The Cambrian trilobites of Jordan – taxonomy, systematic and stratigraphic significance

OLAF ELICKI1 AND GERD GEYER2

1 Geological Institute, TU Bergakademie Freiberg, Bernhard-von-Cotta-Straße 2, 09599 Freiberg, Germany. E-mail elicki@geo.tu-freiberg.de
2 Institut für Geographie und Geologie, Lehrstuhl für Geodynamik und Geomaterialforschung, Bayerische Julius-Maximilians-Universität Würzburg, Am Hubland, 97074 Würzburg, Germany; and Department of Earth Sciences (Palaeobiology), Uppsala University, Villavägen 16, 752 36 Uppsala, Sweden. E-mail: gerd.geyer@uni-wuerzburg.de

ABSTRACT:


Marine carbonates and siliciclastic rocks of the Burj Formation in Jordan include paucispecific trilobite associations of the (traditional) Lower–Middle Cambrian boundary interval. Comprehensive new material of these trilobites allows a review of their taxonomy and systematic positions as well as a refined morphological description and a reconsideration of previous interpretations of their stratigraphic position and thus the correlation of the fossiliferous beds. In addition to the classic species Kingaspis campbelli (King, 1923) and Redlichops blanckenhorni Richter and Richter, 1941, Timnaella? orientalis (Picard, 1942) and Hesa problematica Richter and Richter, 1941, the discussed trilobites include Issalia gen. nov. with Issalia scutalis gen. nov., sp. nov., Tayanaspis gen. nov. with Tayanaspis bulbosus gen. nov., sp. nov., Uhaymiria gen. nov. with Uhaymiria glabra gen. nov., sp. nov., Cambriunicornia? jafnaensis sp. nov., Myopsolenites palmeri (Parnes, 1971), M. hyperion sp. nov., and Enixus cf. antiquus (Chernysheva, 1956). Myopsolenites boutiouiti Geyer and Landing, 2004 is now regarded as a junior synonym of Myopsolenites altus (Liñán and Gozalo, 1986). A detailed discussion of the correlation with a focus on global aspects provides clues for the utility of potential index fossils for the global Cambrian Series 3 and Stage 5.

Key words: Cambrian; Trilobita; Stratigraphy; Global correlation; Dead Sea; Jordan; Israel; Spain; Morocco; Poland; South China; Siberia.

INTRODUCTION

Cambrian trilobites from Jordan have been known since the early years of the twentieth century, when the German geologist Max Blanckenhorn (1861–1947) discovered a number of specimens near the Dead Sea in 1908. He also reinterpreted trilobite remains sampled by E. Hull near the end of the nineteenth century close to the southern tip of the Dead Sea. A limited number of studies, such as those by Blanckenhorn (1910), Dienemann (1915), Richter and Richter (1941), Parnes (1971) and Rushton and Powell (1998), provided an overview of the oligospecific faunal composition and the obviously small stratigraphic window in which the known species occurred. However, most of the earlier studies suffered from a limited amount of material and incomplete knowledge on the fine-scale stratigraphy and lateral facies changes in the region. This led to difficulties in understanding the ontogenetic variation and precise systematic position of the taxa as well as a stratigraphic
assignment and correlation into other areas. Such correlations were mostly biased and, in addition, neglected the context of facies architecture and continental geotectonic evolution.

This study is based on copious additional material which gives a more complete portrait of the species and suggests modifications of the taxonomy and stratigraphic position. Nevertheless, new species and informally described additional forms suggest that the preservational window provides us merely with an imperfect glance at the biota during this interval in the region.

Spelling of geographic and stratigraphic Arabic local terms used in this paper is adopted from the 1:50,000 geological map sheets. It should be noted that syntax varies among the different maps and their explanations because of the lack of a transliteration standard.

Figured specimens are housed in the collection of the Geological Institute of the TU Bergakademie Freiberg under collection number FG-602.

LOCALITIES

The material described herein has been collected during several field seasons over the last decade by one of us (OE) and his working group. Additional material was collected by GG but is not used for the documentation. For the first time, trilobites from the southern Dead Sea area and from northern Wadi Araba were sampled in situ from the bedrock so that their stratigraphic position within the succession is fixed. All former trilobite finds came either from only a single stratigraphic horizon of the Wadi Zerqa Ma’in section, northern Dead Sea shore, or from allochthonous material from the valleys at the southern margin of the Dead Sea. The material described in this study comes from four localities and sections (Text-fig. 1):

1. Wadi Zerqa Ma’in (“1” in Text-fig. 1, “B” in Text-fig. 2 left and Text-fig. 2 right, Text-fig. 3)
Northeastern shore of the Dead Sea, near the mouth of Wadi Zerqa Ma‘in; 31°37’56” N, 35°34’26” E.

The classic Cambrian locality of the northeastern Dead Sea is located about 1 km north of the mouth of Wadi Zerqa Ma‘in and exposes an approximately 80 m thick succession. The incomplete succession has been reinvestigated and described in detail by Shinaq and Elicki (2007). It consists of seven lithostratigraphic units, which represent the higher part of the Burj Formation, followed by strata of the Umshirin Formation. The marine Burj Formation shows four shallow marine to marginal marine sandstone and minor silstone units (partly with *Cruziana* ichnofossil assemblages) subdivided by three levels of shallow subtidal carbonates, each of them only a few metres thick. Trilobites of this locality come mainly from the lowermost of the exposed carbonate units. *Kingaspis campbelli* and *Enixus cf. antiquus* come from two separate bioturbated limestones of the lowest carbonate interval *sensu* Shinaq and Elicki (2007). *Enixus cf. antiquus* occurs sporadically in a fossil-bearing, distinctly bioturbated part at the top of a cross-bedded oolite. *Kingaspis campbelli* occurs in huge numbers in a bioturbated bioclastic grainstone to rudstone facies about one metre above the massive oolite (Shinaq and Elicki 2007). The siliciclastic heterolithic unit on top of this carbonate level is the type locality and stratum of *Cruziana salomonis* (Seilacher 1990) and contains abundant specimens of this trace fossil. About 43 m upsection (third and youngest carbonate interval *sensu* Shinaq and Elicki 2007), crania of *Kingaspis campbelli* were found in carbonates of a thinly bedded alternation of sandstone and ooid-bearing bioclastic grainstone. The second carbonate interval in between the two above-mentioned carbonate levels also contains trilobite remains, which are seen in thin sections, but no specimens could be cracked out from this level.

2. Wadi Issal (“2” in Text-fig. 1, “C” in Text-fig. 2)
Southeastern Dead Sea area; 31°11’22” N, 35°33’08” E.
Trilobite material from the southeastern Dead Sea area comes from Wadi Issal and Wadi Uhaymir (Text-fig. 1). Only the upper part of the Burj Formation is exposed at Wadi Issal. The trilobite remains [*Issalia scutalis* gen. nov., sp. nov., *Myopsolenites palmeri* (Parnes, 1971)] come from the basal beds of the Hanneh Member. Cranidia and other remains are generally disarticulated and occur in sandstone horizons of shallow marine origin.

3. Wadi Uhaymir (“4” in Text-fig. 1, “A” in Text-fig. 2, Text-fig. 4)
Southeastern Dead Sea area; 31°9’11” N, 35°33’37” E.

At Wadi Uhaymir (‘Wadi Tayan locality’ *sensu* Elicki et al. 2002; see below), many trilobite finds come from a distinct carbonate level within the Numayri Member, about 12.5 m below the transition to the overlying Hanneh Member. The most prolific fossil horizon is an approximately 10 cm thick hash layer, which overlies an alternation of fine-grained limestone beds and marlstones (each a few centimetres thick). The hash layer is a bioturbated bioclastic floatstone with a sharp base. The trilobites [*Timnaella? cf. orientalis* (Picard, 1942), *Tayanaspis bulbosus* gen. nov., sp. nov., *Uhaymiria glabra* gen. nov., sp. nov., *Myopsolenites salomonis* sp. nov., *Myopsolenites palmeri* (Parnes, 1971), genus and species undeterminate 1, genus and species undeterminate 2] occur mostly as disarticulated sclerites and are accompanied by common cone-in-cone nested monospecific hyolith assemblages (*Hyolithes kingi* Richter and Richter, 1941) and some brachiopods. The bioclasts are sporadically strongly current-oriented in small erosional channels (Shinaq and Elicki 2007; OE, unpubl. data). Interestingly, various ontogenetic stages occur among the large number of trilobite specimens. Larger carapaces, which occur quite rarely, functioned as a shelter and have capped smaller fossil remains, indicating significant runoff after high-energy deposition of this stratum. Upsection, the hash layer changes into an oncoid limestone with abundant brachiopods (nearly exclusively *Trematosia*). The depositional environment of this part of the Burj Formation at Wadi Uhaymir has been interpreted by Elicki et al. (2002) as a shallow marine setting with interfingered of lagoonal sediments and open-marine oolite shelf facies affected by occasional storm events.

4. Wadi Umm Jafna (“6” in Text-fig. 1, “A” in Text-fig. 2)
Ghawr Fifa area of the northeastern Wadi Araba; 30°56’39” N, 35°29’58” E.
The trilobite specimens from the Wadi Umm Jafna section (*Cambriuncornia? jafnaensis* sp. nov. and incomplete sclerites of two undeterminable species) come from bioclastic wackestones to floatstones with intraclasts of the Numayri Member, which occur about 15 m below the transition into the overlying Hanneh Member. The fossiliferous bioclastic beds are a few centimetres thick and intercalated between platy limestones. The two lithotypes alternate repeatedly over a vertical distance of several metres. Such a depositional facies is similar to that observed in the Wadi Uhaymir section in an interval 16 to 20 m below the trilobite hash layer.
Wadi Issal, Wadi Uhaymir and Wadi Umm Jafna are newly discovered fossil localities. The Wadi Uhaymir section is very close to an offshoot from Wadi At Tayan, leading Elicki et al. (2002) in describing sedimentary facies types from this locality to term it the ‘Wadi Tayan locality’. Rushton and Powell (1998) assumed Wadi At Tayan as being more-or-less synonymous with Wadi Rimeileh, an area from where King (1923) reported trilobite remains of ‘a distinctly Asaphid type’ [= Myopsolenites palmeri (Parnes, 1971)] from green micaceous siltstones to claystones. King (1923) located Wadi Rimeileh ‘about 1.6 km south of Wadi Issal’. As already noted by Elicki (2007), the litho- and biofacies characteristics of King’s samples from Wadi Rimeileh are identical to those of the Hanneh Member of the nearby Wadi Issal locality. Because of this obvious accordance, a distinctly more southerly position of Wadi At Tayan (about 5 km south of Wadi Issal), and the absence there of King’s lithofacies (OE, unpubl. data), Wadi Rimeileh (mouth at 31°10’22” N, 35°32’12” E) is apparently not synonymous with Wadi At Tayan, but represents a valley unnamed on the 1:50,000 geological map sheet and situated between Wadi Issal and Wadi At Tayan at exactly the geographic position given by King (1923) (Text-fig. 1). In addition, according to new observations (OE, unpubl. data), the Burj Formation in this area is exclusively represented by the Hanneh Member siltstones and sandstones, which is in accordance with the 1:50,000 geological map sheet. This supports the geographic locations discussed above.

GEOLOGICAL SETTING

The Dead Sea Rift Valley, which separates the Arabian Plate from the African Plate, is situated at the northern part of the Cenozoic Great Rift Valley but represents an extensional structure related to a triple junction active in late Proterozoic to early Cambrian times (Bender 1968a; Husseini 1989; Sharland et al. 2001). Crystalline basement rocks of the Arabian–Nubian Shield (Aqaba Complex) are exposed at its southern edge in the southern Wadi Araba, and are unconformably overlain to the north by Neoproterozoic (Araba Complex) and/or early Palaeozoic sedimentary rocks (Ram Group) (e.g. Bender 1968b; Segev 1984; Amireh et al. 1994; Elicki 2007; Schneider et al. 2007).

Cambrian rocks in the Middle East region are known from scattered, isolated outcrops in the Dead Sea area, the Wadi Araba and southward to Wadi Rum in Jordan, as well as from the Timna region in the southern Negev Desert, Israel, and from minor outcrops on the Sinai Peninsula, Egypt.

Onlapping the Arabian–Nubian Shield in the south, the thickness of the Cambrian rock succession increases northward to nearly 700 m in the Dead Sea area, and to probably more than 1000 m in the subsurface of northern Jordan and southern Syria (Bender 1968b; Powell 1989; Shinaq and Bandel 1992; Best et al. 1993).

In the study area at the northeastern and the southeastern edge of the Dead Sea and in the northeastern Wadi Araba (Text-fig. 1), the Cambrian succession (Text-fig. 2) starts with conglomeratic, fluvial and alluvial plain siliciclastics of the Umm Gaddah Formation (up to 60 m thick, late Ediacaran to Early Cambrian; Amireh et al. 2008) and Salib Formation (probably more than 200 m thick, Early Cambrian; Selley 1972; Powell 1989; Amireh et al. 1994; Makhlouf 2003). Deposition of the Salib Formation was occasionally influenced by marine ingresses in its upper part and the
formation is overlain by the ?late Early to Middle Cambrian marine Burj Formation (up to 120 m in the Dead Sea area), followed by predominantly continental siliciclastics of the Umm Ishrin Formation. In southern Jordan, the siliciclastic rocks of the Abu Kusheiba Formation represent lateral facies types equivalent to strata of the Burj Formation. The Cambro(?)-Ordovician mainly fluvial Disi Formation overlies the Umm Ishrin Formation and wedges out from south to north in the Wadi Dana area.

The main marine incursion of this succession is represented by the Burj Formation, which is a mixed siliciclastic-carbonate unit with a complex architecture (Rushton and Powell 1998; Elicki 2007; Shinaq and Elicki 2007). It is subdivided into three members termed the Tayan (Tayan Siltstone), Numayri (Numayri Dolomite Shale) and Hanneh (Hanneh Siltstone) members. The Tayan Member consists of transgressive siltstones and sandstones with sporadic dolomite intercalations and attains a thickness of up to 21 m (Elicki 2007). The known fossil content is restricted to simple trace fossils and sporadic stromatolitic horizons (Powell 1989; Rushton and Powell 1998; Elicki 2007). The overlying Numayri Member (up to about 120 m in the Dead Sea area; Andrews 1991) consists of shallow marine limestones, dolostones and few marly siliciclastics. The carbonates are commonly rich in shelly fossils (trilobites, hyoliths, brachiopods, echinoderms, sponges and others; Elicki 2011). Distinct horizons yield stromatolites (Shinaq and Bandel 1992; Rushton and Powell 1998; Elicki et al. 2002; Shinaq and Elicki 2007). The sandstone-dominated siliciclastic Hanneh Member has yielded a few trilobite remains at its base and rich trace fossil assemblages (Makhlof and Abed 1991; Amireh et al. 1994; Rushton and Powell 1998; Mángano et al. 2007; Shinaq and Elicki 2007; Hofmann et al. 2012; Mángano et al., in press). The Burj Formation is interpreted as a marginal to shallow marine succession deposited during a relatively short marine ingression on the southerly exposed basement rocks or on their continental detritus (e.g., Amireh et al. 1994; Rushton and Powell 1998; Elicki 2007).
STRATIGRAPHY AND FOSSIL OCCURRENCES

The northernmost known fossiliferous Cambrian outcrop near the mouth of Wadi Zerqa Ma’in (see Shinaq and Elicki 2007 and Text-figs 1, 2) was discovered by K.A. Campbell in the beginning of the twentieth century. Campbell’s material and additional trilobites from the southern Dead Sea area have been studied by King (1923), Richter and Richter (1941), and Picard (1942).

The Cambrian of the western margin of the Dead Sea Rift Valley in the southern Negev in Israel was examined by Parnes (1971). Roughly equivalent strata were first studied in the Timna area and at Har ‘Amram (Parnes 1971). These strata of the southern Negev yield trilobite assemblages with taxa which are, in part, in need of a revision (Geyer and Landing 2000).

In the Cambrian of the Dead Sea and Timna areas, trilobites and other shelly fossils are known exclusively from a fairly thin interval at the traditional Lower–Middle Cambrian boundary. Older early Cambrian and younger mid to possibly late Cambrian rocks are interpreted as mainly fluvial sequences (e.g., Bender 1968b; Selley 1972; Powell 1989; Pflüger 1990; Makhlouf and Abed 1991; Makhlouf 2003; Schneider et al. 2007). Nevertheless, at least one marine incursion, represented by a thin interval with trace fossils generated by arthropods (Diplichnites, Cruziana, Rusophycus and others), have been identified from regions such as the northern Wadi Rum near the Jordanian–Saudi Arabian border (Geyer and Landing 2000; OE, unpublished data).

The fossiliferous rocks of the traditional Lower–Middle Cambrian boundary interval belong to a depositional sequence composed of the Umm Gaddah, Salib, Burj, Umm Ishrin and Disi formations (Selley 1972; Powell 1989; Amireh et al. 2001, 2008), which can be traced westward into the Timna region, where the three middle units are termed the Amudei Shelomo, Timna, and Shehoret formations (Karez and Key 1965; Weissbrod 1981; Segev 1984).

Simplified, the marine incursion (Burj Formation) reflects a transgressive-regressive cycle (Rushton and Powell 1998; Elicki 2007), with the shelly fossils marking the marine highstand. This development reflects a
Hawke Bay-type transgression at the Lower–Middle Cambrian turnover.

The trilobite faunas from the Burj and Timna formations have long been known, but the oligospecific assemblages have been imperfectly studied (Richter and Richter 1941; Parnes 1971; Rushton and Powell 1998). Rushton and Powell (1998) presented a meticulous analysis of the trilobites of the Burj Formation. They distinguished between a level with *Kingaspis campbelli* and *Enixus cf. antiquus* (as *Palaeolenus antiquus*) in the upper part of the Numayri Member and a slightly lower level with *Tayanaspis bulbosus* (as *Realaspis sp. nov.*), *Redlichops blanckenhorni* and *Myopsolenites palmeri* (as *Onaraspis palmeri*). However, the *Kingaspis campbelli* faunule is known only from limestones of the classic Wadi Zerqa Ma’in locality. The taphonomic aspects of these limestones (composed of high energy shell accumulations) suggest that the beds represent an ecosтратigraphic horizon. The *Redlichops* faunule is known only from other southwardly located sites and occurs in quite variable carbonate-dominated rocks with notable shaly intercalations, which are interpreted to have been deposited in environments with generally low to moderate wave energy. Fairly intermittent trilobite occurrences contrast with few accumulations of various brachiopods (Cooper 1976). The puzzling occurrence of *Hesa problematica* (see below) also belongs to this level. In conclusion, the depositional and taphonomic aspects do not allow the *Kingaspis campbelli* faunule and the *Redlichops* faunule to be regarded as distinct biostratigraphic levels.

It should be noted that the Burj-equivalent Timna Formation of the Negev region is also divided into three members, termed the Hakhlil, Sosgun, and Mikhrot members (Segev 1984). The known trilobites from the formation include *Myopsolenites palmeri* and *Timnaella* spp., and all come from the Hakhlil Member, which contains extremely shallow marine to peritidal and subaerial to locally lacustrine deposits, with the most unrestricted marine deposits close to the top. The report of *Myopsolenites palmeri* from the Mikhrot Member (Parnes 1971) appears to be erroneous and based on local stratigraphic complications (E. Landing and G. Geyer, unpubl. data). In addition to the trilobites, these beds are locally rich in trace fossils such as *Cruziana*, “*Monocraterion*” and *Planolites* (Geyer and Landing 2000 and unpubl. data).

**CORRELATION**

The potential of the trilobite occurrences in the Dead Sea and Timna areas to enhance regional and even intercontinental correlation have been widely ignored. The reason was simply that the trilobites were either recognized as endemics or mismatched with genera or species erroneously identified from elsewhere. Even with the new taxa described by Rushton and Powell (1998) and in this study, correlation is not straightforward.

The classic *Kingaspis campbelli* was the first trilobite which became well known from the area and was utilized for correlation. This species was the first strongly effaced trilobite from a wider stratigraphic interval, and similarly effaced trilobites from the traditional Lower and Middle Cambrian boundary interval were frequently interpreted as close relatives, or even regarded as synonyms, and used as a basis for correlation. These largely arbitrary correlations are not discussed herein. The study of the ellipsocephaloid trilobites of the Moroccan Atlas ranges (Geyer 1990b) unfolded a complex taxonomic scheme of the group and showed that the trilobites previously grouped under *Kingaspis* Kobayashi, 1935 in fact represented at least three different genera. Even the restricted concept of *Kingaspis*

<table>
<thead>
<tr>
<th>Locality</th>
<th>Wadi Zerqa Ma’in</th>
<th>Wadi Issal</th>
<th>Wadi Uhaymir</th>
<th>Wadi Umm Jafna</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Kingaspis campbelli faunule</strong></td>
<td><em>Kingaspis campbelli</em></td>
<td><em>Enixus cf. antiquus</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Redlichops faunule</strong></td>
<td><em>Issalia scutalis</em></td>
<td><em>Myopsolenites palmeri</em></td>
<td><em>Redlichops blanckenhorni</em></td>
<td><em>Cambrunicornia? jafaensis n. sp.</em></td>
</tr>
<tr>
<td></td>
<td><em>Tayanaspis bulbosus</em></td>
<td><em>Uhaymiria glabra</em></td>
<td><em>Myopsolenites hyperion</em></td>
<td>Genus and sp. undet. 1</td>
</tr>
<tr>
<td></td>
<td><em>Myopsolenites palmeri</em></td>
<td></td>
<td><em>Myopsolenites palmeri</em></td>
<td>Genus and sp. undet. 2</td>
</tr>
</tbody>
</table>

Text-fig. 5. List of Cambrian trilobites described from Jordan
Cambrian Trilobites of Jordan

Included species which occur in at least three different zones in the Moroccan Cambrian. *Kingaspis campbelli* was identified from the easternmost Anti-Atlas close to the Moroccan–Algerian border, but unfortunately in rocks which cannot be placed unequivocally into one distinct zone. According to the existing zonation for the Anti-Atlas ranges (Geyer 1990a), the species occurs either in the upper part of the *Morocconus* Zone (formerly the *Cephalopyge* Zone) or in the overlying *Ornamentaspis frequens* Zone (Geyer 1990b), which are coeval with part of the lower (but not lowest) range of *Paradoxides* s.1. and thus the base of the traditional ‘Acadobaltic’ Middle Cambrian (compare Geyer and Landing 2004; Geyer 2005; Geyer and Peel 2011).

Specimens of *Kingaspis* from the Iberian Chains, northern Spain, as well as specimens from the Láncara Formation of the Cantabrian Mountains have been identified as *K. campbelli* (Liñán et al. 2003; Dies et al. 2004). As outlined by Geyer and Landing (2004), this identification is certainly erroneous (see discussion under *K. campbelli*), but was maintained by Gozalo et al. (2007) and further used for direct correlation. The specimen from the Valdoré section figured by Gozalo et al. (2007, fig. 4A) shows the shell exterior with features characteristic of *Kingaspoides* Geyer, 1990 rather than *Kingaspis*. The Iberian material from Aragón occurs in the local *Protolenus dimarginatus* Zone, where specimens notoriously suffer from notable tectonic distortion.

1The name *Cephalopyge* Geyer, 1988 for the trilobite genus and the eponymous zone is a junior homonym of *Cephalopyge* Hanel, 1905, a Recent phylliroid nudibranch. The senior author of this paper had been aware of this and had started preparation of a short manuscript in which the name *Cephalopyge* Geyer, 1988 would have been replaced and some of the morphological details of the trilobite *Cephalopyge* further scrutinized. In a sort of nomenclatural piracy, a colleague recently suggested the new name *Morocconus* without having had any contact with any of the authors affected by the article in question (Özdikmen H. 2009, Nomenclatural changes for Twenty trilobite [sic] genera. *Mun. Ent. Zool.* Vol. 4, No. 1, 155–171)
The eponymous *Protolenus (Hupeolenus) dimarginatus* Geyer, 1990 is a species first described from the *Morocconus* Zone of southern Morocco, where it most probably spans its lower and middle part and appears to range into the top of the underlying *Hupeolenus* Zone of Morocco, which is primarily characterized by other species of *Protolenus (Hupeolenus)* such as *P. (H.) hupei* Geyer, 1990 and *P. (H.) termierelloides* Geyer, 1990 (e.g., Geyer 1990a; Geyer and Landing 2006). Gozalo et al. (2007, p. 367) erroneously state that this species, as well as *Protolenus (P.) interscriptus*, have been “defined [...] in the *Hupeolenus* zone.” Both were in fact described from the *Morocconus* Zone of Morocco, and *P. (P.) interscriptus* is restricted to this zone (Geyer 1990b).

Summarized, there is little likelihood that *Protolenus (Hupeolenus) dimarginatus* and *Kingaspis campelli* could occur together in the Moroccan sections. Furthermore, *Protolenus (Hupeolenus) termierelloides* is also identified from the *Protolenus (Hupeolenus) dimarginatus* Zone of the Iberian Chains (Dies et al. 2004, as “cf. *termierelloides*”; Gozalo et al. 2007). This species ranges in Morocco from the acme in the *Hupeolenus* Zone into the top of the underlying *Hupeolenus* Zone of Morocco, which is primarily characterized by other species of *Protolenus* and *Kingaspis campelli*. Further, the *Morocconus* Zone is overlain by the *Protolenus jilocanus* Zone (formerly the *Hamatolenus ibericus* Zone) and the *Acadoparadoxides mureroensis* Zone, which marks the base of the Middle Cambrian in Iberia.

**Paradoxides** (A.) *mureroensis* Sdzuy, 1957 has not yet been described and figured from the Moroccan Atlas ranges but occurs there in some sections in the *Morocconus* Zone. Its first occurrence is always well above the base of the zone (G. Geyer and A. Vincent, unpubl. results). In contrast, *Paradoxides* (A.) *nobilis* Geyer, 1997 has its first occurrence at or even below the base of the *Morocconus* Zone, and that species has been erroneously synonymized with *P. (A.) mureroensis*, which led to the report of the latter species as existing in Morocco (e.g., Sdzuy 1995; Sdzuy et al. 1999; Liñán et al. 2002). Hence, some of the identifications of the Spanish material appear to obscure the quite obvious correlation between Iberia and southern Morocco, which are reinforced by depositional patterns with distinct transgressive events (Álvaro et al. 2003; Landing et al. 2006).

Another trilobite linking between Iberia, Morocco and Jordan is *Myopsolenites*. The genus was unfortunately established in an unusually brief manner (Öpik 1975), which has created unnecessary confusion (see Geyer and Landing 2004, and discussion below). Despite notable morphological differences, Gozalo and Liñán (1997), Rushton and Powell (1998), Gozalo et al. (2007) and Dies Álvarez et al. (2007) synonymized *Myopsolenites* with the endemic Australian genus *Omaraspis*. As discussed below, both genera are interpreted herein to represent members of the Bathynotidae; a small family which has a strong and short acme at the traditional Early–Middle Cambrian boundary interval. In addition to *Myopsolenites palmeri*, *M. hyperion* sp. nov. is another species which occurs in Jordan, and additional, morphologically similar species are known from southern Morocco, the Holy Cross Mountains in Poland, the Iberian Chains and the Cantabrian Mountains in Spain. All of the Moroccan material comes from the *Morocconus* Zone (Geyer and Landing 2004). The material from the Iberian Chains was found in the *Protolenus jilocanus* Zone (Gozalo et al. 2007). However, based on new material published by Dies et al. (2007), it can be shown that the Spanish species *Myopsolenites antiquus* (Liñán and Gozalo, 1986) is a senior synonym of the Moroccan *Myopsolenites boutioui* Geyer and Landing, 2004 (see discussion below under *Myopsolenites* in the systematic section). The above-mentioned species plus additional faunas enable an apparently precise correlation between the Moroccan Atlas ranges and the Iberian Chains, with a fairly reliable correlation into other regions of West Gondwana and into the Holy Cross Mountains (Text-fig. 6). However, these correlations deviate notably from those presented by Dies et al. (2007, fig. 5). The reason for this is either that the precise stratigraphic ranges of these and additional accompanying trilobite species are not known, or that the correlations from the Iberian successions into other areas are incorrect.

A species of *Enixus* (formerly *Schistocephalus*) occurs in association with *Kingaspis campbelli* in the Dead Sea area. The species is dealt with herein as *Enixus cf. antiquus* and has been identified as *Palaeolenus antiquus* by Rushton and Powell (1998). *Enixus antiquus* is the index fossil of the lower Amgan *Schistocephalus antiquus* Zone. Our tentative identification refers to minor differences which are difficult to quantify taxonomically. However, regardless of these differences, it can be assumed that the form permits a direct stratigraphic correlation into the lower part of the Siberian Amga Stage. Complications arise, however, if the former genus *Schistocephalus* is merged with *Palaeolenus* Mansuy, 1912 and *Megapalaeolenus* Chang, 1966, as suggested by Lin and Peng (2004). The morphological concepts and taxonomic consequences are discussed below under *Enixus*. Stratigraphically, there exists a morphological lineage from species of *Palaeolenus* to species of *Megapalaeolenus*, which indicates that a considerable amount of time is involved so that a collective genus *Palaeolenus* has little significance for correlation. More important is that another probable
species of Enixus has been found in the Morocconus Zone of the High Atlas mountains, associated with Clavigellus Geyer, 1994, a genus which was found in the P. (A.) muroeroensis level in the lower Láncara Formation of the Cantabrian Mountains (Gozalo et al. 2007), in the lower part of the Campo Pisano Formation of Sardinia, together with the oldest Acodaparadoxides specimens (Elicki and Pillola 2004), and in the Çal Tepe Formation of the Amanos Mountains, Turkey, where it occurs with P. (A.) cf. muroeroensis (Dean and Özgül 1994; Dean 2005). Palaeolenus medius (Bednarczyk, 1970) has been found in the Holy Cross Mountains, in strata which also yielded Paradoxides (A.) cf. muroeroensis (Zylińska and Masiak 2007).

Yuan et al. (2009) used the taxonomic lumping of Schistocephalus with Palaeolenus and a supposed similarity of Enixus antiquus with Megapalaeolenus deprati (Mansuy, 1912) to correlate the base of the Amgan Stage roughly with the upper part of the Megapalaeolenus Zone, or the Articcephalus chaueuai Zone in South China. The presence of Ovatoryctocara granulata Chernysheva, 1962, in the Henson Gletscher Formation of North Greenland, in strata above the level with Articcephalus chaueuai Bergeron, 1899, enables to falsify the correlation suggested by Yuan et al. (see comprehensive discussion in Geyer and Peel 2011). Ovatoryctocara granulata, a candidate GSSP marker for the base of the proposed Cambrian Series 3 (to replace the traditional Middle Cambrian), is a fairly common trilobite in the lower Amgan of the Yudoma–Olenek facies region, where it defines the lowermost Amgan biozone and thus the base of the Middle Cambrian series in Siberia (e.g., Korovnikov 2001; Shabanov et al. 2008; Naimark et al. 2011). The species is also present in Guizhou, South China, where it occurs in the Ovatoryctocara granulata–Bathynotus holopygus Zone of the Duyunian (Yuan et al. 1997, 2001, 2002). In addition, specimens of O. granulata have been found in the uppermost Brigs Formation of southeastern Newfoundland, Canada, a part of Western Avalonia (Fletcher 2003). The single occurrence at Easter Cove, southeastern Newfoundland, is underlain by shales in which Hamatolenus (Hamatolenus) cf. meridionalis Geyer, 1990 has been found [Fletcher 2006, pl. 27, fig. 35, described as “Hamatolenus (H.) sp. aff. H. (H.) marocanus” (Neltner, 1938)]. Hamatolenus (H.) meridionalis is known from the lower Morocconus Zone of the shaly facies of the Jbel Wawrmast Formation in the western Anti-Atlas (Geyer 1990b). A further constituent of the uppermost Brigs Formation of southeastern Newfoundland is Condylopyge eli Geyer, 1997, a trilobite first described from the Morocconus Zone of the Moroccan Anti-Atlas, and Morocconus notabilis, the index fossil of that Moroccan zone was also found in southeastern Newfoundland in the Brigs Formation (Fletcher 2003, 2006). Therefore, the Morocconus Zone of the Moroccan Atlas ranges and equivalent strata in western Avalonia correlate at least in part with the Ovatoryctocara granulata Zone in Siberia and northern Greenland and the Ovatoryctocara granulata–Bathynotus holopygus Zone of the Duyunian in South China. A detailed discussion is provided by Geyer and Peel (2011). The Kingaspis campbelli and Redlichops faunules of Jordan are also best correlated with the Morocconus Zone, possibly with its upper part. They would thus be equivalents of the O. granulata level with the acme of Bathynotus species and the Paradoxides (A.) muroeroensis Zone of Iberia. If Ovatoryctocara granulata is selected to mark a GSSP for the base of the Cambrian Series 3 and Stage 5, the trilobite occurrences in Jordan would thus indicate the basal strata of those units.

**SYSTEMATIC PALAEONTOLOGY**

Superfamily Redlichioidea Poulsen, 1927
Family uncertain

Genus Redlichops Richter and Richter, 1941

**TYPE SPECIES:** Redlichia (Redlichops) blanckenhorni Richter and Richter, 1941; by original designation.

**DISCUSSION:** As detailed by Rushton and Powell (1998), the traditional systematic position of the endemic monotypic genus Redlichops within the Subfamily Pararedlichiiinae (Zhang 1966; Zhang et al. 1980) cannot be maintained. Clearly distinguishing characters are the moderately wide (rather than distinctly slender) interocular area, the shape of the palpebral lobes, and primarily, the character of the glabella. In particular, the pattern of the glabellar furrows, with S3 distant from the axial furrows, the shallow but widened median portion of the occipital furrow, as well as the parafrontal band and the anterior progression of the eye ridges, are all characters of advanced redlichioids. Taking into account the huge amount of material of Redlichops blanckenhorni collected without a pygidium attributable to the species, we must assume that Redlichops is micropygid. These characters do not allow Redlichops to be placed into an established family with any degree of confidence.

Redlichops blanckenhorni Richter and Richter, 1941
(Text-figs 7 and 8)

1910. Ptychoparia; Schmidt in Blanckenhorn, p. 412.


1912. Ptychoparia; Blanckenhorn, p. 129.

1915. Ptychoparia sp.; Dienemann, p. 25.

v 1941. Redlichia (Redlichops) blanckenhorni n. sp.; Richter and Richter, p. 15–18, pl. 2, figs 1, 2, 5?, 6a (only).

Text-fig. 7. Redlichops blanckenhorni Richter and Richter, 1941; all specimens from the Wadi Uhaymir section. 1-15 – cranidia. 1 – FG–602–003, incomplete cranidium, dorsal view, × 3; 2, 3 – FG–602–009a, incomplete cranidium, dorsal views; 2, entire specimen, × 3; 3, detail, × 5; 4-6 – FG–602–058a, semigerontic cranidium, dorsal, left lateral and anterior views, × 3; 7 – FG–602–024c, incomplete cranidium, dorsal view, × 3; 8 – FG–602–027, incomplete cranidium, dorsal view, × 3; 9, 10, 13 – FG–602–016a, cranidium of young individual, left lateral, dorsal, and anterior views, × 5 each; 11, 12, 14, 15 – FG–602–039a, incomplete cranidium; 11, anterior view, × 3; 12, oblique right lateral view, × 3; 14, dorsal view, × 3; 15, detail, dorsal view, showing granulation of glabella, left fixigena and eye ridge.

Note muscular pits at occipital furrow and close to S1 and swelling on frontal lobe, × 8
Redlichia (Redlichops) blanckenhorni n. sp.; Richter and Richter, pl. 2, figs 3, 6b.

v 1959. Redlichops (R.) blanckenhorni; Poulsen in Harrington et al., p. O201, fig. 141,8.


1998. Redlichops blanckenhorni Richter and Richter, 1941; Rushton and Powell, p. 138–139, figs 7–9, 11–13 (only).


DESCRIPTION OF ADULT MORPHOLOGY: Cranidium with width/length ratio c. 1.3 to 1.4. Anterior margin gently curved. Palpebral lobes extending beyond tips of anterior sections of facial suture, with ocular furrows at about the distance of anterolateral corners from midline.

Glabella more than 75 % of cranidial length and 36 to 39 % of cranidial width across occipital ring: sides straight to S3, tapering progressively forward; frontal lobe curved, with tendency to slightly less curved oblique anterolateral sections; posterior section of frontal lobe faintly expanded due to fusion with projections of eye ridges, maximum width of frontal lobe c. 85 % of cranidial length. Palpebral furrows quite narrow, weakly developed into barely recognizable parafrontal band. Well-preserved specimens exhibiting anteriorly directed caecal lines on these preocular areas, indicating that at least three vascular threads extend from this region into the marginal area of the glabellar front. Caeca rarely preserved in distal portions of fixigenae, with vessels setting off from an arched trunk parallel to palpebral lobe (Text-fig. 7.15).

Palpebral lobes gently curved, subequal in width throughout, generally with moderate transverse convexity, distal portions with tendency to be more raised towards posterior, with most elevated parts close to posterior tips; breadth 18 to 26 % of maximum cranial width, exsagittal length extending to almost half of cranial length. Palpebral furrows quite narrow, weakly defined, well-developed posteriorly to eye ridges. Eye ridges projecting from palpebral lobes, slightly narrower but without change in elevation or transverse furrow; roughly three-times longer than wide; swinging abruptly forward next to glabella, projecting into a shallow lobe, surrounding frontal lobe of glabella; barely recognizable in width and elevation anterior to glabellar front.

Anterior branches of cephalic suture diverging moderately strongly from mid-section of eye ridges, swinging medially in a gentle curvature in anterior third of preocular field to cut anterior border in distinctly oblique angle, located distinctly more ventrally than anterior end of sensory organ.

Axial furrows varying considerably in depth and width, generally shallow, with maximum width next to anterior sectors of occipital ring and adjacent to L2, proceeding into shallow groove around frontal lobe. Fixigenae with low convexity, including longitudinal shallow depression along centre and slightly raised interior areas with highest elevations next to L2 and posterior part of L3; extend across centre of palpebral lobes to c. 50–55 % maximum width of occipital ring; exsagittal length slightly less than 40 % (36–39 %) of cephalic length next to axial furrows. Highest elevation projecting into faint lobe, fused with anterior part of L3 and developed into barely recognizable parafacial band. Well-preserved specimens exhibiting anteriorly directed caecal lines on these preocular areas, indicating that at least three vascular threads extend from this region into the marginal area of the glabellar front. Caeca rarely preserved in distal portions of fixigenae, with vessels setting off from an arched trunk parallel to palpebral lobe (Text-fig. 7.15).

Palpebral lobes gently curved, subequal in width throughout, generally with moderate transverse convexity, distal portions with tendency to be more raised towards posterior, with most elevated parts close to posterior tips; breadth 18 to 26 % of maximum cranial width, exsagittal length extending to almost half of cranial length. Palpebral furrows quite narrow, weakly defined, well-developed posteriorly to eye ridges. Eye ridges projecting from palpebral lobes, slightly narrower but without change in elevation or transverse furrow; roughly three-times longer than wide; swinging abruptly forward next to glabella, projecting into a shallow lobe, surrounding frontal lobe of glabella; barely recognizable in width and elevation anterior to glabellar front.

Anterior branches of cephalic suture diverging moderately strongly from mid-section of eye ridges, swinging medially in a gentle curvature in anterior third of preocular field to cut anterior border in distinctly oblique angle, located distinctly more ventrally than anterior end of sensory organ.
of visual surface. Posterior branches short, diverging strongly laterally and ventrally, with lateral tips of posterior border extending to nearly the level of palpebral lobes’ maximum distance from midline.

Preglabellar field sagittally c. 6–7 % of cephalic length, with median plectrum becoming more and more inconspicuous during ontogeny, incompletely fused with parafrontal band, and creating an indentation of posterior margin of anterior border (Text-fig. 7.14). Precocular fields weakly convex, sloping ventrally, occasionally with faint and fairly irregular caeca.

Anterior border sagittally of c. 13 to 15 % of cephalic length, defined posteriorly by relatively steep slope toward border furrows, medial surface of low convexity. Anterior border furrow a relatively narrow, poorly defined groove. Posterior border convex, weakly sigmoidal, of more-or-less equal breadth throughout. Posterior border furrow obsolescent.

The surface of cranidium with different types of ornamentation/prosopon. Anteriorly and laterally directed convex parts of glabella covered with fine- to medium-sized granules, as well as anterior portions of eye ridges. Ocular areas, precocular fields, anterior and posterior border generally covered with fine granules. Parts of glabella, adjacent to lateral and occipital furrows and proximal portions of palpebral lobes, smooth. Distal and medial part of anterior border with terrace lines, forming long scarps parallel to anterior cephalic margin in anterior position and becoming progressively shorter rearward.

On the exterior of the test, specimen FG–602–039a shows traces of muscle scars as imprints of the ventral surfaces. These scars are all pairs with symmetric arrangement in the glabella, situated on the anterior rim of the occipital ring, on both sides of S1 and S2, and as unpaired scars on the glabellar midline in L3 and the frontal lobe (Text-fig. 7.15).

Librigena with narrow extraocular area, about as wide as or slightly wider (tr.) than adjacent part of lateral border. Lateral border extends into a long genal spine (in adults at least as long as the anterior part of the librigena). Anterior and posterior sections of facial suture in accordance to that of the cranidium. Narrow ocular socle present along ocular suture. Lateral border and genal spine with longitudinal terrace lines, extraocular area with radial caeca. Doublure completely underlies borders, densely covered with longitudinal terrace lines.

Thoracic segments known only from small fragments; pleurae apparently almost twice the width of the axial rings. Pleural furrow distinctly delimited toward the anterior and less distinctly toward the posterior. Posterior rim of the pleurae more or less straight, curving slightly into an apparently moderately long spine.

ONTOGENETIC VARIABILITY: Small holaspids individuals of Redlichops blanckenhorni differ from large adult specimens primarily in the relative proportions and in the expression of furrows. The most conspicuous differences of the juveniles (compared to adults) are:

Cranidium transversely wider (compared to adults), length less than three-quarters width. Glabella wider (compared to adults), width across occipital ring more than 35 % cranial width, with sides subparallel or weakly tapering forward (in undeformed, non-flattened specimens), more than three-quarters cephalic length. Occipital ring sagittally broader (up to 20 % cephalic length). Glabellar furrows shallower, less clearly impressed.

Fixigenae less structured morphologically, sunken between palpebral lobes and glabella, with only minor bacculae. Palpebral lobes long (exsag. length more than 40 % cephalic length) and broad (width tr. more than 8 % cranial width across centre of palpebral lobes); eye ridges broad. Preglabellar field narrow, often developed as a sunken area created by confluent axial and anterior border furrows. Anterior border wide (up to more than 15 % cephalic length), highly convex.

Late larval stages of Redlichops blanckenhorni have cranidia with a subparallel to slightly tapering glabella and with transverse, medially connected furrows, its frontal lobe reaching the anterior border furrow; occipital ring extending into a stout thorn; clearly convex fixigenae with an interior bacculate section and a more or less clearly visible oblique depression; palpebral lobes with posterior tips at the level of L1 or even more anteriorly and anterior tips laterally of the frontal lobe; short, subnodular eye lobes; anterior border narrow, thread-like.

DISCUSSION: For affinities of Redlichops blanckenhorni see discussion under Genus Redlichops.

Rushton and Powell (1998, fig. 10) described and figured a rostro-hypostomal plate, which they attributed to Redlichops blanckenhorni. This specimen would have placed the species under trilobites with conterminant hypostomal condition. However, the specimen must be attributed to Myopsolenites palmeri (Parnes, 1971) (see below for discussion).

Superfamily Ellipsocoephaloidea Matthew, 1887
Family Ellipsococephalidae Matthew, 1887
Timnaella Parnes, 1971
TYPE SPECIES: Timnaella eyali Parnes, 1971; by original designation.

DISCUSSION: The genus Timnaella was introduced by Parnes (1971) for small ellipsocephaloid-type trilobites from the Timna area, southern Negev. Parnes (1971) introduced two formal species (T. eyali and T. wingi) and referred additional material to as Timnaella sp. 1 and Timnaella? sp. 2. All of his material came from the Hakhil Formation and is poorly preserved in a relatively coarse, rose-coloured to pinkish sandstone so that the details of the furrows and the surface ornament are unknown. Timnaella eyali is known from one and T. wingi from two incomplete cranidia. The additional material is even less perfectly preserved and partly characterizes immature individuals so that some may indeed belong to T. eyali and T. wingi, respectively. Both species are in need of additional material for a more precise characterization.

The species first described as Protolenus orientalis by Picard (1942) from the Wadi Numayri, Jordan, and transferred to Resserops (Richterops) orientalis by Parnes (1971) shares most significant characters with Timnaella eyali and is thus transferred to Timnaella with certain caveats (see discussion below).

Timnaella? orientalis (Picard, 1942)

v 1942. Protolenus orientalis sp. nov.; Picard, p. 1–4, pl. 2, fig. 1.
non 1942. Protolenus orientalis sp. nov.; Picard, pl. 2, fig. 2.

v 1971. Resserops (Richterops) orientalis (Picard); Parnes, p. 186, 188–189, pl. 1, figs 1–4.
non 1971. Resserops (Richterops) orientalis (Picard); Parnes, pl. 1, fig. 5 (only).

? 1971. Strenuella (S.) sp. 2; Parnes, p. 193–194, pl. 1, figs 10, 11.
1976. Resserops (Richterops) orientalis; Cooper, p. 273.

DISCUSSION: The history of Timnaella? orientalis (Picard, 1942) is complicated and has not yet led to a satisfactory understanding of the species. Picard (1942) figured only a single cranidium as a simple line drawing (Picard 1942, pl. 2, fig. 1) with obvious minor inaccuracies that made an unequivocal determination difficult. The incomplete pleura figured by Picard (1942, pl. 2, fig. 2) as a second specimen of “Protolenus orientalis sp. nov.” shows the ventral surface and probably is a pleura of Myaspsoleslites. Parnes (1971) restudied and refigured Picard’s type material, which gives a fairly good impression of the morphology, despite the imperfect preservation in oolitic limestone and Parnes’ photograph with very shallow incident light. One of the authors (GG) was able to examine this original material in the collections in Jerusalem.

The holotype cranidium is preserved as part and counterpart, with the positive being an internal mould. The counterpart has some portions of the recrystallized shell and steinkern adhering to the exterior surface. The cranidium is fairly small and has a long, tapering glabella, the frontal lobe of which reaches the anterior border furrow. The glabellar furrows appear to be directed backward, with S3 not clearly visible. The occipital furrow is transglabellar. The occipital ring carries a median node.

Palpebral lobes are long, their posterior tips reaching or nearly reaching the posterior border furrow; gently arched, defined by a distinct palpebral furrow, projecting into fairly short, oblique eye ridges. Fixigenae comparatively narrow (tr.), about 60 % of maximum glabellar width, traversed by a shallow and broad, slightly curved depression, which creates a slightly raised proximal portion close to the axial furrows and a less raised narrow lunar ridge close to the palpebral furrows.

Preocular areas of exsagittal are narrow, sloping anteriorly. Anterior border furrow a distinct groove. Anterior border narrow, distinctly raised to form a brim-like structure.

DISCUSSION: Timnaella? orientalis (Picard, 1942), originally described as a species of the formerly collective genus Protolenus Matthew, 1887b, is clearly distinguished from that genus by numerous characters, most of all the different type of glabella (see Geyer 1990b and Westrop and Landing 2000, for detailed description of Protolenus). Parnes (1971) transferred the species to Resserops (Richterops), which is now treated as Perrector (Richterops). Indeed, the cranidium is quite reminiscent of species of this subgenus, but the similarities are superficial and of little taxonomic significance. The genus and subgenus includes fairly large redlichioiid trilobites with a long thorax and at least one macropleural segment as well as a pygidium of comparable size to the cranidium. In all of the species, the anterior border is quite broad (sag.) and bar-like because it overlies a distinct rostral plate, to which the hypostome is attached (see holotype of Perrector (Richterops) falloti (Hupé, 1953) in Hupé 1953a or Chang et al. 1997, fig. 293.3). In addition, the palpebral lobes in the Resseropinae are relatively broad and have a distinct adaxial curvature close to the posterior tips,
and a transverse depression of the type seen in *Timnaella orientalis* is also unknown among the Resseropinae.

Rushston and Powell (1998) placed the species tentatively under *Realalaspis*, emphasizing the similarity of the palpebral lobes, and also noted some resemblance to *Hesa problematica*.

The species cannot be assigned to any known genus with certainty. The visible characters show a fairly good match with those seen in *Timnaella eyali* and *T. wingi*, despite the differences in preservation. Fundamental differences can be seen in the quite narrow (sag.) anterior border, the apparently quite prominent anterior border and the narrower fixigenae with a more clearly developed depression. *Timnaella eyali* has a similarly narrow anterior border, which appears to be less upturned, a less clearly tapering glabella, and wider fixigenae with a better developed exsagittal convexity. In addition, the connection of the palpebral lobes with the eye ridges shows a small angular change. *Timnaella wingi* has wider fixigenae and a broader preglabellar depression. Imperfectly preserved material of immature cranidia figured by Parnes (1971, pl. 1, figs. 10, 11) as *Strenuella (S)* sp. 2 possible represents the same species.

*Timnaella? cf. orientalis* (Picard, 1942)
(Text TX.9.4, 9.5, 9.8–9.10)

**MATERIAL:** Approximately 7 cranidia, reposited under FG–602–007a, FG–602–014b, FG–602–023, FG–602–24b, FG–602–051b. All from the Wadi Uhaymir section, Jordan.

**DESCRIPTION:** Cranidium with width/length ratio c. 0.75. Anterior margin gently curved, with median arccuation. Palpebral lobes extending beyond tips of anterior sections of facial suture. Glabella length c. 80 % of cranidial length in adult individuals (up to more than 90 % in immature individuals) and c. half cranidial width across occipital ring in adult individuals (c. 40 % in immature individuals); tapering slightly forward, with approximately straight sides; frontal lobe with shallow arccuation in dorsal view; glabella moderately convex, with tendency to form a median longitudinal ridge in transverse section. Three pairs of lateral glabellar furrows, all of them shallow; S1 moderately long, rearwardly curved; S2 moderately long, thin furrow more-or-less normal to axis; S3 very short and at the anterolateral corners of glabella. Occipital furrow consisting of moderately well impressed, faintly arcuate lateral furrows with a tendency to bifurcate; median section obsolescent. Occipital ring length sagittally up to c. 15 % of cephalic length, with moderately curved posterior margin and subterminal node (imperfectly preserved on the specimens).

Axial furrows varying in depth, usually moderately deep and moderately wide. Fixigenae composed of slightly convex crescentic portions adjacent to axial furrows and to palpebral furrows, separated by shallow and weakly defined slightly curved longitudinal depression; width across centre of palpebral lobes up to three-quarters maximum width of occipital ring; exsag. length next to axial furrows c. 45–50 % of cephalic length.

Palpebral lobes moderately long, c. 40 % of cephalic length in adult specimens (up to c. 48 % in immature individuals), gently curved at ocular suture, about equal in transverse width throughout, transversely convex, but more strongly sloping toward palpebral furrow; anterior tips at level of S2 in adult specimens, up to level of S3 in young individuals; posterior tips at anterior level of occipital ring, short distance from posterior border. Palpebral furrows moderately well developed. Eye ridges project from the anterior ends of palpebral lobes obliquely forward, delimited from palpebral lobes by a shallow transverse depression, relatively short; develop into two faint projections at axial furrow of which the posterior thread appears to be faintly connected with anteriormost part of L3 at anterolateral corners of the glabella, the anterior thread develops into a low indistinct parafrontal band, which is well separated from the frontal lobe.

Anterior branches of the cephalic suture not well preserved, clearly diverging from anterior ends of ocular suture, curve inward at anterior border furrow to clip off a considerable part of the anterior border. Posterior branches not preserved, short.

Preglabellar field sagittally very narrow to absent. Preocular fields narrow, slightly convex areas sloping ventrally from eye ridges. Anterior border clearly convex, well defined in small individuals, less so with growing size, length sagittally c. 5–8 % cephalic length. Anterior border furrow well defined in front of preocular areas, merely a change in convexity in front of glabella. Posterior border with low to moderate convexity, with double curvature from axial furrows, sloping ventrally posterior to palpebral lobes. Posterior border furrow a shallow groove, poorly delimited from fixigenae.

Librigena known only from immature, imperfectly preserved specimens. Extraocular area narrow. Lateral margin fairly weakly curved. Lateral border convex, extended into a long genal spine along ocular suture. Posterior margin of librigena broad (tr.), forms an acute angle with adaxial margin of genal spine.

Thorax and pygidium not known with certainty.
Surface of the cranidium in adult specimens covered by small granules except for glabellar furrows, axial furrows, and palpebral lobes. Small individuals smooth. DISCUSSION: The material described above probably represents the species described as *Protolenus orientalis* by Picard (1942) (see discussion above).
The type material from Israel, however, is imperfectly preserved in a way that does not allow the well preserved material from the Wadi Uhaymir section to be placed with certainty under the same species.

Subfamily Ellipsocephalinae Matthew, 1887

*Issalia* gen. nov.

**TYPE SPECIES:** *Issalia scutalis* gen. nov., sp. nov.

**ETYMOLOGY:** Named after the Wadi Issal area, Jordan, in which the type material was collected.

**DIAGNOSIS:** Genus of the Subfamily Ellipsocephalinae with glabella c. 80 % cephalic width, tapering forward, frontal lobe moderately wide, subtruncate; three pairs of weakly impressed, roughly parallel and progressively less well developed lateral glabellar furrows; fixigenae weakly convex, with faint oblique depression; palpebral lobes long, reaching toward posterior border furrow, separated from eye ridges by weak depression; eye ridges meet dorsal furrows at level of S3; preglabellar field, preocular areas and anterior border fused to form a wide platform with the anterior border slightly raised toward the anterior margin.

**DISCUSSION:** *Issalia* gen. nov. is a typical genus of the Ellipsocephalinae, with the basic pattern of three pairs of lateral glabellar furrows, a slightly tapering glabella, long palpebral furrows which are slightly arched against the eye ridges, and inconspicuously subdivided fixigenae. These characters create the impression that the type species could be grouped among trilobites dealt with under classic genera such as *Lusatitops* Richter and Richter, 1941, *Pseudolenus* Hupé, 1953a, *Latoucheia* Hupé, 1953a, *Proamphyx* Frech, 1897 or *Comluella* Hupé, 1953a. However, careful examination of this notoriously difficult group has shown that morphoclines towards a progressive smoothing occurred independently at least twice in the traditional late Early to early Middle Cambrian (Geyer 1990b) and that the distinction of exterior morphology from internal moulds as well as the longitudinal morphology across the eye ridges are crucial issues. In this context, *Issalia* does not belong to the ellipsocephaloid-kingaspidoid clade, as clearly indicated by the morphology of the frontal lobe, being slightly narrower (tr.) than L3 and with sub-rounded to obliquely truncated anterolateral corners. A major feature that distinguishes *Issalia* from other non-ellipsocephaloid-kingaspidoid trilobites is the platform or brim-like anterior part of the cranidium composed of the superficially fused preglabellar field, preocular areas and anterior border, with the anterior border weakly raised towards the anterior.

*Issalia* gen. nov. resembles the Moroccan genus *Planolimbus* Geyer, 1990b from the lower Middle Cambrian of the Anti-Atlas Mountains. However, it clearly derives from kingaspidoid ancestors and can be distinguished easily from the latter by a number of relevant characters, primarily the pattern of strongly arched lateral glabellar furrows. A closer systematic affinity appears to exist to *Timnaella* Parnes, 1971, as illustrated by *T. eyali* Parnes, 1971. However, the genera are easily distinguished by the almost subtruncate glabella and the low convexity of the anterior cranidial region in *Issalia*.

*Issalia scutalis* gen. nov., sp. nov.  
(Text-fig. 9.1–9.3)

**ETYMOLOGY:** From the Latin *scutum*, shield; a reference to the unit resulting from the fused preglabellar field, preocular areas and anterior border and to the low overall convexity of the cranidium.

**HOLOTYPE:** Incomplete cranidium, FG–602–095a.

**PARATYPES:** Incomplete cranidium, FG–602–107, and single librigena, FG–602–080. All material from Wadi Issal, Jordan.

**DESCRIPTION:** Cranidium with width/length ratio of about 0.8. Anterior margin gently curved. Palpebral lobes slightly extending beyond tips of anterior sections of facial suture. Glabella c. 80 % cranidial length and c. 45 % cranial width across occipital ring; slightly tapering forward, with approximately straight sides; frontal lobe about one-third maximum cranidial width, with semitruncate anterior margin; glabella moderately con-
vex, with tendency to have a median arcuation anteriorly in transverse section. Three pairs of lateral glabellar furrows, all of them shallow to moderately well impressed; S1 and S2 moderately rearward directed and fairly long, separated at short distance over the midline; S3 a short depression close to the axial furrows. Occipital furrow moderately well impressed and moderately wide, with a faint median section. Occipital ring sagittally up to 20 % cephalic length, with a moderately curved posterior margin and a probably short subterminal spine (only imperfectly preserved on the specimens).

Axial furrows varying from shallow next to proximal areas of fixigenae to moderately deep between eye ridges and glabella. Fixigenae with small bacculae on posteroaxial proximal portions and weakly raised distal portions paralleling palpebral lobes, separated by shallow and weakly defined longitudinal depression; width across centre of palpebral lobes half maximum width of occipital ring; exsag. length next to axial furrows in the order of 40 % cephalic length.

Palpebral lobes moderately long (c. 40 % cephalic length), gently curved at ocular suture, transverse width about equal throughout, with distinct transverse convexity; anterior ends at level of S3 or slightly behind, posterior tips at mid level of occipital ring, almost reaching posterior border. Palpebral furrows moderately well developed. Eye ridges project obliquely forward from anterior ends of palpebral lobes, delimiting from palpebral lobes by a shallow transverse depression, swing forward as low, poorly defined ridges; posterior section of eye ridges transversal furrows as a low swelling to meet anteriormost part of L3.

Anterior branches of the cephalic suture not well preserved, but clearly diverging from anterior ends of ocular suture, curve inward at anterior border furrow to clip off a considerable curvature. Posterior branches not preserved, short.

Preglabellar field a sagittally narrow area, grading into the anterior border without a clear definition. Preoccular fields moderately wide, slightly convex areas sloping ventrally from eye ridges.

Anterior border a low, slightly convex area, length sagittally c. 5 % cephalic length. Anterior border furrow a comparatively broad, poorly defined depression. Posterior border convex, apparently more-or-less straight throughout. Posterior border furrow a quite shallow groove, poorly defined from fixigenae.

Librigena with very narrow extraocular area, which is narrower (tr.) than adjacent part of lateral border. Lateral border slightly convex, poorly delimited from border furrow, extends into a fairly long genal spine (about as long as the anterior part of the librigena). Anterior section of facial suture in accordance to that of the cranium, swings adaxially at border furrow to describe a considerable curvature. Posterior section of facial suture short, almost straight and almost normal to axis. Narrow eye socle along ocular suture.

Thorax, and pygidium unknown.

Surface of the cranium smooth except for small terrace ridges on anterior border. Preglabellar field and preoccular areas traversed by faint caeca originating from eye ridges and arranged in a “centrifugal”, pseudopalmate pattern. Librigena smooth on extraocular area, with fine terrace ridges on lateral border.

DISCUSSION: See discussion under genus.

_Tayanaspis_ gen. nov.

**TYPE SPECIES:** _Tayanaspis bulbosus_ gen. nov., sp. nov., by monotypy.

**ETYMOLOGY:** After Wadi Tayan, Jordan, where most of the present material was collected.

**DIAGNOSIS:** Genus of the Subfamily Ellipsocephalinae with glabella moderately long, tapering forward, frontal lobe narrow; fixigenae sublongitudinally divided by arched furrow, with adaxial portion conspicuously swollen in adult individuals, abaxial portion faintly convex; palpebral lobes and eye ridges fused to form arched lobe, fused with frontal lobe and with branch proceeding into parafrontal band; preglabellar field and preoccular areas forming continuous groove; anterior border more-or-less equally curved convex brim.

**DISCUSSION:** Rushton and Powell (1998, p. 139) had a limited amount of material of the type species of this new genus and assigned it to the genus _Realaspis_ Sdzuy, 1961 without referring to a described species or introducing a new species. In their discussion, they particularly emphasized the “broad interocular area of the fixigenae and the short palpebral lobe” and also the morphological similarity of the pygidia with those of the Subfamily Metadoxinae. These were taken as arguments for the placement of the species, albeit the authors were the first to suggest a placement of _Realaspis_ under the Metadoxinae. In addition, Rushton and Powell (1998) stressed the similarity of the cranium with that of _Realaspis strenoides_ Sdzuy, 1961, the type species of _Realaspis_, but emphasized marked differences. Among the differential characters mentioned by Rushton and Powell (1998), particularly the absence of a swollen area on the fixigena and a rather moderate relief in the cranium, are such obvious differences in the character set that a close similar-
ity between the type species of Realaspis and the new species described here can be excluded.

In fact, Realaspis is based on poorly preserved material from the Sierra Morena region in southern Spain (Sdzuy 1961). The holotype, a distorted internal mould of a cranidium, does not allow recognition of the details of the exterior, with the glabella having lost nearly all furrows. What is recognizable, however, is that the palpebral lobes are much shorter than in the Jordanian material, distinctly oblique to the axis, broader than the low eye ridges, and located on a slight abaxial swelling of the fixigenae. Thus, the Spanish form is certainly not congeneric with the Jordanian one. A close phylogenetic relationship with Tayanaspis is at best indicated only by the similarity of the pygidium, albeit notable differences exist in the number of segments (as shown by the axial furrows) and the overall shape with fairly angular anterolateral corners in Realaspis. It should be mentioned that the late Klaus Sdzuy himself stated that he was unhappy with the taxon after having introduced it and conceded that the much retouched photos in the original publication did not give a realistic impression of the morphology of the type species (oral comm. to GG, November 1995). Additional species, which were later attributed to Realaspis, suffer from this same deficiency and, in fact, a morphologically fairly heterogeneous assemblage of species has been included under this genus.

Despite the question of a close relationship between Tayanaspis and Realaspis, they are certainly not closely related with Metadoxides Bornemann, 1891, and thus are not members of the Subfamily Metadoxidinae. Metadoxides belongs to a small group of lower Cambrian trilobites characterized by a tapering glabella, fairly short palpebral lobes continuing into eye ridges which are broadly attached to the frontal lobe of the glabella, fairly wide (tr.) pleurae without much transverse convexity, and a mostly triangular pygidium continuing the rearward tapering of the posterior thorax. Only its taxa from Sardinia allow the Subfamily Metadoxidinae to be characterized with sufficient morphological criteria. As a consequence, most of the genera shown in the revised Treatise volume (Chang et al. 1997) appear to be wrongly placed. This accounts primarily for Onaraspis Opik, 1968 (see discussion below under Bathynotidae), Churkinia Palmer, 1968, and Minusinella Repina, 1960 (in Khalfin 1960).

Tayanaspis bulbosus gen. nov., sp. nov.
(Text-figs 9.11, 10 and 11)


11 – FG-602-037a, fragment of thoracic segment, dorsal view, × 5; 12, 15 – FG-602-001, incomplete pygidium, dorsal and posterior views, × 4; 13 – FG-602-025d, pygidium, partly exfoliated, dorsal view, × 6; 14 – FG-602-057c, incomplete librigena, dorsal view, × 6; 16 – FG-602-025a, pygidium, partly exfoliated, dorsal view, × 6; 17 – FG-602-018b, partial pygidium, internal mould, dorsal view, × 6
terior rim, forming very low bacculae-type swellings close to the posterior tips of the palpebral lobes; extending across centre of palpebral lobes to c. 60 % maximum width of occipital ring; exsag. of about one-third cephalic length next to axial furrows.

Palpebral lobes gently curved at ocular suture, slightly reduced in width towards anterior and posterior tips, generally with moderate transverse convexity, distal portions tend to be more raised towards the posterior, with about equal elevation throughout; breadth 22 to 24 % of maximum occipital width, exsagittal length about 25 % cranidial length; anterior ends at level of S3, posterior tips at about level of occipital furrow, separated by short distance from posterior border. Palpebral furrows shallow, weakly defined topographically but distinguished from distal area of fixigenae by lack of granules. Eye ridges project from palpebral lobes, together forming a fairly homogeneous area, slightly narrower and progressively less elevated to fade towards axial furrows; roughly three times longer than wide; anterior section of eye ridges swings forward next to glabella to project into a shallow lobe that surrounds the frontal lobe of the glabella, barely recognizable in width and elevation anterior to glabellar front on the exterior of the test.

Anterior branches of facial suture diverge moderately strongly from anterior ends of visual surface and swing medially in a gentle curvature in the anterior third of the preocular field to cut anterior border at a distinctly oblique angle. Posterior branches fairly short, forming a gentle curve, but only slightly diverging in ventral view because of a strong ventral deflection, with lateral tips of posterior border extending to nearly the level of the palpebral lobes’ maximum distance from midline.

Preglabellar field sagittally narrow, only c. 3 % cephalic length, developed as a concave groove in sagittal section, with faint median plectrum. Preocular fields slightly concave, sloping ventrally in the distal one-third.

Anterior border highly convex, sagittal length c. 10 % cephalic length, defined posteriorly by relatively steep slope towards border furrow. Anterior border furrow developed only as a steep slope. Posterior border convex, weakly sigmoidal, of more-or-less equal breadth throughout, but with highest elevation posterior to distal one-third of fixigenae. Posterior border furrow fairly shallow, poorly defined anteriorly.

Librigena with narrow extraocular area, about as wide as or slightly wider (tr.) than adjacent part of lateral border. Lateral border extends into a moderately long genal spine (shorter than the anterior part of the librigena). Anterior and posterior sections of facial suture in accordance to that of the cranidium. Narrow eye socle along ocular suture. Lateral border and genal spine with longitudinal to fairly oblique terrace lines. Doulure completely underlies borders.

Thorax not known from a complete adult, of at least 12 segments. Axial rings strongly convex in transverse section. Median portion sagittally convex, with distinct, prominent median node. Anterior margin of axial ring generally straight or slightly sigmoidal, but with distinct rearward bend in segment 1. Posterior margin generally straight, with a distinct forward curvature toward axial furrows.

Pleurae with anterior margin straight and more-or-less normal to axis up to a stout fulcral process about three-fifths the distance between dorsal furrows and pleural tips, curving rearward abaxially to the fulcral process. Anterolateral portion forms strongly deflected facet abaxial to fulcral socket. Posterior margin of pleurae slightly curved, distal portion with slight forward swing to form shallow concave area at the base of the pleural spine, with small dorsoventral flexure about to two-thirds the distance between dorsal furrow and pleural tip that forms an indistinct socket corresponding to the fulcral process of the adjacent tergite. Except for tergite 1, fulcral sockets and corresponding processes are in a progressively adaxial position in posterior tergites. Pleural furrows well impressed, well defined, widest close to axial furrows, progressively decreasing abaxially to form slender, slightly curved triangular areas; abaxial tip slightly distant from base of pleural spine. Pleural tip formed by a posterior curvature of the anterolateral margin, producing a fairly long, obliquely directed pleural spine that grows progressively rearward. Pleurae of first tergite slightly modified, with short abaxial portions terminated by fulcral processes corresponding to cephalic posterior border. Abaxial part of pleurae at first thoracic tergite with strong deflection of a moderately large facet.

Pygidia from the same beds as the cranidia are assigned to the same taxon. These pygidia have comparatively strongly curved anterior and posterior rims. The length/width ratio is usually slightly less than 1.0. The highly convex axis is stout and tapers rearwards. It has a length of about 83 to 85 % of the pygidial length. Its terminal axial piece is well rounded. Apart from the sagittally very narrow articulating half-ring, one axial ring is defined by a clearly visible furrow, an additional axial furrow faintly impressed. The width of the axis is less than one third of the pygidial width. The highly convex pleural fields have three pairs of interpleural furrows. A very shallow plectrum between terminal axial piece and posterior border counterfeits an additional pair of interpleural furrows. Pleural furrows are lacking.
The border furrow is shallow and weakly delimited. It becomes slightly deeper where it joins the interpleural furrows. The border is low, its surface broad and weakly convex. Behind the posterior axial piece the border has a weak indenture or at least a straight, clipped portion.

Surface of the cranidium with different types of ornamentation/prosopon. Highly elevated parts of glabella, palpebral lobes, and proximal areas of fixigenae smooth; lateral parts of glabella, distal areas of fixigenae, posterior border and border furrow, preglabellar field and preocular areas densely covered with fine to medium-sized granules. Distal and medial part of anterior border carries well developed terrace lines roughly parallel to the anterior margin. Thoracic axial rings and pleura (except for smooth pleural furrow) covered with granules, pleural spines with terrace ridges (Text-fig. 11.11).

ONTOGENY: Tayanaspis bulbosus gen. nov., sp. nov. shows a striking ontogenetic development, which is mainly created by a change of the fixigenal morphology and, less clearly, of the glabella and the anterior border. Starting at meraspids with normal, distinctly convex fixigenae (Text-figs 10.12, 10.13), there is a tendency during individual development to reduce the convex genae to narrow, lenticular bulbous lobes close to the dorsal furrows, whereas the rest of the fixigenae forms a nearly flat platform or even develops weakly convex lobes adjacent to the palpebral furrows (Text-fig. 10.8). In early meraspids, the glabella is nearly parallel-sided and with a distinct occipital ring (Text-fig. 10.12; no occipital node!). It becomes progressively more tapered with a rounded frontal lobe. Fully grown adult specimens have a tapering glabella with slightly curved sides and a somewhat flattened frontal lobe with subangular anterolateral corners (Text-fig. 10.6). The anterior border starts as an inconspicuous, relatively narrow (sag.) blade and grows in breadth during growth. It finally develops a faint kink in front of the glabella. Palpebral lobes and eye ridges form subevenly homogeneous, subequally curved, highly convex lobes of subequal width (Text-fig. 10.14) early in the ontogenetic series. However, the eye ridges become progressively narrower than the palpebral lobes in large individuals (Text-fig. 10.6, 10.7).

Interestingly, the small holaspis specimen FG–602–042a shows a small but distinct remnant of an intergenal spine (Text-fig. 10.7).

The Wadi Uhaymir locality yielded a number of small complete specimens preserved as external moulds in silty shale. Because of preservational reasons, details are often difficult to recognize. Nevertheless, the number of thoracic segments in these late meraspid to earliest holaspis specimens appears to be constantly 11 or 12. Text-fig. 11.7 shows a typical specimen with the pleural spines growing progressively in length rearward to attain a length of several times the length of the rest of the pleura in the posterior half of the thorax and with a progressive rearward direction. Further noteworthy is the extreme forward curvature of the extended posterior border so that the genal angle in this specimen is opposite L2 and includes an acute angle.

DISCUSSION: The most conspicuous character of Tayanaspis orientalis is the pronounced swelling on the fixigenae. Rushton and Powell (1998) referred to line-drawings of an as yet undescribed bigotiid species presented by Sdzuy (1978) from about the lowermost trilobite-bearing strata of the Moroccan Atlas ranges. This material has only superficial similarity, being minute, micropygid and having a quite different type of glabella. Much more worthy of comparison are the Moroccan ellipsocephaloid genera from the latest early Cambrian/late Banian Sectigena Zone of Morocco, such as Sectigena Geyer, 1990b or Termierella Hupé, 1953a. Both share with Tayanaspis the distinct swelling on the fixigenae, but have additional swollen areas adjacent to the palpebral lobes and the anterior border. Remarkably, similar genera with swollen portions of the fixigenae lived contemporaneously on other Cambrian continents, such as Chorbusulina Lazarenko, 1962 and Charaulaspis Lazarenko, 1962. However, all of these genera are obviously micropygid and probably represent a convergent morphological evolution.

Hesa Richter and Richter, 1941

TYPE SPECIES: Hesa problematica Richter and Richter, 1941, by original designation.

Hesa problematica Richter and Richter, 1941

DISCUSSION: Richter and Richter (1941, p. 18–19, figs 7–11) described and figured fragmentary trilobite material on which they based a new genus and species termed Hesa problematica. This material included a fragment of a large cranidium, an incomplete librigena, and fragmentary thoracic segments. The material, all collected by Blanckenhorn in 1908, came from the mysterious “Chirbet el-Bursch” locality and was partly referred to as fragments of Paradoxides by Blanckenhorn and Schmidt (see Blanckenhorn 1912, p. 411 and 412, respectively). The Richters designated the fragmentary cranidium as the holotype. This specimens preserves only slightly more than the frontal lobe of the glabella, part of the eye.
ridge, the preglabellar field and part of the preocular areas and anterior border. Investigation of the type material suggests that the reconstruction by Richter and Richter (1941, pl. 2, fig. 9) is incorrect. The Richters (1941, p. 19) referred to the fragment of a posterior part of a glabella which was not figured but present in the repository. They noted that all parts show a distinct granulation, including the occipital furrow. In fact, the well impressed parts of the occipital furrow are devoid of granules, and the different parts show some variation in the size and density of the granulation. It appears that the main reason for assigning the different sclerites like parts of a puzzle to the same species was based on the granulation and their large size.

In fact, the holotype is too insufficiently preserved to be useful for defining a taxon. The other sclerites almost certainly do not belong to the same species and were erroneously assigned to Hesa problematica. It is therefore suggested herewith to restrict Hesa problematica to its holotype. The holotype was re-illustrated by Rushton and Powell (1998, fig. 14).

Kingaspis Kobayashi, 1935

**TYPE SPECIES:** Anomocare campbelli King, 1923; by original designation.

Kingaspis campbelli (King, 1923)  
(Text-fig. 12)

1923. Anomocare campbelli; King, p. 511, figs 2, 3.
1935. Kingaspis campbelli [sic] (King); Kobayashi, p. 196, 305, pl. 23, figs 9, 10
v 1941. Palaeolenus campbelli King [sic]; Richter and Richter, p. 19–20, pl. 2, figs 12–16.
1941. Anomocare campbelli King; Picard, p. 3.
1953. Kingaspis campbelli (King); Hupé, text-fig. 63A.
non 1953. Kingaspis campbelli (King); Hupé, p. 254, pl. 9, figs 1–3.

v 1971. Kingaspis campbelli (King); Parne, p. 201–202, pl. 2, figs 28, 29.
1989. Kingaspis campbelli (King); Gil Cid and Jago, p. 93–94.
v 1990. Kingaspis campbelli (King 1923); Geyer, p. 104–105, pl. 15, fig. 11, pl. 17, figs 8–10.
non 1999. Kingaspis campbelli (King, 1923); Dies et al., p. 221, 222, figs 1, 2.
non 2004. Kingaspis campbelli (King, 1923); Dies et al., p. 284.
non 2004. Kingaspis campbelli; Liñán et al., p. 35.
v 2007. Kingaspis campbelli (King, 1923); Shinaq and Elicki, p. 255, 256, 261, 263, fig. 7.3, 7.4.
2007. Kingaspis (Kingaspis) campbelli (King, 1923); Gozalo et al., p. 366.
non 2007. Kingaspis (Kingaspis) campbelli (King, 1923); Gozalo et al., p. 364, 365, 367, figs 4A, 5F.
2007. Kingaspis campbelli (King, 1923); Dies Álvarez et al., p. 420, 426.
non 2008. Kingaspis (K.) campbelli; Gozalo et al., p. 143.
non 2008. Kingaspis (Kingaspis) campbelli King, 1923; Liñán et al., p. 12, 16, 28, 34, 36, figs 4, 12, 15, 17.
2009. Kingaspis campbelli (King, 1923); Żylińska and Szczepanik, p. 421, 423.
2011. Kingaspis (Kingaspis) campbelli; Gil Cid et al., p. 54, 55, 56.

DESCRIPTION OF ADULT MORPHOLOGY: Cranidium with width/length ratio of about 0.85. Anterior margin gently and evenly curved. Palpebral lobes extending slightly beyond tips of the anterior sections of the facial suture, anterior end of palpebral lobes opposite or slightly posterior to S3. Cranidium of fairly even convexity, similar to a spherical segment, with only glabella slightly elevated, palpebral lobes somewhat upturned, occipital lobe developed as low transverse ridge, and fixigenae less clearly convex.

Glabella 81 to 84 % cranidial length and a maximum width across L1 of 42 to 45 % cranidial width across centre of the palpebral lobes; lateral sides clearly concave in dorsal view, with maximum constriction at or slightly anterior to S2; expansion at or slightly posterior to frontal lobe attains almost the width across L1, expansion results from a fusion with projections of the eye ridges. Front with low to moderate curvature in dorsal view. Three to four pairs of lateral glabellar furrows, all of them faintly impressed to obsolescent on the exterior of the test, all of them quite short and clearly separated at midline; S1 and S2 moderately directed rearward; S3 slightly longer and slightly more oblique rearwards. Occipital furrow very shallow, forming a shallow and sagittally relatively broad depression medially, lateral sections short and obsolescent, swinging backwards, terminate distant from axial furrows. Occipital ring sagittally of about 10 % cephalic length, forming a transversely subelliptical to sublenticular low ridge with a moderately curved posterior margin.

Axial furrows obsolescent, only developed as minor changes in convexity on the shell exterior. Fixigenae low, forming weakly convex subsemicircular areas; extend across centre of palpebral lobes to slightly more than half maximum width of glabella; exsagittally of slightly more than 45 % cephalic length next to axial furrows.

Palpebral lobes small, with fairly low curvature at ocular suture, with moderate transverse convexity, but distal portions more raised and thus slightly upturned in respect to the overall convexity of the cephalon; breadth only about 6 to 8 % of maximum glabellar width, exsagittal length in large individuals about 22 to 24 % cranidial length; situated at a fairly posterior position: anterior end opposite S3 or slightly posterior to its level, posterior tips opposite S1. Palpebral furrows a very shallow, weakly defined, almost straight and somewhat obliquely directed depression. Eye ridges very low, obsolescent ridges, only recognizable as minor changes in convexity on the shell exterior, projecting from near anterior ends of palpebral lobes obliquely forward to merge with the anterolateral corners of the glabella.

Anterior branches of the facial suture weakly diverging from anterior ends of visual surface, faintly curved. Posterior branches moderately long, weakly diverging from posterior ends of palpebral lobes, almost straight in dorsal view for most of the distance, with only minute curvature shortly anterior to posterior margin, posterolateral tips extending to about the level of the palpebral lobes' maximum distance from midline.

Preglabellar field sagittally c. 11 to 14 % cephalic length, follows more-or-less the sagittal curvature of the glabella, in some species obsolescent median plecnum faintly indicated. Precocular fields likewise slightly concave, sloping ventrally.

Anterior border sagittally of c. 10 % cephalic length, poorly defined by obsolescent border furrows. Posterior border consists of an exsagittally narrow, more-or-less straight, barely elevated adaxial section, and somewhat nodular, elevated fulcral extension from which the abaxial section is developed as a simple deflected flap defined by a slightly forward curvature of the cephalic margin. Posterior border furrow obsolescent.

Librigena with relatively narrow ocular platform and wide lateral border separated by change in convexity only. Lateral border extends into a moderately long and broad genal spine of roughly two-thirds the length of the rest of the librigena. Lateral margin with very low curvature for most of the distance so that the genal spines are directed obliquely rearward. Straight posterior margin swings back into the genal spine in a gentle curvature so that the base of the genal spine is conspicuously broad. Doublure wide (Text-fig. 12.6).

Thorax known only from disarticulated thoracic segments. Axial rings wide, moderately convex in transverse section and with low convexity in sagittal section; anterior margin defined by slightly rearward swinging, well impressed axial furrow in normal segments, but obviously with distinct rearward bend in segments 1 and 2 (Text-fig. 12.11). Posterior margin more-or-less straight. Articulating half-ring moderately wide (sag.) and distinctly convex, acutely tapering laterally, not reaching to the axial furrows. Axial furrows developed as distinct changes in convexity only, curved in dorsal view.

Pleurae short, transverse width slightly greater that half width of axial ring, in anterior tergites with anterior margin straight and normal to axis up to a weakly developed fulcral process about half way between dorsal furrows and pleural tips, curves rearward abaxial to the fulcral process. Anterolateral portion developed as a strongly deflected facet abaxial to fulcral socket. Posterior margin of pleurae with straight adaxial section normal to axis up to a slight elevation about half distance between dorsal furrows and pleural tips that forms an indistinct socket corresponding to the fulcral process of the adjacent tergite; posterior margin of pleura faintly back-
ward directed abaxially of the socket. Pleural tip developed as triangular truncated ends, without spines. Pleural ribs low to moderately convex, divided by oblique pleural furrow developed as a wide and poorly defined groove, which commences somewhat distant from axial furrows and fades distant from pleural tips; anterior rib extends into the pleural tip to create slightly elevated part; posterior rib somewhat triangular, does not reach pleural tip.

Surface of cephalon smooth except for few low terrace ridges close to the anterior and lateral margin and the genal spine. Thoracic segments smooth except for a few small, wrinkled terrace ridges close to the pleural tips.

**ONTOGENETIC VARIABILITY AND PHYLOGENETIC IMPLICATIONS:** Juvenile specimens of *Kingaspis campbelli* differ in several respects from adult specimens. Most of these differences focus on the convexity and the proportions of the cephalon. Particularly noteworthy is that the individual parts of the cranidium show their own convexity in juvenile specimens so that the glabella is elevated above moderately convex fixigenae and separated from the preglabellar front. The palpebral lobes are wider and more clearly upturned, and the eye ridges form recognizable low ridges. The posterior border furrow is moderately well developed as a groove increasing in exsagittal width toward the facial suture; the posterior border is somewhat upturned. No distinct differences in the morphological expression compared to adult individuals are evident for the glabellar furrows, the occipital ring, and the anterior border. However, the glabellar furrows, particularly S1 and S2, are visible for a longer distance and thus more strongly recurved, and the preglabellar field carries weakly developed caeca on internal moulds.

The differences in the proportions have an effect on the glabella, which is relatively longer (up to nearly 85% cephalic length in juvenile specimens) and has its widest portion across the anterolateral corners. The glabellar front is more strongly curved, and the occipital ring is sagittally broader and reaches 17% cephalic width in the material and shows a tendency to a subterminal node (Text-fig. 12.14, 12.18). The palpebral lobes are slightly longer (exsag. up to 30% cephalic length) and clearly wider as well as more elevated and more strongly curved at the rim of the visual surface.

These differences between adult and juvenile morphology of *Kingaspis campbelli* reflect in total a progressive smoothing of the surface with the tendency to attain the morphology of a spherical segment. The ef-facement recapitulates the supposed phylogenetic development within the clade, particularly the development from *Kingaspidoideae* to *Kingaspis* which represents one evolutionary pathway among the Ellip-socephalinae. Apart from minor characters, juvenile cranidia of *Kingaspis* resemble those of adult *Kingaspidoideae* Geyer, 1990b so that the distinct modifications during later stages determine the generic identity. This development is a paradigm for Ernst Haeckel’s basic concept of his “law of recapitulation” and certainly one of the few examples of this rule known from early and middle Cambrian trilobites. It provides the direction of the phylogenetic changes and thus indicates indeed that *Kingaspis* is derived from *Kingaspidoideae* and thus phylogenetically younger. In terms of heterochrony, however, the more or less equal size of *Kingaspidoideae* and *Kingaspis* combined with an increasing development of shape places its development under the category acceleration, despite the fact that the newly acquired reduced relief would have been regarded as an indication for neoteny without knowing the direction of the evolutionary development. In addition, this development is a stratigraphic argument that provides a further prove for the age assignment of the Wadi Zerqa Ma’in faunule.

**DISCUSSION:** As mentioned earlier (Geyer and Landing 2004), specimens identified from Spain as *Kingaspis campbelli* by Lihán et al. (2003) are considerably distorted and usually do not show the precise morphology of the external surface. As detailed by Geyer (1990b), kingaspidoidean genera such as *Kingaspis* Kobayashi, 1935; *Kingaspidoideae* Hupé, 1953a; and *Ornamentaspis* Geyer, 1990b, can be identified with certainty and used in precise correlations only if the detailed morphology of the external and internal surfaces is known. For this reason, the Spanish material is not precisely determinable. However, the internal moulds of the specimens identified as *Kingaspis* (K.) *campbelli* from the Iberian Chains represent a species different from *K. campbelli*. This species has pronounced triangular projections of the anterolateral corners of the glabella created by the protruding eye ridges. As noted by Geyer and Landing (2004), one principal character in adult specimens of *K. campbelli* is that the anterior facial suture meets the margin posterior to the anterolateral corners of the glabella — a character that is distinctly different in the Spanish material. Gozalo et al. (2007) noted that this character appears to be variable in the specimens figured by Rushton and Powell (1998, figs 21–26). However, these specimens are not all oriented with the palpebral lobes in a horizontal line. In addition, the specimens from the Iberian Chains have a more elevated glabella and show palpebral lobes in a more anterior position.
Uhaymiria gen. nov.

TYPE SPECIES: Uhaymiria glabra gen. nov., sp. nov., by monotypy.

ETYMOLOGY: Named after the Wadi Uhaymir, Jordan, in which the type material was collected.

DIAGNOSIS: Genus of the Subfamily Ellipsocephalinae with glabella c. 85% cephalic length, slightly tapering forward, frontal lobe moderately wide, subtruncate; three pairs of weakly impressed, roughly parallel glabellar furrows; occipital ring broad (sag.), more than one-fourth glabellar length, with subterminal thorn; fixigenae weakly convex, simple; palpebral lobes reaching posterior border furrow, separated from eye ridges by weak depression only; eye ridges oblique, meet dorsal furrows at level of S3 and tend to bifurcate to produce obliquely forward-directed low ridges which separate the preglabellar field and preocular areas; preglabellar field and preocular areas low, flat to faintly concave; anterior border wide, slightly convex, poorly delimited from preglabellar field.

DISCUSSION: Uhaymiria gen. nov. is a genus typical of the Ellipsocephalinae with fairly effaced morphology. The basic pattern of three pairs of lateral glabellar furrows on the slightly tapering glabella is difficult to recognize. The long palpebral furrows which are slightly arched against the eye ridges, and fixigenae without particular characters also belong to this basic bauplan resulting in the usual mosaic pattern of ellipsocephaloid characters. However, Uhaymiria does not belong to the notoriously “smooth” ellipsocephaloid group with Kingaspis Kobayashi, 1935, Ellipsocephalus Zenker, 1833 or Ornamentaspis Geyer, 1990. This is unequivocally displayed by its subtruncate glabellar front and the broad (sag.) occipital ring. Other diagnostic characters include the strange bifurcation of the eye ridges with obliquely forward-directed anterior branches which extend to the anterior border as faint lines and separate the preglabellar field from the preocular areas.

Uhaymiria glabra gen. nov., sp. nov. (Text-fig. 13)

ETYMOLOGY: From the Latin glabrum, smooth; a reference to the effaced morphology and the low overall convexity of the cranidium.

HOLOTYPE: Incomplete cranidium, FG–602–067c.


DIAGNOSIS: Diagnosis of genus (because of monotypy).

Text-fig. 13. 1-4 – Uhaymiria glabra gen. nov., sp. nov. 1; specimens from the Wadi Uhaymir section. 1, 4 – FG–602–067c, partial cranidium, dorsal view, × 3; 2 – detail of anterior border showing terrace ridges and pits, × 12; 2, 3 – FG–602–008a, partial cranidium, left lateral and dorsal views, × 1.5
DESCRIPTION: Cranidia 21.0 and 19.9 mm long, with width/length ratio of about 0.85. Anterior margin gently curved. Palpebral lobes extend laterally beyond anterior sections of the facial suture. Glabella moderately convex, slightly less than 80 % cranidial length and approximately half cranidial width across occipital ring; slightly tapering forward, with more-or-less straight sides; frontal lobe wide, slightly more than 40 % cranidial width, with very gently curved anterior margin. Three pairs of obsolescent lateral glabellar furrows, indicated only as faint semi-lateral depressions. Occipital furrow an indistinct to obsolescent transverse groove. Occipital ring sag. broad, up to 20 % cephalic length and more than quarter glabellar length, with a moderately curved posterior margin and a short subterminal thorn.

Axial furrows shallow next to proximal areas of fixigenae, poorly defined between eye ridges and glabella. Fixigenae with low convexity, their width across centre of palpebral lobes equals c. 45 % width of occipital ring; exsag. length next to axial furrows about 45 % cephalic length.

Palpebral lobes moderately long (roughly one-third cephalic length), gently curved at ocular suture, transverse width about equal throughout or with slightly narrower (tr.) anterior portion and with moderate transverse convexity; anterior ends at supposed level of L3, posterior tips at about level of anteriormost part of occipital ring, at short distance from posterior border. Palpebral furrows shallow and narrow, weakly defined. Eye ridges project from palpebral lobes, delimited from palpebral lobes by a narrower and less elevated portion, weakly defined, form long elliptical swellings oblique to axis; anterior ends of eye ridges well separated from glabella but with faint trifurcation into threads progressively towards the posterior part of the frontal lobe, towards the anterior part of the frontal lobe, and obliquely forward in the direction of the anterior border (Text-fig. 13.1). This low anterior branch separates the preglabellar field from the preocular areas.

Anterior branches of the cephalic facial suture subparallel from anterior ends of the visual surface, gently curved adaxially when reaching anterior border. Posterior branches not well preserved, moderately long, probably clearly diverging in dorsal view.

Preglabellar field a sagittally broad, faint depression grading into the anterior border. Preocular fields exsag. fairly broad, ventrally sloping areas.

Anterior border low, barely convex, poorly defined, characterized mainly by the presence of terrace ridges, sagittal length c. 5–7 % cephalic length. Anterior border furrow obsolescent. Posterior border convex, slightly backwardly directed and wider abaxially. Posterior border furrow a shallow, poorly defined and fairly narrow groove.

Librigena, thorax, and pygidium unknown.

Surface of cranidium smooth. Anterior border with distinct terrace ridges. Between the terrace ridges are fine punctae (Text-fig. 13.3) which are interpreted to represent the positions of sensory organs.

DISCUSSION: See discussion under genus.

Subfamily Protoleninae Richter and Richter, 1948

*Cambrunicornia* Geyer, 1990b

TYPE SPECIES: *Cambrunicornia vanlooyi* Geyer, 1990; by original designation.

*Cambrunicornia? jafnaensis* sp. nov.

(Text-fig. 14)
DERIVATION NOMINIS: Named after the type locality at Wadi Umm Jafna.

HOLOTYPE: Incomplete cranidium, FG–602–078, from Wadi Umm Jafna section, Jordan.

DIAGNOSIS: Species tentatively assigned to Cambrunicornia with a more-or-less evenly tapering glabella of c. 80 % cephalic length, with three pairs of protolenid-type glabellar furrows; frontal lobe narrow, c. quarter maximum cranidial width, subtriangular, without swelling or spine; fixigenae with small bacculae on posteroproximal portions and weakly raised distal portions separated by shallow longitudinal depression.

DESCRIPTION: Cranidium with width/length ratio of about 0.8. Anterior margin apparently gently curved. Palpebral lobes clearly extending laterally beyond tips of the anterior sections of the facial suture. Glabella slightly more than 80 % cranidial length and 43 % cranidial width across occipital ring; distinctly tapering forward, with approximately straight sides; frontal lobe narrow, almost precisely quarter maximum cranidial width, with subarcuate anterior margin; glabella highly convex. Three pairs of lateral glabellar furrows present, all of them distinctly impressed; S1 and S2 directed moderately rearward and fairly long, indistinctly separated across midline; S3 a short depression close to the axial furrows. Occipital furrow a distinctly impressed, wide and indistinctly delimited furrow with shallow median section. Occipital ring sagittally of c. 13 % cephalic length, with a moderately curved posterior margin and a probably short subterminal spine (only imperfectly preserved on internal mould).

Axial furrows varying from shallow next to proximal areas of fixigenae to moderately deep between eye ridges and glabella. Fixigenae with small bacculae on postero proximal portions and weakly raised distal portions paralleling palpebral lobes, separated by shallow and weakly defined longitudinal depression; width across centre of palpebral lobes half maximum width of occipital ring; exsag. length next to axial furrows slightly less than half cephalic length.

Palpebral lobes moderately long (c. 40 % cephalic length), gently curved at ocular suture, transverse width about equal throughout, with distinct transverse convexity; anterior ends at level of S3, posterior tips at about level of anterior limits of occipital furrow, distinctly distant from posterior border. Palpebral furrows shallow and narrow, weakly defined. Eye ridges project from palpebral lobes, delimited from palpebral lobes by a narrower and less elevated portion, fade towards axial furrows; anterior section of eye ridges appears to swing forward next to glabella, but progression not clearly recognizable.

Anterior branches of facial suture not well preserved, probably subparallel. Posterior branches not preserved, moderately long, probably clearly diverging in dorsal view.

Preglabellar field a sagittally narrow area, developed as a low medial groove grading into the anterior border. Preocular fields narrow and slightly convex, ventrally sloping areas.

Anterior border poorly preserved, only defined by the presence of relatively coarse terrace ridges, sagittally of c. 5 % cephalic length. Anterior border furrow obsolescent. Posterior border convex, more-or-less straight and with little variation in breadth over the preserved section, sloping ventrally. Posterior border furrow a moderately deep groove.

Librigena, thorax, and pygidium unknown.

Surface of the cranidium densely covered with moderately coarse granules except for the smooth lateral portions of the glabellar furrows. The size of the granules varies slightly, being coarsest on the elevated parts of the glabella and slightly more than half this size on the palpebral lobes and the adjacent palpebral furrows and distal parts of the fixigenae. Anterior border with terrace ridges.

DISCUSSION: Species similar to Cambrunicornia? jafnaensis sp. nov. are placed under Cambrunicornia Geyer, 1990, a genus which appears to include species with a fairly wide morphologic range. Geyer (1990b) introduced the genus for two species with a node or small spine on the frontal lobe in adult specimens, which creates the reference for the name (“Cambrian unicorn”). These species share the more-or-less evenly tapering glabella with protolenid-type glabellar furrows, a sagittally narrow, sunken preglabellar field prograding into a low and fairly indistinct anterior border. In addition to the two formally introduced species of Cambrunicornia from Morocco, C. agdziensis Geyer, 1990 and the type species C. vanlooyi Geyer, 1990, the genus is known from incomplete sclerites of a new species from a fauna of earliest middle Cambrian age of the Tröbitz Formation of Germany (Elicki and Geyer 2010). The new species from Jordan agrees in morphology with these species, but lack the node or spine on the glabellar frontal lobe and has a Protolenus-type of fixigenae which differs from the Ornamentaspis-type known from the Moroccan species of Cambrunicornia.

Family Bathynotidae Hupé, 1953b

DISCUSSION: Onaraspis and the Onaraspis clade

The key for a correct understanding, however, may be the morphologically unique genus *Bathynotus* Hall, 1860. *Bathynotus* is a trilobite genus known from Laurentia, South China, Australia and Siberia, and therefore suitable to enhance the resolution of intercontinental correlation in the traditional Lower–Middle Cambrian boundary interval. Such stratigraphical considerations have been described in detail by Geyer and Peel (2011).

Webster (2009) presented comprehensive morphometric data from representatives of all nine previously described species and, based on multivariate and bivariate analyses, recognised a total of only four species, *B. holopygus* (Hall, 1859) (from eastern Laurentia and South China), *B. granulatus* Lermontova, 1940 (including its junior synonym *B. elongatus* Zhao *et al.*, 1987, known from the Siberian Altay-Sayan foldbelt, South China, and western Laurentia), *B. kueichouensis* Lu in Wang, 1964 (from South China and Australia) and *B. namanensis* Lermontova, 1940 (from the Siberian Platform).

The family Bathynotidae was treated earlier as a distinct suborder, Bathynotina Lochman-Balk, 1959, of the Redlichiida (Lochman-Balk 1959), whereas during the last two decades the family was usually treated as of uncertain higher systematic position. *Bathynotus* and thus the Bathynotidae share many characters with taxa of the Redlichiida, namely a tapering glabella with a comparatively simple set of lateral glabellar furrows; long and crescentic palpebral lobes, an opisthoparian facial sutures; librigenae with long genal spines; and a fairly wide thoracic axis with axial nodes. However, most of these characters are plesiomorphic and simply portray a “primitive” character set.

A particular character of the dorsal carapace shared with few Redlichiine trilobite taxa, however, is the thorax consisting of 13 thoracic segments with the 11th segment being macrospinose. A comparable thoracic morphology is only known in the Metadoxididae and the Saukiandidae. The ventral morphology of the Bathynotidae, as shown by *Bathynotus holopygus*, is also characteristic. The hypostome of this species is conterminant and without a recognizable rostral plate so that oblique anterior sutures separate the hypostome from the anterior doublure, giving the hypostome a roughly pentagonal shape. Whittington (1988) suggested that the triangular area in front of the median body in the hypostome of *Bathynotus* may represent a remnant of the rostral plate, fused with the hypostome. If this interpretation is correct, the sutures with the anterior doublure would be connective sutures separating the doublure from the unrecognizable rostral plate. In nearly all other trilobites with small rostral remnants, the typical pattern for such sutures is a divergence towards the anterior rather than a convergence toward an apex, so this feature clearly distinguishes *Bathynotus* from all well known Redlichiida. This considered, Whittington (1997) emphasized the unique character of this sutural contact. It should be noted that a similar discussion with some additional comparisons is placed on Sam Gon’s trilobite website under http://www.trilobites.info/bathynotidae.htm.

Although the hypostomal attachment is unique among the trilobites, the newly discovered material of *Myopsolenites palmeri* (described below; Text-fig. 14.8–10) exhibits a conterminant hypostome with a completely fused rostral plate. The connective sutures with the ventral doublure are generally straight in exsagittal direction, but have a slight median convergence.

The pygidial morphology is exemplified by such material as of *Bathynotus* sp. indet. from the Pioche Formation of Lincoln County, Nevada, USA (Webster 2009, fig. 4), which shows a remarkable conformance with pygidia of *Myopsolenites* from Jordan. In addition, specimens of immature individuals from the Wadi Uhaymir section described below under *Myopsolenites* hyperion sp. nov. exhibit a morphology which very much resembles that of imperfectly preserved immature specimens of *Bathynotus* spp. from South China, again indicating a close phylogenetic relationship of *Myopsolenites* and *Bathynotus*.

Considering Whittington’s (1988) arguments, and the strikingly similar dorsal morphology, *Myopsolenites* and the *Onaraspis*-clade in general is best placed in the Family Bathynotidae. As pointed out by Żylińska and Masiak (2007), other possible relatives which are as yet insufficiently known include Conomicmacca *plana* (Matthew, 1895) from the *Protolemus elegans* Zone of southern New Brunswick, Canada (Westrop and Landingham 2000), *Bathynotellus yermolaevi* Lermontova, 1940 from Novaya Zemlya, Russia (Lermontova 1940), and *Rectifrontinella oliae* Konstantinenko, 2001 from the Carpathian Foreddeep in the Ukraine (Drygant and Konstantinenko 2001).

It should also be noted that *Bathynotus* has been placed under the Chengkouaspidae by Jell and Adrain (2003), which was regarded by these authors as a redlichiod family. Nevertheless, most East Asian taxa exhibit characters which appear to mark them as a primordial lineage of corynexochoid trilobites (which would include the Acanthomicmacca clade).
Despite the extremely limited knowledge on the ventral morphology of most Redlichida, a few older species with a similar conterminant hypostomal pattern, such as in the emuellid Balcoracania Pocock, 1970 and particularly in Saukianda (as shown by material from the Issafen Syncline, Moroccan Anti-Atlas), make it the most parsimonious solution to place the Bathynotidae as a higher taxon under the Redlichida.

Genus Myopsolenites Öpik, 1975

TYPE SPECIES: Myopsolenus palmeri Parnes, 1971; Timna Formation, Timna area, southern Israel.

EMENDED DIAGNOSIS: Bathynotid genus characterized by a cephalon with low overall convexity; glabella tapering forward, front subtruncate, reaching anterior border; three pairs of lateral glabellar furrows, S1 longer than S2 and S3, and with a tendency to be connected mediadly; anterior border with low convexity, slightly sloping rearward from anterior rim, anterior border furrow weakly impressed. Thorax in adult individuals with twelve to fourteen segments, with tenth, eleventh or twelfth segment conspicuously macropleural and thoracic tergites posterior to the macropleural segment reduced in size. Pygidium shield-like, with long, multisegmented axis; pleural fields with shallow to modestly developed furrows; lateral and posterior border obsolete, lateral and posterior margins smooth, usually gently curved. Rostrohypostomal plate with broadly connected rostral and hypostomal units; lateral sutures parallel or slightly diverging forward.

DISCUSSION: As detailed by Geyer and Landing (2004), the genus Myopsolenites Öpik, 1975, was based on a species described by Parnes (1971) as Myopsolenus palmeri from the Timna Formation of southern Israel. Myopsolenus is now regarded as a subgenus of the protolenoid genus Hamatolenus Hupé, 1953a (see Geyer 1990b and Geyer and Landing 2004). Öpik (1975) recognized significant generic (and supra-generic) differences, and proposed the new genus Myopsolenites for the Israeli species. Unfortunately, he named the new genus without a thorough discussion and only in a caption of a table (Öpik 1975, p. 8, table caption) so that this naming has been misinterpreted as a type error for the (morphologically significantly different) Genus Myopsolenus Hupé, 1953 (see Geyer 1990b, p. 175, and Żylińska and Masiak 2007 for further discussion).

The type material of Myopsolenites palmeri (Parnes, 1971) from the Hakhlil Member of the Timna Formation was based on incomplete and poorly preserved specimens, which were restudied by one of the authors (GG). The material is now housed in the Geological Survey of Israel, Jerusalem. This species shows the same general type of thorax and pygidium as the Australian Onaraspis somniuma Öpik, 1968, but differs in the cephalic morphology and the overall size (see below for further discussion). Although a precise correlation of the strata with M. palmeri was unknown, the presence of the genus in northwestern Gondwana significantly expanded the geographic distribution of Onaraspis-clade trilobites.

Over the last decades the known geographic range became even larger. Together with recent finds in Morocco (Geyer and Landing 2004), the clade now appears to have had its maximum diversity in West Gondwana. The previously known specimens of Myopsolenites palmeri from the southern Negev are relatively poorly preserved and do not unequivocally preserve the morphology of the species. What can be seen is that the glabella progressively tapers forward to a subtruncate front. The eye ridges commence at S3 and run more-or-less straight posteriorly and abaxially. The well-elevated palpebral lobes are oblique to the axis and moderately long. The oc- cipital ring carries a stout terminal node. The anterior bor- der is slightly elevated anteriorly, but weakly defined to- wards the preglabellar field. The holotype (Parnes 1971, pl. 3) is a fairly complete but slightly disarticulated spec- imen (due to the moulting process) that is more than 120 mm in length. Due to the cranium being dislocated from the thorax, the exact number of thoracic segments cannot be counted with certainty. However, the specimen’s taphonomically modified condition suggests that the thoracic segments are all preserved, with only the anteri- most tergite disconnected from the rest of the thorax. Therefore, the thorax seems to have twelve segments, with a macropleural tenth followed by two tergites of re- duced size between it and the pygidium. The pygidium is quite well known from additional material. It has rela- tively gently curved lateral margins, shallow interpleural furrows only on the anterior parts of the pleural fields, and successively shallower furrows across the axis. An additional small specimen from the same locality at Har ‘Am- ram (Parnes 1971, pl. 4, figs 35–36) also has twelve thor- acic segments. The best preserved type specimens were collected at Har ‘Amram and have been considered to come from the Mikhrot Member of the Timna Formation. Additional material of the type lot from the overlying Hakhlil Member of the Timna Formation in Timna Na- tional Park suggests that the stratigraphy at Har ‘Amram has been incorrectly interpreted, and that the specimens may indeed come from the Hakhlil Member. Additional material collected by G. Geyer and E. Landing at Timna clearly shows the cranial features, such as a stout ter- minal occipital node; a relatively straight, well impressed
occipital furrow with a shallower part medially; posterior lateral glabellar furrows (S1) that are not clearly deeper than S2 or S3; low, almost straight eye ridges that are connected with and progressing into a parafrontal band; and moderately curved, medium-sized palpebral lobes, the posterior ends of which are clearly distant from the posterior border furrow. In summary, the small number of thoracic segments, the comparatively short (exsag.) palpebral lobes, and the shallow lateral glabellar furrows S1 serve to distinguish Myopsolenites palmeri from the three other species assigned to this genus (Table 1).

A species first described as Perrector? altus Liñán and Gozalo, 1986 from the “Bilbilian Stage” (abandoned designation; now upper Banian through lower Agdzian; see Geyer and Landing 2004) of the Iberian Chains, northern Spain, has been reassigned to Myopsolenites altus by Geyer and Landing (2004) and to Onaraspis altus by Dies (2004). Geyer and Landing (2004, p. 200; 2007, p. 22) argued that the species was difficult to characterize on the basis of the type material from the Valdemiedes Formation at the Rambla de Valdemiedes due to strong distortion, but emphasized the great similarity with the Moroccan species Myopsolenites boutiouiti Geyer and Landing, 2004. Minor differences were seen in the more robust genal spines and the somewhat less curved palpebral lobes of the Anti-Atlas specimens, but the distinguishing characters were accepted from the original description by Liñán and Gozalo (1986, p. 44), who had described M. altus as having a thorax consistently with fourteen segments, and with three segments posterior to the macropleural eleventh segment. In contrast, exoskeletons of adult individuals of M. boutiouiti have no more than thirteen thoracic segments, and with only two posterior to the macropleural eleventh segment. Geyer and Landing (2004, p. 200) emphasized that “additional, better preserved material of M. altus will probably reveal additional characters that distinguish the species”. Myopsolenites altus was redescribed by Dies Álvarez et al. (2007), who noted a “thorax composed of thirteen segments” (p. 424) and the tenth thoracic segment as being “macroleural with a thin, long spine extending to the pygidium” [sic!]. Indeed, some of the figured specimens indicate substantial differences against the original descriptions by Liñán and Gozalo (1986) and that of Dies Álvarez et al. (2007) as pointed out by A. Vincent (written comm. to GG, 2009). Figure 2 in Dies Álvarez et al. (2007) shows several fairly complete exoskeletons, of which the left specimen in text-fig. 2C has a macroleural eleventh segment followed by two segments with small pleurae. The specimens in text-figs 2D, 2G, and fig. 3A in Dies Álvarez et al. (2007) show the same configuration whereas the specimen in text-fig. 2E shows that the macroleural segment is the eleventh. The situation of the right specimen in text-fig. 2C is not unequivocally determinable from the published photograph. The pleural spines of the macroleural segments are apparently somewhat more slender than those seen in the slightly larger specimens described as M. boutiouiti from the Anti-Atlas. However, these differences may be attributed to intraspecific variability, ontogenetic differences, and/or the different degree of deformation. Consequently, we regard M. boutiouiti Geyer and Landing, 2004 as a junior synonym of M. altus (Liñán and Gozalo, 1986). The biosstratigraphic implications for correlation between Iberia and the Moroccan Atlas regions are discussed above under Correlation.

Another species of Myopsolenites was described as “Jakutus kielcensis” by Bednarczyk (1970) from the Brzechów area in the Świętokrzyskie (Holy Cross) Mountains of southern Poland (Geyer and Landing 2004, Żylińska and Masiak 2007). This species differs from M. altus in having distinctly shorter eye ridges and, as a result, narrower fixigenae and palpebral lobes that are somewhat more oblique to the axis. The pygidium of M. kielcensis is narrower, with distinctly less curved lateral margins so that the outline is more-or-less “heart-shaped”, in contrast to the subrounded to subparabolic pygidium of M. altus. In addition, the almost smooth lateral border is much narrower (tr.) in the pygidium of M. kielcensis than that in M. altus.

<table>
<thead>
<tr>
<th>Species</th>
<th>S1</th>
<th>palpebral lobes</th>
<th>thoracic segments</th>
<th>pygidial outline</th>
</tr>
</thead>
<tbody>
<tr>
<td>M. palmeri</td>
<td>shallow, disconnected</td>
<td>moderately long</td>
<td>12–13</td>
<td>subtriangular</td>
</tr>
<tr>
<td>M. hyperion</td>
<td>shallow, disconnected</td>
<td>relatively short</td>
<td>(12–)13</td>
<td>subparabolic</td>
</tr>
<tr>
<td>M. altus</td>
<td>well impressed, transglabellar</td>
<td>moderately long</td>
<td>13</td>
<td>subparabolic</td>
</tr>
<tr>
<td>M. kielcensis</td>
<td>well impressed, transglabellar</td>
<td>relatively short</td>
<td>unknown</td>
<td>subtriangular</td>
</tr>
</tbody>
</table>

Table 1. Comparative morphologies of four Myopsolenites species
MODE OF LIFE: The characters of the dorsal carapace (particularly those of the thoracic segments) suggest that the species of *Myopsolenites* from Jordan described below, were able to considerably incline the body and to curve the thorax concavely in longitudinal section. This leads to the assumption that the ventral appendages were able to dig in the substrate and possibly created *Rusophycus*-type resting trails. Prerequisites of such a mode of life are strongly developed ventral appendages, the suggested presence of which is in perfect accordance with paired bulbous structures on the axial rings close to the axial furrows, which obviously depict the attachment sites of strong dorsoventral muscles to the ventral/internal surface of the shell. *Rusophycus*- and *Cruziana*-type traces were found in the predominantly siliciclastic rocks coeval with the *Myopsolenites palmeri*-bearing beds in the Timna area, southern Israel (Geyer and Landing, unpubl. results). Those *Cruziana* traces with multiple distinct scratch marks of sharp claws can be identified as *Cruziana salomonis* Seilacher, 1990. Its size and occasional imprints of pygidial margins fit perfectly with the characters seen in *Myopsolenites*.

*Myopsolenites hyperion* sp. nov.  
(Text-figs 9.6?, 9.7?, 15)

ETYMOLOGY: Named after Hyperion, a member of the Greek divine gender of the Titans.


DIAGNOSIS: Species of *Myopsolenites* with moderately tapering glabella with shallow, medially disconnected S1 and weakly curved front; palpebral lobes moderately long in adults; adult thorax with twelve to thirteen segments; occipital and thoracic rings with subterminal nodes; pygidium with gently curved lateral and posterior margins, either subparabolic to heart-shaped in outline or subelliptical with median indentation of the posterior margin.

DESCRIPTION: Medium- to large-sized (large complete holaspid up to 18 cm in the studied material) trilobites. Cranidium transversely subrectangular, overall convexity modest, length about 70 to 80 % maximum width across centre of palpebral lobes. Glabella moderately convex, length in adult specimens slightly more than 75 % cephalic length; continuously tapers forward. Frontal lobe sagittally short, with shallow anterior curvature. Three pairs of lateral furrows developed, all shallow and progressively indistinct from S1 to S3 and progressively more forwardly directed, enhanced in visibility on the exterior of the shell by differences in ornamentation. S3 commences at short distance from dorsal furrows, indistinct. Occipital furrow a wide transverse depression but relatively shallow in median sagittal section, lateral sections deeper and slightly backward-directed from axial furrows, median portion almost straight. Occipital ring up to c. 23 % cephalic length, posterior rim gently curved, with terminal occipital node.

Dorsal furrows moderately wide, very shallow, weakly delimited from fixigenae. Fixigenae faintly convex, subtriangular, exsagittally up to 40 % maximum cephalic length and transversely narrow, less than 20 % maximum cranidial width (across centre of palpebral lobes). Posterolateral projections of fixigenae narrow and laterally extended, slope ventrally. Palpebral lobes slightly convex in large individuals, moderately long and with gentle curvature (less than one-third cephalic length), about 7–9 % cranidial width and c. 14–17 % maximum glabellar width, weakly oblique to exsagittal axis, posterior tips clearly distant from posterior border furrow in adults. Palpebral furrows narrower (tr.) than palpebral lobes, shallow in large individuals. Eye ridges clearly oblique to axis, directed c. 25 degrees anteriorly from anterior tips of palpebral lobes in adults, forming moderately elevated ridges in small specimens but fading in height both anteropproximally and posterodistally. Eye ridges curve forward adaxially into narrow bands that form a narrow and indistinct parafourantal band anterior to the glabellar front.

Anterior branches of facial suture moderately long, diverge moderately to meet anterolateral border and curve forward toward anterolateral margin, resulting in a long and distinctly adaxially curved anterior section.
Posterior branches diverge strongly, curve gently from posterior tips of palpebral lobes. Premarginal field narrow and forms more-or-less a sunken, poorly defined area between glabella and anterior border furrow. Preocular fields shallow, slope ventrally from eye ridges.

Anterior border moderately wide, in front of glabella sag. up to one-tenth cephalic length, fairly low, slightly ascending from preglabellar field. Anterior border furrow shallow to obsolete. Posterior border moderately wide and moderately convex, adaxial section normal to axis up to a small socket that corresponds to fulcral process at first thoracic segment, then runs somewhat rearward and is dorsally deflected. Posterior border furrow moderately wide close to dorsal furrows, deepens from midway of the fixigenal width.

Thorax of adults consists of thirteen segments. Thorax subdivided by a macropleural segment which always appears to be the third from the rear. Greatest width in normal segments at about fifth segment, progressively tapering backward. Rhachis conical, widest at second or third tergite, progressively tapering backward. Width of axis at first thoracic segment c. 40 % transverse width of first segment, width at fifth segment c. 36 %, width at tenth segment c. 32 % thoracic width; axial ring at first segment about as wide as occipital ring.

Axial rings moderately convex in transverse section; consist of wide median portion and fairly indistinct narrow (tr.) anterolateral portions. Median portion sagittally weakly convex, with distinct median node which is spherical on the anterior and middle tergites and faintly and progressively extended sagittally on tergites 10 to 12. Anterior margin of axial ring generally straight or slightly sigmoidal, but with distinct rearward bend in segment 1 and in the terminal segment. Posterior margin generally straight, with a slight forward curvature close to the axial furrows. Doublure often visible due to compression, consists of a moderately wide lenticular strip at the posterior rim of the axial ring. Articulating half-ring moderately wide (sag.) and distinctly convex.

Normal pleurae of anterior tergites with anterior margin straight and normal to axis up to a stout fulcral process about halfway between dorsal furrows and pleural tips, curves rearward abaxially to the fulcral process. Anterolateral portion forms strongly deflected facet abaxial to fulcral socket. Posterior margin of pleurae S-shaped, distal portion slightly swinging forward to form shallow concave area close to the pleural spine, with small dorsoventral flexure half way between dorsal furrow and pleural tip that forms an indistinct socket that corresponds to the fulcral process of the adjacent tergite. Except for tergite 1, fulcral sockets and corresponding processes are in a progressively more adaxial position in posterior tergites. Pleural furrows moderately deep, fairly well defined, widest medially, with progressively lenticular shape posteriorly; abaxial tip slightly distant from base of pleural spine. Pleural tip formed by a relatively strong posterior curvature of the anterolateral margin and a concave curvature of the distal part of the posterior margin which produce a moderately long, fairly backward-directed pleural spine that grows progressively rearward. Pleurae of first tergite slightly modified, with short abaxial portions terminated by fulcral processes corresponding to cephalic posterior border. Abaxial part of pleurae at first thoracic tergite with strong posterior curvature of anterior margin to form strongly deflected, large facet, rearward curvature of distal posterior margin and a comparatively small pleural spine (Text-fig. 15.3). Pleural furrow with wider area adaxially.

Macropleural eleventh (or sometimes tenth) segment with axial ring comparable in morphology to that in other thoracic segments, distal portions strongly modified: pleurae greatly expanded, with anterior margin of pleurae similar to normal pleurae and posterior margin directed backward from axial furrows; regular forward swing of posterior margin results in an obliquely falcate outline of the distal portion of the pleura; mid-pleural areas attain nearly double (exsag.) width of normal pleurae. Pleural spines extend outward and rearward to almost the level of the posterior tip of the pygidial axis.

Presence of a macropleural segment affects shape of adjacent posteriormost two tergites. Pleurae in first tergite posterior to macropleural segment reduced to small subtriangular or slightly falcate areas with the anterior margins curved but strongly truncate and posterior margins slightly rearward directed; pleural spine absent. Pleurae of posteriormost tergite lanceolate, with slightly curved anterior and posterior margins; pleural spine absent (Text-fig. 15.5).

Pygidium large, subelliptical in outline with median concave indentation of posterior margin in large individuals; length from articulating furrow to posterior margin about two-thirds maximum width (tr.). Lateral margins smooth, diverge from anterolateral corners to reach maximum width about one-fourth distance from anterior margin, then swinging gently inward. Articulating half-ring narrow (sag.).

Axis of almost 90 % pygidial length, with elongate subtriangular outline, consists of semi-fused axial rings, with c. five axial rings recognizable in juvenile specimens, c. three ill-defined rings recognizable in adult specimens, followed by a large terminal axial piece. Posterior part of axis with elevated, chevron-shaped marginal lobe that creates a weakly defined subtriangular depression.

Pleural areas fused to form a relatively homogeneous, weakly convex subtriangular platform. Three to
four weakly defined interpleural furrows recognizable. Border furrow poorly defined, visible as a slope from pleural fields and a subtle change in convexity towards border. Border low, moderately wide.

Surface of test generally covered with moderately large to large, densely spaced granules without a visible central perforation. Furrows tend to be less densely covered or smooth. The relative size and density of the granules decrease with increasing size of the specimens. Glabellar furrows tend to be smooth in large specimens so that shallow glabellar furrows are often easily recognizable by the differences in the prosopon. More-or-less smooth are the dorsal surfaces of the palpebral lobes. Anterior border covered by long terraces ridges parallel to the anterior margin. Only its posteriormost part shows a slightly transition from terrace ridges to granulation. Remarkably, small but distinct pits are scattered between the terrace ridges (Text-fig. 15.2). These are interpreted herein to have been the location of sensory organs.

The axial rings and thoracic pleurae are covered with granules, whereas the pleural furrows tend to be smooth. The anterior pleural fields carry more and more densely spaced granules than the posterior pleural fields. The pleural spines are covered with obliquely-directed, sparsely dichotomic terrace ridges arranged more-or-less parallel to the anterolateral margin of the pleural spine.

Pygidial axis and pleural fields of the pygidium covered with granules which vary slightly in relative size and density, indicating the pattern of axial furrows. Pygidial border with small and relatively few terrace ridges close to and oblique to the pygidial margin.

**DISCUSSION:** *Myopsolenites hyperion* sp. nov. is a close relative to *Myopsolenites altus* (Liñán and Gozalo, 1986). In particular, the species share a thorax consisting of a macropleural eleventh segment and wide pleural areas that provide the pygidium with a subparabolic outline. The fine granulation with its fine spatial variation also resembles that seen in well preserved specimens of *Myopsolenites altus* from the Moroccan Anti-Atlas (Geyer and Landing 2004, fig. 5). However, *M. hyperion* is clearly distinguished from *M. altus* by its shallow, medially disconnected S1 and a weakly curved front, and by shorter palpebral lobes. The occipital and thoracic rings have subterminal nodes, and the pygidium shows gently curved lateral and posterior margins. Differential growth leads to the presence of a median indentation of the pygidial posterior margin in very large specimens.

*Myopsolenites hyperion* is clearly distinguished from *M. palmeri* (Parnes, 1971) by its shorter glabella; by longer palpebral lobes; by transversely shorter thoracic pleurae with shorter pleural spines; by the subtriangular rather than subparabolic outline of the pygidium with more clearly developed furrows; and by the coarser and more regularly arranged granulation of the test.

**EMENDED DIAGNOSIS:** Species of *Myopsolenites* with moderately tapering glabella with shallow, medi ally disconnected S1 and weakly curved front; palpebral}

**Myopsolenites palmeri** (Parnes, 1971)
(Text-figs 16, 17)

1923. Asaphid; King, p. 511.


1975. *Myopsolenites palmeri*; Öpik: p. 8, caption of fig. 3.


?2007. *Onaraspi* sp.; Gozalo et al., fig. 4E, 4I
Text-fig. 16. *Myopsolenites palmeri* (Parnes, 1971). 1–16 – FG–602–021a, pygidium with compressed rhachis, dorsal view, Wadi Uhaymir; 1, entire sclerite, × 2; 2, detail showing prosopon, × 5; 3–4, 8–9 – FG–602–010a, incomplete pygidium, right lateral, dorsal and oblique posterior views, Wadi Uhaymir, × 2; 5–7, 9–10 – FG–602–043a, pygidium, Wadi Uhaymir; 5, dorsal view, × 2; 6, left lateral view, × 2; 7, detail showing posterolateral pleural region with prosopon of coarse obliquely directed granules, × 9; 9, oblique posterior view, × 2; 10 – FG–602–035, slab with incomplete pygidium (FG–602–035a) in the centre, Wadi Uhaymir, × 2; 11 – FG–602–097, incomplete semigerontic pygidium, preserved in sandstone, partly exfoliated sculptural steinkern, dorsal view, Wadi Issal, × 1.5; 13, 14 – FG–602–045a, incomplete cranidium, dorsal view; 13, detail showing prosopon of coarse granules and short, wrinkled terrace ridges oblique to the pygidial margin grading into granules, Wadi Uhaymir, × 5; 14, dorsal view of entire sclerite and fragment of thoracic segment, × 2; 15, – FG–602–004a, fragment of thoracic axial ring, Wadi Uhaymir. Note differences in the nature of the axial furrow and slightly longitudinal course of axial node, × 5; 16 – FG–602–007b, fragment of thoracic pleura showing transition from granules to terrace ridges, Wadi Uhaymir, × 3.

lobes moderately long in adults; fixigenae moderately wide; adult thorax with thirteen segments; occipital and thoracic rings with posteromedian to subterminal nodes; thoracic pleura transversely short, with small and acute, posteriorly directed spines; pygidium with gently curved lateral and posterior margins, subparabolic to heart-shaped in outline; prosopon of coarse granules.


DESCRIPTION: Medium- to large-sized trilobites. Cranidium transversely subrectangular, overall convexity moderate, length c. 70 % maximum width across centre of palpebral lobes. Glabella moderately convex, length in adult specimens more than 85 % cephalic length; continuously tapering forward. Frontal lobe sagittally short, with shallow anterior curvature which is progressively less pronounced with increasing size. Three pairs of lateral furrows, all shallow and progressively indistinct from S1 to S3 and progressively more forward directed, enhanced in visibility on the exterior of the shell by differences in ornamentation. S1 commences at dorsal furrows, faintly curved and slightly backwardly directed, clearly separated medially on glabella. S2 commences short distance from dorsal furrows, shallow, more-or-less transverse. S3 commences short distance from dorsal furrows, indistinct, transverse to somewhat forwardly directed. Occipital furrow a wide transverse depression but shallow in median sagittal section, lateral sections deeper and faintly backward-directed from axial furrows, median portion almost straight. Occipital ring up to 23 % cephalic length in adults, posterior rim gently curved in large individuals, with subterminal occipital node.

Dorsal furrows moderately wide, very shallow, weakly delimited from fixigenae. Fixigenae faintly convex, subtriangular, exsagittally up to c. 45 % maximum cephalic length and transversely moderately wide, about 20 % maximum cranidial width (across centre of palpebral lobes). Posterolateral projections of fixigenae narrow and laterally extended, sloping ventrally. Palpebral lobes with slight changes in morphology during ontogeny; slightly convex in large individuals, relatively weakly convex and upturned distally in juveniles; moderately long and with gentle curvature in large individuals (c. 45 % cephalic length); about 5–7 % cranidial width and ca. 11–15 % maximum glabellar width, weakly oblique to exsagittal axis, posterior tips located short distance from posterior border furrow in adults. Palpebral furrows narrower (tr.) than palpebral lobes, shallow. Eye ridges clearly oblique to axes, directed c. 25–30 degrees anteriorly from anterior tips of palpebral lobes, curve forward into narrow bands that form a narrow and indistinct parafrontal band anterior to the glabellar front.

Anterior branches of facial suture moderately long, diverge moderately to meet anterolateral border and curve forward toward anterolateral margin. Posterior branches diverge strongly, curve gently from posterior tips of palpebral lobes. Preglabellar field a narrow sunken area between glabella and anterior border furrow. Preocular fields shallow, slope ventrally from eye ridges.

Anterior border moderately wide, in front of glabella sag. about 7 to 9 % cephalic length, fairly low, slightly ascend from preglabellar field. Anterior border furrow shallow and obsolescent. Posterior border narrow, moderately convex, slightly rearward directed and dorsally deflected. Posterior border furrow moderately wide close to dorsal furrows, poorly defined anteriorly.

Thorax of adults consists of thirteen segments, whereas the holotype from Har ‘Amram, southern Negev, has twelve thoracic segments. Thorax subdivided by a macroleural segment which always appears to be the third from the rear. Greatest width in normal segments at about fifth or sixth segment, slightly and progressively tapering backward. Rhachis conical, widest at second or third tergite, progressively tapering rearward. Width of axis at first thoracic segment c. 42–45 % transverse width of first segment, width at fifth segment c. 45–48 %, width at tenth segment c. 40 % thoracic width; axial ring at first segment about as wide as occipital ring.

Axial rings moderately convex in transverse section; consist of wide median portion and fairly indistinct narrow (tr.) anterolateral portions. Median portion sagittally weakly convex, with distinct posteromedian node which is spherical on the anterior and middle tergites and faintly and progressively extended sagittally on tergites 10 to 12 (Text-fig. 16.14). Anterior margin of axial ring generally straight or slightly sigmoidal, but with distinct rearward bend in segment 1. Posterior margin generally straight, with a slight forward curvature close to the axial furrows. Axial furrow with narrow (sag.) lateral and distinctly wider median portion, covered by granules (Text-fig. 16.15). Articulating half-ring relatively wide.
normal ribbon-like sections separated by shallow transverse furrows which are often poorly recognizable on the shell exterior, and nodular lateral sections. Terminal axial piece with chevron-shaped marginal lobe that creates a shallow subtriangular depression.

Pleural areas narrow, subtriangular. Five to six low pleural ribs recognizable. Border furrow poorly defined, visible as a slope from pleural fields and a subtle change in convexity towards border. Border low, moderately wide.

Rostrohypostomal plate with broadly connected rostral and hypostomal units. Rostral plate highly convex in sag. direction, slightly constricted medially; lateral sutures parallel or slightly diverging forward. Hypostome slightly tapering backward. Lateral border convex, moderately broad (tr.), swings distinctly distally and ventrally to meet lateral sutures; delimited from posterior border by inconspicuous nodular swellings and a slight change in direction; posterior border imperfectly preserved in the present specimens, convex, parallels gentle curvature of the posterior margin. Lateral and posterior border furrows moderately deep and moderately broad. Hypostomal middle body consists of large more-or-less elliptical anterior lobe and much smaller crescent-shaped posterior lobe, separated by a shallow, weakly defined furrow; with pair of maculae at anterior section of middle furrow. Anterior part of hypostome developed as a broad (sag.), curved depression between rostral plate and middle body.

Surface of test generally covered with relatively large, densely spaced granules. These granules are relatively high, but do not show any central perforation. Furrows tend to be less densely covered or smooth. The relative size and density decreases with increasing size of the specimens. Glabellar furrows are covered with slightly smaller and slightly less densely arranged granules so that the shallow glabellar furrows are often more easily recognizable by the differences in the prosopon than by its actual topographic alteration. Dorsal surfaces of the palpebral lobes more-or-less smooth. The anterior border is covered by long terraces ridges parallel to the anterior margin. Only its posteriormost part shows a slightly transition from terrace ridges to granulation. The rostral plate carries coarse terrace ridges parallel to the anterior margin. The hypostomal middle body and border furrows are covered by granules, lateral and posterior border by terrace ridges. The sunken connective area between hypostome and rostral plate shows a progressive development from granules via branched ridges in a fingerprint-type manner to the parallel terrace ridges on the rostral plate.

Axial rings and thoracic pleurae covered with granules, whereas the pleural furrows tend to be smooth. The
pleural spines are covered with obliquely directed, sparsely dichotomic terraces ridges.

The pygidial axis and the pleural fields of the pygidium are covered with granules which vary slightly in relative size and density, indicating the pattern of axial furrows. Some specimens show a distinct oblique rearward direction of the granular tips. The pygidial border shows short, wrinkled terrace ridges strongly oblique to the pygidial margin grading into granules of the normal prosopon (Text-fig. 16.13).

DISCUSSION: The new material of *Myopsolenites palmeri* from Jordan shows several morphological details which were unknown in the type material and which help to define the species more precisely. These include the pattern of the glabellar furrows, the morphology of the thoracic segments and the pygidial axis, and the detailed morphology of the prosopon/ornamentation of the test. All these are described above and need no repetition here. Several aspects have been inaccurately described earlier and necessitate a modification of the species’ diagnostic features. They include the number of thoracic segments. The holotype from the Timna area is a carapace with 12 thoracic segments only. Complete individuals from the newly described Jordanian material have 13 segments so that the number varies as in the ontogeny of *Myopsolenites altus*. Another character not seen properly in the material from southern Israel is the morphology of the node on the occipital ring and the axial rings. *Myopsolenites altus* has a large median node extending sagittally to form a short ridge. *Myopsolenites palmeri*, in contrast, has on the anterior and middle part normal median nodes with a circular shape and without a sagittal extension relatively close to but not directly at the posterior margin. In the posterior thoracic segments, the nodes show a weak but progressive tendency toward a longitudinal elongation (Text-fig. 16.14).

The differences from *Myopsolenites hyperion* sp. nov. have been listed above under the discussion of that species. A closer morphologic relationship appears to exist with *Myopsolenites kielcensis* (Bednarczyk, 1970). This species is from the lowermost traditional Middle Cambrian of the Świętokrzyskie (Holy Cross) Mountains in Poland and was originally described as *Jakatrus? kielcensis*. It has been based only on two incomplete cranidia with a surface granulation and a single heart-shaped pygidium (Bednarczyk 1970, pl. 2, figs 1–3). The species has been assigned to *Myopsolenites* by Geyer and Landing (2004) and redescribed by Żylińska and Masiak (2007). Striking differences between *M. palmeri* and *M. kielcensis* exist in the broader, more strongly tapering glabella with a clearly recognizable L1 and a deeper occipital furrow in *M. kielcensis*; distinctly shorter palpebral lobes in the species from the Holy Cross Mountains; and narrower fixigenae. The pygidium of *M. kielcensis* is much narrower and has better developed furrows.

Gozalo et al. (2007, fig. 4E, 4I) and Dies Álvarez et al. (2007, fig. 3C, 3E) figured as *Onaraspis aff. kielcensis* a fragmentary cranidium from the Valdoré section, Cantabrian Mountains, which shares most of the recognizable characters with *Myopsolenites palmeri*. Unfortunately, the slight distortion and incomplete preservation does not permit a confident determination.

Genus and species indeterminate 1
(Text-fig. 18)

MATERIAL: Single incomplete thorax, FG–602–061a, from the Wadi Uhaymir locality, Jordan.

DESCRIPTION: Incomplete thorax of 16 segments, divided into anterothoracic and opisthothoracic section by macropleural segments and differences in width of the rhachis. Axial rings generally moderately convex in transverse section; composed of broad median portion and slightly swollen lateral portions. Lateral portions clearly recognizable in the anterothoracic segments, indicating the attachment sites of strong
dorsoventral muscles. Median portion sagittally weakly convex, with distinct slightly longitudinally elongate posteromedian node. Anterior margin of axial ring generally straight or slightly sigmoidal, but with a notable rearward bend in segments 1 through 3. Posterior margin almost straight, with a forward curvature close to the axial furrows. Articulating half-ring sag, narrow where visible in the present specimen.

Anterothorax in the present specimen composed of 7 segments. Segment 1 through segment 3 with axial ring of c. 40% total transverse width of the segment; pleurae with anterior margin straight and normal to axis up to a stout fulcral process about half-distance between dorsal furrows and pleural tips. Anterolateral portion forms small facet. Posterior margins of pleurae slightly S-shaped, distal portion swinging into a very small concave area close to the pleural spine. Pleural furrows moderately deep, fairly well defined, widest medially; abaxial tip short distance from base of pleural spine. Pleural tip formed by a strong posterior curvature of the anterolateral margin, developing into a short, acute and strongly backward-directed pleural spine. Segment 4 macropleural, with axial ring similar in morphology to that in the thoracic segments anterior to it, distal portions modified to greatly expanded pleurae, with anterior margin nearly straight and normal to axis, posterior margin curved backward from axial furrows and extended into posterior margin of long pleural spine; abaxial exsag. width of pleural areas nearly twice the width at axial furrow; pleural furrow extended into the base of the pleural spine, defined anteriorly by a torus-like ridge which drops markedly to a low anterolateral part of the pleura with a sharply angular tip.

Opisthothorax in the present specimen composed of eight segments, all of them clearly narrower (tr.) than those of the anterothorax, its axis rapidly tapering from thoracic segment 8 rearward to a subparallel posterior section of the rachis. Anterolateral swollen areas of axial rings less conspicuous than in the anterothorax. Segment 8 with acutely subtriangular or slightly falcate pleurae, which are directed slightly rearward; its anterolateral margins slightly curved, posterior margins directed slightly rearward; pleural tips of segment 8 extended into short thorns, delimited from posterior pleural margin by a shallow notch. Segment 9 through segment 16 with axial rings of c. 45% total transverse width of the segment. Pleurae with anterior margin directed slightly backward, curved to extend into short to moderately long posterolaterally directed pleural spines; posterior margins of pleurae directed faintly backward, straight to slightly S-shaped, distal portion swinging into a small concave area close to the pleural spine or forming an indistinct notch. Pleural furrows moderately deep, fairly well defined; abaxial tip extend into the base of the pleural spine and end short distance from posterior margin. Pleural spines directed posterolaterally, but with a progressively rearward direction from segment 9 through 16.

Surface covered by fine granules.

DISCUSSION: The single specimen is characterized by its two macropleural spines. Such a morphology is only known from genera that are otherwise clearly differentiated from the specimen by the majority of morphological criteria, and which are unknown from this part of the Cambrian world.

Other characters include: a relatively straight abaxial margin of the ‘ordinary’ pleurae extended into a short, bristle-type pleural spine; a subterminal elongate pleural node; bulbous anterolateral portions of the axial rings; and a prosopon of small granules. These characters are found, at least partly, in the two species of Myopsolenites described above. The pleural morphology of specimen FG–602–061a matches Myopsolenites palmeri (Parnes, 1971), which, however, does not show a macropleural segment and is easily distinguished by other characters such as a broader axis, etc. Myopsolenites hyperion sp. nov. has one macropleural segment, but in a more posterior position, and has a different type of articulation leading to another shape of the pleural tips.

The size of specimen FG–602–061a indicates that it is much smaller than the adult individuals of all known species of Myopsolenites, but still too large to
show an ontogenetic stage early enough to explain such differences in thoracic morphology. The specimen does not provide any clues for an unusual morphological expression induced by external factors or for a unique individual mutation. As long as no cephalic and pygidia are known, it has to be considered that it represents a bathyurid genus and species which is not formally described.

**Family Palaeolenidae Hupé, 1953**

*Enixus* Ödikmen, 2009

(syn. *Schistocephalus* Chernysheva, 1956)

**NOMENCLATURAL NOTE: Schistocephalus** Chernysheva, 1956 is a junior homonym of *Schistocephalus* Creplin, 1829, a Recent cestode tapeworm (Platyhelminthes, Cestoda) of the Order Pseudophyllidea living on fish, fish-eating birds and rodents. Ödikmen (2009) proposed replacing the generic name for the trilobites by *Enixus*.

**DISCUSSION:** The genus *Schistocephalus* Chernysheva, 1956, long a traditional genus of index fossils in Siberia and the Sayan-Altay Fold Belt, has been revised and is now often treated as a junior synonym of *Palaeolenus* Mansuy, 1912. The type species of *Enixus*, *E. enigmaticus* (Chernysheva, 1956), has a distinct morphology with an almost parallel-sided (or subtly expanding) glabella and four pairs of glabellar furrows, all of them more or less connected over the glabellar midline, and with S3 and S4 much more closely spaced than the others. Other described species of *Enixus* such as *E. juvenis* (Chernysheva, 1956), or *E. amzassiensis* (Fedynina, 1971) (in Chernysheva 1971) agree well with this morphology. Others, such as *E. antiquus* (Chernysheva, 1956), *E. tchernishevae* (Bognibova, 1971) (in Chernysheva 1971), and *E.? ornatus* (Geyer, 1998), differ in having a clearly expanding glabella with the glabellar furrows not joined over the midline.

How far this disparity in morphology requires a taxonomic separation is a matter of debate and personal opinion. As long as lumping of taxa under genera with wide morphological concepts is not used for precise correlation by means of the species furnished under these genera, this modus operandi is equally proficient than splitting the organismic world into small compartments. Nevertheless, small units have, a priori, the benefit to provide a plain view of the morphological characteristics of its members. However, the situation becomes difficult to handle if a broad morphological concept is applied and forces a chain of subsequent synonymizations. Unfortunately, this happened in the case of relatives of *Enixus*.

A first step was the amalgamation of *Palaeolenus* Mansuy, 1912 and *Megapalaeolenus* Zhang, 1966, both genera being typical of the traditional Lower–Middle Cambrian boundary interval in South China. In fact, *Megapalaeolenus* was first thought to be an ancestor of *Palaeolenus*. There is indeed a morphological series leading towards the *Palaeolenus* morphology, which forced authors such as Lin and Peng (2004) and Luo et al. (2007) to regard *Megapalaeolenus* as a junior synonym of *Palaeolenus*. The distinction between *Megapalaeolenus* and *Palaeolenus* was mainly based on the differences between a clavate and an almost subparallel shape of the glabella. This concept was primarily affected by erroneous assignments of species with subparallel glabellas such as *Megapalaeolenus magnus* Zhu, 1980 (in Zhang et al. 1980), *M. expansus* Zhang and Zhu, 1980 (in Zhang et al. 1980) and *M. longispinatus* Zhang and Zhu, 1980 (in Zhang et al. 1980). The species in total show a gradual morphological development which parallels differences between the species furnished under *Enixus/Schistocephalus*: the species of *Enixus* show a similar morphological shift during their evolution, but starts in Siberia with earlier species having a clavate glabella to later species with a glabella with subparallel sides. It is thus logical either to unite all three genera under *Palaeolenus*, or to maintain two separate genera: those with a club-shaped glabella and separated glabellar furrows under *Megapalaeolenus*, and those with fairly parallel-sided glabellas and transglabellar furrows under *Palaeolenus* (with *Enixus/Schistocephalus* as a younger synonym). However, the situation is not readily applicable and complicated insofar as the expected morphocline in Siberia does not match that in South China, but appears to be reversed. Consequently, the stratigraphic order of *Palaeolenus*-type trilobites in South China and Siberia may also be interpreted as a phylogenetic lineage from *Palaeolenus* and related genera via *Megapalaeolenus* to *Schistocephalus*.

In addition, the concept of lumping all of the above-mentioned genera under *Palaeolenus* does not pay much attention to the specific convexity, or relief, of the carapace. Logically, a reduced relief affects the expression or implementation of certain characters so that gradual reduction of the relief may lead to convergent morphological characterization. In this context, it needs to be emphasized that *Gigoutella* Hupé, 1953 is certainly related to *Palaeolenus* and *Enixus*, but synonymizing the genus with *Palaeolenus* as tentatively suggested by Lin and Peng (2004), Gozalo et al. (2007), and Yuan et al.
(2009) means to ignoring clear differences in convexity. The situation is further complicated by species such as Palaeolenus medius (Bednarczyk, 1970) from the Paradoxides insularis Zone of the Holy Cross Mountains, Poland, which has a Palaeolenus-type configuration but represents another, somewhat deviant morphotype (compare Żylińska and Masiak 2007).

An even more delicate problem not discussed earlier is that pygidia are not known from the Siberian material, but are from the Chinese genus Palaeolenus (s. str.). These pygidia are small, trapezoidal, and have a large axis. Rushton and Powell (1998) figured a cranidial fragment from the Wadi Zerqa Ma’in section which morphologically agrees fairly well with the Chinese pygidia.

As an example of the resulting mantras, the synonymization of Enixus/Schistococephalus with Palaeolenus was used by Yuan et al. (2009, p. 218) to correlate the occurrences of Enixus antiquus on the Siberian Platform with those of Palaeolenus/Megapalaeolenus deprati in South China. The authors thus suggested that “the base of the Amgan Stage can roughly correlate with the upper part of the Megapalaeolenus Zone or Arthropoideus chauveaui Zone in South China”, which is incorrect as shown by other, more reliable biostatigraphic data (see Geyer and Peel 2011).

The problem of elaborating the most suitable taxonomic concept for the group is difficult and exceeds the scope of this study. However, lumping all species of Enixus together with Megapalaeolenus and Gigoutella under Palaeolenus certainly does not satisfy the phylogenetic progressions of the clade. Therefore, it appears to be the most sensible solution to retain Enixus for the moment as an independent genus.

Enixus cf. antiquus (Chernysheva, 1956)
(Text-fig. 19)

1998. Palaeolenus antiquus (Chernysheva, 1956); Shinaq and Elicki, p. 255, 256, 259, 263, fig. 7.2.
2007. Palaeolenus antiquus (Chernysheva, 1956); Gozalo et al., p. 366.
2007. Palaeolenus antiquus; Dies Álvarez et al., p. 426.


DESCRIPTION: Cranidium slightly wider than long, length/width ratio c. 0.92 to 0.94 in adults. Axial furrows shallow on the shell exterior, moderately deep on internal moulds, weakly delimited against fixigenae, tapering forward.

Glabella somewhat clavate, maximum width 1.25 times occipital width in large specimens; length reaches 85 % of cephalic length in adults, width across occipital ring slightly less than 40 % cranidial width across centre of palpebral lobes. Glabella moderately convex, frontal lobe rounded, but some large specimens with a clearly shallower section medially (Text-fig. 19.2); four pairs of moderately well impressed lateral glabellar furrows: S1 relatively wide, with clear onset at axial furrows, with slightly rearward bend, weakly bifurcated medially but only connected medially by shallow depression; S2 narrower than S1, slightly curved, in total view normal to axis, deepest slightly distant from axial furrows, disconnected medially; S3 shallow on shell exterior but clearly visible on internal moulds, directed slightly forward, commencing slightly distant from axial furrows, disconnected medially and ending in shallow drop-shaped depression that suggests short bifurcation sections; S4 short but clearly recognizable on both exterior and interior of the shell, directed slightly forward, commencing at axial furrows, at short distance from and with only small angle against S3. Frontal lobe with two faint parafrONTAL lobes, both recognizable with certainty on internal moulds only; anterior one narrowing, complete, commencing short distance anterior to eye ridges and surrounding entire front with about equal breadth; second parafrONTAL lobe posterior to anterior one, thus in a slightly more elevated position, wider, recognizable on well preserved internal moulds only. Occipital furrow more or less straight, with wider median section that suggests slight forward bend medially. Occipital ring moderately long (sagittal length 11–14 % cephalic length), without medial node or with only faint tubercle.

Fixigenae subelliptical in outline; maximum width about 60 % width of occipital ring, maximum length about 36 to 38 % of cephalic length. Palpebral lobe relatively narrow, moderately convex in transverse section, a moderately curved arc of about equal width throughout; posterior end obliquely clipped. Palpebral furrow a narrow crescent-shaped groove, which extends to about double the palpebral lobe width in the centre (opposite L2); extends anteriorly with a faint kink into a shallow and weakly defined furrow that defines the palpebral lobes against fixigenae; shallow groove runs from this sharp bend toward the anterior facial suture and separates palpebral lobe from eye ridge. Eye ridge exsagittally about as wide as palpebral
Text-fig. 19. 1-14 – *Enixus cf. antiquus* (Chernysheva, 1956); all specimens from the Wadi Zerqa Ma’in locality. 1, 4, 9 – FG–602–073, cranidium, partly exfoliated, dorsal, left lateral and anterior views. Note vascular threads on anterior part of frontal lobe, × 3; 2, 5 – FG–602–094, cranidial fragment of individual with unusually broad frontal lobe, dorsal and anterior views, × 6; 3, 6, 8 – FG–602–082a, immature cranidium, dorsal, left lateral and anterior views, × 6; 7, 10 – FG–602–048b, partial cranidium, internal mould, dorsal and left lateral views, × 4; 11 – FG–602–047c, fragmentary cranidium of juvenile specimen, largely exfoliated, dorsal view, × 6; 12 – FG–602–036b, incomplete small cranidium, dorsal view, × 6; 13 – FG–602–047a, incomplete cranidium of juvenile specimen, dorsal view, × 6; 14 – FG–602–036a, incomplete cranidium of immature specimen, dorsal view, × 6
lobe, weakly arched against palpebral lobe; clearly oblique to axis and faintly curved, faintly bilobate with a slightly narrower (exsag.) anterior section (recognizable on well preserved internal moulds only); distinguished from preocular field by greater relief and a marked slope.

Anterior branches of facial suture diverge moderately, ventrally inclined, slightly curved toward sagittal line from half-distance to border furrow and with a distinct curvature towards sagittal line midway through anterior border (which thus suffers a distinct cut-off anteriorly). Anterior wings thus extend slightly less distally than palpebral lobes (despite specimens with seemingly more widely extending wings which are affected). Posterior branches of facial suture diverge strongly, are moderately curved, clearly ventrally deflected, extend more distally than palpebral lobes.

Preglabellar field in front of glabella slightly narrower that anterior border. Preocular fields more than twice as wide as anterior border, with distinct slope from eye ridges, and with low convexity.

Anterior border does not fit into the general convexity of the cephalon; surrounds preglabellar field and preocular fields as a low to moderately raised ribbon with weakly convex surface; tapers slightly toward facial suture due to long clipped portions near margin. Anterior border furrow appears moderately sharp, but is shallow, slightly better defined laterally. Posterior margin of cranidium slightly S-shaped; posterior border slightly broader distally. Posterior border furrow deep near dorsal furrow, widens distally, and runs slightly oblique to axis.

Exterior of test finely and densely punctuate, and with irregularly spaced, small granules on elevated areas of the cranidium such as glabella (except furrows), fixigenae, eye ridges, and posterior border. Anterior border with terrace lines in the anterior half. Preglabellar field and adaxial parts of preocular fields on interior of test with faint, densely spaced radially arranged caeca (visible on well preserved internal moulds only). Larger imprints of vessels occasionally visible, commencing from anterior of eye ridges (Text-fig. 19.2, 19.5).

Pygidium known only from very incomplete sclerites, small, subelliptical in outline, length to width ratio presumably c. 0.6.

Axis highly convex, stout, tapers rearward and reaches posterior border furrows; length c. 85 % of pygidial length. Terminal axial piece rounded posteriorly, about half axial length, with a pair of low protuberances about half-length and a shallow median depression close to the posterior margin. Two obvious axial rings defined by sharp transverse furrows. Width of axis slightly less than two-thirds pygidial width.


ONTOGENETIC VARIABILITY: Immature specimens of Enixus cf. antiquus differ from large adult specimens mainly in the relative proportions. The most conspicuous differences of the juveniles (compared to adults) are:

Glabella highly elevated, sagittally of 83–86 % cephalic length. Glabellar furrows less impressed on the shell exterior. Relative length of frontal lobe larger than in adults. Transverse width across L1 c. 77–81 % width across frontal lobe (c. 74–78 % in large individuals). Anterior border in immature specimens more strongly elevated and sag. and exsag. broader than in adults.

DISCUSSION: The material from Wadi Zerqa Ma’in provides an ontogenetic series of cranidia which clearly defines a distinct form of Enixus Ödikmen, 2009 (= Schistocephalus Chernysheva, 1956). It has been discussed earlier (Geyer 1998, p. 383) that Enixus was known from material of the Siberian Platform and the adjacent regions long before it was found in the Moroccan Atlas region (Geyer 1998) and in the Dead Sea area of Jordan (Rushton and Powell 1998). If the type species of Enixus, E. enigmaticus (Chernysheva, 1956), is taken as the paradigm of the morphology of the genus, both the Moroccan and the Jordanian material is not particularly typical of Enixus, and neither are some of the Siberian species assigned to Enixus. The enigmaticus clade is characterized by lateral labellar furrows S3 and S4 that are closely spaced and well developed close to the midline of the glabella, whereas the antiquus clade has the lateral labellar furrows best developed close to the axial furrows. Among the latter is Enixus antiquus (Chernysheva, 1956), introduced by Chernysheva (1956) with the type species and now index fossil of the lower Amgan Enixus antiquus Zone, which defines the basal Middle Cambrian in the Siberian Anabar-Sinsk facies region. Both the Moroccan and the Jordanian specimens differ markedly from the type species, E. enigmaticus, but are at least similar to E. antiquus. Geyer (1998) has contributed to this fact and assigned the Moroccan species E. ornatus (Geyer, 1998) to the genus with proviso only.

Rushton and Powell (1998) identified the Jordanian material as “Palaeolenus antiquus”, which not only greatly facilitates the correlation from Jordan to Siberia, but also provides a correlation tool into the South China continent. Indeed, the holotype selected by Chernysheva (1956; see Sdzuy 1995, pl. 1, fig. 9) is similar to large
specimens from the Wadi Zerqa Ma’in locality. The specimens from Jordan are so similar to a few of the Siberian specimens described from the River Amga sections (e.g., Chernysheva 1961, pl. 3, fig. 2) that this identification appears to be well justified at first glance. A major problem remains in that the Siberian material of *Enixus antiquus* is quite heterogeneous. The vast majority of the specimens described and figured as *E. antiquus* (compare e.g., Egorova et al. 1976, pl. 23, figs 11–14, pl. 25, figs 1, 2, pl. 28, fig. 1) have an only weakly clavate glabella with a much less expanded frontal portion, its lateral sides moderately diverging forward to form a club-shaped glabella. Those forms have a transversely wider occipital ring and slightly narrower fixigenae than the clavate forms. They are further distinguished from almost all of the specimens from Wadi Zerqa Ma’in by transversely wider palpebral lobes and more strongly projecting anterior wings. It is noteworthy that the sagittal breadth of the preglabellar field varies considerably within the Siberian specimens but does not show a relationship to the glabellar shape. Two incomplete specimens from Wadi Zerqa Ma’in show a broader (or shorter) frontal lobe of the glabella. There is little doubt, however, that these specimens belong to the same species. Nevertheless, they illustrate the difficulties in portraying the morphological characteristics.

Because of the absence of precise stratigraphical and locality information in Chernysheva’s earlier publications on *Enixus* (as *Schistocephalus*; Chernysheva 1956, 1961), it is impossible to assess whether the recognizable morphological differences between specimens are probably due to taxonomic or preservational differences or if the material indeed belongs to a single species with a large and unusual type of variation. In any case, subsequent studies dealing with Siberian specimens of *Enixus antiquus* do not illustrate specimens that accurately match the Wadi Zerqa Ma’in type morphology (e.g., Egorova et al. 1976).

In addition, an apparently clearly distinguishing character between the specimens from Jordan and those from Siberia was already described by Rushton and Powell (1998, p. 143) but not taken into consideration: it is the absence of a distinct occipital node as seen in the Siberian specimens. The cranidia from Wadi Zerqa Ma’in have at best a tiny swelling on the occipital ring.

Gozalo et al. (2007, fig. 4F–4H) figured two incomplete cranidia from the Valdoré section of the Cantabrian Mountains as *Palaeolenus* sp. These specimens apparently represent a species best placed under *Enixus* and appear to resemble an internal mould with a fairly low convexity found at Wadi Zerqa Ma’in (Text-fig. 19.7, 19.10). However, the wider glabella (across L1) with a broad anterior border carrying terrace ridges only in the anterior half and other characters differentiate these specimens from the Jordanian material.

**Family uncertain**

**Genus and species indeterminate 2**

(Text-fig. 20)

**MATERIAL:** Single fragment of a pygidium, FG–602–005, from the Wadi Uhaymir locality, Jordan.

**DESCRIPTION AND DISCUSSION:** The specimen is a fragment of a comparatively large pygidium having a more or less an incomplete rhachis with a quite unusual morphology. Preserved is the c. 6 mm long rhachis consisting of 8 rings in this specimen. These axial rings consist of a strongly convex, almost regularly transverse median section of c. 80 to 85 % width of the entire axis and narrow lateral portions separated from the former by a change in convexity and somewhat irregularly wrinkled narrow furrows as well as slightly rearward deviation with respect to the median section. The long median sections of the axial rings are delimited from each other by moderately well developed furrows, which fade progressively backwards. The course and composition of these furrows are complicated by irregularly and obliquely directed wrinkles, which also become
Acknowledgements

The material for this study was collected during the course of several field campaigns by O.E. and his Cambrian working group, funded by a research grant from the Deutsche Forschungsgemeinschaft (DFG) to O.E. Collection was also made possible by the generous support from the Natural Resources Authority (NRA) of Jordan. Furthermore, O.E. is grateful for joint field work with Rafie Shinaq (Yarmouk University, Irbid, Jordan), Klaus Bandel (Hamburg University), Thomas Wotte (Münster University) and Thomas Biener (now Ludwigshafel, Germany). Comparative trilobite specimens from the Negev and Jordan were collected by G.G. during field research campaigns with E. Landing, New York State Museum, Albany, NY, USA. Due to various circumstances, this study developed into a long term project supported technically by several institutions.

Particular thanks are due to P. Ahlberg (Lund) and A. Żylińska (Warsaw) for extremely thorough reviews of the manuscript and important suggestions for modifications. In addition, we are indebted to Anthony Vincent, Aberdeen, for important observations and for sharing his unpublished field data from the Moroccan Anti-Atlas.

REFERENCES

Chernysheva, N. E. (Ed.) 1971. Amginskiy yarus Altae-Sayan-skoy oblasti. Trudy Sibirskogo nauchno-issled. institut geologii, geofiziki i mineral’no-gorya, Serya paleon-


112–157. Gosgeoltelkhizdat; Moskau, Leningrad. [In Russian]


Yuan Jin-liang, Zhao Yuan-long, Li Yue and Huang You-zhuang 2002. Trilobite fauna of the Kaili Formation (uppermost Lower Cambrian-lower Middle Cambrian) from southeastern Guizhou, South China, pp. 1–422. Earth, Shanghai Scientific and Technical Publisher; Shanghai. [In Chinese with English summary]


*Manuscript submitted: November 15th 2011*

*Revised version accepted: January 10th 2013*