Polytene Chromosome Map and Inversion Polymorphism in
Drosophila mediopunctata

Galina Ananina, Alexandre A Peixoto*, Wilma N Souza, Louis B Klaczko+

Departamento de Genética e Evolução, Instituto de Biologia, Universidade Estadual de Campinas, Unicamp, Caixa Postal 6109, 13083-9790 Campinas, SP, Brasil *Departamento de Bioquímica e Biologia Molecular, Instituto Oswaldo Cruz-Fiocruz, Rio de Janeiro, RJ, Brasil

Drosophila mediopunctata belongs to the tripunctata group, and is one of the commonest Drosophila species collected in some places in Brazil, especially in the winter. A standard map of the polytene chromosomes is presented. The breakpoints of the naturally occurring chromosomal rearrangements are marked on the map. The distribution of breaking points through the chromosomes of D. mediopunctata is apparently non-random. Chromosomes X, II and IV show inversion polymorphisms. Chromosome II is the most polymorphic, with 17 inversions, 8 inversions in the distal region and 9 in the proximal region. Chromosome X has four different gene arrangements, while chromosome IV has only two.

Key words: linkage disequilibrium - gene order - cytogenetic analysis - chromosome mapping

Drosophila mediopunctata is a Neotropical species with a wide geographical distribution, found in many parts of Brazil and in El Salvador (Val et al. 1981). In some areas, particularly in the South or at high altitudes during the winter, it may be the commonest of the Drosophila species collected (Saavedra et al. 1995). D. mediopunctata belongs to the tripunctata group (Frota-Pessoa 1954), which is the second largest group of the Neotropical region (Vilela 1992). Its identification is relatively easy, contrasting with most species of the group that are difficult to identify (Vilela 1992). Various aspects of its biology make D. mediopunctata an interesting model for studies of the genetics of natural populations (for a review and discussion, see Klaczko 1995).

D. mediopunctata has 5 pairs of acrocentric chromosomes and a pair of dots (Kastritis 1966). Chromosomes X, II and IV show inversion polymorphisms. Chromosome II is the most polymorphic with 17 inversions. Chromosome X has only two gene arrangements (Carvalho et al. 1989, Klaczko et al. 1990, Peixoto & Klaczko 1991).

The presence of chromosome inversion polymorphisms is typical of various species of Drosophila (Krimbas & Powell 1992, 2000). These polymorphisms are often under selection, being dynamic with seasonal, geographical and long-term frequency variations (Sperlish & Pfriem 1986, Anderson et al. 1991). In D. mediopunctata we also found seasonal, micro- and macrogeographic variations in the frequencies of the karyotypes (Klaczko 1995). Moreover, there are differences among karyotypes for characters associated with fitness, such as size and shape (Bitner-Mathé et al. 1995).

Kastritis (1966) made a map of the polytene chromosomes of D. mediopunctata using a single isofemale line. However, he was able to identify two inversions on chromosome II. We now present a more detailed map, locating the breaking points of all the inversions.

MATERIALS AND METHODS

Strains used and field collection - To analyse the gene arrangements, we used D. mediopunctata isofemale lines maintained in our laboratory (Laboratório de Genética, Ecologia e Evolução de Drosofilídeos, Unicamp, Brazil). They were originally set up with flies collected in various places in Brazil (Atibaia, SP; Campinas, SP; Itatiaia, MG; Juiz de Fora, MG; Jundiaí, SP; Porto Alegre, RS). This ensured the sampling of the most common inversions. To obtain the missing arrangements, we made field trips to the Reserva Santa Genebra (Campinas, SP) and to the Japi Mountains (Jundiaí, SP). However, the mapping of some very rare inversions (DL, PA8, PC5 and PC3) was based only on the analysis of old chromosome preparations and of photographs from the laboratory collection.

In the field collections, we used fermented banana baits or traps, as described in Peixoto and Klaczko (1991) and Medeiros and Klaczko (1999). The analysis methodology followed the procedure described by Peixoto and Klaczko (1991).

Preparation of the polytene chromosomes - The polytene chromosomes were prepared using third instar larvae, following the method described by Ashburner (1989), with 1N HCl and lacto-acetic orcein.

Map drawing - Various drawings of D. mediopunctata polytene chromosomes were made using a camera lucida (1700x magnification). To prepare the consensus map, we used up to 10 drawings for each region from different nuclei.
RESULTS AND DISCUSSION

Cytological map - The cytological map of the five acrocentric chromosomes of *D. mediopunctata* is shown in Fig. 1. We followed the convention of Bridges (1935) in *D. melanogaster*, dividing the polytene chromosomes of *D. mediopunctata* in a total of 100 regions, numbered from 1 to 100. Each chromosome has 20 segments and a remarkable band at the beginning of each segment. The numerical regions were subdivided in smaller fragments named with letters.

In the map published by Kastritsis (1966) each chromosome was divided in 10-11 segments named with letters. Our map is more detailed, but we preserved, whenever possible, Kastritsis divisions. However, we adopted numbers for the description of inversions, since it is more convenient, except for the second chromosome (see below).

In the map presented, there are 372 bands in chromosome X; 422 in chromosome II; 446 in chromosome III; 440 in chromosome IV; and 431 in chromosome V; with a total of 2,111 bands.

In spite of the misleading simplicity, the exact determination of the number of bands in the cytological maps remains an unsolved problem. In species of Diptera, the numbers vary from 1,500 to 5,000. It is unlikely that this is due to a fundamental genomic variation. Of course, there are objective and subjective factors that affect the determination of the number of bands. We now know that artifacts of fixation with acetic acid cause the appearance of double bands and consequently an overestimate of the number of bands. Moreover, the identification of bands depends also on the physiological state of the cell, on the degree of polytenization and on the stretching of the chromosomes. Using electron microscopy in three species, (*Chironomus tentans, D. melanogaster* and *D. hydei*) the best estimates of the number of bands were obtained as being around 3,500. This number is considered typical for all Diptera. But, in chromosomes normally stretched, 2,500 bands can usually be identified (Zhimulev 1996). Thus, the number we obtained in our work (2,111) corresponds to the normal stretching of the chromosomes.

Description of the gene arrangements - The most convenient way to describe a gene arrangement is by comparison to a standard. For chromosomes X, III, IV and V we chose the commonest configuration in natural populations as the standard. For the second chromosome there are too many arrangements and no single one can be considered the most representative of the species. We chose DI-PC0 as standard, since both DI and PC0 are, respectively, the probable ancestors for the distal and proximal regions (Peixoto & Klaczko 1991). The breaking points for each of the inversions are shown in Fig. 1.
Chromosome II is the most polymorphic for inversions. Its complexity can be simplified by assigning the inversions to two groups according to the chromosome region: distal and proximal. Figs 2A, D show, respectively, the double heterokaryotypes DA-PA0/DI-PB0 and DS-PC0/DI-PB0 which are characteristic of the configurations found in nature and where one can clearly see the two regions. The distal group includes 8 inversions: DI, DS, DV, DP, DA, DL, DR and DJ. In the proximal region, there are 9 inversions: PC0, PC1, PC2, PC3, PC4, PC5, PB0, PA0 and PA8. Figs 2B, C, E show, respectively the heterokaryotypes DP/DI, DV/DI, and PA0/PC0.

In almost all combinations between proximal and distal inversions there is no overlap between the two regions. Thus, in principle recombination between them can occur. However, there is a very strong linkage disequilibrium and consequently, the number of configurations found in nature is reduced (Peixoto & Klaczko 1991).

The distal inversions and their relation to DI are shown diagrammatically in Fig. 1. Inversion DJ is presented as a separated arrangement, but it is nearly always found associated to DV which is also a distal inversion. This latter can be found isolated with good frequency. The combination DV+DJ was previously called DT (Peixoto & Klaczko 1991).

The proximal inversions can be divided in three phylads (Dobzhansky 1970): PC, PB and PA. The phylad PC (Fig. 1) is made up of six gene arrangements: PC0, PC1, PC2, PC3, PC4 and PC5. These latter five can be considered derived from PC0.

The PA phylad has two gene arrangements: PA0 and PA8 (Fig. 1). PA0 is complex, since it is an intrachromosomal transposition of two inverted fragments: In(2)PA0 [= In(2) 28A; 36A; 36C; 37C]. The heterokaryotypes PC0/PA0 and PB0/PA0 display complicated configurations (with segments 36A-36C always unpaired).

PB0 is isolated in the PB phylad (Fig. 1). Compared to PC0, it is an inverted insertion transposition in the second chromosome: In(2)PB0 [= In(2) 28A; 28B; 36C].

The arrangements PA0, PB0 and PC0 are overlapping inversions. This allows us to make inferences about their phylogenetic relations (Dobzhansky & Sturtevant 1938). If compared, one can not relate any two of these inversions through a single inversion event. However, it is possible to assume and draw a hypothetical arrangement (PH) that is a parsimonious explanation relating the three gene inversions (Fig. 3) as it was done with the chromosomes of D. pseudoobscura (Dobzhansky 1970).

Chromosome X has four gene arrangements: Standard, 1, 2 and 3 (Fig. 1). There is a strong linkage disequilibrium between In(X)1 and In(X)2. They are often found together (Peixoto & Klaczko 1991).

In chromosome IV, aside from the Standard gene arrangement, we found only one inversion, In (IV)1 (Fig. 1).

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![Fig. 2: the most frequent inversions of *Drosophila mediopunctata*. A: DA-PA0/DI-PB0; B: DP/DI; C: DV/DI; D: DS-PC0/DI-PB0; E: DA-PA0/DS-PC0. The open arrows point to the distal region and the closed arrows to the proximal region. Bar = 100 µm](image)
The distribution of breaking points through the chromosomes of *D. mediopunctata* is apparently non-random. The second chromosome is the most saturated. In this chromosome, we found a subterminal inversion \([\text{In}(2)\text{DS}, \text{Fig. 1}]\), but no subbasal inversions. The majority of the breaking points are found in its distal half portion. In some cases, different inversions share cytologically the same breaking point (Fig. 1).

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