Disruption of Adenosine 2A Receptor Exacerbates NAFLD through Increasing Inflammatory Responses and SREBP1c Activity

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This material is the result of work partly supported by resources at the Central Texas Veterans Health Care System. The views expressed in this article are those of the authors and do not necessarily represent the views of the Department of Veterans Affairs.

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ABSTRACT

Adenosine 2A receptor (A$_{2A}$R) exerts protective roles in endotoxin- and/or ischemia-induced tissue damages. However, the role for A$_{2A}$R in non-alcoholic fatty liver disease (NAFLD) remains largely unknown. We sought to examine the effects of global and/or myeloid cell-specific A$_{2A}$R disruption on the aspects of obesity-associated NAFLD and to elucidate the underlying mechanisms. Global and/or myeloid cell-specific A$_{2A}$R-disrupted mice, as well as control mice were fed a high-fat diet (HFD) to induce NAFLD. Also, bone marrow-derived macrophages and primary mouse hepatocytes were examined for inflammatory and metabolic responses. Upon feeding an HFD, both global A$_{2A}$R-disrupted mice and myeloid cell-specific A$_{2A}$R-deficient mice revealed increased severity of HFD-induced hepatic steatosis and inflammation compared with their respective control mice. In in vitro experiments, A$_{2A}$R-deficient macrophages exhibited increased proinflammatory responses, and enhanced fat deposition of wild-type primary hepatocytes in macrophage-hepatocyte co-cultures. In primary hepatocytes, A$_{2A}$R deficiency increased the proinflammatory responses and enhanced the effect of palmitate on stimulating fat deposition. Moreover, A$_{2A}$R deficiency significantly increased sterol regulatory element-binding protein 1c (SREBP1c) abundance in livers of fasted mice and in hepatocytes upon nutrient deprivation. In the absence of A$_{2A}$R, SREBP1c transcription activity was significantly increased in mouse hepatocytes. Taken together, these results demonstrate that disruption of A$_{2A}$R in both macrophage and hepatocytes accounts for increased severity of NAFLD, likely through increasing inflammation and through elevating lipogenic events due to stimulation of SREBP1c expression and transcription activity.
**Key words:** Adenosine 2A receptor, non-alcoholic fatty liver disease, obesity, inflammation, sterol regulatory element-binding protein 1c

**Abbreviations**

ACC1, acetyl-CoA carboxylase 1  
A2AR, adenosine 2A receptor  
ALD, alcoholic fatty liver disease  
AMPK, AMP-activated protein kinase  
BMDM, bone marrow-derived macrophages  
BSA, bovine serum albumin  
CD, chow diet  
ChIP, chromatin immunoprecipitation  
ChREBP, carbohydrate-responsive element-binding protein  
CPT1a, carnitine palmitoyltransferase 1a  
DMEM, Dulbecco’s modified Eagle’s medium  
FAS, fatty acid synthase  
FBS, fetal bovine serum  
GAPDH, glyceraldehyde 3-phosphate dehydrogenase  
GTT, glucose tolerance test  
H&E, hematoxylin and eosin  
HFD, high-fat diet  
LFD, low-fat diet  
IL-1β, interleukin 1β  
IL-6, interleukin 6  
ITT, insulin tolerance test  
LPS, lipopolysaccharide  
JNK, c-Jun N-terminal kinases  
MCD, methionine- and choline-deficient diet  
NAFLD, non-alcoholic fatty liver disease  
NASH, non-alcoholic steatohepatitis  
NFκB, nuclear factor kappa B  
PBS, phosphate-buffered saline  
P-Akt, phosphorylated Akt  
PP65, phosphorylated p65 subunit of NFκB  
Pp46, phosphorylated JNK1 (p46)  
RQ, respiratory quotient  
SREBP1c, sterol regulatory element-binding protein 1c  
TG, triglycerides  
TNFα, tumor necrosis factor α  
WAT, white adipose tissue
INTRODUCTION

Hepatic steatosis is a hallmark of non-alcoholic fatty liver disease (NAFLD) (1, 2). When the liver develops overt inflammatory damage, simple steatosis progresses to non-alcoholic steatohepatitis (NASH) (2, 3). The latter is the advanced form of NAFLD and is considered as a leading causal factor of cirrhosis and hepatocellular carcinoma (4, 5). In addition, NAFLD critically contributes to the development of dyslipidemia and significantly increases the incidence of atherogenic cardiovascular diseases (6), thereby serving as a key component of metabolic syndrome (7).

Numerous studies from both human subjects and rodent models demonstrate that obesity significantly increases the incidence of NAFLD (8-10). Accordingly, obesity-associated inflammation is accepted as a critical factor that initiates or exacerbates NAFLD. For instance, inflammation can cause hepatic insulin resistance, which, in turn elevates hepatic steatosis, at least in part, through increasing the expression of genes for lipogenic enzymes such as acetyl-CoA carboxylase 1 (ACC1) and fatty acid synthase (FAS), and decreasing the expression of genes for fatty acid oxidation including carnitine palmitoyltransferase 1a (CPT1a) (11-13). In addition, proinflammatory mediators, e.g., cytokines, can exert direct effects on hepatocytes to increase lipogenic events (14), and act on both hepatocytes and liver macrophages/Kupffer cells to accelerate liver inflammation. This exemplifies how inflammation serves as “a second hit” to drive the progression of simple steatosis to NASH. However, exactly how inflammation is regulated in the context of NAFLD pathophysiology remains largely unclear.
Adenosine receptors, including A1, A2A, A2B and A3, belong to the superfamily of G-protein-coupled receptors, and mediate various physiological functions of adenosine (15). Among the four adenosine receptors, A2A R displays powerful anti-inflammatory effects in immune cells such as macrophages and neutrophils (15, 16). Several studies in animal models have indicated that A2A R deficiency exacerbates concanavalin A- or endotoxin-induced liver damage, which is attributable to prolonged and enhanced expression of proinflammatory cytokines including tumor necrosis factor alpha (TNFα) and interleukin-6 (IL-6) (17). Also, there is a study showing that A2A R deficiency exacerbates the severity of aspects of alcoholic fatty liver disease (ALD) in mice (18). Additionally, A2A R activation displays beneficial effects on aspects of NASH in rodents (19, 20), which is largely mediated through the anti-inflammatory effect of A2A R. In contrast, treatment of mice with an A2A R antagonist increased the severity of CCl4-induced liver fibrosis; although the same treatment reversed the effect of ethanol on exacerbating CCl4-induced liver fibrosis (21). Collectively, these findings suggest that A2A R has a protective role in liver damage. However, whether and how A2A R coordinates hepatocyte and macrophage metabolic and/or inflammatory responses to alter NAFLD development and progression remains poorly understood. It is also not clear whether and how A2A R regulates hepatocyte lipogenic events, whose increase is of particular importance in the pathogenesis of hepatic steatosis. The present study aimed to examine how global and/or myeloid cell-specific A2A R disruption influences NAFLD aspects in mice and provides the primary evidence to support a protective role for the A2A R in both macrophages and hepatocytes in the pathogenesis of obesity-associated NAFLD. In addition, A2A R has a previously unidentified role in repressing sterol regulatory element-binding protein 1c (SREBP1c), which contributes to the anti-steatotic effect of A2A R.
MATERIALS AND METHODS

Animal experiments

Wild-type (WT) C57BL/6J were obtained from Jackson Laboratory (Bar Harbor, ME). A2AR+/−, A2AR+/−, and A2AR+/+ mice were generated as described (22). Myeloid cell-specific A2AR-disrupted mice were generated using the Cre-LoxP strategy. Briefly, homozygous myeloid cell-specific A2AR-disrupted (LysMCre+A2AR+/F/F) mice, heterozygous A2AR-disrupted (LysMCre+A2AR+/+/F/F) mice, and LysM control (LysMCre+A2AR+/+/F/F) mice were used. All mice were maintained on a 12:12-h light-dark cycle (lights on at 06:00) and subjected to studies involving chow-diet, high-fat diet (HFD, 60% fat calories) and low-fat diet (LFD, 10% fat calories) as detailed in Supplementary Information (SI). All study protocols were reviewed and approved by the Institutional Animal Care and Use Committee of Texas A&M University.

Cell culture and treatment

Bone marrow cells were prepared from A2AR-disrupted mice and WT mice, as well as LysMCre+A2AR+/F/F mice and LysMCre+A2AR+/+/F/F mice, and differentiated into macrophages (BMDM) and examined for the proinflammatory activation as described (23). Primary hepatocytes were isolated from free-fed mice (24, 25) and examined for metabolic and inflammatory responses. For macrophage-hepatocyte co-culture study, bone marrow cells were prepared from A2AR-disrupted mice and WT mice at 6 days prior to hepatocyte isolation. After differentiation, BMDM were trypsinized and added to WT primary mouse hepatocytes at a ratio of 1:10 based on the published method (12). Some hepatocytes were incubated in the absence of macrophages and served as the control. After incubation for 48 hr, the co-cultures and control
hepatocytes were subjected to the selected assays detailed in SI. Additional experiments involving hepatocytes were also detailed in SI.

**Histological, biochemical and molecular assays**

Liver sections were subjected to histological and immunohistochemical assays. Plasma parameters were measured using metabolic assay kits and ELISA kits. Also, tissue and/or cell samples were subjected to selected assays including Western blots analysis, real-time PCR, the reporter assays, and chromatin immunoprecipitation (ChIP) assay. Details are provided in SI.

**Statistical Methods**

Numeric data are presented as means ± SE (standard error). Statistical significance was assessed by unpaired, two-tailed ANOVA or Student’s *t* tests. Differences were considered significant at the two-tailed *P* < 0.05.

**RESULTS**

**HFD feeding increases liver A2AR abundance and the proinflammatory status**

Tissue distribution results revealed that the liver is one of the organs in which A2AR abundance was at the highest levels (Figure 1A). Next, we examined the pathophysiological relevance of hepatic expression of A2AR in C57BL/6J mice, which display obesity-associated NAFLD upon feeding HFD (9, 10). HFD-fed C57BL/6J mice displayed a significant increase in liver A2AR mRNAs compared to LFD-fed mice (Figure 1B). HFD-fed mice also displayed significant increases in liver protein levels of A2AR, as well as CD39 and CD73 (ectoenzymes responsible
for extracellular adenosine production) (Figure 1C). By immunohistochemistry in liver sections, HFD-fed mice revealed increased A₂AR expression in liver cells including hepatocytes compared with LFD-fed mice (Supplemental Figure S1). Additionally, HFD-fed C57BL/6J mice displayed significant increases in liver phosphorylation states of Jun N-terminal kinase (JNK) p46 and nuclear factor kappa B (NFκB) p65 as well as mRNA and protein levels of TNFα, interleukin 1 beta (IL-1β), and/or IL-6 compared with LFD-fed mice (Figure 1D,E). These results suggest an association between A₂AR amount and diet-induced liver inflammation.

A₂AR disruption exacerbates HFD-induced hepatic steatosis and inflammation

The role of A₂AR in obesity-associated NAFLD remains largely unknown, and was examined using both male and female A₂AR-disrupted (A₂AR⁻/⁻ and/or A₂AR⁺/+ ) mice, and their WT (A₂AR⁺/+ ) littermates (Supplemental Figure S2A) upon HFD feeding for 12 weeks. After the feeding period, hepatic A₂AR expression was examined and verified for A₂AR disruption (Supplemental Figure S2B). Regardless of the gender, A₂AR⁻/⁻ mice displayed much more severe phenotype of insulin resistance and glucose intolerance compared with A₂AR⁺/+ mice and/or A₂AR⁺/± mice; although A₂AR disruption exacerbated HFD-induced obesity in male, but not female mice (Supplemental Figures S3-S6). Next, we examined the effect of A₂AR disruption on NAFLD pathology. Among male mice, HFD-fed A₂AR-disrupted mice revealed a greater increase in liver weight compared with control mice and this increase was gene-dose-dependent (Figure 2A). Consistently, the severity of HFD-induced hepatic steatosis in A₂AR⁻/⁻ mice or A₂AR⁺/+ mice was greater than that in A₂AR⁺/+ mice. Also, HFD-induced hepatic steatosis in A₂AR⁺/+ mice was much pronounced compared to A₂AR⁺/+ mice as demonstrated by H&E or Oil Red O staining in liver sections (Figure 2B). Additionally, both plasma and hepatic levels of
triglycerides in HFD-fed $A_2\alpha R^{-/-}$ mice were greater than their respective levels in HFD-fed $A_2\alpha R^{+/+}$ mice and/or $A_2\alpha R^{+/+}$ mice (Supplemental Figure S7A,B). Among female mice, $A_2\alpha R$ disruption caused a slight increase in the severity of HFD-induced hepatic steatosis without significantly altering liver weight (Supplemental Figure S7C).

Because HFD-induced phenotype in male mice was more pronounced, we examined other liver phenotypes, i.e., inflammation and insulin sensitivity, mainly in male mice. Upon HFD feeding, $A_2\alpha R$-disrupted and control mice displayed comparable numbers of macrophages/Kupffer cells in the liver (Figure 2B). However, either HFD-fed $A_2\alpha R^{-/-}$ mice or HFD-fed $A_2\alpha R^{+/+}$ mice revealed a significant increase in liver proinflammatory signaling through JNK p64 and/or NFκB p65 (Figure 2C). The same trends were also observed in TNFα and IL-1β mRNAs (Figure 2D). Since increased lipogenesis is key to the development of hepatic steatosis, we examined the mRNA levels of ACC1 and FAS, which are key enzymes that stimulate liver lipogenesis by converting acetyl-CoA to malonyl-CoA and synthesizing long-chain fatty acids (26, 27). Compared with those in livers of HFD-fed $A_2\alpha R^{+/+}$ mice, the mRNA levels of ACC1 and FAS in either HFD-fed $A_2\alpha R^{-/-}$ mice or HFD-fed $A_2\alpha R^{+/+}$ mice were significantly elevated, and these elevations were $A_2\alpha R$ gene-dependent (Figure 2D). Additionally, hepatic mRNA levels of SREBP1c in HFD-fed $A_2\alpha R$-disrupted mice were significantly elevated in an $A_2\alpha R$ gene-dependent manner (Figure 2D). We also measured hepatic mRNA levels of CPT1a, a master regulator that transfers acyl CoA into mitochondria for oxidation (28, 29). Compared with the control, CPT1a mRNAs in HFD-fed $A_2\alpha R^{-/-}$ mice were slightly higher than in HFD-fed $A_2\alpha R^{+/+}$ mice or $A_2\alpha R^{+/+}$ mice (Figure 2D), suggesting a compensatory response to increased hepatic steatosis. Since SREBP1c stimulates lipogenic gene expression, we analyzed hepatic SREBP1c
expression. Both cytosolic and nuclear SREBP1c levels were significantly increased in HFD-fed $A_2AR^{-/-}$ mice compared to the values of HFD-fed $A_2AR^{+/+}$ mice (Figure 2E). When insulin signaling was analyzed, insulin-stimulated insulin receptor phosphorylation and Akt phosphorylation (S473) were significantly decreased in HFD-fed $A_2AR$-disrupted mice compared to $A_2AR^{+/+}$ mice; the decreases were $A_2AR$ gene-dose-dependent (Figure 2F). These results suggest that $A_2AR$ disruption exacerbates diet-induced hepatic steatosis and inflammation in male mice. Moreover, $A_2AR$ disruption enhances liver lipogenic events.

**Myeloid cell-specific $A_2AR$ disruption exacerbates aspects of NAFLD**

Macrophages critically regulate liver inflammation and hepatic steatosis. We sought to address a role for the $A_2AR$ in macrophages (myeloid cells) in the pathophysiology of NAFLD. We fed homozygous myeloid cell-specific $A_2AR$ disrupted (LysMCre$^+$-$A_2AR^{F/F}$) mice, heterozygous myeloid cell-specific $A_2AR$ disrupted (LysMCre$^+$-$A_2AR^{F/+}$) mice, and their WT (LysMCre$^+$-$A_2AR^{+/+}$) littermates (Supplemental Figure S8) an HFD and examined systemic insulin sensitivity and NAFLD aspects. Compared with HFD-fed LysMCre$^+$-$A_2AR^{+/+}$ mice, HFD-fed LysMCre$^+$-$A_2AR^{F/F}$ mice and HFD-fed LysMCre$^+$-$A_2AR^{F/+}$ mice revealed similar increases in the severity of systemic insulin resistance although all mice displayed similar body weight and consumed comparable amount of foods (Supplemental Figures S9 and S10A,B). However, liver weight as well as the severity of HFD-induced hepatic steatosis in LysMCre$^+$-$A_2AR^{F/F}$ or LysMCre$^+$-$A_2AR^{F/+}$ mice was greater compared to HFD-fed LysMCre$^+$-$A_2AR^{+/+}$ mice (Figure 3A,B).

Additionally, plasma and hepatic levels of triglycerides in HFD-fed LysMCre$^+$-$A_2AR^{F/F}$ mice were significantly greater than their respective levels in HFD-fed LysMCre$^+$-$A_2AR^{+/+}$ mice or LysMCre$^+$-$A_2AR^{F/+}$ mice (Supplemental Figure S10C,D). The severity of HFD-induced hepatic
steatosis in LysMCre\(^+\)-A\(_2\)AR\(^{F/F}\) mice was lower compared to that observed in global A\(_2\)AR-deficient (A\(_2\)AR\(^{-/-}\)) mice. When liver proinflammatory signaling was examined, the phosphorylation states of JNK p46 and/or NF\(\kappa\)B p65 and the mRNA levels of cytokines in HFD-fed LysMCre\(^+\)-A\(_2\)AR\(^{F/F}\) mice were greater than in HFD-fed HFD-fed LysMCre\(^+\)-A\(_2\)AR\(^{+/+}\) mice or LysMCre\(^+\)-A\(_2\)AR\(^{F/+}\) mice (Figure 3C, D). Consistent with increased hepatic steatosis, liver mRNA expression of ACC1 and FAS in HFD-fed LysMCre\(^+\)-A\(_2\)AR\(^{F/F}\) mice was also greater than their respective levels observed in HFD-fed LysMCre\(^+\)-A\(_2\)AR\(^{+/+}\) or LysMCre\(^+\)-A\(_2\)AR\(^{F/+}\) mice (Figure 3D). Taken together, these results suggest that A\(_2\)AR disruption in macrophages (myeloid cells) is sufficient to exacerbate diet-induced NAFLD.

**A\(_2\)AR-disrupted macrophages display increased proinflammatory responses and exacerbate hepatocyte fat deposition and cytokine expression**

Since LysMCre\(^+\)-A\(_2\)AR\(^{F/F}\) mice revealed increased severity of diet-induced NAFLD, we sought to verify a direct role played by the A\(_2\)AR in macrophages in regulating hepatocyte responses *in vitro*. Initially, we isolated bone marrow cells from A\(_2\)AR\(^{-/-}\) and/or A\(_2\)AR\(^{+/+}\) mice, differentiated the cells into macrophage (BMDM), and analyzed BMDM proinflammatory responses. Under LPS-stimulated conditions, the phosphorylation states of JNK p46 and NF\(\kappa\)B p65 as well as the secretion of TNF\(\alpha\) and IL-6 were significantly increased in A\(_2\)AR\(^{-/-}\) BMDM compared to the values observed in A\(_2\)AR\(^{+/+}\) BMDM (Figure 4A,B). Similarly, LPS-induced phosphorylation states of JNK p46 and NF\(\kappa\)B p65 in BMDM from LysMCre\(^+\)-A\(_2\)AR\(^{F/F}\) mice were significantly increased compared with those in BMDM from LysMCre\(^+\)-A\(_2\)AR\(^{+/+}\) mice (Figure 4C).
Next, we co-cultured WT primary mouse hepatocytes with A\textsubscript{2}\textalpha{}R$^{-/-}$ or A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM. Under palmitate-stimulated conditions, hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM accumulated more fat and revealed significantly higher levels of triglycerides than hepatocytes co-cultured in the absence of BMDM or in the presence of A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM (Figure 5A). When the expression of genes/enzymes related to fat metabolism was analyzed, ACC1, FAS and SREBP1c mRNAs in hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM were significantly higher than their respective levels in hepatocytes co-cultured in the absence of BMDM or in the presence of A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM (Figure 5B). In contrast, CPT1a mRNAs in hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM were significantly lower than those in hepatocytes co-cultured in the absence of BMDM or in the presence of A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM. When the expression of cytokines was analyzed, TNF$\alpha$ and IL-1$\beta$ mRNAs in hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM were significantly higher than their respective levels in hepatocytes co-cultured in the absence of BMDM or in the presence of A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM under basal conditions (Figure 5C, top panel). Strikingly, under LPS-stimulated conditions, TNF$\alpha$, IL-1$\beta$, and IL-6 mRNAs in hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{+/+}$ or A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM were markedly higher than their respective levels in hepatocytes cultured in the absence of BMDM (Figure 5C, bottom panel). Among co-cultures, TNF$\alpha$, IL-1$\beta$, and IL-6 mRNAs in hepatocytes/A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM co-cultures were also significantly higher than their respective levels in hepatocytes/A\textsubscript{2}\textalpha{}R$^{+/+}$ BMDM co-cultures. Additionally, the phosphorylation states of Akt in hepatocytes co-cultured with A\textsubscript{2}\textalpha{}R$^{-/-}$ BMDM were significantly lower than those in control hepatocytes or control co-cultures (Figure 5D). Taken together, these results suggest that the A\textsubscript{2}\textalpha{}R in macrophages protects against macrophage proinflammatory activation. The latter has detrimental effects on increasing hepatocyte fat deposition and proinflammatory responses and on impairing hepatocyte insulin sensitivity.
A2AR disruption exacerbates hepatocyte fat deposition and proinflammatory responses

A2AR appeared to play a protective role in NAFLD. Next, we examined the direct effects of A2AR disruption on hepatocyte responses. When fat deposition was analyzed, hepatocytes from A2AR−/− mice accumulated much more fat and revealed significantly higher levels of triglycerides than hepatocytes from A2AR+/− mice or A2AR+/+ mice under palmitate-stimulated conditions (Figure 6A). When proinflammatory signaling was analyzed, the phosphorylation states of JNK p46 and NFκB p65 in hepatocytes from A2AR−/− mice were significantly stronger than those in hepatocytes from A2AR+/− mice or A2AR+/+ mice under LPS-stimulated conditions (Figure 6B). When insulin sensitivity was analyzed, the states of insulin-stimulated Akt phosphorylation in hepatocytes from A2AR−/− or A2AR+/− mice were significantly lower than those in hepatocytes from A2AR+/+ mice (Figure 6C). Consistently, A2AR inhibition increased hepatocyte fat deposition and proinflammatory responses, and decreased hepatocyte insulin signaling (Supplemental Figure S11). Taken together, these results indicate a direct role of A2AR in protecting hepatocytes from palmitate-induced fat deposition and from LPS-induced proinflammatory responses.

A2AR disruption increases the expression and transcription activity of SREBP1c

A2AR disruption exacerbated hepatic steatosis and increased lipogenic events. To gain mechanistic insights for A2AR regulation of lipogenesis, we examined the effect of A2AR disruption on SREBP1c in mice and in isolated primary mouse hepatocytes. Since feeding increases the amount and/or transcription activity of SREBP1c, we examined whether A2AR disruption alters the effect of feeding on SREBP1c. Compared with fasting, feeding caused a
significant increase in the abundance of both cytosolic and nuclear SREBP1c in livers of A2A R+/+ mice, but not A2A R−/− mice. Strikingly, the abundance of either cytosolic or nuclear SREBP1c in livers of A2A R−/− mice was greater than that in livers of A2A R+/+ mice under fasted conditions, but not fed conditions (Figure 7A). Consistently, the abundance of cytosolic or nuclear SREBP1c was increased in hepatocytes from A2A R−/− mice compared with that in hepatocytes from A2A R+/+ mice under conditions mimicking fasting (Figure 7B). We then performed the reporter assay and observed that SREBP1c transcription activity in hepatocytes from A2A R−/− mice was significantly higher than that in hepatocytes from A2A R+/+ mice under basal conditions (in the absence of insulin) (Figure 7C). Similar changes were also observed in hepatocytes under insulin-stimulated conditions. However, insulin increased SREBP1c transcription activity in hepatocytes from A2A R+/+ but not A2A R−/− mice. Next, we performed the ChIP assay. The binding of SREBP1c to SRE/E (a binding element in FAS promoter region) was significantly increased in nuclear extracts of hepatocytes from A2A R−/− mice compared with that observed in hepatocytes from A2A R+/+ mice (Figure 7C). Also, the mRNAs of key target genes of SREBP1c, e.g., glucokinase and FAS, were greater in hepatocytes from A2A R−/− mice compared to those of hepatocytes from A2A R+/+ mice (Figure 7D). These results suggest that A2A R is capable of repressing SREBP1c expression under fasted conditions and suppressing SREBP1c transcription activity.

We also examined the phosphorylation status of AMP-activated protein kinase (AMPK), a regulator that exerts a powerful anti-steatotic effect. Compared with those in HFD-fed A2A R+/+ mice or A2A R−/− mice, hepatic phosphorylation states of AMPK in HFD-fed A2A R−/− mice were significantly decreased (Figure 7E). Since cellular ADP: ATP ratio critically influences AMPK phosphorylation, we performed cellular assays and observed that the ADP: ATP ratios were
significantly lower in hepatocytes from A2AR−/− mice than those of hepatocytes from A2AR+/+ mice or A2AR+/− mice (Figure 7F). These results were consistent with changes in the status of hepatic steatosis.

DISCUSSION

We established an association between A2AR and obesity-associated NAFLD using HFD-fed WT mice, in which increased liver proinflammatory responses were accompanied by increased liver abundance of A2AR. Considering that A2AR activation exerts powerful anti-inflammatory effects (19, 20, 30, 31) whereas A2AR deficiency exacerbates proinflammatory responses (32), we speculated that the increase in liver A2AR abundance in HFD-fed WT mice was a defensive response. As substantial evidence, global A2AR deficient mice displayed a significant increase in the severity of HFD-induced liver inflammation and other NAFLD aspects. Of importance, this evidence from A2AR deficient mice establishes a cause-and-effect relationship between A2AR disruption and NAFLD. In global A2AR-deficient mice, increased NAFLD aspects may be attributable to the lack of A2AR in macrophages, hepatocytes, or both. Considering the importance of macrophages in regulating inflammation, we speculated that the A2AR in macrophages was needed for protecting mice from obesity-associated NAFLD and we found this was the case. Therefore, our three lines of in vivo evidence makes it conceivable that A2AR acts through, at least partially, suppressing inflammation to protect against obesity-associated NAFLD.
A2AR activation was previously shown to ameliorate methionine- and choline-deficient diet (MCD)-induced NASH in rats (19). A follow-up study confirmed the beneficial effect of A2AR activation in MCD-fed mice (20), which was associated with decreased proinflammatory responses in both hepatocytes and lymphocytes. These results suggest anti-inflammation as an essential mechanism by which A2AR activation improves NASH. However, the studies by Carini et al. did not address how macrophages respond to A2AR activation in the context of protecting against NASH. Indeed, the observed decreases in cytokine production by hepatic mononuclear cells of MCD-fed mice upon A2AR activation could be secondary to the anti-inflammatory effects of A2AR activation on hepatocytes and/or macrophages. Considering that macrophages have been implicated to critically determine the development and progression of obesity-associated NAFLD (11, 12), it is necessary to address how the A2AR in macrophages alters NAFLD aspects. In the present study, we confirmed that A2AR disruption exacerbates macrophage proinflammatory activation. Moreover, we revealed, for the first time, that myeloid cell-specific A2AR deficiency exacerbated HFD-induced hepatic steatosis and inflammation. Because A2AR was disrupted only in myeloid cells, increased hepatic steatosis and inflammation in HFD-fed LysMCre+\( \rightarrow \)A2AR\(^{−/−}\) mice appeared to be secondary to the consequences of A2AR disruption in myeloid cells. As supporting evidence, hepatocytes co-cultured with A2AR-disrupted macrophages revealed significant increases in the expression of proinflammatory cytokines and in palmitate-induced fat deposition and in the expression of ACC1, FAS, and SREBP1c compared with hepatocytes co-cultured with control macrophages. Because the initial differences among co-cultures existed solely in macrophages (e.g., the presence or absence of A2AR), the outcomes of hepatocyte lipogenic events were attributable to the effects secondary to A2AR disruption-associated increase in macrophage proinflammatory activation. The latter has
been previously shown to generate macrophage-derived factors, i.e., proinflammatory cytokines, to promote hepatocyte lipogenic events (14), enhance hepatocyte proinflammatory responses, and decrease hepatocyte insulin signaling (33). Therefore, the results from myeloid cell-specific A2AR deficient mice and macrophage-hepatocyte co-cultures demonstrated that A2AR has a role in coordinating macrophage actions on hepatocytes to protect against NAFLD aspects.

A2AR also critically regulates hepatocyte metabolic and inflammatory responses. In support of this, A2AR disruption brought about deleterious effects on palmitate-induced fat deposition and on LPS-induced proinflammatory responses in primary mouse hepatocytes. In addition, treatment of WT mouse hepatocytes with an A2AR antagonist generated effects mimicking A2AR disruption. These two lines of evidence not only recapitulated the findings in HFD-fed A2AR-disrupted mice, but also served as complementary evidence to support the anti-inflammatory and anti-steatotic effects of A2AR activation (19, 20). However, it remains to be explored, within hepatocytes, whether and how the anti-inflammatory effect of A2AR interplays with the anti-lipogenic effect of A2AR. Given the co-existence of these effects of A2AR, it is possible that A2AR acts through suppressing hepatocyte inflammatory signaling to inhibit lipogenic events. Alternatively, A2AR may act through two parallel pathways to suppress hepatocyte proinflammatory responses and to inhibit hepatocyte fat deposition.

It is a significant finding that A2AR disruption elevated hepatic SREBP1c abundancy. Additionally, increased liver abundancy of SREBP1c in HFD-fed A2AR-deficient mice was accompanied with increases in hepatic expression of lipogenic enzymes and in the severity of hepatic steatosis. These results are consistent with an established paradigm concerning the
critical contribution of lipogenesis, at increased levels, to the development of hepatic steatosis (34-36). Physiologically, hepatic SREBP1c is stimulated by feeding and repressed by fasting (36), which accounts for increased hepatic fat content under fed states and decreased hepatic fat content under fasted states, respectively. Strikingly and interestingly, in the present study, A₂AᵡR deficiency increased liver SREBP1c abundance under fasted conditions, where SREBP1c expression and activity should be suppressed physiologically. Consistently, A₂AᵡR disruption increased the abundance of SREBP1c in hepatocytes upon nutrient deprivation. These findings support a repressive effect A₂AᵡR on SREBP1c. Additional to altering SREBP1c abundance, A₂AᵡR disruption also decreased hepatic AMPK phosphorylation states, likely through a mechanism involving decreases in hepatocyte ADP: ATP ratio because AMPK is activated by increased ratios of AMP: ATP and ADP: ATP. Considering the role of AMPK in phosphorylating and inactivating SREBP1c (37), A₂AᵡR deficiency-associated decreases in AMPK phosphorylation appeared to also contribute to the effects of A₂AᵡR deficiency on increasing hepatic lipogenesis and on exacerbating hepatic steatosis. Indeed, at the cellular level, A₂AᵡR disruption increased SREBP1c transcription activity and enhanced the binding of SREBP1c to the promoter region of FAS. Collectively, our findings suggest that A₂AᵡR signaling pathways are involved in repression of SREBP1c, which in turn contributes to the effects of A₂AᵡR on suppressing lipogenic events, and on protecting against the development of hepatic steatosis.

In summary, we validated a protective role for A₂AᵡR in obesity-associated NAFLD. Mechanistically, this role of A₂AᵡR is attributable to the direct effects of A₂AᵡR on altering inflammatory and metabolic responses of macrophages and hepatocytes. Specifically, A₂AᵡR
suppresses macrophage and hepatocyte proinflammatory responses. A<sub>2A</sub>R suppression of macrophage activation also generates secondary effects on inhibiting the inflammatory responses and lipogenic events of hepatocytes. Lastly, A<sub>2A</sub>R can directly suppress hepatocyte fat deposition, which is attributable to the effects of A<sub>2A</sub>R on repressing SREBP1c expression under fasted states and on suppressing SREBP1c transcription activity. Therefore, targeting A<sub>2A</sub>R to suppress inflammation and lipogenesis is a viable therapeutic strategy for the treatment of NAFLD and inflammation-associated human liver diseases.

**FUNDING**

This work was supported in whole or in part by grants from the American Diabetes Association (1-17-IBS-145 to C.W.) and the National Institutes of Health (DK095862 to C.W. and HL095556 to Y.H). This work was also supported in part by the Dr. Nicholas C. Hightower Centennial Chair of Gastroenterology from Scott & White, a VA Research Career Scientist Award to G.A. and the NIH grants DK062975 and DK054811 to Drs. G.A, F.M. and S.G. Also, C.W. is supported by the Hatch Program of the National Institutes of Food and Agriculture (NIFA). Y.C. is supported by China Scholarship Council.

**AUTHOR CONTRIBUTION**

CONFLICT OF INTEREST

This material, in part, is the result of work supported with resources and the use of facilities at the Central Texas Veterans Health Care System, Temple, Texas. The content is the responsibility of the author(s) alone and does not necessarily reflect the views or policies of the Department of Veterans Affairs or the United States Government.

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FIGURE LEGENDS

Figure 1. HFD feeding increases hepatic A2A R abundance and enhances liver proinflammatory responses
Wild type C57BL/6J mice were fed as described in Methods. (A) Tissue distribution of A2AR. SI, small intestine; WAT, white adipose tissue. (B) Liver A2AR mRNA levels. AU, arbitrary unit. (C) Liver amount of A2AR, CD39 (ecto-nucleoside triphosphate diphosphohydrolase 1), and CD73 (ecto-5'-nucleotidase). (D) Liver proinflammatory signaling. (E) Liver cytokine mRNA (left panel) and protein (right two panels) levels. For A, C, and D, liver lysates were used for Western blot analysis. Blots were quantified using densitometry (bar graphs in C and D). For B and E, liver mRNAs were quantified using real-time RT-PCR. For E, liver protein levels of cytokines were quantified using ELISA kits. For all bar graphs, data are means ± SE. n = 6 - 8. Statistical difference between HFD and LFD: *, P < 0.05 and **, P < 0.01 in B and E (right two panels) or in bar graphs of C, D, and E (left two panels) for the same protein or gene.

Figure 2. A2AR disruption exacerbates HFD-induced hepatic steatosis and inflammation

Male A2AR+/− mice, A2AR+/− mice, and A2AR+/+ mice, at 5 - 6 weeks of age, were fed an HFD for 12 weeks. Age- and gender-matched A2AR+/+ mice were fed an LFD for 12 weeks and served as normal control. (A) Liver weight. (B) Representative images of H&E, Oil-Red-O, and/or F4/80 staining for liver sections. (C) Liver proinflammatory signaling. (D) Liver mRNA levels. (E) Liver SREBP1c abundance. pSREBP1c, precursor SREBP1c; nSREBP1c, nuclear SREBP1c. (F) Liver insulin signaling. Prior to tissue harvest, HFD-fed mice were injected with or without insulin (1 U/kg) into the portal vein for 5 min. IR, insulin receptor. For C, E, and F, liver lysates (C and F), as well as liver cytosolic and nuclear fractions (E) were subjected to Western blot analysis. Bar graphs, quantification of blots. For A - F, numeric data are means ± SE. n = 10 - 12 (A) or n = 8 - 10 (C - F). Statistical difference between A2AR+/−-HFD and A2AR+/+-HFD: *, P < 0.05 and **, P < 0.01 in bar graphs of A, C, and E, in D within the same gene, or in bar graphs of...
F under insulin-stimulated condition; statistical difference between $A_2AR^{-/-}$-HFD and $A_2AR^{+/+}$-HFD: †, $P < 0.05$ and ††, $P < 0.01$ in bar graphs of A, C, and E, in D within the same gene, or in bar graphs of F under insulin-stimulated condition; statistical difference between $A_2AR^{-/-}$-HFD and $A_2AR^{+/+}$-HFD: †, $P < 0.05$ and ††, $P < 0.01$ in bar graphs of A and C, in D within the same gene, or in bar graphs of F under insulin-stimulated condition.

Figure 3. Myeloid cell-specific $A_2AR$ disruption exacerbates HFD-induced hepatic steatosis and proinflammatory responses

Male LysMCre$^+$-A$_2$AR$^{F/F}$ mice, LysMCre$^+$-A$_2$AR$^{F/+}$ mice, and LysMCre$^+$-A$_2$AR$^{+/+}$ mice, at 5 - 6 weeks of age, were fed an HFD for 12 weeks. (A) Liver weight. (B) Representative images of H&E, Oil-Red-O, and/or F4/80 staining for liver sections. (C) Liver lysates were examined for the proinflammatory signaling using Western blot analysis. Bar graphs, quantification of blots. AU, arbitrary unit. TLR4, Toll-like receptor 4. (D) Liver mRNA levels were quantified using real-time RT-PCR. For A, C, and D, numeric data are means ± SE. n = 8 - 10. Statistical difference between LysMCre$^+$-A$_2$AR$^{F/F}$ and LysMCre$^+$-A$_2$AR$^{F/+}$: *, $P < 0.05$ and **, $P < 0.01$ in A or in bar graphs of C and D within the same protein or gene; statistical difference between LysMCre$^+$-A$_2$AR$^{F/F}$ and LysMCre$^+$-A$_2$AR$^{F/+}$: †, $P < 0.05$ and ††, $P < 0.01$ in bar graphs of C and D within the same protein or gene; statistical difference between LysMCre$^+$-A$_2$AR$^{F/+}$ and LysMCre$^+$-A$_2$AR$^{+/+}$: †, $P < 0.05$ and ††, $P < 0.01$ in A or in bar graphs of C and D within the same protein or gene.

Figure 4. $A_2AR$ disruption aggravates macrophage proinflammatory activation.
Bone marrow-derived macrophages (BMDM) were prepared as described in Methods. (**A, B**) Proinflammatory signaling (A) and cytokine production (B) of BMDM from A<sub>2A</sub>R<sup>−/−</sup> mice and A<sub>2A</sub>R<sup>+/+</sup> mice. For A, BMDM were treated with or without lipopolysaccharide (LPS, 100 ng/ml) for 30 min prior to harvest. Cell lysates were subjected to Western blot analysis. Bar graphs, quantification of blots. For B, cytokine concentrations in BMDM conditioned-media. (**C**) Proinflammatory signaling of BMDM from LysMCre<sup>+</sup>-A<sub>2A</sub>R<sup>F/F</sup> mice and LysMCre<sup>+</sup>-A<sub>2A</sub>R<sup>+/+</sup> mice. Cells were treated and analyzed as described in A. Bar graphs, quantification of blots. For A - C, numeric data are means ± SE. n = 4 - 6. Statistical difference between A<sub>2A</sub>R<sup>−/−</sup> and A<sub>2A</sub>R<sup>+/+</sup>: *, P < 0.05 and **, P < 0.01 in bar graphs of A (under LPS-stimulated condition) or in B; statistical difference between LysMCre<sup>+</sup>-A<sub>2A</sub>R<sup>F/F</sup> and LysMCre<sup>+</sup>-A<sub>2A</sub>R<sup>+/+</sup>: †, P < 0.05 and ††, P < 0.01 in bar graphs of C (under LPS-stimulated condition).

**Figure 5.** A<sub>2A</sub>R disruption exacerbates the effects of macrophages on increasing hepatocyte fat deposition, cytokine expression, and insulin resistance

Macrophage-hepatocyte co-cultures were performed as described in Methods. A set of primary hepatocytes were incubated without macrophages and followed by the same treatments as co-cultures. (**A**) Hepatocyte fat deposition. The cells were treated with palmitate (Pal, 250 µM, conjugated in bovine serum albumin (BSA)) or BSA for the last 24 hr of the 48 hr incubation period, and stained with Oil Red O for 1 hr. Bar graphs, quantification of fat content and triglyceride levels. (**B, C**) The mRNA levels of genes related to fat metabolism (B) and proinflammatory cytokines (C) were examined using real-time RT-PCR. Prior to harvest, cells were treated without or with LPS (20 ng/ml) for 6 hr. (**D**) Insulin signaling. Prior to harvest, cells were treated with insulin (100 nM) or PBS for 30 min. Cell lysates were subjected to Western
blot analysis. Bar graph, quantification of blots. For A - D, numeric data are means ± SE, n = 4 - 6. Statistical difference between co-cultures with $A_2A R^{-/-}$ BMDM and co-cultures with $A_2A R^{+/+}$ BMDM: *, $P < 0.05$ and **, $P < 0.01$ in A, in B and C within the same gene, or in bar graph of D under insulin-stimulated condition; statistical difference between co-cultures with $A_2A R^{-/-}$ BMDM and hepatocytes cultured without BMDM (None): †, $P < 0.05$ and ††, $P < 0.01$ in A, in B and C within the same gene, or in bar graph of D under insulin-stimulated condition; statistical difference between co-cultures with $A_2A R^{+/+}$ BMDM and hepatocytes/None: ‡, $P < 0.05$ in bar graph of D under insulin-stimulated condition.

Figure 6. $A_2A R$ disruption exacerbates hepatocyte fat deposition and proinflammatory responses and impairs hepatocyte insulin sensitivity

Primary hepatocytes were isolated from chow-diet-fed male $A_2A R^{-/-}$ mice, $A_2A R^{+/+}$ mice, and $A_2A R^{+/+}$ mice, at 10 - 12 weeks of age. (A) Hepatocyte fat deposition. After attachment, hepatocytes were treated with palmitate (Pal, 250 µM) or BSA for 24 hr, and stained with Oil Red O for 1 hr. Bar graphs, quantification of fat content and triglyceride levels. (B, C) Hepatocyte proinflammatory (B) and insulin (C) signaling. Prior to harvest, cells were treated with LPS (100 ng/ml), insulin (100 nM), or PBS for 30 min. Cell lysates were subjected to Western blot analysis. For A - C, numeric data are means ± SE, n = 4 - 6. Statistical difference between $A_2A R^{-/-}$ vs. $A_2A R^{+/+}$: *, $P < 0.05$ and **, $P < 0.01$ in A, in B under LPS-stimulated condition, or in C under insulin-stimulated condition; statistical difference between $A_2A R^{-/-}$ and $A_2A R^{+/+}$: †, $P < 0.05$ in A or in B under LPS-stimulated condition; statistical difference between $A_2A R^{+/+}$ and $A_2A R^{+/+}$: ‡‡, $P < 0.01$ in C under insulin-stimulated condition.
Figure 7. A<sub>2A</sub>R deficiency elevates hepatic abundance of SREBP1c under fasting/nutrient deprivation and enhances SREBP1c transcription activity

(A) Liver SREBP1c abundance. Male A<sub>2A</sub>R<sup>−/−</sup> mice and A<sub>2A</sub>R<sup>+/+</sup> mice, at 10 - 12 weeks of age, were free-fed or fasted for 18 hr. (B) Hepatocyte SREBP1c abundance. Primary hepatocytes were incubated in M199 in the absence of fetal bovine serum for 24 hr. For A and B, liver (A) or hepatocyte (B) cytosolic and nuclear proteins were subjected to Western blot analysis. (C) Hepatocyte SREBP1c transcription activity. Left panel, primary hepatocytes were incubated in M199 in the absence of fetal bovine serum and transfected with a reporter construct in which luciferase expression is under the control of SRE sequences on fatty acid synthase (pFAS-SRE-luc) or a control (pGL3-luc) for 24 hr; right panel, primary hepatocytes were treated as described in B. Hepatocyte chromatin was immunoprecipitated with antibodies against SREBP1c. The resultant DNA were analyzed for SRE/E sequences of FAS promoter. (D) Hepatocyte mRNAs. Cells were treated as described in B. (E) Liver AMPK phosphorylation states. Male A<sub>2A</sub>R<sup>−/−</sup> mice, A<sub>2A</sub>R<sup>−/+</sup> mice, and A<sub>2A</sub>R<sup>+/+</sup> mice were fed as described in Figure 2. (F) Hepatocyte ADP/ATP ratio. Cells were treated as described in B. For B - D and F, primary hepatocytes were isolated from chow-diet-fed male A<sub>2A</sub>R<sup>−/−</sup> mice and A<sub>2A</sub>R<sup>+/+</sup> mice, at 10 - 12 weeks of age. For A, B, and E, blots were quantified using densitometry. For all bar graphs, data are means ± SE, n = 4 - 6 (in A - C) or n = 6 - 8 (in D - F). Statistical difference between A<sub>2A</sub>R<sup>−/−</sup> and A<sub>2A</sub>R<sup>+/+</sup>: *, P < 0.05 and **, P < 0.01 in A and B for the same fraction, in C for the same construct under the same condition, in D for the same gene, or in E and F; statistical difference between insulin-treated A<sub>2A</sub>R<sup>−/−</sup> and A<sub>2A</sub>R<sup>+/+</sup>; †, P < 0.05 in E and F; statistical difference between insulin-treated A<sub>2A</sub>R<sup>−/−</sup> and A<sub>2A</sub>R<sup>−/+</sup>; ‡, P < 0.05 in E and F; statistical difference between insulin-treated A<sub>2A</sub>R<sup>−/−</sup> and A<sub>2A</sub>R<sup>−/+</sup> cells and
control-treated $A_2AR^{+/+}$ cells (transfected with pFAS-SRE-luc): †, $P < 0.05$ in C; statistical
difference between $A_2AR^{+/–}$ and $A_2AR^{+/+}$: ‡‡, $P < 0.01$ in F.
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