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Systematic assessment of the Panopeidae and broader Eubrachyura (Decapoda: Brachyura) using mitochondrial genomics

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Abstract

This study provides a broad phylogenetic analysis for the Eubrachyura, with the inclusion of three new Panopeidae mitochondrial genomes: *Eurypanopeus depressus* (flatback mud crab) (15,854bp), *Panopeus herbstii* (Atlantic mud crab) (15,812bp) and *Rhithropanopeus harrisii* (Harris, or 'white-fingered' mud crab) (15,892bp). These new mitogenomes were analyzed alongside all available brachyuran mitochondrial genomes (n = 113), comprising 80 genera from 29 families, to provide an updated phylogenetic analysis of the infra-order Brachyura ("true crabs"). Our analyses support the subsection Potamoida within the Eubrachyura as the sister group to Thoracotremata. The family Panopeidae aligns with the family Xanthidae to form the Xanthoidea branch, which is supported by current morphological and genetic taxonomy. A unique gene arrangement termed 'XanGO' was identified for the panopeids and varies relative to other members of the subsection Heterotremata (within the Eubrachyura) via a transposition of the *trnV* gene. This gene arrangement is novel and is shared between several Xanthoidea species, including *Etisus anaglyptus* (hairy spooner crab), *Atergatis floridus* (brown egg crab), and *Atergatis integerrimus* (red egg crab), suggesting that it is a conserved gene arrangement within the Xanthoidea superfamily. Our study further reveals a need for taxonomic revision of some brachyuran groups, particularly the Sesarmidae. The inclusion of panopeid mitogenomes into the greater brachyuran phylogeny increases our understanding of crab evolution and higher level Eubrachyuran systematics.

Keywords

Xanthidae, Panopeus, Eurypanopeus, Rhithropanopeus, mud crab, marine, genomics

1. Introduction

Brachyura ("true" crabs) is the largest subgroup of the Decapoda (Crustacea). It is a ubiquitous group, whose members thrive in terrestrial and aquatic habitats but are particularly prevalent in marine environments (Tsang 2009; Jia et al. 2018; Tan et al. 2018; Ma et al. 2019; Tan et al. 2019). Marine Brachyura boast a broad range of morphological and ecological diversity, leading to a complex taxonomy (Yong-kun et al. 2014). Historically, brachyurans were divided into two sections: the Eubrachyura and the Podotremata, with the Eubrachyura being further divided into the subsections: Heterotremata and Thoracotremata (Guinot 2013). The sub-sectioning of these Eubrachyura is based almost entirely on typological morphology (particularly the genital openings) and has been subject to debate with regards to monophyly. Due to the morphological complexity of this group, genetic tools and analytical methods are typically used to resolve systematic discrepancies (Basso et al. 2017; Bai et al. 2018; Jia et al. 2018; Tan et al. 2018; Wang et al. 2018).

High throughput sequencing (HTS) has proven effective in advancing and resolving taxonomies (Tan et al. 2018). Early studies on brachyuran mitogenomics relied on long PCR and primer walking techniques to read and assemble the mitogenome (Yamauchi 2003; Miller 2005). The rapid sequencing and assembly of the mitochondrial genome (mitogenome) using HTS has proven to be a powerful tool for conducting phylogenetic studies of eukaryotes (Gan et al. 2018). The small size of mitogenomes (~ 14-16 kb), potential for high mutation rate, and a simple closed structure, make them an ideal marker for inferring an organism's mitogenetic phylogeny (Boore 1999). Brachyura has been classified into 93 families with over 7000 species (Ng 2008); for 111 species (representing 28 families) the complete mitogenomes are known (NCBI and Supplementary material 1). The Brachyuran genomes that have been sequenced thus far all are between 10-25 kb in length. Much of these data represent three brachyuran families [Portunidae (n = 15); Varunidae (n = 14); Sesarmidae (n = 10)]. The remaining 25 families have < 5 mitogenomes sequenced per family, many of them only having 1 (see Supplementary material). Recent publications using mitogenomics have challenged the validity of the morphologically founded Eubrachyuran subsections of Heterotremata and Thoracotremata. For example, freshwater crabs in the family Potamidae fall into the Heterotremata based on morphology but based on mitogenomics align with members of the Thoracotremata (Basso 2017; Bai et al. 2018; Tan et al. 2018).

Mitogenomes also offer insights into gene arrangement, which can have diagnostic properties at different systematic levels (Boore 1998; Boore 2000; Moret 2001; Perseke et al. 2008; Zhuang 2010; Babbucci 2014; Mindell 2016; Nakjima et al. 2016; Zhang 2020). Within the Malacostraca, mitogenome gene arrangements are conserved within certain groups (Shen 2011; Tan et al. 2019), which allows for simple comparisons at different taxonomic levels. Grouping the Crustacea with the Insecta to form the Pancrustacea has strong support based on the near-identical arrangement of the shared genes across taxa (Boore 1998). However, gene arrangement can vary greatly within crustacean orders. Specifically, in Brachyura many species share a gene arrangement that is thought to be ancestral to Brachyura (termed BraGO). However, recent research has shown that most groups deviate from this pattern forming new arrangements at the family and subfamily levels (Basso 2017; Tan et al. 2018; Wang et al. 2018; Wang 2020a; Wang 2020b). BraGO differs from the Pancrustacea genomic order (PanGO) by a transposition of the gene trnH from a location between the nad5 and nad4 genes to a location between the trnD and trnF genes (Basso 2017). To date, 20 different gene arrangements have been identified within the Brachyura (Basso 2017; Tan et al. 2018; Zhang 2020a), but many brachyuran groups remain un-sequenced and this syntenic diversity could be much higher.

An example of an understudied brachyuran group is the superfamily Xanthoidea (Brachyura), which boasts high diversity across the world's oceans (Karasawa 2006). Species within this superfamily share a high level of morphological similarity and are often poorly described both morphologically and genetically (Ng 2008; Thoma 2014). The number of families and subfamilies within the Xanthoidea has changed drastically in recent years (Lai 2011). Two common families, the Xanthidae and Panopeidae, share several morphological features that can lead to systematic confusion and difficulty in identifying them beyond the family level (Shih 2011). Both families are found in temperate and tropical shallow intertidal and subtidal zones, but xanthid crabs have a circumtropical distribution while panopeids are only found in the Americas, excluding global invasions (Thoma 2014). To date, there are only four mitogenomes available for the Xanthidae and none for the Panopeidae, whose systematics have primarily relied upon a select number of genes or morphological keys (Williams 1984; Schubart 2000). Studies using conventional PCR to amplify and sequence mitochondrial and nuclear markers revealed that the genera Eurypanopeus and Panopeus are polyphyletic (Schubart 2000). Similarly, studies on the panopeid genus Hexapanopeus using 12S and 16S genes as markers have also suggested that this genus is polyphyletic (Thoma 2009). Later studies using three mitochondrial markers (COI, 12S and 16S) and three nuclear markers [18S, enolase (ENO) and Histone H3 (H3)] revealed that Xanthoidea is monophyletic, but its two families are not and are in need of taxonomic revision (Thoma 2014).

In this study, we enhance understanding of brachyuran systematics by adding three complete mitogenomes for the Panopeidae: *Eurypanopeus depressus, Panopeus herbstii* and *Rhithropanopeus harrisii* from their native range along the Atlantic coast of North America. The genetic composition, genetic similarity and gene arrangement of these three panopeid species are described relative to other brachyuran mitogenomes, allowing us to update the brachyuran mitogenomic phylogeny and explore brachyuran-wide classification. A new gene arrangement for the superfamily Xanthoidea is described as well as a renaming of previously reported gene arrangements suggested for other Brachyura.

2. Materials and methods

2.1. Specimen collection and dissection

Three species of panopeid mud crabs were collected for this study. First, an individual Eurypanopeus depressus was sampled on December 1, 2018 from Hoop Pole Creek, a polyhaline site located in Atlantic beach, North Carolina (NC), USA. The individual was hand-collected at low tide from an intertidal oyster reef and then brought back to the lab for dissection. Second, an individual Panopeus herbstii was sampled on August 12, 2019 from Middle Marsh (Beaufort, NC), another polyhaline site, using a passive sampler attached to a wooden stake that had been driven into the sediment. The sampler design is a small plastic milk crate $(19 \times 22 \times 16 \text{ cm})$ filled with autoclaved oyster shell (Roche 2007). Third, an individual Rhithropanopeus harrisii was sampled on February 5, 2020 from Mallard Creek (Washington, NC), a mesohaline site, using the same passive sampling design as above, but this time attached to a small fishing dock. Crabs were brought back to the lab and anesthetized prior to dissection in a -20°C freezer. Dissections for all three species were carried out using a sterilized razor blade, and part of the hepatopancreas and gills were removed and placed into separate tubes for later DNA extraction.

2.2. DNA extraction, sequencing and assembly

The DNA extractions were conducted on the hepatopancreas and gill tissue of E. depressus, P. herbstii, and R. harrisii using a Zymo DNA extraction kit, according to manufacturer's protocols. The DNA samples were shipped on dry ice to Novogene, California, who conducted library preparation using the NEBNext Ultra DNA Library Prep Kit. The library was loaded on to a NovaSeq 6000 (Illumina) system using the 150 bp NovaSeq 600 SP reagent kit (300 cycles) for paired end metagenomic sequencing for each individual sample. The resulting data were delivered to the University of Florida for bioinformatic analysis. The data were quality checked and trimmed using Trimmomatic v.0.36 (Bolger 2014) using default parameters. The paired and unpaired reads were assembled using SPAdes v.3.13.0 (Bankevich et al. 2012) with default parameters and k-mer lengths: 21, 33, 55, 77 and 99. The resulting datasets provided a series of contigs that were compared to the NCBI nr database using BLASTx. The mitochondrial genomes of *E. depressus* (574.088X coverage), *P. herbstii* (122.084X coverage) and *R. harrisii* (400.520X coverage) were each identified and circularized. Confirmation of their sequence coverage was conducted using CLC genomics workbench v.12.

The circularized mitogenomes were annotated using MITOS (Bernt 2013). Using the MITOS output, the location of the cox1 gene was determined and the sequences were re-annotated with the cox1 gene at the start of the genome. The putative amino acid and rRNA sequences determined by MITOS were checked using BLASTn and BLASTp (Tables 1–3). The completed genomes were then annotated graphically using Circa (http://omgenomics.com/circa). The genomes are deposited in GenBank under accession numbers MN399962 (*E. depressus*), MT024989 (*R. harrisii*), and MT024990 (*P. herbstii*).

2.3. Phylogenetic and mitochondrial gene order assessment

There were 112 brachyuran mitogenomes (see Supplementary material 1) obtained from the GenBank database (NCBI) for phylogenetic comparison using the Brachyura taxonomic ID (txid6752) and filtering results to yield DNA sequences of 10,000-25,000 bp (search date: January 2020). The amino acid and nucleotide sequences were retrieved and annotated from these genomes (13 and 15 sequences, respectively) using the Mitophast pipeline (Tan et al. 2015) which downloads each gene in the mitochondrial genome as separate files. The amino acid and nucleotide sequences files were then aligned individually using MAFFT in Geneious (v10.0.2), trimmed to the smallest sequence and concatenated using Geneious. Phylogenetic analyses were conducted in IQtree (Trifinopoulos 2016), which computed the most appropriate evolutionary model (mtMet+F+I+G4) according to BIC for both the amino acid sequences and the nucleotide sequences. A maximum likelihood tree using the amino acid sequences was created using 1000 bootstrap replicates and an SH-aLRT branch test (Guindon 2010) over 3733 positions; the tree had a log score of -157295.9511. A maximum likelihood tree using the nucleotide sequences was created using 1000 bootstrap replicates and an SH-aLRT branch test over 13790 positions; the tree had a log score of -598390.2632. The resulting trees were annotated using FigTree (http://tree.bio.ed.ac.uk/ software/figtree). Both trees were rooted with the mitogenomes of Coenobita brevimanus (KY352233), C. rugosus (KY352235) and C. perlatus (KY352234).

All previously reported gene orders for the Brachyura were annotated according to Basso et al. (2017) and Tan et al. (2018). Pairwise comparisons of the gene orders were performed using CREx software (Bernt et al. 2007) at common intervals. The nomenclature for the gene orders follows Basso et al. (2017). MITOS was used to determine the putative location of the control region (CoRe) for *E. depressus*, *P. herbstii*, *R. harrisii*, *Etisus* *anaglyptus, Leptodius sanguineus, Atergatis floridus*, and *A. integerrimus* through manual examination of the start and stop codons of the open reading frames to look for intergenic spacers. The CoRe for the crabs in the family Potamidae were obtained from Genbank. The CoRes for *Echinoecus nipponicus* and *Pilumnus vespertilio* were determined using MITOS following the same method as for the Panopeidae.

3. Results

3.1. The mitochondrial genomes of panopeid crabs

The mitochondrial genomes of the panopeid crabs used in this study were closed circular molecules containing 13 protein coding genes, 22 tRNA genes, 2 rRNA genes, and a single control region (CoRe) (Fig. 1). The *E. depressus* mitogenome was 15,854 bp in length. The *P. herbstii* mitogenome was 15,812 bp in length. The *R. harrisii* mitogenome was 15,967 bp in length. As with most brachyurans, the *rrnL* (16S) and *rrnS* (12S) genes were located on the negative strand, as are the *nad5*, *nad4*, *nad4L*, *nad1* and 8 tRNA genes (Table 1).

The nucleotide composition of the complete *E. depressus* mitochondrial genome was as follows: A=5442 (34.32%), T=5509 (34.75%), G=1652 (10.42%), C=3251 (20.51%). The A+T and G+C contents were 69.07% and 30.93%, respectively. The protein coding regions include 7 NADH dehydrogenases (*nad1–nad6* and *nad4L*), three cytochrome c oxidases (*cox1–cox3*), 2 ATPases (*atp6* and *atp8*) and 1 cytochrome *b* (*cob*) and account for 10,838 bp of the mitogenome. The 22 rRNA genes present in the mitogenome range in size from 62 (*trnD)–*71 (*trnL1*) bp in length, and the ribosomal RNA genes *rrnL* (16S) and *rrnS* (12S) have a length of 1393 bp and 817 bp, respectively. The 13 protein coding genes and majority of the ncRNA sequences showed similarity among the panopeid crabs used in this study (Table 1).

Figure 1. Annotated Circa plots for the circular mitochondrial genomes of Eurypanopeus depressus, Panopeus herbstii and Rithropanopeus harrisii. Each mitogenome is represented by a thick circular black line near the centre of the plot. Protein coding genes are on the outside of this line (negative = dark violet, positive = maroon). Non-coding RNA genes are on the inside of this line (negative = light violet, positive = light maroon). The genome sizes are written in the centre of each plot. The protein coding gene names are represented in the outer most circle (dark grey). The ncRNA gene names are listed in the second internal circle (light grey). The green rectangle labelled "CoRe" indicates the putative control region of the mitochondrial genomes. Figure 1 layout: Portrait. Associated with section 3.1



Table 1. Nucleotide and protein similarity data for the protein-coding and non-coding genes of the Eurypanopeus depressus mitochondrial genome. The data represented were acquired from BLASTn and BLASTp outputs via comparison against the complete non-redundant database. The accession number of the specific nucleotide or amino acid sequence are provided in addition to the species, if known, belonging to the sequence isolate. The similarity (%), coverage comparison (%) and e-value are all provided. MCG = mitochondrion, complete genome.

niətor4 noizzəzəs	Present study		Present study			Present study	Present study	Present study		Present study								Present study	Present study	Present study			Present study	Present study
Protein -value	0.0		1e-170			6e-18	6e-158	0.0		1e-79								0.0	0.0	8e-66			6e-90	0.0
Protein eover	100		100			100	100	100		66								98	66	100			100	100
Protein similarity	99.8		99.11			90.20	99.10	99.61		96.49								92.75	96.15	100.00			92.73	98.68
Protein hit	cytochrome c oxidase subunit I [Rhith- ropanopeus harrisii]		cytochrome c oxidase subunit II [Pa- nopeus herbstii]			ATP synthase F0 subunit 8 [Panopeus herbstii]	ATP synthase F0 subunit 6 [Panopeus herbstii]	cytochrome c oxidase subunit III [Rhithropanopeus harrisii]		NADH dehydrogenase subunit 3 [Panopeus herbstii]								NADH dehydrogenase subunit 5 [Rhithropanopeus harrisii]	NADH dehydrogenase subunit 4 [Panopeus herbstii]	NADH dehydrogenase subunit 4L [Panopeus herbstii]			NADH dehydrogenase subunit 6 [Rhithropanopeus harrisii]	cytochrome b [Panopeus herbstii]
Gene noissoos	Present study	Present study	Present study	Present study	Present study		Present study	Present study	Present study			Present study		Present study	Present study	Present study	Present study	Present study	Present study	Present study				
Gene e-value	0.0	5e-23	0.0	2e-27	3e-32		0.0	0.0	1e-30		1e-28	4e-31		9e-31	6e-29	9e-26	9e-26	0.0	0.0	1e-107	2e-29	1e-31	4e-147	0.0
Сепе Соvегаде (%)	100	100	100	100	100		100	100	100		100	100		100	100	100	100	98	66	98	95	100	98	66
Gene similarity (%)	88.05	92.42	88.99	95.52	100		87.80	89.83	98.41		98.41	98.44		98.51	95.45	95.31	95.31	88.79	87.30	91.21	98.39	98.46	85.51	86.77
tid ənəƏ	Rhithropanopeus harrisii MCG	Panopeus herbstii MCG	Rhithropanopeus harrisii MCG	Panopeus herbstii MCG	Rhithropanopeus harrisii MCG		Panopeus herbstii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	1	Panopeus herbstii MCG	Rhithropanopeus harrisii MCG	I	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG				
Strand	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+						+		+	+
ənəÐ	cox1	trnL2(tta)	cox2	trnK(aaa)	trnD(gac)	atp8	atp6	cox3	trnG(gga)	nad3	trnA(gca)	trnR(cga)	trnN(aac)	trnS1(aga)	trnE(gaa)	trnH(cac)	trnF(ttc)	nad5	nad4	nad4L	trnT(aca)	trnP(cca)	nad6	cob
puJ	1515	1598	2277	2357	2420	2573	3238	4032	4100	4448	4518	4582	4649	4718	4786	4871	4935	6574	8046	8318	8408	8473	8970	10118
Start	1	1535	1606	2291	2358	2421	2576	3256	4038	4107	4456	4519	4583	4652	4721	4808	4872	4943	6721	8043	8345	8409	8476	8982
Genome							ວບ	ແດນຈອ	lsirt	ouoyoo	im	snss	səлd	əp s	məd	ouv	dAn	$n_{\overline{J}}$						

Ргоtеіп яссеззіоп		Present study		l	l					Present study			
Protein -value		0.0								0.0			
Protein 2076r		100								66			
Protein similarity		98.67								90.62			
Protein hit		NADH dehydrogenase subunit 1 [Rhithropanopeus harrisii]								NADH dehydrogenase subunit 2 [Panopeus herbstii]			
Gene noizzəcor	NC_039618.1	Present study		KT959469.1	EU863325.2	Present study	Present study	Present study	NC_037201.1	Present study		NC_037201.1	NC_042208.1
9n9D e-value	2e-15	0.0		0.0	0.0	3e-26	4e-31	2e-30	1e-22	0.0		2e-29	4e-12
Gene Coverage (%)	97	66		98	44	100	100	100	100	100		100	100
Gene similarity (%)	95.85	88.95		91.03	99.73	95.45	97.06	95.65	98.53	82.18		96.88	90.77
tid ənəD	Echinoecus nipponicus voucher MABIK CR00241788 MCG	Rhithropanopeus harrisii MCG		Eurypanopeus depressus voucher USNM 16S RNA gene	<i>Eurypanopeus depressus</i> voucher ULLZ 3976 12S ribosomal RNA gene, partial sequence; mitochondrial	Rhithropanopeus harrisii MCG	Panopeus herbstii MCG	Rhithropanopeus harrisii MCG	Atergatis floridus MCG	Rhithropanopeus harrisii MCG		Panopeus herbstii MCG	Etisus anaglyptus MCG
Strand	+						+		+	+	+		
ənəĐ	trnS2(tca)	nad1	trnL1(cta)	rmL	IIIS	trnV(gta)	trnl(atc)	trnQ(caa)	trnM(atg)	nad2	trnW(tga)	trnC(tgc)	trnY(tac)
рид	10183	11134	11242	12610	13522	14204	14489	14555	14646	15621	15723	15787	15852
Start	10117	10235	11171	11217	12705	14140	14422	14487	14579	14659	15656	15724	15788
Genome		əu	ioua	atrial go	uoqootim su	ssəл	dəp	snə	dou	vdAn _H			

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Protein Protein	Present study		Present study			Present study	Present study	Present study		Present study								Present study	Present study	Present study			Present study
niətorA 9ulrv-9	0.0		1e-170			6-e18	6e-158	0.0		1e-79								0.0	0.0	8e-66			7e-89
Protein Cover	100		100			100	100	100		66								66	66	100			66
Protein similarity	100.00		99.11			90.20	99.10	98.46		96.49								93.64	96.15	100.00			91.52
Protein hit	cytochrome c oxidase subunit I [Rhith- ropanopeus harrisii]		cytochrome c oxidase subunit II [<i>Eurypanopeus depressus</i>]			ATP synthase F0 subunit 8 [Eurypa- nopeus depressus]	ATP synthase F0 subunit 6 [Eurypa- nopeus depressus]	cytochrome c oxidase subunit III [<i>Eurypanopeus depressus</i>]		NADH dehydrogenase subunit 3 [<i>Eurypanopeus depressus</i>]								NADH dehydrogenase subunit 5 [Eurypanopeus depressus]	NADH dehydrogenase subunit 4 [<i>Eurypanopeus depressus</i>]	NADH dehydrogenase subunit 4L [<i>Eurypanopeus depressus</i>]			NADH dehydrogenase subunit 6 [Rhithropanopeus harrisii]
onsd noizessoor	Present study	Present study	Present study	Present study	Present study		Present study	Present study	Present study		Present study	Present study		Present study	Present study	Present study		Present study	Present study	Present study		Present study	Present study
Gene e-value	0.0	6e-23	0.0	2e-29	3e-27		0.0	0.0	6e-29		1e-28	6e-29		4e-29	2e-28	2e-27		0.0	0.0	6e-106		1e-31	2e-130
Gene Coverage (%)	100	100	100	100	100		100	100	100		100	100		100	100	100	I	66	66	98	I	100	97
Gene similarity (%)	87.52	92.42	88.24	95.52	95.24		87.80	89.32	96.88		98.41	96.88		97.01	95.45	96.88		87.93	85.54	90.84		98.46	83.54
tirl ənəƏ	Rhithropanopeus harrisii MCG	Eurypanopeus depressus MCG	Rhithropanopeus harrisii MCG	Eurypanopeus depressus MCG	Rhithropanopeus harrisii MCG		Eurypanopeus depressus MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG		Eurypanopeus depressus MCG	Rhithropanopeus harrisii MCG		Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Rhithropanopeus harrisii MCG		Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG		Rhithropanopeus harrisii MCG	Eurypanopeus depressus MCG
Strand	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+						+		+
ənəÐ	cox1	trnL2(tta)	cox2	trnK(aaa)	trnD(gac)	atp8	atp6	cox3	trnG(gga)	nad3	trnA(gca)	trnR(cga)	trnN(aac)	trnS1(aga)	trnE(gaa)	trnH(cac)	trnF(ttc)	nad5	nad4	nad4L	trnT(aca)	trnP(cca)	nad6
puI	1515	1600	2278	2358	2421	2574	3239	4033	4102	4450	4520	4583	4651	4719	4785	4868	4935	6550	8047	8319	8409	8474	8974
Start	1	1535	1607	2292	2359	2422	2577	3257	4039	4109	4458	4521	4584	4653	4722	4805	4869	4943	6725	8044	8346	8410	8477
Genome							ə	monəū) lsi	гриоцэ	otil	N !!!	ısqл	əy s	nəde	ouv	I						

Ргоtеіп яссеззіоп	Present study		Present study			l					Present study			
Protein 9ulav-9	0.0		0.0								0.0			
Protein 207er	100		66								100			
Protein similarity	98.68		97.00								90.62			
Protein hit	cytochrome b [Eurypanopeus depres- sus]		NADH dehydrogenase subunit 1 [<i>Eurypanopeus depressus</i>]								NADH dehydrogenase subunit 2 [<i>Eurypanopeus depressus</i>]			
Gene noizzeoor	Present study	Present study	Present study		KT959516.1	EU863296		Present study	Present study	NC_042208			Present study	Present study
9n9D e-value	0.0	4e-26	0.0		0.0	0.0		4e-31	2e-30	9e-25	I		4e-31	2e-24
Gene Coverage (%)	66	100	98		37	44		100	100	100			100	100
Gene similarity (%)	87.13	92.65	89.40		100.00	99.46		97.06	95.65	98.55			98.44	92.31
tid ənəƏ	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG		Panopeus herbstii voucher USNM: 16S RNA gene, mitochondrial	Panopeus herbstii voucher ULLZ 8457 12S ribosomal RNA gene, partial sequence; mitochondrial		Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Etisus anaglyptus MCG			Rhithropanopeus harrisii MCG	Rhithropanopeus harrisii MCG
Strand	+	+						+		+	+	+		
Gene	cob	trnS2(tca)	nad1	trnL1(cta)	rrnL	ITINS	trnV(gta)	trnl(atc)	trnQ(caa)	trnM(atg)	nad2	trnW(tga)	trnC(tgc)	trnY(tac)
рид	10119	10184	11135	11239	12584	13502	14190	14423	14489	14609	15581	15685	15748	15812
Start	8983	10118	10230	11171	11194	12683	14124	14357	14421	14541	14622	15619	15685	15749
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	niətorA noize930218	Present study		Present study			Present study	Present study	Present study		Present study								Present study	Present study	Present study		
	niətorA 9.1kv-9	0.0		8-e170			2e-16	1e-156	0.0		9e-73								0.0	0.0	2e-64		
	Protein cover	100		100			100	100	100		100								86	100	100		
	Protein Vinilarity	100.00		99.11			84.31	97.74	99.61		94.74								92.75	94.80	96.74		
•	Protein hit	cytochrome c oxidase subunit I [Pa- nopeus herbstii]		cytochrome c oxidase subunit II [Pa- nopeus herbstii]			ATP synthase F0 subunit 8 [Panopeus herbstii]	ATP synthase F0 subunit 6 [Eurypa- nopeus depressus]	cytochrome c oxidase subunit III [Eurypanopeus depressus]		NADH dehydrogenase subunit 3 [<i>Eurypanopeus depressus</i>]								NADH dehydrogenase subunit 5 [Eurpanopeus depressus]	NADH dehydrogenase subunit 4 [<i>Eurypanopeus depressus</i>]	NADH dehydrogenase subunit 4L [Eurypanopeus depressus]		
	Gene acces- sion	LN810615		Present study	Present study	Present study		Present study	Present study	Present study		Present study	Present study		Present study	Present study	Present study	Present study	Present study	Present study	Present study	Present study	Present study
	9n9Đ e-value	0.0		0.0	7e-29	3e-32		0.0	0.0	1e-30		1e-28	4e-31		9e-31	6e-29	2e-27	8e-28	0.0	0.0	1e-107	2e-29	1e-31
	Gene Co- verage (%)	65		100	100	100	I	100	100	100		100	100		100	100	100	100	66	66	98	95	100
	Gene simi- larity (%)	99.39		88.99	95.52	100.00		88.54	89.86	98.41		98.41	98.44		98.51	95.45	96.88	95.31	88.79	87.30	91.21	98.39	98.46
)	tid ənəƏ	Rhithropanopeus harrisii mitochondrial partial COI gene for cytochrome oxidase subunit 1, isolate R617-8	1	Eurypanopeus depressus MCG	Panopeus herbstii MCG	Eurypanopeus depressus MCG	1	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	1	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	1	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Panopeus herbstii MCG	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Panopeus herbstii MCG
	Strand	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+						+	
-	Gene	cox1	trnL2(tta)	cox2	trnK(aaa)	trnD(gac)	atp8	atp6	cox3	trnG(gga)	nad3	trnA(gca)	trnR(cga)	trnN(aac)	trnS1(aga)	trnE(gaa)	trnH(cac)	trnF(ttc)	nad5	nad4	nad4L	trnT(aca)	trnP(cca)
))	рид	1515	1599	2278	2357	2420	2573	3238	4032	4100	4448	4517	4581	4648	4717	4786	4866	4930	6554	8037	8309	8400	8465
-	Start	-	1535	1607	2292	2358	2421	2576	3256	4038	4107	4455	4518	4582	4651	4721	4803	4867	4941	6712	8034	8336	8401
	Genome					ຈແ	Genor	ondrial	Mitoch	[ətə	Idmo	<u>115</u>	เมต	y sn	ədo	uvd	олүз	įųу					

Protein noiseoos	Present study	Present study		Present study								Present study			
niətor4 9. value	6e-90	0.0		0.0								2e-168			
Protein Protein	100	100		100								98			
Protein similarity	92.73	98.68		98.67								91.28			
Protein hit	NADH dehydrogenase subunit 6 [<i>Eurypanopeus depressus</i>]	cytochrome b [Panopeus herbstii]		NADH dehydrogenase subunit 1 [Eurypanopeus depressus]								NADH dehydrogenase subunit 2 [<i>Eurpanopeus depressus</i>]			
-eene acces- nois	Present study	Present study	Present study	Present study		KT959486.1	EU863280	Present study	Present study	Present study	Present study	Present study		Present study	Present study
Gene Gene	3e-148	0.0	4e-24	0.0		0.0	0.0	3e-26	4e-31	3e-27	1e-29	0.0		4e-31	2e-24
Gene Co- Verage (%)	98	100	100	66		37	44	100	100	100	100	98		100	100
Gene simi- larity (%)	85.45	87.13	92.65	89.40		100.00	98.90	95.45	97.06	92.75	97.10	82.16		98.44	92.31
tift ənəƏ	Eurypanopeus depressus MCG	Panopeus herbstii MCG	Panopeus herbstii MCG	Panopeus herbstii MCG	1	Rhithropanopeus harrisii voucher USNM 12S ribosomal RNA gene, partial sequence; mitochondrial	Rhithropanopeus harrisii vou- cher ULLZ 3995 12S ribosomal RNA gene, partial sequence; mitochondrial	Eurypanopeus depressus MCG	Eurypanopeus depressus MCG	Panopeus herbstii MCG	Panopeus herbstii MCG	Eurypanopeus depressus MCG	1	Panopeus herbstii MCG	Panopeus herbstii MCG
Strand	+	+	+						+		+	+	+		
ene	nad6	cob	trnS2(tca)	nad1	trnL1(cta)	ımL	Smr	trnV(gta)	trnI(atc)	trnQ(caa)	trnM(atg)	nad2	trnW(tga)	trnC(tgc)	trnY(tac)
puI	8962	10107	10175	11123	11228	12583	13499	14208	14496	14562	14687	15680	15764	15827	15892
Start	8468	8974	10109	10224	11160	11184	12683	14143	14430	14494	14620	14700	15697	15764	15828
Genome			ວເ	nonsD	lsin	e Mitochond	təlqmoD <i>ilsi</i>	лрү	snə	dou	vdo.	иңің			



Figure 2. Maximum-likelihood phylogenetic relationships derived from 112 species of brachyuran crabs, using 13 concatenated amino acid sequences (cox1-cox3, cob, atp6, atp8, nad1, nad5, nad4, nad4-L). Some families have been collapsed for increased clarity (triangles). Black circles on nodes represent an SH-aLRT and bootstrap support of greater than 90/90. Stars (*) indicate areas on the tree with taxonomic conflicts related to previous literature. The symbol α indicates the family Xanthidae; β indicates the family Panopeidae. See Supplementary material 1 for a list of the species used and their accession numbers involved in this analysis.

The nucleotide composition of the complete P. herbstii mitochondrial genome was as follows: A=5520 (34.91%), T=5687 (35.97%), G=1627 (10.29%), C=2980 (18.85%). The A+T and the G+C contents were 70.87% and 29.13%, respectively. The protein coding region contains 7 NADH dehydrogenases (nad1-nad6 and nad4L), three cytochrome c oxidases (cox1-cox3), 2 ATPases (atp6 and atp8) and 1 cytochrome b (cob) and accounts for 10947 bp of the mitogenome of P. herbstii. The 22 rRNAs present in the mitogenome range in size from 63 (trnD, trnA, trnR) - 69 (trnL1, trnQ, trnM) bp in length, and the ribosomal RNA genes rrnL (16S) and rrnS (12S) have a length of 1392 bp and 820 bp, respectively. All 13 protein coding genes showed high similarity to the panopeid crabs used in this study. The ncRNAs all showed similarity to decapod crustaceans with the majority having high similarity with the Panopeidae (Table 2).

The nucleotide composition of the complete *R. harrisii* mitochondrial genome was as follows: A=5595 (34.21%), T=5873 (37.00%), G=1556 (9.82%), C=2866 (17.99%). The A+T and the G+C contents were 72.20% and 27.080%, respectively. The protein coding region contains 7 NADH dehydrogenases (*nad1–nad6* and *nad4L*), three cytochrome c oxidases (*cox1–cox3*), 2 AT-Pases (*atp6* and *atp8*) and 1 cytochrome b (*cob*) and account for 10,848 bp of the mitogenome. The 22 rRNAs present in the mitogenome range in size from 63 (*trnD*, *trnG*, *trnA*) – 69 (*trnL1*, *trnQ*) bp in length, and the ribosomal RNA genes *rrnL* (16S) and *rrnS* (12S) have a length of 1400 bp and 817 bp, respectively. The 13 protein coding genes showed high similarity with the panopeid crabs used in this study (Table 3).

3.2. Phylogenetics

To establish where the panopeid crabs align within the Eubrachyrua, amino acid and nucleotide sequences from 112 mitogenomes comprising 77 genera from 28 families of brachyuran crabs were used along with the three new mitogenomes (Fig. 2). Two sequences that are publicly available for brachyurans were not included in our analysis due to inconsistences with the sequences. (1) The protein sequences for *Gecarcoidea natalis* contained ambiguous amino acid identifications, resulting in poor alignment with other members within the superfamily Grapsoidea. (2) The protein sequences for *Pyrhila pisum* aligned poorly with other members of the Brachyura; however, there were no missing protein codes. When tested in BLASTp, the proteins for *P. pisum* yielded low identity with other brachyurans; < 60% identity in most cases.

Four distinct clades were identified (Fig. 2). One clade belongs to crabs in the subsection Heterotremata (n=40), a second belongs to crabs in the subsection Thoracotremata (n=44) and a third belongs to crabs in the section Podotremata (n=7). The fourth clade belongs to the 'Old World' freshwater crabs in the superfamilies Potamoidea and Gecarcinucoidea (n=20). This fourth clade forms a subsection termed Potamoida, a sister group to Thoracotremata. The split between the Heterotremata and the Potamoida/Thoracotremata clades is well supported using both amino acid sequences (Sh-aLRT/UFBoot: 100/100) as well as nucleotide sequences (Sh-aLRT/UF-Boot:100/100). The Potamoida and Thoracotremata split is also well supported using both sequence types (amino



Figure 3. Maximum-likelihood phylogenetic relationships derived from 112 species of brachyuran crabs, using 15 concatenated nucleotide sequences (*cox1–cox3*, *cob*, *atp6*, *atp8*, *nad1*, *nad5*, *nad4*, *nad4–L*, *rrnL*, *rrnS*). Some families have been collapsed for increased clarity (triangles). Black circles on nodes represent an SH-aLRT and bootstrap support of greater than 90/90. Stars (*) indicate areas on the tree with taxonomic conflicts related to previous literature. The symbol α indicates the family Xanthidae; β indicates the family Panopeidae. See Supplementary material 1 for a list of the species used and their accession numbers involved in this analysis.

acids- Sh-aLRT/UFBoot: 98.9/99; nucleotides- Sh-aLRT/UFBoot: 92.9/98).

The panopeid crab species E. depressus, P. herbstii and R. harrisii formed a branch for the family Panopeidae (Fig. 2 and Fig. 3; " β ") aligned alongside the xanthid branch to form the superfamily Xanthoidea (amino acids-Sh-aLRT/UFBoot: 100/100; nucleotides- Sh-aLRT/UF-Boot: 100/100). The xanthid branch contains members of the Xanthidae family: E. anaglyptus, A. floridus and A. integerrimus (Fig. 2 and Fig. 3; "a"). When considering its amino acid sequences, the crab species Epixanthus frontalis from the family Oziidae aligns with the Xanthoidea superfamily with moderate support (Sh-aLRT/ UFBoot: 89.4/94) (Fig. 2). The nucleotide sequences for E. frontalis show a similar pattern; however, Leptodius sanguineus is part of the branch with middling support (Sh-aLRT/UFBoot: 66.7/93) (Fig. 3). Based on amino acid comparison, L. sanguineus (considered a member of the Xanthidae) aligns between E. frontalis and members of the Pilumnidae, on a branch separate from other xanthid crabs (Sh-aLRT/UFBoot: 16/66) (Fig. 2). The amino acid phylogeny suggests that the hydrothermal vent crabs in the family Xenograpsidae align with the terrestrial crabs in the family Ocypodidae (Sh-aLRT/UFBoot: 83.2/72), yet the nucleotide sequences suggest that the xenograpids form their own branch alongside of the sesarmid crabs (Sh-aLRT/UFBoot: 100/99).

The family Sesarmidae (10 mitogenomes) appears to be polyphyletic. Rather than grouping together, the genus *Chiromantes* is split, where *C. dehaani* aligns with *Sesarma neglectum* (amino acids- Sh-aLRT/UFBoot: 99.5/100; nucleotides- Sh-aLRT/UFBoot: 100/100), and *C. haematocheir* aligns with *Sesarmops sinensis* (amino acids- Sh-aLRT/UFBoot: 99.7/100; nucleotides- Sh-aLRT/UFBoot: 100/100) (Fig. 2 and Fig. 3).

3.3. Gene arrangement among the Brachyura, incorporating the Panopeidae

The gene arrangements for the panopeid crabs E. depressus, P. herbstii and R. harrisii (Fig. 3) correspond in synteny to other sequenced xanthid species: E. anaglyptus, A. floridus and A. integerrimus. This gene arrangement differs from both the PanGO and BraGO, where the rrnL and rrnS are adjacent to each other and the trnV is transposed past the CoRe (Table 1). The gene order for the xanthid crab species L. sanguineus reported by Tan et al. (2018) is different from the other xanthids presented in this study, following the basic BraGO pattern rather than the shared pattern of the superfamily Xanthoidea. Based on the CREx test, the new gene arrangement XanGO shares 870 common intervals with PanGO and 988 common intervals with BraGO (Fig. 3), suggesting it to be a low-level rearrangement relative to the common gene arrangements. The new XanGO is most different to the MaVaGO, sharing only 80 common intervals.

BraGO Br	
GeoGO <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>VOO</u> <u>V</u>	σ
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Xango Xango Xango Xango	racotren
Coxi C	Tho

	PanGo	BraGo	XanGo	MajGO	MaVaGO	SesGO	XenGo	SinGO	GeoGO	DamGO	DynGO	HuaGO	PotGO	PilGO	SomGO
PanGo	1326														
BraGo	1188	1326													
XanGo	1122	1256	1326												
MajGO	290	300	292	1326											
MaVaGO	122	118	124	156	1326										
SesGO	1122	1256	1188	296	124	1326									
XenGo	136	146	140	242	58	152	1326								
SinGO	680	782	776	276	116	786	132	1326							
GeoGO	600	690	670	232	116	708	102	644	1326						
DamGO	876	994	934	296	124	1064	150	766	654	1326					
DynGO	942	1064	1060	308	120	1124	136	782	690	886	1326				
HuaGO	1058	1188	1118	296	126	1256	144	792	728	1002	1188	1326			
PotGO	996	1122	1124	292	124	1190	140	782	690	944	1188	1188	1326		
PilGO	548	634	630	228	108	620	106	574	878	582	598	598	596	1326	
SomGO	194	194	186	240	74	190	134	148	126	288	202	190	186	122	1326

Figure 4. Gene orders (-GO) found among brachyuran crabs. Red boxes indicate protein coding genes. Blue boxes indicate tRNA's. Green boxes indicate rRNAs. Purple boxes indicate the control region (CoRe). The red lines along the bottom of the gene orders represents areas within the gene order that are located on the negative strand. Not shown are the 9 unique gene orders for the freshwater crabs (see Zhang et al. 2020). The CREx results are listed for the different gene orders. In the associated table, gene orders with high similarity (> 1000) have red boxes while those with low similarity (< 200) have blue boxes. Intermediate similarity remains white.

The mitogenomes of the crabs in the family Panopeidae all shared a ~600 bp long intergenic spacer between the *rrnS* and *trnV* ncRNA genes (*E. depressus*, 618 bp; *P. herbstii*, 622 bp; *R. harrisii*, 644 bp) representing the control region (CoRe) (Fig. 1). The CoRe in the panopeid mitogenomes are A + T skewed (78.40–80.22%) and contain the repeated motifs TA (125–107), AT (112–104), TAA (47–39), TTA (30–40), ATA (43–35) and TAT (41–37). The mitogenomes of the xanthid crabs used in this study also have similar sized intergenic spacers in this region, suggesting that this is the putative location of the CoRe for members of the superfamily Xanthoidea. The CoRe

nucleotide sequence for all species within Xanthoidea were isolated and run through BLASTn, resulting in a lack of any significant similarity, suggesting high mutability.

4. Discussion

This study provides the first mitochondrial genomes for three members of the Panopeidae and an updated concatenated mito-phylogenetic analysis for the Eubrachyura (excluding nuclear genetic data), informing upon the systematics of multiple families and higher taxonomic rankings. In addition, the mitochondrial genomes for members of the Panopeidae are identified with a consensus gene arrangement shared with other Xanthoidea (XanGO). These results advance our systematic understanding of the brachyurans through the exploration of mitochondrial genomics and gene synteny rearrangement events.

4.1. Xanthid systematics considering panopeid mitogenomic data

The mitogenomes of the panopeid crabs E. depressus, P. herbstii and R. harrisii support the position of the Panopeidae within the Heterotremata, helping to build/support the branch belonging to the superfamily Xanthoidea (Ng 2008). Along this branch, the Xanthidae and Panopeidae form sister groups, additionally supported by previous genetic data using five or less mitochondrial and nuclear genes (Thoma 2009; Lai 2011; Thoma 2014). The genera within these families have been historically identified as polyphyletic (Thoma 2009) and the limited number of mitogenomes available makes it difficult to determine their validity. We acknowledge that the families Xanthidae and Panopeidae both occur in two forms: sensu stricto and sensu lato (Ng 2008). There are 4 publicly available mitogenomes for the Xanthidae (GenBank) and we provide 3 additional mitogenomes for the Panopeidae. We have treated these families in their simple form due to the lack of genetic information to split them further. As more mitogenomes become available, the validity of the two forms should be revisited.

Several taxonomic conflicts appear when considering mitogenetics surrounding the Xanthoidea. First, based on morphology and limited mitogenome availability, the genus *Leptodius* is considered a member of the family Xanthidae. However, despite this genus having 12 separate species, only one mitogenome (the species *L. sanguineus*) is available for analysis. Previous studies showed that *L. sanguineus* aligns closely with other members of the Xanthidae (Sung 2016; Karagozlu 2018; Xie 2018; Ma 2019), but these studies use fewer brachyuran mitogenomes in their analysis prior to our study. When considering all mitogenomes available for the Brachyura in our investigation, *L. sanguineus* aligns more closely with the members

of the family Oziidae, rather than the Xanthidae. This interesting observation merits further exploration.

4.2. Mitogenomic gene arrangements across the Brachyura

Gene arrangement changes were once thought to be rare (Boore 2000) but with greater availability of mitogenome sequencing, it appears that changes in gene arrangements can be common across groups. For example, gene order is conserved within Osteichthyes and some subgroups of Mammalia, while it varies strongly in e.g. Ctenophora (Arafat 2018), Mollusca (Guerra 2018), Hymenoptera (Dowton 1999) and Anomura (Tan et al. 2018). For Crustacea, some species within the Stomatopoda, Amphipoda and Dendrobranchiata still carry the PanGO ground pattern of Pancrustacea (Shen 2011), while no sequenced species within the Brachyura have retained this gene order. Studies on gene order rearrangement are ongoing with some hypothesizing that the evolution to living within harsh environments, such as the deep sea or hydrothermal vents, can lead to new gene synteny (Nakajima 2016; Gan 2018; Tan et al. 2019).

The brachyurans include several families found in the deep sea. Two of them are represented herein: Bythograeidae and Xenograpsidae. Bythograeidae possess the Bra-GO arrangement plesiomorphic for Brachyura, while Xenograpsidae have their own gene arrangement (XenGO). In contrast, the freshwater crab family Potamidae has 9 different gene arrangements (Zhang 2020). Brachyuran crabs represent both cases: the adaptation from a marine to a freshwater environment was likely harsh and may have resulted in several new gene arrangements, while in contrast, the evolution of crabs to the deep-sea benthos resulted in some retaining the ancestral gene order in the face of a new environmental extreme. Therefore, when considering crabs, living within harsh environments does not seem to be the only answer to gene arrangement plasticity, but perhaps requires consideration at the finer scale of environmental adaption. Similar findings have been reported by Tan et al. (2019) who found little evidence for linking gene order rearrangements with adaptations to extreme environments, concluding that these cues are poorly understood and merit a more detailed approach.

A comparison of the eubrachyuran subsections shows that Heterotremata has a higher diversity of gene arrangements than Thoracotremata. Both subsections share species whose gene arrangement follows the basic BraGO pattern. Aside from the BraGO, Thoracotremata only has 3 unique gene arrangements while Heterotremata has 8 unique gene arrangements (including the herein newly established XanGO). This does not include the gene arrangements for the freshwater crabs in the superfamilies Potamoidea and Gecarcinucoidea. The freshwater crabs have more unique gene arrangements than the known Heterotremata.

The panopeid crabs *E. depressus*, *P. herbstii* and *R. harrisii* all have the trnV gene transposed from between the rrnL and rrnS genes to a location past the CoRe.

This differs from the PanGO, BraGO, SesGO, XenGO, DamGO, MajGO and DynGO, which all have the trnV gene located between the rrnL and rrnS genes, with the CoRe following the rrnS gene. The xanthid crabs E. anaglyptus, L. sanguineus, A. floridus and A. integerrimusi all share the latter gene arrangement, suggesting that it might be a conserved arrangement within Xanthoidea and thus support our interpretation of the new Xanthoidea gene arrangement (XanGO). The intergenic spacer found between the rrnS and trnV genes in panopeids appears to be the putative location of the CoRe for these species and is shared with xanthid species, E. anaglyptus, A. floridus and A. integerrimus. All have similarly sized intergenic spacers (600-750 bp long) at this location, suggesting that this may be the location of the CoRe across Xanthoidea. Apart from L. sanguineus, the Xanthidae all follow the new gene arrangement XanGO. Leptodius sanguineus follows the plesiomorphic brachyuran gene arrangement BraGO and based on its amino acid sequences, it groups more closely with the family Pilumnidae than the members of the Xanthidae or the panopeids presented here; however, nodal support is low, meriting further study and sequencing of closer relatives. Higher nodal support is offered with the nucleotide tree, where L. sanguineus groups with Epixanthus frontalis from Oziidae rather than with the xanthids. Based on the molecular taxonomy and its gene arrangement, the placement of L. sanguineus within Xanthidae appears to be invalid and in need of revision, adding to our explanation above.

The mitogenome analysis we performed also supports the renaming of two gene arrangements and confirms the correct gene sequence for another. Two mitogenomes were available for the pilumnid crabs, Echinoecus nipponicus and Pilumnus vespertilio. They follow the gene arrangement reported by Tan et al. (2018) and differ from BraGO in having the trnL gene transposed from its location between the cox1 and cox2 genes to a location between the second *trnL* and *rrnL* genes. This gene arrangement was reported by Tan et al. (2018) as number 12, but we propose Pilumnidae gene order (PilGO) to follow the original gene nomenclature determined by Basso et al. (2017). Similarly, the gene arrangement reported as number 5 by Tan et al. (2018) we rename to the Somanniathelphusa gene order (SomGO). Basso et al. (2017) report the gene arrangement GeoGO as having the trnL gene between the cox1 and cox2 genes, but based on the gene arrangement listed in Genbank, this is nonconcurrent. The correct gene arrangement was reported by Tan et al. (2019) and is supported here with the addition of the mitogenome for Geothelphusa sp. (MG674171), where the trnL gene is located between nad1 and the second trnL gene. This corrected nomenclature should be incorporated into further taxonomic assessments.

4.3. Conclusions

This study provides an updated mitophylogeny for the Brachyura, utilizing all available mitogenomes, along with the first mitogenomes for the Panopeidae, a high-

ly abundant group of ecologically important estuarine crabs with a limited phylogenetic understanding. Our data support the subsection, Potamoida, within the Eubrachyura. The addition of E. depressus, P. herbstii and R. harrisii mitogenomes provides a greater phylogenetic understanding of a group that has been taxonomically challenging in the past. Moreover, the addition of mitogenomes from the Panopeidae further supports the split of the Xanthoidea into multiple families. The novel gene arrangement we describe within the Heterotremata, increases the total number of unique gene arrangements within this subsection to eight. Whilst our results clarify some phylogenetic relationships, they also highlight the need for further study of the genus Leptodius which appears to be incorrectly placed within the subfamily Xanthoidea. Greater sequencing efforts will provide more comparative data for these underrepresented crab groups, and should include the incorporation of nuclear genetic data where possible.

5. Author contributions

AMHB collected the crabs used in the study. JB performed the extraction and bioinformatic processing/assembly of the mitogenomes. LAJ and JB performed the phylogenetics and gene similarity assessments. Gene order analysis and annotation was performed by LAJ and JB. LAJ, AMHB, KAM, DCB and JB contributed to the writing of the manuscript.

6. Competing interests

The authors declare no competing interests.

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Supplementary material 1

Table S1

Authors: Jennings et al. (2021)

Data type: .docx

Explanation note: NCBI accession numbers for species used to conduct phylogenetic analysis.

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