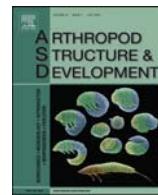




Contents lists available at ScienceDirect



Arthropod Structure & Development

journal homepage: www.elsevier.com/locate/asd

The evolution of a key character, or how to evolve a slipper lobster

Joachim T. Haug ^{a,*}, Denis Audo ^b, Sylvain Charbonnier ^c, Ferran Palero ^d, Gilles Petit ^c, Pierre Abi Saad ^e, Carolin Haug ^a^a LMU Munich, Department of Biology II and GeoBio-Center, Großhaderner Str. 2, 82152 Martinsried-Planegg, Germany^b Université de Rennes, EA 7316, 263 Avenue du General Leclerc CS 74205, 35042 Rennes Cedex, France^c Muséum National d'Histoire Naturelle, Paris UMR 7207 CR2P, CNRS-MNHN-UPMC, case postale 38, 57 rue Cuvier, 75231 Paris Cedex 05, France^d Institut Sophia Agrobiotech (INRA), 400 route des Chappes, BP 167, 06903 Sophia Antipolis Cedex, France^e Memory of Time, Citadel Area, Byblos, Lebanon

ARTICLE INFO

Article history:

Received 28 April 2015

Accepted 15 August 2015

Available online xxx

Keywords:

Achelata

Character evolution

Scyllarida s. str.

Verscyllarida

Autofluorescence imaging

Limestones

ABSTRACT

A new fossil lobster from the Cretaceous of Lebanon, *Charbelicaris maronites* gen. et sp. nov., is presented here, while the former species '*Cancrinus*' *libanensis* is re-described as *Paracancrinus libanensis* comb. nov. *P. libanensis* is shown to be closer related to the contemporary slipper lobsters than to *Cancrinus claviger* (lithographic limestones, Jurassic, southern Germany). A finely-graded evolutionary scenario for the slipper-lobster morphotype is reconstructed based on these fossil species and extant forms. The evolutionary changes that gave rise to the current plate-like antennae of Scyllaridae, a key apomorphy of this group, are traced back through time. The antenna of what is considered the oldest slipper lobster became petaloid and consisted of about 20 fully articulated elements. For this group the name Scyllarida *sensu lato* tax. nov. is introduced. In a next evolutionary step, the proximal articles became conjoined and a lateral extension appeared on peduncle element 3. The entire distal petaloid region is conjoined already at the node of Verscyllarida tax. nov. In modern slipper lobsters, Neoscyllarida tax. nov., the distal region is no longer petaloid in shape but asymmetrical. The study also emphasizes that exceptionally preserved fossils need to be documented with optimal documentation techniques to obtain all available information.

© 2015 Elsevier Ltd. All rights reserved.

1. Introduction

Distinct morphotypes have been discussed repeatedly within Decapoda in recent years, with a focus on evolutionary scenarios leading to these morphotypes. A particularly 'hot' topic has been the process of 'carcinisation', or the evolution of the 'crab' morphotype, respectively, the evolution of a crab-like habitus (e.g. McLaughlin and Lemaitre, 1997; McLaughlin et al., 2004; Hiller et al., 2010; Tsang et al., 2011; Scholtz, 2014). In evolutionary terms, any form of crab has evolved from an ancestral macruran or "lobster" morphotype. An interesting and much less studied case of departure from the simple "lobster" morphotype are the slipper lobsters, or scyllarids, easily differentiable from other decapod lobsters due to their unique morphology (e.g. Jones, 1990). The

group Scyllaridae forms a sharply characterised natural group, distinguished by wide and flat segments of the peduncle of the antennae (a2, second antennae) and by having the antennal flagellum transformed to a single broad and flat element without a trace of articulation. Scyllarid lobsters can be considered to represent a distinct 'morphotype' (note that the term used here is more or less equivalent what other authors term 'bauplan'; yet this has a quite difficult meaning in German), and this is especially apparent when comparing them to their closest relatives, the palinurid lobsters (e.g. Palero et al., 2009; Bracken-Grissom et al., 2014). Scyllaridae and Palinuridae form together Achelata (Scholtz and Richter, 1995), characterized by the lack of true claws (i.e. no index or fixed finger is opposed to the dactylus, or it is very short) and the presence of a phyllosoma larva. Contrary to the most familiar lobster types, slipper lobsters are dorsoventrally flattened, so that their cephalo-thoracic dorsal shield (also called carapace) is sometimes very wide, often square-shaped (in dorsal view) or even anteriorly widened. The dorsoventrally flattened body and, more

* Corresponding author.

E-mail addresses: joachim.haug@palaeo-evo-devo.info (J.T. Haug), denis.audo@univ-rennes1.fr (D. Audo), scharbonnier@mnhn.fr (S. Charbonnier), Ferran.Palero@sophia.inra.fr (F. Palero), carolin.haug@palaeo-evo-devo.info (C. Haug).

importantly, the distinctive antennae of scyllarid lobsters must have evolved in their own evolutionary lineage.

The structure of the highly specialized, shovel-like antennae of scyllarid lobsters is remarkable. According to Holthuis (1991) the basal element of the antennal peduncle is conjoined with the carapace; so it is only visible ventrally (treated as peduncle element 1 in the following). The second and third element are conjoined to a single element which is short and broad and may bear teeth on the anterior margin (treated as peduncle element 2). In most scyllarids this element is free. In *Scyllarus*, however, it is immovably connected with the antennular somite and the carapace, thereby forming an integral part of the orbit (Holthuis, 1985). The fourth element is large, broad and flat; it usually bears teeth on its margins (treated as peduncle element 3). The fifth segment is narrow and small; it may bear teeth on the inner margin (treated as peduncle element 4). The last element of the antenna, which represents the flagellum, is very broad and flat, sometimes with teeth on the margin. This element, together with the fourth one, is what makes the antenna into a broad, shovel-like organ for which the vernacular names 'shovel nose shrimp' and 'bulldozer lobster' are given to species of scyllarids. By providing "intermediate forms", fossils have the potential to provide further insights in the origin of key synapomorphies, such as the scyllarid antenna. Phylogenetically expressed, early fossil representatives of a group often have only part of the apomorphies of the modern group, while still lacking others. In this way the sequence in which the specializations of a morphotype evolved can be broken down into several, more gradual evolutionary transformations (examples in Senter, 2010; Haug et al., 2010).

The Upper Jurassic Solnhofen lithographic limestones (ca. 150 million years old, southern Germany) yielded the fossils of *Archaeopteryx lithographica*, an important early representative of birds still lacking some autapomorphies of the modern representatives, thus representing an example of a fossil providing an "evolutionary step in between". But the Solnhofen lithographic limestones also yielded an important early representative of slipper lobsters, which provides insight into the evolution of their specific morphotype. *Cancrios claviger* resembles spiny lobsters in general habitus. Yet, its antennae have a rather stout petaloid shape and comprise only about 20 elements. Förster (1973, 1984, 1985) therefore suggested that *C. claviger* represents an evolutionary step towards slipper lobsters. Haug et al. (2009a) could add additional insights into the evolutionary transition from the morphotype of spiny lobsters to *C. claviger*. Early juveniles of *C. claviger* still possess an elongate multi-annulated antenna and develop the petaloid shape gradually, by widening of the proximal elements and finally loss of the distal feeler-like part. Yet, the further evolutionary transition from the morphology of *C. claviger*, which mainly appears like a highly specialized spiny lobster, to the very distinct morphotype of slipper lobsters is still unclear.

We present new fossils from the Cretaceous of Lebanon that further close the gap of knowledge about the evolution of the morphotype of slipper lobsters. Based on these we reconstruct a possible scenario for the evolution of the slipper lobster morphotype.

2. Material and methods

2.1. Material

Most material is part of the palaeontological collections of the Muséum national d'Histoire naturelle, Paris, France (acronym: MNHN.F). The material was collected during a field campaign in 2011 (SC, DA, GP) in the Late Cretaceous Lagerstätten of Lebanon: Hadjoula and Hakel (Cenomanian, ~95 Ma) and Sahel Alma

(Santonian, ~85 Ma). Comparative specimens are housed in the palaeontological collections of the Museo civico di Storia naturale di Milano (acronym: MSNM).

Lebanese Lagerstätten are well-known for their exquisitely preserved crustacean faunas. For details see: Brocchi 1875; Fraas 1878; Dames 1886; Glaessner 1945; Roger 1946; Garassino 1994, 2001; Larghi 2004; Garassino and Schweigert 2006; Ahyong et al., 2007; Feldmann 2009; Garassino et al., 2009; Feldmann and Charbonnier 2011; Pasini and Garassino 2011; Petit and Charbonnier 2012; Charbonnier et al., 2013; Haug et al., 2013. For geological setting, see Ferry et al. (2007) and Audo and Charbonnier (2012, 2013).

2.2. Methods

Specimens from Paris have been documented using the macro-fluorescence settings described by Haug et al. (2011a) and Haug and Haug (2011). In most cases grey scale images of fluorescence images were produced by deleting blue and green channel, desaturating the red channel and optimizing the histogram. In some cases a better contrast was achieved by the following procedure: saturation dialog was opened, brightness of 'blue' and 'cyan' was set to '0', the brightness of 'red' was set to '100'. The resulting image was desaturated, and the histogram was optimized. Some specimens were additionally documented as stereo images (see e.g. Haug et al., 2011b, 2012a). Stereo images had their brightness values inverted to enhance the contrast.

Specimens from Milano were imaged by a Nikon D700 digital camera equipped with a Sigma 105 mm macro-lens with or without extension tube. Fossils display autofluorescence under UV light (yellow) and green light (orange) (see Haug et al., 2009b). Macro-fluorescence was therefore used to document their anatomy. Under cross-polarized light, spotlights and lens were equipped with linear polarizing filters arranged so that reflexions would be minimal. Under UV light, a portable UV light set to short wave UV (emission peak around 254 nm) was used. Under green–orange fluorescence, spotlights were equipped with medium blue–green gelatine filter and camera macro-lens equipped with a red filter to cancel all light but orange fluorescence. All raw image files were processed in Adobe Camera Raw and Gimp to adjust white balance and levels.

2.3. Description

The description of the here presented specimens is provided as a descriptive matrix (Haug et al., 2012b). This is thought to allow the taxonomically interested reader to access all necessary information, without centering this paper on this aspect.

2.4. Phylogenetic reconstruction

The present study focuses on the evolution of a series of characters of the antenna which resolve without character conflict. For this reason, there was no need to use computer-based phylogenetic analyses. This situation also has the advantage to avoid "hiding" apomorphies in a large matrix where they would be mixed with various characters, and the contribution of each character might end up difficult to assess. We will therefore concentrate on discussing the primary homology assumptions and the resulting evolutionary scenario.

3. Taxonomy

Eucrustacea *sensu* Walossek, 1993
Malacostraca Latreille, 1802
Decapoda Latreille, 1802

Reptantia Boas, 1880

Achelata Scholtz and Richter, 1995.

Scyllarida sensu lato tax. nov. (see discussion)

Scyllarida sensu stricto tax. nov. (see discussion)

Paracancrinos gen. nov.

Derivatio nominis: “*Cancrinos*” for the similarities to this genus and “para” for stressing the difference to it.

Type species: *Cancrinos libanensis* Garassino and Schweigert, 2006, by monotypy.

Diagnosis: as for the species.

Remarks: The erection of this new genus becomes necessary as the species *libanensis* Garassino and Schweigert, 2006 was originally ascribed to the genus *Cancrinos* Münster, 1839. Yet, it shares one key character with other scyllarids that *C. claviger* is lacking: the non-separation of the proximal elements of the petaloid region of the antenna (see below). In consequence *Cancrinos* would become paraphyletic. Therefore, the species *C. libanensis* Garassino and Schweigert, 2006 has to be excluded and (following Linnean taxonomy) demands for the erection of its own genus.

Paracancrinos libanensis (Garassino and Schweigert, 2006) comb. nov.

Type material: holotype MSNM i26597 (Figs. 1 and 2A–C), paratype MSNM i26593 (Figs. 1 and 2D,E).

Additional material: two specimens MNHN.F.A48952 (Figs. 1, 3A and 4A), A48973 (Fig. 5A,B).

Type locality: Hadjoula Lagerstätte, Lebanon (Late Cretaceous, Cenomanian).

Amended diagnosis: Small-sized achelatan with a carapace of rectangular shape (dorsal view); strong and stout petaloid flagellum of antenna; distally with fully articulated elements, proximal elements conjoined.

Description: Full description is provided as descriptive matrix in supplement.

Remarks: Our description differs in certain aspects from the original one. This is most likely facilitated by the additional material, but also by the application of macro-fluorescence imaging. Most important, the petaloid part of the antenna has fully jointed articles only distally, further proximally these are conjoined (Figs. 3B and 5D,E). Garassino and Schweigert (2006) described the species as lacking a rostrum. Our new material as well as re-investigation of the type material indicates the presence of a short rostrum (Fig. 5B). Furthermore the authors described the uropod exopod as lacking a diaeresis (subdivided by a joint). The morphology of the typical achelatan uropod makes this interpretation difficult, as the distal part of the uropod exopod is soft, while the proximal part is strongly sclerotized (Figs. 2E, 3A,C, and 5F). The decision comes down to the question whether this situation is homologous to a diaeresis or not. We leave this question open, but want to point to this difficulty. A further difference to the original description is the relation of lengths (carapace length vs. pleon length). Garassino and Schweigert (2006) stated that the “abdomen” (pleon) is as long as the carapace. Yet as the anterior rim of the carapace is difficult to assess (due to the strong antennal peduncles) this was most likely a measuring error. The length of the carapace equals about the length of four pleomeres. Furthermore, fluorescence light microscopy revealed some aspects of the tail fan that differ from the original description. The rim of the hard part of the telson is straight, not convex (Fig. 2E). Carinae on telson and

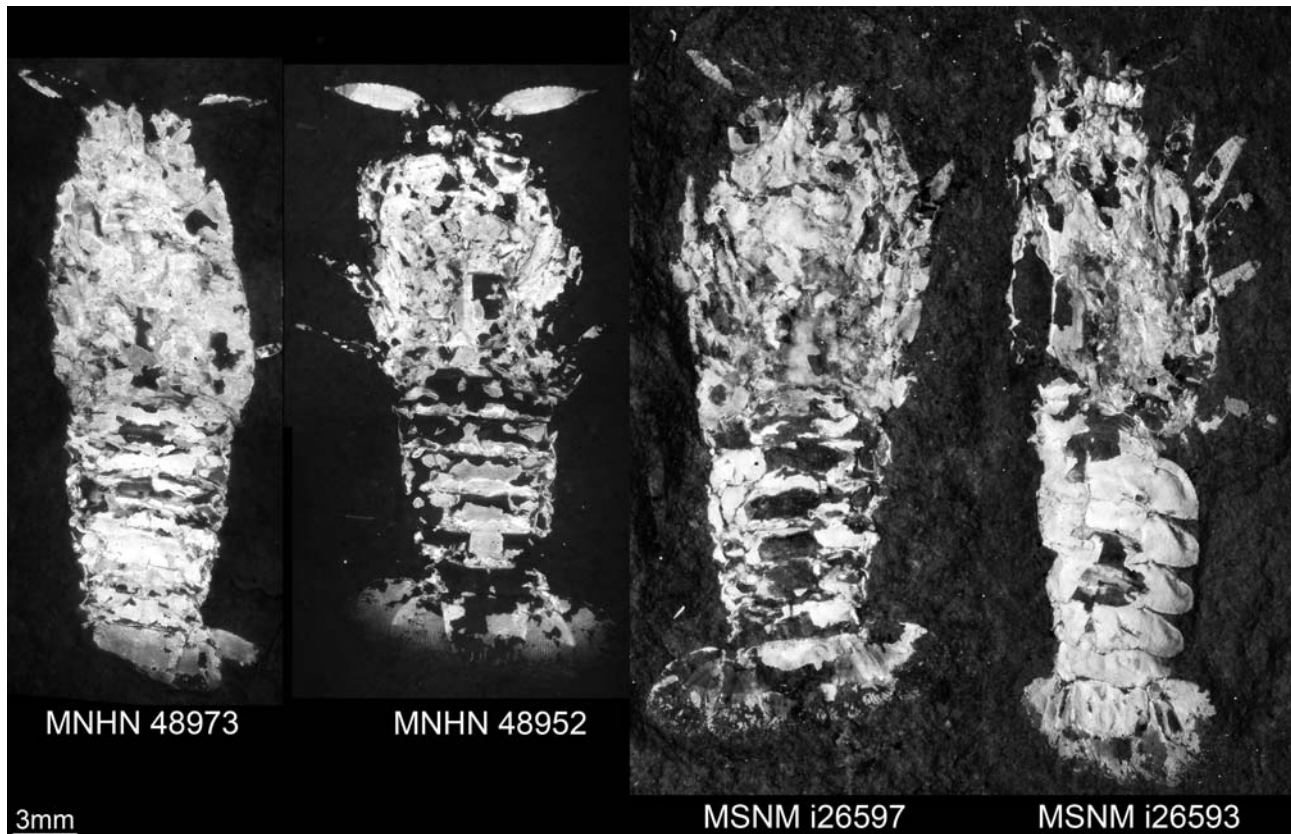


Fig. 1. Overview of the material of *Paracancrinos libanensis* (Garassino and Schweigert, 2006). New material, representing an earlier instar (MNHN.F.A48973, A48952). Type material, representing a later instar (holotype MSNM i26597, paratype MSNM i26593).

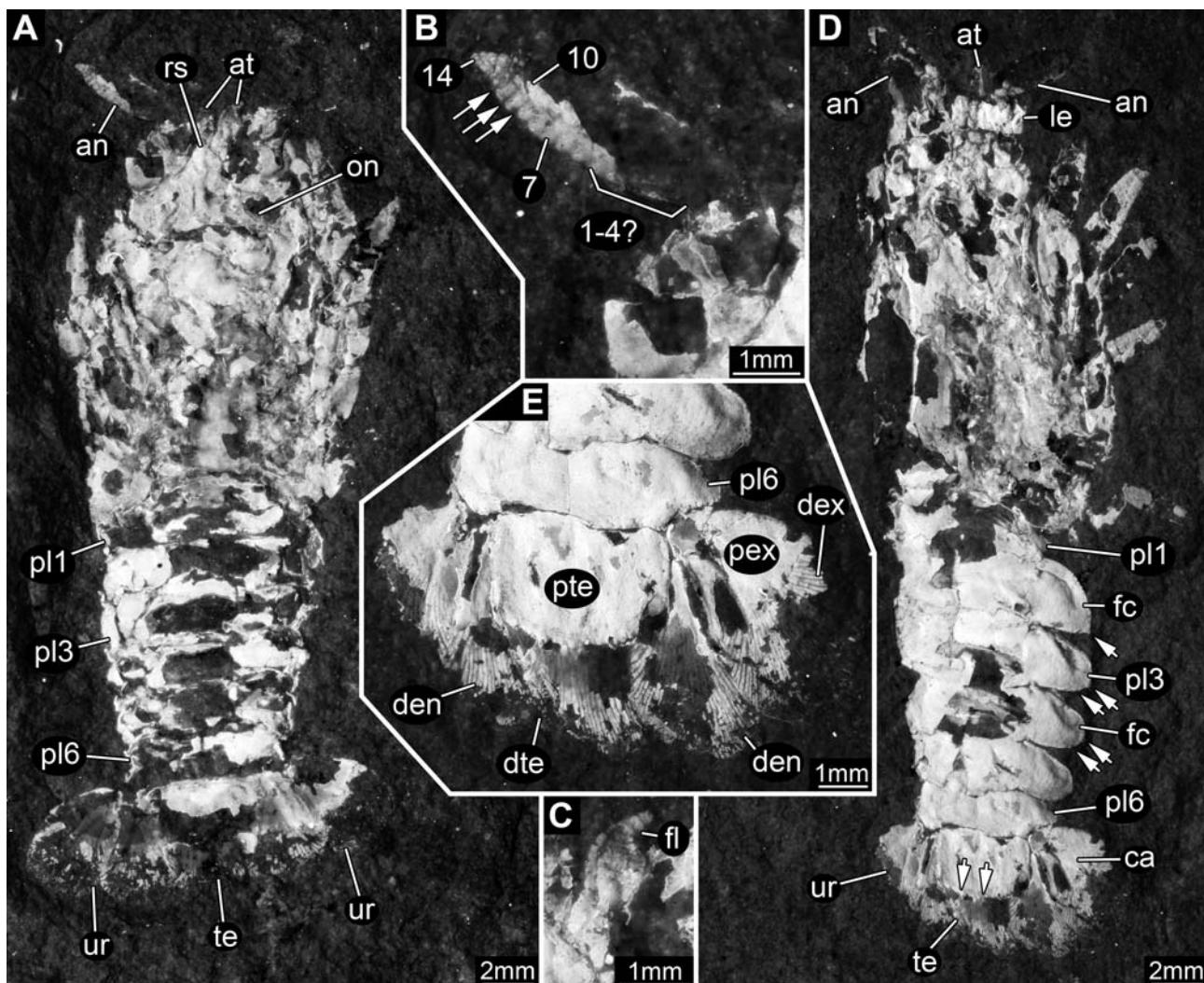


Fig. 2. *Paracancrius libanensis* (Garassino and Schweigert, 2006), later instar, type material. A–C. Holotype MSNM i26597. A. Overview. B. Close-up of antenna. Arrows mark borders between articles. C. Close-up on antennula. D, E. Paratype MSNM i26593 48939. D. Overview. Arrows mark spines. E. Close-up on tail fan. Abbreviations: 1–14 = antennal articles; an = antenna; at = antennula; ca = carina; den = distal uropod endopod; dex = distal uropod exopod; dte = distal part of telson; fl = flagellum; le = lateral extension; on = optical notch; pex = proximal uropod exopod; pl 1–6 = pleomeres; pte = proximal region of telson; rs = rostrum; te = telson; ur = uropod.

uropod exopod are not apparent; possibly these are compressions of uropod endopod and exopod rims. Last point, what Garassino and Schweigert (2006) described as “fringed” might have referred to the stiffeners of the softer part of endopod and exopod. A further point is the length and robustness of the thoracopods 4–8 (=pereiopods 1 to 5 in decapod crustaceans). We find these difficult to assess due to preservation, and neither support nor disprove whether thoracopod 4 (pereiopod 1) is shorter and stouter than the others (Fig. 4B).

Verschyllarida tax. nov. (see discussion)

Charbelicaris gen. nov.

Type species: *Charbelicaris maronites*.

Etymology: in honor of Youssef Antoun Makhlof also known as Saint Charbel, hermit of Mount Lebanon, near Hadjoula.

Diagnosis: as for the species.

Charbelicaris maronites sp. nov.

Etymology: from the Maronites, an ethnoreligious group in Levant who established the Maronite Church from Lebanon; used as a noun in apposition.

Type material: holotype MNHN.F.A48955 (Fig. 6A–C), paratype MNHN.F.A48984 (Fig. 6D–G).

Type locality: Hadjoula Lagerstätte, Lebanon (Late Cretaceous, Cenomanian).

Diagnosis: Small-sized achelatan with a carapace of rectangular shape (dorsal view); strong and stout petaloid flagellum of antenna; all elements conjoined. Third article of antenna laterally extending into leaf-shaped structure.

Description: Full description is provided as descriptive matrix in supplement.

Remarks: Specimens like the here described ones appear to have been treated among collectors and sellers as *Palibacus praecursor* (Dames, 1886). Yet, the material of *Pb. praecursor* described by Förster (1984) has a strongly more derived antenna and carapace morphology (work in preparation). The material of Förster appears to be conspecific with the incomplete type described by Dames

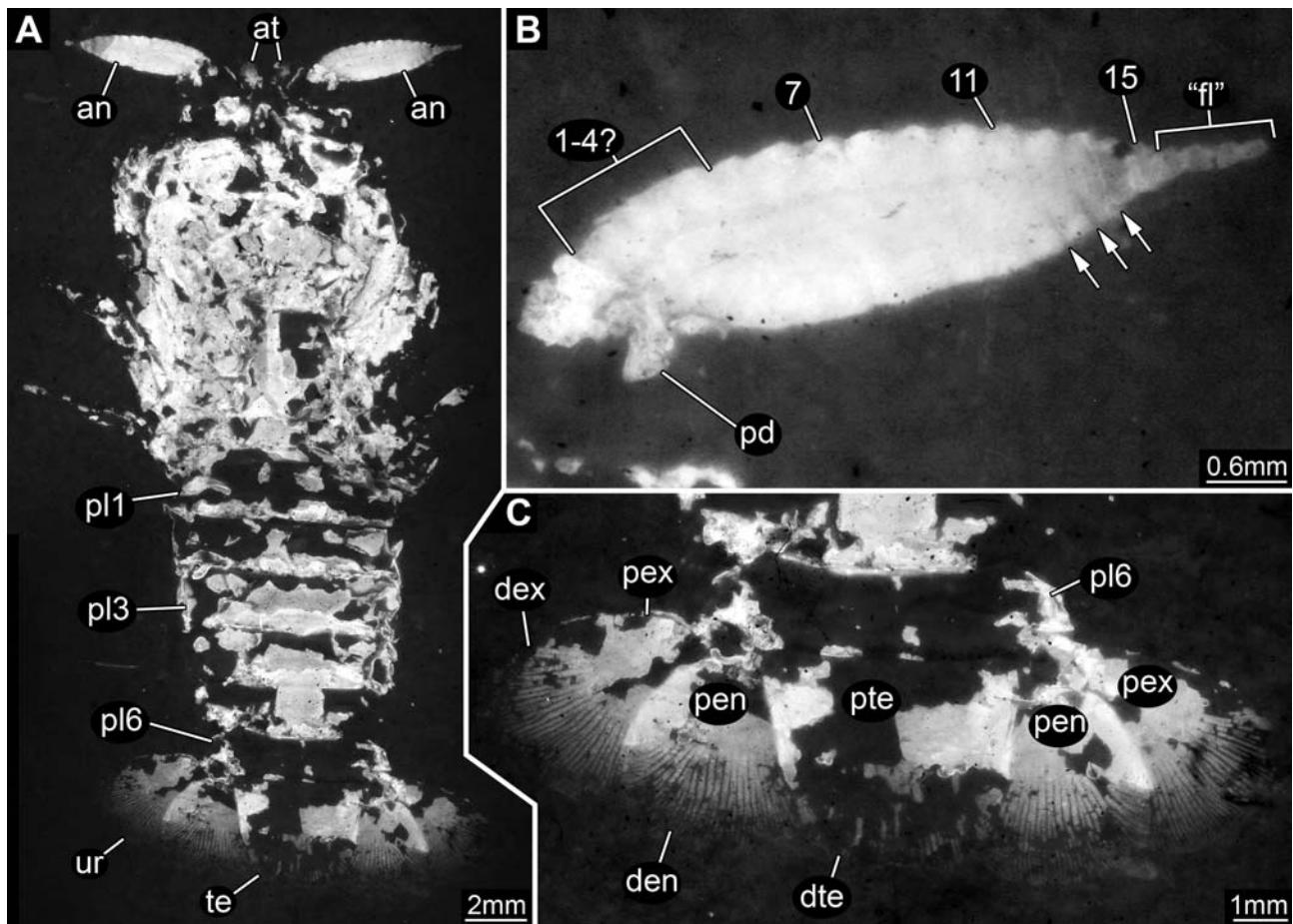


Fig. 3. *Paracancrios libanensis* (Garassino and Schweigert, 2006), early instar MNHN.FA48952. A. Overview. B. Close-up of petaloid region. Note the conjoined proximal region (1–4?), the distal articles (marked by arrows) and the distal short flagellum ("fl"). C. Close up of tail fan. Note the proximal hard region of telson (pte) and uropod rami (pen, pex) and the distal soft regions (dte, den, dex). Abbreviations: 1–15 = antennal articles; an = antenna; at = antennula; den = distal uropod endopod; dex = distal uropod exopod; dte = distal part of telson; "fl" = flagellum; pd = peduncle element; pen = proximal uropod endopod; pex = proximal uropod exopod; pl 1–6 = pleomeres; pte = proximal region of telson; te = telson; ur = uropod.

(1886). Hence, the here described specimens represent a yet unnamed species, now called *Ch. maronites*.

4. Discussion

4.1. Preservation and degree of detail

The preservation of the here described specimens is exceptional. It is not restricted to sclerotized hard parts. Also soft details like those of the distal parts of the tail fan are perfectly preserved. Yet, comparing our observations to earlier ones indicates that even exceptionally preserved details demand for the right documentation method. Looking at the original image of the antenna of the types of *P. libanensis* (Garassino and Schweigert, 2006; their Fig. 6) one gets the impression of a complete subdivision, including that of the proximal articles. This impression is most likely caused by shadows. These are caused by slight elevations that might indicate a former or future (see below) subdivision. Only the fluorescence images reveal the differences of the articulated situation of the distal articles and the conjoined condition of the proximal ones. Hence, fluorescence imaging again proves to be valuable in revealing important details (for further discussions, see for example Haug et al., 2009b, 2011a; Kerp and Bomfleur, 2011, and references therein).

4.2. Ascription to *Paracancrios libanensis*

The newly described specimens differ in some aspects of their morphology from the two type specimens of *P. libanensis*. We interpret these minor differences as caused by ontogenetic variance. The two new specimens are slightly smaller than the two type specimens. Comparing their size indicates that the two new specimens are probably one instar younger than the type specimens.

The new specimens of *P. libanensis* possess a short flagellate part distally on the petaloid region of the antenna (Fig. 3B) which appears to be absent in the type specimens (Fig. 2). In the ontogeny of *C. claviger* a long flagellate antenna at first becomes widened proximally during late juvenile development, hence roughly resembling the condition in the specimens at hand. The distal flagellum finally gets lost in *C. claviger*, retaining only the proximal (then petaloid) part (Haug et al., 2009a). Hence, this demonstrates that it is likely that the same type of ontogenetic loss is also present in the here described case. This interpretation provides a possible explanation for the fact that the antennae in the type specimens have fewer articles.

As all specimens come from the same locality, we cannot argue with geographic or time differences for basing a new species on the two new specimens. An interpretation as an ontogenetic difference seems more parsimonious than erecting a new species based on these differences.

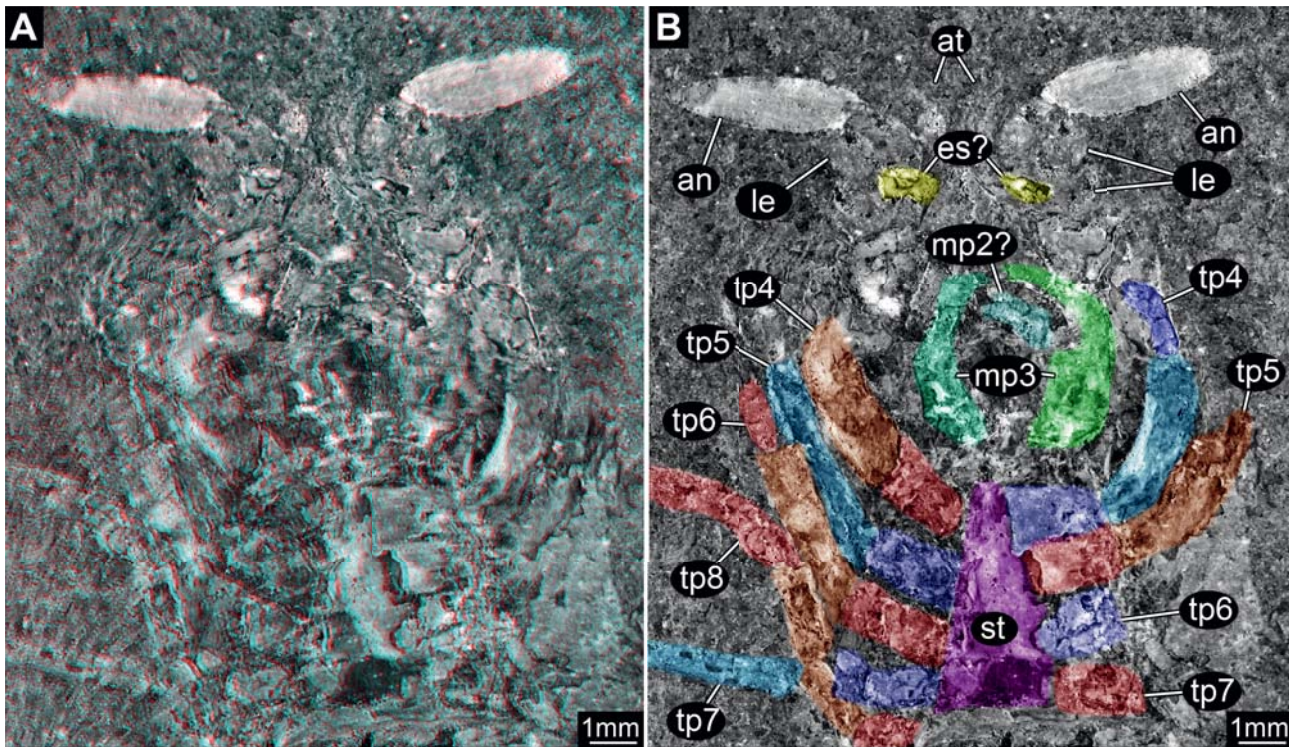


Fig. 4. *Paracancrius libanensis* (Garassino and Schweigert, 2006), early instar MNHN.F.48952, continued. A. Red–cyan stereo-anaglyph (please use red–cyan glasses to view, red left, cyan right) of the ventral region of the cephalothorax. B. Colour-marked version of A. Abbreviations: an = antenna; at = antennula; es? = possible eye stalks; le = lateral extension; mp2? = possible maxilliped 2; mp3 = maxilliped 3; st = thoracic sternum; tp4–8 = thoracopods.

4.3. Evolutionary scenario: interpreting antennal morphologies

The antennal morphologies of the species described here are significant for understanding the evolution of the peculiar scyllarid antenna (Fig. 7). *C. claviger* possesses a distal petaloid-shaped region of the antenna, comprising about 20 articles of the flagellum. The peduncle elements 3 and 4 are symmetrically shaped. This condition is still very similar to that found in Palinurida (interpreted as retained plesiomorphies), besides the short and petaloid shape of the distal part.

A relatively similar condition is seen in *P. libanensis*. Also the newly found short rostrum could represent a rather plesiomorphic character as it resembles that seen, e.g., in species of the palinurid group *Palinurellus*. Yet, certain aspects of the antenna of *P. libanensis* appear further derived than in *C. claviger*. The petaloid region resembles that of *C. claviger*; it most likely also has about or slightly less than 20 articles. Yet, unlike in *C. claviger* only the distal articles appear to be fully separated from each other. In the proximal region single articles are recognizable by indentations along the sides, yet, they appear to be continuous along the main surface (Fig. 3B). We interpret this partially conjoined condition of articles as leading to a fully conjoined condition (see further below). Furthermore, peduncle elements 3 and 4 are asymmetrical; they are slightly drawn out laterally (Figs. 4B and 5B).

In *Ch. maronites* the condition is further derived. The antenna is already clearly recognizable as a “typical” scyllarid antenna. Only a single element forms the distal petaloid part, no subdivisions are apparent (Fig. 6B,E,F). Yet, the general shape and the indentations along the rim strongly resemble the condition seen in *P. libanensis*. We therefore see this as a support of the assumption that the distal petaloid parts of *C. claviger* and *P. libanensis* are homologous to the

petaloid region in *Ch. maronites* and modern scyllarids, and that the single petaloid structure is a product of fusion, or better non-separation of several elements. The indentations are consequently interpreted as an evolutionary “left-over” of this subdivision. Spines along the margin most likely evolved within modern scyllarids and are likely not homologous to these indents. The lateral edge of peduncle element 3 is drawn out into a petaloid extension with few teeth (Fig. 6E,F). Also this lateral extension is interpreted to be homologous to the lateral extension seen in *P. libanensis* (Fig. 3B).

In modern scyllarids, the antennal morphology is further derived. The distal part is, in most modern species, no longer symmetrical: this is most likely a ground pattern feature for the group (autapomorphy). The distal tip appears to point further laterally in modern scyllarids, which means that the median edge appears longer than the lateral edge. Furthermore, the lateral extension is in most species (and probably in the ground pattern) further elongated than in *Ch. maronites*.

4.4. New systematics

Haug et al. (2009a) suggested the name Scyllaridae *sensu lato* for the group comprising *Cancrios* and modern slipper lobsters, which were suggested to be named Scyllaridae *sensu stricto*. This was intended to represent a strict phylogenetic naming. Yet, Karasawa et al. (2013) rejected these suggested group names, re-establishing a superfluous category (“empty bracket”), Cancriinidae, obviously mistaking the strict phylogenetic terms of Haug et al. (2009a) for taxonomic ones. To overcome this misunderstanding, we suggest here new terms for the different monophyletic groups in the evolutionary lineage towards modern slipper lobsters, emphasizing their phylogenetic meaning. Following Scholtz and

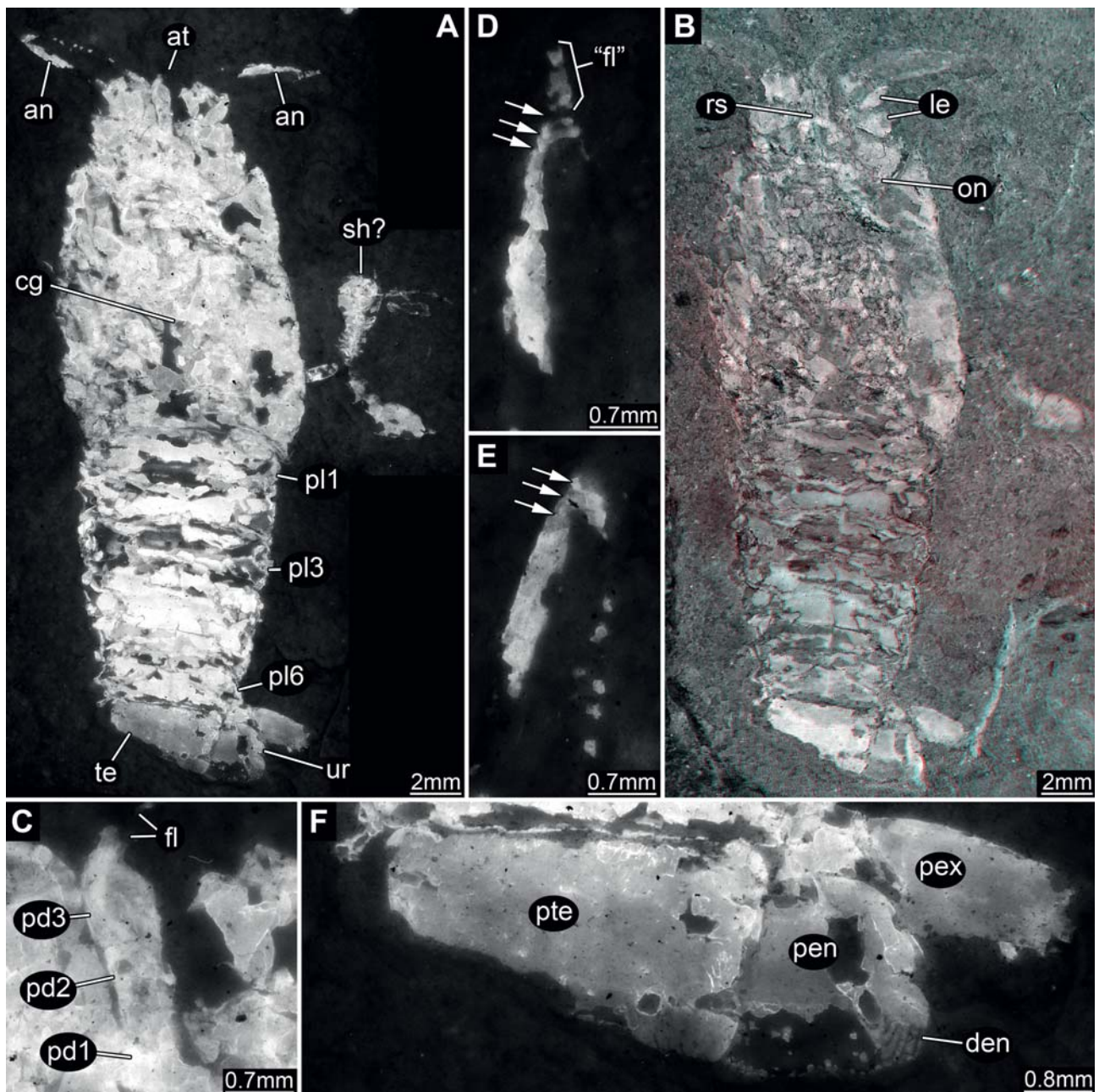


Fig. 5. *Paracancrinos libanensis* (Garassino and Schweigert, 2006), early instar MNHN.FA48973. A. Overview. B. Red–cyan stereo-anaglyph (please use red–cyan glasses to view, red left, cyan right) of the specimen. C. Close up of antennula. D, E: Close-up of antenna. Note short flagellum ("fl"). Arrows mark boundaries between articles. F. Close-up of tail fan. Abbreviations: an = antenna; at = antennula; cg = cervical groove; den = distal uropod endopod; fl = flagellum; le = lateral extension; on = optical notch; pd1–3 = peduncle element; pen = proximal uropod endopod; pex = proximal uropod exopod; pl1–6 = pleomeres; pte = proximal region of telson; rs = rostrum; sh? = possible small shrimp; te = telson; ur = uropod.

Richter (1995), any endings related to specific ranks (e.g. -idae) are transformed to a neutral ending (-a).

Scyllarida sensu lato tax. nov. is autapomorphically characterized by the petaloid shape of the distal part of the antenna in the adult stage, which consists of about 20 well-separated elements. The monophyletic group comprises *C. claviger* and *Scyllarida sensu stricto* tax. nov. This characterization of *Scyllarida sensu lato* causes the difficulty that there is currently no character "left" for characterizing (diagnosing) a monophyletic group *Cancrienos*, or the species *C. claviger*. This is due to the relatively high resolution of the here discussed phylogeny.

Scyllarida sensu stricto tax. nov. is autapomorphically characterized by the partially conjoined condition of the proximal flagellar antennal articles. The known representatives additionally are characterized by the lateral extension of antennal peduncle element 3. The monophyletic group comprises *P. libanensis* and *Verscyllarida* tax. nov.

Verscyllarida tax. nov. is autapomorphically characterized by the fully conjoined condition of the distal antennal articles. All known representatives additionally show the further lateral extension of the third peduncle element of the antenna. The monophyletic group comprises *Ch. maronites* and *Neoscyllarida* tax. nov.

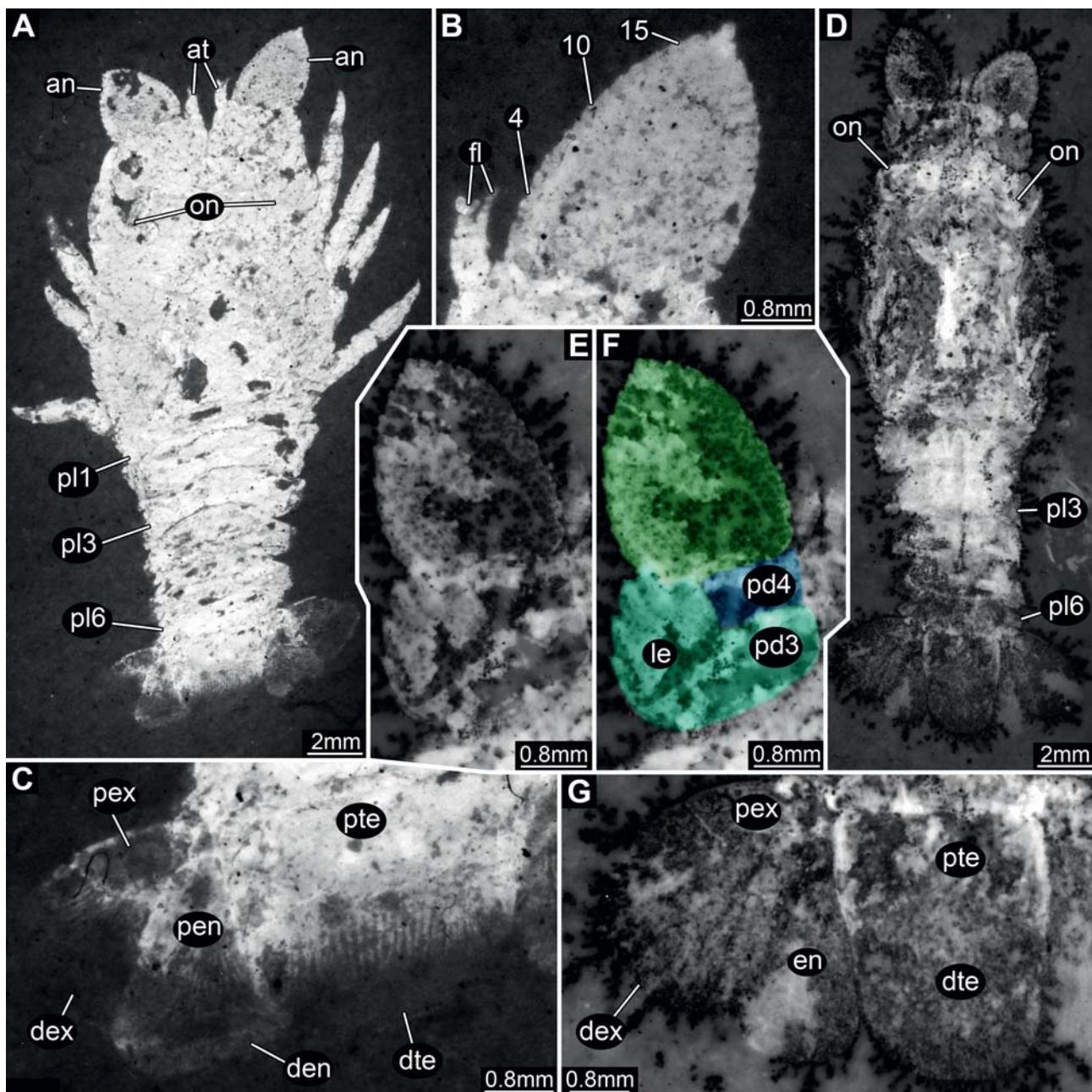


Fig. 6. *Charbelicaris maronites* gen. nov., sp. nov. from the Hadjoula Lagerstätte (Cretaceous, Lebanon). A–C. Holotype MNHN.F.A48955. A. Overview. B. Close-up of antennula and antenna. C. Close-up of tail fan. D–G. Paratype MNHN.F.A48984. D. Overview. E. Close-up of antenna. F. Colour-marked version of E. G. Close-up of tail fan. Abbreviations: 1–15 = former antennal articles; an = antenna; at = antennula; den = distal uropod endopod; dex = distal uropod exopod; dte = distal part of telson; fl = flagellum; le = lateral extension; on = optical notch; pen = proximal uropod endopod; pex = proximal uropod exopod; pl1–6 = pleomeres; pte = proximal region of telson.

Neoscyllarida tax. nov. is characterized by a shape change of the distal antennal region. It is no longer petaloid but more rounded, most likely by allometric growth of the original inner edge. Additionally, the lateral extensions of peduncle element 3 are more drawn out. Neoscyllarida comprises all modern representatives = crown-group Scyllaridae.

5. Conclusions and outlook

The here presented case supports that important evolutionary novelties in scyllarids occurred during the Jurassic and Cretaceous (Förster, 1973, 1984). Due to the exceptional preservation of several

Lagerstätten in the Jurassic and Cretaceous we have direct access to these evolutionary changes. In the days of evo-devo it is especially fortunate that such fossils, e.g. of achelate lobsters, even give insights into the ontogeny (Polz, 1971, 1972, 1973, 1987, 1995, 1996; Tanaka et al., 2009; Haug et al., 2011c, 2015; Audo and Charbonnier, 2012) and thus facilitate a palaeo-evo-devo approach (Haug et al., 2009a, 2013; Haug and Haug, 2013). More scyllarid and other achelatan fossils from the Mesozoic are currently worked up, and will likely provide new and additional insights.

To summarise, our new finds on Mesozoic fossils demonstrate:

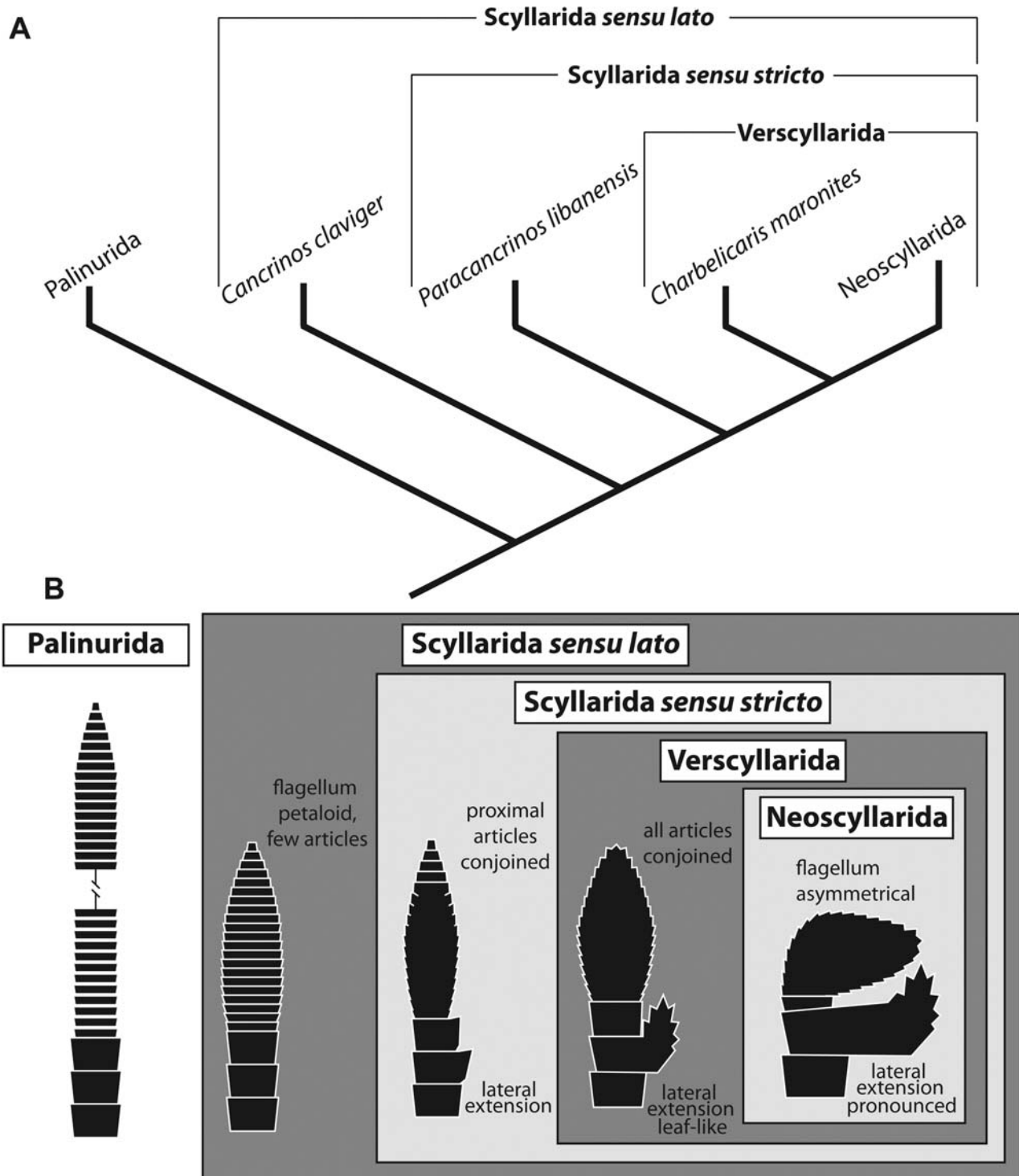


Fig. 7. Phylogeny and evolution of Achelata. A. Phylogram of Achelata. B. Evolutionary scenario of the antenna of the early lineage towards modern slipper lobsters (antennae depicted without peduncle element 1). In Palinurida the antenna is relatively slender, elongate and composed of numerous elements. In Scyllarida sensu lato the distal part of the antenna is petaloid and composed of about 20 elements, which are fully articulated. In Scyllarida sensu stricto the distal region is petaloid and composed of about 15 elements; only the distal elements are fully articulated. The peduncle elements are moderately laterally extended; optical notches are present. In Verscyllarida the petaloid region is fully conjoined, the lateral extension of the third peduncle article is leaf-shaped. In Neoscyllarida the petaloid region is asymmetrical and the lateral extension is more pronounced.

- the importance of applying the right documentation method to exceptionally preserved fossils,
- the step-wise evolution of the scyllarid morphotype, documented by these,
- that fossils are extremely important to resolve the stepwise evolution of a distinct modern morphotype.

Acknowledgements

This study was supported by numerous institutions and persons, which are all heartily thanked for. We thank Jonas Keiler, Rostock, and an anonymous reviewer for their comments on the manuscript. The visit of JTH to the MNHN was kindly funded by a

grant from the European Commission's (FP6) Integrated Infrastructure Initiative programme SYNTHESYS FR-TAF-2590. CH has been supported by the German Academic Exchange Service (DAAD) with a return fellowship. JTH has been kindly funded by the Alexander von Humboldt Foundation with a Feodor Lynen return fellowship. This study is part of the project 'Palaeo-Evo-Devo of Malacostraca' of JTH kindly funded by the German Research Foundation (DFG) under Ha 6300/3-1. Field campaign in Lebanon was granted by the MNHN-ATM program Biodiversité actuelle et fossile. Crises, stress, restaurations et panchronisme: le message systématique. JTH and CH want to express their thanks to their former host Steffen Harzsch, Greifswald and to J. Matthias Starck, Munich for their support. We thank all people involved in providing open source, open access or low cost software, which also was used during this study, such as Open Office, GIMP, or Inkscape.

Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.asd.2015.08.003>.

References

- Ahyong, S.T., Garassino, A., Gironi, G., 2007. *Archaeosculda phoenicia* n. gen., n. sp. (Crustacea, Stomatopoda, Pseudosculdidae) from the upper Cretaceous (Cenomanian) of Lebanon. Atti Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 148 (1), 3–15.
- Audo, D., Charbonnier, S., 2012. New nisto of slipper lobster (Decapoda: Scyllaridae) from the Hadjoula Lagerstätte (Late Cretaceous, Lebanon). J. Crust. Biol. 32, 583–590.
- Audo, D., Charbonnier, S., 2013. Late Cretaceous crest-bearing shrimps from the Sahel Alma Lagerstätte of Lebanon. Acta Palaeontol. Pol. 58, 335–349.
- Bracken-Crisson, H.D., Ahyong, S.T., Wilkinson, R., Yang, Z., Briehnholz, J., Palero, F., Chan, T.-Y., Bendall, M., Feldmann, R., Schweitzer, C., Chu, K.H., Felder, D.L., Robles, R., Kim, D., Tsang, M., Martin, J., Crandall, K.A., 2014. Emergence of the lobsters: phylogenetic relationships, morphological evolution and divergence time comparisons of a fossil rich group (Achelata, Astacidea, Glypheidea, Polychelida). Syst. Biol. 63, 457–479.
- Brocchi, P., 1875. Note sur une nouvelle espèce de crustacé fossile (*Penaeus libanensis*). Bull. Soc. Géol. Fr. 3, 609–610.
- Charbonnier, S., Garassino, A., Schweigert, G., Simpson, M., 2013. A worldwide review of fossil and extant glypheid and litogastrid lobsters (Crustacea, Decapoda, Glypheidea). Mém. Mus. Natl. Hist. Nat. Paris 205, 1–304.
- Dames, W., 1886. Ueber einige Crustaceen aus dem Kreideablagerungen des Libanon. Z. Dtsch. Geol. Ges. 38, 551–576.
- Feldmann, R.M., 2009. A new cirolanid isopod (Crustacea) from the Cretaceous of Lebanon: dermoliths document the pre-molt condition. J. Crust. Biol. 29, 373–378.
- Feldmann, R.M., Charbonnier, S., 2011. *Ibacus cottreai* Roger, 1946, reassigned to the isopod genus *Cirolana* (Cymothoidae). J. Crust. Biol. 31, 317–319.
- Ferry, S., Merran, Y., Grosheny, D., Mroueh, M., 2007. Relations entre les marges septentrionale et méridionale de la Téthys au Crétacé. Carn. Géol. 6, 38–42.
- Förster, R., 1973. Untersuchungen an oberjurassischen Palinuridae (Crustacea, Decapoda). Mitt. Bayer. Staatsamml. Paläo. Hist. Geol. 13, 31–46.
- Förster, R., 1984. Bärenkrebse aus dem Cenoman des Libanon und dem Eozän Italiens. Mitt. Bayer. Staatsamml. Paläo. Hist. Geol. 24, 57–66.
- Förster, R., 1985. Evolutionary trends and ecology of Mesozoic decapod crustaceans. Trans. R. Soc. Edinb. 76, 299–304.
- Fraas, O., 1878. Geologisches aus dem Libanon. Jahresh. Vereins Naturk. Württ. 34, 257–391.
- Garassino, A., 1994. The macruran decapod crustaceans of the upper Cretaceous of Lebanon. Paleo. Lombar. Nuova Ser. Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 3, 3–27.
- Garassino, A., 2001. New decapod crustaceans from the Cenomanian (Upper Cretaceous) of Lebanon. Atti. Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 141 (2), 237–250.
- Garassino, A., Schweigert, G., 2006. *Cretasergestes sahelalmae* n. gen., n. sp. (Crustacea, Decapoda, Sergestoidae) and *Cancrius libanensis* n. sp. (Crustacea, Decapoda, Palinuroidea) from the upper Cretaceous (Cenomanian) of Lebanon. Atti Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 147, 69–78.
- Garassino, A., De Angeli, A., Pasini, G., 2009. A new hermit crab (Crustacea, Anomura, Paguroidea) from the late Cretaceous (Cenomanian) of Lebanon. Atti Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 150 (2), 215–228.
- Glaessner, M.F., 1945. Cretaceous Crustacea from mount Lebanon, Syria. Ann. Mag. Nat. Hist. 11, 694–707.
- Haug, J.T., Haug, C., Waloszek, D., Maas, A., Wulf, M., Schweigert, G., 2009a. Development in Mesozoic scyllarids and implications for the evolution of Achelata (Reptantia, Decapoda, Crustacea). Palaeodiversity 2, 97–110.
- Haug, C., Haug, J.T., Waloszek, D., Maas, A., Frattigiani, R., Liebau, S., 2009b. New methods to document fossils from lithographic limestones of southern Germany and Lebanon. Palaeontol. Electron 12 (3) art.6T. http://palaeo-electronica.org/2009_3/193/index.html.
- Haug, J.T., Haug, C., Maas, A., Kutschera, V., Waloszek, D., 2010. Evolution of mantis shrimps (Stomatopoda, Malacostraca) in the light of new Mesozoic fossils. BMC Evol. Biol. 10 art. 290, 17 pp.
- Haug, J.T., Haug, C., Kutschera, V., Mayer, G., Maas, A., Liebau, S., Castellani, C., Wolfram, U., Clarkson, E.N.K., Waloszek, D., 2011a. Autofluorescence imaging, an excellent tool for comparative morphology. J. Microsc. 244, 259–272.
- Haug, C., Mayer, G., Kutschera, V., Waloszek, D., Maas, A., Haug, J.T., 2011b. Imaging and documenting gammarideans. Int. J. Zool. <http://dx.doi.org/10.1155/2011/380829> art. 380829.
- Haug, J.T., Haug, C., Waloszek, D., Schweigert, G., 2011c. The importance of lithographic limestones for revealing ontogenies in fossil crustaceans. Swiss J. Geosci. 104 (Suppl. 1), S85–S98.
- Haug, J.T., Haug, C., 2011. Fossilien unter langwelligem Licht: Grün-Orange-Fluoreszenz an makroskopischen Objekten. Archaeopteryx 29, 20–23.
- Haug, J.T., Mayer, G., Haug, C., Briggs, D.E.G., 2012a. A Carboniferous non-onychophoran lobopodian reveals long-term survival of a Cambrian morphotype. Curr. Biol. 22, 1673–1675.
- Haug, J.T., Briggs, D.E.G., Haug, C., 2012b. Morphology and function in the Cambrian Burgess Shale megacheiran arthropod *Leanchoilia superlata* and the application of a descriptive matrix. BMC Evol. Biol. 12 art. 162.
- Haug, J.T., Audo, D., Charbonnier, S., Haug, C., 2013. Diversity of developmental patterns in achelate lobsters—today and in the Mesozoic. Dev. Genes. Evol. 223, 363–373.
- Haug, J.T., Haug, C., 2013. An unusual fossil larva, the ontogeny of achelatan lobsters, and the evolution of metamorphosis. Bull. Geosci. 88, 195–206.
- Haug, J.T., Audo, D., Haug, C., Abi Saad, P., Petit, G., Charbonnier, S., 2015. Unique occurrence of polychelidan lobster larvae in the fossil record and its evolutionary implications. Gondwana Res. 28, 869–874.
- Hiller, A., Viviana, C.A., Werding, B., 2010. Hypercarcinisation: an evolutionary novelty in the commensal porcellanid *Allopeltostethes spinifrons* (Crustacea: Decapoda: Porcellanidae). Nauplius 18, 95–102.
- Holthuis, L.B., 1985. A revision of the family Scyllaridae (Crustacea: Decapoda: Macrura). I. Subfamily Ibacinae. Zool. Verhandl 218, 1–130.
- Holthuis, L.B., 1991. FAO Species Catalogue. Marine Lobsters of the world. An annotated and illustrated catalogue of species of interest to fisheries known to date. FAO Fish. Synop. 125 (13), 292. Rome.
- Jones, C.M., 1990. Morphological characteristics of bay lobsters, *Thenus*, Leach species (Decapoda, Scyllaridae) from North-Eastern Australia. Crustaceana 59, 265–275.
- Karasawa, H., Schweitzer, C.E., Feldmann, R.M., 2013. Phylogeny and systematics of extant and extinct lobsters. J. Crust. Biol. 33, 78–123.
- Kerp, H., Bomfleur, B., 2011. Photography of plant fossils—new techniques, old tricks. Rev. Palaeobot. Palynol. 166, 117–151.
- Larghi, C., 2004. Brachyruran decapod Crustacea from the upper Cretaceous of Lebanon. J. Paleo. 78, 528–541.
- McLaughlin, P.A., Lemaitre, R., 1997. Carcinization in the Anomura – fact or fiction? I. Evidence from adult morphology. Contrib. Zool. 67, 79–123.
- McLaughlin, P.A., Lemaitre, R., Tudge, C.C., 2004. Carcinization in the Anomura – fact or fiction. II. Evidence from larval, megalopal and early juvenile morphology. Contrib. Zool. 73, 165–205.
- Palero, F., Crandall, K.A., Abelló, P., Macpherson, E., Pascual, M., 2009. Phylogenetic relationships between spiny, slipper and coral lobsters (Crustacea, Decapoda, Achelata). Mol. Phylogen. Evol. 50, 152–162.
- Pasini, G., Garassino, A., 2011. First direct predatory evidence of an eel-like fish on *Pseudosculda* cfr. *P. laevis* (Crustacea, Stomatopoda, Unipeltata) from the Late Cretaceous (Cenomanian) of Lebanon. Atti Soc. Ital. Sci. Nat. Mus. Civ. St. Nat. Milano 152 (1), 69–78.
- Petit, G., Charbonnier, S., 2012. Fossil sponge gemmules, epibionts of *Carpopenaeus garassinoi* n. sp. (Crustacea, Decapoda) from the Sahel Alma Lagerstätte (Late Cretaceous, Lebanon). Geodiversitas 34, 359–372.
- Polz, H., 1971. Eine weitere Phyllosoma-Larve aus den Solnhofener Plattenkalken. N. Jb. Geol. Paläontol. Mh. 1971, 474–488.
- Polz, H., 1972. Entwicklungsstadien bei fossilen Phyllosomen (Form A) aus den Solnhofener Plattenkalken. N. Jb. Geol. Paläontol. Mh. 1972, 678–689.
- Polz, H., 1973. Entwicklungsstadien bei fossilen Phyllosomen (Form B) aus den Solnhofener Plattenkalken. N. Jb. Geol. Paläontol. Mh. 1973, 284–296.
- Polz, H., 1987. Zur Differenzierung der fossilen Phyllosomen (Crustacea, Decapoda) aus den Solnhofener Plattenkalken. Archaeopteryx 5, 23–32.
- Polz, H., 1995. Ein außergewöhnliches Jugendstadium eines palinuriden Krebses aus den Solnhofener Plattenkalken. Archaeopteryx 13, 67–74.
- Polz, H., 1996. Eine Form-C-Krebslarve mit erhaltenem Kopfschild (Crustacea, Decapoda, Palinuroidea) aus den Solnhofener Plattenkalken. Archaeopteryx 14, 43–50.
- Roger, J., 1946. Invertébrés des couches à poissons du Crétacé supérieur du Liban. Mém. Soc. Géol. Fr. Nouv. Sér. 51, 1–92.
- Scholtz, G., 2014. Evolution of crabs – history and deconstruction of a prime example of convergence. Contrib. Zool. 83, 87–105.
- Scholtz, G., Richter, S., 1995. Phylogenetic systematics of the reptantian Decapoda (Crustacea, Malacostraca). Zool. J. Linn. Soc. 113, 289–328.

- Senter, P., 2010. Using creation science to demonstrate evolution: application of a method for visualizing gaps in the fossil record to a phylogenetic study of coelurosaurian dinosaurs. *J. Evol. Biol.* 23, 1732–1743.
- Tanaka, G., Smith, R.J., Siveter, D.J., Parker, A.R., 2009. Three-dimensionally preserved decapod larval compound eyes from the Cretaceous Santana formation of Brazil. *Zool. Sci.* 26, 846–850.
- Tsang, L.M., Chan, T.-Y., Ahyong, S.T., Chu, K.H., 2011. Hermit to king, or hermit to all: multiple transitions to crab-like forms from hermit crab ancestors. *Syst. Biol.* 60, 616–629.