Primary sequence and epigenetic determinants of in vivo occupancy of genomic DNA by GATA1

Ying Zhang1,2, Weisheng Wu1,3, Yong Cheng1,4, David C. King1,5, Robert S. Harris1, James Taylor6, Francesca Chiaromonte1,7 and Ross C. Hardison1,4,*

1Center for Comparative Genomics and Bioinformatics, Huck Institutes of Life Sciences, 2Graduate Programs in Genetics, 3Graduate Programs in Cell and Developmental Biology, 4Department of Biochemistry and Molecular Biology, 5Graduate Programs in Bioinformatics and Genomics, The Pennsylvania State University, University Park, Pennsylvania, PA 16802, 6Department of Biology, Emory University, Atlanta, GA 30333 and 7Department of Statistics, The Pennsylvania State University, University Park, Pennsylvania, PA 16802, USA

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ABSTRACT

DNA sequence motifs and epigenetic modifications contribute to specific binding by a transcription factor, but the extent to which each feature determines occupancy in vivo is poorly understood. We addressed this question in erythroid cells by identifying DNA segments occupied by GATA1 and measuring the level of trimethylation of histone H3 lysine 27 (H3K27me3) and monomethylation of H3 lysine 4 (H3K4me1) along a 66 Mb region of mouse chromosome 7. While 91% of the GATA1-occupied segments contain the consensus binding-site motif WGATAR, only ~0.7% of DNA segments with such a motif are occupied. Using a discriminative motif enumeration method, we identified additional motifs predictive of occupancy given the presence of WGATAR. The specific motif variant AGATAA and occurrence of multiple WGATAR motifs are both strong discriminators. Combining motifs to pair a WGATAR motif with a binding site motif for GATA1, EKLF or SP1 improves discriminative power. Epigenetic modifications are also strong determinants, with the factor-bound segments highly enriched for H3K4me1 and depleted of H3K27me3. Combining primary sequence and epigenetic determinants captures 52% of the GATA1-occupied DNA segments and substantially increases the specificity, to one out of seven segments with the required motif combination and epigenetic signals being bound.

INTRODUCTION

A fundamental paradigm in regulation of gene expression is the binding of a regulatory protein to a specific DNA sequence, which then leads to activation or repression by a variety of mechanisms. The specific DNA sequence recognized by a protein is its binding site, which can be characterized as a motif—either a specific string or a position-specific weight matrix, often a consensus of sequences at multiple binding sites. The binding sites for many regulatory proteins have been determined by sequencing DNA segments with a high affinity for the protein in solution. Binding-site motifs tend to be quite short (hexamers are common), and thus they occur frequently in any long DNA sequence—much more frequently than specific occupancy is observed in vivo. Therefore, an enduring problem is to identify other determinants of occupancy in vivo (1).

Gene regulation involves transcription factor interactions with both primary DNA sequence elements and the chromatin structure of the regions that contain these elements. In particular, histone modifications play a strong role in transcriptional regulation, and are likely to be significant contributors to determining in vivo occupancy. Specific classes of regulatory elements have been shown to be accompanied by distinctive histone modifications, for example trimethylation of lysine 27 of histone H3 (H3K27me3) is correlated with repression of gene expression (2), and monomethylation of lysine 4 of histone H3 (H3K4me1) is associated with enhancers (3).

High throughput methods for mapping the positions of DNA segments cross-linked to proteins and immunoprecipitated from chromatin, namely ChIP-chip and ChIP-seq (4,5), are used to determine comprehensively the DNA segments occupied by particular proteins or having particular chromatin modifications in vivo. Thus careful examination of DNA sequences and epigenetic marks in the occupied segments is expected to reveal the determinants of occupancy in vivo, show the extent to which primary sequence can contribute to the specificity of occupancy, and explore the ability of histone modification in chromatin to explain additional specificity.

*To whom correspondence should be addressed. Tel: +1 814 863 0113; Fax: +1 814 863 7024; Email: rch8@psu.edu

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The cis-regulatory modules studied in eukaryotes consist of binding sites for multiple proteins (6,7), and thus binding site motifs for other proteins that commonly co-occupy DNA segments with the protein of interest are good candidates for determinants of binding specificity in addition to the primary binding site motif. In order to pursue this strategy, the protein of interest must have an identifiable primary binding site that is present in most of the occupied DNA segments. This is not always the case, e.g. the cognate consensus motif was not found in the majority of DNA segments in mammalian cells occupied by transcription factors Sp1, c-Myc and p53 (8) and E2F1 (9). However, occupancy by the transcription factor GATA1 in vivo is almost invariably associated with the primary consensus binding-site motif WGATAR (10). Thus we have chosen to search for additional discriminative motifs that help determine specificity of occupancy by GATA1.

The transcription factor GATA1 is a zinc finger protein that is required for normal hematopoiesis and plays a role in regulating most of the genes that define the mature erythroid phenotype (11,12). Early work identified WGATAR as the consensus motif bound by GATA1 (13–16). Some (17) but not all (18) investigations using in vitro site selection assays indicated that GATA1 also has high affinity for non-consensus motifs in solution. While directed studies of individual cis-regulatory modules have shown binding in vitro of GATA1 to DNA that deviates from the consensus motif (19–21), other studies find the non-consensus motifs to be poor predictors of enhancer activity (22). Even limiting the analysis to the consensus binding site motif WGATAR, only a small fraction of all such motifs are bound in vivo (23–25).

We searched for other determinants of GATA1 occupancy in vivo by generating a set of 314 DNA segments that are occupied by GATA1 in the mouse erythroid cell line G1E-ER4. These were discovered by immunoprecipitating DNA fragments associated with GATA1 in vivo, followed by hybridization to a high-density tiling array of non-repetitive DNA sequences (ChIP-chip, 4) along a large segment (66 Mb) of mouse chromosome 7. Randomly sampled ChIP-chip positive regions were re-tested in a quantitative PCR (qPCR) assay using independent ChIP material, yielding very high validation rates for GATA1 occupancy. This dataset also provides a large number of reliably unbound segments to be used for studies of discriminative features in the sequence.

The motif WGATAR is well known as a primary determinant of binding, occurring in 91% of the bound regions we discovered. However, the motif alone provides poor specificity, as ~0.7% of DNA segments that have this motif are occupied. Thus, we focused on additional sequence motifs found in DNA segments containing WGATAR that could distinguish bound from unbound. We constructed ‘background’ sets of unbound DNA segments with a distribution of WGATAR motifs and GC content indistinguishable from the one observed for the ‘foreground’ (bound) DNA segments. We employed the computer program Discriminating Matrix Enumerator, or DME (26), to identify enriched motifs (represented by scoring matrices) in the foreground set compared with a series of background sets of unbound sequences. These over-represented motifs were then matched to known binding sites for other transcription factors, and their discriminatory power evaluated both against an independent testing set and against the bulk genome. While several discriminatory motifs were discovered in this way, they did not fully explain the specificity of occupancy by GATA1. However, when histone modifications around the DNA segments were examined in addition to motif combinations, the specificity of occupancy significantly increased from 1 out of 147 segments containing a WGATAR motif to 1 out of 7 segments that have high H3K4me1 signal and contain the combinations of discriminatory motifs.

**MATERIALS AND METHODS**

**GATA1 ChIP-chip and peak calling**

A previous study from this laboratory identified 63 DNA segments occupied by GATA1 in mouse erythroid cells (10). That study employed chromatin immunoprecipitation (ChIP) by anti-estrogen receptor (anti-ER) antibody to enrich for DNA segments occupied by the hybrid GATA1-ER protein in the G1E-ER4 cell line, 24 h after activation of GATA1-ER with estradiol (12). Peaks of hybridization of this ChIP material to a NimbleGen high-density array tiling across 66 Mb of mouse chromosome 7 were further tested by large-scale independent validation using qPCR to identify the 63 high-quality occupied DNA segments (10). In the current study, we used antibody against the GATA1 portion of the GATA1-ER hybrid protein (Ab GATA-1 N6; sc-265), and hybridized the new ChIP material in duplicate to the NimbleGen microarray. These new data showed peaks on the DNA segments previously demonstrated to be occupied by GATA1, and they had low signal in the regions where many false positives were found in the previous study. Thus we applied three peak-calling programs to the new GATA1 ChIP-chip data to predict occupied DNA segments: Mpeak (27), TAMALPAIS (28) and PASS (29). For Mpeak and TAMALPAIS, application of the most stringent thresholds (+3SD for Mpeak, L1 for TAMALPAIS) identified 238 peaks (union and merge). For PASS, we allowed one false discovery in the peaks called from the 66 Mb region (corresponding to a false discovery rate of 0.4%), and identified 90 additional peaks. A 30 kb segment of mouse chromosome 7 is covered almost continuously by ChIP hybridization signal rather than showing the usual discrete peak. The nature of GATA1 binding to this interval is not clear and requires further study. Thus peaks residing in this 30 kb segment were not included in the analysis. This left 304 DNA segments containing in vivo binding sites for GATA1 along the 66 Mb locus. These are validated at a very high rate, regardless of the program used to call them. Of the 304 peaks, 101 were tested independently by qPCR (including the 63 previously published), and 99 (98%) were validated. The non-validated regions were then removed from the
estradiol (10^2 U/ml erythropoietin and 50 ng/ml kit ligand. Beta-
G1E-ER4 cells were cultured in Iscove’s modified
ment for the negative control regions.
validation
levels on intervals from the regions with no peak calls
for real-time PCR. The abundance of GATA1 on DNA
controls. For each qPCR experiment, the enrichment
samples for qPCR along with the experimental tests and
Zfpm1 upstream from
Zfpm1 (an occupied segment in the first intron of the gene
qPCR validation
Thirty-eight peaks (in addition to the 63 already validated)
were randomly selected from the total of 304 GATA1
peaks in the 66 Mb region of mouse chromosome 7.
The enrichment of GATA1 in these regions was tested by real-time qPCR. The templates examined by qPCR
are GATA1-ChIPed DNA and the ChIP input DNA,
both amplified twice by GenomePlex Complete Whole
Genome Amplification (WGA) Kit. A serial dilution of
the ChIP input DNA was used to build standard curves
for real-time PCR. The abundance of GATA1 on DNA
intervals was measured by the relative enrichment of
GATA1-ChIPed DNA compared with the input DNA.
To control for variation between the several qPCR
experiments needed for the assays, a positive standard
(an occupied segment in the first intron of the gene
Zfpm1) and a negative standard (a DNA segment
upstream from Zfpm1) were included in each plate of
samples for qPCR along with the experimental tests and
controls. For each qPCR experiment, the enrichment
levels for the DNA segments were normalized by subtracting
the enrichment of the negative standard from the
enrichment on the tested segment and then divided by the
difference between the enrichment of the positive
standard and the negative standard, using the equation
(S−N)/(P−N), where S, P and N are the qPCR enrichment
values for the tested region, the positive standard
and the negative standard, respectively. Enrichment
levels on intervals from the regions with no peak calls
(negative controls) were tested to build a validation
threshold. The enrichment of GATA1 on these three
negative controls was normalized in the same way as
above. The validation threshold is set to be the mean
plus two standard deviations of the normalized enrichment
for the negative control regions.

ChIP-chip analysis of histone modifications
G1E-ER4 cells were cultured in Iscove’s modified
Dulbecco’s media (IMDM) with 15% fetal calf serum,
2 U/ml erythropoietin and 50 ng/ml kit ligand. Beta-
estradiol (10^-7 mol/l) was added to the culture to
activate the G1E-ER4 cells for 24 h.
The ChIP assay was performed as described previously
(10). The antibody against histone H3 trimethylated
on K27 was purchased from Millipore (catalog number
07-449) and the antibody against histone H3
monomethylated on K4 was purchased from Abcam
(catalog number ab8895). The ChIP DNA from induced
G1E-ER4 cells after cross-linking and immunoprecipitation by antibody was amplified using Whole
Genome Amplification (WGA) kit from Sigma. The
amplified DNA was hybridized to the NimbleGen array
16 from the mm6 version of the high-density tiling array,
which covers 69 577 286–133 051 535 position on mouse
chromosome 7 (mm6 assembly). The tiling array
contains 50 bp-long oligonucleotide probes spaced 50 bp
apart (100 bp from the start of one to the start of the
next) in the non-repetitive DNA within the above
genomic region.
The ChIP-chip data called can be viewed and down-
loaded from our customized genome browser (http://
main.genome-browser.bx.psu.edu/), mouse February
2006 assembly, group: Erythroid Gene Regulation
group, composite track: Eryth TF ChIP chr7, tracks:
‘ChIP with antibody against GATA1 portion of
GATA1-ER in cells with GATA1 restored and activated’
replicates 1 and 2.

Collection of positive and negative regions for
identification and evaluation of enriched motifs
We processed the 302 peaks into 314 intervals of 500 bp
each, which covers the amplicons used in qPCR verifica-
tion. These 314 intervals are the foreground set to evaluate
the sequence patterns that might contribute to the in vivo
occupancy of GATA1. This foreground set was randomly
split into two sets of equal size, comprising a foreground
training set (157 intervals) used for the identification
of enriched motifs and a foreground testing set (157
intervals) used to evaluate the predictive power of the
identified motifs. Two genomic features were controlled
in the background datasets of unoccupied DNA
fragments; namely, G + C content and occurrence of
WGATAR motifs. Three background datasets, with
distributions for both G + C content and occurrences of
WGATAR motif very similar to those in the foreground
training set, were randomly selected for use in the identi-
fication of enriched motifs (background training sets). One
more background set was formed for evaluation of
enriched motifs (background testing set)—its distributions
for G + C content and occurrences of WGATAR were
matched to those in the foreground testing set (Figure 2B).

Identification of significantly enriched motifs
DME2 (beta version 2008_08_30) was used to identify
enriched motifs of size 6 in the training datasets. DME2
is the beta version of a previously published program
DME (26), which exhaustively searches a finite space of
possible matrices, and tests the relative overrepresentation of
the matrices in the foreground set compared with the
background. The identified motifs are not only enriched in
the foreground set but they also have high specificity
(measured by information content). To ensure the
identified motifs were robust with respect to the choice of
background, DME was run independently using each
background training set, and motifs identified in all three
runs were selected. Corresponding motifs between runs
were identified by high matrix similarity determined
using matcompare (30) (most had identical consensus sequences and similar enrichment scores among the three runs computed by DME2). As expected, almost all the motifs discovered by this process show a level of sensitivity and specificity against the foreground training set that exceeds that of non-discriminators (Supplementary Figure S1).

Measuring discriminatory power of motifs

The discriminatory power of each motif (and combination of motifs) was evaluated in terms of sensitivity and specificity. The sensitivity is defined as the fraction of the 157 occupied intervals in the foreground test set that contain the motif (or motifs). The specificity reflects the ability of a classifier to reject unoccupied DNA segments, and is defined as the fraction of 157 unoccupied segments in the background test set that do not contain the motif (or motifs).

Matching words to a library of known transcription factor binding sites

Enriched motifs were first compared with a customized library composed of scoring matrices from the latest Jaspar library (31) and two curated scoring matrices for EKLF and GATA1 binding sites, using the program matcompare (30). To supplement the motifs in the Jaspar library, we also applied a string-matching program to compare the enriched motifs to the consensus sequences for known transcription factor binding sites (TFBSs), using a non-redundant set (32) derived from TRANSFAC (33), the Jaspar library (31), and custom consensus sequences for binding sites for three erythroid transcription factors: EKLF_bs (CCNCACCCW), GATA1_bs (WGATAR), CP2 bs (CCWG half site). The program scores exact matches as 1 and mismatches as −1, and it gives fractional positive scores to matches at a degenerate position other than N, e.g. matches to a 2-fold degenerate site have a score of 0.5, etc. The TFBS with the highest match score to a motif is considered the best match of the motif. This approach produced 14 matches of previously described TFBS motifs to one or more over-represented motifs (Supplementary Table S6). Motif matches that violated known rules for critical positions in a TFBS were then removed by inspection. For example, the word GTGAGC is a significant match to the consensus binding site GTGGGGCGNR for EGR. However, the nucleotide G at the fourth position is critical for the binding, so this motif was removed from the list because it violated this rule.

RESULTS

Binding specificity of GATA1 along mouse chromosome 7 in erythroid cells

With new GATA1 ChIP-chip data from mouse erythroid G1E-ER4 cells, we mapped 302 DNA segments occupied by GATA1 along 66 Mb of mouse chromosome 7 (see ‘Materials and Methods’ section). A total of 101 GATA1 peaks were tested independently by qPCR, including the 63 previously published (10); none of the false positives from the earlier ChIP-chip data were peaks in the new data. Of these 101 peaks, 99 (98%) were validated. Figure 1 shows the qPCR results for 38 GATA1 ChIP-chip peaks not previously tested; all but two are validated. Importantly, the peak calls that were uniquely identified by each of the programs (see ‘Materials and Methods’ section) are also validated at a high rate, and thus we pooled all the peak calls into a set of 302 occupied DNA segments.

![Figure 1](https://example.com/figure1.png)

Figure 1. Relative enrichment in ChIP material from induced G1E-ER4 cells for the sampled ChIP-chip positive hits tested by quantitative PCR (qPCR). The grey bars are the controls for qPCR assay. The green bars are the positive ChIP-chip peaks called by at least two out of the three peak calling programs (see ‘Materials and Methods’ section). The blue bars are the peaks only called by program PASS. The salmon bars are the peaks called only by program Mpeak. The purple bars are the peaks called only by Tamalpais. The line is drawn at the mean relative enrichment of the negative controls plus two standard deviations, which is used as the threshold for qPCR validation.
Based on the qPCR validation rate of 98%, we estimated that 296 segments are occupied by GATA1 in this 66 Mb region. The non-repetitive portion of the 66 Mb region on mouse chromosome 7 was divided into 67,420 intervals of 500 bp each in which a ChIP-chip peak could be found (the ‘chip-able’ portion, see Supplementary Data ‘Segmentation of interrogated ChIP regions into 500 bp windows’). Of these intervals, 43,594 contain at least one WGATAR motif. Because our goal is to identify and test motifs enriched in the GATA1-occupied segments, we selected three background training sets and one background testing set for each run of DME2. In each run, we evaluated the occurrence of all possible matrices of size 6 for over-representation in the foreground training set compared with one of the background training sets. Motifs that are similar in the over-representation score and matrix composition in the three runs of comparisons using DME2 were combined and reported as the enriched motifs in the GATA1-occupied segments. These motifs were evaluated for their ability to discriminate occupied segments from unoccupied ones using the testing datasets. (B) Distribution of G+C content and occurrences of WGATAR motif in the training datasets and (C) in the testing datasets. The foreground training set and the background training sets have similar G+C content and occurrences of WGATAR motif; so do the foreground and background testing sets.

Figure 2. Identification and evaluation of motifs enriched in the GATA1-occupied segments. (A) The overall procedure for identifying and testing motifs enriched in GATA1-occupied segments is illustrated. The 314 DNA intervals of 500 bp that are occupied by GATA1 comprise the foreground set. It was randomly split into two groups of 157 bound intervals, one used for the identification of enriched motifs and the other used for evaluation of the enriched motifs. Three background training sets and one background testing set were selected. In each run of DME2, the occurrence of all possible matrices of size 6 was evaluated for over-representation in the foreground training set compared with one of the background training sets. Motifs that are similar in the over-representation score and matrix composition in the three runs of comparisons using DME2 were combined and reported as the enriched motifs in the GATA1-occupied segments. These motifs were evaluated for their ability to discriminate occupied segments from unoccupied ones using the testing datasets. (B) Distribution of G+C content and occurrences of WGATAR motif in the training datasets and (C) in the testing datasets. The foreground training set and the background training sets have similar G+C content and occurrences of WGATAR motif; so do the foreground and background testing sets.

DNA segments with potential binding sites are not occupied indicates that the ChIP data are highly specific.

Significantly enriched motifs in DNA segments occupied by GATA1

The 302 GATA1-occupied segments were divided into 314 DNA segments of 500 bp; these constitute the foreground set. Half of these comprise the foreground training set used to identify sequence patterns that might contribute to the in vivo occupancy of GATA1 (Figure 2A), and the other half is used for testing. Almost all (91%) segments comprising the foreground set contain at least one instance of the motif WGATAR. Because our goal is
to find additional, more specific, discriminative patterns, we required DNA intervals in the training and testing background (unbound) sets to have the same distribution of WGATAR occurrences as in the foreground sets (Figure 2B). The background sets were also matched to the foreground sets in the distribution of G + C content.

DME2 (26) was used to evaluate the over-representation of motifs in the foreground training set compared with each of three background training sets independently (Figure 2A). Twenty-nine motifs were found in all three runs of DME; these are the discriminatory motifs that are robust to the choice of the background set (Table 1). These motifs all had similar levels of enrichment in the foreground compared with each of the background sets, as given by the enrichment score (Table 1).

We also analyzed the sequences of the GATA1-occupied DNA segments using other published motif discovery tools, namely YMF (34), DEME (35), Weeder (36), MEME (37), AlignACE (38), CLOVER (39). They all identified words that match the WGATAR binding site motif for GATA1, and some also discovered GC-rich stings (see Supplementary Data, ‘Motifs identified by direct enumeration of words’).

Table 1. Motifs discovered by DME2 as discriminating GATA1-bound from unbound DNA segments

| Consensus motif (combined from three runs) | Enrichment scorea | S0  | Sp  |
|-------------------------------------------|-------------------|-----|-----|
|                                           | Run 1             | Run 2 | Run 3 |
| GMTAAG                                    | 342 460 413       | 0.516 | 0.726 |
| AGATTA                                    | 219 266 358       | 0.331 | 0.86  |
| AGATAA                                    | 196 254 196       | 0.637 | 0.548 |
| GTTAA                                     | 130 189 94        | 0.217 | 0.885 |
| TTTGAG                                    | 156 156 108       | 0.293 | 0.79  |
| AAAAAAC                                   | 115 104 81        | 0.369 | 0.688 |
| TAAACA                                    | 94 189 201        | 0.261 | 0.79  |
| AAGATA                                    | 231 162 81        | 0.408 | 0.643 |
| GAAAGA                                    | 212 118 94        | 0.503 | 0.541 |
| GTAACT                                    | 71 118 47         | 0.185 | 0.854 |
| GGTTAG                                    | 265 217 229       | 0.166 | 0.866 |
| CAGTTA                                    | 224 177 141       | 0.229 | 0.803 |
| CAACAG                                    | 84 60 24          | 0.293 | 0.732 |
| GCTTAC                                    | 153 153 200       | 0.153 | 0.854 |
| AGCACA                                    | 24 48 24          | 0.389 | 0.599 |
| AACAGG                                    | 168 181 132       | 0.293 | 0.688 |
| ATACAG                                    | 71 59 59          | 0.223 | 0.739 |
| ACTCTA                                    | 24 130 71         | 0.178 | 0.777 |
| GATACA                                    | 130 118 189       | 0.14  | 0.803 |
| AACTCA                                    | 130 200 248       | 0.217 | 0.72  |
| Highly similar in three runs              |                   |      |
| CCGCC                                     | 236 311 261       | 0.312 | 0.866 |
| CCGGCC                                    | 187 303 239       | 0.783 | 0.338 |
| GCCAGG                                    | 340 402 288       | 0.688 | 0.401 |
| GCCCG                                     | 179 71 200        | 0.586 | 0.484 |
| GAGGCC                                    | 88 88 137         | 0.586 | 0.427 |
| GGCAGG                                    | 138 376 242       | 0.386 | 0.395 |
| GCTCAG                                    | 85 19 73          | 0.51  | 0.465 |
| GGAAG                                      | 194 217 205       | 0.414 | 0.548 |
| GGAAGT                                    | 193 132 212       | 0.49  | 0.433 |

Motifs that are not effective discriminators against an independent testing set are shaded in grey.

The enrichment score is the relative over-representation score returned by DME2 (26).

Discriminatory power of each enriched motif

The effectiveness of each motif discovered by DME2 was evaluated by calculating its sensitivity and specificity against the testing sets of foreground and background segments (note these are independent sets, i.e. the testing sets do not overlap with the training sets; Figure 2A and C). Both the sensitivity and specificity of enriched motifs cover wide ranges (Table 1). The discriminatory performance of the motifs can be visualized in an ROC-type graph (Figure 3A), in which sensitivity is plotted against 1-specificity. An ideal discriminator would produce a point in the extreme upper left (0,1), while points along the diagonal reflect no discriminatory power. As shown in Figure 3A, the motifs represented by the consensus sequences GATAAG and AGATAA have the best combination of sensitivity and specificity, capturing 52–64% of the occupied DNA segments while rejecting 55–73% of the unbound DNA segments. In contrast, several of the motifs enriched in the foreground training set do not discriminate effectively in the testing sets; these are represented by the black dots to the right of the diagonal in the ROC graph (Figure 3A), and they have a light gray background in Table 1. As expected, all 29 motifs discriminate effectively in the training sets (Supplementary Figure S1); the failure of some to discriminate in the testing sets illustrates the need to use independent testing sets for evaluation.

Enriched motifs matching known transcription factor binding sites

Having found a collection of enriched motifs associated with GATA1 occupancy (and validated against the testing set), we searched for matches to known binding sites for mammalian transcription factors (TFBs) as compiled in the Jaspar library (31), in a collection of non-redundant motifs (32) compiled from the TRANSFAC library (33), or in a set of custom motifs for binding sites for erythroid transcription factors, such as EKLF binding sites and CP2 half sites (see ‘Materials and Methods’ section). After filtering to remove motif matches that were not consistent with nucleotides known to be critical for binding (see ‘Materials and Methods’ section), TFBS motifs for six families of proteins remained (Table 2). These include not only GATA1, but also the binding site motifs for proteins previously demonstrated to interact with GATA1, such as EKLF, Sp1 and CP2 (40,41) (see ‘Discussion’ section).

Preference for specific variants of WGATAR for occupancy in vivo

The distribution of WGATAR motifs in the DNA segments comprising the background training sets is the
same as in the foreground sets. Therefore, we initially were surprised to find a variant of the motif, AGATAA, in the list of highly enriched motifs that were validated against the testing set (Table 1). This observation suggested that this particular variant of the WGATAR motif is preferred for binding in vivo. Further analysis strongly supports this conclusion. The AGATAA variant outnumbers the other three variants of WGATAR in the foreground set, comprising 40% of all instances of WGATAR. Also, 69% of the occupied segments contain this motif. In contrast, AGATAA is less dominant in the unoccupied segments, comprising 33% of all instances of WGATAR. Only 46% of the unoccupied segments contain it (Table 3).

The prevalence of variants of WGATAR was also examined in a dataset compiled from a collection of literature reports on 36 human and mouse erythroid cis-regulatory modules (CRMs) that have been shown to be bound by GATA1 in vivo by ChIP (see Supplementary Data, ‘Curated datasets of in vivo occupied segments by GATA1’ and Supplementary Table S5). None overlap with our 314 foreground intervals. In these occupied DNA segments, AGATAA outnumbers the other variants of WGATAR, comprising 42% of the instances of WGATAR and occurring in 62% of the occupied segments or erythroid CRMs (Table 3). In addition, the variant TGATAA also occurs frequently, comprising 35% of the instances of WGATAR in this dataset and occurring in 66% of the occupied segments with a WGATAR (Table 3). These observations indicate that the two purine nucleotides are not equally preferred in the sixth position of the motif in vivo, and that WGATAA is a better consensus than WGATAR for predicting in vivo occupancy.

![Diagram](https://example.com/diagram.png)

**Figure 3.** Evaluation of the discriminatory power of enriched motifs, motif combinations and histone modifications in the testing set. This graph takes the form of a receiver–operator characteristic (ROC) plot. (A) The sensitivity and (1-specificity) is plotted for the 29 enriched motifs discovered by DME2 (Table 1, black dots), the 10 combinations of enriched motifs that have the top performance (red dots), the 10 prominent TFBSs (the 7 TFBSs in Table 2 column 5 plus the general WGATAR consensus GATA1 binding site motif and two specific variants, WGATAA and AGATAA; blue dots), combinations of TFBSs (green dots), and histone modification status (orange dots, labeled C for low H3K27me3 and D for high H3K4me1). The dots for several key motifs and combinations are designated by their consensus sequence. The performance for combinations containing matches to TFBS motifs are labeled with the transcription factor followed by the suffix ‘bs’, e.g. SP1_bs. (B) The sensitivity and (1-specificity) is plotted for all combinations of two to four enriched motifs discovered by DME2 (grey dots). The top 10 motif combinations with best performance are colored red.

**Combinations of motifs improve discrimination**

Each discriminatory motif with high specificity but low sensitivity (producing the black dots in the lower left of ROC graph in Figure 3A) tends to capture a different set of GATA1-bound DNA segments. Thus evaluating the union of the intervals containing the discriminatory motifs is expected to improve discrimination. Therefore we computed sensitivity and specificity against the testing datasets for all combinations of two to four of the 29 motifs (Figure 3B). In this analysis, sensitivity is the fraction of bound DNA intervals in the foreground testing set that contains any one or more of the motifs in a combination. Specificity is the fraction of unbound intervals in the background testing set that contains none of the motifs in a combination. As expected, the motif combinations evaluated by this method cover a broad area of the ROC graph, and the greatest density is in the higher sensitivity, lower specificity portion. No discrete group of motif combinations stands out from the rest, but rather the better performing combinations are part of a continuum of discrimination. This indicates that no small group of motifs is substantially better than others for distinguishing GATA1-bound from unbound DNA. While we will focus on the top 10 combinations for further analysis, it is important to note that many other combinations are almost equivalently good.

The top 10 motif combinations with the best performances (colored in red in Figure 3A and B, and listed in Supplementary Table S7) are all distributed along the upper left boundary of the cloud in the ROC graph.
plot in Figure 3B; in fact, they are better discriminators than any of the single motifs shown in Figure 3A. Almost all the top combinations contain at least one match to the preferred GATA1 binding site motif (e.g. AGATAA, GATAAG, AAGATA). Another motif occurring frequently in the top combinations is GGGCGG, which is similar to the binding site motif for SP1 (GGGCGGR). Thus a particularly effective combination that discriminates occupied from unoccupied segments is the presence of any of the three motifs GATAAG, AGATTA, or CCCCCC, as shown by motif combination B in Figure 3A.

We also evaluated the discriminatory power of the TFBS motifs that match the enriched motifs discovered by DME2 (Table 2) and combinations of those TFBSs. As shown in Figure 3A, no single TFBS motif is an effective discriminator for GATA1 binding. Because WGATAR is almost ubiquitous in the foreground testing set (91%) and is required in the background testing set, it has no discriminatory power and sits right on the diagonal (Figure 3A). However, two more specific variants of WGATAR, i.e. WGATAA and AGATAA, do have some discriminatory power, achieving some specificity at the cost of reduced sensitivity (Figure 3A). No other single TFBS motif has better discriminatory power; as with the DME2-discovered motifs, they tend to capture different sets of bound intervals with high specificity (blue dots in the lower left in Figure 3A). After pairing a WGATAR motif with a second motif of either an AGATAA motif or the TFBS motif for SP1 or the TFBS motif for EKLF, 62% of the occupied DNA segments can be captured, and 57% of the unoccupied DNA segments can be rejected (dot A in Figure 3A).

Good prediction of occupancy by multiple instances of WGATAR

Given that multiple motifs can improve discrimination, we wished to examine how well multiple instances of WGATAR would predict occupancy. This cannot be addressed by comparisons to the background training and testing sets, because these were selected to match the foreground sets in terms of the distribution of WGATAR occurrences (Figure 2). Thus we compared the distribution of WGATAR occurrences in the 314 GATA1-occupied DNA segments with that in all the 67,106 unoccupied DNA segments in the non-repetitive portion of the interrogated region of chromosome 7. The average number of instances of the WGATAR motif is higher in the occupied DNA segments (2.3 compared with 1.1 in the unoccupied DNA segments, Table 3). Furthermore, the distribution of WGATAR occurrences per DNA segment is shifted considerably higher for occupied versus unoccupied DNA segments (Figure 4). The difference between the distributions is significant by both a Student’s $t$-test and a Wilcoxon rank order test ($P$-values < 2.2 x 10^{-16}). In particular, a much higher fraction of GATA1-bound DNA intervals has multiple WGATAR motifs (67%) compared with unoccupied intervals (31%). Thus multiple occurrence of WGATAR is a feature with strong predictive power.

### Table 2. TFBS motifs significantly enriched in the GATA1-bound intervals

| Protein family | Core binding consensus | Matched words | Candidate TF | Known consensus |
|----------------|------------------------|---------------|--------------|-----------------|
| GATA           | GATA                   | AGATAA, GATAAG, AAGATA | GATA         | WGATAR          |
| KLF            | GC-box                 | CGCCCGG       | SP-1 or EKLF | YCCGCCG or CCNCMCCCW |
| Grainyhead     | CCWG                   | GCCAGC, GCCCGC | CP2          | CCWG            |
| Winged helix   | GTTA                   | AGTTTAG, GTTTAG | HNF1-like    | RGT TAMWNATT    |
| IRF            | GAAANN                 | GAAAGA        | ICSBP        | AAANYGAAAS      |

### Table 3. Presence of all WGATAR variants within in vivo GATA1-occupied DNA segments

| Motifs | 314 GATA1-occupied intervals | 67,106 unoccupied intervals | Curated in vivo bound intervals (36) |
|--------|-------------------------------|---------------------------|-------------------------------------|
|        | Occurrences of motif | Segments with motif | Occurrences of motif | Segments with motif | Occurrences of motif | Segments with motif |
| AGATAA | 294 | 200 | 24,865 | 20,122 | 25 | 18 |
| AGATAG | 167 | 116 | 16,687 | 14,413 | 9 | 6 |
| TGATAA | 180 | 138 | 20,197 | 16,675 | 21 | 19 |
| TGATAG | 93 | 83 | 14,524 | 12,895 | 5 | 5 |
| WGATAA | 474 | 257 | 45,062 | 31,115 | 46 | 26 |
| WGATAR | 734 | 288 | 76,273 | 43,306 | 60 | 29 |
Biologically, this observation is consistent with the reported self-association of the GATA1 protein (42,43).

### Histone modifications around GATA1-occupied DNA segments

Given the strong role played by histone modifications in transcriptional regulation, we compared the modification status between GATA1-occupied and unoccupied DNA segments. Along the 66 Mb region of mouse chromosome 7 in G1E-ER4 cells, we measured by ChIP-chip the level of H3K27me3 (associated with repression of expression) (2), and H3K4me1 (associated with enhancer activity) (3).

In aggregate, DNA segments in the foreground training sets are strongly depleted for H3K27me3 for at least 2 kb on either side of the WGATAR motifs (Figure 5, upper panel). A small positive signal for H3K27me3 is observed for the WGATAR motifs in the background training sets. In striking contrast, the GATA1-occupied segments comprising the foreground training set (and surrounding sequences) are strongly enriched for H3K4me1, whereas the unoccupied segments in the background training sets show low signals (Figure 5, middle panel). For comparison, the aggregated hybridization signal for GATA1 ChIP-chip shows a peak at the WGATAR motif in the occupied DNA segments but a uniformly low signal throughout the unoccupied DNA segments (Figure 5, lower panel). Hybridization signals are low for all three features around randomly sampled positions in the 66 Mb region of chromosome 7 (Figure 5).

The substantial difference in histone modification status observed in aggregate in Figure 5 is also observed in individual DNA segments. Two examples illustrate the high level of H3K4me1 and depletion of H3K27me3 in a GATA1-occupied DNA segment (Figure 6A) and the

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Distributions of the frequency of occurrence of a WGATAR motif in GATA1-occupied and unoccupied DNA segments. The number of occurrences of the WGATAR motif in each of the 314 occupied segments and the 67,106 unoccupied segments in the 66 Mb locus was counted. The distributions of these occurrences are shown as box-plots for each class of DNA segments. The differences between occupied and unoccupied DNA segments are significant by Student’s $t$-test and Wilcoxon Rank Order test, both with $P$-values much <2.2E–16.

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Histone modifications around DNA segments occupied and unoccupied by GATA1. Each graph plots the mean of the ChIP-chip hybridization signal for tri-methylated histone 3 lysine 27 (top panel), mono-methylated histone 3 lysine 4 (middle panel), or GATA1 (bottom panel) for GATA1-occupied DNA segments in the foreground training set (red) or unoccupied DNA segments in the background training sets (blue). Each DNA segment was centered on a WGATAR motif, the hybridization signal was aggregated in 50 bp bins extending for 2000 bp on each side for all DNA segments in the designated class; the mean for each bin is plotted in the graph. The mean hybridization signals for each feature around random sampled genomic locations in the 66Mb region on chromosome 7 are shown in grey.
opposite pattern in an unoccupied DNA segment (Figure 6B). While both DNA segments contain four WGATAR motifs, the presence of a match to an EKLF binding site motif and characteristic histone modifications distinguish the occupied from unoccupied DNA segment. The histone modification status distinctive for occupied versus unoccupied DNA segments is observed consistently across the testing datasets. The distribution of mean H3K27me3 signal is significantly lower for the occupied DNA segments in the foreground testing set compared with the unoccupied ones in the background testing set (one tailed Student’s \( t \)-test, \( P \)-value \( < \times 10^{-16} \)).

### Histone modifications provide strong discriminatory power

The dramatic separation between GATA1-occupied and unoccupied DNA segments for H3K4me1 signal suggests that it should have strong discriminatory power for occupancy. Based on the distributions shown in Figure 6D, we partitioned DNA segments into those with a high average H3K4me1 signal (average hybridization signal of at least 1) versus those with lower signal. Eighty-two percent of the GATA1-occupied segments, and only 12% of the unoccupied DNA segments containing WGATAR motifs, were labeled as high H3K4me1 (Table 4). Indeed, H3K4me1 is the best single discriminatory feature examined in this study; sensitivity and specificity for this feature on the testing foreground and background sets are compared with those for other features in Figure 3 (dot D). The level of H3K27me3 signal (partitioning DNA segments into two classes, those with a mean signal \( \geq 0 \) and those with a mean signal of at least 0, chosen to obtain good separation based on the distributions shown in Figure 6C) provides a discriminatory power comparable to the primary sequence motifs (dot C in Figure 3). These associations are also observed when the chromosome was segmented based on histone modification status by applying hidden Markov model approaches, which are not dependent on an arbitrary threshold (Supplementary Data,chr7:

### Table 4. GATA1 occupancy is associated with different chromatin status

| Histone modification status | Number of occupied DNA segments (foreground testing set, 157 total) | Number of unoccupied DNA segments (background testing set, 157 total) |
|----------------------------|---------------------------------------------------------------|---------------------------------------------------------------|
| H3K4me1 +                  | 129                                                           | 19                                                           |
| H3K27me3 +                 | 44                                                            | 79                                                            |

A + indicates a DNA segment with a mean hybridization signal of at least one for H3K4me1 or at least zero for H3K27me3.
The initial number of DNA intervals (set 1) is the number of ∼500 bp windows in the non-repetitive portion of the 66 Mb locus (see ‘Materials and Methods’ section). The intervals that have one or more matches to the GATA1-bs are set 2. Intervals were then searched for those with matches to the additional binding site motifs (Set 18) returns a set of DNA fragments of which 43 594 contain at least one WGATAR motif (Set 2). This has almost all the DNA intervals occupied by GATA1 (288 of the 314), but of course the vast majority of such intervals are not occupied (Figure 7, upper graph). Thus the positive predictive value (PPV) of a WGATAR motif is quite low. Requiring additional TFBS motifs reduces the number of occupied segments captured (Sensitivity, middle graph in Figure 7), but the reduction in total number of intervals is greater (lower graph in Figure 7) so that the fraction of intervals bound by GATA1 increases (upper graph in Figure 7). Allowing any from a combination of motifs substantially increases the number of occupied segments captured, with only a small reduction in the PPV.

Requiring a low level of H3K27me3 captures even more occupied segments than does combinations of motifs, but the fraction of intervals occupied is quite low. Requiring a high level of H3K4me1 is very effective, with sensitivity better than any combination of motifs and a PPV substantially higher than any other feature. Requiring both epigenetic marks further improves the PPV, but with some loss of sensitivity. This trend is even more pronounced when both diagnostic epigenetic marks and a motif combination are required (green bars in Figure 7). The best balance of PPV and sensitivity is obtained with H3K4me1 and motif combinations. The 1669 segments meeting these criteria captured 162 occupied segments (Sensitivity = 0.52, Set 15 in Table 5). This set of DNA segments gives one true occupied segment in about 5 intervals with a WGATAR being occupied.

DISCUSSION

Primary sequence determinants of occupancy by GATA1

The protein GATA1 has a high affinity in solution for sequences containing the motif WGATAR, but previous studies left it unclear whether this was true for segments occupied in vivo. Our results show that the WGATAR motif is significantly associated with occupancy: 91% of the occupied segments have the motif and unoccupied segments have substantially fewer of these motifs. Many mutagenesis studies have also shown the importance of the WGATAR motif for regulated expression of reporter genes (13,22,23,42–44). These results, combined with the in vitro affinity and our demonstration of the motifs in the preponderance of segments occupied in vivo, make a strong case for the WGATAR motif as a critical determinant of in vivo binding by GATA1. Furthermore, our studies show that two variants of this motif, AGATAA and TGATAA, are more frequently found in...
the \textit{in vivo} bound segments than are the motif variants that end in G.

However, the presence of the WGATAR motif is not sufficient to determine occupancy by GATA1 in erythroid cells. In G1E-ER4 cells, only about one in 147 DNA segments on chromosome 7 that potentially could be occupied actually are bound by GATA1 \textit{in vivo}. Some of the other DNA segments may be occupied by GATA1 in myeloid lineages, not in erythroid cells, presumably regulating genes that are specifically activated or repressed by GATA1 in those other cell types. Thus some of the specificity could be determined by combinatorial actions of different transcription factors with GATA1. Our investigation of motifs associated with occupancy supports this model. Some of the most frequently occurring motifs are good matches for the half-site for binding CP2, which has been shown to participate with GATA1 in regulation by multiple erythroid CRMs (40). Other enriched motifs match to the binding sites for either Sp1 or EKLF. These are Krüppel-like zinc finger proteins that have been shown to interact with GATA1 and to play important roles in erythroid regulation (41).

Another major determinant of occupancy by GATA1 is the presence of multiple WGATAR motifs. This suggests that multiple molecules of GATA1 may be bound to such segments \textit{in vivo}. GATA1 is known to self-associate through its zinc finger domains (45), and a mutant GATA1 defective in self-association activity can only partially rescue the \textit{Gata1–0.5} mutant mouse (46). This shows that the specific interaction of multiple GATA1 protein molecules is needed for the regulatory activity of GATA1. The fact that most of the occupied segments also have multiple binding motifs suggests that this self-association \textit{in vivo} may be driven both by multiple binding sites and specific protein-protein interaction domains.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure7.png}
\caption{Ability of primary sequence and epigenetic signals to determine occupancy of DNA by GATA1. Beginning with the 67,420 DNA segments of length 500 bp in the non-repetitive portion of the 66 Mb region of mouse chromosome 7, groups of DNA segments having the feature(s) listed along the x-axis were examined for the fraction of the DNA segments bound by GATA1 (top panel), the number of GATA1-occupied DNA segments (GATA1os) captured within the group (middle panel; maximum is 314), and the total number of DNA segments in the group (bottom panel). The features are binding site motifs recognized by transcription factors (labeled GATA1-bs, e.g.) and combinations of motifs (red bars), histone modifications (blue bars), and combinations of histone modifications and motifs (green bars).}
\end{figure}
Epigenetic determinants of occupancy by GATA1

Combinations of WGATAR with other TFBS motifs can achieve an improvement in accuracy of determining GATA1 occupancy in vivo. Combinations of motifs can restrict the potential targets to one true occupied segment in 75 segments with combinations of the discriminatory motifs identified in this study (Table 5, set 9). In order to find additional signals that allow the protein GATA1 to distinguish the one real binding site among the 75 with multiple motifs, we examined histone modifications in chromatin. It is possible that the majority of the GATA1-occupied DNA segments are in regions of the chromosome that are active in erythroid cells. If so, then these regions may be in an accessible chromatin conformation, and one would expect to find chromatin marks associated with open chromatin in these regions. Our studies confirm this.

Mono-methylation of H3K4 is known to be associated with enhancers and open chromatin (3). We find that this is the feature with the strongest discriminatory power of any single feature we have examined. In contrast to the GATA1 occupancy signals that are centered and most enriched right at the WGATAR motifs in the occupied segments (Figure 6, lower panel), signals of histone modifications cover a larger region (at least 2000 bp away from the centered position). One interpretation of these data is that a small set of DNA segments are marked by histone modifications that directly or indirectly facilitate binding of sequence-specific binding proteins such as GATA1 to DNA. These chromatin marks may be in place early in a differentiating cell lineage. In future experiments, it will be informative to examine the histone modification status of chromatin in G1E cells prior to restoration and activation of GATA1. Some of the DNA in the H3K4me1 regions may have that status prior to restoration of the transcription factor. This would support a role for the epigenetic marks in directing the selective binding of GATA1.

We note that one contributor to the effective discrimination by H3K4me1 is the large number of DNA segments occupied by GATA1. Other transcription factors that bind much less frequently may not show this discriminatory power of histone modification because a smaller fraction of the modified segments will be occupied.

Comparison with other enumeration-based motif discovery tools

The challenge in using enumeration methods for motif discovery is to adequately evaluate the significance of the enrichments obtained after the enumeration process. The enrichments are computed based on motif frequencies in a foreground set compared with some background set. DME2 (26) enumerates all possible score matrices of a pre-defined length in both a foreground and a background dataset and then optimizes the over-representation score, but it makes no assumptions about the background distributions. In contrast, several other enumeration-based methods compare the observed frequencies with those predicted by theoretical distributions, such as hypergeometric (47) or binomial (48). The YMF program (34) enumerates all possible words up to a certain length and compares the observed occurrences with simulations of background sequences based on Markov models. While these approaches have merit, their utility may be limited by how well the chosen distributions fit the real background.

Another positive feature of DME is the flexibility in choosing background sets for finding discriminatory motifs. Almost all the segments occupied by GATA1 have the consensus motif WGATAR, and this motif is common in the bulk genomic DNA segments. Thus we customized our used of DME by requiring background DNA segments not only to be unoccupied, but also to contain this primary motif. This approach facilitates the discovery of additional motifs. Using the DME pipeline, we found 19 motifs with significant discriminatory power that were also validated against a testing set. Furthermore, many of them matched binding sites for transcription factors previously implicated in working with GATA1 in regulating gene expression. Many of the other methods we used, whether enumeration-based or probabilistic, only used the foreground set of intervals, and invariably these returned few fewer discriminatory motifs. Thus the ability to input user-defined foreground and background datasets is an advantage in motif discovery.

Novel patterns

About half of the enriched, validated motifs do not match to a known transcription factor binding-site in our compiled library. The lack of a TFBS match may reflect the limitation of current knowledge on TFBSs, or the enriched motifs may be derived from functional sequence patterns other than binding sites, for example, partial sequences of microRNAs. Further investigation of these novel motifs could lead to new insights into determinants of specificity of GATA1 occupancy (e.g. binding site motifs for a protein that co-binds with GATA1) and mechanisms of GATA1-dependent regulation of gene expression.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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