Positional distribution of human transcription factor binding sites

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ABSTRACT

We developed a method for estimating the positional distribution of transcription factor (TF) binding sites using ChIP-chip data, and applied it to recently published experiments on binding sites of nine TFs: OCT4, SOX2, NANOG, HNF1A, HNF4A, HNF6, FOXA2, USF1 and CREB1. The data were obtained from a genome-wide coverage of promoter regions from 8-kb upstream of the transcription start site (TSS) to 2-kb downstream. The number of target genes of each TF ranges from few hundred to several thousand. We found that for each of the nine TFs the estimated binding site distribution is closely approximated by a mixture of two components: a narrow peak, localized within 300-bp upstream of the TSS, and a distribution of almost uniform density within the tested region. Using Gene Ontology (GO) and Enrichment analysis, we were able to associate (for each of the TFs studied) the target genes of both types of binding with known biological processes. Most GO terms were enriched either among the proximal targets or among those with a uniform distribution of binding sites. For example, the three stemness-related TFs have several hundred target genes that belong to ‘development’ and ‘morphogenesis’ whose binding sites belong to the uniform distribution.

INTRODUCTION

Elucidating the basic principles that underlie regulation of gene expression by transcription factors (TFs) is a central challenge of the postgenomic era. Reliable experimental and computational identification of TF-binding motifs is an essential step towards this goal. In spite of major technological advances that generated rapidly improving high-throughput measurements of both gene expression (1) and TF binding (2), and intense parallel bioinformatic efforts that produced a large variety of computational methods (3–6) aimed at identifying functionally important TF-binding motifs, very basic questions remain unresolved. Perhaps one of the most pressing outstanding issues concerns the relative importance of proximal versus distal regulatory regions [with respect to the transcription start site (TSS)] in higher organisms.

While for prokaryotes the region in the close vicinity of the TSS is known (7) to play a central role in binding TFs that regulate gene expression, for eukaryotes the prevalent opinion is to the contrary; even though arguments supporting the special role of the proximal region have been presented for yeast (7)—it is believed that distal regulatory regions are most significant, especially for mammals (8,9). Most recently several bioinformatical studies have claimed that even in mammals the proximal region dominates transcriptional regulation in general (10,11) or for particular biological contexts (12). There is no known estimate, however, of the relative abundance of distal as compared with proximal functional binding sites of TFs. There is no clear answer to simple questions such as the abundance of dual-action TFs, that under different conditions and in different pathways switch from proximal to distal regulatory binding. Conversely—do different genes, that belong to a particular biological function or pathway, exhibit the same positional bias in binding TFs that regulate their expression?

The work presented here is an attempt to answer some of these questions by means of analysis of a large number of experimentally derived (13,14) TF binding sites.

To this end, we developed a method for estimating the positional distribution of TF binding sites on the basis of ChIP-chip data, and applied it to recently published experiments on binding sites of nine TFs (13,14), obtained from a genome-wide coverage of promoter regions from 8-kb upstream of the TSS to 2-kb downstream. Even though binding detected by ChIP-chip (in cell lines) is not synonymous to in vivo functional binding that regulates transcriptional activity, knowing the positional distribution of binding sites does contain important, interesting and yet unexplored information.

The resulting estimated binding site distribution reveals an unexpected picture: it is closely approximated by a mixture of two components. One is a sharp peak, localized within 300-bp upstream of the TSS, and the second...
component is a distribution of almost uniform density within the tested region (−8 kb to 2 kb). These two components appear in all nine TFs studied, but their relative weights do depend on the TF. Such a mixture of two distributions suggests that there might be two distinct groups of binding sites which differ in their biological function or in the mechanism by which their function is achieved. Indeed we found that the three TFs, OCT4, SOX2 and NANOG, that constitute a control unit that governs the genetic program of embryonic stem cells (15), communicate with hundreds of genes involved in morphogenesis and development via uniformly distributed binding sites. On the other hand, the internal connections between these three TFs are of both kinds: the corresponding binding sites on the NANOG promoter are from the component proximal to the TSS, whereas on the other two promoters they are from the more distal uniform component. Further analysis and experiments are needed to elucidate other characteristics of the two kinds of binding sites and their possibly differing roles.

**MATERIALS AND METHODS**

**The data and platform**

We used ChIP-chip data from two studies. The first (13) aimed at mapping the binding sites of three TFs, NANOG, OCT4 and SOX2, known to play central roles in the maintenance of key properties of embryonic stem cells. The second study (14) concentrated on six TFs known to be expressed in the liver and believed to be critical for the biology of hepatocytes: HNF1A, HNF4A, HNF6, FOXA2, USF1 and CREB1. For six of the nine TFs there is data from 2 biological replicas; for the remaining three—HNF6, USF1 and CREB1 only single replicas were available.

Both studies used human cells and the same custom designed DNA microarrays (code-named 10array) developed in the Young lab (16), containing 60mer oligonucleotide probes. The probes cover regions that extend from 8-kb upstream to 2-kb downstream of the TSS of about 18,000 annotated human genes. On the average, there is approximately one probe every 280 bp in the covered region. A full account of the technique can be found in Supplementary Material of (13) and (14) or on the web site accompanying those publications (17).

Here, we review only those components of the technique that are essential for understanding our analysis.

After immobilizing the proteins and fragmenting the DNA (into fragments of length of 550 bp on average), part of the resulting material is used for immunoprecipitation (IP), while the other part is reserved for control. The IP-enriched DNA extract is labeled with red fluorescent dye, while the control whole-cell DNA extract is labeled with green. The whole-cell extract (WCE) is assumed to contain any piece of the genome with equal probability (concentration), as opposed to the immunoprecipitated DNA extract that is significantly enriched by DNA fragments to which the TF of interest was bound. Both DNA extracts are applied to the microarray to allow competitive hybridization. The fluorescence intensity is then measured using red and green filters separately for each probe.

**Data analysis pipeline: identifying regions with bound TFs**

**Normalization and preprocessing.** These procedures (described in the Supplementary Material) were used to assign each probe a score (referred to below as $M$-score) indicative of the probability of the presence of a binding site in its vicinity.

**Smoothed $M$-scores.** This score has unit variance and approximately normal distribution; hence $M$-score cutoffs can be interpreted in terms of probability. Since the spacing between adjacent probes was mostly within the resolution limit of chromatin IP (i.e. spacing was comparable to the DNA fragment lengths), a binding event on probe $i$ was called on the basis of the $M$-scores of a triplet of consecutive probes and the value of their ‘triplet $M$-score’,

$$M_i^{(3)} = \frac{M_{i-1} + M_i + M_{i+1}}{\sqrt{3}}$$

Under the assumption of statistical independence of $M_i$ and $M_{i\pm1}$, the smoothed variable $M_i^{(3)}$ is also approximately normally distributed with unit variance (see Section 1.3.3 of the Supplementary Material for verification of this). Using calls from a triplet of probes helps to filter out spurious signals from single isolated probes.

**Identifying bound triplets.** The filtering criterion (described in detail in Section 1.3 of the Supplementary Material) contains four different $P$-value-like cutoff parameters, $t_1$, $t_2$, $t_3$ and $t_n$, and an overall control parameter ($com$).

A triplet centered on $i$ was labeled as bound if it passed the following criteria:

1. $M_i^{(3)} > (com) \cdot t_3$, AND
2. either (2.1) or (2.2),
   
   (2.1) $M_i > (com) \cdot t_2$ AND $[M_{i-1} > (com) \cdot t_2$
   OR $M_{i+1} > (com) \cdot t_3$],
   
   (2.2) $M_i > (com) \cdot t_1$ AND $[M_{i-1} > (com) \cdot t_n$
   OR $M_{i+1} > (com) \cdot t_n$]

See Supplementary Material for the rationale of these criteria, adopted from (13).

In order to avoid the arbitrariness often present when selecting a cutoff significance level, each of the four $P$-value-like cutoffs was initially assigned some reasonable value, adopted from (13), and these were then multiplied by the overall control parameter ($com$) (abbreviation for ‘cutoff multiplier’). The cutoff multiplier was varied from 0.1 to 500 and for each value the whole data analysis pipeline was run (lower multiplier value means stricter cutoff). For each TF, we selected a ‘natural’ value for the cutoff, as described in the Results section.

For each triplet of probes that passed the filters, the region between the two flanking probes (Figure 1) was marked as a bound region. Overlapping bound regions were collapsed into a single bound region.

The preprocessing steps described above resulted in lists of several hundred to several thousand bound regions for each TF. Each bound region is several hundred base pair long (700 on average).
data, the blue curve is the resulting intensity of the red (IP) and green (WCE) channels from NANOG.

Coverage plots

In order to estimate the distribution of binding sites as a function of distance from TSS, promoters containing bound regions were aligned relative to the TSS nearest the bound region and a coverage number was calculated for each nucleotide location (defined with respect to its closest TSS). It is somewhat similar to a histogram, but since the bound regions have different lengths, a simple histogram could not be used. The coverage number of a particular position, at a given distance from the aligned TSSs, is the number of bound regions that contain the nucleotide at this position. That is, we count how many bound regions cover a point at distance \( x \) from the TSS, adding them up for all the genes tested. Figure 2 illustrates this concept. The genomic locations of the genes were taken from the RefSeq genes table from UCSC genome browser, build hg17.

Coverage plots that were obtained from the experimental data for the nine TFs studied are presented in Figure 3. Note that in order to highlight the similarities between the different coverage plots, in each of these figures the coverage numbers are normalized by the area under the curves (the number of detected binding sites of the nine TFs varied between about 100 and 4000, see Table 1).

Obtaining binding site distributions by deconvolution of coverage plots

The genomic locations of the binding sites of a TF are not known; the aim of our analysis is to find, for each of the TFs, that distribution of binding sites (as a function of distance from the TSS), which provides the best fit for the corresponding (experimentally determined) coverage plot. The fit is obtained by a simulation of the entire process of TF-binding events and their identification by the experiment.

Denote by \( Q(x) \) the probability that the nucleotide at distance \( x \) from the TSS belongs to a binding site of a given TF. We choose, for the TF that is studied, a particular \( Q(x) \) from a family of distributions described below. A gene is randomly selected from the list of all possible targets, and a binding site is placed on its promoter at a location \( x \) selected at random from \( Q(x) \). Ten thousand binding sites are generated independently this way, each characterized by a genomic address and a strength parameter \( U \), which represents the binding affinity of the site. The value of \( U \) is sampled at random from a shifted gamma distribution

\[
p(U) = \frac{(U - s)^{k-1}}{\theta^k \Gamma(k)} \exp\left(-\frac{U - s}{\theta}\right)
\]

with parameters: shift \( s = 3 \), shape \( k = 2 \) and scale \( \theta = 3 \), based on the model derived in Ref. (16).

The number of binding sites that were generated for a simulation (10 000) was chosen so that the resulting simulated coverage plot is not noisier than the experimental one. The precise value of the number of sites generated has a minimal effect on the results of our simulations.

Since the locations of all probes are known, we can calculate for each probe a simulated \( M \)-score, determined by its distance \( d \) to the nearest of the 10 000 simulated binding sites and by its strength parameter \( U \):

\[
M\text{-score} = f(d) \cdot U
\]

\[
f(d) = \sum_{l=0}^{\infty} \frac{1}{l!} \sum_{a=d}^{l} p(a)p(l - a)
\]

The influence function \( f(d) \) used to calculate the \( M \)-score was adapted from the Supplementary Material of Ref. (18); \( p(a) \) is the probability that the DNA was cut at a distance \( a \) from the nearest binding site. The distribution \( P_l(l) \) of the DNA fragment lengths \( l \) has been measured (18) and was approximated by a shifted gamma distribution (18). This distribution is related to \( p(a) \) by a convolution,

\[
p_l(l) = \sum_{a=0}^{l} p(a)p(l - a)
\]

Since convolution of two identical shifted gamma distributions is again a gamma distribution (with twice the mean and shift), \( p(a) \) was also taken to be a shifted gamma distribution. The following parameters were used for the simulation: shift \( s = 50 \text{ bp} \), shape parameter \( k = 2 \) and scale parameter \( \theta = 60 \). The simulated \( M \)-scores were...
then fed into the analysis pipeline described above as if they were derived from real raw data and a coverage number plot was generated. We performed simulations using binding site distributions $Q$ parameterized as described below and searched for a $Q$ that yielded a good approximation to the experimentally derived coverage plot. Since the simulation is computationally intensive (about 55 s on Intel P4 2.4 GHz 1 GB RAM for a single run), any systematic fitting method would be difficult to implement and our forward-fit method has to be viewed as an approximate deconvolution.

Figure 3. The fitted deconvolved binding site distributions (blue) and the corresponding simulated ones (cyan) compared with experimental (red) coverage number plots.

Table 1. Summary of the fitted distributions

<table>
<thead>
<tr>
<th>TF</th>
<th>com</th>
<th>$N_c$</th>
<th>$N_g$</th>
<th>$N_{g^*}$</th>
<th>$X_p$</th>
<th>$D_f$</th>
<th>$D_c$</th>
<th>$W_u/W_p$</th>
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<tbody>
<tr>
<td>NANOG_M3</td>
<td>100</td>
<td>2467</td>
<td>2394</td>
<td>1683</td>
<td>−180</td>
<td>260</td>
<td>827</td>
<td>6.5</td>
</tr>
<tr>
<td>OCT4_M4</td>
<td>100</td>
<td>1546</td>
<td>1536</td>
<td>623</td>
<td>−120</td>
<td>260</td>
<td>914</td>
<td>5.7</td>
</tr>
<tr>
<td>SOX2_M3</td>
<td>5</td>
<td>314</td>
<td>341</td>
<td>1271</td>
<td>−200</td>
<td>165</td>
<td>624</td>
<td>5.7</td>
</tr>
<tr>
<td>FOXA2_M2</td>
<td>50</td>
<td>1066</td>
<td>1126</td>
<td>890</td>
<td>−130</td>
<td>260</td>
<td>862</td>
<td>6.1</td>
</tr>
<tr>
<td>HNF1A_M2</td>
<td>50</td>
<td>1052</td>
<td>1097</td>
<td>1016</td>
<td>−180</td>
<td>212</td>
<td>802</td>
<td>5.7</td>
</tr>
<tr>
<td>HNF4A_M1</td>
<td>30</td>
<td>3889</td>
<td>3637</td>
<td>4519</td>
<td>−180</td>
<td>212</td>
<td>860</td>
<td>5</td>
</tr>
<tr>
<td>HNF6_M1</td>
<td>5</td>
<td>782</td>
<td>886</td>
<td>1306</td>
<td>−240</td>
<td>283</td>
<td>1321</td>
<td>6.1</td>
</tr>
<tr>
<td>CREB_M1</td>
<td>10</td>
<td>1008</td>
<td>1290</td>
<td>2197</td>
<td>−150</td>
<td>212</td>
<td>708</td>
<td>1</td>
</tr>
<tr>
<td>USF1_M1</td>
<td>1</td>
<td>111</td>
<td>151</td>
<td>1632</td>
<td>−200</td>
<td>212</td>
<td>606</td>
<td>1</td>
</tr>
</tbody>
</table>

Com, CutOff Multiplier; it represents the $P$-value cutoffs selected for the TF as described later. $N_c$, number of bound regions (bound regions are defined in Methods section). $N_g$, number of bound genes (a bound gene is defined here as any gene for which there is a bound probe within 10-kb upstream to 3-kb downstream of its TSS). $N_{g^*}$, number of bound genes as previously reported by (13, 14). $X_p$, peak position relative to TSS (in base pairs). $D_f$, width in base pairs of the fitted peak at half max (2.36 σ). $D_c$, width of the peak of the measured coverage plot at half max. $W_u/W_p$, ratio of weights of the uniform component and of the peak in the distribution, it is also the ratio of the number of binding sites that are distributed uniformly and the number of binding sites that are localized within the peak.

Family of binding site distribution functions tested

The most prominent feature of the experimental coverage number graphs (Figure 3) is the peak at around 150 bp upstream of the TSS. The structure away from this peak, namely the rapid decrease to zero at $−8$ kb and $+2$ kb (the edges of the genomic region covered by probes), is due to microarray design and is consistent with a uniform binding site distribution (see Results section). Therefore, we modeled $Q(x)$ as the sum of a uniform distribution and one or
more Gaussians (up to five, but usually one or two sufficed). The centers and widths of the Gaussians and the weights of all the components were the parameters that were varied to identify the distribution that gave best agreement with the experimental coverage plot.

**Gene Ontology enrichment analysis**

For a group of genes of interest $G$ (such as those targets of a TF whose binding is close to the TSS), we performed Gene Set Enrichment Analysis (GSEA) (19). Admittedly over-representation of a specific Gene Ontology (GO) category among the genes of $G$ does not prove regulation of these genes by the relevant TF, but co-regulation is a most plausible reason for the observed enrichment. To exclude the possibility that the over-representation of a GO group was due to a chance fluctuation in a random set of stray binding sites, the observed enrichment $P$-values were submitted to a stringent false discovery rate (FDR) analysis (20).

**RESULTS**

Coverage plots for the nine tested TFs exhibit a prominent peak near the TSS

The coverage plots obtained from the experimental data when the processing and analysis steps described above were implemented are shown in Figure 3. It is important to remember that the number of identified targets depends on the thresholds and cutoff parameters that were used; the dependence of the coverage plots on these parameters and the manner in which they were determined for each TF are discussed in detail below.

The coverage plots obtained from the experiments are in red; for six out of the nine TFs the experiments were repeated twice, and coverage plots were prepared for each repeat. In four out of the six cases the two repeats are in good agreement with each other; in all six cases we chose to adopt the repeat with the higher peak (see Supplementary Figure S5). For each of the nine TFs studied the coverage plot exhibits a sharp peak close to the TSS. This strong peak near the TSS is the most prominent feature of these plots. As explained in the Methods section, these coverage plots were normalized, in order to emphasize that the peak near the TSS is shared by all nine TFs studied.

Binding site distributions obtained by fitting coverage plots

In Figure 3, we present for each of the nine TFs studied the ‘optimal’ binding site distribution $Q(x)$ that produced a good fit to the experimentally obtained coverage plots. The quality of the fit can be assessed from the same figure. Good fits were obtained when a mixture of a uniform background distribution with one (or more) Gaussians was used. In several cases mixing a single sharp Gaussian, located near the TSS, sufficed; even when more than one was needed, the first one, located upstream close to the TSS, was the most prominent by far. The parameters of each of the binding site distributions, obtained for the nine TFs tested, are summarized in Table 1, together with numbers of bound regions (see Methods section) and bound genes. A bound gene is defined here as any gene for which there is a bound probe in the interval from 10-kb upstream to 3-kb downstream of its TSS. Note that a single bound probe can give rise to more than one bound gene (sense and antisense).

The number of bound genes varies from about 150 for USF1 to 3600 for HNF4A. Note that for several TFs it differs considerably from the numbers reported previously by Refs (13,14). These differences are mainly due to the different values of the $P$-value thresholds, selected as described later. The position of the peak is close to $-200$ bp for all TFs, and its width varies between 165 and 260. Using these numbers and by inspection of the deconvolved BS distributions of Figure 3, we identified for each target gene the ‘proximal region’ as the interval $[-300, +300]$ bp (on both sides of the TSS), for all the TFs studied. If a particular region $R$ was identified as proximal for gene $G_1$ and distal for $G_2$, a binding event in $R$ is counted as proximal binding to $G_1$ and distal to $G_2$. Note that the width of the fitted distribution of binding sites is about 25% of the width obtained from the coverage plots; hence deconvolution of the coverage plots indeed sharpened significantly the resolution at which ChIP-chip data can be used to identify binding site positional bias.

The picture that emerges indicates that a TF may have two classes of binding sites, that probably differ in their biological function and the mechanism by which this function is achieved.

Comparison of binding sites within the peak and outside using GO

We turned to look for such a difference of functions, using GO (21) annotations of genes bound by the nine TFs studied. For this end, the genes bound by a particular TF were split into two disjoint groups—one contained genes that have a probe detected as bound, located within the gene’s proximal region, and the other contains the rest of the TF’s putative targets—i.e. genes with one or more bound probe, none of which lie within the gene’s proximal region. The first group is assumed to include most of the genes that have a binding site on their promoter within the peak. Both groups were subjected separately to hypergeometric GO enrichment analysis (see Section 2.5 of the Supplementary Material for details of the calculation of $P$-values and FDR correction), using only the ‘biological process’ type of GO annotations. Results of this analysis are depicted graphically on Figure 4. Genes bound by HNF6, one of the six liver-associated TFs, were not enriched by any GO group. For the other TFs we do find association of biological processes with binding location; the GO groups are divided roughly into two subsets, (i) containing a variety of categories related to metabolism, RNA processing and splicing, cell cycle and more (see Supplementary Table S4 for a full list) and (ii) mainly regulatory groups and developmental processes. The GO categories of subset (i) are enriched mainly by target genes of the liver-associated TFs and the binding sites are close to the TSS. The GO
categories of type (ii) are enriched mostly by genes that bind the three stemness-related TFs, with binding sites far from the TSS. The finding that different GO terms are enriched in different groups supports the possibility of different functions associated with binding sites within and outside the peak. We list below a few selected observations.

For example, the group of genes with probes bound by OCT4, far from the TSS, is enriched with a GO term ‘organ morphogenesis’ with an FDR corrected P-value of $4.2 \times 10^{-6}$, while the group of genes with probes bound by OCT4 close to the TSS is not enriched with the same term.

For some TFs such as USF1 and CREB1 there are enriched terms only in the group of genes with bound probes close to the TSS. The situation is reversed for OCT4. HNF4A has several GO terms enriched in both the far and the close groups. NANOG has some terms like mitosis enriched only in the close group, other terms, like morphogenesis, are enriched only in the far bound gene group, and yet others like RNA metabolism are enriched in both (Table 2).

It is interesting to note that for genes bound by the stem cell TFs NANOG, OCT4 and SOX2, development-related GO categories are enriched only among the genes with a binding site far from the TSS (Table 2 and Supplementary Table S4).

Proximal and distal binding sites in transcriptional circuitry

It is interesting to analyze the different roles played by the two kinds of TF binding sites in the transcriptional circuits in which they participate. Figure 5 presents the connections within the three stem cell-related TFs and their connections to the ‘external world’ of transcriptional targets. Interestingly, NANOG itself is the target of exclusively proximal internal binding, whereas SOX2 and OCT4

<table>
<thead>
<tr>
<th>GO category</th>
<th>FDR corrected P-values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Close to TSS</td>
</tr>
<tr>
<td>Development</td>
<td>1.00E+00</td>
</tr>
<tr>
<td>Transcription</td>
<td>6.37E-02</td>
</tr>
<tr>
<td>Nucleobase, nucleoside, nucleotide and nucleic acid metabolism</td>
<td>6.95E-09</td>
</tr>
<tr>
<td>Anatomical structure development</td>
<td>1.00E+00</td>
</tr>
<tr>
<td>RNA metabolism</td>
<td>2.25E-06</td>
</tr>
<tr>
<td>Biopolymer metabolism</td>
<td>5.91E-09</td>
</tr>
<tr>
<td>RNA biosynthesis</td>
<td>3.88E-02</td>
</tr>
<tr>
<td>Transcription, DNA-dependent morphogenesis</td>
<td>3.68E-02</td>
</tr>
<tr>
<td>Regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolism</td>
<td>1.00E+00</td>
</tr>
<tr>
<td>Regulation of transcription</td>
<td>4.06E-01</td>
</tr>
<tr>
<td>Transcription from RNA polymerase II promoter</td>
<td>4.13E-01</td>
</tr>
<tr>
<td>Regulation of cellular metabolism</td>
<td>4.16E-01</td>
</tr>
<tr>
<td>Regulation of metabolism</td>
<td>5.76E-01</td>
</tr>
</tbody>
</table>

Notice that there are several development-related categories that are not enriched in the group of genes with binding site close to the TSS. See Supplementary Material for details about the calculation of P-values and FDR correction and a full list of enriched categories.
have distal internal binding sites. As to external binding, the GO processes of development, morphogenesis, regulation of transcription and sensory organ development are controlled by binding sites from the distal, uniformly distributed class (of all three TFs).

Another interesting observation concerns the circuit of liver-related TFs: most of the internal interactions between the TFs in this group are either through binding close to the TSS or within the gene.

Having discussed in detail the main characteristics of the coverage plots and the fitted distributions obtained from their deconvolution, as well as the biological observations concerning the two kinds of binding sites we found, we turn to some technical details that must be addressed. First, we rule out two possible trivial sources of the strong peak we found; next, present a purely computational test of the main result and describe the manner in which the $P$-value thresholds (that were used to identify binding events of the TFs studied) were set.

Addressing several possible concerns about the analysis

We describe here several possible reasons and artifacts that could have misled us to reach the conclusion described above.

The effect of probe density

The first question to consider is whether the strong peak reflects nothing but the density of probes represented on the chip. Even though our simulations generate binding site distributions that fit the data (coverage plots) using the actual genomic locations of the probes placed on the chip, we wish to demonstrate here clearly that the distributions we found, and especially the sharp peak near the TSS, are not due to the probe distribution.

Clearly, if all probes are placed in a narrow region near the TSS, the coverage plot will have nonzero values in this region only. Indeed, since the microarray does not cover promoter regions outside the interval $[-8, 2]$ kb from the TSS, we get zero coverage outside this region. Additionally, the probe density within the covered region is not uniform, as can be seen on Figure 6 (red curve): it is higher near the TSS. In order to understand how this probe density variation influences the coverage number, we performed a simulation of the measurement process starting with a hypothetical TF with a uniform distribution of binding sites as a function of distance from the TSS. The blue curve on Figure 6 is the average of the coverage number plots obtained from 100 such simulations. Comparison of this simulated curve with that generated from real data of HNF1A is presented on Figure 6. Clearly, the peak of the real data is much sharper and narrower. Hence, the prominent peaks observed for all nine TFs cannot be attributed to the probe density variation, while the gradual decrease further away from the TSS most probably reflects just that.

The effect of GC-content variation

A second possible artifact that could in principle produce such a peak is the nonuniform GC content of the promoters of human genes. As seen in Figure 7A, the GC content increases from 45% far from the TSS to a peak value of about 65% near the TSS. Higher GC content means stronger binding of the DNA fragments to the corresponding probes and hence higher $M$-scores and, possibly, a higher density of detected binding sites. Since the bias introduced by the GC content is, on the average, similar for the IP and the WCE, we expect that taking the ratio of the intensities of the same probe in the two channels reduces significantly the effect of the GC variation on our measured signal. Indeed, as expected, the fluorescent intensity of the microarray probes in both channels is highly correlated...
with their GC content (correlation coefficient of about 0.7 in the data used here). On the other hand, correlation coefficients between the M-score and GC content are very low (order of 0.01) for most TFs, demonstrating that working with the M-score (which is basically a scaled log ratio of intensities in the two channels) successfully cancels the variation introduced by the GC content. As can be seen on Figure 7B, for NANOG the coverage number peak is about 150-bp upstream from the TSS, while the GC content peak, as well as the peaks in intensity of the red and green channels, are about 70-bp downstream. This significant shift in peak position serves as convincing evidence that the sharp peaks in coverage number plots are not an artifact caused by GC-content variation.

A computational test

To obtain further evidence for the fact that the peaks of the coverage plots and the resulting fits for binding site distribution are not due to some artifact of the method, we performed a purely computational test. We used the database underlying the ‘TFBS Conserved’ track in the UCSC genome browser (22,23), http://genome.ucsc.edu/cgi-bin/hgTrackUi?g=tfbsConsSites. It was generated using the TRANSFAC (24) collection of positional score matrices (PSSMs) representing the binding preferences of TFs. The database contains the locations and scores of TF binding sites conserved in the human/mouse/rat alignment. In general, the number of conserved binding sites in this database is too small to construct meaningful positional histograms for most TFs, but for HNF1A, USF1 and CREB1 there was a large enough number. The resulting histograms are very similar to what we found from the ChIP-chip experiments (see Supplementary Figure S8). Since these coverage plots are derived in a purely computational way, they are not influenced by GC concentration in the same way as hybridization-based experiments.

Are the identified binding sites functional?

As stated in the Introduction, the binding events detected by ChIP-chip in cell lines may not necessarily correspond to functional binding, that actually regulates transcriptional activity, that takes place in vivo. Performing in vitro and in vivo experiments is the only way to establish beyond doubt the functionality of a binding site. Using in silico bioinformatic methods to deduce functionality are at odds with the spirit and aims of this work, in which we tried to limit the analysis to experimentally derived binding events.

We did try to address specific concerns, in particular regarding a possible reasonable suspicion about functionality of the distal binding events. Over the 10-kb long DNA strands scanned for binding one may (and will) have ‘stray’ binding sites because of purely statistical reasons. We tried several tests, direct and indirect, to rule out the suspicion that our results reported above were based on such nonfunctional statistical binding events. As a sanity check of the assumed functionality of the distal binding sites, we investigated the promoters of a group of housekeeping genes [derived from (25)]. Housekeeping genes are believed to be proximally regulated; we found that housekeeping genes had OCT4 and NANOG binding sites, and, as expected, these had a much stronger tendency for proximal binding than the full genome-wide set of bound promoters. The weight ratio $W_u/W_p$ dropped from ~6 (Table 1) to about 1.5 (see Supplementary Figure S14). Another (experiment-based) test is described below; by lowering the threshold for identification of a binding event from the data, we move from a regime where the identified binding events are dominated by strongly bound functional sites, to one where weaker stray statistical binding events constitute the majority. Since the origins of the two types of binding are very different, the number of detected binding sites should behave differently, as a function of the varying threshold.
in the two regimes. Observation of such a difference (change of slope, apparent discontinuity, etc.) is indicative of the fact that we indeed have two different types of binding sites, one of which is statistical and the other—most probably functional. We have demonstrated that for most of the studied TFs indeed such a crossover was observed for the distal binding sites (for which statistical binding occurs with high probability). These results are presented below and in Supplementary Figures S11. Since for seven out of the nine TFs studied the weight of the distal uniform distribution is about six times the weight of the proximal one (Table 1), such a crossover induces a similar trend in the total number of binding events as a function of cutoff can be divided into three different types.

Selecting the P-value cutoffs for each TF

As described in the Methods section, we used a single parameter to control the cutoff values of the four P-values that were used to decide whether a probe was considered bound or not by the TF. This CutOff Multiplier is referred to as com in the various figures and their legends. It was varied between 0.1 and 500 (lower values mean stricter cutoff, i.e. more rigorous filtering and smaller number of regions identified as bound). The numbers of bound regions and genes that were identified for each TF are reported in Table 1, which also contains a first column the value of com used for each TF. Obviously the number of identified bound regions depends on the value of com, and we discuss here the manner in which we selected the values that were used. A related question concerns the extent to which the coverage plots, and in particular the sharp peak near the TSS, depend on com.

As described above, the general underlying assumption we make is that for low values of com we have very few false positives but many false negatives. As com increases, more binding events are identified, until at some point the resulting filter loses its meaning and the additional binding events are dominated by noise. Hence, we are looking for a change of the behavior of either the number of binding events as a function of com, or of some other important property of the resulting coverage plots.

The observed behavior of coverage number plots as a function of cutoff can be divided into three different types. For five TFs the shape remained almost invariant as the cutoff multiplier increased, and deteriorated quickly beyond some ‘critical’ value, above which the coverage plot resembled the one simulated for a hypothetical TF with uniform distribution of binding sites (Figure 6). As shown in Figure 8, HNF1A belongs to this type (the other four are FOXA2, HNF6, HNF4 and SOX2, see Supplementary Figure S9). As shown in Supplementary Figure S10, the total number of bound regions also exhibits a fairly sharp anomaly for these five TFs at the critical value of com (either a change of slope or apparent discontinuity). The critical value of com differs between TFs and may be different even between experimental replicas for the same TF (Figure 6B).

A different behavior was exhibited by coverage number plots for NANOG (Figures 6C and D) and OCT4 (Supplementary Figure S9). For these TFs the peak value initially increases with com until a maximum is reached around com = 100, and then decreases.

The peaks of the coverage number plots of the remaining two TFs, USF1 and CREB1, decreased monotonically with com without any apparent discontinuities. It is interesting to note that these two TFs have the highest peaks, with coverage numbers nearly zero outside the peak (Figure 3).

For the five TFs that exhibit the first type of behavior we selected com just below the critical value. The rationale is that for relatively stringent cutoffs we get coverage number plots that correspond to a relatively clean list of binding sites with few false positives. It can be assumed that while the relative coverage number plot does not change with loosening cutoff, the growing list of binding sites maintains a noise level similar to the initial one, until the critical value, beyond which many false binding sites enter the list and the noise level rises affecting the coverage number plot, and we choose com at a value before that happens.

The cutoffs for NANOG and OCT4 were selected to get the highest peak. Since the peak heights and number of bound genes for USF1 and CREB1 exhibited no obvious discontinuity (see Supplementary Figures S9 and S10) and therefore provided no hint for the selection of cutoffs, we selected rather conservative values of com (1 for USF1 and 10 for CREB1). This resulted in relatively small numbers of genes detected as bound by USF1 (and to some extent, by CREB1), compared with other TFs and to what was reported by Refs (13,14).

DISCUSSION

We used ChIP on chip data for nine TFs to pose and answer questions regarding the genome-wide distribution of binding sites with respect to the various genes’ TSS. From the experiment we extracted coverage plots, from which we estimated the distribution of binding sites. This step was performed by using the experimental DNA fragment length distribution, the distribution of binding strengths and the actual addresses of the probes on the genome. The main result of this analysis is that the distribution of binding sites can be expressed as the sum of a very narrow peak close to the TSS, and a uniform background distribution.

We ascertained that our results are not due to various artifacts. For example, the effect of the nonuniform GC content (high near the TSS) on hybridization efficiency was assessed by a careful comparison with purely in silico results (available only for a subset of the TFs). The distortion caused by the nonuniform distribution of probes on the chip (denser near the TSS) was also taken into account. Thresholds of binding calls were set by a careful analysis of the dependence of the numbers of binding sites and the relative weights of the two components of the distribution on variation of the threshold. For most TFs, we observed a fairly sharp change of behavior of these quantities, allowing us to identify the value of the threshold at which such changes set in, indicating a change in the strength of the contaminating noisy background signal. The number of target genes was found to be large. Even though most of the TFs studied were known
to be hubs of transcriptional networks, the fact that the number of target genes of a TF is on the order of thousands (and that this seems to be the rule, rather than the exception!) seems to be fairly surprising.

Finally, we performed a functional analysis of the two types of genes: those that are regulated by binding sites proximal to the TSS and those whose binding sites belong to the uniform component of the binding site distribution. For the three TFs that regulate and govern embryonic stemness, we observed that the target genes associated with morphogenesis, development and regulation of transcription predominantly belong to the class with uniformly distributed regulatory binding sites. The molecular reasons behind this and the role this bias plays needs to be explored in the future.

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REFERENCES


