Structural analysis of the glucocorticoid receptor ligand-binding domain in complex with triamcinolone acetonide and a fragment of the atypical coregulator, small heterodimer partner

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Novel Glucocorticoid Receptor-Ligand Structure

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Number of Text Pages: 19
Number of Tables: 1
Number of Figures: 6
Number of Words in Abstract: 234
Number of Words in Introduction: 785
Number of Words in Discussion: 774

Abbreviations: Activator Function (AF); Activator Protein-1 (AP-1); Dexamethasone (Dex); Differential Scanning Fluorimetry (DSF); DNA Binding Domain (DBD); Glucocorticoids (GCs); Glucocorticoid Receptor (GR); GR Binding Sequences (GBS); Inverted Repeat Binding Sequences (IR-GBS); Isopropyl-thio-D-galactopyranoside (IPTG); Ligand Binding Domain (LBD); Maltose Binding Protein (MBP); Molecular Dynamics (MD); Mometasone Furoate (MF); Nuclear Factor Kappa-Beta (NF-κB); Nuclear Receptor (NR); Nuclear Receptor Coactivator-2 (TIF2); Protein Data Bank (PDB); Root Mean Square Deviation (RMSD); Small Heterodimer Partner (SHP); Triamcinolone acetonide (TA); Tobbaco Etch Virus (TEV)
Abstract

The synthetic glucocorticoids (GCs) dexamethasone, mometasone furoate, and triamcinolone acetonide have been pharmaceutical mainstays to treat chronic inflammatory diseases. These drugs bind to the glucocorticoid receptor (GR), a ligand-activated transcription factor and member of the nuclear receptor superfamily. GR is widely recognized as a therapeutic target for its ability to counter pro-inflammatory signaling. Despite the popularity of GCs in the clinic, long-term use leads to numerous side effects, driving the need for new and improved drugs with less off-target pharmacology. X-ray crystal structures have played an important role in the drug-design process, permitting the characterization of robust structure-function relationships. However, steroid receptor ligand-binding domains (LBDs) are inherently unstable and their crystallization has required extensive mutagenesis to enhance expression and crystallization. Here, we utilize an ancestral variant of GR as a tool to generate a high-resolution crystal structure of GR in complex with the potent glucocorticoid triamcinolone acetonide (TA) and a fragment of the small heterodimer partner (SHP). Using structural analysis, molecular dynamics and biochemistry, we show that TA increases intramolecular contacts within the LBD to drive affinity and enhance stability of the receptor-ligand complex. These data support the emerging theme that ligand-induced receptor conformational dynamics at the mouth of the pocket play a major role in steroid receptor activation. This work also represents the first GR structure in complex with SHP, which has been suggested to play a role in modulating hepatic GR function.
Introduction

The nuclear receptor superfamily is comprised of a family of ligand-regulated transcription factors that are critical for maintaining specific gene expression profiles across a number of biological processes (Nagy and Schwabe, 2004). The glucocorticoid receptor (GR), the founding member of the nuclear receptor superfamily, is ubiquitously expressed and both up- and down-regulates thousands of genes involved in immunity, metabolism, and inflammation (Kadmiel and Cidlowski, 2013). GR has a modular architecture consisting of five primary domains: an N-terminal region that contains an activator function (AF-1) surface that interacts with coregulators; a highly conserved zinc finger DNA-binding domain (DBD); a flexible hinge region; and a ligand-binding domain (LBD), which binds ligands that modulate receptor activity. Ligands drive conformational changes within the LBD that modulate a second activation function surface (AF-2), which in turn enables selective interaction with coregulators (Carson-Jurica et al., 1990; Gronemeyer and Moras, 1995; Kumar and Thompson, 1999; Nagy and Schwabe, 2004). Unliganded GR resides in the cytosol bound to chaperone proteins. Upon binding to glucocorticoids, GR undergoes a conformational change resulting in an exchange of chaperones and translocation to the nucleus (O'Malley and Tsai, 1992; Pratt and Toft, 1997). In the nucleus, GR interacts with the genome via multiple mechanisms to then recruit transcriptional coregulator proteins, which serve to promote or repress transcription (Darimont et al., 1998; Hu and Lazar, 1999). The simplistic view of GR signaling suggests that GR will bind directly to DNA at canonical GR binding sequences (GBS) as a homodimer to activate transcription or can bind as a monomer to inverted repeat GBS (IR-GBS/nGREs) to repress transcription (Hudson et al., 2013; Luisi et al.,
GR can also regulate transcription in a DNA-independent manner by interacting directly with the pro-inflammatory transcription factors activator protein-1 (AP-1) and nuclear factor kappa-beta (NF-κB) through protein-protein interactions to block their activity (Mordacq and Linzer, 1989; Schule et al., 1990; Yang-Yen et al., 1990). This mechanism, known as tethering, is the major mechanism targeted by pharmaceutical companies for anti-inflammatory therapies (De Bosscher et al., 2000).

Since their discovery in 1948, glucocorticoids (GCs) have been the most efficacious treatment for chronic inflammatory diseases such as rheumatoid arthritis and asthma (Cato and Wade, 1996). GC-bound GR is unmatched in its anti-inflammatory action, resulting in its prevalence in the clinic and worldwide sales that surpass 10 billion dollars per year (Clark and Belvisi, 2012). Yet despite the effectiveness of GCs, long-term exposure leads to numerous debilitating side effects including weight gain, muscle wasting, and development of Cushing’s syndrome (Schacke et al., 2002). These adverse effects have generally been attributed to GR’s role in transactivation (Schacke et al., 2002), spurring studies focused on the molecular mechanisms of GR-ligand interactions to support the development of more selective GCs.

Triamcinolone Acetonide (TA) is a highly potent synthetic glucocorticoid that has been FDA-approved to treat allergic rhinitis as a nasal spray (Nasacort®; Sanofi, Bridgewater, New Jersey) (Doggrell, 2001; Fredman and Tenenhaus, 2013). It is also used to treat macular edema as a result of diabetes (KENALOG) and as an over-the-counter cream to treat skin lesions (Bristol-Myers Squibb Company; Princeton, New Jersey) (Fredman and Tenenhaus, 2013). In addition to its prolific clinical uses, TA has been widely used in the laboratory setting to probe GR biology since it shows a higher
affinity than dexamethasone (Dex) and promotes the soluble expression and purification of the full length receptor for \textit{in vitro} biochemical characterization (Bain et al., 2012; Derendorf and Meltzer, 2008; Nehme et al., 2009; Robblee et al., 2012).

Soluble expression and purification of human GR LBD is challenging and requires mutation at several sites to promote solubility and purification (Gebhardt et al., 2013). These mutations, designed via sequence alignments to other steroid receptor LBDs, enable expression in \textit{E. coli} and greatly enhance crystallization. However, their effect on ligand selectivity, conformational dynamics and activity remains unknown, resulting in uncertainty when constructing robust structure-function relationships. Recently, resurrected ancestral proteins have been used as a tool not only to study evolutionary biology, but also to generate x-ray crystal structures of novel complexes (Bridgham et al., 2009; Colucci and Ortlund, 2013; Kohn et al., 2012). Here, we utilize AncGR2, the ancestral precursor to the modern GR, which recapitulates both the structure and function of its extant counterpart while exhibiting better expression, purification, and crystallization in the laboratory setting along with a higher tolerance to mutagenic analysis, which is critical for probing structure-function relationships (Bridgham et al., 2009; Kohn et al., 2012; Ortlund et al., 2007; Thiruchelvam et al., 2011). AncGR2 and hGR LBDs share 78.6 \% sequence identity with 100 \% of the residues within the ligand binding pocket conserved. This makes it an ideal surrogate for hGR structure-function studies, and allowed us to generate the first x-ray crystal structure of the GR LBD complexed with TA and with a peptide from the atypical coregulator, small heterodimer partner (SHP).
**Materials and Methods**

**Protein Expression and Purification**

AncGR2 LBD (GenBank accession number EF631976.1) was cloned into a pMALCH10T vector, transformed into *Escherichia coli* strain BL21(pLysS), and expressed as a maltose-binding protein (MBP) with a hexahistidine tag. Cultures (9 L in Terrific Broth) were grown at 37 °C until an OD$_{600}$ of 0.6. Protein expression was induced with the addition of 0.5 mM isopropyl-1-thio-D-galactopyranoside (IPTG) and 25 µM TA or 50 µM dexamethasone and grown for 4 hours at 32 °C. Cells were lysed in a buffer containing 20 mM Tris-HCl (pH 7.4), 300 mM NaCl, 25 mM imidazole, and 5 % glycerol via sonication on ice. Fusion protein was purified by affinity chromatography (GE Healthcare His-Trap FF) and the MBP tag was removed by Tobbaco Etch Virus (TEV) protease cleavage with an additional two passes over HisTrap FF media (GE Healthcare). After purification, protein was dialyzed overnight at 4 °C into buffer containing 20 mM HEPES (pH 8.0), 200 mM NaCl, 10 % glycerol, and 50 mM CHAPS. Protein was concentrated to 3-5 mg/mL, flash frozen in liquid N$_2$, and stored at -80 °C.

**Protein Crystallization, Data Collection, and Structure Determination**

AncGR2 LBD-TA was concentrated to 4.75 mg/mL and incubated with a peptide derived from SHP (NH2-QGASRPAILYALLSSSLK-OH) at two-fold molar excess. Crystals were grown by hanging drop vapor diffusion at 18 °C in drops containing 1 µL of AncGR2-TA-SHP and 1 µL of 100 mM HEPES pH 7.5, 1.5 % glycerol, and 25 % PEG 300. Crystals were flash frozen in liquid nitrogen, using a cryopreservative consisting of 100 mM HEPES pH 7.5, 10 % glycerol, and 40 % PEG 300. Data were
collected remotely from the South East Regional Collaborative Access Team (SER-CAT) at the Advanced Photon Source (APS), 22ID beamline (Argonne National Laboratories, Chicago, IL). Data were processed and scaled using HKL-2000 (Otwinowski and Minor, 1997) and phased by molecular replacement using Phaser-MR (Adams et al., 2010). The structure was phased using the previously solved AncGR2 – dexamethasone – nuclear receptor coactivator 2 (TIF2) peptide complex as a search model (3GN8) (Bridgham et al., 2009). Structure refinement and validation was performed using PHENIX (v1.11.1) and model building was performed in COOT (Adams et al., 2010; Emsley et al., 2010). PDB Redo was used iteratively to optimize refinement parameters and geometry (Joosten et al., 2009). PyMOL (v1.8.2) was used to visualize structures and generate figures (Schrödinger, LLC).

$R_{\text{factor}}$ for the final model are 21.5 % and 25.2 % for $R_{\text{work}}$ and $R_{\text{free}}$, respectively. MolProbity was used for model validation, indicating that 98 % of the residues fall in the most favored regions of the Ramachandran plot with none in disallowed regions (Adams et al., 2010; Davis et al., 2007). The overall MolProbity score was 0.70, placing the structure in the 100th percentile for overall geometric quality among protein crystal structures of comparable resolution. The remaining data collection and refinement statistics can be found in Table 1. Coordinates and structure factors have been deposited in the Protein Data Bank (PDB) with the accession number, 5UFS.

*Ligand Binding & Competition Assays*
Hexahistidine-tagged MBP-fused AncGR2 LBD-Dex was dialyzed overnight into buffer containing 150 mM NaCl, 10 mM HEPES pH 7.4, 3 mM EDTA, 5 mM DTT, and 0.005 % Tween-20. All binding experiments were performed using this buffer. Binding affinity for dexamethasone-fluorescein was measured with 12 nM dexamethasone-fluorescein and protein concentrations from $10^{-10}$ to $10^{-5}$ M. Polarization was monitored on a Biotek Neo plate-reader at an excitation/emission wavelength of 485/528 nm (Winooski, VT). Three technical replicates and three biological replicates were conducted and graphs are a compilation of all data collected. Binding data were fit with a one-site binding curve in GraphPad Prism v7 (GraphPad, Inc). Competition assays were performed at a protein concentration 1.2 times the binding affinity for dexamethasone and in the presence of 12 nM dexamethasone-fluorescein and $10^{-10}$ to $10^{-5}$ M of competing ligand. Three technical replicates and three biological replicates were conducted and graphs are a compilation of all data collected. GraphPad Prism v7 was used to analyze data using a one-site, fit $K_i$ curve.

**Differential Scanning Fluorimetry (DSF)**

DSF assays for protein-ligand complexes (**Figure 2d**) were performed with 5 µM of AncGR2 LBD (expressed and purified with varying ligands), buffer containing 20 mM Tris pH 7.4, 300 mM NaCl, and 5 % glycerol, and a final 1:1000 dilution of SYPRO® orange dye (Sigma) to a final volume of 20 µL. Reactions with coregulator peptides (**Figure 5e**) consisted of 5 µM protein, 5 µM peptide, buffer, and SYPRO dye as above. The peptide sequences used were as follows: SHP NR Box 1 (NH2-QGASRPAILYALLSSSLK-OH) and TIF2 NR Box 3 (NH2-KENALLRYLLDKDD-
OH). For all DSF experiments reactions were heated from 25 °C to 95 °C in 0.5 °C increments every minute and fluorescence was monitored using the ROX filter (602 nm) with a StepOne Plus Real Time PCR System (ThermoFisher). Two technical replicates and three biological replicates were conducted and graphs are a compilation of all data collected. Data was normalized by subtracting each data point from a dye-buffer only control reaction. Data was fit using a Boltzmann sigmoidal curve to determine the melting temperature ($T_m$), defined as 50 % unfolding.

**ProSMART Analysis**

ProSMART alignment tool provides a conformation-independent structural comparison of two proteins (Nicholls et al., 2014). Pairwise comparisons were conducted between AncGR2 LBD-TA-SHP and AncGR2 LBD-Dex-TIF2 (PDB 3GN8); AncGR2 LBD-TA-SHP and AncGR2 LBD-MF-TIF2 (PDB 4E2J); AncGR2 LBD-Dex-TIF2 and AncGR2 LBD-MF-TIF2 (Bridgham et al., 2009; Kohn et al., 2012). The comparisons generate a Procrustes score, which is mapped onto the LBD structures. This score is the r.m.s.d. of the central residue of two corresponding structural fragments of length $n$, where $n$ is an odd number of amino acids.

**Molecular Dynamics Simulations**

Three complexes were prepared for molecular dynamics simulations using AncGR2 LBD: i) TA-TIF2 ii) Dex-TIF2 (PDB 3GN8) and iii) MF-TIF2 (PDB 4E2J). The structure for TA-TIF2 was created by replacing SHP peptide with TIF2 and an all-atom minimization of the structure. The complexes were solvated in an octahedral box of
TIP3P (Jorgensen et al., 1983) water with a 10 Å buffer around the protein complex. Na\(^+\) and Cl\(^-\) ions were added to neutralize the protein and achieve physiological conditions, to a final concentration of 150 mM. All systems were set up using xleap in AmberTools (Case and Kollman, 2012) with the parm99-bsc0 forcefield (Pérez et al., 2007). Parameters for TA, MF and Dex were obtained using Antechamber (Wang et al., 2001) in AmberTools. All minimizations and simulations were performed with Amber14 (Case et al., 2014). Systems were minimized with 5000 steps of steepest decent followed by 5000 steps of conjugate gradient minimization with 500 kcal/mol·Å\(^2\) restraints on all atoms. Restraints were removed from all atoms excluding the atoms in both the ligand and the TIF2 peptide, and the previous minimization was repeated. The systems were heated from 0 to 300 K using a 100-ps run with constant volume periodic boundaries and 5 kcal/mol·Å\(^2\) restraints on all protein and ligand atoms. Twelve ns of MD equilibration was performed with 10 kcal/mol·Å\(^2\) restraints on protein and ligand atoms using the NPT ensemble. Restraints were reduced to 1 kcal/mol·Å\(^2\) for an additional 10 ns of MD equilibration. Then restraints were removed and 500 ns production simulations were performed for each system in the NPT ensemble. A 2 fs timestep was used and all bonds between heavy atoms and hydrogens were fixed with the SHAKE algorithm (Ryckaert et al., 1977). A cut-off distance of 10 Å was used to evaluate long-range electrostatics with Particle Mesh Ewald (PME) and for van der Waals forces. 25,000 evenly spaced frames were taken from each simulation for analysis.

MD trajectories were analyzed with various tools. Structural averaging and analysis were performed with the CPPTRAJ module (Roe and Cheatham III, 2013) of AmberTools (v14). Hydrogen bonds were identified using HBPLUS (McDonald and
Thornton, 1994) with default criteria (minimum angles: D-H-A 90.0, H-A-AA 90.0, D-A-AA 90; maximum distances D-A 3.9Å, H-A 2.5Å, D = Donor, A = acceptor, AA= atom attached to acceptor). Root mean square fluctuations (RMSF) analysis was performed on Cα atoms of protein residues. RMSF was computed relative to the reference structure (crystal structure) for each frame in the trajectory. CPPTRAJ was used to calculate distances between Cα atoms or hydrogen bonding atoms over trajectories. All distances between residue pairs were calculated using Cα atoms. The MMTSB toolset was used to perform a cluster analysis with a 2 Å RMSD cutoff (Feig et al., 2004).

Results

Structural analysis of AncGR2-Triamcinolone/SHP complex

To understand how TA interacts with GR, we determined the x-ray crystal structure of AncGR2 LBD – TA complex bound to a fragment of the coregulator SHP. Crystals formed in the C2 space group with two AncGR2 LBD – TA – SHP complexes in the asymmetric unit. Data sets were collected to 2.1 Å with 98.6 % completeness. Full data collection and refinement statistics are summarized in Table 1.

The AncGR2 LBD is bound by one molecule of TA and the SHP coregulator peptide (Figure 1a). The LBD adopts a canonical fold with 11 α-helices and 4 β-strands that fold into three layers of a helical sandwich bundle (Figure 1a) (Kumar and Thompson, 1999). This folding creates a hydrophobic ligand binding pocket (LBP) encompassing TA (Figure 1b), which is supported by unambiguous electron density (Figure 1c).
TA is coordinated via extensive hydrophobic contacts and a series of specific hydrogen-bonding interactions (Figure 1d). This hydrogen bonding network includes a water molecule, which is a common feature for SR LBD-ligand complexes and is seen in all AncGR2 LBD-ligand structures to date (Bridgham et al., 2009; Kohn et al., 2012). This water hydrogen bonds with the terminal oxygen of the A-ring. The amide of Gln39 and guanidinium group of Arg80 not only hydrogen bond to the water molecule but also form hydrogen bonds to the A-ring 3-keto oxygen (Figure 1d,2a). Molecular dynamics simulations (MD) of ligand-hydrogen bonding interactions show that over a 500 ns simulation, these hydrogen bonds alternate between being direct or water-mediated (Figure 2a). In addition, there is a water-mediated hydrogen bond that occurs between the Arg80 and Gln39 residues that persists for 47% of the simulation. Both of these hydrogen-bonding interactions have a joint effect in keeping TA in the correct orientation within the LBP over the duration of the simulation. Asn33 hydrogen bonds to the C-ring 11-hydroxyl. Finally, Asn33 and Thr208 hydrogen bond to the 21-hydroxyl located off the C-17 position on the D-ring. These hydrogen bonds are well supported via MD and persist throughout the entirety of the simulations (Figure 2b).

The C-17 acetonide moiety on TA increases the affinity and stability for GR LBD over Dex.

Synthetic GCs typically show increased affinity and selectivity for the receptor over GR’s endogenous ligand, cortisol. Corticosteroids, like most cholesterol-derived steroid hormones, contain three 6-carbon rings (A, B, C) and one 5-carbon ring (D) (Figure 3a). Dex varies from cortisol by the addition of a C1-C2 double bond in ring A, a
C-16 $\alpha$-methyl, and a C-9 $\alpha$-fluoro group. TA varies from Dex by the addition of the 16,17 acetonide moiety. MF varies from Dex by the presence of a chloro group at C-9 instead of a fluoro group, a chloro group instead of a 21-hydroxyl off the C-17 D-ring, and the large furoate moiety off the C-17 position. The C-17 furoate moiety is more polar and labile than the TA acetonide moiety and is cleaved \textit{in vivo}. The active metabolite, mometasone, contains a $\alpha$ hydroxyl at the C17 position due to loss of the fuorate moiety; thus, conclusions based on analysis of the GR-MF complex cannot be reliably correlated with its \textit{in vivo} potency. However, MF has proven a useful tool to probe GR LBD biochemistry \textit{in vitro} and is therefore included in our analysis. For synthetic GCs, the steroid backbone halogen substitutions are thought to increase potency and the substitutions at position 17 account for the largest functional differences (Bikowski et al., 2006; Derendorf and Meltzer, 2008). These substitutions on TA and MF drive the increased affinity and reported potencies over Dex (Grossmann et al., 2004; Nehme et al., 2009).

Similar to previous reports, we show that TA displaces bound fluorescent Dex (FL-Dex) from the AncGR2 LBD with a $K_i$ of 3.2 nM ($K_d$ for FL-Dex = 38 nM) (\textbf{Fig 3b}) (Bledsoe et al., 2002; Grossmann et al., 2004; Nehme et al., 2009; Yoneda et al., 1995). Furthermore, we show there is an 8 $^\circ$C increase in thermal stability of the LBD when bound by TA or MF over Dex (\textbf{Figure 3c}). To understand how TA has such a high affinity and potency for GR, we compared our structure to the previously solved AncGR2-Dex-TIF2 and AncGR2-MF-TIF2 complexes (PDB 3GN8, 4E2J) (Bridgham et al., 2009; Kohn et al., 2012) (\textbf{Figure 4}).
Superposition of the different ligand-bound structures revealed only subtle differences in the RMSD’s of main chain carbon alphas. The difference between AncGR2-TA and AncGR2-Dex is 0.346 Å, and between AncGR2-TA and AncGR2-MF is 0.349 Å. Within the ligand-binding pocket, the same side chains participate in hydrogen bonding interactions, including a water molecule in all structures (Figure 4a). This is not surprising when comparing Dex and TA, as the ligands are almost identical except for the acetonide moiety, which is contacted by hydrophobic interactions. However, MF-bound GR makes one fewer hydrogen bonding interaction; the C17 position contains a chloro group rather than a hydroxyl, altering the interaction with Thr208. Yet, as with MF, the acetonide C-17 addition is oriented almost 90° from the steroid backbone. This allows for the ligand to expand and fill the binding pocket as well as generate additional hydrophobic contacts between GR and the ligand. These large substitutions on TA and MF cause a 100 Å³ (1.1-fold) and 200 Å³ (1.3-fold) increase, respectively, in pocket volume compared to Dex (Figure 4b). Therefore, the differences in potency and affinity must be due to changes in hydrophobic contacts and long-range allosteric changes generated by the ligands. To investigate this possibility, we combined in-depth structural analysis with MD simulations.

**TA increases intramolecular contacts within the LBD to drive affinity and stability**

To identify how differential ligands affect local structure, we performed pairwise comparisons of the three AncGR2 ligand structures using ProSMART (Nicholls et al., 2014) (Figure 5a). Each chain is compared separately and the final models are colored by their Procrustes score, which defines the similarity of an aligned fragment according
to the legend provided in Figure 4. The largest differences are seen in the mouth of the ligand binding pocket, encompassing the loops after H1, H3, and H6, and the transition between H9 and H10. Differences in the H9-H10 loop are not surprising, as this loop is often difficult to model and is not supported by strong electron density in any structure. However, there are meaningful differences in local residue conformation at the entrance to the ligand binding pocket, where TA constricts the opening of the pocket to 4.6 Å, compared to 6.3 Å for Dex and 10.1 Å for MF as measured by the distance between Ser25 on H3 and Pro106 on H6. (Figure 5b). This TA-driven pocket constriction is also observed in MD simulations. The average Ser25-Pro106 distances in the simulations are 6.4 Å, 10.5 Å and 11.3 Å respectively for TA, Dex and MF. To link differential contacts with ligand-induced differences in receptor motion, we used MD to calculate the root mean square fluctuations between the key LBD-ligand complexes over 500 ns simulations. To focus only on ligand-induced dynamic changes, we first replaced SHP with TIF2 to mitigate peptide-driven effects. The MD trajectories showed less motion at residues 101-109 (Helix 6-7) in the TA complex compared to both Dex and MF, suggesting TA introduces a stabilizing effect (Figure 5c,d).

To obtain the structure most representative of the dominant conformation sampled in each trajectory, clustering was performed with the MMTSB tool. For each AncGR2 complex, the structure with the lowest RMSD in the most populated cluster was used as a representative for the complex. Comparison of these structures shows a rearrangement of the residues in the binding pocket of the TA bound structure as a result of stabilizing, hydrophobic interactions between the acetonide group and Met29 and Met108 (Figure 5e). The interaction causes a shift of other residues (e.g. Met103) and shifts in the overall
positions of H6 and H7, bringing them closer (Figure 5f,g). This movement is responsible for the closer distances observed between Ser25 and Pro106 in the TA-bound structure. Similar effects are observed in AncGR2 LBD-TA-SHP and LBD-TA-TIF2 structures, indicating that the TA ligand drives this effect.

**Recognition of the SHP NR Box 1 LXXLL Motif**

Coregulator proteins interact with SRs via an α-helix containing a short LXXLL motif (L-leucine, X-any amino acid) (Jenkins et al., 2001; Millard et al., 2013). These coregulators bind to the same hydrophobic groove on the surface of SR LBDs, the AF-2 (Heery et al., 1997; Vandevyver et al., 2014). The AF-2 is comprised of helices 3, 4, and 12 and is generally held in place by a charge clamp formed by a lysine on H3 and a glutamate on H12 that interacts with the helix dipole (Bledsoe et al., 2002; Vandevyver et al., 2014). GR is known to interact with a wide variety of coregulator proteins, yet structural analysis has been limited to the typical coregulators of the p160 SRC family, specifically SRC-2 (TIF2, GRIP-1). Here, we show GR complexed with 11 residues of the SHP NR Box 1 peptide. This is the first GR LBD structure with this coregulator.

SHP is an atypical orphan nuclear receptor comprised only of an LBD that co-folds with the transcriptional corepressor EID1 (Lee et al., 2007; Seol et al., 1996; Zhi et al., 2014). SHP has been shown to act as a potent corepressor for numerous nuclear receptors including ER, RXR, LRH-1, and GR (Borgius et al., 2002; Johansson et al., 2000; Johansson et al., 1999; Lee et al., 2000; Ortlund et al., 2005). To modulate receptor activity, SHP utilizes two canonical NR Box motifs to bind to the AF-2 surface, competing directly with coactivators (Lee et al., 2000). SHP’s repression of GR activity
has been postulated to have important biological implications in the liver and pancreas (Borgius et al., 2002).

The overall AncGR2-TA-SHP structure shows the AF-H in the active conformation and the SHP peptide bound at the AF surface (Figure 6a). The peptide inserts leucine side chains into the surface and is further stabilized by a charge clamp interaction, which is conserved across NRs. Lys48 on H3 hydrogen bonds with the backbone carbonyl of Leu24, and Glu224 hydrogen bonds to the free amide nitrogen of Ile20 (Figure 6b). The terminus of the peptide is stabilized by a hydrogen bond involving Lys48 interacting directly with the terminal Ser27 residue. The peptide is further stabilized by Met62, which makes van der Waals contacts with Tyr22. As in previous NR LBD-SHP structures, this interaction holds the tyrosine residue in the center of the peptide helix, which is further stabilized between the aromatic face and the Ser26 sidechain. The NR Box 1 peptide contains an arginine residue at the first position, but there was not strong enough density to model in this side chain; therefore it is modeled as an alanine residue. The SHP NR Box 1 and 2 peptides look most similar to the LXXLL motif seen in the coactivator PGC-1α (Figure 6d). It has been postulated that SHP antagonizes PGC-1α activation of GR. This repression of GR was shown to inhibit PEPCK, implying a role for SHP in modulating GR function within the liver (Borgius et al., 2002).

The SHP peptide in our structure interacts with the same AF-2 surface seen in the previously solved AncGR2 LBD-TIF2 NR Box 3 structure (Figure 6c) (Kohn et al., 2012). Since SHP interacts with GR via a classical NR coactivator motif, similar interactions are expected. In both GR-peptide complexes, the charge clamp formed by
Lys48 and Glu224 is conserved. Despite these peptides having sequence differences, both peptides stabilize AncGR2 LBD to similar levels (Figure 6d,e).

**Discussion**

In the 1940s, GR’s endogenous ligand, cortisol, was identified as a potent suppressor of inflammation (Kendall, 1951). Since then, synthetic GCs with dramatically improved affinity, potency and selectivity have become the most widely used treatment for anti-inflammatory therapies (Clark and Belvisi, 2012). Targeting GR is not challenging, however, current GCs drive the activation of metabolic, homeostatic and growth pathways in addition to immunosuppression (Cato and Wade, 1996; De Bosscher et al., 2000; Schacke et al.). X-ray crystal structures are required to build robust structure-function relationships; however, this has proved difficult due to the instability of recombinantly expressed, purified human GR LBD (hGR). To circumvent this issue, we utilized the AncGR2 LBD derived from the phylogenetically reconstructed GR present in the ancestor of bony vertebrates (Bridgham et al., 2009; Colucci and Ortlund, 2013; Kohn et al., 2012). This ancestral receptor has been extensively used to understand the evolution of corticoid selectivity in modern GR; it displays the same ligand selectivity and agonist response as the human receptor but shows enhanced expression, solubility, crystallizability, and tolerance to mutation (Kohn et. al., 2012; Carrol et. al., 2011; Bridgham et. al., 2009; Ortlund et. al., 2007). Therefore, AncGR2 represents a powerful tool to explore novel GR-ligand complexes that would otherwise be difficult to probe.(Bridgham et al., 2009; Colucci and Ortlund, 2013; Kohn et al., 2012).
The synthetic glucocorticoid TA is used in the clinic to treat allergic rhinitis, macular edema, and skin lesions (Fredman and Tenenhaus, 2013; Veritti et al., 2012). TA is more hydrophobic than other synthetic GCs, making it ideal for topical use and allowing for a prolonged duration of action (Doggrell, 2001; Fredman and Tenenhaus, 2013). Furthermore, TA has gained popularity in the laboratory due to its increased affinity for GR over other synthetic compounds (Figure 3c) such as dexamethasone (Veritti et al., 2012; Yoneda et al., 1995), and for its ability to promote the expression and solubility of the intact receptor. These improvements in affinity, potency, and bioavailability over other GCs makes TA a critical ligand to study. We show that TA significantly increases the thermal stability of the LBD relative to dexamethasone, (Figure 3d) causing GR to assume a more compact structure with smaller conformational fluctuations near the pocket (Figure 5). Driving these effects with TA is the bulky hydrophobic C17 acetonide moiety, which generates additional contacts relative to other ligands and repositions the H6-H7 loop to constrict the pocket (Figure 5d-e). The bound TA maintains interactions with the hydrogen bond network critical for GR specificity and transactivation (Figure 1d, 2, 4a), but enables greater intramolecular contacts, which likely explains the increased affinity and stability (Figure 5). Ligand-driven perturbation of intramolecular contacts and dynamics at the mouth of the ligand binding pocket has been shown to be essential for selective modulation of other nuclear receptors such as ERα, PPARγ, and LRH-1 (Gee and Katzenellenbogen, 2001; Kojetin and Burris, 2013; Musille et al., 2012; Srinivasan et al., 2013). Taken together, we have elucidated the structural mechanisms driving TA’s enhanced affinity and ability to stabilize the GR LBD, which will inform future glucocorticoid design.
This work also offers the first visualization of GR’s interaction with the atypical NR, SHP, which is part of a transcriptional corepressor complex that targets NRs in the active conformation (Bavner et al., 2002; Borgius et al., 2002; Macchiarulo et al., 2006; Zhi et al., 2014). SHP accomplishes this by utilizing two LXXLL motifs to mimic coactivators and bind directly to the AF surface. SHP is expressed at high levels in the liver and has been shown to interact directly with GR to modulate its function (Borgius et al., 2002). We demonstrate that indeed, SHP binds to the AF surface on GR and makes the conserved charge clamp interaction, similar to the previously solved structure of AncGR2 in complex with the coregulator TIF2 (Figure 6b). Furthermore, the GR LBD–TA complex was stabilized to similar levels whether in complex with SHP or TIF2 (Figure 6e). SHP has been suggested to play a role in modulating hepatic GR function and thereby metabolism (Borgius et al., 2002). It is likely that increased levels of SHP permit direct competition with coregulators for the GR-agonist-DNA complex (Bavner et al., 2002; Borgius et al., 2002; Macchiarulo et al., 2006; Zhi et al., 2014). However, further studies are needed to test this mechanism and to determine its physiological and clinical relevance for hepatic GR action.

Developing improved GCs with less off target side effects, will require linking ligand-induced receptor motions with selective coregulator interactions to drive GR to specific promoters (Nehme et al., 2009; Sacta et al., 2016; Weikum et al., 2017). Obtaining structural and dynamic information, as presented here, is vital to this effort.
Acknowledgements

X-ray data were collected at Southeast Regional Collaborative Access Team (SER-CAT) 22-ID beamline at the Advanced Photon Source, Argonne National Laboratory. Supporting institutions may be found at www.ser-cat.org/members/html. Use of the Advanced Photon source was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract No. W-31-109-Eng-38.

Authorship Contributions

Participated in research design: Weikum, Okafor, Ortlund.

Conducted experiments: Weikum, Okafor, D’Agostino, Ortlund.

Data Analysis: Weikum, Okafor, D’Agostino, Ortlund.

Wrote or contributed to the manuscript: Weikum, Okafor, D’Agostino, Colucci, Ortlund.
References


Footnotes

Weikum is supported by a predoctoral NRSA awarded by a National Institutes of Health General Medical Sciences [1F31GM113397-01A1]. Ortlund is supported by NIH grant [R01DK095750], AHA grant [14GRNT20460124], and a W. M. Keck Foundation Medical Research Grant. Okafor is funded by a National Institutes of Health General Medical Sciences Institutional Research and Career Development Award [K12 GM000680].
Figure Legends

**Figure 1: Structure of the AncGR2-TA-SHP NR Box 1 Complex.** (a) Cartoon representation of the overall structure of AncGR2 LBD (blue) in complex with TA (Green) and SHP (Pink) (b) Close-up view of ligand binding pocket (side chains are shown as sticks with α-carbons shown as spheres) Residues that participate in hydrophobic contacts are shown in faded blue and side chains that make hydrogen bonding interactions are shown darker blue with Hydrogen bonds denoted by red dashes. (c) Omit map (F₀-Fᵣ) contoured at 2.5 σ around the ligand. (d) Schematic of hydrogen bonding network. Hydrogen bonds are shown in red. (e)

**Figure 2: MD Simulations Support LBP-Hydrogen Bonding Interactions**

(a) Graph of hydrogen bonding distances between the guanidinium hydrogen atom on Arg80 and the amide hydrogen atom on Gln39 with the 3-keto oxygen on the A ring of TA during the 500 ns MD simulation. These hydrogen atoms either make contacts with the water molecule within the pocket or make direct interactions with the ligand. The A ring of TA is labeled in each view of the ligand (b) MD analysis supports the hydrogen bonding interactions seen between Asn33 and Thr308 with the ligand.

**Figure 3: TA readily competes Dex out of the binding pocket and highly stabilizes the LBD.** (a) GR’s endogenous ligand cortisol and a myriad of synthetic glucocorticoids including Dex, TA, and MF. (b) Fluorescent Dex (FL-Dex) binds AncGR2 LBD with a $K_d$ of 38 nM. Binding was measured via fluorescence polarization and graphs are fit using a one site binding curve to calculate $K_d$ values, error bars indicate standard error of
the mean (SEM; n=9). (c) TA competes FL-Dex out of the ligand binding pocket with a $K_i$ of 3.2 nM. Graphs are fit using a one site, fit $K_i$ equation to calculate $K_i$ values, error bars indicate SEM (n=9). (d) DSF monitors the thermal stability of different AncGR2 ligand complexes. Both TA and MF-bound LBD increased the thermal stability by 8 °C. Graphs are fit using the Boltzmann sigmoidal equation, which calculates $T_m$ as 50 % unfolding. Error bars error bars indicate SEM (n=6).

**Figure 4: Comparison of Glucocorticoid Ligands bound to AncGR2 LBD.** (a) Overlay of TA (blue), Dex (purple), MF (green) within the pocket. Insets to the right show TA (blue), Dex (purple) and MF (green) hydrogen bonding networks. Hydrogen bonds are shown in red. (b) Mesh depicting ligand pocket volume induced by different glucocorticoid ligands. TA and MF have large additions at the C-17 position, which the GR LBP expands to accommodate.

**Figure 5: Structural comparison of different GR-ligand complexes.** (a) ProSMART analysis of pairwise comparisons of different structures of GR-ligand complexes. Areas shown in white were not used in the comparison. Structures are colored by Procrustes score of the central residue of an aligned fragment pair according to the legend shown on the right. The following comparisons were made: AncGR2 LBD-TA-SHP *versus* AncGR2 LBD-Dex-TIF2 (PDB 3GN8); AncGR2 LBD-TA-SHP *versus* AncGR2 LBD-MF-TIF2 (PDB 4E2J); AncGR2 LBD-Dex-TIF2 *versus* AncGR2 LBD-MF-TIF2. (b) Structural overlay of the mouth of the ligand binding pocket, which was one of the areas with the largest variance from the analysis done in (a). TA induces a narrowing of the
pocket entrance, indicated by a 4.6 Å distance between Ser25 on H3 and Pro106 on H6. Dex has a 6.3 Å and MF has a 10.1 Å distance across the same area. (c) 500 ns molecular dynamics simulations looking at RMSFs between various GR-ligand complexes. ** indicate areas of significant differences in RMSF that are not obvious due to flexible loops, where higher RMSF values are to be expected. (d) The AncGR2 LBD-TA structure shows the acetonide group on TA makes hydrophobic contacts (black) with Met29 and Met108 (e) The hydrophobic contacts seen in d causes a shift in the position of other side chains within the ligand binding pocket, such as Met103. Structural overlays of AncGR2-ligand complexes show this is unique to the TA bound complex. (f) The shift in side chains causes an overall rearrangement of H6/H7, explaining the constricted pocket entrance seen in the TA structure.

Figure 6: AncGR2-TA complex is bound by atypical coregulator, SHP. (a) Overall structure of AncGR2 with SHP represented as sticks and ribbon (pink). The peptide is bound in the canonical AF-2 surface made by H3, H4, and the AF-H/H12. The peptide is held in place by charge clamp formed by Glu224 and Lys48, shown in sticks (blue). (b) Zoom in of SHP peptide. (c) Overlay of SHP (faded pink) with TIF2 peptide (purple) from the AncGR2-Dex (PDB 3GN8) structure. (d) Comparison of LXXLL motifs found in various coregulators. (e) DSF monitors the thermal stability AncGR2 LBD-TA in complex with the SHP NR Box 1 and TIF-2 NR Box 3 peptide. There is little difference between the two complexes.
### Table 1. Data Collection and Refinement Statistics

<table>
<thead>
<tr>
<th>Data Collection</th>
<th>AncGR2 LBD-TA + SHP</th>
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</thead>
<tbody>
<tr>
<td><strong>Space Group</strong></td>
<td>C2</td>
</tr>
<tr>
<td><strong>Cell Dimension</strong></td>
<td>a=87.0, b=52.8, c=126.0</td>
</tr>
<tr>
<td><strong>Resolution (Å)</strong></td>
<td>2.11 (2.19-2.11)</td>
</tr>
<tr>
<td>$R_{pim}$</td>
<td>6.2 (37.0)</td>
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<tr>
<td>$I/\sigma$</td>
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<tr>
<td><strong>Completeness</strong></td>
<td>98.6 (93.2)</td>
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<tr>
<td><strong>Redundancy</strong></td>
<td>3.6 (3.0)</td>
</tr>
</tbody>
</table>

**Refinement**

- **Resolution**: 2.11
- **No. Reflections**: 31942
- **$R_{work}/R_{free}$**: 21.5/25.2
- **No. Atoms**
  - Protein: 4158
  - Ligands: 62
  - Water: 119
- **$B$-factors**
  - Protein: 44.9
  - Ligands: 31.8
  - Water: 41.0
- **R.m.s. deviations**
  - Bond lengths (Å): 0.002
  - Bond angles (°): 0.52
- **PDB Code**: 5UFS

*Data collected from a single crystal; values in parentheses are for the highest-resolution shell.*
Figures

Figure 1

(a) Structure of the complex between SHP and AF-H, showing the binding of Triamcinolone Acetonide.

(b) Close-up view of the interactions between the ligand and the receptor.

(c) Stereo view of the complex showing the orientation of the ligand.

(d) Molecular details of the interaction, including distances.
Figure 2

Distances are consistent with alternating between direct (< 3 Å) and water-mediated Hbonds (3-5 Å) for the following pairs:

- Arg80 - A ring (3-Keto O)
- Gln39 - A ring (3-Keto O)
- Asn33 - D ring (21-OH)
- Thr208 - D ring (21-OH)
- Asn33 - C ring (11-OH)
Figure 3

a) Structures of various glucocorticoids: Cortisol, Dexamethasone, Triamcinolone Acetonide, and Mometasone Furoate.

b) Graph showing polarization (mP) against concentration of GR 2 LBD [M].

- $K_d = 38 \pm 3.75 \text{ nM}$

(c) Graph showing polarization (mP) against log concentration of Triamcinolone Acetonide.

- $K_i = 3.2 \pm 1.2 \text{ nM}$

(d) Graph showing fluorescence against temperature (°C).

- AncGR2-TA: $T_m = 55.1 \pm 0.15 \text{ °C}$
- AncGR2-Dex: $T_m = 47.1 \pm 0.12 \text{ °C}$
- AncGR2-MF: $T_m = 55.5 \pm 0.13 \text{ °C}$
- No protein: $T_m$
Figure 4

a

Arg80
Gln39
Asn33
Thr208
TA  Dex  MF

b

Arg80
Gln39
Asn33
Thr208
TA  Dex  MF

Gln39
Arg80
Asn33
Thr208
TA  Dex  MF

Gln39
Arg80
Asn33
Thr208
TA  Dex
MF

Gln39
Arg80
Asn33
Thr208
TA  Dex
MF

Gln39
Arg80
Asn33
Thr208
TA  Dex
MF

Gln39
Arg80
Asn33
Thr208
TA  Dex
MF

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Figure 5

(a) Structural comparison of TA-SHP, Dex-TIF2, and MF-TIF2. 
(b) Close-up view of specific interacting residues. 
(c) Root mean square fluctuations (Å) with significant differences indicated. 
(d) Comparison of secondary structure elements. 
(e) Distance between Met29 and Met108. 
(f) Met108 and Met103 interactions. 
(g) Overall structure comparison.
Figure 6

a

SHP AF-H/H12

H3

H4

b
c

Glu224

Lys48

Asn742

Ala743

Leu744

Leu745

Arg746

Tyr747

Leu748

Leu749

Asp750

Lys751

Asp752

d

SHP NR Box 1 APAILYALLSS

SHP NR Box 2 VPSILKKILLE

PGC-1α EPSLLKLLLLA

TIF-2 NR Box 3 NALLRYLLDKD

LXXLL Motif

e

Fluorescence

Temperature (°C)

AncGR1-TA + SHP 55.0 ± 0.15 °C

AncGR1-TA + TIF2 55.8 ± 0.13 °C