Structural Insights into the Ligand Binding Domain of the Glucocorticoid Receptor: A Molecular Dynamics Study

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Supporting Information

ABSTRACT: The glucocorticoid receptor (GR) is a ligand-binding dependent transcription factor that ultimately regulates vital biological processes and inflammation response through specific gene expression control, thus representing a notable drug target to explore. Structurally, its ligand binding domain (LBD) harbors the region for the ligand-dependent transcriptional activation function 2 (AF-2), a majorly hydrophobic groove formed by residues from helices H3, H4, and H12, where the H12 position plays a critical role in AF-2 spatial conformation and GR function as a whole. However, the exact mechanisms underlying how regulatory ligands control the H12 structure and dynamics are yet to be elucidated. In this work, we have explored the correlation between ligand identity and GR LBD H12 behavior through different molecular dynamics (MD) simulations. After building diverse GR LBD systems in agonist and nonagonist states, we studied each system’s response in the absence or the presence of an agonist ligand (dexamethasone) or an antagonist ligand (RU486) using classical MD simulations. We complemented them with steered MD for assessing the transition between those states and with the Umbrella Sampling method for free-energy evaluation. On the one hand, successfully obtaining fully folded nonagonist GR LBD states from the partially unfolded crystal GR LBD/RU486 underlines the role of the H1 in the GR LBD folding pathway. On the other hand, our results describe the H12 as a dynamic ensemble of conformations whose relative population is in the end determined by the interacting ligand: while dexamethasone privileges only a few poses (determined by a potential energy surface with a deep minimum), RU486 favors a wider H12 conformational amplitude, as indicated by a flatter potential landscape. By characterizing the H12 conformation in different conditions, we provide novel GR LBD models that represent potential targets for rational glucocorticoid drugs design, with the aim of accurately modulating GR activity.

1. INTRODUCTION

Growth, apoptosis, developmental behavior, and metabolism are among the biological features regulated by a group of steroid hormones named glucocorticoids (GCs). Given their powerful anti-inflammatory and immunosuppressive activities, they are administered as a treatment for asthma, dermatitis, rheumatoid arthritis, and other pathologies of autoimmune origin, while their proapoptotic nature extends GCs application to hematological neoplasms including various types of leukemias, lymphomas, and myelomas. This entitles GCs to be one of the most frequently prescribed medications worldwide. However, despite their evident benefits, GCs’ clinical use is often limited by their severe long-term use side effects, e.g., osteoporosis, diabetes, hypertension, and obesity. Hence, developing new steroids that can dissociate therapeutic effects from adverse effects appears both promising and challenging, since it demands unraveling the molecular mechanisms underlying GCs physiological and pharmacological actions.

The glucocorticoid receptor (GR) is a transcription factor classified within the nuclear receptor (NR) superfamily and depicted as a modular protein with three prominent and conserved domains: an N-terminal transactivation domain, a central DNA-binding domain (DBD), and a C-terminal ligand-binding domain (LBD). The N-terminal domain contains the AF-1 transcriptional activation domain required for transcriptional enhancement and association with basal transcription factors. The DBD is composed of two zinc finger motifs which play a critical role in receptor homodimerization and DNA binding. The LBD consists of 11 α-helices (H1; H3—H12) and four small β-strands folding into a globular structure, often described as a net, enclosing a central hydrophobic ligand-binding pocket (LBP, Figure 1). Besides, LBD also harbors a second dimerization interface and the domain for the ligand-dependent transcriptional activation function 2 (AF-2). Particularly, the AF-2 domain is a majorly hydrophobic groove formed by residues from helices H3, H4, and H12, where the H12 position plays a critical role in the AF-2 spatial context.

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conformation and GR function. Indeed, the AF-2 domain interacts with specific nonpolar residues present in particular coregulators’ amino acid motifs (LxxLL and I/LxxIL for coactivators and corepressors families respectively, where x is any amino acid) arranged on one side of their amphipathic helix (Figure 1).7

Hitherto, a central issue within the study of NRs is how regulatory ligands control the structure and dynamics of the H12.8,9 Initially, the analysis of the first crystallographic structures of isolated LBD led to propose the so-called induced-fit hypothesis—also known as the mouse-trap mechanism—to explain the molecular basis of action. This hypothesis postulates that, upon ligand binding, an inactive state of the receptor suffers a conformational change toward an agonist state. In the absence of a ligand (apo state), the H12 of several LBDs was described as a stable helix located away from the LBD body.10 However, further studies revealed that the H12 of apo-LBDs cannot be detached from the LBD surface and concluded that those observed positions were probably a consequence of artificial crystal packing interactions.11 Thus, in contrast to the induced-fit hypothesis, the conformational selection mechanism12 proposes that the receptor’s apo state comprises an ensemble of conformations in which the H12 swings along different well-folded positions, and where ligand-binding drives the relative population of each subensemble. In this sense, by using fluorine-19 (19F) nuclear magnetic resonance (NMR), Chrisman et al.13 recently proved that the H12 of the nuclear receptor peroxisome proliferator-activated receptor gamma (PPARγ) consists of a dynamic ensemble of conformations where the relative populations of subensembles correlate with the receptor’s activity. Its AF-2 domain, including the H12, exchanges relatively fast between many conformations for the apo-receptor. In turn, agonist and inverse-agonist binding consolidates the complex’s ensemble into structurally distinct active or inactive states, which favors coactivator or corepressor binding, respectively.

For the GR LBD, more than 40 crystallographic structures are archived in the Protein Data Bank (PDB, www.rcsb.org), mostly bound to ligands that stabilize the H12 in an agonist conformation.14 For nonagonist states, two structures of RU486 (Figure 1) complexes were resolved,15,16 providing information on the antagonistic mode of action of this antiglucocorticoid. In pdb:1nhz, the dimethylaniline group of RU486 appears to be preventing the H12 from adopting an agonistic conformation. However, this structure is incomplete and shows a fully deployed H12 bound to another GR LBD subunit. As described above, this is attributable to a crystallization technique artifact. In turn, the case of pdb:3h52 tetrameric structure is more informative: the receptor is observed with the H12 in three different positions (all consistent with nonagonist states) where some allow the union of a peptide corresponding to a corepressor motif. Unfortunately, lacking residues at the H11−H12 region and an incomplete folding of the H1 and H1−H3 loop in the four monomers of pdb:3h52 prevent its direct use for further molecular modeling studies.

During the last decades, computational modeling techniques have enriched the study of proteins and constitute a profitable tool that complements and expands experimental data.17,18 A prominent example is the case of molecular dynamics (MD) simulation that can render protein’s overall behavior in solution, thus allowing the study of protein inherent flexibility. Nevertheless, its robustness is limited in the presence of large energetic barriers between possible relevant conformations. To overcome this weakness, enhanced sampling methods have been developed, which expand and improve the understanding of the potential energy surface of a given system.19 In the case of NRs, MD simulation allowed for obtaining valuable

Figure 1. Left panel: three-dimensional structure of the GR LBD (pdb:1m2z, chain A) showing the H3, H4, and H12 helices forming the AF-2 domain which interacts with the NCoA2 peptide (in green). The position of the ligand binding pocket (LBP) is indicated as a cyan surface. A schematic of the secondary structure is shown below (numbers indicate first and final residues of each helix). Right panel: 2D and 3D structures of the glucocorticoid dexamethasone (dex) and the antiglucocorticoid mifepristone (RU486). The dimethylaniline moiety of RU486 is highlighted.
information regarding the ligand binding mode, the binding and unbinding pathways, and their molecular basis of action. Moreover, research on the dynamic association between the H12 position and the action of ligands bound to the LBP was recently conducted for other steroid receptors: progesterone, androgen, and estrogen receptors. However, to our knowledge, no studies were dedicated to thoroughly analyzing the conformational equilibria of the H12 in the GR LBD.

The above concerns encouraged us to explore through different MD simulation methods the correlation between ligand identity and the GR LBD H12 behavior. With this purpose, we constructed and studied several systems. The absence or the presence of an agonist ligand (dexamethasone) or an antagonist ligand (RU486) was combined with GR LBD both in agonist and nonagonist states. We envisaged that dissimilar starting conformations for the classical MD (cMD) analysis could offer a wider panorama on the H12 dynamics, while including a combination of steered MD (SMD) and the Umbrella Sampling (US) method would provide additional insights into the molecular determinants involved in the transition between those states.

2. RESULTS

2.1. Construction of GR LBDs in Nonagonist States.

The first aim of this work focused on the in silico construction of GR LBDs in nonagonist states, to obtain viable structures for MD simulation studies. As previously mentioned, monomers from pdb:3h52 have several remarkable H12 conformations, but inconveniently, their N-terminal end (H1 and H1−H3 loop) is unfolded from the LBD body and packaged in other monomers of the unit cell (Table 1 and Figure 2a). Although at first sight this phenomenon was associated with a crystallization artifact, a recent study by Suren et al. using single-molecule force spectroscopy showed

| PDB ID/chain | H1 and H1−H3 loop | missing residues | peptide coregulator | receptor state |
|--------------|-------------------|------------------|---------------------|---------------|
| 1m2z/A       | folded            |                  | NCoA2               | GR LBD^Ag    |
| 3h52/A       | unfolded          | 762−767 (H12 C-terminal loop) | NCoR1               | GR LBD^Ant1   |
| 3h52/B       | unfolded          |                  | NCoR1               | GR LBD^Ant2   |
| 3h52/C       | unfolded          | 738−754 (part of H11; H11−H12 loop; part of H12) | NCoR1               | GR LBD^Ant3   |
| 3h52/D       | unfolded          | 742−750 (H11−H12 loop) and 762−765 (H12) | NCoR1               | GR LBD^Ant4   |

Figure 2. a) Overall view of the GR LBD/RU486 tetramer (pdb:3h52): chain A in blue, B in green, C in magenta, and D in orange. H1 helices are schematized as cylinders, and NCoR1 peptides are shown in white. b) Superposition of chain A of pdb:1m2z (red) and chain B of pdb:3h52 (green). c) Absolute value of the difference between the Phi and Psi angles of the H1 and H1−H3 loop segment in pdb:1m2z (chain A) and pdb:3h52 (chain B). d) Contact map between residues from the H1 and H1−H3 loop and the remainder residues of the receptor for the equilibrated complexes. e) Main contacts between the H1 and H1−H3 loop and the remainder of the receptor. Cα of the involved residues are shown as spheres.
that the H1 folding, in fact, comprises the last step in the overall folding process and occurs only upon ligand binding. Thus, monomers of pdb:3h52 structure could presumably correspond to the open state described by these authors, constituting an intermediate state receptive to ligand binding. On this basis, we considered it feasible to obtain completely folded states of nonagonist systems starting from pdb:3h52 structures.

The structural alignment between pdb:1m2z and pdb:3h52 reveals a large congruence among most GR LBD helices (Figure 2b). Indeed, their conformation converges from residue 555 (H3) onward, except only for the H11−H12 region where the helices exhibit slightly different positions (see below). After comparing H1 and H1−H3 loop backbone dihedral angles Phi and Psi between the fully folded and the partially unfolded structures, it was noticed that only three angles at the end of the H1−H3 loop (PhiAsp554, PsiAsp554, and PhiSer555) diverged (Figure 2c). Apart from the pointed differences, no major changes were found along the rest of the loop or helices H1 and H3. Thus, after having established the structural similarity between both complexes, we obtained fully folded nonagonist state systems by replacing the coordinates of residues 528 to 554 in pdb:3h52 chains with those from the nonagonist state systems by replacing the coordinates of similarity between both complexes, we obtained fully folded H1 and H3. Thus, after having established the structural similarity between both complexes, we obtained fully folded nonagonist state systems by replacing the coordinates of residues 528 to 554 in pdb:3h52 chains with those from the corresponding residues of chain A in the pdb:1m2z structure. Each monomer from pdb:3h52 was then analyzed separately. In chain A (termed here as GR LBDAnt1), there were 6 missing residues following the H12 C-terminal loop (Table 1), which we added using the Modeller software. In chain B (termed here as GR LBDAnt2), the GR LBD was fully resolved. The structural alignment between both nonagonist GR LBD states here constructed (GR LBDAnt1 and GR LBDAnt2) and the agonist system (GR LBDAg) displays differences in the H12 position (Figure 3a). On the other hand, chain C (termed here as GR LBDAnt3) revealed a noteworthy H12 position displaced to the opposite side of the RU486 11β-substituent compared with the GR LBDAnt1 and GR LBDAnt2, partially occupying the AF-2 groove (Figure 3b). Unfortunately, this helix was poorly resolved, with as much as 17 residues missing along the H11 C-terminal end, the H11−H12 loop, and part of H12. We fruitlessly attempted to model this large portion of the receptor with Modeller, but no convincing structures were achieved. Lastly, chain D, with 13 unresolved residues along the H11−H12 loop and H12, showed a H12 configuration equivalent to the observed in chain A (GR LBDAnt1). Preliminary MD simulations from these two models showed a high structural instability, thus, in view of the above, neither chain C nor D was further considered in this study.

2.2. Initial Evaluation of Nonagonist GR LBDs. GR LBDAnt1/RU486 and GR LBDAnt2/RU486 complexes were energetically minimized and equilibrated through classical MD simulations. The Ramachandran plots of the resulting structures show that nearly all residues are present in the favored regions, few are present in the permitted ones, and only one is present in a forbidden region (Figure S1). Besides, the secondary structure of the modified section remaining comparable to that of GR LBDAg, the H1 rearrangement was notably stabilized in the designated position, and the H1−H3 loop did not suffer substantial structural deformations. Moreover, the analysis of the contacts between the N-terminal end and the remainder of the receptor revealed a high similarity between the original GR LBDAg and the constructed GR LBDAnt1 and GR LBDAnt2 systems (Figure 2d and e).

Finally, a proper H12 helical conformation was also conserved (see below).

2.3. Classical Molecular Dynamics Simulation of GR LBD Systems. Once the quality of the initial GR LBD models was evaluated, we simulated the systems GR LBDAg, GR LBDAnt1, and GR LBDAnt2 through 500 ns cMD to study the dynamic behavior under different conditions (Table 2), i.e., in the presence of the corresponding ligand (dex for GR LBDAg, RU486 for GR LBDAnt1 and GR LBDAnt2), in the absence of a ligand (apo systems), and in the presence of the opposite ligand (RU486 for GR LBDAg, dex for GR LBDAnt1 and GR LBDAnt2).

2.3.1. Stability and Fluctuation Pattern of cMD Trajectories. The stability of cMD trajectories was then assessed by monitoring backbone atoms’ RMSD relative to the initial structures. It was noticed that while during the first 200 ns the protein structure evidenced fluctuations, all GR LBDAg and GR LBDAnt2 systems were stabilized thereafter in constant values, which were less than 2.2 Å (Figure 4a). In turn, the GR LBDAnt1 systems suffered significant deviations all along the simulation, and only for the GR LBDAnt1/RU486 the trajectory seemed to converge. On the other hand, the RMSD of the H12 Ca atoms (Figure 4b) showed that from 200 ns onward the H12 position was stabilized in all systems, except for GR LBDAnt1/apo. Specifically, in systems GR LBDAg and GR LBDAnt1, the stabilized H12 was located close to the departing position when their original ligands were involved (dex and RU486, respectively). For GR LBDAg, the introduction of RU486 slightly deviated the H12, while the removal of the
ligand considerably deflected it. Likewise, in GR LBDAnt1, the absence of a ligand provoked a remarkable distortion of the H12 position. The RMSF calculated over the last 300 ns from the simulation (Figure S2) revealed, besides loops’ inherent mobility, some regions of secondary structure with meaningful differences among systems. Regarding the C-terminal end, the GR LBDAg/dex complex showed the least mobility, and the ligand’s substitution by RU486 clearly increased the fluctuation of the H11−H12 loop and the H12 first residues. A different RMSF profile was observed for the GR LBDAnt systems, with a large mobility in the C-terminal end of the H12. Among them, the GR LBDAnt/RU486 appeared the most rigid; GR LBDAnt/apo presented an increment in H12 mobility, and GR LBDAnt/dex had the H11−H12 loop fluctuation augmented. In turn, the RMSF pattern of GR LBD Ant2 complexes resembled those of the GR LBDAg systems, with some increased fluctuation for the GR LBD Ant2/dex H12 residues.

2.3.2. Evolution of the H12 Conformation in the cMD Trajectories. A detailed RMSF global analysis highlighted the great stability of helices H3 and H10−H11 along the cMD, particularly of the H3 C-terminal end (residues 568 to 579; hereinafter, portion “B” or H3B) and the H10−H11 N-terminal residues 711 to 721 (hereinafter, H10). This peculiarity allowed us to select these virtually invariant segments as a spatial reference to analyze the mobility of the H12 over time. Three vectors of equal direction than segments H3B, H10, and helix H12 were defined, and the resulting angles were calculated along the complete cMD. In accordance with the cited RMSD analysis, the temporal evolution of these angles evidenced that all systems converged after the 200 ns (Figure S3) with the exception of GR LBD Ant1/apo that presented great instability at the H3B−H12 angle.

For visualization purposes, the last 300 ns of each cMD trajectory were then examined using box plots (Figure 5). As expected, the H3B−H10 angle was mainly conserved among the 9 systems, particularly within GR LBDAg and GR LBDAnt2 complexes. Conversely, angles H3B−H12 and H10−H12 displayed a more condition-dependent behavior. This was particularly evident for the GR LBDAnt1 systems, where large changes were observed in both angles when RU486 was removed from the LBP or replaced by dex. Although less pronounced, the GR LBDAnt2 systems also showed certain differences in the evolution of those angles. Lastly, for GR LBDAg systems, both angles suffered only narrow deviations, and solely the imposed presence of RU486 markedly disturbed angle H3B−H12.

For further characterization of the stabilized H12 conformation, we calculated the spatial distribution of the angles H3B−H12 and H10−H12 over the last 300 ns of the simulation and plotted them as 2D histograms (Figure 6a). There, different H12 states achieved by each system are clearly displayed. In the case of the GR LBDAg, there was a tight distribution of H3B−H12 and H10−H12 angles, around 75° and 98°, respectively, corresponding to the GR LBDAg/dex state. H12 disposition was not perceptibly altered after dex removal, but its substitution by RU486 slightly withdrew the receptor from the agonist state. However, the H12 did not turn toward any of the positions observed for complexes GR LBDAnt1 or GR LBDAnt2 but in the opposite direction, interestingly resembling that of the GR LBD Ant3 state (cf. Figure 3b). This led us to evaluate GR LBDAg/RU486 possible additional conformational changes. We extended its trajectory

Table 2. Simulated Systems and Average Distances between “Charge Clamp” Residues

| system | receptor | ligand   | Glu755-Lys579 | Glu755-Asp590 | Asp590-Lys579 |
|--------|----------|----------|---------------|---------------|---------------|
| 1      | GR LBD Ag | Dex      | 18.4          | 15.0          | 11.8          |
| 2      | GR LBD Ag | none     | 18.3          | 15.0          | 11.7          |
| 3      | GR LBD Ag | RU486    | 17.8          | 12.9          | 11.9          |
| 4      | GR LBD Ant1 | RU486 | 22.2          | 21.6          | 12.0          |
| 5      | GR LBD Ant1 | none | 21.5          | 20.4          | 11.7          |
| 6      | GR LBD Ant1 | Dex | 20.5          | 20.2          | 11.9          |
| 7      | GR LBD Ant2 | RU486 | 20.1          | 16.6          | 11.6          |
| 8      | GR LBD Ant2 | none | 18.5          | 15.5          | 11.4          |
| 9      | GR LBD Ant2 | Dex | 18.4          | 15.2          | 11.4          |

“Distance between the Cα of the indicated residues.
up to 1000 ns and compared the resulting angle distribution with GR LBD\(^{\text{Ag}}\)/dex and LBD\(^{\text{Ag}}\)/RU486 300−500 ns (Figure S4). It was then confirmed that the H12 sustained its drift in the mentioned direction, partially occupying the AF-2 groove. This emerging state was termed pseudo-GR LBDAnt\(^{3}\) and incorporated within our pool of systems.

For the GR LBDAnt\(^{1}\)/RU486 complex, histograms showed a tight angle distribution at 30° for H3B−H12 and 122° for H10−H12 angles (Figure 6a). By contrast, for the apo and dex systems, the angle distribution approaches that of GR LBDAnt\(^{2}\)/RU486. Thus, this H12 conformation presumably describes an intermediate state between the antagonist state GR LBDAnt\(^{1}\)/RU486 and the agonist state GR LBDAg/dex. In this sense, GR LBDAnt\(^{2}\)/apo and GR LBDAnt\(^{2}\)/dex histograms were remarkably similar to that of GR LBDAg/dex and manifested how the intermediate state evolves toward the agonist state when RU486 is removed or replaced by dex.

A schematic diagram depicting the evolution of each simulated system is shown in Figure 6b. The complexes directly derived from the crystallized structures conserve the H12 in a position similar to the original one, while variations in LBP condition modify the H12 destination during the MD. Figure 7 shows representative snapshots of the main configurations observed in the H10−H12 and H3B−H12 angle configurational space. It can be observed that while the C-terminal half of the H12 occupies a similar position in all the systems, its N-terminal half explores different states swinging from virtually blocking the AF-2 domain to releasing and enlarging it.

2.3.3. Dimension of the AF-2 Domain. To estimate how H12 conformational changes influence the AF-2 configuration, we calculated the average distance among Cα atoms residues Glu755 (H12), Lys579 (H3), and Asp590 (H4) (Table 2 and Figure 8). These residues, localized within each of the AF-2 domain conforming helices, participate in anchoring the coregulator to the receptor through polar interactions (“charge clamps”). Hence, the relative position between these three residues determines the accessibility through the opening dimension of the AF-2. As expected, the Lys579-Asp590 distance remained equal in all simulated systems, indicating the permanence in the relative disposition of this coregulator’s anchor point. In contrast, large variations characterized the distances involving the H12 residue. Differences of up to 4.4 and 7.5 Å were calculated for the Lys579-Glu755 and Asp590-Glu755 pairs, respectively, with GR LBD\(^{\text{Ag}}\)/RU486 and GR LBDAnt\(^{1}\)/RU486 at the extremes. These findings evidence that the AF-2 dimensions and interactions ultimately depend on H12 spatial configuration.

2.3.4. Ligand Binding Mode. A detailed analysis of the ligand binding mode was pursued calculating hydrogen bond interactions between the polar moieties of dex or RU486 and the polar residues of the GR LBP. Figure S5 displays the hydrogen-bond occupancy (HBO) between ligand and receptor atoms for the last 300 ns of the trajectories. Globally, in the 6 holo-systems these ligand−receptor interactions occur in well characterized zones, i.e., the C-3 carbonyl group and the 17-hydroxyl group; in addition, dex presents long-lasting hydrogen bonds through its 11- and 21-hydroxyl groups. The C-3 carbonyl group of both ligands participates in a hydrogen bonding network that involves Gln570 (H3) and Arg611 (H5). In the case of dex, while for the GR LBDAg/dex and GR LBD Ant1/dex complexes the interaction is mainly mediated by a water molecule (explaining in part the small HBO values for the Arg611), larger direct ligand−receptor contacts are observed for the GR LBDAnt2/dex complex. For RU486 systems, the agonist state presents similar values for those interactions, opposite to the nonagonist states which show little interaction with Gln570 but a more sustained one with Arg611.

Regarding the 17-hydroxyl group, a persistent hydrogen bond is formed with Gln642 (H7) in the GR LBDAnt1/dex and, to some extension, the GR LBD Ag/RU486 systems. It is
noticeable that the crystal-derived structures only maintain sporadic interaction with this residue, suggesting that those contacts within the mentioned complexes are forced due to the size-occupancy relationship of the LBP and each ligand: a smaller agonist LBP would shift RU486 orientation toward Gln642, while a larger LBP conformation would prevent dex from interacting through its 11- and 21-hydroxyl groups (see below), thus favoring this bond.

For dex 11- and 21-hydroxyl groups, large HBO values are detected in the agonist GR LBDAg state, particularly between the couples 11-hydroxyl/Asn564 (H3) and 21-hydroxyl/Thr739 (H11). Although less frequently, these are also present in the GR LBDAnt1/RU486 complex; on the contrary, as mentioned before, no interactions between these polar groups and LBP residues are formed in the antagonist GR LBDAnt1 state. These findings suggest that variations in the H12 position are related with variations in the mode in which 11- and 21-hydroxyl are recognized by the LBP, plus strengthening the notion that the GR LBDAnt resembles an intermediate state between GR LBDAg and GR LBDAnt.

On the other hand, we identified the contact residues for the RU486 dimethylaniline moiety along each GR LBD/RU486 simulation (Figure S6). In the three systems, this voluminous group contacts with Gly567 and Gly568 of the H3 and Trp600 localized between the H4 and H5. The third contact point, however, differs for each complex: Phe749 for GR LBDAg/RU486, Leu753 for the GR LBDAnt1/RU486, and Ile756 for the GR LBDAnt2/RU486. Remarkably, the last two belong to the H12, while the first is located within the H11−H12 loop, correlating the contacts established by H12 residues with the RU486 dimethylaniline moiety and the conformation adopted by the H12.

On the whole, it is noteworthy that all systems bond with at least one residue of the H3 chain (which has been earlier proven to be practically invariant), while other interactions are ultimately related to the GR LBD (and particularly H12).
configuration. This becomes more evident when ligands are switched. On the other hand, the number of interactions is positively correlated with the stability observed in the MDs. Indeed, systems with larger interactions evidenced less fluctuations than those with few contacts along their trajectories, with GR LBD\textsuperscript{LBD\textsuperscript{apo}}/dex representing the most stable complex.

2.4. Steered Molecular Dynamics Simulation. Through cMD simulations we have found that the H12 can adopt numerous positions depending on LBP conditions (i.e., ligand identity and occupancy), with the GR LBD\textsuperscript{apo}/RU486 (or pseudo-GR LBD\textsuperscript{Ant3}) at one extreme of the conformational space and the GR LBD\textsuperscript{Ant1}/RU486 at the other. Considering the already mentioned limitations of cMD, we alternatively employed steered MD simulations (SMD) to explore putative transitions between those two states. In SMD, an external force is applied to drive the system along a selected reaction coordinate or “collective variable” (CV) at a constant velocity.26 The election of an adequate CV capable to describe the pathway between two states of the protein is mandatory and usually very challenging.

At first, considering our cMD results, angles H12–H3B and H12–H11A were taken as CV to displace the H12 in the absence of ligand, but trajectories proceeded with considerable deformation in several regions of the receptor. Then, we tested a combination of distances between Cα atoms of key residues from H3, H11, and H12, also obtaining inadequate pathways. Other geometrical descriptors, such as combinations of distances and angles of the main variable residues, were equally assessed, with no proper results. Finally, when using the RMSD of H3, H11, and H12 Cα residues as CV, we obtained a smooth path between the extreme states without constitutional deformations and with well conserved secondary structure elements. Figure 9a shows the temporal evolution of the CV, and Figure 9b shows the superposition of snapshots extracted at regular intervals from the 100 ns SMD trajectory. As it can be observed, along the transition from the pseudo-GR LBD\textsuperscript{Ant3} to the GR LBD\textsuperscript{Ant1}, the H12 visits multiple intermediary conformations, notably including those corresponding to the GR LBD\textsuperscript{apo} and GR LBD\textsuperscript{Ant2} original states.

SMD trajectories were also simulated for dex and RU486 complexes, finding a similar behavior to the apo system, with straightforward pathways in which both the GR LBD integrity and secondary structure were conserved. All SMD trajectories were also analyzed by monitoring the temporal evolution of the H3B–H12 and H10–H12 angles (Figure S7), finding only some minor discrepancies among all systems. For example, the H3B–H12 angle values in the first half of the simulation were larger in the RU486 complex than in the apo and dex systems, as was expected due to the bulky substituent of the antiglucocorticoid. On the other hand, the H10–H12 angle showed differences between dex and RU486 when the GR LBD\textsuperscript{Ant2} state was reached. As a control, SMD trajectories were also carried out in the reverse direction (from the GR LBD\textsuperscript{Ant1} to the pseudo-GR LBD\textsuperscript{Ant3} states) in the three conditions (apo, dex, and RU486), finding a very similar behavior to that described in the direct direction (data not shown).

In summary, the one-dimensional path coordinate RMSD successfully illustrates a complete dynamic picture of the conformational space sampled by the H12 in the GR LBD. Hence, this CV was then employed to obtain an energetic profile of the configurational space.

2.5. Umbrella Sampling Analysis. After validating a suitable CV for our systems, we performed an Umbrella Sampling (US) analysis27 using the RMSD of H3, H11, and H12 Cα atoms as CV. The three SMD trajectories (apo, dex, and RU486) were sampled every 0.2 Å distance along the CV, obtaining 25 snapshots for each system. These were then equilibrated and simulated for 3 ns. The last 2 ns of each window were integrated with the Umbrella Integration method.28

The free energy obtained through US was then plotted as a function of the CV, showing a clear difference between GR LBD dex and GR LBD RU486 systems (Figure 9c). Indeed, in the presence of dex, a deep minimum is described at CV = 3.8 Å, which corresponds to the H12 position in the agonistic conformation (GR LBD\textsuperscript{apo} original state). Large energetically barriers are observed when the H12 is displaced toward both GR LBD\textsuperscript{Ant2} and pseudo-GR LBD\textsuperscript{Ant3} H12 positions, indicating that this ligand anchors the H12 in a well-defined and stable position. This finding sustains the large rigidity observed in the cMD simulations and explains the full agonistic activity of this ligand.

In contrast, RU486 provides a flatter free-energy curve, in accordance with the wider flexibility exhibited by the RU486 systems in cMD simulations. A broad global minimum is observed at CV = 5 Å, concordant with the pseudo-GR LBD\textsuperscript{Ant3} H12 conformation. In turn, GR LBD\textsuperscript{Ant3} and GR LBD\textsuperscript{Ant2} H12 positions do not correlate with any noticeable minima but to plain regions of the free energy surface. The GR LBD/ apo system also provides a flattened free energy curve with a slight global minimum at CV = ∼4 Å.

3. CONCLUSION
The GR LBD conform to a compact globular structure whose end-terminal helices play a crucial role in the activity of this
receptor’s domain. Thus, following an approach of concatenated MD simulations of different types, we focused on deciphering the correlation between ligand identity and the GR LBD H12 dynamic behavior. Even when only two commercial ligands were evaluated, we consider performing SMD simulations followed by US analysis a powerful strategy and a potential tool to satisfactorily understand the molecular basis of action of glucocorticoids with different activity profiles.

Several GR LBD systems had to be generated starting from the structure of the nonagonist GR LBD/RU486 complex (pdb:3hs52). This complex crystallized as a tetramer with the H1 unfolded in the four monomers (resembling the H1 situation described for the native GR LBD apo)\(^25\) and required an in silico refolding in all cases. The straightforward extension of stable closed structures upon simulation supports the idea that the interaction between monomers during the GR LBD/RU486 crystallization process prevented the H1 from adopting its complete final shape. Previous studies assessing the ligand’s access/release to the receptor have mostly spotlighted the H6–H7 plasticity as an integral part of the ligand-entry mechanism.\(^3\) Our findings, in turn, lead us to consider the GR LBD/RU486 unfolding H1, far from being a crystallization artifact, a determinant in ligand association and dissociation from the GR and a very auspicious element in the understanding of this process.

A detailed depiction of how two paradigmatic glucocorticoids affect the H12 configurational space is also presented. Moreover, a dynamic GR LBD H12 capable of exploring a wide range of clearly identified and characterized conformations is described. Our results reveal pronounced H12 reorientations depending on the LBP conditions, while the remainder of the GR LBD entails only modest changes in the N-terminal H3 and the C-terminal H11. Together, these findings fit within the conformational-selection model in explaining GR agonism and antagonism.

Potential surface analysis delineated a deep minimum for the GR LBD dex system, indicating that among all possible conformations the agonist dex privileges only a few poses, providing high rigidity to the receptor–ligand complex. As a consequence, the AF-2 domain remains virtually fixed, suitable for recognizing coactivators’ LxxLL motifs, as present in NCoA\(_\alpha\). Alternatively, in the presence of the antagonist RU486, the H12 conformational amplitude appears to be wider, as indicated by a flatter potential surface energy with no prominent minimum. Antagonist complexes’ larger flexibility in comparison to agonist systems has also been described for the androgen\(^23\) and the estrogen receptors.\(^24\) Besides, RU486 would also withdraw the H12 from the agonist state and place it on either side of the ligand’s bulky substituent. From these observations, we propose that in the absence of coregulators, the H12 lowest-energy state would be reached when this helix directly occupies the AF-2 groove. In turn, the presence of corepressors would stabilize conformations that extend the AF-2 dimensions. Therefore, H12 conformational selection is a dynamic equilibrium ultimately determined by both the identity of the ligand and the presence of corepressors.

Finally, by characterizing novel H12 putative conformations different from those already crystallized, we provide a whole series of GR LBD systems that represent potential targets for the rational design of new glucocorticoid drugs, with the ultimate aim of finely modulating the GR activity.

4. COMPUTATIONAL METHODS

4.1. Initial GR LBD Structures. Five GR LBD structures were initially built for the MD simulations (Table 1). The GR LBD\(^\alpha\) coordinates were taken from the crystal structure of the GR LBD/dex complex (residues 528 to 777 of chain A from pdb:1m2z). GR LBD\(^\alpha1\) and GR LBD\(^\alpha2\), and GR LBD\(^\alpha3\) were constructed by combining coordinates from two crystal structures: residues 528 to 554 of chain A of the GR LBD/dex complex (pdb:1m2z) and residues 555 to 777 of chain A, B, or C of the crystal structure of the GR LBD/RU486 complex (pdb:3hs52). The missing regions in GR LBD\(^\alpha1\) and GR LBD\(^\alpha3\) (Table 1) were modeled with the Modeller software.\(^30\) In all cases, the introduced mutations were reverted to retrieve the wild-type GR LBD sequence. Using the GR LBD\(^\beta\), GR LBD\(^\alpha1\), and GR LBD\(^\beta2\) coordinates, 9 systems (3 apo and 6 holo) were prepared for classical MD simulations (Table 2). For the GR LBD\(^\beta\)/dex, GR LBD\(^\alpha1\)/RU486, and GR LBD\(^\beta2\)/RU486, the crystal coordinates of the ligands were conserved. Ligand’s coordinates were directly removed to obtain apo systems, while for the cross-systems GR LBD\(^\beta\)/RU486, GR LBD\(^\alpha1\)/dex, and GR LBD\(^\beta2\)/dex, the HF/6-31G** optimized structures of the steroids were introduced into the GR LBD models by superimposing the skeleton carbon atoms of C and D rings with the corresponding atoms of the crystallized steroid. For the force field parameters of the ligands, RESP (restraint electrostatic potential) atomic partial charges were computed using the HF/6-31G** method in the quantum chemistry program Gaussian 09\(^31\) on the corresponding optimized structures.

4.2. Classical Molecular Dynamics Simulation. Molecular dynamics were performed with the AMBER 18 software package.\(^32\) Ligand parameters were assigned according to the general AMBER force field (G Aff) and the corresponding RESP charges using Antechamber. The FF14SB force field parameters were used for receptor residues. Complexes were immersed in an octahedral box of TIP3P water molecules using the Tleap module, giving final systems of about 35,000 atoms. Systems were initially optimized and then gradually heated up to a final temperature of 300 K. All MD production runs of 500 ns were performed at 300 K in a NPT ensemble using periodic boundary conditions and the particle mesh Ewald method (grid spacing of 1 Å) for treating long-range electrostatic interactions with a uniform neutralizing plasma. Temperature regulation was done by using a Langevin thermostat with a collision frequency of 2 ps–1. The SHAKE algorithm was used to keep bonds involving H atoms at their equilibrium length, allowing the use of a 2 fs time step for the integration of Newton’s equations.

4.3. Steered Molecular Dynamics Simulation. The RMSD of C\(_\alpha\) atoms of H3 (556 to 579), H10–H11 (711 to 740), and H12 (751 to 766) residues was employed as a collective variable (CV) as it is implemented in the Amber package.\(^33\) The cMD-equilibrated GR LBD\(^\alpha1\)/apo system was used as a reference (CV = 0 Å) to drive the cMD-equilibrated pseudo-GR LBD\(^\alpha1\)/apo (CV = 5.1 Å) with a force harmonic constant of 300 kcal/mol Å\(^2\) in a 100 ns long SMD trajectory. A similar strategy was used to obtain the GR LBD/dex and GR LBD/RU486 SMD trajectories.

4.4. Umbrella Sampling. Twenty-five snapshots with equidistant CV 0.2 Å apart were extracted from the SMD trajectories and prepared for Umbrella Sampling simulations using parameters as for CMD. Production runs of 3 ns were
performed as for cMD. Only the last 2 ns were integrated through the Umbrella Integration method.28

4.5. Analysis of Results. Trajectories were analyzed with CPPTRAJ,34 and representative snapshots were visualized with VMD.35 The root mean square fluctuations (RMSF) of Cα receptor atoms and boxplots were calculated over the last 300 ns of the cMD trajectories. The angle between two α-helices was computed using a vector for each helix. The vector of an α-helix was defined by two points determined by the center of mass of four Cα atoms. Residues 568–571 and 576–579 were chosen for the starting and ending points of the H3B vector, residues 712–715 and 718–721 were chosen for the starting and ending points of the H10 vector, and residues 751–754 and 761–764 were chosen for the starting and ending points of the H12 vector. Graphical representations of α-helices were obtained with Bendix.36 Ramachandran plots were obtained with the RAMPAGE server.37 Hydrogen bond occupancy (HBO) for each interaction was calculated with the hbond command of CPPTRAJ using the default parameters for the distance cutoff (acceptor to donor heavy atom less than 3.0 Å) and for the angle cutoff (135°).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jcim.9b00776.

Figure S1: Ramachandran plots of equilibrated GR LBDapo/RU486 and GR LBDmut2/RU486 complexes. Figure S2: Root mean square fluctuation (RMSF) measured over last 300 ns of cMD simulations. Figure S3: Time evolution of angle confined between H3B–H12, H10–H11, and H3B–H10 helices. Figure S4: Distribution of H3B–H12 and H10–H12 angles. Figure S5: Comparison of dex and RU486 ligand binding mode: schematic representation of polar interactions between ligands and LBP of GR LBD. Figure S6: Residues contacting dimethylamino moiety of RU486 in GR LBD/RU486 systems. Figure S7: Time evolution of H3B–H12 and H10–H12 angles for GR LBD/dex, GR LBD/ apo, and GR LBD/RU486 systems along SMD simulations (PDF)

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Notes

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