Unexpectedly curved spines in a Cambrian trilobite: considerations on the spinosity in *Kingaspoidoides spinirecurvatus* sp. nov. from the Anti-Atlas, Morocco, and related Cambrian ellipsocephaloids

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Abstract

The new ellipsocephaloid trilobite species *Kingaspoidoides spinirecurvatus* has a spectacular morphology because of a unique set of two long and anteriorly recurved spines on the occipital ring and the axial ring of thoracic segment 8. Together with the long genal spines this whimsical dorsally directed spine arrangement is thought to act as a non-standard protective device against predators. This is illustrated by the body posture during different stages of enrolment, contrasting with the more sophisticated spinosities seen in later trilobites, which are discussed in brief. *Kingaspoidoides spinirecurvatus* from the lower–middle Cambrian boundary interval of the eastern Anti-Atlas in Morocco has been known for about two decades, with specimens handled as precious objects on the fossil market. Similar, but far less spectacular, spine arrangements on the thoracic axial rings are known from other ellipsocephaloid trilobites from the Anti-Atlas of Morocco and the Franconian Forest region of Germany. This suggests that an experimental phase of spine development took place within the *Kingaspoidoides* clade during the early–middle Cambrian boundary interval.

Keywords Cambrian · Trilobita · Systematics · Biostratigraphy · West Gondwana · Morocco

Introduction

The development of spines in trilobites reflects an evolutionary arms race over most of the Palaeozoic. This is particularly signposted by a palpable bloom of spiny trilobites from the late Silurian to the end of the Devonian. Spines are present among the earliest trilobites at the base of Cambrian Stage 3, but these include genal spines and more or less pronounced pleural spines only (as in genera such as *Eofalotaspis*, *Fallotaspis* or *Archaeaspis*). All of these are best regarded as representing part of a beneficial cover by the dorsal exoskeleton in species that needed to perform a more or less distinct inclination of the body along the length axis to allow pronounced enrolment, in some species possibly combined with improved stability on a soft substrate. Early Cambrian trilobites generally retained such a “primordial” organization and spinosity. However, small to medium-sized spines on the occipital ring, axial rings or the lateral and posterior margins of the pygidium developed progressively, apparently as a response to environmental factors, and all of them were nearly upright to more or less gently rearwardly directed. It is generally accepted that new selective pressures from predators prompted animals to craft innovative protective devices, which in several species included longer and more strongly curved spines.

A simple evolutionary solution of enhanced protection was offered in amplifying structures that already existed in predecessors. This solution was frequently performed, but is often not an ideal answer to environmental stresses.
Nevertheless, here we portray a new Cambrian trilobite species with characters of the genus *Kingaspidoides*, which was fairly common in the Cambrian continents West Gondwana and Baltica. However, this species, *Kingaspidoides spinirecurvatus* sp. nov., is easily distinguished from all other well-known species of its genus, and all other ellipsoccephaloid trilobites known thus far, by having two long spines on the axis that curve anteriorly rather than rearwardly as in all other Cambrian trilobites known to date. Several specimens of *Kingaspidoides* are known that appear to indicate an experimental period in respect to the development of occipital and axial spines in the lower–middle Cambrian boundary beds in West Gondwana, with *K. spinirecurvatus* as an extreme form of a morphological experiment with regard to an apparently “wrong” direction of the curvature in the two extravagant spines.

**Trilobite spines as a bioconstructional system**

Spiny trilobites occur particularly frequently in Devonian strata of the Moroccan Anti-Atlas, with a number of species having spines curved in a way that is difficult to understand in terms of their functional morphology. One example for such spines that serve as an organized device is *Koneprusia dahmani* Chatterton, Fortey, Brett, Gibb and McKellar, 2006, which has rearwardly curved genal and occipital spines as well as long dorsally directed spines on the axial rings, but also extended pleural spines which are distinctly curved towards the axis rather than in a rearward direction. However, this entire set of spines forms a spiny cage, which unmistakably was aimed at repelling attacks of possible aggressors (Fig. 1a–c). A similar strategy can be postulated for the quite different spine apparatus developed in *Dicranurus monstrosus* (Barrande, 1852). This species is equipped with a pair of almost spirally coiled occipital spines, as well as abaxially directed genal spines and pleural spines pointing in different directions, but progressively rearward in the posterior thoracic segments.

A special case of axial spine armour is developed in such species as *Psychopyge praestans* Morzadec, 2001 and *P. hammerorum* Chatterton, Fortey, Brett, Gibb and McKellar, 2006. Well preserved specimens of these species show that not all axial rings bear distinct spines (Chatterton et al. 2006, pl. 24, figs. 2, 3). Axial rings with long rearwardly directed spines are separated by two or even three segments lacking axial spines. In addition, a conspicuously rearwardly curved occipital spine is followed by three non-spinose axial rings, and segment 4 with only a relatively small axial spine that corresponds spatially with the tip of the occipital spine (Fig. 2a, b).

More stunning, however, are trilobites with few, but well developed, spines that apparently cannot be considered as a near-optimal solution for a defensive strategy (as shown for *Koneprusia*, above) and are far from forming an apparatus. Such morphotypes include numerous species of *Cyphaspis*, such as *C. smeeki* Van Viersen and Holland, 2016, *C. heisingi* Van Viersen and Holland, 2016, *C. ihmadii* Van Viersen and Holland, 2016, and *C. eberhardiei* Chatterton, Fortey, Brett, Gibb and McKellar, 2006, all with only two long genal spines and a single long, curved axial spine on segment 6 or 7, respectively (e.g., Fig. 1d–f). With 12–14 thoracic segments in total, these species are reminiscent of the morphology of the Cambrian species *Kingaspidoides spinirecurvatus* described herein.

**Spinosity in *Kingaspidoides spinirecurvatus* and other species of the *Kingaspidoides* clade**

*Kingaspidoides spinirecurvatus* sp. nov. is characterized by a long occipital spine and a long axial spine on segment 7 or 8, both of ca. two-thirds the cephalic length or up to cephalic length, and both being curved anteriorly. The occipital spine is usually slightly less curved and generally describes an arc of roughly one-third of a circle, the axial spine is usually slightly more strongly bent and describes an arc in the order of 40 percent of a circle. On a stretched-out individual, these spines are generally subvertically oriented (Figs. 3a, 4a–c, i, 8, 9f–i). They do not suggest an obvious purpose because the distance between them and their positions on the exoskeleton appear to be unsuitable for an effective protective function. However, *K. spinirecurvatus* is frequently found in various types of enrolment, ranging from specimens with the cephalon slightly inclined against the thorax to specimens with (in lateral view) a concave curvature of the thorax to a near-spherical enrolment (Figs. 3c, 4d–h, j, 9a–e). In these inclined or enrolled postures, it becomes obvious that the long extended genal spines play a role in the functional significance of the spine apparatus. The three exemplary postures depicted in Fig. 3 illustrate strongly differing situations for the spatial relations of these prominent spines.

If the cephalon is inclined at 20° and more against the thorax, the tips of the genal spines reach back to the level of the macrospinose segments 7 or 8 and form a lateral barrier in the anterior part of the thorax (Fig. 3a). The efficacy of this lateral protective structure is slightly enhanced by a notable adaxial curvature of the genal spines’ distal portions.

A concave curvature of the trilobite body is commonly believed to represent a resting stage (e.g., Seilacher 1959; Bergström 1983; Figs. 3b, 8). *Rusophycus*-type traces result from this situation during the lifetime of the animals and provide evidence for short-distance burrowing activities.
In *K. spinirecurvatus*, this posture of the body moves the occipital spine back, with its terminal part being directed more perpendicularly to the length axis. The long axial spine with its pronounced curvature, in contrast, is moved forward so that its tip approaches the rearwardly inclined occipital spine. The genal spines are directed posterodorsally and cannot be deemed helpful for protection in this situation.

In a strongly enrolled position, each thoracic segment is inclined at an angle similar to that of its neighbouring segments, whereas the anterior thoracic segments are inclined at a higher angle to the cephalon. Due to the relatively large cephalon, the animals were not able to perform a complete spherical enrolment, and near-completely enrolled specimens are very rare. One specimen that comes close to the maximum enrolment of the species is known and provides insight into the direction of the spines and the positions of their tips in that posture (Fig. 4d, e, h). In this case, the occipital spine, the genal spines, and the macro-axial spine present different angles with respect to the various axes. However, if the centre of the enrolled body is taken as a reference point, the tips of the spines form a series separated by similar angles of 70–80° from each other. The spaces between the resulting equidistant positions of the spines are smaller than the average size of supposed predators from this interval in the Cambrian so that this posture could act as a simple barrier. This appears to be the best possible mechanical solution of a defensive structure against a large predator, being composed solely of four spines, and being based on parsimonious, economic material input with few adaptive characters.
It needs to be emphasized that another specimen of *Kingaspidoides spinirecurvatus* shows a modified version of post-mortem configuration in which the small pygidium forms a flap underneath the posterior thoracic segments (Fig. 9i). Such a configuration is known from several other ellipsocephaloids (see an example of *Hamatolenus (Myopsolenus) magnus* (Hupé, 1953) in Geyer 1990, pl. 44, fig. 12; similarly known from *Ellipsocephalus hoffii* (Schlotheim, 1823) from the Wuliuan of Bohemia).

In conclusion, the presence of two anteriorly curving long spines on the occipital ring and on a posterior thoracic segment, in combination with long, straight (in lateral aspect) genal spines provide a solution for a simple protective system that works under different situations of inclination of the body, in trilobites with a relatively low number of thoracic segments and a small pygidium. The curvature of the occipital spine has to be forward to enable a considerable concave curvature of the thorax to have the spine directed dorsally. The curvature of the axial spine is necessary to maintain a protective device during strong enrolment.

That the morphology of *Kingaspidoides spinirecurvatus* is a possible and extreme response to supposed predatorial stress can be deduced from additional species/forms of *Kingaspidoides* and related genera in the Jbel Wawrmast Formation of the Moroccan Anti-Atlas. The period of deposition of this formation obviously formed a temporal and spatial playground for different morphological experiments. Most of the common species of *Kingaspidoides* do not have long genal, occipital, or axial spines. Such a typical morphology is developed in *Kingaspidoides angustigena* (Geyer, 1990) and *Cambrosaurura usitata* (Geyer, 1990) from the *Morocconus notabilis* Biozone of the Jbel Ougnate region of Morocco, with only relatively short genal spines, a small occipital node, and minute axial nodes, which grow in size in segments 7–9 or 10. Sometimes these nodes are extended into minute spines that cause the axial rings to break off when the rock is split (Fig. 5e). Rare findings show that *Kingaspidoides angustigena* had a single medium-sized axial spine on thoracic segment 8 (Fig. 5a, b, d), at apparently the same position as the long axial spine in *K. spinirecurvatus*, in addition to the medium-sized or long occipital spine that is more often preserved (Geyer, 1990, pl. 31, figs. 5, 8).

One example of character alteration is shown by extremely diverging and broadened genal spines in *Kingaspidoides* sp. A (Fig. 5c, f) which could be considered as an adaptation to fine-grained and very soft substrate. However, the specimens of this species are found in rocks that do not indicate a pronounced soupy substrate present during deposition.

Although the morphologies of the cranidium in *Kingaspidoides* spp. and closely related genera such as...
Ornamentaspis, Cambrosaurura, and Kingaspis are often difficult to distinguish at the species level, the thoraces of the species often show pronounced differences (see Cederström et al., under review). Similar thoracic characters occur in several combinations in different genera, which indicates a mosaic pattern induced by rapid diversification in the ellipsocephaloid clade as suggested earlier (Geyer 1990, 2017). These differences in thoracic character have not been discussed in detail for the group, but some remarks are presented in Cederström et al. (under review).

Among these forms with different patterns of occipital, genal, and thoracic axial spines is a single specimen of a new species from probably the same carbonate bed as Kingaspidoïdes spinirecurvatus of the Jbel Ougnate region of Morocco with a spine pattern that offers an explanation for the evolutionary development of the spinosity towards K. spinirecurvatus. This specimen (Fig. 6a–d), similar to K. spinirecurvatus in most morphological characters, has a moderately long, rearwardly directed and slightly curved occipital spine and nearly straight axial spines of different lengths on segments 4–6 and probably 7. It appears as if their lengths grow from segment 4 to 6 and then diminish.

An additional axial spine of considerable strength and length is developed on thoracic segment 9, which points strongly rearward in partly enrolled carapaces (Fig. 6c).

A similar configuration is seen in another form of Kingaspidoïdes from coeval strata in the Franconian Forest region, Germany. One specimen of this form shows the presence of a slightly curved, moderately long occipital spine, the absence of spines on thoracic segments 1 and 2, and the presence of strongly developed axial spines on segments 3–8, with growing length from 3 to 5 and slightly decreasing in length in segments 5 through 8 (Fig. 6f). A very similar form exists in the Brèche à Micmacca Member (Morocconus notabilis Biozone) of the Jbel Ougnate region. It has moderately long, slightly curved axial spine on segments 3–6, but lacks spines on segments 1, 2 and in the posterior third of the thorax (segments 7–11) (Fig. 5g, h).

Cambrosaurura robusta Geyer in Cederström et al., under review from the Brèche à Micmacca Member of the Jbel Ougnate area has a thick, moderately long and obliquely rearwardly directed occipital spine, but lacks distinct spines on the thoracic axial rings. Nevertheless, the axial rings of some of the thoracic segments show bulbous protuberances resembling short spines. The size of these nodes differs from segment to segment, reaching the maximum size in segments 7 through 9 (Fig. 6e), resembling the series of short spines seen in Kingaspidoïdes sp. B (Fig. 5g, h) and particularly Kingaspidoïdes sp. D (Fig. 6f).

A new, undescribed species of Kingaspis from the Jbel Wawrmast Formation of the Jbel Ougnate region is known only from sparse material and a single more or less complete dorsal exoskeleton (Fig. 6g). It has a long occipital spine and an exceedingly long axial spine on thoracic segment 6, the tip of which reaches back to well posterior to the rear end of the pygidium. No other spines are developed in the thoracic axial rings, but the genal spines are also of considerable length. This configuration of spines is reminiscent to that known from some species of Cyphaspis. It suggests a similar protective strategy although Kingaspis sp. has a distinctly larger body size.

In summary, these different arrangements of axial spines on the thoracic segments indicate less spectacular arrangements apparently developed as protective devices against predatorial stress. They appear to indicate an experimental phase within the Kingaspidoïdes clade during the early–middle Cambrian boundary interval.

Locality information, stratigraphical occurrence, and taphonomy

The localities/areas from which the commercially traded specimens are originating is generally kept confidential. The specimens are collected from two different areas. The main excavating site lies near the western margin of Jbel Ougnate, eastern Anti-Atlas, in the vicinity of Jbel Ba Had-dou near Tizi n’Tfarkhin, approximately at 31°18.5’ N 5°16’ W (Fig. 7), and most of the specimens on the fossil market with slightly scratched surfaces seem to originate from this area. A second site with quarries exists near Tizi n’Izem, approximately at 31°30.5’ N 4°50.0’ W, near the northern edge of Jbel Ougnate.

The strata in which the specimens occur comprise an interval that belongs to the regional upper part of the Brèche à Micmacca Member which forms the lower part of the Jbel Wawrmast Formation (Geyer 1989; Geyer and Landing 2006; Fig. 7). Due to regional differences in subsidence, the Brèche à Micmacca Member is particularly thickly developed in the Jbel Ougnate area, which in turn has provided favourable conditions for fossilization. This led to the establishment of several quarries in which Cambrian trilobites are recovered for commercial use, such as near Tarhoucht (Geyer 1993; Geyer and Vincent 2015), near Assemame (Devaere et al. 2014), or near Tizi n’Tfarkhin. However, these quarries are established in the lower and middle part of the Brèche à Micmacca Member, which belongs to the Morocconus notabilis Biozone. This part of the succession is rich in large specimens of Acidodaparadoxides species (such as A. briareus and A. levissetii) as well as species of Hamatolenus (such as H. vincenti and H. marocanus) (Geyer 1993; Geyer and Vincent 2015; Geyer et al. 2019), which are frequent objects on the fossil market. The fossils from these strata are characterized by the ochre or red-brown colour of the objects on a yellow sandstone matrix due to the shell substance being largely leached. In contrast, trilobite
The limestone beds of the upper part of the Bréche à Micmacca Member that bear the specimens are typical for the carbonate layers that mark the top of shallowing upward cycles and characterize a maximum flooding interval (see detailed explanations in Heldmaier 1998), and explanation for the lower part of the Bréche à Micmacca Member in Geyer and Vincent (2015), with some of the beds overgrown by microbial mats. These uppermost Bréche à Micmacca Member carbonates are usually developed as medium-grey shell hash beds with packstone to wackestone signature, which yield either complete, or more often fragmented, trilobite sclerites, but also brachiopod shells and several other fossil remains such as hyoliths, helcionelloids, and echinoderm ossicles (Geyer et al. 1995). Others are completely devoid of fossils. The trilobite, brachiopod, and hyolith sclerites in the lower beds are secondarily phosphatized in a style that allowed preservation of fine details on the surfaces, whereas the phosphatization leads to homogeneous layers of at least eight specimens prepared from a single slab, with sizes ranging from ca. 27–36 mm length of an outstretched dorsal carapace. a–c BOM 1152a, complete dorsal exoskeleton, dorsal, oblique lateral and anterior views. d, e, h BOM 1152c, enrolled dorsal exoskeleton, oblique posterodorsal, lateral and oblique lateral views. f BOM 1152d, enrolled dorsal exoskeleton, dorsal view of cephalon. g BOM 1152f, incomplete dorsal exoskeleton, anterior view of cephalon. i BOM 1152d, 1152a, 1152b, and 1152c, dorsal exoskeletons. From Ornamentaspis frequens Biozone, Jbel Wawrmast Formation, upper Brèche à Micmacca Member, Jbel Ougnate region, eastern Anti-Atlas, Morocco, most probably from sample locality near Tizi n’Tfarkhin.

Systematic palaeontology

The material used in this study is reposited in the collections of the TU Bergakademie Freiberg (FG), the Naturmuseum Senckenberg, Frankfurt a. M. (SMF), the collection of Bommel, Bize-le-Minervois (BOM), the collection of Gérard Barbe, Reims (CGB), the collections of the Department of Earth and Atmospheric Sciences, University of Alberta, Edmonton (UA), and the Institut royal des Sciences naturelles de Belgique (IRSNB), respectively, under the listed collection number.

‘Class’ Trilobita Walch, 1771
‘Order’ Redlichiida Richter, 1932
‘Superfamily’ Ellipsoscephalacea Matthew, 1887
‘Family’ Ellipsoscephalidae Matthew, 1887
‘Subfamily’ Ellipsoscephalinae Matthew, 1887

‘Genus’ Kingaspoides Hupé, 1953, emend. Geyer, 1990

Type species. Kingaspis (Kingaspoides) armatus Hupé, 1953 (= Kingaspis (Kingaspis) brevifrons Hupé, 1953).

Discussion. Kingaspoides is among the ellipsoscephaline genera which are difficult to differentiate from morphologically similar genera of the clade. The genus has been discussed in detail by Geyer (1990), with an update by Cederström et al. (under review).

As emphasized by Cederström et al. (under review), Kingaspoides is characterized by a distinct reduction of the glabellar and genal convexity toward a Kingaspis-like aspect of the dorsal surface of the cephalic cuticle, whereas internal moulds show distinct axial furrows, well defined glabellar frontal margins and fixigenae which are domed independently from the glabellar convexity. Despite the numerous specimens of Kingaspoides species known from the Moroccan Atlas ranges, well preserved pygidia were unknown from the Moroccan material. Well preserved pygidia of Kingaspoides are known from Kingaspoides frankenwaldensis (Wurm, 1925) from the Franconian Forest, Germany (Geyer 2017) and particularly from the Scanian K. nordenskoeldi (Linnarsson, 1883). These pygidia reconfirm the volucent type of pygidial morphology.

The similar genus Ornamentaspis has an analogous cranial morphology on internal moulds, but shows a morphology of the exterior of the cuticle with a well demarcated glabella with a subtriangular anterior margin of the frontal lobe and fixigenae with a convexity independent from that of the glabella. In addition, the glabella has anterolateral corners without recognizable extensions into the eye ridges on the exterior. The palpebral lobes of Ornamentaspis are subevenly curved and form an arc subparallel to the elongated rise.

\[\text{Fig. 4} \text{ Kingaspoides spinirecurvatus sp. nov., paratypes, cluster of at least eight specimens prepared from a single slab, with sizes ranging from ca. 27–36 mm length of an outstretched dorsal carapace. a–c BOM 1152a, complete dorsal exoskeleton, dorsal, oblique lateral and anterior views. d, e, h BOM 1152c, enrolled dorsal exoskeleton, oblique posterodorsal, lateral and oblique lateral views. f BOM 1152d, enrolled dorsal exoskeleton, dorsal view of cephalon. g BOM 1152f, incomplete dorsal exoskeleton, anterior view of cephalon. i BOM 1152d, 1152a, 1152b, and 1152c, dorsal exoskeletons. From Ornamentaspis frequens Biozone, Jbel Wawrmast Formation, upper Brèche à Micmacca Member, Jbel Ougnate region, eastern Anti-Atlas, Morocco, most probably from sample locality near Tizi n’Tfarkhin.} \]
that forms the most elevated part of the fixigena, whereas the palpebral lobe in *Kingaspoidoides* forms a torus-shaped bar with an elevation independent from the fixigenal morphology, so that the palpebral lobe in *Kingaspoidoides* may be curved or almost straight. The palpebral furrow in *Kingaspoidoides* is thus not evenly curved as in *Ornamentaspis* and frequently shows a nearly straight adaxial margin. On internal moulds of *Ornamentaspis* it proceeds into well
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recognizable depressions which separate the palpebral lobes from the eye ridges, whereas such a depression is less well developed in *Kingaspidoides*.

*Latikingaspis* Geyer, 1990 (type species *Kingaspis* (*Kingaspis*) *alatus* Hupé, 1953) differs from both *Kingaspidoides* and *Ornamentaspis* primarily in having broad axial furrows and librigenae and a relatively slender glabella. Its palpebral lobes have the characters seen in the species of *Kingaspidoides*, but are slightly shorter. The frontal region of the cranidium in *Latikingaspis* consists of an amalgamated unit of the anterior border and the pre-glabellar field/preocular areas, but is shorter than in *Kingaspidoides*.

Other genera of the Ellipsococephalinae are more easily differentiated from *Kingaspidoides*. *Ellipsococephalus* Zenker, 1833, for example, can be distinguished, amongst other features, by its narrow pre-glabellar field and the correspondingly long glabella, shorter and less convex palpebral lobes, librigenae without a genal spine, and particularly by its transversely narrow, sublenticular pygidium with a relatively indistinctly defined axis. *Kingaspidoides*, in contrast, has librigenae with distinct genal spines and a volucent/subordinate pygidium with a well-defined, longitudinally elongate axis.

The differences between *Kingaspidoides* and other genera of the Ellipsococephalinae (such as *Germaropyge* Šnajdr, 1957, *Ellipsostrenua* Kautsky, 1945, *Kingaspis* Kobayashi, 1935, *Mesetaia* Hupé, 1953, *Kymataspis* Geyer, 1990, *Planolimbus* Geyer, 1990, *Elatius* Orlowski, 1985, or *Cambrosaurura* Geyer in Cederström et al. (under review) are discussed in detail in Cederström et al. (under review).

*Kingaspidoides spinirecurvatus* sp. nov.

Figures 3, 4, 8, 9

**Holotype.** TU Bergakademie Freiberg collection FG 688.

**Type locality.** Near Tizi n’Tfarkhin, western Jbel Ougnate, eastern Anti-Atlas, Morocco.

**Type stratum.** Jbel Waermast Formation, upper part of Brèche à Micmacca Member; *Ornamentaspis frequens* Biozone.

**Paratypes.** Collection Patrick Bommel, BOM 1153, seven more or less complete dorsal exoskeletons. Collection Gérard Barbe, CGB T. 41, complete dorsal exoskeleton. Collection Devoille, DEV C 15.2b.

**Diagnosis.** Species of *Kingaspidoides* with moderately elevated glabella of ca. 80% cephalic length (except occipital spine) and 35–40% cranidial width across centre of palpebral lobes, frontal lobe of glabella with indistinct arcuation of anterior margin; lateral glabellar furrows S1 through S4 indistinct on the exterior of the cuticle; occipital ring extended into a long and dorsally curved spine; palpebral lobes relatively short (ca. one-quarter cephalic length), poorly defined from fixigenae on the exterior of the cuticle; anterior area convex on sagittal line, strongly deflected ventrally; anterior border furrow obsolescent. Thorax of adult individuals composed of ca. 14 segments; axial rings with low median node; segment 8 with long, curved median spine on axial ring, recurved forward to almost meet with occipital spine when thorax performs a concave curvature. Surface of cuticle smooth.

**Description.** Cranidium with a length-to-width ratio (occipital spine and genal spines excluded) of 68–73% in the studied specimens. Glabella (exclusive of occipital spine) of 78–83% cephalic length in the studied specimens, maximum width across L1 of 35–38% maximum cranidial width across mid-length of palpebral lobes; glabellar length slightly affected by the grade of dorsoventral compaction that affects the convexity of the pre-glabellar part of the cranidium. Glabella mostly faintly tapering forward from occipital furrow, with width across frontal lobe ca. 85% width across L1, but distinctly tapering in two of the studied specimens (Fig. 9k); gently convex in transverse section, but without the typical almost bar-like elevation seen in most species of *Kingaspidoides*; faint sagittal crest-line visible on internal moulds of three of the studied specimens (Fig. 9o); frontal lobe with faint anterolateral geyer, anterior margin in dorsal view with almost even curvature, but occasionally with faint median kink; four pairs of faint lateral glabellar furrows, S1–S3 recognizable as shallow depressions on internal moulds, all disconnected medially. S4 short, mostly obsolescent. Occipital furrow moderately broad (sag. and exsag.), almost straight, clearly shallow on the exterior of the cuticle. Occipital ring moderately broad (sag.), exclusive of occipital spine of ca. 15% of cephalic length, extended into a long terminal spine; this spine is directed dorsally with a distinct concave curvature so that the tip points more or less exactly upright.

Axial furrows relatively broad and shallow, poorly defined from fixigenae.

Fixigenae posterior to eye ridge gently convex in transverse section, but genal sector generally sloping slightly ventrally in abaxial direction; near axial furrow of ca. 45% cephalic length (exsag.; occipital spine excluded), transverse width across mid-length of the palpebral lobes of ca. 65% maximum glabellar width and only 22–24% maximum cranidial width.

Palpebral lobe variably developed, but without considerable reduction in length during growth, relatively short for a species of *Kingaspidoides*, in adult individuals exsag. of 23–26% cranidial length (occipital spine excluded), width in the order of 6–8% cranidial width across centres of palpebral

lobes; weakly convex in transverse section on the exterior of the cuticle, gently convex (tr.) on internal moulds, with slightly crescentic shape, distinctly curved along suture toward visual surface, less curved toward palpebral furrow, with anterior end at approximately the level of the anterior part of L3, in a faintly more adaxial position than posterior end; posterior end approximately at level of L2; better defined and narrower on internal moulds; considerably elevated above librigenal platform (Fig. 9j). Palpebral furrow a shallow depression, poorly defined on the exterior of the cuticle, relatively distinct on internal moulds; extends on internal moulds anteroproximally into a shallow and poorly
defined furrow that separates the eye ridge from the fixigena posterior to it. Eye ridge a low, obliquely transverse and weakly curved torus, poorly marked on the exterior of the cuticle, separated from palpebral lobe by a constriction and a weakly defined depression.

Anterior area of the cranidium uniformly convex on the exterior of the cuticle, with anterior border generally not defined or obsolescent both on the exterior and interior of the cuticle. Anterior area of ca. 20% cephalic length (occipital spine excluded), gently convex, moderately well demarcated from glabellar front on internal moulds, but separated only by changes in convexity on the exterior of the cuticle; grades laterally into preocular areas without any recognisable distinction. Anterior border on the sagittal line poorly marked in most individuals, of only ca. 5% cephalic length in dorsal view, of nearly equal breadth (sag., exsag.) throughout, distinctly narrower (sag., exsag.) than pre-glabellar field.

Posterior border faintly defined on the exterior of the cuticle, moderately wide, relatively narrow close to the axial furrow and growing in width abaxially to a moderately broad (exsag.) lobe for most of its extension on internal moulds, slightly to moderately convex, transverse to the length axis in the proximal and median part, its course swinging slightly anteriorly in the distal part, with a small process serving for articulation with first thoracic segment posterior to palpebral furrow. Posterior border furrow a shallow, moderately broad (exsag.) groove.

Anterior branches of the facial suture are difficult to recognize in most specimens, weakly divergent from anterior ends of ocular suture, straight for most of its course, with slight adaxial curvature at approximately the supposed position of the anterior border. Posterior branches are equally difficult to recognize, moderately divergent, comparatively weakly curved.

Librigena moderately wide (tr.), with a moderately long to long, relatively strong, genal spine of roughly equal length of the anterior part of the librigena. Genal spine weakly curved, with a moderately broad base that defines a gentle curvature from the posterior margin to the adaxial rim of the genal spine. Lateral margin gently and evenly curved, at most with a faint bend at the base of the genal spine. Palpebral section of the suture moderately large, corresponding to an eye length of ca. one-quarter cephalic length. Librigenal field fairly narrow in dorsal view, slightly convex, slopes gently downward from the palpebral section toward the weakly defined border furrow. Lateral border slightly convex in transverse profile, subequal to the librigenal field’s transverse width.

Hypostome and rostral plate unknown.

Thorax of 14 segments (possibly 15 segments in two of the studied specimens) in adult individuals. Axial rings of segment 3–7 and 9–14 of more or less uniform shape (sag. and exsag.), length and width reduced progressively posteriorly to less than half maximum width across segment 3, whereas the reduction in sagittal and exsagittal directions are less affected by allometric growth. Transverse width of average axial ring ca. 35% of overall width of the segment, growing slightly in relative width to ca. 45% in the posterior segments. Moderately broad (sag. and exsag.) articulating furrow with distal apodemat pit. Surface of axial ring gently convex in sagittal and exsagittal profile. Each axial ring with an (usually low) axial node, which apparently tends to be better developed and more prominent in the mid-thoracic segments. Greatest overall width (tr.) of thorax at segments 1–3, decreasing in width (tr.) rearward, with slightly but progressively backward directed distal parts of pleurae, particularly posterior to segment 10. Pleural furrows moderately broad to broad (exsag.), forming furrows of narrow lenticular outline slightly oblique to transverse axis, which fade a short distance from pleural tips (e.g., Fig. 9k). Pleural tips blunt, somewhat obliquely subtruncate, but with minute, slightly posterolaterally directed spine or thorn. In anterior view, the pleurae have a distinct geniculation adaxial to the transverse mid-length, which corresponds to the articulation of the adjacent segments by means of fulcral points. As a result, the anterior flanks of the distant pleural portions have an almost flat, anteriorly tilted articulating facet, whereas the posterior margin of the distal pleura turns slightly forward to allow a limited overlap of adjacent segments during inclination of the segments (Fig. 9k, m).
The first/anterior thoracic segment differs in having an axial ring with an anterior margin that swings backward to allow a distinctive dorsal flexure of the cephalon against the thorax. Its pleurae are straight in the proximal sector, but show a well-developed fulcral point to fit into the socket of the cephalon’s posterior border; distal part of pleura reduced in size, with a somewhat backward-directed anterior margin and an obliquely ventrally deflected articulating facet. This configuration is repeated in a less distinct manner in the second thoracic segment.

Segment 8 with a slightly broadened axial ring, which extends into a prominent, large, conspicuously curved spine. This spine is posterodorsally directed from its origin but strongly curved in lateral view so that its tip is directed anterodorsally (Fig. 9m).

Pygidium incompletely known, only preserved in two specimens (Fig. 9i, n), but clearly of volucent morphology, with segments having a similar morphology as the posterior thoracic segments; considerably convex in sagittal and transverse profile. Axis clearly defined, tapering rearward, convex in transverse profile, consisting of at least five axial rings, apparently reaching to posterior margin. Pleural areas similar in morphology to thoracic segments, with well-developed pleural and interpleural furrows, but without lateral border (Fig. 9n).

Surface of cuticle smooth, finely punctate in places.

**Dimensions.** Fully grown cranidia of *Kingaspidoides spinirecurvatus* are between 25 and 40 mm long (holotype: 32.8 mm), with 20–32 mm width between the tips of the genal spines (26.3 mm in the holotype). The width between the ocular sutures at the palpebral lobe is 17.4 mm in the holotype. The length of the thoracopygidial part of the exoskeleton is usually between 25 and 30 mm (22.3 mm in the holotype).

**Comparison.** *Kingaspidoides spinirecurvatus* is easily distinguished from all other species of the genus *Kingaspidoides* by its two long occipital and axial spines. The species is typical for the genus *Kingaspidoides* in having fixigenal-palpebral units sloping ventrally toward the suture with little difference in convexity between the interior and exterior surfaces of the exoskeletal cuticle (other than in the species of *Ornamentaspis*), in a fairly uniformly vaulted and ventrally deflected anterior area, and in having a glabella with subparallel sides. The most similar cranidium can be seen in large individuals of *Kingaspidoides destombesi* (Geyer, 1990), which is distinguished by a simple, moderately long and more or less straight occipital spine and a distinct Bertillon pattern on most of the cranidium.

*Kingaspidoides spinirecurvatus* also has relatively strongly developed genal spines that fit into the spine apparatus discussed above. The cuticular surface is finely punctate (Fig. 8d) and certainly lacks any distinct granules or Bertillon patterns that occur in a number of species from...
Unexpectedly curved spines in a Cambrian trilobite: *Kingaspidoides spinirecurvatus* sp. nov. from Morocco.

**Fig. 8** *Kingaspidoides spinirecurvatus* sp. nov. **a-c, e–h** Holotype, FG 688, complete exoskeleton with slightly concave flexure; **a** dorsal view; **b** oblique lateral view with spine configuration and comparatively well-defined palpebral lobes; **c** anterior view; **e, f** dorsolateral and lateral views illustrating fulcral points and well developed facets on thoracic pleura; **g** oblique posterior view showing posterior part of the thorax and damaged, slightly ventrally inclined pygidium; **h** oblique lateral view of middle and posterior part of the thorax and pygidium with slightly broadened axial ring of thoracic segment 8 and subequal curvature of axial spine. From *Ornamentaspis frequens* Biozone, Jbel Wawrmast Formation, upper part of Brèche à Macoma Member, sample locality near Tizi n’Tfarkhin, Jbel Ougnate region, eastern Anti-Atlas, Morocco. **d** Paratype, Collection Gérard Barbe, CGB T. 41, detail of glabella showing finely punctate surface and lateral glabellar furrows, oblique anterolateral view; from unknown locality, Jbel Ougnate region, eastern Anti-Atlas, Morocco. All scale bars 5 mm.
Unexpectedly curved spines in a Cambrian trilobite: *Kingaspidoides spinirecurvatus* sp. nov. from Morocco

West Gondwana, but examination of perfectly preserved specimens is needed to further discriminate the precise surface design.

**History of discovery and commercial use.** *Kingaspidoides spinirecurvatus* sp. nov. is hitherto known from a number of specimens on the fossil market, sold under various names, mostly as "*Kingaspis* sp." or "*Kingaspidoides* sp.". Three specimens were recognized in 1988 from small fragments of cranidia from the Tizi n’Izem locality of the Jbel Ougnate region, eastern Anti-Atlas (sample horizon TN1-N2 of Geyer 1990) that showed the unique occipital spine with its anterior curvature, but excluded from a formal description in Geyer (1990) because of expected subsequent better preserved and more conclusive material. Another fragment of a specimen was found from the scree unequivocally coming from the upper part of the Bréche à Micmacca Member or the lowest Tarhoucht Member of the Bou Tiouit section near Tarhoucht, thus roughly coeval with the material from the Tizi n’Izem section. Unfortunately, these specimens have been lost during relocation of samples.

The species is easily recognized by the two recurved spines on the occipital ring and thoracic segment 8, which unequivocally distinguish it from all other species of the genus *Kingaspidoides*. This configuration of the spines is also unknown from all species of the closely related genera, such as *Ornamentaspis*, *Kymataspis*, *Ellipsostrepsina*, or *Cambrosaurura*.

Commercial exploitation of the species started in the 1990s, and a number of specimens were offered and sold on the fossil market. However, the species remained relatively rare, and most specimens were damaged, incomplete and/or partly faked. Commercially traded specimens are almost exclusively prepared mechanically from relatively thin, partly nodular limestone beds by means of chisels, needles and vibrottools, which cause considerable scratching of the shell and split-off particles. The conspicuous occipital and axial spines break off during preparation and are glued to the rest of the carapace at the end of the preparation process, sometimes replaced by spines that originate from other specimens, even other trilobite species or other materials. In a number of cases, the spines are mounted in a wrong direction on the rest of the exoskeleton so that the occipital spine or the axial spine appears to curve rearward.

**Preservation.** The specimens known of *Kingaspidoides spinirecurvatus* sp. nov. are all preserved in carbonate horizons of variable lithology, but generally slightly argillaceous or with considerable clay content. The cuticle is preserved as a black phosphatized coat. This preservation is typical for skeletal fossils from the uppermost part of the Bréche à Micmacca Member and the lower part of the Tarhoucht Member of the Jbel Wawrma Formation in a number of sections in the eastern Anti-Atlas, and differs considerably from the normal preservation of skeletal fossils of the Jbel Wawrma Formation, which are generally preserved as orange to brownish coatings resulting from the weathered calcareous substance of the shells and cuticles in the yellowish-green fine-grained sandstones of the lower and middle parts of the Bréche à Micmacca Member.

Typical carbonate beds from which the specimens of *Kingaspidoides spinirecurvatus* are recovered are slightly nodular and contain a plethora of trilobite and brachiopod sclerites, with an unusually high number of articulated specimens. The limited, but occasionally evident fracturing of these sclerites indicates deposition under high energy conditions, but with only a very limited transport distance. Nevertheless, these conditions appear to be limited to only a few localities, with fracturing of fossil shells and sclerites being a general condition in nearby locations of the same beds.

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