The early evolutionary history of neo-sex chromosomes in Neotropical grasshoppers, Boliviacris noroestensis (Orthoptera: Acrididae: Melanoplinae)

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Abstract. Neo-sex chromosomes are an important component of chromosome variation in Orthoptera, particularly South American Melanoplinae species, which have proven to be outstanding experimental model systems to study the mechanism of sex chromosome evolution in this group of insects. In terms of their origin, most derived sex chromosome mechanisms involve a Robertsonian fusion (i.e. translocation) between the ancestral X chromosome and an autosome. In the grasshopper, Boliviacris noroestensis Ronderos & Cigliano (1990) (Orthoptera: Acrididae: Melanoplinae), our results point to a small degree of differentiation (conserved homology between the XR arm and the neo-Y) of the neo-XY chromosomes, which may be of recent evolutionary origin. However, a simple centric fusion model does not explain their origin, mainly because of the observed reduction in the fundamental number (FN) of arms. We propose two models which, we hope, clarify the genesis of B. noroestensis neo-sex chromosomes. Records of karyotype variation in related species due to multiple rearrangements support our models. We propose a possible adaptive advantage for neo-sex chromosome carriers, such changes perhaps representing the primary force that increases their frequency within natural populations compared with non-fused translocated forms, and occurring without apparent detriment to the microevolutionary forces that may also act, at least at the beginning of the evolutionary history of individuals bearing such neo-sex chromosomes.

INTRODUCTION

Neo-sex chromosomes are an important component of chromosome variation in Orthoptera (grasshoppers and crickets) and especially in the superfamily Acridoidea. South American melanopline grasshoppers have proven to be an ideal model system for studies of karyotypic evolution in this globally-widely spread and often economically-important group of insects; thus most cytogenetic studies have been performed, and most cases of derived sex chromosome changes described, using this system (Hewitt, 1979; Mesa et al., 1982; Castillo et al., 2010a, b). During the origin of neo-sex chromosomes, Robertsonian (Rb) fusions are always involved, a condition which can be exploited to infer stages in the Y chromosome degeneration process (Charlesworth, 2002; Charlesworth et al., 2005; Bachtrog, 2013). South American melanoplines are of interest in this respect due to a high frequency of heteromorphic species when compared with species of the same subfamily from the rest of the world (Castillo et al., 2010b). Moreover, this subfamily also includes a large number of species which display extreme reductions in chromosome number, mainly due to the occurrence of centric fusions; however, other rearrangements have also played a significant role in karyotype re-structuring (Mesa et al., 1982; Bidau, 1990; Bidau & Martí, 1995; Martí & Bidau, 1995; Colombo et al., 2005; Castillo et al., 2010b).

Robertsonian X-A fusions have occurred several times in the evolutionary history of the Melanoplinae (Mesa et al., 1982, 2001; Bidau & Martí, 2001; Colombo et al., 2005; Castillo et al., 2010b). Indeed, not only do they show a high frequency of derived systems, but also the observed neo-XY chromosome systems fall within a continuum of evolutionary scenarios showing at one end, a small degree of differentiation where the XR (fused autosome) arm of the neo-X still conserves high homology with the neo-Y; on the other end there is an almost complete loss of homology between the neo-Y and XR. This is evidenced by an extremely reduced synaptic zone and eventually XR heterochromatinization (White, 1973; Hewitt, 1979; Castillo et al., 2010a). Recent work on melanopline species using C-f-1 DNA fraction mapping point to a relatively restricted spreading of this repetitive DNA in neo-sex chromosomes, which contrasts with the repetitive DNA accumulation expected after recombination restriction (Palacios-Gimenez et al., 2013). Research also suggests different accumulation/diversification patterns of repetitive DNAs of neo-Y chromosomes in related species (Palacios-Gimenez et al., 2013); such empirical data is evidence for the loss of selection pressure in chromosomal regions where recombinatio-
tion is abolished, leading to a high rate of genetic diversification (Palacios-Giménez et al., 2013). Between these two extreme cases, a continuum of neo-sex chromosome evolutionary stages and those pertaining when the process is apparently complete are found (Mesa, 1962a, b; Sáez, 1963; Díaz & Sáez, 1968; Sáez & Pérez-Mosquera, 1977; Cardoso & Dutra, 1979; Mesa et al., 1982; Martí & Bidau, 2001; Carbonell & Mesa, 2006). In general terms, there are few records of recently established neo-sex chromosomes, probably due to the fact that the new fixed system has, in the initial stage at least, an accelerate process of differentiation. According to Castillo et al. (2010b), a phylogenetically-distant group (Ommexechidae) includes two cases, namely South American Tetrisochecephalus willemsei Gurney & Liebermann and Neuquenina fictor (Rehn) (both Orthoptera: Acridoidea: Ommexechidae), where chiasmata analyses evidenced more than four crossing-over events between the XR and the neo-Y (Mesa, 1961). In the case of Melanoplinae, spontaneous neo-XY mutants are also expected to behave this way, as it was observed in Leiotettix sanguineus Bruner (Orthoptera: Acridoidea: Acrididae) (Mesa & Mesa, 1967; Castillo et al., 2010b) but no records of established systems with this characteristics were found.

In the last years cytogenetic information encompassing the origin, structure and meiotic behaviour of melanopline neo-sex chromosomes has increased notably (Mesa et al., 1982; Castillo et al., 2010a, b; Palacios-Giménez et al., 2013). In general terms, the information provided by recent work has begun to clarify some questions concerning the mechanisms involved in the origin of these sex chromosome changes, but different outcomes could be inferred according to several recent papers (Kaiser & Bachtrog, 2010; Bidau et al., 2011; Palacios-Giménez et al., 2013). Besides, researchers also extrapolate theoretical and empirical data from other biological models in order to shed light on how sex could be determined in Orthoptera (White, 1973; Pannell & Pujol, 2009; Kaiser & Bachtrog, 2010; Bidau et al., 2011).

Within the framework of derived sex chromosomes in recent and advanced evolutionary stages, the main objective of the present study is to describe the karyotype of Boliviacris noroestensis Ronderos & Cigliano (1990) and to discuss the origin of its sex chromosome system in the context of Neotropical Melanoplinae neo-sex chromosomes.

**MATERIAL AND METHODS**

Male and female adult of B. noroestensis grasshoppers were sampled from two localities: Sauzalito, Chaco in Argentina (n = 2♂) (24.424°S, 61.683°W, 170 m a.s.l.) and Filadelfia, Boquerón in Paraguay (n = 5♂; n = 8♀) (22.573°S, 59.879°W, 132 m a.s.l.) from 2008 to 2012. Voucher specimens were deposited in the collection of the Laboratorio de Genética Evolutiva of the Facultad de Ciencias Exactas Quimicas y Naturales, IBS, CONICET-UNAM.

Male meiotic preparations were performed by squashing testes follicles in ferric hematoxylin. Mitotic metaphase chromosomes from female gastric caeca were obtained following the procedure described by Castillo et al. (2011).

Silver staining of kinetochores and chromatid cores were performed according to the procedure of Rufas (1985). Briefly, air-dried male meiotic preparations were incubated in 2 × SSC at 60°C for 10 min and stained with 50% AgNO 3 in dH2O (pH adjusted to 3.5 with formic acid). Microscopic observation of silver stained preparations involved bright field and Nomarski interference optics. C-banding was performed following the protocol of Sumner (1972), with modifications.

Chiasmata were scored at metaphase I with ten cells from each of seven males, classified following inspection as proximal (P), interstitial (I) and distal (D), corresponding to their presence in the first (para-centromeric), second, or third portion of the chromosome arm, respectively, when divided into three equal regions. We used White’s (1973) terminology to refer neo-sex chromosomes.

**RESULTS**

Boliviacris noroestensis showed a karyotype 2n = 20♂/20♀, FN = 21♂/22♀, with all autosomes telocentric and a neo-XY sex-chromosome mechanism. The male karyotype had nine autosomal pairs grouped in three large (L1–L3), three medium-sized (M4–M6), and three small (S7–S9), plus a heteromorphic sex chromosome pair (Fig. 1a, b). As is usual within some melanopline groups, the species appeared to show an S7 megameric autosomal bivalent (Fig. 2b).

All chromosomes in female mitotic metaphase (Fig. 1c) showed small pericentromeric heterochromatic C-positive blocks (Fig. 1d). Furthermore, we found evidence for variations in the distribution of distal C positive blocks. However, a constant pattern was observed in L2, M5 and S7 pairs, showing the same size of pericentromeric blocks; M4 was polymorphic for a prominent paracentromeric C positive block (Fig. 1d).

Concerning the morphology and structure of sex chromosomes, they comprise a submetacentric neo-X and a telocentric neo-Y, the latter being distinguished from autosomes in mitosis due to its large size (Fig. 1a). Also, in
metaphase I we observed that the XR arm is twice the size of the XL arm (Fig. 2c).

In all specimens analyzed, the XL arm showed the typical allocyclic behaviour of the acridoid X chromosome during late pachytene and diplotene (Fig. 2a, b). In addition, using conventional staining we observed a similar condensation pattern in the neo-Y proximal-interstitial region, while the XR and the interstitial-distal region of the neo-Y still conserved complete homology as well as euchromatic behaviour (Fig. 2b). However, at the diplotene stage, we identified the proximal-interstitial region using C-banding and C-banding combined with DAPI staining (Fig. 3a, b).

In males, pachytene nuclei revealed an incomplete synaptic configuration between the interstitial-distal region of the XR arm and the proximal-interstitial region of the neo-Y (Fig. 2a). In metaphase I neo-X and neo-Y centromeres were distantly localized from the pairing region, thus two configurations could be observed in this phase. In addition to a high frequency of distal-interstitial chiasmata (78%) we also observed interstitial chiasmata (22%) between neo-X and neo-Y in all the cells analyzed (n = 70) (Fig. 2c).

In female mitotic metaphases and male diplotenes, the neo-X centromeric region was identified by C-banding (Figs 1d, 3a). Additionally the proximal-interstitial region of the neo-Y was found to display a large heterochromatic block (Fig. 3a, b).

Upon performing the scaffold technique with silver impregnation, neo-XY chromosome kinetochores and the fibrous network of non-histone proteins (scaffolds) were observed at metaphase I. The XL arm of the neo-X showed the typical interchromatidic axis of the ancestral X chromosome without any particular structure, as with those observed in differentiated neo-sex chromosome systems [e.g. Ronderosia bergi (Stål) (Orthoptera: Acrididae: melanoplinae)] (Castillo et al., 2010a). The XR and the neo-Y showed a typical autosomal scaffold configuration (Fig. 2c).

**DISCUSSION**

The occurrence of neo-sex chromosomes has been well documented in Orthoptera and in other insect orders with X0/XX systems, such as Coleoptera (Viturri et al., 2003) and Lepidoptera with ZW/ZZ systems (i.e. with heterogametic females) (White, 1973; Hewitt, 1979; Traut, 1999; Traut et al., 2007; Castillo et al., 2010b). While the Z0/ZZ condition is ancestral, present in all basal lineages of Lepidoptera, cases of Z0/ZZ among more “advanced” insect species are sporadic and considered to be due to secondary loss of the W chromosome (Sahara et al., 2012). As a matter of fact in some cases, X0/XX systems are thought to be the last stages of sex chromosome differentiation. However, they are not as such “dead ends” of sex chromosome evolution. Empirical evidence of chromosome
rearrangement, like autosome translocations, creates a neo-Y as in Heteroptera (Jacobs, 2004; Bressa et al., 2009) or neo-W as in Lepidoptera (Traut & Marec, 1997; Traut et al., 2007). From then on, a new cycle of sex chromosome differentiation is found to begin (Traut, 1999; Traut et al., 2007). In Orthoptera, the most accepted hypothesis for the origin of most neo-chromosome systems is a centric fusion between a standard telocentric X and a telocentric autosome, producing the neo-X chromosome with the homologue (unfused autosome), becoming then the neo-Y (Mesa, 1962a,b; Sáez, 1963; Díaz & Sáez, 1968; Sáez & Pérez-Mosquera, 1977; Cardoso & Dutra, 1979; Mesa et al., 1982; Martí & Bidau, 2001; Carbonell & Mesa, 2006; Castillo et al., 2010a,b). Additionally, there are some records where more than one re-structuring event is involved in their origin [e.g. *Dichroplus viittatus* Bruner (Orthoptera: Acrididae: Melanoplinae) and *R. bergi*] (Bidau & Martí, 2001; Carbonell & Mesa, 2006; Castillo et al., 2010a).

The neo-sex chromosome system spectrum seen in Neotropical South American grasshoppers is wide and only a few of such events have been described in the initial step of the differentiation process, including here in the case of *B. noroestensis*. In three species examined, a recent replacement of the X0-XX sex chromosome system took place (*N. fictor, T. willemsei, L. sanguineus*) (Mesa, 1961; Mesa et al., 2001). Recently arisen neo-sex chromosomes will be initially represented by a pair of sex chromosomes, which still conserves extensive homology, synapic ability and the chance of free recombination along the fused autosome (XR) and its homologue, now the neo-Y (Castillo et al., 2010a).

Based on our observations, the single centric fusion model cannot explain neo-sex chromosome origin of *B. noroestensis*. In the first place, the autosome size involved in the fusion is twice the length of a typical L1 pair. Besides, if a single X-A Rb fusion was involved, the ancestral FN = 23/24 should be maintained. Our results revealed not only a reduction in the chromosome number but also in the number (FN) of arms. Furthermore, we observed a sub-metacentric neo-X chromosome with a large XR arm, similar in size to the neo-Y, at early-diplotene. In metaphase I, the scaffold structure of the neo-XY in all analyzed cells revealed two disjunctive configurations; despite ‘cross’-shaped configurations due to interstitial chiasmata which could distort segregation, although anomalous spermatids were not observed among a sample of 700 inspected. The standard X chromosome of grasshoppers is invariably allocyclic during male meiosis and strongly negatively heteropycnotic at metaphase I. This behaviour is conserved even when the X is fused to an autosome. Thus, the XL arm has a typical positive heteropycnotis but the XR maintains an euchromatic state (White, 1973; Hewitt, 1979; John, 1983). Additionally, a different condensation pattern was observed in the neo-Y: in the diplotene stage, its proximal-intestinal region shows a particularly condensed pattern while the interstitial-distal region is euchromatic.

Our results lead us to ask the following questions: (1) Are the neo-sex chromosomes of *B. noroestensis* in the initial state of the differentiation process? (2) If so, is a recent origin for *B. noroestensis* neo-sex chromosomes thus suggested? Probably we should suggest such a situation, based on melanopline neo-sex chromosome data, where in few cases the XR arm and neo-Y still seem to conserve their homology. Besides this, the conserved homology which allows for more than one chiasma at metaphase I suggests a recent evolutionary origin of neo-sex chromosomes. Even so, how could we explain the FN reduction? Moreover, we could not explain the observed chromosome number of the species using only a single chromosome rearrangement event. We propose rather that the atypical meiotic features
of neo-sex chromosomes of *B. noroestensis* imply a complex origin through several chromosomal rearrangements. In melanoelines there is a single case where neo-XY chromosomes are the result of multiple chromosome rearrangements and where Rb fusions are not involved (e.g. *D. vitattus*) (Bidau & Martí, 2001).

In light of these factors, we propose two models which may explain the origin of *B. noroestensis* neo-sex chromosomes. Firstly, a tandem fusion between two autosome pairs from the L group of the standard karyotype took place and became completely established, producing an unusually large telocentric autosome (*L₂*) (Fig. 4a). In the next step, a centric fusion between *L₃* and the X chromosome produced a large submetacentric neo-X; the homologue of the fused autosome became the neo-Y (Fig. 4c). The second model involves a Rb fusion between two autosome pairs from the L group that became fixed, thereby producing a metacentric pair; in the next step a pericentric inversion involving the new metacentric pair occurred, producing a large telocentric autosome *L₄* (Fig. 4b). Lastly, a centric fusion between *L₅* and the X chromosome produced a submetacentric neo-X; again, the homologue of the fused autosome became the actual neo-Y (Fig. 4c).

It is possible that the first model could be the better explanation, being the most parsimonious hypothesis because it involves less chromosome rearrangements in the evolution of neo-sex chromosomes. Additionally, it is difficult to imagine this chromosome arrangement becoming established in large populations. Based on cytogenetic records in Orthoptera, it is possible to propose an explanation of what probably occurred during their origin (White et al., 1967; John & Weissman, 1977). Even under the most favourable conditions, i.e. regularity in terms of synopsis and chiasma formation, tandem fusion heterozygotes will produce 50% of aneuploid gametes carrying both a deficiency or duplication. If this situation is compared with that of Robertsonian neo-XY heterozygotes, which produce a high number of euploid gametes, as occurs in normal male gametogenesis, selection is more drastic in the former case (White et al., 1967; White, 1973). It is possible that tandem fusion achieved initial success in small isolated populations (John & Weissman, 1977); if this is so, the low vagility of the species could have accelerated the process, thereby increasing the frequency of such chromosome forms in a few generations.

Either way, current cytogenetic evidence in the genus cannot assist in understanding the evolution of this chromosomal arrangement if the neo-XY sex chromosome system of *B. noroestensis* occurred in a remote ancestor or if different chromosome rearrangements were incorporated independently in *Boliviacris* species. Hence, neo-sex chromosomes could provide a certain degree of adaptive advantage to the carrier organism, or sex; in any case, it could be an important mechanism by which increase of the frequency of this karyotypic variant within natural populations versus the previous arrangement might occur without detriment of the ongoing micro-evolutionary forces (such as genetic drift in small populations), pertaining at least at the beginning of the evolutionary history of neo-sex chromosomes (White, 1978; Rieseberg, 2001).

We can also make some final assessment concerning how probable these models are and implications of this neo-sex system in the context of Melanoplinae sex chromosome evolution. In South American Melanoplinae, there are three records of FN variation, especially including reductions. This feature could be explained only by multiple centric fusions events and also pericentric inversions, as were earlier proposed for *Dichroplus silveiraguidoi* Liebermann and *D. pratensis* Bruner (Orthoptera: Acrididae: Melano- plinae) (Cardoso et al., 1974; Sáez & Pérez-Mosquera, 1977; Mesa et al., 1982). Besides, Mesa et al. (1982) described a 2n = 19/20 (M/F) species (determined as *Chlorus* sp.) with all telocentric chromosomes and an X0/XX sex chromosome determination mechanism. Moreover, White et al. (1967) reported the only case known which involves tandem fusion in the neo-sex chromosome origin. Within *Dichroplus*, there is one record of multiple rearrangements, including a sex chromosome-autosome tandem fusion (*D. vittatus*) (Bidau & Martí, 2001) which supports their probable occurrence in the presently analyzed species. *Boliviacris noroestensis* neo-sex chromosomes are an example of a recently established sex mechanism, mainly because the observed homology between a large distal-interstitial region of the XR and the neo-Y. Despite this cytogenetic feature which allows interstitial chiasma, the low frequency of this karyotype within natural populations indicates regular meiotic behaviour of the system. Interstitial-distal chiasma, as observed, would facilitate a convergent orientation and a normal segregation of the sex bivalent, whilst the neo-Y proximal-interstitial region (positively heteropycnotic) which was evidenced at early meiosis, points to an interruption in the degeneration process where previous rearrangement took place.

We conclude that *B. noroestensis* neo-sex chromosomes have recently arisen through a complex mechanism and, as such, represent the first example within the Melanoplinae. However, it is well known that chromosomal rearrangements like tandem fusions need time to reach fixation within populations and all the cytogenetic consequences that structural changes (very large asymmetry) cause to their carriers. We can also infer, from cytogenetic data in related species, that several rearrangements could be involved in the chromosome number variation and FN reduction in the studied species.

Finally, as shown in earlier taxonomic work by Ronderos & Cigliano (1990), *B. noroestensis* is related to several Melanoplinae species originally assigned to the genus *Dichroplus*. From molecular and morphological data, Colombo et al. (2005) tested the monophyly of *Dichroplus*, and besides so doing, showed the evolutionary relationships between this group and related genera of Dichroplini. They also proposed an explanation to the karyotype diversification, suggesting that X-autosomal centric fusions took place several times during the evolution of this group (Colombo et al., 2005). From the optimization of chromosomal characters on the topology derived from the combined morpho-
logical and molecular analyses, centric fusions appeared in several independent branches among the phylogenetic trees thus produced. The evidence of multiple chromosome rearrangements (A-A/X-A) involved in the origin of typi-
cal karyotype as well as neo-sex chromosome systems in

D. vittatus, R. bergi and D. silveiraguidoi largely support our models, as described above.

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