Tight Association between Macrophages and Adipocytes in Obesity: Implications for Adipocyte Preparation

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Objective: To determine the cellular architecture of the inflammatory infiltrate in adipose tissue from obese mice, and identify the source of inflammatory cytokines in adipose tissue at a single cell level.

Methods: Adipose tissue from diet-induced obese mice was digested by collagenase treatment and fractionated by density centrifugation to obtain an adipocyte floating layer and a pellet of stromal vascular cells. The cellular architecture of the adipocyte-macrophage interaction in both intact white adipose tissue (WAT) and the separated density gradient floating layer fraction was analyzed by confocal immunohistochemistry. Cytokine expression was detected by semi-quantitative real time PCR and immunohistochemical analysis.

Results: Three dimensional image analysis of WAT and the separated “adipocyte” floating layer revealed lipid-engorged macrophages, macrophages in contact with lipid droplets and sheath-like assemblies of macrophages surrounding adipocytes. The macrophages immunostained for TNFα and to a lesser extent for the immunoregulatory cytokine IL-10. TNFα staining was associated only with macrophages indicating that macrophages and not adipocytes are the source of TNFα expression in the adipocyte floating layer.

Conclusion: Macrophages form assemblies that tightly adhere to and cover adipocytes and lipid droplets. TNFα found in low density adipocyte preparations is due to contamination with macrophages.

Introduction

In obesity, adipose tissue is in a state of chronic low-grade inflammation (1,2) with elevated numbers of immune cells, including macrophages and T cells, and dysregulated expression of cytokines/chemokines [reviewed in Refs. (3,4)]. The inflammatory components have direct and indirect effects on metabolic pathways and insulin sensitivity. Of note, TNFα causes insulin resistance, at least in part by blunting insulin signaling (5). Thus, it is important to understand the cellular architecture and interactions within this specific microenvironment.

The cellular infiltrate in adipose tissue from obese animals is organized in cellular assemblies known as crown-like structures (CLSs) and contains a large number of macrophages (6). CLS macrophages are found predominantly around dying adipocytes, and appear to engulf adipocyte-derived lipid fragmented into small droplets (7). Mechanistic details of these events are still unknown as are the signals that orchestrate these events. To address these questions further, more information is required about the architecture of the cellular infiltrate surrounding adipocytes.

In an attempt to identify the cellular source of TNFα, Hotamisligil et al. (5) separated adipocytes from non-adipocytes based on buoyancy and found that the floating cells, presumed to contain adipocytes, expressed more TNFα mRNA than stromal vascular cells (SVCs) found in the pellet. Based on a carefully prepared adipocyte fraction, Fain et al. (8) concluded that non-adipocytes, most likely macrophages, were the major source of TNFα in human adipose tissue. The work was also supported by grants from the NIH; R01 DK054254, R01 DK083850, and R01 HL112248 to SMN.

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tissue. Although the majority of studies suggest that macrophages are the main source of TNFα in adipose tissue, adipocytes are still considered candidate producers of TNFα (6,9). A definitive identification of the source of TNFα and other cytokines requires an analysis at the single cell level and a more detailed analysis of the cellular architecture of the adipose tissue inflammatory infiltrate.

In this study, whole adipose tissue as well as density fractionated tissue preparations (10) from obese mice were analyzed by confocal immunohistochemistry to generate 3D images of the adipose tissue inflammatory infiltrate. This shows that buoyancy alone does not separate adipocytes from other types of cells. Adipocyte-associated macrophages, and not the adipocytes themselves, stain positive for TNFα. Previous reports, reviewed in Ref. (11), of separated adipocytes producing TNFα may be indicative of the inability to separate adipocytes from macrophages even after digestion and separation into single adipocytes.

Methods
Mice and Diets
Ceacam1 null mice (Cc1<sup>−/−</sup>), genetically predisposed to developing obesity and metabolic syndrome, were propagated on a C57BL/6 genetic background (12,13). C57BL/6 (The Jackson Laboratory, Bar Harbor, ME) and Cc1<sup>−/−</sup> male mice were bred in house and used throughout the study. Mice had free access to water and one of three diets; regular diet (RD) (Lab Diets #5001), high-fat diet (HF), or Western high-fat/high carbohydrate diet (HF/HC) (Research Diets Inc., New Brunswick, NJ, #D12451 and D12079B, respectively). Feeding began at 2 months of age and continued for 1-4.5 months. Weight measurements were taken at the initiation of feeding and prior to sacrifice and tissue harvesting. Visceral adiposity was calculated as the percentage of peri-gonadal visceral white adipose tissue (WAT) mass relative to body weight after overnight fasting. Both strains of mice were considered together for analysis and divided into: non-obese mice fed a RD diet with less than 3.5% visceral adiposity and obese mice fed a HF or HF/HC diet with 5-7% visceral adiposity. All procedures were approved by the Institutional Animal Care and Use Committee.

Tissue Histology and Adipocyte Cell Measurement
Small pieces of adipose tissue were removed and fixed for >24 hours in Z-fix buffered zinc formalin fixative (Anatech, Battle Creek, MI). Tissue was processed, paraffin-embedded, and cut into 5 µm sections and H&E stained.

RNA Isolation and Semi-Quantitative Real-Time PCR Analysis
Adipose tissue and isolated cell populations were homogenized using one-time-use generators and a Powergen 125 tissue homogenizer (ThermoFisher Scientific, Pittsburg, PA). Frozen tissue samples (30-50 mg) were stored in RNAlter (Qiagen, Valencia, CA). Messenger RNA (mRNA) was isolated and purified using RNeasy mini kits (Qiagen) according to the manufacturer’s protocol. cDNA was synthesized using M-MLV Reverse Transcriptase (Thermo-Fisher Scientific) and was used in triplicate semi-quantitative real-time PCR reactions (qRT-PCR) to measure the relative amount of mRNAs. Real time amplification was obtained via Absolute qRT-PCR SYBR mastermix and a CFX96 system Thermocycler (Bio-Rad, Hercules, CA) used per manufacturer’s instructions. Forward and reverse primers were designed using the Primer Express 1.5 software (Applied Biosystems, Foster City, CA) or identified using PrimerBank (http://pga.mgh.harvard.edu/primerbank/, Harvard Medical School). Glyceraldehyde-3 phosphate dehydrogenase (GAPDH) was used as an endogenous control to normalize the amount of starting cDNA. The delta Ct quantification method was employed. Mouse primer sequences (forward/reverse) were as follows:

GAPDH: CCAAGTTGTCTCTCTGCAGCT/ATACCAGGAAATAGGCTTACAAAGT; TNFα: CATCCTCTCCTAACATTGAGTACAA/TTGGGAGTACACAGGGTACACCC; IL-10: GGTTGCCAGCGTTATCGGA/ACCTGCTCCACCTGCTTGCT; F4/80: CTTTGCTATGGCGCTCCAGTC/CAAGGGACAGGTTATACGTTG; Adiponectin: AGCCGCTTATATGTATCGCTC/TGGCGTCAGAGTTTATCGTG; Ceacam1 C. Further digestion time was used as an endogenous control to normalize the amount of starting cDNA. The delta Ct quantification method was employed. Mouse primer sequences (forward/reverse) were as follows:

GAPDH: CCAAGTTGTCTCTCTGCAGCT/ATACCAGGAAATAGGCTTACAAAGT; TNFα: CATCCTCTCCTAACATTGAGTACAA/TTGGGAGTACACAGGGTACACCC; IL-10: GGTTGCCAGCGTTATCGGA/ACCTGCTCCACCTGCTTGCT; F4/80: CTTTGCTATGGCGCTCCAGTC/CAAGGGACAGGTTATACGTTG; Adiponectin: AGCCGCTTATATGTATCGCTC/TGGCGTCAGAGTTTATCGTG; Ceacam1

Adipose Tissue Separation
SCVs and adipocyte fractions were isolated from adipose tissue following a method modified from Lumeng et al. (10). One gram of visceral adipose tissue from each mouse was pooled, finely minced, and suspended in PBS containing magnesium and calcium, 1.5% fatty acid and endotoxin free bovine serum albumin (BSA), 5 mM glucose, and 100 Units of penicillin and 100 μg streptomycin. Minced tissue was centrifuged at 500 rcf for 5 minutes to pellet red blood cells (RBC). Collagenase (≥125 U/mg, Sigma, St. Louis MO) was added to a final concentration of 1.0 mg/ml, and the tissue was routinely digested in a shaking water bath (200 Hz) for 45-50 minutes at 37°C. Further digestion time resulted in lysed adipocytes. Undigested material was removed by straining through a 100 µm sieve (BD Biosciences, San Jose, CA) and SVC were pelleted by centrifugation at 500rcf for 5 minutes. Remaining RBCs in the SVC fraction were lysed with erythrocyte lysing buffer (Lonza, Walkersville, MD) for 5 minutes at room temperature. Floating adipocytes were removed and washed with PBS three times. Purified cell fractions were either lysed for mRNA isolation and further analyzed by qRT-PCR, or analyzed by fluorescence activated cell sorter (FACS) or laser scanning confocal microscopy (LSCM).

Magnetic Separation of Adipose Tissue
Macrophages from SVC
Macrophages within the SVC pellet were purified using anti-F4/80-PE antibody 2.0 µg/ml (eBioscience, San Diego, CA) and EasySep PE selection kit (Stemcell Technologies, Vancouver, BC) per manufacturer’s protocol. Purity of the separated cell population was >95% as confirmed by FACS analysis of F4/80<sup>+</sup> cells versus total SVC using a BD FACSscan and CELLquest software (BD Biosciences, San Jose, CA) (FACS data not shown).

Fixation and Antibody Staining of Visceral WAT and Floating Layer Adipocytes
Adipose tissue for microscopy was harvested in 2-3 mm pieces and fixed using Z-fix formalin (Anatech) for 12 hours before being

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stored in PBS at 4°C. Adipose tissue and floating layer cells were permeabilized in 1% Triton X-100 for 10 minutes. For simple staining, fixed adipose tissue and floating layer cells were stained with rat anti-mouse F4/80 (Invitrogen, Carlsbad, CA) to mark macrophages and detected with donkey anti-rat IgG conjugated to Alexa Fluor488 (Invitrogen) or IL-10 was stained with anti-mouse IL-10 biotin conjugate (Invitrogen) and an Alexa647 conjugated streptavidin (Invitrogen). Tissues were incubated with primary antibodies overnight then washed with staining buffer before application of secondary stains. Secondary staining reagents were applied for 2 hours then washed three times in PBS before imaging. For complex five-color staining, fixed tissue and floating layer adipocytes were stained utilizing directly conjugated fluorescent anti-mouse monoclonal antibodies: anti-TNFα conjugated FITC (eBioscience), anti-IL-10 conjugated PE (BD Bioscience) and anti-F4/80 conjugated Alexa Fluor647 (Invitrogen) in PBS staining buffer with 1% BSA. Antibodies were incubated with samples overnight at room temperature with rocking. All stained samples were then counterstained for 15 minutes at room temperature with 5 μM BODIPY 558/568 (Molecular Probes, Inc. Eugene, OR) to visualize lipid and/or 40 μM Hoechst 33342 (Invitrogen) to visualize nuclei. All samples were washed three times with PBS before visualization. Adipose tissue samples were placed directly on a coverslip with buffer and visualized. For the floating layer samples, the aqueous layer was removed down to 20 μl of sample which was then mixed with 50 μl Fluormount-G (Southern Biotech, Birmingham, AL) and mounted using a coverslip.

Laser Scanning Confocal Microscopy

Samples were imaged using a Leica TCS SP5 laser scanning microscope (Leica Microsystems, Bannockburn, IL) equipped with conventional solid state and a Ti-sapphire tunable multi-photon laser (Coherent, Santa Clara, CA). Images were acquired in the 3D XYZ plane in 1-5 μm steps with a 20X objective (NA 0.70) or 63X objective (NA 1.40) using the sequential scan mode to eliminate any spectral overlap in the individual fluorophores. Specifically, FITC was excited at 488 nm with collection at 500-558 nm. The BODIPY 558/568 dye (Molecular Probes, Inc.) was excited at 561 nm and collected at 567-609 nm. AlexaFluor647 (Invitrogen) was excited at 633 nm and collected at 644-713 nm. The Hoechst dye was excited with the multi-photon (MP) laser tuned to 790 nm with collection at 420-500 nm. Selected images are a 2D representation of the 3D LSCM image stack as labeled.

Statistical Analysis

The two-way ANOVA comparisons and Bonferroni post-tests were performed using GraphPad Prism 5.0 (GraphPad Software, La Jolla,
CA). Additionally, direct comparisons were made utilizing a t-test via Microsoft Excel (Microsoft Corp., Redmond, WA).

**Results**

**Macrophages Appear in CLS with Obesity and Take Up Lipid**

Hematoxylin and eosin staining of visceral WAT from obese mice displayed cell accumulation around adipocytes known as CLS (Figure 1a), while non-obese mice remained CLS-free. Adipocyte size increased in obese mice compared to non-obese mice (Figure 1a) consistent with previous observations (14).

LSCM analysis of the adipose tissue from obese animals showed that macrophages are a major component of the CLS as determined by F4/80 staining (Figure 1b) consistent with previous publications (15,16). The lipid staining inside some of the CLS cells clearly demonstrated that macrophages have lipid in their cytoplasm (Figure 1c) Structures such as these can be seen in Nishimura et al. (16), but with CD68 staining.

**WAT from Obese Mice Contained Macrophages, but not Adipocytes, that Expressed TNFα**

Figure 2 shows a CLS containing macrophages. Some macrophages stained only with F4/80, without TNFα or IL-10 (Figure 2; compare F4/80 with overlay). The majority of F4/80 positive macrophages however, also stained for TNFα (Figure 2; compare F4/80 with TNFα and overlay) and occasionally macrophage areas gave distinctive staining for F4/80 and TNFα as well as IL-10 (Figure 2 overlay; arrows). Overall, most macrophages (>11) expressed TNFα while a minority (~2) expressed both TNFα and IL-10. None of the macrophages, within the CLS shown in Figure 2, produced IL-10 alone.

Previous reports have suggested that adipocytes are a source of TNFα (6,9,11,17). Figure 3a and b demonstrate that adipocytes did not express TNFα even under obese conditions. The only TNFα expression is associated with F4/80 positive macrophages as shown by co-localization (Figure 3b overlay; arrows).

**Separated Floating Layer “Adipocytes” from Obese Mice were Persistently Associated with Macrophages Markers**

We sought to further clarify the cellular origin of TNFα and IL-10 cytokines from obese visceral adipose tissue by utilizing adipose tissue digestion and cellular separation. Adipose tissue was separated as described (10,14) into a floating adipocyte layer and a pelleted SVC layer known to contain multiple types of cells, including macrophages (18).

Only the floating adipocyte layer in both non-obese and obese mice showed expression of the adipocyte marker adiponectin (19,20), whereas the SVC fraction was essentially negative (Figure 4a). Surprisingly, expression of the macrophage markers F4/80 and CD68 was significantly higher in the adipocyte floating layer compared to the SVC layer of separated obese adipose tissue (Figure 4b). This is the opposite of what is expected if the SVC layer in obese mice contained all the macrophages. There was little expression of either F4/80 or CD68 in the non-obese adipocyte floating layer or SVC cell preparations (Figure 4b). When macrophages were purified from the SVC fraction there was a significant increase of F4/80 expression in obese mice compared to non-obese mice (Figure 4b). This is
unlikely to be caused by adipocyte expression of F4/80, as mature cultured 3T3-L1 adipocytes were tested by qRT-PCR and were found not to express F4/80 or other macrophage markers (data not shown). In the floating layer and in the SVC fraction, the expression of F4/80 and CD68 was higher in obese compared to the non-obese mice. Even after collagenase digestion and washing, a significant number of macrophages remained in the floating layer prepared from obese mice as determined by F4/80 and CD68 markers.

Macrophages Remained Adherent to Single Adipocytes or Lipid Droplets in the “Adipocyte” Floating Layer

The qRT-PCR experiments described above suggested that the floating adipocyte layer contained a significant number of macrophages. To determine how macrophages interact with adipocytes we stained the floating layer cells and analyzed them by LSCM. Adipocytes in the adipocyte floating layer were digested to single adipocytes (Figure 5a). Macrophages, as indicated by F4/80 staining, remained adherent to and even appear to surround several isolated single adipocyte cells obtained from obese mice (Figure 5a). This was not observed in non-obese mice. Using 63X magnification the distinctive oval shaped adipocyte nucleus is highly visible as indicated by white arrows (Figure 5b). The two macrophage nuclei stain more densely and are indicated by the blue arrow (Hoechst dye). In Figure 5b (obese panel), the F4/80 staining very brightly close to the macrophage nuclei but becomes fainter as the macrophage “spreads out” over the adipocyte. The adipocyte nucleus is visible underneath the macrophage cover. The obese adipocyte with attached macrophages was analyzed further in Figure 5c. This single section in the z-stack displays the F4/80 and lipid staining and shows two separate cells one to the left (the full cell is not shown) and the one in the center. The cell in the center is segmented with one segment covered by macrophages and being pinched off from the remaining segment not covered by macrophages. The segmented structure is clearly a single cell since only one adipocyte nucleus is seen. Both segments are connected by a bridge of cytoplasm indicated by the yellow arrow in Figure 5c. In contrast, the cell to the left is clearly separate and surrounded by F4/80+ macrophages (red arrow). The macrophages surrounding the cell in the center appear to be pinching off part of the adipocyte.

Similar structures can be seen in Figure 6b. As the 2D projection of macrophage-adipocyte interactions shows (Figure 6b), an oval nucleus belonging to the adipocyte is under a sheath of macrophages, which partially cover the adipocyte (Figure 6b; white arrows in the image stack overlay and single horizontal section right hand side). The adipocyte appears deformed, indicating that macrophages adhere to each other and exert a compressive force on the adipocyte. Several comma shaped dense macrophage nuclei appear on the

FIGURE 3 Adipocytes do not express TNFα. Whole adipose tissue was stained with BODIPY 558/568 for lipid (gray), Hoechst for nuclei (blue), anti-F4/80 for macrophages (red), anti-TNFα (green). All images were captured using LSCM with a 63X objective and are 2D projections of a 3D image stack. (a) Negative TNFα staining throughout the adipocytes visualized. Nuclei not shown. (b) TNFα staining was associated only with macrophages (arrows). No adipocytes stain for TNFα. Yellow in the overlay image indicates co-localization of TNFα and F4/80.
Despite the segmentation of the adipocytes, a single adipocyte nucleus was observed within a continuous cytoplasm in both cases. The segmented lipid seen in Figure 6c is most likely also due to this process. This intimate interaction between adipocytes and macrophages occurs in “adipocyte” floating layer cells obtained from obese mice.

**Macrophages in the Adipocyte Floating Layer are Responsible for TNFα Expression**

Floating layer preparations from non-obese and obese mice were analyzed for TNFα and IL-10 expression by qRT-PCR. There was little expression of either TNFα or IL-10 in the non-obese mice. IL-10 increased significantly by 10-fold and TNFα increased significantly by 40-fold in floating layer cells obtained from obese mice. The amount of TNFα was significantly greater than that of IL-10.

Macrophages and adipocytes in the floating layer from obese mice were analyzed by LSCM in order to determine which cells were expressing TNFα. TNFα staining consistently coincided with F4/80 macrophage staining (Figure 6b) and there was no TNFα stain associated with the adipocyte itself. In multiple experiments at different time points of feeding, we never observed floating adipocytes that stained positive for TNFα. There is a loss of grey lipid mirroring the heaviest areas of TNFα staining leaving black clear tracks (Figure 6b, lipid panel). IL-10 staining was negative for both the macrophages and the adipocyte shown.

In addition to macrophages adherent to adipocytes, isolated macrophages were observed as well in the floating layer. Invariably, such macrophages had either engulfed significant amounts of lipid or they were in contact with lipid droplets, which would explain why they were found in the low-density fraction. Figure 6c illustrates macrophages stained with F4/80 in the floating adipocyte layer that are surrounding, or are engorged with lipid. The lipid areas shown in Figure 6c are too small (25-50 μm) to be full adipocytes (100-300 μm, see scale in Figures 5a-c and 6b overlay) and lack an adipocyte nucleus. High-resolution confocal analysis of the
FIGURE 5 Adipocytes in the floating layer were fully liberated from adipose tissue, yet macrophages remained bound to single cell adipocytes recovered from obese mice. Floating layer cells were stained with anti-F4/80 (red) to stain macrophages, BODIPY 558/568 (grey) to stain lipid and Hoechst dye (blue) to stain nuclei. Anti-F4/80 primary antibody was visualized using anti-rat IgG Alexa647 conjugated secondary antibody. A sample from obese mice was also stained using the secondary antibody only and showed no background staining (data not shown) (a) Images show liberated, intact adipocytes after collagenase digestion and washing from non-obese and obese mice. The top panels show increased F4/80 covering adipocytes from obese mice when compared to non-obese mice. Images were collected with 20× objective. (b) Individual adipocyte images from obese and non-obese mice collected with 63× objective. In the obese condition, macrophages are seen covering individual intact adipocytes. An intact adipocyte was confirmed by the presence of an adipocyte nucleus (white arrow), while macrophage nuclei are morphologically different and stain more densely (blue arrow). (c) A one micron section taken from the middle of the 3D z-stack of the macrophage covered adipocyte in Figure 5b reveals that the large, single adipocyte is effectively being pinched in half by the macrophages surrounding it. This can be seen by the continuation of lipid (gray) between the attached circular bodies and the break in F4/80 staining on the right-hand side of the frame (yellow arrow) in contrast to the distinct border (red arrow) of macrophages covering the separate adipocytes on the left-hand side of the frame.
adipocyte floating layer revealed repeated z-stacks of macrophages with engulfed lipid. The single macrophage (single blue comma shaped nucleus indicated by the blue arrow) depicted in Figure 6c is in tight contact with a lipid droplet (grey lipid covered faintly by F4/80 red stain and flattened contact interface). There are smaller areas of lipid inside the body of the macrophage. The macrophage is expressing TNFα (Figure 6c), but not IL-10. The pattern of TNFα staining (Figure 6c) did not coincide with lipid staining (Figure 6c) indicating that the TNFα is only associated with the macrophage.

Figure 6d illustrates another example of a macrophage having surrounded or engulfed lipid. In this case, the macrophage is clearly expressing IL-10, but also a small amount of TNFα as shown in the single section views. IL-10 secreting macrophages in the floating layer as seen in Figure 6d did not occur as frequently as macrophages expressing TNFα alone, reflecting the qRT-PCR results.

**Discussion**

The confocal immunohistochemical analysis in this study has revealed novel insights into the cellular architecture of the inflammatory infiltrate of obese adipose tissue and the interaction between macrophages and adipocytes and adipocyte-derived lipid. The data show that macrophages are in tight contact with adipocytes or lipid droplets, an interaction that survives tissue disruption, collagenase digestion and density gradient centrifugation. This suggests that macrophages are not simply trapped by surrounding adipocytes, but are retained in the floating layer, possibly by physical attachment to low density adipocytes or lipid droplets. Tight contact with adipocytes is consistent with a role for macrophages in clearing dying adipocytes (7).

In addition to adhering to adipocytes or lipid droplets, macrophages were observed to form contacts with each other when surrounding the much larger adipocytes. Although reminiscent of multinuclear giant cells found in granulomatous disorders (21), the sheath-like structures partially covering adipocytes have not been observed previously. Sheaths of macrophages impinge on the adipocyte and the macrophage-macrophage interaction appears to have sufficient strength to compress the adipocyte in the contact area. Most likely,
the compressed adipocytes in Figures 5c and 6b, visible here in 3D, represents CLS detected in histological sections with 6b representing an earlier time point in the progression of coverage.

Macrophages in contact with lipid structures that are smaller than an adipocyte were observed as well. Most likely, these are lipid droplets or lipid-filled fragments of adipocytes. Although it is not clear whether macrophages can break up lipid droplets, ingest and ultimately degrade ingested lipid, our data, like others (7), clearly show that macrophages do have mechanisms to take up lipid in adipose tissue. In this study, lipid-laden macrophages in CLS were observed by LSCM imaging of WAT. Lipid uptake may constitute another mechanism, in addition to contact with low-density lipid structures that reduces the density of macrophages to allow them to float during density gradient centrifugation. In a previous study, density gradient centrifugation was used to isolate lipid-laden macrophages (22). The mechanism of lipid uptake and transport that operates in adipose tissue is currently unclear. Cinti et al. (7) suggested that the residual lipid, in necrotic adipocytes, fragmented into smaller lipid droplets which can be phagocytosed by CLS macrophages.

It has been suggested that adipocytes can grow only up to a threshold above which they will generate a stress signal or die (23). In fact, increased adipocyte cell death has been observed in obese mice and humans (7,24). Cinti et al. (7) determined that the adipocytes which were surrounded by macrophages in adipose tissue forming CLS were in fact dead. In the experiments described here, this condition occurred in obese mice with 5-7% adiposity. This suggests that CLS are a physiologic response to obesity exceeding a threshold.

The 3D analysis of the cellular architecture of the inflammatory infiltrate allowed unequivocal assignment of TNF-α to macrophages. In all the experiments performed (1-4-5-month feeding of HF or HF/HC diets to generate obese mice), we did not observe a single adipocyte expressing TNF-α in intact WAT or in the separated floating layer. All TNF-α staining was associated with macrophages. In WAT macrophages expressed TNF-α, and the vast majority of the macrophages in the separated floating layer expressed TNF-α. IL-10 expression was observed to a lesser extent, but again was associated with macrophages. A high ratio of TNF-α to IL-10 expression was reflected in cytokine mRNA levels. There are reports of simultaneously elevated inflammatory TNF-α (25,26) and immunoregulatory IL-10 (27,28) in obese individuals. Co-existence of TNF-α and IL-10 with an imbalance in favor of TNF-α are hallmarks of chronic inflammatory conditions (6).

In vitro, TNF-α is cytotoxic to both cultured adipocytes and pre-adipocytes (29-33). TNF-α has also been shown to activate lipolysis in adipocytes [reviewed in Ref. (34)] via the TNF receptor I (TNFR1) (35,36). Indeed, in this study, lipid staining of adipocytes was visibly diminished adjacent to sites where macrophage expression of TNF-α was heaviest. The lipotoxic and lipolytic activity of macrophage products and the macrophage activating activity of responding adipocytes establish a self-reinforcing vicious cycle maintaining chronic adipose tissue inflammation (18).

As observed by others previously, CLSs contain activated macrophages (15,16). The majority of macrophages in the floating layer were pro-inflammatory expressing TNF-α. Activated macrophages express adhesion molecules allowing for tight adherence (37,38). In order to deform adipocytes as shown in this study, macrophages need to tightly adhere to each other as well as the adipocyte target. The tight contact of macrophages with each other and with adipocytes has implications for future studies of inflamed adipose tissue (39). The only procedures that would recover adipocytes from inflamed areas of the tissue would require disrupting macrophage/macrophage and macrophage/adipocyte interactions. There is currently no known procedure to achieve this and allow the preservation of cell integrity. The molecular nature of the tight adherence of macrophages to other macrophages and to adipocytes warrants further investigation.

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