

# Segmental Duplication, Microinversion, and Gene Loss Associated with a Complex Inversion Breakpoint Region in *Drosophila*

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## Abstract

Chromosomal inversions are usually portrayed as simple two-breakpoint rearrangements changing gene order but not gene number or structure. However, increasing evidence suggests that inversion breakpoints may often have a complex structure and entail gene duplications with potential functional consequences. Here, we used a combination of different techniques to investigate the breakpoint structure and the functional consequences of a complex rearrangement fixed in *Drosophila buzzatii* and comprising two tandemly arranged inversions sharing the middle breakpoint:  $2m$  and  $2n$ . By comparing the sequence in the breakpoint regions between *D. buzzatii* (inverted chromosome) and *D. mojavensis* (noninverted chromosome), we corroborate the breakpoint reuse at the molecular level and infer that inversion  $2m$  was associated with a duplication of a  $\sim 13$  kb segment and likely generated by staggered breaks plus repair by nonhomologous end joining. The duplicated segment contained the gene *CG4673*, involved in nuclear transport, and its two nested genes *CG5071* and *CG5079*. Interestingly, we found that other than the inversion and the associated duplication, both breakpoints suffered additional rearrangements, that is, the proximal breakpoint experienced a microinversion event associated at both ends with a 121-bp long duplication that contains a promoter. As a consequence of all these different rearrangements, *CG5079* has been lost from the genome, *CG5071* is now a single copy nonnested gene, and *CG4673* has a transcript  $\sim 9$  kb shorter and seems to have acquired a more complex gene regulation. Our results illustrate the complex effects of chromosomal rearrangements and highlight the need of complementing genomic approaches with detailed sequence-level and functional analyses of breakpoint regions if we are to fully understand genome structure, function, and evolutionary dynamics.

**Key words:** inversion, breakpoint, *Drosophila*, BAC, shotgun sequencing, transposable elements.

## Introduction

Changes in the structure of chromosomes have long been recognized as significant for the evolution of eukaryotes (Sturtevant 1917; Dobzhansky 1947; Sperlich and Pflüger 1986), and their role in speciation, adaptation, evolution of sex chromosomes, and also in human and animal disease is widely documented (Murphy et al. 2005; Lindsay et al. 2006; Darai-Ramqvist et al. 2008; Hoffmann and Rieseberg 2008; Chen et al. 2010; Kirkpatrick 2010). In recent years, a renewed interest in the study of chromosomal rearrangements has arisen as a consequence of the availability of genome sequences (Hoffmann and Rieseberg 2008; Hurler et al. 2008; Kirkpatrick 2010). The increased resolution of comparative genomic approaches has revealed that structural variants in general and chromosomal inversions in particular are much more common than previously thought (Coghlan et al. 2005; Feuk et al. 2005; Hoffmann and Rieseberg 2008; Kirkpatrick 2010). Genomic comparisons are also revealing that rather than being simple two-breakpoint rearrangements changing gene order, inversions

may often entail gene duplications that can lead to genetic novelty and adaptation (Ranz et al. 2007; Kaessmann 2010; Furuta et al. 2011). Given the prevalence of chromosomal rearrangements and their associated functional consequences, understanding the mechanisms that generate them and their functional impact is necessary in order to understand the structure, function, and evolution of eukaryotic genomes.

The analyses of the chromosomal inversion breakpoints identified so far suggest that several mechanisms underlie their generation. For example, inversions can be generated by ectopic recombination between “homologous sequences” present at different chromosomal sites in opposite orientation (Petes and Hill 1988; Lim and Simmons 1994). The “homologous sequences” can be, for example, copies of transposable elements (TEs) that belong to the same TE family (Cáceres et al. 1999; Evans et al. 2007; Delprat et al. 2009), segmental duplications (Coulibaly et al. 2007), or tandemly arranged short repeats (Richards et al. 2005; Lobo et al. 2010). Inversions can also be generated by two simultaneous breaks in the same chromosome and repair by nonhomologous end joining (NHEJ), which

sometimes leads to the duplication of sequences present at the breakpoints (Kehrer-Sawatzki et al. 2005; Matzkin et al. 2005; Sharakhov et al. 2006; Ranz et al. 2007). Because only a limited number of inversion breakpoints have been characterized so far, it is likely that additional analyses may elucidate novel molecular mechanisms generating them. Increasing our understanding of the mechanisms generating rearrangements should also improve our ability to design strategies to detect rearrangements, which is particularly interesting for rearrangements associated with human diseases (Chen et al. 2010).

The increased resolution of the techniques used to study chromosomal rearrangements also suggests that chromosome breakage is nonrandom. Breakpoint reuse, a term used as a shorthand for close clustering of breakpoints (Pevzner and Tesler 2003) appears to be a common phenomenon in eukaryotes (Pevzner and Tesler 2003; Bourque et al. 2004; Murphy et al. 2005; Gordon et al. 2007). However, there is some controversy about whether the identified cases of breakpoint reuse are real cases of reuse or a consequence of the limited resolution of the genomic studies since breakpoints are usually only assignable to regions spanning a few hundred kilobases (Kehrer-Sawatzki and Cooper 2008; Sankoff 2009). Here, we will use the terms “breakpoint clustering” and “breakpoint reuse” nonsynonymously to refer to the apparent reuse of breakpoints due to clustering and to the actual reuse at the molecular level, respectively. Therefore, a precise identification and characterization of sequences at inversion breakpoint sites and of adjacent sequences is needed in order to shed light on the nonrandom distribution of rearrangement breakpoints.

Detailed analyses of inversion breakpoint sequences are also needed in order to elucidate the functional consequences of chromosomal inversions. Several hypotheses have been proposed to explain the adaptive significance of inversions. First, inversions might have functional consequences due to the mutational impact of breakpoints (Sperlich and Pfriem 1986). Inversions might affect the expression, structure, and function of genes by removing or exchanging the regulatory regions of genes adjacent to the breakpoints, by disrupting the genes spanning the breakpoints, or by creating chimeric genes (Puig et al. 2004; Kehrer-Sawatzki and Cooper 2007; Hurles et al. 2008). The functional impact of inversions could also be due to their recombination-reduction effect (Sturtevant and Beadle 1936; Navarro and Ruiz 1997; Navarro et al. 1997). In heterozygotes, crossing-over and recombination are reduced inside the inverted segment and the alleles of the genes inside the inversion tend to be inherited as a single block. Accordingly, an inversion might be adaptive if it captures a favorable allele combination (Dobzhansky 1970; Charlesworth 1974) or a set of locally adapted genes (Kirkpatrick and Barton 2006). An inversion could also be favored if it captured a set of genes relatively free of recessive deleterious mutations (Nei et al. 1967). Alternatively, functional consequences of inversions might be due to their role in the formation and maintenance of selfish gene complexes that increase in frequency in populations

through meiotic drive (Novitski 1967; Kirkpatrick and Barton 2006; Presgraves et al. 2009).

*Drosophila* is a good model organism to study the molecular mechanisms and the functional consequences of chromosomal inversions. Chromosomal inversions are particularly abundant in *Drosophila* both as intraspecific polymorphisms and as fixed differences between species (Krimbas and Powell 1992). Within the *Drosophila* genus, the species of the repleta group are particularly good models to study genome evolution because, besides the wealth of information on genome rearrangements and the availability of genomic resources, the species of this group have been used for studies of ecological adaptation and speciation for 70 years (Spencer 1941; Wharton 1942; Barker 1982; Manfrin and Sene 2006). Finally, the wealth of genetic tools and functional information available for *Drosophila* (Roy et al. 2010) facilitate the analysis of the potential functional consequences of inversion breakpoints in species of this genus.

Here, we used the Bacterial Artificial Chromosome (BAC) library and physical map of the *Drosophila buzzatii* genome (González et al. 2005) and the genome sequence of *D. mojavensis* (Clark et al. 2007), both species belonging to the repleta group, to isolate and characterize the breakpoints of a complex rearrangement of chromosome 2 fixed in all species of the *buzzatii* complex. This rearrangement comprises inversions  $2m$  and  $2n$  that were described by cytological studies as arranged in tandem and sharing the middle breakpoint (Ruiz and Wasserman 1993; González et al. 2007). The aims of this study are: 1) to determine the mechanisms that generated  $2m$  and  $2n$  fixed inversions, 2) to elucidate whether the described breakpoint clustering is in fact a breakpoint reuse at the sequence level, and 3) to assess potential functional effects of these inversions. To accomplish these aims we combined the classical *in situ* hybridization technique with shotgun sequencing of BAC clones, comparative genomics analyses and bioinformatic and experimental gene expression analyses. The combined results of these analyses allowed us to shed light not only into the mechanism that generated  $2mn$  rearrangement but also on its functional consequences.

## Materials and Methods

### *Drosophila* Stocks

*D. buzzatii* stock *st-1* (González et al. 2005), *D. repleta* stock 15084-1611.06 from Siboney, Cuba (University of California [UC] San Diego *Drosophila* Stock Center) and *D. mojavensis* stock 15081-1352.22 from Catalina Island, CA (UC San Diego *Drosophila* Stock Center) were used in this work. Additionally, the genomes of the 12 *Drosophila* species sequenced were used as reference in the comparative analyses (Clark et al. 2007).

### *D. buzzatii* BAC Clones

BAC clones putatively encompassing inversion breakpoints were selected from the genomic library CHORI-225 (<http://bacpac.chori.org>), using the physical map of the *D. buzzatii* genome (González et al. 2005, 2007) available at the

Genome Sciences Centre website (<http://www.bcgsc.ca/ice>). Clones containing breakpoints were identified because they produce a single signal when hybridized to *D. buzzatii* inverted chromosomes and two signals when hybridized to the noninverted chromosomes of *D. repleta*.

### BAC Walking

Primers were designed in the *D. mojavensis* genome preferentially in coding regions using PRIMER software (supplementary table S6 in additional file 3, Supplementary Material online). Amplification reactions were carried out in a total volume of 50  $\mu$ l: 50–200 ng of DNA as template, 40 pmols of each primer, 200  $\mu$ M of dNTPs, 1.5 mM Mg<sub>2</sub>Cl, and 1 U of *Taq* DNA Polymerase (Platinum *Taq* DNA Polymerase High Fidelity; Invitrogen; Carlsbad, CA). When polymerase chain reactions (PCRs) were carried out using DNA from a species different from that used to design the primers as template, the annealing temperature of the PCR was set 5 °C lower than the suggested temperature to account for putative nucleotide mismatches.

### In Situ Hybridization

In situ hybridizations were carried out as in Montgomery et al. (1987). Probes were labeled with biotin-16-dUTP (Roche Diagnostics; Mannheim Germany) by random primed labeling and hybridized to *D. buzzatii* and *D. repleta* salivary gland chromosome squashes. Interspecific hybridizations were performed at 25 °C, while the intraspecific ones were performed at 37 °C. The localization of the hybridization signals was done using the cytological maps of *D. buzzatii* (Ruiz and Wasserman 1993; González et al. 2005, 2007). Two types of probes were used, BAC clones from the *D. buzzatii* CHORI-225 library (González et al. 2005) and PCR products generated in the chromosome walk (supplementary table S7 in additional file 3, Supplementary Material online).

### DNA Sequencing

PCR products were purified using QIAquick gel extraction kit (Qiagen; Valencia, CA, USA) and directly sequenced with the same primers used to amplify them. When no optimal sequencing chromatograms were obtained, the PCR products were cloned into the pGEM-T easy vector (Vector Systems I of Promega; Madison, WI) and sequenced using M13 forward and reverse primers. These PCR products were sequenced at MacroGen sequence service (<http://www.macrogen.com>). The same primers used to amplify the breakpoints were used for sequencing.

BAC ends were sequenced using T7 and SP6 primers of the vector *pTARBAC2.1* at GATC-biotech sequence service (<http://www.gatc-biotech.com>). Two BAC clones were fully sequenced: BAC 1N19 (AC breakpoint) and 20O19 (EB breakpoint). Sequences have been submitted to GenBank (accession numbers JQ611723 and JQ611724). Random shotgun sequencing was provided by MacroGen (<http://www.macrogen.com>). Details of the sequencing process are given in supplementary table S9 in additional file 3 (Supplementary Material online). Gap closure was carried out in our lab. Two gaps were closed in BAC clone 20O19

using primer pair A (5' TGTCGACTATAGTTAAGCGT 3' and 5' GGCAGTAGTCGTCGATTAT 3') and primer pair B (5' GTGAGGCAATGCGTAACATT 3' and 5' CTTCTTGCTACGCATAATCT 3'). One gap was closed for BAC clone 1N19 using primer pair (5' CTACGCAGATAAGCAGGCTT 3' and 5' AACTGTCAGCAGCAACGTGT 3').

### Similarity Searches and Sequence Annotation

Similarity searches against the *D. mojavensis* genome (CAF1 assembly released in February 2006) (Clark et al. 2007) were done using the BAC-end sequences as queries and PCR products generated in the chromosome walk and in the amplifications of the breakpoint regions. These searches were done with BlastN implemented at DroSpeGe (Gilbert 2007) and bl2seq implemented at NCBI (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). The two sequenced BAC clones were completely annotated. The annotation was done using the online software gene predictors GeneID (Parra et al. 2000) and GENESCAN (Tiwari et al. 1997). Gene predictions were compared with the GLEANR models of the *D. mojavensis* genome annotation available at DroSpeGe (Gilbert 2007) and to *D. melanogaster* annotations available at FlyBase (<http://flybase.org>) (Tweedie et al. 2009). Alignments were performed using MUSCLE (Edgar 2004) available at the EBI website (<http://www.ebi.ac.uk>) and modified as needed to adjust splicing, start and stop codons and frames.

For the annotation of TEs and repeat sequences in the two sequenced BAC clones, similarity searches were carried out against the TE database in the DPDB website (Casillas et al. 2005), RepeatMasker (Repbase library based) (Jurka et al. 2005), and DroSpeGe (ReAs library based) (Gilbert 2007). This limited our search to TEs that are conserved among species. We also perform similarity searches against the nonredundant nucleotide database at the NCBI website (Tatusova and Madden 1999) to identify any potential TE sequence not included in TE databases. After this general search, a more detailed analysis of the breakpoint regions was performed using bl2seq (Tatusova and Madden 1999). In each case, our sequence was used as query and the sequence of the TE producing the best hit in the previous search was used as subject. Only hits with a minimum size of 40 bp and  $E$  value  $\leq 1 \times 10^3$  were considered significant.

RepeatMasker also allowed us to look for AT-rich regions and low complexity repeats such CCG repeats. Bl2seq was also used to search for palindromic sequences. BlastN and bl2seq were used to search for Conserved Noncoding Sequences (CNSs). Overrepresented conserved Transcription Factor Binding Sites (TFBSs) were searched using Whole Genome rVISTA implemented in <http://genome.lbl.gov/vista> webpage.

### Dating the Duplication

We aligned the CDS of *Dmoj/CG4673* and *Dbuz/CG4673* and estimated Ks (number of synonymous substitutions per synonymous site) using DnaSP (Librado and Rozas 2009). Divergence time between this two species is 11.3 myr (Russo et al. 1995; Tamura et al. 2004) (Oliveira DCSG, Almeida FC, O'Grady P, Etges WJ, Armella MA, DeSalle R,



personal communication). The rate of neutral evolution ( $r$ ) per site per million years ( $r = Ks/2T$ ) is therefore  $0.0187 \times 10^{-6}$ . We then estimated the number of changes between the two duplicated regions (taking into account both coding and noncoding sequences), and we used  $r$  to estimate  $T$ .

### RNA Extraction and RT-PCR

Total RNA was isolated from different developmental stages (0–1.30, 0–2.15, 0–2.30, and 0–22 h embryos, larvae, pupae, adult males and females) of *D. buzzatii* with RNeasy Kit (Quiagen; Valencia, CA) and treated with 1 unit of DNase I (Applied Biosystems/Ambion; Austin, TX) for 30 min at 37 °C to eliminate DNA contamination. cDNA was synthesized with First Strand cDNA Synthesis Kit (Roche Diagnostics; Mannheim, Germany) using ~500 ng of the DNase I-treated RNA and Oligo-dT primer (Promega; Madison, WI). RT-PCRs were performed with volumes as previously described for PCR adding 1  $\mu$ l of the retrotranscriptase reaction. As a control of the reverse transcription reaction, *Gapdh* (438 bp) was amplified with H1 and H2 primers (data not shown) (Puig et al. 2004). Reactions without retrotranscriptase (RT-) were carried out to control for DNA contamination (data not shown).

The name of the primers used for RT-PCR corresponds to the region where they were designed for example primer 5R was designed in exon 5 (supplementary table S3 in additional file 3, Supplementary Material online). Only one RT-PCR was performed for *D. buzzatii* CG4673 with primers designed in exons 5 and 6. Four different RT-PCRs were performed for *D. buzzatii*  $\Psi$ CG4673 that attempted to amplify the regions that showed homology with exons 2, 3, 7, and to the 3' UTR region (supplementary table S3 in additional file 3, Supplementary Material online). Position 3' of these primers were located in a mismatch to amplify specifically  $\Psi$ CG4673 in AC breakpoint. We sequenced the five RT-PCR products, and we confirmed that they corresponded to exons 5–6 of CG4673 and exons 2, 3, 7, and to the 3' UTR of  $\Psi$ CG4673.

### Rapid Amplification of Complementary DNA End

Rapid Amplification of Complementary DNA End (RACE) experiments were carried out with DNase I-treated total RNA from embryos and adults of *D. buzzatii* using specific adapters and primers from FirstChoice RLM-RACE (Applied Biosystems/Ambion; Austin, TX, USA). For CG4673, 5' RACE primers were designed in the first exon (1L and 1Lint; supplementary table S3 in additional file 3, Supplementary Material online) and primers for 3' RACE in the last exon (8R and 8Rint; supplementary table S3 in additional file 3, Supplementary Material online). For  $\Psi$ CG4673, 5' RACE primers were designed in exon 2 (2L and 2Lint; supplementary table S3 in additional file 3, Supplementary Material online) and primers for 3' RACE in exon 7 (7R and 7Rint; supplementary table S3 in additional file 3, Supplementary Material online).

### Northern Blot

Northern blot hybridization was performed as described in De et al. (1990). RNA from embryos, larvae, pupae, adult

males and females was used. Two different membranes were hybridized one with 5–10  $\mu$ l (~1,300 ng/ $\mu$ l) of total RNA and the other one with 1–3  $\mu$ l (~1,300 ng/ $\mu$ l) of total RNA. Samples were loaded with 3V of Northern Formaldehyde Load Dye (Applied Biosystems/Ambion; Austin, TX). Running buffer 10 $\times$  for formaldehyde gels (Applied Biosystems/Ambion; Austin, TX, USA) was used to run the gels in denaturalizing conditions. The probe was amplified with primers 5R and 7L (supplementary table S3 in additional file 3, Supplementary Material online). Hybridization was performed with 5  $\mu$ l (15.3 ng/ $\mu$ l) of the [<sup>32</sup>P]UTP-labeled probe. Hybridization was carried out for 16 h at 65 °C with Perfect Hyb Plus buffer (Sigma–Aldrich; St. Louis, MO). Two washes with 2 $\times$  saline sodium citrate/ 0.1% sodium dodecyl sulfate were done before exposition.

### Double-Stranded RNA Detection

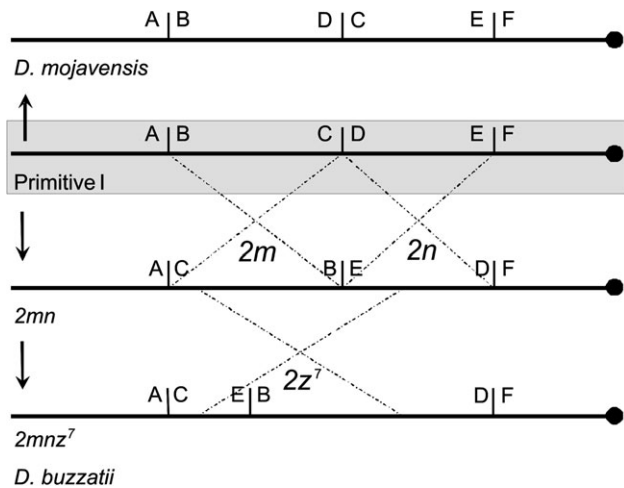
Detection of double-stranded RNA (dsRNA) was carried out as in Aravin et al. (2007). RT-PCRs were performed with RNA of embryos and adults of *st-1* line of *D. buzzatii*. Total RNA was treated with RNase ONE (Promega; Madison, WI) to degrade single-stranded RNA (D+). RNase-treated RNA was denatured and retrotranscribed as described for RT-PCR. Negative controls without retrotranscriptase (RT-), without RNase (D-), without the denaturalizing step (N) and using Oligo-dT primer (T) instead of random primers were performed. dsRNA detection were carried out with the primers 2R and 2L, 3R and 3L, 7R and 7L, and 3' UTR-R, 3' UTR-L and *Gapdh* primers H1 and H2 as described for RT-PCR (supplementary table S3 in additional file 5, Supplementary Material online).

## Results

### High-Resolution Mapping and Isolation of Inversion Breakpoints

Following previous works (Wesley and Eanes 1994; Cáceres et al. 1999), the three breakpoint regions of inversions  $2m$  and  $2n$  were tentatively designated as AB, CD, and EF to refer to breakpoints in the noninverted chromosome, and AC, BE, and DF to describe breakpoints in the inverted chromosome (fig. 1). In both cases, A represents the most distally located region relative to the centromere, whereas F stands for the most proximal region.

To clone and sequence the  $2m$  and  $2n$  breakpoint regions, we used the BAC-based physical map of the *D. buzzatii* genome (González et al. 2005) and the genome sequence of *D. mojavensis*, the closest relative to *D. buzzatii* whose genome has been fully sequenced (Clark et al. 2007). We followed a procedure similar to that devised by Prazeres da Costa et al. (2009) comprising four experimental steps. 1) We used in situ hybridization to identify *D. buzzatii* BAC clones encompassing the breakpoints (table 1 and supplementary fig. S1 in additional file 1, Supplementary Material online). 2) We sequenced the ends of the identified BAC clones, and we located the sequences in the *D. mojavensis* genome, which allowed us to narrow down the position of the breakpoints to ~100–150 kb regions (fig. 2).



**Fig. 1.** Schematic representation of chromosome 2 arrangements in Primitive I (the “repleta” group ancestor), *D. buzzatii* and *D. mojavensis*. Three inversions took place between Primitive I and *D. buzzatii*. Inversions  $2m$  and  $2n$  share the middle breakpoint. Inversion  $2z^7$  took place on a  $2mn$  chromosome inverting BE breakpoint region. Seven inversions (not shown in this scheme) took place from Primitive I to *D. mojavensis* ( $2c$ ,  $2f$ ,  $2g$ ,  $2h$ ,  $2q$ ,  $2r$ , and  $2s$ ) and as a result, inversion breakpoint region CD is inverted in *D. mojavensis*.

3) We walked along the *D. mojavensis* genome by designing DNA probes and hybridizing them to the *D. buzzatii* chromosomes, which allowed us to further delimit the breakpoint regions in *D. mojavensis* (fig. 2): AB breakpoint region was located in  $\sim 10$  kb fragment between *msi* (A) and *Ssadh* (B); CD breakpoint was located in a  $\sim 1.5$  kb region between *scrib* (C), and *Or98b* (D); and EF breakpoint region was located in a  $\sim 2.1$  kb region between *Wsck* (E) and *CG8147* (F). 4) We isolated the breakpoint regions in *D. buzzatii* by PCR using primers designed in *D. mojavensis* in the expected orientation according to figure 1 (Supplementary tables S8 in additional file 1, Supplementary Material online). Although the last step was attempted for the three breakpoints with different primers, it only worked out for DF breakpoint region. In *D. buzzatii*, the region between *Or98b* (D) and *CG8147* (F), and containing DF breakpoint is 4.4 kb long (GenBank accession number JQ611725). Comparison of this 4.4 kb region with CD and EF regions of *D. mojavensis* further narrowed down the breakpoint to a  $\sim 3.1$  kb segment. For breakpoint regions AC and BE, we had to resort to an alternative step. 5) We sequenced the whole BAC clones containing the AC and BE inversion breakpoints (fig. 3).

BAC clone 1N19 is  $\sim 134$  kb long and contains complete coding sequence (CDS) of three genes and partial CDS for another two genes (fig. 3; GenBank accession number JQ611723). Clone 20O19 is  $\sim 143$  kb long and contains full CDS of 22 genes and partial CDS of another one (fig. 3; GenBank accession number JQ611724). Almost all genes (with the exception of *CG4673*; see below) have a similar exon-intron structure and identical orientation in *D. buzzatii* and *D. mojavensis* (fig. 3). By comparing these two BAC clones to *D. mojavensis* genome sequence, we confirmed that each

**Table 1.** *D. buzzatii* BAC Clones Containing the Breakpoints of Inversions  $2m$  and  $2n$ .

Breakpoint Region	BAC	Size (base pairs) <sup>a</sup>	Cytological Coordinates in <i>D. buzzatii</i> <sup>b</sup>	Cytological Coordinates in <i>D. repleta</i>
AC	1N19	138,197	D3e/F2a	D3e and F2a
AC	14B19 <sup>c</sup>	140,515	D3e/F2a	D3e and F2a
AC	15L19	133,077	D3e/F2a	D3e and F2a
BE	20O19 <sup>c</sup>	142,697	D3e/F6h	D3e and F6h
BE	22B03	128,945	D3e/F6h	D3e and F6h
BE	2N19	191,034	D3e/F6h	D3e and F6h
DF	16H04 <sup>c</sup>	124,185	F6h/F2a	F6h and F2a
DF	14E21	195,292	F6h/F2a	F6h and F2a
DF	8C14	156,485	F6h/F2a	F6h and F2a

<sup>a</sup> Estimates of BAC insert sizes taken from the *D. buzzatii* BAC-based physical map available at the Genome Sciences Centre (<http://www.bcgsc.ca/ice>).

<sup>b</sup> Cytological coordinates refer to the cytological map of *D. repleta* chromosome 2.

<sup>c</sup> These clones were identified by González et al. (2007).

clone contains a single synteny interruption (breakpoint). Instead of being located between *msi* (A) and *scrib* (C), as expected according to *D. mojavensis* chromosome annotation (fig. 2), in *D. buzzatii* AC breakpoint was located in a  $\sim 15$  kb region between *CG12250* (A) and *scrib* (C). This is so because, in *D. buzzatii*, *CG12250* and *msi* are overlapping instead of nested genes (fig. 3). Manual reannotation of *CG12250* in *D. mojavensis* revealed that *CG12250* and *msi* are also overlapping in this species (fig. 3). Comparison of this 15-kb region with AB and CD regions of *D. mojavensis* allowed us to further narrow down the breakpoint to a  $\sim 700$  bp segment. Finally, in *D. buzzatii*, BE breakpoint is located between genes *Wsck* (E) and *Ssadh* (B) in a  $\sim 8$  kb region (fig. 3). Comparison of this  $\sim 8$  kb region with AB and EF regions of *D. mojavensis* located the breakpoint in a  $\sim 1.1$  kb segment. A more detailed description of this experimental procedure is given in supplementary additional file 2 (Supplementary Material online).

### Sequence Analysis of Breakpoint Regions in *D. buzzatii* and *D. mojavensis*

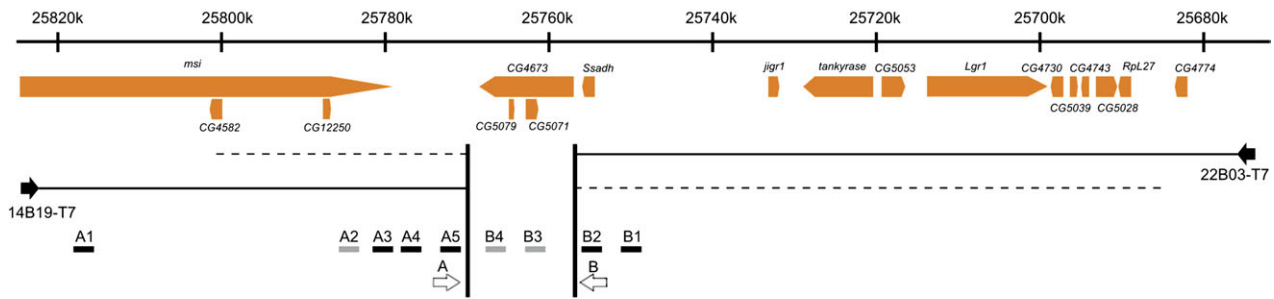
We performed a detailed analysis of the three breakpoint regions including annotation of 1) genes, 2) gene-associated sequences, 3) TE insertions, and 4) other structural features (fig. 4).

#### Gene Annotation

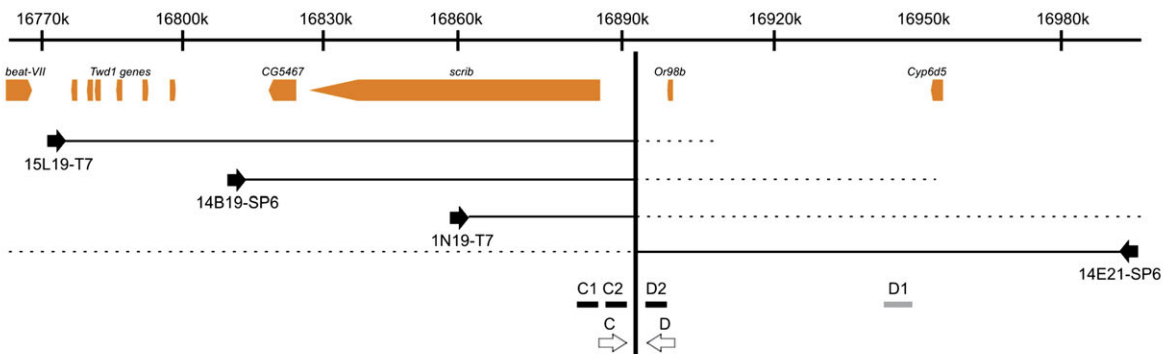
In *D. mojavensis*, the AB breakpoint region is located between *CG12250* and *Ssadh* and contains three genes: *CG5079* and *CG5071* nested within intron 6 of *CG4673* (fig. 4).

In *D. buzzatii*, the AC breakpoint region comprises the region between *CG12250* and *scrib*, which is 15,343 bp long (fig. 4). Only one putatively functional ORF was detected in this region corresponding to *CG5071*, located 6,585 bp upstream of the *CG12250* start codon. In *D. buzzatii*, *CG5071* is a 2,004-bp long gene and encodes a 668 aa-long protein. Thus, in *D. buzzatii*, the coding region of this gene is considerably longer than in *D. mojavensis* (225 aa) and similar to that of *D. melanogaster* annotated CDS *CG5071*-RB (680 aa).

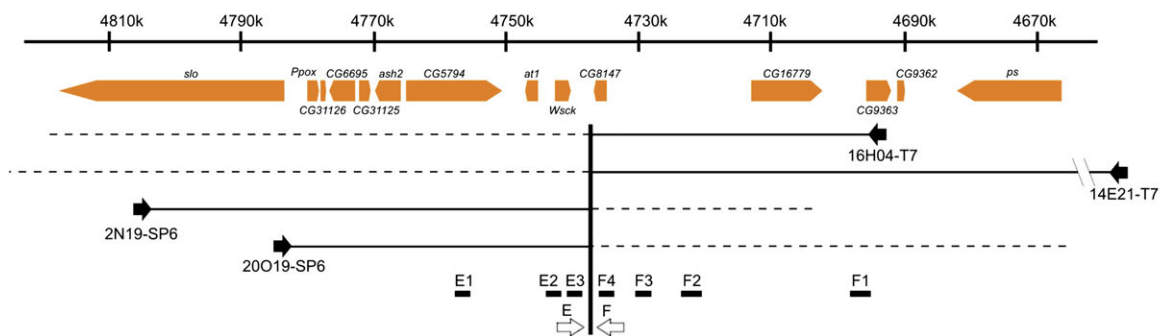
### AB breakpoint



### CD breakpoint



### EF breakpoint

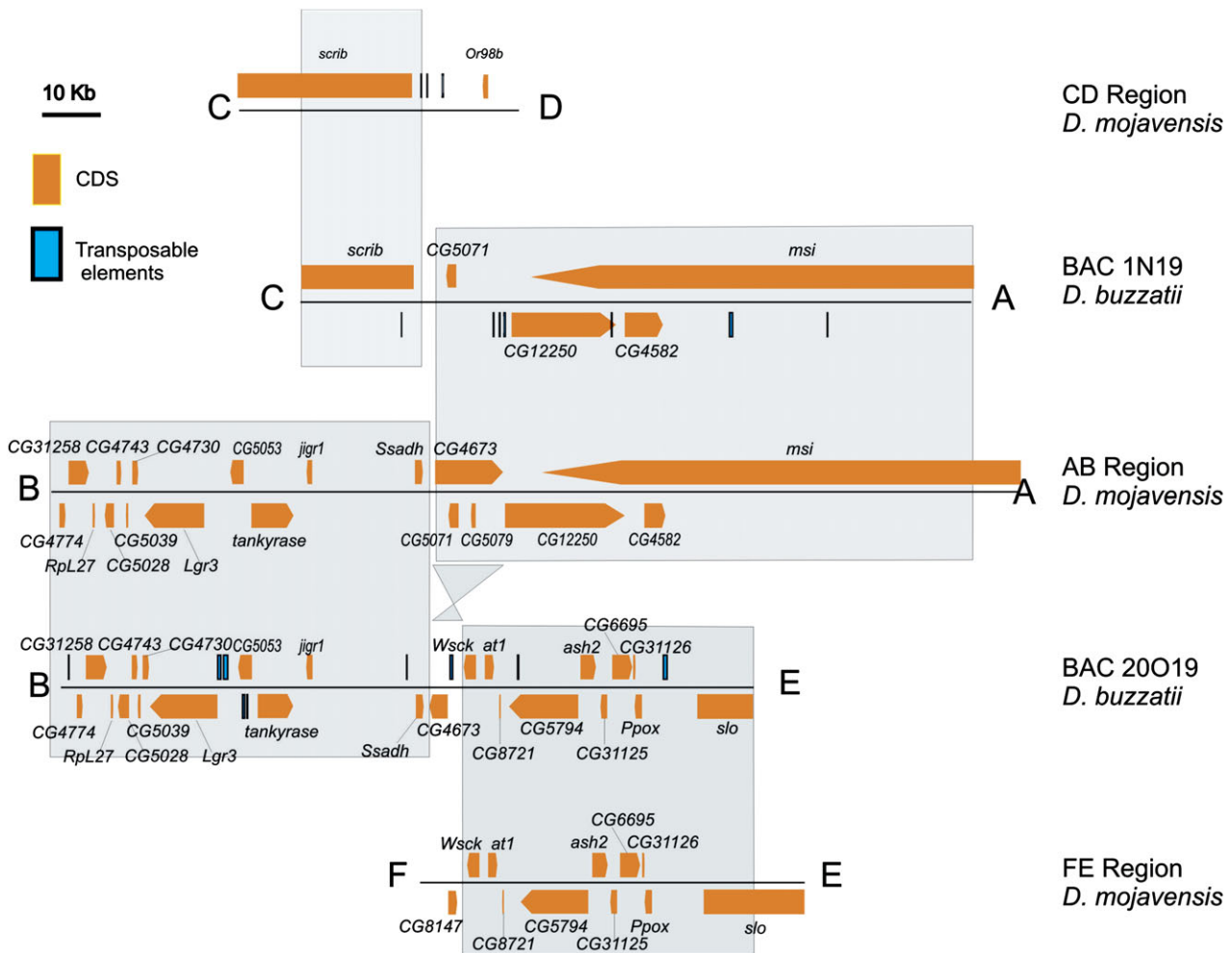


**Fig. 2.** Schematic view of the BAC walking through *D. mojavensis* genome to map the breakpoints of inversions 2*m* and 2*n*. Nucleotide coordinates above the gene structure map are from scaffold\_6540 of *D. mojavensis* (Schaeffer et al. 2008). Black arrows show where the sequenced BAC ends were anchored. The discontinuous lines show the length of the BAC clones and therefore the maximum distance where the breakpoints can be located. Black boxes indicate successfully hybridized PCR probes, while gray boxes are probes that did not yield any hybridization signal. White arrows show the location of the different primers designed to attempt the amplification of the breakpoints in *D. buzzatii* (not drawn to scale).

A small fragment with similarity to *Dmoj*/CG5079 was detected in *D. buzzatii* AC region (fig. 4). However, this fragment contains multiple stop codons and was therefore considered to be a pseudogene ( $\Psi$ CG5079). Additionally, 16 small blocks (45–273 bp long) showing 65.2–88.9% identity to *Dmoj*/CG4673 were detected in this region (fig. 4 and supplementary table S1 in additional file 3, Supplementary Material online). The distance between the most remote CG4673 hits in the AC breakpoint region is 12,585 bp, which is similar to the distance between the orthologous hits in *D. mojavensis* (12,578 bp). However, we could not identify the start or stop codon of

CG4673 in this region of the *D. buzzatii* genome and concluded that a degenerated, seemingly nonfunctional, copy of CG4673 ( $\Psi$ CG4673) is present in this breakpoint region (fig. 4).

The *D. buzzatii* EB breakpoint region is located in the 8,242-bp long intergenic region between *Wsck* and *Ssadh* (fig. 4). A seemingly complete copy of gene CG4673 is present between *Wsck* and *Ssadh* (supplementary table S2 in additional file 3, Supplementary Material online). This gene putatively encodes three transcripts that are similar in exon number and exon size to those present in *D. mojavensis* and *D. melanogaster* (supplementary table S2 in additional file 3,



**FIG. 3.** Annotation of the sequenced *D. buzzatii* BACs 1N19 and 20O19 by comparison with the homologous regions in scaffold 6540 of *D. mojavensis*. Shaded rectangles include homologous regions between the two species. See [supplementary additional data file 2](#) (Supplementary Material online) for further information.

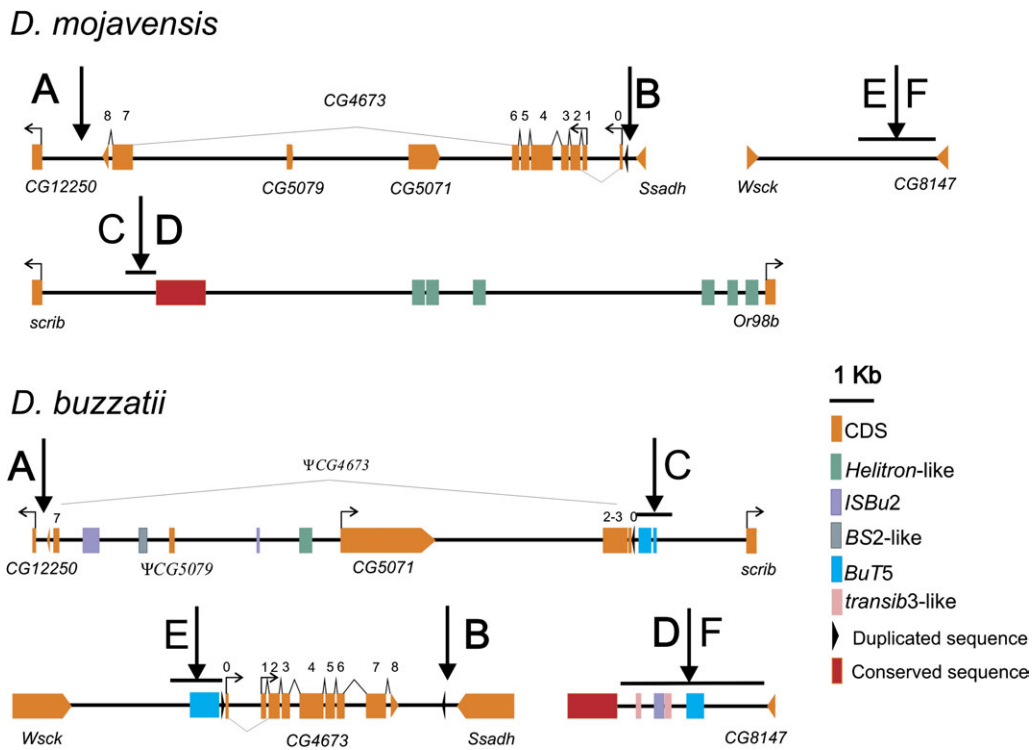
Supplementary Material online). However, *D. buzzatii* gene is considerably shorter than *D. mojavensis* and *D. melanogaster* genes due to a 20-fold size reduction of intron 6 (table 2 and supplementary table S2 in additional file 3, Supplementary Material online). The CG4673 coding sequence is highly conserved ( $\sim 79\%$  identity with *D. melanogaster* and  $\sim 90\%$  identity with *D. mojavensis*; table 2). Estimates of the ratio of nonsynonymous substitutions per nonsynonymous site to synonymous substitutions per synonymous site ( $\omega$ ) average 0.05 indicating that the three transcripts are subject to strong purifying selection (table 2).

The presence of CG4673 in EB region reveals that the region containing this gene was duplicated in the *D. buzzatii* lineage. However, no traces of the two genes (CG5079 and CG5071) nested within intron 6 of *Dmoj*\CG4673 were found within *Dbuz*\CG4673 intron 6 suggesting that these genes were deleted after the duplication. Two pieces of evidence suggest that the duplication of the region containing CG4673 gene occurred at the same time as the inversion. First, this duplication is present in other species of the *buzzatii* complex that also have the *2mn* rearrangement (data not shown) but it is not present in the genome sequence of the other 12 *Drosophila*

species that lack this rearrangement (Clark et al. 2007). Second, based on sequence analyses, we estimated that the duplication took place  $\sim 8$  Ma, which also agrees with the duplication taking place at the same time as the inversion, since the inversion must have happened prior to the divergence of the species of the *buzzatii* complex ( $\sim 8.4$  Ma) and after the divergence between the *buzzatii* complex species from *D. mojavensis* ( $\sim 11.3$  Ma) (Russo et al. 1995; Tamura et al. 2004) (Oliveira DCSG, Almeida FC, O'Grady P, Etges WJ, Armella, R. DeSalle MA, personal communication).

Furthermore, other than the duplication, a microinversion also happened in this genomic region. Taking into account the orientation of the genes in *D. mojavensis*, we would expect *Ssadh* and CG4673 to be in the same orientation in *D. buzzatii*. However, the orientation of CG4673 is reversed relative to what is expected. Note also that the duplicated region is flanked by a 121-bp long duplicated sequence with 72.7% identity ( $E$  value =  $2 \times 10^{16}$ ) (fig. 4). However, this duplicated sequence is found only once in *D. mojavensis* AB region and in *D. buzzatii* AC region, and in both cases, it is located upstream of CG4673 (fig. 4) suggesting that it was not duplicated before the microinversion.





**FIG. 4.** Annotation of breakpoint regions of inversions 2*m* and 2*n* for *D. mojavensis* (AB, CD, and EF) and *D. buzzatii* (AC, EB, and DF). CDS, TE fragments, duplicated and conserved sequences are shown. Lines under black arrows show the breakpoint junctions.

Finally, DF breakpoint in *D. buzzatii* is located in a 4,400 bp segment between *Or98b* and *CG8147* (fig. 4). When the DF sequence was compared with the CD region of *D. mojavensis*, two blocks with significant similarity were detected, one is 408 bp long with 87.7% identity ( $E$  value =  $2 \times 10^{147}$ ) and another one is 93 bp long with 70.9% identity ( $E$  value =  $9 \times 10$ ). Although these conserved sequences do not correspond to the coding region of *Or98b* or any other known gene, they are located upstream of *Or98b* in *D. mojavensis* and might contain enhancers or other regulatory elements for this gene.

*Gene-Associated Sequences*

We looked for highly CNSs and TFBSs in the breakpoint regions to check whether the inversion disrupted or changed their location. Neither CNSs nor TFBSs were found in any of the three breakpoint regions.

*TE Annotation*

In *D. mojavensis*, six blocks of similarity with known *Helitrons* (Feschotte and Pritham 2007) were found in the CD

region, while no similarities to any TE were found either in AB or EF regions (fig. 4). In the three breakpoint junctions of *D. buzzatii*, we found small blocks of similarity with *BuT5* element (fig. 4 and table 3) (Casals et al. 2006). In the *D. buzzatii* AC breakpoint region, two other fragments similar to *ISBu2*, two similar to *BS2*, and finally, two more similar to *Helitron1* were detected upstream of *CG5071*, between the most remote fragments of  $\Psi$ *CG4673* (fig. 4). *ISBu* elements are also *Helitrons* (Yang and Barbash 2008). *BS2* is a *LINE* element (Capy 1998) and the only class I element found in these regions. Finally, another fragment similar to *ISBu2* and two fragments with identity with *transib3* (Kapitonov and Jurka 2003) of *D. melanogaster* were annotated in the DF breakpoint junction.

*Annotation of Other Structural Features*

Using several bioinformatic tools, we searched for structural features in the *D. buzzatii* breakpoint regions, such as AT-rich regions, palindromic sequences, or CCG repeats, that may contribute to chromosomal instability (Kurahashi

**Table 2.** Comparison of *CG4673* Gene of *D. buzzatii* (EB breakpoint region) with *D. mojavensis* and *D. melanogaster*<sup>a</sup>.

Region	<i>Dbuz</i> (base pairs)	<i>Dmoj</i> (base pairs)	<i>Dmel</i> (base pairs)	<i>Dbuz/Dmoj</i>			<i>Dbuz/Dmel</i>		
				Identity (%)	Ka/Ks	$\omega$	Identity (%)	Ka/Ks	$\omega$
RA transcript	3,015	11,726	12,290	—	—	—	—	—	—
RA CDS <sup>b</sup>	1,968	1,962	1,959	90.6	0.0235/0.4218	0.0557	79.6	0.0723/1.4327	0.0505
RB/RD transcript <sup>c</sup>	3,701	12,418	13,317	—	—	—	—	—	—
RB/RD CDS	1,881	1,875	1,875	90.0	0.0259/0.4363	0.0593	78.9	0.0728/1.4874	0.0489

<sup>a</sup> In *D. melanogaster*, *CG4673* encodes three strongly supported transcripts, RA, RB, and RD. In *D. mojavensis*, only RA transcript is annotated; however, a detailed analysis of this region allowed us to annotate the other two transcripts.

<sup>b</sup> Not including UTRs.

<sup>c</sup> Transcripts RB and RD differ in the UTR regions.



**Table 3** TEs detected at the breakpoint regions of inversions 2*m* and 2*n* in *D. buzzatii*.

Region	TE copy <sup>a</sup>	TE class	Coordinates in <i>D. buzzatii</i>	Identity <sup>b</sup> (%)	E-value
AC	<i>ISBu2</i> ( <i>Dbuz</i> \AF368867)	II	1024-1371	306/425 (72.0)	6e-74
	<i>BS2</i> ( <i>Dmel</i> \X77571)	I	2169-2238	55/79 (69.6)	1e-04
			2295-2349	44/55 (80.0)	2e-07
			4766-4819	45/56 (80.3)	3e-08
	<i>ISBu2</i> ( <i>Dbuz</i> \AF368867)	II	4766-4819	45/56 (80.3)	3e-08
	<i>Helitron1</i> ( <i>Dmoj</i> \AY645947)	II	5626-5733	78/109 (71.5)	4e-07
	<i>BuT5</i> ( <i>Dbuz</i> \AF368894)	II	5782-5852	54/71 (76.0)	8e-09
12877-12930			41/54 (75.9)	1e-06	
<i>BuT5</i> ( <i>Dbuz</i> \AF368894)	II	13127-13166	34/41 (82.9)	1e-06	
		13218-13278	43/61 (70.5)	6e-04	
BE	<i>BuT5</i> ( <i>Dbuz</i> \AF368894)	II	5158-5276	88/123 (71.5)	7e-12
			5307-5686	266/404 (65.8)	8e-24
			5730-5787	49/65 (75.4)	2e-06
DF	<i>Transib3</i> ( <i>Dmel</i> \AE003193)	II	1419-1518	73/101 (72.3)	9e-10
			2042-2172	94/133 (70.7)	2e-11
			1824-2027	182/204 (89.2)	5e-68
	<i>ISBu2</i> ( <i>Dbuz</i> \AY900631)	II	1824-2027	182/204 (89.2)	5e-68
	<i>BuT5</i> ( <i>Dbuz</i> \AF368894)	II	2522-2700	122/179 (68.2)	9e-11
			2787-2836	40/55 (72.7)	0.009

<sup>a</sup> The species and the accession number that allowed us to identify each TE copy is given in parentheses.

<sup>b</sup> Only hits with a minimum size of 40 bp and E-value  $\leq 1e-03$  are included in this list.

et al. 2007; Zhang and Freudenreich 2007; Kolb et al. 2009). We did not find any sequence that suggests that the breakpoint regions are predisposed to breaks.

### Analysis of the Breakpoint Regions in the 12 *Drosophila* Sequenced Genomes

The gene arrangements at the breakpoint regions of inversions 2*m* and 2*n* were also investigated in the 12 *Drosophila* sequenced genomes (supplementary fig. S2 in additional file 1, Supplementary Material online). The noninverted arrangement of *D. mojavensis* (AB, CD, and EF) was also found in *D. pseudoobscura* and *D. persimilis* (*Sophophora* subgenus) and *D. virilis* and *D. grimshawi* (*Drosophila* subgenus) suggesting that this is probably the ancestral gene arrangement of the genus. The order and orientation of the genes in the AB breakpoint region is conserved in all 12 sequenced species suggesting that this breakpoint region has been interrupted in *D. buzzatii* and its close relatives only. Region CD has been rearranged two other times: once in the lineage leading to the melanogaster group and once in the lineage leading to *D. willistoni*. Other than in *D. buzzatii*, the EF breakpoint region has also been rearranged in the lineage leading to the melanogaster group of species.

### Expression Analysis of *CG4673* and $\Psi$ *CG4673* in *D. buzzatii*

As a first approximation to the study of the possible functional consequences of 2*mn* rearrangement, we analyzed the expression of *CG4673* and  $\Psi$ *CG4673*. Specifically, we used a combination of bioinformatic and experimental techniques in order to shed light on the expression, molecular structure, and function of *D. buzzatii* *CG4673* gene and  $\Psi$ *CG4673* (see Materials and Methods).

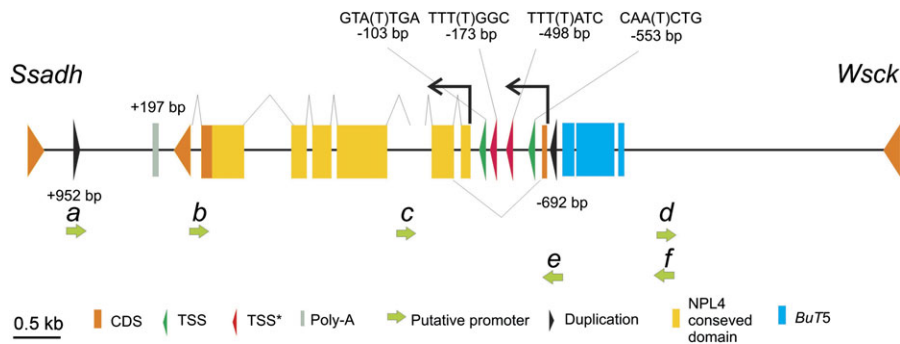
#### Transcription Analyses

We first performed RT-PCR to check whether *CG4673* and/or  $\Psi$ *CG4673* are transcribed (supplementary table S3 in additional file 3, Supplementary Material online). Specific

primers for *CG4673* and for  $\Psi$ *CG4673* were used (see Materials and Methods). RT-PCR reactions performed without retrotranscriptase yield no signal indicating the absence of DNA contamination (results not shown). Amplification products were obtained for both *CG4673* and  $\Psi$ *CG4673* in embryos, larvae, pupae, and adults, revealing that both genes are transcribed in *D. buzzatii* (supplementary fig. S3 in additional file 1, Supplementary Material online).

A Northern Blot was also carried out to detect how many *CG4673*-related transcripts are present in *D. buzzatii* (supplementary fig. S4 in additional file 1, Supplementary Material online). *Drosophila buzzatii* total RNA extracted from different developmental stages was hybridized with a probe amplified with primers designed in exons 5 and 7 of *CG4673*. A clear and unique signal of approximately 2,100 bp, which corresponds to the expected size of *CG4673* transcripts ( $\sim 2,000$  bp; table 2), was observed in the hybridization with total mRNA of *D. mojavensis*. This result indicates that the probe did not hybridize nonspecifically with any other similar sequence, alternative transcript or unspliced mRNA.

A strong hybridization signal similar in size ( $\sim 2,100$  bp) to that obtained in *D. mojavensis* was observed in all the lanes containing *D. buzzatii* mRNA (supplementary fig. S4 in additional file 1, Supplementary Material online). A smaller hybridization signal ( $\sim 1,400$  bp) appears to be partially hidden in the majority of lanes. Finally, another hybridization signal ( $\sim 3,600$  bp) was observed in the lane containing mRNA from pupae. We performed a second Northern Blot hybridization with a lower concentration of mRNA in an attempt to get more defined hybridization bands (supplementary fig. S4 in additional file 1, Supplementary Material online). However, in this second Northern Blot, the  $\sim 2,100$  bp signal was observed only in eggs and pupae. The smaller hybridization signal ( $\sim 1,400$  bp) was clearly observed in pupae, and the  $\sim 3,600$ -bp hybridization signal was not present. Overall, three different transcripts related with *CG4673* (3,600, 2,100, and 1,400 bp long) were observed in *D. buzzatii*.



**Fig. 5.** Structure of the gene *CG4673* in *D. buzzatii*. Polyadenylation signal (Poly-A), NPL4 domain, putative promoters (a–f), and putative TSS are shown. TSS refers to putative start sites found with Neural Network Promoter Prediction software. TSS\* refers to putative start sites inferred from conservation with close species (for details, see text).

We hypothesized that two of the three identified transcripts correspond to the transcripts detected with RT-PCR: the 2,100-bp long transcript corresponds to the functional copy of *CG4673* gene and the 3,600-bp long transcript to an unspliced form of  $\Psi$ *CG4673* (note that the splice sites are not conserved in  $\Psi$ *CG4673*). The third transcript detected in the Northern Blot hybridization (~1,400 bp) could be transcribed from the putative promoter annotated in the duplicated sequence (see below) that is located downstream and oriented toward *CG4673* in *D. buzzatii* BE breakpoint (fig. 3). This antisense transcript would therefore be complementary with both transcripts found in the RT-PCR experiments and the annealing of the sense and antisense transcripts would form dsRNA molecules.

We tested whether we could detect dsRNA by performing RT-PCR with mRNA of *D. buzzatii* previously digested with RNase. RNase only digests single-strand RNA molecules and therefore, RT-PCR would only amplify if dsRNA were present. RT-PCRs were attempted with primers that specifically amplify exons 2, 3, and 7 of  $\Psi$ *CG4673*. While no amplification was observed for  $\Psi$ *CG4673* exon 2, both exons 3 and 7 did amplify when digested samples were used (supplementary fig. S5 in additional file 1, Supplementary Material online). Therefore, exons 3 and 7 of  $\Psi$ *CG4673* of *D. buzzatii* would be protected from digestion with RNase suggesting that dsRNA molecules are present in *D. buzzatii*.

#### Transcription Start and Termination Sites

To identify the transcription start sites (TSS) of *CG4673*, we used Neural Network Promoter Prediction software (Reese 2001). We detected two putative TSS and their corresponding TATA boxes located 30 bp upstream of them (fig. 5). These TSS and TATA boxes were not conserved in *D. mojavensis*, *D. grimshawi*, or *D. virilis*. We also identify putative TSS from sequence conservation with closely related species (*D. mojavensis*, *D. grimshawi*, and/or *D. virilis*). When the sequence upstream of *CG4673* of *D. buzzatii* was aligned with that of these species, two different conserved regions that could also act as TSS were identified with their corresponding TATA box and BR elements (binding site for transcription factors located immediately upstream of the TATA box; fig. 5). 5' RACE experiments confirmed the

presence of several TSS. We obtained a single band product but sequencing of this band resulted in overlapping chromatograms ~200 bp upstream of the start codon.

Using 3' RACE, we identified the transcription end site of *CG4673* in *D. buzzatii* 196 bp downstream of the stop codon (fig. 5). We also attempted to determine the start and end of the putative transcript of  $\Psi$ *CG4673*. However, neither RACE experiments nor software analysis detected the putatively start or end of  $\Psi$ *CG4673* transcript.

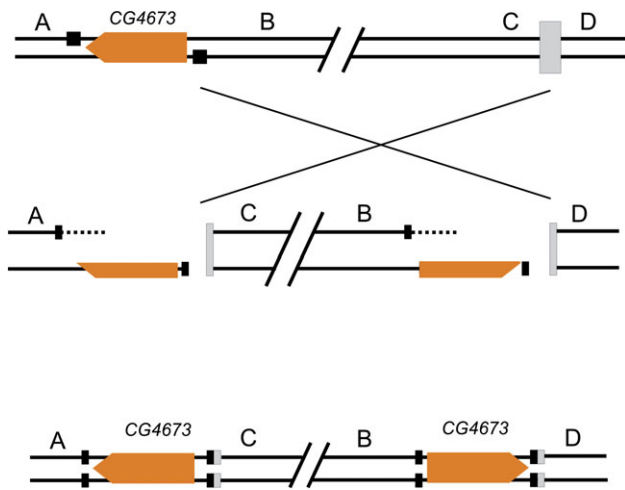
#### Promoter Detection

We used the default threshold of McPromoter software (Ohler 2006) to find putative promoters in the 8.2 kb region between *Ssadh* and *Wsck* genes where *CG4673* is located in *D. buzzatii* BE region. Two promoters, promoter *e* (score = 0.0407) and promoter *f* (score = 0.0506), were identified upstream and in the same orientation as *CG4673* (fig. 5). Promoter *e* is located in the 121-bp duplicated sequence. Another four putative promoters were found in reverse orientation (fig. 5). Promoter *b* (score = 0.0351) and promoter *c* (score = 0.0367) were annotated in exonic sequence and are likely artifacts (note that these are the two promoters with lower scores). Promoter *d* (score = 0.0445) was located upstream of *CG4673* and oriented toward *Wsck* gene. The most significant promoter, promoter *a* (score = 0.0555) was located in the 121-bp duplicated region in the 3' intergenic region. Therefore, both duplicated regions in the BE breakpoint contain potential promoters.

No putative promoters were detected with McPromoter software in direct or reverse orientation for  $\Psi$ *CG4673* (AC breakpoint region).

#### *CG4673* Function

*Dbuz/CG4673* has a 79% nucleotide sequence identity with *Dmel/CG4673* (table 2). In *D. melanogaster*, *CG4673* has a zinc finger and a NPL4 domain (supplementary Table S4 in additional file 3, Supplementary Material online), and it is involved in nuclear transport and in the ubiquitin-proteasome degradation pathway (Bays and Hampton 2002). We searched for conserved domains in the putative protein encoded by *Dbuz/CG4673* (Marchler-Bauer et al. 2009). We found similarity with a COG5100 domain (exons 2–7; *E* value =  $3 \times 10^{103}$ ), which corresponds to the NPL4 protein (Bays et al. 2001; Botta et al. 2001). This domain is



**Fig. 6.** Schematic representation of the mechanism generating inversion  $2m$ : single strand breaks (black squares) and double strand breaks (gray squares) followed by NHEJ repair.

composed of two subdomains: an active NPL4 domain (pfam05021,  $E$  value =  $9 \times 10^{92}$ ) and a zinc finger of the NPL4 superfamily (pfam05020,  $E$  value =  $1 \times 10^{65}$ ). The conservation of the functional domains in *D. melanogaster* and *D. buzzatii* proteins, suggest that CG4673 is also involved in nuclear transport and in the ubiquitin-proteasome degradation pathway in *D. buzzatii* (supplementary table S4 in additional file 3, Supplementary Material online).

## Discussion

Detailed analyses of inversion breakpoints can help to explain not only processes related to chromosome evolution, such as whether breakpoint use is random or nonrandom, but also provide insight into the functional consequences of inversions. In this work, we have sequenced the inversion breakpoints of the  $2mn$  rearrangement fixed in all species of the *buzzatii* complex. Our results shed light on the mechanism generating this rearrangement, corroborate the breakpoint reuse at the molecular level, and start uncovering its functional consequences.

### Generation of Inversions $2m$ and $2n$

Chromosomal inversions may be generated by diverse mechanisms and the features observed in the inversion breakpoints often provide clues for the mechanisms involved, supporting some of them and rejecting others. In this work, we found that the  $2m$  inversion is associated with a  $\sim 13$  kb duplicated segment present at both breakpoints and was likely generated by staggered breaks followed by repair by NHEJ as depicted in figure 6. The phylogenetic distribution of the duplication and the estimate of the age of the duplication (based on sequence divergence analyses) are consistent with the duplication co-occurring with the inversion.

The hybrid element insertion model (Gray et al. 1996; Gray 2000; Zhang and Peterson 2004) could explain inversion  $2m$  and its associated duplication also as a single event

if the participating *BuT5* copies were originally inserted 5' and 3' of CG4673 in homologous chromosomes (supplementary fig. S6 in additional file 1, Supplementary Material online). However, this mechanism has only been described for a few transposon families and under particular conditions. Furthermore, *BuT5* copies might have inserted in the  $2m$  inversion breakpoint regions as secondary colonizers after the generation of the inversion as it has been reported previously (Cáceres et al. 2001; Casals et al. 2003; Delprat et al. 2009). Therefore, although an involvement of *BuT5* in the generation of inversion  $2m$  cannot be completely ruled out, we find the evidence unconvincing.

Alternative mechanisms, such as duplicative transposition of the  $\sim 13$  kb duplicated segment into the CD region followed by an event of ectopic recombination between the duplicated segments (inverting the region between the duplications) or duplication of the  $\sim 13$  kb region first and then the inversion breakpoint happening right in between of the two duplicated copies, are also possible. However, these two mechanisms are less parsimonious since they would require two independent events (the duplication and the inversion). Moreover, transposition is not as common as other types of rearrangements in *Drosophila* (Ranz et al. 2003; González et al. 2004; Bhutkar et al. 2007).

We have found few clues to explain the generation of inversion  $2n$ . This inversion could have been generated by double strand breaks at both breakpoints and repair by NHEJ. As a matter of fact, as inversion  $2m$  and  $2n$  have never been found in isolation, we can not rule out the possibility that the two inversions were generated concurrently by the same mechanism (a three-point breakage followed by NHEJ repair). On the other hand,  $2n$  might have resulted from ectopic recombination between *BuT5* copies, as fragments of this transposon were found at both  $2n$  breakpoints (BE and DF). However, the evidence is weak because these fragments are highly degraded and currently lack target site duplications.

The fixed inversions studied here are old (between  $\sim 8.4$  and 11.3 Ma) and some footprints of the events that generated them are not apparent anymore. This shortcoming could be overcome by studying younger polymorphic inversions. However, the study of fixed inversions should provide insights into what features make inversions successful whereas the evolutionary fate of polymorphic inversions is not known (as they might eventually be lost).

### Breakpoint Reuse

Our molecular characterization of  $2mn$  breakpoint regions has corroborated the breakpoint reuse previously suggested in this region (Ruiz and Wasserman 1993; González et al. 2007). If  $2m$  and  $2n$  were independent inversions but separated by a short distance, a small segment of the original CD region would be found in the middle breakpoint region of the derived chromosome. If  $2m$  and  $2n$  were overlapping by a short segment, this small segment would have been transferred between the distal and proximal breakpoints. None of these expectations hold true, that is, we could neither detect a small segment remaining in the



middle breakpoint nor a small segment exchanged between distal and proximal breakpoints (fig. 4).

The detailed analysis of the *2mn* breakpoint regions in *D. buzzatii* allowed us to identify an additional rearrangement, a microinversion, that took place in the EB breakpoint region. Furthermore, the comparison of the three *2mn* breakpoint regions in the 12 *Drosophila* species sequenced (Clark et al. 2007) revealed that while AB region is conserved in the 12 species, CD and EF regions have been rearranged at least once during the evolution of the *Drosophila* genus (supplementary fig. S2 in additional file 1, Supplementary Material online) suggesting that these breakpoint regions have been repeatedly reused during evolution.

Although breakpoint clustering at the cytological level is common in *Drosophila* (González et al. 2007), only two other breakpoint reuse regions have been previously characterized at the sequence level (Richards et al. 2005; Ranz et al. 2007). Recurrent breakage might be revealing structural instability of these particular regions. However, the analyses of the breakpoint regions in *D. buzzatii* did not provide evidence for sequence features previously associated with fragile regions, such as AT-rich regions, CCG repeats, or palindromic sequences (Kurahashi et al. 2007; Zhang and Freudenreich 2007; Kolb et al. 2009). We neither found evidence for a high density of TE insertions in these regions. Alternatively, breakpoint reuse might be due not to an increase rate of generation of inversions in particular regions but to differential survival of inversions. Future analyses are necessary to unveil the actual causes of breakpoint reuse.

### Functional Analyses of the Breakpoint Regions

The detailed analyses of the breakpoint regions also allowed us to shed light on the possible functional consequences of the *2mn* rearrangement. We found that none of the breakpoints disrupt a coding region (fig. 4). Breakpoints could also affect the expression of nearby genes by disrupting, changing the location, or creating *cis*-regulatory elements. Although we did not find any evidence for CNSs or TFBSs in the breakpoint regions, the existence of species-specific regulatory regions cannot be ruled out since no polymorphism data is available for *D. buzzatii*.

Inversion *2m* is associated with a 13 kb duplication of the region located between CG12250 and *Ssadh* and containing genes *CG4673*, *CG5071*, and *CG5079* (fig. 4). *CG5071* and *CG5079* are nested within intron 6 of *CG4673* and are flanked by the same two genes in all the other *Drosophila* genomes sequenced (Clark et al. 2007) indicating that this is the ancestral organization. However, the duplication of the region located between CG12250 and *Ssadh* and containing *CG4673*, *CG5071*, and *CG5079* did not eventually lead to the duplication of any functional gene. In one of the duplicated fragments (AC breakpoint; fig. 4), *CG4673* and *CG5079* are now pseudogenes, while in the other duplicated fragment (BE breakpoint; fig. 4) *CG5071* and *CG5079* have been lost. Therefore, *2mn* rearrangement eventually led to the net loss of one gene (*CG5079*). Although *CG5079* is not associated with any gene ontology

terms, RNAi analyses indicate that it might play a role in cell growth and viability (Boutros et al. 2004) suggesting that the loss of this gene might have functional consequences in *D. buzzatii* (supplementary table S4 in additional file 3, Supplementary Material online).

The ancestral copy of *CG4673* has seemingly lost its function, while the derived copy is functional. This derived copy is much shorter than the ancestral one due to the loss of the two nested genes initially located in intron 6. Gene expression levels are negatively correlated with intron length, likely as a result of selection for transcriptional speed or economy (Castillo-Davis et al. 2002; Seoighe et al. 2005). As *CG4673* is a highly expressed gene in *D. melanogaster*, we hypothesized that the loss of ~9 kb in *D. buzzatii* increased the efficiency of transcription of this gene resulting in the conservation of the derived copy while the ancestral copy was allowed to degenerate.

Although the functional benefits of nested genes are not clear, gains of nested gene structures are more common than losses in *Drosophila* and other metazoans (Assis et al. 2008). It has been hypothesized that nested host gene structures present a mechanism for the coordinated regulation of functionally related gene pairs. However, many nested genes exhibit functions and expression patterns distinct from those of host genes suggesting that the gain of nested gene structures is neutral (Assis et al. 2008; Kumar 2009). This might also be the case for *CG4673* and its two nested genes: *CG4673* is involved in nuclear transport and in protein degradation while *CG5071* has enzymatic activity and *CG5079* might play a role in cell growth and viability (supplementary table S4 in additional file 3, Supplementary Material online). The level of expression of these three genes is not correlated (supplementary table S4 in additional file 3, Supplementary Material online). As hypothesized above, in *D. buzzatii* the loss of the nested structure in the derived copy might have contributed to the efficiency of expression of *CG4673* leading to the degeneration of the ancestral copy of this gene.

*CG4673* is flanked by a 121-bp long duplication that has promoter capability (fig. 5). We suggest that this 121 bp duplication was generated concurrently with the microinversion as a result of staggered single-strand breaks and NHEJ repair because the duplication is present only once in *D. mojavensis* AB and *D. buzzatii* AC regions (fig. 4). We further hypothesized that this 121-bp long sequence could be driving the transcription of both sense and antisense transcripts since the two copies are oriented toward *CG4673*, and we found that this was the case. Antisense transcripts can exert both positive and negative effects on gene regulation at different levels (Faghihi and Wahlestedt 2009; Werner and Swan 2010). For example, antisense transcription, but not the antisense RNA molecule itself, can modulate transcription of the sense RNA through transcriptional collision (Prescott and Proudfoot 2002). Antisense transcripts can also affect gene expression through DNA–RNA interactions or through the formation of RNA duplexes (Faghihi and Wahlestedt 2009). We indeed found evidence for the existence of *CG4673* dsRNA in *D. buzzatii*. However,



further analyses are necessary in order to elucidate whether dsRNA formation has an inhibitory influence on the sense transcript, for example, by inhibiting its translation, or on the other hand increases the expression of the gene, for example, by reducing mRNA decay (Faghihi and Wahlestedt 2009). In any case, our findings suggest that besides the *2mn* rearrangement, the microinversion of *CG4673* also had functional consequences. We also found evidence for transcription of  $\Psi$ *CG4673*. Transcription of pseudogenes is not uncommon (Zheng and Gerstein 2007). Whether this transcription is spurious or has a biological function remains to be determined.

Finally, in *D. buzzatii*, *CG5071* CDS is considerably longer than in *D. mojavensis* (668 aa and 225 aa long, respectively) but similar to *D. melanogaster* transcript RB (680 aa long). In *D. melanogaster*, this gene is involved in protein folding and shows moderately high expression in males (supplementary table S4 in additional file 3, Supplementary Material online). Sequence conservation between *D. buzzatii* and *D. melanogaster* proteins (74% identity) suggest that the function of this gene might be conserved in *D. buzzatii*.

In summary, the *2mn* rearrangement fixed in all species of the *buzzatii* complex is currently associated with the loss of one gene (*CG5079*), the loss of a nested gene structure (*CG4673* does not longer contain any other gene), and changes in gene structure and gene regulation (*CG4673*). Any of these changes or the combination of them might have been adaptive and therefore might have driven this rearrangement to fixation. Alternatively, this rearrangement could have increased in frequency through meiotic drive. *CG4673* is involved in nuclear transport, and one of the best studied meiotic drive systems in *Drosophila* (SD system) comprises, as the main part of the driving chromosomes, a truncated duplicate of a nuclear transport protein, *Sd-RanGAP*, a short satellite region and often chromosomal rearrangements that restrict recombination between its components (Kusano et al. 2003; Presgraves et al. 2009).

## Conclusion

Our detailed analyses of the breakpoints of the *2mn* rearrangement present in all the species of the *buzzatii* complex have shown that other than the expected gene order change, this complex rearrangement was associated with changes in gene number, gene structure, and with the generation of an additional microinversion that has potentially functional consequences since it leads to the generation of an antisense transcript. These results indicate that 1) functional consequences of inversions are much more complex than those expected from a simple two-break rearrangement and that 2) in order to fully describe the genomic consequences of structural variants, genomic approaches should be complemented with detailed sequence-level and functional analyses of the breakpoint regions.

## Supplementary Material

Supplementary Figures S1 to S6, Supplementary Data, and Supplementary Table S1\_to\_S9 are available at *Molecular*

*Biology and Evolution* online (<http://www.mbe.oxfordjournals.org/>).

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