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Identification of a binding motif specific to HNF4 by comparative analysis of multiple nuclear receptors

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ABSTRACT

Nuclear receptors (NRs) regulate gene expression by binding specific DNA sequences consisting of AG[G/T]TCA or AGAACA half site motifs in a variety of configurations. However, those motifs/ configurations alone do not adequately explain the diversity of NR function in vivo. Here, a systematic examination of DNA binding specificity by proteinbinding microarrays (PBMs) of three closely related human NRs-HNF4a, retinoid X receptor alpha (RXRa) and COUPTF2-reveals an HNF4-specific binding motif (H4-SBM), xxxxCAAAGTCCA, as well as a previously unrecognized polarity in the classical DR1 motif (AGGTCAxAGGTCA) for HNF4a, RXRa and COUPTF2 homodimers. ChIP-seq data indicate that the H4-SBM is uniquely bound by HNF4 α but not 10 other NRs in vivo, while NRs PXR, FXR α , Rev-Erb α appear to bind adjacent to H4-SBMs. HNF4-specific DNA recognition and transactivation are mediated by residues Asp69 and Arg76 in the DNA-binding domain; this combination of amino acids is unique to HNF4 among all human NRs. Expression profiling and ChIP data predict \sim 100 new human HNF4 α target genes with an H4-SBM site, including several Co-enzyme A-related genes and genes with links to disease. These results provide important new insights into NR DNA binding.

INTRODUCTION

Nuclear receptors (NRs) are ligand-dependent transcription factors (TFs) that play important roles in nearly every aspect of human physiology (1). They are linked to human disease, are popular drug targets and play a major role in regulating the expression of genes responsible for drug metabolism (2-4). Therefore, proper elucidation of their target genes has important clinical ramifications. NRs regulate gene expression primarily by first binding specific DNA response elements in the regulatory regions of target genes. Basic rules for NR DNA binding that consist of distinct half sites motifs for steroid (AGAACA) and non-steroid (AGGTCA) receptors and half site configurations—direct (DR, $\rightarrow \rightarrow$), inverted (IR, $\rightarrow \leftarrow$), everted repeats (ER, $\rightarrow \leftarrow$) and non-repeats (nRs) as well as spacing between the repeats—were established early on (5-8). This dogma, the 'DR rule', successfully drove the identification of individual NR target genes in the pre-genomic era. However, there remains considerable overlap in the apparent binding specificity of many NRs, which could be due to the high degree of conservation among the NR DNA-binding domains (DBDs) and/or to the fact that NR DNA binding specificity has not been analyzed in a systematic, global fashion.

The magnitude of the complexity and intricacy of DNA binding specificity is illustrated by the fact that for a 13-nt-long motif, such as a direct repeat of AGGT CA with a spacing of 1 nt (DR1, AGGTCAxAGGTCA), there are $4^{13}/2$ (~34 million) different potential DNA sequences. If a given NR binds just 0.01% of those sites for a specificity of 1 in 10000, that still yields 3400 unique sequences, all of which could have just a minor variation on the DR1 consensus. The question then arises as to how many of those different sequences will bind a given NR and whether there are motifs specific to different NRs.

There are three well-characterized subfamilies of receptors that bind DR1s as homodimers—HNF4 α (NR2A1), COUPTF2 (NR2F2) and RXR α (NR2B1). They are all very highly conserved, closely related and regulate a variety of metabolic genes in common tissues such as

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The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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liver, kidney and intestine (9,10). HNF4 α is a constitutive activator that binds an endogenous ligand (linoleic acid) (11). It is considered to be a master regulator of liver-specific gene expression, including genes involved in intermediary metabolism as well as xenobiotic and drug metabolism (12–15). HNF4 α is linked to several human diseases including diabetes, hemophilia, hepatitis, atherosclerosis and inflammatory bowel disease (16,17).

RXR α is also expressed primarily in the liver, as well as kidney, gut, muscle and skin (10). While it is best known as a heterodimeric partner of other NRs such as PPAR (NR1C), RAR (NR1B), FXR (NR1H), PXR (NR1I2), TR (NR1A) and VDR (NR1I1) (7,18), it binds directly to the synthetic ligand 9-cis retinoid acid and has been shown to activate transcription in the absence of an ectopically expressed partner (19-23). COUPTF2 is also present in the liver and, like $RXR\alpha$, is fairly ubiquitously expressed. However, it acts primarily as a repressor of transcription (24,25); it remains an orphan receptor in that its endogenous ligand has not yet been identified, although it has been shown to bind and respond to retinoids (26). Like HNF4a, COUPTF2 binds DNA well as a homodimer but unlike HNF4a it can also heterodimerize with RXR α (21,25,27). It was recognized early on that HNF4a, RXRa and COUPTF2 all share common binding sites (that roughly resemble DR1s) in the promoters of certain genes and consequently compete for regulation of those genes (28-30). However, it was also noted that there are a couple of binding sites in other genes that were not bound by all three NRs (28,31,32). While these results suggested the existence of NR-specific binding motifs, common versus unique binding features were never identified and the extent of the overlap remained obscure.

New genome-scale technologies, both in vivo and in vitro, now allow us to address the issue of NR binding specificity in an appropriately global fashion (33). For *in vivo* binding, chromatin immunoprecipitation followed by deep sequencing (ChIP-seq) enables robust and accurate identification of TF binding regions, but the resolution is not sufficient to identify the exact site to which TFs bind; this must be done by statistical inference (34). For in vitro binding, protein-binding microarrays (PBMs), can assay the binding of TFs to 10000 s of DNA probes in a high throughput fashion (35). PBM data are extremely useful for mining ChIP-seq data to identify the precise location of TF binding and to predict new target genes by cross-referencing. We recently applied a modified PBM approach to HNF4a and identified 100 s of new direct targets of HNF4 α by combining the PBM results with genome-wide location and expression profiling data (13).

Here, we apply the PBM approach to the problem of distinguishing the binding specificity of HNF4 α , RXR α and COUPTF2. We show that RXR α homodimers bind DNA very well in the PBM and that they have a specificity nearly identical to COUPTF2 homodimers, with a clear preference for the 3' half site of a DR1 motif. The PBMs also identified an HNF4 α -specific binding motif (H4-SBM) (CAAAGTCCA) that was verified *in vivo* in ChIP-seq data. Finally, we determine the amino acid residues in the HNF4 α DBD (Asp69 and Arg76) responsible for the unique binding specificity of HNF4 α and identify ~100 new human target genes that contain the HNF4–specific motif. These results have important implications not just for HNF4 α , RXR α and COUPTF2, but for the entire NR superfamily as well.

MATERIALS AND METHODS

PBM design, assay and data processing

Two different custom arrays in 8×15 k format ordered from Agilent Technologies (Santa Clara, CA, USA) were used to test binding specificity (PBM2) (13) and half site preferences (PBM6.1) (see Supplementary Figure S2 for design and Supplementary Table S4 for complete sequences of PBM6.1). Each of the ~ 3000 sequences was spotted four (PBM6.1) to five (PBM2) times in each of the eight grids. The PBM assay was performed as previously described (13) with minor modifications: 1.6 µM Cy5 dUTPs (Enzo Life Sciences) was used to label the double-stranded DNA on the slide; \sim 0.8–1.2 µg of each full length NR in crude nuclear extracts from transfected Cos-7 cells was applied to the PBM; and bound NRs were detected with primary antibodies for human HNF4a, RXRa, COUPTF2 and RARa from R&D Systems (Catalogue # PP-H1415, PP-K8508, PP-H7147 and PP-H1920-00, respectively) at a 1:100 dilution overnight at room temperature, and then secondary antibody (Dam-Dylight-Cy3, Jackson ImmunoResearch) at a 1:50 dilution for 1 h. After three washes in PBS-0.1% Tween-20, the slide was dried and scanned at 633 (Cy5) and 543 nm (Cy3). Data extraction and normalization was performed as previously described (13). The binding threshold of each NR was set to two SD from the mean or the quantile 0.95 of the random controls, whichever was higher. DNA motifs (represented using position weight matrices, PWM) were generated using SeqLogo (36) and Weblogo v2.8.2 (37). Hypergeometric tests are performed using phyper in R. While the HNF4 α PBM2 was repeated, our previous raw data for the HNF4a PBM2 (13) was used for analysis since the signal intensities were closer to that of RXRα and COUPTF2.

Motif mining of ChIP-seq peaks

DNA sequences in ChIP-seq peak regions were extracted by Cisgenome and analyzed for relative enrichment level (fold enrichment) compared to matched control regions as described in (38). The length of each control region was determined by the average length of all peaks in a data set and the total number of control regions was five times as many as ChIP-seq peaks. If multiple data sets were available for an NR, the average was used as the final score and SD was given (Supplementary Table S1). The empirical P value of the enrichment score for each motif was determined using the enrichment distribution of 1000 6- or 9-nt random genomic sequences. Coordinates of peak centers were either given in the preprocessed BED or WIG files or obtained by reprocessing the data (in the case of ERR β and Rev-Erb α). Density plots were generated by R; variances between plots were calculated using a two-sided *F*-test. Motif mining was performed by Gibbs Motif Sampler (39) implemented in Cisgenome, using 3000 iterations and the third-order Markov Chain model. The best AGGTCA-like motif was selected from 10 candidates with the highest scores.

Identification of HNF4-specific target genes

DNA sequences from -10 to +10 kb of each gene in the human genome (hg19) were downloaded from the Ensembl genome browser (http://uswest.ensembl.org/ index.html) using BioPerl API. The occurrences of HNF4-specific binding sequences, obtained from either the PBM data or support vector machine (SVM) prediction (see details in Supplementary Data), were identified by Seqmap (40) using exact match. A total of 3000 random 13-mers were selected randomly from the human genome. The difference between two frequency distributions was calculated using the Student's t-test in R. The online search engine for HNF4-specific binding sequence was developed using Perl and cgi, and is hosted by the Bioinformatics Core of Institute for Integrative Genome Biology at University of California http://nrmotif.ucr.edu/NRBSScan/ Riverside at at H4SBM.htm. HNF4a ChIP-seq peaks from HepG2 cells (41) were examined and only those peaks containing at least one HNF4-specific binding site were selected as HNF4-specific peaks. If a gene had at least one HNF4specific peak within 10kb of its transcription start site (TSS), then this gene was considered as an HNF4specific candidate target. Only down-regulated candidate genes in HNF4 RNAi assay (in HepG2 cells) (13) were

chosen as final targets. Gene Ontology (GO) analysis was performed using DAVID (42).

RESULTS

PBMs reveal a polarity in the DR1 motif for RXR α , COUPTF2 and HNF4 α

Custom PBMs were used to examine the DNA binding specificity of human HNF4 α 2 (referred to as HNF4 α), RXRa and COUPTF2. They bound 1371, 1285 and 1530 unique DNA sequences, respectively, although there was considerable overlap, especially between RXRa and COUPTF2 (Figure 1A). Notably, while RXR α in the absence of a heterodimeric partner typically does not bind DNA in shift gels (22,23,27), it bound very well in the PBM in both the presence and the absence of its ligand 9-cis retinoid acid; control experiments verified that the amount of endogenous NRs such as RARa was too low to be detected in the PBMs (see Supplementary Figure S1 for details and Supplementary Table S4 for a list of all sequences). There are reports of RXRa activating transcription or binding DNA in the absence of added NR partners (19-23,43-45), and recent ChIP-seq experiments reveal considerable RXR binding in the absence of a PPAR partner (46), suggesting that RXR homodimers bind DNA in vivo as well as in vitro.

All bound sequences were ranked according to their normalized signal intensities (binding scores) and DNA motifs were generated for strong, medium and weak binders [binding affinity is linearly correlated with the binding signal from PBMs (47)]. Motifs of the top 10%



Figure 1. DNA binding specificity of human HNF4 α , RXR α and COUPTF2 determined by PBMs. (A) Work flow and overall results of PBM experiment. Total number of distinct DNA sequences (sites) bound by each NR is given as well as the number of overlapping and unique sequences (Venn diagram). (B) DNA motifs for the top 10% of binders as well as top 33% (strong), middle 33% (medium) and bottom 33% (weak) of binders. (C) Schematic representation of NR binding orientation on original and reverse complement (RC) DNA probes. Orange triangles, DBDs of 5' and 3' NR monomers. The canonical DR1 half site AGGTCA (red) is close to the free end of the original probe, while the corresponding sequence in the RC probe is close to the slide in the PBM. (D) Motifs of the 30 strongest and 30 weakest binders of the original or RC probes for COUPTF2 as indicated. (See Supplementary Figures S1B for the results of HNF4 α and RXR α , and Figure S2A for details in probe design.).

strongest binders showed that both the 5' and 3' half sites of the DR1 motif contain essential positions for binding, reinforcing the notion that these three NRs bind as homodimers on the PBM (Figure 1B). We also noted that the strong binders for RXRa and COUPTF2 had more conserved positions in the 3' than in the 5' half site, suggesting that the 3' half site may be more critical for DNA binding. Since, the 3' half site is farther from the glass slide in the PBM, it was possible that the observed polarity was an artifact of the PBM design. Therefore, we designed a second PBM in which 30 strong binders of COUPTF2 with AGGTCA half sites on the 3' side were compared with 30 weak binders with AGGTCA on the 5' side; a third group of probes had weak AGGTCA -like motifs in both half sites. The reverse compliment (RC) of each probe was also examined (Supplementary Figure S2A). If COUPTF2 DNA binding is affected by the distance from the slide, then the RC probes of corresponding strong binders will become weak binders, and vice versa (Figure 1C). We found that the strongest binders for both the original and the RC probes were the same sequences with a clear preference for AGGTC A in the 3' half site, even when it was close to the slide; conversely, the weakest binders had the AGGTCA motif in the 5' half site regardless of the distance to the slide (Figure 1D). Similar results were observed for RXRa and HNF4 α (Supplementary Figure S2B). We also examined the effect of the distance of the entire 13-mer from the slide as well as the free end of the probe and found that the strongest binding by all three NRs

resulted when the 13-mer is 27 nt from the slide and 5 nt from the free end of the probe (Supplementary Figure S2C and S2D). All together, these results suggest that *in vitro* RXR α , COUPTF2 and HNF4 α prefer the 3' half site of the DR1 motif.

PBMs identify an HNF4α-specific binding motif

The DNA binding specificity of HNF4a, RXRa and COUPTF2 was further analyzed using scatter plots (Figure 2A). RXRa and COUPTF2 shared very similar binding specificities with an R^2 of 0.84 for all bound sequences. In contrast, HNF4a showed a different profile with >200 DNA sequences bound exclusively by HNF4a. The consensus motif of HNF4-specific binders had a subtle vet consistent change at position 10 (p10) and p11, where GT was replaced by TC. Together with seven other conserved positions, it defined the HNF4specific binding motif (H4-SBM) xxxxCAAAGTCCA (x refers to any nucleotide; the nucleotides that differ from the canonical AGGTCA are underlined) (Figure 2B). Hypergeometric plots using all the binders indicate that TC at p10-p11 are disfavored by both RXRa and COUPTF2 (Figure 2C). Interestingly, the probes that were bound by RXRa and COUPTF2 but not HNF4a strongly resembled a canonical DR1 motif (Figure 2B).

We next determined that all 81 of the PBM probes bearing the H4-SBM were in fact HNF4-specific binders; they all bound HNF4 α but none bound RXR α or COUTF2 (Figure 2D, left panel). There



Figure 2. Identification of an HNF4-specific DNA binding motif. (A) Scatter plots of PBM results of the indicated NRs. Each spot is the average of \sim 5 replicates for each unique DNA sequence (\sim 3000 total) on the PBM; only probes with a coefficient of variation less than 0.3 were used. Each plot is divided into four quadrants by the binding thresholds of the NRs. Common binding sequences of RXR α and COUPTF2 show a linear correlation with a R^2 of 0.84. Specific targets of RXR α /COUPTF2 and HNF4 α are in the blue and red quadrants, respectively. (**B**) Shown are motifs derived from the RXR α /COUPTF2 and HNF4 α -specific sequences in the blue and red quadrants in the scatter plots in (A), respectively. (**C**) Hypergeometric plots showing the over- and under-representation for each nucleotide at pl to pl3 (in four colors) of RXR α and COUPTF2 in bound sequences on PBM. *Y*-axis indicates log(*P*-value) calculated using phyper test in *R*. Positive and negative values indicate over-representation and under-representation, respectively. The maximum log*P* is set as 50. As shown, 'TC' at pl0–pl1 is strongly disfavored by RXR α and COUPTF2. (**D**) Box plots showing rankings of all PBM probes containing the 9-nt sequence indicated at the top of each plot. The position of the 9-nt sequence in the DR1 motif is indicated; pl–p9 is on the 5' side and p5–pl3 is on the 3' side. The 'TC' in the HNF4-specific half site sequence is in red. Bound probes of HNF4 α (H), COUPTF2 (C), and RXR α (R) have scores below 1371, 1530 and 1285, respectively.



Figure 3. In vivo DNA binding specificity of HNF4 α and RXR α . (A) Analysis of HNF4 α and RXR α ChIP-seq data from mouse liver and pre-adipocyte cells, respectively. Shown are consensus binding motifs in overlapping and non-overlapping ChIP-seq peaks. The dimeric status of RXR α (hetero- versus homodimer) in the ChIP peaks is unknown although only a small portion of peaks overlap with PPAR γ peaks (49). Numbers refer to number of ChIP-seq peaks analyzed for each category. (See Supplementary Table S1 for references for the data sets.) (B) Fold enrichment of 16 half site sequences in ChIP-seq peaks of HNF4 α and RXR α described in (A). Variations in the half site sequences are underlined. Average enrichment scores of all data sets for the same protein are used and SDs are indicated by

were two exceptions (AGGTCAAAGTCCA and GGG TCAAAGTCCA) that bound RXR α and COUPTF2 weakly; they are hybrids of the canonical DR1 and the HNF4-specific half site. In contrast, the probes containing AGTCCA only in the 5' half site were not completely specific for HNF4 α (Figure 2D, middle panel) while those with the canonical AGGTCA in the 3' half site preferred COUPTF2 and RXR α (Figure 2D, right panel). Finally, a similar analysis of 16 H4-SBM-like motifs with all possible permutations at p10 and p11 confirmed that the H4-SBM xxxCAAAG TCCA is the only motif recognized by HNF4 α specifically (Supplementary Figure S3).

H4-SBM is bound by HNF4a but not RXRa in vivo

Since the DNA in the PBM does not necessarily have exactly the same conformation as DNA wrapped in nucleosomes, we determined whether the H4-SBM bound by HNF4 α in vitro is also bound in vivo by analyzing 26 publicly available genome-wide occupancy profiles for 14 NRs (Supplementary Table S1). Since data from the same tissue/cell type were not available for all the NRs and since we are concerned with binding motifs and not specific target genes, we combined and compared ChIP-seq data for a given NR across different conditions/cell types for greater statistical power. First, we compared HNF4 α ChIP'ed from mouse liver (48) and RXRa ChIP'ed from mouse 3T3-L1 (pre-adipocyte) cells (49). In 12097 peaks unique to HNF4a, the motif mined by Gibbs Motif Sampler (39) was found to be very similar to the H4-SBM, whereas the motif in 4285 peaks unique to RXRa was more similar to the canonical DR1 (Figure 3A). Interestingly, there were 736 ChIP peaks for HNF4 α in liver and RXR α in pre-adipocytes that overlapped; the motif mined from those peaks had minor variations from both the H4-SBM and the canonical DR1 but was nearly identical for RXR α and HNF4 α .

We next compared the enrichment of the 16 different half site sequences with all possible permutations at p10 and p11 in ChIP-seq peaks of RXR α and HNF4 α . The H4-specific half site sequence AGTCCA had a 2-fold higher enrichment level compared to control regions (P = 0.001) in the HNF4 α peaks but not the RXR α

Figure 3. Continued

error bars. Empirical P values for enrichment levels (*P < 0.05, **P < 0.001, ***P < 0.0001) are estimated by comparison with the enrichment levels of 1000 6-nt random genomic sequences. (C) Enrichment of particular sequences near the center of the HNF4 α and RXRa ChIP-seq peaks described in (A). Distribution of the indicated DNA sequences containing AGGTCA (left) or AGTCCA (right) are shown in density plots. The x-axis indicates the distance (in base pairs) from the peak center. Density plots in black lines are derived from five 6-nt random genomic sequences as controls. (D) Binding polarities of HNF4a and RXRa in vivo. Fold enrichment of 9-nt motifs in the HNF4a and RXRa ChIP peaks described in (A) are shown in bar plots. Each 9-nt motif contains a 6-nt canonical DR1 half site or HNF4-specific half site sequence (underlined) and a 3-nt extension (p7-p9) that defines the polarity of the half site [half site sequence on the 3' side (blue) or the 5' side (red)]. Error bars indicate SD among multiple data sets.

peaks (Figure 3B). In contrast, the canonical AGGTCA half site showed high and comparable enrichment in both the HNF4 α and RXR α peaks. These results suggest that the H4-specific half site is preferred by HNF4 α but not RXR α *in vivo*.

Since ChIP-seq peaks typically cover a region much larger than the size of a TF binding site, it is frequently assumed that the binding site of a protein is close to the peak center (peak summit). Therefore, we compared the distance from the peak center to the canonical and HNF4-specific half site in the ChIP peaks. In both the HNF4 α and RXR α peaks, the canonical half site AGGT CA was significantly enriched around the peak center compared to random sequences (P < 0.001); adding 1 nt at a time to the 5' side of the motif significantly increased the enrichment (Figure 3C, left panels). In contrast, when the HNF4-specific half site was analyzed, the enrichment was observed in HNF4 α peaks but not the RXR α peaks (right panels), confirming that the H4-SBM (CAAAG TCCA) is bound *in vivo* by HNF4 α but not RXR α .

Finally, in order to determine whether there was any evidence of half site polarity *in vivo*, the enrichment levels of 9-mer sequences with the canonical (AGGTCA) and the HNF4-specific (AGTCCA) half site on both the 5' and 3' side were compared in the HNF4 α and RXR α ChIP-seq peaks. HNF4 α had a clear preference for the 3' side for both half site sequences (Figure 3D, left panel, blue versus red bars). While a similar result was observed for the canonical half site in the RXR α peaks, the HNF4-specific motif was not enriched in either position for RXR α (right panel). These results further confirm that AGTCCA is not bound by RXR α *in vivo* and that the polarity for the half site preference observed in the PBM for RXR α and HNF4 α is also observed *in vivo*.

H4-SBM is exclusive to HNF4a in vivo

To determine whether the H4-SBM is recognized by any other NR aside from HNF4 α , we analyzed ChIP-seq data from 12 additional NRs that are known to be expressed in one or more of the same tissues as HNF4 α -PPAR γ (NR1C3) (46,49,50), PPAR8 (NR1C2) (51), VDR (NR1I1) (52), FXRa (NR1H4) (53,54), PXR (NR1I2) (55), ERRβ (NR3B2) (56), RARα (NR1B1) (57), ERα (NR3A1) (58), LXR^β (NR1H2) (59), LRH-1 (NR5A2) (60), Rev-Erba (NR1D1) (61). The glucocorticoid receptor (GR, NR3C1) (46,62,63), which prefers a different half site AGAACA (64,65), served as a negative control (Supplementary Figure S4A). Motifs of consensus sequences derived from the 12 NR ChIP-seq peaks did not reveal any binding to the H4-SBM (Supplementary Figure S4B). When 6-mer half site sequences were examined, the H4-specific half site (AGTCCA) was found to be enriched in PXR peaks but not in peaks for the remaining 11 NRs (Figure 4A and Supplementary Figure S5). In contrast, there was an enrichment of the canonical half site AGG TCA in 8 of the 12 NRs (Supplementary Figure S5), which served as a positive control. We also examined the half site sequences as a function of distance from the peak center. There was an enrichment in AGGTCA at the peak center for all tested NRs except GR. The HNF4-specific half site, however, was only enriched in peak centers for HNF4 α and three other NRs: Rev-Erb α , FXR α and PXR (Supplementary Figure S6).

Since searching with the 6-mer (AGTCCA) does not provide any information about its position within a 13-nt motif (i.e. 5' or 3' side), the search was repeated with the 9-mer H4-SBM (CAAAGTCCA). We found a 13.5-fold enrichment in HNF4a peaks and a considerable enrichment (>3-fold) in FXRa, PXR and Rev-Erba peaks: the remaining NRs continued to show no enrichment (Figure 4B). Furthermore, the ChIP-seq peaks for FXRa, PXR and Rev-Erba ChIP-seq showed considerable overlap with HNF4 α peaks (Figure 4C and Supplementary Table S5). Since ChIP cannot distinguish between direct and indirect binding, it was possible that FXRa, PXR and Rev-Erba were binding to the H4-SBM through HNF4 α . To address this, we compared the enrichment level of the HNF4-specific half site (AGTCCA) and the full H4-SBM (CAAAGTCCA) in the peaks of each of the three NRs that overlapped with the HNF4 α peaks to those that did not overlap. If a particular sequence is a direct binding target of a NR, then similar enrichment levels are expected in both overlapping (sector C) and non-overlapping peaks (sector A) (Figure 4D, left). On the other hand, if the NR requires HNF4 α to bind the sequence, then the overlapping peaks should have a higher enrichment level than the non-overlapping peak (Figure 4D, right). The results show that both AGTCCA and CAAAGTCCA are enriched in the overlapping peaks but not the non-overlapping peaks for all three NRs (Figure 4E). The fold enrichment for H4-SBM was in fact 3- to 6-fold higher in the overlapping peaks compared to the non-overlapping peaks. Furthermore, the HNF4-specific half site was the only 1 of 16 candidate 6-mers that showed such an enrichment for these NRs (Supplementary Figure S7C). A similar analysis with the peaks unique to HNF4a (sector B) verified the methodology (Supplementary Figures S7B and S7D). These results suggest that while PXR, FXRa and Rev-Erba are localized to regions in the genome that contain the HNF4-specific motif, they may do so only when HNF4 α is also present in the same region. While we cannot definitively rule out direct interactions between these NRs and HNF4α, AGGTCA-like motifs were found in about half of the overlapping peaks that contain the H4-SBM (Supplementary Figure S7E-S7H). This suggests that PXR, FXRa and Rev-Erba may bind canonical AGGT CA-containing motifs cooperatively with HNF4 α in regions that contain the H4-SBM. For example, on the Cvp7al promoter there are overlapping ChIP-seq peaks for HNF4a, PXR and Rev-Erba but the only H4-SBM is at the center of the HNF4 α peak (Supplementary Figure S7I).

HNF4-specific DNA recognition is mediated by two residues in the DBD, Asp69 and Arg76

To determine the molecular basis of HNF4-specific DNA recognition, protein–DNA interactions were analyzed in the DBD structures of HNF4 α and RXR α (66,67).



Figure 4. Comparison of *in vivo* DNA binding specificity of multiple NRs. (A) Fold enrichment of the canonical DR1 half site sequence (blue) and the HNF4-specific half site sequence (red) in ChIP peaks of 14 NRs. ChIP-seq data for different isoforms of a NR or multiple data sets for the same protein are combined (see Supplementary Table S1 and text for references); SD is indicated by error bars. Empirical *P* values for enrichment levels (**P* < 0.001, *** *P* < 0.0001) are estimated using enrichment levels compared to 1000 6-nt random genomic sequences. Only *P* values for AG<u>TC</u>CA are indicated. (See Supplementary Figure S5 for enrichment values for 14 additional half site sequences.) (B) Fold enrichment in ChIP-seq peaks of 14 NRs as in (A) but for the H4-SBM. *P* values are estimated using 1000 9-nt random genomic sequences. (C) Overlap between ChIP-seq peaks among select NRs in mouse liver. Given is the percent of overlapping ChIP-seq peaks between the NRs indicated on the top and the side. For example, 59.6% of FXR α peaks overlap with HNF4 α peaks, while only 32.2% of HNF4 α peaks overlap with FXR α peaks. (D) Schematic representation of direct and indirect binding of a NR to the H4-SBM. Left: direct binding to the H4-SBM is characterized by the even distribution of H4-SBM sites in overlapping and non-overlapping peaks (sectors C and A, respectively). Right: indirect binding of a NR to the H4-SBM via HNF4 α would yield a greater number of H4-SBM sites in the overlapping peaks. (E) Enrichment of the HNF4-specific half site (left) and the HA-SBM (right) in ChIP-seq peaks for the indicated NRs as described in (C). Blue bars indicate fold enrichment in non-overlapping peaks (sector C). *P* values in (A)] are estimated using ratios of 1000 6- and 9-nt random genomic sequences, respectively, for two analyzed motifs.

Among all the residues making contacts with the DNA, only four residues differ between RXR α and HNF4 α . In HNF4 α , two of these residues, Asp69 and Arg76, lie in the first DNA recognition helix of the DBD and interact with side chains of nucleotides at p3 and p4 in both half sites of the DR1 motif (Figure 5A). These two residues are completely conserved in HNF4 genes across all species from human down to Trichoplax, except for *Caenorhabditis elegans* which contains ~260 HNF4-like genes (Supplementary Figure S8A). Residues at equivalent positions in RXR α and COUPTF2 are Glu and Lys, respectively (Figure 5B).

To examine the effect of Asp69 and Arg76 on HNF4 α binding specificity, single and double point mutations D69E, R76K and D69E/R76K were introduced into the HNF4 α DBD to convert it into an RXR/COUPTF2-like DBD. In the PBM, the D69E mutant selectively abolished binding to the H4-SBM while R76K seemed to decrease binding in a non-selective fashion; interestingly, the double mutant D69E/R76K yielded a profile nearly identical to that of D69E (Supplementary Figure S8B), suggesting that both Asp69 and Arg76 are necessary for optimal discrimination between H4-SBM and other sites. Importantly, the HNF4 α D69E/R76K double mutant also altered the binding profile to more closely resemble that of

RXR α /COUPTF2; none of the HNF4-specific binders were bound by the double mutant (Figure 5C and D). While some common binders of HNF4 α and RXR α / COUPTF2 were also affected by the mutations (Supplementary Figure S8C and S8D), most of them bear the AGTTCA motif in the 3' half site that is also preferred by HNF4 α but not RXR α *in vivo* (Figure 3B). Finally, HNF4 α and HNF4 γ are the only human NRs with an aspartic acid at residue 69 and an arginine at residue 76 [Figure 5B and (7)], suggesting that the H4-SBM may be truly specific to HNF4 in the entire NR superfamily.

HNF4α activates gene expression using HNF4-specific binding sites

To determine whether the HNF4 α binding site preference results in a functional outcome (i.e. gene expression), luciferase reporter assays were performed using a known HNF4 α /RXR α response element from the human *APOA1* promoter. As predicted, HNF4 α activated gene expression using both the wild-type response element (WT-RE) AGGGCAgGGGTCA and an HNF4-specific mutant (MUT-RE) AGTCCAgGGTCCA; in contrast RXR α activated expression only from the WT-RE (Figure 6A). In addition, COUPTF2, a repressor, competed with



Figure 5. Molecular basis of HNF4-specific DNA recognition. (A) Superimposed crystal structures of HNF4 α DBD (3CBB, cyan) and RXR α DBD (1BY4, yellow). In the HNF4 α structure, only D69 and R76 interact with the third and fourth base pairs of the AGTCCA half site. The equivalent residues of RXR α are E and K, respectively. (B) Sequence alignment of the indicated human NRs in the DNA recognition helix of the first zinc finger. Numbers 69 and 76 refer to the residues in human HNF4 α 2. (C) Scatter plots of PBM data as in Figure 2A of the WT and mutant HNF4 α compared to WT RXR α . Binding sequences specific to HNF4 α WT are indicated as red spots. All the red spots shift to the non-binding quadrant in the HNF4 α double mutant D69E/R76K. (D) As in (C) but comparison of HNF4 α WT and D69E/R76K mutant to COUPTF2.

HNF4 α on the WT-RE but not the MUT-RE (Figure 6B). These results verify that HNF4 α is capable of activating gene expression using the H4-SBM, while RXR α and COUPTF2 cannot functionally compete with HNF4 α for the H4-SBM response element.

The effect of the D69E mutation was also examined in the luciferase assay using an endogenous promoter from a known HNF4 α target gene, *APOC2* (31). WT HNF4 α successfully activated the expression of the WT promoter containing an AGGCCAaAG<u>TCCT</u> motif whereas the D69E mutant failed to do so (Figure 6C).



Figure 6. Specificity of the HNF4-specific binding motif in cell-based reporter gene assays. (A) Luciferase reporter assay in HEK293T cells 500 ng of indicated reporter construct co-transfected with (pGL4.23.hApoA1.DR1.WT or pGL4.23.hApoA1.DR1.MUT) plus (+, 50 ng of expression vector pcDNA3.1.hHNF4α2 or pMT2.hRXRa) or empty vector (-, pcDNA3.1). 9-cis-retinoic acid (1µM) (or ethanol as vehicle control) was added as indicated. Right: diagram of the wild-type (WT) and mutated (MUT) response elements in the reporter construct for (A) and (B). Shown are relative light units (RLU) normalized to β-galactosidase activity. Bars are average of three replicates. Error bars indicate SEM. (B) As in (A) but with the expression vector (+, pcDNA3.1.hHNF4 α) and two different amounts of pMT2.hCOUPTF2 (400 and 800 ng). Empty vector was added to adjust for amount of total DNA transfected. (C) As in (A) but with the indicated reporter constructs (pGL4.10.hAPOC2. WT or pGL4.10.hAPOC2.MUT) plus the expression vector (+, pcDNA3.1hHNF4a or pcDNA3.1hHNF4a.D69E carrying the D69E mutation in its DBD). Right: diagram of the WT and MUT hHNF4a response element in the human APOC2 promoter.

In contrast, both the WT and D69E mutant activated gene expression when the H4-specific response element was mutated to a more canonical DR1 element AGGCC AaAG<u>GT</u>CT (Figure 6C, reverse complement sequences



Figure 7. Prediction of HNF4-specific binding sequences and target genes. (A) Distribution of HNF4-specific binding sequences in HNF4 α ChIP-seq peaks in HepG2 cells (41) are shown in density plots. Red line, ~200 HNF4-specific PBM sites; blue line, ~3000 predicted HNF4-specific sites; green line, the HNF4-specific half site sequence; black line, five 6-nt random genomic sequences used as controls. *P* value is calculated by the *F*-test. (**B**) Box plots of predicted and verified HNF4-specific sites compared to ~3000 random sequences in the regulatory regions of human genes (-10 to +10 kb from TSS). Both the predicted and verified HNF4-specific stepsecific sequences occur at a statistically significant high frequency compared to random sequences (*P* < 0.001 in Student's *t*-test). (**C**) Strategy used to search for potential HNF4-specific target genes. See text for details. (**D**) Classification of ~100 predicted human HNF4 α target genes based on the analysis in (C). All of these genes have at least one HNF4-specific site in an HNF4 α ChIP-seq peak in HepG2 cells (41) and were down regulated by HNF4 α RNAi in HepG2 cells (13). See Supplementary Table S4 for a complete list of the 137 genes identified in (C) as well as Gene Ontology results. *, genes previously identified in (13) as predicted HNF4 α target.

are shown in the figure to maintain the orientation in the genome). This result confirms that Asp69 is responsible for the HNF4-specific DNA binding.

Functionality of HNF4-specific binding sequences in vivo

In order to determine whether the H4-SBM is functional under physiological conditions, we trained an SVM model on \sim 200 verified H4-SBM sites from the PBM to predict all 13-nt sequences that can be bound specifically by HNF4 α (variations in the less conserved positions of the consensus H4-SBM, CAAAGTCCA, such as p6-p8 and p13 can be tolerated, Figure 1B). This method generated ~3000 high confidence, predicted HNF4-specific binding sequences. (See Supplementary Table S3 for a list of all H4-SBM sequences and http://nrmotif.ucr.edu/ NRBSScan/H4SBM.htm for a web-based, HNF4-specific binding sequence search tool.)

The predicted H4-SBM sites were first validated by comparing their position in ChIP-seq peaks to the verified H4-SBM sites and the AGTCCA half site (Figure 7A). The predicted and verified H4-SBM sites

were then examined across all protein-coding genes in the human genome relative to the TSS (+1), -10 to +10 kb. (HNF4 binding sites are symmetrically distributed on both side of TSS, Supplementary Figure S9). The results show comparable occurrence frequencies for the predicted and verified sites (P > 0.05) that were significantly higher than those of random sequences (P < 0.001) (Figure 7B). Finally, we used the predicted H4-SBM sites to search HNF4 α ChIP-seq and expression profiling data to identify potential HNF4a target genes. A total of 730 genes in the human liver cell line HepG2 were found to have at least one predicted H4-SBM site in a HNF4a ChIP-seq peak within -10 to +10 kb of +1 (Figure 7C). Among these genes, 137 were down-regulated by HNF4 α RNA interference (RNAi) in HepG2 cells (13), suggesting that these genes are direct targets of HNF4a. GO analysis showed that the H4-specific genes were enriched in a variety of metabolic processes (e.g. lipid, carbohydrate, xenobiotic/drug metabolic processes; homeostasis; transport), typical of classical HNF4a targets (Supplementary Table S3). There were also genes in some of the new categories of HNF4 α targets that we identified in our previous study (13), such as immune response, signal transduction, apoptosis and cell structure (Figure 7D). That study, however, did not identify HNF4-specific motifs or targets. All told, this analysis identified ~ 100 new predicted, direct HNF4a target genes that have an H4-SBM in a ChIP peak and are down regulated by an HNF4α RNAi.

Interestingly, several genes associated with acyl Coenzyme A metabolism (acyl-CoA synthases, thioesterases, ligases and a co-factor for a desaturase) were identified as putative HNF4α targets with an H4-SBM site (ACSM2B, EHHADH, ACOT2, ACSF2, SLC27A2, CYB5A, AGXT). Others have shown that HNF4 α binds not only acyl Co-A binding protein but also fatty acyl thioesters of Co-enzyme A that appear to act as modulators of HNF4a function (68-70). Acyl Co-A is well known for donating an acetate group in acetylation reactions of lipids and proteins. HNF4 α itself has been shown to be acetylated in its hinge region (71), as have many of the enzymes encoded by HNF4a target genes-e.g. PEPCK (PCK1), a classical HNF4 α target (72) that catalyzes the rate limiting step in gluconeogenesis, and the new H4-SBM target reported here, EHHADH, which encodes enoyl-coenzyme A hydratase/3-hydroxyacylcoenzyme A that catalyzes two steps in fatty acid oxidation (73). While it is not only possible but very likely that these H4-SBM genes are also regulated by other TFs, including other NRs via other sites in the promoters, these results nonetheless reinforce the notion that there is an important relationship between HNF4 α and processes involving Co-enzyme A. Finally, we note that several of the newly identified H4-SBM containing HNF4 α target genes are linked to human diseases (LIPA, EHHADH, AGXT, PIPOX, HGD, CYB5A, PDZK1, F11), expanding the clinical relevance of HNF4a. (See Supplementary Table S3 for a complete list of H4-SBM target genes from both a -10 to +10 kb and a -2 to +1 kb analysis.)



Figure 8. New insights into NR DNA binding from PBM and ChIP-seq analysis. (A) Half site motif for the indicated NRs along with the amino acid sequence of the P box region in the DNA recognition helix of the first zinc finger in the DBD showing the residues relevant to this work. GR, MR, AR and PR: glucocorticoid, mineralocorticoid, androgen and progesterone receptors, respectively). The only NRs that have P boxes that are not depicted are TLX (NR2E1) with an Asp (D) and Lys (K) and PNR (NR2E3) with an Asn (N) and Lys (K) at positions 69 and 76, respectively (HNF4a numbering). (B) RXRa, COUPTF2 and HNF4a homodimers all bind classical DR1-like motifs (AGGTCAxAGGTCA) while HNF4a homodimers also bind an HNF4-specific binding motif (H4-SBM, xxxxCAAAGTCCA) that is not bound by RXRa or COUPTF2 in the PBMs, nor 11 other NRs in ChIP-seq. (C) RXRa has a preference for the 3' half site in a DR1-like motif in RXR:RXR homodimers as well as in RXR:RAR and RXR:PPAR homodimers, suggesting that an exchange between the 5' RXRa monomer and RAR or PPAR monomers could occur without disrupting the binding of the 3' $RXR\alpha$ monomer. COUPTF2 also prefers the 3' half site of the DR1-like motifs, suggesting it could compete with RXRa homo- or heterodimers on these response elements. (D) PXR, FXRa and RevErba ChIP peaks from mouse liver were all found to be enriched for the H4-SBM but only in the presence of HNF4 α , suggesting that they may occupy their own binding sites near HNF4_α-bound H4-SBM.

DISCUSSION

The dogma in the NR field is that NRs recognize DNA targets based on one of two motif modules, AGGTCA (non-steroid receptors plus ER) and AGAACA (all other steroid receptors) (Figure 8A). While there are additional rules for spacing and orientation [as well as some variations on the half site motif, such as AGTTCA (7)] that distinguish the different receptors, those rules do not adequately explain the diversity of NR function *in vivo*. We addressed this issue by comparing the binding specificity of three highly related NRs—HNF4 α , RXR α and COUPTF2—on ~3000 different variations of a canonical

DR1 (AGGTCAxAGGTCA) using PBMs and full length receptors in crude nuclear extracts. We identify a new motif module for HNF4 α , AG<u>TC</u>CA and provide additional insight into NR DNA binding (Figure 8A–D).

NR DNA binding polarity

In contrast to most gel shift results, including our own (27), we observed excellent binding of RXR α in the PBM in the absence of an ectopically expressed partner. The RXR α binding was nearly identical to that of COUPTF2, and both receptors exhibited a polarity in binding with a preference for the 3' half site of the DR1 (Figure 8B and C). To our knowledge, this is the first report to examine polarity in DNA binding using PBMs, and the first report of polarity for NR homodimers. Since RXR α homodimers can activate transcription in the presence of the 9-*cis* retinoic acid and since COUPTF2 generally represses transcription, this suggests that the competition already observed between these two NRs on a few genes may in fact be a much broader phenomenon.

While binding polarities of NR homodimers have not been reported previously, the concept in heterodimers is not new. For example, RXR binds the 5' half site in DR3. DR4 and DR5 motifs as heterodimers with VDR, TR and RAR, respectively (74–76). This polarity was also noted in crystal and solution structures of RXR heterodimers with RAR and VDR (77,78). However, on DR1 motifs the polarity of the RXR heterodimers is reversed with RXR in heterodimers with RAR and PPAR occupying the 3' half site in DR1 motifs (78-82). Since our results now show that the RXR α homodimer also prefers the 3' half site in DR1-like motifs, this suggests that there could be an exchange between the 5' RXR α monomer in the homodimer and PPAR or RAR monomers (Figure 8C). The net result would be a replacement of the RXR α homodimer with a heterodimeric complex, perhaps while the 3' RXR α monomer remains bound to the DNA. This would suggest a new paradigm for RXRa dimer exchange and a potential new role for RXRa homodimers as placeholders for RXR α heterodimers on DR1-like motifs, but must be experimentally proven. A thorough analysis of RXR heterodimer DNA binding specificity also needs to be done.

An NR-specific DNA binding motif

Comparison of the HNF4 α PBM results to that of RXR α and COUPTF2 allowed us to identify *in vitro* an HNF4-specific binding motif (xxxxCAAAG<u>TC</u>CA) that we had not identified previously when we analyzed HNF4 α alone (13). *In vivo* analysis showed that ~8 and 42% of HNF4 α ChIP-seq peaks contain CAAAG<u>TC</u>CA and AG<u>TC</u>CA, respectively (motif variations not considered). Comparable results were observed for CAA AGGTCA and AGGTCA (Supplementary Figure S10), suggesting a similar importance of AGGTCA- and AG<u>TC</u>CA-based motifs for HNF4 α binding. The H4-SBM was not bound by RXR α or COUPTF2 in the PBM, nor eight other NRs in ChIP-seq data (Figure 8B). However, three NRs (FXR α , PXR and Rev-Erb α) were associated with the H4-SBM in vivo, most likely via their own binding sites, not the H4-SBM (Figure 8D). While there are a few reports of these NRs regulating the same target genes as HNF4 α via their own binding sites (83-87), our results suggest that this may be a much broader phenomenon. The considerable overlap between PXR, FXRa, Rev-Erba and HNF4a peaks in ChIP-seq data also demonstrates the complexity of NR-mediated regulation and the difficulty of precisely identifying binding sites for a given TF in ChIP-seq peaks. This complexity is increased by protein-protein interactions between HNF4 α and other NRs such as PXR and FXR and competition for co-regulators (12). Our analysis shows how this problem can be overcome at least in part with a better understanding of binding specificities generated by the PBMs.

While this is the first identification and genome-wide analysis of an HNF4-specific binding motif, there were previous reports using classical methods suggesting that such a motif might exist. A survey of the literature identified 28 response elements that had been examined for responsiveness to HNF4 α , RXR α and/or COUPTF2. Six of those elements carry the H4-SBM TC instead of the DR1-like GT at p10 and p11, and all six were responsive to HNF4 α but not RXR α or COUPTF2. In contrast, the remaining 22 response elements did not have a TC at p10-11 and were responsive to RXR α or COUPTF2 (Supplementary Table S2). However, since these findings were generated by different groups over a period of years, an HNF4-specific motif was never identified.

In addition to the H4-SBM, we also identified a pair of residues in the HNF4 DBD that is responsible for the H4-SBM binding (Asp69 and Arg76) (Figure 8A). Asp69, which has the greatest effect on HNF4-specific binding, is in the P box which was shown previously to be responsible for the different half site of GR and related receptors (AGAACA) (5,88). Arg 76 is in the helix that contacts the DNA but to our knowledge has not been previously associated with DNA binding specificity. HNF4 is the only human NR with the combination of Asp69 and Arg76 and all HNF4 DBDs, except some of the HNF4-like genes in C. elegans, have these residues, suggesting that the H4-SBM may be truly unique to HNF4. There is only one other NR that has an Asp at position 69, TLX (NR2E1), but it also has a Lys at position 76 instead of an Arg (7). Since the R76K mutant of HNF4 lost most of its DNA binding activity, we assume that TLX would not be able to recapitulate the HNF4-specific binding, although that remains to be determined.

Finally, it is of interest to note that HNF4, RXR and COUPTF are among the most ancient of all the NRs (9). Since the biologically least complex metazoans currently in existence all have at least two NR genes (one HNF4-like and one RXR/COUPTF-like), it has been proposed that the NR family evolved from a now extinct early metazoan that contained a single NR gene, which was most similar to HNF4 (89). Intriguingly, the HNF4 DBDs of primitive metazoans all have the Asp69-Arg76 pair of mammalian HNF4. In conclusion, our results highlight a complexity of NR DNA binding specificity that was previously under appreciated. They also demonstrate the usefulness of the PBM approach to more accurately define that complexity and thereby more precisely identify NR target genes *in vivo*. It will be of interest to compare the binding specificity of other NRs in a similar fashion.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–5, Supplementary Figures 1–10, Supplementary Materials and Methods, and Supplementary References [16,27,30,32,39,41,48,53,54, 66,67,90–91].

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