Increased CD36 protein as a response to defective insulin signaling in macrophages

Chien-Ping Liang, Seongah Han, Haruka Okamoto, Ronald Carnemolla, Ira Tabas, Domenico Accili, and Alan R. Tall

Divisions of Molecular Medicine and Endocrinology, Department of Medicine, Columbia University, New York, New York, USA.

Accelerated atherosclerosis is a major cause of morbidity and death in insulin-resistant states such as obesity and the metabolic syndrome, but the underlying mechanisms are poorly understood. We show that macrophages from obese (ob/ob) mice have increased binding and uptake of oxidized LDL, in part due to a post-transcriptional increase in CD36 protein. Macrophages from ob/ob mice are also insulin resistant, as shown by reduced expression and signaling of insulin receptors. Three lines of evidence indicate that the increase in CD36 is caused by defective insulin signaling: (a) Treatment of wild-type macrophages with LY294002, an inhibitor of insulin signaling via PI3K, results in an increase in CD36; (b) insulin receptor knockout macrophages show a post-transcriptional increase in CD36 protein; and (c) administration of thiazolidinediones to intact ob/ob mice and ob/ob, LDL receptor–deficient mice results in a reversal of macrophage insulin receptor defects and decreases CD36 protein. The last finding contrasts with the increase in CD36 that results from treatment of macrophages with these drugs ex vivo. The results suggest that defective macrophage insulin signaling predisposes to foam cell formation and atherosclerosis in insulin-resistant states and that this is reversed in vivo by treatment with PPAR-γ activators.

Introduction

The complications of atherosclerosis in coronary, cerebral, and peripheral arteries are the major reason for hospital admission in diabetics and account for about 80% of the mortality of this condition (1). Although diabetes is often associated with dyslipidemia (2, 3), the excess risk is disproportionate to lipid abnormalities and the root causes of accelerated atherosclerosis in diabetes are not well understood. Some evidence suggests that insulin resistance, commonly seen in overweight individuals with the metabolic syndrome, is an important cause of both diabetes and increased atherosclerosis risk (4). Insulin resistance has been related to dyslipidemia, hypertension, and hypercoagulability, all factors that may promote atherosclerosis (2, 3). Potentially insulin resistance could also be important at the arterial cellular level (5). Defective insulin signaling in endothelial cells leads to impaired NO activity (6), and this is probably proatherogenic. Macrophages also express insulin receptors (IRs) (7), but the relationship of insulin signaling to macrophage foam cell formation has received little attention.

Thiazolidinediones, activators of PPAR-γ, have emerged as an important class of drugs in the treatment of type II diabetes. PPAR-γ activators markedly improve insulin resistance and decrease atherosclerosis in mouse models, and may also decrease atherosclerosis in humans (8–11). Although the in vivo mode of action of PPAR-γ activators is not well understood, PPAR-γ is highly expressed in adipocytes and macrophages (11), suggesting these cells could be important targets. Paradoxically, however, PPAR-γ activators increase the expression of the proatherogenic molecule CD36 in cultured macrophages (12). These proatherogenic effects of PPAR-γ activators may be counteracted by an increase in ABCA1, a macrophage transporter that promotes efflux of cholesterol to lipid-poor apolipoproteins, perhaps offsetting increased uptake of modified low-density lipoprotein (LDL) via CD36 (11, 13, 14). However, the importance of this mechanism is uncertain, and some studies have failed to find any effect of PPAR-γ activators on ABCA1 levels or lipid efflux in macrophages (15, 16).

Obese (ob/ob) mice are genetically deficient in leptin and have been widely used as a model of insulin resistance and diabetes (17). In this study, we made the initial observation that macrophages from ob/ob mice have increased uptake of modified LDL, due to post-transcriptional upregulation of CD36 protein. Unexpectedly, the increase in CD36 is caused by defective insulin signaling in macrophages, suggesting that macrophage insulin resistance promotes foam cell formation. Consistent with this hypothesis, in vivo treatment with PPAR-γ activators improved insulin resistance and decreased macrophage CD36 protein and uptake of oxidized LDL (oxLDL).

Methods

Animals. Obese ob/ob and lean+/+, and ob/+ and LDL receptor–deficient (LDLR−/−) on a C57BL/6J background were obtained from the Jackson Laboratory (Bar Harbor, Maine, USA). Mice deficient in both leptin and LDLR (ob/ob LDLR−/−) were generated by crossing of ob/+ and LDLR−/− mice as described (18). All experiments were performed with animals fed regular rodent chow diet, except that in the experiments of in vivo rosiglitazone treatment, ob/ob LDLR−/− and LDLR−/− mice were fed a Western-type diet (TD88137; Harlan-Teklad, Madison, Wisconsin, USA). At least three mice of each group were used in the experiments of this study. All experimental procedures were carried out in accordance with Columbia University’s institutional guidelines for use of laboratory animals.

Macrophage and monocyte isolation. Thioglycollate-elicited peritoneal macrophages from at least three ob/ob and lean−/+ mice or from transgenically rescued IR KO mice and littermate mice (19, 20) were collected by peritoneal lavage, and pooled macrophages of each strain were cultured as described (21), except that 0.2% bovine serum
albumin (BSA) was supplemented in DMEM (Invitrogen, Carlsbad, California, USA) containing 1,000 mg/l of D-glucose unless otherwise specified. After 1–2 hours, the cells were washed three times with 1× phosphate-buffered saline (PBS; Invitrogen) and were used immediately for experiments as described below. For the isolation of mouse peripheral blood monocytes, the flotation method (22) with OptiPrep density gradient medium (Sigma-Aldrich, St. Louis, Missouri, USA) was used. Blood from lean (+/−) and ob/ob mice was collected and mixed well with OptiPrep in Tricine-buffered saline (TBS; 0.85% NaCl and 10 mM Tricine-NaOH, pH 7.4). The pooled samples were centrifuged at 1,000 g for 30 minutes at room temperature. The cells were harvested from the meniscus downwards and diluted into TBS. After being spun at 400 g for 10 minutes, the cells were resuspended and cultured as described above.

Acetylated LDL (acLDL) uptake and cholesteryl ester formation. We prepared acLDL and labeled it with 125I and 3H as described (21, 23). The assays of cell association, degradation, and cholesteryl ester formation with macrophages following loading of 125I-acLDL were performed as previously described (23, 24).

Binding assay for oxLDL. 125I-labeled oxLDL was prepared as described (25). Labeled oxLDL (10–25 µg/ml or in amounts as indicated in Figure 1E) in DMEM/0.2% BSA with or without the addition of unlabeled oxLDL (40-fold excess; for estimates of nonspecific binding) were incubated for 2–3 hours at 4°C with macrophages with or without prior treatments as indicated in figure legends. In some experiments, macrophages were preincubated for 30 minutes at 4°C with fucoidan (Sigma-Aldrich), mouse anti–CD36 IgA (Cascade BioScience, Winchester, Massachusetts, USA), or control mouse IgA (Biodesign, Saco, Maine, USA) in DMEM/0.2% BSA prior to the addition of oxLDL. After cells were washed on ice three times with ice-cold PBS, cell-associated 125I radioactivity was determined as described above.

Ex vivo treatments of macrophages and monocytes. Fresh peritoneal macrophages or blood monocytes were treated at 37°C in DMEM/0.2% BSA with insulin (Sigma-Aldrich), wortmannin (Sigma-Aldrich), LY294002 (Biomol, Plymouth Meeting, Pennsylvania, USA), TNF-α (Sigma-Aldrich), oleic acid– or palmitic acid–BSA, rosiglitazone (GlaxoSmithKline, Research Triangle Park, North Carolina, USA), troglitazone (Parke-Davis, Morris Plains, New Jersey, USA), calpeptin (Calbiochem, La Jolla, California, USA), chloroquine (Sigma-Aldrich), lactacystin (Sigma-Aldrich), or corresponding vehicle controls for 1 day, unless otherwise specified; or in DMEM/10% FBS for 1–2 days; or in glucose-free DMEM/0.2% BSA with the indicated concentrations of glucose or mannitol (Sigma-Aldrich) for 1–2 days. Fresh wortmannin was added every 4–8 hours. Complexes of oleic acid or palmitic acid (Sigma-Aldrich) with BSA were prepared as described (26). After treatments, oxLDL binding assays were carried out, or total RNA and cell lysates were prepared for further analysis.

Pulse-chase labeling of macrophages. Peritoneal macrophages were incubated with methionine- and cysteine-free DMEM/0.2% BSA for 1.5 hours at 37°C. The cells were then incubated with Pro-Mix 35S cell-labeling mix (0.2 mCi/ml; Amersham, Piscataway, New Jersey, USA) for 20 minutes or 4 hours at 37°C in methionine- and cysteine-free DMEM/0.2% BSA, followed by washing of cells three times with PBS. Labeled macrophages were chased for the indicated times at 37°C in DMEM/0.2% BSA. Subsequently, cells were rinsed three times with PBS and total cell lysates were prepared as described (21) with a lysis buffer (50 mM HEPES, 150 mM NaCl, 1.5 mM MgCl2, 2.5 mM EDTA, 1% Triton X-100, and 0.1% sodium deoxycholate, pH 7.4) with protease inhibitor cocktail (Roche, Indianapolis, Indiana, USA). Equal amounts (100 µg) of total lysates were then used to immunoprecipitate 35S-labeled CD36 with mouse anti–CD36 IgA (1 µg) overnight at 4°C. Protein L-Agarose (30 µl; Santa Cruz Biotechnology, Santa Cruz, California, USA) was added, followed by incubation for another 3 hours at 4°C. After centrifugation agarose beads were washed four times with ice-cold lysis buffer without 0.1% sodium deoxycholate. Precipitated proteins were eluted from beads, separated by SDS-polyacrylamide gel electrophoresis (PAGE), electrotransferred to nitrocellulose membrane (Bio-Rad, Hercules, California, USA), and analyzed by autoradiography. The intensities of 35S-labeled CD36 were quantified by densitometric analysis (Molecular Dynamics, Piscataway, New Jersey, USA).

Internalization assay. CD36 internalization in macrophages was assessed with reversible surface protein biotinylation as described (27). Fresh macrophages were incubated with sulfo-NHS-SS-biotin (0.5 mg/ml; Pierce) for 40 minutes at 4°C in PBS with 1 mM MgCl2 and 0.5 mM CaCl2. Cells were washed three times with PBS and were then cultured in DMEM/0.2% BSA for the indicated times at 37°C, followed by surface biotin removal with or without glutathione (150 mM; Sigma-Aldrich) twice for 20 minutes at 4°C. After neutralization with iodoacetamide (50 mM; Sigma-Aldrich), macrophages were lysed with a lysis buffer and CD36 was immunoprecipitated as described above. After SDS-PAGE and membrane transfer, biotinylated CD36 was detected with streptavidin-HRP conjugate (Pierce) and quantified by densitometric analysis. Percent internalization was calculated as described (27) after the intensity of the CD36 band was corrected for input with β-actin.

Insulin-dependent tyrosine-phosphorylation assay. After 5 hours of serum starvation, macrophages were labeled with 12P (0.2 mCi/ml; Parkin-Elmer, Boston, Massachusetts, USA) for 2.5 hours at 37°C in DMEM/0.2% BSA then treated with insulin at the indicated concentrations at 37°C. After 10 minutes, cells were frozen by liquid nitrogen to rapidly quench any further phosphorylation. Total cell lysates were prepared as described above with the addition of phosphatase inhibitor cocktail (Sigma-Aldrich), and 100-µg samples were used to immunoprecipitate IRs with anti-IR IgG (Oncogene, Boston, Massachusetts, USA) or tyrosine-phosphorylated proteins with anti–phosphotyrosine IgG agarose (Santa Cruz Biotechnology). Immunoprecipitated IRs or phosphoproteins were separated by SDS-PAGE and transferred to nitrocellulose membrane. Insulin-dependent tyrosine-phosphorylated proteins were identified by autodigestion and Western analysis (see below). For the detection of tyrosine-phosphorylated IR or IR substrate 2 (IRS-2), the membrane was probed with anti–phosphotyrosine–HRP conjugate (Santa Cruz Biotechnology) or anti–IRS-2 (Upstate Biotechnology, Lake Placid, New York, USA), respectively.

In vivo treatment of mice with rosiglitazone. Obese ob/ob mice were injected intraperitoneally once daily with rosiglitazone (3 µg/g body weight) or PBS (saline control) for 17 days. For LDLR−/− and ob/ob LDLR−/− mice, animals were fed 4–5 g Western diet per mouse with or without rosiglitazone supplementation (3 µg/g body weight) each day for 3 weeks. At different times, blood was collected and blood glucose and insulin levels were measured with kits from Sigma-Aldrich and Crystal Chem Inc. (Chicago, Illinois, USA), respectively. Peritoneal macrophages from treated and control mice pre-elicited with thioglycollate were then obtained as described above and were used immediately for oxLDL binding assay or for preparation of RNA and total cell lysates.

Western analysis. Western analysis using total lysates (25–50 µg) from macrophages, monocytes, or liver was performed as described above.
(21) except that primary antibodies against CD36 (Santa Cruz Biotechnology and Cascade BioScience), scavenger receptor-A (SR-A) (TransGenic Inc., Kumamoto, Japan), scavenger receptor BI (SR-BI) (Novus Biologicals, Littleton, Colorado, USA), IR β-subunit (Santa Cruz Biotechnology), IRS-2 (Upstate Biotechnology), phosphotyrosine (Santa Cruz Biotechnology), or β-actin (Sigma-Aldrich) were used. Adipose tissue was homogenized in the lysis buffer described above and clarified lysates were prepared as described (21).

Northern analysis. Northern analysis using total RNA (5–20 µg) isolated from peritoneal macrophages was carried out as described (28). The membranes were hybridized with randomly primed 32P-labeled probes of mouse CD36, SR-A, or β-actin cDNA fragments.

Real-time quantitative PCR analysis. Total RNA was reverse transcribed at 42°C with Superscript II (Life Technologies). For CD36 and β-actin mRNA measurements, cDNA was mixed with TaqMan Universal PCR Master Mix (Perkin-Elmer Biosystems) according to the manufacturer’s protocol. The following primers were used: CD36, 5′-FAM-TCACCCCTCAGAATCCAGAACAACCA-BHQ-1-3′, forward, 5′-TCCAGCCATGCTTTCG-3′, and reverse, 5′-TGGAGTTATTTTTCAGTGAGA-3′ (FAM, 6-carboxy-fluorescein; BHQ-1, Black Hole Quencher-1); β-actin, 5′-JOE-CACTGCCG-CATCCTCTCCTCC-BHQ-1-3′, forward, 5′-AGAGGAAATGCT-GCGTGAC-3′, and reverse, 5′-CAATAGTGATGACCTGGCCGT-3′ (JOE reporter dye from Molecular Probes, Invitrogen; primers synthesized by Biosearch Technologies Inc., Novato, California, USA). Detection of specific PCR products was performed in triplicate using the Mx4000 Multiplex Quantitative PCR System (Stratagene) with 1 cycle at 50°C for 2 minute and 1 cycle at 95°C for 10 minutes followed by 45 cycles at 95°C for 15 seconds and 60°C for 1 minute. For the determination of mRNA levels of other genes, SYBR Green PCR Core Reagents (Perkin-Elmer Biosystems) were used as described by the manufacturer. Quantitative PCR was performed in triplicate using the Mx4000 Multiplex Quantitative PCR System with 1 cycle at 95°C for 10 minutes followed by 40 cycles of 90°C for 30 seconds, 58°C for 30 seconds and 72°C for 1 minute. The relative amounts of specific target amplicons for each primer set were estimated by a standard curve method using Mx4000 software (v.3.01; Stratagene) and were normalized to the copy number of mouse β-actin transcripts.

Statistical analysis. Results are expressed as means ± SE.

Results

Increased uptake of modified LDL in ob/ob macrophages associated with increased levels of CD36 and SR-A. We compared the binding and uptake of LDL modified by acetylation or oxidation in pooled primary macrophages from ob/ob and wild-type mice immediately after removing the macrophages from the peritoneal cavity. Using acLDL, there was two- to threefold increased cell association (Figure 1A), degradation (Figure 1B), and cholesteryl ester formation (Figure 1C) in ob/ob macrophages compared with wild-type macrophages. A number of different macrophage receptors recognize acLDL, including CD36, SR-A, and SR-BI. Western blotting with specific antibodies against each of these receptors revealed increases in CD36 (2.5 ± 0.4-fold) and SR-A protein (2.4 ± 0.3-fold), but reduced levels of SR-BI (0.6 ± 0.1) (Figure 1D). Measurements of mRNA by Northern analysis showed no increase in the mRNA of either SR-A or CD36 (0.8 ± 0.2) (Figure 1E).
Macrophages from ob/ob mice also showed increased binding of oxLDL (Figure 1E), consistent with increased levels of SR-A and CD36. To assess the relative importance of SR-A and CD36 in binding of oxLDL, macrophages were incubated with fucoidan, an inhibitor of SR-A-dependent uptake (29), or with CD36 monoclonal antibodies (24, 25). Figure 1F shows that binding attributable to both CD36 and SR-A was increased in mice versus wild-type mice, with the larger effect due to CD36. Control isotype-matched antibodies did not reduce binding of oxLDL (data not shown). These studies reveal an increase in binding, uptake, and degradation of modified LDL in ob/ob macrophages, primarily reflecting a post-transcriptional increase in CD36 protein.

Increased cholesteryl ester formation (Figure 1C) indicates a propensity for foam cell formation in ob/ob macrophages and could reflect both increased uptake of modified LDL and potentially decreased cholesterol efflux. However, despite the reduced levels of SR-B1, cholesterol efflux to apoA-I or HDL was similar in ob/ob and wild-type macrophages (data not shown), suggesting no significant defect in efflux. Thus, the major focus of subsequent studies was to elucidate the mechanism of the post-transcriptional increase in CD36 protein in ob/ob macrophages.

Increased CD36 results from decreased protein turnover. For investigation of the mechanism of increased CD36 protein expression, metabolic turnover (pulse-chase) studies of CD36 were performed in macrophages from ob/ob and wild-type mice. Short-term pulse labeling studies showed that the synthesis of CD36 protein was slightly reduced in ob/ob macrophages compared with wild-type macrophages (Figure 2A). In contrast, CD36 protein turnover was markedly delayed in ob/ob macrophages (Figure 2B), indicating that the accumulation of CD36 protein in these cells reflects decreased catabolism. The half-lives for CD36 protein in wild-type and ob/ob macrophages were approximately 16 and 33 hours, respectively.

Decreased turnover of CD36 in ob/ob macrophages could reflect either decreased internalization from the cell surface or decreased degradation of CD36 after internalization. We used the approach of cell surface reversible biotinylation (27) with nonpermeant glutathione to measure the rate of internalization of CD36. The sample at 0 hours confirmed an increase in cell surface CD36 in ob/ob macrophages, as shown by biotinylation (Figure 3A, glutathione –). With this approach, cell surface biotinylated CD36 becomes pro-

Figure 2
CD36 protein turnover is decreased in ob/ob macrophages. Pooled WT or ob/ob macrophages were pulse-labeled with [35S]methionine/cysteine cell-labeling mix for 20 minutes (A) or 4 hours (B), then chased in medium with methionine/cysteine for the times indicated. [35S]-labeled CD36 in total lysates was immunoprecipitated with anti-CD36 and subjected to SDS-PAGE followed by transfer to nitrocellulose membrane and autoradiography. [35S] intensities were quantified by densitometric analysis, and each number was normalized for input (measured by Western analysis of β-actin in lysates), then to the relative intensity at chase time 0 hours. For 20-minute and 4-hour labeling experiments, one experiment representative of two and four independent experiments, respectively, is shown.

Macrophages were isolated from five to seven mice of each strain.

Figure 3
CD36 is internalized similarly in WT and ob/ob macrophages and is increased in lysosome- and proteasome-inhibited WT but not in ob/ob macrophages. (A) CD36 in ob/ob macrophages is endocytosed at a rate similar to that of WT cells. Cell surface proteins were biotinylated in pooled macrophages of five mice of each strain. One experiment representative of two independent experiments is shown. Biotinylated CD36 was allowed to internalize for different times, as indicated. Surface biotin was then removed with glutathione (+). Cells without glutathione (–) were used for the measurement of total biotinylated CD36 left in the cells at the time point. The total biotinylated CD36 at time 0 was used as a reference (100%) for the quantification of internalized CD36 (all corrected for input with β-actin). Percent internalization was calculated as described (27). (B) CD36 protein is increased by lysosomal and proteasomal inhibitors in WT but not in ob/ob macrophages. Pooled macrophages isolated from five mice of each strain were treated with calpeptin (30 μg/ml), chloroquine (50 μM), or lactacystin (1 μM) for 1 day. Total lysates were prepared, and Western analysis was performed. One experiment representative of three independent experiments is shown.
ob/ob macrophages, the dose-response curve was shifted to the right, as shown by markedly reduced signaling at 1 and 10 nM insulin (Figure 5B; plasma insulin concentrations are about 4 nM in ob/ob mice). To assess downstream signaling events, we measured phosphorylation of IRS-2 (IRS-1 was undetectable in macrophages) and found phosphorylation was markedly reduced (Figure 5C); furthermore, the dose-response of IRS-2 phosphorylation was also shifted to the right in ob/ob macrophages (data not shown). These results indicate reduced IR phosphorylation and downstream signaling in ob/ob macrophages.

Defective insulin signaling in macrophages leads to increased CD36. The finding of reduced levels and tyrosine phosphorylation of both IRSs and IRS-2 indicates defective insulin signaling in ob/ob macrophages, resulting from increased CD36.

**Figure 4**

**A** Short-term treatment with insulin or PI3 kinase inhibitors, or long-term treatment with glucose or fatty acids, does not increase CD36 protein expression in WT macrophages. (A–D) Pooled WT macrophages (A, B, and D) or WT and ob/ob monocytes (C) isolated from three to four mice were incubated with insulin (1 µM) for 0.5 hours or with the PI3 kinase inhibitors wortmannin (200 nM) or LY294002 (20 µM) for 2 hours (A), with glucose or mannitol at the indicated concentrations for 2 days (B), with 400 mg/dl of glucose or mannitol for 1 day (C), or with oleic acid– or palmitic acid–BSA complexes for 1 day (D). Troglitazone (TRG; 1 µM) was used as a positive control. Treated cells were then used for oxLDL binding assays with the SR-A ligand fucoidan (50 µg/ml) added to the binding buffer or for total protein extraction followed by Western analysis. One experiment representative of two (A, B, and D) or three (B) independent experiments is shown. M0, macrophage; FA, fatty acid; Conc., concentration.
phages. We used three different approaches to test the hypothesis that defective insulin signaling could lead to increased CD36 protein in macrophages. In the first, we exposed macrophages to long-term treatment (24 hours) with high-dose insulin (100 nM), a procedure that is known to cause decreased IRs and defective signaling in other cell types (30). This treatment reduced IR levels (0.5 ± 0.2), increased CD36 protein (1.5- to 2-fold), and increased binding of oxLDL (1.5- to 2-fold) in wild-type but not \( \text{ob/ob} \) macrophages (Figure 6A). The lack of response in \( \text{ob/ob} \) macrophages reflects the fact that IRs are already markedly depleted (Figure 5A) and is consistent with the idea that downregulation of IRs is the major mechanism that accounts for the increase in CD36.

PI3 kinase is a major downstream effector of IR signaling via either IRS-1 or IRS-2 (31). Thus, in the second approach we treated macrophages with wortmannin or LY294002, two different inhibitors of PI3 kinase. Both treatments resulted in a dramatic induction of CD36 protein (two- to fivefold, depending on the doses of inhibitors used) in wild-type but no \( \text{ob/ob} \) macrophages and a...
parallel increase in binding of oxLDL to wild-type macrophages (Figure 6B). In contrast, CD36 mRNA was minimally induced by PI3 kinase inhibition (1.2- to 1.5-fold; data not shown). TNF-α, another factor that may induce insulin resistance in mice (32), did not result in increased CD36 protein (Figure 6C). When ob/ob macrophages were cultured for 24 or 48 hours in FBS, there was partial or complete reversal in the defective expression of IRS and CD36, respectively (Figure 6D), and a partial reversal in CD36 protein turnover (data not shown), suggesting that the defect in ob/ob macrophages was not cell autonomous. These results are consistent with the idea of defective IR signaling as the underlying cause of increased CD36 in ob/ob macrophages.

In a third genetic approach, we used macrophages from transgenically rescued IR KO mice. These mice express normal IRSs exclusively in the liver, islets, and brain, and are not diabetic (19, 20). Macrophages from these mice lacked IRs, showed normal IRS-2 (Figure 7A), and showed increased binding of oxLDL and increased CD36 (2.8 ± 0.6-fold) and SRA protein (3.6 ± 1.1-fold) but not mRNA (Figure 7), similar to ob/ob mice. Treatments with fucoidan and anti-CD36 indicated that increased CD36 in IR KO mice was the major cause of increased binding of oxLDL (Figure 7C). Thus, genetic deficiency of IRs in macrophages leads to the same defect in CD36 protein expression as genetic obesity, consistent with defective insulin signaling through the macrophage IR as the major underlying mechanism.

PPAR-γ activators decrease macrophage CD36 protein in vivo but not ex vivo. If insulin resistance is the underlying reason for increased CD36 in ob/ob macrophages, then reversal of insulin resistance in vivo by treatment with thiazolidinediones should lead to a reversal of the macrophage phenotype. Treatment of ob/ob mice with rosvitiglazone (RSG) improves insulin resistance and normalizes oxLDL binding and CD36 expression in macrophages. (A) In vivo rosvitiglazone (RSG) treatment reduces plasma glucose levels and oxLDL binding to macrophages in ob/ob mice. Plasma glucose levels (left) and macrophage oxLDL binding (right) were assessed in ob/ob mice treated with or without RSG or control saline. At the times indicated, plasma glucose levels from these mice were measured. After 3 weeks, macrophages were collected and an oxLDL binding assay was performed with the addition of fucoidan (50 µg/ml) in the binding buffer. (B) Differential patterns of CD36 protein expression in ob/ob macrophages in response to thiazolidinedione treatments in vivo and ex vivo. Left, Western analysis of CD36 protein expression in ob/ob macrophages in response to thiazolidinedione treatments in vivo and ex vivo. Left, Western analysis of CD36 protein expression in ob/ob and WT macrophages treated ex vivo with RSG or troglitazone (TRG) at the indicated doses for 1 day. Right, the expression of CD36 and IR β-subunit (IRβ) protein in macrophages from ob/ob mice described in A was determined by Western analysis. Macrophage CD36 protein was normalized in ob/ob mice, though its mRNA was increased by rosvitiglazone in vivo, as measured by Northern analysis. Macrophage experiments were performed with pooled cells isolated from three to five mice of each strain. One experiment representative of two independent experiments is shown.
rosiglitazone, a PPAR-γ activator, resulted in a marked reduction of blood glucose levels (threelfold; Figure 8A) and of plasma insulin levels (data not shown), indicative of decreased insulin resistance in these animals (33). This was associated with a reversal of the macrophage phenotype with upregulation of IRs (Figure 8B), decreased binding of oxLDL (Figure 8A), and decreased CD36 protein but not CD36 mRNA (CD36 mRNA was actually increased; Figure 8B). In contrast, ex vivo treatment of ob/ob or wild-type macrophages with thiazolidinediones resulted in two- to fourfold increased CD36 protein (Figure 8B), as reported (12). These results suggest that in vivo improvement of insulin resistance by PPAR-γ activators led to decreased macrophage CD36 expression, predominating over any possible increase in CD36 related to direct effects of PPAR-γ activators in the macrophage.

To assess the potential relevance of these observations to atherogenesis, similar experiments were conducted in LDLR−/− and ob/ob LDLR−/− mice, since the latter show markedly increased atherosclerosis compared with the former (18). As shown in Figure 9A, this confirmed a reduction in CD36 (40% decrease) and SR-A protein (60% decrease), along with decreased plasma glucose and insulin levels (data not shown), in the ob/ob LDLR−/− mice; the effect was not as dramatic as that shown in Figure 8, probably because we had to use a reduced dose of rosiglitazone due to toxicity at the higher dose in ob/ob LDLR−/− mice. In contrast, in the control LDLR−/− mice, CD36 protein was induced threefold by rosiglitazone treatment (Figure 9A). This is consistent with a reversal of obesity-associated insulin resistance as the major mechanism responsible for decreased CD36 in the ob/ob LDLR−/− mice. In contrast to the response in macrophages, both CD36 mRNA (three- to fourfold; data not shown) and protein (two- to fourfold; Figure 9B) were increased in adipose tissue from rosiglitazone-treated ob/ob LDLR−/− mice, indicating a differential response in macrophages and adipose tissue.

Discussion

This study documents defective insulin signaling in macrophages from ob/ob mice and provides evidence that this leads to a post-transcriptional increase in CD36. As CD36 is a key molecule in the recognition of modified LDL (34), our findings suggest that insulin resistance at the level of the macrophage predisposes to atherosclerosis and represents a new mechanism to explain increased atherogenesis in insulin resistant states such as metabolic syndrome and type II diabetes. Moreover, this abnormality is reversed by treatment of ob/ob and ob/ob LDLR−/− mice with PPAR-γ activators, providing an explanation for the known antithrombogenic effects of these agents that is closely linked to their main mechanism of action; that is, improvements in insulin signaling (8, 9).

Surprisingly, there is only sparse evidence in the literature indicating the presence of IRs and insulin signaling in macrophages. Older studies showing high-affinity binding of insulin to monocytes and macrophages implied the existence of such receptors (7); moreover, reduced insulin binding to monocytes of obese people has been documented (35), suggesting a depletion of IRs similar to that observed in ob/ob mice in this study. However, there is a relative paucity of information on macrophage insulin signaling and its relationship to normal or abnormal macrophage functions (36–38), in contrast to the detailed studies carried out with other insulin-sensitive tissues (31). Our studies revealed abundant IRs and insulin signaling via IRS-2 in normal macrophages. The physiological significance of insulin signaling remains unclear, as the constitutive GLUT-1 glucose transporter, rather than the insulin-inducible GLUT-4 (39, 40), is the main isofrom expressed in macrophages, and insulin had no direct effect on glucose uptake by mouse macrophages (data not shown). While a role for insulin signaling in macrophage phagocytosis could be suspected, studies of insulin effects on this process have yielded inconsistent results (7, 41).

Our studies link defective insulin signaling in macrophages to an increase in CD36 protein. Part of the evidence for this was that IR-deficient macrophages obtained from IR-transgenic/ KO mice also displayed increased CD36 protein, but reduced CD36 mRNA, similar to ob/ob macrophages. However, like ob/ob mice, the IR KO animals have insulin resistance, so these findings alone do not rule out the possibility that the increase in CD36 was secondary to insulin resistance in the whole animal rather than a direct effect of genetic deficiency of IRs in macrophages. However, the downregulation of IRs by high-dose insulin treatment or treatment with PI3 kinase inhibitors in cultured normal macrophages also led to increased CD36 protein, providing further evidence that CD36 is increased as a direct result of decreased insulin signaling in the macrophage. Also consistent with this hypothesis, a number of alternative explanations, such as direct effects of leptin, glucose, or fatty acids on macrophages, were excluded in cell culture experiments. IR-transgenic/ KO mice expressing human IRs only in liver, islets, and brain do not have hyperglycemia (20) and their plasma free fatty acids are normal (data not shown), providing additional evidence to support the direct ex vivo evidence that increased glucose and fatty acids did not lead to an increase in CD36 protein in macrophages.

The mechanism responsible for decreased IRs and signaling in ob/ob mice is unknown but could reflect changes in humoral factors associated with insulin resistance, such as hyperinsulinemia, and altered levels of plasma factors such as adiponectin (42–44) or resistin (45). Surprisingly, insulin resistance led to a delay in the cellular catabolism of CD36 protein. Studies with inhibitors suggest a defect in degradation of CD36 in lysosomes and proteasomes of ob/ob
macrophages. As this is associated with an increase in cell surface CD36 that does not reflect a defect in internalization, increased recycling of CD36 to the cell surface is implied. Thus, a defect in insulin signaling appears to lead to a defect in receptor trafficking. The resultant buildup of CD36 protein leads to increased binding, uptake, and degradation of modified LDL in ob/ob macrophages (Figure 1). Presumably the modified LDL dissociates from CD36 in early endosomes following internalization and enters lysosomal degradation pathways, while CD36 itself undergoes recycling.

In an earlier study, Griffin et al. (46) reported a post-transcriptional increase in CD36 in diabetic macrophages, similar to our findings. However, this effect was attributed to increased CD36 protein translation, related to hyperglycemia. For unknown reasons, we could not confirm such an effect of increased glucose levels on CD36 protein expression in mouse peritoneal macrophages or blood monocytes. Subsequent studies have not confirmed the increase in CD36 expression ex vivo by glucose in human blood monocytes (47). Moreover, broader considerations call into question the link between hyperglycemia and atherosclerosis. Hyperglycemia is not well correlated with atherosclerosis risk in humans. For example, in the UK Prospective Diabetes Trial, higher levels of hemoglobin A1C levels, reflecting poorer control of hyperglycemia, were associated with a much higher incidence of microvascular disease but only slightly increased macrovascular disease (3, 4). In the majority of studies in mouse models of atherosclerosis, hyperglycemia per se does not increase atherosclerosis (48, 49). Thus, we have made an important advance on the original observation of increased CD36 in diabetic macrophages by linking this phenotype to defective macrophage insulin signaling. Our findings may be relevant to the increased atherogenic risk in metabolic syndrome, a highly prevalent condition associated with insulin resistance in the absence of frank hyperglycemia.

CD36 has emerged as a key proatherogenic molecule (34, 50), potentially mediating binding of LDL containing as few as two or three appropriately modified phospholipid molecules (51). Mice deficient in both CD36 and apoE show reduced atherosclerosis compared with apoE KO controls (50). Although not the primary focus of the present study, ob/ob and IR-deficient mice also showed upregulation of macrophage scavenger receptor A via post-transcriptional mechanisms; this could also predispose to atherosclerosis. SR-A recognizes only LDL that has been extensively modified, and studies with SR-A KO mice have indicated somewhat variable atherogenic results (52). There appear to be multiple pathways by which macrophages take up modified forms of LDL, including the uptake of aggregated LDL produced by arterial sphingomyelinase or other factors. These pathways may involve engulfment and prolonged cell surface contact with immobilized, aggregated LDL, and require remodeling of the actin cytoskeleton, a process known to be regulated by PI3 kinase signaling (24). Such pathways could also potentially be regulated by changes in PI3 kinase activity associated with macrophage insulin resistance.

PPAR-γ activators have emerged as an important treatment of type II diabetes. Since the discovery that PPAR-γ activators, including hydroxycotadecadienoic acids present in oxLDL, increase transcription of CD36, there has been concern that PPAR-γ activators could worsen atherosclerosis. This concern has only been partly allayed by mouse atherosclerosis studies (8–11) and a single small human study suggesting benefit on carotid intimal thickening (53). One potential beneficial effect of PPAR-γ activators in macrophages is a decrease in inflammatory responses (54); indeed, these drugs cause a widespread decrease in expression of macrophage inflammatory genes (55). Our studies indicate an additional possibly indirect effect of these agents; that is, a decrease in insulin resistance that leads to an improvement in insulin signaling and a decrease in macrophage CD36. However, this beneficial effect was seen only in a model of severe insulin resistance, that is, the ob/ob mouse, whereas LDLR-deficient mice fed a Western diet that have milder insulin resistance actually showed an increase in CD36 protein after rosiglitazone treatment (Figure 9A). This suggests that direct effects of thiazolidinedione on CD36 gene expression may predominate in the setting of mild insulin resistance. Thus, the effects of these drugs on atherogenesis could differ depending on the degree of insulin resistance. Our findings focus attention on macrophage insulin signaling as a key parameter potentially determining atherogenic responses and suggest that further studies of this pathway may be rewarding for understanding of both the normal functions of macrophages and also defects in insulin signaling that could contribute to impairment of innate immunity and to atherosclerosis in metabolic syndrome and diabetes.

Acknowledgments

We thank Wengen Chen for help in cholesterol efflux experiments. This work was supported by NIH grants HL22682 and DK58282.

Received for publication July 18, 2003, and accepted in revised form December 15, 2003.

Address correspondence to: Chien-Ping Liang, Division of Molecular Medicine, Department of Medicine, Columbia University, 630 West 168th Street, New York, New York 10032, USA. Phone: (212) 305-4899; Fax: (212) 305-5052; E-mail: CL534@columbia.edu.