Anti-obesity effect of a small molecule repressor of RORγ

Mi Ra Chang, Yuanjun He, Tanya Khan, Dana S. Kuruvilla, Ruben Garcia-Ordonez, Cesar Corzo, Thaddeus J Unger, David W. White, Susan Khan, Li Lin, Michael D. Cameron, Theodore M. Kamenecka, and Patrick R. Griffin

Department of Molecular Therapeutics (M.R.C., Y.H., R.G-O, D.K., C.C., T.K., S.K., L.L, M.D.C., T.M.K., P.R.G)

The Scripps Research Institute, Jupiter, FL- 33458.USA

Ember Therapeutics (T.J.U, D.W.W.)

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Corresponding Author: Patrick R. Griffin, The Scripps Research Institute, Scripps, Florida, 130 Scripps Way, Jupiter, FL 33458, pgriffin@scripps.edu

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Abbreviations: Diet induced obesity mice (DIO), RORγ (Retinoic acid receptor-related orphan receptor γ), Hormone sensing lipase (HSL), Uncoupling protein 1 (UCP1), Comprehensive lab animal monitoring system (CLAMS), Inguinal white adipose tissue (iWAT)
ABSTRACT

The orphan nuclear receptor RORγ is a key regulator for TH17 cell differentiation and it regulates metabolic and circadian rhythm genes in peripheral tissues. Previously it was shown that the small molecule inverse agonist of RORγ SR1555 suppressed TH17 differentiation and stimulated iTreg cells. Here we show that treatment of cultured pre-adipocytes with SR1555 represses the expression of RORγ while leading to increased expression of FGF21 and adiposQ. Chronic administration of SR1555 to obese diabetic mice resulted in a modest reduction in food intake accompanied with significant reduction in fat mass resulting in reduced body weight and improved insulin sensitivity. Analysis ex vivo of treated mice demonstrates that SR1555 induced expression of the thermogenic gene program in fat depots. Further studies in cultured cells showed that SR1555 inhibited activation of hormone sensitive lipase and increased fatty acid oxidation. Combined, these results suggest that pharmacological repression of RORγ may represent a strategy for treatment of obesity by increasing thermogenesis and fatty acid oxidation, while inhibition of hormone sensitive lipase activity results in a reduction of serum free fatty acids leading to improved peripheral insulin sensitivity.
INTRODUCTION

The percentage of the global population categorized as overweight or obese has increased dramatically over the last few decades. This trend is predicted to continue as developed and developing nations increasingly adopt more sedentary lifestyles and gain easier access to high calorie diets. The metabolic syndrome is associated with obesity and patients with this syndrome are at a significant increased risk of suffering from cardiovascular disease and stroke (Moller & Kaufman, 2005). There are two major underlying drivers for the development of metabolic syndrome: excess adiposity (obesity) and Type 2 diabetes mellitus (T2DM) (Grundy et al, 2005). T2DM is a chronic metabolic disorder that results in part by the inability of the body to respond adequately to circulating insulin, a state of insulin resistance. In the obese state, free fatty acids (FFAs) are elevated in plasma and in all insulin responsive organs including skeletal muscle, liver and endothelial cells. As such, elevated FFAs are linked with the development of the metabolic syndrome. Obesity is also closely associated with a low grade state of inflammation characterized by elevated pro-inflammatory cytokines in blood and tissues (Tataranni & Ortega, 2005).

Treatments for metabolic syndrome included modification of diet and increased exercise (Grundy et al, 2004). However, pharmacologic intervention is typically required as weight loss and exercise often are not sufficient due to poor compliance and confounding genetic factors (Bouchard, 1988; Moller et al, 1996). Members of the nuclear receptor (NR) superfamily are ligand controlled
transcription factors that regulate a wide range of metabolic, endocrine and immunologic functions and this protein superfamily has proven to be a rich source of targets for the development of therapeutics for a wide range of human diseases including inflammation and diabetes. A subset of NRs are classified as orphan receptors due to lack of a characterized or agreed upon endogenous ligand (Kliewer et al, 1999). The retinoic acid receptor-related orphan receptor (ROR; NR1F) subfamily was identified based on sequence similarities to the retinoic acid and retinoid X receptors (Becker-Andre et al, 1993; Giguere et al, 1994). The T cell specific gamma isoform, RORγt, has been the focus of considerable attention due to its role in the development of T helper 17 cells (TH17) and in the pathology of autoimmune disease. While RORγt is highly expressed in immune cells and thymus, RORγ variants are expressed in the liver, skeletal muscle, adipose tissue, and kidney (Jetten, 2009), and its expression can be induced in macrophages during acute inflammatory responses (Barish et al, 2005; Chang et al, 2014; Gu et al, 2008). Furthermore, genetic deletion of RORγ results in profound effects on adipose depots with increased adipocyte number, yet reduced hypertrophy, accompanied by improved insulin sensitivity. These RORγ-deficient mice were shown to be resistant to diet-induced insulin resistance (Meissburger et al, 2011).

Recently, synthetic modulators of the ROR subfamily have been described (Kojetin & Burris, 2014; Marciano et al, 2014) including a series of reports on the development of potent and selective inverse agonists of RORγ. These studies demonstrate their utility in reducing the severity of inflammation in mouse models.
of multiple sclerosis (experimental autoimmune encephalomyelitis, EAE) (Solt et al, 2011), and rheumatoid arthritis (collagen-induced arthritis, CIA) (Cascao et al, 2012; Chang et al, 2014; Huh et al, 2011). One such compound, SR1555, was previously shown to inhibit TH17 cell development while increasing the frequency of T regulatory (Treg) cells (Solt et al, 2012). Thus, SR1555 was evaluated in the CIA mouse model (Chang et al, 2014). In addition to observing a reduction in joint inflammation, effects on body weight and adiposity were noted. Based on this observation SR1555 was evaluated in the diet-induced obesity (DIO) murine model of obesity and T2DM and the results of these studies are presented in this report.
MATERIALS AND METHODS

Chemicals. SR1555 (Fig. 1A) was synthesized as previously described (Solt et al, 2012).

Lanthascreen TR-FRET Competitive Binding assay for PPARγ: The PPARγ competitive binding assay (Invitrogen) was performed according to the manufacturer’s protocol. A mixture of 0.5 nM GST–PPARγ LBD, 5 nM Tb-GST-antibody, 5 nM fluormone Pan-PPAR Green, and serial dilutions of compound beginning at 10 µM downwards was added in 384-well low-volume plates (Greiner) to a total volume of 18 µl (2% DMSO in all wells). DMSO at 2% concentration was used as a no-ligand control. All dilutions were made in TR-FRET PPAR assay buffer. Experiments were performed in triplicate and incubated for 2 h in the dark before analysis in Perkin Elmer ViewLux ultra HTS microplate reader. TR-FRET signal was measured by excitation at 340 nm with emission at 520 nm for fluorescein and 490 nm for terbium. Data were plotted as the TR-FRET ratio 520/490 nm using GraphPad Prism software.

Animals: Seventeen-week-old male C57BL/6 DIO mice were purchased from Jackson Laboratories. All procedures were approved and conducted in accordance to the Scripps Florida Institutional Animal Care and Use Committee. Mice were placed on a high-fat diet (HFD; 60% kcal derived from fat) for seven weeks until they reached an average weight of ~40 g. Prior to and during compound administration mice were maintained on HFD. Animals were sham dosed with vehicle for three days prior to compound administration. SR1555 was administered intraperitoneally (i.p.) twice a day (BID) at 10 or 20 mg/kg or per oral
(p.o.) at 20 mg/kg once-a-day (QD). Body weight and food intake were monitored daily. Pre- and post-experiment whole body composition analysis was performed on vehicle only and SR1555 treated mice using nuclear magnetic resonance (NMR, Minispec mq 7.5 NMR analyzer, Bruker Optics). Blood samples were collected for plasma cholesterol, triglyceride, free fatty acid and insulin measurements. For insulin tolerance tests, mice were injected (i.p.) with 10 mg/kg SR1555 for 18 days, and fasted 6 hrs before i.p. injection of Insulin (0.75 U/kg, Sigma aldrich). Blood glucose levels were measured at 0, 15, 30, 60, 90, 120 min using a glucometer (LifeScan). mice were allowed to recover from the stress associated with ITT for 7 days prior to performing the a glucose tolerance test (GTT). Here mice were fasted overnight before injection of 2 g/kg D-glucose i.p. and blood glucose levels were measured at 0, 15, 30, 60, 90, 120 min using a glucometer (LifeScan).

**In vitro adipocyte differentiation:** 3T3-L1 cells and C3H10T1/2 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS (Gibco BRL). Twenty-four hours post-confluence, the 3T3-L1 cells were differentiated with IBMX (0.5mM), dexamethasone (1μM), insulin (10 μg/ml) and 10% FBS for forty-eight hours. After induction of adipogenesis, the cells were maintained with insulin (10 μg/ml) and 20μM of SR1555 for three or four days. The stromal vascular fraction (SVF) within adipose tissue was isolated from the epididymal adipose tissue of C57BL/6 mice to avoid contamination from non-adipose tissues as previously described (Koh et al, 2007). To exclude blood contamination in the adipose tissue, systemic perfusion with heparinized PBS
was performed before harvesting or washing isolated adipose tissues with PBS. Then, the adipose tissues were incubated in Hanks balanced salt solution (HBSS; Sigma-Aldrich) containing 0.2% collagenase type 2 for 60 minutes at 37°C with constant shaking. After inactivation collagenase activity with 10% fetal bovine serum (FBS) containing Dulbecco modified eagle medium (DMEM), the cell suspension was filtered through a 40μm nylon mesh (BD Biosciences) followed by centrifugation at 420g for 5 minutes. Floating adipocytes and supernatant were removed from the SVF pellet. The SVF pellet was washed and re-suspended in sterilized PBS.

**Quantitative real-time PCR:** Total RNA was extracted from 3T3-L1 cells or tissues using TRIzol reagent (Invitrogen). The RNA was reverse-transcribed using the ABI reverse transcription kit. Quantitative PCR reactions were performed with SYBR green fluorescent dye using an ABI9300 PCR machine. Relative mRNA expression was determined by the ΔΔ-Ct method normalized to GAPDH levels. The sequences of primers used in this study are found in **Supplemental Fig. 1.**

**Mitotracker-FACS quantification:** C2C12 cells were washed with PBS, trypsinized and incubated at 37°C for 20 min with 100 nM MitoTracker Green FM and Red GM dyes (Molecular Probes). Mitotracker Green probe preferentially accumulates in mitochondria, allowing estimation of mitochondrial quantity. Mitotracker Red probe is a red-fluorescent dye that stains mitochondria in living cells and its accumulation is dependent on the membrane potential (Molecular Probes). Samples were washed 3 times in PBS and subjected to flow cytometric
analysis on a BD LSRII instrument (Becton Dickinson, San Jose, CA), and the results were analyzed using FlowJo software (Tree Star).

Oxygen Consumption: Immortalized SVF cells were seeded at $1 \times 10^4$ cells per well in a differentiation medium containing DMEM with 10% FBS on XF-96 well culture plates (Seahorse). Cellular oxygen consumption rate was measured using the Seahorse XF96 at four to five days differentiation with 20μM SR1555. The concentration of fatty acid (oleic acid, Linoleic acid), oligomycin, FCC, rotenone were 1mg/mL, 10mg/mL, 6mM, 3mM, respectively.

Resting whole-body metabolic parameters: Daily average whole–body $VO_2$ (ml/kg/h), and cage activity (movement counts) were measured in DIO mice using a comprehensive laboratory animal monitoring system (CLAMS: Oxymax series; Columbus Instruments, Columbus, OH, USA). Each sealed chamber (CLAMS unit) is equipped with an O2 electrochemical sensor, a CO2 infrared sensor, and infrared beam activity sensors. The units allow the volume of O2 consumed ($VO_2$; ml/kg/h) to be measured. Infrared beam interruptions in both horizontal (X) and vertical (Z) directions were measured to quantify the motor activity of mice in both directions. Total horizontal beam breaks are summed to provide total horizontal activity count or XTOT. Two or more consecutive horizontal beam breaks were recorded as ambulatory activity count (XAMB). All vertical beam breaks were summed to provide total vertical activity count or ZTOT. CLAMS studies began after animals were acclimated to the metabolic chambers, and $VO_2$ data were collected 16-min intervals over a 5 day period.
under a consistent environmental temperature (22°C). During these studies, mice were allowed free access to food and water.

**Western blot analysis:** Primary antibodies against phosphor-HSL (S660), total HSL and beta-actin were purchased from Cell Signaling Technology (CST). A rabbit fast western blotting kit (Pierce, Thermo scientific) was used for detection of primary antibody signal.

**Plasma concentration of FGF21 and Insulin:** Blood samples were collected in heparin tubes on ice and centrifuged at 4°C. Endogenous FGF21 and Insulin levels were determined by a specific mouse enzyme-linked immunosorbent assay (ELISA) (Millipore, Billerica, MA) following manufacturer’s instructions.

**Plasma Lipid Profile:** Blood samples were collected in heparin tubes on ice and centrifuged at 4°C. Cholesterol, triglyceride (TAG) and free fatty acid (FFA) were quantitated using a Cobas c311 clinical chemistry analyzer (Roche Diagnostics).

**Pharmacokinetics:** The oral pharmacokinetic profile of SR1555 was determined in male C57Bl6 mice. SR1555 was formulated at a concentration of 1 mg/ml in 15% cremophore EL in water and dosed by oral gavage to three mice at a final dose of 20 mg/kg. Blood was collected in lithium-heparin coated capillary tubes using a microsampling technique at 15, 30, 60, 120, 240, 360, 480, and 1440 minutes and plasma was generated using standard centrifugation methods. Plasma concentrations were determined via LC-MS/MS on an ABSciex 5500. SR1555 was detected using the mass transition 461→333, and concentrations were determined by comparison to a nine point standard curve between 2 and
2000 ng/ml prepared in mouse plasma. Pharmacokinetic analysis was done with WinNonLin, Pharsight inc. using a noncompartmental model. All procedures described are covered under existing protocols and have been approved by the Scripps Florida IACUC to be conducted in the Scripps vivarium, which is fully AAALAC accredited.

**Statistical analysis:** The data are reported as the mean ± SEM. Statistical significance was evaluated using Student's unpaired t-test. P values less than 0.05 were considered significant.
RESULTS

SR1555 alters adipogenesis. Previously, SR1555 (Fig. 1A) was shown to be a RORγ-specific inverse agonist that inhibited T_H17 cell development and increased the Treg cell population (Solt et al, 2012). Since genetic deletion of RORγ results in profound effects on adipose tissue we investigated the effects of SR1555 on adipogenesis using differentiated 3T3L1 cells. Treatment of these cells with SR1555 resulted in decreased expression of RORγ (Fig. 1B). Consistent with observations from RORγ-depleted mice, SR1555 treatment led to increased expression of FABP4 (fatty acid binding protein; aP2) and Adiponectin (AdipoQ) but it had no effect on the expression of PPARγ, a key adipogenic regulator. Interestingly, the expression of FGF21 (fibroblast growth factor 21) was increased in SR1555 treated 3T3L1 adipocytes. FABP4 is a PPARG (NR1C3) target gene so we tested the ability of SR1555 to bind to this nuclear receptor. Consistent with results from broad selectivity profiling and as shown in Supplemental Fig. 2A, in a TR-FRET competitive binding assay, SR1555 was not able to displace labeled ligand at all concentrations tested. Furthermore, SR1555 was not able to transactivate the expression of luciferase in a reporter gene assay at all concentrations tested (Supplemental Fig. 2B).

RORγ-deficient mice are resistant to obesity and insulin resistance (Meissburger et al, 2011). Based on these observations, we sought to determine the effects of pharmacological repression of RORγ in the diet induced obesity (DIO) mouse model of obesity and diabetes. Mice were fed a HFD until reaching obesity (avg. weight of ~40g). DIO mice were then treated with vehicle (15%
cremophor, i.p.) for three days prior to initiating administration of SR1555. As shown in Supplemental Fig. 3, bolus administration of SR1555 to mice at 5mg/kg, 10mg/kg i.p., and 20mg/kg by oral gavage afforded plasma concentration of compound four hours post administration in excess of 4, 8, and 16μM, respectively. The calculated PK parameters are presented in Supplemental Fig. 4. With the intent of only partially repressing RORγ and based on the plasma exposure from the PK studies shown in Supplemental Fig. 3, SR1555 was administered i.p. to obese mice at 5mg/kg or 10mg/kg twice a day (BID). Body weight and food intake were recorded daily. Following 20 days of compound administration, DIO mice receiving vehicle only maintained body weight but animals treated with SR1555 at 5mg/kg had lost 15% of their body weight (Fig. 2A). SR1555 demonstrated a dose-dependent impact on body weight as mice administered compound at 10mg/kg lost an additional 8% body weight (23% loss). Compound treatment did cause a modest reduction in food intake by DIO mice as compared to control animals (Fig. 2B). However, no change in food intake was observed in lean mice suggesting SR1555 is not driving weight loss by systemic toxicity. It is possible that SR1555 treatment results in an improvement in leptin sensitivity.

Whole body composition of DIO mice was analyzed using NMR prior to the start of vehicle or compound administration and again at the conclusion of the study. As shown in Fig. 2C, lean mass was unaffected by SR1555 treatment; however, there was a statistically significant loss of fat mass in the 10mg/kg SR1555 treated DIO mice. On Day 18 of treatment, mice were fasted for 8 hours
and then subjected to an insulin tolerance test (ITT). As shown in Fig. 3A fasting insulin was significantly reduced in both the 5 and 10 mg/kg arms of the study. Fig. 3B illustrates that glucose disposal was improved with a quick decline in plasma glucose 15 min post insulin administration in the 10mg/kg arm of the study. Statistically significant differences in plasma glucose were observed at 30, 60 and 90 min post insulin administration. Following the ITT, compound administration was terminated and mice were allowed to recover from the stress of fasting. At seven days post compound administration a glucose tolerance test (GTT) was performed to determine if SR1555 treatment results in improved disposal of plasma glucose in response to a bolus intake of a glucose solution. As shown in Fig. 3C, DIO mice treated with SR1555 retained improved insulin sensitivity seven days post compound treatment. Analysis of plasma lipids at this time point showed that SR1555 treatment reduced cholesterol and triglyceride levels below vehicle only treated mice and to a level similar to that in control lean mice maintained on a normal chow diet (Fig. 3D). Free fatty acid levels were also reduced in SR1555 treated animals (Fig. 3D). Although SR1555 increased expression of FGF21 in adipocytes, compound treatment in DIO mice did not result in an increase in plasma FGF21 (data not shown) released from the liver.

We next investigated whether whole-body energy expenditure and ambulatory activity were impacted by SR1555 treatment. DIO mice were acclimated to single animal housing for seven days prior to the start of this study. SR1555 was then administered p.o. at 20mg/kg for 7 days before placing the animals in Comprehensive Lab Animal Monitoring System (CLAMS) chambers.
Once inside the CLAMS chambers, mice were administered SR1555 orally each day at 20 mg/kg and metabolic parameters and activity were evaluated over a seven-day period.

As shown in Fig. 4A, oxygen consumption (VO2) in compound treated DIO mice was increased during both the light and dark cycles as compared to the vehicle only treated group. Furthermore, body temperature was significantly increased in SR1555 treated mice (Supplemental Fig. 5A and B). No difference in XTOT and XAMB was observed when comparing SR1555 treated mice to vehicle only treated mice; however, ZTOT was increased in compound treated DIO mice (Fig. 4B, C, D). It is possible that this increase in vertical activity may reflect increased exploration of the environment (attention), cognitive enhancement, or perhaps anxiety. Regardless, the data obtained using CLAMS suggests that SR1555 stimulated activity and increased energy expenditure in DIO mice.

**Induction of thermogenesis by SR1555.** Uncoupling protein 1 (UCP1) is critical for heat generation and the protein is selectively expressed in brown and beige adipose cells. Classical brown adipose tissue (BAT) and inducible white adipocytes (beige cells) are capable of increasing energy expenditure through the uncoupling of oxidative metabolism from ATP (Wu et al, 2012; Wu et al, 2013). Also, previous physiological studies have shown that diet can induce thermogenesis in order to maintain body weight (Rothwell & Stock, 1997). Therefore, we asked whether SR1555 induces thermogenic genes in adipose depots in DIO mice. Inguinal white adipose tissue (iWAT) and brown adipose
tissue (BAT) were collected from DIO mice. Induction of UCP1 protein in iWAT and overall adipose size were smaller in SR1555 treated iWAT (Fig. 5A). Additionally, gene expression of UCP1, PRDM16 and FGF21 were increased in BAT from SR1555 treated mice as compared to tissue from vehicle only treated mice. The expression of 18S gene was used as a control to determine relative expression of thermogenic genes (Fig. 5B).

FABP4 (aP2) gene expression was also increased by SR1555 treatment and this phenotype was similar to that observed with genetic depletion of RORγ (Meissburger et al, 2011). Induction of UCP1 gene expression was also observed in the brown cell adipogenic process using C3H10T1-2 cells (Fig. 5C) where the effect of SR1555 treatment was similar to that of TZD treatment which was the positive control for the assay (Ohno et al, 2012). Furthermore, SR1555 increased the mitochondrial membrane potential and mass in C2C12 cells (Fig. 6A). Taken together, we conclude that SR1555 induces the thermogenic gene program and alters mitochondrial potential in adipose and muscle cell lines.

**Fatty Acid Oxidation and Lipolysis.** The reduction of FFAs observed in obese mice treated with SR1555 could be the result of increased FFA storage, increase degradation of FFAs by β-oxidation, decrease in lipolysis of triglycerides, or a combination of these events. The effect of SR1555 on fatty acid oxidation was monitored in immortalized stromal vascular fraction (SVF) cells. Differentiation is achieved *in vitro* by sequential application of adipogenic factors to pre-adipocytes over a period of several days. Following differentiation, adipocytes were characterized using the XF Analyzer from Seahorse Bioscience to measure four
essential parameters of mitochondrial function: basal respiration, ATP turnover, proton leak, and maximal respiration. Fatty acid oxidation in mitochondria is mainly controlled by CPT-1, which facilitates transport of long chain fatty acids into mitochondria. CPT-1 was unchanged in SR1555 treated SVF cells. However, oxygen consumption was increased at four independent time points following addition of fatty acid mixture (oleic acid and linoleic acid) whereas no increase was observed in DMSO only treated SVF cells (Fig. 6B). Next, we evaluated the activity of hormone sensitive lipase (HSL) in differentiated 3T3L1 cells following SR1555 treatment (Supplemental Fig. 6). Elevated plasma levels of free fatty acids (FFAs) are thought to play a major role in the pathogenesis of insulin resistance and type 2 diabetes by inhibiting glucose uptake and utilization by muscle and causing increased glucose output by the liver (Claus et al, 2005; Girousse et al, 2013). Here we observed that HSL gene expression and phosphorylated HSL (S660) were reduced in fully differentiated 3T3L1 cells. Therefore, we suggest that decreasing phosphorylated HSL might drive the reduction of FFA release from white adipocytes.
DISCUSSION

The nuclear receptor superfamily of ligand regulated transcription factors has proven to be a rich source of targets for the development of therapeutics for a wide range of human diseases. Within the NR1F subfamily of NRs, RORyt has garnered much attention due to its role in Th17 cells. However, its contribution to obesity and insulin resistance has been recently demonstrated (Meissburger et al, 2011; Takeda et al, 2014). The discovery that the potent LXR agonist T0901317 also functioned as an inverse agonist for ROR\(\alpha\) and ROR\(\gamma\) (Kumar et al, 2010) led to an explosion in medicinal chemistry efforts focused on the NR1F subfamily of receptors. This effort is exemplified by publications of the ROR\(\alpha/\gamma\) agonist SR1078 (Wang et al, 2010), the selective ROR\(\alpha\) inverse agonist SR3335 (Kumar et al, 2011), the dual ROR\(\alpha/\gamma\) inverse agonist SR1001 (Solt et al, 2011), and the selective ROR\(\gamma\) inverse agonist SR2211 (Kumar et al, 2012) and SR1555 (Solt et al, 2012). Here we demonstrate that the selective ROR\(\gamma\) inverse agonist SR1555 recapitulates many of RORG depletion phenotype in terms of adiposity and metabolic parameters. Specifically, SR1555 treatment of obese and diabetic mice modulated adipocyte function leading to body weight loss exclusively as a result of reduced adiposity leading to improved insulin sensitivity and glucose disposal.

It is possible that SR1555-mediated improvement in metabolic phenotype involves the modulation of Treg development. Previously it was shown that in cultured cells SR1555 repressed Th17 cell differentiation with concomitant
expansion of inducible Tregs (Solt et al, 2012), and induction of Tregs have been shown to be associated with metabolic disorders (Bhat et al, 2014; Michalek et al, 2011). Additionally, obese individuals with insulin resistance display a sharp reduction of Treg’s (DeFuria et al, 2013), and depletion of Treg’s in mice using an anti-CD25 monoclonal antibody exacerbates insulin resistance (Eller et al, 2011). Previously, we had shown that SR1555 was capable of suppressing inflammatory cytokine production such as IL-6 and TNF-α in LPS-stimulated RAW264.7 cells (Chang et al, 2014). Therefore, it is possible that the anti-inflammatory actions of SR1555 are associated with its ability to improve adiposity and metabolic parameters.

Treatment of 3T3L1 cells with SR1555 resulted in inhibition of the enzyme hormone-sensitive lipase (HSL; EC 3.1.1.79). HSL is highly expressed in adipose tissue and is a major contributor to the formation FFAs in fat. Release of these FFAs provides a source of energy for most tissues. HSL activity is the rate-limiting step in catecholamine-induced lipolysis and the level of HSL expression is directly related to the lipolytic capacity of mature fat cells (Large et al, 1998). These initial findings suggested that increased HSL activity may contribute to the increased plasma FFAs observed in obesity. Interestingly, partial inhibition of lipolysis in adipose tissue in HSL haplo-insufficient mice (HSL⁺⁻) resulted in improved insulin sensitivity (Girousse et al, 2013). Analysis of the promoter region of HSL (LIPE) suggests it contains four putative RORE sequences (Supplemental Fig 7. A and B). Thus, it is plausible that HSL expression and activity are regulated by RORγ. Previous CHiP-Seq data demonstrated that
RORγ binds to the promoter of family members of HSL. It remains to be determined if SR1555 mediated repression of HSL activity is direct or indirect.

Furthermore, SR1555 treatment increased gene expression and protein levels of UCP1, altered mitochondrial potential, and increased the level of fatty acid oxidation in both brown adipocytes and beige cells suggesting that RORγ regulates the oxidative phosphorylation process in mitochondria. In summary, the selective RORγ inverse agonist SR1555 may provide a useful starting point for the development of peripheral acting therapeutics for the treatment of diabetes and obesity.

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AUTHORSHIP CONTRIBUTIONS:

Participated in research design: Chang, Kamenecka, White, Griffin, Cameron

Conducted experiments: Chang, Garcia-Ordonez, Kuruvilla, T. Khan, S. Khan, Lin, Unger

Contributed new reagents or analytic tools: He, Kamenecka

Performed data analysis: Chang, Unger, White, Griffin, Cameron

Wrote or contributed to the writing of the manuscript: Chang, Corzo, Griffin, Cameron
References


Kojetin DJ, Burris TP (2014) REV-ERB and ROR nuclear receptors as drug targets. *Nat Rev Drug Discov* **13**: 197-216


Solt LA, Kumar N, He Y, Kamenecka TM, Griffin PR, Burris TP (2012) Identification of a selective RORgamma ligand that suppresses T(H)17 cells and stimulates T regulatory cells. ACS chemical biology 7: 1515-1519


Tataranni PA, Ortega E (2005) A burning question: does an adipokine-induced activation of the immune system mediate the effect of overnutrition on type 2 diabetes? Diabetes 54: 917-927


FOOTNOTES

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COI statement: None of the authors have conflicts of interest with the work presented in this manuscript. P.R.G. was a cofounder of Ember Therapeutics but they have recently shutdown. Two authors (T.J.U. and D.W.W.) were but are no longer employees of Ember Therapeutics.
Figure Legends

**Figure 1. SR1555 impacts adipogenic markers.** A) The chemical structure of SR1555 and B) gene expression analysis in differentiated adipocytes. 3T3-L1 cells were grown to confluence, incubated with dexamethasone, insulin and isobutylmethylxanthine (IBMX) for 2 days to induce differentiation. Cells were then treated with 20μM SR1555 and insulin for 5 days after which the expression of thermogenic genes and several nuclear receptors were analyzed by qPCR. GAPDH expression was used for normalization. Values are the mean ± SEM of 3 samples per group. * = P < 0.05, *** = P < 0.001 versus DMSO, as determined using Student’s unpaired t-test.

**Figure 2. SR1555 administration mice are resistant to obesity.** A) Body weight of HFD controls and SR1555-treated obese mice, B) Food intake, and C) whole body NMR. Twenty-one week old DIO mice were administered SR1555 i.p. at 5mg/kg or 10mg/kg for 20 days. Body weight and food intake were monitored daily. Percentage fat was measured just prior to the start of compound administration and again after 17 days of compound administration. Values are the mean ± SEM of 10 samples per group. * = P < 0.05, ** = P < 0.01, *** = P < 0.001 versus vehicle or before start dosing, by Student’s unpaired t-test.

**Figure 3. Metabolic parameters in DIO mice were improved by SR1555 administration.** A) cholesterol (CHOL), Triglyceride (TRIG) and free fatty acids (FFA) were measured in SR1555 treated DIO mice after 20 days dosing. B) Fasting Insulin and Insulin (0.75 U/kg) tolerance test (ITT, n=10) were measured
on DIO mice and C) glucose tolerance test GTT performed 7 days following last dose of SR1555. Values are the mean ± SEM of 10 samples per group. * = P < 0.05, ** = P < 0.01, *** = P < 0.001 versus vehicle, by Student's unpaired t-test.

Figure 4. *In vivo* measurement of energy expenditure and motor activity A) total oxygen consumption (VO2) B) total horizontal motor activity (XTOT), C) total vertical motor activity (ZTOT), and D) ambulatory activity count (XAMB). Mice were placed into the CLAMS for 5 days after one week of SR1555 treated obese mice. Values from all animals were pooled an average for an N=5 per group. Data were analyzed based on light and dark cycles. The light cycle occurred from 7:00am~7:00pm and dark cycle occurred from 7:00pm~7:00am in the metabolic cages. Values are the mean ± SEM of 10 samples per group. * = P < 0.05, ** = P < 0.01 versus vehicle, by Student's unpaired t-test.

Figure 5. Improvement of thermogenic genes by SR1555 and UCP1 induction. A) Immunohistochemical examination of the presence of UCP1. Inguinal adipose tissues from vehicle treated DIO mice or 10mg/kg SR1555 treated DIO mice were stained brown for UCP1 immunoreactivity. B) mRNA expression of thermogenic genes in brown fat tissue of mice fed with either Vehicle treated DIO or 10mg/kg SR1555 treated DIO. C) UCP1 mRNA expression in adipogenic C3H10T1/2 cell line. C3H10T1/2 cells were induced adipogenesis with Dex, Insulin, IBMX for 2 days then maintained with insulin and each compound for 5 days.
Figure 6. Fatty acid oxidation and modulation of mitochondrial respiration.
A) Functional mitochondria content in C2C12 cells was analyzed by flow cytometry using JC-1 and red/green JC-1 (the high-low mitochondrial membrane potential) MFI ratio (right, n= 3 per condition). Mitochondria content was measured by flow cytometry analysis using Mito Tracker (MT) Green staining (n=3 per condition). B) SVF cells were grown to confluence, incubated with dexamethasone, insulin, isobutylmethylxanthine, T3, and Rosiglitazone for 2 days to induce adipocyte differentiation, and then cultured with 20µM SR1555, insulin and T3. Oxygen consumption rates (OCR) were measured by Seahorse XF-96 flux analyzer (n= 10, 3 different condition). Values are the mean ± SEM of 10 samples per group. * = P < 0.05 versus DMSO, by Student’s unpaired t-test.
Figure 1

A

B
Figure 6

A

B

FA  oligomycin  FCCP  Rotenone

OCR (pMol/min)

0  5  11  17  24  30  36  42  48  54  60  66  72  78  84  90  96

time (min)

DMSO  SR1555

*  **  ***