RESEARCH ARTICLE

Phylogenetic positions of some species of the genus *Macrobrachium* Bate, 1868 (Crustacea, Decapoda, Palaemonidae) in Sri Lanka

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Abstract: *Macrobrachium* species, are an economically important freshwater prawn group in Sri Lanka. These species are recognized by their local names and many synonyms can be found for one species. Identification of taxonomic positions and species boundaries within the genus is important to obtain reliable information for application in aquaculture and biodiversity conservation programmes.

Approximately 471bp partial sequences from mitochondrial 16S rRNA using seven Macrobrachium species were collected during the present study. Six of them were collected from the southern part of Sri Lanka and their phylogenetic positions among the relevant species that have been recorded within the region were determined. The analysis resulted in five clades, of which three showed monophyletic lineages (M. australe group, M. latimanus group, and M. latidactylus group). M. malcolmsonii is in a sister clade to M. rosenbergii while M. scabriculum joined with M. idae. The estimated intraspecific nucleotide divergence level varied from 0 - 6.07% while it varied from 5.21 - 10.84% at the interspecific level. The phylogenetic positions of the samples recorded within the region are discussed.

Keywords: Freshwater prawns, *Macrobrachium*, mitochondrial 16S gene region, phylogenetic relationships, Sri Lanka.

INTRODUCTION

Freshwater prawns of the genus *Macrobrachium* Bate, 1868¹ are important macro invertebrates in freshwater and estuarine systems throughout the tropical and warm temperature areas of the world. This genus can be ecologically separated into two groups. Most species are widely distributed and require specific saline concentrations to complete larval development (euryhaline species) and others are land locked species

with limited distributions and complete their entire life cycle in freshwater^{2,3}. So far more than 200 species have been described around the world and there are more to be described. *Macrobrachium* species have a high commercial value in the fisheries sector ⁴. A study⁵ listed 86 *Macrobrachium* species that are economically important and among them at least 11 species have gained great commercial value in different countries. The giant freshwater prawn *M. rosenbergii* ⁶ is farmed commercially, both within its natural range and outside.

Taxonomically, this genus is one of the most challenging decapod crustacean groups. The most distinguishable morphological characters of this genus are the rostrum and the second percopod, which are highly variable among species⁷. Most of the studies on Macrobrachium species are based on morphology^{2,3,5,7,8} and in recent years few studies have been published using molecular data. Some previous studies 9-12 have produced important clarifications for Australian Macrobrachium species based on 16S mitochondrial gene region. Further, based on the same gene region another study ³ suggested multiple origins of Macrobrachium species, region wise. Subsequently, two researchers^{14,15} carried out phylogenetic studies using multiple gene regions. For these studies data were collected from a wide geographical range mainly from the south and south-eastern Asian regions. However, Sri Lankan Macobrachium samples were not included. Collectively, the above studies produced interesting information and insight regarding Macrobrachium taxonomy.

In Sri Lanka, *Macrobrachium* species are an economically important group. Many species are

recognized by their local names and many synonyms can be found for one species, which makes studies more complicated. Therefore, identification of species boundaries within this genus is important to obtain information for applications in aquaculture and biodiversity conservation.

Two detailed taxonomic studies are recorded from Sri Lankan *Macrobrachium* species. One study¹⁶ described six species from different geographic locations and the other⁸ described twelve species collected from all over the country. However, these recorded species are only briefly described and their morphological characters are not properly illustrated using figures. To date no taxonomic studies at molecular level have been conducted with this genus in Sri Lanka. Therefore, the objective of the present study was to collect partial sequences of mitochondrial 16S ribosomal gene region from seven *Macrobrachium* species collected mainly from the southern part of Sri Lanka to determine their phylogenetic positions among the relevant species that have been recorded within the region.

METHODS AND MATERIALS

Seven *Macrobrachium* species were identified based on morphological features⁴ and used in this study. Among them 6 species (*M. rosenbergii*, *M. scabriculum*¹⁷, *M. idea*¹⁷, *M. australe*¹⁸, *M. latimanus*¹⁹ and *M. latidactylus*²⁰) were collected from the southern part of Sri Lanka mainly from the streams and small rivers of 2 major river basins: Nilwala and Walawe. *M. malcolmsonii* ²¹ was collected from the Gal Oya river system in the eastern part of Sri Lanka (Figure 1). Three to four specimens from each species were used to extract DNA. Reference samples were stored at the Department of Zoology, University of Ruhuna, Matara for further studies. Sampling localities are given in Table 1.

DNA was extracted using the easy DNA extraction kit (QIAGEN, USA). A fragment of the 16S rRNA mitochondrial gene was amplified by Polymerase Chain Reaction (PCR) using primers 1471- 5'CCTGTTTANCAAAAACAT3' and 1472-

Table 1: Sampling localities and gene bank accession numbers for sequences

Species (abbre:)	Location	Sample code	Genbank accession numbers
M. rosenhergii	Sri Lanka ¹	SriA	FJ595480*
(M. rose)	Sri Lanka ^{1,2}	SriB	FJ595481*
()	India	Ind	DO004836
	Thailand	Tha	AY203908
	Papua New Guinea	Pan	AY203906
	Australia	Aus	AY203918
M malcolmsonii	Sri Lanka ³	Sri	GU987055*
(M male)	India	Ind	AY730050
M scabriculum	Sri Lanka ¹	Sri	GU987059*
(M scab)	India	Ind	AY730055
M idea	Sri Lanka ^{1,2}	Sri	GU987058*
(M idae)	Taiwan	Twn	DO194930
(minut)	Australia	Aus	AY282777
M. australe	Sri Lanka ¹	Sri	GU987057*
(M.aust)	Philippine	phi	DO194905
()	Taiwan	Twn	DO194904
	New Guinea	New	DO681290
M. latimanus	Sri Lanka ^{1,2}	Sri	GU987056*
(M.lati)	Japan	Jap	DO194938
()	Taiwan	Twm	DO194936
	Philippine	Phi	DO194937
M. latidactvlus	Sri Lanka ¹	Sri	GU987060*
(M.latid)	Taiwan	Twn	EU493140
()	Thailand	Tha	DO194946
	Philippine	phi	DO194945
	China	Cha	DO194943
	Malaysia	Mal	DO194944
Palaemon			FM986647
(Palmon)			

* Sequences derived from this study

Sampling localities: 1-Nilwala river basin, 2- Walawe river basin, 3- Gal Oya basin (refer to Figure 1)

	28																												.38 -
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	26																										'	2 0.2	5 16.
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	24																									2.13	2.13	1.95	16.8
le 1.	23																								2.38	0.65	0.21	0.43	16.6
in Tabl	22																							1.95	1.29	1.73	1.73	1.52	16.4
given	21																						7.38	7.59	7.38	7.38	7.38	7.16	17.4
ns are	20																					0.22	7.59	7.81	7.59	7.59	7.59	7.38	17.7
viation	19																				0.22	0.00	7.38	7.59	7.38	7.38	7.38	7.16	17.4
Abbre	18																			2.82	3.04	2.82	7.59	7.81	7.59	7.59	7.59	7.38	16.6
egion.	17																	,	8.05	7.83	8.05	7.83	8.26	9.34	8.04	8.69	9.13	8.91	18.1
gene r	16																	2.39	8.48	8.26	8.48	8.26	9.13	9.35	8.92	9.35	9.35	9.57	19.2
e 16S	15																0.00	2.39	8.48	8.26	8.48	8.26	9.13	9.35	8.92	9.35	9.35	9.57	19.2
o for th	14															1.52	1.52	1.30	8.23	8.05	8.26	8.05	8.91	9.78	8.69	9.13	9.56	9.35	19
group	13													,	9.77	9.53	9.53	9.53	9.32	8.88	9.10	8.88	9.54	9.97	9.11	9.76	9.76	9.54	17.2
nd out	12													0.86	8.90	8.69	8.69	9.12	8.88	8.45	8.67	8.45	9.10	9.54	8.67	9.32	9.32	9.10	16.9
tions a	Ξ												0.00	0.86	8.90	8.69	8.69	9.12	8.88	8.45	8.67	8.45	9.10	9.54	8.67	9.32	9.32	9.10	17
popula	10											6.73	6.73	6.52	10.7	10.2	10.2	10.7	8.46	7.82	8.03	7.82	8.23	8.44	8.23	8.66	8.66	8.44	18.7
mong	6										0.43	6.96	6.96	6.75	10.9	10.5	10.5	10.9	8.72	7.63	7.84	7.63	8.47	8.63	8.47	8.91	8.9	8.69	18.7
nce) a	8									8.06	7.82	8.03	8.03	8.46	10.2	10.2	10.2	9.76	9.54	8.68	8.89	8.68	8.89	8.89	8.89	9.11	9.11	8.89	19
o' dista	٢								0.00	8.06	7.82	8.03	8.03	8.46	10.2	10.2	10.2	9.78	9.54	8.68	8.89	8.68	8.89	8.89	8.89	9.12	9.12	8.89	19
cted ' ₁	9							5.21	5.21	9.79	9.55	8.03	8.03	8.68	10.2	9.56	9.56	9.99	9.11	8.46	8.68	8.46	10.2	9.75	86.6	86.6	9.54	9.75	19.8
incorre	5						0.65	5.42	5.42	10.22	9.99	8.46	8.46	9.11	9.99	9.78	9.78	10.21	9.33	8.68	8.89	8.68	10.19	9.54	9.93	9.75	9.32	9.54	19.84
ence (ı						86	21	.07	.07	58	.12	59	59	.03	0.21	66	66	0.21	33	0.41	0.63	0.41	0.41	0.84	0.41	0.19	0.62	0.41	0.32
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uence	3			'	0.0	5.8	5.2	6.0	6.0	9.5	9.1	7.5	7.5	8.0	10.	9.6	9.6	10.	9.3) 10.	1 10.) 10.) 10.	2 10.) 10.	10.	1 10.	9 10.	4 20.
of seq	7			0.22	0.22	6.07	5.42	6.29	6.29	9.79	9.33	7.82	7.82	8.25	96.99	9.78	9.78	96.99	9.11	10.19	10.4	10.19	10.19	10.62	10.19	9.97	10.4	10.19	20.5
ntage o	1		0.22	0.00	0.00	5.86	5.21	6.07	6.07	9.58	9.12	7.59	7.59	8.03	10.21	9.99	9.99	10.21	9.33	10.41	10.63	10.41	10.41	10.84	10.41	10.19	10.62	10.41	20.32
Perce		e,SriA	e,SriB	e,Ind	e, Tha	e, Pap	e, Aus	lc, Sri	lc, Ind	ıb, Sri	ıb, İnd	e, Sri	e,Twn	e, Aus	st, sri	st,Phi	st,Twn	s,New	i,Sri	Jap.	,Twn	.Phi	d,Sri	d,Twn	d,Tha	d,Phi	d,Cha	d,Mal	u
ıble 2:		M.ros	M.ros	M.ros	M.ros	M.ros	M.ros	M.ma	M.må	M.scá	M.scá	M.ida	M.ida	M.ida	M.au	M.au	M.au	M.au	M.lat.	M.lati	M.lat	M.lati	M.lati	M.lati	M.lati	M.lati	M.lati	M.lat	Palmo
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Macrobrachium species in Sri Lanka



Figure 1: Sampling localities of the current study



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5'AGATAGAAACCAACCTGG3' described in a previous study²² which were initially designed for freshwater crayfish. Double stranded PCR products were obtained in a total reaction volume of 25 µL, containing 5 µL of 10X PCR buffer, 0.4 mM of each dNTP, 0.8 µM of each primer, 4 mM MgCl,, 1 unit of Taq polymerase and 2 µL of DNA extract. PCR amplification was carried out using the following temperature regime: an initial denaturation step of 95°C for 5 min, followed by 30 cycles of 95°C for 30 s, an annealing temperature of 50°C for 30 s and an extention of 72°C for 30 s. This was followed by an additional extension of 72°C for 3 min. PCR products were purified using a QIAGEN QIAquick PCR purification kit, with a final elution volume of 50 µL per individual. The quality of the PCR products was visualized on 1.5% agarose gels. Sequencing was carried out in both directions using the same primer pairs for PCR. Sequences were obtained using the Big Dye Terminator version 3.1 protocol (Applied Biosystems, USA) and analyzed using an ABI 3130xl Genetic Analyzer (with KB base caller; Applied Biosystems, USA).

Sequence chromatograms were viewed and edited manually using a combination of Edit View and SeqPup programmes²³. Once edited, multiple alignments were performed using Clustal X programme²⁴ with multiple alignment parameters of gap penalty equal to 10-15 and gap extension penalty equal to 3-5. Positions of uncertain alignment were excluded to produce a stable data set. Sequences were then imported into PAUP 4.0b10 programme²⁵ for phylogenetic analysis. Additional sequences for the recorded species within the region and out group species ²⁶ were collected from Genebank (Table 1) and used to justify the phylogenetic positions of Sri Lankan species among them. Sequences derived for this study were deposited in Genebank (Accession number GU987055 - GU987060 and FJ595480 -FJ595481) (Table 1).

Bayesian analyses (BA) were performed with MrBayes version 3²⁷ using the model selected by MrModeltest programme²⁸. Markov chain Monte Carlo (MCMC) chains were run for 1x10⁶ generations, and trees were saved each 100 generations (with the 1st 1000 trees being discarded as 'burn in'). The probability values greater than 95% were considered as significant support for relationships. The neighbour-joining analyses [Minimum Evolution option (ME)] were performed using distance calculated under the same model of evolution as the Bayesian analysis. Maximum Parsimony (MP) analyses were performed with gaps treated as missing data and heuristic search option was used with tree bisection-reconnection (TBR) branch swapping and 100 stepwise random additions. Bootstrapping was performed with

1000 replicates for all analyses. The level of support for each analysis is indicated on clads of the phylogenetic tree.

RESULTS

Sequences of approximately 471bp in length were obtained from 16S rRNA gene. The mean nucleotide composition was A= 36.58%, T= 30.57%, C=22.57%, G=10.28%. This indicates that the 16S rRNA region of the mtDNA is the adenosine and thymine rich in the palaemonids. On the basis of the ModelTest, the GTR+I+G model of sequence evolution was chosen and the parameters specified by this model [(A-C)=1.1231, (A-G) 20.5595. (A-T)=2.0745, (C-G)=0.0001, (C-T)=9.5918, (G-T)=1.0000 and gamma distribution parameter=0.2301] were used for further analysis. Pairwise distances (uncorrected 'p' distance) for the data set are given in Table 2.

In this study, the estimated intraspecific nucleotide divergence level varied from 0 - 6.07% while it varied from 5.21 - 10.84% for the interspecific level. All three methods of phylogenetic analysis (BA, MP, ME) produced identical tree topologies. Five clades were derived: *M. australe* group, *M. malcolmsonii* + *M. rosenbergii* group, *M. latimanus* group, *M. latidactylus* group and *M. scabriculum* + *M. idae* group. However, the generated dendogram did not resolve the deeper level phylogenetic relationships (Figure 2).

DISCUSSION

This study was based on mitochondrial 16S rRNA gene region of seven *Macrobrachium* species collected from different geographical regions. The analysis resulted in five clades and four of them produced monophyletic lineages. *M. malcolmsonii* made a sister clade to *M. rosenbergii* while *M. scabriculum* joined with *M. idea* (Figure 2).

Congeneric crustacean species commonly exhibit significant differences at the 16S rRNA mtDNA gene ranging from 2 - 17% sequence divergence²⁹⁻³². This divergence range is also supported by other recent studies conducted on *Macrobrachium* species^{9-11,14}. Therefore, the divergence levels that became evident in this study between and among species are typical for those observed between crustacean species (Table 2).

M. australe showed the lowest intraspecific nucleotide divergence that varied from 0 - 2.4%. *M. australe* group is defined with high support and

contains two clades. The southeast Asian samples were grouped together while Sri Lankan *M. australe* is grouped with New Guinean sample with lower support (0.77). The nucleotide divergence level between Sri Lankan and New Guinean sample was 1.3% while the Sri Lankan sample varied from the others with 1.5% divergence level.

M. malcolmsonii formed a sister clade which was basal to M. rosenbergii. The nucleotide divergence level between the two species varied from 5.2 - 6.4%. Two haplotypes were found within Sri Lankan M.rosenbergii. The M. rosenbergii samples collected from the South and South-East Asia were grouped together to accept the hypothesis of eastern and western division of this species along the Huxley's line suggested by a previous study 33. The neucleotide sequence divergence between the two clades varied from 5.2 - 6%. This value was quite similar to the divergence level found between M. rosenbergii and M. malcolmsonii samples. M. malcolmsonii is so far reported from the South Asian region and it is worthwhile to conduct further phylogenetic studies using more gene regions to reveal their relationship.

In M. latimanus group, the Sri Lankan sample was positioned basal to the other samples. This separation of Sri Lankan M. latimanus from other samples (nucleotide divergence level 3%) was greatly supported by high bootstrap values. This result was also supported by a previous study14, which reported up to 3.2% intra specific sequence divergence levels for Macrobrachium species. Within the M. latidactylus group, Sri Lankan and Thailand samples have given basal support to the clade. The Sri Lankan sample differed from the inner group by 1.5 - 2.% nucleotide divergence level while this level was 1.2 - 2.2% for the Thailand sample. M. idae made a sister clade to M. scabriculum with lower support. Within the M. idae clade, two Asian samples were similar to each other and showed 0.9% nucleotide divergence level from the Australian sample. M. scabriculum is so far reported from South Asian region and the Sri Lankan sample differs from the Indian sample by 0.4% nucleotide divergence level.

Detailed morphological study of Sri Lankan *Macrobrachium* taxonomy has been conducted previously⁸. In this study, many species have been collected from outside the southern Sri Lanka. From the twelve species described in the above study, the present study has found six species in the southern part of Sri Lanka. However, the previous studies have not highlighted the phylogenetic relationships of the species. The current study is the first molecular based study of the genus *Macrobrachium* in Sri Lanka. More taxon

sampling with additional gene regions are required to better understand the phylogenetic relationships among Sri Lankan samples.

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