Re-appraisal of thylacocephalans (Euarthropoda, Thylacocephala) from the Jurassic La Voulte-sur-Rhône Lagerstätte

Thomas LAVILLE 1,2, Marie-Béatrice FOREL 2 & Sylvain CHARBONNIER 3

1,2,3 Muséum national d’Histoire naturelle, Centre de Recherche en Paléontologie – Paris (CR2P, UMR 7207), Sorbonne Université-MNHN-CNRS, 57 rue Cuvier, 75005 Paris, France.

Abstract. Thylacocephalans are enigmatic euarthropods, known at least from the Silurian to the Cretaceous. Despite remaining uncertainties concerning their anatomy, key features can be recognised such as a shield enveloping most of the body, hypertrophied compound eyes, three pairs of raptorial appendages and a posterior trunk consisting of eight up to 22 segments bearing appendages and eight pairs of gills. Well-known for its euarthropod diversity, the La Voulte-sur-Rhône Lagerstätte (Callovian, Middle Jurassic, France) has provided many remains of four thylacocephalan species so far: Dollocaris ingens, Kilianicaris lerichei, Paraostenia voultensis and Clausocaris ribeti. In this paper, we study the type material as well as undescribed material. The re-description of La Voulte thylacocephalans reveals an unexpected diversity, with the description of two new species, Austriocaris secretanae and Paraclausocaris harpa gen. et sp. nov., and of specimens of Mayrocaris, a taxon originally described from Solnhofen Lagerstätten. We also reassign Clausocaris ribeti to Ostenocaris. The reappraisal of La Voulte thylacocephalans also provides important insight into the palaeobiology of Thylacocephala. New key anatomical features are described, such as an oval structure or a putative statocyst, which indicate a nektonic or nektobenthic lifestyle. Finally, we document a juvenile stage for Paraostenia voultensis.

Keywords. Euarthropoda, Jurassic, anatomy, disparity, morphometrics.
compound eyes, three pairs of raptorial appendages, a set of eight pairs of gills, and from eight up to 22 posterior trunk segments bearing appendages (Schram 2014). Known from at the least the Silurian to the Cretaceous (Schram 2014), thylacocephalans have a scarce fossil record in the Jurassic. They have only been reported from four different European localities (Table 1): Osteno Lagerstätte (Italy, Sinemurian, Early Jurassic: Arduini et al. 1980; Pinna et al. 1982, 1985), Strawberry Bank Lagerstätte (England, Toarcian, Early Jurassic: Williams et al. 2015), La Voult-sur-Rhône Lagerstätte (France, Callovian, Middle Jurassic: Van Straelen 1923a; Secrétan & Riou 1983; Secrétan 1985; Charbonnier et al. 2010) and Solnhofen Limestones (Germany, Tithonian, Late Jurassic: e.g., Oppenheim 1888; Polz 1990, 1994, 2001; Braig et al. 2019). Despite this limited distribution during the Jurassic, 11 thylacocephalan species have been described so far (Table 1).

In addition to this important diversity, Jurassic representatives are among the best-preserved thylacocephalans. While most species are essentially described based on their shield, Jurassic representatives are often preserved with soft-parts. For instance, the organisation of the digestive, respiratory and visual systems has been reported for Dollocaris ingens Secrétan, 1985 (La Voult Lagerstätte; Vannier et al. 2016). Moreover, cephalic appendages have been described from Solnhofen representatives of Clausocraris lithographica (Oppenheim, 1888) (see Laville et al. 2021a) and Mayrocaris bucculata Polz, 1994 (see Haug et al. 2014). These anatomical details are important for resolving the phylogenetic affinities of Thylacocephala (e.g., Haug et al. 2014; Vannier et al. 2016; Broda & Zatoń 2017). Even though many morphological details point out to a pancrustacean affinity, the position of Thylacocephala relative to other pancrustacean ingroups is uncertain. Affinities with cirripeds (Pinna et al. 1982, 1985), malacostracans (Secrétan 1985; Vannier et al. 2016) and remipeds (Haug et al. 2014) have been suggested but no consensus exists on this topic.

In this paper, we revise known species from the La Voult-sur-Rhône Lagerstätte and describe new species. This descriptive work provides new important insight into the diversity, morphology and palaeobiology of thylacocephalans. Furthermore, the exceptional preservation of the La Voult specimens clarifies key aspects of the anatomy of thylacocephalans.

**Geological setting**

The La Voult-sur-Rhône Lagerstätte (Middle Jurassic, early Callovian, Gracils Biozone) is located in Ardèche, France and belongs to the eastern sedimentary cover of the Massif Central (Fig. 1A–B). Following the onset of the transgression in the late Bathonian, the Tethys Ocean was characterised by a high eustatic level during the Callovian (Jacquin & Graciansky 1999; Hallam 2001). This event was associated with an important period of rifting which led to the overflow of epicontinental seas on the peri-tethyan domain and to well-developed terrigenous platforms with only few, small carbonate platforms in Western Europe (Enay et al. 1993). The deposits of the La Voult-sur-Rhône Lagerstätte are particularly linked to this context. During the Callovian, La Voult-sur-Rhône was located on the Ardèche palaeomargin, adjacent to the submerged Hercynian Massif Central and the Subalpine Basin. This latest was connected to the Tethys Ocean (Enay 1993).

The La Voult Lagerstätte corresponds to about 5–6 m of marls, mainly composed of a clay mineralogy (60–70% of montmorillonite), and topped by 15 m of iron carbonate deposits (Fig. 1C). Described for the first time by Fournet (1843), the La Voult Lagerstätte has yielded a well-diversified non-vertebrate fauna with few vertebrate remains (actinopterygians, sarcopterygians and reptiles; Charbonnier 2009). This Lagerstätte is famous for its fossils with soft-tissues preservation, especially among pancrustaceans and cephalopods (Wilby et al. 1996). Fossils are either preserved in 3D in sideritic nodules uncovered from the basal marls, or compressed in other marly layers (Fischer et al. 2003; Charbonnier et al. 2014). Jauvion et al. (2019) studied in detail the preservation sequence of Thylacocephala in nodules from La Voult and highlighted the important role of the early precipitation of fluorapatite and pyrite in soft-tissue preservation and the various mineralogies associated with the different tissue types.
Table 1. Diversity of Jurassic thylacocephalans.

<table>
<thead>
<tr>
<th>Series</th>
<th>Age</th>
<th>Locality</th>
<th>Species</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Toarcian</td>
<td>Strawberry Bank Lagerstätte, Ilminster, Somerset, UK</td>
<td>No described or figured specimens</td>
<td>Williams <em>et al.</em> 2015</td>
</tr>
<tr>
<td>Middle Jurassic</td>
<td>Callovian</td>
<td>La Voult-sur-Rhône, Ardèche, France</td>
<td><em>Austriocaris secretanae</em> sp. nov.</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Dollocaris ingens</em> Van Straelen, 1923</td>
<td>Van Straelen 1923; Secrétan &amp; Riou 1983; Secrétan 1985; Charbonnier <em>et al.</em> 2010; Vannier <em>et al.</em> 2016</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Kilianicaris lerichei</em> Van Straelen, 1923</td>
<td>Van Straelen 1923; Secrétan &amp; Riou 1983; Secrétan 1985; Charbonnier <em>et al.</em> 2010; This paper</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Mayrocaris</em> sp.</td>
<td>This paper</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td><em>Ostenocaris ribeti</em> (Secrétan, 1985)</td>
<td>Secrétan &amp; Riou 1983; Secrétan 1985; Charbonnier <em>et al.</em> 2010; This paper</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Paraclausocaris harpa</em> gen. et sp. nov.</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Paraostenia voutensis</em> Secrétan, 1985</td>
<td>Secrétan &amp; Riou 1983; Secrétan 1985; Charbonnier <em>et al.</em> 2010; This paper</td>
</tr>
<tr>
<td>Late Jurassic</td>
<td>Tithonian</td>
<td>Solnhofen Lagerstätten, Bavaria, Germany</td>
<td><em>Clausocaris lithographica</em> (Oppenheim, 1888)</td>
<td>Oppenheim 1888; Polz 1988, 1989, 1990, 1993; Haug <em>et al.</em> 2014; Laville <em>et al.</em> 2021a</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td><em>Dollocaris michelorum</em> Polz, 2001</td>
<td>Polz 2001</td>
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<td></td>
<td></td>
<td></td>
<td><em>Falcatacaris bastelbergeri</em> Braig, Haug J.T, Schädel &amp; Haug C., 2019</td>
<td>Braig <em>et al.</em> 2019</td>
</tr>
</tbody>
</table>
Sixty species have been described, including bivalves, cephalopods (Fischer & Riou 1982, 2002; Fuchs 2014; Kruta et al. 2016), annelids (Alessandrello et al. 2004), pycnogonids (Charbonnier et al. 2007b), pancrustaceans (cumaceans: Bachmayer 1960; mysidaceans: Secrétan & Riou 1986; decapods: Van Straelen 1922, 1923b, 1925; Carriol & Riou 1991; Charbonnier et al. 2013; Audo et al. 2014; Jauvion

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**Fig. 1.** Geological settings of La Voult-sur-Rhône Lagerstätte. **A.** Location of La Voult Lagerstätte. **B.** Geological map of the La Voult area. **C.** Stratigraphic column of the La Voult Lagerstätte, with the distribution of thylacocephalan taxa. Abbreviations: 1 = *Austriocaris secretanae* sp. nov.; 2 = *Dollocaris ingens* Van Straelen, 1923; 3 = *Kilianicaris lerichei* Van Straelen, 1923; 4 = *Mayrocaris* sp; 5 = *Ostenocaris ribeti* (Secrétan, 1985) comb. nov.; 6 = *Paraclusocaris harpa* gen. et sp. nov.; 7 = *Paraostenia voultensis* Secrétan, 1985. Scale bars: B = 500 m; C = 1 m. A–B modified after Charbonnier et al. (2010), C modified after Charbonnier (2009).
et al. 2016, 2017, 2020; Audo & Schweigert 2018), brachiopods, echinoderms (e.g., Villier et al. 2009) and hemichordates (Alessandrello et al. 2004). Euarthropoda Lankester, 1904 is the most diverse taxon, representing nearly 50% of all taxa described from La Voulté (Charbonnier 2009). Among euarthropods, thylacocephalans are the most abundant fossils, representing 32.3% of the euarthropods recovered in nodules (Charbonnier et al. 2010). Up to now, four species had been described: *Dollocaris ingens* Van Straelen, 1923, *Kilianicaris lerichei* Van Straelen, 1923, *Paraostenia voultensis* Secrétan, 1985 and *Clausocaris ribeti* Secrétan, 1985.

Based on structural, sedimentary and faunal evidences, the La Voulté Lagerstätte corresponds to the upper part of the bathyal zone, near the slope-basin transition, with a water depth most probably exceeding 200 m (Elmi 1967; Charbonnier et al. 2007a). However, recent discoveries place doubts on this statement, providing arguments for an illuminated and thus shallower environment (Vannier et al. 2016).

**Material and methods**

**Material**

We investigated 211 specimens in this study (exhaustive list in Supp. file 1), most being preserved in compression. The studied specimens are deposited in the collections of the Muséum national d’histoire Naturelle, Paris, France (197 specimens; acronym: MNHN.F), the Observatoire des Sciences de l’Univers de Grenoble, France (three specimens; acronym: OSUG-UJF-ID), the Museo Civico di Storia Naturale, Milano, Italy (11 specimens; acronym: MSNM).

We do not include the species *Dollocaris ingens* in this study, as it has been extensively studied in the past (Secrétan & Riou 1983; Secrétan 1985; Fröhlich et al. 1992; Charbonnier 2009; Vannier et al. 2016; Jauvion et al. 2019).

**Documentation methods**

**General view**

Six specimens (MNHN.F.A29301, A29310, A29327, A53329, A53335, A53342) were imaged using a Keyence VHX-6000 digital microscope at the Zoomorphologie group, Ludwig-Maximilians Universität (LMU), München. Three different illumination settings were used: cross-polarised ring-light illumination, cross-polarised co-axial illumination or co-axial illumination. A 50 times magnification was used. The built-in high dynamic range (HDR) function was used to cope with issues of under- or over-exposure. Every image is thus a composite of several images taken under different exposure times. To overcome limitations in depth of field and in field of view, several adjacent images were recorded, each with a Z-stack of images.

All other MNHN specimens were documented using either a Nikon d700 camera equipped with a Nikon AF-S Nikkor 35 mm f/1.8g ED lens or a Nikon d5300 camera equipped with a Nikon AF-S Nikkor 35 mm f/1.8g ED lens and a Hoya HD polarising filter. Specimens were illuminated with a Kaiser RB 218n HF lighting unit. The camera was controlled for remote shooting using DigiCamControl (Duka Istvan, MIT license). To overcome limitations in depth of field, some specimens were recorded with a Z-stack of images.

MSNM specimens were recorded using a Pentax KS-2 camera equipped with a Pentax D-FA Macro 50 mm f/2.8 lens.

Pictures of anatomical details were taken either with a Canon EOS 60D camera equipped with a Canon MPE-65 mm macro lens and a Hoya HD polarising filter, illuminated with a Kaiser RB 218n HF lighting unit or with an Axio ZOOM.V16 stereo microscope equipped with a DeltaPix Invenio-20EIII camera under cross-polarised ring light at the Centre de Recherche en Paléontologie – Paris (CR2P).
Fluorescence imaging
Specimen MNHN.F.A53342 was photographed with a Keyence BZ-9000 inverse epifluorescence microscope using green light at the Zoomorphologie group, LMU, München (TRITC filter, band-pass width: 532–554 nm, with a maximal excitation wavelength of 543 nm; detection ranges: 570–613 nm) with lens magnification of ×2. Together with the magnification of the camera, this results in a magnification of about 20×. To overcome limitations in depth of field and in field of view, several adjacent images were recorded, each with a Z-stack of images. Only the luminescence was collected.

Image processing
For specimens documented with the Keyence VHX-6000 digital microscope, focus stacking and panoramic stitching were performed with the software implemented in the digital microscope. For the other specimens, Z-stacks of fluorescence images were digitally fused to single in-focus images using CombineZP (Alan Hadley, GNU).

For the specimen documented under fluorescence settings using Keyence BZ-9000 inverse epifluorescence microscope, stitching of the adjacent images was made with the Photomerge plug-in from Adobe Photoshop CS5.

Post-processing of every image (histogram optimisation, contrast, brightness and sharpness adjustment) was performed with GIMP 2.10 (GNU). Additionally, areas of interest were colour-marked to highlight morphological structures. Line drawings were made using Adobe Illustrator CS6.

Morphometric analyses
Measurements
Shield, eyes and appendages were measured directly on digital photographs with ImageJ2 (Schneider et al. 2012). Descriptive statistics are given in Table 2, which was computed using the gtsummary R package (Sjoberg et al. 2021). Measurement data are available at the following link (Supp. file 2): https://doi.org/10.5281/zenodo.6341180.

Measurement abbreviations
Aad = anterodorsal angle
Aav = anteroventral angle
Apd = posterodorsal angle
Apv = posteroventral angle
H = height
Ha = anterior shield height
Hmax = maximum shield height
Hp = posterior shield height
L_n = length of podomere n (appendages)
Lr = length of the rostrum
Ls = length of the shield
Lw = length of the shield without the rostrum
w = width
w_n = width of podomere n (appendages; Fig. 2)

Simple allometry
The allometry between shield height (Hmax) and shield length (Ls) was studied by running a standardised major axis analysis (SMA) for each species using the R package smatr ver. 3.4.8 (Warton et al. 2012). A major axis analysis is a type-II regression, which is used to find a line of best fit that summarises the relationships between two variables but not to predict the value of one variable using the other as in
a linear regression (Warton et al. 2006). The major axis is the line that minimises the sum of squares of the shortest distances from the data points to the line. Log-transformed data were used to study the relationships between both variables (see Packard 2018 and Pélabon et al. 2018 for a discussion on the use of log-transformed data in allometry studies). Significance of the correlation for each group was tested using an analysis of variance (ANOVA) under a α-level of 0.05. This analysis was not performed for Mayrocaris sp. and Austriocaris secretanae sp. nov. as the number of specimens available for measurements was no more than two for each species.

The allometry between shield height (Hmax) and shield length (Ls) for Paraostenia voultensis was studied in more details as the qualitative morphological description (see Systematic Palaeontology) distinguished two morphotypes. Comparison between elevation and slope of both morphotypes was respectively done using a Wald test (Rao 1973) and a likelihood ratio test (Flury 1984) for α = 0.05.

Complex allometry
To have a complete view of the morphological variability of Paraostenia voultensis, we studied the complex allometry using a principal component analysis (PCA). The first principal component (PC1) is considered an estimate of the allometric line as it represents the line of best fit under a least-square criterion for multivariate dataset (Jolicoeur 1963). Log-transformed data were used for this analysis. Not all measures being assessable on all the specimens due to preservation, we applied an iterative PCA method, also known as EM-PCA algorithm (Kiers 1997), to estimate the missing values for the PCA. The Burnaby back-projection method was applied to remove the effect of size on the data (Burnaby 1966).

Estimation of missing values was done with the R package missMDA ver. 1.18 (Josse & Husson 2016). Burnaby back-projection method algorithm developed by Blankers et al. (2012) and modified by Eberle et al. (2014) was used for the size correction.

The PCA was performed with the R package FactoMineR ver. 2.4 (Lê et al. 2008) and the results were plotted using R packages factoextra ver. 1.0.7 (Kassambara & Mundt 2019) and cowplot ver. 1.1.1 (Wilke 2019). Table of eigenvalues and factor loadings was done with gridExtra ver. 2.3 (Auguie 2017).

![Fig. 2. Scheme of measurements for thylacocephalans. A. Shield. B. Raptorial appendages. Abbreviations: Aad = anterodorsal angle; Aav = anteroventral angle; Apd = posterodorsal angle; Apv = posteroventral angle; Ha = anterior shield height; Hmax = maximum shield height; Hp = posterior shield height; Ln = length of podomere (appendages); Lr = length of the rostrum; Ls = length of the shield; Lw = length of the shield without the rostrum; w_n = width of podomere n (appendages).](image-url)
Table 2 (continued on next page). Descriptive statistics on the shield measurements. Abbreviations: Aad = anterodorsal angle; Aav = anteroventral angle; Apd = posterdorsal angle; Apv = posteroventral angle; Ha = anterior shield height; Hmax = maximum shield height; Hp = posterior shield height; Inf = unknown data; Lr = length of the rostrum; Ls = length of the shield; Lw = length of the shield without the rostrum; N = number of measured specimens; NA = non-applicable; SD = standard deviation.

<table>
<thead>
<tr>
<th>Measurements</th>
<th>Austriocaris secretanae sp. nov. (N = 4)</th>
<th>Kilianicaris lerichei (N = 15)</th>
<th>Mayrocaris sp. (N = 2)</th>
<th>Ostenocaris ribeti (N = 8)</th>
<th>Paraclausocaris harpa gen. et sp. nov. (N = 9)</th>
<th>Paraostenia voultensis (N = 44)</th>
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</thead>
<tbody>
<tr>
<td>Ls (mm)</td>
<td>Mean (SD) 66.29 (47.98)</td>
<td>37.27 (8.69)</td>
<td>18.27 (1.25)</td>
<td>17.16 (5.46)</td>
<td>33.73 (12.34)</td>
<td>24.27 (7.76)</td>
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<td>Lr (mm)</td>
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<td>13.96 (14.65)</td>
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<td>8</td>
<td>9</td>
<td>44</td>
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<tr>
<td>Lw (mm)</td>
<td>Mean (SD) 27.38 (NA)</td>
<td>19.18 (12.46)</td>
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<td>NA (NA)</td>
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<td>Ha (mm)</td>
<td>Mean (SD) 13.22 (8.38)</td>
<td>10.76 (3.46)</td>
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<td>7.64 (1.82)</td>
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<tr>
<td>Hmax (mm)</td>
<td>Mean (SD) 29.03 (11.91)</td>
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<tr>
<td>Hp (mm)</td>
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<td>6.19 (1.21)</td>
<td>4.10 (0.34)</td>
<td>2.66 (0.70)</td>
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Table 2 (continued). Descriptive statistics on the shield measurements. Abbreviations: Aad = anterodorsal angle; Aav = anteroventral angle; Apd = posterodorsal angle; Apv = posteroventral angle; Ha = anterior shield height; Hmax = maximum shield height; Hp = posterior shield height; Inf = unknown data; Lr = length of the rostrum; Ls = length of the shield; Lw = length of the shield without the rostrum; N = number of measured specimens; NA = non-applicable; SD = standard deviation.

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<th>Austriocaris secretanae sp. nov. (N = 4)</th>
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<th>Mayrocaris sp. (N = 2)</th>
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<th>Paraostenia voutensis (N = 44)</th>
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<tbody>
<tr>
<td>Aav (°)</td>
<td>Mean (SD)</td>
<td>108.10 (7.17)</td>
<td>140.07 (5.38)</td>
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<td>Mean (SD)</td>
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Clustering analyses
In addition to the SMA to compare the allometry of the two morphotypes of Paraostenia voultensis, we ran a K-means analysis to see if the morphotypes can be recovered using a quantitative morphological method. K-means is a partitioning clustering analysis in which a dataset is separated into a set of K groups, where K is specified by the analyst (here K = 3). Samples are classified in multiple groups, using the minimisation of the within-cluster sum-of-squares (also known as inertia). Each group is represented by its centroid, which corresponds to the means of the samples assigned to the group. Prior to the analysis, we analysed the clustering tendency of the dataset using the Hopkins statistic and the visual assessment of cluster tendency method using the factoextra package. The optimal number of clusters was assessed using the NbClust function from the R package NbClust ver. 3.0 (Charrad et al. 2014). The K-means analysis was run with functions implemented in the R environment (R Core Team 2021).

Results
Systematic palaeontology

Class Thylacocephala Pinna, Arduini, Pesarini & Teruzzi, 1982

Original diagnosis (Schram 1990)
These are arthropods with a small to large bilobed carapace enclosing the entire body. Compound eyes very well developed as either large, sessile bodies situated in optic notches or as organs that virtually cover the surface of a protrudent, sac-like cephalon. Body appendages are of two types: anteriorly 3 pairs of subchelate, raptorial limbs, and posteriorly a set of from 8 to 16 pairs of apparently paddle-like limbs. Eight sets of well-developed gills are associated with the subchelate limbs. The posterior trunk is marked with pronounced pleurites, probably developed internally as an endorphragmal skeleton, associated with the posterior limbs.

Remarks
We decide not to follow the attempt of classification proposed by Schram (2014) as it has been shown to present many issues (e.g., polyphyly of Concavicarida Briggs & Rolfe, 1983 and Conchyliocarida Secrétan, 1983, problems with the definition of autapomorphies for many taxa; see Hegna et al. 2014 and Laville et al. 2021a, 2021b for detailed explanations).

Type species
Austriocaris carinata Glaessner, 1931 by monotypy.

Original diagnosis (Rolfe 1969)
Carapace only known, posterior excavation arcuate or irregular, anterior margin concave or produced into small beak; posteroventral or ventromedian V-shaped furrow; surface smooth with mid-dorsal granulations or posteroventral pits, or dorso-ventrally striate.

Emended diagnosis
Thylacocephala with an optic notch occupying only the ventral part of the anterior margin, presence of a straight mid-dorsal line and a sharp postero-ventral corner.

Remarks
Rolfe (1969) inversed the orientation of the shield of Austriocaris, the posterior being identified as the anterior. This influenced his diagnosis, with the optic notch being described as a posterior notch.
Austriocaris secretanae sp. nov.
urn:lsid:zoobank.org:act:1AE5E1E4-F48D-4D36-BD5B-3CDC7225DE95
Figs 3–4

Dollocaris ingens – Secrétan 1985: 381–386, fig. 5a–b. — Charbonnier et al. 2010: 119, fig. 6d; 2014: 375, fig. 4a.

Diagnosis
Austriocaris with stout and long raptorial appendages, the third pair having spines on the second and third podomeres of the distal part, and a posterior trunk consisting of 12 segments bearing rod-like appendages. Absence of ventro-lateral structure.

Etymology
In reference to Sylvie Secrétan (CNRS, Paris), for her important contribution to the study of La Voult-sur-Rhône thylacocephalans.

Material examined
Holotype
FRANCE • La Boissine, La Voult-sur-Rhône, Ardèche; Callovian, Middle Jurassic; MNHN.F.R50963.

Paratypes
FRANCE • 6 specs; same collection data as for holotype; MNHN.F.A29249, A29257, A29286, A84029, R06203, R50971 • 3 specs; same collection data as for holotype; MSNM i7591, i13276, i13603.

Measurements
See Table 2.

Description
Shield morphology
Sub-oval shield in lateral view with a short, sinuous anterior margin, a sharp antero-dorsal corner, a straight and horizontal dorsal midline, endings posteriorly in a sharp postero-dorsal corner, a posterior margin divided into a very short and straight dorsal part antero-ventrally oriented and a straight, sub-vertical ventral part, a rounded postero-ventral corner, a vertical margin divided into a posterior half, steeply descending antero-ventrally and a horizontal anterior part merging approximately at the posterior third of the length, a rounded antero-ventral corner (Fig. 3A–D). Rim of all free margins delimiting a marginal fold.

Shield macro-ornamentation
Shield with a dorso-lateral carina located on its anterior area and a medio-lateral carina protruding from the postero-ventral angle (Fig. 3). Dorso-lateral carina horizontal until mid-length of the shield where it folds down on itself. Presence of a row of 33 pores on the top of the carina (Fig. 3E). Medio-lateral carina horizontal until mid-length then becoming antero-ventrally oriented. Rim of all free margins delimiting the marginal fold. Shield also ornamented with horseshoe-like ridges (Fig. 4A, C).

Sensory structures
Large, oval, stalked compound eyes protruding from the anterior margin and occupying the entire margin (Fig. 3H–I). Eyes tilted in the same orientation as the anterior margin, formed by hexagonal ommatidia (w = ~65 µm; l = ~70 µm, about 196 per mm²; Fig. 4A, F).
Cephalic appendages
Cephalic appendages not preserved.

Raptorial appendages
Presence of three pairs of stout raptorial appendages recognised by preservation of muscles, protruding from the anterior part of the ventral margin (Fig. 3A, C). Raptorial appendages consisting in two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages (Fig. 3H–I), the shortest, consisting of a proximal part with two rectangular podomeres: podomere 1 being 4.2 times as long as wide; podomere 2 being 4.7 times as long as wide. Distal part of the appendages formed by three rectangular podomeres: podomere 3 being 3.4 times as long as wide, bearing six spines on its medial margin; podomere 4 being 2.3 times as long as wide; podomere 5, the shortest, being 1.4 times as long as wide and ending in a cluster of three spines.

Second pair of raptorial appendages consisting of a proximal part with two podomeres (Fig. 3A, C): podomere 1 being 4.7 times as long as wide; podomere 2, the longest, being 5.3 times as long as wide. Distal part of the appendage consisting of three podomeres: podomere 3 being 3.2 times as long as wide; podomere 4 being 3.1 times as long as wide; podomere 5 being 1.5 times as long as wide and ending distally in a cluster of five spines.

Third pair of raptorial appendages, the longest, consisting of a proximal part formed by two rectangular podomeres (Figs 3A–C, 4I–J): podomere 1 being 3.2 times as long as wide; podomere 2, the longest, being 7.9 times as long as wide. Distal part consisting of three podomeres: podomere 3 being 5.2 times as long as wide; podomere 4 being 4.5 times as long as wide and bearing eight spines on its medial margin, the first two being close to the proximal margin (Fig. 4I–J), followed by two stout spines, surrounding a more gracile one, and then by three short spines and a long, stout spine. Distal part of the third raptorial appendages ending with a square-shaped podomere (podomere 5), bearing a cluster of three spines.

Oval structure
Long, oval structure protruding from the anterior part of the ventral margin, between raptorial and trunk appendages (Fig. 3A–D, H–I). Two elongate and sinuous rectangular structures visible on this feature. The nature of this structure is unknown.

Posterior trunk
Posterior trunk consisting of eleven elongate, rectangular structures (i.e., pleural parts of the segments; Figs 3A–D, 4D, G), slightly curved upward at their distal end and surrounding a long rectangular structure (i.e., tergal part of posterior trunk segments). Last pleural part thinner than the other ones. Presence of a square-shaped element (i.e., telson) at the posterior extremity of the trunk, bearing a circular structure formed by two circular parts: an outer one, delimiting the entire feature, and an inner smaller one.

Gills
Four long lamellar gills preserved under the posterior area of the shield below the medio-lateral carina (Fig. 3H–I).
Fig. 3. Austriocaris secretanae sp. nov. **A**–**F**. Holotype MNHN.F.R50963. **A**. General view (part). **B**. General view (counter-part). **C**. Drawing (part). **D**. Drawing (counter-part). **E**. Close-up of pores. **F**. Close-up of muscles of raptorial appendages. **G**. Paratype MNHN.F.A29257. **H**–**I**. Paratype MNHN.F.R06203. **H**. General view. **I**. Drawing. Abbreviations: am = anterior margin; ce = compound eye; dlc = dorso-lateral carina; dm = dorsal midline; g = gills; mlc = medio-lateral carina; os = oval structure; p1–5 = podomeres of raptorial appendages; pm = posterior margin; po = pores; RA1–3 = raptorial appendages; sp = spines; st = stalk; t = posterior trunk; TA8 = posterior trunk appendage 8; TP1–11 = pleural parts of posterior trunk; VLG = ventro-lateral groove; VM = ventral margin. Scale bars: **A**–**D**, **G**–**I** = 10 mm; **E** = 1 mm; **F** = 5 mm. Photos: **A**–**B**, **G**: L. Cazes; **E**–**F**: T. Laville. **H**: C. Haug & Joachim T. Haug.
Remarks
The specimens described above display typical characters of *Austriocaris* such as the well-developed optic notch restricted to the ventral part of the anterior margin, the large triangular rostrum, a straight dorsal mid-line ending in a spinous postero-dorsal corner. They differ from the type species, *Austriocaris carinata*, by the hook-shaped and tuberculate dorso-lateral carina, by the presence of horseshoe-like ridges and by the absence of a Y-shaped ventro-lateral carina. Thus, we consider these specimens as belonging to a new species: *Austriocaris secretanae* sp. nov.

Genus *Kilianicaris* Van Straelen, 1923

Type species
*Kilianicaris lerichei* Van Straelen, 1923 by monotypy.

Original diagnosis (literal translation from Van Straelen 1923a)
The overall shape is more angular, sub-quadratic in the posterior region and with a large rostrum expressed anteriorly. Tergal carina rises, especially in the rostral region. Lateral carinas are quite low. The rostrum is proportionally wider than long. The spiny posterior process is poorly delineated. Likewise for the postero-lateral processes that are slightly expressed.

Emended diagnosis
Thylacocephala with a rounded shield in lateral view, a rounded rostrum, a convex posterior part of the ventral margin, a falciform, tuberculate, dorso-lateral carina, eleven styliform, posterior trunk appendages, including elongate terminal trunk appendages.

Remarks
Schram (2014) suggested that *Kilianicaris* might be related to *Ostenocaris* Arduini, Pinna & Teruzzi, 1984, forming the Ostenocarididae Schram, 2014. He proposed that both taxa “have fused the posterior-most 8 trunk somites of the series of 16 into a posteriorly directed lobe”. This lobe corresponds to the putative telson that we describe in most of La Voult-sur-Rhône taxa (see below) and is therefore not only present in *Kilianicaris* and *Ostenocaris*. In addition to this misconception, many morphological differences exist between both taxa. As pointed out by Schram (2014), *Ostenocaris* lacks a rostrum while *Kilianicaris* has a rounded rostrum. Moreover, they present differences in the overall shape of the shield (trapezoidal for *Ostenocaris* vs rounded for *Kilianicaris*), in the morphology of their anterior margin (proverse vs straight), in the ornamentation of their dorsal margin (serrate and carinate in *Kilianicaris*), in the morphology of their dorso-lateral carina and in the number of posterior trunk appendages (9 vs 11).

*Kilianicaris lerichei* Van Straelen, 1923

Figs 5B, 6–9


Material examined

Holotype
FRANCE • La Boissine, La Voulte-sur-Rhône, Ardèche, France; Callovian, Middle Jurassic; OSUG-UJF-ID.11545.

Paratype
FRANCE • same collection data as for holotype; OSUG-UJF-ID.1752.

Additional material
FRANCE • 48 specs; same collection data as for holotype; MNHN.F.A29246, A29253, A29301 to A293016, A29318, A29374, A29375, A53334, A53335, A53338, A53339, A53342, A83991 to A85013, A84073, R50942, R50961, R50967, R50969, R50972, R50973, R50975, R50976 • 1 spec.; same collection data as for holotype; OSUG-UJF-ID.1751 • 3 specs; same collection data as for holotype; MSNM i7593, i7596, i20704.

Measurements
See Table 2.

Description

Shield morphology
Rounded shield in lateral view with a straight and sub-vertical anterior margin, an antero-dorsal corner expressed as a large, rounded and horizontal rostrum, a slightly convex dorsal midline bearing a dorsal carina in its posterior third (Fig. 6). Presence of serrations along the first two thirds of the dorsal midline, until the dorsal carina (Fig. 6H–I). Postero-dorsal corner expressed as a small spine; straight and vertical posterior margin ending in a rounded postero-ventral corner; ventral margin divided into a posterior half steeply descending antero-ventrally and an anterior one ascending antero-dorsally; both parts merged approximately at mid-length; rounded antero-ventral corner. Rim of all free margins forming a marginal fold.

Shield macro-ornamentation
Dorso-lateral carina protruding from the posterior margin, running straight until the anterior third of the shield and forming a depression (Figs 6–7); carina then ascending to the dorsal midline and merging with the dorso-lateral carina from the other side of the shield; carina associated with a row of at least 19 pores (d = ~0.5 mm) in its posterior part (Fig. 8H–K). Marginal fold delimited by a ventro-lateral groove (Fig. 6C–D).

Sensory structures
Oval, stalked compound eyes protruding from the anterior margin and occupying the entire margin (Figs 7F–G, 9E, G).

Cephalic appendages
Cephalic appendages not preserved.

Raptorial appendages
Presence of three pairs of raptorial appendages recognised by preservation of muscles, protruding from the ventral margin (Figs 9H–I, 11). Raptorial appendages consisting of two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages (Fig. 9A–D), the shortest, consisting of a proximal part with two podomeres: a rectangular podomere 1 being 3.4 times as long as wide; a stout podomere 2 being 5.9 times
as long as wide. Distal part of the appendages formed by three podomeres: podomere 3 being 4 times as long as wide; podomere 4 being 5.6 times as long as wide and bearing two medial spines; podomere 5, the most distal and shortest one, being 4.8 times as long as wide.

Second pair of raptorial appendages consisting of a proximal part with two podomeres (Fig. 9): podomere 1 not well preserved; podomere 2 being 4.7 times as long as wide (Fig. 9A–B, D). Distal part of the appendage consisting of three podomeres: podomere 3, the longest, being 5.2 times as long as wide and bearing two medial spines; podomere 4 being 5.3 times as long as wide and bearing four medial spines; podomere 5 being 3.5 times as long as wide and ending distally in a cluster of five long spines, the medial one being the smallest.

Third pair of raptorial appendages, the longest, consisting of a proximal part with two poorly preserved rectangular podomeres (Fig. 9). Distal part consisting of three podomeres: podomeres 3 being 4.8 times as long as wide; podomere 4 being 6.1 times as long as wide; podomere 5 being 5.3 times as long as wide and ending in a cluster of two spines.

**Oval structure**
Long oval structure, protruding from the anterior part of the ventral margin, between raptorial and trunk appendages (Fig. 9E–F).

![Fig. 5. Scatter plots of length of the shield against maximal height of the shield. A. Kilianicaris lerichei Van Straelen, 1923. B. Ostenocaris ribeti (Secrétan, 1985) comb. nov. C. Paraclausocaris harpa gen. et sp. nov. D. Paraostenia voultensis Secrétan, 1985. Abbreviations: Hmax = maximal shield height; Ls = length of the shield.](image-url)
Posterior trunk

Eleven, elongate rectangular structures (i.e., pleural part of posterior trunk segments) surrounding a long rectangular structure (i.e., possible tergal part; Fig. 8A–D). Pleural parts slightly curved downward at their distal end (Fig. 8D). Last pleura thinner than the others. Posterior extremity of the trunk expressed as a rhomboidal element (i.e., telson) bearing a rounded structure, formed by two circular parts (Fig. 6C–D): an outer one, delimiting the entire feature, and an inner smaller one.

**Fig. 8. Kilianicaris lerichei** Van Straelen, 1923. **A–B.** Specimen MNHN.F.A29375. **A.** General view. **B.** Drawing. **C–D.** Specimen MNHN.F.A29309. **C.** General view. **D.** Close-up of posterior trunk. **E–G.** Specimen MNHN.F.A29246. **E.** General view. **F.** Close-up of oval structure. **G.** Close-up of posterior trunk. **H–K.** Specimen MNHN.F.A29374. **H.** General view. **I.** Drawing. **J.** Close-up of dorso-lateral carina. **K.** Close-up of pores. Abbreviations: am = anterior margin; b = body; dlc = dorso-lateral carina; dm = dorsal midline; pm = posterior margin; ra₂ = raptorial appendage 2; tls = telson; tp₁₋₁₁ = pleural part of posterior trunk segments; tta = terminal trunk appendages; ttaen = endopod of terminal trunk appendages; ttaex = exopod of terminal trunk appendages; vlg = ventro-lateral groove; vm = ventral margin. Scale bars: A–C, E, H–I = 10 mm; D, G = 5 mm; F, J = 4 mm; K = 1 mm. Photos: T. Laville.
Posterior trunk appendages
Eleven pairs of trunk appendages, postero-dorsally oriented, protruding from the posterior part of the ventral margin (Fig. 8). First ten appendages formed by two parts: a square-shaped proximal one with a rounded distal margin and a styliform distal one (Fig. 6E). Last pair of trunk appendages formed by two paddle-like parts and being longer than the rest (Fig. 8A–B).

Gills
Eight long lamellar gills preserved under the central part of the shield below the dorso-lateral carina (Fig. 6H–I). Gills with an elliptic shape, becoming arcuate near the dorsal midline.

Genus *Mayrocaris* Polz, 1994

Type species
*Mayrocaris bucculata* Polz, 1994 by monotypy.

Original diagnosis (literal translation from Polz 1994)
Carapace subtrapezoid, without a clearly delineated optic notch, however, with a pronounced subquadratic notch; ventral line from here slopping down at 45° angle, almost equally ascending from the middle of the carapace, forming, together with the lowest part of the anterior margin, a characteristic U-shaped bulge; hinge line on the whole slightly convex, terminating in a pointed spine; rostrum inconspicuous; carapace surface roughly sculptured only by a short horizontal, very delicate ridge, and by eight small bases of spines, micro-relief in the form of single, very small arched lines on an ultrafine faveolated surface; anterior lobes larger than in comparable genera, protruding from the carapace not only anteriorly, but even projecting over its antero-ventral section; abdomen segmented. Kind of large appendages not clausocarid, course of anterior carapace margin conchyliocarid.

Emended diagnosis
Thylacocephala with a slightly developed optic notch, a serrate dorsal margin consisting of a carina, a rounded postero-dorsal spine, a posterior notch, and an anterior part of the ventral margin oriented antero-dorsally and bearing a square notch.

*Mayrocaris* sp.

![Fig. 10](image)

*Dollocaris* sp. – Secrétan & Riou 1983: text-fig. 14c.

Material examined
FRANCE • 11 specs; La Boissine, La Voult-sur-Rhône, Ardèche, France; Callovian, Middle Jurassic; MNHN.F.A29366, A84014 to A84022, R50964.

Measurements
See Table 2.

Description
Shield morphology
Sub-trapezoidal shield in lateral view with an anterior margin formed by a weakly developed optic notch bent in its central part, a sharp antero-dorsal corner, a slightly convex dorsal midline ascending until a carina located in the last third of the margin (Fig. 10); dorsal midline with small serrations in the anterior part of the midline; posterior margin consisting of a notch delimited dorsally by a postero-dorsal corner
expressed as a spine with a rounded tip and ventrally by a rounded postero-ventral corner; ventral margin divided into a steep posterior half descending antero-ventrally and an anterior one slightly ascending antero-dorsally and bearing a pronounced subquadratic notch; ventral margin ending anteriorly as a rounded antero-ventral corner. Rim of all free margins delimiting a marginal fold.

**Shield macro-ornamentation**
Presence of a medio-lateral carina in the central part of the shield, above the gills (Fig. 10A–D). One rounded pore, formed by a canal-like structure, sitting close to the anterior end of the carina (Fig. 11A–D).

**Sensory structures**
Oval, stalked compound eyes protruding from the optic notch and occupying all of it (Fig. 10).

**Cephalic appendages**
One antenniform structure protruding from the anterior margin of the shield, becoming thinner distally (Fig. 10I–J). Three rectangular elements preserved distally.

**Raptorial appendages**
Presence of three pairs of raptorial appendages recognised by preservation of muscles, protruding from the ventral margin, posterior to the presumed cephalic appendages. Raptorial appendages consisting of two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages protruding from the ventral notch (Fig. 10E–F). Proximal part formed by two rectangular podomeres (podomeres 1 and 2). Distal part consists of at least one podomere (podomere 3).

Second pair of raptorial appendages consisting of a proximal part with at least two rectangular podomeres (Fig. 10E–F). Distal part of the second raptorial appendages ending distally in a cluster of three spines.

Third pair of raptorial appendages ending distally in a cluster of five spines (Fig. 10E–F).

**Posterior trunk**
Long, tubular-like structure consisting of longitudinal muscles, extending from the anterior part of the shield to the posterior notch, interpreted as part of the posterior trunk (Fig. 10A–D). Nine trunk segments preserved under the shield, consisting of a rectangular pleural part, slightly curved downward at its distal end (Fig. 10H). Pleural parts decreasing in length toward the last segment. Presence of a circular structure at the posterior extremity of the trunk (Fig. 10A–D).

**Posterior trunk appendages**
Remnants of nine pairs of trunk appendages protruding from the posterior part of the ventral margin; appendages postero-dorsally oriented and poorly preserved, preventing their morphology from being determined (Fig. 10A–D).

**Gills**
Eight lamellar gills preserved under the central part of the shield, below the lateral carina (Fig. 10A–D). Gills becoming more arcuate toward the posterior part of the shield.

**Anterior muscles**
Presence of a bundle of radially arranged muscles close to the antero-ventral corner of the shield (Fig. 10A–D).
Remarks
The morphology of Mayrocaris sp. from La Voult-sur-Rhône Lagerstätte is similar to that of Mayrocaris bucculata from Solnhofen-type Lagerstätten (Laville et al. 2021a): presence of a carina and of serrations on the dorsal midline, postero-dorsal corner expressed as a rounded spine, posterior and ventral notches, morphology and orientation of trunk segments, raptorial and trunk appendages, and of gills. The main difference is the ornamentation: no scale-like structures are visible on the shield of the La Voult specimens. However, this might be due to poor preservation. Until better-preserved specimens are discovered, these specimens cannot be unambiguously assigned to a particular species.

Genus Ostenocaris Arduini, Pinna & Teruzzi, 1984

Type species
Ostenia cypriformis Arduini, Pinna & Teruzzi, 1980 by monotypy.

Diagnosis
Thylacocephalan with a sinuous anterior margin ending in a sharp antero-dorsal corner. Large oval eyes protruding from the anterior margin. Medio and dorso-lateral carina present on the shield. Large raptorial appendages bearing short spines.

Other included species
Ostenocaris ribeti (Secrétan, 1985) comb. nov. – Callovian, Middle Jurassic, France.

Remarks
Arduini et al (1980) did not provide a diagnosis for Ostenocaris. They just mentioned that the description of Ostenocaris was “coinciding with that of the type species” (Arduini et al. 1980: 362). Concerning the type species, they just described the original species without providing a proper diagnosis.

Ostenocaris ribeti (Secrétan, 1985) comb. nov.
Figs 5C, 11–12

Clausia ribeti Secrétan, 1985: 388, fig. 5d, i–j.
Clausia sp. – Secrétan & Riou 1983: 85, pl. IV figs 2, 4–5.
Clausia ribeti – Arduini 1992: 266.

Original diagnosis (Secrétan 1985)
The new species differs from the type species Clausia lithographica Oppenheim in its much stronger anterior appendages. Like the type species, C. ribeti is small, has a rounded carapace and large eye with oval lateral aspect. The appendages, however, have very fine paintbrush like extremities and are very much shorter and very much more massive than these of C. lithographica.

Emended diagnosis
Ostenocaris with a tuberculate, club-shaped dorso-lateral carina and hook-shaped first raptorial appendages.

Material examined
Holotype
FRANCE • La Boissine, La Voult-sur-Rhône, Ardèche, France; Callovian, Middle Jurassic; MNHN.F.R50981.
Additional material
FRANCE • 11 specs; same collection data as for holotype; MNHN.F.A29349, A29365, A29368, A29372, A84023 to 84028, R50970.

Measurements
See Table 2.

Description

Shield morphology
Sub-oval shield in lateral view with a sinuous and proverse anterior margin, a sharp, antero-dorsal corner, a convex dorsal midline with a short flattening near the rounded postero-dorsal corner, a straight and vertical posterior margin, a rounded postero-ventral corner a ventral margin consists of a posterior half steeply descending antero-ventrally and of a horizontal anterior one (Figs 11–12). Rim of all free margins delimiting a marginal fold.

Shield macro-ornamentation
Shield adorned by two large, tuberculate carinae on its lateral sides: a club-shaped horizontal dorso-lateral carina and a sinuous medio-lateral carina (Figs 11A, 12A–C, G).

Sensory structures
Two oval, sessile compound eyes protruding from the optic notch, occupying most (Fig. 11).

Cephalic appendages
One elongate structure, consisting of at least three elements, lying in front of the raptorial appendages (Fig. 12A–C).

Raptorial appendages
Presence of three pairs of raptorial appendages recognised by preservation of muscles, protruding from the ventral margin, posterior to the presumed cephalic appendages (Fig. 12A–D). Raptorial appendages consisting of two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages, the shortest, having a hook-shaped morphology (Fig. 12D). Proximal part consisting of two rectangular podomeres: podomere 1 being 2.5 times as long as wide; podomere 2 being 8 times as long as wide. Distal part formed by three rectangular podomeres: podomere 3 being 3.4 times as long as wide; podomere 4 being 3.4 times as long as wide; podomere 5 being 1.7 times as long as wide and ending in a spiny tip.

Second pair of raptorial appendages consisting of a proximal part with two rectangular podomeres (Fig. 12D): podomere 1 not being well preserved; podomere 2 being 4.6 times as long as wide. Distal part of the appendage consisting of three podomeres: podomere 3 being 3.8 times as long as wide; podomere 4 being 5 times as long as wide; podomere 5 being 2.7 times as long as wide and ending in a spiny tip.

Third pair of raptorial appendages, the longest, consisting of a proximal part with two long rectangular podomeres (Fig. 12D): podomere 1 being 1.8 times as long as wide; podomeres 2 being 5.2 times as long as wide. Distal part consisting of three podomeres: podomere 3 being 4.7 times as long as wide; podomere 4 being 5 times as long as wide and bearing a row of spine on its medial and lateral sides; podomere 5 being slightly curved and 2.9 times as long as wide.
**Posterior trunk**

Presence of seven rectangular posterior trunk segments, with only three rectangular pleural parts preserved (Fig. 11F–G). Trunk appendages not preserved.

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**Fig. 11. Ostenocaris ribeti** (Secrétan, 1985) comb. nov.  
A. Holotype MNHN.F.R50981.  
B. Specimen MNHN.F.A29349.  
C. Specimen MNHN.F.A29368.  
D–E. Specimen MNHN.F.A84023.  
D. General view.  
E. Drawing.  
F–G. Specimen MNHN.F.A29365.  
F. General view.  
G. Drawing. 
Abbreviations: am = anterior margin; b = body; ce = compound eye; dlc = dorso-lateral carina; dm = dorsal midline; g = gills; mlc = medio-lateral carina; p₁₋₅ = podomeres of raptorial appendages; ra₂–₃ = raptorial appendages; tp₁₋₇ = pleural part of the posterior trunk; vm = ventral margin. Scale bars: 5 mm. Photos: A: J. Falconnet (RECOLNAT – MNHN); B–C, F: T. Laville; D: L. Cazes.
Gills
Six oval, short gills located below the median part of the shield (Fig. 11F–G).

Remarks
New observations of the holotype of *Clausocaris ribeti* indicate that this species has the general morphology of *Ostenocaris*: a proverse anterior margin with a sharp antero-dorsal angle, a convex dorsal midline, a straight posterior margin, a marginal fold on all free margins, presence of rows of spines on

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the medial and lateral end of podomere 4 of the third pair of raptorial appendages. This species also displays major differences with *Clausocaris*: absence of an optic notch and of serrations on the dorsal margin, presence of a posterior margin, differentiation of the ventral margin into a posterior and an anterior parts, short and stout raptorial appendages. Thus, we propose the new combination *Ostenocaris ribeti* (Secrétan, 1985) comb. nov.

Here, it is also important to note that the paratype of *C. ribetii* (MNHN.F.R50978) is no longer considered to be a representative of this species. Based on its morphology, it is now considered to belong to *Paraostenia voultenis*. As pointed out by Laville et al. (2021c), there has been a lot of confusion between *Paraostenia* Secrétan, 1985 and *Ostenocaris* due to the wrong assignment of specimens of *Paraostenia* from the Osteno Lagerstätte, Italy, to *Ostenocaris* (see Arduini et al. 1980). Both taxa differ from each other based on the morphology of their anterior margin (sinuous for *Ostenocaris* vs large optic notch for *Paraostenia*), of their dorsal margin (presence of carina in *Paraostenia*) and of their posterior margin (straight vs concave).

*Ostenocaris ribeti* is the second species of *Ostenocaris* described. The main difference with the type species is the size: *O. ribeti* is only 1.7 cm long while *O. cypriformis* can reach a length of 20 cm. *Ostenocaris ribeti* also differs from *O. cypriformis* by its large, tuberculate and elongate dorso-lateral and medio-lateral carinae and its hook-shaped first raptorial appendages. Its dorsal midline also appears less convex.

**Genus *Paraclausocaris* gen. nov.**

*Paraclausocaris* harpa gen. et sp. nov. by monotypy.

**Diagnosis**

Thylacocephala with a well-developed optic notch restricted to the ventral margin, a sharp antero-ventral angle, small sinuous ridges on the shield and raptorial appendages 3 having spines on the second and third podomeres of the distal part.

**Etymology**

From the Latin ‘par, paris’ ('close to’) and *Clausocaris* Polz, 1989.

**Remarks**

*Paraclausocaris* gen. nov. has many similarities with *Clausocaris*: a cone-shaped shield in lateral view, a dorsal midline with sharp, short serrations and long, thin raptorial appendages relative to the shield. However, it also displays some differences, including a well-developed posterior margin and an anterior margin with an optic notch limited to the ventral part of the margin for *Paraclausocaris*, while in *Clausocaris* the posterior margin is absent and the optic notch occupies the whole margin. As a result of these differences, we erect *Paraclausocaris*.

One last morphological trait should also be discussed here: the number of posterior trunk appendages. In *Paraclausocaris* gen. nov., we recognize nine pairs of posterior trunk appendages. Arduini (1992) described eight posterior trunk appendages in *Clausocaris pinnai*, but based on pictures from Dalla Vecchia (2012), there may actually be nine as in *Paraclausocaris*. However, it is different from the number reported by Polz (1990) or Haug et al. (2014) for *Clausocaris lithographica*. They respectively reported 10 or 11 posterior trunk appendages for this species. Interspecific variation in the number of
posterior trunk appendages is not uncommon among Thylacocephala (Schram 2014). However, it is quite unusual to find such variation among closely related forms: for instance, eight posterior trunk appendages are usually reported in Concavicaris species (Schram 2014). Thus, it will be necessary to revise C. pinnai to confirm its assignment to Clausocaris.

**Paraclausocaris harpa** gen. et sp. nov.

*urn:lsid:zoobank.org:act:4F007D44-251E-496D-818B-9C3065947B0D*

Figs 5A, 13–15

*Clausia* sp. – Secrétan & Riou 1983: 85–86, pl. 4 figs 2, 17.

*Clausia ribeti* – Secrétan 1985: 388, fig. 5j.

*Clausocaris ribeti* – Polz 1989: 73. — Charbonnier 2009: 158, fig. 250. — Charbonnier et al. 2010: 118, fig. 5d; 2014: 375, fig. 5c.

*Dollocaris ingens* – Charbonnier et al. 2010: 113, fig. 3e.

**Diagnosis**

Same as for *Paraclausocaris*.

**Etymology**

Referring to the shape of the shield, which is similar to the shape of a harp.

**Material examined**

**Holotype**

FRANCE • La Boissine, La Voult-sur-Rhône, Ardèche; Callovian, Middle Jurassic; MNHN.F.A29295.

**Paratypes**

FRANCE • 22 specs; same collection data as for holotype; MNHN.F.A29148, A29149, A29264, A29267 A29295, A29323, A29327, A29345, A29348, A29353, A29356, A29358, A29360, A53341, A83986 to A83990, A84072, R50955, R50974.

**Measurements**

See Table 2.

**Description**

**Shield morphology**

Cone-shaped shield in lateral view with an anterior margin formed by a symmetric and concave optic notch in its ventral part and with a convex dorsal part, a sharp antero-dorsal corner, a slightly convex dorsal midline with small serrations in the last two thirds of the midline and ending in a sharp postero-dorsal corner (Figs 13A, C, H, 14G–H); straight posterior margin antero-ventrally oriented, ending in a rounded postero-ventral corner; ventral margin divided in a concave posterior part steeply descending antero-ventrally and in a horizontal anterior part slightly rising to the antero-ventral corner. Rims of all free margins forming a marginal fold.

**Shield macro-ornamentation**

Small sinuous ridges covering the entire dorsal area of the shield (Fig. 13F). Central part of the shield adorned with a straight dorso-lateral carina (Fig. 14A–B). Shield forming a hump near the antero-ventral corner (Fig. 14A–B). Thin ventro-lateral groove delimiting the marginal fold. Four rounded, canal-like pores visible in the posterior part of the shield, above the pleural part of trunk segments (Fig. 13F–G). Pores forming a rim surrounding a hollow part.
Sensory structures
Two stalked, circular compound eyes protruding from the optic notch and occupying most of the anterior margin (Fig. 14A–B). Eyes consisting of 136 hexagonal ommatidia per mm² (w = ~73 µm; h = ~130 µm; Figs 13E, 14E, 15H).

Cephalic appendages
One elongate, antenniform structure (~15 mm) lying in the anterior part of the body (Figs 13A, C, 14C–D).

Raptorial appendages
Three pairs of long and slender raptorial appendages recognised by the preservation of muscles, protruding from the ventral margin, posterior to the presumed cephalic appendages. Raptorial appendages consisting of two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages (Figs 13H–I, 15F–G), the shortest, consisting of a proximal part with two podomeres: podomere 1 being 3.4 times as long as wide; podomere 2 being 4.6 times as long as wide and longer than podomere 1. Distal part of the appendages formed by three podomeres: podomeres 3 being 4.3 times as long as wide; podomere 4 being 3.3 times as long as wide; podomere 5, the most distal one, being 3 times as long as wide, ending in a cluster of three long spines.

Second pair of raptorial appendages consisting of a proximal part with two podomeres: podomere 1 being 3.4 times as long as wide, podomere 2 being 6.2 times as long as wide (Figs 14A–B, 15F, I). Distal part of the appendage consisting of three podomeres: podomere 3 being 4.7 times as long as wide; podomere 4 being 5.5 times as long as wide; podomere 5, being 3.2 times as long as wide and ending distally in a cluster of three long spines.

Third pair of raptorial appendages, the longest, consists of a proximal part formed by two podomeres: podomere 1 being 3.6 times as long as wide; podomere 2, the longest of the appendages, being 6 times as long as wide (Fig. 15A, F–G). Distal part consisting of three podomeres: podomere 3 being 4.4 times as long as wide; podomere 4 being 5.7 times as long as wide and bearing three long spines on its medial end; podomere 5 being 3.3 times as long as wide and ending with a cluster of three long spines.

Oval structure
Long oval structure, protruding from the anterior part of the ventral margin, posterior to raptorial appendages (Figs 13H–I, 15A, C, G–H).

Posterior trunk
Posterior trunk consists of nine rectangular segments, slightly tilted anteriorly. Segments consisting of a dorsal square-shaped structure (i.e., tergal part of the segments), surrounded on their lateral sides by nine elongate rectangular structures (i.e., pleural part; Fig. 15G–H, J), slightly curved downward at their distal end (Fig. 15A, D). Segments increasing in height towards the penultimate segment. Most posterior segment shorter than the previous one.

Posterior trunk ending posteriorly in a rhomboidal element (i.e., telson), bearing a circular structure formed by two circular parts (Fig. 14A–D): an outer one, delimiting the entire feature, and an inner smaller one.

Posterior trunk appendages
Nine pairs of posterior trunk appendages protruding from the posterior part of the ventral margin (Fig. 13A–D). First appendage formed by two parts: a square-shaped dorsal one (i.e., endopod) and a
Fig. 13. *Paraclausocaris harpa* gen. et sp. nov. A–G. Holotype MNHN.F.A29295. A. General view of the part. B. General view of the counter-part. C. Drawing of the part. D. Close-up of the posterior trunk. E. Close-up of cephalic appendages. F. Close-up of the ornamentation. G. Close-up of pores. H–I. Paratype MNHN.F.A29148. H. General view. I. Drawing. Abbreviations: ca = cephalic appendages; ce = compound eye; dm = dorsal midline; en = putative endopod of posterior trunk appendage; ex = putative exopod of posterior trunk appendage; g = gills; mu = muscles; on = optic notch; os = oval structure; $p_1$–5 = podomeres of raptorial appendages; po = pore; r$alpha_1$–3 = raptorial appendages; st = stalk; t = trunk; $ta_2$–8 = posterior trunk appendages; tls = telson; tp$1$–9 = pleural part of posterior trunk segments; ts$1$–7 = trunk segments; vm = ventral margin. Scale bars: A–D, F = 5 mm; E = 2 mm; G = 1 mm; H–I = 10 mm. Photos: A–B: Lilian Cazes; D–H: T. Laville.
Fig. 14. *Paraclausocaris harpa* gen. et sp. nov. A–E. Paratype MNHN.F.A29327. A. General view. B. Drawing. C. Close-up of cephalic appendage. D. Close-up of cephalic appendage (colour-marked). E. Close-up of compound eyes. F–G. Paratype MNHN.F.A29149. F. General view. G. Close-up of posterior part. Abbreviations: ca = cephalic appendage; ce = compound eye; dm = dorsal midline; dlc = dorso-lateral carina; g = gills; hu = hump; ls = left side of the shield; mu = muscles; on = optic notch; os = oval structure; \( p_{2,5} \) = podomeres of raptorial appendages; \( r_{1-3} \) = raptorial appendages; rs = right side of the shield; vlgy = ventro-lateral groove. Scale bars: A–B, G = 5 mm; C–D = 2 mm; E = 500 µm; F = 10 mm. Photos: T. Laville.
ventral oval one (i.e., exopod). Trunk appendages 2–8 consist of at least two elements: a square-shaped proximal one and a rectangular distal one. Last pairs of trunk appendages being much longer than the rest (Fig. 14F–G). No square-shaped element discernible.

**Gills**
Eight lamellar gills preserved under the central part of the shield below the lateral carina (Fig. 14A–B). Gills arcuate with an elliptic shape.

**Anterior muscles**
Bundle of radially arranged muscles preserved in the antero-medial part of the shield, above the antero-ventral hump (Fig. 13A, C).

**Remarks**
Some specimens previously assigned to *Clausocaris ribeti* (now *Ostenocaris ribeti* comb. nov.) are now ascribed to *Paraclausocaris harpa* gen. et sp. nov. (e.g., MNHN.F.A29149; Fig. 13H–I; Charbonnier et al. 2010). Indeed, they differ from the type material (and thus from *Ostenocaris*) by their well-developed optic notch, their serrated dorsal margin, their concave posterior part of the ventral margin and their elongate raptorial appendages relative to the shield size.

**Genus Paraostenia** Secrétan, 1985

**Type species**
*Paraostenia voultensis* Secrétan, 1985 by monotypy.

**Original diagnosis** (Secrétan 1985)
The voluminous and rounded eyes, the gills and the anterior and posterior appendages are as in *Dollocaris*. The form of the carapace is nearly identical to that of *Ostenia* (Arduini et al. 1980 amend Arduini & Brasca 1984) differing in form from *Dollocaris* in the outline which is proportionally higher and shorter and devoid of suborbital spine below the orbital opening which is less deep and more gaping.

**Emended diagnosis**
Thylacocephala with a symmetric optic notch, a sharp antero-dorsal corner, a dorsal carina, a concave posterior margin, rounded postero-dorsal and postero-ventral corners, long and gracile appendages.

**Other included species**

*Paraostenia voultensis* Secrétan, 1985
Figs 5D, 16–20

*Paraostenia voultensis* Secrétan, 1985: 388–389, fig. 5c–h, l.
Original diagnosis (Secrétan 1985)
As for genus.

Emended diagnosis
Paraostenia with a serrate dorsal margin and a dorso-lateral row of pores.

Material examined
Holotype
FRANCE • La Boissine, La Voulte-sur-Rhône, Ardèche; Callovian, Middle Jurassic; morphotype 2; MNHN.F.R50977.

Paratype
FRANCE • 1 spec.; same collection data as for holotype; MNHN.F.R50979.

Additional material
FRANCE • 73 specs; same collection data as for holotype; morphotype 2; MNHN.F.A29150, A29231, A29256, A29300 A29320 to A29322, A29326, A29328 to A29338, A29340, A29341, A29344, A29346, A29347, A29351, A29354, A29355, A29362, A29369, A29370, A53132, A53134, A53135, A53312, A53328 to A53330, A53336, A53340, A84005 to A84008, A84038 to A84041, A84043 to A84058, A84060 to A84063, A84065 to A84071, R50966, R50978 • 4 specs; same collection data as for holotype; morphotype 2; MSNM i7597, i18928, i25037, i25038 • 9 specs; same collection data as for holotype; morphotype 1; MNHN.F.A29339, A29354, A29357, A29361, A53131, A84039, A84040, A84047, A84055, A84059.

Measurements
See Table 2.

Description
Shield morphology
Square-shaped to sub-trapezoidal shield in lateral view with an asymmetric anterior margin formed on a well-developed optic notch, a rounded or sharp antero-dorsal corner, a convex dorsal midline is slightly convex, with a large fin-like (Fig. 18D–E) or flat dorsal carina, a rounded postero-dorsal corner, a vertical or postero-ventrally tilted, concave posterior margin, a rounded postero-ventral corner and a ventral margin divided into a posterior half steeply descending antero-ventrally and a horizontal anterior one (Figs 16, 17A–D). Rim of all free margin delimiting a marginal fold (Fig. 16B, D).

Shield macro-ornamentation
Presence of one straight and large medio-lateral carina, running along almost the entire length of the shield and divided into two parts by a longitudinal narrow furrow (Figs 16A, C, 21A–B). Presence of seven small serrations dorsally oriented on the anterior side of the dorsal carina and of seven serrations dorsally oriented, longer than the anterior ones on the posterior side (Fig. 18D–E).

At least 13 pores forming a sinuous row in the dorsal area of the shield, above the medio-lateral-carina (Figs 16A, C, 18D–F). Horseshoe-like ornamentation or field of punctuations covering the ventral area of the shield (Fig. 19A, C, J). Fields of punctuations also covering the surface of the dorsal area (Fig. 19E–G). Punctuations in the ventral area most probably correspond to degraded horseshoe-like ornamentation.

Presence of a hump either on the medio-ventral area of the shield or near the antero-ventral corner (Figs 16A, C, 19H–J).
Sensory structures
Rounded, stalked, compound eyes protruding from the optic notch and occupying most of it (Figs 16, 18B–C). Eyes formed of hexagonal ommatidia (w = ~80 µm; l = ~80 µm; about 236 per mm²; Fig. 17F).

Cephalic appendages
Antenniform structure only perceptible by the muscle fibres protruding from the optic notch and decreasing in diameter distally (Fig. 17E).

Raptorial appendages
Presence of three pairs of slender raptorial appendages recognised by the preservation of the muscles, protruding from the ventral margin, posterior to the presumed cephalic appendages. Raptorial appendages consisting of two parts: a proximal postero-ventrally oriented and a distal one, antero-ventrally oriented. Articulation between both parts forming an elbow.

First pair of raptorial appendages (Figs 16, 19A, C), the shortest, consisting of a proximal part with two podomeres: podomere 1 being 3.7 times as long as wide; podomere 2 being 4.7 times as long as wide and being longer than podomere 1. Distal part of the appendages formed by three podomeres: podomeres 3 being 4.4 times as long as wide; podomere 4 being 4.8 times as long as wide; podomere 5, the most distal one, being 3.1 times as long as wide.

Second pair of raptorial appendages consisting of a proximal part with two rectangular podomeres (Fig. 18M, O); podomere 1 not being well preserved; podomere 2 being 6.1 times as long as wide. Distal part of the appendage consisting of three podomeres: podomere 3 being 5.6 times as long as wide and bearing one spine on its medial side; podomere 4 being 5.2 times as long as wide and bearing three spines on its medial side; podomere 5 being 2.8 times as long as wide and ending distally in a cluster of three long spines.

Third pair of raptorial appendages, the longest, consisting of a proximal part formed by two podomeres, podomere 2 being 5.9 times as long as wide (Fig. 18I–K). Distal part consisting of three podomeres: podomere 3 being 4.4 times as long as wide, bearing two spines on the distal part of its medial side; podomere 4 being 6.4 times as long as wide and bearing nine spines on its medial side, with the first four arranged in pairs; podomere 5 being 4.3 times as long as wide and ending distally with a cluster of three long spines.

Oval structure
Oval structure protruding from the ventral margin, behind the last pair of raptorial appendages (Fig. 18G–H). Structure associated with a lamellar structure in one specimen (Fig. 17G).

Posterior trunk
Nine rectangular pleural parts of trunk segments protruding from the posterior part of the ventral margin (Figs 16B, D, G, 20). In some specimens, pleural part increasing in size until tp₉, then decreasing until tp₁₀(Fig. 16B, D, G). In other specimens, pleural parts decreasing in size posteriorly with last one (tp₉) thinner than the rest (Fig. 17H). Proximally, they have a postero-dorsal orientation while they are curved downward distally. Trunk ending in a square segment, i.e., the telson, bearing a rounded structure (Fig. 17I–J). Field of setae visible on the telson.

Gills
Eight arcuate, lamellar gills preserved below the medio-lateral carina (Fig. 19A–B).
**Fig. 16.** Juvenile of *Paraostenia voultensis* Secrétan, 1985 (morphotype 1). A–C. Specimen MNHN.F.A84040. A. General view. C. Drawing. B–D. Specimen MNHN.F.A29357. B. General view. D. Drawing. E. Specimen MNHN.F.A29354. F. Specimen MNHN.F.A84039. G. Specimen MNHN.F.A29261. Abbreviations: ce = compound eye; dc = dorsal carina; dm = dorsal midline; hu = hump; mlc = medio-lateral carina; on = optic notch; p\(_{1-5}\) = podomeres of raptorial appendages; pm = posterior margin; po = pore; ra\(_{2-3}\) = raptorial appendages; sp = spines; tp\(_{1-7}\) = pleural part of posterior trunk segments; vlg = ventro-lateral groove; vm = ventral margin. Scale bars = 5 mm. Photos: T. Laville.
Fig. 17. Adult of *Paraostenia voultensis* Secrétan, 1985 (morphotype 2; MNHN.F.A29300). **A**, **C–H**, **J**. Counter-part. **A**. General view. **C**. General view (colour-marked). **D**. Drawing. **E**. Close-up of cephalic appendage (colour-marked). **F**. Close-up of compound eye. **G**. Close-up of oval structure. **H**. Close-up of posterior trunk. **J**. Close-up of telson. **B**, **I**. Part. **B**. General view. **I**. Close-up of telson. Abbreviations: ca = cephalic appendage; ce = compound eye; cs = circular structure; dc = dorsal carina; dm = dorsal midline; g = gills; mlc = medio-lateral carina; on = optic notch; os = oval structure; pm = posterior margin; po = pore; ra2–3 = raptorial appendages; t = trunk; tls = telson; ta1–7 = posterior trunk appendages; tta = terminal trunk appendages. Scale bars: A–D = 10 mm; E, G–H = 5 mm; F = 500 µm; I–J = 1 mm. Photos: T. Laville.
Remarks

Paraostenia voultensis displays an important variability for some features. For instance, some specimens have a trapezoidal shield with a vertical posterior margin and a horseshoe-like ornamentation (Figs 17–20), while others have a square-shaped shield with a postero-ventrally tilted posterior margin and no ornamentation (Fig. 16). Moreover, it seems there is a slight difference in total length (Fig. 21A): the histogram shows a bimodal distribution that might indicate the presence of two different morphotypes.

Morphometry

In order to characterise the variability of Paraostenia voultensis and to confirm the presence of two morphotypes, we ran various morphometric and statistical analyses. In this section, we consider separately both P. voultensis putative morphotypes in order to test for their differences. Morphotype 1 corresponds to the small specimens with a square-shaped shield, a postero-ventrally tilted posterior margin, and no ornamentation. Morphotype 2 corresponds to bigger specimens having a square-shaped shield with a postero-ventrally tilted posterior margin and no ornamentation.

Descriptive statistics

The measurements made on Paraostenia voultensis allow us to test the homogeneity of the sample. With the exception of the length of the shield (Ls), histograms appear normally distributed for all measurements (Fig. 21A). According to the results of the Shapiro-Wilk test (W > 0.95; p > 0.13), all measurements probably follow a normal law.

Simple allometry

For Paraostenia voultensis (Fig. 21; Table 3), a strong and significant correlation has been found for morphotype 2 ($R^2 = 0.892$, p-value < 0.05) while it is non-significant for morphotype 1 ($R^2 = 0.605$, p-value = 0.222). The slope (likelihood ratio statistic equal to 0.4298, p-value = 0.5121) and the elevation (Wald statistic equal to 0.7178, p-value = 0.39687) are equal between the allometric line of both morphotypes. This indicates that there is no difference in static allometry of both morphotypes and that they are located on a similar allometric line.

For all other species, a strong and significant positive correlation has been found between shield length and shield height ($R^2 > 0.86$, p-value < 0.02; Table 3).

Complex allometry

The first two principal components account for 89.0% of the variance (Fig. 22A). PC1 accounts mostly for variation in height and angle measurements while PC2 accounts mostly for variation in shield length (Supp. file 3). Size measurements appear to be correlated between them (Fig. 22B), but not with angles measurements, which also appear correlated between them.

Table 3. Coefficients of standardised major axis analyses and dependence parameters.

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Fig. 21. Frequency distribution and simple allometry of shield measurements for *Paraostenia voultensis* Secrétan, 1985. 

A. Histograms of shield measurements. 

B. Standardised major axis (no groups). 

C. Standardised major axis (morphotypes 1 and 2). 

D. Residuals vs fitted data plot. 

E. Q-Q plot. 

Abbreviations: Aad = antero-dorsal angle; Aav = antero-ventral angle; Apd = postero-dorsal angle; Apv = postero-ventral angle; Ha = anterior height; Hmax = maximal shield height; Hp = posterior height; Ls = shield length; SMA = standardised major axis.
Clustering analysis (Paraostenia voutensis)

Based on the VAT, the dataset presents a good clustering tendency that is not confirmed by the Hopkins statistic (0.57), which indicates a low clustering tendency. An optimal number of three clusters for the K-Means analysis was determined. Results of the prediction are presented in Figure 22. Cluster 1 corresponds to specimens identified as morphotype 1 while specimens identified as morphotype 2 are split between clusters 2 and 3.

Discussion

Intraspecific variability in Paraostenia voutensis

The clustering analysis confirms that at least two morphotypes of Paraostenia voutensis co-occur in La Voulté. All specimens of morphotype 1 are part of the same cluster (1 in Fig. 22A) while specimens assigned to morphotype 2 are split into two other clusters (2–3 in Fig. 22A). There are no clear morphological differences between clusters of morphotype 2 and we thus consider these specimens as belonging to the same morphotype.

The dimorphism noticed here could have different origins: sex-related differential growth (sexual dimorphism), interspecific differential growth (interspecific variation) or ontogeny. Here, it is unlikely that the morphotypes represent two different taxa or sexual dimorphism. Indeed, in case of sexual dimorphism or interspecific variation, we would have noticed differences in allometric shape due to different growth (e.g., Fairbairn 1997; Gidasweski et al. 2009). Instead, both morphotypes share a common allometric line, which most probably represents the ontogenetic allometry of P. voutensis. This corresponds to the covariation among traits across ontogenetic stages of an individual or in a population (mean ontogenetic allometry) of a given species (Gould 1966; Klingenberg 1998; Pélabon et al. 2013). In a classic case, the various ontogenetic stages are distributed along the common allometric line. For P. voutensis, all the specimens of morphotype 1 form a cluster in the lower part of the allometric line, indicating a smaller absolute size, while morphotype 2 specimens form a cluster in the upper part of the allometric line, indicating a larger absolute size. Based on these results, it is most probable that the two morphotypes correspond to two different ontogenetic stages of the same taxa, with morphotype 1 being a juvenile stage while morphotype 2 would represent an adult stage.

Another interesting fact here is the similarity in the number of segments and appendages described in both ontogenetic stages. As there is only size difference between both stages, P. voutensis displays, at least partly during its post-embryonic development, an epimorphic growth, i.e., a growth between stages without increment in segment numbers (Minelli & Fusco 2013).

Morphological and anatomical interpretations

Compound eyes

With the exception of the oldest species (Thylacares brandonensis: Haug et al. 2014), thylacocephalans possess hypertrophied compound eyes. These eyes have been initially interpreted as cephalic pouches, i.e., an extension of the cephalon containing the stomach (Arduini et al. 1980; Pinna et al. 1985). However, Fröhlich et al. (1992) and Vannier et al. (2016) later demonstrated that these presumed cephalic pouches were actually compound eyes. Vannier et al. (2016) were even able to describe the micro-structure of the ommatidia of Dollocaris ingens: they are formed of a corneal lens which sits on the top of a crystalline cone, and of rhabdom surrounded by four to five retinula cells. The eyes described here for other La Voulté thylacocephalans seem to have a comparable morphology: they consist of densely packed, hexagonal ommatidia. However, the ommatidia are larger than those of D. ingens and less densely packed. Here, it was not possible to determine the visual parameters of the eyes as they are poorly preserved compared to those of D. ingens.
In La Voulte thylacocephalans, with the exception of *Ostenocaris ribeti*, the compound eyes are followed by longitudinal muscles that are attached to the posterior edge of the eyes. This suggests the presence of a massive, tubular stalk. Presence of a stalk is not uncommon in thylacocephalans: it has been documented multiple times (e.g., Haug et al. 2014; Laville et al. 2021a), including for *D. ingens* (Secrétan 1985). Usually, eye stalks are considered to be long and narrow connecting structures, but here, the stalks are quite massive and short. This is not uncommon among arthropods, for which stalks can be variously developed from a short, massive structure to a narrow, elongate and flexible structure (e.g., Schoenemann et al. 2011; Strausfeld et al. 2016).

**Cephalic appendages**

Several flagellate structures have been previously interpreted as antennules or antennae in thylacocephalans. Haug et al. (2014) and Laville et al. (2021a) described two antenniform structures from specimens of *Mayrocaris bucculata* (Late Jurassic, Solnhofen Lagerstätten, Germany). They interpreted these structures as antennula and antenna, although an interpretation as a biflagellate antennula was not ruled out. Antenniform structures were also described in Triassic Malagasy thylacocephalans (Laville et al. 2021c). In our study, we describe similar flagellate appendages in specimens of *Mayrocaris* sp, *Ostenocaris ribeti*, *Paraostenia voulensis*, and *Paraclausocaris harpa* gen. et sp. nov. These structures are often located on the side of the large compound eyes but, except for *Mayrocaris* sp., their exact morphology and origin are unknown, preventing further interpretation of their nature.

Secrétan & Riou (1983: fig. 3) previously mentioned cephalic appendages, including antennae, in La Voulte thylacocephalans (*Dollocaris ingens*). On two specimens (MNHN.F.R50937, R50938), they described putative bases of several cephalic appendages. After re-examining these specimens, we can conclude that the first three elements, interpreted as the bases of the mandibula, the maxillula and the maxilla, actually belong to raptorial appendages. Concerning the first element, i.e., the putative antenna, it is unlikely to be an antenna due to its position just anterior to the raptorial appendages. It is probably a part of a masticatory appendages (i.e., the mandibula, the maxillula or the maxilla).

**Adductor muscles and muscle scar**

A bundle of radially arranged muscles is often found in the antero-ventral area of the shield of *Mayrocaris* sp. and *Paraclausocaris harpa* gen. et sp. nov. (Figs 10, 14). Similar structures have been described in other thylacocephalan taxa. Schram et al. (1999) interpreted similar muscles in *Protozooea hilgendorfi* Dames, 1886 as gastric muscles. In *Ostenocaris cypriformis* (Arduini et al. 1980), a similar bundle associated with a circular mark is present on the shield. Here, the muscles are interpreted as adductor muscles and thus, the rounded mark is believed to be a muscle scar. This would be consistent with the position and the morphology of the muscles found in *Mayrocaris* sp and *Pc. harpa*.

Another structure found on the shield of thylacocephalans might be interpreted as a muscle scar. A large rounded protuberance, located on the central area of the shield has been described as muscle scar on multiple species of thylacocephalan species (*Ligulacaris parisiana* [see Charbonnier et al. 2019]; *Dollocaris ingens* [see Secrétan & Riou 1983]; *Paraostenia voulensis* [see Secrétan 1985]; *Victoriacaris muhienisi* [see Hegna et al. 2014]; *Paradollocaris vannieri*, *Globulocaris garassinoi*, *Keelicaris deborae* [see Charbonnier et al. 2017]; *Mayrocaris bucculata* [see Laville et al. 2021a]). However, no muscles are found associated with these protuberances, making unlikely that they are muscle scars. Instead, Secrétan (1985) suggested that this protuberance might serve as a “pocket” for the raptorial appendages. The raptorial appendages, which are protruding from the shield just under the protuberance, could have been folded in the gap formed by the protuberance. This would be consistent with the fact that thylacocephalans might have been ambush predators and thus were lying on the seafloor, waiting for prey (Vannier et al. 2016). In modern ambush predators, such as stomatopods, the raptorial appendages are retracted under the margin of the shield when the organism is resting (Kunze & Anderson 1981).
This would also be consistent with the fact that thylacocephalans were swimming, at least occasionally. Having their raptorial appendages hidden under the shield would have given a more streamlined shape to the body, and thus the organism would have swum more efficiently.

**Fig. 22.** Complex allometry for *Paraostenia voultensis* Secrétan, 1985 (morphospace). **A.** Principal component analysis projection of morphospace on principal components 1 and 2. **B.** Correlation circle of variables for principal components 1 and 2. Abbreviations: Aad = antero-dorsal angle; Aav = antero-ventral angle; Apd = postero-dorsal angle; Apv = postero-ventral angle; Ha = anterior height; Hmax = maximal shield height; Hp = posterior height; Ls = shield length.
Oval structure

In *Austriocaris secretanae* sp. nov., *Kilianicaris lerichei*, *Paraclausocaris harpa* gen. et sp. nov. and *Paraostenia voultensis*, a large oval sclerotised structure is protruding between the raptorial appendages and the posterior trunk appendages (Fig. 23). This feature has already been reported by Secrétan & Riou (1983) in *Dollocaris ingens* and by Polz (1990: 108) in *Clausocaris lithographica* from Solnhofen Lagerstätten (Fig. 24), where it was referred to as “additional leaf-shaped limbs”. It has a rounded distal margin ornamented with a row of setae. It appears associated with the posterior trunk, as seen in the specimens on which the posterior trunk is detached from the shield. The structure does not appear segmented or related to any appendage, making unlikely that it represents an appendage, or a part of an appendage. Therefore, it probably represents an enlarged sclerite or a modified posterior trunk segment.

In terms of function, with its large surface with setose margins, this oval feature could have acted as a paddle, producing a thrust for swimming, similarly to the phyllopodous appendages of branchiopods (Hessler 1985). Another interpretation may be that of ventilatory function. Indeed, the oval structure resembles scaphognathite of decapod pancrustaceans, which corresponds to the exopod of the maxilla. It has a well-developed oval morphology and has setose margins (e.g., Boxshall & Jaume 2009).

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**Fig. 23.** Reconstruction of La Voulté thylacocephalans. **A.** *Paralausocaris harpa* gen. et sp. nov. **B.** *Kilianicaris lerichei* Van Straelen, 1923. **C.** *Austriocaris secretanae* sp. nov. **D.** *Paraostenia voultensis* Secrétan, 1985. Scale bars = 10 mm. Reconstructions: Alexandre Lethiers (made with Blender).
beating, it produces a water flow that brings water into the shield and helps ventilate the gills. In La Voulte-sur-Rhône thylacocephalans, the oval structure is located just beneath the gills, which would be a good position for ventilating them. Therefore, the oval structure might be analogous to the scaphognathite.

**Posterior trunk appendages**

Already mentioned in multiple other thylacocephalan species (Schram 1990; Haug et al. 2014; Laville et al. 2021a), a long and probably biramous terminal trunk appendage is also quite common in La Voulte thylacocephalans (*Kilianicaris lerichei*, *Paraclausocaris harpa* gen. et sp. nov.). A more interesting observation is the possible presence of other biramous posterior trunk appendages. Indeed, one specimen of *Pc. harpa* seems to display a biramous first posterior trunk appendage (Fig. 13A, C). Such detail has never been observed in thylacocephalans before.

**Telson**

Two particular posterior trunk segments drew our attention during the study of the La Voulte thylacocephalans: the 10th segment in *Paraclausocaris harpa* gen. et sp. nov. and *Paraostenia voultensis* (Figs 13, 17) and the 12th in *Austriocaris secretanae* sp. nov. and *Kilianicaris lerichei* (Figs 4, 6). These segments have a peculiar morphology compared to the rest of the posterior trunk segments: they have a squared shape, do not bear any appendage and display a unique rounded structure on their dorsal surface. On one specimen of *P. voultensis* (MNHN.F.A29300), setae can be observed on its distal part. Dorsally to these particular segments, the segments change orientation and appear symmetrical to the ventral ones. These ‘dorsal’ segments probably correspond to the other pleural parts of the ‘ventral’ segments. Thus, these peculiar segments represent the terminal segment of the posterior trunk. Based on their position and their morphology, the presence of marginal setae and the absence of appendages attached to these segments, we interpret them as the telson. It probably corresponds to the “reduced, unsegmented abdomen” of *Ostenocaris cypriformis* (Pinna et al. 1985).

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**Fig. 24. Clausocaris lithographica** (Oppenheim, 1888). A–B. Specimen JM SOS 972. A. General view. B. Close-up of oval setose structure. Scale bars: A = 5 mm; B = 2 mm. Photos: T. Laville.
On the telson, a large, circular structure can be distinguished. This structure consists of two circles: a large one, occupying most of the width of the telson, surrounding a smaller one. Such a structure was already mentioned by Secrétan & Riou (1983) for Dollocaris ingens. A rounded structure on the telson could be the anus. However, in pancrustaceans, the anus is usually located ventrally or at the posterior end on the telson (Bowman 1971). Here, the circular structure is dorsally located which makes it unlikely to be the anus. The circular structure could also be a tubercle or the basis of a spine, although it is unlikely as it does not rise from the telson and looks more depressed. One hypothesis would be that it represents a statocyst, a balance organ common among malacostracans (Espeel 1985; Sekiguchi & Terazawa 1997; Meland et al. 2015; Bober et al. 2018), and already described in fossil mysids (e.g., Voicu 1974; Schlacher et al. 1992; Wittmann et al. 1993; Vicente & Cartanyà 2017). It corresponds to a rounded invagination of soft-tissue that encloses ambient water and a rounded biomineralised statolith. This can give the impression of being formed by two concentric circles, such as what is found in La Voult-sur-Rhône thylacocephalans. It usually serves for stabilising the body in the water and for orientation while swimming. In macrostylid isopods, two symmetric statocysts can be found on the dorsal surface of the telson (Bober et al. 2018). Therefore, it would not be surprising to find such an organ on the telson of thylacocephalans. As it is, better-preserved specimens remain necessary before being able to determine precisely the nature of this circular structure.

**Pores**

Two different kinds of pores are recorded in La Voult-sur-Rhône thylacocephalans: 1) canal-like pores (Mayrocaris sp., Paraclusocaris harpa gen. et sp. nov. and Paraostenia voulensis), which are formed by a rim surrounding a hollow part (Figs 13, 19); 2) unrimmed pores on carina (Austriocaris secretanae sp. nov.; Kilianicaris lerichei). The canal-like pores are quite common among thylacocephalans (Lange & Schram 2002; Broda & Zatoń 2017; Laville et al. 2021a). They have been suggested to be sensory structures acting in a similar way as the sensory dorsal organs of pancrustaceans (Broda & Zatoń 2017). Among species with canal-like pores in La Voulte Lagerstätte, one is of particular interest: *P. voulensis*. Indeed, Secrétan (1985) reported that the canal-like pores forming a sinuous line enclosed somewhat of a capsule, that has been interpreted by Rolfe (1985) as putative photophores. While studying the numerous specimens of *P. voulensis*, no capsules have been found in the pores. Moreover, the morphology of these pores is different from the bioluminescent pores of halocyprid myodocops (Angel 1968), which form tubercles. In *P. voulensis*, the pores are flat and not raised. For these reasons, we consider the photophores hypothesis as unlikely.

The second kind of pores only occurs on the carina of *Austriocaris secretanae* sp. nov. and *Kilianicaris lerichei*. They lack a rim and appear filled with sediment. Unrimmed pores have been reported by Broda & Zatoń (2017) and interpreted as setal lumina, i.e., a simple pore (lumen) in the cuticle, that provides intracuticular articulation to the setae (Garm 2004). However, the pores found in *A. secretanae* and in *K. lerichei* (~100 µm) are much bigger than the previously mentioned setal lumina (~1 µm). Another hypothesis could be that the pores represent the bases of broken spines. Spines on the lateral area of the shield have only been reported on putative thylacocephalan cuticular fragments from the Middle Devonian of Czech Republic (Mergl 2020). On these remains, the spines, when broken, form pits. If these pits would have been filled by sediment, they could probably have a similar morphology as the pores found on *A. secretanae* and on *K. lerichei*. We cannot conclude definitively on the nature of such pores but the spine hypothesis should be kept in mind.

**Phylogenetic affinities of thylacocephalans**

Usually considered as pan/eucrustaceans (Haug et al. 2014; Vannier et al. 2016), a recent phylogenetic analysis has positioned thylacocephalans as stem-mandibulates in a clade with two enigmatic Palaeozoic euarthropods bearing raptorial appendages (Pulsipher et al. 2022): Acheronauta stimulapis Pulsipher et al., 2022 and Captopodus poschmanni Kühl & Rust, 2012. A position within mandibulates is not surprising as it
now seems certain that thylacocephalans had antenniform appendages (possibly antennula and/or antenna) and mandibula. However, a placement as a stem-mandibulates is more unexpected. Indeed, in addition to the cephalic appendages, several anatomical details suggest a close relationship with pancrustaceans: presence of proximal endites on raptorial appendages (Haug et al. 2014), sensory dorsal organs (Broda & Zatoń 2017), similarities of organic systems with these of pancrustaceans (Vannier et al. 2016).

Moreover, some uncertainties on the anatomy of thylacocephalans, in particular concerning the raptorial appendages, seems to preclude any firm conclusion as to their phylogenetic affinities. Indeed, the nature of raptorial appendages is still debated. They have been interpreted as being antennula, antenna and mandibula (Arduini et al. 1980), mandibula, maxillula and maxilla (Vannier et al. 2016) or as maxillula, maxilla and maxillipeds (Schram 1990; Haug et al. 2014). Some authors also suggested that they were entirely part of the trunk (Secrétan 1985; Rolfe 1985; Polz 1993). These various interpretations impact the tagmatisation of thylacocephalans. If the raptorial appendages are mostly or entirely considered as cephalic, it would imply that thylacocephalans consist of two regions, a cephalon and an undifferentiated trunk, which would be similar to Ca. poschmanni and to Ac. stimulapsis. In the case of thoracic raptorial appendages, thylacocephalans would have been formed by three tagma: a cephalon, an anterior trunk bearing raptorial appendages and a posterior trunk. In their phylogenetic analysis, Pulsipher et al. (2022) favoured the interpretation of the raptorial appendages of thylacocephalans as the maxillula, the maxilla and a maxilliped, and thus of a two-tagma body. They did not test alternative interpretations which would probably have led to a different phylogenetic position for thylacocephalans. Pending confirmation of the nature of the raptorial appendages, the results of Pulsipher et al. (2022) should thus be taken with caution, leaving the question of the phylogenetic affinities of thylacocephalans open.

Palaeoecology

Sensing

The antenniform appendages of La Voult-sur-Rhône thylacocephalans appear quite small in comparison to their hypertrophied compound eyes (Fig. 14A–D). The combination of large eyes and short antenniform appendages is common among thylacocephalans (Haug et al. 2014; Laville et al. 2021c). As mentioned by Schram & Koenemann (2022), an analogy can be made with hyperiid amphipods, which also have large eyes and atrophied antennulae and antennae (e.g., Bowman & Gruner 1973; Stenvers et al. 2021). With hypertrophied compound eyes that allow for an acute vision, the antennula might have been less useful and thus, would have been replaced by eyes as the primary sensory structure. Moreover, a complex sensory system formed of the canal-like pores has been reported in thylacocephalans (Broda & Zatoń (2017). Such sensory system may have supplemented the eyes, thus reducing the importance of antenniform appendages as sensory structures.

Locomotion

Several lifestyles have been proposed for thylacocephalans: endobenthic (Briggs & Rolfe 1983), epibenthic crawler (Pinna et al. 1982; Secrétan 1985),necktobenthic (Vannier et al. 2016) or nektonic (Rolfe 1985; Schram et al. 1999; Haug et al. 2014). It is most likely that thylacocephalans adopted several lifestyles during their evolution.

Concerning La Voult-sur-Rhône thylacocephalans, several features seems to indicate a nektonic or nektobenthic lifestyle (Fig. 23). First, they appear to have a streamlined shield (Vannier et al. 2016), as exemplified in specimens preserved in 3D or dorso-ventrally compressed (Fig. 7). This type of shield is likely to have reduced the drag effect. In addition to the streamlined aspect, the shield appears flexible and thin: it is often wrinkled but not broken and the underlying structures can often be distinguished. This combination is often indicative of a nektonic or nektobenthic lifestyle in extant pancrustaceans such as in Nebalia Leach, 1814 (Vannier et al. 1997) and halocyprid myodocops (Vannier & Chen 2000).
La Voulte thylacocephalans also seem to have a complex circulatory and respiratory system. They display eight large gills, allowing for efficient gas exchange that might have been ventilated by the oval structure mentioned above. The gills are connected by efferent and afferent vessels to the heart, which itself is connected to large arteries (Vannier et al. 2016; Schram & Koenemann 2022). According to Vannier et al. (2016), this system might be associated to a complex integumental circulatory system, such as what is known in extant myodocopids (Abe & Vannier 1995). This complex circulatory and respiratory system is indicative of an active lifestyle, consistent with a nektonic or nektobenthic lifestyle.

Some structures on the posterior trunk could also be related to a nektonic or nektobenthic lifestyle. The first of these structures is the combination of the square-shaped and flattened telson, with long terminal trunk appendages (Fig. 23). This recalls the tailfan of extant malacostracans which plays an important role for locomotion, either allowing for escape reaction (Neil & Ansell 1995), or for elevating from the seafloor (Vannier et al. 1997). A similar function is likely in thylacocephalans. Another structure of the posterior trunk might play a role in locomotion, the putative statocyst. In mysids and decapods, the statocyst has an important function during locomotion, as it allows for stabilisation of the body and for directional swimming (Neil 1975). If this circular structure is confirmed to be a statocyst, it probably played an important role for swimming in thylacocephalans.

Finally, the only missing information to confirm the swimming ability of La Voulte thylacocephalans is the morphology of the trunk appendages. Indeed, except for Austriocaris secretanae sp. nov. for which they appear rod-like (Fig. 4E, H), they are poorly preserved, preventing from determining their exact morphology and thus, from confirming the swimming ability of thylacocephalans. Despite these uncertainties, the morphological evidence largely supports an active nektonic or nektobenthic lifestyle for La Voulte thylacocephalans.

Diversity of Jurassic thylacocephalans

With seven different taxa described, the La Voult-sur-Rhône Lagerstätte is the most taxonomically diverse thylacocephalan fauna known so far (Fig. 23). Looking at the distribution of thylacocephalan species along the stratigraphic column of this outcrop (Fig. 1), it appears that these thylacocephalans form a unique assemblage. Indeed, the distribution of all taxa spans the entire stratigraphic layer and more importantly, they co-occur in layer g-8-9 (Fig. 1C).

In terms of diversity, the description of two new species, Austriocaris secretanae sp. nov. and Paraclausocaris harpa gen. et sp. nov. from La Voult Lagerstätte greatly increase the specific diversity of Jurassic thylacocephalans. Even though only four thylacocephalan-bearing localities are known in the Jurassic, they preserve an important specific diversity with 11 species described (Table 1). Another Jurassic species was previously assigned to thylacocephalans: Rugocaris indunensis Tintori, Bigi, Crugnola & Danini, 1986. The affinities of this species with thylacocephalans has always been doubtful (e.g., Schram 2014). Indeed, this species displays a unique morphology among thylacocephalan: it has a sub-triangular shield with rugations. Recently, we were informed that the holotype and only specimen of R. indunensis actually represents a fragment of a cephalopod (G. Teruzzi, pers. com.). Thus, we no longer consider this species as a thylacocephalan.

In addition to the important specific diversity, Jurassic thylacocephalans also preserve a large morphological diversity, with unique anatomical details and a huge diversity of shield and appendage morphologies. The Jurassic is also most probably the period with the longest thylacocephalans, as exemplified by Austriocaris secretanae sp. nov. or O. cypriformis (see Vannier et al. 2006).

In terms of palaeogeographic distribution, all known Jurassic localities are restricted to the Tethys Ocean, as are most of the Triassic thylacocephalan-bearing outcrops (Ehiro et al. 2019). This can
explain the similarities between these Jurassic faunas, being geographically close. Indeed, the La Voultè thylacocephalan assemblage shares many taxa with other Jurassic assemblages: *Mayrocaris* and *Dollocaris* are also known from the Solnhofen Lithographic Limestones (Polz 1988, 1989, 1997, 2001) while *Paraostenia* and *Ostenocaris* are also known from the Osteno Lagerstätte (Arduini et al. 1980; Pinna et al. 1985). However, the small number of known localities throughout the Jurassic prevents a biogeographic synthesis of thylacocephalans during the Jurassic.

**Conclusion**

The reappraisal of the La Voultè thylacocephalans provides important insight into the diversity and morphology of Jurassic thylacocephalans. It reveals an unexpected diversity, with the description of two new species, *Austriocaris secretanae* sp. nov. and *Paraclausocaris harpa* gen. et sp. nov., and the occurrence of *Mayrocaris* sp.

La Voultè thylacocephalans also greatly improve our knowledge of the anatomy of the group. We recognised new key anatomical features, such as an oval structure or putative statocyst on the telson, which provide additional pieces of evidence for nektonic or nektobenthic lifestyle in La Voultè thylacocephalans.

Finally, the re-investigation of La Voultè thylacocephalans also provides information concerning their ontogeny. A juvenile stage of *Paraostenia voultensis* is reported for the first time, which indicates that this species had, at least partly during its post-embryonic development, an epimorphic growth.

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**Supplementary material**

**Supp. file 1.** List of the material studied. [https://doi.org/10.5852/ejt.2023.898.2295.9935](https://doi.org/10.5852/ejt.2023.898.2295.9935)
