Cytotaxonomy of the subgenus *Artibeus* (Phyllostomidae, Chiroptera) by characterization of species-specific markers

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Abstract

The genus *Artibeus* represents a highly diverse group of bats from the Neotropical region, with four large species occurring in Brazil. In this paper, a comparative cytogenetic study was carried out on the species *Artibeus obscurus* Schinz, 1821, *A. fimbriatus* Gray, 1838, *A. lituratus* Olfers, 1818 and *A. planirostris* Spix, 1823 that live sympatrically in the northeast of Brazil, through C-banding, silver staining and DNA-specific fluorochromes (CMA₃ and DAPI). All the species had karyotypes with 2n=30,XX and 2n=31,XY₂ and FN=56. C-banding showed constitutive heterochromatin (CH) blocks in the pericentromeric regions of all the chromosomes and small CH blocks at the terminal region of pairs 5, 6, and 7 for all species. Notably, our C-banding data revealed species-specific autosomic CH blocks for each taxon, as well as different heterochromatic constitution of Y₂ chromosomes of *A. planirostris*. Ag-NORs were observed in the short arms of chromosomes 5, 6 and 7 in all species. The sequential staining AgNO₃/CMA₃/DA/DAPI indicated a positive association of CH with Ag-NORs and positive CMA₃ signals, thus reflecting GC-richness in these regions in *A. obscurus* and *A. fimbriatus*. In this work it was possible to identify interspecific divergences in the Brazilian large *Artibeus* species using C-banding it was possible provided a suitable tool in the cytotaxonomic differentiation of this genus.
Introduction

The genus *Artibeus* Leach, 1821 has been divided into two main groups based on body size. The species with larger body size have been classified as subgenus *Artibeus* and the species with smaller body size as the subgenus *Dermanura* Gervais, 1856. In addition, a new subgenus *Koopmania* Owen, 1991 was proposed by Owen (1987, 1991) to set apart one of the species, *A. concolor* Peters, 1865. However, this subgenus was later disregarded by Van De Bussche et al. (1998) based on morphological, enzymatic, molecular and karyotypic analysis. The taxonomic classification proposed by Hoofer et al. (2008) recognized *Artibeus* as a distinct subgenus from *Dermanura*. Recently, Marchán-Rivadeneira et al. (2010) described the genus *Artibeus* as constituted by 11 large-body size species including *A. concolor* as the basal taxon. Its distributional range is restricted to the Neotropical region. The genus is widely distributed from Mexico to northern Argentina, including the Antillean islands in the Caribbean (Simmons 2005).

The extensive similarity of morphometric characters, high degree of shape diversity and overlapping of natural habitats have hindered accurate identification of the large *Artibeus* along their distribution, particularly in the Neotropical region (Haynes and Lee 2004, Hollis 2005). A typical example is the northeastern region of Brazil where four species of the large *Artibeus* (*A. obscurus* Schinz, 1821, *A. fimbriatus* Gray, 1838, *A. lituratus* Olfers, 1818 and *A. planirostris* Spix, 1823) were formally recorded living in sympatry (Taddei et al. 1998, Araújo and Langguth 2010). In this region, similarity in morphometric measurements (e.g. cranial distances and external dimensions) and geographical variation of *A. planirostris* are main reasons for confusing taxonomy (Guerrero et al. 2003, Simmons 2005).

Since the systematic classification of subgenus *Artibeus* remains subject of several discussions concerning phylogenetic relationships and actual taxonomic status of species, the use of complementary information may help to define species more precisely (Larsen et al. 2010). For other Mammalian groups, such as primates, felines and rodents, classical and molecular cytogenetic analysis have been successfully allied to taxonomic studies to identify species since chromosomes are not affected by adaptation process to different feeding niches as cranial and general gross anatomies (Granjon and Dobigny 2003, Garcia and Pessoa 2010).

In this work a karyotypic characterization of *A. obscurus* Schinz, 1821, *A. fimbriatus* Gray, 1838, *A. lituratus* Olfers, 1818 and *A. planirostris* Spix, 1823 from northeastern of Brazil was performed by the cytogenetic techniques – conventional analysis, C-banding, Ag-NOR and triple staining CMA3/DA/DAPI. The data were helpful to carry a comparative analysis of those species, in terms of interspecific differences, and also to provide a better identification of them.
Material and methods

Based on literature (Handley 1989, Taddei et al. 1998), the following characters were used to diagnose the species: presence of fur on the forearms, structure of legs and interfemoral membrane; color of body, dorsal and ventral fur; facial stripes; form of nose leaf and its relationship with the upper lip; shape of the pre- and postorbital process and postorbital constriction and the presence or absence of the 3rd molar. The identification process also included the following 10 measurements: length of forearm, condylobasal length; length of maxillary tooth-row; length of lower tooth-row; length of mandible; breadth across upper canines; mastoidal breadth; zygomatic breadth; postorbital constriction; breadth across upper molars.

After identification, cytogenetic studies were carried out on 53 Artibeus specimens from the state of Pernambuco, northeastern Brazil. Voucher specimens are deposited in the Mammalian collection at the Department of Systematic and Ecology, Federal University of Paraíba, João Pessoa, Paraíba, Brazil. The specimens studied were six males and eight females of Artibeus obscurus; two males and four females of A. fimbriatus; eight males and five females of A. planirostris; ten males and ten females of A. lituratus captured at different sites across the Pernambuco State: Igarassu (07°50′02″S, 34°54′21″W), Água Preta (08°42′27″S, 35°31′50″W), Rio Formoso (08°39′50″S, 35°09′32″W), Ipojuca (08°24′00″S, 35°03′45″W) and Recife (08°03′14″S, 34°52′51″W) (see also Appendix).

Metaphase spreads were obtained from bone marrow cells according to conventional procedures and staining with Giemsa. C-banding and silver staining were performed according to Sumner (1972) and Howell and Black (1980), respectively. Triple staining CMA3/DA/DAPI was carried out according to Santos and Souza (1998a).

For sequential staining (AgNO3/CMA3/DA/DAPI), the slides stained by silver nitrate were distained after photographing (Dos Santos Guerra 1991) and re-stained by CMA3/DA/DAPI. Photomicrographs were taken using Leica DMLB photomicroscope for C-banding and silver staining. Sequential staining images were captured by IM50 capture system.

Results

All four species shared the same diploid number (2n=30, gap XX; 2n=31, gap XY1Y2) and fundamental number FN=56. Chromosomes were meta-submetacentric (1-4, 8-14), subtelocentric (5, 6, 7 and X) and two small acrocentric (Y1 and Y2). Except for the size of Y1 and Y2 chromosomes, it was not found any intraspecific variation between species analyzed with conventional staining.

C-banding revealed constitutive heterochromatin (CH) in the pericentromeric region of all the autosomes and small heterochromatic blocks were observed in the terminal region of chromosome pairs 5, 6 and 7 (Fig. 1a–d). The karyotype of A. obscurus (Fig. 1a) exhibited interstitial blocks in the short and long arms of pair 1, as well as in the long arms of pairs 2, 5, 6 and in the terminal region of the short arm of pair 9. The A. planirostris karyotype
Figure 1. C-banding of *A. obscurus* (a) *A. fimbriatus* (b) *A. lituratus* (c) and *A. planirostris* (d) karyotypes. The arrowheads indicate a particular set of CH blocks in each species. Bar = 5 µm.

(Fig. 1d) has the same CH pattern but lacks interstitial blocks in the short arm of chromosome 1. Absence of an interstitial block on the chromosome 6 distinguished karyotype of *A. fimbriatus* from the other investigated karyotypes (Fig. 1). In all the material examined the long arms of the X chromosomes were more darkly stained when compared with the euchromatin of the autosomes. The Y2 appeared almost entirely heterochromatic in all species, except for *A. planirostris* which showed pericentromeric and distal blocks (Fig. 1d). The pattern of the Y1 could not be determined with precision due to its punctiform size.

Table 1 shows exhibits the C-banding pattern in chromosomal complement in all species analyzed.

Silver staining (Ag-NORs) showed three pairs of NORs in the terminal region of the short arms in chromosomes 5, 6 and 7 in all species. As a result of remarkable variation in expression and activity, Ag-NORs were counted up to 100 nuclei, which were randomly selected, and the mean number of Ag-NORs per nucleus was determined for each case (Table 2).

The sequential staining AgNO3/CMA3/DA/DAPI showed a correlation between CMA3 positive regions and Ag-NORs in the karyotypes of *A. obscurus* and *A. fimbriatus* (Fig. 2a–f). Karyotypes of both species had presented CH blocks associated with Ag-NORs sites, reflecting GC-richness in these heterochromatics clusters. In addition, positive CMA3 signals were observed in the pericentromeric regions of certain autosomes, particularly in pairs 1, 2 and 6 of *A. obscurus* and in pair 6 of *A. fimbriatus* (Fig. 2b,e). On the other hand, a uniform pattern was observed in all the chromosomes after DA/DAPI staining (Fig. 2c–d).
Figure 2. Sequential staining of *A. obscurus* (a–c) and *A. fimbriatus* (d–f) karyotypes with AgNO₃/CMA₃/DA/DAPI. (a,d) Ag-NORs, (b,e) CMA₃, (c–f) DA/DAPI. Bar = 5 µm.
Discussion

Our data regarding diploid number, chromosome morphology and sex determination system obtained for *Artibeus obscurus*, *A. fimbriatus*, *A. lituratus* and *A. planirostris* karyotypes are in agreement with those previously described in the literature (Baker and Hsu 1970, Gardner 1977, Baker et al. 2003). The autosomal complements presented are morphologically similar to other, except by length of the sex chromosomes Y1 and Y2, which varies from punctiform elements to well-defined acrocentric chromosomes, as firstly described by Hsu et al. (1968) for *A. lituratus*, *A. jamaicensis* Leach, 1821 and *A. toltecus* Saussure, 1860.

The multiple sex chromosome system XY1Y2 has been widely reported within the genus *Artibeus*, e.g. *A. aztecus* Andersen, 1906, *A. glaucus* Thomas 1893, *A. toltecus*, *A. concolor*, *A. cinereus* Gervais, 1856, *A. hirsutus* Andersen, 1906, *A. inopinatus* Davis et Carter, 1964 and *A. jamaicensis*, and for other 23 species of family Phyllostomidae, as predominant type of sex-determining mechanism in this group (Baker and Hsu 1970, Varella-Garcia et al. 1989, Wetterer et al. 2000, Baker et al. 2003, Noronha et al. 2009). In mammals, this sexual system has been reported in marsupials, insectivores (shrews), carnivores (mongoose), rodents (hamsters) and in artiodactyls (gazelles) (reviewed in Gruetzner et al. 2006). Its origin involves a single sex chromosome-autosome translocation, which in meiosis leads to one sexual trivalent structure formed by XY1Y2 (Rodrigues et al. 2003, Noronha et al. 2004).

The CH distribution was evaluated and intercompared in the large *Artibeus* and with others phyllostomatids, pointing out an extensive similarity of CH pattern local-

### Table 1. Heterochromatin pattern in chromosomal complement in *Artibeus* species

| Species         | Pericentromeric | Terminal | C-banding       | Distal |
|-----------------|-----------------|----------|-----------------|--------|
| *A. obscurus*   | +               | 5p, 6p, 7p, 9p | 1*, 2q, 5q, 6q | Y1 e Y2 |
| *A. fimbriatus* | +               | 5p, 6p, 7p | -               | Y1 e Y2 |
| *A. lituratus*  | +               | 5p, 6p, 7p | 6q              | Y1 e Y2 |
| *A. planirostris* | +            | 5p, 6p, 7p, 9p | 1q, 2q, 5q, 6q | Y1   | Y2   |

(p) = short arm; (q) = long arm; * = both p and q; + = all chromosomes; - = absent

### Table 2. Frequency analyzes of active NORs in the large species of genus *Artibeus.*

| Species          | Active NOR number per cell | Total of cells analyzed |
|------------------|---------------------------|-------------------------|
|                  | 1  | 2  | 3  | 4  | 5  | 6  |                  |
| *A. obscurus*    | 0  | 17 | 53 | 71 | 12 | 16 | 169               |
| *A. fimbriatus*  | 0  | 14 | 40 | 58 | 27 | 22 | 161               |
| *A. lituratus*   | 0  | 7  | 30 | 36 | 22 | 24 | 119               |
| *A. planirostris*| 0  | 14 | 28 | 47 | 11 | 25 | 125               |
| Total            | 0  | 52 | 151| 212| 72 | 87 | 574               |

(%) total 0 9.06 26.31 36.94 12.54 15.16
ized in the pericentromeric region (Rodrigues et al. 2000, Barros et al. 2009, Sbragia et al. 2010). Additionally, CH blocks were also found in the terminal region of chromosome pairs 5, 6 and 7 that has been considered a characteristic shared by subfamily Stenodermatinae (Souza and Araújo 1990, Santos and Souza 1998b, Silva et al. 2005).

On the other hand, a particular set of CH blocks was observed in *A. obscurus*, *A. fimbriatus*, *A. lituratus* and *A. planirostris*. This finding allowed the individualization and differentiation of each species for karyotype comparison (Fig. 1). *A. fimbriatus* and *A. lituratus* karyotypes showed a closer CH distribution differing only by one heterochromatin block. Furthermore *A. obscurus* and *A. planirostris* karyotypes presented more interstitial heterochromatin.

The occurrence of intrageneric variation on CH distribution had been described only in sporadic cases among phyllostomatids whose extensive karyotypic conservation is widely known. In turn, the genus *Artibeus* is widely cited as a chiropteran group that exhibits low rate of karyotype evolution whereas: (1) most of species had same diploid number (30/31) and (2) G-banding patterns are essentially identical (Baker and Bickham 1980, Baker et al. 2003).

The other parameter evaluated intercomparison was the NORs localization by silver staining. The Ag-NORs were situated on the subtelocentric autosomes 5, 6 and 7 of all species. The data obtained for *A. lituratus*, *A. planirostris* and *A. fimbriatus*, together with the new data of *A. obscurus*, were similar those described by Santos et al. (2002). These authors employed FISH with 18S ribosomal probe allied to silver staining to investigate the precise localization of rDNA sites, and discovered a non-correlation between the number and distribution of the NORs in *A. cinereus* Gervais, 1856, being the first report on silent NORs in bats. They also had distinguished two rDNA sites patterns for *Artibeus* genus: 1) in the distal regions of the short arms of pairs 5, 6 and 7 (*A. lituratus, A. jamaicensis* Leach, 1821 and *A. fimbriatus*) and 2) in the interstitial region of the long arms of pairs 9, 10 and 13 (*A. cinereus*). In addition, *A. fimbriatus* had one NOR in the interstitial region on the long arm of pair 5, that it was not observed in this work, which may indicate a chromosomal polymorphism for this species.

As only active NORs could be visualized in our data, the variation in Ag-NORs activity for cell was also investigated (Table 2). In the most of cells analyzed (> 500), the frequency of active NORs was 3 or 4 black spots (26.31 to 36.94 %). Such variability is in accordance with other studies on a NOR sites activity in Phyllostomidae bats that presents multiple NORs (Morielle and Varella-Garcia 1988, Souza and Araújo 1990, Santos et al. 2002).

The association between NORs and CH by GC-specific fluorochromes staining presented in this work for *A. obscurus* and *A. fimbriatus*, has also been reported to *A. lituratus, A. jamaicensis, Desmodus rotundus* Geoffroy, 1810, *Diphylla ecaudata* Spix, 1823 and *Lonchorhina aurita* Tomes, 1863. On the other hand, *Carollia perspicillata* Linnaeus, 1758, *Molossus molossus* Pallas, 1766, *M. ater* Peters, 1865, *Molossops planirostris* Peters, 1865, *Phylllostomus discolor* Wagner, 1843 and *Trachops cirrhosus* Spix, 1823 NORs and CH were CMA3 neutral. The reason for that is probably in heterogeneity of
base composition of the intergenic regions related to NORs. In some cases, the triple staining with CMA3/DA/DAPI has also enhanced the patterns of R-bands with CMA3, an uniform staining with DA/DAPI or a weak G-banding pattern, as it has been observed in some bat's families (Santos and Souza 1998a, 1998b, Santos et al. 2001, Leite-Silva et al. 2003, Barros et al. 2009).

**Conclusion**

Classical and molecular cytogenetic markers, associated to taxonomic studies, have provided a better understanding of phylogenetic relationships and the mechanisms responsible for chromosomal divergence in the different taxa in the order Chiroptera. Cytogenetic analysis of all Brazilian species of the subgenus *Artibeus* allowed us to reveal the conservative and specific chromosomal features among their karyotypes. Furthermore, it was possible to identify intrageneric and interespecific divergences in a group that up to today has been characterized by showing extensive karyotypic conservation. The cytogenetic techniques herein employed, demonstrated the usefulness of C-banding in the identification and correct individualization of the large *Artibeus* that live sympatrically in the northeastern of Brazil, thus providing an important tool in the cytotaxonomic differentiation of this genus.

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Appendix

Família Phyllostomidae

Artibeus obscurus
M 318 (3214) – Saltinho (Rio formoso)
M 319 (3206) – Saltinho (Rio formoso)
M 336 (3235) – Saltinho (Rio formoso)
M 337 (3238) – Saltinho (Rio formoso)
M 340 (3216) – Saltinho (Rio formoso)
M 344 (3237) – Saltinho (Rio formoso)
M 359 (3230) – Saltinho (Rio formoso)
M 381 (3181) – Saltinho (Rio formoso)
M 397 (3186) – Dois irmãos (Recife)
M 455 (3189) – Saltinho (Rio formoso)
M 478 (3179) – Saltinho (Rio formoso)

A. fimbriatus
M 346 (3215) – Saltinho (Rio formoso)
M 382 (3184) – Saltinho (Rio formoso)
M 395 (3177) – Dois irmãos (Recife)
M 453 (3175) – Saltinho (Rio formoso)
M 479 (3192) – Saltinho (Rio formoso)

A. lituratus
M 221 (3418) – Igarassu
M 379 (3178) – Saltinho (Rio formoso)
M 446 (3191) – Saltinho (Rio formoso)
M 454 (3182) – Saltinho (Rio formoso)
M 475 (3188) – Saltinho (Rio formoso)
M 476 (3180) – Saltinho (Rio formoso)

A. planirotris
M 118 (3424) – Igarassu
M 124 (3212) – Igarassu
M 137 (3423) – Aldeia (Camaragibe)
M 188 (3428) – Igarassu
M 262 (3220) – Água Preta (Fazenda Camarão)
M 263 (3218) – Água Preta (Fazenda Camarão)
M 393 (3202) – Dois irmãos (Recife)
M 400 (3176) – Dois irmãos (Recife)
M 401 (3196) – Dois irmãos (Recife)