Blockade of endogenous interleukin 12 results in suppression of murine streptococcal cell wall arthritis by enhancement of interleukin 10 and interleukin 1Ra

Leo A B Joosten, Monique M A Helsen, Wim B van den Berg

Abstract

Objective—The goal of this study was to investigate the role of endogenous interleukin 12 (IL12) in acute murine streptococcal cell wall (SCW) arthritis.

Methods—C57black/6 mice were injected intraperitoneally with rat anti-murine IL12 (C17.8), shortly before induction of arthritis by intra-articular injection of 25 µg SCW fragments into the right knee joint. Joint swelling and chondrocyte synthetic function was analysed several days after induction of SCW arthritis. Local cytokine profile was determined, protein by using ELISA and mRNA by RT-PCR technology. To confirm the findings at later time points, tissue chamber model of inflammation was used. Histology was performed to examine cell influx and cartilage damage.

Results—Suppression of joint swelling was noted at days 2 and 4, whereas no suppressive effect of anti-IL12 was found at day 1. Severe inhibition of chondrocyte proteoglycan synthesis was seen at day 1 in both arthritic control and anti-IL12 treated mice. However, chondrocyte function was restored at day 4 of arthritis in the anti-IL12 injected animals, but not in the arthritic controls. Moreover, cell influx in synovial tissue and joint cavity was reduced by anti-IL12 treatment. Neutralisation of IL12 reduced the local levels of IL18, IL12 and interferon γ, when examined shortly after induction of SCW arthritis, whereas tumour necrosis factor a levels were not affected. In contrast, IL10 and IL1Ra protein and mRNA levels were strongly up regulated in synovial tissues after IL12 blockade. Enhancement of IL10 and IL1Ra by anti-IL12 was confirmed in a tissue chamber model with SCW induced inflammation.

Conclusions—This study indicates that IL12 is a pro-inflammatory cytokine during onset of acute SCW arthritis. Balances of proinflammatory and anti-inflammatory cytokines were strongly improved by anti-IL12 treatment.

Interleukin 12 (IL12), originally described as cytotoxic lymphocyte maturation factor and natural killer cell stimulatory factor is a 70 kDa heterodimeric cytokine consisting of two subunits of 35 kDa and 40 kDa. It is produced by phagocytic cells, dendritic cells, B-lymphocytes and NK cells upon stimulation with lipopolysaccharide (LPS), bacteria, and parasites. IL12 generates the development of naïve T cells into Th1 cells and stimulates interferon γ (IFNγ) secretion by differentiated Th1 cells. Through major induction of IFNγ and tumour necrosis factor α (TNFα), IL12 plays a crucial part in LPS lethal shock and generalised Schwartzman reaction. During bacterial infections, IL12 is considered to be a principal protective cytokine, where it bridges innate resistance and antigen specific immunity. Recently, it was demonstrated that IL12 deficient mice are unable to control bacterial infections with Leishmania major and were defective in IFNγ production and Th1 responses. In contrast, IL12 may unmask Th1 dependent autoimmune responses and may be a crucial intermediate in the often suggested link between bacterial infections and expression of autoimmune diseases. For instance arthritis has been associated with bacterial infections of the throat (rheumatic fever), the gastrointestinal tract (reactive arthritis), inflammatory bowel disease like Crohn’s disease, and Lyme disease. In all of these cases IL12, induced by the bacterial organisms, may be the pivotal cytokine involved in the onset of arthritis. Rheumatoid arthritis (RA) is associated with Th1 immune response, however typical Th1 cytokines, like IL2, IFNγ and TNFβ were hardly detected in synovial fluid or tissues. Recently, IL12 was detected in synovia of RA patients, and macrophage-like cells seems to be the main producers.

Several bacterial arthritis models have been described and beside adjuvant arthritis (AA), induced by M. tuberculosis, streptococcal cell wall (SCW) arthritis is one of the important experimental animal models. Chronic arthritis can be induced in susceptible rat strains by an intraperitoneal injection of a sterile solution of bacterial cells or cell wall fragments from the group A streptococci. A reactivation model of local SCW arthritis, induced by intra-articular injection of SCW into a rat ankle joint, can be achieved by intravenous rechallenge with SCW fragments. SCW arthritis can also be induced in mice after immunisation with SCW fragments in complete Freund’s adjuvant. After intra-articular injection of SCW fragments into the knee joint a chronic erosive arthritis will occur. In contrast, one single injection of SCW fragments into a knee joint of a naïve mouse
leads to an acute joint inflammation. It has been shown that TNFα and IL1 play different parts in the latter SCW arthritis model. While TNFα mediates only joint swelling, IL1 is crucial regarding cartilage destruction and inflammatory cell influx. Furthermore, it was demonstrated that endogenous IL10 controls the severity of SCW arthritis, because anti-IL10 treatment aggravated SCW arthritis. Later on, at days 4 and 7 macrophage-like cells were noted in the synovial tissue. The maximum influx of inflammatory cells was seen at days 2 and 4 after the induction of SCW arthritis. 

In this study, we investigated whether IL12, produced by synovial macrophages after injection of bacterial cell wall fragments, is involved in the onset and severity of acute SCW arthritis. Therefore we injected anti-murine IL12 antibodies before intra-articular application of SCW fragments. Joint swelling was monitored and joint pathology was examined by histology. Furthermore, local cytokine profile was determined by ELISA and RT-PCR technology. Because of the fact that synovial cytokine levels were only detectable in the first 24 hours of acute SCW arthritis, we further analysed the effect of endogenous IL12 blockade in a tissue chamber model of SCW inflammation. This study indicates that blockade of IL12 during onset of acute SCW arthritis suppresses secondary joint swelling, cell influx and restoration of chondrocyte PG synthesis by up-regulation of IL10 and IL1Ra levels.

Methods

Animals

Male C57/Bl6 mice were obtained at our university breeding facilities in Nijmegen. The mice were housed in filter top cages, and water and food were provided ad libitum. The mice were used at the age of 10–12 weeks.

Materials

Ethidiumbromide, rat Igs, and bovine serum albumin were purchased from Sigma Chemicals, St Louis, USA. DNEM medium, RPMI 1640 medium, FCS, Taq DNA Polymerase, 100 bp DNA marker, TRIzol Reagent, and Agarose were obtained from Life Technologies, Breda, the Netherlands. GAPDH, IL1β, IL10, IL12, IFNγ and TNFα primers were purchased from Eurogentec, Seraing, Belgium. Murine IL1Ra capture (MAP480) and detection (BAF480) antibodies and recombinant murine IL1Ra were obtained from R&D Systems, Abingdon, UK. IL10 (CMC9102) and INFγ (CMC4034) ELISA cytosets were obtained from Biosource, Camarillo, CA, USA. Anti-mIL12 antibodies (capture 80–3891–01, detection 80–3892–01) were purchased from Genzyme, Cambridge, MA, USA. Murine IL12 was kindly provided by Dr S Wolf, Genetic Institute Inc, Cambridge, MA, USA.

Anti-IL12 Antibodies

Hybridoma cells producing rat anti-murine IL12 antibodies (C17.8) were kindly provided by G Trinchieri, Wistar Institute, Philadelphia, USA. Hybridoma cells (3.106) were injected in nude Balb/C mice and after three weeks ascites fluid was collected. Thereafter Igs were isolated using a protein-G column. Anti-murine IL12 antibodies (0.5 mg) were injected in a volume of 0.2 ml phosphate buffered saline (PBS).

SCW Preparation and Induction of SCW Arthritis

Streptococcus pyogenes T12 organisms were cultured overnight in Todd-Hewitt broth. Cell walls were prepared as described previously. The resulting 10,000 × g supernatant was used throughout the experiments. These preparations contained 11% muramic acid. Unilateral arthritis was induced by intra-articular injection of 25 µg SCW (rhamnose content) in 5 µl

Figure 1 SCW arthritis, joint swelling, chondrocyte proteoglycan (PG) synthesis, and cytokine profile. (A) SCW arthritis was induced by injection of 25 µg SCW fragments. Joint swelling was measured by 99mTc uptake method and is expressed as right/left ratio. A ratio >1.10 is considered as inflammation. Inhibition of chondrocyte PG synthesis was determined by 35S-sulphate incorporation. The data represent the mean (SD) of at least seven mice per group. (B) Cytokine levels were determined either by RIA or ELISA in patellae washouts at several time points. The data represent the mean cytokine level of washouts of six patellae per time point.
PBS into the right knee joint of naive mice. As a control, PBS was injected into the left knee joint.

**ANTI-IL12 TREATMENT**

Two hours before induction of acute SCW arthritis, mice were given an intraperitoneal injection of 0.5 mg purified rat anti-murine IL12. As previously described, this dose has been shown to be effective in vivo. As control the same amount of normal rat IgG was injected. For time course experiments, mice were injected intraperitoneally at −2 hours, day 1 and day 2 after the induction of acute SCW arthritis. To neutralise IL12 during SCW inflammation in the tissue chambers, mice were intraperitoneally injected with anti-IL12 two hours before injection of SCW fragments into the tissue chambers.

**MEASUREMENT OF JOINT INFLAMMATION**

SCW arthritis was quantified by the $^{99m}$Tc uptake method. This method measures by external gamma counting the accumulation of a small radioisotope at the site of inflammation attributable to local increased blood flow and tissue swelling. The severity of inflammation is expressed as the ratio of the $^{99m}$Tc uptake in the right (inflamed) over the left (control) knee joint. All values exceeding 1.10 were assigned as inflammation.

**CHONDROCYTE PROTEOGLYCAN SYNTHESIS DETERMINATION**

Patellae with minimal surrounding tissue, were placed in RPMI 1640 medium with glutamax, gentamicin (50 µg/ml) and $^{35}$S-sulphate (0.74 MBq/ml). After three hours of incubation at 37°C in a carbon dioxide incubator, patellae were washed in saline three times, fixed in 4% formaldehyde and subsequently decalcified in 5% formic acid for four hours. Patellae were punched out of the adjacent tissue, dissolved in 0.5 ml Luma Solve at 65°C (Omnilabo) and after addition of 10 ml Lipolula (Omnilabo) the $^{35}$S content was measured by liquid scintillation counting. Values are presented as percentage $^35$S incorporation of the left control joint.

**CYTOKINE PRODUCTION IN A MOUSE TISSUE CHAMBER MODEL OF INFLAMMATION**

Autoclaved sterile Teflon (Teflon-PFA, kind gift of Novartis, Basel, Switzerland) tissue chambers (20 mm × 8 mm diameter perforated by 135 regularly spaced 1 mm holes) were implanted subcutaneously under aseptic conditions through a small incision into the backs of mice. The incisions were closed by wound clips and the chambers were tested for sterility after seven days by culturing a 10 µl sample on LB agar for 24 hours at 37°C. Thereafter SCW inflammation was induced by injection of 100 µl SCW fragments (4 mg/ml rhamnose content). At days 1 and 2 after induction of inflammation samples of tissue chambers fluid (±200 µl) were taken. The samples were centrifuged for five minutes at 1000 g and stored at −80°C until cytokine determination.

**CYTOKINE MEASUREMENTS**

To determine the levels of IL1β, IL10, IL12, IFNγ or TNFα in patellae washouts, patellae were isolated from inflamed knee joints as previously described. Patellae were cultured in RPMI 1640 medium (200 µl/patella) for one hour at room temperature. Thereafter, supernatant was harvested and centrifuged for five minutes at 1000 g. IL1β and TNFα were determined by RIA. IL1Ra, IL10, IL12 and IFNγ levels was measured by ELISA. The sensitivity of the ELISA for IL1Ra, IL10, IL12, IFNγ and the RIA for IL1β and TNFα is 80, 40, 20, 10, 20 and 40 pg/ml, respectively. The same assays were used for determination of cytokine levels in tissue chamber fluid samples.

**RNA ISOLATION**

Mice were killed by cervical dislocation, immediately followed by dissection of the patella with adjacent synovium. From six patella specimens synovium biopsy specimens were taken. Two biopsy specimens with a diameter of 3 mm were punched out, using a biopsy...
punch (Stiefel, Wachtersbach, Germany): one from the lateral and one from the medial side. Six patella specimens per experimental group were taken and three lateral and three medial biopsy specimens were pooled, to yield two samples per group. The synovium samples were immediately frozen into liquid nitrogen. Synovium biopsy specimens were ground to powder using a micro-dismembrator II (B Braun, Melsungen, Germany). Total RNA was extracted in 1 ml TRIzol reagent, an improved single step RNA isolation method based on the

Table 1  Joint pathology after neutralisation of endogenous IL12

<table>
<thead>
<tr>
<th>Days of arthritis</th>
<th>Treatment</th>
<th>Cell influx synovial tissue</th>
<th>Cell influx joint cavity</th>
<th>Matrix PG depletion</th>
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</thead>
<tbody>
<tr>
<td>Day 2</td>
<td>Control (Igs)</td>
<td>0.8 (0.2)</td>
<td>1.8 (0.4)</td>
<td>1.0 (0.2)</td>
</tr>
<tr>
<td>Day 2</td>
<td>Anti-IL12</td>
<td>0.4 (0.3)</td>
<td>0.7 (0.3)*</td>
<td>0.9 (0.3)</td>
</tr>
<tr>
<td>Day 4</td>
<td>Control (Igs)</td>
<td>1.3 (0.3)</td>
<td>1.9 (0.4)</td>
<td>1.7 (0.4)</td>
</tr>
<tr>
<td>Day 4</td>
<td>Anti-IL12</td>
<td>1.0 (0.3)</td>
<td>0.6 (0.3)*</td>
<td>1.2 (0.3)</td>
</tr>
</tbody>
</table>

Mice were given an intraperitoneal injection injected with either rat Igs (0.5 mg) or anti-IL12 (0.5 mg) two hours before induction of an unilateral arthritis by intra-articular injection of 25 µg SCW fragments into the right knee joint. Histology was taken at days 2 and 4 of SCW arthritis and performed as described in Methods. Data represent the mean (SD) of two identical experiments, each experiment consisted of at least seven mice per group. *p<0.05, Mann-Whitney U test compared with control.

Figure 3  Effect of IL12 blockade on cell influx and cartilage damage in the knee joint. (A) Arthritic knee joint of the rat Igs treated control group. Note the marked joint inflammation and cell influx in the joint cavity. Bar=80 µM. (B) Section of the control group stained for matrix proteoglycan. Note the loss of matrix PG in the cartilage layer (arrows). Bar=40 µM. (C) Knee joint of a mouse treated with anti-IL12 antibodies. Strong reduction of cell influx can be seen in synovial tissue and joint cavity. Bar=80 µM. (D) Section of anti-IL12 treated knee joint stained for PG loss. No reduction of matrix PG depletion was seen after IL12 blockade. Bar=40 µM. In all experiments, (A and C) Haematoxylin and eosin staining, (B and D) Safranin O staining. All specimens were taken at day 2 after induction of SCW arthritis.
method described by Chomczynski and Sacchi.26

PCR AMPLIFICATION
One microgram of synovial RNA was used for RT-PCR. Messenger RNA was reverse transcribed to cDNA using oligoT primers and 1/20 of the cDNA was used in one PCR amplification. PCR was performed at a final concentration of 200 mM dNTPs, 0.1 mM of each primer, and 1 unit Taq-polymerase in standard PCR buffer. The mixture was overlaid with mineral oil and amplified in a thermocycler (Omnigene, Hybaid Ltd, UK). Message for GAPDH, IL1β, IL10, IL12, IFNγ and TNFα was amplified using the primers as previously described.27 Samples of 5 µl were taken from the reaction tubes after a certain number of cycles. The cycle curves were performed by two cycle intervals. PCR products were separated on 1.6% agarose and stained with ethidiumbromide. The results are presented as differences in PCR cycles between mRNA isolated from inflamed synovium compared with normal synovium. GAPDH levels were very consistent in all samples, not warranting any further correction of mRNA values for IL1β, IL10, IL12, IFNγ and TNFα.

HISTOLOGICAL EXAMINATION
Mice were killed by ether anaesthesia. Thereafter, whole knee joints were removed and fixed for four days in 4% formaldehyde. After decalcification in 5% formic acid the specimens were processed for paraaffin wax embedding. Tissue sections (7 mm) were stained with haematoxylin and eosin or safranin O. Histopathological changes were scored using the following parameters. Infiltration of cells was scored on a scale of 0–3, depending on the amount of
inflammatory cells in the synovial cavity and synovial tissues. Proteoglycan depletion was determined using safranin O staining. The loss of proteoglycans was scored on a scale of 0–3, ranging from full stained cartilage to destained cartilage or complete loss of articular cartilage. Histopathological changes in the knee joints were scored in the patella/femur region on five semi-serial sections of the joint, spaced 70 mM apart. Scoring was performed on decoded slides by two observers, as described elsewhere.

**STATISTICAL ANALYSIS**

Differences between experimental groups were tested using the Mann-Whitney U test, unless stated otherwise.

**RESULTS**

**MURINE SCW INDUCED ARTHRITIS**

One single injection of SCW fragments into a mouse knee joint leads to an acute inflammation, characterised by joint swelling and inhibition of chondrocyte proteoglycan (PG) synthesis in patellar cartilage. Significant knee joint swelling was found up to day 7 after injection of SCW fragments. Marked inhibition of chondrocyte PG synthesis was observed at days 1, 2 and 4, whereas at day 7 an overshoot of chondrocyte PG synthesis was noted (fig 1A).

To examine the cytokine pattern during the first stage of this type of experimental arthritis we analysed synovial tissue washouts (fig 1B). TNFα and IL1β levels were increased shortly after injection of SCW fragments. At 90 minutes the highest levels (mean (SD) 450 (40) pg/ml) of TNFα were found, whereas the maximum IL1β levels were found at four hours (2520 (280) pg/ml). IL12 levels were detectable at 1.5, 4, 24 and 48 hours after induction of arthritis, with the peak at four hours (160 (30) pg/ml). Protein levels of IFNγ reached a maximum at four hours, 50 (15) pg/ml. As described previously, massive influx of polymorphonuclear cells (PMNs) in the joint cavity and synovial membrane was seen at days 1 and 2.

**REDUCTION OF SCW ARTHRITIS BY BLOCKING ENDOGENOUS IL12**

To investigate the effect of neutralisation of endogenous IL12 on acute SCW arthritis, we gave an intraperitoneal injection of rat anti-murine IL12 antibodies two hours before injection of SCW fragments. At day 1 no reduction of joint swelling was found. In contrast, marked suppression of joint swelling was seen at days 2 and 4 (fig 2A). Severe inhibition of chondrocyte PG synthesis was noted at day 1 in the arthritis control group treated with rat Igs. Blocking of IL12 had no
significant effect on chondrocyte PG synthesis at day 1. In line with the reduction of joint swelling at days 2 and 4, anti-IL12 treatment increased the chondrocyte PG synthesis (fig 2B). At day 4, the chondrocyte PG synthesis reached values found in normal cartilage (left joint), whereas in the arthritic control group (rat Iggs) 30% inhibition was seen. Histological examination showed that anti-IL12 treatment significantly reduced the number of inflammatory cells in the joint cavity at day 2 of SCW arthritis (fig 3A/C, table 1). Infiltration of synovial membrane with inflammatory cells was reduced by IL12 blocking although this did not differ significantly from the control group. No significant effect of anti-IL12 was seen on cartilage PG depletion as depicted in figure 3B/D. Marked loss of PG was found in both control and anti-IL12 groups (table 1).

KINETICS OF ANTI-IL12 INDUCED SUPPRESSION OF SCW ARTHRITIS

To examine whether IL12 is only crucial at the onset of acute SCW arthritis or is involved as well at later stages of this type of joint inflammation we injected anti-IL12 at several time points. As shown above anti-IL12 treatment before induction of SCW arthritis suppressed both joint swelling and inhibition of chondrocyte PG synthesis (fig 4A/B). When anti-IL12 treatment was delayed until day 1 of arthritis we found nearly the same effects on joint swelling and chondrocyte function as treatment before induction of arthritis. No suppressive effect on joint swelling or inhibition of chondrocyte PG synthesis was observed anymore when anti-IL12 was given at day 2.

EFFECT OF IL12 BLOCKADE DURING SCW ARTHRITIS ON CYTOKINE PATTERN

Analysis of local cytokine production at 1.5 and 4 hours after induction of arthritis showed that TNFα levels were not changed after anti-IL12 treatment, whereas IL1β, IL12 and IFNγ levels were significantly reduced (fig 5A/B). Of great interest, IL10 and IL1Ra levels were markedly increased because of IL12 elimination (fig 5C). RT-PCR analysis of synovium biopsy specimens showed that mRNA levels of TNFα, IL1β, IL1Ra, IL10, IL12 and IFNγ were up regulated at days 1 and 2 of SCW induced arthritis, when compared with normal (non-inflamed) synovium (fig 6). Anti-IL12 treatment slightly decreased mRNA levels for IL1β and TNFα, but marked reduction was found of mRNA levels for IFNγ, IL12 and TNFα (figs 6, 7). This was mainly noted at day 2 after treatment with anti-IL12. In line with cytokine levels in synovium washouts, both IL1Ra and IL10 mRNA levels were increased at day 1 and 2 of SCW arthritis after blocking endogenous IL12. When ratios of relative mRNA levels were calculated a strong shift in IL1Ra/IL1β and IL10/TNFα balances was found (fig 6G/H).

NEUTRALISATION OF IL12 DURING SCW INFLAMMATION IN A TISSUE CHAMBER MODEL RESULTED IN INCREASED LEVELS OF IL10 AND IL1Ra

As we cannot detect large amounts of cytokines in patella washouts at days 1 and 2 after induction of SCW, we analysed the impact of IL12 blockade on cytokine levels in a tissue chamber model. High levels (ng/ml) of various cytokines could be measured after injection of SCW fragments in a tissue chamber (fig 8). Anti-IL12 treatment of mice bearing such tissue chambers showed that neutralisation of IL12 reduced IL1β, IFNγ and IL12 levels, whereas TNFα levels were not changed (fig 8B, C, D, A). In line with mRNA and protein levels in synovial tissue, IL10 and IL1Ra levels were up regulated after anti-IL12 treatment (fig 8F). Table 2 shows that ratios of either mRNA or protein for IL1Ra/IL1β and IL10/TNFα were increased because of anti-IL12 treatment. This shows that the balance of anti-inflammatory (for example, IL1Ra, IL10) and

<table>
<thead>
<tr>
<th>Day of SCW arthritis</th>
<th>Treatment</th>
<th>Protein level*</th>
</tr>
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<tbody>
<tr>
<td>Day 1</td>
<td>Control (Igs)</td>
<td>0.64</td>
</tr>
<tr>
<td>Day 1</td>
<td>Anti-IL12</td>
<td>2.27</td>
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<tr>
<td>Day 2</td>
<td>Control (Igs)</td>
<td>0.60</td>
</tr>
<tr>
<td>Day 2</td>
<td>Anti-IL12</td>
<td>3.55</td>
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Cytokine ratios were calculated after treatment with either rat Igs or anti-IL12. *For protein ratios we used levels of cytokine found in the tissue chamber model of inflammation (see fig 8).
Role of endogenous interleukin 12 in murine streptococcal cell wall arthritis

Discussion

SCW arthritis is an experimental model of arthritis in which macrophages play an important part. It has been shown that bacterial cell wall fragments (peptidoglycan) induced production of IL12 by macrophages in vitro.29

Here we showed that endogenous IL12 is involved in the persistence of acute SCW induced arthritis. Suppression of joint swelling was found at days 2 and 4 and not at day 1 after intra-articular injection of SCW fragments. This shows that IL12 is not involved in joint swelling seen at the initial stage. As shown previously, TNFα is the major cytokine regarding acute joint swelling in SCW arthritis.18 30 The lack of effect at day 1 is in line with the fact that shortly after onset of acute SCW arthritis TNFα levels were not affected by anti-IL12 treatment. Moreover, TNFα levels were not influenced by anti-IL12 treatment at later time points both in a tissue-chamber model of inflammation. The pivotal role of TNFα in the early joint swelling was corroborated by elegant studies of SCW arthritis in TNFα deficient mice.31 The major role of TNFα in early joint swelling was also noted in murine collagen induced arthritis.29 32 The suppressive effect of anti-IL12 on joint swelling in sub-acute stages of arthritis could be explained on the one hand by the reduced levels of IFNγ, IFNγ itself being involved in joint swelling and on the other hand reduction of NO.33 In respect to joint swelling, endogenous IL12 is involved between 24–48 hours after onset of SCW arthritis, but seems again less dominant at later stages, as anti-IL12 treatment started at day 2 was ineffective.

Inhibition of chondrocyte PG synthesis in the arthritic cartilage is a common feature seen in experimental arthritis models and IL1 is the pivotal cytokine in this inhibition.34 In this study it was shown that elimination of IL12 during onset results in restoration of chondrocyte function at days 2 and 4 of acute SCW arthritis. IL1β levels were reduced by anti-IL12 treatment, but significant levels were still found in synovial tissue washouts. At days 1 and 2 reduction of IL1 activity was also observed in tissue chamber fluid after IL12 blockade. This was in line with findings in tumour bearing mice treated with anti-IL12 antibodies in which decreased IL1β levels were reported.35 Recently, it was demonstrated that chondrocyte PG synthesis induced by IL1 was mediated by NO as NOS2 deficient mice were found at days 2 and 4 and not at day 1 after injection of SCW. IL10 is the major cytokine regarding acute joint swelling and on the other hand cytokines is clearly improved after elimination of endogenous IL12, also at later time points. It has been shown that bacterial cell wall fragments (peptidoglycan) induced production of IL12 by macrophages in vitro.29

Here we showed that endogenous IL12 is involved in the persistence of acute SCW induced arthritis. Suppression of joint swelling was found at days 2 and 4 and not at day 1 after intra-articular injection of SCW fragments. This shows that IL12 is not involved in joint swelling seen at the initial stage. As shown previously, TNFα is the major cytokine regarding acute joint swelling in SCW arthritis.18 30 The lack of effect at day 1 is in line with the fact that shortly after onset of acute SCW arthritis TNFα levels were not affected by anti-IL12 treatment. Moreover, TNFα levels were not influenced by anti-IL12 treatment at later time points both in a tissue-chamber model of inflammation. The pivotal role of TNFα in the early joint swelling was corroborated by elegant studies of SCW arthritis in TNFα deficient mice.31 The major role of TNFα in early joint swelling was also noted in murine collagen induced arthritis.29 32 The suppressive effect of anti-IL12 on joint swelling in sub-acute stages of arthritis could be explained on the one hand by the reduced levels of IFNγ, IFNγ itself being involved in joint swelling and on the other hand reduction of NO.33 In respect to joint swelling, endogenous IL12 is involved between 24–48 hours after onset of SCW arthritis, but seems again less dominant at later stages, as anti-IL12 treatment started at day 2 was ineffective.

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Blockade of IL1 activity by application of IL1Ra completely prevents inhibition of chondrocyte PG synthesis during acute SCW arthritis as described previously.42 Apart from reduction of the cytokines IL1, IFNγ and NO the restoration of chondrocyte PG synthesis could be the result of enhancement of IL1Ra and IL10 levels by anti-IL12. IL12 exposure to mononuclear cells reduced the IL1Ra production by these cells in vitro.43 Administration of IL10 during acute SCW arthritis was shown to increase chondrocyte function whereas anti-IL12 aggravated inhibition of PG synthesis at days 2 and 4 after injection of SCW.44 IL10 is an inhibitor of IL1 and NO production and it up regulates IL1Ra production in several cell types. Reduction of IL1β, IL12 and IFNγ by IL12 blockade was not only found shortly after onset of acute SCW arthritis, but also at later stages (days 1 and 2). The latter was examined in a tissue-chamber model of inflammation. As shown previously, high levels of several cytokines could be detected in the fluid that can be easily isolated from the chamber.22 We found some differential effect of IL12 in synovium and tissue chambers. This could be attributable to the fact that synovial cytokine levels were measured during onset, and in tissue chambers cytokines were determined in established SCW inflammation.

Histological examination showed that neutralisation of IL12 increased the number of inflammatory cells in joint tissues. Recently, it has been reported that overexpression of IL12 with an adenoviral vector leads to upregulation of proinflammatory chemokines, including MIP-1α, MIP-2 and MCP-1.45 These chemokines direct leucocyte migration into inflamed tissue and blockade of IL12 may lead to decreased expression. Furthermore, it was shown that IL10 down regulated ICAM expression and anti-IL10 treatment of collagen induced arthritis increased expression of MIP-1α and MIP-2.46 47 Therefore, the reduced number of inflammatory cells in the joint might also be explained by markedly increased IL10 and decreased IL12 levels in the anti-IL12 treated animals. Finally, IL1 itself induces production of pro-inflammatory chemokines and expression of adhesion molecules.48 A shift in the balance of IL1Ra/IL1β was noted after IL12 elimination, and this may also contribute to reduce cell influx.

Recently, three pathways of IL12 production by either macrophages or dendritic cells have been described.49 Direct stimulation by bacterial agents, triggering by T cell dependent immune responses and activation by the matrix glycosaminoglycan hyaluran.50 The latter is mediated by binding of low molecular...
fragments of matrix proteoglycans to adherence surface molecule CD44 and these PG fragments are abundantly present in an inflamed joint. As IL12 promotes Th1 responses, it may be possible that IL12 generates a local immune response in the joint towards cartilage autoantigens like CII, gp-39 and aggregan.19 In other models of autoimmune inflammation such as collagen induced arthritis, experimental colitis, diabetes in non-obese diabetic mice and experimental allergic encephalomyelitis, an essential role of IL12 was observed.20−22 Elimination of IL12 in these animal models by anti-IL12 treatment leads to suppression or prevention of disease expression. We recently found that intra-articular IL12 gene transfer with an adenoviral vector aggravates acute SCW arthritis and local IL12 expression can promote conversion of an acute SCW arthritis to a chronic destructive arthritic process (manuscript in preparation). This suggests that IL12 not only plays a part in the onset of arthritis but could also determine the chronicity of the disease. Whether generation of joint specific Th1 response occurred in these animals is still under investigation. This study shows that IL12 is a primary pro-inflammatory cytokine during onset of bacterial induced arthritis and blockade of IL12 ameliorates joint swelling, leads to restoration of chondrocyte function, and reduces influx of leucocytes. Anti-IL12 treatment could be efficacious in the treatment of patients with arthritis.

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5 Seder RA, Gazzinelli RT, Sher A, Paul WE. Interleukin-12 acts directly on CD4+ T-cells to enhance priming for interferon-gamma production and diminishes interleukin-4 inhibition of such priming. Proc Natl Acad Sci USA 1993;90:1088–92.


