CLINICAL SCIENCE

Rituximab as therapy to induce remission after relapse in ANCA-associated vasculitis


Handling editor: Josef S Smolen

► Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10.1136/annrheumdis-2019-216863).

For numbered affiliations see end of article.

Correspondence to: Dr Rona M Smith, University of Cambridge, Cambridge, UK; ronasmith@doctors.net.uk

PM and DJ are joint senior authors.

Received 4 February 2020
Revised 15 April 2020
Accepted 29 April 2020
Published Online First 24 June 2020

ABSTRACT

Objectives Evaluation of rituximab and glucocorticoids as therapy to induce remission after relapse in ANCA-associated vasculitis (AAV) in a prospective observational cohort of patients enrolled into the induction phase of the RITAZAREM trial.

Methods Patients relapsing with granulomatosis with polyangiitis or microscopic polyangiitis were prospectively enrolled and received remission-induction therapy with rituximab (4×375 mg/m2) and a higher or lower dose glucocorticoid regimen, depending on physician choice: reducing from either 1 mg/kg/day or 0.5 mg/kg/day to 10 mg/day by 4 months. Patients in this cohort achieving remission were subsequently randomised to receive one of two regimens to prevent relapse.

Results 188 patients were studied: 95/188 (51%) men, median age 59 years (range 19–89), prior disease duration 5.0 years (range 0.4–34.5). 149/188 (79%) had previously received cyclophosphamide and 67/188 (36%) rituximab. 119/188 (63%) of relapses had at least one major disease activity item, and 54/188 (29%) received the higher dose glucocorticoid regimen. 171/188 (90%) patients now an established induction strategy in ANCA-associated vasculitis.

Key messages

What is already known about this subject?

► Rituximab is increasingly being used as a remission induction agent in ANCA-associated vasculitis.

What does this study add?

► This large prospective cohort provides further efficacy and safety data for the use of rituximab in patients specifically with relapsing disease.

How might this impact on clinical practice or future developments?

► Rituximab in conjunction with glucocorticoids is now an established induction strategy in ANCA-associated vasculitis.

INTRODUCTION

Granulomatosis with polyangiitis (GPA) and microscopic polyangiitis (MPA) are the major subgroups of antineutrophil cytoplasmic antibody (ANCa)-associated vasculitis (AAV). These conditions are characterised by leucocyte infiltration of blood vessel walls, fibrinoid necrosis and vascular damage and are usually associated with the presence of circulating ANCA.1 Prior to the availability of effective treatment, AAV had a mortality of 93% within 2 years, primarily due to renal and respiratory failure.2 The introduction of glucocorticoids and cyclophosphamide, which became established treatment for this disease in the 1980s, markedly improved survival, inducing remission at 1 year in approximately 80% of patients. However, relapsing disease is common with over 50% of patients experiencing a relapse within 5 years and the majority suffering treatment-related toxicity.1,3

B-lymphocytes have been implicated in the pathogenesis of AAV. Rituximab is a murine/human chimeric monoclonal antibody directed against
the CD20 antigen found on the surface of B-lymphocytes and results in B cell depletion. Rituximab was shown to be non-inferior to cyclophosphamide for induction of remission in AAV and superior to cyclophosphamide for the treatment of relapsing disease.6 7 Rituximab became a licenced therapy for remission induction of AAV in 2011.

Fixed-interval, repeat-dosing of rituximab was shown to be superior to azathioprine as a maintenance strategy following induction of remission cyclophosphamide in a trial of 117 patients with predominantly newly diagnosed AAV.8 The optimal strategy to maintain remission following induction of remission with rituximab, especially for treatment of relapse, is not clear. RITAZAREM was an international, randomised, controlled trial designed to assess whether rituximab is superior to azathioprine for the maintenance of remission following induction of remission with rituximab and glucocorticoids in patients with relapsing AAV. In this trial, fixed-interval, repeat doses of rituximab were compared with daily azathioprine for maintenance of remission.

Since all patients received rituximab for induction of remission in the RITAZAREM trial, this is the largest reported prospective cohort of patients with relapsing AAV to receive treatment with rituximab for induction of remission. This first report outlines the efficacy and safety of rituximab with either higher or lower dose glucocorticoids for induction of remission in a large prospective cohort of patients with relapsing AAV.

METHODS

The details of the RITAZAREM protocol have been previously published.9 In summary, RITAZAREM trial has three phases:

1. An *induction phase* (months 0–4): eligible patients enrolled at time of disease relapse received rituximab (4 weekly doses of 375 mg/m²) and glucocorticoids.
2. A *maintenance phase* (months 4–24): 4 months after enrolment, participants who achieved remission (defined as a Birmingham Vasculitis Activity Score for Wegener’s granulomatosis (BVAS/WG) ≤1 and prednisone/prednisolone dose ≤10 mg/day) were randomised in 1:1 ratio to receive 1000 mg rituximab at 4 monthly fixed intervals or daily azathioprine (2 mg/kg/day).
3. A *follow-up phase*: clinical follow-up after completion of therapy with either rituximab or azathioprine (minimum of 12, maximum of 24 months).

This paper reports on the first, induction phase of the trial, prior to randomisation.

Participants

Participants were aged over 15 years and had a diagnosis of GPA or MPA according to Chapel Hill Consensus Conference definitions10 and a current or historical positive test for PR3-ANCA or MPO-ANCA. All patients had disease relapse defined by one major or three minor disease activity items on the BVAS/WG and had previously achieved remission following at least 3 months of induction therapy, with a combination of glucocorticoids and an immunosuppressive agent (cyclophosphamide, rituximab, methotrexate or mycophenolate mofetil).

Key exclusion criteria included the receipt of any biological B cell depleting agents within the previous 6 months, alemtuzumab or antithymocyte globulin within the previous 12 months, or intravenously administered immunoglobulin, plasma exchange...
or anti-tumour necrosis factor (TNF) treatment within the previous 3 months. Patients with other multisystem autoimmune diseases, such as eosinophilic granulomatous with polyangiitis, systemic lupus erythematosus, antiglomerular basement membrane disease or cryoglobulinaemic vasculitis or history of malignancy within the past 5 years were also excluded.

Participants were recruited from 29 centres in seven countries.

Interventions and induction therapy

Rituximab

Rituximab 375 mg/m²/week was administered in four doses.

Glucocorticoids

Investigators chose from one of two glucocorticoid regimens taking into consideration disease severity and local prescribing practices. Schedule 1A had a glucocorticoid starting dose of 1 mg/kg/day (maximum 60 mg daily) and 1B had a starting dose of 0.5 mg/kg/day (maximum 30 mg daily), both tapering to 10 mg daily by month 4. Deviation from the protocol-specified tapering glucocorticoid regimen was defined as a 25% higher or lower glucocorticoid dose, averaged over 2 weeks. Patients could also receive a maximum cumulative dose of 3000 mg intravenous methylprednisolone, between 14 days prior to enrolment and 7 days after enrolment.

Other treatments

Prophylaxis to prevent pneumocystis (carinii) jiroveci infection and/or to prevent osteoporosis were recommended according to local practice. Plasma exchange could be administered during the induction period following local practice. However, rituximab was not administered within 48 hours before a plasma exchange treatment.

Assessments

Evaluations (including clinical, biochemical and patient-reported outcomes) were performed at 0, 1.5, 3 and 4 months.

Power calculation

Enrolment was set to be open until at least 160 patients were randomised at their month 4 visits. It was anticipated that 190 patients would be required in order to randomise 160 patients. Details of how the sample size was determined have been previously published.

Definitions

Remission was defined as a BVAS/WG of 1 or less with a prednisone/prednisolone dose of 10 mg/day or less by 4 months.

Statistical methods

Continuous variables are expressed as medians and IQRs. Categorical variables are presented as percentages and frequencies. A set of univariate logistic regression analyses to predict remission at month 4 for candidate factors was performed. Estimates of marginal ORs, with 95% CIs and p values are presented. The statistical comparisons were not formally powered or prespecified in the protocol so these results must be interpreted with caution. Data were analysed using R V.3.6.1.

RESULTS

Baseline demographics

188 patients were enrolled into the trial. Patient disposition throughout the 4-month induction period is shown in the consort diagram (figure 1) and baseline demographics in table 1. Ninety-five out of 188 (51%) patients were male, with a median age of 59 years (range 19–89) and prior disease duration of 5.0 years (range 0.4–34.5). One hundred and forty-nine out of 188 (79%) patients had previously received cyclophosphamide (median dose 9 g (range 0.15–301) and 67/188 (36%) had received rituximab (median dose 3910 mg (range 1000–16000)). At enrolment, 60/188 (32%) patients were on an oral immunosuppressive agent: (35/188 (19%) azathioprine; 12/188 (6%) mycophenolate mofetil; and 13/188 (7%) methotrexate), each of which were stopped as per protocol. One hundred and thirty-seven out of 188 (73%) had a history of previous cyclophosphamide (median dose 9 g (range 0.15–301) and 67/188 (36%) had received rituximab (median dose 3910 mg (range 1000–16000)).

<table>
<thead>
<tr>
<th>Table 1 Baseline demographics</th>
<th>Total (n=188)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years: median (range)</td>
<td>59 (19–89)</td>
</tr>
<tr>
<td>Male, number (%)</td>
<td>95 (51)</td>
</tr>
<tr>
<td>Race, number (%)</td>
<td>168 (89.4)</td>
</tr>
<tr>
<td>White</td>
<td>13 (6.9)</td>
</tr>
<tr>
<td>Asian</td>
<td>3 (1.6)</td>
</tr>
<tr>
<td>Black</td>
<td>1 (0.5)</td>
</tr>
<tr>
<td>Other</td>
<td>3 (1.6)</td>
</tr>
<tr>
<td>Disease duration, years: median (range)</td>
<td>5.0 (0.4–34.5)</td>
</tr>
<tr>
<td>Prior treatment with cyclophosphamide</td>
<td>149 (79.3)</td>
</tr>
<tr>
<td>Cumulative dose, grams (g): median (range)</td>
<td>9 (0.15–301)</td>
</tr>
<tr>
<td>Prior rituximab therapy</td>
<td>67 (35.6)</td>
</tr>
<tr>
<td>Cumulative dose, grams (g): median (range)</td>
<td>3910 (1000–16000)</td>
</tr>
<tr>
<td>Glucocorticoid induction regimen, number (%)</td>
<td></td>
</tr>
<tr>
<td>1 mg/kg/day starting dose (1A)</td>
<td>54 (28.7)</td>
</tr>
<tr>
<td>0.5 mg/kg/day starting dose (1B)</td>
<td>134 (71.3)</td>
</tr>
<tr>
<td>ANCA type, number (%)</td>
<td></td>
</tr>
<tr>
<td>ANCA negative</td>
<td>137 (72.9)</td>
</tr>
<tr>
<td>MPO-ANCA</td>
<td>51 (27.1)</td>
</tr>
<tr>
<td>Relapse type on entry into trial, number (%)</td>
<td></td>
</tr>
<tr>
<td>Severe</td>
<td>119 (63.3)</td>
</tr>
<tr>
<td>Non-severe</td>
<td>69 (36.7)</td>
</tr>
<tr>
<td>BVAS/WG: median (range)</td>
<td>5 (3–14)</td>
</tr>
</tbody>
</table>

ANCA, antineutrophil cytoplasmic antibody; BVAS/WG, Birmingham Vasculitis Activity Score for Wegener’s granulomatosis.
Vasculitis

Treatment exposure

The median total dose of rituximab in the induction phase was 2937 mg (range 1552–4320 mg) and cumulative oral glucocorticoid exposure in the 4-month induction phase was 3010 mg (2485–7875 mg) in the 1A higher dose induction regimen and 1960 mg (1715–3535 mg) in the 1B lower dose induction regimen. There was no difference in cumulative glucocorticoid exposure between patients that achieved and did not achieve remission (median dose 1960 mg in both groups (1A range: 1715–3010; 1B range: 1715–7875). Twenty-five per cent of patients deviated from the specified glucocorticoid tapering regimen at some point in the induction phase.

Disease response

One hundred and seventy-one out of 188 (90%) patients achieved remission at month 4 (figure 2). Of the 17 patients who did not achieve remission by month 4, 13 (76%) had PR3-ANCA positive disease and 10 (59%) had ear, nose and throat involvement at baseline. Fourteen out of 17 (82%) patients who did not achieve remission had severe (at least one major BVAS/WG item) disease and 5/17 patients (29%) received the higher glucocorticoid dosing regimen. Seven out of 17 (41.2%) non-responders had previously received rituximab, median cumulative dose of 4125 mg (1000–8930), which was comparable with responders (60/171 (35.1%), and cumulative dose 3910 mg (1500–16000)). At month 4, three patients had ongoing ENT disease activity; three had pulmonary manifestations; two had active renal disease; and four had other features of active disease (fatigue, pachymeningitis and headache). None of the following baseline variables were predictive of disease response: age, ANCA type at enrolment, glucocorticoid induction regimen, presence of ear, nose and throat or renal involvement (online supplementary table 1), although it is notable non-severe disease was associated with an OR of 2.93, 95% CI 0.915 to 13.1 for subsequent response. Of the 17 patients who did not progress in the trial, only 6/188 (3.2%) had a failure to achieve disease control at month 4, four died in the induction phase, two were withdrawn by their investigator (diagnosis of a new malignancy and occurrence of serious adverse events (SAEs)), three withdrew consent, one required additional therapy not permitted in the protocol and one failed screening and did not receive induction therapy.

Biochemical parameters

Median B cell count fell from 0.12×10^9/L (12%) (range 0–3.49 (0%–46%)) at baseline to 0×10^9/L (0%) (range 0–1 (0%–3%)) at month 4. There was no difference in median B cell counts between responders and non-responders. There were modest reductions in C reactive protein levels (median 2.65 mg/L (0–165) at baseline; 1.2 mg/L (0–183) at month 4) and erythrocyte sedimentation rate (21.5 mm/hour (1–149) to 12.5 mm/hour (2–100)) following treatment with glucocorticoids and rituximab. Serum creatinine remained stable (92.5 µmol/L (37.1–472) at baseline and 97.3 µmol/L (42-542) at month 4). One hundred and thirty out of 188 (69.1%) patients tested positive for ANCA at baseline and 81/188 patients (43.1%) at month 4. There was a greater proportion of PR3-ANCA positive patients who became ANCA negative (53.2% to 33.1%) compared with MPO-ANCA patients (14.9% to 12.4%) (figure 3). The two individuals who switched from being ANCA negative at baseline to PR3-ANCA positive at month 4 entered remission.

Safety

Forty-one SAEs occurred in 27 patients, including 13 severe infections (nine chest, three urinary and one gastrointestinal infection) in seven patients. Five out of 13 infections occurred within 4 weeks of the first induction dose of rituximab. In addition, there were 86 non-severe infections in 59 patients (online

Figure 2  Disease response according to baseline BVAS/WG score. Figures represent the number of individuals according to disease status. In addition to those displayed on the graph: at month 1.5, two individuals had severe disease, and four were withdrawn/missing. At month 3, one individual had severe disease and one limited disease. At month 4, one individual had severe disease, three had limited disease and three had persistent disease. Withdrawn/missing includes all participants who did not attend a study visit either due to death, withdrawal from trial or a missed visit. BVAS/WG, Birmingham Vasculitis Activity Score for Wegener’s granulomatosis.
supplementary table 2). Fifty-one patients had an IgG level less than 5 g/L at some point during the induction phase (table 2). Four patients (2.1%) died in the induction phase; causes of death included: pneumonia (2), cerebrovascular accident (1) and active vasculitis (1).

**DISCUSSION**

These data from the induction phase of the RITAZAREM trial, the largest reported prospective cohort of patients with relapsing AAV, demonstrate that rituximab, in conjunction with glucocorticoids, is effective at reinducing remission in patients with AAV who have relapsed, regardless of previous therapy. A high proportion of patients (171/188, 90%) achieved remission by 4 months, and it is notable that 71% of patients received the lower dose glucocorticoid regimen. Although there are retrospective series, the only previous prospective data on induction of remission for this subgroup of patients with ANCA-associated vasculitis was from the RAVE trial that observed a higher rate of remission in 50 relapsing patients treated with rituximab when compared with 50 relapsing patients treated with cyclophosphamide. Thus, these data confirm and extend the data on the efficacy of rituximab for relapsing GPA/MPA and supports a recommendation of rituximab for this indication.

The higher remission rate found in RITAZAREM versus RAVE may be due in part to the different definitions of remission. In RITAZAREM, remission was defined as a BVAS/WG ≤1 with a prednisolone dose ≤10 mg/day. The RAVE trial observed a lower remission rate of 64% at 6 months but required a BVAS/WG of zero and complete glucocorticoid withdrawal. The stricter definition of remission in RAVE, together with differences in trial design, and the enrolment in RAVE of a more severely affected patient population (median BVAS/WG 8.5 (5–13) for patients treated with rituximab), makes direct comparison between RITAZAREM and RAVE difficult. In the current study, only 6 of the 17 patients who did not achieve remission (3.2% of the whole study population) clearly represented failure of the therapy. The remainder were withdrawn from the study protocol either due to investigator or participant decision (seven patients, 3.7%) or died (four patients, 2.1%) within the induction phase. In this cohort, no baseline variables studied were predictive of failure of treatment response, although the small numbers of non-responders make it difficult for such an analysis to be definitive.

Induction regimens in AAV have been associated with high rates of SAEs, and these are more frequent and problematic than failures to control disease activity, thus improvements in the safety of induction regimens are required. In RITAZAREM, SAEs occurred in 14.3% of patients, which is a lower rate than seen in the RITUXVAS trial in which 42% of patients treated with rituximab experienced at least one SAE and the RAVE trial in which 22% of patients experienced at least one grade 3 adverse event. In the treatment of AAV, concomitant use of glucocorticoids is a major contributor to SAEs, especially infective risks, and two glucocorticoid regimens were permitted in this study to suit physician preference. The choice of glucocorticoid regimen was not randomised, and thus may have been subject to bias, so the...
relative efficacy of these two regimens cannot be completely analysed. Nonetheless, these two regimens appeared similarly effective with the lower dose approach providing approximately two-thirds of the total oral glucocorticoid exposure, and thus reduced dose glucocorticoids can be recommended as a treatment option for this indication.

The key strength of the study lies in the number of patients recruited, making this the largest cohort of patients with relapsing AAV to be studied in a clinical trial, facilitating the collection of high-quality efficacy and safety data on a complex patient population. This is a typical population of patients relapsing with AAV, enriched for patients with PR3-ANCA positivity, with median prior disease duration of 5 years, prior exposure to cyclophosphamide and/or rituximab in the majority and a degree of established chronic damage, meaning that results are broadly applicable. A potential weakness of this study was the option for investigators to choose, rather than randomly assigning the glucocorticoid dosing regimen in a blinded manner. Prescribing practices for use of glucocorticoids in AAV vary, necessitating a pragmatic approach to trial design. However, investigators were required to select the dosing regimen at enrolment, and tapering schedules were standardised.

Achieving a negative serum ANCA test following induction therapy is associated with a lower subsequent risk of relapse in AAV.12 17 In the current study, despite 90% of patients achieving remission at month 4, 46% remained positive for serum ANCA at month 4, supporting data from the RAVE trial, in which 53% of patients treated with rituximab remained positive for ANCA at 6 months.12 Follow-up in the randomised phase of the RITAZAREM trial will provide further insight into the significance of changes in ANCA levels.

These data from the first phase of RITAZAREM demonstrate that rituximab, in conjunction with even relatively low doses of glucocorticoids, is highly effective at reinducing remission in patients with AAV who have relapsed, with a safety profile similar to or better than previous studies.

Author affiliations
1University of Cambridge, Cambridge, UK
2Cambridge University Hospitals NHS Foundation Trust, Cambridge, UK
3Mayo Clinic, Rochester, Minnesota, USA
4Cambridge Clinical Trials Unit, Cambridge University Hospitals NHS Foundation Trust, Cambridge, UK
5University Hospitals of Leicester NHS Trust, Leicester, UK
6NIHR Leeds Musculoskeletal Biomedical Research Unit, Leeds Teaching Hospitals Trust, Leeds, UK
7Department of Renal Medicine, Karolinska University Hospital and Karolinska Institute, Stockholm, Sweden
8Ipswich Hospital NHS Trust, Ipswich, UK
9University of Toronto, Toronto, Ontario, Canada
10University of Leicester, Leicester, UK
11University of North Carolina, Chapel Hill, North Carolina, USA
12East Kent Hospitals University NHS Foundation Trust, Canterbury, UK
13Cedars-Sinai Medical Center, Los Angeles, California, USA
14University of Miyazaki, Miyazaki, Japan
15Chiba University, Chiba, Japan
16University of Michigan, Ann Arbor, Michigan, USA
17University of Birmingham, Birmingham, UK
18Kyoto University, Kyoto, Japan
19McMaster University, Hamilton, Ontario, Canada
20Dudley Group NHS Foundation Trust, Dudley, UK
21University of Utah Vasculitis Center, Salt Lake City, Utah, USA
22Ko sin University School of Medicine, Tokyo, Japan
23Cleveland Clinic Foundation, Cleveland, Ohio, USA
24Rheumatology, Nottingham University Hospital, Nottingham, UK
25Nuffield Department of Orthopaedics, Rheumatology and Musculoskeletal Science (NORMS), University of Oxford, Oxford, UK
26Okayama University Hospital, Okayama, Japan

Acknowledgements The RITAZAREM trial is directed by the European Vasculitis Society and the Vasculitis Clinical Research Consortium (VCRC). The primary sponsor is Cambridge University Hospitals NHS Foundation Trust, and there are collaboration and data-sharing agreements with the University of Pennsylvania and the University of Miyazaki and Okayama University in Japan. Additional RITAZAREM coinvestigators were: Dr Y Arimura (Kyorin University, Japan); Dr M Clarkson (Cork University Hospital, Ireland); Dr J de Zoya (North Shore Hospital, Auckland, New Zealand); Dr T Endo (Kitano Hospital, Japan); Dr Y Hamano (Tokyo Metropolitan Geriatric Hospital, Japan); Dr H Kono (Teikyo University Hospital, Tokyo, Japan); Dr S Lawman (Brighton Royal Sussex County Hospital, UK); Dr E Muso (Kitano Hospital, Japan); Dr K Sada (Okayama University, Japan); Dr R Smith (Ipswich Hospital, UK); Dr K Suzuki (Teikyo University, Japan); Dr T Tsukamoto (Kitano Hospital, Japan); Dr S Uchida (Teikyo University Hospital, Tokyo, Japan); Dr A Vaglio (University of Parma, Italy); and Dr R Watts (Ipswich Hospital, UK).

Contributors RMS, DJ and PM conceived and designed the study. US, RBJ and SB were also involved in study design. RMS, SB, NM, DJ and PM analysed the data and interpreted the results. RMS wrote the manuscript with support from DJ and PM. All authors collected data and contributed critical appraisal to the final manuscript.

Funding RITAZAREM is funded by grants from Versus Arthritis (formerly Arthritis Research UK) (Grant number 18706) and Roche/Genentech (ZMA28150). The Vasculitis Clinical Research Consortium (VCRC) (USA AR057319 and U01 AR5187404) is part of the United States National Institutes of Health Rare Diseases Clinical Research Network, an initiative of the Office of Rare Diseases Research, National Center for Advancing Translational Science (NCATS). The VCRC is funded through collaboration between NCATS, and the National Institute of Arthritis and Musculoskeletal and Skin Diseases, and has received funding from the National Center for Research Resources (USA RR02197) and the National Heart, Lung, and Blood Institute. This research was also supported by the National Institute for Health Research, Cambridge Biomedical Research Centre and the Cambridge Clinical Trials Unit.

Competing interests RMS reports grants from Roche during the conduct of the study. RBJ reports grants from GlaxoSmithKline, personal fees from ChemoCentryx outside the submitted work. US reports grants from Genentech, during the conduct of the study, grants from Chemocentryx, personal fees from BMS, personal fees from AstraZeneca, personal fees from Genentech, personal fees from Merck/MSD, personal fees from Abbvie outside the submitted work. CKC reports grants from GlaxoSmithKline, consulting and travel grants from Roche outside the submitted work. CL reports grants from Genentech outside the conduct of the study. NK reports personal fees and non-financial support from Roche, non-financial support from Bristol Meyers Squibb outside the submitted work. RK reports grants and personal fees from Roche, grants from Arthritis Research UK, grants from Genentech, grants from ChemoCentryx, personal fees from Roche, personal fees from GlaxoSmithKline, grants from Roche, outside the submitted work. RA reports grants from Roche, outside the submitted work. FB reports grants from Roche, outside the submitted work. LG reports grants and personal fees from Roche, outside the submitted work. RC reports grants from Roche, outside the submitted work. MK reports grants and personal fees from Roche, outside the submitted work. EY reports grants from Roche, outside the submitted work.

The VCRC is funded through collaboration between NCATS, and the National Institute of Arthritis and Musculoskeletal and Skin Diseases, and has received funding from the National Center for Research Resources (USA RR02197) and the National Heart, Lung, and Blood Institute. This research was also supported by the National Institute for Health Research, Cambridge Biomedical Research Centre and the Cambridge Clinical Trials Unit.
GSK, personal fees from Inflarx, personal fees from Medpace, personal fees from MedImmune, personal fees from Roche, outside the submitted work. HM reports personal fees from Abbvie, personal fees from Boehringer-Ingelheim, personal fees from Teijin, outside the submitted work. PM reports personal fees from Kiniksa, personal fees from ChemoCentryx, personal fees from Celgene, personal fees from Insmed, outside the submitted work. PN reports other from ChemoCentryx, other from Inflarx, other from Omeros, other from Aurinia outside the submitted work. CP reports grants and personal fees from Roche, personal fees from ChemoCentryx, grants and personal fees from GlaxoSmithKline, personal fees from Sanofi, personal fees from Inflarx outside the submitted work. FP reports grants from Vifor Pharma outside the submitted work. RS reports grants from GlaxoSmithKline, grants from chemocentryx, grants from Roche/Genentech, grants from BPI, personal fees from GlaxoSmithKline, personal fees from chemocentryx. RS reports personal fees from Bristol-Myers Squibb, other from Alexion outside the submitted work. VT reports other from Abbvie, other from Amgen, other from Boehringer-Ingelheim, other from Calilddd, other from ChemoCentryx, other from FMC, other from Retrophin outside the submitted work. PM reports personal fees from Abbvie, grants and personal fees from AstraZeneca, personal fees from Biogen, grants and personal fees from Bristol-Myers Squibb, grants and personal fees from Boehringer-Ingelheim, grants and personal fees from Celgene, grants and personal fees from chemocentryx, CSL Behring, grants and personal fees from Genentech/Roche, grants and personal fees from Genzyme/Sanofi, grants and personal fees from GlaxoSmithKline, grants and personal fees from Inflarx, personal fees from Insmed, personal fees from Jannsen, personal fees from Kiniksa, grants from Kypha, personal fees from Sparrow, grants from TerumoBCT outside the submitted work. DJ reports grants from Roche/Genentech, during the conduct of the study; grants from Sanofi-Genzyme, grants and personal fees from Chemocentryx, grants and personal fees from GSK, grants from Roche/Genentech, personal fees from Takeda, personal fees from Insmed, personal fees from Astra-Zeneca, personal fees from Inflarx-Rx, personal fees from Chugai, personal fees from Boehringer-Ingelheim outside the submitted work.

Patient and public involvement Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Methods section for further details.

Patient consent for publication Not required.

Ethics approval An initial favourable ethical opinion was granted by NRES Committee East of England – Cambridge South: REC reference: 12/EE/0230 on 24 July 2012. US approvals: Cedars-Sinai Medical Center Institutional Review Board: Pro00031367; Cleveland Clinic Institutional Review Board: 13-666; Hospital for Special Surgery Institutional Review Board: 13-114; Mount Sinai Hospital Research Ethics Board: 12-0231-A; St. Joseph’s Hospital Hamilton Integrated Research Ethics Board: 13-017; University of Pittsburgh Institutional Review Board: PRO130020329; University of Pennsylvania Office of Regulatory Affairs: 816166; The Mayo Clinic, University of Michigan, University of North Carolina and the University of Utah all deferred to the University of Pennsylvania Ethics board and fall under the University of Pennsylvania approval number. Japanese ethics committee numbers: University of Miyazaki 2013-126; Chiba University 97; Kitano Hospital P14-01-002; Okayama University m05002; Koyin University H26-031; Teikyo University 14-031; TMIHG 260201.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request. Deidentified participant data can be requested from the corresponding author.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution 4.0 Unported (CC BY 4.0) license, which permits others to copy, redistribute, remix, transform and build upon this work for any purpose, provided the original work is properly cited, a link to the licence is given, and indication of whether changes were made. See: https://creativecommons.org/licenses/by/4.0/.