Article

Structural Features and Phylogenetic Implications of Crinoid Echinoderms Based on Thirteen Novel Mitochondrial Genomes

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Abstract: Crinoids, as integral echinoderms, play a crucial ecological role in benthic communities, serving as significant indicators reflecting the health of marine ecosystems. However, the phylogenetic relationships within crinoids are unclear. More molecular data can help to facilitate biodiversity assessment and elucidate evolutionary relationships by the phylogenetic tree. In this study, 13 complete mitochondrial genomes of the Crinoidea class were sequenced, annotated, and compared with other same class species available on NCBI. The results reveal five different gene order patterns among these mitochondrial genomes, indicating that crinoids have undergone gene rearrangements during evolution. The complete mitochondrial genome length of crinoids ranges from 15,772 bp to 16,850 bp. High A + T content, ranging from 64.5% to 74.2%, was observed. Additionally, our analysis of protein-coding genes highlights a preference for A + T nucleotides, along with specific start and stop codon usage, offering insights into codon bias and its implications for protein synthesis and function. The phylogenetic topology shows that the stalkless crinoid and stalked crinoid are distinct, and the phylogenetic trees generated based on maximum likelihood and Bayesian inference are almost identical at the family and order topology levels. The phylogenetic relationships of each family were fully clarified in four orders. A total of eleven positive selection sites were detected within six genes: cytb, nad2, nad3, nad4, nad4L and nad5. This study reveals the phylogenetic relationships of crinoid species, the mitochondrial gene differences, and the selective pressure on the evolution of stalked crinoids. This study significantly enhanced the crinoid mitochondrial genome database and contributed to a better understanding of the phylogenetic relationships among crinoid echinoderms.

Keywords: crinoids; echinoderm; mitochondrial genome; mitochondrial structure character

1. Introduction

Crinoids have been present in oceanic ecosystems for more than 500 million years, displaying a remarkable range of diversity across various marine environments, from tropical to polar seas and from intertidal zones to the depths of the ocean [1,2]. Based on their attachment type, crinoids are categorized into two main types: stalked crinoids and stalkless crinoids. Stalked crinoids [3] have a stalk throughout life and adopt a fixed way of life. Their body shape resembles a plant with three distinct sections: root, stem, and crown. They primarily inhabit the deep seas. Stalkless crinoids [4] (order: Comatulida) are stationary as adults but can freely move across the seafloor. Their varied shapes reflect
adaptations to different habitats and niches. Most of them inhabit coastal shallow sea rocks or hard bottoms [5].

Crinoids, intriguing marine animals within the Echinodermata phylum, have captured the interest of scientists for centuries. Historically significant to evolutionary biology, crinoids provide insights into the Cambrian explosion and subsequent echinoderm evolution [6]. Research on extant crinoids has been particularly robust from the early to mid-20th century, with more than 100 publications focusing on morphology, taxonomy, and classification [7,8]. The advent of molecular techniques has marked a new era in crinoid studies, with molecular phylogenetic methods shedding light on the evolutionary relationships among existing species [5,9–11]. Recent molecular studies, particularly concerning crinoid development, have underscored their unique life history and evolutionary trajectory [12–14]. For instance, Summers et al. conducted a molecular phylogenetic analysis on the Comatulidae family, reassessing morphological features for classification and proposing a new taxonomy [5,15]. This involved integrating molecular, morphological, behavioral, and biogeographic data to identify key traits and describe species. However, the current phylogeny of crinoids is undergoing significant revisions due to its instability [16]. Morphological traits are less reliable for classifying juvenile crinoids, which are smaller and more variable [17]. Robust molecular systematics are therefore critical for establishing consistent phylogenies.

Mitochondrial genomes have become invaluable in elucidating phylogenetic relationships and advancing evolutionary biology, thanks to features such as maternal inheritance and rapid evolutionary rates [18,19]. While mitochondrial research in crinoids has not been as extensive as in other groups, recent advancements in sequencing technology and enhanced sampling have started to bridge this gap. To date, complete mitochondrial genomes have been cataloged for two orders—Cyrtocrinida and Comatulida—and six families therein, including Sclerocrinidae, Colobometridae, Comatulidae [20], Antedonidae [21], Mariametridae [22] and Pentametrocrinidae [23].

To clearly understand crinoid phylogeny, this study presents the sequencing and annotation of mitochondrial genomes from 13 crinoid species spanning a range of depths and habitats. Our analysis aims to build upon the existing body of work by comparing new mitogenomic data with those from previous studies, which have typically been analyzed independently. Integrating new and existing datasets allows us to revisit and clarify the evolutionary history of crinoids. Our phylogenetic analysis supports the established idea that crinoids were ancestrally stalked, with stalkless forms evolving from these ancestors [11]. Interestingly, previous studies have shown that certain stalkless crinoids can secondarily become stalked.

By providing a detailed exploration of crinoid mitochondrial evolution, this study lays the groundwork for future research into the developmental and evolutionary aspects of crinoid biology.

2. Materials and Methods

2.1. Sample Collection and Mitogenome Sequencing

The samples were collected from various locations, including the Indian Ocean Ninety Degree Ridge, Carlsberg Ridge Seamount, Wuzhizhou Island, Sanya, Hainan, the East China Sea, Weizhou Island, Beihai, Guangxi, Hainan Xisha Ganquan Island, and the waters near Fuqing, Fujian, and Hainan Xisha Yongxing Island (Table 1). In this study, species within the order Comatulida have been classified as stalkless crinoids, whereas species belonging to the orders Hyocrinida, Isocrinida, and Cyrtocrinida are described as stalked crinoids.
Table 1. Species information, collection methods, and sampling locations.

<table>
<thead>
<tr>
<th>Sampling Location</th>
<th>Collection Method</th>
<th>Species Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indian Ocean Ninety Degree Ridge</td>
<td>TV grab</td>
<td>Thaumatocrinus sp.</td>
</tr>
<tr>
<td>Carlsberg Ridge Seamount</td>
<td>TV grab</td>
<td>Hyocrinidae sp.</td>
</tr>
<tr>
<td>Wuzhizhou Island, Sanya, Hainan</td>
<td>Diving collection</td>
<td>Comanthus parvicirrus</td>
</tr>
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<td>Wuzhizhou Island, Sanya, Hainan</td>
<td>Diving collection</td>
<td>Comaster schlegelii</td>
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<td>Wuzhizhou Island, Sanya, Hainan</td>
<td>Diving collection</td>
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</tr>
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<td>Wuzhizhou Island, Sanya, Hainan</td>
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<td>Capillaster sp.</td>
</tr>
<tr>
<td>East China Sea</td>
<td>Benthic trawl</td>
<td>Metacrinus rotundus</td>
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<td>Ptilometra sp.</td>
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<tr>
<td>Weizhou Island, Beihai, Guangxi</td>
<td>Diving collection</td>
<td>Zygometra comata</td>
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<td>Hainan Xisha Ganquan Island</td>
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<td>The waters near Fuzing, Fujian</td>
<td>Benthic trawl</td>
<td>Tropiometa macrodiscus</td>
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<tr>
<td>Hainan Xisha Yongxing Island</td>
<td>Diving collection</td>
<td>Comatella stelligera</td>
</tr>
</tbody>
</table>

After the surface mud stains were washed away, the samples were fixed with 100% ethanol and stored at −20 °C. They were then transported to the laboratory for further processing. Crinoid barbs were separated and cleansed with sterilized distilled water. The total DNA was subsequently extracted using the EZNATM MicroElute Genomic DNA Kit (OMEGA, Buffalo, NY, USA), and its quality was verified by electrophoresis on a 1% agarose gel. Novogene Technology Co., Ltd. (Beijing, China) was engaged to construct a second-generation data database for the DNA that passed quality inspection. Using a Covaris ultrasonic breaker (Covaris, East Sussex, UK), the DNA was fragmented into pieces of 300–500 bp, which were then end-repaired by a double-strand enzyme. An ‘A’ base was added to the 3’ end of each fragment before attaching the double-strand sequencing adapters. This was followed by electrophoretic purification and bridge PCR amplification to finalize the next-generation sequencing library. Once the library passed quality control, it was calibrated according to effective concentration and the desired output volume for sequencing. Finally, sequencing was carried out using the Illumina Novaseq 2000 platform (PE150, 10×) in a paired-end configuration.

2.2. Mitogenome Assembly and Annotation

The raw sequence data were initially processed using Trimmomatic version 0.39 [24] (code: java -jar trimmomatic-0.39.jar PE -phred33 R1.fq R2.fq R1_clean.fq R1_un.fq R2_clean.fq R2_un.fq R2ILLUMINACLIP:TruSeq2-PE.fa:2:30:10:8:true LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:100) to trim adapters and low-quality sequences. Subsequent splicing and assembly were conducted using MitoZ version 3.0 [25]. The MITOS web service [26] was employed for the prediction of protein-coding genes (PCGs), transfer RNAs (tRNAs), and ribosomal RNAs (rRNAs). We performed a BLAST search [27] against the NCBI Nucleotide Collection (NR/NT) database based on the predicted mitochondrial gene. Alignment and subsequent correction of the protein-coding sequences were performed using MEGA X [28], followed by a comparison with previously published sequences of crinoid mitochondrial genomes. Complete assemblies of 13 mitochondrial genomes have been submitted to the GenBank database (see Table 2).

Table 2. Species information from mitochondrial phylogenetic analysis (the accession numbers of mitochondrial genomes provided in this study are in bold).

<table>
<thead>
<tr>
<th>Order</th>
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<td>Crossaster papposus</td>
<td>MW046047</td>
<td>[35]</td>
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</table>

2.3. Comparative Mitogenome Analyses

The crinoid mitochondrial genome sequence was analyzed using MEGA X [28] and PhyloSuite v.1.2.3 [36], including relative synonymous codon usage (RSCU), base composition, and amino acid composition. The relative usage of synonymous codons is calculated as RSCU = S × Nc/Na, where ‘S’ represents the number of synonymous codons that encode the same amino acid, ‘Nc’ is the frequency of a particular codon within the genome, and ‘Na’ is the total count of all codons that encode that amino acid across the genome. This ‘Na’ serves as a measure of how frequently each synonymous codon is used relative to others that encode the same amino acid [37]. The calculation of the RSCU value reflects changes in coding efficiency and gene expression, which aids in understanding the evolution of the genetic code and the adaptive differences among populations [38]. The skewness of nucleotide composition was calculated using the following formulas: AT-skew = (A – T)/(A + T) and GC-skew = (G – C)/(G + C) [39]. To examine gene structure and variation across crinoid families, the 13 new mitochondrial genomes were compared to known crinoid sequences. Gene order rearrangements were analyzed using the online program CREx (http://pacosy.informatik.uni-leipzig.de/crex, accessed on 10 November 2023) [40].

2.4. Phylogenetic Analysis

Traditionally, echinoderm phylogeny has relied on morphology and fossils. Now, DNA and protein sequences are the main data for establishing phylogenies. Outgroups are added to determine the root. To study the phylogeny within the crinoid class, based on 13 PCGs and two ribosomal RNAs of 25 species of crinoids—using A. squamata, E. cordatum, and C. papposus as outgroups—the phylogenetic trees of crinoids were constructed using Bayesian inference (BI) and maximum likelihood (ML). MAFFT v.7.505 software [41] was used for auto multiple-sequence alignment, and MACSE v.2 [42] was subsequently used to refine the alignment of the 15 genes. Gblocks v.0.91b software [43] was then utilized to eliminate inconsistent positions and divergent regions in the aligned sequences [44], making the final alignment more suitable for phylogenetic analysis. We utilized the concatenation function of PhyloSuite v.1.2.3 [36] to generate
concatenated files for the 15 genes. The optimal evolutionary model GTR + F + I + G4 was selected using ModelFinder (part of IQ-TREE v.1.6.8) [45]. Bayesian inference analysis using MrBayes 3.2 [46] was used for phylogenetic reconstruction with the following settings: 1,000,000 generations, four Markov-chain Monte Carlo chains, trees sampled every 100 generations, and an initial 25% of trees discarded as burn-in. Effective sample sizes (ESS) were monitored to ensure sufficient mixing and convergence of the MCMC chains. All parameter ESS values exceeded 200 after burn-in, indicating sufficient sampling. After the burn-in phase, we used the remaining trees to generate a 50% majority-rule consensus tree. Phylogenetic analyses were conducted with IQ-TREE v.1.6.8 [47], employing the maximum likelihood (ML) method with automatic model selection and 1000 ultrafast bootstrap replicates for tree support. Finally, the generated BI and ML phylogenetic trees were visualized using the iTOL (http://itol.embl.de/, accessed on 15 November 2023) [48].

2.5. Selection Pressure Analysis

Selection pressure analysis is a pivotal method for investigating how biological functional genes adapt to specific environmental conditions or factors. In our study, selection analyses were conducted using CODEML within the PAML 4.10 suite [49]. The nonsynonymous to synonymous substitution ratio (dN/dS) serves as a crucial metric for quantifying selective pressures acting on genetic sequences [50]. It is generally considered that synonymous mutations are not subject to natural selection, while nonsynonymous mutations are affected by natural selection. Typically, a dN/dS ratio greater than 1 indicates positive selection, suggesting that advantageous mutations are being selected for; a dN/dS ratio of 1 implies no selection, corresponding to neutral evolution; and a dN/dS ratio less than 1 but greater than 0 suggests purifying selection. The smaller the dN/dS value, the stronger the purifying selection pressure, indicating a more conserved amino acid sequence [51]. In these analyses, we designated stalked crinoids (n = 3) as the foreground group and stalkless crinoids (n = 22) as the background group. The rationale for selecting three stalked crinoids as the foreground clade for positive selection analysis is predicated on the unique genetic adaptations they may possess compared to their stalkless counterparts. Stalked crinoids, with their fixed lifestyle and close association with specific deep-sea substrates, may be subject to distinct metabolic and adaptive selection pressures on their mitochondrial genes.

We computed the dN/dS ratio for 13 mitochondrial protein-encoding genes, using a Bayesian phylogenetic tree of 25 crinoids as the guide. A single-proportion model was applied to estimate a uniform dN/dS ratio across all phylogenetic branches, and a branch-site model was employed to detect signatures of positive selection on the foreground branch specifically. The most suitable model within the nested set was determined using a likelihood ratio test (LRT), with the significance of the model appraised through a chi-square test ($\chi^2$), comparing the $2\Delta\ln L$ value against the chi-square distribution for the single-proportion model. Upon passing the LRT and exhibiting a statistically significant p value, the Bayesian Empirical Bayesian (BEB) approach [52] was utilized to identify sites of positive selection, denoting those with a posterior probability (PP) greater than 0.9 as candidates for positive selection. We identified amino acid sequences from positively selected sites. These sequences were used to model the three-dimensional structure of mitochondrial proteins. The proteins studied are the products of protein-coding genes (PCGs) from M. rotundus. For the modeling, we employed homology modeling techniques provided by the Swiss Model Server (https://swissmodel.expasy.org, accessed on 20 October 2023) [53–55].

3. Results

3.1. Mitogenome Structure and Organization

In this study, we expanded the crinoid mitogenome database by annotating 13 new species and analyzed the mitochondrial genomes of 25 species across 4 orders and 11 families. The lengths of the 25 complete mitochondrial genomes of crinoids range from 15,772 bp to 16,850 bp. All mitogenomes are circular and double-stranded molecular structures with
37 genes (22 tRNAs, 13 PCGs, 2 rRNAs). Among the 13 protein-coding genes, 10 are located on the heavy strand (H-strand), namely cox1, cox2, cox3, atp6, atp8, nad3, nad4, nad4L, nad5, and cytb, while the remaining genes are situated on the light strand (L-strand), including nad1, nad2, and nad6. Both ribosomal RNA genes (rrnL and rrnS) are found on the L-strand. All species showed a higher AT than GC content. The nucleotide composition of all protein-coding genes, as well as the rRNA genes, demonstrates a clear preference for A and T nucleotides.

The average contents of A, G, T, and C bases in these 25 mitogenomes were 25.08%, 16.82%, 46.48%, and 11.61%, respectively. The G base content of N. richeri and Metacrinus rotundus was higher than the A base content, while the remaining 23 species showed a higher A base content than the G base content. The A/T content ranged from 64.5% to 74.2% (Table S1), exhibiting an AT bias.

Among the 25 crinoids examined, including 13 that were newly sequenced, five distinct mitochondrial gene arrangement patterns have been identified (Figure 1). The predominant arrangement, found in 21 crinoid species, is categorized as pattern 1. In this pattern, the sequences of rrnL and rrnS genes are notably conserved, with rrnL located between trnG and trnY, and rrnS situated between trnE and trnF. Compared to pattern 1, the trnL gene in Aneissia pinguis (pattern 2) undergoes an inversion, shifting from the L-strand to the H-strand. Pattern 3 is represented by Neogymnocrinus richeri, where a translocation of the nad4L gene is observed. In Tropiometra macrodiscus (pattern 4), the gene clusters trnR-nad4L translocate from downstream of cox1 gene to the upstream position of trnS1 gene. Notably, the gene order of Antedon mediterranea (pattern 5) is basically consistent with that of Tropiometra macrodiscus, except for the translocation of trnA and trnV genes.

<table>
<thead>
<tr>
<th>Pattern</th>
<th>Gene Arrangement</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>cox1 (\rightarrow) nad4L, cox2, K, atp8, atp6, cox3, S2, nad4, H, S1, nad5, nad6, cytb, P, Q, N, L, A, V, C, Y, M, D, T, E, rrs, T, L2, G, rrs, Y, nad2, E, nad1</td>
</tr>
<tr>
<td>2</td>
<td>cox1 (\rightarrow) nad4L, cox2, K, atp8, atp6, cox3, S2, nad4, H, S1, nad5, nad6, cytb, P, Q, N, L, A, V, C, Y, M, D, T, E, rrs, T, L2, G, rrs, Y, nad2, E, nad1</td>
</tr>
<tr>
<td>3</td>
<td>cox1 (\rightarrow) cox2, K, atp8, atp6, nad4L, cox3, S2, nad4, H, S1, nad5, nad6, cytb, P, Q, N, L, A, V, C, Y, M, D, T, E, rrs, T, L2, G, rrs, Y, nad2, E, nad1</td>
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<tr>
<td>4</td>
<td>cox1 (\rightarrow) cox2, K, atp8, atp6, cox3, S2, nad4, H, S1, nad5, nad6, cytb, P, Q, N, L, A, V, C, Y, M, D, T, E, rrs, T, L2, G, rrs, Y, nad2, E, nad1</td>
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<td>5</td>
<td>cox1 (\rightarrow) cox2, K, atp8, atp6, cox3, S2, nad4, H, S1, nad5, nad6, cytb, P, Q, N, L, A, V, C, Y, M, D, T, E, rrs, T, L2, G, rrs, Y, nad2, E, nad1</td>
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</tbody>
</table>

**Figure 1.** Five gene arrangement patterns of 25 crinoids (the symbol “—” denotes that the gene is located on the light strand (L-strand). In comparison to the gene order of pattern 1, sequence fragments that are rearranged in the other patterns are highlighted in green and blue marks. This figure was generated using the online program CREx.

3.2. Protein-Coding Genes

Among the 25 mitogenomes, 9 genes (cox1-3, cytb, nad1-4, and nad6) use “ATG” as the start codon, while the remaining 3 genes (atp8, nad4L, nad5) have “GTG” codons (Table S2).

Eight PCGs (atp6, atp8, cox2-3, nad3, nad4, nad4L, and nad5-6) terminate with complete stop codons (“TAA” or “TAG”), while five genes (cox1, cytb, nad1-2, and nad4) have incomplete stop codons (“TA-” or “T-”) in some crinoids. More specifically, the cox1 and cytb genes use “TA”, while cytb, nad1, nad2, and nad4 have terminal “T” (Table S3).

RSCU analysis indicates a marked preference for specific codons, including UCU, GUU, GCU, and GCU, across the protein-coding genes (PCGs) of 25 crinoid mitogenomes (Figure 2). Notably, the UCU codon is highly favored in Hyocrinidae sp., and UUA is favored in Comatella stelligera, both exhibiting significant overexpression with an RSCU value of 3.94, suggesting a strong codon bias. In contrast, the UUA codon, which codes for leucine (Leu2), is underrepresented in Neogymnocrinus richeri and Metacrinus rotundus when compared to other crinoids, indicating a unique codon usage pattern in these species that could be reflective of evolutionary or adaptive differences.
3.2. Protein-Coding Genes

Among the 25 mitogenomes, 9 genes (\textit{cox1-3}, \textit{cytb}, \textit{nad1-4}, and \textit{nad6}) use “ATG” as the start codon, while the remaining 3 genes (\textit{atp8}, \textit{nad4L}, and \textit{nad5}) have “GTG” codons (Table S2).

Eight PCGs (\textit{atp6}, \textit{atp8}, \textit{cox2-3}, \textit{nad3}, \textit{nad4L} and \textit{nad5-6}) terminate with complete stop codons (“TAA” or “TAG”), while five genes (\textit{cox1}, \textit{cytb}, \textit{nad1-2}, and \textit{nad4}) have incomplete stop codons (“T A-” or “T-”) in some crinoids. More specifically, the \textit{cox1} and \textit{cytb} genes use “TA”, while \textit{cytb}, \textit{nad1}, \textit{nad2}, and \textit{nad4} have terminal “T” (Table S3).

RSCU analysis indicates a marked preference for specific codons, including UCU, UUA, GUU, CCU, and GCU, across the protein-coding genes (PCGs) of 25 crinoid mitogenomes (Figure 2). Notably, the UCU codon is highly favored in \textit{Hyocrinidae} sp., and UUA is favored in \textit{Comatella stelligera}, both exhibiting significant overexpression with an RSCU value of 3.94, suggesting a strong codon bias. In contrast, the UUA codon, which codes for leucine (Leu2), is underrepresented in \textit{Neogymnocrinus richeri} and \textit{Metacrinus rotundus} when compared to other crinoids, indicating a unique codon usage pattern in these species that could be reflective of evolutionary or adaptive differences.

Figure 2. Relative synonymous codon usage (RSCU) and codon distribution in PCGs of 25 crinoid mitogenomes (the symbols in brackets on the X-axis represent the amino acid corresponding to each codon and use the one-letter nomenclature for that amino acid). Stalked crinoids in the Y-axis are highlighted with a blue background. Orange: high RSCU value; green: low RSCU value. This figure was generated using Origin 2018 [56].

3.3. Phylogenetic Analyses

Phylogenies were constructed using 13 PCGs and 2 rRNAs from 25 crinoid taxa and outgroups, with Bayesian inference (BI) and maximum likelihood (ML) approaches (Figures 3 and 4). Except for the outgroup and the Ptilometridae family, the phylogenetic trees generated by the two approaches are nearly identical at the family and order levels of topology. The phylogenetic relationship among the four orders is (Isocrinida + (Cystocrinida + (Hyocrinida + Comatulida))). As shown in Figure 4, the order Comatulida has three main phylogenetic clades: (1) Pentametrocrinidae, (2) Comatulidae, and (3) a clade represented by the remaining families, including Ptilometridae, Tropiometridae, Antedonidae, Himerometridae, Stephanometridae, and Colobometridae.

More specifically, \textit{Thaumatocrinus} sp. and \textit{Thaumatocrinus naresi} are separately aggregated into the first clade of the Comatulida order (Figures 3 and 4). In the second clade, the Comatulidae family is divided into two branches, namely the Comatellinae subfamily (\textit{Comatella nigra} and \textit{Comatella stelligera}) and the Comatulinae subfamily (the other five genera). Furthermore, the Anneissia genus seems to be a polyphyletic lineage due to the separation of its species into \textit{Anneissia intermedia}. 
The phylogenetic relationship among the four orders is (Isocrinida + (Cyrtocrinida + (Hyocrinida + Comatulida))). As shown in Figure 4, the order Comatulida has three main phylogenetic clades: (1) Pentametrocrinidae, (2) Comatulidae, and (3) a clade represented by the remaining families, including Ptilometridae, Tropiometridae, Antedonidae, Himerometridae, Stephanometridae, and Colobometridae.

Figure 3. Bayesian phylogenetic tree constructed based on 13 PCGs and 2 rRNA genes (the species marked in red are the species sequenced in this study; posterior probabilities values are indicated above nodes). This figure was generated using iTOL (http://itol.embl.de/, accessed on 15 November 2023).

For the third phylogenetic clade within the Comatulida order, the Himerometridae, Stephanometridae, and Colobometridae families are supported as a monophyletic sister group, receiving strong bootstrap support in the ML analysis and high posterior probabilities in the BI analysis. Moreover, the evolutionary relationship of the Ptilometridae family within this third clade is ambiguous, and we need more molecular data to illuminate this.

Notably, the Antedonidae family is polyphyletic, with its species divided into three main branches based on the BI analysis. For instance, *Antedon mediterranea* and *Tropiometra macrodiscus* are depicted as an isolated lineage, reflecting their independent evolutionary histories. For the Florometra genus, two species fail to cluster together, but instead separately group with other genera. These results hint that feather stars have a complex evolutionary relationship that may warrant further investigation to elucidate.
Figure 4. Maximum likelihood phylogenetic tree constructed based on 13 PCGs and 2 rRNA genes (the species marked in red are the species sequenced in this study; bootstrap support values were indicated above nodes). This figure was generated using iTOL (http://itol.embl.de/, accessed on 15 November 2023).

3.4. Relaxed Selective Constraint and Positive Selection on the Mitochondrial Genes

Incorporating the three stalked crinoids as the foreground clade, the results of the single-ratio model showed that all PCGs exhibited a dN/dS ratio <1 (Table S4 and Figure 5), indicating purifying selection. Most PCGs have similar dN/dS ratios in both types of crinoids, suggesting comparable evolutionary pressures. Particularly noteworthy are the dN/dS ratios for atp8 in the foreground group, which decreased significantly to 0.0013 from 0.1256 in the background, suggesting an intensified purifying selection in the stalked crinoids.

In contrast, nad2 exhibited a slight increase in its dN/dS ratio from 0.06 in the background group to 0.0722 in the foreground, suggesting a relaxation in purifying selection, though the value still denotes purifying selection overall. This subtle change hints at potential variations in metabolic demands or mitochondrial function between the two groups. Other genes such as nad3, nad4, and nad5, while also under purifying selection, showed smaller differences in their dN/dS ratios between the foreground and background groups, which does not suggest positive selection but rather minor variations in the strength of purifying selection.
crinoids, suggesting comparable evolutionary pressures. Particularly noteworthy are the 
\( \frac{dN}{dS} \) ratios for \( atp8 \) in the foreground group, which decreased signifi-
cantly to 0.0013 from 0.1256 in the background, suggesting an intensi-
fi ed purifying selection in the stalked 
crinoids.

Figure 5. The ratio of nonsynonymous to synonymous substitutions (\( \frac{dN}{dS} \)) for 13 PCGs in 25 crinoids (with stalked crinoids as the foreground group and stalkless crinoids as the background 
group), generated using Origin 2018 [56].

The existing literature and preliminary studies suggest that these adaptive changes 
could play a pivotal role in the evolutionary trajectory of their mitochondrial genes [57]. Consequently, we focused on identifying signals of adaptive positive selection within this 
group’s mitochondrial genes. Notably, when the analysis was conducted with 22 stalkless 
crinoids as the foreground clade, no positive selection sites were detected, further sub-
stantiating our hypothesis that stalked crinoids may have undergone specific adaptive 
evolutionary processes.

Branch-site model analysis revealed eleven positively selected sites across six mito-
chondrial PCGs in stalked crinoids (Table 3). Four of these sites showed Bayes Empirical 
Bayes posterior probability over 0.99, indicating very strong positive selection signals. 
These included serine at site 2020 of \( \text{nad2} \); serine at site 2344 of \( \text{nad3} \); lysine at site 2568 of \( \text{nad4} \); and serine at site 3244 of \( \text{nad5} \). The remaining seven sites also showed high posterior 
probabilities between 0.954 and 0.987. The high probabilities of positive selection suggest 
these amino acid replacements likely conferred selective advantages during the evolution 
of stalked crinoids.

Table 3. Positively selected sites and genes in lineages under positive selection as indicated by 
Bayesian Empirical Bayesian (BEB) analysis. * BEB posterior probability > 95%; ** BEB posterior 
probability > 99%.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Site</th>
<th>Probability</th>
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<tbody>
<tr>
<td>cytb</td>
<td>1492 I</td>
<td>0.955 *</td>
</tr>
<tr>
<td>nad2</td>
<td>2020 S</td>
<td>0.996 **</td>
</tr>
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<td></td>
<td>2044 R</td>
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</tr>
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<td></td>
<td>2174 C</td>
<td>0.977 *</td>
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<td>nad3</td>
<td>2344 S</td>
<td>0.995 **</td>
</tr>
<tr>
<td>nad4</td>
<td>2495 F</td>
<td>0.977 *</td>
</tr>
<tr>
<td>nad4L</td>
<td>2568 K</td>
<td>0.999 **</td>
</tr>
<tr>
<td></td>
<td>2691 S</td>
<td>0.987 *</td>
</tr>
<tr>
<td>nad5</td>
<td>3124 N</td>
<td>0.967 *</td>
</tr>
<tr>
<td></td>
<td>3244 S</td>
<td>0.994 **</td>
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<tr>
<td></td>
<td>3403 K</td>
<td>0.954 *</td>
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To comprehensively assess the effects of positive selection sites and the alterations in physicochemical properties resulting from amino acid substitutions, we mapped the positive selection sites, identified by CODEML in *M. rotundus* species, onto the corresponding mitochondrial protein structure. Remarkably, our findings revealed that a significant proportion of the identified positively selected sites were situated within or near functional regions (Figure 6). This emphasizes their potential role in evolutionary adaptation.

Table 3. Positively selected sites and genes in lineages under positive selection as indicated by Bayesian Empirical Bayesian (BEB) analysis. * BEB posterior probability > 95%; ** BEB posterior probability > 99%.

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<td>3124</td>
<td>0.967 *</td>
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Figure 6. Atomic-level structures of computationally predicted mitochondrial protein-coding genes (*nad2*–*nad5*, *nad4L*, and *cytb*) of *M. rotundus*. Positively selected sites inferred by CODEML are shown in red; all sites identified by these methods are shown in orange and highlighted with arrows.

4. Discussion

4.1. Mitochondrial Genome Structural Characteristics

In our phylogenetic analysis, *N. richeri* and *M. rotundus* emerge as basal taxa, providing a unique vantage point to explore the ancestral states of nucleotide composition. Notably, we observed a pronounced evolutionary shift from guanine (G) to adenine (A) bases within these lineages, indicative of an overarching trend towards an AT bias. This bias is not merely a relic of past mutational events but appears to be a dynamic feature sustained over evolutionary timescales.

Mitochondrial DNA (mtDNA) tends to mutate faster than nuclear DNA, often resulting in a higher occurrence of adenine (A) and thymine (T) bases, which contributes to a condition known as AT bias. The tendency for AT enrichment may partly derive from the susceptibility of mtDNA to oxidative damage coupled with distinctive aspects of its replication process. Sequences with a higher AT content tend to be less stable compared to those rich in guanine (G) and cytosine (C), predisposing them to a more frequent mutational occurrence. Consequently, this can substantially alter the base composition of mtDNA. While this shift may be attributable in part to the known vulnerability of the light strand to mutation during replication, leading to the observed base composition [39], it is plausible that selective pressures have also played a role [58]. Moreover, this finding aligns with previous studies that have reported a similar AT bias in crinoid mitochondrial genomes [20,21], suggesting that the light strand’s proclivity for higher mutation rates could be a widespread phenomenon [59].
Echinoderm mitogenomes typically utilize an “ATG” start codon [60] and incomplete stop codons [61]. This study showed that not onlynad4L and nad5 [20,21] but also atp8 initiated with “GTG”. In other echinoderms, atp8 was also found to have “GTG” as the initiation codon [62–64]. Of the 13 sequenced mitogenomes in this study, all PCGs except cyt b terminated with complete stops (“TAA” or “TAG”). Cyt b in six species (Comanthus parvicirrus, Comanthus sp., T. macrodiscus, Capillaster sp., Annessi. bennetti, and Hyocrinidae sp.) used an incomplete “T” stop codon.

The balance between base mutation bias and natural selection is expressed as codon bias [65,66]. In 23 crinoid species except N. richeri and M. rotundus, the content of the A base was higher than that of the G base. In N. richeri and M. rotundus, the UUA codon for leucine Leu2 is under-expressed relative to other stalked crinoids. We hypothesize that the imbalance in nucleotide composition, specifically the asymmetry between the strands in mitochondrial DNA, contributes to the observed preferences in codon usage among these crinoid species.

Here, we expand upon the significance of asymmetric PCG codon usage. The asymmetric usage of codons, where H-strand genes prefer G/T terminal codons and L-strand genes favor A/C terminal codons [67], may have substantial implications for mitochondrial genome function and evolution in crinoids. Such biases in codon usage can influence gene expression levels, potentially due to the differential availability of tRNA and the efficiency of the translation process [68]. Additionally, these biases could impact the speed and accuracy of protein synthesis, affecting protein folding and function [69]. In the context of mitochondrial genomes, these codon biases may reflect the organism’s adaptations to specific energetic and metabolic demands [38]. The under-expression of the UUA codon for leucine (Leu2) in N. richeri and M. rotundus may indicate the adaptation of their mitochondrial genomes to their unique ecological niches, characterized by factors such as temperature, salinity, and available energy resources. These environmental factors could exert selective pressures that favor certain mitochondrial codon usages over others, leading to more efficient protein synthesis in the specific habitats in which these crinoids thrive.

In our study, 25 species of crinoid mitogenomes exhibited five arrangement patterns. Mitogenome arrangement can reveal the phylogenetic relationship between different species [70,71]. In recent years, several studies on the arrangement of mitochondrial genes in echinoderms have been reported [62,72]. Crinoidea [62] and Ophiuroidea [73,74] exhibit greater gene order variability than other echinoderms. The mitogenome order varies most in Crinoidea; some crinoids (e.g., A. mediterranea and N. richeri) no longer showed conserved gene blocks present in the other four echinoderm classes [75].

4.2. Phylogenetic Relationships

At present, echinoderm phylogenetics traditionally focuses on interphyla and interclass relationships. Crinoids occupy a pivotal position in the phylogenetic tree of echinoderms [34], yet their intraclass evolutionarv relationships, especially through the lens of mitochondrial genomes, remain underexplored. Our study aims to fill this gap by utilizing mitochondrial genome sequences to construct phylogenetic trees, thus providing a new insight into the intraclass relationships among crinoids using this molecular approach. In this study, we used A. squamata, E. cordatum, and C. papposus as outgroups, and constructed Bayesian and ML phylogenetic trees with 25 species.

Inferred phylogenetic relationships from the ML analysis, based on the concatenated fifteen-gene complete dataset (PCGs and rRNA genes of mitochondrial genomes) within the Himerometroididea [76], corroborate previous findings suggesting a close kinship among Himerometridae, Stephanometridae, and Colobometridae. The phylogenetic relationships within the Antedonidae family are complex. Currently, related studies based on molecular data (COI, 16S, 28S, and ITS), morphology, and multivariate analyses show that the Antedonidae family is a polyphyletic group [11]. Our study also extends this insight by demonstrating that two species from the Antedonidae and Tropiometridae families exhibit a close genetic relationship, clustering as an isolated lineage. In addition, they both exist as
translocations of nad4L and trnR, suggesting that these translocations may be conserved in this lineage. Moreover, two species of the Florometra genus within the Antedonidae family separately group with other genera. Meanwhile, this same inconsistency is obvious for Anneissia species from the Comatulidae family. This genetic closeness and inconsistency, however, present a conundrum regarding the taxonomic delineations within Antedonidae and Comatulidae, as our data suggest a potential revision of their classification may be necessary.

The widespread distribution and diverse habitats of the Crinoidea class have given rise to an extensive variety of morphological features, demonstrating notable variation even among species that are closely related. This morphological plasticity often complicates species’ identification and may lead to misclassifications. The current findings emphasize the utility of molecular data in providing clarity and resolving ambiguities that morphological analyses alone may not adequately address [77]. Such discrepancies highlight the need for a taxonomic reassessment that incorporates both molecular and morphological data to ensure accurate species classification and a deeper understanding of crinoid diversity [78,79].

4.3. Positive Selection Site

Natural selection is widely recognized as a key evolutionary force [80]. Mitochondrial genes are considered to undergo purifying selection to maintain function [81]. Positively selected genes drive adaptive evolution [82]. The dN/dS ratio indicates the evolutionary rate [83–86]. There remains a significant debate concerning whether positive selection or purifying selection prevails.

All 13 PCGs exhibited a dN/dS ratio < 1, signifying prevalent negative selection that acts to remove deleterious mutations while preserving conserved amino acid sequences. This observation is consistent with the known functions of mitochondrial genes, as changes in these genes can significantly impact an organism’s survival. Notably, the dN/dS ratio for atp8 was significantly higher in stalkless crinoids (0.1256) compared to stalked crinoids (0.0013). This difference may suggest that stalkless crinoids either face reduced selective constraints or have advantageous mutations in the atp8 gene that confer adaptive benefits. Such advantageous mutations in the atp8 gene could have significant implications for the evolution of the mitochondrial genome. They might lead to compensatory evolutionary changes in other mitochondrial genes, a phenomenon we refer to as ‘compensatory draft feedback’, where alterations in one part of the genome necessitate or drive changes in another to maintain overall mitochondrial function and efficiency [87,88]. Species under significant negative selection may evolve faster to maintain mitochondrial function. Therefore, the atp8 gene may play a crucial role in the mitogenomic evolution of stalkless crinoids. Compared to stalkless crinoids, stalked crinoids showed lower dN/dS ratios for nad3, nad4, and nad5, suggesting stronger negative selection on more essential functions. The ineffectiveness of negative selection allows for the accumulation of deleterious mutations [89].

Divergent PCG selection sites between crinoid groups distributed across cyt b, nad2, nad3, nad4L, nad4, and nad5 indicate adaptive diversification. Positively selected sites within the cyt b gene, particularly in regions critical to its function, suggest that deep-sea crinoids may have evolved adaptations to their extreme habitats. These modifications could enhance the efficiency of oxidative phosphorylation [90], allowing these organisms to optimize energy production in low-oxygen environments. Nad genes encode electron transport proteins [91], with nad2, nad4, and nad5 integral to proton pumping [92]. Furthermore, changes in the Nad genes, which encode components of the electron transport chain, may result in proteins that are better suited to the unique features of deep-sea life [93] such as high pressure and cold temperatures. Such adaptations could lead to increased metabolic efficiency, allowing these crinoids to survive and function effectively where resources are scarce and conditions are challenging. The deep sea features anoxia, darkness, high pressure, and low temperatures [94]. Under these extreme environmental conditions,
organisms may require improved and adapted energy metabolism. Positive selection sites in nad family genes, found in a variety of deep-sea species such as corals [95], sea cucumber [96], mussels [97], and starfish [98], suggest specific evolutionary adjustments. These sites may alter the structure and function of the NADH dehydrogenase complex, which could lead to variations in the efficiency of the electron transport chain. For example, modifications in the gene products might improve the organism’s ability to generate energy under the low-oxygen conditions of the deep sea. Enhanced energy production could, in turn, influence the organism’s resilience to the high pressures and cold temperatures characteristic of these environments.

5. Conclusions

This study markedly extends the crinoid mitochondrial genome database by adding 13 newly sequenced genomes, thus providing new resources for future research. Such expansions in the database not only facilitate the identification of species-specific mitochondrial markers but also underpin the comparative studies necessary for resolving the complex phylogenetic relationships within the Echinodermata. The identification of five mitochondrial gene arrangement patterns offers profound insights into the evolutionary processes of crinoids. These patterns represent genomic structural variations that may correlate with key ecological and evolutionary transitions in crinoid history, such as changes in habitat or feeding strategies. By examining the variations in gene arrangements alongside the phylogenetic information, we can begin to unravel the adaptive significance behind these mitochondrial configurations. Arrangement comparisons of the mitochondrial genome may be useful for phylogenetic reconstruction. Moreover, additional molecular data are essential to deepen our understanding of how mitochondrial gene arrangement contributes to the evolution of crinoids.

Our selective pressure analysis revealed positive selection in several genes—cytb, nad2, nad3, nad4, nad4L, and nad5—indicating adaptive evolution at the molecular level. These genes, integral to the respiratory chain, may have evolved under selective pressures to optimize energy production, which is critical for crinoid survival in diverse marine environments.

The monophyly of crinoid families remains a topic of debate, with in-depth phylogenetic studies being scarce. Based on the mitochondrial genome data, our phylogenetic reconstruction reveals the polyphyly of certain genera and families within the Comatulida order. This result not only offers molecular insights into crinoid phylogenetic relationships but also uncovers the intricate tapestry of their evolutionary history. Providing a statistical backbone to these findings, such as bootstrap values and posterior probabilities, strengthens our confidence in these phylogenetic interpretations and invites further investigation into the evolutionary mechanisms at play. As we amass new mitochondrial genome data, it is equally important to collect more nuclear data, such as ITS, 18S rRNA, 28S rRNA, and Histone H3. Combining both mitochondrial and nuclear sequence data is likely to yield more comprehensive and accurate results.

Furthermore, our study sheds light on the potential of mitochondrial markers in species identification and phylogenetics in crinoids, asserting their utility in this domain. Practical applications of these enhanced mitochondrial markers may include the development of more precise conservation strategies or the implementation of mitogenomic tools in monitoring biodiversity and ecosystem health.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/jmse12030361/s1, Table S1: Base composition of mitochondrial genomes of 25 crinoids; Table S2: Putative start codon of 13 PCGs in 25 crinoids; Table S3: Putative terminal codon of 13 PCGs in 25 crinoids; Table S4. The ratio of nonsynonymous to synonymous substitutions (dN/dS) for 13 PCGs in 25 crinoids with stalked crinoids as the foreground group and stalkless crinoids as the background group.
Author Contributions: Conceptualization, Q.X. and X.H. (Xuebao He); Funding acquisition, Q.X.; Investigation, Z.L.; Project administration, Q.X.; Validation, Y.L.; Visualization, Y.D.; Writing—original draft, Q.X., M.L. and Y.S.; Writing—review and editing, X.H. (Xuying Hu), Q.Z. and B.L. All authors have read and agreed to the published version of the manuscript.

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Conflicts of Interest: The authors declare no conflicts of interest.

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