Obesity aggravates the joint inflammation in a collagen-induced arthritis model through deviation to Th17 differentiation

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http://dx.doi.org/10.3858/emm.2012.44.7.047
Accepted 10 April 2012
Available Online 19 April 2012

Abbreviations: AIA, adjuvant-induced arthritis; BMI, body mass index; CCP, cyclic citrullinated peptide; CIA, collagen-induced arthritis; DIO, diet-induced obesity; EAE, experimental autoimmune encephalomyelitis; HFD, high fat diet; MLR, mixed lymphocyte reaction; OA, osteoarthritis; RA, rheumatoid arthritis

Abstract

White fat cells secrete adipokines that induce inflammation and obesity has been reported to be characterized by high serum levels of inflammatory cytokines such as IL-6 and TNF-α. Rheumatoid arthritis (RA) is a prototype of inflammatory arthritis, but the relationship between RA and obesity is controversial. We made an obese inflammatory arthritis model: obese collagen-induced arthritis (CIA). C57BL/6 mice were fed a 60-kcal high fat diet (HFD) from the age of 4 weeks and they were immunized twice with type II collagen (CII). After immunization, the obese CIA mice showed higher arthritis index scores and histology scores and a more increased incidence of developing arthritis than did the lean CIA mice. After treatment with CII, mixed lymphocyte reaction also showed CII-specific response more intensely in the obese CIA mice than lean CIA. The anti-CII IgG and anti-CII IgG2a levels in the sera of the obese CIA mice were higher than those of the lean CIA mice. The number of Th17 cells was higher and the IL-17 mRNA expression of the splenocytes in the obese CIA mice was higher than that of the lean CIA mice. Obese CIA mice also showed high IL-17 expression on synovium in immunohistochemistry. Although obesity may not play a pathogenic role in initiating arthritis, it could play an important role in amplifying the inflammation of arthritis through the Th1/Th17 response. The obese CIA murine model will be an important tool when we investigate the effect of several therapeutic target molecules to treat RA.

Keywords: arthritis, experimental; inflammation; mice; obesity; Th17 cells

Introduction

Obesity has been considered to be one of the most important risk factors for osteoarthritis (OA) in knees. According to the longitudinal studies, hand OA is also related to obesity (Carman et al., 1994; Oliveria et al., 1999; Grotle et al., 2008). Obesity plays a prominent role due to the overloading of joints and the over-production of pro-inflammatory molecules that can exert detrimental effects on the articular tissues. Activated white adipose tissue increases the synthesis of pro-inflammatory cytokines such as IL-6, IL-1, IL-8, TNFα and IL-18, while regulatory cytokines such as IL-10 are decreased (Iannone and Lapadula, 2010). Recent results indicate that IL-6 gene is associated with fat mass in Caucasian men (Andersson et al., 2010). Adipocytes cross-talk with other cells by producing...
and secreting peculiar proteins named adipokines and these are called leptin, resistin and visfatin (Iannone and Lapadula, 2010). Researchers are interested in the relationship between obesity and inflammatory arthritis due to a new concept that obesity induces inflammation.

Two case-control studies, one from UK and the other from the Pacific Northwestern US, have suggested a possible association between a high body mass index (BMI) and the risk of rheumatoid arthritis (RA) (Voigt et al., 1994; Symmons et al., 1997). Merete et al. proposed that obesity was selectively associated with risk of anti-cyclic citrullinated peptide (CCP) positive RA (Pedersen et al., 2006). On the other hand, some investigators have suggested that obesity does not influence the likelihood of developing RA. A high BMI is associated with a less severe disease outcome for anti-CCP positive patients with RA (van der Helm-van Mil et al., 2008). In patients with established RA, both a very low BMI and a very high BMI and the percentage of body fat are independently associated with increased disease activity and physical dysfunction (Stavropoulos-Kalinoglou et al., 2009). Unfortunately, epidemiological studies also have shown contradictory findings.

Although both inflammatory arthritis such as RA and obesity have been reported to be characterized by high serum levels of inflammatory cytokines, the frequency of one disorder was not increased in the other. Obesity is not a manipulable variable in human studies and only observation study is possible. It is also hard to assess some effective factors in cohort studies. It is very hard to confirm that obesity is a major risk factor of RA or that obesity has an association with inflammatory arthritis such as RA.

Therefore, we suggest a new murine model, which is called the obese collagen-induced arthritis (CIA) model. In obese CIA mice, arthritis was induced by CII collagen using diet-induced obesity (DIO) mice. We compared the obese CIA mice and the lean CIA mice for the severity and incidence of arthritis, the presence of autoantibodies, expression of IL-17 and T cell differentiation. This animal model suggests that obesity be associated with aggravating inflammatory arthritis. We can perform research on the alterations of the cytokine profiles and the immune reaction and experiment on diverse therapeutic interventions using this CIA animal model.

**Figure 1.** The obesity CIA enhanced induction of arthritis score and incidence of arthritis. The mice were given either standard chow or a high fat diet from 4 weeks of age. Mice sacrifice was done the first time the mouse weighed 30 g. HFD group exhibited significantly higher serum glucose, cholesterol and triglyceride concentrations than standard chow group. (A) Glucose, triglyceride, and insulin levels. Values are means \( \pm \) S.E.M. \((n = 6)\). The mean values for the arthritis scores are indicated on the graph. \( P < 0.01 \) compared with the saline control. A representative result of at least three independent experiments is shown. (B) Joints sections from the obese CIA and CIA mice were stained with H&E, Safranin O and Toluidine blue (Original magnification of the H&E: 40 \( \times \), Safranin O and Toluidine blue: 200 \( \times \)).
Results

Comparison of the arthritis score and the incidence of arthritis between the obese CIA and CIA mice

We made an obese model. The normal contained 16.0% fat calories, whereas 60% of the calories of HFD were fat calories. Mice sacrifice was done the first time the mouse weighed 30 g. HFD increased glucose ($P = 0.0331$), total cholesterol ($P = 0.0237$) and triglyceride ($P = 0.0095$) levels in the serum as compared with the lean mice group (Figure 1A).

First, we made an obese inflammatory arthritis model. The C57BL/6 (wild-type) mice and the C57BL/6 mice fed a high fat diet (60 Kcal) were immunized with chicken type II collagen (CII) and the development of arthritis was followed for 10 weeks. Both the C57BL/6 (CIA) mice and the C57BL/6 mice fed a high fat diet (obese CIA) mice developed severe arthritis. However, the mice fed a high fat diet had significantly higher arthritis severity scores and a higher incidence of arthritis than that did the C57BL/6 mice at all time points (Figures 1B and 1C). We assessed synovial inflammation ($P = 0.0086$) by the H&E stained sections and cartilage damage ($P = 0.0050$) by safranin-O and toluidine blue staining in the knee joints. The histology of the swollen joints in the obese CIA mice showed a massive infiltration of mononuclear and multinuclear cells into the joint and there was synovial pannus formation and progressive articular destruction as compared to that of the CIA mice (Figure 1D).

T cell responses in the obese CIA and CIA mice

The proliferative recall response to CII was performed to assess the degree of the T cell response between the obese CIA and CIA mice. The cells obtained from the spleens of the CIA mice with obesity showed increased T cell proliferation in response to anti-CD3 plus anti-CD28 stimulation as well as in response no stimulation as compared with those cells obtained from the CIA mice (Figure 2).

Antigen-specific autoantibodies in the obese CIA and CIA mice

The concentration of IgG anti-CII, which reflects the Th1 response, was significantly greater in the obese CIA mice than that in the CIA mice ($P = 0.0105$) (Figure 3A). Additionally, the level of CII-specific IgG2a in the serum, which was higher in the obese CIA mice as compared to that of the CIA mice ($P = 0.0045$) (Figure 3B).

Quantification of intracellular IL-17 in the splenocytes of the obese CIA and CIA mice

We then performed intracellular IL-17A staining on the splenocytes from the obese CIA and CIA mice. The IL-17 mRNA levels in the splenocytes of the obese CIA mice were higher than that of the CIA mice ($P = 0.0491$) (Figure 4A). Consistent with the above results, there were more IL-17-secreting cells in the CD4+ cell fraction in the obese CIA mice than that in the CIA mice (Figure 4B).

Discussion

DBA/1 strain mice have been widely used for CIA studies, including the studies involved in the assessment of potential arthritis therapies (Malfait et
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Figure 4. Obesity induces IL-17 in the CIA mice. (A) The expression of IL-17 in the splenocytes obtained from each mouse was determined by real time PCR. The data is expressed as the mean ± SD. *P < 0.05, **P < 0.01 compared with the CIA mice. (B) The splenocytes from the mice of each group were cultured with 25 ng/ml PMA and 250 ng/ml ionomycin for 4 h. The expression of IL-17 was determined by intracellular flow cytometry analysis. The data is representative of three independent experiments. (C) Immunohistochemical assessment of IL-17 expression in mice joint with CIA and obese (original magnification, 400 ×).

al., 2001). Usually, the DBA/1 strain mice show well induced CIA compared to that of the C57BL/6 mice. Though this study was performed with the C57BL/6 strain, the obese CIA mice well manifested arthritis. It suggested that obesity was related to overcome the resistance to develop CIA in C57BL/6 strain and to be susceptible to inflammation and autoimmunity. Obese CIA model in C57BL/6 strain itself was good evidence that obesity influence on developing autoimmune reaction. Diet-induced obesity (DIO) with C57BL/6 strain induced a good osteoarthritis model expressing high leptin, adiponectin, and IL-1α. DIO showed biochemical, neurobehavioral, musculoskeletal, inflammatory and structural knee joints change (Griffin et al., 2010). This model exhibits that obesity has possibility to accelerate inflammation process through inflammatory adipokines and cytokines.

The C57BL/6 mice were fed a 60 Kcal high fat diet from the age of 4 weeks and they were immunized twice with CII. After immunization with CII, the obese CIA mice (C57BL/6 strain) showed a higher arthritis index score and histology score, and there was a more increased incidence of developing arthritis than that of the control (the C57BL/6 strain lean CIA mice) (Figures 1B-1D). This result is important evidence that obesity has an amplifying action in the development of inflammatory arthritis.

The purified splenic lymphocytes from obese CIA mice or lean CIA mice, 10 weeks after the first immunization with CII, were or were not stimulated with anti-CD3. In both conditions (with or without anti-CD3 treatment), the mixed lymphocyte reaction (MLR) of the obese CIA mice was more increased than that of the lean CIA mice (Figure 2). It is well known that T cells have a critical role in developing CIA. In the condition of the treatment with CII, CII specific response was shown in MLR. The splenocytes were proliferated in response to CII antigen. This result represents that obesity operates immune systems through activation of CII specific T cells. Fat cells secret some materials and these react with the immune system and particularly T cells.

Anti-CII IgG and anti-CII IgG2a (autoantibodies of CIA mice) were checked in the serum of both the obese and lean CIA mice. The levels of autoantibody of the obese CIA mice were higher than those of the lean CIA mice (Figure 3). IgG2a type antibodies were produced by the Th1 response. Thus, obesity may cause the T cells of the CIA mice to differentiate toward Th1 cells.

Some investigators have demonstrated that obesity selectively promotes an expansion of the Th17 T cell subtype, which is a subset with prominent pro-inflammatory roles in experimentally induced encephalomyelitis (EAE) model (Winer et al., 2009). The T cells from DIO mice expand the Th17 cell pool and produce progressively more IL-17 than that of the lean mice (Winer et al., 2009). Moreover, IL-17 plays an important role in the pathogenesis of RA and CIA. The studies that have employed strategies to neutralize or delete IL-17 have shown that Th17 cells have a pathogenic role in CIA (Nakae et al., 2003; Lubberts et al., 2004; Ju et al., 2008; Sumarac-Dumanovic et al., 2009). Both obesity and CIA activate Th17 cells and this produces
IL-17. As expected, we observed a more increased number of Th17 cells in the obese CIA mice than that in the lean CIA mice. There was a higher IL-17 mRNA expression in the splenocytes of the obese CIA mice than that of the lean CIA mice (Figure 4A). Moreover, in immunohistochemistry, there was high expression in synovium of obese CIA compared with lean CIA mice (Figure 4C). We suggest obesity amplifies the joint inflammation of CIA mice through Th17 T cell differentiation and IL-17 production in joint synovium.

It is well established that obesity promotes Th1 deviation and induces inflammatory condition. However, it is not certain that Th17 cell is related with obese-induced inflammation and autoimmunity. Although the precise mechanisms are not clear, EAE model suggested Th17 cells and IL-17 in obese mice have an important role aggravating disease (Winer et al., 2009). According to our results, obese CIA mice show more severe arthritis and an increased incidence of arthritis than do lean CIA mice. This is probably due to Th1 and Th17 differentiation. Sarkar et al. proposed that the Th1/Th2/Th17 balance is important to initiate autoimmunity (Sarkar et al., 2009). Both CIA mice and DIO mice are biased towards Th1 and Th17 differentiation. It is no wonder that the obese CIA mice deviate more to Th1 and Th17 T cells than do the lean CIA mice. We suggest a hypothesis that Th17 cells may be influenced by adipokines and Th17 cells may have some adipokine receptors on cell surface. Clarifying these mechanisms of relationship between adipokines and Th17 cells or IL-17 would be another study.

We made an animal model that represented RA with obesity. In the future, we will investigate the interactions between several cytokines and adipokines using this obese CIA model. Obesity studies have revealed that adipokines are important factors in inflammation. The upregulated resistin at local sites of inflammation and the link between serum resistin, inflammation and the disease activity score suggest a role for resistin in the pathogenesis of RA (Senolt et al., 2007). Bokarewa et al. demonstrated that resistin is a molecule that is accumulated at the site of inflammation and that recombinant resistin injected intraarticularly caused a dose-dependent increase of arthritis (Bokarewa et al., 2005). Neutralization of resistin will be an important therapeutic target in the future. Visfatin, which is also known as pre-B cell colony-enhancing factor, is also another candidate for inducing inflammation in RA. The visfatin gene expression in the synovial tissue and PBMCs of RA patients was significantly higher than that of the controls (Matsui et al., 2008). The positive correlation between the levels of visfatin and resistin suggests that visfatin plays a role in the inflammation of RA (Straburzynska-Lupa et al., 2010). In the future, we will test the adipokines in obese CIA mice and try to find therapeutic modalities through blocking the inflammatory adipokines (Seo et al., 2011).

Other investigators have created adjuvant-induced arthritis (AIA) in obese rats. They tested the effect of the cannabinoid CB1 receptor antagonist rimonabant on AIA in obese rats. They also showed that the inflammation reaction in obese rats is more severe than that in lean rats. The same as in obese AIA rats, it is possible to conduct several therapeutic trials using obese CIA mice (Croci et al., 2007).

In conclusion, obesity plays an addictive role in the inflammation of an inflammatory arthritis model through CII specific T cell differentiation -Th17 T cells differentiation. IL-17 is a pivotal cytokine to accelerate joints inflammation in obesity. The obese CIA model has value that we can control and investigate the obese condition in an inflammatory arthritis model and we were able to pinpoint several therapeutic target molecules.

Methods

Animals and diet
C57BL/6 mice (SLC, Inc., Shouzouka, Japan) that were 4 weeks old were housed in polycarbonate cages. They were fed 60 Kcal fat-derived calories and standard mouse chow (Ralston Purina, St Louis, MO) and water ad libitum. All the experimental procedures were examined and approved by the Animal Research Ethics Committee of the Catholic University of Korea.

Biochemical parameters
The levels of total serum cholesterol were measured using commercial kits from Wako Co. (Osaka, Japan), and glucose, triglycerides were measured using commercial kits from Asan Pharmaceutical Co. (Hwaseong-si Gyeonggi-do, Korea).

Induction of CIA
C57BL/6 mice (4 weeks) were purchased from Charles River Breeding Laboratories. CIA was induced as previously described (Inglis et al., 2008). The CIA immunization was done the first time when the mouse weighed 30 gram. In brief, An emulsion was formed by dissolving 2 mg/ml chick collagen type II (CII; Chondrex, Redmond, WA) overnight at 4°C in 0.5M acetic acid, followed by mixing this with an equal volume of CFA (Chondrex, Redmond, WA). The mice were intradermally injected with the emulsion at two sites at the base of the tail and a slightly more anterior location. A second injection as a booster was done 14 days after the primary immunization.
Mixed leukocyte reaction (analysis on CII specific T cell response)
A CII-specific T cell proliferative response was achieved as previously described (Rosloniec et al., 1996).
Briefly, cells were prepared from the spleens of the obese CIA and CIA mice with arthritis ten weeks after the primary immunization. A single cell suspension was prepared, and 2 × 10^6 cells/well in 96-well flat bottom plates were cultured in the absence or presence of plate-bound CD3 1 μg/ml at 37°C for 72 h, followed by the incorporation of 25 μCi/ml [3H]-thymidine (GE Healthcare, Piscataway, NJ) for the last 16 h of the indicated total culture interval. Then the radioactivity was measured with a Micro Beta (Pharmacia Biotech, Piscataway, NJ).

Clinical assessment of arthritis
The severity of arthritis was determined by three independent observers. The mice were observed three times a week for the onset and severity of joint inflammation for up to 10 weeks after the primary immunization. The severity of arthritis was assessed on a scale of 0–4 with the following criteria, as was described previously (Rosloniec et al., 1996): 0 = no edema or swelling, 1 = slight edema and erythema limited to the foot or ankle, 2 = slight edema and erythema from the ankle to the tarsal bone, 3 = moderate edema and erythema from the ankle to the tarsal bone and 4 = edema and erythema from the ankle to the entire leg. The arthritic score for each mouse was expressed as the sum of the scores of three limbs.

Histopathological analysis of arthritis
The mouse joint tissues were fixed with 10% formalin, decalcified in EDTA and embedded in paraffin. The sections were dewaxed using xylene and dehydrated in a graded series of alcohol solutions. The sections were then stained with Harris hematoxylin and eosin (H&E), Safranin O and Toluidine blue to detect proteoglycans. The H&E stained sections were scored for inflammation and bone erosion. Inflammation was scored according to the following criteria: 0 = no inflammation, 1 = slight thickening of the lining layer or some infiltrating cells in the underlying layer, 2 = slight thickening of the lining layer plus some infiltrating cells in the underlying layer, 3 = thickening of the lining layer plus some infiltrating cells in the underlying layer, 4 = synovium highly infiltrated with many inflammatory cells. Cartilage damage was determined using safranin-O staining and Toluidine blue and the extent of cartilage damage was scored according to the following criteria (Rosloniec et al., 1996): 0 = no destruction, 1 = minimal erosion limited to single spots, 2 = slight to moderate erosion in a limited area, 3 = more extensive erosion and 4 = general destruction.

Immunohistochemistry was performed using the Vectastain ABC kit (Vector Laboratories, Burlingame, CA) and incubated with anti-CD4 PerCP and they were permeabilized with cytofix/Cytoperm solution (BD). Intracellular staining for flow cytometry was performed with the anti-IL-17 PE (BD Pharmingen) for 4 h. The cells were washed in Cytoperm buffer and in FACS buffer before data acquisition on a FACS Calibur (BD) and the analysis was done using FlowJo software (Tree Star, Ashland, OR).

Measurement of type II collagen specific antibodies
The serum levels of type-II-collagen-specific IgG2a and IgG1 were measured by enzyme-linked immunosorbent assay (ELISA), as previously described, with minor modifications (Rosloniec et al., 1996). Briefly, microtiter plates were coated with type II collagen (4 μg/ml in PBS) at 4°C overnight and this was followed by a blocking step for 30 min at room temperature. The serum samples were then diluted 1:8000 in Tris buffered saline (pH 8.0) that contained 1% chicken serum albumin and 0.5% Tween-20, and the samples were incubated in the microtiter plates for 1 h, after which time the plates were washed five times. The concentrations of IgG2a and IgG1 were measured using mouse IgG2a and IgG1 ELISA Quantitation Kits (Bethyl Laboratories, Montgomery, TX), respectively. Standard serum from the arthritic mice was added to each plate in serial dilutions, and a standard curve was constructed to assign arbitrary units to the levels of anti-type-II-collagen IgGa and IgG1. The absorbance values were determined with an ELISA microplate reader operating at 450 nm.

Cell preparation and culture
Ten weeks after the primary immunization, the mouse spleens were collected for cell preparation and they were washed twice with PBS. The spleens were minced and the red blood cells were lysed with 0.83% ammonium chloride. The cells were filtered through a cell strainer and then they were centrifuged at 1300 rpm at 4°C for 5 min. The cell pellets were resuspended in RPMI 1640 medium and plated in 48-well plates (Corning, NY) at a concentration of 1 × 10^6 cells/well.

Intracellular staining for flow cytometry
Cell pellets were prepared from the spleens of the obese CIA and CIA mice with arthritis ten weeks after the primary immunization. The spleenocyte cells were stimulated with 50 ng/ml PMA and 500ng/ml ionomycin (both from Sigma, St. Louis, MO) and GolgiStop (BD Pharmingen) for 4 h. The cells were stained with anti-CD4 PerCP and they were permeabilized with cytofix/Cytoperm solution (BD). Intracellular staining with IL-17 PE was performed in Cytoperm Buffer (BD) according to the manufacturer’s protocol. The cells were washed in Cytoperm buffer and in FACS buffer before data acquisition on a FACs Calibur (BD) and the analysis was done using FlowJo software (Tree Star, Ashland, OR).

Expression of mRNA as measured by RT-PCR
The mRNA was extracted using RNAzolB (Biotex Laboratories, Houston, TX) according to the manufacturer’s instructions. Reverse transcription of 2 μg aliquots of total mRNA was carried out at 42°C using the Superscript reverse transcription system (TaKaRa, Shiga, Japan). The expression of IL-17 mRNA was determined by performing real-time PCR with SYBR Green I. A LightCycler (Roche Diagnostics Mannheim, Germany) was used for the quanti-
tative analysis of the IL-17 mRNA. The following sense and antisense primers for each molecule were used (5' → 3'): IL-17 sense CCT CAA AGC TCA GCG TGT CC, IL-17 antisense GAG CTC ACT TTT GCG CCA AG and β-actin sense GAATGCGTGGCTGACATCAAG, β-actin antisense TGTAGTTCTAGGATGCCACAG. The reaction mixtures contained 2 μl of LightCycler FastStart DNA mastermix for STBR Green I, 0.5 μM of each primer, 4 μl MgCl$_2$ and 2 μl of template DNA. All the capillaries were sealed, centrifuged at 500 g for 5 s and then amplified in a LightCycler instrument with activation of polymerase (95°C for 10 min) as follows: 45 cycles of 10 s at 95°C, 10 s at 60°C, and 10 s at 72°C. The double-stranded PCR products were measured during the 72°C extension step by detection of the fluorescence associated with the binding of SYBR Green I to the product. The fluorescence curves were analyzed with LightCycler software v.3.0. The expressed levels of IL-17 mRNA were calculated and corrected for the values of the endogenously expressed housekeeping gene (β-actin) controls.

Statistical analysis
The results are expressed as means ± S.D. (or means ± S.E.M.). Groups of data were compared using the Mann-Whitney U-test. Differences were considered statistically significant at $P < 0.05$.

Acknowledgements
This research was supported by a grant of the Korea Health Technology R&D Project, Ministry for Health, Welfare & Family Affairs, Republic of Korea (grant number A092258).

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