Repeated infusions of infliximab, a chimeric anti-TNFα monoclonal antibody, in patients with active spondyloarthropathy: one year follow up

E Kruithof, F Van den Bosch, D Baeten, A Herssens, F De Keyser, H Mielants, E M Veys

Background: In a pilot study, the anti-tumour necrosis factor α monoclonal antibody, infliximab, induced a rapid and significant improvement in global, peripheral, and axial disease manifestations of patients with active spondyloarthropathy.

Objective: To determine whether repeated infusions of infliximab would effectively and safely maintain the observed effect.

Methods: Safety and efficacy of a maintenance regimen (5 mg/kg infliximab every 14 weeks) was evaluated using the measurements reported in the pilot study. Of the 21 patients, 19 completed the one year follow up for efficacy; two patients changed to another dosing regimen after week 12 owing to partial lack of efficacy. However, they are still being followed up for safety analysis.

Results: After each re-treatment a sustained significant decrease of all disease manifestations was observed. Before re-treatment, symptoms recurred in 3/19 (16%) at week 20, in 13/19 (68%) at week 34, and in 15/19 (79%) at week 48. No withdrawals due to adverse events occurred. Twelve minor infectious episodes were observed. Twelve patients (57%) developed antinuclear antibodies; in four of them (19%) anti-dsDNA antibodies were detected. However, no lupus-like symptoms occurred.

Conclusion: In this open study of infliximab in patients with active spondyloarthropathy, the significant improvement of all disease manifestations was maintained over a one year follow up period without major adverse events. Although recurrence of symptoms was noted in a rising number of patients before each re-treatment, no loss of efficacy was observed after re-treatment.

The group of spondyloarthropathies (SpA) comprises related chronic autoimmune disorders of the joint with common clinical, radiological, and genetic characteristics. Entities belonging to this group are ankylosing spondylitis (AS), reactive arthritis, psoriatic arthritis (PsA), undifferentiated SpA (uSpA), arthritis associated with inflammatory bowel disease, and acute anterior uveitis. Juveniles as well as adults can be affected by one of the diseases in the group of SpA. This common rheumatic condition has a global prevalence of 0.5–1%, with, however, important racial and geographical differences and a clear male predominance. Quality of life may be severely compromised by pain and loss of anatomical and functional integrity. The actual therapeutic options for the management of the SpA are still limited and often unsatisfactory. Non-steroidal anti-inflammatory drugs and physiotherapy are considered the cornerstone of the treatment of AS. Patients with peripheral disease manifestations may benefit from sulfasalazine. For PsA, different disease modifying treatments have been suggested to be efficacious, including sulfasalazine, methotrexate, and cyclosporin.

Recently, the use of biological treatments that block tumour necrosis factor α (TNFα) has opened new perspectives for the treatment of patients with SpA. Infliximab (Remicade, Centocor, Malvern, PA, USA) is a chimeric anti-TNFα monoclonal IgG1 antibody that has been shown to be efficacious in treatment resistant Crohn’s disease and rheumatoid arthritis (RA). We note a significant improvement in articular as well as axial disease manifestations in four patients with Crohn’s disease with associated SpA, treated with infliximab. In patients with poly-articular PsA, a clinically significant benefit with infliximab as well as with other TNFα blocking agents was shown. In view of these observations we designed an open label trial with infliximab to investigate the therapeutic potential of TNFα blockade in the different subtypes of SpA.

In this pilot study the effect of a loading dose regimen of three infusions of 5 mg/kg infliximab was evaluated over 12 weeks. The treatment induced a rapid and significant improvement in global, peripheral, and axial disease manifestations in all 21 patients, without major side effects. Simultaneously, successful treatment of active AS was also reported in an open label trial of 11 patients. All patients of the initial open label trial were re-treated every 14 weeks with an infusion of 5 mg/kg infliximab. In this paper we report the safety and efficacy of this maintenance regimen.

PATIENTS AND METHODS

Study design

This study was an extension protocol of the initial 12 week pilot study. The study design consisted of an induction regimen of three infusions of infliximab (5 mg/kg) at weeks 0, 2, and 6, followed by a maintenance regimen every 14 weeks with the same dose. The study protocol, the information for the patients, and the consent form were approved by the ethics committee of Ghent University Hospital. All patients enrolled in the trial signed an informed consent form before any study related procedure.

Patient selection criteria

Patients with active SpA, fulfilling the European Spondylarthropathy Study Group criteria for SpA, were enrolled in the
The open pilot study.

Evaluation of efficacy was performed at weeks 20, 22, 28, 34, and 50 using the same measurements as reported in the previous study.

Drug administrations

Infliximab at a dose of 5 mg/kg in 250 ml NaCl 0.9% was prepared by the hospital pharmacy. The total dose was given over a period of at least two hours. Aseptic procedures were used during the preparation and administration of the study drug. Patients were re-treated every 14 weeks (weeks 20, 34, and 48).

Clinical evaluation

Evaluation of efficacy was performed at weeks 20, 22, 28, 34, 36, 42, 48, and 50 using the same measurements as reported in the open pilot study. In addition, patients were asked at each visit about recurrence of symptoms, including occurrence of night pain, morning stiffness, pain in peripheral joints, or axial pain.

Safety evaluation

During the infusion the patient was monitored for adverse effects, and vital parameters (blood pressure, pulse, temperature) were measured. At each visit patients were asked about side effects, and routine laboratory tests were performed, which consisted of a full blood count with white blood cell differentiation, and biochemical screening, including electrolytes, urea, creatinine, total protein, liver function tests, and urine analysis. Furthermore, screening for antinuclear antibodies (ANA) by indirect immunofluorescence on HEp-2 cells and, when positive, additional testing for anti-double stranded DNA antibodies (anti-dsDNA) by the Crithidia luciliae assay test was performed on each visit. ANA and anti-dsDNA were considered positive in a patient when present on at least two occasions.

Statistical analysis

Results were presented as the median and range. Significance of the change from baseline was measured by the Wilcoxon signed ranks test (p<0.05).

RESULTS

Table 1 shows the demographic and descriptive characteristics of the study group. Values are given as median and range.

<table>
<thead>
<tr>
<th>Total (n=21)</th>
<th>AS (n=3)</th>
<th>AS + PA (n=7)</th>
<th>PsA (n=9)</th>
<th>uSpA (n=2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>49 (26–73)</td>
<td>33 (26–65)</td>
<td>44 (28–59)</td>
<td>49 (30–73)</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>17/4</td>
<td>3/0</td>
<td>6/1</td>
<td>6/3</td>
</tr>
<tr>
<td>Disease duration (years)</td>
<td>17 (1–42)</td>
<td>15 (9–40)</td>
<td>17 (10–33)</td>
<td>19 (1–42)</td>
</tr>
<tr>
<td>Swollen joint count [0–66]</td>
<td>3 (0–12)</td>
<td>0</td>
<td>2 [1–4]</td>
<td>9 (1–12)</td>
</tr>
<tr>
<td>Axial morning stiffness (min)</td>
<td>–</td>
<td>90 (60–120)</td>
<td>90 (30–240)</td>
<td>–</td>
</tr>
<tr>
<td>Axial night pain (0–3)</td>
<td>–</td>
<td>3 (2–3)</td>
<td>3 [1–3]</td>
<td>–</td>
</tr>
<tr>
<td>Sedimentation rate (mm/1st h)</td>
<td>44 (10–101)</td>
<td>26 (10–74)</td>
<td>44 (10–101)</td>
<td>27 (13–51)</td>
</tr>
<tr>
<td>C reactive protein (mg/l)</td>
<td>40 (7–290)</td>
<td>33 (29–64)</td>
<td>92 (7–290)</td>
<td>34 (7–61)</td>
</tr>
<tr>
<td>HLA-B27 (+/-)</td>
<td>14/5</td>
<td>3/0</td>
<td>6/1</td>
<td>4/3</td>
</tr>
</tbody>
</table>

AS, ankylosing spondylitis; AS + PA, ankylosing spondylitis with peripheral arthritis; PsA, psoriatic arthritis; uSpA, undifferentiated spondyloarthropathy.

Table 1 Demographic and descriptive characteristics of the study group. Values are given as median and range.

Efficacy evaluation

Global (n=19) and peripheral (n=16) assessments were performed at weeks 20, 22, 28, 34, 36, 42, 48, and 50, and were compared with the baseline value. Axial assessments (n=11) were carried out at weeks 20, 34, 48, and 50. Table 2 summarises the results (data for weeks 28 and 42 are not shown). At week 50 a significant decrease (p<0.05) for all variables was observed, comparable with the effect seen at week 12. When the global assessments were analysed at different times, this significant decrease was maintained for all variables, except for erythrocyte sedimentation rate (ESR) and CRP at week 48, immediately before the third re-treatment, reflecting a trend towards relapse in some patients. A similar trend was noted for morning stiffness of the spine, spinal pain assessed by the patient on aVAS, and the Dougadoas functional index.

Figure 1 shows the evolution over this one year period for patient global assessment (100 mm VAS), CRP (mg/l), swollen joint count, and spinal pain assessment (100 mm VAS).

When patients were asked about recurrence of symptoms, reflecting a possible disease relapse, 3/19 (16%) patients reported recurrence of symptoms at week 20, 13/19 (68%) reported a relapse at week 34, and 15/19 (79%) reported a relapse at week 48. However, two weeks after each re-treatment, at weeks 22, 36, and 50, none of the 19 patients reported such disease manifestations. The moment of recurrence of symptoms varied, though for most of these patients 3/3 at week 20, 12/13 at week 34, and 12/15 at week 48 symptoms occurred between weeks 10 and 14 after re-treatment. Nevertheless, none of them reported a full relapse comparable with baseline symptoms.

initial trial. Patients with AS were classified according to the modified New York criteria. Disease modifying drugs were not allowed during the study.

Concurrent drugs

During this extension study, efforts were made to keep the dose and schedule of non-steroidal anti-inflammatory drugs or prednisolone, or both, stable; however, modifications were allowed at the discretion of the treating rheumatologist.

Drug administrations

Infliximab at a dose of 5 mg/kg in 250 ml NaCl 0.9% was prepared by the hospital pharmacy. The total dose was given over a period of at least two hours. Aseptic procedures were used during the preparation and administration of the study drug. Patients were re-treated every 14 weeks (weeks 20, 34, and 48).

Clinical evaluation

Evaluation of efficacy was performed at weeks 20, 22, 28, 34, 36, 42, 48, and 50, and were compared with the baseline value. Axial assessments (n=11) were carried out at weeks 20, 34, 48, and 50. Table 2 summarises the results (data for weeks 28 and 42 are not shown). At week 50 a significant decrease (p<0.05) for all variables was observed, comparable with the effect seen at week 12. When the global assessments were analysed at different times, this significant decrease was maintained for all variables, except for erythrocyte sedimentation rate (ESR) and CRP at week 48, immediately before the third re-treatment, reflecting a trend towards relapse in some patients. A similar trend was noted for morning stiffness of the spine, spinal pain assessed by the patient on aVAS, and the Dougadoas functional index.

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Table 2

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>Week 12</th>
<th>Week 20</th>
<th>Week 22</th>
<th>Week 34</th>
<th>Week 36</th>
<th>Week 48</th>
<th>Week 50</th>
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<tr>
<td><strong>Global assessment</strong></td>
<td>n=21</td>
<td>n=21</td>
<td>n=19</td>
<td>n=19</td>
<td>n=19</td>
<td>n=18</td>
<td>n=19</td>
<td>n=19</td>
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<tr>
<td>Patient global (0–100)</td>
<td>57 (27–100)</td>
<td>10 (1–73)***</td>
<td>10 (0–94)***</td>
<td>6 (0–34)***</td>
<td>24 (1–97)***</td>
<td>5.5 (0–41)***</td>
<td>22 (2–95)*</td>
<td>14 (2–47)**</td>
</tr>
<tr>
<td>Patient pain (0–100)</td>
<td>70 (23–100)</td>
<td>10 (1–86)***</td>
<td>14 (1–96)***</td>
<td>8 (1–37)***</td>
<td>29 (1–100)***</td>
<td>7.5 (1–37)***</td>
<td>26 (2–100)*</td>
<td>15 (1–60)**</td>
</tr>
<tr>
<td>Doctor global (0–100)</td>
<td>57 (12–79)</td>
<td>10 (5–75)***</td>
<td>14 (1–38)***</td>
<td>14 (1–32)***</td>
<td>20 (1–60)**</td>
<td>10 (5–23)***</td>
<td>21 (7–73)*</td>
<td>12 (4–20)**</td>
</tr>
<tr>
<td><strong>Peripheral assessment</strong></td>
<td>n=18</td>
<td>n=18</td>
<td>n=16</td>
<td>n=16</td>
<td>n=16</td>
<td>n=15</td>
<td>n=16</td>
<td>n=16</td>
</tr>
<tr>
<td>Morning stiffness (min)</td>
<td>90 (10–300)</td>
<td>5 (0–60)***</td>
<td>10 (0–300)**</td>
<td>–</td>
<td>25 (0–120)*</td>
<td>–</td>
<td>25 (0–120)**</td>
<td>7.5 (0–90)**</td>
</tr>
<tr>
<td>Pain peripheral joints (0–100)</td>
<td>75 (20–99)</td>
<td>12 (0–84)***</td>
<td>15 (4–31)***</td>
<td>–</td>
<td>28 (0–80)***</td>
<td>–</td>
<td>34 (1–91)*</td>
<td>11.5 (1–34)**</td>
</tr>
<tr>
<td>Tender joint count (0–68)</td>
<td>6 (1–18)</td>
<td>0 (0–5)***</td>
<td>1 (0–6)***</td>
<td>0.5 (0–5)***</td>
<td>0.5 (0–8)***</td>
<td>0 (0–4)***</td>
<td>1 (0–12)*</td>
<td>0 (0–4)**</td>
</tr>
<tr>
<td>Night pain (0–3)</td>
<td>3 (1–3)</td>
<td>1 (0–1)**</td>
<td>1 (0–2)*</td>
<td>–</td>
<td>1 (0–3)*</td>
<td>–</td>
<td>1 (0–3)*</td>
<td>0 (0–1)**</td>
</tr>
<tr>
<td>Pain spine (0–100)</td>
<td>57 (4–95)</td>
<td>11 (1–60)**</td>
<td>15.5 (3–62)**</td>
<td>–</td>
<td>17 (0–81)*</td>
<td>–</td>
<td>22 (1–88) NS</td>
<td>10 (1–65)**</td>
</tr>
<tr>
<td>BASFI (0–100)</td>
<td>77.5 (9.9–94.2)</td>
<td>22.3 (1.0–63.2)**</td>
<td>36.8 (1–74.4)**</td>
<td>–</td>
<td>37.9 (4.7–73.7)**</td>
<td>–</td>
<td>45.6 (5.8–75.3)*</td>
<td>17.9 (3.9–63.9)**</td>
</tr>
<tr>
<td>Spinal pain (0–3)</td>
<td>2 (0–3)</td>
<td>0 (0–2)*</td>
<td>0 (0–3)*</td>
<td>–</td>
<td>0.5 (0–3) NS</td>
<td>–</td>
<td>0 (0–3) NS</td>
<td>0 (0–3)*</td>
</tr>
<tr>
<td>Occiput to wall (cm)</td>
<td>12 (3.7–21.5)</td>
<td>7.7 (2.5–22.0)**</td>
<td>9 (3.5–21)*</td>
<td>–</td>
<td>8.5 (3.8–21)*</td>
<td>–</td>
<td>8.5 (4–20)*</td>
<td>10 (2.5–20)*</td>
</tr>
<tr>
<td>Chest expansion (cm)</td>
<td>1.8 (1.5–3.2)</td>
<td>3.8 (1.7–5.5)**</td>
<td>3.5 (1.7–5.5)**</td>
<td>–</td>
<td>3.2 (1.5–5.6)**</td>
<td>–</td>
<td>4.5 (1.8–5.5)**</td>
<td>4.5 (2.0–6.0)**</td>
</tr>
<tr>
<td>Fingers to floor (cm)</td>
<td>24 (5.5–38.0)</td>
<td>15 (0–20.3)**</td>
<td>10.5 (0–33.8)**</td>
<td>–</td>
<td>15.6 (7.2–36.5)**</td>
<td>–</td>
<td>17.5 (0–31)*</td>
<td>17.4 (0–31)*</td>
</tr>
<tr>
<td>Dougados articular index (0–30)</td>
<td>6 (1–15)</td>
<td>0 (0–4)**</td>
<td>0.5 (0–4)**</td>
<td>–</td>
<td>2 (0–11)*</td>
<td>–</td>
<td>1.5 (0–8)**</td>
<td>0 (0–4)**</td>
</tr>
</tbody>
</table>

- *p < 0.05
- **p < 0.01
- ***p < 0.001
- NS, not significant (determined by Wilcoxon signed ranks test compared with baseline).

Safety evaluation and adverse drug reactions

No patients were withdrawn from the study owing to adverse events. Infliximab infusions were well tolerated. During the one year follow up one mild self limiting, infusion related adverse event was observed. Nineteen of the 21 patients had an adverse event. Minor events comprised nausea (n=3), dizziness (n=2), headache (n=2), fatigue (n=3), palpitations (n=1), and transient epigastric pain (n=3). One patient with PsA developed eczema of hands and feet. Another patient had an episode of erosive lichen of the mouth mucosa, which disappeared after local treatment. In total, 12 infectious episodes were noted: eight patients had an episode of self limiting upper respiratory tract infection, whereas in four patients the infections (one otitis media, one vaginal candidiasis, one tooth abscess, and one pyelonephritis) required antibiotic or antymycotic treatment. None of these infections were life threatening, nor did they require admission to hospital. No malignancies were reported. Laboratory tests performed at week 50 indicated a significant increase in serum aspartate aminotransferase (median score at baseline 17 U/l (range 12–36), median score at week 50, 28 U/l (range 12–55), p<0.001) and serum alanine aminotransferase (median score at baseline 21 U/l (range 7–55), median score at week 50, 35 U/l (range 12–83), p<0.001), but these median values were still in the normal range. Furthermore, these laboratory findings were not clinically significant. A significant increase in haemoglobin was found at week 50 (median score at baseline 132 g/l (range 101–159), median score at week 50, 148 g/l (range 133–161), p<0.001) compared with baseline. Initially, nine (42%) patients had a normocytic normochromic anaemia; however, after one year’s maintenance treatment the haemoglobin level and packed cell volume had normalised in all patients (53). During the one year follow up 12/21 (57%) patients developed ANA; in four of these patients (19%) anti-dsDNA were detected. However, no abnormalities in peripheral blood count or complement nor any symptoms suggestive for lupus-like syndromes occurred.

**DISCUSSION**

In this study we report the safety and efficacy of a maintenance regimen of 5 mg/kg infliximab every 14 weeks in patients with active 5pA. The patients included in this extension protocol were initially treated with three infusions of infliximab in a 12 week open label pilot study after this loading dose regime, a rapid and significant improvement in global, peripheral, and axial disease manifestations was seen, without major side effects.
From this one year follow up study the following conclusions can be reached. Firstly, a statistically significant decrease of global, peripheral, and axial disease manifestations was seen after one year of treatment in the 19 evaluated patients. Two patients changed to another dosing regimen owing to partial lack of efficacy of the drug. Improvements in BASMI, chest expansion, fingers to floor, and occiput to wall distance in the group of patients with AS, paralleled global improvement of the disease activity in these patients. It is unclear if this improvement in spinal metrology is an indication of an effect on the progressive axial ankylosis which is typical for AS. This study was designed to evaluate the clinical efficacy of infliximab in SpA, and not to evaluate the effect on ankylosis in AS: therefore one cannot draw conclusions from this small subgroup. In order to detect such a structural improvement, homogeneous patient groups, adequate sample size, and sensitive outcome measurements, including radiological assessments evaluated over longer periods, are mandatory.

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Secondly, recurrence of symptoms, such as axial night pain, morning stiffness, or synovitis, was reported by three patients (16%) at week 20, 13 patients (68%) at week 34, and 15 patients (79%) at week 48, with the moment of recurrence between 10 and 14 weeks after re-treatment. This relapse was furthermore documented by the fact that ESR, CRP, axial morning stiffness, and axial pain assessed by the patient, were no longer statistically significant when compared with baseline, immediately before the third re-treatment, at week 48. Together with observations made in other open label studies, these results indicate that a maintenance regimen of 5 mg/kg infliximab every 14 weeks could not control the inflammatory disease activity continuously. Adjustment of the maintenance regimen seems warranted; however, it is not clear whether this should be achieved by increasing the dose or shortening the interval between doses.
Thirdly, re-treatment of patients whose symptoms recurred induced an improvement comparable with the response seen after the induction scheme reported in the 12 week pilot study. All variables of disease activity were again significantly improved two weeks after every re-treatment.

Fourthly, the treatment with infliximab in this patient group was shown to be safe and the infusion procedure was generally well tolerated. Only one mild self limiting infusion reaction was seen, and no major infectious episodes or malignancies occurred. Previous studies indicate that treatment with TNFα inhibitors, such as infliximab, can induce the development of ANA in patients with RA.39 So far no data on the occurrence of such antibodies in patients with SpA treated with infliximab have been reported: here, up to 12/21 (57%) patients developed ANA; in four of them (19%) anti-dsDNA antibodies were also detected. However, no significant laboratory changes in peripheral blood count or complement occurred in these patients; no patients developed proteinuria, nor where there any other lupus-like symptoms. The overall frequency of autoantibodies is comparable with the data reported in RA.44 However, we have to remember that the background for developing such antibodies—in RA versus SpA—is different. Occurrence of ANA in RA is not considered unusual and a prevalence up to 60% has been seen; in SpA and AS, however, only scarce data are available, showing a prevalence ranging from 11.4% to 19.1%.45 In these studies a bias attributed to the use of sulfasalazine is likely. A retrospective analysis of a patient group with SpA disclosed a prevalence of 6.8%.46 ANA positive patients before sulfasalazine treatment with no anti-dsDNA antibodies detected. After two to five years of sulfasalazine treatment, the incidence rose to 24% and 4%, respectively for ANA and anti-dsDNA antibodies.

In conclusion, this one year open pilot study suggests that infliximab is a safe and effective drug in the treatment of SpA. Regardless of the fact that these data need further confirmation in ongoing double blind, placebo controlled trials, the extent of the clinical improvements, their consistency throughout the study group, the parallel changes in laboratory indices of inflammation, and the maintenance of this therapeutic effect over one year, are encouraging. Further accurate follow up of this patient group is mandatory for the optimisation of the maintenance regimen and for the evaluation of longer term safety with regard to infections, malignancies, and the occurrence of ANA.

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REFERENCES


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