Transgenic tools for targeted chromosome rearrangements allow construction of balancer chromosomes in non-melanogaster Drosophila species

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Abstract

Perhaps the most valuable single set of resources for genetic studies of Drosophila melanogaster is the collection of multiply inverted chromosomes commonly known as balancer chromosomes. Balancers prevent the recovery of recombination exchange products within genomic regions included in inversions and allow perpetual maintenance of deleterious alleles in living stocks and the execution of complex genetic crosses. Balancer chromosomes have been generated traditionally by exposing animals to ionizing radiation and screening for altered chromosome structure or for unusual marker segregation patterns. These approaches are tedious and unpredictable, and have failed to produce the desired products in some species. Here, I describe transgenic tools that allow targeted chromosome rearrangements in Drosophila species. The key new resources are engineered reporter genes containing introns with yeast recombination sites and enhancers that drive fluorescent reporter genes in multiple body regions. These tools were used to generate a doubly inverted chromosome 3R in Drosophila simulans that serves as an effective balancer chromosome.

Keywords: transgenes; chromosome rearrangements; chromosome inversion; nonmodel species; Drosophila; reporter genes; fluorescent reporters

Introduction

Balancer chromosomes are inverted chromosome segments that prevent inheritance of the products of recombination events and suppress recombination near inversion breakpoints (Crown et al. 2018) and thus allow maintenance of deleterious mutations in stable stocks (Hentges and Justice 2004; Kaufman 2017). A single chromosomal inversion can serve as a balancer chromosome, but only for alleles located close to inversion breakpoints. Balancer chromosomes have therefore traditionally been constructed through serial introduction of new inversions in the context of previous inversions (Kaufman 2017). Multiply inverted chromosomes have been the basis of all widely used balancer chromosomes in Drosophila melanogaster, giving rise to the prefixes of many widely used balancers (FM, first multiple; SM, second multiple; TM, third multiple) and balancers have facilitated many D. melanogaster experiments.

In plants, C. elegans, and mouse and human cells, inversions have been generated using CRISPR-Cas9 mutagenesis to induce simultaneously 2 double-strand breaks, which occasionally resolve as an inversion (Li et al. 2015; Iwata et al. 2016, 2019; Zhang et al. 2017; Korabliev et al. 2017; Djejima et al. 2018; Schmidt et al. 2019). Similar efficient approaches have not been reported in Drosophila, and I have failed to recover inversions in D. melanogaster using this kind of approach to generate inversions on the order of 10 kb–1 Mbp. Recently, Ng and Reed reported successful isolation of a single inversion event in D. melanogaster using transgenic sources of Cas9 and gRNAs (Ng and Reed 2019). The efficiency of this approach is not yet known. The success of this method depended on a clever positive selection scheme that allowed survival of only animals that carried an inversion. It is not yet clear whether this method can be modified to incorporate markers or selection to allow identification of inversion events in a more general setting.

Inversions have traditionally been generated in Drosophila species by exposing flies to ionizing radiation that generates chromosome breaks, which are then incorrectly repaired. Balancer chromosomes would be valuable for diverse Drosophila species, but perhaps most useful for Drosophila simulans, a sister species to D. melanogaster. Comparisons between D. melanogaster and D. simulans have provided fundamental insights into genetic evolution ever since D. simulans was discovered by Alfred Sturtevant in 1919 (Barbash 2010). However, while irradiation is effective to generate chromosome inversions in multiple species, for reasons that are not clear, it has proven difficult to induce inversions by mutagenesis in D. simulans. Several studies have found that irradiation and chemical mutagenesis induce rearrangements at a lower rate in D. simulans than in D. melanogaster (Lemke et al. 1978; Inoue 1988), although Woodruff and Ashburner (1978) observed irradiation-induced chromosome breaks at equal frequencies in each species, indicating that D. simulans retains mechanisms...
required for chromosome repair. In addition, inversion polymorphisms are nearly absent in D. simulans populations but common in D. melanogaster populations (Aulard et al. 2004). I am aware of several previous unpublished efforts to generate inversions in D. simulans using irradiation, but apparently none yielded useful balancer chromosomes.

One inversion stock is available in D. simulans with chromosome breaks at cytological locations 81F1 (approximately 0.91 Mbp on 3R) and 89E (9.75 Mbp), in the proximal region of chromosome 3R (Coyne and Sniegowski 1994). The distal break apparently disrupts Ultrabithorax (Ubx), since 1 breakpoint is at 89E, the cytological location of Ubx, and the inversion carries an Ubx allele. This inversion was probably generated by E. H. Grell (Coyne and Sniegowski 1994), perhaps using irradiation, though no details are available. This inversion can be stably maintained on modified chromosomes, for non-plasmonic construction of balancer chromosomes, and other could be adapted for use in multiple D. simulans on 3R over 15 generations and thus will serve as an effective balancer chromosome.

This doubly inverted chromosome arm prevented recombination and red) could be used to visualize fluorescent molecules easily in fly eyes on a binocular microscope. I therefore first searched in body regions of the adult fly. The use of these reagents is illustrated by the construction of a D. simulans doubly inverted 3R. These transgenic reagents could be adapted for use in multiple Drosophila species, thus simplifying the construction of balancer chromosomes, and other modified chromosomes, for non-melanogaster Drosophila species.

**Methods**

**Design**

An autosomal balancer chromosome should contain multiple overlapping inversions, a dominant marker, and a recessive lethal allele (Kaufman 2017). It is straightforward to introduce recessive lethal alleles onto existing chromosomes with chemical mutagenesis or CRISPR mutagenesis, so I designed reagents that would accomplish the first 2 goals, a chromosomal region containing overlapping inversions and a dominant marker. I exploited a collection of existing mapped pBac transgenic D. simulans strains that each carry a dominant fluorescent marker (3XP3-EYFP) adjacent to a site-specific attP recombination site (Stern et al. 2017). I integrated new reagents into these attP sites that would be modified during inversion events but leave the 3XP3-EYFP markers unaltered, ultimately generating inverted chromosomes carrying 3XP3-EYFP (Fig. 1). I thus required enough markers to follow all of the transgenes in a single experiment. Generating a double inversion requires, at a minimum, 6 markers: 1 dominant marker that remains in the balancer chromosome; 4 markers to identify the sites targeted for recombination; and 1 marker for the recombinase. I performed these initial experiments in D. simulans flies carrying w- and used recombinase transgenes carrying mini-white. Since the YFP/GFP channel is occupied by the attP landing sites, only 2 other channels (blue and red) could be used to visualize fluorescent molecules easily in fly eyes on a binocular microscope. I therefore first searched for additional enhancers that would drive high levels of fluorescence in other body regions.

**New reporter constructs**

Reporter constructs were generated to test the ability of synthetic combinations of characterized transcription factor binding sites for Tinman (Kimbrill et al. 2002) and Mef2 (Andres et al. 1995) to drive a fluorescent reporter in specific anatomical domains. gBlocks (IDT) were synthesized carrying 3 or 9 binding sites of each transcription factor binding site and homology arms complementary to the plasmid sequences, allowing Gibson cloning (Gibson et al. 2009) of these engineered enhancers upstream of eGFP reporter genes in piggyBac plasmids carrying an attP site-specific recombination sequence, allowing integration either through piggyBac transposition or attP-attB site-specific integration (Supplementary Table 1). A second set of Tinman and Mef2 transcription factor binding site constructs were generated with GAGA binding sites (Wilkins and Lis 1998, Leibovitch et al. 2002) to attempt to boost expression levels (Supplementary Table 1).

Reporter constructs were generated to test 145 and 285 bp enhancers of the Actin5C gene and 453 and 678 bp enhancers of the Myosin Heavy Chain gene (Supplementary Table 1). The i1 enhancer (Masumoto et al. 2012) was PCR amplified from a plasmid provided by Rory Coleman and Gibson cloned upstream of fluorescent reporter genes.

The full list of plasmids generated for this project are listed in Supplementary Table 2. All plasmids were re-sequenced in their entirety and annotated full-plasmid sequences are provided as Supplementary Material. All plasmids are available from Addgene.org.

**Generation of transgenic strains**

Plasmids were introduced into the D. melanogaster attP2 site-specific recombination site or various D. simulans attP sites by standard injection protocols performed by Rainbow Transgenics. For D. simulans injections, we compared coinjection of attP plasmids with a “helper” plasmid carrying phiC31 integrase under the control of a heat-shock inducible promoter vs mRNA encoding phiC31 integrase. PhiC31 integrase mRNA was generated as follows. p(T7-phiC31-nos-3’UTR) plasmid DNA was linearized by digestion with EcoR1 and electrophoresed in a 1% agarose gel in 40 mM Tris, 20 mM Acetic Acid, 1 mM EDTA buffer. The gel extracted linearized DNA was cleaned with the Zymoclean Gel DNA recovery kit (Zymo) and used as template in mMESSAGE mMACHINE T7 Transcription Kit (Thermo Fisher). RNA was cleaned with a LiCl2 precipitation followed by cleanup with a MEGAclean Transcription Clean-Up kit (ThermoFisher). Embryos were injected with 200ng/μl of plasmid DNA and 500ng/μl of mRNA.

**Reporter gene expression patterns**

Photomicrographs of expression of reporter constructs in living adults were generated with an Olympus MBX10 macro zoom microscope with microscope magnification and exposure conditions held constant across all photographs. A subset of the reporter lines were selected to examine potential reporter gene expression in the central nervous system. Central nervous systems were dissected, antibody stained, and imaged by Project Technical Resources at Janelia Research Campus as described previously (Aso et al. 2014). In brief, brains and VNCs were dissected in Schneider’s insect medium and fixed in 2% paraformaldehyde (diluted in the same medium) at room temperature for 55 min. Tissues were washed in PBT (0.5% Triton
X-100 in phosphate buffered saline) and blocked using 5% normal goat serum before incubation with antibodies. DsRed was detected by staining with rabbit anti-dsRed (Clontech 632496, 1:1,000) followed by CyTM2-conjugated goat antirabbit (Jackson ImmunoResearch 115-225-166, 1:1,000). After staining and post-fixation in 4% paraformaldehyde, tissues were mounted on poly-L-lysine-coated cover slips, cleared, and embedded in DPX as described previously. Image Z-stacks were collected at 1 um intervals using an LSM710 confocal microscope (Zeiss, Germany) fitted with a Plan-Apochromat 20x/0.8 M27 objective.

**Generation of doubly inverted D. simulans chromosome 3R**

Four plasmids carrying different reporter genes and yeast recombination sites were integrated separately into existing attP landing sites in D. simulans (Stern et al. 2017): p{3XP3-dsRed::FRT, attB} into 367-simD-311.3 at 3R : 1,235,923 bp; p{Mhc-F4-678-EGFP::KDRT, attB} into 1392-simD-925.4 at 3R : 10,884,926; p{3XP3-mTurquoise::FRT-RC, attB} into 1029-simD-234.5 at 3R : 17,243,204, and p{Mhc-F4-678-DsRed::KDRT, attB} into 1048-simD-299.6 at 3R : 10,843,913. All genome coordinates are relative to D. simulans genome assembly NCBI. GCA_016746395.1. All 4 transgenes were recombined onto a single chromosome arm. Flies carrying all 4 transgenes on a single chromosome were crossed to flies carrying a source of heat-shock inducible Flippase p{MUH-HES-FLP1::PEST, w+, attB} integrated into 1173-simD-523.7 at X : 13,484,041. Plastic vials containing developing larvae in their food media were placed in a 37°C bead bath (Lab Armor) for 1 h, followed by 1 h at room temperature, and a second 37°C heat shock for 1 h. Male progeny were crossed to a w strain of D. simulans and offspring of this cross were screened for loss of 3XP3-DsRed and 3XP3-mTurquoise, but retention of MHC-EGFP and MHC-DsRed, which indicates inversion between the FRT sites. Flies carrying this singly inverted chromosome were then crossed to flies carrying a source of heat-shock inducible KD recombinase p{MUH-HES-KD::PEST, w+, attB} integrated into 2021-simD-299.6 at 2L : 6,583,842. This attP landing site is derived from 1048-simD-299.6, but has had its 3XP3-EYFP marker inactivated by CRISPR-Cas9 mutagenesis of the EYFP gene (Stern et al. 2017). Larvae from this cross were heat shocked as described above, and adults were crossed to a w strain of D. simulans. Offspring were screened for loss of MHC-EGFP and MHC-DsRed, but presence of 3XP3-EYFP, which is present in all 4 landing sites with integrated recombinase plasmids, but not in the landing...
site with the KD-recombinase integrated. These progeny carried presumptive double-inversions on chromosome 3R.

**Testing for recombination with balancer chromosome**

To determine whether the doubly inverted chromosome arm might serve as a useful balancer chromosome, flies carrying this chromosome, which could be recognized by the 3XP3-EYFP markers, were backcrossed to a u^w^ strain of *D. mauritiana* for 15 generations. Ten independent backcrosses were performed and DNA was prepared from 1 individual of the 15th generation from each backcross and subjected to multiplexed-shotgun genotyping to estimate chromosome ancestry (Andolfatto et al. 2011).

**Results**

**An expanded palette of enhancers useful for fluorescent reporters in Drosophila**

Tracking multiple transgenes, especially in species with few existing genetic reagents, requires development of new markers. For multiple reasons, I favored use of fluorescent markers. Fluorescent markers can be used in many species (Stern et al. 2017), they can be used in flies with normal eye color (Horn and Wimmer 2000), and they are unlikely to disrupt behavior, whereas eye color and pigmentation markers commonly used in *D. melanogaster* disrupt behavior (Kyriacou et al. 1978; Zhang and Odenwald 1995; Suh and Jackson 2007; Anaka et al. 2008; Massey et al. 2019).

I first attempted to engineer synthetic enhancers similar to the 3XP3 enhancer, which consists of 3 binding sites for the *Drosophila* Eyeless transcription factor, called P3 sites (Sheng et al. 1997). The 3XP3 enhancer drives strong expression in the eye and has become a popular enhancer to mark transgenes in many nonmodel species (Horn et al. 2000, 2002). As expected, the 3XP3 enhancer drove strong expression in neurons of the optic lobe and in many other regions of the brain and ventral nervous system (Supplementary Fig. 1, a–d), which may limit its utility for some purposes.

I first generated concatemers of transcription factor binding sites for the Tinman and Mef2 transcription factors, which I hypothesized might drive expression in the heart and the muscles, respectively (Andres et al. 1995; Kimbrell et al. 2002). The Tinman binding site constructs did not drive strong expression of fluorescent reporters that could be detected in adults (Fig. 2, a and b), even after addition of GAGA binding sites (Leibovitch et al. 2002) which were hypothesized to open chromatin to allow improved access from Tinman transcription factors (Fig. 2c). The Mef2 binding site 3X and 9X concatemers drove weak expression in a subset of flight and leg muscles (Fig. 2, d and e). The 3XMef2 constructs also drove extensive expression in the nervous system (Supplementary Fig. 1, e–h). These synthetic enhancers did not drive adult expression that was judged useful for most experiments.

One hundred and forty-five and 285 bp fragments from the *D. melanogaster* Actin 88F gene were tested. No transgenic animals were recovered with the 145 bp enhancer, suggesting that this enhancer did not support strong expression. The 285 bp enhancer drove strong expression in thoracic muscles (Fig. 2f), but also drove some weak expression in eyes (which is not easily seen in Fig. 2f). This enhancer drove no detectable expression in neurons of the brain or ventral nervous system in either sex (Supplementary Fig. 1, i–l).

**Confirmation of balancer efficiency**

To determine the efficiency of j3RM1 to balance the third chromosome, *D. simulans* flies carrying j3RM1 were crossed to *D. mauritiana*, and virgin female offspring were backcrossed to *D. mauritiana* for 15 generations (Fig. 3b). Crosses were made to *D. mauritiana* because *D. simulans* and *D. mauritiana* chromosomes are homosequential and display recombination throughout the genome (True et al. 1997). In addition, the species genomes are sufficiently divergent that even small genomic regions from each species can be identified unambiguously (Meiklejohn et al. 2018). Flies from 10 independent backcrosses were subjected to light shotgun genome resequencing and ancestry of genomic regions...
was estimated using Multiplexed Shotgun Genotyping (Andolfatto et al. 2011). Flies from all 10 crosses displayed heterozygosity for D. simulans and D. mauritiana DNA across the entire right arm of chromosome 3 (Fig. 3b), indicating that j3RM1 prevented recovery of recombination events on 3R for 15 generations.

**Discussion**

The doubly inverted D. simulans chromosome 3R, called j3RM1, prevented recombination over 15 generations in 10 independent crosses (Fig. 3b). This suggests that a single pair of overlapping inversions is sufficient to block inheritance of recombinant
chromosomes to an extent that will make this balancer chromosome useful for many kinds of experiments. In our experience, j3RM1 appears to have reduced viability relative to the un-inverted chromosome, but the chromosome is not homozygous lethal. However, it would be straightforward to add lethal alleles either through classical mutagenesis or CRISPR-Cas9 editing, and these manipulations are underway in our laboratory.

While the FRT and KDRT sites utilized here supported recombination between sites located approximately 16 and 20Mbp apart, inversion events were rare. Although we did not quantify the inversion frequency precisely, we recovered approximately 1 inversion per several thousand flies, a frequency not too different than observed previously for FRT sites located ≥10Mbp apart (Golic and Golic 1996). There may be several ways to increase the efficiency with which inversions are generated. For example, the heat-shock source of recombinases could be replaced with a stable germ-line source of recombinase. This might increase efficiency both by increasing the concentration of recombinase in cells and by exposing all germ cells to high levels of recombinase.

An alternative approach to increase efficiency might be to exploit homologous recombination to drive inversions. Egli et al. (2004) demonstrated that homologous recombination generated inversions in D. melanogaster at a frequency of 1%–4%, independent of the distance between targeted sites. This is at least an order of magnitude more frequent than recombinase catalyzed inversions recovered in this study. Egli et al. targeted 2 double-strand breaks using a rare-cutting restriction endonuclease that recognized a restriction site engineered into an intron of a yellow transgene. This general approach could be expanded by exploiting multiple rare-cutting restriction endonucleases (Taylor et al. 2012) or Cas9 gRNA sites. The target sites and long homology regions could be incorporated into the Gateway cloning sites of the plasmids described here.

Alternative uses of these reagents

The fluorescent marker reagents described here simplified the generation of flies carrying many different transgenes, which we required to drive multiple inversions. One can envision many other potential uses of these markers. For example, many neurogenetic experiments performed in D. melanogaster require assembly of genotypes carrying many transgenes, most or all which are normally marked with mini-white. These complex crossing schemes are enabled, of course, by balancer chromosomes. However, the availability of up to 9 dominant fluorescent reporters, using 3 colors to mark 3 body regions, may enable the construction of complex transgene genotypes in other species without requiring balancer chromosomes. The fact that the act88F, MHC, and ie1 reporters drive limited expression in the nervous system (Supplementary Fig. 1) may make these more useful markers for neurogenetic studies than 3XP3 reporters.

Most transgenic reagents used in D. melanogaster have employed visible markers that rescue eye color or cuticle pigmentation mutants. However, most or all of these mutations can influence development and behavior. For example, mini-white transgenes can influence male behavior (Zhang and Odenwald 1995; Anaka et al. 2008) and pigmentation mutations can alter behavior (Suh and Jackson 2007; Massey et al. 2019). Fluorescent markers may have fewer or weaker influences on development and behavior than transgenes carrying native Drosophila genes.

The reagents described here can be used to drive inversions, as illustrated, but could also be adapted to generate targeted deletions, as has proven useful in D. melanogaster (Ryder et al. 2004). The reagents can be adopted readily for other species.

Fig. 3. Testing for suppression of recombination by balancer j3RM1. a) D. simulans flies carrying j3RM1 were crossed to a u- strain of D. mauritiana. Females of subsequent generations were backcrossed for 15 generations. b) Chromosomal ancestry estimates for single individuals from 10 independent 15-generation backcross experiments. DNA from each individual was subjected to light shotgun sequencing and ancestry was estimated using the Hidden Markov Model of MSG (Andolfatto et al. 2011). The y axis of each plot represents the probability that a chromosomal region is homozygous for either D. simulans (up, red regions) or D. mauritiana (down, blue regions). Individual DNA markers are shown as red and blue carpets along the top and bottom axes. Regions without support for homozygosity for either species are estimated to be heterozygous for the 2 species. Above the top plot, the approximate locations of the 4 markers (colored triangles) and 2 inverted segments (gray bars, labeled as In-1 and In-2) are indicated. In all individuals, the entire right arm of 3R is heterozygous, indicating the absence of recombination events within this region represented in progeny. In all individuals, the region of heterozygosity extends to the leftmost breakpoint. The chromosomal arms and scale bar in Mbp are indicated at bottom. For the scale bar, 10 individual segments of 1 Mbp each are shown.
carrying attF landing sites (Stern et al. 2017). Alternatively, the reagents could be introduced directly into targeted genomic locations using CRISPR-Cas9 homology directed repair to allow inversions and deletions of specific chromosomal regions. This might provide a means to simplify reversal of naturally occurring inversions, making these regions accessible to further genetic studies (Schmidt et al. 2020).

**Data availability**

All new data are available at Figshare. Genome re-sequencing data from backcross experiments are at https://figshare.com/articles/dataset/In3RG1_1_1_fastq.gz/19064738. Full plasmids sequences are available at https://figshare.com/articles/dataset/Full_plasmid_sequences/19070825. All plasmids are available from addgene.org, plasmids # 182163–182199. All transgenic flies and the balancer chromosome j3RM1 are available from the author upon request.

Supplemental material is available at G3 online.

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**Conflicts of interest**

None declared.

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