Adipocyte SIRT1 knockout promotes PPARγ activity, adipogenesis and insulin sensitivity in chronic-HFD and obesity

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ABSTRACT

Objective: Adipose tissue is the primary site for lipid deposition that protects the organisms in cases of nutrient excess during obesogenic diets. The histone deacetylase Sir2 (SIRT1) inhibits adipocyte differentiation by targeting the transcription factor peroxisome proliferator activated receptor gamma (PPARγ).

Methods: To assess the specific role of SIRT1 in adipocytes, we generated Sirt1 adipocyte-specific knockout mice (ATKO) driven by aP2 promoter onto C57BL6 background. Sirt1+/+ aP2Cre+ (ATKO) and Sirt1+/+ aP2Cre− (WT) mice were fed high-fat diet for 5 weeks (short-term) or 15 weeks (chronic-term). Metabolic studies were combined with gene expression analysis and phosphorylation/acetylation patterns in adipose tissue.

Results: On standard chow, ATKO mice exhibit low-grade chronic inflammation in adipose tissue, along with glucose intolerance and insulin resistance compared with control fed mice. On short-term HFD, ATKO mice become more glucose intolerant, hyperinsulinemic, insulin resistant and display increased inflammation. During chronic HFD, WT mice developed a metabolic dysfunction, higher than ATKO mice, and thereby, knockout mice are more glucose tolerant, insulin sensitive and less inflamed relative to control mice. SIRT1 attenuates adipogenesis through PPARγ repression and, in the ATKO mice adipocyte PPARγ was hyperacetylated. This high acetylation was associated with a decrease in Ser273-PPARγ phosphorylation. Dephosphorylated PPARγ is constitutively active and results in higher expression of genes associated with increased insulin sensitivity.

Conclusion: Together, these data establish that SIRT1 downregulation in adipose tissue plays a previously unknown role in long-term inflammation resolution mediated by PPARγ activation. Therefore, in the context of obesity, the development of new therapeutics that activate PPARγ by targeting SIRT1 may provide novel approaches to the treatment of T2DM.

Keywords Obesity; SIRT1; PPAR03B3; Glucose homeostasis; Insulin resistance; Phosphorylation

1. INTRODUCTION

Obesity is a complex metabolic disorder associated with insulin resistance, glucose intolerance and hepatic steatosis. In this syndrome, the white adipose tissue plays an important role as the primary storage site for lipid deposition. Increased caloric intake causes an expansion of adipocyte size, with subsequent activation of stress pathways, leading to metabolic deterioration and decreased insulin sensitivity. This process is associated with increased infiltration of inflammatory cells into the adipose tissue, which contribute to the development of insulin resistance [1,2].

Sirtuin 1 (SIRT1, one of the seven mammalian sirtuins) is the ortholog of the yeast protein Silent information regulator 2 (Sir2) and a nuclear NAD+−dependent class III histone deacetylase class III which engages in reciprocal co-regulation of many different binding partners [3,4]. SIRT1 is regulated by cellular NAD+ levels and several studies have shown that its expression and activity increases during fasting or calorie restriction whereas decreases by over nutrition in rodents and humans [3,5,6]. Thus, SIRT1 couples host metabolic status to gene expression regulation through deacetylation of histones, transcription factors, and transcriptional co-regulators. In this way, SIRT1 controls metabolic homeostasis via the integration of diverse actions across different tissues, including brain, liver, pancreas and adipose tissue [3,7].
In adipose tissue, SIRT1 controls lipolysis and inhibits inflammation by repressing PPARγ and NFκB activity [8,9]. Furthermore, SIRT1 expression is inversely correlated with macrophage infiltration in subcutaneous fat from obese humans [10]. Thus, activation of SIRT1 has been proposed as a therapeutic target for the treatment of obesity and associated metabolic disorders [11,12]. However, this proposition is contradictory to the proven benefits of enhancing PPARγ activity in obese/diabetic individuals [13], and inhibition of PPARγ activity in insulin resistant subjects requires careful study.

In the current work, we sought to better understand the physiological role of adipocyte SIRT1 with the aim of resolving this contradiction and clarifying its therapeutic potential by generating an adipocyte-specific knockout mouse (ATKO). We report clear evidence that selective adipocyte deletion of SIRT1 exacerbates the detrimental effects of acute HFD-feeding. Strikingly, we find protective effects of SIRT1-deletion in the context of chronic HFD exposure. These beneficial outcomes of SIRT1-deletion were associated with increased PPARγ activity resulting from hyperacetylation and dephosphorylation of PPARγ Ser273 along with reduced CDK5 activity. Thus we propose that, in the context of chronic HFD/obesity, inhibition of SIRT1 in adipocytes can result in improved metabolic functions.

2. MATERIALS AND METHODS

2.1. Animals
We backcrossed mice carrying Sirt1 floxed alleles (fl/fl mice) [14] onto C57BL/6 background for more than 6 generations. Mice were bred with transgenic mice harboring Cre recombinase driven by αP2 promoter [15,16] to create the following genotypes: WT (Sirt1\textsuperscript{fl/fl} αP2Cre\textsuperscript{−/−}) or ATKO (Sirt1\textsuperscript{fl/fl} αP2Cre\textsuperscript{−/−}). Mice were housed on a 12-h light/dark cycle and given ad libitum access to food and water. Mice were fed either with NC (13.5% fat; LabDiet) or HFD (60% fat; D12492, Research Diets) starting at 8 weeks of age. All protocols were approved by the Animal Subjects Committee of the University of California, San Diego.

2.2. Adipocytes, stromal vascular fraction, and peritoneal macrophages isolation
Adipocytes and SVs were prepared from collagenase-digested adipose tissue, as described previously [16]. Peritoneal macrophages were prepared from WT and ATKO mice as previously described [17].

2.3. RNA isolation and real-time PCR
Total RNA was isolated with TRIzol reagent (Invitrogen) and cDNA was synthesized with Transcriptor First Strand cDNA Synthesis Kit (Roche Applied Science). Semi-quantitative PCR was performed with MyTaq DNA Polymerase (Bioline) and quantitative real-time PCR (qPCR) with iTaq SYBRgreen Supermix (BioRad). We calculated relative gene expression levels by ∆∆Ct method using 36B4 (Rplp0) as internal control. Expression of genes regulated by the phosphorylation of PPARγ was described before by Choi et al. [18]. Primers are summarized in Table S1.

2.4. Western blot analysis
Proteins from total tissue lysates were prepared as previously described [19], separated by SDS-PAGE and probed with different primary antibodies against Sirt1, Anti-phospho-Akt (Ser473), total Akt, PPARγ, phospho-CDK5 (Tyr15), CDK5, and Hsp90β (Santa Cruz); Acetyl (Lys379) p53, p53 and NF-кB p65 (Cell Signalling); Acetyl (Lys310) NF-кB p65 and FGFr2 (Abcam); Anti-phospho- PPARγ (Ser 273) was a kind gift from Dr. Bruce Spiegelman. For immunoprecipitation (IP), eWAT was homogenized in IP Lysis/Wash Buffer and 500 µg of total protein were incubated with the antibody-conjugated magnetic cross-linked beads for 1 h at RT. The immuneprecipitates were eluted according to the manufacturer’s instructions (Thermo Scientific). In order to avoid interferences with denatured rabbit IgG heavy chains, we used a mouse anti-rabbit IgG conformation specific as a secondary antibody (L27A9, Cell Signaling). The antibody-conjugated magnetic beads were prepared with 5 µg of PPARγ (H100) (Santa Cruz). Blots were visualized using a CCD camera in a Molecular Imaging System (Chemidoc XRS+, Bio-Rad).

2.5. Metabolic studies
Glucose and insulin tolerance tests were performed on mice fasted for 6 h by intraperitoneal injection of 1 g/kg dextrose (Hospira) or 0.6 U/kg insulin (Novolin R, Novo-Nordisk) respectively. Blood glucose concentrations were measured at 0, 15, 30, 60, 90 and 120 min after dextrose or insulin injection. Plasma levels of insulin were measured by ELISA (ALPCO). Hyperinsulinemic-euglycemic clamps were performed after 6 h of fasting using a method described previously [20]. Plasma FFA levels were measured enzymatically (WAKO Chemicals). Acute insulin challenge experiments were performed on anesthetized mice fasted for 6 h. Basal samples of liver, muscle, and fat were taken and vessels were ligated. Five minutes after injection via inferior vena cava with 0.6 U/kg of insulin, the remaining liver, muscle, and fat were snap frozen for subsequent protein extraction. Glucose uptake in eWAT explants was performed as previously described [21].

2.6. Confocal microscopy and determination of adipocyte cell size
The method for immunofluorescence study of mouse adipose tissue was followed as described [22]. Adipocyte area was measured from each section using Image Gauge v4.0 software (Fujifilm). Frequency, histograms and Gaussian distributions were calculated using Excel 2013 (Microsoft) and GraphPad Prism v6.0 (GraphPad Software).

2.7. Statistical analysis
Results are presented as the mean ± SEM. Statistical significance was determined using GraphPad Prism v6.0 (GraphPad Software) or Excel 2013 (Microsoft) at P < 0.05. Student t-test was used to compare differences between two groups. Two-way ANOVA and Bonferroni posttest were used to analyze multiple experimental groups.

3. RESULTS

3.1. Adipocyte-specific Sirt1 knockout mice (ATKO)
In order to investigate the specific role of adipocyte SIRT1 on the development of insulin resistance, we generated an adipocyte-specific knockout mouse (ATKO) using the Cre-lox system. We bred mice carrying the floxed Sirt1 allele, containing two loxP sites flanking exon 4, which encodes the deacetylase domain, to transgenic mice expressing Cre recombinase driven by the α2P promoter [10,14,15]. In this model, restricted Cre expression in both white and brown adipose tissue results in a smaller SIRT1 protein with an intact C terminus but without deacetylase activity [23]. Floxed Sirt1 and Cre genotypes were determined from genomic DNA (Figure 1A), and the generated genotypes were denominated WT (Sirt1\textsuperscript{fl/fl} α2PCre\textsuperscript{−/−}) or ATKO (Sirt1\textsuperscript{fl/fl} α2PCre\textsuperscript{−/−}). Growth and fertility between ATKO mice and their control littermates were comparable. We verified by PCR from DNA that the excision of exon 4 occurs only in fat tissues and is restricted to adipocytes. We have found no-Cre mediated target gene recombination in liver, spleen, kidney, muscle or peritoneal macrophages (∆4 ATKO, Figure 1B,C). The exon 4 deletion was predominantly detected in mature adipocytes and not in cells from the
stromal vascular fraction (SVF) (Figure 1D,E). To confirm the magnitude of the SIRT1 deacetylase site deletion, we measured the acetylation status of NF-κB, one well-known SIRT1 target. In adipocytes, increased p65 acetylation was detected in ATKO mice (Figure 1F). In addition, WT and ATKO mice did not show any differences in gene expression levels of Sirt2 and Sirt3 in white and brown adipose tissues (Figure 1G).

3.2. Adipocyte Sirt1 deletion causes insulin resistance on chow diet

Due to the described role of SIRT1 as an inhibitor of inflammatory pathways and modulator of insulin sensitivity in macrophages and adipocytes [9,24], we analyzed the metabolic phenotype of mice with adipocyte specific deletion of Sirt1. Initially the body weights of ATKO and WT mice fed NCD were comparable until 23 weeks of age after which the ATKO mice gained significantly more weight than WT controls (Figure 2A). At 15 weeks of age, we harvested fat depots from WT and ATKO mice fed NCD and found that ATKO mice showed increased subcutaneous, epididymal, and brown adipose tissue (Figure 2B). Before the divergence in body weight (Figure 2C), we conducted glucose tolerance and insulin tolerance tests. ATKO mice at 19 weeks of age were hyperinsulinemic, with impaired glucose and insulin intolerance compared to WT mice (Figure 2D,E and F). Indeed, elevated basal insulin levels could be detected in ATKO mice as soon as 8 weeks of age although no differences in glucose tolerance were observed at this age (Figure 2D,E). We also measured insulin-stimulated glucose uptake in primary adipocytes, and we confirmed that the ability of insulin to enhance glucose transport was reduced in ATKO versus WT mice (Figure 2G).

Figure 1: Quantification of adipocyte-specific deletion of Sirt1 in mice (ATKO). A: DNA genotyping. Upper panel, Sirt1lox/lox (floxed-ATKO) and Sirt1+/+ (WT) DNA expression. Lower panel, Cre-Recombinase (all mice are Sirt1lox/lox). C- Cre negative control. C+ Cre positive control. B: Semi-qPCR showing the Sirt1 floxed exon 4 (∆4 ATKO) in several adipose depots and other different tissues from WT and ATKO mice. 36b4 (Rplp0) expression was used as housekeeping gene. C: Sirt1 mRNA expression in intraperitoneal macrophages and epididymal white adipose tissue (eWAT). D: Semi-qPCR showing Sirt1 mRNA expression in adipocytes or Stromal vascular fraction (SVF) from WT and ATKO eWAT. E: Protein expression of SIRT1 determined by Western Blot in adipocytes isolated from eWAT. Hsp90 expression was used as loading control. The Western blots shown are representative of three independent experiments (**P ≤ 0.01 vs. WT). G: Relative mRNA expression of Sirt2 and Sirt3 in different adipose tissues from WT and ATKO mice after 15 weeks of HFD.
3.3. The impact of adipocyte SIRT1 deletion on the metabolic response to HFD is time-dependent

Next, we studied the metabolic effects of adipocyte Sirt1 deletion during short term and chronic HFD. WT and ATKO mice were fed 60% HFD for 5 weeks (short term; S) or 15 weeks (chronic; C) starting at 8 weeks of age. Body weight (BW) and adipose tissue mass were increased in ATKO mice compared to WT (Figure 3A,B), and this was evident in the first week of HFD and maintained throughout the treatment period. No differences in food intake were noted between the groups (Figure 3C). To further evaluate body composition changes, Dual energy X-ray absorptiometry (Dexa Scan) analyses were performed after 15 weeks of HFD (Figure 3D). ATKO mice exhibited increased total body area (TBA), fat mass, percentage of fat, and decreased bone mineral density (BMD), with no changes in lean body mass (LBM) compared to WT mice.

During the short term HFD feeding, ATKO mice showed exaggerated glucose intolerance, hyperinsulinemia and a reduced hypoglycemic effect of insulin compared to WT mice (Figure 4A–D). However, after chronic HFD feeding (15 weeks), ATKO mice were relatively more glucose tolerant and displayed an increased hypoglycemic response to insulin compared to WT (Figure 4A–D). In addition, basal insulin levels plateaued in the ATKO mice from short to chronic HFD, while they continued to increase in the WT mice with the development of obesity (Figure 4D), so that the ATKOs were no longer more hyperinsulinemic compared to WT. In other words, the metabolic dysfunction progressively worsened going from short term to chronic HFD in WT mice, whereas it plateaued, or improved, in the ATKO group. The result is that ATKO mice are less glucose tolerant and more insulin resistant than WT mice on short term HFD, but more glucose tolerant and less insulin resistant than WT mice on chronic HFD.

To further assess the difference in insulin sensitivity on chronic HFD, we performed hyperinsulinemic-euglycemic clamp studies in body weight matched WT and ATKO mice (Figure 4E). The glucose infusion rate (GIR) required to maintain euglycemia, and the ability of insulin to suppress hepatic glucose production (HGP) were both significantly increased in ATKO mice compared to WT mice. Insulin suppression of free fatty acid levels was slightly increased in ATKO mice (but not statistically significant). There was no impact on basal and insulin-stimulated glucose disposal rate (GDR and IS-GDR), suggesting muscle insulin sensitivity is not altered between groups. Consistent with the glucose clamp studies, insulin-stimulated Akt phosphorylation was significantly increased in liver and adipose tissue in ATKO mice compared with WT mice (Figure 4F,G). Thus, ATKO mice clearly display improved insulin sensitivity relative to WT mice after chronic HFD feeding, despite the increase in body weight and adiposity.

3.4. ATKO mice show reduced eWAT inflammation after chronic HFD

Inflammation is a key causative factor in insulin resistance [1]. Consistent with the relative insulin sensitive state of the chronic HFD ATKO mice, we found decreased inflammation both systemically (Figure 5A) and locally in epididymal adipose tissue (Figure 5B). ATKO mice had lower circulating levels of MCP-1, TNF-α and PAI-1 (Figure 5A) and lower expression of proinflammatory genes Mcp-1, Tnf-α (Figure 5B). Furthermore, increased expression of Il-10, an important anti-inflammatory gene and Arginase, a macrophage M2-associated marker, were detected in ATKO epididymal adipose tissue (Figure 5B,F).

We also measured the levels of FGF21 in plasma after chronic HFD feeding. FGF21 is a circulating factor, upregulated by fasting and HFD in liver and adipose tissue, that is associated with insulin sensitivity [25], plays a role in beiging of adipose tissue [26] and also is selectively repressed by SIRT1 in adipocytes [27]. SIRT1 attenuates PPARγ activity in adipocytes resulting in decreased FGF21 production [27]. In our model, ATKO mice displayed higher...
circulating levels of FGF21 (Figure 5A) as well as up-regulation of \(Fgf21\) gene expression in eWAT (Figure 5B). The elevated \(Fgf21\) expression, and particularly the increased circulating FGF21 levels, appear to provide at least one component for the insulin sensitivity in the chronic HFD ATKO mice. A similar upregulation was observed for \(Ucp1\), a FGF21 target gene [28], in eWAT from ATKO mice fed HFD for an extended period of time (Figure 5C), as well as increased expression of other brown fat-selective genes, including \(PGC1\)-\(a\), \(Cox7a1\) (Figure 5C).

To investigate the effect of ATKO on macrophage infiltration in eWAT, we performed immunostaining for the macrophage marker F4/80 and visualized the number of crown-like structures (CLS). Short term feeding of HFD to ATKO mice resulted in an increased number of CLSs (Figure 5D) and increased expression of macrophage markers CD68, CD11b, and CD11c (Figure 5F) compared with WT control mice. Interestingly, after chronic HFD feeding, ATKO mice exhibited reduced macrophage accumulation compared to WT, fully consistent with their relative insulin sensitivity at this time point (Figure 5E,F).

Taken together, these results show that while inflammation continues to progress in WT mice between short term and chronic HFD feeding, this progression fails to occur in ATKO mice, resulting in a relative reduced inflammatory status compared with WT mice at the chronic time point.

3.5. ATKO mice show increased adipocyte hypertrophy/hyperplasia during HFD

In the early stages of HFD feeding, hypertrophy drives adipose tissue expansion in both eWAT and sqWAT. However, at more prolonged stages of HFD, only eWAT retains adipogenic capacity [29,30]. We studied the adipocyte-architecture in short term and chronically fed HFD ATKO and WT mice. The eWAT adipocytes from short term HFD-fed ATKO mice were significantly larger than from WT HFD fed mice (Figure 6A,B and C). During chronic HFD, the eWAT from ATKO mice underwent a significant change in adipocyte-architecture (Figure 6D). Adipocytes from ATKO mice were smaller (Figure 6B,E) with an overall 25% decrease in average cell size compared to WT (Figure 6F). This suggests that hyperplasia plays a major role in eWAT expansion in ATKO mice, and that deletion of \(Sirt1\) partially protects from the hypertrophy seen in WT eWAT. These effects were specific to eWAT as no significant differences in cell size were observed in sqWAT or lipid droplet size in BAT (Figure 6G).

3.6. ATKO mice display PPAR\(\gamma\) hyperactivity resulting from increased acetylation and reduced Ser273 phosphorylation

Enhanced adipogenic and anti-inflammatory responses in the adipose tissue can be induced by hyperactivity of PPAR\(\gamma\), a well characterized SIRT1 target. SIRT1 inhibits adipocyte differentiation by deacetylation...
of PPARγ [8,31,32]. Consistent with this, ATKO mice showed enhanced PPARγ acetylation after both, short term and chronic HFD feeding (2.5 and 4.2 times higher respectively, Figure 7A). Interestingly, WT mice fed chronic HFD also displayed enhanced PPARγ acetylation (2.4x), most likely as a result of SIRT1 downregulation in adipose tissue [10]. Furthermore, we analyzed a subset of genes previously reported to be repressed by PPARγ [33], showed lower expression in ATKO mice during short term and chronic HFD, while in WT mice the downregulation is progressive (Figure 7B).

PPARγ activity is also regulated by phosphorylation of serine 273 by CDK5 [34]. CDK5 itself is activated by phosphorylation at tyrosine 15 and binding to the co-activator p25. In obesity and diabetes, pro-inflammatory signals lead to increased levels of cytoplasmic p25 with consequent activation of CDK5 and phosphorylation of PPARγ.
Since PPARγ Ser273 phosphorylation is inhibitory, whereas dephosphorylation confers an active insulin sensitizer transcriptional program [16,18], we hypothesized that the enhanced insulin sensitivity observed in chronic HFD-fed Sirt1 ATKO mice might be related to PPARγ phosphorylation status.

In normal chow and short term HFD fed mice, pSer273-PPARγ and pTyr15-CDK5 levels were increased consistent with the insulin resistant phenotype (Figure 7C,D). However, pSer273-PPARγ and pTyr15-CDK5 levels were reduced in ATKO compared to WT mice after 15 weeks of HFD feeding (Figure 7C,D). Furthermore, the set of genes
Figure 6: ATKO mice show higher hyperplasia during chronic HFD. A: Immunofluorescence of eWAT from short term HFD feeding. Adipose tissue was stained with Caveolin (blue) as adipocyte membrane marker. Scale bar is 200 μm. The images below are 4x magnification corresponding to the white box on the upper images. B-C: Quantification, area and adipocyte size frequency measurement from eWAT immunofluorescence images. D-F: Same analysis in WT and ATKO eWAT from chronic HFD feeding mice. Values are expressed as means ± SEM. (n = 4 mice/group and 2 images/mouse were analyzed). G: Hematoxylin & Eosin of eWAT, sqWAT, and BAT from WT and ATKO mice from chronic HFD-fed mice. Fat tissues were fixed with 4% PFA, embedded in paraffin blocks and sectioned in the UCSD Mouse Phenotype Service Core (Moore Cancer Center, UCSD). Bright field photographs were taken using a Zeiss Observer.Z1 microscope. Scale bar is 100 μm. (n = 4 mice/group and 2 images/mouse were analyzed).
repressed by phosphorylated PPARγ described by Choi et al. were downregulated in ATKO mice fed NCD or short term HFD (Figure 7E,F) but were upregulated in ATKO following chronic HFD feeding (Figure 7G). Thus, we suggest that protection of ATKO mice from long-term impact of HFD feeding is due to PPARγ hyperacetylation and dephosphorylation, resulting in sustained PPARγ activity. This leads to reduced inflammation, decreased CDK5 phosphorylation and upregulation of genes involved in insulin sensitization (summarized in the Graphical Abstract).

4. DISCUSSION

It is well known that obesity leads to a state of chronic, sub-acute tissue inflammation with increased accumulation of adipose tissue...
macrophages (ATMs) and this is a key driver of the insulin resistance syndrome [1, 2, 36]. Recently we have found that moderate and severe obesity were associated with a significant reduction in Sirt1 mRNA expression and an increase in ATF content in human adipose tissue [10]. It has been reported that Sirt1 can exert anti-inflammatory effects and knockdown of Sirt1 in vivo using antisense-oligonucleotides (ASOs) promotes inflammation and AT accumulation [10]. However, in these studies, macrophage-deficient mice showed a similar induction of inflammation after treatment with Sirt1 ASO, suggesting that in addition to its influence on ATMs, SIRT1 activity might also be important in adipocytes. To address this question, we studied the role of adipocyte SIRT1 on inflammation, glucose metabolism and insulin resistance in adipocyte-specific Sirt1 knock-out mice (ATKO) in the context of an acute or chronic HFD/obesity. Given the important anti-inflammatory effects of SIRT1 [9, 24], we found, as expected, that NCD fed ATKO mice showed impaired glucose tolerance and were insulin resistant compared to WT controls. When challenged with a HFD, ATKO mice initially become more inflamed and developed glucose intolerance, hyper-insulinemia and insulin resistance faster than WT mice. Surprisingly, as the time period of HFD feeding continued, the ATKO mice phenotype plateaued while the WT mice continued to develop metabolic disease. Therefore, ATKO mice fed chronic HFD showed reduced inflammation, improved glucose tolerance and enhanced insulin sensitivity, relative to their control WT mice. This phenotype was unexpected, and we were able to trace the mechanisms to increased PPARγ activity, higher adipogenesis and reduction of inflammation during the latter stages of HFD. Thus, in obesity, adipocyte PPARγ was largely hyperacetylated and unphosphorylated with a constitutively active state in the ATKO mice [34]. This was also associated with upregulation of brown fat-selective genes and induction of a variety of WAT target genes associated with insulin sensitization [18, 33]. A decrease in the adiponectin secretion has been associated with insulin resistance in mouse models of impaired insulin sensitivity [37], and this effect was reproduced in our experiments in the WT HFD-fed mice. However, no significant changes were detected in ATKO HFD-fed mice between short or chronic HFD (data not shown), this may be due to the involvement of Sirt1, FOXO1 and C/EBPβ interaction in the adiponectin gene expression [38, 39], and therefore, the reduction of adiponectin plasma levels in WT HFD-fed mice could be related to the SIRT1 downregulation previously described in 3T3-L1 adipocytes and obesity/HFD [10, 40].

Much attention has been given to the initiators of inflammation in obesity, with relatively little focus on the resolution of inflammation. Normally, tissue inflammation is resolved through a series of complex immune mechanisms [41]. In obesity, the chronic state of inflammation could be, at least in part, a reflection of inadequate resolution. Thus, in the normal obese state, resolution mechanisms come into play, but they might be inadequate to dissipate the inflammation and can only maintain a sub-acute chronic inflammatory state. Wernstedt Asterholm et al. have recently shown that adipose tissue inflammation is an adaptive response, essential for storage of excess nutrients and contributes to a healthy expansion and remodeling of the adipose tissue [42]. Also, Wang et al. have found that eWAT adipose tissue, but not sqWAT, undergoes hypertrophy first and hyperplasia later during chronic HFD [29]. In our model, during short term HFD, the ATKO mice displayed increased inflammation, coupled with eWAT hypertrophy compared to the WT (Figures 5 and 6). However, the population of adipocytes after chronic HFD is phenotypically different from those in the short term HFD and, therefore, by increasing adipogenesis, hyperplasia plays a major role in eWAT expansion in ATKO mice during chronic HFD (Figure 6). The exact mechanism underlying the proliferation of smaller, more insulin sensitive, adipocytes due to the Sirt1 deletion in eWAT remains unclear, but we did find increased expression of Cd24 in ATKO mice during chronic HFD, which is a marker of adipocyte progenitor cells [43, 44] (Figure 7H). This is also consistent with increased adipocyte PPARY activity [45, 46]. Indeed, it is known that PPARγ exhibits ligand-independent activation by protein acetylation through downregulation of SIRT1 [32], and the eWAT from ATKO mice showed high levels of acetylated PPARγ during chronic HFD. Although Bouant et al. recently found that SIRT1 improves glucose homeostasis by enhancing BAT function [47], we did not find significant differences in cell size or lipid droplet size in BAT (Figure 6G). Interestingly, Kusminski et al. found that during the initial stages of a HFD challenge, the outer mitochondrial membrane protein MitoNEET is necessary to induce the browning program to dissipate surplus dietary energy for up to 12 weeks [48]. However, after 12 weeks HFD, the adipocytes lose their ability to maintain the browning program and this is associated with adipocyte death [49]. This mechanism could relate to the increased expression of Cd24 in ATKO mice. In our model, during chronic HFD, ATKO mice exhibit a decrease in the inflammatory state with improvement in glucose tolerance. This raises the possibility that downregulation of SIRT1 plays a previously unknown role in long-term inflammation resolution mediated by PPARγ activation. An interesting mechanism that relates to the more insulin sensitive state in the ATKO mice after long-term HFD, could be related to the regulation of PPARγ activity by phosphorylation. It is known that PPARγ exhibits ligand-independent insulin sensitizing effects in the unphosphorylated state. Thus, Spiegelman et al. and others have demonstrated that PPARγ S273 phosphorylation is increased in the obese state and that CDK5, which is also activated in obesity and diabetes, can phosphorylate PPARγ at this site [18, 35]. Unphosphorylated PPARγ induces a set of target genes which can promote insulin sensitivity. Indeed, treatment of cells or mice with classical and non-classical thiazolidinediones (TZDs) such as rosiglitazone or MRL24 respectively, prevents PPARγ phosphorylation, inducing a comparable target gene expression signature, along with systemic insulin sensitization [18]. Moreover, SIRT1 can be phosphorylated at serine 47 (S47) by CDK5, which inhibits its anti-inflammatory functions in endothelial cells and blocks the anti-senescence activity of SIRT1 [50]. This post-translational regulation of SIRT1 may contribute to the inflammatory phenotype in WT mice. In our studies, the reduction in inflammation of the chronic HFD ATKO mice might explain the decrease in CDK5 Tyr15 and PPARγ S273 phosphorylation. This process was associated with induction of the same set of PPARγ target genes reported by Spiegelman et al. [18]. Consistent with the studies of Accili et al., who showed that decreased PPARγ phosphorylation can be mediated by increased acetylation and therefore a reduction in inflammation, we also show that during chronic HFD adipocyte Sirt1 deletion leads to hyperacetylation of PPARγ and low profile of phosphorylation, [51]. The regulation of PPARγ activity by phosphorylation in S273 (negatively) and acetylation in K286 and K293 (positively) are different mechanisms and they are not related directly. While the acetylation of PPARγ is p300/SIRT1 dependent, the phosphorylation in S273 is CDK5 dependent. Furthermore, CDK5 activity is regulated by its phosphorylation at Y15 and by calcium-dependent proteases (Calpains) that are highly activated in obesity, diabetes, atherosclerosis, and inflammation [35, 52]. We have described that SIRT1 inhibits NFkB in macrophages and plays a role in adipose tissue inflammation [10]. NFkB induces expression of PTPIB, a negative regulator of insulin sensitivity [53]; and SIRT1 represses PTPIB in insulin resistant obese mice improving insulin
Neuronal Sirt1 deletion is protective against metabolic disease [23], and one recent report describes that inhibition of SIRT1 by RNAi or HDAC inhibitors promotes adipogenesis, and enhances insulin sensitivity [32]. We traced this phenotypic change to the acetylation/phosphorylation status of PPARγ (summarized in the Graphical Abstract). Future experiments comparing the gene expression pattern of the chronic HFD- SIRT1 ATKO mice with the pattern induced by TZD treatment, would be of interest in order to highlight commonalities and differences. These studies should inform potential SIRT1-based therapies, since such treatments would be chronic in nature and might have different effects over the long haul compared to the short term.

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CONFLICT OF INTEREST

None declared.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.molmet.2015.02.007

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