Genomic Regions Flanking E-Box Binding Sites Influence DNA Binding Specificity of bHLH Transcription Factors through DNA Shape

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SUMMARY

DNA sequence is a major determinant of the binding specificity of transcription factors (TFs) for their genomic targets. However, eukaryotic cells often express, at the same time, TFs with highly similar DNA binding motifs but distinct in vivo targets. Currently, it is not well understood how TFs with seemingly identical DNA motifs achieve unique specificities in vivo. Here, we used custom protein-binding microarrays to analyze TF specificity for putative binding sites in their genomic sequence context. Using yeast TFs Cbf1 and Tye7 as our case studies, we found that binding sites of these bHLH TFs (i.e., E-boxes) are bound differently in vitro and in vivo, depending on their genomic context. Computational analyses suggest that nucleotides outside E-box binding sites contribute to specificity by influencing the three-dimensional structure of DNA binding sites. Thus, the local shape of target sites might play a widespread role in achieving regulatory specificity within TF families.

INTRODUCTION

Transcriptional regulation is effected primarily by sequence-specific transcription factors (TFs) that recognize short DNA sequences (5–15 bp long) in the promoters or enhancers of the genes whose expression they regulate (Bulyk, 2003). Determination of the DNA recognition properties of TFs is essential for understanding how these proteins achieve their unique regulatory roles in the cell.

TFs are typically annotated according to the structural class of their DNA binding domains. Members of a particular class (i.e., paralogous TFs) often have similar DNA binding preferences (Badis et al., 2009). However, despite apparently shared binding specificities, individual TF family members often exhibit nonredundant functions. In some cases, differences in the core DNA binding site motifs have been shown to contribute to differential in vivo binding by closely related TFs (Busser et al., 2012; Fong et al., 2012; Grove et al., 2009; Wei et al., 2010). However, in many cases, the DNA motifs of paralogous TFs are virtually identical, and still the proteins select different genomic targets in vivo. In these cases, interactions with protein cofactors are thought to be responsible for differential in vivo DNA binding of paralogous TFs. However, such cofactors can be difficult to identify, and only a few conclusive examples are known (e.g., Hollenhorst et al., 2009; Mann and Chan, 1996; Slattery et al., 2011). Another factor that determines in vivo TF binding is the local chromatin environment (Arvey et al., 2012; Lelli et al., 2012; Thurman et al., 2012; Zhou and O’Shea, 2011). Nevertheless, protein cofactors and chromatin context are unlikely to completely explain differential binding specificity of paralogous TFs.

Here, we investigate a potential mechanism through which TFs with highly similar DNA binding motifs can achieve differential binding in vivo. Several studies have indicated that nucleotides flanking TF binding sites (i.e., nucleotides outside the core DNA binding site motif) can affect binding specificity (Leonard et al., 1997; Morin et al., 2006; Nagaoka et al., 2001; Rajaram and Kerppola, 1997). Thus, we investigated whether the genomic context of putative TF binding sites differentially affects binding of paralogous TFs.

In this case study, we examined S. cerevisiae basic-helix-loop-helix (bHLH) TFs Cbf1 and Tye7. These factors have highly similar DNA binding motifs (Maclsaac et al., 2006; Zhu et al., 2009) but interact with different sets of genomic regions in vivo (Harbison et al., 2004) (Figure 1). Importantly, these differences in in vivo DNA binding are not due to the TFs being active under different conditions, in which the accessibility of potential DNA binding sites might be different (as has been observed for other
bHLH factors; Fong et al., 2012). Instead, the Cbf1 and Tye7 ChIP-chip data (Harbison et al., 2004) were both collected from the same culture conditions (YPD), in which the two proteins had access to the same E-box (CAnnTG) binding sites. Thus, mechanisms other than chromatin accessibility contribute to differential in vivo DNA binding by these two TFs.

Using custom-designed “genomic-context protein binding microarrays” (gcPBMs), we analyzed binding of Cbf1 and Tye7 to their putative E-box binding sites centered within native genomic sequences. Our gcPBM data show that when placed within genomic flanking sequences, E-box sites are bound with different preferences by these two proteins. Importantly, these differences in binding are observed not just in vivo but also in vitro, where cofactors or histones are not present. Thus, the DNA sequence itself is responsible for differential binding by these two TFs.

Notably, the identified differences in DNA binding preferences between Cbf1 and Tye7 are not apparent from these proteins’ binding site motifs (Figure 1). Therefore, to further investigate the source of the binding differences, we used the gcPBM data in a regression analysis to build computational models of the DNA binding specificities of Cbf1 and Tye7. Compared to traditional DNA motif models (i.e., position weight matrices, PWMs), these models are more accurate in predicting in vitro DNA binding. Examination of the sequence features that are important for our regression models revealed that features located in the genomic sequences flanking the E-boxes contribute to DNA binding specificity. Our results show that differences in the intrinsic sequence preferences of related TFs, even when they occur outside the core DNA binding site motif, can contribute to differential TF-DNA binding. Importantly, these differences in intrinsic sequence preferences, as identified through our in vitro studies, can partly explain differential DNA binding in vivo.

DNA sequences flanking the E-box motif, which were found to affect binding of Cbf1 and Tye7, do not typically form base-specific contacts with bHLH proteins (De Masi et al., 2011). Thus, we hypothesized that these sequences contribute to binding specificity indirectly by influencing the three-dimensional structure of the DNA binding sites. A role of DNA shape in achieving binding specificity of TFs has been suggested for Drosophila Hox proteins (Joshi et al., 2007; Slattery et al., 2011) and other protein families (Rohs et al., 2009, 2010). However, for these examples, DNA shape was a result of the nucleotide sequence within the TF binding site. Here, we found that nucleotides flanking Cbf1 and Tye7 binding sites alter structural properties of their DNA targets and, thus, contribute to their differential binding preferences. This finding reveals a mechanistic explanation for the role of nucleotides that are located outside of a binding motif to TF binding specificity. Moreover, this finding suggests why TFs bind in vivo to only a subset of available target sites with identical core motifs. Future studies will investigate the generality of our findings within the bHLH family as well as other TF families. Our results...
suggest that the local shape of DNA binding sites might play a critical role in achieving regulatory specificity within TF families.

RESULTS

**S. cerevisiae** TFs Tye7 and Cbf1 Recognize Highly Similar DNA Sequence Motifs Despite Binding Different Target Sites In Vivo

TFs from the bHLH protein family recognize DNA binding sites containing the E-box motif (CAnnTG) (Atchley and Fitch, 1997), with different family members sometimes having different preferences for the two central base pairs of the E-box (De Masi et al., 2011; Fong et al., 2012; Grove et al., 2009). In *S. cerevisiae*, the bHLH family comprises eight TFs that have diverse functions. Among these TFs, Cbf1 and Tye7 are most similar in terms of their DNA binding specificities (Figure 1A) (Cherry et al., 2012; Maclsaac et al., 2006; Zhu et al., 2009), with both having a strong preference for the E-box CACGTG. However, the sets of in vivo targets bound by Cbf1 and Tye7, as determined by ChiP-chip (Harbison et al., 2004), barely overlap (Figure 1B), and the two TFs regulate different processes: Cbf1 is involved in methionine biosynthesis and chromatin remodeling (Cai and Davis, 1990; Kent et al., 2004), whereas Tye7 plays a major role in the regulation of glycolytic genes (Nishi et al., 1995). It is currently unclear how two TFs with highly similar DNA binding motifs attain their regulatory specificities.

The Cbf1 and Tye7 DNA binding motifs, although very similar, are not identical. For this reason, we first asked whether the small differences in these motifs (Figure 1A) can explain, at least in part, their differential binding in vivo. Using DNA motifs derived from in vivo (ChiP-chip) and in vitro (PBM) data, we computed an area under the curve (AUC)-based enrichment score (see Experimental Procedures) (Gordán et al., 2009) for the enrichment of Cbf1 and Tye7 motifs in vivo DNA binding data (Harbison et al., 2004), where a value of 1.0 corresponds to perfect enrichment, and a value of 0.5 corresponds to the enrichment of a random motif. If the DNA motifs can explain, even in part, the differential in vivo binding, then we would expect the Cbf1 motif to be significantly more enriched in the Cbf1 ChiP-chip data and the Tye7 motif to be significantly more enriched in the Tye7 ChiP-chip data. However, we find that the motifs of both of these TFs are equally well enriched in both the Cbf1 and Tye7 ChiP-chip data sets (Figure 1C), which indicates that the information in the existing PWMs does not explain why these TFs bind different sites in vivo. A similar enrichment analysis that included the *S. cerevisiae* bHLH protein Pho4, which also has a strong preference for the E-box CACGTG, revealed that the Pho4 PWM was not significantly enriched in the Cbf1 or Tye7 ChiP-chip data (Gordán et al., 2009). The same study showed that the Tye7 PWM was not significantly enriched in the Pho4 ChiP-chip data, and the Cbf1 PWM was only marginally enriched (in agreement with previous studies of Pho4 and Cbf1; Zhou and O'Shea, 2011). Thus, differences in the PWMs of Pho4 versus Cbf1/Tye7 can explain, at least in part, the differences in their in vivo DNA binding. However, the Cbf1 and Tye7 PWMs are too similar to explain why these two TFs interact with distinct sets of E-box sites in vivo.

An alternative way to represent the DNA binding specificities of TFs utilizes data generated by universal PBMs. PBM experiments performed on universal arrays (Berger et al., 2006) provide measurements of TF binding to all possible 8 bp sequences (8-mers), as well as a measure of the PBM enrichment score (E-score) for each 8-mer. E-scores range from −0.5 to +0.5, with higher values corresponding to higher sequence preference; typically, E-scores >0.35 correspond to specific TF-DNA binding (Berger et al., 2006; Gordán et al., 2011). We compared previously published 8-mer E-scores for Cbf1 and Tye7 (Zhu et al., 2009) and found that, although they are correlated, the binding specificities of the two TFs are not identical (Figure 1D); there are many 8-mers that are strongly preferred by only one of these two TFs. We did not observe such differences between universal PBM experiments performed for the same factor (Cbf1) on two different universal array designs (Figure 1D). This suggests that Cbf1 and Tye7 have slightly different specificities in vitro.

**Tye7 and Cbf1 Bind with Different Specificities to Putative DNA Binding Sites in Their Genomic Context**

To further investigate the differences in the in vitro DNA binding specificities between Cbf1 and Tye7, we designed a custom PBM containing putative Cbf1 and Tye7 DNA binding sites in their native genomic context (Figures 2A–2C). In this array design, termed gcPBM, we initially focused on genomic regions bound in vivo by either of the two TFs, defined as regions with p < 0.005 in Cbf1 or Tye7 ChiP-chip data (Harbison et al., 2004). To identify putative TF binding sites in the *S. cerevisiae* genome, we used universal PBM data for Cbf1 and Tye7 (Zhu et al., 2009) to search for DNA sites containing two consecutive, overlapping 8-mers with E-scores >0.35 (Busser et al., 2012). Next, we selected 30 bp genomic regions centered at the putative binding sites to create a set of “ChiP-chip bound” probes for our gcPBM. Similarly, we created a set of “ChiP-chip unbound” probes by searching for putative Cbf1 and Tye7 binding sites in the genomic regions not bound in the ChiP-chip experiments.

For two proteins with identical specificities, we expect their in vitro DNA binding signals (here, the natural logarithm of the PBM fluorescence signal intensity) to be highly correlated. However, comparison of the in vitro DNA binding specificities of Cbf1 and Tye7 for their putative ChiP-chip bound sites (Figure 2D) clearly shows that the two TFs interact differently with these genomic sites. Importantly, even when we extend the comparison to include the ChiP-chip unbound probes, we observe the same trend (Figure 2E). Finally, although Cbf1 and Tye7 were tested at the same concentration (200 nM) on the array, Cbf1 bound with higher affinity to a larger number of probes than did Tye7. To ensure that the generally higher-affinity binding by Cbf1 is not the reason for the low correlation between in vitro DNA binding by these two TFs, we repeated the PBM experiment at a lower concentration of Cbf1 (100 nM). As expected, we saw a lower overall PBM signal for Cbf1, but the differences in DNA binding specificity between Cbf1 and Tye7 were maintained (Figure S1). In conclusion, our gcPBM data show that, despite having highly similar DNA binding motifs, the two TFs exhibit different binding preferences for their putative genomic binding sites.
Base Pairs Flanking the E-Box Binding Site Contribute to DNA Binding Specificity In Vitro

The DNA binding signal observed in our gcPBM experiments reflects the specificities of Cbf1 and Tye7 for E-box binding sites and their genomic flanks. Henceforth, we will refer to the two base pairs immediately upstream and downstream of the E-box as the “proximal flanks” and the base pairs more than two positions away from the E-box as the “distal flanks” (Figure 3A). Previous studies of bHLH DNA binding specificity focused either on the core E-box or the 2 bp proximal flanks (e.g., De Masi et al., 2011; Fong et al., 2012; Grove et al., 2009; Maerkl and Quake, 2007; Wang et al., 2012). Our analyses of the gcPBM data revealed that in addition to the E-box site and the proximal flanks, the distal flanks also contribute to the differential DNA binding specificities of Cbf1 and Tye7.

We first investigated whether the central two base pairs in the E-box binding sites are responsible for the different binding preferences. Analysis of the binding of these two TFs for all possible E-boxes revealed that the 2 bp central spacer does not appear to be the cause of the binding specificity differences, and as expected, both proteins have a strong preference for the E-box CACGTG (Figure S2). Thus, in our analyses of the gcPBM data, we focused primarily on genomic regions containing this E-box.

Our gcPBM data indicate that not all CACGTG sites across the genome are bound equally well by Tye7: depending on the flanking genomic regions, this E-box is bound in vitro with a wide range of affinities, ranging from highly preferential to nonspecific binding (Figure 3B). We observed a similar trend for Cbf1 (Figure S3). Even when we expanded the binding sites to include the 1 bp or 2 bp proximal flanks, we still observed wide variation in Cbf1 and Tye7 binding signal (Figures 3B, 3C, and S3), which indicates that the distal flanks contribute significantly to DNA binding specificity. Importantly, the wide range of binding affinities is not due to probes containing different numbers of binding sites because the probes comprise a single binding site located in the center of the probe (see Experimental Procedures). Thus, the differences in TF-DNA binding observed for probes that contain identical E-boxes and proximal flanks (e.g., ATCACGTGAA in Figure 3C) are due to contributions from the distal flanks.

Regression-Based Models Can Accurately Predict In Vitro DNA Binding of Cbf1 and Tye7

To understand what features in the genomic flanks contribute to the DNA binding specificities of Cbf1 and Tye7, we performed a regression analysis of the gcPBM data. We used support vector regression (SVR) (Drucker et al., 1997) to train linear models that use sequence features derived from the proximal and distal flanks to predict the DNA binding signal observed with gcPBMs (Figures 4A and 4B). Because both Cbf1 and Tye7 bind DNA as homodimers, and their E-box binding sites are palindromic, we combined the two flanking regions 5′ and 3′ of the E-box motif...
Figure 3. Flanking Sequences Contribute to Cbf1 and Tye7 DNA Binding Specificity

(A) Proximal or distal flanks surrounding the E-box (A) result in (B) variation in Tye7 DNA binding signal for probes that contain the preferred E-box CACGTG, or any of the possible 8-mers centered at this E-box. Numbers in parentheses indicate number of probes containing each 6-mer or 8-mer. (C) Wide variation in Tye7 DNA binding signal for probes that contain the preferred E-box CACGTG, or any of the possible 8-mers centered at this E-box. Numbers in parentheses indicate number of probes containing each 6-mer or 8-mer. The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines extend to 1.5 times the interquartile range from the box.

See also Figure S3.
SVR models accurately predict the individual DNA binding specificities of these TFs.

Next, to investigate how the various sequence features contribute to differences in DNA binding specificity between Cbf1 and Tye7, we compared the feature weights computed from the regression models for these TFs (Figure 4D). Although the two sets of weights are positively correlated ($R^2 = 0.32$), there are numerous differences between them, resulting from both proximal and distal flanks. For example, Tye7 disfavors the nucleotide C at position 4 (i.e., immediately downstream of the E-box), whereas Cbf1 actually prefers a C at this position (see feature “4-C” in the upper-left quadrant of Figure 4D). Unlike this difference, which is apparent in their DNA binding site motifs (Figure 1A), most differences in feature weights are subtle, in that they cannot be inferred from the motifs, and the individual contributions of the corresponding features are small. However, taken together, these features can accurately predict the different DNA binding specificities of Cbf1 and Tye7, as illustrated by the accuracy of the SVR models on both our initial gcPBM and the validation PBM. This suggests that the features represented by the distal flanks might not correspond to direct recognition by Cbf1 and Tye7 but, rather, might contribute to TF-DNA binding specificity indirectly by influencing the three-dimensional DNA structure. To further investigate this hypothesis, we performed a detailed DNA shape analysis of the sequences bound by Cbf1 and Tye7 in gcPBMs.

**Figure 4. Regression Analysis of gcPBM Data**

(A) For each 30 bp probe, we combined the two flanking regions, and we generated 1-mer, 2-mer, and 3-mer features. We used ε-SVR to train linear models that predict the PBM log signal intensity of each probe based on its sequence features. Positions are numbered starting from the center of the CACGTG core.

(B) Leave-one-out cross-validation analysis indicates that regression models for Cbf1 and Tye7 accurately predict PBM signal intensity.

(C) Analysis of the sequence features with the largest positive and negative weights in SVR models shows that base pairs in both the proximal and distal flanks are important for predicting DNA binding specificity. Bar plots show the top 20 positive and negative weights. For brevity, feature names are shown only for the top positive/negative weight and then for every other weight among the top 20.

(D) Features show numerous differences between Cbf1 and Tye7. See also Figure S4 and Table S1.

**DNA Shape Features Are Characteristic for bHLH Binding Sites**

We used a high-throughput (HT) DNA shape prediction approach (Slattery et al., 2011) to analyze differential DNA shape preferences selected by Cbf1 and Tye7 as a function of the in vitro binding signal (i.e., PBM log signal intensity). This DNA shape prediction method derives structural features of DNA (e.g., groove width and helical parameters) by mining Monte Carlo (MC) trajectories using a sliding pentamer window (see Experimental Procedures). Groove width in B-DNA is measured over a region of four base pairs and thus is affected by the sequence composition of at least half a helical turn (Rohs et al., 2005). In contrast, helical parameters describe DNA shape at dinucleotide resolution and give rise to groove geometry (Joshi et al., 2007). We analyzed both groove geometry and helical parameters.

Minor groove width and propeller twist (Figure 5A) and roll and helix twist (Figure S5) reflect the unique shape of E-boxes (CAnnTG), with minor groove widening at both CpA (TpG) base pair steps due to weak stacking interactions and the tendency of these dinucleotides (at positions -2/-3 and +2/+3) to open the minor groove. Propeller twist, roll, and helix twist further indicate a distinct conformation of the E-box. Our analysis of these features shows differences between high- and low-affinity binding. For example, minor groove width tends to be wider for high-compared to low-binding affinity sites, and propeller twist can distinguish binding preferences of Tye7 versus Cbf1 (Figure 5A).
Figure 5. DNA Shape Analysis
(A) Heatmaps show the average minor groove width (left) and propeller twist (right) for sequences derived from the gcPBM. Sequences were sorted in decreasing order of gcPBM signal intensity for either Cbf1 (top) or Tye7 (bottom) and grouped into 50 bins. Average DNA shape parameters were computed within each bin.
(B) Different proximal flanks surrounding the common CACGTG E-box are preferred by Tye7 and Cbf1. Sequences located in the upper-left triangle are preferentially bound by Tye7, and 10-mers located in the lower-right triangle are preferentially bound by Cbf1. Dashed lines indicate respective cutoffs of a difference of ≥30 in rank between Tye7 preferred (red) and Cbf1 preferred (blue). Lighter-colored dots exhibit larger differences.
(C) DNA shape variation due to flanks surrounding CACGTG selected preferentially by Cbf1 (light blue) or Tye7 (light red). Asterisks (*) indicate positions with significant differences (p < 0.05, Mann-Whitney U test) in the minor groove width (upper) or propeller twist (lower) between the sequences preferred by Cbf1 or Tye7. The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines show the range between the 5th and 95th percentiles. The symmetry of the box plots is due to the shape predictions having been performed for the combined flanks.
(D) Incorporating DNA shape features improves binding intensity predictions in comparison to using DNA sequence (1-mers) alone. The improvement is similar to that obtained by adding 2-mer and 3-mer features.

See also Figure S5.
Figure 6. Differences in the In Vitro DNA Binding Preferences of Cbf1 and Tye7 Are Important for Differential In Vivo Binding

(A) Overlap between sets of genomic regions bound by Cbf1 and Tye7 in ChIP-chip data in rich medium (YPD).

(B) Scatterplot of Tye7 versus Cbf1 PBM log signal intensity for 30-mer probes that occur in genomic regions bound in vivo only in Tye7, red, only in Cbf1 (blue), or in both data sets (gray).

(C) Cbf1 and Tye7 in vitro binding signal (i.e., natural logarithm of gcPBM probe intensity) for 30-mer probes selected from genomic regions bound only by Cbf1 (blue) or only by Tye7 (red) in vivo. The differences in PBM log signal intensity between the two sets of 30-mer probes are statistically significant by Kolmogorov-Smirnov (KS) tests. The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines extend to 1.5 times the interquartile range from the box.

See also Figure S6.

DNA Shape Features in Flanking Regions Are Distinct for Binding Sites Preferred by Cbf1 versus Tye7

Because our previous analysis of PBM data indicated that Tye7 and Cbf1 both bind preferentially to the E-box CACGTG (Zhu et al., 2009), we hypothesized that specificity for distinct binding sites arises from 5' and 3' flanking sequences. Therefore, we collected all the sequences from our gcPBM data that contained the E-box CACGTG, and then we compared the ranked log signal intensities for Tye7 and Cbf1 for these probes. Next we analyzed the groups of sequences bound preferentially either by Tye7 or Cbf1, defined as gcPBM probes with a difference ≥ 30 in rank between the two TF sets (shown by dashed lines in Figure 5B). Next, for both sets of sequences, we predicted DNA structural features and analyzed them for variation in DNA shape due to different flanks. We performed this analysis for both strands of the double helix and averaged the results because of the palindromicity of the CACGTG E-box. Our results indicate that both of these TFs select sites with distinct minor groove geometry (Mann-Whitney U test, p = 0.03, 0.008, 8.7 × 10⁻⁷, and 5.08 × 10⁻⁷ at positions 6, 4, 3, and 2, respectively) and propeller twist (p = 0.02, 0.01, 0.04, 0.02, and 1.1 × 10⁻³ at positions 9, 8, 7, 4, and 2) (Figure 5C). Differing flanking regions of the E-box (positions −3 to +3) being selected by Tye7 versus Cbf1 (Figure 5B). We observed similar statistically significant distinctions in roll (at dinucleotide positions 1/2, 3/4, 4/5, and 5/6) and helix twist (at dinucleotide positions 2/3, 3/4, and 10/11) (Figure S5).

Incorporation of DNA Shape Features Improves Binding Intensity Predictions in Comparison to Using DNA Sequence Alone

If DNA shape distinguishes binding targets selected by Cbf1 and Tye7, the use of structural features should also improve binding affinity predictions. To test this hypothesis, we incorporated structural features into our linear SVR approach. We found that adding DNA shape features (minor groove width, roll, propeller twist, and helix twist) leads to an improvement in binding specificity predictions similar to those obtained by adding 2-mer and 3-mer features: R² = 0.72 and 0.89 using 1-mer and DNA shape features (Figure 5D) compared to R² = 0.74 and 0.88 using 1-mer, 2-mer, and 3-mer features, for Cbf1 and Tye7, respectively (Figures 4B and 5D). Incorporating DNA shape features in addition to 2-mer and 3-mer features did not improve the prediction accuracy any further. This suggests that 2-mers and 3-mers implicitly contain structural information, whereas DNA shape implicitly contains interdependencies between nucleotides at different positions of the binding site. Using structural features instead of 2-mer and 3-mer features has the advantage that the total number of features is much smaller, and thus, regression algorithms other than SVR can be used successfully to learn accurate models of DNA binding specificity. To illustrate this point, we used L2-regularized linear regression and obtained highly accurate predictions: R² = 0.7 and 0.87 for Cbf1 and Tye7, respectively, using 1-mer and DNA shape features (Figure 5D; Experimental Procedures).

Genomic Sequences Flanking the E-Box Motif Contribute to Explaining the Differences in In Vivo DNA Binding between Cbf1 and Tye7

Both our regression analysis based on DNA sequence features and our DNA shape analysis show that Cbf1 and Tye7 interact differently with their putative genomic binding sites. To assess whether these differences contribute to differential DNA binding by these two TFs in vivo, we examined whether the DNA sequences preferred in vivo by a particular TF also have higher TF binding signal in vitro (Figure 6). Figure 6B shows a scatterplot of Cbf1 versus Tye7 in vitro binding signal for the 30-mer PBM probes selected from genomic regions bound in vivo by either of the two TFs (Harbison et al., 2004). We colored the data points based on in vivo specificity: blue for PBM probes selected from the 37 regions bound only by Cbf1 in vivo, red for PBM probes selected from the 67 regions bound only by Tye7 in vivo, and gray for PBM probes selected from the 11 genomic
regions bound by both Cbf1 and Tye7 in vivo. Next, for each TF, we compared the in vitro signal for PBM probes bound uniquely by only one TF in vivo (i.e., blue versus red data points) and found that DNA sequences preferred in vivo by a particular TF also have higher binding signal for that TF in vitro (Figure 6C) (Kolmogorov-Smirnov p = 0.00078 for Cbf1 and 0.003 for Tye7). We performed a similar analysis focusing on the PBM probes containing the E-box CACGTG and observed the same trend (Figure S6; Extended Experimental Procedures). Our results suggest that subtle differences in the intrinsic sequence preferences of Cbf1 and Tye7 observed in vitro on gcPBMs partially explain differential DNA binding in vivo observed in ChIP-chip data.

**DISCUSSION**

This study shows that subtle differences in the intrinsic preferences of paralogous TFs for sequences flanking the core DNA binding site motif can contribute to differential DNA binding in vivo. Using the *S. cerevisiae* TFs Cbf1 and Tye7 as our model system, we show that, when tested in vitro in their native genomic flanking sequences, putative DNA binding sites of Cbf1 and Tye7 are bound differentially by the two proteins. As expected, the differences between the intrinsic sequence preferences of the two TFs observed in vitro on our gcPBMs do not fully explain the differences in in vivo DNA binding observed in ChIP-chip data (Harbison et al., 2004). Other mechanisms might be used in vivo to provide additional specificity. For example, Cbf1 interacts with Met4 and Met28 to regulate genes involved in sulfur metabolism (Lee et al., 2010; Siggers et al., 2011). In addition, Cbf1 has chromatin-remodeling properties (Kent et al., 2004) that may allow it to bind certain CACGTG sites that are inaccessible for Tye7 due to nucleosome occupancy. However, to fully understand how these different mechanisms are used, it is important to have a better characterization of the intrinsic sequence preferences of the two TFs.

The analyzed structural features characterize free DNA (i.e., DNA not bound by the proteins) and thus reflect the intrinsic properties of the E-box binding sites and their genomic sequence context. Analysis of DNA shape shows that a widening of the minor groove characterizes the E-box in its unbound state, as we observed for sites selected by Tye7 and Cbf1. The same observation was made for the crystal structures of E-boxes in complex with the yeast TF Pho4 (Shimizu et al., 1997) and mammalian bHLH TFs (Brownlie et al., 1997; Ma et al., 1994). This suggests that DNA shape features observed in complexes of bHLH factors and their DNA targets are inherent to DNA binding sites and thus may constitute previously underappreciated, widely used signals in *cis*-regulatory sequences recognized by TFs. This form of intrinsic DNA shape recognition was previously observed for Hox proteins (Joshi et al., 2007; Slattery et al., 2011) and other TFs (Rohs et al., 2009). In addition to reporting this observation for E-box binding sites, we show here that structural variations due to different flanking sequences of E-boxes are a source of differences in DNA binding specificity among bHLH TFs. Consequently, we demonstrate that the integration of DNA shape and sequence leads to improved binding intensity predictions, similar to the use of 2-mers and 3-mers, compared to sequence (1-mers) alone.

In this study, we expressed both TFs as full-length proteins, so residues within or outside the DNA binding domain may play a role in the protein–DNA interactions. bHLH factors are known to select the E-box CAnnTG through DNA contacts by their His5 and Glu9 residues from each monomer of the bHLH dimers, which recognize the CpA (TpG) base pair steps (Shimizu et al., 1997). Based on cocrystal structures of a human bHLH factor and the yeast factor Pho4 bound to DNA (Shimizu et al., 1997), modeling, and mutagenesis studies, we showed previously that the Arg13 side chains of bHLH dimers select C/G base pairs at the two central positions of the CACGTG E-box through the formation of base-specific hydrogen bonds with the guanine bases at positions −1 and +1 (De Masi et al., 2011). Because the yeast bHLH factors Tye7, Cbf1, and Pho4 all have His5, Glu9, and Arg13 residues, the CACGTG motif is the E-box that is most preferred by all of these TFs. However, the reason why Tye7, Cbf1, and Pho4 prefer different sequences flanking the common E-box motif CACGTG is likely due to the length and sequence variation of the loop that separates the H1 and H2 helices in the bHLH protein (Figure 7). Cocrystal structures are not available for either Cbf1 or Tye7 bound to DNA, but crystal structures of Pho4 (Shimizu et al., 1997) and the human homolog of Cbf1, the upstream stimulatory factor (USF), have been solved in complex with DNA (Ferré-D’Amaré et al., 1994). The crystal structures of Pho4 and USF bound to DNA illustrate that the conformations of the respective loops between the H1 and H2 helices in both bHLH monomers can give rise to different DNA recognition in the regions flanking the E-box. The two loops of the Pho4 homodimer each form an additional 3 helix, whereas the USF loops are fully extended (Figure 7). Although base-specific contacts by bHLH factors are restricted to the E-box, the extended loops of both USF monomers lead to phosphate and other nonspecific contacts further upstream and downstream from the E-box, which can also be detected in DNase I footprints (Hesselberth et al., 2009; Neph et al., 2012). We suggest that these additional contacts outside the E-box may result in the selection of different flanking sequences through DNA shape features. In addition, structural differences in the flanking regions affect the ability of DNA to deform upon protein binding in order to optimize bHLH-DNA contacts and protein-protein interactions within the bHLH dimer.

In summary, our combined experimental and computational analysis of DNA sequence and shape preferences of yeast bHLH factors demonstrates that Cbf1 and Tye7 share the same E-box as a result of highly specific base contacts in the major groove, whereas they prefer different DNA flanking sequences because of structural features that enhance bHLH loop-DNA phosphate contacts that optimize the induced fit within the complex. Thus, this study demonstrates that bHLH factors use a combination of two different mechanisms of protein-DNA recognition: “base readout” and “shape readout” (Harris et al., 2012; Rohs et al., 2010); base readout in the major groove conserves the E-box, whereas local DNA shape readout in the flanking regions appears to enable distinct DNA binding preferences among paralogous TFs. It will be interesting to investigate if other TF families utilize DNA shape readout in similar ways because this could be an important mechanism...
through which closely related TFs recognize different DNA target sites and perform different regulatory roles in the cell.

**EXPERIMENTAL PROCEDURES**

**Enrichment of DNA Binding Site Motifs in ChIP-Chip Data**

Using Cbf1 and Tye7 DNA binding motifs derived from both in vivo (ChIP-chip) (MacIsaac et al., 2006) and in vitro (PBM) (Zhu et al., 2009) data, we computed the AUC enrichment, as described previously (Gordan et al., 2009), for each motif in the ChIP-chip data sets Cbf1_YPD and Tye7_YPD, which correspond to Cbf1 and Tye7, respectively, tested in rich medium, yeast peptone dextrose (YPD) (Harbison et al., 2004). In brief, from each ChIP-chip data set, we selected the “bound” and “unbound” probes, defined as probes with p < 0.005 and p > 0.5, respectively. Next, for each probe, we computed the probability of it being bound by a TF with a particular DNA motif. We used the scores for the bound and unbound probes to generate an ROC curve and took the AUC as a measure of enrichment of the motif in the ChIP-chip data.

**Protein Expression and Purification**

GST-Cbf1 and GST-Tye7 (Zhu et al., 2009) were overexpressed in E. coli BL21 (DE3) cells (New England BioLabs) and purified by FPLC (ÄKTAprime plus) using GSTrap FF affinity columns (GE Healthcare). Anti-GST western blots were performed to assess protein quality and concentration. See Extended Experimental Procedures for further details.

**gcPBM Design**

We designed a custom oligonucleotide array in 4x44K format (Agilent Technologies; AMADID #029393) containing putative Cbf1 and Tye7 DNA binding sites. Briefly, we represent three categories of 30 bp genomic sequences on our gcPBM: (1) ChIP-chip bound probes, (2) ChIP-chip unbound probes, and (3) negative control probes. ChIP-chip bound probes corresponded to genomic regions bound in vivo by Cbf1 or Tye7 (ChIP-chip p < 0.005 in rich medium [YPD]; Harbison et al., 2004) and containing at least two consecutive 8-mers with universal PBM E-scores >0.35 (Zhu et al., 2009). All putative binding sites occurred at the same position within the probes on the array. ChIP-chip unbound probes corresponded to genomic regions with ChIP-chip p > 0.5 and at least two consecutive 8-mers at a more stringent universal PBM E-score threshold of 0.4. Negative control probes corresponded to S. cerevisiae intergenic regions with a maximum 8-mer E-score of <0.3. We also designed probes that contain, within constant flanking regions, all 10 bp sequences that occur within the ChIP-chip bound probes and contain the E-box CACGTG but are flanked by synthetic rather than native genomic sequence. The reported PBM signal intensity for each probe is the median PBM signal intensity over four replicate spots. The validation array (Agilent Technologies; AMADID #041711) contains 30 bp genomic sequences from our initial custom array, with zero through four mutations designed at various positions in the genomic sequences. Details are provided in Extended Experimental Procedures.

**PBM Experiments and Data Analysis**

Custom-designed arrays were synthesized (AMADID #029393 and #041711), converted to double-stranded DNA arrays by primer extension, and used in PBM experiments essentially as described previously (Berger and Bulyk, 2009; Berger et al., 2006). PBM data quantification was performed as previously described (Berger and Bulyk, 2009; Berger et al., 2006). See Extended Experimental Procedures for details.

**SVR Analysis**

SVR was trained separately for Cbf1 and Tye7. For each TF, we first selected ChIP-chip bound and ChIP-chip unbound probes centered at the E-box CACGTG. To ensure that no additional binding sites occur in the regions

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**Figure 7. Sequence and Structure Comparison of bHLH TF-DNA Complexes**

(A) Assignment of secondary structure elements of S. cerevisiae Tye7, Cbf1, and Pho4, and human USF shows the sequence and length variation of the loops between α helices H1 and H2. The helical regions were either derived from crystal structures (Pho4 and USF) (Shimizu et al., 1997; Ferre-D’Amare et al., 1994) or predicted from amino acid sequence (Cbf1 and Tye7) (Cole et al., 2008). (B and C) In complex with their target sites, (B) yeast Pho4 and (C) human USF form base-specific contacts with the E-box, whereas the loops between the H1 and H2 helices of the bHLH motifs adopt different conformations. The Pho4 loop regions form additional short α helices, whereas the USF loops are fully extended. The bHLH TF-DNA complexes shown are based on crystal structures with Protein Data Bank IDs 1A0A (B) and 1AN4 (C).
flanking CACGTG, we selected probes (280 for Cbf1 and 312 for Tye7) for which the maximum PBM 8-mer E-score in the flanks was <0.3. Next, for each selected sequence, we computed the number of occurrences of each 1-mer, 2-mer, and 3-mer in the combined flanks (Figure 4A), or the corresponding DNA shape features. We thus obtained sparse feature matrices for each of the two TFs. As target features for the SVR analyses, we used the natural logarithm of the Cbf1 and Tye7 PBM fluorescence signal intensities. We used the 
\text{-}\text{SVR} algorithm implemented in the LIBSVM toolkit (Chang and Lin, 2011) for all SVR analyses. We performed a grid search using 10-fold and leave-one-out cross-validation to determine the best values for parameters c and C (see Extended Experimental Procedures). Using these parameters, we trained the final SVR models using all 280 sequences for Cbf1 and all 312 sequences for Tye7 and used them to predict the PBM log signal intensities for all probes on the validation array. We also performed an SVR analysis using the 312 sequences selected for Tye7 but shuffling the PBM log signal intensities; the best R² on randomized sets of sequences was <0.1 (Figure S4A).

**High-Throughput DNA Shape Prediction**

DNA shape parameters were derived from a high-throughput (HT) prediction approach (Slattery et al., 2011) based on mining data from Monte Carlo (MC) simulations (Joshi et al., 2007; Rohs et al., 2005) of 2,121 different DNA fragments. These MC simulations were analyzed with CURVES (Lavery and Sklenar, 1989) to calculate average minor groove width and helical parameters as a function of sequence. The resulting structural features were used to describe the average conformation of each of the 512 unique pentamers. The average conformation at the central base pair (for groove width and propeller twist) or the two central base pair steps (for roll and helix twist) of each unique pentamer was used to characterize a pentamer. A query table for pentamers was assembled using these data, and a sliding pentamer window was implemented to compute structural features for any DNA sequence. We validated our HT method for DNA shape predictions based on a comparison with all crystal structures of protein-DNA complexes available in the Protein Data Bank with a DNA duplex of at least one helical turn (10 bp) and no chemical modifications as specified elsewhere (Bishop et al., 2011). Spearman’s rank correlation coefficients are 0.67 for minor groove width, 0.55 for propeller twist, 0.63 for roll, and 0.54 for helix twist. Comparison with solution-state NMR structures of the Dickerson dodecamer in its unbound state using residual dipolar coupling (Wu et al., 2003) yields excellent quantitative agreement with our predictions for the four discussed parameters, with Pearson correlation coefficients of 0.84 for minor groove width, 0.79 for propeller twist, 0.93 for roll, and 0.49 for helix twist.

**Statistical Analysis of DNA Shape Parameters**

For Cbf1 and Tye7 separately, the selected sequences were grouped into 50 bins according to their ranked natural log signal intensity from gcPBM data. To extract the effect of the flanking sequences, the probes were filtered by the criterion of sharing the E-box motif CACGTG. The signal intensity ranks for all those probes were compared, and flanks bound preferentially by Tye7 or Cbf1 were identified as a difference ≥30 in rank between the two TFs (Figure 5B). The statistical significance of differences in the predicted groove width and helical parameters of these two distinct groups at each position was determined by the Mann-Whitney U test.

**Regularized Linear Regression Analysis Using DNA Sequence and Shape Features**

We trained L2-regularized linear regression models using sequence (1-mer) features alone or in combination with DNA shape features (minor groove width, roll, propeller twist, and helix twist). A 10-fold cross-validation was performed to assess their performance. In each round of cross-validation, the optimal regularization parameter λ was selected using an embedded 10-fold cross-validation on the training data set.

**ACCESSION NUMBERS**

The PBM data reported in this paper have been deposited in the Gene Expression Omnibus under accession number GSE44604.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, six figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2013.03.014.

**LICENSING INFORMATION**

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**EXTENDED EXPERIMENTAL PROCEDURES**

### Protein Expression and Purification

Full-length CBFI and TYE7 open reading frames cloned into the Gateway pDEST15 (N-terminal GST-tag) expression vectors (Invitrogen) were obtained from (Zhu et al., 2009).

GST-Cbf1 and GST-Tye7 were overexpressed in *E. coli* BL21 (DE3) cells (New England Biolabs) and purified by FPLC (AKTAprime plus) using 1 ml GSTrap^TM^ FF affinity columns (GE Healthcare). Samples were concentrated by centrifugation using Amicon Ultra filters (Millipore). Glycerol was added to a final concentration of 10%, and proteins were stored at −80 °C until further use. Western blots were performed for each protein to assess quality and to approximate protein concentration by visual inspection relative to a dilution series of recombinant GST standard (Sigma), as described previously (Zhu et al., 2009).

### gcPBM Design

We designed a custom DNA microarray (Agilent Technologies, Inc.; AMADID #029393) containing putative Cbf1 and Tye7 DNA binding sites within ~44,000 probes. The probes are 60 bp long and have the general form CAT(N)_{30}TTCGTCTTGATT CGCTTGACGTGCTG, where the last 24 nucleotides are complementary to a common primer used to double-strand the oligonucleotide array by primer extension (Berger et al., 2006) and (N)_{30} corresponds to 30-bp sequences from the *S. cerevisiae* genome. The constant 3-bp array sequences immediately flanking the genomic sequences (CAT and TTC) were selected so that they do not inadvertently create putative Cbf1 or Tye7 binding sites outside of the selected genomic regions. We chose to represent three categories of 30-bp genomic sequences on our gcPBM: 1) “ChIP-chip bound” probes, 2) “ChIP-chip unbound” probes, and 3) negative control probes. To generate the “ChIP-chip bound” probes, we first identified the genomic regions bound in vivo by Cbf1 or Tye7 (i.e., sequences with ChIP-chip p < 0.05 in rich medium (YPD) (Harbison et al., 2004)), and then searched those sequences for putative TF binding sites, defined as 10-bp sites that contain at least two consecutive 8-mers with PBM E-score > 0.35 according to the Cbf1 or Tye7 universal PBM data reported by (Zhu et al., 2009). These 10-bp sites were aligned to 10-bp long Cbf1 and Tye7 DNA binding site motifs (derived from the DNA motifs of (Zhu et al., 2009)) that contained the E-box site and 2-bp flanking regions; this ensured that all putative binding sites occurred at the same location within the probes on the microarray. Finally, we considered the 30-bp genomic sequences centered at the putative Cbf1 or Tye7 binding sites. The “ChIP-chip unbound” probes were selected similarly to “ChIP-chip bound” probes, except that: (1) they were based on genomic regions with ChIP-chip p > 0.5 (i.e., not bound significantly), and (2) we required at least two consecutive 8-mers at a more stringent E-score cutoff of 0.4. The negative control probes were selected from *S. cerevisiae* intergenic regions, with the criterion that the maximum E-score over all 8-mers for Cbf1 or Tye7 in each negative control probe was < 0.3, to ensure that these probes do not contain any putative Cbf1 or Tye7 binding sites. In addition to the three categories of 30-bp genomic sequences (i.e., “ChIP-chip bound” probes, “ChIP-chip unbound” probes, and negative control probes), we also designed probes that contain, within constant flanking regions, all 10-bp sequences that satisfy the following criteria: 1) they occur within the “ChIP-chip bound” probes, and 2) they contain the E-box CACGTG. These probes have the form CCTAGTCTGATGCTAGCTG (followed by the priming sequence GCCTTAGTCGGAGTACGCTG), and were specifically designed to compare binding of Cbf1 and Tye7 to putative 10-bp binding sites when these sites do not occur within native genomic flanks (Figure 6C). Each probe is represented at 4 separate spots on the custom array. The reported PBM signal intensity for each probe is the median over the PBM signal intensity values over the 4 replicate spots. To design the “validation” array (Agilent Technologies, Inc.; AMADID #041711), we selected 30-bp genomic sequences from our initial custom array, and introduced 1-bp, 2-bp, 3-bp, or 4-bp mutations at various positions in the original genomic sequences; both the wild-type sequences and the resulting mutant sequences are represented on the validation array.

### PBM Experiments and Data analysis

Custom-designed microarrays were synthesized (Agilent Technologies, AMADID #029393 and #041711), converted to double-stranded DNA arrays by primer extension, and used in PBM experiments essentially as described previously (Berger et al., 2006; Berger and Bulyk, 2009). All PBM data reported in this study are from experiments performed either on a fresh slide or a slide that had been stripped exactly once (Berger et al., 2008). Microarray scanning and quantification were performed using ScanArray (Perkin Elmer) for AMADID #029393, and GenePix 4400A (Molecular Devices) for AMADID #041711. In both cases data normalization was performed using masliner (Microarray LINEar Regression) (Dudley et al., 2002) and the Universal PBM Data analysis Suite (Berger and Bulyk, 2009) as previously described (Berger et al., 2006; Berger and Bulyk, 2009).

### SVR Analysis

Support Vector Regression (SVR) was run separately for Cbf1 and Tye7. For each TF, we first selected “ChIP-chip bound” and “ChIP-chip unbound” probes centered at the E-box CACGTG. To ensure that no additional binding sites occur in the regions flanking CACGTG, we selected probes (280 for Cbf1, and 312 for Tye7) for which the maximum E-score over all the 8-mers in the flanks was < 0.3. Next, for each selected sequence we combined the two flanking regions (see Figure 4A) and computed the number of occurrences of each 1-mer, 2-mer and 3-mer in the combined flanks. For example, for a given 30-bp sequence, the feature “6-TTT” will be 0 if TTT does not occur at position 6 in either of the flanks, 1 if TTT occurs at position 6 in one of the flanks, and...
2 if TTT occurs at position 6 in both flanks. The positions are numbered starting from the center of the CACGTG core (Figure 4A). We thus obtained a sparse feature matrix for each of the two TFs. As target features for the SVR analyses, we used the natural logarithm of the Cbf1 and Tye7 PBM fluorescence signal intensities. Our goal was to use SVR to train linear models that can predict the PBM log signal intensity for each probe based on the sequence features derived from that probe. We used the ε-SVR algorithm implemented in the libSVM toolkit (Chang and Lin, 2011) for all SVR analyses. We performed a grid search using 10-fold and leave-one-out cross-validation to determine the best values for parameters C and ε: C = 0.005 and ε = 0.1 for Cbf1; C = 0.1 and ε = 0.35 for Tye7 (see below). Using these parameters, we trained the final SVR models using all 280 sequences for Cbf1 and all 312 sequences for Tye7. These models were used to predict the PBM log signal intensities for all the probes on the “validation” array. We also performed an SVR analysis using the 312 sequences selected for Tye7, but shuffling the PBM log signal intensities, to determine the range of R² values we would obtain on randomized sets of sequences; the best R² on randomized sets of sequences was < 0.1 (Figure S4A).

Parameter Search in the SVR Analyses

The ε-SVR implementation in the libSVM package (Chang and Lin, 2011) was used for all SVR analyses. Analyses were run separately for Cbf1 and Tye7, using different sets of features as input. For each transcription factor (TF) and each set of input features, we selected the best values for parameters C and ε by doing a grid search and using cross-validation to evaluate each set of parameters. We started with 10-fold cross validation and tried a wide range of parameters for C (0.0001, 0.00025, 0.0005, 0.00075, 0.001, 0.0025, 0.005, 0.0075, 0.01, 0.025, 0.05, 0.075, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, 0.4, 0.45, 0.5, 0.55, 0.6, 0.65, 0.7, 0.75, 0.8, 0.85, 0.9, 0.95, 1, 1.05, 1.1, 1.15, 1.2, 1.25, 1.3, 1.35, 1.4, 1.45, 1.5). Next, we narrowed down the parameter range for C (0.001, 0.0025, 0.005, 0.0075, 0.01, 0.025, 0.05, 0.075, 0.1, 0.25, 0.5, 0.75, 1, 2.5, 5, 7.5, 10, 25, 50, 75, 100) and ε (0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, 0.4, 0.45, 0.5) and used leave-one-out cross-validation to find the best parameter values within these sets. After obtaining the best parameter values, we ran ε-SVR again with the selected parameters using leave-one-out cross-validation, and this time we stored the predicted gcPBM log signal intensity for every input sequence and we calculated the average weights of all the features in the regression model.

For both Cbf1 and Tye7, we tested SVR models using: 1) 1-mer features, 2) 1-mer and 2-mer features, 3) 1-mer, 2-mer and 3-mer features, and 4) 1-mer, 2-mer, 3-mer and 4-mer features. In a 10-fold cross-validation test, using the 280 Cbf1 sequences (see main text) and the parameters optimized using the grid search described above, we obtained R² values of: 0.60, 0.67, 0.74, and 0.71 for the four models, respectively. For the 312 Tye7 sequences, the R² values were 0.77, 0.78, 0.88, and 0.86 for the four models, respectively. Thus, the models using 1-mers, 2-mers and 3-mers performed best. The fact that the models using 4-mers did not improve the prediction accuracy is not surprising given that they have a very large number of features compared to training examples, and are thus prone to overfitting the training data.

Design of Validation PBM

To test the accuracy of the regression models trained on sequences from the gcPBM, we introduced mutations at various positions in both the proximal and the distal flanks of the 30-bp gcPBM probes, and generated a “validation” PBM containing three categories of probes: 1) “feature mutations” probes, 2) “flank mutations” probes, and 3) negative control probes.

To generate the “feature mutations” probes, we focused on the sequence features that had large positive or negative weights in the Cbf1 or Tye7 regression models. Since we can only fit a limited number of probes on a custom PBM, we sorted the features in decreasing order of their SVR weights and we selected: 1) proximal features (i.e., features that start at position 4 or 5) that are among the top 10 or bottom 10 features, and 2) distal features (i.e., features that start at a position > 5) that are among the top 20 and bottom 20 features. Next, for each feature we selected the sequences for which the weight was > 0. Features for which no such sequences were found were removed from the analysis. If the number of corresponding sequences for a selected feature was larger than 10, we selected 10 of those sequences randomly, otherwise we considered all sequences for that feature. Finally, we made point mutations for the 1-bp features and double mutations for the 2-bp features selected as described above, and added both the wild-type and the mutant sequences to the microarray design. The results of this analysis are presented in Figure S4F, as squared Pearson correlation coefficients (R²) between the predicted and the measured PBM log signal intensities over the probes corresponding to each selected feature. The high R² values (between 0.68 and 0.97 for Cbf1, and 0.64 and 0.92 for Tye7) show that our models are accurate at predicting the PBM log signal intensities. In addition to evaluating our predictions using the Pearson correlation coefficient, we also computed, for each selected feature, the fraction of mutated probes for which we correctly predicted whether the mutation will cause an increase or decrease in PBM signal intensity compared to wild-type sequence. As shown in Figure S4F, these numbers vary from 83% and 100% for features selected according to the Cbf1 model, and from 67% to 99% for features selected according to the Tye7 model. These validation results can be used in the future to improve the regression models by focusing on the sequences with the largest prediction errors.

To generate the “flank mutations” probes, we selected 5 genomic regions bound either by Cbf1 or Tye7 in vivo and generated all possible mutations of their 4-bp proximal flanks. The selected sequences are:

1. CATGTCCGGCCTACGTGGTTATGATGTAAG,
2. ATTTGTCACTAGTCAGTGATTTACATTTAAG,
3. ACGCGTTCGCTTCAGTGATGTTCTTTTC.
Analyses of the “flank mutations” probes are presented in Figures S4C–S4E. Similarly to the initial gcPBM, the “validation” microarray contains negative control probes selected so that they do not contain any putative Cbf1 or Tye7 binding sites. The “validation” array contains a total of 237 negative control sequences, and each sequence is present in 4 different replicate spots. All the other sequences on the “validation” array are present in 6 different replicate spots.

After running the PBM experiment on the “validation” array, we compared the PBM log intensity values for the wild-type sequences against the PBM log intensity values obtained for the same sequences in the initial gcPBM experiment, and we adjusted the PBM log intensity on the “validation” array to be in the same range as the values on the original array.

Statistical Analysis of the Differences in gcPBM Signal for 30-mer Probes Bound by Cbf1 versus Tye7 In Vivo

To assess whether differences in the in vitro binding preferences of Cbf1 and Tye7 contribute to differential DNA binding by these TFs in vivo, we examined whether the sequences preferred in vivo by each TF also have higher TF binding signal in vitro (Figure 6). As shown in Figure 6B, we first separated the gcPBM 30-mer probes based on in vivo specificity: blue for 30-mer probes selected from the 37 regions bound only by Cbf1 in vivo, and red for 30-mer probes selected from the 67 regions bound only by Tye7 in vivo (Figure 6B also shows, in gray, 30-mer probes selected from the 11 genomic regions bound by both Cbf1 and Tye7 in vivo; these probes were not used in the statistical analyses described here). Next, for each TF we filtered out the low-affinity gcPBM probes, defined as probes with log signal intensity less than two standard deviations above the mean signal observed for negative control gcPBM probes. We eliminated these probes from the analysis because they are likely to be bound nonspecifically by the TFs. Using the specifically-bound gcPBM probes, we next compared the in vitro (gcPBM) signal for probes bound only by Cbf1 in vivo versus probes bound only by Tye7 in vivo (Figures 6B and 6C). We used the Kolmogorov-Smirnov test to determine whether the differences in in vitro binding signal between the two sets of probes (i.e., red versus blue points in Figures 6B and 6C) are statistically significant.

SUPPLEMENTAL REFERENCES


Figure S1. Comparison of In Vitro Binding of Cbf1 versus Tye7 to ChIP-chip Bound and ChIP-chip Unbound Probes on gcPBMs, Related to Figure 2
Concentrations of Cbf1 or Tye7 in the PBM binding reactions were 100 nM and 200 nM, respectively. The plots show the natural logarithm of the normalized PBM fluorescence signal intensities, with higher numbers corresponding to higher affinity binding.
Figure S2. Universal PBM-Derived E-Box Specificities of Cbf1 and Tye7, Related to Figure 1

Box plots depict PBM E-scores (y-axis) for all 8-mers that contain a given E-box (x-axis). For each box plot, the central horizontal bar shows the median of the E-score distribution, the box’s edges mark the 25th and 75th percentiles, and the whiskers represent the most extreme points of the distribution which were not determined to be outliers. For convenience, we include a horizontal bar at an E-score value of 0.35, above which we consider 8-mers to correspond to specific TF-DNA binding (Berger et al., 2006; Gordan et al., 2011).
Figure S3. Variation in Cbf1-DNA Binding Signal due to Sequences Flanking the E-Box CACGTG, Related to Figure 3

(A) Variation in Cbf1-DNA binding signal for probes that contain the preferred E-box CACGTG (top box plot), or any of the possible 8-mers centered on this E-box (middle 10 box plots). The numbers in parentheses indicate how many probes contain each 6-mer or 8-mer, respectively (in either the forward or reverse complementary orientation). The negative control probes (bottom box plot) were chosen so that the maximum 8-mer E-score across the entire probe is < 0.3, i.e., these probes do not contain any putative Cbf1 binding sites. Thus, the distribution of gcPBM log signal intensities for the negative control probes corresponds to nonspecific Cbf1-DNA binding.

(B) We observe a wide variation in DNA binding signal even when we restrict the analysis to probes containing specific 10-mers (i.e., the E-box plus 5' and 3' proximal flanks). The box plot illustrates two examples of 10-mers showing a wide variation in Cbf1 binding signal. For comparison, we also show the variation among 4 replicate spots (bottom box plot) containing the same 30-bp sequence, which contains a TTCACGTGAT binding site (CCAGTGAAATTTCACGTGAT). The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines extend to 1.5 times the interquartile range from the box.
Figure S4. Performance of SVR Models, Related to Figure 4

(A) Distribution of $R^2$ values (squared Pearson correlation coefficient) for 100 randomized data sets derived from the Tye7 data set. The parameters for the support vector regression were optimized using the same parameter search as for the real data set. The (*) indicates the $R^2$ obtained for the real data set.

(B) Accuracy of SVR models trained on flanking regions of different lengths. The best accuracy was obtained when 11-bp flanks were used to predict Cbf1 binding, and when 5-bp flanks were used to predict Tye7 binding.

(C) One category of probes on the validation microarray ("flank mutations" probes) was derived from 5 genomic regions containing CACGTG sites. We kept the E-box and the distal flanks constant, and generated all possible sequences in the 4-bp proximal flanks.

(D) Table shows Pearson correlation $R^2$ between the predicted and observed gcPBM log signal intensities for probes containing mutations at various positions relative to the E-box (e.g., to compute the $R^2$ for mutations at position 4 we selected those sequences that match one of the wild-type sequences at all positions except position 4, with the mutation occurring in either of the two flanks). Predictions were made using SVR models derived for Cbf1 and Tye7 from the 12-bp flanking regions (see main text for details).

(E) Box plots show the variation in gcPBM log signal intensity for Cbf1 and Tye7 across the mutated sequences generated from each of the genomic sequences in (C). Each box corresponds to one of the 5 genomic regions. The wide range of gcPBM log signal intensities obtained for each sequence shows the influence of the 4-bp proximal flanks on DNA binding. The variation we observe among the medians of the five gcPBM log signal intensity distributions (corresponding to the 5 sequences) shows the influence of the distal flanks. The differences between the Cbf1 and Tye7 boxplots show that the two TFs interact differently with these DNA sequences. The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines extend to 1.5 times the interquartile range from the box.

(F) Performance of the Cbf1 and Tye7 SVR models on genomic sequences containing mutations at various positions in the flanking regions.
Figure S5. DNA Shape Analysis, Related to Figure 5

(A) Heat maps show the average roll and helix twist for the DNA sequences on the gcPBM. The sequences were sorted in decreasing order of the PBM signal intensity for either Cbf1 (left) or Tye7 (right), and grouped into 50 bins. Average DNA shape parameters were computed over the sequences in each bin.

(B) Box plots show the DNA shape variation due to flanks selected by Cbf1 (light blue) or Tye7 (light red). We compared the DNA shape parameters at individual positions between two groups of sequences that share the CACGTG E-box but are bound with higher affinity by either Cbf1 or Tye7 (see text for details). Asterisks (*) mark the positions at which we observed a significant difference (Mann-Whitney U p < 0.05) in roll or helix twist between the sequences preferred by the two TFs. The E-box is located at positions –3 to +3. The symmetry of the box plots is due to the shape predictions performed for both DNA strands, thus combining the left and right flanks.

The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines show the range between the 5th and 95th percentiles.
Figure S6. In Vitro Binding Differences in the Sequence Preferences of Cbf1 and Tye7 Partially Explain Differential In Vivo DNA Binding, Related to Figure 6

(A) Tye7 in vitro binding signal (gcPBM log signal intensity) for CACGTG-containing 30-mer probes that occur in Tye7_YPD (red), or in Cbf1_YPD but not in Tye7_YPD (blue). The difference in Tye7 PBM log signal intensity between the two sets of 30-mer probes is statistically significant (Kolmogorov-Smirnov (KS) p = 0.003612), with probes bound by Tye7 in vivo having an overall higher DNA binding signal in vitro. We observed a similar trend for Cbf1, with probes bound by Cbf1 in vivo having a higher Cbf1 PBM signal than probes not bound in vivo by Cbf1 (p = 0.04; not shown).

(B) As in (A), except that CACGTG-containing 10-mers were tested within constant 10-bp DNA flanks on the gcPBM. The difference in Tye7 PBM log signal intensity between the two sets of 10-mers is not statistically significant (KS p = 0.1696).

The boxes show the range between the 25th and 75th percentiles, the line within each box indicates the median, and the outer lines extend to 1.5 times the interquartile range from the box.