The Use of in Vitro Peptide Binding Profiles and in Silico Ligand-Receptor Interaction Profiles to Describe Ligand-Induced Conformations of the Retinoid X Receptor $\alpha$ Ligand-Binding Domain

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It is hypothesized that different ligand-induced conformational changes can explain the different interactions of nuclear receptors with regulatory proteins, resulting in specific biological activities. Understanding the mechanism of how ligands regulate cofactor interaction facilitates drug design. To investigate these ligand-induced conformational changes at the surface of proteins, we performed a time-resolved fluorescence resonance energy transfer assay with 52 different cofactor peptides measuring the ligand-induced cofactor recruitment to the retinoid X receptor-$\alpha$ (RXR$\alpha$) in the presence of 11 compounds. Simultaneously we analyzed the binding modes of these compounds by molecular docking. An automated method converted the complex three-dimensional data of ligand-protein interactions into two-dimensional fingerprints, the so-called ligand-receptor interaction profiles. For a subset of compounds the conformational changes at the surface, as measured by peptide recruitment, correlate well with the calculated binding modes, suggesting that clustering of ligand-receptor interaction profiles is a very useful tool to discriminate compounds that may induce different conformations and possibly different effects in a cellular environment. In addition, we successfully combined ligand-receptor interaction profiles and peptide recruitment data to reveal structural elements that are possibly involved in the ligand-induced conformations. Interestingly, we could predict a possible binding mode of LG100754, a homodimer antagonist that showed no effect on peptide recruitment. Finally, the extensive analysis of the peptide recruitment profiles provided novel insight in the potential cellular effect of the compound; for the first time, we showed that in addition to the induction of coactivator peptide binding, all well-known RXR$\alpha$ agonists also induce binding of corepressor peptides to RXR$\alpha$.

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Abbreviations: AF, Activation function; ARES, automatic residue extraction system; 9-cis RA, 9-cis retinoic acid; 3D, three-dimensional; DHA, docosa hexaenoic acid; ER, estrogen receptor; GST, glutathione S-transferase; H12, helix 12; LBD, ligand binding domain; LBP, ligand binding pocket; MI, modulation index; MD, molecular dynamics; NCoR, nuclear receptor corepressor; NR, nuclear receptor; PDA, pentadecanoic acid; RAR, retinoic acid receptor; RIP, receptor-interacting protein; RXR, retinoid X receptor; SMRT, silencing mediator of retinoid and thyroid hormone receptor; SNRM, selective nuclear receptor modulator; SRC, steroid receptor coactivator; TR-FRET, time-resolved fluorescence resonance energy transfer; TTNPB, 4-[(E)-2-(5,6,7,8-tetrahydro-5,5,8,8-tetramethyl-2-napthalenyl-1-propenyl) benzoic acid.

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contains a LXXLL sequence motif (NR box) (7–9). In contrast, an antagonist displaces H12, resulting in the dissociation of coactivators and the association of corepressors. Corepressors bind into the cofactor binding groove similarly to coactivators, but with a LXX(I/H)I/XXX(I/L) sequence motif (10–12). The only available crystal structure with a corepressor shows that the corepressor peptide position is slightly tilted so that it partially overlaps with the agonistic location of H12 (10). Subsequently, the receptor-cofactor complex binds as a homodimer, heterodimer, or monomer to NR-specific response elements in the promoter region of target genes, which results in up- or down-regulation of those genes.

The above described mechanism suggests that agonists and antagonists induce two distinct conformational states of the LBD, i.e. stabilizing H12 in the agonist or antagonist position, respectively. These distinct conformations lead to the interaction with coactivators or corepressors and result in two distinct patterns of gene expression. However, it was observed that different agonists induce different gene expression profiles (13, 14). This may suggest that H12 does not have two distinct positions but can take any preferential position between the distinct agonist position and antagonist position. It is the combination of both compound and cofactor that determines the position of H12 and therefore the effect on gene expression. This suggests the existence of a more subtle communication pathway between the LBP and the cofactor binding groove of the NR. It has been shown for the estrogen receptor (ER) α and the peroxisome proliferator-activated receptor-γ (13, 15) that the binding of different ligands results in distinct patterns of cofactor binding. The cofactors in those studies are mimicked by helical peptides of 20–25 amino acids, containing the LXXLL coactivator motif or the LXX(I/H)I/XXX(I/L) corepressor motif. These different patterns of peptide binding, so-called peptide recruitment profiles, are most likely caused by different conformations at the NR surface, in particular the cofactor binding groove. These studies show that peptide recruitment profiles are very useful to probe the conformation at the NR surface and to study the position of H12 in the presence of ligand and cofactor peptides. Moreover, compounds can be clustered according to their similarity in peptide recruitment profiles. It is believed that compounds in one peptide recruitment cluster induce similar conformations at the NR surface, which may result in transcription of the same set of target genes. From a structure-based drug design perspective it is therefore important to know which interactions between ligand and receptor cause a specific NR surface conformation, i.e. peptide recruitment profile. In other words, molecular understanding of the communication between LBP and cofactor binding groove may be one step further toward the design of NR drugs with a certain gene transcription profile.

In this paper we describe a methodology that helps to obtain this molecular understanding of ligand-induced conformations of the NR surface. We first measured peptide recruitment profiles for a set of different ligands. Second, we studied whether these different ligands actually have distinct binding modes using the available X-ray structures and docking. The different binding modes of compounds were represented by so-called ligand-receptor interaction profiles, thereby converting complex three dimensional (3D) data into two-dimensional fingerprints. These ligand-receptor interaction profiles were used to cluster compounds that bind in a similar manner. Finally we compared the clusters of the peptide recruitment profiles with the clusters of the ligand-receptor interaction profiles to reveal residue positions that are involved in signaling between the LBP and the surface of the LBD.

As a representative of the NRs we used the retinoid X receptor (RXR) α. RXRs (16) are unique within the NR superfamily because they form heterodimers with many other NRs and also function autonomously, as homodimers (17). The receptors are involved in important processes such as the regulation of carbohydrate and lipid metabolism, cell differentiation, proliferation, and morphogenesis. RXRs are regulated by retinoids, which are derivatives of vitamin A. The natural ligand of these receptors is 9-cis retinoic acid (9-cis RA) (18). Another class of RXR ligands comprises fatty acids, e.g. oleic acid, docosa hexaenoic acid (DHA), and phytanic acid (19–21). Finally, there are many synthetic ligands known for RXR (22), some of which are currently available as drugs for the treatment of (skin) cancers and dermatological diseases, such as psoriasis and acne.

In this study the induced peptide recruitment profiles of 10 well-described RXR ligands [and one retinoic acid receptor (RAR) ligand, TTNPB] were determined using a time-resolved fluorescence resonance energy transfer (TR-FRET) assay with 52 peptides. Simultaneously, the binding modes of these ligands in the pocket of RXRα were determined by modeling and molecular dynamics (MD) simulations. Clustering of the peptide recruitment profiles revealed that the compounds induced three distinct conformations at the surface of the LBD. Clustering of the binding modes showed two distinct modes of ligand binding for the RXR compounds. We successfully combined ligand binding profiles and peptide recruitment profiles to reveal structural determinants of signaling from the LBP to the surface of the LBD.

RESULTS

To investigate whether different RXR ligands induce different conformations at the RXRα LBD surface, we performed a peptide recruitment assay with 52 peptides (10⁻⁷ M) and 11 different ligands (10⁻⁵ M). The resulting peptide recruitment profiles are depicted in Fig. 1. Figure 1A shows the raw fluorescence intensity data. This figure shows that approximately half of the
Fig. 1. Peptide Recruitment Profiles of RXRα LBD Induced by Different Ligands
A, Binding of 52 biotinylated peptides (0.1 μM) to GST-labeled hRXRα LBD (10 nM) in the absence (dotted line) and presence of 10 μM 9-cis RA (solid line). The SEM of two separate experiments, each performed in duplicate, is indicated by the error bars.
B, Peptide recruitment profiles of 11 RXR ligands: the association and dissociation of 52 peptides to hRXRα LBD is represented as log(MI) values. The modulation index (MI) was obtained by dividing the fluorescence intensity in the presence of ligand by the intensity in the absence of ligand. Each line represents the average peptide recruitment profile for one of the 11 compounds. The SEM for each profile is given in supplemental Fig. 1 published as supplemental data on The Endocrine Society’s Online Journals web site. The average peptide recruitment profile of 9-cis RA is highlighted as a black solid line. Peptides have been divided in coactivators or corepressors on the basis of their sequence motif.
peptides already bind to the RXRα surface in the absence of ligand (dotted line). The majority of these peptides are derived from coactivators. This suggests that H12 is stabilized in the agonist position in the absence of ligand, which is also observed in several NR LBD apo crystal structures (e.g., Refs. 23–27). Most likely, these peptides bind so well in the cofactor binding groove that they stabilize H12 in the agonist position (and vice versa). Figure 1A also shows the peptide recruitment profile in the presence of 9-cis RA, the natural ligand of RXRα (solid line). For many peptides, the fluorescence intensity, which is a measure of the peptide binding to the LBD, is higher in the presence of 9-cis RA. This indicates that the affinity of these peptides is increased due to the binding of 9-cis RA in the LBP.

Because the basal ligand-independent signal varies between different peptides, it is difficult to easily compare the effect of the ligand. For this purpose, the log\[modulation index (MI)\] was calculated (see TR-FRET Assay in Materials and Methods for more details), and the resulting peptide recruitment profile of 9-cis RA is depicted in Fig. 1B (black line). This representation directly shows whether a compound is associative or dissociative. Values above zero indicate recruitment of peptides, and thus an associative effect of the ligand on peptide binding. In contrast, values below zero indicate dissociation of peptides and therefore a dissociative effect of the compound on peptide binding. A log(MI) value of approximately zero means that the peptide does not bind or that the binding of a peptide is hardly changed by the ligand.

Peptides that Do Not Bind in the Absence and Presence of 9-cis RA

The peptide recruitment profile of 9-cis RA in Fig. 1B (dark line) shows that 25 peptides have a log(MI) value of approximately zero (log(MI) < -0.1). Of these 25 peptides, 12 also do not bind in the absence of 9-cis RA (Fig. 1A), indicating that these peptides are not compatible with the RXRα cofactor binding groove. These 12 peptides include six peptides with a corepressor motif (NcoR_1, NCoR_2, BL29, HR_1, HR_2, BT_1) and six peptides with a coactivator motif (SRC3_4, ASC2_2, ARA70, Ppt4-1, LXRα_H12, and FHL2). A sequence alignment of the six nonbinding corepressor peptides revealed no sequence similarity that explains why these peptides do not bind (data not shown). The alignment of the six nonbinding coactivator peptides (Fig. 2A) showed that three coactivator peptides do not have a LXXLL motif but a FXXL(F/Y) motif. Several studies showed that the FXXL(F/Y) motif is preferred by the androgen receptor to interact with coactivators, whereas other NRs show no affinity for peptides with this motif (28-30). Moreover, all six peptides possess a polar residue at the −1 position relative to the coactivator motif (LXXLL). This observation agrees with previous studies, which showed that a hydrophobic residue is preferred at this position to facilitate proper coactivator binding to various NR LBDs (31, 32). This hydrophobic residue at the −1 position is locked in an aromatic region of RXRα that is induced by the peptide (33). A polar residue at the −1 position is therefore unfavorable.

Peptides that Bind in the Absence of Ligand but Are Not Inducible by 9-cis RA

Figure 1, A and B, shows that the remaining 13 peptides equally bind to RXRα with and without 9-cis RA. These data suggest that these peptides bind so strongly to the RXRα LBD that this interaction cannot be further enhanced by 9-cis RA. This is confirmed by the dose-response curves of one of these peptides [receptor interacting protein (RIP)140_3], shown in Fig. 3A. The estimated Kₐ values of the dose-response curves in the absence and presence of 9-cis RA are nearly equal (152 nM in the absence and 155 nM in the presence of compound), which indicates that there is no effect on the peptide binding by 9-cis RA.

Figure 2B shows the sequence alignment of these 13 strong binding peptides. Except RIP140_3, all peptides contain both the LXXLL motif and the favorable hydrophobic residue at the −1 position (31–33). Obviously, this is one of the essential structural properties that coactivator peptides need: to bind so strongly to the LBD that binding of 9-cis RA is no longer required. RIP140_3 is the only peptide without the hydrophobic residue at the −1 position. This suggests that other residues in the coactivator peptide compensate for the absence of the hydrophobic residue. One possible explanation could be the lysine at position 2 that may have a favorable electrostatic interaction with aspartate 295 in the coactivator binding groove. This is corroborated by the observation that the four top-ranked peptides (in the absence of ligand) all contain a positive charge or H-bond donor at position 2 in the coactivator motif.

Peptides Whose Binding Is Inducible by 9-cis RA

Finally, there is a set of 17 peptides that weakly or not bind in the absence of ligand and bind (log(MI) > 0.30, i.e., 2-fold increase in fluorescence intensity) significantly better in the presence of 9-cis RA. Some of these peptides have been described in previous studies to interact with RXRα. For example, our observation that RIP140_7 is significantly enhanced in binding to the receptor upon addition of 9-cis RA agrees well with a study of Farooqui et al. (34). Also the observation that the interaction of SHP_2 with the LBD is enhanced with 9-cis RA is in agreement with previous studies (35). Surprisingly, upon addition of the agonist 9-cis RA, there is also a significant increase in the affinity of four corepressor peptides [NCoR_3L, silencing mediator of retinoid and thyroid hormone receptor (SMRT)_ID2, PR_H12 and BN2]. To verify the effect of 9-cis RA on the affinity of SMRT_ID2, the dose-response curves were measured, and Fig. 3B shows that
**Fig. 2.** Categorization of Peptides Based on the Effect of 9-cis RA on their Binding to RXR/LBD

**A**

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**Fig. 2.** Categorization of Peptides Based on the Effect of 9-cis RA on their Binding to RXR/LBD

Sequence alignment of A, Coactivator peptides that do not bind to RXR/LBD in the absence of ligand nor in the presence of 10 μM 9-cis RA; B, coactivator peptides that equally bind to RXR/LBD in the absence of ligand and in the presence of 10 μM 9-cis RA (log(MI) < 0.1) and in addition show a significantly high fluorescence intensity (> 3500); C, coactivator and D, corepressor peptides that show an increase in affinity for the RXR/LBD upon addition of 10 μM 9-cis RA (log(MI) > 0.3). Sequences are aligned manually by their coactivator motifs or corepressor motifs. The amino acids that are part of these motifs are indicated in bold. The peptides in panels B and C/D are ordered by increasing fluorescence intensities in the absence and presence of 9-cis RA, respectively. P.D. pept., Phage display peptide.
the binding of SMRT_ID2 improves upon addition of 9-cis RA, i.e. the estimated $K_d^{app}$ changes from 325 to 153 nM. It has been shown before that the second interaction domain of SMRT binds to RXR/H9251 in the absence of ligand (36) and that a synthetic agonist LG100268 induces SMRT binding to RXR/H9252 (37). However, for the first time, we demonstrate that the affinity of SMRT_ID2 increases upon addition of the natural agonist, 9-cis RA. Similar peptide recruitment data have been described for agonists in peroxisome proliferator-activated receptor (38). From a structural perspective, knowing that H12 adopts the agonist position in all 9-cis RA structures (6, 33, 39–41), this leads to the question what the role of this helix is in the
recruitment of the corepressor peptides. The only crystal structure of an NR LBD with a corepressor peptide [SMRT_ID2 (10)] showed that the peptide binds in the coactivator binding groove and that H12 is not in the agonist position. The N-terminal part of SMRT_ID2 relocates H12 toward the N terminus of H3 and partially occupies the agonistic location of H12. The recruitment of both coactivator and corepressor peptides by 9-cis RA suggest that 9-cis RA induces a conformational change in H12 that is stabilized in the agonist position by coactivator peptides and in the antagonist position by corepressor peptides.

The sequence alignment of the 13 coactivator peptides and the four corepressor peptides that do not bind in the absence of 9-cis RA, but do bind significantly better to RXRα in the presence of 9-cis RA, are shown in Fig. 2, C and D, respectively. The sequence alignment of the coactivator peptides shows that the residue types at position −1 and 2 are less conserved as compared with the residue types at these positions for peptides that already bind to RXRα in the absence of 9-cis RA (Fig. 2B). This suggests that coactivator peptides without a hydrophobic residue at position −1 and without a positively charged or H-bond donating residue at position 2 can only bind to the LBD when H12 is sufficiently stabilized by the ligand. With a sufficiently stable H12, the structural composition of the peptides becomes less critical for binding to the receptor, i.e. more sequential variation of the peptides is allowed.

**Clustering of Peptide Recruitment Profiles of Various RXRα Ligands**

Figure 1B shows the peptide recruitment profiles of a total of 11 different ligands, and 9-cis RA is depicted as a reference (solid black line). The 10 other ligands include five fatty acids, three synthetic agonists (targetin, LG100268, LG100324), a homodimer antagonist (LG100754), and a RAR agonist TTNPB (Fig. 4). Figure 1B clearly shows that some compounds induce peptide association in a similar manner as 9-cis RA,
whereas other compounds induce peptide dissociation (negative log MI values). Dissociation of peptides is confirmed by dose-response curves. For example, Fig. 3C shows that the estimated $K_{\text{app}}$ of RIP140_3 increases from 152 nM to 1129 nM upon addition of DHA. This indicates that the firm binding of RIP140_3 in the absence of ligand can be antagonized by a ligand such as DHA. This suggests that DHA destabilized H12 and thereby altered the optimal conditions for coactivator binding.

To identify which compounds induce identical or unique peptide recruitment profiles, we performed a hierarchical clustering (see Data Analysis in Materials and Methods) on the 11 ligand-induced peptide recruitment profiles (Fig. 5). The dendrogram of the hierarchical clustering of these peptide recruitment profiles was used to determine a boundary to cluster compounds that induce similar peptide recruitment profiles. Figure 6 shows the resulting four different clusters with the corresponding peptide recruitment profiles of the individual compounds in these clusters. On the basis of the shape of the peptide recruitment profiles, we refer to these clusters as a dissociative profile (1), silent profiles (2 and 3), and an associative profile (4).

The natural ligand of RXR, 9-cis RA, clusters together with targetin, LG100268, LG100324, and methoprene acid. These five compounds form the largest cluster (Fig. 6, cluster 4, 9-cis RA; black line). All compounds in this cluster improve the binding of coactivator and corepressor peptides. The enhancement of the binding of coactivator peptides is in agreement with previous studies, which demonstrate that these compounds act as agonists in RXR-mediated transcription pathways (18, 42–44). The recruitment of the four corepressor peptides by 9-cis RA (see above) is also observed for the other agonists in this cluster.

The second largest cluster contains three compounds that comprise three of the five fatty acids in the ligand set (phytanic acid, oleic acid, and DHA). These fatty acids deteriorate the binding of coactivator peptides and enhance the binding of the corepressor peptides NCoR_3L and, to a lesser extent, SMRT_ID2, indicating that these fatty acids are classical antagonists. However, the corepressor peptide BN2 is dissociated upon binding of these classical antagonists, suggesting that BN2 binds as a coactivator via its IXXL motif.

The three remaining compounds are in clusters 2 and 3 (Fig. 6). Cluster 2 contains the homodimer antagonist LG100754 and the fatty acid pentadecanoic acid; cluster 3 contains the RAR agonist TTNPB. Compared with the peptide recruitment profiles of the compounds in clusters 1 and 4, these compounds show negligible association or dissociation of most peptides. This raises the question whether these compounds actually bind to the receptor. Pentadecanoic acid binds in the LBP as has been demonstrated by x-ray (45). To determine the binding of LG100754 and TTNPB, we performed a peptide recruitment competition assay with CBP as peptide and 9-cis RA as ligand. In the absence of LG100754, the $EC_{50}$ of 9-cis...
RA is 3 nM (Fig. 7A). In the presence of 10⁻⁷ M LG100754, the EC₅₀ is increased to 43 nM, indicating that LG100754 competes with 9-cis RA for the same binding site. At even higher concentrations LG100754 (10⁻⁵ M), 9-cis RA is completely substituted by LG100754. These data indicate that LG100754 binds in the LBP but hardly affects the peptide binding profile, i.e., the compound does not induce a conformational change in the LBD that results in a significant dissociation or association of peptides.

The same competition assay was performed with TTNPB (Fig. 7B). In the presence of increasing concentrations of TTNPB, the EC₅₀ of 9-cis RA is hardly changed, indicating that TTNPB does not compete with 9-cis RA for the same binding site. Therefore we omitted this compound from further discussion.

Correlation between the Peptide Recruitment Profiles and Ligand Interaction Profiles

To investigate whether there are also differences between the binding modes of ligands, we first analyzed the binding mode of all compounds that have been cocrystallized with human (h)RXRα LBD. These compounds are the natural ligand 9-cis RA (6, 33, 39–41), its isomer all-trans retinoic acid (46), the synthetic agonists BMS649 (47) and L79 (48), and two fatty acids, DHA (47) and pentadecanoic acid (PDA) (45). The comparison of different ligand binding modes in RXRα has already been described for BMS649, 9-cis RA, and DHA by Egea et al. (47). We used a different approach by calculating so-called ligand-receptor interaction profiles, which are believed to represent the binding mode of the ligands. This facilitates the analysis of ligand binding in large sets of crystal structures in an automatic manner. Figure 8 shows the ligand-receptor interaction profiles of the six cocrystallized ligands with 9-cis RA highlighted as solid black line (see Materials and Methods for more details on the methodology to calculate these profiles). The ligand-receptor interaction profile describes the number of contacts between each residue in the LBD and the ligand. Residues in the LBD have been assigned so-called 3D numbers to allow easy comparison of different structures (49). It should be noted that the crystal structures used in this study are derived from multiple independent sources, which could lead to differences in binding modes of the ligand in the LBP.
in the different crystals. We therefore averaged the contacts between 9-cis RA and hRXR in the five available crystal structures (6, 33, 39–41) and calculated the SEM of the average number of contacts in the ligand-receptor interaction profiles (supplemental Fig. 2 published as supplemental data on The Endocrine Society’s Journals Online web site at http://mend.endojournals.org). This figure shows that the SEM is relatively small, indicating that the binding modes of 9-cis RA as resolved from multiple independent sources is highly similar. Figure 8 shows that the six ligands have interactions with residues in H3, H5, the β-sheet, H7, H10, and H11.

**Fig. 7.** Competition Studies of LG100754 and TTNPB with 9-cis RA

Dose-response curves of 9-cis RA were measured in a TR-FRET assay with GST-hRXRα LBD and CBP_1 (0.1 μM) in the presence of increasing concentrations of LG100754 (panel A) and TTNPB (panel B). The SEM of two separate experiments, each performed in duplicate, is indicated by the error bars.

**Fig. 8.** Ligand-Receptor Interaction Profiles of Compounds in hRXRα Crystal Structures

A ligand-receptor interaction profile describes for each position in the LBD the number of interactions with the ligand. The profile of 9-cis RA is highlighted as a solid black line. The profiles of the five other compounds that are cocrystallized with hRXRα (all-trans retinoic acid, DHA, PDA, BMS649, and L79) are shown in light gray. The secondary structures of the LBD that correspond with the 3D numbers are displayed below the graph. The average number of contacts and the SEM in five available crystal structures (6, 33, 39–41) of hRXRα with 9-cis RA are given in supplemental Fig. 2.
and H10. The most important ligand binding positions are 335, 339, 556 and 1057, which is in agreement with other members of the NR family (50). The ligand-receptor interaction profiles of the six compounds in Fig. 8 show that the number of contacts between various residues and the ligand differ among the six ligands. This suggests that the ligands have distinct binding modes, which was already described by Egea et al. (47) for three of these six ligands. We therefore performed a hierarchical clustering (see Data Analysis in Materials and Methods) on the six ligand-receptor interaction profiles, which resulted in four different clusters. The corresponding ligand-receptor interaction profiles of the individual compounds in these clusters are shown in Fig. 9.

9-cis RA, BMS649, and L79 form the largest cluster (Fig. 9, cluster 4, 9-cis RA; black line). The compounds in this cluster are described as agonists. BMS649 and L79 are synthetic agonists that better fill the LBP as compared with 9-cis RA. The natural ligand only occupies 59% of the total LBP volume (51). One part of the pocket that is not occupied by 9-cis RA is near residues at 3D positions 548 and 549 (W305 and N306 in hRXRα, respectively). The synthetic agonists are designed to better fill this side of the pocket, which is reflected in the ligand-receptor interaction profiles that show higher number of contacts between the ligand and positions 549 and 552 (Fig. 9, cluster 4) than that observed in the other clusters.

The three remaining clusters each contain one ligand (Fig. 9, cluster 1–3). Cluster 1 contains all-trans retinoic acid, which is the natural ligand of RAR. The x-ray structure of this compound with RXRα crystallized as a tetramer with H12 of each monomer in the coactivator groove of the adjacent monomer (46). The formation of a tetramer also led to a displacement of the N-terminal part of H3 and the C-terminal part of H10. Therefore, the ligand-receptor interaction profile of this compound is dissimilar to the other profiles. Clusters 2 and 3 both contain a fatty acid, PDA and
DHA, respectively. The major difference in the ligand-receptor interaction profile that separates these two clusters is the high number of contacts of DHA (47) with position 1057 (Fig. 9, cluster 3), which is not observed in the ligand-receptor interaction profiles of any other compound. Clustering of the ligand-interaction profiles of these six ligands suggests that there are four different binding modes of RXR compounds in the available crystal structures, which can be detected by this method.

Because the RXRα crystal structure data are limited to six compounds, of which only two are in the peptide recruitment data set, we manually modeled each of the 10 RXRα compounds that were profiled in the peptide recruitment assay and studied the interaction between the receptor and the ligand by MD (see MD Stimulations of Manually Docked Compounds in the RXRα LBD in Materials and Methods). TTNPB was excluded from the initial set, because this compound does not bind to the RXRα LBD (see above). Figure 10 shows the ligand-receptor interaction profiles that were calculated from model structures of the 10 compounds. To identify how unique or similar the different binding modes of the ligands in the LBP are, we performed the same clustering procedure (see Data Analysis in Materials and Methods) on the ligand-receptor interaction profiles as was applied to the profiles of the compounds in the crystal structures. From the dendrogram (data not shown) two distinct ligand-receptor interaction profile clusters could be identified. The corresponding profiles of the individual compounds in both clusters are shown in Fig. 11.

The first cluster contains oleic acid, DHA, phytanic acid, PDA, and LG100754. The second cluster contains all agonists: 9-cis RA (black line), targetretin, LG100324, LG100268, and methoprene acid. A clear difference between the two clusters is the number of contacts of the compounds with residues in the C-terminal part of H10, in particular 3D positions 1057 and 1061, which was already observed by Egea et al. in comparing the ligand binding modes in crystal structures with 9-cis RA, DHA, BMS649, and oleic acid (47). In general, compounds from the agonist cluster have a lower number of contacts with these residues than the compounds in the other cluster. To reveal a structural explanation for the differences in binding mode of compounds from the agonist cluster and the other cluster, we superposed hRXRα crystal structures, each with a compound from one of the clusters (9-cis RA from the agonist cluster and DHA from the other cluster; Fig. 12). Figure 12 clearly shows the differences of interaction between each of the two compounds and the H10 region. Although the side chain of cysteine 1057 (corresponding to residue 432 in hRXRα) is in the same orientation in both structures, leucine 1061 (436 in hRXRα) showed a movement of its side chain. In the crystal structure with DHA (47), the side chain of this leucine is shifted toward H12, whereas in the crystal structure with 9-cis RA this residue is pointing into the pocket of the LBP. The movement of this side chain toward H12 suggests a destabilization of this helix. This destabilization is reflected in the DHA-induced peptide recruitment profile, which shows a dissociation of most coactivators and confirms destabilization of H12 by interaction of ligands with residue 1061 (436 in hRXRα).

The clustering of the ligand-receptor interaction profiles does not distinguish between compounds that were dissociative (DHA, oleic acid, and phytanic acid) or silent (LG100754 and PDA) in their induced peptide recruitment profiles. We therefore compared the individual ligand-receptor interaction profile of LG100754 with the profiles of representative compounds from the two other clusters (9-cis RA for the associative, DHA for the dissociative peptide recruitment cluster; Fig. 11). In addition, the binding mode of LG100754 was compared with that of 9-cis RA by superposition of the x-ray RXRα structure with 9-cis RA [1FM6 (39)] and the coordinate set of RXRα with LG100754 of the frame with the lowest interaction energy (Fig. 13). Interestingly, the binding mode of LG100754 in the LBP is different from 9-cis RA in three areas. The first area is near the tryptophan at position 548 (3D no. 305 in
The propoxy group of LG100754 points toward this residue, whereas the other ligands do not occupy this space. This leads to a higher number of contacts in the ligand-receptor interaction profile of LG100754 at 3D residue positions 548, 549, 552, and 553 (corresponding to residues 305, 306, 309, and 310 in hRXRα) compared with the profiles of the other two compounds (DHA and 9-cis RA, Fig. 11). The second area is in the N-terminal part of H3. The tetra-hydronaphthalene moiety of the homodimer antagonist is shifted toward this part of the LBD, leading to a higher number of contacts with 3D residue positions 331 and 332 (corresponding to residues 264 and 265 in hRXRα). The third area comprises residues in H7. The binding mode of LG100754 shows a lower number of contacts between LG100754 and 3D residue positions 738 and 739 (corresponding to residues 345 and 346 in hRXRα) than 9-cis RA and DHA.

DISCUSSION

We hypothesized that the binding mode of a ligand in the LBP determines the conformation at the receptor surface through allosteric coupling. For this purpose, we determined whether compounds that bind differently in the LBP act differently in cofactor recruitment, by comparing the clustering based on ligand-receptor interaction profiles with the clustering of the compounds based on peptide recruitment profiles. We observed two distinct binding modes of ligands in the LBP by means of clustering of the ligand-receptor interaction profiles whereas we observed three distinct conformations of the RXRα LBD by means of clustering of the peptide recruitment profiles. These three distinct conformational changes of the RXRα LBD can be described as an associative, dissociative, or silent peptide recruitment profile. We observed that the five
Compounds in the associative peptide recruitment profile cluster are the same five compounds in one of the two clusters of the ligand-receptor interaction profiles. This indicates that there is indeed a correlation between the binding mode of a ligand and the conformation at the surface. However, the ligand-receptor interaction profiles do not yet discriminate between compounds that induce minor conformational changes (silent peptide recruitment profile) and compounds that destabilize H12 (dissociative peptide recruitment profile).

In this discussion we will focus on two issues. First, we discuss how well the data from a peptide recruitment assay reflect the effect of a compound in in vitro transactivation assays as described in the literature. Second, we give a structural explanation for the three distinct conformational changes at the RXRα surface, i.e. the structural mechanism underlying the associative, dissociative, and silent peptide recruitment profile.

**Peptide Recruitment Assay vs. in Vitro Transactivation Assay**

Current models of NR activation suggest that agonist compounds promote the association of a coactivator and the dissociation of a corepressor. Agonist compounds stabilize H12 in an active conformation, thereby facilitating the binding of a coactivator protein and reducing the affinity of corepressors. On the other hand, antagonists displace H12 from the active position, which results in enhancement of the binding of corepressors and reduction of the binding of coactivators. Finally, the so-called selective nuclear receptor modulators (SNRMs) induce such a conformation of H12 that both coactivator and corepressor can bind (52, 53). This may lead to an increase or decrease in gene transcription depending on the concentration of coactivators and corepressors in a cell. In this study, however, the coactivators and corepressors are represented by short peptide fragments, and the ligand-induced recruitment or dissociation of these fragments was measured by peptide recruitment. This raises two questions:

**How Well Does a Peptide Recruitment Profile Reflect the Effect of a Compound in a Cell?**

We observed that compounds in the associative cluster recruited not only coactivator peptides, but also corepressor peptides. In the context of the above described models for NR activation, our data indicate that 9-cis RA and other RXR agonists are not classical agonists, but actually act as SNRMs. The observation that these SNRMs induce coactivator and corepressor binding suggests that these compounds only partially stabilize H12 in the agonist position. This offers H12 sufficient conformational freedom to take the antagonist position in the presence of a corepressor peptide and the agonist position in the presence of a coactivator peptide. The final agonist or antagonist response of the compound in a particular cell type depends on the concentrations of cofactors in this cell type (52, 53). Because several studies have shown that these compounds act as agonists in various cell-based assays (18, 42–44), this suggests that these cell types...
have a higher concentration of coactivators than corepressors.

Compounds in the dissociative cluster reduced the binding of coactivator peptides whereas they increased the binding of corepressor peptides, i.e., the profile of a typical antagonist. This suggests that these compounds behave as full antagonists in cell-based assays. However, several studies show that fatty acids activate rather than inhibit RXRα-mediated gene transcription (19, 21). One possible explanation is that concentrations of corepressors in these cell-based assays may have been too low to compete with high levels of coactivators and that other cell types with high corepressor concentrations are needed to demonstrate the antagonist activity of fatty acids.

Compounds in the silent cluster did not induce a change in binding affinity for either coactivator or corepressor peptides, suggesting that there will be no change in basal gene transcription. Most likely, these compounds will be silent antagonists due to competition with the natural ligand 9-cis RA under physiological conditions.

In general, the correlation between the peptide recruitment profile of a compound and its cellular effect should be carefully interpreted, because it cannot be excluded that the ligand induced a conformational change at other locations at the surface of the LBD that are not detected by means of the current peptide recruitment assay. For example, it has been shown that ligands also induce conformational changes that lead to a different interaction with dimer partners, which results in a different cellular response (54, 55). We also cannot exclude that peptides in the peptide recruitment profile bind outside the classical coactivator binding groove, as has been suggested to alternative peptide binding positions as has been described in the literature. For example, the full-length cofactor recruitment of SRC3, FHL2, SRC1, and RIP140 (33, 34, 59, 60) is reflected by the recruitment of peptides derived from these coactivators. In addition, we noticed that the coactivator peptides can be classified as peptides that already strongly bind to the LBD in the absence of an agonist, peptides that only bind in the presence of a ligand, and peptides that do not bind. Peptides with a hydrophobic residue at position −1 in the peptide sequence did already bind strongly to the LBD in the absence of ligand, indicating that these coactivators force H12 in an agonist position, even in the absence of ligand. This suggests that these RXRα-cofactor complexes are continuously active in gene transcription even without ligand.

Peptides that lack a hydrophobic residue at position −1 did bind only to the RXRα LBD in the presence of 9-cis RA. This indicates that the cellular action of these coactivators is activated only when 9-cis RA is present. This suggests that these coactivators may be more important for ligand-induced gene transcription processes as compared with coactivators that always bind to RXRα. However, binding experiments with full-length cofactors and NRs are needed to study this in more detail.

**Peptide Recruitment Profiles and Underlying Structural Mechanisms**

**Associative and Dissociative Peptide Recruitment Profiles.** To determine the structural mechanism underlying the induced association/dissociation of peptides, we focused on compounds that induce an associative peptide recruitment profile and on compounds that induce a dissociative peptide recruitment profile. Comparison of the ligand-receptor interaction profiles of both clusters revealed that the compounds that induce a dissociative peptide recruitment profile have a significant higher number of contacts with residues 1057 and 1061 (3D nos. C432 and L436 in hRXRα, respectively) in the C-terminal part of H10 (47). This suggests that compounds that are in contact with the C terminus of H10 destabilize the agonistic binding mode of H12, resulting in dissociation of coactivators and therefore act as antagonists. A similar mechanism for antagonism has been described for ER (61). The differences in binding of 9-cis RA and DHA near the C terminus of H10 were already observed by Egge et al. (47), but for the first time we were able to link these structural interactions between ligand and receptor with the ligand-induced association and dissociation of coactivator and corepressor peptides.

**Silent Peptide Recruitment Profile.** Clustering of the ligand-receptor interaction profiles could not distinguish between compounds that induced a dissociative or silent peptide recruitment profile. However, the proposed ligand binding mode of LG100754 (Fig. 13) shows that there are three plausible explanations for the observation that this compound acts as a silent antagonist.

1. The shift of LG100754 toward H3 excludes a high number of contacts with the C-terminal part of H10, which is needed to destabilize H12 (see above).
2. W548 in H5 is described in stabilizing H12 via a water molecule (6), and therefore the position of this residue is important. Ligands protruding into the area near this residue could disrupt the stabilization of H12, as was already shown by the docking of the retinoid antagonist HX503 (6). The binding mode of LG100754 suggests that the orientation of the propoxy group toward W548 is such that H12 is neither stabilized nor destabilized. This hypothesis is strengthened by the observation that ligands that are identical to LG100754, but have a methoxy or ethoxy group instead of a propoxy group, act as agonists or partial agonists, respectively (54, 55).
3. The N-terminal part of H3 is described in direct
interactions with H12 and the C-terminal part of H11. In addition, an analysis of ligand-receptor contacts in all NR LBDs revealed that partial agonists interact more strongly with this part of H3 (50). LG100754 has a significant higher number of contacts with H3 that could destabilize H12. Together, this suggests that the homodimer antagonist has a unique binding mode, which neither stabilizes nor destabilizes H12 and results in a minor effect on peptide binding in the peptide recruitment assay.

**Final Conclusion**

In summary, this work shows for one class of compounds that there is a correlation between the binding mode of a ligand (represented by a ligand–receptor interaction profile) and the conformational change it induces at the surface of the RXRα LBD (represented by a peptide recruitment profile). We therefore conclude that clustering of ligand–receptor interaction profiles is a very useful tool to discriminate compounds that may induce different conformations. We also conclude that compounds can be easily classified by their peptide recruitment profiles. However, the translation to a cellular effect is difficult. These peptide recruitment profiles suggest that, depending on the cofactor concentration in the cell, five well-known RXRα agonists actually behave like SNRMs, whereas three weak agonistic fatty acids may act as antagonists and two compounds act as silent antagonists.

We therefore envision that the methods described in this paper can be of great value for drug design. With these methods new compounds can be profiled based on their ligand binding properties and their peptide recruitment properties. Moreover, the combination of both profiles will lead to useful insights in the working mechanism of a NR LBD, which in turn aid the design of compounds with a desired effect.

**MATERIALS AND METHODS**

**Plasmid Construction and Protein Purification**

cDNA coding for the ligand-binding domain of human RXRα (hRXRα, amino acids 221–462) was cloned into the EcoRI and XhoI sites of pGEX-4T-1 (Amersham, Piscataway, NJ). The protein was expressed as a fusion protein with glutathione S-transferase (GST) in *Escherichia coli* DH5α and purified by affinity chromatography.

**Ligands**

Figure 4 shows the structures of the ligands that were used in the peptide recruitment assay. Oleic acid, phytanic acid, DHA, pentadecanoic acid (PDA), methoprene acid, TTNPB, and 9-cis RA (all purchased from Sigma-Aldrich) and LG100324, LG100268, LG100754, and targetrin (all synthesized in house) were diluted in dimethyl sulfoxide to a final concentration of 10 μM.

**Peptides**

Supplemental Table 1, published as supplemental data on The Endocrine Society’s Online Journals web site, shows the sequences of the 52 N-terminal biotinylated peptides (Neo-system S.A., Strasbourg, France) derived from 15 different cofactors, five NRs, and various phage display peptide libraries (28, 62). Thirty three peptides contain the LXXLL coactivator motif, whereas three peptides possess a FXLX motif and two peptides have a FXXLY motif. Twelve repressor peptides in the set are described by the consensus sequence (L/I/V)XXX(L/I/V)XXX(L/I/V), whereas two repressor peptides (NcoR_4 and NCoR_4M) lack this consensus sequence. Due to similarity between the consensus sequences of a coactivator and corepressor motif, it is possible that some peptides contain both coactivator and corepressor motifs.

**TR-FRET Assay**

A TR-FRET assay was performed with 52 peptides in the absence and presence of ligand. Each reaction mixture (pH 7.2) contains 50 mM Tris, 50 mM KCl, 1 mM EDTA, 1 mM dithiothreitol, 0.1 mg/ml BSA, 10 nM RXRα LBD, 0.1 μM biotinylated peptide, 10 μM ligand, 8 mM allopregnanolone-labeled Streptavidin (PerkinElmer Life Sciences, Boston, MA), and 1.25 nm LANCE Eu-W1024-labeled anti-GST antibody (PerkinElmer Life Sciences). Each experiment was carried out in duplicate in 384-well plates. Plates were centrifuged and incubated for 24 h at 4°C. Fluorescence at 665 nm was measured on a Victor (Wallac, Inc., Gaithersburg, MD).

To directly visualize the effect of a ligand on recruitment of the different peptides, the MI was calculated, i.e., the ratio between the fluorescence intensity in the presence of a ligand and the fluorescence intensity in the absence of ligand. An MI of 2.0 indicates a 2-fold increase of the amount of peptide bound, whereas a MI of 0.5 indicates a 2-fold decrease in the amount of peptide bound. This nonlinear behavior was translated into a linear positive signal for an increase in peptide binding and a linear negative signal for a decrease in peptide binding by calculating the $10^{\log(MI)}$. Log(MI) values above zero indicate recruitment of peptides and thus an associative effect of the ligand on peptide binding. In contrast, negative values indicate dissociation of peptides and therefore a dissociative effect of the compound on peptide binding. A log(MI) value of around zero means that a peptide does not bind and/or that the binding is not changed by the ligand.

To determine the apparent $K_a$ values of specific peptides, the binding curves were determined in a peptide concentration range between 0.1 nM and 1 μM. Calculations were performed in GraphPad Prism (GraphPad Software, Inc., San Diego, CA; nonlinear regression curve fit, one-site binding model). To determine EC$_{50}$ values of ligands, we measured ligand dose-response curves with ligand concentrations ranging from 31.6 pm to 1 μM.

**MD Simulations of Manually Docked Compounds in the RXRα LBD**

MD simulations were performed using QUANTA/CHARMm (version 3.1b; Ref. 63) following a protocol that was described previously by Kouwijzer et al. (64). The hRXRα LBD of 1F68 (chain A) was selected as template structure. The initial binding modes of six ligands were obtained from available x-ray structures of either hRXRα or hRXRβ by superposition of the structures onto the template structure. The superposed hRXRα structure with DHA [1MV9 (47)] was used to construct the initial binding mode of phytanic acid, and the superposed hRXRβ structure with LG100268 (37) was used for all others. LG100754, LG100324, and targetrin (supplied as Table 2, published as supplemental data on The Endocrine Society’s Online Journals web site). The complexes were protonated, charges were assigned, and ligand atom types and bond
orders were corrected. The MD run started with a heating phase of 10 psec, followed by a 100-psec run at 400 K during which time the coordinate sets were saved every picosecond. These coordinate sets were energy minimized, and for each of these 100 frames the average interaction energy between ligand and protein was calculated. The 10 coordinate sets with the lowest average interaction energy were used to calculate an average ligand-receptor interaction profile.

**Ligand-Receptor Interaction Profiles**

The number of contacts between ligands and residues in the RXRa LBD was calculated by the Automatic Residue Extraction System (ARES), as was previously described for the analysis of ligand binding in all NR LBDs (50). For each unique receptor-ligand combination, ARES automatically creates a so-called ligand-receptor interaction profile. A contact is included in the profile when the distance between the Van der Waals’ surfaces of the atoms of a ligand and a residue is less than 1 Å. A unique number is assigned to each residue position that is structurally conserved within the family of the NR LBDs (49). This 3D number starts with one or two digits that indicate the helix number; similarly B and L reflect residues in the β-sheet and loops, respectively. Residues that are not structurally conserved and therefore do not have a 3D number all got the number zero. Contacts in multiple copies of the same LBD-ligand complex were normalized. If, for example, a LBD-ligand complex observed two times in one PDB file and three times in another, all contacts in all five chains are summed up and divided by 5.

**Data Analysis**

The TR-FRET assay and the contact analysis by ARES yielded peptide recruitment profiles and ligand-receptor interaction profiles, respectively. Spotfire DecisionSite (Spotfire, Somerville, MA) was used to cluster both types of profiles by a hierarchical unweighted clustering routine with a similarity measure that was based on (cosine) correlation. Resulting dendrograms were used to determine the boundary to separate clusters.

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