Mitochondrial DNA and karyotypic data confirm the presence of *Mus indutus* and *Mus minutoides* (Mammalia, Rodentia, Muridae, *Nannomys*) in Botswana

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Abstract

We use a combination of cytochrome *b* sequence data and karyological evidence to confirm the presence of *Mus indutus* and *Mus minutoides* in Botswana. Our data include sampling from five localities from across the country, including one site in northwestern Botswana where both species were captured in syntopy. Additionally, we find evidence for two mitochondrial lineages of *M. minutoides* in northwestern Botswana that differ by 5% in sequence variation. Also, we report that *M. minutoides* in Botswana have the 2n=34 karyotype with the presence of a (X.1) sex-autosome translocation.

Keywords

Africa, rodent, distribution, karyotype, sex-autosome translocation, cytochrome *b*

Introduction

Delineating geographic distributions of African *Mus* (subgenus *Nannomys* Peters, 1876) in Sub-Saharan Africa has been especially challenging due to a combination of incomplete taxon sampling throughout the region as well as uncertainties in species
identification resulting from their highly conserved morphology. Despite morphological similarities, African pygmy mice (*Nannomys*) are characterized by a high degree of chromosomal variation, including chromosomal rearrangements such as Robertsonian translocations, pericentric inversions, heterochromatin additions, and tandem fusions (see summary in Britton-Davidian et al. 2012). Additionally, whole-arm translocations (WARTs) and novel sex-chromosome determination have been documented in populations in South Africa (Veyrunes et al. 2007, 2013).

Britton-Davidian et al. (2012) produced the most complete phylogenetic analysis of *Nannomys* to date, which included previously published sequences of nine species (Suzuki et al. 2004; Chevret et al. 2005; Veyrunes et al. 2005; Kan Kouassi et al. 2008; Veyrunes et al. 2010; Mboumba et al. 2011; and others). Their phylogeny illuminated the diversity of taxa within this subgenus (including at least one unnamed species from Chad), and clearly indicated that further phylogenetic investigations are necessary to clarify species diversity within *Nannomys*. Their comprehensive review surmised that there are at least 18 species of African pygmy mice and estimated that eight species occur within southern Africa (Britton-Davidian et al. 2012). In addition, their study highlighted important gaps in both geographic and taxonomic sampling for this subgenus, particularly within southern Africa. Included in this underrepresented southern African group is *Mus minutoides*, one of the most widespread pygmy mice, with a distribution encompassing most of Sub-Saharan Africa.

Within the southern African country of Botswana, the taxonomy of *Mus* has never fully been resolved. Early assessments of the regional mammalian fauna (Smithers 1971) concluded that two native forms of *Mus* exist within Botswana: the widespread *Mus minutoides indutus* (Thomas, 1910)—which was later elevated to specific status (Petter and Matthey 1975; Musser and Carleton 1993, 2005)—and an arid-adapted form with large ears and a white band of fur near the rump (referred to as *Leggada* sp. in Smithers 1971) restricted to northwestern Botswana and a single record from Sekhuma Pan in the Jwaneng District of southern Botswana (Petter 1978). This latter species was later described as *M. setzeri* Petter, 1978. De Graaff’s (1981) assessment of *Nannomys* in southern Africa concluded that all records for Botswana conform to *M. minutoides*, although he acknowledged that *M. m. indutus* and the desert form (*M. setzeri*) may be distinct species that require further study. More recent evaluations describe allopatric distributions for *M. indutus* and *M. minutoides* and only acknowledge the former within the boundaries of Botswana (Skinner and Chimimba 2005, Hapbold and Veyrunes 2013). These recent assessments estimated the geographic range for *M. minutoides* as extending from the southwest cape in South Africa through the Zambezian woodlands in the east (Fig. 1a, dark grey). Monadjem (2013a) stated that *M. indutus* replaces *M. minutoides* in the western part of the Zambezian woodlands and extends throughout Botswana and into neighboring countries (Fig. 1a, light grey). Although Britton-Davidian et al. (2012) proposed that the range of *M. minutoides* greatly differs from the map published by Monadjem (2008b), and including the countries of Angola, Botswana, Namibia, Zambia, and Zimbabwe, verified records from their study were only presented for South Africa, Swaziland, and Zimbabwe. However, records
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of *M. minutoides* have recently been confirmed for Angola and Namibia (Lamb et al. 2014), providing additional support for the extended range map proposed by Britton-Davidian et al. (2012)

Regarding chromosomal rearrangement in southern Africa, *M. minutoides* from South Africa exhibit Robertsonian fusions with two major monophyletic groups showing either a diploid number of 2n=18 – where all of the acrocentric chromosomes are fused to produce metacentric elements, or a 2n=34 – where sex-chromosome translocations have been reported (Veyrunes et al. 2010). Additionally, WARTs have been documented in several populations exhibiting the 2n=18 karyotype in South Africa, which has contributed significantly to reported chromosomal variation, with at least four different cytotypes within this clade (Veyrunes et al. 2007). Currently, the geographic distributions of the 2n=18 and 2n=34 forms of *M. minutoides* are not known outside of the country of South Africa (Veyrunes et al. 2010).

Our objective was to utilize material from recent collecting efforts and molecular techniques to accurately delimit which species of *Nannomys* occur within Botswana. Further, we describe karyotypes for individuals from this region and make comparisons with previously published data from South Africa.

**Figure 1.** Distributions for three species of *Nannomys* in southern Africa. Dark grey indicates distribution for *Mus minutoides*, light grey for *M. indutus*, and stippled pattern for *M. setzeri*, adapted from Monadjem (2008a), Monadjem (2008b), and Monadjem and Coetzee (2008), respectively. Five trapping localities in Botswana (a); black crosses indicate captures for *M. minutoides* and grey triangles for *M. indutus*. Records from northwestern Botswana, Ngamiland District (b). Locality of syntopic records for *M. indutus* and *M. minutoides* at Koanaka Hills site (c).
Materials and methods

Our mitochondrial phylogeny was generated from combining previously published sequences deposited on GenBank (Appendix) with those derived from sequencing new specimens collected during field trips to Botswana conducted in 2008, 2009, and 2011 (Table 1, Appendix). We collected 16 specimens of *Mus* from five localities in Botswana including: Gcwihaba Caves (20°00.99'S; 21°15.89'E); Kang (23°32.10'S; 22°32.76'E); Koanaka Hills (20°09.60'S; 21°11.61'E); Lepokole Hills (21°49.59'S; 28°23.94'E); and Tsabong (25°56.57'S; 22°25.44'E) (Fig. 1a–c). Specimens were collected using Sherman live traps, pitfall traps, or Museum Special snap traps. Standard external measurements were recorded in the field (Table 1). Specimens were preserved as skins with complete skeletons (SSPS), skulls only, or as whole bodies in alcohol (alc.) and deposited at the at the Natural Science Research Laboratory (NSRL) at the Museum of Texas Tech University, Lubbock, Texas, USA or the Botswana National Museum, Gaborone, Botswana. Tissue samples were preserved in 95% ethanol, lysis buffer, or flash frozen in liquid nitrogen for future genomic analyses (2011 material) and deposited in the NSRL. Field collecting methods followed taxon specific guidelines for wild mammals (Sikes et al. 2012) as outlined by the Animal Care and Use Committee of the American Society of Mammalogists (Gannon et al. 2007; Sikes et al. 2011).

Genomic DNA was extracted using a DNeasy Blood and Tissue Kit (Qiagen Inc., Chatsworth, California). The complete cytochrome *b* gene (*cytb*, 1140 nucleotides) was amplified following methods outlined in Veyrunes et al. (2010). Cycle sequencing reactions were performed with BigDye terminator version 3.1 and were electrophoresed on an ABI 3100-Avant (Applied Biosystems, Foster City, California). Sequences were edited and aligned using SEQUENCHER version 4.9 (Gene Codes Corporation, Ann Arbor, Michigan). Novel sequences (GenBank accession nos. KF184308-KF184323) were aligned with previously published sequences deposited on GenBank using only individuals that exhibited unique haplotypes (Appendix). The final alignment was trimmed to exclude regions with large amounts of missing data due to the large number of GenBank sequences in the alignment that were partial *cytb* sequences. Therefore, a total of 741 base pairs of the *cytb* gene (the first 7 codons and last 126 codons were removed from the analysis) were used in the final alignment for the phylogenetic analysis including 125 individuals.

Appropriate models of evolution were examined using MEGA version 5 (Tamura et al. 2011). Phylogenetic relationships were estimated using Bayesian inference with the program MRBAYES version 3.2 (Huelsenbeck and Ronquist 2001). Four independent Markov chains were run for 50 million generations and trees were logged every 1000th iteration. Log-likelihood values were examined in the program TRACER version 1.5 (Rambaut and Drummond 2007) and the first 5,000 trees were discarded as burn-in. An additional phylogeny was estimated using the Maximum-likelihood method with the program PhyML version 3.0 (Guindon et al. 2010) with a BIONJ starting tree (Gascuel 1997) and 1,000 bootstrap replicates. Kimura 2-parameter genetic distances were calculated using MEGA version 5 (Tamura et al. 2011).
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Specimens were karyotyped in the field using bone marrow after 1 h of \textit{in vivo} incubation with Velban (Sigma-Aldrich, St. Louis, Missouri), following the methods described in Baker et al. (2003). \textit{Mus indutus} males were not karyotyped in this study because both males captured died in snap traps. Fluorescent \textit{in situ} hybridization (FISH) experiments were performed using Star*FISH © biotin-labeled mouse chromosome X paints (Cambio), following the manufacturer’s instructions and using Cy3-conjugated streptavidin (Invitrogen) for signal detection.

In order to assess the nature of the X-autosome translocation of the specimens that exhibited the translocation, we compared the X-chromosome of our specimens with those from South Africa using images of inverted DAPI-banding, and G-banding (Seabright 1971). Images were captured using the GENUS SYSTEM version 3.7 (Applied Imaging Systems, San Jose, California) through an Olympus BX51 epi-fluorescence microscope. Cy3 and DAPI (4’,6-diamidino-2-phenylindole) signals were pseudocolored yellow and red, respectively.

Results

The model with the lowest AICc (Akaike Information Criterion, corrected) and BIC (Bayesian Information Criterion) scores was the General Time Reversible (GTR) model using a discrete gamma distribution (+G) and a fraction of invariable sites (+I). Overall, the two methods of phylogenetic analysis resulted in similar tree topologies, except that the Maximum-likelihood analysis recovered weak support for the south + east \textit{M. minutoides} clade (Fig. 2). Additionally, the relationship between \textit{M. indutus}, \textit{M. sp.}, \textit{M. mattheyi}, \textit{M. haussa}, and the portion of the phylogeny that includes \textit{M. minutoides} and \textit{M. musculoides} was unresolved in the Maximum-likelihood analysis, though it was well-supported using Bayesian inference.

Sixteen \textit{cytb} sequences were generated from specimens from Botswana, corresponding to two species. Seven individuals are phylogenetically related to \textit{M. minutoides} from South Africa and nine individuals cluster with \textit{M. indutus}. Five individuals, captured from the same locality in the Koanaka Hills region of northwestern Botswana, represent two clades within \textit{M. minutoides} that are 5% different in \textit{cytb} sequence variation (Fig. 2). Six of the individuals of \textit{M. indutus} were collected in the Koanaka Hills alongside both of these lineages of \textit{M. minutoides} (Fig. 1c).

Karyotypes for individuals in the \textit{M. minutoides} clade exhibited a diploid number of 34 and fundamental number (as defined by Veyrunes et al. 2004 as the total number of chromosomal arms per diploid genome, instead of number of autosomal arms) of FN=36 (Fig. 3a–d, Table 2). All autosomes were acrocentric in morphology, including the pair 13, which presented a small short arm in some metaphase spreads. The metacentric X chromosome is the largest element of the chromosome complement, followed by the subtelocentric Y chromosome, which is comparable in size with the first autosomal pair. Individuals in the \textit{M. indutus} clade exhibited diploid and fundamental numbers of 36 (Fig. 3e–f, Table 2). All chromosomes had an acrocentric morphology. Due to
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The lack of male karyotyped specimens, the Y chromosome morphology could not be determined. The FISH with *Mus* X whole chromosome probe allowed the detection of an X-autosome translocation on the karyotypes of *M. minutoides* specimens (Fig. 3b, d), but not for individuals of *M. indutus* (Fig. 3e). Banding results indicate that individuals of *M. minutoides* from Botswana share the same sex-chromosome translocations (X.1) and (Y.1) as *M. minutoides* from South Africa, although differential condensation of the South African chromosomes makes direct comparison difficult (Fig. 3a and b).

**Figure 2.** Cytochrome *b* gene tree generated from 741 base pairs including 125 taxa using Bayesian inference. Grey boxes indicate species of interest: *Mus minutoides* and *Mus indutus*. Clades that include *Mus* from Botswana are enlarged to the right of the phylogeny. Diploid and fundamental numbers are shown for individuals sampled in this study and Veyrunes et al. (2005). Identification includes GenBank number and general locality. Support values at nodes are Bayesian posterior probabilities followed by Maximum-likelihood bootstrap support; dashes indicate regions of the tree where Maximum-likelihood analysis resulted in a polytomy.
Discussion

Efforts to resolve the geographic distributions of African pygmy mice remain in a state of flux. Regional studies involving DNA sequence data and karyotypes, such as presented here, contribute to a broader understanding of this complex genus. Historical
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(see Schmidt et al. 2008) and recent (Ferguson et al. 2010) bioinventories have resulted in extensive collections of Mus from Botswana, but there has been little consensus as to whether both M. minutoides and M. indutus occur in the country.

Mitochondrial sequence and cytogenetic data confirm the presence of both M. minutoides and M. indutus in Botswana. These specimens represent the first DNA sequences for these two species in Botswana, which we also made available for use in a recent paper by Lamb et al. (2014). Despite previous suggestions that M. minutoides and M. indutus occur in allopatry, our results confirm that these two species occur in sympatry and even syntopy in northwestern Botswana. Interestingly, we also found two lineages of M. minutoides in northwestern Botswana (Koanaka Hills) that were 5% different in cytb sequence variation. We hypothesize that these two mitochondrial lineages were separated in the past and have now come back together in a region of secondary contact in the arid savannah region near the Kalahari Desert, a hypothesis that should be tested with broader sampling and using additional genetic markers.

Also of interest is the fact that no M. setzeri were collected from either the Koanaka Hills or Gcwihaba Caves although their current range – as delimited by Monadjem and Coetzee (2008) and Skinner and Chimimba (2005) – includes this region of Botswana. We compared our specimens with M. setzeri deposited at the National Museum of Natural History, Smithsonian Institution, Washington D.C., USA and found no evidence that any of our individuals correspond to this conspicuous form. Our failure to capture M. setzeri, in spite of concerted trapping efforts in this region (> 2600 Sherman trap nights, > 280 pitfall trap nights during June 2008 and July 2009 seasons), is in agreement with Monadjem (2013b) who pointed to the scarcity of this species in collections as evidence for true ecological rarity. Further sampling is clearly warranted to more accurately delimit the exact geographic boundaries of Nannomys species both within Botswana and throughout the broader Southern African Subregion (Skinner and Chimimba 2005).

Mus minutoides in Botswana exhibit the 2n=34 karyotype with the diagnostic (X.1) and (Y.1) sex-autosome translocations that have also been documented in specimens from South Africa (Veyrunes et al. 2010), Zambia, Kenya (Castiglia et al. 2002, 2006), Central African Republic, and Ivory Coast (Jotterand-Bellomo 1984, 1986). Veyrunes et al. (2004) propose that 2n=34 with the 1 sex chromosome translocation is the ancestral karyotype for M. minutoides and our results provide further support for an early (X.1) translocation before the radiation of M. minutoides over a large geographic area. Furthermore, the 2n=34 cytotype is reported in several locations in northern South Africa, but not in southern South Africa or in other countries to the north, including Botswana. The fact that our sampling localities included individuals from the easternmost and northwestern regions of Botswana might be an indicator that this is the predominant cytotype in the country, likely extending into the bordering countries of Zambia, Zimbabwe, and Namibia.

We found that three of our gender identifications made in the field (Table 2, “Gender Field”) did not match the identifications made from karyotype assessments (Table 2, “Gender Lab”) indicating the potential for X*Y females. Therefore, we attempted to
Table 2. Individuals of *Mus (Nannomys)* collected in Botswana including GenBank number, final species identification, gender determined in the field, museum preparation type (Alcoholic=alc; skin, skull, postcranial skeleton=SSPS; or Skull only), collection date, total length (TL), tail length (T), hind foot (HF), ear (E), weight in grams, karyotype, and sex-chromosome.

| Genbank No. | Species     | Gender “Field” | Prep. Type | Coll. Date  | TL | T  | HF | E | Weight (g) | Karyotype         | Gender “Lab” |
|-------------|-------------|----------------|------------|-------------|----|----|----|----|------------|-------------------|---------------|
| KF184315    | *M. indutus*| Female         | SSPS       | 16-Jul-09   | 95 | 42 | 13 | 13 | 4.5        | 2n=36, FN=36     | XX            |
| KF184316    | *M. indutus*| Female         | SSPS       | 22-Jul-09   | 85 | 40 | 10 | 10 | 2.9        | none              | -             |
| KF184317    | *M. indutus*| Female         | SSPS       | 27-Jul-09   | 101| 43 | 14 | 11 | 5.1        | none              | -             |
| KF184318    | *M. indutus*| Female         | SSPS       | 22-Jul-09   | 14 | 45 | 12 | 10 | 6.3        | 2n=36, FN=36     | XX            |
| KF184319    | *M. indutus*| Female         | SSPS       | 15-Jul-09   | 110| 45 | 13 | 11 | 6.75       | 2n=36, FN=36     | XX            |
| KF184320    | *M. indutus*| Male           | SSPS       | 18-Aug-11   | 98 | 45 | 15 | 10 | 4          | none              | -             |
| KF184321    | *M. indutus*| Female         | Alc        | 25-Aug-11   | 75 | [23]| 14 | 10 | 3          | none              | -             |
| KF184322    | *M. indutus*| Female         | Skull Only | 17-Aug-11   | 109| 40 | 15 | 11 | 5          | none              | -             |
| KF184323    | *M. indutus*| Male           | SSPS       | 20-Jul-09   | 86 | 42 | 14 | 12 | 4          | none              | -             |
| KF184308    | *M. minutoides*| Female      | SSPS       | 20-Jul-09   | 107| 43 | 14 | 12 | 5.5        | 2n=34, FN=36     | XX            |
| KF184309    | *M. minutoides*| Male       | SSPS       | 24-Jul-09   | [80]| [23]| 13 | 11 | 4.6        | 2n=34, FN=36     | XY            |
| KF184310    | *M. minutoides*| Female     | SSPS       | 16-Aug-11   | 93 | 45 | 13 | 10 | 3.5        | 2n=34, FN=36     | XY            |
| KF184311    | *M. minutoides*| Male       | SSPS       | 26-Jun-08   | 102| 47 | 14 | 12 | 5.8        | none              | -             |
| KF184312    | *M. minutoides*| Female     | SSPS       | 15-Jul-09   | 111| 52 | 15 | 11 | 5.5        | 2n=34, FN=36     | XX            |
| KF184313    | *M. minutoides*| Female     | SSPS       | 20-Jul-09   | 99 | 44 | 12 | 9  | 4          | 2n=34, FN=36     | XY            |
| KF184314    | *M. minutoides*| Female     | SSPS       | 26-Jul-09   | 96 | 47 | 14 | 10 | 3.7        | 2n=34, FN=36     | XY            |
Mitochondrial DNA and karyotypic data confirm the presence of *Mus indutus*... examine these specimens for the possibility of sex reversal in *M. minutoides*, which has been documented in other countries (Veyrunes et al. 2013). Although we have tried to identify the X* chromosome in our samples through X chromosome morphology assessment as well as DAPI banding patterns, the particular high degree of condensation of the chromosomes in our *in vivo* bone marrow preparations did not allow us to ascertain the nature of the X chromosomes of two of these three specimens. For one of the individuals, both the morphology and banding patterns of the X chromosome do not seem to correspond to those of the derivative X* chromosome (Fig. 3a), indicating that field misidentification of sex might have been the case for that specimen (the reproductive organs can no longer be clearly seen on the prepared skin of this specimen). Additionally, there were no evident X chromosome polymorphisms in the XX female specimens, which would be expected in populations where X*Y females were present. Due to our small sample, and the relative low frequency of the X* found in populations outside South Africa, we were not able to rule out the presence of the X polymorphism in Botswana. Further collecting efforts, together with an in depth sex determination study, including high quality chromosome preparations suitable for G-banding studies, will be needed to shed further light on this issue.

Our data presented here agree with previous molecular phylogenies of *Nannomys*, with well-defined clades representing *M. minutoides* and *M. indutus* exhibiting diploid and fundamental numbers consistent with those reported in the literature. Veyrunes et al. (2010) detected a wide range of chromosomal variation for *M. minutoides* in South Africa, with one particular clade presenting 2n=34, FN=36. Our *M. minutoides* samples display chromosome conservation as well as sequence similarity to the South African clade bearing karyotypic stasis, indicating that these specimens might be part of a widespread group chromosomally and genetically isolated from the karyotypically diverse 2n=18 *M. minutoides* clade. *Mus indutus* on the other hand exhibits a karyotype not very divergent from the proposed ancestral karyotype for *Nannomys* (2n=36 with all acrocentric chromosomes; Veyrunes et al. 2004), similar to many of the basal lineages included in recent molecular phylogenies (see Britton-Davidian et al. 2012).

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### Appendix

Individuals included in the molecular phylogeny representing eleven species with the country of origin, GenBank number and the original citation for the original description. RCA = Central African Republic.

| Species | Country | Genbank No. | Reference |
|---------|---------|-------------|-----------|
| *Mus (Nannomys)* | | | |
| *baoulei* | Benin | EU603991-92 | Kan Kouassi et al. 2008 |
| | Guinea | EU603995 | Kan Kouassi et al. 2008 |
| | Ivory Coast | EU603993-94, 98 | Kan Kouassi et al. 2008 |
| *bufo* | Burundi | DQ789905 | Mboumba et al. 2011 |
| *hausta* | Chad | AY875071 | Veyrunes et al. 2005 |
| | Mali | AY698877 | Chevret et al. 2005 |
| | Niger | AY875072-73 | Veyrunes et al. 2005 |
| | Senegal | AY875074 | Veyrunes et al. 2005 |
| *indutus* | Botswana | KF184315-23 | This paper |
| | South Africa | AY698874 | Chevret et al. 2005 |
| | South Africa | AY875070 | Veyrunes et al. 2005 |
| *mattheyi* | Burkina Faso | AY877114 | Veyrunes et al. 2005 |
| | Guinea | EU603970-73 | Kan Kouassi et al. 2008 |
| | Mali | AY698876 | Chevret et al. 2005 |
| | Mali | AY875066-67 | Veyrunes et al. 2005 |
| | Senegal | AB125781 | Suzuki et al. 2004 |
| | Senegal | AY875068 | Veyrunes et al. 2005 |
| | Togo | AY875069 | Veyrunes et al. 2005 |
| *minutoides* | Botswana | KF184308-14 | This paper |
| | Congo | DQ789929 | Mboumba et al. 2011 |
| | Gabon | DQ789911, 20, 26 | Mboumba et al. 2011 |
| | Guinea | AY875076-77 | Veyrunes et al. 2005 |
| | Guinea | EU603936-37, 60-61, 64-65 | Kan Kouassi et al. 2008 |
| | Ivory Coast | EU603925-28, 30-33,35, 45, 47,49, 54-56, 58, 999, 001-02, 005 | Kan Kouassi et al. 2008 |
| | Kenya | AY875084 | Veyrunes et al. 2005 |
| | Kenya | AY057816 | Lundrigan et al. 2002 |
| | RCA | DQ789938-39 | Mboumba et al. 2011 |
| | South Africa | AY875078-80 | Veyrunes et al. 2005 |
| | South Africa | FN985222-24 | Veyrunes et al. 2010 |
| | Tanzania | AY875081 | Veyrunes et al. 2005 |
| *musculoides* | Cameroon | HM635855-56 | Dobigny et al. 2011 |
| | Guinea | EU603968-69 | Kan Kouassi et al. 2008 |
| | Guinea | DQ789902 | Mboumba et al. 2011 |
| | Ivory Coast | EU603967 | Kan Kouassi et al. 2008 |
| | Ivory Coast | DQ789901 | Mboumba et al. 2011 |
| | Mali | Z96069 | Barome et al. 1998 |
| | Mali | AY698875 | Chevret et al. 2005 |
| | Mali | AY875075 | Veyrunes et al. 2005 |
| | Mali | JX292892-93 | Schwan et al 2012 |
Mitochondrial DNA and karyotypic data confirm the presence of *Mus indutus*...  

| **Mus** (Nannomys) | **Country** | **Genbank No.** | **Reference** |
|--------------------|-------------|-----------------|---------------|
| setulosus          | Cameroon    | EU603989        | Kan Kouassi et al. 2008 |
|                    | Cameroon    | DQ789900        | Mboumba et al. 2011 |
|                    | Gabon       | AJ698873        | Chevret et al. 2005 |
|                    | Guinea      | AJ875083        | Veyrunes et al. 2005 |
|                    | Guinea      | EU603976, 78, 82-83, 86 | Kan Kouassi et al. 2008 |
|                    | Ivory Coast | EU603974-75, 77, 79-81, 84-85, 88, 97 | Kan Kouassi et al. 2008 |
|                    | Ivory Coast | GU830865, 67, 69 | Coulibaly-N’golo et al. 2011 |
|                    | RCA         | AJ875082        | Veyrunes et al. 2005 |
|                    | RCA         | EU603990        | Kan Kouassi et al. 2008 |
| sorella            | RCA         | DQ789904        | Mboumba et al. 2011 |
| M. sp.             | Chad        | AJ875085        | Veyrunes et al. 2005 |
| tenellus           | Ethiopia    | DQ789903        | Mboumba et al. 2011 |