EXTENDED REPORT

Anaemia may add information to standardised disease activity assessment to predict radiographic damage in rheumatoid arthritis: a prospective cohort study

Burkhard Möller,1 Almut Scherer,2 Frauke Förger,1 Peter M Villiger,1 Axel Finckh,3 on behalf of the Swiss Clinical Quality Management Program for Rheumatic Diseases

ABSTRACT

Objective Anaemia in rheumatoid arthritis (RA) is prototypical of the chronic disease type and is often neglected in clinical practice. We studied anaemia in relation to disease activity, medications and radiographic progression.

Methods Data were collected between 1996 and 2007 over a mean follow-up of 2.2 years. Anaemia was defined according to WHO (Hb haemoglobin<12 g/dl, Hct haematocrit<36%, or alternative criteria. Anaemia prevalence was studied in relation to disease parameters and pharmacological therapy. Radiographic progression was analysed in 9731 radiograph sets from 2681 patients in crude longitudinal regression models and after adjusting for potential confounding factors, including the clinical disease activity score with the 28-joint count for tender and swollen joints and erythrocyte sedimentation rate (DAS28ESR) or the clinical disease activity index (cDAI), synthetic antirheumatic drugs and antitumour necrosis factor (TNF) therapy.

Results Anaemia prevalence decreased from more than 24% in years before 2001 to 15% in 2007. Erosions progressed significantly faster in patients with anaemia (p<0.001). Adjusted models showed these effects independently of clinical disease activity and other indicators of disease severity. Radiographic damage progression rates were increasing with severity of anaemia, suggesting a ‘dose-response effect’. The effect of anaemia on damage progression was maintained in subgroups of patients treated with TNF blockade or corticosteroids, and without non-selective nonsteroidal anti-inflammatory drugs (NSAIDs).

Conclusions Anaemia in RA appears to capture disease processes that remain unmeasured by established disease activity measures in patients with or without TNF blockade, and may help to identify patients with severe progressive disease.

INTRODUCTION

Anaemia in rheumatoid arthritis (RA) is prototypical of anaemia of chronic disease (ACD), characterised by low serum iron concentrations in conjunction with normal or increased storage iron.1 ACD without functional iron deficiency is normochromic, normocytic, and usually mild.2 Haemoglobin concentrations were a component of earlier RA disease activity measures until they were omitted during validation of the original RA disease activity score (DAS) due to low correlation with other disease activity measures, despite moderate correlation with radiographic outcome.3

The significantly higher need for joint replacement and higher mortality rates in the anemic patients of the British early RA cohort (ERAS) challenged the concept of causal ACD treatment by aiming at remission only with synthetic disease-modifying antirheumatic drugs (DMARDs).4 Inhibition of human erythroid colony-forming units in the bone marrow downstream of tumour necrosis factor-α (TNF-α)5 and anaemia6 is reversible upon TNF-α blockade, even in otherwise refractory RA patients. Here, we question whether anaemia in RA is still relevant in terms of prevalence and clinical consequences in the era of TNF-α blockade, one of the most efficacious and common biological RA treatments.

METHODS

Patients and setting

This longitudinal population-based cohort study is nested within the Swiss clinical quality management (SCQM) database, which is described elsewhere.7 Patient registration in SCQM is restricted to board-certified rheumatologists to follow the effectiveness and safety of antirheumatic therapies. More than two-thirds of all practicing rheumatologists in Switzerland contributed patients, representing ~10% of all hypothetical Swiss RA patients.8 Approximately 60% of patients come from private practices, 20% from university hospitals, and 20% from non-academic centres. Ethical approval for patient enrolment into the SCQM programme and related studies was obtained from the Swiss Academy of Medical Sciences review board.

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Written informed consent was provided by all 4377 patients.8,9 Approximately 60% of patients come from private practices, 20% from university hospitals, and 20% from non-academic centres. Ethical approval for patient enrolment into the SCQM programme and related studies was obtained from the Swiss Academy of Medical Sciences review board. Written informed consent was provided by all 4377 participants with classification criteria-guided RA diagnosis, and at least one haemoglobin assessment between the start of the registry in 1996 and a major system revision in 2007.

Anaemia definitions

We used the WHO definition of anaemia with lower normal haemoglobin limits of 12 g/dl in women and 13 g/dl in men.11 To explore the effect of anaemia severity, we used the less stringent anaemia definition proposed by Beutler et al,12 using thresholds of 12.2 g/dl in white women and
of 13.7 g/dl or 13.2 g/dl in white men aged 20–59 years, or 60 + years, respectively, in sensitivity analyses. ‘More severe anaemia’ was arbitrarily defined (in the absence of official definitions) by haemoglobin levels lower than 11 g/dl in women and 12 g/dl in men. Haemoglobin concentrations, white blood cell counts, and platelet counts were derived from certified local laboratories.

Covariates
Registry documentation provided the American College of Rheumatology core set of disease activity measures, which allowed to calculate the DAS including the 28-joint count for tender and swollen joints and erythrocyte sedimentation rate (DAS28ESR) and the clinical disease activity index (cDAI). Other covariates were age and gender, rheumatoid factor (RF) status, and RA disease duration. RA therapies, including TNF-α blockers, methotrexate with or without folic acid supplementation, and other synthetic DMARDs, corticosteroids, cyclo-oxygenase-II selective (Coxib type) NSAIDs, non-selective NSAIDs and proton-pump inhibitors (PPI). Generic statisticians on the presence of haematological, renal, or gastrointestinal comorbidities was also available, without detailed information.

Outcome
Radiographic evaluation of erosions was centrally performed by an independent assessor following the Ratingen score, a validated method measuring quintiles of the total joint bone surface in 38 joints in the hands and forefeet. Radiographic data were dated method measuring quintiles of the total joint bone surface and expressed in percent of the maximal 190-point score and changes over time reported in absolute differences. The minimal detectable change for this method is ~3.3% of the maximum score.

Statistical analysis
Descriptive statistics included the mean and SD for normally distributed variables, and the median and IQR for non-parametric continuous variables. Sporadic missing covariates were replaced by the population mean (<5%). Between-group differences in disease characteristics from baseline to registration were compared using a Student t test for normally distributed variables, the Wilcoxon rank sum test for non-normally distributed continuous variables, and Pearson’s χ² test for dichotomous variables. We examined potential time trends in the prevalence and incidence of anaemia and the haemoglobin normalisation rates with Cox regression analyses. Patients were defined as ‘anaemic’ according to the chosen definition between two radiographs by subnormal haemoglobin concentrations in more than 50% of determinations. We used mixed-model random-slope regression for longitudinal data to analyse radiographic damage progression over time. Unless otherwise stated, data were adjusted for all baseline covariates (table 1) and the baseline and time-varying DAS28ESR or cDAI. Patients may have switched between the anaemic and non-anaemic groups according to their time-dependent haemoglobin concentration. We chose the best-fitting parsimonious covariance structure using a series of likelihood ratio tests based on restricted maximum likelihood estimates. We examined whether time best fit the data as a linear trend or as a quadratic function. Finally, we selected the best-fitting mean response model of the outcome, controlling for other covariates for the respective outcome. All statistical tests were two-sided performed with Stata V 12 for Windows (Stata Statistical Software, StataCorp, Texas, USA), and were evaluated at the 0.05 significance level.

Subgroup analyses
Subgroups were analysed in order to further address potential treatment effects. Here, we performed statistical models for the radiographic outcome after having selected either patients with...
RESULTS
Baseline characteristics of all included patients are listed in table 1. More than 80% of subsequent consultations were scheduled after approval of TNF-α blocking agents in Switzerland for patients after failure of at least one conventional DMARD. Anti-TNF-α and other DMARD therapies were often intensified after inclusion, and the prevalence of TNF-α inhibitors and second-line biologics progressively reached 35% and 2% of patients, respectively. Use of methotrexate or other conventional DMARDs was registered at least once in 64% and 60% of the patients, respectively, over time.

Haemoglobin concentrations
Patients had a mean of 4.4 haemoglobin assessments over a mean follow-up period of 2.8 years. The mean baseline haemoglobin concentration was 12.7 (SD±1.4 g/dl) in women and 14.0±1.6 g/dl in men. Haemoglobin levels increased by 0.08±0.04 g/dl (p=0.04) after the first year, by 0.14±0.07 g/dl (p=0.06) after the second year, and by 0.16±0.09 g/dl (p=0.077) after the third year. Individual haemoglobin fluctuations in either direction (mean 1.6±1.2 g/dl) were more substantial. Higher haemoglobin concentrations were significantly associated with higher haemoglobin concentrations at baseline (0.04, 95% CI 0.03 to 0.05, p<0.001) and male gender (1.11, 95% CI 1.03 to 1.19, p<0.001), but negatively associated with higher DAS28ESR (−0.20 g/dl, 95% CI −0.23 to −0.18, p<0.001), higher erosion scores (−0.005, 95% CI −0.007 to −0.003, p<0.001), and treatment with non-selective NSAIDs (−0.16, 95% CI −0.20 to −0.12, p<0.001), with or without PPI cotherapy. Haemoglobin levels were not associated with baseline renal or gastrointestinal comorbidity.

Prevalence of anaemia
Anaemia11 occurred with similar frequency in men and women, oscillating around 24%, before a continuous negative time trend started in 2001 to reach 14.7% in 2007. No anaemia at any visit was found in 69.2% of patients, and persistent anaemia at all visits occurred in 10% of patients.

Incidence and correction of anaemia
The annual anaemia incidence rate was 7.6% in 2531 initially non-anaemic patients. Patients with higher baseline DAS28ESR (HR 1.14, 95% CI 1.07 to 1.22, p<0.001), RF positivity (HR 1.49, 95% CI 1.16 to 1.92, p=0.002), registered haematological comorbidity (HR 1.76, 95% CI 1.23 to 2.53, p=0.002) or corticosteroid therapy (HR 1.31, 95% CI 1.03 to 1.67, p=0.028) became significantly more often anaemic. PPI comedication was often used in incident anaemia (HR 2.63, 95% CI 0.97 to 7.14, p=0.057) among patients with non-selective NSAID therapy (HR 1.29, 95% CI 1.06 to 1.57, p=0.013). By contrast, coxibs, gastrointestinal comorbidities or renal comorbidities were not associated with different anaemia incidence. Gastrointestinal toxicities were attributed to NSAIDs in 12 cases, but only one clinically manifest bleeding was reported. Haemoglobin concentrations normalised after a median of 1.3 (IQR: 0.9–3.2) years in 800 initially anaemic patients.11 Anaemia correction was slower in patients with reported haematological disorder (HR 0.60, 95% CI 0.48 to 0.75, p<0.001) or under non-selective NSAIDs (HR 0.73, 95% CI 0.60 to 0.89, p=0.002), but was accelerated after anti-TNF-α (HR 1.22, 95% CI 1.02 to 1.45, p=0.028) or PPI therapy (HR 7.77, 95% CI 1.86 to 32.39, p=0.005).

Reporting of anaemia and haematological comorbidities
Treating rheumatologists reported baseline haematological comorbidity in only 19% of anaemic patients. Indicators of bone marrow insufficiency or other relevant blood cell abnormalities (white blood cell count below 3 g/l and thrombocytopenia below 50 g/l) were reported in less than 0.4% or less than 0.2% of all cases, respectively. Reactive elevations in these blood cell counts, platelets ≥400 g/l (3.6%), or leukocytes ≥10 g/l (13.8%) were more frequent. Patient’s treatment with iron, vitamin B12, or erythropoietin was reported in nine, 15 and 14 cases, respectively. Malignancies were reported in 1.3% of all patients.

Anaemia and erosions
This relationship was studied in 9731 sets of radiographs from hands and forefoot in 2681 patients, of whom 627 individuals were anaemic at baseline according to WHO definition. The less stringent anaemia definition12 was fulfilled in 1487 individuals, and 506 patients complied with the definition of ‘more severe anaemia’. In crude analyses and after statistical adjustments for potential confounding baseline parameters and the time-dependent DAS28ESR or cDAI, anaemic patients experienced significantly faster radiographic progression (p<0.001) than their non-anaemic counterparts, irrespective of the employed anaemia definition (figure 1). A ‘dose-response effect’ was suggested by the average 2.0% annual damage progression of the maximal score in WHO-defined anaemic patients versus 1.2% in non-anaemic patients; these values were 1.4% vs 1.1% according to the higher threshold of normal haemoglobin concentrations,12 but 2.4% vs 1.2% according to the definition of ‘more severe anaemia’. Significant baseline determinants of radiographic progression in all these statistical models were baseline erosion scores (mean ∼1.0 in all models, p<0.001), gender to the advantage of the male (mean 0.3–0.4%, dependent on the model, p=0.024–0.038), methotrexate therapy (mean 0.2–0.3% p=0.012–0.043), and non-selective NSAID therapy (mean ∼0.3% in all models, p=0.013–0.024).

Subgroup analyses
We obtained similar results as in the total patient population after exclusion of patients with non-selective NSAID therapy (figure 2A,B). Anaemia effects on erosive progression were reproduced in another independently selected patient population with anti-TNF therapy (figure 2C,D). By contrast with mild anaemia,12 WHO defined and ‘more severe anaemia’ exhibited similarly significant effects on x-ray progression (p<0.001) in these subgroups. Anaemia effects on radiographic outcome went insignificant after exclusion of all patients with steroid registration, but remained significant in the complementary population (p<0.001, data not shown). Finally, mean joint damage progression was faster in anaemia after selection of patients with DAS28ESR defined remission at their last observation (p=0.026, data not shown).

DISCUSSION
In this study, WHO-defined anaemia was present in approximately every fifth patient at baseline, and at least in every third patient over their observation time, demonstrating ongoing...
Figure 1  Erosive progression over time in all 2681 evaluable rheumatoid arthritis (RA) patients. Progression trajectories were adjusted for differences in patient characteristics at their first x-ray (table 1), (A,C) DAS including the 28-joint count for tender and swollen joints and erythrocyte sedimentation rate (DAS28ESR) or (B,D) clinical disease activity index where indicated, plus anti-inflammatory therapies and covariates of RA severity. Values are mean and 95% CI. Erosion scores progressed significantly faster in patients with anaemia than in non-anaemic RA patients. The kinetics of damage progression accelerated with the grade of anaemia severity. (A and C) For comparison, the lower margin of the shaded area represents mean damage progression in clinical remission (DAS28ESR<2.6), and the upper margin disease progression in patients with high clinical disease activity (DAS28ESR>5.1). Patient numbers at given time points are the same in left (A,C) and right hand figures (B,D).

Figure 2  Erosive progression over time was analysed in patients with and without anaemia,11 (A,B) without NSAID and (C,D) with antitumour necrosis factor-α therapy. Values are mean and 95% CI. The progression of erosions (mean±95% CI) was adjusted for potential confounders (figure 1, table 1). Patient numbers at given time points are the same in left (A,C) and right hand figures (B,D).
relevance of anaemia despite decreasing time trends. We confirmed that anaemia in RA is usually mild, but still only partially resolved by available therapeutics and of remarkable importance for radiographic outcome.

SCQM services a typical RA registry cohort in terms of RF status, anaemia frequency and severity,\textsuperscript{18} other extra-articular RA manifestations,\textsuperscript{20} comorbidities,\textsuperscript{21} DMARD and TNF-\(\alpha\) blocker prescription rates.\textsuperscript{22} SCQM is unique among the national registries in providing centralised radiographic assessment for RA patients. The necessary compromise was the abandonment of the evaluation of joint space narrowing, but the Ratingen erosion score provides good reliability and less susceptibility to ceiling effects in advanced disease, since it uses a true ordinal rating system.\textsuperscript{16}

This large real-life observational study confirmed that mean joint damage progression rates are significantly lower in clinical remission than in active disease, but this goal was achieved at the end of observation in only 30\% of patients. Articular inflammation can be detected by ultrasound and MRI in the absence of clinically detectable inflammation,\textsuperscript{24} which could explain the dissociation of disease activity from radiographic outcome.\textsuperscript{22,27} In the present study, anaemia-related progression of joint damage was not restricted to the patients with clinically active disease.\textsuperscript{14} Thus, anaemia may be a useful surrogate for subclinical RA disease activity.

The well established relationship between inflammation and anaemia was confirmed in this study by significant associations between lower haemoglobin concentrations and higher DAS28ESR and by faster haemoglobin normalisation after TNF-\(\alpha\) blockade. ACD in RA may be caused by a shortened red blood cell lifespan, pathologic iron homeostasis induced by hepcidin, and blunted response to erythropoietin.\textsuperscript{1} Interestingly, erythropoietin treatment reduced disease activity in RA patients\textsuperscript{28} and tissue damage in collagen-induced arthritis models.\textsuperscript{30} Our results are consistent with anaemia as an indicator of deficient anti-inflammatory properties of erythropoietin.\textsuperscript{31} Hepcidin may trigger functional iron deficiency upon induction by TNF-\(\alpha\), interleukin-6, and other inflammatory stimuli\textsuperscript{1}, resulting in reduced intestinal iron uptake at the mucosal barrier, and iron retention in the reticulo-endothelial system via internalisation of the same exclusive cellular iron exporter ferroportin on both cell types. While elevated hepcidin levels could likewise be controlled by TNF-\(\alpha\) blockade and manifest iron deficiency\textsuperscript{32}, its plasma concentrations appear not to be predictive for RA joint damage.\textsuperscript{33}

The association between anaemia and erosions was not observed in a smaller patient population,\textsuperscript{34} but in the large ERAS inception cohort, anaemic patients underwent a more aggressive course of joint destruction.\textsuperscript{2} The SCQM-RA cohort includes patients with early and established disease, with lower mean disease activity at inclusion than the ERAS cohort, and a substantial proportion of patients received anti-TNF-\(\alpha\) therapy, yet the consequences of anaemia on RA outcome appear to be similar in these two populations.\textsuperscript{3}

The present study is limited by missing biological markers of iron deficient erythropoiesis, such as erythrocyte indices or parameters of iron metabolism. Iron-deficiency anaemia is increasing in industrialised countries with age and with risk factors for bleeding,\textsuperscript{35} as seen here with an increased prevalence and delayed recovery of anaemia in patients taking non-selective NSAIDs, in contrast with Coxib-type NSAIDs.\textsuperscript{36} One can estimate, on the basis of prevalence data in young Swiss males,\textsuperscript{13} that iron deficiency contributes to anaemia prevalence in about 30\% of the cases before, and in 50\% after, market introduction of TNF blockers. With this assumption, the prevalence of deficient erythropoiesis may still be underestimated, as iron deficiency was rapidly uncovered by erythropoietin stimulation in the vast majority of patients with longstanding and treatment-resistant active RA.\textsuperscript{28} Here, it appears necessary to point out that a higher proportion of similar cases with established disease, no matter of being clinically active on a very short term, but with subsequent functional iron deficiency on a longer time perspective would not be a confounding factor, but support the prognostic value of anaemia assessment.

Observational studies have inherent limitations, such as incomplete data, bias by indication and confounding factors. We do not believe there is a specific selection bias by anaemia in this register, and the most important confounding factors have been measured and could be adjusted for in the analysis. Corticosteroids may carry an increased risk of anaemia due to toxic mucosal drug effects\textsuperscript{37} or a bias by indication. As increased rates in radiographic damage in association with anaemia were not seen in patients without corticosteroids, this constellation differs from that in NSAID-treated patients. Divergent reporting of haematological conditions and true anaemic haemoglobin levels were explainable by the mild degree in an explained context. Missing adjustments for C-reactive protein concentrations appear less critical with regard to the superior validity of composite scores over single items.\textsuperscript{1,4} We were unable to adjust for anticyclic citrullinated peptide antibody status as this information was missing in 40\% of patients,\textsuperscript{38,39} nor do not know the shared epitope status in this large cohort, which may control immune responses\textsuperscript{30,41} and determine haematopoietic progenitor cell senescence.\textsuperscript{42,43}

In summary, these results indicate that anaemia may serve as predictor of worse outcome in RA patients. Our data suggest that anaemia and erosion progression are associated independently of common disease activity outcome measures. This report may add clinical background to recent discoveries at the nexus of inflammation, haematopoiesis, and iron metabolism,\textsuperscript{30,31} and highlights the clinical implications of anaemia in RA. Diagnosis of anaemia in RA should prompt a thorough search for subclinical disease activity, after exclusion of other frequent causes.

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Contributors BM planned this study, performed statistical analyses and is responsible for the overall content as guarantor. AS tested the results in alternative statistical models. FF provided important intellectual content to this study. PV provided administrative support. AF gave advice and supervised all statistical procedures. All authors approved the final manuscript. Physicians and institutions who contributed patients to the SCQM registry without fulfilling the requirements for authorship are listed under (http://www.scqm.ch/institutions).

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REFERENCES

1 Weiss G, Goodnough LT. Anemia of chronic disease. N Engl J Med 2005;352:1011–23.
2 Wilson A, Yu HT, Goodnough LT, et al. Prevalence and outcomes of anemia in rheumatoid arthritis: a systematic review of the literature. Am J Med 2004;116(Suppl 1):S50–57.
3 van der Heijde DM, van’t Hof MA, van Riel PL, et al. Validity of single variables and composite indices for measuring disease activity in rheumatoid arthritis. Ann Rheum Dis 1992;51:177–81.
4 Young A, Koduri G. Extra-articular manifestations and complications of rheumatoid arthritis. Best Pract Res Clin Rheumatol 2007;21:907–27.
5 Means RT Jr, Desespois EN, Krantz SB. Inhibition of human colony-forming-unit erythroid by tumor necrosis factor requires accessory cells. J Clin Invest 1990;86:538–41.
6 Doyle MK, Rahman MU, Han C, et al. Treatment with infliximab plus methotrexate improves anemia in patients with rheumatoid arthritis independent of improvement in other clinical outcome measures—a pooled analysis from three large, multicenter, double-blind, randomized clinical trials. Semin Arthritis Rheum 2009;39:123–31.
7 Ulte E, Fransen J, Langenegger T, et al. Clinical quality management in rheumatoid arthritis: putting theory into practice. Swiss Clinical Quality Management in Rheumatoid Arthritis. Rheumatology (Oxford) 2000;39:542–9.
8 Myasoedova E, Crowson CS, Kremers HM, et al. Is the incidence of rheumatoid arthritis rising? results from Olmsted County, Minnesota, 1955–2007. Arthritis Rheum 2010;62:1576–82.
9 Neovius M, Simard JF, Aukland K. Nationwide prevalence of rheumatoid arthritis and penetration of disease-modifying drugs in Sweden. Ann Rheum Dis 2011;70:624–9.
10 Arnett FC, Edworthy SM, Bloch DA, et al. The American Rheumatism Association 1987 revised criteria for the classification of rheumatoid arthritis. Arthritis Rheum 1988;31:15–24.
11 DeMaeyer E, Adels-Tegman M. The prevalence of anemia in the world. World Health Stat Q 1985;38:302–16.
12 Beutler E, Waalen J. The definition of anemia: what is the lower limit of normal of the blood hemoglobin concentration? Blood 2006;107:1747–50.
13 Felson DT, Anderson JJ, Boers M, et al. The American College of Rheumatology preliminary core set of disease activity measures for rheumatoid arthritis clinical trials. The Committee on Outcome Measures in Rheumatoid Arthritis Clinical Trials. Arthritis Rheum 1993;36:729–40.
14 Prevoo ML, van’t Hof MA, Kuper HH, et al. Modified disease activity scores that include twenty-eight-joint counts. Development and validation in a prospective longitudinal study of patients with rheumatoid arthritis. Arthritis Rheum 1995;38:44–8.
15 Aleataha D, Smolen J. The Simplified Disease Activity Index (SDAI) and the Clinical Disease Activity Index (CDAI): a review of their usefulness and validity in rheumatoid arthritis. Clin Exp Rheumatol 2005;23:S100–6.
16 Rau R, Wassenberg S, Herborn G, et al. A new method of scoring radiographic change in rheumatoid arthritis. J Rheumatol 1998;25:2094–107.
17 Fiandt A, Moller B, Dudler J, et al. Evolution of radiographic joint damage in rituximab-treated versus TNF-treated rheumatoid arthritis cases with inadequate response to TNF antagonists. Ann Rheum Dis 2012;71:1680–5.
18 Wolfe F, Michaud K. Anemia and renal function in patients with rheumatoid arthritis. J Rheumatol 2006;33:1516–22.
19 Han C, Zhao N, Rahman MU, et al. A case-control study of anaemia in patients with rheumatoid arthritis treated with disease-modifying antirheumatic drugs in an adult population in the US: prevalence and impact on healthcare utilisation. J Med Econ 2008;11:255–64.
20 Turesson C, O’Fallon WM, Crowson CS, et al. Occurrence of extraarticular disease manifestations is associated with excess mortality in a community based cohort of patients with rheumatoid arthritis. J Rheumatol 2002;29:62–7.
21 Hyrich K, Symmons D, Watson K, et al. Baseline comorbidity levels in biologic and standard DMARD treated patients with rheumatoid arthritis: results from a national patient register. Ann Rheum Dis 2006;65:895–8.
22 Ziegler S, Huscher D, Karberg K, et al. Trends in treatment and outcomes of rheumatoid arthritis in Germany 1997–2007: results from the National Database of the German Collaborative Arthritis Centres. Ann Rheum Dis 2011;69:1803–8.
23 Furnt DE, Pagan AI, Harrold LR, et al. Greater likelihood of remission in rheumatoid arthritis patients treated earlier in the disease course: results from the Consortium of Rheumatology Researchers of North America registry. Arthritis Care Res (Hoboken) 2011;63:856–64.
24 Brown AK, Conaghan PG, Karim Z, et al. An explanation for the apparent dissociation between clinical remission and continued structural deterioration in rheumatoid arthritis. Arthritis Rheum 2008;58:2956–67.
25 Mulherin D, Fitzgerald O, Brenihan B. Clinical improvement and radiological deterioration in rheumatoid arthritis: evidence that the pathogenesis of synovial inflammation and articular erosion may differ. Br J Rheumatol 1996;35:1263–8.
26 Møltenaer ET, Voskuyl AE, Djinot H, et al. Progression of radiologic damage in patients with rheumatoid arthritis in clinical remission. Arthritis Rheum 2004;50:36–42.
27 Smolen JS, Han C, van der Heijde DM, et al. Radiographic changes in rheumatoid arthritis patients attaining different disease activity states with methotrexate monotherapy and infliximab plus methotrexate: the impacts of remission and tumour necrosis factor blockade. Ann Rheum Dis 2009;68:823–7.
28 Kaltwasser JP, Kessler U, Gottschalk R, et al. Effect of recombinant human erythropoietin and intravenous iron on anemia and disease activity in rheumatoid arthritis. J Rheumatol 2001;28:2430–6.
29 Peeters HR, Jongen-Lawrenc M, Vreugdenhil G, et al. Effect of recombinant human erythropoietin on anemia and disease activity in patients with rheumatoid arthritis and anaemia of chronic disease: a randomised placebo controlled double blind 52 weeks clinical trial. Ann Rheum Dis 1996;55:739–44.
30 Cuzzoora S, Mazzeno E, di Pada R, et al. Erythropoietin reduces the degree of arthritis caused by type II collagen in the mouse. Arthritis Rheum 2005;52:940–50.
31 Nairz M, Scholl A, Moschen AR, et al. Erythropoietin contrastingly affects bacterial infection and experimental colitis by inhibiting nuclear factor-kappaB-inducible immune pathways. Immunol 2011;34:61–74.
32 van Santen S, van Dongen-Lases EC, de Vegt F, et al. Heparin and hepcidin content parameters in the diagnosis of iron deficiency in rheumatoid arthritis patients with anemia. Arthritis Rheum 2011;63:3672–80.
33 Sellam I, Korti S, Fellahi S, et al. Serum hepcidin level is not an independent surrogate biomarker of disease activity or of radiographic progression in rheumatoid arthritis: results from the ESPOIR cohort. Ann Rheum Dis 2013;72:312–14.
34 Nikolaisen C, Figenschau Y, Nosset JC. Anemia in early rheumatoid arthritis is associated with interleukin 6-mediated bone marrow suppression, but has no effect on disease course or mortality. J Rheumatol 2008;35:380–6.
35 Pang WW, Schier SL. Anemia in the elderly. Curr Opin Hematol 2012;19:133–40.
36 Chan FK, Lanas A, Scheiman J, et al. Celaecibo versus omeprazole and dicitofenc in patients with osteoarthritis and rheumatoid arthritis (CONDOR): a randomised trial. Lancet 2010;376:173–9.
37 Laine L. Approaches to nonsteroidal anti-inflammatory drug use in the high-risk patient. Gastroenterology 2001;120:594–606.
38 van der Woude D, Syversen SW, van der Voort EJ, et al. The ACPO isotype profile reflects long-term radiographic progression in rheumatoid arthritis. Ann Rheum Dis 2011;70:1461–4.
39 de Vries-Bouwstra JK, Goekoop-Ruiterman YP, Verpooten GN, et al. Progression of joint damage in early rheumatoid arthritis: association with HLA-DRB1, rheumatoid factor, and anti-citrullinated protein antibodies in relation to different treatment strategies. Arthritis Rheum 2008;58:1293–8.
40 Gourraud PA, Boyer IF, Barretche T, et al. A new classification of HLA-DRB1 alleles differentiates predisposing and protective alleles for rheumatoid arthritis structural severity. Arthritis Rheum 2006;54:593–9.
41 Schonland SO, Lopez C, Widdmann T, et al. Premature telomeric loss in rheumatoid arthritis is genetically determined and involves both myeloid and lymphoid cell lineages. Proc Natl Acad Sci USA 2003;100:13471–6.
42 Colmegna I, Diaz-Borjon A, Fuji H, et al. Defective proliferative capacity and accelerated telomeric loss of hematopoietic progenitor cells in rheumatoid arthritis. Arthritis Rheum 2008;58:990–1000.