Considering new lessons about the use of IL-6 inhibitors in arthritis

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ABSTRACT
Interleukin (IL)−6 represents one of several possible targets for the treatment of rheumatoid arthritis. Drugs targeting IL-6 can be divided into monoclonal antibodies against IL-6 itself and monoclonal antibodies against the IL-6 receptor. Both types of agent inhibit both classical signalling through membrane-bound IL-6 receptor, and trans-signalling via formation of a complex between IL-6 and soluble IL-6 receptor. The IL-6 receptor blockers tocilizumab and sarilumab inhibit the low affinity binding of IL-6 to its receptor. The anti-IL-6 agents clazakizumab and vobarilizumab also block binding of IL-6 to the receptor, while olokizumab blocks the higher affinity interaction of the IL-6-receptor complex with gp130. The doses and dosing intervals of the biologics targeting different elements vary, but no major differences in efficacy or safety have yet been seen between the two approaches, although more studies are needed in this area. In addition to the different blocking actions of monoclonal antibodies, we consider therapeutic strategies including the timing of IL-6 blockade and the use of monotherapy versus the addition of methotrexate.

A panel of international experts in the field of rheumatology recently came together to consider the mechanisms by which the IL-6 pathway is targeted.

CONSIDERING TARGET MOLECULES IN RHEUMATOID ARTHRITIS
Several molecules have been targeted in the treatment of rheumatoid arthritis. For tumour necrosis factor (TNF)α, four drugs targeting the ligand (adalimumab, certolizumab pegol, golimumab, and infliximab) and one targeting the receptor (etanercept) have been approved. In the case of IL-6, tocilizumab and sarilumab (both approved) target the receptor and four drugs that target the ligand have results from phase II or III clinical trials (sirukumab, olokizumab, clazakizumab, and vobarilizumab). Following a negative review from the FDA, sirukumab development and registration has been stopped. An additional modality based on the selective blockade of IL-6 trans-signalling is in early stage clinical trial development (olamkicept). In addition, abatacept targeting CD80/86, rituximab targeting CD20, and anakinra targeting the IL-18 receptor are approved, and mavrilimumab targeting the granulocyte macrophage colony stimulating factor (GM-CSF) receptor has positive trial results. Proof of concept has also been demonstrated for several other targets, including IL-18 ligand, IL-12/23 (p40), IL-23 (p19), IL-17 ligand, and B cell activating factor, although benefits have so far been either marginal or insufficient to merit progression to general use.

As well as pro-inflammatory cytokines, such as TNFα, IL-6, and GM-CSF, B cells and T cells play an important role in the pathogenesis of RA.1–3 Activated B cells and T cells work together to interact with synovial cells to produce a large number of inflammatory mediators, including pro-inflammatory cytokines (TNFα and IL-6), induce matrix metalloproteinases, promote synovial fibroblast proliferation, and induce osteoclast differentiation.4 This hypothesis is corroborated in a substantial number of clinical trials that show the clinical efficacy of abatacept, an inhibitor of co-stimulatory moieties on T cells. Inflammation is believed to result from or be perpetuated by autoantibodies such as anti-citrullinated peptide antibody and rheumatoid factors. B cells are thought to be responsible for the production of these autoantibodies. The pathogenic role of CD20+ B cells has been validated by the efficacy of rituximab, a monoclonal antibody against CD20, in clinical trials.

The role of pro-inflammatory cytokines varies from one disease to another. This has given rise to the notion of context-dependent role for individual cytokines in chronic disease progression. Consequently, the cytokine network established in these conditions varies considerably and implies that some clinical conditions are more susceptible to intervention with specific biological drugs than others. For example, biological drugs that target TNFα, IL-6, and GM-CSF are more efficacious in treating rheumatoid arthritis than blockers of the IL-17 or IL-23 pathway.1 In contrast, inhibition of IL-17 and IL-23 is more appropriate for the treatment of psoriasis.3

Two decades ago, it was discovered that IL-6 concentrations were decreased in 80%–90% of patients with rheumatoid arthritis 2 hours after an infusion of infliximab, suggesting that IL-6 is immediately downstream of TNFα in the cytokine cascade in rheumatoid arthritis.1 In addition, methotrexate acts to further inhibit IL-6.6–8 Blocking TNFα reduces the signalling through the IL-6 receptor, which leads to decreased production of other molecules such as soluble intercellular adhesion molecule-1 and vascular endothelial growth factor.

CONSIDERING SIGNAL TRANSDUCTION THROUGH IL-6 RECEPTOR
When IL-6 binds to membrane-bound IL-6 receptor, the signal is transmitted through gp130 on the cell membrane [classical signalling]. In trans-signalling, IL-6 binds to soluble IL-6 receptor (sIL-6R) and the resulting IL-6/sIL-6R complex combines with gp130 to transmit the signal (figure 1).9 Classical signalling via membrane-bound IL-6 receptor occurs only in the haematopoietic system, whereas trans-signalling via sIL-6R can occur in any type of cell. The binding of IL-6.
Review


Figure 1  Signal transduction through IL-6R.9

to its receptor is of much lower affinity than the binding of the IL-6/sIL-6R complex to gp130, meaning that the former is easier to block.

Agents that target the IL-6 receptor, such as tocilizumab and sarilumab, inhibit both IL-6 classical and trans-signalling, as do agents such as clazakizumab and olokizumab that target the IL-6 ligand.10 Tocilizumab and sarilumab act on domain 2 of the IL-6 receptor to inhibit binding of IL-6 without interfering with binding to gp130 at domain 3 (figure 2).11 Monoclonal antibodies against the IL-6 molecule may block the binding of IL-6 receptor at site 1, as is the case for clazakizumab, or interfere with the fully-functioning receptor complex by blocking the interaction with gp130 at site 2 or site 3, as with olokizumab [site 3] (figure 2). These different blocking actions result in differences in efficacy and pharmacokinetics; IL-6 inhibitors that act at site one cause more pronounced increases in systemic IL-6 levels than do inhibitors that bind site 2 or site 3.11

CONSIDERING THE DETERMINANTS OF EFFICACY OF BIOLOGICS

One key factor determining the efficacy of biologics is the appropriateness of the target, and both IL-6 and the IL-6 receptor are appropriate targets.
in rheumatoid arthritis. Another important determining factor is trough concentration of the drug (that is, the concentration immediately before the next dose), which should be above the minimum efficacious concentration. The trough concentration is affected by the amount of target molecule, the dose and dosing interval of the biologic, the affinity of Fc receptors for clearing antibodies, and the presence of antidrug antibodies. Other efficacy determining factors include adverse drug reactions and transfer to the site of inflammation.

In a cohort of patients with rheumatoid arthritis (n=61) at Keio University School of Medicine, Tokyo, baseline sIL-6R concentrations were shown to predict DAS-28(REMD) remission at week 24 of tocilizumab treatment.12 Significantly more patients with an sIL-6R concentration below 72.6 ng/mL achieved remission compared with those with higher sIL-6R. In patients with rheumatoid arthritis, sIL-6R concentration is slightly higher than in healthy people, whereas the IL-6 concentration is much higher in patients with rheumatoid arthritis than in healthy people.9 However, even though the relative magnitude of IL-6 in patients with rheumatoid arthritis versus healthy people is greater than for sIL-6R, the IL-6 concentration is 10–100 times lower than sIL-6R concentration (5–500 pg/mL v 40–800 mg/mL). This means that it is much more difficult to neutralise sIL-6R with a given amount of a biologic than it is to neutralise IL-6. In general, less monoclonal antibody is needed when the target is IL-6 than when the IL-6 receptor is targeted, and less is needed when targeting IL-6 site 1 than site 3 (table 1). Appropriate selection of dose and dosing interval means that similar American College of Rheumatology 20% improvement criteria [ACR20] responses have been achieved at 12 weeks with clazakizumab, which acts at site 1, and with olokizumab acting at site 3.14–16

We suggest that drugs’ pharmacokinetics and pharmacodynamics may be important in determining whether patients who do not respond well or do not tolerate one biologic may go on to respond to another drug targeting the same cytokine, its receptor, or another molecule involved in the same pathway. For example, in clinical practice, the efficacy of intravenous tocilizumab (8 mg/kg) can be adjusted by shortening or extending intervals of regular 4-week administration in patients with rheumatoid arthritis.17

### CONSIDERING REGULATORY T CELLS

A study of 39 patients with rheumatoid arthritis who received tocilizumab as their first biologic found that the proportion of regulatory T cells increased over 52 weeks of treatment, and the change in these cells was inversely correlated with the change in Clinical Disease Activity Index (CDAI) score [figure 3]. The greatest change in the proportion of regulatory T cells was seen in patients who achieved CDAI remission at 52 weeks.18 This implies that IL-6 receptor blockade leads to an increase in regulatory T cells, which is associated with maintenance of well-controlled disease from 6 to 12 months. Whether the same is true for agents targeting the IL-6 ligand is not known. However, IL-6 is known to inhibit the action of transforming growth factor β in inducing regulatory T cells and also plays a role in the production of naturally occurring regulatory T cells,19 so blocking IL-6 should also have this effect. Research is needed to confirm the clinical expansion of regulatory T cell populations during IL-6 blocking or IL-6R intervention therapy.

### CONSIDERING THERAPEUTIC STRATEGIES IN RHEUMATOID ARTHRITIS

#### The timing of the introduction of IL-6 blockade

Looking at the Disease Activity Score-28 (DAS28) remission rate at 24 weeks in the tocilizumab monotherapy arm of various studies, U-Act-Early stands out with a rate of 76% compared with 49% in SATORI, 40% in ADACTA, 39% in FUNCTION, and 55% in SURPRISE.13–20–23 This may be because IL-6 receptor blockade with tocilizumab was introduced as early as 4 weeks’ disease duration in U-Act-Early compared with 0.5–8.5 years in the other studies. The baseline DAS28 was lower in U-Act-Early than in the other studies, and no joint damage was present.20 This suggests that earlier introduction of an IL-6 receptor inhibitor leads to a higher DAS28 remission rate. Studies are needed to determine whether the same is true for IL-6 ligand inhibitors. Of note, ACR and European League Against Rheumatism (EULAR) do not regard DAS28 <2.6 as a surrogate of clinical remission but rather the ACR-EULAR remission definition; the limitations of using DAS28 to assess outcomes when applying drugs that directly target the acute phase response irrespective of clinical improvement are detailed in key manuscripts from Smolen et al.24 For example, IL-6/IL-6 receptor blocking agents may prevent the induction of acute phase reactants such as C reactive protein (CRP). As DAS28 is a composite score based partly on the measurement of CRP, the effect of these agents may be exaggerated if considering DAS28 alone. Furthermore, comparisons of agents with different mechanisms of action should be considered cautiously; the use of composite measures (e.g. CDART) that do not incorporate information on acute phase reactants may be more appropriate for this purpose.

#### Monotherapy versus combination with methotrexate

The latest EULAR recommendations for rheumatoid arthritis management state that if clinical targets such as remission are not achieved with methotrexate or another synthetic disease modifying antirheumatic drug (DMARD), addition of a biological drug or a Janus kinase inhibitor should be considered.27 However, if methotrexate is not tolerated and so biologic monotherapy

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**Table 1** Biologics against IL-6 and IL-6 receptors: dose and interval tested in phase II and III clinical trials

<table>
<thead>
<tr>
<th>Target molecules</th>
<th>Biologics</th>
<th>Antibody</th>
<th>Route of administration</th>
<th>Dose</th>
<th>Intervals</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-6R (D2)</td>
<td>Tocilizumab</td>
<td>Humanised</td>
<td>IV (8 mg/kg)</td>
<td>480 mg</td>
<td>4w</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>SC</td>
<td>162 mg</td>
<td>1–2w</td>
</tr>
<tr>
<td>IL-6R (D2)</td>
<td>Sarilumab</td>
<td>Human</td>
<td>SC</td>
<td>150 mg</td>
<td>2w</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>200 mg</td>
<td></td>
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<tr>
<td>IL-6 (site 1)</td>
<td>Sirukumab</td>
<td>Human</td>
<td>SC</td>
<td>50 mg</td>
<td>2–4w</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>100 mg</td>
<td></td>
</tr>
<tr>
<td>IL-6 (site 1)</td>
<td>Clazakizumab</td>
<td>Humanised</td>
<td>SC</td>
<td>25 mg</td>
<td>4w</td>
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<td>100 mg</td>
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<td></td>
<td>200 mg</td>
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<tr>
<td>IL-6 (site 3)</td>
<td>Olokizumab</td>
<td>Humanised</td>
<td>SC</td>
<td>60 mg</td>
<td>2–4w</td>
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<td>120 mg</td>
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<td></td>
<td></td>
<td>240 mg</td>
<td></td>
</tr>
<tr>
<td>IL-6/sIL-6R complex</td>
<td>Olamkicept</td>
<td>Human</td>
<td>IV</td>
<td>300 mg</td>
<td>2w</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>600 mg</td>
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</tbody>
</table>

IV, intravenous; SC, subcutaneous; w, weeks.
must be considered, significantly better outcomes have been shown with anti-IL-6 receptor monotherapy than with anti-TNF monotherapy. The SURPRISE study looked at tocilizumab monotherapy versus addition of tocilizumab to methotrexate in patients who did not respond adequately to methotrexate. The primary endpoint was DAS28(ESR) remission at 24 weeks, with DAS28(ESR) remission and change from baseline in modified total Sharp score (mTSS) at 52 weeks as secondary endpoints. At 24 weeks, the DAS28(ESR) remission rate was 69.6% in the add-on arm and 55.0% in the switch arm (P=0.028). However, by week 52 there was no significant difference between the arms (72.2% v 70.3%). Conversely, in the ACT-RAY study, which assessed the 1-year efficacy of tocilizumab plus methotrexate or placebo, a significant difference in mean DAS28 score was seen at week 24 (3.13 for switch v 2.88 for add-on; P=0.05), but the difference was not significant at week 52 (2.63 v 2.58; P=0.39).

In the SURPRISE study, change from baseline in mTSS did not differ significantly between the groups, but the rate of clinically relevant radiographic progression (CRRP; mTSS ≥3) tended to be higher in switch than in add-on patients (15% v 7%; P=0.07). Radiographic progression in the patients with CRRP was larger in the switch arm than in the add-on arm (9.0/year v 5.0/year; P=0.04). The difference between the groups in mean CRP concentration in the CRRP patients was significant for the first 24 weeks (1.56 for switch v 0.49 for add-on; P=0.001) but not for the following 28 weeks (0.10 v 0.04; P=0.1).

The results from the SURPRISE study suggest that adding tocilizumab to methotrexate achieves more rapid DAS28 remission than tocilizumab monotherapy and prevents structural damage. However, in terms of safety, the proportion of patients with at least one adverse event or serious adverse event was higher in the add-on arm (60.0% and 13.9%, respectively) than in the switch arm (45.0% and 8.1%, respectively). Gastrointestinal, hepatic, and respiratory adverse events were all higher in the add-on arm. In light of these findings in Japanese patients, adding tocilizumab to methotrexate for the first 6 months and then tapering the methotrexate to achieve tocilizumab monotherapy might be a useful strategy.
CONSIDERING TARGETING THE IL-6 PATHWAY—
A SUMMARY

The TNF-IL-6 cytokine cascade plays an important role in the pathogenesis of rheumatoid arthritis (figure 4). Provenance and peer review in the round table. Zeneca KK., Eli Lilly Japan KK, Novartis Pharma KK, Janssen Pharmaceutical KK, outside other from Taisho Toyama Pharmaceutical Co Ltd, Nipponkayaku Co Ltd, other from Astra Co, Ltd, Pfizer Japan Inc, grants from Takeda Pharmaceutical Co, Ltd, Teijin Pharma Ltd, and personal fees from Bristol–Myers KK, Chugai Pharmaceutical Co Ltd, Daiichi Sankyo Samsung, Sanofi, UCB during the conduct of the study; TT reports grants, personal fees Gilead, Glaxo, ILTOO, Janssen, Lilly, Medimmune, MSD, Novartis-Sandoz, Pfizer, Roche, submitted work; IM reports grants from Roche and Refereron, during the conduct of the study; grants and personal fees from Abbvie, AstraZeneca, Celgene, GSK, Janssen, Lilly, Novartis, Pfizer, Roche, grants and personal fees from Refereron, outside the submitted work; JS reports grants from Abbvie, Astra-Zeneca, Janssen, Lilly, MSD, Pfizer, Roche, personal fees from Abbvie, Amgen, Astra-Zeneca, Astro, BMS, Celgene, Celltrion, Chugai, Gilead, Glaxo, ILTOO, Janssen, Lilly, Medimmune, MSD, Novartis-Sandoz, Pfizer, Roche, Samsung, Sanofi, UCB during the conduct of the study; TT reports grants, personal fees and other from Astellas Pharma Inc, Abbvie GK, Mitsubishi Tanabe Pharma Co, grants and personal fees from Bristol–Myers KK, Chugai Pharmaceutical Co Ltd, Daiichi Sankyo Co Ltd, Pfizer Japan Inc, grants from Takeda Pharmaceutical Co Ltd, Teijin Pharma Ltd, Asahikasei Pharma Corp, Eisai Co Ltd, AYUMI Pharmaceutical Corporation, grants and other from Taisho Toyama Pharmaceutical Co Ltd, Nipponkayaku Co Ltd, other from AstraZeneca KK, Eli Lilly Japan KK, Novartis Pharma KK, Janssen Pharmaceutical KK, outside the submitted work; the participants/authors received personal fees for their participation in the round table.

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REFERENCES


