This article was downloaded by: [Ferdowsi University of Mashhad]
On: 15 November 2010
Access details: Access Details: [subscription number 912974446]
Publisher Taylor \& Francis
Informa Ltd Registered in England and Wales Registered Number: 1072954 Registered office: Mortimer House, 3741 Mortimer Street, London W1T 3JH, UK

## NATURAL <br> HISTORY

## Journal of Natural History

Publication details, including instructions for authors and subscription information:
http://www.informaworld.com/smpp/title $\sim$ content=t713192031

## Phylogenetic relationships of Mesobuthus eupeus (C.L. Koch, 1839) inferred from COI sequences (Scorpiones: Buthidae)

Omid Mirshamsia ${ }^{\text {ab }}$; Alireza Saria; Elahe Elahiac; Shidokht Hosseinie ${ }^{\text {d }}$
${ }^{\text {a }}$ School of Biology, College of Science, University of Tehran, Tehran, Iran ${ }^{\text {b }}$ Department of Biology, Faculty of Sciences, Ferdowsi University of Mashhad, Mashhad, Iran ${ }^{\text {c }}$ Centre of Excellence in Biomathematics Statistics and Computer Science, College of Science, University of Tehran, Tehran, Iran ${ }^{\text {d Biology Department, Faculty of Sciences, Shiraz University, Shiraz, Iran }}$

Online publication date: 15 November 2010

To cite this Article Mirshamsi, Omid, Sari, Alireza, Elahi, Elahe and Hosseinie, Shidokht(2010) 'Phylogenetic relationships of Mesobuthus eupeus (C.L. Koch, 1839) inferred from COI sequences (Scorpiones: Buthidae)', Journal of Natural History, 44: 47, $2851-2872$
To link to this Article: DOI: 10.1080/00222933.2010.512400
URL: http://dx.doi.org/10.1080/00222933.2010.512400

## PLEASE SCROLL DOWN FOR ARTICLE

```
Full terms and conditions of use: http://www.informaworld.com/terms-and-conditions-of-access.pdf
This article may be used for research, teaching and private study purposes. Any substantial or
systematic reproduction, re-distribution, re-selling, loan or sub-licensing, systematic supply or
distribution in any form to anyone is expressly forbidden.
The publisher does not give any warranty express or implied or make any representation that the contents
will be complete or accurate or up to date. The accuracy of any instructions, formulae and drug doses
should be independently verified with primary sources. The publisher shall not be liable for any loss,
actions, claims, proceedings, demand or costs or damages whatsoever or howsoever caused arising directly
or indirectly in connection with or arising out of the use of this material.
```


# Phylogenetic relationships of Mesobuthus eupeus (C.L. Koch, 1839) inferred from COI sequences (Scorpiones: Buthidae) 

Omid Mirshamsi ${ }^{\text {a,b* }}$, Alireza Sari ${ }^{\text {a }}$, Elahe Elahia ${ }^{\text {a,c }}$ and Shidokht Hosseinie ${ }^{\text {d }}$<br>${ }^{a}$ School of Biology, College of Science, University of Tehran, Tehran, Iran; ${ }^{b}$ Department of Biology, Faculty of Sciences, Ferdowsi University of Mashhad, Mashhad, Iran; ${ }^{c}$ Centre of Excellence in Biomathematics Statistics and Computer Science, College of Science, University of Tehran, Tehran, Iran; ${ }^{\text {dBiology Department, Faculty of Sciences, }}$ Shiraz University, Shiraz, Iran

(Received 15 March 2010; final version received 21 July 2010)


#### Abstract

In this study, the first molecular phylogenetic analysis of Mesobuthus eupeus in Iran is presented based on sequence data of a $\sim 700$-base-pair fragment of cytochrome C oxidase, subunit I. Phylogenetic relationships were inferred using parsimony, maximum likelihood and Bayesian inference. The results support monophyly of M. eupeus, but there is a clear divergence between northern and southern clades. The northern clade includes four subspecies - M. e. eupeus, M. e. philippovitschi, M. e. afghanus and M. e. thersites; whereas the southern clade is comprised of two others - M. e. phillipsi and M. e. kirmanensis. Accordingly, possible scenarios for the evolution and phylogeographic structure of M. eupeus based on the geological history of the Iranian Plateau were proposed. The observation of two distinct lineages supports the proposal that M. eupeus might be a species complex composed of species with highly similar morphological features.


Keywords: Mesobuthus eupeus; scorpions; Buthidae; subspecies; cytochrome C oxidase subunit I; Iran

## Introduction

Scorpions possess several characteristics notably different from other arthropods. Based on some traits including longevity, age to maturity, complex courtship behaviour, viviparous embryonic development, maternal care, post-embryonic development, low metabolic rate and a degree of social behaviour, they are more similar to long-lived vertebrates (Polis and Sissom 1990; Gantenbein and Largiadèr 2002; Lourenço 2000). Scorpions are traditionally considered to be living fossils and are known for being morphologically highly conserved animals (Sissom 1990). Although these unusual features have been well known for some time, biochemical and molecular methods have only recently been applied to this taxon. DNA sequence evidence was first presented by Gantenbein et al. (1999) for assessing the phylogeny of the genus Euscorpius. The phylogeographical studies on euscorpiids (Gantenbein et al. 1997, 1999, 2000; Gantenbein and Scholl 1998; Gantenbein, Fet and Barker 2001; Huber et al. 2001; Fet et al. 2003; Salomone et al. 2007) and buthids (Gantenbein et al. 1999, 2000, 2003, 2005; Gantenbein, Soleglad and Fet 2001; Gantenbein and Keightley 2004; Gantenbein and Largiadèr 2002, 2003; Parmakelis et al. 2006) using

[^0]nuclear and mitochondrial markers revealed that many previously described species are in fact constituted by sibling species not easily differentiable based on morphological features. In some instances extensive genetic divergence had evolved in the absence of a significant morphological differentiation. For example, on the basis of molecular evidence, Euscorpius (Euscorpius) carpathicus (Linnaeus 1767) is now considered a complex of at least seven sibling species (Gantenbein and Largiadèr 2002; Vignoli et al. 2005; Salomone et al. 2007).

Scorpions of the genus Mesobuthus Vachon, 1950 represent a useful terrestrial model for the study of molecular evolution (Gantenbein and Keightley 2004). This genus has recently been the subject of molecular studies for intra- and interspecific phylogeographic and phylogenetic studies (Gantenbein et al. 2003; Gantenbein and Keightley 2004; Parmakelis et al. 2006). The genus Mesobuthus was erected when Vachon (1950) initiated the revision and splitting of the traditional genus Buthus Leach, 1815. The taxonomic composition of this genus is still controversial and there is no consensus on the number of species comprising the genus Mesobuthus. Some described species may in fact be composed of sibling species, and still novel species are being described. For instance, separate species such as Mesobuthus cyprius from Cyprus (Gantenbein et al. 2000) and Mesobuthus songi from China (Lourenço et al. 2005) were described. In addition, Mesobuthus nigrocinctus has been recorded from Israel and Turkey (Fet et al. 2000; Teruel 2000; Karataş and Karataş 2001; 2003). It is widely accepted that the genus Mesobuthus includes at least 12 species (Fet and Lowe 2000; Gantenbein et al. 2000). The modern distribution of the genus Mesobuthus extends from the Balkans through China, and its occurrence in northern parts of central Asia represents the northern limit of scorpion distribution in Asia (Fet 1994; Gromov 2001).

The scorpion Mesobuthus eupeus (C. L. Koch, 1839) is the type species of the genus Mesobuthus. This species is the most widely dispersed species of the genus Mesobuthus and one of the most dispersed members of the family Buthidae. It occurs in eastern and central parts of Turkey, Armenia, Azerbaijan, Georgia, southern Russia, Syria, Iraq, Iran, Afghanistan, Pakistan, Central Asia, southern Mongolia and northern China (Birula 1917; Farzanpay 1987; Vachon and Kinzelbach 1987; Fet 1989; Kovařík 1997; Fet et al. 2000; Gromov 2001; Shi et al. 2007). This extensive geographic distribution is accompanied by morphological variations on the basis of which several subspecies have been described (Pocock 1889; Birula 1900, 1905, 1917; Vachon 1952). As early as 1917 Birula grouped the described subspecies of M. eupeus into two 'sections' or species groups, M. eupeus and M. thersites. Moreover, he recognized three "natio" within the nominotypical subspecies, M. e. eupeus. Among the described subspecies, 14 are considered formally valid (Fet and Lowe 2000). But the taxonomic status and relationships between subspecies of $M$. eupeus has not been recently examined and revisions may be appropriate. The morphological characteristics used for determination and assessing the relationships of these subspecies are inconclusive and vague (Birula 1900, 1905). With respect to the M. eupeus subspecies recorded from Iran, authors are in disagreement (Farzanpay 1987; Fet 1989) and some authors believe that M. eupeus is a species complex (Gantenbein et al. 2003).

Mitochondrial DNA markers can be used for resolving taxonomic ambiguities in M. eupeus by a molecular approach. Mitochondrial DNA sequences are used in the present study with the objective of describing the evolutionary lineages of this species. Additionally, the times of divergence between lineages are estimated. Finally, possible
processes that shaped the current distribution of M. eupeus in the Iranian Plateau are discussed.

## Materials and methods

## Sample collection

According to Prendini $(2001,2005)$ and Lamoral $(1979)$, members of the genus Mesobuthus are habitat generalist lapidicolous scorpions that shelter under stones or any other available covers. Here, most specimens were collected at daytime by rock rolling in the field and a few were caught at night using the ultraviolet light detection method (Lowe et al. 2003). A portable ultraviolet flashlight equipped with inidium gallium nitride light-emitting diodes was used for specimen collection at night. After collection, ethanol was injected into the specimens' bodies and they were preserved in $70-96 \%$ ethanol. Geographic coordinates of most collection sites were recorded using a hand held global positioning system (Garmin ${ }^{\mathrm{TM}}$ ). Geographic coordinates of a small number of sites were assessed by reference to the gazetteers and official topographic maps of Iran.

A distribution map of collection sites was created using ARCVIEW GIS 3.1 (Environmental System Research Institute, Redlands, CA, USA), by superimposing locality records on layers depicting political boundaries and topography. The topographic contour layer was based on the GTOPO30 raster grid coverage, available on the U.S. Government Public Information Exchange Resource at http://edc.usgs.gov/ products/elevation/gtopo30.

## Ingroup and outgroup taxa

Fifty-nine adult specimens of M. eupeus (Figure 1, Table 1) were collected from different localities in Iran. Additionally, 31 sequences were retrieved from GenBank (Table 1). Details of the origin of samples and the accession numbers of sequences used in this research or retrieved from databases are given in Table 1. All vouchers and DNA extracts have been deposited in the Zoological Museum, University of Tehran.

Three buthid species, Androctonus australis (Linnaeus 1758), Buthus occitanus (Amoreux 1789) and Buthus mardochei Simon, 1878 are frequently used as outgroups in buthid phylogenetic studies (Gantenbein and Largiadèr 2003; Gantenbein et al. 2003; Parmakelis et al. 2006). The species closest to these species that is found in Iran is the Old World buthid, Androctonus crassicauda and it was used as outgroup in the present study. Additionally, sequences of three other Mesobuthus species, M. gibbosus, M. cyprius and M. caucasicus were included (Gantenbein and Keightley 2004; Gantenbein et al. 2005).

## Molecular laboratory methods

DNA was isolated from the specimens, using the GenNetBio ${ }^{\mathrm{TM}}$ genomic DNA Extraction kit following the manufacturer's instructions (Seoul, South Korea). Polymerase chain reaction (PCR) was subsequently performed to amplify a fragment of approximately 700 base pairs (bp) of the mitochondrial cytochrome C oxidase, subunit I (COI) gene. Primers used were LCO1490: $5^{\prime}$-GGTCAACAAATCATCATA AAGATATTGG-3' (Folmer et al. 1994) and Nancy: 5'-CCCGGTAAAATTAAA


Figure 1. Collection sites of Mesobuthus eupeus (■). Triangles ( $\mathbf{\Delta}$ ) shown in (A), (B) and (C) denote origin of sequences of Mesobuthus species retrieved from GenBank. Names of collection sites are given in Table 1. Localities of two of the outgroup specimens used in the study are also shown in $(\mathrm{A} ; \bullet)$.
Table 1. Mesobuthus eupeus specimens and outgroups used in the present study.

| Map code* | Specimen $\dagger$ | Species | Museum code | Geographic location | Accession numbers of COI sequences $\dagger$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | Me MZ1099 | M. eupeus | ZUTC-arach-1099 | Kiasar - Mazandaran province | HM567346 |
| 2 | Me MZ1083 | M. eupeus | ZUTC-arach-1083 | Kiasar - Mazandaran province | HM567348 |
| 3 | Me MZ1085 | M. eupeus | ZUTC-arach-1085 | Kiasar - Mazandaran province | HM567349 |
| 4 | Me MZ1086 | M. eupeus | ZUTC-arach-1086 | Kiasar - Mazandaran province | HM567350 |
| 5 | Me MZ1092 | M. eupeus | ZUTC-arach-1092 | Kiasar - Mazandaran province | HM567347 |
| 6 | Me RK1056 | M. eupeus | ZUTC-arach-1056 | Neishabour - Pivezhan | HM567364 |
| 7 | Me RK1057 | M. eupeus | ZUTC-arach-1057 | Neishabour - Pivezhan | HM567366 |
| 8 | Me RK1011 | M. eupeus | ZUTC-arach-1011 | Mashhad - Moghan village | HM567362 |
| 9 | Me RK1015 | M. eupeus | ZUTC-arach-1015 | Mashhad - Moghan village | HM567361 |
| 10 | Me RK1016 | M. eupeus | ZUTC-arach-1016 | Mashhad - Moghan village | HM567363 |
| 11 | Me RK1055 | M. eupeus | ZUTC-arach-1055 | Neishabour - Pivezhan | HM567367 |
| 12 | Me NI1112 | M. eupeus | ZUTC-arach-1112 | Esphehan province - Niasar | HM567380 |
| 13 | Me RK1040 | M. eupeus | ZUTC-arach-1040 | ca. 10 km south of Mashhad | HM567358 |
| 14 | Me RK1051 | M. eupeus | ZUTC-arach-1051 | ca. 10 km south of Mashhad | HM567358 |
| 15 | Me RK1053 | M. eupeus | ZUTC-arach-1053 | ca. 10 km south of Mashhad | HM567359 |
| 16 | Me RK1054 | M. eupeus | ZUTC-arach-1054 | ca. 10 km south of Mashhad | HM567360 |
| 17 | Me RK1008 | M. eupeus | ZUTC-arach-1008 | 5 km south of Mashhad (northeast Iran) | HM567357 |
| 18 | Me RK1032 | M. eupeus | ZUTC-arach-1032 | Sarakhs road - Robate-Sharaf | HM567375 |
| 19 | Me RK1039 | M. eupeus | ZUTC-arach-1039 | Sarakhs road - Robate-Sharaf | HM567377 |
| 20 | Me RK1033 | M. eupeus | ZUTC-arach-1033 | Sarakhs road - Robate-Sharaf | HM567372 |
| 21 | Me RK1035 | M. eupeus | ZUTC-arach-1035 | Sarakhs road - Robate-Sharaf | HM567374 |
| 22 | Me FA1123 | M. eupeus | ZUTC-arach-1123 | Kenar tapeh - Kazeroon road | HM567338 |
| 23 | Me FA1124 | M. eupeus | ZUTC-arach-1124 | Kenar tapeh - Kazeroon road | HM567339 |
| 24 | Me SB1132 | M. eupeus | ZUTC-arach-1132 | Baluchistan - Bampur | HM567381 |
| 25 | Me SB1133 | M. eupeus | ZUTC-arach-1133 | Baluchistan - Bampur | HM567368 |

Table 1. (Continued).

| Map code* | Specimen $\dagger$ | Species | Museum code | Geographic location | Accession numbers of COI sequences $\dagger$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 26 | Me SB1134 | M. eupeus | ZUTC-arach-1134 | Baluchistan - Bampur | HM567382 |
| 27 | Me SB1135 | M. eupeus | ZUTC-arach-1135 | Baluchistan - south of Zahedan | HM567369 |
| 28 | Me MZ1003 | M. eupeus | ZUTC-arach-1003 | Southwest Chalus - Kordichal | HM567352 |
| 29 | Me MZ1004 | M. eupeus | ZUTC-arach-1004 | Southwest Chalus - Kordichal | HM567353 |
| 30 | Me MZ1005 | M. eupeus | ZUTC-arach-1005 | Southwest Chalus - Kordichal | HM567354 |
| 31 | Me MZ1006 | M. eupeus | ZUTC-arach-1006 | Southwest Chalus - Kordichal | HM567355 |
| 32 | Me MZ1007 | M. eupeus | ZUTC-arach-1007 | Southwest Chalus - Kordichal | HM567356 |
| 33 | Me YZ1138 | M. eupeus | ZUTC-arach-1138 | Yazd-Mehriz-Marvast road | HM567370 |
| 34 | Me SE1130 | M. eupeus | ZUTC-arach-1130 | Semnan - Biarjomand | HM567373 |
| 35 | Me SE1131 | M. eupeus | ZUTC-arach-1131 | Semnan - Mohammad Abad | HM567351 |
| 36 | Me AZ1118 | M. eupeus | ZUTC-arach-1118 | Bazargan - Gerik road | HM567335 |
| 37 | Me AZ1119 | M. eupeus | ZUTC-arach-1119 | Bazargan, Oskanlu - Kalibar road | HM567337 |
| 38 | Me AZ1120 | M. eupeus | ZUTC-arach-1120 | Buralan - Poldasht road | HM567336 |
| 39 | Me RK1001 | M. eupeus | ZUTC-arach-1001 | Khorasan: Gonabad - 10 km northeast Kakhk | HM567371 |
| 40 | Me RK1145 | M. eupeus | ZUTC-arach-1145 | Khorasan: Gonabad - 10 km northeast Kakhk | HM567390 |
| 41 | Me RK1146 | M. eupeus | ZUTC-arach-1146 | Khorasan: Gonabad - 6 km south Kakhk | HM567391 |
| 42 | Me RK1147 | M. eupeus | ZUTC-arach-1147 | Khorasan: Gonabad - 6 km south Kakhk | HM567392 |
| 43 | Me RK1148 | M. eupeus | ZUTC-arach-1148 | Khorasan: Gonabad - 6 km south Kakhk | HM567393 |
| 44 | Me KR1158 | M. eupeus | ZUTC-arach-1158 | Jiroft - Khatoon Abad village. (southern Iran) | HM567383 |
| 45 | Me SK1106 | M. eupeus | ZUTC-arach-1106 | Khorasan: Nehbandan - Mighan village | HM567378 |
| 46 | Me SK1107 | M. eupeus | ZUTC-arach-1107 | Khorasan: Nehbandan - Mighan village | HM567379 |
| 47 | Me KH1160 | M. eupeus | ZUTC-arach-1160 | Khuzestan province - Bagh Malek | HM567385 |
| 48 | Me KH1161 | M. eupeus | ZUTC-arach-1161 | Khuzestan province - Bagh Malek | HM567386 |
| 49 | Me KH1162 | M. eupeus | ZUTC-arach-1162 | Khuzestan province - Bagh Malek | HM567387 |
| 50 | Me KH1163 | M. eupeus | ZUTC-arach-1163 | Khuzestan province - Bagh Malek | HM567388 |

HM567389 HM567340 HM567341 HM567343 HM567345 HM567344 HM567342 HM567384 n AJ783586
AJ783595 －
 ＜合

 AJ783593合 O AJ550711 $\stackrel{\rightharpoonup}{4}$ oi Lossfy



Khuzestan province－Bagh Malek Fars province－Kazeroon－Borazjan road Fars province－Firouz Abad Fars province－Ghir－o－Karzin road Fars province－Ghir－o－Karzin road Fars province－Ghir－o－Karzin road Fars province－Ghir－o－Karzin road Neishour，Aliab Turkmenistan：Badkhyz Nature Reserve Turkmenistan：Badkhyz Nature Reserve Turkmenistan：Badkhyz Nature Reserve Turkmenistan：northwest Chemenibit Turkmenistan：northwest Chemenibit Turkmenistan：northwest Chemenibit Turkmenistan：Repetek，Karakum Turkmenistan：Repetek，Karakum Turkmenistan：west Kazarma
Turkmenistan：Kushka River valley Turkmenistan：Kushka River valley Turkey：Guelsehir Turkey：Cemilkoey Turkey：Cemilkoey
Greece：Igoumenitsa Greece：Igoumenitsa
Greece：Litochoro Greece：Litochoro
Greece：Mathia Greece：Kalampaka Greece：Petalia Turkey：Hacibectas


[^1]ZUTC－arach－1164 ZUTC－arach－1164
ZUTC－arach－843
ZUTC－arach－851 ZUTC－arach－8632 ZUTC－arach－8632
ZUTC－arach－8693 ZUTC－arach－8691 ZUTC－arach－1159
 NA そて Z で Z でて Z で を そ ぞで Z NA を


[^2]
Table 1. (Continued).

| Map code* | Specimen $\dagger$ | Species | Museum code |  | Accession <br> numbers of COI <br> sequences $\dagger$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| 82 | MgCAa | M. gibbosus | NA | Turkey: Avanos | AJ783459 |
| 83 | Mcl136 | M. caucasicus | ZUTC-arach-1136 | Iran: Sistan and Baluchistan- Bampur | HM567334 |
| 84 | McKZb1 | M. caucasicus | NA | Kazakhstan: northwest of Baigakum | AJ550693 |
| 85 | Mc KZc1 | M. caucasicus | NA | Kazakhstan: Kyzylkum Desert | AJ783602 |
| 86 | Mc TUj2 | M. caucasicus | NA | Turkmenistan: Badkhyz Nature Reserve | AJ783614 |
| 87 | Mc TUo1 | M. caucasicus | NA | Turkmenistan: Repetek Nature Reserve | AJ783515 |
| 88 | Mc TUi2 | M. caucasicus | NA | Turkmenistan: Chagaly, Akhal Region | AJ783509 |
| 89 | Mcy1 | M. cyprius | NA | Cyprus: Roudia bridge | AJ550698 |
| 90 | Mcy2 | M. cyprius | NA | Cyprus: Kantara | AJ550699 |
| 91 | Mcy3 | M. cyprius | NA | Cyprus: Tepebasi | DQ310849 |
| 92 | Ac1110 | Androctonus | ZUTC-arach-1110 | Iran: Nehbandan | HM567333 |
|  |  | crassicauda |  |  |  |

[^3]ATATAAACTTC-3' (Simon et al. 1994). Each PCR contained $2.5 \mu 110 \times$ PCR buffer ( 100 mm Tris- $\mathrm{HCl} \mathrm{pH} 8.3,1.5 \mathrm{~mm} \mathrm{MgCl} 2,500 \mathrm{~mm} \mathrm{KCl}$ ), $2 \mu \mathrm{dNTP}$ mix ( 2.5 mm of each dNTP); $0.6 \mu 1$ ( $=6 \mathrm{pmol}$ ) of each primer, $0.125 \mu \mathrm{~T}$ Taq DNA polymerase (5 units/ $\mu$ l, Takara Bio Inc., Shiga, Japan), and $1-5 \mu 1$ DNA template. The PCR thermal regimen consisted of one cycle of 2.5 min at $94^{\circ} \mathrm{C}$; five cycles of 45 seconds at $94^{\circ} \mathrm{C}, 45$ seconds at $45^{\circ} \mathrm{C}$, and 1 min at $72^{\circ} \mathrm{C}$; 35 cycles of 45 seconds at $94^{\circ} \mathrm{C}$, 1 min at $51^{\circ} \mathrm{C}$ and 1 min at $72^{\circ} \mathrm{C}$; and a final cycle of 10 min at $72^{\circ} \mathrm{C}$. Sequencing was performed using ABI Big Dye terminator chemistry and an ABI Prism 3700 instrument (Applied Biosystems, Foster City, CA, USA). The sequences were analysed using Sequencher version 4.1.1 (GeneCodes Corporation, Ann Arbor, MI, USA). Sequence qualities were visually checked and the infrequent sequences containing ambiguous callings were not used.

Sequence alignment was performed using ClustalX (version 1.8, Thompson et al. 1997) under its default parameters. Validity of callings for variant nucleotides at highly conserved positions was confirmed visually. All novel COI sequences were deposited in GenBank [http://www.ncbi.nlm.nih.gov with the accession numbers HM567333 to HM567393].

## Estimation of intraspecific and interspecific sequence divergence

DNASP 5 (Rozas et al. 2003) and Arlequin 2.00 (Schneider et al. 2000; Excoffier et al. 2005) software was used to compare sequence characterizations, nucleotide diversities and levels of divergence between M. eupeus, M. caucasicus, M. gibbosus and M. cyprius species. Intraspecific sequence divergences among M. eupeus sequences were also calculated using MEGA4 (Tamura et al. 2007).

## Phylogenetic analyses

Phylogenetic relationships were analysed by maximum parsimony, maximum likelihood and Bayesian inference using Paup (Swofford 1998) and MrBayes v.3.1 (Ronquist and Huelsenbeck 2003) software. Parsimony analyses with PAUP (version 4.0) were performed using a heuristic search with equal weighting of all characters, 1000 random stepwise addition and tree bisection-reconnection, (TBR) branch swapping. The robustness of inferred relationships and nodal supports was checked with 1000 bootstrap replications.

For maximum likelihood analyses, the best substitution model was selected with modeltest (version 3.7, Posada and Crandall 1998). Parameters from this best model were used for subsequent likelihood analyses in PAUP with 100 random stepwise additions and TBR branch swapping. Finally, to assess the nodal support of the resulting clades, 250 bootstrap pseudoreplicates were performed.

Bayesian analysis, was carried out with MrBAYES (version 3.2, Ronquist and Huelsenbeck 2003) using the same substitution model used for maximum likelihood analysis, this method was recently proposed as a relatively faster method for estimating phylogenetic trees (Holder and Lewis 2003). The Monte Carlo-Markov chain length was $2,500,000$ generations and trees were sampled every 100 generations. Bayesian topology and posterior probabilities were computed by majority rule consensus after burning of all pre-asymptotic tree scores.

The Bayesian approach was also used for analysis of alignment partitioned by codon positions. The best substitution models for this purpose were selected using PhyML v.2.4.4 (Guindon and Gascuel 2003). The molecular clock hypothesis (i.e. equal rates across all branches) was tested with a $\chi^{2}$ approximated likelihood-ratio test using PaUp (Felsenstein 1981; Huelsenbeck and Crandall 1997; Huelsenbeck and Rannala 1997; Swofford et al. 1996). The trees were compared using two degrees of freedom for operational taxonomic units (OTUs -2 ) i.e. $92-2=90$.

## Morphological comparisons

To test whether genetic divergence is in line with morphological divergence of M. eupeus clades, a comparison of general morphology was performed. A total of 17 morphometric ratios were determined for representatives of each clade. Measurements were taken with a $>0.02-\mathrm{mm}$ accurate graticule on an Olympus BHZ stereomicroscope following Stahnke (1970) and Lamoral (1979). Abbreviations of morphometric ratios used are as follows: $\mathrm{Ca} \_1 / \mathrm{aw}$, carapace length to anterior width; $\mathrm{Ca} \_1 / \mathrm{pw}$, carapace length to posterior width; $\mathrm{Ca} \_$aw/pw, carapace anterior width to posterior width; ca_x/y, the distance between anterior margin of carapace and anterior edge of median eyes to the distance between anterior edge of median eyes and posterior margin of carapace; Mt-I_1/w, metasomal segment I length to width; Mt-I_1/h, metasomal segment I length to height; Mt-II_1/w, metasomal segment II length to width; Mt-II_1/h, metasomal segment II length to height; Mt-III_1/w, metasomal segment III length to width; Mt-III_1/h, metasomal segment III length to height; Mt-IV_1/w, metasomal segment IV length to width; Mt-IV_1/h, metasomal segment IV length to height; MtV_l/w, metasomal segment V length to width; Mt-V_1/h, metasomal segment V length to height; $\mathrm{CH} \_1 / \mathrm{ml}$, chela length to manus length; $\mathrm{Tl} \_1 / \mathrm{w}$, telson length to width; Tl_L/h, telson length to height.

Principal component and canonical discriminant analyses were applied to M. eupeus and M. caucasicus specimens collected from different localities of Iran using SPSS ver. 13 and PAST 1.91 (Hammer et al. 2001). Principal component analysis was performed to determine whether any of the geographic populations are morphologically distinct. Also, to show discrimination between populations, canonical discriminant analysis using Mahalanobis distance was performed.

## Results

## Sequence characteristics and levels of variations

In total, 59 individuals of $M$. eupeus, one specimen of $M$. caucasicus and one specimen of $A$. crassicauda were sequenced. The COI sequences belonging to M. gibbosus (nine sequences), M. cyprius (three sequences) and Central Asian specimens of M. caucasicus (five sequences) and M. eupeus ( 14 sequences) were retrieved from GenBank. The average fragment length of sequences we sequenced and those obtained from GenBank were, respectively, 650 bp and 496 bp . Considering all sequences, frequencies of A, G, C and T were, respectively, $19.7 \%, 26.7 \%, 12.6 \%$ and $41.2 \%$.

Estimates of $\%$ variation among sequences were based on a fragment length of 672 nucleotides, which is the longest length used in the alignments. In the alignment of all sequences, including all Mesobuthus and outgroup species, 395 (58.3\%) positions
were completely conserved and 277 ( $41.7 \%$ ) positions were variable. One hundred and eighty-three ( $27.8 \%$ ) of the positions were parsimony informative. Overall nucleotide diversity $\left(\mathrm{P}_{i}\right)$ for the 92 sequences was 0.076 . Among the 59 M. eupeus specimens, in the $672-b p$ fragment, 196 (29.16\%) variable sites and 476 ( $70.83 \%$ ) completely conserved sites were observed. Of the variable sites, $158 \mathrm{bp}(23.51 \%)$ were parsimony informative. $\mathrm{P}_{i}$ for the 59 organisms was 0.0617 . To assess interspecific divergence, genetic distances between M. eupeus and three other Mesobuthus species were calculated using net between group average using MEGA4. The average distances between M. eupeus and M. caucasicus, M. cyprius and M. gibbosus were, respectively, $7.2 \%$, $10 \%$ and $8.4 \%$. Uncorrected pairwise Kimura two-parameter (K2P) distance distances for all 92 sequences were also calculated using the maximum composition likelihood method. Values for intraspecific distance within the genus Mesobuthus ranged from $0 \%$ to $15 \%$ and values for interspecific distances ranged from $7 \%$ to $20 \%$.

## Best-fit nucleotide substitution model

The best-fit model of sequence evolution selected for COI determined by MODELTEST 3.7 under Akaike information criterion was GTR $+\Gamma+\mathrm{I}(\ln \mathrm{L}=5901.9956$; Akaike information criterion 11,823.991). Estimates for the model parameters employed in maximum-likelihood searches included estimated base frequencies $\left(\pi_{\mathrm{A}}=0.1970, \pi_{\mathrm{C}}=0.1262, \pi_{\mathrm{G}}=0.2656, \pi_{\mathrm{T}}=0.4113\right)$, rate parameter estimates $([\mathrm{A}<>\mathrm{C}]=0.1933 ;[\mathrm{A}<>\mathrm{G}]=13.5717 ;[\mathrm{A}<>\mathrm{T}]=1.4370 ;[\mathrm{C}<>\mathrm{G}]=0.5092$; $[\mathrm{C}<>\mathrm{T}]=3.1889 ;[\mathrm{G}<>\mathrm{T}]=1.000$ ), gamma distribution shape parameter $(\alpha=1.2841)$ and proportion of invariant sites (pinvar $=0.5145$ ). This model was subsequently used for Bayesian phylogenetic inference of $C O I$ sequences. For partitioned sequence data by codon positions, GTR $+\Gamma$, F81 and HKY $+\Gamma$ substitution models were used for first, second and third codon positions, respectively.

The molecular clock hypothesis was tested with the $\chi^{2}$ approximated likelihoodratio test with $\mathrm{df}=92-2=90$. This test makes use of parameters described above. The estimated $p$-value ( $2 \delta=2\left(\ln \mathrm{~L}_{0}-\ln \mathrm{L}_{1}\right) ; \ln \mathrm{L}_{0}=6008.82, \ln \mathrm{~L}_{1}=5901.99 ; \mathrm{df}=90$, $p=0.1089$ ) supported the molecular clock hypothesis.

## Phylogenetic analyses

Maximum likelihood, parsimony and Bayesian analyses all produced phylogenetic trees that were almost congruent. All predicted the existence of an Eastern and a Western clade, subclades A and B within the M. eupeus clade, and further divisions of subclades A and B as described below. The topology resulting from maximum likelihood analysis is presented in Figure 2. Division into two major Mesobuthus clades labelled Eastern and Western is evident. The Western clade is constituted by M. gibbosus and M. cyprius, whereas the Eastern clade is constituted by M. eupeus and M. caucasicus. The M. eupeus clade itself is divided into subclades A and B , and subclades $A$ and $B$ are each further subdivided, respectively, into clades $A_{1}, A_{2}$ and $B_{1}, B_{2}$. Further subdivision of $B_{2}$ itself into $B_{2.1}$ and $B_{2.2}$ may be warranted. A majority rule consensus tree resulted from parsimony analysis (length 1092, Consistency Index (CI): 0.3425 ; Retention Index (RI): 0.7985; Rescaled Consistency Index (RC): 0.2735 and 183 parsimony informative characters), and was similar in topology to the presented maximum likelihood tree (maximum parsimony tree not shown). In the parsimony


Figure 2. Fifty per cent majority rule consensus tree resulting from maximum likelihood analysis. Numbers below branches indicate the posterior probabilities of the nodes in the Bayesian inference analysis. Numbers above branches are the bootstrap values of the nodes in the maximum likelihood analysis. The estimated separation time for calibration point ( $\pm$ standard deviations) is also displayed. The geographic distributions of inferred clades are shown on the embedded map.
consensus tree, there were no nodes that contradicted those presented in the maximum likelihood tree. Topologies resulting from Bayesian analysis using complete and partitioned sequence data based on codon positions of nucleotides were also similar to topologies described above.

Table 2. K2P genetic distances between Mesobuthus eupeus clades, M. caucasicus, M. cyprius, M. gibbosus and Androctonus crassicauda.

|  | $\mathrm{A}_{1}$ | $\mathrm{~A}_{2}$ | $\mathrm{~B}_{1}$ | $\mathrm{~B}_{2.1}$ | $\mathrm{~B}_{2.2}$ | M.ca | M.cy | M.g |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Clade $\mathrm{A}_{1}$ |  |  |  |  |  |  |  |  |
| ${\text { Clade } \mathrm{A}_{2}}^{\text {Clade } \mathrm{B}_{1}}$ | 0.05 |  |  |  |  |  |  |  |
| ${\text { Clade } \mathrm{B}_{2.1}}^{0.05}$ | 0.06 | 0.06 | 0.05 |  |  |  |  |  |
| Clade $\mathrm{B}_{2.2}$ | 0.06 | 0.07 | 0.05 | 0.04 |  |  |  |  |
| M. caucasicus | 0.07 | 0.07 | 0.07 | 0.08 | 0.07 |  |  |  |
| M. cyprius | 0.09 | 0.10 | 0.10 | 0.11 | 0.10 | 0.08 |  |  |
| M. gibbosus | 0.08 | 0.08 | 0.09 | 0.09 | 0.08 | 0.07 | 0.05 |  |
| Outgroup | 0.13 | 0.13 | 0.13 | 0.14 | 0.13 | 0.13 | 0.13 | 0.11 |

Notes: $\mathrm{A}_{1}, \mathrm{~A}_{2}, \mathrm{~B}_{1}, \mathrm{~B}_{2.1}$ and $\mathrm{B}_{2.2}$ represent Mesobuthus eupeus clades shown in Figure 2. M.ca, Mesobuthus caucasicus; M.cy, Mesobuthus cyprius; M.g, Mesobuthus gibbosus; and Outgroup, Androctonus crassicauda
In the analysis, sequences of all specimens of respective clades shown in Figure 2 were used.

## Genetic distances and age estimation

Genetic distances between $M$. eupeus clades $\mathrm{A}_{1}, \mathrm{~A}_{2}, \mathrm{~B}_{1}, \mathrm{~B}_{2.1}$ and $\mathrm{B}_{2.2}$, three Mesobuthus species, M. caucasicus, M. gibbosus and M. cyprius and the outgroup species $A$. crassicauda were calculated (Table 2). K2P and net between group averages were used in the calculations. The genetic distances between the M. eupeus clades ranged between $4 \%$ and $7 \%$, and the maximum distance was observed between clades $\mathrm{A}_{2}$ and $\mathrm{B}_{2.2}$. The distances between M. eupeus and other Mesobuthus species ranged from $7 \%$ to $11 \%$. Distances between the other Mesobuthus species were $5 \%$ to $8 \%$. Finally, as expected, the distances between Mesobuthus species and the outgroup species $A$. crassicauda were in the range of $11 \%$ to $14 \%$.

Inspection of the geographic distribution of clades shown in Figure 2 showed that those of clades A and B, respectively, clustered in the southern and northern regions of Iran (Figure 2). The Zagros Mountains separating these two regions may well serve as a geographic barrier. To test the role of Zagros formation in the separation of these clades, the age of calibration point that represents the node at which clades A and B were separated, was estimated (Figure 2). The mean divergence between the clades A and B was $6.00 \pm 0.8 \%$ (mean $\pm \mathrm{SE}$ ). For estimation of divergence time, calculations were performed assuming divergence rates of $1.3 \%$ and $1.9 \%$ per million years. The published divergence rates of $C O I$ for Arthropoda ranged between these figures. The estimated divergence times between clades A and B based on the $1.3 \%$ and $1.9 \%$ values were, respectively $4.61 \pm 0.61$ million years ago (mya) and $3.15 \pm 0.42$ mya.

## Morphological comparisons

Principal component analysis based on 17 morphological ratios showed a deep divergence between M. eupeus and M. caucasicus, as expected (Figure 3). The first three components resulting from the analysis explained $45.15 \%, 16.88 \%$ and $15.64 \%$ of the


Figure 3. Principal component analysis of Mesobuthus eupeus and Mesobuthus caucasicus based on morphological ratios. (A) and (B) plot positions of the species, respectively, with respect to the first and second components and with respect to the first and third components (see text). Mesobuthus eupeus and M. caucasicus are encircled according to species identification in phylogenetic tree shown in Figure 2: ■, M. eupeus; $\bullet$, M. caucasicus.
total variation. Figure $3(\mathrm{~A}, \mathrm{~B})$ clearly show that the first component was most effective in defining the divergence between the two species. Variables Mt5_1/h, Mt5_1/w and $\mathrm{Tl} \_1 / \mathrm{w}$ had the highest positive loadings for this component, and two variables $\mathrm{Ca} \_$aw/pw and $\mathrm{Ca} \_\mathrm{X} / \mathrm{Y}$ had negative loadings.


Figure 4. Principal component analysis of Mesobuthus eupeus based on morphological ratios. The specimens are plotted with respect to first and second components (see text). Mesobuthus eupeus specimens are encircled within clades according to phylogenetic tree shown in Figure 2:
$\bullet$, M. eupeus clade $\mathrm{A}_{1} ; \mathbf{X}$, M. eupeus clade $\mathrm{A}_{2} ; \boldsymbol{*}$, M. eupeus clade $\mathrm{B}_{1} ; \mathbf{\square}$, M. eupeus clade $\mathrm{B}_{2}$.

Principal component analysis based on the same 17 morphological ratios was also performed only for M. eupeus species (Figure 4). Notably, the analysis showed divergence between $M$. eupeus clade A and M. eupeus clade B. As expected, the divergence between these M. eupeus clades was notably less than between M. eupeus and M. caucasicus (Figure 3). For the M. eupeus species analysis, the first three components explained $28.83 \%, 26.02 \%$ and $15.09 \%$ of the total variation. Here, the second component was more effective in defining the divergence, and variables Ch_l/ml, Tl_l/w and Tl_1/h had positive loadings. Canonical discriminant analysis based on the 17 morphological ratios resulted in discrimination between M. eupeus clade A, M. eupeus clade B and M. caucasicus (Figure 5).

## Discussion

A remarkable feature of the tree presented in Figure 2 is the poor resolution for certain clades. As stated by Rokas et al. (2002), the first reason for poor resolution is that the trees might represent an adaptive radiation, if the rate of speciation for a given time period is relatively high, interlineage differentiations are expected to be low. This would result in poorly resolved phylogeny, which, however, is an exact representation of the historical relationships between the taxa. Moreover, employing several loci which diverged at different rates and/or addition of more taxa, is probably an alternative way to obtain well-resolved phylogenetic trees. However, Gantenbein and Keightley (2004) used nine nuclear genes ( 3856 bp ) in the study of the genus Mesobuthus in the eastern Mediterranean region, but the phylogenetic tree produced by their data set was also weakly supported. Therefore, despite the weak statistical


Figure 5. Canonical discriminant analysis of Mesobuthus eupeus and Mesobuthus caucasicus species based on morphological ratios. Mesobuthus eupeus and M. caucasicus are grouped according to species identification shown in Figure 2: $\boldsymbol{\square}$, M. eupeus clade A; $\mathbf{A}$, M. eupeus clade B; $\boldsymbol{O}^{\text {, M. caucasicus; } \boldsymbol{*} \text {, group centroid. }}$
support for major clades, the low nodal support of the resulting tree (Figure 2), is an accurate representation of the historical associations between the taxa analysed.

The COI sequence data presented here confirm the monophyly of M. eupeus as sister taxon of M. caucasicus (Figure 2). Furthermore, as expected, M. gibbosus and M. cyprius constituted a "Western clade" ( $89 \%$ bootstrap $)$ well separated from the "Eastern clade" including M. eupeus and M. caucasicus ( $54 \%$ bootstrap). In the topology observed, the phylogenetic relationships within the Western clade were not clearly resolved as M. gibbosus was paraphyletic with respect to M. cyprius, suggesting that they may be synonymous. The latter species was described by Gantenbein et al.(2000) based on allozyme data and morphological evidence from Cyprus. Also, the data are consistent with M. gibbosus consisting of more than one species as proposed by Parmakelis et al. (2006). As the M. gibbosus and M. cyprius sequence data were retrieved from GenBank, we will not make a taxonomic decision on M. cyprius in this study.

With regards to M. eupeus, this taxon is a morphologically complicated species with 14 formally valid subspecies, of which five (Farzanpay 1987) to nine (Fet 1989; Fet and Lowe 2000) subspecies were described from Iran. It was considered that
further phylogenetic analyses may result in elevation of some subspecies to the species level (Gantenbein et al. 2003). The COI data presented here confirmed the monophyly of M. eupeus. Additionally, all the analyses suggest subdivision of the species into distinct clades. Notably, the major subdivisions are supported by morphological comparisons and geographic distributions. They are also potentially explainable according to the geomorphological history of the Iranian Plateau. The most distinct division of $M$. eupeus shown by the resulting topology is between clade A and clade B . The phylogenetic tree is also compatible with further subdivision of clades $A$ and $B$, respectively, into clades $A_{1}$ and $A_{2}$, and clades $B_{1}$ and $B_{2}$. Clade $B_{2}$ is potentially further divisible into $\mathrm{B}_{2.1}$ and $\mathrm{B}_{2.2}$ (Figure 2). The clade $\mathrm{A}_{2}$ which was originally described as a separate species, Buthus phillipsi, was later downgraded to a subspecies of M. eupeus as M. e. phillipsi (Farzanpay 1987; Fet and Lowe 2000). Sequence divergence comparisons support these divisions, as the values for pairwise comparisons between $M$. eupeus clades ( $4-7 \%$ ) approach those observed between distinct Mesobuthus species (5-11\%) (Table 2). The highest divergence (7\%) was observed between members of $\mathrm{A}_{2}$ and $\mathrm{B}_{2.2}$ clades.

Clade A members were all collected from the southern parts of Iran, whereas those of clade B were from a belt extending from southwestern Turkey (nominotypical subspecies M. e. eupeus) to the north (M. e. philippovitschi), northeastern Iran and southeastern Turkmenistan (M. e. thersites). The habitats of clade A are highly to moderately arid and distinctly different from the habitats of clade B. The habitats of clade B include humid environments on northern slopes of the Alborz Mountains, high altitude locations in northeastern and northwestern Iran and xeric habitats in the east of Iran. In general, specimens from the northern and northwestern parts of the range of distribution (Clade B) tend to be darker in colouration than specimens from further southern regions (Clade A). In the resultant topologies the nominotypical subspecies, M. e. eupeus, which is found in Turkey, Caucasus and northwest Iran, is highly supported, but the other northern sequences showed a more complicated case. Therefore, the taxonomic validity of the other northern subspecies, namely M. e. philippovitschi should be revised carefully based on the morphological characteristics and geographic data. Furthermore, the sequences from northeast and east of Iran could be assigning to the subspecies, M. e. thersites and M. e. afghanus. Finally, the relationship between southern sequences (Clade A) is concordant with two subspecies namely M. e. phillipsi (southwest) and M. e. kirmanensis (southeast).

Earlier morphological comparisons of $M$. eupeus had suggested existence of various subspecies (Birula 1900, 1905; Farzanpay 1987; Fet 1994). Reanalysis of morphological characteristics here by principal component and canonical discriminant analyses not only obviously distinguished between M. eupeus and M. caucasicus, but also supported the existence of M. eupeus clade A and M. eupeus clade B (Figures 3 and 4). The members of clade B are distinguished from those of clade A by having longer metasomal segments (I-III), relatively shorter fixed fingers and more inflated vesicles.

Topographic barriers that may have been responsible for distinction of the clades proposed here need to be considered. Our COI-based phylogenetic tree is compatible with clade A being ancestral to clade B (Figure 2). As distribution of clade A was distinctly within southern Iran and that of clade B was in northern Iran, a south to north distributional expansion in past history is implicated. This gradient may have been instigated by progressive expansion of aridization toward the north, making northern
regions of Iran favourable habitats for scorpion species and promoting the migration northward. The proposed palaeoclimatic changes and aridization occurred during the Tertiary (Fet 1994; Gantenbein et al. 2003).

The present-day geographic isolation of clade A and clade B may have been affected by topographic barriers that evolved during or after the distributional expansion process. The relevant vicariant events here may have been the uplifting and formation of the Zagros and Alborz Mountain ranges, which occurred in the late Tertiary approximately 5-10 mya (Macey et al. 1998; Gök et al. 2003; Ramezani Oomali et al. 2008). The formation of Kavir and Lut deserts as a consequence of geomorphological events on the Iranian Plateau may act as barriers to dispersal. The effects of geomorphological features on the distribution of various African species of Hottentotta, Parabuthus, Uroplectes and Opistophthalmus have been previously reported (Lamoral 1979; Fet et al. 1998; Prendini 2005). Mountain systems have played the most important role in shaping the present distribution of various taxa, including the gekkonid lizards of the genus Assacus (Macey, et al. 1998; Rastegar Pouyani 2006). Rastegar Pouyani (2006) proposed that fragmentation resulting from uplifting of Zagros Mountains in the late Miocene or early Pliocene is a general scenario for the fauna of the Iranian Plateau.

Fossil evidences of Mesobuthus are lacking. However, time of separation of clade A and clade B can be estimated based on an assumed molecular clock. The time of divergence was estimated at 3.15-4.61 mya based on published divergence rates of COI (Quek et al. 2004). The 4.61-mya estimate may be more accurate because it is based on the relatively slower sequence divergence rate of $1.3 \%$ per million years. Variations in sequence divergence rates mainly depend on generation time and the rate of metabolism (Towler et al. 2001). Within arthropods, scorpions gave the longest generation time and they had the lowest known rate of metabolism among all animals (Martin and Palumbi 1993), suggesting that their sequence divergence rate should be relatively low. The estimated age of 4.61 mya (and even of 3.15 mya) for clade $A$ and $B$ corresponds to the early and middle Pliocene geological events of the Iranian Plateau and is consistent with the proposal that the formation of Zagros in the south and consequent uplifting of Alborz may have affected their evolution.

In conclusion, the phylogenetic analysis and sequence divergence data presented here based on COI sequence data indicate two distinct lineages within M. eupeus suggesting that it may be a species complex consisting of at least two species. Although, considerable morphological conservation is apparent for M. eupeus, statistical analysis of morphological features are consistent with the possible phylogenetic history being considered. Geographic distribution of M. eupeus and palaeogeographic evidences also support the proposal. However, as the molecular analysis was based only on the partial sequence of a single gene, a taxonomic decision about M. eupeus is not now warranted. We suggest further molecular testing using other genes and a detailed revision of diagnostic and morphological characteristics before a taxonomic decision is made.

## Acknowledgements

The authors thank Ghasem M. Kashani, Afshin Faghiri and Siavash Taravati for their help during field activities. We dedicate this paper to our friends and student colleagues, Ehsan Entezari and Mostafa Tarahomi, who also assisted in sample collection, but have since passed away
because of an unfortunate accident during fieldwork. We are grateful to Shahrokh Navidpour (Razi Reference Laboratory of Scorpions), Peter Jäger (Senckenberg Museum, Frankfurt) and Elise-Ann Leguin (Museum National d'Histoire Naturelle, Paris) for the loan of specimens. We are grateful to Lorenzo Prendini, Frantisek Kovařík, Victor Fet and Mansour Aliabadian for invaluable discussions during the preparation of this manuscript. Financial support was provided by the Office of Research Affairs, University of Tehran, Iran.

## References

Birula AA. 1900. Beiträge zur Kenntniss der Scorpionenfauna Ost-Persiens (1. Beiträg). Bull Acad Imp Sci, St-Pétersb. Series 5,12(4): 355-375.
Birula AA. 1905. Miscellanea scorpiologica.VIII. Bemerkungen ueber die SkorpioneSammlung des Kaukasischen Museum zu Tiflis. Bull Acad Imp Sci, St. Pétersb. 10:119-131.
Birula AA. 1917. Arthrogastric Arachnids of Caucasia. Part I. Scorpions. Zap Kavk Mus, sér. A. 5:1-170 (in Russian).

Excoffier L, Laval G, Schneider S. 2005. Arlequin ver. 3.0: An integrated software package for population genetics data analysis. Evol Bioinform. 1:47-50.
Farzanpay R. 1987. Knowing scorpions. No. 312, Biology 4. Teheran: Central University Publications (in Farsi with Latin index).
Felsenstein J. 1981. Evolutionary trees from DNA sequences: a maximum likelihood approach. J Mol Evol. 17:368-376.
Fet V. 1989. A catalogue of scorpions (Chelicerata: Scorpiones) of the USSR. Riv Mus Sc Nat Bergamo. 13:73-171.
Fet V. 1994. Fauna and zoogeography of scorpions (Arachnida: Scorpiones) in Turkmenistan. In: Fet V, Atamuradov KI, editors. Biogeography and ecology of Turkmenistan (Monographiae Biologicae 72). Dordrecht-Boston: Kluwer Academic Publishers. pp. 525-534.
Fet V, Gantenbein B, Gromov AV, Lowe G, Lourenço WR. 2003. The first molecular phylogeny of Buthidae (Scorpiones). Euscorpius 4:1-12.
Fet V, Hendrixson BE, Sissom WD, Levy G. 2000. First record for the genus Mesobuthus Vachon, 1950 in Israel: Mesobuthus nigrocinctus (Ehrenberg, 1828), comb. n. (Scorpiones: Buthidae) from Mt. Hermon. Israel J Zool. 46:157-169.
Fet V, Lowe G. 2000. Family Buthidae C.L. Koch, 1837. In: Fet V, Sissom WD, Lowe G, Braunwalder ME, editors. Catalog of the scorpions of the world (1758-1998). New York: New York Entomological Society. pp. 54-286.
Fet V, Polis GA, and Sissom WD. 1998. Life in sandy deserts: the scorpion model. J Arid Environ. 39:609-622.
Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R. 1994. DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Mol Mar Biol Biotech. 3:294-299.
Gantenbein B, Büchi L, Braunwalder ME, Scholl A. 1997. The genetic population structure of Euscorpius germanus (Scorpiones: Chactidae). Proceedings of the 17th European Colloquium of Arachnology; 14-18 July 1997; Edinburgh.
Gantenbein B, Fet V, Barker MD. 2001. Mitochondrial DNA markers reveal a deep, divergent phylogeny in Centruroides exilicauda (Wood, 1863) (Scorpiones: Buthidae). In: Fet V, Selden PA, editors. Scorpions 2001: In Memoriam Gary A. Polis. Burnham Beeches, Bucks, UK: British Arachnological Society. pp. 235-253.
Gantenbein B, Fet V, Gantenbein-Ritter IA, Balloux F. 2005. Evidence for recombination in scorpion mitochondrial DNA (Scorpiones: Buthidae). Proc R Soc Lond. B. 272:697-704.
Gantenbein B, Fet V, Gromov AV. 2003. The first DNA phylogeny of four species of Mesobuthus (Scorpiones, Buthidae) from Eurasia. J Arachnol. 31:412-420.

Gantenbein B, Fet V, Largiadèr CR, Scholl A. 1999. First DNA phylogeny of the genus Euscorpius Thorell, 1876 (Scorpiones, Euscorpiidae) and its bearing on the taxonomy and biogeography of this genus. Biogeographica (Paris) 75:59-72.
Gantenbein B, Keightley PD. 2004. Rates of molecular evolution in nuclear genes of east Mediterranean scorpions. Evolution. 58:2486-2497.
Gantenbein B, Kropf C, Largiadèr CR, Scholl A, 2000. Molecular and morphological evidence for the presence of a new buthid taxon (Scorpiones: Buthidae) on the Island of Cyprus. Rev Suisse Zool. 107:213-232.
Gantenbein B, Largiadèr CR. 2002. Mesobuthus gibbosus (Scorpiones: Buthidae) on the island of Rhodes-hybridisation between Ulysses'stowaways and native scorpions? Mol Ecol. 11:925-938.
Gantenbein B, Largiadèr CR. 2003. The phylogeographic importance of the Strait of Gibraltar as a gene flow barrier in terrestrial arthropods: a case study with the scorpion Buthus occitanus as model organism. Mol Phylogenet Evol. 28:119-130.
Gantenbein B, Scholl A. 1998. Allozymes show an unusually high genetic differentiation of Euscorpius germanus (Scorpiones, Chactidae) populations. Rev Suisse Zool. 105(4):748-749.
Gantenbein B, Soleglad ME, Fet V. 2001. Euscorpius balearicus Caporiacco, 1950, stat. nov. (Scorpiones:Euscorpiidae): molecular (allozymes and mtDNA) and morphological evidence for an endemic Balearic Islands species. Org Divers Evol. 1: 301-320.
Gök R, Sandvol E, Turkelli N, Seber D, Barazangi M. 2003. Sn attenuation in the Anatolian and Iranian Plateau and surrounding regions. Geophys Res Lett. 30(24):1-13.
Gromov AV. 2001. On the northern boundary of scorpions (Arachnida: Scorpiones) in Central Asia. In: Fet V, Selden PA, editors. Scorpions 2001: In Memoriam Gary A. Polis. Burnham Beeches, Bucks, UK: British Arachnological Society. pp. 301-307.
Guindon S, Gascuel O. 2003. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst Biol. 52:696-704.
Hammer A, Harper DAT, Ryan PD. 2001. PAST: Paleontological statistics software package for education and data analysis. Paleont Elec. 4(1):1-9.
Holder M, Lewis PO. 2003. Phylogeny estimation: traditional and Bayesian approaches. Nat Rev. 4: 275-284.
Huber D, Gantenbein B, Fet V, Scherabon B. 2001. Euscorpius carpathicus (L., 1767) in Austria (Scorpiones: Euscorpiidae): phylogenetic position clarified by mitochondrial DNA analysis. In: Fet V, Selden PA, editors. Scorpions 2001: In Memoriam Gary A. Polis. Burnham Beeches, Bucks, UK: British Arachnological Society,pp. 273-278.
Huelsenbeck JP, Rannala B. 1997. Phylogenetic methods come of age: testing hypotheses in an evolutionary context. Science 276:227-232.
Huelsenbeck JP, Crandall KA. 1997. Phylogeny estimation and hypothesis testing using maximum likelihood. Annu Rev Ecol Syst. 28:437-466.
Karataş A, Karataş A. 2001. First record of Mesobuthus eupeus (C.L. Koch, 1839) from Central Anatolia (Scorpiones: Buthidae). In: Fet V, Selden PA, editors. Scorpions 2001: In Memoriam Gary A. Polis. Burnham Beeches, Bucks, UK: British Arachnological Society. pp. 297-299.
Karataş A, Karataş A. 2003. Mesobuthus eupeus (C.L. Koch, 1839) (Scorpiones: Buthidae) in Turkey. Euscorpius. 7:1-6.
Kovařík F. 1997. Results of the Czech biological expedition to Iran. Part 2. Arachnida: Scorpiones, with descriptions of Iranobuthus krali gen. n. et sp. N. and Hottentotta zagrosensis sp. N. (Buthidae). Acta Soc Zool Bohem. 61:39-52.
Lamoral BH. 1979. The scorpions of Namibia. Ann Natal Mus. 23(3):497-784.
Lourenço WR. 2000. Reproduction in scorpions, with special reference to parthenogenesis. Proceedings of the 19th European Colloquium of Arachnology; 17-22 July 2000; Århus, Denmark.

Lourenço WR, Qi JX, Zhu MS. 2005. Description of two new species of scorpions from China (Tibet) belonging to the genera Mesobuthus Vachon (Buthidae) and Heterometrus Ehrenberg (Scorpionidae). Zootaxa 985:1-16.
Lowe G, Kutcher SR, Edwards D. 2003. A powerful new light source for ultraviolet detection of scorpions in the field. Euscorpius 8:1-7.
Macey JR, Schulte JA, Ananjeva NB, Larson A, Rastegar-Pouyani N, Shammakov SM, and Papenfuss TJ. 1998. Phylogenetic relationships among agamid lizards of the Laudakia caucasia species group: testing hypotheses of biogeographic fragmentation and an area cladogram for the Iranian Plateau. Mol Phylogenet Evol. 10(1):118-131.
Martin AP, Palumbi SR. 1993. Body size, metabolic rate, generation time, and the molecular clock. Proc Natl Acad Sci USA. 90:4087-4091.
Parmakelis A, Stathi I, Chatzaki MS, Simaiakis S, Spanos L, Louis C, Mylonas M. 2006. Evolution of Mesobuthus gibbosus (Brullé, 1832) (Scorpiones: Buthidae) in the northeastern Mediterranean region. Mol Ecol. 15:2883-2894.
Pocock RI. 1889. Arachnida, Chilopoda and Crustacea. In: On the Zoology of the Afghan commission delimitation. Trans Linnean Soc Lond Zool sér. 2. 5(3):110-122.
Polis GA, Sissom WD. 1990. Life history. In: Polis GA, editor. Biology of scorpions. Stanford, CA: Stanford University Press161-223.
Posada D, Crandall KA. 1998. MODELTEST: testing the model of DNA substitution. Bioinformatics 14:817-818.
Prendini L. 2001. Substratum specialization and speciation in southern African scorpions: the Effect Hypothesis revisited. In: Fet V, Selden PA, editors. Scorpions 2001: In Memoriam Gary A. Polis. Burnham Beeches, Bucks, UK: British Arachnological Society. pp. 113-138.
Prendini L. 2005. Scorpion diversity and distribution in southern Africa: pattern and process. In: Huber BA, Sinclair BJ, Lampe KH. editors. African biodiversity: molecules, organisms, ecosystems. Proceedings of the 5th International Symposium on Tropical Biology, Museum Alexander Koenig, Bonn. New York: Springer Verlag. pp. 25-68
Quek SP, Davies SJ, Itino T, Pierce NE. 2004. Codiversification in an ant-plant mutualism: stem texture and the evolution of host use in Crematogaster (Formicidae: Myrmicinae) inhabitants of Macaranga (Euphorbiaceae). Evolution 58:554-570.
Ramezani Oomali R, Shahriari S, Hafezi Moghaddas N, Omidi P, Eftekharnejahd J. 2008. A model for active tectonics in Kopet Dagh (North-East Iran). World Appl Sci J. 3(2):312-316.
Rastegar-Pouyani N. 2006. Systematics of the genus Assacus (Sauria: Gekkonidae) on the Zagros Mountains, Iran. Paper presented at: Herpetologia Bonnensis II. Proceedings of the 13th Congress of the Societas Europaea Herpetologica; October 2005; Germany.
Rokas A, Nylander JAA, Ronquist F, Stone GN.2002. A maximum likelihood analysis of eight phylogenetic markers in gallwasps (Hymenoptera: Cynipidae): Implications for insect phylogenetic studies. Mol Phylogenet Evol. 22:206-219.
Ronquist F, Huelsenbeck JP. 2003. Mrbayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19:1572-1574.
Rozas J, Sanchez-DelBarrio JC, Messeguer X, Rozas R. 2003. DnaSP, DNA polymorphism analyses by the coalescent and other methods. Bioinformatics 19:2496-2497.
Salomone N, Vignoli V, Frati F, Bernini F. 2007. Species boundaries and phylogeography of the "Euscorpius carpathicus complex" (Scorpiones: Euscorpiidae) in Italy. Mol Phylogenet Evol. 43:502-514.
Schneider S, Roessli D, Excoffier L. 2000. Arlequin, Version 2000: a Software for Population Genetics Data Analysis Genetics and Biometry Laboratory. Geneva: University of Geneva.
Shi CM, Huang ZS, Wang L, He LJ, Hua YP, Leng L, Zhang DX. 2007. Geographical distribution of two species of Mesobuthus (Scorpiones, Buthidae) in China: insights from systematic field surveys and predictive models. J Arachnol. 35:215-226.

Simon C, Frati F, Beckenbach A, Crespi B, Liu H, Flook P. 1994. Evolution, weighting, and phylogenetic utility of mitochondrial gene sequences and a compilation of conserved polymerase chain reaction primers. Ann Entomol Soc Am. 87:651-701.
Sissom WD. 1990. Systematics, biogeography and paleontology. In: Polis GA. editor. Biology of scorpions. Stanford, CA: Stanford University Press. pp. 64-160.
Stahnke HL. 1970. Scorpion nomenclature and mensuration. Entomol News. 81:297-316.
Swofford DL, Olsen GJ, Waddell PJ, Hillis DM. 1996. Phylogenetic inference. In: Hillis DM, Moritz C, Mable BK, editors. Molecular systematics. Sunderland (MA): Sinauer Associates. pp. 407-510.
Swofford DL. 1998. PAUP* Phylogenetic analysis using parsimony (*and other methods). Version 4. Sunderland (MA): Sinauer Associates.
Tamura K, Dudley J, Nei M, Kumar S. 2007. MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0. Mol Biol Evol. 24:1596-1599.
Teruel R. 2000. First record of Mesobuthus eupeus (Koch, 1839) from western Turkey (Scorpiones:Buthidae). Rev Iberica Aracnol. 5(31):75-76.
Thompson JD, Gibson TJ, Plewniak F, Jeanmougin F, Higgins DG. 1997. The ClustalX windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Res. 24:4876-4882.
Towler WI, Ponce Saavedra J, Gantenbein B, Fet V. 2001. Mitochondrial DNA reveals a divergent phylogeny in tropical Centruroides (Scorpiones: Buthidae) from Mexico. Biogeographica 77(4):157-122.
Vachon M. 1950. Études sur les Scorpions. III(suite). Déscription des Scorpions du Nord de l'Afrique. Arch Inst Pasteur Alger. 28:152-216.
Vachon M. 1952. Etudes sur les scorpions. Alger: Institut Pasteur d'Algérie. 482 pp.
Vachon M, Kinzelbach R. 1987. On the taxonomy and distribution of the scorpions of the Middle East. Proceedings of the Symposium on the Fauna and Zoogeography of the Middle East; 1985; Mainz.
Vignoli V, Salomone N, Caruso T, Bernini F. 2005. The Euscorpius tergestinus (C.L. Koch, 1837) complex in Italy: Biometrics of sympatric hidden species (Scorpiones: Euscorpiidae). Zool Anz. 244:97-113.


[^0]:    *Corresponding author. Email: mirshams@ferdowsi.um.ac.ir

[^1]:    Turkey：Avanos

[^2]:    Me KH1164
    $\infty$
    $\sum_{i}^{\infty}$
    $\sum_{i}^{\infty}$
    $\sum_{i}^{\infty}$
    $i$
    N
    M
    힝
    $\stackrel{N}{0}$ n Me RK1180 Me TUil Me TUr1 im＠L等完 $\stackrel{\pi}{5}$ N S N N究 N $\sum^{0}$ 뭉 $\sum_{0}^{\circ}$ त्र Nㅣㄴ $\sum_{2}^{6}$ $\sum_{i}^{\circ} \sum_{i}^{0}$这 $\sum_{0}^{0} \underbrace{0}_{0}$

[^3]:    Notes: COI, cytochrome C oxidase, subunit I; NA, not available; ZUTC, Zoological Museum, University of Tehran.
    *Map codes correspond to locations shown in Figure 1.
    $\dagger$ All non-Mesobuthus eupeus (Me) specimens served as outgroups.
    $\ddagger$ Obtained from: http://www.ncbi.nlm.nih.gov/

