Comparison of two etanercept regimens for treatment of psoriasis and psoriatic arthritis: PRESTA randomised double blind multicentre trial

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ABSTRACT

Objectives To compare the efficacy over 12 weeks of two different etanercept regimens in treating the skin manifestations of psoriasis in patients who also have psoriatic arthritis and to evaluate efficacy and safety over an additional 12 weeks of open label etanercept treatment.

Design Randomised double blind multicentre outpatient study.

Setting 98 outpatient facilities in Europe, Latin America, and the Asia Pacific region.

Participants 752 patients with both psoriasis (evaluated by dermatologists) and psoriatic arthritis (evaluated by rheumatologists).

Interventions During the blinded portion of the study, participants were randomised to receive etanercept 50 mg twice weekly (n=379) or 50 mg once weekly (n=373) for 12 weeks by subcutaneous injection. All participants then received open label etanercept 50 mg once weekly for 12 additional weeks, while remaining blinded to the regimen.

Main outcome measures The primary efficacy end point was the proportion of participants achieving “clear” or “almost clear” on the physician’s global assessment of psoriasis at week 12. Secondary efficacy analyses included psoriasis area and severity index, American College of Rheumatology responses, psoriatic arthritis response criteria, and improvement in joint and tendon disease manifestations.

Results At week 12, 46% (176/379) of participants receiving etanercept 50 mg twice weekly achieved a physician’s global assessment of psoriasis of “clear” or “almost clear” compared with 32% (119/373) in the group treated with 50 mg once weekly (P=0.001). In contrast, an equally high percentage of participants in both groups achieved psoriatic arthritis response criteria (77% (284/371) in the twice weekly/once weekly group versus 76% (282/371) in the once weekly/once weekly group). Participants treated with 50 mg twice weekly/once weekly had greater mean reductions from baseline in the psoriasis area and severity index at week 12 compared with those who received 50 mg once weekly/once weekly (71% vs 62%, P=0.001), with less difference at week 24 (78% v 74%, P=0.110). Joint and tendon disease manifestations improved from baseline in both groups to a similar extent. No new safety signals were seen in either etanercept treatment group, and no significant difference in the safety profiles was observed.

Conclusions In participants with active psoriasis and psoriatic arthritis, initial treatment of the psoriasis with etanercept 50 mg twice weekly may allow for more rapid clearance of skin lesions than with 50 mg once weekly. A regimen of 50 mg once weekly seems to be appropriate for treatment of joint and tendon rheumatic symptoms. The choice of regimen should be determined by the clinical needs of the individual patient.

Trial registration Clinical trials NCT00245960.

INTRODUCTION

The major manifestation of psoriasis is chronic inflammation of the skin characterised by scaling and erythematous plaques that may be painful or severely pruritic.1 Recommended treatment for the management of psoriasis includes topical treatments, ultraviolet light therapy, oral retinoids, methotrexate, ciclosporin, and biological agents.1 In many psoriasis patients, an inflammatory arthritis develops, with a distinct clinical picture. Psoriatic arthritis is distinguished by a chronic inflammation of joints and entheses, the point at which the collagen fibres of ligaments or tendons become mineralised and integrated into bone tissue.2 Enthesitis (the inflammation of entheses) and dactylitis or “sausage digit” (the uniform swelling of an entire digit) are frequent components of the clinical picture of psoriatic arthritis.2 The cutaneous symptoms of psoriatic arthritis usually appear a decade or more before the joint symptoms, enthesitis, and dactylitis.3 The disease affects men and women equally and has a worldwide distribution.4,5 Although incidence of psoriatic arthritis is less than 1% in the general population, the prevalence of psoriatic arthritis in patients with psoriasis is estimated to be as high as 30%.6,7 The goal of treating the arthritis component of psoriatic arthritis...
is to reduce inflammation related joint swelling and pain and to inhibit radiological progression, thereby preserving function and improving quality of life. Historically, treatment options for psoriatic arthritis have favoured non-steroidal anti-inflammatory drugs and disease modifying antirheumatic drugs. However, the literature to support the effectiveness of these agents is scant. Biological agents have changed the management of psoriatic arthritis by showing clinical as well as radiographic efficacy.

Analyses of skin, joint synovium, and synovial fluid from patients with psoriasis and psoriatic arthritis have indicated that T cells and cytokines, such as tumour necrosis factor alpha, may play an important role in this disease. Etanercept, a fully human tumour necrosis factor soluble receptor fusion protein that antagonises the effects of endogenous tumour necrosis factor, has been approved as a treatment option for patients with rheumatoid arthritis, juvenile idiopathic arthritis, and ankylosing spondylitis, as well as both moderate to severe plaque psoriasis and active psoriatic arthritis.

Etanercept is approved in the European Union for the treatment of psoriasis with either intermittent (50 mg twice weekly for 12 weeks, followed by 50 mg weekly) or continuous dosing (50 mg once weekly for 24 weeks) and has a favourable safety profile with no observed dose dependent toxic effects. In a double blind phase 3 trial, the exposure adjusted rates of adverse events and infections in patients treated with etanercept were similar to those for placebo in adults with psoriasis (n=618) for 96 weeks. Long term evaluation of the safety of etanercept in patients with psoriasis has found no signs of dose related or cumulative toxicity over time in registry data (up to 156 weeks). Moreover, etanercept treatment in patients with psoriatic arthritis has been found to be within the range of cost effectiveness estimates considered to represent value in the NHS by the National Institute for Health and Clinical Excellence. Patients with this combination of skin disease and arthritis present a management challenge, as they have two serious disease manifestations. However, similarities in their pathological processes present an opportunity to use a single treatment to effectively treat both components.

The aim of the PRESTA (Psoriasis Randomized Etanercept STudy in Subjects with Psoriatic Arthritis) trial was to determine the efficacy of two different etanercept regimens not previously studied in patients with both moderate to severe psoriasis and active psoriatic arthritis. In an effort to optimise patients’ care, PRESTA paired dermatologists and rheumatologists in a cooperative strategy to assess the impact of etanercept treatment on both skin and arthritic manifestations.

**METHODS**

Study population

Patients were eligible for this study if they were aged at least 18 with active but clinically stable plaque psoriasis involving at least 10% of the total body surface area and a physician’s global assessment of psoriasis of moderate to severe at screening and at baseline. Additionally, all participants were required to have active psoriatic arthritis, defined as at least two swollen joints, at least two tender or painful joints, joint pain (including axial) for at least three months before screening, and negative serum rheumatoid factor within six months before screening. In most cases, a rheumatologist did the rheumatologic assessments and diagnosed psoriatic arthritis; when this was not possible, joint evaluations were done by trained assessors. Female participants were required to have a negative pregnancy test at baseline, and all participants were required to use a medically acceptable form of contraception throughout the trial.

Patients were excluded if they had other active skin conditions that would interfere with study evaluations; a tender, swollen joint not assessed by a rheumatologist as psoriatic arthritis; severe comorbidities; recent

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**Table 1** Assessment tools

| Assessment tool                                      | Rating                                   | End point                                                |
|------------------------------------------------------|------------------------------------------|----------------------------------------------------------|
| Physician’s global assessment (PGA) of psoriasis     | 0 (clear, no lesions) to 5 (severe)      | PGA of psoriasis of “clear” (0) or “almost clear” (1)   |
| Psoriasis area and severity index (PASI)             | 0 (no lesions) to 72 (severe lesions on 100% of body) | PASI improvement ≥75% and ≥90% at weeks 12 and 24 |
| Psoriatic arthritis response criteria (PsARC)        | Improvement in 2/4 PsARC criteria; no criteria could worsen | Proportion of participants achieving PsARC at weeks 12 and 24 |
| PGA of arthritis                                     | 0 (no arthritis activity) to 100 (severe disease) | % improvement from baseline                              |
| American College of Rheumatology (ACR) response     | % reduction in tender and swollen joint counts plus 3/5 other parameters | ACR ≥20, ACR ≥50, and ACR ≥70 |
| Enthesitis at baseline                               | No of tendons showing enthesitis (0-4), based on Achilles tendons and plantar fasciae bilaterally | Proportion with improvement in ≥1 tendon/ligament insertion |
| Dactylitis at baseline                               | Rate each digit 0 to 3; total score for hands and feet 0 (none) to 60 (severe) | % change from baseline based on 60 point scale |

Fig 1 | Flow of participants through study. *Two patients were enrolled and had study drug dispensed but not administered; they were not included in safety or efficacy analyses. BIW=twice weekly; QW=once weekly
serious infection (within one month); or tuberculosis infection (appropriate screening and treatment of tuberculosis in the setting of anti-tumour necrosis factor treatment was based on guidelines of the local country). Prohibited treatments included all forms of ultraviolet light therapy, psoralen plus ultraviolet A radiation within 28 days before baseline, and ultraviolet B radiation within 14 days before baseline. Therapeutic sunbathing was prohibited from after the baseline visit to week 24 of the study. Participants were not to have received systemic psoriasis treatment, ciclosporin, or disease modifying antirheumatic drugs within 28 days before starting the study drug, with the exceptions of ≤20 mg/week of methotrexate or ≤50 mg/day of acitretin if the patient had been receiving a stable dose of either for at least eight weeks before starting the study drug. Changing the dose of either agent during the study was permitted only if required for the participant’s safety. Participants were not to have used topical vitamin A or vitamin D analogue preparations or anthralin within 14 days. Topical corticosteroids of low to moderate strength, and in stable doses and formulations, were permitted only for use on the scalp, axillae, or groin. Use of any tumour necrosis factor inhibitor, including etanercept, at any time before enrolment was not permitted. Participants were not to receive an injectable corticosteroid within 28 days before screening or during the study; however, oral corticosteroids (prednisone ≤10 mg/day or equivalent) for the inflammatory arthritis were permitted as long as the dose did not change within 28 days of baseline. Non-steroidal anti-inflammatory drugs were allowed if the dose remained stable from 14 days before baseline and throughout the study.

All elements of informed consent were explained to the participant’s safety. Participants were not to receive any systemic corticosteroids, biologics, or disease modifying antirheumatic drugs during the study. Non-steroidal anti-inflammatory drugs were allowed if the dose remained stable from 14 days before baseline and throughout the study.

Study design
Patients who had both moderate to severe plaque psoriasis and psoriatic arthritis were enrolled from 98 international sites into this randomised multicentre study. The study consisted of a 12 week double blind treatment period followed by a 12 week open label treatment period and a two week post-treatment follow-up.

We randomly assigned participants to one of two etanercept treatment regimens. In the double blind period, one group (n=379) received etanercept 50 mg administered subcutaneously twice weekly for 12 weeks and a second group (n=373) received etanercept 50 mg subcutaneously once weekly and matching placebo administered once weekly for 12 weeks. In the subsequent open label period, participants in both groups received etanercept 50 mg once weekly for 12 weeks; patients and investigators remained blinded to their treatment during the first period throughout the study. Participants who did not achieve improvement of at least one unit from baseline on the physician’s global assessment of psoriasis at week 12 were deemed treatment failures and were withdrawn from the study, unless the investigator determined that the treatment was providing improvement in joint symptoms.

The primary efficacy end point of the study was the proportion of participants who achieved “clear” or “almost clear” on the physician’s global assessment of psoriasis at week 12. This measure was reported on a scale of 0 to 5, with a rating of 0 indicating clear skin, 1 being almost clear, and 5 indicating severe skin symptoms. The physician’s global assessment of psoriasis is considered to be similar to the evaluation methods used in clinical practice, with comparable reliability and lower intra-rater variation than the psoriasis area and severity index.

Secondary end points included physician’s global assessment of psoriasis at week 24, as well as achievement of 75% and 90% improvement in psoriasis area and severity index, mean improvement in psoriasis

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Table 2 | Demographics and baseline clinical data. Values are numbers (percentages) unless stated otherwise

| Characteristics                        | Etanercept 50 mg BIW/QW (n=379) | Etanercept 50 mg QW/QW (n=373) |
|----------------------------------------|---------------------------------|---------------------------------|
| Mean (SD) age (years)                  | 46 (11)                         | 47 (11)                         |
| Male sex                               | 243 (64)                        | 230 (62)                        |
| White ethnicity                        | 333 (88)                        | 335 (90)                        |
| Mean (SD) body mass index (kg/m²)      | 28 (5)                          | 28 (6)                          |
| Mean (SD) duration of psoriasis (years)| 19 (12)                         | 19 (11)                         |
| Mean (SD) duration of psoriatic arthritis (years) | 7 (7) | 7 (7) |
| PGA-psoriasis                          | 3.6 (0.7)                       | 3.6 (0.7)                       |
| PASI                                    | 20 (11)                         | 19 (10)                         |
| Mean (SD) affected body surface (% area)| 31 (22)                        | 30 (22)                         |
| Mean (SD) No of swollen joints         | 12 (15)                         | 13 (15)                         |
| Mean (SD) No of tender joints          | 19 (18)                         | 19 (18)                         |
| Previous methotrexate use†             | 120 (32)                        | 150 (40)                        |
| Previous topical steroids‡             | 218 (58)                        | 181 (49)                        |
| Mean (SD) C reactive protein (mg/l)    | 15.3 (25.5)                     | 16.2 (27.7)                     |

BIW=twice weekly; PASI=psoriasis area and severity index; PGA=physician’s global assessment; QW=once weekly. *Within six months before screening. **Fisher exact test, two tailed. ***Fisher exact test, two tailed.
Table 3 | Soft tissue and articular manifestations. Values are numbers (percentages, 95% CI) unless stated otherwise

| Participants achieving ACR response | Etanercept 50 mg BIW/QW (n=379) | Etanercept 50 mg QW/QW (n=373) |
|--------------------------------------|---------------------------------|---------------------------------|
| ACR 20 week 12                        | 239/360 (66.4, 61.3 to 71.3)    | 219/360 (60.8, 55.6 to 65.9)    |
| ACR 20 week 24                        | 249/361 (69.0, 63.9 to 73.7)    | 258/360 (71.7, 66.7 to 76.3)    |
| ACR 50 week 12                        | 161/360 (44.7, 39.5 to 50.0)    | 146/360 (40.6, 35.4 to 45.8)    |
| ACR 50 week 24                        | 187/361 (51.8, 46.5 to 57.1)    | 193/360 (53.6, 48.3 to 58.9)    |
| ACR 70 week 12                        | 73/360 (20.3, 16.2 to 24.8)     | 79/360 (21.9, 17.8 to 26.6)     |
| ACR 70 week 24                        | 125/361 (34.6, 29.7 to 39.8)    | 132/360 (36.7, 31.7 to 41.9)    |

Participants achieving psoriatic arthritis response criteria

Week 12: 284/371 (76.0, 71.3 to 80.3) 282/371 (76.0, 71.3 to 80.3)
Week 24: 303/372 (81.5, 77.1 to 85.3) 299/372 (80.4, 76.0 to 84.3)

Enthesitis

Enthesitis at baseline 153 (40.4) 134 (35.9)
Improved week 12 109/148 (73.6, 67.9 to 78.8) 91/130 (70.0, 61.3 to 77.7)
Improved week 24 114/141 (80.0, 73.4 to 87.0) 100/123 (81.3, 73.3 to 87.8)

Dactylitis

Dactylitis at baseline 158 (41.7) 160 (42.9)
Mean score at baseline 7.93 8.16
Week 12:
Mean (SD) score 2.06 (5.51) 2.52 (7.69)
Mean % change from baseline 74.3 78.4
Week 24:
Mean (SD) score 1.42 (5.12) 1.80 (7.15)
Mean % change from baseline 84.5 84.8

ACR=American College of Rheumatology; BIW=twice weekly; QW=once weekly.

Continuous and ordinal end points by using analysis of covariance stratified by geographical region and using baseline as covariate or analysis of variance if the baseline value was not available.

The modified intention to treat population included all randomised participants who took at least one dose of the test drug and had at least one post-baseline efficacy evaluation. We did efficacy and safety analyses on the modified intention to treat population. Two additional patients were enrolled and had study drug dispensed, but the drug was not administered, and they were not included in the safety or efficacy analyses. Efficacy analyses used the last observation carried forward method for imputation of missing data. We used the data analysis software UNIX SAS version 9.1.3 for statistical analyses.

RESULTS

Participants’ characteristics

We randomised 754 participants; 752 participants (379 in the etanercept 50 mg twice weekly/once weekly group and 373 in the etanercept 50 once weekly/once weekly group) comprised the modified intention to treat population and 92% (695) completed the study (fig 1). Baseline demographic and disease characteristics were balanced between treatment groups (table 2). Participants had a mean age of 46.5 years. Most participants were men (473/752; 63%), and most were white (668/752; 89%). The mean duration of psoriasis was 18.9 years, and the mean duration of psoriatic arthritis was 7.0 years. In general, the extent and severity of arthritis and psoriatic symptoms were similar across treatment groups. Rheumatologists diagnosed the psoriatic arthritis and did the rheumatic assessments 92% of the time; when this was not possible, joint evaluations were done by trained assessors (6%) or by dermatologists (2%). The mean doses of etanercept over 24 weeks were 74.6 (SD 11.4) mg in the twice weekly/once weekly group and 50.0 (4.7) mg in the once weekly/once weekly group. Mean concentrations of C reactive protein were high at baseline in both groups (15.3 (SD 25.5) mg/l in the twice weekly/once weekly group and 50.0 (4.7) mg/l in the once weekly/once weekly group. No statistically significant differences existed...
between groups in the proportions of participants receiving concomitant treatment for psoriasis.

**Efficacy**

**Skin**

A significantly greater proportion of participants in the twice weekly/once weekly group (46%; 176/379) achieved a status of “clear” or “almost clear” for physician’s global assessment of psoriasis at week 12 compared with those in the once weekly/once weekly group (32%; 119/373) (P<0.001) (fig 2). By week 24, the proportions were similar (56% (214/379) v 50% (187/373), P=0.104). The mean percentage improvement from baseline in the physician’s global assessment of psoriasis at week 12 was significantly greater in the twice weekly/once weekly group than in the once weekly/once weekly group (52% v 43%, P<0.001). At week 24, the mean percentage improvement from baseline in physician’s global assessment of psoriasis was similar for both groups (57% v 53%, P=0.420) (fig 3).

At week 12, the mean improvement from baseline in the psoriasis area and severity index was significantly greater in the twice weekly/once weekly group than in the once weekly/once weekly group (71% v 62%, P<0.001) (fig 4); however, at week 24, the change from baseline was similar in the two groups (78% v 74%, P=0.110). A significantly greater proportion of participants in the etanercept 50 mg twice weekly/once weekly group than in the 50 mg once weekly/once weekly group achieved at least 75% improvement in the psoriasis area and severity index (55% (207/377) v 36% (135/371) at week 12, P<0.001; 70% (265/377) v 62% (231/371) at week 24, P<0.026). The within group changes from baseline in physician’s global assessment of psoriasis and psoriasis area and severity index were statistically significant at all study visits in both etanercept groups (P<0.001 for each).

**Joint and tendon rheumatic manifestations**

The proportions of participants who achieved American College of Rheumatology (ACR) 20, 50, and 70 responses were similar in the two groups at weeks 12 and 24. At week 24, 69% of the twice weekly/once weekly group and 72% of the once weekly/once weekly group had ACR 20 responses (P=0.379), 52% and 54% achieved ACR 50 responses (P=0.594), and 35% and 37% achieved ACR 70 responses (P=0.530) (table 3, fig 5). Six participants who failed to meet the inclusion criteria for active psoriatic arthritis of at least two painful and two swollen joints at baseline were excluded from this analysis. The proportions of participants who achieved psoriatic arthritis response criteria were similar in the two groups at week 12 and remained stable at week 24. At week 12, 77% in the twice weekly/once weekly group and 76% in the once weekly/once weekly group achieved psoriatic arthritis response criteria, as did 82% and 80% at week 24 (table 3). The percentage improvements from baseline in physician’s global assessment of arthritis were similar in the two groups at weeks 12 and 24. At week 12, 60% for the twice weekly/once weekly group and 62% for the once weekly/once weekly group achieved psoriatic arthritis response criteria, as did 82% and 80% at week 24 (table 3). The percentage improvements from baseline in physician’s global assessment of arthritis were similar in the two groups at weeks 12 and 24. At week 24, the corresponding reductions were 73% and 74% (P=0.760).

At baseline, enthesitis was present in 287 participants and dactylytis was present in 318 participants (table 3). Participants with enthesitis at baseline had more extensive skin and joint involvement and higher C reactive protein concentrations than did those who did not present with enthesitis at baseline (table 4). The

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**Table 4** Disease characteristics in participants with and without enthesitis at baseline. Values are mean (SD) score; percentage improvement

| Enthesitis at baseline | No enthesitis at baseline | Enthesitis at baseline | No enthesitis at baseline |
|-----------------------|--------------------------|-----------------------|--------------------------|
| Physician’s global assessment of psoriasis | | | |
| Baseline | 57.56** (20.19) | 45.85 (19.89) | 55.13** (20.86) | 46.95 (20.31) |
| Week 12 | 23.03* (21.09) | 15.20 (16.87) | 22.89 (21.15) | 15.86 (15.82) |
| Week 24 | 14.80 (19.16) | 10.48 (16.77) | 15.58* (17.72) | 11.04 (13.24) |
| Painful joints | | | |
| Baseline | 28.34** (21.05) | 12.75 (11.54) | 28.43** (21.40) | 14.03 (12.33) |
| Week 12 | 13.33 (18.48) | 5.06 (7.23) | 13.70* (18.03) | 5.29 (7.42) |
| Week 24 | 9.12 (16.24) | 3.79 (6.77) | 8.71 (15.20) | 6.92 (6.82) |
| Swollen joints | | | |
| Baseline | 18.17** (19.92) | 7.73 (7.55) | 19.48** (20.35) | 9.03 (8.97) |
| Week 12 | 7.38 (14.70) | 2.12 (3.55) | 7.85 (14.79) | 5.97 (3.93) |
| Week 24 | 4.77 (12.69) | 1.35 (3.25) | 4.42 (10.49) | 1.68 (3.36) |
| C reactive protein | | | |
| Baseline | 16.17 (26.16) | 14.88 (24.97) | 17.34 (22.79) | 15.52 (29.76) |
| Week 12 | 5.29 (7.12) | 4.66 (2.70) | 5.98 (7.43) | 5.88 (8.55) |
| Week 24 | 5.13 (4.77) | 5.76 (11.62) | 5.76 (6.67) | 5.60 (5.04) |

*P<0.05, enthesitis versus no enthesitis within treatment arm.
**P<0.01, enthesitis versus no enthesitis within treatment arm.
number of sites of enthesitis, determined by manual pressure on the tendon insertion, decreased from baseline in both treatment groups (table 3). Similarly, among participants with dactylitis at baseline, comparable decreases occurred in the mean number of toes and fingers with objective dactylitis in the two etanercept groups at weeks 12 and 24 (table 3).

Concomitant treatment
Only 25% of participants in this trial received concomitant methotrexate treatment; the mean dosage was 12.7 (SD 4.3) mg/week. In this subset of participants, some benefit of combination therapy was apparent at week 12 for skin but not joint symptoms and only in those who received etanercept 50 mg twice weekly during this period.

C reactive protein
Mean concentrations of C reactive protein were significantly decreased from baseline to a similar extent in both groups. Concentrations decreased from 15.3 (SD 25.5) mg/l at baseline to 5.5 (9.5) mg/l by week 24 in the 50 mg twice weekly/once weekly group and from 16.2 (27.7) mg/l to 5.7 (4.9) mg/l in the once weekly/once weekly group. Interestingly, participants who presented with enthesitis at baseline had higher C reactive protein concentrations than did those who did not have enthesitis at baseline. Baseline C reactive protein concentrations were 16.2 (26.2) mg/l in the 50 mg twice weekly/once weekly group and 17.3 (22.8) mg/l in the once weekly/once weekly group among participants with enthesitis at baseline. In participants who did not present with enthesitis, the baseline C reactive protein concentrations were 14.9 (25.0) mg/l and 15.5 (29.78) mg/l in the two groups (table 4).

Safety
Etanercept was well tolerated in both treatment groups over 24 weeks; we found no significant differences between the groups in the incidence of adverse events. The most commonly reported treatment emergent adverse events were upper respiratory tract infection, injection site reaction, pharyngitis, and headache. A total of 15 (4%) participants in the twice weekly/once weekly group and 11 (3%) in the once weekly/once weekly group reported serious adverse events, including serious infections. Five (0.7%) serious infections were reported: two (0.3%) in the twice weekly/once weekly group and three (0.8%) in the once weekly/once weekly group (table 5). No cases of tuberculosis, other opportunistic infections, or demyelinating disorders were reported. Four malignancies were reported: two skin carcinomas and one breast carcinoma in the twice weekly/once weekly group and one skin carcinoma in the once weekly/once weekly group. No participant died during the study.

DISCUSSION
In participants with active psoriasis and psoriatic arthritis, we found that initial treatment of the psoriasis with etanercept 50 mg twice weekly may allow for more rapid clearance of skin lesions than a 50 mg weekly regimen. The recommended dose regimens of etanercept for psoriasis and psoriatic arthritis are different. In the European Union, the summary of product characteristics recommends that psoriasis can be treated with either 50 mg weekly or 50 mg twice weekly for 12 weeks followed by 50 mg weekly, whereas the recommended etanercept dose for psoriatic arthritis is 50 mg weekly. The greater effect of the twice weekly/once weekly regimen on skin manifestations at 12 weeks seen in this study was similar to what has been found in the treatment of psoriasis in the absence of arthritis. The results also suggest that the skin manifestations may benefit from 50 mg twice weekly initially and that more than 12 weeks of treatment may be needed to achieve a maximal response— in this case 24 weeks.

The effect on skin was greater than that seen in the first study of etanercept in patients with both psoriasis and psoriatic arthritis and may reflect the lower baseline severity of psoriasis in that trial. In that study, the analysis of skin improvement was done only for patients with 3% or more body surface area involvement compared with the far more severe involvement seen in the PRESTA trial, in which the mean body surface area involved was more than 30%. Demonstrating a 75% improvement in the psoriasis area and severity index may be more difficult in patients with less severe disease.
WHAT IS ALREADY KNOWN ON THIS TOPIC

Dermatologists and other practitioners treating patients with plaque psoriasis are in an ideal position to screen for psoriatic arthritis and provide therapeutic management or referral.

Etanercept is approved for treatment of moderate to severe plaque psoriasis and active psoriatic arthritis on the basis of its efficacy in treating both skin and joint symptoms.

WHAT THIS STUDY ADDS

For patients with plaque psoriasis and psoriatic arthritis, etanercept 50 mg twice weekly was superior to 50 mg once weekly for skin manifestations at week 12 but similar for joint manifestations.

Both regimens achieved significant improvement from baseline in skin, joint, and enthesal disease components at week 24 without notable differences in safety.

Either etanercept dose regimen can be used in the treatment of psoriasis with or without the presence of psoriatic arthritis, allowing for individualized care.

Recent studies using similar dosing regimens in psoriasis alone support a greater and faster response with the use of etanercept 50 mg twice weekly during the first 12 weeks. The results of the PRESTA trial suggest that 50 mg twice weekly followed by 50 mg weekly is an appropriate dose regimen for treating skin symptoms in these patients, whether or not they have concomitant psoriatic arthritis. The 50 mg twice/once weekly regimen allowed for a faster cutaneous response and may be preferable in patients with more severe skin disease. On the other hand, at no time point was the twice weekly/once weekly regimen more advantageous in treating joint or tendon symptoms than the 50 mg once weekly dose regimen that is approved for psoriatic arthritis. The challenge of treating patients with both active psoriasis and active psoriatic arthritis is to optimize the treatment of both disease manifestations to give the best overall outcome.

The noteworthy improvements in psoriatic arthritis response criteria and the high percentage of participants who achieved American College of Rheumatology 20/50/70 responses were similar to the results of the original registration study using a 25 mg twice weekly regimen, suggesting that a 50 mg once weekly dose is comparable in efficacy to 25 mg twice weekly. Dissociation clearly existed with regard to the optimal dosages for the skin lesions at week 12. However, when the dosage was decreased in the second 12 weeks of the trial, both skin and joint symptoms continued to improve, and the 50 mg once weekly/once weekly group achieved responses similar to those of the 50 mg twice weekly/once weekly group at week 24.

Enthesitis and dactylitis, which are clinically important components of psoriatic arthritis, improved equally well on both etanercept regimens. This study confirms previous data suggesting that patients with enthesitis have more severe disease than those without these extra-articular problems. The response to etanercept in this study is consistent with the results of a previous study that evaluated the effect of infliximab on enthesitis and dactylitis in patients with psoriatic arthritis. This is the first definitive demonstration that etanercept significantly improves both of these important extra-articular disease manifestations of psoriatic arthritis, even at the lower doses commonly used to treat arthritis alone, compared with the infliximab studies in psoriatic arthritis which used the higher 5 mg/kg dosage.

Under the conditions of this study, the higher dose of etanercept improved skin manifestations more rapidly than did the lower dose but did not seem to provide an additional advantage in treating joint or enthesal symptoms. The explanation for this differential effect on skin and joints is unclear. The ideal dosing for psoriatic arthritis is apparently more similar to the regimen used in rheumatoid arthritis than to that used in psoriasis. These two different organ systems may have dissimilar autoimmune inflammatory environments, allowing for differences in local concentrations of tumour necrosis factor or in disease burdens or a subtle difference in tissue penetration of drug, although little information is available to support any particular mechanism.

Although trials of anti-tumour necrosis factor agents have been done in psoriatic arthritis, PRESTA is unique in its collaboration between dermatologists and rheumatologists for the evaluation of both skin and joint symptoms in this complex population. The advantage of the cooperative strategy between specialists in this trial can be supported by the consistent measurement of outcomes for both psoriasis and psoriatic arthritis compared with previous disease specific trials.

Conclusion

The results of this study indicate that, although significant differences in skin responses were seen at week 12 between the 50 mg twice weekly/once weekly and 50 mg once weekly/once weekly dosages, 50 mg weekly is a sufficient dose for treatment of joint symptoms alone. Both regimens achieved significant improvement from baseline in skin, joint, and enthesal disease components at week 24. Furthermore, these improvements were achieved without any notable differences in safety.

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Competing interests: WS has received fees for speaking/consulting from Abbott, Schering-Plough, Wyeth, and Janssen-Ciaglia. J-PO has received fees for speaking/conferences/consulting from Schering-Plough, Abbott, Merck-Serono, Centocor, Wyeth, Janssen-Ciaglia, MedPharma, Laboratorios Pierre-Fabre, Galderma Laboratorios, and Leo Pharma. BK has served on advisory boards for Schering-Plough and Roche; has received funds for research/travel/conferences from Wyeth, Centocor, Abbott, Schering-Plough, Roche, and Bristol-Myers Squibb; and has served on a speaker panel for Bristol-Myers Squibb. OB has received fees from Wyeth, Schering-Plough, Abbott, Roche, Chugai, and Bristol-Myers Squibb. DR, RD, JE, CM, and BF are all employees of Pfizer.

Ethical approval: The protocol and its amendments received independent ethics committee or institutional review board approval, and regulatory review and approval before site initiation and recruitment of patients. All participants signed and dated an approved informed consent form.

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1. Menter A, Gottlieb A, Feldman SR, Van Voorhees AS, Leonardi CL, Gordon KB, et al. Guidelines of care for the management of psoriasis and psoriatic arthritis: section 1. Overview of psoriasis and guidelines of care for the treatment of psoriasis with biologics. J Am Acad Dermatol 2008;59:826-50.

2. Gottlieb A, Korman NJ, Gordon KB, Feldman SR, Lebwohl M, Koo JYM, et al. Guidelines of care for the management of psoriasis and psoriatic arthritis: section 2. Psoriatic arthritis: overview and guidelines of care for treatment with an emphasis on the biologics. J Am Acad Dermatol 2008;58:851-64.

3. Gladman DD. Psoriatic arthritis. Dermatol Ther 2004;17:390-63.

4. Feuchtenberger M, Kleinef H, Tony HP, Kerlitz C. Psoriatic arthritis: therapeutic principles. Clin Dermatol 2008;26:460-3.

5. Saad AA, Symmons DPM, Noyce PR, Ashcroft DM. Risks and benefits of tumor necrosis factor-alpha inhibitors in the management of psoriatic arthritis: systematic review and metaanalysis of randomized controlled trials. J Rheumatol 2008;35:883-90.

6. Gottlieb AB, Chao C, Dann F. Psoriasis comorbidities. J Dermatol Treat 2008;19:5-21.

7. Mease P. Management of psoriatic arthritis: the therapeutic interface between rheumatology and dermatology. Curr Rheumatol Rep 2008;10:349-54.

8. Ritchlin CT, Kavanaugh A, Gladman DD, Mease PJ, Hellwell P, Boehnecke WH, et al. Treatment recommendations for psoriatic arthritis. Ann Rheum Dis 2008;67:1387-94.

9. Amgen Inc and Wyeth Pharmaceuticals. Enbrel® (etanercept) prescribing information. Amgen Inc and Wyeth Pharmaceuticals, 2008.

10. Mease PJ, Goffe BS, Metz J, Van der Steen F, Burge DJ. Etanercept in the treatment of psoriatic arthritis: a randomized trial. Lancet 2000;356:385-90.

11. Tyring S, Gordon KB, Poulin Y, Langley RG, Gottlieb AB, Dunn M, et al. Long-term safety and efficacy of 50 mg of etanercept twice weekly in patients with psoriasis. Arch Dermatol 2007;143:719-26.

12. Driessen RJ, Bierma van de Kerf PCM, de Jong EMGJ. Three-year registry data on biological treatment for psoriasis: the influence of patient characteristics on treatment outcome. Br J Dermatol 2009;160:760-75.

13. Bravo Vergel Y, Hawkins NS, Claxton K, Asseburg C, Palmer S, Woolacott N, et al. The cost-effectiveness of etanercept and infliximab for the treatment of patients with psoriatic arthritis. Rheumatology 2007;46:1729-35.

14. Persikowski T, Petterson U. Severe psoriasis—oral therapy with a new retinoid. Dermatologica 1978;157:239-44.

15. Feldman SR, Krueger GG. Psoriasis assessment tools in clinical trials. Ann Rheum Dis 2005;64(suppl 2):S65-8.

16. Langley RG, Ellis CN. Evaluating psoriasis with psoriasis area and severity index, psoriasis global assessment, and lattice system physician’s global assessment. J Am Acad Dermatol 2004;51:563-9.

17. Ashcroft DM, Wan Po AL, Williams HC, Griffiths CM. Clinical measures of disease severity and outcome in psoriasis: a critical appraisal of their quality. Br J Dermatol 1999;141:185-91.

18. Finlay AY. Current severe psoriasis and the rule of tens. Br J Dermatol 2005;152:861-7.

19. Clegg DO, Reda DJ, Mejias E, Cannon GW, Weisman MH, Taylor T, et al. Comparison of sulfasalazine and placebo in the treatment of psoriatic arthritis. Arthritis Rheum 2002;46:328-46.

20. Van de Kerkhof PC, Segaat S, Lahfa M, Lutter TA, Karolyi Z, Kaszuba A, et al. Once-weekly administration of etanercept 50 mg is efficacious and well tolerated in patients with moderate-to-severe plaque psoriasis: a randomized controlled trial with open-label extension. Br J Dermatol 2008;159:1177-85.

21. Ortonne J-P, Griffiths C, Dauwe M, Lugert TA, Karolyi Z, Fazekas K, and J-PO, BK, and OB recruited and enrolled participants, collected data, and drafted the report. DR and JE designed the trial, wrote protocols for the participating institutions, interpreted the data, and drafted the report. CM interpreted the data, served as medical monitor during the trial, and drafted the report. All authors edited the final version of the manuscript. WS is the guarantor.

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