Effectiveness of Golimumab in Clinical Management of Patients with Rheumatoid Arthritis

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Abstract

Background and Objectives  Limited data are available regarding the use of golimumab (100 mg) every 4 weeks, with or without methotrexate (MTX). The aim of this retrospective analysis was to evaluate the effectiveness and safety of golimumab following usual clinical practice in Japanese patients with rheumatoid arthritis (RA) according to the recommendations given in the Japanese package insert.

Patients and Methods  Japanese RA patients with moderate-to-high disease activity, according to the 28-joint disease activity score based on C-reactive protein (DAS28-CRP) criteria, despite treatment with MTX or another biological agent, were enrolled. Patients were assigned to 50 mg golimumab plus MTX or 100 mg golimumab monotherapy every 4 weeks for 24 weeks. All patients were given MTX if it was not contraindicated. The primary endpoint was the proportion of patients achieving clinical remission (defined as a DAS28-CRP <2.3 or a simplified disease activity index [SDAI] score <3.3) at 24 weeks.

Results  Most patients received combined 50 mg golimumab plus MTX (41/43). In these patients, the primary endpoint, clinical remission, was attained in 83% of patients according to DAS28-CRP criteria (p < 0.001) and 69% according to SDAI criteria (p < 0.001) by week 24. Adverse events were reported in 11.6% of patients receiving golimumab.

Conclusions  Golimumab (50 mg) plus MTX effectively reduced the signs and symptoms of RA and was generally well tolerated in patients with an inadequate response to MTX and other biological agents.

1 Introduction

In recent years, methotrexate (MTX) therapy at high dose levels and tumor necrosis factor (TNF) inhibitor therapy have been applied to treatment of rheumatoid arthritis (RA). Anti-TNF therapy, either alone or in combination with MTX (apart from infliximab, which should only be used in combination with MTX), is recommended in patients with active RA with inadequate response to MTX or another disease-modifying antirheumatic drug (DMARD) or combination of DMARDs or another anti-TNF agent [1–3]. These new methods of treatment are expected to yield not only the alleviation of disease activity, but also structural improvement of the affected joints and improvement in daily life for patients. The three most widely used anti-TNF agents in Japan are infliximab, etanercept, and adalimumab, and numerous reports have been published on these agents [4–6].

Golimumab (GLM), a new human anti-TNF antibody agent created using transgenic mice, has been shown to exert effectiveness comparable to that of existing anti-TNF antibody agents when injected subcutaneously at 4-week intervals [7–13]. This drug was introduced in Japan in September 2011, thus providing a new treatment option for Japanese patients with RA. GLM can be administered either as monotherapy at a dosage of 100 mg or in combination with MTX at dosages of 50 or 100 mg every 4 weeks [14]. It is indicated not only in patients who have not previously received treatment with biological agents but also in patients who have experienced difficulties with infliximab or adalimumab therapy; for example, problems with neutralizing antibodies.

In Japan, there have been no published reports on the use of GLM in clinical practice to date. When patients are enrolled into clinical studies, age and disease activity are
often taken into account to ensure safety and continued use of the investigational agent, so the populations studied differ from the population managed in real life. Therefore, this analysis evaluates the use of GLM in patients with RA receiving real-life clinical care at our clinic.

2 Methods

2.1 Subjects

This retrospective analysis included patients with baseline moderate-to-high disease activity according to a 28-joint disease activity score based on C-reactive protein (DAS28-CRP) >3.2 despite treatment with MTX or another biological agent. Patients had RA according to American College of Rheumatology (ACR) criteria [15] and had received GLM treatment as outpatients at our hospital, the Shono Rheumatology Clinic, Fukuoka, Japan, between September 2011 and April 2012. Consent in writing was obtained from each patient in advance.

2.2 Treatment

Patients received combination therapy with GLM plus MTX, with GLM administered at a dose of 50 mg or 100 mg every 4 weeks plus MTX administered at a dose of up to 8 mg/week; or GLM monotherapy, with GLM administered at 100 mg every 4 weeks, for a total of 24 weeks. All patients were prescribed MTX if it was not contraindicated. GLM was administered subcutaneously in accordance with the Japanese package insert [14].

2.3 Outcome Measures

The primary endpoint of this retrospective analysis of effectiveness was to evaluate the proportion of patients achieving remission defined as a DAS28-CRP <2.3 or a simplified disease activity index (SDAI) score <3.3. Mean changes in the DAS28-CRP from baseline to 4 weeks were also evaluated.

Safety was evaluated on the basis of adverse events and laboratory test data. For each parameter, additional stratified analyses were conducted, dividing the patients into two groups; that is, bio-naive patients who had not received biological agents prior to receiving GLM, and patients who had received prior biological agents (i.e., those switching from other biological agents to GLM).

2.4 Statistical Analysis

All data were included for efficacy and safety analyses. The last observation carried forward (LOCF) method was used to allow for missing data. Comparison of groups was performed using the Student’s t test with statistical significance set at $p < 0.05$.

3 Results

3.1 Patient Baseline Demographics and Clinical Characteristics

Of all patients studied, 18 were bio-naive cases and 25 had received prior biological agents, including infliximab ($n = 4$), etanercept ($n = 10$), adalimumab ($n = 6$), and tocilizumab ($n = 5$). Of the 25 patients previously treated with biological agents, 19 had received one prior biological agent and 6 had received two or more agents.

Table 1 shows the baseline demographics and disease characteristics of the patients enrolled into the study. Patient characteristics were generally well balanced between bio-naive patients and those who had received a prior biological agent, except the proportion of women was slightly greater (96.0 vs 83.3 %) and disease duration was slightly longer (122.6 vs 105.3 months) in the bio-switching group.

3.2 Interventions

In total, 41 patients received GLM at a dose of 50 mg every 4 weeks in combination with MTX (mean dose 6.23 mg/week) and 2 patients received GLM monotherapy at a dose of 100 mg every 4 weeks. Four patients were unsatisfied with the inconvenience of self-injection and injection pain of prior biological treatment, despite sufficient clinical response; therefore, those patients in clinical remission at baseline were switched to GLM treatment as a result of patients’ wishes. Of the 43 patients, 35 completed the 24-week treatment period.

3.3 Effectiveness

Remission rates, defined as the proportion of patients achieving a DAS28-CRP <2.3 and an SDAI score <3.3, steadily improved over the course of treatment with GLM (Fig. 1). After 8 weeks of treatment, 71.4 % of patients achieved a DAS28-CRP <2.3 and 50.0 % achieved an SDAI score <3.3. After 8 weeks of treatment, the DAS28-CRP and SDAI remission rates were higher in patients who had not received prior biological agents versus those who had (55.6 vs 50.0 % and 61.1 vs 41.7 %, respectively).

The mean DAS28-CRP 4 weeks after the start of treatment was significantly improved compared with the pretreatment score [mean DAS28-CRP at week 4 = 1.80 vs 4.14 (range 1.28–7.04) at baseline; $p < 0.001$].
Improvements in DAS28-CRP and SDAI scores throughout the treatment period were similar in bio-naïve patients and those who had received prior biological agents (Figs. 2, 3). Changes in DAS28-CRP and SDAI scores at 4 and 8 weeks were statistically significant compared with baseline in both bio-naïve patients and those who had received prior biological agents (all \( p < 0.001 \)).

### 3.4 Tolerability

GLM was generally well tolerated with no unexpected safety issues observed. Adverse events (shown in Table 2) were reported in five patients, most of whom were receiving GLM (50 mg) in combination with MTX (6 or 8 mg). Two patients reported fractures (one ankle and one femur); one patient was hospitalized due to renal impairment, chest pain, dyspnea, bronchial asthma, acute upper respiratory tract inflammation, and bronchitis; one patient (treated with GLM monotherapy at 100 mg) experienced venous thromboembolism and lower limb edema; and one patient reported renal impairment, hepatic function, and nephrogenic anemia. Consistent with other GLM safety data reported in Japanese clinical trials, no unknown adverse event was reported in this clinical analysis. All adverse events were resolved with treatment.

### Table 1 Baseline demographics and disease characteristics in bio-naïve patients and patients who had received prior biological agents

|                          | Total (\( n = 43 \)) | Bio-naïve (\( n = 18 \)) | Prior biologicals (\( n = 25 \)) |
|--------------------------|-----------------------|--------------------------|----------------------------------|
| **Sex** [\( n (\% \)] ]  |                       |                          |                                  |
| Female                   | 39 (90.7)             | 15 (83.3)                | 24 (96.0)                        |
| Male                     | 4 (9.3)               | 3 (16.7)                 | 1 (4.0)                          |
| **Age [years]**          | 59.1 (32–79)          | 55.8 (37–79)             | 61.4 (32–76)                     |
| **Disease duration [months]** | 115.3 (7–708)       | 105.3 (7–708)            | 122.6 (12–252)                  |
| **DAS28-CRP**            | 4.14 (1.28–7.04)      | 4.16 (2.61–6.39)         | 4.12 (1.28–7.04)                |
| **SDAI**                 | 22.2 (2.81–62.30)     | 22.30 (6.70–56.29)       | 22.20 (2.81–62.30)              |
| **CDAI**                 | 20.92 (2.50–60.90)    | 20.94 (6.50–56.00)       | 20.90 (2.50–60.90)              |
| **Tender joint count [0–68]** | 6.3 (0–24)           | 6.7 (1–22)               | 6.0 (0–24)                       |
| **Swollen joint count [0–68]** | 5.9 (0–22)           | 5.4 (0–18)               | 4.8 (0–22)                       |
| **mHAQ [0–24]**          | 0.65 (0–2)            | 0.44 (0–2)               | 0.72 (0–2)                       |
| **CRP [mg/dL]**          | 1.5 (0.1–13.5)        | 1.6 (0.1–13.5)           | 1.4 (0.1–8.4)                    |
| **RF positive [\( n (\% \)] ]** | 34 (79.0)           | 13 (72.2)                | 21 (84.0)                        |
| **ACPA positive [\( n (\% \)] ]** | 22 (51.1)           | 11 (61.1)                | 11 (44.0)                        |

All values are presented as means with ranges given in parentheses unless specified otherwise.

**ACPA** anti-citrullinated protein autoantibody, **CDAI** clinical disease activity index, **CRP** C-reactive protein, **DAS28-CRP** disease activity score 28 based on C-reactive protein, **mHAQ** modified health assessment questionnaire, **RF** rheumatoid factor, **SDAI** simplified disease activity index.

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**Fig. 1** Remission rate in 43 patients with rheumatoid arthritis treated with golimumab alone or in combination with methotrexate. Remission was defined as a 28-joint disease activity score based on C-reactive protein (DAS28-CRP) of <2.3 or a simplified disease activity index (SDAI) score of <3.3. DAS28-CRP remission and DAS28-CRP plus SDAI remission (ALL) are shown. BL baseline, W weeks

Improvements in DAS28-CRP and SDAI scores throughout the treatment period were similar in bio-naïve patients and those who had received prior biological agents (Figs. 2, 3). Changes in DAS28-CRP and SDAI scores at 4 and 8 weeks were statistically significant compared with baseline in both bio-naïve patients and those who had received prior biological agents (all \( p < 0.001 \)).

**Fig. 2** Changes in mean 28-joint disease activity score based on C-reactive protein (DAS28-CRP) in bio-naïve or previously treated patients with rheumatoid arthritis receiving golimumab alone or in combination with methotrexate. The dashed line represents the defined remission cutoff value of 2.3. BL baseline, W weeks

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Discussion

The present analysis in Japanese patients with RA in real-life clinical care revealed high effectiveness and safety of GLM alone or in combination with MTX, with significant improvements in mean DAS28-CRP and SDAI scores observed in bio-naive patients 16 weeks after the start of treatment ($p < 0.001$). The reason for the high remission rate was considered to be the difference in average patient body weight between western countries and Japan (75 vs 50 kg, respectively).

These effectiveness data are consistent with efficacy data from clinical studies [7–10, 12, 13, 16]. Most GLM studies are designed to permit rescue of patients at 16 weeks with alternative pharmacological therapy for those meeting the nonresponse criteria for early escape [8–10, 12, 13].

Similar to the GO-FORTH study [13], our clinical analysis involved patients treated with MTX at 8 mg/week, which is the maximum dose approved in Japan at the time that the patients were receiving treatment [17]. This is lower than the current recommended MTX dose in RA [3, 14, 18] and lower than the MTX dose used in combination with GLM in other published studies [7, 9, 10]. Despite the low doses of MTX used, overall remission rates with GLM were high. Evidence suggests that, for patients receiving MTX who fail to achieve clinical remission based on SDAI or disease activity score scales, increasing the MTX dose needs to be considered at this time [17]. In the GO-FORWARD study, GLM was shown to be effective in patients who showed lower responses or who were refractory to prior MTX therapy [9, 10].

In the present retrospective analysis, manifestation of effectiveness appeared to be delayed in the bio-switching group compared with the bio-naive group, suggesting the necessity for longer follow-up when evaluating effectiveness in patients who switch between biological therapies.

In a post-hoc analysis of the effectiveness in relation to the reasons for switching, the effectiveness did not differ significantly according to the reason (data not shown). This suggests that patients undergoing switching will respond to this therapy, regardless of the reasons for switching. This supports findings by Smolen et al. [12] that switching from other anti-TNF agents to GLM was effective regardless of the reasons for switching, indicating that GLM can serve as the second anti-TNF agent when patients are switched from another TNF agent. Of the five anti-TNF agents available, including certolizumab pegol, all have different affinities to TNF-α; therefore, switching from one anti-TNF agent to another is likely to be effective.

Expression of antibodies to anti-TNF antibody agents such as infliximab, adalimumab, and certolizumab pegol monotherapy is not uncommon; however, incidences of anti-GLM antibodies in the GO-FORWARD [9] and GO-FORTH [13] studies were remarkably low. Because GLM is prepared by the transgenic mouse technique, it is an antibody with high affinity for the antigen [19], which means that formation of unstable proteins or aggregations, which can serve as immunogens, is unlikely. Studies of GLM (100 mg) monotherapy were conducted in Caucasian and South American countries in GO-FORWARD [9, 10] and in Japan in GO-FORTH [13] and GO-MONO [16], and showed that GLM is an appropriate biological agent for preventing the loss of effectiveness in Caucasian, South American, and Japanese populations receiving long-term RA treatment [9, 10, 13, 16]. As a result of findings from the GO-FORTH [13] and GO-MONO [16] studies, in addition to the 50-mg dose, GLM (100 mg) every 4 weeks—as monotherapy and in combination with MTX—has been approved in Japan. Further studies at this dose level in larger numbers of patients are necessary.

Apart from the usual limitations relating to observational data and retrospective analyses, particularly with
regard to selection and enrolment bias, a major limitation of our analysis is the small patient numbers, especially for patients receiving GLM (100 mg) monotherapy. In addition, evaluation of levels of anti-GLM antibodies and the effects of GLM on structural joint damage in this real-life setting would have been useful; however, this was not evaluated in the original study.

Despite suppression of joint destruction being an important factor in improving the quality of life of patients with RA, the number of published reports evaluating this effect in patients treated with GLM is small, because of its relatively recent development. Emery et al. [7] reported that treatment with GLM suppressed joint destruction significantly 52 weeks after the start of treatment, and further long-term observation is needed. However, due to the short follow-up period in our analysis, such observation was not possible.

In the present analysis, there were no serious adverse events arising from the use of GLM, although deterioration in renal function was reported in two patients. An association with the development of malignant tumors has been suggested with GLM, and further clinical confirmation is warranted [20]. However, long-term observation of the patients in our study is needed before any definite conclusions can be made.

It is important to select a type of biological agent taking into account the lifestyle of individual patients. Despite reported problems with pain and administration site reactions, subcutaneous injection of drugs offers greater convenience than intravenous infusion, which requires physical immobilization for many hours at a hospital, and a longer dosing interval is also advantageous. Because GLM contains only small amounts of stimulating acidic additives and requires only a small volume of dosing solution, reported incidences of pain and administration site reactions are low [14].

5 Conclusion

In the present analysis, GLM plus MTX or GLM monotherapy used in clinical practice in Japanese patients with RA was confirmed to have high effectiveness and safety, comparable with existing biological agents. Thus, we conclude that GLM is a promising new alternative for the treatment of RA in Japanese patients showing poor response, those in whom the use of other biological agents is contraindicated, and cases where the use of MTX in combination with biological agents is difficult.

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Conflict of Interest The author has no conflicts of interest to declare.

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