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The phylogenetic position of the enigmatic Atlantic forest-endemic spiny mouse *Abrawayaomys* (Rodentia: Sigmodontinae)

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Abstract

Background: The phylogenetic position of the sigmodontine genus *Abrawayaomys*, historically assigned to the tribe Thomasomyini or considered a sigmodontine incertae sedis, was assessed on the basis of nuclear and mitochondrial DNA sequences obtained from four individuals from different localities in the Atlantic forest of Brazil. Sequences of *Abrawayaomys* were analyzed in the context of broad taxonomic matrices by means of maximum-likelihood (ML) and Bayesian analyses (BA).

Results: The phylogenetic position of *Abrawayaomys* differed depending on the gene analyzed and the analysis performed (interphotoreceptor retinoid-binding protein (IRBP) ML: sister to Thomasomyini; IRBP BA: sister to Akodontini; cytochrome (Cyt) *b* ML: sister to *Neotomys*; and Cyt *b* BA: sister to Reithrodontini). With the sole exception of the BA based on Cyt *b* sequences, where the *Abrawayaomys-Reithrodon* clade had strong support, all sistergroup relationships involving *Abrawayaomys* lacked any significant support.

Conclusions: As such, *Abrawayaomys* constitutes the only representative so far known of one of the main lineages of the sigmodontine radiation, differing from all other Atlantic forest sigmodontine rodents by having a unique combination of morphological character states. Therefore, in formal classifications, it should be regarded as a Sigmodontinae incertae sedis.

Keywords: Akodontini; Atlantic forest; Cricetidae; Phylogeny; Thomasomyini

Background

With about 86 living genera, cricetids of the subfamily Sigmodontinae are one of the most diversified and taxonomically complex groups of mammals. Predominantly distributed in South America, sigmodontines also reach Central and North America, and one extant genus is endemic to the Galapagos Islands (D'Elía 2003a). Remarkably, new sigmodontine genera are still being erected on the basis of both revisionary museum work and from newly discovered species (e.g., Pardiñas et al. 2009a; Percequillo et al. 2011; Pine et al. 2012; Alvarado-Serrano and D'Elía 2013; see comments in D'Elía and Pardiñas 2007). Similarly, statuses of several sigmodontine taxonomic forms at

the species level are unclear (e.g., Alarcón et al. 2011; Bonvicino et al. 2012).

Sigmodontine genera have been arranged into different groups, most of which have been given the formal rank of tribes (e.g., Reig 1980). In the last two decades, phylogenetic analyses using either morphological or molecular data or both were used to set the limits and contents of these groups and determine the timing of their diversification (e.g., Braun 1993; Smith and Patton 1999; Steppan 1995; D'Elía 2003b; Pacheco 2003; Weksler 2003; D'Elía et al. 2003, D'Elía et al. 2006a, b; Martínez et al. 2012; Parada et al. 2013; Salazar-Bravo et al. 2013). Those studies caused a number of major reconsiderations on the limits and contents of these groups. Currently, 12 extant genera are considered as Sigmodontinae incertae sedis (see the most recently published classification in D'Elía et al. (2007) and the modification prompted by the description of a new

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genus by Alvarado-Serrano and D'Elía (2013)); one of these is *Abrawayaomys* Cunha and Cruz 1979.

Abrawayaomys is a poorly known sylvan sigmodontine genus that stands out within the sigmodontine radiation due to its spiny pelage and unusual craniodental morphology. It is found in the Atlantic forest of Argentina and Brazil and is known from a handful of trapped specimens and a few osteological remains gathered from owl pellets (Pardiñas et al. 2009b). Two species are recognized: the type species Abrawayaomys ruschii Cunha and Cruz 1979 and the recently described Abrawayaomys chebezi (Pardiñas et al. 2009a, 2009b).

Abrawayaomys displays a striking combination of morphological features that was referred by Musser and Carleton (2005, p. 1,088) in the following terms 'Diagnostic traits seem to combine aspects of Neacomys, Oryzomys, and Akodon, and both Reig (1987) and Smith and Patton (1999) acknowledged the enigmatic affinities of Abrawayaomys as uncertain.' Recently, Pardiñas et al. (2009b) evaluated the morphology of Abrawayaomys in detail and noted a certain resemblance to the bauplan of the tribe Akodontini but also to the thomasomyine genera Chilomys and Rhagomys. Similarities with Akodontini were regarded as convergences since Abrawayaomys was assigned to Thomasomyini given that independent sets of data placed Chilomys and Rhagomys in Thomasomyini (D'Elía et al. 2006a; Salazar-Bravo and Yates 2007) and because morphology-based phylogenetic analyses placed Abrawayaomys in this tribe (Pacheco 2003; Salazar-Bravo and Yates 2007). The lack of molecular data for Abrawayaomys has prevented an assessment of its position within the available comprehensive sigmodontine trees generated on the basis of mitochondrial and nuclear DNA sequences (e.g., Engel et al. 1998; Smith and Patton 1999; D'Elía 2003b; Weksler 2003; D'Elia et al. 2006a, b).

In recent fieldwork in the states of Rio de Janeiro (Pereira et al. 2008), Minas Gerais (Passamani et al. 2011), and São Paulo (see below) in southeastern Brazil, four specimens referred to as *A. ruschii* were collected. Based on mitochondrial and nuclear DNA sequences gathered from these specimens, we present the first phylogenetic analyses to test previous hypotheses concerning the placement of *Abrawayaomys* within the sigmodontine radiation. In addition, we provide some taxonomic judgments based on the resulting phylogeny and comments on the evolution of some morphological traits.

Methods

DNA sequences corresponding to the cytochrome (Cyt) *b* gene and the first exon of the nuclear interphotoreceptor retinoid-binding protein (IRBP) gene were used as evidence. We sequenced four specimens of *A. ruschii* deposited in the following Brazilian collections: Museu Nacional, Univ. Federal do Rio de Janeiro (MN 67557; Brazil, Rio

de Janeiro, Aldeia Sapucai) (Pereira et al. 2008); Museu de Zoologia, Univ. de São Paulo (MZUSP 32319; Brazil, São Paulo, Biritiba-Mirim; BO 27; Brazil, São Paulo, Estação Ecológica de Boracéia); and Coleção de Mamíferos da Univ. Federal de Lavras (CMUFLA 906; Brazil, Minas Gerais, Caeté) (Passamani et al. 2011). Sequences were gathered following the protocol of Pardiñas et al. (2003) and D'Elía et al. (2006b). We found minor differences among Cyt b sequences (see 'Results' below), but in the phylogenetic analyses, we used that of specimen MN 67557 because it was the only complete one (i.e., 1,140 bp; the other being 801 bp long). We found no variation in IRBP sequences of the four specimens analyzed, and therefore, Abrawayaomys was represented by a single terminal (MN 67557) in the phylogenetic analyses. New sequences were submitted to GenBank [GenBank: JX949182 to JX949189].

To appraise the phylogenetic position of Abrawayaomys within the radiation of the Sigmodontinae, we sought to ensure that sigmodontine diversity was represented as comprehensively as possible. According to the current classification, our sampling only lacked for both matrices the incertae sedis genera *Phaenomys* and *Wilfredomys*; the akodontines Gyldenstolpia and Podoxymys; the ichthyomyines Anotomys, Chibchanomys, Neusticomys, and Ichthyomys; and the oryzomyine Mindomys. In addition, the IRBP matrix lacked the thomasomyine Chilomys. Meanwhile, the ichthyomyine Rheomys, the oryzomyine Microakodontomys, and the thomasomyine Aepeomys were also missing from the Cyt b matrix. Therefore, both the IRBP (1,181 characters) and Cyt b (1,134 characters) matrices respectively included representatives of 76 and 74 sigmodontine genera (including sequences of the recently described genus Neomicroxus). The IRBP matrix lacked a sequence for Neusticomys available in GenBank [GenBank: EU649036] because a recent inspection of it indicated that it may be in fact a composite of an ichthyomyine and an oryzomyine sequence. Although the monophyly of the Sigmodontinae is well corroborated (e.g., Engel et al. 1998; Steppan et al. 2004; Parada et al. 2013), the identity of its sister group is unclear. Therefore, we integrated the outgroup with two representatives of each of the other four main lineages that, together with the Sigmodontinae, compose the family Cricetidae: arvicolines (Arvicola and Microtus), cricetines (Cricetulus and Phodopus), neotomines (Neotoma and Scotinomys), and tylomyines (Nyctomys and Tylomys). All taxa represented in the analyses, along with the GenBank accession numbers of their DNA sequences, are listed in Table 1.

Alignment was done with Clustal X (Thompson et al. 1997) using default parameters for all alignment parameters. Uncorrected genetic distances (*p* distances) with pairwise deletions were computed using MEGA 5 (Tamura et al. 2011). Each matrix was subjected to maximum-likelihood (ML) (Felsenstein 1981) and Bayesian analyses (BA)

Table 1 List of taxa and the DNA sequences of which were included in the phylogenetic analyses

Tribe	Species	IRBP	Cyt b
Abrotrichini	Abrothrix longipilis	AY163577	U03530
Abrotrichini	Chelemys macronyx	emys macronyx AY277441	
Abrotrichini	Geoxus valdivianus	AY277448	AY275116
Abrotrichini	Notiomys edwardsii	AY163602	U03537
Abrotrichini	Pearsonomys annectens	AY851749	AF108672
Akodontini	'Akodon' serrensis	EF626799	AY273908
Akodontini	Akodon azarae	AY163578	DQ444328
Akodontini	Bibimys chacoensis/labiosus	AY277435	DQ444329
Akodontini	Blarinomys breviceps	AY277437	AY275112
Akodontini	Brucepattersonius soricinus	AY277439	AY277486
Akodontini	Deltamys kempi	AY277444	AY195862
Akodontini	Juscelinomys huanchacae	AY277453	AF133667
Akodontini	Kunsia tomentosus	AY277455	AY275120
Akodontini	Lenoxus apicalis	AY277456	U03541
Akodontini	Necromys lasiurus	AY277459	AY273912
Akodontini	Oxymycterus nasutus	AY277468	EF661854
Akodontini	Scapteromys tumidus	AY163637	AY275133
Akodontini	Thalpomys cerradensis	AY277481	AY273916
Akodontini	Thaptomys nigrita	AY277482	AF108666
Ichthyomyini	Rheomys raptor	AY163635	-
Incertae sedis	Abrawayaomys ruschii	JX949185	JX949189
Incertae sedis	Andinomys edax	JQ434400	AF159284
Incertae sedis	Chinchillula sahamae	JQ434409	JQ434422
Incertae sedis	Delomys sublineatus	AY163582	AF108687
Incertae sedis	Euneomys chinchilloides	AY277446	AY275115
Incertae sedis	Irenomys tarsalis	AY163587	U03534
Incertae sedis	Juliomys pictipes	AY163588	AF108688
Incertae sedis	Neomicroxus latebricola	QCAZ4160	QCAZ4160
Incertae sedis	Neotomys ebriosus	HM061605	HM061604
Incertae sedis	Punomys kofordi	JQ434414	JQ434426
Oryzomyini	Aegialomys xanthaeolus	GQ178247	EU579479
Oryzomyini	Amphinectomys savamis	AY163579	EU579480
Oryzomyini	Cerradomys scotti	EU649040	EU579482
Oryzomyini	Drymoreomys albimaculatus	EU649042	EU579487
Oryzomyini	Eremoryzomys polius	AY163624	EU579483
Oryzomyini	Euryoryzomys macconnelli	AY163620	EU579484
Oryzomyini	Handleyomys intectus	AY163584	EU579490
Oryzomyini	Holochilus brasiliensis	AY163585	EU579496
Oryzomyini	Hylaeamys megacephalus	AY163621	EU579499
Oryzomyini	Lundomys molitor	AY163589	EU579501
Oryzomyini	Melanomys caliginosus	EU649052	EU340020
Oryzomyini	Microakodontomys transitorius	EU649054	-
Oryzomyini	Microryzomys minutus	AY163592	AF108698
01,2011,111			

Table 1 List of taxa and the DNA sequences of which were included in the phylogenetic analyses (Continued)

were included in the phylogenetic analyses (Continued)								
Oryzomyini	Nectomys squamipes	AY163598	EU340012					
Oryzomyini	Nephelomys albigularis	EU649057	EU579505					
Oryzomyini	Nesoryzomys fernandinae	EU649058	EU579506					
Oryzomyini	Oecomys bicolor	AY163604	EU579509					
Oryzomyini	Oligoryzomys fulvescens	AY163611	DQ227457					
Oryzomyini	Oreoryzomys balneator	AY163617	EU579510					
Oryzomyini	Oryzomys palustris	AY163623	EU074639					
Oryzomyini	Pseudoryzomys simplex	AY163633	EU579517					
Oryzomyini	Scolomys ucayalensis	AY163638	EU579518					
Oryzomyini	Sigmodontomys alfari	AY163641	EU340016					
Oryzomyini	Sooretamys angouya	AY163616	EU579511					
Oryzomyini	Tanyuromys aphrastus	JF693878	JF693877					
Oryzomyini	Transandinomys talamancae	AY163627	EU579514					
Oryzomyini	Zygodontomys brevicauda	AY163645	EU579521					
Phyllotini	Andalgalomys pearsoni	EU649038	AF159285					
Phyllotini	Auliscomys pictus	AY277434	JQ434420					
Phyllotini	Calomys lepidus	AY163580	EU579473					
Phyllotini	Eligmodontia typus	AY277445	AF108692					
Phyllotini	Galenomys garleppi	JQ434410	JQ434423					
Phyllotini	Graomys griseoflavus	EU649037	EU579472					
Phyllotini	Loxodontomys micropus	AY277457	AY275122					
Phyllotini	Phyllotis xanthopygus	AY163632	AY275128					
Phyllotini	Salinomys delicatus	JQ434415	EU377608					
Phyllotini	Tapecomys primus	JQ434416	AF159287					
Phyllotini	Phyllotini n. gen.	JQ434417	JQ434425					
Reithrodontini	Reithrodon auritus	AY163634	EU579474					
Sigmodontini	Sigmodon alstoni	EU635698	AF293397					
Thomasomyini	Aepeomys lugens	DQ003722	-					
Thomasomyini	Chilomys instans	-	AF108679					
Thomasomyini	Rhagomys longilingua/rufescens	DQ003723	AY206770					
Thomasomyini	Rhipidomys macconnelli	AY277474	AY275130					
Thomasomyini	Thomasomys aureus	AY277483	U03540					
Wiedomyini	Wiedomys pyrrhorhinos	AY163644	EU579477					
Outgroup	Arvicola terrestris	AY277407	AY275106					
	Cricetus cricetus	AY277410	AY275109					
	Microtus socialis	FM162055	AY513830					
	Neotoma albigula	AY277411	AF108704					
	Nyctomys sumichrasti	AY163603	AY195801					
	Phodopus sungorus	AY163631	JN015007					
	Scotinomys xerampelinus	AY277416	AF108706					
	Tylomys nudicaudus	AY163643	DQ179812					

Sequences of the genera *Bibimys* and *Rhagomys* of each gene were gathered from different species. GenBank accession numbers for each gene (IRBP and Cyt b) are indicated in the last two columns. Tribal assignations follow D'Elía et al. 2007 (see also D'Elía and Pardiñas 2007) and results of the present study.

(Rannala and Yang 1996). The ML analysis was conducted in Treefinder (Jobb et al. 2004; Jobb 2008). The best fitting models of nucleotide substitution (IRBP: TVM[Optimum, Empirical]:G[Optimum]:5; Cyt b: GTR[Optimum, Empirical]:G[Optimum]:5) (see Jobb 2008) were selected with the Akaike information criterion in Treefinder using the 'propose model' routine. The best tree was searched under the model of nucleotide substitutions previously selected using search algorithm 2 implemented in Treefinder version March 2011; nodal support was estimated with 1,000 bootstrap pseudoreplicates (BS). The BA was conducted using MrBayes 3.1.2 (Ronquist and Huelsenbeck 2003) by means of two independent runs with three (IRBP) or seven (Cyt b) heated and one cold Markov chains each. Considering the model selected by Treefinder and models specified in MrBayes, a model with six categories of a base substitution, a gamma-distributed rate parameter, and a proportion of invariant sites was specified for both matrices; all model parameters were estimated using MrBayes. Runs were allowed to proceed for 20 million (IRBP) and 28 million (Cyt b) generations, and trees were sampled every 1,000 (IRBP) and 2,000 (Cyt b) generations. Log-likelihood values were plotted against the generation time to check that runs converged on a stable log-likelihood value. The first 25% of sampled trees were discarded as burn-in; the remaining trees were used to compute a 50% majority rule consensus tree and obtain posterior probability (PP) estimates for each clade.

Results

Cyt b gene sequences of Abrawayaomys gathered from the three specimens collected in the states of São Paulo and Rio de Janeiro varied by $0.5\% \sim 0.7\%$; while comparisons involving the Cyt b sequence of the specimen from Minas Gerais ranged $2.6\% \sim 3.0\%$. As mentioned above, IRBP sequences of the four specimens analyzed were identical.

Abrawayaomys, as represented by sequences of specimen MN 67557, was highly divergent from all compared sigmodontines for both analyzed genes. For the Cyt b gene, observed pairwise values involving Abrawayaomys ranged 19.0% \sim 25.9% for comparisons with Neotomys and Lundomys. For the IRBP gene, observed pairwise values involving Abrawayaomys ranged $3.42\% \sim 6.65\%$ for comparisons with Brucepattersonius and Rheomys.

Phylogenetic analyses recovered general results congruent with those of previous IRBP- and/or Cyt b-based studies (e.g., D'Elia et al. 2006a, b; Percequillo et al. 2011; Martínez et al. 2012; Parada et al. 2013; Salazar-Bravo et al. 2013). For descriptions and discussions of these findings (e.g., a strongly supported Sigmodontinae, a strongly supported Oryzomyalia, the polyphyly of the *Reithrodon* group, and differences between gene trees), we refer the reader to those studies because herein we focus on the phylogenetic position of *Abrawayaomys*.

Depending on the gene analyzed, the phylogenetic position of *Abrawayaomys* varied from being sister to Akodontini (IRBP BA; PP = 0.62; Figure 1), sister to Thomasomyini (IRBP ML; BS < 50), sister to Reithrodontini (Cyt b BA; PP = 0.98; Figure 2), or sister to *Neotomys* (Cyt b ML; BS < 50). As noted, with the sole exception of the BA based on Cyt b sequences, where the *Abrawayaomys-Reithrodon* clade had strong support, all sister-group relationships involving *Abrawayaomys* lacked any significant support. The most inclusive and well-supported clade containing *Abrawayaomys* was that corresponding to Oryzomyalia (*sensu* Steppan et al. 2004).

Discussion

The diversity of sigmodontine forms has long captivated students of New World mammals, but at the same time, it has seriously defied those attempting to classify them according to their evolutionary history. Problems range from species boundaries to relationships among species and genera, and limits and contents of higher taxa (e.g., tribes). These issues have direct implications for the study of the history of the diversification of the group, which in turn arguably constitutes one of the most controversial debates in muroid systematics (Voss 1993; D'Elía 2003b).

Herein, we showed that the phylogenetic position of Abrawayaomys varies from being sister to Akodontini, Thomasomyini, Reithrodontini, or Neotomys, depending on the gene analyzed (IRBP or Cyt b) and the analysis performed (ML or BA). Given that previous studies showed discrepancies between the topologies of different sigmodontine gene trees (e.g., Feijoo et al. 2010; Teta et al. 2011), the fact that different datasets (i.e., IRBP or Cyt b matrices with slightly different taxonomic sampling) provide different relationships for Abrawayaomys is not unexpected. Importantly, with the exception of the BA based on Cyt b sequences (sister to Reithrodontini), no sister relationship involving Abrawayaomys was recovered that had good support. In light of the analytical results, Abrawayaomys cannot be placed with certainty in any more inclusive clade than that of Oryzomyalia (sensu Steppan et al. 2004).

Pacheco (2003, p. 130), in a phylogenetic analysis based on morphological characters, found the genus *Abrawayaomys* to be sister to *Rhagomys* within the Thomasomyini clade. This relationship was supported by the following character states: a broad zygomatic plate, an interorbital region convergent with the supraorbital margins squared or weakly beaded, a long jugal, the absence of mesolophids (but see below), a masseteric crest anterior to the procingulum of the first lower molar, and a deeply excavated retromolar region of the mandible. Pacheco (2003) highlighted the retromolar region condition, i.e., broad and fenestrated, as a synapomorphy of *Abrawayaomys + Rhagomys*. We agree with Pacheco

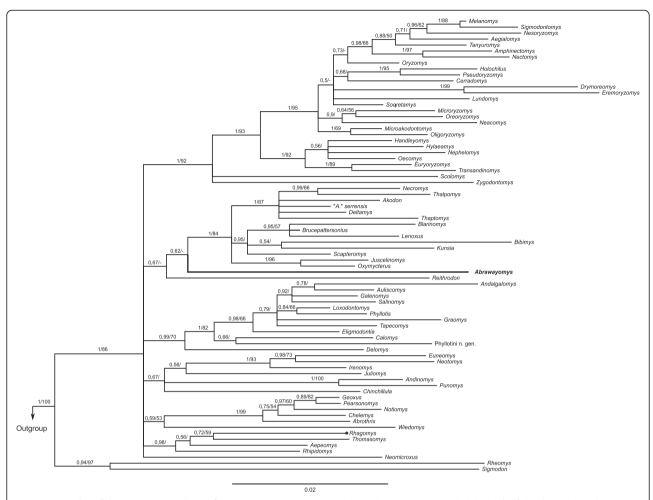


Figure 1 Results of the Bayesian analysis of IRBP gene sequences. Numbers indicate posterior probability (left of the diagonal) and ML bootstrap values (right of the diagonal) of adjacent nodes. Only bootstrap values of >50% are shown. Dashes signal nodes that were not recovered in the ML topology (ln = -9.764.8433). Dots at terminals indicate genera with species possessing dorsal spines.

(2003) (see Musser and Carleton 2005) in the general resemblance between Abrawayaomys and Rhagomys and to a lesser extent to the remainder of the Thomasomyini, but we suggest that this similarity is not remarkable and more important and that these shared character states are not synapomorphies of a putative Abrawayaomys and Rhagomys clade. Almost all of those character states listed by Pacheco (2003) are also present in many sigmodontines, indicating the large amount of homoplasy existing within this group. In addition, we assert that the molar morphologies of both genera are quite distinct, having only the widespread brachyodont condition in common. Rhagomys has very well-developed mesolophs/phids (cf. Luna and Patterson (2003) vs. Pacheco (2003)), procingula of the first upper molars clearly crossed by a deep anteromedian flexus, well-developed posterolophs, a slightly reduced third lower molar with respect to the second lower molar, and several other traits found among taxa displaying the dental bauplan of the pentalophodont type

(sensu Hershkovitz 1962), which is clearly distinguishable from the unequivocally tetralophodont molar of Abrawayaomys (cf. Pardiñas et al. 2009b). Additional differences between these two genera are more than trenchant, including incisive foramina and palate extensions, parapterygoid plate morphology, and carotid circulatory pattern (Table 2) (see also Pardiñas et al. 2009b, Table three). Similarly, several trenchant character states are present in Abrawayaomys and representatives of other sigmodontine tribes. The external morphology Abrawayaomys resembles that of many akodonts (cf. Pereira et al. 2008, Figure one), although it has a moderately longer tail, at least in some individuals. A morphological description of the stomach (Finotti et al. 2003) suggests a hemiglandular-unilocular type, a widespread condition among sigmodontines (Carleton 1973). Finally, Pacheco (2003) also indicated that *Abrawayaomys* has a peculiar genal vibrissa (called genal vibrissa 2), which is also present in the oryzomyine Oecomys and

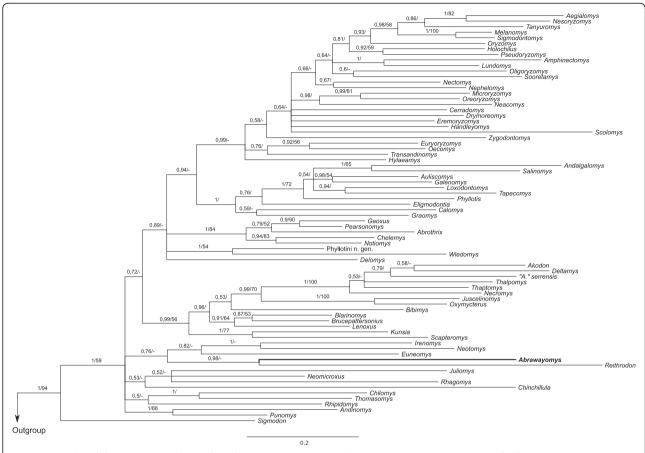


Figure 2 Results of the Bayesian analysis of Cyt b gene sequences. Numbers indicate posterior probability (left of the diagonal) and ML bootstrap values (right of the diagonal) of adjacent nodes. Only bootstrap values of >50% are shown. Dashes signal nodes that were not recovered in the ML topology (ln = -33,444.453).

the akodontine Kunsia, and a reduced fifth pedal digit. Similarly, karyotypic evidence sheds no conclusive light on the phylogenetic position of Abrawayaomys, as the diploid number of 2n = 58 (Pereira et al. 2008) found in A. ruschii is also present in several distantly related oryzomyine species, such as Euryoryzomys lamia (Andrades-Miranda et al. 2000), Holochilus brasiliensis (Yonenaga-Yassuda et al. 1987), Nectomys squamipes (Yonenaga-Yassuda et al. 1987), Oecomys trinitatis (Patton et al. 2000), Oligoryzomys chacoensis (Myers and Carleton 1981), and Sooretamys angouya (Andrades-Miranda et al. 2000). Taking all this evidence as a whole, Abrawayaomys cannot be placed with certainty in any clade less inclusive than the large clade Oryzomyalia (see also Voss 1993). Therefore, in light of all of the evidence at hand, in formal classifications, Abrawayaomys should be kept as an incertae sedis sigmodontine (D'Elía et al. 2007). Results of future phylogenetic analyses, especially those including sequences of two other Atlantic forest inhabitants, Phaenomys and Wilfredomys, which were also considered part of the tribe Thomasomyini (e.g., Pacheco 2003) and are now regarded as Sigmodontinae incertae sedis (e.g., D'Elía et al. 2007), may prompt changes in this classification.

The prevailing biogeographic view is that the Andes played a major role in sigmodontine diversification, in which the main sigmodontine lineages originated there and later colonized lowlands of South America (Reig 1984, 1986; see also Salazar-Bravo et al. 2013). However, Abrawayaomys, an Atlantic forest endemic, is the sole living representative of one of the main sigmodontine lineages (i.e., those classified at the tribal rank in formal classifications or those genera that do not belong to any recognized tribe). A similar scenario was found for two other Atlantic forest endemics, the genera Delomys and Juliomys (Figures 1 and 2) (Voss 1993; D'Elía et al. 2006a; see also the classification in D'Elía et al. 2007). The other main sigmodontine lineages are distributed outside the Atlantic forest (e.g., Abrotrichini and Sigmodontini) or, even when present in this biome, are not endemic to it (e.g., Akodontini and Oryzomyini). As mentioned above, the Atlantic forest-endemic *Phaenomys* and mostly Atlantic forest-resident Wilfredomys have not been included in any molecular-based phylogenetic analysis. Until their

Table 2 Morphological comparisons among Abrawayaomys and other members of the Sigmodontinae

Character	Abrawayaomys	Rhagomys ^b	Phaenomys ^d	Thomasomys ^f	Aepeomys ^g	Akodon	Reithrodon
Plantar pads	6	6	6	6	6	6	4
Hindfoot surface	Smooth?	Smooth	Squamated	Smooth	Smooth	Squamated	Squamated
Mammae	ба	6	8	6	6	8	8
Spines	Present	Present ^c	Absent	Absent	Absent	Absent	Absent
Relation of tail length (LT) to head-body length (HB)	LT < = > HB	LT≤HB	LT >> HB	LT < = > HB	LT > HB	LT≤HB	LT < < HB
Rostrum	Short	Short	Long	Long	Long, rostral tube developed	Moderate	Moderate
Interorbit	Hourglass-shaped or slightly convergent, with rounded margins	Convergent, with beaded margins	Hourglass-shaped ^e , with beaded margins	Hourglass-shaped, with rounded margins	Hourglass-shaped, with rounded margins	Hourglass-shaped, with rounded margins	Symmetrically constricted, with parallel margins
Palate	Typically short	Long	Short	Short	Short	Short	Long
Mesopterygoid fossa	Not fenestrated	Not fenestrated	Not fenestrated	Not fenestrated	Not fenestrated	Fenestrated	Fenestrated
Alisphenoid strut	Typically present	Present	Absent	Present	?	Typically present	Present
Tegmen tympani	Overlaps squamosal	Overlaps squamosal	Overlaps squamosal	Overlaps squamosal	Overlaps squamosal	Overlaps squamosal	Does not overlag
Carotid circulation	Pattern 1	Pattern 3	Pattern 1	Pattern 1	Pattern 1	Pattern 1	Pattern 3
Capsular process	Present	Present	Absent	Absent	Absent	Typically present	Present
Retromolar fossa	Enlarged	Enlarged	Not enlarged	Not enlarged	Not enlarged	Not enlarged	Not enlarged
Molar design	Intermediate to alternate, crested	Opposite, crested	Opposite, crested	Opposite, crested	Opposite, crested	Intermediate, crested to terraced	Alternate, plane
Anteromedian flexus	Patent	Patent	Patent	Patent	Patent	Patent	Not patent
Mesoloph on M1	Present, small	Present, large	Present, large	Present, large	Present, large	Present, small	Absent
M3 reduction to M2	Much reduced	Moderately	Weakly	Moderately	Moderately to reduced	Much reduced	Weakly
Incisors	Orthodont to proodont	Orthodont	Opisthodont	Opisthodont	Opisthodont	Typically opisthodont	Opisthodont
Incisive foramina	Reaching anterior face M1	Very short	Reaching anterior face M1	Reaching anterior face M1	Reaching anterior face M1	Reaching protocone M1	Reaching protocone M1
Subsquamosal foramen	Present	Present	Absent	Present	Present	Present	Present
Number of ribs	12	13	12	13	13	13	12
Gall bladder	Absent	Absent	Present	Present	Present	Typically present ^h	Present

^aInguinal, abdominal, and postaxial pairs (Pardiñas, unpublished data). ^bBased on *Rhagomys longilingua*, after Luna and Patterson (2003). ^c*Rhagomys rufescens* lacks spines (cf. Luna and Patterson 2003). ^dData from Voss et al. (2002) and Pardiñas (unpublished data). ^eScored as 'convergent' by Pacheco (2003, p. 43). ^fVariation in this genus is remarkable (cf. Pacheco 2003); herein, we follow Voss (1993). ^gBased on *Aepeomys lugens*, after Voss et al. (2002). ^hAkodon montensis lacks gall bladder (cf. Geise et al. 2004).

phylogenetic position is assessed, it is unclear whether they in fact represent additional main sigmodontine lineages almost endemic to the Atlantic forest or simply constitute additional genera belonging to other already identified main lineages of the Sigmodontinae either already known from the Atlantic forest or not. Whatever this result is, the finding that at least three unrelated main sigmodontine lineages, those currently represented by Abrawayaomys, Delomys, and Juliomys, are endemic to the southern Atlantic forest supports early claims (Smith and Patton 1999; D'Elía 2003b; see also Salazar-Bravo et al. 2013) highlighting the role of the Atlantic forest in harboring sigmodontine phylogenetic diversity. Future studies should be designed to test if these lineages originated in the Atlantic forest or simply invaded it after originating elsewhere.

Spines of varying hardness and architectures are present in several rodents (Chernova and Kuznetsov 2001) and are conspicuous in some Neotropical groups, such as porcupines (Erethizontidae) and spiny rats (Echimyidae). The vast majority of sigmodontine rodents have soft fur, but a few genera and species have dorsal spines. These spines are present in both species of Abrawayaomys, both species of Scolomys, all eight species of Neacomys, but only in one of the two species of Rhagomys. Rhagomys longilingua from the Andes has spiny fur, but Rhagomys rufescens from the Atlantic forest has soft fur (Luna and Patterson 2003). Considering the phylogeny portrayed here, we concluded that these spines are the result of evolutionary convergence, and this trait evolved at least four times in sigmodontines (Figure 1). None of these four genera are sister groups, and each one of them shares a more recent common ancestor with soft-furred genera or species (in the case of Rhagomys). The functional significance of spines remains unknown. Unlike porcupine quills, spines of muroid or echimyid rodents are insufficiently rigid to provide much protection against predators, including snakes, birds, and mammals, which are known to prey heavily on many spiny species (Hoey et al. 2004). Patterson and Velazco (2008) suggested a thermoregulatory interpretation based on the geographic distribution of echimyid rodents: the spiniest members occur in tropical lowland forests, while many of the softest-haired members of the family range into high elevations or latitudes, but they also noted several exceptions. The same pattern does not occur in sigmodontine rodents, since soft-furred taxa occur at all latitudes and elevations throughout the Neotropics. Regarding spiny taxa, both Neacomys and Scolomys are lowland forest genera found at lower latitudes (Patton et al. 2000); Abrawayaomys also occurs in lowland forests, but at higher latitudes in the Atlantic forest (Pardiñas et al. 2009b); and the spiny R. longilingua is found in Andean cloud forests at 1,900 ~ 2,100 m in elevation (Luna and Patterson 2003), while the soft-furred R.

rufescens occurs mostly in montane Atlantic forest at $500 \sim 1,000$ m in elevation (Steiner-Souza et al. 2008). Therefore, we still lack robust hypotheses for the ecological role, if any, of spiny fur in sigmodontine rodents, but the character distribution mapped on a phylogenetic tree presented here is the first step toward understanding its evolutionary importance.

Conclusions

Phylogenetic analyses show that *Abrawayaomys* constitutes the single representative so far known of one of the main lineages of the radiation of Sigmodontinae. In addition, it differs from all other Atlantic forest sigmodontine rodents by having a unique combination of morphological character states. Therefore, in formal classifications, it should be regarded as a Sigmodontinae incertae sedis.

Finally, the observed variation of the four Cyt *b* sequences analyzed and their geographic pattern, where the one gathered from the northernmost-collected specimen (in the state of Minas Gerais) was the most divergent, are enticing to further explore variation of a larger sample of *A. ruschii* sequences. Such a study would clarify into which of the already known phylogeographic patterns of Atlantic forest mammals (e.g., Colombi et al. 2010;Ventura et al. 2012; Valdez and D'Elía 2013; see reviews in Martins 2011; Costa and Leite 2012) would *A. ruschii* fit, or if this species presents a so far undescribed pattern. Similarly, such a study would help assess the alpha diversity of *Abrawayaomys* (Pardiñas et al. 2009b). Now, that *A. ruschii* is becoming more frequent in specimen collections, such a study seems feasible.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

KV and GD conceived and coordinated the study. KV, MJJS, LG, YLRL, and YY-Y gathered the DNA sequences. UFJP and YLRL performed the morphological assessment. GD carried out the phylogenetic analyses. GD, KV, and UFJP drafted the manuscript. All authors read and approved the final manuscript.

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