Profiling of promoter occupancy by PPARα in human hepatoma cells via ChIP-chip analysis

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ABSTRACT
The transcription factor peroxisome proliferator-activated receptor α (PPARα) is an important regulator of hepatic lipid metabolism. While PPARα is known to activate transcription of numerous genes, no comprehensive picture of PPARα binding to endogenous genes has yet been reported. To fill this gap, we performed Chromatin immunoprecipitation (ChIP)-chip in combination with transcriptional profiling on HepG2 human hepatoma cells treated with the PPARα agonist GW7647. We found that GW7647 increased PPARα binding to 4220 binding regions. GW7647-induced binding regions showed a bias around the transcription start site and most contained a predicted PPAR binding motif. Several genes known to be regulated by PPARα, such as ACOX1, SULT2A1, ACADL, CD36, IGFBP1 and G0S2, showed GW7647-induced PPARα binding to their promoter. A GW7647-induced PPARα-binding region was also assigned to SREBP-targets HMGCS1, HMGCR, FDFT1, SC4MOL, and LPIN1, expression of which was induced by GW7647, suggesting cross-talk between PPARα and SREBP signaling. Our data further demonstrate interaction between PPARα and STAT transcription factors in PPARα-mediated transcriptional repression, and suggest interaction between PPARα and TBP, and PPARα and C/EBPα in PPARα-mediated transcriptional activation. Overall, our analysis leads to important new insights into the mechanisms and impact of transcriptional regulation by PPARα in human liver and highlight the importance of cross-talk with other transcription factors.

INTRODUCTION
Peroxisome proliferator-activated receptors (PPARs) are ligand-activated transcription factors (TFs) that play an important role in the regulation of numerous biological processes, including lipid metabolism, adipocyte differentiation, cell proliferation and inflammation (1). Three different highly conserved PPAR isotypes have been identified: PPARα (NR1C1), PPARβ/δ (NR1C2) and PPARγ (NR1C3). The PPARα isotype has been shown to govern expression of numerous genes involved in fatty acid oxidation, ketogenesis, gluconeogenesis, cholesterol catabolism and lipoprotein metabolism (2,3). Additionally, PPARα has anti-inflammatory effects by suppressing pro-inflammatory genes (4,5). Consistent with its prominent function in lipid metabolism, PPARα is activated by various fatty acids and fatty acid derivatives, as well as by synthetic agonists such as fenofibrate, WY14,643 and GW7647 (6). Analogous to several other nuclear receptors, PPARs form heterodimers with retinoid X receptors (RXRs) (7), which occurs independently of ligand or DNA binding (8). PPARs bind to DNA by recognizing specific cis-acting PPAR responsive elements (PPREs) present in the regulatory regions of PPAR target genes. The consensus PPRE consists of a direct repeat of the hexameric sequence AGGTCA separated by one less conserved spacer nucleotide. PPARα was shown to bind to the 5′ motif of the PPRE, whereas RXR binds to the 3′ motif (9). Full activation of gene transcription by the DNA-bound PPAR–RXR complex is ultimately dependent on the formation of a larger transcription initiation protein complex via recruitment of a number of
co-activator proteins and RNA polymerase II (10). Additionally, PPARz has been shown to down-regulate gene expression by interfering with the activity of other TFs (11,12).

Recent studies using chromatin immunoprecipitation (ChIP) in combination with genomic tilling microarrays or sequencing (ChIP-chip or ChIP-Seq) have provided important new information on the requirements for DNA-binding by PPARs and other nuclear receptors. One of the most interesting findings was that genes that are activated by PPARγ show an enrichment of modules consisting of a PPRE-like motif together with a C/EBP binding element (13,14). It was found that knocking-down of either PPARγ or C/EBP reduced expression of several PPAR target genes, which was further reduced when these genes where knocked down simultaneously (13). A similar enrichment of a TF module was shown for the estrogen receptor (ER), which clustered together with forkhead, Oct1 and C/EBP motifs (15), as well as for the androgen receptor (AR) and the glucocorticoid receptor (GR) (16,17). These ChIP-chip studies thus reveal a complex interplay between nuclear receptors and other TFs. This cross-talk might be an important mechanism in gene regulation by nuclear receptors and may be responsible for transcriptional regulation of specific sets of genes.

While several studies have mapped the genomic binding regions of PPARγ, there are no reports available that describe the use of ChIP-chip or ChIP-Seq to investigate genomic binding of PPARz. Accordingly, in the present study we generated genome-wide maps of PPARz binding regions in the HepG2 human hepatoma cell line using ChIP combined with human promoter tiling arrays. We used HepG2 cells because they represent the most widely used cellular model for human liver cells, despite the modest effects of PPARz activation on gene expression. Our analysis was targeted towards the discovery of promoter sites showing increased PPARz binding in response to PPARz activation by ligand. To investigate the relation between promoter occupancy by PPARz and regulation of gene expression, results of ChIP-chip analysis were coupled to gene expression data.

MATERIALS AND METHODS

Cell culture

HepG2 cells were grown in phenol red-free Dulbecco’s modified medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 2 mM l-glutamine, 0.1 mg/ml streptomycin and 100 U/ml penicillin. Cells were split the day before experiments. Cells were kept at 37°C and 5% CO₂. The following day cells were treated with either 100 nM of the PPARz agonist GW7647 or control vehicle (DMSO). Cells used for ChIP-chip analysis were harvested after 2h of GW7647 treatment. Cells used for gene expression analysis were harvested after 6h of GW7647 treatment.

ChIP

ChIP was performed as described previously (18). Briefly, protein–DNA complexes were cross-linked in 1%

formaldehyde for 10 min at room temperature. Cross-linking was stopped by addition of 1 M glycine to a final concentration of 125 mM for 5 min. Cells were washed twice with PBS, scraped and collected by centrifugation. The cell pellet was dissolved in lysis buffer (1% SDS, 10 mM EDTA, 50 mM Tris–HCl, pH 8.0). Extracts were sonicated using the Bioruptor (Diagenode) at high power until DNA fragments of ~500–1000 bp were formed. Sonicated chromatin was diluted in five volumes of dilution buffer (0.01% SDS, 1.1% Triton X-100, 1.2 mM EDTA, 167 mM NaCl, 16.7 mM Tris–HCl, pH 8.0). The immuno-complexes were precipitated using antibodies against PPARz, STAT1, STAT3 or STAT6 or normal serum IgG (sc-9000, sc-592, sc-482, sc-981, sc-2027, Santa Cruz Biotechnologies). Precipitated complexes were reverse cross-linked and proteins digested with proteinase K (Fermentas) overnight at 65°C. DNA fragments were purified using the Affymetrix GeneChip Clean-up module. Purified DNA was used for qPCR and ChIP-chip analysis. ChIP experiments were run in quadruplicate.

ChIP-chip

To obtain sufficient DNA for hybridization, purified ChIP DNA was amplified and reamplified with the WGA (re)amplification kit (Sigma-Aldrich). Amplified DNA was fragmented with DNase I to ~50 bp fragments. The fragments were labeled with biotin according to Affymetrix instructions. Biotinylated DNA fragments were hybridized to Human Promoter 1.0R Arrays (Affymetrix). Arrays were washed and scanned according to instructions from manufacturer. Arrays were run in quadruplicate for both untreated and GW7647 treated cells.

Quantitative real-time PCR

Quantitative real-time PCR was performed on enriched regions of LPIN1, HMGR and AGPAT9. Equal amounts of amplified ChIP material was used and measurements were performed on the iCycler (BioRad) using platinum Taq polymerase (Invitrogen) and the double stranded DNA dye SYBR green. The following primers where used: LPIN1: sense primer 5’-ATTGGGAGGTGTGTTGTGGTATG-3’ and anti-sense primer 5’-ATAACAAATGCTGGCAAACG-3’. AGPAT9: sense primer 5’-CATCTAATACACAAACCAAGG-3’ and anti-sense primer 5’-AGCCAAACAAAGACTATTCG-3’. HMGR: sense primer 5’-AGCGTGATTGTGGGTCATATGG-3’ and anti-sense primer 5’-GTGTAAATGGCTCCGGTAC-3’. qPCRs were performed in duplicate and on all ChIP samples used for ChIP-Chip. Other primer sequences are available upon request.

Data analysis ChIP-chip

Affymetrix microarray CEL files were acquired and normalized using the Model-based analysis of tiling-arrays (MAT) algorithm (19). MAT was also used to identify PPARz binding regions induced by GW7647 treatment. Tiling-array probe intensities from ChIP performed on
HepG2 cells treated for 2 h with GW7647 were compared with probe intensities from vehicle-treated HepG2 cells. The analysis was performed with a MAT score of 2.4 or higher, which was based on MAT scores found for promoters of several known PPARα target genes. The human NCBIv36 (hg18) was used as a mapping file. In our analysis we used MaxGap of 400 bp and a bandwidth of 400 bp.

Transcriptomics
Treatment of HepG2 cells with GW7647 for gene expression analysis by transcriptomics was performed in triplicate. Total RNA was extracted from HepG2 cells with TRIzol reagent (Invitrogen) and subsequently purified using the SV Total RNA Isolation System (Promega). RNA quality was measured on an Agilent 2100 bioanalyzer (Agilent Technologies) using 6000 Nano Chips according to manufacturer’s instructions. RNA was judged as suitable for array hybridization only when samples showed intact bands corresponding to the 18S and 28S rRNA subunits, displayed no chromosomal peaks or RNA degradation products and had a RNA integrity number above 8.0. Five micrograms of RNA were used for one cycle cRNA synthesis (Affymetrix). Hybridization, washing and scanning of Affymetrix human genome 133 2.0 plus arrays was carried out according to standard Affymetrix protocols. Scans of the Affymetrix arrays were processed using packages from the R/Bioconductor project. Arrays were normalized with quantile normalization and expression levels of probe sets were calculated using the robust multichip average quantile normalization and expression levels of probe sets were calculated using the robust multichip average method. Differentially expressed probe sets were identified using Limma and genes were considered to be significantly changed when raw *p*-value <0.05 and fold-change >1.2. Genes regulated with an annotated ChIP-chip peak in its promoter region are listed in Supplementary Table S1.

Characterization of PPARα binding regions
For annotation of the ChIP peaks to adjacent genes, we used the Genomatix tool RegionMiner. The genes with the shortest distance to the ChIP enriched binding site were selected. For identification of new TF modules the software tool FrameWorker was used (20). GW7647-induced PPARα binding regions that were assigned to the top 25 differentially regulated genes (both up- and down-regulated) were analyzed. The minimal occurrence of a module in the binding regions studied was set at 20% and a maximal distance bandwidth variation between the motives was set at 75 bp. Another Genomatix tool, ModelInspector, was used to scan all the PPARα binding regions linked to GW7647-regulated genes. As a control, we scanned approximately all human promoter regions present in the Genomatix promoter database with sizes between 1 and 1.5 kb, which corresponds with the average size of the binding sites identified by ChIP-chip. We used a two-proportion *z*-test to analyze significant enrichment of modules. Samples with *p*-values values below 0.05 were considered significantly different.

Motif analysis
To investigate the presence of *de novo* DNA motifs, the top 25 PPARα binding regions linked to up- or down-regulated genes were loaded into the MEME tool (http://meme.sdsc.edu/meme4_1/cgi-bin/meme.cgi). The length for the output motif was varied between 12 and 20 bp. Enriched matrices of output motifs were compared to existing motif matrices available in both the TRANSFAC and the JASPAR database with the use of the STAMP tool (http://www.benoslab.pitt.edu/stamp/index.php).

Data release
The ChIP-chip and expression array data have been submitted to GEO and the data will be released upon publication.

RESULTS
Mapping of PPARα binding regions to adjacent transcripts
PPARα represents a ligand-induced TF that in the absence of ligand does not seem to act as transcriptional repressor, unlike several other nuclear receptors (21). For the analysis presented here we assumed that induction of PPARα target gene expression by PPARα ligand is associated with increased binding of PPARα to the DNA, which is supported by ample experimental data (22–27). Therefore, to identify the complete repertoire of hepatic PPARα target genes, we treated HepG2 cells with the synthetic ligand GW7647 for 2 h and performed ChIP-chip analysis using an antibody against PPARα, with vehicle-treated HepG2 cells serving as control. After amplification and fluorescent labeling, the immunoprecipitated chromatin templates were hybridized to Affymetrix human promoter tiling arrays. These arrays cover promoter regions of 7.5 kb upstream and 2.5 kb downstream of the transcription start site (TSS). Statistical analysis of four replicate experiments provided evidence for increased binding of PPARα to 4220 binding regions in response to treatment with GW7647.

We used the Genomatix tool RegionMiner to map the genes closest to the 4220 GW7647-induced PPARα binding regions in four genomic directions and selected the gene with the shortest distance to the binding region. The 4220 PPARα binding regions were linked to 3670 unique genes, of which 2875 were present on the expression arrays used for gene expression analysis. Analogous to other nuclear receptors, PPARα binding regions showed a bias towards the TSS (Figure 1A) (15,16, 28,29). To further analyze the location of PPARα binding regions, the online tool PinkThing (http://pinkthing.cmbi.ru.nl/cgi-bin/index50.pl) was used, which categorizes binding regions based on distance relative to the TSS (Figure 1B). We found that 46% of the binding regions were located within introns, which matches very well with a similar analysis done for PPARγ (14). Since we used promoter arrays, binding regions located to introns...
are in most cases present in the first intron. It is possible that some of these intronic binding regions actually surround the TSS of an alternative splice variant of a gene. Most other binding regions were identified upstream of the TSS with 26% within 5 kb and 11% between 5 and 25 kb. The more distal binding regions are not covered by promoter tiling arrays and therefore binding regions were not expected for the categories 3' far (5–25 kb) and distant (>25 kb). Hence, the 1 and 2% of binding regions located to the 3' far and distant categories, respectively, likely reflect misclassification by the PinkThing tool.

Several genes that are established targets of PPARα, such as SULT2A1, ACOX1, ACADL, CD36, IGFBP1 and G0S2 (18, 27, 30–34), showed GW7647-induced PPARα binding regions in the promoter (Figure 1C).
However, many other known PPARα target genes did not show any ligand-induced binding regions in their promoter. For these genes, PPARα is likely bound to a binding site not covered by the promoter array or, less likely, their PPRE may be located in repetitive sequences, which are excluded from Affymetrix human promoter tiling arrays using RepeatMasker. For example, the PPARα target gene ANGPTL4 carries a ligand-induced PPARα binding site in the third intron (24), which is out of the range of 2.5 kb downstream of the TSS covered by the array. As a result, no significant peak is present within the ANGPTL4 promoter (Figure 1C). Finally, it cannot be excluded that for some PPAR targets the degree of promoter occupancy by PPARα is not influenced by PPARα agonist and therefore no signals are detected, since our analysis concentrated on GW7647-induced PPARα binding regions.

Overlap between ChIP-chip and expression array

To investigate the relation between genes assigned to a GW7647-induced PPARα binding region and expression of that particular gene, we performed expression microarray analysis on HepG2 cells treated with GW7647 for 6 h. Confirming activation of PPARα, established PPARα targets CYP1A1, ADFP and TRIB3 were significantly induced by GW7647 (Figure 2A). Genes were considered significantly regulated if the mean fold change

Figure 2. Overlap between GW7647-induced PPARα binding and GW7647-induced changes in expression. (A) Significant induction of PPARα targets by GW7647 treatment. (B) Number of genes significantly altered upon GW7647 treatment as determined by microarray analysis using criteria: fold change >1.2 and q-value <0.05. (C) Overlap between genes assigned to GW7647-induced PPARα binding regions and genes altered after treatment with GW7647 as determined by transcriptomics. (D) Percentage of GW7647-induced PPARα binding regions linked to either up- or down-regulated genes that contain at least one V$PERO$ site, as determined using Genomatix. Similar analysis was done for all GW7647-induced PPARα binding regions as well as a control set of promoter regions in the Genomatix promoter database with similar size range as the binding regions identified by ChIP-chip (1000–1500 bp).
Motifs and module searches

To search for specific DNA motifs in GW7647-induced PPARα binding regions we performed a de novo motif search with MEME (http://meme.sdsc.edu/meme4_1/cgi-bin/meme.cgi). For this analysis we used the binding regions of the 25 most significantly up-regulated genes that were assigned to a GW7647-induced PPARα binding site (Figure 4). Identified motifs were compared to the two major motif databases TRANSFAC and JASPAR to search for similarities to existing TF binding motifs using STAMP (http://www.benoslab.pitt.edu/stamp/index.php). One of the identified motifs matched the PPARα–RXRα motif in the TRANSFAC database. While the classical AGGTCA motif of a PPRE was present, a clear DR1-type tandem repeat was not found. This is in line with the earlier findings indicating that the PPRE 5′ motif is less conserved than the 3′ motif (14,36), suggesting that the site we found is most likely a 3′ motif. A second motif had a similarity hit with the C/EBPβ motif in the JASPAR database, indicating that C/EBPβ motif is enriched in PPARα binding regions. Interestingly, the C/EBPβ motif was recently shown to be enriched in PPARγ binding regions and important for regulation of PPARγ target genes (13).

TFs often work in conjunction with other TFs to regulate DNA transcription. Accordingly, we examined whether other TF binding motifs are enriched together with the PPAR binding motif within the set of GW7647-induced PPARα binding regions linked to up- or down-regulated genes, using the Genomatix tool Frameworker. The PPAR matrix VSPERO was set and screening was performed for binding motifs of other TFs. Within the binding regions of the 25 most significantly up-regulated genes linked to a GW7647-induced PPARα binding region, we identified a highly significant ($p = 6.94 \times 10^{-8}$) module composed of the TATA binding protein (TBP, OSTBP) in combination with the PPAR binding motif VSPERO (Figure 5A). Within the binding regions of the 25 most significantly down-regulated genes linked to a GW7647-induced PPARα binding region, we identified a highly significant ($p = 1.04 \times 10^{-8}$) module composed of the binding motif for the TF signal transducers and activators of transcription (STAT) family (VSSTAT) together with VSPERO (Figure 5A). To further analyze for the presence of these specific modules, we used ModelInspector to scan all the PPARα binding regions linked to GW7647-regulated genes. As a control, we also scanned all human promoter areas present in the Genomatix promoter database with a size between 1 and 1.5 kb. As shown in Figure 5B, the module STAT-PERO is significantly enriched in the binding regions linked to genes that are down-regulated after 6 h GW7647 treatment compared to binding regions linked to genes up-regulated by GW7647 treatment and to all human promoter regions present in the Genomatix promoter database. In fact, the STAT-PERO module was found in eight out of 46 genes that were linked to a GW7647-induced PPARα binding region and were down-regulated by GW7647. Similarly, the TBP-PERO module is significantly enriched in binding regions linked to up-regulated genes compared to binding regions linked to down-regulated genes and all human promoter regions. These results suggest interaction between PPARα and STAT in PPARα-mediated transcriptional repression and interaction between PPARα and TBP in PPARα-mediated transcriptional activation.

Members of the STAT family bind to similar DNA sequences. Expression profiling indicated that expression of STAT3 was highest in HepG2 cells, followed by STAT1 and STAT6 (data not shown). Accordingly, we focused on
those three proteins to experimentally validate the suggested interaction between PPARα and STATs on the promoter of the aforementioned eight genes. HepG2 cells were treated with GW7647 for 2h and ChIP performed using antibodies against STAT3, STAT1 and STAT6 with vehicle-treated HepG2 cells serving as control. Precipitated chromatin was subsequently amplified using primers around the predicted STAT-PERO site. Consistent with interaction between PPARα and STATs, for seven out of eight genes STAT1 was released in response to GW7647 treatment (Figure 5C and Supplementary Figure S1). A similar picture emerged for STAT3. In contrast, STAT6 showed minor binding which generally was not altered by GW7647. These data show that PPARα activation and DNA binding causes the release of STAT3 and STAT1 from gene promoters concurrent with down-regulation of gene expression.
Biological clustering

Finally, within up-regulated genes that were assigned to GW7647-induced PPARα binding regions we analyzed for functional biological clusters using DAVID (http://niaid.abcc.ncifcrf.gov/home.jsp). We found significant over-representation of genes in the biological cluster of sterol and lipid biosynthetic process (Table 1), in line with binding of PPARα to the putative promoter of several SREBP targets. Within down-regulated genes that were assigned to GW7647-induced PPARα binding regions we found significant over-representation of genes in the biological cluster of humoral and innate immune response, which is consistent with the known suppressive effect of PPARα on inflammation (Table 1).

DISCUSSION

It is well established that PPARα is a major transcriptional regulator of fatty acid metabolism in liver. Numerous genes involved in fatty acid oxidation and other metabolic processes have been identified as direct target genes of PPARα in mouse, characterized by the presence of a functional PPRE (2,33). In the past few years, genomic binding regions of several nuclear receptors including PPARγ have been mapped (13,14). However, no reports are available that were aimed at mapping the binding regions for PPARα. In our study we used ChIP-chip to investigate ligand-induced PPARα binding to genomic regions within HepG2 human hepatoma cells.

A total number of 4220 ligand-induced PPARα binding regions were identified, which were assigned to 3670 unique genes. Although this number may appear exceptionally high, the threshold for inclusion was set based on inspection of a number of known PPARα binding regions. Our assignment of PPARα binding regions relative to the nearest gene showed a distance-to-TSS distribution centered around the TSS. Categorizing these distances revealed that the majority of the binding regions are located within an intron or in the 5' proximal region. These findings are in line with various studies showing that TF binding regions are normally distributed around the TSS (29,37–39). The preponderance of binding regions around the TSS may also reflect DNA looping allowing nuclear receptors bound to distal sites to contact the basal transcription machinery. Previous genome-wide analysis of ERα and PPARγ binding sites indicate that a large fraction of the binding regions are located distal from the TSS (13,15). Since we used promoter tiling arrays, our analysis was unable to detect binding to distant binding regions. Some of these distal elements have been shown to indeed function as functional PPREs (24,40–42). However, many distant binding regions are likely misclassified and are actually located in proximal promoters of new transcripts (43,44). Furthermore, many regions identified by ChIP-based methods do not show any effect on the expression of the gene closest to that region. These issues raise questions about the functional relevance of many of these binding regions (45).

We compared our ChIP-chip data with mRNA expression data collected by expression microarray and found...
that ~18% of the genes induced by ligand were linked to a GW7647-induced PPARα binding region. The other 82% may not represent direct target genes, may bind PPARα equally in absence and presence of ligand, may bind PPARα via a site located outside the range covered by the array, or the actual PPARα binding region was linked to a different gene.

Importantly, the overwhelming majority of PPARα binding regions were linked to genes that were not significantly altered upon GW7647 treatment, even after more prolonged treatment (data not shown). It is well recognized that the majority of binding regions found by ChIP-chip or ChIP-Seq experiments do not have any effect on the genes they were linked to (45). This could be partially due to miss-annotation, which is inevitable when assigning to the nearest gene. However, since the number of GW7647-induced binding regions far exceeds the number of GW7647-regulated genes many regions must bind PPARα without any impact on gene regulation. These findings are consistent with the ‘scanning model’ proposed for PPARs, which states that PPARs scan the genome and transiently bind to PPRE-like sequences.

Figure 5. Enrichment of TF modules in PPARα binding regions. (A) The binding regions of the 25 most significantly up-regulated genes assigned to GW7647-induced PPARα binding regions were analyzed for TF modules using the Genomatix tool Frameworker. Two modules were identified: TBP-PERO and STAT-PERO in the binding regions linked to up- or down-regulated genes, respectively. (B) The modules TBP-PERO and STAT-PERO were scanned for the relative presence in all GW7647-induced PPARα binding regions located near transcriptionally regulated as well as all human promoter regions present in the Genomatix database. A two-proportion z-test was used to analyze significant enrichment of modules. p-values values below 0.05 were considered significantly different. (C) Loss of DNA binding by STAT3 and STAT1 upon PPARα activation. HepG2 cells were treated with GW7647 for 2h and ChIP performed using antibodies against STAT3, STAT1 and STAT6 with vehicle-treated HepG2 cells serving as control. Precipitated chromatin was subsequently amplified using primers around the predicted STAT-PERO site found in four genes that were linked to a GW7647-induced PPARα binding region and were down-regulated by GW7647. Significant differences are indicated with an asterisk (Student’s t-test, p<0.05).
without inducing any transcription. According to this model, PPARs only start transcription upon binding to a bona fide PPRE from a PPRE-like sequence is not well understood. Recent ChIP-chip studies on several TFs revealed that clusters of different TF binding elements are enriched in proximity of the binding region for the TF under study (13,15,16,46–48). We searched for these types of clusters in PPARx binding regions that were linked to differentially regulated genes and found several modules of TF binding elements in combination with a PPRE. In PPARx binding regions linked to up-regulated genes we found enrichment for a module composed of the binding sequence for TBP together with that of PPARx. Since TBP is an important component of the basal transcriptional machinery, this finding suggests binding of PPARx adjacent to the core promoter. A second interesting module we found was a combination of a PPARx binding sequence and a STAT binding sequence, which was strongly enriched in PPARx binding regions linked to genes that were down-regulated by PPARx activation. STAT TFs function downstream in the signaling pathway of a large number of cytokines, growth factors and hormones (49). Follow-up analysis by normal ChIP showed that binding of STAT3 and also STAT1 to these genes was reduced upon GW7647 treatment. Our data suggest that loss of STAT binding is dependent on binding of PPARx adjacent to the STAT binding site. Inhibition of STAT-dependent transcriptional activity and DNA binding has been previously documented for PPARγ and PPARβ/σ (50–54), and may also partially account for the anti-inflammatory action of PPARx activation (17). Overall, down-regulation of gene expression by PPARx activation in HepG2 cells may be partially mediated by interfering with binding of STAT3 and STAT1 to the DNA.

Using Genomatix we found that at least one VSPERO site was present in 81% of the PPARx binding regions linked to the significantly up-regulated genes. A similarly high percentage was found for all the GW7647-induced PPARx binding regions resulting from the ChIP-chip analysis. While this result would suggest that the majority of PPAR binding detected by our ChIP-chip analysis conforms to the general paradigm of PPRE-dependent DNA binding, it should be emphasized that VSPERO sites are found at relatively high frequency throughout the genome and may have limited specificity. As ChIP is capable of detecting any type of binding of PPARx to DNA, which includes indirect binding via other TFs, it is difficult to provide a good estimate of the relative importance of PPRE-dependent and -independent binding to DNA. Binding to DNA of TFs in the absence of a consensus motif has been a common observation in recent ChIP- and ChIP-seq studies. Several explanation may account for this apparent discrepancy as elaborated by Farnham (45), including binding at a distal site that contains a consensus motif and looping to the site in question through protein–protein interactions; ‘piggy-back’ binding that is mediated by protein–protein interactions with a second factor and that does not involve the DNA binding domain of the first factor; or assisted binding to a site that is similar to the consensus site, which is enhanced by protein–protein interaction with another site specific DNA binding factor or with a specifically modified histone.

Within the set of GW7647-induced PPARx binding regions linked to up-regulated genes, no established direct PPARx targets were present. One problem is that HepG2 cells, despite their broad use, poorly reflect gene regulation by PPARx in other cultured cells such as rat FAO hepatoma cells (55) and primary human hepatocytes. Indeed, we found remarkably little overlap in gene regulation by PPARx agonist between HepG2 cells and primary human hepatocytes (van der Meer et al., manuscript in preparation). Unfortunately, ChIP-chip analysis in primary human hepatocytes is practically unfeasible. Clearly, an ideal system to study PPARx-dependent gene regulation in human is lacking.

The TFs SREBP-1 and SREBP-2 are important regulators of hepatic lipid and cholesterol synthesis (56). Previously, it was shown that PPARx is involved in the normal circadian regulation of target genes of SREBPs, including HMGCR (57). Furthermore, synthetic PPARx agonists were found to induce expression of SREBP targets in liver, which was completely abolished in SREBP-1−/− mice and PPARx−/− mice (58,59). The effect of PPARx agonists on SREBP targets was attributed to increased activation of SREBP-1c via enhanced proteolytic cleavage, and was not mediated by changes in SREBP-1 mRNA (58). In the present article, we find induction of expression of several SREBP targets by PPARx activation in human hepatoma HepG2 cells. Furthermore, in these cells PPARx agonist stimulated binding of PPARx to the putative promoter of SREBP targets HMGCS1, HMGCR, FDFT1, SC4MOL and LPIN1 genes. These data suggest important cross-talk between PPARx and SREBP signaling. The exact nature of the cross-talk requires further investigation but one possibility is that PPARx is recruited to promoters of SREBP targets via direct physical interaction with SREBP. It could be envisioned that through this interaction PPARx may promote SREBP activity, perhaps by assisting with recruitment of transcriptional co-activators. In addition to the above-mentioned genes, a PPARx binding region was assigned to the AGPAT9 gene, which can be suspected to be a SREBP target as well. LPIN1 and AGPAT9 encode enzymes that catalyze the second and

| Biological process                                    | p-value |
|-------------------------------------------------------|---------|
| Up-regulated genes                                    |         |
| Sterol biosynthetic process                           | 1.1E-4  |
| Apoptosis                                             | 1.9E-2  |
| Developmental process                                 | 2.2E-2  |
| Lipid biosynthetic process                            | 4.0E-2  |
| Down regulated genes                                  |         |
| γ-Hexachlorocyclohexane                               | 1.8E-3  |
| Humoral immune response                               | 1.2E-2  |

Up-regulated genes assigned to GW7647-induced PPARx binding regions were analyzed for functional biological clusters using DAVID.

Table 1. Enrichment of biological processes in GW7647-regulated genes assigned to a PPARx binding region
third step in the triacylglycerol synthesis pathway. So far, several genes involved in either fatty acid or triacylglycerol synthesis have been identified as direct PPARα target genes including Δ5 and Δ6 desaturases and malonyl enzyme [summarized in refs (2,3)]. Indeed, the involvement of PPARα in lipogenesis appears to be much more extensive than previously understood (60). Although this notion is seemingly at odds with increased hepatic TG observed upon PPARα deletion, regulation of fatty acid metabolism by PPARα is probably more subtle than generally envisioned. According to data presented here, PPARα may also indirectly impact lipid biosynthesis via cross-talk with SREBP.

In aggregate, it can be concluded that ChIP-chip represents a powerful tool to investigate whole genome binding of PPARα. Our data indicate that PPARα agonists trigger the binding of PPARα to a large number of genomic sites and provide novel insights into the mechanisms of PPARα-dependent transcriptional regulation.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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