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Fenretinide Derivatives Act as Disrupters of Interactions of Serum Retinol Binding Protein (sRBP) with Transthyretin and the sRBP Receptor

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Supporting Information

ABSTRACT: Serum retinol binding protein (sRBP) is released from the liver as a complex with transthyretin (TTR), a process under the control of dietary retinol. Elevated levels of sRBP may be involved in inhibiting cellular responses to insulin and in generating first insulin resistance and then type 2 diabetes, offering a new target for therapeutic attack for these conditions. A series of retinoid analogues were synthesized and examined for their binding to sRBP and their ability to disrupt the sRBP—TTR and sRBP—sRBP receptor interactions. A number inhibit the sRBP—TTR and sRBP—sRBP receptor interaction and perhaps offering a new therapeutic intervention against type 2 diabetes and its development. Shortening the chain length of the FEN derivative substantially abolished binding to sRBP, indicating that the strength of the interaction lies in the polyene chain region. Differences in potency against the sRBP—TTR and sRBP—sRBP receptor interactions suggest variant effects of the compounds on the two loops of sRBP guarding the entrance of the binding pocket that are responsible for these two protein—protein interactions.



INTRODUCTION

Transport of retinol [ROH (Figure 1)] in the blood is controlled by the lipocalin serum retinol binding protein (sRBP, 21 kDa), which is synthesized and released from the liver as a complex with a second protein, transthyretin (TTR).^{1–3} sRBP consists of an N-terminal coil, eight antiparallel β -strands (A-H) forming a barrel, and a short α -helix close to the C-terminus [Protein Data Bank (PDB) entry 1RBP⁴].¹ ROH is accommodated in the barrel with the β -ionone ring positioned innermost, the polyene chain fully extended, and the hydroxyl end group nearly solvent-exposed near the A–B, C–D, and E–F loops (Figure 1). Circulating in the plasma, sRBP is bound to TTR, a homotetramer of \sim 56 kDa. Each TTR monomer consists of eight antiparallel β -strands (A–H) arranged like a Greek key β -barrel. One tetramer of TTR can bind two sRBP molecules in vitro;⁵ however, because of a significantly higher concentration of TTR compared to that of sRBP, each tetramer forms a 1:1 complex in plasma.⁶ This association of the sRBP-ROH complex with TTR is believed to prevent its filtration through the kidney glomeruli.³ In the blood, the uncomplexed or free form of sRBP interacts with a receptor in the plasma membrane of virtually all cells in a ROH-sensitive manner.^{7,8} Only the TTRdissociated holo-sRBP is able to bind to the receptor with high

affinity, because of steric hindrance of access to the sRBP–sRBP receptor binding site.^{7–10} Binding of sRBP to its receptor is followed by the release of retinol and its uptake into the cell via the sRBP receptor.

A notable paper¹¹ has chronicled the evidence of the involvement of sRBP in obesity, insulin resistance, and type 2 diabetes. The level of sRBP is elevated in insulin resistant mice and humans with obesity and type 2 diabetes.¹² sRBP is mostly secreted by the liver but can also come from other tissues, including adipocytes.¹³ The source of this excess sRBP is thought to be visceral adipose tissue, hence the link to obesity. sRBP acting on peripheral tissues such as skeletal muscle is postulated to attenuate insulin sensitivity, as indicated by reduced IRS1 phosphorylation and PI3K activity.¹¹ The mechanism of this proposed effect is unknown but should involve a sRBP-binding entity of some kind.

Fenretinide [4-hydroxy(phenyl)retinamide (FEN) (Figure 1)] is a synthetic retinoid and has been investigated for potential use in the treatment of cancer, cystic fibrosis, rheumatoid arthritis, acne, psoriasis, and Stargardt's disease.^{14,15} Previously, FEN has

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Figure 1. (A) Three-dimensional image of ROH (blue) in sRBP (PDB entry 1RBP⁴) and FEN (yellow) in sRBP (PDB entry 1FEL¹⁶) with contacting residues labeled and illustrated in stick format. Images are shown with a 6° rotation around the *y* axis. Images produced using Pymol.³⁹ (B and C) Two-dimensional (2D) depiction of ROH and FEN binding poses as depicted using MOE.²⁰ (D) Legend for the 2D interactions in the binding pose as depicted using MOE.

been shown to compete with retinoids for binding to sRBP (PDB entry 1FEL), 16 disrupting sRBP-TTR complexes and resulting

in urinary secretion of sRBP and systemic depletion of ROH.¹⁷ Administration of FEN has been observed to exert therapeutic

effects in mouse models of obesity and diabetes.¹⁵ Long-term FEN treatment prevents high-fat diet-induced obesity, insulin resistance, and hepatic steatosis.¹⁸

The hypothesis we are exploring is that agents that reduce sRBP levels or disrupt the protein interactions that sRBP undergoes will prevent the genesis of insulin resistance and consequently that of type 2 diabetes. In this work, we developed a series of retinoid analogues that were first evaluated experimentally for binding to sRBP through fluorescence measurements. Subsequently, their ability to disrupt protein-protein interactions was examined through pull-down, timeresolved fluorescence resonance energy transfer (TR-FRET) and surface plasmon resonance (SPR) assays of sRBP-TTR complexes and SPR assays of sRBP with solubilized membranes containing the sRBP receptor. The latter is a novel focus in an effort to identify compounds that exhibit a dual or different effect. Compounds that exhibit differences in their ability to inhibit sRBP-TTR and sRBP-sRBP receptor interactions may provide an approach to examining both the validity and the mechanism of action of sRBP levels in the context of insulin resistance and type 2 diabetes.

RESULTS

Retinoid Analogues. We were largely interested in establishing if the retinoids could bind to sRBP without necessarily penetrating deep into the binding pocket. We also wished to investigate the level of variability allowed at the region of the molecule near the entrance to the binding cavity, thought to be the effective moiety with regard to alterations in conformation. We were also interested in determining if steric hindrance was the sole and main mechanism for inhibition or whether a conformational change was also required.

Traditional medicinal chemistry around both ends of a known chemical scaffold (ROH/FEN) was utilized to develop a novel series of retinoid analogues (Figure 2). The target compounds 1-68 were prepared as outlined in Schemes 1-3 (see Experimental Section), and all chemical structures are shown in the Supporting Information.

The approach of the analysis of these compounds was structured on three levels. The first binding to sRBP was monitored using quenching of the fluorescence of the intrinsic tryptophan in sRBP and/or inhibition of the FRET response of



Figure 2. 2D image of Fenretinide (FEN), which differs from ROH by a phenylamide in place of the hydroxyl group. (A) Area where the variations were introduced at the β -ionone ring end of FEN (compounds **2–22**). (B) Polyene end of FEN where variations were introduced (compounds **1** and **23–68**).

Scheme 1

bound retinol. The second involved the sRBP-TTR interaction as monitored by pull-down and SPR responses. The third monitored sRBP-sRBP receptor interactions in solubilized membrane preparations using SPR. Only compounds that provided some evidence of binding to sRBP progressed to the second and third stages.

Binding of Compounds to sRBP. To study the interaction between retinoid analogues and sRBP, the quenching of the intrinsic tryptophan (Trp) fluorescence of the apoprotein following excitation at 280 nm was measured in the presence of increasing concentrations of the compounds. ROH binds to the ligand binding site of the sRBP and produces a quenching of the Trp fluorescence. Those compounds that absorbed in the emission region of Trp reduced this emission in a dose-dependent manner, indicative of their specific interaction with aposRBP (Figure 3). The fluorescence intensities of the compounds in an N-acetyltryptophanamide solution were measured to correct the titration curves. The change in fluorescence intensity was used to calculate the apparent dissociation constants (K_d) for the protein-compound interactions as described by Cogan et al.¹⁹ The K_d values for selected compounds are listed in Table 1. A K_d of 182 \pm 4 nM was obtained for the ROH-sRBP complex, in agreement with that reported in the literature for the protein isolated from human plasma.¹⁹ The other compounds had an affinity in the same range (Table 1).

Effect of Compounds on the sRBP-TTR Interaction Determined by a Pull-Down Assay. The ability of the 68 library compounds to disrupt the interaction between sRBP and its partner, TTR, was first examined using a pull-down assay. The purified ROH-sRBP-TTR complex, as indicated in Experimental Section, was utilized. Both proteins were incubated together in the presence of ROH and the various compounds, at 10 and 100 μ M, respectively. The difference in concentration between ROH and the compounds allowed for the detection of their inhibitory effect. After incubation with the different compounds, the flow-throughs were run in SDS gels and the free sRBP disrupted from the complexes was detected by silver staining. A representative result for ROH and FEN is shown in Figure 4A. FEN, which is known to disrupt the interaction between sRBP and TTR, was used as a control to which potential positive compounds could be compared. A small quantity of free sRBP was always obtained in the negative control flow-through (no compound added), as a consequence of the relatively weak interaction ($K_{\rm d} \approx 0.3 \ \mu {\rm M}$) between the two proteins.²¹ FEN produced an estimated 3-5-fold increase in the amount of sRBP released from the complex. The shorter retinoid analogues, 2-22, which did not exhibit absorption in the Trp emission region, also did not show the ability to disrupt the sRBP-TTR interaction and were disregarded in further studies. A number of compounds based on scaffold B (1 and 23-68) showed a similar disruptive effect on the complex with FEN. The dose-dependent effect of the positive compounds was similarly assessed using increasing concentrations of the analogues, as shown for one selected compound, **58** (Figure 4B).



sRBP-TTR Interaction Assessed by TR-FRET. To estimate the potency of the different inhibitors selected in the pull-down assay, we continued our analysis of the compounds using a TR-FRET assay based on a methodology developed previously.²² The sRBP used in this assay was a recombinant protein expressed with a His tag in *Pichia pastoris*,²³ to which an europium anti-His antibody was bound. Native TTR was labeled with biotin. When both proteins bind, and after excitation of europium with light at 340 nm, energy is transferred from europium to streptavidin-conjugated allophycocyanin (APC) and the emission of light can be detected at 665 nm. A stronger signal is observed as the affinity between both proteins increases with ROH concentration. The TR-FRET signal was measured against increasing concentrations of ROH and a constant concentration of inhibitor. The system was tested first with FEN from 0.1 to 50 μ M, and from this experiment, an appropriate concentration of 10 μ M was selected to study the inhibitory effect of the compounds. A set of ROH dose-response curves in the presence of inhibitory compounds is shown in Figure 5. The compounds demonstrated a competitive antagonist effect, as previously shown for FEN,²² with a shift of the ROH response curve to the right. FEN produced a 30-fold increase in the EC_{50} (Table 2). Additionally, compounds 1 and 58 had a strong effect, with a 15-fold increase, while the rest of the compounds tested in the TR-FRET assay produced an only slight increase in the EC₅₀ (2-3-fold).

sRBP-TTR Interaction Assessed by Surface Plasmon Resonance. We sought to further expand our understanding of the best retinoid analogues by studying their ability to disrupt the interaction between sRBP and TTR in real time using a surface plasmon resonance (SPR) assay. As shown in Figure 6, the only compound that strengthens the interaction between sRBP and TTR is ROH, as expected. None of the analogues strengthened the interaction with TTR, but all of them, with the exception of compound 1, were less disruptive than FEN itself and hence exhibited qualitatively the same results that were seen in the TR-FRET analysis.

sRBP-Receptor Interaction Assessed by Surface Plasmon **Resonance.** The ability of the compounds to disrupt the interaction between sRBP and solubilized membranes was also examined using the SPR binding assay. Detergent-treated HEK293 membrane fractions were preincubated with compounds and then injected over sRBP immobilized on the sensor chip surface. All the novel retinoid compounds tested had a negative effect on the stability of the sRBP-membrane protein complex and consequently produced faster dissociation (Figure 7). However, FEN had little or no effect. All experiments were repeated at least three times with no significant differences. Note

Scheme 2

that background binding to other components of the solubilized membrane will be present.

DISCUSSION

For some time, there has been great concern about the link among obesity, insulin resistance, type 2 diabetes, and cardiovascular disease. A recent scientific discovery provided a way forward toward preventing the avalanche of disability due to type 2 diabetes. This was the observation that elevated sRBP levels may attenuate the response of the cell to insulin, thus increasing resistance and presaging type 2 diabetes.¹¹ A second important element was the much earlier demonstration and functional characterization of a plasma membrane receptor for sRBP^{24,25} through which the effects of sRBP might be mediated. The connection with insulin action is a novel one, suggesting cross talk between RBP and insulin, because higher levels of the former are seen to inhibit the intracellular response to insulin.¹³

There has been some controversy about whether sRBP levels are statistically correlated with the development particularly of type 2 diabetes, with reports both confirming $^{26-29}$ and failing to confirm³⁰⁻³² the link. The reasons for these differences are not clear, the only suggestion put forward so far being some variation in the methodologies used for sRBP estimation,²⁹ but very significantly, the first studies on isolated human adipocytes confirmed that an elevated sRBP level



Figure 3. Titration of apo-sRBP with a selection of compounds. For quenching of the protein fluorescence, the excitation and emission wavelengths were 280 and 345 nm, respectively. The titration system consisted of 0.5 mL of 2 μ M purified apo-sRBP in PBS (pH 7.4). Solutions of ROH, FEN, or compounds in ethanol were added to the cuvette until a final concentration of 7.5 μ M was reached. The curves were corrected for the inner filter effect of the compounds.







 a Data shown are average values \pm the standard deviation (SD) from three independent experiments.

does indeed attenuate insulin-induced IRS-1 and ERKI/2 phosphorylation.¹³ Moreover, as found in the whole-animal studies, the serine residue on IRS-1 specifically affected was that closely identified with insulin signaling. Considering all this, it seems possible that reducing the "activity" of elevated levels of sRBP may restore normal insulin sensitivity and counteract type 2 diabetes.

The hypothesis we are exploiting is that compounds that reduce sRBP levels or disrupt its protein interactions will prevent the genesis of insulin resistance and consequently that of type 2 diabetes and its cardiovascular complications. Administration of FEN has been observed to exert therapeutic effects in mouse models of obesity and diabetes.^{15,18} Long-term FEN treatment prevents high-fat diet-induced obesity, insulin resistance, and hepatic steatosis.¹⁸ Additionally, retinoic acid (RA) treatment was determined to reduce body weight and basal serum glucose, serum ROH, and sRBP levels coupled with improved insulin sensitivity and to decrease the ROH:sRBP ratio.³³ In this work, a series of retinoid analogues were synthesized and examined for their ability to disrupt the interaction of sRBP with TTR and, for the first time, with its receptor. As a first step, we assessed the binding of these compounds to sRBP by fluorescence titration, a well-established method.¹⁹ The fluorescence emission data were used to calculate K_d values according to the method of Cogan et al.¹⁹ The values obtained were consistent with previous work, 19,34 with a $K_{\rm d}$ for ROH binding to sRBP of 182 \pm 4 nM

and comparable values for analogues 1 and 23–68. This is in agreement with earlier findings, which indicated that modifications of the functional hydroxyl end group of ROH (see Figure 2 and schemes in Experimental Section) do not change substantially the affinity for apo-sRBP.^{17,19,35} Compounds 2–22 did not show any absorption in the Trp emission region, and their binding to sRBP could not be assessed by this method. It has been shown that, to interact specifically with sRBP, retinoids must possess an intact trimethylcyclohexenyl group that establishes close contacts with the side chains of the internal β -barrel cavity.³⁵ The lack of a cyclohexene ring or having a shorter structure may limit the binding affinity of compounds 2–22 for sRBP, indicating that the strength of the interaction lies in the polyene chain region.

In the available crystal structures, the ROH hydroxyl group is near the protein surface, in the region of the entrance loops surrounding the opening of the binding cavity. It participates in polar interactions and becomes fully buried in the holo-sRBP– TTR complex.⁴ However, the cyclohexene ring and the isoprene tail of FEN take the place of the corresponding portions of ROH, while the hydroxyphenyl amide group protrudes from the cavity toward the solvent, replacing the ROH hydroxyl group and a water molecule hydrogen bonded to it.¹⁶

Subsequent analysis focused on compounds 1 and 23-68 with an examination of their effect on the interaction between sRBP and TTR. For this purpose, we developed a pull-down



Figure 4. Pull-down assay for sRBP after incubation with compounds. (A) The ROH–sRBP–His-TTR complex was incubated in the presence of $10 \,\mu$ M ROH and $100 \,\mu$ M FEN. After incubation, the complexes and free sRBP were separated by a pull-down assay and the flow-through ractions: lane 1, control ($10 \,\mu$ M ROH, no compound); lane 2, $10 \,\mu$ M retinol and $100 \,\mu$ M FEN (positive control for complex disruption). (B) Dose–response pull-down assay for **58**. Lane 1 was the control ($10 \,\mu$ M ROH, no compound). Lanes 2–6 contained 10 μ M ROH and different concentrations of **58** (1, 10, 20, 50, and 100 μ M, respectively).



Figure 5. Time-resolved FRET for the sRBP–TTR interaction. Assays were conducted with a fixed concentration (10 μ M) of compound and increasing concentrations of ROH. sRBP and TTR concentrations were 5 and 100 nM, respectively. TR-FRET signals are the ratios between 665 and 615 nm emissions. The curves were generated using a sigmoidal dose–response equation in Prism (GraphPad Software Inc.). Results shown are from a representative experiment performed in duplicate \pm the standard deviation.

assay that could detect free sRBP released from the ROH-sRBP-TTR complex after its disruption in the presence of compounds.

Table 2. EC_{50} Values for ROH-sRBP-TTR Interaction in the Presence of Compounds^{*a*}

compound	EC ₅₀ (nM)
no compound	71 ± 11
Fenretinide	2372 ± 186
1	1260 ± 177
23	245 ± 35
41	171 ± 21
48	145 ± 5
58	954 ± 114
62	154 ± 17

 a Data shown are average values \pm the standard deviation from three independent experiments.



Figure 6. Sensorgram showing interactions of His-sRBP with TTR in the presence of selected compounds. SPR response curves reflect the interaction of TTR with immobilized sRBP. The responses were recorded as a function of time and are expressed in resonance units (RU).



Figure 7. Sensorgram showing interactions of His-sRBP with solubilized HEK293 cell membranes in the presence of selected compounds. SPR response curves reflect the interaction of HEK293 cell solubilized membrane preparations passed over immobilized sRBP. The responses were recorded as a function of time and are expressed in resonance units (RU).

The crystal structure of holo-sRBP complexed with TTR revealed that ROH participates in the interaction of sRBP with TTR,⁴ increasing the affinity of the sRBP–TTR interaction

6-fold, from 1.2 to 0.3 μ M.²¹ This relatively low affinity, even for the holoprotein, is physiologically significant: because holo-sRBP complexed with TTR lacks affinity for its membrane receptor,^{9,10} a small amount of uncomplexed holo-sRBP is needed to deliver ROH to the cells through this specific receptor. On the other hand, FEN, with a structure similar to that of ROH but containing a bulky phenylamide in place of the hydroxyl group, drastically interferes with the sRBP-TTR interaction because of steric hindrance in the contact regions between sRBP and TTR and also changes in the position of the loops surrounding the entrance of the binding site. In agreement with these observations, a small amount of free sRBP was always detected in the control without compounds after incubation of the holosRBP-TTR complex in the pull-down assay. In comparison, a larger amount of free sRBP (estimated to be 3-5-fold) could be measured after incubation of the complex with FEN. Compounds 2-22 did not show the ability to disrupt the sRBP-TTR interaction and will be disregarded in further studies. However, a number of compounds based on scaffold B [with variations in the outer end (1 and 23-68)] showed a similar disruptive effect on the complex with FEN and probably act through the same mechanism. Interestingly, some of the larger groups at the polyene end of the retinoid analogues extend farther than FEN but do not disrupt the sRBP-TTR interaction, indicating that the retinoid analogues may not function solely as steric blockers but perhaps induce conformational changes in the loop regions. This shall be further examined in the crystal structure work.

We cannot calculate accurate K_i values because of the nature of the pull-down assay. In all runs (tested at least three times) of this assay, the band intensities were very similar to those for FEN, indicating a disruptive effect with a potency similar to that of FEN. From this qualitative analysis, we could draw no definitive conclusions regarding the strength of the disruption of the best analogues, and hence, additional studies were pursued to validate our initial findings. We used a TR-FRET assay, a technique that has been recently applied to the study of sRBP-TTR interactions, to characterize their inhibitory behavior.²² The selected compounds produced a decrease in the strength of the basal interaction between sRBP and TTR in the absence of ROH and a right shift of the ROH curve compared to the control (no compound). This is in agreement with the competitive behavior described for FEN and retinyl acetate.²² The TR-FRET results were also validated using a secondary binding assay, SPR, which yielded qualitatively similar results, with FEN and compound 1 exhibiting a more disruptive sRBP-TTR interaction effect, while the other compounds were clustered together in both assavs.

With the SPR assay using the HEK293 cell membranes and immobilized His-sRBP, ROH had a positive effect on the interaction observed in real time and the compounds all had a disruptive effect, with FEN exhibiting the smallest disruption of the set. We have given values for the disruption of the RBP-TTR interaction from FRET assays, but it is not possible to generate other than relative potencies for the sRBP-receptor interactions because of the nature of SPR with solubilized whole membrane preparations. The usual problem is that accurate values can be obtained only when pure preparations of a receptor are used.

Using our three-tiered assay approach, including analyzing binding to sRBP and SPR studies of the sRBP–TTR and sRBP–sRBP receptor interaction, we were able to show that

most sRBP binders exhibited a dual effect. Some compounds inhibited TTR more than the receptor interaction and a few the reverse. It is known that FEN reduces sRBP levels and improves insulin sensitivity,^{11,16,36} probably by disrupting the ROH–sRBP– TTR complex promoting clearance of sRBP.^{11,17} Our SPR data suggest that FEN might have a dual effect on the RBP insulin resistance link: (i) lowering sRBP levels and (ii) interacting directly with the insulin pathway inside the cell, its uptake having been mediated by the sRBP receptor. This is supported by the observation that FEN only "mildly" interferes with the interaction of the sRBP–ROH complex with the membrane compared to the strong disruption of that interaction caused by other compounds.

CONCLUSIONS

In this work, a number of active retinoid-like molecules that inhibit sRBP-TTR and sRBP-receptor interactions as well as or better than FEN, the antitumor positive control, have been identified, thereby presenting a potential novel dual mechanism of action. Thus, the compounds discovered may offer a new therapeutic intervention against type 2 diabetes and its development.

EXPERIMENTAL SECTION

Biology. *Materials. Escherichia coli* strain BL21(DE3), *P. pastoris* strain KM71H (*aox1::ARG4, arg4*), expression vector pPICZα A, the Zero Blunt TOPO PCR cloning kit, and europium-labeled anti-His tag antibodies were purchased from Invitrogen. HEK293 cells (human embryonic kidney) were from the ATCC. SureLight streptavidin-conjugated allophycocyanin (APC) was from PerkinElmer. The EZ-Link Sulfo-NHS-LC-Biotinylation kit was from Pierce. ROH, FEN, native human TTR, and synthetic oligonucleotide primers were from Sigma. Polymerase chain reaction (PCR) amplification of coding sequences was performed with PfuTurbo Hotstart DNA polymerase from Stratagene. Ni-NTA superflow resin and cobalt affinity gel were purchased from Qiagen and Sigma, respectively. Plasmid isolation and gel extraction kits were from Qiagen. Biacore materials were obtained from GE Healthcare. Retinoid analogues were synthesized by Sygnature Chemical Services. Other reagents used were molecular biology grade.

Expression and Purification of Recombinant Human sRBP in E. coli. The mature form of human sRBP was expressed without tags using vector pMMHa. NdeI and BamHI restriction sites flanking the coding sequence of mature human sRBP (corresponding to residues 19–201) were constructed by PCR using the sRBP cDNA originally cloned into the pQE-30 vector.⁹ The oligonucleotide primers designated to amplify the insert were NdeI (forward) 5'-TACATATGGAGCGCGACTGCC-GAGTG-3' and BamHI (reverse) 5'-TAGGATCCCTACAAAAGGT-TTCTTTCTGATCTGC-3'. The PCR product was inserted into the pCR-Blunt II TOPO vector, digested with NdeI and BamHI, and introduced into the same restriction sites of the pMMHa vector under the control of the T7 promoter.

BL21(DE3) cells transformed with expression plasmid pMMHasRBP were grown in 2×YT medium (1.6% tryptone, 1% yeast extract, and 0.5% NaCl) with 100 μ g/mL ampicillin. When the cell culture reached an optical density of 0.6 at 600 nm, protein expression was induced with 1 mM IPTG. After overnight incubation at 37 °C, the cells were harvested by centrifugation at 7000g for 10 min at 4 °C and subjected to disruption by sonication. The resulting suspension was centrifuged at 10000g for 10 min at 4 °C and the pellet (sRBP inclusion bodies) suspended at a concentration of 1 mg/mL sRBP in 5 M guanidinium chloride with 10 mM DTT at pH 9.0 to help in the reduction and solubilization of recombinant protein. After overnight incubation at 25 °C, the refolding of sRBP was started by dilution of the denatured protein to a concentration of 0.25 mg/mL in cold refolding buffer [25 mM Tris-HCl (pH 9.0), 0.3 mM cystine, 3 mM cysteine, and 1 mM EDTA] with a 10-fold molar excess of ROH.³⁷ The guanidinium chloride concentration at this step was 1 M. After refolding, the protein solution was dialyzed against PBS buffer, and then refolded sRBP was purified by affinity chromatography with His-TTR (see below).

Expression and Purification of Recombinant Human sRBP in P. pastoris. The sRBP was expressed with an N-terminal His tag using the *P. pastoris* expression system as described previously.²³ The sRBP coding sequence was amplified by PCR using the oligonucleotide primers EcoRI (forward) *S'*-GAATTCCATCATCATCATCATCATCATGAGCGCGACT-GCCGAGTG-3' and XbaI (reverse) *S'*-TCTAGACTACAAAAGGTTT-CTTTCTGATCTGCC-3'. The forward primer included the sequence for a six-His tag. The PCR product was ligated in the pCR-Blunt II TOPO vector and subsequently introduced into the EcoRI and XbaI sites of the pPICZα A vector under the control of the AOX1 promoter.

P. pastoris cells were transformed with expression vector pPICZα A-His-sRBP and grown at 30 °C in phosphate-buffered YP medium [1% yeast extract and 2% peptone (pH 7.5)] in the presence of 2% glycerol (w/v) until the OD₆₀₀ reached 6.0. The cells were harvested by centrifugation at 7000g for 10 min and used to inoculate YP medium to an OD₆₀₀ of 1.0 supplemented with 1% methanol (v/v) to induce expression. For each of 2 days, methanol was added to a final concentration of 1% (v/v). After induction for 48 h, the culture supernatant containing the His-sRBP was incubated with Ni-NTA resin and the bound proteins eluted with elution buffer [20 mM Na₂HPO₄, 500 mM NaCl, and 250 mM imidazole (pH 7.4)]. The purified His-sRBP was dialyzed against PBS buffer and then concentrated by ultrafiltration.

Expression and Purification of Recombinant Human His-TTR. The coding sequence of human TTR in vector pMMHa was a gift from J. W. Kelly.³⁸ To facilitate the subsequent purification of the holo-sRBP–TTR complex, we expressed TTR with an N-terminal His tag. The coding sequence of mature TTR (corresponding to residues 21–147) was inserted into the BamHI and HindIII restriction sites in the pQE-30 vector. The oligonucleotide primers used in the PCR were BamHI (forward) 5'-GGATTCGGCCCTACGGGCACCG-3' and HindIII (reverse) 5'-AAGCTTTCATTCCTTGGGATTGGTGACG-3'. The PCR product was ligated in the pCR-Blunt II TOPO vector, digested with BamHI and HindIII, and inserted into the pQE-30 vector.

The expression of this protein in *E. coli* was performed as for sRBP. After overnight induction with IPTG, the cells were collected and sonicated. The supernatant containing the soluble TTR was incubated with Ni-NTA resin and the recombinant protein eluted with elution buffer. The purified His-TTR was dialyzed against PBS buffer.

Purification of the ROH–sRBP–TTR Complex. Purified His-TTR was added to the refolded, untagged holo-sRBP solution and the mixture incubated overnight at 4 °C in the presence of 100 μ M ROH. Then, the protein solution was applied to a cobalt affinity column equilibrated with buffer I [20 mM Na₂HPO₄, 500 mM NaCl, and 10 mM imidazole (pH 7.4)]. The column was washed with the same buffer and then eluted with elution buffer. Fractions were analyzed by SDS–PAGE and Coomassie staining.

Fluorimetric Binding Assays. The binding of compounds to HissRBP was monitored by titration of the intrinsic fluorescence emission of the protein in the presence of increasing concentrations of compounds. The quenching of protein fluorescence due to the transfer of energy to the ligand was evaluated using excitation and emission wavelengths of 280 and 350 nm, respectively, in a Cary Eclipse fluorescence spectrophotometer (Varian). Purified His-sRBP expressed with the *P. pastoris* system was diluted to a concentration of 2 μ M in PBS buffer, and small increments of compound solutions in ethanol were added. The system was mixed and allowed to equilibrate for 5 min before the fluorescence emission of the compound—His-sRBP complex was recorded. The final concentration of ethanol never exceeded 2%. A solution of *N*-acetyl-L-tryptophanamide was used as a blank. The apparent dissociation constants (K_d) of compounds from His-sRBP were estimated as described by Cogan et al.¹⁹

sRBP—TTR Interaction Assessed by Pull-Down Assay. The capacity of the compounds that bind to sRBP to disrupt the interaction between this protein and its partner TTR was examined first by a pull-down assay. In this assay, the purified holo-sRBP—TTR complex [0.2 mg/mL in PBS buffer (pH 7.4)] was incubated in the presence of 10 μ M retinol and each compound at 100 μ M, for 1 h at 37 °C. Then, 20 μ L of cobalt resin was added to separate by pull-down assay the fraction of free sRBP from the fraction bound to TTR. The flow-throughs, containing the free sRBP, were examined by SDS—PAGE and silver staining. A control with FEN, which is known to disrupt the interaction between sRBP and TTR, was included in the assay for comparison.

Time-Resolved FRET. The TR-FRET assay was based on the previously published protocol of Coward et al.²² The reaction mixtures were prepared in white 96-well plates (Greiner) in a final volume of 100 μ L. Native TTR was biotinylated according to the manufacturer's manual. Each well contained 5 nM apo-His-sRBP, 100 nM biotinylated TTR, 100 nM SureLight streptavidin-conjugated APC, 1 nM europium-labeled anti-His tag antibody, 10 μ M inhibitor, and increasing concentrations of ROH in TR-FRET buffer [20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 0.01% BSA, and 0.02% Tween 20]. Reaction mixtures were incubated for 2 h at room temperature and measured on a POLARstar Omega microplate reader (excitation wavelength of 337 nm, emission wavelengths of 615 and 665 nm).

Surface Plasmon Resonance (SPR). Interaction analyses were performed in SPR running buffer (HBS with 50 μ M EDTA) at a flow rate of 10 μ L/min, at a constant temperature (25 °C), using a Biacore 3000 system (GE Healthcare). Ni-NTA sensor chips were used for all experiments, and general procedures were performed according to the manufacturer's instructions for this sensor chip. Briefly, each cycle consisted of surface activation with 20 µL of nickel (2 min), followed by a 5 min injection of recombinant sRBP (His-tagged, 200 nM solution, prepared in SPR running buffer), followed by a 1 min pulse of a specific concentration of compound (diluted in SPR running buffer), culminating with (i) a 5 min injection of untagged, native TTR (1 μ M in running buffer incubated with each compound at $10 \,\mu\text{M}$) or (ii) a 5 min injection of solubilized HEK293 cells membranes⁹ (50 μ g/mL in running buffer incubated with each compound at 10 μ M). Finally, sensor chip regeneration was achieved by stripping all proteins from the surface, performing a 2 min injection of 0.3 mM EDTA in HBS, at the end of each cycle (maximal number of regenerations per chip, 100). Sensor chips were used for 2 weeks, or until the level of nonspecific binding increased (\geq 5%). All experiments were performed in parallel with an inactivated flow cell not coated with protein. Each independent experiment was repeated three to five times.

Data Processing. Prism 5.00 (GraphPad Software Inc.) was used to determine EC_{50} values. All SPR sensorgrams were corrected for bufferinduced refractive index changes at an uncoated reference surface, analyzed using BIAEVALUATION version 3.2 (GE Healthcare).

Chemistry. For the determination of compound purity, the system consisted of a Waters HPLC and mass spectrometer system and an Agilent Scalar 5 μ m C18 50 mm × 4.6 mm column. Detection was achieved using an electrospray ionization source (positive or negative ion) and a UV detector at 254 and 215 nm. Mobile phase A was 0.1% aqueous formic acid, and mobile phase B was 0.1% formic acid in MeCN. The flow rate was 2.5 mL/min. The following gradient was used: 5% B from 0 to 0.1 min, 5 to 95% B from 0.1 to 5 min, 95% B from 5 to 5.5 min, 95% B from 5.5 to 5.6 min, flow rate increased to 3.5 mL/min, 95% B from 5.6 to 6.6 min, 95 to 5% B from 6.6 to 6.75 min, 5% B from 6.75 to 6.9 min, 5% B from 6.9 to 7 min, flow rate reduced to 2.5 mL/min. The majority of the library supplied was >95% pure.

Preparation. Retinoic acid (1 g, 3.33 mmol) and EDCI (0.957 g, 4.99 mmol) were stirred in DCM (20 mL), and methyl 2-(4-aminophenoxy)-acetate (0.724 g, 3.99 mmol) was added (Scheme 1). After 16 h, the reaction mixture was applied to a silica cartridge (80 g) and eluted with an EtOAc/isohexane mixture (1:3, v/v) to afford the target compound as a yellow oil that solidifies on standing.

1, methyl 2-{4-[(2*E*,4*E*,6*E*,8*E*)-3,7-dimethyl-9-(2,6,6-trimethylcyclohex-1-en-1-yl)nona-2,4,6,8-tetraenamido]phenoxy}acetate: $t_{\rm R} = 3.02$ min; m/z 464 (M + H)⁺ (ES⁺); yield 0.5226 g, 34%; 100% pure.



Preparation. 4-Hydroxyaniline (43.6 mg, 0.399 mmol) was added to a solution of *cis*-retinoic acid (100 mg, 0.333 mmol) and EDCI (96 mg, 0.499 mmol) in DCM (1 mL) (Scheme 2), and the reaction mixture was stirred at room temperature overnight.

The reaction mixture was diluted with water and passed through a phase separator. The organic layer was applied to a 4 g silica cartridge and eluted with 0 to 100% EtOAc in cyclohexane. This gave a yellow solid.

23, (2*Z*,4*E*,6*E*,8*E*)-*N*-(4-hydroxyphenyl)-3,7-dimethyl-9-(2,6,6-trimethylcyclohex-1-en-1-yl)nona-2,4,6,8-tetraenamide: $t_{\rm R}$ = 2.82 min; *m*/*z* 392 (M + H)⁺ (ES⁺); yield 13.6 mg, 10%; 94% pure.



Preparation. Amine (0.4 mmol, 1.2 equiv) was added to retinoic acid (100 mg, 0.333 mmol) and EDCI (96 mg, 0.499 mmol) in DCM (1 mL) (Scheme 3). The reaction mixtures were then stirred at room temperature for 24 h. All reactions were investigated by LC–MS. Those showing the desired product were quenched with water, and the mixture was passed through a phase separator. Products were purified by preparative HPLC.

41, (2*E*,4*E*,6*E*,8*E*)-*N*-(2,3-dihydroxypropyl)-3,7-dimethyl-9-(2,6,6-trimethylcyclohex-1-en-1-yl)nona-2,4,6,8-tetraenamide: $t_{\rm R}$ = 4.72 min; *m*/*z* 374 (M + H)⁺ (ES⁺); yield 7.8 mg, 6%; 92% pure.



48, (2*E*,4*E*,6*E*,8*E*)-*N*-(2-hydroxyethyl)-3,7-dimethyl-9-(2,6,6-trimethyl-cyclohex-1-en-1-yl)nona-2,4,6,8-tetraenamide: $t_{\rm R}$ = 2.60 min; *m*/*z* 344 (M + H)⁺ (ES⁺); yield 12.2 mg, 11%; 90% pure.



Amine (0.4 mmol, 1.2 equiv) was added to Retinoic acid (100 mg, 0.333 mmol), and EDCI (96 mg, 0.499 mmol) in DCM (1 mL). The reaction mixtures were then stirred at room temperature for 24 h. All reactions were investigated by LC–MS. Those showing the desired product were quenched with water, and the mixture was passed through a phase separator.

All products were purified on 4 g silica cartridges eluting with a 0 to 100% gradient (5% MeOH in EtOAc) in isohexane.

58, (2*E*,4*E*,6*E*,8*E*)-*N*-[4-(dimethylamino)phenyl]-3,7-dimethyl-9-(2,6,6-trimethylcyclohex-1-en-1-yl)nona-2,4,6,8-tetraenamide: $t_{\rm R}$ = 3.22 min; *m*/*z* 419 (M + H)⁺ (ES⁺); yield 15.3 mg, 11%; 91% pure.



62, (2*E*,4*E*,6*E*,8*E*)-3,7-dimethyl-1-morpholino-9-(2,6,6-trimethylcy-clohex-1-en-1-yl)nona-2,4,6,8-tetraen-1-one: $t_{\rm R}$ = 3.09 min; m/z 370 (M + H)⁺ (ES⁺); yield 10.8 mg, 9%; 98% pure.



ASSOCIATED CONTENT

Supporting Information. Chemical structures of compounds 1–68. This material is available free of charge via the Internet at http://pubs.acs.org.

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ABBREVIATIONS USED

APC, allophycocyanin; FEN, Fenretinide; ROH, retinol; RU, resonance unit; SDS—PAGE, sodium dodecyl sulfate—polyacrylamide gel electrophoresis; SPR, surface plasmon resonance; sRBP, serum retinol-binding protein; TR-FRET, time-resolved fluorescence resonance energy transfer; TTR, transthyretin

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