Intra- and interspecific shell variability of the genus
Urocythereis Ruggieri, 1950 (Ostracoda: Hemicytheridae)
in the La Strea Bay (Ionian Sea, Italy)

Giuseppe AIELLO¹*, Diana BARRA², Roberta PARISI³

¹²³ Dipartimento di Scienze della Terra, dell’Ambiente e delle Risorse, Università degli
Studi di Napoli Federico II, Largo San Marcellino 10, 80138 Napoli, Italy.

*Corresponding author: aie64llo@hotmail.com
² Email: dibarra@unina.it
³ Email: rohyparisi@alice.it

Abstract. The variability of the reticulum pattern, ornamentation and outline of the Urocythereis populations of the La Strea Bay is analysed. The results show that the shell features of the form U. distinguenda (Neviani, 1928) (= U. oblonga Brady, 1866) have to be included within the high variability range of U. margaritifera (G.W. Müller, 1894). Consequently, it is suggested that in the upper infralittoral waters of the inlet two (and not three, as stated in previous investigations) species of the genus Urocythereis presently live. A second form, displaying a relatively low variability, is described as a new species, U. ilariae sp. nov.

Keywords. Intraspecific variability, shell morphology, Urocythereis, celation, outline analysis.

Introduction

During a study of the Recent ostracods of the La Strea Bay more than two thousand valves of Urocythereis were collected and assigned to three species (Aiello et al. 2006). Most specimens were assigned to U. margaritifera (G.W. Müller, 1894) and U. distinguenda (Neviani, 1928), while a third form was left in open nomenclature and named Urocythereis sp. 1.

The presence of shells exhibiting transitional characters between the former two species (then included in U. margaritifera) indicated an unsolved taxonomic issue. For a correct evaluation of the degree of intraspecific variability in Urocythereis, we have re-examined sub-recent specimens recovered in this relatively small (length: 2.5 km, maximum width: 1 km) inlet, also known as Porto Cesareo Lagoon. It
is in its northern part connected with the Ionian Sea, and is characterised by a subtropical biocoenosis (Parenzan 1976, 1983, 1984). The environmental setting is entirely included in the upper part of the infralittoral zone, with water depth not exceeding five meters. Ecological parameters vary over seasons and sectors of the bay (Fig. 1).

It can be supposed that environmental factors influence, at least in part, the development of different shell morphs. Consequently, the studied specimens may represent slightly different ecomorphotypes belonging to dead assemblages.

**Fig. 1.** Location map of the study area and sampling stations and range of selected physico-chemical parameters of the La Strea Bay (data from Belmonte & Rubino 1988; Mercurio *et al.* 2000; Gherardi *et al.* 2001).
With the aim to minimize the unavoidable taxonomist’s subjectivity, we have employed two morphological methods of discrimination: the first is the comparison of ornament morphology, taking into account the features of the reticulation especially; the other is the morphometric analysis of the outline by means of the computer program Morphomatica (Linhart et al. 2006).

To be more precise, our intent is: 1) to define intrademic, non-polytypic (*sensu* Sylvester-Bradley 1976) variations of shell features, namely the morphological variability of the *Urocythereis* demes - panmictic populations (freely interbreeding, sharing the same gene pool, relatively confined) of the La Strea Bay; 2) to assess how many *Urocythereis* species are present in the Recent/sub-Recent thanatocoenoses of the bay; we consider this attempt as a step toward a better understanding of the complex morphology of the genus from a paleontological viewpoint; 3) to characterize groups of fossae, stable within species and variable among species (e.g., Al-Furaih 1977).

**Intraspecific variability in ostracod shells**

The studies on intraspecific morphological variations include investigations on soft parts (*i.a.*, Rossetti & Martens 1996; Martens et al. 1998; Yin et al. 1999), internal features of smooth valves (*i.a.*, Aiello et al. 2007, and references therein), surface ornamentation (*i.a.*, Keen 1982; Pey pouquet et al. 1988; Neil 2000) and outline (Baltanas & Danielopol 2011, and references therein). Intraspecific variability can be continuous or discontinuous, and continuous variations may seem very simple to analyse if compared with polymorphism *sensu* Ford (1940), that is “the occurrence together in the same locality of two or more discontinuous forms of a species in such proportions that the rarest of them cannot be maintained merely by recurrent mutation” (see Clark 1976 for discussion and extensive references). Yet, when we deal with morphologically highly variable taxa, also non-polymorphic species highlight complex taxonomic problems.

Here, we focus on the continuous variation of shape and ornament in *Urocythereis*, with special regard to reticulation. *Urocythereis* species may show all five types of continuous variation of fine sculpture (in position, form, strength of expression, size and number of elements) described by Liebau (1971).

High-degree variability in ostracod shell sculpture, derived from both genetic and environmental factors, is a recurrent problem experienced by authors dealing with studies on shallow marine assemblages (e.g., Hartmann 1982), which always include some phenotypically plastic species. Phenotypic plasticity, that is the ability of an organism to express different phenotypes under different environmental conditions (see, e.g., West-Eberhard 1989), may represent a fundamental resource for effective adaptation to coastal environments, characterized by wide ranges and rapid changes of ecological parameters, and is a major drawback for systematic investigations. Seemingly distinct features occurring in some infra-littoral ostracod taxa lead taxonomists to propose a large number of specific names, which frequently have not stood the test of time. In other cases, taxonomists “lumped” together very different forms (see, e.g., a brief discussion on the genus *Carinocythereis* in Aiello & Szczechura 2001).

**The genus Urocythereis**

The genus *Urocythereis* was originally described by Ruggieri in 1950, with the intent to accommodate three strictly related hemicytherid species, with subrectangular valves and an amphidont hinge provided with a reniform posterior tooth in the right valve. These species were *Cytherina favosa* Roemer, 1838, designated as type species, *Cythereis margaritifera* G.W. Müller, 1894 and *Cythereis distinguenda* Neviani, 1928, from the Pliocene to Recent of the Mediterranean area. Since then, dozens of species have been included in *Urocythereis*, especially from the Neogene to Recent of the Mediterranean area and the eastern Atlantic. Aiello et al. (2004) assigned the south-western Atlantic species, previously included in *Urocythereis*, to the new genus *Ruggiericythere*, due to the different structure of the reticulation-ridges system. The allied genus *Yezocythere* Hanai & Ikeya, 1991 includes eastern Asian
species with a low subcentral tubercle, lacking in *Urocythereis*. Even though the authors recognized a second distinctive feature, the smaller size of the median frontal scar, the value of this character at genus level remained questionable. The Neogene to Recent Eastern African and Asian records of *Urocythereis* (e.g., *U. pohangensis* Huh & Whatley, 1997; *U. salebrosa* Ahmad, Neale & Siddiqui, 1991), as well as its presence in Paleogene deposits (e.g., *U. bertelsae* Kielbowicz, 1988), are, in our opinion, questionable. In our present state of knowledge, *Urocythereis* is a Cenozoic genus possibly restricted to the infralittoral and uppermost circalittoral zone (0–40 m, Athersuch 1977) of the Eastern Atlantic-Mediterranean area. The taxonomy of this genus is complex due to the unusually wide variability of the shell ornamentation and, consequently, species limits within the genus remain partly undefined.

A major contribution to the knowledge of the genus is the study by Athersuch (1977), who provided a revision of all the then known living and some fossil species, in addition to an excellent iconography. He described detailed ornamental patterns of five allied species (*U. distinguenda* Neviani, 1928), *U. margaritifera*, *U. favosa* (Roemer, 1838), plus *U. neapolitana* and *U. britannica* Athersuch, 1977, described as new species) based on “nine arbitrary groups” of fossae. The author used these patterns to demonstrate differences among groups of species, one being later described by Ruggieri & Russo (1980) as a new genus, *Nonurocythereis*. These groups are not effective for discrimination at species level. The outline of the muri, as well as the shape and dimension of the fossae, and their degree of anastomosis, are rarely steady, so that reticulation varies “even between members of the same species” (Athersuch 1977). Uncertainty is increased due to the frequent occurrence of celtation (Sylvester-Bradley & Benson 1971), a morphological noise able to conceal, in some cases almost completely, ornamental patterns. In a similar way, the relationship between normal pores and reticulation, used by authors for ostracod morphology studies (e.g., Hunt & Yasuhara 2010), is blurred by various degrees of development of shell calcification. Normal pores are well distinct in internal view, not in external view, and consequently the relationship between pores and reticulum pattern is difficult to establish.

**Reticulation**

Shell reticulation represents a complex of features that, since the early contributions of Pokorny (1969a, 1969b), Liebau (1969, 1971) and Benson (1971, 1972), has been considered by ostracod workers a fascinating object of study for taxonomy, evolution and related research fields. Ornament pattern definition has been carried out especially on hemicytherid and trachyleberid taxa, frequently showing a well-defined and more or less stable reticulation. Interspecific variations, in related species, are sometimes limited to a few fossae and highly consistent in each species (Okada 1982b). The analysis of the fossal pattern has been used for species discrimination (e.g., Al-Furaih 1977) and in studies on the evolution and ontogeny of Trachybelleridoidea (*i.a.*, Hunt 2007a, 2007b; Hunt & Yasuhara 2010; Tanaka et al. 2011). Jones (1988) showed the possibility to extend the reticulation analysis to Paleozoic taxa.

Okada (1981, 1982a, 1982b) extensively studied the relationship of the epidermal cells to the reticulation pattern and he recognized a direct correspondence between fossae and the underlying epidermal cells. Since the muri represent the boundary of adjacent cells, the mesh arrangement reflects cell lineages, commonly considered as genetically determined (*i.a.*, Liebau 1971; Benson 1972; Irizuki 1993; Hunt 2007a).

Reticulated species frequently show a pattern of ridges where homologous fossae can be recognized by their constancy in shape, number and arrangement and by the presence of landmarks such as pores, ridges or spines (*i.a.* Benson 1972; Liebau 1991; Hunt 2007b; Hunt & Yasuhara 2010). In *Urocythereis*, the number of fossae is relatively stable within species, while their shape and arrangement vary widely, pores are frequently obscured by calcification and the features of anatomical landmarks, for example muscle scars, can be difficult to define. Furthermore, alae, carinae or tubercles are not present in *Urocythereis*. Consequently, an unambiguous identification of fossae and an accurate quantitative study
of their variations (e.g., Hunt & Yasuhara 2010 on *Poseidonamicus*) has proved unattainable. Ruiz *et al.* (2006) hypothesized a direct influence of environmental factors on the variability of *Urocythereis* reticulation.

**Celation as a taxonomic issue**

The development of an outer layer of calcite, the tegmen, overlapping the ornament of the valves, has been named “celation” by Sylvester-Bradley & Benson (1971). The authors considered the tegmen as a secondary thickening paving over the outer surface and thus obscuring, to some degrees, primary reticulation. Benson (1972) hypothesized that the development of the tegmen represents the architectural response to the requirement of a high shell strength. In deep and cold waters, this lightweight solution may represent a case of parsimony in design balancing the scarcity of available skeletal material with the development of high resistance to compression. Athersuch (1977) observed that the development of celation (not recorded in juveniles) in *Urocythereis distingueda* is related neither to depth nor to other known ecological parameters, and suggested that the growth of the tegmen depends from the age of the individuals: an intense celation would be typical of older adults. Another hypothesis considers celation as a response to calcium saturated bottom waters, pertaining to the “environmentally-cued polymorphism” described by Peypouquet *et al.* (1980, 1981, 1988). Anyway, because celation varies continuously, we agree with Neil (2000) on preferring the term “environmentally-cued variation”. In this particular case the terms “celation” and “agradation” (“aggradation” in Neil 2000) are equivalent. These two hypotheses are not mutually exclusive.

The ecological demands of an infralittoral genus can hardly be compared with the bathyal and abyssal taxa studied by Benson (1972), who supposed that the secondary development of the top of the muri in shallow water species, such as *Cythere lutea* O.F. Müller, 1785 and *Hemicythere villosa* (Sars, 1866), has to be an adaptation to high energy sedimentary regimens. The attested preference of *Urocythereis* for sandy bottoms (e.g., Athersuch *et al.* 1989; Aiello *et al.* 2006) and relatively agitated waters supports this interpretation. It is understood that celation may be a consequence of simple genetic variation, or a mixture of genetic and environmental effects. In any case, the occurrence of specimens with more or less celated shells poses a taxonomic problem, whether we are dealing with two species (in this case *U. margaritifera* and *U. distingueda*), with individuals of the same species that have possibly built their valves in different physico-chemical conditions, or, alternatively, with younger and older adults of the same species.

**How many *Urocythereis* species occur in the La Strea Bay?**

The assignment of the *Urocythereis* specimens of the La Strea Bay to the above mentioned three species by Aiello & Barra (in Aiello *et al.* 2006) followed the original description and figures of G.W. Müller (1894) and subsequent literature for *U. margaritifera*, and Athersuch (1977) for *U. distingueda*. *Cythereis margaritifera* was described by G.W. Müller in his monograph on the ostracods living in the Gulf of Naples, and later recorded from Pliocene to Recent in the Mediterranean area. *Cythereis distingueda* is the name proposed by Neviani (1928) to replace the pre-occupied *Cythere oblonga* Brady, 1866 (non *C. oblonga* M’Coy, 1844). Athersuch (1977) discussed nomenclatural problems and designated a neotype for *U. distingueda* from the shallow waters of Cyprus. This author figured *U. margaritifera* specimens from Müller’s collection, showing individuals with rounded fossae as well as with fossae reduced to punctae and foveolae (*sensu* Athersuch *et al.* 1989) because of celation. In the same paper the specimens assigned to *U. distingueda* show a marked celation with small-sized irregular punctae-foveolae.

A third form present in the La Strea Bay bottom samples was left in open nomenclature and named *U. sp. 1* because of the resemblance with the specimens from the beach sands of Tripoli figured by
Barra (1997). This previously undescribed species has a marked affinity with *U. exedata* Uliczny, 1969, especially in the very distinct ocular riblet, a feature homologous to the prominent “ocular ridge” described by Benson (1972) on some *Bradleya* species. The new species shows a distinct ornamental pattern and specific features. Conversely, it seems very difficult to define an unambiguous boundary between *U. distinguenda* and *U. margaritifera*, necessitating the examination of “intermediate” morphs, in order to verify the existence of two separate species rather than a single highly variable species.

**Material and methods**

All the 32 samples were taken within the depth range of 0.12–4.78 m bsl (Aiello *et al.* 2006), from the Recent bottom sediments of the La Strea Bay (Table 1). The samples contained 2133 valves and 128 carapaces, including both adults and young instars. All the specimens were studied and 150 shells were

<table>
<thead>
<tr>
<th>Samples</th>
<th>Longitude E</th>
<th>Latitude N</th>
<th>Depth (m)</th>
<th>Grain size</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>17°53.67'</td>
<td>40°15.92'</td>
<td>0.43</td>
<td>FS</td>
</tr>
<tr>
<td>A2</td>
<td>17°53.76'</td>
<td>40°15.83'</td>
<td>1.30</td>
<td>MS-FS</td>
</tr>
<tr>
<td>B1</td>
<td>17°53.57'</td>
<td>40°15.64'</td>
<td>3.21</td>
<td>CS-FS</td>
</tr>
<tr>
<td>B2</td>
<td>17°53.64'</td>
<td>40°15.71'</td>
<td>2.43</td>
<td>CS-MS</td>
</tr>
<tr>
<td>B3</td>
<td>17°53.73'</td>
<td>40°15.78'</td>
<td>2.40</td>
<td>CS</td>
</tr>
<tr>
<td>B4</td>
<td>17°53.87'</td>
<td>40°15.88'</td>
<td>1.40</td>
<td>CS</td>
</tr>
<tr>
<td>B5</td>
<td>17°53.93'</td>
<td>40°15.91'</td>
<td>0.90</td>
<td>CS-MS</td>
</tr>
<tr>
<td>C1</td>
<td>17°53.52'</td>
<td>40°15.34'</td>
<td>4.40</td>
<td>CS</td>
</tr>
<tr>
<td>C2</td>
<td>17°53.61'</td>
<td>40°15.40'</td>
<td>4.78</td>
<td>CS</td>
</tr>
<tr>
<td>C3</td>
<td>17°53.72'</td>
<td>40°15.46'</td>
<td>4.50</td>
<td>CS-FS</td>
</tr>
<tr>
<td>C4</td>
<td>17°53.83'</td>
<td>40°15.53'</td>
<td>3.40</td>
<td>CS</td>
</tr>
<tr>
<td>C5</td>
<td>17°53.92'</td>
<td>40°15.61'</td>
<td>2.58</td>
<td>CS-FS</td>
</tr>
<tr>
<td>C6</td>
<td>17°54.02'</td>
<td>40°15.66'</td>
<td>1.73</td>
<td>CS</td>
</tr>
<tr>
<td>C7</td>
<td>17°54.10'</td>
<td>40°15.72'</td>
<td>-</td>
<td>CS</td>
</tr>
<tr>
<td>D1</td>
<td>17°53.85'</td>
<td>40°15.33'</td>
<td>1.52</td>
<td>MS-FS</td>
</tr>
<tr>
<td>D2</td>
<td>17°54.01'</td>
<td>40°15.38'</td>
<td>2.30</td>
<td>CS-MS</td>
</tr>
<tr>
<td>D3</td>
<td>17°54.17'</td>
<td>40°15.49'</td>
<td>1.51</td>
<td>CS-FS</td>
</tr>
<tr>
<td>D4</td>
<td>17°54.32'</td>
<td>40°15.59'</td>
<td>1.11</td>
<td>CS-FS</td>
</tr>
<tr>
<td>E3</td>
<td>17°54.01'</td>
<td>40°15.07'</td>
<td>1.42</td>
<td>MS-FS</td>
</tr>
<tr>
<td>E4</td>
<td>17°54.25'</td>
<td>40°15.16'</td>
<td>1.50</td>
<td>CS-MS</td>
</tr>
<tr>
<td>E5</td>
<td>17°54.26'</td>
<td>40°15.23'</td>
<td>1.49</td>
<td>FS</td>
</tr>
<tr>
<td>E6</td>
<td>17°54.45'</td>
<td>40°15.28'</td>
<td>1.30</td>
<td>MS-FS</td>
</tr>
<tr>
<td>F1</td>
<td>17°54.28'</td>
<td>40°14.84'</td>
<td>0.50</td>
<td>CS-FS</td>
</tr>
<tr>
<td>F2</td>
<td>17°54.34'</td>
<td>40°14.88'</td>
<td>0.94</td>
<td>MS-FS</td>
</tr>
<tr>
<td>F3</td>
<td>17°54.45'</td>
<td>40°14.97'</td>
<td>1.12</td>
<td>FS</td>
</tr>
<tr>
<td>F4</td>
<td>17°54.60'</td>
<td>40°15.04'</td>
<td>1.50</td>
<td>MS-FS</td>
</tr>
<tr>
<td>G1</td>
<td>17°54.47'</td>
<td>40°15.63'</td>
<td>0.20</td>
<td>MS-VFS</td>
</tr>
<tr>
<td>G2</td>
<td>17°54.52'</td>
<td>40°14.67'</td>
<td>0.70</td>
<td>MS</td>
</tr>
<tr>
<td>G3</td>
<td>17°54.57'</td>
<td>40°14.71'</td>
<td>0.80</td>
<td>MS-VFS</td>
</tr>
<tr>
<td>G4</td>
<td>17°54.66'</td>
<td>40°14.77'</td>
<td>0.65</td>
<td>MS-VFS</td>
</tr>
<tr>
<td>H</td>
<td>17°54.69'</td>
<td>40°14.61'</td>
<td>0.26</td>
<td>VFS</td>
</tr>
<tr>
<td>I</td>
<td>17°54.69'</td>
<td>40°14.51'</td>
<td>0.12</td>
<td>FS</td>
</tr>
</tbody>
</table>

Table 1. Coordinates, depth and granulometry of the studied samples. Abbreviations: VSF = Very Fine Sands, FS = Fine Sands, MS = Medium Sands, CS = Coarse Sands.
selected for analyses and iconography. This study deals exclusively with adult specimens. Young instars were considered only to evaluate the possible occurrence of celation in juveniles.

The analysis of the reticulation pattern and celation variations is based on scanning electron microscopy (SEM) micrographs (carried out at Cisag, Università di Napoli Federico II). The above mentioned high variability of the reticulation in *Urocythereis* species lead us to select some areas of the shell, where the fossae-muri pattern was detectable with a minimum degree of uncertainty.

**Fig. 2.** Scheme of the five fossal groups used for reticulation pattern analysis. White arrow indicates the pre-ocular bridge. **A.** *Urocytheris margaritifera* (G.W. Müller, 1894), ABMC 2014/057. **B.** *Urocytheris ilariae* sp. nov., ABMC 2014/047.
The Morphomatica test was performed on drawings in transmitted light. They were carried out by means of a Visopan Reichert, with a magnification of 201 times. The pictures were drawn with a technical pen (thickness of the line: 0.30 mm) on tracing paper. The studied specimens are housed in the Aiello Barra Micropaleontological Collection (ABMC), Distar, Università degli Studi di Napoli Federico II.

Table 2. Catalogue number and sampling stations of the specimens used in the Morphomatica analysis.

<table>
<thead>
<tr>
<th>Catalogue number</th>
<th>Samples</th>
<th>Catalogue numbers</th>
<th>Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABMC 2014/125</td>
<td>C4</td>
<td>ABMC 2014/142</td>
<td>C5</td>
</tr>
<tr>
<td>ABMC 2014/126</td>
<td>C4</td>
<td>ABMC 2014/143</td>
<td>C5</td>
</tr>
<tr>
<td>ABMC 2014/127</td>
<td>D3</td>
<td>ABMC 2014/144</td>
<td>D3</td>
</tr>
<tr>
<td>ABMC 2014/128</td>
<td>D3</td>
<td>ABMC 2014/145</td>
<td>D3</td>
</tr>
<tr>
<td>ABMC 2014/129</td>
<td>E6</td>
<td>ABMC 2014/146</td>
<td>D3</td>
</tr>
<tr>
<td>ABMC 2014/130</td>
<td>E6</td>
<td>ABMC 2014/147</td>
<td>D3</td>
</tr>
<tr>
<td>ABMC 2014/131</td>
<td>E6</td>
<td>ABMC 2014/148</td>
<td>E6</td>
</tr>
<tr>
<td>ABMC 2014/132</td>
<td>E6</td>
<td>ABMC 2014/149</td>
<td>E6</td>
</tr>
<tr>
<td>ABMC 2014/133</td>
<td>E6</td>
<td>ABMC 2014/150</td>
<td>E6</td>
</tr>
<tr>
<td>ABMC 2014/134</td>
<td>H</td>
<td>ABMC 2014/151</td>
<td>G2</td>
</tr>
<tr>
<td>ABMC 2014/135</td>
<td>H</td>
<td>ABMC 2014/152</td>
<td>G2</td>
</tr>
<tr>
<td>ABMC 2014/136</td>
<td>A2</td>
<td>ABMC 2014/153</td>
<td>G2</td>
</tr>
<tr>
<td>ABMC 2014/137</td>
<td>A2</td>
<td>ABMC 2014/154</td>
<td>H</td>
</tr>
<tr>
<td>ABMC 2014/138</td>
<td>A2</td>
<td>ABMC 2014/155</td>
<td>H</td>
</tr>
<tr>
<td>ABMC 2014/139</td>
<td>B3</td>
<td>ABMC 2014/156</td>
<td>H</td>
</tr>
<tr>
<td>ABMC 2014/140</td>
<td>C4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Reticulation pattern

In a highly variable genus such as Urocythereis the purpose of naming the position of all the meshes seems difficult to achieve (in Urocythereis especially in the ventral area and in the posterior half of the shell). Consequently, we identified five arbitrary fossal groups where the pattern can be recognized, both in different species and specimens of the same species, as homologous without uncertainties (Fig. 2).

These groups were selected with the aim to define the structure of inter- and intra-specific reticulation pattern variations. We chose muri/fossae sets that could be reliably considered homologous in U. margaritifera and U. exedata. The two species are quite different and we hope that these groups may be used for future comparisons among other species of Urocythereis.

Fossal groups are defined as follows:

a) Pre-ocular fossa (Pre-ocular bridge): presence of a preocular connection between the anterior marginal rim and the ocular riblet, as figured in Fig. 2.

b) Anterodorsal group: a group of fossae located in the subocular area (B1, B2, B3, B4, C1, C2, C3, C4) as numbered in Figs 2 and 3.

c) D3-D4-E9 group: these fossae form a triangular area situated in the anterocentral area, behind the fourth concentric anterior ridge as figured in Figs 2 and 4.

d) Dorsal median group: Athersuch (1977) defined this group as “two vertical rows of fossae directly beneath the post-ocular sinus and directed towards the sub-central area” (α1–α6, β1–β5 in Figs 2 and 5);

e) Caudal group: the group of fossae (Cg1–Cg4) located on the caudal process area as figured in Figs 2 and 6.


Outline analysis

A quantitative analysis of ostracod morphology has been carried out by means of different methods, including techniques of outline analysis (for a partial review, see Baltanás & Danielopol 2011). In recent years, ostracod researchers seem leaning towards using the computer program Morphomatica, based on a B-spline algorithm specifically adapted to ostracod valve outline (Baltanás et al. 2003; Linhart et al. 2006), which provides valuable results in specific discrimination (i.a., Iepure et al. 2007; Danielopol et al. 2008; Ligios & Gliozzi 2012; Mazzini et al. 2014). The method has been described, for example, in Gross et al. 2008. This technique has been used mostly on smooth or weakly ornamented ostracod taxa, whereas we tested its effectiveness by using it to discriminate strongly reticulated forms. The material consists of 20 left valves of the Urocythereis margaritifera-distinguenda group and 11 left valves of U. ilariae sp. nov. (Table 2). On the basis of the shell features a clear discrimination between female and male valves is not possible. The selection of the valves showing a higher h/l ratio would be, in absence of discontinuity, a mere example of circular reasoning. Consequently, we selected the best-preserved valves, with no regard for their supposed sex, well-aware of the resulting limitations. Due to the presence of unevenly preserved marginal denticleation we chose, unlike the usual procedure using transmitted light microscope photographs, to analyse and digitize drawings generated using Visopan Reichert at a magnification of ×201. The results of Morphomatica were processed using Cluster analysis, non-metric multidimensional scaling (n-MDS) and analysis of similarities (One-way ANOSIM Pairwise Test) by means of the program PRIMER 6 (Clarke & Gorley 2006).

Results

Celation, variability and specific assignment of Urocythereis sp. 1

The morphological variations in Urocythereis sp. 1 appear limited when compared with the U. margaritifera-distinguenda group. Celation is rare and the fossae are never hidden by secondary calcification; the structure of reticulation is somewhat constant. The basic pattern shows a close resemblance to the Pliocene form described by Uliczny (1969) as Urocythereis favosa exedata and elevated to the rank of species by Mostafawi & Matzke-Karasz (2006). Some stable shell features of U. sp. 1 are different from those of the Pliocene taxon, leading us to describe a new species, named U. ilariae sp. nov. as discussed in the systematic section.

Fossae shape and size variations and celation in the U. margaritifera-distinguenda group

Brady (1866) described a species from Recent sponge-sands of the Eastern Mediterranean as Cythere oblonga, characterized by “somewhat distant, oblong, pittings”. The name C. oblonga was, however, preoccupied by C. oblonga M’Coy, 1844; therefore, in 1928 Neviani assigned a new name, Cythereis distinguenda, recognized by Athersuch (1977, 1982) as the “next available name for Brady’s species”.

In previous investigations, the present authors used the size of the fossae to discriminate between *U. margaritifera* and *U. distinguenda*, being relatively large in the former species and very small in the latter one. The finding of specimens exhibiting transitional features makes their specific attribution difficult. The following morphs can be defined on the basis of the size and shape of the fossae:

Morph a (commonly assigned to *U. margaritifera*): medium-sized fossae, equidimensional, except in the anterodorsal zone where they are medium- to small-sized; generally rounded, elongated or fused in the anteroventral area (e.g., Fig. 16A, H);

Morph b (commonly assigned to *U. margaritifera*): medium-sized fossae rounded and equidimensional on the whole surface of the shell (e.g., Figs 2A; 16B);

Morph c (“transitional” form): mesh size from medium to very small, generally rounded (e.g., Fig. 17E);

Morph d (“transitional” form): fossae mainly elongated, from small to medium-sized (e.g., Fig. 17B);

Morph e (“transitional” form): the anterior area shows fossae very small or completely celled; the remaining fossae are small and rounded (e.g., Fig. 17D);

Morph f (commonly assigned to *U. distinguenda*): small-sized fossae; narrow and elongated in the anterior part of the valve, rounded in the posterior part and mixed in the central area (e.g., Fig. 16D);

Morph g (commonly assigned to *U. distinguenda*): small-sized fossae, rounded or elongated in the posterior half of the shell; very small (reduced into puncta) to completely celled in the anterior half (Fig. 17C);

Morph h (commonly assigned to *U. distinguenda*): size of fossae ranging from small to very small; elongated or rounded (Fig. 16G).

It has to be noted that in this latter group the muri are generally densely or sparsely papillate; in specimens showing intense celeration, papillae occur on the tegmen and are very sparse or lacking in correspondence with the underlying muri, revealing the reticulation pattern (Fig. 7). A similar feature was observed by Benson (1972) in the species *Poseidonamicus nudus* Benson, 1972.

In our opinion, these data suggest that *U. margaritifera* and *U. distinguenda* are morphotypes of the same species.

**Reticulation variability**

The fossal groups show the following variations:

**a. Pre-ocular fossa (pre-ocular bridge)**

In *U. margaritifera/distinguenda*, a murus located in the antero-dorsal corner, in pre-ocular position, generally connects the anterior marginal rim and the ocular riblet, which delimitate the marginal furrow (Figs 3A, C–E; 19D), forming a pre-ocular fossa. The pre-ocular bridge is more or less well defined in *U. margaritifera/distinguenda*, including celled specimens, where it is highly developed and the pre-ocular fossa is reduced (Fig. 3B). In some specimens of *U. margaritifera/distinguenda*, the bridge is absent and the furrow is continuous up to the eye tubercle (Fig. 3F).

The pre-ocular fossa never occurs in *U. ilariae* sp. nov. (Fig. 3G–H).

**b. Anterodorsal group. Fossal pattern B1–B4 – C1–C4**

In *U. ilariae* sp. nov., the presence of a murus between the fossae B5 and B4, connecting the second and the third marginal riblets, is a stable feature and a possible landmark when comparing species
of *Urocythereis*. It is subdued in some very rare ciliated specimens (Fig. 18G). A composite fossa, including the anastomized fossae B1, B2, B3, B4, C1 and C2, shows slight intraspecific variability (Figs 2B; 3G–H); in some specimens (Fig. 18C, F) B4 and C4 are connected.

In *U. margaritifera/distinguenda*, B and C fossae never connect. The fossae B1 and B2 may be divided (Fig. 17G), partially fused (Figs 2; 16B) or merged (Fig. 3E–F). Fossae B2-B3-B4 are never joined, C1 and C2 are usually divided (Fig. 3C, F) and can show some degrees of anastomosis (Fig. 3D–E). In some specimens ciliation may simulate mesh subdivision. For example, in Fig. 3B the C2 mesh is almost completely closed by tegmen and small foveolae mirror the underlying fossa. Fossae C3 and C4 are never fused.

c. D3-D4-E9 group

In *U. margaritifera/distinguenda*, D4 and E9 are usually distinct (Fig. 4B–F), rarely joined together (Fig. 4A). In a few specimens, D4 is partly divided, and D4 and E9 appear as an elongated subhorizontal fossa (Fig. 4A). D3 is constantly distinct.

---

**Fig. 7.** Detail of papillae in heavily ciliated specimen of *U. margaritifera* (G.W. Müller, 1894), ABMC 2014/071.
In *U. ilariae* sp. nov., the fossae D3-D4-E9 form a steady triangular pattern with D4-E9 fused and D3 is situated above them. In very rare celled specimens, D4 and E9 are distinct (Fig. 18G).

d. Dorsal median group. Fossal pattern α1–α6, β1–β5

In *U. margaritifera/distinguenda*, an ideal scheme of this group can be represented as two rows: the anterior row consists of six fossae, the posterior row includes five fossae, named α1–α6 and β1–β5 (Fig. 5A–F), respectively. This is seemingly in contrast with the scheme of Athersuch with 5+4 fossae; in fact, the upper two fossae (α1 and β1), just below the post-ocular sinus, may be clearly observed in dorsal view (Fig. 19D). In lateral view they appear very reduced or even completely hidden (Figs 5E–F; 16H; 17B). In the posterior row, between the 4th (β4) and 5th (β5) fossa, a small smooth area, the “focus” of Athersuch (1977), is present. The lowermost fossae, herein named α6 and β5, pertain, following Liebau's scheme (1971), to the E-ring encircling the muscle-scar node.

We observed the following variations:
- fossa α3 divided in two subfossae (Fig. 16F);
- fossa α5 divided in two (Fig. 16F, H) or three (Fig. 17F) subfossae;
- fossa α6 divided in two (Figs 5F; 17G);
- fossa β3 partially connected with the centrodorsal fossae (Figs 16C; 17A).

The shape of the meshes shows a wide variability, partly due to cellation, ranging from large rounded fossae to small pits, including narrow foveolae (Fig. 5A–F).

In *U. ilariae* sp. nov., the dorsal median group shows a different pattern (Fig. 5G–H). In comparison with the ideal scheme, the uppermost fossae (α1, β1) are absorbed in the post-ocular sinus; α2 and β2 are located under the post-ocular sinus; α2 is alternatively isolated (Figs 5G; 19B) or fused with β2 (Fig. 18B). The fossa β2, in turn, is fused with a posterior fossa (Figs 5G; 19B). The α3-β3/α4-β4 meshes may coalesce in part, generally forming a butterfly shaped quadruple fossa (Fig. 5G); in some specimens α4 is fused with an anterior fossa (Fig. 5H).

e. Caudal group

In *U. margaritifera/distinguenda*, it consists of a basic pattern of three well-defined fossae, two marginal and one internal with various shapes in different specimens (Fig. 6A–F). They may appear as a single triple fossa (Fig. 17H) or as a single plus one double fossa (Fig. 2A). In the latter case, anastomosis partially hides the identity of the fossae. Cellation may completely obscure this pattern (Figs 6B; 17C).

In *U. ilariae* sp. nov., a fourth upper fossa (Cg4) merges with the caudal group, forming a multiple L-shaped or elongated fossa; it can be more or less fused with another internal elongated multiple fossa (Fig. 6G–H).

Our observations indicate that the reticulation pattern in the forms previously attributed to *U. margaritifera* and *U. distinguenda* are identical. In *U. ilariae* sp. nov., well distinct patterns of homologous fossae can be recognized.

Outline analysis

The valve outline of *U. margaritifera* (i.e., *U. margaritifera* “morph margaritifera” plus *U. margaritifera* “transitional morph”), *U. distinguenda* (i.e., *U. margaritifera* “morph distinguenda”) and *U. ilariae* sp. nov. were analysed and compared. In Fig. 8A–C the “non normalized area” (size dependent) mode results show that the variability of *U. ilariae* sp. nov. (Fig. 8A) is very limited. By contrast, in *U. margaritifera* (Fig. 8B) the variability is relatively wide, linked to the length of the valves. Conversely, with the “normalized area” (size independent) mode, a low variability is observed in *U. margaritifera* (Fig. 8C).
9B) while it is relatively high in *U. ilariae* sp. nov. (Fig. 9A). *U. distinguenda* shows an intermediate variability in both modes (Figs 8C; 9C).

Comparisons of the mean outline of the three forms indicate that *U. ilariae* sp. nov. is positively separated from *U. margaritifera* and *U. distinguenda*, while the discrimination of the latter forms is difficult. In “normalized area” mode (Fig. 10B) *U. ilariae* sp. nov. shows a shorter length and a different shape in the caudal and in the anterodorsal regions. Its shorter length is more evident in the mean outlines deriving from “non normalized area” mode analysis (Fig. 10A).

**Fig. 8.** Comparison of valve outlines obtained by Morphomatica analysis. Results for “non-normalized area” mode. A. *U. ilariae* sp. nov. B. *U. margaritifera* (G.W. Müller, 1894). C. *U. distinguenda* (Neviani, 1928).

**Fig. 9.** Comparison of valve outlines obtained by Morphomatica analysis. Results for “normalized area” mode. A. *U. ilariae* sp. nov. B. *U. margaritifera* (G.W. Müller, 1894). C. *U. distinguenda* (Neviani, 1928).

**Fig. 10.** Comparison of mean outlines of *U. ilariae* sp. nov. (green), *U. margaritifera* (G.W. Müller, 1894) (light blue) and *U. distinguenda* (Neviani, 1928) (dark blue) obtained by the Morphomatica analysis. A. Results for “non-normalized area” mode. B. Results for “normalized area” mode.
Fig. 11. Cluster analysis performed on valve outlines obtained by Morphomatica analysis. 

A. Dendrogram for “non-normalized area” mode.

B. Dendrogram for “normalized area” mode.
Cluster analysis for “normalized area” mode reveals two main clusters, the first (A1) includes three valves of *U. ilariae* sp. nov., the second (A2) consists of all the remaining specimens (Fig. 11B). This second cluster is subdivided into two subclusters: A2a, including nine valves pertaining to *U. distinguenda* and three specimens of *U. margaritifera* and A2b, where all the three forms are present (eight *U. ilariae* sp. nov., two *U. distinguenda* and six *U. margaritifera* valves).

The “non normalized area” mode analysis (Fig. 11A) discriminates two main clusters: B1, consisting of three valves of *U. margaritifera* and four of *U. distinguenda*, and B2 including the remaining specimens.

The Anosim Pairwise Test applied both to the “non normalized area” (Fig. 12A) and “normalized area” mode (Fig. 12B) displays that *U. ilariae* sp. nov. and *U. distinguenda* are separated on the basis of the outline analysis (R = 0.603; R = 0.628). Conversely, *U. margaritifera* cannot be discriminated from *U. ilariae* sp. nov. or from *U. distinguenda*.

N-Mds analysis provided comparable results (Fig. 13). The distribution areas of *U. ilariae* sp. nov. and *U. distinguenda* are well defined for both “non normalized area” mode (Fig. 13A) and “normalized area” mode (Fig. 13B), whereas the distribution of *U. margaritifera* specimens overlaps both *U. ilariae* sp. nov. and *U. distinguenda*.

![Table](image)

<table>
<thead>
<tr>
<th></th>
<th>R</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>U. ilariae</em> vs <em>U. margaritifera</em></td>
<td>0.263</td>
<td>0.006</td>
</tr>
<tr>
<td><em>U. ilariae</em> vs <em>U. distinguenda</em></td>
<td>0.603</td>
<td>0.001</td>
</tr>
<tr>
<td><em>U. margaritifera</em> vs <em>U. distinguenda</em></td>
<td>0.070</td>
<td>0.165</td>
</tr>
</tbody>
</table>

**Fig. 12.** Analysis of similarities (One-way Anosim Pairwise Test). A. R statistic and p value for “non-normalized area” mode. B. R statistic and p value for “normalized area” mode.
Fig. 13. Non-metric multidimensional scaling (n-MDS) analysis. A. Results for “non-normalized area” mode. B. Results for “normalized area” mode.
**Taxonomy**

Classification follows Liebau (2005) and, for the attribution to subfamily, Hartmann & Puri (1974).

- **Subclass Ostracoda Latreille, 1802**
- **Superorder Podocopomorpha Kozur, 1972**
- **Order Podocopida Sars, 1866**
- **Suborder Cytherocopina Gründel, 1967**
- **Infraorder Nomocytherinina Liebau, 1991**
- **Superfamily Trachyleberidoidea Sylvester-Bradley, 1948**
- **Family Hemicytheridae Puri, 1953**
- **Subfamily Urocythereidinae Hartmann & Puri, 1974**

**Genus Urocythereis** Ruggieri, 1950

**Urocythereis ilariae** sp. nov.

urn:lsid:zoobank.org:act:607A810F-773C-4F8F-91e7-D2eB5223CB7D


**Urocythereis favosa** (Roemer) subsp. – Wouters 1973: 385, pl. 2, fig. 7.

**Urocythereis** sp. – Bonaduce, Ciampo & Masoli 1976: 46, pl. 22, fig. 9.

? **Urocythereis** sp. – Athersuch 1977: pl. 17, fig. 2.

**Urocythereis aff. U. favosa** (Roemer) – Arbulla, Pugliese & Russo 2001: fig. 3t.

? **Urocythereis** sp.l – Barra 1997: 82, pl. 4, fig. 6.

**Urocythereis** sp.l – Aiello et al. 2006: tables 3, 10. — Aiello, Barra & Parisi 2013: fig. 1d.

**Diagnosis**

A large reticulate species of *Urocythereis*, subrectangular in lateral view, inflated-ovate in dorsal view.

Reticulum with large polygonal-rounded, frequently coalescing, large fossae separated by broad muri.

In the anteroventral area the muri form distinct riblets running parallel to the margin.

**Etymology**

In honour of our friend and colleague Ilaria Mazzini, in recognition of her important contribution to ostracodology.

**Type material** (4 carapaces, 43 valves: 29 adults and 14 juveniles)

**Holotype**

IONIAN SEA: ABMC 2014/03

**Paratypes**


**Stratum typicum**

Recent.
Locus typicus
La Strea Bay (Porto Cesareo Lagoon), Southern Italy, Ionian Sea, sampling station E4, 17°54′25″ N, 40°15′59″ E, depth 1.5 m bsl.

Description
Measurements (holotype): LV: L = 0.85 mm, H = 0.44 mm (Fig. 18A).

Large (L = 0.85–0.90 mm) species of *Urocythereis*, characterized by large fossae and strongly developed muri, subrectangular in lateral view, inflated-ovate in dorsal view. Valves strongly calcified and thick. Dorsal margin gently, unevenly convex, ventral margin weakly sinuous; anterior end broadly rounded, denticulate in the lower part; upper part of the posterior margin concave, lower part of the posterior margin convex, variably denticulate, forming short blunt caudal process located below mid-height. Maximum height at anterior cardinal angle, greatest length below mid-height. Surface of valves coarsely reticulate. Fossae, showing subrounded or irregular shape, coalesce, especially in marginal areas, forming both multiple anastomized elongated fossae and deep sulci parallel to margin. The corresponding muri tend to form a system of concentric riblets. Marginal rim starts from anterior part of dorsal margin, behind eye tubercle (Fig. 19C), and ends in posteroventral angle. Second riblet, constantly well developed, runs parallel to margin of valve except posterior end. This ocular riblet is connected with eye tubercle and rises above dorsal margin. Marginal rim and second riblet not connected. Third riblet, irregularly developed, delimits anteriorly the reticulum *stricto sensu* from subocular area to posterior part of ventral area and is connected with second riblet anteriorly, at mid height, through single radial murus; in the ventral area second and third riblets converge and, in lateral view, they seem apparently to be connected, but ventral view (Fig. 19A) shows they remain separate. The fossae between second and third riblet mainly anastomized. Fourth riblet fully part of reticulum, and shows a rather regular parallel trend only in anterocentral area. Surface of central area irregularly reticulate with subrounded/polygonal fossae with a low degree of anastomosis. Conversely, fossae located in proximity of caudal process coalesce following a longitudinal trend. Rare specimens show celation, never fully developed. Muri smooth, not papillate (Fig. 19E).

Hinge holamphidont (*sensu* Scott 1961): in left valve posterior hinge socket elongate and curved; anterior element formed by ovate-rounded (or elongate) tooth and elongate socket; median bar smooth; its posterior thickening forms, in some cases, barely defined toothlet; right valve hinge complementary, with faintly crenulate teeth (Figs 18I–J; 19G–J).

Inner lamella, marginal pore canals and muscle scar pattern (Fig. 19F) characteristic of genus (details in Athersuch 1977).

Distribution
The species occurs in the Recent of the Mediterranean: Gulf of Naples (Bassiouni 1965), Sardinia (Arbulla *et al.* 2001), South Adriatic Sea (Bonaduce *et al.* 1976) and possibly Libya (see section Remarks); it has previously been recorded in fossil associations from the Tyrrenian (upper Pleistocene) of Tunisia only (Wouters 1973). Distribution data are summarized in Fig. 14.

Remarks
*U. ilariae* sp. nov. has previously been assigned to *U. favosa* (Bassiouni 1965; Wouters 1973), type species of the genus *Urocythereis* (neotype figured by Athersuch 1977). The reticulation of *U. favosa* differs from that of *U. ilariae* sp. nov. in the different style of fossal anastomosis. This is mostly evident, for example, in the anterodorsal zone, where the continuous depressed area formed by the fossal pattern C1–C2/B1–B4 is present in *U. ilariae* sp. nov. and absent in the Pliocene species.
The shell characters of *U. exedata*, described by Uliczny (1969) as a subspecies of *U. favosa* (SEM micrographs in Mostafawi & Matzke-Karasz 2006), show a close resemblance to those of *U. ilariae* sp. nov., especially in the structure of the ocular riblet, homologous to *Bradleya*’s “ocular ridge” (Benson 1972). The Pliocene species probably represents an ancestor of the living form. The two species differ in some reticulum features. In the anteroventral area of *U. ilariae* sp. nov., the third and the fourth riblets are connected ventrally and anteriorly; consequently they delimit the merged C fossae, forming an anteroventral furrow enclosed by muri. Conversely, in *U. exedata* the anteroventral area is characterized

![Distribution of *U. ilariae* sp. nov. and *U. margaritifera* (G.W. Müller, 1894).](image)

**Fig. 14.** Distribution of *U. ilariae* sp. nov. and *U. margaritifera* (G.W. Müller, 1894). Circles = Recent; triangles = Pleistocene; squares = Pliocene; filled = certain; empty = uncertain.
by a segment of the third riblet encircled by an elongated ring made up of B and C anastomized fossae, anteriorly and ventrally connected. In the anterodorsal area of *U. ilariae* sp. nov., the third concentric riblet is more or less developed in different specimens (Figs 2B; 18D, F), while in *U. exedata* in the anterodorsal area the fossae of the B group coalesce with C and D fossae, the muri follow a radial trend and consequently the third concentric riblet is virtually absent.

The assignment of the North-African form, figured by Athersuch (1977) as *Urocythereis* sp. and by Barra (1997) as *Urocythereis* sp. 1, to *U. ilariae* sp. nov. needs further investigation. At the current state of knowledge we are inclined to interpret the morphological differences between the central Mediterranean species and the Lybian deme as the beginning of an allopatric speciation.
Distribution

The species is widely distributed in the infralittoral waters of the Eastern Mediterranean (Brady 1866; Barbeito-Gonzalez 1971; Doruk 1974; Athersuch 1977, 1979; Kubanç 1995; Tunoglu 1999; Perçin-Paçal & Balkis 2012), the Tyrrenian Sea (G.W. Müller 1894; Puri 1974) and the southern Mediterranean (Athersuch 1977; Lachenal 1989; Barra 1997). Recordings from the Black Sea are uncertain: the specimen figured by Stancheva (1989) is a young instar, and Schornikov (1969) reported Müller’s original drawings. The species is present in the southern part of the Adriatic Sea; the findings in the central and northern Adriatic are doubtful (Uffenorde 1972; Bonaduce et al. 1976; Breman 1976).

Fossil specimens have been reported from the Upper Pleistocene–Holocene of the Gulf of Gabès (Lachenal 1989), the Pleistocene of Southern Italy (Ruggieri 1953; Aiello et al. 2012), Zakynthos (Tsapralis 1981), the Northern Peloponnesus (Mostafawi 1994) and, possibly, Rhodes (Sissingh 1972) and from the Pliocene of Cephalonia (Uliczny 1969). Distribution data are summarized in Fig. 14.

The presence of the species in Miocene sediments (Şafak et al. 1999) has to be confirmed by further studies.

Remarks

The analysis of the shell features of the *Urocythereis* population in the La Strea Bay and comparisons with the literature have convinced us that *U. margaritifera* and *U. distinguenda* (= *U. oblonga*) are two morphotypes of the same species. In particular, we consider the latter “species” as the celated variation of the former. Celation is not expressed homogeneously on the valves in all the specimens; consequently, also “transitional” shells show different morphs.

The original illustration by Müller (1894: pl. 32, fig. 26) shows anteroventral fossae horizontally merged; the lectotype reported by Athersuch (1977: pl. 13, fig. 2) and the Libyan specimen figured by Barra (1997, as *U. sp. 2*) shows the same feature. Presently, we do not regard this character as diagnostic, due to the observed variability.

---

**Fig. 15.** Scheme of the hidden reticulation evidenced in a celated specimen, LV, sample E6, ABCM 2014/053. Same specimen as Fig. 16G.
Athersuch (1977) figured some appendages, including the right copulatory appendage, of *U. distinguenda* and the left copulatory appendage of *U. margaritifera*. Regarding the discrimination of the two species, the author stated that in *U. margaritifera* the ductus ejaculatorius is short and is contained within the area of the appendage, whereas in *U. distinguenda* the duct is much longer and passes beyond the ventral margin. Examination of Athersuch’s illustrations (figs 4.d; 5.i) and a comparison with Müller’s drawing of the male copulatory appendage of *Cythereis margaritifera* (pl. 32, fig. 32) show that only very subtle differences are present and they represent, in our opinion, intraspecific variations.

The maximum mesh size is reached in the subspecies *Urocythereis margaritifera alba* Uliczny, 1969. This form, in which celation is not developed, does not occur in the La Strea Bay. In some specimens the SEM micrographs revealed a feeble trace of the muri underlying the secondary calcification, as shown in Fig. 15. The comparison of this hidden reticulation with the specimens figured by Uliczny (1969), Breman (1976) and Aiello et al. (2012) suggests that *U. m. alba* is a non-celate morphotype of *U. margaritifera*. The North Adriatic form figured by Uffenorde (1972) (very similar to the Pliocene specimen figured by Şafak et al. 1999 as *U. margaritifera*) shows some tiny differences in the reticulation pattern and the assignment to *U. margaritifera* is queried.

The Libyan form figured by Athersuch (1977: pl. 17, fig. 5) as *U. sp.* and by Barra (1997) as *U. sp.* 3 fits the *U. margaritifera* morph c (Fig. 17E) well.

The left valve, figured by Aranki (1987) from the western Mediterranean shallow waters, shows some minor differences in the reniform outline and in some details of the reticulum. The relationships between Mediterranean and Atlantic forms (i.e., between *U. margaritifera* and *U. britannica*, frequently reported as *U. oblonga*) need further investigations.

Ruggieri (1953) hypothesized that *U. favosa* and *U. margaritifera* might be conspecific, the latter species representing a subspecies of the former. In spite of the similarity of the two forms, some features of the reticulum seem to allow a separation of the two species. In *U. favosa* (neotype figured in Athersuch 1977) the fossae B3 and B4 are merged, as well as C3 and C4; in *U. margaritifera* they are distinct; in *U. favosa* the fossae D2-D1 and α4 coalesce while in *U. margaritifera* they are distinct. In the caudal group the fossa Cg4 in *U. margaritifera* is separate, whereas in *U. favosa* the arrangement of the fossae is similar to that in *U. iliares* sp. nov. In the La Strea Bay, and possibly in the Recent of the Mediterranean, *U. favosa* s.s. is not recorded and we prefer to retain them as separate species.

Some *Urocythereis* spp. from the Pliocene of Rhodes have been described by Terquem (1878) and figured by Mostafawi (1989). They are distinct from *U. margaritifera* in some characters of the reticulum. By contrast, the specimen from Cephalonia figured by Mostafawi & Matzke-Karasz (2006) as *U. crenulosa* fits well with some specimens of *U. margaritifera* from Porto Cesareo.

The Atlantic forms reported as *U. britannica* and (erroneously) as *U. oblonga* (e.g., Guillaume et al. 1985; Ruiz et al. 2006) show a high variability and a complex affinity with *U. margaritifera* and *U. favosa* and they have not been considered in the present study.

**Discussion**

The comparison of five selected fossae-muri homologous groups convinced us that at present two species of *Urocythereis* live in the Porto Cesareo Lagoon and not three as previously reported by Aiello et al. (2006, 2013). Reticulum analysis demonstrates that the ornamentation patterns of *U. margaritifera* and *U. distinguenda* (sensu Aiello et al. 2006) are basically identical, and that the differences between morphs are similar, for example, to the continuous (non-polymorphic s.s.) variations described by Neil (2000) for the hemicytherid species *Chapmanella flexicostata* (Chapman, 1914).
Our observations confirm only partly Athersuch’s statement that celation is present exclusively in adult individuals of *Urocythereis*. Actually, we have recorded a few penultimate instars showing a certain degree of celation (Fig. 16I, K). In young A-2 instars, celation is not recorded or is very poorly developed (Fig. 16J).

The results of the morphometric analysis by means of the Morphomatica program need further interpretation.

The **ANOSIM** Pairwise Test and n-Mds analysis confirm that the outlines of *U. margaritifera* and *U. distinguenda* cannot be discriminated, as showed by the reticulum analysis, supporting the idea that they are different morphs of the same species. Furthermore, the **ANOSIM** Pairwise Test and n-Mds analysis display a clear separation between *U. ilariae* sp. nov. and *U. margaritifera “morph distinguenda”*, while they are not able to discriminate between *U. ilariae* sp. nov. and *U. margaritifera “morph margaritifera”*, in spite of the evidence that they show different specific reticulation features, due to the wide outline variability of the latter form.

By contrast, the comparison of the mean outlines derived from the “non normalized area” mode allows the discrimination between *U. ilariae* sp. nov. and *U. margaritifera s.l.*

We interpret the peculiar distribution of *U. ilariae* sp. nov. and celled *U. margaritifera (“morph distinguenda”) in two different ways. The first is to separate them, due to the obvious attribution to a different species showing a relatively low variability in outline and reticulation. On the other hand, the distribution of *U. margaritifera “morph distinguenda”* needs a different explanation. We suggest that the outlines of celled specimens show a certain degree of separation due to the secondary calcification of the valves. According to the Van Morkhoven’s investigations (1962), in specimens of *Cypris pubera* a rich calcium diet induced secondary thickening of the shell. The morphological alterations developed to such a point that, in the author’s words, “most likely they would have been classified as a distinct species, if the history of the individuals had not been known”.

In some *Urocythereis* species, the age effect could somewhat replace the food-related secondary calcification observed by Van Morkhoven.

Our favourite solution to the “*U. margaritifera - U. distinguenda* problem” is to consider the *distinguenda* morph as the result of a supposed gerontic process, possibly in waters supersaturated with CaCO$_3$ (see Peypouquet *et al.* 1988, and references therein), which caused both celation and a slight outline modification, partly recognized by Morphomatica.

The presence of celation in juveniles seems to contradict this interpretation, nonetheless it has been considered that the presence of long A-2 and A-1 stages duration has been recorded, as for example by Hagerman (1969) in *Hirschmannia viridis* (O.F. Müller, 1785). We suggest that overwintering juveniles (Horne 1983 and references therein) could in some cases develop the same gerontic features as the adults.

**Conclusions and additional remarks**

In the present investigation we attempted to examine the variability of the *Urocythereis* shell characters of the species occurring in the La Strea Bay. Our results show that we deal here with two species: *U. ilariae* sp. nov., displaying a low variability and a low degree of celation, and *U. margaritifera*, a species with a high (continuous, non polymorphic) variability and able to express intense celation. The specific name *U. distinguenda* has previously been assigned, in our opinion, to celled specimens of *U. margaritifera (“aggrading” morph, *sensu* Peypouquet *et al.* 1988), and it is not even a valid subspecies. Furthermore, we have considered *U. margaritifera alba* as a “fully non-celated” (“degrading” *sensu* Peypouquet *et
al. 1988) morph of *U. margaritifera*. In the La Strea Bay this form is absent, probably due to the persisting supersaturation of calcium carbonate, as demonstrated by the high percentage of miliolids in taphocoenoses of the bay, ranging from 12.16% to 35.08% (Aiello et al. 2006). Actually, miliolids are benthic foraminifera preferring waters rich in calcium carbonate (e.g., Jorissen 1988) and very rare in low pH environments (e.g., Aiello et al. 2012 and references therein). In such a calcium super-saturated environment, all the adult specimens, and rare penultimate instars, show at least a moderate celation. Since the La Strea Bay is populated by a subtropical biocoenosis, our observations are in agreement with the statement of Peypoqu et al. (1988), who pointed out the relationship between aggradation and high water temperature.

In *U. ilariae* sp. nov., the less developed celation highlights different behaviour in *Urocythereis* species. Consequently, we suggest that three main factors influence the tegmen development: a genetically determined (specific) ability to express celation, the saturation in CaCO₃ of the bottom waters and the age of the organism (adult or penultimate instar). *Urocythereis margaritifera, U. m. alba and U. distinguenda* are morphotypes derived by environmentally cued variability.

The definition of the variability in a single population may provide, in our opinion, a step toward a broader understanding of the variability within *Urocythereis* through time and space.

Morphological variations of shallow marine ostracod species living in different areas (e.g., Ishizaki 1975; Hartmann 1982) may be interpreted as the beginning of an allopatric speciation, and future investigations of the *U. marginitifera / U. britannica* group could discriminate these species or, conversely, demonstrate that they are subspecies or morphotypes. A more complex problem is the definition of the relationship of the Miocene Atlantic *Urocythereis* pertaining to the *U. favosa* group (e.g., *U. favosa* in Nascimento 1988) that possibly recolonized the Mediterranean at the end of the Salinity Crisis with the Pliocene–Recent forms. An estimation of the total number of *Urocythereis* species in the Mediterranean Neogene and their phylogenetic relationships represent open questions.

**Acknowledgment**

The authors thank Alexander Liebau for some useful suggestions in the very beginning of this investigation, Dave Horne and an anonymous reviewer for their helpful comments and Roberto de’ Gennaro (Cisag, Università di Napoli Federico II) who took the SEM micrographs.

**References**


Athersuch J. 1979. The ecology and distribution of the littoral ostracods of Cyprus. *Journal of Natural History* 13: 135–160. [http://dx.doi.org/10.1080/00222937900770111a](http://dx.doi.org/10.1080/00222937900770111a)


---

*Manuscript received: 13 October 2015*

*Manuscript accepted: 14 December 2015*

*Published on: 3 May 2016*

*Topic editor: Rudy Jocqué*

*Desk editor: Kristiaan Hoedemakers*

---

Printed versions of all papers are also deposited in the libraries of the institutes that are members of the EJT consortium: Muséum national d’Histoire naturelle, Paris, France; Botanic Garden Meise, Belgium; Royal Museum for Central Africa, Tervuren, Belgium; Natural History Museum, London, United Kingdom; Royal Belgian Institute of Natural Sciences, Brussels, Belgium; Natural History Museum of Denmark, Copenhagen, Denmark.