Olokizumab, a monoclonal antibody against interleukin 6, in combination with methotrexate in patients with rheumatoid arthritis inadequately controlled by methotrexate: efficacy and safety results of a randomised controlled phase III study

Evgeniy Nasonov, Saeed Fatenejad, Eugen Feist, Mariana Ivanova, Elena Korneva, Diana G Krechikova, Aleksey L Maslyanskiy, Mikhail Samsonov, Rumen Stoilov, Elena V Zonova, Mark Genovese

ABSTRACT

Objective To evaluate the efficacy and safety of olokizumab (OKZ) in patients with active rheumatoid arthritis despite treatment with methotrexate (MTX).

Methods In this 24-week multicentre, placebo-controlled, double-blind study, patients were randomised 1:1:1 to receive subcutaneously administered OKZ 64 mg once every 2 weeks, OKZ 64 mg once every 4 weeks, or placebo plus MTX. The primary efficacy endpoint was the proportion of patients achieving an American College of Rheumatology 20% (ACR20) response at week 12. The secondary efficacy endpoints included percentage of subjects achieving Disease Activity Score 28-joint count based on C reactive protein ≤3.2, Health Assessment Questionnaire Disability Index at week 12, ACR50 response and Clinical Disease Activity Index ≤2.8 at week 24. Safety and immunogenicity were assessed throughout the study.

Results A total of 428 patients were randomised. ACR20 responses were more frequent with OKZ every 2 weeks (63.6%) and OKZ every 4 weeks (70.4%) than placebo (25.9%) (p<0.0001 for both comparisons). There were significant differences in all secondary efficacy endpoints between OKZ-treated arms and placebo. Treatment-emergent serious adverse events (TESAEs) were reported by more patients in the OKZ groups compared with placebo. Infections were the most common TESAEs. No subjects developed neutralising antidrug antibodies.

Conclusions Treatment with OKZ was associated with significant improvement in signs, symptoms and physical function of rheumatoid arthritis without discernible differences between the two regimes. Safety was as expected for this class of agents. Low immunogenicity was observed.

Trial registration number NCT02760368.

INTRODUCTION

Rheumatoid arthritis (RA) is a chronic inflammatory disease that if left inadequately treated can lead to significant disability, morbidity and mortality. Current guidelines recommend a treat to target strategy in order to attain acceptable level of disease control and prevent long-term disability. A number of effective therapies with different modes of action are currently available for RA; however, many patients with active RA fail to achieve defined targets of therapy, namely low disease activity or remission.

The proinflammatory cytokine interleukin 6 (IL-6) plays a significant role in the pathogenesis of RA and two anti-IL-6 receptor (IL-6R) antibodies have been shown to be relatively safe and effective...
Olokizumab (OKZ) is an anti-IL-6 monoclonal antibody that binds directly to IL-6 at a specific site and neutralises its activity through blocking hexamer formation of the extracellular signalling complex inhibiting transmembrane signalling. In early clinical studies it was shown that OKZ resulted in a rapid reduction in the level of IL-6 and C reactive protein (CRP) that lasted over an extended period of time due to OKZ’s long half-life of approximately 31 days. OKZ in doses ranging from 60 mg to 240 mg administered every 2 weeks or every 4 weeks was relatively safe and effective in reducing signs and symptoms of RA in two phase II randomised controlled trials in patients with RA who had failed to respond to antitumour necrosis factor (anti-TNF) therapy. Based on findings from these two studies, as well as information from earlier studies, two doses of OKZ, 64 mg every 2 weeks and 64 mg every 4 weeks, were selected for advancement to phase III. The lowest two doses tested in phase II were chosen to achieve efficacy while minimising potential adverse effects. Here we report the full results of the first completed phase III study of OKZ in patients with active RA despite treatment with methotrexate (MTX).

**METHODS**

**Study design**

This phase III, randomised, double-blind, placebo-controlled, parallel-group, multicentre trial was conducted at 42 hospitals in Russia, Belarus and Bulgaria from May 2016 to April 2019. Written informed consent was obtained from each patient. After week 24, patients had the choice of either enrolling into an ongoing open-label study or entering the safety follow-up period.

**Patient inclusion and exclusion criteria**

Adults were eligible for inclusion if they had active RA (swollen joint count ≥ 66; joint count ≥ 6 (68-joint count) and CRP > 6 mg/L) classified by the American College of Rheumatology/European League Against Rheumatism 2010 revised classification criteria for at least 12 weeks prior to screening and had an inadequate response to treatment with MTX for at least 12 weeks at a dose of 15–25 mg/week (or ≥ 10 mg/week if intolerant to higher doses). The dose and route of administration of MTX must have been stable for at least 6 weeks. Exclusion criteria were other inflammatory or rheumatic diseases and Steinbrocker class IV functional capacity. Also excluded were those who had a prior exposure to IL-6 or IL-6R inhibitors, Janus kinase inhibitors, those treated with cell-depleting agents or those concurrently on disease-modifying antirheumatic drugs (DMARDs) other than MTX. Prior use of biologic DMARDs was an exclusion criterion with the exception of subjects who discontinued anti-TNF therapy due to reasons other than lack of efficacy. Non-steroidal anti-inflammatory drugs and glucocorticoids in doses less than or equal to 10 mg/day prednisone or equivalent were allowed if their doses were stable during the 2 weeks prior to study enrolment. Patients with a history of malignancies within the last 5 years (successfully treated carcinoma of the cervix in situ, basal cell carcinoma and squamous cell carcinoma of the skin were allowed if beyond 1 year prior to screening), recurrent infections, primary or secondary immunodeficiency, hepatitis B or C, active tuberculosis (TB) or other uncontrolled medical conditions, or prespecified abnormal laboratory values were excluded. Patients with latent TB infection were allowed to participate if they had started appropriate anti-TB therapy at least 30 days prior to randomisation (see online supplemental material for additional selection criteria).

**Randomisation and blinding**

Patients were randomised 1:1:1 to receive subcutaneous injections of OKZ 64 mg every 2 weeks, OKZ 64 mg once every 4 weeks, or placebo (PBO) for 24 weeks with continuation of their background MTX using an automated randomisation system. Subjects who discontinued the randomised treatment earlier were required to continue the study without study treatment administration; patients could discontinue study treatment but completed the study.

All patients, investigators, clinical site staff, contract research organisation’s staff and the sponsor’s staff directly involved in the study were blinded. Joint assessments were performed by independent assessors, blinded to study drug assignment and all other study assessments (see online supplemental material for additional details).

**Rescue medication**

Starting at week 14, non-responders, defined as subjects in any treatment group who did not improve by at least 20% in both swollen and tender joint counts (66–68 joints), were prescribed rescue medication (sulfasalazine and/or hydroxychloroquine) in addition to their study treatment (see online supplemental material for details of the prior and concomitant medications).

**Endpoints**

The primary endpoint was the proportion of patients achieving the American College of Rheumatology 20% (ACR20) response at week 12. Ranked secondary endpoints were percentage of subjects achieving Disease Activity Score 28 based on C reactive protein (DAS28-CRP) < 3.2 at week 12, improvement in physical ability from baseline to week 12 measured by the Health Assessment Questionnaire Disability Index (HAQ-DI), ACR50 response at week 24 and percentage of subjects with Clinical Disease Activity Index (CDAI) ≤ 2.8 (remission) at week 24. Quality of life was assessed using several questionnaires including Short Form-36 (SF-36) Physical Component Summary (PCS), Mental Component Summary (MCS) and total scores, and the Functional Assessment of Chronic Illness Therapy-Fatigue Scale (FACIT-F).

Standard safety monitoring, including assessment of adverse events, serious adverse events and laboratory tests via the central laboratory, was performed regularly. Determination of antidrug antibodies (ADA) in plasma samples was done using electrochemiluminescence assay (Covance...
Laboratories, Harrogate, North Yorkshire, UK). For detection of neutralising ADAs, a cell-based assay was used (Eurofins BioPharma Product Testing Munich, Planegg/Munich, Germany).

An independent external Data and Safety Monitoring Board reviewed the safety data throughout the study. Major adverse cardiovascular events (MACE) were adjudicated by a Cardiovascular Adjudicated Committee and were defined as cardiovascular death or death from undetermined cause, non-fatal myocardial infarction, non-fatal stroke, transient ischaemic attack, hospitalisation for unstable angina requiring unplanned revascularisation and coronary revascularisation procedures.

Statistical analyses
The ACR20 response at week 12 for each of the active treatment groups was compared with PBO using a $2 \times 2$ $\chi^2$ test for equality of proportions. To control the overall type I error rate at a one-sided $\alpha=0.025$, Bonferroni adjustment was used for the tests related to each of the OKZ dose regimens versus PBO (ie, one-sided $\alpha=0.0125$ for each dose). A gate-keeping strategy with a fixed order of hypothesis was used for the primary and secondary endpoints within each OKZ dose regimen independently (figure 1).

To detect a difference between at least one OKZ dose regimen and PBO, a sample size of 420 patients randomised in a 1:1:1 ratio was estimated to ensure sufficient disjunctive power (100% for testing the primary hypothesis (ACR20 at week 12) and 98% for the secondary endpoint of DAS28-CRP $<3.2$ rate at week 12).

The secondary endpoints that were binary in nature were analysed as per the primary endpoint. For analyses of binary

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**Figure 1** Gate-keeping strategy. $p_{\text{Sup,q2w}}$ and $p_{\text{Sup,q4w}}$ represent p values from a one-sided test of superiority versus placebo for OKZ dose regimens 64 mg q2w and q4w. ACR, American College of Rheumatology response; CDAI, Clinical Disease Activity Index; DAS28-CRP, Disease Activity Score 28 based on C reactive protein; HAQ-DI, Health Assessment Questionnaire Disability Index; OKZ, olokizumab; q2w, every 2 weeks; q4w, every 4 weeks; Wk, week.
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Enrollment

- 785 patients assessed for eligibility (screened)
- 428 patients randomized

Allocation

- 143 patients randomized to OKZ q2w + MTX and received allocated intervention
- 142 patients randomized to OKZ q4w + MTX and received allocated intervention
- 143 patients randomized to PBO + MTX. 142 received intervention, 1 did not (randomized by mistake)

Treatment

- Dropout
  - Lost to follow-up n=14/9.8%
  - Discontinued by sponsor n=0
  - AE n=7/4.9%
  - Subject refusal n=7/4.9%
  - Lack of efficacy n=0
  - Withdrawal if IC n=7/4.9%
  - Other n=0
- Completed treatment n=129/90.2%
  - Subjects with rescue n=14/9.9%
  - More than 1 rescue n=0
  - Completed study n=130/90.9%
  - Entered OLE n=122/86.3%

- Dropout
  - Lost to follow-up n=11/7.7%
  - Discontinued by sponsor n=0
  - AE n=5/3.5%
  - Subject refusal n=6/4.2%
  - Lack of efficacy n=2/1.4%
  - Withdrawal if IC n=3/2.1%
  - Other n=1/0.7%
- Completed treatment n=131/92.3%
  - Subjects with rescue n=10/7.0%
  - More than 1 rescue n=0
  - Completed study n=134/94.4%
  - Entered OLE n=127/88.4%

- Dropout
  - Lost to follow-up n=8/5.6%
  - Discontinued by sponsor n=1
  - AE n=1/0.7%
  - Subject refusal n=6/4.2%
  - Lack of efficacy n=1/0.7%
  - Withdrawal if IC n=5/3.5%
  - Other n=0
- Completed treatment n=134/93.7%
  - Subjects with rescue n=61/43%
  - More than 1 rescue n=5/3.5%
  - Completed study n=132/92.3%
  - Entered OLE n=126/88.1%

Analysis

- Efficacy analysis (ITT population) - 143 (100%)
  - Safety analyzed - 143 (100%)
- Efficacy analysis (ITT population) - 142 (100%)
  - Safety analyzed - 142 (100%)
- Efficacy analysis (ITT population) - 143 (100%)
  - Safety analyzed - 142 (99.9%)

Figure 2 Patient disposition. AE, adverse event; IC, informed consent; ITT, intention-to-treat; MTX, methotrexate; OKZ, olokizumab; OLE, open-label extension; PBO, placebo; q2w, every 2 weeks; q4w, every 4 weeks.

RESULTS

Disposition

A total of 428 patients were randomised to OKZ 64 mg every 2 weeks (n=143), OKZ 64 mg every 4 weeks (n=142) or PBO (n=143). One patient failed screening, was randomised in error to the PBO group and was withdrawn once the error was discovered, before receiving study treatment; the safety population consisted of 427 subjects (figure 2). The three treatment groups were well balanced for baseline demographic and disease characteristics (table 1).

A total of 92.1% (n=394) of subjects completed the treatment period: 92.3% (n=131) in OKZ every 4 weeks, 90.2% (n=129) in OKZ every 2 weeks and 93.7% (n=134) in the PBO group. The most common reasons for treatment discontinuation were withdrawal of informed consent and adverse events (figure 2).

A higher proportion of patients in the PBO group (43%) received rescue medication(s) compared with patients on OKZ every 4 weeks (7%) or OKZ every 2 weeks (9.8%).

At week 24 of the study, 122 (85.3%) patients on OKZ every 2 weeks, 127 (89.4%) on OKZ every 4 weeks and 126 (88.1%) on PBO were enrolled in the open-label extension study.

Efficacy

The primary efficacy endpoint, ACR20 response rate at week 12, was 70.4% in OKZ every 4 weeks and 63.6% in OKZ every 2 weeks, both significantly greater than 25.9% in the PBO group (p<0.0001 for both comparisons) (table 2). Separation of the ACR20 response in the OKZ treatment groups from PBO was seen starting around week 2 and plateauing at week 12 (figure 3).

The secondary endpoint of DAS28-CRP <3.2 at week 12 was achieved by 33.6% and 38.7% of patients on OKZ every 2 weeks and every 4 weeks, respectively, significantly higher than those in the PBO group (3.5%, p<0.0001 for both comparisons) (table 2, figure 3).

Significant improvements in physical function as assessed with HAQ-DI were observed at week 12 for subjects in both OKZ
Table 1  Demographic and other baseline characteristics (ITT population)*

| Characteristics, mean (SD) unless otherwise specified | OKZ every 2 weeks N=143 | OKZ every 4 weeks N=142 | PBO N=143 |
|-------------------------------------------------------|-------------------------|-------------------------|-----------|
| Age (years)                                           | 52.0 (11.8)             | 49.1 (12.1)             | 52.7 (11.3)|
| Female (%)                                            | 81.1                    | 83.1                    | 83.9      |
| Duration of RA (years)                                | 8.7 (8.0)               | 7.3 (7.0)               | 8.4 (7.8) |
| MTX dose (mg)†                                        | 16.1 (3.4)              | 16.3 (3.4)              | 16.1 (3.7)|
| Duration of prior MTX use (weeks)                     | 201.5 (223.1)           | 157.4 (165.6)           | 210.1 (208.2)|
| Glucocorticoid use, n (%)                             | 52 (36.4)†              | 50 (35.2)‡              | 41 (28.7)†|
| Prednisone dose or equivalent (mg)                    | 7.6 (6.0)               | 6.1 (2.3)               | 6.6 (2.4) |
| Prior exposure to TNF inhibitors, n (%)                | 0                       | 0                       | 4 (2.8)   |
| BMI (kg/m²)                                           | 26.6 (5.1)              | 26.4 (5.5)              | 26.9 (5.0)|
| RF+ (>15 IU/mL), n (%)                                | 115 (80.4)              | 122 (85.9)              | 127 (88.8)|
| Anti-CCP+, >10 IU/mL, n (%)                           | 110 (76.9)              | 115 (81.0)              | 117 (81.8)|
| CRP (mg/L)§                                           | 23.5 (23.1)             | 22.7 (22.7)             | 25.8 (28.7)|
| TJC¶                                                  | 24.4 (11.4)             | 22.2 (10.3)             | 24.0 (11.3)|
| SJC¶                                                  | 14.8 (6.5)              | 14.5 (6.7)              | 14.6 (6.9) |
| DAS28-CRP§                                            | 6.0 (0.7)               | 5.9 (0.7)               | 6.0 (0.8) |
| CDAI score (0–76)                                     | 40.5 (9.8)              | 38.7 (9.4)              | 40.4 (10.5)|
| HAQ-DI score                                         | 1.74 (0.47)             | 1.64 (0.50)             | 1.78 (0.49)|
| PtGA (VAS) (mm)                                      | 70.4 (16.0)             | 68.5 (14.5)             | 69.6 (15.9)|
| Pain (VAS) (mm)                                      | 70.2 (16.3)             | 67.4 (18.5)             | 68.3 (17.6)|
| PGA (VAS) (mm)                                       | 70.5 (13.9)             | 66.4 (14.2)             | 68.0 (14.3)|

Pain: patient assessment of pain.
*All patients with exception of one were Caucasian.
†100% patients were on MTX.
‡Upper limit of normal: >6 mg/L.
§Joint counts were assessed based on 66–68 joint counts.
| anti-CCP+, anti-citrullinated peptide positivity; BMI, body mass index; CDAI, Clinical Disease Activity Index; CRP, C reactive protein; DAS28-CRP, Disease Activity Score 28 based on C reactive protein; HAQ-DI, Health Assessment Questionnaire Disability Index; ITT, intention-to-treat; MTX, methotrexate; N, number of subjects; OKZ, olokizumab; PBO, placebo; PGA, Physician Global Assessment of Disease Activity; PtGA, Patient Global Assessment of Disease Activity; RA, rheumatoid arthritis; RF+, rheumatoid factor positivity; SJC, swollen joint count; TJC, tender joint count; TNF, tumour necrosis factor; VAS, Visual Analogue Scale.

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Table 2  Efficacy results in the intent-to-treat population (NRI)

|                   | OKZ every 2 weeks | OKZ every 4 weeks | PBO |
|-------------------|-------------------|-------------------|-----|
|                   | N=143             | N=142             | N=143 |
| ACR20 response, n (%), week 12 (primary endpoint) | 91 (63.6)*       | 100 (70.4)*       | 37 (25.9)* |
| ACR50 response, n (%), week 24 | 61 (42.7)*     | 69 (48.6)*       | 11 (7.7) |
| ACR70 response, n (%), week 24 | 28 (19.6)     | 32 (22.5)       | 3 (2.1) |
| DAS28-CRP<3.2, n (%), week 12 | 48 (33.6)*    | 55 (38.7)*      | 5 (3.5) |
| HAQ-DI week 12 | 0.97 (0.4)       | 1.03 (0.4)       | 0.38 (0.3) |
| CDAI ≤2.8, n (%), week 24 | 12 (8.4)‡   | 11 (7.7)‡      | 0 |
| DAS28-CRP<2.6, n (%), week 24 | 31 (21.7)   | 40 (28.2)      | 5 (3.5) |
| DAS28-CRP, change from baseline, week 24 | 6 (4.2)‡     | 8 (5.8)‡       | 1 (0.7) |
| Treatment comparison vs PBO | 1.7 (0.1)     | 1.7 (0.1)       | 1.7 (0.1) |
| 97.5% CI for LMS difference | −1.7 to −1.0 | −2.0 to −1.4 | 7 (4.9) |
| Treatment comparison vs PBO | −0.47 to −0.21 | −0.49 to −0.23 | 0 |
| CDAI <10†, n (%), week 12 | 1 (0.7)‡  | 1 (0.7)‡       | 0 |

*P value difference from PBO <0.0001.
†Results for other than primary and secondary endpoints were not tested for significance.
‡P value difference from PBO <0.001.

AUCR50 response at week 24 was achieved by 48.6% of patients on OKZ every 4 weeks, 42.7% on OKZ every 2 weeks and 7.7% on PBO (p<0.0001 for comparisons of OKZ groups vs PBO) (table 2, figure 3).

Dissemination of evidence as CDAI ≤2.8 was achieved at week 24 by 7.7% of patients on OKZ every 4 weeks and by 8.4% on OKZ every 2 weeks. No subjects achieved this endpoint in the PBO group (p=0.0003 for OKZ every 4 weeks vs PBO and p=0.0002 for OKZ every 2 weeks vs PBO comparisons) (table 2, figure 3). The percent mean changes in ACR response criteria parameters and CDAI score parameters are presented in online supplemental figure 1. The number of missing observations for key efficacy outcomes is presented in online supplemental table 1. The results of the primary and ranked secondary endpoints were confirmed by predefined sensitivity analyses and a post-hoc linear mixed model analysis (data available on request).

Subgroup analyses of the ACR20 response did not show influence of country, gender, age, weight, body mass index, baseline disease severity, time since diagnosis, duration of prior MTX use, or anticyclic citrullinated peptide and rheumatoid factor status on the efficacy of OKZ (data available on request).

In parallel with the main efficacy endpoints, there were marked increases (improvement) in SF-36 mental component scores from baseline to week 24 of approximately 8.9, 6.2 and 2.5 in patients on OKZ every 4 weeks, OKZ every 2 weeks and PBO, respectively. Corresponding values for SF-36 physical component scores were 8.7, 7.8 and 3.5. Likewise, FACIT-F improvements were 10.6, 8.5 and 3.7 (table 3). Other quality of life measures showed similar trends in improvement (table 3, online supplemental table 2).
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Safety

Two hundred and twenty-six patients (52.9%) reported treatment-emergent adverse events (TEAE) with similar incidences across the treatment groups (table 4).

Most TEAEs were mild to moderate in severity and non-serious, leading to study treatment discontinuation in 3.5%, 4.9% and 0.7% of patients on OKZ every 4 weeks, OKZ every 2 weeks and PBO, respectively. The most common TEAEs were investigations reported for 35.9% of patients on OKZ every 4 weeks, 35.0% on OKZ every 2 weeks and 18.3% on PBO, and infections reported for 14.1% on OKZ every 4 weeks, 15.4% on OKZ every 2 weeks and 16.2% on PBO. Injection site reactions were reported by two subjects (1.4%) in each OKZ group. A total of 20 treatment-emergent serious adverse events (TESAEs) were reported.

Incidences of TESAEs were numerically higher in patients on OKZ every 4 weeks and OKZ every 2 weeks, compared with PBO: 5.6%, 5.6% and 2.8%, respectively. The most frequently reported serious events were serious infections: 2.8% in patients on OKZ every 2 weeks and 1.4% on PBO (no serious infections were reported for OKZ every 4 weeks). One TEAE leading to death was reported in the study, sepsicaemia due to Staphylococcus aureus and toxic shock syndrome in the OKZ group every 2 weeks. There were no reports of gastrointestinal perforations or anaphylaxis.

As reported with other anti-IL-6 therapies, there were early rises in mean serum lipids noted from week 4, with a plateau that reached around week 8 (figure 4); however, no MACE was observed. Likewise, early decreases in mean blood platelets and neutrophils were seen, with a plateau reached at week 4. No
Immunogenicity

Positive confirmed ADA tests at any time post baseline were reported in six subjects (4.4%) on OKZ every 2 weeks and in nine subjects (6.6%) on OKZ every 4 weeks. No subjects had a positive result for neutralising antibodies.

DISCUSSION

CREDO 1 trial, a phase III study of OKZ in patients with active RA despite MTX, achieved the primary and all ranked secondary efficacy endpoints. This study evaluated two effective doses with a frequency of injection of once per 2 weeks and once per month, and both regimens of OKZ were superior to PBO in reducing signs and symptoms and improving disability and quality of life over a period of 24 weeks. The onset of efficacy of OKZ was rapid as detected by differences in ACR20 response rates between OKZ and PBO that were apparent within 2 weeks from the start of treatment. The study was specifically designed and sized to detect differences between OKZ doses in one variable could be by chance, especially since they were not consistently detected across efficacy endpoints. ACR20 was used as the primary endpoint due to its widely accepted and validated value in assessing the efficacy of drugs in RA over many years. While higher levels of response such as ACR50 or ACR70 responses could have been chosen as the primary outcome, use of ACR20 allows for easier comparisons with other compounds evaluated in the past that used ACR20. While ACR20 was the primary endpoint, the study included ACR50 as a ranked secondary endpoint, all of which confirmed the results of the ACR20 analysis. In this study patients had relatively high disease activity at baseline, making it more difficult to achieve ACR20 status, all of which confirmed the results of the ACR20 analysis. In this study patients had relatively high disease activity at baseline, making it more difficult to achieve ACR20 status, all of which confirmed the results of the ACR20 analysis. In this study patients had relatively high disease activity at baseline, making it more difficult to achieve ACR20 status, all of which confirmed the results of the ACR20 analysis. In this study patients had relatively high disease activity at baseline, making it more difficult to achieve ACR20 status, all of which confirmed the results of the ACR20 analysis.

Table 3  Patient-reported outcome measures at months 3 (12 weeks) and 6 (24 weeks) *

| Week 12 | OKZ every 2 weeks N=143 | OKZ every 4 weeks N=142 | PBO N=142 | Week 24 | OKZ every 2 weeks N=143 | OKZ every 4 weeks N=142 | PBO N=142 |
| --- | --- | --- | --- | --- | --- | --- | --- |
| PtGA | −30.6 (1.7) | −31.0 (1.7) | −13.1 (1.8) | −32.1 (1.9) | −36.3 (2.0) | −19.4 (1.9) |
| Pain | −23.9 to −12.0 | −23.4 to −12.4 | −18.8 to −6.6 | −23.0 to −10.6 | −24.5 (1.9) | −37.1 (2.1) | −21.4 (2.1) |
| Pain, patients with >30% improvement, n (%) | 94 (65.7) | 86 (60.6) | 37 (25.9) | 96 (67.1) | 95 (66.9) | 57 (39.9) |
| HAQ-DI | −0.55 (0.05) | −0.65 (0.05) | −0.28 (0.05) |
| SF-36 PCS | 6.7 (0.6) | 6.0 (0.6) | 2.2 (0.6) | 7.8 (0.7) | 8.7 (0.7) | 3.5 (0.7) |
| SF-36 MCS | 6.5 (0.7) | 7.0 (0.7) | 3.5 (0.8) | 6.2 (0.8) | 8.9 (0.8) | 2.5 (0.8) |
| EQ-5D score | 19.7 (1.7) | 18.7 (1.7) | 7.4 (1.7) | 20.9 (2.0) | 23.6 (2.0) | 8.3 (2.0) |
| FACIT-F | 8.2 (0.7) | 8.7 (0.7) | 3.6 (0.7) | 8.5 (0.8) | 10.6 (0.8) | 3.7 (0.8) |

Pain: patient’s assessment of arthritis pain.

*With the exception of pain, n (%): LSM change from baseline (SE), treatment comparison vs placebo LSM difference (SE), and 97.5% CI for LSM difference are presented.

†Secondary endpoint (refer to table 2).

Discussion includes reference to other studies and clinical trials.

Disability is an important aspect of RA that originates from joint pain and joint damage and should be directly assessed in RA clinical trials. One of the secondary endpoints in the study was assessment of disability using the HAQ-DI questionnaire. The study showed that both regimens of OKZ resulted in significantly more improvement in disability than PBO. Additionally, patients had grade 3 or higher neutropaenia in accordance with the Common Terminology Criteria for Adverse Events version 4.0. Elevations in serum alanine aminotransferase values above 3× ULN at any time during the study were seen in 11.4%, 9.2% and 5.0% of patients on OKZ every 2 weeks, OKZ every 2 weeks and PBO, respectively, with no concomitant elevations in serum bilirubin above 2× ULN. Selected abnormal haematology and chemistry assessments are presented in online supplemental tables 3 and 4.
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| System organ class (preferred term) | OKZ every 2 weeks N=143, n (%) | OKZ every 4 weeks N=142, n (%) | PBO N=142, n (%) |
|-----------------------------------|-------------------------------|-------------------------------|-----------------|
| Number of subjects with at least one TEAE reported for 4% of subjects in any treatment group | 83 (58.0) | 81 (57.0) | 62 (43.7) |
| Investigations | 50 (35.0) | 51 (35.9) | 26 (18.3) |
| ALT increased | 25 (17.5) | 33 (23.2) | 11 (7.7) |
| AST increased | 16 (11.2) | 22 (15.5) | 10 (7.0) |
| White cell count decreased | 7 (4.9) | 6 (4.2) | 4 (2.8) |
| Neutrophil count decreased | 6 (4.2) | 7 (4.9) | 3 (2.1) |
| Blood cholesterol increased | 6 (4.2) | 4 (2.8) | 3 (2.1) |
| Gamma-glutamyltransferase increased | 3 (2.1) | 6 (4.2) | 4 (2.8) |
| Infections and infestations | 22 (15.4) | 20 (14.1) | 23 (16.2) |
| Nasopharyngitis | 4 (2.8) | 3 (2.1) | 6 (4.2) |
| Upper respiratory tract infection | 2 (1.4) | 6 (4.2) | 4 (2.8) |
| Blood and lymphatic system disorders | 17 (11.9) | 18 (12.7) | 15 (10.6) |
| Leucopenia | 8 (5.6) | 7 (4.9) | 4 (2.8) |
| Neutropaenia | 5 (3.5) | 9 (6.3) | 2 (1.4) |
| Anaemia | 4 (2.8) | 3 (2.1) | 6 (4.2) |
| Metabolism and nutrition disorders | 9 (6.3) | 7 (4.9) | 3 (2.1) |
| Musculoskeletal and connective tissue disorders | 6 (4.2) | 7 (4.9) | 6 (4.2) |
| Skin and subcutaneous tissue disorders | 8 (5.6) | 3 (2.1) | 2 (1.4) |
| Number and percentage with at least one key TESAE | 8 (5.6) | 8 (5.6) | 4 (2.8) |
| Investigations | 2 (1.4) | 4 (2.8) | 1 (0.7) |
| ALT increased | 2 (1.4) | 4 (2.8) | 1 (0.7) |
| AST increased | 0 | 3 (2.1) | 0 |
| Infections and infestations | 4 (2.8) | 0 | 2 (1.4) |
| Subcutaneous abscess | 2 (1.4) | 0 | 0 |
| Gastroenteritis | 0 | 0 | 0 |
| Pneumonia | 0 | 0 | 1 (0.7) |
| Pulmonary tuberculosis | 1 (0.7) | 0 | 0 |
| Staphylococcal sepsis | 1 (0.7) | 0 | 0 |
| Toxic shock syndrome | 1 (0.7) | 0 | 0 |
| Herpes zoster | 0 | 0 | 0 |
| Hepatobiliary disorders | 0 | 1 (0.7) | 0 |
| Drug-induced liver injury | 0 | 1 (0.7) | 0 |
| Neoplasms benign, malignant and unspecified (including cysts and polyps) | 0 | 1 (0.7) | 0 |
| Cervix carcinoma stage II | 0 | 1 (0.7) | 0 |
| Gastrointestinal disorders | 0 | 1 (0.7) | 0 |
| Obstructive pancreatitis | 0 | 1 (0.7) | 0 |
| Gastrointestinal perforation | 0 | 0 | 0 |
| Vascular disorders | 0 | 1 (0.7) | 0 |
| Diabetic vascular disorder | 0 | 1 (0.7) | 0 |
| Venous thromboembolism | 0 | 0 | 0 |
| Death | 1 (0.7) | 0 | 0 |

All AEs were collected from the signature of the informed consent form until the last visit of the subject in the study (up to 22 weeks after the final dose of study treatment) regardless of relationship to study treatment, thus up to approximately 44 weeks.

A TAE is defined as an AE that first occurred or worsened in severity after the first dose of the study treatment.

%, percentage of subjects calculated relative to the total number of subjects in the population.

MedDRA (Medical Dictionary for Regulatory Activities, V.21.1) was used to code AEs.

AE, adverse event; ALT, alanine aminotransferase; AST, aspartate aminotransferase; n, number of subjects with events; N, number of subjects; OKZ, olokizumab; PBO, placebo; TEAE, treatment-emergent adverse event; TESAE, treatment-emergent serious adverse event.

in this patient population and investigational setting, 89 (62.2%) and 94 (66.2%) patients treated with OKZ had improvement in their HAQ-DI score with more than minimally detectable difference of 0.22, compared with 63 (47.6%) in the PBO group.

Chronic arthritis can have a profound effect on patients' quality of life.20 In this study it was shown that the improvements seen in signs and symptoms and disability of RA were mirrored by positive effects on quality of life measures including SF-36 and FACIT-F. SF-36 is a multidomain questionnaire that assesses different aspects of a person's life, summarised into PCS and MCS. Treatment with OKZ resulted in improvements across all of these domains (table 3). Certain mental ailments such as sleep disorders and fatigue in RA may be linked to high levels of circulating IL-6.21,22 OKZ treatment resulted in marked improvements in fatigue, consistent with its mechanism of action as an inhibitor of IL-6.
CREDO 1 trial also evaluated the safety of OKZ over 24 weeks and confirmed that OKZ has a safety profile similar to approved anti-IL-6R antagonists and no unexpected safety findings.23 24

As expected, there were more adverse events observed in the OKZ-treated patients, but they were mostly mild to moderate with few serious adverse events and no unexpected safety findings and relatively low number of dropouts due to an adverse event. In this relatively small study few serious infections, including opportunistic infection (pulmonary TB) and one fatal event, were reported for OKZ every 2 weeks and none for OKZ every 4 weeks.

There are several limitations to the study. First, there was no active comparator in this study, limiting the ability to compare with other agents. Second, the study did not include radiographic assessments. An analysis of RA trials of anti-TNF biologics showed a trend towards decreasing rate of radiographic progression, possibly due to more effective patient management, and to reliably show a positive radiographic effect one must include large numbers of patients on PBO, a possible ethical issue.25 Third, this study was conducted in a limited geographical location with limited racial diversity and its findings should be confirmed in other phase III controlled trials that include a more diverse patient population.
CONCLUSION

In this first phase III trial of OKZ in patients with active RA treatment with an adequate dose of MTX, OKZ demonstrated significant improvements in signs and symptoms of RA, including in disability and quality of life measures, compared to the RCT initiation, the study protocol was approved by the ethics committee and conducted in accordance with CONSORT 2010 Statement: updated guidelines for reporting parallel-group randomised trials (https://www.bmj.com/content/340/bmj.c332) and Good Publication Practice guidelines (http://annals.org/aim/article/2424869).

Contributors

CJSC R-Pharm was involved in the study design, collection and analysis, interpretation of data, and checking of information provided in the manuscript. MS, SF and KE were involved with study conceptualisation and conducted the data analysis. All authors had unrestricted access to study data and contributed to the interpretation of the results. All authors were responsible for all content and editorial decisions.

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Competing interests

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Patient consent for publication

Not required.

Ethics approval

The study was reviewed by the Ethics Council of the Ministry of Health of the Russian Federation (extract of minutes #122, date: 5 April 2016). Prior to the RCT initiation, the study protocol was approved by the ethics committee and regulatory authorities. The study was conducted in accordance with the ICH GCP and the Declaration of Helsinki requirements.

Provenance and peer review

Not commissioned; externally peer reviewed.

Data availability statement

Data are available upon reasonable request. All data relevant to the study are included in the article or uploaded as supplementary information. Data in the form of pdf files can be provided upon reasonable request. The requests should be submitted to Sofia Kuzkina at R-Pharm International at the following email address: kuzkina@rpharm.ru.

Supplemental material

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ORCID iDs

Evgenei Nasonov http://orcid.org/0000-0002-1598-8360
Saeed Falnejad http://orcid.org/0000-0002-3005-9439
Mark Genovese http://orcid.org/0000-0001-5294-4503

REFERENCES

1 Singh JA, Saag KG, Bridges SL, et al. 2015 American College of rheumatology guideline for the treatment of rheumatoid arthritis. Arthritis Rheumatol 2016;68:1–26.
2 McInnes IB, Schett G. The pathogenesis of rheumatoid arthritis. N Engl J Med 2011;365:2205–19.
3 Smolen JS, Landewé RBM, Bijlmer JW, et al. EULAR recommendations for the management of rheumatoid arthritis with synthetic and biological disease-modifying antirheumatic drugs: 2019 update. Ann Rheum Dis 2020;79:685–99.
4 Winthrop KL, Weinblatt ME, Bathon J, et al. Unmet need in rheumatology: reports from the targeted therapies meeting 2019. Ann Rheum Dis 2020;79:88–93.
5 Dayer J-M, Choy E. Therapeutic targets in rheumatoid arthritis: the interleukin-6 receptor: Rheumatology 2010;49:15–24.
6 Hunter CA, Jones SA, IL-6 as a keystone cytokine in health and disease. Nat Immunol 2015;16:448–57.
7 Nakashima Y, Kondo M, Harada H, et al. Clinical evaluation of tocilizumab for patients with active rheumatoid arthritis refractory to anti-TNF biologics: tocilizumab in combination with methotrexate. Mod Rheumatol 2010;20:343–52.
8 Genovese MC, Fleischmann R, Kivitz AJ, et al. Sandumab plus methotrexate in patients with active rheumatoid arthritis and inadequate response to methotrexate: results of a phase III study. Mod Rheumatol 2015;25:1424–37.
9 Genovese MC, McKay JD, Nasonov EL, et al. Interleukin-6 receptor inhibitor with tocilizumab reduces disease activity in rheumatoid arthritis with inadequate response to disease-modifying antirheumatic drugs: the tocilizumab in combination with traditional disease-modifying antirheumatic drug therapy study. Arthritis Rheum 2008;58:2968–80.
10 Shaw S, Broume T, Meier C, et al. Discovery and characterization of olokizumab: a humanized antibody targeting interleukin-6 and neutralizing gp130-signaling. Mabs 2014;6:774–82.
11 Kretos K, Golor G, Jullion A, et al. Safety and pharmacokinetics of olokizumab, an anti-IL-6 monoclonal antibody, administered to healthy male volunteers: a randomized phase I study. Clin Pharmacol Drug Dev 2014;3:388–95.
12 Genovese MC, Fleischmann R, Furst D, et al. Efficacy and safety of olokizumab in patients with rheumatoid arthritis with an inadequate response to TNF inhibitor therapy: outcomes of a randomised phase IIb study. Ann Rheum Dis 2014;73:1607–15.
13 Takeuchi T, Tanaka Y, Yanamaka H, et al. Efficacy and safety of olokizumab in Asian patients with moderate-to-severe rheumatoid arthritis, previously exposed to anti-TNF therapy: results from a randomized phase II trial. Mod Rheumatol 2016;26:15–23.
14 Aleatah D, Neogi T, Silman AJ, et al. 2010 rheumatoid arthritis classification criteria: an American College of Rheumatology/European League against rheumatism collaborative initiative. Arthritis Rheum 2010;62:2569–81.
15 Rubin D. Multiple Imputation for Nonresponse in Surveys. Hoboken, NJ, USA: John Wiley & Sons, Inc. 1987.
16 Yazici Y, Curtis JR, Ince A, et al. Efficacy of tocilizumab in patients with moderate to severe active rheumatoid arthritis and a previous inadequate response to disease-modifying antirheumatic drugs: the rosy study. Ann Rheum Dis 2012;71:198–205.
17 Orabi A-M, Bingham CO. Patient reported outcomes in rheumatoid arthritis clinical trials. Curr Rheumatol Rep 2015;17:1–10.
18 Barber CEH, Zell J, Yazdany J, et al. 2019 American College of rheumatology recommended patient-reported functional status assessment measures in rheumatoid arthritis. *Arthritis Care Res* 2019;71:1531–9.

19 Kingsley G, Scott IC, Scott DL. Quality of life and the outcome of established rheumatoid arthritis. *Best Pract Res Clin Rheumatol* 2011;25:585–606.

20 Gilbert AL, Lee J, Ehrlich-Jones L, et al. A randomized trial of a motivational interviewing intervention to increase lifestyle physical activity and improve self-reported function in adults with arthritis. *Semin Arthritis Rheum* 2018;48:732–40.

21 Choy EHS, Calabrese LH. Neuroendocrine and neurophysiological effects of interleukin 6 in rheumatoid arthritis. *Rheumatology* 2018;57:1885–95.

22 Mihara M, Hashizume M, Yoshida H, et al. IL-6/IL-6 receptor system and its role in physiological and pathological conditions. *Clin Sci* 2012;122:143–59.

23 Genovese MC, van der Heijde D, Lin Y, et al. Long-Term safety and efficacy of sarilumab plus methotrexate on disease activity, physical function and radiographic progression: 5 years of sarilumab plus methotrexate treatment. *RMD Open* 2019;5:e000887.

24 Hushaw LL, Sawaqde R, Sweis G. Critical appraisal of tocilizumab in the treatment of moderate to severe rheumatoid arthritis. *Ther Clin Risk Manag* 2010;6:143–52.

25 Rahman MU, Buchanan J, Doyle MK, et al. Changes in patient characteristics in anti-tumour necrosis factor clinical trials for rheumatoid arthritis: results of an analysis of the literature over the past 16 years. *Ann Rheum Dis* 2011;70:1631–40.