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Ben Thuy Natural History Museum Luxembourg

Vivienne Maxwell Smith College

Sara B. Pruss Smith College, spruss@smith.edu

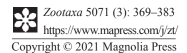
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A new phosphatized ophiuroid from the lower Triassic of Nevada and its position in the evolutionary history of the Ophiuroidea (Echinodermata)

BEN THUY1*, VIVIENNE MAXWELL2 & SARA B. PRUSS2

¹Natural History Museum Luxembourg, Department of Paleontology, 25 rue Münster, 2160 Luxembourg City, Luxembourg.

²Department of Geosciences, Smith College, Northampton, MA 01063, USA.

smith.edu; https://orcid.org/0000-0002-2597-0498

*corresponding author: sthuy@mnhn.lu; https://orcid.org/0000-0001-8231-9565

Abstract

The Lower Triassic fossil record of brittle stars is relatively rich, yet most records published to date are based on poorly preserved or insufficiently known fossils. This hampers exhaustive morphological analyses, comparison with recent relatives or inclusion of Early Triassic ophiuroid taxa in phylogenetic estimates. Here, we describe a new ophiuroid from the Lower Triassic of Nevada, preserved as phosphatized skeletal parts and assigned to the new taxon *Ophiosuperstes praeparvus* gen. et sp. nov Maxwell, V. & Pruss. S.B. This unusual preservation of the fossils allowed for acid-extraction of an entire suite of dissociated skeletal parts, including lateral arm plates, ventral arm plates, vertebrae and various disk plates, thus unlocking sufficient morphological information to explore the phylogenetic position of the new taxon. Bayesian phylogenetic inference suggests a basalmost position of *O. praeparvus* within the Ophintegrida, sister to all other sampled members of that superorder. The existence of coeval but more derived ophiuroids suggests that *O. praeparvus* probably represents a member of a more ancient stem ophintegrid group persisting into the Early Triassic.

Key words: brittle star, end-Permian mass extinction, phylogeny, fossil

The Late to Early Triassic was a pivotal interval in the evolution of marine communities (e.g. Brayard *et al.* 2017). It witnessed the most extensive mass extinction event so far recorded in Earth history (Sepkoski 1981; Erwin 1993). As a consequence, many of the surviving organismal clades are believed to have undergone a drastic bottleneck followed by a post-extinction radiation (e.g. Twitchett & Oji 2005). However, our understanding of the evolutionary change associated with the end-Permian mass extinction and the subsequent recovery directly depends on the completeness of the fossil record available. In recent years, a number of new fossil discoveries have challenged previous paradigms on faunal change around the Permian-Triassic boundary, including an unexpectedly rapid post-Permian recovery of some marine groups (Brayard *et al.* 2017; Doguzhaeva *et al.* 2018; Botting *et al.* 2019; Brayard *et al.* 2019; Charbonnier *et al.* 2019; Romano *et al.* 2019; Saucède *et al.* 2019) and the previously unnoticed survival of Paleozoic holdovers (Thuy *et al.* 2017; Thompson *et al.* 2018; Hagdorn 2018).

Ophiuroids, or brittle stars, are one of the five extant echinoderm classes to have survived the end-Permian mass extinction. With respect to their Late to Early Triassic fossil record, they appear to be the best sampled of the five classes, with more than 14 species known from around the boundary interval (Chen & McNamara 2006). A critical re-evaluation of these fossil occurrences, however, debunked most of them as poor, based on insufficiently preserved specimens and/or superficial descriptions, thus precluding their inclusion in phylogenetic analyses (Thuy *et al.* 2019). In fact, for an ophiuroid fossil to be available for phylogenetic studies, it has to be known in sufficient detail to allow comparison with recent relatives (Thuy & Stöhr 2016). The lack of phylogenetically informative ophiuroid fossils from the Late to Early Triassic is particularly unfortunate because the current consensus on the evolution of the class indicates a mid-Permian origin for the crown-group Ophiuroidea followed by a rapid radiation of the major extant clades (O'Hara *et al.*, 2014, 2017; Thuy & Stöhr 2016, 2018).

Here, we describe new ophiuroid fossils from the Lower Triassic of Nevada (Fig. 1). The extraordinary preservation of the fossils, involving replacement of the original skeletal calcite by calcium phosphate (apatite),

allowed for acid-extraction of an entire suite of dissociated skeletal plates from limestone, unlocking sufficient morphological information to allow inclusion of the new ophiuroid in a phylogenetic analysis.

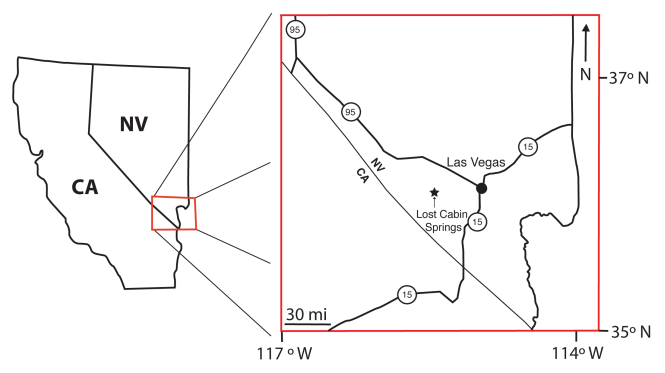


FIGURE 1: Locality map of the Virgin Limestone Member at the Lost Cabin Springs Locality, Southern Nevada, Western United States (36°4′57.18′N, 115°39′12.05′W is near the base of the section). (modified from Maxwell, 2020).

Geological Context

During the Early Triassic, the western United States was situated at a tropical paleolatitude (Reif & Slatt, 1979). The Lower Triassic Moenkopi Formation extends throughout the Colorado Plateau and into southern Nevada and its depositional environment gradually shifts from continental fluvial sedimentation in Arizona to more intertidal and shallow marine environments in Utah and Nevada (Reif & Slatt, 1979). The Moenkopi Formation in southern Nevada was deposited in a shallow water setting on the eastern margin of the Panthalassa sea (Marzolf, 1993) (Fig. 2). The non-marine Timpoweap Member or the Lower Red Member is overlain by the Virgin Limestone Member, which is in turn overlain by the evaporitic Shnabkaib Member (Reif & Slatt, 1979). The Virgin Limestone Member of the Moenkopi Formation is a mixed carbonate-siliciclastic succession (Poborski, 1953). At the Lost Cabin Springs locality, it is constrained to the Spathian Stage (late Early Triassic) (Poborski, 1953) and was deposited in a storm-dominated subtidal paleoenvironment (Schubert & Bottjer, 1995; Pruss *et al.*, 2005). There is an abundance of unusual facies present in the Virgin Limestone Member, including microbialite facies, such as large microbial reef mounds that span the length of the Lost Cabin Springs outcrop area (Pruss & Bottjer, 2004), wrinkle structures (Pruss *et al.*, 2004), and flat-pebble conglomerate facies (Pruss *et al.*, 2005). Of eight fossiliferous samples, ophiuroids were only present in Bed LC–18–34 (Maxwell *et al.*, 2020). Approximately 115 meters from the base, this bed is a thin 3.5 m-thick fossiliferous packstone with isolated crinoidal ossicles overlying a small microbial build-up.

Material and Methods

In 2018 and 2019, samples of fossiliferous packstone were collected and dissolved in 200 – 400 ml of 10% glacial acetic acid solution buffered with ammonium acetate, and only one sample (LC–18–34) produced abundant ophiuroid fragments. The insoluble residues were sieved into the following size fractions: >400 μ m, >250 μ m and >177 μ m, and the ophiuroid elements were common components of the >400 μ m and >250 μ m fractions. The sieved samples were examined under the Nikon SMZ645 stereoscopic microscope, and an Olympus BHS BH-2 light microscope

was used to image these fossils. The best preserved ophiuroid elements (n=71) were coated with gold and palladium by a Hummer V Sputter coater and imaged with the FEI Quanta 450 Scanning Electron Microscope (SEM) at Smith College and with the Jeol Neoscope JMC-5000 SEM at the Natural History Museum Luxembourg. The elemental composition of the fossils was analyzed using EDS (Energy Dispersive Spectroscopy) Team software at Smith College.

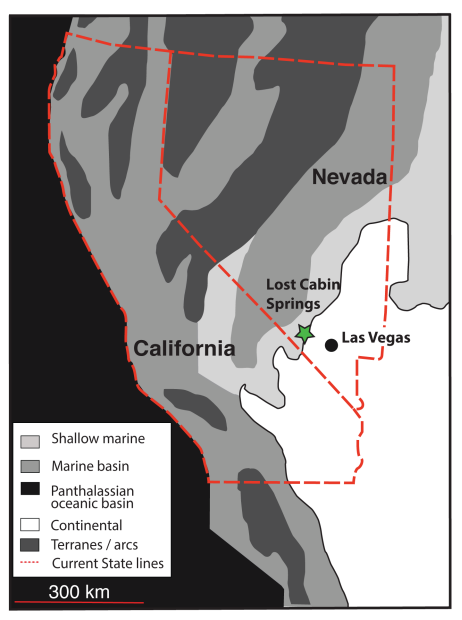


FIGURE 2: Early Triassic paleogeographic reconstruction of the western US. The Lost Cabin Springs Locality is denoted with a green star and the various shades depict the depositional environment during the Early Triassic (modified from Hoffmann *et al.*, 2013).

Specimens here included were deposited in the collections of the Natural History Museum (MnhnL OPH) and the Muschelkalkmuseum Ingelfingen (MHI).

The terminology used herein follows Stöhr *et al.* (2012) and Thuy & Stöhr (2011, 2016). We adopt the classification proposed by O'Hara *et al.* (2017, 2018). The extreme fragility of the skeletal plates, however, precluded detachment and re-mounting of the lateral arm plates. As a result, we were unable to illustrate the external and internal sides of the same lateral arm plates using scanning electron microscopy, as is a common practice in ophiuroid descriptions (Thuy & Numberger-Thuy 2021).

We scored the skeletal plates following Thuy & Stöhr (2016, 2018), referring to the same character definitions and acronyms and using the character matrix elaborated by Thuy & Stöhr (2016) and modified by Thuy & Stöhr

(2018) but excluding *Inexpectacantha acrobatica* Thuy, 2005 because of its ambiguous taxonomic affinities (Thuy & Numberger-Thuy 2021). The character matrix including the scores of the ophiuroid material described in the present paper is shown in Table 1. Bayesian inference analysis was performed using MrBayes (Huelsenbeck & Ronquist 2001) which relies on a modified version of the Juke-Cantor model for morphological data as outlined by Lewis (2001), with variable character states from 2 to 10 (Wright & Hillis 2014). Only variable characters were sampled, and we compensated for character selection bias by letting MrBayes search for parsimony informative characters (Mkpars model) (Wright & Hillis 2014). All character states were assumed to have equal frequency, and prior probabilities were equal for all trees. We assumed that evolutionary rates vary between sites according to a discrete gamma distribution. Average standard deviations of split frequencies stabilized at about 0.008 after 3 million generations (mgen), sampled every 1,000 generations. The first 25% of the trees were discarded as burnin. The consensus trees were examined with the software FigTree v. 1.4.2 by Rambaut (http://tree.bio.ed.ac.uk/software/figtree/). Confidence intervals of 95–99% were regarded as strong support for a node to be true, and at least 90% probability as good support.

Results

Preservation of the ophiuroid remains

The ophiuroid fragments described herein were found in insoluble residues of one packstone sample (LC–18–34). The insoluble residues from this sample were dominated by ophiuroid fragments with very few other fossils preserved. The most abundant fragments in this sample were vertebrae, which were brown to amber in color. These were all likely preserved as stereomic molds, with minerals infilling and replacing parts of the skeletal elements, so the stereomic structure is well preserved and visible in SEM imagery.

Phylogenetic analysis

The Bayesian estimate resulted in a well-resolved tree (Fig. 3) with good support for most of the clades. The ophiuroid taxon described herein (*Ophiosuperstes praeparvus* gen. et sp. nov.) holds a basal position in the tree, sister to the Ophintegrida clade. This node gains high posterior probability (96 %), suggesting a robust phylogenetic relationship. The Ophintegrida clade is almost fully resolved, including a basal split into an Ophiacanthida-Ophioscolecida clade (but including the Ophioleucida) and an Amphilepidida clade. The other Triassic ophiuroid included in the tree, *Aplocoma agassizi*, holds a more derived position than *Ophiosuperstes praeparvus*, being situated at the base of the Ophiodermatina.

Systematic palaeontology Class Ophiuroidea Gray, 1840 Subclass Myophiurida Matsumoto, 1915 Infraclass Metophiurida Matsumto, 1913 Superorder Ophintegrida O'Hara, Hugall, Thuy, Stöhr & Martynov, 2017 Order unknown Family unknown Genus Ophiosuperstes nov.

	10 000000000	000000110 00000000
0000200012		0000010110 00002
0101200012		0200100110 01012
0001000012		130011010- 00010
0001000012		130011000- 00010
0001000001		0000000110 00010
0000000000		000000010- 00000
0001000000		??0010010- 00010
002110		02001????? 002
1000100000		0000110110 10001
002110		021110100- 002
0011200110		02111????? 00113
1011200101		0100100110 1011
0000000012		0100100110 0000
66666666		???0?????? ??????
100000013		000000110 1000
1000210012		0300100010 1000
1101111013		0000100110 1101

	1 - 10	11 - 20	21 - 30	31 - 40	41 - 50	51 - 60	61 - 70	71 - 80	81 - 90	91 - 100	101 - 110	111 - 120	121 - 130
Ophienigma sninilimhatum	100000310	0000202211	31100020	0201000113	0-20202011	0100110110	1001010012	00010-2001	1101001010	1020010010	02120001		1011101
Jummannum												00010100	
Ophiochiton fastigatus	10010000	0010002212	31000021	1421100023	0120002011	0100100110	0001200012	?0010-2000	1012011110	1110010102	02120001	 11110000	1000113- -3
Ophiochondrus stelliger	11011-0110	0120101012	30000121	1201000210	0-10002011	0000010110	100000011	?101101002	1101001011	0031023001	10100001	 20100120	1000102- -1
Ophiocopa spatula	11001-0100	0003101211	30000110	1321000221	0-10002011	0000000110	1000010013	11100-2000	1101001011	1032023000	10100001	 20000120	1010112- -2
Ophiocoma echinata	11011-0110	0100121011	20001010	1001000221	1-20013100	0400100110	1001111013	02010-2000	1000001010	1020020100	22100001	 20000120	1010213- -3
Ophiocomina nigra	11011-0110	0100101211	31000021	1201100223	1020110011	0000110110	0001000013	01010-2001	1101001011	1012020000	10100001	 20000120	1010113- -2
Ophiocten sericeum	1000000	0000100211	5012110100	1400100010	0110020111	000000010-	0001100012	00010-2101	0001001210	0012020100	3103110003	000	2000310- -1
Ophioderma longicauda	11011-0110	0100121012	10001020	1021000113	0-21010011	0000100110	1101111000	02100-0000	1000011010	1120022101	32112001	 20000110	1000210- -1
Ophiodoris malignus	11001-00	0000002211	3101000010	0021000121	0-20002011	0100100110	0001200012	00010-1000	1112011110	1020010101	02120001	 10110000	1010113- -3
Ophiolepis superba	20001-00	0010110212	31100000	1301000121	0-20202011	0100100110	0011100000	0?010-0000	1000011010	1010022101	32120011	 11110000	1000200- -0
Ophioleuce seminudum	1001000110	0100002011	3101000000	1321000110	0-00101011	0000000110	100112?001	01010-3000	1101011010	1121123002	01113001	 20001200	10101110
Ophiolimna bairdi	11011-0210	0100101011	30000110	1201000023	0-10102011	0000000110	1000020012	11010-2001	1101001011	1012020001	10100001	 20001120	1000112- -2
Ophiolycus purpureus	11011-1100	0100101012	30000021	1221000003	1-10110011	0000110110	0?1110?012	0100111002	0101011011	1100000101	00100001	 20000110	10101111
Ophiomusium Iymani	2000000	0020100202	5012020100	1311000020	0-10200011	0000000110	0000000000	0000-00000	1000011110	0010021100	0013211013	111	0100010- -0
Ophiomyces delata	01011-0300	1112	00000000	0321110112	0110101011	0100010111	100000012	01110-1000	0111001010	1112020100	01112001	 20001200	10001112
Ophiomyxa pentagona	311-20	0000121211	20001010	1001000010	0-21000011	0000110110	0001100000	010010101-	0001010	1100003100	00112011	 20000110	1000210- -1
Ophionereis porrecta	10011-00	0000002212	3101000010	0021000121	0-20003110	0300100110	1011210012	00010-1000	1013001110	1020010100	02120001	 10110000	1010113- -3
Ophiopallas paradoxa	10011-0110	0100102012	3101010100	0321000021	0-00200011	0000100110	1001121001	0101111000	1?01011010	1121121002	01120001	 20001200	10101110

	1 - 10	11 - 20	21 - 30	31 - 40	41 - 50	51 - 60	61 - 70	71 - 80	81 - 90	91 - 100	101 - 110	111 - 120	121 - 130
Ophiopholis	10101-0211	10101-0211 0110202110 41000021	41000021	1101000222	1-20113100	1300111010	1011210012	0101111000	0103011010	0103011010 1020010010	22120011	:	1001213
aculeata												20100100	
Ophioplax	10001-00	0000202012	3101000110	1321000121	0-10002011	0100100110	0001200012	00000-1000	1012011210	1010010101	02120001	1	1010113-
lamellosa												10110000	-3
Ophiopsila	11011-00	0010302211	31000011	1021000112	1-20203100	0300110110	0001200012	0001200012 00100-0004	1102011110	1100010000	02110011	:	00110-1
guineensis												11111000	
Ophioscolex	311-20	1210	30000020	1200000203	1-00111011	0000100110 002012	002012	01010-201-	0011011	1100020102	02100001	:	00101110
glacialis												20100000	
Ophiotholia	110???00 1212	1212	00000000	0321110112	0110101011	0100000111	0000000012	01010-1000	0111001011	1112020101	01112001	1	00001111
spatifer												20101000	
Ophiothrix	1110010301	1110010301 0020002110 41000020	41000020	010102	2-10113100	1300111110	1001110013	0101211002	0003011010	1020010010	22120011	;	1001203
fragilis												20100100	
Ophiotreta	11011-0110	0110101010 31001010	31001010	1321000111	1-10110011	0000100110	0001000013	?1110-2000	1101011011	1012020000	10100001	:	1000113-
valenciennesi												20000120	-2
Ophiozonella	10000000	10000000 0010100212 31100010	31100010	1301000120	0-20002111	0200100110 1001111012 00010-2000	1001111012	00010-2000	1000011010	1120023101	32133011	-	1000200-
longispina												11110000	0-
Ophiura	10000000	0010100211	5112120100	1400100113	1110020011	00000010-	0001100001	00100-2100	0011011210	0011020100	3200111013	000	2000300-
ophiura													-
Palaeocoma	100????110	0020100010	5012120100	1311000110	0-1001??11	0000110110	0001100000	0001100000 00100-2000	0010001010	1010021101	3213111013	000	1100010-
milleri													
Sigsbeia	20100000	00201?020? ???0?020	???0?020	1101002010	0-1????????	??00101110	1011010100	1011010100 00100-3100	1200010-10	1020012101	02033001	1	1000100-
murrhina												11110000	0-
Praeparvus	66666666666	0???100???	<i>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</i>	<i>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</i>	?????????11	0020200110	005777900	111111111111111111111111111111111111111	???0011010	1010020102	02130001	!	1100000-
superstesii												20000200	0-

Etymology. Ophiosuperstes n. gen. is derived from the Latin adjective *superstes* meaning survivor, referring to the occurrence in the aftermath of the end-Permian mass extinction. Gender masculine.

Type and only species. Ophiosuperstes praeparvus sp. nov.

Diagnosis. as for the species, by monotypy.

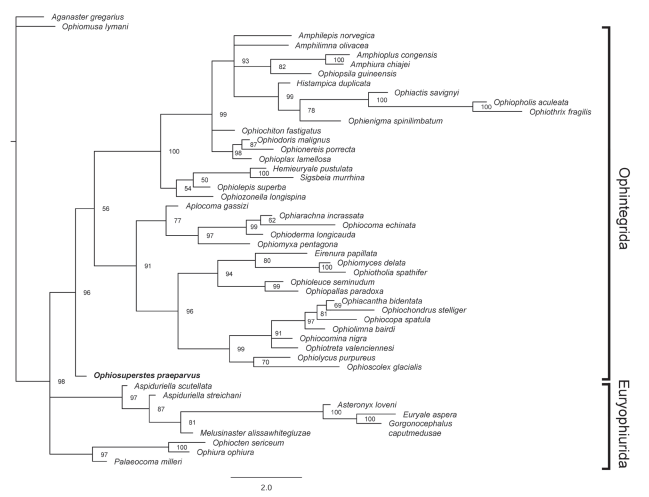


FIGURE 3: Morphology-based phylogenetic tree inferred using MrBayes, showing the position of *Ophiosuperstes praeparvus* gen. et sp. nov. (marked in bold). Numbers at nodes indicate posterior probabilities.

Ophiosuperstes praeparvus **gen. et sp. nov.** Figs. 4–5

Etymology. praeparvus is Latin for very small or minute, as these microfossils were extracted from a small shelly fossil-style assemblage.

Holotype. MnhnL OPH177

Paratypes. MnhnL OPH178 – 190

Type locality. Lost Cabin Springs locality, southern Nevada (Maxwell et al. 2020).

Type stratum. Bed LC-18-34 within the Virgin Limestone Member of the Moenkopi Formation, Spathian, lower Triassic.

Diagnosis. Ophintegrid ophiuroid with stout lateral arm plates of rounded outline, very weak oblique elongated spur on ventro-proximal tip of outer surface, otherwise devoid of spurs or conspicuous outer surface ornamentation; with up to six small, vertical spine articulations slightly sunken into distal edge of lateral arm plate, composed of arched, shifted dorsal and ventral lobes merged at their proximal tips and encompassing the muscle and nerve openings; no sigmoidal fold; tentacle openings developed as small notches lined by a shallow groove in proximal to median lateral arm plates, and reduced to a within-plate perforation in distal lateral arm plates; ventral arm plates trapezoid; radial shields rounded isosceles triangular, without extensions or incisions.

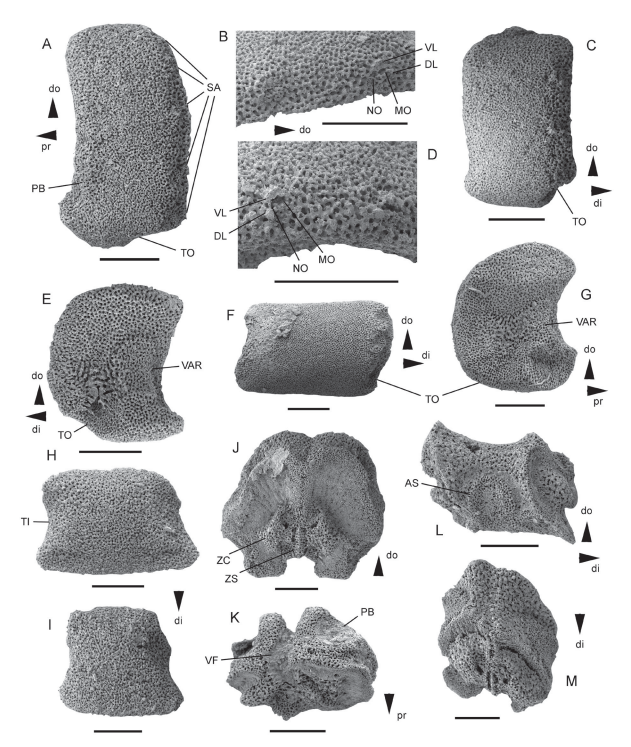


FIGURE 4: Dissociated arm plates of the ophiuroid *Ophiosuperstes praeparvus* gen. et sp. nov., from the Virgin Limestone Member of the Moenkopi Formation, Spathian, lower Triassic, Lost Cabin Springs locality, southern Nevada. A–B: holotype (OPH177), proximal lateral arm plate in external view (A) and with detail of spine articulations (B). C–D: paratype (OPH178), median lateral arm plate in external view (C) and with detail of spine articulations (D). E: paratype (OPH179), median lateral arm plate in internal view. F: paratype (OPH181), distal lateral arm plate in external view. G: paratype (OPH180), median lateral arm plate in internal view. H: paratype (OPH182), proximal ventral arm plate in external view. I: paratype (OPH182), proximal ventral arm plate in distal view. K: paratype (OPH183), median to distal ventral arm plate in external view. J: paratype (OPH184), proximal vertebra in lateral view. K: paratype (OPH185), proximal vertebra in ventral view. L: paratype (OPH186), median vertebra in lateral view. M: paratype (OPH187), median vertebra in dorso-distal view. Abbreviations: AS: articular structure; di: distal; DL: dorsal lobe; do: dorsal; MO: muscle opening; NO: nerve opening; pr: proximal; PB: podial basin; SA: spine articulations; TO: tentacle opening; VAR: vertebral articular ridge; VF: ventral furrow; VL: ventral lobe; ZC: zygocondyle; ZS: zygosphene. All scale bars equal 0.2 mm.

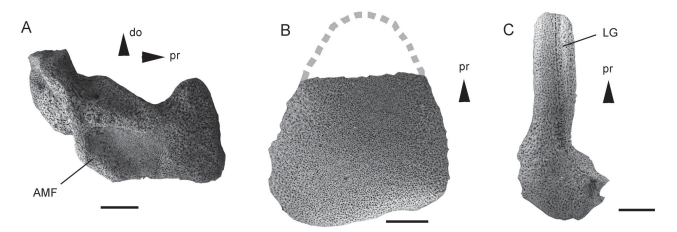


FIGURE 5: Dissociated disc plates of the ophiuroid *Ophiosuperstes praeparvus* gen. et sp. nov., from the Virgin Limestone Member of the Moenkopi Formation, Spathian, lower Triassic, Lost Cabin Springs locality, southern Nevada. A: paratype (OPH188), oral plate in abradial view. B: paratype (OPH189), radial shield in external view, with outline of missing proximal tip reconstructed using a dashed grey line. C: paratype (OPH190) adradial genital plate in dorsal view. Abbreviations: AMF: abradial muscle fossa; do: dorsal; LG: longitudinal groove; pr: proximal. All scale bars equal 0.2 mm.

Description of holotype. MnhnL OPH177 is a dissociated proximal lateral arm plate of stout, rounded outline, approximately two times higher than long, with rounded dorsal and distal edges and protruding ventro-proximal portion; outer proximal edge evenly concave, lined by a relatively broad but poorly defined band of slightly more coarsely meshed stereom with a faint horizontal striation but without spurs other than a very weak oblique elongated spur on the ventro-proximal tip of the outer surface (Fig. 4A); outer surface devoid of constriction, covered by a relatively finely meshed stereom with trabecular intersections transformed into very small tubercles; five relatively small, equal-sized spine articulations along the distal edge, with a slight dorsalward increase in the size of the gaps between the spine articulations, at the same level as the outer surface stereom; spine articulations (Fig. 4B) almost vertical, composed of arched and slightly shifted dorsal and ventral lobes encompassing a small muscle opening and a slightly smaller nerve opening; ventral edge with a small but well developed tentacle notch, lined by a shallow, poorly defined groove; inner side of lateral arm plate (not figured) with a large, well-defined, single vertebral articular ridge composed of the same stereom as the remaining inner side; inner distal edge without spurs; small but clearly defined tentacle notch; no perforations discernible.

Paratype supplements and variation.

Lateral arm plates: median lateral arm plates (MnhnL OPH178 – OPH180) similar to holotype with respect to general outline, outer surface stereom (Fig. 4C), outer proximal edge, shape and position of spine articulations (Fig. 4D) and morphology of inner side (Fig. 4E, G) but with lower height/length ratio, fewer spine articulations (three to four) and smaller tentacle notch. Distal lateral arm plate (MnhnL OPH181) longer than high, of rounded rectangular outline with straight dorsal and ventral edges, with three closely-spaced spine articulations and with tentacle openings developed as within-plate perforation close to ventralmost spine articulation (Fig. 4F).

Vertebrae (MnhnL OPH184 – 187) roughly disc-shaped (proximal vertebrae, Fig. 4J, K) to cylindrical (median and distal vertebrae, Fig. 4L, M), with concave lateral saddle showing a single, tongue-shaped articular surface (Fig. 4L) with the lateral arm plate; large, straight dorsal muscle fossae and much smaller, straight ventral ones; distal face with dorsalwards converging zygocondyles and with a small zygosphene protruding only very little beyond the ventral edge of the zygocondyles (Fig. 4J); deep and broad ventral furrow with small podial basins (Fig. 4K).

Ventral arm plates (MnhnL OPH182 – 183) trapezoid, with weakly convex distal edge, concave lateral edges and narrower, rectangular proximal portion (Fig. 4H, I); no spurs or conspicuous outer surface ornamentation.

Oral plate (MnhnL OPH188) longer than high, slender, fragile, with abradial muscle fossa developed as central depression (Fig. 5A) and small adradial muscle attachment area lining ventral edge of adradio-distal articular surface.

Radial shield (MnhnL OPH189) with proximal portion broken, rounded isosceles triangular in extrapolated outline (Fig. 5B), no signs of extensions or incisions.

Adradial genital plate (MnhnL OPH190) bar-like, with a longitudinal groove and a swollen distal end (Fig. 5C).

Remarks. The ophiuroid remains described in the present paper include various types of skeletal components, i.e. lateral and ventral arm plates, vertebrae, oral plates, genital plates and fragmentary radial shields. The ossicles of every component type show very little variation other than the changes depending on the position within the arm (e.g. Thuy & Stöhr 2011). In particular, the lateral arm plates all belong to the same species, as indicated by the similarities in plate proportions, outer surface ornamentation, spine articulation morphology and position, and vertebral articular structure morphology (Thuy & Stöhr 2011). Furthermore, the various types of ossicles are of compatible size and, in some cases, show corresponding articulation surfaces, e.g. the vertebral articular ridge on the inner side of the lateral arm plates matching the lateral articular ridge of the vertebrae. We therefore conclude that all the ophiuroid remains belong to the same species.

The spine articulations of the lateral arm plates described in the present paper are composed of small muscle and nerve openings encompassed by a ventral and dorsal lobe. This configuration precludes assignment to the Euryophiurida, whose spine articulations lack dorsal and ventral lobes and have the muscle and nerve openings separated by a vertical ridge (O'Hara *et al*, 2018), and instead suggests assignment to the Ophintegrida. Within that superorder, affinities are not clear-cut because the lateral arm plates and the other skeletal parts lack characters that are uniquely found in one of the orders. This observation is in line with our phylogenetic estimate in which the ophiuroid described herein holds a basalmost position at the stem of the Ophintegrida.

Comparison with other fossil ophiuroids is hampered by the lack of detailed morphological analyses. Most descriptions of Triassic ophiuroids published to date are superficial and poorly illustrated, omitting important characters such as lateral arm plate microstructures that have been identified as systematically relevant (e.g. Thuy & Stöhr 2016; O'Hara *et al.* 2018). *Praeaplocoma hessi* Broglio Loriga & Berti Cavicchi, 1972 from the lower Triassic of Italy shares some superficial similarities with the ophiuroid described herein, in particular regarding the outline of the radial shields and the shape of the ventral arm plates. The lateral arm plates of *P. hessi*, however, are bulging. The most important difference, however, was revealed by scanning electron microscopy of newly collected specimens from the Dolomites, close to the type area of the species (Twitchett *et al.* 2005), showing that the spine articulations of *P. hessi* lack dorsal and ventral lobes and instead have their muscle and nerve openings separated by a vertical ridge (Fig. 6). This observation places *Praeaplocoma* in the superorder Euryophiurida and allows a clear distinction from the ophiuroid described herein.

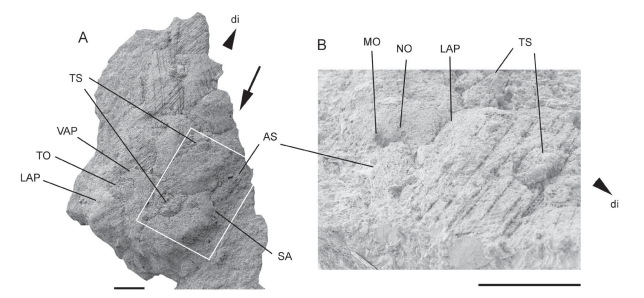


FIGURE 6: Arm fragment of *Praeaplocoma hessi* Broglio Loriga & Berti Cavicchi, 1972, (MHI 1307/nnn) from the Werfen Formation, Lower Triassic, of Weißhorn, Italy. A: arm fragments in ventral view. B: detail of part marked by white rectangle in A. Abbreviations: AS: arm spine; di: distal; LAP: lateral arm plate; MO: muscle opening; NO: nerve opening; SA: spine articulation; TO: tentacle opening; TS: tentacle scale; VAP: ventral arm plate. All scale bars equal 0.2 mm. Arrow in A indicates direction of view shown in B; refer to tentacle scales (TS) for better correspondence between A and B.

Shoshonura brayardi Thuy, 2019 from the lower Triassic of Idaho is another confirmed basal member of the Ophintegrida. It differs from the ophiuroid described herein in the more bulging lateral arm plates and the deeper tentacle notches of the ventral arm plates. Furthermore, the spine articulations are typical of the ophintegrid order Ophiacanthida. *Arenorbis squamosus* (Picard, 1858) from the middle Triassic of central Europe has lateral arm plates with several spurs along the outer proximal edge, and a slightly raised distal portion with larger, unambiguously ophiodermatid-like spine articulations.

Closest similarities are shared with the middle to upper Triassic genus *Aplocoma* d'Orbigny, 1852, in particular with respect to the shape of the lateral arm plates. In contrast to the ophiuroid described herein, however, *Aplocoma* has parallel rather than dorsalward converging zygocondyles on the distal vertebral articulations, ventral arm plates with a pointed rather than straight proximal edge, lateral arm plates with a ventralwards protruding ventro-distal tip and an outer proximal edge with spurs but lacking a fine horizontal striation, and spine articulations that are oblique rather than vertical and that have a weakly developed sigmoidal fold. These characters not only differentiate *Aplocoma* from the ophiuroid described herein but also suggest a more derived position of that genus within the Ophintegrida. Our phylogeny corroborates this assumption and places the type species of *Aplocoma*, *A. agassizi*, at the stem of the order Ophiodermatida, albeit with low support.

Because the ophiuroid described herein differs from all other unambiguously diagnosed fossil ophiuroids, we assign it to the new genus and species *Ophiosuperstes praeparvus*. For the time being, we prefer to eclipse the numerous poorly diagnosed lower Triassic ophiuroid taxa (e.g. Detre & Mihály 1987; Chen *et al.* 2004; Chen & McNamara 2006), because the morphological information available for these taxa is too superficial for a meaningful comparison. A detailed re-evaluation taking into account recent progress in ophiuroid systematics is necessary in order to determine whether the respective type specimens yield sufficient morphological information for an unambiguous diagnosis. Therefore, rather than extrapolating or perpetuating these questionable taxa, we prefer to introduce a new, unambiguously diagnosed taxon as a basis for future comparison.

Discussion

The results of the phylogenetic estimate suggest that *Ophiosuperstes praeparvus* gen. et sp. nov. holds a basalmost position within the Ophintegrida, which is one of the two superorders of the living Ophiuroidea (O'Hara *et al.* 2017, 2018). For a long time, the basal radiation of the modern ophiuroids was supposed to have taken place in the Early Triassic, following an extreme bottleneck at the end-Permian mass extinction (e.g. Smith *et al.* 1995). Molecular and palaeontological data published in recent years, however, show that the diversification of the modern ophiuroids was well under way long before the end-Permian mass extinction (e.g. O'Hara *et al.* 2014, 2017; Thuy *et al.* 2015, 2019). The existence of coeval but more derived ophiuroids such as *Shoshonura brayardi* Thuy, 2019 suggests that *Ophiosuperstes praeparvus* gen. et sp. nov. is probably a persisting member of a more ancient stem ophintegrid group.

The discovery of *Ophiosuperstes praeparvus* gen. et sp. nov. was only possible thanks to the highly unusual preservation of its fossils. In fact, elemental composition analysis using EDS confirmed that the skeletal elements are composed of calcium phosphate, implying that the original high-Mg calcite of the ophiuroid skeleton was completely replaced by apatite. Thanks to phosphatization, the ophiuroid ossicles were left unscathed by the dissolution of the hard, fossiliferous limestone. There has been some debate surrounding the mechanisms behind phosphatization during this time period, in particular phosphogenesis driven by high-energy mixing events (Freeman *et al.* 2013; Milam *et al.* 2017) versus phosphatization occurring in beds with low sedimentation rates and/or sedimentary hiatuses (Freeman *et al.* 2019,). Maxwell *et al.* (2020) observed a link between small shell size and phosphatization, and suggest phosphatization resulted from the combination of warm pore water with low levels of oxygen and the small particle size of the ophiuroid fragments.

Generally, the original high-Mg calcite of echinoderm ossicles recrystallizes into low-Mg calcite during diagenesis, filling the stereom pores and transforming the original porous ossicle into a massive calcite crystal (Gorzelak *et al.* 2016). The fact that stereom pores are preserved to the finest detail even inside the skeletal plates suggests either that phosphatization took place before calcite recrystallisation, or that recrystallisation exceptionally led to a perfect textural stereom preservation. This is in line with previous reports of phosphatized echinoderms, where excellent stereom preservation prevailed (Pisera 1994; Svensson 1999). To our knowledge, this is the first case of an unambiguous phosphatized ophiuroid.

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Author contributions

VM and SBP collected the material and processed the samples; BT and VM picked the skeletal remains and produced the scanning electron microscope images; BT performed the phylogenetic analysis; VM and SBP performed the elemental composition analysis; VM and BT compiled the figures; BT took the lead in writing the manuscript; all authors contributed to the manuscript.

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