

**Redescription of the enigmatic long-tailed rat *Sigmodontomys aphrastus*
(Cricetidae: Sigmodontinae) with comments on taxonomy and
natural history**

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Abstract.—*Sigmodontomys aphrastus*, the long-tailed rat, is an exceedingly rare rodent species from montane regions of Central and South America of which very little is known ecologically or systematically. It has been variously placed in the genera *Oryzomys*, *Nectomys*, and *Sigmodontomys* based on the five previously known specimens. Two new individuals were collected in northwestern Costa Rica's Cordillera de Tilarán. These new specimens and the other five known specimens are used to redescribe the species, detail measurements of external and cranial morphology, and compare *S. aphrastus* to similarly appearing sympatric species (*Nephelomys albigularis* and *N. devius*) and proposed closely related species (*Sigmodontomys alfari*, *Mindomys hammondi*, and *Melanomys caliginosus*). New ecological data is presented and the general knowledge of its natural history is summarized. The phylogenetic relatedness of *S. aphrastus* with purported sister taxa remains unresolved until combined molecular and morphological analyses are conducted.

Resumen.—*Sigmodontomys aphrastus*, la rata de cola larga, es una especie rara de rodedor de las montañas de America Central y Sur de la cual se conoce muy poco acerca de su ecología y sistemática. Esta especie ha sido clasificada en el género *Oryzomys*, *Nectomys*, y *Sigmodontomys* basado en solo cinco especímenes. Dos nuevos individuos fueron recolectados en el noreste de Costa Rica en la Cordillera de Tilarán. Estos dos nuevos especímenes y junto con los cinco anteriores son usados a describir nuevamente la especie, detallar sus medidas morfométricas externas y craneales, y comparar *S. aphrastus* con especies simpátricas similares (*Nephelomys albigularis* and *N. devius*) y especies que han sido propuestas como dentro del mismo clado (*Sigmodontomys alfari*, *Mindomys hammondi*, y *Melanomys caliginosus*). Presentamos nuevos datos ecológicos y resumimos el conocimiento de su historia natural. La relación filogenética de *S. aphrastus* con las especies propuestas como del mismo clado no sera resuelto hasta que analisis moleculares y morfologicos sean llevados a cabo.

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One of the most poorly known rodents in the Western Hemisphere is the long-tailed rat *Sigmodontomys aphantus* (Harris 1932). Little is known about its natural history, ecology, and systematics, and that is reflected in its complicated taxonomic history. Harris (1932) described this distinctive species as a member of *Oryzomys* Baird, 1857 based on the then single known specimen but made no comparisons to other species. Hershkovitz (1944) transferred it to the genus *Nectomys* Peters, 1861 and suggested that it closely resembled Alfaro's rat, *Nectomys alfari* (= *Sigmodontomys alfari* J. A. Allen, 1897). Later, Hershkovitz (1948) associated it with *Nectomys hammondi* [= *Mindomys hammondi* (Thomas, 1913)]. Hall & Kelson (1959:565) assigned *aphrastus* to their *Oryzomys devius*-group (i.e., *Nephelomys*) but reported that the "systematic position of this 'species' is doubtful." In the second edition of *Mammals of North America*, Hall (1981:618) listed *aphrastus* as a species of *Oryzomys* "provisionally placed between *O. albigularis* and *O. capito*." He reported it as known only from the type locality and apparently considered the taxonomic status uncertain because quotation marks were placed around the word species. [Hall's use of *O. albigularis* corresponds to the species we now recognize as *Nephelomys devius* (Bangs, 1902) and his use of *O. capito* corresponds to the species we now recognize as *Transandinomys talamancae* (J. A. Allen, 1891); see Weksler et al. 2006 for updated nomenclature of oryzomyine genera]. Ray (1962), in an unpublished dissertation on fossil Caribbean oryzomyines, associated *aphrastus* with *Sigmodontomys alfari*. This position was followed by Musser & Carleton (1993, 2005), who provisionally assigned *aphrastus* to the genus *Sigmodontomys*. Recently, Weksler (2006) recovered *S. aphantus* as a member of a clade that also included *Sigmodontomys alfari* and *Melanomys caliginosus* (Tomes,

1860) using cladistic analyses of morphological characters (with and without nuclear sequence data). Although phylogenetic relationships within this clade varied among analytic permutations, *S. aphantus* was recovered as sister species to *M. caliginosus* in the total evidence analysis of morphological and molecular data.

This taxonomic uncertainty is perhaps due to the paucity of specimens, lack of adequate description, and meager ecological information available for the species. *Sigmodontomys aphantus* had been known from only five specimens. Harris (1932) described the species based on a single adult female collected in 1931 by Austin Smith from the Pacific foothills of Costa Rica's Cordillera de Talamanca at 1220 m. The holotype (UMMZ 62875) is represented by a study skin and a broken, inadequately cleaned skull (see Fig. 4). P. Menas took the second specimen, an adult male, in northwestern Ecuador on Volcán Pichincha in 1953. This individual is in the Museum of Comparative Zoology at Harvard University and was first identified as *Sigmodontomys alfari*, but later determined to be *O. aphantus* by Guy G. Musser (pers. comm.). In 1980, Ronald H. Pine and Robert J. Izor caught an adult male and an immature male in the Cordillera de Chiriquí of western Panama (Méndez 1993; R. H. Pine, pers. comm.). The fifth specimen is an adult female caught later in 1980 by Robert S. Voss on the Pacific slopes of Volcán Pichincha in Ecuador (Musser & Williams 1985, Voss 1988).

Our recent fieldwork in the Monteverde Cloud Forest Reserve in northwestern Costa Rica resulted in the collection of two new specimens of *S. aphantus*. Here, we redescribe the species, report external and cranial measurements, and present new ecological data. Additionally, we compare *S. aphantus* with the sympatric and similar appearing *Nephelomys albigularis* and *N. devius*, with taxa it has

been associated with previously, *Sigmodontomys alfari* and *Mindomys hammondi*, and provide comparative photographs of the crania and dentition of these species. Finally, we compare morphological characteristics with the newly proposed sister-taxon, *Melanomys caliginosus* (see Weksler 2006). Reconstruction of phylogenetic relationships is beyond the scope of the current manuscript and will be examined with morphological and molecular analyses in a future manuscript.

Materials and Methods

Our trapping efforts were undertaken over several years in and around the Monteverde Cloud Forest Reserve (Fig. 1 in McCain 2004), a 10,500 h private reserve in Alajuela and Puntarenas provinces in northwestern Costa Rica's Cordillera de Tilarán. At 1250–1300 m in the Peñas Blancas Valley, McCain set 130 traps per night for seven consecutive nights during the dry season (2002) and two late wet seasons at different sites (2000, 2001), and for five consecutive nights during the early wet season (2001) for a total of 3380 trap nights. The trap types (and numbers) used were: standard folding Sherman live traps ($3 \times 3.5 \times 9$ in) (73), large folding Sherman live traps ($3 \times 3.75 \times 12$ in) (40), Victor snap traps (10), and pitfalls (7). At 1500–1550 m along the continental divide, McCain set the same array of traps during the early wet, two late wet (2 sites), and a dry season for a total effort of 3380 trap nights. The same trapping effort was conducted at 1800–1840 m, 1000–1050 m, and 750–800 m. See McCain (2003, 2004) for additional details of trapping procedures. Timm ran pitfall traps along the continental divide at 1550 m from 12–22 Oct 2000, with a total of 22 trap nights with small pitfall traps (6.5 cm diameter) and 10 trap nights with larger pitfall traps (14 cm diameter). Additionally, Timm has trapped extensively on the Pacific



Fig. 1. Verified distribution of *Sigmodontomys aphrastus* in lower Central America and northern South America based on the seven known specimens. The new specimens reported herein from Monteverde in Costa Rica's Cordillera de Tilarán are represented by the star (★); the type locality at San Joaquín de Dota in the Cordillera de Talamanca is represented by the triangle (△); the specimens from the Cordillera de Chiriquí of western Panama and from Volcán Pichincha in northwestern Ecuador are both represented by closed circles (●). See Materials and Methods section for details on localities and museum catalog numbers.

slopes of the Monteverde region over the past two decades with both Sherman live traps and pitfalls at elevations ranging from 1100 m to the continental divide at 1840 m on Cerro Amigos.

We personally examined six of the seven known specimens of *Sigmodontomys aphrastus*. Philip Myers provided us with excellent photographs and with measurements of the type specimen (UMMZ 62875), which we were not able to examine personally. External and cranial comparisons were made between specimens of *S. aphrastus* and those of *S. alfari* ($n = 29$), *Mindomys hammondi* ($n = 2$), *Nephelomys albigularis* ($n = 5$), and *N. devius* ($n = 18$). Nineteen cranial dimensions were measured using dial calipers

graduated to 0.01 mm. Three specimens of *S. aphantus* have damaged skulls, so only eight measurements are used in multivariate analyses (here designated in bold). The majority of cranial measurements used are detailed and illustrated by Musser et al. (1998), which include length of upper molar row (**CLIM1–3**), width of M1 (**BM1**), breadth across bony palate at M1–M1 (**BBP**), length of bony palate (**LBP**), postpalatal length (**PPL**), length and breadth of incisive foramen (**LIF**, **BIF**), length of upper diastema (**LD**), breadth of zygomatic plate (**BZP**), occipitonasal length (**ONL**), breadth of rostrum (**BR**), interorbital breadth (**IB**), zygomatic breadth (**ZB**), and lambdoidal breadth (**LB**). Height of braincase (**HBC**) was measured following Musser (1979); length of rostrum (**LR**) was measured at an angle from the tip of the nasals to the posterior margin of the zygomatic notch. The breadth of the second and third molars (**BM2**, **BM3**) and the breadth across the bony palate across M3–M3 (**BBP3**) were measured also.

We used principal component analysis (PCA) to assess which cranial characteristics are important in explaining the variation among the specimens and to visually display morphometric differentiation between the species. PCA on the correlation matrices of the eight cranial characters was run using SAS (SAS Inc. 1999–2001). Correlation matrices are used because there is no reason to suspect that higher variability within a variable indicated higher overall importance. We used the latent root criterion for determination of meaningful components of variation in the PCA analyses (McGarigal et al. 2000). This criterion applies to PCAs of correlation matrices and assumes that eigenvalues less than one are not considered important, because they account for less variability on average than in any single original variable.

Specimens from the following institutions were used in this study: American Museum of Natural History, New York (AMNH); Field Museum, Chicago (FMNH); Museo Nacional de Costa Rica, San José (MNCR); Museum of Comparative Zoology, Harvard University, Cambridge (MCZ); National Museum of Natural History, Washington, D.C. (USNM); University of Kansas Natural History Museum, Lawrence (KU); University of Michigan Museum of Zoology, Ann Arbor (UMMZ). One specimen of *S. aphantus* obtained at Monteverde (KU 159021) will be deposited in the Museo Nacional de Costa Rica, San José.

Results

As part of our ongoing research on Central American mammals, two individuals of *Sigmodontomys aphantus*, a sigmodontine species previously unknown from northwestern Costa Rica's Cordillera de Tilarán, were captured in the Monteverde Cloud Forest Reserve. Our identification is based on the comparison of external and cranial characters of these two specimens with those of four of the five previously known specimens of *S. aphantus*, with those shown in photographs and measurements of the holotype, and with those of a variety of other oryzomyines. The only previously known specimen from Costa Rica is the holotype, which was collected in the Cordillera de Talamanca at San Joaquín de Dota, San José Province, in 1931. There are now three specimens of *S. aphantus* known from Costa Rica, two from Panama, and two from Ecuador (Fig. 1). These seven specimens of this poorly known species range in age from juvenile to old adult with worn teeth; all specimens were taken at middle elevations, and the species is documented from northwestern Costa Rica's Cordillera de Tilarán to Volcán Pichincha on the western slope of the Andes in northern Ecuador. The eleva-

tional range of these seven specimens is 1220 to 2000–2500 m.

As judged by tooth eruption and wear, the youngest individual (KU 159021) is the juvenile caught in Costa Rica with M3 only partially erupted, and the oldest, with considerable tooth wear, is a specimen from Volcán Pichincha, Ecuador (MCZ 50396). External measurements, based on collectors' notes and tags, and nineteen cranial dimensions for all known specimens of *S. aphrastus*, a representative sample of *S. alfari*, *Nephelomys devius*, and a single *Mindomys hammondi* are shown in Table 1.

Comparing *S. aphrastus*, *S. alfari*, and *N. devius*, the principal components analysis produced eigenvalues ranging between 2.802 and 0.062. According to the latent root criterion, only the first three principal components are significant (McGarigal et al. 2000). Eigenvalues and unit eigenvectors for the first three components of variation are presented in Table 2. The first two principal components are a contrast between the breadth of bony palate, the distance between the first and last molars (BBP1, BBP3), and the length of bony palate (LBP) with the length of the tooth row (CLIM1–3), the size of the last molar (BM3), and the contrasting breadth of the zygomatic plate (BZP). The differential morphology of three species can be distinguished clearly in a plot of the first two principal components (Fig. 2).

Redescription of *Sigmodontomys aphrastus* (Harris, 1932)

The dorsal pelage is long, thick, and soft. In most specimens, it is smooth and shiny but more woolly and fluffy in the adult female from Ecuador and in the young adult from Panama. Ventral fur is sparser and shorter than the dorsal fur, and is paler gray with pronounced ochraceous highlights. Laterally, most specimens have noticeably more orangish highlights than

dorsally. Dorsal-ventral countershading is distinct but not as strong as in many oryzomyines. All hairs, both dorsally and ventrally, have slate gray bases. Most specimens have a rich, dark, and grizzled fur dorsally of deep browns with orange highlights. Some have deeper browns across the rump and head, whereas others have more orangish highlights across the face and shoulders. The young adult from Panama has an overall pale gray pelage with occasional yellowish highlights, but the juvenile from Costa Rica is darker overall than the adults. The adult from Monteverde has white around the mouth and rhinarium, and the old adult male from Volcán Pichincha has many large, conspicuous, white blotches scattered irregularly, both dorsally and ventrally. Pinnae are small (not reaching the eye when laid forward) and sparsely to moderately covered with blackish, dark brown, or reddish-brown hairs both internally and externally. Mystacial and superciliary vibrissae are very long (extend posteriorly beyond caudal margins of pinnae), dark, and numerous. Manual claws are small and unkeeled; several specimens have noticeable deep orange pigmentation at tips. Hind feet are long and robust with sparse to moderate coverage of pale brown to dark brown hairs dorsally and naked ventrally. Feet lack well-developed natatory fringes and interdigital webs. The feet are a pale brown coloration with small, dark squamae visible dorsally. Plantar surface is densely covered with distinct squamae distal to thenar pad (heel is smooth); the hypothenar pad is absent or vestigial (extremely reduced). Interdigital and thenar pads are large and fleshy, with interdigitals 2 and 3 set apart from 1 and 4 as pairs. The second and third interdigital pads are lightly pigmented as compared to the other pads. Digits dI–dIV are long and of nearly equal length, with claw of dI extending just beyond base of phalange 1 of dII and claw of dV extending beyond middle of phalange 2 of dIV. Sparse tufts

Table 1.—External and cranial measurements (mm) of all seven *Sigmodontomys aphrastus* specimens known, in order of tooth wear from least to greatest, with comparisons to average measurements of *Sigmodontomys affari* and *Nephelomys devius* (previously *Oryzomys*). All measurements taken by authors, except for the type specimen, UMMZ 62875 (by Philip Myers).

Measurements	KU 159021	USNM 541201	UMMZ 153808	USNM 541200	UMMZ 62875	KU 161003	MCZ 50396	<i>S. aphrastus</i> Mean (SD)	<i>S. affari</i> Mean (SD, n)	<i>N. devius</i> Mean (SD, n)
Head & body length	84	116	142	140	152	133	—	127.83 (24.58)	161.17 (19.19, 25)	159.47 (12.06, 15)
Tail length	152	176	205	217	235	213	—	199.67 (30.30)	135.40 (18.63, 24)	185.50 (9.21, 16)
Length of hind foot	31	35	37	37	40	37	—	36.17 (3.00)	35.06 (1.74, 25)	36.83 (1.82, 18)
Length of ear	13	17	18	17	—	18	—	16.60 (2.07)	18.70 (2.00, 10)	21.94 (1.95, 18)
Length of upper toothrow	5.54	6.17	5.65	5.78	5.84	6.00	5.79	5.82 (0.210)	5.50 (0.260, 29)	5.85 (0.122, 18)
Breadth of M1	1.88	2.12	1.93	2.12	1.94	2.08	2.10	2.02 (0.103)	1.94 (0.105, 29)	1.70 (0.042, 18)
Breadth of BM2	1.78	1.94	1.88	1.93	2.10	2.04	1.90	1.94 (0.105)	1.80 (0.080, 29)	1.67 (0.040, 18)
Breadth of BM3	1.46	1.60	1.54	1.65	1.76	1.57	1.62	1.60 (0.094)	1.33 (0.885, 28)	1.47 (0.050, 18)
Breadth across M1–M1	5.66	6.68	6.80	7.12	6.71	7.10	7.26	6.76 (0.535)	7.13 (0.538, 28)	7.37 (0.346, 18)
Breadth across M3–M3	5.86	6.29	7.13	7.24	6.83	7.05	7.11	6.79 (0.517)	6.29 (0.428, 24)	7.04 (0.338, 18)
Length of bony palate	5.52	7.10	6.49	7.19	6.47	7.42	7.09	6.90 (0.687)	7.86 (0.574, 28)	7.95 (0.300, 18)
Postpalatal length	7.45	8.81	10.37	9.86	—	10.45	10.32	9.59 (1.241)	10.59 (1.170, 26)	12.85 (0.704, 18)
Length of incisive foramen	3.75	4.02	—	4.82	5.34	5.32	4.98	4.70 (0.671)	4.68 (0.464, 28)	5.56 (0.458, 18)
Breadth of incisive foramen	1.51	1.65	—	2.08	2.93	2.16	1.91	2.04 (0.502)	1.83 (0.155, 28)	2.87 (0.264, 18)
Length of upper diastema	5.67	7.04	—	8.68	—	8.94	8.09	7.68 (1.342)	8.19 (0.802, 28)	9.75 (0.613, 18)
Breadth zygomatic plate	2.46	2.52	3.08	2.84	3.24	3.23	2.67	2.86 (0.327)	3.86 (0.647, 29)	3.62 (0.217, 18)
Occipitonasal length	26.05	29.33	—	32.65	—	32.85	31.67	30.51 (2.859)	33.91 (2.477, 27)	37.19 (1.338, 18)
Height of braincase	8.21	9.82	9.59	10.29	—	9.88	—	9.56 (0.795)	10.09 (0.607, 26)	10.46 (0.471, 17)
Breadth of rostrum	5.38	5.78	—	6.61	6.83	7.11	6.31	6.34 (0.654)	7.00 (0.723, 26)	7.32 (0.456, 18)
Interorbital breadth	4.83	5.38	—	5.11	5.50	5.50	4.50	5.14 (0.406)	6.24 (0.490, 28)	5.31 (0.260, 18)
Zygomatic breadth	14.44	16.33	—	18.58	—	18.63	—	17.00 (2.013)	18.18 (1.200, 25)	19.22 (0.832, 18)
Lambdoidal breadth	10.87	11.69	12.56	13.34	—	13.06	—	12.30 (1.018)	12.66 (0.639, 27)	13.51 (0.406, 18)
Length of rostrum	7.30	8.46	—	9.72	—	10.14	8.97	8.90 (1.087)	10.13 (0.927, 27)	12.37 (0.547, 18)

Table 2.—Eigenvectors and eigenvalues for the first three axes of the principal components analysis and canonical coefficients and eigenvalues for the two canonical axes of *Sigmodontomys aphrastus*, *S. alfari*, and *Nephelomys devius* using eight cranial variables (see Materials and Methods for abbreviations).

Measurement	PC1	PC2	PC3	CA1	CA2
CLIM1-3	0.283243	0.485201	0.002927		
BM1	-0.206149	-0.069189	0.660632	2.47016	0.23220
BM2	-0.114680	0.127876	0.686633		
BM3	0.206730	0.566823	0.154536	-0.74303	1.64358
BBP1	0.515285	-0.185868	0.168765	-0.86774	0.09020
BBP3	0.498870	0.247484	0.032380		
LBP	0.460968	-0.279109	0.043562	0.17556	-1.19733
BZP	0.307419	-0.498453	0.191787		
Eigenvalue	2.8017	2.2820	1.8477	6.8494	3.4426

of rather short (almost as long as the claw), brown unguis hairs at bases of claws on dII-dV may be present or absent. Tail is much longer than head and body (1.5 × HBL or more) and covered sparsely with short, stout, black or brown hairs with conspicuous hexagonal epidermal scales; reduced terminal tuft of hairs present. The tail is unicolored, uniformly brown, except for patches of slightly paler skin ventrally at base and white tips on the adult and juvenile from Monteverde. Mammary complement consists of eight teats in inguinal, abdominal, postaxial, and pecto-

ral pairs. For a summary of external measurements see Table 1.

Skull has a short, stout rostrum flanked by very shallow zygomatic notches and a slightly inflated braincase with well-developed temporal crests (Figs. 3A, B, 4). Lambdoidal and nuchal crests are developed in older adults. Interorbital region is anteriorly convergent with strongly beaded supraorbital margins. Posterior margin of zygomatic plate is anterior to M1 alveolus, and anterior margin is straight, without an anterodorsal spinous process. Zygomatic arches converge anteriorly and are wider at the squamosal root. Jugal present and large with the maxillary and squamosal zygomatic processes widely separated and not overlapping in lateral view. Nasal bones have acutely angled posterior margins that extend posteriorly behind lacrimals. Lacrimals usually have longer maxillary than frontal sutures. Posterior wall of the orbit is smooth. Frontosquamosal suture is anterior to frontoparietal suture (dorsal facet of frontal in broad contact with squamosal). Parietals have broad lateral expansions. The interparietal bone is large, truncated anteriorly, and broadly triangular posteriorly with lateral wings. Basicranial flexion is weakly pronounced and foramen magnum orientation is mostly caudad. Incisive foramina are narrow and short, not extending posteriorly to level of M1 alveoli, widest at

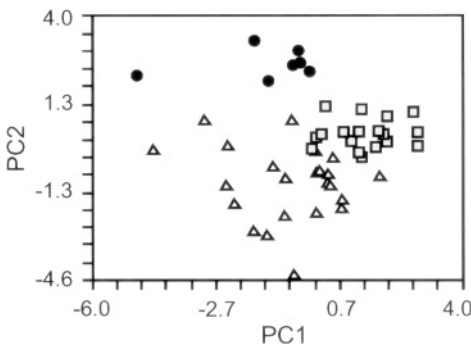


Fig. 2. Plot of specimen scores on the first two axes of the principal components analysis (PCA) for all known specimens of *Sigmodontomys aphrastus* (●), 23 *S. alfari* (■), and 18 *Nephelomys devius* (△) based on 8 cranial and dental characters. The specimens of *S. aphrastus* are arrayed from left to right mainly by age, as the youngest specimen (KU 159021) is on the far left and the adults are grouped on the right.

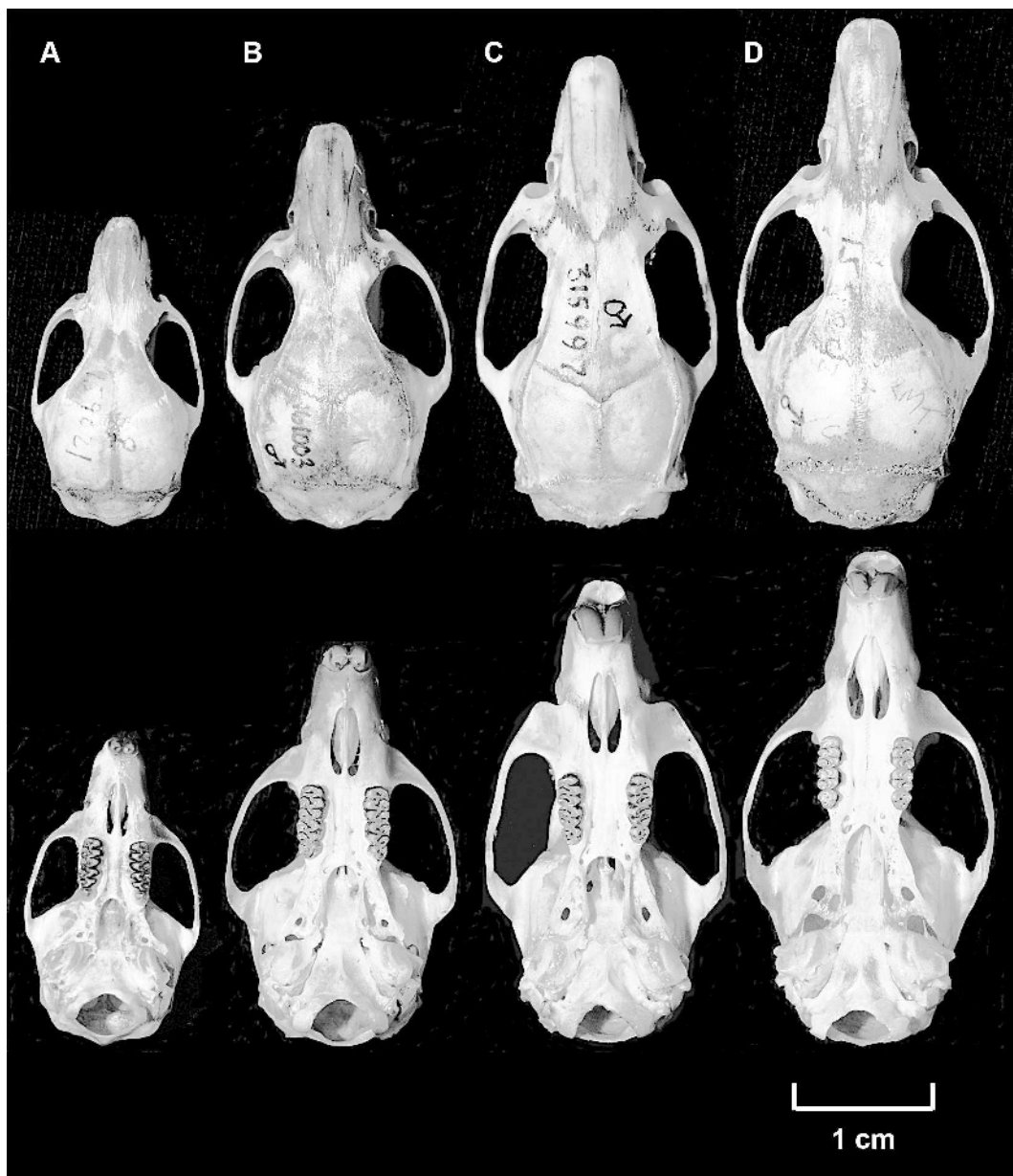


Fig. 3. Dorsal and ventral cranial views (ca. $1.75\times$) of the two new specimens of *Sigmodontomys aphrastus* from the Monteverde Cloud Forest Reserve, Costa Rica: (a) the juvenile female (KU 159021); (b) the adult male (KU 161003); (c) a specimen of *S. alfari* from Almirante, Panama (USNM 315997); and (d) a specimen of *Nephelomys devius* from the Monteverde Cloud Forest Reserve, Costa Rica (KU 159002).

midlength, and taper symmetrically anteriorly and posteriorly. Auditory bullae are of an intermediate size relative to the exposed petiotic bone. Bony palate between the molar rows is smooth or weakly sculpted, short, and narrow. Posterolat-

eral palatal pits are simple and small, usually unrecessed in fossae. Mesopterygoid fossa penetrates anteriorly between maxillae; bony roof of mesopterygoid fossa is usually perforated by narrow sphenopalatine vacuities (the juvenile

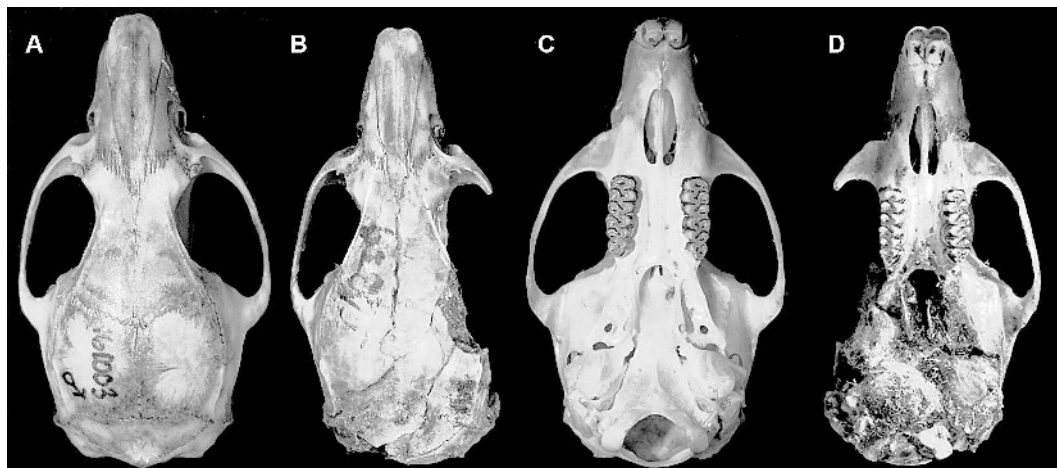


Fig. 4. Dorsal and ventral cranial views (ca. 1.75 \times) of the adult specimen of *Sigmodontomys aphrastus* from the Monteverde Cloud Forest Reserve, Costa Rica with the holotype from San Joaquín de Dota, Costa Rica: (a) dorsal view of new adult male (KU 161003); (b) dorsal view of adult female holotype (UMMZ 62875); (c) ventral view of new adult male (KU 161003); and (d) ventral view of adult female holotype (UMMZ 62875).

from Monteverde has a completely ossified mesopterygoid roof). Alisphenoid strut is absent (buccinator-masticatory foramen and accessory foramen ovale are confluent), and alisphenoid canal has a large anterior opening. Stapedial foramen and posterior opening of alisphenoid canal is small, squamosal-alisphenoid groove and sphenofrontal foramen are absent, and secondary anastomosis of internal carotid crosses dorsal surface of pterygoid plate (carotid circulatory pattern 3 of Voss 1988). Posterior suspensory process of the squamosal is absent. Postglenoid foramen is large and rounded. Subsquamosal fenestra is vestigial or absent. Periotic is exposed posteromedially between ectotympanic and basioccipital but usually not extending anteriorly to carotid canal. Mastoid is unfenestrated. Capsular process of lower incisor alveolus is absent or reduced. Superior and inferior masseteric ridges are conjoined anteriorly as single crest below m1. The coronoid processes are long, thin, pointed, and posteriorly angled dorsally, reaching the level of the rounded condyloid processes.

Upper incisors are ungrooved, opisthodont, and covered with orange enamel anteriorly with smoothly rounded enamel bands. Maxillary toothrows are mostly parallel with a slight divergence posteriorly and molars are large, complex, and bunodont (Fig. 5). Labial and lingual flexi of M1 and M2 are deeply interpenetrating; labial flexi are convoluted (wrinkled) and enclosed by a cingulum. First upper molar (M1) anterocone not divided into labial and lingual conules (anteromedian flexus absent). Anteroloph is well developed and fused with the anterostyle on labial cingulum and fused with anterocone (anteroflexus reduced or absent). Protostyle is absent. Mesolophs are present on all upper molars. Paracone is usually connected by an enamel bridge to posterior moiety of protocone, and median mure is connected to the protocone. Second upper molar (M2) protoflexus is absent, mesoflexus present as single internal fossette (a second very small labial fossette is present in the adult from Panama), and paracone has no accessory loph. Third upper molar (M3) has a posteroloph and diminutive hypo-

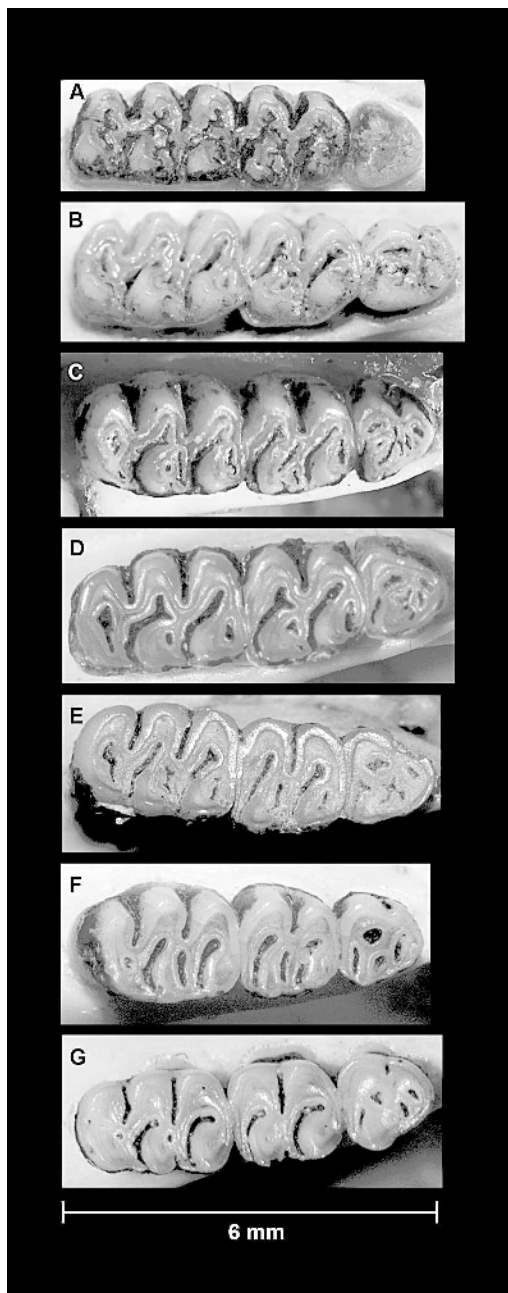


Fig. 5. Molar tooth rows (upper right) of five *Sigmodontomys aphrastus*: (a) the juvenile female from the Monteverde Cloud Forest Reserve, Costa Rica (KU 159021), (b) the immature male from Chiriquí, Panama (USNM 541201), (c) the holotype—an adult female from San Joaquín de Dota, Costa Rica (UMMZ 62875), (d) the adult male from the Monteverde Cloud Forest Reserve, Costa Rica (KU 161003), and (e) the adult male from Guarumos, Ecuador (MCZ 50396); (f) a specimen of

flexus (the latter tending to disappear with moderate to heavy wear). Accessory labial root of M1 is present. First lower molar (m1) anteroconid has no anteromedian flexid (Fig. 6). Both the anterolabial cingulum and a small anterolophid are present on all lower molars (the latter disappear with moderate to heavy wear). Ectolophid is absent on m1 and m2. Mesolophid is present and distinct on m1 and m2. Posteroflexid present on m3. Accessory lingual and labial roots are present on m1. For cranial measurements see Table 1.

Stomach is unilocular-hemiglandular with glandular epithelium extending into corpus. Gall bladder is absent.

Discussion

Comparisons with other taxa

Sigmodontomys aphrastus was consistently recovered as a member of a clade composed of *Melanomys caliginosus* and *Sigmodontomys alfari* in the phylogenetic analyses of Weksler (2006); comparisons with these taxa are provided here. In addition, we compare it with *Mindomys hammondi* and *Nephelomys devius* (see Weksler et al. 2006 for new classification of oryzomyines). Although these taxa are not closely related to *S. aphrastus* (Weksler 2003, 2006), they occur at the same localities where *S. aphrastus* was collected, are externally similar to it, and have been considered to be taxonomically close to *S. aphrastus* historically (Hershkovitz 1948, Hall & Kelson 1959). Table 3 summarizes key morphological comparisons among these taxa.

Sigmodontomys aphrastus differs from *S. alfari* by its considerably smaller body

←

S. alfari from Almirante, Panama (USNM 315997); and (g) a specimen of *Nephelomys devius* from the Monteverde Cloud Forest Reserve, Costa Rica (KU 159002).

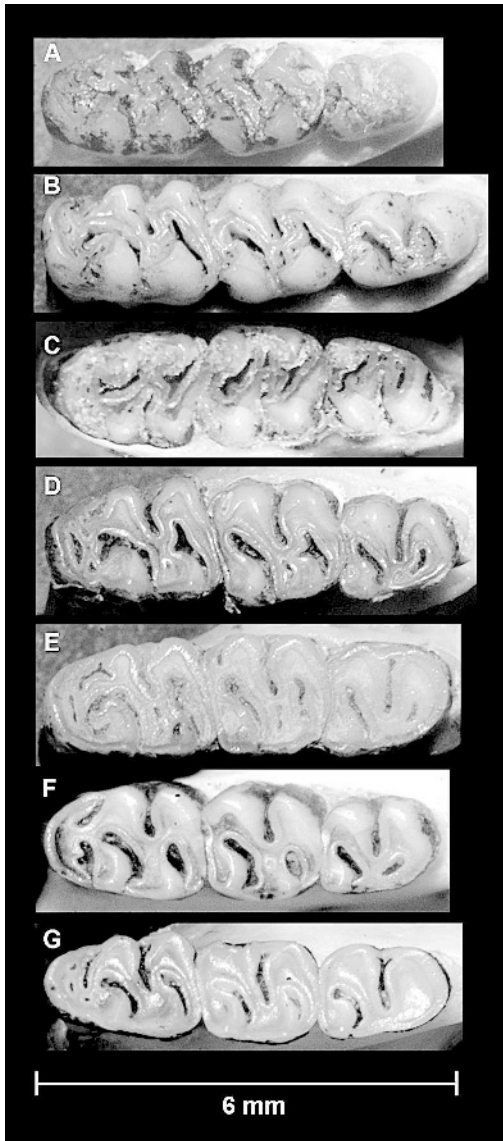


Fig. 6. Molar tooth rows (lower right*) of five *Sigmodontomys aphrastus*: (a) the juvenile female from the Monteverde Cloud Forest Reserve, Costa Rica (KU 159021), (b) the immature male from Chiriquí, Panama (USNM 541201), (c) the holotype—an adult female from San Joaquín de Dota, Costa Rica (UMMZ 62875; *lower left), (d) the adult male from the Monteverde Cloud Forest Reserve, Costa Rica (KU 161003), and (e) the adult male from Guarumos, Ecuador (MCZ 50396); (f) a specimen of *S. alfari* from Almirante, Panama (USNM 315997); and (g) a specimen of *Nephelomys devius* from the Monteverde Cloud Forest Reserve, Costa Rica (KU 159002).

size (adults average 33.2 mm smaller) and much longer tail (average 64.3 mm longer) with a small terminal tuft (Table 1). The tail of *S. alfari* is slightly longer than or subequal to head-and-body length and lacks the terminal tuft. *S. aphrastus* has much longer vibrissae; in *S. alfari* the vibrissae do not extend posteriorly behind the pinnae as in *S. aphrastus*. *Sigmodontomys aphrastus* lacks the interdigital webs on hind feet that are present in *S. alfari*. The second and third digital pads are lightly pigmented in *S. aphrastus*, whereas they are heavily pigmented in *S. alfari*. The hexagonal tail scales of *S. aphrastus* are smaller, thinner, and not as visible to the naked eye as in *S. alfari*. Additionally, the pelage of *S. aphrastus* is somewhat longer, softer, and has fewer orangish or yellowish highlights than does that of *S. alfari*. *Sigmodontomys aphrastus* and *S. alfari* both have large, complex molars with four molar roots in M1 and three in m1, small auditory bullae, defined post-orbital ridge, subsquamosal fenestra, derived carotid arterial circulation, and well-developed supraorbital crests. *Sigmodontomys aphrastus* has a less robust and a broader skull with a more inflated braincase than does *S. alfari*. The zygomatic arches in *S. alfari* are more squared and with more prominent supraorbital beading and temporal ridges than in *S. aphrastus*. Nasal bones in *S. aphrastus* have acutely angled posterior margins, while nasals of *S. alfari* have blunt posterior margins. The zygomatic plate is broader and forms a deeper notch in *S. alfari*. *Sigmodontomys aphrastus* has a well developed jugal, which is reduced or absent in *S. alfari*. In *S. aphrastus* the bony palate between the tooth rows is narrower and shorter, extending to the posterior borders of M3s only in the oldest individuals and M3s are only slightly smaller than the M2s; whereas in *S. alfari* the palatal bone extends well beyond the M3s by almost a molar length and the M3s are considerably smaller

Table 3.—Morphological characteristics distinguishing *Sigmodontomys aphrastus* from *Sigmodontomys alfari* and three closely related or sympatric species, including *Melanomys caliginosus*, *Mindomys hammondi*, and *Nephelomys devius*.

Morphological Characteristic	<i>Sigmodontomys aphrastus</i>	<i>Sigmodontomys alfari</i>	<i>Melanomys caliginosus</i>	<i>Mindomys hammondi</i>	<i>Nephelomys devius</i>
Superciliary vibrissae ¹ :	Long	Short	Short	Long	Short
Hypothenar pad:	Absent or vestigial	Absent or vestigial	Distinct	Distinct	Distinct
Squamae on plantar surface of hind feet:	Distinct	Distinct	Distinct	Sparse, indistinct	Smooth
Interdigital webbing:	Sparse & short	Sparse & short	Sparse & short	Sparse & short	Dense & long
Tail:	Absent	Present	Absent	Absent	Absent
	Much longer than HBL ² , unicolorous, small terminal tuft	Subequal to HBL, unicolorous, no terminal tuft	Shorter than HBL, unicolorous, no terminal tuft	Much longer than HBL, unicolorous, no terminal tuft	Longer than HBL, bicolorous, no terminal tuft
Nasal posterior terminus:	Acutely pointed	Blunt	Blunt	Blunt	Blunt
Interorbital shape:	Cuneate & beaded	Cuneate & beaded	Cuneate & beaded	Cuneate & beaded	Hourglass & smooth
Zygomatic notch:	Shallow	Deep	Deep	Shallow	Deep
Jugal:	Developed	Reduced or absent	Reduced	Developed	Reduced or absent
Carotid circulation ³ :	Pattern 3	Pattern 3	Pattern 3	Pattern 1	Pattern 1
Subsquamosal fenestra:	Absent or vestigial	Absent or vestigial	Absent or vestigial	Absent or vestigial	Present
M1/m1 accessory roots:	Present	Present	Present	Absent	Only labial on m1
Molar flexi penetration:	Deep	Shallow	No penetration	Deep	Deep
M1 anterocone:	Undivided	Undivided	Undivided	Undivided	Divided
Connection of paracone & protocone:	Posterior moiety	Posterior moiety	Median moiety	Posterior moiety	Posterior moiety
Fossettes in M2 mesoloph:	1 internal	1 internal	1 internal	2 internal	1 internal
M3 hypoflexus:	Reduced	Reduced	Reduced	Developed	Reduced
m1 ectolophid:	Absent	Absent	Absent	Present	Present
m2 anterolabial cingulum:	Present	Absent	Present	Present	Present
Gastric glandular epithelium:	Extending to corpus	<i>Condition unknown</i>	Limited to antrum	Limited to antrum	Limited to antrum

¹ Long = extending posteriorly behind pinnae; short = not extending posteriorly behind pinnae.
² HBL = combined head and body length.
³ Pattern 1 = presence of stapedial and sphenofrontal foramina and the squamosal-alisphenoid groove; pattern 3 = secondary anastomosis of internal carotid crosses dorsal surface of pterygoid plate (Voss 1988).

than the M2s. *Sigmodontomys alfari* also has more complex palatal pit morphology than does *S. aphrastus*. Additionally, the capsular processes are considerably smaller in *S. aphrastus*; they form broad shelves in *S. alfari*. *Sigmodontomys aphrastus* has more lophodont upper molars (the labial and lingual flexi do not interpenetrate deeply on the upper molars of *S. alfari*) and has an anterolabial cingulum on m2 that is absent in *S. alfari*.

Sigmodontomys aphrastus differs from *Melanomys caliginosus* by its larger body size (adults average 6.3 mm larger) and much longer tail (average 65.7 mm longer) with small terminal tuft (lengths of *Melanomys* from Reid 1997). The tail of *Melanomys* is shorter than the length of head and body, and lacks a terminal tuft. They have distinctly different fur coloration: *S. aphrastus* has brownish pelage with orange highlights, whereas the fur in *Melanomys* is uniformly dark, almost black. *Sigmodontomys aphrastus* has much longer vibrissae; the vibrissae do not extend posteriorly behind the pinnae in *Melanomys*. *Melanomys* has a distinct hypothenar pad on the hind foot, which is absent or vestigial in *S. aphrastus*. The nasal bones of *S. aphrastus* have acutely angled posterior margins, whereas *Melanomys* has blunt posterior margins. In *Melanomys*, the zygomatic plate is broader and forms a deeper notch. *Sigmodontomys aphrastus* has a developed jugal, which is reduced or absent in *Melanomys*. *Sigmodontomys aphrastus* also has more lophodont upper molars (the labial and lingual flexi do not interpenetrate deeply on the upper molars of *Melanomys*). The paracone and protocone are connected by median moiety in *Melanomys* but have a posterior connection in *S. aphrastus*.

Sigmodontomys aphrastus differs from *Mindomys hammondi* by its smaller body size (the full-grown adult male *S. aphrastus* from Monteverde is about the same size of a young *M. hammondi*) and has

a proportionally longer tail with small terminal tuft. The tail of *Mindomys* is slightly longer than the head and body length, lacks terminal tuft, and has rectangular scales, not hexagonal ones as in *S. aphrastus*. *Mindomys* has shorter and woollier fur with paler brown coloration with a faint orange tint across the spine and less ochraceous highlights laterally than *S. aphrastus*. *Mindomys* has a distinct hypothenar pad on the hind foot, which is absent or vestigial in *S. aphrastus*. *Sigmodontomys aphrastus* has distinct plantar squamae, whereas the sole of the hind foot is sparsely covered with indistinct squamae in *Mindomys*. The cranium of *M. hammondi* is considerably larger and more robust, with similarly proportioned molars and palatal bones as compared to *S. aphrastus*. The braincase is smooth, more inflated, and has less of an interorbital constriction than does *S. aphrastus*. The nasal bones of *S. aphrastus* have acutely angled posterior margins, whereas *Mindomys* has blunt posterior margins. Convex-shaped incisive foramina occur in *S. aphrastus*, whereas the incisive foramina are tear-drop shaped in *Mindomys*. *Sigmodontomys aphrastus* has the derived carotid circulation pattern, and *Mindomys* has the stapedial and sphenofrontal foramina and the squamosal-alisphenoid groove. *Sigmodontomys aphrastus* retains accessory roots on M1/m1, which are normally absent on these teeth in *Mindomys* and has two internal fossettes at the mesoloph of M2 (only one fossette is present on the M2 of *Mindomys*). *Mindomys* has a developed hypoflexus on the M3, which is reduced in *S. aphrastus*, and an ectolophid on m1, which is absent in *S. aphrastus*.

Sigmodontomys aphrastus externally resembles in dorsal fur coloration, general size, and facial features species of *Nephelomys*, such as *N. albigularis* and *N. devius*, common rodents in mid and high elevations in the Andean and lower Central American cordilleras. However, *S. aph-*

rastus differs from *Nephelomys* in several traits, the following are the most noteworthy. *Sigmodontomys aphantus* has a much longer unicolorous tail with small terminal tuft, whereas the tail of *Nephelomys* is bicolored, slightly longer than head-and-body length, and lacks a terminal tuft. *Sigmodontomys aphantus* has longer fur that is occasionally more lax than in *Nephelomys*, smaller ears, and much longer superciliary vibrissae (the vibrissae do not extend posteriorly behind the pinnae in *Nephelomys*). *Nephelomys* retains a distinct hypothenar pad on the hind foot, which is absent or vestigial in *S. aphantus*. *Sigmodontomys aphantus* has much wider hind feet and distinct plantar squamae, whereas the sole of the hind foot is entirely smooth in *Nephelomys*. Only sparse tufts of short ungual hairs are present on pedal digits II–V of *S. aphantus* but are denser and longer in *Nephelomys*. In comparison with *S. aphantus*, *N. albigularis* and *N. devius* have a much longer, narrower skull with an elongated rostrum, greater constriction at the interorbitals, a larger interparietal, distinctly smaller and less complex, rectangular molars. Nasal bones of *S. aphantus* have acutely angled posterior margins, whereas *Nephelomys* has rounded or squared posterior margins. Interorbital regions are cuneate with beaded margins in *S. aphantus*, but the interorbital region is hourglass-shaped with smooth supraorbital margins in *Nephelomys*. *Sigmodontomys aphantus* has a shallower zygomatic notch, whereas the zygomatic plate is broader and forms a deeper notch in *Nephelomys*. *Sigmodontomys aphantus* retains a developed jugal, which is reduced or absent in *Nephelomys*, and displays a derived carotid circulation pattern absent in *Nephelomys*. The subsquamosal fenestra is always present and developed in *Nephelomys* but absent or extremely reduced in *S. aphantus*. *Sigmodontomys aphantus* retains accessory roots on M1 (normally

absent in *Nephelomys*) but lacks the M1 anteromedian flexus and the ectolophid on m1 (both present in *Nephelomys*). Of the cranial measurements taken, all measurements are larger for *N. devius* than for *S. aphantus*, except breadth of the molars (Table 1).

Taxonomic comments.—Cadena et al. (1998:11) assigned a recently collected specimen (ICN 13663) to the genus *Sigmodontomys*, stating it “...probably represents an undescribed species of the genus.” The specimen was collected in cloud forest at 1300 m in the Colombian Chocó of western Nariño Department. They characterized the single adult male as having an extremely long tail; glossy, buffy, brown dorsal pelage; long, prominent vibrissae; cuneate interorbital with weakly developed supraorbital ridges; and oval (rather than rectangular) molars. Although these traits are shared by *Sigmodontomys aphantus*, the undescribed species has traits that are clearly at odds with the two currently recognized species of *Sigmodontomys*: narrow hind feet, short palate without posterolateral palatal pits, and a well-developed alisphenoid strut. In fact, the latter two characters are in contrast with 2 putative oryzomyine synapomorphies (Voss and Carleton 1993): absence of alisphenoid and presence of long palate with prominent posterolateral palatal pits. Although some oryzomyines possess one or the other trait, there is no oryzomyine with both alisphenoid strut and short palate (Weksler 2006). Both states are commonly observed among thomomyine rodents (Pacheco 2003). Further examination of this Chocó exemplar is necessary for its taxonomic clarification.

The phylogenetic relationships between *Sigmodontomys aphantus*, *S. alfari*, and *Melanomys* varied among the analytic permutations of Weksler (2006). *Sigmodontomys aphantus* was recovered as sister species to *S. alfari* in the analysis of morphology-only matrix when coding

polymorphic character-states as intermediate steps between fixed character-states (Weksler 2006, Fig. 35). In the total evidence analysis of morphological and molecular data, *Melanomys caliginosus* and *S. aphrastus* are recovered as sister species, although molecular data was not included for *S. aphrastus*. Additional data are necessary for the resolution of this unresolved area of oryzomyine phylogeny, particularly the inclusion of molecular data for *S. aphrastus* which are now available from the adult specimen recently collected at Monteverde (Weksler et al. in press).

Ecology of *Sigmodontomys aphrastus*

Habitat.—Both specimens from Monteverde were taken in ground sets. The juvenile female was captured along the continental divide at 1550 m and the adult male was captured at 1250 m on the Caribbean slope of Cordillera de Tilarán in northwestern Costa Rica in 2000 and 2002, respectively. Both specimens were taken in intact, mature forests that are classified as lower montane rain forest and premontane rain forest, respectively (Haber 2000). The lower montane rain forest along the continental divide at 1550 m is dominated by cloud forest vegetation with a variable and broken canopy at about 5–15 m with pronounced epiphyte and moss growth and a moderately dense ground cover of mosses, ferns, herbs, and shrubby vegetation. This forest experiences substantial rainfall during the wet season (May–December) and heavy misting for most of the year. The forest along the continental divide where the juvenile female was captured is within a few meters of the elfin forest (3–5 m canopy) on an exposed ridge molded by the high winds and driven precipitation coming from the Caribbean.

A lush evergreen forest characterizes premontane rain forest vegetation (700–1400 m at Monteverde), where canopy height and continuity decrease with ele-

vation, and epiphyte and moss loads increase with elevation. The forest where the adult male was caught had a tall, somewhat broken canopy (15–30 m) with heavy epiphyte and moss growth. There was a multi-layered understory of ferns, shrubs, herbs, young trees, and monocots. Rainfall is still considerable during the rainy season, although the year-round dense mists and wind-driven rain are not as pervasive as at the continental divide. The habitat directly around the trap where the adult male was captured was a large, decomposing (~1 m diameter) fallen tree trunk with several young trees growing out of it. Overhead the canopy was closed and the ground cover was sparse with occasional plants but a deep litter layer around the trap. Within a few meters was very dense ground cover of low vegetation. For a broad overview of the plant communities in the vegetation zones at Monteverde, see Haber (2000), and for more detailed climatic data see Clark et al. (2000).

The holotype was caught at San Joaquín de Dota, Costa Rica, in undisturbed premontane rain forest on the Pacific slope of the Cordillera de Talamanca at 1220 m. Both Ecuadorian specimens were from apparently undisturbed forest on the Pacific slope of Volcán Pichincha, the first from upper montane rain forest between 2000–2500 m near Guarumos, and the second from lower montane rain forest near Mindo (Voss 1988). The two Panamanian specimens were trapped in disturbed, fairly open, relatively dry habitat, which included mixed grass, weeds, brush, oaks, and other trees (R. H. Pine, pers. comm.). No *S. aphrastus* have been captured in disturbed habitats in Monteverde, despite Timm's extensive trapping efforts in a wide variety of disturbed habitats within and near the community over two decades, and the species has not been detected anywhere along the Pacific slopes of the Monteverde region where

we have trapped extensively. All known specimens of *S. aphantus*, except the two specimens from Panama, were taken in intact forest.

Community composition.—Along the continental divide at 1500–1550 m where the juvenile female *S. aphantus* was trapped, we also obtained the following terrestrial or scansorial species: *Cryptotis nigrescens*, *Heteromys nubicolens*, *Nephelomys devius*, *Nyctomys sumichrasti*, *Peromyscus nudipes*, *Reithrodontomys gracilis*, *Reithrodontomys* sp., and *Scotinomys teguina*. *Peromyscus nudipes* dominated numerically (51% of individuals captured), followed by *Heteromys nubicolens* (24%), *Scotinomys teguina* (15%), and *Nephelomys devius* (9%); all other species were relatively rare. At 1250–1300 m on the Caribbean slope where the adult male was obtained, we captured *Marmosa mexicana*, *Cryptotis nigrescens*, *Heteromys nubicolens*, *Nephelomys devius*, *Nyctomys sumichrasti*, *Peromyscus nudipes*, *Reithrodontomys gracilis*, and *Scotinomys teguina*. Again, *Peromyscus nudipes* dominated numerically (50% of individuals captured), followed by *Heteromys nubicolens* (21%), *Nephelomys devius* (17%), and *Scotinomys teguina* (10%); all other species were relatively rare. Two *Heteromys nubicolens* were caught in the same trap (baited with cracked corn, mixed seeds, and vanilla) as the adult male *S. aphantus* over the seven days the trap was out. For additional information on other mammals that occur in the Reserve and general ecology of species in this area see Timm & LaVal (2000) and McCain (2003, 2004, 2006).

At San Joaquín de Dota, Costa Rica, the following species are known to occur, based on specimens in the University of Michigan collection obtained by Austin Smith when he acquired the holotype of *S. aphantus*—*Marmosa mexicana*, *Microsciurus alfari*, *Sciurus granatensis*, *Oligoryzomys vegetus*, *Peromyscus nudipes*, *Reithrodontomys mexicanus*, and *Tylomys*

watsoni (Goodwin 1946, McPherson 1985, Carleton & Musser 1995). In Panama, R. H. Pine and R. J. Izor obtained the following species of terrestrial mammals in the vicinity of where they captured the two specimens of *S. aphantus* in 1980: *Marmosa robinsoni*, *Oligoryzomys fulvescens*, *Peromyscus nudipes*, *Reithrodontomys mexicanus*, and *Sigmodon hirsutus* (R. H. Pine, pers. comm.). And in Ecuador at Mindo, R. S. Voss collected or observed *Chironectes minimus*, *Didelphis* sp., *Marmosa* sp., and the rodents, *Mindomys hammondi*, *Handleyomys alfari*, and *Nephelomys moerex* along with his one specimen of *S. aphantus* (Voss 1988).

To date, no specimens of *S. aphantus* and *S. alfari* have both been taken from the same locality or similar elevations; the two species appear to be elevationally separated, with *S. aphantus* occurring above 1200 m, and *S. alfari* occurring at and/or below 900 m (Reid 1997, Musser et al. 1998).

Natural history.—Of the seven known specimens of *Sigmodontomys aphantus*, three are females and four are males. All were adults, except for the juvenile female caught by the authors in Monteverde and a young adult male from Panama. The adult male caught at Monteverde had descended testes measuring 7×5 mm, and the adult male from Panama had testes measuring 5 mm in length. Mammary morphology consists of eight teats in inguinal, abdominal, postaxial, and pectoral pairs; other reproductive specifics remain unknown. We collected one species of tick (*Ixodes sinaloa*—2 nymphs, 5 larvae) from the adult male *S. aphantus* from Monteverde; these are the first parasites reported for this species. Muscle tissue and all major organs of the adult male were preserved in ethanol for future study. The juvenile was captured in a pitfall trap (14 cm diameter) and the adult male was taken in a standard Sherman live trap on the ground. Arrays of

traps at these sites and at other Caribbean slope sites between 750–1840 m were placed both on the ground and in the trees (1–3 m above ground; McCain 2004). To date, no *S. aphantus* have been caught arboreally. In McCain's (2004) trapping efforts, half of the Sherman live traps were baited with peanut butter and oats, and the other half were baited with a mixture of grain sorghum, millet, sunflower seeds, and vanilla extract. The adult male *S. aphantus* was caught in a Sherman with the later bait type. Victor traps were baited with peanut butter and oats. Pitfall traps were not baited by McCain or Timm. The stomach and stomach contents were preserved, but no diet analysis was performed. Given our extensive trapping efforts using various sizes and types of traps in the Monteverde area over the last 20+ years, it is surprising that only two *S. aphantus* have been detected. Judging from our trapping efforts, as well as those of others, *S. aphantus* occurs only on the Caribbean slopes there and either occurs in very low densities or is reluctant to enter traps. It is plausible that the detection of so few individuals in the Cordillera de Tilarán may be the result of limited trapping efforts between 1100–1500 m on the Caribbean slopes. Before the concerted trapping effort of McCain (2003, 2004, 2006) from 850–1840 m, minimal trapping was conducted there, mostly around shelters at 840 and 950 m.

The Monteverde Cloud Forest Reserve is a region of considerable endemism and also the northernmost locality known for *Cryptotis nigrescens*, *Nephelomys devius*, and *Reithrodontomys creper* (Timm & LaVal 2000) and the southernmost locality for a new species of *Heteromys* (*H. nubicolens* Anderson & Timm 2006) as well as for the shrew *Cryptotis merriami* (Woodman 2000). *Sigmodontomys aphantus* is classified as species with "reduced populations" by Costa Rica's Ministerio del Ambiente y Energía

(MINAE 1999, Rodríguez-Herrera et al. 2002). Clearly, additional fieldwork is necessary to document the distribution, population size, and general ecology of *S. aphantus* in Monteverde and throughout its currently known range at mid-elevations in Central and South America.

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Appendix 1

Specimens Examined

Locality information not provided by the collector appears in brackets. Where original elevation was reported in feet, we either provide that datum or the metric equivalent to the nearest 10 meters. Those specimens used in multivariate analyses are denoted with an asterisk (*).

Mindomys hammondi (2).—ECUADOR: *Napo*; Concepción [00°48'S, 77°25'W] (1♀, MCZ 52543) [The locality for this specimen may be in error. Concepción is at ca. 300–500 m in the eastern lowlands (see Paynter 1993), whereas all other known specimens of *M. hammondi* are from the Mindo area of the western slopes of Volcán Pichincha. Paynter (1993) reports that the Olallas collected at Concepción on 25–26; the date on the single specimen of *M. hammondi* obtained by the Olallas is 27 July 1929]. *Pichincha*; Mindo, 4200 ft [1,380 m] [00°03'S, 78°46'W] (1♂, UMMZ 155827) (Voss 1988).

Nephelomys albigularis (5).—PERU: *Amazonas*: Goncha, 8,500 ft [2790 m] (1♀, KU 79328). VENEZUELA: *Aragua*; Rancho Grande, 1100 m [10°22'N, 67°41'W] (Voss 1988) (3♂♂, 1♀, KU 120296, 120297, 120298, 120299).

Nephelomys devius (36).—COSTA RICA: *Alajuela*; Monteverde, Monteverde Cloud Forest Reserve, Cerro Amigos, 1750–1840 m [10°19'N, 84°48'W] (5♂♂, 2♀♀, KU 159043, 159044, 160221, 160222, 160982, 160983, 160984); Monteverde, Monteverde Cloud Forest Reserve, Peñas Blancas Trail to Refugio Aleman, 1000–1050 m [10°18'N, 84°47'W] (1♂, KU 160985); Monteverde, Monteverde Cloud Forest Reserve, Sendero Pantanosa, 1580 m [10°18'N, 84°48'W] (1♂, KU 143401); Monteverde, Monteverde Cloud Forest Reserve, Sendero Peñas Blancas, 1250–1300 m [10°18'N, 84°47'W] (5♂♂, 2♀♀, KU 159045, 159046, 159047, 159048, 159050, 159051, 159052). *Cartago*; Cordil-

lera de Talamanca, upper Río Macho watershed, Villa Mills, ca 2750 m [9°33'N, 83°43'W] (1♂, KU 143311); Refugio Nacional de Fauna Silvestre Tapantí, near Río Dos Amigos (1♀, KU 142137*); San Vito, Coto Brus, Estación Biológica Las Alturas, 1500 m [8°57'N, 82°50'W] (1♀, KU 158634). *Puntarenas*; Monteverde, Cfemalebell's Woods, 1500 m [10°18'N, 84°48'W] (1♀, KU 143655*); Monteverde, Cerro Amigos, 1,760 m [10°19'N, 84°48'W] (3♂, KU 142060*, 142061*, 142062*); Monteverde, Monteverde Cloud Forest Reserve, 1580 m [10°18'N, 84°48'W] (1♀, KU 143403*); Monteverde, Monteverde Cloud Forest Reserve, Brillante Trail, 1500–1560 m [10°18'N, 84°48'W] (3♂♂, 1♀, KU 160956*, 160958*, 160959*, 160960*); Monteverde, Monteverde Cloud Forest Reserve, Cerro Amigos, 1800–1820 m [10°19'N, 84°48'W] (1♂, KU 160961*); Monteverde, Monteverde Cloud Forest Reserve, Investigators' Trail, 1550 m [10°18'N, 84°48'W] (4♂♂, KU 159,000*, 159001*, 159002*, 159003*); Monteverde, Monteverde Cloud Forest Reserve, Sendero Chomogo, ~1660 m [10°18'N, 84°48'W] (1♀, KU 143402*). *Puntarenas*–*Guanacaste* border; Monteverde, Cerro Amigos, 1790 m [10°19'N, 84°48'W] (1♂, 1♀, KU 143404*, 143405*).

Sigmodontomys alfari (29).—COLOMBIA: *Caldas*; Samaná, Río Hondo [05°17'N, 75°15'W] (1♂, FMNH 71639); *Córdoba*; Socorré, upper Río Sinú, 110 m [09°24'N, 75°49'W] (1♀, 1♂, FMNH 69192, 69195). COSTA RICA: *Cartago*; Perálta, El Sauce [09°58'N, 83°37'W—for Perálta] (1♂, AMNH

123305); Perálta, Santa Teresa (1♂, AMNH 141877). HONDURAS: *Gracias a Dios*; Río Coco [14°45'N, 84°03'W] (2♂♂, USNM 392876, 392877). NICARAGUA: *Matagalpa*; Río Grande (1♂, 1♀, AMNH 28547, 28549). PANAMA: *Bocas del Toro*; Almirante [09°18'N, 82°24'W] (4♂♂, USNM 315996, 315997, 315998, 315999). *Darién*; Boca de Río Paya, Cana, 2000 ft [660 m] [07°55'N, 77°31'W] (1♀, FMNH 53998); 1800 ft [590 m] [07°55'N, 77°31'W] (6♂♂, USNM 178625, 178626, 178630, 178631, 178632, 178633); Tacarcuna, 2650 ft [870 m] [08°07'N, 77°17'W] (8♂♂, AMNH 37900, 37901, 37904, 37905, 37908, 37909, 37912, 37914); Tacarcuna Village, 3200 ft [1050 m] [08°05'N, 77°17'W] (1♂, USNM 310586).

Sigmodontomys aphrastus (6).—COSTA RICA: *Alajuela*; Monteverde, Monteverde Cloud Forest Reserve, Sendero Peñas Blancas, 1300 m [10°18'N, 84°47'W] (1♂, KU 161003). *Puntarenas*; Monteverde, Monteverde Cloud Forest Reserve, Sendero Brillante, 1550 m [10°18'N, 84°48'W] (1♀, KU 159021). ECUADOR: *Pichincha*; Guarumos on the “W. slope of Mt. Pichincha, 4200 ft” (1♂, MCZ 50396) [The specimen tag records the elevation of Guarumos as 4200 ft. Paynter & Traylor (1977) and Paynter (1993) place Guarumos at 1500–2000 m [00°04'S, 78°38'W]; however, IGM (1972) places Guarumos at 2600 m [00°03'S, 78°38'W]; Mindo, 4200 ft [1380 m] [00°03'N, 78°48'W] (1♀, UMMZ 155808) (Voss 1988). PANAMA: *Chiriquí*; 24 km NNE San Félix, ca. 1275–1325 m (2♂♂, USNM 541200, 541201).