How early should ankylosing spondylitis be treated with tumour necrosis factor blockers?

J Sieper, M Rudwaleit

Recognition and treatment of ankylosing spondylitis (AS) in the early stages of the disease has yet to be established. This paper considers the evidence available and the questions that need to be answered regarding the benefits of early diagnosis and treatment with tumour necrosis factor (TNF) blockers in AS. The authors conclude that AS can and has to be diagnosed earlier than is being done at present, before radiological changes are evident, and the potential of TNF blockers to induce long term remission if given early enough needs to be clarified.

Ankylosing spondylitis (AS) is an important chronic inflammatory disease with a prevalence between 0.5% and 1.0%, which starts in about 80% of patients before the age of 30 years, thus early in life. Although temporary remissions and mild courses are part of the disease spectrum about 30% of patients have a constantly high disease activity over one year, as judged by a high disease activity index and elevated C-reactive protein. Furthermore, it has been estimated that at least 30% of patients do develop severe spinal restriction during the natural course of the disease. Socioeconomic costs can be high and increase with higher disease activity and lower function. Thus, effective early treatment should be focused on patients with high disease activity and on those who are likely to develop radiological progression and rapid functional deterioration during the natural course of their disease.

Until recently the treatment options for AS were limited. Regular physiotherapy and treatment with non-steroidal anti-inflammatory drugs (NSAIDs) were the only available symptomatic treatment. Other treatment options such as disease modifying antirheumatic drugs (DMARDs) or steriods, which are quite effective in other chronic inflammatory diseases such as rheumatoid arthritis, have no or only a limited effect. About 75% of patients with AS show a good or very good response to a full dose of NSAIDs in 48 hours, in contrast with only 15% of patients with mechanical back pain. Furthermore, a recent study has shown that patients with AS treated continuously over two years with a daily dose of NSAIDs had less radiological progression compared with those patients who took NSAIDs irregularly—that is, on demand. Thus because of the good efficacy for acute symptoms and because of the potential to retard long term damage, AS patients with inflammatory back pain (in contrast with patients with mechanical low back pain) should be probably treated long term with NSAIDs—once the diagnosis is made and the disease is active.

However, about 20–50% of AS patients still have active disease despite treatment with NSAIDs. For these patients, tumour necrosis factor (TNF) blocking agents have meant a breakthrough in treatment. Both the TNF blocking agents, infliximab and etanercept, have shown a surprisingly large and rapid effect on every aspect of active AS: disease activity including acute phase reactants, pain and morning stiffness, function, spinal mobility, peripheral arthritis, and enthesisitis. In the different studies investigating these two compounds a 50% improvement of the disease activity was demonstrated in about half of the treated patients who were refractory to NSAID therapy and physiotherapy. Preliminary data indicate that the third TNF blocking agent, adalimumab, may be similarly effective. In parallel with a clear improvement in signs and symptoms, acute bony inflammation in the spine and sacroiliac joints, as detected by magnetic resonance imaging (MRI), was reduced impressively by this treatment.

Thus, this high efficacy in patients otherwise refractory to treatment raises the question: How early in the course of their disease should these patients be treated with TNF blockers? In the last two decades an impressive paradigm shift in the treatment of rheumatoid arthritis took place. For this disease it has been demonstrated that with early and aggressive treatment even remission can be achieved in a substantial proportion of patients, and that radiological progression can be stopped and even healing of bony erosions occurs.

With regard to early treatment of AS with TNF blockers some questions need to be answered, and these will be addressed in this article.

**HOW CAN ANKYLOSING SPONDYLITIS BE DIAGNOSED EARLIER?**

A major obstacle to any early treatment is the big gap of 5–10 years between the first (chronic) symptoms of the disease and the diagnosis of the disease. One reason for this is certainly the low awareness of AS among non-rheumatologists. It is also a major challenge for any physician in primary care to think of and to identify patients with inflammatory spine disease among the large group of patients with chronic back pain. However, the relative late appearance of radiographic sacroiliitis, by up to several years after the first symptoms, is another important reason for this delay because usually the diagnosis is made according to the modified New York criteria; for this radiological sacroiliitis grade 2 bilaterally or grade 3 or 4 unilaterally is mandatory. However, it has become obvious, specially through the detection of acute sacroiliitis (and also inflammation of the spine) by MRI, that there is a continuum in spinal inflammation from a pre-radiographic to a radiographic stage (fig 1). We have argued recently that this should be treated as one disease, that radiological sacroiliitis should not be an essential diagnostic parameter but rather a measure of severity or chronicity, and that a diagnosis in the pre-radiographic stage can be made if a combination of clinical, laboratory, and imaging (especially MRI) parameters is applied.

Currently the ASessment in Ankylosing Spondylitis (ASAS) group is developing a new classification/diagnostic criteria for AS, which will also include the
early (pre-radiographic) forms of AS—an essential and crucial step for an early diagnosis.

ARE PATIENTS EARLY IN THEIR DISEASE SIMILARLY ACTIVE COMPARED WITH LATER IN THE DISEASE?

We have recently started a prospective cohort study of patients with early spondyloarthritides with special focus on AS (German Spondyloarthritis Inception Cohort (GESPIC)) in Germany. In this cohort we have included 120 AS patients (with radiographic sacroiliitis) with a disease duration between 5 and 10 years, 105 patients with a disease duration of less than 5 years and 200 patients with a predominant axial spondylarthropathy (SpA) without radiographic sacroiliitis. The mean disease activity index (the Bath Ankylosing Spondylitis Disease Activity Index) and the mean level of pain in all three groups were exactly the same, demonstrating that the degree of signs and symptoms is already high early in the disease and independent of the presence or absence of radiological changes. Thus to alleviate the signs and symptoms, effective therapies have to be offered to patients early in the disease. Furthermore, on analysis of 99 AS patients treated with TNF blockers we were able to show that patients with a shorter disease duration have a better response rate compared with those patients with a longer disease duration: the percentage of patients with an at least 50% improvement of their disease activity was 73% for a disease duration less than 10 years, 58% for a disease duration between 10 and 20 years and only 31% for a disease duration of more than 20 years (fig 2). This most probably reflects that symptoms early in disease are mainly due to inflammation whereas later in the disease symptoms are caused by a mixture of inflammation, structural spinal damage, and secondary damage to soft tissue such as muscles and ligaments.

CAN RADIOLOGICAL PROGRESSION IN ANKYLOSING SPONDYLITIS BE STOPPED?

Another major argument in favour of early treatment would be cessation of radiological progression, as has been shown in rheumatoid arthritis, in order to prevent disability. At the present time we only have preliminary evidence—from a comparison of radiological progression of the spine over a two year period between a group of patients treated with infliximab and the natural course of the disease in a control group over the same time—that this might be the case. The strongest evidence for such a potential of TNF blockers comes from MRI follow up studies during treatment with infliximab and etanercept. These studies have shown that acute inflammatory lesions in the spine and sacroiliac joints can be effectively suppressed (fig 3), also suggesting that bony destruction and bony proliferation can be prevented. However, such a link between acute inflammation and structural damage, although likely, has yet to be proved.

ARE THERE PROGNOSTIC FACTORS THAT COULD PREDICT BAD OUTCOME?

An effective treatment early in disease should be offered preferentially to patients with a bad prognosis. However, currently only a few studies are addressing this question. Amor et al undertook an analysis of 151 AS patients followed up retrospectively for 10 years and identified presence of hip arthritis, elevated erythrocyte sedimentation rate (ESR), and a bad response to NSAIDs as variables with the highest likelihood ratio to predict bad outcome. van der Heijde et al prospectively followed up 137 AS patients from the OASIS cohort and correlated measures at baseline with radiological progression of the spine after four years. They found that radiological damage at baseline, male sex, hip arthritis, and extraspinal manifestations such as uveitis and peripheral arthritis correlated best with structural damage. More data are expected in the near future from the analysis of ongoing cohort studies concentrating on patients with early disease.

CAN REMISSION BE INDUCED IN ANKYLOSING SPONDYLITIS IF TREATMENT WITH TNF BLOCKERS IS STARTED EARLY?

Treatment with TNF blockers has to be continued long term in most of the patients with established AS. We have recently shown that a relapse occurs after as short as a few weeks to a few months when treatment with infliximab or etanercept was discontinued in AS patients with a mean disease duration of more than 10 years. No data are available at the moment regarding whether remission is sustained after withdrawal of TNF blockers if patients are treated earlier, as has been suggested in rheumatoid arthritis. However, we have shown that remission can be achieved in a higher percentage of patients with shorter compared with longer disease duration (in 35% of patients with less than 10 years since first symptoms, but in none of the patients with more than 20 years and in only 24% of those with a disease duration between 10 and 20 years). Thus, patients early in the course of their disease clearly show a higher rate of remission if treated with infliximab or etanercept. Data are needed about the potential of such treatment to induce long term remission if patients are treated early.
HOW CAN PATIENTS WITH EARLY ANKYLOSING SPONDYLITIS BE BETTER IDENTIFIED IN PRIMARY CARE?

Although there is some evidence that patients early in the course of their disease might benefit most from treatment with TNF blockers there are some open questions, which have been discussed here. These will only be addressed if these patients have a chance to be treated earlier. Thus, strategies have to be developed to alert the primary care physician when to think of inflammatory spine disease in patients with chronic back pain and when to refer these patients to the rheumatologist for a definite diagnosis. Most of the data on treatment of early rheumatoid arthritis have been raised in the last 10–15 years following the introduction of early arthritis clinics in many parts of the world. We have also recently proposed screening measures for early referral which are easy to apply. Such measures have to be sensitive and specific for the disease under consideration, easy to apply by the non-specialist, and should not be too expensive. We have calculated that a diagnosis of axial SpA can be made in every third to fifth patient who fulfils the criteria for the disease. The disappearance of inflammation in the spine and sacroiliac joints during treatment, as detected by MRI, is a demonstration of the great efficacy of the TNF blockers and also suggests that structural damage may also be prevented. Whether and how these new treatments also have the potential to induce long term remission if given early enough has to be shown in the future.

SUMMARY

In summary, AS can and has to be diagnosed earlier than is being done at present, even before radiological changes are evident. Therapies such NSAIDs and TNF blockers are most effective for the signs and symptoms that are caused by inflammation. This is probably a major reason why these drugs are so effective if used early in the course of the disease. The disappearance of inflammation in the spine and sacroiliac joints during treatment, as detected by MRI, is a demonstration of the great efficacy of the TNF blockers and also suggests that structural damage may also be prevented. Whether and how these new treatments also have the potential to induce long term remission if given early enough has to be shown in the future.

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