | 2.4 | 20.0 | 20 |
|                  | Tetractine I  | Basal  | 270.0  | 356.5 | 79.8 | 500.0 | 30.0  | 35.5 | 6.0 | 50.0 | 20 |
|                  |               | Apical | 90.0   | 138.4 | 39.6 | 210.0 | 30.0  | 32.6 | 5.6 | 50.0 | 19 |
|                  | Tetractine II | Basal  | 140.0  | 168.3 | 22.3 | 220.0 | 15.0  | 17.1 | 1.9 | 20.0 | 20 |
|                  |               | Apical | 42.5   | 109.9 | 75.0 | 350.0 | 5.0   | 6.1  | 1.7 | 10.0 | 20 |

Tetractines I: Equiangular and equiradiate to subregular. This is the largest category of tetractine. Basal actines are conical and sharp. The apical actine is short, conical, sharp and smooth (Fig. 12C).

Tetractines II: Equiangular and equiradiate. Actines are slightly conical and sharp (Fig. 12D). The apical actine is short, conical, sharp and smooth.

*Ecology:* Specimens found in a submarine cave.

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: Already considering that A. sertularia is in fact Leucosolenia sertularia, Ascandra oceanusvitae is the eighth species of

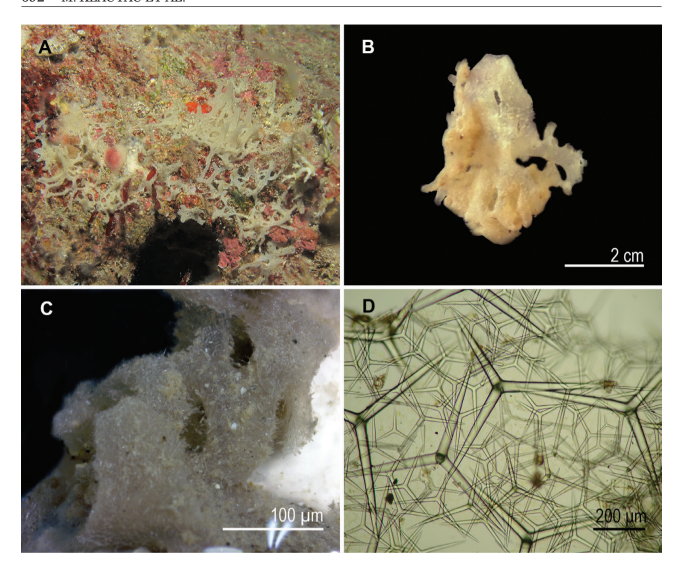


Figure 11. Ascandra oceanusvitae (holotype; UFRJPOR 8944). A, specimen in vivo. B, fixed specimen. C, hispid surface. D, tangential section of a tube.

Ascandra with diactines. As its diactines are fusiform, it can be differentiated from A. falcata, A. mascarenica and A. minchini, which do not have fusiform diactines. The fusiform diactines of A. oceanusvitae are similar to those of A. atlantica, A. contorta, A. densa and A. izuensis.

The new species can be differentiated from *A. contorta* and *A. densa* by, for example, the cormus, which is more tightly anastomosed in *A. contorta* and *A. densa*. Besides, it can also be differentiated by the presence of only one category of triactines and tetractines in both species, while *A. oceanusvitae* has two categories of triactines and of tetractines.

Consequently, the most similar species to the new one are *A. atlantica* from Cape Verde and *A. izuensis* from Japan. Although similar, they can be differentiated. *Ascandra oceanusvitae* has two size categories of

triactines, while *A. atlantica* and *A. izuensis* have only one category (the equivalent to the small category). Even though the largest category of triactine is rare in *A. oceanusvitae*, this new species can also be differentiated from the other two species by the size of the other spicules, which are much larger in the new species (Table 5).

GENUS LEUCALTIS HAECKEL, 1872

Type species: Leucaltis clathria Haeckel, 1872.

Diagnosis: 'Leucaltidae with a body composed of large, ramified and anastomosed tubes. Each tube has a distinct cortex, a choanoderm composed of elongated and ramified choanocyte chambers, and a central

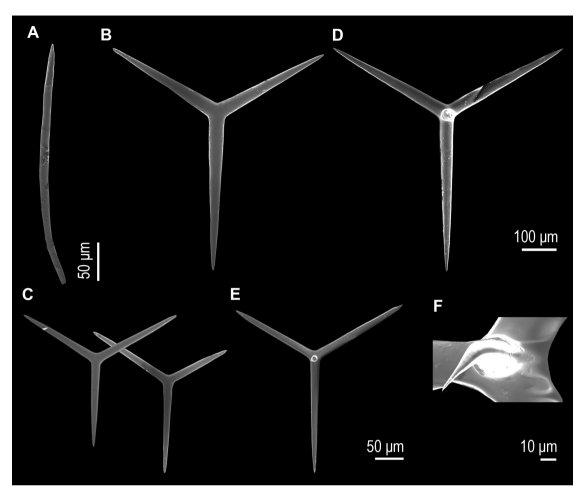


Figure 12. Ascandra oceanusvitae (holotype; UFRJPOR 8944). A, diactine. B, triactine I. C, triactines II. D, tetractine I. E, tetractine II. F, apical actine of a tetractine II.

atrium. The choanoderm and the atrial wall have a secondary skeleton composed of small triactines and tetractines' (Borojević *et al.*, 2002).

# LEUCALTIS NODUSGORDII (POLÉJAEFF, 1883) (FIGS 13, 14; TABLE 6)

Synonyms: Heteropegma nodusgordii Poléjaeff, 1883: 45 (only the Torres Strait specimens); von Lendenfeld, 1885: 1107 (only the Australian specimens); Dendy, 1892: 113; Dendy, 1905: 230; Jenkin, 1908: 453. Leucaltis bathybia var. mascarenica Ridley, 1884: 628 (to be confirmed). Clathrina latitubulata Carter, 1886: 515 (to be confirmed). Heteropegma latitubulata Dendy, 1892: 114 (to be confirmed). Leucaltis clathria Dendy, 1913: 16; Dendy & Frederick, 1924: 483; Hôzawa, 1940: 136; Tanita, 1943: 393; Lévi, 1998: 75; Wörheide & Hooper, 1999: 876; Borojević & Klautau, 2000: 190. Leuconia paloensis (non Tanita, 1943), Colin & Arneson, 1995: 61; Gosliner et al., 1996: 17; Erhardt

& Baensch, 1998: 21. Leucaltis nodusgordii Van Soest & De Voogd, 2015: 39, 2018: 66.

Type locality: Torres Strait, Australia.

Material examined: UFRJPOR 8930 (= 171109-PAE2-TP4). Photos: TP6029-6039. Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18-20 m depth.

Colour: White alive and light beige after fixation (Fig. 13A, B).

Morphology and anatomy: Sponge formed by large, individualized, anastomosed tubes (Fig. 13A, B). Each tube has its own cortex. Oscula located at the top of elevated tubes (Fig. 13A, B). The specimen was full of reproductive elements (Fig. 13C, D). The aquiferous system is composed of irregular, elongated and ramified choanocyte chambers

**Table 5.** Spicule measurements of  $Ascandra\ oceanusvitae$ ,  $Ascandra\ atlantica\ (after\ Klautau\ \&\ Valentine,\ 2003)$  and  $Ascandra\ izuensis$ . H=holotype, P=paratype

| Specimen               | Spicule       | Actine | Length | (µm)  |       |       | Width | $(\mu m)$ |      |       | N  |
|------------------------|---------------|--------|--------|-------|-------|-------|-------|-----------|------|-------|----|
|                        |               |        | Min    | Mean  | SD    | Max   | Min   | Mean      | SD   | Max   |    |
| UFRJPOR 8944 (H)       | Diactine      |        | 216.2  | 374.6 | 81.4  | 562.1 | 8.1   | 11.8      | 1.9  | 16.2  | 20 |
|                        | Triactine I   |        | 113.6  | 245.7 | 128.9 | 432.4 | 27.0  | 36.8      | 6.1  | 43.2  | 03 |
|                        | Triactine II  |        | 151.2  | 185.1 | 16.1  | 216.0 | 13.5  | 17.6      | 1.9  | 21.6  | 20 |
|                        | Tetractine I  | Basal  | 345.6  | 462.2 | 62.3  | 583.2 | 32.4  | 47.0      | 6.3  | 54.0  | 20 |
|                        |               | Apical | 108.1  | 141.7 | 19.9  | 162.2 | 27.0  | 45.8      | 24.1 | 110.3 | 11 |
|                        | Tetractine II | Basal  | 140.4  | 189.1 | 22.2  | 232.2 | 13.5  | 18.2      | 2.5  | 21.6  | 20 |
|                        |               | Apical | 10.8   | 14.2  | 3.4   | 18.9  | 1.4   | 2.0       | 0.8  | 2.7   | 04 |
| UFRJPOR 8946 (P)       | Diactine      | _      | 275.0  | 375.7 | 45.9  | 435.0 | 10.0  | 11.7      | 2.4  | 15.0  | 15 |
|                        | Triactine I   |        | -      | -     | -     | -     | -     | -         | -    | -     | -  |
|                        | Triactine II  |        | 157.5  | 176.3 | 17.7  | 230.0 | 12.5  | 13.9      | 2.4  | 22.5  | 20 |
|                        | Tetractine I  | Basal  | 202.7  | 314.9 | 53.5  | 418.9 | 24.3  | 32.7      | 4.9  | 40.5  | 20 |
|                        |               | Apical | -      | -     | -     | -     | -     | -         | -    | -     | -  |
|                        | Tetractine II | Basal  | 160.0  | 184.5 | 14.1  | 205.0 | 12.5  | 14.5      | 1.5  | 17.5  | 20 |
|                        |               | Apical | 15.0   | 16.3  | 1.8   | 17.5  | 5.0   | 5.0       | 0.0  | 5.0   | 2  |
| Ascandra atlantica (H) | Diactine      |        | 125.0  | 201.8 | 37.5  | 250.0 | -     | 7.5       | 0.8  | -     | 19 |
|                        | Triactine     |        | 75.0   | 108.3 | 16.5  | 140.0 | -     | 10.0      | 1.5  | -     | 30 |
|                        | Tetractine I  | Basal  | 87.5   | 165.0 | 30.5  | 225.0 | -     | 23.0      | 4.0  | -     | 30 |
|                        |               | Apical | 40.8   | 69.1  | 23.8  | 102.0 | -     | 18.9      | 7.0  | -     | 13 |
|                        | Tetractine II | Basal  | 85.0   | 106.6 | 13.5  | 145.0 | -     | 9.8       | 1.5  | -     | 30 |
|                        |               | Apical | 10.2   | 35.0  | 13.4  | 51.0  | -     | 7.8       | 2.8  | -     | 14 |
| Ascandra izuensis (H)  | Diactine      | _      | 270.0  | -     | -     | 550.0 | 17.0  | -         | -    | 30.0  | -  |
|                        | Triactine     |        | 85.0   | -     | -     | 120.0 | 8.0   | -         | -    | 14.0  | -  |
|                        | Tetractine I  | Basal  | 120.0  | -     | -     | 165.0 | 14.0  | -         | -    | 18.0  | -  |
|                        |               | Apical | 140.0  | -     | _     | 190.0 | 10.0  | -         | _    | 14.0  | -  |
|                        | Tetractine II | Basal  | 60.0   | -     | -     | 95.0  | 8.0   | -         | -    | 10.0  | -  |
|                        |               | Apical | 70.0   | -     | -     | 100.0 | 6.0   | -         | -    | 8.0   | -  |

(Fig. 13D). There is a thick cortex composed of large triactines and tetractines, tangential to the surface (Fig. 13D, E). The choanosome has few spicules, tiny triactines and tetractines (Fig. 13D). The apical actine of the cortical tetractines penetrates the choanosome and, sometimes, the atrium. The atrial skeleton is composed of tiny triactines and tetractines (Fig. 13F).

## Spicules (Table 6)

Cortical triactines: Large, equiangular and equiradiate or sagittal, with a curve between the paired actines. Actines are conical and sharp (Fig. 14A).

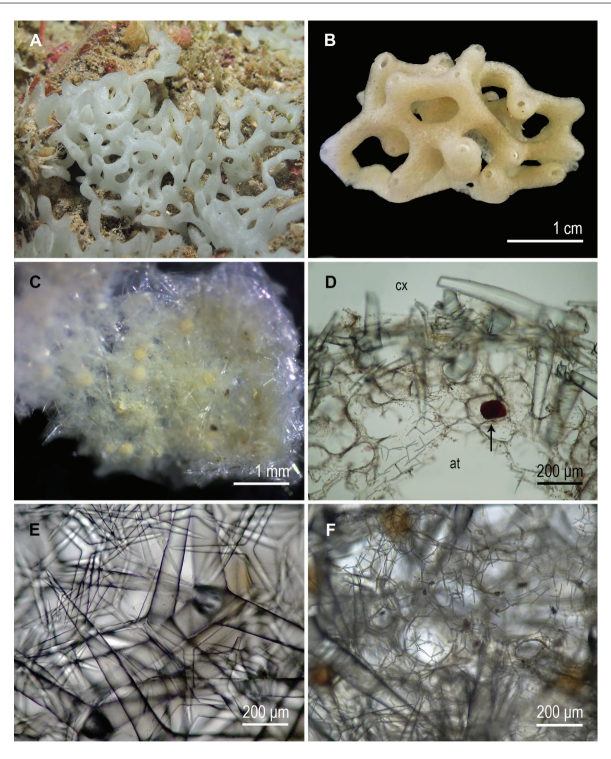
Cortical tetractines: Large, equiangular and equiradiate or sagittal. Actines are conical and sharp. The apical actine is longer than the basal ones (Fig. 14B).

Choanosomal and atrial triactines: Tiny, equiangular and equiradiate (Fig. 14C) or sagittal (Fig. 14D). Actines are cylindrical with rounded tips.

Choanosomal and atrial tetractines: Tiny, equiangular and equiradiate (Fig. 14E) or sagittal (Fig. 14F). Actines are cylindrical with rounded tips. The apical actine of the tetractines is conical and sharp and can be long.

Ecology: The specimen from Réunion was associated with two white nudibranchs and specimens of *J. indica* (UFRJPOR 8931) and *Ute insulagemmae* (Fig. 5A). *Leucaltis nodusgordii* has been found from 0 to 88 m depth in the Seychelles (Van Soest & De Voogd, 2018). Its colour alive varies from white to pink and lavender.

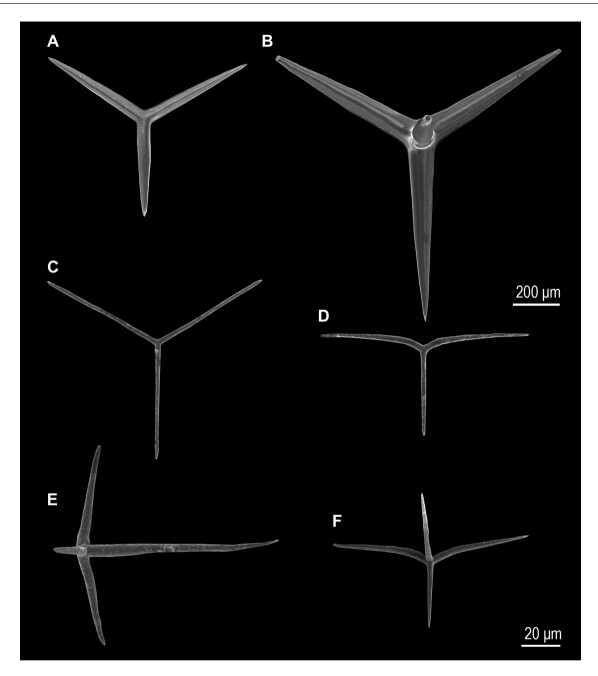
Geographic distribution: Torres Strait Northern Great Barrier Reef Ecoregion – Torres Strait (Poléjaeff,



**Figure 13.** *Leucaltis nodusgordii*. (UFRJPOR 8930). A, specimen *in vivo*. B, fixed specimen. C, choanosome full of yellow reproductive elements. D, cross-section (arrow points to a reproductive element). E, tangential section of the cortex. F, tangential section of the atrial wall. cx = cortex, at = atrium.

1883; von Lendenfeld, 1885; Dendy, 1892). Houtman Ecoregion – Houtman Abrolhos (Dendy & Frederick, 1924). Bassian Ecoregion – near Port Phillip Heads (Carter, 1886; Dendy, 1892); Bass Strait, Tasmania;

Westernport Bay, Victoria (Wörheide & Hooper, 1999). Central and Southern Great Barrier Reef Ecoregion – Egmont Reef (Dendy, 1913); Heron Island, GBR (Wörheide & Hooper, 1999). South India and Sri



**Figure 14.** Leucaltis nodusgordii. (UFRJPOR 8930). A, cortical triactine. B, cortical tetractine. C, regular choanosomal triactine. D, sagittal choanosomal or atrial triactine. E, regular choanosomal tetractine. F, sagittal choanosomal or atrial tetractine.

Lanka Ecoregion – Sri Lanka (Dendy, 1905); East African Coral Coast – Wasin (Jenkin, 1908); Seychelles Ecoregion – Amirantes group (Ridley, 1884; Dendy; 1913; Hôzawa, 1940; Van Soest & De Voogd, 2018); Mahé (Van Soest & De Voogd, 2018). Cargados Carajos/Tromelin Island Ecoregion – Cargados Carajos Atoll (Dendy, 1913). South Kuroshio – Naha, Okinawa Prefecture, Japan (Tanita, 1943). New Caledonia Ecoregion – Canal Woodin; Le Banc Gail (Borojević

& Klautau, 2000). Sulawesi Sea/Makassar Strait Ecoregion—North Sulawesi; Solomon Sea Ecoregion — Papua New Guinea (Van Soest & De Voogd, 2015). East African Coral Coast Ecoregion — Zanzibar (Van Soest & De Voogd, 2015). Mascarene Islands Ecoregion — La Réunion (this work).

Taxonomic remarks: Since Dendy (1913), Heteropegma nodusgordii was considered a synonym of L. clathria.

| Table 6. | Spicule measurements | of Leucaltis | nodusgordii ( | (UFRJPOR 8930) |
|----------|----------------------|--------------|---------------|----------------|
|----------|----------------------|--------------|---------------|----------------|

| Specimen     | Spicule       | Actine   | Length | (µm)  |       |       | Width | (µm) |      |       | N  |
|--------------|---------------|----------|--------|-------|-------|-------|-------|------|------|-------|----|
|              |               |          | Min    | Mean  | SD    | Max   | Min   | Mean | SD   | Max   |    |
| UFRJPOR 8930 | Triactine I   | Paired   | 210.0  | 368.3 | 72.1  | 475.0 | 30.0  | 46.8 | 7.8  | 60.0  | 20 |
|              |               | Unpaired | 225.0  | 374.5 | 81.9  | 500.0 | 25.0  | 48.3 | 9.6  | 60.0  | 20 |
|              | Tetractine I  | Basal    | 320.0  | 685.0 | 161.5 | 950.0 | 60.0  | 93.0 | 20.0 | 120.0 | 20 |
|              |               | Apical   | 160.0  | 265.0 | 73.3  | 320.0 | 70.0  | 72.5 | 5.0  | 80.0  | 04 |
|              | Triactine II  | Paired   | 50.0   | 61.5  | 4.9   | 70.0  | 2.5   | 2.6  | 0.6  | 5.0   | 20 |
|              |               | Unpaired | 25.0   | 60.1  | 10.5  | 72.5  | 2.5   | 2.6  | 0.6  | 5.0   | 20 |
|              | Tetractine II | Paired   | 40.0   | 63.2  | 7.0   | 72.5  | 2.5   | 3.6  | 1.3  | 5.0   | 20 |
|              |               | Unpaired | 42.5   | 60.8  | 10.5  | 87.5  | 2.5   | 3.6  | 1.3  | 5.0   | 20 |
|              |               | Apical   | 15.0   | 22.5  | 9.4   | 37.5  | 2.5   | 4.0  | 1.3  | 5.0   | 10 |

Nonetheless, Klautau et al. (2013) showed that specimens from Australia identified as L. clathria were molecularly not conspecific with those from the Caribbean, type locality of this species. Later, considering this molecular analysis and their observation of large (cortical) sagittal triactines only in Caribbean specimens of L. clathria, Van Soest & De Voogd (2015) also concluded that specimens from the Indo-Pacific and the Atlantic were not conspecific. Consequently, they decided to name the Indo-Pacific specimens L. nodusgordii. In another paper, Van Soest & De Voogd (2018) maintained their proposition, although they could not find well-resolved clades separating these two species in their phylogenetic tree. This may be related to the use of C-LSU marker, which is too conserved for separating closely related species such as L. clathria and L. nodusgordii, which has been confirmed by our molecular analyses (Fig. 2), whereas a more variable marker such as ITS allowed to clearly separate clades (Fig. 3).

However, morphologically differentiating *L. nodusgordii* from *L. clathria* remains a challenge. Van Soest & De Voogd (2015) stated that these species could be differentiated by the presence of large (cortical) sagittal triactines, which were present only in *L. clathria*. Still, these large (cortical) sagittal triactines and tetractines were found in the specimen from Réunion (*L. nodusgordii*). On the other hand, atrial tetractines with long apical actines were also found in our specimen, and on a picture from a specimen analysed by Van Soest & De Voogd (2018: fig. 37E). And so far, such an unusual trait could not be observed by us in specimens of *L. clathria* in previous studies.

The record of *L. nodusgordii* in La Réunion is particularly interesting because as until recently (Van Soest & De Voogd, 2015), this species was considered a synonym of *L. clathria*, we did not know

if the distribution of *L. nodusgordii* was limited to the Eastern part of the Indian Ocean or if this species occurred also further West. With our study, we confirmed that *L. nodusgordii* is widely distributed along the Indian and Pacific Oceans, while *L. clathria* seems restricted to the Atlantic.

Family Leucascidae Dendy, 1892 Genus *Leucascus* Dendy, 1892

Type species: Leucascus simplex Dendy, 1892.

Diagnosis: 'Copiously branched and anastomosed choanocyte tubes covered by a continuous membrane. The exhalant aquiferous system is represented by a well-developed atrium delimited by a membrane with no choanoderm, supported by a specific skeleton of triactines and/or tetractines. The size of the spicules of the choanocyte tubes and of the cortical and atrial membranes is similar. Apical actine of the tetractines ornamented with spines. Solenoid aquiferous system' (Cavalcanti et al., 2013).

## LEUCASCUS TENUISPINAE SP. NOV.

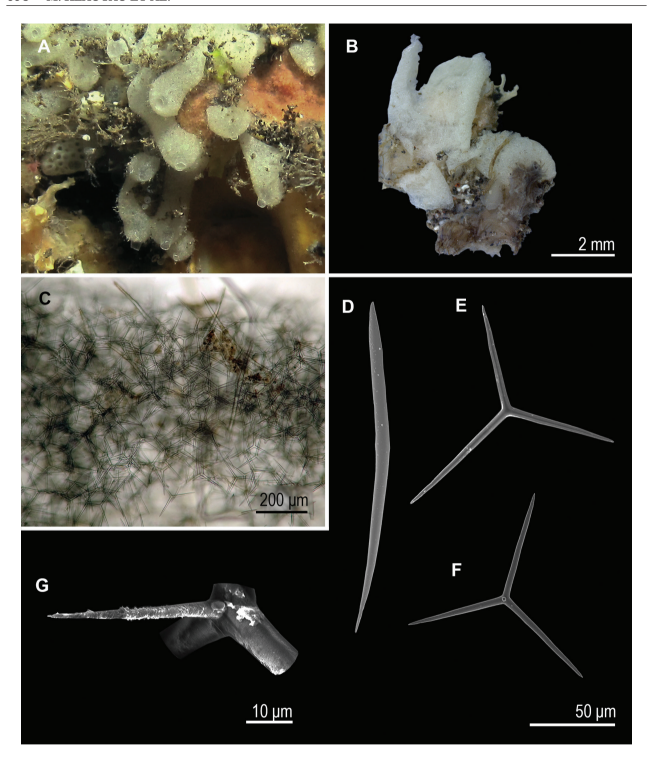
(FIG. 15; TABLE 7)

Zoobank registration: urn:lsid:zoobank. org:act:4EB114CF-5870-4338-95FE-09BE9C398196.

*Diagnosis:* Subglobular *Leucascus.* S skeleton composed of fusiform diactines, triactines and rare tetractines with short spines.

*Etymology:* For the microspines on the apical actine of the tetractines (*tenues* = thin; *spinae* = spines).

Type locality: Portail Caves, La Réunion, Indian Ocean.



**Figure 15.** Leucascus tenuispinae (holotype; UFRJPOR 8927). A, specimen in vivo. B, fixed specimen. C, tangential section of the surface. D, diactine. E, triactine. F, tetractine. G, apical actine of a tetractine with short spines.

8.2

8.1

2.8

0.7

0.0

0.3

10.8

8.1

4.1

20

20

17

| Specimen         | Spicule  | Actine | Length | (µm)  |      |       | Width | (μm) |     |      | $\overline{N}$ |
|------------------|----------|--------|--------|-------|------|-------|-------|------|-----|------|----------------|
|                  |          |        | Min    | Mean  | SD   | Max   | Min   | Mean | SD  | Max  |                |
| UFRJPOR 8927 (H) | Diactine |        | 324.3  | 433.2 | 97.7 | 648.6 | 13.5  | 16.8 | 2.6 | 21.6 | 14             |

105.0

105.2

72.3

7.3

7.1

11.8

121.5

116.1

97.2

6.8

8.1

2.7

94.5

94.5

51.3

**Table 7.** Spicule measurements of *Leucascus tenuispinae*. H = holotype

Basal

Apical

Type material: Holotype – UFRJPOR 8927 (= 171110-ESB2-TP2). Photos: TP6114-6116. Portail Caves, La Réunion, Indian Ocean, coll. T. Pérez, 10 November 2017, 25 m depth.

Triactine

Tetractine

*Colour:* White alive and after fixation (Fig. 15A).

Morphology and anatomy: Sponge subglobular with a single apical osculum with membrane (Fig. 15A). A cortical membrane covers the anastomosed tubes. It was not possible to clearly observe the anastomosis because the specimen was compressed (Fig. 15B). The atrium is surrounded by a thin membrane. Aquiferous system solenoid. The sponge surface is hispid by the presence of diactines crossing it (Fig. 15A). There are also few trichoxeas, many triactines and rare tetractines (Fig. 15C).

## Spicules (Table 7)

*Diactines:* Fusiform but with one end thicker than the other. They are slightly curved and undulated (Fig. 15D).

*Triactines:* Equiangular and equiradiate, sometimes subregular. Actines are slightly conical with sharp to blunt tips (Fig. 15E).

Tetractines: Equiangular and equiradiate. Actines are slightly conical with sharp to blunt tips (Fig. 15F). The apical actine is needle-like and covered by delicate short spines (Fig. 15G).

*Ecology:* Specimens were associated with colonial ascidians.

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: Currently there are 11 accepted species of Leucascus in the world: L. albus Cavalcanti et al., 2013; L. clavatus Dendy, 1892; L. digitiformis Klautau, Lopes et al., 2020; L. flavus; L. leptoraphis

(Jenkin, 1908); *L. lobatus* Rapp, 2004; *L. neocaledonicus* Borojević & Klautau, 2000; *L. protogenes* (Haeckel, 1872 sensu Dendy, 1891); *L. roseus* Lanna et al., 2007; *L. schleyeri* Van Soest & De Voogd, 2018; and *L. simplex* Dendy, 1892 (type species of the genus). From these, only *L. clavatus*, a species from Australia, and *L. schleyeri*, from South Africa, have diactines. The new species can be easily differentiated from them mainly by the fusiform shape of the diactines, which are club-shaped in *L. clavatus* and banana-shaped in *L. schleyeri*.

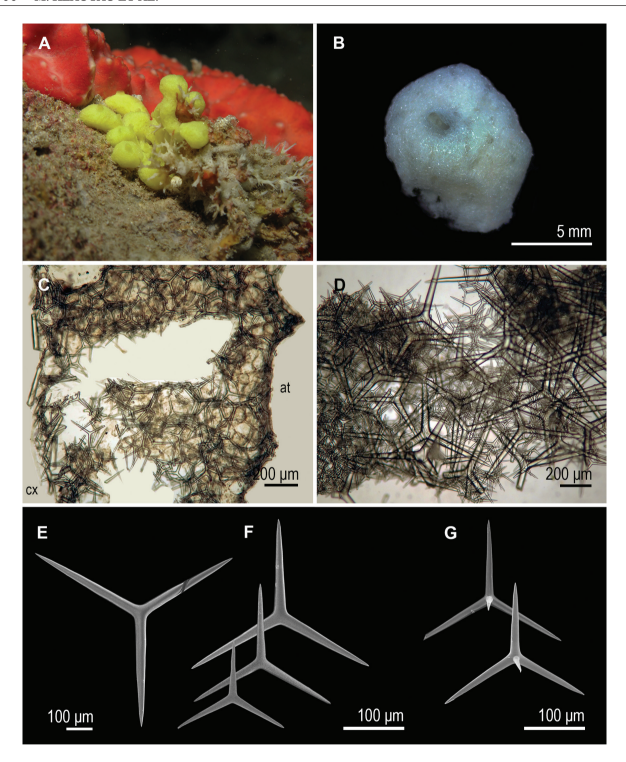
# Family Leucettidae Dendy & Row, 1913 Genus *Leucetta* Haeckel, 1872

Type species: Leucetta primigenia Haeckel, 1872.

Diagnosis: 'Leucettidae with a homogeneous organisation of the wall and a typical leuconoid aquiferous system. There is neither a clear distinction between the cortex and the choanoskeleton, nor the presence of a distinct layer of subcortical inhalant cavities. The atrium is frequently reduced to a system of exhalant canals that open directly into the osculum' (Borojević et al., 2002).

# LEUCETTA CHAGOSENSIS DENDY, 1913 (FIG. 16; TABLE 8)

Synonyms: Leucetta chagosensis Dendy, 1913: 10; Dendy & Row, 1913: 733; Dendy & Frederick, 1924: 482; Burton, 1963: 241; Borojević, 1967: 2; Pulitzer-Finali, 1982: 89; Gosliner et al., 1996: 16; Lévi, 1998: 77; Wörheide & Hooper, 1999: 882; Borojević & Klautau, 2000: 194; Wörheide et al., 2002: 1753, 2005: 379; Baine & Harasti, 2007: 15; Wörheide et al., 2008: 1; Voigt et al., 2012: 101; Van Soest & De Voogd, 2015: 51, 2018: 76; George et al., 2020: 285; Klautau et al., 2020: 279; Pasnin et al., 2020: 6. Leucetta sp. Colin & Arneson, 1995: 60 (photo 230). Ascoleucetta sagittata Cavalcanti et al., 2013: 308; Van Soest & De Voogd, 2015: 49; Lopes et al., 2018: 59; Klautau et al., 2020: 265.



**Figure 16.** Leucetta chagosensis. (UFRJPOR 8939). A, specimen in vivo. B, fixed specimen. C, cross-section. D, tangential section of the cortex. E, triactine I. F, triactines II. G, tetractines. cx = cortex, at = atrium.

Table 8. Spicule measurements of Leucetta chagosensis (UFRJPOR 8939)

| Specimen     | Spicule                     | Actine          | Length         | (µm)            |              |                | Width        | (µm)         |            |              | $\overline{N}$ |
|--------------|-----------------------------|-----------------|----------------|-----------------|--------------|----------------|--------------|--------------|------------|--------------|----------------|
|              |                             |                 | Min            | Mean            | SD           | Max            | Min          | Mean         | SD         | Max          |                |
| UFRJPOR 8939 | Triactine I<br>Triactine II |                 | 315.0<br>130.0 | 410.0<br>164.8  | 57.9<br>14.6 | 500.0<br>195.0 | 25.0<br>10.0 | 38.3<br>14.3 | 7.9<br>2.5 | 55.0<br>20.0 | 20<br>20       |
|              | Tetractine                  | Basal<br>Apical | 77.5<br>25.0   | $124.1 \\ 42.1$ | 22.5<br>9.0  | 155.0<br>55.0  | 10.0<br>5.0  | 13.1<br>6.2  | 2.9<br>1.3 | 17.5<br>7.5  | 20<br>13       |

Type locality: Salomon, Chagos Archipelago, Indian Ocean.

Material examined: UFRJPOR 8939 (= 171109-PAE2-TP15). Photos: TP6059-6060. Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18-20 m depth.

*Colour:* Lemon-yellow alive and white after fixation (Fig. 16A, B).

Morphology and anatomy: Sponge formed by several globose parts, each one with a single apical osculum surrounded by membrane (Fig. 16A, B). Surface smooth and transparent. Canals can be seen below the surface. Although the atrium is reduced, it obviously receives a large number of exhalant canals. The aquiferous system is leuconoid (Fig. 16C). The skeleton is composed of two size categories of cortical triactines (Fig. 16D). The small triactines form the main part of the choanosomal skeleton, which has also rare small tetractines surrounding the canals. The atrial skeleton is composed of the small triactines and the rare tetractines.

## Spicules (Table 8)

*Triactines I:* Large, equiangular and equiradiate. Actines are conical and sharp (Fig. 16E).

*Triactines II:* Small, equiangular and equiradiate. Actines are conical and blunt to sharp tips (Fig. 16F).

Tetractines: Small, equiangular and equiradiate. Actines are conical and blunt to sharp (Fig. 16G). The apical actine is short, conical and sharp.

Ecology: Leucetta chagosensis was found down to 25 m (Van Soest & De Voogd, 2018).

Geographic distribution: INDIAN OCEAN: Chagos Ecoregion – Salomon, Chagos Archipelago (Dendy, 1913); Houtman Ecoregion – Houtman Abrolhos (Dendy & Frederick, 1924); Maldives Ecoregion - Maldives (Wörheide et al., 2008; Van Soest & De Voogd, 2018; Pasnin et al., 2020); Arabian (Persian) Gulf Ecoregion – Red Sea (Wörheide et al., 2008; Voigt et al., 2012; Van Soest & De Voogd, 2018; Pasnin et al., 2020); India (Van Soest & De Voogd, 2018; George et al., 2020). Seychelles Ecoregion – Seychelles (Van Soest & De Voogd, 2018); Western and Northern Madagascar Ecoregion – Madagascar (Van Soest & De Voogd, 2018); Mascarene Islands Ecoregion - Rodrigues (Van Soest & De Voogd, 2018; Pasnin et al., 2020); La Réunion (Trentin & Massé, 2018; Pasnin et al., 2020). PACIFIC OCEAN: Banda Sea Ecoregion – Indonesia; Sulawesi Sea Ecoregion - Sulawesi; Lesser Sunda Ecoregion - Bali (Wörheide et al., 2002, 2008; Pasnin et al., 2020); Eastern Philippines Ecoregion – Philippines (Wörheide et al., 2008; Voigt et al., 2012; Van Soest & De Voogd, 2015, 2018; Pasnin et al., 2020); Great Barrier Reef, Australia; Tweed Moreton Ecoregion - Queensland, Australia (Wörheide & Hooper, 1999; Wörheide et al., 2002, 2004, 2005, 2008; Voigt et al., 2012; Pasnin et al., 2020); Mariana Islands Ecoregion - Guam, South Kuroshio Ecoregion - Taiwan; Central Kuroshio Current Ecoregion - Japan (Wörheide et al., 2002, 2008; Pasnin et al., 2020); Bismarck Sea Ecoregion and Solomon Sea Ecoregion - Papua New Guinea, West Caroline Islands Ecoregion - Palau Islands (Wörheide et al., 2002, 2005, 2008; Voigt et al., 2012; Van Soest & De Voogd, 2015; Pasnin et al., 2020); Vanuatu Ecoregion - Vanuatu; Fiji Islands Ecoregion - Fiji; Samoa Islands Ecoregion - Samoa (Wörheide et al., 2002, 2004, 2008; Pasnin et al., 2020); Society Islands Ecoregion – Society Island, French Polynesia; Tuamotus Ecoregion - Tuamotu Islands, French Polynesia (Wörheide et al., 2002, 2005, 2008; Klautau et al., 2020; Pasnin et al., 2020); New Caledonia Ecoregion - New Caledonia (Borojević, 1967; Lévi, 1998; Borojević & Klautau, 2000); Coral Sea Ecoregion - Coral Sea (Wörheide et al., 2008).

Taxonomic remarks: Leucetta chagosensis is a widespread species in the Indo-Pacific Ocean, which likely represents a species complex (Wörheide *et al.*, 2002, 2005, 2008; Voigt *et al.*, 2012; Pasnin *et al.*, 2020).

Descriptions of this species show variable morphologies, including variation in the external shape (globular to pyriform to massive lobate), colour after fixation (white or brown) and, perhaps, the spicules' size (Table 8; e.g. Dendy, 1913; Wörheide & Hooper, 1999; Borojević & Klautau, 2000; Van Soest & De Voogd, 2015, 2018; Klautau *et al.*, 2020). However, some characters seem to be constant, such as the bright yellow colour when alive, the large osculum and the conspicuous subsuperficial inhalant canals, running almost parallel.

Comparing measurements retrieved from the literature (Dendy, 1913; Wörheide & Hooper, 1999; Borojević & Klautau, 2000; Van Soest & De Voogd, 2015, 2018; Klautau et al., 2020) and this work, including new measurements of the holotype given by Alencar (2012), we observed that only the size of the spicules given by Van Soest & De Voogd (2018) was different, being much larger than all the others. These authors did not mention if their measurements were based on one specimen or several but, curiously, in their description of specimens from the Red Sea and the Indian Ocean, we also found other morphological differences. They mentioned the presence of numerous tetractines in the atrium, whereas in other descriptions of *L. chagosensis* these spicules were referred as rare. Moreover, they mentioned the presence of a reduced atrium, while other authors mentioned a wide atrial cavity in this species. In that work, specimens from Indonesia, the Red Sea, the Maldives and Rodrigues grouped in the same clade, together with L. villosa from Australia and Ascoleucetta sagittata Cavalcanti et al., 2013 from Indonesia (Van Soest & De Voogd, 2018). In this work, we also obtained a well-supported clade unifying specimens from Australia, French Polynesia, Indonesia, the Red Sea, Rodrigues and La Réunion, including A. sagittata, synonymized to L. chagosensis by Van Soest & De Voogd (2018) and L. villosa (Figs 2, 3).

Although Van Soest & De Voogd (2018) suggested that A. sagittata is a synonym of L. chagosensis, they considered L. villosa a valid species because of morphological evidence that might separate them. These characters are: colour alive (bright yellow in L. chagosensis and yellow-brownish in L. villosa), absence of sagittal spicules surrounding the osculum of L. villosa and its 'characteristic hairy surface' (Wörheide & Hooper, 1999). As colour in sponges can be variable, we should be careful about this character and, about the absence of sagittal spicules, we see some variation among L. chagosensis specimens. Regarding the 'hairy surface' that gave the name to the species, we analysed the holotype of L. villosa and found this to be colonial hydrozoans (Alencar, 2012). The same slight differences can be

seen between *L. chagosensis* and *A. sagittata*, as they can be differentiated only by the presence of sagittal spicules in the cortex and in the atrium of the latter and by the presence of large triactines only in the cortex.

Moreover, in a phylogeographic analysis performed with L. 'chagosensis' across the Indo-Pacific, L. villosa was considered to be possibly a hybrid with L. chagosensis (Wörheide et al., 2008). Based on the mentioned morphological characters, L. 'chagosensis' seems to be variable and based on genetic studies, there is not enough differentiation to distinguish L. villosa from L. chagosensis. Therefore, we agree with Van Soest & De Voogd (2018) that L. sagittata is synonymous of *L. chagosensis* and we propose the same for *L. villosa*. Of course, it is possible that *L. chagosensis* is a species complex. A recent work using more variable markers (Pasnin et al., 2020) found five lineages restricted to different biogeographic regions. The authors said they did not find morphological differences, but no morphological analyses were presented and they did not propose to split L. chagosensis. Hence, for the moment we consider A. sagittata and L. villosa synonyms of *L. chagosensis*.

ORDER MURRAYONIDA VACELET, 1981
FAMILY LELAPIELLIDAE BOROJEVIĆ *ET AL.*, 1990
GENUS *LELAPIELLA* VACELET, 1977

Type species: Lelapiella incrustans Vacelet, 1977 (by monotypy).

*Diagnosis:* Lelapiellidae encrusting and without fused spicules. Radial growing. Cortical skeleton composed chiefly of a layer of large tripods, diactines may be present. Choanosome with triactines and tetractines. Tracts of diactines may be present (modified from Borojević *et al.*, 2002).

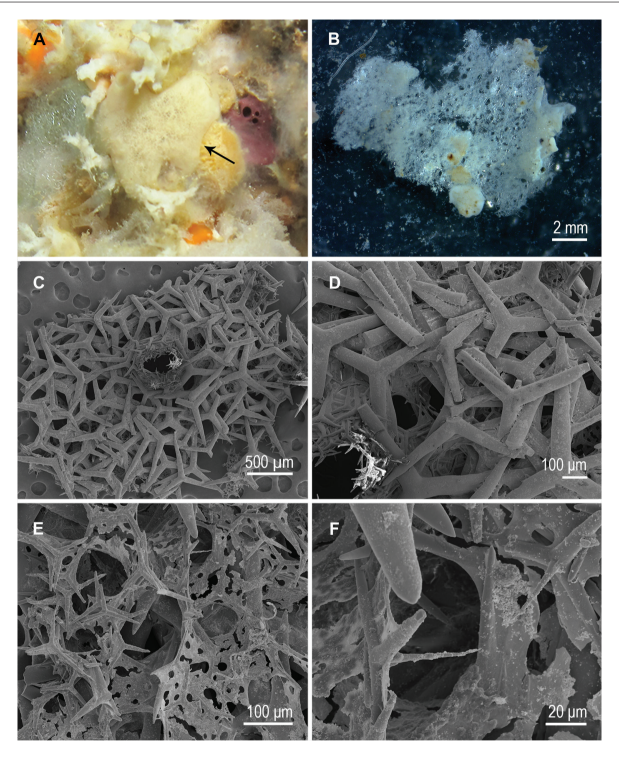
## LELAPIELLA TERTIA SP. NOV.

(FIGS 17, 18; TABLE 9)

Zoobank registration: urn:lsid:zoobank.org:act: AF5A1C7E-AA51-4D4B-A330-CAD0D653719E.

*Diagnosis: Lelapiella* with cortical tripods and choanosomal triactines and tetractines. Diactines and spicules tracts absent.

*Etymology:* From the Latin word *tertius*, which means third, because this is the third recognized species of the genus.



**Figure 17.** Lelapiella tertia (holotype; UFRJPOR 8937). A, specimen in vivo (arrow points to the sponge). B, fixed specimen. SEM: C, fragment of the specimen. D, tangential section of the surface. E, choanosome. F, tetractines in the choanosome.

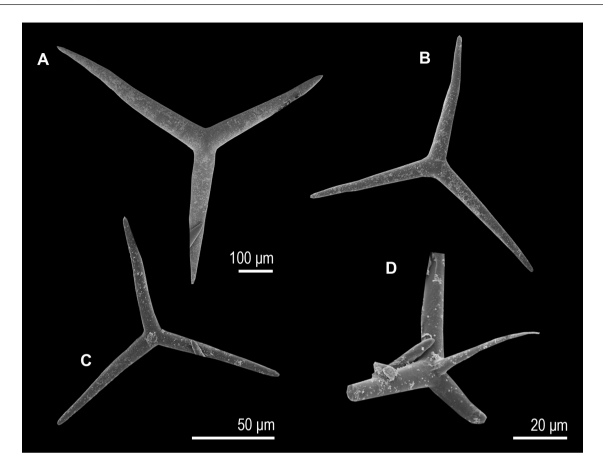


Figure 18. Lelapiella tertia (holotype; UFRJPOR 8937). A, tripod. B, triactine. C, tetractine. D, apical actine of a tetractine.

Type locality: Passe de l'Hermitage, La Réunion, Indian Ocean.

Type material: Holotype – UFRJPOR 8937 (= 171109-PAE2-TP13). Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

*Colour:* Sponge white to beige in life and white in ethanol (although we have not observed this species during the collection, it corresponds probably to the sponge in Fig. 17A).

Morphology and anatomy: Sponge encrusting, with radial growing and oscula on the top of short cones (Fig. 17A–C). Aquiferous system leuconoid. Surface smooth. The skeleton is composed of large cortical tripods (Fig. 17D) and choanosomal triactines (Fig. 17E) and few tetractines (Fig. 17F). Sagittal triactines and tetractines similar to those of the choanosome support the oscula. Trichoxeas are also present.

Spicules (Table 9)

*Tripods:* Equiangular and equiradiate to sagittal. Huge. Actines are conical, stout with rounded to blunt tips (Fig. 18A).

*Triactines:* Equiangular and equiradiate to sagittal. Actines are conical and blunt (Fig. 18B).

Tetractines: Equiangular and equiradiate to sagittal. Actines are conical and blunt (Fig. 18C). The apical actine is conical, slightly curved and undulated, sharp and smooth (Fig. 18D).

*Ecology:* This sponge was growing close to *Ascandra mascarenica* sp. nov. (UFRJPOR 8936), also protected from sunlight and associated with a bryozoan and a hydrozoan.

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: Lelapiella inscrustans, a species from Madagascar (Western Indian Ocean), is the

**Table 9.** Spicule measurements of *Lelapiella tertia*, *L. incrustans* and *L. sphaerulifera*. Abbreviations: c = curved; H = holotype; o = from the osculum; r = reniform; st = straight, from the tracts

| Specimen               | Spicule       | Actine   | Length | (µm)  |       |       | Width | (µm) |      |      | N  |
|------------------------|---------------|----------|--------|-------|-------|-------|-------|------|------|------|----|
|                        |               |          | Min    | Mean  | SD    | Max   | Min   | Mean | SD   | Max  |    |
| L. tertia sp. nov. (H) | Tripod        |          | 118.9  | 490.8 | 150.3 | 713.5 | 32.4  | 75.7 | 16.0 | 97.3 | 20 |
| _                      | Triactine     |          | 67.5   | 82.1  | 7.5   | 94.5  | 6.8   | 9.9  | 1.6  | 13.5 | 20 |
|                        | Tetractine    | Basal    | 75.6   | 82.9  | 6.1   | 99.9  | 8.1   | 10.0 | 1.1  | 10.8 | 20 |
|                        |               | Apical   | 27.0   | 41.3  | 8.6   | 54.0  | 4.1   | 5.4  | 0.7  | 8.1  | 20 |
| $L.\ incrustans\ (H)$  | Diactine (st) |          | 80.0   | -     | -     | 200.0 | 6.0   | -    | -    | 20.0 | -  |
|                        | Diactine (c)  |          | 50.0   | -     | -     | 160.0 | 10.0  | -    | -    | 15.0 | -  |
|                        | Diactine (r)  |          | -      | 35.0  | -     | -     | -     | 12.0 | -    | -    | -  |
|                        | Tripods       |          | 65.0   | -     | -     | 400.0 | 32.0  | -    | -    | 60.0 | -  |
|                        | Triactine (o) | Paired   | 40.0   | -     | -     | 60.0  | 7.5   | -    | -    | 20.0 | -  |
|                        |               | Unpaired | 37.0   | -     | -     | 55.0  | 10.0  | -    | -    | 15.0 | -  |
|                        | Tetractine    | Basal    | 75.0   | -     | -     | 80.0  | 10.0  | -    | -    | 12.0 | -  |
|                        |               | Apical   | 50.0   | -     | -     | 80.0  | 10.0  | -    | -    | 12.0 | -  |
| L. sphaerulifera       | Diactine (st) |          | 126.0  | 193.2 | -     | 297.0 | 13.0  | 17.2 | -    | 27.0 | -  |
|                        | Diactine (c)  |          | 84.0   | 95.8  | -     | 106.0 | 18.0  | 19.1 | -    | 21.0 | -  |
|                        | Diactine (r)  |          | 26.0   | 30.2  | -     | 36.0  | 17.0  | 18.2 | -    | 20.0 | -  |
|                        | Tripods       |          | 171.0  | 226.2 | -     | 285.0 | 36.0  | 46.0 | -    | 57.0 | -  |
|                        | Tetractine    | Basal    | 78.0   | -     | -     | 100.0 | -     | 9.0  | -    | -    | -  |
|                        |               | Apical   | 10.0   | -     | -     | 20.0  | -     | -    | -    | -    | -  |

type species of Lelapiella (Vacelet, 1977). In the same paper, a subspecies, Lelapiella inscrustans sphaerulifera, from Moorea (French Polynesia) was also described and later elevated to the species rank (Van Soest & De Voogd, 2015). Both species are morphologically similar, being encrusting, circular and having large cortical tripods, choanosomal tetractines, triactines, straight diactines forming tracts, curved diactines and reniform diactines. According to Vacelet (1977), the creation of a subspecies was justified by differences in the size and shape of the spicules. For example, L. inscrustans harbours tripods with 'nipples' and few reniform diactines, whereas L. inscrustans sphaerulifera presents abundant smooth tripods and reniform diactines (Vacelet, 1977). Studying specimens from Indonesia, Van Soest & De Voogd (2015) identified them as L. sphaerulifera, elevating the subspecies to species status, although not explaining why. We are not certain that the specimens from Indonesia are L. sphaerulifera. Vacelet (1977) mentioned that his subspecies presented 'nipples' in some tripods. although some of them were smooth. Besides, he found triactines in his specimens from French Polynesia. Van Soest & De Voogd (2015) did not

mention the presence of 'nipples' in the tripods from the Indonesian specimens and showed images of a smooth tripod. Moreover, they did not mention the presence of triactines in their specimens.

Because we consider that *L. sphaerulifera* is a distinct species, the sponge we are describing from La Réunion has to be the third of the genus. Indeed, *Lelapiella tertia* can be differentiated from *L. inscrustans* and *L. sphaerulifera* by skeleton composition and also by spicules size (Table 9). The new species does not have diactines, except for some trichoxeas. It is interesting that, similarly to *L. inscrustans* and *L. sphaerulifera*, the new species is encrusting, grows radially, its oscula are on the top of short cones and its skeleton has tripods. However, the new species does not have the characteristic tract of diactines of *L. inscrustans* and *L. sphaerulifera*, which convinced us to propose a new diagnosis for the genus.

Because our analyses corroborated that *Lelapiella* is a Calcinea (Borojević *et al.* 1990; Dohrmann *et al.*, 2006; Voigt *et al.*, 2012) and that it is not a Murrayonida (Dohrmann *et al.*, 2006; Voigt *et al.*, 2012; Fig. 5), we propose to synonymize Murrayonida with Clathrinida.

SUBCLASS CALCARONEA BIDDER, 1898
ORDER LEUCOSOLENIIDA HARTMAN, 1958
FAMILY GRANTIIDAE DENDY, 1892
GENUS APHROCERAS GRAY, 1858
Type species: Aphroceras alcicornis Gray, 1858.

*Diagnosis:* 'Grantiidae with leuconoid organisation. The cortex is supported, at least in part, by giant longitudinal diactines' (Borojević *et al.*, 2002).

APHROCERAS SEYCHELLENSIS (HÔZAWA, 1940) COMB. NOV. (FIGS 19–21; TABLE 10) Basionym: Leucandra seychellensis Hôzawa, 1940: 158; Burton, 1963: 519.

Type locality: Seychelles, Indian Ocean.

Material examined: UFRJPOR 8926 (= 171109-PAE2-TP2). UFRJPOR 8934 (= 171109-PAE2-TP9). UFRJPOR 8935 (= 171109-PAE2-TP10). Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll. T. Pérez, 9 November 2017, 18–20 m depth.

*Colour:* White alive but with a grey region around the oscula (Fig. 19A, B). White after fixation (Fig. 19C, D).

Morphology and anatomy: Sponge amorphous, massive, with several large and small apical oscula (Fig. 19A–D) surrounded by membrane (Fig. 19D) and

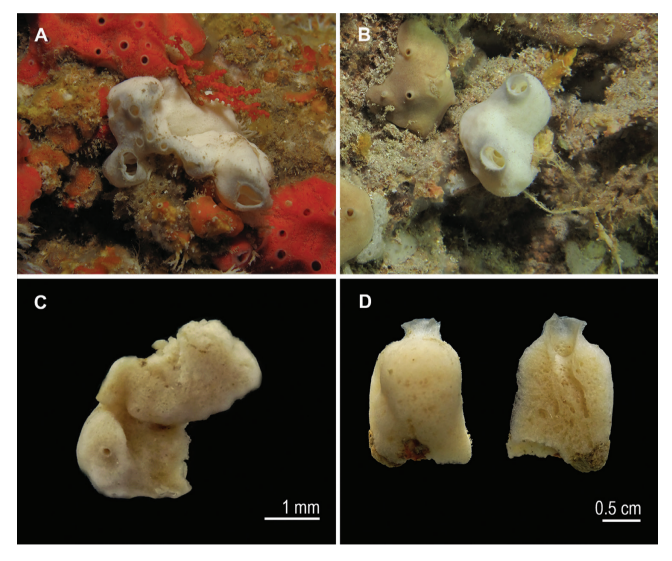
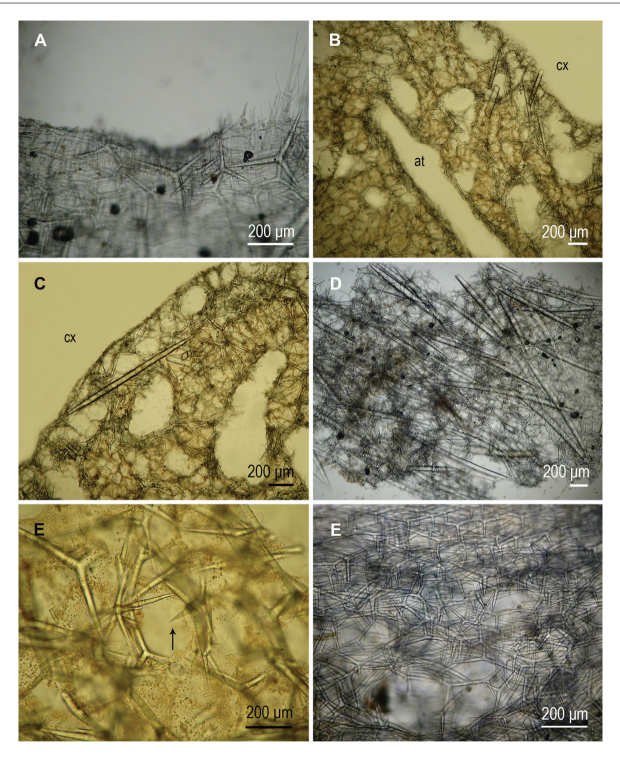


Figure 19. Aphroceras seychellensis. A, specimen in vivo (UFRJPOR 8926). B, specimen in vivo (UFRJPOR 8934). C, fixed specimen (UFRJPOR 8926). D, fixed specimen (UFRJPOR 8934).



**Figure 20.** Aphroceras seychellensis (UFRJPOR 8926). A, oscular margin. B, cross-section. C, cross-section evidencing subcortical lacunae. D, tangential section of the cortex. E, choanosomal canal surrounded by a triactine and a tetractine (arrow points to the apical actine of the tetractine). F, tangential section of the atrial surface. cx = cortex, at = atrium.

a short crown of trichoxeas (Fig. 20A). The crown is supported by T-shape triactines with the unpaired actine thinner than the paired ones (Figs 20A, 21B). The huge cortical diactines protrude near the oscula

but lie tangentially under the surface in most of the body. As a result, the surface is smooth. The atrium can be reduced or large (Fig. 19D). It is possible to see the lacunae and canals below the cortex (Fig. 20B). The

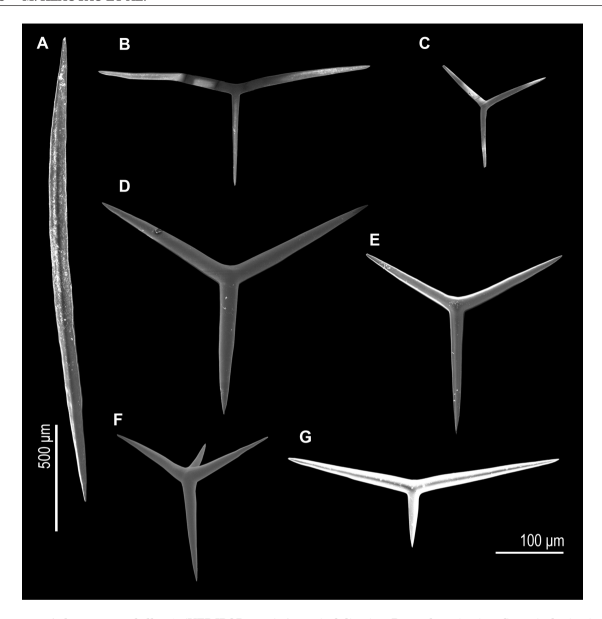


Figure 21. Aphroceras seychellensis (UFRJPOR 8926). A, cortical diactine. B, oscular triactine. C, cortical triactine. D, choanosomal triactine. E, choanosomal triactine. F, tetractine of the canals. G, atrial triactine.

aquiferous system is leuconoid. The cortical skeleton is composed of longitudinal giant diactines and small triactines (Fig. 20B–D). The choanosome has dispersed triactines, and triactines and tetractines surrounding the canals (Fig. 20C, E). The triactines tend to reduce the angle between the paired actines when they are not surrounding the canals (i.e. less sagittal). The atrial skeleton has only triactines and they are larger than the cortical ones (Fig. 20F).

# Spicules (Table 10)

*Diactines:* Giant and fusiform, with both tips sharp (Fig. 21A).

Cortical triactines: Small and sagittal. Actines are cylindrical and sharp (Fig. 21C).

Choanosomal triactines: Variable size, subregular to sagittal. Actines are conical or slightly conical and sharp. (Fig. 21D, E).

Choanosomal tetractines (canals): Sagittal, with curved paired actines. Actines are slightly conical to conical and sharp. The apical actine is conical, sharp and short (Fig. 21F).

Atrial triactines: Strongly sagittal. The paired actines are straight and much longer than the unpaired one. They are conical and sharp (Fig. 21G).

**Table 10.** Spicule measurements of *Aphroceras seychellensis* from La Réunion and from Seychelles (Hôzawa, 1940). H = holotype

| Specimen         | Spicule               | Actine   | Lengtl | n (µm) |       |        | Width | (µm) |      |       | N  |
|------------------|-----------------------|----------|--------|--------|-------|--------|-------|------|------|-------|----|
|                  |                       |          | Min    | Mean   | SD    | Max    | Min   | Mean | SD   | Max   |    |
| UFRJPOR 8926     | Cortical diactine     |          | 729.7  | 1695.0 | 406.1 | 2243.2 | 45.9  | 67.1 | 15.1 | 94.5  | 14 |
|                  | Cortical triactine    | Paired   | 59.4   | 93.4   | 19.6  | 121.5  | 5.4   | 7.3  | 1.5  | 10.8  | 20 |
|                  |                       | Unpaired | 51.3   | 77.9   | 20.2  | 113.4  | 5.4   | 7.2  | 1.6  | 10.8  | 20 |
|                  | Choanosomal triactine | Paired   | 94.5   | 169.4  | 45.6  | 243.0  | 13.5  | 21.5 | 4.9  | 32.4  | 20 |
|                  |                       | Unpaired | 81.0   | 142.7  | 31.9  | 221.4  | 13.5  | 21.5 | 4.1  | 29.7  | 20 |
|                  | Canal tetractine      | Paired   | 124.2  | 153.3  | 18.4  | 183.6  | 16.2  | 18.9 | 2.7  | 24.3  | 09 |
|                  |                       | Unpaired | 105.3  | 123.9  | 19.3  | 167.4  | 16.2  | 19.2 | 1.6  | 21.6  | 09 |
|                  |                       | Apical   | 51.3   | 58.8   | 5.7   | 70.2   | 10.8  | 13.5 | 1.1  | 16.2  | 13 |
|                  | Atrial triactine      | Paired   | 129.6  | 175.1  | 26.0  | 243.0  | 13.5  | 17.7 | 2.6  | 24.3  | 20 |
|                  |                       | Unpaired | 45.9   | 75.3   | 17.2  | 113.4  | 13.5  | 15.5 | 1.9  | 18.9  | 20 |
| UFRJPOR 8934     | Cortical diactine     | -        | 918.9  | 1621.6 | 422.7 | 2432.4 | 54.0  | 86.4 | 16.2 | 113.4 | 20 |
|                  | Cortical triactine    | Paired   | 59.4   | 83.4   | 16.5  | 113.4  | 5.4   | 7.1  | 1.4  | 9.5   | 20 |
|                  |                       | Unpaired | 40.5   | 75.9   | 17.8  | 105.3  | 5.4   | 6.9  | 1.4  | 8.1   | 20 |
|                  | Choanosomal triactine | Paired   | 86.4   | 172.9  | 34.9  | 240.3  | 16.2  | 26.1 | 5.3  | 40.5  | 20 |
|                  |                       | Unpaired | 91.8   | 141.9  | 28.3  | 216.0  | 16.2  | 25.4 | 3.9  | 32.4  | 20 |
|                  | Canal tetractine      | Paired   | 124.2  | 153.9  | 21.1  | 183.6  | 16.2  | 19.4 | 3.5  | 24.3  | 05 |
|                  |                       | Unpaired | 89.1   | 112.3  | 23.1  | 151.2  | 16.2  | 19.4 | 2.3  | 21.6  | 05 |
|                  |                       | Apical   | 54.0   | 60.9   | 6.4   | 70.2   | 10.8  | 13.1 | 1.0  | 13.5  | 07 |
|                  | Atrial triactine      | Paired   | 143.1  | 199.4  | 28.0  | 245.7  | 16.2  | 22.5 | 4.0  | 29.7  | 20 |
|                  |                       | Unpaired | 40.5   | 66.2   | 12.1  | 83.7   | 16.2  | 21.0 | 3.0  | 27.0  | 20 |
| Hôzawa 1940 (H)* | Cortical diactine     | -        | 800.0  | _      | _     | 1050.0 | 45.0  | _    | _    | 65.0  | -  |
|                  | Cortical triactine    |          | -      | 70.0   | _     | -      | -     | 6.0  | _    | -     | -  |
|                  | Choanosomal triactine | Paired   | 140.0  | _      | _     | 240.0  | 20.0  | _    | _    | 30.0  | -  |
|                  |                       | Unpaired | 130.0  | _      | _     | 230.0  | 20.0  | _    | _    | 30.0  | -  |
|                  | Canal tetractine      | Apical   | -      | 50.0   | -     | -      | -     | 14.0 | -    | -     | -  |
|                  | Atrial triactine      | Paired   | 130.0  | _      | _     | 180.0  | 8.0   | _    | -    | 10.0  | -  |
|                  |                       | Unpaired | 50.0   | -      | -     | 90.0   | 6.0   | -    | -    | 8.0   | -  |

<sup>\*</sup>Hôzawa (1940) also gave, for the cortical triactines, the following measurements: 260.0 µm length/ 20 µm width. We consider that this measurement belongs to the large choanosomal triactines.

*Ecology:* The specimen was attached to a coral and an undetermined red sponge (Fig. 19A).

Geographic distribution: Seychelles Ecoregion – Seychelles (Hôzawa, 1940); Mascarene Islands Ecoregion – La Réunion (this study).

Taxonomic remarks: Our specimens from La Réunion mostly match the description given by Hôzawa (1940) for Leucandra seychellensis, a species from the Seychelles. The only differences were the size of the diactines, larger and thicker in our specimens, and the atrial triactines, thicker in the specimens from La Réunion (Table 10). As we also observed some differences in spicule measurements among our specimens, we did not consider these differences strong enough to not consider the specimens from La Réunion conspecifc with Hôzawa's specimen. The other difference we found concerns the

presence of large triactines in the cortex. Hôzawa (1940) mentioned the presence of small and large triactines in the cortex and in the choanosome. We observed that, although these large triactines are sometimes near the cortex in our specimens, they are mainly part of the choanosome. Therefore, we described the cortex as having only small triactines (and the diactines) and the choanosome as having the large triactines, small triactines (mainly surrounding canals) and tetractines only surrounding canals. Therefore, we identified our specimens as conspecific with those of Hôzawa.

The presence of diactines running parallel to the cortical surface, and not protruding through it, indicates that *L. seychellensis* is not *Leucandra* but *Aphroceras*. Therefore, we propose here to transfer this species to the genus *Aphroceras*. Moreover, Hôzawa (1940) pointed out that the Mediterranean *L. crambessa* Haeckel, 1872 is also similar to *A. seychellensis* because of large parallel

diactines in the cortical skeleton. Both species, however, differ mainly in the composition of the atrial skeleton: tetractines are present in  $L.\ crambessa$ , whereas  $A.\ seychellensis$  only has triactines. Considering the positioning of the diactines of  $L.\ cambressa$ , this species also has to be transferred to Aphroceras.

GENUS LEUCANDRA HAECKEL, 1872

Type species: Sycinula egedii Schmidt, 1870.

*Diagnosis:* 'Grantiidae with sylleibid or leuconoid organization. Longitudinal large diactines, if present, are not restricted to the cortex, but lie obliquely across the external part of the sponge wall and protrude from the surface of the sponge' (Borojević *et al.* 2002).

### LEUCANDRA ORNATA SP. NOV.

(FIGS 22-25; TABLE 11)

Zoobank registration: urn:lsid:zoobank.org:act:2F2F2C38-A216-4B1F-8A04-7CAA93B8F0EC.

Diagnosis: Leucandra with spined microdiactines in the cortical and in the atrial skeletons. The cortical microdiactines are frequently organized forming bouquets. Large cortical diactines are also present, along with cortical triactines. The choanosome contains regular triactines and sagittal triactines and tetractines. The atrial skeleton has microdiactines with spines and tetractines.

*Etymology:* From the Latin *ornatum*, ornate or decorated, for the presence of many diactines on the surface of this species.

Type locality: Cap La Houssaye Reef, La Réunion, Indian Ocean.

Type material: Holotype – UFRJPOR 8928 (= 171108-CAH2-TP2, Photos: TP5973–5975). Cap La Houssaye Reef, La Réunion, Indian Ocean, coll. T. Pérez, 8 November 2017, 10 m depth.

Colour: White alive (Fig. 22A) and after fixation (Fig. 22B).

Morphology and anatomy: Sponge vase-shaped with apical osculum surrounded by a crown of trichoxeas (Figs 22A, B, 24A) supported by sagittal triactines and tetractines (Fig. 23A). Surface hispid because of several large diactines protruding through the surface (Figs 22A, B, 23C, 24B). The aguiferous system is leuconoid. The cortical skeleton is composed of tangential triactines, large diactines and microdiactines protruding through the surface (Figs 22A, B, 23C, D, 24C). The cortical microdiactines form bouquets or lie tangentially to the cortex (Figs 23D, 24C). The choanosome contains large and small triactines. These small triactines are found spread in the choanosome or surrounding the canals, although tetractines are the main spicules in the canals (Fig. 23E). The atrial skeleton is composed of tetractines and microdiactines tangential to the atrium (Figs 23F, 24D).

Spicules (Table 11)

Cortical diactines: Huge and fusiform, with both tips sharp (Fig. 25A).

Cortical microdiactines: They are fusiform and slightly bent with sharp tips. The tip that lay outside

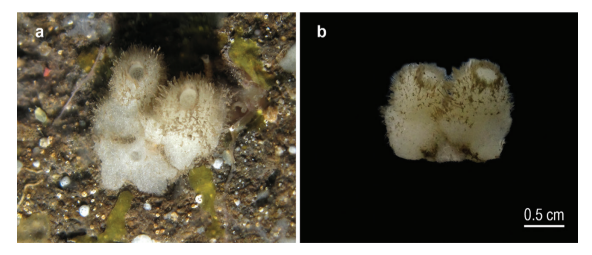
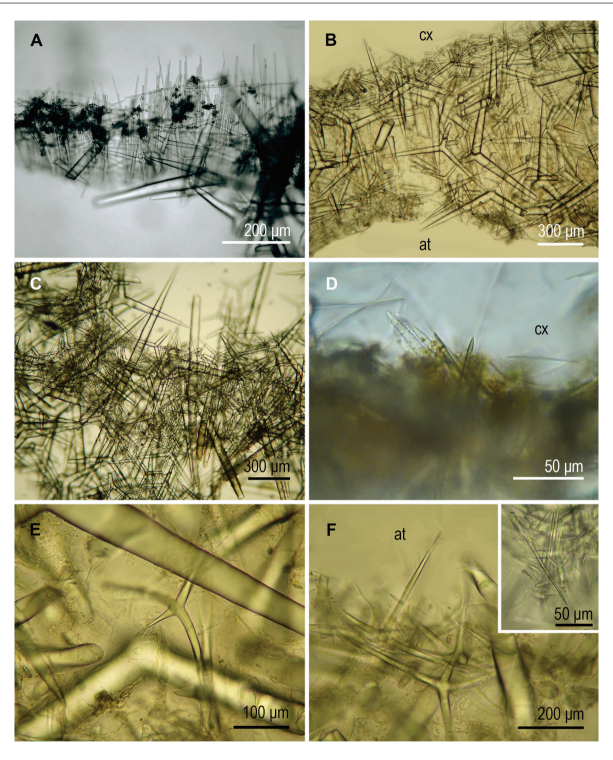
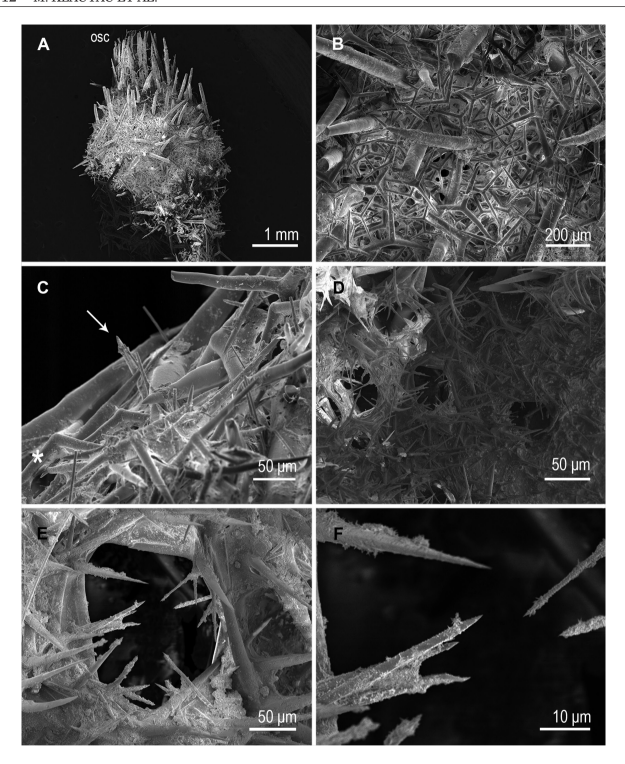


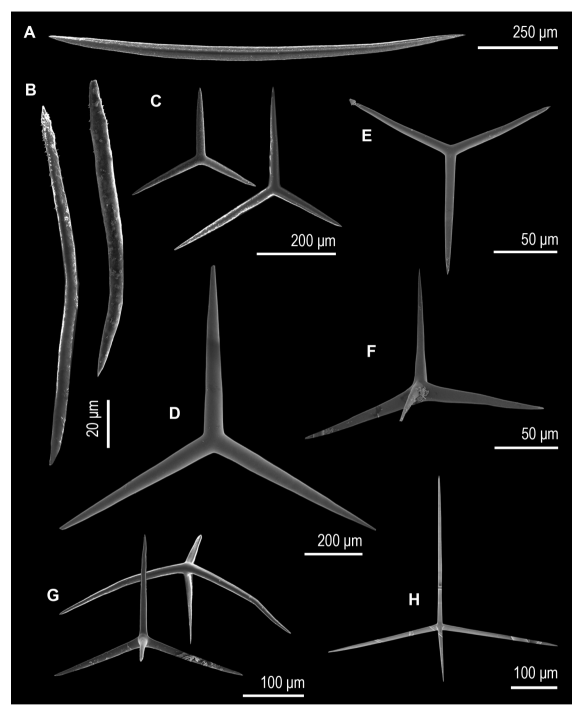
Figure 22. Leucandra ornata (holotype; UFRJPOR 8928). A, specimen in vivo. B, fixed specimen.



**Figure 23.** *Leucandra ornata* (holotype; UFRJPOR 8928). A, oscular margin. B, cross-section. C, tangential section of the cortex. D, detail of the cortex showing diactines II organised in bouquet. E, detail of the choanosome showing a tetractine of the canals. F, cross-section of the atrium (microdiactines in the inset). cx = cortex, at = atrium.



**Figure 24.** *Leucandra ornata* (holotype; UFRJPOR 8928). SEM: A, entire specimen. B, tangential section of the cortex. C, detail of the cortex showing a microdiactine (arrow) and a cortical triactine with elevated centre (asterisk). D, tangential section of the atrium. E, detail of an exhalant canal with microdiactines. F, detail of the microdiactines with spines. osc = osculum.



**Figure 25.** Leucandra ornata (holotype; UFRJPOR 8928). A, diactines. B, microdiactines. C, cortical triactines. D, choanosomal large triactine. E, canal triactine. F, tetractine of the canals. G, atrial tetractines. H, tetractine of the osculum.

**Table 11.** Spicule measurements of  $Leucandra\ ornata$  and the original measurements of  $L.\ riojai$ . H = holotype

| Specimen         | Spicule                | Actine   | Length (µm) | m)     |       |        | Width (µm) | mr)  |      |       |    |
|------------------|------------------------|----------|-------------|--------|-------|--------|------------|------|------|-------|----|
|                  |                        |          | Min         | Mean   | SD    | Max    | Min        | Mean | SD   | Max   | N  |
| UFRJPOR 8928 (H) | Cortical diactine      |          | 1162.3      | 1389.3 | 193.6 | 1621.8 | 54.0       | 62.1 | 5.4  | 67.5  | 05 |
|                  | Cortical microdiactine |          | 81.0        | 171.4  | 127.9 | 399.9  | 5.4        | 9.0  | 4.5  | 18.9  | 60 |
|                  | Cortical triactine     | Paired   | 118.9       | 264.0  | 99.7  | 486.5  | 16.2       | 25.4 | 8.9  | 43.2  | 20 |
|                  |                        | Unpaired | 97.3        | 233.2  | 99.1  | 464.8  | 13.5       | 26.2 | 10.1 | 54.0  | 20 |
|                  | Choanosomal triactine  |          | 540.5       | 671.2  | 58.6  | 783.8  | 37.8       | 61.9 | 8.7  | 75.6  | 30 |
|                  | Canal triactine        | Paired   | 162.2       | 194.6  | 28.6  | 216.2  | 16.2       | 18.0 | 1.6  | 18.9  | 03 |
|                  |                        | Unpaired | 205.4       | 234.2  | 25.0  | 248.6  | 16.2       | 18.9 | 2.7  | 21.6  | 03 |
|                  | Canal tetractine       | Paired   | 162.2       | 227.6  | 23.4  | 259.4  | 16.2       | 20.5 | 2.2  | 24.3  | 20 |
|                  |                        | Unpaired | 173.0       | 231.3  | 26.3  | 270.3  | 16.2       | 20.5 | 2.2  | 24.3  | 20 |
|                  |                        | Apical   | 54.0        | 76.8   | 10.7  | 91.8   | 10.8       | 17.1 | 2.9  | 21.6  | 20 |
|                  | Atrial tetractine      | Paired   | 129.7       | 230.3  | 41.4  | 324.3  | 13.5       | 16.6 | 2.8  | 21.6  | 20 |
|                  |                        | Unpaired | 97.3        | 202.7  | 41.5  | 270.3  | 13.5       | 16.3 | 2.2  | 21.6  | 20 |
|                  |                        | Apical   | 108.0       | 143.1  | 20.6  | 197.1  | 13.5       | 15.8 | 1.6  | 18.9  | 20 |
|                  | Atrial microdiactine   |          | 97.2        | 114.1  | 11.7  | 143.1  | 2.7        | 2.9  | 0.7  | 5.4   | 20 |
| L. riojai        | Cortical diactine      |          | 1600.0      | 1      |       | 2500.0 | 0.06       |      | •    | 100.0 | •  |
|                  | Cortical microdiactine |          | 100.0       | 1      |       | 240.0  | •          |      | •    | •     | •  |
|                  | Cortical triactine     |          | 160.0       | ı      |       | 210.0  | 12.0       |      | •    | 14.0  | •  |
|                  | Choanosomal triactine  |          | 180.0       | ı      |       | 240.0  | 20.0       |      | •    | 30.0  | ٠  |
|                  | Atrial tetractine      |          | ı           | 1      | ı     | 1      |            | ı    |      | ٠     | •  |
|                  | Atrial microdiactine   |          | 100.0       | 1      | 1     | 240.0  | ı          | ı    | 1    | 1     | ı  |

the sponge has delicate spines. These spicules are distributed in bouquets or lay tangentially to the cortex (Fig. 25B).

Cortical triactines: Equiangular and equiradiate. Actines are conical and sharp (Fig. 25C).

Choanosomal triactines: Huge. Equiangular and equiradiate. Actines are conical and sharp (Fig. 25D).

*Triactines of the canals:* Small, rare. Sagittal. Actines are conical and sharp (Fig. 25E).

Tetractines of the canals: Small. Sagittal. Actines are conical and sharp (Fig. 25F). The apical actine is conical, sharp and smooth.

*Atrial tetractines:* Small. Sagittal. Actines are conical and sharp. The apical actine is conical, sharp, smooth and curved. It is thicker than the basal actines (Fig. 25G).

Atrial microdiactines: Thin, one tip is arrow-shaped and the other is sharp (Figs 23F, 24E). Small spines are present (Fig. 24F).

Oscular tetractines: Sagittal. Actines are cylindrical and sharp. The unpaired actine is longer than the paired ones. The apical actine is cylindrical, sharp, smooth and strongly curved. Consequently, it is similar to a chiactine (Fig. 25H).

*Ecology:* This sponge was found in a submarine cave, among a sponge community mostly dominated by encrusting growth forms.

Geographic distribution: Mascarene Islands Ecoregion – La Réunion (this work).

Taxonomic remarks: Many species of Leucandra have microdiactines with spines in the atrial skeleton but, considering the whole skeleton composition of *L. ornata*, the most similar species is L. riojai Ferrer-Hernandez, 1918. This globulose species occurs in the Mediterranean Sea (Spain) and, although it has the same skeleton compostion, we consider that some of the differences are enough to separate it from our new species. For example, Ferrer-Hernandez (1918) did not mention the presence of tetractines in the exhalant canals. Besides, our choanosomal triactines seem to be more regular than those of *L. riojai*. The size of the spicules is also different, being much larger in the new species, except for the large diactines, which are bigger in L. riojai (Table 11). Besides, in our species the cortical microdiactines are frequently organized in bouquets, while in L. riojai this kind of organization was not mentioned.

GENUS UTE SCHMIDT, 1862

Type species: Ute glabra Schmidt, 1864.

Diagnosis: 'Grantiidae with syconoid organization. The cortex is supported by giant longitudinal diactines, and the choanoskeleton is articulated, composed of several rows of triactines with occasional tetractines. There are no radial fascicles of diactines' (Borojević et al., 2002).

UTE INSULAGEMMAE VAN SOEST & DE VOOGD, 2018
(FIGS 26–28; TABLE 12)

Synonym: Ute insulagemmae, Van Soest & De Voogd, 2018: 92.

 $\begin{array}{lll} \textit{Material} & \textit{examined:} & \textit{UFRJPOR} & 8925 \\ (=171109\text{-PAE2-TP1}, \textit{Photos:} \textit{TP6017-6018}, \textit{TP6051}). \\ \textit{UFRJPOR} & 8938 & (=171109\text{-PAE2-TP14}, \textit{Photo:} \\ \textit{TP6052}). & \textit{UFRJPOR} & 8943 & (=171109\text{-PAE2-TP18}, \\ \textit{Photo:} & \textit{TP6050} & \textit{or} & \textit{TP6057}). & \textit{UFRJPOR} & 8948 \\ (=171109\text{-PAE2-TP23}, \textit{Photo:} & \textit{TP6061}). & \textit{Passe de l'Hermitage Reef, La Réunion, Indian Ocean, coll.} \\ \textit{T.Pérez}, 9 & \textit{November 2017}, 18-20 & \textit{m depth.} & \textit{UFRJPOR} \\ \textit{8929} & (=171110\text{-ESB2-TP4}, & \textit{Photos:} & \textit{TP6129-6130}). \\ \textit{UFRJPOR} & 8942 & (=171110\text{-ESB2-TP5}, & \textit{Photo:} \\ \textit{TP6131}). & \textit{Portail Caves, La Réunion, Indian Ocean, coll.} & \textit{T. Pérez, 9 November 2017, 25 m depth.} \\ \end{array}$ 

Type locality: Bijoutier Island, Amirantes, Seychelles, Indian Ocean.

Colour: This species can have several colours (Fig. 26). Some were mostly pink but white at the base (UFRJPOR 8925; Fig. 26A) or white with a pink sponge budding (UFRJPOR 8929) or even lemonyellow (UFRJPOR 8938; Fig. 26B), white but pink near the osculum (UFRJPOR 8942) or completely white (UFRJPOR 8943, 8948; Fig. 26C). They all become beige in ethanol (Fig. 26D).

Morphology and anatomy: Sponge with a shape of vase with one single apical osculum surrounded by a crown of trichoxeas and with a delicate sphincter (Fig. 26A–D). The crown is supported by T-shape triactines (Fig. 27A). Frequently there are sponges budding from another and they keep connected to each other (Fig. 26D). Surface smooth, covered by huge longitudinal diactines (Fig. 27B–D). These diactines protrude through the surface near the osculum (Fig. 27A). The surface is perforated by ostia surrounded by tiny curved triactines, giving a honeycomb appearance to it (Fig. 27C, D). The aquiferous system is syconoid (Fig. 27B). The cortical

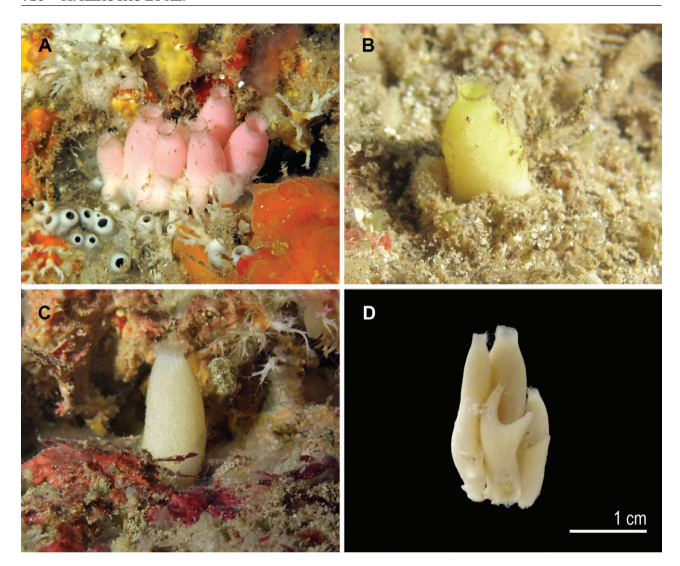


Figure 26. Ute insulagemmae. A, pink chromotype (UFRJPOR 8925). B, yellow chromotype (UFRJPOR 8938). C, white chromotype (UFRJPOR 8948). D, fixed specimen.

skeleton is composed of the giant longitudinal diactines and by the tiny triactines of the ostia (Fig. 27C). Below, the articulated tubar skeleton is composed of triactines pointing their unpaired actine to the surface (Fig. 27D, E). The subatrial skeleton has triactines and few tetractines and the atrial skeleton has only tetractines (Fig. 27E). The atrium is hispid (Fig. 27E, F).

Spicules (Table 12)

*Diactine:* Giant, fusiform with both tips sharp (Fig. 28A).

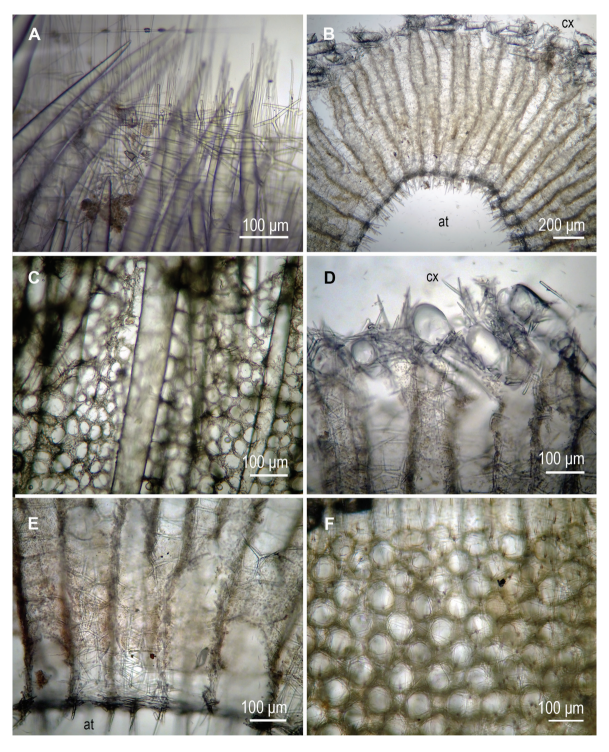
Cortical triactine: Tiny and sagittal. These spicules surround the ostia, consequently they have curved actines. Actines are conical and sharp and the unpaired actine is shorter than the paired ones (Fig. 28B).

Tubar triactines: Sagittal. The unpaired actine is longer than the paired ones and each paired actine has a different length. They are conical and sharp (Fig. 28C).

Subatrial triactine and tetractine: Sagittal. The unpaired actine is longer than the paired ones. They are conical and sharp. The apical actine of the tetractines is short, conical, sharp and curved, pointing into the atrium (Fig. 28D, E).

Atrial tetractine: Sagittal. The basal actines are shorter and thinner than the apical one. They are conical and sharp. The apical actine is long, straight, conical and sharp (Fig. 28F).

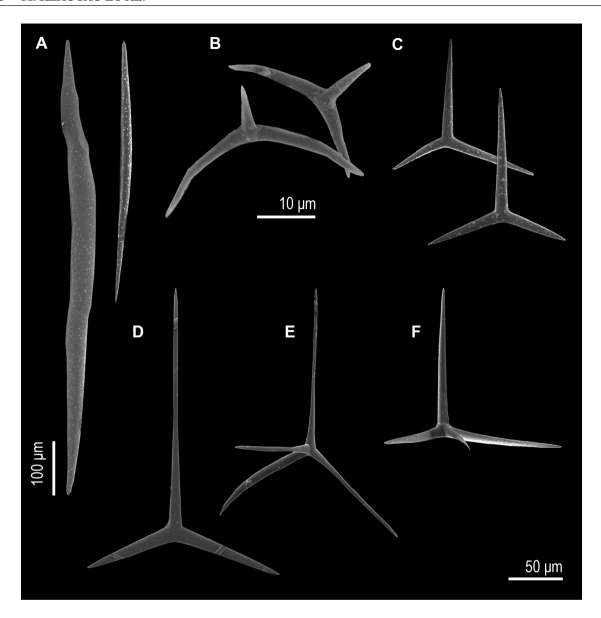
*Ecology:* Observed in caves, some specimens being associated with ophiuroids. It was found in the Seychelles from 6 to 18 m deep.



**Figure 27.** Ute insulagemmae (UFRJPOR 8928). A, oscular margin. B, cross-section. C, tangential section of the cortex. D, detail of the cortex. E, detail of the tubar, subatrial and atrial skeletons. F, tangential section of the atrial surface. cx = cortex.

Geographic distribution: Seychelles Ecoregion – Seychelles (Van Soest & De Voogd 2018); Mascarene Islands Ecoregion – La Réunion (this study).

Taxonomic remarks: Ute insulagemmae was described by Van Soest & De Voogd (2018) based on a 'pale orange'specimen collected in the Seychelles.



**Figure 28.** *Ute insulagemmae* (UFRJPOR 8928). A, diactines. B, cortical triactines. C, tubar triactines. D, subatrial triactine. E, subatrial tetractine. F, atrial tetractine.

Morphologically, except for the colour, our specimens match the description of the specimen from the Seychelles, including mostly the size of the spicules (Table 12). Considering all the analysed specimens, the diactines showed the largest size variation among the specimens, indicating that in this species spicule size can be variable. Colour is also variable, as shown by our molecular results.

# DISCUSSION

With this inventory, the number of known calcareous sponge species from the Mascarenes ecoregion increases

from 16 to 27, while that of La Réunion went from three to 14 (Table 13). Considering the two subclasses, Calcinea and Calcaronea, calcinean sponges continue to be the most diverse, with 17 species out of 27. Curiously, we did not find any of the species previously found in La Réunion (*T. stylifera*, *P. hindei* and *L. microraphis*) (Bourmaud, 2003) and of the 11 species found, seven are new to science, one of them belonging to a new genus (*Janusya*). The high number of species is remarkable, considering that our collections resulted from only seven short dives, most of them with only one sponge specialist, and in a limited area of La Réunion.

The knowledge about the occurrence of Calcarea in La Réunion seems to be fragmented. Among the

Table 12. Spicule measurements of *Ute insulagemmae* (UFRJPOR 8925). H = holotype

| Specimen                                  | Spicule              | Actine   | Length (µm) | mm)    |        |        | Width (µm) | (mm) |      |       |    |
|---|----------------------|----------|-------------|--------|--------|--------|------------|------|------|-------|----|
|   |                      |          | Min         | Mean   | SD     | Max    | Min        | Mean | SD   | Max   | N  |
| UFRJPOR 8925 (pink and white at the base) | Diactine             |          | 1540.5      | 2354.1 | 686.5  | 3783.8 | 54.1       | 83.8 | 14.9 | 108.1 | 20 |
|   | Cortical triactine   | Paired   | 17.5        | 24.6   | 4.5    | 35.0   | 2.5        | 2.5  | 0    | 2.5   | 20 |
|   |                      | Unpaired | 7.5         | 15.4   | 5.4    | 25.0   | 2.5        | 2.5  | 0    | 2.5   | 20 |
|   | Tubar triactine      | Paired   | 62.5        | 75.1   | 7.0    | 0.06   | 5.0        | 8.0  | 1.3  | 10.0  | 20 |
|   |                      | Unpaired | 85.0        | 97.3   | 7.9    | 115.0  | 7.5        | 8.4  | 1.2  | 10.0  | 20 |
|   | Subatrial triactine  | Paired   | 62.5        | 95.5   | 17.4   | 142.5  | 5.0        | 8.6  | 2.9  | 15.0  | 20 |
|   |                      | Unpaired | 115.0       | 163.4  | 37.6   | 230.0  | 5.0        | 7.3  | 2.4  | 12.5  | 20 |
|   | Subatrial tetractine | Paired   | 80.0        | 113.1  | 16.4   | 137.5  | 7.5        | 9.7  | 9.0  | 10.0  | 20 |
|   |                      | Unpaired | 125.0       | 182.0  | 38.0   | 240.0  | 5.0        | 5.3  | 8.0  | 7.5   | 20 |
|   |                      | Apical   | 30.0        | 39.9   | 9.7    | 67.5   | 5.0        | 5.5  | 1.0  | 7.5   | 20 |
|   | Atrial tetractine    | Paired   | 55.0        | 91.0   | 20.1   | 100.0  | 5.0        | 0.6  | 2.9  | 12.5  | 05 |
|   |                      | Unpaired | 37.5        | 63.0   | 23.3   | 100.0  | 5.0        | 0.6  | 2.2  | 10.0  | 05 |
|   |                      | Apical   | 87.5        | 106.9  | 12.1   | 130.0  | 5.0        | 7.8  | 1.4  | 10.0  | 20 |
| UFRJPOR 8938 (lemon yellow)               | Diactine             |          | 345.0       | 1428.3 | 1145.2 | 4500.0 | 15.0       | 66.5 | 41.6 | 125.0 | 20 |
|   | Cortical triactine   | Paired   | 15.0        | 30.0   | 7.9    | 45.0   | 2.5        | 3.0  | 1.03 | 5.0   | 20 |
|   |                      | Unpaired | 10.0        | 15.6   | 4.2    | 25.0   | 2.5        | 2.5  | 0.0  | 2.5   | 20 |
|   | Tubar tractine       | Paired   | 67.5        | 88.0   | 14.3   | 112.5  | 7.5        | 10.1 | 1.5  | 12.5  | 20 |
|   |                      | Unpaired | 55.0        | 84.9   | 14.0   | 102.5  | 7.5        | 6.6  | 1.7  | 12.5  | 20 |
|   | Subatrial triactine  | Paired   | 55.0        | 82.3   | 17.8   | 117.5  | 5.0        | 8.8  | 1.9  | 12.5  | 20 |
|   |                      | Unpaired | 112.5       | 153.9  | 38.6   | 250.0  | 5.0        | 8.4  | 2.0  | 12.5  | 20 |
|   | Subatrial tetractine | Paired   | 62.5        | 87.5   | 16.5   | 107.5  | 7.5        | 8.3  | 1.7  | 12.5  | 10 |
|   |                      | Unpaired | 92.5        | 160.0  | 34.7   | 212.5  | 7.5        | 8.3  | 1.2  | 10.0  | 10 |
|   |                      | Apical   | 25.0        | 34.5   | 7.5    | 50.0   | 7.5        | 8.0  | 1.1  | 10.0  | 10 |
|   | Atrial tetractine    | Paired   | 47.5        | 86.2   | 16.7   | 107.5  | 5.0        | 8.2  | 1.5  | 10.0  | 15 |
|   |                      | Unpaired | 20.0        | 83.5   | 14.7   | 100.0  | 7.5        | 8.3  | 1.2  | 10.0  | 15 |
|   |                      | Apical   | 17.5        | 29.0   | 29.0   | 100.0  | 5.0        | 7.5  | 1.5  | 10.0  | 12 |
|   |                      |          |             |        |        |        |            |      |      |       |    |

| Table 12. Continued  |                      |          |             |        |       |        |            |      |      |       |    |
|----------------------|----------------------|----------|-------------|--------|-------|--------|------------|------|------|-------|----|
| Specimen             | Spicule              | Actine   | Length (µm) | mm)    |       |        | Width (µm) | (mm) |      |       |    |
|                      |                      |          | Min         | Mean   | SD    | Max    | Min        | Mean | SD   | Max   | ×  |
| UFRJPOR 8943 (white) | Diactine             |          | 200.0       | 1360.0 | 727.3 | 2500.0 | 15.0       | 66.5 | 33.1 | 125.0 | 20 |
|                      | Cortical triactine   | Paired   | 20.0        | 32.1   | 8.2   | 47.5   | 2.5        | 3.3  | 1.2  | 5.0   | 20 |
|                      |                      | Unpaired | 7.5         | 19.9   | 8.9   | 35.0   | 2.5        | 2.6  | 9.0  | 5.0   | 20 |
|                      | Tubar triactine      | Paired   | 57.5        | 78.0   | 14.0  | 107.5  | 7.5        | 10.8 | 2.8  | 17.5  | 20 |
|                      |                      | Unpaired | 57.5        | 88.8   | 17.7  | 125.0  | 7.5        | 10.6 | 2.0  | 15.0  | 20 |
|                      | Subatrial triactine  | Paired   | 50.0        | 84.6   | 17.7  | 112.5  | 5.0        | 10.1 | 3.8  | 15.0  | 20 |
|                      |                      | Unpaired | 102.5       | 163.3  | 48.4  | 275.0  | 7.5        | 6.6  | 3.0  | 15.0  | 20 |
|                      | Subatrial tetractine | Paired   | 85.0        | 95.8   | 16.6  | 115.0  | 7.5        | 9.2  | 1.4  | 10.0  | 03 |
|                      |                      | Unpaired | 130.0       | 145.8  | 17.7  | 165.0  | 7.5        | 8.3  | 1.4  | 10.0  | 03 |
|                      |                      | Apical   | 25.0        | 49.2   | 25.0  | 75.0   | 7.5        | 9.5  | 1.4  | 10.0  | 03 |
|                      | Atrial tetractine    | Paired   | 75.0        | 93.8   | 26.5  | 112.5  | 7.5        | 7.5  | 0.0  | 7.5   | 02 |
|                      |                      | Unpaired | 62.5        | 8.89   | 8.8   | 75.0   | 7.5        | 7.5  | 0.0  | 7.5   | 02 |
|                      |                      | Apical   | 20.0        | 82.7   | 37.6  | 140.0  | 7.5        | 7.7  | 0.7  | 10.0  | 13 |
| ZMA Por 11562 (H)    | Diactine             |          | 256.0       | 1509.0 | ٠     | 2520.0 | 13.0       | 68.1 | 1    | 116.0 | 1  |
|                      | Cortical triactine   | Paired   | 21.0        | 35.0   | •     | 51.0   | 2.5        | 3.6  | 1    | 4.0   | 1  |
|                      |                      | Unpaired | 5.0         | 9.4    | •     | 18.0   | 2.0        | 4.1  | 1    | 0.9   | 1  |
|                      | Tubar triactine      | Paired   | 0.69        | 85.1   | •     | 117.0  | 7.0        | 8.8  | 1    | 13.0  | 1  |
|                      |                      | Unpaired | 52.0        | 79.5   | •     | 112.0  | 8.0        | 10.2 | 1    | 13.0  | 1  |
|                      | Subatrial triactine  | Paired   | 78.0        | 97.0   | 1     | 112.0  | 0.6        | 12.2 | 1    | 14.0  | 1  |
|                      |                      | Unpaired | 91.0        | 168.0  | •     | 203.0  | 0.6        | 12.4 | 1    | 15.0  | 1  |
|                      | Subatrial tetractine | Paired   | 0.97        | 109.0  | •     | 126.0  | 10.0       | 11.3 | 1    | 12.0  | 1  |
|                      |                      | Unpaired | 171.0       | 202.0  | •     | 241.0  | 8.0        | 10.2 | 1    | 11.0  | 1  |
|                      |                      | Apical   | 49.0        | 60.4   | •     | 71.0   | 0.9        | 7.1  | 1    | 8.0   | 1  |
|                      | Atrial tetractine    | Paired   | 0.89        | 104.0  | •     | 126.0  | 0.9        | 8.3  | 1    | 10.0  | 1  |
|                      |                      | Unpaired | 61.0        | 107.6  | •     | 165.0  | 7.0        | 8.6  | 1    | 11.0  | 1  |
|                      |                      | Apical   | 40.0        | 115.0  | 1     | 201.0  | 0.9        | 8.4  |      | 11.0  | '  |

Table 13. Species with reported occurrences in the Mascarenes Island ecoregion

| Species  | Locality   | Reference  |
|--|------------|--|
| subclass Calcinea                                    |            |  |
| Arturia compacta (Schuffner, 1877)                   | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,<br>2018         |
| Ascandra mascarenica sp. nov.                        | La Réunion | This study   |
| Ascandra oceanusvitae sp. nov.                       | La Réunion | This study   |
| Borojevia pirella Van Soest & De Voogd, 2018         | Rodrigues  | Van Soest & De Voogd, 2018                             |
| Clathrina luteoculcitella Wörheide & Hooper, 1999    | Rodrigues  | Van Soest & De Voogd, 2018                             |
| Clathrina rodriguesensis Van Soest & De Voogd, 2018  | Rodrigues  | Van Soest & De Voogd, 2018                             |
| Ernstia naturalis Van Soest & De Voogd, 2015         | Rodrigues  | Van Soest & De Voogd, 2018                             |
| Janusya indica gen. nov. sp. nov.                    | La Réunion | This study   |
| Lelapiella tertia sp. nov.                           | La Réunion | This study   |
| Leucaltis mauritiana Schuffner, 1877                 | Mauritius  | Schuffner, 1877 – it is considered a taxon inquirendum |
| Leucaltis nodusgordii (Poléjaeff, 1883)              | La Réunion | This study   |
| Leucascus tenuispinae sp. nov.                       | La Réunion | This study   |
| Leucetta chagosensis Dendy, 1913                     | La Réunion | This study   |
| Leucetta microraphis Haeckel, 1872                   | La Réunion | Bourmaud, 2003   |
| Leucetta sulcata Van Soest & De Voogd, 2018          | Rodrigues  | Van Soest & De Voogd, 2018                             |
| Pericharax orientalis Van Soest & De Voogd, 2015     | Mauritius  | Van Soest & De Voogd, 2018                             |
| Soleneiscus intermedius sp. nov.                     | La Réunion | This study   |
| Subclass Calcaronea                                  |            |  |
| Aphroceras seychellensis (Hôzawa, 1940) comb. nov.   | La Réunion | This study   |
| Leucandra claviformis Schuffner, 1877                | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,<br>2018         |
| Leucandra echinata Schuffner, 1877                   | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,<br>2018         |
| Leucandra falcigera Schuffner, 1877                  | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,<br>2018         |
| Leucandra ornata sp. nov.                            | La Réunion | This study   |
| Plectroninia hindei Kirk, 1900                       | La Réunion | Vacelet, 1977; Van Soest & De Voogd, 2018              |
| Sycettusa sycilloides (Schuffner, 1877)              | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,<br>2018         |
| Sycon oscari Van Soest & De Voogd, 2018 (as Sycandra | Mauritius  | Schuffner, 1877; Van Soest & De Voogd,                 |
| tabulata Schuffner, 1877)                            |            | 2018   |
| Tulearinia stylifera Vacelet, 1977                   | La Réunion | Vacelet, 1977  |
| Ute insulagemmae Van Soest & De Voogd, 2018          | La Réunion | This study   |

seven investigated localities, calcareous sponges were found in only three of them: Cap La Houssaye Reef, with one species (Leucandra ornata sp. nov.); Passe de l'Hermitage Reef, with eight species (Aphroceras seychellensis, Ascandra mascarenica sp. nov., A. oceanusvitae sp. nov., Janusya indica sp. nov., Leucaltis nodusgordii, Leucetta chagosensis, Lelapiella tertia sp. nov. and Ute insulagemmae); and Portail Caves, with three species (Leucascus tenuispinae sp. nov., Soleneiscus intermedius sp. nov. and Ute insulagemmae). One single site (Passe de l'Hermitage) presents most of the collected samples, confirming

that calcareous sponges require specific conditions to thrive. Further studies in other sites of the island could help understand the environmental requirements of calcareous sponges that may contribute to such distribution.

The species found in this work represent important findings for the systematics of Calcarea. Some of the new additions highlight inconsistencies in the traditional taxonomy, such as the paraphyly of orders and families in both Calcinea and Calcaronea (e.g. Voigt *et al.*, 2012; Klautau *et al.*, 2013; Lopes *et al.*, 2018). Considering the calcinean groups,

with the inclusion of *Lelapiella tertia* sp. nov. in the analysis, we once more recovered the paraphyly of Murrayonida (*Murrayona* and *Lelapiella*) and its placement among representatives of Clathrinida. For that reason, we opted to consider Murrayonida a junior synonym of Clathrinida, leaving Calcinea with a single monophyletic order. We were able to separate the problematic *Arturia l.s.* in two groups: the 'true' *Arturia* clade and the new genus *Janusya*, both with morphological and molecular support. However, some issues with other non monophyletic taxonomic groups, e.g. Leucaltidae (here represented by *Leucaltis* and *Ascandra*) and Clathrinidae, could still not be solved.

The two DNA markers currently available (C-LSU and ITS) for phylogenetic studies on Calcarea proved helpful for the integrative taxonomy. However, in some cases C-LSU provided inconclusive results because of the low evolutionary rate, being unable to separate close related species. For the subclass Calcaronea, the scenario is even harder, because only the C-LSU can be used, and it results in phylogenies that strongly conflict with the current morphological taxonomy. New markers are extremely important to provide a robust phylogenetic backbone and thus allow reorganization of taxa whose current markers render taxonomic decisions inconclusive.

## ACKNOWLEDGEMENTS

This work is part of the International Associated Laboratory MARRIO. It was supported by the Centre National de la Recherche Scientifique (CNRS), the Brazilian National Research Council (Conselho Nacional de Desenvolvimento Científico e Tecnológico [CNPq]) and by the Coordination for the Improvement of Higher Education Personnel (Coordenação de Aperfeiçoamento de Pessoal de Nível Superior [CAPES]). MK received a fellowship from CNPq (grant number: 305451/2017-0). MVL and GT received scholarships from CAPES (PROTAX, Edital 001/2015) and PIBIC, respectively. DEAL Réunion helped to finance the expedition, while the laboratory ENTROPIE of the Université de La Réunion provided material and equipements during the collections. TP had an internationally recognized certificate of compliance constituted from information on the permit or its equivalent made available to the Access and Benefit-sharing Clearing-House: TREL1928896S/253 (collection permission number). We thank the editors and anonymous reviewers for their constructive comments.

### REFERENCES

Alencar AR. 2012. Revisão taxonômica e filogenia da família Leucettidae de Laubenfels, 1936 (Porifera, Calcarea,

- Calcinea). Unpublished D. Phil. Thesis, Universidade Federal do Rio de Janeiro.
- Alvizu A, Eilertsen MH, Xavier JR, Rapp HT. 2018. Increased taxon sampling provides new insights into the phylogeny and evolution of the subclass Calcaronea (Porifera, Calcarea). Organisms Diversity & Evolution 18: 279–290.
- Azevedo F, Cóndor-Luján B, Willenz P, Hajdu E, Hooker Y, Klautau M. 2015. Integrative taxonomy of calcareous sponges (subclass Calcinea) from the Peruvian coast: morphology, molecules, and biogeography. Zoological Journal of the Linnean Society 173: 787–817.
- Azevedo F, Padua A, Moraes F, Rossi A, Muricy G, Klautau M. 2017. Taxonomy and phylogeny of calcareous sponges (Porifera: Calcarea: Calcinea) from Brazilian midshelf and oceanic islands. Zootaxa 4311: 301–344.
- Baine M, Harasti D. 2007. The marine life of Bootless Bay, Papua New Guinea. Port Moresby: Motupore Island Research Centre
- Bidder GP. 1898. The skeleton and the classification of calcareous sponges. *Proceedings of the Royal Society of London* 64: 61–76.
- Borojević R. 1966. Éponges calcaires des côtes de France. II. Le genre Ascandra Haeckel emend. Archives de Zoologie Expérimentale et Générale 107: 357–368.
- Borojević R. 1967. Éponges calcaires recueillies en Nouvelle-Calédonie par la Mission Singer-Polignac. Éditions da la Fondation Singer-Polignac, Paris 2: 1–12.
- Borojević R, Klautau M. 2000. Calcareous sponges from New Caledonia. *Zoosystema* 22: 187–201.
- Borojević R, Boury-Esnault N. 1986. Une novelle voie d'évolution chez les éponges Calcinea: description des genres Burtonella n.g. et Levinella n.g. Bulletin du Muséum National d'Histoire Naturelle, Paris 8: 443–455.
- Borojević R, Boury-Esnault N. 1987. Calcareous sponges collected by N.O. Thalassa on the continental margin of the Bay of Biscaye: I. Calcinea. In: Vacelet J, Boury-Esnault N, eds. *Taxonomy of Porifera from the NE Atlantic and Mediterranean Sea. NATO ASI Series, G13.* Berlin, Heidelberg: Springer-Verlag, 1–27.
- Borojević R, Boury-Esnault N, Vacelet J. 1990. A revision of the supraspecific classification of the subclass Calcinea (Porifera, Class Calcarea). Bulletin du Muséum National d'Histoire Naturelle (Paris) 2: 243–246.
- Borojević R, Boury-Esnault N, Vacelet J. 2000. A revision of the supraspecific classification of the subclass Calcaronea (Porifera, class Calcarea). Zoosystema 22: 203–263.
- Borojević R, Boury-Esnault N, Manuel M, Vacelet J. 2002. Order Clathrinida Hartman, 1958. In: Hooper JNA, Van Soest RWM, eds. Systema Porifera: a guide to the classification of sponges. New York: Kluwer Academic/ Plenum Publishers, 1141–1152.
- **Bourmaud C. 2003.** Inventaire de la biodiversité marine récifale à la Réunion. Rapport de l'Association Parc Marin et du Laboratoire d'Ecologie Marine, ECOMAR, de la Réunion.
- **Bowerbank JS. 1862.** On the anatomy and physiology of the Spongiadae. Part II. *Philosophical Transactions of the Royal Society of London*. London: The Royal Society, **152:** 747–836, pls XXVII–XXXV.

- **Bowerbank JS. 1866.** A monograph of the British Spongiadae, II. London: Ray Society.
- **Brøndsted HV. 1931.** Die Kalkschwämme. *Deutschen Südpolar Expedition 1901–3* **20:** 1–47.
- **Burton M. 1963.** A revision of the classification of the calcareous sponges. London: British Museum (Natural History).
- Carter HJ. 1878. On Teichonia, a new Family of calcareous sponges with description of two species. Annals and Magazine of Natural History 5: 35-40.
- Carter HJ. 1886. Descriptions of the sponges from the neighbourhood of Port Phillip Heads, South Australia, continued. Annals and Magazine of Natural History 15-18: 431-441.
- Cavalcanti FF, Rapp HT, Klautau M. 2013. Taxonomic revision of *Leucascus* Dendy, 1892 (Porifera: Calcarea) with revalidation of *Ascoleucetta* Dendy & Frederick, 1924 and description of three new species. *Zootaxa* 3619: 275-314
- Colin PL, Arneson C. 1995. Tropical Pacific invertebrates. A field guide to the marine invertebrates occurring on tropical pacific coral reefs, seagrass beds and mangroves. Beverly Hills: Coral Reef Research Foundation.
- Cóndor-Luján B, Louzada TS, Hajdu E, Klautau M. 2018. Morphological and molecular taxonomy of calcareous sponges (Porifera: Calcarea) from Curaçao, Caribbean Sea. Zoological Journal of the Linnean Society 182: 1–67.
- **Dendy A. 1891.** A monograph of the Victorian sponges. I. The organisation and classification of the Calcarea Homocoela, with description of the Victorian species. *Transactions of the Royal Society of Victoria* **3:** 1–81.
- **Dendy A. 1892.** Synopsis of the Australian Calcarea Heterocoela, with a proposed classification of the group and descriptions of some new genera and species. *Proceedings of the Royal Society of Victoria* **5:** 69–116.
- **Dendy A. 1905.** Report on the sponges collected by Professor Herdman at Ceylon in 1902. Reports on the Pearl Oyster Fisheries of the Gulf of Manaar 3: 59–246.
- Dendy A. 1913. Report on the calcareous sponges collected by
  H. M. S. 'Sealark' in the Indian Ocean. Transactions of the Linnean Society of London, Zoology 16: 1–29.
- **Dendy A, Frederick LM. 1924.** On a collection of sponges from the Abrolhos Islands, Western Australia. *Journal of the Linnean Society of London, Zoology* **16:** 1–29.
- Dendy A, Row H. 1913. The classification and phylogeny of the calcareous sponges, with a reference list of all the described species, systematically arranged. *Proceedings of* the Zoological Society of London 1913: 704–813.
- Dohrmann M, Voigt O, Erpenbeck D, Wörheide G. 2006. Nonmonophyly of most supraspecific taxa of calcareous sponges (Porifera, Calcarea) revealed by increased taxon sampling and partitioned Bayesian analysis of ribosomal DNA. *Molecular Phylogenetics and Evolution* 40: 830–843.
- $\label{eq:continuous} \textbf{Erhardt}\, \textbf{H}, \textbf{Baensch}\, \textbf{HA.}\, \textbf{1998}. \textit{Meerwasser atlas}\, 4. \textit{Wirbellose}.$  Melle: Mergus Verlag, 1214.
- **Felsenstein J. 1985.** Confidence limits on phylogenies: an approach using the bootstrap. *Evolution* **39:** 783–791.
- Ferrer-Hernandez F. 1918. Esponjas del litoral de Asturias. Trabajos del Museo Nacional de Ciencias Naturales 36: 1–39.

- Fontana T, Cóndor-Luján B, Azevedo F, Pérez T, Klautau M. 2018. Diversity and distribution of Calcareous sponges (subclass Calcinea) from Martinique. *Zootaxa* 4410: 331–369.
- George AM, Van Soest RWM, Sluka RD, Lazarus S. 2020. A checklist of marine sponges (Porifera) of peninsula India. Zootaxa 4885: 277–300.
- Gosliner TM, Behrens DW, Williams GC. 1996. Coral reef animals of the Indo-Pacific: animal life from Africa to Hawaii exclusive of the vertebrates. Monterey: Sea Challengers.
- Grant RE. 1836. Animal kingdom. In: Todd RB, ed. The encyclopedia of anatomy and physiology, Vol. 1. London: Sherwood, Gilbert and Piper, 107–118.
- **Gray JE. 1858.** Description of *Aphroceras*, a new genus of calcareous Spongiadae brought from Hong-Kong by Dr. Harland. *Proceedings of the Zoological Society of London* **26**: 113–114.
- **Haeckel E. 1870.** Prodromus eines Systems der Kalkschwämme. *Jenaische Zeitschrift* 5: 176–191.
- **Haeckel E. 1872.** *Die Kalkschwämme. Eine monographie, Vols* 1–3. Berlin: Reimer.
- **Hartman W. 1958.** A re-examination of Bidder's classification of the Calcarea. *Systematic Zoology* **7:** 97–110.
- Hess PN, Russo CAM. 2007. An empirical test of the midpoint rooting method. Biological Journal of the Linnean Society 92: 669–674.
- Hillis D, Morritz C, Mable B. 1996. Molecular systematics, 2nd edn. Sunderland: Sinauer Associates.
- Hoang DT, Chernomor O, von Haeseler A, Minh BQ, Vinh LS. 2017. UFBoot2: improving the ultrafast bootstrap approximation. Molecular Biology and Evolution 35: 518–522.
- **Hôzawa S. 1940.** Reports on the calcareous sponges obtained by the zoological institute and museum of Hamburg. *Science Reports of the Tohoku Imperial University* **15:** 131–163.
- Huelsenbeck JP, Ronquist F. 2001. MRBAYES: Bayesian inference of phylogenetic trees. *Bioinformatics* 17: 754–755.
- Jenkin CF. 1908. The Calcarea of the National Antarctic Expedition. Natural History Reports 4: 182–311.
- **Katoh K**, **Standley DM. 2013.** MAFFT multiple sequence alignment software v.7: improvements in performance and usability. *Molecular Biology and Evolution* **30:** 772–780.
- Katoh K, Toh H. 2008. Improved accuracy of multiple ncRNA alignment by incorporating structural information into a MAFFT-based framework. *BioMedCentral Bioinformatics* 9: 212.
- Kirkpatrick R. 1900. XLIII. Description of sponges from Funafuti. *Journal of Natural History* 6: 345–362.
- Kirkpatrick R. 1910. On a remarkable pharetronid sponge from Christmas Island. Proceedings of the Royal Society Series B 83: 124–133.
- Klautau M, Borojević R. 2001. Calcareous sponges from Arraial do Cabo–Brazil (I: the genus *Clathrina*). *Zoosystema* 23: 395–410.
- Klautau M, Valentine C. 2003. Revision of the genus Clathrina (Porifera, Calcarea). Zoological Journal of the Linnean Society 139: 1-62.

- Klautau M, Azevedo F, Cóndor-Luján B, Rapp H, Collins A, Russo C. 2013. A molecular phylogeny for the order Clathrinida rekindles and refines Haeckel's taxonomic proposal for calcareous sponges. *Integrative and Comparative Biology* 53: 447–461.
- Klautau M, Imešek M, Azevedo F, Pleše B, Nikolić V, Cetković H. 2016. Adriatic calcarean sponges (Porifera, Calcarea), with the description of six new species and a richness analysis. European Journal of Taxonomy 178: 1–52.
- Klautau M, Lopes MV, Guarabyra B, Folcher E, Ekins M, Debitus C. 2020. Calcareous sponges from the French Polynesia (Porifera: Calcarea). Zootaxa 4748: 261–295.
- Lanna E, Rossi AL, Cavalcanti FF, Hajdu E, Klautau M. 2007. Calcareous sponges from São Paulo State, Brazil (Porifera: Calcarea: Calcinea) with the description of two new species. Journal of the Marine Biological Association of the United Kingdom 87: 1553–1561.
- **de Laubenfels MW. 1936.** A discussion of the sponge fauna of the Dry Tortugas in particular and the West Indies in general, with material for a revision of the families and orders of the Porifera. *Tortugas Laboratory Occasional Papers* **467:** 1–225.
- von Lendenfeld R. 1885. A monograph of the Australian sponges. III. The Calcispongiae. Proceedings of the Linnean Society of New South Wales 9: 1083–1150.
- Lévi C. 1986. Laocaetis perion nov. sp., spongiaire hexactinellide Craticulariidae de l'océan Indien. Bulletin du Muséum National d'Histoire Naturelle 8: 437–442.
- **Lévi C. 1998.** Sponges of the New Caledonian lagoon. Paris: Éditions ORSTOM.
- Lopes MV, Cóndor-Luján B, Azevedo F, Pérez T, Klautau M. 2018. A new genus of calcareous sponge discovered in the Caribbean Sea: *Bidderia* gen. nov. (Porifera, Calcarea, Calcinea). *Zootaxa* 4526: 56–70.
- Manuel M, Borojević R, Boury-Esnault N, Vacelet J. 2002. Class Calcarea Bowerbank, 1864. In: Hooper JNA, Van Soest RWM, Willenz P, eds. *Systema Porifera*. Boston: Springer.
- Miklucho-Maclay N. 1868. Beiträge zur kenntniss der spongien. I. Über Guancha blanca, einen neven Kalkschwämm. Jenaische Zeitschrift 4: 221–240.
- Miller MA, Pfeiffer W, Schwartz T. 2012. The CIPRES science gateway: enabling high-impact science for phylogenetics researchers with limited resources. Proceedings of the 1st Conference of the Extreme Science and Engineering Discovery Environment: Bridging from the extreme to the campus and beyond. Chicago: ACM Press.
- Minchin EA. 1900. The Porifera. In: Ray Lankester E, ed. A treatise on zoology, Vol. 2. London: Adam & Charles Black.
- Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, von Haeseler A, Lanfear R. 2020. IQ-TREE 2: new models and efficient methods for phylogenetic inference in the genomic era. Molecular Biology and Evolution 37: 1530-1534.
- Pasnin O, Voigt O, Wörheide G, Murillo Rincón AP, Von der Heyden S. 2020. Indo-Pacific phylogeography of the lemon sponge *Leucetta chagosensis*. *Diversity* 12: 466.
- Pérez T, Díaz MC, Ruiz C, Cóndor-Luján B, Klautau M, Hajdu E, Lôbo-Hajdu G, Zea S, Pomponi SA,

- Thacker RW, Carteron S, Tollu G, Pouget-Culevier A, Thélamon P, Marechal JP, Thomas OP, Ereskovsky AV, Vacelet J, Boury-Esnault N. 2017. How a collaborative integrated taxonomic effort has trained new spongiologists and improved knowledge of Martinique Island (French Antilles, eastern Caribbean Sea) marine biodiversity. *PLoS One* 12: e0173859.
- Poléjaeff N. 1883. Report on the Calcarea dredged by the HMS Challenger during the years 1873–1876. Report on the Scientific Results of the Voyage Challenger (Zoology) 8: 1–76
- Pulitzer-Finali G. 1982. Some new or little-known sponges from the Great Barrier Reef of Australia. Bollettino dei Musei e degli Istituti Biologici dell'Universitá di Genova 48: 87–141.
- Rapp HT. 2004. The first record of the genus *Leucascus*Dendy, 1892 from the Atlantic Ocean, with description of *Leucascus lobatus* sp. nov. (Porifera, Calcarea, Calcinea) from
  Greenland. *Steenstrupia* 28: 1–9.
- Rapp HT. 2006. Calcareous sponges of the genera *Clathrina* and *Guancha* (Calcinea, Calcarea, Porifera) of Norway (NE Atlantic) with the description of five new species. *Zoological Journal of the Linnean Society* 147: 331–65.
- Rapp HT, Janussen D, Tendal OS. 2011. Calcareous sponges from abyssal and bathyal depths in the Weddell Sea, Antarctica. Deep Sea Research Part II: Topical Studies in Oceanography 58: 58–67.
- Ridley SO. 1884. Spongiida. In: Report on the zoological collections made in the Indo-Pacific Ocean during the voyage of HMS Alert, 1881–2. London: British Museum (Natural History), 366–482.
- Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19: 1572–1574.
- Russo CAM. 1997. Efficiencies of different statistical tests in supporting a known vertebrate phylogeny. Molecular Biology and Evolution 14: 1078–1080.
- Sambrook J, Fritsch EF, Maniatis T. 1989. Molecular cloning: a laboratory manual. New York: Cold Spring Harbor Laboratory Press.
- Schleyer MH, Benayahu Y, Parker-Nance S, Van Soest RWM, Quod JP. 2016. Benthos on submerged lava beds of varying age off the coast of Reunion, western Indian Ocean: sponges, octocorals and ascidians. *Biodiversity* 17: 93–100.
- Schmidt O. 1862. Die Spongien des Adriatischen Meeres. Leipzig: Wilhelm Engelmann.
- Schmidt O. 1864. Supplement der Spongien des Adriatischen Meeres. Enthaltend die Histologie und systematische Ergänzungen. Leipzig: Wilhelm Engelmann.
- Schmidt O. 1870. Grundzüge einer Spongien-Fauna des atlantischen Gebietes. Leipzig: Wilhelm Engelmann.
- Schuffner O. 1877. Beschreibung einiger neuer Kalkschwämme. Jenaische Zeitschrift 4: 403–433.
- Spalding MD, Foz HE, Allen GR, Davidson N, Ferdaña ZA, Finlayson M, Halpern BS, Jorge MA, Lombana A, Lourie SA, Martin KD, McManus E, Molnar J, Recchia CA, Robertson J. 2007. Marine ecoregions of

- the world: a bioregionalization of coastal and shelf areas. *Bioscience* **57:** 573–583.
- Tanita S. 1942. Calcareous sponges collected in the Kanto District, Japan. Science Reports of the Tôhoku Imperial University 17: 17-69.
- Tanita S. 1943. Studies on the Calcarea of Japan. Science Reports of the Tôhoku Imperial University 17: 353–490.
- **Thacker AG. 1908.** On collections of the Cape Verde Islands fauna made by Cyril Crossland, M.A. The calcareous sponges. *Proceedings of the Zoological Society of London* **49:** 757–782.
- **Trentin F**, **Massé L. 2018.** Première approche de la biodiversité des éponges à la Réunion. *Rapport Vie Océane*. La Réunion: University of La Réunion.
- Vacelet J. 1977. Éponges pharétronides actuelles et sclérosponges de Polynésie française, de Madagascar et de la Réunion. Bulletin du Muséum National d'Histoire Naturelle de Paris 444: 345–368.
- Vacelet J. 1981. Éponges hypercalcifées ('Pharétronides' 'Sclérosponges') des cavités des récifs coralliens de Nouvelle-Calédonie. Bulletin du Muséum National d'Histoire Naturelle 3: 313–351.
- Van Soest RWM, De Voogd NJ. 2015. Calcareous sponges of Indonesia. Zootaxa 3951: 1–105.
- Van Soest RWM, De Voogd NJ. 2018. Calcareous sponges of the western Indian Ocean and Red Sea. Zootaxa 4426: 1–160.
- Van Soest RWM, Kaiser K, Van Syoc R. 2011. Sponges from Clipperton Island, East Pacific. Zootaxa 2839: 1–46.
- Van Soest RWM, Boury-Esnault N, Vacelet J, Dohrmann M, Erpenbeck D, De Voogd NJ, Santodomingo N, Vanhoorne B, Kelly M, Hooper JNA. 2012. Global diversity of sponges (Porifera). PLoS One 7: e35105.
- Van Soest RWM, Hooper JN, Butler PJ. 2020. Every sponge its own name: removing Porifera homonyms. Zootaxa 4745: 1–93.
- Voigt O, Wörheide G. 2016. A short LSU rRNA fragment as a standard marker for integrative taxonomy in calcareous sponges (Porifera: Calcarea). *Organisms, Diversity & Evolution* 16: 53–64.

- Voigt O, Wülfing E, Wörheide G. 2012. Molecular phylogenetic evaluation of classification and scenarios of character evolution in calcareous sponges (Porifera, Class Calcarea). *PLoS One* 7: 1–16.
- Voigt O, Erpenbeck D, Gonzalez-Pech RA, Al-Aidaroos AM, Berumen ML, Wörheide G. 2017. Calcinea of the Red Sea: providing a DNA barcode inventory with description of four new species. *Marine Biodiversity* 47: 1009–1034.
- Wörheide G, Hooper JNA. 1999. Calcarea from the Great Barrier Reef. I: cryptic Calcinea from Heron Island and Wistari Reef (Capricorn-Bunker group). *Memoirs of the* Queensland Museum 43: 859–891.
- Wörheide G, Hooper JNA. 2003. New species of Calcaronea (Porifera: Calcarea) from cryptic habitats of the southern Great Barrier Reef (Heron Island and Wistari Reef, Capricorn-Bunker Group, Australia). Journal of Natural History 37: 1–47.
- Wörheide G, Hooper JNA, Degnan BM. 2002. Phylogeography of western Pacific Leucetta 'chagosensis' (Porifera: Calcarea) from ribosomal DNA sequences: implications for population history and conservation of the Great Barrier Reef World Heritage Area (Australia). Molecular Ecology 11: 1753–1768.
- Wörheide G, Nichols S, Goldberg J. 2004. Intragenomic variation of the rDNA internal transcribed spacers in sponges (phylum Porifera): implications for phylogenetic studies. *Molecular Phylogenetics and Evolution* 33: 816–830.
- Wörheide G, Solé-Cava AM, Hooper JNA. 2005. Biodiversity, molecular ecology and phylogeography of marine sponges: patterns, implications and outlooks. *Integrative and Comparative Biology* 45: 377–385.
- Wörheide G, Epp SL, Macis L. 2008. Deep genetic divergences among Indo-Pacific populations of the coral reef sponge *Leucetta chagosensis* (Leucettidae): founder effects, vicariance, or both? *BMC Evolutionary Biology* 8: 24.

## SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's web-site.

Figure S1. ITS sequences aligned through the online version of the program MAFFT v.7 using the strategy Q-INS-i.

**Figure S2.** C-LSU sequences of Calcinea aligned through the online version of the program MAFFT v.7 using the strategy Q-INS-i.

**Figure S3.** C-LSU sequences of Calcaronea aligned through the online version of the program MAFFT v.7 using the strategy Q-INS-i.

**Figure S4.** ML tree based on the C-LSU sequences of the subclass Calcinea rooted with the subclass Calcaronea. **Figure S5.** ML tree based on the C-LSU sequences of the subclass Calcaronea rooted with the subclass Calcinea.