Genetic variations and phylogenetic relationship of genus *Uromastyx* from Punjab Pakistan

N. Ismat1 Ø, S. Malik1 Ø, M. Rashid2 Ø, A. Javid1* Ø, A. Hussain1 Ø, S. M. Bukhari1 Ø, S. Suleman1 Ø, R. Noor1 Ø, S. Husaain1 Ø, N. Ismat1 Ø, M. Hussain1 Ø, S. Ghafoor1 Ø, G. Mustafa1 Ø and W. Ali1 Ø

1University of Veterinary and Animal Sciences, Department of Wildlife and Ecology, Lahore, Pakistan
2University of Veterinary and Animal Sciences, Faculty of Fisheries and Wildlife, Lahore, Pakistan
3The University of Lahore, Institute of Molecular Biology and Biotechnology – IMBB, Pakistan

Abstract

During the present study, specimens were collected from selected sites of Cholistan desert and Kalabagh Game Reserve, Punjab province, Pakistan. Each captured specimen was tagged with voucher number and morphometric measurements were taken. The average snout to vent length was 172.559±1.40 mm and average weight was 92.1±1.30 g. The DNA of *Uromastyx hardwickii* was amplified and sequenced using 16S rRNA primer set. The obtained DNA sequence has shown reliable and clear species identification. After trimming ambiguous bases, the obtained 16S rRNA fragment was 520 bp while 16S rRNA fragments aligned with closely matched sequence from NCBI comprised of 510 bp. Closely matched sequences of genus *Uromastyx* were retrieved from NCBI in blast searches. Neighbour-joining tree of genus *Uromastyx* was constructed based on p-distance using MEGA X. The mean intraspecific variation was 0.095±0.01 while intraspecific variation was ranging from 0-1%. Similarly, interspecific variation of *Uromastyx hardwickii* with *Saara asmussi, Uromastyx alfredschmidti, Uromastyx geyri, Uromastyx thomasi, Uromastyx alfredschmidti* was 0-12%, 0-19%, 0-19%, 0-20%, 12-19% respectively. The newly produced DNA was submitted to NCBI and accession number was obtained (MW052563.1). Results of current study provided information about the molecular and morphological identification of Genus *Uromastyx*. In our recommendation, comprehensive molecular based identification of Pakistan’s reptiles is required to report any new or subspecies from country.

Keywords: Kalabagh Game Reserve, *Uromastyx hardwickii, Uromastyx asmussi*, Palearctic region, IUCN redlist.

Resumo

Durante o presente estudo, os espécimes foram coletados em locais selecionados do deserto do Cholistan e da Reserva de Caça de Kalabagh, província de Punjab, Paquistão. Cada espécime capturado foi etiquetado com o número do comprovante e medidas morfométricas foram realizadas. O comprimento médio do focinho à cloaca foi de 172.559 ± 1.40 mm, e o peso médio foi de 92.1 ± 1.30 g. O DNA de *Uromastyx hardwickii* foi amplificado e sequenciado usando o conjunto de primer 16S rRNA primer set. A sequência de DNA obtida mostrou identificação de espécies confiável e clara. Após o corte de bases ambiguas, o fragmento de rRNA 165 obtido tinha 520 pb, enquanto os fragmentos de rRNA 16S alinhados com a sequência próxima do NCBI composta por 510 pb. Sequências semelhantes do gênero *Uromastyx* foram recuperadas do NCBI em pesquisas de explosão. A árvore de união de vizinhos do gênero *Uromastyx* foi construída com base na distância-p usando MEGA X. A variação intraespecífica média foi de 0.095 ± 0.01, enquanto a variação intraespecífica foi de 0-1%. Da mesma forma, a variação interspecífica de *Uromastyx hardwickii* com *Saara asmussi, Uromastyx alfredschmidti, Uromastyx geyri, Uromastyx thomasi, Uromastyx alfredschmidti* foi de 0-12%, 0-19%, 0-19%, 0-20%, 12-19% respectivamente. O DNA recém-produzido foi submetido ao NCBI e o número de acesso foi obtido (MW052563.1). Os resultados do estudo atual forneceram informações sobre a identificação molecular e morfológica do Gênero *Uromastyx*. Em nossa recomendação, a identificação de base molecular abrangente de répteis do Paquistão é necessária para relatar qualquer nova ou subsespécie do país.

Palavras-chave: Reserva de caça de Kalabagh, *Uromastyx hardwickii, Uromastyx asmussi*, região paleártica, lista vermelha da IUCN.
1. Introduction

The changes that reptiles faced over millions of years results in massive diversity in their morphology, behavior, ecology history of life and in strategies against their predators. Due to all these features reptiles in terms of evolutionary and ecological research now become the central model system (Rasmussen et al., 2011). We used the term paraphyletic group for reptiles which include all non-avian taxa such as Sphenodontia, Squamata, Testudines and Crocodylia in this group. The total numbers of living reptile species are approximately 9546, out of these the crocodiles are 25 (0.3%), turtles are 327 (3.4%) and tuatara is 1 (0.01%) and the remaining 9,193 (96.3%) are the squamates species (lizards, snakes and amphisbaenians) (Hay et al. 2010).

Reptiles are the mixture of Palearctic, Ethiopian and Indo-Malayan forms in Pakistan and are represented by 195 reptiles, and consists of 23 families including Dermochelyidae, Chelonidae, Emydidae, Testudinidae, Trionychidae, Gavialidae, Crocodylidae, Agamidae, Chamaeleonidae, Gekkonidae, Eublepharidae, Scincidae, Lacertidae, Varanidae, Uromastycidae, Tylopholidae, Leptotyphlopidae, Colubridae, Boidae, Elapidae, Hydrophididae, Elapidae, Crotales and Viperidae (Khan et al. 2010). Out of these thirteen species are endemic to country. The Lizards are the prominent group of reptiles in Pakistan and represented by eight families viz., Scincidae, Uromastycidae, Varanidae, Agamidae, Chamaeleonidae, Eublepharidae, Lacertidae and Gekkonidae (Khan, 2004, 2006). According to IUCN (IUCN Red List of Threatened Species, 2009) 1,677 reptiles are included in Red List. Out of these, 469 species are near to extinction and 22 are already extinct. Furthermore, only 40% of reptiles’ species have their Conservation status and about 4,000 were not assessed yet (IUCN Red List of Threatened Species, 2009).

The identification and taxonomic classification of reptiles is very challenging task. Recently, a molecular technique which is used for the identification is known as DNA barcoding which is used for finding phylogenetic relationships. Before 2005 DNA barcoding of amphibians and reptiles was very difficult as compared to other taxa such as fishes, birds and mammals (Hebert et al., 2004). In DNA barcoding, a short sequence of DNA is compared with reference database for the identification of species (Hajibabaei et al., 2007). The genomic regions that are mostly used include 12S rRNA, 16S rRNA, Cytb and COI consisting (Hebert et al., 2004; Nagy et al., 2012).

Habitat loss and illegal trading have effect the survival and existence of many reptile species worldwide especially in Pakistan. One of them is Indian spiny-tailed lizard (Uromastyx hardwickii). U. hardwickii is medium to large size lizard reported from North Africa to North Western Indian desert. Spiny tailed lizard is ground dweller and herbivorous. In Pakistan, it is distributed in Southern Baluchistan, Indus valley, extended to Las bela and Chagai desert. The major threats to these species is its commercial exploitation for meat, skin and oil. About 367,000 specimens were legally traded between 1977 and 2005 (Knapp, 2004).

Genus Uromastyx is represented by 17 species. Most of species within genus Uromastyx were identify on the basis of external morphology and heir phylogenetic relationship is still under debate (Amer and Kumazawa, 2005; Wilms and Schmitz, 2007). Pakistan is represented by two species of genus Uromastyx namely Uromastyx hardwickii and Uromastyx asmsussi. Present study was therefore planned to identify members of genus Uromastyx through mitochondrial genes and sort out taxonomy problems of these taxa.

2. Materials and methods

2.1. Sample collection

A total of 10 specimens are collected from selected sites of Cholistan desert and Kalabagh Game Reserve, Punjab province. Each captured specimen was tagged with voucher number and morphometric measurements were taken following Ali et al. (2017). A few specimen of each sampling site (n=3) were euthanized and preserved in 75% ethanol for molecular characterization.

2.2. Morphometric measurement

The following morphometric measurement were taken SVL=snout vent length, TAL=tail length, HL=head length, HW=head width, BW=body width, IIL=interlimb length, FC=fore claw length, HC=hind claw length, HLS=hind limb Span, FLS=forelimb span, HLF=hind limb longest finger, FLF=forelimb longest finger, W=weight and TL=total length (Ali et al. 2017).

2.3. DNA extraction and amplification

DNA was extracted using phenol chloroform method (Ali et al., 2020). The quality of DNA was checked on 1.5% agarose gel in the post-graduate lab, Department of Wildlife and Ecology, University of Veterinary and Animal Sciences, Pakistan. The amplification was done by using cytochrome oxidase I (COI) and 16S rRNA primers sets. PCR reaction was performed in 0.2 ml PCR tube and 25µL reaction mixtures. To prepare 25 µL reaction mix 6 µL double distilled water, 1 µL (25 mM) Primer F,1µL (25 mM) primer R, 12 µL Master mix and 5 µL of DNA template was mixed. In each reaction a negative control was also run using sterilized water as the template. The following steps were performed for the amplification of gene, 3 minutes denaturing at 94°C followed by 40 cycles of for 30 sec at 94°C, primer annealing for 30sec at 42-55°C depend on the primer’s annealing temperature and elongation for 1 minutes at72°C, with final 10 minutes at 72°C and infinity.

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2.4. Data analysis

The obtained DNA sequences were analyzed and edited in Bioedit software 7.0 and aligned using Clustal X.
Genetic variations of genus *Uromastyx* (Hussain et al., 2020; Tamura et al., 2013). The consensus sequence of each sample were subjected to BLAST (basic local alignment tool search) analysis to compare the percentage identity of sequence by searching in the public DNA databases. The closely related sequences were obtained and incorporated in the neighbor-joining (NJ) tree analyses with bootstrap value of 100 replicates using MEGA 10. Genetic distances within and between species were calculated using Mega 6.0 based on p-distance.

3. Results

The specimens of *Uromastyx hardwickii* (Figure 1) were collected from selected sites of District Bahawalnagar and Kalabagh Game Reserve, Punjab province, Pakistan (Figure 2) during field surveys extended from May to August, 2020.

3.1. Morphological identification and measurements

The average snout to vent length was 172.559±1.40 mm, tail length was 126.55±0.9 mm, head length was 33.05±1.9, head width was 24.75±1.1, body weight was 42.6±1.4, IIL was 33.175±1.6, FC was 18.675±1.1, HC was 27.525±1.4, HLS was 126.525±1.2, FLS was 17.625±0.8, HLF was 17.625±0.8, FLS was 12.1±2.2 and average weight was 92.1±1.30 g (Table 1).

The description of captured specimens from study area is as follows:

![Figure 1. *Uromastyx hardwickii* captured from District Bahawalnagar and Kalabagh Game Reserve, Punjab province, Pakistan.](image)

![Figure 2. Map of study area.](image)
I. Genus *Uromastyx* (Merrem, 1820)

The subfamily Uromastyicinae consist of two genera Saara Gray, 1845 and Uromastyx Merrem, 1820. Within the Uromastyicinae, Uromastyx is the largest genus. Its habitat is desert and semi-desert area. With the passage of years, its taxonomy has been changed since 1980 and includes four species and three subspecies. The members of Uromastyicinae are commonly called spiny-tailed lizards as their tail is covered with spines. On the basis of presence and absence of intercalaries between the tail and whorls, two genera, *Saara* and *Uromastyx* are differentiated. Pakistan is represented by two species of the genus *Uromastyx* namely *Uromastyx hardwickii* and *Uromastyx asmussi*.

i. *Uromastyx hardwickii*

*Uromastyx hardwickii* commonly called spiny-tailed lizard and found in dry regions of northwest India and Pakistan. It also found in Afghanistan. Its maximum SVL is 233 mm. Around the mid body, 190–275 scales are found and 24–42 scales are present near the ear opening on the fourth left toe. 15–21 scales are present. On either side, 12–19 pre-anofemoral pores are present. Between the gular- and inguinal fold, 112–157 scales are present. The *U. hardwickii* is medium to large size lizard reported from North Africa to North Western Indian desert. Spiny-tailed lizard is a ground dweller and herbivorous. In Pakistan, it is distributed in Southern Baluchistan, Indus valley, extended to Las Bela and Chagai desert. The major threats to this species are its commercial exploitation for meat, skin, and oil. About 367,000 specimens were legally traded between 1977 and 2005.

3.2. Amplification and sequencing

During the present study, DNA of *Uromastyx hardwickii* was amplified and sequenced using 16S rRNA primer set. The obtained DNA sequence showed reliable and clear species identification. After trimming ambiguous bases, the obtained 16S rRNA fragment of *Uromastyx hardwickii* was 520 bp while 16SrNRA fragments aligned with NCBI sequences comprised 510 bp (Table 2). The newly produced DNA was submitted to NCBI and accession number was obtained (MW052563.1).

3.3. Genetic diversity and variation

The mean intraspecific variation was 0.095±0.019 while intraspecific variation of *Uromastyx hardwickii* was ranging from 0-1%. Similarly, interspecific variation of *Uromastyx hardwickii* with *Saara asmussi*, *Uromastyx alfredschmidti*, *Uromastyx geyri*, *Uromastyx thomasi*, *Uromastyx alfredschmidti* was 0-12%, 0-19%, 0-19%, 0-20%, 12-19% respectively (Table 3).

3.4. Phylogenetic analysis

Recently few DNA barcoding studies of Asian amphibians and reptiles have been carried out and sequences for related species were available at NCBI. Closely matched sequences

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**Table 1. Weight (g) and external body measurements (mm) of *Uromastyx hardwickii* captured from study area.**

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<th>Specimen 3</th>
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<th>Specimen 5</th>
<th>Specimen 6</th>
<th>Specimen 7</th>
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**Table 2. The details of successful amplified specimens and Genbank accession number.**

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<th>Species</th>
<th>Voucher number</th>
<th>GenBank accession number</th>
</tr>
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<td>Uromastyx hardwickii</td>
<td>ZMUVAS40</td>
<td>MW052563.1</td>
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</table>
Genetic variations of genus Uromastyx were retrieved from NCBI in blast searches. Neighbour-joining tree of genus Uromastyx was constructed based on 16S rRNA sequence using MEGA X (Figure 3). The analysis involved 11 nucleotide sequences. All ambiguous positions were removed for each sequence pair. There were a total of 510 positions in the final dataset.

4. Discussion

The information related to DNA barcoding of reptiles is very less as compared to other animals. The genes that are used for this purpose is Cytochrome C Oxidase I (COI), cytochrome b (Cytb), 12S rRNA and 16S rRNA (Nicolas et al., 2012; Xia et al., 2012). In the past due to the problems that researcher faced in case of COI sequences amplification and evaluation of reptiles and amphibians prefer to use the 16S ribosomal RNA gene for DNA barcoding. Vasconcelos et al. (2016) used COI gene to find the intra-specific of reptiles which was ranging between 13.8– 54.4%. Sharma et al. (2018) isolate the DNA from the feces of Indian Spiny Tailed Lizard Saara hardwickii. They carried out research in Rajasthan in June 2016 and used the 16S rRNA gene in order to find out genetic variation in S. hardwickii. Phylogenetic relationship indicated that due to illegal trading, India and
global samples shared ancestry as at seven bp insertion (126-128) and deletion present at 2(49-252).

Arida (2017) used different mitochondrial DNA regions for the identification of monitor lizard in Indonesia. Previous Studies on the Varanus genus shows that the mitochondrial markers that were used on the biogeography and on systematics were 12s rRNA and 16s rRNA. The combination of nuclear and mitochondrial genes also used by others (Ziegler et al., 2007; Fuller et al., 1998; Fitch et al., 2006; Doughty et al., 2014; Francis et al., 2010; Nagy et al., 2012).

Adam et al. (2009) used mitochondrial DNA sequences of about 1,181 bp to described phylogeny of nineteen species of African lizards. The species of East and West Africa has monophyletic radiations which were supported along with a clade that contain 2 species of sabah region.

5. Conclusions and recommendations

During present study, different mitochondrial genes COI and 16s rRNA were used to identify genus Uromastyx from study area. The DNA of Uromastyx Hardwickei successfully amplified using 16s rRNA primer set. Results of current study provided information about the molecular and morphological identification of Genus Uromastyx. Comprehensive molecular based identification of Pakistan’s reptiles is required to report any new or subspecies from country.

References


