Management of ankylosing spondylitis with infliximab

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Abstract: Ankylosing spondylitis (AS) is a systemic inflammatory rheumatic disease responsible for back pain, stiffness and progressive loss of functional capacity with limited therapeutic options. Regular physical exercises together with the use of nonsteroidal antiinflammatory drugs are the two recognized treatment options in AS. Infliximab is a chimeric anti-tumor necrosis factor-α monoclonal antibody that has been demonstrated to be highly effective in the treatment of AS, providing clinical amelioration at both axial and peripheral skeleton. Infliximab also improves quality of life, function, biological parameters (acute phase reactants) and inflammatory lesions of the spine as detected by magnetic resonance imaging. It is given at a 5 mg/kg dosage, as an infusion at weeks 0, 2, 6, and every 6 to 8 weeks after. Open-label and placebo-controlled trials have well demonstrated its high level of efficacy, with an improvement of the disease activity of at least 50% in 60%–80% of patients. In a large placebo-controlled trial, Assessment in Ankylosing Spondylitis Response Criteria (ASAS20) responders were observed in 61.2% of patients receiving infliximab compared to 19.2% of patients under placebo. Long-term efficacy is maintained when infliximab is administered every 6–8 weeks. Consensus international guidelines for the initiation and the use of this expensive treatment are available. Some questions remain, notably whether this treatment has disease-controlling properties.

Keywords: anti-TNFα, infliximab, ankylosing spondylitis

Introduction
For the past ten years, the therapeutic management of ankylosing spondylitis (AS) has considerably changed. In fact, the introduction of anti-TNFα agents in AS provides new (and until now, unmet) therapeutic perspectives for the patients. This review analyzes the available data on clinical, radiological, and biological efficacy of infliximab, the first anti-tumor necrosis factor-α (anti-TNFα) used in AS, and discusses its place in the treatment of AS. Some questions persist, notably whether this treatment has disease-controlling properties.

Ankylosing spondylitis: The disease
Clinical features
Ankylosing spondylitis is a systemic and chronic inflammatory rheumatic disease of the axial skeleton (spine and sacroiliac joints). The disease mainly affects young male subjects, with onset of symptoms between the ages of 20 and 30 years. The rheumatological symptoms include inflammatory back pain, stiffness and restriction of spinal
Epidemiology and impact of the disease

The prevalence of the disease has been estimated to range between 0.1% and 1.1%. A German study evaluating the prevalence of AS in German HLA-B27 blood donors estimated it at 1.9% of the Berlin population. An epidemiological study in France (EPIRHub study) evaluated the rate of standardized prevalence of SpA to 0.30% and AS to 0.08%.  

Ankylosing spondylitis is a chronic progressive disease leading to a limited range of motion of the spine, a loss of functional capacity, a clinical feature increasing with disease duration. Some indicators of prognosis have been identified: the presence of extraarticular disease (uveitis), a hip involvement, the stage of the disease at the time of diagnosis, the degree of patient compliance, the level of socioeconomic status, and education. Thus, the disease has socioeconomic consequences, with a higher rate of days of sick leave, loss of work, and higher mortality compared to age- and sex-matched general population.

Assessment of the disease

Ankylosing spondylitis is a chronic inflammatory disease and like other chronic rheumatic diseases, various domains can be evaluated and especially, pain, inflammation, function, mobility, fatigue, global assessment, quality of life, concomitant treatments, radiographic changes, and others. According to international experts in AS assessment (Assessment in Ankylosing Spondylitis [ASAS] working group), several domains have been selected and are considered as clinically important in assessing symptomatic outcome in AS: physical function, pain, spinal mobility, patient global assessment, spinal stiffness, fatigue, and inflammation. For each of these domains, there is a number of instruments which could be used (Table 1). Each of these instruments have the required properties ie, reliability, validity, reproductibility, and sensibility to changes and are commonly used in clinical trials for evaluating symptomatic or disease-modifying treatments in AS.

The ASAS working group has defined criteria for evaluating symptomatic improvement in AS, using four of the five outcome domains ie, physical function, pain, patient global assessment and inflammation, and consists of an improvement by 20% and by 10 units (on a scale of 0–100 mm) in each of the three domains with no worsening of the fourth. This defines the ASAS20 criteria of short term improvement in AS which could be compared to the American College of Rheumatology (ACR) 20 criteria of improvement used in rheumatoid arthritis (RA).
Traditional therapeutic approach

Regular physical exercise associated with the administration of nonsteroidal anti-inflammatory drugs (NSAIDs) is considered to be the cornerstone of therapeutic management in AS.18–20 Physical exercise may slow down the progression of functional disability and NSAIDs are considered as the gold standard in AS drug therapy.21,22 NSAIDs clearly demonstrated substantial relief of symptoms, pain and stiffness in patient with axial or peripheral disease. However, the main problem of NSAIDs is their poor gastrointestinal tolerability. Another problem is that NSAIDs are not enough effective in some cases. In addition, there are limited alternative therapeutic options. Indeed, second-line treatments such as sulfasalazine and methotrexate are poorly effective in AS, particularly in patients with axial disease or enthesitis.23,24 Leflunomide or anakinra do not work on axial or peripheral disease. However, Pamidronate has been proposed in some NSAID refractory patients but studies gave contradictory results.27,28 Systemic corticosteroids are not effective in AS while local corticosteroid injections (in the sacroiliac joints or enthesal structures) may be helpful in some cases.29

Thus, before the introduction of TNFα-blocking agents, medical treatment options in AS were limited and there was an unmet need for new therapeutic options. Anti-TNFα agents have revolutionized this therapeutic management.

Infliximab

Rationale for targeting TNFα in AS

TNFα is a pro-inflammatory cytokine produced by monocytes/macrophages and activated T cells. It is responsible for lymphocyte activation, release of other cytokines (IL-1, IL-6), prostaglandins, and metalloproteinases. There are substantial data suggesting a role for TNFα in AS physiopathology. Indeed, TNFα RNA messenger has been found to be abundantly expressed in the sacroiliac joints from patients with AS3 and the serum levels of TNFα and other inflammatory cytokines have been found to be increased compared to patients with mechanical back pain.30,31 In addition, patients with AS (and also SpA) have sub-clinical gut involvement with histological lesions resembling those in Crohn’s disease.32 Anti-TNFα therapy was effective in Crohn’s disease patients.33 Another argument for targeting TNFα in AS and SpA is supported by the results of clinical trial with thalidomide in AS, a drug interfering with the production of TNFα.34 Taken together, these data give the rationale for the use of TNFα-targeting treatment in AS.35–38

Structure and pharmacokinetic properties

Infliximab (Remicade®; Centocor Ortho Biotech Inc., Horsham, PA) is a chimeric monoclonal antibody that is 75% human and 25% mouse and consists of the constant region of human IgG1κ coupled to the Fv region of high-affinity neutralizing murine anti-human TNFα antibody. Its molecular size is 149 kDa. It binds to soluble TNFα monomers and trimers and membrane-bound TNFα with high affinity, avidity, and specificity. Infliximab may promote both lysis of TNFα-producing cells and T lymphocyte apoptosis. Infliximab also inhibits binding of soluble TNFα to TNF receptors and may dissociate TNFα/TNF receptor complexes. However, infliximab does not bind to lymphotxin β. Infliximab is administered IV over at least two hours. The molecule has a half life of 8–12 days, although it has been detected in the serum 28 days after the infusion.39,40

Clinical efficacy of infliximab in ankylosing spondylitis

Infliximab has been tested in AS and/or SpA in 10 studies, including seven open31–38 and three placebo-controlled

Table 1 Specific instruments for each relevant domain in ankylosing spondylitis16

<table>
<thead>
<tr>
<th>Domain</th>
<th>Instrument</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>BASFI or Dougados Functional Index</td>
</tr>
<tr>
<td>Pain</td>
<td>VAS, last week, spine at night and VAS, last week, spine, due to AS</td>
</tr>
<tr>
<td>Spinal mobility</td>
<td>Chest expansion and modified Schober and occupit to wall distance</td>
</tr>
<tr>
<td>Patient global assessment</td>
<td>VAS, last week</td>
</tr>
<tr>
<td>Stiffness</td>
<td>Duration of morning stiffness, spine, last week</td>
</tr>
<tr>
<td>Peripheral joints and entheses</td>
<td>Number of swollen joints (44 joint count); no preferred instrument for entheses</td>
</tr>
<tr>
<td>Acute phase reactants</td>
<td>ESR</td>
</tr>
<tr>
<td>Radiograph of the spine</td>
<td>Anteroposterior + lateral spine (lumbar + cervical) + X-ray pelvis (sacroiliac, hips)</td>
</tr>
<tr>
<td>Fatigue</td>
<td>No preferred instrument</td>
</tr>
</tbody>
</table>

**Abbreviations:** AS, ankylosing spondylitis; VAS, visual analog scale; BASFI, Bath Ankylosing Spondylitis Functional index; ESR, erythrocyte sedimentation rate.
trials\textsuperscript{48-50} (Tables 2 and 3). These different trials enrolled between 11 and 279 patients. Patients received three infusions of infliximab 5 mg/kg at weeks 0, 2, and 6 (except in one Canadian study which administered infliximab 3 mg/kg,\textsuperscript{44} followed by a maintenance regimen every 6, 8, or 14 weeks in the extension or follow-up studies.\textsuperscript{51,52} The study duration ranged from 12 weeks to one year. All these studies assessed the effects of infliximab on clinical parameters (spinal pain, swollen and tender joint scores, enthesal count, Bath Ankylosing Spondylitis Disease Activity Index [BASDAI] and Bath Ankylosing Spondylitis Functional Index [BASFI], quality of life questionnaire) and laboratory tests (erythrocyte sedimentation rate [ESR], C-reactive protein [CRP]). In some studies, the primary outcome measures were the 50% improvement in the BASDAI index or the ASAS20 response for improvement. Imaging methods using dynamic MRI were used in two studies to evaluate the impact of the treatment on enthesal lesions, synovitis, and sacroiliitis.\textsuperscript{42,43} Open-label studies were conducted in different countries (Germany with 11 AS patients, Belgium with 21 SpA patients, France with 50 AS patients, Spain with 40 SpA patients, Canada with 21 AS patients, and Greece with 25 AS patients) (Table 2). Altogether, open-label studies showed that infliximab gave a dramatic improvement in all clinical and laboratory variables of AS patients. An improvement of at least 50% was achieved in up to 60%–80% of patients. The beneficial effects were observed early after the beginning of the treatment, within two weeks after the first infusion of infliximab. Both axial and peripheral symptoms but also enthesopathic manifestations improved. In the German study, symptoms recurred around six weeks after the three infliximab infusions.\textsuperscript{52} This led the authors to propose a six-week interval between infusions in the subsequent randomized controlled trial.

Table 2

<table>
<thead>
<tr>
<th>Author (Country) (Reference)</th>
<th>Patients (N)</th>
<th>Mean age</th>
<th>Treatment protocol</th>
<th>Outcome parameters</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brandt (Germany)\textsuperscript{41}</td>
<td>11 AS</td>
<td>36</td>
<td>IFX 5 mg/kg Week 0, 2, 6 Follow up: 12 weeks</td>
<td>Spinal pain (VAS) BASDAI, BASFI, ESR, CRP, IL-6, SF-36, MRI</td>
<td>BASDAI improvement by 70% at week 4 and improvement in all end-points</td>
</tr>
<tr>
<td>Van den Bosch (Belgium)\textsuperscript{42}</td>
<td>21 SpA (10 AS)</td>
<td>49</td>
<td>IFX 5 mg/kg Week 0, 2, 6 Follow up: 12 weeks</td>
<td>Patient and physician global assessment of disease activity, BASDAI, BASFI, ESR, CRP</td>
<td>Improvement in all end-points</td>
</tr>
<tr>
<td>Stone (Canada)\textsuperscript{42}</td>
<td>21 AS</td>
<td>37.9 8.7</td>
<td>IFX 5 mg/kg Week 0, 2, 6 Follow up: 14 weeks</td>
<td>Spinal pain, patient and physician global assessment of disease activity, BASDAI, BASFI, ESR, CRP</td>
<td>More than 60% improvement in BASDAI, BASFI, HAQ, pain, fatigue, acute phase reactants</td>
</tr>
<tr>
<td>Maksymowych (Canada)\textsuperscript{44}</td>
<td>21 AS</td>
<td>42.5 13.8</td>
<td>IFX 3 mg/kg Week 0, 2, 6 and the every 8 weeks Follow up: 14 weeks</td>
<td>BASDAI, BASFI, BAS-G, BASMI, ESR, CRP, MRI</td>
<td>More than 50% improvement in BASDAI</td>
</tr>
<tr>
<td>Breban (France)\textsuperscript{45}</td>
<td>50 AS</td>
<td>Median 35 Median 13</td>
<td>IFX 5 mg/kg Week 0, 2 Follow up: 24 weeks</td>
<td>Fatigue, BASDAI, BASFI, global pain, CRP, ASAS20</td>
<td>94% ASAS20 responders</td>
</tr>
<tr>
<td>Temekonidis (Greece)\textsuperscript{46}</td>
<td>25 AS</td>
<td>36</td>
<td>IFX 5 mg/kg Week 0, 2, 6 and every 8 weeks for 12 months</td>
<td>BASDAI, BASFI; patient global assessment of pain, ESR, CRP</td>
<td>Significant improvement in BASDAI, CRP. Reduction of patient global pain by 20% in 92%</td>
</tr>
<tr>
<td>Collantes-Estevez (Spain)\textsuperscript{47}</td>
<td>40 SpA (34 AS)</td>
<td>41</td>
<td>IFX 5 mg/kg Week 0, 2, 6 and every 8 weeks for 38 weeks</td>
<td>BASDAI, BASFI, patient pain assessment, quality of life, 50% improvement in BASDAI</td>
<td>50% improvement of BASDAI in 60% of patients</td>
</tr>
</tbody>
</table>

Abbreviations: IFX, infliximab; BASDAI, Bath Ankylosing Spondylitis Disease Activity Index; BASFI, Bath Ankylosing Spondylitis Functional Index; BASMI, Bath Ankylosing Spondylitis Metrology Index; VAS, visual analog scale; HAQ, health assessment questionnaire; ESR, erythrocyte sedimentation rate; MRI, magnetic resonance imaging; ASAS, Assessment in Ankylosing Spondylitis group.
• The group from Ghent published the results of a placebo controlled trial of infliximab treatment in a series of 40 active SpA patients (including 19 AS patients).\textsuperscript{40} Twenty patients received infliximab 5 mg/kg at weeks 0, 2, 6, and every 8 weeks thereafter and 20 patients the placebo at the same interval. At week 12, there was a significant improvement of disease activity, physician global, patient global, peripheral arthritis and acute phase reactants in the infliximab group compared to the placebo group. Response to infliximab was seen already at week 2. One patient under infliximab developed disseminated tuberculosis.

• Similar results were obtained in a 12-week placebo-controlled study from the German group.\textsuperscript{48} Seventy patients participated in this trial, 35 receiving infliximab at weeks 0, 2, and 6, and 35 received the placebo. The primary outcome was a 50% improvement of the BASDAI which was achieved at week 12 in 53% of the patients treated by infliximab and only 8% of the patients on placebo. The other results showed that quality of life (assessed by short from SF-36), physical function (assessed by BASFI), spinal mobility (assessed by Bath Ankylosing Spondylitis Metrology Index [BASMI]) significantly improved in the infliximab group. Peripheral arthritis and enthesitis also improved. The concentration of CRP in serum was identified as a probable indicator of response to infliximab since patients with high concentrations of CRP had the best rate of 50% improvement in disease activity.

• The largest placebo-controlled trial with infliximab enrolled 279 patients (201 received infliximab 5 mg/kg and 78 received the placebo).\textsuperscript{50} Treatment was given at weeks 0, 2, 6, and every 6 weeks over 24 weeks. The primary outcome parameter in this study was the proportion of patients with a 20% improvement response according to the ASAS criteria. At week 24, 61.2% of patients in the infliximab group were ASAS20 responders compared to 19.2% in the placebo group. Other parameters (BASDAI, BASFI, BASMI, SF-36, number of swollen joints and acute phase reactants) improved similarly in the infliximab group. Similarly to other placebo-controlled studies, patients receiving infliximab showed significant improvement as early as week 2 and this response was maintained over the 24 week study.

### Efficacy of infliximab assessed by imaging

The good clinical responses observed in the patients receiving infliximab paralleled the imaging amelioration as evaluated by magnetic resonance imaging (MRI).\textsuperscript{43,44} The modifications in MRI changes were observed at the different pathogenic levels of AS, ie, sacroiliac and peripheral joints and enthesal structures. In the German-controlled trial, MRI with STIR

### Table 3 Randomized placebo-controlled trials evaluating the efficacy of infliximab in ankylosing spondylitis

<table>
<thead>
<tr>
<th>Authors (Country) Reference</th>
<th>Patients (N)</th>
<th>Mean age Mean disease duration</th>
<th>Treatment protocol</th>
<th>Outcome parameters</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Braun (Germany)\textsuperscript{48}</td>
<td>70 AS 39.8 15.6</td>
<td>IFX 5 mg/kg or placebo Week 0, 2, 6 follow up 12 weeks</td>
<td>BASDAI, BASFI, BASMI, quality of life</td>
<td>50% improvement in BASDAI: 53% (IFX) vs 8% (placebo) Significant improvement in all end points in the IFX group</td>
<td></td>
</tr>
<tr>
<td>Van den Bosch (Belgium)\textsuperscript{48}</td>
<td>40 SpA 46.5 7.2</td>
<td>IFX 5 mg/kg or placebo Week 0, 2, 6 follow up 12 weeks</td>
<td>Patient and physician global assessment of disease activity; ESR, CRP, tender and swollen joint counts, morning stiffness, spinal pain, BASDAI, BASFI</td>
<td>Significant improvement in all end points in the IFX group</td>
<td></td>
</tr>
<tr>
<td>Van der Heijde (international)\textsuperscript{50}</td>
<td>279 AS 41 11 IFX 13.2 Placebo</td>
<td>IFX 5 mg/kg or placebo Week 0, 2, 6, 12 and 18 Follow up: 24 weeks</td>
<td>ASAS20 responders BASDAI, pain, BASFI, BASMI, enthesis index, swollen joint, CRP, SF-36</td>
<td>ASAS20 responders: 61.2% IFX vs 19.2% placebo Significant improvement in all assessed variables in the IFX group</td>
<td></td>
</tr>
</tbody>
</table>

**Abbreviations:** IFX, infliximab; BASDAI, Bath Ankylosing Spondylitis Disease Activity Index; BASFI, Bath Ankylosing Spondylitis Functional Index; BASMI, Bath Ankylosing Spondylitis Metrology Index; vAS, visual analog scale; ESR, erythrocyte sedimentation rate; ASAS, Assessment in Ankylosing Spondylitis group.
sequences showed a significant regression by 60% of active spinal inflammatory lesions in the infliximab patients compared with 20% deterioration in the placebo group.53

Another imaging method to evaluate the efficacy of TNFα blockers in AS is ultrasonography: in two patients with refractory heel pain related to enthesopathic lesions at the insertion of the Achilles tendon and plantar fascia, infliximab dramatically improved the symptoms. Inflammatory lesions were shown using power Doppler sonography and after infliximab administration, a progressive recovery of normal entheseal structures was observed in both patients.54

Ankylosing spondylitis is characterized by osteoporosis.55 It was demonstrated that bone mineral density (BMD) (assessed by dual energy X-ray absorptiometry [DEXA]) improved after six months of infliximab treatment in a series of 29 SpA patients: lumbar spine BMD increased by 3.6% and hip BMD by 2.2%.56

Effects of infliximab on laboratory and immunological parameters
In both open-label and placebo-controlled trials, it was evident that laboratory parameters of inflammation and ESR and CRP levels improved with a rapid decrease after starting the treatment.48–50

In addition, changes on cytokine profile were examined during infliximab therapy by means of circulating cytokine levels or intracellular cytokine protein expression in peripheral blood T cells.57 The circulating levels of TNFα or other inflammatory cytokine such as IL-6 were found to be increased in AS or SpA in general, as well as serum IL-10, an antiinflammatory cytokine.30,31 The changes in cytokine T cell expression during infliximab therapy was analyzed by Zou and colleagues. In this study, 10 patients with AS received infliximab at weeks 0, 2, and 6. The percentage of CD4+ CD8+ T cells positive for intracellular TNFα and interferon-γ (IFNγ) was evaluated at baseline and at weeks 6 and 12. In these patients, infliximab induced a significant reduction in the number of CD4+ CD8+ T cells expressing TNFα and IFNγ, while there were no changes for IL-10 or IL-4 production.58 In addition, the same author analyzed the changes on cytokine T cell expression after 12 weeks of etanercept (another TNFα-blocking agent) administration. By contrast, the results did not show a downregulation of TNFα or IFNγ but an increase in the percentage of CD4+ CD8+ T cells positive for these cytokines.59 The author explained their results by the fact that infliximab can downregulate Th1 cytokine while etanercept binds soluble TNFα and upregulates Th1 protein production after neutralization of peripheral TNFα. Different results were obtained in another study on the T cell cytokines in peripheral blood of SpA patients.50

Before infliximab treatment, patients with SpA were found to have an impaired Th1 cytokine profile compared with healthy controls and RA patients, as suggested by a decreased percentage of mononuclear cells positive for intracellular IFNγ and IL-2. TNFα blockade induced restoration of Th1 cytokines with a significant and persistent increase in IFNγ and IL-2. These data suggest that infliximab modifies the cytokine profile in SpA patients and can reverse a state of energy of Th1 cells. Overall, these descriptions of cytokine protein expression in peripheral blood mononuclear cells during anti-TNFα therapy are not clear. This may reflect the complexity of cytokine regulation and may also be related to the artificial nature of stimulating cytokine expression before measurement.

Serum matrix metalloproteinase 3 (MMP-3) and macrophage colony-stimulating factor-1 (MCSF-1) were identified as laboratory parameters of disease activity in AS.61 In addition, serum MMP-3 is also an independent predictor of radiological progression in AS.62 When patients with AS were treated by infliximab, serum MMP-3 decreased while there were no changes in the serum levels of MCSF-1.63 Another study found similar results with a decreased in serum MMP-3 in a series of 12 patients receiving infliximab.61

Histological changes induced by infliximab administration in AS
In the Ghent cohort was particularly examined the impact of infliximab on synovial tissue obtained by needle arthroscopy from inflamed peripheral joints.64 In eight patients (3 AS, 1 undifferentiated SpA, 4 psoriatic arthritis) receiving infliximab was performed a series of knee synovial biopsies. Consecutive histological analysis of the synovial tissues showed that, after the treatment, the synovial lining thickness decreased with a reduction of the inflammatory infiltrate (neutrophils, macrophages, and also CD4+ T cells) and CD55+ fibroblast-like synoviocytes. A reduction of synovial vascularization and a decreased of endothelial expression of certain but not all adhesion molecules (vascular cell adhesion molecule-1 but not intracellular adhesion molecule-1 or E-selectin) were also observed. Of interest, the overall degree of cellular infiltrate did not change since neutrophils and CD68+ macrophages decreased while CD20+ lymphocytes and plasma cells increased. Another study found that the synovial expression of metalloproteinase MMP-3 and tissue inhibitor of matrix metalloproteases (TIMP-1) significantly decreased after infliximab treatment.63
Long term efficacy of infliximab in AS

The initial published studies reported the effects of infliximab on AS or SpA symptoms over a short period of 12 weeks. One study in Greece was conducted over a longer period (12 months).46

Data about long term treatment with infliximab in AS and SpA are now available: the Ghent open-label trial investigating the therapeutic potential of infliximab in SpA was extended to one year and the patients of the initial open trial were retreated every 14 weeks during one year.51 This study showed that the significant improvements of disease manifestations were all maintained without major side effect or loss of efficacy. The moment of recurrence of SpA symptoms was observed between 10 and 14 weeks after the last infusion, indicating that infliximab has a transient efficacy and therefore, the optimal maintenance regimen was certainly below this interval of time.

Prolonged infliximab efficacy has also been demonstrated in patients from the German study who went on to receive infliximab at a six-week interval for up to five years.52,65–67 After one year of therapy, 78% of patients were still being treated and around 50% of them still achieved 50% improvement in BASDAI score.52 After 2, 3, and 5 years, 70%, 62%, and 55% continued treatment, respectively.55–67 These studies confirm that long-term treatment with infliximab remains very efficacious in patients with active AS, with persistence of a low disease activity and durable clinical response in the different assessed variables (function, mobility, peripheral arthritis, enthesitis and quality of life). At two years, ASAS20 response was observed in 73% of patients.68

In another study from Greece, persistent clinical response was also observed in 35 patients with AS over a four-year period, with an infliximab survival rate of 77.9%.69 Another cohort from Greece reported the same adherence to treatment after one (94%) and two (89%) years.70

In France, continuation of infliximab over time was also estimated in an open, observational two-year extension study after an open-label study of three infusions of treatment in refractory AS. Patients received infliximab infusions at a variable interval (mean: 11.6 ± 9 weeks), according to the rheumatologist opinion. At year 2, continuation rate was 74%.71 Thus, sustained clinical efficacy and high continuation rate over time were observed with prolonged infliximab treatment.

Safety and tolerance

Adverse events associated with infliximab treatment have been recently reviewed.72 The main safety concerns of infliximab are infusion reactions, infections, especially opportunistic infections including tuberculosis, development of auto-antibodies and auto-immune diseases, risk of lymphoma, and more rarely occurrence of demyelinating diseases and congestive heart failure.50,72 Placebo-controlled studies and long-term extensions of these studies, and post-marketing surveillance give important information on the adverse events of infliximab and other TNFα antagonists.

During the clinical trials with infliximab in AS, mild side effects were recorded including headache, dizziness, paresthesia, fatigue, upper respiratory infections (with a frequency which did not differ between infliximab and placebo), other minor infections (cystitis, sore throat, rhinitis, sinusitis, pharyngitis), abdominal pain, development of anti-nuclear antibodies in single patients. In general, there were no infusion reactions or delayed type hypersensitivity with infliximab infusions. In the controlled study by Braun, three serious adverse events were observed leading to withdrawal from the study: one case of systemic tuberculosis, one case of bronchocentric allergic granulomatosis, and one case of neutropenia.48 Maksymowycz reported two serious adverse events: one septic osteomyelitis, and one case of severe hypersensitivity reaction.44 A Belgian controlled study observed one case of tuberculosis and one septic arthritis.49 During the one-year follow-up study of this cohort from Ghent, 57% of SpA patients developed antinuclear antibodies while anti-double stranded DNA occurred in 19% of these patients.51 The German cohort evaluated the frequency of antinuclear antibodies to 25%.52 In the ASSERT study, adverse events were reported by 82.2% of patients receiving infliximab and 72% receiving placebo. Forty-three percent of patients in the infliximab group reported infection compared with 36% in the placebo group.70

In all of these studies testing the efficacy of infliximab, there were no life-threatening events and no cases of malignancy. Considering the whole group of AS and SpA patients included in these clinical trials (total = 500), serious infections such as tuberculosis occurred in only two cases.

In the systematic safety follow-up of the Ghent cohort including 107 SpA patients treated with infliximab, corresponding to a total of 191.5 patient years, eight severe infections were observed, including two tuberculosis, three retropharyngeal abscesses, one extensive wound infection, one septic arthritis, and one unprecised sepsis.73

Risk of lymphoma is a serious preoccupation in patients receiving anti-TNFα agents. In rheumatoid arthritis, there is an increased risk for lymphoma development which is related to the disease activity. Whether this risk is increased...
by anti-TNFα treatment or not remains still debated. The risk for malignant lymphoma in AS has been evaluated in a population-based case control study conducted in Sweden. In this study and in the absence of anti-TNFα agents, patients hospitalized with AS do not show increased risk for lymphoma (relative risk [RR]: 1.0; 95% CI: 0.6–1.7). In the different clinical trials with infliximab in AS and the long-term surveillance of these patients, a limited number of lymphoma were reported (four cases by the French three-year prospective RATIO observatory). In addition, one exceptional case of lymphoma has been reported in a patient with AS treated by etanercept.

**Impact of infliximab on extraarticular manifestations of ankylosing spondylitis**

Acute anterior uveitis occurs in 20%–30% of AS patients. Infliximab was shown to be effective in the treatment of acute onset HLA-B27 associated anterior uveitis and resistant SpA-related uveitis. During the clinical trials with infliximab in AS patients, a small proportion of patients developed acute uveitis under infliximab compared with patients in the placebo group. Using a registry-based study, the cases of uveitis occurring associated with the three available TNFα-blocking agents (infliximab, etanercept, and adalimumab) were reviewed. After correction for the estimated number of cases treated by each treatment, the results showed that etanercept was associated with a greater number of uveitis than infliximab or adalimumab. Another study estimated the incidence of anterior uveitis in AS patients receiving infliximab or etanercept. Data were collected from placebo-controlled and open-label studies. The frequency of anterior uveitis was higher in patients receiving placebo compared to patients under anti-TNFα agents (15.6 versus 6.8 per 100 patient-years). Of interest, flares of anterior uveitis occurred less frequently in patients treated with infliximab (3.4/100 patient-years) than in those treated with etanercept (7.9/100 patient-year), but the difference was not significant. Collectively, these data suggest that infliximab (and other TNFα blockers) is associated with a decrease incidence of anterior uveitis, a phenomenon that was more evident with the monoclonal antibodies.

Psoriasis may be associated with AS. A randomized placebo-controlled trial demonstrated that infliximab was effective in psoriatic arthritis and in the treatment of skin lesions of psoriasis. Infliximab, on the contrary of etanercept is effective in Crohn’s disease. Patients with AS have subclinical gut involvement resembling Crohn’s disease at a high frequency (40%–60%). In this sense, infliximab is a treatment of choice for AS patient with concomitant Crohn’s disease. A retrospective analysis of open-label and placebo-controlled trials evaluated the incidence of flares or new onset of IBD in patients with AS treated by anti-TNFα. Data were analyzed from 419 AS patients exposed to infliximab (618 patient-years) or etanercept (625 patient-years) or adalimumab (132 patient-years) and 434 placebo patients (150 patient-years). The results showed that new onset and flare of IBD were infrequent in patients receiving anti-TNFα agents. In addition, infliximab prevented better flare of IBD than etanercept. However, the incidence of new onset of IBD was statistically not different from placebo for all anti-TNFα agents.

**Current guidelines for initiating infliximab in ankylosing spondylitis**

Infliximab has been approved in the USA (in 2004) for reducing signs and symptoms in patients with active AS and in Europe (in 2003) for treatment in patients with severe axial AS and elevated laboratory parameters of inflammation and who are refractory to conventional treatments. Infliximab is also licensed for the treatment of Crohn’s disease, rheumatoid arthritis, psoriatic arthritis, and psoriasis without arthritis. A relevant question is which patient can benefit from infliximab, a high-cost treatment?

In 2002, the ASAS working group proposed a consensus statement for the use of TNFα blockers in AS patients. These recommendations were updated in 2006 and involved different levels concerning diagnosis, failure of standard treatments, disease activity, monitoring and criteria for withdrawal. Patients should respond to the modified New York criteria. Standard treatment failure corresponds to the trial of at least three months of standard NSAIDs, with at least two different NSAIDs. Failure of intraarticular joint corticosteroid injection and sulfasalazine for four months in the case of peripheral AS are required. For enthesopathic symptoms, failure of steroid injections is also required. Disease activity is defined based on the BASDAI, with a score higher than 4/10. The disease must be active for at least four weeks and the expert opinion should be useful. For monitoring the patients, the following clinical variables must be recorded: patient assessment, pain, spinal mobility, BASDAI, BASFI, swollen joint count, and laboratory parameters of inflammation (ESR and CRP levels). There was an agreement to state that biologic agents should be discontinued in patients who do not have a 50%
or a two-point absolute improvement in the BASDAI after 6–12 weeks. This consensus statement may be used as guidelines for clinical decision in initiating (or interrupting) anti-TNFα therapy in AS.

The Canadian rheumatology association has also proposed a consensus for infliximab (or etanercept) initiation, based upon literature analysis. Anti-TNFα agents are indicated for reducing signs and symptoms of AS with moderate to severe activity and inadequate response to maximal doses of at least two NSAIDs taken for three months and either sulfasalazine or methotrexate if the patient has peripheral arthritis.87

In France, the CRI (Club Rhumatismes et Inflammation), a subgroup of the French Society for Rheumatology, published in 2006 the guidelines for the use of anti-TNFα agents in patients with AS.88 These guidelines were similar to the ASAS recommendations, but with some differences: three NSAIDs are required before considering the patient to be refractory to conventional treatments and patients without radiological sacroilitis (required for satisfying the modified New York criteria) but with evidences of inflammation at the spine or sacroiliac joints documented by MRI may benefit from anti-TNFα agents.

In clinical practice, rheumatologists have their own opinion on the criteria for initiating anti-TNFα therapy. In a Dutch rheumatologist study, the more important variables for starting anti-TNFα in AS were rate of development of functional impairment, physician global assessment of disease activity, presence of hip arthritis, and physician global assessment of disease severity.89

In addition, patients with poor prognostic factors may be considered as reasonable candidates for anti-TNFα therapy and infliximab. Thus, the clinicians can use the prognostic factors for AS previously identified13,14 for screening the patients who could be treated by anti-TNFα agents.

Predictors of good response to anti-TNFα agents (infliximab and etanercept) have been identified by analyzing the results from randomized controlled trials: a short disease duration, a young age, a low BASFI, an elevated CRP, and a high BASDAI are predictor of a BASDAI improvement >50%.90 In addition, MRI gives also insights in clinical response: spine and sacroiliac joint MRI from patients who participated in controlled studies were analyzed with respect to the presence or not of inflammatory lesions. The likelihood to achieve a BASDAI 50 response was found better in patients with a high MRI spinal score, a short disease duration, and a high CRP.91

However, these factors may help the physician to predict a good clinical response, but it does not mean that the initiation of infliximab or another TNFα antagonist may be restricted to these patients.

Discussion
Collectively, all these data strongly supports that infliximab is highly effective in the treatment of AS patients. This efficacy has been demonstrated in open-label and adequately designed placebo-controlled trials in the different aspects of the disease. Indeed, infliximab is effective in both spinal and peripheral manifestations, in sacroilitis or enthesitis, in reducing signs and symptoms of AS with an improvement of all ASAS core set (disease activity, function, pain, stiffness, quality of life), as well as laboratory parameters of inflammation or active inflammatory lesions detected on MRI. Clinical response is retained for up to five years with maintenance therapy.92

The global safety profile of infliximab seems favorable, but data are supported by a short-term use. Long-term surveillance data begins to be available but more time is needed to know the safety for a longer period of time (especially for a disease which will be treated as long as the treatment is effective and well tolerated). Currently, there are only a few cases of opportunistic infections that have been reported in AS under infliximab and lymphoma is also a rare complicating situation.

What are the pending questions with the use of infliximab in AS?

- **What is the optimal interval of infusion?** According to the clinical trials and the extension protocol studies, the recommended infliximab regimen is an intravenous infusion of 5 mg/kg at weeks 0, 2, and 6, followed by maintenance infusions at six- or eight-week intervals.40,92
- **What is the recommended infliximab dosage?** Most studies evaluated the efficacy of infliximab in AS at a 5 mg/kg dosage. One study tried the medication at a lower dosage, 3 mg/kg, with favorable results.44 However, in a small study involving six patients with SpA, response to 3 mg/kg was inferior to 5 mg/kg.43
- **Do we need to associate methotrexate with infliximab in clinical practice?** This question is relevant since anti-chimeric antibodies may occur with the use of infliximab.40 In RA, it is thought that methotrexate (MTX) reduces the incidence of anti-chimeric antibodies and this associated medication may lower the incidence of acute infusion reaction to infliximab and finally, prevent progressive loss of efficacy. However, we do not have proof that MTX may be useful in AS patients treated by infliximab. One randomized controlled trial conducted...
in the UK evaluated the response to MTX (7.5–10 mg weekly) + infliximab (5 mg/kg given at weeks 0, 2, 6, and then at weeks 14 and 22) compared with MTX + placebo. A higher proportion of patient in the MTX + infliximab group reached an ASAS20 response compared to the MTX + placebo group (50% versus 21%), and the association of MTX did not allow to prolong the response to infliximab. Indeed, in this study, due to a longer interval between infliximab infusions (eight weeks after the induction treatment regimen at weeks 0, 2, and 6), some patients had a flare of their disease.\(^6\) Another multicenter study conducted in France specifically evaluated the need for the patient to be treated continuously by infliximab or only in case of relapse, and the potential benefit of associated MTX treatment. 247 patients participated in this study: 124 received infliximab (5 mg/kg) every six weeks and 123 received on-demand treatment (based upon symptom recurrence). In this latter group, 62 patients received associated treatment with MTX and 61 infliximab alone. At week 58, a greater proportion of patients treated continuously achieved an ASAS20 response than patients in the on-demand group. The association of MTX to infliximab did not improve the proportion of ASAS20 responders. Thus, this study indicates that infliximab is more efficacious when administered continuously (every six weeks) and that the addition of MTX provides no substantial benefit.\(^5\)

A third study in UK confirms these results: in a randomized placebo controlled study, 38 AS patients received either infliximab + MTX or infliximab + placebo. The ASAS 20 response did not differ between the two groups as well as the improvement in MRI spinal score.\(^6\)

- **Does infliximab have an impact on the structural progression of the disease?** Infliximab may suppress active signs of inflammation on MRI, suggesting that the treatment has the potential to slow down the progression of the disease. In other words, infliximab could prevent the development of (new) syndesmophytes and therefore, has a structural effect. In fact, preliminary analysis suggests that infliximab is capable to decelerate progression of spinal structural changes. In the German cohort, patients receiving infliximab for up to two and four years were scored for radiological changes using the modified Stokes Ankylosing Spondylitis Spinal Score (mSASSS) and were compared to published data from the historical OASIS cohort who had no prior use of anti-TNF\(\alpha\) agents.\(^7\) The results showed that the rate of progression of the mSASSS score in patients under infliximab was lower compared to patients from the OASIS cohort (mean mSASSS changes over four years in the infliximab patients were 1.6 ± 2.6 units compared to 4.4 units in OASIS). As previously described, patients who have definite changes at baseline developed more chronic changes than those without damage at baseline. However, if the change of the mean mSASSS in comparison with baseline was significantly different at two years of follow-up, this difference did not persist after four years. Collectively, and although there is no adequate control group in this study, these data suggest that infliximab has the potential to decelerate the progression of spinal structural damage in AS.

However, these data were not confirmed by another radiographic analysis from the patients of the ASSERT trial.\(^8\) In this randomized placebo-controlled study (including 279 AS patients), 201 had radiographic analyses at baseline and after two years of infliximab therapy. The radiographic findings (assessed by the mSASSS) of these patients were compared to those from the OASIS cohort. After two years, the median or the mean change of the mSASSS score did not differ between the two patient populations, suggesting that infliximab has no effect on the progression of the disease. Again, there is no true control group in this study. In addition, the mSASSS evaluates structural damage at the cervical and lumbar spine, without considering the thoracic spine which is the preferential level of syndesmophyte development. And finally, the mSASSS score has a low sensitivity to changes and a minimal period of two years is required to appreciate a radiological progression. For all these reasons, the radiographic analysis from the ASSERT study (and from the German cohort) is to interpret with caution.

- **Infliximab is associated with paradoxal and unexplained effects:**
  - Infliximab is approved in the treatment of psoriasis and psoriatic arthritis. However, there are some unexpected reports of exacerbation or new onset of psoriasis in patients who had previously no skin disease. This paradoxical effect has been described with the three available anti-TNF\(\alpha\) including infliximab. 120 cases were recently reviewed. The prevalence of this phenomenon has been estimated to range between 1.5% to 5% of patients treated by TNF\(\alpha\) antagonists. Underlying diseases may be AS, psoriatic arthritis, RA, or Crohn’s disease.
The clinical features included psoriasis, palmo-plantar pustular lesions, psoriasis of the nail or psoriasiform exanthema. The delay between initiation of the anti-TNFα antagonist and appearance of skin lesions varied from the first administration to 63 months. Various outcomes were reported after cessation or switching the TNFα antagonist, with or without anti-psoriatic adjuvant therapy. The underlying pathomechanisms of these psoriasis-induced lesions remain unclear and certainly involved different factors (infection, change in cytokine balance).39

- Infliximab is effective in the treatment of Crohn’s disease. However, there are exceptional cases of Crohn’s disease development in AS patients receiving etanercept100–102 but also infliximab (Toussirot, personal observation). However, one may remark that the introduction of the TNFα blockers may demask an undiagnosed but underlying IBD in these patients who are predisposed to have inflammation of the gut.

- Finally, various infectious granulomatous diseases have been described in patients with AS or RA with the use of TNFα antagonists such as sarcodiosis, mainly with etanercept but also infliximab.103,104

Conclusion
Infliximab is effective in the treatment of active AS and has demonstrated in the short term and medium term, significant improvements of signs and symptoms of AS. Some questions persist such as the real impact of this treatment on the progression of radiological damage. On the contrary of NSAIDs,105 the results of the published studies failed to demonstrate such an effect and thus, we cannot currently consider infliximab (and other TNFα antagonists as well) as a disease-controlling anti-rheumatic therapy. Nevertheless, due to the high level of efficacy obtained in the different domains of the disease, infliximab represents undoubtedly a considerable advancement in the therapeutic management of AS, and this medication has “really changed the life of the patients.”

Disclosure
The authors report no conflicts of interest in this work.

References


