Weight loss and lipolysis promote a dynamic immune response in murine adipose tissue

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Obesity elicits an immune response characterized by myeloid cell recruitment to key metabolic organs, including adipose tissue. However, the response of immune cells to nonpathologic metabolic stimuli has been less well studied, and the factors that regulate the metabolic-dependent accumulation of immune cells are incompletely understood. Here we characterized the response of adipose tissue macrophages (ATMs) to weight loss and fasting in mice and identified a role for lipolysis in ATM recruitment and accumulation. We found that the immune response to weight loss was dynamic; caloric restriction of high-fat diet–fed mice led to an initial increase in ATM recruitment, whereas ATM content decreased following an extended period of weight loss. The peak in ATM number coincided with the peak in the circulating concentrations of FFA and adipose tissue lipolysis, suggesting that lipolysis drives ATM accumulation. Indeed, fasting or pharmacologically induced lipolysis rapidly increased ATM accumulation, adipose tissue chemoattractant activity, and lipid uptake by ATMs. Conversely, dietary and genetic manipulations that reduced lipolysis decreased ATM accumulation. Depletion of macrophages in adipose tissue cultures increased expression of adipose triglyceride lipase and genes regulated by FFA, and increased lipolysis. These data suggest that local lipid fluxes are central regulators of ATM recruitment and that once recruited, ATMs form lipid-laden macrophages that can buffer local increases in lipid concentration.

Introduction

Obesity activates a complex immune program in which differentiation, activation, and recruitment of lymphoid and myeloid cells to key metabolic tissues are central features (1–11). In mammals, expansion of adipose tissue mass induces accumulation of adipose tissue macrophages (ATMs), which produce proinflammatory molecules, including TNF-α, SAA3, and CCL2 (MCP-1) (2, 12). ATMs contribute to both local and systemic inflammation and modulate metabolic phenotypes, including insulin resistance (13). Genetic and pharmacologic manipulations that reduce ATM content or alter their inflammatory state in obese rodents modulate local inflammation and are associated with reduced insulin resistance (14). For example, Ccr2 deficiency or antagonism reduces ATM recruitment and partially protects mice from obesity-induced insulin resistance (15). Similarly, myeloid cell–specific deletion of inflammatory pathway–regulating IKK-β reduces obesity-induced inflammation and diet-induced insulin resistance (6, 16).

The metabolic factors that regulate the immune response to obesity and the accumulation of macrophages and other immune cells in adipose tissue remain poorly defined. In weight-stable individuals and rodents or those gaining weight, there is a strong, positive correlation between adipocyte size and ATM content (2, 17, 18). This correlation suggests that macrophage accumulation occurs in response to a process associated with increased adipocyte volume. Cinti and colleagues have suggested that adipocyte necrotic-like death, which they hypothesize is driven by hypertrophy and accelerated by obesity, is the primary stimulus that regulates ATM accumulation (19). Indeed, massive adipocyte apoptosis in a transgenic model of inducible lipoatrophy leads to rapid accumulation of ATMs (20) and suggests that adipocyte death can drive ATM accumulation. However, recent studies find that the rate of adipocyte death is not increased in obese individuals (21). Another compelling hypothesis implicates hypoxia in ATM recruitment. In this model, adipocyte hypertrophy creates areas of microhypoxia that activate inflammatory programs important in the remodeling of vasculature. These pathways include JNK1-regulated chemokine release (22–24). However, in a recent study, adipose tissue expression of Hif1α, the primary factor mediating the hypoxia response, and of Vegfa, a key downstream Hifα target, were negatively correlated with ATM content (17).

Increases in adipose tissue mass and adipocyte volume have other broad metabolic consequences including reduced mitochondrial function, increased ER stress, impaired insulin signaling, and higher rates of basal lipolysis (25–28). A clue to the function of ATMs and their regulation may come from the observation that with increasing adiposity, ATMs form multinucleated syncytia that contain large lipid droplets (19, 29), suggesting that in obesity, ATMs phagocytose or take up excess lipid. The tight coupling of adipocyte size with macrophage accumulation and lipid uptake suggests that excess lipids may be critical for ATM accumulation. We therefore hypothesized that the immune system and macrophages respond directly to alterations in metabolic function and substrate fluxes and specifically that obesity-induced increases in basal lipolysis (by virtue of increased fat cell volume) (25) will increase local extracellular lipid concentrations and drive ATM accumulation. Consistent with this hypothesis, visceral adipose tissue depots — compared with abdominal subcutaneous depots — have increased basal rates of lipolysis and contain more ATMs (18, 30, 31). If increased lipolysis drives ATM accumulation, then altering lipolysis should alter ATM accumulation in a predictable manner. Total lipolysis is the sum of (a) basal lipolysis, which is
determined in large measure by adipocyte triglyceride content, and (b) demand lipolysis, which is the hormonally regulated release of FFA in response to nutritional demands. Obesity increases adipocyte size and therefore basal lipolysis. Negative energy balance leads to the mobilization of adipose tissue triglyceride stores and activates demand lipolysis; therefore, in obese individuals during early weight loss, when adipocyte triglyceride content remains high, both basal and demand lipolysis are high, and if our hypothesis is correct, ATM content should be elevated.

While detailed time-course studies of macrophage accumulation in adipose tissue during weight gain have been performed, the kinetics of ATM accumulation during weight loss remain poorly defined. A steady increase in weight gain and adipose tissue mass lead to a proportional increase in ATM content (2, 5, 18). After sustained...
Weight loss induces a transient accumulation of ATMs. To understand how the immune system — and specifically ATMs — responds to weight loss, we characterized the metabolic and inflammatory phenotypes of obese mice that were subjected to moderate caloric restriction. Nine-week-old male C57BL/6J mice were fed a high-fat diet (60% of the calories derived from fat; Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI42845DS1) until they reached 40 grams of body mass. These animals were then subjected to caloric restriction (they received 70% of their ad libitum food intake) to induce gradual weight loss. Mice were sacrificed and tissues collected after 0, 3, 7, 14, 21, 42, and 60 days of caloric restriction. A group of age-matched lean-chow diet–fed mice was studied as a control (Supplemental Figure 1A). At the start of caloric restriction, all high-fat diet–fed mice had similar body composition, fasting blood glucose, and serum insulin concentrations (Supplemental Table 2). We rotated the assignment of mice to different caloric restriction groups, ensuring that the mean age of each group at sacrifice was not significantly different (Supplemental Figure 2A). As expected, this protocol of moderate caloric restriction induced a gradual reduction in weight and fat mass without affecting lean mass significantly (Supplemental Figure 1B, C and D, and Supplemental Figure 2B). Total fat mass, as measured by NMR spectroscopy, was measurably reduced by day 14. Mice continued losing weight throughout the intervention and at the end of the caloric restriction they had lost 27.9% of their initial body mass (40.8 ± 0.3 g vs. 29.4 ± 3.8 g; P < 0.001). All adipose depots decreased in mass at a similar rate (Supplemental Figure 1D), and the decrease in fat mass was due to a decrease in adipocyte size and not to a decrease in adipocyte number (Supplemental Figure 1, C–F). Decreases in

Results

Weight loss induces a transient accumulation of ATMs. To understand how the immune system — and specifically ATMs — responds to weight loss, we determined directly whether fasting and pharmacologically induced adipose tissue lipolysis affects ATM accumulation in adipose tissue, we measured ATM content in high-fat–fed obese mice placed on moderate caloric restriction and monitored the kinetics of macrophage accumulation over a 2-month period. We determined directly whether fasting and pharmacologically induced adipose tissue lipolysis increases ATM accumulation. Conversely, we reduced the rate of adipose tissue lipolysis by isocaloric substitution of carbohydrate for dietary fat and studied the effects of fasting in Atgl/Pnpla2–deficient mice to assess whether reduced lipolysis decreases ATM accumulation. We studied the effects of inducing lipolysis on ATMs in vivo and in intact adipose tissue in vitro to determine whether lipolysis per se activates FFA uptake by ATMs. Finally, we depleted macrophages from adipose tissue in vitro and found that in the absence of ATMs, adipose tissue expression of Atgl/Pnpla2 lipase and lipolysis are increased. Our results support a model in which adipose tissue lipolysis drives ATM accumulation and recruited macrophages buffer local increases in lipid concentration through phagocytosis and perhaps through modulation of adipocyte metabolism.

Figure 2

Measures of lipolysis correlate with ATM content. (A) Serum concentrations of FFA during weight loss induced by caloric restriction. Black bars represent high-fat diet–induced obese mice that underwent caloric restriction for different time intervals. White bar represents lean mice that were fed a chow diet and did not undergo caloric restriction. n = 5–6 mice/group. (B) Correlation of macrophage content (% macrophages) and serum FFA concentration in mice during weight loss; the square value of the Pearson’s correlation coefficient is shown. n = 5–6 mice/group. Each data point represents the % of macrophages in murine perigonadal adipose tissue at different serum FFA concentrations. (C) And serum FFA concentration in mice during weight loss; the square value of the Pearson’s correlation coefficient is shown. n = 5–6 mice/group. Each data point represents the % of macrophages in murine perigonadal adipose tissue at different serum FFA concentrations. (D) FFA release from explants of perigonadal adipose tissue incubated under basal conditions. Explants were isolated from high-fat diet–induced obese mice that were ad libitum fed or underwent caloric restriction for 3 or 42 days. (E) Glycerol release from explants of perigonadal adipose tissue incubated under basal conditions. Explants were isolated from high-fat diet–induced obese mice that were ad libitum fed or underwent caloric restriction for 3 or 42 days. All data are represented as mean ± SD. *P < 0.05, versus day 0.
adipocyte size were accompanied, as expected, by a reduction in leptin gene expression and leptin serum concentration (Supplemental Figure 3, A and B).

During weight gain and in cross-sectional studies of weight-stable humans, there is a positive, nearly linear relationship between adiposity and markers of local and systemic inflammation (13). Following weight loss of approximately 17% in patients 3 months after bariatric surgery, ATM content and adipose tissue inflammation are reduced and insulin sensitivity improved (32, 33). However, there have been few studies of the relationship between adiposity and measures of inflammation during dynamic weight loss. A recent study by Langin and colleagues suggests that during early weight loss in humans, the expression of inflammatory genes is not decreased (34).

To provide greater temporal resolution of the relationship between adiposity and metabolic and inflammatory phenotypes during weight loss, we measured fasting blood glucose, serum insulin, and adipose tissue expression of inflammatory genes in our cohort of mice during 2 months of continuous weight loss. Fasting blood glucose and serum insulin concentrations remained elevated during the first week of weight loss but thereafter significantly decreased, concomitantly with the reduction in percentage of body fat (Supplemental Figure 4). In contrast, the reduction in inflammatory gene expression in perigonadal adipose tissue was not uniform, and several classes of genes were identified based on their expression pattern during weight loss. The expression of some inflammatory genes such as Saa3 decreased early during weight loss, preceding the improvement in glucose homeostasis, whereas the gene expression...
of other prototypical M1 inflammatory genes, including Tnf (Tnfa) remained elevated during the entire period when the animals were in negative energy balance (Supplemental Figure S5). Circulating inflammatory proteins also fell into several classes in response to negative energy balance, with concentrations of some inflammatory molecules, e.g., resistin, falling early during weight loss and before metabolic improvement, while the concentrations of others, e.g. PAI-1, remained unchanged throughout caloric restriction and weight loss (Supplemental Figure 3, C and D).

In contrast to the other classes of genes, the adipose tissue expression of macrophage/myeloid cell–specific genes Emr1 (F4/80), Cd68, and Csf1r increased after 3 days of caloric restriction (Figure 1A). By 60 days of weight loss, however, the expression of Emr1 and Cd68 was decreased to levels below those present prior to the start of weight loss. The late reduction in macrophage-specific gene expression was consistent with previous studies that had examined the effects of long-term weight loss. To determine whether the initial increase and ultimate reduction in macrophage/myeloid-specific gene expression was due to alterations of gene expression or to a change in macrophage number, we performed immunohistochemistry using an antibody that recognizes the macrophage antigen F4/80 (EMR1). Consistent with the gene expression studies, macrophage number in perigonadal adipose tissue increased during the first week of weight loss. Three days after the beginning of caloric restriction, perigonadal adipose tissue had 47% more macrophages than adipose tissue from ad libitum–fed control mice (percentage of macrophages per total cells: 38.6% ± 4.1% vs. 26.3% ± 7.4%; P < 0.01; Figure 1, B and C). We and others have previously shown that there is a strong positive correlation between adiposity and ATM content (2, 18). However, during weight loss, this relationship is lost. Instead, during an initial period of weight loss (days 0–7 in our study), there are more ATMs found in leaner mice (Figure 1D), whereas later during weight loss, the more typical positive correlation is found (Figure 1D). Identical relationships were found when ATM content was plotted versus adipocyte size (data not shown). ATMs, and specifically CD11c+ ATMs and CD11c+ crown-like structures (CLS), are most tightly associated with adipose tissue inflammation and systemic insulin resistance. Almost all CD11c+ cells in the perigonadal adipose tissue of mice during early weight loss were also F4/80+ (CD11c+ ATMs) (Supplemental Figure 6). Although the total number of ATMs increased during early weight loss, the number of CD11c+ ATMs and CLS did not increase (Supplemental Figure 7), consistent with accumulation of a CD11c+ population of ATM during early weight loss and a lack of increase in markers of inflammation or an impairment in insulin resistance during this same period (Supplemental Figures 4 and 5).

The early initial increase in ATM content was not unique to perigonadal adipose tissue. Consistent with previous findings in high-fat diet–induced obese mice, in subcutaneous adipose tissue, the ATM content was lower than in perigonadal adipose tissue (2, 32). However, after 3 days of caloric restriction, the ATM content doubled in subcutaneous depots (macrophages as a percentage of all cells: 10.0% ± 3.1% vs. 20.2% ± 7.4%; P < 0.05; Figure 1, E and F). In both perigonadal and subcutaneous depots, ATM numbers decreased progressively after 3 days of negative energy balance so that after 42 days of caloric restriction, ATM content was significantly lower than in adipose tissue from high-fat–fed mice that had never been caloricologically restricted (Figure 1, B, C, E, and F).

During weight gain, it has been suggested that ATM accumulation is driven by adipocyte necrosis, tissue remodeling, or micro-hypoxia. The initial increase in ATMs in response to weight loss...
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Ad lib fed Atgl+/+ Fasted Atgl+/+

Perigonadal adipose tissue

Ad lib fed Atgl−/− Fasted Atgl−/−

Perigonadal adipose tissue

% of macrophages

Ad lib Fasted ***

Atgl+/+ Atgl−/−

C

Emr1 Casgr1

Relative expression (AU)

Ad lib Atgl+/+ Fasted Atgl−/−

Figure 5
ATGL/PNPLA2 deficiency limits ATM accumulation during fasting. (A) Immunohistochemical staining of F4/80-expressing (EMR1) macrophages in perigonadal adipose tissue sections from Atgl+/+(top panel) and Atgl−/−(lower panel) mice that were either ad libitum fed (left) or fasted (right). Arrows indicate ATMs. Scale bars: 100 μm. (B) Macrophages as a percentage of all cells in lean ad libitum–fed Atgl+/+ and fasted Atgl−/− mice. n = 4–5 mice/group. ***P < 0.001, versus ad libitum fed. (C) Expression of macrophage-specific genes in perigonadal adipose tissue of lean ad libitum–fed and fasted Atgl+/+ mice. n = 4–5 mice/group. All data are represented as mean ± SD.

was not associated with a reduction in adipocyte number, as might have been expected, if adipocyte necrosis was driving the accumulation of ATMs (Supplemental Figure 1F). Nor did the peak in ATMs coincide with an upregulation of a transcriptional program of adipose tissue remodeling (Supplemental Figure 8).

Measures of lipolysis correlate with ATM content. The initial increase in ATMs was associated with changes in circulating FFA concentrations and measures of lipolysis. Basal lipolysis in adipose tissue is the release of FFA from adipocytes, which occurs in the absence of negative energy balance. Basal lipolysis is increased in adipose tissue of obese individuals and correlates positively with adipocyte size (25). Demand lipolysis is the hormonally and autonomically driven release of FFA from adipocyte triglycerides that is activated by negative energy balance, when FFA are mobilized from adipose tissue for systemic use as substrates (25). A model in which lipolysis regulates ATM accumulation is consistent with previous observations of weight-stable or weight-gaining individuals: obese mice with large adipocytes have higher basal adipose tissue lipolysis and greater ATM content than lean animals. During early weight loss when adipocyte size has not changed significantly, basal lipolysis remains high and demand lipolysis increases, and thus we predict there is a net increase in total lipolysis. But as adipose tissue mass and adipocyte size decrease, basal lipolysis is also reduced and the net efflux of lipids from adipose tissue decreases. Serum FFA concentrations correlate with total rates of lipolysis and fatty acid fluxes in adipose tissue (25, 35). Therefore, if lipolysis does have a role in the accumulation of ATMs, we predicted that serum FFA would correlate with ATM content in our calorically restricted mice. Indeed, serum FFA concentrations were higher on day 3 of caloric restriction, before significant weight loss, and coincident with the peak in macrophage content (Figure 2A). Consistent with our hypothesis, a positive correlation existed between serum FFA concentrations and the percentage of ATMs in perigonadal adipose tissue throughout the duration of caloric restriction (Figure 2B).

In adipose tissue, the rate-limiting step in both basal and demand lipolysis is regulated by the enzyme encoded by Atgl/Pnpla2. The expression of Atgl/Pnpla2 is regulated by nutritional status and is closely correlated with rates of adipose tissue lipolysis (36). Consistent with there being a peak of adipose tissue lipolysis on day 3 of caloric restriction, the adipose tissue expression of Atgl/Pnpla2 increased on day 3 of caloric restriction and returned to baseline by day 42 (Figure 2C). In contrast, the expression of hormone-sensitive lipase encoded by Hsl/Lipe is not correlated with nutritional status. Hsl/Lipe mRNA levels are downregulated during acute fasting and increase only after prolonged food deprivation (36). Consistent with these data, we did not observe any changes in Hsl/Lipe levels during caloric restriction (Supplemental Figure 9A).

Circulating FFA concentrations and Atgl/Pnpla2 expression provided indirect measures of adipose tissue lipolysis. To directly measure lipolysis in adipose tissue during caloric restriction, the rates of release of nonesterified FFA and glycerol were measured in perigonadal adipose tissue from mice during caloric restriction. Consistent with our indirect measures, lipolysis was increased in adipose tissue from mice following 3 days of caloric restriction compared with adipose tissue from ad libitum–fed mice (Figure 2, D and E). FFA and glycerol release were reduced after 42 days. These data demonstrate a positive correlation between adipose tissue lipolysis and ATM content. To determine directly whether increasing or decreasing lipolysis alters ATM accumulation, we performed a series of dietary, pharmacological, and genetic manipulations.

Lipolysis induces ATM accumulation. If lipolysis drives the accumulation of ATMs in adipose tissue, then fasting, which rapidly
increases adipose tissue hydrolysis of triglycerides, should also increase ATM content. Perigonadal adipose tissue was collected from high-fat–fed obese C57BL/6J mice that were either fasted for 24 hours or fed ad libitum. Fasting induced an increase in serum FFA concentration (Figure 3A) and led to a rapid accumulation of ATMs. Compared with adipose tissue from ad libitum–fed mice, adipose tissue from fasted mice contained 65% more ATMs (percentage of macrophages per total cells: 22.9% ± 3.5% vs. 37.9% ± 3.5%; P < 0.01; Figure 3B–D). Fasting-induced ATM accumulation was not limited to obese mice. In lean mice, the expression of macrophage-specific genes, Emr1 and Csf1r, was increased by 3- and 4-fold respectively after a 24-hour fast (Figure 3E). Consistent with our observations that caloric restriction did not induce an accumulation of CD11c+ ATMs, the expression of Itgax (Cd11c) was unchanged by fasting (Figure 3E).

Activation of the β3-adrenergic receptor (Adrb3), which in mice is almost exclusively expressed by adipocytes, increases adipocyte lipolysis. Granneman and colleagues had previously noted that β3-adrenergic stimulation leads to the accumulation of myeloid–appearing cells in adipose tissue (37). To determine whether pharmacological activation of lipolysis would increase ATM accumulation, we injected lean C57BL/6J mice (body weight = 24.5 ± 0.8 g) with the β3-adrenergic agonist CL316,243 twice, 4 hours apart. Adipose tissue explants were collected 14 hours after the final injection (Figure 3F). Compared with control-treated mice, pharmacological induction of lipolysis rapidly increased ATM content in perigonadal adipose tissue more than 5-fold, to levels typical of obese mice (Figure 3G–J). Macrophages were seen both isolated and in clusters (Figure 3G–I). Thus, among the hormonal signals activated by fasting, β3-adrenergic induction of adipocyte lipolysis is sufficient to induce rapid macrophage recruitment. Similar to our observations in early weight loss and fasting, the β3-adrenergic–induced recruited macrophages do not increase the inflammatory phenotype of adipose tissue (Supplemental Figure 9E).

Adipose triglyceride lipase (also known as desnutrin or patatin-like phospholipase domain–containing protein 2) (Atgl/Pnpla2) regulates both basal and demand lipolysis in adipose tissue. Animals deficient in ATGL/PNPLA2 are severely impaired in their ability to mobilize FFAs, have very low basal lipolysis, and are unable to mount demand lipolysis in response to fasting, despite having an intact hormonal and autonomic response to fasting (36). To provide genetic evidence that lipolysis is a critical determinant of ATM content, we studied the effects of fasting in Atgl/Pnpla2–/– mice. Consistent with our hypothesis, we found that ad libitum–fed Atg1/Pnpla2–/– mice have fewer than 3% ATMs and that following a fast, there was no increase in ATMs (Figure 5A, A and B) or macrophage–specific gene expression (Figure 5C). These data argue that the lipase ATGL/PNPLA2 is required for ATM recruitment and accumulation in adipose tissue.

Adipose tissue lipolysis induces macrophage migration. The rate of mitosis in adipose tissue of fasted mice was very low (<2%) and not different from the rate of mitosis in adipose tissue from ad libitum–fed mice (Supplemental Figure 9, B and C), suggesting that lipolysis–dependent increase in ATMs was not due to proliferation but as a consequence of myeloid cell recruitment. To determine whether lipolysis increases the release of adipose tissue chemottractants for macrophages, we performed a migration assay with

Reducing lipolysis during weight loss or fasting decreases ATM accumulation. If our hypothesis is correct, then manipulations that inhibit lipolysis during early weight loss or fasting should reduce or prevent ATM accumulation. While diets high in fats increase dietary lipids, they are also ketogenic during negative energy balance. Compared with high-fat diets during negative energy balance, diets high in carbohydrate content increase circulating insulin/glucagon ratio and reduce lipid mobilization from and lipolysis rates in adipose tissue (38). Therefore, we repeated the caloric restriction intervention, adding a weight–matched group of mice that were calorically restricted on an isocaloric high–carbohydrate diet (Supplemental Table 1). Mice were maintained on caloric restriction for 3 days, and mice in both groups were equally restricted (30% fewer calories than their ad libitum consumption). At the end of caloric restriction, there was no difference in weight between the high–carbohydrate and high–fat diet calorically–restricted groups. However, as expected, the serum FFA concentration of mice calorically restricted on a high–carbohydrate diet was 36% lower than the mice calorically restricted on a high–fat diet (0.38 ± 0.11 vs. 0.59 ± 0.07 mmol/l; P < 0.05) (Figure 4A). Consistent with our hypothesis, during weight loss, ATM content was 35% lower (31.1% ± 4.1% vs. 20.1% ± 5.2%; P < 0.05) in perigonadal adipose tissue in mice fed a high–carbohydrate diet compared with those fed a high–fat diet (Figure 4, B and C).

Figure 6 Lipolysis induces macrophage migration. Perigonadal adipose tissue explants were isolated from lean mice that were either ad libitum fed or were fasted for 24 hours. Explants from fasted animals were incubated with or without isoproterenol treatment (10 μM). (A) FFA concentration was measured in the explant–conditioned media. (B) The chemotactic activity of control medium, medium supplemented with MCP-1/CCL2 (50 ng/ml), and explant–conditioned medium were measured using a standard migration assay for BMDMs. Data are represented as mean ± SD. n = 4, 5–8 replicates per sample. *P < 0.05; **P < 0.01.
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Adipose tissue explants from fasted and ad libitum–fed mice. Peri-
gonadal adipose tissue explants were collected from lean C57BL/6J
mice that were either ad libitum fed or fasted for 24 hours. Explants
from ad libitum–fed mice were incubated under basal conditions or
with the addition of isoproterenol to induce lipolysis. As expected,
compared with adipose tissue isolated from ad libitum–fed mice,
adipose tissue from fasted mice or adipose tissue treated with iso-
proterenol increased lipolysis as evidenced by an increase in the
release of FFA (Figure 6A). With this increase in lipolysis, there was
a proportional increase in the chemotactic activity of adipose tissue
explants toward bone marrow–derived macrophages (BMDMs). This
increase in adipose tissue chemoattractant activity was comparable
to that induced by MCP-1/CCL2 (Figure 6B). We also found that
in contrast to fasting wild-type mice, fasting CCR2-deficient mice
did not increase the macrophage-specific gene expression (Supple-
mental Figure 9D). However, there was no increase in adipose tissue
expression of known CCR2 ligands during a fast (data not shown).
These data are consistent with the recruitment of ATMs during a
fast being a multi-step process in which accumulation of ATM pre-
ursors in the circulation is CCR2 dependent but transit into fat
depot does require CCR2 or its ligand.

Weight loss and lipolysis activate a program of lipid uptake by ATMs. A
primary function of macrophages is the phagocytosis of tissue-
specific products in a manner that maintains tissue homeostasis.
For example, in bone, osteoclast (the multinucleated macrophage
of bone) resorption of matrix is necessary to maintain bone health
(39). By analogy, the accumulation of ATMs during periods of
elevated lipolysis may permit uptake or phagocytosis of excess
local lipids and participate in the turnover of lipid in adipose tis-

Figure 7
Fasting acutely induces lipid droplet formation in ATMs. (A) Adipose tissue expression of genes whose products, CD36 (Cd36) and scavenger
receptor A (Msr1), are implicated in lipid uptake by macrophages were measured in perigonadal adipose tissue during weight loss induced by
caloric restriction. n = 5–6 mice/group. *P <0.05; ***P < 0.001, versus day 0. (B) Stromal vascular cells (SVCs) isolated from perigonadal adipose
tissue of high-fat diet–induced obese mice that were fed ad libitum (left panel) or were fasted for 24 hours (right panel) were stained for neutral
lipid with oil red O. Scale bars: 50 μm. (C) Number of lipid droplets in macrophages from perigonadal adipose tissue of high-fat diet–induced
obese mice fed ad libitum or fasted for 24 hours. n = 5 mice/group. ***P < 0.001. (D) Expression of genes (in the stromal vascular fraction) encod-
ing proteins involved in lipid uptake, utilization, and export. n = 5 mice/group. *P < 0.05. All data are represented as mean ± SD.
The expression of genes involved in lipid uptake, storage (response to lipolysis per se, we established an in vitro system to unilocular droplets. Consistent with a primary function of ATMs an adrenergic stimulus (isoproterenol). In the absence of adipose tissue (Figure 7D).

In addition, coculture with adipose tissue, and more profoundly with the addition of isoproterenol stimulation, induced Ccr2 gene expression in the SVC fraction, potentially reflecting an activation of a chemotactic macrophage program (Figure 8B). Histologically, induction of lipolysis led to the accumulation of lipid within SVCs (Figure 8, C and D) and increased the appearance of multinucleated lipid-laden macrophages typically seen in adipose tissue of obese individuals (Figure 8E).

**Macrophage depletion increases adipose tissue lipolysis.** The recruitment of ATMs and the uptake of lipid during periods of lipolysis led us to hypothesize that ATMs play a role in regulating local concentrations of FFA. To determine whether ATMs affect lipolysis or lipid release, we depleted ATMs from adipose tissue that harbored a high concentration of macrophages. Epididymal adipose tissue explants were collected from high-fat diet–induced obese mice that were fasting for 24 hours. The explants were treated with liposome-encapsulated clodronate to deplete ATMs, and control explants from the same animals were treated with liposome-encapsulated PBS. Clodronate is a biphosphonate that induces macrophage apoptosis when phagocytosed in liposomes.

and others that form multinucleated giant cells that contain large unilocular droplets. Consistent with a primary function of ATMs being the uptake of lipid during periods of increased release of FFA from adipocytes, the expression in adipose tissue of 2 macrophage lipid transport receptors, Cd36 and Morl, is increased during the initial period of weight loss (Figure 7A). By day 42 of caloric restriction, when adipose tissue lipolysis is reduced, the expression of these 2 genes returns to levels seen in the never–weight-reduced mice. To determine whether lipolysis acutely induces accumulation of lipid within ATMs, we studied lipid droplet content of adipose tissue (right panel) were stained for neutral lipid with oil red O. Lipid-containing cells are marked with arrows. Scale bars: 50 μm.

**Figure 8**
Induction of adipose tissue lipolysis activates lipid uptake by ATMs. (A) SVCs isolated from perigonadal adipose tissue of high-fat diet–induced obese mice were cultured either alone or with perigonadal adipose tissue pieces (harvested from lean animals) with or without isoproterenol treatment (10 μM) to induce lipolysis in the adipose tissue fraction. The gene expression of Adfp and Cd36 in SVCs was measured. n = 5 mice/group. Data are represented as mean ± SD. (B) The expression of the chemokine receptor Ccr2 was measured in SVCs treated as described in A. n = 5 mice/group. (C) SVCs treated with isoproterenol cultured alone (left panel) or with adipose tissue (right panel) were stained for neutral lipid with oil red O. Lipid-containing cells are marked with arrows. Scale bars: 15 μm.

Adipose tissue coculture

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As expected, clodronate treatment reduced by approximately 80% the macrophage content as measured by expression of the macrophage marker Emr1 and the macrophage-expressed scavenger receptor Msr1. Macrophage depletion increased the expression of the lipase Atgl/Pnpla2 by 2.5-fold and increased the expression of fatty acid–regulated genes Fabp4/aP2, Acadl, and Dgat1. The induction of a lipase and genes required for fatty acid metabolism suggested that macrophage depletion increases lipolysis and FFA substrates. Indeed, compared with control-treated adipose tissue explants, macrophage-depleted explants had a 27% higher rate of lipolysis as measured by glycerol release. To assess whether macrophage depletion might have a similar effect on FFA metabolism in vivo, we injected lean C57BL/6J mice intraperitoneally with liposome-encapsulated clodronate to deplete ATMs from intra-abdominal adipose tissue depots and subsequently fasted them for 24 hours. Clodronate treatment compared with control (liposome-encapsulated PBS) treatment reduced macrophage-specific gene expression by more than 90% (Supplemental Figure 10C). Intra-abdominal ATM depletion increased fasting serum (FFA) by 69% compared with control liposome-injected mice (Figure 9C). These data provide preliminary evidence that macrophages function in part to attenuate lipolysis-induced release of FFA.

Discussion

Obesity engenders a complex immune response in which macrophage accumulation in adipose tissue is a characteristic feature. The factors that regulate ATM accumulation are not well defined, and indeed the effects of other metabolic perturbations on ATM accumulation and function have largely been unexplored. In other settings, the recruitment of myeloid-macrophage precursors occurs in response to perturbation of tissue function, typically in response to injury or a foreign pathogen, but also in response to perturbations in lipid fluxes (40). For example, in the arterial wall, local excess cholesterol and lipoproteins drive, through a 2-step process of recruitment and lipid uptake, the accumulation of lipid-laden macrophages (foam cells) (41). Lipid-filled macrophages are also a characteristic finding in adipose tissue from obese individuals (19, 29). We therefore hypothesized that increased lipid fluxes may also regulate the accumulation of ATMs within adipose tissue depots. Here we have shown that the accumulation of macrophages in adipose tissue is an acute response to weight loss and is regulated by adipose tissue lipolysis. These observations are consistent with previous studies of lipolysis in obesity. Total basal lipolysis is chronically elevated in adipose tissue from obese compared with adipose tissue from lean individuals (25) and in intra-abdominal compared with subcutaneous adipose tissue (30, 31). It is in these circumstances and depots, in which lipolysis is higher, that ATM content is also elevated (2, 5, 18, 33).

The role of ATMs in the development of obesity-induced adipose tissue inflammation and its associated pathological sequelae, including insulin resistance, has been intensively studied. Our findings that ATMs rapidly increase lipid uptake in response to adipocyte lipolysis suggest that ATMs may serve an adaptive function, at least over short periods of time, by taking up excess lipid and egress from the tissue. However, in the development of atheromas, where elevated concentrations of cholesterol in vessel walls drive the recruitment of macrophage precursors into subendothelial spaces, the efficient uptake and clearance of lipid from the vasculature is adaptive, as long as macrophage clearance of cholesterol is not overwhelmed. However, in the development of atheromas, the accumulation of cholesterol outstrips the ability of macrophages to clear the lipid; macrophages ingest cholesterol but do not migrate from their anatomic location. Thus, cholesterol-laden macrophages over time form foam cells that remain in the vessel wall and become a central pathogenic component of atherosclerotic lesions (41). During weight loss, both the elevation in lipolysis and the increase in ATMs are transient. This argues that ATMs engage in uptake of excess lipid and egress from the tissue.
However, in obesity, a chronic elevation in lipolysis and local FFA concentrations provides a constant signal for macrophage accumulation. In time, the phenotype of ATMs in obesity, unlike what is seen during weight loss, is altered so that a CD11c+ population, which has been described as more inflammatory, predominates and leads to an impairment of local metabolism.

To understand the functions of ATM, there have been many efforts to delineate the production of inflammatory molecules by ATMs. However, as “professional” phagocytes, macrophages are also efficient in the uptake of a remarkable array of molecules, ranging from small lipids and colonies of pathogens to dying and dead cells (42). The presence of lipid droplets within ATMs, including large unilamellar droplets within multinucleated giant cells, has suggested to some that ATMs primarily function to phagocytose necrotic adipocytes (19). The rapid appearance of lipid drops in ATMs during a fast, even in lean animals, suggests that the recruitment of macrophages to adipose tissue, at least during negative energy balance, is part of a coordinated response that would reduce local extracellular concentrations of FFA.

FFA and other lipids have been found to regulate the activation state and immune function of myeloid cells and macrophages. As extracellular signals, fatty acids, especially saturated fatty acids, activate classical inflammatory responses in macrophages and other immune cells through engagement of pattern recognition receptors, including TLRs (43–45). Our findings demonstrate that lipolysis and increased FFA concentrations can also regulate macrophage accumulation and do so without activation of proinflammatory or M1-polarized state. The more than 40% increase in macrophages during the initial phase of weight loss occurs without increase in inflammatory gene expression or circulating adipokine concentration and supports a model in which lipids, apart from their involvement in ATM activation, play an important and distinct role in ATM recruitment.

The accumulation of ATMs in adipose tissue is only part of the immune response to obesity. T cells are also recruited to adipose tissue during the development of obesity (1, 7–9, 11). If T cells play a role in adipose tissue comparable to that played in athromas, subpopulations of T cells may regulate the recruitment and function of ATMs and other myeloid cells. The complexity of the adipose tissue immune response to metabolic perturbations is further increased by the heterogeneity of ATM populations. The 2 largest classes of ATMs can be distinguished based on expression of the antigens F4/80, CD11b, and CD11c; one population expresses all 3 antigens (FBC cells) and a second expresses only F4/80 and CD11b (FB cells) (46, 47). Data presented here suggest that primarily FB cells accumulate during negative energy balance. This is in contrast to the development of obesity when FBC cells predominate and may explain why ATM accumulation during early weight loss is not accompanied by increases in adipose tissue inflammation and insulin resistance. The ontogeny, functions, and in particular the metabolism of FFA and other lipids of individual subpopulation ATMs still remain to be defined and will likely provide mechanistic insights into differences in the immune response to obesity and weight loss.

Our results support a model in which adipose tissue lipolysis drives ATM recruitment and lipid uptake (Figure 10). These findings suggest that ATMs may play a role in tempering extracellular increases in FFA concentrations during periods of high lipolysis and may thus protect local adipocyte function. In the lean state, adipocytes store little lipid, basal lipolysis is limited, and ATMs are few. When lipolysis is activated and FFA concentrations increase acutely, macrophages accumulate rapidly in adipose tissue, without a significant initial increase in inflammation. Once recruited, ATMs phagocytose excess lipid, possibly reducing adipocyte stress. During weight loss, an increase in demand lipolysis increases local FFA concentrations and thus ATM recruitment. However, progressively, as triglyceride stores decrease and basal lipolysis falls, ATM content is reduced. In obesity, excess accumulation of lipid by large adipocytes increases basal lipolysis, and thereby, net release of FFA. Macrophages are recruited to the adipose tissue but unlike weight loss, in which lipolysis eventually abates, chronic stimulation of ATMs leads to local inflammation and altered metabolic function.

Methods
Mice and weight loss protocol. Male C57BL/6j mice were obtained from the Jackson Laboratory at 9 weeks of age and housed individually in ventilated Plexiglas cages within a pathogen-free barrier facility that maintained a 12-hour light/12-hour dark cycle. Mice were fed a high-fat diet (D12492; Research Diets Inc.) (see Supplemental Table 1 for diet composition details). Food pellets were placed on feeding racks (Wenzel) to improve the accuracy of food intake measurements. Food intake was measured daily for each mouse individually for 1 month before the initiation of caloric restriction. Caloric restriction was initiated when mice weighed 40–41 g. We rotated the assignment of mice to different calorie restriction groups so that the weight of each group was not significantly different at sacrifice (Supplemental Figure 2). During caloric restriction, each mouse received 70% of its ad libitum consumption. The Journal of Clinical Investigation http://www.jci.org
Inc.). For the high-carbohydrate diet caloric restriction experiment, mice were switched to D12450B (Research Diets Inc.) (see Supplemental Table 1 for diet composition details). Food intake was adjusted daily based on mouse weight. Mice were fed at the beginning of the dark and light cycle (they received two-thirds of their food during the dark and one-third during the light cycle. Body composition measurements were performed with the miniSpec TD NMR analyzer (Bruker). Blood samples for baseline insulin measurements were obtained by submandibular bleeding. Blood glucose was measured using a FLASH FreeStyle Glucometer (Abbott). Mice were sacrificed by CO₂ asphyxiation and cervical dislocation after 4-hour fasting during the fifth and sixth hours of the light cycle. Serum samples for final insulin, FFA, and cytokine measurements were collected by cardiac puncture. Abdominal subcutaneous, epididymal, perirenal, and mesenteric adipose tissues were excised, and the right adipose tissue depots and the entire mesenteric depot were weighed for each animal. Tissue samples were frozen in liquid nitrogen and stored at −80°C prior to RNA extraction and immunohistochemical analysis. The experiment was performed on 2 separate occasions (n = 40 per experiment). All animal experiments were approved by the Columbia University IACUC.

**Fasting experiments.** Lean C57BL/6J mice (8 to 9 weeks old) fed a standard pellet diet (PicoLab Rodent Diet 20; Purina Mills Inc.) were fasted for 24 hours starting on the second hour of the light cycle. Weight andagematched ad libitum-fed C57BL/6J mice were used as controls. Perigonadal adipose tissue was excised, and samples were stored as described above. For fasting experiments on high-fat diet-induced obese mice 24- to 28-week-old mice were used. Mice were placed on a high-fat diet (D12492/Research Diets Inc.) starting at 6 weeks of age. They underwent a 24-hour fast starting on the second hour of the light cycle.

Five 8-week-old Ccr2−/− and five 8-week-old Ccr2+/+ mice (litterates) fed a standard pellet diet (PicoLab Rodent Diet 20; Purina Mills Inc.) were fasted for 24 hours starting on the second hour of the light cycle. Five weight- and age-matched Ccr2−/− and 5 Ccr2+/+ mice (littermates) were used as controls. Perigonadal adipose tissue was excised, and samples were stored as described above.

**CL316,243 treatment of mice.** Ten 8-week-old C57BL/6J mice fed the standard pellet diet (see above) were injected intraperitoneally with 1 mg/kg of CL316,243 or saline twice (4 hours between each injection). Mice were sacrificed 14 hours after the second injection. Perigonadal adipose tissue was excised and samples were stored as described above.

**BrdU treatment of mice.** Ten 8-week-old C57BL/6J mice fed the standard pellet diet (see above) were injected intraperitoneally with 0.133 mg/g of BrdU (Sigma-Aldrich) or vehicle twice (3 hours between each injection). 1.5 hours after the first injection, we removed food from the experimental group. The control group was fed ad libitum. Both groups were sacrificed 24 hours later. Perigonadal adipose tissue was excised, and samples were stored as described above.

**Atg1−/− and Atg5−/− mice.** Perigonadal adipose tissue samples from Atg1−/− and Atg5−/− mice were collected from 8- to 9-week-old lean mice that were ad libitum fed or fasted for 16 hours.

**Metabolic analyses.** Serum insulin levels were determined using an ultra-sensitive insulin ELISA (Merckia Inc.). Serum PAI-1 and resistin were measured by using the LINCOplex Mouse Adipokine Panel (Millipore). Serum leptin was measured by using the quantikine mouse leptin ELISA (R&D systems). FFA were measured using the HR NEFA series (Wako Diagnostics). Blood samples for serum isolation were collected after a 4-hour fast during the fifth and sixth hours of the light cycle. The serum samples collected represented in Figure 2A were also analyzed for several hormones and cytokines and were therefore placed on wet ice before further processing. The serum sample represented in Figure 3A and Figure 4A were used only for FFA analysis and were immediately frozen in liquid nitrogen before analysis (to minimize triglyceride lipolysis).

**Immunohistochemistry.** Adipose tissue samples were fixed for 48 hours at room temperature in zinc-formalin fixative (Anatech Ltd.), incubated in 70% ethanol for 24 hours, and subsequently embedded in paraffin. 5-μm sections, cut at 50-μm intervals, were mounted on charged glass slides, deparaffinizied in xylene, and stained for expression of F4/80 with a rat anti-mouse F4/80 monoclonal antibody (Abd Serotec). Sections were incubated with the primary antibody for 80 minutes at room temperature (1:100 dilution). Rat IgG2a (Invitrogen) was used as isotype control (1:50 dilution). Subsequently, a biotinylated anti-rat secondary antibody was used at 1:200 dilution followed by the Avidin DH: biotinylated horseradish peroxidase H complex (Vector Laboratories) and development in chromogen substrate 3,3′-diaminobenzidine (Vector Laboratories). Slides were counterstained with hematoxylin. For each individual adipose tissue depot, 5–10 different high-power fields from each of 3 different sections were analyzed. The total number of nuclei and the number of nuclei of F4/80-expressing cells were counted for each field. The fraction of F4/80-expressing cells for each sample was calculated as the sum of the number of nuclei of F4/80-expressing cells divided by the total number of nuclei in sections of a sample. Cross-sectional area was determined for each adipocyte in each field using image analysis software Image-Pro Plus (Media Cybernetics Inc.). For each mouse, 800–1000 adipocytes were counted. Adipocyte number was calculated from fat pad weight and adipocyte volume (48).

For detection of BrdU-positive cells, sections were prepared as described above and immunohistochemistry was performed using a monoclonal biotinylated anti-BrdU (BrdU staining kit; Invitrogen). Slides were counterstained with hematoxylin. For each individual adipose tissue depot, 10 different high-power fields from each of 3 different sections were analyzed. The total number of nuclei and the number of nuclei of BrdU-positive cells were counted for each field. The fraction of BrdU-positive cells for each sample was calculated as the sum of the number of nuclei of BrdU-positive cells divided by the total number of nuclei in sections of a sample.

For detection of CD11c-positive cells, frozen sections (10 μm) were used. Sections were stained for expression of CD11c with a hamster anti-mouse CD11c antibody (Abd Serotec). Sections were incubated with the primary antibody for 60 minutes at room temperature (1:100 dilution). Hamster IgG was used as isotype control (1:100 dilution) (Abd Serotec). Subsequently, a biotinylated anti-hamster secondary antibody was used at 1:200 dilution (30 minutes) followed by the Avidin DH: biotinylated horseradish peroxidase H complex (Vector Laboratories) and development in chromogen substrate 3,3′-diaminobenzidine (Vector Laboratories). Slides were counterstained with hematoxylin. The total number of nuclei and the number of CD11c-positive CLS were counted for each field. The fraction of CD11c-positive CLS for each sample was calculated as the sum of the number of positive CLS divided by the total number of nuclei in sections of a sample.

For double-immunofluorescence staining of F4/80- and CD11c-expressing cells, frozen sections were used. The sections were incubated with the primary antibodies overnight at 4°C (1:100 dilution). Subsequently, a donkey anti-rat Cy3 IgG (Jackson ImmunoResearch) and a goat anti-hamster Alexa Fluor 488 IgG were added (Invitrogen). Sections were incubated for 45 minutes at room temperature (1:300 dilution). Fluorescent microscopy was performed using a Nikon Eclipse 80i equipped with a Retiga Exi camera and X-Cite 120 Fluorescent Illumination System.

**Quantitative RT-PCR.** RNA was extracted from frozen adipose tissue using an acid-phenol reagent (TRIZol; Invitrogen). RNA was isolated from SVCs using the QIAGEN RNeasy MiniKit (QIAGEN) and used as template for cDNA synthesis using Superscript III reverse transcriptase (Invitrogen) and random hexamer primers. Quantitative RT-PCR assays were carried out using DNA Engine Opticon 2 system instruments (Bio-Rad) and PCR SYBR Green I.
QuantiTect Master Mix (QIAGEN). The mRNA expression of all genes reported is normalized to the cyclophilin B (Ppib) gene expression. Every reaction was performed in duplicate, and the data were analyzed with the 2-DDCT method (49). All primers used are listed in Supplemental Table 3.

In vitro lipolysis of isolated perigonadal adipose tissue (explants). Perigonadal fat pads were surgically removed from high-fat-fed C57BL/6J mice (n = 5/group), which were either ad libitum fed or were undergoing caloric restriction. Subsequently, they were washed several times with PBS. Tissue pieces (~100 mg total weight) were incubated in DMEM (Invitrogen) containing 2% fatty acid–free BSA (Sigma-Aldrich) at 37°C. Aliquots were collected after 120 minutes and investigated for FFA and glycerol content. FFA and glycerol were measured using the HR NEFA series (Wako Diagnostics) and the free glycerol determination kit (Sigma-Aldrich), respectively. We measured adipocyte numbers in parallel samples from weight- and fat-matched ad libitum–fed and calorically restricted mice.

Coculture experiments. To isolate SVCs, perigonadal adipose tissue was isolated from high-fat diet–induced obese C57BL/6J male mice immediately after CO2 asphyxiation. Tissues were handled using sterile techniques and minced into fine (<10 mg) pieces. For SVC isolation, minced samples were moved and DMEM containing 2% fatty acid–free BSA (Sigma-Aldrich) were added with 20% liposomes containing either clodronate or PBS. After 6 hours, fresh media were added with 20% liposomes containing either clodronate or PBS, reaching a final volume of 1 ml. Explants were maintained for 48 hours at 37°C in a CO2 incubator. Subsequently, the media containing liposomes were moved and DMEM containing 2% fatty acid–free BSA (Sigma-Aldrich) was added. Aliquots were collected after 120 minutes and investigated for glycerol content. Glycerol was measured using the free glycerol determination kit (Sigma-Aldrich). Explants were also used for RNA isolation.

Statistics. Data are presented as mean ± SD. All P values were calculated using 2-tailed distribution, 2-sample unequal variance Student’s t test. All calculations were performed using Microsoft Excel and Statistica (Statsoft Inc.).

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