OBJECTIVE—Obesity induces a program of systemic inflammation that is implicated in the development of many of its clinical sequelae. Hepatic inflammation is a feature of obesity-induced liver disease, and our previous studies demonstrated reduced hepatic steatosis in obese mice deficient in the C-C chemokine receptor 2 (CCR2) that regulates myeloid cell recruitment. This suggests that a myeloid cell population is recruited to the liver in obesity and contributes to nonalcoholic fatty liver disease.

RESEARCH DESIGN AND METHODS—We used fluorescence-activated cell sorting to measure hepatic leukocyte populations in genetic and diet forms of murine obesity. We characterized in vivo models that increase and decrease an obesity-regulated CCR2-expressing population of hepatic leukocytes. Finally, using an in vitro co-culture system, we measured the ability of these cells to modulate a hepatocyte program of lipid metabolism.

RESULTS—We demonstrate that obesity activates hepatocyte expression of C-C chemokine ligand 2 (CCL2/MCP-1) leading to hepatic recruitment of CCR2 expressing myeloid cells that promote hepatosteatosis. The quantity of these cells correlates with body mass and in obese mice represents the second largest immune cell population in the liver. Hepatic expression of CCL2 increases their recruitment and in the presence of dietary fat induces hepatosteatosis. These cells activate hepatic transcription of genes responsible for fatty acid esterification and steatosis.

CONCLUSIONS—Obesity induces hepatic recruitment of a myeloid cell population that promotes hepatocyte lipid storage. These findings demonstrate that recruitment of myeloid cells to metabolic tissues is a common feature of obesity, not limited to adipose tissue. Diabetes 59:916–925, 2010

Obesity induces hepatic accumulation of triglycerides (TGs) and, in a large proportion of obese individuals, leads to the development of nonalcoholic fatty liver disease (1). Liver TG storage in the context of obesity is achieved through a complex program coordinated by transcriptional regulators, including SREBP1c (2), PPAR-γ (3,4), Foxo1 (5), and XBPl (6). Obesity also increases delivery of free fatty acids (FFAs) to the liver from adipose tissue, which are esterified and deposited as TG in cytoplasmic lipid droplets, and decreases TG export by promoting endoplasmic reticulum stress (7).

Another cardinal feature of obesity and nonalcoholic fatty liver disease is hepatic inflammation. Similar to its effects on adipose tissue, obesity induces hepatic inflammation, as reflected by increased production of proinflammatory cytokines and acute-phase reactants and by activation of NF-κB and JNK regulated pathways (8,9). In addition, myeloid-macrophage populations are key mediators of obesity-induced inflammation. Recent data suggest a role for local immune cells in the regulation of hepatic insulin sensitivity in response to obesity. Deletion of Ppard within hematopoietic cells polarizes the liver resident macrophages, Kupffer cells (KCs), toward an M1@classically activated state and reduces systemic insulin sensitivity in high-fat–fed mice (10,11). Despite the evidence of a close relationship between hepatosteatosis and inflammation, this inflammatory state remains poorly characterized.

In adipose tissue, inflammation is caused in part by the recruitment and activation of CCR2+ adipose tissue macrophages. Consistently, mice overexpressing CCL2, the primary ligand of CCR2, in adipose tissue accumulate more adipose tissue macrophages (ATMs) and are insulin resistant (12,14). Interestingly, CCR2 deficiency protects mice from nonalcoholic fatty liver disease after high-fat feeding, implying that a CCR2-bearing cell is critical for development of hepatosteatosis. Studies in humans have shown that hepatic expression of CCL2 is increased in obesity and nonalcoholic fatty liver disease (16). These observations lead us to hypothesize that obesity induces recruitment of a CCR2+ myeloid cell population that promotes hepatic steatosis.
from each of the following groups: C57BL/6J males, high-fat-fed C57BL/6J males, calorie-restricted C57BL/6J males, calorie-restricted B6.Cg-Ay/" females, B6.Cg-Ay/" females, and exercised male B6.Cg-Ay/" males, as well as five male B6.Cg-Ay/" males. Three from each of the following groups were included: C57BL/6J females, B6.Cg-Ay/" females, and exercised B6.Cg-Ay/" females, which were injected with Rnase, pioglitazone, B6.Cg-Ay/" females, and exercised B6.Cg-Ay/" females with pioglitazone (Takeda NA), and B6.Cg-Ay/" females treated with CI316243 (Sigma-Aldrich). Data and experimental details have been deposited in the GEO database (www.ncbi.nlm.nih.gov/geo).

Liver perfusion. Perfusion was performed as described previously with slight modification (19). Mice were anesthetized, and the vena cava was catheterized using a 23-gauge catheter. The thoracic vena cava was then clamped, and the perfusion was initiated at 5 ml/min with buffers described by Liver perfusion.

Immunophenotyping. Liver cells were cooled on ice and counted using a hemocytometer. They were centrifuged at 500 g for 5 min and resuspended in FACS buffer at a concentration of 7 × 10^6 cells/ml. Cells were incubated in the dark for 30 min in FcBlock (BD Pharmingen) and then for an additional 45 min with fluorophore-conjugated antibodies or isotype controls. The antibodies used were as follows: CD11b-PE and APC, CD11c-PE and PE-TR, CD3e-PE and FITC, and NK1.1-APC (eBioscience), B220-PE-TR (Caltag Laboratories), CD45.1-PE and Cy5.5 (BD Pharmingen), and F4/80-PE (AbD Serotec). After incubation with primary antibodies, cells were washed twice, resuspended in FACS buffer containing 4',6-diamidino-2-phenylindole dihydrochloride (DAPI), and analyzed on an LSRII flow cytometer (Becton Dickson) equipped with four excitation lines. Data analysis was performed using FlowJo software (Tree Star). All samples were gated on DAPI^- cells before quantification. Cells were sorted into Dulbecco’s modified Eagle’s medium (DMEM) containing 20% fetal bovine serum on a BD FACS Aria Cell Sorter.

Tissue culture. Hepatocytes were isolated from dissociated livers by three 50g centrifugations followed by centrifugation over Percoll density gradient for 10 min at 50g. Hepatocytes were plated on Collagen I (Sigma-Aldrich) coated 24-well plates and were washed after 2 h in DMEM 10% fetal bovine serum, 1% penicillin/streptomycin. After 2 more hours, cells were washed and media were replaced with Hepatocyte (Invitrogen), a proprietary formulation with a glucose concentration of 4.5 g/l, and co-cultured with recruited myeloid cells (RMCs) for 18 h. RMCs were sorted from obese mice after 20–25 weeks of high-fat feeding and cultured in macrophase medium (DMEM containing 20% L-cell conditioned medium, 10% fetal bovine serum, 1% penicillin/streptomycin) on cell culture inserts for 24 h before hepatocyte co-culture.

Adenoviral preparation. Recombinant adenovirus encoding human CCL2 or LacZ (Vector BioLabs) was amplified in HEK293 cells (ATCC) and purified with an Adeno-X Maxi Purification Kit (Clontech Laboratories). The 7-week-old male mice were injected with recombinant adenovirus intravenously at a dose of 1.1 × 10^11 pfu/g in a total volume of 0.1 ml. Mice were fasted for 6 h before intraperitoneal injection of human regular insulin (Humulin R; Eli Lilly). After injection, glucose was measured with the FreeStyle blood glucose meter (Abbott Laboratories). Mice were re-euthanized and blood samples were collected from the tail vein to measure blood glucose levels.

Metabolic studies. Fasting glucose and insulin measurements were taken after a 6-h fast. Glucose was measured with the FreeStyle blood glucose meter (Abbott Laboratories). Insulin was measured using Ultrasensitive Mouse Insulin ELISA (Alpco Diagnostics).

The insulin tolerance test was performed 3 weeks after infection. Mice were fasted for 6 h before intraperitoneal injection of human regular insulin (0.75 units/kg) and insulin-stimulated glucose levels were measured after 6 h by tail-vein puncture. Blood was collected from the tail vein of mice that were injected with insulin (5 units/kg) through tail vein after a 6-h fast, and livers were removed after 3 min. Homogenized proteins were immunoblotted for total Akt and phospho-Akt (Cell Signaling Technology). For SREBP1c Western blotting, nuclear and cytoplasmic protein fractions were isolated from whole tissue (Active Motif) and immunoblotted using anti-SREBP1c antibody (SC13551; Santa Cruz Biotechnology, Santa Cruz, CA).

Measurement of hepatic TG. TG content was quantified using a colorimetric assay kit (Thermo Fisher Scientific) as previously described (3). Livers were snap-frozen and stained with oil red O to visualize lipids. Aria Cell Sorter.

Statistical analysis. Data are presented as mean ± SD. For all comparisons, P values were calculated using a Student’s t test (two-tailed distribution, unequal variance). To assess associations, Pearson correlation coefficients were calculated.

RESULTS

Chemokines direct the trafficking of immune cells to sites of inflammation. To determine whether obesity activates a chemokine-driven program in liver, we measured expression of chemokines and chemokine receptors in the livers of a cohort of mice with varying degrees of adiposity. We identified only one chemokine ligand and receptor pair whose expression was positively correlated with body mass in mice (supplemental Tables 1 and 2). Expression of the chemokine receptor Ccr2 (r = 0.46, P < 0.01) and Ccl2 (r = 0.40, P < 0.01) correlated positively with body mass (Fig. 1A). In addition, Ccr2 expression correlated with expression of known acute-phase response genes and inflammatory cytokines (supplemental Table 3), suggesting that Ccr2 is part of the larger obesity-induced inflammatory program in the liver.

The recruitment of macrophages to adipose tissue and other inflamed tissue is regulated by a subset of genes that includes chemokines, chemokine receptors, adhesion molecules, myeloid markers, and inflammatory cytokines. We looked for a similar expression profile in the livers of leptin-deficient obese mice. We confirmed the findings from our microarrays that hepatic Ccl2 and Ccr2 expression is increased in obese mice (Fig. 1B). In addition, these mice expressed greater amounts of adhesion molecules (Icam1, Vcam1) required for recruitment of immune cells to tissues, and markers expressed by macrophage and populations (Irigam, Emr1). Expression of the potent inflammatory cytokine Tnf was elevated as well. When hepatocytes and the immune cell (CD45^+ ) fractions were isolated from livers of obese mice, Ccr2 expression was detected exclusively in the immune cell population (Fig. 1C). Furthermore, hepatocyte expression of Ccl2 was induced ninefold in hepatocytes isolated from obese compared with hepatocytes from lean mice (Fig. 1D).

Despite these findings, we and others have previously shown that KCs are not increased by obesity (8,18). To determine whether obesity recruited non-KC immune cells to the liver, we cataloged immune cells using FACS (Fig. 2A) and accounted for >5% of hepatic immune cells. Among myeloid cells, KCs express the F4/80 (Emr1) macrophage antigen at high levels and were the largest immune cell population. However, we found another population of myeloid cells (CD11b^+ ) that expressed F4/80 at low levels (Fig. 2B). Obesity modestly reduced the population of KCs (CD45^-F4/80^+) but lead to a doubling in the distinct CD45^-/CD11b^+ /F4/80^dim cells from 10.0 to 19.7% (Fig. 2C) in C57BL/6J Lep^obob. Only a small fraction of
Leptin genes involved in myeloid cell recruitment (mice. As in the logged immune cells from high-fat diet (HFD)-fed obese features of obesity or specific to leptin deficiency, we cata-

Fig. 1). Numbers in obese compared with lean mice (supplemental 

bution of immune cells but an increase in its absolute 

css suggest that KCs are larger and more granular than 

ally distinct from KCs. Forward and side light scatter 

body mass (Fig. 2). Consistent with our hypothesis, their number correlated with 

Fig. 1. Obesity induces a chemotactic program in the liver. A: Body 

mass correlates with hepatic expression of Ccl2 and Ccr2 among mice with varying body mass due to genetic mutations, sex, and diet (n = 45, 
P < 0.01 for Ccr2 and Ccl2). B: Obesity induces hepatic expression of genes involved in myeloid cell recruitment (n = 8–9). [C] 57BL/6J 

Lep+/+; □, Lep ob/ob. C: Hepatic immune cells (CD45+) but not hepatocytes express Ccr2 (n = 4). D: Hepatocyte expression of Ccl2 is induced in obese leptin-deficient mice compared with lean mice (n = 4). **P < 0.01; ***P < 0.001; ****P < 0.0005. AU, arbitrary units.

these expressed the granulocyte marker Ly6G (2–4% of cells). In addition, we confirmed a previous report (21) that leptin deficiency reduces the number of NKT cells (CD45+/CD3ε+/NK1.1+). The increase in the CD45+/CD11b+/F4/80dim population did not represent a redistribution of immune cells but an increase in its absolute numbers in obese compared with lean mice (supplemental Fig. 1).

To determine whether these changes were general features of obesity or specific to leptin deficiency, we cataloged immune cells from high-fat diet (HFD)-fed obese mice. As in the Lepob/ob model, HFD-fed C57BL/6J mice had reduced KC numbers and increased CD45+/CD11b+/F4/80dim cells compared with LPD-fed control mice (Fig. 2D); therefore, these changes are common features of obesity and not the consequence of a specific diet or genetic alteration. We refer to the CD45+/CD11b+/F4/80dim cells as “recruited myeloid cells” (RMCs), and consistent with our hypothesis, their number correlated with body mass (Fig. 2E).

This population was morphologically and transcriptionally distinct from KCs. Forward and side light scatter profiles suggest that KCs are larger and more granular than RMCs (Fig. 3A). Likewise, isolated RMCs are smaller than resident KCs and contain less cytoplasm. Nonetheless, both cell types share the typical mononuclear myeloid morphology, with oval/kidney-shaped nuclei (Fig. 3B).

The transcriptional profile of RMCs was consistent with their being a monocytic-myeloid population recruited to the liver through CCR2. As expected based on our sorting criteria, KCs expressed high levels of Emr1 (F4/80), while RMCs expressed higher amounts of Itgam (CD11b) (supplemental Fig. 2). We hypothesized that the recruitment of RMCs is responsible for the increase in Ccr2 expression noted in livers of obese mice. Indeed, RMCs expressed Ccr2 at >100 times the level of KCs, indicating that this population is primarily responsible for hepatic Ccr2 expression (Fig. 3C).

Among monocytic lineages of myeloid cells, CCR2 expression is associated with the classically inflammatory (M1 polarized) phenotype. As expected, RMCs expressed greater amounts of Tnf than KCs (Fig. 3D). Conversely, the expression of the M2 markers Arg1 and Jag1 were higher in KCs. However, not all markers of M2 polarization were higher in KCs compared with RMCs. The expression of Chi3l3(Ym1), a heparin binding lectin that is induced by M2 polarization (22), was >30 times higher in RMCs than KCs (Fig. 3D). Although these data suggest that the activation/polarization state of KCs and RMCs cannot simply be described as M1 or M2, they do argue that activation phenotypes of KCs and RMCs are distinct.

ATM accumulation in part depends on CCR2 (12). The elevated expression of CCL2 in obese liver, and the high level of expression of CCR2 in the RMCs, suggest that this chemokine ligand/receptor pair may be important for obesity-induced accumulation of RMCs. To test this possibility, bone marrow isolated from CCR2-deficient (C57BL/6J Ccr2−/−) or control (C57BL/6J Ccr2+/+) mice was adoptively transferred into irradiated C57BL/6J mice yielding CCR2-chimeric (CCR2KO/WT) and control (CCR2WT/WT) mice (schematic Fig. 4A). After recovery and short-term HFD feeding, there was no difference in body mass, fasting serum insulin, or blood glucose between CCR2KO/WT or CCR2WT/WT mice (Fig. 4B). However, CCR2WT/WT recipients accumulated twice the number of RMCs than weight-matched CCR2KO/WT mice (10.5 vs. 5.5%, respectively, Fig. 4C). As would be expected for a recruited and rapidly turning-over population of cells, the percentage of donor-derived RMCs in CCR2WT/WT and CCR2KO/WT recipients was 85% (± 6.3%) and 88% (± 7.7%), respectively, 8 weeks after bone marrow transplantation. In contrast, only 10% (± 22.4%) and 35.6% (± 20.6%) of KCs were donor-derived in CCR2WT/WT and CCR2KO/WT recipients, respectively. Bone marrow transplantation significantly reduces the susceptibility of mice to obesity (23). However, after becoming obese (body mass >40 g, ~8 months), CCR2KO/WT accumulated less hepatic TG than weight-matched CCR2WT/WT mice (132 ± 54 vs. 204 ± 58 mg/g, respectively, P < 0.05; n = 5) (supplemental Fig. 4). Thus, obesity-induced accumulation of RMCs and hepatic steatosis both in part depend on hemopoietic CCR2.

In mice, CCR2 has four known ligands (CCL2, CCL7, CCL8, and CCL12) that can induce chemotaxis in receptor-bearing cells (24); however, hepatic expression of only Ccl2 was elevated in obesity (supplemental Fig. 3). This suggests that liver-derived CCL2 is responsible for RMC recruitment in obesity. To test whether hepatic expression of CCL2 was sufficient to recruit RMCs to the liver, mice were injected with a recombinant adenovirus (Ad-hCcl2) designed to induce expression of human CCL2 (schematic,
CCL2 binds and activates murine CCR2, albeit at a reduced affinity (25). Expression of human Ccl2 was detected in serum at 21 days and was expressed exclusively in the liver (Fig. 5C and D). One week after infection, RMCs were increased in livers of mice expressing human Ccl2 but not in the livers of control mice infected with a LacZ-containing adenovirus (Fig. 5E). As expected based on Fig. 2D, among Ad-LacZ–injected mice, those on an HFD accumulated more RMCs than those on an LFD (6.3 ± 0.6 vs. 11.7 ± 2.8%, \( P < 0.05 \)). These data support the hypothesis that obesity-induced expression of CCL2/MCP1 is sufficient to increase recruitment of RMCs.

Genetic deficiency of Ccr2 in HFD-fed mice reduces hepatic steatosis (12). To test whether CCL2-dependent RMC recruitment was sufficient to replicate obesity-induced hepatic steatosis, we measured the effects of
hepatic Ad-hCcl2 in mice. Body mass, insulin sensitivity, and serum lipids were not changed in mice overexpressing hepatic hCCL2 (Fig. 5B, Fig. 6A, and supplemental Fig. 6A and B). We found that in LFD-fed mice, increased hepatic RMC content did not affect hepatic TG concentration (Fig. 6C and D); thus, RMCs are not sufficient to induce steatosis. However, adenosine-mediated expression of Ccl2 for 3 weeks in HFD-fed mice did increase hepatic TG content by 65% (P < 0.01; Fig. 6C and D) compared with HFD-fed controls. This occurred without evidence of liver hepatomegaly, steatohepatitis, or elevation of serum transaminase concentration (Fig. 6B and supplemental Fig. 5A and B). Thus, RMCs can induce the accumulation of hepatic TG, but only in the presence of excess dietary lipids.

Hepatic expression of CCL2 in mice fed an HFD activates a lipogenic program that included upregulation of lipogenic and TG synthesis genes (Fig. 7A); thus, RMCs are not sufficient to induce steatosis. Indeed, CCL2 activates a lipogenic program that included upregulation of lipogenic and TG synthesis genes (Fig. 7B). Obesity-induced steatosis is driven in part by hyperinsulinemia associated with hepatic insulin resistance (1); insulin’s inhibition of gluconeogenesis is impaired, whereas insulin-regulated lipogenesis remains intact, resulting in increased TG production (2). These derangements in insulin signaling did not appear to contribute significantly to RMC-induced steatosis. Insulin tolerance tests in Ad-hCcl2 mice were not different from controls (supplemental Fig. 6A), and differences in homeostasis model assessment of insulin resistance (HOMA-IR) values did not reach significance (P = 0.2 HFD Ad-LacZ vs. Ad-hCcl2). Similarly, hepatic overexpression of hCCL2 did not impair hepatic insulin signaling in HFD-fed mice (supplemental Fig. 6C). These data suggest that RMC lead to increased lipogenic gene expression and TG deposition as a result of modulating SREBP1c-dependent and -independent transcriptional activity, likely in a manner not driven by alterations in insulin signaling. Dissociation of hepatic insulin resistance and steatosis is also seen when TG esterification genes are overexpressed in liver. Modest hepatic overexpression of Dgat2 or Gpam increases hepatic steatosis in mice without increasing fasting glucose or insulin (26,27). Conversely impaired hepatic lipid storage can lead to liver damage (28). Thus, TG formation can be adaptive and over the short term protect the liver from the adverse effects of increased FFAs. If RMCs facilitate TG esterification, they could be predicted to activate a program of RMC recruitment. Indeed, 6-h infusion of oleate increased hepatic expression of CCL2 (supplemental Fig. 9).

To test whether RMCs could activate a hepatocyte
program of TG esterification, we co-cultured primary RMCs from HFD-fed obese mice with primary hepatocytes from lean mice for 18 h. The expression of both Dgat2 and Gpam was induced in hepatocytes co-cultured with RMCs (Fig. 8). In contrast, lipogenic and β-oxidation programs were unaffected by short-term co-culture with RMCs (Fig. 8). The esterification of fatty acid into TG requires glycerol-3-phosphate as a substrate, and recent evidence suggests that production of glycerol-3-phosphate from pyruvate (glyceroneogenesis) is the dominant source of liver TG glycerol (29). Co-cultured RMCs increase hepatocyte expression of Pck1 (PEPCK) without modulating the expression of G6pc. Whereas coordinately increasing Pck1 and G6pc drives gluconeogenesis in hepatocytes, isolated upregulation of Pepck would drive pyruvate to glycerol rather than to glucose.

If the most proximal effect of RMCs on liver is to drive expression of genes involved in lipid storage, one would expect to find these differences in LFD-fed Ad-hCcl2–treated mice, despite there being no difference in TG per se. In fact, expression of Gpam and Dgat2 was elevated in standard diet–fed mice in which RMC numbers were increased by hepatic CCL2 expression, 2 weeks after adenoviral infection (supplemental Fig. 7).

**DISCUSSION**

The response of the immune system to foreign pathogens and injury is critical for restoring tissue function after an insult. However, recent studies suggest that the immune system has also evolved to respond to metabolic stresses and that immune cells have a role in maintaining metabolic homeostasis (30). Most of these studies have focused on populations of ATMs and their long-term maladaptive contributions to insulin resistance and impaired glucose homeostasis. Recent findings have implicated impairment of M2 polarization of KCs in the development of insulin resistance (11). Here we have identified a population of hepatic myeloid cells, distinct from KCs, that are recruited to the liver in proportion to body mass/adiposity in a CCR2/CCL2-dependent fashion.

We have provided three lines of evidence demonstrating a key role for these cells in the pathogenesis of obesity-induced steatosis. First, CCR2 deficiency impairs RMC accumulation and likewise reduces lipogenic gene expression and hepatic TG accumulation. It is difficult to assess the relative contributions of other CCR2-dependent populations, such as ATMs, to this phenotype. Other studies have directly implicated inflammatory signals from adi-
pose tissue in modulating hepatic TG content and insulin sensitivity. Nevertheless, the proximity of RMCs to hepatocytes suggests that they contribute to the regulation of hepatic TG metabolism.

Second, hepatic CCL2 expression leads to recruitment of RMCs, induction of lipogenic genes, and, when accompanied by high-fat feeding, hepatic steatosis. However, when CCL2 is overexpressed in LFD-fed mice, RMCs are recruited (Fig. 5E) and expression levels of Gpam and Dgat2 are elevated (supplementary Fig. 7), but this is not associated with hepatic TG accumulation. These observations not only corroborate our primary hypothesis but imply that the mechanism through which RMCs induce steatosis is by elevating hepatic lipogenic gene expression. This alone is insufficient to drive lipid accumulation without the availability of excess circulating metabolic substrates (e.g., FFAs) to be channeled into this metabolic pathway; consequently, RMC recruitment is insufficient to induce steatosis in LFD-fed subjects.

Lastly, co-culture experiments suggest that a primary mechanism by which RMCs effect TG accumulation is by activating a transcriptional program that facilitates FFA esterification into TG. Over the short term, this effect may provide an adaptive mechanism to reduce local concentrations of the more biologically active FFAs and favor storage of more inert TG. However, in the setting of chronic lipid excess, this would compromise hepatocyte metabolic function. During the preparation of this work, an article appeared that provides parallel evidence for the changes in hepatic myeloid cell populations (31). The authors furthermore provide an alternative mechanism by which hepatic myeloid cells induce liver inflammation,
namely by promoting NKT cell apoptosis, although it is unclear whether this would lead to metabolic dysfunction.

Accumulation of RMCs in part depends on the chemokine receptor CCR2. This suggests that this population derives from CCR2-expressing M1 polarized circulating monocytes. Likewise, RMCs express TNF to a greater degree than KCs, an alternatively activated tissue resident macrophage population that is critical in maintaining hepatic insulin sensitivity in obese animals (11). However, expression of genes considered markers of M2-activated macrophages such as Chi3l3 and Mrc was inconsistent, with RMCs being categorized simply as M1 polarized. A likely conclusion based on this observation is that the activation state of the RMCs cannot be defined strictly as either M1 or M2 polarized.

While RMCs are distinct from KCs, it is unclear whether they are ontologically related. If RMCs are indeed precursors to KCs, our data would imply that obesity accelerates the turnover of KCs, and RMCs are recruited at an accelerated rate to replenish the KC population. However, there appears to be a reciprocal relationship between the size of the RMC and KC populations: manipulations that increase RMCs, i.e., obesity or hepatic CCL2 expression, reduce the KC population. Therefore, if RMCs do serve as precursors to KCs or a subpopulation of KCs, the relationship would appear to be complex and our data would suggest that accelerated accumulation of RMCs can drive the efflux of KCs.

While we have demonstrated that RMCs contribute to the development of obesity-induced hepatic steatosis, hepatic TG storage is a feature of other diseases, including alcoholic fatty liver disease and lipodystrophy. If RMCs promote hepatocyte lipid storage more generally, RMCs may contribute to these disorders as well. We demonstrated that infusing oleate increases hepatic expression of

FIG. 6. RMCs regulate hepatic TG accumulation in the presence of excess lipid. Hepatic expression of hCCL2 and expansion of the RMC content did not alter insulin sensitivity (A) or liver mass (B) of mice on LFD or HFD (n = 4). C: Increased recruitment of RMCs did not increase hepatic TG in LFD-fed mice but did induce steatosis in mice with excess lipid intake (Ad-LacZ, Ad-hCcl2). D: Hepatic expression of hCCL2 and recruitment of RMCs increased histologically visible hepatic steatosis and lipid droplets in HFD-fed mice (bar = 50 microns). *P < 0.05.
RMCs promote expression of lipogenic genes. A: Increased recruitment of RMCs through hepatic expression of hCCL2 increased lipogenic gene expression in HFD-fed mice (\(\square\), Ad-LacZ; \(\blacksquare\), Ad-hCcl2). B: Deficiency of CCR2 in weight-matched mice fed an HFD for 9 weeks reduced lipogenic gene expression (\(\square\), CCR2\(^{+/+}\); \(\blacksquare\), CCR2\(^{-/-}\) (n = 13). Data represent mean ± SD. *P < 0.05; **P < 0.01. AU, arbitrary units.

FIG. 7. RMCs promote expression of lipogenic genes. RMCs from obese HFD-fed mice induce in hepatocytes from lean mice a program that would favor fatty acid esterification (\(\square\), control; \(\blacklozenge\), RMCs). Data represent mean ± SD. *P < 0.05; ***P < 0.005. AU, arbitrary units.

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