A karyotype comparison between two species of bordered plant bugs (Hemiptera, Heteroptera, Largidae) by conventional chromosome staining, C-banding and rDNA-FISH

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Abstract
A cytogenetic characterization, including heterochromatin content, and the analysis of the location of rDNA genes, was performed in Largus fasciatus Blanchard, 1843 and L. rufipennis Laporte, 1832. Mitotic and meiotic analyses revealed the same diploid chromosome number 2n = 12 + X0/XX (male/female). Heterochromatin content, very scarce in both species, revealed C-blocks at both ends of autosomes and X chromosome. The most remarkable cytological feature observed between both species was the different chromosome position of the NORs. This analysis allowed us to use the NORs as a cytological marker because two clusters of rDNA genes are located at one end of one pair of autosomes in L. fasciatus, whereas a single rDNA cluster is located at one terminal region of the X chromosome in L. rufipennis. Taking into account our results and previous data obtained in other heteropteran species, the conventional staining, chromosome bandings, and rDNA-FISH provide important chromosome markers for cytotaxonomy, karyotype evolution, and chromosome structure and organization studies.

Keywords
Largus, Heteroptera, C-banding, rDNA-FISH, holokinetic chromosomes, karyotype comparison

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Introduction

All species of Hemiptera studied so far present holokinetic chromosomes (i.e. without a primary constriction). Kinetic activity is restricted to the chromosome ends and the chromosomes can be regarded as telokinetic during male meiosis, but holokinetic activity is recognized in mitosis. Meiotic behaviour is slightly different depending on whether we are dealing with autosomal bivalents, sex chromosomes, m chromosomes or autosomal univalents. As a rule, autosomal bivalents are chiasmatic and segregate reductionally, whereas sex and m chromosomes are achiasmatic and divide equationally at first male meiotic division. Besides, sex chromosomes do not present a defined position at metaphases I and II. Several reports on C-positive heterochromatin in true bugs showed that C-bands are terminally located (Ueshima 1979, Manna 1984, Papeschi and Bressa 2006).

At present, the seven species cytogenetically studied of Largidae possess a low diploid chromosome number, ranging between 11 and 17 autosomes, a X0/XX sex chromosome system (male/female), except for one species, and a pair of m chromosomes, excluding the genus Largus Hahn, 1831 (Ueshima 1979, Manna 1984, Manna et al. 1985, Mola and Papeschi 1993, Bressa et al. 2005).

The genus Largus comprises 61 taxonomically described species and most of them are distributed in America, where its geographic distribution ranges from the north of the United States to the south of Argentina. Although they are more diverse and abundant in tropical and subtropical areas, in Argentina there are only seven species recorded (Melo and Dellapé 2013, Rosas and Brailovsky 2016). At cytogenetic level, Largus rufipennis Laporte, 1832 is the only species analysed to this date, using only conventional methods (Mola and Papeschi 1993, Bressa et al. 1998, 2005). It possesses a male diploid number of 2n = 13 = 12 + X0 and very large chromosomes. The partial karyotype analyses allowed detecting several Argentinean populations with different number of autosomal univalents, variable chiasma frequency, and the presence/absence of B chromosomes.

The main aim of this study was to describe the karyotype of L. fasciatus Blanchard, 1843 and examine the structure of its holokinetic chromosomes by means of C-banding and fluorescent in situ hybridization (FISH) with 18S rDNA probes. Using these data we performed a detailed comparison of the content and distribution of constitutive heterochromatin and the location of rDNA gene clusters between L. fasciatus and L. rufipennis collected from several fields in Argentina.

Material and methods

Insects

Adults and nymphs of L. fasciatus (12 males and 2 females) and L. rufipennis (6 males) were collected from 1995 to 2009 in several fields from Buenos Aires and Entre Ríos in Argentina (Table 1). The collected adults were taxonomically determined by María del
Table 1. Species, locality, geographical coordinates, and number of adults’ collected and examined of *Largus* for chromosomal analyses discriminated by gender.

| Species          | City and Province from Argentina       | Coordinates (DMS)          | N° of individuals |
|------------------|---------------------------------------|---------------------------|------------------|
| *Largus fasciatus* | Tornquist, Buenos Aires               | 38°05’45”S, 62°13’25”W   | 11 males, 2 females |
|                  | Isla Martín García, Buenos Aires      | 34°11’03”S, 58°14’58”W   | 1 male           |
|                  | Sierra de los Padres, Buenos Aires    | 37°56’50”S, 57°46’40”W   | 3 males          |
|                  | Santa Catalina, Buenos Aires          | 34°46’11”S, 58°27’19”W   | 1 male           |
|                  | Ceibas, Entre Ríos                    | 33°30’02”S, 58°48’16”W   | 1 male           |

Carmen Coscarón (Facultad de Ciencias Naturales y Museo, Universidad Nacional de La Plata) and specimens were deposited in the Museo Argentino de Ciencias Naturales Bernardino Rivadavia (MACN, Buenos Aires, Argentina).

**Chromosome preparations**

The captured specimens were swollen in freshly prepared fixative (methanol: glacial acetic acid, 3:1). In the laboratory their gonads were dissected out in 70% ethanol. Cells of gonads were dissociated in a drop of 45% acetic acid, prepared by the squash technique, and stored at -20°C until use. Chromosome preparations were removed from freezer, dehydrated in an ethanol series, and air-dried. For mitotic and meiotic analyses, the chromosome preparations were stained with 5% Giemsa solution following conventional procedures. Heterochromatin content and distribution were analysed by means of C-bands according to Papeschi (1988), and the pre-treated slides were stained with 4′6-diamidino-2-phenylindole (DAPI; Fluka BioChemika, Sigma Aldrich Production GmbH, Buchs, Switzerland) for a better resolution (Poggio et al. 2011).

**Fluorescence in situ hybridization**

Spread chromosome preparations were made in a drop of 60% acetic acid with the help of tungsten needles and the spreading on the slide was performed using a heating plate at 45°C as described in Traut (1976). Unlabelled 18S ribosomal DNA (rDNA) probes were generated by polymerase chain reaction (PCR) using universal arthropod primers: forward 5′-CCTGAGAAACGCGCTACCACATC-3′ and reverse 5′-GAGTCTCCTCGTTCTCAGGA-3′ (Whiting 2002). Total genomic DNA of *Dysdercus albofasciatus* Berg, 1878, obtained by standard phenol-chloroform-isoamyl alcohol extraction, was used as a template. PCR was done following the profile described in Fuková et al. (2005). The PCR product showed a single band of about 1,000 bp on a 1% agarose gel. The band was excised from the gel and purified by using a QIAquick Gel Extraction Kit (Quiagen GmbH, Hilden, Germany). The 18S rDNA fragment was re-amplified by PCR and then labeled with biotin 14-dATP by nick translation using a BioNick Labeling System (Invitrogen, Life Technologies Inc.,
San Diego, CA, USA). FISH with a biotinylated 18S rDNA probe was carried out following the procedure described in Sahara et al. (1999) with several modifications described by Fuková et al. (2005) and Bressa et al. (2009).

**Microscopy, photographs and image processing**

Preparations were observed under high power magnification using a Leica DMLB epi-fluorescence microscope equipped with a Leica DFC350 FX CCD camera and Leica IM50 software, Version 4.0 (Leica Microsystems Imaging Solutions Ltd., Cambridge, UK). Black-and-white images of chromosomes were recorded separately for each fluorescent dye with the CCD camera. Images were pseudo-coloured (light blue for DAPI and red for Cy3) and processed with Adobe Photoshop CS6 Version 6.1 (1999–2012) software.

**Results**

Based on the observation of metaphase I autosomal bivalents (AA) and the identification of the sex univalent we described the male karyotype of *L. rufipennis* as 2n = 6AA + X0 (see Mola and Papeschi 1993), and the chromosome complement of *L. fasciatus* as 2n = 6AA + X0/XX (male/female sex chromosomes) (Fig. 1). In both species, the autosomes decrease gradually in size and the X chromosome is the smallest of the complement having an equal or nearly equal diameter in all directions (Fig. 1). From diakinesis onwards, the X is negatively heteropycnotic (Fig. 1a, b, d, e). The X chromosome in *L. rufipennis* is slightly longer than in *L. fasciatus* (Fig. 1b, e). At metaphases I and II, the autosomes are arranged forming a ring and with the X located outside of it (Fig. 1b–e).

The C-banding pattern in *L. rufipennis* and *L. fasciatus* showed discrete C-positive bands terminally located in all autosomes and the X chromosomes, which were observed in all stages of mitosis and meiosis (Fig. 2).

FISH experiments with the 18S rDNA probe revealed differences in the location of the probe signals between both species analysed (Fig. 3). In *L. rufipennis* a cluster of rDNA genes was located at one end of the X chromosome (Fig. 3a–b), whereas in *L. fasciatus* the hybridization signals were located at a subterminal position in an autosomal bivalent (Fig. 3c–d).

**Discussion**

*Largus rufipennis* and *L. fasciatus*, the two species herein analysed, showed similar karyotypes composed of six pairs of autosomes, a simple sex chromosome system (X0/XX), and the same location and distribution of constitutive heterochromatin. The main cytogenetic difference between both species was detected in the location of the rDNA
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**Figure 1.** Karyotypes of *L. rufipennis* (a–b) and *L. fasciatus* (c–e) stained with 5% Giemsa. a diakinesis b metaphase I c oogonial metaphase (2n = 12 + XX) d diakinesis e metaphase I. X = sex chromosome. Scale bar: 10 μm.

clusters. Two signals were located at a subterminal position of an autosomal bivalent of *L. fasciatus* but only one signal was observed at one end of the X chromosome of *L. rufipennis*. Taking into account the data on the chromosomal location of rDNA clusters in other heteropteran species along with our results, the NORs chromosome location varies among several congeneric species, i.e. *Belostoma* Leach, 1815 (Nepomorpha), *Triatoma* Laporte, 1832, *Panstrongylus* Berg, 1879, *Rhodnius* Stål, 1859 (Cimicomorpha), and *Dysdercus* Guérin-Méneville 1831 (Pentatomomorpha) (Papeschi and Bressa 2006, Morielle-Souza and Azeredo-Oliveira 2007, Bressa et al. 2009, Chirino et al. 2013, Pita et al. 2013, Chirino and Bressa 2014, Grozova et al. 2014, Panzera et al. 2014), and also among different species of Tingidae (Cimicomorpha) and several species belonging to different families of Pentatomomorpha (Bardella et al. 2013, 2016, Golub et al. 2015, 2016). The analysis in a wide number of species shows that 5S, 18S, and 45S rDNA remain mainly among the autosomes, although in some species the NORs are located in the sex and m chromosomes. This might be due to the fact that NORs can be easily translocated to other chromosomes changing their number and
position. Consequently, the number and location of rDNA loci (determined by FISH and/or Ag-NOR banding) constitutes an important chromosome marker, which can be useful for studies on cytotaxonomy, karyotype evolution, and chromosome structure and organization for heteropteran species. Therefore, rearrangements involving rDNA-repositioning seem to be involved in the species’ evolutionary history, indicating a particular genome dynamics for this marker.

From the cytogenetic point of view, Largidae is an interesting heteropteran family because of its low diploid chromosome number and the large chromosome size observed in most of the species (Ueshima 1979, Manna 1984, Manna et al. 1985, this study). The six karyologically analysed species of the subfamily Larginae, *Largus* and *Macrochraia* Guérin-Méneville, 1835, are characterized by the absence of an m chromosome pair, the possession of an X0/XX sex chromosome mechanism, and a number of autosomes that varies between 10 and 14. Conversely, all the studied species belonging to Physopeltinae possess 12 autosomes, two m chromosomes, and different sex chro-
mosomes systems (X0 or X1X2Y) (see references in Papeschi and Bressa 2006). Based on the presence of a Y chromosome in very primitive heteropteran species, Nokkala and Nokkala (1983, 1984) and Grozeva and Nokkala (1996) suggested that the X0 system is a derived condition from the ancestral XY that is present in the majority of the species cytogenetically analysed. In Larginae, the X0 sex chromosome system most probably originated through the loss of the Y chromosome. The finding of a pair of m chromosomes in three species of Physopeltinae (Ueshima 1979, Manna et al. 1985) led us to suggest that this pair of chromosomes might be involved in the ancestral karyotype of the family Largidae. Then, the absence of m chromosomes and the presence of sex chromosome system X0 in species of Larginae could be considered as derived characters, which arose during karyotype evolution.

The use of different cytogenetic techniques will be very useful in further integrative studies because a group-level taxonomy followed by a reliable association among different data sets is fundamental to allow a more precise evaluation of the processes involved in the karyotype evolution and the interrelationships among different species.

Figure 3. Location of rDNA genes in chromosomes of L. rufipennis (a–b) and L. fasciatus (c–d) by FISH with 18S rDNA probes (red signals). Chromosomes were counterstained with DAPI (blue). a metaphase I b telophase II c spermatogonial prometaphase d diakinesis. X = sex chromosome. Scale bar: 10 μm.
Conclusions

Taking into account the data on the number and location of rDNA clusters in *L. ru- fipennis* and *L. fasciatus*, we can observe two different patterns of rDNA distribution. As a result, the rDNA clusters revealed by rDNA-FISH are very useful tools for the study of the karyotype structure and chromosome evolution in groups with holokinetic chromosomes due to it can contribute to understand the karyotype evolution and taxonomic relationships among several taxa.

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References

Bardella VB, Fernandes T, Vanzela ALL (2013) The conservation of number and location of 18S sites indicates the relative stability of rDNA in species of Pentatomomorpha (Heterop- tera). Genome 56(7): 425–429. https://doi.org/10.1139/gen-2013-0140

Bardella VB, Fernandes JAM, Cabral-de-Mello DC (2016) Chromosomal evolutionary dy-namics of four multigene families in Coreidae and Pentatomidae (Heteroptera) true bugs. Molecular Genetics and Genomics 291(5): 1919–1925. https://doi.org/10.1007/s00438- 016-1229-5

Bressa MJ, Papeschi AG, Mola LM, Larramendy ML (1998) Meiotic studies in *Largus rufipen- nis* (Castelnau) (Largidae, Heteroptera). II. Reciprocal translocation heterizygosity. Caryolo- logia 51: 253–264. doi:10.1080/00087114.1998.10797417

Bressa MJ, Larramendy ML, Papeschi AG (2005) Heterochromatin characterization in five species of Heteroptera. Genetica 124(2–3): 307–317. https://doi.org/10.1007/s10709-005-4524-3

Bressa MJ, Papeschi AG, Vítková M, Kubíčková S, Fuková I, Pigozzi MI, Marec F (2009) Sex chromosome evolution in cotton stainers of the genus *Dysdercus* (Heteroptera: Pyrrhocoridae). Cytogenetic and Genome Research 125(4): 292–305. https://doi. org/10.1159/000235936

Chirino MG, Papeschi AG, Bressa MJ (2013) The significance of cytogenetics for the study of karyotype evolution and taxonomy of water bugs (Heteroptera, Belostomatidae) native to Argentina. Comparative Cytogenetics 7(2): 9–27. https://doi.org/10.3897/CompCytoge-n.v7i2.4462

Chirino MG, Bressa MJ (2014) Karyotype evolution in progress: a new diploid number in *Be- lostoma candidulum* (Heteroptera: Belostomatidae) from Argentina leading to new insights
into its ecology and evolution. European Journal of Entomology 111(2): 165–174. https://doi.org/10.14411/eje.2014.027

Fuková I, Nguyen P, Marec F (2005) Codling moth cytotgenetics: karyotype, chromosomal location of rDNA, and molecular differentiation of sex chromosomes. Genome 48(6): 1083–1092. https://doi.org/10.1139/g05-063

Golub NV, Golub VB, Kuznetsova VG (2015) Variability of 18rDNA loci in four lace bug species (Hemiptera, Tingidae) with the same chromosome number. Comparative Cytogenetics 9(4): 513–522. https://doi.org/10.3897/CompCytogen.v9i4.5376

Golub NV, Golub VB, Kuznetsova VG (2016) Further evidence for the variability of the 18S rDNA loci in the family Tingidae (Hemiptera, Heteroptera). Comparative Cytogenetics 10(4): 517–528. https://doi.org/10.3897/CompCytogen.v10i4.9631

Grozeva S, Nokkala S (1996) Chromosomes and their meiotic behavior in two families of the primitive infraorden Dipsocoromorpha (Heteroptera). Hereditas 125(1): 31–36. https://doi.org/10.1111/j.1601-5223.1996.t01-1-00031.x

Grozeva S, Anokhin B, Kuznetsova VG (2014) Chapter 8. Recent advances in cytogenetics of bed bugs: FISH mapping of the 18S rDNA and TTAGG telomeric loci in Cimex lectularius Fabricius, 1803 (Hemiptera, Heteroptera, Cimicidae). In: Sharachov I (Ed.) Protocols for chromosome mapping of arthropod genomes, CRCPress, Taylor & Francis, Boca Raton, Florida, 283–322.

Manna GK, (1984) Chromosomes in evolution in Heteroptera. In: Sharma AK, Sharma A (Eds) Chromosomes in evolution of eukaryotic groups. CRCPress, Boca Raton, Florida, 189–225.

Manna GK, Ueshima N, Dey SK, Ded-Mallick S (1985) Market Sex chromosomal variations between Indian and a Japanese species of Physopelta (Largidae, Heteroptera). Cytologia 50: 621–630. https://doi.org/10.1508/cytologia.50.621

Melo MC, Dellapé PM (2013) Catalogue of the Pyrrhocoroidea (Hemiptera: Heteroptera) from Argentina. Revista de la Sociedad Entomológica Argentina 72(1–2): 55–74. http://www.scielo.org.ar/pdf/resea/v72n1-2/v72n1-2a06.pdf [In Spanish]

Mola LM, Papeschi AG (1993) Meiotic studies in Largus rufipennis (Castelnau) (Largidae, Heteroptera): frequency and behaviour of ring bivalents, univalents and B chromosomes. Heredity 71: 33–40. doi:10.1038/hdy.1993.104

Morielle-Souza A, Azeredo-Oliveira MTV (2007) Differential characterization of holocentric chromosomes in triatomines (Heteroptera, Triatominae) using different staining techniques and fluorescent in situ hybridization. Genetics and molecular research 6(3): 713–720. https://science.report/pub/16044094

Nokkala S, Nokkala C (1983) Achiasmatic male meiosis in two species of Saldula (Saldidae, Hemiptera). Hereditas 99(1): 131–134. https://doi.org/10.1111/j.1601-5223.1983.tb00737.x

Nokkala S, Nokkala C (1984) The occurrence of the X0 sex chromosome system in Dictytonota tricornis (Schr.) (Tingidae, Hemiptera) and its significance for concepts of sex chromosome system evolution in Heteroptera. Hereditas 100(2): 299–301. https://doi.org/10.1111/j.1601-5223.1984.tb00130.x
Panzera F, Ferreiro MJ, Pita S, Calleros L, Pérez R, Basmadjián Y, Guevara Y, Frédérique Brenière S, Panzera Y (2014) Evolutionary and dispersal history of *Triatoma infestans*, main vector of Chagas disease, by chromosomal markers. Infection, Genetics and Evolution 27: 105-113. https://doi.org/10.1016/j.meegid.2014.07.006

Papeschi AG (1988) C-banding and DNA content in three species of *Belostoma* (Heteroptera) with large differences in chromosome size and number. Genetica 76: 43–51. https://doi.org/10.1007/BF00126009

Papeschi AG, Bressa MJ (2006) Evolutionary cytogenetics in Heteroptera. Journal of Biological Research 5: 3–21. http://www.jbr.gr/papers20061/01-Alba.pdf

Pita S, Panzera F, Ferrandis I, Galvao C, Gomez-Palacio A, Panzera Y (2013) Chromosomal divergence and evolutionary inferences in Rhodniini based on the chromosomal location of ribosomal genes. Memórias do Instituto Oswaldo Cruz 108(3): 376–382. https://doi.org/10.1590/S0074-02762013000300017

Poggio MG, Bressa MJ, Papeschi AG (2011) Male meiosis, heterochromatin characterization and chromosomal location of rDNA in *Microtomus lunifer* (Berg, 1900) (Hemiptera: Reduviidae: Hammacerinae). Comparative Cytogenetics 5(1): 1–22. https://doi.org/10.3897/compcytogen.v5i1.1143

Rosas C, Brailovsky H (2016) Revision of the genus *Largus* (Hemiptera: Heteroptera: Largidae) from Mexico. Revista Mexicana de Biodiversidad 87: 347–375. https://doi.org/10.1016/j.rmb.2016.05.001

Sahara K, Marec F, Traut W (1999) TTAGG telomeric repeats in chromosomes of some insects and other arthropods. Chromosome Research 7: 449–460. https://doi.org/10.1023/A:1009297729547

T raut W (1976) Pachytene mapping in the female silkworm *Bombyx mori* L. (Lepidoptera). Chromosoma 58: 275–284. https://doi.org/10.1007/BF00292094

Ueshima N (1979) Hemiptera II: Heteroptera. Animal Cytogenetics 3 Insecta 6. Gebrüder Bornträger, Stuttgart, 117 pp.

Whiting MF (2002) Phylogeny of the holometabolous insect orders: molecular evidence. Zoologica Scripta 31(1): 3–15. doi:10.1046/j.0300-3256.2001.00093.x