PPARγ is a nexus controlling alternative activation of macrophages via glutamine metabolism

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The nuclear receptor peroxisome proliferator-activated receptor γ (PPARγ) is known to regulate lipid metabolism in many tissues, including macrophages. Here we report that peritoneal macrophage respiration is enhanced by rosiglitazone, an activating PPARγ ligand, in a PPARγ-dependent manner. Moreover, PPARγ is required for macrophage respiration even in the absence of exogenous ligand. Unexpectedly, the absence of PPARγ dramatically affects the oxidation of glutamine. Both glutamine and PPARγ have been implicated in alternative activation (AA) of macrophages, and PPARγ was required for interleukin 4 (IL4)-dependent gene expression and stimulation of macrophage respiration. Indeed, unstimulated macrophages lacking PPARγ contained elevated levels of the inflammation-associated metabolite itaconate and express a proinflammatory transcriptome that, remarkably, phenocopied that of macrophages depleted of glutamine. Thus, PPARγ functions as a checkpoint, guarding against inflammation, and is permissive for AA by facilitating glutamine metabolism. However, PPARγ expression is itself markedly increased by IL4. This suggests that PPARγ functions at the center of a feed-forward loop that is central to AA of macrophages.

[Keywords: macrophage; alternative activation; PPARγ; glutamine; rosiglitazone; metabolism]

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The nuclear receptor peroxisome proliferator-activated receptor γ (PPARγ) is the master regulator of adipogenesis and fat cell function [Rosen et al. 1999; Rangwala and Lazar 2004] and the target of anti-diabetic thiazolidinedione drugs [Lehmann et al. 1995]. Although its expression is predominant in adipocytes, PPARγ is also expressed in macrophages [Ricote et al. 1998; Chawla et al. 2001a; Lefterova et al. 2010]. Indeed, mice lacking macrophage PPARγ are prone to atherosclerosis [Chawla et al. 2001b; Babaev et al. 2005], and PPARγ ligands have anti-atherosclerotic properties [Nagy et al. 1998; Tontonoz et al. 1998; Akiyama et al. 2002].

In macrophages, PPARγ has been shown to play important roles in inflammation and metabolism [Hévener et al. 2007; Odegaard et al. 2007]. PPARγ is induced by interleukin 4 (IL4), a canonical inducer of the anti-inflammatory macrophage alternative activation (AA) pathway [Ricote et al. 1998; Bouhlel et al. 2007], and PPARγ ligands have anti-inflammatory properties [Huang et al. 1999; Li et al. 2000a; Stienstra et al. 2008]. Studies of its metabolic role have focused largely on lipid metabolism, and PPARγ has been shown to control lipid uptake [Chawla et al. 2001a; Odegaard et al. 2007], extrusion [Chawla et al. 2001b; Chimenti et al. 2001; Akiyama et al. 2002], and intracellular metabolism [Odegaard et al. 2007]. These properties have been linked to the role of PPARγ in inflammation and prevention of metabolic diseases [Chawla 2010].

Here, studying the regulation of macrophage metabolism by the thiazolidinedione rosiglitazone [Rosil], we report an unexpected role for PPARγ in glutamine metabolism. Glutamine is required for macrophage AA [Jha et al. 2015; Davies et al. 2017; Liu et al. 2017], and there is a remarkable overlap between the effects of glutamine depletion and the loss of PPARγ, providing a molecular and metabolic basis for the requirement of PPARγ for IL4-stimulated AA. IL4 also induces PPARγ, creating a feed-forward loop controlling AA, with PPARγ as the nexus.

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Results

Rosi activates a PPARγ-dependent gene program in macrophages

We generated homozygous macrophage-specific PPARγ knockout (MPKO) mice by crossing LysM-Cre mice to C57BL/6j PPARγ floxed mice. Quantification of mRNA isolated from thioglycollate-induced peritoneal macrophages showed almost complete depletion of PPARγ mRNA (Fig. 1A), and Western blotting confirmed loss of macrophage PPARγ protein (Fig. 1B). These cells were used to assess the role of PPARγ in the action of Rosi, a potent insulin sensitizer in the thiazolidinedione class that is activating ligands for PPARγ potent insulin sensitizer in the thiazolidinedione class. We next investigated whether Rosi induces PPARγ-dependent gene programs in macrophages. (Fig. 1A), and Western blotting confirmed loss of macrophage PPARγ protein (Fig. 1B). These cells were used to assess the role of PPARγ in the action of Rosi, a potent insulin sensitizer in the thiazolidinedione class that is activating ligands for PPARγ (Socco et al. 2014) but has been suggested to have other cellular actions.

Figure 1. PPARγ binds at the genome to control macrophage enhancer RNA (eRNA) and gene transcription. (A) Confirmation of PPARγ knockout assessed with quantitative PCR (qPCR) analysis of PPARγ mRNA in control and MPKO macrophages. Data are shown as mean ± standard error. n = 6. [*] P < 0.05. (B) Western blot showing expression of PPARγ in control and MPKO macrophages with RAN as a loading control. (C) Heat map showing z-transformed RPKM (reads per kilobase per million mapped reads) values of all statistically significant (P < 0.05) Rosi-responsive genes in control macrophages treated with either vehicle or 1 µM Rosi and corresponding expression in MPKO macrophages. Data are shown as biological replicates. n = 3–4. (D) Total PPARγ ChIP-seq (chromatin immunoprecipitation [ChIP] combined with high-throughput sequencing) tag count in reads per million [RPM] within 2 kb of up-regulated, down-regulated, and unregulated eRNA sites identified by GRO-seq [global run-on sequencing] in response to Rosi treatment. [***] P < 0.0001. Differentially expressed eRNAs were defined with edgeR as having fold change > 1.5 and false discovery rate (FDR) < 0.05 in Rosi-treated macrophages as compared with the untreated control. (E) PANTHER gene ontology analysis of the Rosi-induced genes identified in red in D with significantly increased PPARγ occupancy at Rosi-induced enhancer RNAs [eRNAs] exhibited a strong link to metabolism and metabolic processes (Fig. 1E).

PPARγ knockout macrophages have reduced respiration

We next investigated whether Rosi’s activation of metabolic pathways altered macrophage respiration. Indeed, Rosi increased both basal and maximal oxygen consumption rates (OCR) of control macrophages, and, consistent with the dependence of the Rosi-stimulated transcriptomes on PPARγ, this effect was abrogated in MPKO macrophages (Fig. 2A, B). Moreover, the baseline and maximal OCRs of MPKO macrophages were markedly attenuated relative to control even in the absence of exogenous PPARγ ligand (Fig. 2A, B). The striking effect of PPARγ depletion on macrophage respiration was independent of cell viability (Supplemental Fig. S1A) and mitochondrial density or number (Supplemental Fig. S1B).

PPARγ is not required for respiration from fatty acid or glucose

We next explored whether PPARγ was required for the use of specific energy sources for macrophage respiration. We first examined fatty acid oxidation, which has been reported to be defective in macrophages lacking PPARγ (Odegaard et al. 2007). Addition of palmitate to control macrophages in standard medium increased their respiration (Fig. 3A). Surprisingly, this was also the case for MPKO macrophages (Fig. 3A), whose defective basal and maximal OCRs were both largely rescued by palmitate (Fig. 3B).

We next considered whether the MPKO macrophages would be defective in glucose utilization. However, glucose uptake was not impaired relative to control macrophages (Supplemental Fig. S2A). Moreover, control and MPKO macrophages respired similarly well with glucose as the only fuel added to nutrient-free medium; in both cases, glucose initially suppressed respiration fueled by internal catabolic processes (potentially by activating

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we performed a normalized comparison of substrate effects on MPKO OCR relative to control macrophages, which makes clear the extent to which MPKO respiration is fueled by the fatty acid and glucose substrates but not by glutamine or αKG (Fig. 4C). Note that these experiments used fluoro-carbonyl cyanide phenylhyrazone (FCCP) at 2 µM, which was sufficient to maximally induce respiration of control macrophages with either glucose or glutamine as the only fuel provided [Supplemental Fig. S3]. Moreover, when respiration was attenuated by inhibition of fatty acid oxidation and glucose metabolism using etomoxir and UK5099, respectively, 10 mM glutamine largely restored oxidative capacity in control macrophages but had little effect on MPKO macrophages [Supplemental Fig. S4A,B], consistent with the conclusion that they are defective in glutamine oxidation.

**PPARγ is required for respiration from glutamine**

Our observation that PPARγ controls macrophage respiration from glutamine was intriguing because (1) glutamine is an important fuel source required for AA of macrophages [Jha et al. 2015; Liu et al. 2017], (2) PPARγ has been shown to be induced by IL4 [Huang et al. 1999], a classical inducer of AA, and (3) PPARγ has been reported to be
required for AA (Bouhlel et al. 2007; Odegaard et al. 2007). We confirmed that PPARγ was induced by IL4 in control macrophages [Fig. 5A]. Indeed, IL4 regulated hundreds of macrophage genes, including those involved in the AA response [such as Irf4, Ym1, and Ccl17] and genes critical to cell metabolism and the TCA cycle [such as Idh1 and Ccsl] [Fig. 5B]. Remarkably, most of the IL4 effect was abrogated in the absence of PPARγ [Fig. 5B]. Thus, although IL4 induces PPARγ expression, PPARγ is required for IL4 action.

Transcripts induced by IL4 in control but not MPKO macrophages were strongly enriched for metabolism-related gene pathways, including those implicated in the TCA cycle and aKG metabolism [Fig. 5C]. aKG is produced from glutamine and is critical for metabolism of AA macrophages (Palmieri et al. 2017). Indeed, IL4 stimulated the respiration of control macrophages but not MPKO macrophages [Fig. 5D], with PPARγ deletion impairing both basal and maximal respiration [Fig. 5E].

**Unstimulated MPKO macrophages have an inflammatory character that phenocopies the effect of glutamine depletion**

We next performed metabolic profiling on unstimulated control and MPKO macrophages. Itaconate stood out as the molecule with the largest and most statistically significant increase in the MPKO macrophages [Fig. 6A,B]. Itaconate is a metabolite with anti-microbial and immunomodulatory properties generated from the TCA cycle intermediate aconitate by the enzyme IRG1 (Michelucci et al. 2013; Jha et al. 2015). Indeed, Irg1 expression was

**Figure 4.** MPKO macrophages have impaired glutamine metabolism. (A) OCR in control and MPKO macrophages in nutrient-depleted medium with and without 2 mM glutamine as a sole fuel substrate. (B) OCR in control and MPKO macrophages in nutrient-depleted medium with only DM-aKG as a fuel substrate. (C) Basal OCR in the presence of only either 10 nM palmitate (Palm), 10 mM glucose (Glu), 2 mM glutamine (Gln), or 2 mM DM-aKG, shown as a percentage of basal respiration of nutrient-treated control OCR. Data are representative of at least three independent experiments. Data points are represented as mean ± SEM from n = 6–12 technical replicates. [*] P < 0.05.

**Figure 5.** PPARγ directs an IL4-mediated metabolic gene program. (A) Expression of PPARγ mRNA measured by qPCR in control macrophages treated with IL4 for 24 h. n = 5 vehicle; n = 6 IL4-treated. [*] P < 0.05. (B) Heat map showing z-transformed RPKM values of all statistically significant (P < 0.05) IL4-responsive genes in control macrophages and corresponding expression in MPKO macrophages. Selected IL4-induced genes involved in immunity and metabolism are listed at the right. Data are shown as biological replicates. n = 3–4. (C) PANTHER gene ontology analysis of genes increased with IL4 treatment in control macrophages but not in MPKO macrophages. (D) OCR in control and MPKO macrophages treated with and without 20 ng/mL IL4 for 24 h prior to assay. (E) Quantification of basal and maximal respiration in IL4-treated macrophages. Data are representative of at least three independent experiments. OCR data points are represented as mean ± SEM from n = 6–12 technical replicates. [*] P < 0.05.
increased in MPKO macrophages (Fig. 6C). Moreover, expression of *Idh1*, which directs the conversion of isocitrate to aKg in the TCA cycle, was decreased in MPKOs (Fig. 6D). The increase in *Irg1* and decrease in *Idh1* in MPKO macrophages likely leads to catapleurotic removal of TCA intermediates. In addition, the accumulation of itaconate may also impair oxidative metabolism by inhibiting succinate dehydrogenase (Cordes et al. 2016).

Itaconate accumulation occurs during M1-type inflammatory activation of macrophages induced by the classic inflammatory stimulus lipopolysaccharide (LPS) (Cordes et al. 2016). Consistent with this, MPKO macrophages secreted less anti-inflammatory IL10 (Fig. 6E) and more proinflammatory TNFα (Fig. 6F) relative to controls, indicating that, in the basal state, MPKO macrophages have an inflammatory phenotype. Indeed, there was significant overlap between a subset of gene changes induced by loss of PPARγ in unstimulated macrophages and those caused by LPS-treated wild-type macrophages (Fig. 6G,H). The overlapping genes and pathways suggest that PPARγ serves as a metabolic checkpoint in unstimulated macrophages that inhibits the inflammatory phenotype and favors AA (Fig. 6I).

Gene set enrichment analysis (GSEA) independently revealed that PPARγ-dependent IL4-responsive genes were strongly enriched for TCA cycle and respiratory electron transport pathways (Fig. 7A), with the greatest effect on key TCA enzyme *Idh1* (Fig. 7B). Indeed, there was a strong positive correlation globally between the up-regulation of gene expression by Rosi and IL4, including the TCA cycle genes (Fig. 7C). Since the respiratory defect in MPKO macrophages was due largely to the inability to oxidize glutamine and also because the MPKO macrophages are biased away from AA and toward inflammation, we compared the transcriptomes of IL4-stimulated glutamine-depleted macrophages (Jha et al. 2015), IL4-stimulated MPKO macrophages, and Rosi-stimulated MPKO macrophages to assess the potential of PPARγ in regulating AA-mediated glutamine metabolism. Remarkably, the effect of PPARγ knockout on the IL4 response was highly similar to that of glutamine depletion (Fig. 7D). Moreover, a similar transcriptional landscape, which includes the AA markers *Irf4* and *Ym1*, was found in Rosi-treated MPKO macrophages, suggesting that PPARγ regulates a shared group of mediators required for not only IL4 but also Rosi response (Fig. 7D). These data suggest that the function of PPARγ to support glutamine metabolism is central to its role in macrophage AA.

**Discussion**

We showed here that PPARγ controls macrophage glutamine metabolism, providing a link between transcription, AA, and metabolism. PPARγ is both required for IL4 stimulation of AA and amplified by IL4 signaling and thus serves as a nexus in a feed-forward pathway to AA (Fig. 7E).

The strong metabolic phenotype of the MPKO macrophages suggests that PPARγ is active in the absence of exogenous ligand either constitutively or due to endogenous ligands. Nonetheless, PPARγ-dependent gene regulation and metabolic activity are increased by Rosi treatment,
indicating that the basal state is not fully active and amenable to pharmacological manipulation. These activities of PPARγ and Rosi likely play a major role in the anti-diabetic and anti-atherosclerotic effects of Rosi in macrophages. Our finding that PPARγ binds at Rosi-regulated enhancers near genes that are induced by the PPARγ ligand strongly suggests that PPARγ directly regulates the genes that it activates.

The overlapping transcriptomes of LPS-treated macrophages and unstimulated MPKO macrophages support a model in which PPARγ acts as an anti-inflammatory checkpoint. Macrophage itaconate has been identified as a LPS-responsive metabolite that is capable of initiating metabolic remodeling and limiting AA cytokine production [Strelko et al. 2011; Lampropoulou et al. 2016; Mills et al. 2018]. This creates a break in the TCA cycle associated with the inflammatory phenotype and reduced oxidative respiration mirrored in the MPKO macrophages [Jha et al. 2015]. The elevation of itaconate in unstimulated MPKO macrophages thus provides further evidence of a dual role for PPARγ as both anti-inflammatory and permissive for AA.

Our work confirms previous reports that IL4 increases macrophage respiration [Vats et al. 2006]. Those studies of Balb/c macrophages chronically (96 h) treated with IL4 showed a role of PPARγ in fatty acid oxidation to promote oxidative phosphorylation in AA [Vats et al. 2006; Odegaard et al. 2007], while other studies suggest that fatty acid oxidation is not required for AA of C57BL/6 macrophages [Nomura et al. 2016; Gonzalez-Hurtado et al. 2017]. More recent studies demonstrate that glutamine metabolism is required for AA [Jha et al. 2015], likely via metabolism to αKG [Liu et al. 2017; Palmieri et al. 2017]. In this regard, although much of the literature on PPARγ in macrophage metabolism has focused on lipids [Castrillo and Tontonoz 2004], the importance of PPARγ in controlling glutamine metabolism fits with its known importance in AA [Huang et al. 1999; Odegaard et al. 2007].

Macrophages use glutamine at high rates in part to provide energy for oxidative phosphorylation [Newsholme et al. 1986; Jha et al. 2015; Davies et al. 2017]. Thus, the reduced expression of TCA enzymes in macrophages lacking PPARγ likely contributes to impaired glutamine metabolism. In particular, IDH1 was shown recently to strongly influence the activation state of the macrophage [Jha et al. 2015; O’Neill 2015]. Decreased TCA cycle function in unstimulated MPKO macrophages directly opposes the metabolic reprogramming required for AA. The requirement of PPARγ for glutamine metabolism and AA as well as the normal increase in PPARγ upon AA stimulation place PPARγ at the center of a feed-forward loop required for IL4 to induce AA.

Materials and methods

Animals

PPARγf/f and LysMCre C57BL/6 mice were purchased from Jackson Laboratories. Mice were housed in a temperature-controlled specific pathogen-free facility under 12-h light/dark cycles. Adult
male and female mice between the ages of 8 and 16 wk old were used in all experiments. Animal care and use procedures followed the guidelines of the Institutional Animal Care and Use Committee of the University of Pennsylvania in accordance with the guidelines of the National Institutes of Health.

Cell culture

Thioglycollate-elicited macrophages were harvested from the peritoneal cavity of mice. Four days after peritoneal injection with 3 mL of autoclaved 3% thioglycollate (Sigma-Aldrich), peritoneal macrophages were harvested from controls and MPKOs by lavage using Delbucco’s phosphate-buffered saline (PBS) [Invitrogen] and cultured overnight in DMEM supplemented with 10% fetal bovine serum [Tissue Culture Biologics], 100 U/mL penicillin, and 100 µg/mL streptomycin [Invitrogen]. When indicated, cultured macrophages were treated with DMSO vehicle, 1 µM Rosi [Cayman Chemical], or 20 ng/mL IL-4 [Peprotech] for 24 h. For metabolomics assays, control and MPKO macrophages were pooled from four mice and cultured in DMEM supplemented with 10% dialyzed fetal bovine serum [Tissue Culture Biologics], 100 U/mL penicillin, and 100 µg/mL streptomycin [Invitrogen].

Gene expression analysis

RNA was isolated from cells using TRIzol [Invitrogen] and purified with the Rneasy minikit [Qiagen]. RT–PCR was performed using 500 ng–1 µg of RNA [Applied Biosystems] following the manufacturer’s instructions. Quantitative PCR (qPCR) was performed on cultured macrophages using the primers PPARG (F: AGAAGCGGTGACACTGATTT, R: AAGGTTCCACA GAGCTTATTCC) and 18s (F: AGTCCGGCCCTTTGAC ACA; R: CGATCCGGGGGCTCATA) with SYBR Green master mix [Applied Biosystems] on the QuantStudio 6 Flex real-time PCR system [Applied Biosystems]. qPCR analysis was performed using the standard curve method, and all genes were normalized to the housekeeping gene 18s. For RNA sequencing (RNA-seq), RNA integrity was examined using an Agilent 2100 Bioanalyzer. RNA samples with RNA integrity number >7 were used for quantification with KAPA library quantification kit (Roche). RNA cleanup and library preparation were performed with the TRUEseq stranded total RNA library preparation kit according to the protocol provided by the manufacturer’s instructions [Illumina].

RNA-seq data processing

High-throughput sequencing data were either generated internally by the Functional Genomics Core at the University of Pennsylvania or downloaded from published Gene Expression Omnibus data sets cited in the text. All publicly obtained raw data, including the IL4-treated glutamine-depleted and LPS-stimulated macrophages [Jha et al. 2015], were subjected to the same analysis pipeline to minimize processing variations. Sequencing reads were aligned to the University of California at Santa Cruz (UCSC) mm9 genome using TopHat [version 2.1.9] [Kim et al. 2013] with the following parameters: –g 1 - -library-type fr-first-strand. Read counts were then obtained with featureCounts per the standard instructions [Liao et al. 2014]. Differentially expressed genes [cutoff defined as fold change > 1.5, false discovery rate [FDR] < 0.05 > 0.5 RPKM [reads per kilobase per million mapped reads]] were identified using edgeR with design matrices normalized for variations in size sequencing, depth, and dispersion [Robinson et al. 2010]. All gene ontology analyses were performed using PANTHER gene ontology enrichment analysis [Mi et al. 2013]. Heat maps were generated in R with the “gplots” package.

Liquid chromatography–mass spectrometry (LC-MS) analysis

Extracts were prepared as described in Lu et al. [2017]. In brief, cultured thioglycollate-elicited macrophages pooled from four control or four MPKO mice and further divided into four technical replicates were incubated on ice in a 40:40:20 acetonitrile: methanol:H2O + 0.5% formic acid solvent solution for 5 min before the cell lysates were collected and neutralized with a 15% NH4HCO3 in water solution for 15 min at ~20°C. The lysates were spun at maximum speed for 15 min at 4°C, and the supernatant was frozen on dry ice and stored at ~80°C before analysis. Cell extracts were analyzed using a quadrupole orbitrap mass spectrometer (Q Exactive, Thermo Fisher Scientific) coupled to a hydride interaction chromatography via electrospray ionization. LC separation was on a XBridge BEH amide column (2.1 mm × 150 mm, 2.5-µm particle size; Waters) using a gradient of solvent A (20 mM ammonium acetate, 20 mM ammonium hydroxide in 95:5 water:acetonitrile at pH 9.45) and solvent B (acetonitrile). Flow rate was 150 µL/min, column temperature was 25°C, autosampler temperature was 5°C, and injection volume was 10 µL. The LC gradient was 0 min and 90% B, 2 min and 85% B, 3 min and 75% B, 7 min and 75% B, 8 min and 70% B, 9 min and 70% B, 10 min and 50% B, 12 min and 50% B, 13 min and 25% B, 14 min and 25% B, 16 min and 0% B, 21 min and 0% B, 22 min and 90% B, and 25 min and 90% B. Autosampler temperature was 5°C, and injection volume was 3 µL. The mass spectrometer was operated in negative ion mode to scan from 70 to 1000 m/z at 1 Hz and a resolving power of 140,000 [Jang et al. 2018]. Data were analyzed using MAVEN software [Melamud et al. 2010].

ELISAs

Cytokine secretion into the supernatant of thioglycollate-elicited cultured macrophages was measured with mouse Quantikine ELISA kits [TNFa [MTA00B] and IL10 [M1000B]] according to the manufacturer’s instructions. Cytokine quantities in the supernatant were normalized to cell protein measured with BCA.

Immunoblotting

Primary antibodies for PPARγ (Santa Cruz Biotechnology) and RAN (BD Biosciences) were detected by a secondary horseradish peroxidase-conjugated antibody [Sigma] and an enhanced chemiluminescent substrate kit [PerkinElmer Western Lightning]. Immunoblots were imaged with Odyssey Clx infrared imaging system [Li-Cor] and uniformly contrasted.

ChIP-seq

ChIP was performed as described previously [Steger et al. 2008] using 200 µg/0.1 mL PPARγ [anti-PPARγ [H100], Santa Cruz Biotechnology, sc7196x, lot K1010] antibody. ChIP DNA was prepared for sequencing according to the amplification protocol provided by Illumina. Next-generation sequencing of ChIP-seq libraries was performed by the Functional Genomics Core at the University of Pennsylvania.
ChIP-seq data processing

Sequencing reads of PPARγ immunoprecipitated DNA of biological replicates for wild-type and PPARγ knockout macrophages (as input) were aligned to the UCSC mm9 genome using Bowtie1 (Langmead et al. 2009). Mapped reads were filtered for duplicates, biological replicates were pooled for read counts, and downstream peak calling and processing were done with HOMER version 4.6 per the instructions (Heinz et al. 2010). Peaks are defined as reads per million (RPKM) > 1, fold change immunoprecipitation/input > 4, and FDR < 0.001.

GRO-seq

GRO-seq was performed using thioglycollate-elicited peritoneal macrophages pooled from 20 mice and treated with DMSO or 1 µM Rosi 24 h prior to the harvest. Cells were washed twice with ice-cold PBS and then resuspended in cold swelling buffer (10 mM Tris at pH 7.5, 2 mM MgCl2, 3 mM CaCl2) for 5 min on ice. Cells were then resuspended and centrifuged at 400g for 10 min. The nuclei were then extracted using lysis buffer (swelling buffer with 10% glycerol and 1% Igepal). After washing twice with lysis buffer, the nuclei were resuspended in freezing buffer (50 mM Tris at pH 8.3, 40% glycerol, 5 mM MgCl2, 0.1 mM EDTA). For each library, 1.5 × 107 nuclei were used. The nuclear run-on reaction was incubated in run-on buffer (10 mM Tris at pH 8.0; 5 mM MgCl2, 1 mM DTT; 300 mM KCl; 200 U/mL Superscript-In; 1% Sarkosyl; 500 mM ATP, GTP, and BrUTP; 2 mM CTP) for 7 min at 30°C. Nascent transcripts were enriched with anti-BrUTP antibodies. After 10 min of RNA hydrolysis and reverse transcription, all products were used for GRO-seq library preparations in parallel to reduce batch effects as reported previously (Step et al. 2014).

GRO-seq data processing and eRNA analysis

FASTA file of trimmed (adapter and poly-A sequences) reads were generated prior to mapping. Bowtie1 was used for alignment to the UCSC mm9 genome with the following parameters: -n 1 -m 1. Downstream processing was done with HOMER version 4.6. Gene body transcription was calculated from processed .BED files. Differential expression of Rosi-modulated genes was identified using HOMER's findPeaks function according to the method per the manufacturer’s instructions (FDR < 0.05; RPMK > 0.5) eRNA analysis was done as described previously (Step et al. 2014). Briefly, HOMER’s findPeaks function was used to identify putative transcript on both positive and negative strands. Start sites within 1 kb of each other are defined as bidirectional. eRNA tags were counted similarly to gene body transcription except for the application of a 2-kb window around known enhancers to account for both positive and negative tags.

Accession numbers

All sequencing data, including RNA-seq, ChIP-seq, and GRO-seq, were deposited to the Gene Expression Omnibus database under the superseries GSE111105, which includes RNA-seq data (GSE111102), ChIP-seq data (GSE111104), and GRO-seq data (GSE111103).

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Flow cytometry

Thioglycollate-elicited peritoneal macrophages from control or PPARγ knockout mice were FC-blocked with CD16/32 (2.4G2, 1:50), BD Bioscience) and rat IgG (1:50, Fisher Scientific) prior to staining with anti-mouse fluorochrome-conjugated monoclonal antibodies [clone and concentration shown; BD Bioscience, BioLegend, eBioscience] specific for CD11b (1:50; BD Biosciences, F4/80 (BM8; 1:250). Cells were also stained with 100 nM MitoTracker red or 750 nM MitoTracker deep red (Thermo Fisher). Compensation was performed with cells or OneComp eBeads (Thermo Fisher). Cells were analyzed with a BD LSR II running DiVa software [BD Bioscience] and analyzed with FlowJo software [version 10.4, Tree Star].

Metabolic assays

Cells were uniformly plated in XF96 plates overnight before 24 h of stimulation with DMSO-vehicle, 1 µM Rosi, or 20 ng/mL IL4. OCR and ECAR were measured with the Seahorse XF96 extracellular fluid analyzer (Agilent). Experiments were conducted in XF medium [nonbuffered DMEM containing 10 mM glucose [Invitrogen], 2 mM L-glutamine [Invitrogen], 2 mM sodium pyruvate [Invitrogen]], with OCR measured basally and in response to sequential addition of 2 µM oligomycin, 2 µM FCCP, and 0.5 µM rotenone + antimycin A. Fatty acid oxidation was measured in medium supplemented with a final concentration of 10 nM free palmitate bound to BSA or an equal volume of BSA control (Agilent). Glycolysis was measured as ECAR at baseline and after the addition of 0.2 µM oligomycin to impair respiration. Fuel preference was measured after 1 h of incubation in Seahorse XF base medium (Agilent) with no fuel substrates and then immediately before the assay supplemented with either 10 mM glucose, 4 mM glutamine, 4 mM DM-αKG (Sigma-Aldrich), or 10 nM free fatty acids in the form of palmitate BSA with unbound BSA as a control. All Seahorse XF data were normalized to total well protein quantified with Pierce BCA protein kit (Thermo Fisher). All calculations for respiration were corrected for nonmitochondrial respiration. Glucose uptake was measured with 1 µCi/mL 2-deoxy-D-[3H] glucose (Sigma-Aldrich) in cultured macrophages following 2 h of incubation in serum-free medium. 3H glucose uptake was detected with a PerkinElmer Tri-Carb 4910 TR scintillation counter.

Author contributions: V.L.N. and H.C.B.N. were involved in project planning, experimental work, data analysis, and manuscript preparation. J.C.G.-C. performed experimental work and data analysis. E.R.B. was involved in project planning. W.Y.H. was involved in experimental work. J.R.D. was involved in project planning and experimental work. J.M.M. was involved in experimental work. D.A.H. was involved in project planning and
experimental work. M.A.L was involved in project planning, data analysis, and manuscript preparation.

References


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