Global analysis of *in vivo* Foxa2-binding sites in mouse adult liver using massively parallel sequencing

Elizabeth D. Wederell¹, Mikhail Bilenky², Rebecca Cullum¹, Nina Thiessen², Melis Dagpinar², Allen Delaney², Richard Varhol², YongJun Zhao², Thomas Zeng², Bridget Bernier², Matthew Ingham², Martin Hirst², Gordon Robertson², Marco A. Marra², Steven Jones² and Pamela A. Hoodless^{1,*}

¹Terry Fox Laboratory and ²Michael Smith Genome Sciences Centre, BC Cancer Agency, Vancouver, Canada

Received April 14, 2008; Revised May 13, 2008; Accepted May 30, 2008

ABSTRACT

Foxa2 (HNF3ß) is a one of three, closely related transcription factors that are critical to the development and function of the mouse liver. We have used chromatin immunoprecipitation and massively parallel Illumina 1G sequencing (ChIP-Seq) to create a genome-wide profile of in vivo Foxa2-binding sites in the adult liver. More than 65% of the ~11.5 k genomic sites associated with Foxa2 binding, mapped to extended gene regions of annotated genes, while more than 30% of intragenic sites were located within first introns. 20.5% of all sites were further than 50 kb from any annotated gene, suggesting an association with novel gene regions. QPCR analysis demonstrated a strong positive correlation between peak height and fold enrichment for Foxa2-binding sites. We measured the relationship between Foxa2 and liver gene expression by overlapping Foxa2binding sites with a SAGE transcriptome profile, and found that 43.5% of genes expressed in the liver were also associated with Foxa2 binding. We also identified potential Foxa2-interacting transcription factors whose motifs were enriched near Foxa2-binding sites. Our comprehensive results for in vivo Foxa2-binding sites in the mouse liver will contribute to resolving transcriptional regulatory networks that are important for adult liver function.

INTRODUCTION

Forkhead box A2 (Foxa2, HNF3 β) is one of three closely related transcription factors, whose expression is an initiating factor in the earliest stages of liver specification during embryonic development and normal liver

homeostasis in the adult liver (1-3). Foxa1 (HNF3 α), Foxa2 and Foxa3 (HNF3γ) were initially identified in liver cells by their ability to interact with the promoters of two important liver-expressed genes-transthyretin (Ttr) and alpha-1-antitrypsin (A1AT) (4). The three members of this group contain a highly conserved winged-helix DNA-binding domain and are critically important in mouse liver development and maintenance. While mutational analysis has shown that the three FoxAs are functionally redundant in the mouse liver, with all three able to bind to the same sequence (5), Foxa1 or Foxa2 are most critical for the specification of the liver (6). Foxa2 is the first of the three family members to be expressed in the embryo prior to gastrulation, with Foxal and Foxa3 expression following at the onset of gastrulation (1). All three are expressed in the embryonic endoderm cells that constitute the precursor cells for all gut organs, and all three remain present and active in the adult liver. In the adult, Foxa2 plays a major role in glucose and lipid metabolism, targeting genes such as glucose-6-phosphatase, catalytic (G6pc) and tyrosine aminotransferase (Tat) (3,7). In addition to its role as an important transcription factor regulating gene expression in liver development and function, Foxa2 can also open compacted chromatin, allowing for the activation of transcription from silenced genes, and so can act as a pioneer transcription factor in liver signaling cascades (8).

In the adult, the liver is involved in the maintenance of general homeostasis. It is comprised primarily of hepatocytes and has many varied roles including regulation of metabolic activity, cholesterol production and secretion, production of plasma proteins, chemical detoxification, drug and alcohol elimination and, during embryonic development, site of red blood cell production. The cellular functions of the normal liver are governed through the dynamic interaction of a central group of transcription factors that regulate liver-specific gene expression. This group includes

^{*}To whom correspondence should be addressed. Tel: +604 675 8133; Fax: +604 877 0712; Email: hoodless@bccrc.ca

^{© 2008} The Author(s)

HNF4, Foxa1, Foxa2, Foxa3, Tcf1/2 (HNF1α/β), Onecut1 (HNF6) and C/EBPa. These transcription factors interact in highly complex ways that are continually regulated throughout prenatal and postnatal development (9).

Until recently, the transcriptional regulatory functions of FoxA proteins were studied on a gene-by-gene basis. However, the advent of chromatin immunoprecipitation coupled with microarray technology (ChIP-chip) has allowed the identification of in vivo binding locations on a larger scale. Several groups have used ChIP-chip to investigate the in vivo binding sites of important liver transcription factors, including Foxa2 (9-17). Most microarray studies have targeted promoter regions that are closely associated with transcriptional start sites (TSS) of genes; however, it has been increasingly recognized that transcription factors have the ability to regulate gene activity from greater distances. A study in the hepatoma cell line HepG2, using ChIP combined with a microarray that covered the ENCODE regions representing 1% of the human genome, demonstrated that in vivo transcription factor-binding sites (TFBS) often occur outside of traditional promoter regions (14). Furthermore, recent genome-wide ChIP-sequencing experiments have highlighted the benefits of binding site analyses of transcription factors that are not restricted to specific genomic regions or by the need for annotated genes (18–21). Newer approaches to massively parallel sequencing (22-24) have the ability to more accurately detect transcription factor binding in any genomic region at much lower cost than earlier global sequencing methods. Since Foxa2 acts not only as an important transcriptional regulator in liver development and maintenance but also as competence factor in accessibility of DNA by other transcription factors, understanding its genomic wide binding locations is imperative to understanding of gene regulation in the liver.

This study presents the first genome-wide investigation of in vivo Foxa2 binding target regions in the adult mouse liver. We combined the power of a secondgeneration DNA sequencer with ChIP and quantitative PCR (qPCR) validation to show not only high coverage of well-characterized Foxa2 target sites and genes, but also novel aspects of Foxa2 interaction with the mouse genome. We show that the majority of Foxa2-binding sites are within 10kb upstream of TSS and within the first introns of genes. In addition, we provide evidence that Foxa2 binds a large number of targets in unannotated genomic regions. We also show that Foxa2's apparent influence on gene expression remains extensive in the adult liver, with close to half of abundantly liver-expressed genes being associated with at least one Foxa2-binding site. Finally, we have demonstrated the presence of additional transcription factor motifs within Foxa2-binding site sequences that represent potential Foxa2-interacting partners. We present analyses of the expression of each transcription factor in the liver and function of the group of genes with which each motif is associated. We have identified a large number of novel Foxa2 in vivo binding sites, which may be involved in specific gene regulation or permissive gene regulation through the opening of chromatin. We have also confirmed

Foxa2 binding to many novel sites using ChIP-qPCR indicating the high quality of our dataset. By identifying Foxa2 binding locations in combination with gene expression and bioinformatics analysis of potential cointeracting transcription factors, we provide novel insights into Foxa2 transcriptional networks in the adult mouse liver.

METHODS

ChIP

Adult female C57Bl/6J mouse livers were homogenized in 1% formaldehyde and incubated at room temperature (RT) for 10 min, prior to the addition of glycine to 0.125 M followed by incubation at RT for 5 min. Cells were pelleted, washed and resuspended in 5 volumes ChIP cell lysis buffer (10 mM Tris-Cl, pH 8.0, 10 mM NaCl, 3 mM MgCl₂, 0.5% NP-40) containing protease inhibitor cocktail tablets (Roche, Laval, QC, Canada). Cells were then rehomogenized, incubated on ice for 5 min and repelleted. The pellet was resuspended in 3 volumes ChIP nuclear lysis buffer (1% SDS, 5 mM EDTA, 50 mM Tris-Cl, pH 8.1) supplemented with protease inhibitor cocktail tablets (Roche). Cells were sonicated on ice-water (Sonicator 3000, Misonix, Farmingdale, NY, USA) for 20 cycles of 30s on, 40 s off and 60 µg of chromatin was precleared with Protein G beads (100 µl, Active Motif), Protein Inhibitor Cocktail (0.5 µl, Active Motif) and ChIP dilution buffer (to 250 µl, 0.01% SDS, 1.1% Triton X-100, 167 mM NaCl, 16.7 mM Tris-Cl, pH 8.1). Beads were precipitated and 3 μg of Foxa2 [HNF-3β (M-20): sc-6554, Santa Cruz] antibody or normal rabbit IgG (sc-2027, Santa Cruz, Santa Cruz, CA, USA) was added to supernatants. Fresh Protein G beads were also blocked with 1 mg/mg BSA and 0.1 mg/mg herring sperm DNA in ChIP dilution buffer. Following overnight incubation, rocking at 4°C, the samples were incubated with the blocked beads for 4h rocking at 4°C. The beads were then precipitated and washed in low salt buffer (0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM Tris-Cl, pH 8.1, 150 mM NaCl), high salt buffer (low salt buffer with 500 mM NaCl), lithium chloride buffer (0.25 M LiCl, 1% NP-40, 1% deoxycholate, 1 mM EDTA, 10 mM Tris-Cl, pH 8.1) and twice with TE buffer. Bead complexes were eluted twice by addition of 125 µl elution buffer (1% SDS, 0.1 M NHCO₃) and rotation for 15 min at RT. NaCl was added to 0.192 M to reverse crosslink and samples were incubated overnight at 65°C. Samples were then incubated with Proteinase K (Invitrogen, Carlsbad, CA, USA) and RNaseA (Sigma-Aldrich, Oakville, ON, Canada) for 2h at 50°C. DNA was purified by two rounds of phenol-chloroform extraction and ethanol precipitation and resuspended in 50 µl dH_2O .

Sequencing

Foxa2-bound DNA (4.7 ng) was purified by SDS-PAGE to obtain 100-300 bp fragments and sequenced on an Illumina 1G sequencer according to Robertson et al. (24). Briefly, size fractionated DNA was extracted and a single adenosine was added using Klenow exo- (3' and 5'

exo minus; Illumina). Illumina adaptors were then added and DNA was subjected to 20 cycles of PCR according to manufacturer's instructions. We then purified DNA and performed cluster generation and 27 cycles of sequencing on the Illumina cluster station and 1G analyzer following the manufacturer's instructions.

Sequencing analysis and peak production

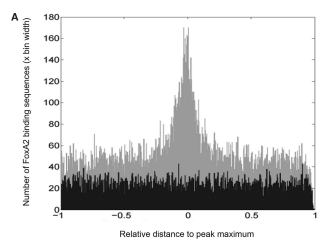
Resulting sequences were mapped to the NCBI Build 36 (mm8) reference mouse genome to produce 13 984 706 mapped reads that were extended to 200 bp length XSETs and overlapped to create peaks according to Robertson et al. (24). To further define the peak dataset, each group of XSETs that represented DNA fragments with identical fragment start coordinates was 'collapsed' to a single XSET. Peaks generated from the resulting filtered reads were thresholded at a peak height of 10, based on the stabilization of the FoxA-like binding sequences enrichment curve (Supplementary Figure 1), creating a high confidence set of Foxa2-binding sites with an estimated false discovery rate of 0.05. The dataset containing all 11 475 peaks can be downloaded from www.bcgsc.ca/data/chipseq.

Foxa2 enrichment analysis

A set of 42 DNA sequences known to bind FoxA proteins in mouse, human or rat was extracted from the literature. The sites, with supporting evidence, are available at the ORegAnno database (www.oreganno.org) (25). Because sequence lengths ranged from 7 to 22 bp, we created a Foxa2 model by adding flanking sequences. We aligned the flanked sequences by scanning them with the Foxa2 position weight matrix (PWM) from the JASPAR database (jaspar.cgb.ki.se) (26). For each flanked sequence, we retained the top scoring sub-sequence that contained at least 5 bp from the binding site originally reported in the literature. The logo for the resulting 42 aligned sequences, 39 of which were unique, is shown in Supplementary Figure 2. To assess the extent to which ChIP-Seq peaks were enriched in Foxa2-like binding sequences, we used a custom Java algorithm to scan peak sequences with a 10-mer representation of each known functional site. We permitted either no mismatches or up to one mismatch at positions that had an information content below 0.6 bits [Supplementary Figures 1 and 7 in Robertson et al. (24)]. We also scanned peak sequences with a PWM constructed from the aligned site sequences, using the above algorithm (Figure 1B). The score assigned to a tested DNA sequence was the ratios of the probabilities of observing a given nucleotide from the PWM model, and from the background model, for all positions in the matrix. We assumed the background probabilities to be 0.25 for all four nucleotides.

Quantitative real-time PCR

Quantitative real-time PCR (qPCR) was performed on an ABI 7900HT Fast Real-Time PCR System using 2x SYBR® Green Master PCR Mix according to the manufacturer's instructions (both from Applied Biosystems, Foster City, CA, USA). The fold enrichment of each target site was calculated as 2 to the power of the cycle



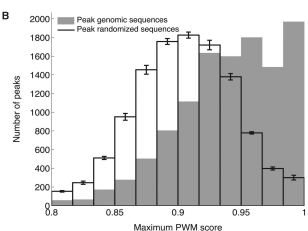


Figure 1. (A) Distances from sites similar to Foxa2-binding sequences to the peak maximum, normalized to peak width. This graph shows that Foxa2-binding sequence sites tended to cluster around peak maxima. (B) Peak enrichment in Foxa2-like sequences, as assessed with PWM scan scores. The Foxa2 PWM used was created from the aligned sequences used to generate the sequence logo shown in Supplementary Figure 2 (see Methods section). The graph indicates that peak genomic sequences contained higher scoring Foxa2-binding sequences than randomized peak sequences; that is, peaks were enriched relative to a random expectation.

threshold (cT) difference between an IgG immunoprecipitated sample and a Foxa2 immunoprecipitated sample. We selected primer sets for six negative regions, all of which showed low enrichment between 0.8 and 1.2. Forty-two other validation sites were chosen over a range of subgroups as defined in the results (41 positive and 1 negative). All primers are listed in Supplementary Table 3.

SAGE

The adult mouse liver long Serial Analysis of Gene Expression (longSAGE) library (SM100) was produced as previously described (27). Briefly, a LongSAGE library was created from 4.7 µg of DNase-treated total RNA using the I-SAGE Long kit and protocol (Invitrogen) and sequenced on capillary DNA sequencers (Applied Biosystems). Tag sequences were mapped to the genome using NCBI Build 36 mouse assembly. SAGE tag sequences for this library are publicly available at www.mouseatlas.org.

SiRNA analysis

SiRNA experiments were performed in the Hepa1-6 hepatocyte cell line. Cells were grown and maintained in Dulbecco's modified Eagle's medium (DMEM) with 10% FBS. Control and mouse Foxa2 siRNA pools were obtained from Santa Cruz and cells were treated with 100 nM siRNA overnight in serum-free DMEM. Media was replaced the following day with DMEM with 10% FBS and cells were cultured for 72 h, prior to total RNA isolation by using Trizol according to the manufacturer's instructions. RNA was reverse transcribed using Superscript III (Invitrogen) to obtain cDNA for use in QPCR reactions. Results presented are an average of three independent experiments.

Transcription factor comotif analysis

The MATCH v2.6 algorithm was used with a profile filter that specified low false positives and good quality models to scan sequences derived from the 3016 peaks that contained at least one exact 10-mer match to one of the 39 known functional sequences, as well as sequences from the 2494 peaks that did not contain such a match (28). A total of 363 vertebrate TRANSFAC Professional v9.3 PWMs were assessed. Scanned sequences extended ±250 bp from each putative Foxa2-binding site under a peak, or for peaks without exact match hits, from the location of the peak maximum. The statistical significance of the number of scan hits per kilobytes for each model was estimated by generating and scanning 100 third-order Markov random sequences from a mouse genomic model outside of open reading frames (rsat.ulb.ac.be/rsat), each of which had a length equal to the combined length of all scanned peak sequences (29). For each TFBS model, we calculated an empirical P-value and z-score from peak versus random scan results, and used the P-value, and then the z-score, to rank the 363 models.

RESULTS

Identification and characterization of Foxa2-binding sites in vivo

To determine locations of in vivo Foxa2 binding in the mouse adult liver, we sequenced Foxa2-bound DNA fragments by combining chromatin immunoprecipitation and massively parallel Illumina 1G sequencing (ChIP-Seq) (22–24). The sequences were then mapped to the mouse genome (NCBI Build 36, UCSC mm8) (see Methods section). Foxa2-bound regions are represented by peaks consisting of overlapping mapped sequences. To determine a high confidence subset of peaks at a minimal peak height, we determined the number of peaks containing a Foxa2-binding sequence. We extracted and aligned a set of 42 sequences (39 of which are unique) that are known from the literature to bind FoxA proteins in mouse, human or rat (see Methods section). This resulted in a 10-mer binding sequence model (Supplementary Figure 1, position 2–11). We scanned all peaks, allowing a maximum of 1 bp mismatch, and assessed the relationship between peak height and the fraction of peaks

Table 1. Global peak properties and peak locations relative to UCSC mm8 known genes

Minimum peak height	10			
# Peaks	11 475			
# Individual DNA reads under peaks	285 176			
Average (median) peak height	16 (13)			
Average (median) peak length (bp)	727.5 (654)			
# Peaks within ±1 kb TSS ^a	694 (6.0%)			
# Peaks in 10 kb upstream TSS region ^a	1800 (15.7%)			
# Intragenic peaks ^a	5575 (48.6%)			
# Peaks in 1st intron ^a	1852 (16.1%)			
	(33.2% of intragenic)			
# Peaks within extended gene	7808 (68.0%)			
region of annotated genes ^a				
# Peaks >50 kb from a gene	2356 (20.5%)			
# Genes (extended gene region)	5060			
with peaks				
# Genes (extended gene region)	2781 (43.5% of			
with peak and SAGE tag	genes with tag)			

^aPeak maximum falls within these regions.

While relatively few peaks (6%) had a maximum within 1 kb of a TSS, a higher than expected percentage of peaks (33.2%) had a maximum within a first intron. A peak was associated with a gene if the peak's maximum was within a region that extended from 10kb upstream from the gene's TSS to 1 kb downstream of its transcription termination site. Genes with a peak and SAGE tag refers to peaks with maxima within the extended region of a gene that also had SAGE tag(s).

containing Foxa2-binding sequences (Supplementary Figure 3A). As anticipated, with increasing peak heights, the percentage of peaks that contained a Foxa2-binding site sharply increased until a peak height of 10, where the curve of enrichment began to level out. Similar comparisons of randomized peak sequences did not show the same increase in binding sequences with respect to peak height. The level of plateau of peaks occurred at peak height 10, resulting in a dataset of 11 475 high confidence peaks of height 10 or above, of which, 8981 (78%) contained at least one Foxa2-binding sequence (FDR = 0.05). Overall, we identified 22 758 individual binding sequences, with an average of 2.5 binding sequences per peak (Supplementary Figure 3B). The peaks had an average width of 728 bp and an average height of 16 (Table 1). Foxa2-binding site were preferentially found close to the peak maximum (Figure 1A) and most peaks had a Gaussian-like shape, suggesting that the location of the Foxa2-binding site responsible for a peak can be inferred from the shape of the peak and the location of the peak height maximum. Peaks were distributed across the mouse genome with approximately equal numbers of peaks per chromosome when normalized for chromosome length (Supplementary Figure 2A). To further confirm that our peaks were enriched in potential Foxa2 binding sequences, we created a PWM based on our aligned literature-derived Foxa-binding sequences and scored each peak (see Methods section). Significantly, the number of peaks containing a high PWM score was highly enriched in our dataset over randomized peak sequences (P=0) (Figure 1B).

We compared the locations of the high-confidence peaks to UCSC mm8 annotated genes, repetitive regions and conserved genomic elements and found that 68% of peaks were located within an extended gene region

encompassing 10kb upstream of the TSS to 1kb downstream of the 3'UTR. Interestingly, only 6.0% of the peaks were located within 1kb upstream or downstream of the TSS of annotated genes, while 15.7% of the peaks were located within 10 kb upstream of the TSS and 16.1% were localized to a first intron (Table 1). We also observed that 20.5% of all peaks were more than 50kb from any annotated gene (Table 1). Of these peaks, 40% overlapped conserved genomic regions, suggesting that either Foxa2 can act at remote distances from genes or that a number of Foxa2-regulated genes have vet to be annotated. Only 5% of these remote peaks overlapped microsatellite repeats (data not shown). This level of overlap is consistent with only 4% of peaks in the total dataset overlapping microsatellite repeats (data not shown), indicating a low occurrence of peaks over sequence repeat regions. Thus, the Foxa2-binding sites we observed are not preferentially located close to an annotated TSS but are observed at a range of locations across the mouse genome.

Peaks were associated with known Foxa2-target genes and binding sites

To validate our high-confidence peak set in silico, we looked for peaks that overlapped known, characterized Foxa2-binding sites in the mouse liver. Of a set of 16 binding sites described in vitro for the mouse liver (representing 10 target genes) (30-42), our dataset demonstrated peaks overlapping nine sites that were associated with six genes. To confirm that those previously published sites over which we did not see a peak were in fact negative for Foxa2 binding in the adult liver, we investigated the *Hhex* and *PTG* Foxa2-binding sites by qPCR on replicate ChIP experiments. Neither of these sites showed enrichment by qPCR (Hhex = 0.7 average enrichment, PTG = 2.2 average enrichment), indicating little or no Foxa2 binding. This suggests that some Foxa2 sites characterized in vitro are not always occupied in the adult mouse liver in vivo. Of those sites associated with a peak, the location of the site corresponded closely to the location of the peak maximum. From a set of 50 genes known to be regulated by FoxA proteins (where the exact binding site may or may not be characterized) (3,4,9,34,42–67), we observed peaks in the extended gene regions of 36 genes (72%) (Supplementary Table 1). Overall, our success at identifying known Foxa2 targets indicates that our dataset has high coverage of bona fide in vivo Foxa2-binding sites across the genome.

Of the nine well-characterized sites over which we saw peaks, we tested the fold enrichment of five by ChIPqPCR. The sites chosen were located in the Foxa2, transthyretin (Ttr) and cytosolic phosphoenolpyruvate carboxykinase 1 (Pck1) promoters, the albumin (Alb) enhancer and the hepatocyte nuclear factor 4 alpha $(HNF4\alpha)$ enhancer (Figure 2A). Enrichment was observed for all sites, with the greatest enrichment seen at the $HNF4\alpha$ enhancer (52.9-fold) (Figure 2A). Interestingly, the $HNF4\alpha$ promoter site, which has also been reported to be bound by Foxa2, was not overlapped by a peak of height 10 or above and showed enrichment of only 2.5 by qPCR, confirming data shown by Hatzis et al. (36) that the $HNF4\alpha$ promoter site is likely not occupied by Foxa2 in the adult mouse liver in vivo. We also tested additional, previously uncharacterized, sites in the promoter/enhancer regions of Foxa2 regulated genes over which peaks of height 10 and above were observed. These included a site in the Alb gene, 5' to the known enhancer and promoter sites (Figure 3A), as well as peaks in the alpha fetoprotein (Afp), vitronectin (Vtn) and tyrosine aminotransferase (*Tat*) genes. All four showed enrichment by qPCR, ranging from 14- to 30-fold (Figure 3B), indicating that we were able to identify additional binding sites associated with established Foxa2-target genes.

Many genes expressed in the liver contained Foxa2-binding sites

To correlate Foxa2 binding with gene expression, we determined the number of genes that are both transcribed in the liver and potentially regulated by Foxa2. We generated a transcriptional profile of active genes by constructing a longSAGE library from adult mouse liver, using the protocol described previously (27,68). We sequenced this library to a depth sufficient to detect abundant and moderately abundant transcripts (66 272 high quality tag counts). At this depth, many transcripts are represented by only one tag (singletons). However, these have previously been shown to represent bona fide transcripts (27). These tag counts represented 14 026 tag sequence types that mapped to 6396 annotated genes (NCBI Build 36, UCSC mm8). We then overlapped data from this library to our ChIP-Seq dataset to identify genes that were both expressed and associated with in vivo Foxa2 binding (Supplementary Figure 4). We found that 2781 of these genes had one or more Foxa2-binding site(s), suggesting that 43.5% of genes expressed in the adult liver can be associated with an active Foxa2-binding site.

We used Gene Ontology (GO) analysis (FatiGO, fatigo.bioinfo.cipf.es) to identify shared functional properties for the group of expressed genes that were associated with at least one Foxa2 peak (Supplementary Table 2) (69). Six of the top 10, hierarchical level 5, Biological Process categories were associated with metabolism, indicating that these genes were primarily involved in the main biological function of the adult liver. However, we also noted that 'transcription' ranked fourth in the top ten Biological Process categories (16.8% of genes), and that 'transcription factor activity' ranked sixth in the top ten, hierarchical level 5, Molecular Function categories (6.5% of genes). This suggests that those liverexpressed genes potentially regulated by Foxa2 also involved in transcriptional processes and supports the role of Foxa2 as a major transcription factor regulating gene expression in the liver.

We further characterized ChIP-Seq's ability to identify novel Foxa2-binding sites using ChIP-qPCR on biological triplicates. We tested peaks that ranged in heights and locations, including peaks both with and without a Foxa2-binding sequence. While all peaks chosen were associated with genes that our SAGE results indicated were transcribed in the liver, we divided the genes into those that had published evidence for expression or

Α	Gene name	Refseq ID	Peak?	Peak height	Fold Enrichment	Reference
	Foxa2	NM_010446	Yes	19	35.4	Samadani and Costa, 1996
	Albumin	NM_009654	Yes	17	16.1	Bossard and Zaret, 1998, Jackson et al., 1993
	HNF4α (P)	NM_008261	No	N/A	2.5	Bailly et al., 2001
	HNF4α (En)	NM_008261	Yes	43	52.9	Hatzis and Talianidis, 2001
	Ttr	NM_22139	Yes	22	24.5	Costa and Garyson, 1991, Qian et al., 1995
	Pck	NM_011044	Yes	11	3.7	O'Brien et al., 1995

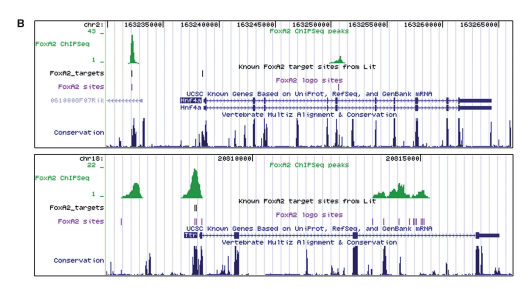


Figure 2. Characterized target Foxa2-binding sites. (A) Table of six characterized target sites in five target genes showing peak presence or absence, peak height and fold enrichment by qPCR. All three sites that contained a peak also showed enrichment above 2-fold. (B) UCSC genome browser mm8 screenshots of four target sites at two of the target genes. Of the $HNF4\alpha$ enhancer and promoter sites, only the enhancer was covered by a peak in our adult liver dataset. The Ttr gene has two closely related promoter sites from different publications, both of which were overlapped by a single peak in our dataset. An additional track displaying the Foxa2-like binding sequence sites from scanning our 10-mer model is also shown ('Foxa2 logo sites').

function in the liver and those that did not (70–77). Of the 16 sites tested, 14 showed an enrichment of >3-fold by qPCR (Figure 4A and B). Of peaks associated with genes with characterized roles in the liver, the greatest enrichment of 30-fold was seen for ApoA2, which is a member of the apolipoprotein family, involved in metabolism. Among the genes lacking characterized roles in the liver, the greatest enrichments of 22.7- and 14.8-fold were seen for the undefined Gtl3 and AI132487 genes, respectively. Of the 16 sites characterized by qPCR, only two did not show enrichment above 3-fold: Cask (does not contain a Foxa2-binding sequence) and Dock8 (contains a Foxa2-binding sequence) (1.2- and 1.5-fold enrichment, respectively). Interestingly, 10 out of the 11 peaks that did not contain Foxa2-binding sequences did show enrichment by qPCR, indicating that the presence of such a sequence within DNA is not essential Foxa2 binding. This observation suggests that there are unidentified Foxa2-binding sequences located within these peaks, or that Foxa2 is localized to these regions through protein protein interactions.

As shown earlier, >15% of genes associated with Foxa2-binding sites are involved in transcription as classified by GO. This observation supports Foxa2 having a role in regulating downstream transcription factors, consistent with results that show regulatory relationships between a core group of six liver transcription factors (9,12,13). To identify novel regulatory cascades involving Foxa2, we measured qPCR fold enrichments for 10 Foxa2 peaks that were associated with genes encoding transcription factor proteins. Those genes with characterized roles in the liver included GATA binding protein 4 (Gata4), GATA binding protein 6 (Gata6), nuclear receptor subfamily 0, group B, member 2 (Nr0b2), one cut domain family member 2 (Onecut2), MAD homolog 1 (Smad1) and Cbp/p300-interacting transactivator, with Glu/Asp-rich carboxy-terminal domain, 2 (Cited2) (78–82). Those without known roles in the liver included TG-interacting factor (TGIF), retinoic acid receptor, beta (Rarb), hairy and enhancer of split 1 (Hes1) and zinc-finger protein 84 (Zfp84). Peak heights ranged from the minimum threshold of 10 up to 28; qPCR enrichment was observed for all 10 peaks, ranging from 5.1 for *Gata4* to 33.7 for *Rarb* (Figure 4C and D). These results further supported a role for Foxa2 in transcriptional cascades in the liver and suggest that

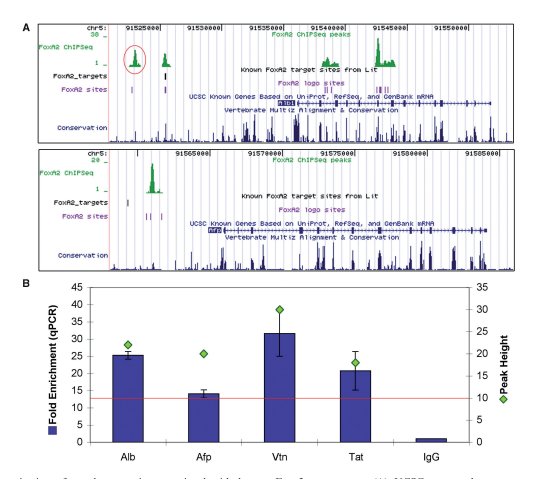


Figure 3. Characterization of novel target sites associated with known Foxa2-target genes. (A) UCSC genome browser screenshots of peaks associated with the Alb and Afp genes. The Alb novel site is an additional location to the characterized target site; both sites were overlapped by peaks, indicating binding by Foxa2 in the liver. The Afp novel site is an alternate location 5' to the site characterized in embryonic liver. (B) QCPR fold enrichment for four novel target sites in characterized Foxa2-target genes averaged over three biological replicate Foxa2 ChIP experiments. Both the fold enrichment and the peak height are shown; the red line indicates the minimum peak height threshold for our dataset (i.e. 10 XSETs). All enrichments were calculated using IgG enrichment as a control in all graphs showing qPCR results. All four novel target sites had fold enrichments above 13.

peaks associated with transcription factor genes in our dataset had a high probability of representing in vivo Foxa2 cross-transcription factor promoter binding sites.

One of the strengths of our genome-wide analysis of Foxa2-binding sites was that it had the capability of identifying novel gene regions that were potentially regulated by Foxa2. As mentioned earlier, 20.5% of Foxa2binding sites were located >50 kb from any annotated gene (Table 1). We used qPCR to determine the fold enrichment of six of these peaks, whose heights ranged from 45 to 79. As for the groups described earlier, the peaks included both those with and without Foxa2binding sequences. All six peaks overlapped, or were near, regions of high inter-species sequence conservation and none of the six were located over microsatellite repeats. Figure 5A shows two example peaks; one close to a mouse EST (unknown #4) and the other adjacent to several highly conserved regions (unknown #5). Fold enrichments by qPCR ranged from 19.1 for unknown #5 to 75.9 for unknown #6 (Figure 5B).

In summary, we tested 41 identified peaks by ChIP coupled with qPCR, corresponding to heights ranging from 10 to 79 from our high-confidence dataset. We also tested six negative sites selected from gaps between peaks of height 2. None of the six negative sites showed qPCR enrichment, while 39 of 41 peaks (95%) showed enrichments greater than 3-fold. Both of the peaks that showed no enrichment by qPCR (Cask and Dock8) had a height of 11, close to our minimal height cutoff. However, eight of the 10 sites chosen for qPCR investigation that had peak heights of 10 or 11 did show enrichment, indicating that, even at and near the peak height threshold of 10, at least 80% of peaks represented in vivo Foxa2-binding sites. We plotted the mean fold enrichment of three biological replicates for each qPCR-validated site against the height of the corresponding peak (Figure 6). Significantly, we observed a strong positive correlation ($R^2 = 0.698$, P = 1.5E-13), demonstrating that peak height was a good indicator of the likely fold enrichment in biological replicates as measured by qPCR. We also tested an additional 33 sites by ChIP-qPCR consisting of both sites from peaks below the height 10 threshold cutoff, as well as sites predicted to show Foxa2 binding from previous publications where no peak was seen (e.g. Hhex) (data not shown). From the 81 total sites tested, using peaks of height 10 and above as true positives, we calculated

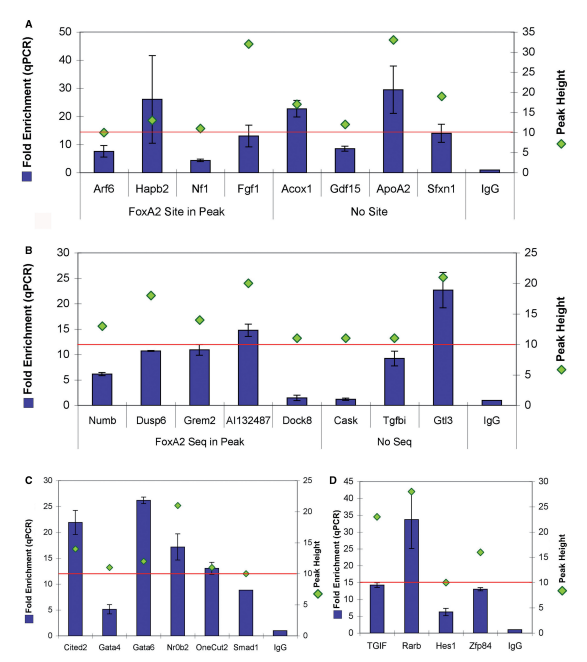


Figure 4. QPCR fold enrichment of annotated genes containing Foxa2-binding site peaks. All genes shown contained SAGE tag(s) indicating that they were expressed in the adult liver. QPCR fold enrichment and peak height are shown for each target; the red line indicates the peak height threshold of 10 XSETs. As previously, IgG enrichment served as the control. (A) Annotated genes that have characterized roles in liver function. Peaks are divided into those that contained a Foxa2-binding sequence and those that did not. All eight targets showed fold enrichments above 4. (B) Annotated genes that have not previously been shown to be expressed in the liver, with peaks divided into those that contained a Foxa2-binding sequence and those that did not. Dock8 and Cask, which did not show enrichment, were the only targets out of 41 peaks tested that were not enriched. The peak height of these two targets was close to the peak height threshold. The remaining six targets all showed fold enrichments above 6. (C) Transcription factors that have a characterized role in liver function in literature. All six targets showed fold enrichments above 5. (D) Transcription factors with uncharacterized roles in the liver. Expression of these transcription factors in the liver is a novel finding. All four targets showed fold enrichments above 5.

a false negative rate of 8.9% and a positive predictive value (PPV) of 94.4%. Together, these results indicate that our dataset, which included peaks that varied widely with respect to location within a gene, height and the presence or absence of a Foxa2-binding sequence. represents a high confidence set of Foxa2-binding sites in the adult mouse liver.

Foxa2 knockdown affects target gene expression

To determine whether Foxa2 binding affects gene expression levels of targets identified by ChIP-Seq, we knocked down Foxa2 in the mouse hepatocellular carcinoma cell line Hepa1-6 by using siRNA pools. This cell line has properties of the developing liver and thus does not

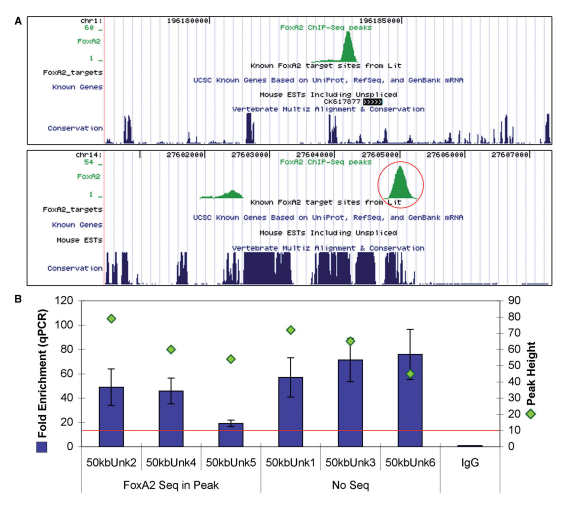


Figure 5. Examples of peaks located >50 kb away from annotated genes. (A) UCSC genome browser mm8 screenshots of two remote peaks in regions containing no known genes. The first peak (unknown #4) lies close to a mouse EST, while the second peak (unknown #5) overlaps and is close to highly conserved sequence regions. (B) Graph of qPCR results of all six peaks lying in unknown gene regions. Four peaks contained Foxa2-binding sequences and two peaks did not. As previously, both fold enrichment by qPCR and peak height are shown. Fold enrichments are relative to IgG control. All six peaks showed fold enrichments above 19.

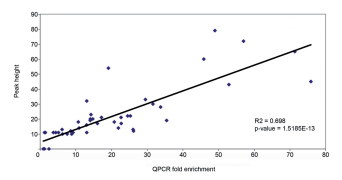


Figure 6. Positive correlation between fold enrichment by qPCR and peak height. Each data point represents the average fold enrichment of three separate Foxa2 ChIP experiments. As indicated, taller peaks showed greater fold enrichments. For this dataset, peak height can be considered to be a predictor of expected fold enrichment.

express all targets investigated above. However, we were able to identify a subset of targets which were expressed. We investigated the expression of nine of these, including both known (Alb, HNF4α, HNF6) and novel (HNF1β, Hes1, Fgf1, Vtn, Cited2, Gtl3) targets, following Foxa2 knockdown. As shown in Figure 7, we were able to decrease Foxa2 expression by $\sim 50\%$, while not significantly affecting levels of Foxa1 or Foxa3 gene expression. Foxa2 protein knockdown was also confirmed by western blotting (data not shown). The knockdown of Foxa2 resulted in significant decrease in expression of six of the nine targets (Alb, HNF4\alpha, HNF6, Hes1, Fgf1, Vtn) and an increase in expression of one—HNF1B. Expression of Cited2 and Gtl3 remained unchanged upon Foxa2 knockdown. These data suggest that many of the Foxa2-binding sites identified can function to regulate gene expression.

Foxa2 peaks contained binding motifs for potential coregulatory transcription factors

Foxa2 can cooperate with other transcription factors to regulate gene expression. To identify potential cofactors, we used 363 vertebrate TFBS models from TRANSFAC Professional v9.3 to scan sequences under the 3016 peaks that contained an exact match to our 10-mer Foxa2binding sequence (see Methods section). The sequences

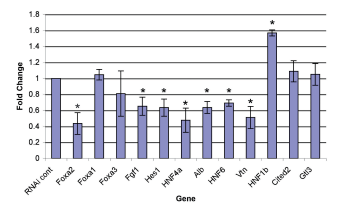


Figure 7. Foxa2 knockdown in Hepa1-6 cells and its effect on expression of targets identified by ChIP-Seq. Foxa2 siRNA treatment results in ~50% knockdown of Foxa2 expression compared to control siRNA. Upon Foxa2 knockdown, expression of both known (Alb, HNF4α) and novel (HNF6, Hes1, Fgf1, Vtn) decreased. Expression of HNF1β increased, while expression of Cited2 and Gtls3 remained unchanged. The expression of Foxa1 was unaffected by Foxa2 knockdown, indicating the specificity of the Foxa2 siRNA. *P < 0.05.

scanned extended 250 bp on either side of a peak's Foxa2binding sequence(s). The 20 TFBS models with the highest enrichment rankings are shown in Table 2; each of these had an estimated enrichment P < 0.05. While the list included Fox TFBS models, as expected, it also included the transcription factors HNF4, HNF1, CoupTF, GATA4 and Pax6, which are all known to cooperate with Foxa2 (12,83,84). In addition, the list identified nine novel potential cofactors that included a wide variety of transcription factor types. We determined whether individual transcription factors were expressed in the mouse adult liver using a combination of SAGE data or reverse transcription coupled with PCR. Twelve out of the top 20 transcription factors were expressed in the liver, and a further four transcription factor families were represented by TFBS models.

We generated distribution plots of the distance between TFBS model sites and the Foxa2-binding sequence and found that 14 out of the top 20 had a significantly nonuniform distribution (P = 1.60E-181-0.03). GATA4 motifs had a relatively uniform distribution across the 500 bp surrounding the Foxa2-binding sequences (Figure 8A), while Pax6 motifs were primarily excluded from the region of the Foxa2-binding sequence out to a distance of \sim 40 bp (Figure 8B). In contrast, HNF1 motifs were clustered around the Foxa2-binding sequence (Figure 8C). The distribution of cotranscription factor motifs relative to Foxa2 sequences may reflect the type of interaction that occurs between the two transcription factors.

We used the same approach to identify motifs that were enriched in the 2494 peaks that contained no Foxa2binding sequence match, to either the 10-mer model ± 1 bp mismatch or the full 14-mer model ± 2 bp mismatch (24). We examined 500 bp sequences that were centered on each peak's maximum for the presence of each of the 363 TRANSFAC TFBS motifs. The HNF3beta (i.e. Foxa2) TRANSFAC model was assigned a very low rank (251 out of 363 models) indicating that our

site-based scanning approach for identifying Foxa2binding sequences was consistent with a TRANSFAC PWM scanning approach. Ten of the 20 highest ranked TFBS models in peaks without Foxa2-binding sites were also among the top 20 models in the peaks that contained Foxa2-binding sequences, and the genes for six of these transcription factors were expressed in the mouse adult liver (Table 2). Of the ten transcription factors corresponding to models that were exclusive to peaks without a Foxa2-binding sequence, five were expressed in the liver. These data provide strong evidence for transcription factor interactions and identify several transcription factors as future targets for investigating Foxa2 interaction.

DISCUSSION

ChIP-Seq identified Foxa2-binding sites throughout the genome

Using ChIP-Seq, we have identified over 11 000 highly confident in vivo Foxa2-binding sites in the adult liver. We were able to show a strong positive correlation between peak height in the ChIP-Seq data and the fold enrichment of binding sites as measured using ChIPqPCR. This demonstrates that peak height can be an indicator of expected fold enrichment and supports our definition of a high-confidence dataset. The stringency used in analysis of Foxa2-binding sequences identified high affinity, high quality Foxa2-binding regions. However, it is likely that a more relaxed binding motif would have identified a greater number of lower confidence Foxa2-binding sequences in peaks.

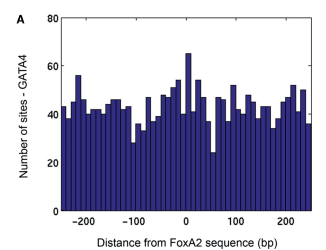
A previous study using ChIP-chip to investigate TFBS in the liver identified 574 Foxa2-bound promoters using a microarray that tiled 10 kb around the TSS of 4000 mouse genes (11). Of the 579 ChIP-Seq peaks present in the regions defined in the ChIP-chip study, >50% of them overlapped with Foxa2-bound regions in the thresholded arrayed dataset (Supplementary Figure 5) (11). This level of overlap is consistent with previous comparisons between ChIP-Seq and ChIP-chip (18). However, since ChIP-Seq can cost effectively sample the entire genome, it has the ability to identify binding sites that occur in regions not often included in promoter microarrays. For example, our study identified over 2300 peaks occurring greater than 50 kb away from any annotated gene. We further observed that 952 of these peaks overlapped regions of high conservation, suggesting that the functions of many of these Foxa2-binding sites have been evolutionarily conserved. These peaks may regulate nearby genes that have not yet been annotated or they may mark enhancer regions that are located far from the genes that they regulate. Alternatively, these sites could represent regions where Foxa2 binds to regulate the chromatin state (8). A further possibility is that these sites are nonfunctional, representing either fortuitous binding or regulatory regions that have been lost through evolution. Further analysis of these regions is necessary to fully define the roles of these Foxa2-binding sites. These remote peaks demonstrate that genome-wide analysis of

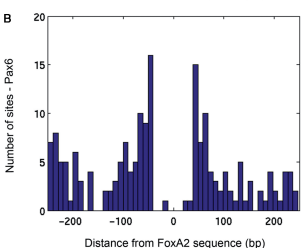
Table 2. The top 20 enrichment-ranked TRANSFAC v9.3 TFBS models that co-occurred within ±250 bp of Foxa2-binding sequences within peaks or occurred in within ± 250 bp of a peak maximum in peaks without Foxa2 sites

TFBS model	Model #	No. of peaks	Z-score	TF in liver?	No. of genes	Level 5 GO Biological Process	Percentage (%)
Peaks with exact Fo	xA2 sequence	•					
FOX_Q2	M00809	995	231.3	Family	621	Cellular metabolic process	25.0
PAX6_Q2	M00979	138	161.3	No	102	Biopolymer modification/Cellular protein metabolic process	29.0
HFH3_01	M00289	448	140.9	No	302	RNA metabolic process/Cellular protein metabolic process	21.4
MAZR 01	M00491	139	106.3	Yes (PCR)	83	RNA metabolic process	30.8
HNF3B 01	M00131	569	82.4	Yes (tag)	340	Cellular protein metabolic process	26.2
HNF4_Q6_01	M01031	791	58.8	Yes (tag)	462	Cellular protein metabolic process	24.0
MAZ Q6	M00649	245	56.8	Yes (tag)	162	Regulation of cellular metabolic process	29.4
HNF4_DR1_Q3	M00764	754	53.3	Yes (tag)	462	RNA metabolic process	22.9
SMAD3 Q6	M00704 M00701	97	44.0	Yes (PCR)	64	Cellular protein metabolic process	33.3
COUPTF Q6	M01036	479	43.8	Family	270	Cellular protein metabolic process	21.3
PAX Q6	M00808	706	41.5	Family	433	Cellular protein metabolic process	26.5
		844			433 465		20.3
HNF1_Q6	M00790		41.2	Yes (tag)		Regulation of cellular metabolic process	
GATA4_Q3	M00632	1285	37.6	Yes (tag)	700	Cellular protein metabolic process	24.9
HNF1_C	M00206	1029	34.8	Yes (tag)	549	Cellular protein metabolic process	29.7
EVI1_04	M00081	1028	33.5	Yes (PCR)	565	Cellular protein metabolic process	25.9
PPARG_03	M00528	522	32.2	Yes (tag)	310	RNA metabolic process	19.3
LDSPOLYA_B	M00317	240	30.5	No	129	RNA metabolic process	28.8
FREAC2_01	M00290	34	30.5	No	23	Regulation of cellular metabolic process/RNA meta- bolic process/Transcription	43.8
KROX Q6	M00982	13	29.0	Family	11	Cell surface receptor linked signal transduction	50
FOXJ2_01	M00422	124	27.6	Yes (tag)	75	RNA metabolic process/Regulation of cellular meta- bolic process	23.2
Peaks with no FoxA		100					
MAZR_01	M00491	138	82.9	Yes (PCR)	77	Cellular protein metabolic process	29.2
PAX6_Q2 MAZ_Q6	M00979 M00649	69 269	57.0 47.0	No Yes (tag)	44 164	Intracellular signaling cascade RNA metabolic process/Transcription/Regulation of	31.8 31.2
						cellular metabolic process	
KROX_Q6	M00982	25	39.0	Family	20	Regulation of cellular metabolic process/RNA meta- bolic process/Transcription/Cellular protein meta-	33.3
HNE4 OC 01	M01021	576	21.7	Vac (to a)	261	bolic process	25.7
HNF4_Q6_01	M01031			Yes (tag)	361	Cellular protein metabolic process	
UF1H3BETA_Q6	M01068	2	19.9	No Var (tan)	1	15 different terms ^a	100
HNF4_DR1_Q3	M00764	537	17.4	Yes (tag)	321	Cellular protein metabolic process	21.9 19.2
SMAD3_Q6	M00701	51 5	15.5 14.9	Yes (PCR)	31 6	Intracellular signaling cascade/RNA metabolic process	33.3
NRF1_Q6	M00652	3	14.9	Yes (tag)	0	Response to oxidative stress/Carboxylic acid metabolic process/Cellular protein metabolic process/ Mitochondrion organization and biogenesis/ Macromolecule biosynthetic process/DNA metabolic process	33.3
COUPTF Q6	M01036	333	14.7	Family	225	Cellular protein metabolic process	26.6
MIF_01	M00279	6	12.3	Yes (PCR)	4	Cytoskeleton organization and biogenesis/ Establishment of cellular localization	100
SMAD4_Q6	M00733	46	11.8	Yes (tag)	29	Phosphate metabolic process/Cellular protein metabolic process	28.6
ATF_B	M00338	4	11.7	Family	3	RNA metabolic process/Regulation of cellular meta- bolic process/Transcription	100
PPARG 03	M00528	423	11.1	Yes (tag)	246	Cellular protein metabolic process	24.8
PAX_Q6	M00808	483	10.9	Family	287	Cellular protein metabolic process	30.9
SF1_Q6	M00727	118	10.9	Yes (tag)	83	Regulation of cellular metabolic process	32.6
BRN2 01	M00727 M00145	75	10.8	No	21	Regulation of cellular metabolic process	50
RREBI_01	M00257	9	10.4	Yes (tag)	7	Regulation of cellular metabolic process/RNA meta- bolic process/Transcription	50
SP3 Q3	M00665	59	10.0	Yes (tag)	36	Cellular protein metabolic process	47.6
LHX3 01	M00510	11	8.3	No	5	Regulation of cellular metabolic process/RNA meta-	66.7
LII/LU UI	14100210	1.1	0.5	110	J	bolic process/Transcription	00.7

Included in the table are: (i) the number of peaks that contained the TFBS model; (ii) the Z-score, which represents the difference between the MATCH v2.6 PWM scan enrichment of real peak sequences and the mean for randomized sequences, divided by the standard deviation for randomized sequences (see Methods section); (iii) an indication of whether the transcription factor is expressed in the adult liver, based either on SAGE tag(s) from our adult liver library or RT-PCR amplification in the adult liver; (iv) the number of annotated genes that contained at least one peak within its extended gene region (10 kb upstream of the TSS to 1kb downstream of the TTS); (v) the 'level 5' Gene Ontology Biological Process category assigned to the largest subgroup within these gene groups and (vi) the corresponding percentage of genes in the subgroup. TFBS models that co-occurred with a Foxa2-binding sequence included both transcription factors that have previously been demonstrated to interact with Foxa2 (HNF4 and members of the Coup-TF family) and also novel potential interacting transcription factors (Smad3 and Maz). Ten of the TFBS models that co-occurred with Foxa2-like binding sequences in peaks containing such sequences also occurred in peaks that did not contain Foxa2-like binding sequences.

^aRegulation of biological process; cellular metabolic process; macromolecule metabolic process; primary metabolic process; cell organization and biogenesis; regulation of metabolic process; regulation of cellular process; organelle organization and biogenesis; biopolymer metabolic process; nucleobase, nucleoside, nucleotide and nucleic acid metabolic process; chromosome organization and biogenesis; regulation of cellular metabolic process; RNA metabolic process; transcription; DNA metabolic process.





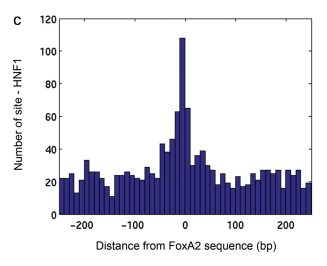


Figure 8. The distribution of TFBS sequence models relative to the corresponding Foxa2-binding sequence for three of the top 20 cotranscription factors in peaks containing a Foxa2-binding sequence. (A) The distribution of GATA4 TFBS models (Transfac model M00632). Motifs for this transcription factor show a uniform distribution across the 500 bp sequence. (B) The distribution of Pax6 TFBS models (Transfac model M00979). Pax6 motifs clustered at a distance of \sim 40 bp away from the Foxa2-binding sequence (P=1.1E-3). (C) The spatial distribution of HNF1 TFBS models (TRANSFAC model M00790). HNF1 motifs are spatially concentrated near the Foxa2-binding sequence (P=5.7E-9).

TFBS is critical for a full understanding of *in vivo* gene regulatory mechanisms.

Identification of *in vivo* Foxa2-binding sites suggests mechanisms of gene regulation

We observed that close to 45% of genes shown to be expressed in the liver by SAGE are associated with a Foxa2 peak, suggesting that a large number of these genes are potentially regulated by Foxa2. Although Foxa2 is known to be a major regulator of the hepatocyte phenotype (14), the extent of coverage of these liverexpressed genes by in vivo Foxa2-binding sites is surprising. Our results may actually underestimate the number of genes regulated by Foxa2, since genes that are expressed at levels not detected by SAGE, and therefore not represented by a tag, are not included. Of interest, Foxa2 has been shown to function by controlling DNA accessibility for transcriptional machinery in the embryo. Through its ability to bind to core histone proteins present in closed chromatin, it can render DNA accessible to additional transcription factor binding and therefore initiate transcriptional activity (8). The large number of binding sites for Foxa2 in the liver may reflect a role for Foxa2 in maintaining the competence for gene expression critical for hepatocyte function rather than a direct regulatory relationship. Of interest, we observed Foxa2 binding in the second intron of Gata4 and the cooccurrence of the GATA4 TFBS model with Foxa2-binding sites, suggesting a coregulatory interaction between these two factors. Given that GATA4 is also able to bind closed chromatin with Foxa2, this supports the role for both in the regulation of chromatin structure (8). Another possibility for the large number of Foxa2-binding sites observed close to liver expressed genes may also represent fortuitous binding of Foxa2 in regions of open chromatin rather than functional binding.

While our analysis suggested a major role for Foxa2 in liver function, deletion of Foxa2 specifically in the liver has shown this transcription factor is not essential for adult liver functions (85). However, combination knockout and conditional knockout studies have shown that, at least in the embryo, either Foxa2 or the closely related family member, Foxal, is necessary for embryonic liver development (86). In addition, the three FoxA family members recognize similar binding sequences (5), suggesting that at least Foxa1 and Foxa2 can function redundantly. The level of redundancy between these three family members reflects the large roles that they play in the formation and functioning of the liver. Our analysis of Foxa2 knockdown in the Hepa1-6 cell line indicates that Foxa2 directly regulates the level of gene expression for at least some of novel gene targets identified that are associated with the Foxa2-binding sites. Additionally, given that Foxal expression levels were unaffected by Foxa2 knockdown in these experiments, this suggests that for a subset of genes Foxal is unable to completely replace reduction of Foxa2 with respect to gene expression regulation. These genes may be sensitive to reduction of FoxA, since Foxal is expressed at lower levels than Foxa2 in these cells. Alternatively, these genes may be preferentially

regulated by Foxa2. The expression of some genes associated with Foxa2 binding was not affected by Foxa2 knockdown, indicating one of several possibilities suggested above including replacement of Foxa2 function by Foxa1, a role for Foxa2 in maintaining open chromatin rather than direct regulation of gene expression, or the presence of other transcription factors which are more critical to the regulation of these genes.

In addition to the global analysis of Foxa2-binding sites, our data suggest novel mechanisms of regulation at the level of individual genes. For example, we observed that the $HNF4\alpha$ enhancer site that binds Foxa2 is located in the first intron of a Riken gene (0610008F07Rik) on the opposite chromosomal strand, encoding a hypothetical, unclassified protein. This Riken gene is expressed in our adult liver SAGE library. Thus, this Foxa2-binding site may be a dual regulatory site involved in the promotion of transcription of two genes, a phenomenon which has been previously documented in other systems (87). Another example is seen with Afp, where there is no peak over the characterized Foxa2 site in the enhancer region that regulates embryonic expression of Afp. However, a peak was observed lying between the characterized site and the Afp TSS, \sim 5 kb upstream of the Afp TSS. Since Afp is not expressed in adult liver, this indicates that the binding of Foxa2 is not sufficient for Afp transcription and suggests that Foxa2 binding at this alternative site may act in a repressive manner. Interestingly, previous studies have demonstrated that the Afp gene is repressed in the adult liver via a competitive mechanism between FoxA and p53 (88). The occurrence of alternate sites for the same transcription factor acting as an inducer or a repressor depending on the site it binds to, has been well documented in literature. For example, the transcription factor Sox9 occupies different inductive or repressive binding sites on the collagen 2a1 gene depending on the activation state of chondrocytes (89).

Foxa2 functions in a transcriptional network

A number of ChIP-chip studies that characterize crossregulation between the promoters of several well characterized liver transcription factors have shown a high level of interaction between a core group of thirteen transcription factors important to liver development and function—HNF1α, HNF1β, HNF4α, Foxa1, Foxa2, HNF6, C/EBPγ, COUP-TFII, LRH1, USF1, PXR, FXRα and GATA6 (9,12,14). Our data showed *in vivo* Foxa2-binding sites within the extended gene region of nine of these 13 core transcription factors, including three transcription factors that had not previously been shown to be bound by Foxa2, indicating a high level of potential cross regulation by Foxa2 in the liver. We also documented the presence of Foxa2-binding sites associated with many additional transcription factors expressed in the adult liver. These results suggest that in the adult liver, Foxa2 is involved in additional transcriptional cascades that contribute to liver function.

In addition to Foxa2-binding sites in the promoters of other transcription factors, Foxa2 has been shown to interact with transcription factor proteins to coordinately regulate gene expression. Studies have indicated simultaneous occupation of gene promoters by Foxa2 in combination with HNF4α, HNF6, CREB1, USF1, HNF1α or GATA4 suggesting cooperative interaction between Foxa2 and these transcription factors (12,31). To extend our knowledge of transcription factors that could interact with Foxa2 in the adult liver, we determined the 20 most highly enriched cooccurring TFBS models that occurred within 250 bp of Foxa2 sites. This group included many transcription factors known to interact with Foxa2, including HNF4, HNF1 and GATA4, as well as novel transcription factors such as Maz, Mazr and Smad3, all of which are expressed in the adult liver as shown by SAGE or RT-PCR. Of interest, several TFBS models were observed in both peaks with and without Foxa2binding sequences. These included Pax6, Maz, Mazr, Smad3, HNF4 and Pparg, of which all but Pax6 are expressed in the adult liver. One possibility for the absence of a Foxa2-binding site in some peaks is a physical regulatory interaction, whereby Foxa2 itself is bound not directly to the DNA, but to a cointeracting transcription factor at a close enough distance to enable protein—protein chemical cross-linking. Smad3 is thought to be primarily involved in acute liver damage via TGFβ signaling, while Pparg is involved in lipid metabolism (90,91). Additionally, Pparg may inhibit TGFβ/Smad3 signaling in liver fibrosis (92). Evil has been previously shown to be expressed in the adult liver; however, neither Maz nor Mazr have been reported to be expressed in the liver. The functional role of these three transcription factors in liver gene regulation is currently unknown (93). The fact that many of these top cooccurring transcription factors are expressed and have roles in the adult liver further supports the likelihood that they can interact with Foxa2.

Although Pax6 is not expressed in the liver, it is expressed in the closely related pancreatic tissue. Notably, the Pax6 TFBS model was found amongst the top 20 enriched cooccurring transcription factors in both peaks with and without an exact Foxa2-binding site. In the pancreas, Foxa2 and Pax6 interact to regulate the activation of genes such as glucagon and Pdx1 (83,94). Of interest, the location of the Pax6 TFBS model in relation to Foxa2-binding sequences showed that the Pax6-binding site was consistently greater than 40 bp away from Foxa2 in either direction (Figure 7B). Thus, although Foxa2- and Pax6-binding sites are found in close proximity, at least 40 bp must be present between the two sites, suggesting that Pax6 and Foxa2 can bind together to regulate gene expression in the pancreas or that noncompetitive inhibitory mechanisms limit the binding of both factors simultaneously.

This study has presented novel Foxa2 regulatory mechanisms in the mouse adult liver by determining genome wide binding sites. We have also shown a high level of coverage of known functional binding sites as well as identifying several potential novel interactions with annotated genes and unannotated genomic regions. Finally, we have shown a potentially wide-ranging role for Foxa2 in regulating gene expression in the function of the adult liver. Overall, this study demonstrates the advantages of using ChIP-Seq technology to characterize

transcription factor binding in tissue, and paves the way for further investigations of in vivo transcription factor networks.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

The authors gratefully acknowledge the support of the MORGEN project team managers, Amanda Kotzer and Joanne Johnson. This study received funding support from the Genome British Columbia and Genome Canada. P.A.H., M.M. and S.J. are Scholars of the Michael Smith Foundation for Health Research. M.M. is a Terry Fox Young Investigator. P.A.H. is a Canadian Institutes for Health Research New Investigator. Funding to pay the Open Access publication charges for this article was provided by Genome Canada and Genome BC.

Conflict of interest statement. None declared.

REFERENCES

- 1. Ang, S.L., Wierda, A., Wong, D., Stevens, K.A., Cascio, S., Rossant, J. and Zaret, K.S. (1993) The formation and maintenance of the definitive endoderm lineage in the mouse: involvement of HNF3/forkhead proteins. Development, 119, 1301-1315.
- 2. Rausa, F.M., Tan, Y., Zhou, H., Yoo, K.W., Stolz, D.B., Watkins, S.C., Franks, R.R., Unterman, T.G. and Costa, R.H. (2000) Elevated levels of hepatocyte nuclear factor 3beta in mouse hepatocytes influence expression of genes involved in bile acid and glucose homeostasis. Mol. Cell. Biol., 20, 8264-8282.
- 3. Wolfrum, C., Asilmaz, E., Luca, E., Friedman, J.M. and Stoffel, M. (2004) Foxa2 regulates lipid metabolism and ketogenesis in the liver during fasting and in diabetes. Nature, 432, 1027-1032.
- 4. Costa, R.H., Grayson, D.R. and Darnell, J.E.Jr. (1989) Multiple hepatocyte-enriched nuclear factors function in the regulation of transthyretin and alpha 1-antitrypsin genes. Mol. Cell. Biol., 9, 1415-1425
- 5. Lai, E., Prezioso, V.R., Smith, E., Litvin, O., Costa, R.H. and Darnell, J.E.Jr. (1990) HNF-3A, a hepatocyte-enriched transcription factor of novel structure is regulated transcriptionally. Genes Dev., **4**, 1427–1436.
- 6. Lee, C.S., Friedman, J.R., Fulmer, J.T. and Kaestner, K.H. (2005) The initiation of liver development is dependent on Foxa transcription factors. Nature, 435, 944-947.
- 7. Friedman, J.R. and Kaestner, K.H. (2006) The Foxa family of transcription factors in development and metabolism. Cell Mol. Life Sci., 63, 2317-2328.
- 8. Cirillo, L.A., Lin, F.R., Cuesta, I., Friedman, D., Jarnik, M. and Zaret, K.S. (2002) Opening of compacted chromatin by early developmental transcription factors HNF3 (FoxA) and GATA-4. Mol. Cell, 9, 279-289.
- 9. Kyrmizi, I., Hatzis, P., Katrakili, N., Tronche, F., Gonzalez, F.J. and Talianidis, I. (2006) Plasticity and expanding complexity of the hepatic transcription factor network during liver development. Genes Dev., 20, 2293-2305.
- 10. Gao, H., Falt, S., Sandelin, A., Gustafsson, J.A. and Dahlman-Wright, K. (2008) Genome-wide identification of estrogen receptor {alpha} binding sites in mouse liver. Mol. Endocrinol., 22, 10–22.
- 11. Odom, D.T., Dowell, R.D., Jacobsen, E.S., Gordon, W., Danford, T.W., MacIsaac, K.D., Rolfe, P.A., Conboy, C.M., Gifford, D.K. and Fraenkel, E. (2007) Tissue-specific transcriptional regulation has diverged significantly between human and mouse. Nat. Genet., 39, 730-732.

- 12. Odom, D.T., Dowell, R.D., Jacobsen, E.S., Nekludova, L., Rolfe, P.A., Danford, T.W., Gifford, D.K., Fraenkel, E., Bell, G.I. and Young, R.A. (2006) Core transcriptional regulatory circuitry in human hepatocytes. Mol. Syst. Biol., 2, 2006.0017.
- 13. Odom, D.T., Zizlsperger, N., Gordon, D.B., Bell, G.W., Rinaldi, N.J., Murray, H.L., Volkert, T.L., Schreiber, J., Rolfe, P.A., Gifford, D.K. et al. (2004) Control of pancreas and liver gene expression by HNF transcription factors. Science, 303, 1378-1381.
- 14. Rada-Iglesias, A., Wallerman, O., Koch, C., Ameur, A., Enroth, S., Clelland, G., Wester, K., Wilcox, S., Dovey, O.M., Ellis, P.D. et al. (2005) Binding sites for metabolic disease related transcription factors inferred at base pair resolution by chromatin immunoprecipitation and genomic microarrays. Hum. Mol. Genet., 14, 3435-3447.
- 15. Rubins, N.E., Friedman, J.R., Le, P.P., Zhang, L., Brestelli, J. and Kaestner, K.H. (2005) Transcriptional networks in the liver: Hepatocyte nuclear factor 6 function is largely independent of Foxa2. Mol. Cell. Biol., 25, 7069-7077.
- 16. Smith, A.D., Sumazin, P. and Zhang, M.Q. (2007) Tissue-specific regulatory elements in mammalian promoters. Mol. Syst. Biol., 3,
- 17. Teufel, A., Weinmann, A., Krupp, M., Budinger, M. and Galle, P.R. (2007) Genome-wide analysis of factors regulating gene expression in liver. Gene, 389, 114-121.
- 18. Lin, C.Y., Vega, V.B., Thomsen, J.S., Zhang, T., Kong, S.L., Xie, M., Chiu, K.P., Lipovich, L., Barnett, D.H., Stossi, F. et al. (2007) Whole-genome cartography of estrogen receptor alpha binding sites. PLoS Genet., 3, e87.
- 19. Loh, Y.H., Wu, Q., Chew, J.L., Vega, V.B., Zhang, W., Chen, X., Bourque, G., George, J., Leong, B., Liu, J. et al. (2006) The Oct4 and nanog transcription network regulates pluripotency in mouse embryonic stem cells. Nat Genet, 38, 431-440.
- 20. Wei, Č.L., Wu, Q., Vega, V.B., Chiu, K.P., Ng, P., Zhang, T., Shahab, A., Yong, H.C., Fu, Y., Weng, Z. et al. (2006) A global map of p53 transcription-factor binding sites in the human genome. Cell, **124**, 207–219.
- 21. Zeller, K.I., Zhao, X., Lee, C.W., Chiu, K.P., Yao, F., Yustein, J.T., Ooi, H.S., Orlov, Y.L., Shahab, A., Yong, H.C. et al. (2006) Global mapping of c-myc binding sites and target gene networks in human B cells. Proc. Natl Acad Sci USA, 103, 17834-17839.
- 22. Barski, A., Cuddapah, S., Cui, K., Roh, T.-Y., Schones, D.E., Wang, Z., Wei, G., Chepelev, I. and Zhao, K. (2007) High-resolution profiling of histone methylations in the human genome. Cell, 129, 823-837.
- 23. Johnson, D.S., Mortazavi, A., Myers, R.M. and Wold, B. (2007) Genome-wide mapping of in vivo protein-DNA interactions. Science, 316, 1497-1502.
- 24. Robertson, G., Hirst, M., Bainbridge, M., Bilenky, M., Zhao, Y., Zeng, T., Euskirchen, G., Bernier, B., Varhol, R., Delaney, A. et al. (2007) Genome-wide profiles of STAT1 DNA association using chromatin immunoprecipitation and massively parallel sequencing. Nat. Methods, 4, 651-657.
- 25. Montgomery, S.B., Griffith, O.L., Sleumer, M.C., Bergman, C.M., Bilenky, M., Pleasance, E.D., Prychyna, Y., Zhang, X. and Jones, S.J. (2006) ORegAnno: an open access database and curation system for literature-derived promoters, transcription factor binding sites and regulatory variation. Bioinformatics, 22, 637-640.
- 26. Vlieghe, D., Sandelin, A., De Bleser, P.J., Vleminckx, K., Wasserman, W.W., van Roy, F. and Lenhard, B. (2006) A new generation of JASPAR, the open-access repository for transcription factor binding site profiles. Nucleic Acids Res., 34, D95-D97.
- 27. Siddiqui, A.S., Khattra, J., Delaney, A.D., Zhao, Y., Astell, C., Asano, J., Babakaiff, R., Barber, S., Beland, J., Bohacec, S. et al. (2005) A mouse atlas of gene expression: large-scale digital geneexpression profiles from precisely defined developing C57BL/6J mouse tissues and cells. Proc. Natl Acad. Sci. USA, 102, 18485-18490.
- 28. Kel, A.E., Gossling, E., Reuter, I., Cheremushkin, E., Kel-Margoulis, O.V. and Wingender, E. (2003) MATCH: a tool for searching transcription factor binding sites in DNA sequences. Nucleic Acids Res., 31, 3576-3579.
- 29. van Helden, J., Andre, B. and Collado-Vides, J. (2000) A web site for the computational analysis of yeast regulatory sequences. Yeast, 16, 177-187.

- 30. Bailly, A., Torres-Padilla, M.E., Tinel, A.P. and Weiss, M.C. (2001) An enhancer element 6kb upstream of the mouse HNF4alpha1 promoter is activated by glucocorticoids and liver-enriched transcription factors. Nucleic Acids Res., 29, 3495-505.
- 31. Bossard, P. and Zaret, K.S. (1998) GATA transcription factors as potentiators of gut endoderm differentiation. Development, 125,
- 32. Cheng, A., Zhang, M., Crosson, S.M., Bao, Z.Q. and Saltiel, A.R. (2006) Regulation of the mouse protein targeting to glycogen (PTG) promoter by the FoxA2 forkhead protein and by 3',5'-cyclic adenosine 5'-monophosphate in H4IIE hepatoma cells. Endocrinology, 147, 3606-3612.
- 33. Costa, R.H. and Grayson, D.R. (1991) Site-directed mutagenesis of hepatocyte nuclear factor (HNF) binding sites in the mouse transthyretin (TTR) promoter reveal synergistic interactions with its enhancer region. Nucleic Acids Res., 19, 4139-4145.
- 34. Denson, L.A., McClure, M.H., Bogue, C.W., Karpen, S.J. and Jacobs, H.C. (2000) HNF3beta and GATA-4 transactivate the liverenriched homeobox gene, hex. Gene, 246, 311-320
- 35. Groupp, E.R., Crawford, N. and Locker, J. (1994) Characterization of the distal alpha-fetoprotein enhancer, a strong, long distance, liver-specific activator. J. Biol. Chem., 269, 22178-22187.
- 36. Hatzis, P. and Talianidis, I. (2001) Regulatory mechanisms controlling human hepatocyte nuclear factor 4alpha gene expression. Mol. Cell. Biol., 21, 7320-7330.
- 37. Jackson, D.A., Rowader, K.E., Stevens, K., Jiang, C., Milos, P. and Zaret, K.S. (1993) Modulation of liver-specific transcription by interactions between hepatocyte nuclear factor 3 and nuclear factor 1 binding DNA in close apposition. Mol. Cell. Biol., 13, 2401–2410.
- 38. Nitsch, D., Boshart, M. and Schutz, G. (1993) Activation of the tyrosine aminotransferase gene is dependent on synergy between liver-specific and hormone-responsive elements. Proc. Natl Acad. Sci. USA, 90, 5479-5483.
- 39. O'Brien, R.M., Noisin, E.L., Suwanichkul, A., Yamasaki, T., Lucas, P.C., Wang, J.C., Powell, D.R. and Granner, D.K. (1995) Hepatic nuclear factor 3- and hormone-regulated expression of the phosphoenolpyruvate carboxykinase and insulin-like growth factorbinding protein 1 genes. Mol. Cell. Biol., 15, 1747-1758.
- 40. Qian, X., Samadani, U., Porcella, A. and Costa, R.H. (1995) Decreased expression of hepatocyte nuclear factor 3 alpha during the acute-phase response influences transthyretin gene transcription. Mol. Cell. Biol., 15, 1364-1376.
- 41. Samadani, U. and Costa, R.H. (1996) The transcriptional activator hepatocyte nuclear factor 6 regulates liver gene expression. Mol. Cell. Biol., 16, 6273-6284.
- 42. Shimizu, S., Miyamoto, Y. and Hayashi, M. (2002) Cell-type dependency of two Foxa/HNF3 sites in the regulation of vitronectin promoter activity. Biochim. Biophys. Acta, 1574, 337-344.
- 43. Antes, T.J. and Levy-Wilson, B. (2001) HNF-3 beta, C/EBP beta, and HNF-4 act in synergy to enhance transcription of the human apolipoprotein B gene in intestinal cells. DNA Cell Biol., 20, 67-74.
- 44. Arenzana, N. and Rodriguez de Cordoba, S. (1996) Promoter region of the human gene coding for beta-chain of C4b binding protein. hepatocyte nuclear factor-3 and nuclear factor-I/CTF transcription factors are required for efficient expression of C4BPB in HepG2 cells. J. Immunol., 156, 168-175.
- 45. Beck, S., Sommer, P., dos Santos Silva, E., Blin, N. and Gott, P. (1999) Hepatocyte nuclear factor 3 (winged helix domain) activates trefoil factor gene TFF1 through a binding motif adjacent to the TATAA box. DNA Cell Biol., 18, 157-164.
- 46. Berg, D.T., Gerlitz, B., Sharma, G.R., Richardson, M.A., Stephens, E.J., Grubbs, R.L., Holmes, K.C., Fynboe, K., Montani, D., Cramer, M.S. et al. (2006) FoxA2 involvement in suppression of protein C, an outcome predictor in experimental sepsis. Clin. Vaccine Immunol., 13, 426-432.
- 47. Ceelie, H., Spaargaren-Van Riel, C.C., De Jong, M., Bertina, R.M. and Vos,H.L. (2003) Functional characterization of transcription factor binding sites for HNF1-alpha, HNF3-beta (FOXA2), HNF4alpha, Sp1 and Sp3 in the human prothrombin gene enhancer. J. Thromb. Haemost., 1, 1688-1698.
- 48. Cha,J.Y., Kim,H., Kim,K.S., Hur,M.W. and Ahn,Y. (2000) Identification of transacting factors responsible for the tissuespecific expression of human glucose transporter type 2

- isoform gene. Cooperative role of hepatocyte nuclear factors lalpha and 3beta. J. Biol. Chem., 275, 18358-18365.
- 49. Christoffels, V.M., Grange, T., Kaestner, K.H., Cole, T.J., Darlington, G.J., Croniger, C.M. and Lamers, W.H. (1998) Glucocorticoid receptor, C/EBP, HNF3, and protein kinase A coordinately activate the glucocorticoid response unit of the carbamoylphosphate synthetase I gene. Mol. Cell. Biol., 18, 6305-6315.
- 50. Crowe, A.J., Sang, L., Li, K.K., Lee, K.C., Spear, B.T. and Barton, M.C. (1999) Hepatocyte nuclear factor 3 relieves chromatinmediated repression of the alpha-fetoprotein gene. J. Biol. Chem., **274**, 25113-25120.
- 51. Delesque-Touchard, N., Park, S.H. and Waxman, D.J. (2000) Synergistic action of hepatocyte nuclear factors 3 and 6 on CYP2C12 gene expression and suppression by growth hormoneactivated STAT5b. Proposed model for female specific expression of CYP2C12 in adult rat liver. J. Biol. Chem., 275, 34173-34182.
- 52. Harnish, D.C., Malik, S., Kilbourne, E., Costa, R. and Karathanasis, S.K. (1996) Control of apolipoprotein AI gene expression through synergistic interactions between hepatocyte nuclear factors 3 and 4. J. Biol. Chem., 271, 13621-13628.
- 53. Jung, D., Hagenbuch, B., Fried, M., Meier, P.J. and Kullak-Ublick, G.A. (2004) Role of liver-enriched transcription factors and nuclear receptors in regulating the human, mouse, and rat NTCP gene. Am. J. Physiol. Gastrointest. Liver Physiol., 286, G752-G761.
- 54. Lantz, K.A., Vatamaniuk, M.Z., Brestelli, J.E., Friedman, J.R., Matschinsky, F.M. and Kaestner, K.H. (2004) Foxa2 regulates multiple pathways of insulin secretion. J. Clin. Invest., 114, 512-520.
- 55. Lemaigre, F.P., Durviaux, S.M. and Rousseau, G.G. (1993) Liverspecific factor binding to the liver promoter of a 6-phosphofructo-2kinase/fructose-2,6-bisphosphatase gene. J. Biol. Chem., 268, 19896-19905.
- 56. Li,C., Zhu,N.L., Tan,R.C., Ballard,P.L., Derynck,R. and Minoo,P. (2002) Transforming growth factor-beta inhibits pulmonary surfactant protein B gene transcription through SMAD3 interactions with NKX2.1 and HNF-3 transcription factors. J. Biol. Chem., 277, 38399-38408.
- 57. Raynal, J.F., Dugast, C., Le Van Thai, A. and Weber, M.J. (1998) Winged helix hepatocyte nuclear factor 3 and POU-domain protein brn-2/N-oct-3 bind overlapping sites on the neuronal promoter of human aromatic L-amino acid decarboxylase gene. Brain Res. Mol. Brain Res., 56, 227-237.
- 58. Roux, J., Pictet, R. and Grange, T. (1995) Hepatocyte nuclear factor 3 determines the amplitude of the glucocorticoid response of the rat tyrosine aminotransferase gene. DNA Cell Biol., 14, 385-396.
- 59. Sawaya, P.L. and Luse, D.S. (1994) Two members of the HNF-3 family have opposite effects on a lung transcriptional element; HNF-3 alpha stimulates and HNF-3 beta inhibits activity of region I from the clara cell secretory protein (CCSP) promoter. J. Biol. Chem., 269, 22211-22216.
- 60. Scassa, M.E., Guberman, A.S., Ceruti, J.M. and Canepa, E.T. (2004) Hepatic nuclear factor 3 and nuclear factor 1 regulate 5-aminolevulinate synthase gene expression and are involved in insulin repression. J. Biol. Chem., 279, 28082-28092.
- 61. Unterman, T.G., Fareeduddin, A., Harris, M.A., Goswami, R.G., Porcella, A., Costa, R.H. and Lacson, R.G. (1994) Hepatocyte nuclear factor-3 (HNF-3) binds to the insulin response sequence in the IGF binding protein-1 (IGFBP-1) promoter and enhances promoter function. Biochem. Biophys. Res. Commun., 203, 1835-1841.
- 62. Vallet, V., Antoine, B., Chafey, P., Vandewalle, A. and Kahn, A. (1995) Overproduction of a truncated hepatocyte nuclear factor 3 protein inhibits expression of liver-specific genes in hepatoma cells. Mol. Cell. Biol., 15, 5453-5460.
- 63. Verschuur, M., de Jong, M., Felida, L., de Maat, M.P. and Vos, H.L. (2005) A hepatocyte nuclear factor-3 site in the fibrinogen beta promoter is important for interleukin 6-induced expression, and its activity is influenced by the adjacent -148C/T polymorphism. J. Biol. Chem., 280, 16763-16771.
- 64. Wang, J.C., Stromstedt, P.E., O'Brien, R.M. and Granner, D.K. (1996) Hepatic nuclear factor 3 is an accessory factor required for the stimulation of phosphoenolpyruvate carboxykinase gene transcription by glucocorticoids. Mol. Endocrinol., 10, 794-800.
- 65. Williamson, E.A., Wolf, I., O'Kelly, J., Bose, S., Tanosaki, S. and Koeffler, H.P. (2006) BRCA1 and FOXA1 proteins coregulate the

- expression of the cell cycle-dependent kinase inhibitor p27(Kip1). Oncogene, 25, 1391-1399.
- 66. Wiwi, C.A. and Waxman, D.J. (2005) Role of hepatocyte nuclear factors in transcriptional regulation of male-specific CYP2A2. J. Biol. Chem., 280, 3259-3268.
- 67. Zhu, Q.S., Xing, W., Qian, B., von Dippe, P., Shneider, B.L., Fox, V.L. and Levy, D. (2003) Inhibition of human m-epoxide hydrolase gene expression in a case of hypercholanemia. Biochim. Biophys. Acta, 1638, 208-216.
- 68. Khattra, J., Delaney, A.D., Zhao, Y., Siddiqui, A., Asano, J., McDonald, H., Pandoh, P., Dhalla, N., Prabhu, A.L., Ma, K. et al. (2007) Large-scale production of SAGE libraries from microdissected tissues, flow-sorted cells, and cell lines. Genome Res., 17, 108-116.
- 69. Al-Shahrour, F., Díaz-Uriarte, R. and Dopazo, J. (2004) FatiGO: a web tool for finding significant associations of gene ontology terms to groups of genes. Bioinformatics, 20, 578-580.
- 70. Brannan, C.I., Perkins, A.S., Vogel, K.S., Ratner, N., Nordlund, M.L., Reid, S.W., Buchberg, A.M., Jenkins, N.A., Parada, L.F. and Copeland, N.G. (1994) Targeted disruption of the neurofibromatosis type-1 gene leads to developmental abnormalities in heart and various neural crest-derived tissues. Genes Dev., 8, 1019-1029.
- 71. Choi-Miura, N.H., Otsuyama, K., Sano, Y., Saito, K., Takahashi, K. and Tomita, M. (2001) Hepatic injury-specific conversion of mouse plasma hyaluronan binding protein to the active hetero-dimer form. Biol. Pharm. Bull., 24, 892-896.
- 72. Duncan, S.A. (2000) Transcriptional regulation of liver development. Dev. Dyn., 219, 131-142.
- 73. Fleming, M.D., Campagna, D.R., Haslett, J.N., Trenor, C.C.III and Andrews, N.C. (2001) A mutation in a mitochondrial transmembrane protein is responsible for the pleiotropic hematological and skeletal phenotype of flexed-tail (f/f) mice. Genes Dev., 15, 652-657.
- 74. Fan, C.Y., Pan, J., Chu, R., Lee, D., Kluckman, K.D., Usuda, N., Singh,I., Yeldandi,A.V., Rao,M.S., Maeda,N. et al. (1996) Hepatocellular and hepatic peroxisomal alterations in mice with a disrupted peroxisomal fatty acyl-coenzyme A oxidase gene. J. Biol. Chem., 271, 24698-24710.
- 75. Hsiao, E.C., Koniaris, L.G., Zimmers-Koniaris, T., Sebald, S.M., Huynh, T.V. and Lee, S.J. (2000) Characterization of growthdifferentiation factor 15, a transforming growth factor beta superfamily member induced following liver injury. Mol. Cell. Biol., 20,
- 76. Jung, J., Zheng, M., Goldfarb, M. and Zaret, K.S. (1999) Initiation of mammalian liver development from endoderm by fibroblast growth factors. Science, 284, 1998-2003.
- 77. Suzuki, T., Kanai, Y., Hara, T., Sasaki, J., Sasaki, T., Kohara, M., Maehama, T., Taya, C., Shitara, H., Yonekawa, H. et al. (2006) Crucial role of the small GTPase ARF6 in hepatic cord formation during liver development. Mol. Cell. Biol., 26, 6149-6156.
- 78. Clotman, F., Jacquemin, P., Plumb-Rudewiez, N., Pierreux, C.E., Van der Smissen, P., Dietz, H.C., Courtoy, P.J., Rousseau, G.G. and Lemaigre, F.P. (2005) Control of liver cell fate decision by a gradient of TGF beta signaling modulated by onecut transcription factors. Genes Dev., 19, 1849-1854.
- 79. Kim, J.Y., Kim, H.J., Kim, K.T., Park, Y.Y., Seong, H.A., Park, K.C., Lee, I.K., Ha, H., Shong, M., Park, S.C. et al. (2004) Orphan nuclear receptor small heterodimer partner represses hepatocyte nuclear factor 3/Foxa transactivation via inhibition of its DNA binding. Mol. Endocrinol., 18, 2880-2894.
- 80. Qu,X., Lam,E., Doughman,Y.Q., Chen,Y., Chou,Y.T., Lam,M., Turakhia, M., Dunwoodie, S.L., Watanabe, M., Xu, B. et al. (2007)

- Cited2, a coactivator of HNF4alpha, is essential for liver development. EMBO J., 26, 4445-4456.
- 81. Shen, H., Huang, G., Hadi, M., Choy, P., Zhang, M., Minuk, G.Y., Chen, Y. and Gong, Y. (2003) Transforming growth factor-beta1 downregulation of Smad1 gene expression in rat hepatic stellate cells. Am. J. Physiol. Gastrointest. Liver Physiol., 285, G539-G546.
- 82. Zhao, R., Watt, A.J., Li, J., Luebke-Wheeler, J., Morrisey, E.E. and Duncan, S.A. (2005) GATA6 is essential for embryonic development of the liver but dispensable for early heart formation. Mol. Cell. Biol., 25, 2622-2631.
- 83. Gauthier, B.R., Schwitzgebel, V.M., Zaiko, M., Mamin, A., Ritz-Laser, B. and Philippe, J. (2002) Hepatic nuclear factor-3 (HNF-3 or Foxa2) regulates glucagon gene transcription by binding to the G1 and G2 promoter elements. Mol. Endocrinol., 16,
- 84. Legraverend, C., Eguchi, H., Strom, A., Lahuna, O., Mode, A., Tollet, P., Westin, S. and Gustafsson, J.A. (1994) Transactivation of the rat CYP2C13 gene promoter involves HNF-1, HNF-3, and members of the orphan receptor subfamily. Biochemistry, 33,
- 85. Sund, N.J., Ang, S.L., Sackett, S.D., Shen, W., Daigle, N., Magnuson, M.A. and Kaestner, K.H. (2000) Hepatocyte nuclear factor 3beta (Foxa2) is dispensable for maintaining the differentiated state of the adult hepatocyte. Mol. Cell. Biol., 20, 5175-5183.
- 86. Lee, C.S., Friedman, J.R., Fulmer, J.T. and Kaestner, K.H. (2005) The initiation of liver development is dependent on foxa transcription factors. Nature, 435, 944-947.
- 87. Zanotto, E., Shah, Z.H. and Jacobs, H.T. (2007) The bidirectional promoter of two genes for the mitochondrial translational apparatus in mouse is regulated by an array of CCAAT boxes interacting with the transcription factor NF-Y. Nucleic Acids Res., 35, 664-677.
- 88. Nguyen, T.T., Cho, K., Stratton, S.A. and Barton, M.C. (2005) Transcription factor interactions and chromatin modifications associated with p53-mediated, developmental repression of the alpha-fetoprotein gene. Mol. Cell. Biol., 25, 2147-2157.
- 89. Kypriotou, M., Fossard-Demoor, M., Chadjichristos, C., Ghayor, C., de Crombrugghe, B., Pujol, J.P. and Galera, P. (2003) SOX9 exerts a bifunctional effect on type II collagen gene (COL2A1) expression in chondrocytes depending on the differentiation state. DNA Cell Biol., **22**, 119–129.
- 90. Gauldie, J., Bonniaud, P., Sime, P., Ask, K. and Kolb, M. (2007) TGF-beta, Smad3 and the process of progressive fibrosis. Biochem. Soc. Trans., 35, 661-664.
- 91. Schadinger, S.E., Bucher, N.L., Schreiber, B.M. and Farmer, S.R. (2005) PPARgamma2 regulates lipogenesis and lipid accumulation in steatotic hepatocytes. Am. J. Physiol. Endocrinol. Metab., 288, E1195-E1205.
- 92. Zhao, C., Chen, W., Yang, L., Chen, L., Stimpson, S.A. and Diehl, A.M. (2006) PPARgamma agonists prevent TGFbeta1/ Smad3-signaling in human hepatic stellate cells. Biochem. Biophys. Res. Commun., 350, 385-391.
- 93. Wimmer, K., Vinatzer, U., Zwirn, P., Fonatsch, C. and Wieser, R. (1998) Comparative expression analysis of the antagonistic transcription factors EVI1 and MDS1-EVI1 in murine tissues and during in vitro hematopoietic differentiation. Biochem. Biophys. Res. Commun., 252, 691-696.
- 94. Samaras, S.E., Cissell, M.A., Gerrish, K., Wright, C.V., Gannon, M. and Stein, R. (2002) Conserved sequences in a tissue-specific regulatory region of the pdx-1 gene mediate transcription in pancreatic beta cells: Role for hepatocyte nuclear factor 3 beta and Pax6. Mol. Cell. Biol., 22, 4702-4713.