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Specific Activation of an I-Like Element in Drosophila Interspecific Hybrids

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Abstract

The non-long terminal repeat (LTR) retrotransposon I, which belongs to the I superfamily of non-LTR retrotransposons, is well known in Drosophila because it transposes at a high frequency in the female germline cells in I-R hybrid dysgenic crosses of Drosophila melanogaster. Here, we report the occurrence and the upregulation of an I-like element in the hybrids of two sister species belonging to the repleta group of the genus Drosophila, D. mojavensis, and D. arizonae. These two species display variable degrees of pre- and postzygotic isolation, depending on the geographic origin of the strains. We took advantage of these features to explore the transposable element (TE) dynamics in interspecific crosses. We fully characterized the copies of this TE family in the D. mojavensis genome and identified at least one complete copy. We showed that this element is transcriptionally active in the ovaries and testes of both species and in their hybrids. Moreover, we showed that this element is upregulated in hybrid males, which could be associated with the male-sterile phenotype.

Key words: hybrids, non-LTR retrotransposon, repleta group, phylogeny.

Introduction

Virtually all species harbor transposable elements (TEs), sequences that are able to move within genomes and that differ in structure, activity, and copy number between populations and species. TEs are classified in two broad classes: Class I and Class II, based on their mode of transposition. Class I is composed of two types of elements that transpose through RNA intermediates: elements with long terminal repeats (LTRs) and elements without these direct repeats, the non-LTR elements, which can further be divided into two types: the long interspersed nuclear elements (LINEs) and the short interspersed nuclear elements (SINEs). Class II (DNA transposons, rolling circle transposons, and miniature inverted repeat elements) is composed of TEs that transpose directly through a DNA molecule (Wicker et al. 2007; Kapitonov and Jurka 2008).

The LINEs are among the oldest genetic elements in eukaryotes and are the most abundant in the human genome (Fawcett et al. 1986; Simonelg et al. 1988; Bucheton et al. 1992; Han 2010). They consist of a 5′-untranslated region (UTR), which has promoter activity, two open-reading frames (ORFs) separated by a spacer, and a 3′-UTR with a poly-A tail. Our study focused on one I-like element, a LINE element first described in Drosophila melanogaster at the beginning of the 1970s (Picard 1976). The I element (originally called the I factor) is responsible for the hybrid dysgenesis syndrome in D. melanogaster females. This species presents two types of strains based on the presence and activity of this element: The I strains, harboring complete and functional I elements, and the R strains, with nonfunctional I elements. Crosses between I males (Inducer) and R females (Reactive) result in high rates of transposition in the germline of female offspring, known as SF females. These females are sterile because most of the laid eggs fail to hatch due to the very high frequency of I element transposition, which causes genetic abnormalities such as failure in meiotic divisions and chromosomal rearrangements (Bucheton et al. 1976, 1984; Bucheton
The number of \( I \) copies in \( D. melanogaster \) is approximately 30, but only five functional sequences are known to be dispersed on the chromosomal arms. The complete and functional copies are 5.4 kb long, possess two long ORFs with cysteine-rich motifs, apurinic–apyrimidic endonucleases, reverse transcriptase (RT), and RNase H domains, and the 3′-end has several TAA repeats. These copies are expressed in the nurse cells of dysgenic ovaries, and the transcripts are transported into the oocytes, where retrotransposition occurs (Pelisson and Picard 1979; Fawcett et al. 1986; Vaury et al. 1990; Seleme et al. 1999, 2005). In \( Drosophila \), piwi-interacting RNAs (piRNAs) are important posttranscriptional regulators that are responsible for TE mRNA degradation (Khurana and Theurkauf 2010; Senti and Brennecke 2010; Kelleher et al. 2012; Akkouche et al. 2013; Dufourt et al. 2013). Specifically, the piRNAs are implicated in maternal \( I \) element control in \( D. melanogaster \) (Brennecke et al. 2008; Chambeyron et al. 2008).

The implication of TEs in hybrid dysgenesis is one of the examples in which transposition may drive population isolation and is often proposed as one of the first steps of speciation (Kidwell and Novy 1979; Fontdevila 2005). However, the TE dynamics of expression and transposition have not been well described during this process. Hence, several questions remain unanswered regarding this subject. If TEs act in the first steps of speciation, what happens when crossing very closely related species that still have incomplete reproductive isolation? In particular, are \( I \)-like elements associated with the incompatibility observed, and are they related to high rates of hybrid transposition as reported in other species (Ungerer et al. 2009; Cavallini et al. 2010; Moschetti et al. 2010; Kelleher et al. 2012; Vela et al. 2014)? In this report, we made a first investigation to check whether \( I \)-like elements could potentially be involved in hybrid sterility.

\( Drosophila mojavensis \) and its sister species analyzed in this study—\( D. arizonae \)—are a perfect pair of species with which to address these questions. They are found in the desert of the southwestern United States and Mexico and share sympatric areas in southern Arizona and the State of Sonora (Mexico). These two species present three interesting features that facilitate the investigation of the above questions: 1) Their hybrids can be produced in the laboratory; 2) the genome of \( D. mojavensis \) has been sequenced, which allow us to analyze the sequences of its TEs; and 3) these species exhibit variable degrees of pre- and postzygotic isolation (Wasserman and Koepfer 1977; Reed and Markow 2004; Massie and Markow 2005). Moreover, the male offspring of \( D. arizonae \) females and \( D. mojavensis \) males is sterile, but in the reciprocal cross, the hybrids have motile sperm depending on the origin of the \( D. mojavensis \) population that is used in the crosses (Reed and Markow 2004). Additionally, the strong prezygotic isolation is higher between flies from sympatric areas than between the allopatric flies, but the isolation level depends in part on the direction of the crosses and the geographic origin of the populations (Wasserman and Koepfer 1977; Ruiz et al. 1990; Reed and Markow 2004; Massie and Markow 2005).

We found four potential full-length copies of \( I \)-like elements and reconstructed their phylogenetic relationships with other \( I \) family members in \( Drosophila \) species, thus establishing the vertical and ancestral inheritance of the \( I \) element of \( D. mojavensis \) from a common ancestor shared with the \( D. melanogaster \) group. We also showed that the \( D. mojavensis \) \( I \)-like sequences are transcriptionally and transpositionally regulated in the hybrid female germline. However, a significant increase in transcription was detected in testes from hybrids coming from \( D. arizonae \) mothers. This is the first report that shows the activity of \( I \)-like elements in the male germline, and it suggests a link with the male-sterile phenotype observed in the male hybrids of the studied species.

### Materials and Methods

#### Genomic Analyses

To investigate the occurrence of \( I \)-like elements in the \( D. mojavensis \) sequenced genome, we used the canonical \( I \) sequence from \( D. melanogaster \) (\( I_{DM} \)), which is available in the Repbase library Rel. 16.06, v.4 (Fawcett et al. 1986). This sequence was blasted against the scaffolds of the \( D. mojavensis \) genome (version-70, 13/dmoj_caf1) on the Ensembl MetaZoa platform using BLASTN \( D. mojavensis \) (http://metazo.ensembl.org/index.html, last accessed July 4, 2014).

The hits were manually curated, and only the largest and putative complete copies (more than 2,000 bp) were selected. We then selected the \( D. mojavensis \) copies that were over 80% identical between them. The sequences with the above characteristics were selected and analyzed using the ORF-FINDER web tool (http://www.ncbi.nlm.nih.gov/projects/gorf/, last accessed July 4, 2014) to identify putative coding regions. The ORFs identified were compared and confirmed by BLAST search for protein sequences at the NCBI website, and the conserved protein domains were checked on the conserved domain database (http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml, last accessed July 4, 2014). The sequence with the most complete structure was selected and hereafter referred to as “Reference Copy” or “RC” (fig. 1).

The RC was used to search for \( I \)-like element copies in the \( D. mojavensis \) genome with identity (ID) higher than 80% and length greater than 200 bp. This criterion was chosen based on the 80-80-80 rule for TE family determination, as proposed by Wicker et al. (2007) for the identification of TEs belonging to the same family. The position of each copy (start and end)
on the genome scaffolds, the length of each fragment aligned by BLAST, and the number of the unidentified nucleotides (N) were recorded (table 1). We performed multiple alignments of the gag and RT domains (nt) identified in the copies, and we determined the ID of each domain, based on those of the RC, using Bioedit version 7.0.4.1 (Hall 1999). A drawing of the copies is shown in figure 2 and represents the integrity of the putatively full-length copies compared with RC. To confirm that we recovered unique copies in the genome, the 5’- and 3’-end flanking sequences of each copy were retrieved, and their identities were determined compared with the RC flanking sequences. All copies corresponded to different insertions.

To check the functional organization of all putatively full-length copies of D. mojavensis, the 5’-end sequences were compared with the sequence of the I promoter of D. melanogaster (Minchiotti et al. 1997) using BLAST2seq.

The chromosomal location of the most complete I copies was inferred from the scaffold coordinates of each copy, and the correspondence of scaffolds with D. mojavensis polytene chromosome maps was performed using information found in Flybase.org (http://flybase.org/maps/chromosomes/maps.html, last accessed July 4, 2014) and in Schaeffer et al. (2008). For an intrachromosomal distribution of the copies, each D. mojavensis chromosome (X, 2, 3, 4, 5, and 6) was divided into three regions: Distal (10% of the sequence), central (80% of the sequence), and proximal (or centromeric, 10% of sequence) segments in relation to the position of the centromere, to test the distribution of the copies in these regions (Marzo et al. 2013).

The nomenclature used to identify each copy followed Wicker et al. (2007); for example, RII_DMJA1 consists of RII followed by the number of each copy within each clade. For an intrachromosomal distribution of the copies, each D. mojavensis chromosome (X, 2, 3, 4, 5, and 6) was divided into three regions: Distal (10% of the sequence), central (80% of the sequence), and proximal (or centromeric, 10% of sequence) segments in relation to the position of the centromere, to test the distribution of the copies in these regions (Marzo et al. 2013).

Evolutionary Analyses of RT Sequences

We reconstructed the evolutionary relationships between the D. mojavensis/D. arizonae I-like sequences with all the I superfamily members of Drosophila species deposited in the Repbase library (http://www.girinst.org/repbase/, last accessed July 4, 2014). Each retrieved element was analyzed using ORF FINDER (http://www.ncbi.nlm.nih.gov/orf/gorf.html, last accessed July 4, 2014), and only the amino acid sequences of the second ORF were used for further analysis. We identified the RT domain in each protein sequence using the structural domain bank from the Pfam platform (http://pfam.sanger.ac.uk/, last accessed July 4, 2014). The Drosophila Repbase RT domains were aligned with the D. mojavensis and D. arizonae RT domains using MEGAS (Tamura et al. 2011). For D. mojavensis (strains #15081-1352.01 and #15081-132.26 from US San Diego Drosophila Stock Center) and D. arizonae (strains #15081-1271.17 and #15081-1271.18 from US San Diego Drosophila Stock Center), we amplified and sequenced the 627 bp RT sequence (RT_IF: 5’-GCCTAGTCATCCTATC-3’ and RT_IR: 5’-TGTTAGC GCCGTTTGTATT-3’), corresponding to the position 2,813 and 3,198 in the D. mojavensis RC sequence. Polymerase chain reaction (PCR) cycling parameters were: 94 °C for 3 min, 40 cycles of 94 °C for 45 s, 56 °C for 45 min, and 72 °C for 1 min, followed by a 72 °C for 7 min. The fragments obtained were purified directly from the PCR product, using the GFX PCR DNA and Gel Band Purification Kit (GE Healthcare), and cloned with the TOPO TA Cloning Kit (Invitrogen). Five randomly chosen clones were sequenced using the M13 universal primers.

We performed the amino acid phylogenetic reconstruction using the maximum-likelihood method, JTT model, and 500 bootstrap replicates as implemented in PHYML 3.0 (Guindon et al. 2010) at the ATGC: Montpellier Bioinformatics platform (http://www.atgc-montpellier.fr/, last accessed July 4, 2014). Two sequences of other LINE superfamilies were used as out-groups: The D. melanogaster Fw non-LTR retrotransposon (GenBank accession number: M17214.1) and the D. yakuba Helena non-LTR retrotransposon (GenBank accession number: AF012049.1).

RNA Extraction and RT–quantitative PCR

The fly samples were obtained from the US San Diego Drosophila Stock Center. We used strains of D. mojavensis...
Table 1
Summary of the I-like Element Copies in the Drosophila mojavensis Genome

| Copy            | Acc.       | Scaff.  | Start S. | End S. | Start Q. | End Q. | Or.     | Chr.arms | Region | N   | % (ID) | % (ID Flanq.) | Leng. | Integ. | % (ID) |
|-----------------|------------|---------|----------|--------|----------|--------|---------|----------|--------|------|--------|---------------|-------|--------|--------|
| RII_DMJA1(RC)a  | CH933806.1 | 6540    | 12533998 | 12539232 | –        | 2(E)   | Central | 0        | 5,382  | 5,382 | 0.999  | 0.433         | Comp. |        |        |
| RII_DMJA2a      | CH933808.1 | 6496    | 21912581 | 21918262 | 1785     | 1052   | 5(C)    | Central | 1,027  | 4,505 | 0.999  | 0.433         | Comp. |        |        |
| RII_DMJA3       | CH933807.1 | 6500    | 11558188 | 11563761 | 1        | 5382   | + 3(B)  | Central | 321    | 4,948 | 0.9981 | 0.43           | Comp. |        |        |
| RII_DMJX1       | CH933815.1 | 6482    | 430575   | 438784  | 1        | 5382   | + N/A   | N/A      | 5,092  | 2,808 | 0.996  | 0.447         | Comp. |        |        |
| RII_DMJX2       | CH935525.1 | 12      | 175      | 1389    | 1        | 4456   | – N/A   | N/A      | 0      | 1,390 | 0.99   | B/#           | Def.  |        |        |
| RII_DMJX3       | CH937882.1 | 97      | 808      | 1490    | 4702     | 5382   | – N/A   | N/A      | 0      | 681   | 0.99   | B/#           | Def.  |        |        |
| RII_DMJX4       | CH933809.1 | 6680    | 15471148 | 15472992 | 3547     | 5382   | + 4(D)  | Central | 0.997  | 1,848 | 0.99   | 0.39          | Def.  |        |        |
| RII_DMJX5       | CH93813.1  | 6498    | 3343118  | 3344746  | 3755     | 5382   | + N/A   | N/A      | 4049   | 1,629 | 0.996  | 0.391         | Def.  |        |        |
| RII_DMJX6       | CH933808.1 | 6496    | 9570337  | 9570944  | 4774     | 5382   | + 5(C)  | Central | 0.998  | 609   | 0.99   | 0.417         | Def.  |        |        |
| RII_DMJX7       | CH93814.1  | 6308    | 2487507  | 2487762  | 5128     | 5382   | – X(A)  | Telomeric | 0.984  | 256   | 0.984  | B/#           | Def.  |        |        |

NOTE.—Acc., accession number in GenBank; Scaff., scaffold; Start S., scaffold start hit; End S., scaffold end hit; Start Q., query start hit; End Q., query end hit; Or., orientation of the copy in genome; Chr.arms, chromosome arms and conventional numbering with Muller syntenic elements in parentheses; Region, chromosome region; N, number of unidentified nucleotides; % ID, nucleic percent ID to RC; % ID Flanq., nucleic percent ID the 5'-flanking region to RC flanking region; Leng., total alignment length; Integ., copy integrity: Comp., putatively complete; Def., defective copies; N/A, not annotated; #, not computed.

*aSequence used in phylogeny (fig. 4).

Fig. 2.—Structure of the putatively complete copies of the I-like non-LTR retrotransposon identified in the Drosophila mojavensis genome. DMJA1 to X1, copies shown in table 1; shaded gray boxes, unidentified nucleotides (N); red boxes, 5'-end promoter position; green boxes, ORF1; blue boxes, ORF2; scale size range of the copies is presented in base pairs.

(Anza Borrego Desert, CA; stock number: 15081-1352.01) and D. arizonae (Metztitlan, Hidalgo, Mexico; stock number: 15081-1271.17) for crosses. These two allopatric strains present incomplete prezygotic isolation, and we could obtain enough F1 hybrids for both reciprocal crosses. In crosses between D. arizonae females and D. mojavensis males, F1 males are sterile. These populations were maintained at 25 °C for mass cultures. One- to two-day-old virgins (females and males) were isolated from the parental lines and split in vials. Subsequently, the crosses were performed (D. mojavensis females × D. arizonae males and the reciprocal cross) using ten couples per cross. F1 hybrids were isolated, and 5-day-old females and males were used for DNA and RNA extraction.

The flies were dissected, and their tissues were placed in 1× phosphate buffer saline. The RNA was extracted from pools of 20 pairs of ovaries and from 30 testes from parental and hybrids. Two independent sets of crosses were performed. For each, three replicates were performed to extract total RNA using the RNeasy kit (Qiagen) and treated with DNase (the DNA-free kit; Ambion). One microgram of the total RNA was then converted into cDNA using the Thermoscript Invitrogen kit primed with a mix of oligo(dT)20 and random primers.

The expression level of the RT of the I-like elements was measured by RT–quantitative (qPCR) using the following primers:

Forward 5' -ATC CAC TCT TCA ACG GCA TC-3'
Reverse 5' -TGG ACG ATA TGG TGC AAA TG-3'

We verified by sequencing (the TOPO TA cloning kit; Invitrogen) that no sequence polymorphism was present in the primers that could affect the PCR efficiency. Five clones of each strain were sequenced (Applied Biosystems 3730XL), and the sequences obtained were aligned using BLAST alignment at the NCBI website. The cDNA was diluted 50 times, and the relative mRNA level was quantified using SYBR green qPCR in a LightCycler 480 apparatus (Roche Diagnostics). The transcript quantity was estimated relative to the rp49 reference gene of D. mojavensis using the primers described by Granzotto et al.
Fluorescent In Situ Hybridization

I transcripts were detected by in situ hybridization in 5-day-old ovaries of parents and hybrids according to Akkouche et al. (2012). Ovaries were hybridized with a 952-bp riboprobe that corresponded to the RT and RNA-H domains in the RC and included a T3 promoter site in the reverse primer (forward: 5'-GCA ATA CAA CCG GCC GTA ACA-3', reverse: 5'-TGG CTG TGG ATT TGG CTG TGA ATT AAC CCT CAC TAA AGG GA-3'). For the in vitro transcription, a digoxigenin (DIG)-RNA labeling mix (Roche) was used. After hybridization and stringency washes, an anti-DIG POD antibody mix (ROCHE) was used for chemiluminescent detection, with a fluorescence amplification (the Tyramide Signal Amplification Kit; PerkinElmer). The DNA was labeled by SYTOX Green Nucleic Acid Stain (Life Technology), and the visualization was performed with a Zeiss LSM510 Meta confocal microscope. Tissues without anti-DIG treatment were used as a negative control, since no label is expected without this secondary antibody. Five-day-old testes hybridization was performed with a Zeiss LSM510 Meta confocal microscope. The product was denatured for 5 min in 92 °C and cooled at room temperature. The linker HindIII–MSEB was ligated to digested DNA with a T4 DNA ligase (10 units), enzyme buffer (5×), and digested DNA in a 50-μl reaction for 3 h at 23 °C. DNA was amplified with a primer adapter (LNP 5'-GAA TTC GTC AAC ATA GCA TTT CT-3') and a forward primer inside of the 3'-end of the I element (5'-TAA CGG TAG ATT AAC CCT CAC TAA AGG GA-5') (supplementary fig. S1, Supplementary Material online).

The TD technique also allowed us to have an estimation of the copy number in each strain. The genomic DNA (100 ng/μl) was digested with HindIII (10 U/μl, 3 h at 37 °C). The MSBE adapter was linked with the HindIII enzyme adaptor site in one double-stranded reaction (HindIII–MSEB) using 100 μM of Hindlink, 100 μM of MSBE, SSC 20×, and 1 M Tris in total volume of 333 μl. The product was denatured for 5 min in 92 °C and cooled at room temperature. The linker HindIII–MSEB was ligated to digested DNA with a T4 DNA ligase (10 units), enzyme buffer (5×), and digested DNA in a 50-μl reaction for 3 h at 23 °C. DNA was amplified with a primer adapter (LNP 5'-GAA TTC GTC AAC ATA GCA TTT CT-3') and a forward primer inside of the 3'-end of the I element (5'-TAA CGG TAG ATT AAC CCT CAC TAA AGG GA-5') (supplementary fig. S1, Supplementary Material online). The TD method was tested with D. mojavensis genomic DNA, extracted from a single flies, that was digested by HindIII and amplified by PCR with HindIII–MSEB and I primers. Then, two fragments (842 and 450 bp), identified in silico and observed in electrophoresis, were cloned (with five clones for each fragment), amplified by PCR with M13 universal primers, and sequenced. The alignment of these sequences against the D. mojavensis RC confirmed the veracity of the fragments obtained by enzyme digestion and amplification for the I element.

Results

Our study revealed the presence of an I-like element in D. mojavensis and D. arizonae that is closely related to the I element found in the genomes of the D. melanogaster subgroup. In silico and in vivo analyses showed the occurrence of complete and transcriptionally active copies in both species and in their hybrids. The I Element of D. mojavensis

We detected 10 I-like copies in the D. mojavensis genome sequence (table 1). One potentially complete copy, called RC (RII_DMJ1A1), is 5,382-bp long and contains two ORFs (ORF 1 and ORF 2) with no internal stop codons and with all the expected protein domains for this type of non-LTR retrotransposon (Malik et al. 1999; fig. 1). It has 62.42% nucleic ID with the D. melanogaster I element and is only 7 bp longer. We identified three other copies that are most likely complete: RII_DMJ2A, RII_DMJ3A, and RII_DMJ1X (table 1 and fig. 2), which show high ID and conserved protein domains compared with the RC, but their sequences were interrupted with unidentified nucleotides (N). Similarly, the RII_DMJX5

(2009). The RT–qPCR experiments were performed with technical triplicates. Only RT–qPCR experiments with efficiencies greater than 1.9 were retained.
copy, with 1,629 bp aligned against RC, may also be complete because it has a long N stretch at its 5’-end. The other copies were small and fragmented by insertions and deletions. The identification of the relative chromosome position of these copies, as established by analyzing the polytene chromosome maps found in Flybase (Schaeffer et al. 2008), revealed that three of the four putative full-length copies, RII_DMJA1, RII_DMJA2, and RII_DMJA3, were inserted in the central portion of different chromosome arms corresponding to chromosomes 2, 3, and 5, respectively.

Moreover, we looked for the I element promoter in the D. mojavensis copies by aligning the flanking sequences of each copy with the promoter sequence of the I element of D. melanogaster (Minchiotti et al. 1997; Han 2010). We identified a putative promoter in the RC that has the same length (41 bp) and is well conserved (76% ID) compared with that of the D. melanogaster I element. This promoter region is identical in the four putative full-length copies found in the D. mojavensis genome (fig. 3). The four longest I sequences have a poly-A tail in the 3’-UTR, with an average of four TAAA repeats.
Phylogenetic Relationship between the D. mojavensis I-Like Element and the I Elements of Other Drosophila Species

The phylogenetic relationship of the different I-like elements was reconstructed using the RT protein sequences (see Materials and Methods for details) (fig. 4). It allowed us to distinguish four major clades of sequences (A–D). Different I-like sequences are present in species of the melanogaster group of the subgenus Sophophora (see supplementary fig. S3, Supplementary Material online, for the species phylogeny), which are included in all four clades (fig. 4). Clade A can be divided into two groups of sequences (Clade A1, bootstrap: 70% and Clade A2, bootstrap: 99%). Inside Clade A1, which branches in two smaller clades (A1a and A1b), all the I-like sequences of the D. mojavensis and of D. arizanae cluster in a highly supported clade (bootstrap: 100%), which clusters with the D. virilis sequence in a well-supported monophyletic group (A1b: 99%). This clade is closely related to the group of species belonging only to the Sophophora subgenus: Clade B includes only species from the melanogaster group, whereas Clades C and D also include sequences from D. willistoni (the willistoni species group). The phylogeny reveals that different I-like families are harbored by the same species, as is the case for D. kikkawai, whose sequences branch into three different clades (A1, A2, and C).

Activity of the Complete I Copies

We observed I element mRNA in ovarian cells by immunofluorescence and detected the presence of transcripts in both parents and hybrids. The I transcripts had the same localization in both species, with an accumulation in the nurse cells and no labeling of the oocytes. We did not detect any signal in the follicle cells (fig. 5). In testes, we observed labeling in parental lines and both types of hybrids (fig. 6). Transcripts were specifically located in primary and meiotic spermatocyte cells (Fuller 1998). Since in situ hybridization is not a quantitative technique, to investigate whether the levels of expression were equal between the different lines (parents and hybrids), we performed expression analyses by RT–qPCR. The results showed transcription of the I elements in D. mojavensis and in D. arizanae, both in ovaries and testes (fig. 7). The levels of transcription between the parental lines and the reciprocal male and female hybrids were not significantly different between ovaries from parents and hybrids (two-way analysis of variance [ANOVA]). However, expression levels were significantly higher in hybrid testes produced by crosses between D. arizanae females and D. mojavensis males than in the reciprocal cross ($F(3, 9) = 84.78, P < 0.0001$). We obtained the same results with crosses and expression analyses performed again 6 months later (supplementary fig. S4, Supplementary Material online).

To detect transposition, we performed TD analyses. This experiment needs to be performed with individual crosses, in order to follow new transpositions that arrived in the germ line cells, and that will be present in all the offspring of the two parents. Unfortunately, these experiments could not be performed for both reciprocal crosses because individual crosses between single D. arizanae females with single D. mojavensis males did not give any offspring. This cross is only possible with a mass of individuals. TD was thus only performed in one cross direction (supplementary table S1 and fig. S1, Supplementary Material online). TD also allows determining the copy number. In the individuals that we analyzed, D. mojavensis females had one copy of I-like element and D. arizanae males had from three to five copies of the element. In agreement with the RT–qPCR results, no transposition was detected in the F1 tissues, which indicates that the I element does not transpose in the parental gonads. Additionally, no transposition was detected in the F2 generation, indicating that neither the I sequences in F1 females nor in D. arizanae males induce transposition (data not shown). However, given the limitations of TD technique, we cannot exclude that somatic transposition occurred, but we were unable to measure this with confidence.

Discussion

Identification and Phylogenetic Analysis of I-Like Elements

In this report, we identified and characterized I-like sequences in the sequenced genome of D. mojavensis. In addition to several defective and divergent sequences, we identified four putatively complete copies in this species, but only one showed all typical structural domains of the I superfamily (Malik et al. 1999). This finding suggests that this copy may also have transposition ability. In D. melanogaster, two intact ORFs are a requisite for protein encoding and complete retrotransposition of the /factor (Selleme et al. 2005). The presence of unidentified nucleotides in some of the detected copies in D. mojavensis suggests the possibility that other sequences are potentially full-length copies. One argument favoring this hypothesis is the conservation of the promoter region that we identified, which is typically not conserved between species (Minchiotti et al. 1997). This promoter region is more conserved between the copies from D. mojavensis and D. melanogaster (76%), than the RT sequences (66%), which is puzzling. Further work should be done to understand if this could be related to the testis-specific expression in the hybrid male offspring.
Non-LTR retrotransposons are common elements in Drosophila genomes and are often described as vertically inherited in this group of species (Malik et al. 1999; Granzotto et al. 2011). I-like element sequences have been found in many species, and their distribution correlates with the phylogenetic relationships between species (Bucheton et al. 1992), which indicates that they are old components of Drosophila genomes. However, I elements with a structure strikingly similar to that of D. melanogaster (I_DM) occur only in the species of the melanogaster subgroup (D. simulans, D. mauritiana, and D. sechellia). The most widely supported hypothesis is that ancestral I elements were lost from the genome of D. melanogaster but were then reintroduced recently because older strains of D. melanogaster contain only defective and immobile I sequences, and all the strains collected after 1930 have both types of sequences (defective and active mobile I elements). This observation is in agreement with the hypothesis that the complete and active I factor progressively invaded

![Immunofluorescence for the I element mRNA probe (RNA in red and DNA in green) in the egg chamber of the parental and hybrid crosses. Oo, oocyte; NC, nurse cells; Fc, follicle cells. The arrows indicate mRNA of the detected I transcripts.](image)

**Fig. 5.** — Immunofluorescence for the I element mRNA probe (RNA in red and DNA in green) in the egg chamber of the parental and hybrid crosses. Oo, oocyte; NC, nurse cells; Fc, follicle cells. The arrows indicate mRNA of the detected I transcripts.
D. melanogaster after this period (Kidwell 1983). The process by which \( I \) elements have reinvaded the D. melanogaster genome is unknown. Horizontal transfers from another species (Simonelig et al. 1988) is one of the possibilities, but more complex hypotheses, such as the reactivation of sequences sequestered in the heterochromatin, should also be considered.

The phylogeny placement of the \( I \)-like elements from D. mojavensis is generally in agreement with the species phylogeny. All the \( I \)-like sequences of D. mojavensis and D. arizonae branch together with the sequence of D. virilis, as expected based on the species phylogeny because these species belong to the virilis–repleta radiation of the subgenus Drosophila suggesting a vertical inheritance of these sequences in this Drosophila radiation. However, when we analyze the global phylogeny of the \( I \)-like element, we cannot exclude that some horizontal transfer events may have occurred between other Drosophila species. For example, this could have been the case between D. ficusphila and D. bipectinata (see Clade A1a in fig. 4).

Activity of the \( I \) Element

We measured the transcriptional activity of the \( I \) elements by RT–qPCR and fluorescent in situ hybridization (FISH) experiments in the parental lines and their hybrids. The level of \( I \) element mRNA was low in ovaries and testes of the parental lines. In the parental ovaries, the \( I \) element transcripts accumulate in the nurse cells. This type of labeling is also observed for the \( I \) element in Drosophila ovaries (Seleme et al. 2005; Chambeyron et al. 2008). However, transcription of the \( I \) element was never reported in testes in this species. Our RT–qPCR analysis shows that the \( I \) element is transcribed in testes of both D. mojavensis and D. arizonae; this was confirmed by FISH, which showed labeling in primary and meiotic spermatocytes. In the reciprocal hybrids between the two species, the levels of expression were identical for the female germline tissues, with no difference between parents and F1 hybrids. FISH experiments also confirmed this result because no change in the localization of the transcripts and no labeling occurred in the Sophophora subgenus, which was accompanied by sequence loss and horizontal transfer events. In the Drosophila subgenus, we can only assume that random loss of the \( I \) families lead to only one remaining family, which is described in this article and which is closely related to the \( I \) factor from D. melanogaster.

**Fig. 6.**—Immunofluorescence for the \( I \) element mRNA probe (RNA in red and DNA in green) in testes of the parental and hybrid lines. Yellow arrows indicate detected mRNA transcripts. Sc.Mt.c, sperm and mitotic cells; first spty, primary spermatocytes; Me.sp., meiotic spermatocytes; R.sp, round spermatids; Di.sp, differentiating spermatids; El.sp, elongating spermatids; horizontal arrow indicates the direction of differentiation (stem cells are located at the apical tip and germ cells move toward the basal end as they differentiate).
in the oocytes were observed. For the male germline, the results were quite different depending on the direction of the cross. F1 males from *D. arizonae* male parents showed no difference in the expression of the *I* element in testes. However, when *D. mojavensis* males were used as parents, a significant increase in the *I* element expression in testes was observed. The fact that F1 males in this particular cross are sterile (Ruiz et al. 1990) suggests a role for TEs in the male-sterile phenotype. Furthermore, it is important to determine whether the high expression in F1 hybrids observed in RT-qPCR is derived from complete copies of *D. mojavensis* males. Hybrid dysgenesis in *D. melanogaster* involves transposon-naive females and active transposon-containing males in several different systems: The *I*-R system, the *P*-M system, and even the *H*-E system (Bucheton et al. 1976; Picard et al. 1978; Kidwell and Novy 1979; Kidwell 1983; Streck et al. 1986). However, at least in the *I*-R and *P*-M systems, it is the maternal line that is affected.

The presence of *I* transcripts only in the nurse cells in ovaries of *D. mojavensis*, *D. arizonae*, and their hybrids suggests that the *I* element in both species is most likely regulated by the piRNA pathway. In *Drosophila* ovaries, Chambeyron et al. (2008) showed that there is an association between the accumulation of piRNA and the regulation of the *I* element transcripts in the nurse cells, where the transcripts are processed, before being transported to the oocytes, where the retrotransposition occurs. Such findings suggest a posttranscriptional regulation of the *I* element described here before its transportation to the oocytes, as reported in *D. melanogaster* (Brennecke et al. 2008; Chambeyron et al. 2008).

In testes, the *I* transcripts in parents and hybrids are located in primary spermatocytes, where high levels of transcription for genes encoding proteins for spermatogenesis and genes encoding male germline-specific isoforms are detected (Fuller 1998). The transcripts of many genes are stable and are present for days after meiosis but then decrease in the postmeiotic spermatids stage, during which proteins can be detected (Kuhn et al. 1988; Fuller 1998). The localization of the *I*-like transcripts observed in this study (fig. 7) corresponds to the transcriptional dynamics described for their genes. Other elements have transcripts detected in primary spermatocytes. Transcripts of 412 retrotransponson were not detectable in ovaries by in situ hybridization in *D. melanogaster* and *D. simulans* populations but were detected in the male germ-line in primary spermatocytes (Borie et al. 2002). Furthermore, no association has been found between expression in soma and in testes, which indicates independent regulation (Borie et al. 2002). The same localization patterns were observed for the retrotransposons 1731, GATE, *mdg1*, and *copia* (Haoudi et al. 1997; Kogan et al. 2003; Morozova et al. 2004; Kalmykova et al. 2005). The relatively high number of transcribed elements, including the *D. mojavensis* *I* element, in primary spermatocytes reveals the importance of these cells in the transcription of these TEs.

In this work, we have investigated the hypothesis that *I*-like elements (which are implicated in the hybrid dysgenic
incompatibility of *D. melanogaster* could be activated in hybrids between *D. mojavensis* and *D. arizonae*, two recently diverged species from the *Drosophila* subgenus of the genus *Drosophila*. We showed that I elements are specifically activated in testes from a specific direction of the intercross, and we suggest that this activation may be associated with the observed male sterility. Moreover, we showed that this activation is sex specific because the female germline does not seem to be affected by this phenomenon. The impact of TEs in the male sterility observed in interspecific crosses is becoming clearer and needs further investigations. Studies using closely related species will be helpful to understand the first steps of the male sterility process and the role of TEs.

**Supplementary Material**

Supplementary figures S1–S4 and table S1 are available at *Genome Biology and Evolution* online (http://www.gbe.oxfordjournals.org/).

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