Glucocorticoids (GCs) are commonly prescribed drugs, but their anti-inflammatory benefits are mitigated by metabolic side effects. Their transcriptional effects, including tissue-specific gene activation and repression, are mediated by the glucocorticoid receptor (GR), which is known to bind as a homodimer to a palindromic DNA sequence. Using ChIP-seq in mouse liver under endogenous corticosterone exposure, we report here that monomeric GR interaction with a half-site motif is more prevalent than homodimer binding. Monomers colocalize with lineage-determining transcription factors in both liver and primary macrophages, and the GR half-site motif drives transcription, suggesting that monomeric binding is fundamental to GR’s tissue-specific functions. In response to exogenous GC in vivo, GR dimers assemble on chromatin near ligand-activated genes, concomitant with monomer evacuation of sites near repressed genes. Thus, pharmacological GCs mediate gene expression by favoring GR homodimer occupancy at classic palindromic sites at the expense of monomeric binding. The findings have important implications for improving therapies that target GR.

[Supplemental material is available for this article.]

Glucocorticoids (GCs) are cholesterol-derived steroid hormones that control metabolic and homeostatic processes essential for mammals. Natural GCs and their synthetic analogs are among the most widely prescribed drugs in the world because of their anti-inflammatory and immunosuppressive properties that are important for the treatment of rheumatoid arthritis, cerebral edema, allergic reactions, and asthma, and they are utilized to prevent organ transplant rejection and graft-versus-host disease (Ito et al. 2006a,b; Kirwan and Power 2007). Unfortunately, the development of major metabolic side effects, including diabetes, obesity, hypertension, osteoporosis, and muscle atrophy, can cause treatment to end (Rosen and Miner 2005). Both natural and synthetic GCs bind and activate GR (NR3C1), a nuclear receptor transcription factor (TF) ubiquitously expressed in the body (Heitzer et al. 2007; Evans and Mangelsdorf 2014). Upon ligand binding, GR translocates from the cytoplasm to the nucleus where it can either activate or repress the expression of target genes in a cell-type-specific manner. GR can regulate transcription by binding as a homodimer to its palindromic recognition sequence, termed a GC response element (GRE), yet the molecular mechanisms leading to transcriptional activation versus repression are unclear. Better understanding of this is the focus of intense effort, as the clinical benefits of GC treatment are thought to be due to the transcriptional repression of cytokines and chemokines in immune cells, whereas the unwanted side effects may be linked to the activation of metabolic genes in other tissues (Beck et al. 2009).

A popular model proposes that DNA binding of the GR homodimer transactivates expression, whereas protein–protein interactions tethering monomeric GR to other DNA-bound TFs, such as NFKB1 or JUN-FOS heterodimers (AP-1), mediate transrepression (Glass and Saijo 2010). Early support came from studies of GRdim mice initially developed to separate transactivation and transrepression functions. These mice carry an amino acid substitution in the D-loop of the DNA-binding domain of GR (A465T), which was designed to reduce GR homodimer formation and DNA
binding (Reichardt et al. 1998). In agreement with the transactivation mechanism, GRdim mice show a reduced ability to activate transcription in the liver in response to exogenous ligands (Frijters et al. 2010). However, later studies were inconsistent with expectations for transrepression by revealing that GRdim mice exhibit a diminished response to GC treatment in inflammatory paradigms such as TNF-induced inflammation (Vandevyver et al. 2012), LPS- and CLP-induced sepsis (Kleiman et al. 2012; Silverman et al. 2013), antigen-induced rheumatoid arthritis (Bascant et al. 2011, 2012), allergic contact dermatitis (Kleiman and Tuckermann 2007), and experimental autoimmune encephalomyelitis, a mouse model for multiple sclerosis (Schweingruber et al. 2014). This was in part due to the inability of GRdim to induce GR-dependent anti-inflammatory genes (Vandevyver et al. 2012). Although its DNA-binding properties are compromised relative to GR (Gebhardt et al. 2013; Watson et al. 2013), the inability of GRdim to dimerize and bind DNA has been challenged by in vitro studies (Jewell et al. 2012; Presman et al. 2014; Sedwick 2014). Complicating matters further, recent cis-regomic analyses in primary macrophages (Uhlenhaut et al. 2013) and the liver (Grøntved et al. 2013) have revealed similar enrichment of GR at recognition sequences near both ligand-activated and ligand-repressed genes, suggesting factors other than GR occupancy as important regulatory determinants. As a whole, these studies highlight the need for improved understanding of GR binding to native chromatin and how it pertains to transactivation and transrepression.

To this end, and to gain insight into the impaired hepatic gene expression and anti-inflammatory responses of GRdim mice, we compared chromatin immunoprecipitation (ChIP) with deep sequencing (ChIP-seq). ChIP with lambda exonuclease digestion and sequencing (ChIP-exo), and transcriptomic analyses in liver tissue isolated from wild-type (WT) and GRdim mice. Our study reveals extensive genomic occupancy of monomeric and dimeric GR that is specified by DNA motifs and colocalized TEs. Exogenous GC treatment of mice uncovers a mechanism whereby GR dimers transactivate at the expense of monomers, resulting in both increased and decreased GC-mediated gene expression.

Figure 1. GR, but not GRdim, occupies the canonical palindromic motif as a dimer in liver. (A) Top-ranked de novo motifs from HOMER for the GR cistromes from WT and GRdim mice. See Supplemental Material for a comprehensive list of motifs. (B) Scatter plots comparing sequence tags from 14,940 GR ChIP-seq peaks with at least two reads per million (RPM) in any condition in livers isolated from WT and GRdim mice killed at either 6 a.m. (top) or 6 p.m. (bottom). Blue and red highlight WT-selective and common sites, respectively. (C) GR binding upstream of the tyrosine aminotransferase (Tat) gene. The 5′ ends of forward- and reverse-stranded sequence tags are indicated in red and blue, respectively, for the ChIP-exo tracks. Tracks are RPM normalized. (D) Distance distribution for opposite-stranded peaks with at least 0.2 RPM from GR ChIP-exo in liver isolated at 6 a.m. from WT and GRdim mice is shown for WT-selective sites. The number of peak pairs and prominent peak distances are indicated. Schematic of opposite-stranded peaks is shown at top. (E) GR ChIP-exo for WT-selective sites in liver isolated at 6 a.m. MEME top-ranked de novo sequence with a hit count of at least 5% is shown at the top. See Supplemental Material for a full list of motifs. Average profiles (midline) and density heatmaps (bottom) of the raw sequence tags are shown for both mouse models. Red and blue indicate the 5′ ends of the forward- and reverse-stranded tags, respectively.
Results

GR, but not GR<sub>dim</sub>, occupies the palindromic motif as a dimer at a minority of genomic sites in mouse liver

To compare genomic occupancy by GR and GR<sub>dim</sub> under physiological conditions, we performed ChIP-seq in liver, where GR regulates diverse processes ranging from hepatic neonatal growth to glucose and lipid homeostasis (Rose and Herzig 2013). Liver tissue was isolated from WT and GR<sub>dim</sub> mice during the morning and evening to control for potential circadian effects of serum corticosterone levels superimposed over an ultradian rhythm, stress effects, and local GC production by hydroxysteroid 11-beta dehydrogenase 1 (Conway-Campbell et al. 2012). Biological replicates revealed highly correlated occupancy for GR at 6 a.m. and 6 p.m. in either WT or GR<sub>dim</sub> mice, with stronger occupancy at 6 p.m. (Supplemental Fig. 1). De novo motif analyses revealed enrichment for different GR motifs (Fig. 1A), and comparison of GR binding sites in WT versus GR<sub>dim</sub> mice revealed a bifurcated distribution (Fig. 1B), suggesting that GR and GR<sub>dim</sub> occupy two classes of sites categorized by relative binding strength. We identified 2110 sites that preferentially bind GR versus GR<sub>dim</sub>, termed WT-selective, and a major class of 11,108 common sites that are occupied similarly by GR and GR<sub>dim</sub> (Supplemental Fig. 2). Nearly indistinguishable behavior was observed for the WT-selective and common sites at 6 a.m. and 6 p.m. (Supplemental Fig. 3). These data demonstrate that GR<sub>dim</sub> binds the genome in vivo, and the absence of substantial GR<sub>dim</sub>-specific peaks indicates that it occupies a subset of GR binding sites.

To define the occupancy properties of GR and GR<sub>dim</sub> in more detail, we performed ChIP-exo using biological replicates from WT and GR<sub>dim</sub> mice to map sites at a higher resolution than afforded by ChIP-seq. The technique uses lambda exonuclease to trim DNA to the point where a crosslinked TF blocks further enzymatic digestion sites in WT versus GR<sub>dim</sub> mice revealed a bifurcated distribution (Fig. 1B), suggesting that GR and GR<sub>dim</sub> occupy two classes of sites categorized by relative binding strength. We identified 2110 sites that preferentially bind GR versus GR<sub>dim</sub>, termed WT-selective, and a major class of 11,108 common sites that are occupied similarly by GR and GR<sub>dim</sub> (Supplemental Fig. 2). Nearly indistinguishable behavior was observed for the WT-selective and common sites at 6 a.m. and 6 p.m. (Supplemental Fig. 3). These data demonstrate that GR<sub>dim</sub> binds the genome in vivo, and the absence of substantial GR<sub>dim</sub>-specific peaks indicates that it occupies a subset of GR binding sites.

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GR and GR<sub>dim</sub> monomers occupy the liver genome at half-site motifs and tethered sites

Remarkably, the sites commonly bound by GR and GR<sub>dim</sub> comprise a major subset of the GR liver cisotrome yet lack association with the palindromic motif. ChIP-exo at these sites identified peak pairs separated by 11 and 25 bp that emerged from the background in both WT and GR<sub>dim</sub> mice at 6 a.m. (Fig. 2A) and 6 p.m. (Supplemental Fig. 5A). De novo examination of peak pairs with
GR and GRdim occupy the liver genome at tethered sites. (A) MEME de novo sequences from 6 a.m. common site peak pairs separated by 20–30 bp and with a hit count of at least 5%. See Supplemental Material for a comprehensive list of motifs. (B) GR ChIP-exo at 6 a.m. common sites, with average profiles and density heatmaps for the ONECUT1 (left) and FOXA (right) motifs shown for both mouse models. GR common sites cobound by ONECUT1 or FOXA2 were interrogated. Red and blue indicate the 5' ends of the forward- and reverse-stranded tags, respectively. (C) Distribution of the GR half-site motif relative to neighboring motifs at common sites cobound by ONECUT1 (top) or FOXA2 (bottom). Results for ONECUT1 and FOXA2 liver sites without GR are shown for comparison. (D) Half-site-facilitated tethering. The GR half-site motif is represented in orange. X indicates formaldehyde crosslinking between proteins or protein–DNA. Two formaldehyde crosslinking events between GR and a DNA-crosslinked TF are necessary to detect sites where GR appears bound to noncanonical motifs.

short spacing (5–15 bp) identified the GR half-site and FOXA motifs as the only sequences present in at least 5% of the sites (Fig. 2B; Supplemental Fig. 5B). Consistent with the low frequency of short-spaced peak pairs, a minority of the common sites has peak pairs flanking the GR half-site motif at a fixed distance (Fig. 2C; Supplemental Fig. 5C, D). This could suggest that GR and GRdim occupy most of the common sites independently of the half-site motif. However, it is also possible that nuclear receptor monomers may not perform well in the ChIP-exo assay. In agreement with this, reexamination of a ChIP-seq study with the estrogen receptor (Serandour et al. 2013) revealed tighter peak pairs flanking the full-versus the half-site motifs (Supplemental Fig. 6). Moreover, in support of sequence-specific binding, the GR half-site motif is centrally enriched at ChIP-seq peaks from common sites (Fig. 2D), and its frequency within a peak scales with GR-occupancy strength (Fig. 2E). As a whole, the data indicate that GR and GRdim occupy the genome through sequence-specific interaction with the half-site motif, but experimental limitations preclude a direct assessment of its prevalence.

Examination of peak pairs with 20–to-30-bp spacing revealed centrally enriched ONECUT1, FOXA, and GR-half motifs at both 6 a.m. (Fig. 3A) and 6 p.m. (Supplemental Fig. 5B) for the common sites. Presence of the ONECUT1 and FOXA motifs raised the possibility of formaldehyde crosslinking between GR or GRdim and DNA-crosslinked ONECUT1 or a FOXA factor(s). To address this, we used published ChIP-seq data sets for ONECUT1 (Laudadio et al. 2012) and FOXA2 (Li et al. 2012) in liver to examine the GR ChIP-exo reads at common sites that are also bound by ONECUT1 or FOXA2. Opposite-stranded peaks flank the ONECUT1 motif with the same spacing found by ONECUT1 ChIP-exo (Wang et al. 2014), while an asymmetric profile of positioned peaks observed previously for FOXA2 ChIP-exo (Serandour et al. 2013) is present at the FOXA motif (Fig. 3B; Supplemental Fig. 5D). Their signal strength is low, most likely reflecting transient protein–protein interactions, yet their tight positioning indicates occupancy. Intriguingly, GR half-site motifs are located nearby the ONECUT1 and FOXA motifs (Fig. 3C), suggesting that GR recruitment is mediated and/or stabilized through sequence-specific interaction with the DNA. DNA crosslinking of both GR and a neighboring TF is unlikely to explain these results because this would preclude formation of a peak pair of uniform length with enrichment for a centered motif. It is interesting that the motifs for HNF4A and CEBP proteins are highly enriched by ChIP-seq but not by ChIP-exo. GR half-site motifs are located nearby the HNF4A and CEBP motifs at cobound common sites (Supplemental Fig. 7A), but positioned peak pairs are not prominent (Supplemental Fig. 7B). These data could reveal selectivity for GR–TF interactions in liver, such that GR preferentially associates with FOXA and ONECUT1 versus HNF4A and CEBP TFs. However, it is formally possible that GR interacts with each of these factors but that HNF4A and CEBP TFs do not produce robust ChIP-seq profiles. The general applicability of ChIP-exo remains to be determined given that only a handful of studies on a small number of TFs have been published to date. As a whole, the data indicate that GR and GRdim monomers occupy the liver genome via sequence-specific binding to the GR half-site motif and through interaction with DNA-bound TFs. The latter, termed half-site-facilitated tethering, is mediated and/or stabilized by binding to the half site (Fig. 3D). Half-site-facilitated tethering may occur generally, as examination of previously described genomic sites in pituitary-derived AtT-20 cells with proposed tethering of GR to STAT3 (Langlais et al. 2012) shows enrichment for the GR half-site motif (Supplemental Fig. 8).

GR monomers bind near lineage-determining TFs in liver

GR ChIP-seq and ChIP-exo from WT and GRdim mice implicate functions for hepatic lineage TFs, prompting us to investigate whether GR dimers and monomers differentially colocalize with them. De novo motif analyses revealed that the dimeric (WT-selective) ChIP-seq peaks are most enriched for the palindromic GR motif, while the monomeric (common) sites contain the recognition sequences for liver TFs along with the GR half site (Fig. 4A). When directly compared to each other as foreground and background data sets, the dimeric sites showed strong enrichment for only the palindromic GR motif, while the monomeric sites were modestly enriched for the HNF4A motif as the top-ranked sequence (see Supplemental Material). This indicates that the GR-palindromic motif distinguishes dimeric from monomeric sites and implies that liver TFs colocalize with monomeric sites more
Monomeric GR colocalizes with lineage-determining TFs in liver. (A) HOMER de novo motif analyses for the dimeric and monomeric GR binding sites from liver ChIP-seq. The four top-ranked lineage TF motifs are shown relative to the top-ranked GR sequence. See Supplemental Material for a comprehensive list of motifs. (B) Box plots interrogating the co-occupancy of liver TFs at GR binding sites. (C) Distribution plots showing relative to colocalized HNF4A, CEBPB, ONECUT1, and/or FOXA2. All TF combinations were examined.

Figure 4. Monomeric GR colocalizes with lineage-determining TFs in liver. (A) HOMER de novo motif analyses for the dimeric and monomeric GR binding sites from liver ChIP-seq. The four top-ranked lineage TF motifs are shown relative to the top-ranked GR sequence. See Supplemental Material for a comprehensive list of motifs. (B) Box plots interrogating the co-occupancy of liver TFs at GR binding sites. (C) Distribution plots showing relative to colocalized HNF4A, CEBPB, ONECUT1, and/or FOXA2. All TF combinations were examined.

than dimeric. Consistent with the latter, interrogation of published ChIP-seq data sets revealed stronger co-occupancy of the hepatic TFs HNF4A, FOXA2, CEBPB, and ONECUT1 at monomeric versus dimeric sites (Fig. 4B) and that the fraction of monomeric, but not dimeric, sites increases along with the number of colocalizing TFs (Fig. 4C). Collectively, the data suggest that monomeric versus dimeric occupancy has a greater dependency on liver-determining TFs.

GRdim partitions the GR cistrome similarly in primary macrophages and liver

To investigate the general nature of our findings, we performed GR ChIP-seq in primary bone marrow–derived macrophages isolated from WT and GRdim mice and activated with lipopolysaccharide (LPS). Sites commonly bound by both GR and GRdim make up the majority of the GR cistrome and are enriched for myeloid-specific and inflammatory-acting TF motifs in addition to the GR half site, whereas WT-selective sites are most enriched for the palindromic GR motif (Fig. 5A; Supplemental Fig. 9). Direct comparison to each other as foreground and background data sets showed modest enrichment for the CEBP motif at common sites and strong enrichment for only the palindromic GR motif at WT-selective sites (see Supplemental Material). In addition, examination of published ChIP-seq data sets showed that the co-occupancy of macrophage TFs with GR is stronger at common versus WT-selective sites (Fig. 5B). Together, these findings indicate similar properties for GR binding sites in macrophages and liver. To explore a transcriptional function for monomeric sites, we placed several WT-selective and common GR binding sites into luciferase reporters and assayed their activity in response to cotransfection of GR or GRdim (Fig. 5C). While GR drives more transcription than GRdim at the WT-selective sites, GRdim compares favorably at the common regions, agreeing with earlier studies demonstrating in vitro activity for GRdim (Adams et al. 2003; Meijising et al. 2009; Jewell et al. 2012). Mutation of the GR half-site sequence within the common regions destroys GR and GRdim activity, demonstrating that transcriptional activation by monomeric GR requires a half-site motif. To test this in a more physiological setting, we compared the expression of genes regulated by a GR ligand in LPS-stimulated WT macrophages to that in GRdim macrophages. As shown in Figure 5D, ligand has a reduced ability to modulate induced and repressed genes in GRdim macrophages, consistent with earlier findings in liver (Frijters et al. 2010). Collectively, the data suggest comparable functions for GR monomers and dimers in primary macrophages and liver.

Exogenous GC redistributes GR from monomeric to dimeric sites at ligand-regulated genes

To explore potential differences between GR monomers and dimers at pharmacological exposure to GC, we examined GR occupancy in liver isolated at 6 a.m. from WT and GRdim mice treated for 24 h with prednisolone, a synthetic GC used to treat a wide range of inflammatory and auto-immune conditions (Czock et al. 2005). Most GR binding sites were unaffected by prednisolone (Supplemental Fig. 10A), yet occupancy was increased at 889 sites and decreased at another 626 sites in WT mice, with minimal changes occurring at these sites in GRdim mice (Fig. 6A). Encouragingly, the top-ranked gene ontology (GO) categories associated with the gained sites involve glucose metabolism (Fig. 6A; Supplemental Fig. 10B), consistent with GR function in liver (Rose and Herzig 2013). Gained sites are strongly enriched for the GR palindromic motif and a robust dimer signature in WT but not GRdim mice (Fig. 6B; Supplemental Fig. 10C), indicating that the exogenous ligand primarily stimulates dimeric occupancy. In addition to GR, CEBPB is also recruited to the gained sites (Fig. 6C), agreeing with earlier work demonstrating collaborative binding interactions between GR and CEBPB (Grantved et al. 2013) and indicating that the genomic occupancy of multiple TFs can be indirectly affected by ligand. Lost sites, in contrast, are highly enriched for the GR half motif. Indeed, modest association with the STAT motif agrees with our finding of GR half-site motifs near sites of interactions between GR and STAT TFs in published data sets from liver (Tronche et al. 2004) and AtT-20 cells (Langlais et al. 2012), supporting the model of half-site-facilitated tethering.

We performed ChIP-seq for RNA polymerase II (RNAPII), which is recruited to active enhancers on a global scale (Szutorisz et al. 2005; Koch et al. 2008; Kim et al. 2010), to explore a potential transcriptional function for the prednisolone-regulated sites. A direct correlation was found between occupancy changes to RNAPII and GR in response to prednisolone. Increased and decreased RNAPII associates with increased and decreased GR, respectively (Fig. 6D; Supplemental Fig. 10D), suggesting that GR occupancy
modulates the activity of these regions. To investigate their association with gene expression, we mapped gained or lost sites to prednisolone-regulated genes (Frijters et al. 2010). Figure 6E shows that prednisolone-induced genes were enriched more than sixfold for gained versus lost sites, suggesting that induced occupancy by GR dimers drives activated transcription. Moreover, dimer sites without prednisolone-regulated occupancy were also colocalized with increased RNAPII (Fig. 6D) and enriched near induced genes (Fig. 6E), suggesting multiple mechanisms mediate prednisolone’s effects. Repressed genes had a greater tendency to harbor lost sites, suggesting that prednisolone stimulates GR dimers to activate particular genes at the expense of others that lose monomers (Supplemental Fig. 10E). GRdim redistribution is diminished relative to GR, likely due to its inability to form dimers on chromatin, and provides a rationale for the reduced ability of exogenous GC to regulate liver genes in GRdim versus WT mice (Frijters et al. 2010).

Discussion

GR has been a focus of extensive study for decades, yet how it activates some genes while repressing others remains an open question. The two main mechanisms of transactivation and transrepression propose that the monomeric/dimeric state of the receptor determines its transcriptional function such that GR dimers activate through binding of inverted repeats, whereas monomers repress by tethering to DNA-bound TFs. Our data reveal a strong association between dimeric GR occupancy and activated gene expression on a genome-wide scale in both liver tissue and primary macrophages. However, the discovery of monomeric occupancy near transcriptionally active genes, and the ability of monomeric-binding sites to drive transcription in vitro, indicate an expanded transactivation mechanism incorporating GR monomers. In support of this, similar monomeric function has recently been described in vitro for human U2OS osteosarcoma cell lines carrying stably integrated GR alleles (Schiller et al. 2014). Revision of the transrepression mechanism may also be in order to include binding of GR monomers to half sites. Direct binding to so-called negative GREs by GR monomers has been proposed to mediate repression (Surjit et al. 2011; Hudson et al. 2013), but we find little interaction between GR and this sequence motif, consistent with earlier studies (Grøntved et al. 2013; Uhlenhaut et al. 2013) and indicating that this scenario does not play a prominent role in macrophages or liver. Rather, we observe ligand-dependent redistribution of GR and RNAPII from monomeric to dimeric GR binding sites in liver. The model supposes that GR and RNAPII are available in limiting amounts so that gain of occupancy at one set of sites leads to loss at another. Note that prednisolone, whose potency is four to five times that of corticosterone, was delivered at a pharmacological dosage that exceeds even the highest physiological GC level by at least two orders of magnitude (Hermann et al. 2009). Thus, we believe that exogenous GC, at pharmacological levels, is required for redistribution of GR from monomeric sites that are occupied under physiological levels of hormone.

Sequence-specific binding by GR monomers is compatible with tethering as a mode of action. Tethering was originally described using reporter assays with constructs lacking GR motifs and overexpressed TFs. It is thought to occur when GR targets a DNA-bound TF to indirectly bind DNA, and is generally invoked to explain ChIP-seq peaks lacking the expected motif for the immunoprecipitated TF. However, tethered sites often reside at regions bound by multiple TFs such that an unambiguous determination of the tethering factor is not possible. Using the superior resolution of ChIP-exo, we identify a FOXA protein(s) and ONECUT1 as GR tethering partners in liver. That a GR half-site motif is positioned nearby these sites suggests a role for sequence-specific binding. We favor a model termed half-site facilitated tethering, where sequence-specific interaction of GR monomers to half-site motifs promotes and/or stabilizes transient contacts between monomers and nearby TFs. These are captured by formamide treatment, but whether the protein–protein crosslinks are formed with monomers or off the DNA cannot be determined by our data. However, little or no evidence for

![Sequence Motif P-value Hit % Bg %](image1)

**Figure 5.** GRdim partitions the GR cistrome in primary macrophages. (A) Top-ranked de novo motifs from HOMER for the WT-selective and common GR binding sites from ChIP-seq in primary macrophages. Comprehensive motif results are reported in the Supplemental Material. (B) Box plots interrogating the co-occupancy of macrophage TFs at GR binding sites. (C) Regions named after the nearest gene that include WT-selective and common sites were assayed by a luciferase reporter in the absence (+) or presence (−) of dexamethasone (Dex) and cotransfection of empty vector (E.V.), GR, GRdim, or the DNA-binding mutant GRhalf. (D) Response of WT-regulated genes (up or down) from LPS- and dexam-treated macrophages in GRdim-derived macrophages under the same treatment.
Glucocorticoids (GCs) redistribute GR from monomeric to dimeric sites at regulated genes. (A) Density heatmap of ChIP-seq reads from GR peaks at 6 a.m. with at least 2 RPM in any condition. Prednisolone (pred) treatment of WT and GRdim mice spanned 24 h, and regulated sites in WT mice have at least a twofold difference compared with untreated controls. Top-ranked GO terms for the gained and lost sites are indicated on the left. MEME top-ranked de novo sequences are indicated on the right. See Supplemental Material for a comprehensive list of motifs. (B) GR ChIP-exo at gained sites in WT mice with or without Pred treatment. Average profiles (top) and density heatmaps (bottom) show the S’ ends of the forward-stranded (red) and reverse-stranded (blue) sequence tags. (C) Box plot comparing CEBPβ occupancy at sites with increased GR binding (gained) after Pred treatment. (D) Scatter plot comparing sequence tags from GR ChIP-seq in WT liver with and without Pred treatment, with sites containing up- or down-regulated RNAPII occupancy in response to Pred highlighted in red or blue, respectively. (E) Fraction of Pred-regulated genes with a transcription start site within 100 kb of a gained or lost GR binding site (top) or a non-Pred-regulated dimeric or monomeric GR binding site (bottom).

Figure 6. Glucocorticoids (GCs) redistribute GR from monomeric to dimeric sites at regulated genes. (A) Density heatmap of ChIP-seq reads from GR peaks at 6 a.m. with at least 2 RPM in any condition. Prednisolone (pred) treatment of WT and GRdim mice spanned 24 h, and regulated sites in WT mice have at least a twofold difference compared with untreated controls. Top-ranked GO terms for the gained and lost sites are indicated on the left. MEME top-ranked de novo sequences are indicated on the right. See Supplemental Material for a comprehensive list of motifs. (B) GR ChIP-exo at gained sites in WT mice with or without Pred treatment. Average profiles (top) and density heatmaps (bottom) show the S’ ends of the forward-stranded (red) and reverse-stranded (blue) sequence tags. (C) Box plot comparing CEBPβ occupancy at sites with increased GR binding (gained) after Pred treatment. (D) Scatter plot comparing sequence tags from GR ChIP-seq in WT liver with and without Pred treatment, with sites containing up- or down-regulated RNAPII occupancy in response to Pred highlighted in red or blue, respectively. (E) Fraction of Pred-regulated genes with a transcription start site within 100 kb of a gained or lost GR binding site (top) or a non-Pred-regulated dimeric or monomeric GR binding site (bottom).

Methods
Animal care
GRdim mice were back-crossed for at least four generations to the BALB/c background. Twelve-week-old male GRdim and GR+/+...
littermate controls were maintained in a 12-h dark–light cycle. For ChIP-seq in liver, 6 a.m. and 6 p.m. tissues were collected at the beginning of the light and dark phases, respectively. A third group of mice was treated for 24 h with subcutaneous applied prednisolone pellets (12 mg/kg; Innovative Research of America). All mice were kept under standardized conditions with water and food ad libitum in a specific pathogen-free animal facility at the Leibniz Institute for Age Research (Fritz Lipmann Institute, Jena) and at the University of Ulm. Animal experiments were performed using procedures in accordance with the Thüringer Landesamt für Lebensmittelsicherheit und Verbraucherschutz and the Regierungspräsidium in Tübingen, Germany.

Cell and tissue preparation

Bone-marrow–derived macrophages were isolated and differentiated in culture as previously described (Barish et al. 2005). Cells were treated overnight with 1 μM dexamethasone (Dex; Sigma) and/or LPS (100 ng/ml, Sigma) for 3 h. Liver tissue isolated from deceased mice was immediately frozen in liquid nitrogen. To process for ChIP, the tissue was thawed, minced, and crosslinked in 1% formaldehyde/PBS. Cells were disrupted and nuclei purified by Dounce homogenization in a hypotonic buffer (20 mM HEPES/NaOH at pH 7.5, 0.25 M sucrose, 3 mM MgCl2, 0.2% NP-40, 3 mM 2-mercaptoethanol, complete protease inhibitor cocktail). To prepare ChIP extracts, nuclei were suspended in SDS lysis buffer (50 mM HEPES/NaOH at pH 7.5, 1% SDS, 10 mM EDTA, complete protease inhibitor), incubated for 10 min at 4°C, and subjected to microtip probe sonication under conditions optimized for IP efficiency.

ChIP-seq and ChIP-exo

ChIP-seq in primary macrophages was performed as described earlier (Uhlenhaut et al. 2013). ChIP in liver was performed with sonicated, nuclear extract prepared from formaldehyde-crosslinked tissue. Each experimental condition, e.g., genotype (WT or GRdim), time of harvest (6 a.m. or 6 p.m.), treatment (with or without prednisolone), was assayed in duplicate with separate mice for ChIP-seq, and the same biological replicates were examined by ChIP-exo. The following antibodies were used: GR (PA1-511A, Pierce; sc-1004, Santa Cruz), RNPAP1 (sc-899, sc-9001, Santa Cruz), and CEBPB (sc-150, Santa Cruz). ChIP-seq libraries were produced and sequenced according to Illumina protocols as previously described (Steiger et al. 2010), while ChIP-exo libraries were prepared according to Serandour et al. (2013). Sequencing was performed with multiplexed libraries. Details for ChIP-seq and ChIP-exo data processing are reported in the Supplemental Material.

RNA-seq

RNA isolation from primary macrophages was carried out after overnight incubation with dexamethasone and 6-h LPS treatment using Qiagen RNeasy kit. Libraries were made per manufacturer’s instructions using Illumina TruSeq sample prep kits and sequenced using an Illumina HiSeq.

Luciferase reporter assays

Luciferase assays were carried out as described elsewhere (Uhlenhaut et al. 2013). In short, candidate enhancers were amplified from mouse genomic DNA using oligos that added 5’ XhoI and 3’ HindIII or BgIII restriction sites. Oligo sequences were reported in the Supplemental Information. Cis-regulatory elements are cloned into pGL4.23 (Promega). After transfection, CV-1 cells were treated overnight with 1 μM Dex or ethanol, and reporter activity was determined with Promega’s ONE-Glo system.

Data access

Genome-wide data sets from this study have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession numbers GSE59752 (liver), GSE59764 (macrophage ChIP-seq), and GSE68160 (macrophage RNA-seq).

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