



[2, 3]. After that, mitochondria coevolved with different hosts and underwent both neutral modifications and adaptive responses that led to the diversity observed today in mitogenomes [4]. In bilaterians, mitogenomes were considered to be extremely compact and normally organized into a single circular molecule ranging in size from 14 to 20 kb [5]. Bilaterian mitogenomes typically contain the same set of 37 genes (13 protein-coding genes encoding different subunits of enzyme complexes for the oxidative phosphorylation (OXPHOS) system, 2 ribosomal RNAs (*rrnS* and *rrnL*), 22 transfer RNA genes) and no introns [5–7]. In the last few years, high-throughput sequencing techniques and extensive sampling for phylogenetic and population genetic studies have accelerated the sequencing of mitogenomes and uncovered the great diversity of structural features [8]. An increasing number of mitogenomes seem to deviate dramatically from typical bilaterian mitogenomes and present wide variation in genome size, and many of them are much larger than 20 kb. Many molluscs, especially bivalves, display an unusual amount of variation in mitogenome structure and size, even among closely related species [6, 9, 10].

Arcidae, known as Ark shell or blood cockles, are an economically-important group of bivalves and have a long evolutionary history, dating back to the Lower Ordovician ~ 450 Mya [11]. Interestingly, mitogenomes of Arcidae species both within and between species reveal a high variability in size, ranging from 19 to 56 kb in length [12]. For example, reported mitogenomes of *Scapharca broughtonii* are 46,985 bp [13] and 48,161 bp [14], the recently published *Scapharca gubernaculum* mitogenome is 45,697 bp [15], which are 2–3 times larger than other bilaterians. The largest Arcidae mitogenome comes from *Scapharca kagoshimensis* (46.7–56.2 kb) and is the largest bilaterian mitogenome yet recorded, out of approximately 86,900 mt-DNAs from more than 11,600 species [8, 12]. In addition, large mitogenomes are also found in sea scallop *Placopecten magellanicus* (31–41 kb) [16] and the clavagellid mussel *Bryopa lata* (32 kb) [17].

The large genome sizes in ark shells and sea scallop are not primarily a result of duplications of control region sequences and coding sequences like model organisms, but rather the expansion of unassigned regions (i.e., non-coding regions that are functionally unassigned) [16, 18]. A previous analysis [19] of 2656 complete mitogenomes showed that some bivalves have a proportion of unassigned regions (URs) that are significantly different from all other groups and show the highest median percentage of URs in Metazoans. According to the mutation pressure theory [20], fast evolving organelle genomes experience more selection pressure for genome reduction, but some bivalve mitogenomes seem to contradict this theory. In some Arcidae species, URs account for more

than 50% of the entire mitogenomes. Tandem repeats, inverted repeats and transposable elements in unassigned regions (URs) have been shown to contribute to the large size of these mitogenomes [12, 13, 16], but it does not mean they are the main cause of huge expansion in URs. Data from previous studies show that repeat families and transposable elements are not the main components of large URs, which only account for 6–31% of URs in different Arcidae species, though they have a significantly positive correlation with mitogenome size [12, 13]. This suggests that there are other components influencing the size of URs. One possible explanation is that retention of the URs in bivalve mitogenomes is caused by the presence of functional sequences and/or structures. However, to date, much of the work on URs of Arcidae has focused on nucleotide-level analyses to observe sequence characteristics (e.g., tandem repeats, inverted repeats), mitogenome expansion remains poorly understood and needs further study with different perspectives.

Mitochondria are long known for bioenergetics, but they also have novel non-OXPHOS-related adaptations and functions [21]. With the increasing number of published mitogenomes, non-standard gene contents have been found in different animal groups, and additional mitochondrial protein-coding genes have been identified and annotated in mitogenomes of metazoan, particularly in invertebrates. For example, additional mitochondrial protein-coding genes were first discovered in the octocoral *Sarcophyton glaucum* [22], which was a homolog of *mutS* and hypothesized to originate either through bacterium or viral infection by horizontal gene transfer [23, 24]. In cnidarians, sponges and placozoans, protein-coding genes with non-OXPHOS functions (e.g., *dnaB*, *tatC*) have been also reported [25, 26]. Surprisingly, nine additional mtDNA-encoded protein genes have been described in humans [27–30], one of which is a 75 bp ORF in the mitochondrial 16S rRNA that acts as a neuroprotector, an antiapoptotic agent, and a cytoprotector [31, 32]. These discoveries indicate that there are additional functional sequences in mitochondria, maybe related to its diverse functions.

Moreover, multiple ORF sequences have also been found in the mitogenomes of bivalves. A novel ORF was discovered with no sequence- or domain-based homology to mitochondrial genes in the mitogenome of pearl-lip oyster *Pinctada maxima* but has domain-based homology to the nuclear genome [33]. Mitochondrial ORFans (open reading frames having no detectable homology and no known function) also have been identified in marine and freshwater bivalves (Mytiloidea, Nuculanoida, Unionoida, and Veneroida) with doubly uniparental inheritance (DUI) of mitochondrial DNA [34, 35]. In these cases, products are exported from the

organelle and may be involved in functions other than energy production [34–40]. These studies indicate that traditional bivalve mitochondrial non-coding regions have sequences or unassigned regions that potentially perform biological functions. The structure of some special ORFs in the mitogenome of *Tegillarca granosa* (Arcidae) have been briefly investigated [41], but the origin of Arcidae ORFs in large URs remain unclear. In addition, a fundamental question regarding the size of mitogenomes in Arcidae bivalves is whether there are ORFs in large URs that perform functions and have a connection with expanded size of mitogenomes.

Here, we sequenced and annotated five new mitogenomes (four *S. broughtonii* and one *S. cornea*) and assembled a mitogenome of *S. kagoshimensis* from NCBI data (SRX8857271). Six transcriptomes of *S. broughtonii* were sequenced and analyzed to identify conserved functional ORFs. Multiple samples from the same species were used to detect intraspecific variation in mitogenome length and the presence of ORFs. To better understand how URs expand and evolved in Arcidae, we present a comparative analysis of 32 complete mitogenomes (6 new assemblies and 26 published assemblies from NCBI) of Arcidae species to highlight both unique features and characteristics shared among different species, with an emphasis on characterizing large URs and ORFs. Then, we investigated the origin and duplication of ORFs and their correlation with mitogenome expansion, and particularly with the expansion and function of mitochondrial large URs in Arcidae.

## Results

### Mitogenome assembly, annotation and features

Complete mitogenomes sequences of four *S. broughtonii*, one *S. kagoshimensis* and one *S. cornea* had sequence lengths more than 40 kb (Table 1). Four new *S. broughtonii* mitogenomes sequences varied in size from 44,327 bp to 48,560 bp, close to previously published *S. broughtonii* mitogenome (46,985 bp) [13]. The length of *S. kagoshimensis* mitogenome reported here was 54,157 bp, slightly smaller than the previously reported *S. kagoshimensis* (56,170 bp) [12], and *S. cornea* is 46,362 bp long. *S. broughtonii* and *S. kagoshimensis* mitogenomes vary dramatically in length within species. All mitogenomes consisted of 12 protein-coding genes (all taxa lacked *atp8*), two ribosomal RNA genes (*rrnS* and *rrnL*) and 27–33 tRNA genes (Additional file 1: Table S1). *Atp8* has never been found in Arcidae species [12, 18, 41, 42]. All protein-coding genes and rRNA genes in the six mitogenomes are encoded on the same strand and share the same gene order. In addition, a duplication of *cox2* was observed in all six mitogenomes (Additional file 1: Table S2), located between *cob* and *cox2*. The two copies have different

length: the *cox2* is 666–720 bp long (221–239aa), while the *cox2-b* is 1179–1431 bp long (392–476aa). This indicates that *cox2-b* have acquired an extension after *cox2* duplication. Four different start codons (ATG, ATA, ATT, GTG) were observed but most protein-coding genes start with the codon ATG, and stop with the TAA and TAG codons. The organization of tRNAs was variable across the six mitochondrial genomes sequenced here. All mitogenomes are composed of four major segments: two coding regions and two major unassigned regions (Fig. 1).

There is little variation in length of coding regions and great variation in URs (Fig. 2). These newly sequenced complete mitogenomes were deposited in GenBank (Accession numbers: OM807131–OM807136).

### Characterization of unassigned regions in mitogenomes

All available mitogenomes of *S. broughtonii*, *S. kagoshimensis* and *S. cornea* were characterized by large unassigned regions separated into two principal blocks. The first block (UR1) was located between *cox2* and *nad6*, and the second block (UR2) was located between *nad2* and *cox1*. We refer to these as “shared large unassigned regions”. An assessment of shared URs in nine mitogenomes (Table 2) revealed that the length of UR1 is relatively stable between 9831 and 10,120 bp except in *S. cornea* where it was 14,034 bp. UR2 length was highly variable from 15,167 to 28,331 bp, and the intergenic DNA (URs between genes in coding blocks) ranged from 1507 to 2925 bp. Overall, the total length of URs was 28,844 to 41,101 bp, accounting for 65.1–73.2% of these nine mitogenomes. Tandem repeats in URs showed significant variation in number (2–26 copies) and sequence length (1–273 nt) (Additional file 1: Table S3). The total length of tandem repeats varied from 759 bp to 4942 bp and takes up only 2–15% of URs. In addition, an examination of all Arcidae complete mitogenomes showed that unassigned regions (i.e., repeats, ORFs) were highly variable and responsible for expansion and variation in Arcidae mitogenomes (Additional file 1: Table S4). In comparison, a higher proportion of unassigned sequences was observed in the ark shell of *Scapharca* (>60%) and *Tegillarca* (>50%), both exceeding 16 kbp, and thus there was a strong positive correlation between mitogenome size and proportion of unassigned region.

### Novel ORFs in mitochondrial unassigned regions

TransDecoder (<https://github.com/TransDecoder/TransDecoder/wiki>) predicted eleven ORFs in mitochondrial unassigned regions that might code proteins from the *S. broughtonii* transcriptome. The analysis of mtDNA transcriptome expression (Fig. 3) showed that 12 mitochondrial coding genes have higher transcription level than all ORFs but ORF8 and ORF21

**Table 1** Mitochondrial (mt) genomes analyzed in this study, including newly assembled mitogenomes and those from Genbank

| Species                            | Subfamily        | Length (bp) | SRA              | Locality                 |
|------------------------------------|------------------|-------------|------------------|--------------------------|
| New mt genomes                     |                  |             |                  |                          |
| <i>Scapharca broughtonii</i> (1)   | Anadarinae       | 44,333      |                  | Qingdao, Shandong, China |
| <i>Scapharca broughtonii</i> (2)   | Anadarinae       | 44,327      |                  | Qingdao, Shandong, China |
| <i>Scapharca broughtonii</i> (3)   | Anadarinae       | 46,191      |                  | Qingdao, Shandong, China |
| <i>Scapharca broughtonii</i> (4)   | Anadarinae       | 48,560      |                  | Qingdao, Shandong, China |
| <i>Scapharca cornea</i>            | Anadarinae       | 46,362      |                  | Philippines              |
| <i>Scapharca kagoshimensis</i> (1) | Anadarinae       | 54,157      |                  | Qingdao, Shandong, China |
| Species                            | Family/Subfamily | Length (bp) | GenBank acc. no. | Publication              |
| GenBank mt genomes                 |                  |             |                  |                          |
| <i>Anadara crebricostata</i>       | Anadarinae       | 36,671      | MN316632         | Kong et al., [12]        |
| <i>Anadara transversa</i>          | Anadarinae       | 18,780      | MN326817         | Kong et al., [12]        |
| <i>Anadara vellicata</i>           | Anadarinae       | 34,147      | KP954700         | Sun et al., [42]         |
| <i>Lunarca ovalis</i>              | Anadarinae       | 19,620      | MN366010         | Kong et al., [12]        |
| <i>Potiarca pilula</i>             | Anadarinae       | 28,386      | KU975162         | Sun et al., [43]         |
| <i>Scapharca broughtonii</i> (5)   | Anadarinae       | 48,161      | KF667521         | Hou et al., [14]         |
| <i>Scapharca broughtonii</i> (6)   | Anadarinae       | 46,985      | AB729113         | Liu et al., [13]         |
| <i>Scapharca globosa</i>           | Anadarinae       | 33,405      | MN366011         | Kong et al., [12]        |
| <i>Scapharca gubernaculum</i>      | Anadarinae       | 45,697      | MN061840         | Sun et al., [15]         |
| <i>Scapharca inaequalis</i>        | Anadarinae       | 45,859      | MN366012         | Kong et al., [12]        |
| <i>Scapharca kagoshimensis</i> (2) | Anadarinae       | 56,170      | MN366013         | Kong et al., [12]        |
| <i>Scapharca kagoshimensis</i> (3) | Anadarinae       | 46,713      | KF750628         | Sun et al., [18]         |
| <i>Tegillarca</i> sp.              | Anadarinae       | 50,104      | MN366016         | Kong et al., [12]        |
| <i>Tegillarca granosa</i>          | Anadarinae       | 31,589      | KJ607173         | Sun et al., [41]         |
| <i>Tegillarca nodifera</i>         | Anadarinae       | 38,672      | MN366014         | Kong et al., [12]        |
| <i>Arca navicularis</i>            | Arcinae          | 18,004      | MN326818         | Kong et al., [12]        |
| <i>Arca zebra</i>                  | Arcinae          | 44,651      | MN366003         | Kong et al., [12]        |
| <i>Barbatia lima</i>               | Arcinae          | 17,479      | MN366005         | Kong et al., [12]        |
| <i>Barbatia virescens</i>          | Arcinae          | 24,871      | MN366006         | Kong et al., [12]        |
| <i>Trisidos semitorta</i> (1)      | Arcinae          | 19,461      | MN366015         | Kong et al., [12]        |
| <i>Trisidos semitorta</i> (2)      | Arcinae          | 19,613      | KU975161         | Sun et al., [43]         |
| <i>Cucullaea labiate</i> (1)       | Cucullaeidae     | 25,845      | KP091889         | Feng et al., [44]        |
| <i>Cucullaea labiate</i> (2)       | Cucullaeidae     | 20,481      | MN366007         | Kong et al., [12]        |
| <i>Glycymeris formosana</i>        | Glycymerididae   | 19,027      | MN366008         | Kong et al., [12]        |
| <i>Glycymeris yessoensis</i>       | Glycymerididae   | 17,903      | MN366009         | Kong et al., [12]        |
| <i>Arcopsis adamsi</i>             | Noetiidae        | 18,716      | MN366004         | Kong et al., [12]        |
| Outgroup                           |                  |             |                  |                          |
| <i>Mizuhopecten yessoensis</i>     | Pectinidae       | 20,964      | FJ595959.1       | Wu et al., [45]          |
| <i>Pinctada maxima</i>             | Pteriidae        | 16,994      | NC_018752.1      | Wu et al., [46]          |
| <i>Crassostrea gigas</i>           | Ostreidae        | 18,224      | AF177226.1       | N/A                      |

have a similar transcription level to *nad4*. *Nad6* had the highest transcription level of all PCGs and ORFs. In comparison, ORF104 and ORF127 showed a very low transcription level, which was considerably lower than other ORFs. The mapped read counts and TPM values of PCGs and ORFs have been recorded in Additional file 2. The ORFs shared the same location and order in four *S. broughtonii* mitogenomes (Additional file 1: Table S1). All were located in URs on the heavy

strand (as all standard coding genes) of *S. broughtonii* mitogenomes, including six in UR1 and five in UR2. The eleven ORFs showed remarkable conservation in all samples: their start and stop codons were the same, respectively, except ORF127 in *S. broughtonii* (1), ORF21 in *S. broughtonii* (2) and ORF8 in *S. broughtonii* (3) (Table 3). The longest length of ORFs was 1983 bp (ORF87, 660aa), the shortest length of ORFs was 238 bp (ORF21, 79aa). Notably, the ORF8 was variable

in its length (561 bp to 636 bp). Although *atp8* has not been previously reported in Arcidae, the lengths of ORF21 (238 bp) and ORF103 (244 bp) are close to that of *atp8* found in other bivalves (102–339 bp) and these ORFs may be candidates for *atp8*. Moreover, we found potential ORF duplication events in four *S. broughtonii* mitogenomes. Amino acid sequence comparisons using a combination of sequence and position similarity revealed that there are five additional ORFs that share a large degree of similarity within the eleven ORFs (Additional file 1: Table S5). Results showed that 47% amino acid identities were observed between ORF8 and ORF11, 39% for ORF8 and ORF7, 30% for ORF8 and ORF78. The highest identity (67% amino acid identity) was observed between ORF87 and ORF127. ORF104 had at least 28% amino acid identities with other three ORFs. These findings showed that ORFs have high similarity, suggesting that duplication events have occurred between them. However, no significant amino acid sequence similarity was detected with known proteins for the 14 new lineage-specific ORFs using BLAST [47] and PSI-BLAST against NRDB and UniProt (Additional file 1: Table S6).

To establish whether these ORFs were taxonomically restricted to *S. broughtonii*, or if they were an evolutionary feature of ark shell, we screened for the presence of 16 ORFs (11 predicted ORFs and 5 duplicated ORFs, see Table 3 and Additional file 1: Table S5) in 32 mitogenomes of Arcidae (Table 1) with BLAST (Additional file 1: Table S7 and Additional file 3). None of the ORFs were similar to mitochondrial PCGs. Complete ORFs and their duplications could only be annotated in 13 mitogenomes, including 6 *Scapharca* and 1 *Anadara* species (Additional file 1: Table S8 and Fig. 2), suggesting that most of the ORFs were specific to the *Scapharca* lineage (except for *S. globosa*). With one exception, ORF21 was annotated in 12 species, which had a wider distribution. All 16 ORFs were verified in the mitogenomes of *S. broughtonii* and *S. kagoshimensis* except for *S. kagoshimensis* (3). Although the mitogenome of *S. kagoshimensis* (3) have a long URs, only five complete ORFs could be annotated, which may be due to poor assembly quality of the mitogenome. The result showed that 15 ORFs were found in the mitogenome of *S. inaequalis* except for ORF 11. In the mitogenomes of *S. gubernaculum* and

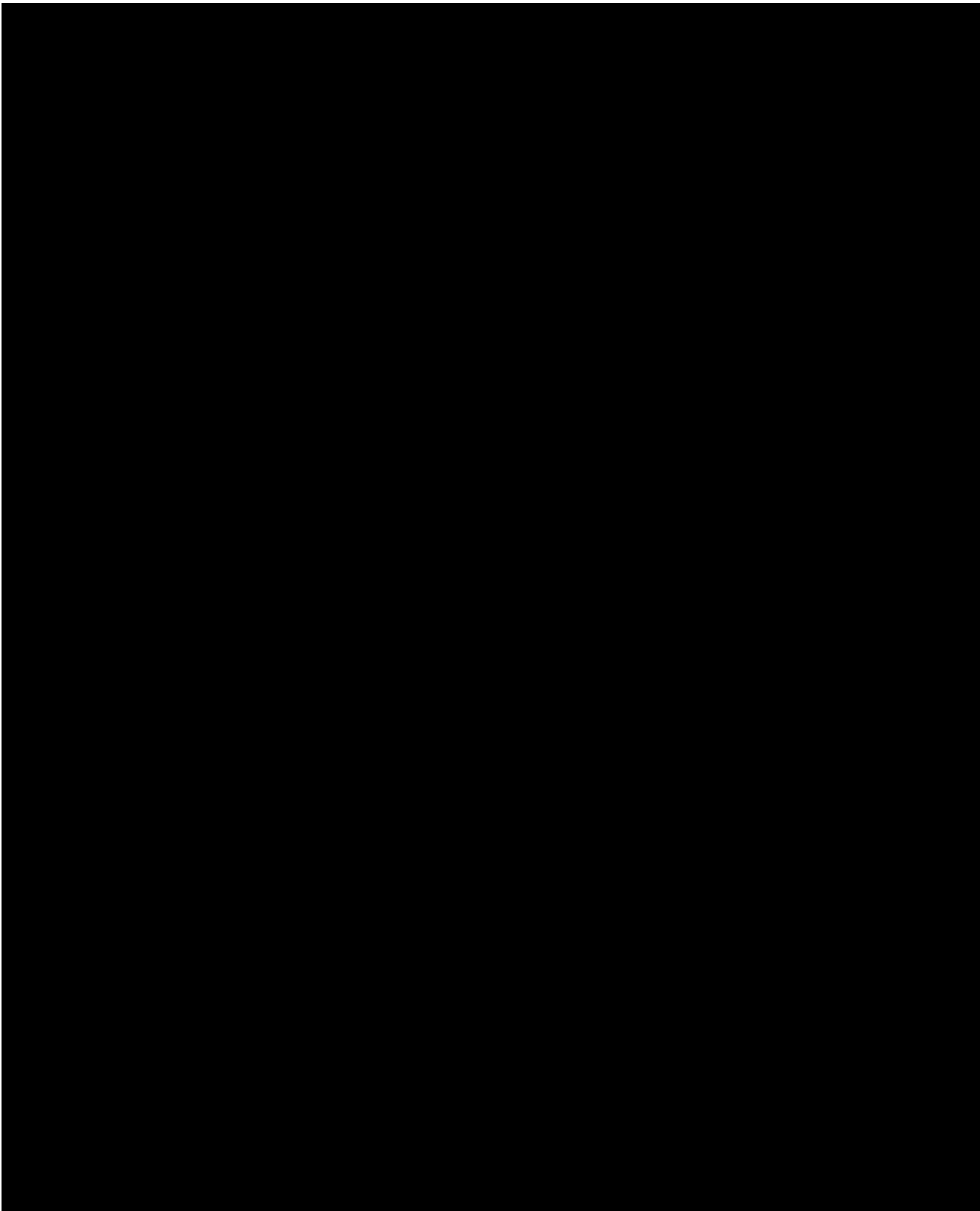
*S. cornea*, 14 ORFs were found. Only ORF8, ORF21 and ORF86 were found in *S. globosa* mitogenome. Most of ORFs were unique to the genus *Scapharca* and *A. crebri-costata*, but the partial sequence (about 120aa) of ORF87 was similar to fragments of other genera mitogenomes (Additional file 1: Table S7), which suggests that ORF87 may have had a wider distribution in Arcidae (Fig. 2). Sequence comparisons both within and between lineage-specific ORFs revealed high variability in length, but some ORF lengths were conservative such as ORF10 and ORF11 (Additional file 1: Table S8). The total length of all ORFs identified in *S. broughtonii* was accounting for 23–26% of mitogenomes and 35–40% of URs, indicating ORFs are one of the main components of the large URs. In addition, many large ORFs (>2000 bp) were found in Arcidae mitogenomes (Additional file 1: Table S9).

#### Ka/Ks analysis for putative novel Arcidae mitochondrial ORF proteins

To estimate the degree of selection (either neutral, positive, or purifying) and genetic conservation on 12 protein coding genes and 16 ORFs, the number of nonsynonymous substitutions per nonsynonymous sites (Ka) relative to the number of synonymous substitutions per synonymous sites (Ks) was calculated. According to our results (Fig. 4 and Additional file 1: Table S10), the mean value for Ka was also different between protein-coding genes and ORFs (0.0137 vs. 0.0624;  $P < 0.001$ ), suggesting that the ORFs accumulated more non-synonymous mutations. Twelve protein-coding genes and all ORFs were found to be under strong purifying selection ( $Ka/Ks < 1$ ). The Ka/Ks of protein-coding genes was between 0.0470 (*cob*) to 0.1447 (*nad4L*). A low Ka/Ks has been a common finding for mtDNA-encoded protein genes in animals and is explained by the elimination of mildly deleterious mutations. The ORFs had a wide range of Ka/Ks ranging from 0.1267 (ORF103) to 0.615042 (ORF40), but the Ka/Ks of some ORFs (ORF5, ORF103, ORF127) were close to some protein-coding genes (*nad4L*, *nad5*). The mean value of Ka/Ks in ORFs (0.2834) was significantly higher than that of protein-coding genes (0.0874) ( $P < 0.001$ ), suggesting that ORFs have been under less selective constraints than mitochondrial protein-coding genes. In addition, the level of ORF sequence

(See figure on next page.)

**Fig. 1** Maps of the six mitogenomes sequenced in this study. The corresponding species name and length are given inside each genome map. The outer ring comprises all standard and putative coding sequences, identified with the following color code: blue, genes encoding electron transport chain and ATP-synthase subunits; yellow, tRNA genes; green, rRNA genes (see Additional file 1: Table S1 for details). All the genes are encoded on the same strand. The middle ring represents GC content (dark grey). The inner ring represents scale



### Genus *Scapharca*

### Genus

conservation between mitogenomes from alignments confirmed that these ORFs have conserved regions (Additional file 4).

**Conserved secondary structures in ORF protein sequences**  
To assess whether mitochondrial ORFs possess ORFs that could have functional importance in Arcidae bivalves, we investigated the structure of these ORFs. Transmembrane (TM) helices were identified using three different programs. Twelve of 16 ORFs in *S. broughtonii* were predicted to have at least one TM-helices with both Phobius [48] and TMHMM 2.0 programs [49]. The third, TOPCONS [50], had a stricter criterion because it gives a consensus result for the protein from five different topology prediction (details see Additional file 1: Table S11). All three programs predicted two TM-helices for all ORF127 and ORF87 in different Arcidae species with 100% confidence. The number of TM-helices of other ORFs was predicted to be unstable across Arcidae species. Four or five TM-helices were found in ORF40 of different species with two software. The remaining 13 ORFs returned variable TM predictions and possessed one to four predicted transmembrane domains in different mitogenomes. In addition, signal peptide (SP) was found in the N-terminus

of five ORFs (ORF40 in *S. cornea* and *S. gubernaculum*, ORF5 in *S. broughtonii* and *S. kagoshimensis*, ORF7 in *S. kagoshimensis* (1), ORF86 in *A. crebricostata*, ORF86-b in *S. inaequivalvis*, ORF87 in *S. cornea* and *S. gubernaculum*) (Additional file 1: Table S11). Notably, ORF40 of *S. cornea* and *S. gubernaculum* were predicted to have a SP and five TM-helices, and SPs are located 1–17 aa from the N'end of the predicted peptide, with a cleavage site between 17 and 18 aa. Eleven of 14 ORF5 in different mitogenomes were predicted to have a SP, and they are almost in the same position. A SP were found only in ORF7 of *S. kagoshimensis* (1), but not in other Arcidae mitogenomes. The remaining 11 ORFs were not predicted to have a SP. Moreover, some ORFs (e.g., ORF21 in *S. broughtonii* (2) (4) (5) (6), *S. cornea* and *S. gubernaculum*) were predicted to have a Rossmann fold, a tertiary fold found in proteins that bind nucleotides, such as enzyme cofactors FAD, NAD<sup>+</sup>, and NADP<sup>+</sup> (Additional file 1: Table S12).

### Function prediction of ORF protein sequences

As these ORF sequences did not show any obvious homology with known proteins, we performed an in-depth comparative analysis using multiple programs to

**Table 2** Detailed information of unassigned regions of Arcidae mitogenomes. The unit of length is the bp

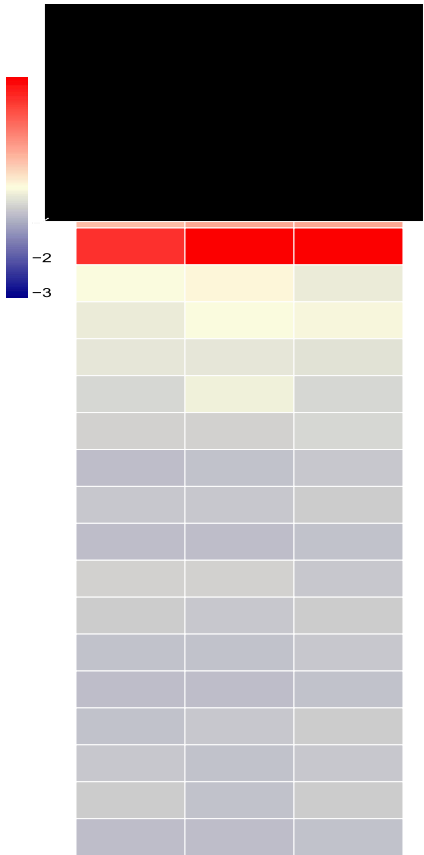
| Species                            | Mitogenome size | The total length of URs | Proportion of URs | UR1 location  | UR1 size | Proportion of UR1 | UR2 location          | UR2 size | Proportion of UR2 |
|------------------------------------|-----------------|-------------------------|-------------------|---------------|----------|-------------------|-----------------------|----------|-------------------|
| <i>Scapharca broughtonii</i> (1)   | 44,333          | 28,908                  | 65.21%            | 18,057–27,994 | 9938     | 22.42%            | 1–861436,903–44,333   | 16,045   | 36.19%            |
| <i>Scapharca broughtonii</i> (2)   | 44,327          | 28,844                  | 65.07%            | 17,914–27,851 | 9938     | 22.42%            | 1–847136,760–44,327   | 16,039   | 36.18%            |
| <i>Scapharca broughtonii</i> (3)   | 46,191          | 30,806                  | 66.69%            | 18,033–27,968 | 9936     | 21.51%            | 1–859036,753–46,191   | 18,029   | 39.03%            |
| <i>Scapharca broughtonii</i> (4)   | 48,560          | 32,984                  | 67.92%            | 25,845–35,677 | 9833     | 20.25%            | 1–16,53244,063–48,560 | 21,030   | 43.31%            |
| <i>Scapharca broughtonii</i> (5)   | 48,161          | 33,004                  | 68.53%            | 9314–19,144   | 9831     | 20.41%            | 27,798–48,161         | 20,364   | 42.28%            |
| <i>Scapharca broughtonii</i> (6)   | 46,985          | 31,809                  | 67.70%            | 9314–19,146   | 9833     | 20.93%            | 27,673–46,985         | 19,313   | 41.11%            |
| <i>Scapharca kagoshimensis</i> (1) | 54,157          | 38,394                  | 70.89%            | 38,928–48,778 | 9851     | 25.66%            | 2805–29,551           | 26,747   | 69.67%            |
| <i>Scapharca kagoshimensis</i> (2) | 56,170          | 41,101                  | 73.17%            | 30,713–40,833 | 10,120   | 18.02%            | 1–21,32949,169–56,170 | 28,331   | 50.44%            |
| <i>Scapharca kagoshimensis</i> (3) | 46,713          | 32,982                  | 70.61%            | 9395–19,345   | 9951     | 21.30%            | 28,023–46,713         | 18,691   | 40.01%            |
| <i>Scapharca cornea</i>            | 46,362          | 30,708                  | 66.24%            | 22,735–36,777 | 14,034   | 30.27%            | 1–13,57144,767–46,362 | 15,167   | 32.71%            |
| <i>Scapharca gubernaculum</i>      | 45,697          | 30,079                  | 65.82%            | 9164–21,106   | 11,943   | 26.14%            | 29,095–45,697         | 16,603   | 36.33%            |
| <i>Scapharca globosa</i>           | 33,405          | 18,631                  | 55.77%            | 14,724–17,921 | 3198     | 9.57%             | 1–437226,555–33,405   | 11,223   | 33.60%            |
| <i>Anadara crebricostata</i>       | 36,671          | 21,991                  | 59.97%            | 23,496–27,840 | 4345     | 11.85%            | 1–593336,253–36,671   | 6352     | 17.32%            |
| <i>Anadara vellicata</i>           | 34,147          | 20,659                  | 60.50%            | 9130–9424     | 295      | 0.86%             | 18,084–34,147         | 16,064   | 47.04%            |

predict the function of ark shell mitochondrial ORFans. Results obtained for all ORFans with all programs for protein function prediction were summarized in Additional file 1: Table S13, which included the most frequent categories of hits for molecular functions, biological processes and cellular components for the mitochondrial ORFans [51, 52], and detailed motifs and domains information (HHpred, [53]; I-TASSER). Overall, the most common hits for all ORFs were proteins involved in oxidoreductase activity, nucleic acid or protein binding (e.g., helicase/hydrolase activity) and metal ion binding (e.g., nickel cation/cobalt ion binding). Some other hits were proteins with membrane association and transporter activity, for example involved in transport

across membrane, establishment of protein localization, and most of all involved in intracellular transport (e.g., ORF10, ORF7, ORF104, ORF106, ORF21). Some ORF proteins pointed to a role in ATP binding, for example in cellular macromolecule metabolic process (e.g., ORF127).

In particular, most sequences analyzed returned predictions that the proteins were involved in oxidoreductase activity and metabolic process, and the predicted subcellular localizations for these ORFs were different, with some being membranes and organelles (endoplasmic reticulum and nucleus) and some being soluble outside the cell (Additional file 1: Table S14). For ORF104, the highest probability matches included proteins that have a role in obsolete oxidoreductase activity, acting on



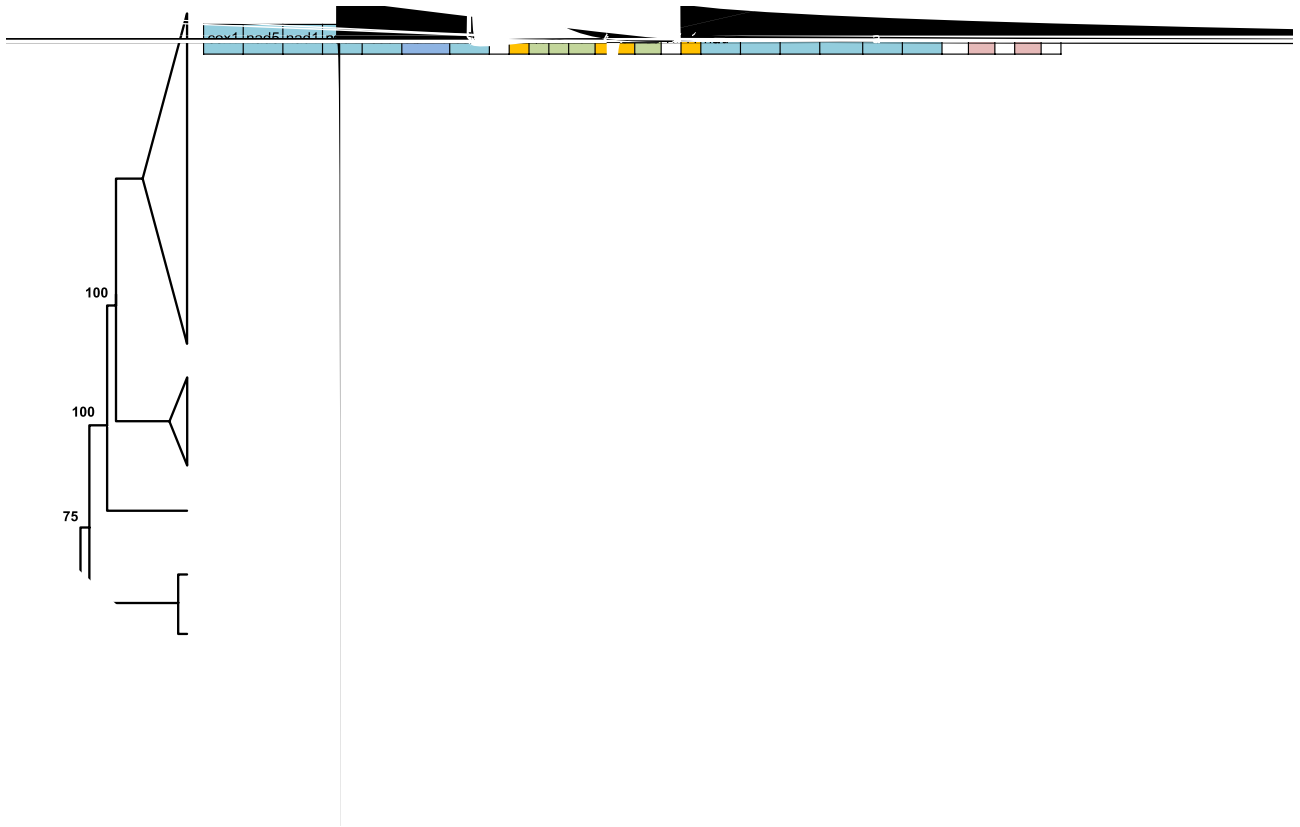


... oxygen as acceptor and obsolete cytochrome oxidase activity. In addition to these hits also included proteins related to nucleic acid binding and oxidase activity participate in cellular processes such as response to stress and nucleic acid metabolic processes. Additionally, several hits of ORF11 appeared to have glucan 1,4-alpha-glucosidase activity (a step in the tricarboxylic acid cycle), obsolete metabolic process and drug metabolic process. ORF48 pointed to a role in cellular respiration (e.g., one-carbon metabolic process (e.g., ORF49) and metabolic process (e.g., ORF5). However, the previous study [38] proposed a viral mitochondrial ORFans in DUI bivalves, therefore scanned our results for protein function predicted with all programs to highlight the hits related to biological process and proteins (Additional file S13). However, BLAST results for all ORFs

(Additional file 1: Table S6) showed that most of the ORF hits are non-viral and viral-related hits have low probability (e.g., ORF7, E-value = 2.2). Hits with high probability values were bacterial or metazoan proteins (e.g., ORF40, Hypothetical protein, *Sepia pharaonic*, ORF87, MCP signaling domain protein, *Clostridium argentinense*). The same ORF in different species did not produce consistent blast result.

#### Phylogenetic analyses and ancestral state reconstruction

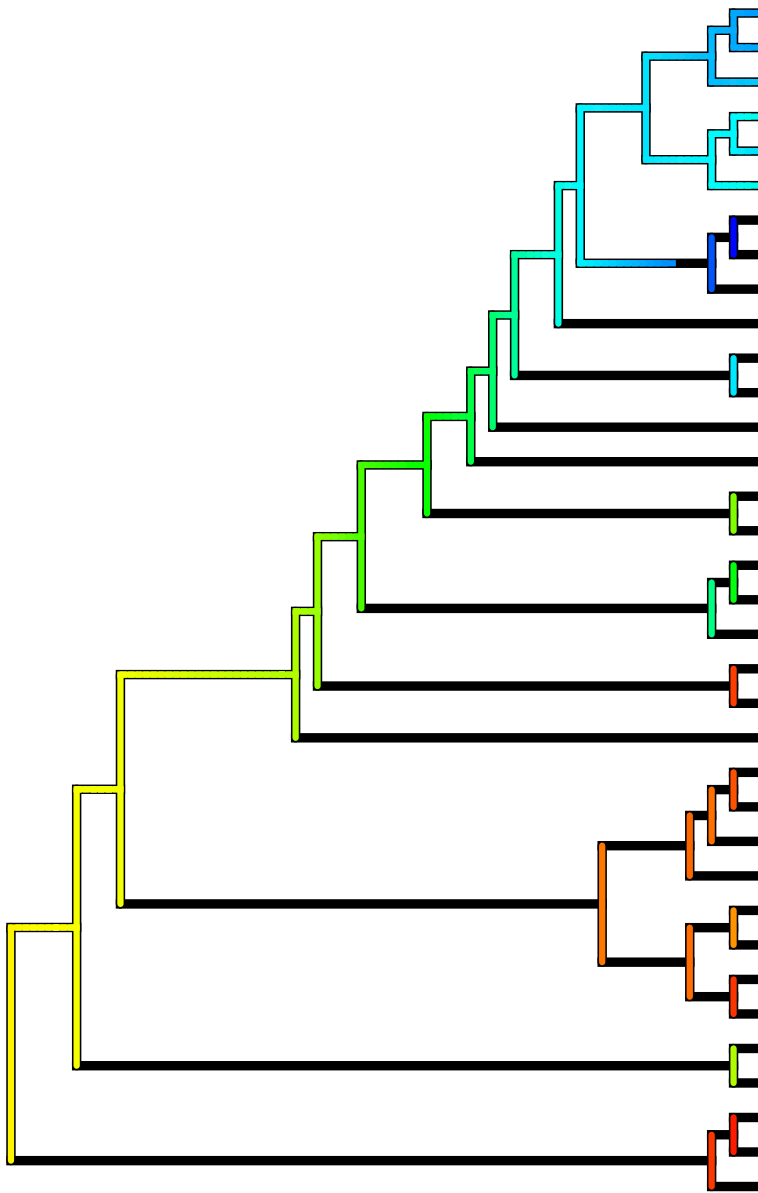
The molecular phylogeny of Arcoidea was reconstructed based on the mitogenome data sets using ML (Figs. 5 and 6). After removing ambiguously aligned positions, the concatenated alignment of amino acid sequences from thirty-five taxa had a total length of 3057 positions (Table 1). Arcidae was found to be polyphyletic with three well-supported lineages. The first lineage included the subfamily Anadarinae and the sister taxon *Barbatia lima*; the second comprised two *Trisidos* species and *Barbatia virescens*; and two *Arca* species formed the



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#### Novel ORFs in Arcidae mitogenomes and their possible origins

Mitogenomes of Arcidae species with large URs have novel lineage-specific ORFs which do not show significant amino acid sequence similarity to known proteins in the database, but they have special secondary structures and meaningful hits for function prediction. The ORF

amino acid sequences are conserved among different mitogenomes and found in extra-genic regions, always inside the large URs. Traditionally, URs are generally regarded as vestiges of pseudogenes generated by random deletions after gene duplication [34, 57–59], while our further examination of the two shared large URs in Arcidae reveals two categories of sequences. The first

category contains sequences exhibiting many features typically associated with mitochondrial control regions, such as presence of repeat units and sequences that can form hairpin structures and stem–loop. The second category contains sequences that possess many ORFs of considerable length. Previous research [12] on the first category found significantly positive correlations between Arcidae mitochondrial genome size and proportion of tandem repeats and proportion of inverted repeats. However, the ORFs identified turned out to be one of the main components of the large URs rather than repeat families and transposable elements in our study. Some ORFs were conserved in the lineage containing *Scapharca* and *A. crebricostata* (Fig. 2), suggesting that they may have emerged before the speciation of *Scapharca* lineage and *A. crebricostata*. Mitochondrial URs of some ark shells are probably inserted in multiple independent evolutionary events [12], and if so these lineage-specific ORFs may arise from independent insertion events.

Previous studies [35, 38–40, 60] suggested mitochondrial ORFs could originate from different processes: (1) the duplication and subsequent modification of extant mitochondrial genes, (2) transfer of DNA from nuclear genomes to mitochondrion, (3) the insertion of viral sequences into the host mitogenome. In our study, the ORF sequences analyzed do not show homology with any known Arcidae mitochondrial protein, so they unlikely originated from recent duplication events from existing mtDNA genes. The ORF sequences we identified in *S. broughtonii* were not found in a high-quality nuclear genome of *S. broughtonii* [61] (Additional file 5), indicating they may not have been transferred from nuclear genomes. Moreover, in our analysis, hits similar to ORF sequences are mainly from proteins of bacteria, fungi, parasites and viruses (Additional file 1: Table S6 and Table S13). Since most of hits have no significant similarity or probability, they are not sufficient to determine the origin of the ORFs. The results showed that 9 of 16 ORFs in *S. broughtonii* are possibly involved in the process of viral entry into host cell and viral release from host cell, which provide clues to speculate on a possible viral origin. However, BLAST results including all Arcidae species showed that most of the ORF hits are to non-viral sequences, and the probability of some viral hits (< 60%) were low. Hits with high probability values (E-values < 0.005) are bacterial or metazoan proteins, suggesting that other organisms or other processes [62] may be the source of these ORF genes. Consequently, our results do not support a viral origin of Arcidae ORFs, although previous studies [38, 40] have suggested that some bivalve mitochondrial ORFs may originate from viruses.

In Arcidae mitogenomes, if ORFs have experienced rapid evolution since their origin, they may diverge to

the extent that homology to mitochondrial proteins, nuclear sequences and viral sequences is not discernable.

The ka/ks results showed that some ORF sequences contained mostly non-synonymous mutations (Additional file 1: Table S10), indicating the rapid evolution of these mitochondrial ORF genes. Fast rate of evolution may erase evidence of ORF sequence similarities (homology) among species, so we cannot fully exclude the possibility that the ORFs are derived from mitochondrial duplications, nuclear genome or even viral sequences. Of these possible origins, mitochondrial duplication interpretation is more plausible. Gene duplication is thought to be the most common mechanism underlying the origin of most novel genes [63]. In our study, large-scale duplication events (both mitochondrial genes and ORFs) were found, suggesting that duplications are prone to occur in Arcidae mitogenome. Also, this explains why many identified ORFs have TM-helices and the most common hits of them involved in metabolic process. In considering possible alternatives, these ORFs may originate from duplication of mitochondrial genes, but also call for further studies to investigate their origin.

#### The insertion and duplication of mitochondrial novel ORFs and implications for UR size evolution

According to ancestral state reconstruction (Fig. 6), the mitogenome size of the common ancestor of Arcidae is relatively small (i.e., < 20 kb) like in most metazoans. This result suggests Arcidae mitogenomes have experienced multiple expansions (insertion and duplication events) with some lineages forming very large mitogenomes. For example, UR1 (regions between *cox2* and *nad6*) is only 295 bp in the mitogenome of *A. crebricostata* (Table 2). However, we found that the UR1 of *S. globosa* mitogenome is 3197 bp, which contains the ORF sequences similar to ORF86 and ORF8. Based on phylogenetic relationships and mitogenome structure of Arcidae (Fig. 5), the inserted sequence may be the origin of the UR1 of *Scapharca* lineage and *A. crebricostata*. Subsequently, the expansion of UR1 is obvious in other *Scapharca* species (reaching above 10 kbp) (Table 2) because of the duplications of some ORFs (Fig. 5). In bivalve, many mitochondrial gene duplication events have been found such as *nad2* duplication in the oyster genus *Crassostrea* (Bivalvia, Ostreidae) [46] and *cox2* duplication in several DUI bivalve species [64, 65]. In our results, *S. broughtonii* mitogenomes contained a duplication of the *cox2* gene, named *cox2-b* (Fig. 5 and Additional file 1: Table S2), a feature that has been also observed in the mitogenome of other closely related species (*Scapharca* lineage and *A. crebricostata*, except for *S. globosa*). Coincidentally, ORF duplications were found in the *Scapharca* species that have *cox2* duplication with exception of *S. globosa*.

For example, according to ORF distribution and phylogenetic relationship (Fig. 5 and Additional file 6), ORF8, ORF7, ORF11 and ORF78 (>30% identity at the protein level) might originate from the same ORF gene and the ORF7 gene in *A. crebricostata* could have been the original copy. In addition, duplication of ORF5, ORF21, ORF104, ORF46, ORF86 and ORF87 were also found in different Arcidae mitogenomes, which suggest that a large-scale duplication event has occurred rather than just *cox2* duplication. Meanwhile, these duplicated ORF sequences make mitogenomes larger. Most of the ORFs from duplication were over 500 bp in length, which adds at least 8217 bp to the mitogenomes of *S. broughtonii*. For example, ORF87 and its duplication of ORF127 are both over 1800 bp, which have a huge impact on mitogenome size. In conclusion, the duplication of *cox2* and ORFs cause Arcidae mitogenomes to expand further. We think that duplication events of ORFs play an important role in multiple subsequent expansions of Arcidae mitogenome.

However, most of ORFs we identified in *S. broughtonii* are only found in *Scapharca* and *Anadara* (Fig. 2). Possible explanation is that the expansion of *Scapharca* and *Anadara* was the result of independent insertion and duplication events, which results in the ORF sequences different from other genera. To further explore expansions caused by ORFs, we reinvestigated all Arcidae mitogenomes and found 54 large ORFs (LORFs) of unknown structure and function in 18 Arcidae mitogenomes (Additional file 1: Table S9). These LORFs range in length from 933 to 5187 bp, and the average length is 2200 bp. In the mitogenomes of *Scapharca*, all LORFs were found in UR2, indicating that the emergence of LORFs may be related to UR2 expansion. Interestingly, many LORFs were also found in *Anadara* and *Tegillarca* mitogenomes, and some of them have a fragment of amino acid sequences similar to the regions of ORF87 in *Scapharca* (Fig. 2 and Additional file 1: Table S7), which implies that these LORFs might have a common origin with ORF87 in an earlier expansion event. Because of large lengths, LORFs were easily disrupted by mutations and harder to maintain. Therefore, we think that the existence of LORFs might be significant and can provide clues to explore the independent expansions of other Arcidae mitogenomes in future.

#### Predicted functions for ark shell mitochondrial ORFs

In our study, there are multiple lines of evidence indicating potential functionality of 14 novel lineage-specific ORFs in *S. broughtonii*, likely as expressed proteins. In other taxa, lineage-specific ORF genes are involved in important adaptive processes and key biological functions [36, 38]. Herein, multiple lines of evidence suggest these UR ORFs are functional in nature. Transcriptome

analysis indicated that the nine ORFs (Table 3 and Fig. 3) are transcribed in mitogenomes of *S. broughtonii*. Secondly, the *ka/ks* analysis showed that all the ORFs are under purifying selection (Fig. 4 and Additional file 1: Table S10). Thirdly, because novel ORFs do not show significant similarity to known proteins, we performed multiple analyses of their structure to predict the function. Our results for secondary structure prediction show that most of the ORFs are predicted to have functional domains (Additional file 1: Table S11), which is a significant support for identifying these ORFs as protein-coding genes. For example, we observe that four conserved TM-helices are present in ORF40 proteins of *S. broughtonii*. In addition, ORF40 returned hits to proteins with membrane association (e.g., proteins involved in tail-anchored membrane protein insertion into ER membrane), and the predicted subcellular localizations with DeepLoc for ORF40 are also endoplasmic reticulum and membrane.

The ORF127 protein is very long in length and has two stable TM-helices among different Arcidae species. Our prediction results, together with transcriptome and *ka/ks* analysis, showed that these lineage-specific ORF proteins that occur in Arcidae mitogenomes may have underlying functions. For example, ORF21 and ORF103 may be candidates for *atp8*, which has not been annotated in many bivalves [6] and is not reported in Arcidae. The two ORFs are approximately the same length as *atp8* and are transcribed in *S. broughtonii*. They are under strong purifying selection and their relative solvent accessibility is similar to the *atp8* gene (Additional file 7). ORF21 is conserved at the amino acid level and has a broader distribution than ORF103 (Fig. 2). ORF103 is predicted to have TM-helices, but ORF21 does not. Compared to ORF21, we believe that ORF103 is a more likely candidate for *atp8* because the TM-helix is important for *atp8* function. However, ORF21 and ORF103 do not show homology with any bivalve *atp8* genes. Therefore, we cannot determine if either of these two ORFs is an *atp8* gene homolog.

Moreover, based on the functional prediction analysis, we speculated that some of novel ORF proteins in mitochondrial URs of Arcidae may have acquired new functions. A previous study showed that ORFans in bivalves with DUI may have a viral origin and be involved in the maintenance of sperm mitochondria during embryo development [38, 40]. Our results for molecular function prediction show that the ORFs have different functions hits, the most common hits involved in metabolic process (e.g., ATP association activity, oxidoreductase activity, glucan 1,4- $\alpha$ -glucosidase activity). Several Arcidae species are limited locomotive and more tolerant to hypoxia, such as *S. kagoshimensis*, which lead to low metabolic rate [43, 66, 67]. The large mitogenome size

in these bivalves may be correlated with their metabolic rates because the relaxed selective constraints of mitogenomes may be energy-related [12, 68]. In addition, genes participating in ATP and lipid metabolism under selection were found to be important in thermal adaptation for oysters [69]. Lineage-specific genes could participate more in lineage-specific adaptation [70], therefore, we speculated that the functions of lineage-specific ORF genes are related to low metabolic rate or thermal adaptation, which may provide new insights into the function of large URs in Arcidae mitogenomes.

Finally, the possibility that ORFs may be pseudogenes and do not perform any function cannot be ruled out. According to the results, some ORFs are unstable in the number of TM-helices such as ORF104, ORF106 and ORF46. The number of TM-helices in a given ORF vary among Arcidae species, indicating that these ORFs are not conserved at the secondary structure level. These results suggest a possibility that ORFs may have different adaptations in various Arcidae species. Another possibility is that the ORFs could be pseudogenes or in the process of pseudogenization after insertion and duplication events. In many cases duplicated genes are subject to pseudogenization [1], which appears to be the most likely fate for mitochondrial gene duplications. But compared to pseudogenes, the ORFs we identified are more conserved at the nucleotide sequence level and some of them may be transcriptionally active, so they are likely to be in the process of pseudogenization but more research is needed.

#### Dynamic changes in intraspecific mitogenome size

Different mitogenomes in the same Arcidae species vary dramatically in length. The previous studies have demonstrated that tandem repeats are potentially a main factor leading to variation of intraspecific mitogenome size [12, 16]. For example, mitogenomes of *S. broughtonii* ranges in size from about 47 kb to ~ 50 kb due to variation in the number of tandem repeats [13]. The four *S. broughtonii* mitogenomes we assembled also have many tandem repeats (Additional file 1: Table S3). However, variation of these tandem repeat size does not fully explain the variation (44,327–48,560 bp) of the URs of *S. broughtonii* mitogenomes because the latter is larger. By observing URs length (Table 2), UR1 in *S. broughtonii* and *S. kagoshimensis* are almost the same length, and UR2 is highly variable and responsible for the variation of mitogenomes in length. We propose that variation of the length of *S. broughtonii* and *S. kagoshimensis* mitogenomes may be caused by incomplete assembly of the UR2 because of their complex content. Large-scale repeat sequences are difficult to sequence using conventional Sanger and short-read sequencing methods

[71]. Theoretically, repeats extend beyond read length, mitogenome assemblies are limited within the boundaries of repetitive elements [72]. Although long-range PCR can be used to amplify DNA regions of several kilobases, sequencing through repetitive regions often results in ambiguous and/or erroneous sequence reads as a consequence of self-priming of randomly-amplified repeat-segments, chimeras and/or jumping PCR artefacts [73, 74]. For example, the *S. kagoshimensis* mitogenome (KF750628) were assembled using long-PCR into a circle [18], but some protein-coding genes and ORFs are fragmented, which suggest a low-quality assembly. Hence, future studies can focus on long-read sequencing for Arcidae mitogenome assembly, which can achieve read lengths of 80 kb to >1 Mb, enabling repetitive and structurally complex DNA elements to be resolved with confidence [75, 76].

#### Conclusions

In this study, we found 14 special ORFs in the large unassigned regions in mitogenomes of *S. broughtonii*. Interestingly, these putative additional proteins have also been found in other species of genus *Scapharca*. We present multiple lines of evidence supporting the functionality of ORFs at amino acid and nucleotide levels and discuss their possible origin. These results support novel features for Arcidae mitochondrial genomes: the presence of lineage-specific mitochondrial ORFs with transcriptional activity and potential functional significance. Moreover, our study reveals that the insertion and duplication events of ORFs play an important role in multiple expansions of Arcidae mitogenome. Although other bilaterian taxa have expansion regions in their mitochondrial genome, those in Arcidae are most extreme. Thus, Arcidae may provide significant insights on how bilaterian mitochondrial genomes evolve in terms of size variation, gene complement, and gene organization.

#### Methods

##### Specimen collection and sequencing

Adult *S. broughtonii* specimens were obtained from populations near Qingdao (the tuandao market), Shandong Province, China. One specimen of *S. cornea*, was collected from a local market in the Philippines. After collection, all specimens were immediately preserved at  $-80^{\circ}\text{C}$  or in 95% ethanol. Species were identified using morphology and genetic distance of Arcidae mitochondrial *Cox1*. Total genomic DNA of four *S. broughtonii* and one *S. cornea* individuals was extracted from adductor muscle using the TIANamp Marine Animals DNA Kit according to the manufacturer's instructions and sequenced on an Illumina HiSeq X using  $2 \times 150$  paired-end (PE) library. Total mRNA extraction of six *S.*

*broughtonii* individuals was performed from adductor muscle using a Trizol protocol. All procedures were carried out on ice and quickly to avoid RNA degeneration.

The extracted mRNA was sequenced with a paired-end (PE) library with an insert size of 250 bp. The sequencing of genomic DNA and RNA was both performed by Tianjin Novogene Bioinformatics Technology Co., Ltd., China.

#### Mitochondrial genome assemblies and annotation

Raw data from four *S. broughtonii*, one *S. kagoshimensis* (SRX8857271) and one *S. cornea* individuals were filtered using Trimmomatic 0.39 [77] for removal of TruSeq adapter sequence and trimming low-quality bases from the ends of each read. Clean short reads were assembled de novo using Novoplasty 4.0 [78], Mitoz v2.3 [79] with the all module and SPAdes v3.11.1 [80] with k-mer of 21, 33, 55, 77 and the `-careful` flag, respectively.

Then, assembly results were searched using BLAST [47] against a nucleotide database constructed from the complete mitogenome of *S. broughtonii* (AB729113) to find the mitochondrial contigs. Some partial mtDNA contigs were recovered in the assembly results of SPAdes. In order to bridge contigs together into a single contig, Price1.0 [81] was applied to extend and join these partial contigs with default settings by iteratively adding sequence reads to the contig ends. Finally, mitogenomes from different assemblers were assessed using Quast 5.0 [82] based on genome fraction, total aligned length, duplication ratio, and level of completeness. The above programs were installed on Linux system via Bioconda [83]. All the newly sequenced mitogenome sequences have already been deposited in GenBank, and the accession numbers are listed in Table 1.

Locations of the protein-coding genes (PCGs), transfer RNAs (tRNAs) and ribosomal RNAs (rRNAs) were determined initially with the MITOS web server using the invertebrate mitochondrial genetic code and validated using MFannot [84]. ORF Finder (<https://www.ncbi.nlm.nih.gov/orfinder/>) and BLAST were employed to examine and adjust manually gene boundaries by comparison with the published Arcidae mitochondrial genes since MITOS seems to underestimate gene length.

The visualization of mitogenomes was performed using CGView [85]. Large URs structure was defined using blastn with manual alignments. ORFs in large URs of six mitogenomes were identified with ORF Finder using the invertebrate mitochondrial genetic code.

#### Transcriptome analysis

Quality assessment of RNA reads from six *S. broughtonii* individuals was carried out using FastQC v0.11.9 [86].

Trimmomatic 0.39 was employed for removal of adaptor sequences and low-quality positions. Then HISAT2 [87] was used to align clean reads from each individual to the reference mitogenomes that had been newly assembled.

The mapped reads were sorted, indexed using Samtools [88] and were assembled using Stringtie [89] in a reference-based approach. Next, we used BLAST to search the StringTie gene sets against the database of UniProtKB/Swiss-Prot proteins [90], and identified the protein domain with PFAM [91]. TransDecoder v5.5.0 was used to predict ORFs (>20 aa) that had coding potential from assembled transcripts. The results from BLASTP and hmmscan [92] searches were used to inform the final TransDecoder prediction step.

To estimate expression levels, we summarized information on annotated mitochondrial PCGs and predicted ORFs and used Artemis [93] to make a GTF file (*S. broughtonii* (4) as a reference mitogenome). FeatureCounts v2.0.1 [94] was used to count mapped reads on BAM files with the option "`--primary, -t exon, -g gene_id`". All other parameter values were left at their defaults. BAM files were from the HISAT2 results of three *S. broughtonii* transcriptomes, which are from the same experimental batch. Finally, we used a R pipeline (Additional file 2) to estimate expression levels in TPM for each mitochondrial PCGs and ORFs. Heatmap and boxplot were made with R package pheatmap and ggplot2 [95].

#### Novel mitochondrial ORFs and sequence conservation

To assess the presence of novel ORFs in other Arcidae mitogenomes in which they were not annotated, and at the same time validate annotations, we used BLAST to align with E-values < 0.00001 and ORF Finder to search for all possible ORFs  $\geq 75$  nucleotides long under the invertebrate mitochondrial genetic code from the 32 Arcidae mitogenomes, and then translated them into the corresponding proteins. The duplicated ORFs were classified based on BLAST results and molecular phylogeny. Because ORF protein sequences vary little within the same species, only one sequence was used for analyzing. For comparative purposes, alignments of the putative novel mitochondrial proteins in different Arcidae species were run with MAFFT v7.475 [96]. Finally, we used Jalview [97] to visualize the alignments. Then we used a ML method implemented in KaKs\_Calculator v2.0 [98] to estimate ratios of nonsynonymous and synonymous substitution rates (Ka/Ks) in mitochondrial PCGs and ORFs between sister pairs of *S. broughtonii*. These comparisons facilitated a better understanding of the selection pressures acting on protein coding genes and ORFs. ORF names were given according to the result of ORF Finder in *S. broughtonii*.



### Protein structural and functional analysis

The above-mentioned ORFs were translated and analyzed at the amino acid level. Putative transmembrane (TM) helices were identified using a variety of protein signature recognition methods implemented by the following programs: Phobius [48], TMHMM 2.0 [49] and TOPCONS [50]. Evidence of signal peptides (SPs) was sought using Phobius, TOPCONS, SignalP 4.0 [99] and TargetP 2.0 [100]. HHpred [53] were used to search for known functional sequence motifs and domains. Cofactory 1.0 [101] was used to identify Rossmann fold sequence domains and predicts their specificity for the cofactors FAD, NAD or NADP. Subcellular localizations (e.g., cell membrane, cytoplasm, nucleus, etc.) were predicted using DeepLoc-1.0 [102] and Euk-mPloc 2.0 (Cell-Ploc 2.0 package) [103].

The following procedures were used to predict the function of ORF proteins: (1) we performed BLASTp, tBLASTx, and PSI-BLAST searches against NCBI entire non-redundant protein database (NRDB) with default parameters [47] and BLAST searches against UniProt (UniProtKB reference proteomes + Swiss-Prot) with default parameters. (2) we used HHpred for profile HMM – profile HMM comparisons, which compares HMM profiles with databases of HMMs representing proteins with annotated protein families (e.g., PFAM, SMART, CDD, COGs, KOGs) or known structure (e.g., PDB, SCOP). (3) I-TASSER, which uses a hierarchical protein structure modeling approach that is based on the secondary-structure enhanced profile–profile threading alignment to predict protein tertiary structure and function [52]. (4) PredictProtein, which predicts aspects of protein structure (secondary structure, solvent accessibility, transmembrane helices and strands, coiled-coil regions, disulfide bonds and disordered regions) and function (identification of functional regions, homology-based inference of Gene Ontology terms, comprehensive subcellular localization prediction, protein-protein binding sites, protein-polynucleotide binding sites and predictions of the effect of point mutations [non-synonymous SNPs] on protein function) [51]. For HHpred, I-TASSER and PredictProtein, only top three results were recorded.

### Phylogenetic analysis and ancestral state reconstruction

We used 12 mitochondrial PCGs to reconstruct the phylogenetic history of 35 mitogenomes: six new assemblies and twenty-six published assemblies of Arcidae and out-group species. Thirty-five taxa representing 26 species were included in the phylogenetic analyses and presented in Table 1

**Authors' contributions**

NZ, YL, KMH, LK and QL designed and coordinated the research. NZ, LK and QL collected the sample. NZ performed sequencing and bioinformatics analysis. NZ and LK analyzed the data and drafted the manuscript. NZ, YL, KMH, LK and QL participated in revising the manuscript drafts during the writing process. All authors read and approved the final manuscript.

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**Availability of data and materials**

The data underlying this article are available in the article and in its Supplementary Material as well as in the GenBank Nucleotide Database and can be accessed with OM807131- OM807136. Sequencing data files are available through the NCBI Sequence Read Archive (BioProject: PRJNA809524).

**Declarations****Ethics approval and consent to participate**

Not applicable. Samples (deceased) were either purchased from or given by commercial fisherman.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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