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Molecular mechanism of allosteric communication in the human PPAR α -RXR α heterodimer

Tuomas Venäläinen, Ferdinand Molnár, Chris Oostenbrink, Carsten Carlberg, And Mikael Peräkylä⁶

ABSTRACT

The peroxisome proliferator-activated receptor α (PPAR α) is a nuclear receptor (NR) that forms a heterodimeric transcription factor complex with the retinoid X receptor α (RXRα). The phenomenon that the heterodimer can be activated by both PPARa and RXRa ligands, while both ligands have a synergistic effect on its activity suggests that there is an allosteric communication within the heterodimer. In this study, the molecular mechanism of this allosteric signaling was studied by molecular dynamics (MD) simulations and some of the residues involved in this communication were tested experimentally. Multiple MD simulations were done for the PPARa-RXRa heterodimer ligand-binding domains (LBDs) without ligands, with agonistic ligand bound to RXR\u03c0 or PPAR\u03c0, and ligand bound to both receptors. Fluctuation calculations and structural clustering analysis of the heterodimer MD simulations showed that ligand binding to RXRa decreases fluctuations of large parts of PPARa, most notably helices 3 and 4 at the coactivator binding site, which presumably stabilizes the coactivator binding to heterodimer complex. The dynamics of helix 8-9 loop and helix 10/11 located at the heterodimeric interface were affected by RXRa ligand binding, suggesting that these parts of the dimer are involved in allosteric communication. Experimental data complemented this view by showing that a large set of residues at the heterodimerization surface has a role in the communication. These results provided evidence that RXRα ligand binding-induced stabilization of PPARα coactivator binding site has a role in the permissive and synergistic activation of the PPARa-RXRa heterodimer.

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Key words: molecular dynamics; nuclear receptors; protein-protein interactions; point mutations; activation.

INTRODUCTION

Allosteric communication of proteins plays a key role in regulation of many biological systems. Allosteric regulation is needed, for example, in mediating signals from outside of the cell into the cell by G-protein coupled receptors, 1 cooperative binding of oxygen molecules into hemoglobin 2 or the regulation of gene expression by nuclear receptors (NRs). Common to these examples is that a small molecule binding to one part of the protein controls the biological activity in another part, which is located far away from the small molecule binding site or even in another protein. In this work, allosteric activation in a NR heterodimer of the peroxisome proliferator-activated receptor (PPAR) α and the retinoid X receptor (RXR) α is studied with molecular dynamics (MD) simulations and experimentally using reporter gene assays.

NRs form a superfamily of ligand-inducible transcription factors that regulate gene expression of a large variety of target genes. Natural NR ligands form a wide group of small lipophilic molecules including steroid hormones, fatty acids and xenobiotics, many of them important in cell regulation and metabolism. 4–6 This makes NRs interesting as drug targets. Binding of an agonist to the ligand-binding pocket (LBP) of a NR leads to a conformational change in the ligand-binding domain (LBD) leading to an interaction

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with coactivator proteins.⁷ The largest conformational change occurs in the C-terminal helix 12, which acts as a molecular switch allowing either coactivator or corepressor protein binding. Agonist binding favors a conformation, in which helix 12 is docked against the helices 3 and 11 forming one side of the activation function 2 (AF-2) surface and facilitating coactivator binding.⁸ Antagonists prevent binding of helix 12 against the core of the LBD and formation of the coactivator binding surface. Without an agonist or with an antagonist, many of the NRs bind a corepressor protein and their target genes are silenced.4

The outcome of the NR action depends on the coregulator proteins bound. Often there is a balance between coactivator and corepressor proteins, which is controlled by helix 12. The conformation of helix 12 in turn is modulated by ligand binding. Thus, the final outcome depends on the relative concentrations of the coregulatory proteins, ligand binding, ligands' properties, and the nature of the receptor itself. Many of the NRs like retinoic acid receptor α (RAR α), vitamin D receptor (VDR), thyroid hormone receptor (TR), and steroid hormone receptors are not active without an agonist, whereas others, such as PPARs and constitutive androstane receptor (CAR) have significant constitutive activity. 9,10

Although the conformation of helix 12 is the main determinant of NR LBD controlled activation and repression, other more subtle structural and dynamical factors have been observed. Bruning et al. showed by the amide H/D exchange (HDX) method that binding of intermediate and partial agonists lead to considerably lower stabilization of PPARy helix 12, but unexpectedly, they stabilized other parts of the LBD, mainly the β-sheet area and helix 3.11 It was also shown that stabilization of helices 3 and 11 is not sufficient to stabilize helix 12. Similar observations were made by computer simulations on the glucocorticoid receptor (GR) and its mutants. In these MD simulations, an increased biological activity was found to correlate with reduced fluctuations of helices 3, 4, and 5.12 MD simulations have also suggested that ligand binding modifies LRH-1-cofactor interactions without causing overall changes in to protein structure. 13 In addition, by a new class of PPAR α/γ agonists it could be shown that direct stabilization of helix 12 by ligand binding is not necessary for PPAR agonism. 14,15 These ligands were shown by X-ray crystallography to adopt a novel binding mode lacking contacts with helices 11 or 12. Taken together, these results suggest that a helix 12 independent mechanism contributing to NR activation exists.

Many NRs function as heterodimers with RXR.4,7 Heterodimer formation is required for efficient contact to DNA binding sites, so-called response elements, in primary NR target genes. On the basis of the heterodimer's response to ligands, NRs can be classified to be permissive or nonpermissive.³ Permissive receptors can be activated also with RXR ligand and they give full response when ligands are bound to both receptors of the heterodimer, whereas nonpermissive receptors do not respond to RXR ligand, but give a full response with their own ligand.³ Conditionally permissive receptors display synergy, or increased activity with two ligands, but cannot be activated with RXR ligand alone. 16

To better understand the mechanism of permissivity, its dynamic nature and synergistic activation of NRs in general at the atomic level, we studied the PPARα-RXRα heterodimer with different combinations of ligands using MD simulations. PPARα was chosen for this study because of its key role in lipid metabolism and its highest constitutive activity among the three PPAR subtypes.¹⁷ From the MD simulations, we have identified residues involved in the communication between the two monomers in the PPARα-RXRα complex and our model has been investigated experimentally using reporter gene assays. This study provided new information of ligandinduced intermonomeric allosteric communication in the PPARα-RXRα heterodimer, which may be applied to the other NR heterodimers with permissive activation profile.

METHODS

Constructing the simulations systems

Because there were no crystal structures available for the PPARα-RXRα heterodimer, a PPARγ-RXRα heterodimer structure was used as template. In this original PPARγ-RXRα structure (Protein Data Bank (PDB) code 1K74¹⁸) 9-cis retinoic acid (9cRA) is bound to the RXRα, GW409544 (GW) is bound to the PPARγ, and two coactivator peptides are bound to the complex. From the template, PPARγ was replaced with the PPARα structure (PDB code 1K7L¹⁸). For the simulations, the coactivator peptides were removed, because they were expected to stabilize the receptor into the active conformation and to mask the effects of the ligands.

The resulting PPARα-RXRα heterodimer contained 499 amino acid residues. Crystallographic water molecules from the PPARγ-RXRα heterodimer were kept, when they did not overlap with PPAR α residues. Side chains of PPARa residues at the heterodimer interface were rotated to the same conformations, as observed in the PPARγ-RXRα heterodimer. This could be determined unambiguously, because the residues of the heterodimer interface of the two PPAR subtypes are highly conserved. Four PPARα-RXRα MD simulation systems were constructed: (1) apo heterodimer: the unliganded LBD heterodimer, (2) 9cRA heterodimer: 9cRA bound to the RXRα LBD with PPARα unliganded, (3) GW heterodimer: GW bound to the PPARa LBD with RXRa unliganded, and (4) 9cRA + GW heterodimer: ligands bound to both RXRα and PPARα LBDs. The dimeric complexes were placed in a periodic box of 7.3 \times 9.9 \times 8.3 nm containing \sim 17,000 SPC water molecules. The net charge of the system (-12 with two ligands) was neutralized with randomly positioned Na⁺-ions.

MD simulations

All the simulations were performed using the GRO-MACS program¹⁹ with the GROMOS 45a3 force field.²⁰ The missing parameters for the ligands were derived by analogy to the amino acid parameters in the GROMOS force field (Supporting Information Tables S1-S10). The LINCS²¹ method was used to constrain all the bond lengths of the protein and the ligands and the SETTLE algorithm²² was used to constrain all the bond lengths and angles of the water molecules. The time step for the leap-frog algorithm was set to 2 fs. A twin range cut-off was used for nonbonded interactions updating the neighbor list within the short range cut-off radius at 8 Å every 10 steps, at which times interactions in the intermediate range (14 Å) were also calculated. A reaction field contribution²³ with a relative dielectric permittivity of 62.0²⁴ was added to the electrostatic energies and forces to account for long-range interactions outside the cut-off sphere. During the MD simulations rotation and translation of the protein was removed every 10 steps.

Initial velocities were sampled randomly from a Maxwell-Bolzmann distribution at 100 K. The water molecules of the simulation systems were first heated at constant volume from 100 to 300 K in 120 ps with position restraints on the protein heavy atoms. The protein was then equilibrated at 300 K at constant pressure conditions for 250 ps while slowly releasing protein restraints. Finally, the systems were equilibrated for 300 ps at constant pressure. Temperature and pressure were controlled with the Berendsen weak coupling algorithm²⁵ with coupling constants of 0.1 and 0.5 ps, respectively. After that, production simulations of 2 ns were started. Ten parallel MD simulations for each simulation system were started after the equilibration phase by randomly assigning new starting velocities. Coordinates were stored every 0.1 ps. Average structures of the last 500 ps of the production trajectories were used in structural analysis and the last 1 ns was used for the fluctuation and structural clustering calculations. Other sampling times were also tested with no change in the conclusions.

To validate our methods, ten parallel simulations were performed for the PPARα and RXRα monomers using the same computational protocols as in the heterodimer simulations. The results are presented in the Supporting Information.

Compounds

The mouse CAR agonist TCPOBOP was kindly provided by Dr. P. Honkakoski (University of Kuopio). 1α,25(OH)₂D₃ and the PPARδ agonist L763483 were a gift from Drs. L. Binderup and M. W. Madsen (Leo Pharma, Ballerup, Denmark). The 9cRA, methoprene acid, GW7647 and the 25-OH cholesterol were obtained from Biomol (Copenhagen, Denmark). 9cRA and 1α,25(OH)₂D₃ was dissolved in ethanol and 2-propanol, respectively, whereas the other compounds were dissolved in dimethyl sulfoxide (Me₂SO). Further dilutions were made in Me₂SO.

Protein expression vectors

Full-length cDNAs for human PPAR α , ²⁶ human RXR α , ²⁷ human VDR, ²⁸ human LXR β , ²⁹ and human RARα³⁰ were subcloned into the T7/SV40 promoterdriven pSG5 expression vector (Stratagene), mouse CAR³¹ into the CMV promoter-driven vector pCMX and that for human PPAR832 into the CMV promoterdriven vector pcDNA3 (Invitrogen). The point mutants of PPARα and RXRα were generated by using the Quick-Change site-directed mutagenesis kit (Stratagene) according to the manufacturer's instructions. All mutations were confirmed by sequencing.

Reporter gene constructs

Four copies of the human CPTI gene DR1-type RE (core sequence GTAGGGAAAAGGTCA),³³ four copies of the rat ANF gene DR3-type RE (core sequence AGAGGT CATGAAGGACA),34 one copy of the PBREM element from mouse CYP2B10 gene promoter (core sequence GAGTGTCAGGCAAGTTGATGGTGCACTGTGCCAAGG TCAGGAAAGTACAGA),³⁵ four copies of the human iNOS gene promoter DR4-type RE (core sequence GTGGTTCATGCCGGTTCA)³⁶ and two copies of the consensus DR5(T/T)-type RE (AGAGTTCACCGAAAGT TCA) were individually fused with the thymidine kinase (tk) minimal promoter driving the firefly luciferase reporter gene.

Transient transfection and luciferase reporter assays

HEK293 human embryonic kidney cells were seeded into 6-well plates (200,000 cells/well) and grown overnight in phenol red-free Dulbecco's modified Eagle's medium (DMEM) supplemented with 5% charcoalstripped fetal bovine serum. Plasmid DNA containing liposomes were formed by incubating 1 µg of an expression vector for wt and mutant PPARα and/or RXRα and 1 μg of reporter plasmid bearing four copies of the human CPTI DR1-type RE with 10 µg of DOTAP (Roth, Karlsruhe, Germany) for 15 min at room temperature in a total volume of 100 μL. After dilution with 900 μL of phenol red-free DMEM, the liposomes were added to the cells. Phenol red-free DMEM supplemented with 500 µL of 15% charcoal-stripped fetal bovine serum was added 4 h after transfection. At this time, 10 nM 9cRA, 100 nM of PPARα selective agonist GW7647, the combination of

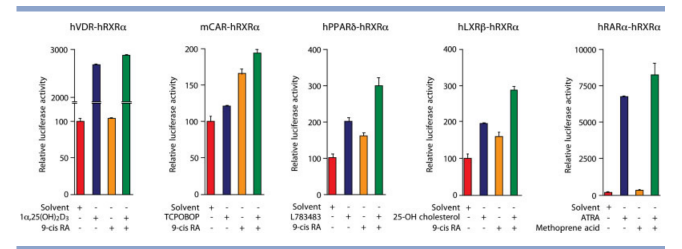


Figure 1

Differential response of RXR heterodimers to ligands. From left: Nonpermissive heterodimer pSG5-hVDR on the tk-[rANF DR3 VDRE]_s-luc reporter plasmid in HEK293 cells in the presence of endogenous RXR, treated with 100 nM 9cRA RXR agonist and 100 nM 1α,25(OH)₂D₃. Permissive heterodimers: Activation of tk-PBREM-luc reporter by pCMX-mCAR in HEK293 cells and by treatment with 100 nM 9cRA and the 1 μM mouse CAR agonist TCPOBOP, pcDNA3-hPPARδ on the tk-[hmCPTI DR1 PPRE]₄-luc reporter plasmid in HaCaT cells in the presence of endogenous RXR, treated with 100 nM 9cRA RXR agonist and 100 nM L783483, tk-[hiNOS DR4 LXRE]₄-luc reporter in the presence of pSG5-hLXRβ in HEK293 cells and by treatment 100 nM 9cRA and 10 μM LXR agonist 25-OH cholesterol. Conditionally permissive heterodimer: Activation of a tk-[consensus DR5 T/T RARE]2-luc reporter by pSG5-hRAR α in HaCaT cells and by treatment with 1 μ M methoprene acid (RXR α specific agonist with no activity on RAR) and 10 nM RAR agonist ATRA. The adequate ligand conditions are indicated in color bars: Apo (red); 9cRA (orange); RXR partner's ligand (blue); 9cRA + RXR partner's ligand (green).

9cRA and GW7647 or solvent was also added. The cells were lysed 16 h after the onset of stimulation using the reporter gene lysis buffer (Roche Diagnostics), and the constant light signal luciferase reporter gene assay was performed as recommended by the supplier (Canberra-Packard, Groningen, The Netherlands). The luciferase activities were normalized with respect to protein concentration. A two-tailed, paired Student's t-test was performed, and P values were calculated in reference to the solvent control (*P < 0.05; **P < 0.01; ***P < 0.001).

RESULTS

Differential response of RXR heterodimers to ligands

PPARs belong to the permissive group of NRs since in cell-based assay the PPAR-RXR heterodimer is activated also when only 9cRA is present, whereas the nonpermissive VDR-RXR complex shows no extra activation with 9cRA compared to its apo state. Conditionally permissive heterodimers, such as RAR-RXR, show very week activation when RXR specific ligand, for example, methoprene acid, is bound to the heterodimer (Fig. 1).

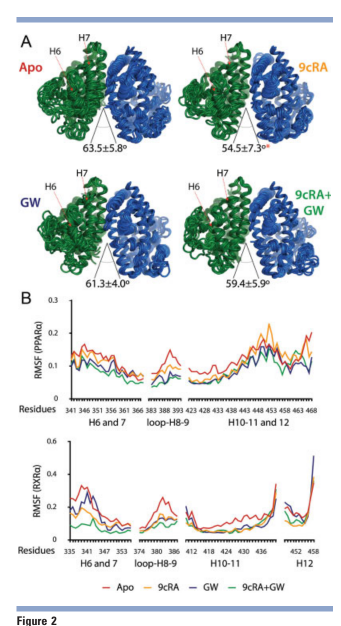
PPARa-RXRa LBD heterodimer stability in the simulations

To investigate the stability of the simulations, the atom-positional root-mean-square deviation (RMSD) of the $C\alpha$ atoms relative to the crystal structure (Supporting Information Fig. S4) were calculated. The RMSD values in all heterodimer simulations were clearly lower (by $\sim 0.5 \text{ Å}$) than those in the monomer simulations (Supporting Information Figs. S2 and S3) being mostly <2Å. This indicates that the MD simulations were stable and that heterodimer formation stabilizes the LBDs. Heterodimer simulations with 9cRA bound to RXRa seem to stay closer to the starting structure than the simulations without 9cRA.

Ligand binding restricts the movements of the heterodimer

To further investigate the ligand effects on the heterodimer flexibility, average structures of the last 500 ps of the parallel MD simulation were compared. The 10 parallel average structures of each ligand condition were superimposed using the $C\alpha$ atoms (Fig. 2). In overall, the apo heterodimer showed higher differences between the average structures compared to conditions where ligands were present [Fig. 2(A)]. Helices 6 and 7 of the RXRα have the highest flexibility in the apo and GW bound heterodimers and the lowest when 9cRA and both 9cRA and GW are present. Interestingly, PPARα was more rigid in all conditions compared to RXRα.

The tilt angle of helices 10–11 (K449 $/C\alpha(PPAR\alpha)$ -D466/ $C\alpha(PPAR\alpha)$ - F438/ $C\alpha(RXR\alpha)$) in the heterodimer is lower when ligands are bound. Binding of 9cRA has a larger effect on the angle than GW binding [Fig. 2(A)]. However, there is no significant change in the dihedral angle between the residues K449/Cα(PPARα)-D466/



Ligand binding restricts the movements of the heterodimer. The 10 superimposed average structures for every ligand condition from the last 500 ps of the MD simulations are visualized in tube representation. (A) Below the structures the angle between the helices 10-11 of PPARα and RXRα is indicated and the helices 6 and 7 are highlighted with red arrows. The graphical representation of the structural differences (RMSF data) of the heterodimeric complex for every residue at the heterodimerization interface is shown in detail. (B) The different ligand conditions are displayed in colored lines: Apo (red); 9cRA (orange); GW (blue); 9cRA + GW (green).

 $C\alpha(PPAR\alpha)$ -L430/ $C\alpha(RXR\alpha)$ - F438/ $C\alpha(RXR\alpha)$ indicating that the tilting takes place only about one axis. The PPARα's helix 12 (the distance between P458/Cα and Y468/Cα) was significantly shorter in the presence of GW (17.22 \pm 1.13 Å*) compared to apo protein (18.27 \pm 0.97 Å). This is most likely due to stabilization of PPARα's C-terminus by direct interaction with helix 10/

11 of RXR α in the apo heterodimer, whereas in the presence of GW the inter-molecular stabilization is partly replaced by intra-molecular interactions.

To obtain a quantitative measure of the structural differences, atom-positional root-mean-square fluctuations (RMSFs) around the average positions of Cα atoms were measured for the 10 parallel average structures [Fig. 2(B)]. The RMSF values of the helices 7, 10–12 and the helix 8,9 loop at the heterodimer interface show that differences between the parallel structures are highest in the apo heterodimer, and lowest in the GW + 9cRA heterodimer. It is also seen, that binding of a ligand to one monomer leads to more regular behavior of the other monomer.

In conclusion, binding of ligands leads to more regular behavior of the heterodimer, which is seen in the smaller differences between the average structures of the parallel MD simulations. Narrowing of the tilt angle of the helices 10/11 is likely to be connected with the ligand binding -induced changes in the behavior of the heterodimer.

Ligand binding increases packing at the heterodimerization surface

Analyses of the average structures of the last 500 ps of the simulations show that ligand binding pushed helices 7 and 11 of PPARα and RXRα closer to the partner protein, which decreased their separation across the heterodimer interface. For example, the distance between the two helix 11 residues K431 (RXR α) and T438 (PPAR α) was on average 1.0 Å shorter in the 9cRA simulations than in the apo simulations. Another helix 11-helix 11 distance, E434 (RXR α)- Q442 (PPAR α), was 1.8 Å shorter in the 9cRA simulations than in the apo simulations, and 0.9 A shorter in the GW simulations than in the apo simulations. This was also seen in the angle of the helices 10/11 [Fig. 2(A)].

To further analyze the structural change, the dimerization surface area of each PPARα-RXRα system was calculated. The solvent accessible surface area of PPARα-RXRα heterodimer was calculated for every 1 ps and subtracted from the sum of surface areas of PPAR α and RXRα calculated separately. The average surface areas buried upon heterodimer formation calculated from 10 parallel MD simulations were 1740 \pm 130 Å², 1700 \pm 90 Å², 1750 \pm 70 Å², and 1780 \pm 50 Å² in the apo-, 9cRA-, GW- and GW + 9cRA systems, respectively. According to the *t*-test, only when 9cRA simulations were compared to GW + 9cRA simulations the dimerization areas were found to be different at 95% confidence level. These results suggest that ligand binding induced conformational changes in the helices 7 and 11 may increase the dimerization surface and thus stabilize the heterodimer partner. However, this increase is small and could not be seen in the case of 9cRA binding, even though the change in the helix 11-helix 11 angle was

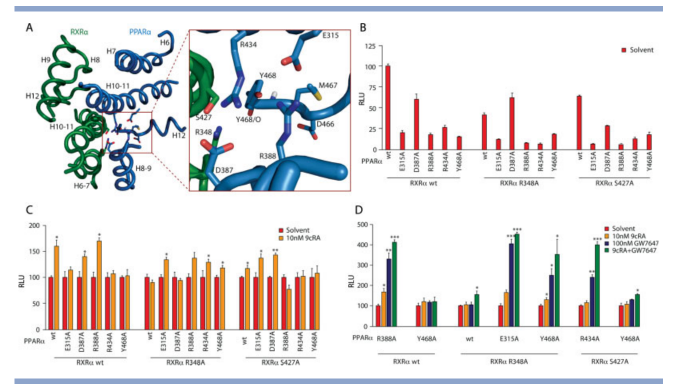


Figure 3

Allosteric communication at the heterodimerization interface. The heterodimerization interface of PPARa (in green)-RXRa (in blue) is shown. Location of amino acid residues R348 and S427 in RXRα, and E315, D387, R388, R434, and Y468 in PPARα mutated in this work are shown in detail. (A) Basal and ligand-induced activities of the wild type (wt) and mutated PPARa-RXRa heterodimers were measured using extracts from HEK293 cells that were transiently transfected with a reporter construct containing four copies of the human CPTI DR1-type RE and expression vectors for wt and mutant PPARα and/or RXRα. Cells were treated for 16 h with solvent (B), 10 nM 9cRA (C), 100 nM of PPARα selective agonist GW7647 or with the combination of 9cRA and GW7647 (D). The data were normalized to the activity of the wt heterodimer (B) or to the solvent treated condition (C and D). Columns represent the mean of at least three experiments, and bars indicate S.D. A two-tailed, paired Student's t test was performed, and p values were calculated in reference to the solvent control (*, P < 0.05, **, P < 0.01, ***, P < 0.001). Red color column bars indicate basal activities in apo state (B, C, D), the orange column show the ligand inducibilities when 9cRA was present (C, D) and the blue and green columns bars indicate the ligand conditions when GW7647 and 9cRA + GW7647 were present, respectively.

largest in that case [Fig. 2(A)]. On the basis of the standard deviations of the average surface areas, the apo systems showed the most irregular behavior of the MD simulations.

In summary, the MD simulations showed that ligand binding leads to changes in the PPARα-RXRα LBD heterodimer, which bring helices 11 of the two receptors closer to each other. The result of this conformational change is an increase in the intensity of packing at the dimerization surface and possibly slightly increased dimerization surface area.

Direct PPAR α helix 12 stabilization by RXR α helices 7 and 11

The C-terminus of apo PPARα's helix 12 was seen to adopt a position in which it is closer to RXRα than in a liganded PPARα. On the other hand, binding of 9cRA to RXRα relocated helices 7 and 11 of RXRα towards helix 12 of PPARα, which stabilized its position by a hydrogen bond between K431 of RXRα and R348 of PPARα. This mechanism may explain at least part of the RXR\alpha ligand-induced activation of PPARα, as it has been suggested by Gampe et al. on the basis of crystal structures of the PPARγ-RXRα LBD heterodimer.³⁷

Allosteric communication at the heterodimerization interface

The heterodimerization interface of PPARα-RXRα consists of the helices H6-7, H8-9, H10/11, and helix 12 [Fig. 3(A)]. The information exchange between the monomers takes place through this interface and it can be disturbed by mutating the residues in these parts of the proteins.

The effect of an alanine mutation depends on the relative importance of the mutated residue, the nature of the interaction of that residue with other parts of the complex and resulting changes on the protein structure. Ideally, alanine mutation on the heterodimerization interface reveals the relative importance of the mutated residue on the communication between the monomers. However, the resulting change in the activation profile of

Table I Interactions Between Residues at the Heterodimerization Interface

	Ligand conditions in the MD simulations			
Interaction (PPAR α -RXR α)	Apo (%)	9cRA (%)	GW (%)	9cRA + GW (%)
H7 H8-9 K364-D379	30	30 (+10)	10 (+30)	60 (+30)
H8-9 H7 G386/O-K356	0	0	20	0
H8-9 H7 D387-R348	10	0	10	0
H8-9 H7 P389/NH-E352	0	0	10	0
H9 H7 E398-K356	20	10	0	40 (+ 20)
H9 H9-10 R409-E401	10	0	0	0
H9 H10 E402-K417	10	0	0	0
H10 H9 K425/O-E394	(+50)	(+40)	(+90)	60 (+10)
H10 H9 Q428-R393	0	0	20	10
H10 H9 Q428-Y397	0	0	10	0
H10 H10 A431/0-R426	0	10	20	0
H10 H10 D432-R426	20	60	40	20
H10 H10 R434-P423/0	0	40	80	80
H10 H10 R434-S427	60	60	90	90
H10 H7 R434-E352	0	10	0	0
H11 H10 Q435-R426	40	70	70	80 (+20)
H11 H10-11 Q442-E434	0	10	0	0
H12 H7 D466/O-R348	0	10	0	0
H12 H7 M467/O-R348	0	30	0	0
H12 H10 Y468-S427	0	10	0	0
H12 H7 Y468/O-R348	40	10 (+30)	0	10
H12 H10 Y468/0-K431	10	50 (+20)	20 (+10)	10 (+10)

The table lists all the hydrogen bond capable interaction (distance <3.5 Å) of the residues located at the heterodimerization interface. The values (in %) represent the occurrence of the interactions in 10 independent MD simulations, and the values in brackets express the additional possibilities where the distance of the hydrogen bond is between 3. 5 and 4.0 Å.

the complex can be weaker, if the mutation is compensated by other contacts, or stronger, if mutation of a large residue into alanine leads to structural changes that modify also other contacts between the monomers. We expect an alanine mutation on a hydrophobic residue to be more easily compensated than a mutation of a hydrogen bond forming residue, because the strength of a hydrogen bond depends largely on the angle and the distance between the donor and the acceptor. The resulting effect of the mutation of a hydrogen bond forming residue is expected to be stronger and to tell more about the importance of that residue on the allosteric communication, although the compensating hydrogen bonds with back bone or with other residues or other structural changes cannot be avoided.

The average structures of the MD simulations were examined to identify candidate residues, which are able to form hydrogen bonds at the heterodimerization interface (Table I). In the table, the percentage of occurrence of a hydrogen bond is indicated in all the four ligand conditions. Some interactions such as R434 (PPARα)-P423/C=O (RXR α) or R434 (PPAR α)- S427 (RXR α) are seen under most of the conditions indicating the significance of these residues in the intermonomeric interaction. The interactions, which are formed occasionally (indicated with smaller % value in Table I) may compensate a mutation.

To examine the information flow through the dimerization surface, the following amino acid residues were substituted with alanine residue: E315, D378, R388, R434, and Y468 found in human PPARα and R348 and S427 in RXRα [Fig. 3(A)]. These residues are located at the heterodimerization interface in H6-7 and H10-11 of RXRα and H8-9, H10-11 and H12 of PPARα. For instance, the residue R348 (RXR α) in the apo state is able to make contacts with four different residues in PPARα (D387, D466/C=O, Y468/Oterm1, and M467/ C=O) showing some alternative options for the communication. We also suspect that this residue may be involved in the intra-helical stabilization of the RXRa H10/11. The drop of the basal activity of the heterodimer to 41% compared to the wild-type receptor is the combined effect of all these functions connected to R348 (RXR α) [Fig. 3(B)]. The PPAR α residues E315, D387, R388, and R434 are placed in the close proximity to the heterodimerization interface and also stabilize directly the PPARα helix 12 through the contact with Y468 [Fig. 3(A)]. This interaction is more visible in the apo state of the heterodimer, when the high basal activity is mediated also through this stabilization. The critical role of these residues is highlighted by a marked drop of the basal activities when mutated to alanine [Fig. 3(B)]. Although a PPARα mutant in combination with a RXRα mutant usually results in an additional decrease in the basal activity [Fig. 3(B)], the activity of PPARα Y468A mutant is not affected further when combined with R348A or S427A RXRα mutants. A particularly interesting profile can be seen for the mutation D387A: with wild-type RXRα the basal activity is decreased to 59% and in combination with R348A (RXRα) it is 61.5% compared to the wild-type heterodimer. However, in combination with S427A the basal activity is as low as 28% due to the indirect effect of the D387 (PPARα)- R434 (PPARα)- S427 (RXR α) interaction, where the R434 (PPAR α)- S427 (RXRα) contact is also compromised and the additional hydrogen bond between R434 (PPARα)- P423/C=O cannot rescue the abolished interaction. This results in a condition, where R434 (PPARa) cannot adopt its correct conformation for the effective allosteric communication [Figs. 3(A,B)].

Next, the combinations of all mutants were tested in the presence of 10 nM 9cRA [Fig. 3(C)]. Unsaturated concentration of 9cRA was used in order to see subtle differences between the mutant profiles. Although the basal activities of the mutants showed high differences [Fig. 3(B)], the 9cRA induction was normalized to the solvent for comparison of the levels of induction by 9cRA. The obtained results show induction profiles varying from almost wild-type level (PPARα mutations D387A and R388A) to total loss of induction (for example PPARα mutations E3135A, R434A, and Y468A). Introducing a second mutation can result in decrease [R348A (RXRα) combined with D387A or R388A of PPARa, S427A

(RXRα) combined with R338A of PPARα] or increase [R348A (RXRα) combined with E315A, R434A or Y468A of PPARα, S427A (RXRα) combined with E315 of PPAR α] of the induction level compared to the single mutation. The results show that balanced cowork of multiple residues on the heterodimerization interface is needed for successful communication between the monomers.

Finally some of the PPARα-RXRα mutant combinations were chosen to test the full panel of ligand conditions. The concentration of 9cRA was 10 nM and the concentration of available PPARa selective agonist GW7647 was 100 nM. The reasons for choosing this compound were that GW409544 is commercially not available and both ligands are strong PPARa selective agonists. In addition, the choice of this agonist seems to be in agreement with the recently published PPARy-RXRα crystal structures, which showed no differences between the domain organization of the full length heterodimer when bound to different type of ligands.³⁸ It must be also noted that the MD simulations were done with GW409544, since PPARα has been cocrystallized with this compound and the docking of GW7647 could have led to errors in the initial structure used for MD simulations. The different profiles obtained with full panel of ligand combinations largely vary from phenotypes resembling the wild-type heterodimer [R388A $(PPAR\alpha)$ - wt $(RXR\alpha)$, E315A $(PPAR\alpha)$ - R348A $(RXR\alpha)$, and Y468A (PPARα)- R348A (RXRα)], to profiles where only 9cRA signaling was affected [R434A (PPARα)-S427A (RXRα)]. Also profiles where some of the communication was selectively diminished [wt (PPARα)-R348A (RXR α) and Y468A (PPAR α)- S427A (RXR α)] and profiles where all the communication was blocked [Y468A (PPAR α)- wt (RXR α)] [Fig. 3(D)] were observed.

Two main conclusions can be drawn from this mutation data: (1) All the residues mutated in this study, also those located far away from the helix 12, have a role in the permissive activation of the PPARα-RXRα heterodimer. (2) The effects of the mutations are not additive: combining two point mutations can reverse the effects of the single point mutations alone. In case of PPARα agonist and when 9cRA and GW7647 are combined the situation is even more complex. It has been shown that under these condition PPARa agonists contribute largely to the stabilization of helix 12 activating the heterodimer [Fig. 3(D)].¹⁰

9cRA binding to RXR α increases the stability at the coactivator binding site of PPAR α

RMSFs around the average positions of $C\alpha$ atoms were measured from the last 1 ns of the 2 ns MD simulations. For the fluctuation analysis, the structures were fitted on the more rigid part of the protein (residues 282-324 and 372-434 of PPARα and 274-316 and 364-426 of RXRα) excluding the residues around the LBP. The fitting and the calculation of RMSFs were done separately for each monomer. The statistical significance of the ligand binding induced difference of average RMSF values of each residue was tested with the classical two-tailed t-test. In Figure 4, the areas of the proteins that showed decreased fluctuations in the simulations upon ligand binding are colored blue and the areas, which showed increased fluctuations upon ligand binding are colored red. Residues colored in dark blue and dark red showed averaged fluctuations, which were different in the simulations with and without ligand at 95% confidence level (P-value <

The method was validated performing same analysis for the monomer simulations and comparing the result to the NMR³⁹ and amide HDX¹¹ measurements of ligand binding to PPARy. The results indicate that the MD simulations of this study are capable of describing the main features of the ligand-induced stabilization of the PPARα LBD. For details see Supporting Information (Fig. S5).

The RMSF analyses showed that binding of 9cRA to the RXR\alpha LBP stabilizes helices 3 and 10 and the helix 8–9 loop of PPAR α [Fig. 4(A,B)]. At the coactivator protein binding site of PPARα, the fluctuations are smaller when both ligands bind to the heterodimer [Fig. 4(E)] compared to GW-bound PPARα [Fig. 4(C)]. When both ligands are bound, large areas in the upper part of the RXRα LBD, namely the helices 5, 7, and 10/11, and the PPARα helix 8–9 loop, helix 10/11 and helices 3 and 4 at the coactivator peptide binding site of PPARα are more stable [Fig. 4(E,F)]. This decrease in fluctuations in the PPARα LBD is a result of conformational and dynamical changes at the dimerization surface caused by ligand binding to the RXRα LBP.

Binding of 9cRA to RXRa decreases fluctuations of RXRα helices 5, 7, 10 and the N-terminal half of helix 11 [Fig. 4(A,B) and Fig. 4(E,F) compared to Fig. (C,D)]. The ligand-induced decrease in fluctuations forms a continuous area of residues extending from RXRα over the heterodimer interface to the helix 10/11 and the helix 8-9 loop, which are located at the heterodimer interface, and further to helix 4/5 and helix 3 of PPARa. When GW is bound to PPARα, similar increase of stability is seen in the N-terminal part of helix 10/11, the helix 8–9 loop and helix 4/5 of RXRα, but in contrast to PPARα, the fluctuations of helix 3 at the coactivator binding site of RXRα were increased. Thus, the MD simulations showed that 9cRA binding to RXRα stabilizes the coactivator binding site of PPARα by a helix 12 independent allosteric mechanism and that this allosteric communication is somewhat asymmetrical as binding of PPARa ligand leads to a weaker and somewhat different effect on $RXR\alpha$.

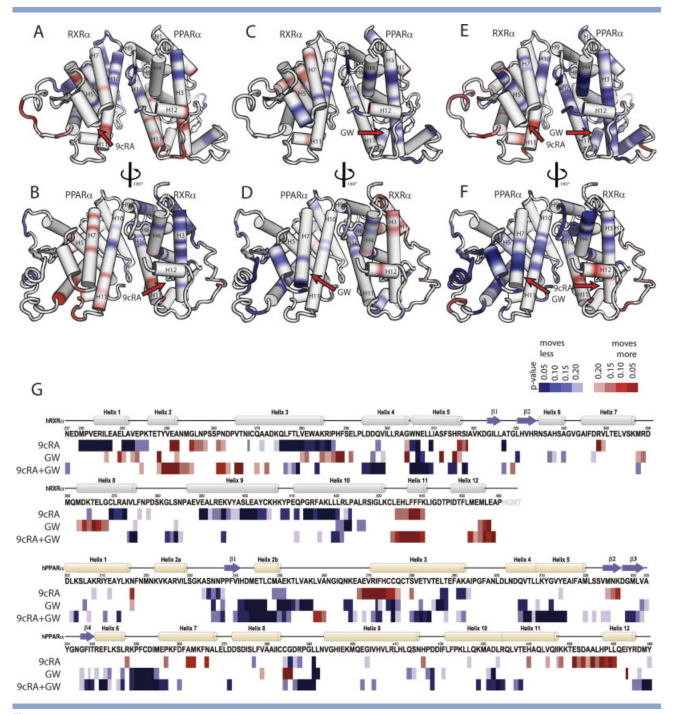


Figure 4

RMSF analyses of the PPARα-RXRα heterodimer MD simulations. Colors: Decrease in the fluctuations upon ligand binding in confidence levels from 80 to 95% (P-values 0.20 to 0.05) is indicated with light blue to blue, and similar increase in the fluctuations with orange to red. Areas where the change in the fluctuations was below the confidence level of 80% are colored white. Coactivator binding site is between the helices 3, 4, and 12. The amount of stabilization varies up to 20%, and the highest decreases of fluctuations are located on the dark blue areas with the lowest P-values. (A,B) Effect of 9cRA binding to RXRα compared to apo simulations. (B: rotated view, PPARα on the left.) Ligand binding to RXRα stabilizes RXRα, but also helix 3 at the coactivator binding site of PPARα. (C,D) Effect of GW binding to PPARα compared to apo simulations. GW stabilizes residues around the LBP and the coactivator binding site. (E,F) Simulations with two ligands compared to apo simulations. The stabilization at the coactivator binding site (E) is stronger than the stabilization with only PPARa ligand (C). (G) The same data shown on the sequence. 9cRA: simulations with 9cRA compared to apo simulations. GW409544: simulations with GW409544 compared to apo simulations. 9cRA + GW409544: Simulations with both ligands compared to apo simulations. The averages of the actual RMSF values are shown in Supporting Information Figure S7 and S8.

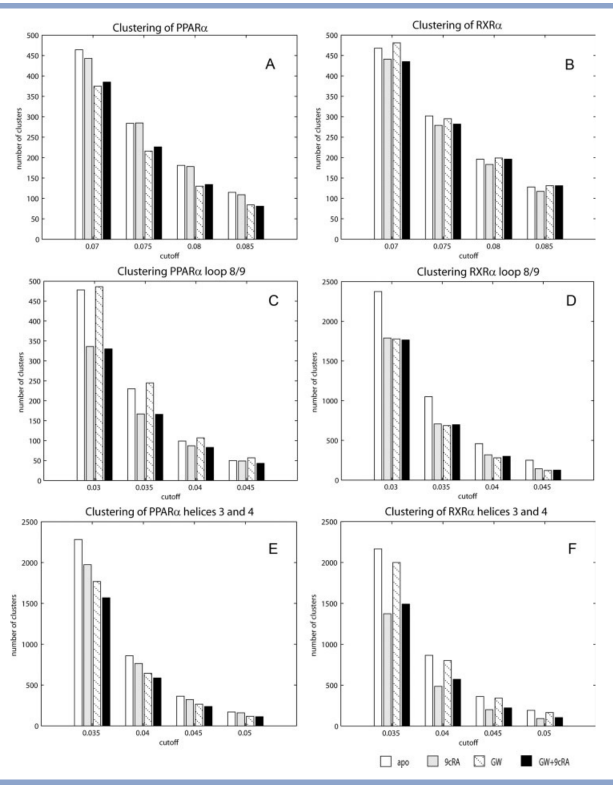


Figure 5

RMSD clustering of LBD, loop 8/9 and helices 3–4 at the coactivator peptide binding site of PPAR α and RXR α in the heterodimer MD simulations. Red: no ligands, orange: 9cRA bound to RXR α , blue: GW409544 bound to PPAR α , green: both ligands. (A) Clustering of the whole PPAR α LBD show smallest number of clusters in simulations with PPAR α ligand (blue and green). (B) Clustering of the whole RXR α LBD show smallest number of clusters in the simulations with RXR α ligand (orange and green). (C) Clustering of PPAR α loop 8/9 gives similar pattern than B, indicating that the PPAR α loop 8/9 is reacting on RXR α ligand, but not on PPAR α ligand. (D) Clustering of RXR α loop 8/9 shows that it reacts on PPAR α ligand if no ligand is bound to RXR α (red compared to blue bar), but also on RXR α ligand (red compared to orange bar). Only weak stabilization with both ligands compared to only RXR α ligand is seen. (E) Clustering of helices 3 and 4 at the coactivator binding site of PPAR α . Both RXR α and PPAR α ligands can stabilize the binding site, and the stabilization is additive: after strong stabilization with PPAR α ligand (red and blue bars) the coactivator binding site is still stabilized by RXR α ligand (green bar). (F) Clustering of helices 3 and 4 at the coactivator binding site of RXR α . This area is stabilized by PPAR α and RXR α ligands, but if RXR α ligand is bound PPAR α ligand cannot stabilize the RXR α coactivator binding site any further.

Preorganization of the coactivator binding site of PPAR α by RXR α ligand binding

The dynamics of the PPARα-RXRα heterodimer was further analyzed with a cut-off-based RMSD clustering method.⁴⁰ In this method, a small number of clusters indicate small mobility or regular motions between a small number of conformations. The composition of the clusters does not necessarily represent any reasonable minimum energy structure, so the contents of the clusters were not further analysed, but the number of clusters was used as a measure of mobility or preorganization.

The clustering was performed for the whole LBD, for the coactivator binding sites and for the helix 8-9 loop located between the coactivator binding site and the heterodimerization interface. For the whole LDBs, larger cut-off values were needed than for clustering of the coactivator binding sites or the helix 8-9 loop, because larger number of atoms and higher flexibility led to larger RMSD values between the structures. All the clusterings were performed with several cut-off values to estimate the sensitivity of the method to the cut-off value. Parallel trajectories, 10 for each system, were pooled to maximize the amount of data in the clustering.

The clustering of the PPARα residues 202-229 and 268-468, excluding the flexible loop area showed a clear difference between systems with and without GW bound to PPAR α : in the GW and GW + 9cRA simulations the PPAR α had a smaller number of clusters than in the apo or 9cRA simulations with all the tested cut-off values [Fig. 5(A)]. This indicates, in line with the fluctuation analyses, that ligand binding to PPARα clearly restrains the movements of PPAR α . Similar results were obtained when RXRa residues 227-241 and 264-458 were clustered: in the 9cRA and GW + 9cRA simulations the RXRα had a smaller number of clusters than in the apo or GW simulations [Fig. 5(B)]. However, 9cRA binding to RXRα had a less pronounced effect on RXRα dynamics than GW binding had on the PPARa LBP. With the two largest cut-off values the difference in the number of clusters in structures from apo- and GW + 9cRA simulations disappeared. In conclusion, the overall movements of the receptor as a whole are controlled by its own ligand, being more notable for PPAR α than RXR α .

Next, clustering was performed for the helix 8–9 loop, which is located between the heterodimer interface and helix 3 of the coactivator binding site. The effect of ligand binding on the number of helix 8-9 loop clusters of PPARα (residues 383–394) was, surprisingly, similar to that obtained for the RXR\(\alpha\) LBD [Fig. 5(B)]: the number of clusters was smallest in the simulations with 9cRA [Fig. 5(C)], whereas binding of ligand to PPARα LBP did not affect the number of clusters at all. The helix 8-9 loop of RXRα behaves differently: the largest number of clusters was seen in the apo simulations, whereas in all the other simulation systems the number of clusters was

similar [Fig. 5(D)]. In addition, there was clearly a smaller number of clusters for the helix 8-9 loop of PPARα than for the same loop in RXRα, indicating higher rigidity for the helix 8–9 loop of PPARα. Thus, this analysis shows that the helix 8-9 loop of PPARα is controlled by RXRa ligand 9cRA and GW has no effect on the loop. The corresponding RXRα loop (residues 376-387) reacts also to 9cRA, presumably making further stabilization with GW difficult when 9cRA is already bound to the RXR α LBP [Fig. 5(D)].

Clustering was also performed for helices 3 and 4 (residues 282-293 and 302-311) of the coactivator binding sites of PPAR α , which was found to be stabilized by RXR α ligand binding in the RMSF analyses [Fig. 4(A)]. Clustering of helices 3 and 4 showed that compared to the apo simulations, ligand binding to PPARα LBP decreases the number of clusters notably [Fig. 5(E)]. However, exactly the same is true for the binding of 9cRA to the RXRα LBP and, interestingly, the effects of these two ligands on helix 3 and 4 clusters of PPARα is additive. Corresponding, although clearly smaller, stabilization of RXRa's coactivator binding site (residues 274-285 and 294–303) was seen by PPARα ligand when compared to the apo case, but no decrease was seen when 9cRA was already bound to RXR α [Fig. 5(F)].

The RMSD clustering analysis shows that even though GW controls the structure and motions of PPAR α as a whole, 9cRA binding to RXR α has a strong stabilizing effect on the coactivator peptide binding site of PPARα and on the helix 8-9 loop, which is located between the coactivator binding site and RXRa. The corresponding helix 8–9 loop of RXRα is also stabilized by the PPARα ligand, and stabilization by PPARa ligand is also seen at the coactivator binding site of RXRα. However, the helix 8–9 loop of RXRα also reacts to its own ligand, and cannot be further stabilized by PPARα ligand when 9cRA is already bound. This is seen at the coactivator binding site of RXR α as well: the coactivator binding site of RXRα is slightly more stable with PPARα ligand than without, but when 9cRA is bound, no extra stabilization is gained by PPARα ligand binding.

If the ligand binding induced changes seen in the clustering analysis are really connected to the permissive and synergistic activation of PPARα-RXRα system, similar intermonomeric communication should not be seen in the nonpermissive NR heterodimer. To test this, calculations were done for the nonpermissive VDR-RXRα heterodimer with 9cRA as a RXR\alpha ligand and vitamin D3 as a VDR ligand using the same MD simulation and clustering protocols that were used for the PPARα-RXRα heterodimer. It was found that RXRa showed similar behavior in all the ligand conditions in both heterodimer systems, whereas 9cRA binding did not induce notable changes in the behavior of VDR. The difference was especially notable in the helix 8-9 loop, which in the case of

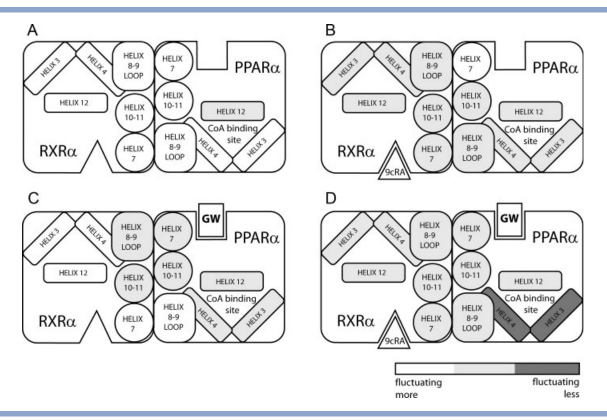


Figure 6 Model of PPARα-RXRα heterodimer activation. (A) Stability of PPARα helix 12 makes the system constitutively active. (B) 9cRA binding leads to permissive activation by stabilizing the coactivator binding site. (C) Binding of GW also stabilizes the coactivator binding site. Even thought not seen in these simulations, stabilization of the helix 12 can play a big role in the activation, depending on the ligand type. (D) Combined effects of both ligands lead to the maximum activation.

VDR did not at all respond to 9cRA binding to RXRα (Supporting Information Fig. S6).

DISCUSSION

The activity of permissive NRs can be regulated also by the ligand of the partner receptor, which is often RXRα, and the heterodimeric complex gives full response when both monomers bind their respective ligands. This type of complex functionality requires the presence of long-range allosteric communication between the LBP of one NR and the LBD of the other NR. At the example of the PPARα-RXRα heterodimer the mechanism of allosteric coupling was investigated in this study. We used MD simulations of PPARα and RXRα LBD monomers as well as PPARα-RXRα LBD heterodimers with different combinations of 9cRA (RXRα ligand) and GW (PPARα ligand) occupying the LBPs of the receptor monomers.

The conformation and dynamics of the C-terminal helix 12 is known to be important for the activity of most of the NRs. Stabilization of helix 12 in the agonistic conformation against the main body of the receptor is needed for recruitment of coactivator proteins. However,

when helix 12 is stabilized into the correct position by an agonist, or when the agonistic position is stable without an agonist, other parts of the coactivator binding site of NR are important for activation as well. This is shown by computer simulations of a mutated GR, 12 by X-ray structures of PPARα/γ agonists that do not make direct contacts with helix 11 or 12^{14,15} and by HDX measurements, which showed that PPARy activation by partial agonists does not necessarily correlate with helix 12 stabilization. 11 In the latter case, the helix 12 independent activation seems to be related to stabilization of helices 3 and 4, which form the two other sides of the coactivator binding cleft, and the β-sheet region. In addition, the distinct functional properties of selective PPARy modulators have been linked to a helix 12 independent mechanism and subtle changes influencing the dynamics of the LBD.41

In our computer simulations of the PPAR α -RXR α heterodimer an allosteric signal transmission was seen between the LBP of RXRα and the coactivator binding site of PPARα. Namely, when 9cRA was bound to the RXRα LBP, notable stabilization of helices 3 and 4 at the coactivator binding site of PPARa was observed [Figs. 5(A) and 6(E)]. It was also seen that even though binding of GW stabilizes the coactivator site of PPARα, binding of 9cRA to RXRα can stabilize it further. The stabilization was seen in the reduced RMSF values and smaller number of clusters in the structural clustering analysis. The binding of RXRα ligand led to small changes in the average positions of helix 7 and helix 11, seen in decreased helix 11 - helix 11 and helix 7-C-terminus distances over the heterodimer interface. Thus, these structural changes increased packing of the heterodimerization interface and enhanced propagation of the ligandinduced structural/dynamical changes from RXRa LBD over the interface to PPAR α .

The effect of RXR α ligand binding on PPAR α is clearly seen in the behavior of the PPARa's helix 8–9 loop, which is located at the heterodimer interface between RXR α and the coactivator binding site of PPAR α . Although the PPAR α LBD as a whole is stabilized by its own ligand [Fig. 5(A)], the clustering analysis showed that the movements of the PPARα helix 8-9 loop is controlled by RXR α ligand binding [Fig. 5(C)].

Similarly to PPARα, the helix 8–9 loop and the coactivator binding site of RXRa reacts to ligand binding to PPARα LBP [Fig. 5(D,F)]. However, the helix 8–9 loop of RXRα reacts also to RXRα ligand binding [Fig. 5(D)]. In addition, it is also visible that ligand binding to the PPARα LBP, if 9cRA already occupies RXRα LBP, does not further stabilize the helix 8-9 loop [Fig. 5(D)]. The stabilization of RXR\alpha coactivator binding site is markedly different from that observed for PPARα: a notable decrease in the number of clusters is seen only when RXRα's own ligand is bound, and no further decrease is caused by PPARα ligand binding [Fig. 5(F)]. The functional model from this comparison shows that although there are some similarities between RXRα and PPARα with respect to ligand binding to partner protein, a synergistic effect is seen only in the case of PPARα. In addition, the absence of RXRα ligand binding-induced changes in the simulations of the nonpermissive RXRα-VDR system supports the conclusion that a helix 12 independent mechanism seen in the PPARα-RXRα simulations has a role in allosteric activation of NRs.

On the basisi of the results of this work, we created a simplified model of PPARα-RXRα heterodimer activation (Fig. 6). If LBPs of both PPARα and RXRα are unoccupied, stability of PPARα helix 12 makes the system constitutively active [Fig. 6(A)]. Binding of 9cRA decreases fluctuations of large parts of RXR α [Fig. 6(B)]. Increased packing at the heterodimer interface and decreased fluctuations of RXRα helices 10–11 and 7 stabilize those areas of PPARα, which they are in contact with: helix 10/ 11 and the helix 8–9 loop. The helix 8–9 loop is in contact with the helix 4 of the coactivator binding site. The fluctuations at the coactivator binding site are decreased, which preorganizes the system for coactivator binding and the system displays permissive activation. Binding of GW also stabilizes the coactivator binding site [Fig.

6(C)]. Helix 12 stabilization upon GW binding was not seen in these simulations, but depending on the type of ligand, helix 12 is expected to be more stable as well (not shown in the model). 11 Helices 3 and 4 of RXRα are not stabilized. Combined effect of GW and 9cRA lead to strong stabilization of the coactivator binding site and the system displays synergistic effect [Fig. 6(D)].

The recent long-awaited crystal structure of the fulllength PPARγ-RXRα complex on DNA provided important data on the domain organization of the heterodimer in 3D space leading to functional implications, but it was unable to provide a direct explanation for the phantom ligand effect or for receptor permissivity.³⁸ The MD simulations suggested that the allosteric signaling between the monomers results from subtle structural and dynamical changes of the whole heterodimerization surface rather than on individual residues. This observation was supported by mutation data presented here: large set of mutations on the heterodimerization interface affect signal transmission.

The data in this study suggests that allosteric activation of the PPARα-RXRα heterodimer is due to relatively small structural and dynamical ligand binding-induced changes rather than large conformational changes in the heterodimer structure. This mechanism is reminiscent of the dynamic allostery model of the cooperative oxygen binding to hemoglobin proposed by Yonetani and Laberge.² According to this model, the dynamics of hemoglobin rather than quaternary structural changes play the central role in regulation of oxygen affinity. The cooperativity is due to a decrease in hemoglobin backbone flexibility caused by hydrogen bonding or steric hindrance by the ligand, which leads to increased reactivity of the heme. Analysis of long MD trajectories by methods based on cross-correlation and various amino acid network analyses have been used to obtain detailed knowledge about allosteric mechanism and communication pathways in protein systems. 42-46 Such analysis methods could also be applied to obtain more detailed picture on the allosteric mechanism and identify possible residue level communication pathways in NR systems. It is likely that increased possibilities to use MD simulations for large proteins will reveal that protein motions have a more widespread role in the mechanisms of allosteric systems. The understanding of allosteric mechanisms may in fact, as suggested in the recent paper from Lee et al., lead to creation of regulatory activities into engineered multidomain protein systems.⁴⁷

The statistical coupling analysis (SCA) is a sequence analysis method designed to detect coevolution of position pairs in protein sequences.⁴⁸ Because coevolution of two positions is likely to be driven by functional interaction of the residues, it is likely that such residue pairs are involved in allosteric networks. It has been shown that results of SCA are consistent with the known allosteric mechanisms in several protein families. 48,49 SCA has been applied to

study the allosteric ligand activation (i.e., permissivity) in RXRα heterodimers of several NRs and predicted a group of amino acid residues outside the helix 12 that are important for allosteric signaling in NR heterodimers.³ Functional analysis of these residues demonstrated that they belonged to an allosteric network as their mutation weakened or destroyed RXRa agonist induced allosteric activation of several permissive NRs.³ These amino acid residues are located mainly in helices 3 and 4 and the helix 8-9 loop. The same region, the upper half of PPAR α , was seen to be stabilized by 9cRA binding to RXRα LBP. Also, the number of clusters of helix 8-9 loop and the helices 3 and 4 were observed to decrease in response to ligand binding to RXR α LBP in this study.

The results of this study suggest that permissive and synergistic activation of the PPARα-RXRα heterodimer is mediated by the same mechanism: stabilization of the PPARα coactivator peptide binding site, which is a result of conformational changes in both LBDs caused by ligand binding. The overall ability of a NR heterodimer to respond to RXRα ligand can be dependent on the stability or rigidity of its coactivator binding site. HDX measurements show that this stability can be modulated by ligands. 11 It would be interesting to see, whether PPARy shows reduced synergistic activation with the helix 3 stabilizing partial agonists, such as BVT.13, when compared to ligands that do not stabilize helix 3.

CONCLUSIONS

The allosteric activation of the PPARα-RXRα heterodimer LBDs was studied with MD simulations in the apo form, and in complex with 9cRA, GW and with both 9cRA and GW. Ligand binding to RXRα was seen to decrease fluctuations on a continuous area of residues extending from the ligand binding pocket of RXRa through the helix 8–9 loop of PPARα, which is located at the heterodimerization interface, to the helices 3 and 4 at the coactivator binding site of PPAR α . The function of the residues identified from MD simulation was tested experimentally, providing evidence that the exchange of information within the heterodimer involves a complex network with alternative, substitutable interactions. These results suggest a helix 12 independent mechanism for permissive and synergistic activation of NR heterodimers by coactivator binding site stabilization.

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