The zinc finger protein CLAMP promotes long-range chromatin interactions that mediate dosage compensation of the Drosophila male X-chromosome

Abstract

Background: Drosophila dosage compensation is an important model system for defining how active chromatin domains are formed. The male-specific lethal dosage compensation complex (MSLc) increases transcript levels of genes along the length of the single male X-chromosome to equalize with that expressed from the two female X-chromosomes. The strongest binding sites for MSLc cluster together in three-dimensional space largely independent of MSLc because clustering occurs in both sexes. CLAMP, a non-sex specific, ubiquitous zinc finger protein, binds synergistically with MSLc to enrich the occupancy of both factors on the male X-chromosome.

Results: Here, we demonstrate that CLAMP promotes the observed three-dimensional clustering of MSLc binding sites. Moreover, the X-enriched CLAMP protein more strongly promotes longer-range three-dimensional interactions on the X-chromosome than autosomes. Genome-wide, CLAMP promotes three-dimensional interactions between active chromatin regions together with other insulator proteins.

Conclusion: Overall, we define how long-range interactions which are modulated by a locally enriched ubiquitous transcription factor promote hyper-activation of the X-chromosome to mediate dosage compensation.

Keywords: Dosage compensation, Three-dimensional genome organization, Transcription, Chromatin, Drosophila, MSL complex

Highlights

- CLAMP promotes long-range interactions on the Drosophila male X-chromosome
- CLAMP differentially regulates 3D interactions in active versus inactive chromatin
- Enrichment of insulator proteins is observed at CLAMP-regulated 3D genomic interaction sites

Introduction

Three-dimensional chromatin domains are important for coordinating gene regulation. Recent work has provided new insight into how silent chromatin domains are formed, for example, through phase separation [1, 2], but less is understood regarding the formation of hyper-active chromatin domains [3]. Dosage compensation in Drosophila provides one of the few model systems for studying the formation of a large hyper-active chromatin domain: approximately one thousand active genes along the length of the single male X-chromosome are coordinately upregulated twofold [4–6].

In heterogametic species, dosage compensation is essential to correct transcriptional imbalance of X-linked
genes between the sexes [7] and to correct for dosage imbalance between the single X-chromosome and paired autosomes. Diverse dosage compensation mechanisms have evolved across species, but an essential conserved step is distinguishing the X-chromosome from autosomes for specific regulation.

In *Drosophila*, the male-specific lethal complex (MSLc) forms only in males and is responsible for increasing transcript levels of X-linked genes along the length of the single male X-chromosome 1.4 fold, helping to equalize gene expression with that of females [4–6]. MSLc consists of five proteins MSL1, MSL2, MSL3, maleless (MLE) [8–10], males absent on the first (MOF) [11], and one of two functionally redundant long non-coding RNAs known as RNA on the X 1 and 2 (roX1 and roX2) [12, 13]. MSLc first targets X-linked genomic elements known as “high-affinity” (HAS) or “chromatin entry” sites (CES), which also include the roX loci [12, 14–16].

Within CES, MSLc is recruited to GA-rich 21-bp elements known as MSL recognition elements (MREs) [14]. Accumulation of MRE sequences on the X-chromosome occurred by expansion of GA-rich sequences and transposon insertion [17, 18]. However, MRE sequences are not X-chromosome specific and are only approximately twofold enriched on the X-chromosome compared with autosomes [14], suggesting they are not sufficient for X-chromosome targeting. Although the MSL2 component of MSLc has a low affinity for MREs, MSL complex requires synergy with an essential, non-sex specific, zinc finger adapter protein known as chromatin-linked adapter for MSL proteins (CLAMP) in order to stabilize its binding to MREs [19–21].

Synergy between CLAMP and MSLc, which has been demonstrated both in vivo and in vitro [20, 21], enhances the occupancy of both factors on the male X-chromosome. Maternally deposited CLAMP is present on chromatin throughout the genome before MSLc assembles at the maternal–zygotic transition [22–24] and regulates chromatin accessibility of the X-chromosome and transcription of X-linked genes [19, 25]. Therefore, it is likely that CLAMP functions as an early transcription factor to enhance X-chromosome accessibility and promote MSLc targeting.

After initial targeting to CES by CLAMP, MSLc generates a hyper-active chromatin domain by localizing to the bodies of active genes and increasing their transcript levels through modulating transcription elongation [14, 15, 25–27]. MSLc has been shown to take advantage of pre-existing three-dimensional chromatin organization to target the X-chromosome [28, 29]. Chromosome conformation capture techniques have demonstrated that CES cluster three-dimensionally in both males and females and form long-range interactions with other X-linked active chromatin regions within the nucleus independent of MSLc [28, 29]. However, the mechanism by which CES cluster remained unknown.

We hypothesized that CLAMP promotes clustering of CES based on the following lines of evidence: (1) in contrast to MSLc, CLAMP is required to globally increase the accessibility of the entire male X-chromosome [25]; (2) CLAMP is part of two insulator protein complexes (Kaye et al., 2017; Bag et al., 2019), acts as an insulator protein in several functional assays [30], and promotes recruitment of the CP190 insulator protein. CP190 also promotes CLAMP recruitment and therefore there is a synergistic relationship between these two proteins [30]. Insulator proteins mediate chromatin interactions across the genome to regulate specialized chromatin domains throughout development [31–35]. However, it was not known whether CLAMP regulates the formation of three-dimensional interactions within the genome.

We used genome-wide chromosome conformation capture (Hi-C) analysis complemented by circular chromosome conformation capture with high-throughput sequencing (4C-seq) to test the hypothesis that CLAMP regulates clustering of CES and three-dimensional organization of the X-chromosome. We discovered that CLAMP promotes long-range interactions on the male X-chromosome more strongly than on autosomes. Furthermore, we demonstrate that CLAMP primarily promotes long-range interactions within active chromatin regions, including CES. We also show that enrichment of several insulator proteins is increased at loci where CLAMP regulates genomic interactions. Overall, we demonstrate that the X-enriched CLAMP protein regulates long-range three-dimensional interactions between CES to target MSLc to the male X-chromosome. Synergy between CLAMP and MSLc [20, 21] increases the occupancy of both factors to specifically hyper-activate approximately one thousand X-linked genes in males.

Results

CLAMP promotes long-range three-dimensional interactions on the X-chromosome more strongly than on autosomes

In order to understand how CLAMP regulates the three-dimensional organization of the genome, we performed in situ chromosome conformation capture with high-throughput sequencing (in situ Hi-C) [36] using HindIII, a 6-bp cutter restriction enzyme, in *Drosophila* male Schneider’s line 2 (S2) [37] cultured cells after validated RNAi depletion of either gfp (control) or clamp [4, 19, 20]. We performed two biological replicates for each experimental condition. We confirmed depletion of CLAMP protein after clamp RNAi by Western blot (Fig. 1A) and used GenomeDISCO [38] to determine replicate concordance. Reproducibility between replicates was high (Fig. 1B).
We visualized our Hi-C interaction maps by combining replicates for each condition (Fig. 1C; Additional file 3: Table S1). To compare the clamp RNAi and control gfp RNAi contact maps, we also generated a differential interaction map (Fig. 1C). In the differential map for the two conditions (clamp/gfp RNAi), we observed an increase in interaction frequency (red) directly along the diagonal (i.e., shorter-range interactions). Many of the more pronounced off-diagonal differences are in regions proximal to centromeres, which will be discussed later. In general, we observe decreased interaction frequency (blue) moving away from the diagonal (i.e., longer-range interactions) across all chromosomes when CLAMP is depleted and this decrease is more widespread on the X-chromosome compared with autosomes.

To quantify differences between our gfp and clamp RNAi interaction matrices, we calculated the log2 ratio of distal to local interactions (DLR) [39] (Fig. 2, Additional file 2: Fig. S1A). We defined local interactions as those that span less than 250 kb and distal as interactions that span more than 250 kb based on the average size of a TAD in Drosophila. After depletion of CLAMP, there is a change in the ratio between long-range and short-range interactions that is different on the X-chromosome compared with autosomes. The X-enriched decrease in the DLR after clamp RNAi is also observed when DLR is measured using a local vs distal cutoff of 100 kb instead of 250 kb (Additional file 2: Fig. S1B). Moreover, this X-enriched change in three-dimensional interactions (Fig. 2, Additional file 2: Fig. S1A,B) is consistent with previous MNase accessibility analysis demonstrating that global chromatin accessibility of the male X-chromosome but not autosomes decreases after clamp RNAi [25]. Therefore, our Hi-C data support a model in which CLAMP alters the three-dimensional organization of the male X-chromosome more than autosomes.

Previous Hi-C studies in multiple cell lines and embryos found that the strongest MSLc binding sites (CES) interact with each other and other genomic regions along the X-chromosome more frequently than expected by chance [28, 29]. To confirm these observations within our own Hi-C data, we investigated whether CES interact frequently with other genomic locations. We used Fit-Hi-C [40] to determine high-confidence intra-chromosomal contacts (see "Methods" section) from our control gfp RNAi Hi-C maps (Fig. 1D; Additional file 5: Table S3). Consistent with previous reports [28, 29], approximately 60% (3,090) of the high-confidence X-chromosome interactions identified at 20-kb resolution involve CES; 45% would be expected by chance, even when restricting our analysis to only active chromatin regions as controls ($p=5.7e−107$, $\chi^2$ test) (Fig. 1E). Therefore, our data are consistent with prior reports that CES interact more frequently with other regions of the genome than expected by chance [28, 29]. Moreover, we demonstrate that CES interact more frequently with other regions of the genome even when compared with other active chromatin regions that are known to cluster together.

To more quantitatively define specific regions throughout the genome where CLAMP regulates three-dimensional interactions, we compared intra-chromosomal interaction frequencies after clamp RNAi with those after gfp RNAi using diffHic [41]. DiffHic uses edgeR to model biological variability between replicates and perform differential analysis between conditions [41, 42]. We found 2552 significantly (FDR < 0.05) differential interactions (DIs) at 30-kb resolution (Additional file 2: Fig. S2A, Additional file 6: Table S4). We classified these DIs by the directionality of their log2 ratio. Notches on all box plots represent 95% confidence intervals around the median line; whiskers represent 1.5 IQR (inter-quartile range) and outliers have been omitted. Interactions that decrease in contact probability after clamp RNAi are defined as CLAMP-promoted (55%), whereas interactions that increase in contact probability after clamp RNAi are defined as CLAMP-repressed (45%). The log2 ratio was significantly larger for CLAMP-promoted X-linked interactions (Fig. 3A; Additional file 2: Fig. S2B) than other remaining interaction types throughout the genome. Therefore, CLAMP more strongly promotes three-dimensional interactions on the X-chromosome compared to on autosomes.

To define the properties of DIs mediated by CLAMP, we measured the genomic distance between DI anchors. The linear distance spanned by CLAMP-promoted X-linked interactions is significantly longer than those that CLAMP promotes on autosomes (Fig. 3B; Additional file 2: Fig. S2C). In contrast, the length span of interactions repressed by CLAMP is significantly shorter on the X-chromosome than on autosomes (Fig. 3B; Additional file 2: Fig. S2C).
Fig. 1 (See legend on previous page.)
file 2: Fig. S2C), consistent with the X-biased decrease in DLR after depleting CLAMP (Fig. 2, Fig S2A, B). Therefore, CLAMP promotes long-range interactions on the X-chromosome. We also measured the proximity of CLAMP-promoted and CLAMP-repressed DI anchors to CES and two randomized classes of sites within regions...
of either active or inactive chromatin based on the 9-state chromatin state model for S2 cells [43] (Fig. 3C). We found that CLAMP-promoted regions are closer to active chromatin regions than CLAMP-repressed regions. In contrast, CLAMP-repressed regions are closer to inactive chromatin than active chromatin regions. Also, CES are in closer proximity to CLAMP-promoted DI anchors and more distant from CLAMP-repressed DI anchors than other active chromatin regions (Fig. 3C). Overall, CLAMP promotes long-range contacts on the X-chromosome that enhance three-dimensional interactions involving active chromatin and CES.

CLAMP promotes three-dimensional interactions at active chromatin regions including CES

Next, we defined the relationship between DIs that are regulated by CLAMP and the enrichment of the CLAMP protein. First, we used available CLAMP ChIP-seq data [44] from S2 cells to generate a high-confidence list of CLAMP peaks and average profiles of CLAMP peak enrichment at DI anchors (see "Methods" section). We found that CLAMP-promoted DI anchors more frequently contain CLAMP peaks than CLAMP-repressed DI anchors on both the X-chromosome and autosomes (Fig. 5A). Therefore, CLAMP-promoted contacts are more likely to be directly linked to CLAMP function than CLAMP-repressed contacts. Furthermore, we found that 60% of CLAMP-promoted DI anchors on the X-chromosome are occupied by CLAMP, which is greater than CLAMP occupancy at CLAMP-repressed anchors on the X-chromosome or CLAMP-promoted and CLAMP-repressed anchors on autosomes. While CLAMP frequently binds to DI anchors, the ChIP-seq occupancy of CLAMP is not more enriched at these genomic locations where CLAMP regulates three-dimensional genomic interactions compared to CLAMP sites that do not occur at DI anchors (Fig. 5A). Therefore, we hypothesize that the presence of additional cofactors within active and/or inactive chromatin modulates the ability of CLAMP to influence three-dimensional interactions.

To test this hypothesis, we first measured the chromatin states, as defined by the *Drosophila* 9-state chromatin model [43], that are present at CLAMP-promoted and CLAMP-repressed DI anchors. We found that across all chromosomes, 60.5% of the chromatin states present within CLAMP-promoted DI anchors represent active chromatin and the remaining 39.5% of chromatin states represent inactive chromatin (Additional file 2: Fig S3D). In contrast, 70% of the chromatin states present within CLAMP-repressed DI anchors represent inactive states including enrichment for Polycomb-mediated repression, while the remaining 30% of chromatin states represent active chromatin across all chromosomes (Additional file 2: Fig S3E). Additionally, at CLAMP-repressed DI anchors, there is an enrichment of the chromatin state corresponding to pericentromeric heterochromatin (Additional file 2: Fig. S2E), which correlates with our visual observation that many pronounced off-diagonal changes in our differential Hi-C maps occur in regions proximal to centromeres (Fig. 1C).

To further quantify the relationship between DI anchors and chromatin states, we computed Jaccard similarity coefficients as ratios ranging from 0 to 1 [45]: the larger the Jaccard coefficient (i.e., closer to 1), the more similar two sets of genomic regions are to each other. CLAMP-promoted DI anchors have more similarity to active chromatin states (Jaccard coefficient: 0.328) than inactive chromatin states (Jaccard coefficient: 0.220). In contrast, CLAMP-repressed DI anchors have more similarity to inactive chromatin states (Jaccard coefficient: 0.297) than active chromatin states (Jaccard coefficient: 0.090). Therefore, CLAMP primarily promotes interactions within active regions; interactions that form after *clamp* RNAi are often within inactive regions of the genome. Moreover, the ability of CLAMP to promote genomic interactions within active chromatin regions (Additional file 2: Fig. S2D; Fig. S2E) is consistent with its ability to activate gene expression and open chromatin on the active male X-chromosome more frequently than on autosomes [25].

CLAMP and MSLc both mediate three-dimensional interactions at CES

We and others previously reported that CLAMP and MSLc function synergistically to target each other to the X-chromosome and increase occupancy of both factors in vivo and in vitro [20, 21]. In addition, MSLc modulates chromatin accessibility specifically within CES, in contrast to CLAMP which not only functions at CES, but also enhances chromatin accessibility of the entire male X-chromosome [25, 28]. Furthermore, long-range interactions at several CES occur in an MSL2-dependent manner [29, 46]. We hypothesized that both CLAMP and MSL complex function to regulate three-dimensional interactions at CES. Therefore, we also processed and performed differential interaction analysis with available Hi-C data (GSE58821) from replicated dilution Hi-C experiments that compare RNAi depletion of two MSLc components (MSL2 and MSL3) with a matched control (*gfp* RNAi) experiment [28] (Fig S1B). In contrast to our *clamp* RNAi experiments, we did not identify significant DIs following *msl2* RNAi, and *msl3* RNAi resulted in only 1 significant DI when compared to matched *gfp* controls (Additional file 3: Table S4). Therefore, MSLc does not
modulate three-dimensional interactions detectable by dilution Hi-C consistent with previous reports (Ramírez et al. 2015).

To validate our Hi-C findings that CLAMP regulates three-dimensional organization and generate a higher resolution subset of DIs, we performed circularized chromosome conformation capture with high-throughput sequencing (4C-seq) at four CES in close proximity to regions containing many DIs identified by Hi-C. In addition, we assessed the function of MSL2 and the GAF protein (encoded by the trl gene), which is a GA-binding zinc-finger protein similar to CLAMP. CLAMP and GAF are present in the same insulator complex [44, 47–49]. However, in contrast to CLAMP, GAF has only a modest function in MSLc recruitment and is depleted within CES because CLAMP outcompetes GAF for binding to the long GA-rich sequences that are present within CES [44, 50]. Therefore, we hypothesized that CLAMP and MSL2 but not GAF regulate three-dimensional interactions at CES.

To test this hypothesis, we performed 4C-seq experiments in biological duplicate in Drosophila S2 cells following previously validated RNAi depletion of gfp (control), msl2, clamp and trl [20, 44] and confirmed depletions by qRT-PCR (Additional file 2: Figs. S3A; Additional file 3: Table S1; Additional file 4: Table S2). We identified high-frequency cis-interacting regions and performed differential interaction analysis for each viewpoint using the 4C-ker pipeline [51] (Additional file 2: Fig. S3). Consistent with our Hi-C analysis, we found that 59% of high-confidence cis-interactions link our four CES viewpoints to regions that also contain a CES (Additional file 2: Fig S4B; Additional file 5: Table S3). After visualizing differential interactions for each viewpoint (Additional file 6: Table S4), we pooled all identified DI anchors from each 4C-seq viewpoint to increase the number of DIs for further analysis.

Next, we compared the number of DIs obtained after clamp and trl RNAi. We identified 199 total DIs after clamp RNAi and only 17 DIs after trl RNAi (Fig S4C). Therefore, even though both CLAMP and GAF are part of the same insulator complex [49], CLAMP has a stronger role than GAF in modulating three-dimensional interactions involving CES. These data are consistent with an enrichment of CLAMP versus GAF at CES [44]. Next, we measured chromatin state occurrence within CLAMP-promoted and CLAMP-repressed 4C-seq DI anchors and found similar chromatin states to those identified in our differential Hi-C analysis (Additional file 2: Figs. S3F, S3E-F). Therefore, our 4C-seq data demonstrate that CLAMP has a stronger role than GAF in regulating three-dimensional interactions at CES and validate a role for CLAMP in promoting contacts between active chromatin regions.

In addition, we measured the role of CLAMP and MSL2 in regulating three-dimensional interactions at CES by comparing our 4C-seq data after clamp and msl2 RNAi treatments. In contrast to our differential Hi-C interaction analysis which did not identify any DIs after msl2 RNAi, we observed 285 DIs after msl2 RNAi from our CES viewpoints, consistent with prior findings [29, 46] (Fig S4C). The discrepancy between the Hi-C and 4C-seq analyses for MSL2 may be due to the dilution Hi-C technique performed by Ramirez et al. which contains more technical noise because there is higher potential for spurious contacts during in-solution ligation compared with in situ ligation [36, 52, 53]. Furthermore, the Hi-C and 4C techniques have very different resolutions. We also measured X-specific chromatin state occurrence for 4C-seq DI anchors identified after msl2 RNAi and we find similar chromatin states to those identified at CLAMP-dependent 4C-seq DI anchors (Additional file 2: Fig. S3E-F). Overall, both CLAMP and MSL2 regulate three-dimensional interactions at CES, consistent with prior observations that CLAMP and MSLc modulate chromatin accessibility [25, 28] and promote each other’s occupancy [20, 21] locally at CES.

To further define the relationship between CLAMP and MSLc in mediating three-dimensional interactions, we integrated our Hi-C and 4C data with previously generated chromatin accessibility data from a Micrococcal Nuclease (MNase)-seq experiment performed in S2 cells under the same RNAi conditions (GSE99894) [25]. In MNase-seq experiments, a MNase accessibility score (MACC) greater than zero indicates that a region of chromatin is relatively accessible while a MACC score less than zero indicates that a region is relatively inaccessible compared to the average genomic accessibility. We found that CLAMP-promoted and CLAMP-repressed DIs from both Hi-C and 4C occur within regions where the chromatin is relatively accessible (MACC > 0) under control gfp and msl2 RNAi conditions (Fig. 4A, B). However, significant decreases in chromatin accessibility after clamp RNAi are observed at CLAMP-promoted and CLAMP-repressed DI anchors on the X-chromosome and CLAMP-repressed DIs on autosomes (Fig. 4A, B). Therefore, CLAMP regulates chromatin accessibility at regions where it regulates three-dimensional interactions.

Next, we asked whether CLAMP or MSL2 regulates chromatin accessibility at sites where MSL2 regulates three-dimensional interactions. We determined that MSL2-dependent 4C-seq DIs show significant decreases in chromatin accessibility after clamp RNAi and modest but not statistically significant changes after msl2 RNAi.
(Fig. 4C). Therefore, CLAMP but not MSL2 regulates chromatin accessibility at regions where MSL2 regulates three-dimensional interactions, consistent with the synergy between the two factors. Overall, the function of CLAMP in modulating three-dimensional interactions is linked to its role in altering chromatin accessibility.

**Insulator proteins have differential occupancy at genomic locations at which CLAMP regulates three-dimensional interactions**

CLAMP has been physically and functionally linked with two different insulator complexes containing either the insulator proteins GAF [49] or Su(Hw) [30]. Therefore, we hypothesized that the function of CLAMP in regulating three-dimensional interactions is mediated by differential occupancy of insulator proteins at DI anchors. To test this hypothesis, we generated high-confidence peaks and average profiles for insulator proteins from the following publicly available ChIP-seq data generated in S2 cells for the insulator proteins GAF (GSE107059), CP190, Su(Hw), Mod(mdg4) and dCTCF (GSE41354). We restricted our analysis to CLAMP-dependent DIs identified by Hi-C in order to make genome-wide comparisons and compared ChIP-seq enrichment of each factor at peaks that occur within DI anchors (DIs) to enrichment at the remaining set of peaks that occur outside of DI anchors (non-DIs).

Overall, we observed enhanced occupancy of all insulator proteins at genomic anchor points where CLAMP regulates three-dimensional interactions (DIs) compared with regions where it does not regulate three-dimensional interactions (non-DIs) (Fig. 5B–F). Furthermore, we discovered the following differences in factor enrichment: (1) overall the enrichment patterns of GAF are similar to that of CLAMP, which correlates with their presence in the same insulator complex [44]. Also, GAF enrichment as measured by ChIP-seq is depleted at CLAMP-promoted DI anchors compared with those outside of DI anchors or CLAMP-repressed anchors (Fig. 5B). (2) Su(Hw) is bound to fewer CLAMP-repressed DI anchors on autosomes compared with those on the X-chromosome, although its average enrichment at bound sites is similar (Fig. 5C). (3) Mod(mdg4), dCTCF and CP190 are all more frequently bound at CLAMP-promoted DI anchors. However, their enrichment is higher at CLAMP-repressed DI anchor sites versus CLAMP-promoted anchors. Furthermore, CP190 binds to a larger majority of CLAMP-promoted DIIs than any of the other factors tested consistent with a previously reported role for CLAMP in modulating CP190 recruitment [30] (Fig. 5D, E, F).

Overall, we observe different patterns of insulator protein occupancy within CLAMP-promoted and CLAMP-repressed DI anchors, compared to peak locations where we do not detect CLAMP-dependent three-dimensional interactions (non-DIs). CLAMP is also a member of two different insulator protein complexes [30, 44]. Therefore, CLAMP may modulate recruitment and/or function of insulator protein complexes to regulate three-dimensional interactions at DI anchors. In fact, very recent work demonstrated that CLAMP regulates the occupancy of the CP190 protein [30].

**Discussion**

Overall, our data provide key insight into how the *Drosophila* male X-chromosome forms a specific hyper-active chromatin domain in three-dimensions. CLAMP de-compacts the genomic architecture of the X-chromosome by promoting long-range interactions and preventing formation of short-range interactions. CLAMP not only tethers MSLc to the X-chromosome [20], but also promotes long-range three-dimensional interactions involving MSLc binding sites. Previously, we demonstrated that CLAMP and MSLc function...
synergistically to increase transcript levels of X-linked genes [20, 54]. Here, we show that both CLAMP and MSLc regulate the three-dimensional organization of the X-chromosome: CLAMP regulates three-dimensional interactions along the length of the entire X-chromosome while MSLc acts locally at several CES.

Throughout the genome, CLAMP promotes long-range interactions between active chromatin regions, including CES. However, we observe that CLAMP has a stronger function in mediating three-dimensional interactions on the X-chromosome than on autosomes. CLAMP is an ancient zinc-finger protein that is highly conserved across Diptera and is present on chromatin in the earliest stages of development before the more recently evolved MSLc [18, 20, 22, 23]. Transposons containing CLAMP recognition sequences transposed onto the ancient

Fig. 4 (See legend on previous page.)
X-chromosome and GA-repeats present at splice-junctions expanded, which together increased the density of CLAMP occupancy at CES [17, 18]. The enhanced role of CLAMP on the X-chromosome compared with autosomes is likely due to the increased density of CLAMP binding sites within CES on the X-chromosome [18]. Furthermore, CLAMP interacts with several insulator protein complexes [30, 49], which are known to modulate three-dimensional genomic interactions and contain CP190. We observe that the enrichment of insulator proteins is increased at sites where CLAMP functions to regulate three-dimensional interactions, compared to non-interacting control regions. Therefore, it is likely that CLAMP regulates three-dimensional organization by modulating the ability of insulator proteins to drive the formation of three-dimensional interactions. For example, CLAMP is known to alter the occupancy of CP190, a component of several different insulator complexes [30].

Overall, we demonstrate that CLAMP, a transcription factor enriched on the male X-chromosome due to synergy with MSLc [20, 21], regulates long-range genomic interactions on the male X-chromosome including those involving CES. Therefore, CLAMP-mediated three-dimensional interactions promote the formation of the three-dimensional active chromatin domain critical for dosage compensation. Further work will be required to define the functional relationships between CLAMP and all of the known insulator proteins.

**Methods**

**Cell culture conditions**

*Drosophila* S2 cells were maintained at 25°C in Gibco Schneider's *Drosophila* media (ThermoFisher Scientific) supplemented with 10% heat-inactivated fetal bovine serum (FBS) and 1.4X antibiotic–antimycotic (ThermoFisher Scientific). Cells were passaged every 2–3 days to maintain appropriate density.

**RNAi treatment**

*Generation of dsRNA for RNAi treatment*

Generation of dsRNA targeting *gfp* (control), *clamp*, *msl2* and *trl* for RNAi have been previously validated and described [4, 19, 20, 55]. PCR product was used as template to generate dsRNA with an ambion T7 MEGAscript kit (ThermoFisher Scientific); dsRNAs were purified following DNase treatment with a Qiagen RNeasy kit (Qiagen).
RNAi treatment

RNAi was performed in T75 tissue culture flasks. A total of $7 \times 10^6$ S2 cells were suspended in 6 mL of Schneider’s *Drosophila* media (without FBS) and added to a T75 culture flask containing 67.5ug of *gfp*, *clamp*, *msl2*, or *trl* dsRNA suspended in 3 mL of Invitrogen UltraPure water (ThermoFishcer Scientific). Cells were serum starved for 45 min at room temperature, then 10.5 mL of Schneider’s *Drosophila* media supplemented with 10% FBS was added. Cells were incubated for a total of 6 days as described previously [20, 44]. After 6 days, samples were collected and RNAi knockdown was validated via western blotting or qRT-PCR. The remaining cells were collected by centrifugation and resuspended to $5 \times 10^6$ cells/ml in fresh non-FBS media. Fresh formaldehyde solution (36.5–38% in H2O; Sigma Aldrich) was then added to obtain a final concentration of approximately 1% formaldehyde. Cell suspensions were then incubated at RT on a rocking platform for 10 min; 2.5 M glycine was then added (final concentration of 0.125 M) to quench the formaldehyde; the cells were incubated as before for an additional 5 min. The cell suspension was then immediately placed on ice for 15 min. The cell suspension was then aliquoted into individual tubes (5 million cells per tube). The cell suspensions were centrifuged at 4C for 5 min at 3000 RPM, supernatant was removed and then flash frozen in liquid nitrogen and stored at -80C for future processing.

Knockdown validation

For Hi-C experiments knockdown of CLAMP was validated using the Western Breeze kit (Invitrogen). Antibodies used for detection were a previously described custom rabbit anti-CLAMP (1:1000, Abcam) [22]. Mouse anti-actin (1:400,000, Sigma Aldrich) was used as a loading control.

Quantitative real-time PCR

To determine transcript abundance of *clamp, msl2*, or *trl*, RNA was calculated using the $2^{-\Delta\Delta Ct}$ method [56] using RNA extracted from 500ul of cells following the 6-day incubation using an RNaseasy Plus RNA extraction kit (Qiagen). Gene targets were amplified from cDNA using previously validated primers for *clamp, msl2, trl*, and three internal control genes (*gapdh, rpl32*, and *ras64h*) using triplicate technical replicates for each biological replicate per condition. Samples were normalized to the control *gfp* RNAi condition.

Hi-C experimental procedure

Hi-C libraries of two independent biological replicates per RNAi condition (*clamp* and *gfp*) were generated as follows:

Cell lysis and restriction enzyme digestion

Approximately 10 million formaldehyde crosslinked (crosslinking procedure described above) S2 cells were resuspended in 500ul of fresh cold lysis buffer (10 mM Tris–HCL pH 8.0, 10 mM NaCl, 10ul of 50X Protease inhibitors cocktail, 0.2% Igepal CA630 in UltraPure water) and incubated on ice for 30 min. Cell suspensions were then centrifuged for 5 min at 3000 RPM at 4C and the supernatant removed. The cell pellet was resuspended with 300ul of cold 1 x NEBuffer2. Cell suspensions were again centrifuged at 3000 RPM at 4C and the supernatant was removed. Cells were resuspended in 95ul of 1X NEBuffer2 and 5ul of 10% SDS was added. The cell suspension was homogenized by gentle pipetting and then incubated at 65C for 10 min. The cell suspension was then placed on ice and 200ul of 1X NEBuffer2 along with 60ul of 10% Triton X-100 (to quench the SDS) was added. The cell suspension was incubated at 37C for 15 min. Lysis efficiency and nuclei integrity was checked via microscope. The samples were centrifuged for 5 min at 3000 RPM at 4C and the supernatant was removed. The pellet was resuspended in 300ul of 1X NEBuffer2 and 400 U of HindIII restriction enzyme was added. The samples were incubated overnight at 37C. The following morning an additional 200U of HindIII was added to each of the samples and they were incubated an additional 2 h at 37C.

End-repair, labeling, in-nuclei ligation, and crosslink reversal

The samples were centrifuged for at 3000 RPM for 5 min at 4C and the supernatant removed. The samples were resuspended with 250ul of 1X NEBuffer and 50ul of the following mix was added: 1.5ul of 10 mM dATP, 1.5ul of 10 mM dGTP, 1.5ul of 10 mM dCTP, 37.5ul of 0.4 mM Biotin-11-dCTP, 1ul of 50U/ul DNA Polymerase I Large (Klenow) fragment, and 7ul of UltraPure water. The sample were incubated at 37C for 45 min and then incubated at 65C for 15 min. The samples were centrifuged for 5 min at 3000 RPM and the supernatant discarded. The pellet was resuspended in 1.195 mL of the following ligation mix: 120ul of 10X T4 DNA Ligase Buffer, 100ul of 10% Triton X-100, 12ul of 10 mg/ml BSA and 963ul of UltraPure water. 5ul of 2000U/ul T4 DNA ligase was then added to each sample and the samples were incubated at 16C overnight. The samples were centrifuged for 5 min at 3000 RPM and supernatant removed. The samples were resuspended in 400ul of 1X NEBuffer2 and 10ul...
of 10 mg/mL RNase A was added and the samples were incubated for 15 min at 37°C at 200 RPM shaking, 20 ul of 10 mg/ml Proteinase K was added and the samples were incubated overnight at 65°C to reverse crosslinks at 300 RPM shaking. The next morning an additional 20 ul of Proteinase K was added again and the samples were incubated for 2 h at 65°C.

DNA purification
The samples were cooled to room temperature and 400 ul of an equal volume of phenol/chloroform/isoamyl alcohol was added to each sample and the solution was mixed vigorously. The samples were centrifuged for 5 min at 13,000 RPM and the upper aqueous phase was transferred to a new tube. DNA was precipitated by adding 40 ul of NaAc pH 5.2 and 1 mL of 100% ethanol. The samples were incubated for 30 min at –80°C and then centrifuged at 13,000 RPM for 30 min at 4°C. The supernatant was discarded and pellet was washed with 1 mL of ethanol 70%. The sample was centrifuged for 15 min at 13,000 RPM at 4°C. The supernatant was discarded and the pellet was air-dried for 5 min. The pellet was resuspended in 1X TE buffer.

DNA shearing and size selection, biotin pull-down and sequencing library preparation were performed as described in [36]. Multiplexed libraries were sequenced on an Illumina Hi-Seq 2500 configured for 150-bp paired-end reads.

4C-seq experimental procedure
4C-seq libraries were generated from S2 cells. Nuclear extraction and crosslinking were carried out as described in Hi-C experimental procedure above. The remainder of the 4C-seq procedure was carried out as previously described [57]. Csp6I, DpnII or NlaIII were used as primary or secondary restriction enzymes. Primer sequences for each viewpoint and condition were generated using 4C primer design (https://mnlab.uchicago.edu/4Cpd/) and are listed in Additional file 4: Table S2. Independent biological replicates of multiplexed libraries were sequenced on separate lanes of an Illumina Hi-Seq 2500 configured for 150-bp paired-end reads.

Datasets
CES locations were obtained from GSE9271. Hi-C following RNAi against msl2, msl3, or gfp was obtained from GSE58821. MACC data were obtained from GSE99894. S2 cell chip-seq data sets for CP190, SuHw, Mod, and dCTCF GSE41354. S2 cell chip-seq data sets for GAF and CLAMP were obtained from GSE107059.

Raw and processed sequencing data generated are deposited to NCBI GEO under accession GSE130546.

Hi-C analysis
Hi-C data were processed using the dm6 Drosophila reference genome [58] using HiC-Pro pipeline version 2.7.8 [59] for read mapping (MIN_MAPQ=15), filtering and quality checks to generate valid read pairs and interaction matrices. GenomeDisco version 1.0.0 [38] was utilized to determine concordance between biological replicates. Custom scripts were used to convert Hi-C-Pro interaction matrices to a format suitable for input into GenomeDisco. For matrix visualizations valid pairs were processed into.hic files using the hicpro2juicebox script provided with Hi-C-Pro. Matrices were visualized using Juicebox [60] with KR matrix balancing.

PCA and DLR analysis
Valid pairs for biological replicates were merged and imported into Homer version 4.10 [61] using the command makeTagDirectory with the parameter -format HiCsummary. Principal component analysis (PCA) was performed using Homer with the command runHiCpca.pl at resolutions of 10, 20 and 50 kb. Distal to local log2 ratio analysis was performed using Homer with the command analyzeHiC with the following parameters: -res 5000 -window 15,000 -compactionStats auto -dlrDistance 250,000. Distal to local log2 ratio differential analysis was performed using the command subtractBedGraphsDirectory.pl with the parameter -center.

Identification of high-confidence long-range interactions
20-kb resolution contact matrices corresponding to the control (gfp RNAi) conditions were generated and iterative corrected using HiC-Pro. The resultant contact matrices were then adapted for Fit-Hi-C version 2.0.5 [40] using the hicpro2Fit-Hi-C.py script provided with Hi-C-Pro. Raw contact matrices and ICE biases were used as input into Fit-Hi-C to identify significant contacts with the following parameters: -L 5,000,000 -U 15,000,000 -v -b 100 -p2. Significant contacts were filtered for those with q<0.01 and the intersection of significant contacts (q<0.01) from the replicates was used as the high-confidence interaction set.

Identifying differential interactions (DIs)
diffHiC version 1.14.0 [41] which uses the edgeR [42] package for differential statistics was utilized to identify differential interactions. Valid pairs for from HiC-Pro for each replicate was imported into diffHic using the savePairs function. Low-abundance read pairs were filtered out and the resulting data were normalized with TMM normalization and trended biases were removed. Differential interactions were identified using a bin size of 30 kb, an FDR target of 0.05, and the functions diClusters,
combineTests, and getBestTest. No threshold was applied for log ratio.

4C-Seq analysis
4C sequencing reads were demultiplexed based on index sequences and inline barcodes. Sequencing reads corresponding to the reading primer were aligned to a reduced genome of unique sequences adjacent to restriction enzyme sites derived from the dm6 Drosophila reference genome [58] using Bowtie2 version 2.3.0 [62] with the following parameters: -N 0 -5 24 -3 101 –very-sensitive, which removes barcode and primer sequences and trim the read length to 25 bp prior to mapping. The aligned reads were then processed as described for use with the 4C-Ker pipeline [51] which uses a three-state hidden Markov model to find chromosome-wide interactions and DESeq2 [63] to perform differential analysis. Quality checks were performed within the 4C-Ker pipeline. Cis analysis was performed for each viewpoint using k=10 and p-value cutoff of 0.05 was used for differential analysis. Viewpoints were visualized using IGV [64].

Identification of high-confidence long-range interactions
A set of high-confidence interactions for each viewpoint was determined by intersecting the significant interactions individually detected in each replicate of the control (gfp RNAi) condition using BEDTools version 2.27.1 [45].

Comparison of 4C and Hi-C high-confidence interactions
High-confidence Hi-C interactions involving each 4C-seq viewpoint was determined by intersecting the viewpoint coordinate sequence with the Hi-C high-confidence interactions set. This set of interactions was then intersected with the 4C-seq high-confidence interactions set to determine overlapping pairwise high-confidence interactions identified in both experiments. All intersections were performed using BEDTools.

Chromatin state annotation
Chromatin states were from the modENCODE project (DCC id: modENCODE_3363) [43, 65] and were lifted over to dm6 with the USCS liftOver tool (http://genome.ucsc.edu) [66]. Ratios were normalized to account for the per chromosome abundance of each individual chromatin state. Jaccard similarity scores were computed using BEDTools.

Distance calculations
Distances to nearest CES were calculated using BEDTools. Controls were conducting by 100 permutations of randomly shuffling CES restricting reshuffled regions to either active or inactive regions as defined by chromatin state [43].

Chromatin accessibility (MACC) analysis
MACC data generated following RNAi knockdown of gfp (control), clamp, or msl2 were lifted over to dm6 with the USCS liftover tool. MACC data were intersected with genomic regions of interest using BEDTools to determine corresponding MACC scores. Other analyses were performed in the python programming environment (https://www.python.org).

Chip-seq data processing and generation of high-confidence peak sets
For each ChIP-seq dataset used, the respective sequencing reads were downloaded and mapped to release 6 D. melanogaster genome (dm6) [58] using Bowtie2 with parameter -N 1. Reads with a MAPQ < 30 and PCR duplicate reads identified using Picard MarkDuplicates version 2.9.2 [67] were removed using SAMtools version 1.9 [68]. Reads mapped to dm3 blacklisted regions (https://sites.google.com/site/anshulkundaje/projects/blacklists) lifted over to dm6, using USCS liftOver, were also removed. In cases where biological replicates were not available the aligned reads were split into pseudoreplicates. MACS2 version 2.1.1 [69] was used to identify peaks with the following parameters: –nomodel -B –SPMR –keep-dup all -f AUTO -g dm -p 0.01. MACS2 was also used generate fold enrichment (default parameters) bedGraphs for each factor. In order to reduce the number of false positive peaks the irreproducible discovery rate (IDR) was calculated using IDR version 2.0.3 [70] using the MACS2 peak score calculated for each replicate experiment (biological replicate in the case of GAF and CLAMP; pseudoreplicate in the case of CP190, SuHw, Mod, dCTCF) as input to IDR. Peaks with an IDR < 0.01 were retained (In cases where there were more than two biological replicates pairwise IDR comparison of each replicate was made and the longest resulting peak list was used). BEDTools and USCS bedGraphToBigWig [71] tool was used to convert bedGraphs to bigwig format. Average profiles were generated using deepTools version 3.1.0 [72].

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13072-021-00399-3.
Acknowledgements
We thank Dr. Leila Rieder for critical reading of the manuscript and Dr. John Urban and Dr. Nicola Neretti for advice on computational analysis.

Authors' contributions
WJ performed all experiments and analyses. EL provided advice on experiments. WJ and EL wrote the manuscript. All authors read and approved the final manuscript.

Funding
This research was supported by NIH grant: 5R35GM126994 to EL. WJ was funded by a NSF GRFP Fellowship and the HHMI Gilliam Fellowship.

Availability of data and materials
All data are available on NCBI GEO (see "Methods" section). Furthermore, processed source data are attached to this manuscript in supplementary files.

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Both authors consent for this work to be published.

Competing interests
We declare no competing interests.

Received: 13 January 2021  Accepted: 16 May 2021
Published online: 29 June 2021

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