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**Title:** Treat to target in rheumatoid arthritis: opportunities and outcomes  
**Issue Date:** 2013-09-24
Chapter 5

The association of treatment response and joint damage with ACPA status in recent onset RA: a subanalysis of the 8-year follow-up of the BeSt study

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Annals of the Rheumatic Diseases 2012 Feb;71(2):245-8
ABSTRACT

Objective Anti-citrullinated protein antibodies (ACPA) are suggested to identify different subsets of patients with rheumatoid arthritis (RA). The authors compared the clinical and radiological response to Disease Activity Score (DAS)-steered treatment in patients with RA positive or RA negative for ACPA.

Methods In the Behandel Strategieën (BeSt) study, 508 patients with recent onset RA were randomized to four treatment strategies aimed at a DAS ≤2.4. Risks of damage progression and (drug-free) remission in 8 years were compared for ACPA-positive and ACPA-negative patients, using logistic regression analysis. Functional ability and DAS components over time were compared using linear mixed models.

Results DAS reduction was achieved similarly in ACPA-positive and ACPA-negative patients in all treatment strategy groups, with a similar need to adjust treatment because of inadequate response. Functional ability and remission rates were not different for ACPA-positive and ACPA-negative patients. ACPA-positive patients had more radiological damage progression, especially after initial monotherapy. They had a lower chance of achieving (persistent) drug-free remission.

Conclusion Clinical response to treatment was similar in ACPA-positive and ACPA-negative patients. However, more ACPA-positive patients, especially those treated with initial monotherapy, had significant radiological damage progression, indicating that methotrexate monotherapy and DAS ≤2.4 steered treatment might be insufficient to adequately suppress joint damage progression in these patients.
INTRODUCTION

Anti-citrullinated protein antibodies (ACPA) are highly specific antibodies for rheumatoid arthritis. Patients positive for ACPA have been shown to have higher disease activity, worse functional ability and more joint damage in observational and/or non-disease activity-steered studies. ACPA-positivity was found to be predictive of not achieving remission. ACPA-negative and ACPA-positive RA may be different diseases with different risk factors and clinical course and may require different therapeutic strategies. Possibly ACPA-positive and ACPA-negative patients also respond differently in a tight control treatment strategy where medication is adjusted based on the aim of achieving low disease activity. Therefore, we compared the changes in Disease Activity Score (DAS), functional ability and radiological damage over time in ACPA-positive and ACPA-negative patients with early RA treated according to the same disease activity steered protocol.

METHODS

Patients

Eight-year follow-up data of all 484 patients with known ACPA-status included in the BeSt (Dutch acronym for Behandel Strategieën, “treatment strategies”) study were analyzed. This is a multi-center randomized trial designed to compare four treatment strategies in 508 patients with recent-onset RA; initial monotherapy, step-up combination therapy (both starting with methotrexate monotherapy for ≥6 months), initial combination therapy with methotrexate, sulfasalazine and prednisolone and initial combination therapy with methotrexate and infliximab. Treatment was assessed every 3 months and adjusted if the DAS was >2.4. If the DAS was ≤2.4 for ≥6 months, medication was tapered to monotherapy in maintenance dose. Starting 2 years after inclusion, patients on monotherapy maintenance dose, who were in remission (DAS <1.6) for ≥6 months, stopped the last disease modifying anti-rheumatic drug (DMARD). Treatment was restarted if the DAS increased to ≥1.6. A more detailed description of the study protocol was published previously.

Study endpoints

ACPA-status was determined with the CCP2 test using baseline sera (n=119) and sera collected during the first years of follow-up (n=365). The DAS and Health Assessment Questionnaire (HAQ) were used to assess treatment response. Drug-free remission was defined as a DAS <1.6 and not using any DMARD. All available radiographs of hands and feet at year 0-1-2-3-4-5-6-7-8 were scored using the Sharp-van der Heijde score.
(SHS) by two independent readers, blinded for patient identity and time order (inter-
observer intraclass correlation coefficient 0.96), to assess joint damage. For DAS and DAS
components, areas under the curve (AUC) were calculated, only for years with complete
data. For years with ≤2 missing values, the last observation carried forward was used to
calculate the AUC, to avoid exclusion of these data.

Statistical analysis
Baseline characteristics and clinical parameters were compared using the χ² test, Student’s t test or Mann-Whitney U test. HAQ and DAS components over time were com-
pared using linear mixed models with ACPA-status and time as categorical variables and
HAQ or DAS component respectively at baseline, adjusted for baseline gender, smoking
habits, age and SHS with a Toeplitz covariance structure. Spearman’s correlation coeffi-
cient test was used to analyze the correlations after 8 years. ORs for achieving (drug-free)
remission, of restarting medication and of joint damage progression were calculated for
ACPA-positive patients using logistic regression analyses, adjusted for gender, smoking
habits, baseline age, DAS and SHS. ORs were converted to RRs to find a more accurate
estimation of the effect size.13
To examine the influence of treatment strategy, we used generalized estimating
equations with an auto-regressive covariance structure, time as categorical variable,
baseline SHS, DAS, age, gender, smoking habits, ACPA-status, treatment strategy
and ACPA*treatment strategy with yearly damage progression as outcome. To assess
the possible difference in the association between disease activity and joint damage
progression for ACPA-positive and -negative patients, we used generalized estimating
equations with these components but with treatment strategy replaced by yearly AUC
DAS or AUC DAS component (with baseline DAS component instead of baseline DAS).

RESULTS
Treatment response
ACPA-positive patients had a lower baseline DAS and HAQ and a higher SHS. Disease
activity over time was similar in both ACPA-groups.(figure 1a) Functional ability was not
different for ACPA-positive and ACPA-negative patients (p=0.9).(figure 1b) This similar
treatment response in both ACPA-groups was seen both in patients initially treated with
methotrexate monotherapy and with combination therapy (p=0.8 and p=0.9).(figure S1)
ACPA-positive patients did have a significantly higher (4.5mm/hr) erythrocyte sedimen-
tation rate (ESR).(figure 1c) Disease activity and functional ability showed a moderate
correlation after 8 years: rs:0.5 (p<0.001). The rates of achieving remission at least once
or of ≥1 year consecutively were not different: RR of 1.0 (95% CI 0.9;1.1) and 0.9 (95% CI
0.7;1.1), respectively. ACPA-positive patients were less likely to achieve drug-free remis-

sion, with a RR of 0.4 (95% CI 0.3;0.7) and more likely to lose remission and having to
restart DMARD: RR 2.3 (95% CI 1.4;3.0). Similar results were seen for patients who were
both ACPA and RF positive or negative.
The median number of treatment steps (2 (IQR 1-4) vs 1 (IQR 1-4)) that patients had
failed on and the proportions of patients who had dropped out before year 8 were not
significantly different for ACPA-positive and ACPA-negative patients in the whole cohort,
or when stratified for initial treatment strategy.

Figure 1: DAS (a), HAQ (b), erythrocyte sedimentation rate (ESR) (c), patient visual analogue scale global
health (VAS) (d), Ritchie Articular Index (e) and Swollen Joint count (f) over 8 years for anti-citrullinated
protein antibody (ACPA)-positive and ACPA-negative patients
ACPA-positive patients showed more radiological damage progression than ACPA-negative patients. (figure 2a) The RR for progression >5 points (SHS) was 3.8 (95% CI 2.5;5.0), 3.7 (95% CI 1.9;6.3) for >15 points, 3.2 (95% CI 1.4;6.4) for >25 and 6.2 (95% CI 1.5;20.3) for >35 points. Similar results were seen for patients who had been in remission for ≥1 year (figure S2) and for patients who were both ACPA and RF positive or negative. ACPA was a predictor of joint damage progression independent of RF.

Figure 2: Probability plots of joint damage progression over 8 years for anti-citrullinated protein antibody (ACPA)-positive and ACPA-negative patients, (A) all patients, (B) initial treatment monotherapy (groups 1 and 2), (C) initial combination treatment (groups 3 and 4)
Figure S1: DAS and HAQ over 8 years for anti-citrullinated protein antibody (ACPA)-positive and ACPA-negative patients.

Figure S2: Probability plot of joint damage progression over 8 years for anti-citrullinated protein antibody (ACPA)-positive and ACPA-negative patients who have been in remission for ≥1 year consecutively.

Figure S1: DAS and HAQ over 8 years for anti-citrullinated protein antibody (ACPA)-positive and ACPA-negative patients.
The association of ACPA-status with joint damage progression was significantly influenced by initial treatment strategy (monotherapy or combination treatment). Figures 2b, 2c) The difference in SHS between ACPA-positive and -negative patients initially treated with combination therapy was 1.7 points smaller than the difference in SHS for ACPA-positive and -negative patients initially treated with monotherapy (p<0.001). Second, the association was influenced by disease activity. ACPA-positive patients showed 1.8 points more increase in SHS per point of the DAS (p=0.001), 0.1 points per mm/hr ESR (p=0.003), 0.2 per tender joint (p=0.02) and 0.1 per swollen joint (p=0.005). The association with VAS global health was not influenced by ACPA-status. Joint damage and functional ability at year 8 did not show a significant correlation.

DISCUSSION

Response to DAS-targeted treatment was similar in ACPA-positive and ACPA-negative patients in terms of reduction of disease activity including remission percentages, and improvement of functional ability, although ACPA-positive patients had a higher ESR over time. ACPA-positive patients did show more joint damage progression, in particular in patients treated with initial methotrexate monotherapy. ACPA-positivity also was a predictor for not achieving and for losing drug-free remission.

To our knowledge, we are the first to report on disease activity in ACPA-positive and ACPA-negative patients in a disease activity steered treated cohort. In previous non-disease activity steered studies of patients with similar disease duration, ACPA-positive patients did show higher disease activity. In a study of 273 patients with recent-onset RA with 6 years of follow-up, similar functional ability was found for ACPA-positive and ACPA-negative patients after correction for disease activity and RF, but ACPA-positive patients had more joint damage, which is in line with our results. The relatively short follow-up period may account for these findings, as radiological joint damage shows a weak correlation with functional ability in the first years after the diagnosis of RA, but a moderate correlation after 12 years, while disease activity shows a stable, moderate correlation with functional ability from baseline onwards. In our tight controlled cohort we found a moderate correlation between functional ability and disease activity but no significant correlation with radiological joint damage after 8 years. Longer follow-up will show whether radiological joint damage will significantly contribute to functional disability with longer disease duration.

Our observation that ACPA-positivity is a predictor for not achieving drug-free remission and for relapsing if drug-free remission was achieved, is an extension on similar results after 5 years of treatment. The results are also in line with the findings of Balsa et al., who found that ACPA-positivity was a predictor for not achieving drug-free remission for
≥5 years, and of van der Woude et al. who found that ACPA-positivity was a predictor for not achieving drug-free remission for ≥1 year. It might be wise to take ACPA-status into consideration when contemplating cessation of medication.

In conclusion, DAS-targeted therapy is equally effective in reducing disease activity, achieving remission and improving functional ability in ACPA-positive and ACPA-negative patients with recent-onset RA. Still, ACPA-positive patients had more radiological damage, especially patients initially treated with methotrexate monotherapy. This suggests that in ACPA-positive patients, initial methotrexate monotherapy is insufficient to suppress joint damage progression even if subsequent treatment is DAS-targeted. This is in line with our previous findings and the European League against Rheumatism recommendations, which suggest that in patients with poor prognostic factors such as ACPA-positivity, starting with combination therapy might be considered. It may also mean that for ACPA-positive patients, the target of DAS ≤2.4 might not be stringent enough. The differences in joint damage progression and systemic inflammation indicate that the inflammatory mechanisms in ACPA-positive and ACPA-negative RA might have different mediators.

Table 1: baseline characteristics for ACPA-negative and ACPA-positive patients and drop-out at year 8

<table>
<thead>
<tr>
<th></th>
<th>ACPA – N=184</th>
<th>ACPA + N=300</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male gender (%)</td>
<td>48 (26)</td>
<td>111 (37)</td>
<td>0.013</td>
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<tr>
<td>Age (mean, SD)</td>
<td>55 (15)</td>
<td>54 (13)</td>
<td>0.5</td>
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<tr>
<td>Smoker (%)</td>
<td>51 (28)</td>
<td>117 (39)</td>
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<tr>
<td>RF pos (%)</td>
<td>59 (32)</td>
<td>258 (86)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Treatment strategy (%)</td>
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<tr>
<td>Sequential monotherapy</td>
<td>40 (22)</td>
<td>80 (27)</td>
<td></td>
</tr>
<tr>
<td>Step-up combination therapy</td>
<td>45 (25)</td>
<td>69 (23)</td>
<td></td>
</tr>
<tr>
<td>Initial combination therapy with pred</td>
<td>56 (30)</td>
<td>68 (23)</td>
<td></td>
</tr>
<tr>
<td>Initial combination therapy with IFX</td>
<td>43 (23)</td>
<td>83 (28)</td>
<td></td>
</tr>
<tr>
<td>Symptom duration, wks (median, IQR)</td>
<td>22 (13-41)</td>
<td>25 (14-56)</td>
<td>0.06</td>
</tr>
<tr>
<td>DAS (mean, SD)</td>
<td>4.6 (0.9)</td>
<td>4.3 (0.8)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HAQ (mean, SD)</td>
<td>1.5 (0.7)</td>
<td>1.3 (0.7)</td>
<td>0.02</td>
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<tr>
<td>SHS (median, IQR)</td>
<td>1.5 (0.0-6.1)</td>
<td>4.0 (1.0-10.5)</td>
<td>&lt;0.001</td>
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<tr>
<td>Number of treatment steps failed on before year 8</td>
<td>1 (1-4)</td>
<td>2 (1-4)</td>
<td>0.2</td>
</tr>
<tr>
<td>Drop-out at year 8</td>
<td>54 (29)</td>
<td>84 (28)</td>
<td>0.7</td>
</tr>
</tbody>
</table>

ACPA – Anti-Citrullinated Protein Antibodies RF – Rheumatoid Factor DAS – Disease Activity Score HAQ – Health Assessment Questionnaire SHS – Sharp-van der Heijde Score
REFERENCE LIST


