In this study, we investigated the chromosomes of three species of *Sicarius* spiders from the Brazilian Caatinga, using classical and molecular cytogenetic techniques. Based on the phylogenetic approach, we also discussed about the variation of diploid number, types of sex chromosome system and changes in the localization of ribosomal genes of Scytodoidea. *Sicarius* are Synspermiata spiders that together with the genera *Loxosceles* and *Hexophthalma* constitute the family Sicariidae. In this group, the available cytogenetic data showed a low diploid number range (2n♂ = 18 to 2n♂ = 23) and the presence of only multiple sex chromosome systems (X0, Y and X0, Y0). Mitotic metaphase cells exhibited 2n♂ = 16+X0, Y for *Sicarius cariri* and *S. ornatus*, and 2n♂ = 18+XY for *S. tropicus*. In these species, silver impregnation revealed nucleolar organizer region (Ag-NOR) on the terminal region of pair 1. In *S. ornatus* and *S. tropicus*, the results obtained with fluorescent in situ hybridization (FISH) using 18S rDNA probe were similar to Ag-NOR, however in *S. cariri*, the ribosomal sites were localized in the terminal region of the X sex chromosome. In this work, we presented the first description of a simple sex chromosome system for Sicariidae, helping to understand how the XY sex chromosome system evolved from the X0, Y0 system. Additionally, FISH data incongruous with Ag-NOR indicate that the cytogenetic studies in Sicariidae allow investigating the relation between the karyotype evolution and the distribution and the activity of rDNA genes.

Key words: karyotype, mitosis, nucleolar organizer region, rDNA, *Sicarius*
INTRODUCTION

The spider family Sicariidae is considered of medical importance in the world (Lotz, 2012), including sedentary species, which can be ground-dwelling hunters or web-weavers (Dias et al., 2010). Sicariidae includes 171 species distributed into three genera: Hexophthalma composed of eight species, Sicarius, with 21 species, and Loxosceles, the most diversified genera with 142 representatives (World Spider Catalog, 2021). This latter genus is well known due to the toxicity of its venom, causing skin necrosis, renal failure and haemolysis (Silva et al., 2004; Vetter, 2008). Hexophthalma spiders occur only in southern Africa while Loxosceles presents widest distribution, with species described in America, Africa, Mediterranean Europe and Asia; however, the largest diversity of species is recorded in the American continent (World Spider Catalog, 2021). Sicarius is distributed in South and Central America and is restricted to xeric habitats, mainly deserts and tropical dry forests (Magalhaes et al., 2013).

For many years, in Brazil only one Sicarius species was known, S. tropicus (Mello–Leitão, 1936). However, recently Magalhaes et al. (2013, 2017) described other species from this country, namely S. boliviensis Magalhaes, Brescovit & Santos, 2017, S. cariri Magalhaes, Brescovit & Santos, 2013, S. diadorm Magalhaes, Brescovit & Santos, 2013, S. jequitinhonha Magalhaes, Brescovit & Santos, 2017, S. ornatus Magalhaes, Brescovit & Santos, 2013, and S. saci Magalhaes, Brescovit & Santos, 2017. The monophyly of Sicariidae is well supported by morphological (Platnick et al., 1991; Binford et al., 2008; Labarque y Ramírez, 2012; Magalhaes et al., 2013, 2017) and molecular data (Wheeler et al., 2017 contra Binford et al., 2008). Some characteristics considered as synapomorphies for siciariids are modifications in chelicerae setae, tarsal claws, abdominal entapophysis, and the venom protein sphingomyelinidase D, which is responsible for the envenomation symptoms (Binford y Wells, 2003; Magalhaes et al., 2017).

Sicariidae belongs to the monophyletic superfamil Scytodoidea composed by (Sicariidae (Drymusidae + Periogopidae) + (Ochyroceratidae + Scytodidae)) (Labarque y Ramírez, 2012; Wheeler et al., 2017). In this group, Scytodoidea is the most diverse, including a total of five genera and 245 known species (World Spider Catalog, 2021), but only five of them belonging to Scytodes were analyzed from the cytogenetic point of view (Araujo et al., 2021). The scytodids present a high variability in diploid number, from 2n=13 to 2n=31, but a simple and conserved sex chromosome system of the XO type. The exception is Scytodes globula Nicolet, 1849 that revealed an intraspecific variation due to the occurrence of XO and XX,0 systems (Diaz y Saez, 1966; Rodríguez–Gil et al., 2002; Araujo et al., 2008).

Ochyroceratidae possess 168 species described into 10 genera, but only a North American undetermined species of Ochyrocerca was cytogenetically analysed, exhibiting 2n=13 and XO sex chromosome system (Král et al., 2006). The family Drymusidae includes 17 species with chromosomal data only for Izithunzi capense (Simon, 1893), from South Africa, with 2n=37+X,X,Y (Král et al., 2006). Periegopidae is known only by three species from Queensland and New Zealand (World Spider Catalog, 2021), and there are no cytogenetic data for this family.

The family Sicariidae has karyotype information for 15 representatives, showing low diversity in the diploid number (2n=18 to 2n=23) and the occurrence of only multiple sex chromosome systems of the XX, Y and X,0 types (Araujo et al., 2021). Hexophthalma only has the diploid number 2n=20 described for females of an undetermined species (Král et al., 2019). The genus Loxosceles presents 12 species chromosomally characterized, in which the following diploid numbers were identified: 2n=18 in L. reclusa Gertsch & Mulaik, 1940; 2n=19 in L. spinulosa Purcell, 1904; 2n=20 in L. rufipes (Lucas, 1834); 2n=20–21 in L. rufescens (Dufour, 1820); 2n=23 in L. amazonica Gertsch, 1967, L. gaucha Gertsch, 1967, L. hirsuta Mello–Leitão, 1931, L. intermedia Mello–Leitão, 1934, L. laeta (Nicolet, 1849), L. puerto Martins, Knysak & Bertani, 2002, L. simillis Moenkhau, 1898 and L. variegata Simon, 1897. All these species showed XX, Y sex chromosomes system, except L. rufipes and L. reclusa that exhibited XX,0 system (Beçak y Beçak, 1960; Diaz y Saez, 1966; Hetzler, 1979; Silva, 1988; Tugmon et al., 1990; Oliveira et al., 1996, 1997; Silva et al., 2002; Král et al., 2006; Kumbiçak, 2014; Araujo et al., 2020).

In the genus Sicarius, only two species were investigated, S. tropicus (2n=19, XX, Y) from Brazil and an undetermined species (2n=21, XX, Y) from Cusco, Peru (Franco y Andía, 2013; Araujo et al., 2021), more likely to be Sicarius boliviensis, owing to the sampling locality (Magalhaes et al., 2017). Nevertheless, the cytogenetic data of S. tropicus could be considered preliminary because the karyotype information is restricted to a brief description of diploid number and sex chromosome system (Franco y Andía, 2013).

A cytogenetic analysis of three Sicarius species from the Brazilian fauna was accomplished in the present study, using standard staining, silver impregnation to reveal the active nucleolar organizer regions (NORs), and fluorescent in situ hybridization (FISH) with 18S rDNA probe to map the number and localization of the major ribosomal genes. Among the 21 Scytodoidea spiders karyotyped, only 10 species were examined regarding to the NOR distribution (Král et al., 2006; Araujo et al., 2008, 2020). Additionally, based on the phylogenetic approach, we discussed about the chromosome evolution of Scytodoidea, focusing in the variation of diploid number, types of sex chromosome system and change in the localization of ribosomal genes.
MATERIALS AND METHODS

A sample of 35 specimens was analyzed in this work. The data concerning the number of individuals and the collection localities in Brazil are shown in Table 1. The vouchers were deposited in the arachnid collection of the Instituto Butantan, São Paulo, (IBSP; curator A.D. Brescovit); Coleções Taxonômicas of the Universidade Federal de Minas Gerais, Belo Horizonte (UFMG; curator A.J. Santos), and Coleção de História Natural of the Universidade Federal do Piauí, Florianópolis (CHNUFPI; curator L.S. Carvalho), in Brazil.

The cytological preparations were obtained following the procedures of Araujo et al. (2005). The chromosome slides were stained with 3% Giemsa solution (3% commercial Giemsa solution and 3% phosphate buffer pH 6.8, in distilled water), silver-impregnated (Howell y Black, 1980) to detect the NORs and submitted to FISH with 18S rDNA probes to localize the major ribosomal gene. The morphological classification of chromosomes followed the nomenclature proposed by Levan et al. (1964).

The 18S rDNA probes were obtained by PCR using the DNA of Physocyclus globosus (Taczanowski, 1874) (Pholcidae) and the primers 18S-F 5’ CGAGCGCTTTTATTAGACCA and 18S-R 5’ GGTTCACTACGGAAAACCTT (Forman et al., 2013). Probes were labeled with 11-dUTP–digoxigenin by PCR. The FISH technique was performed according Pinkel et al. (1986). The chromosomal DNA was denatured in 70% formamide for 5 min at 70°C and the hybridization solution was denatured in a thermal cycler for 10 min at 95°C. Probes were detected with anti-digoxigenin antibody conjugated to rhodamine. Chromosome spreads were counterstained with 4′–6–diamidino–2–phenylindole (DAPI) and the slides were mounted with antifading solution. The images were captured using a Zeiss Imager A2 microscope, coupled to a digital camera and the Axio Vision software.

The ancestral condition of diploid number, sex chromosome system and number of rDNA sites was reconstructed in Mesquite (Maddison y Maddison, 2011), using the maximum parsimony approach and the phylogenetic proposal of Wheeler et al. (2017). The chromosome data were obtained from the present study and spider cytogenetic database (Araujo et al., 2021).

Table 1. Sicarius species cytogenetically analyzed in this work, including the number of specimens and collection localities in Brazil. PI=state of Piauí; SE=state of Sergipe; PB=state of Paraíba.

| Species         | Number of individuals | Locality                                                                 |
|-----------------|-----------------------|--------------------------------------------------------------------------|
| Sicarius cariri | 3♂/1♀                 | Parque Nacional da Serra da Capivara (8°49’48.0"S, 42°33’16.0"W), São Raimundo Nonato, PI |
|                 |                       | (7°9’39.4”S, 41°28’2.5”W), Picos, PI                                       |
|                 | 1♀                    | Horto Florestal (4°16’19.0”S, 41°41’18.9”W), Piripiri, PI               |
|                 | 2♂                    | Povoado Saquinho (2°46’2.5”S, 41°48’19.3”W), Ilha Grande do Piauí, PI  |
|                 | 2♀                    | Parque Nacional da Serra das Confusões (8°56’16.9”S, 43°51’48.1”W), Cristino Castro, PI |
|                 | 15♂/1♀                | Parque Municipal Pedra de Castelo (5°12’5.9”S, 41°41’14.0”W), Castelo do Piauí, PI |
| Sicarius ornatus| 2♂                    | Parque Nacional da Serra de Itabaiana (10°44’57.3”S, 37°20’20.1”W), Itabaiana, SE |
| Sicarius tropicus| 3♂                    | Reserva Particular de Patrimônio Natural Fazenda Almas (7°23’16.9”S, 36°48’31.8”W), São José dos Cordeiros, PB |
|                 | 3♂                    | Parque Municipal Pedra de Castelo (5°12’5.9”S, 41°41’14.0”W), Castelo do Piauí, PI |
RESULTS

Chromosome characterization

Mitotic metaphase cells of male and female specimens of *S. cariri* showed the diploid number and sex chromosome system 2n=16+X,Y and 2n=16+X,X,Y, respectively (Fig. 1A-B). All the chromosomes presented metacentric morphology, with exception of submetacentric pair 2. Regarding to the size, the autosomal chromosomes could be classified into three categories: large (pairs 1 and 2), medium (pairs 3 and 4) and small (pairs 5 to 8). In spermatogonial cells, the X and Y sex chromosome were easily identified as unpaired elements, which corresponded to the largest and smallest chromosomes of the karyotype. The X chromosome showed an intermediary size between the 2nd and 3rd autosomal pairs (Fig. 1A).

*Sicarius ornatus* also presented 2n♂=19. In the karyotype, three unpaired chromosomes were identified, which were similar to the sex chromosomes of *S. cariri*. Thus, *S. ornatus* should also display a X,Y sex chromosome system. However, in this species, all autosomal pairs revealed metacentric morphology, the X and Y sex chromosomes were submetacentric and the Y was a tiny acrocentric chromosome (Fig. 1C). The autosomal pair 1 exhibited large size, compared to the medium-sized pairs 2 to 5 and the smallest elements of the karyotype, pairs 6 to 8. The X chromosome presented a similar size to the pair 1, the Y chromosome was larger than the pair 2 and the Y was the smallest chromosome of the karyotype (Fig. 1C).

In *S. tropicus*, the mitotic metaphase cells evidenced 2n♂=20, with two unpaired chromosomes, one large and other small-sized. The karyotype comparison with *S. cariri* and *S. ornatus* and the analysis of meiotic cells permitted us interpreted these unpaired elements as X and Y sex chromosomes. In *S. tropicus*, all chromosomes presented metacentric morphology. The pair 1 was

![Figure 1. Karyotypes of three Sicarius species. A-B. Sicarius cariri with 2n♂=16+X,Y and 2n♀=16+X,X,Y, respectively. C. Sicarius ornatus, 2n♂=16+X,Y. D. Sicarius tropicus, 2n♂=18+XY. In all species, the chromosomes were predominantly metacentrics. Scale bar=5μm.](image)
large-sized, the pairs 2, 3 and 4 medium-sized and the pairs 5 to 9 small-sized. The X and Y sex chromosomes corresponded to the largest and the smallest elements of the karyotype, respectively (Fig. 1D).

The analysis of meiotic cells of *S. cariri* and *S. tropicus* revealed, in the pachytene, autosomal chromosomes completely paired and a single and very small element, interpreted as X chromosome (Fig. 2A-B). For both species, the Y sex chromosomes were not identified in this meiotic substage. Diplotene cells of *S. tropicus* presented 9 autosomal bivalents, with up to two interstitial or terminal chiasmata, and one heteromorphic bivalent, formed by the end-to-end paired XY chromosomes (Fig. 2C). Nuclei in metaphase II of this species showed the haploid sets n=9+X and n=9+Y (Fig. 2D).

Silver-impregnated mitotic metaphase nuclei of the three *Sicarius* species revealed active NORs on the long arm terminal region of pair 1 (Fig. 3A-F). In *S. cariri*, the 18S rDNA sites were located in the long arm terminal region of the X sex chromosome (Fig. 4A). In this species, the incongruence between the results of Ag-NOR and FISH were observed among the cells of a same individual as well as in cells of different specimens. In *S. ornatus* and *S. tropicus*, the ribosomal cistrons occurred only in the long arm terminal region of the 1st autosomal pair (Fig. 4B-E), confirming the results of silver impregnation.
Figure 3. Mitotic metaphase cells of Sicarius species submitted to Giemsa-staining (A, C, E) and silver impregnation (B, D, F) to reveal the nucleolar organizer regions (arrowhead). A–B. Sicarius cariri. C–D. Sicarius ornatus. E–F. Sicarius tropicus. The cells showed in C and D are with incomplete diploid set. Scale bar=5μm.
Chromosome evolution

The maximum parsimony analyses revealed 2n=18–22 as the ancestral autosomal number for Scytodoidea and Sicariidae (Fig. 5A). Overall, the karyotype evolution occurred through independent decreased in autosomal number, with the exception of two species of Scytodidae (*Scytodes fusca* – 30 autosomes and *Scytodes* sp. – 26 chromosomes, without description of the sex chromosome system) and Drymusidae (*Izithunzi capense*). The only exception is an unidentified species of the genus *Scytodes*, in which the sex chromosome system was not described. Within Sicariidae, the only change in the number of X sex chromosome was reported in *S. tropicus* with a XY sex chromosome system. The loss of Y chromosome was recorded only in *L. reclusa* and *L. ruifera*.

Despite the low number of species characterized, the presence of three or four rDNA sites seems to be the ancestral condition for Scytodoidea (Fig. 5D). However, this state was frequently changed during the evolution of this group. In *L. amazonica* and *L. puerto*, these changes involved the increase of the number of major rDNA cistrons while in *Sicarius* species seems to be occurred a decrease in the number of these sites. The analyses also showed that in Ochyroceratidae + Scytodidae and *Sicarius* species, the ancestral rDNA number is lower than those observed in Scytodoidea.
DISCUSSION

The diploid number 2n=19, the X1X2Y sex chromosome system and the chromosomal morphology predominantly metacentric herein observed in S. cariri and S. ornatus are similar to those previously described for S. tropicus and only one species of the genus Loxosceles, L. spinulosa (Král et al., 2006; Araujo et al., 2020). Additionally, the X1X2Y sex chromosome system verified in S. cariri and S. ornatus is the most common in Sicariidae, occurring in 12 out of the 15 species cytogenetically characterized so far (Araujo et al., 2020). The tendency of decreasing of the diploid number verified in some Scytodoidea species is the main mechanism of chromosome evolution for spiders and has been reported in many studies accomplished with related species (Stávale et al., 2010; Araujo et al., 2020; Ávila Herrera et al., 2021). In an elegant cytogenetic work with many Pholcidae spiders, in which data of molecular and paleontological studies were discussed, Ávila Herrera et al. (2021) suggested that the X1X2Y sex chromosome system possesses an ancient origin in spiders and could have arise before the emergence of Araneomorphae lineage.

The karyotype found here for S. tropicus (2n=18+XY) differed from that registered for other population of this same species (2n=16+X1X2Y) (Araujo et al., 2021), and the description of a simple sex chromosome system of the XY type is original for Sicariidae. The high similarity regarding to the size of the Y chromosome among the Sicarius species having the X1X2Y and XY systems indicates that the evolution of the XY system occurred through rearrangements involving only the X chromosome. The XY system probably had origin from the X1X2Y system, in which the ancestral and metacentric X1 and X2 chromosomes were pericentrically inverted, originating subtelo-acrocentric chromosomes, such as those verified in Sicarius sp. (Franco y Andía, 2013). In a subsequent event, the X1 and X2 chromosomes were fused, converting the X1X2Y into a XY system. This hypothesis regarding XY sex chromosome evolution was proposed by Král et al. (2006) and Ávila Herrera et al. (2021), analyzing the behavior of the XY sex chromosomes during the meiosis of Diguetia albolineata (O. Pickard–Cambridge, 1895) (Diguetidae) and Wujigarra sp., (Pholcidae) respectively. In these species as well as in S. tropicus analyzed here, the X and Y chromosomes exhibited only one end-to-end association during prophase I, without the presence of chiasma. The present study in Sicarius species filled in an important gap in the hypothesis of Král et al. (2006)

Figure 5. Chromosome evolution in Scytodoidea spiders obtained after Mesquite analysis. A. Autosomal number. B. Number of X sex chromosome. C. Presence of sex chromosome system including a Y chromosome. D. Number of chromosomes with NOR or rDNA sites.
about the evolution of sex chromosomes systems in basal clades of Araneomorphae, taking into account that the hypothetic X,X,Y system with subtelo-acrocentric X and X chromosomes was exclusively observed in Sicarius sp. (Franco y Andía, 2013).

The differences related to diploid number and sex chromosome system observed in S. tropicus (present study; Araujo et al., 2021) may represent an interpopulationual variation, indicating that the karyotype 2n=18+XY is not well established in all populations of this species or it had an independent origin in the populations analyzed by us. Magalhaes et al. (2014), performing a phylogeographic study in S. cariri, using sequence data of nuclear and mitochondrial genes, revealed highly structured populations, which might be evolving independently. It is possible that S. tropicus populations are also strongly structured geographically, which could explain the differences in the karyotypes. Alternatively, the specimens initially described by Araujo et al. (2021) as S. tropicus could correspond to another species of the genus Sicarius, considering that the cytogenetic study accomplished by Araujo preceded the taxonomical and systematic revision of the genus Sicarius (Magalhaes et al., 2013, 2017).

The supposed stability of number and localization of NORs in spiders has knocked down with the increase of cytogenetic studies. In an analysis of NORs in 30 Pholcidae spiders, Ávila Herrera et al. (2021) revealed a great diversity of number of this site, which can occur in autosomes and/or X sex chromosome. The results obtained herein using FISH with rDNA probe only in three Sicarius species revealed the presence of ribosomal cistrons in autosomes (S. ornatus and S. tropicus) and X, sex chromosome (S. cariri). It is interesting to emphasize that this difference of localization of rDNA in autosome/sex chromosome occurs in species with similar karyotype characteristics, indicating that the changes involving the ribosomal genes can be independent of the differentiation of the sex chromosome system. In S. cariri, the localization of active NORs and 18S rDNA showed incongruous data, considering that the silver-impregnated regions were visualized on the terminal sites of the 1st autosomal pair, such as in S. ornatus and S. tropicus, but the FISH evidenced a bright signal in the terminal region of the X, sex chromosome. Therefore, in S. cariri the silver impregnation might have evidenced false Ag-NORs, taking into account that this technique reveals the NORs indirectly. This occurs due to the affinity of the silver nitrate by acidic proteins associated with the rRNAs or heterochromatic regions (Sanchez et al., 1995; Lorite et al., 1997; Dobigny et al., 2002; Kasahara, 2009; Kavalco et al., 2009; Reis et al., 2012). On the other hand, the impregnation of the terminal region of pair 1 of S. cariri, which is certainly carrier of 18S rDNA genes in the two other closely related species, S. ornatus and S. tropicus, might suggest the presence of cryptic NORs in S. cariri, such as those reported by Cabrero y Camacho (2008) in some grasshopper species. The silver impregnation on pair 1 of S. cariri can represent a vestigial locus of rDNA gene for this species, which was translocated to the X, sex chromosome; this vestigial rDNA is very small to be detected by the FISH technique but it retains its transcriptional activity.

In conclusion, the data shown herein expanded the knowledge of the karyotype diversity already registered for sicariid spiders. Moreover, we identified an intriguing variation when the results of Ag-NOR and FISH were compared. Therefore, the Scytodidae spiders are not only interesting for cytogenetic studies due to the variability in the sex chromosome system, but also because they are suitable for investigating karyotype evolution in spiders and its relationship to the distribution and activity of rDNA genes.

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