Clinical efficacy and safety of abatacept in methotrexate-naive patients with early rheumatoid arthritis and poor prognostic factors


ABSTRACT

Objectives: To assess the efficacy and safety of abatacept in methotrexate-naive patients with early rheumatoid arthritis (RA) and poor prognostic factors.

Methods: In this double-blind, phase IIIb study, patients with RA for 2 years or less were randomly assigned 1:1 to receive abatacept (~10 mg/kg) plus methotrexate, or placebo plus methotrexate. Patients were methotrexate-naive and seropositive for rheumatoid factor (RF), anticyclic citrullinated protein (CCP) type 2 or both and had radiographic evidence of joint erosions. The co-primary endpoints were the proportion of patients achieving disease activity score in 28 joints (DAS28)-defined remission (C-reactive protein) and joint damage progression (Genant-modified Sharp total score; TS) at year 1. Safety was monitored throughout.

Results: At baseline, patients had a mean DAS28 of 6.3, a mean TS of 7.1 and mean disease duration of 6.5 months. 96.5% and 89.0% of patients were RF or anti-CCP2 seropositive, respectively. Over 1 year, a significantly greater proportion of abatacept plus methotrexate-treated patients achieved remission (41.4% vs 23.3%; p<0.001) and there was significantly less radiographic progression (mean change in TS 0.63 vs 1.06; p = 0.040) versus methotrexate alone. Over 1 year, the frequency of adverse events (84.8% vs 83.4%), serious adverse events (7.8% vs 7.9%), serious infections (2.0% vs 2.0%), autoimmune disorders (2.3% vs 2.0%) and malignancies (0.4% vs 0%) was comparable for abatacept plus methotrexate versus methotrexate alone.

Conclusions: In a methotrexate-naive population with early RA and poor prognostic factors, the combination of abatacept and methotrexate provided significantly better clinical and radiographic efficacy compared with methotrexate alone and had a comparable, favourable safety profile.

In rheumatoid arthritis (RA), persistent synovitis, early erosions and the presence of rheumatoid factor (RF) and anti-cyclic citrullinated protein (CCP) type 2 antibodies are prognostic indicators of joint destruction and loss of function. The initiation of intensive treatment early in the course of disease is now an accepted paradigm in the treatment of RA, with an increasing emphasis on tight disease control and clinical remission as a treatment goal.

Studies comparing biological agents in combination with methotrexate compared with methotrexate alone have demonstrated significant benefit when treatment is initiated early. These trials have also highlighted that, although methotrexate monotherapy can be efficacious, it does not provide optimal disease control in a proportion of patients.

Abatacept is a soluble, fully human, recombinant fusion protein that selectively modulates the CD80/CD86:CD28 co-stimulatory signal for T-cell activation. The ability of abatacept to modulate the activation of T cells, including naive T cells, and the role of T cells in initiating disease suggests that abatacept has the potential to impact the progression of pathology early in the course of disease.

The sustained efficacy and safety of abatacept has previously been demonstrated in patients with moderate-to-severe established RA who have had an inadequate response to methotrexate and/or anti-tumour necrosis factor agents. Here, we report 1-year data from a study that assessed the efficacy, safety and tolerability of abatacept plus methotrexate compared with methotrexate alone, in methotrexate-naive patients with early RA (<2 years). The patients in this study represented a particularly poor prognosis population, because the inclusion criteria required all patients to have erosions and to be seropositive for RF and/or anti-CCP2, which are associated with poor radiological outcomes.

PATIENTS AND METHODS

Patients

Eligible patients were 18 years of age or older, with RA for 2 years or less, at least 12 tender and 10 swollen joints, C-reactive protein (CRP) 0.45 mg/dl or greater, RF and/or anti-CCP2 seropositivity and radiographic evidence of bone erosion of the hands/wrists/feet. Patients were either methotrexate-naive or had previous exposure of 10 mg/week or less for 3 weeks or less, with none administered for 3 months before providing informed consent (there were no requirements relating to the reason for discontinuation of previous methotrexate therapy).

Study design

This was a multi-national, randomised, double-blind, 2-year study (ClinicalTrials.gov identifier: NCT00122382). The protocol and patients’ informed consent received institutional review board/independent ethics committee approval, and the study was conducted in accordance with...
the Declaration of Helsinki and was consistent with International Conference on Harmonisation Good Clinical Practice. Patients, sites and the site conducting radiographic assessment remained blinded to treatment assignments until the end of the study.

Patients were randomly assigned 1:1 to receive abatacept (10 mg/kg according to weight range) plus methotrexate or placebo plus methotrexate for 1 year by intravenous infusion on days 1, 15 and 29, and every 4 weeks thereafter. Methotrexate was initially dosed at 7.5 mg/week and subsequently increased to 15 mg at week 4 and to 20 mg at week 8, at which dose it was maintained until study completion. Dose reduction was permitted to a minimum of 15 mg/week due to toxicity or intolerability.

Women who were pregnant or breastfeeding were excluded, and patients were required to practice effective contraceptive measures for the study duration. Patients were excluded if they had active *Mycobacterium tuberculosis* (tuberculosis) requiring treatment within 3 years. Patients with a positive purified protein derivative test were eligible if treatment for latent tuberculosis had been initiated (according to local guidelines) and there was no evidence of active tuberculosis by chest x ray at enrollment.

Stable low-dose oral corticosteroids (≤10 mg prednisone equivalent, daily) were permitted, plus up to two corticosteroid “pulses” (>10 mg prednisone or equivalent oral corticosteroids for at least three consecutive days or injectable corticosteroids) in any 6-month period. After 6 months, the addition of one non-biological disease-modifying antirheumatic drug (DMARD) was permitted.

**Clinical assessments**

**Efficacy assessments**

**Co-primary endpoints**
Remission at year 1, defined as a disease activity score in 28 joints (DAS28; CRP) of less than 2.6; structural damage at year 1, measured using the Genant-modified Sharp scoring system total score (TS) with a maximum possible score of 290.

**Secondary endpoints**
These were assessed at year 1 and included American College of Rheumatology (ACR) 50 responses; major clinical response (MCR; ACR70 maintained for ≥6 consecutive months); DAS28 (CRP) scores; Genant-modified Sharp erosion score (ES; maximum possible 145) and joint-space narrowing score (JSN; maximum possible 145); physical function (defined as an improvement of ≥0.3 units from baseline in the health assessment questionnaire disability index; HAQ-DI) and health-related quality of life (HRQoL) measured using the short form (SF)-36, with an improvement of 3 units or more considered to be clinically meaningful.

The proportion of patients achieving ACR70 and ACR90 responses and the proportion of patients without radiographic progression (change in TS from baseline of ≤0) were also evaluated.

**Safety assessments**
All patients who received one dose of abatacept were evaluated, and adverse events and serious adverse events were classified using the Medical Dictionary for Regulatory Activities version 10.1. Autoimmune and infusional events were taken from a predefined list of events of interest. Acute infusional events were any adverse event occurring within 1 h after initiation of infusion.

**Statistical analysis**

**Power calculation**
It was estimated that 500 patients randomly assigned 1:1 to each group would yield 99% power and detect a 20% difference in DAS28 (CRP)-defined remission rates at the 5% level (two-tailed test). A response rate of 15% at year 1 in the methotrexate group and an overall 15% dropout rate was assumed. Based on the hierarchical testing procedure for the co-primary endpoints, this sample size also allowed the detection of a treatment difference of 1.6 (common SD 5) with a power of 90% for mean change from baseline in TS.

**Data analysis**
Unless otherwise stated, efficacy analyses were performed on all patients randomly assigned and treated: for DAS28 (CRP)-defined remission, ACR and HAQ-DI responses; patients who discontinued were considered non-responders subsequent to discontinuation. For analysis of the co-primary endpoints, patients who received more than one corticosteroid pulse (oral >10 mg/day prednisone equivalent for ≥5 consecutive days, intramuscularly or intravenously) during year 1, or who received more than two intra-articular injections (corticosteroid...
TABLE 1 Summary of baseline demographics, clinical characteristics and concomitant medications

<table>
<thead>
<tr>
<th></th>
<th>Abatacept + methotrexate (n = 256)</th>
<th>Placebo + methotrexate (n = 253)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age in years, mean (SD)</td>
<td>50.1 (12.4)</td>
<td>49.7 (13.0)</td>
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<tr>
<td>Gender, % female</td>
<td>76.6</td>
<td>78.7</td>
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<tr>
<td>Race, % white</td>
<td>78.9</td>
<td>86.6</td>
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<tr>
<td>Geographical region, n (%)</td>
<td></td>
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<tr>
<td>North America, n (%)</td>
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<tr>
<td>South America, n (%)</td>
<td>46 (18.0)</td>
<td>40 (15.8)</td>
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<td>Europe, n (%)</td>
<td>103 (40.2)</td>
<td>102 (40.3)</td>
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<tr>
<td>Rest of world, n (%)</td>
<td>88 (34.4)</td>
<td>95 (37.5)</td>
</tr>
<tr>
<td>Disease duration, mean months (SD)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tender joints, mean (SD)</td>
<td>31.3 (14.8)</td>
<td>30.8 (14.0)</td>
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<tr>
<td>Swollen joints, mean (SD)</td>
<td>22.9 (11.3)</td>
<td>21.9 (10.1)</td>
</tr>
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<td>CRP levels, mg/dl, mean (SD)</td>
<td>3.1 (3.1)</td>
<td>3.6 (5.0)</td>
</tr>
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<td>DAS28 (CRP)*, mean (SD)</td>
<td>6.2 (7.5)</td>
<td>6.7 (7.1)</td>
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<tr>
<td>HAQ-DI (0–3), mean (SD)</td>
<td>1.7 (0.7)</td>
<td>1.7 (0.7)</td>
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<tr>
<td>RF positive, n (%)</td>
<td>246 (96.1)</td>
<td>245 (96.8)</td>
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<tr>
<td>Anti-CCP2 positive, n (%)</td>
<td>236 (92.2)</td>
<td>217 (85.8)</td>
</tr>
<tr>
<td>RF and anti-CCP2 positive, n (%)</td>
<td>227 (88.7)</td>
<td>211 (83.4)</td>
</tr>
<tr>
<td>Total x ray score, mean (SD)</td>
<td>7.5 (9.7)</td>
<td>6.7 (8.8)</td>
</tr>
<tr>
<td>Total x ray score, median (range)</td>
<td>3.4 (0.0–57.3)</td>
<td>3.8 (0.0–54.6)</td>
</tr>
<tr>
<td>Erosion score, mean (SD)</td>
<td>5.4 (6.1)</td>
<td>4.8 (5.4)</td>
</tr>
<tr>
<td>Erosion score, median (range)</td>
<td>2.9 (0–29.1)</td>
<td>3.1 (0–36.9)</td>
</tr>
<tr>
<td>Joint space narrowing, mean (SD)</td>
<td>2.1 (4.2)</td>
<td>1.9 (4.0)</td>
</tr>
<tr>
<td>Joint space narrowing, median (range)</td>
<td>0.2 (0–28.2)</td>
<td>0.5 (0–28.0)</td>
</tr>
<tr>
<td>Total patients on antirheumatic concomitant medications at randomisation, n (%)</td>
<td>243 (94.9)</td>
<td>236 (93.3)</td>
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<tr>
<td>Corticosteroids, oral and/or injectable</td>
<td>131 (51.2)</td>
<td>124 (49.0)</td>
</tr>
<tr>
<td>Corticosteroids, oral &lt;10 mg/day</td>
<td>95 (37.1)</td>
<td>86 (34.0)</td>
</tr>
<tr>
<td>NSAID</td>
<td>203 (79.3)</td>
<td>201 (79.4)</td>
</tr>
<tr>
<td>Other non-biological DMARD</td>
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<td></td>
</tr>
<tr>
<td>Chloroquine</td>
<td>7 (2.7)</td>
<td>10 (4.0)</td>
</tr>
<tr>
<td>Hydroxychloroquine</td>
<td>3 (1.2)</td>
<td>4 (1.6)</td>
</tr>
<tr>
<td>Sulfasalazine</td>
<td>4 (1.6)</td>
<td>5 (2.0)</td>
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</tbody>
</table>
| *n = 252 for methotrexate; †n = 254 for abatacept plus methotrexate, n = 251 for methotrexate; †n = 253 for abatacept plus methotrexate.*

Sensitivity analysis
For the proportion of patients without radiographic progression, an analysis was performed in which all patients who received additional non-biological DMARD were considered non-responders. For DAS28 (CRP)-defined remission and the proportion of patients without radiographic progression, additional analyses were performed in which all patients who received additional non-biological DMARD and/or a steroid pulse were considered non-responders. For mean change from baseline to year 1 in TS, ES and JSN, an analysis was performed considering only completers.

Comparisons between treatment groups
For DAS28 (CRP)-defined remission, ACR, MCR and HAQ-DI responder rates at year 1, a continuity-corrected χ² test was used. For change from baseline for DAS28 (CRP), HAQ-DI and SF-36, comparisons were based on an analysis of covariance model, including treatment as the main factor and baseline value as a covariate. A non-parametric analysis of covariance model was used for change from baseline to year 1 in TS, ES and JSN.

RESULTS
Baseline demographics and characteristics, and patient disposition
Five hundred and nine patients (predominantly South American (40.3%) and European (36.0%)) were randomly assigned to receive abatacept plus methotrexate (n = 256) or methotrexate alone (n = 253). Demographics and baseline clinical characteristics were comparable between groups, with an overall mean disease duration of 6.5 months (table 1). At baseline, patients had a high disease activity evidenced by an overall mean DAS28 (CRP) score of 6.3, and mean tender and swollen joint counts of 31.0 and 22.4, respectively. In total, eight (3.1%) patients in the abatacept plus methotrexate group and two (0.8%) patients in the methotrexate alone group had received methotrexate before...
randomisation. The mean baseline TS was 7.1. A total of 491 (96.5%) and 453 (89.0%) patients was seropositive for RF and anti-CCP2, respectively; 86.1% of patients were seropositive for both. A total of 232 (90.6%) patients in the abatacept plus methotrexate group and 227 (89.7%) patients in the methotrexate group completed year 1 (fig 1). Fewer patients receiving abatacept plus methotrexate discontinued due to lack of efficacy (0% vs 3.2%) or adverse events (3.5% vs 4.3%), than those receiving methotrexate alone.

Concomitant antirheumatic medications
Concomitant medications at baseline are shown in table 1. At month 6, the mean dose of methotrexate was 18.9 mg/week (SD 3.2) in the abatacept plus methotrexate group and 18.9 mg/week (SD 3.4) in the methotrexate group. At year 1 the mean dose of methotrexate was 18.1 mg/week (SD 3.0) in the abatacept plus methotrexate group and 19.0 mg/week (SD 3.8) in the methotrexate group. The addition of non-biological DMARD was permitted after month 6: six (2.3%) patients in the abatacept plus methotrexate group and 17 (6.7%) patients in the methotrexate group received additional non-biological DMARD. Over 1 year, 136 (53.1%) patients in the abatacept plus methotrexate group and 136 (53.8%) patients in the methotrexate group received oral steroids, and the mean dose was 6.7 mg (SD 3.2) versus 8.3 mg (SD 7.1), respectively. A total of 10 (3.9%) patients in the abatacept plus methotrexate group and 20 (7.9%) patients in the methotrexate group received oral steroid pulses (>10 mg); the mean dose of these was 18.2 mg (SD 4.7) versus 27.7 mg (SD 16.7), respectively.

Efficacy
Clinical efficacy
A significantly higher proportion of patients in the abatacept plus methotrexate group achieved DAS28 (CRP)-defined remission compared with the methotrexate group by day 57 (fig 2A), and a significant difference was maintained up to year 1 (41.4% vs 23.3%, respectively, at year 1; p<0.001). When patients for whom a non-biological DMARD was initiated were considered non-responders, the proportion of responders was 41.0% (95% CI 35.0 to 47.0) in the abatacept plus methotrexate group compared with 21.7% (95% CI 16.7 to 26.8) in the methotrexate group. At year 1, disease activity was significantly reduced with abatacept plus methotrexate compared with methotrexate alone (adjusted mean changes from baseline in DAS28 (CRP) were −3.22 (SE 0.09) and −2.49 (SE 0.09), respectively, p<0.001). The proportion of patients achieving ACR50 or ACR70 responses was significantly higher in the abatacept plus methotrexate compared with the methotrexate group by day 57 (fig 2B and C), and a significant difference was maintained up to year 1. For abatacept plus methotrexate compared with methotrexate alone at year 1, the proportion of patients achieving ACR50 was 57.4 versus 42.3% (p<0.001), ACR70
42.6 versus 27.3% (p < 0.001) and ACR90 16.4 versus 6.7% (p = 0.001; fig 2B–D). Over 1 year, 27.3% of patients achieved an MCR versus 11.9% of patients receiving methotrexate alone (p < 0.001).

Radiographic progression
Baseline radiographic scores are shown in table 1. At month 6 and year 1, observed data were available for 234 (91.4%) and 218 (85.2%) patients in the abatacept plus methotrexate group and 235 (92.9%) and 223 (88.1%) patients in the methotrexate group, respectively. Figure 3 presents mean changes in Genant-modified Sharp scores at month 6 (based on observed data) and year 1 (linear extrapolation applied). At year 1, changes from baseline in TS and ES were significantly lower in the abatacept plus methotrexate group compared with the methotrexate group (p = 0.040 and p = 0.033, respectively; fig 3A and B); mean changes in JSN scores were minimal and were comparable between the two groups (p = 0.353; fig 3C). Results were comparable when only completers were considered at the 1-year time point (mean changes from baseline in the abatacept plus methotrexate vs the methotrexate groups were 0.57 vs 1.05 (p = 0.020) for TS, 0.46 vs 0.89 (p = 0.025) for ES and 0.11 vs 0.16 (p = 0.246) for JSN, respectively). The median change from baseline in TS, ES and JSN was 0 for both groups. The cumulative probability plot (fig 3D) shows the distribution of change from baseline in TS over 1 year. At year 1, the proportion of patients with no radiographic progression (TS (≤0) was 61.2% (95% CI 55.0 to 67.5) in the abatacept plus methotrexate group versus 52.9% (95% CI 46.6 to 59.2) in the methotrexate group, with an estimated difference of 8.3% (95% CI −1.0 to 17.5). When patients for whom a non-biological DMARD was initiated were considered progressors, the proportion of patients without progression was 59.1% (95% CI 52.9 to 65.3) compared with 48.3% (95% CI 42.1 to 54.6) for the abatacept plus methotrexate versus methotrexate groups, and the estimate of difference was 10.7% (95% CI 1.4 to 20.0). When patients who received steroid pulses were also considered progressors, the proportion of patients without progression was 58.3% (95% CI 52.1 to 64.5) compared with 47.1% (95% CI 40.8 to 53.4) and the estimate of difference was 11.2% (95% CI 1.8 to 20.5).

Physical function and HRQoL
At year 1, 184 (71.9%) patients in the abatacept plus methotrexate group compared with 157 (62.1%) patients in the methotrexate group achieved a change from baseline of 0.3 units or greater in HAQ-DI (p = 0.024 for the abatacept plus methotrexate versus methotrexate groups). The adjusted mean change from baseline in HAQ-DI was −0.96 (SE 0.04) for the abatacept plus methotrexate group and −0.76 (SE 0.04) for the methotrexate group, and the adjusted treatment difference did not cross zero (−0.20; 95% CI −0.31 to −0.08). Significant improvements in the SF-36 were also observed: the adjusted mean change from baseline was 11.68 (SE 0.62) versus 9.18 (SE 0.63) for the physical component summary and 8.15 (SE 0.64) versus 6.34 (SE 0.64) for the mental component summary.
Table 2 Summary of safety at year 1

<table>
<thead>
<tr>
<th></th>
<th>Abatacept + methotrexate (n = 256)</th>
<th>Placebo + methotrexate (n = 253)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse events</td>
<td>217 (84.8)</td>
<td>211 (83.4)</td>
</tr>
<tr>
<td>Discontinuations due to adverse events</td>
<td>8 (3.1)</td>
<td>11 (4.3)</td>
</tr>
<tr>
<td>Serious adverse events</td>
<td>20 (7.8)</td>
<td>20 (7.9)</td>
</tr>
<tr>
<td>Discontinuations due to serious adverse events</td>
<td>3 (1.2)</td>
<td>3 (1.2)</td>
</tr>
<tr>
<td>Infections</td>
<td>132 (51.6)</td>
<td>139 (54.9)</td>
</tr>
<tr>
<td>Serious infections</td>
<td>5 (2.0)</td>
<td>5 (2.0)</td>
</tr>
<tr>
<td>Autoimmune events</td>
<td>6 (2.3)</td>
<td>5 (2.0)</td>
</tr>
<tr>
<td>Acute infusional events</td>
<td>16 (6.3)</td>
<td>5 (2.0)</td>
</tr>
<tr>
<td>Malignancies</td>
<td>1 (0.4)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>Deaths</td>
<td>2 (0.8)</td>
<td>4 (1.6)</td>
</tr>
</tbody>
</table>

Extended report

Safety

At year 1, the frequency of adverse events, serious adverse events and discontinuations as a result of serious adverse events was comparable between groups (table 2). The proportion of adverse events that led to discontinuation was 5.1% in the abatacept plus methotrexate group compared with 4.3% in the methotrexate group. The most frequently reported adverse events (>10% of patients in the abatacept plus methotrexate group) were nausea, upper respiratory tract infection and headache. There were six deaths; two (0.8%) in the abatacept plus methotrexate group and four (1.6%) in the methotrexate group. Of the two deaths in the abatacept plus methotrexate group, one patient had pneumonia and severe gastrointestinal bleeding and the other had an acute myocardial infarction.

The most frequent infections were upper respiratory tract infection in 26 (10.2%) versus 26 (10.3%) patients, nasopharyngitis in 21 (8.2%) versus 26 (10.3%) patients and influenza in 19 (7.4%) versus 25 (9.1%) patients in the abatacept plus methotrexate versus methotrexate groups, respectively. Serious infections were experienced by five (2.0%) patients in the abatacept plus methotrexate group (pneumonia, gastroenteritis, cellulitis, pseudomonal lung infection and postoperative wound infection, one patient each) and five (2.0%) patients in the methotrexate group (pneumonia, three patients; gastroenteritis, one patient; and breast cellulitis and staphylococcal infection, both in the same patient). No patients in the abatacept plus methotrexate group discontinued due to an infection. No opportunistic infections or cases of tuberculosis were reported in either group.

One malignancy was reported—pancreatic cancer in the abatacept plus methotrexate group. The patient was 67 years of age and had received 12 infusions of abatacept (750 mg) before the event. The event was considered unlikely to be related to study medication and abatacept was discontinued.

Six patients (2.3%) in the abatacept plus methotrexate group and five patients (2.0%) in the methotrexate group had adverse events that were recorded under the category of autoimmunity disorders. Sjogren’s syndrome, sicca syndrome, systemic lupus erythematosus, psoriasis and atrophic gastritis occurred in one patient in each group; erythema nodosum occurred in one patient in the abatacept plus methotrexate group.

Sixteen patients (6.5%) in the abatacept plus methotrexate group experienced acute infusional events compared with five patients (2.0%) in the methotrexate group. The most common events were dizziness (five (2.0%) vs two (0.8%) patients for abatacept plus methotrexate vs methotrexate alone, respectively); all other events were reported in two or less than two patients in either group. All acute infusional events were mild or moderate in intensity, except for one case of severe urticaria in the abatacept plus methotrexate group.

Two abatacept patients became pregnant (protocol violations). One patient had received one abatacept infusion, had a positive urine pregnancy test on day 1 and experienced a spontaneous abortion between days 1 and 30. One patient received nine infusions of abatacept and had a positive urine pregnancy test on day 253. Pregnancy was confirmed by ultrasound and was subsequently terminated by induced abortion on day 281. Both patients were discontinued from the study as a result of pregnancy.

DISCUSSION

This is the first trial to examine the impact of the modulation of T-cell co-stimulation with abatacept in patients with early RA. Both primary endpoints were met. Abatacept plus methotrexate was significantly better at inducing DAS28 (CRP)-defined remission and reducing radiographic progression compared with methotrexate alone. Sensitivity analyses demonstrated that background medications did not impact the primary endpoints. Consistent with previous studies of abatacept in patients with established RA,22 23 clinical responses (ACR criteria) and clinically meaningful improvements in both physical function and HRQoL were significantly higher in the abatacept plus methotrexate group compared with the group receiving methotrexate alone. In addition, the safety and tolerability of abatacept plus methotrexate was generally comparable to methotrexate alone.

The clinical benefits observed with abatacept plus methotrexate were coupled with significantly lower rates of radiographic progression compared with the methotrexate-alone group. In a patient population with established RA and an inadequate response to methotrexate from the Abatacept in Inadequate Responders to Methotrexate (AIM) study, the rate of structural damage progression was significantly lower during the second year of treatment compared with the first year.19 The 2-year results from the current trial will determine whether this effect is also observed in patients with early RA. In contrast to the lower rates of progression in TS and ES in the abatacept plus methotrexate group, rates of JSN progression were minimal in both groups and were not significantly different. Consistent with other trials in patients with early RA,6–9 baseline values and (p = 0.005 for the physical component summary and p = 0.046 for the mental component summary for the abatacept plus methotrexate versus methotrexate groups).

(p = 0.005 for the physical component summary and p = 0.046 for the mental component summary for the abatacept plus methotrexate versus methotrexate groups).
the progression of JSN scores were low in both treatment arms, and it is possible that there was not enough power to detect small differences in rates of progression between the two groups.

On-drug remission is now an important and realistic therapeutic goal, which is increasingly reflected in the design of clinical trials for RA. This abatacept study and an early RA etanercept trial are the first studies of biological agents to use DAS28-defined remission as a primary endpoint. Remission was also the primary endpoint of an early RA trial comparing combination non-biological DMARD therapy with monotherapy, although in that trial remission was defined using the ACR criteria. The clinical and radiographic efficacy benefits reported here, including the proportion of patients achieving DAS28 (CRP)-defined remission, are comparable with those observed in other similar trials of early RA with biological therapies. This is particularly encouraging because of the poor prognostic status of the patients in this study, as all were required to have erosions and to be RF or anti-CCP2-seropositive at baseline. In fact, approximately 90% of the patients were anti-CCP2 positive. These prognostic factors all correlate with poor long-term outcomes and an aggressive disease course.

As a result of the chronic long-term nature of RA, the safety and tolerability of a therapeutic agent is of critical importance. In this study, the overall frequency of adverse events and serious adverse events (including serious infections and autoimmune events) was similar between treatment groups, and in some cases was numerically lower in the abatacept plus methotrexate group. Infusional reactions occurred more frequently in the abatacept plus methotrexate group and were mostly mild in severity. Across both groups there were no cases of tuberculosis or opportunistic infections. One case of cancer was reported across treatment arms (pancreatic cancer in the abatacept plus methotrexate group). These data are consistent with the safety findings in previous studies of abatacept-treated patients with RA of a longer disease duration. The safety and tolerability of abatacept plus methotrexate in this study was further supported by a high retention rate (>90%) and a low rate of discontinuations for safety reasons.

These data should be interpreted within the context of the trial. As these short-term, double-blind data are restricted to 1 year, the capacity to assess longer-term structural changes or detect infrequent safety-related events may be limited. It will be interesting to examine this group of poor prognosis patients over the second year of this 2-year trial in order to monitor whether the clinical, functional and radiographic benefits observed here are maintained or improved, as has been observed with abatacept in patients with established RA. As this study focused on a poor prognosis patient group, it will be interesting to examine this group of poor prognosis patients who received treatment with abatacept plus methotrexate in a two-year prospective study. The safety and tolerability profile comparable to methotrexate alone, these data support the early use of abatacept in RA patients with moderate to severe disease.

Acknowledgements: The authors would like to thank Helen Clarke, MSc, Medicus International, for her editorial assistance.

Funding: Editorial assistance was funded by Bristol-Myers Squibb, Princeton, NJ, USA.

Competing interests: RW has received consulting fees and speaker’s bureau fees from Bristol-Myers Squibb and Schering-Plough and research support from UCB; AD was a member of the national board as a consultant for Bristol-Myers Squibb (2003–7) and Abbott (2003–5); JW has received consulting fees from Bristol-Myers Squibb; JF is a member of advisory boards for Wyeth, Schering-Plough, Bristol-Myers Squibb and Roche and has received lecture fees from Abbott, Wyeth, Roche, Bristol-Myers Squibb and Schering-Plough; WG is a consultant for Bristol-Myers Squibb, Abbott Immunology, General Electric, Esata and Schering-Plough, has received honorarium from Bristol-Myers Squibb, Abbott Immunology, General Electric, Schering-Plough and Wyeth and has received research support from Abbott Immunology and Wyeth; BH is a consultant and has received a grant/research support for Abbott Canada, Argenz Canada, Bristol-Myers Squibb Scandinavia, Roche and Schering-Plough; WS is a speaker and principal investigator for Argenz, Wyeth, Abbott, Bristol-Myers Squibb, Centocor, Genentech and Biogen Idec; HG is a consultant for CCBR-SYNARC, Bristol-Myers Squibb, Wyeth, Roche, Servier, GlaxoSmithKline, Merck, Biogen Idec and Genentech and has stocks in CCBR-SYNARC; CP is an employee and has stocks in Synarc Inc; J-CB is an employee of BMS and owns stock options; AC and RH are employees of Bristol-Myers Squibb; JB is a consultant for Centocor and Riley, is on the advisory board for Abbott and Argenz, has received research support for Argenz, Biogen Idec and Bristol-Myers Squibb and has received honorarium for Abbott, Argenz, Centocor and Novartis. MR, SN, PD and SHP have nothing to disclose.

Ethics approval: The protocol and patients’ informed consent received institutional review board/independent ethics committee approval, and the study was conducted in accordance with the Declaration of Helsinki and was consistent with International Conference on Harmonisation good clinical practice.

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Ann Rheum Dis 2009 68: 1870-1877 originally published online January 5, 2009
doi: 10.1136/ard.2008.101121

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