Phylogenomic resolution of sea spider diversification through integration of multiple data classes

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Abstract

Despite significant advances in invertebrate phylogenomics over the past decade, the higher-level phylogeny of Pycnogonida (sea spiders) remains elusive. Due to the inaccessibility of some small-bodied lineages, few phylogenetic studies have sampled all sea spider families. Previous efforts based on a handful of genes have yielded unstable tree topologies. Here, we inferred the relationships of 89 sea spider species using targeted capture of the mitochondrial genome, 56 conserved exons, 101 ultraconserved elements, and three nuclear ribosomal genes. We inferred molecular divergence times by integrating morphological data for fossil species to calibrate 15 nodes in the arthropod tree of life. This integration of data classes resolved the basal topology of sea spiders with high support. The enigmatic family Austrodecidae was resolved as the sister group to the remaining Pycnogonida and the small-bodied family Rhynchothoracidae as the sister group of the robust-bodied family Pycnogonidae. Molecular divergence time estimation recovered a basal divergence of crown group sea spiders in the Ordovician. Comparison of diversification dynamics with other marine invertebrate taxa that originated in the Paleozoic suggests that sea spiders and some crustacean groups exhibit resilience to mass extinction episodes, relative to mollusk and echinoderm lineages.

Keywords: arthropods | Pycnogonida | mitogenome | ultraconserved | diversification

Introduction

Pycnogonida (sea spiders), the sister group to the remaining Chelicerata, are exclusively marine arthropods ranging from one to 750 mm in leg span (fig. 1). The morphological architecture of sea spiders is unusual, with a typically small body that is dwarfed by the much longer legs (hence, the alternate name “Pantopoda”, or “all legs”), into which diverticula of major organ systems emanate. Sea spiders are found throughout the world’s oceans from the intertidal zone to abyssal depths, but are especially abundant and diverse in polar benthic communities. In contrast to many invertebrate groups that flourish in the tropics, the peak of sea spider diversity is concentrated in the Southern Ocean, which also harbors multiple cases of gigantism in distantly related species (Arnaud and Bamber 1987; Griffiths et al. 2011).

Pycnogonids typically have four pairs of walking legs attached to the small body; the cephalon bears an anterior proboscis, composed of three longitudinal antimeres with the triradiate pharynx lumen in their center; and three pairs of cephalic appendages, the chelifores, palps, and ovigers.
Extant Pycnogonida lack a segmented opisthosoma (abdomen or posterior tagma), and thus also lack segmentally iterated opisthosomal respiratory organs that are found in other chelicerates. Sea spiders respire instead via cuticular gas exchange, with peristaltic contractions of the gut facilitating oxygen transport through the body (Woods et al. 2017; Lane et al. 2018). Ovigers, a type of modified leg unique to Pycnogonida, are used for grooming and by the males to carry egg masses (fig. 1M) (Arnaud and Bamber 1987). A remarkable exception from the conserved body architecture are genera with supernumerary body segments (resulting in 10-legged species), which occur in three families (Colossendeidae, Pycnogonidae, and Nymphonidae), and two genera (Dodecolopoda and Sexanymphon) are even characterized by 12 legs (Arango and Wheeler 2007). Beyond this, the cephalic appendages show generally a high degree of variation. Families are often distinguishable by the number of articles in the palps and ovigers (Arango 2002), and in some lineages, adults may lack one or more cephalic appendage types altogether.

Early conceptions of sea spider phylogeny envisioned a gradual reductive trend characterized by unidirectional, stepwise losses of appendage types (Hedgpeth 1947; Fry 1978; Munilla 1999). Phylogenetic investigations of sea spider relationships based on anatomical data (Arango 2002) or combined analyses of morphology and molecular sequence data (Arango and Wheeler 2007) suggested instead that reduction of appendages occurred independently across the phylogeny, but tree topologies were highly discordant between data partitions. Subsequent approaches to infer sea spider relationships under model-based approaches (Nakamura et al. 2007; Arabi et al. 2010; Sabroux et al. 2018) were repeatedly frustrated by the instability of basal relationships, which are attributable to two possible causes.

First, efforts to infer the phylogeny of Pycnogonida have been based on a small number of loci (one to six genes) that evolve at high rates (e.g., mitochondrial genes cytochrome c oxidase subunit I and 16S rRNA) or those that evolve at uninformatively low rates (e.g., nuclear ribosomal genes) (fig. 2) (Arango and Wheeler 2007; Nakamura et al. 2007; Arabi et al. 2010; Sabroux et al. 2018). Separately, mitochondrial genes of sea spiders exhibit well-known lineage-specific compositional biases (Arabi et al. 2010). Datasets based on fast-evolving mitochondrial genes have exhibited limited utility in resolving Paleozoic relationships of various invertebrate groups, and the placement of sea spiders within Chelicerata specifically (Rota-Stabelli et al. 2010; Masta et al. 2010).

Second, previous phylogenetic studies have poorly sampled (with respect to number of loci), or omitted entirely, two small-bodied families of sea spiders, Austrodecidae and Rynchothoracidae (Arabi et al. 2010; Sabroux et al. 2018). Austrodecidae (fig. 1G) are distinguished from other sea
spiders by the annulation of the proboscis. Little is known about their biology, as most austrodecids are small-bodied species (< 5 mm) that are infrequently encountered (Arnaud and Bamber 1987; Griffiths et al. 2011). Even less understood are species of Rhynchothoracidae (fig. 1H), which typically do not exceed one millimeter in length (Arnaud and Bamber 1987; Arango 2002).

The lack of a robust sea spider phylogeny has hindered inferences of major macroevolutionary trends in the group (Brenneis and Scholtz 2015; Brenneis et al. 2017, 2018). While phylotranscriptomic approaches have proven effective for resolving relationships of various chelicerate groups (Sharma et al. 2014; Santibáñez-López et al. 2019), the inaccessibility of rare sea spider lineages has obviated RNA-Seq-based approaches, as cryptic and small-bodied species are often not identified in ethanol-preserved samples until weeks to years after their initial collection. Moreover, sea spider specimens are often covered with epibionts and the sea spider digestive system extends into all but the two most distal podomeres of the legs. Thus, RNA-Seq-based approaches carry high risks of contaminations from epibionts and gut contents, especially for small-bodied species.

Resolving sea spider relationships thus requires an approach that can overcome several challenges to on-target data collection. To surmount these challenges, we undertook a target capture sequencing approach toward generating a robustly resolved phylogenetic backbone for Pycnogonida. We present here the first phylogenomic tree of sea spiders sampling all extant families. To assess whether whole genome duplications discovered in some chelicerate orders have affected orthology inference in sea spiders, we separately generated developmental transcriptomes of this group and investigated the incidence of Hox duplications across Pycnogonida. To place this branch of the Tree of Life in a temporal context, we inferred for the first time the age of the crown group Pycnogonida using a node dating approach under a Bayesian inference framework.

Results

**Phylogenomic relationships of Pycnogonida**

Maximum likelihood analyses of Matrices 1-4 consistently recovered the monophyly of Pycnogonida with maximal nodal support (bootstrap resampling frequency [BS]) (fig. 3). Austrodecidae was recovered in all analyses as the sister group to the remaining sea spiders with support (BS = 81-94%). At the next internal node, we recovered a clade comprised of Colossendeidae, Pycnogonidae, and Rhynchothoracidae (BS = 100%), with unambiguous support for the sister group relationship of the latter two families (BS = 96-100%). All other families were consistently recovered as a clade (BS = 85-100%).
As shown in fig. 3, ML analyses of Matrices 1-4 with model-fitting using ModelFinder recovered Endeidae and Phoxichiliidiidae as sister taxa (BS = 60-87%). Endeidae + Phoxichiliidiidae was in turn sister group (BS = 62-89%) to a clade comprised of the reciprocally monophyletic Pallenopsidae and Amмоtheidae (BS = 94-100%). The remaining sea spider lineages consisted of a monophyletic Nymphoniidae nested within Callipallenidae (BS = 80-100%), and this lineage in turn subtended with weak support by a grade comprised of Nymphoneella tapetis (BS = 53-70%) and a paraphyletic Ascorhynchidae sensu stricto (BS = 54-66%). Ascorhynchidae consistently included the putative amмоtheid genus Paranyphon (BS = 89-96%).

Model-fitting strategy did not affect the backbone tree topology, save for the placement of Nymphonella. Assigning a unique GTR + Γ model to each locus for Matrices 1-4 consistently recovered Nymphonella as the sister group of Endeidae. Matrix 4 alternatively recovered Nymphonella as nested within Ascorhynchidae when the partition models were developed using ModelFinder (fig. 3). Barring the placement of Nymphonella and its attendant topological instability, all analyses congruently resolved the monophyly of all families, excepting Rhynchothoracidae (one terminal only), Amмоtheidae (due to the placement of Paranyphon), and Callipallenidae (consistently recovered as paraphyletic with respect to Nymphoniidae). To assess support for non-monophyly of Callipallenidae, we performed AU tests, using a constraint tree that enforced the mutual monophyly of Callipallenidae and Nymphoniidae. The monophyly of Callipallenidae was rejected under both Matrix 1 (ΔlnL = 101.92, p = 1.67 × 10⁻³) and Matrix 4 (ΔlnL = 132.34, p = 3.27 × 10⁻⁴).

Most genera sampled with two or more taxa were also recovered as monophyletic. Exceptions included Decolopoda (a result attributable to the lack of any loci that include both Decolopoda species) and Achelia (sampled with three terminals). Our results corroborated the non-monophyly of Colossendeis due to the nested placement of species with supernumerary segments (e.g., the genera Decolopoda and Dodecolopoda; Dietz et al. 2019), and additionally reveal here the non-monophyly of Nymphon due to the nested placement of Boreonymphon and the 10-legged genus Pentanymphon.

Comparative assessment of phylogenetic data classes

Performance measures for phylogenetic data classes consisted of number of taxa sampled, alignment length, GC content (see Arabi et al. 2010), Robinson-Foulds distance (RF), weighted Robinson-Foulds distance (wRF), and evolutionary rate (mean percent pairwise sequence identity; MPSI). Properties of individual loci and matrices are separately provided in tables S4 and S5.
The mean number of taxa per locus was highest for the mitochondrial and nuclear ribosomal genes (68.2 and 85.7, respectively) and comparable for targeted exons, albeit with high variance (mean=50.9, \( \sigma^2=20.9 \)). UCE loci bore the most missing data, with an average of 22.2 terminals per locus. In addition, after trimming, the targeted exons and UCEs had comparable alignment lengths, and were shorter on average than mitochondrial and nuclear ribosomal genes. GC content of nuclear exons and UCEs was also comparable, with the average values intermediate between mitochondrial genes and nuclear ribosomal genes (fig. S1).

Assessment of congruence between gene trees and the pruned species trees showed that targeted exons and UCEs had comparable distributions of RF and wRF distances (fig. S2). Average values of both metrics were intermediate for targeted exons and UCEs, with respect to the lower values of mitochondrial genes and the higher values of nuclear ribosomal genes. Paralleling these distributions, MPSI values of targeted exons and UCEs were intermediate between mitochondrial genes and nuclear ribosomal genes (fig. S2).

Inference of tree topologies based on each of the four data partitions was performed in IQ-TREE with identical heuristics as for the species tree analyses. Comparison of tree topologies reflected the performance metrics measured above; mitochondrial genes and targeted exons retained many of the higher-level relationships recovered by the species tree, such as the monophyly of various families, and the sister group relationships of Callipallenidae + Nymphonidae and Pallenopsidae + Ammolethidae (fig. S3). However, by themselves, none of these partitions could resolve basal relationships with support. Reflecting the high proportion of missing data, the UCE tree topology was largely discordant with the species tree and other partitions, although some higher-level relationships were consistently recovered (e.g., the monophyly of some families; Callipallenidae + Nymphonidae; Pallenopsidae + Ammolethidae). The nuclear ribosomal partition retained some higher-level relationships, but branch lengths subtending interfamilial relationships were close to zero and were not supported (fig. S3).

Assessment of whole genome duplication and paralogy in Pycnogonida

It has recently been shown that whole genome duplications (WGD) have independently occurred in at least two different groups of Chelicerata, the horseshoe crabs (Kenny et al. 2016; Shingate et al. 2020) and the arachnopulmonates (e.g., spiders and scorpions; Sharma et al. 2014b; Schwager et al. 2017; Leite et al. 2018). Due to the unavailability of genomes sampling all chelicerate orders, it is presently unknown whether WGD has also affected Pycnogonida and whether putative orthologs of sea spiders may in fact be paralogous copies. To assess this
possibility, we generated the first developmental transcriptomes of five sea spider species, sampling four families. We surveyed these transcriptomes for the presence of Hox genes, which reliably exhibit duplicates in embryonic transcriptomes of WGD taxa (Sharma et al. 2014b; Gainett and Sharma 2020) and a subset of which has been shown to be expressed in canonical Hox-like domains in the protonymphon stage of sea spiders (Jager et al. 2006). Between six and eight Hox genes were recovered per species from the five sea spider transcriptomes, with gene tree analysis inferring all these to be single-copy (fig. 4). By comparison, an embryonic transcriptome of the mygalomorph spider Aphonopelma hentzi (an exemplar of Arachnopulmonata) that was sequenced in parallel with the embryonic sea spiders, revealed a complement of 18 Hox genes (two copies of all ten Hox genes except for Hox3 and fushi tarazu), consistent with the pattern across arachnopulmonates. Other genes surveyed for potential shared duplication included appendage patterning genes (e.g., dachshund, homothorax; Nolan et al. 2020) and members of the retinal determination gene network (e.g., sine oculis; Optix; Gainett et al. 2020); these were also found to be single-copy across all sea spiders (Dryad Digital Repository). Broadly, these patterns suggest that Pycnogonida is not part of the WGD events exhibited by Xiphosura and Arachnopulmonata.

As a separate approach to validating orthology, we annotated our multiple sequence alignments of targeted exons de novo by identifying the best BLAST hit to the proteome of the common fruit-fly (Drosophila melanogaster). All recovered annotations were consistent with the gene IDs of the single-copy orthologs intially selected for probe design (from Sharma et al. 2014a; Table S6). Taken together with the contents of sea spider embryonic transcriptomes, these results suggest that sea spiders are not included among the chelicerate orders that exhibit WGD and thus do not exhibit the same patterns of paralogy.

**Age and tempo of sea spider diversification**

Phylogenomic estimation of divergence times was performed using 11 outgroup and four ingroup fossil calibrations; fossil taxa and justification are provided in Supplementary Text S2. Dating was performed under a correlated rates clock model and the most complete matrix (Matrix 1) recovered a Late Cambrian to Early Silurian age for the crown group of Pycnogonida (median: 457 Mya; 95% HPD interval: 423-489 Mya) (fig. 5A). Comparable ages were recovered for the same dataset under an uncorrelated rates clock model (fig. S4). Diversification of sea spider families (with Nymphonidae considered as a derived lineage of Callipallenidae) was estimated to have occurred during the Paleozoic under both clock models. Separately, we reran molecular dating analyses under both clock models with two alternative calibration schemes: (1) all calibrations...
except for the fossil *Haliestes dasos* and (2) outgroup calibrations only. Results varied little between analyses (fig. S4). BAMM analysis of sea spider diversification revealed no evidence of rate shifts within Pycnogonida, recovering instead a monotonic and near-constant process of diversification (fig. S5).

**Discussion**

*A phylogenomic view of higher-level sea spider relationships*

Traditional views of sea spider evolution suggested that the sister group of the remaining sea spiders was Nymphonidae (Munila and de Haro 1981), based on its generalized appendage morphology (Hedgpeth 1947), or possibly Ammootheidae (previously thought to include *Eurycyde*; Stock 1994), based on anatomical similarities between ammootheids and fossils like *Palaeoisopus problematicus*. The species tree that we produced decisively recovered the enigmatic family Austrodecidae as the sister group to the remaining sea spiders. Earlier classifications had already pointed to the distinctness of Austrodecidae, segregating the family as sole member of the suborder Stiripasterida (Fry 1978; Bamber 2007). The recovery of a sister group relationship of Pycnogonidae and Rhynchothoracidae has been previously proposed based on morphology (Munilla 1999) and an analysis of 18S rRNA (Nakamura et al. 2007). Potential morphological apomorphies of Pycnogonidae and Rhynchothoracidae include the general habitus (compact body, relatively short legs), shape and configuration of the proboscis, presence of female gonopores on the last leg pair only, and the egg-carrying behavior of males (single pancake-shaped mass into which both ovigers insert). Overall, our dataset established a stable backbone topology for the remaining families (fig. 3). Only the position of *Nymphonella tapetis* engendered topological discordance across the datasets.

Establishing the sequence of basal branching families facilitates reconstruction of evolutionary transformation series for major character systems. Upon reconstructing adult appendage characteristics on our tree topology, we found that adult chelifores are absent in a grade of families at the base of extant Pycnogonida (fig. 6). Given some ambiguity in the state of the chelifore in some fossil taxa, as well as uncertainty about their phylogenetic placement, our results suggest that the presence of an adult chelifore may not be the unambiguous ancestral condition for crown-group sea spiders. In any case, a strictly gradual reduction of cephalic appendages (*sensu* Munilla 1999) has to be refuted for the crown-group, as the traditionally advocated early branching families Nymphonidae and Ammootheidae with full sets of adult appendages are resolved in derived positions (a result consistent with previous studies; Arango 2002; Arango and Wheeler 2007; Arabi...
The sister group relationship of Endeidae and Phoxichilidiidae accords with the shared loss of palps and female ovigers in these families. We also consistently obtained the non-monophyly of Callipallenidae, recapitulating a result that has been suggested, albeit with weak support, in previous analyses of one to six loci (Arango and Wheeler 2007; Nakamura et al. 2007; Arabi et al. 2010). In contrast to previous topologies, we recovered the monophyly of Nymphonidae as a derived lineage within the callipallenids across all datasets and analyses. This result calls for systematic revision of these two families and illustrates another instance of cephalic appendage regain instead of reduction during crown-group evolution. While functional chelifores are shared by both families, adult palps are lacking in the majority of callipallenid taxa (fig. 6), with the exception of few genera where they are present but largely reduced in males only (e.g., Anoropallene). By contrast, all nymphonids possess well-differentiated five-articled adult palps in both sexes (fig. 6; Brenneis et al. 2011). Moreover, this callipallenid-nymphonid relationship also suggests the re-emergence of the pycnogonid-typical protonymphon larva in the nymphonid lineage, having been lost in the more direct development of callipallenids (Brenneis et al. 2011a).

While genera were largely recovered as monophyletic, those defined to accommodate the condition of supernumerary segments (e.g., Decolopoda, Pentanymphon) were nested within larger genera of eight-legged species (e.g., Colossendeis, Nymphon). The exception was Pentapycnon, although our sampling of the family Pycnogonidae remains too sparse to test the monophyly of Pycnogonum (ca. 100 described species). Lineages with supernumerary segments are remarkable in an evolutionary context; in other Chelicerata, segment number tends to be fixed within a given extant order, as exemplified by such groups as scorpions and harvestmen. Changes in the segmentation of the prosoma (the appendage-bearing tagma of chelicerates) are especially rare (e.g., the synzipsorousines Offacolus kingi and Weinbergina opitzi; Dunlop and Lamsdell 2017), yet all sea spiders are distinguished from the remaining Chelicerata by the presence of an additional oviger-bearing segment (the post-tritocerebral segment, which bears a walking leg in all other chelicerates). The mechanistic basis of supernumerary segment addition is currently an enigma, due to limited developmental genetic resources for sea spiders (Jager et al. 2006; Brenneis et al. 2008). Future work should prioritize the gap segmentation genes of sea spiders, especially those functionally linked to gap segmentation phenotypes in chelicerate model species (Schwager et al. 2009; Pechmann et al. 2011; Setton and Sharma 2018).
Nymphonella and the composition of Ascorhynchidae

Surprisingly, the position of the putative ascorhynchid Nymphonella tapetis was unstable to analytical treatment of the dataset, a result that cannot be attributed to missing data alone (N. tapetis was represented in 47-59% of loci across Matrices 1-4). Nymphonella is clearly distinguished from other sea spiders by anterolaterally projecting chelifores, annulated distal podomeres of the first walking leg, and a bulbous proboscis (fig. 1F). Previous efforts to infer the placement of Nymphonella have recovered limited support for its placement or for the monophyly of Ascorhynchidae, though Nymphonella has only been represented by 18S rRNA in such studies (Sabroux et al. 2018; Miyazaki et al. 2015). Nymphonella may therefore constitute a rogue taxon, an inference corroborated by its variable placement across data partitions generated in this study (fig. S3).

By contrast, we recovered support for a nested placement of the putative ammothid genus Paranympphon [32] within Ascorhynchidae sensu stricto (represented by the genera Eurycyde and Ascorhynchus). Paranympphon has never been previously sequenced, but like Nymphonella, this terminal exhibited topological incongruence across data classes, being recovered within ammothids by the mitochondrial data partition (albeit being represented therein by only three mitochondrial genes), and with ascorhynchids by UCEs and exons (13 and 17 loci, respectively).

Given the uncertainty surrounding the composition of Ascorhynchidae, future systematic efforts should target deeper sequencing of the genera Nymphonella (three described species) and Paranympphon (six described species).

Combining data classes overcomes limitations of individual partitions

Historical efforts to infer higher-level relationships of sea spiders have leaned heavily on the Sanger-sequenced nuclear ribosomal and mitochondrial markers, but these have not yielded a stable sea spider phylogeny to date (Arango and Wheeler 2007; Arabi et al. 2010; Sabroux et al. 2018). Ideally, new markers should exhibit evolutionary rates intermediate between slow-evolving nuclear ribosomal genes and fast-evolving mitochondrial markers. Targeted exons and UCEs sequenced in this study exhibit rates that propitiously fall exactly within this desired range (fig. S2).

Problematically for the UCE dataset, capture efficiency of UCE probes was much lower than other data partitions (fig. S1). This phenomenon appears to be partly linked amount and quality of genomic DNA in a given extraction, as reported in a previous study on spiders with the same UCE probe set, resulting in low recovery rates for extractions with small quantities of DNA (Wood et al. 2018). Low quantity of DNA is unavoidable for such minute and rare lineages as Rhynchothorax and
austrodecids. However, coverage inefficiency of the arachnid UCE probe set was systemic for sea spiders, even when ample DNA was available for large-bodied species, suggesting that the Arachnida 1.1Kv1 probe set is not universally effective across chelicerate orders (e.g., a single mite exemplar in Starrett et al. 2017 similarly exhibited low yields of UCE loci, by comparison to other arachnid taxa). As a result, of an initial 230 UCEs amplified, we discarded 56% of alignments where fewer than six sea spider terminals were obtained. The remaining 101 UCE loci bore high proportions of missing data, in contrast to the targeted exons. Similar outcomes have been reported for palpimanoid spiders, a group that had the advantage of a well-annotated theridiid spider genome for validation of probe design (Wood et al. 2018).

Nevertheless, we observed high informativeness of the UCE loci despite missing data, as inferred from distributions of RF and weighted RF distances. The concatenated UCE tree, while incongruent with the species tree, recovered such higher-level groupings as Nymphonidae nested within Callipallenidae, and Colossendeidae sister group to Rhynchothoracidae + Pycnogonidae. Given the promise of this phylogenetic data class, efforts to improve the recovery of UCE datasets in sea spiders should target the generation of high-quality sea spider genomes, with downstream improvements in the design of sea spider-specific UCE probes. Such strategies have been shown to overcome limitations inherent to the arachnid UCE bait set for spiders (Kulkarni et al. 2019).

The first molecular dating of Pycnogonida reveals ancient diversification and monotonic evolutionary rates

Generally, the fossil record for many invertebrate groups (especially marine arthropods) is scarce. The appearance of a Cambrian fossil resembling a sea spider early developmental instar, together with the exquisite preservation of the fossil Haliestes dasos, clearly point to an ancient origin of sea spiders before the Silurian (Supplementary Text S1). However, a fossil record dating to the Paleozoic is not dispositive of ancient diversification of the crown group. Evolutionary relicts like Xiphosura (horseshoe crabs) and Nautiloidea appeared early in the fossil record (oldest fossils belonging to the Ordovician for Xiphosura; Late Cambrian for Nautiloidea), but survived to the present as a small number of species that diverged relatively recently (Obst et al. 2012; Combosch et al. 2017). Molecular divergence time estimation for such groups as Crinoidea (feather stars) and Ophiuroidea (brittle stars) have shown that both these echinoderm groups diversified in the wake of the end-Permian mass extinction (Rouse et al. 2013; O’Hara et al. 2014), having survived this extinction episode as a single lineage that subsequently recovered some fraction of its diversity (a “revenant” taxon sensu Sharma and Wheeler 2013). By contrast, marine invertebrate groups like
Holothuroidea (sea cucumbers) (Miller et al. 2017) and Protobranchia (protobranch bivalves) (Sharma et al. 2013) diversified in the Paleozoic but retain the signature of the end-Permian mass extinction in their dated phylogenies, which manifests as a low diversification rate (the plateau of an anti-sigmoidal curve in a log-lineage through time plot) until the beginning of the Mesozoic (fig. 5B) (Crisp and Cook 2005; Sharma et al. 2013).

To date, molecular divergence time estimation has never been performed for sea spiders. It was previously postulated that sea spiders diversified relatively recently (in the Mesozoic), but this speculation was based solely on the branch length (i.e., substitutions per site) subtending Pycnogonida in a molecular phylogeny, rather than a parametric molecular dating approach or the use of fossil calibrations (Arabi et al. 2010). Our divergence time estimation unambiguously recovered Cambrian to Ordovician age of sea spider diversification, a result that is independent of the choice of clock model or use of ingroup fossil calibrations. Ancient diversification of Pycnogonida during the Ordovician is consistent with their fossil record (e.g., Jurassic sea spiders that are assignable to families; Supplementary Text S1), and further suggests that Devonian sea spiders with opisthosomal segments (e.g., Flagellopantopus, Palaeoisopus) constitute stem lineages that diverged from extant Pycnogonida before the Ordovician and thereafter went extinct. A parallel evolutionary history has been reconstructed for spiders and their extinct sister group Uraraneida, which was recently shown to have survived at least until the Cretaceous (Wang et al. 2018; Huang et al. 2018).

Comparison of lineage accumulation through time for selected marine invertebrate groups reveals the marked difference between the evolutionary history of sea spiders and other Paleozoic fauna (fig. 5B) (Obst et al. 2012; Lins et al. 2012; Korn et al. 2013; Rouse et al. 2013; Sharma et al. 2013; O’Hara et al. 2014; Miller et al. 2017; Wolfe et al. 2019). In contrast to groups like Protobranchia and Crinoidea, Pycnogonida exhibited a static diversification regime, with a monotonic process of slowing diversification rate since initial divergence, and no evidence of rate shifts (fig. 5, S6), by comparison to other Paleozoic taxa. Notably, the use of log-lineage through time plots as the sole window for examining evolutionary history has been recently criticized for non-identifiability; it has been shown mathematically that myriad combinations of cladogenetic and extinction rate regimes can engender the same net diversification rate curves (Louca and Pennell 2020). Fossil datasets, when available and densely sampled, were proposed as more reliable arbiters of the accuracy of log-lineage through time trajectories, and thus potential solutions to this impasse (Louca and Pennell 2020). Fortuitously, such direct comparisons between net diversification curves and the fossil record are available for a subset of taxa that exhibit a detailed
fossil record (fig. 5B). Specifically, dynamics of generic diversity of fossil echinoderms substantiate the signature of susceptibility to the end-Permian extinction event in dated phylogenies, with coincident accumulation of lineages in extant phylogenies and the fossil record in the wake of the mass extinction episodes (Figure 3 of Rouse et al. 2013; Figure 5 of Miller et al. 2017). Taken together with the absence of inferred rate shifts for sea spider in BAMM analyses (fig. S6), our results indicate the absence of evidence for radical changes in diversification dynamics of Pycnogonida during most of the Phanerozoic.

The lack of any signature of major mass extinction events in the early (pre-Mesozoic) evolutionary history of extant sea spider and crustacean lineages is unexpected. In contrast to groups like bivalves and echinoderms, sea spiders do not form large calcified hard parts (valves or tests) whose deposition was severely affected by cataclysmic historical environmental changes (e.g., during the end-Permian event) (Krishnan 1955). However, this differing composition of hard parts does not account for the Paleozoic origin and post-Permian diversification of aquatic arthropod groups like Xiphosura and Notostraca (Obst et al. 2012; Korn et al. 2013). Moreover, the cuticle of decapods is indeed biomineralized, with denser cuticle associated with higher calcification in benthic Decapoda (Amato et al. 2008). Apropos, the log-lineage through time trajectory for Decapoda does indeed exhibit a small decline in lineage accumulation rate immediately preceding the Triassic (fig. 5B), but this result was only observed under divergence time estimation under one of two clock models (Wolfe et al. 2019).

We therefore postulate that the inferred resilience of sea spiders is not exclusively due to their lack of a calcified exoskeleton. Patterns of sea spider species richness observed today could be attributable to higher diversification rates in cooler regions, as has previously been shown in other marine arthropod groups like Anomura (Bracken-Grissom et al. 2013; Davis et al. 2016). In deep-sea isopods, diversification in the deep sea occurred in parallel with anoxic events, with the earliest radiation dated to the early Permian and subsequent episodes of rapid colonization and radiation (Lins et al. 2012). Such processes could partly explain the oddities of sea spider biogeography (such as the concentration of their diversity in Antarctica), but the role that the Southern Ocean has played as a potential center of endemism is not clear based on our results.

In the present study, which is focused on higher-level phylogenetic relationships, we lack sufficiently broad geographic sampling to assess the biogeography of sea spiders and a putative role for the polar regions as the driver of sea spider diversity in lower latitudes. Population genomic works have begun tackling such questions at shallower taxonomic scales (within species or species groups), with high levels of precision (Dietz et al. 2015, 2019; Dömel et al. 2015, 2019;
Harder et al. 2016; Soler-Membrives et al. 2017; Collins et al. 2018), in tandem with challenging comparative physiological experiments in Southern Ocean species that address the dynamic evolution of body size in Pycnogonida (Woods et al. 2009; Shishido et al. 2019). Denser sampling of extant sea spider diversity in future studies will augment the predictive power of such approaches in deciphering the evolutionary history of this ancient group.

**Materials and Methods**

*Species sampling and locus selection*

Specimens of all sea spider families were obtained from field expeditions and museum collections, as well as material collected during multiple deep sea cruises. A list of taxa sampled is provided in table S1. DNA was extracted using the Qiagen DNeasy Blood and Tissue Kit (Valencia, CA, US) from one to five specimens, prioritizing tissue from the propodus and main claw (gut-free leg podomeres) where possible. Taxonomic sampling consisted of 89 sea spiders; outgroups consisted of 14 Arachnida (including one Xiphosura, which has recently been shown to be possibly nested within the arachnids; Ballesteros and Sharma 2019; Ballesteros et al. 2019), three Myriapoda, three Pancrustacea, and one Onychophora.

Probes were designed using available references and databases, given the absence of a published sea spider genome, following the pipeline established by Breinholt et al. (2018). Probes for mitochondrial genomes were designed using a newly-sequenced mitogenome of *Pycnogonum litorale* and five published sea spider mitogenomes: *Tanystylum orbiculare* (GU370074.1), *Ammothea hilgendorfi* (GU370075.1), *Achelia bituberculata* (NC_009724.1), *Nymphon unguiculatum-charcoti* complex sp. (GU370076.1), and *Nymphon gracile* (NC_008572.1). From previous work on the phylogeny of Chelicerata (Sharma et al. 2014a), we identified 56 nuclear exons that (1) occurred in single copy across Panarthropoda, (2) were present for at least one of two sea spider Illumina transcriptomes used as templates for probe design (*Anoplodactylus insignis* and *Pycnogonum litorale*; Ballesteros and Sharma 2019); and (3) exhibited sufficient conservation of intron-exon boundaries in the genomes of arthropods for 125-bp probe design with 3 × tiling. Probes design was guided by conservation of intron-exon boundaries in the genomes of *Drosophila melanogaster*, *Strigamia maritima*, and *Parasteatoda tepidariorum*. For ultraconserved element (UCE) sequencing, we deployed probes from the UCE Arachnida 1.1Kv1 bait set (Starrett et al. 2017). Probe design, synthesis, automated library preparation, and paired-end sequencing (2 × 150 bp) on the Illumina Hi-Seq 2500 platform were performed through RAPiD Genomics (Gainesville,
Sequencing efficiency is provided in table S2; details of locus selection and orthology validation are provided in Supplementary Text S1. All probe sequences are made available on the Dryad Digital Repository.

**Bioinformatic and phylogenomic operations**

Raw reads were assembled using Trinity v.2.7 (Grabherr et al. 2011; Haas et al. 2013) using a path reinforcement distance of 75. Reference sequences of sea spiders (mitogenomes and nuclear exons) and/or euchelicerates (UCEs) were used to separate loci into alignments. We included the nuclear ribosomal genes 5.8S rRNA, 18S rRNA, and 28S rRNA, which were obtained as bycatch; on-target amplification of these bycatch loci was verified using BLASTn against available sea spider ribosomal sequences in GenBank. Outgroup taxa were subsequently added into the alignments using available transcriptomes from our previous works (for nuclear exons; Sharma et al. 2014a), from GenBank-accessioned data (mitochondrial genes/genomes and nuclear ribosomal genes), and from a subset of the taxa in chelicerate UCE alignments (Starrett et al. 2018). Due to the unavailability of the same phylogenetic data classes among the genomic resources of some data-poor outgroups, we created chimeric terminals for such terminals as _Eremobates_ and _Ricinoides_. The nature of these congeneric chimeras is that sequence data for nuclear exons and ribosomal genes are drawn from the transcriptomes of one species (e.g., _Eremobates cf. transfugoides_; _Ricinoides atewa_), but the mitochondrial genome is drawn from a congener (e.g., _Eremobates cf. palpisetulosus_; _Ricinoides karschii_). The list of chimeric outgroup terminals is detailed in table S2.

Multiple sequence alignment was performed using MUSCLE v.3.8.31 (Edgar 2004). For ribosomal genes, alignments were performed on nucleotide sequences. For nuclear and mitochondrial exons, alignments were performed using peptide translations and subsequently reverted to nucleotides. We discarded the mitochondrial genes 12S rRNA, 16S rRNA, and tRNAs, due to their high rate of evolution, as well as off-target capture of non-sea spider sequence for these regions, which we presumed to have resulted from sequencing of gut content and/or epibionts.

Alignments were trimmed using trimal v.1.2 (Capella-Gutiérrez et al. 2009) to cull overhanging ends. Each alignment was filtered to ensure the inclusion of a minimum of six sea spider terminals and four outgroup terminals (resulting in the exclusion of the mitochondrial protein ATP8 and 129 out of 230 UCE alignments). Gene trees were inferred using IQ-TREE v.1.6 (Nguyen et al. 2014) with the best-fitting model selected by ModelFinder (-m MFP) (Kalyaanamoorthy et al. 2017). Gene tree topologies were visually inspected for potential paralogs, which were summarily discarded from alignments (principally found in the UCE alignments previously generated for arachnids).
Orthology was validated using BLAST-based annotation; the annotations for all sequenced loci is provided in.

Four matrices were constructed for our main analyses and all datasets were treated as nucleotide alignments. Matrices 1-3 were constructed using taxon occupancy thresholds of 50%, 33%, and 25%, respectively. Matrix 4 consisted of loci stipulated to sample at least one Austrodecidae (the putative sister group to the remaining sea spiders). The final alignments were analyzed as concatenated supermatrices using IQ-TREE v.1.6 with best-fitting models per locus. Due to ongoing debate over the benefits of model-fitting, we additionally analyzed our datasets using a unique GTR + $\Gamma_4$ model for each locus as well. Nodal support was estimated using bootstrap resampling frequency with 1000 ultrafast bootstrap replicates in IQ-TREE. In addition, Approximately Unbiased (AU) tests of monophyly were performed using in-built tools in IQ-TREE for selected phylogenetic hypotheses using Matrices 1 and 4.

For assessment of dataset performance, we inferred the normalized Robinson-Foulds (RF) and weighted Robinson-Foulds (wRF) distances between the species tree inferred using the most complete matrix (Matrix 1) and each gene tree. To overcome missing data, the species tree was recursively pruned of tips to match the terminals of each gene tree prior to calculation of RF and wRF distances, and the distances were normalized using the maximal possible distance between each gene tree and the pruned species tree. Other metrics of performance consisted of locus length, taxon sampling, GC content, and mean pairwise sequence identity (MPSI; a proxy for evolutionary rate). Tree manipulations and calculations of performance metrics were performed using R packages ape v.5.3 (Paradis and Schliep 2019) and phytools v.0.6-99 (Revell 2012).

Bayesian analyses were trialed using PhyloBayes-mpi v.1.7 (Lartillot and Philippe 2004), but these analyses failed to reach convergence after several months of computation and were ultimately discarded.

**Embryonic transcriptomes**

Ovigerous spider males with prominent egg masses were hand collected from field sites between 2015-2017 for five sea spider species: *Phoxichilidium cf. femoratum* (Phoxichilidiidae), *Tanystylum orbiculare* (Ammotheidae), *Pallenella flava* (Callipallenidae), *Stylopallene cheilorhynchus* (Callipallenidae), and *Nymphon molleri* (Nymphonidae). Egg masses were separated from the males with fine forceps and immediately transferred to RNAlater (ThermoFisher). Due to asynchrony of development within egg masses, these samples represented a range of embryonic stages, from early embryogenesis through protonymphon larvae. As a control sample, we included
a range of embryonic stages from the tarantula *Aphonopelma hentzi*; field collection and processing of embryos of this spider are described in Setton et al. (2019). All samples were removed from RNAlater and transferred to TRIZOL TriReagent for RNA extraction, following our previous procedures (Sharma et al. 2014a). Library preparation, paired end Illumina sequencing, and transcriptome assembly with Trinity also followed our previous approach (Sharma et al. 2014a).

We used the conserved homeodomain sequences of all Hox genes of the spider *Parasteatoda tepidariorum* to retrieve Hox sequences from new libraries, using tBLASTn searches. Putative *A. hentzi* and sea spider sequences Hox homologs were added to multiple sequence alignments recently generated by us (Gainett and Sharma 2020), in addition to all available sea spider Hox sequences previously generated using RACE PCR (Manuel et al. 2006). Gene tree inference was performed using IQ-TREE, treating sequence data as peptides and model-fitting using ModelFinder. The Hox gene tree is shown in fig. S4; the multiple sequence alignment used for Hox orthology inference is provided on the Dryad Digital Repository.

**Phylogenomic dating**

Phylogenomic estimation of divergence times was estimated using a node dating approach with MCMCTree (Yang 2007) on two datasets (Matrices 1 and 3), implementing a likelihood approximation of branch lengths using a multivariate normal distribution (dos Reis and Yang 2019). Fossils used to inform the dating consisted of 11 outgroup and four ingroup node calibrations. All calibrations were implemented as soft minimum and soft maximum ages.

The oldest unequivocal fossil of crown group Pycnogonida, *Haliestes dasos*, dates to the Silurian (424 Mya) (Siveter et al. 2004), whereas the oldest fossil assigned to the pycnogonid (stem) lineage is an early developmental instar from the Upper Cambrian (*Cambropycnogon klausmuelleri*) (Waloszek and Dunlop 2002). *Haliestes dasos* superficially resembles Nymphonidae, clearly lacks an annulated or segmented proboscis, and has been recovered at the base of Pycnogonida in some morphological cladistic analyses (Siveter et al. 2004) As a conservative measure, we calibrated the age of crown group Pycnogonida using a soft minimum age of 424 Mya and a soft maximum age of 501 Mya. Various other fossils (*Flagellopantopus blocki, Palaeothea devonica, Palaeoisopus problematicus, Palaeopantopus maucherii, Palaeomarachne granulata, Pentapantopus vogteli*), some of which exhibit morphologies attributable to stem-group lineages (e.g., a flagellum; opisthosomal segments), were not considered usable for node dating, due to poor preservation and/or an age younger than *Haliestes dasos.*
Sea spider crown group fossils from the Late Jurassic and La-Voulte-sur-Rhône were used to calibrate ingroup nodes: *Palaeopycnogonides gracilis* (Ammotheidae), *Colossopantopodus boissinensis* and *Colossopantopodus nanus* (Colossendidae), and *Palaeoendeis elmii* (Endeidae) (Charbonnier et al. 2007; Sabroux et al. 2019). Each of these fossils was treated as a minimum age calibration for the corresponding family, with a maximum soft bound age of 501 Mya. We abstained from using *Palaeothea devonica*, *Pentapantopus vogteli* and *?Eurycyde golem* in this study, as the poor preservation and lack of characteristic details of these likely crown group fossils preclude reliable placement in any extant genus or family.

For Arachnida, we constrained the crown group of Opiliones using the soft minimum age of 411 Mya, based on the harvestman fossil *Eophalangium sheari*, and the divergence of Eupnoi harvestmen with a soft minimum age of 305 Mya, based on the age of *Macrogyion cronus* (Garwood et al. 2011). The divergence of spiders from the remaining Tetrapulmonata was constrained to at least 386 Mya (age of *Attercopus fimbriunguis*) and the divergence of *Liphistius* at 305 Mya (age of *Palaeothele montceauensis*) (Selden et al. 2008; Wang et al. 2018). Pedipalpi (Amblypygi + Uropygi) was set to a minimum age of 319 Mya, based on the age of the fossil *Parageralinura naufraga* (Tetlie and Dunlop 2008). The divergence of Xiphosura from Rinculei was constrained to a minimum of 445 Mya (age of *Lunataspis aurora*) (Rudkin et al. 2008). The stem-group age of scorpions was constrained to a soft minimum age of 435 Mya and a soft maximum of 514 Mya (based on the ages of *Parioscorpio venator* and *Eramoscorpius brucensis*) (Waddington et al. 2015; Wendruff et al. 2020). The crown group age of scorpions was constrained between 112.6 and 313.7 Mya (based on the ages of *Protoischnurus axelrodurum* and *Compsoscorpius buthiformis*) (Dunlop et al. 2016).

For Mandibulata, we constrained the basal diversification of Chilopoda between 416 and 521 Mya, and of Pleurostigmophora between 382.7 and 521 Mya. Altocrustacea (the node comprising the most recent common ancestor of the malacostracan exemplar *Parhyale hawaiensis* and the hexapods in this analysis) was constrained between 429.8 and 521 Mya. The root of the tree (split of Onychophora and Arthropoda) was constrained between 550 and 636 Mya. Mandibulate and root calibrations are based on various fossils reviewed by Wolfe et al. (2016).

Both the independent rates and correlated rates clock models were used to infer node ages with Matrix 1 (the most complete matrix), under the maximum likelihood tree topology inferred for this matrix with model selection using ModelFinder. We constrained the monophyly of Mandibulata and Phalangida (the non-Cyphophthalmi Opiliones) for node calibration. Four independent chains were run for 10⁶ generations for each analysis, sampling every 100⁶th generation. Convergence diagnostics were assessed using Tracer v.1.7 (Rambaut et al. 2018) and inbuilt tools in MCMCTree, ensuring...
minimum effective sample sizes >200 for summary statistics. As a conservative measure, 1000 trees (10%) were discarded as burnin. Separately, we ran two other dating analyses to assess the impact of sea spider fossils as calibration points: (1) using all fossil calibration points except for *Halieses dasos*; and (2) using only outgroup calibration points.

Analyses of diversification rates through time were performed using Bayesian Analysis of Macroevolutionary Mixtures (BAMM v.2.5.0) (Rabosky 2014). The chronogram inferred using all calibration points and the correlated rates clock model was used as the input for BAMM analysis. Outgroups were removed prior to analysis using phytools. Priors were iteratively tested for the expected number of rate shifts (one to five). Rate shifts were permitted on all branches. Sampling frequency was set to 6.54%, using the number of described species as a proxy for extant diversity. As an exploration of the effect of this parameter, we additionally reran the analysis with sampling frequency set to 1.308% and 0.654% (i.e., inferring total extant diversity to equal five times or ten times the current number of described species). MCMC chains were run for 500 000 generations, sampling event data every 10 000 generations, and discarding 25% of the run as burnin.

**Micro-computed tomography**

Fixed specimens were dehydrated via an ascending ethanol series, incubated in solution of 2% iodine (resublimated; Carl Roth GmbH&Co. KG, Karlsruhe, Germany; cat. #X864.1) in 99.5% ethanol for ca. 48 hours at ambient temperature, briefly rinsed in 99.5% ethanol (3-4x 10 min) and either transferred into a vial in ethanol (“wet scan”) or critical point-dried with a Leica EM CPD300 and glued with the posterior body pole on plastic welding rods (“dry scan”). Scans were performed with an Xradia MicroXCT-200 (Carl Zeiss Microscopy GmbH). Scan settings were individually optimized for each specimen, including objective choice (0.39×, 4×, 10x) according to size. Scans were performed under 40kV/200µA/8W, exposure times ranged from 0.4-1.5 sec. Tomography projections were reconstructed using the XMRsoftware (Carl Zeiss Microscopy GmbH) with TIFF format image stacks as output. All scans were performed with Binning 2 to reduce noise and subsequently reconstructed with Binning 1 (= full resolution) to avoid information loss. Processing and 3D visualization of the image stacks (including highlighting of cephalic appendages and removal of non-target structures) was performed with Imaris (version 7.0.0., Bitplane AG, Switzerland) as described previously (Scholtz and Brenneis 2016).

**Author contributions.** C.P.A, W.C.W., and P.P.S. conceived of the study. Specimens were collected and/or contributed by C.P.A., G.B., S.B., E.C.S, G.F.D., M.P.E., G.C., L.M., A.L.M. R.M., G.G., H.A.W., P.L.G.,
and P.P.S. A new mitogenome of *P. litorale* was contributed by G.S. for bait design. Coordination of sample sorting from the DIVA3 and IceAGE cruises was performed at the DZMB in Hamburg by S.B. Taxonomic identification from IceAGE and DIVA3 collections was performed by C.P.A, G.B., E.V.W.S., and P.P.S. E.V.W.S. and P.P.S performed molecular work. E.V.W.S. performed imaging of specimens used for extraction. G.B. performed μCT scans and their subsequent processing and visualization. J.A.B., C.E.S.L., M.D., S.M., J.S., C.W., and P.P.S performed bioinformatic and phylogenetic analyses. K.F.C., J.T.Z., E.V.W.S., and P.P.S. analyzed embryonic transcriptome data. W.C.W. and P.P.S funded the work. P.P.S drafted the manuscript, and all authors edited and approved the final content.

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**Data availability.** The complete dataset, including sequence alignments, annotated tree files, chronograms, and embryonic transcriptomic assemblies, have been deposited on Dryad Digital Repository under URL

https://datadryad.org/stash/share/HsFKzSdD00rwa5BQSEf9km6dXuURuiAt0wv0w0QVDUG. Raw read data were deposited in NCBI Sequence Read Archive. Loci from targeted sequencing were deposited in GenBank.

**Permitting.** Specimens from multiple deep sea cruises were collected under the auspices of the United States Antarctic Program, in compliance with the US Antarctic Conservation Act. All diplomatic formulars for sampling the Icelandic economic zone were given for cruises M85/3 (IceAGE1) and POS456 (IceAGE2). Specimens from Australia were collected under permit no. 15111 of the Tasmanian Department of Primary Industries, Parks, Water and Environment and Marine Parks permit G08/27858.1.

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Figure and Table Legends

**Figure 1.** Exemplars of sea spider diversity. (a) *Pallenella harrisii* (Callipallenidae). (b) *Nymphon grossipes* (Nymphonidae). (c) *Rhopalorhynchus magdaleneae* (Colossendeidae). (d) Copulating pair of *Pycnogonum litorale* (Pycnogonidae) with UV illumination. (e) *Stylopallene* sp. (Callipallenidae), photograph by Iain Gray. (f) *Nymphonella tapetis* (Ascorhynchidae sensu lato). (g) *Austrodecus glaciale* (Austrodecidae). (h) *Rhynchothorax australis* (Rhynchothoracidae). (i) *Anoplodactylus evansi* (Phoxichilidiidae). (j) *Cilunculus armatus* (Ammotheidae), (k) *Decolopoda australis* (Colossendeidae), photograph by Andrei Utevsky. (l) *Colossendeis megalonyx* (Colossendeidae). (m) Male of *Pallenella* sp. (Callipallenidae) with egg clutch.

**Figure 2.** Historical hypotheses of higher-level sea spider relationships based on molecular sequence data. Nodes lacking nodal support (<70% bootstrap support; <0.90 posterior probability) or conflicting across analyses have been collapsed. Brackets correspond to paraphyletic lineages. In Sabroux et al. (2018), both Nymphonidae and Callipallenidae were recovered as polyphyletic. Abbreviations: 12S, 12S rRNA; 16S, 16S rRNA; 18S, 18S rRNA; 28S, 28S rRNA; COI, cytochrome c oxidase subunit I; H3, histone H3.

**Figure 3.** Phylogenomic relationships of Pycnogonida based on maximum likelihood analysis of Matrix 3 (lnL = -472525.59). Colors of branches correspond to families (right). Numbers on nodes indicate bootstrap resampling frequencies for Matrices 1-4 with model-fitting using ModelFinder. Quotation marks on Ascorhynchidae reflect the inclusion of the putative ammotheid genus *Paranyphon*. Bottom left: Sensitivity plot indicating design of matrices and phylogenomic analyses. Inset (gray background): Alternative placements of *Nymphonella tapetis*.

**Figure 4.** Summary of Hox gene complements of surveyed taxa. Boldface text indicates new embryonic transcriptomes generated for this study. Hox genes for *Carcinoscorpius rotundicauda* are drawn from Shingate et al. (2020). Hox genes for *Endeis spinosa* and *Nymphon gracile* were reported by Manuel et al. (2006) using RACE-PCR. Phylogenetic relationships within Chelicerata are based on Ballesteros and Sharma (2019) and this study. Multiple sequence alignment, gene tree topology, and transcriptomic assemblies are provided in the Dryad Digital Repository.

**Figure 5.** (a) Phylogenomic dating of sea spiders based on the most complete data matrix and a correlated rates molecular clock model. Colors of branches and 95% HPD intervals correspond to families, as in figure 2. Line drawings (inset) correspond to stem-group
fossils *Palaeoisopus problematicus* (top) and *Flagellopantopus blocki* (bottom). 

(b) Log lineage through time trajectories for selected Paleozoic aquatic taxa (sources in text). Branching times are truncated at the Cenozoic to mitigate under-sampling of recent diversity and/or oversampling of intraspecific terminals.

**Figure 6.** Cephalic appendage evolution in sea spiders, with emphasis on the chelifore.

Reconstruction of adult chelifores is mapped on the topology obtained herein, with addition of fossil stem group and crown group representatives. Ancestral state reconstruction is based on equally weighted parsimony. Note the omission of functional adult chelifores in colossendeids with supernumerary segments (a derived state within the genus *Colossendeis*). Specimens in counter-clockwise order from top left: male *Austrodecus glaciale*, male *Endeis spinosa*, female *Phoxichilidium femoratum*, egg-bearing male *Ascorhynchus ramipes*, male *Stylopallene cheilorhynchus*, male *Nymphon gracile*, female *Ammothella longipes*, female *Colossendeis angusta*, female *Pycnogonum litorale*, female *Rhynchothorax australis*.

**Figure S1.** Comparative metrics of dataset performance by partition. Left column: Taxon sampling per locus. Middle column: Alignment length post-trimming. Right column: GC content.

**Figure S2.** Comparative metrics of dataset performance by partition. Left column: normalized Robinson-Foulds distance per locus (from species tree). Middle column: normalized weighted Robinson-Foulds distance per locus (from species tree). Right column: mean pairwise sequence identity (a proxy for evolutionary rate).

**Figure S3.** Maximum likelihood supermatrix analyses of individual data partitions. Colors in branches correspond to families (left). Annotated tree files with nodal support frequencies are available on the Dryad Digital Repository.

**Figure S4.** Chronograms of sea spiders under two different clock models and three different calibration schema, as indicated. Annotated tree files are available on the Dryad Digital Repository.

**Figure S5.** BAMM analysis of sea spiders under three different parameters for taxon sampling frequency. Left: sampling frequency set to 6.54% (i.e., extant diversity assumed to equal number of described species). Middle: sampling frequency set to 1.308% (i.e., extant diversity assumed to equal five times the number of described species). Right: sampling frequency set to 0.654% (i.e., extant diversity assumed to equal ten times the number of described species). Note the lack of rate shifts inferred under all three runs (prior on rate shifts set to 1).
Table S1. Locality and accession data for study specimens.

Table S2. Read counts per terminal.

Table S3. Accession data for sequences.

Table S4. Number of taxa, length, and missing data per locus.

Table S5. Number of taxa, number of loci, alignment length, and missing data per matrix.

Table S6. Validation of locus orthology.
Figure 2

Arango & Wheeler (2007)
Parsimony
59 taxa + 5 fossils
morph. and 6 loci: 18S, 28S, COI, 12S, 16S, H3

Nakamura et al. (2007)
ML, Bayesian inference
59 taxa
one locus: 18S

Arabi et al. (2010)
ML, Bayesian inference
35 taxa
4 loci: 18S, 28S, COI, 16S

Sabroux et al. (2018)
ML, Bayesian inference
135 taxa
2 loci: 18S, COI
Figure 5
