

OPEN Collective behaviour in 480-millionyear-old trilobite arthropods from Morocco

Jean Vannier^{1*}, Muriel Vidal², Robin Marchant³, Khadija El Hariri⁴, Khaoula Kouraiss⁴, Bernard Pittet¹, Abderrazak El Albani⁵, Arnaud Mazurier ⁶ & Emmanuel Martin¹

Interactions and coordination between conspecific individuals have produced a remarkable variety of collective behaviours. This co-operation occurs in vertebrate and invertebrate animals and is well expressed in the group flight of birds, fish shoals and highly organized activities of social insects. How individuals interact and why they co-operate to constitute group-level patterns has been extensively studied in extant animals through a variety mechanistic, functional and theoretical approaches. Although collective and social behaviour evolved through natural selection over millions of years, its origin and early history has remained largely unknown. In-situ monospecific linear clusters of trilobite arthropods from the lower Ordovician (ca 480 Ma) of Morocco are interpreted here as resulting either from a collective behaviour triggered by hydrodynamic cues in which mechanical stimulation detected by motion and touch sensors may have played a major role, or from a possible seasonal reproduction behaviour leading to the migration of sexually mature conspecifics to spawning grounds, possibly driven by chemical attraction (e.g. pheromones). This study confirms that collective behaviour has a very ancient origin and probably developed throughout the Cambrian-Ordovician interval, at the same time as the first animal radiation events.

Modern arthropods provide numerous examples of collective behaviour and group migrations. The pine processionary caterpillars use pheromone trails and stimuli from abdominal setae to travel head to tail in large groups and over long distances in search of pupation sites (e.g.²). Similarly, the non-flying juveniles of the desert locust³⁻⁶ engage in gregarious behaviour to form huge mobile foraging groups in reaction to a set of mechanical, olfactory and visual stimuli associated with serotonin release⁵. Collective behaviour also occurs in marine crustaceans such as spiny lobsters (Palinurus) which perform mass single-file migrations⁷⁻¹¹ across open substrates either in possible response to storm-induced environmental disturbances, or for reaching spawning grounds (Palinurus ornatus¹²). Consistent directional positioning is maintained via tactile contact between followers and/ or possible chemical cues¹³. Linear and unidirectional fossil clusters of conspecific trilobite arthropods occur in the Palaeozoic, which have been assumed to result from feeding, reproduction, moulting or sheltering behaviours¹⁴⁻²⁰. They have been also assumed to be a solution for reducing hydrodynamic drag effects within the moving groups²¹. Much more enigmatic are the chain-like associations of bivalved euarthropods from the early Cambrian Chengjiang biota, which have been interpreted to have resulted from collective behaviour and tentatively compared with modern pelagic tunicate chains 22,23. Some non-linear trilobite clusters are seen as evidence for egg deposition in hatching sites²⁴, others as resulting from hypothetical gatherings for protection or moulting²⁵. Most of these reported cases of linear or multidirectional clusters lack potentially important constraints on their interpretation such as the sedimentary environment where these animal groups lived and were buried.

Here we describe and analyse quantitatively, numerous linear clusters of Ampyx priscus from the Lower Ordovician (upper Tremadocian-Floian, ca 480 Ma) Fezouata Shale of Morocco^{26–29}, and show that these alignments of trilobites do not result from passive transportation and accumulation by currents but from a collective

¹Université de Lyon, Université Lyon 1, ENS de Lyon, CNRS, UMR 5276 Laboratoire de géologie de Lyon: Terre, Planètes, Environnement, Bâtiment Géode; 2, rue Raphaël Dubois, F-69622, Villeurbanne, France. ²Université de Brest, CNRS, IUEM-UBO, CNRS, UMR 6538 Laboratoire Géosciences Océan, rue Dumont d'Urville, F-29280, Plouzané, France. ³Musée Cantonal de Géologie, Université de Lausanne, Bâtiment Anthropole, 1015, Lausanne, Switzerland. ⁴Université Cadi-Ayyad, Département des Sciences de la Terre, Faculté des Sciences et Techniques, BP 549, 40000, Marrakesh, Morocco. ⁵Université de Poitiers, UFR SFA, IC2MP, CNRS, UMR 7285 (HydrASA); 5, rue Albert Turpin, Bâtiment B8, TSA 51106, F-86073, Poitiers, France. *email: jean.vannier@univ-lyon1.fr

behaviour. *Ampyx priscus* was probably migrating in groups and used its long projecting spines to maintain a single-row formation by physical contacts possibly associated with mechano-receptors and/or chemical communication. This group behaviour may have been a response to environmental stress due to periodic storms shown by sedimentological evidence, or was associated with reproduction. This record of linear clustering in early euarthropods suggests that intraspecific group-level patterns comparable to those of modern animals already existed 480 million years ago in the early stages of the Great Ordovician Biodiversification Event³⁰.

Results

The trilobite clusters described here from the upper Tremadocian Fezouata Shale (near Zagora, Morocco) are overwhelmingly dominated by Ampyx priscus^{31,32} with rare occurrences of asaphids and calymenids (Fig. 1, Table 1; Supplementary Figs 1-5, Supplementary Tables 1-3) associated with them. Ampyx priscus Thoral, 1935³¹ is a raphiophorid trilobite characterized by a stout glabellar spine and a pair of very long librigenal spines projecting posteriorly (Supplementary Text), which occurs at various horizons through the Fezouata Shale (Supplementary Fig. 1a), as isolated individuals or in linear clusters (see also specimens figured by Chatterton and Fortey¹⁶, plate 1). Trilobites are preserved as internal or external moulds, and are frequently coated with iron oxide. They show no remains of appendages and internal organs. Transverse thin sections (Fig. 1h) indicate that the genal spines had an original tubular structure with a dorsal and ventral furrow and were probably hollow (e.g. infilling; Fig. 1f,g). The glabellar spine is also ventrally striated near its base (Fig. 1j). The distal part of the genal spines runs almost parallel to the longitudinal axis of the animal (Fig. 2a). In each cluster, trilobites are arranged in a linear fashion with their anterior end facing one direction and lie on the surface of a single bedding plane with the dorsal surface of their exoskeleton directed upwards. Only one specimen out of 105 lies upside down (Fig. 2d). The number of A. priscus specimens in clusters varies from 3 to 22. Clusters with a low number of individuals, may be fragmentary. No specimen is disarticulated (i.e.-cephalon including free cheeks, thoracic segments and pygidium in connection), suggesting that they represent carcasses and not exuviae. Ampyx specimens distribute within a relatively narrow size range (more than 75% with a total length (TL) between 16 and 22 mm; Table 1, Supplementary Table 3, Supplementary Fig. 6) and belong to the holaspis stage characterized by a stable number (six) of thoracic segments (Fig. 1b). They probably represent adult or subadult sexually mature animals. Only one juvenile is found in these clusters (see Fig. 2a, most posterior specimen). The angle between the longitudinal axis of two succeeding individuals (α) is usually low and rarely exceeds 45° (see α mean; Table 1, Supplementary Table 3), and the rotation angles within each cluster are randomly clockwise or anticlockwise. This contributes to forming an overall, relatively straight queue with minor local irregularities (Fig. 2). Trilobites facing a direction opposite ($\alpha > 90^{\circ}$) to that of their preceding associates are extremely rare (ca 2%; see Supplementary Table 3). The distance between individuals (D; measured from occipital rings) is relatively short and rarely exceeds twice the body length (TL; D mean < 45 mm; Table 1, Supplementary Table 3) giving the trilobite clusters a cohesive appearance. Succeeding specimens are frequently in contact with each other via their long glabellar and genal spines. The length of the glabellar spine (LglS) slightly exceeds TL, that of the genal spines (Lges) is at least twice as long as TL. Overlapping individuals are frequent (Fig. 2). Comparable monospecific linear clusters dominated by *Ampyx priscus* also occur in the upper Tremadocian of the Montagne-Noire, (Hérault, southern France) and share important features with their Moroccan counterparts, such as the anterior polarity, a relatively short inter-individual distance and low angle (α) variations (Supplementary Fig. 7).

Polished and thin sections through two rock slabs (Fig. 3; Supplementary Fig. 8) were used to characterize the sedimentary background above and below the trilobite clusters. The sediment is a siltstone evenly composed of well-sorted rounded grains (Fig. 3) of low granulometry (20–30 μ m) with a very low percentage of muscovite, clay minerals, and is locally enriched with very fine layers or patches of organic matter (Fig. 3d, Supplementary Figs 8, 9a,b). The original structure of laminae is disturbed by horizontal or subhorizontal bioturbation (Supplementary Fig. 9c,d) which most probably result from the activity of small epibenthic or very shallow endobenthic invertebrates (e.g. euarthropods, worms). No deep burrows or other centimetric sedimentary structures are associated with these trilobite clusters. Bioturbation is indicative of oxic conditions at and slightly below the water-sediment interface. However, its low level of penetration suggests that the redox boundary was probably not deeper than 1.5–2 cm below the sea bed. Elemental mapping (Supplementary Fig. 10) reveals the dominance of Si, Al and K and the ubiquitous presence of Fe (iron oxides and phyllosilicates) and the absence of Ca and S.

Discussion

In situ preservation. Accumulation layers or shell beds are frequent in the fossil record. They typically consist of disarticulated organisms and heterogenous exoskeletal fragments assembled together and often oriented by currents. The *Ampyx* clusters from the Fezouata Shale have none of these diagnostic features. In contrast, they are made up of articulated monospecific individuals and are not associated with sedimentary structures indicative of sea bottom troughs or burrows. Moreover, the consistent anterior polarity of individuals could hardly be explained by the action of currents. We reject the hypothesis of organisms mechanically accumulated along linear submarine reliefs (e.g. between ripple-marks¹⁴). Instead, these linear clusters most likely represent in situ clusters which retain most of the original position of individuals at the time of their death. This interpretation is strongly supported by geological evidence. Detailed sedimentological analyses^{28,29,33,34} show that the Fezouata Shale is characterized by background sedimentation (mainly argillaceous siltstones) repeatedly disturbed by distal storm sequences made of normally graded, very fine sands and coarse siltstones that locally exhibit small oscillation structures, indicating that the depositional setting was located close to the storm wave base. This facies alternation suggests a depositional setting that lays below the fair-weather wave base and above the storm wave base thus corresponding to marine environments ranging from lower shoreface to lower offshore^{28,29,33,34} (Supplementary Fig. 1b). This style of sedimentary dynamics regulated by storms controlled the settlement of animal communities and also played a major role in the preservation of fossils^{33,34}. In the studied area (near Zagora, Morocco) the

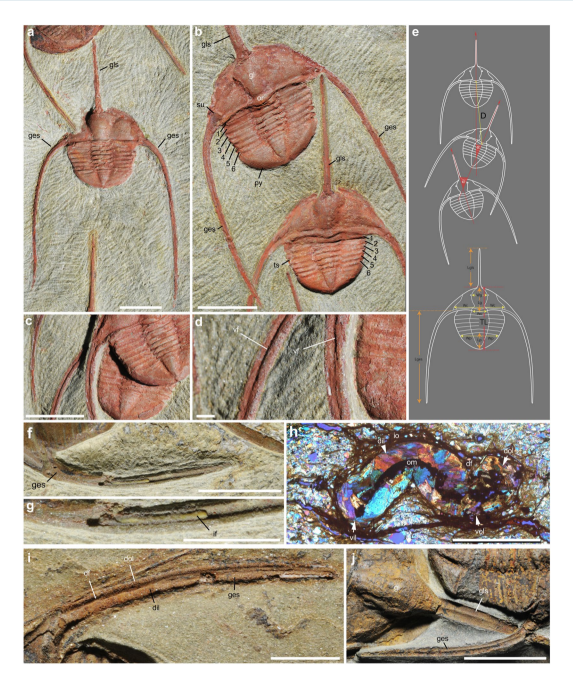


Figure 1. General morphology and parameters of the raphiophorid trilobite *Ampyx priscus* Thoral, 1935, from the Lower Ordovician (Upper Tremadocian-Floian) Fezouata Shale of Morocco (Zagora area). (\mathbf{a} – \mathbf{d}) BOM 2481, overall morphology and details of genal spines. (\mathbf{e}) Parameters used in measurements. (\mathbf{f} , \mathbf{g}) MGL 096718, genal spine showing internal mineralized infilling. (\mathbf{h}) AA.OBZ2.OI.1, transverse thin section through right genal spine (see general view in Supplementary Fig. 8d). (\mathbf{i}) MGL 096727, genal spine. (\mathbf{j}) ROMIP 57013, external mould of glabellar and genal spine showing longitudinal ridge. \mathbf{a} – \mathbf{d} , \mathbf{f} , \mathbf{g} , \mathbf{i} , \mathbf{j} are light photographs. Abbreviations are as follows: α, angle between the longitudinal axis of two successive individuals; D, distance between two successive individuals in clusters (joins central part of occipital rings); df, dorsal furrow; dil, dorsal inner lobe; dol, dorsal outer lobe; ges, genal spine; gl, glabella; gls, glabellar spine; if, mineralized infilling; io, iron oxide; Lg, length of glabella; Lges, length of genal spine; Lgls, length of glabellar spine; Lp, length of pygidium; om, organic matter; py, pygidium; su, suture; TL, total length; vf, ventral furrow; vil, ventral inner lobe; vol, ventral outer lobe; Wc, width of cranidium; Wg, width of glabella; Wo, width of occipital ring; Wp, width of pygidium; 1–6, 1st to 6th thoracic segment. Scale bars: 1 cm in a–c, f, i, j; 5 mm in g,h; 1 mm in d.

raphiophorid trilobite biofacies to which *Ampyx priscus* belongs runs slightly above and below the storm wave base (lower part of shoreface to upper part of offshore; see Supplementary Text and Supplementary Fig. 1b).

The amount of sediment deposited during a storm event was probably sufficient to entomb trilobites and other epibenthic animals *in situ* but not powerful enough to take them away. This process is usually invoked to explain

Clusters	N	TL max	TL min	TL mean	α max	α min	α mean	D max	D min	D mean
LC01	11	19.1	14.9	16.5	34.6	0.1	15.8	43.4	1.7	24.2
LC02	4	14.7	14.2	14.4	15.8	11.1	11.9	36.7	14.8	24.9
LC03	9	26.0	20.1	21.9	21.5	1.5	8.8	43.5	17.8	32.5
LC04	8	19.6	14.9	17.5	12.0	3.7	7.4	71.3	8.6	33.0
LC06	5	18.7	13.0	16.8	14.6	1.0	14.0	65.2	36.3	44.8
LC09	3	22.6	19.4	21.0	48.6	32.9	40.7	31.6	30.7	31.1
LC15	16	23.8	14.1	nd	152.3	0	53.8	90.0	7.6	27.4
LC17	5	20.6	16.8	18.6	38.7	0.7	15.3	23.3	9.4	19.4
LC18	22	20.6	15.6	18.3	60.7	1.5	17.3	46.1	2.5	18.8
LC19	6	20.2	18.9	19.9	26.7	3.6	13.5	29.8	10.0	16.2
LC20	5	21.4	17.6	19.9	13.8	0.2	6.4	48.7	11.8	25.7
LC21	11	21.1	8.6	16.5	15.4	0.2	7.4	71.8	6.4	27.4

Table 1. Summary table of measurements of *Ampyx priscus* linear clusters from the Lower Ordovician (Upper Tremadocian-Floian) Fezouata Shale of Morocco (Zagora area). See Supplementary Tables 1 and 3. Abbreviations are as follows: α , inter-individual angle; D, inter-individual distance; LC, linear cluster; nd, no data; N, number of trilobite (*Ampyx priscus*) specimens; TL, total length.

the exceptional and in situ preservation of marine animals from major Palaeozoic Lagerstätten such as those of the Fezouata Shale^{29,33,34} or the Maotianshan Shale (Chengjiang biota³⁵). Hydrogen sulphide released in water by the storm-generated stirring of anoxic sediment has been proposed to explain the sudden death and in situ preservation of some Devonian trilobite clusters^{21,36}. Similarly, water poisoning may have also participated in rapidly killing Ampyx since neither sedimentary disturbances nor body attitudes indicate strong reaction of trilobites to escape burial. Many trilobites (e.g. 37), including Ampyx priscus (Supplementary Fig. 11) had the capacity to enroll as do modern terrestrial isopods when threatened. The extreme rarity of enrolled specimens in linear clusters would support the hypothesis of a very sudden death either induced by water poisoning or by rapid deposition of sediment which hindered enrollment in most specimens. The absence of sulphur in the sediment surrounding trilobite clusters (see elemental mapping; Supplementary Fig. 10) might result from diagenetic processes. The most plausible scenario that led to the preservation of the *Ampyx* clusters from the Fezouata Shale is the following (Fig. 4). Natural linear clusters of trilobites were entombed by sediment generated by distal storms. The upwards migration of the redox boundary created almost immediate lethal conditions around them and maintained carcasses in anoxic conditions, thus limiting scavenging activities and other degradation processes. In summary, we reject the notion that the linear clusters of trilobites result from passive transportation and accumulation by currents and favour a possible behavioural origin.

Collective behaviour. Various types of trilobite linear clusters have been reported from the Palaeozoic of Morocco¹⁶, Canada¹⁴, Poland^{20,21} and other regions (e.g. Portugal¹⁸), but often lack detailed characterization and key information on their depositional setting. Although some of them tend to be clearly monospecific (e.g. those dominated by Ampyx and $Trimerocephalus^{16,21}$), others are associated with diverse invertebrates (e.g. bivalves, echinoderms, hyoliths) and contain disarticulated exoskeletons, or consist of randomly oriented individuals. Chatterton et al. 14 and Chatterton and Fortey 16 interpreted these clusters as possibly resulting from the congregation of trilobites within sub-horizontal burrows made by other animals such as worms but provide no evidence of burrow margins around the trilobite clusters. Our lithological sections (Fig. 3 and Supplementary Figs 8–10) do not reveal any coloured outlines or disturbances in the sediment surrounding trilobites and thus rule out the hypothesis that trilobites may have sheltered collectively in burrows before being trapped and buried in situ. Chemical cues or visual attraction to food often drive modern animals to gather and form monospecific clusters of scavengers over carcasses. These aggregations result from individual taxic responses (e.g. myodocope ostracods³⁸) and do not strictly correspond to a coordinated group behaviour. The trilobite clusters described here are unlikely to represent such aggregations unless individuals gathered to feed on narrow linear food patches. Moreover, their consistent unidirectional polarity and the lack of associated food remains or traces in sediment make this hypothesis improbable. Another option related to feeding is that *Ampyx* may have fed on suspended particles thus prompting individuals to orientate themselves in the direction of current. If so, we would rather find trilobites lying side by side facing the same direction than aligned in queues.

We propose that the linear clusters formed by *Ampyx priscus* result from the coordinated gathering and locomotion of individuals and therefore suggests a collective and synchronized behaviour. We hypothesize that these trilobites moved in small groups on the seafloor, keeping a single-row formation by physical contacts via their long projecting spines and antennules and/or through chemical communication. Similarly, extant spiny lobsters perform mass single-file migrations by maintaining tactile contact between the tail fan of one individual and the antennular ramus and the tips of the anterior-most walking legs of its follower^{7,10}. Such mechanical contacts appear to be essential for group cohesion and for optimal coordinated locomotion. Experimental studies⁵ have shown that the gregarious behaviour of desert locusts was linked to an increase of neurotransmitter (serotonin) driven by mechanical stimuli, visual or olfactory pathways^{39–41}. Knowing that *Ampyx priscus* was blind, we hypothesize that mechanosensory stimulation via both genal and glabellar spines, or/and chemical cues, may have been a major trigger that maintained group behaviour. No trace fossils seem to be associated with the

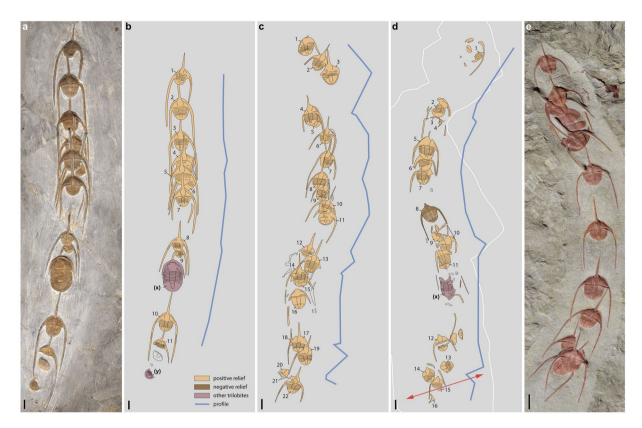


Figure 2. Linear clusters of the raphiophorid trilobite *Ampyx priscus* Thoral, 1935³¹, from the Lower Ordovician (upper Tremadocian-Floian) Fezouata Shale of Morocco (Zagora area). (**a,b**) AA.TER.OI.12 (see Supplementary Fig. 2a). (**c**) MGL 096727 (see Supplementary Fig. 5a). (**d**) AA.TER.OI.13 (see Supplementary Fig. 2b). (**e**) BOM 2461 (see Supplementary Fig. 2f). (**a,e**) are light photographs. Line drawings from photographs. Segmented blue lines in (**b–d**) join the central part of occipital rings of trilobites. Red arrows indicate the position of polished section in Fig. 3. Abbreviations are as follows: (**x**), *Asaphellus* aff. *jujuanus* (asaphid trilobite); (**y**), juvenile asaphid trilobite. Scale bars: 1 cm.

trilobite clusters. Possible reasons for this are that either *Ampyx* made no deep imprints into the sediment or that the substrate conditions were unfavourable to preserving traces.

Microscopic observations were unable to locate potential sites (e.g. sensilla pores⁴²) of mechano- or chemoreception on the exoskeleton of *A. priscus* due to unfavourable conditions of preservation (internal/external moulds with rare cuticular remains). Its remarkably long spiny projections seem to offer optimal conditions for individuals to contact via multiple points (Supplementary Fig. 12) thus increasing the probability for communication via mechanosensory stimulation. This unusual exoskeletal configuration may have facilitated alignment and cohesion within the group. Interestingly, *Ampyxinella (Eoampyxinella) villebruni*, a raphiophorid trilobite affiliated to *Ampyx* (Supplementary Fig. 13), which lacks a frontal glabellar spine, forms linear clusters characterized by a lower directional consistency compared with those of *Ampyx priscus*. Chemical communication plays a crucial role in the inter-individual relationships of a wide range of extant animals including crustaceans (e.g. ^{43–45}). Various types of chemosensory sensilla occur over the crustacean body, some of them such as olfactory aesthetascs concentrating on antennules⁴². Such receptors are assumed to be present in Cambrian mandibulates such as *Waptia* ^{46,47}. Similarly, pores housing a central seta found in Cambrian agnostids⁴⁸ and trilobites (e.g. ⁴⁹) may have had a comparable chemosensory function. Chemical communication is often invoked to explain the group behaviour of other trilobites (e.g. blind phacopid *Trimerocephalus* ^{20,21,50}) and indeed may have played an important role in the collective behaviour of *Ampyx*. However, none of these trilobites provide detailed information on the nature and location of possible receptors.

Fifty-seven percent of the studied clusters from the Fezouata Shale have one and more rarely two non-*Ampyx* trilobite elements which are dominantly complete carapaces of asaphids (mainly *Asaphellus* aff. *jujuanus*), calymenids (*Parabathycheilus* sp.) and dalmanitids; (*Toletanaspis* sp.) (see Supplementary Table 2). Four of these complete trilobites follow the direction of the *Ampyx* cluster in which they seem to be incorporated, whereas the remaining individuals point towards the opposite (2) or intermediate directions (2). In contrast with *Ampyx*, these trilobites had eyes and therefore may have been able to visually detect, track and possibly follow individuals moving in their surroundings. However, the small number of data does not allow for determining whether they may have joined the *Ampyx* groups for some opportunistic reasons or may simply have crossed their migratory path by pure coincidence. As far as we know from the literature⁷⁻¹¹, no other crustacean species participates in the migration of extant spiny lobsters.

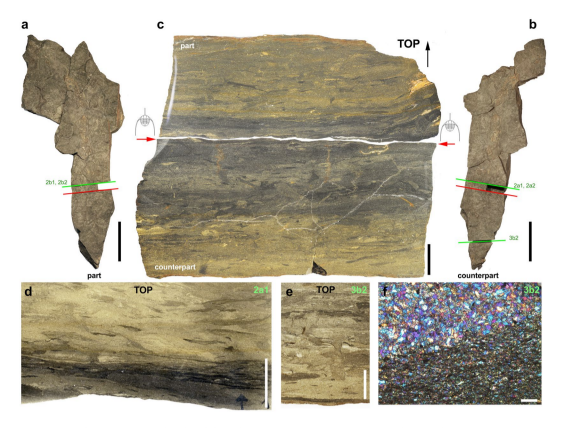


Figure 3. Sediment associated with trilobite clusters. (**a,b**) AA.TER.OI.13 (see also Supplementary Figs 8–10), part and counterpart, from the Lower Ordovician (upper Tremadocian-Floian) Fezouata Shale of Morocco (Zagora area). (**c**) Polished section, general view. (**d,e**) Thin sections showing details of sedimentary structure. (**f**) Thin section showing grain size and local enrichment in organic matter (lower part). Red arrows indicate bedding plane with Ampyx clusters. Location of thin sections indicated by green lines and numbers (see also Supplementary Figs 9–11). Scale bars: 10 cm in a, b; 1 cm in c–e; 50 μ m in f.

Triggers and drivers, functions and benefits

Spawning congregations and synchronized moulting. Congregation of sexually mature individuals are frequent in extant euarthropods and are often related to reproduction or moulting (e.g. ^{12,51}). In modern horseshoe crabs, spawning is synchronous with sexually mature individuals gathering along the shore in large numbers for mating (e.g. ⁵²⁻⁵⁵). This behaviour is synchronized through both visual and pheromone communication ⁵⁶. Ampyx may have performed comparable group migrations to distant spawning grounds during the reproductive season, as proposed by Blazejowki *et al.* ²⁰ for other trilobites. This hypothesis is supported by the fact that Ampyx clusters almost exclusively consist of adult or sub-adult holaspid stages. However, no morphological traits (e.g. body size, spines ^{57,58}) point to co-existing dimorphs within clusters. Synchronized moulting is frequent in modern crustaceans (e.g. krill ⁵⁹) and insects and is assumed to have occurred also in Cambrian euarthropods such as the trilobite Balcoracania dailyi ⁶⁰ from the lower Cambrian (Series 2, Stage 4) of Australia and Canadaspis and Alalcomenaeus ⁶¹ from the middle Cambrian Burgess Shale (Series 3, Stage 5) of British Columbia, Canada. This phenomenon brings together a large number of conspecific individuals but does not create any directional polarity among the group (e.g. linear gatherings). Moreover, synchronized moulting, by definition, releases numerous exuviae which are absent from the Ampyx clusters.

Hydrodynamic cues. Field and laboratory studies on extant spiny lobsters from the Bahamas region have highlighted the possible relation between collective migrations and environmental disturbances⁷⁻¹¹. A drop in water temperature, higher water turbidity and intense current induced by seasonal storms are assumed to be the main triggers for the mass migrations of these crustaceans, which always take place from highly disturbed shallow coastal areas to the edge of oceanic channels. Most migrants are late juveniles and adults but do not congregate for spawning. The hypothesis that hydrodynamic cues may have driven a comparable behaviour in Ampyx is realistic considering that these trilobites were potentially exposed to periodic environmental disturbances generated by storms (see sedimentological evidence above). Ampyx is known to occur preferentially across the storm wave base and preferentially in the lower shoreface and upper offshore environments^{28,29,62} (Supplementary Fig. 1b) suggesting possible migrations from storm-influenced to quieter and deeper areas. We hypothesize that its behaviour may have been a collective response to physical stress. Collective migrations of modern crustaceans are assumed to have important advantages over individual ones. Travelling in queues decreases hydrodynamic drag and saves energy (e.g. spiny lobsters^{8,10}) and also tends to reduce the probability of detection and attacks from predators by creating confusion in their visual perception (e.g.⁶³). Ampyx clusters have a relatively small number of individuals

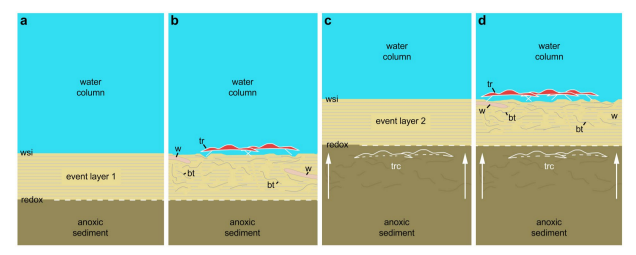


Figure 4. Scenario to explain the *in situ* preservation of the *Ampyx* linear clusters from the Lower Ordovician (Upper Tremadocian-Floian) of Morocco. (a) Deposition of a distal tempestite (event layer 1). (b) Epibenthic (e.g. trilobites) and shallow endobenthic (e.g. possible worms) organisms settle and generate bioturbation above red-ox boundary. (c) Second storm event layer entombs epibenthic fauna *in situ*; red-ox boundary moves upwards (white arrows). (d) New faunal recolonization. According to Vaucher *et al.*³⁴, distal storm deposits are relatively thin (less than 5 cm) and consist of a waning (base) and waxing (top) phases (subdivision not represented in this diagram), and depositional environment is that of the distal lower shoreface with a possible water depth of approximately 30–70 m. Bioturbation is based on polished and thin sections (Fig. 3 and Supplementary Figs 8 and 9). Abbreviations are as follows: bt, bioturbation; tr, trilobite group (*Ampyx*); trc, trilobite carcasses (*Ampyx*); w, worm; wsi, water-sediment interface.

and contrast with the long queues seen in spiny lobsters. This might be explained by a natural tendency of trilobites to form smaller groups or by the dispersion of original larger linear clusters by currents. Hydrodynamic signals (direction of storm back current) possibly relayed by sensory setae over the carapace or appendages may also have played an important role in the polarization of locomotion but tactile contacts and/or chemical communication were probably essential to maintaining group cohesion.

In summary, two options can be seen as the most likely (Fig. 5): (1) that of a collective behaviour triggered by hydrodynamic cues in which mechanical stimulation detected by motion and touch sensors may have played a major role; (2) that of a possible seasonal reproduction behaviour leading to the migration of sexually mature conspecifics to spawning grounds, possibly driven by chemical attraction (e.g. pheromones). These options are not mutually exclusive. *Ampyx* may have alternatively responded to environmental stress and reproduction signals by adopting the same behaviour.

Origin of collective behaviour. *Ampyx* shows that collective behaviour in arthropods has a very deep ancestry back to the lower Palaeozoic. This behaviour was necessarily associated with a communication system between individuals involving motion and mechanical sensors, chemical signals and possibly neurotransmitters (e.g. serotonin⁵). Although this behaviour was not mediated by sight because *Ampyx* was blind, it implies neural complexity and the ability to process signals. Numerous arthropods had already evolved sophisticated brains (e.g. ⁶⁴), and sensory organs such as antennae and compound eyes (e.g. ^{47,65-67}) by the Cambrian although their exact functional capabilities often remain conjectural.

Ampyx shows how a 480-million-year-old euarthropod may have integrated its neural complexity into a temporary collective behaviour related to seasonal reproduction or triggered by environmental cues. This behaviour is likely to have been widespread among trilobites throughout the Palaeozoic (e.g. Devonian¹⁹). Another case of enigmatic collective behaviour has been reported in shrimp-like bivalved euarthropods from the early Cambrian Chengjiang biota, which form seemingly closely tied chain-like monospecific associations (Synophalos^{22,23}, ca 520 Ma). These chains strongly recall the migratory queues of spiny lobsters, processionary moth caterpillars or ants but are interpreted^{22,23} as being assembled in the water column and then deposited on the sea bottom through passive sinking. This scenario would suppose an extremely robust interlocking system between individuals and a synchronized group locomotion that have no counterparts in extant arthropods. Males of extant clam shrimps have specialized organs to clasp the edge of the female's carapace during pairing⁶⁸. However, their behaviour is limited to two sexual partners. Although much information is missing concerning the Synophalos clusters (e.g. appendages, relation to sediment, measurements), it appears more likely that, as with Ampyx, they represent epibenthic arthropods migrating in groups and buried *in situ* by storm-related or turbiditic sedimentary events. Collective behaviour associated with communication and recognition systems probably evolved through natural selection as the Cambrian radiation proceeded, and developed more extensively during the Great Ordovician Biodiversification Event when ecosystems became increasingly complex. Improving the chances of reproduction and survival to environmental stress are among the advantages that such behaviour may have conferred to euarthropods.

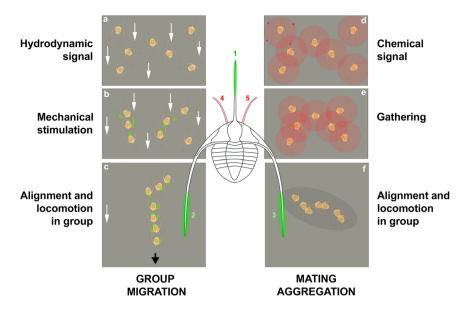


Figure 5. Two non-exclusive hypotheses to explain the linear clusters of *Ampyx priscus* from the Lower Ordovician of Morocco. (**a-c**) Response to oriented environmental stress (e.g. storms); hydrodynamic signal (higher current velocity represented by white arrows) received by motion sensors triggers re-orientation of individuals; mechanical stimulation and/or possible chemical signals cause gathering, alignment and locomotion in group. (**d-f**) Seasonal reproductive behaviour; chemical signals (e.g. pheromones; see red circles and red arrows) cause attraction and gathering of sexually receptive individuals (males and females) and migration to spawning grounds. The alignment of individual may have been controlled by mechanical stimuli (as in **a-c**). Olfactive and mechanical sensors were probably located on the antennules (pink areas 4, 5), and genal and glabellar spines (green areas 1–3), respectively. The exact location of mechanoreceptors is uncertain (possibly on high-relief exoskeletal features such as the glabella).

Methods

Fossils specimens were photographed using a digital camera (D3X-Nikon with Nikon Micro-Nikkor 60 mm lens) and measured (lengths, distances, angles) from high-resolution digital images by using Image J, a public domain processing program. Polished and lithological thin sections were made using standard methods and observed under binocular stereo-microscope (Leica MZ125 and Leica DM750P). Elemental maps were acquired using a Tornado M4 micro-XRF system (Bruker, Germany) equipped with a silicon drift detector and a Rh source operating at $50\,\mathrm{kV}$ and $600\,\mu\mathrm{A}$. A spot size of $40\,\mu\mathrm{m}$ was employed with dwell times of $7\,\mathrm{ms/pixel}$, and mapping was performed under vacuum. Image processing included spectral deconvolution and 3-pixel averaging. Tomographic images of ROMIP 57013 were obtained via the same methods and with the same machine as in Kouraiss $et~al.^{69}$.

Data availability

All figured specimens are deposited in fossil collections: Cadi Ayyad University, Faculty of Sciences and Techniques, Marrakesh, Morocco (AA numbers); Musée Cantonal de Géologie, Lausanne, Suisse (MGL); Museum d'Histoire Naturelle de Marseille, France (MHNM); Royal Ontario Museum, Toronto, Canada (ROMIP); Palaeontological Collections, Patrick Bommel, Bizes-Minervois, France (BOM); Palaeontological Collections, Laurent Lacombe, Ouveillan, France (LAC). Correspondence and requests for materials should be addressed to JV.

Received: 19 March 2019; Accepted: 6 September 2019;

Published online: 17 October 2019

References

- 1. Sumpter, D. J. T. Collective animal behaviour. Princeton University Press, 312 p (2010).
- Fitzgerald, T. D. Role of trail pheromone in foraging and processionary behavior of pine processionary caterpillars *Thaumetopoea pityocampa*. Journal of Chemical Ecology 29, 513–532 (2003).
- 3. Steedman, A. Locust handbook. Overseas Development Natural Resources Administration (Great Britain), 204 p (1990).
- 4. Buhl, J. et al. From disorder to order in marching locusts. Science 312, 1402-1406 (2006).
- 5. Anstey, M. L. *et al.* Serotonin mediates behavioral gregarization underlying swarm formation in desert locusts. *Science* **323**, 627–630 (2009).
- 6. Stevenson, P. A. The key to Pandora's box. Science 323, 594-595 (2009).
- 7. Herrnkind, W. F. Queuing behavior of spiny lobsters. Science 164, 1425 (1969).
- 8. Herrnkind, W. F. Evolution and mechanisms of mass single-file migration in spiny lobster: Synopsis. *Contributions in Marine Science* 27, 197–211 (1985).
- 9. Herrnkind, W. F. & McLean, R. Field studies of homing, mass emigration, and orientation of the spiny lobster, *Palinurus argus*. *Annals of the New York Academy of Science* **188**, 359–377 (1971).

- Herrnkind, W. F. et al. Descriptive characterization of mass autumnal migrations of spiny lobster, Palinurus argus. Proceedings of the Gulf and Caribbean Fisheries 25, 79–98 (1973).
- 11. Kanciruk, P. & Herrnkind, W. F. Mass migration of spiny lobster, *Panulirus argus* (Crustacea: Palinuridae): behavior and environmental correlates. *Bulletin of Marine Science* 28, 601–623 (1978).
- 12. Moore, R. & McFarlane, J. W. Migration of the ornate rock lobster, *Panulirus ornatus* (Fabricius), in Papua New Guinea. *Australian Journal of Marine and Freshwater Research* 35, 197–212 (1984).
- 13. Hay, M. E. Marine chemical ecology: chemical signals and cues structure marine populations, communities, and ecosystems. *Annual Review of Marine Sciences* **2009**, 193–2012 (2009).
- 14. Chatterton, B. D. E. et al. Cryptic behaviour in trilobites: Cambrian and Silurian examples from Canada, and other related occurrences. Special Papers in. Palaeontology 70, 157–173 (2003).
- 15. Cherns, L. et al. Tunneling trilobites: habitual infaunalism in an Ordovician carbonate seafloor. Geology 34, 657-660 (2006).
- 16. Chatterton, B. D. E. & Fortey, R. A. In Advances in trilobite research (eds Rábano, I., Gozalo, R. & García-Bellido, D. 73–79 (Cuadernos del Museo Geominero, No. 9. Instituto Geológico y Minero de España, 2008).
- 17. Paterson, J. R. et al. A. in Advances in trilobite research (eds Rábano, I., Gozalo, R. & García-Bellido, D. 313–318 (Cuadernos del Museo Geominero, No. 9. Instituto Geológico y Minero de España, 2008).
- 18. Gutiérrez-Marco, J. C. et al. Giant trilobites and trilobite clusters from the Ordovician of Portugal. Geology 37, 443-446 (2009).
- 19. Radwanski, A. et al. Queues of blind phacopid trilobites *Trimerocephalus*: a case of frozen behaviour of Early Famennian age from the Holy Cross Mountains, Central Poland. Acta Geologica Polonica 59, 459–481 (2009).
- Blazejowski, B. et al. Ancient animal migration: a case study of eyeless, dimorphic Devonian trilobites from Poland. Palaeontology 59, 743–751 (2016).
- 21. Trenchard, H. et al. Trilobite 'pelotons': possible hydrodynamic drag effects between leading and following trilobites in trilobite queues. Palaeontology 60, 557–569 (2017).
- 22. Hou, X.-G. et al. Collective behavior in an early Cambrian arthropod. Science 322, 224 (2008).
- 23. Hou, X.-G. *et al.* A new arthropod in chain-like associations from the Chengjiang Lagerstätte (Lower Cambrian), Yunnan, China. *Palaeontology* **52**, 951–961 (2009).
- 24. Schwimmer, D. R. & Montante, W. M. A cambrian meraspid cluster: evidence of trilobite egg deposition in a nest site. *Palaios* 34, 254–260 (2019).
- 25. Gutiérrez-Marco, J. C. et al. The nileid trilobite *Symphysurus* from upper Tremadocian strata of the Moroccan Anti-Atlas: taxonomic reappraisal and palaeoenvironmental implications. *Fossils and Strata* 64, 155–171 (2019).
- 26. Van Roy, P. et al. Ordovician faunas of Burgess Shale type. Nature 465, 215-218 (2010).
- 27. Martin, E. L. O. et al. Age and environmental setting of the Lower Ordovician Fezoauta Biota (Zagora, Morocco). Gondwana Research 34, 274–283 (2016).
- 28. Martin, E. L. O. *et al.* In The Fezouata Biota: an exceptional window on the Cambro-Ordovician faunal transition (eds Lefebvre, B., Lerosey-Aubril, R., Servais, Th. & Van Roy, P.). 142–154 (Special Issue, vol. 460 Palaeogeography, Palaeoclimatology, Palaeoecology, 2016)
- Lefebvre, B. et al. In The Fezouata Biota: an exceptional window on the Cambro-Ordovician faunal transition (eds Lefebvre, B., Lerosey-Aubril, R., Servais, Th. & Van Roy, P.). 1–6 (Special Issue, vol. 460 Palaeogeography, Palaeoclimatology, Palaeoecology, 2016).
- Servais, T. et al. The Great Ordovician Biodiversification Event (GOBE): the palaeoecological dimension. Palaeogeography, Palaeoclimatology, Palaeoecology 294, 99–119 (2016).
- 31. Thoral, M. Contribution à l'étude paléontologique de l'Ordovicien inférieur de la Montagne Noire et révision sommaire de la faune cambrienne de la Montagne Noire. Imprimerie de la Charité, Montpellier, 362 pp (1935).
- 32. Vidal, M. Trilobites (Asaphidae et Raphiophoridae) de l'Ordovicien inférieur de l'Anti-Atlas, Maroc. *Palaeontographica A* **251**, 39–77 (1998a).
- 33. Vaucher, R. et al. In The Fezouata Biota: an exceptional window on the Cambro-Ordovician faunal transition (eds Lefebvre, B., Lerosey-Aubril, R., Servais, Th. & Van Roy, P.). 24–34 (Special Issue, vol. 460 Palaeogeography, Palaeoclimatology, Palaeoecology, 2016).
- 34. Vaucher *et al.* A wave-dominated, tide-modulated model for the Lower Ordovician of the Anti-Atlas, Morocco. *Sedimentology* **64**, 777–807 (2017).
- 35. Zhu, M.-Y. Fossilization modes in the Chengjiang Lagerstätte (Cambrian of China): testing the roles of organic preservation and diagenetic alteration. *Palaeogeography, Palaeoclimatology, Palaeoecology* **220**, 31–46 (2005).
- 36. Brett, C. E. et al. Mid-Paleozoic trilobite Lagerstätten: models of diagenetically enhanced obrution deposits. Palaios 27, 326–345 (2012)
- 37. Esteve, J. et al. Sphaeroidal enrolment in middle Cambrian solenopleuropsine trilobites. Lethaia 43, 478-493 (2010).
- 38. Vannier, J. et al. Feeding in myodocopid ostracods: functional morphology and laboratory observations from videos. *Marine Biology* 132, 39–408 (1998).
- Roessingh, P. et al. Effects of sensory stimuli on the behavioural phase state of the desert locust, Schistocerca gregaria. Journal of Insect Physiology 44, 883–893 (1998).
- 40. Simpson, S. J. et al. Gregarious behavior in desert locusts is evoked by touching their back legs. Proceedings of the National Academy of Sciences 98, 98 (7) 3895–3897 (2001).
- 41. Rogers, S. M. et al. Mechanosensory-induced behavioural gregarization in the desert locust Schistocerca gregaria. Journal of Experimental Biology 206, 3991–4002 (2003).
- 42. Hallberg, E. & Skog, M. Chemosensory sensilla in crustaceans. In Chemical Communication in Crustaceans (eds Breithaupt, T. & Thiel, M.). 103–121. Springer (2011).
- 43. Atema, J. Chemical signals in the marine environment: dispersal, detection and temporal signal analysis. *Proceedings of the National Academy of Sciences* **92**, 62–66 (1995).
- 44. Aggio, J. & Derby, C. D. Chemical communication in lobsters. In *Chemical Communication in Crustaceans* (eds Breithaupt, T. & Thiel, M.). 239–256. Springer (2011).
- 45. Breithaupt, T. Chemical communication in crayfish. In *Chemical Communication in Crustaceans* (eds Breithaupt, T. & Thiel, M.). 257–276. Springer (2011).
- 46. Strausfeld, N. J. Waptia revisited, intimations of behaviours. *Arthropod Structure and Development* **45**, 173–184 (2016).
- 47. Vannier, J. et al. Waptia fieldensis Walcott, a mandibulate arthropod from the middle Cambrian Burgess Shale. Royal Society Open Science 5, 172206 (2018).
- 48. Müller, K. J. & Waloszek, D. Morphology, ontogeny, and life habit of *Agnostus pisiformis* from the Upper Cambrian of Sweden. *Fossils and Strata* 19, 1–124 (1987).
- 49. Miller, J. The sensory fields and life mode of *Phacops rana* (Green, 1832) (Trilobita). *Proceedings of the Royal Society of Edinburgh* **69**, 337–367 (1976).
- 50. Karim, T. & Westrop, S. R. Taphonomy and paleoecology of Ordovician trilobite clusters, Bromide Formation, South-central Oklahoma. *Palaios* 17, 394–403 (2002).
- 51. Linnane et al. Movement patterns of the southern rock lobster, Jasus edwardsii, of South Australia. New Zealand Journal of Marine and Freshwater Research 39, 335–346 (2005).

- 52. Sekiguchi, K. Biology of horseshoe crabs. Science House, Tokyo (1988).
- 53. Chiu, H. M. C. & Morton, B. The biology, distribution and status of horseshoe crabs, Tachypleus tridentatus and Carcinoscorpius rotundicauda (Arthropoda: Chelicerata) in Hong Kong: Recommendations for conservation and management. Final Report. The Swire Institute of Marine Science, The University of Hong Kong, 1999.
- 54. Chiu, H. M. C. & Morton, B. The distribution of horseshoe crabs (*Tachypleus tridentatus* and *Carcinoscorpius rotundicauda*) in Hong Kong. *Asian Marine Biology* 16, 185–196.
- 55. Mattei, J. et al. Reproductive Behavior in Horseshoe Crabs: Does density matter? Current. Zoology 56, 634-642 (2010).
- 56. Schwab, R. L. & Brockmann, J. The role of visual and chemical cues in the mating decisions of satellite male horseshoe crabs, *Limulus Polyphemus*. *Animal Behaviour* 74, 837–846 (2007).
- 57. Hughes, N. C. & Fortey, R. A. In Ordovician Odyssey (eds Cooper, J. C., Droser, M. L. & Finney, S. C.), 419–421 (SEPM Pacific Section, Los Angeles, 1995).
- 58. Knell, R. J. & Fortey, R. A. Trilobite spines and beetle horns: sexual selection in the Palaeozoic? Biology Letters 1, 196-199 (2005).
- 59. Tarling, G. A. & Cuzin-Rudy, J. Synchronization in the moulting and spawning activity of northern krill (*Meganyctiphanes norvegica*) and its effect on recruitment. *Limnology and Oceanography* 48, 2020–2033 (2003).
- 60. Paterson, J. R. et al. Taphonomy and palaeoecology of the emuellid trilobite Balcoracania dailyi (early Cambrian, South Australia). Palaeogeography, Palaeoclimatology, Palaeoecology 249, 302–321 (2016).
- 61. Haug, J. T. et al. Demecology in the Cambrian: synchronized moulting in arthropods from the Burgess Shale. BMC Biology 11, 64 (2013).
- 62. Vidal, M. The trilobite biofacies model: a test in the Early Ordovician of the Anti-Atlas, Morocco. Compte-rendu de l'Académie des Sciences Paris, Earth and Planetary Sciences 327, 327–333 (1998b).
- 63. Jeschke, J. M. & Tollrian, R. Prey swarming: which predators become confused and why? Animal Behaviour 74, 387-393 (2007).
- 64. Ma, X.-Y. et al. Complex brain and optic lobes in an early Cambrian arthropod. Nature 490, 259-262 (2012).
- 65. Schoenemann, B. et al. Structure and function of a compound eye, more than half a billion years old. P Proceedings of the National Academy of Sciences 114, 13489–13494 (2017).
- 66. Lee, M. S. Y. et al. Modern optics in exceptionally preserved eyes of Early Cambrian arthropods from Australia. Nature 474, 631–634 (2011).
- 67. Paterson, J. R. et al. Acute vision in the giant Cambrian predator *Anomalocaris* and the origin of compound eyes. *Nature* 480, 237–240 (2011).
- 68. Sigwardt, Z. M. S. & Olesen, J. Mating behaviour in Laevicaudatan clam shrimp (Crustacea, Branchiopoda) and functional morphology of male claspers in a phylogenetic context: A video-based analysis. *Plos One* 9(1): e84021.
- Kouraiss, K. et al. X-ray microtomography applied to fossils preserved in compression: Palaeoscolescid worms from the Lower Ordovician Fezouata Shale. Palaeogeography, Palaeoclimatology, Palaeoecology 508, 48–58 (2018).

Acknowledgements

We thank D. Rudkin (Royal Ontario Museum) and F. Escuillé for the loan of specimens, P. Bommel, L. Lacombe for access to collections, S. Jouve (Museum, Marseille) and B. Lefebvre for photographs, P. Tieffenbach (Univ. Brest) for thin-sections, the PLATeforme INstrumentale d'Analyses (PLATINA, IC2MP, Poitiers) and the Région Nouvelle Aquitaine for access to microtomography, M. Williams (Univ. Leicester, UK) for the critical reading of the MS, M. Laurans (Ifremer, Brest) for information on extant crustaceans, Q. Ou (Univ. Geosciences Beijing) for information and discussion on Cambrian linear clusters, and J.R. Paterson (Univ. New England) and A. W. Owen (Univ. Glasgow) for their insightful reviews. The authors acknowledge support from the Académie Hassan II of Science and Technology.

Author contributions

J.V. designed the research and wrote the paper with his co-authors. E.M. analysed trilobite clusters, M.V. provided data on trilobite morphology, distribution and biofacies; B.P. provided sedimentological analysis; R.M., K.E.H. and K.K. provided fossil specimens and data on geological background, A.E.A. and A. M. performed X-Ray Microtomography.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41598-019-51012-3.

Correspondence and requests for materials should be addressed to J.V.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019