The patterns and variability of colors in extant vertebrates are the result of the conjunct action of more than 150 genes that determine pigmentation. Nucleotide alterations in these genes are responsible for differences in pigmentation, which affect visual attraction, aposematism, and camouflage (Hofreiter and Schöneberg, 2010). Among mammals, the two factors that determine color patterns are the presence and distribution of pigments in skin, eyes, and hair (Hofreiter and Schöneberg, 2010; Abreu et al., 2013). However, chromatic abnormalities can occur due to the excess or deficit of melanin, either partially or totally throughout animal’s body (Acevedo and Aguayo, 2008; Abreu et al., 2013).

In recent zoological studies, chromatic anomalies have been classified into piebaldism, leucism, albinism, and melanism (Fertl and Rosel, 2002; Miller, 2005; Acevedo and Aguayo, 2008), making the terms “partial” and “total” obsolete when referring to different types of albinism (Summers, 2009; Abreu et al., 2013). Fertl and Rosel (2002) characterized piebaldism as the absence of pigments in some parts of an animal’s body. Leucism is characterized by the absence of pigments in most of an animal’s body, while the eyes and body extremities are still pigmented. Furthermore, albinism occurs when there is an absence of pigments throughout the body, including the eyes (Fertl and Rosel, 2002; Miller, 2005; Acevedo and Aguayo, 2008). Finally, melanism is characterized by dark coloration of parts of the body due to high melanin deposition (Hubbard et al., 2010).

Many Neotropical vertebrates have been reported with these anomalies, including fishes (e.g. Sazima and Pombal, 1986; Leal et al., 2013), amphibians (e.g. Sanabria et al., 2010), reptiles (e.g. Silva et al., 2010; Ayala-Monedero and Álvarez-León, 2014), birds (e.g. Cavarzere and Tonetti, 2015) and mammals (e.g. Acevedo and Aguayo, 2008; Toledo et al., 2014). However, these anomalies are considered rare in free-ranging populations (Parsons and Bonderup-Nielsen, 1995) due to the assumed short life expectancy of animals carrying these traits, since they would be more susceptible to predation (Sazima and Di-Bernardo, 1991).

Abreu et al. (2013) reported 198 cases from 26 species of Neotropical mammals that presented deficiencies in their melanin production, with only four cases for the order Rodentia. These authors emphasized the importance of knowing the type and frequency of these phenomena to understand the consequences of the anomalous coloration on the survival of wild mammals and the most susceptible taxa. Herein, we present the first reports of piebaldism in the Akodontini rodent *Akodon cursor* (Winge, 1887) and melanism in *Scapetomys tumidus* (Waterhouse, 1837).

On March 28th 2011, a piebald individual of *A. cursor* (male; head-body length=129mm; tail length=104mm; body mass=55g) was captured at the Sítio Boa Sorte (21°20’20”S, 42°45’43”W, 288m a.s.l.), located on the border of the municipalities of Cataguases and Dona Euzébia, Minas Gerais state. This area lies within an Atlantic Forest fragment characterized as Submontane Seasonal Semideciduous Forest (Veloso et al., 1991). Species identification was confirmed through comparisons with skulls of *A. cursor* specimens from the mammalian collections, as well as through microscopic analyses of the chromosomes after Giemsa stain procedure, following the morphological classification proposed by Levan et al. (1964). The specimen presented diploid number (2n)=14 and fundamental number (NF)=20, which is in agreement with the karyotype of *A. cursor*. Additional morphological and molecular analyses are ongoing to identify the chromosomal changes responsible for the anomalous coloration.
with karyotypic data reported for *A. cursor* (Di-Nizio et al., 2017). Skull, skin, and tissues of the specimen were deposited into the mammalian collection of “Museu de Zoologia João Moojen da Universidade Federal de Viçosa” (MZUFV 4423), Viçosa, Minas Gerais state, Brazil (Figure 1A).

On September 27th 2011, a melanic individual of *S. tumidus* (male; head-body length=136mm; tail length=131mm; body mass=85g) was captured from a marshy area with predominance of herbaceous *Eryngium pandanifolium* in “Bujuru” (31°36′34″S, 51°23′40″W), São José do Norte County, Coastal Plain of Rio Grande do Sul State, in southern Brazil. The individual had full darkish dorsum and ventral pelage, except for a small whitish spot in the gular region. Species identification was confirmed through comparisons with skulls of *S. tumidus* specimens from the mammalian collection and cytochrome *b* (Cytb) sequences obtained from GenBank. Skull, skin, and tissues of the specimen were deposited into the mammalian collection of “Departamento de Genética da Universidade Federal do Rio Grande do Sul” (TR 1588), Porto Alegre, Rio Grande do Sul, Brazil (Figure 1B). Cytb sequence was deposited in GenBank (access number KP233850, haplotype 5; Quintela et al., 2015).

At least 722 native mammal species occur in Brazil, of which more than 34% are rodents (Percequillo et al., 2017). In Brazil, chromatic anomalies in rodents have only been reported for *Trinomys albinsinus* (I. Geoffroy, 1838) (Pessoa and Reis, 1995), *Delomys dorsalis* (Hensel, 1873) (Cademartori and Pacheco, 1999), *Dasyprocta azarae* (Lichtenstein, 1823) (Oliveira, 2009) and *Thrichomys pachyurus* (Wagner, 1845) (Neves et al., 2014). These reports only represent 1.7% of the rodent species that occur in Brazil. Such low number could be associated with the low position that rodents occupy in the trophic network, since they are more exposed to predation (Abreu et al., 2013). This is the first report of chromatic anomalies occurring in *A. cursor* and *S. tumidus*. Such anomalies seem to be rare in both species, considering that a large number of specimens have already been collected.

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