Cytogenetic analysis of three Ctenidae species (Araneae) from the Amazon

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Abstract

Cytogenetic characterization was performed on three wandering spiders: Ctenus amphora Mello-Leitão, 1930, C. crulsi Mello-Leitão, 1930 and C. villasboasi Mello-Leitão, 1949. The three species had similar karyotypes, with 2n = 28 (26 + X0) in males, with sex chromosomes exhibiting positive heteropicnosis in meiotic cells. 18S rDNA mapping revealed gene sites at the terminal region of one chromosomal pair for all species, with one C. crulsi individual, showing markings in two pairs. C. villasboasi showed markers only in the pachytene phase. The distribution pattern of constitutive heterochromatin was found to be characteristic for the genus, with markings in the centromeric region of all chromosomes, suggesting an acrocentric morphology for all chromosomes of the three analysed species. The results support the fusion of sex chromosomes as an evolutionary tendency for this spider group.

Keywords: Meiosis, FISH, NORs, spider, Amazon.

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The order Araneae currently contains 48,455 described species, distributed in 120 families (World Spider Catalog, 2020). In Brazil, there are records of 3,203 species, of which 694 occur in the Amazon Region (Brescovit et al., 2011). Within this group, we focus here on the Ctenidae family, a group which has received increasing attention due to their value as bioindicators of environmental quality, and the use of their venom neurotoxins as therapeutic agents (Rego et al., 2005; Mestre and Gasnier, 2008; Pinheiro et al., 2009).

More than 868 species of spiders have already been characterized cytogenetically and, of these, 12 belong to the family Ctenidae (Araujo et al., 2020). The data for this family show a diploid number varying between 22 and 29 chromosomes for males, and X1X20 and X3X40 sex chromosome systems (Araujo et al., 2014). While these two systems are rare overall, they are common in spiders (Araujo et al., 2012), including the Ctenidae, making it possible, in this Family, to study the sex chromosome behavior during meiosis.

In Ctenidae, with the exception of Asthenoctenus borellii Simon, 1897, whose diploid number is 22, all species have 26 autosomal chromosomes plus two or three sex chromosomes (Araujo et al., 2020). Araujo et al. (2014) suggest that the conversion of the sexual chromosomal system X1X20 to X3X40, and vice-versa, is a relatively common event. However, available cytogenetic data are insufficient to allow inferences concerning evolutionary chromosomal tendencies within the group.

The aim of the current study is to increase the cytogenetic knowledge of the Ctenidae, and so contribute to the discussion concerning mechanisms of chromosome evolution in this family, especially regarding the behavior of sexual chromosomal systems X1X20 and X3X40, in males during meiosis.

A total of 10 individuals (5 males of Ctenus amphora and 5 males of C. crulsi) were collected in a forest fragment surrounding the Federal University of Amazonas (UFAM), in the eastern part of the city of Manaus (03°04’34”S, 59°57’30” W), and 16 individuals (6 males of C. amphora, 8 males of C. crulsi, and 2 males of C. villasboasi) in the Adolpho Ducke Forest Reserve (2°57’42”S, 59°55’40”W) (Figure 1). In order to identify species of Ctenus we used Hofer et al. (1994). Collections were carried out under SISBIO license number 60728-1. Chromosomal preparation of male gonads was conducted according to Araujo et al. (2008). Fluorescent in situ hybridization (FISH) was performed following Pinkel et al. (1986), using 18S rDNA probes (Almeida et al., 2010), that showed 91% homology with probes generated by Rincão et al. (2017) for Ctenus ornatus, and 92% homology for Ctenus crulsi. The sequences were compared in BLASTN, using the National
Center for Biotechnology Information (NCBI) database website (https://blast.ncbi.nlm.nih.gov/Blast.cgi). For C-banding the Sumner (1972) protocol was followed. For Ag-NOR the Howell and Black (1980) protocol was followed. In *C. amphora* (Figure 2A), *C. crulsi* (Figure 2B) and *C. villasboasi* (Figure 2C) 2n = 28, with 26 of the chromosomes being autosomal and two sexual. In pachytene (Figure 3A, D, G) and dipotene (Figure 3B, E, H) phase cells the sex chromosomes showed positive heteropiconis, thus allowing their identification. Determination of the sex chromosomal system was performed by analyzing chromosome segregation during metaphase II of meiosis (Figure 3C, F, I), which shows nuclei with n = 13 and n = 15 (13 + X1X2).

The FISH technique revealed 18S rDNA sites on pair 10 of the chromosomes in the spermatogonial metaphase of *C. amphora* and *C. crulsi* (Figure 2A, B). In addition, a *C. crulsi* individual from the Ducke Reserve was polymorphic compared to the other seven analyzed for this same locus, with four labeled chromosomes (Pairs 4 and 7) in spermatogonial metaphases (Figure 2B - highlighted). *Ctenus villasboasi* showed marking on a pair of chromosomes in metaphase (Figure 2C). However, chromosomal size visualized with this technique was insufficient to allow determination of which pair carried the 18S rDNA site.

C-banding showed the presence of constitutive heterochromatin in the centromeric region of all chromosomes, characterizing the morphology as acrocentric for all three species (Figure 4A-C). Silver nitrate impregnation revealed NORs on pair 10 in the spermatogonial metaphases for *C. amphora* and *C. crulsi* (most individuals) (Figure 2A, B). *C. crulsi* individual from the Ducke Reserve was polymorphic compared to the other seven analyzed for this same locus, with four labeled chromosomes (Pairs 4 and 7) in spermatogonial metaphases (Figure 2B - highlighted). *Ctenus villasboasi* showed marking on a pair of chromosomes in metaphase (Figure 2C). However, chromosomal size visualized with this technique was insufficient to allow determination of which pair carried the 18S rDNA site.

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The karyotypic formula of males (2n=28 - 26+X1X20), found for all species, has been shown to be conserved in *Ctenus* (Araujo et al., 2020). Identification of sexual chromosomes in meiotic cells was facilitated by their high degree of condensation and positive heteropiconis. The literature contains contrasting opinions regarding Ctenidae chromosomal morphology, with Araujo et al. (2014) considering them to be telocentric, while Kumar et al. (2017) and Rincão et al. (2017) consider that they are acrocentric. We considered the three species studied here to be acrocentric, based on C-banding, which showed markings in the centromeric regions of all chromosomes.

All three species showed metaphase II meiotic cells with n = 13 and n = 15 (13+X1X2), which agrees with the information given by Araujo et al. (2014) and Rincão et al. (2017) for Ctenidae species. This feature is common for a X1X20 sex chromosomal system, in which for males, at the end of meiosis I, two sex chromosomes migrate to the same pole cell.

Araújo et al. (2012) collated proposed spider chromosomal evolution theories, which include the X0 sexual chromosomal system giving rise to the X1X20 (White, 1940; Pátiau, 1948), via part of an X chromosome fissioning and attaching to a supernumerary chromosome (Bole-Gowda, 1950), and the sexual chromosomal system X1X2X30 giving rise to X1X20 via fusion of two X chromosomes (Král, 2007).

Taking into account the cytogenetic characterization of species considered basal within the Ctenidae, such as *Nothroctenus* sp. and *Viracucha andicola* Simon, 1906 (Pothow and Brescovit, 2014), and the data presented here for Ctenidae species occupying positions considered to be derived, we believe the diploid number reduction hypotheses based on chromosome fusion to be more parsimonious, since the first group has 2n = 29 and the second group has 2n = 28.
However, Rincão et al. (2020) recently found two individuals of *C. ornatus* showing one supernumerary chromosome and one individual with two supernumerary chromosomes. Those chromosomes showed positive heteropiconosis and behavior similar to sex chromosomes, which the author states may demonstrate conversion of sexual chromosomal system $X_1X_20$ to $X_1X_2X_3X_40$ for the first time in Ctenidae.

The FISH-obtained 18S rDNA tags for the three species in our study confirmed Ag-NOR derived data, and were similar to those described for *C. ornatus* and *C. medius* by Rincão et al. (2017). However, an individual of *C. crulsi* in the current study had two pairs of chromosomes with 18S rDNA labeling (Pairs 4 and 7). These additional markers were shown to be relatively minor when compared to that
found in the other *Ctenus* species of the current study, as well as those available in the literature. Such data suggest that these alterations can be caused by chromosomal rearrangements, insertions by transposable elements or ectopic recombination, all processes that could, potentially, be involved in karyotypic differentiation and new species emergence (Stáhlavsky et al., 2018).

Considering the number of chromosomes with 18S rDNA in *C. crulsi*, we believe the translocation hypothesis to be the most feasible, since this mechanism of chromosome evolution depends on the interchange of segments between two non-homologous chromosomes without loss of genetic material (Gross et al., 2010). Such translocations can be simple, when only the segment of one chromosome passes to the other, or reciprocal, when two chromosomes exchange segments with each other. The translocation model for rDNA is described by Araujo et al. (2015) for the exchange of such segments between autosomal and sexual chromosomes in the genus *Nephila*. This shows that a possible translocation between the two types of chromosomes is possible. In the *C. crulsi* metaphases analyzed here, the sex chromosomes were not evident.

Silver nitrate impregnation showed NOR markers on an autosomal pair for *C. amphora* and *C. crulsi* (most individuals), and on a pachytene bivalent for *C. villasboasi*, a result similar to that found for *C. ornatus* Keyserling, 1877 (Araujo et al., 2014; Rincão et al., 2020), and *C. medius* (Rincão et al., 2020), but which differs from those reported for *C. indicus* Gravely, 1931, by Kumar et al. (2017), and for *G. longipes* by Rincão et al. (2020), who found NOR markings on two chromosome pairs. The NOR distribution pattern is currently known for only five species in the genus *Ctenus*. Therefore, we suggest that additional cytogenetic studies are still needed to establish the plesiomorphic characteristics of the Ctenidae karyotype and, thus, to be able to understand the mechanisms of chromosomal evolution that occurred in this group of spiders.

The pattern of constitutive heterochromatin distribution reported here for *C. amphora*, *C. crulsi* and *C. villasboasi* is similar to those found by Rincão et al. (2017) for *C. medius* Keyserling, 1891, *E. cyclothorax* Bertkau, 1880, *P. nigriventer* Keyserling, 1880, and *V. andicola*, with blocks in the centromeric regions of all the chromosomes.

According to Sumner (1972), C-banding marks centromeric and telomeric regions, possibly marking nucleolar regions and, rarely, intercalated regions of the chromosome. In the metaphases studied here, the sex chromosomes were not totally heterochromatic, thus confirming heteropycnosis of the sex chromosomes in meiosis. Heterochromatic chromosomes remain condensed throughout the cell cycle, whereas heteropycnotic chromosomes may have a higher or lower level of condensation, depending on the stage of cell division in which they are found (John, 1990).

As a result of the current study, the number of Ctenidae species with chromosomal data has been increased to 14. The data obtained for *C. amphora*, *C. crulsi* and *C. villasboasi* extends to seven the number of species analyzed from the genus *Ctenus*, and so allow a conserved karyotype to be

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**Figure 3** - Meiotic cells in pachytene (A, D, G), diplotene (B, E, H) and metaphase II (C, F, I) of: *Ctenus amphora*. (A, B, C); *C. crulsi*. (D, E, F); *C. villasboasi*. (G, H, I). Metaphase II cells shows the division of chromosome between cells poles, with n = 13 and n = 15 (13+X1X2) during segregation. Scale= 10µm.
inferred, as well as the diploid number, chromosomal formula and sex chromosome system plus mapping of the rDNA sites, constitutive of sex chromosomes in meiotic cells in pachytene and diplotene phases is a common feature for these species (Araújo et al., 2012). The results also reinforce theories of chromosome fusion as a possible evolutionary tendency in this family.

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Conflict of Interest

The authors declare that there is no conflict of interest that could be perceived as prejudicial to the impartiality of the reported research.

Author Contributions

JPCPN, LGG, MCG and CHS conceived the study; JPCPN and LGG collected the individuals and conducted the experiments; JPCPN, LGG, MCG, EF and CHS analyzed the data; JPCPN, LGG, EF, MCG and CHS wrote the manuscript. All authors read and approved the final version.
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