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The Bolivian bamboo rat, Dactylomys boliviensis (Rodentia: Echimyidae), a new record for chromosome number in a mammal

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16 recent genera and about 70 living species (Woods 1993), however, new taxa continue to be described (e.g., DA SILVA 1998; Patton et al. 2000). To date no comprehensive phylogenetic analysis is available for the group although great advances have been forthcoming (e.g., LARA et al. 1996; PATTON et al. 2000). The group is highly diversified ecologically and has had a long evolutionary history in South America (PATTERSON and PASCUAL 1972; WOODS 1982). Karyologically, less than half of the species have been analyzed but it is known that there is considerable variation in di-

The family Echimyidae, a highly diverse group of rodents, occurs throughout most

of Central and South America. There are

ploid (from 2n = 14 to 2n = 96) and funda-(from FNa = 18FNa = 144) (Tab. 1). One of the most specialized groups within the Echimyidae is the subfamily of bamboo rats (Dactylomyinae). Woods (1993) placed three genera in the subfamily Dactylomyinae; Dactyloms, Kannabateomys, and Olallamys. The biology and evolutionary relationships of the

mals of Bolivia, many new and important records for the country were collected (An-DERSON 1997). In July of 1992 and May of 1996, we took a total of five specimens of Dactylomys boliviensis (Bolivian bamboo rat) from a locality in the Yungas of La Paz (SALAZAR et al. 1994). Here we report the karyotype of this species, the highest chromosomal number known in a mammal. The individuals were located and collected at night in a dense stand of bamboo and

is known is well summarized in PATTON et

al. (2000). Until recently (ANISKIN 1993)

no information on the chromosomal com-

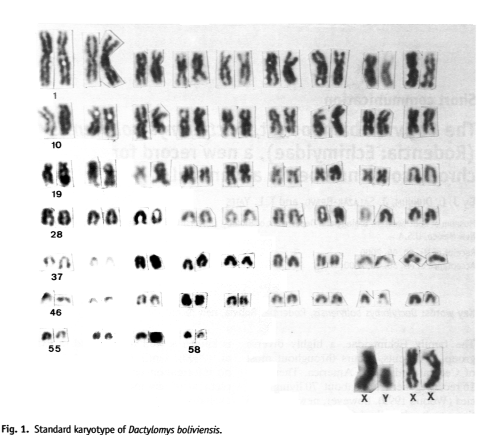
plement of any member of this group was

As part of a long-term survey of the mam-

secondary growth within the village of La Reserva (Departamento La Paz, Nor Yungas, La Reserva, elev. 840 m, 15° 44′ S, 67°31'W) by following their distinctive calls and eye shine. The village of La Reserva lies along Rio La Reserva, a small tributary of the Caranavi River. The village is at the bottom of a valley in the subtropical montane forest that covers most of the eastern Andean slopes between 15° and 17°S Dactylomyinae are poorly known, likely due to their rarity to collectors and subselatitude in the Cordillera Oriental of Bolivia. The foothills at this elevation are covquent scarcity in museum collections. What

numbers

mental



ered with semi-deciduous vegetation inter-

mingled with columnar cacti and bro-

meliads. The forest is drier and sparser than at higher elevations. Compared to forests at higher and lower elevations, the trees are smaller, more highly branched, and most grow in open sun. The east facing slope above the river is steep, with much vegetation, some secondary growth, and banana and tangerine cultivation. Palms and tree ferns are absent (SALAZAR et al. 1994).

tion, some secondary growth, and banana and tangerine cultivation. Palms and tree ferns are absent (SALAZAR et al. 1994). Chromosomal preparations were obtained using the technique described in ANDERSON et al. (1987). Metaphase cells were photographed and scored to determine the diploid (2n) and fundamental numbers (FNa). One of us (JLD) scored 5 slides per animal and over 20 spreads per slide to determine chromosome numbers. The analysis of the morphology of the chromosomes was based on 10 metaphase plates from three

individuals. Nomenclature for chromosome morphology and fundamental number follows PATTON (1967). Chromosome slides, tissue samples, and cell suspensions are deposited in the Division of

Biological Materials, Museum of South-

western Biology (MSB). Voucher speci-

mens are deposited at MSB (MSB 68547,

MSB 85627, NK 40537), the American Mu-

seum of Natural History (AMNH 264887, 264884), and the Colección Boliviana de Fauna (CBF 2608), in La Paz, Bolivia. The standard karyotype of *Dactylomys boliviensis* is highly asymmetrical, composed of 26 pairs of metacentric or sub-metacentric autosomes and 32 pairs of acro-

centric autosomes. The X chromosome is a large sub-metacentric and the Y chromosome is a medium sub-metacentric. The resulting karyotype has a diploid count of 2n = 118 and FNa of 168 (Fig. 1). Chromo-

Table 1.	Diploid (2n) and fundamental number (FN) for members of the family Echimyidae.	

FN

168

2n

118

Taxon

Dactylomys boliviensis

Dactylomys boliviensis, new record for chromosome number

Reference

this report

Dactylomys boliviensis	118	168	this report
Dactylomys dactylinus	94	144	Aniskin (1993)
Echimys blainvillei	50	94	Reig (1989)
Echimys dasythrix	96	102	Lima et al. (1998)
Echimys semivillosus	94	134	Aguilera et al. (1998)
Echimys sp.	90	108	Lima et al. (1998)
Echimys sp.	90	110	Aniskin (1993)
Echimys sp.	90	112	Reig (1989)
Isothrix bistriata	60	116	Patron et al. (2000)
Isothrix bistriata	60	120	Lima et al. (1998)
Isothrix pagurus	22	38	Patton and Emmons (1985)
Isothrix sinnamariensis	28	42	VIE et al. (1996)
Makalata armata	70	120	LIMA et al. (1998)
Makalata didelphoides	66	106	Lima et al. (1998)
Clyomys laticeps	34	60	Reig (1989)
Euryzygomatomys guiara	46	82	Aniskin (1993)
Euryzygomatomys spinosus	46	92	Reig (1989)
Hoplomys gymnurus	46		Aniskin (1993)
Lonchothrix emiliae	60	116	Aniskin (1993)
Mesomys hispidus	60	120	LIMA et al. (1998)
Mesomys hispidus	60	116	Раттом et al. (2000)
Mesomys occultus	42	54	PATTON et al. (2000)
Proechimys albispinus	60	116	LEAL-MESQUITA et al. (1992)
Proechimys amphicoricus	26	44	Reig (1989)
Proechimys brevicauda	28-30	48-50	GARDNER and EMMONS (1984)
Proechimys canicollis	24	44	GARDNER and EMMONS (1984)
Proechimys cuvieri	28	46	MAIA and LANGGUTH (1993)
Proechimys decumanus	30	54	GARDNER and EMMONS (1984)
Proechimys echinothrix	32	69	PATTON et al. (2000)
Proechimys gardneri	40	56	Patton et al. (2000)
Proechimys goeldii	24	44	Patton et al. (2000)
Proechimys quiarae	44-50	72- 76	GARDNER and EMMONS (1984)
Proechimys gularis	30	48	GARDNER and EMMONS (1984)
Proechimys guyannensis	40	54-56	GARDNER and EMMONS (1984)
Proechimys iheringi	62-65	117-124	Reig (1989)
Proechimys kulinae	34	52	Patton et al. (2000)
Proechimys mincae	48	68	GARDNER and EMMONS (1984)
Proechimys oconnelli	32	52	GARDNER and EMMONS (1984)
Proechimys oris	30	52-56	GARDNER and EMMONS (1984)
Proechimys pattoni	40	56	PATTON et al. (2000)
Proechimys poliopus	42	76	GARDNER and EMMONS (1984)
Proechimys quadruplicatus	28	44	Gardner and Emmons (1984)
Proechimys semispinosus	30	50- 54	GARDNER and EMMONS (1984)
Proechimys simonsi	32	58	GARDNER and EMMONS (1984)
Proechimys steerei	24	42	GARDNER and EMMONS (1984)
Proechimys trinitatus	62	80	GARDNER and EMMONS (1984)
Proechimys urichi	62	88	GARDNER and EMMONS (1984)
Proechimys yonenagae	54	104	LEAL-MESQUITA et al. (1992), ROCHA (1995)
Proechimys sp.	34	56	Aniskin (1993)
Proechimys sp.	14-16	18	Reig (1989)
Proechimys sp. (Balta)	40	56	REIG (1989)
Proechimys sp. (Barinas)	62	74	GARDNER and EMMONS (1984)
Thricomys aperoides	26	48	LEAL-MESQUITA et al. (1993)
Thricomys aperoides	30	54	Reig (1989)
Thricomys aperoides	30	50	Aniskin (1993)



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180

160



60

Diploid number (2n)

Eumysopinae

70

80

Fig. 2. Karyogram of known echimyid karyotypes.

20

30

40

Echimyinae

50

10

Dactylomyinae

satellite chromosome found in other echimvid rodents. No chromosomal information is available for Kannabateomys, Olallamys, or D. peruanus. Aniskin (1993) described the karyotype of D. dactylinus (2n = 94, Fn = 144)

from the Loreto Department in Peru. The

some pair nine exhibits the characteristic

karyotype of D. boliviensis differs from that of D. dactylinus by the presence of one additional set of meta or sub-metacentric pairs, and 10 pairs of acrocentric chromosomes although comparisons are difficult due to the fact that Aniskin (1993) did not identify sex chromosomes. At least 14 Robertsonian rearrangements would be necessary to transform the karyotype of one spe-

We compiled a list of all species of echimyid

rodents for which data were available

cies into the other.

I. sinnamariensis). The most speciose and karyologically studied group is the Eumysopinae (represented in this sample by Proechimys, Clyomys, Euryzygomatomys, Hoplomys, Lonchothrix, Mesomys, and Thrichomys). For the most part these fall at the lower end of (Fig. 2). To date, no eumysopids have been found with a 2n > 65.

90

100

(Tab. 1) and created a karyograph (IAMI

and Crozier 1980) based on chromosomal

and fundamental numbers (Fig. 2). A defi-

nite pattern of subfamily grouping is clear

where two species of Dactylomys assume

the highest positions on the plot and the

echimyine rodents (Echimys, Makalata, Iso-

thrix) are positioned at an intermediate le-

vel (with the exception of I. pagurus and

110

120

LIMA et al. (1998) proposed that Robertsonian rearrangements were more important in the evolution of the karyotype of arboDactylomys boliviensis, new record for chromosome number

Prior to our results, the highest chromo-

some number reported for a mammal was

2n = 102 in Tympanoctomys barrerae (Con-

TRERAS et al 1990). These authors also sug-

gested that the family Octodontidae pre-

sented the greatest chromosomal diversity.

While this remains true for Fundamental

number, the Echimyidae now represent the

family with the greatest diversity in diploid

Although Tympanoctomys and Dactylomys

represent terminal branches in two different

families of South American hystricognath

rodents with a long history on this continent,

they also share another characteristic: both

occupy restricted ecological niches and pos-

ses highly specialized life history traits. We

concur with Contreras et al. (1990) in sug-

gesting that the high chromosomal count ap-

number (2n = 14 to 2n = 118).

pears to be a derived character.

(Kruskal-Wallis; P < 0.001). However, in all cases the fundamental number varied more than the diploid number, suggesting that pericentric inversions may be more common. None the less it is quite likely that several processes may have influenced the evolution of the karyotype in this group. Acknowledgements This work was supported by grants from the National Science Foundation and a Research Experience for Undergraduates supplement. We would like to thank the 1992 and 1996 members of the joint CBF-MNK-AMNH-MSB field expe-

real echimyids than other chromosomal re-

arrangements because karyotypes of this

group appeared to show higher levels of

variation in diploid numbers than in funda-

mental numbers. Our data do not support

LIMA et al. (1998). We found statistically

significant differences in the levels of var-

iation between diploid and fundamental

number for the arboreal echimyids (Krus-

kal-Wallis; P < 0.004), terrestrial echimyids

(One-way ANOVA; a = 0.05; P < 0.004),

and for the entire echimyid radiation

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