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A Role for the High-Density Lipoprotein Receptor SR-B1 in Synovial Inflammation via Serum Amyloid-A

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Acute phase apoprotein Serum Amyloid A (A-SAA), which is strongly expressed in rheumatoid arthritis synovial membrane (RA SM), induces angiogenesis, adhesion molecule expression, and matrix metalloproteinase production through the G-coupled receptor FPRL-1. Here we report alternative signaling through the high-density lipoprotein receptor scavenger receptor-class B type 1 (SR-B1). Quantitative expression/localization of SR-B1 in RA SM, RA fibroblast-like cells (FLCs), and microvascular endothelial cells (ECs) was assessed by Western blotting and immunohistology/fluorescence. A-SAA–mediated effects were examined using a specific antibody against SR-B1 or amphipathic α-Helical Peptides (the SR-B1 antagonists L-37pA and D-37pA), in RA FLCs and ECs. Adhesion molecule expression and cytokine production were quantified using flow cytometry and ELISA. SR-B1 was strongly expressed in the RA SM lining layer and endothelial/perivascular regions compared with osteoarthritis SM or normal control synovium. Differential SR-B1 expression in RA FLC lines (n = 5) and ECs correlated closely with A-SAA, but not tumor necrosis factor α–induced intercellular adhesion molecule-1 up-regulation. A-SAA–induced interleukin-6 and -8 production was inhibited in the presence of anti-SR-B1 in human microvascular endothelial cells and RA FLCs. Moreover, D-37pA and L-37pA inhibited A-SAA–induced vascular cell adhesion molecule-1 and intercellular adhesion molecule expression from ECs in a dose-dependent manner. As SR-B1 is expressed in RA synovial tissue and mediates A-SAA–induced pro-inflammatory pathways, a better understanding of A-SAA–mediated inflammatory pathways may lead to novel treatment strategies for RA.

Rheumatoid arthritis (RA) is a chronic progressive autoimmune disease characterized by proliferation of the synovial membrane (SM), which leads to degradation of articular cartilage and subchondral bone. Normal SM consists of a monolayer of synoviocytes including fibroblast-like cells (FLCs) and macrophage-like synoviocytes, which produce a proteoglycan-rich synovial fluid to lubricate the joint and provide nutrition to the avascular cartilage. A critical early event in synovial inflammation is angiogenesis, where new blood vessels develop from existing blood vessels and act as a conduit for the delivery of nutrition and invading immune cells into the joint. Recruitment of immune cells into the joint is mediated by tissue expression of chemokines and by expression of cell surface adhesion molecules such as intercellular adhesion molecule-1 (ICAM-1) and vascular cell adhesion molecule-1 (VCAM-1), which selectively recruit mononuclear cells through their integrin receptors ligands. In RA, activation of the SM transforms the lining layer into a hyperplastic tumor-like ‘pannus’ composed primarily of activated FLCs, macrophage-like synoviocytes, and lymphocytes, which through self perpetuating and persistent pro-inflammatory activation, are capable of destroying adjacent articular cartilage and bone.

Acute phase serum amyloid A (A-SAA) is a highly-conserved acute phase apoprotein whose serum levels increase up to a 1000 fold within hours of an inflammatory stimulus. Unlike other acute phase proteins, which are synthesized primarily in the liver as part of the systemic acute phase response, A-SAA is also markedly expressed at local sites of tissue inflammation. A-SAA is also known to be present at high levels in wound repair and in cancer tissues. A-SAA, at normal serum levels,

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associates with high-density lipoprotein (HDL) forming a heterogenous HDL population containing both A-SAA and apolipoprotein A-1 (ApoA-1). During the inflammatory response, however, A-SAA is dramatically elevated in serum (1 to 1000 μg/ml), at which levels A-SAA displaces ApoA-1 and saturates HDL resulting in high levels of free circulating A-SAA. Our group has demonstrated a strong correlation between serum A-SAA and disease activity in RA. Furthermore, we and others have demonstrated that A-SAA is produced by synovial FLCs and articular chondrocytes, where it is a powerful inducer of matrix metalloproteinases in these cells in vitro. A-SAA induces the secretion of pro-inflammatory cytokines tumor necrosis factor-α (TNF-α), interleukin-1β (IL-1β), and interleukin-8 (IL-8) and acts as a chemoattractant for human monocytes, neutrophils and T cells. A-SAA has high affinity for the extracellular matrix components laminin and fibronectin and has been shown to enhance leukocyte binding to extracellular matrix through a β1 integrin dependent mechanism. We have demonstrated increased ICAM-1/VCAM-1 expression, increased endothelial cell (EC) migration, EC tubule formation, and matrix metalloproteinase 1 production in response to A-SAA in RA FLCs and ECs with an effect equipotent to TNF-α and IL-1β. These studies together indicate that A-SAA may play a primary pathogenic role locally in joint destruction and mediating cartilage destruction.

A-SAA is known to bind to a formyl peptide receptor-like 1 (FPRL-1), a seven-transmembrane G protein–coupled receptor. Through FPRL-1, A-SAA interacts with a diverse array of exogenous and endogenous ligands in the inflammatory process including leukocyte migration, IL-8 induction, and matrix metalloproteinase production. We, and others, have shown overexpression of A-SAA and its receptor FPRL-1 in RA SM, Lipoxin (LXA4) A4, an eicosanoid metabolite with anti-inflammatory actions, has recently been shown to inhibit the pro-inflammatory actions of A-SAA by competitive binding with FPRL-1 and differential regulation of the transcription factor nuclear factor-kappa B (NF-kB). Recent evidence has demonstrated alternative A-SAA binding to and signaling through the HDL receptor human scavenger receptor class B type 1 (SR-B1), also known CD36 and LIMPii analogues-1 (CLA-1), is the human orthologue of rodent SR-B1, which functions as a membrane bound receptor for HDL. A-SAA binding to SR-B1 expressed on a monocyte cell line, induces chemokine production, activates extracellular signal-regulated kinases 1/2 (ERK1/2) and p38 mitogen activated protein kinases, a process that may involve cholesterol efflux. A-SAA binding and signaling through SR-B1 have been shown to be inhibited by co-incubation with ApoA-1 or a synthetic mimetic peptide of ApoA-1 known as L-37PA in a dose-dependent manner. Ligand binding of L-37PA to SR-B1 has also been shown to block the pro-inflammatory cytokine response to LPS in THP-1 cells, although to date this effect has not been demonstrated for intact ApoA-1.

In this study we provide further evidence of a pathophysiological role of A-SAA in driving the pro-inflammatory response in RA through SR-B1. We demonstrate strong expression of SR-B1 in RA SM and differential expression of SR-B1 on RA FLC lines that correlate with A-SAA responsiveness. Furthermore we demonstrate that the pro-inflammatory actions of A-SAA in RA FLCs and human microvascular endothelial cells (HMVECs) can be blocked both by specific antibodies directed against SR-B1 and by synthetic mimetic peptides of ApoA-1. These findings support a novel role for A-SAA as an important mediator of synovial inflammation and suggest a novel strategy for A-SAA as a potential future therapeutic target in the treatment of inflammation through modulation of SR-B1.

Materials and Methods

Patient Recruitment and Arthroscopy

Patients with RA (n = 8), according to the criteria of the American College of Rheumatology, or osteoarthritis (OA; n = 5) were recruited from rheumatology outpatient clinics at St. Vincent’s University Hospital along with one normal healthy control subject. RA patients had clinically active disease including at least one inflamed knee joint. When compared with OA patients, RA patients had statistically higher serum levels of systemic inflammation as measured by C-reactive protein (18 ± 16 RA vs 5 ± 2 OA, mmol/L, P < 0.05) and erythrocyte sedimentation rate (29 ± 7 RA vs 18 ± 7 OA, mm/hr, P < 0.05, mean ± SEM). There was no difference in the age range of patients when RA (59 ± 4 yrs) was compared with OA (55 ± 24 yrs). After approval by the institutional ethics committee, all patients gave written informed consent, before a day case arthroscopy procedure under local anesthetic. Arthroscopy of an inflamed knee joint using a 2.7-mm-diameter needle telescope (Storz, Tuttingen, Germany) was performed and biopsies of inflamed SM were obtained under direct visualization as previously described. Synovial biopsies were obtained from seven RA patients. SM Biopsies were snap frozen in OCT, foil wrapped, and stored at −70°C. Cryostat sections 7 μm thick were cut, dried overnight at 37°C, and stored at −70°C and used for immunohistochemistry. Further biopsies were placed in medium and used to isolate RA primary synovial fibroblasts as outlined below.

Isolation and Culture of RA FLC Cell Lines

Synovial biopsies were digested using 1 mg/ml collagenase (Worthington Biochemical, Freehold, NJ) in RPMI (Gibco-BRL, Paisley, UK) for 4 hours at 37°C in humidified air with 5% CO2. Dissociated cells were grown to confluence in RPMI, 10% fetal calf serum (FCS; Gibco-BRL), 10 ml of 1mmol/L HEPES (Gibco-BRL), penicillin (100 units/ml), streptomycin (100 units/ml), and fungizone (0.25 μg/ml) before passaging. Cells were morphologically homogeneous RA FLCs and were used for experiments between passages 4 and 8.

Isolation and Culture of HMVECs

Dermal-derived HMVECs (Clonetics, San Diego, CA) grown in endothelial basal medium supplemented with 5% FCS, 0.5 ml human epidermal growth factor, 0.5 ml
hydrocortisone, 0.5 ml gentamicin, 0.5 ml bovine brain extract (Clonetics) were used for experiments between passages 3 and 8.

**Immunohistochemical Expression of SR-B1 Receptor in RA Synovial Tissue**

For histological analysis SM samples were placed in cryopreservative embedding media OCT compound (Tissue Tek, Sakura, Finetek, Europe BV, Zoeterwoude, The Netherlands) and immediately frozen in liquid nitrogen. Sections (7 μm) were cut using a cryostat and placed on glass slides coated with 2% 3-aminopropyl-triethoxy-silane (Sigma-Aldrich Ireland Ltd, Dublin, Ireland) dried overnight at room temperature and then stored at −80°C until required for staining. Tissue sections were allowed to reach room temperature, fixed in acetone for 10 minutes, air-dried, and allowed to reach room temperature. Tissue sections were then incubated for 1 hour at room temperature with normal blocking serum (Vectastain Elite Kit, Vector Laboratories Ltd, Peterborough, UK). The sections were incubated with avidin for 15 minutes, rinsed, and then incubated with biotin for 15 minutes. The sections were then incubated for 1 hour with the primary mouse monoclonal antibody for SR-B1 (BD Biosciences) at 1:50 to 1:100 dilutions at room temperature. Secondary antibody (anti-mouse; Vectastain) was prepared and added to the relevant sections and incubated for 1 hour. The secondary antibody was washed off and the slides were incubated with avidin-biotin-peroxidase complex solution for 30 minutes and incubated for 6 minutes with 3,3′-diaminobenzidine and counterstained in hematoxylin stain for 1 minute. The sections were mounted using DPX (BDH Laboratory Supplies, Poole, UK). For negative control, isotype-matched nonimmune IgG was included. Sections were assessed using light microscopy. Sections were specifically examined for evidence of SR-B1 expression by two blinded operators using an established and leukocyte recruitment occur. SR-B1 expression was specifically examined for evidence of SR-B1 expression in articular cartilage and bone occurs, and on vascular ECs and perivascular areas where angiogenesis of articular cartilage and bone occurs, and on vasculature.

**SR-B1 Mediates A-SAA–Induced IL-6 and IL-8 Production in HMVECs and RA FLCs**

To assess the role of SR-B1 on A-SAA–induced IL-6, expression HMVECs (n = 6) and an additional 4 primary RA FLC lines were pre-incubated with purified IgG anti-mouse anti–SR-B1 antibody (10 μg/ml; BD Biosciences) or an isotype matched (IgG1) anti-mouse control (R&D systems) for 2 hours before stimulation with A-SAA (10

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**Immunofluorescent Staining for SR-B1 in HMVECs and RA FLCs**

SR-B1 expression in resting HMVECs and RA FLCs was detected by immunofluorescence microscopy. HMVECs and RA FLCs were grown to 20% confluence in eight-chamber culture slides (Becton Dickenson) in full-strength endothelium basal medium (Clonetics). The cells were then fixed in 4% paraformaldehyde for 20 minutes and blocked using 1% casein solution for 20 minutes. Slides were then incubated with a primary antibody for SR-B1 (mouse monoclonal; 1:100, BD Biosciences) and an appropriate IgG isotype-matched control for 1 hour in a humidified chamber. Chamber slides were then washed in PBS for 5 minutes and then incubated with a Cy2 conjugated secondary antibody (Sigma; 1:200 dilution) for 30 minutes. Slides were washed in PBS for 5 minutes and counterstained with DAPI nuclear stain (Sigma) (1:1000 dilution) for 20 minutes. The cells were then mounted with antifade (Molecular Probes, Eugene, OR) and assessed by immunofluorescence microscopy (×40 magnification).

**SR-B1 Expression in Primary Cell Lines by Western Blot**

For measurement of SR-B1 by Western blot, RA FLC lines (n = 5) or HMVECs 1 × 10⁶/well were seeded into six-well plates in RPMI, 10% FCS for 48 hours. Cells were then washed twice in PBS and lysed in 100 μl SDS sample buffer (62.5 mmol/L Tris/HCl, pH 6.8, 10% glycerol, 2% SDS, 5% mercaptoethanol, and 0.0013% Bromophenol Blue). Lysates were fractionated by SDS/PAGE, 10% acrylamide gels and transferred on to a nitrocellulose membrane (BioTrace®NT, Ponsacola, FL). The membrane was washed for 15 minutes 3× in wash buffer (PBS, 0.1% TWEEN) before being blocked with 5% milk powder in PBA overnight at 5°C. After washing 3× in washing buffer, the membrane was incubated overnight at 5°C with purified monoclonal mouse IgG antibodies for SR-B1 (BD Biosciences) at a 1:500 dilution in 10 ml primary blocking buffer containing PBS, 0.1% TWEEN 20, 2.5% milk powder. The primary antibody was then aspirated off and the nitrocellulose membrane washed 3× in wash buffer. The membrane was then incubated with an enhanced chemiluminescent substrate (Pierce, Rockford, IL) and assessed by immunofluorescence microscopy (×40 magnification).

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**Western Blot**

SR-B1 expression was detected by Western blot using a primary incubation step as described above using a mouse mAb β-actin antibody (Sigma) in 10 ml PBS, 0.1% TWEEN 20, 2.5% milk powder.
Figure 1. SR-B1 protein expression in inflamed RA synovial tissue. RA synovial tissue sections from three representative RA patients (A–C), two OA patients (D and E), and one healthy control (F) were stained with an antibody to SR-B1. A brown stain indicates cells positive for SR-B1 expression. In RA patients SR-B1 expression was demonstrated on (A) synovial intimal blood vessels, (B) the synovial lining and sublining layers, (C) and in synovial vascular ECs (shown by arrow). Minimal staining is seen in OA tissue (D and E) and normal healthy control tissue (F).
RA SM (Figure 1, A–C) compared with OA tissue, which showed minimal staining (Figure 1, D and E) and normal tissue (Figure 1F). Semiquantitative scoring demonstrated increased expression in both the lining layer and sublining of patients with RA compared with OA (1.1 ± 0.52 RA versus 0.5 ± 0.5 OA, lining layer; 1.4 ± 0.72 RA versus 0.8 ± 0.53 OA, sublining). No expression was observed in normal tissue. Immunostaining with an IgG isotype-matched control was completely negative in all samples examined (data not shown).

**SR-B1 Expression on HMVECs by Direct Immunofluorescence**

To confirm the presence of SR-B1 on vascular ECs, SR-B1 immunofluorescence was performed on resting HMVECs grown on culture slides (n = 3). Strong staining for SR-B1 was seen on all HMVECs tested. Staining for IgG control was completely negative (Figure 2, A–D).

**SR-B1 Expression on RA FLCs and HMVECs by Western Blot**

To investigate whether the presence of SR-B1 on RA synovial tissue is due, at least in part, to expression on RA FLCs or ECs, SR-B1 expression was demonstrated on five separate RA FLC lines and on HMVECs. A representative Western blot is shown in Figure 3. SR-B1 expression was demonstrated in all five RA FLC lines studied and on HMVECs (Figure 3A). Variable expression of SR-B1 expression was shown between cell lines and was more marked on RA FLCs than on HMVECs.

The erosive and invasive nature of primary RA FLC lines in vitro have previously been shown to explain different rates of radiographical progression seen between RA patients in vivo.32 Furthermore, high levels of expression of ICAM-1 on RA FLCs are associated with a more aggressive and invasive phenotype.33 To establish whether a relationship exists between SR-B1 expression and responsiveness to A-SAA, ICAM-1 induction after A-SAA stimulation was compared with quantitative baseline SR-B1 expression on four RA FLC lines as shown in Figure 3b. Induction of ICAM-1 by A-SAA correlated closely with baseline SR-B1 expression across RA FLC lines (r² = 0.99, P < 0.05). In contrast, no significant correlation existed between ICAM-1 induction by TNF-α and SR-B1 expression (r² = 0.65, Figure 3c). This suggests that the higher SR-B1 expression in primary cells may predict increased responsiveness to A-SAA.

**A-SAA–Induced IL-6 and IL-8 Production Is Inhibited by Anti-SR-B1 in HMVECs and RA FLCs**

To establish whether a specific antibody SR-B1 could interfere with A-SAA–induced IL-6 and IL-8 production, a further four primary RA FLC lines along with HMVECs were incubated with A-SAA in the presence or absence of anti–SR-B1. Figure 4, A and B demonstrates that A-SAA–induced IL-6 and IL-8 production were significantly inhibited in the presence of anti–SR-B1 (P < 0.05). No inhibition of A-SAA effects on IL-6 and IL-8

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**Figure 2.** SR-B1 staining on resting HMVECs by immunofluorescence. DAPI nuclear stain (A) with corresponding cytoplasmic SR-B1 localized to the same cells (B). DAPI staining (C) with IgG control (D).

**Figure 3.** SR-B1 expression in RA FLCs and HMVECs. A: Representative Western blot showing SR-B1 expression in endothelial cells (ECs) and 5/5 RA FLC lines 80 kDa molecular weight versus β-actin loading control 47 kDa. Bar graph represents quantitative SR-B1 expression by optical density (OD) relative to β-actin OD. A glycosylation band for SR-B1 is seen as previously described. Correlation between SR-B1 expression and ICAM-1 induction in RA FLC lines (n = 4) in response to (B) A-SAA stimulation and (C) TNFα expression. r = Spearman Rank correlation co-efficient. P < 0.05, significant correlation between SR-B1 expression and ICAM-1 induction. SR-B1 expression was calculated before stimulation by calculating SR-B1/actin OD ratios from Western Blots. ICAM-1 measured as Median Flourescene Intensity (MFI), which indicates the level of ICAM-1 expression on a per-cell basis.

**Figure 4.** A-SAA–Induced IL-6 and IL-8 Production Is Inhibited by Anti-SR-B1 in HMVECs and RA FLCs.
production by isometric control IgG1 in HMVECs was observed (Figure 4C).

Figure 5. A–D demonstrates the effect of anti-SR-B1 on A-SAA–induced IL-6 and IL-8 expression in four different primary RA FLC lines. Incubation with anti-SR-B1 inhibits A-SAA–induced IL-6 production in three primary RA FLCs (Figure 5, A–C), with no effect in the fourth RA FLC line. A-SAA–induced IL-8 production in three of four primary RA FLC lines (A and B), with no effect observed in the fourth primary RA FLC line as observed was observed for IL-6. This suggests that SR-B1 mediates both A-SAA–induced IL-6 and IL-8 production, however in comparison with the HMVECs where A-SAA induction and inhibition with anti-CLA-I was consistent, results differed between primary RA FLC lines reflecting the heterogeneity of patients. This is consistent with the immunohistochemistry findings, where two of the eight RA patients had no expression of SR-B1, suggesting it may mediate A-SAA pro-inflammatory effects on some patients but not all.

Amphipathic α-Helical Peptides Inhibit A-SAA–Induced VCAM-1 and ICAM-1 Expression

The effects of the amphipathic α-helical peptides D-37pA and L-37pA on A-SAA–induced cell surface expression of VCAM-1 on HMVECs are shown in Figure 6. A-SAA 0.5 μg/ml stimulation for 6 hours significantly increased VCAM-1 from 20 + 0.8% to 82 + 17% of cells. Co-incubation of cells with both A-SAA 0.5 μg/ml and L-37pA 5 μg/ml significantly inhibited the induction of VCAM-1 expression to 27 + 5% cells (P < 0.001, Figure 6A). Co-incubation of cells with both A-SAA 0.5 μg/ml and D-37pA 5 μg/ml significantly inhibited the induction of VCAM-1 expression to 27 + 5% cells (P = 0.002, Figure 6B). In contrast, co-incubation of HMVECs with A-SAA and the control scrambled peptide L3D-37pA did not attenuate the response of HMVECs to A-SAA (Figure 6C). A dose-dependent inhibition of A-SAA–induced VCAM-1 expression on HMVECs by both D-37pA and L-37pA was demonstrated when used at both 10- and 20-fold excess. No dose-dependent inhibition was seen with the control peptide L3D-37pA (Figure 6D). A-SAA 0.5 μg/ml stimulation for 6 hours significantly increased ICAM-1 from 63 + 0.8% to 90 + 0.7% of cells. Co-incubation of cells with both A-SAA 0.5 μg/ml and L-37pA 5 μg/ml significantly inhibited the induction of ICAM-1 expression to 76 + 0.7% cells (P < 0.001, Figure 6E). Co-incubation of cells with both A-SAA 0.5 μg/ml and D-37pA 5 μg/ml significantly inhibited the induction of ICAM-1 expression to 80 + 0.9% cells (P = 0.02, Figure 6F).

Discussion

RA is a chronic progressive autoimmune disease characterized by proliferation of synovial membrane with deg-
cytokines20,39 – 45 and has been implicated in a wide range of pathological processes including arthritis, bowel disease, malignancy, atherosclerosis, the metabolic syndrome, and type II diabetes.46 –50 We have previously demonstrated that A-SAA promotes angiogenesis, leukocyte recruitment, and matrix degradation,17 key processes in the pathogenesis of RA. It is known that FRP-1/LXA4R protein mediates the pro-inflammatory effects of A-SAA, however recent evidence demonstrates that A-SAA is a potent ligand for SR-B1.23 Together these studies provide strong evidence for an important regulatory role of A-SAA in inflammation and in the immune response.

The current study demonstrates expression of the SR-B1 scavenger receptor in the inflamed synovial membrane of RA joints, with expression lower in patients with inflammatory OA. In particular, we have demonstrated strong expression of SR-B1 on the synovial vascular endothelium, and the hyperplastic lining layer at the leading edge of the synovial pannus, which invades articular cartilage and bone. Furthermore, using Western blot and immunofluorescent microscopy we confirm the expression of SR-B1 on both primary RA FLCs and HMVECs. This pattern of expression of SR-B1 corresponds closely with results of our previous work, which demonstrated expression of A-SAA in a similar histological distribution. To elucidate the extent to which SR-B1 expression relates to A-SAA responsiveness, A-SAA–induced ICAM-1 was correlated with baseline SR-B1 expression. The linear correlation between these variables suggest that the level of SR-B1 expression is an important determinant of A-SAA–induced pro-inflammatory responses in RA FLCs and HMVECs. In contrast, no significant correlation was found between SR-B1– and TNF–α–induced ICAM-1 expression, suggesting that the A-SAA/SR-B1 effect is specific and not merely a marker of cell responsiveness to general pro-inflammatory stimuli. This observation is further supported by Baranova et al, who demonstrated direct A-SAA/SR-B1 binding in human monocyte cells and showed that SR-B1–overexpressing HeLa cells correlated with higher IL-8 production compared with control cells.23

The FPRL-1 receptor has been implicated as the specific receptor mediating the effects of A-SAA in several cell types, however we demonstrate in primary human RA synovial cells for the first time that A-SAA may function through an alternative pathway via SR-B1. To date a direct role for SR-B1 in mediating A-SAA–induced pro-inflammatory effects has not been examined in RA FLCs or HMVECs. To examine this functional relationship we used a specific monoclonal antibody to the SR-B1 receptor. A HMVEC line and four separate primary RA FLC lines were stimulated with A-SAA in the presence or absence of anti–SR-B1, which significantly inhibited A-SAA–induced IL-6 and IL-8 production. Finally, D-37pA and L-37pA, which are homologous to the amphipathic helical region of ApoA-1 (antagonists of SR-B1), inhibited A-SAA-induced adhesion molecule (ICAM-1 and VCAM-1) expression on the HMVEC line. ApoA-1 has previously been shown to be anti-inflammatory by preventing IL-1β and TNF-α secretion from monocytes cells though inhibition of contact mediated activation by T lymphocytes.51 The ApoA-1 mimetic peptide L-37pA acts as a ligand for SR-B1, where it has been shown to block the pro-inflammatory cytokine response to LPS in THP-1 cells.26 To date, however, no direct anti-inflammatory role for ApoA-1 through interaction with SR-B1 has been demonstrated in vivo. Our observation that a specific anti-SR-B1 antibody inhibits A-SAA–induced IL-6 and IL-8...
production provides further evidence that SR-B1 may be an important determinate of the innate inflammatory response in vivo. Further work investigating a broader range of pro-inflammatory responses is now required to confirm both whether other A-SAA-mediated effects occur through SR-B1 and whether the actions of the ApoA-1 mimetic peptides presented here are reflective of an immunomodulatory role for ApoA-1 in vivo.

The evidence that induction of IL-6 may result after stimulation of RA tissue with A-SAA is novel. IL-6 is a key cytokine in the circulation of patients with RA, and recently has been the focus of a new targeted biological therapy. IL-6 had previously been shown to be an inducer of A-SAA, and is synthesized by lymphocytes and RA FLCs, and is detectable in inflamed synovial tissue. Reciprocal upregulation between A-SAA and IL-1 in inflammation has been reported before. Treatment with IL-1 receptor antagonist (IL-1Ra) or with a soluble IL-6 receptor (IL-6R) have both demonstrated efficacy in ameliorating the clinical course of RA.

Recent evidence has demonstrated that A-SAA activates Receptor for Advanced Glycation Endproducts (RAGE) in synovial cells, resulting in activation of the NF-κB signaling pathway. Furthermore, Toll-like Receptor 2 (TLR2) has recently been described as a functional receptor for A-SAA, leading to increased phosphorylation of MAPKs and degradation of IkBα. Together, these findings suggest an important role for A-SAA in the pro-inflammatory cascade of events, interlinked with cytokines, closely associated with synovial membrane inflammation, tissue destruction, and permanent joint damage in RA.

In addition to the well-characterized joint destruction seen in this disease, RA has a well-recognized association with significantly higher rates of cardiovascular mortality. SR-B1 overexpression has previously been demonstrated in atheromatous plaques compared with normal vascular endothelium. Increased cardiovascular risk has been independently associated with elevated levels of systemic inflammatory markers including A-SAA. As part of the systemic inflammatory response A-SAA displaces ApoA-1 from HDL to become the major HDL-associated apoprotein in inflammation. In vitro studies have demonstrated that A-SAA–enriched HDL may be pro-atherogenic through a three- to fourfold higher affinity for macrophages and an increased affinity for uptake into vascular ECs when compared with unenriched HDL. A-SAA has been implicated in a number of diseases characterized both by inflammation and increased cardiovascular mortality, including diabetes, the metabolic syndrome, malignancy, and primary cardiovascular disease. It is intriguing to speculate that A-SAA/SR-B1 interactions may, in addition to a role in joint inflammation, also have a role in the long-term cardiovascular complications described in RA.

In summary, we have demonstrated for the first time the expression of the HDL receptor, SR-B1 in inflamed RA synovial tissue. We have shown that pro-inflammatory responses by A-SAA correlate closely with SR-B1 expression on RA FLCs in vitro, and that the effects of A-SAA stimulation can be down-regulated by SR-B1 antagonist mimetic peptides and a specific anti–SR-B1 antibody. Finally, we have identified new cytokine interactions with A-SAA, which may be attenuated by blocking SR-B1. These findings suggest that modulation of inflammation through SR-B1 may provide a future therapeutic strategy for the treatment of RA and its associated cardiovascular risk.

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