A Swedish register-based, long-term inception cohort study of patients with rheumatoid arthritis – results of clinical relevance

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Purpose: At the end of the twentieth century, the outcome of rheumatoid arthritis (RA) was shown to be unsatisfactory and new therapeutic strategies were introduced. This initiated a register-based long-term study of early RA, the Better Anti-Rheumatic PharmacOTherapy (BARFOT) study. The aims were to evaluate the disease course and to acquire knowledge for improved care.

Patients and methods: BARFOT is a multicentre observational study of patients with early RA, consecutively included 1992–2006. The patients are followed in daily practice according to a structured protocol for 15 years and data recorded in a web-based register. Also, through linkage of the BARFOT register to national registers we have acquired information on comorbidity and mortality.

Results: In all, 2857 patients have been included and over 80 scientific articles have been published. Phenotypic characteristics at disease onset, i.e. gender, smoking habits and autoantibody profiles have been addressed. The disease course over 15 years was described. Early predictors for persistent disease activity, impaired function, joint damage and comorbidities have been identified. Treatment strategies have been studied. A randomized sub-study gave strong support for the treatment of recent RA with low-dose prednisolone in combination with disease-modifying anti-rheumatic drug. Furthermore, the impact of lifestyle factors, such as smoking, alcohol consumption, body weight and physical activity has been addressed.

Conclusion: A register-based study like BARFOT has provided a basis for optimal long-term management of patients with RA. In addition, the register has made it possible to perform a diversity of studies of RA addressing various issues of major relevance to the patients.

Keywords: patient reported outcomes, PRO, disease progression, prognostic factors, lifestyle, observational study, registry

Introduction

Toward the end of the twentieth century, it became increasingly realized that rheumatoid arthritis (RA) was a disease with a high incidence of disability and increased mortality, which brought about major efforts to improve treatment. Treatment strategies turned from “go low and slow” to “early and active” with the aim that this would improve the disease course and outcome. This assumption was supported by clinical trials, but nothing was known about long-term efficacy. Therefore, to respond to this important unmet need, we initiated a longitudinal
observational study including all available incident RA patients to assess treatment and outcome for several years.

In 1992, these considerations resulted in Better Anti-Rheumatic PharmacOTherapy (BARFOT), a long-term observational cohort study of patients with RA engaging six rheumatology centers in southern Sweden. All data were assembled in a register, which was the first in Sweden to enable regular documentation of different aspects of RA and became the prototype for the national Swedish Rheumatology Quality Register (initially named RAMONA- RA monitoring and assessment), which started in 1995.

Unlike other RA registers, the BARFOT register focused on providing information of clinical importance for the patient. Thus, the BARFOT study pays attention to most facets of the disease, besides measures of disease activity and damage also physical function, pain and quality of life.

Through the start of the BARFOT register, it became possible for the participating rheumatologists to follow the disease course in detail in daily practice and thus make well-founded interventions. The structured data collection admitted studying several aspects of disease outcome and predictors. Also, through linkage of the BARFOT register to different national registers we have been able to add information about comorbidity and mortality. Here, some of the results will be reviewed, particularly those with relevance for clinical practice.

Patients and methods
Patients
The patients were consecutively enrolled in the BARFOT multicentre study during 1992–2006. Inclusion criteria were disease duration of 12 months or less, fulfilling the ACR 1987 classification criteria.1 All patients gave their written informed consent to the study, which was performed in accordance with the Helsinki Declaration. The following ethics committees approved the study: Lund university LU 154-95 and 398-01; Göteborg university Gbg M 45-95 and Ö 282-01; Linköping university Li 123-95 and 01-263; Karolinska Institutet KI 153-95 and 02-075; Stockholm EPN 2011/381-31/4 and 2016/297-31/1, all in Sweden.

The BARFOT protocol
A structured protocol was developed by the collaborating rheumatology units in close co-operation with international experts working with newly proposed outcome measures. The patients were assessed at inclusion and at predefined follow-up visits up to 15 years. Besides the fixed times there was an option for closer registrations if required. All units collected the data in a common database. Treatment decisions were made by the responsible rheumatologist except in limited randomized substudies.

At inclusion, patient characteristics such as age, gender, education, social status, smoking habits (current, previous or never smokers), menopausal age, comorbidity and medication and disease duration were registered.

Information of the regular clinical monitoring and radiographic assessments are given in Table 1.

Biomarkers
At all predefined visits, serum samples were biobanked until analyzed. Among later analyses are anti-cyclic citrullinated peptide (anti-CCP2) antibodies (Euro-Diagnostica, Malmö, Sweden) and anti-carbamylated protein antibodies (anti-CarP) (in-house ELISA, Leiden).

Data enrichment by questionnaires
In 2010 and 2017, questionnaires were sent to all patients remaining in the study (2102 and 1542, respectively), with questions concerning lifestyle factors, such as smoking, alcohol habits, diet and physical activity, as well as pain, health-related quality of life and comorbidity.

Data enrichment by register linkage
To secure information about comorbidities and mortality, linkage of the BARFOT register to national health registers, such as the National Patient Register, including data on hospital discharges, and Cause of Death Register, has been performed. Comparisons with the general population have been made through linkage with the national demographic registers in Sweden.

Major results
Disease onset
From 1992 to 2006, a total of 2837 patients were included. Patient and disease characteristics at baseline are shown in Table 2.

Role of autoantibodies
The BARFOT studies have investigated the role of autoantibodies for the clinical profile of the disease. Thus, the most recent study revealed that the proportion of patients with a positive test for rheumatoid factor (RF), anti-CCP2 and anti-CarP was lower in patients with a higher age at disease
An increasing number of these autoantibodies at onset was associated with younger age, current smoking and higher erythrocyte sedimentation rate (ESR). Moreover, ACPA (CCP2) was found to be associated with osteopenia in the femoral neck and/or Ward’s triangle at disease onset, independently of inflammation.

Table 1 Clinical, laboratory and radiographic assessments according to the BARFOT protocol

<table>
<thead>
<tr>
<th>Assessment</th>
<th>Measurement variable</th>
<th>Measurement method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease activity</td>
<td>Patient’s global assessment (PatGA)</td>
<td>VAS (range 0–100 mm, best to worse)</td>
</tr>
<tr>
<td></td>
<td>Patient’s assessment of pain</td>
<td>VAS (range 0–100 mm, best to worse)</td>
</tr>
<tr>
<td></td>
<td>Physician’s global assessment</td>
<td>Five-grade Likert scale</td>
</tr>
<tr>
<td></td>
<td>Number of tender joints</td>
<td>Calculated on 28 joints</td>
</tr>
<tr>
<td></td>
<td>Number of swollen joints</td>
<td>Calculated on 28 joints</td>
</tr>
<tr>
<td></td>
<td>Acute phase response</td>
<td>Erythrocyte sedimentation rate (ESR)</td>
</tr>
<tr>
<td></td>
<td>Acute phase response</td>
<td>C-reactive protein (CRP)</td>
</tr>
<tr>
<td></td>
<td>Disease activity score, DAS28</td>
<td>VAS PatGA, swollen and tender joint count, ESR DAS28&lt;2.6</td>
</tr>
<tr>
<td></td>
<td>Disease remission</td>
<td></td>
</tr>
<tr>
<td>Function</td>
<td>Disability</td>
<td>Swedish version of the Health Assessment Questionnaire (HAQ)</td>
</tr>
<tr>
<td></td>
<td>Physical impairment</td>
<td>The Signals of Functional Impairment (SOFI)</td>
</tr>
<tr>
<td>Joint damage</td>
<td>Radiographs of the hands and feet</td>
<td>Van der Heijde modification of the Sharp score (SHS), with total, erosion and joint space narrowing scores. Larsen score</td>
</tr>
<tr>
<td>Biomarkers</td>
<td>Autoantibodies</td>
<td>Rheumatoid factor (RF) test</td>
</tr>
<tr>
<td></td>
<td></td>
<td>anti-CCP2 test</td>
</tr>
</tbody>
</table>

Abbreviation: DAS28, 28-joint disease activity score.

Table 2 Characteristics of the patients at inclusion divided by gender

<table>
<thead>
<tr>
<th>Numbers</th>
<th>All patients</th>
<th>Women</th>
<th>Men</th>
</tr>
</thead>
<tbody>
<tr>
<td>Numbers</td>
<td>2837</td>
<td>1916</td>
<td>921</td>
</tr>
<tr>
<td>Age (years)</td>
<td>58 (16)</td>
<td>57 (16)</td>
<td>61 (14)</td>
</tr>
<tr>
<td>Disease duration (months)</td>
<td>8 (11)</td>
<td>8 (11)</td>
<td>8 (13)</td>
</tr>
<tr>
<td>Smoking habits</td>
<td>Never (%)</td>
<td>41</td>
<td>49</td>
</tr>
<tr>
<td>RF present (%)</td>
<td>60</td>
<td>61</td>
<td></td>
</tr>
<tr>
<td>Ever (%)</td>
<td>58</td>
<td>58</td>
<td></td>
</tr>
<tr>
<td>anti-CCP2 present (%)</td>
<td>5.25 (1.26)</td>
<td>5.30 (1.26)</td>
<td></td>
</tr>
<tr>
<td>DAS28 (0–9.4)</td>
<td>45 (26)</td>
<td>47 (26)</td>
<td>41 (25)</td>
</tr>
<tr>
<td>PatGA (VAS, 0–100 mm)</td>
<td>36 (26)</td>
<td>36 (26)</td>
<td>36 (26)</td>
</tr>
<tr>
<td>ESR (mm/h)</td>
<td>32.4 (37.0)</td>
<td>30.5 (36.3)</td>
<td>36.5 (38.3)</td>
</tr>
<tr>
<td>CRP (mg/L)</td>
<td>46 (24)</td>
<td>48 (24)</td>
<td>43 (24)</td>
</tr>
<tr>
<td>Pain (VAS, 0–100 mm)</td>
<td>1.02 (0.65)</td>
<td>1.10 (0.65)</td>
<td>0.87 (0.61)</td>
</tr>
<tr>
<td>HAQ (0–3)</td>
<td>8 (6)</td>
<td>8 (6)</td>
<td>9 (6)</td>
</tr>
<tr>
<td>SOFI (0–44)</td>
<td>1.5 (3.5)</td>
<td>1.4 (3.2)</td>
<td>1.8 (4.0)</td>
</tr>
<tr>
<td>ES</td>
<td>3.5 (6.6)</td>
<td>3.6 (6.8)</td>
<td>3.2 (6.1)</td>
</tr>
<tr>
<td>JSN</td>
<td>5.0 (9.0)</td>
<td>5.0 (9.0)</td>
<td>5.0 (9.0)</td>
</tr>
<tr>
<td>SHS total</td>
<td>6.6 (4.0)</td>
<td>6.6 (4.0)</td>
<td>6.6 (4.0)</td>
</tr>
<tr>
<td>Note: Values are means (SD) unless indicated otherwise.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: RF, rheumatoid factor; DAS28, 28-joint disease activity score; PatGA, patient’s global assessment; ESR, erythrocyte sedimentation rate; CRP, C-reactive protein; HAQ, Health Assessment Questionnaire; SOFI, Signals of Functional Impairment; ES, erosion score; JSN, joint space narrowing score; SHS, modified Sharp/van der Heijde score.

onset. An increasing number of these autoantibodies at onset was associated with younger age, current smoking and higher erythrocyte sedimentation rate (ESR). Moreover, ACPA (CCP2) was found to be associated with osteopenia in the femoral neck and/or Ward’s triangle at disease onset, independently of inflammation.
The presentation of RA has changed
To examine if the presentation of RA had changed over the
years, we compared patients with a disease onset 1992–
1999 with those included 2000–2006. Patients included in
the 2000s were older, had a shorter mean disease duration
at inclusion, were more often RF positive, had higher
disease activity score (DAS28), disability measured by
Health Assessment Questionnaire (HAQ), pain measured
by VAS pain and patient’s global assessment (PatGA) as
well as a higher total van der Heijde modification of the
Sharp score, the latter mainly dependent on a higher joint
space narrowing score. The proportion of smokers were
higher in 1996, 29%, than in 2004, 20%.6

At inclusion, the patients addressed their beliefs about
the cause of their RA. Thirty percent of the patients
reported some possible causal event, most frequently
infection, linked to a seasonal trend in the onset of RA.7
The role of infections as possible triggers was also sug-
gested by a two-cohort study, which showed that onset of
RA in the Swedish patients was more frequent in winter
than in other seasons, in contrast to the Dutch cohort.8

Living with RA
Remission
The goal for treatment in RA is remission. Since several sets
of criteria for remission have been proposed, we tried to find
the most appropriate criterion for long-term studies. To that
end, we studied the frequency of achieved remission defined
by three commonly used sets of criteria in 700 patients
included in BARFOT during the 1990s and followed for 8
years.9 Remission was determined by DAS28 remission
criterion (DAS28 Cr), the Simplified Disease Activity
Index criterion (SDAI Cr)10 and the Boolean-based ACR/
EULAR remission criteria (Boolean Cr).11 Remission at all
follow-up visits, i.e. sustained remission, was infrequent,
most common by the DAS28 Cr, 14%, followed by the
SDAI Cr, 5%, and the Boolean Cr, 3%. Progression of radi-
ographic joint damage was least pronounced in patients with
sustained remission, independently of criterion used. Sustained remission was also associated with improved func-
tion, irrespective of criteria.9

Persistently active disease
Patients who did not achieve DAS28 Cr remission at any
visit up to 8 years were defined as having persistently
active disease which afflicted as many as 37% (43%
women vs. 25% men).12 These patients had significantly
greater PatGA, pain, HAQ and radiological damage com-
pared to the non-persistent group.

Living with persistent disease
An interview study focused on the life situation of ten
patients with persistent disease. Living with persistent
RA was revealed as an existence dominated by painful
symptoms, radical changes and limitations in life, a con-
tinual struggle to cope with life and to master the illness,
and a dependency on those who are close by and the world
around.13 This qualitative study nicely conforms with the
quantitative data described above and is a further argument
for great attention and improved care for this group of
patients with a severe disease course.

Pain
Pain in patients with RA is the most common and trouble-
some complaint, which causes functional impairment and
reduced quality of life. The 2010 postal questionnaire
addressed among other things also pain. As many as
34% of the responders reported chronic wide-spread
pain, more often women, a pain related to DAS28 and
HAQ, and less distinctly to acute phase reactants, i.e., ESR
and C-reactive protein.14

In another study, unacceptable pain, i.e. pain VAS 40 mm
or more, was present in 34% of the patients after 15 years. Of
these, 33% were in remission suggesting that many patients
had pain not directly related to inflammation.15

Physical function and disability
In most patients, physical function assessed by HAQ
improves during treatment with disease-modifying anti-
rheumatic drugs (DMARDs). However, after 8 years, the
patients with persistently active disease had significantly
higher frequency of disability (HAQ≥1.0) than the rest of
the patients (52% vs. 18%).12 Furthermore, we have recently shown that improvement in physical function
does not always follow a reduction in disease activity,
suggesting that impaired physical function may also
depend on other factors.16 The Signals of Functional
Impairment performance test measuring hand, shoulder/
arm and lower extremity function was shown to be a
valuable complement to self-reported function.17

Comorbidities
RA is associated with an enhanced risk of cardiovascu-
lar disease (CVD) and mortality. Whether prompt control of
the inflammatory process after disease onset could
improve CVD and mortality outcome was studied in
patients recruited 1993–1999 and followed until 2010. Incident CVD morbidity and mortality data were obtained through register linkage. In summary, we found that a better control of inflammation during the critical first 2 years of disease resulted in cardiovascular and survival benefits. Predictive factors for morbidity and mortality differed by age at disease onset, suggesting a need for age stratification in the evaluation of CVD and mortality risks in early RA.\(^5\)

**Disease course in patients with disease onset in the 1990s and in the 2000s**

In 1999, biologic DMARDs were introduced in Sweden, which inspired us to compare the disease course in patients included before and after this change in treatment options. Over 5 as well as 8 years, the DAS28 decreased significantly more in patients included in the 2000s compared with those included in the 1990s. Of interest, there was no significant difference in improvement in pain or HAQ between the groups, despite the treatment differences.\(^5\)

**Gender differences**

In several of the BARFOT studies, women were found to suffer a more severe disease course than men. In the cohort included in the 1990s, women had compared with men, higher DAS28 and HAQ, both at inclusion and after 2 years.\(^19\) After 2 years, 28% of the women and 40% of the men were in DAS28 remission.\(^19\) After 5 years, the remission rates were 31% for women and 52% for men.\(^20\) Of the women, only 10% achieved sustained remission over 8 years compared to 23% of the men.\(^9\)

Even though women had higher DAS28, HAQ and VAS pain at all visits up to 5 and 8 years compared to men, the radiological scores did not differ between genders at any timepoint.\(^21,22\)

Also, when comparing the patients included in the 1990s with those included in the 2000s the gender differences were significant. At the 5-year follow-up, the remission rate had increased in the group included in the 2000s (women 42% vs. men 64%).\(^5\)

**Prediction of disease outcome**

The disease course and outcome of RA may vary from spontaneous remission to a persistently active disabling disease. Since treatment options today are manifold with varying toxicity, the key issue in early management of RA is adequate prediction of the long-term outcome.

To address this, we performed a series of studies to find early predictors of later radiological damage. Thus, antiflaggrin (which equals anti-CCP1),\(^23,24\) serum cartilage oligomeric matrix protein,\(^25\) survivin (a proto-oncogene),\(^26\) and hand bone loss by digital X-ray radiogrammetry,\(^27\) independently predicted radiological progression after a disease course of 2 and 5 years. We could confirm that anti-CCP2 is a most reliable laboratory predictor of radiological damage.\(^28\) The combination of anti-CCP2 with anti-CarP autoantibodies proved to have additional predictive potency.\(^29\) However, in our nested randomized study, RF and anti-CCP2 did not predict progressive joint damage in patients treated with prednisolone in combination with DMARDs.\(^30\)

However, it is at least equally important to find predictors of the clinical course of RA. In a study over 8 years we found that as many as 37% of the patients suffered a persistently active disease which was independently predicted at onset by female gender, current smoking and disease activity and absence of remission at 6 months.\(^12\) In a collaboration work involving three European RA cohorts, we found that the presence of RF at disease onset was associated with an increased overall mortality, especially deaths related to neoplasm and respiratory disease, while anti-CCP2 was associated with an increase in cardiovascular deaths.\(^31\)

**Lifestyle factors and disease outcome**

Lifestyle factors like smoking, alcohol consumption, body weight and physical activity influence not only cardiovascular health but may also have impact on the disease course and outcome of RA.

**Smoking**

To assess the effects of smoking on disease outcome, 1787 patients, of which 23% of the women and 24% of the men were current smokers, were studied over the first year of the disease. Smoking status was not associated with disease activity at inclusion, but after 1 year smokers had higher disease activity assessed by DAS28 and lower remission rate compared with previous smokers and non-smokers, despite more DMARD treatment during the first 3 months.\(^6\) In the year 2010, the change in smoking habits and its influence on disease activity was assessed by a postal survey. The proportion of smokers had decreased to 17% 6–14 years after diagnosis. However, smoking cessation after the onset of RA did not influence disease activity.\(^32\)
Current smoking was, in a nested control study over the first 2 years of the disease, found to be associated with subsequent development of severe extra-articular RA, most frequently interstitial lung disease and pleuritis. In addition, current smoking was associated with high disease activity, disability and positive RF.33

The influence of smoking on joint damage was addressed in a six-cohort multinational study. In this study, smokers had more severe joint damage on radiographs of hands and feet 1, 2 and 5 years after diagnosis compared to non-smokers. However, this effect was lost when adjusting for anti-CCP2 positivity. This indicates that the effect of smoking on joint damage could be mediated via the effect of ACPA.34

In a follow-up study, with data on three autoantibodies, RF, anti-CCP2 and anti-CarP, smoking was associated with concurrent presence of two or three autoantibodies rather than anti-CCP2 alone. This implies that smoking is a risk factor for breaking tolerance to multiple autoantigens in RA.35

Given the negative effects of smoking on RA described above, we further studied if second-hand exposure to tobacco smoke affected disease activity in patients who had never smoked. No association was found between second-hand exposure and disease activity.36

Alcohol consumption
In 2010, 1238 BARFOT patients answered the questionnaire about alcohol consumption. Of all, 67% were non-hazardous drinkers and 21% were classified as hazardous drinkers.37 Alcohol consumption in women, but not in men, was associated with lower self-reported disease activity and better health-related quality of life than non-drinkers. Current smoking emerged as an independent negative prognostic factor for reported disease activity, which might have counteracted the immunosuppressive effect of alcohol.37

The role of alcohol consumption in RA is complex. On the one hand, alcohol may lower the risk for getting RA,38 and reduce inflammation in established RA,39 on the other hand, alcohol may enhance hepatotoxicity of DMARDs. Discussions regarding alcohol consumption are thus important to make patients aware of its interactions with medication.

What about changes in alcohol habits after diagnosis in our patients? Based on the questionnaire of 2010 it was found that 6% of the patients had stopped drinking. By qualitative content analyses, we found that patients who stopped drinking were older, had lower health-related quality of life, more pain and fatigue and worse physical function compared with those who continued drinking.40 The reasons for stopping drinking alcohol varied but was most often illness. Another reason was a desire to improve health and well-being and, less frequently, the patients referred to work and solidarity with family members.40

Body weight
In 1596 patients from the BARFOT cohort, 13% of the patients were obese at disease onset (body mass index [BMI] ≥30 kg/m²) and 16% in the survey 9 years later.41 Obesity was independently associated with higher disease activity, worse HAQ score, more pain and worse PatGA both at disease onset and at the time of the survey. Also, obesity was independently associated with important comorbidities, such as hypertension, diabetes mellitus and chronic pulmonary disease.41

Interestingly, if the cutoff for obesity was set to ≥28 kg/m², the associations to disease severity gave similar results.41 This implies that unfavorable body composition could be more accurately detected by redefining BMI cutoffs in RA. Further, a higher waist circumference was also associated with a worse disease outcome and could be used in the RA population besides BMI to aid in the prediction of the disease prognosis.41

Physical activity
According to the 2010 questionnaire, 65% of the BARFOT patients met the WHO recommendations for health-enhancing physical activity, women to a larger extent than men,43 virtually the same as in a non-rematoid population in Sweden. Meeting recommended levels of physical activity could not be predicted by function or pain at disease onset, which makes it important to recommend a healthy lifestyle along the course of the disease.

Lifestyle discussions
It is important and recommended to make patients aware of the influence of lifestyle on RA. Based on the 2017 questionnaire it was found that, according to the patients’ recall, smoking had been discussed with 25% of the patients, alcohol consumption with 17%, diet with 23% and physical activity with 49%. Although these figures are higher than previously reported, there is a need for further improvement to comply with the EULAR recommendations of patient-centered standards of care.
Treatment with low-dose prednisolone

An important but controversial topic in the treatment of early RA is use of prednisolone, which gave rise to our randomized prednisolone study. A total of 250 patients were randomized to 7.5 mg prednisolone daily or no prednisolone for 2 years at the start of their first DMARD. We found that the prednisolone-group had lower joint damage progression and higher remission rate than the non-prednisolone group and the frequency of adverse effects was small. A follow-up study demonstrated that remission after 2 years with prednisolone was associated with reduced joint destruction also after 4 years. In a further study, we concluded that the inhibition of progression of joint destruction by prednisolone might depend on decreased activity of matrix metalloproteinases as well as impaired osteoclast activation.

Prednisolone in this low dose had no or minor effects on bone density. Measurement of bone-markers revealed that the suppressive effect on bone synthesis exerted by prednisolone was counteracted by the ability of prednisolone to hamper the inflammatory mediated increase in bone resorption. Concerning cardiovascular risks, treatment with low-dose prednisolone did not influence the thickness or endothelial function of the carotid intima after 5 years. After 10 years, the incidence of ischaemic coronary artery events was similar in the two treatment groups, whereas the long-term risk for incident cerebrovascular events was somewhat higher in the prednisolone group.

Discussion

The primary goal of the BARFOT study was to create a basis for improved treatment of patients with RA. The information acquired by the register served as guidance to clinical practice, primarily with regard to tailored anti-inflammatory treatment. In addition, the structured protocol allowed further studies, which have contributed to increase the knowledge of this complex disease.

RA has been recognized as a separate disease entity but is indeed a multi-faceted disease, in which the identification of novel RA-related autoantibodies has made it possible to distinguish subphenotypes. The present studies, including the recent multi-national cohort studies, suggest that the magnitude of immune response at disease onset influences the clinical presentation of the disease and may have implications for diagnosis and prognosis of RA at different ages. The data also support the hypothesis of a pathophysiological role of ACPA in systemic bone loss.

Although infections with periodontal pathogens and Epstein-Barr virus have been shown to induce autoimmunity and ACPA-production, the prevalence and possible pathogenetic role of common viral infections in the pathogenesis of RA has not yet been established. Many patients believe that an infection might be a possible causal event, linked to onset of RA in the winter season. However, this could not be verified in our dual cohort study, in which joint destruction over time did not differ between patients with disease onset in winter or summer.

A key focus of the BARFOT studies has been to increase knowledge about the disease course from the patient’s perspective. We have shown that despite improved treatment options, a high proportion of patients suffer from a persistently active disease, pain, disability and impaired health over many years after disease onset. Pain emerged as the major complaint, frequently unacceptable, and not always linked to active disease but also present in remission, stressing the need of improved pain assessment and treatment.

There is agreement concerning the importance of achieving remission, but various remission definitions exist. The DAS28 remission criterion, used in BARFOT, identified more patients in sustained remission than did the other applied criteria with no difference in degree of joint damage and disability. This suggests that this criterion may cause fewer misclassifications and may therefore be used in long-term cohort studies until more accurate criteria are available.

Despite modern treatment strategies, many of the BARFOT patients had a persistently active disease. Importantly, several recent studies similarly report a substantial proportion of patients with persistent disease activity in spite of currently available effective treatment options. This emphasizes the importance of very early and persistent management and care of the patients.

The BARFOT studies stress the fact that pain in RA is a major distress causing impaired function and is associated with increased risk of poor outcome, and it also contributes to the global burden of the disease. In addition to inflammation, pain may also be due to other causes. These observations imply that the origin of pain in patients with RA should be identified and adequate treatment given, which has been emphasized in the recent EULAR recommendations for pain management.
Like pain, function did not improve in all patients following a satisfactory reduction in disease activity.\textsuperscript{16} This is in line with other reports, reviewed in Norton et al.\textsuperscript{57} It is thus vital to be aware of the fact that functional impairment as well as pain may have causes other than inflammation and therefore may remain also following successful control of the inflammation.

The higher risk of CVD seen in patients with RA is not fully explained by traditional CVD risk factors,\textsuperscript{58} moreover also inflammation plays a role. Thus, inflammatory cytokines in RA stimulate plaques formation and rupture in CVD.\textsuperscript{59} The role of inflammation for CVD is supported in our studies in which prompt control of inflammation during the critical first 2 years of disease resulted in cardiovascular and survival benefits providing a rationale for our treatment approach to prevent CV in patients with RA.\textsuperscript{18}

Our data suggest that RA is more serious in women than in men.\textsuperscript{12} However, Sokka et al reasoned that this gender difference is valid only for patients with more limited disease activity reflecting “considerable differences in the measures between genders”.\textsuperscript{60} In accordance with that report, the observation of a significant overrepresentation of women among patients with long-standing active disease indicates true worse outcome as to components included in the disease activity measures but not radiologic destruction.\textsuperscript{21,22}

Lifestyle factors like smoking influenced disease severity and development of extra-articular disease, while obesity was associated with high disease activity and comorbidities.

Among lifestyle factors, cigarette smoking is established to increase severity and comorbidity in RA. In addition, during the last years obesity and low physical activity have also been found to affect the disease negatively.\textsuperscript{51–65} All these factors are potentially intervenable, and consequently smoking cessation, weight control and physical activity have been included in the EULAR recommendation for management of early arthritis.\textsuperscript{66}

Reliable predictors of the disease course and outcome are of paramount importance for treatment precision. We have detected predictors for persistently active disease, joint damage and comorbidities. Some of these have been performed as joint studies with other groups, while in others linkage was done with national health registers.

Through the year’s rheumatologists have tried to identify predictors of the future course of early RA. On a group level, we and others have found baseline variables predicting clinical as well as radiological outcome. However, to our knowledge, no single predictor or prediction model has been proven to identify the disease course in an individual patient. Accordingly, a recent literature review found that “No predictors were identified reliably predicting clinical response to methotrexate after 3–6 months in the individual patient: clinical predictors were weak.”\textsuperscript{67} Furthermore, despite much research, a significant number of patients in the present era of biologic DMARDs yet develop refractory disease.\textsuperscript{68} Consequently, there is still an urgent need for improved early prediction of disease outcomes. We will now focus on the immune response to joint specific proteins by identification of unique peptides which can be recognized by antibodies by a specific pathogenic or regulatory function.

The clinical message from the randomized studies of treatment with low-dose prednisolone in combination with synthetic DMARDs was that low-dose prednisolone may be recommended for at least 2 years to patients with recent onset RA. This recommendation is in line with that of the 2019 update of the Swedish Society for Rheumatology RA treatment guidelines. Furthermore, a recent “Systematic Review of International Guidelines and Consensus Statements” reports that most current recommendations of glucocorticoid use in RA are similar to those proposed by our group.\textsuperscript{69}

To conclude, long-term observational studies on RA like BARFOT may serve the dual purpose of providing a basis for optimal long-term management of the patient and of creating a platform for a diversity of studies of the disease which otherwise would have been difficult or impossible to achieve.

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Disclosure

The authors report no conflicts of interest in this work.

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