Adaptive Evolution and the Birth of CTCF Binding Sites in the *Drosophila* Genome

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Abstract

Changes in the physical interaction between cis-regulatory DNA sequences and proteins drive the evolution of gene expression. However, it has proven difficult to accurately quantify evolutionary rates of such binding change or to estimate the relative effects of selection and drift in shaping the binding evolution. Here we examine the genome-wide binding of CTCF in four species of *Drosophila* separated by between ~2.5 and 25 million years. CTCF is a highly conserved protein known to be associated with insulator sequences in the genomes of human and *Drosophila*. Although the binding preference for CTCF is highly conserved, we find that CTCF binding itself is highly evolutionarily dynamic and has adaptively evolved. Between species, binding divergence increased linearly with evolutionary distance, and CTCF binding profiles are diverging rapidly at the rate of 2.22% per million years (Myr). At least 89 new CTCF binding sites have originated in the *Drosophila* melanogaster genome since the most recent common ancestor with *Drosophila simulans*. Comparing these data to genome sequence data from 37 different strains of *Drosophila melanogaster*, we detected signatures of selection in both newly gained and evolutionarily conserved binding sites. Newly evolved CTCF binding sites show a significantly stronger signature for positive selection than older sites. Comparative gene expression profiling revealed that expression divergence of genes adjacent to CTCF binding site is significantly associated with the gain and loss of CTCF binding. Further, the birth of new genes is associated with the birth of new CTCF binding sites. Our data indicate that binding of *Drosophila* CTCF protein has evolved under natural selection, and CTCF binding evolution has shaped both the evolution of gene expression and genome evolution during the birth of new genes.

Introduction

Gene regulation is a major driver in the generation of morphological diversity [1,2]. Transcriptional regulators determine spatial and temporal patterns of mRNA level by binding to cis-regulatory DNA elements. Many previous studies have demonstrated that changes at the level of protein-DNA interactions can account for specific phenotypic differences observed in nature [1,3]. Genome-wide studies have shown that binding of transcriptional regulators evolves substantially between different species [4–9]. Although in *Drosophila*, the binding profiles of some regulatory factors involved in embryonic development, such as the Twist protein, are relatively more conserved [5,8]. Yet it remains an open question to what extent such protein–DNA binding evolution is adaptively and functionally significant or whether it reflects drift. In order to address this question, regulatory factors must be mapped in multiple related species and the results interpreted in the light of intraspecific and interspecific cis-regulatory DNA variation.

Insulator proteins participate in the marking of boundaries for genomic regulatory units by binding to DNA sequences that mediate intra-/interchromosomal interactions [15–19]. These protein–DNA complexes are thought to function as barriers against the spread of heterochromatin or to regulate enhancer–promoter communications by preventing inappropriate interactions, although the precise molecular mechanism by which they act is not known [12,14,15]. Recent studies have suggested that insulator complexes may also participate in the global nuclear organization of active and inactive chromatin domains via mediating intra-/interchromosomal interactions [16–19]. The broad functions possessed by insulator proteins make them a key
Author Summary

A large proportion of the diversity of living organisms results from differential regulation of gene transcription. Transcriptional regulation is thought to differ between species because of evolutionary changes in the physical interactions between regulatory DNA elements and DNA-binding proteins; these can generate variation in the spatial and temporal patterns of gene expression. The mechanisms by which these protein–DNA interactions evolve is therefore an important question in evolutionary biology. Does adaptive evolution play a role, or is the process dominated by neutral genetic drift? Insulator proteins are a special group of DNA-binding proteins—that is, directly by means of repressing genes such as enhancers and promoters. Additionally, insulator proteins can limit the spreading of chromatin condensation and help to demarcate the boundaries of regulatory domains in the genome. In spite of their critical role in genome regulation, little is known about the evolution of interactions between insulator proteins and DNA. Here, we use ChIP-seq to examine the distribution of binding sites for CTCF, a highly conserved insulator protein, in four closely related Drosophila species. We find that genome-wide binding profiles of CTCF are highly dynamic across evolutionary time, with frequent births of new CTCF-DNA interactions, and we demonstrate that this evolutionary process is driven by natural selection. By comparing these with RNA-seq data, we find that gain or loss of CTCF binding impacts the expression levels of nearby genes and correlates with structural evolution of the genome. Together these results suggest a potential mechanism of regulatory re-wiring through adaptive evolution of CTCF binding.

CTCF shows similar binding distributions in intergenic, promoter, intronic, and exonic sequences among the four species (Figure 1C) and in a pattern consistent with previous genomic mapping studies of CTCF in both fly and human [13,20]. Importantly, the position weight matrices of the consensus motifs for CTCF-bound sequences are virtually identical among the four species (Figure 1D). These motifs are also similar to the in vitro identified CTCF consensus motif [27] and to previously identified motifs from Drosophila, human, and other vertebrates [13,20,23]. CTCF protein evolution is highly constrained (Table S2, Figure S5), especially the 253 amino acid DNA binding domain (Figure S5), for which there are only 1, 4, and 36 amino acids diverged between D. simulans, D. yakuba, and D. pseudoobscura, respectively. We calculated the occurrences of each species-specific motif in the CTCF binding sites and obtained similar percentages among the four species at various thresholds (Table S3). These results confirm the conservation of CTCF binding motifs among Drosophila species and indicate that any evolutionary patterns we observe are most likely due to changes in the cis-regulatory target DNA sequences of CTCF.
CTCF Binding Evolves Rapidly

A straightforward way to assess binding conservation or divergence is to directly compare the boundaries of identified peak regions between each species (Table S7, also see Materials and Methods) with genome-wide alignment. However, by using independent analyses in each species, actual conserved binding sites are likely to be identified as diverged between species due to false negatives being scored as “diverged.” To avoid this problem and to quantitatively explore the evolutionary dynamics of CTCF binding profiles across species (Figure 2A), we developed a *D. melanogaster*–centric analysis approach to examine the between-species CTCF occupancy on orthologous DNA sequences in light of within-species binding variation (Figure S6, also see Materials and Methods). In brief, instead of directly comparing the binding region boundaries between each species, the approach we took translated uniquely mapped sequence reads in the non-*D. melanogaster* species into *D. melanogaster* genome (Table S1B), thus allowing quantitative modeling of within- and between-species read number data using an ANOVA-like linear categorical model to partition variances of local read number data in each binding region. We thus identified *D. melanogaster*–specific, non-*D. melanogaster*–specific, and shared binding events for each paired species accordingly (Figure 2B, Table S5). Our method yields highly reliable conservation and divergence information of the *D. melanogaster* binding sites between each species since the False Positive Rate (identifying shared binding sites as *D. melanogaster* specific) of the linear categorical model is estimated to be 0.35% using simulated data (Materials and Methods) and the overall analysis pipeline error rate (False Positive Rate plus False Negative Rate) is estimated to be less than 2% using a “gold standard” data set (Materials and Methods). However, since different *Drosophila* genomes have different assembly and annotation quality, which are all based on and are poorer than the *D. melanogaster* genome, inevitably the translated read number data from some regions of the non-*D. melanogaster* species will be smaller than they would be in an ideal situation. As a result, there is generally reduced power in detecting non-*D. melanogaster*–specific binding events compared to *D. melanogaster*–specific and shared binding events (Figure 2B, Table S4).

Because loss of orthologous sequences among species is often driven by large-scale genome evolution instead of local nucleotide substitutions or small insertion-deletions (indels) [40], diverged CTCF binding events in regions with or without orthologous sequences have different biological interpretations. With a criterion of at least 50% sequence identity for orthology assignment, we identified binding events in orthologous regions between each species pair, which we refer to as two-way orthologous binding (TWOB). Similarly, we identified binding
Figure 2. Diverged CTCF binding between *Drosophila* species. (A) Evolutionary dynamics of CTCF binding profiles at the Bithorax complex region. The four colored wiggle file tracks show the ChIP CDP enrichment scores estimated from our quantitative analysis pipeline for the four species: *D. melanogaster* (blue), *D. simulans* (green), *D. yakuba* (orange), and *D. pseudoobscura* (purple). The four tracks are at the same scale, with the adaptive evolution of CTCF binding sites.
events in regions with orthologous sequence counterparts in all four species and refer to those as four-way orthologous binding (FWOB). With the analysis pipeline described above, we identified 2,267 binding events for D. melanogaster (Figure 2B, Table S5). Since genome assembly imperfections and gaps among the non-D. melanogaster species lead to an underestimate of binding events in these genomes, we used the percentage of diverged binding events with respect to D. melanogaster, which has the best refined genome assembly map, to measure pair-wise binding divergence. Naturally, the rate of binding site evolution must be greater than the error rate (<2%) in order to be detected. For the different species pair-wise comparisons with D. melanogaster, approximately 20%, 30%, and 70% (19.67%, 29.11%, and 74.06% of all binding sites; 17.34%, 28.05%, and 68.37% of TWOBs; and 15.24%, 26.31%, and 68.06% of FWOBs) were identified as diverged from D. simulans, D. yakuba, and D. pseudoobscura, respectively (Figure 2B and 2C, Table S5). These values are not only consistent with each other but also highly comparable to divergence rates estimated with different parameters (Table S5), or with a subset of sites filtered for high input sequence coverage (“high sequence coverage sites”) to ensure that the binding evolution was not an artifact of low sequence coverage in one or more of the species (Table S6) or using alternative methods (Tables S7 and S8). When plotted against species divergence time, these values show a clear linear trend (Figure 2C). We fit a simple linear regression for the TWOBs, and we estimated the divergence rate of CTCF binding as 2.22% per Myr (Student’s test, p<0.05, R-squared=0.99). This divergence rate is lower than synonymous substitution rates (6.34% per Myr, [40]) but substantially higher than the protein sequence divergence rate (1.19% per Myr, [40]) and non-synonymous nucleotide substitution rate (0.4% per Myr, [40]) in Drosophila, indicating that, although constrained, CTCF binding evolves relatively rapidly. This linear pattern of binding divergence remains stable when different peak calling stringencies were applied (Tables S5 and S6).

Estimates of binding divergence/conservation rates can depend on the choice of analysis methods, which have different associated false positive and false negative rates. In order to gain an unambiguous comparison of binding divergence between CTCF and other transcription factors in Drosophila, we also applied our analysis pipeline to previously published Twist comparative data [8] and reciprocally applied the He et al. method [8] to our data (Tables S8, S9, and S10). In both comparisons, we obtained a larger estimate of binding divergence in the CTCF data (Tables S8 and S10), indicating that CTCF binding is evolving faster than binding of the developmental regulatory transcription factor Twist.

New CTCF Binding Sites Originate Frequently

By combining between-species binding appearance and absence results, we grouped CTCF binding events into 15 different evolutionary categories (Figure 2D). Regardless of evolutionary category, we found that CTCF binding events are distributed similarly between various genomic locations (Table S11), showing no biases in binding evolution according to genomic position. We next inferred the evolutionary age of each D. melanogaster binding site by assigning its origination on the Drosophila phylogeny with parsimony (Figure 2D, Table S12). Whether we examined all binding sites or considered only FWOBs, more than 60% (1,533/2,267 for all binding and 655/1,030 for FWOBs) of D. melanogaster binding sites originated after the split of the melanogaster group from pseudoobscura group, and thus less than 40% were inherited from the common ancestor of these two major clades (Figure 2D). Notably 89 FWOBs were newly gained specifically on the D. melanogaster branch (Figure 2D), leading to a conservative estimate of ~36 binding gains per million years (89 binding events/2.5 Myr since last common ancestor of D. melanogaster and D. simulans). Interestingly, there are only 39 D. melanogaster-specific new genes identified [41,42], resulting in an average of ~16 gene gains per Myr. This result indicates that in Drosophila new regulatory elements bound by CTCF are evolving at a higher rate than new genes. In contrast to the large number of newly gained binding sites, there are no FWOBs identified as lost in D. melanogaster (Figure 2D). Although underdetection of non-D. melanogaster binding sites in our analysis could lead to failure in observing D. melanogaster lineage-specific loss, the asymmetric patterns of CTCF binding site gains are also observed along the D. simulans branch (85 gains and five losses, Figure 2D), the D. yakuba branch (113 gains and 21 losses, Figure 2D), or when taking only the three species in the melanogaster group into consideration (Figure S7) or when using only high-sequence coverage sites (Table S13), indicating that gain of binding is evolutionarily favored. This pattern is not likely due to ascertainment biases of highly conserved genomic regions associated with FWOBs, because we also observed large numbers of binding gains and small numbers of binding losses for binding events genome-wide (i.e., the pattern
holds for sites that have not been filtered for sequence conservation) (Figure 2D, Figure S7, Table S13). Such a robust pattern suggests that positive selection may be driving the creation of new sites, which we sought to explore further.

CTCF Binding Evolution Is Correlated with Sequence Evolution

We investigated sequence divergence of cis-DNA elements associated with CTCF binding evolution, since CTCF protein and its binding preference are highly conserved (Figure 1C, Figure S6, Table S2). We examined the 201 bp elements comprised of the summit coordinate of each binding peak plus the two 100 bp flanking regions (i.e., CTCF-201 sites). We found that the median PhastCons scores [43] of the conserved D. melanogaster binding sites are significantly higher than those of the diverged sites (Figure S8A). A similar pattern was observed when we calculated the percentage of between-species sequence identity (Figure S8B), indicating that CTCF binding evolution is correlated with levels of sequence conservation. Because motifs are special sequence features associated with protein–DNA interactions, we next examined the relationships between motif evolution and CTCF binding evolution. For each species pair, we counted the number of species-specific motif occurrences in the corresponding orthologous sequences of each binding site. Binding sites that contained at least one motif in both sequences were defined to have conserved motifs. This is a crude way of defining motif conservation in binding sites, but nonetheless we still observed a significantly higher proportion of conserved binding sites that contain conserved motifs than diverged binding sites (two-sided Fisher’s exact test, p<0.05, Figure 3A), confirming that cis-regulatory target sequences are correlated with CTCF binding evolution.

CTCF Binding Evolution Is Shaped by Natural Selection

We next examined genomic DNA variation associated with CTCF binding events for signatures of selection. We first assessed whether purifying selection may play a role in shaping CTCF binding evolution. Purifying selection acting on polymorphic variants is expected to keep them at lower frequencies in a population, leading to a relatively higher number of segregating sites and therefore a more negatively skewed Tajima’s D value than expected under neutrality [44]. We calculated Tajima’s D using DNA polymorphism data from 37 D. melanogaster inbred lines (www.dpbgp.org) for the core consensus binding motifs (CTCF-motif sites) identified within the D. melanogaster TWOBs and FWOBs. As expected, the distribution of Tajima’s D values for nonsynonymous sites is negatively skewed compared to the synonymous sites (Wilcoxon rank sum test, p<2.2e-16; Figure 3B). This pattern also extends to the CTCF-motif sites when they are compared to synonymous sites in neighboring protein-coding genes (Wilcoxon rank sum test, p<0.0001; Figure 3B). Interestingly, the distribution of Tajima’s D values for CTCF motifs within CTCF binding sites is comparable to 3’UTR and 5’UTR sequences, while it is significantly more negatively skewed than intergenic sequence (Wilcoxon rank sum test, p<0.02). Thus, CTCF binding appears to be subject to stronger purifying selection than synonymous and intergenic genomic sequences. To explore whether these trends varied depending on evolutionary conservation of binding, we separated the TWOB CTCF-motif sites into subgroups associated with conserved binding (conserved TWOB) and diverged binding (diverged TWOB). We observed a more negatively skewed Tajima’s D distribution in the former group (Wilcoxon rank sum test, p<0.01; Figure 3B). Similar analyses of CTCF motifs within FWOB binding sites were carried out by designating binding events with evolutionary age <2.5 Myr as young FWOB and >6 Myr as old FWOB. Again a more negatively skewed Tajima’s D distribution was observed in the old FWOB group (Wilcoxon rank sum test, p=0.11; Figure 3B). We observed similar patterns of Tajima’s D in the CTCF-201 sites (Figure S9) as well as in high-sequence coverage sites (Figure S10). These results indicate that the more conserved CTCF binding sites are subject to stronger purifying selection and therefore are more constrained than the less conserved sites, as one might expect.

Using the same polymorphism data and employing D. yakuba as an outgroup, we counted the number of fixed and polymorphic nucleotides within CTCF motifs present within different classes of binding sites. Overall, significant excesses of fixed nucleotide changes are observed in groups of CTCF binding sites (except FWOB and Old_FWOB groups) when compared to synonymous nucleotide changes at nearby genes (Chi-square test, p<0.001; Tables S14 and S15), indicating that positive selection has shaped CTCF binding evolution. By extending the McDonald-Kreitman test framework [45–47], we estimated α, the proportion of between-species divergence fixed by positive selection for each subgroup of sites (as described earlier for Tajima’s D; Figure 3C). We found that the young FWOB sites show a significantly higher shared α value (0.25, p<0.0001) than the old FWOB sites (0.25 versus −0.0673; log likelihood ratio test for group comparison, p<0.05; Figure 3C), and a similar trend was observed between diverged TWOB (0.2237, p<0.0001) and conserved TWOB sites (0.2237 versus 0.0526; log likelihood ratio test for group comparison, p<0.05; Figure 3C). Since synonymous sites are usually constrained by codon usage [48], we also used a set of pre-characterized small introns that are believed to have evolved neutrally [49] as a neutral reference. Again we observed that the diverged TWOB (0.3598, p<0.0001) and young FWOB sites (0.4265, p<0.0001) shared significantly higher α values than their counterparts (log likelihood ratio test for group comparison, all p<0.001; Figure 3D). The same pattern remains when using D. simulans as an outgroup (Figure S12) or using CTCF-201 sites for the calculation (Figure S11). This trend of higher shared α from more diverged sites is also observed in the high-sequence coverage sites (Figure S13). These observations indicate that gain of new CTCF binding events are likely driven by positive selection. To further confirm the role of positive selection in the birth of new CTCF binding events, we carried out a multilocus HKA test ([50]; Materials and Methods). By comparing the young sites to the old sites as well as the neutral small introns, we observed significantly reduced polymorphism in the young sites, suggesting strong directional positive selection (Tables S16 and S17).

In order to test whether this phenomenon of selection-driven binding evolution is CTCF specific or more general, we applied the same population genetic analysis to the available comparative data for Twist [8]. Consistent with the higher binding conservation level, we observed a stronger purifying selection signal and weaker positive selection signal in Twist binding sites than in CTCF binding sites (Figure S20). Interestingly, we found a similar pattern of a higher positive selection signature in the diverged binding sites than the conserved sites for Twist (Figure S20).

CTCF Binding Evolution Is Associated with Expression Divergence

Since CTCF participates in transcriptional control through organizing and delineating regulatory domains [10–13,19] and gain of CTCF binding appears to be driven by positive selection, naturally we wondered if there were any detectable effects on gene expression. We measured mRNA transcript abundances for WPP samples in D. melanogaster, D. simulans, and D. yakuba using RNA-seq.
and estimated the interspecies expression change for every orthologous gene pair between *D. melanogaster/D. simulans* and *D. melanogaster/D. yakuba* through a generalized linear model framework, cataloging the evolutionary status of each gene as either “stable” or “diverged” (Materials and Methods) thereafter.

We focused on the nearest genes to the *D. melanogaster* TWOB sites and grouped them into genes near conserved TWOB sites and genes near diverged TWOB sites. Since diverged TWOBs resulted from either binding gain in *D. melanogaster* or binding loss in *D. simulans* or *D. yakuba*, regulation of these genes by CTCF might have been altered. Consistent with this hypothesis, the proportion of genes with diverged expression near diverged TWOB sites is significantly greater than near conserved TWOB sites (Fisher’s exact test, \( p < 0.05 \); Figure 4A,B). We obtained a similar result when comparing between genes near young FWOB sites and genes near old FWOB sites (Fisher’s exact test, \( p < 0.05 \); Figure 4A,B). Moreover, the proportions of genes with diverged expression near conserved TWOB and near old FWOB sites are smaller than the genome-wide average (Fisher’s exact test, \( p < 0.05 \); Figure 4A,B).

![Figure 3. Selection on CTCF motif sites.](image)

**(A)** Proportion of binding sites with conserved motifs. The bar plots show proportions of *D. melanogaster*-specific (pink) and shared (green) binding sites that have conserved motifs between each species pair. A binding site is defined as having conserved motifs if there is at least one species-specific motif identified in the corresponding orthologous sequences. The \( p \) value cutoff for FIMO motif searching here is 0.005. For any species pair, the proportion of conserved (here shared) binding sites having conserved motifs is significantly higher than the diverged (here *D. melanogaster*-specific) binding sites. Significance levels: * \( p < 0.05 \); ** \( p < 0.01 \), two-sided Fisher’s exact test. **(B)** Mean Tajima’s \( D \) values for CTCF-motif sites. Tajima’s \( D \) values were calculated using 37 *D. melanogaster* North American strains’ polymorphism data for various groups of CTCF-motif sites, the synonymous and nonsynonymous sites of nearest genes, and randomly sampled 3’ UTR, 5’ UTR, and intergenic 9 bp sites. The center of each filled circle depicts the mean Tajima’s \( D \) value for each group, with the error bar indicating 2 standard deviations. **(C and D)** Estimated shared proportion of adaptation with neutral reference to nearest gene synonymous sites (C) and a set of small introns (D). *D. yakuba* sequences were used as an out-group for estimating alpha values for different groups of CTCF-motif sites using an extension of the MK test framework. The filled colored circles depict the shared alpha value estimated within each group, with the error bar indicating the 95% confidence interval. Label abbreviations: Syn, synonymous sites of nearest genes of CTCF binding sites; Nonsyn, non-synonymous sites of nearest genes of CTCF binding sites; TWOB, CTCF-motif sites associated with two-way orthologous binding events between *D. melanogaster* and the out-group; conserved TWOB, CTCF-motif sites associated with conserved two-way orthologous binding events; diverged TWOB, CTCF-motif sites associated with *D. melanogaster*-specific two-way orthologous binding events; FWOB binding, sites associated with four-way orthologous binding events; Young FWOB, sites associated with FWOBs, for which the age is estimated to be <2.5 Myr; old FWOB, sites associated with FWOBs, for which the age is estimated to be >6 Myr. doi:10.1371/journal.pbio.1001420.g003

Table S1C, Figure S18)
Figure 4A,B). Such correlation is also observed when using microarray data for inferring gene expression divergence (Figure S14) as well as when using high-sequence coverage sites (Figure S15). These observations indicate that CTCF binding evolution impacts gene expression evolution, which previously has been shown to evolve rapidly and to be shaped by selection in these species at the WPP stage [51,52].

Selection on gene expression can lead to adaptive evolutionary signatures in *cis*-regulatory elements. Indeed, in *Drosophila*, adaptive gene expression has been linked to adaptive *cis*-DNA evolution [53]. We thus hypothesized that the stronger positive selection signature observed in the diverged TWOBs might stem from the sites being associated with diverged expression that has more directly been subject to natural selection. We calculated and compared $\alpha$ values for two additional subgroups of TWOB sites: diverged TWOBs near genes with divergent expression and conserved TWOBs near genes with conserved expression. Consistent with our hypothesis, we observed a larger difference in $\alpha$ values between these two subgroups than between all conserved and diverged TWOBs (Figures S16 and S17).

CTCF Binding Evolution Is Correlated with the Origin of New Genes

CTCF binding sites in *Drosophila* have been associated with syntenic break points, consistent with their role in delineating the regulatory architecture of genes [13]. We wished to determine whether CTCF binding evolution correlates with any other genome structural evolution. New genes are defined as genes

![Figure 4. Functional consequences of CTCF binding evolution.](image-url)
recently originated in a clade, and they provide the opportunity to add new functions to a genome [34]. We found that among 42 young genes that are essential for *Drosophila melanogaster* survival [41], eight show the origin of new CTCF binding sites within 5 kb flanking regions. All eight show phylogenetic correspondence between the appearance of newly evolved CTCF binding sites and the appearance of the associated new gene. Seven new genes exhibit a new CTCF binding site near their 3’ end. For example, *Tfha-A*S2 (CG11839) [41] is a newly originated gene through gene duplication in the *melanogaster* subgroup, and a CTCF binding site is observed near its 3’ end in *D. melanogaster*, *D. simulans*, and *D. yakuba* but not in *D. pseudoobscura* (Figure 4C). A similar example is the gene *CheB93a* (CG15503) [41], which originated before the split of *melanogaster* subgroup from *D. ananassae* as a tandem duplicate of its parental gene *CheB93b* (CG31438) (Figure 4D). We also found a CTCF binding site near the 3’ end of the RNA gene *sphaus* (CR34154) [55], which originated in the *D. melanogaster* branch and is implicated in courtship behavior of male flies [56].

The association of newly evolved essential genes with newly evolved CTCF binding sites is highly significant compared to old essential genes with conserved CTCF binding sites (Kolmogorov-Smirnov test, p<1e-6; Figure S19).

How are new binding sites generated? Point mutations are observed in numerous case studies linking cis-regulatory sequence change to phenotypic consequences [3] and therefore are considered as the main source of binding site evolution in many theoretical works [57–59]. In a mammalian CTCF comparative study, Schmidt et al. presented compelling evidence that CTCF binding sites are driven by retrotransposon expansions, especially in the rodent lineages [9]. We investigated whether TEs might also be associated with CTCF binding site evolution in *Drosophila*. For all the CTCF binding sites identified in *D. melanogaster*, only slightly more than 1% (27 out of 2,267) overlap with annotated TEs [60]. This rate is extremely low compared to rodent species in which around 20% of CTCF binding sites are contained within SINE elements [9], indicating that it is not the primary mode of generating binding site diversity in *Drosophila*. However, for the *D. melanogaster* lineage-specific binding sites, approximately 6% (15 out of 261) of these sites overlap with a TE, resulting in a significant excess of new binding sites overlapping with TEs (Fisher’s exact test, p<0.0001; Table S18). Thus it is possible that a minority of newly arisen CTCF binding sites have resulted from TE insertions, but the majority of new binding sites are likely originating through mutation selection processes at the nucleotide level.

**Discussion**

Ever since King and Wilson proposed the importance of gene regulation for phenotypic variation [2], evolution of cis-regulatory elements has been under intensive investigation with an emphasis on enhancers [61,62] and transcription factor binding sites [4–8]. Insulator elements are a special class of cis-elements implicated in many fundamental biological processes including transcriptional regulation [14,15]. Despite their functional importance, the origin and evolution of insulator complexes remained largely uncharted [63,64]. Only very recently was the first comparative ChiP-seq study on CTCF in mammalian species published [9]. Here, we presented a formal evolutionary genetic analysis of CTCF-related insulator elements in multiple *Drosophila* species.

We found that CTCF binding is highly evolutionarily dynamic, with about 70% of binding events diverged between *D. melanogaster* and *D. pseudoobscura*. This high level of evolutionary divergence is consistent with a recent mammalian study, in which the CTCF binding conservation between human and mouse was estimated to be around 30% [9]. While in mammalian species, CTCF binding profiles are more conserved than tissue-specific transcription factors [7,9]; in *Drosophila* species, we observe higher binding divergence of CTCF than the developmental transcription factor Twist [9]. In fact, the high degree of binding divergence observed in liver-specific transcription factor CEBPA and HNF4A has led to a proposal of neutral drift underlying binding evolution [7]. However, the population genetic analysis of binding divergence of both the Twist data [8] and our CTCF data indicates that both purifying and positive selection are active forces in CTCF binding evolution. Although previous studies on *Drosophila* noncoding DNA [46,65] and DNA footprinting-derived TFBS sequences [66] have suggested the role of positive selection, here we present the first genome-wide evidence in support of positive selection using protein-binding-associated DNA mapped in vivo.

Our observation that young binding sites exhibit a signature of positive selection mimics the pattern observed with young genes [41], indicating that the origination of new binding sites is driven by positive selection. Further, the association between CTCF binding divergence and gene expression divergence indicates that change in CTCF binding has functional consequence. The fact that CTCF binding origination in multiple species coincided with new gene appearance also reinforces this functional view of binding change. The binding changes of this insulator protein may well result in regulatory rewiring through structurally redefining regulatory domains. We predict that this might be a universal mechanism in cis-regulatory evolution since CTCF protein is highly conserved across the metazoans [64]. Indeed, in mammalian species, lineage-specific CTCF binding sites are observed to demarcate both chromatin and gene expression domains [9]. Consistent also with the functional relevance of evolutionary changes in CTCF binding profiles, we observed that old and conserved CTCF binding sites are subject to stronger purifying selection and that expression levels of genes near these conserved sites are less likely to diverge. Together these observations indicate that functional constraints maintain conserved binding. This meshes well with the study on Twist [8], in which He et al. found that the most developmentally important genes in early embryo development have the most conserved Twist binding. In summary, we have provided evidence that the evolution of CTCF binding in *Drosophila* species is adaptive.

**Materials and Methods**

**Genomic Data Production**

The sequenced strains of *D. melanogaster*, *D. simulans*, *D. yakuba*, and *D. pseudoobscura* were maintained at room temperature (18–20°C). Whole animal white prepupa (WPP) for both ChiP-seq and expression profiling experiments were collected strictly within a 15-min time interval to ensure developmentally synchronized samples across species.

Triplicate CTCF ChiP-seq experiments in different species were carried out using a previously published [23] and verified CTCF antibody (Figure S1) according to the standard *Drosophila* modEncode ChiP protocol (www.modencode.org) and Illumina sequencing library preparation protocol. Illumina sequencing data were generated at the High-Throughput Genomic Analysis Core (HGAC) at the Institute for Genomics and Systems Biology.

RNA samples were isolated using Trizol, and the integrity of these samples were checked using an Agilent Bioanalyzer. Transcript levels of *D. melanogaster*, *D. simulans*, and *D. yakuba* WPP samples were measured by single-end mRNA-seq performed in triplicate. Additional sets of quadruplicate expression profiling
of *D. melanogaster* and *D. simulans* WPP samples were performed using custom-designed high-density 105K Agilent Gene Expression Arrays. All the genomic data are deposited at GEO under accession number GSE 24449.

**Peak Calling**

Sequence reads were mapped back to the genome with the ELAND algorithm using the following Flybase reference genome versions: *D. melanogaster* r5.3, *D. simulans* r1.2, *D. yakuba* r1.2, and *D. pseudoobscura* r2.2. Any reads with more than two mismatches or more than two “N”s were filtered out; only uniquely mapped reads were used in our later analyses. The raw data wiggle files were generated by counting the number of times each coordinate was sequenced. We used the Affymetrix Integrated Genome Browser (IGB) as well as the Broad Institute Integrative Genomics Viewer (IGV) [67] to visualize the raw data and generate snapshots at various genomic positions, as shown in Figure S2.

We modified the peak calling software QuEST [39] to incorporate triplicate data for CTCF binding peak identification. Briefly, QuEST (version 2.0) was used to generate the CDP (Compiled Density Profile) scores for each paired ChIP-input samples. We then normalized the CDP scores by multiplying corresponding ratio scores generated according to sequencing depth, and for each species, we calculated the mean CDP enrichment score (defined as the mean CDP score of ChIP samples minus mean CDP score of input samples) and mean CDP fold enrichment score (defined as the ratio of mean CDP of ChIP samples over mean CDP score of input samples). We performed a permutation-simulation procedure to empirically find the threshold values for the mean CDP enrichment score. We first permuted the experimental label (“ChIP” or “input”) of the CDP scores and then randomly sampled 10,000 coordinates to obtain their mean CDP enrichment score; we repeated the process 100 times and built a “Null” distribution for mean CDP enrichment scores. The 99th percentile of the positive values of the distribution is taken as our threshold for peak calling, which ensures an FDR<1%. We performed peak segmentation using the threshold in a way similar to TAS (Tiling Analysis Software, Affymetrix). We identified regions with at least 50 bp above the threshold and merged neighboring regions if the distance in between is less than 100 bp. We then filtered out peak regions for which the summit coordinate had a mean CDP fold enrichment score of less than 2. We have also calculated, for each identified peak region, the q value (Poison p value after multiple testing correction) associated with read number enrichment between ChIP and input samples for each species using raw read count data, and all q values<0.001. The summit peak coordinates of each identified peak regions were used to infer genomic positions of all the binding events. We designated a CTCF binding event/site as “intronic” or “exonic” if the summit coordinate is within boundaries of an annotated intron or exon, respectively. The remaining binding events/sites were then categorized into “promoter” or “intergenic” groups based on the distance of the peak summit coordinate to the nearest gene transcription start site (TSS); if the distance is <1 kb, we labeled it as “promoter”; otherwise, “intergenic.”

**Motif Analyses**

Species-specific motifs were de novo generated by running MEME [68] on binding site DNA sequences (i.e., 201 bp sequence surrounding the summit coordinate) using default parameters except for setting a motif length of 9 bp. We have used both the top 2,000 and total binding site sequences to run MEME and obtained similar species-specific motifs. FIMO [69] were used to search for motif occurrences in DNA sequences, and a Perl script was written to parse the FIMO result to get percentages of motif containing at various p value thresholds and to find the best motif and each individual motif in each binding site.

**Binding Divergence Analyses**

1. **Direct comparison of identified binding regions in each species.** We first mapped all the non-*D. melanogaster* species binding regions into the *D. melanogaster* genome using LiftOver [70], with all default parameters except a match of 0.3; then, we counted the percentage of *D. melanogaster* binding regions overlapping with each of the non-*D. melanogaster* LiftOver binding regions as pair-wise conservation rate. We also performed the reciprocal procedure by mapping *D. melanogaster* binding regions to each of the non-*D. melanogaster* genomes and estimated the pair-wise binding conservation rate as the percentage of each non-*D. melanogaster* binding region that overlaps with the LiftOver *D. melanogaster* binding regions.

2. **The *D. melanogaster*-centric quantitative analysis pipeline.** We developed a *D. melanogaster*-centric quantitative analysis pipeline to partition read count data variation within and between species and to directly identify conserved and diverged peaks between each pair of the non-*D. melanogaster* species and *D. melanogaster* (*D. simulans*, *D. yakuba*, and *D. pseudoobscura*) species into *D. melanogaster* genome using LiftOver [70] with all default parameters, except a match of 0.5. We then generated CDP scores using the LiftOver sequence reads for each non-*D. melanogaster* species using QuEST (version 2.0). For each species pair considered, the CDP scores from each sample were then normalized by multiplying a normalizing value calculated as follows:

\[ \text{No. of LiftOver reads} \times \frac{D. melanogaster \text{ genome size}}{Between \text{ species alignable size}} \times 10^8 \]

An ANOVA-like linear categorical model described below was then applied to the normalized CDP scores at each coordinate to obtain species-specific ChIP enrichment score estimates, difference of species-specific ChIP enrichment score estimates (the interaction term), and their associated p values.

\[ Y = B_0 \times \text{Experiment} + B_1 \times \text{Species} + B_2 \times (\text{Experiment} \times \text{Species}) + \varepsilon \]

In the model, Y is defined as the observed sequence data, by inputting the normalized CDP scores transformed from sequence read count data of the two species at a specific genomic coordinate. Experiment here is a categorical variable (dummy variable) indicating the CDP score source as ChIP or input; Species is the other categorical variable, indicating the species source of the CDP score (either *D. melanogaster* or non-*D. melanogaster*); Experiment × Species is the interaction term between experiment types and species types. B0, B1, and B2 are the associated coefficients with the variables to be estimated, and ε is the residual error term. We then smoothed these tracks of ChIP enrichment score estimates as well as the −log10 transformed p values for each chromosome by averaging 100 bp sliding windows with each step moving 1 bp. We first identified candidate regions of ChIP enrichments in both species. We then directly identified *D.
melanogaster-specific binding sites, non–D. melanogaster-specific binding sites, and shared binding sites. For a binding peak to be identified as shared, it must satisfy the following two requirements: (1) the ChIP enrichment scores in both species are above the threshold and (2) the p values associated with ChIP input comparison are significant (−log10 transformed p value >20).

We took a conservative approach in identifying diverging binding sites. For a binding peak to be identified as D. melanogaster–specific, it must satisfy the following three conditions: (1) the ChIP enrichment scores in D. melanogaster are above the chosen threshold (see below and Table S5), but in the non–D. melanogaster species, the score must be below the threshold; (2) the p value associated with the ChIP/input comparison must be significant; and (3) the p value associated with the species-specific ChIP effect must be significant. We similarly identified the non–D. melanogaster–specific binding sites. We have used a set of different thresholds (0.35, 0.4, 0.5, and 1.0; also see Tables S5 and S6) to identify the diverged and conserved binding sites. The results presented in the main text are based on the threshold 0.4, which is empirically determined in the permutation-simulation procedure for D. melanogaster data as described in the section “Peak Calling.” The reason to use this threshold are (1) it is empirically determined and (2) all the non–D. melanogaster data have been translated and normalized to be comparable to the D. melanogaster data. We applied the other thresholds to test the robustness of observed patterns under looser criteria (0.35) and more stringent criteria (0.5 and 1.0).

Using data simulation, we estimated the False Positive Rate (the rate of identifying conserved binding sites as diverged) of the linear model as 0.35%. The simulation was performed by pooling all the D. melanogaster ChIP sample sequence reads together and randomly sampling the same number of reads for each ChIP replicate from the pool to build a simulated ChIP sequence read data set. Similarly, we obtained a simulated input data set. We performed our analysis pipeline with the D. melanogaster input data and the simulated data. Ideally, we would identify all the binding sites as shared between D. melanogaster and the simulated data but found 0.35% of them are identified as diverged.

As an alternative method, we also estimated the overall error rate (False Positive Rate plus False Negative Rate) for misidentifying the pair-wise evolutionary status of D. melanogaster binding sites of the whole analysis pipeline as <2% using a set of 100 randomly sampled CTCF binding sites that were manually curated as a “gold standard.” Briefly, we curated 100 random D. melanogaster binding peaks by manually inspecting the raw data wiggle file. We then looked at the corresponding orthologous regions as well as 2 kb flanking region of the orthologous sequences in each non–D. melanogaster species. If we identified any peak using this method, we defined the D. melanogaster binding peak as shared, and otherwise, not shared. The percentage of discrepancy between human eye curation of raw count data and our analysis pipeline are taken as the overall error rate. From the pipeline with the empirical thresholds, we identified 2,267 binding sites for D. melanogaster, which shows >95% overlap with the binding sites identified previously using triplicate data. We estimated pair-wise binding divergence as the percentage of D. melanogaster binding sites that is not shared with the other species.

### 3. The He et al. method.

We followed the method as described in [8]. Briefly, we randomly picked two out of our three replicates for each species to match the structure of analysis He et al. performed. Since there are more input reads than ChIP reads in our data, we performed random sampling of input reads to match the number of reads in the paired ChIP samples. This is important to gain an accurate estimate of FDR with software MACS [71]. We then applied MACS (version 3.2) to identify binding peaks with the D. melanogaster sequence read data as well as the non–D. melanogaster LiftOver sequence read data. We took the set of binding peaks with a p value <10–21.8 (same p value as He et al. used) in one D. melanogaster replicate as the reference binding sites and compared it to all binding sites identified in other species with p value cutoff 10–2. In order to assess the False Negative Rate, we have also generated two pseudo-ChIP replicates by randomly sampling input sequence reads and performed the same procedure as for other species data.

### Population Genetics Analyses

We downloaded the pre-assembled genome sequences of 37 North American RAL lines from the Drosophila Population Genome Project (www.dpgp.org; Release 1, 30 genome) and filtered out any nucleotide with Phred score <30 as “N.” Combining that data and the D. melanogaster reference genome sequence, we generated the polymorphism data for various sets of genomic sites. We included two different types of CTCF-related genomic sites in our analyses: CTCF-201 bp sites and CTCF-motif sites. The CTCF-201 bp sites comprised all the 201 bp flanking sequences centered at the D. melanogaster peak coordinate identified in our linear categorical model. The CTCF-motif sites comprised all 9 bp motif sequences found by FIMO at a p value = 0.01 within each CTCF-201 bp site concatenated together. The different genomic reference sites were generated by random simulation. For neutral reference, we used the synonymous sites of the nearest genes to the binding sites as well as a set of small intron sequences. The small intron sequences are the 8th–30th nucleotides of introns <65 bp as described in [49], and any of these introns overlapping with known EST were filtered out.

We calculated Tajima’s D values [44] for different sets of noncoding sites using DnaSP 5.0 batch mode [72] and used Polymorphorama [65] for synonymous and nonsynonymous sites of the nearest genes.

For ω estimation, corresponding orthologous DNA sequences in out-group species D. simulans and D. yakuba were used. Orthologous coding sequences of genes were obtained according to the Flybase (www.flybase.org) annotation. Orthologous sequences for noncoding sites were generated using UCSC pair-wise genome alignment [73]. Sequence alignments were performed with ClustalW2 [74]. The number of polymorphic and divergent sites for noncoding sequences as well as synonymous sites of nearest genes [45,46] were obtained using a Perl script implementing the PopGen module of BioPerl, which yielded the same result as DnaSP5.0. By taking binding-associated DNA as “nonsynonymous” sites, we estimated the shared ω with a 95% confidence interval using DoFE 2.0 [47].

We used the multilocus HKA test [50] C code implemented by Jude Hey (http://genfaculty.rutgers.edu/hey/software#HKA) to perform the HKA tests for the following three comparisons: (1) young CTCF-201 sites versus old CTCF-201 sites; (2) young CTCF-201 sites versus neutral small intron sites; and (3) old CTCF-201 sites versus neutral small intron sites. The sum of deviations is calculated by summing up across all loci, and the p values are obtained from 1,000 times of coalescence simulations.

### Expression Data Analyses

For microarray data, probe intensities were extracted using Feature Extraction Software (Agilent). All arrays passed the manufacture’s QC and our additional QC processes, with high linear correlation between probe intensities and actual concentration of Spike-in RNAs (linear regression slope ≈1 and R-squared >0.95) and high correlation between duplicated probes (Pearson’s correlation r >0.98). Any probes flagged by FE were treated as...
missing data. Background subtraction (“normexp” function in “marray” package), log2 transformation, and quantile normalization were performed for each species-specific array set using Bioconductor packages. We took the advantage of the fixed amount of starting Spike-In RNA species in our sample prep experiments and performed the between-species normalization as follows: for each species, we regressed log2 transformed expression measurements of Spike-In probes to the log2 transformed actual RNA concentrations to obtain a regression line; we then subtracted the value of y-axis intercept from each probes. We pooled all probes for each pair of orthology and applied a linear mixture model as follows and categorized the expression level of the gene as “diverged” or “stable” according to the \( p \) value associated with the estimated between-species expression difference. Correction for multiple testing was performed in a FDR approach [75,76] using \( R \) package “qvalue.”

\[
y = B_2 S + B_R (|P|) + \varepsilon
\]

We input \( Y \) as the normalized log2 microarray intensity measurements of a given pair of orthologs between species; \( S \) here is a categorical variable indicating the different species (\( D. melanogaster \) or \( D. simulans \)); \( P \) here is a numerical variable indicating the number of different probes for the genes in microarray design; \( \{ |P| \} \) here indicates the random effects of different probes. \( B_2 \) and \( B_R \) are the coefficients to be estimated. \( \varepsilon \) is the residual error term.

**RNA-seq Data Analyses**

For RNA-seq data, we used Bowtie [77] to map the Illumina sequence reads to the genome as well as the annotated exon–exon conjunctions. The number of mapped reads for each gene in different species is counted the same way as described in [78]. Reproducibility between replicates was assessed by calculating the Spearman’s rank correlation coefficient of RPM (reads per million) values. For each species pair, we pooled the read count data for orthologous genes together, performed upper-quantile normalization [79], and filtered out genes with <5 reads mapped as “NA.” The genes with divergent expression between species were then called through a generalized linear model framework as described in [80] with a multiple testing corrected \( p \) value <0.01 and a log2 fold between species difference >2.

**Association Between Essential Genes and CTCF Binding Sites**

We used a set of 42 \( D. melanogaster \) new essential genes (genes originated in \( Drosophila \) within 25 Myr) as described in [41] and a set of 2,003 old essential genes (genes originated more than 40 Myr ago) for our analysis. The list of old essential genes is a union of two sets: first, a set of 86 old essential genes identified in an RNAi screen as in [41]; second, a set of 1,948 genes with lethal allele phenotypes reported in previous mutagenesis screen studies obtained from the \( Drosophila \) Interaction Database (DroID) [81]. For the new essential gene set, we calculated the proportion of genes that have phylogenetically congruent CTCF binding sites within flanking regions of different length. A CTCF binding site is described as phylogenetically congruent to a gene if and only if the binding site distributes in the exactly same branches on the phylogeny as the gene. For the old essential gene set, we performed 1,000 times of random sampling; each time we randomly picked 42 genes and performed the same procedure as described for the new essential gene and calculated the mean proportion of old genes that have phylogenetically congruent CTCF binding sites within various flanking regions.

**Overlapping with TEs**

We downloaded the annotated TEs in \( D. melanogaster \) from Flybase and calculated the overlap between the 201 bp flanking region of each group of \( D. melanogaster \) binding sites with the annotated TEs.

**Supporting Information**

**Figure S1** Verification of antibody. (A) Alignment of CTCF protein C terminus sequences in the four species. The CTCF_C rabbit antibody used in this article was generated using the \( D. melanogaster \) CTCF protein C terminus sequence as antigen. The C terminus parts of CTCF protein are identical in the \( D. melanogaster \) subgroup species. There are three amino acid changes between \( D. melanogaster \) and \( D. pseudoobscura \), two of them are similar amino acid changes (in blue) and only one is a different amino acid change (in red); (B) Western Blot of CTCF-C antibody used for the ChIP-seq experiments with \( D. pseudoobscura \) white pre-pupa extracts at two different volumes. The size of detected band is consistent with the predicted 91.31 kD molecular weight for \( D. pseudoobscura \) CTCF protein. (PDF)

**Figure S2** CTCF binding profiles at the Bithorax complex region in \( D. melanogaster \) genome. Previous reported canonical CTCF binding sites in the Bithorax complex region are recapitulated in every biological replicate in our ChIP-seq data. From top to bottom, the heights of the wiggle files denote the absolute values of raw data sequence depth for every 10 bp bin calculated using only the uniquely mapped Solexa reads for each of the three ChIP samples—\( D. mel \) ChIP1, \( D. mel \) ChIP2, and \( D. mel \) ChIP3—and their corresponding reference samples—\( D. mel \) input1, \( D. mel \) input2, and \( D. mel \) input3. The seventh panel shows the boundaries of previously identified insulator elements (in sky blue) in this region. (PDF)

**Figure S3** Illustration of the modified QuEST peak calling procedure. (PDF)

**Figure S4** Spearman’s rank correlation between ChIP-seq replicates. The Spearman’s correlation coefficients (rank order correlation) were calculated with CDP scores (compiled density profile, a QuEST transformation of the sequence depth data for peak calling) in the 500 bp flanking region around the peak summit coordinate for each individual binding peak between any two replicates. The box plots show the overall distribution of Spearman’s correlation coefficients for summarized overall combinations of replicate pairs identified in (A) \( D. melanogaster \), (B) \( D. simulans \), (C) \( D. yakuba \), and (D) \( D. pseudoobscura \). (PDF)

**Figure S5** Sequence alignments of CTCF protein DNA binding domain (DBD) in the four species. The 11 yellow colored blocks represent the 11 predicted C2H2 zinc finger domains using online domain finding software provided by the Pfam database. Different color depicts different types of amino acids compared to the consensus ones: amino acids that are identical to the consensus (in black); amino acids that are different but with similar properties to the consensus (in blue); and amino acids that are different and have different properties from the consensus (in red). (PDF)

**Figure S6** Illustration of the \( D. melanogaster \)-centric quantitative analysis pipeline. (PDF)
Figure S7  Evolutionary groups of CTCF binding events in D. melanogaster group. Top panel, representative dynamic binding profiles across the three D. melanogaster group species illustrating examples of the seven mutually exclusive binding statuses. The heights of the binding curve denote the ChiP CDP enrichment score estimated from our analysis pipeline (Figure S6). The y-axes in the three binding curves for each evolutionary group are at the same scale. In the lower table, the first row contains the Boolean conservation score for each evolutionary status, where 1 depicts the existence of the binding event and 0 depicts the absence of binding event; second and third rows, number of binding events falling into each evolutionary group for all possible binding events and FWOBs (four-way orthologous binding).

Figure S8  Sequence conservation of CTCF binding sites. (A) Distributions of median PhastCons scores for CTCF binding sites. The box plots show the distribution of median PhastCons scores for the conserved and diverged 201 bp sites summarized over all three pair-wise comparisons. (B) Percentage of sequence identity for CTCF binding sites. The box plots show the distribution of percentages of sequence identity in the TWOB 201 bp sites summarized over all three pair-wise comparisons. The percentages of sequence identity are calculated using the pair-wise sequence alignments of the 201 bp flanking sequences of the summit coordinates. 

Figure S9  Mean Tajima’s D for CTCF-201 sites. Mean Tajima’s D values were calculated using 37 D. melanogaster North American strains’ polymorphism data for various groups of CTCF-201 sites. The center of each circle depicts the mean value, with the error bar indicating 2 standard deviations. The out-group species used here is D. simulans. Label abbreviations: Syn/Nonsyn, synonymous/non-synonymous site of the nearest gene; inter, randomly sampled 201 bp intergenic regions; 3’UTR, randomly sampled 201 bp 3’UTR regions; 5’UTR, randomly sampled 201 bp 5’UTR regions; TWOB, CTCF-201 bp sites associated with two-way orthologous binding events between D. melanogaster and the out-group; conserved TWOB, sites associated with conserved two-way orthologous binding; diverged TWOB, sites associated with diverged two-way orthologous binding; FWOB, sites associated with four-way orthologous binding; Young FWOB, sites associated with those FWOB with inferred evolutionary age <2.5 Myr; Old FWOB, sites associated with those FWOB with inferred evolutionary age >6 Myr.

Figure S10  Mean Tajima’s D for CTCF-motif and CTCF-201 high-sequence coverage sites. Mean Tajima’s D values for different groups of (A) CTCF-motif and (B) CTCF-201 sites after filtering out sites with input sequence coverage <0.5. The center of each circle depicts the mean value, with error bars indicating 2 standard deviations. The out-group species used here is D. simulans. Label abbreviations are the same as for Figure S9.

Figure S11  Shared proportion of adaptation in CTCF-201 bp sites. Shared χ values estimated for various groups of CTCF-201 bp sites through the extended MK test framework, with (A) D. simulans and (B) D. yakuba as out-group species. The center of each circle in the plot depicts the χ value estimated, with error bars indicating the 95% confidence interval. The label abbreviations are the same as for Figure S10.

Figure S12  Shared proportion of adaptation in CTCF-motif sites. Shared χ values estimated for various groups of CTCF-motif sites through the extended MK test framework with D. simulans as an out-group species. The center of each circle in the plot depicts the χ value estimated, with error bars indicating the 95% confidence interval. The label abbreviations are the same as for Figure 3.

Figure S13  Shared proportion of adaptation in CTCF-motif high-sequence coverage sites. Shared χ values estimated for various groups of CTCF-motif sites after filtering out sites with input sequence coverage <0.5 through the extended MK test framework, with D. yakuba as the out-group species. The center of each circle in the plot depicts the χ value estimated, with error bars indicating the 95% confidence interval. The label abbreviations are the same as for Figure 3.

Figure S14  CTCF binding evolution associated with gene expression evolution inferred from microarray data. The bar plots show the proportion of genes with diverged expression between D. melanogaster/D. simulans associated with different groups of CTCF binding sites: Genome-wide (black), Conserved TWOB (pink), Diverged TWOB (green), Old FWOB (orange), and Young FWOB (light purple). The table below each bar plot shows the number of genes with diverged and conserved gene expression in the corresponding comparisons and associated with the corresponding CTCF binding sites. For each groups of CTCF binding sites, the associated genes are the union of the nearest gene to each binding site. The evolutionary status of gene expression (conserved or diverged) is determined using quadruplicate expression profiling with custom-designed species-specific Agilent 105K microarrays. The label abbreviations are the same as for Figure S15. Significance levels: * p<0.05; **p<0.01, one-sided Fisher’s exact test.

Figure S15  CTCF binding evolution associated with gene expression evolution inferred from RNA-seq data at high-sequence coverage sites. The bar plots show the proportion of genes with diverged expression between (A) D. melanogaster/D. simulans and (B) D. melanogaster/D. yakuba comparisons associated with different groups of CTCF binding sites after filtering out sites with input coverage <0.5. All labels are the same as in Figure 4A and 4B.

Figure S16  Shared proportion of adaptation in different groups of CTCF TWOB sites. Shared χ values estimated for various groups of (A) CTCF-motif sites and (B) CTCF-201 sites through the extended MK test framework using D. yakuba as an out-group. The center of each circle in the plot depicts the χ value estimated, with error bars indicating the 95% confidence interval. The label abbreviations: TWOB, Two-Way Orthologous Binding sites identified between D. melanogaster and the out-group species; diverged TWOB, diverged Two-Way Orthologous Binding sites; conserved TWOB, conserved Two-Way Orthologous Binding sites; conserved TWOB with conserved expression, the subset of conserved Two-Way Orthologous Binding sites for which the expression level of their nearest gene are evolutionarily conserved; diverged TWOB with diverged expression, the subset of diverged Two-Way Orthologous Binding sites for which the expression level of their nearest gene are evolutionarily diverged.

Figure S17  Shared proportion of adaptation in different groups of CTCF TWOB high-sequence coverage sites. Shared χ values estimated for various groups of (A) CTCF-motif sites and (B) CTCF-201 sites through the extended MK test framework using D. yakuba as an out-group. The center of each circle in the plot depicts the χ value estimated, with error bars indicating the 95% confidence interval. The label abbreviations are the same as for Figure S16.
D. yakuba as an out-group. The sites used here for \( \alpha \) estimation are those sites with input sequence coverage \( >0.5 \). All labels and abbreviations are the same as in Figure S16.

**Figure S18** Reproducibility of RNA-seq data. The scatter plot shows the high correlation between gene RPM (number of reads per million) values from two *D. melanogaster* WPP biological samples. The estimated Spearman’s rank order correlation is 0.96.

**Figure S19** Association between essential genes and CTCF binding events. The red dotted line shows the cumulative proportions of 42 new essential genes (originated less than 25 Myr ago) with phylogenetically congruent CTCF binding sites within flanking regions of various lengths. The blue dotted line shows the cumulative average proportions of randomly sampled 42 old essential genes (originated more than 40 Myr ago) with phylogenetically congruent CTCF binding sites obtained from 1,000 simulations. A CTCF binding site is described as phylogenetically congruent to a gene if and only if the binding event appears in the exactly same branches as the gene on the evolutionary tree. The difference between the two cumulative lines is significant, \( p<1e-6 \), Komogorov Smirnov test.

**Figure S20** Selection signatures in Twist-201 bp sites. (A) Mean Tajima’s \( D \) values for Twist-201 bp sites. The center of each circle depicts the mean value, with error bars indicating 2 standard deviations. The out-group species used here is *D. yakuba*. (B) Shared proportion of adaptation (alpha) estimated for Twist-201 bp sites using *D. yakuba* as the out-group. The center of each circle in the plot depicts the \( \alpha \) value estimated, with error bars indicating the 95% confidence interval. The mean Tajima’s \( D \) values as well as alpha values for Twist-201 bp sites are plotted together with CTCF-201 bp sites (as labeled in the figure). TWOB, diverged TWOB, and conserved TWOB for Twist and for CTCF are defined the same way as in Figure 3. The TWOB, diverged TWOB, and conserved TWOB Twist binding sites were identified by applying our analysis method to the Twist comparative data.

**Table S1** Summary of sequence reads. (A) Number of Solexa sequence reads for ChIP-seq experiments. (B) Number of LiftOver reads of non-*D. melanogaster* species. (C) Number of Solexa sequencing reads for RNA-seq experiments.

**Table S2** Ka/Ks ratio for CTCF gene in *Drosophila* species.

**Table S3** CTCF binding site motif enrichment in each species.

**Table S4** Pearson’s correlation coefficients between ChIP replicates.

**Table S5** Diverged and conserved CTCF binding events.

**Table S6** Diverged and conserved CTCF binding events at high-sequence coverage sites.

**Table S7** CTCF binding divergence estimated by direct comparison.

**Table S8** CTCF binding divergence estimated using He et al. method.

**Table S9** Twist binding divergence estimated using our pipeline.

**Table S10** Summary of Twist binding divergence estimated using different methods.

**Table S11** Genomic distribution of different evolutionary groups of CTCF binding events.

**Table S12** Parsimonious age dating of *D. melanogaster* CTCF binding events.

**Table S13** Evolutionary groups of CTCF binding events at high-sequence coverage sites.

**Table S14** Number of fixed and polymorphic mutations in CTCF-associated DNA sequences.

**Table S15** Number of fixed and polymorphic mutations in CTCF-associated DNA sequences at high-sequence coverage sites.

**Table S16** HKA test for old and young sites.

**Table S17** HKA test for old and young high-sequence coverage sites.

**Table S18** Overlapping of CTCF binding sites with TE.

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**Author Contributions**

The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: KPW XN NN. Performed the experiments: XN. Analyzed the data: XN YEZ KPW NN SC ML. Contributed reagents/materials/analysis tools: XN YEZ KPW NN SC ML. Wrote the paper: XN KPW.

**References**


