

Drug Design

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Designed Spiroketal Protein Modulation

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Abstract: Spiroketal structures are structural motifs found in many biologically active natural products, which has stimulated considerable efforts toward their synthesis and interest in their use as drug lead compounds. Despite this, the use of spiroketals, and especially bisbenzannulated spiroketals, in a structure-based drug discovery setting has not been convincingly demonstrated. Herein, we report the rational design of a bisbenzannulated spiroketal that potently binds to the retinoid X receptor (RXR) thereby inducing partial co-activator recruitment. We solved the crystal structure of the spiroketal-hRXR α -TIF2 ternary complex, and identified a canonical allosteric mechanism as a possible explanation for the partial agonist behavior of our spiroketal. Our co-crystal structure, the first of a designed spiroketal-protein complex, suggests that spiroketals can be designed to selectively target other nuclear receptor subtypes.

Spiroketal structures are archetypal spirocyclic compounds,^[1] found abundantly in the CAS registry (Supporting Information), of which many are bioactive natural products.^[2,3] Besides being attractive targets for total synthesis,^[4-6] spiroketal-derived natural products have contributed to a renaissance of thinking towards intelligent library design, grounded in principles of diversity-oriented synthesis (DOS)^[7] and biology-oriented synthesis (BIOS).^[8,9] Spiroketal structures are arguably well-adapted to both philosophical approaches in being biologically relevant^[2,3] while fulfilling the three criteria for molecular diversity set out by Schreiber and co-workers—appendage, stereochemistry, skeletal diversity^[7]—owing in particular to the conformational and configurational flexibility of these

scaffold structures. A BIOS study on spiroketals performed by Waldmann and co-workers^[10-12] clearly placed these structures within a hierarchical classification of bioactive scaffold structures, navigable with the help of cheminformatics tools such as Scaffold Hunter,^[13] and inspired the design of other spiroketal libraries for phenotypic screening.^[13,14]

Recently, 3D-pharmacophore modelling was used to design spiroketal-derived sugars, which showed inhibitory activity towards SGLT2.^[19] To the best of our knowledge, the co-crystal structure of bistramide A bound to actin,^[20] and the bis-spiroketal pinnatoxins A and G^[21] are to date the only examples of spiroketals bound to protein targets. Benzannulated spiroketals are a relevant subtype of spiroketals,^[3] which includes the antibiotic rubromycin family (Figure 1A).^[22]

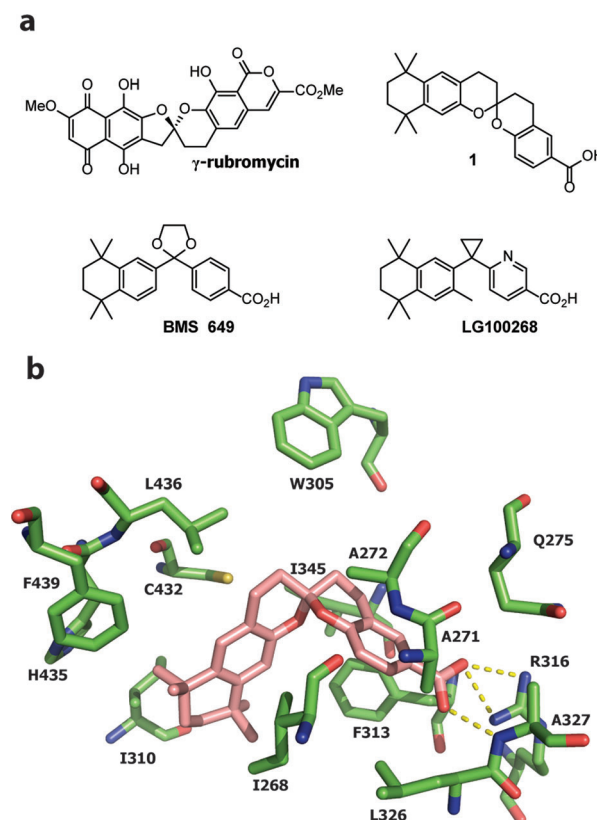


Figure 1. Design of a spiroketal as RXR ligand. a) Molecular structures of γ -rubromycin, [6,6]-bisbenzannulated spiroketal **1** and RXR full agonists, **BMS 649**^[16] and **LG100268**. b) Top-ranked pose of **R-1** generated by docking into the space occupied by **BMS 649** in the hRXR α -**BMS 649** co-crystal structure (PDB code: 1MVC)^[16] using the FlexX docking module in the LeadIT suite^[17] followed by HYDE scoring in SEESAR.^[18] **R-1** skeleton: C = pink, O = red; protein backbone: C = green, N = blue, O = red, S = yellow; dashed lines = H-bonding interactions below 3.3 Å.

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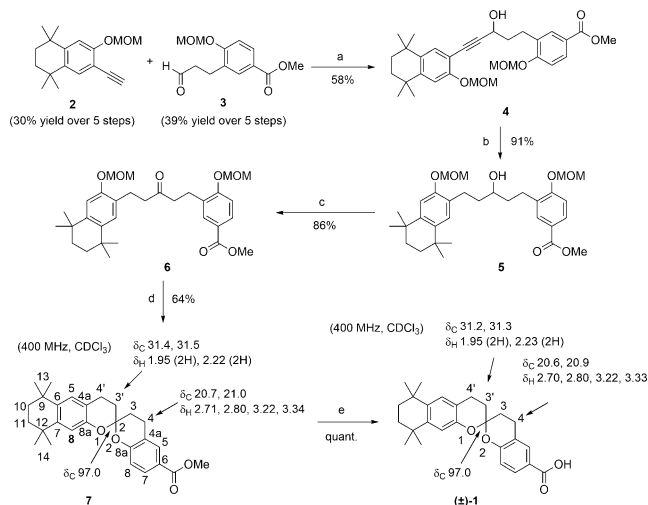
Recent reports have revealed that these molecules function as telomerase inhibitors, with the [6,5]-spiroketal ring system in this case playing an essential role in the rubromycin's pharmacophore.^[22] Despite the abundance of spiroketals and their highlighted potential as medicinal chemistry scaffolds, the structure-based design and structural validation of spiroketals as medicinal chemistry scaffolds has not been clearly demonstrated.

A reported crystal structure of a [6,6]-bisbenzannulated spiroketal^[23] inspired us to consider these structures as nuclear receptor (NR) ligands, which we speculated would possess the correct size, shape, and hydrophobicity to target the L-shaped ligand binding pocket (LBP) of the retinoid X receptor (RXR),^[24] a member of the superfamily of gene transcription factors. RXR plays a central role in hormone-driven cell-signaling events through its ability to heterodimerize with other type II nuclear receptors.^[25] RXR is a drug target for the treatment of cutaneous T-cell lymphoma and is under investigation as a potential treatment for Alzheimer's Disease.^[26] Despite the fact that RXR ligands have been thoroughly investigated,^[27] only a very few RXR partial agonists with limited structural diversity have been characterized, and rules guiding the design of heterodimer-specific RXR ligands are essentially non-existent. In part as a response to these challenges, and in continuation of our group's recent efforts to identify selective RXR^[28] and other NR modulators,^[29] we report the first designed spiroketal protein modulator, as exemplified by RXR modulation.

To assess the potential of bisbenzannulated spiroketals to target RXR, we adopt a classical scaffold-hopping approach^[30] commencing with the co-crystal structure of commercial nanomolar-potent RXR full agonist **BMS 649** (otherwise known as SR11237)^[31] bound to hRXR α (PDB code: 1MVC).^[16] We then replaced the rigid acetal linker present in **BMS 649** with a [6,6]-bisbenzannulated spiroketal linker while retaining the key tetramethyl-tetrahydronaphthyl- and carboxylic acid groups,^[32] to generate **1**. We then modelled and docked both enantiomers of **1** into the space occupied by **BMS 649** in the LBD of the hRXR α -**BMS 649** co-crystal structure^[16] using the FlexX docking module in the LeadIT suite followed by evaluation using the scoring function HYDE in SEESAR (Figure 1). While all attempts at docking the *S*-enantiomer in this PDB structure failed to generate poses, we succeeded in docking the *R*-enantiomer, with the best pose shown in Figure 1B. Although our docking studies did not take into account the thermodynamic preferences of the spiroketal ring system, interestingly, *R*-**1** adopts a diaxial ring conformation, which would be favored owing to bis-anomeric stabilization. In this ring conformation favorable polar interactions are maintained between the carboxylic acid of *R*-**1**, Arg316, and the backbone of Ala327.

To enable an expedient testing of our binding hypothesis, we elected for a racemic synthesis of (\pm)-**1**, with a view to separating the enantiomeric spiroketals by chiral HPLC at a later stage. In reported syntheses of [6,5]-^[33,34] and [6,6]-bisbenzannulated spiroketals,^[3] the spiroketal core is frequently formed under thermodynamically driven conditions through a dehydrative ring cyclisation. Our synthesis of (\pm)-**1** was based on work by Brimble and co-workers on analogous

[6,6]-bisbenzannulated spiroketals^[23] and is described in Scheme 1. We reacted aldehyde **3** with the lithium acetylide of **2** to obtain alkynol **4** in a reasonable 58% yield. After subsequent catalytic hydrogenation of **4**, we performed



Scheme 1. Reagents and conditions: a) *n*-BuLi, THF, -78°C to RT; b) 10% Pd/C, KHCO₃, EtOAc, RT; c) Dess–Martin periodinane, CH₂Cl₂, RT; d) TMSBr, CH₂Cl₂, -30°C to RT; e) NaOH, dioxane/MeOH, 40°C . Characteristic ¹H and ¹³C resonance peaks are summarized for **7** and (\pm)-**1**.^[23]

a Dess–Martin oxidation on the resulting secondary alcohol **5** to generate the spirocyclization precursor, **6**, which we treated with trimethylsilyl bromide to effect a one-pot deprotection/cyclization, which produced the [6,6]-bisbenzannulated spiroketal **7** in a yield of 64%. The synthesis of (\pm)-**1** concluded with a straightforward base-mediated hydrolysis of the methyl ester group. The resonance peak at δ_C 97.0 ppm in the ¹³C-NMR spectra of **7** and (\pm)-**1** is diagnostic for the spirocarbon, while the ¹H resonances corresponding to the two sets of diastereotopic protons H³, H^{3'}, H⁴ and H^{4'} suggest that the spiroketal adopts a diaxial ring conformation (Supporting Information) similar to analogous structures.^[23]

We profiled the RXR α -activity of (\pm)-**1** alongside full agonist **LG100268** (Figure 1A)^[36] in a fluorescence-based cofactor recruitment assay (Figure 2, left and Table 1). As expected,^[28] **LG100268** induced potent recruitment of the D22 peptide^[37] with an EC₅₀ = $0.10 \pm 0.01 \mu\text{M}$. Intriguingly, (\pm)-**1** was also active, with an EC₅₀ = $0.73 \pm 0.06 \mu\text{M}$, and additionally exhibited a partial agonist behavior, as judged by the levelling off of polarization at 53% of the maximum response induced by **LG100268**. We tested (\pm)-**1** against two other LXXLL-derived peptides (Table 1), ribosome display peptide Pro22^[35] and the naturally occurring peptide TIF2,^[38] and observed similar EC₅₀ values but different % efficacies. One of the separated enantiomers, **1-ent1**, was found to approach the potency of **LG100268** in the same FP assay (Figure 2, left and Table 1). Furthermore, **1-ent1** displayed seven-fold higher potency and two-fold higher % efficacy than the other enantiomer, **1-ent2**. The isolated enantiomers did not evidently epimerize under the acidic separation

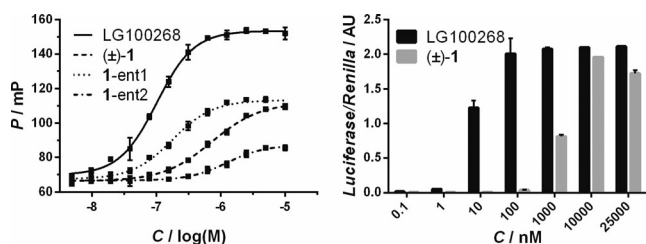


Figure 2. Biochemical and cellular evaluation of (±)-1. Left: Fluorescence polarization assay data showing that full agonist **LG100268** induces binding of the fluorescently labelled D22 peptide in a concentration-dependent manner, while (±)-1, **1-ent1**, and **1-ent2** (separable by chiral HPLC) each exhibit a partial agonist behavior. Right: Cellular activities of **LG100268** and (±)-1 measured in a mammalian two-hybrid luciferase assay. Error bars denote s.d. ($n=3$).

Table 1: Summary of FP and cellular M2H data.

Compound	Peptide ^[a]	Fluorescence Polarization	
		EC ₅₀ (±) [μM]	% _{eff}
LG100268	D22 ^[b]	0.10 (0.01)	100
	TIF2 ^[c]	0.48 (0.02)	100
	Pro22 ^[c]	0.25 (0.01)	100
(±)-1	D22 ^[b]	0.73 (0.06)	53
	TIF2 ^[c]	0.47 (0.06)	28
	Pro22 ^[c]	0.43 (0.01)	59
1-ent1	D22 ^[b]	0.17 (0.02)	54
1-ent2	D22 ^[b]	1.20 (0.33)	26

[a] Peptide sequences: D22 = FAM-LPYEGSLLLKLLRAPVEEV; TIF2 = FITC-Ahx-KHKILHRLQLQSS-NH₂; Pro22 = FITC-Ahx-LTARHPLLMRLLLSPS-NH₂.^[35] [b] See Figure 2. [c] Refer to the Supporting Information for the binding curves.

conditions (Supporting Information) nor was any significant change in EC₅₀ value observed for either **1-ent1** or **1-ent2** in the same FP assay over a 24 h period (Supporting Information) as evidence of the stability of these compounds under the aqueous assay conditions. Our findings are consistent with those from studies on similar benzannulated spiroketals^[39] and the fact that analogous natural products are isolated as single enantiomers.^[40]

To evaluate **1** under more biologically relevant conditions, we compared (±)-1 and **LG100268** in a mammalian two-hybrid (M2H) luciferase assay (Figure 2, right). As expected,^[28] **LG100268** produced a full and potent concentration-dependent response (Figure 2, right). (±)-1 was also active under the assay conditions, though less potent than **LG100268**, and the partial response less emphatic than that observed in the FP assay. The difference in % efficacy observed for (±)-1 in both the in vitro FP and cellular M2H assays might be explained by differences in protein concentration between the two assay formats. Nevertheless, we could conclude that (±)-1 is cell permeable and active toward gene transcription similarly to an established RXR full agonist.

To provide a structural explanation for the observed RXR activity, we co-crystallized (±)-1 with the hRXRα LBD-TIF2 peptide complex and solved the structure to a resolution of 2.17 Å (Figure 3). Globally, the protein adopts a canonical,^[41] agonistic, folded state with the helix co-activator bound to the AF2 (Figure 3A). Closer inspection of the RXR LBP

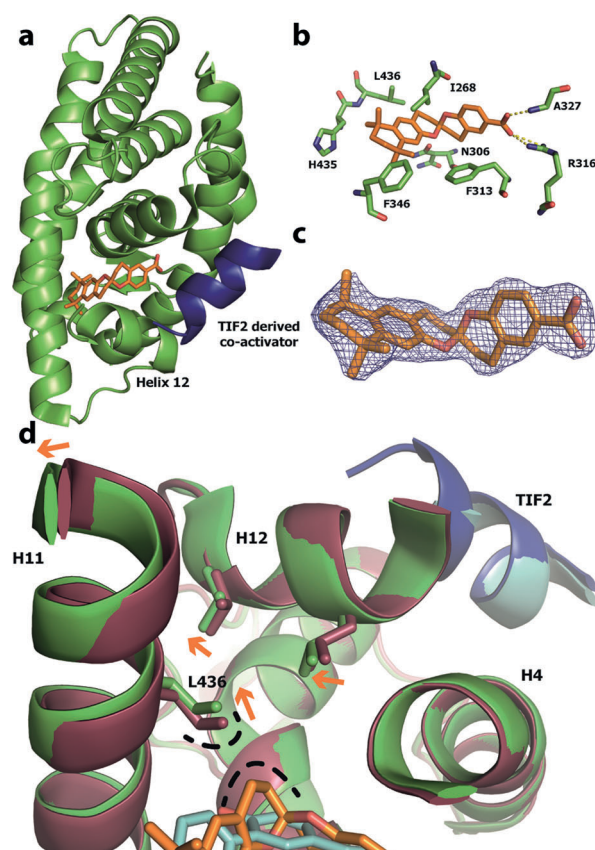


Figure 3. X-ray co-crystallography data. a) Ribbon representation of the X-ray co-crystal structure of **R-1** (orange stick) bound to the ligand-binding pocket (LBP) of hRXRα (green ribbon) with the TIF2 co-activator-derived peptide (blue ribbon) PDB code: 5LYQ. b) Enlarged view of the hRXRα LBP with the amino acid side chains labelled and represented as sticks. c) Superimposition of **R-1** and final $2F_o - F_c$ electron density map (contoured at 1σ). d) Superimposition of the LBP region of hRXRα bound to **R-1** (protein in green, ligand in orange, PDB code: 5LYQ) and a documented full agonist (protein in red, ligand in cyan, PDB code: 40C7).^[28] The TIF2 peptide corresponding to PDB code 40C7 is shown in cyan. Participating helices/residues/atoms are labelled. Orange arrows indicate movement in protein conformation from an agonistic (red) to the folded state induced by **R-1** (green). The loop region connecting helix 11 (H11) to H12-443GDTPID448 is hidden for clarity.

revealed clear electron density indicative of a single spiroketal molecule. On closer inspection, the *R*-enantiomer, with the spiroketal ring system in a bis-anomeric diaxial conformation, fitted best in the electron density (Figure 3C and Supporting Information). The carboxylate group of **R-1** engages in a canonical polar interaction with the side chain of Arg316 and hydrogen bonds with the protein backbone at residue Ala327 (Figure 3B). Interestingly, despite our inability to generate docking poses for **S-1** in our initial model (Figure 1), we were able to dock **S-1** into the space created by **R-1** in the LBP of our hRXRα-**R-1** co-crystal structure (Supporting Information). The contrast between this success and the docking studies highlights an imperfection of molecular docking on account of the dynamic behavior of protein folding and the need for caution when interpreting docking results. Nevertheless, in contrast to **R-1**, our best docking pose

for **S-1** placed the spiroketal ring system in a thermodynamically less-favorable axial–equatorial conformation. Therefore, we logically assign **1-ent1** to be **R-1** and **1-ent2** to be **S-1**, and tentatively speculate that the weaker activity observed for **S-1** in the FP assay may result from a protein-induced fit of the spiroketal to the RXR LBP via the postulated axial–equatorial conformation.

In search of a plausible structural basis for the RXR activity of our spiroketals, we superimposed the co-crystal structure of hRXR α –**R-1**–TIF2 with a previously published crystal structure of hRXR α –TIF2 bound to a potent full agonist (Figure 3D).^[28] Compared to the full agonist, the binding of **R-1** results in a circa 1 Å displacement of the side chain of Leu436 in H11 and the C-terminal region of H11, which would perturb H12 binding, and potentially destabilize coactivator peptide binding as a possible cause of the partial agonist effect observed in the FP data. A similar sequence of side-chain displacements has been cited by Nahoum et al. to explain the partial agonist behavior of structurally different biaryl RXR ligands reported,^[42] thus hinting at a general mechanism for RXR partial agonism.

In conclusion, we report the structure-based design, synthesis, and biochemical as well as structural evaluation of a bisbenzannulated spiroketal as a potent modulator of the RXR α gene transcription factor. Our work includes a rare co-crystal structure of a spiroketal,^[20,21] which is to the best of our knowledge the first of a benzannulated spiroketal bound to a protein target. We believe that the apparent RXR partial agonist behavior of our spiroketal contributes to establishing partial agonism^[43] as a concept for RXR.^[42,44] Furthermore, the high structural homology of the LBP across the NR superfamily,^[41] combined with the structural versatility of spiroketals, suggests that spiroketals can be designed to selectively target other NR subtypes as forerunner to the designed modulation of other protein targets.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: drug design · drug discovery · natural products · spiro compounds · structure elucidation

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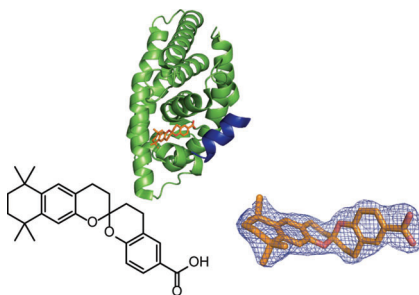
Communications



Drug Design

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Designed Spiroketal Protein Modulation



Spiroketals for drug discovery: The structure-based design, synthesis, and biochemical as well as structural validation of a spiroketal protein modulator are reported. The data suggests that the bisbenzannulated spiroketal functions as a potent partial agonist of the retinoid X receptor thus highlighting the potential of spiroketals as de novo drug lead compounds.