

# Determination and Inference of Eukaryotic Transcription Factor Sequence Specificity

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## SUMMARY

Transcription factor (TF) DNA sequence preferences direct their regulatory activity, but are currently known for only ~1% of eukaryotic TFs. Broadly sampling DNA-binding domain (DBD) types from multiple eukaryotic clades, we determined DNA sequence preferences for >1,000 TFs encompassing 54 different DBD classes from 131 diverse eukaryotes. We find that closely related DBDs almost always have very similar DNA sequence preferences, enabling inference of motifs for ~34% of the ~170,000 known or predicted eukaryotic TFs. Sequences matching both measured and inferred motifs are enriched in chromatin immunoprecipitation sequencing (ChIP-seq) peaks and upstream of transcription start sites in diverse eukaryotic lineages. SNPs defining expression quantitative trait loci in *Arabidopsis* promoters are also enriched for predicted TF binding sites. Importantly, our motif “library” can be used to identify specific TFs whose binding may be altered by human disease risk alleles.

These data present a powerful resource for mapping transcriptional networks across eukaryotes.

## INTRODUCTION

Transcription factor (TF) sequence specificities, typically represented as “motifs,” are the primary mechanism by which cells recognize genomic features and regulate genes. Eukaryotic genomes contain dozens to thousands of TFs encoding at least one of the >80 known types of sequence-specific DNA-binding domains (DBDs) (Weirauch and Hughes, 2011). Yet, even in well-studied organisms, many TFs have unknown DNA sequence preference (de Boer and Hughes, 2012; Zhu et al., 2011), and there are virtually no experimental DNA binding data for TFs in the vast majority of eukaryotes. Moreover, even for the best-studied classes of DBDs, accurate prediction of DNA sequence preferences remains very difficult (Christensen et al., 2012; Persikov and Singh, 2014), despite the fact that identification of “recognition codes” that relate amino acid (AA) sequences to preferred DNA sequences has been a long-standing goal in the study of TFs (De Masi et al., 2011; Desjarlais and Berg, 1992; Seeman et al., 1976). These deficits represent a fundamental limitation in our ability to

analyze and interpret the function and evolution of DNA sequences.

The sequence preferences of TFs can be characterized systematically both in vivo (Odom, 2011) and in vitro (Jolma and Taipale, 2011; Stormo and Zhao, 2010). The most prevalent method for in vivo analysis is currently chromatin immunoprecipitation sequencing (ChIP-seq) (Barski and Zhao, 2009; Park, 2009), but ChIP does not inherently measure relative preference of a TF to individual sequences and may not identify correct TF motifs due to complicating factors such as chromatin structure and partner proteins (Gordán et al., 2009; Li et al., 2011; Liu et al., 2006; Yan et al., 2013). In contrast, it is relatively straightforward to derive motifs from all of the common methods for in vitro analysis of TF sequence specificity, including protein binding microarrays (PBMs), bacterial 1-hybrid (B1H), and high-throughput in vitro selection (HT-SELEX) (Stormo and Zhao, 2010), all of which have been applied to hundreds of proteins (Berger et al., 2008; Enuameh et al., 2013; Jolma et al., 2013; Noyes et al., 2008).

Previous large-scale studies have reported that proteins with similar DBD sequences tend to bind very similar DNA sequences, even when they are from distantly related species (e.g., fly and human). This observation is important because it suggests that the sequence preferences of TFs may be broadly inferred from data for only a small subset of TFs (Alleyne et al., 2009; Berger et al., 2008; Bernard et al., 2012; Noyes et al., 2008). However, these analyses have utilized data for only a handful of DBD classes and species and they contrast with numerous demonstrations that mutation of one or a few critical DBD AAs can alter the sequence preferences of a TF (Aggarwal et al., 2010; Cook et al., 1994; De Masi et al., 2011; Mathias et al., 2001; Noyes et al., 2008), which suggest that prediction of DNA binding preferences by homology should be highly error-prone. To our knowledge, rigorous and exhaustive analyses of the accuracy and limitations of inference approaches to predicting TF DNA-binding motifs using DBD sequences has not been done.

Here, we determined the DNA sequence preferences for >1,000 carefully-selected TFs from 131 species, representing all major eukaryotic clades and encompassing 54 DBD classes. We show that, in general, sequence preferences can be accurately inferred by overall DBD AA identity, suggesting that mutations that dramatically impact sequence specificity are relatively rare. By identifying distinct confidence thresholds for each individual DBD class (i.e., levels of protein sequence identity above which motifs can be assumed to be identical between two proteins), we infer sequence preferences for roughly one-third of all known eukaryotic TFs, based on experimental data for fewer than 2% of them. Cross-validation indicates that ~89% of predicted sequence preferences are as accurate as experimental replicates of the same TF. We demonstrate the functional relevance and utility of both known and inferred motifs by showing that they coincide with ChIP-seq binding peak sequences, are enriched in the promoter regions of diverse eukaryotes, and significantly overlap eQTLs in *Arabidopsis*. We also demonstrate how our data can be used to predict the specific TFs whose binding would be altered by a human disease risk allele. To house the data and the resulting inferences, we have created the Cis-BP database (catalog of inferred sequence binding preferences), freely available at <http://cisbp.cabr.utoronto.ca>.

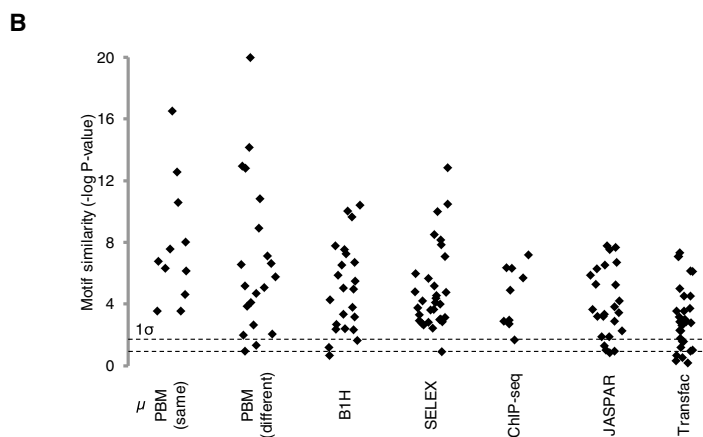
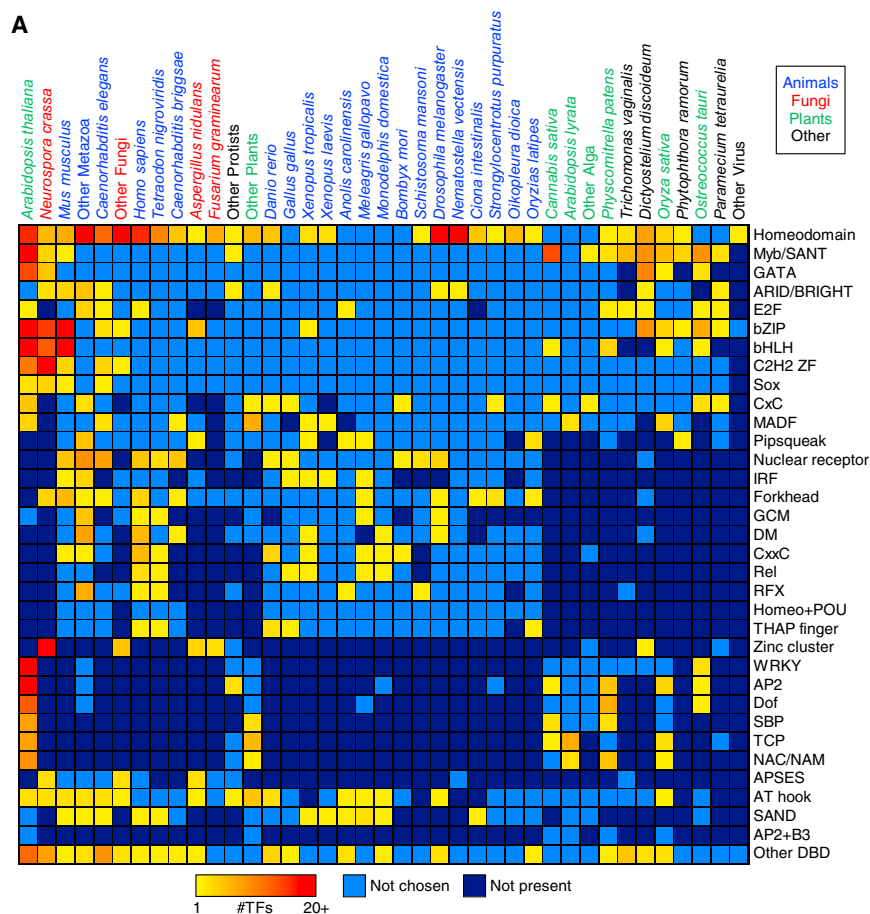
## RESULTS

### PBM Data for >1,000 Diverse Eukaryotic TFs

We sought to examine the relationship between the DBD AA sequence identity and the DNA sequence specificity of any two proteins and to simultaneously broadly survey the sequence specificity of eukaryotic TFs. To identify TFs, we first scanned the AA sequences for each of 81 different types of DBDs for which there is an available Pfam model (Weirauch and Hughes, 2011), using known or predicted proteins in 290 sequenced eukaryote genomes. We identified a total of 166,851 putative TFs that fit these criteria. From these, we selected 2,913 individual TFs to analyze, using several different criteria aimed at achieving the goals of our study, including a relatively even balance among DBD classes and species, a survey of different levels of sequence identity among proteins, and a deeper focus on several model organisms and abundant DBD classes (see Experimental Procedures). Figure S1, available online, depicts the overall scheme.

We analyzed each TF using PBM assays, following procedures previously described (Berger et al., 2006; Weirauch et al., 2013). The PBM technique can be summarized as follows: a GST-tagged DNA-binding protein is “hybridized” to a double-stranded DNA microarray, and subsequent addition of a fluorescently tagged antibody reveals the DNA sequences that the protein has bound and to what degree. Each PBM contains a diverse set of ~41,000 35-mer probes, designed such that all possible 10-mers are present once and only once; thus, all nonpalindromic 8-mers are present 32 times, allowing for a robust and unbiased assessment of sequence preference to all possible 8-mers. Values for individual 8-mers are typically given as both *E* scores (that represent relative rank of intensities and range from -0.5 to + 0.5) (Berger et al., 2006) and *Z* scores (that scale approximately with binding affinity) (Badis et al., 2009). PBMs also allow derivation of position weight matrices (PWMs) up to 14 bases wide (Badis et al., 2009; Berger et al., 2006; Mintseris and Eisen, 2006; Weirauch et al., 2013), as well as International Union of Pure and Applied Chemistry (IUPAC) consensus sequences (Ray et al., 2013). PWMs can be represented as sequence logos (Schneider and Stephens, 1990) that are typically taken as synonymous with “motifs.” Here, we report 8-mer scores, PWMs, and consensus sequences, and use whichever is best suited to individual analyses. For example, while motifs are well suited for visualization of sequence preferences, and convenient for scanning longer sequences, they can underestimate reproducibility of experiments due to the introduction of uncertainties in the process of PWM derivation (Weirauch et al., 2013; Zhao and Stormo, 2011).

We analyzed each of the 2,913 proteins using two different PBM arrays, designated “ME” and “HK” after their designers (Lam et al., 2011). Of these, 1,032 (encompassing 1,017 different TFs) yielded data that satisfy our stringent success criteria, including *E* scores >0.45 on both arrays, and agreement in both 8-mer data and motifs between the two arrays (Berger et al., 2008; Weirauch et al., 2013) (see Experimental Procedures). The distribution of the 1,032 proteins over 131 species and 54 DBD classes is summarized in Figure 1A. Many values in this matrix are zero because the majority of DBD classes are



only present in certain subsets of species (e.g., plant-specific TFs) (Weirauch and Hughes, 2011). PBM failures may be due to any of several causes, including protein misfolding, requirement for cofactors, or bona fide lack of sequence-specific DNA binding activity.

For 123 of the 1,017 examined TFs, there are previously described motifs. In most cases, the new motifs we obtained are highly similar to motifs compiled from the literature (JASPAR and Transfac), derived from PBMs in other studies, derived from

**Figure 1. Overview of the Motif Data Set**

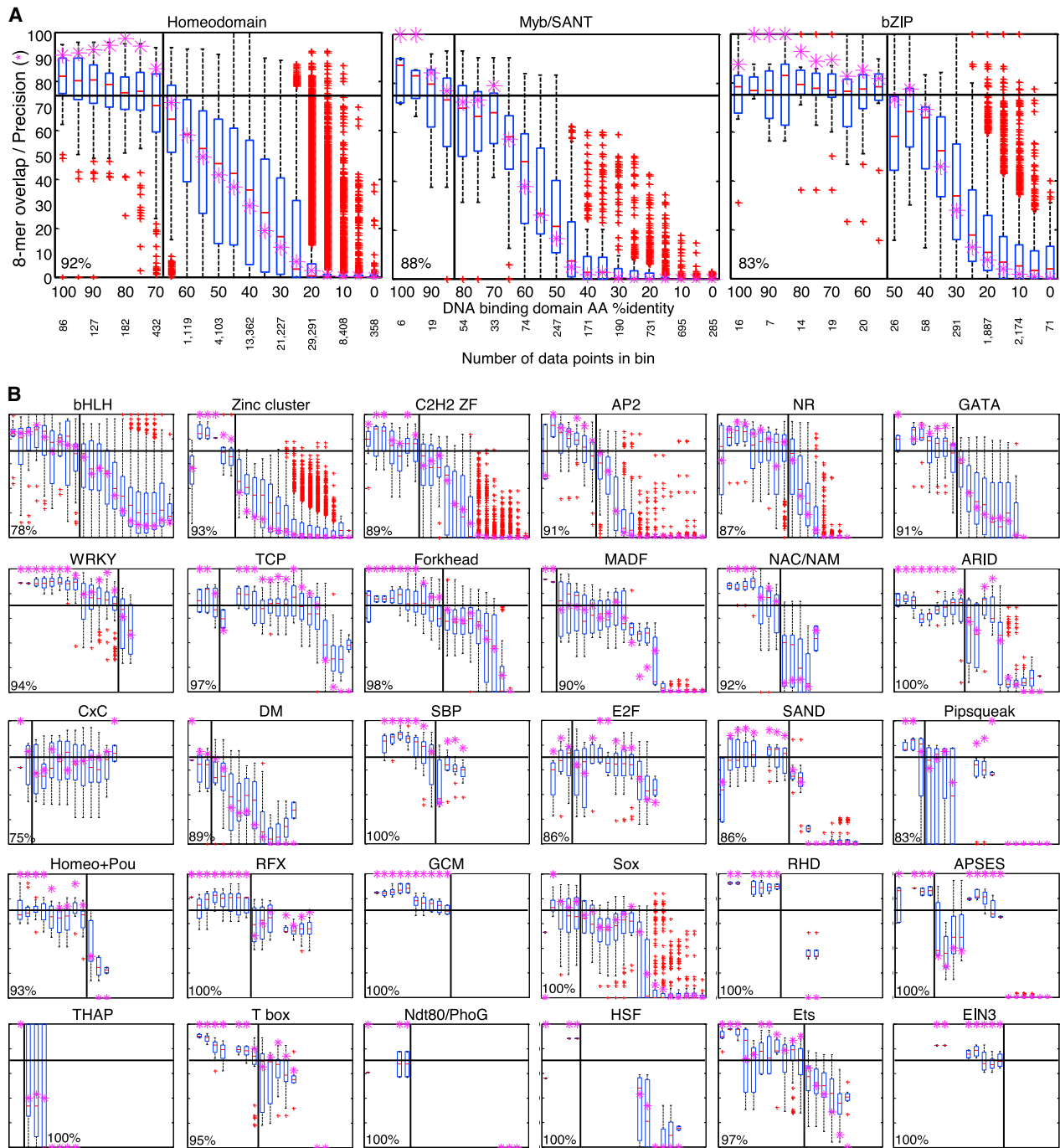
(A) TFs characterized in this study by species and DBD class. TFs with multiple DBD classes are indicated with a “+” (e.g., AP2+B3). DBD classes and species containing fewer than five members are grouped into “Other.” Species are ordered by the total number of TFs with characterized motifs. (B) PBM-derived motifs are similar to previously characterized motifs. We compared new PBM-derived motifs to previously determined motifs for the same TF. p values were calculated using the TomTom PWM similarity tool (Tanaka et al., 2011), with Euclidean distance and default parameter settings. Dashed lines indicate mean (bottom) and mean + 1 SD (top) of p values obtained from 10,000 randomly selected PWM pairs. “PBM (same)” and “PBM (dif)” indicate PBMs from other studies performed using the same, or different array designs as this study, respectively. See also Figure S1 and Tables S1, S2, and S6.

other technologies (B1H and HT-SELEX), or derived from ChIP-seq experiments (Figure 1B; sources provided as Table S1; full motif comparisons provided as Table S2). Importantly, a large majority of the proteins we analyzed (894/1,017) had no previous binding data, and among these, roughly half yielded a motif that is highly different from any previously known motif (see Experimental Procedures). For several DBD classes (CG-1, CxC, GRAS, LOB, and Storekeeper) we characterized sequence specificity for the first time. CxC domains, for example, span a wide range of organisms encompassing plants, animals, and protists and recognize variations of a unique, largely conserved TTTCGAAA motif.

### Inference of TF Sequence Specificity Using Degree of Identity in the DNA-Binding Domain

We next used the PBM data to ask how well the percent protein sequence identity in the DBD between two proteins correlates with similarity in their DNA sequence preferences. As a measure of DNA sequence preference similarity, we calcu-

lated the overlap in high-scoring 8-mers (e.g., *E* score >0.45) (see Experimental Procedures), as the *E* scores are on a uniform scale that facilitates direct comparison. The boxplots shown in Figure 2 and Data S1 illustrate that there are different characteristic relationships between DBD identity and 8-mer overlap for different DBD classes. However, virtually all of the plots display a sigmoidal appearance, such that a particular threshold defines the %AA identity over which the majority (75% or more) of protein pairs have very similar DNA binding preferences (i.e., are



**Figure 2. Motif Inference Thresholds by DBD Class**

(A) Relationship between similarity in DBD AA sequence and DNA sequence preferences. Boxplots depict the relationship between the %ID of aligned AAs and % of shared 8-mer DNA sequences with *E* scores exceeding 0.45, for the three DBD classes with the most PBMs in this study. Red lines indicate the median value of each bin. Edges of the blue boxes indicate the 25th and 75th percentiles. Whiskers extend to the most extreme data points not considered outliers. Red crosses indicate outliers. %ID bins range from 0 to 100, of size 10, in increments of five. Bottom: number of DBD pairs in each bin. Pink asterisks indicate the precision of the corresponding bin (i.e., the fraction of protein pairs with 8-mer similarity at least as high as the 25th percentile of replicates). Horizontal line indicates the 75% precision line used to choose the inference threshold. Vertical lines indicate AA %ID threshold (i.e., the point before the pink asterisks drop below the horizontal line). Percentage in lower left corner indicates cross validation success rate.

(B) Relationship for all DBD classes. Boxplots for all DBD classes for which we could establish an inference threshold, depicted as in (A). DBD classes are ordered by the number of TFs characterized in this study.

NR, nuclear receptor. See also [Figures S2 and S6](#).

at least as similar as the 25th percentile of replicates of the same protein—see [Experimental Procedures](#) ([Figure 2](#)). For homeodomains, this threshold is 70% AA identity ([Figure 2A](#)), almost exactly what we previously reported (65%) ([Berger et al., 2008](#)).

Strikingly, such a threshold can be drawn for virtually every DBD class ([Figure 2](#) and [Data S1](#)). Moreover, exactly the same trends and thresholds are observed in a leave-one-out cross validation prediction framework, in which the high-scoring 8-mers are predicted to be identical to those of the protein with the closest %AA identity (see [Experimental Procedures](#)). Using this framework, we estimate that ~89% of predicted sequence preferences (hereafter, “inferences”) above the thresholds shown in [Figure 2](#) are as accurate as experimental replicates ([Data S1](#)), with similar precision obtained whether inferences are derived from only orthologs (88%) or only paralogs (90%). The distribution of DBD AA similarities among the 1,017 proteins we analyzed is nearly identical to the distribution among all sequenced eukaryotes (data not shown), so the same numbers can be expected for de novo predictions. Furthermore, comparisons among PWMs (i.e., motifs) confirm these thresholds, despite the potential variation contributed by motif derivation ([Weirauch et al., 2013](#); [Zhao and Stormo, 2011](#)). To illustrate this concept, [Figure 3](#) depicts PBM-derived motifs obtained for the Myb/SANT family, grouped according to the relationship of their DBD AA sequences. Darkly shaded regions, which invariably contain nearly indistinguishable motifs, indicate groups of TFs with DBD similarity exceeding the 87.5% threshold for this family.

We conclude that TFs with DBD AA sequence identity above these thresholds will typically have very similar sequence specificity. As previously proposed, this presents a simple approach for broadly predicting the sequence preferences of TFs ([Alleyne et al., 2009](#); [Berger et al., 2008](#); [Bernard et al., 2012](#); [Noyes et al., 2008](#)): the 8-mer data, motifs, and consensus sequences can be directly transferred from the nearest protein above the threshold for which data are available. The data presented here encompass 37 DBD classes with sufficient data (i.e., comparisons in each of the bins in the box plots) to produce thresholds, together representing ~85% of all known eukaryotic TFs. Until more data are available, we propose that a threshold of 70% AA identity (the mean, median, and mode across all DBD classes) can be applied to the remaining DBD classes (boxplots in [Figure S2](#) show the aggregate of all of the classes with no threshold).

### Cis-BP: A Catalog of Direct and Inferred Sequence Binding Preferences

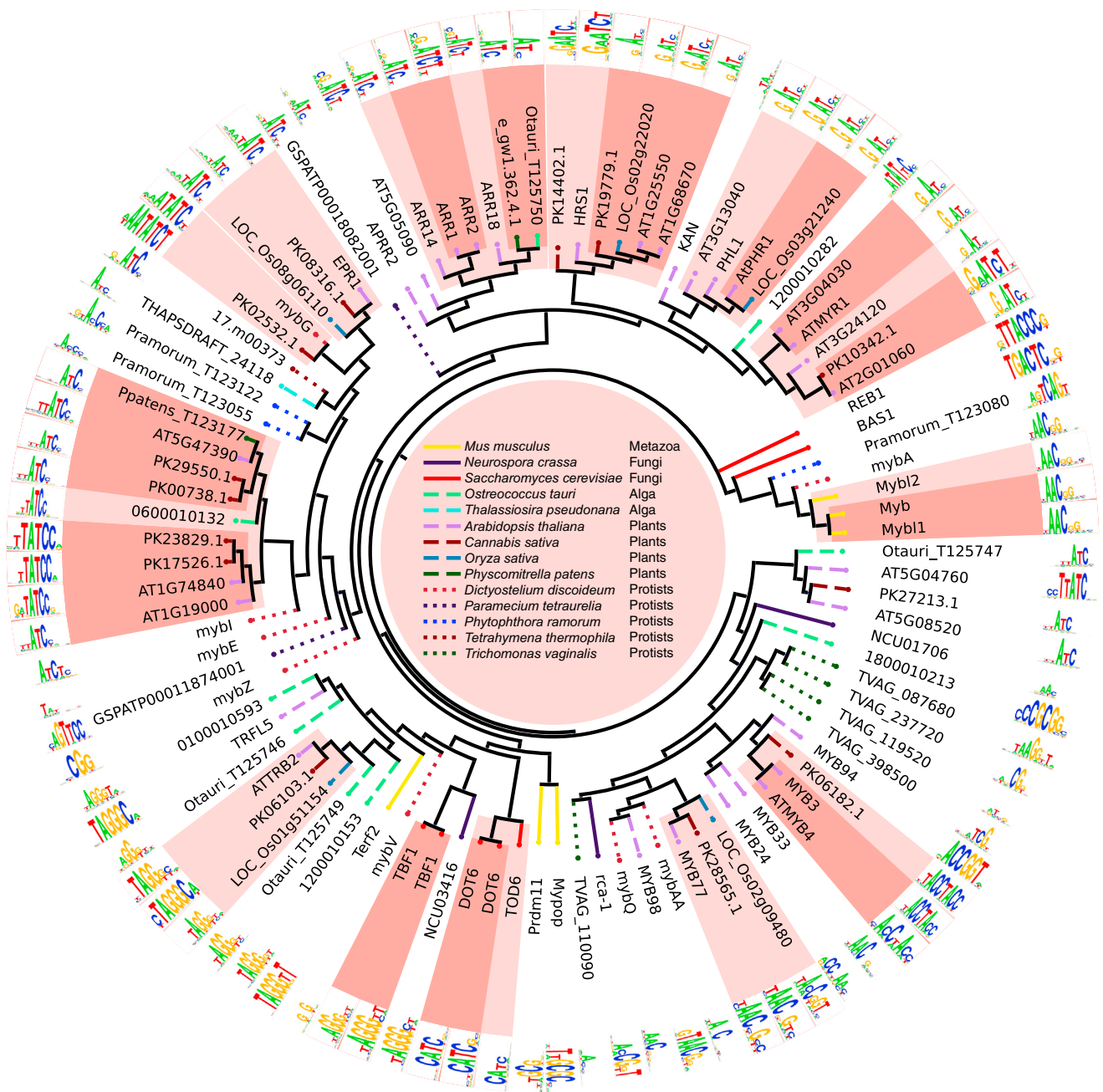
Using the DBD identity thresholds generated above, we globally assigned probable motifs (and other sequence preference data, if available) to TFs from 290 eukaryotic genomes. To do this, we supplemented the PBM data collected in this study with 4,234 additional published motifs (674 from PBMs and 3,560 from other sources—see [Table S1](#)) derived from 1,850 different proteins, mainly in human (623 proteins), mouse (409), fly (316), or yeast (216). Altogether, there are experimentally determined motifs for only 1.7% (2,750) of the 166,851 unique TFs within these genomes. Using the thresholds determined here, it is

possible to infer a motif for roughly one-third of all TFs encoded by sequenced eukaryotes, bringing the total (known + inferred) to 57,165. [Figures 4A](#) and [4B](#) summarize the coverage by DBD class and by species (see [Table S3](#) for all DBD classes and species). Lineages that benefit most from the inference scheme include vertebrates, plants, fungi, and insects, which contain many orthologs conserved in the model species analyzed most heavily. For example, the motif collection for zebrafish (*Danio rerio*), which largely consists of inferred motifs, is as complete as that of mouse and human ([Figure 4B](#)).

To facilitate use of the known and inferred motifs by the scientific community, we created a database called catalogue of inferred sequence preferences of DNA-binding proteins (Cis-BP) (<http://cisbp.cabr.utoronto.ca>). In addition to the new experimental data reported here, Cis-BP contains comprehensive 8-mer binding scores, position weight matrices, and IUPAC consensus motifs from publicly available sources (JASPAR ([Portales-Casamar et al., 2010](#)), Transfac (public data only) ([Matys et al., 2006](#)), FlyFactorSurvey ([Zhu et al., 2011](#)), FactorBook ([Wang et al., 2013](#)), and data from 674 PBM experiments taken from other studies (compiled in UniPROBE) ([Newburger and Bulyk, 2009](#)).

### In-Vitro-Derived Motifs Predict ChIP-Seq Peaks

To examine the relationship between in-vitro-defined motifs (both measured and inferred) and in vivo binding sites, we asked how well the motifs in Cis-BP predict in vivo binding, based on ENCODE human ChIP-seq data (see [Experimental Procedures](#)). To gauge the ability of a motif to discriminate between real ChIP peaks (positives) and peak sequences permuted using an algorithm that maintains all dinucleotide frequencies (negatives), we used the area under the receiver operating characteristic (AUROC) summary statistic, in which perfect discrimination between positives and negatives scores 1.00 and random guessing scores 0.50. Nearly all (111) of the 114 PBM-derived motifs achieved AUROC scores exceeding the 0.50 random expectation level in at least one ChIP data set ([Figure 5A](#) and [Table S4](#)). Strikingly, over one-third of the motifs (43 of 114), in a variety of DBD classes and cell types, achieve AUROCs exceeding 0.90 in at least one ChIP data set, including GABPA in H1-hESC cells (Ets family, AUROC = 0.99), USF2 in GM12878 cells (bHLH, 0.97), and FOS in K562 cells (bZIP, 0.97). Motifs inferred from in vitro data from related proteins achieve AUROC values similar to those obtained using in vitro motifs from exactly the same protein that was ChIPped ([Figure 5A](#)). Overall, PBM-derived motifs are equally accurate as those derived from other sources; the mean AUROCs across the 19 TFs with at least one PBM-derived motif and one Transfac motif are 0.834 and 0.827, respectively ([Figure 5B](#) and [Table S4](#)), and 0.826 versus 0.831 for the 14 TFs with both PBM and HT-SELEX-derived motifs ([Figure 5C](#) and [Table S4](#)). The small number of cases in which motifs derived from these other sources performed better than PBM motifs appear to correspond to multimeric binding that was not detected in the PBM assays. From this analysis, we conclude that there is generally no fundamental discrepancy between motifs obtained from different assays and between the in vivo and in vitro sequence preferences of TFs.



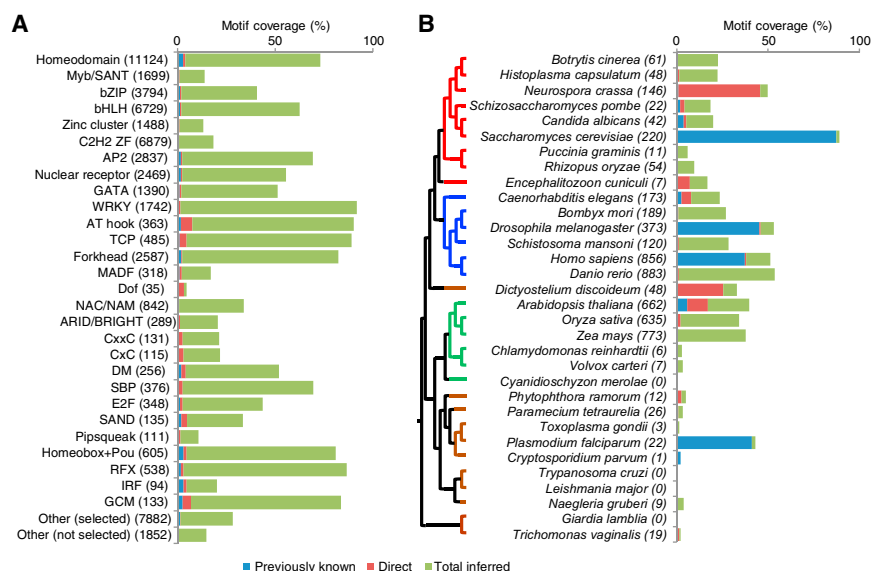
**Figure 3. Overview of Myb/SANT Family Motifs**

PBM-derived motifs from the Myb/SANT family (84 from this study, 13 from other studies) are shown. Tree reflects the percent of identical AAs after alignment. Dark shading, 87.5% AA identity (standard inference threshold); light shading, >70% AA identity (relaxed inference threshold). TBF1 and DOT6 each have two motifs because they were examined in two different studies.

**Motifs Display Positional Bias in Promoters across Eukaryotes**

Enrichment and positional bias of TF motifs in promoters has been reported in several model species and human (FitzGerald et al., 2004; Lee et al., 2007; Mariño-Ramírez et al., 2004; Ohler et al., 2002), and in some cases TFs appear to be involved in determining promoter identity (de Boer et al., 2014; Megraw

et al., 2009). However, it is unknown whether this is a general property of eukaryotic promoters. We found that genomes from vertebrates, plants, fungi, and protists all displayed enrichment of binding sites for their TFs just upstream of the TSS, relative to permuted sequences and also relative to unrelated (control) motifs taken from other lineages (Figure 6). The enrichment involved both directly determined and inferred motifs



**Figure 4. TF Motif Coverage**

TFs with multiple protein isoforms are counted as a single gene.

(A) Motif coverage by DBD class. DBD classes sorted top to bottom by number of TFs characterized in this study. Those with fewer than eight proteins characterized in this study are grouped into “Other.” “Other (selected)” indicates DBD classes selected for characterization in this study. “Other (not selected)” indicates DBD classes not characterized here. “Direct” includes those experimentally characterized in this study, but not previously known. “Total inferred” excludes those experimentally characterized in this or previous studies. (B) Motif coverage by species. Tree at left, phylogenetic relationships between organisms (Baldauf et al., 2000).

See also Table S3.

(Figure 6). Strikingly, the TF classes with motifs enriched in promoters differed between lineages of organisms (Figure S3), indicating that these trends either arose independently in different lineages, or have evolved considerably in most of them. Furthermore, we found little to no correspondence between overall motif enrichment and motif sequence composition (i.e., overall GC-content) (Table S5), and the enrichment was observed whether the promoters were characteristically GC-rich or AT-rich (i.e., predicted to be nucleosome favoring or disfavoring, respectively). We observed similar trends using a reduced set of nonredundant motifs for each organism (Figure S4), showing that the phenomenon is not due to expansion of a small number of TFs families that bind promoters. These observations indicate that, across eukaryotes, the region just upstream of the TSS typically is enriched for binding sites that may function in either promoter definition or gene regulation.

### Arabidopsis Expression Quantitative Trait Loci SNPs Are Enriched for TF Binding Sites

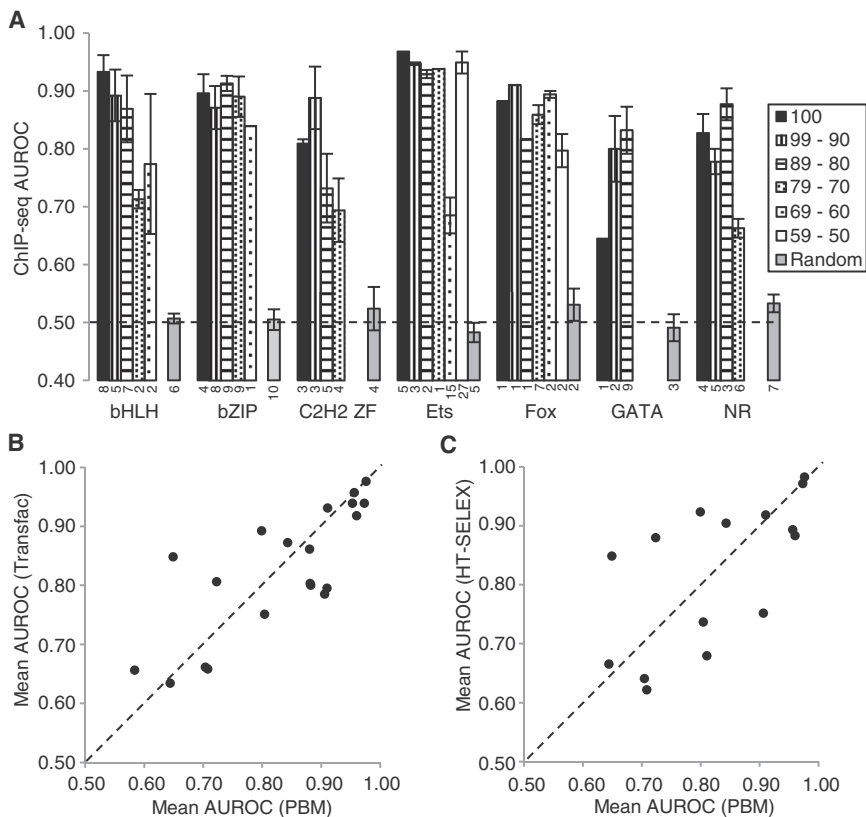
Identifying causal genetic variants and their mechanisms of action is a fundamental challenge in association studies. As our data greatly expand the motif collection in the model plant *Arabidopsis*, we examined a data set of expression quantitative trait loci (eQTLs) defined from *Arabidopsis* genomic sequences and matched seedling RNA sequencing (RNA-seq) taken from 19 strains (Gan et al., 2011). Among highly significant eQTLs found within 1 kb upstream of a TSS, there was a striking enrichment (>3-fold for the SNPs with strongest association) for overlap with TF motif matches (Figure 7A), strongly suggesting that these SNPs impact transcription factor binding. As an example, Figures 7B and 7C show a SNP in the promoter of the *AT5G47250* gene, in which loss of a potential binding site for the VNI2 TF in the “A” allele correlates almost perfectly with a dramatic increase in *AT5G47250*’s expression level, consistent with the well-characterized role of VNI2 as a transcriptional repressor (Yamaguchi et al., 2010; Yang et al., 2011). Both

*VNI2* and *AT5G47250* were associated with the plant defense response to the pathogenic oomycete *H. arabidopsidis*

in a recent genome-wide association study examining 107 different phenotypes (Atwell et al., 2010). Taken together, these results suggest that VNI2 represses the expression of *AT5G47250* in a pathogen response context, a mechanism dependent on the specific SNP present in the *AT5G47250* promoter region. We expect that the Cis-BP motif collection will be useful for similar analyses in other organisms: eQTL analysis can be performed in virtually any species for which there are multiple strains (or individuals) and does not require well-developed genetic systems. In fact, in this analysis, a similar level of enrichment was observed using only a set of 65 motifs inferred from organisms other than *Arabidopsis thaliana* (Figure S5), suggesting that the motif data need not be derived from the organism in question (see also below).

### Identification of TFs Affected by Disease-Associated Genetic Variants

Recent analyses indicate that between 85% and 93% of disease- and trait-associated variants are located in noncoding regions (Hindorf et al., 2009; Maurano et al., 2012), suggesting that many might alter TF binding events. However, the identification of specific TFs whose binding might be affected by a given variant remains a challenging and laborious task. We devised a system that utilizes all available PBM data to produce a ranked list of human TFs whose binding might be affected by any given genetic variant (see Experimental Procedures). To examine the utility of this system for analyzing human disease-associated variants, we collected a set of 15 SNPs whose alleles have been experimentally demonstrated to affect the binding of a specific TF. One of these SNPs affects two TFs, bringing the total of analyzed TF/SNP pairs to 16. Strikingly, in ten of these 16 cases, this procedure ranked the correct TF (or a highly related TF from the same DBD class) in the top five and often number one (Figure 7D and Data S1). We note that most of the novel high-ranking TFs we identify have likely never been experimentally examined and thus might also represent bona fide cases.



**Figure 5. PBM-Derived Motifs Identify In Vivo TF Binding Locations**

(A) AUROC analysis, showing ability of directly determined and inferred motifs to distinguish ChIP-seq peak sequences from scrambled sequences. We identified TFs with available ENCODE ChIP-seq data that also have PBM data available either for that TF, or for related TFs (based on the inference threshold for the DBD class). We then gauged the ability of the PBM-derived motifs to distinguish real ChIP peaks from scrambled sequences (maintaining all dinucleotide frequencies) using the AUROC (see [Experimental Procedures](#)). For each DBD class, results are binned by DBD %AA ID (key indicated at upper right). Numbers below each bar indicate the count in each bin. Error bars indicate SE. “Random” indicates results obtained with a randomly assigned, unrelated TF motif. Fox, Forkhead box. NR, nuclear receptor. [Figure S7](#) shows results obtained using an alternative null model.

(B) Comparison of AUROC for PBM-derived motifs and literature-derived motifs. We identified TFs with ENCODE ChIP-seq experimental data that also have both Transfac and PBM-derived motifs available. For each TF, we calculated the best AUROC obtained by any PBM or any Transfac motif on any of the ENCODE cell line ChIP experiments for that TF. For TFs with multiple motifs from the same source, the plot shows the mean AUROC across the motifs.

(C) PBM-derived motifs versus HT-SELEX-derived motifs. Same as for (B), but including only TFs with motifs available both from PBMs and a recent HT-SELEX study ([Jolma et al., 2013](#)). See also [Table S4](#).

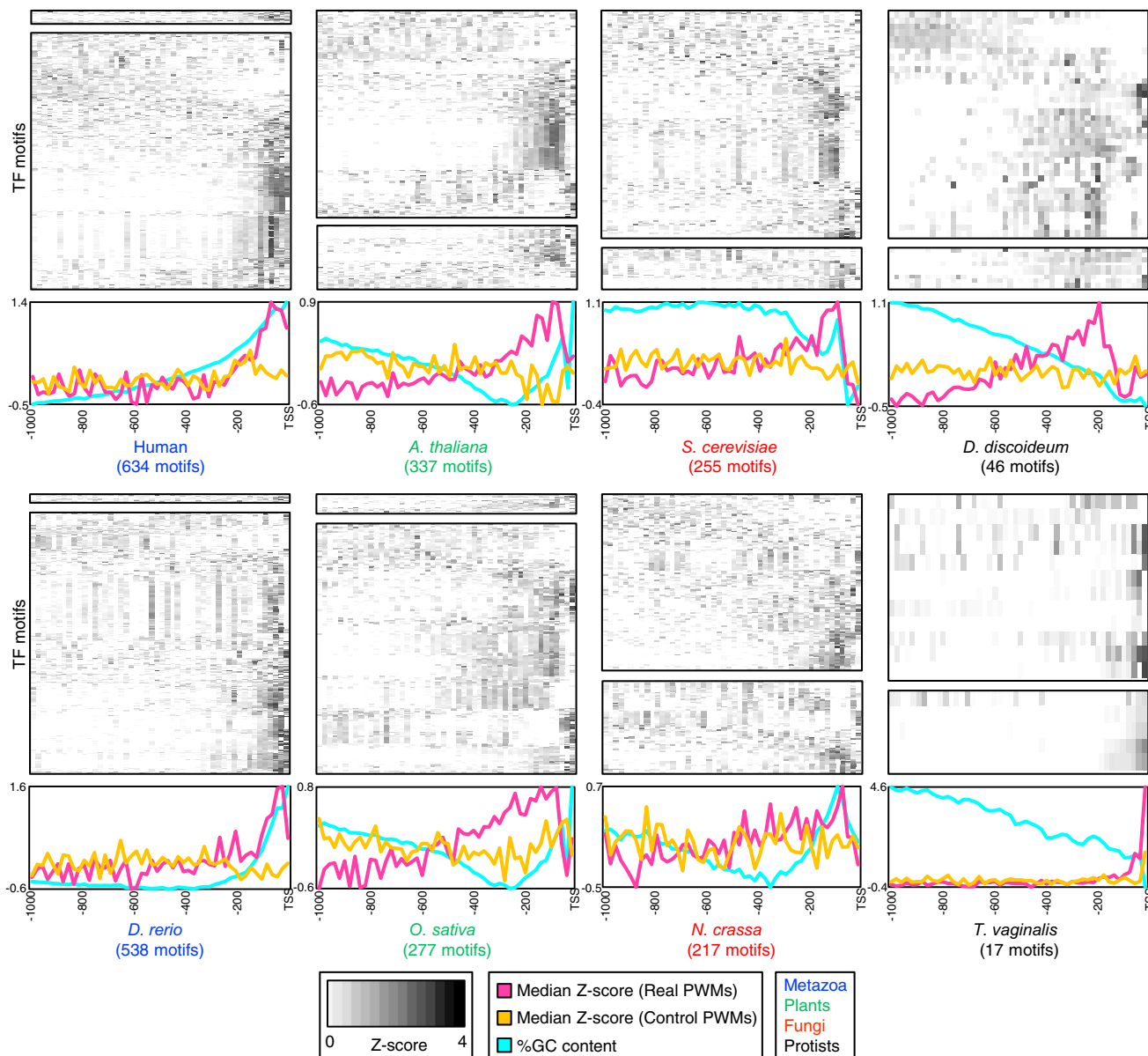
In one example, a recent study implicated rs554219 in estrogen-receptor-positive breast cancer tumors and used a series of experiments to predict and eventually establish that this SNP causes the differential binding of two Ets family TFs, ELK4 and GABPA ([French et al., 2013](#)). Because rs554219 does not overlap ChIP-seq binding peaks from ENCODE or other sources, ELK4 and GABPA were identified using a series of EMSA competition experiments involving known TF binding sites, a laborious process involving substantial guesswork. Our automated computational procedure correctly ranked ELK4 number one and GABPA number two out of all human TFs (their EMSA data demonstrate strong and moderate differential binding of ELK4 and GABPA, respectively). The ELK4 prediction is based on an inference from the pufferfish *Tetraodon nigroviridis* ELK4 protein (81% AA ID to human ELK4), further illustrating the utility of cross-species inferences. Moreover, differential binding of ELK4 to the alleles of rs554219 can also be predicted using data inferred from *Drosophila melanogaster* Ets21C (57% AA ID to human ELK4) and *Caenorhabditis elegans* F19F10.1 (only 33% AA ID to ELK4). Thus, our system (available at <http://cisbp.ccb.utoronto.ca/TFTools.php>) can accurately predict the specific TFs whose binding is affected by risk alleles of disease-associated SNPs, even when using PBM data inferred from distantly related organisms.

## DISCUSSION

We anticipate that the new data collected here—as well as the inferred motifs across eukaryotes—will be an invaluable resource and knowledge base for functional genomics and analysis of gene regulation. In addition to the data itself, the Cis-BP database also contains web-based interfaces to tools for scanning DNA sequences for putative motifs, reporting the TFs with motifs similar to a given motif, predicting the motif recognized by a given TF based on its DBD AA sequence, and identifying TFs that will bind differentially to two different DNA sequences (e.g., disease risk and nonrisk alleles). Cis-BP will enable dissection of regulatory mechanisms from expression data even in the vast majority of species for which there are currently no genetic tools.

The analyses here present a strategy to rapidly populate TF motif collections across the eukaryotes. As we previously proposed for RNA-binding proteins ([Ray et al., 2013](#)), targeting members of the largest groups of uncharacterized proteins for experimental analysis will allow the largest number of inferred motifs to be obtained. Motif inference based on DBD identity alone is only a first approximation, but it is remarkably cost effective: the analyses described here indicate that motifs can be inferred for 34% of all TFs, using data from only 1.7%. We extrapolate that 1,032 additional successful experiments





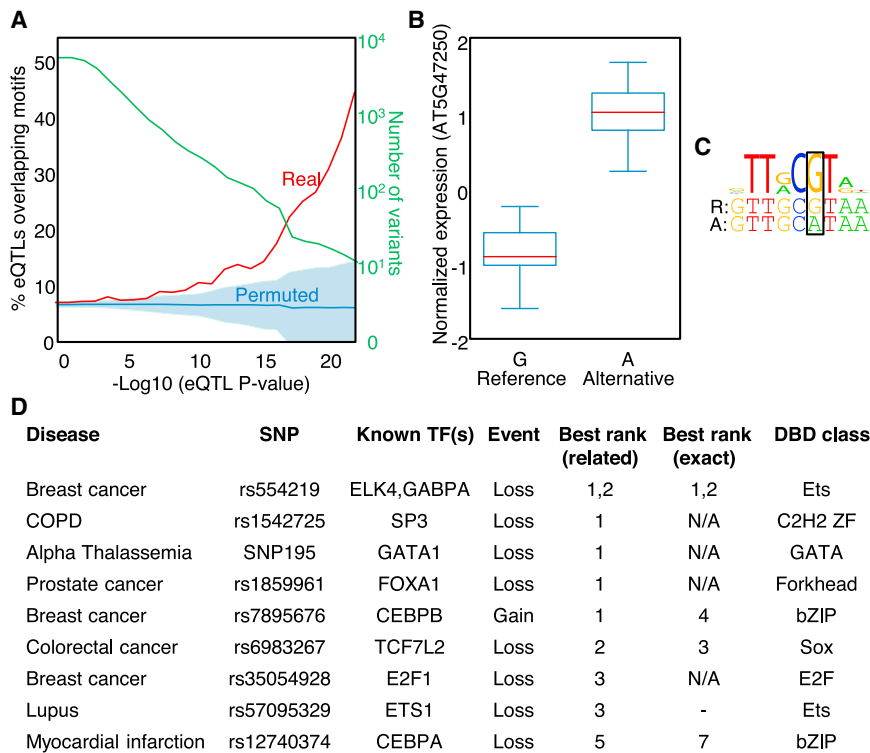
**Figure 6. Positional Bias of Motif Matches in Eukaryotic Promoters**

PBM-derived PWMs (direct, top heatmap; inferred, bottom heatmap) scored in 20 bp bins, normalized to dinucleotide-permuted controls, averaged across all promoters, and displayed as Z scores (see [Experimental Procedures](#)). Each row in the heatmap corresponds to one PWM. Rows were clustered using hierarchical clustering (Pearson correlation, average linkage). Summary plots at the bottom indicate the median Z score, taken across all PWMs from the indicated species (“Real PWMs”), or across a set of PWMs from unrelated lineages (“Control PWMs”) (see [Experimental Procedures](#)). See also [Table S5](#) and [Figures S3](#) and [S4](#).

(<0.5% of all TFs and a number identical to that in this study)—one from each of the largest groups of orthologs and paralogs with no known motifs—would increase coverage across the eukaryotes from 34% to 48%.

The inference scheme described here relies on the high degree of conservation among DBDs. Indeed, our analyses confirm the “deep homology” that has been described for metazoan developmental processes and the TFs that regulate them

(e.g., homeodomains) ([Berger et al., 2008](#); [Carroll, 2008](#); [Noyes et al., 2008](#)) and furthermore indicate that deep homology is a property of the sequence preferences of many TFs in all eukaryotic kingdoms. Our initial analyses (data not shown) suggest that many motifs likely date to the base of metazoans, land plants, angiosperms (flowering plants), or euteleostomi (bony vertebrates), consistent with well-established TF expansions in these lineages ([de Mendoza et al., 2013](#); [Weirauch and Hughes, 2011](#)).



**Figure 7. Overlap of Predicted TF Binding Sites with *cis*-eQTLs**

(A) Number and percentage of *Arabidopsis cis*-eQTLs overlapping motifs, as a function of eQTL significance. Shaded region indicates one standard deviation in the expected distribution (see [Experimental Procedures](#)).

(B) A *cis*-eQTL affecting the expression of the AT5G47250 gene. Boxplots (see [Figure 2](#) legend) indicate the median normalized gene expression level for each allele of the *cis*-eQTL. “Reference” indicates the allele present in the *Arabidopsis* reference genome assembly.

(C) The same *cis*-eQTL “breaks” a putative binding site for the VNI2 transcriptional repressor. Sequence logo depicts the DNA-binding motif we obtained for VNI2. Sequences below indicate the reference (top) and alternative (bottom) alleles of the *cis*-eQTL SNP (boxed) and its flanking bases.

(D) Prediction of human TF binding events altered by disease risk alleles. We created a method for using PBM data to predict TFs whose binding is affected by disease-associated genetic variants and applied it to 16 known examples. Shown here are the ten cases in which we ranked the correct TF (column labeled “exact”) or a highly related TF from the same DBD class (column labeled “related”) within the top five TFs. The “Event” column indicates whether the risk allele results in a “Loss” or “Gain” of binding of the TF. N/A, indicates that PBM data are not available for the corresponding TF; –, indicates that the TF did not receive a rank because both alleles had an *E* score >0.45.

See also [Figure S5](#).

Despite widespread conservation, TF repertoires do change over evolutionary time, and these changes likely shape eukaryotic evolution ([de Mendoza et al., 2013](#)). TFs that tend to diversify, such as the large metazoan C2H2 class ([Stubbs et al., 2011](#)), will present an ongoing challenge to a complete characterization of eukaryotic TF motifs. In other lineages, other classes of TFs have expanded and diversified, including the nuclear receptor class in *C. elegans* ([Maglich et al., 2001](#)), the zinc cluster/GAL4 class in fungi ([Shelest, 2008](#)), and several classes in plants ([Lang et al., 2010](#)). The data described here confirm that the sequence specificities of at least some of these factors have also diversified. Mapping recognition codes represents an alternative approach to more complete cataloguing of TF motifs, and the data presented here provide many new examples for the study of TF-DNA recognition. Exceptions to the simple AA similarity rules described here should also be informative regarding mechanisms of sequence recognition, because they will identify AA residues critical for DNA sequence specificity ([De Masi et al., 2011](#); [Noyes et al., 2008](#)). Ongoing efforts to further the collective knowledge of TF binding specificities will greatly advance our understanding of TF-DNA interactions, as well as our ability to interpret the function of DNA sequences, including understanding the functional impact of natural genetic variants in human and other species.

## EXPERIMENTAL PROCEDURES

Full details are provided in the [Extended Experimental Procedures](#).

## Data Availability

PBM data are available in the Gene Expression Omnibus (GEO) database under accession number GSE53348. PBM data, clone information, and other data from analyses carried out in this study are available on the project web site: <http://hugheslab.cabr.utoronto.ca/supplementary-data/CisBP/>. Additional data (including 8-mer scores, PWMs, sequence logos, and information on TFs) are found on the CisBP web server (<http://cisbp.cabr.utoronto.ca/>).

## Selection, Cloning, and PBM Analysis of TFs

We compiled the predicted proteomes of 290 eukaryotic organisms from a variety of sources and supplemented them with an additional 49 known TFs from organisms without fully sequenced genomes. We scanned all protein sequences for putative DNA-binding domains (DBDs) using the 81 Pfam ([Finn et al., 2010](#)) models listed in [Weirauch and Hughes \(2011\)](#) and the HMMER tool ([Eddy, 2009](#)). Each protein was classified into a family based on its DBDs and their order in the protein sequence. We selected 2,913 individual TFs to analyze, using several different criteria, including a relatively even balance among DBD classes and species, a survey of different levels of sequence identity among proteins, and a deeper focus on several model organisms and abundant DBD classes. For most constructs, we designed primers to clone the region encompassing all DBDs plus the 50 flanking endogenous AAs on either side (or until the termini of the protein) by conventional PCR methods into one of a panel of T7-GST vectors for expression in *E. coli* (referred to hereafter as “plasmid constructs”). PBM laboratory methods were identical to those described in [Lam et al. \(2011\)](#) and [Weirauch et al. \(2013\)](#). Each plasmid was analyzed in duplicate on two different arrays with differing probe sequences (denoted “ME” and “HK”). Calculation of 8-mer Z and E scores was performed as previously described ([Berger et al., 2006](#)). To obtain a single representative motif for each protein, we used a procedure similar to a recent study from our group in which we generated motifs for each array using four different algorithms and chose the best-performing single motif based on cross-replicate array evaluations ([Weirauch et al., 2013](#)).

### Inference Scheme

We established a separate inference threshold for each DBD class. We first aligned the DBD sequences of all constructs within a DBD class using clustalOmega (Sievers et al., 2011). We then calculated the AA %ID for all construct pairs (i.e., the number of identical AAs in the alignments). Within each DBD class, we grouped all PBM construct pairs into bins, based on AA %ID. We used overlapping bins of size 10, ranging from 0 to 100, increasing by 5. We calculated the precision of each bin by comparing the DNA sequence preferences obtained from all characterized protein pairs contained in the bin. We quantified the similarity of the DNA sequence preferences of two proteins as the fraction of shared high-scoring 8-mers. We considered a prediction to be correct only if this fraction exceeded the value obtained at the 25th percentile of experimental replicates (i.e., the fraction of shared 8-mers between ME and HK arrays for the same protein). The proportion of predictions scored as correct (i.e., precision) for each bin of each DBD class is shown as magenta stars in Figure 2 and Data S1.

We chose inference thresholds for each DBD class based on the precision scores of each AA %ID bin. Because we used the 25th percentile threshold to define precision, we would expect a precision of 0.75 or higher in each AA %ID bin. We therefore chose an inference threshold for each DBD class by identifying the final AA %ID bin before precision drops below 0.75 (vertical bars in Figure 2). Similar thresholds were obtained regardless of the  $E$  and  $Z$  score 8-mer thresholds used and also regardless of the replicate overlap percentile considered (i.e., 25th percentile, requiring 0.75 precision or 20th percentile, requiring 0.80 precision) (Figure S6). The final threshold for a DBD class was chosen as the median threshold across the eight 8-mer similarity measures (see Figure S6 and Data S1). We found this scheme to be appropriate for most DBD classes (all of which are depicted in Figure 2). For three DBD classes (IRF, CXXC zinc fingers, and Dof zinc fingers), we could not establish a threshold—these therefore received a threshold of 100%. We used a threshold of 40% for AT-hook TFs, which recognize AT-rich sequences, based on manual inspection of the data (see Data S1). For the remaining classes, with suggestive but insufficient data, we chose a threshold of 70%, which is the mean, median, and mode threshold across all DBD classes.

We used the AA %ID of all pairs of proteins to infer motifs, 8-mer scores, and consensus sequences within each DBD class by simple transfer (i.e., aligning the DBD sequences of all proteins and all constructs in a given DBD class, as described above, and calculating the AA %ID of each protein with each construct). We also evaluated the effectiveness of our inference scheme in a leave-one-out cross validation framework, in which the PBM data for each characterized protein was held out and compared to the PBM data of its nearest neighbor (i.e., the characterized protein with highest AA %ID), using a similar scoring scheme to that used to calculate the precisions.

### Comparison to ChIP-Seq Data

We calculated AUROC scores on real and permuted (maintaining dinucleotide frequencies) ChIP-seq peak sequences following Weirauch et al. (2013). We obtained ENCODE consortium human ChIP-seq data from the UCSC Genome Browser (Rosenbloom et al., 2012). For each ChIP experiment, we extracted the top 500 scoring peak region sequences and scored them (and the permuted sequences) using all direct and inferred PWM models for the given TF. For each PWM/experiment pair, we then calculated the AUROC using these sets of 500 positives and 500 negatives. Similar results were obtained using a negative set consisting of ChIP-bound peaks from unrelated TFs with matched GC-content (Figure S7).

### Positional Bias of Motifs in Eukaryotic Promoters

We obtained the 1,000 bases upstream of transcription start sites (or, if unavailable, translation start sites) and scored the PWMs of each organism at each position. We then placed the resulting scores into 20 bp bins, summed the scores for each bin, and took the average across all promoters for the given species for each bin. To correct for mono- and dinucleotide biases, we also scored shuffled promoter sequences, which were created by shuffling the sequences within each 20 bp bin (while maintaining dinucleotide frequencies). For each PWM, we then calculated the ratio of each bin's real score relative to the score of the shuffled sequence. The resulting ratios were then normalized across all bins for the given PWM using a standard  $Z$  score transformation.

We also calculated  $Z$  scores for a negative control set of TF PWMs for each organism, consisting of a collection of random motifs from species in other clades that were unrelated to any PWM from the given species.

### Arabidopsis eQTL Analysis

We used a publicly available data set (Gan et al., 2011) containing genome-wide RNA-seq variance-stabilized expression levels (Huber et al., 2002) taken from 19 strains of seedling *Arabidopsis thaliana* and matching genome sequences. We identified matches to each *A. thaliana* PWM within the 1,000 bases upstream of each TSS. We calculated the percentage of genetic variants that affect these putative binding sites, as a function of the *cis*-eQTL  $p$  value of the variant (red line, Figure 7). We also created a null distribution (blue line and blue shaded region, Figure 7) to exclude the possibility that the observed percentages might solely be due to the higher density of TF binding sites in promoter regions.

### Human Disease SNP/TF Analysis

We devised a system for utilizing our collection of PBM data to identify candidate human TFs whose binding might be affected by the allelic sequences of genetic variants. In this system, we score each variant (along with its flanking genomic bases) using 8-mer  $E$  scores taken from the 3,132 PBM experiments contained in our database. For each PBM experiment, we identify the highest scoring 8-mer  $E$  score attained by any of the risk allele sequences ( $E_{risk}$ ) and the highest attained by any nonrisk allele ( $E_{nonrisk}$ ). We then identify all PBM experiments where only one of  $E_{risk}$  and  $E_{nonrisk}$  has an  $E$  score value exceeding 0.45 (values above this threshold will likely be strongly bound by the given TF (Berger et al., 2008)) and map these experiments to human using the inference scheme. This procedure thus produces a ranked list of human TFs whose binding is likely to be affected by the alleles of a given SNP (e.g., strongly binding to one allele but not binding to the other).

### ACCESSION NUMBERS

The Gene Expression Omnibus (GEO) accession number for the PBM data reported in this paper is GSE53348.

### SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, seven figures, six tables, and one data file and can be found with this article online at <http://dx.doi.org/10.1016/j.cell.2014.08.009>.

### AUTHOR CONTRIBUTIONS

M.T.W. and T.R.H. conceived of the study, oversaw analyses, and wrote the manuscript. M.T.W. performed most of the computational analyses. A.Y. performed and oversaw most of the experimental analyses. M.T.W. and M.A. developed and implemented the database and web server. A.G.C., A.M.M., I.M., H.Z., A.G., J.C.L., M.G., M.G.L., E.H., and J.R.H. performed additional experiments. H.S.N., M.A., H.vB., and X.C. provided additional computational support. P.D. performed the *Arabidopsis* eQTL analysis of Figure 7. S.L. and M.A. performed the motif location enrichment analysis in Figure 6. K.C. performed motif analyses related to Figure 3. T.M. performed analysis for the ChIP-seq comparisons in Figure 5. S.G., G.S., A.J.M.W., F.Y.B., G.R., L.F.L., and J.R.E. developed and oversaw experimental and computational analyses. All authors provided critical feedback on the manuscript, with extensive contributions from A.M.M., H.vB., J.R.H., A.J.M.W., L.F.L., M.T.W., and T.R.H.

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