EXTENDED REPORT

Methotrexate in combination with other DMARDs is not superior to methotrexate alone for remission induction with moderate-to-high-dose glucocorticoid bridging in early rheumatoid arthritis after 16 weeks of treatment: the CareRA trial

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ABSTRACT

Objectives To compare the efficacy and safety of intensive combination strategies with glucocorticoids (GCs) in the first 16 weeks (W) of early rheumatoid arthritis (eRA) treatment, focusing on high-risk patients, in the Care in early RA trial.

Methods 400 disease-modifying antirheumatic drugs (DMARD)-naive patients with eRA were recruited and stratified into high risk or low risk according to classical prognostic markers. High-risk patients (n=290) were randomised to 1/3 treatment strategies: combination therapy for early rheumatoid arthritis (COBRA) Classic (methotrexate (MTX)+ sulfasalazine+60 mg prednisone tapered to 7.5 mg daily from W7), COBRA Slim (MTX +30 mg prednisone tapered to 5 mg from W6) and COBRA Avant-Garde (MTX+leflunomide+30 mg prednisone tapered to 5 mg from W6). Treatment modifications to target low-disease activity were mandatory from W8, if desirable and feasible according to the rheumatologist. The primary outcome was remission (28 joint disease activity score calculated with C-reactive protein <2.6) at W16 (intention-to-treat analysis). Secondary endpoints were good European League Against Rheumatism response, clinically meaningful health assessment questionnaire (HAQ) response and HAQ equal to zero. Adverse events (AEs) were registered.

Results Data from 98 Classic, 98 Slim and 94 Avant-Garde patients were analysed. At W16, remission was reached in 70.4% Classic, 73.6% Slim and 68.1% Avant-Garde patients (p=0.713). Likewise, no significant differences were shown in other secondary endpoints. However, therapy-related AEs were reported in 61.2% of Classic, in 46.9% of Slim and in 69.1% of Avant-Garde patients (p=0.006).

Conclusions For high-risk eRA, MTX associated with a moderate step-down dose of GCs was as effective in inducing remission at W16 as DMARD combination therapies with moderate or high step-down GC doses and it showed a more favourable short-term safety profile.


INTRODUCTION

While in the past patients with early rheumatoid arthritis (eRA) were treated conservatively, current guidelines recommend treating high-risk patients intensively, early and to target.1–4 A lot of interesting and important pioneering work has already been done, but many questions regarding the optimal dosage and combination of medication in the management of patients with eRA remain unaddressed.4

Trials using early intensive combination strategies with classical disease-modifying antirheumatic drugs (DMARDs) and glucocorticoids (GCs) gave rise to the ‘early window of opportunity’ theory.5–10 This implies that if intensive treatment is initiated early in the disease process and disease activity is rapidly controlled, more patients will go into long-term remission with better functional and radiographic outcomes later on.11–16 Discussion still exists about the optimal way to rapidly induce remission at the individual patient level. Some patients might do equally well on methotrexate (MTX) monotherapy, and even in case of insufficient response, intensifying to triple DMARD therapy or a combination with a biological can rescue patients later on.17 A delay in optimal disease control might indeed not necessarily result in worse outcomes at standard evaluation time points, but unfortunately does not take into account the cumulative disease activity patients have to suffer before arriving at these endpoints. This illustrates that the patient perspective is still understudied in traditional eRA trials.

Guidelines suggest adapting treatment according to prognostic factors.17–18 Unfortunately, this does not guarantee a favourable outcome in daily practice.18 Until better prediction models become available, the most effective approach to use the window of opportunity is to combine classical DMARDs with rapid remission inducing agents like GCs or biologicals.

GCs are commonly used to bridge the onset of the therapeutic effect of DMARDs, to rapidly control inflammation and to prevent radiographic damage.19–21 During the difficult initial treatment

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Conclusions For high-risk eRA, MTX associated with a moderate step-down dose of GCs was as effective in inducing remission at W16 as DMARD combination therapies with moderate or high step-down GC doses and it showed a more favourable short-term safety profile.

weeks, GCs can relieve pain, stiffness and disability, allowing patients to take up again their role in society more rapidly and potentially preventing chronic disease behaviour. The perception on GCs, however, remains ambiguous in both the patient’s and the physician’s mind. Thus many rheumatologists hesitate to prescribe GCs due to fear for side effects. Little is yet known about the optimal initial dose, treatment duration and administration route.

Ample evidence indicates that compared with MTX monotherapy, biologics combined with MTX are more efficacious in eRA. Unfortunately, insufficient clarity exists whether these agents can be used as remission induction agents in bridging strategies just as well as GCs since most trial protocols led to persistent biological use after the induction phase. Moreover, tumour necrosis factor-blocking agents did not demonstrate superior efficacy compared with induction regimens with GCs. Thus, administering GCs could avoid or postpone starting expensive long-term biological therapy.

The debate on the ideal DMARD content of initial RA treatment strategies is still ongoing. Triple therapy (MTX, sulfasalazine (SSZ) and hydroxychloroquine), combination therapy for early rheumatoid arthritis (COBRA)-like schemes (MTX±SSZ+ salazine (SSZ) and hydroxychloroquine), combination therapy for persistent biological use after the induction phase. Strate- ges just as well as GCs since most trial protocols led to persistent biological use after the induction phase. Moreover, tumour necrosis factor-blocking agents did not demonstrate superior efficacy compared with induction regimens with GCs. Thus, administering GCs could avoid or postpone starting expensive long-term biological therapy.

The aim of the current study was to compare in high-risk patients with eRA the efficacy and safety of different initial DMARD combinations and GC bridging schemes, 16 weeks after initiation.

**PATIENTS AND METHODS**

**The CareRA study**

Care in early RA (CareRA—EudraCT number: 2008-007225-39) is a prospective 2-year investigator-initiated multicentre randomised controlled trial rooted in daily practice. The trial is conducted in 13 Flemish rheumatology centres: two academic centres, seven general hospitals and four private practices.

**Patients**

Patients with RA, as defined by the American College of Rheumatology 1987 revised criteria, were recruited between January 2009 and May 2013. The main inclusion criteria were having a disease duration ≤1 year and being DMARD and GCs treatment naive. Disease duration was defined as time elapsed between RA diagnosis and treatment initiation. Patients having contraindications for intensive treatment combinations with GCs as judged by the treating rheumatologist were excluded. See online supplement 1 for a full list of exclusion criteria.

Patients were allocated to a high-risk group based on an algorithm constructed with classical RA prognostic factors: erosions, rheumatoid factor (RF) and/or anticitrullinated protein antibody (ACPA) and disease activity score based on C-reactive protein (CRP) status (DAS28 (CRP)) at screening (figure 1).

**Design**

After risk allocation, high-risk patients were randomised into 1/3 treatment arms:

- **COBRA Classic**: 15 mg MTX weekly, 2 g SSZ daily and a weekly step-down scheme of oral GCs (60-40-25-20-15-10-7.5 mg prednisone). This scheme has a higher dose of MTX than the original cobra schedule, based on experience in daily clinical practice.
- **COBRA Slim**: 15 mg MTX weekly with a weekly step-down scheme of oral GCs (30-20-12.5-10-7.5-5 mg prednisone).
- **COBRA Avant-Garde**: 15 mg MTX weekly, 10 mg leflunomide (LEF) daily and a weekly step-down scheme of oral GCs (30-20-12.5-10-7.5-5 mg prednisone).

The GC dose was tapered down weekly except for the lowest dose (7.5 mg in COBRA Classic and 5 mg in the other arms), which was maintained until week (W) 28. Then, GCs were tapered on a weekly basis by leaving out one daily dose each week over a period of 6 weeks until complete discontinuation. Prophylactic treatment including oral folic acid, calcium and vitamin D supplements was prescribed to all patients. Furthermore, all patients received face-to-face education and ino-material (leaflet, DVD and website) about the disease as well as on the proposed treatment at screening. Additional information was given on demand.

A treat-to-target approach was used in a tight control setting, aiming for a DAS28(CRP) ≤3.2. If patients failed to reach this target, treatment adjustments were made according to protocol from W8 onwards. Treating rheumatologists had the option not to adapt treatment, but in that case they had to motivate their decision based on a predefined list of specific clinical

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**Figure 1** Classification of patients in high or low risk according to classic prognostic factors. ACPA, anticitrullinated protein antibody; DAS28 (CRP), 28 joint disease activity score calculated with C-reactive protein; RF, rheumatoid factor.
conditions. Other treatment adjustments not stated in the protocol could not be implemented by the treating physician. The first adjustment in all treatment arms was a weekly increase in MTX dose to 20 mg. If necessary a second adjustment could be made from 8 weeks after the first adjustment. The second adjustment depended on the treatment arm: an SSZ dose increase to 3 g daily in COBRA Classic, an LEF addition of 10 mg daily in COBRA Slim or an LEF dose increase to 20 mg daily in COBRA Avant-Garde. If patients did not reach the target after two predefined treatment adjustments during the first year, this was considered a strategy failure for efficacy reasons. Intramuscular and intra-articular GC injections were allowed maximally every 8 weeks, but not within 4 weeks preceding W16. Concomitant therapy with non-steroidal anti-inflammatory drugs and analgesics was allowed and registered.

Assessment
Patients were assessed at screening, baseline, W4, W8 and W16. A maximum of 4 weeks were allowed between screening and baseline. In case a treatment adjustment was required according to the protocol at W8, an optional visit was performed at W12. Demographics were registered at screening and clinical parameters, DAS28(CRP) and health assessment questionnaire (HAQ) at every visit (table 1).

Safety and toxicity
At each visit, patients were asked about any adverse events (AE) and medication changes. Each reported AE was registered and evaluated in relation to therapy, seriousness and severity by the treating rheumatologist. In case of toxicity, the protocol predefined schemes for tapering/interrupting the assigned treatment strategy. If toxicity was persistent, this was considered a strategy failure for safety reasons.

Outcomes
The primary outcome of this study was the proportion of patients in remission (DAS28(CRP)<2.6) at W16. Secondary outcomes were the proportion of good European League Against Rheumatism (EULAR) responders (DAS28(CRP) change>1.2 and DAS28(CRP)<3.2), the proportion of patients having a clinically meaningful improvement of the HAQ (HAQ change>0.22) and the proportion of patients having an HAQ equal to zero at W16.

Statistical analysis
The study was designed as a superiority analysis of Classic versus Slim and Avant-Garde versus Slim. Sample-size calculation was based upon the proportion of patients in remission at W16. Eighty-five patients per treatment arm were required for a power of 80% and significance level of 0.05, starting from an estimated clinically relevant difference in effect size of 20%. All patients starting treatment were analysed.

Missing data were handled as follows. Screening variables were used to impute missing baseline variables and vice versa. A maximum likelihood model (by the Expectation–Maximisation algorithm) was applied to impute missing data needed to calculate the DAS28(CRP) at W4, W8 and W16.

An intention-to-treat analysis was performed by \( \chi^2 \) or Kruskal–Wallis test, where appropriate. Area under the curve (AUC) analysis was used to evaluate the DAS28(CRP) over time. All statistical analyses were carried out using the Statistical Package for the Social Sciences (SPSS) V20. A p value <0.05 was considered statistically significant.

RESULTS
A total of 400 patients were screened and 380 patients were included in CareRA. Seventy-five per cent of these patients were included in non-university centres. No differences in demographic and clinical characteristics were observed between screened and included patients. In total, 290 patients were allocated to the high-risk group and randomly assigned to treatment in the COBRA Classic (98), COBRA Slim (98) and COBRA Avant-Garde (94) arm. Randomisation resulted in similar baseline characteristics between groups (table 1).

Figure 2 describes the patient disposition from screening until W16.

Efficacy
Primary outcome
Remission was achieved in 70.4% (68/98) COBRA Classic patients, 73.5% (72/98) COBRA Slim patients and 68.1% (64/94) COBRA Avant-Garde patients (p=0.713) at W16 (figure 3A).

Secondary outcomes
At W16, a good EULAR response was reached in 79.6% of Classic patients, 79.6% of Slim patients and 76.6% of

Table 1

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>COBRA Classic</th>
<th>COBRA Slim</th>
<th>COBRA Avant-Garde</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>98</td>
<td>98</td>
<td>94</td>
</tr>
<tr>
<td>Age (years)</td>
<td>53.2±11.9</td>
<td>51.8±13.1</td>
<td>51.2±12.8</td>
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<tr>
<td>BMI (kg/m²)</td>
<td>26.0±9.43</td>
<td>26.8±4.2</td>
<td>26.5±4.2</td>
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<td>Gender (women)</td>
<td>65.3%</td>
<td>64.3%</td>
<td>69.1%</td>
</tr>
<tr>
<td>Smoking status (ever)</td>
<td>57.1%</td>
<td>59.2%</td>
<td>60.6%</td>
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<td>Alcohol intake (yes)</td>
<td>55.1%</td>
<td>56.1%</td>
<td>54.3%</td>
</tr>
<tr>
<td>Symptom duration (weeks)</td>
<td>33.8±35.5</td>
<td>33.2±38.2</td>
<td>42.2±65.6</td>
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<tr>
<td>Disease duration (weeks)</td>
<td>1.8±3.1</td>
<td>2.6±3.3</td>
<td>3.1±6.3</td>
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<tr>
<td>Employed before symptom onset (yes)</td>
<td>52.0%</td>
<td>65.3%</td>
<td>62.8%</td>
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<td>Employed at screening (yes)</td>
<td>44.9%</td>
<td>53.1%</td>
<td>51.1%</td>
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<tr>
<td>Comorbidities at screening (yes)</td>
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<td>74.5%</td>
<td>64.9%</td>
</tr>
<tr>
<td>Nocturnal pain (yes)</td>
<td>69.4%</td>
<td>72.4%</td>
<td>68.1%</td>
</tr>
<tr>
<td>Morning stiffness (yes)</td>
<td>74.5%</td>
<td>68.4%</td>
<td>58.4%</td>
</tr>
<tr>
<td>RF (yes)</td>
<td>79.6%</td>
<td>83.7%</td>
<td>75.5%</td>
</tr>
<tr>
<td>ACPA (yes)</td>
<td>77.6%</td>
<td>79.6%</td>
<td>77.7%</td>
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<td>Erosions (yes)</td>
<td>32.7%</td>
<td>32.7%</td>
<td>34.0%</td>
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<td>Total TJC</td>
<td>14.7±9.5</td>
<td>13.7±8.2</td>
<td>14.0±9.0</td>
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<tr>
<td>Total SJC</td>
<td>11.9±8.9</td>
<td>10.8±6.5</td>
<td>10.5±6.8</td>
</tr>
<tr>
<td>TJC/28</td>
<td>9.5±6.0</td>
<td>8.5±5.5</td>
<td>8.2±5.5</td>
</tr>
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<td>SJC/28</td>
<td>7.9±6.0</td>
<td>7.1±4.6</td>
<td>7.0±5.1</td>
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<tr>
<td>PGA (0–100)</td>
<td>59.5±21.7</td>
<td>56.2±21.7</td>
<td>54.2±24.3</td>
</tr>
<tr>
<td>Pain (0–100)</td>
<td>59.5±23.6</td>
<td>56.5±21.9</td>
<td>56.9±23.88</td>
</tr>
<tr>
<td>Fatigue (0–100)</td>
<td>50.6±26.0</td>
<td>49.0±21.3</td>
<td>48.6±23.78</td>
</tr>
<tr>
<td>PhGA (0–100)</td>
<td>54.7±18.5</td>
<td>53.1±18.1</td>
<td>51.8±18.2</td>
</tr>
<tr>
<td>ESR</td>
<td>33.5±25.2</td>
<td>32.1±23.3</td>
<td>25.1±17.7</td>
</tr>
<tr>
<td>CRP</td>
<td>19.7±28.9</td>
<td>21.5±33.3</td>
<td>15.1±20.0</td>
</tr>
<tr>
<td>DAS28(CRP)</td>
<td>5.4±1.3</td>
<td>5.2±1.2</td>
<td>5.0±1.3</td>
</tr>
<tr>
<td>DAS28(CRP)</td>
<td>5.0±1.2</td>
<td>4.9±1.1</td>
<td>4.7±1.2</td>
</tr>
<tr>
<td>HAQ (0–3)</td>
<td>1.2±0.7</td>
<td>0.98±0.69</td>
<td>0.99±0.64</td>
</tr>
</tbody>
</table>

ACPA, anticitrullinated protein antibody; BMI, body mass index; CRP, C-reactive protein; DAS28, 28 joint disease activity score; Disease duration, time elapsed between diagnosis of RA and start of treatment; ESR, erythrocyte sedimentation rate; HAQ, health assessment questionnaire; Morning stiffness, being stiff in the morning for at least 45 min; PGA, patient global assessment; PhGA, physician global assessment; RF, rheumatoid factor; SJC, swollen joint count; Symptom duration, time elapsed between onset of symptoms and start of treatment; TJC, tender joint count; COBRA, combination therapy for early rheumatoid arthritis. Data are presented as means±SD or as percentages.
Avant-Garde patients (p=0.844). A clinically meaningful HAQ response was reached in 84.7% of Classic patients, 86.7% of Slim patients and 76.6% of Avant-Garde patients (p=0.271). HAQ was equal to zero in 45.9% of Classic patients, 42.9% of Slim patients and 48.9% of Avant-Garde patients (p=0.700) (table 2).

Likewise, complete case analysis (without missing data imputation) of the primary and secondary outcomes revealed no significant differences between the three treatment arms (data not shown).

Area under the curve
The mean±SD AUC for DAS28(CRP) from baseline to W16 was 10.66±3.41, 11.05±3.39 and 10.72±2.96 for the Classic, Slim and Avant-Garde, respectively (p=0.521) (figure 3B).

Treatment adaptations according to protocol
During the first 16 weeks of therapy, treatment adaptations were performed in 19.4%, 22.4% and 14.9% in the Classic, Slim and Avant-Garde arm, respectively (p=0.407). Of these patients requiring treatment adaptations at W8, 50.0%, 87.5% and 60.0% in the Classic, Slim and Avant-Garde arm, respectively, reached the low-disease activity target at W16 (p=0.086). Of 61 patients requiring per protocol treatment adaptation at W8, 39% (24/61) had no change in therapy because of contraindications or because the treating physician judged the disease sufficiently controlled. These 24 patients received no other medication and stayed on the initial strategy without treatment adaptation. Intra-articular GC injections were given in 3.1% of Classic patients, 5.1% of Slim patients and 5.1% of Avant-Garde patients (p=0.703).

Safety
Therapy-related AEs were registered in 171 out of the 290 patients (59%) during the first 16 weeks of treatment. These were reported in 61.2% of Classic, in 46.9% of Slim and in 69.1% of Avant-Garde patients (p=0.006). The total number of AEs related to Classic, Slim and Avant-Garde treatment was 148, 70 and 130, respectively, with a similar distribution for discomfort and toxicity (table 3).

DISCUSSION
In patients with eRA with unfavourable classical prognostic factors such as RF, ACPA, erosions and/or high-disease activity, MTX associated with a moderate step-down dose of GCs was as effective as DMARD combination therapies with moderate or high step-down GC doses, for remission induction at 16 weeks. Furthermore, the short-term safety profile of MTX associated with a moderate step-down dose of GCs was more favourable.

This finding has two implications. First, in association with a moderate or high GC dose, the combination of MTX with other DMARDs does not seem to be more effective compared with MTX alone, at least in the early treatment stage. Until now only a few studies have addressed the question whether DMARD combinations are superior to MTX monotherapy independent from additional GC bridging in eRA. The
tREACH trial showed that DMARD combination was better than MTX monotherapy, both in association with low-dose GC bridging. In our trial, the COBRA-like moderate-dose or high-dose GC scheme bridged the time lag before full DMARD efficacy, probably erasing any difference between the different DMARD schedules. As a consequence, less medication is needed over time, which might impact AE and possibly also patients’ adherence to treatment. The tight control setting could also correct swiftly for any suboptimal treatment regimen, explaining some of the good efficacy of COBRA Slim. However, only MTX dose adjustment and no step up to combination therapy could be implemented before W16. Furthermore, the proportion of treatment adjustments between the three arms was not significantly different.

Second, a high-dose GC scheme starting at 60 mg prednisone does not seem to improve early clinical outcomes compared with a moderate-dose scheme starting at 30 mg prednisone, regardless of the DMARD strategy used. Thus, a lower cumulative GC dose is still equally effective, perhaps avoiding long-term AEs. Furthermore, the possibility to use a lower dose of GCs, while having the same efficacy, could benefit the implementation of COBRA-like strategies. Rheumatologists appear more reluctant to administer complex therapies with high dosages of GCs, although we showed that this approach is feasible in daily practice.22 23 40–42 den Uyl et al36 reported similar results comparing an attenuated COBRA regimen with the original one in a moderately active eRA population. However, this study lacked decisive evidence, the MTX dose in the classical scheme was suboptimal and the GC scheme in the attenuated COBRA version was cumulatively comparable with the classical one.

Many rheumatologists use low-dose GCs in association with DMARDS for eRA in daily practice, much to their own and their patients’ satisfaction. They have, however, doubts about

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**Figure 3** Remission and disease activity over 16 weeks. (A) The proportion of patients in remission per treatment arm at weeks 4, 8 and 16. (B) The area under the curve of DAS28(CRP) per treatment arm during 16 weeks of treatment. DAS28(CRP), 28 joint disease activity score calculated with C-reactive protein.

<table>
<thead>
<tr>
<th></th>
<th>Cobra Classic</th>
<th>Cobra Slim</th>
<th>Cobra Avant-Garde</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Week 4</td>
<td>63 (64.3%)</td>
<td>60 (61.2%)</td>
<td>60 (70.4%)</td>
<td>0.652</td>
</tr>
<tr>
<td>Week 8</td>
<td>64 (65.3%)</td>
<td>61 (62.2%)</td>
<td>65 (73.5%)</td>
<td>0.652</td>
</tr>
<tr>
<td>Week 16</td>
<td>69 (70.4%)</td>
<td>72 (73.5%)</td>
<td>64 (68.1%)</td>
<td>0.713</td>
</tr>
</tbody>
</table>

Remission was defined as DAS28(CRP) <2.6; DAS28(CRP) = 28 joint disease activity score calculated with C-reactive protein. The proportion of patients per treatment arm was compared via $\chi^2$ testing and no statistically significant differences were found.

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the need for higher GC dosages and prolonged use. The potential advantage of a COBRA-like schedule over low-dose GCs is two-sided. First, a high or moderate dose could have a more radical biological effect on the disease process favouring ‘real’ remission induction. Low-dose GCs show only a slow genomic effect, while higher dosages show both slow genomic and faster non-genomic effects. Second, compared with using GCs only short term and discontinuously, it can be more advantageous over DMARD monotherapy in trials using GCs against GC dosage and lack of knowledge of GC side effects. Certain rheumatologists could have been biased towards a certain therapy and therefore report less therapy-related AEs. Certain patients could also be more motivated for certain treatment regimens than for others.

Another limitation was the superiority design of this study. We opted for this design because in a non-inferiority trial the number of patients needed would be doubled. Hence, we can only state that COBRA Classic and COBRA Avant-Garde are non-superior to COBRA Slim, which is not the same as claiming non-inferiority.

In conclusion, the data presented are positioning classical MTX therapy with bridging GCs at a lower dose than in the original COBRA study as a highly effective and safe remission induction therapy in more than 70% of high-risk patients with eRA and this in a close to daily practice setting applying a treat-to-target strategy.

**Table 2** Clinical outcomes at week 16 per treatment arm

<table>
<thead>
<tr>
<th></th>
<th>COBRA Classic</th>
<th>COBRA Slim</th>
<th>COBRA Avant-Garde</th>
<th>p Value</th>
<th>( \Delta ) between Classic vs Slim (95% CI)</th>
<th>( \Delta ) between Avant-Garde vs Slim (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>98</td>
<td>98</td>
<td>94</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DAS28 (CRP) change</td>
<td>2.8±1.2</td>
<td>2.6±1.2</td>
<td>2.4±1.3</td>
<td>0.140</td>
<td>0.2 (−0.13 to 0.52)</td>
<td>−0.2 (−0.49 to 0.21)</td>
</tr>
<tr>
<td>Remission</td>
<td>70.4%</td>
<td>73.5%</td>
<td>68.1%</td>
<td>0.713</td>
<td>−3.1% (−15.4% to 9.5%)</td>
<td>−5.4% (−18.0% to 7.4%)</td>
</tr>
<tr>
<td>Low-disease activity</td>
<td>84.7%</td>
<td>86.7%</td>
<td>87.2%</td>
<td>0.863</td>
<td>−2.0% (−12.0% to 7.9%)</td>
<td>0.5% (−9.3% to 10.2%)</td>
</tr>
<tr>
<td>Good EULAR response</td>
<td>79.6%</td>
<td>79.6%</td>
<td>76.6%</td>
<td>0.844</td>
<td>0.0% (−11.3% to 11.3%)</td>
<td>−3.0% (−14.7% to 8.7%)</td>
</tr>
<tr>
<td>Moderate EULAR response</td>
<td>98.0%</td>
<td>95.9%</td>
<td>93.6%</td>
<td>0.320</td>
<td>2.1% (−3.6% to 8.2%)</td>
<td>−2.3% (−9.6% to 4.6%)</td>
</tr>
<tr>
<td>HAQ change</td>
<td>0.8±0.6</td>
<td>0.6±0.6</td>
<td>0.7±0.6</td>
<td>0.081</td>
<td>0.2 (0.02 to 0.37)</td>
<td>0.1 (−0.17 to 0.19)</td>
</tr>
<tr>
<td>Clinically meaningful HAQ change</td>
<td>84.7%</td>
<td>76.5%</td>
<td>76.6%</td>
<td>0.271</td>
<td>8.2% (−3.0% to 19.1%)</td>
<td>0.1% (−11.9% to 12.0%)</td>
</tr>
<tr>
<td>HAQ=0</td>
<td>45.9%</td>
<td>42.9%</td>
<td>48.9%</td>
<td>0.700</td>
<td>3.0% (−10.7% to 16.6%)</td>
<td>6.0% (−7.9% to 19.7%)</td>
</tr>
</tbody>
</table>

Remission was defined as DAS28(CRP) <2.6. Low-disease activity was defined as DAS28(CRP) ≤3.2. Good EULAR response was defined as low-disease activity with a DAS28(CRP) change >1.2. Moderate EULAR response was defined as DAS28(CRP) change ≥1.2 and a DAS28(CRP) changes between 0.6 and 1.2. Clinically meaningful HAQ change was defined as an HAQ change >0.22. Data are presented as mean±SD or as percentages. Statistical analysis was performed by \( \chi^2 \) or Kruskal–Wallis test. A p value <0.05 was considered statistically significant.

**Table 3** Number of adverse events per treatment arm during 16 weeks of treatment

<table>
<thead>
<tr>
<th></th>
<th>COBRA Classic</th>
<th>COBRA Slim</th>
<th>COBRA Avant-Garde</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>91</td>
<td>96</td>
<td>91</td>
</tr>
<tr>
<td>AE related to therapy</td>
<td>148</td>
<td>70</td>
<td>130</td>
</tr>
<tr>
<td>Type related AE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Discomfort</td>
<td>111</td>
<td>50</td>
<td>96</td>
</tr>
<tr>
<td>Toxicity</td>
<td>27</td>
<td>10</td>
<td>23</td>
</tr>
<tr>
<td>Infection</td>
<td>5</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Others</td>
<td>4</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>Surgery</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Severity of related AE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>121</td>
<td>64</td>
<td>103</td>
</tr>
<tr>
<td>Moderate</td>
<td>23</td>
<td>5</td>
<td>21</td>
</tr>
<tr>
<td>Severe</td>
<td>4</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Serious AE</td>
<td>2</td>
<td>1</td>
<td>3</td>
</tr>
</tbody>
</table>

AE, adverse event; COBRA, combination therapy for early rheumatoid arthritis.

‘window of opportunity’, is crucial for longer-term outcome at the biological and probably also at a psychosocial level. Long-term disease control and patient-reported outcomes after 1 and 2 years are awaited in CareRA.

The safety analysis strengthened the efficacy outcomes further. The proportion and number of related AEs was comparable in Classic and Avant-Garde, while Slim patients had half the related AEs. GC dosage does not make any difference in the frequency or type of related AEs at this stage. Remarkable are the comparable number of AEs of the combination therapies with different GC dose, underscoring again the prejudice against GC dosage and lack of knowledge of GC side effects.

The first two limitations are related to the design of this study, although unavoidable in a pragmatic trial aiming to reflect daily clinical practice. First, medication adherence was not measured. However, if adherence was lower in a certain trial arm, the same could be expected from this treatment regimen in daily clinical practice. Second, no blinding was implemented. Rheumatologists could have been biased towards a certain therapy and therefore report less therapy-related AEs. Certain patients could also be more motivated for certain treatment regimens than for others.

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Data sharing statement Further analyses are available upon request from DDC/PV, the corresponding authors at KU Leuven.

REFERENCES


Methotrexate in combination with other DMARDs is not superior to methotrexate alone for remission induction with moderate-to-high-dose glucocorticoid bridging in early rheumatoid arthritis after 16 weeks of treatment: the CareRA trial


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