Sequence analysis

Identification of transcription factor binding sites from ChIP-seq data at high-resolution

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ABSTRACT
Motivation: Chromatin-immunoprecipitation coupled to next-generation sequencing (ChIP-seq) is widely used to study the in vivo binding sites of transcription factors (TFs) and their regulatory targets. Recent improvements to ChIP-seq, such as increased resolution, promise deeper insights into transcriptional regulation, yet require novel computational tools to fully leverage their advantages.

Results: To this aim, we have developed peakzilla, which can identify closely spaced TF binding sites at high resolution (i.e. resolves individual binding sites even if spaced closely), as we demonstrate using semi-synthetic datasets, performing ChIP-seq for the TF Twist in Drosophila embryos with different experimental fragment sizes, and analyzing ChIP-exo datasets. We show that the increased resolution reached by peakzilla is highly relevant as closely spaced Twist binding sites are strongly enriched in transcriptional enhancers, suggesting a signature to discriminate functional from abundant non-functional or neutral TF binding. Peakzilla is easy to use as it estimates all the necessary parameters from the data and is freely available.

Availability: The peakzilla program is available from https://github.com/steinmann/peakzilla or http://www.starklab.org/data/peakzilla/.
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Supplementary information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION
Gene expression is mainly regulated at the transcriptional level and achieved through the binding of transcription factors (TFs) to genomic regulatory regions called promoters and enhancers. Chromatin immunoprecipitation followed by sequencing (ChIP-seq) is extensively used to determine transcription factor binding sites (TFBSs) genome-wide (Johnson et al., 2007; Robertson et al., 2007). Compared to ChIP-chip (Ren et al., 2000; Iyer et al., 2001), ChIP-seq has dramatically improved the resolution of the identified TFBSs (from hundreds to only tens of nucleotides). A recent refinement of ChIP-seq, the ChIP-exo method (Rhee and Pugh, 2011), further increases the resolution of ChIP-seq experiments, theoretically to single nucleotides.

The specific features and strategies of TF ChIP-seq data analysis have been well described by several reviews (e.g. Pepke et al., 2009) and protocols (e.g. Bardet et al., 2012). Briefly, as typically only one of both ends of the immunoprecipitated DNA fragments is sequenced, the sequencing reads (or tags) result in a bimodal distribution that is characteristic for true TFBSs (Figure 1a). This distribution is typically used to estimate the average size of the fragments, which subsequently allows the prediction of TFBSs across the genome.

Many computational tools have been developed and are successfully used to predict such binding events (Wilbanks and Facciotti, 2010). However, in our experience, these tools are not optimized to take advantage of recent methodological improvements, which comprise paired-end sequencing, high sequencing depth (e.g. on Illumina HiSeq systems), and - most importantly - an increase in experimental resolution, both of conventional ChIP-seq and of ChIP-exo. Available tools typically merge closely spaced read-density peaks into large regions, which is preferable when analyzing certain chromatin features that mark extended regions (e.g. histone modifications), but means that the ability to distinguish (i.e. resolve) individual closely spaced TFBSs is lost. High resolution and precision (i.e. the correct prediction of the TFBSs’ exact locations as measured for example as the distance from the inferred TFBS [the reported peak summit] to the TF’s sequence motif) are crucial when determining individual TFBSs (e.g. Guo et al., 2012), as promoter and enhancer regions often consist of multiple TFBSs for the same TF (homotypic TFBSs clusters) (Lifanov et al., 2003; Gotea et al., 2010; He et al., 2011) or different TFs (Berman et al., 2002; Schroeder et al., 2004).

To this end, we developed a new computational tool, peakzilla, which fully exploits the bimodal distribution of
sequence reads characteristic of true TF binding events, to identify closely adjacent TF binding sites with high resolution and precision. Peakzilla is not meant for the identification of broad enriched regions (e.g. histone marks) for which we recommend using MACS or similar programs.

We evaluate peakzilla by comparing it to three widely used peakfinders of the first generation such as MACS (Zhang et al., 2008), QuEST (Valouev et al., 2008), and cisGenome (Ji et al., 2008), as well as four methods methods developed more recently for the detection of high resolution peaks such as spp (Kharchenko et al., 2008), SISSRs (Jothi et al., 2008), GPS (Guo et al., 2010) and PeakRanger (Feng et al., 2011). Peakzilla shows superior resolution and precision on conventional ChIP-seq datasets from S. Cerevisiae (Zheng et al., 2010), C. elegans (Zhong et al., 2010), D. melanogaster (He et al., 2011), mouse (Schmidt et al., 2010), and human (Kasowski et al., 2010; Cuddapah et al., 2009) and on recent ChIP-exo datasets (Rhee and Pugh, 2011). We also specifically test the resolution limits of each method using semi-synthetic ChIP-seq datasets and show experimentally that peakzilla fully reflects increased experimental resolution by performing ChIP-seq experiments for Twist in D. melanogaster at normal, medium, and high resolution. These results suggest that peakzilla is best suited for the identification of TFBSs with recent ChIP methods.

2 PEAKZILLA ALGORITHM

Peakzilla uses the bimodal distribution of the reads (Figure 1a) not only to estimate the fragment length but also to weight the read counts during peak calling and to score the candidate TFBSs. This has two important advantages: first, it enables peakzilla to more clearly discriminate between reads from adjacent TFBSs, leading to a substantial increase in resolution compared to treating reads irrespective of their directionality. Second, it avoids false positives that originate from artifacts during library preparation or sequencing, without the need to collapse or down-weight reads that map to identical genomic positions (Figure 1b and Supplementary Figure 1). This is especially important when working with high sequence coverage, as obtained for small genomes (e.g. yeast, flies, or nematodes) and with modern NGS sequencers (Chen et al., 2012). Finally, when using peakzilla on paired-end ChIP-seq data, the estimated fragment size is directly averaged from the mapped reads.

Peakzilla first scans the genome for candidate TFBSs that show high coverage in sequencing reads of the IP sample compared to the control sample (note that the control sample is optional, allowing peakzilla to be used with ChIP-exo). It then scores the candidates by the normalized read count of IP sample minus the control sample if available. To discriminate between artifacts and true TFBSs with recent ChIP methods.
binding events in enriched regions, each candidate TFBS score is further weighted by a distribution score that estimates how well the observed distribution of the reads in the peak region fits to a model for the bimodal read distribution (Figure 1c & d). Indeed, further analyses suggest that candidates which are penalized using this distribution score are likely false since they contain substantially less diverse sequence reads (Figure 1e and Supplementary Figure 6), i.e. their high read densities stem from only a few highly duplicated sequences, which are likely amplification artifacts, and are significantly less enriched in the corresponding TF motif (Figure 1f) and. The method is illustrated in the method section and in Supplementary Figure 3.

3 RESULTS

3.1 Overall performance

To evaluate peakzilla, we compared it to other methods using diverse ChIP-seq datasets from S. cerevisiae, C. elegans, D. melanogaster, mouse, and human. Although the number of peaks called by the different methods is highly dependent on the thresholds and parameters used, the respective genomic regions overlap very well (Supplementary Figure 4), demonstrating the maturity of available tools for ‘peak calling’. For example, all known Twist enhancers are identified by all methods, except for four and three enhancers that are not found by QuEST and GPS, respectively (Supplementary Figure 5). For most TFs tested, of all peaks identified by only one of the methods, those found exclusively by peakzilla are significantly more highly enriched in TF motif occurrences (Figure 2a and Supplementary Figure 6). They also have higher fold enrichments of ChIP over input than peaks found exclusively by any of the other methods (Figure 2b and Supplementary Figure 6).

3.2 High precision of peakzilla peaks

When identifying TFBSs at high resolution, the correct prediction of the precise TFBSs’ location is important and critical for subsequent analyses of sequence features of TF binding. We found that most methods, including peakzilla, place the summits of the peak regions very closely to the nearest motif occurrence (Supplementary Figure 7, which also shows that the fraction of peaks that contain a motif is comparable across methods), arguing that the high resolution of peakzilla’s peak (see below) does not come at the expense of precision.

3.3 Multiple peak regions function as transcriptional enhancers

The main strength of peakzilla is its ability to find peaks at high resolution. As the locations of sequencing reads that originate from a single TFBS are limited by the fragment size used for library preparation, we adjust our search window for counting and scoring sequence reads accordingly, and report peak regions as the average fragment size centered on the summit position. This is different from the large peak regions reported by MACS and to a lesser extent QuEST, CisGenome and PeakRanger (and from reporting only the summit positions as do SISSRs and spp; Figure 3a), and is one of the key features that allow peakzilla to resolve closely spaced peaks up to distances that correspond to half the fragment size. This is the highest resolution that can easily be obtained without losing the ability to uniquely assign reads to individual TFBSs. A further gain in resolution would require the deconvolution of overlapping read-distributions by model fitting, a computationally intensive approach used e.g. by GPS (Guo et al., 2010). Peakzilla splits a substantial amount of MACS peaks (e.g. 22% for Twist) into several peaks, each constituting a putative TFBS (Supplementary Figure 8). Indeed, as expected for independent TFBSs, both the split peaks that correspond to the MACS summits (major peaks) and the minor peaks were significantly enriched for the Twist motif. Thus, many split MACS peaks may represent homotypic clusters of Twist binding sites. In addition, minor peaks frequently contained motifs for the TFs Snail and Dorsal, which are known to cooperate with Twist and might have been co-precipitated after cross-linking (Figure 3b).

Most importantly, MACS peaks split by peakzilla appear to be more often functional than MACS peaks that are not split (one-to-one peaks) and control regions (Figure 3c): TF binding to split peaks is more highly conserved in other Drosophila species (He et al., 2011) and they are significantly more enriched for known
Fig. 3. Functionality of multiple peak regions. Analyses performed on the Twist dataset in *D. melanogaster*. (A) Example of peak split. Peakzilla detects three adjacent peaks, while MACS, QuEST, CisGenome and PeakRanger report a single large peak region, and SISSRs and spp report two peak regions (GPS did not call any peak in that region; we considered all peaks called with standard parameters for each method) (B) Split peaks match motif occurrences. All peakzilla peaks corresponding to a single MACS peak (major: same summit; minor: additional summit) are more highly enriched in Twist motifs than control regions, suggesting that they constitute true independent TFBSs. The same is true for motifs of Snail and Dorsal, which are transcription factors known to cooperate with Twist. (C) Split peaks are highly conserved. (D) Split peaks are enriched for known enhancers. (E) Split peaks are enriched for mesodermal enhancers.

3.4 High resolution of peakzilla

The evaluation of TFBS predicted from ChIP-seq data (i.e. peaks) is complicated by the fact that a ground truth typically does not exist for experimental ChIP-seq datasets. This is especially true for the evaluation of peak calls at different resolutions that predict a single TFBS versus several closely spaced TFBSs. Although the occurrence of TF motifs is a good proxy for independent binding events, we sought to demonstrate the validity of peak splitting more directly by generating semi-synthetic datasets with defined peak-to-peak distances. We found that peakzilla is still able to separate all peak pairs at 150bp without calling any false positives, which is the best resolution compared to all other methods (Figure 4).

Fig. 4. High resolution of peakzilla. We evaluated the different methods on semi-synthetic datasets that contained peak pairs at decreasing peak-to-peak distances (i.e. resolution). For each method, we determined a true positive rate (TPR; number of correct peak calls divided by the total number of true peaks) and false discovery rate (FDR; number of false peak calls divided by the number of total peak calls) and indicate the best resolution reached (in base pairs [bp] below each method’s name).
3.5 Peakzilla leverages increased resolution of ChIP experiments

To directly demonstrate peakzilla’s ability to make use of increased experimental ChIP resolution, we performed conventional ChIP-seq for Twist from Drosophila embryos with increasingly smaller fragment sizes. For this, the chromatin was sonicated into relatively small DNA fragments and then further trimmed by DNase I digestion before ChIP (see Methods). This yielded three ChIP datasets with estimated fragment sizes of 102bp, 72bp, and 49bp, from which we called peaks with the different peak finders (Figure 5a). We expected that with decreasing fragment sizes, the width of the identified peaks should decrease and the resolution, i.e. the ability to resolve closely spaced binding sites, should increase. Indeed, the peak regions reported by peakzilla, but not those reported by most other methods, showed decreased width with decreasing fragment sizes (Figure 5a). More importantly, the resolution, as measured by the minimal peak-to-peak distance (after removing 1% outliers), increased with decreasing fragment sizes for peakzilla, but not for other methods (Figure 5b). SISSRs, GPS, and peakzilla performed well on the small-fragment sample, with peakzilla reaching the highest resolution of all methods. These results demonstrate that the maximum benefit of experimental methods with higher resolution can only be obtained when used together with high-resolution computational methods such as peakzilla.

3.6 Peakzilla as a peak caller for ChIP-exo data

The recently developed ChIP-exo method adds a lambda exonuclease digestion step after ChIP, which trims the 5’ DNA strand until the cross-linked TFBS (Rhee and Pugh, 2011). This digestion end point can be mapped to the genome using the remaining single-stranded overhang. Since each TFBS can be mapped from both sides, the resulting distribution of mapped breakpoints is also bimodal and resembles that of conventional ChIP-seq, with the ‘fragment sizes’ corresponding directly to the actual sizes of the TF footprints. To our knowledge, no computational method has been specifically developed for the analysis of ChIP-exo data. We therefore assessed how well peakzilla and other methods perform on ChIP-exo datasets. We called human CTCF binding sites from published ChIP-seq data (Cuddapah et al., 2009) and ChIP-exo data (Rhee and Pugh, 2011). Peakzilla reported estimated fragment sizes of 98bp for ChIP-seq and 36bp for ChIP-exo (Figure 5c) and had the highest resolution (smallest peak-to-peak distance) among all methods tested (Figure 5d). This suggests that peakzilla is well suited for high-resolution ChIP-seq data, including ChIP-exo.
4 DISCUSSION

Understanding how combinations of TFs bind to DNA to regulate gene expression is one of the most pressing questions of today’s biology. Its importance is witnessed by recent community efforts that aim to determine all functional elements in the genomes of model organisms and the human (e.g. ENCODE (ENCODE Project Consortium, 2004), modENCODE (Celinker et al., 2009), Mouse ENCODE (Mouse ENCODE Consortium et al., 2012)). The availability of high-throughput sequencing at low cost widely promoted the use of the ChIP-seq methodology and an enormous number of datasets for different TFs from various species, developmental stages, or tissues are becoming available. This enables the identification of in vivo binding sites and thus enhancers that contain multiple binding sites for a single TF or multiple different TFs. While it is widely accepted that enhancers are characterized by clusters of TF binding motifs (Berman et al., 2002; Schroeder et al., 2004), it has remained less clear to what extent each of several TFs. While it is widely accepted that enhancers are characterized by clusters of TF binding motifs (Berman et al., 2002; Schroeder et al., 2004), it has remained less clear to what extent each of several clustered TF binding motifs is bound in vivo (Yáñez-Cuna, Dinh, et al., 2012). Similarly, potential constraints on the relative distance or orientation of co-bound TFs have remained unclear, yet might be crucial to understand the molecular mechanisms and to decipher the sequence basis of gene regulation (Yáñez-Cuna, Kvon, et al., 2012).

To experimentally address this question, it is important to resolve closely spaced binding sites and precisely predict their location from ChIP-seq data, challenges which current improvements to the ChIP methodology have started to address (e.g. ChIP-exo (Rhee and Pugh, 2011)). However, whereas many computational tools exist to identify enriched regions (peaks) from ChIP data (‘peak calling’), many of them are not designed to fully leverage these improvements, e.g. the increased resolution or the vastly increased number of deep sequencing reads of modern deep sequencing (Chen et al., 2012). To meet these challenges, especially the need for discovering TFBSs at high resolution, we have developed a new computational method, peakzilla.

The importance of high resolution and precision is also supported by alternative efforts to correctly position predicted TFBSs, for example by taking the location of enriched sequence motifs into account (Guo et al., 2012; Boeva et al., 2010; Wu et al., 2010). Although TFBSs predicted by peakzilla coincide very well with the known sequence motifs of the respective TFs, it is important to note that we chose to predict the TFBS locations from the ChIP-seq data alone, without taking sequence motifs into account: it is well established that within TFBSs, motifs for different TFs can be more highly enriched in TFBSs than motifs for the precipitated TF itself. For example, the binding sites of most TFs in the early Drosophila embryo are highly enriched in motifs for the TF Zelda (Li et al., 2008; Bradley et al., 2010; Satija and Bradley, 2012; Kvon et al., 2012; Yáñez-Cuna et al., 2012; modENCODE Consortium et al., 2010), such that the Zelda motif - which is sometimes more highly enriched than the motif of the TF of interest - might bias the correct prediction of the TFBSs and possibly hinder the study of relative positioning and orientation of TFBSs.

The combination of maximum experimental resolution and a peak caller like peakzilla thus makes full use of recent ChIP-seq approaches and will be invaluable for testing hypotheses on how combinatorial TF binding realizes the developmental blueprint encoded in the regulatory regions of our genomes. Indeed, closely spaced Twist binding sites resolved by peakzilla coincided and were strongly enriched in known enhancers, corroborating the prevalent model that functional enhancers are characterized by clusters of TFBSs. The increasing number of ChIP studies that determine the in vivo binding sites of transcription factors at high resolution will prove invaluable for our understanding of enhancer function and transcriptional regulation.

5 METHODS

5.1 Peakzilla algorithm

Initially, peakzilla reads the coordinate files of the mapped reads of the IP and – optionally – control sample. These can stem from either single-end (BED format) or paired-end (BEDPE format) deep sequencing data.

Peakzilla then first determines the average fragment size of the sequencing library to determine the peak size that should result from a true TFBS. For paired-end data, this corresponds directly to the average fragment size (i.e. the average distance of the two mapped ends of each fragment). For single-end data, the average fragment size is estimated from the shift size of positive and negative reads in the top 200 enriched regions in the ChIP sample as described before. Peakzilla then defines peak size as two times the fragment size, as all reads from the ends of fragments immunoprecipitated due to a single TFBS will on average lie in this region.

In a second step, the distribution of positive and negative strand reads that are to be expected is modeled. By default, two normal distributions are used with standard deviations stdev = peak-size / 5 and locations of their means at 1/4th and 3/4th of the peak-size, respectively. Alternatively, the user can choose to estimate the model empirically from the average distribution of reads within the top 200 candidate peaks in the ChIP.

To call TFBSs peakzilla first scans the genome counting reads within a ‘double-window’: each putative (candidate) TFBS receives the counts of positive strand reads within a window of the fragment size downstream of the candidate TFBS and the negative strand reads within an equivalent window upstream of the candidate TFBS. This scores all candidates with a raw score defined as the normalized read count in the IP sample (normalized to a library size of 1 million reads) minus the normalized read count in the control (i.e. input) sample (note that the correction with a control sample is optional). Final peaks are the candidates with sums that are local maxima at least one fragment length (1/2 peak size) apart from each other. This scanning mode allows for both fast and comprehensive investigation of large genomes at single-base resolution.

To obtain a final peak score, each raw score is corrected with a multiplicative distribution score [0..1] that assesses the fit of the observed read count distribution to the distribution expected from the model (see above). This fit is assessed by a chi-square test and the chi-square p-value becomes the distribution score, which provides a measure of how likely the candidate peak is a true TFBS (distribution score: 1) or the result of a sequencing artifact (distribution score: 0). Note that to robustly estimate the average fragment size we only count distinct reads i.e. remove duplicates (prior to model building). For peak calling however, we did not remove duplicates but use the model to penalize PCR duplicates. This strategy is more sensitive for datasets on small genomes or with high coverage (see main text).

If a control sample is provided, an empirical FDR is calculated for each peak by repeating the peak-calling step (after fragment size estimation) with swapped IP and control sample and scoring the resulting control peaks by the raw and distribution score. This provides for each final peak score
the number of true and control peaks that achieve this score or better and thus an FDR estimate. Peakzilla reports the TFBSs in a BED-like format including the genomic positions, raw-, distribution-, and final score, FDR, and a peak number according to each peak’s rank. In addition, the control peaks and a log are reported.

The method is illustrated in a flowchart in Supplementary Figure 3. Peakzilla can be downloaded from http://github.com/steinmann/peakzilla or

http://www.starklab.org/data/peakzilla/

5.2 Program implementation

Peakzilla is implemented in Python 2 and runs on both the standard CPython and the fast PyPy interpreter. The program is freely available under the terms of the General Public License at http://github.com/steinmann/peakzilla. It runs from the command line under any Linux distribution or OS X. The only required argument is the name of the file with the aligned reads from the ChIP sample and – optionally – from the control sample (both BED format). In addition, the following parameters can be used: -m to specify the number of candidate binding sites to use to estimate fragment size (default: 200); -l to limit the candidate regions to lengths above a certain minimum length (which may be necessary if the dataset contains a large number of strong PCR artifacts; default: off [1]); -f to set a FDR cutoff (default: off [100]); -c to set an enrichment cutoff (default: 2); -s to set a score cutoff (default: 1); -l to specify logfile (default: log.txt); -e to use an empirical estimate derived from the data for the model instead of a normal distribution (default: off); -p to specify that peaks should be mostly between 50-200 bp). During Illumina library preparation, the samples are run on a 2% gel at 90V for ~2 hours, and fragments corresponding to ~50 bp, ~75, and ~100 bp inserts are cut out of the gel (slices are ~25 bp thick). The final libraries are run a BioAnalyzer to measure the actual average insert size. The high-resolution ChIP-seq data for Twist is deposited on GEO under the accession code GSE40664.

5.3 ChIP-seq datasets

Raw sequencing reads for Twist in Drosophila melanogaster (He et al., 2011) (ArrayExpress accession code E-MTAB-376), CEBPA in Mus musculus (Schmidt et al., 2010) (GEO accession code GSE22078) and PHA-4 in Caenorhabditis elegans (Zhong et al., 2010) (GEO accession code GSE14545) were aligned uniquely using bowtie allowing for 3 mismatches to the corresponding genomes (assemblies dm3, mm9 and ce6 respectively). For NFkB in Homo sapiens (Kasowski et al., 2010) (GEO accession code GSE19486), Ste12 in Saccharomyces cerevisiae (Zheng et al., 2010) (GEO accession code GSE19636) and CTCF ChIP-seq (Cuddapah et al., 2009) (GEO accession code GSE12889) and ChiP-exo (Rhee and Pugh, 2011) (which the authors kindly shared) in Homo sapiens, already mapped reads were used (assemblies hg18, sarCer2 and hg18 respectively).

5.4 Semi-synthetic datasets

We generated semi-synthetic control (input) and ChIP samples by subsampling input samples and ChIP peaks: we generated two 30 million bp artificial chromosomes by repeatedly randomly sub-sampling an arbitrarily selected region from the Twist input sample that showed no strong enrichment, one as a semi-synthetic input control and the second as a ChIP background. We next selected several highly ranking peaks from the Twist ChIP sample that were found by all methods, had no other peak nearby, and showed a regular fragment density distribution. We subsampled them to yield peaks with 5-fold or higher over background and placed such peaks as pairs with defined peak-to-peak distances into the background every 30000 bp for a total of 1000 peak-pairs. For each distance between the semi-synthetic peak-pairs, we combined the semi-synthetic chromosome with the experimental ChIP data for peak calling of the combined set (i.e. parameters are estimated primarily on the experimental dataset as it contains higher peaks).

5.5 High-resolution ChIP-seq

Embryo aged 2-4 hours after egg laying (AEL) were processed and immunoprecipitated with Twist antibodies based on the protocol by He et al., 2011 with slight modifications. Sonication occurs in three microfuges, each with ~80 mg chromatin extracts resuspended in 250 µl A2 buffer, in a Bioruptor sonicator for 15 min on high (30s on and off) at 4°C. After 15 min cooling, the sonication step is repeated, followed by high-speed centrifugation at 4°C for 10 min and pooling of the supernatant (the DNA fragments should be mostly between 200-500 bp). 600µl supernatant are then incubated with 120µl DNase I (RNAsae-free from NEB, to 0.3 U/µl final concentration) and 80µl DNase I buffer for 30 min at 37°C. To stop DNAse I activity, A2 buffer with 10% SDS is added to a give a final concentration of 1% SDS. The extract is then directly used for ChIP (the DNA fragments should now be mostly between 50-200 bp). During Illumina library preparation, the samples are run on a 2% gel at 90V for ~2 hours, and fragments corresponding to ~50 bp, ~75, and ~100 bp inserts are cut out of the gel (slices are ~25 bp thick). The final libraries are run a BioAnalyzer to measure the actual average insert size. The high-resolution ChIP-seq data for Twist is deposited on GEO under the accession code GSE40664.

5.6 Peak calling

The format of the mapped reads was adapted to each method. Peakzilla, SIISSRs (Jothi et al., 2008) version 1.4. cisGenome (Ji et al., 2008) version 2.0, Spp (Kharchenko et al., 2011) version 1.8 and GPS (Guo et al., 2010) version 0.10.1 were run with default parameters. MACS (Zhang et al., 2008) version 1.4.1 was run with an mfold parameter 3,30 and the gsize parameter was adapted for each genome. QuEST (Valouev et al., 2008) version 2.4 was run with the following interactive choices: transcription factor binding sites with recommended (or relaxed) peak calling parameters. PeakRanger (Feng et al., 2011) read extension length parameter was run using peakzilla’s estimated fragment length. Both QuEST and PeakRanger could not be used for the CTCF samples without a control dataset.

5.7 Functional analyses

We used the known motif CACATGT for Twist and the motifs from JASPAR (Sandelin et al., 2004): snai1 (sna MA0086.1), dorsal (dl_1 MA0022.1), NFkB (NFkB1 MA0105.1), CEBPA (CEBPA MA0102.2), pha-4 (Foxa2 MA0047.1), Ste12 (STE12 MA0393.1). We searched for motif de novo using MEME (Bailey and Gribskov, 1998) with 31bp peak around peak summits and for occurrences of the known motifs using MAST (Bailey and Gribskov, 1998) from the MEME suite programs version 4.1.1 with a p-value of 10^-4 for Twist, which corresponds to allowing for one mismatch) in an area of 151 bp (average genomic fragment length) around each peak summit. We called a peak conserved when it overlapped with a peak region in all other Drosophila species from He et al., 2011 (D.simulans, D.yakuba, D.erecta, D.ananasae and D.pseuodoobscura). We overlap peaks with known Twist enhancers from He et al., 2011 and the known mesodermal enhancers from Bonn et al., 2012
(only M for mesoderm at stage 5,6 and 7). To create a set of control peaks, we shuffled peaks randomly within the same chromosomes.

COMPETING INTERESTS
The authors declare that they have no competing interests.

AUTHORS’ CONTRIBUTIONS
AFB, JS, JAK, JZ, and AS conceived the project. JZ wrote peakzilla. AFB benchmarked peakzilla and performed all computational analyses. AFB and AS analyzed the data. SB and JZ designed and performed the high-resolution ChIP-seq experiments. AFB, JZ, and AS wrote the manuscript.

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