Diabetes incidence in psoriatic arthritis, psoriasis and rheumatoid arthritis: a UK population-based cohort study

Maureen Dubreuil¹, Young Hee Rho², Ada Man¹, Yanyan Zhu², Yuqing Zhang², Thorvardur Jon Love³,⁴, Alexis Ogdie⁵, Joel M. Gelfand⁶ and Hyon K. Choi¹,²

Abstract

Objective. The objective of this study was to evaluate the incidence of diabetes among patients with PsA and RA in the general population.

Methods. We conducted a cohort study using an electronic medical records database representative of the UK general population (1986–2010). We estimated hazard ratios (HRs) for incident diabetes in PsA, psoriasis and RA cohorts compared with age- and sex-matched comparison cohorts without the corresponding conditions, adjusting for BMI, smoking, alcohol use, co-morbidities and glucocorticoids at baseline.

Results. Cohorts included 4196 persons with PsA, 59,281 with psoriasis and 11,158 with RA, with mean follow-up times of 5.9, 5.8 and 5.5 years, respectively. Incidence rates for diabetes were 7.3, 6.4 and 6.3 cases per 1000 person-years among individuals with PsA, psoriasis and RA, respectively. Age- and sex-matched HRs for diabetes were 1.72 (95% CI 1.46, 2.02) in PsA, 1.39 (95% CI 1.32, 1.45) in psoriasis and 1.12 (95% CI 1.01, 1.25) in RA. After adjustment for BMI, smoking and alcohol, the HRs were attenuated substantially (1.43, 1.24 and 1.00, respectively). With further adjustment for baseline glucocorticoid use and co-morbidities, the HRs were 1.33 (1.09, 1.61) in PsA, 1.21 (1.15, 1.27) in psoriasis and 0.94 (0.84, 1.06) in RA.

Conclusion. This general population study suggests an increased incidence of diabetes in PsA and RA, which is substantially explained by obesity and lifestyle factors. These findings support the importance of managing such factors in PsA and RA patients.

Key words: psoriatic arthritis, psoriasis, rheumatoid arthritis, diabetes.

Introduction

PsA and RA are both chronic, progressive and destructive joint diseases that are associated with systemic inflammation and often lead to significant complications. Diabetes risk may be elevated due to inflammatory arthritis, although studies of diabetes in RA patients have had inconsistent results [1, 2] and diabetes risk, specifically in PsA, has not been previously reported. As psoriatic skin disease (PsO) is associated with diabetes, independent of obesity and other traditional diabetes risk factors [3, 4], diabetes risk in PsA may be elevated due to the systemic inflammatory burden from both skin and joint disease. Furthermore, the prevalence of obesity in PsA patients is greater than in PsO or RA patients, conferring diabetes risk [5]. Therefore, due to underlying systemic inflammation and obesity, we suspected diabetes risk in PsA may be greater than that seen in inflammatory skin disease or joint disease alone. To address these issues, we evaluated the risk of incident diabetes among persons with PsA and RA in a large UK cohort in which data on obesity and lifestyle variables have been collected. To demonstrate

¹Department of Rheumatology, Boston Medical Center. ²Clinical Epidemiology Research and Training Unit, Boston University School of Medicine, Boston, MA, USA. ³Department of Immunology and Centre for Rheumatology Research, Landspítali University Hospital, Fossavogur. ⁴Faculty of Medicine, University of Iceland, Reykjavík, Iceland. ⁵Division of Rheumatology and ⁶Department of Dermatology, University of Pennsylvania, Philadelphia, PA, USA.

Submitted 24 January 2013; revised version accepted 4 September 2013.

Correspondence to: Maureen Dubreuil, Section of Rheumatology and the Clinical Epidemiology Unit, Boston University School of Medicine, 650 Albany Street, Suite 201, Boston, MA 02118, USA. E-mail: maureen.dubreuil@bmc.org
the previously shown association with PsO (as a positive control) [4, 6], we also analysed diabetes risk among individuals with PsO.

Methods

Data source

The Health Improvement Network (THIN) is a database of electronic medical records in which general practitioners (GPs) in the UK have systematically recorded data on approximately 7.3 million patients. Because >92% of the British population is registered with a GP [7, 8], THIN is a population-based cohort representative of the UK general population. THIN includes demographics, details from GP visits, specialist referrals and hospital admissions, results of laboratory tests and additional health information. The Read code classification is used to identify specific clinical diagnoses, and a drug dictionary based on the Multilex classification is used to code medications [9-11]. Health information is recorded with quality control procedures to maintain high data completion rates and accuracy. Thus THIN data represent routine medical practice in a population-based setting.

Study design and cohort definitions

We conducted a cohort analysis of incident diabetes among adults with incident PsA (PsA cohort) compared with age-, sex- and entry year-matched individuals without PsA (comparison cohort) using data from THIN. We conducted equivalent cohort analyses among individuals with incident PsO and RA. Participants were required to be continuously enrolled in the database for 12 months prior to inclusion in the cohort, and those with diabetes prior to study entry were excluded.

PsA and PsO were defined using the earliest diagnostic code for the respective condition using the Read classification [10,11], definitions previously found to have positive predictive values (PPVs) of 85% and 90%, respectively, in THIN [12,13]. Secondary definitions of PsA and PsO included at least two diagnostic codes for the respective conditions.

Our primary definition for RA required one RA diagnostic Read code followed by at least one prescription for a DMARD using THIN drug codes based on the Multilex classification [9]. This RA definition has been found to have a specificity of 96% (vs the 1987 ACR criteria) [14]. Secondary definitions of RA included (i) a single diagnostic record for RA and (ii) at least two visits with RA diagnoses [2].

For the comparison cohorts corresponding to each exposure cohort, we matched up to 10 individuals without the corresponding exposure condition to each exposed subject based on age, sex and year of study entry. Our study period was 1 January 1986 through 31 May 2010. Participants entered the cohort when all inclusion criteria were met or the matched date for subjects in the comparison cohorts. Participants were followed until they developed diabetes, died or the follow-up ended, whichever came first.

Diabetes mellitus outcome assessment

Our outcome of interest was development of diabetes requiring medication treatment. Diabetes was defined by a prescription for a medication used for the treatment of diabetes, including all insulin preparations and oral agents, as done previously [2].

Assessment of covariates

Data on demographics, BMI, smoking and alcohol use were collected from the database using the most recent value before study entry. We calculated the Charlson co-morbidity index using Read diagnostic codes and assessed glucocorticoid use during the 12 months prior to diagnosis of the respective exposure condition (RA, PsA or PsO), or the matched date for subjects in the comparison cohorts.

Statistical analysis

We compared the baseline characteristics between the exposure cohorts (i.e. PsA, PsO and RA cohorts) and corresponding comparison cohorts. We identified incident cases of diabetes during follow-up and calculated incidence rates (IRs) for diabetes. We estimated the cumulative incidence of diabetes in each cohort accounting for the competing risk of death [15]. Examination of log-log survival curves for each of the exposure variables (RA, PsA and PsO) in our model demonstrated that assumptions of proportional hazards were met. We employed Cox proportional hazard regression models to assess the multivariable-adjusted hazard ratios (HRs) after stratifying by matched clusters (age, sex and calendar year of study entry). We imputed missing values for BMI and smoking using a sequential regression method based on a set of covariates as predictors (IVEware for SAS, version 9.2; SAS Institute, Cary, NC, USA) [16]. To minimize random error, we imputed five datasets and then combined estimates from these datasets. The imputed datasets were used for all primary analyses. Our multivariable analyses were further adjusted for baseline assessments of BMI (in kg/m², using the World Health Organization classification), smoking (non-smoker, current smoker and past smoker), use of oral and topical glucocorticoids, and Charlson co-morbidity index. For all HRs, we calculated 95% CIs. All P-values were two-sided. This study was approved by the UK National Health Service Research Ethics Committee (Protocol 12-007) and was judged exempt from review by the Boston Medical Center Institutional Review Board (Protocol H-31390).

Results

Our primary analyses included 4196 individuals with PsA (50% female, mean age 49 years, mean BMI 27.6), 59281 persons with PsO (51% female, mean age 49 years, mean BMI 26.7), 11158 individuals with RA (68% female, mean age 58 years, mean BMI 26.8) and their matched comparison cohorts (Table 1). Relative to the comparison cohorts, the exposed cohorts had a higher baseline prevalence of overweight or obese status, past or current
Table 1 Baseline characteristics of the study populations

<table>
<thead>
<tr>
<th></th>
<th>PsA (n = 4196)</th>
<th>No PsA (n = 41 794)</th>
<th>P-value</th>
<th>PsO (n = 59 281)</th>
<th>No PsO (n = 581 248)</th>
<th>P-value</th>
<th>RA (n = 11 158)</th>
<th>No RA (n = 110 375)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (S.D.), years</td>
<td>48.7 (13.9)</td>
<td>48.7 (13.8)</td>
<td>0.92</td>
<td>49.2 (17.1)</td>
<td>48.9 (17)</td>
<td>&lt;0.001</td>
<td>58.4 (14.2)</td>
<td>58.2 (14.1)</td>
<td>0.33</td>
</tr>
<tr>
<td>Women, n (%)</td>
<td>2104 (50.1)</td>
<td>20965 (50.2)</td>
<td>0.98</td>
<td>30330 (51.2)</td>
<td>297 675 (51.2)</td>
<td>0.02</td>
<td>7582 (68)</td>
<td>75 093 (68)</td>
<td>0.86</td>
</tr>
<tr>
<td>BMI, mean (S.D.), kg/m²</td>
<td>27.6 (5.6)</td>
<td>26.2 (5.1)</td>
<td>&lt;0.001</td>
<td>26.7 (5.4)</td>
<td>25.9 (5.0)</td>
<td>&lt;0.001</td>
<td>26.8 (5.2)</td>
<td>26.3 (5.0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>&lt;18.5, n (%)</td>
<td>33 (0.8)</td>
<td>580 (1.4)</td>
<td>&lt;0.001</td>
<td>893 (1.5)</td>
<td>9115 (1.6)</td>
<td>&lt;0.001</td>
<td>167 (1.5)</td>
<td>15 464 (1.4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>18.5–24.9, n (%)</td>
<td>1170 (27.9)</td>
<td>13 698 (32.8)</td>
<td>&lt;0.001</td>
<td>18 227 (30.7)</td>
<td>185 857 (32)</td>
<td>&lt;0.001</td>
<td>3522 (31.6)</td>
<td>35 626 (32.3)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>25.0–29.9, n (%)</td>
<td>1152 (27.5)</td>
<td>10 580 (25.3)</td>
<td>&lt;0.001</td>
<td>15 772 (26.6)</td>
<td>140 238 (24.1)</td>
<td>&lt;0.001</td>
<td>3238 (29.0)</td>
<td>29 777 (27.0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>≥30.0, n (%)</td>
<td>948 (22.6)</td>
<td>5653 (13.5)</td>
<td>&lt;0.001</td>
<td>9859 (16.8)</td>
<td>71 193 (12.2)</td>
<td>&lt;0.001</td>
<td>2017 (18.1)</td>
<td>15 919 (14.4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unknown, n (%)</td>
<td>893 (21.3)</td>
<td>11 283 (27)</td>
<td>&lt;0.001</td>
<td>14 530 (24.5)</td>
<td>174 845 (30.1)</td>
<td>&lt;0.001</td>
<td>2214 (19.8%)</td>
<td>27 507 (24.9)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Smoking, n (%)</td>
<td>1967 (46.9)</td>
<td>19 730 (47.2)</td>
<td>&lt;0.001</td>
<td>23828 (40.2)</td>
<td>267 871 (46.1)</td>
<td>&lt;0.001</td>
<td>4730 (42.4)</td>
<td>54 696 (49.6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Never</td>
<td>807 (19.2)</td>
<td>5811 (13.9)</td>
<td>&lt;0.001</td>
<td>10 846 (18.3)</td>
<td>78 779 (13.6)</td>
<td>&lt;0.001</td>
<td>2351 (21.1)</td>
<td>17 516 (15.9)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Past</td>
<td>929 (22.1)</td>
<td>9285 (22.2)</td>
<td>&lt;0.001</td>
<td>16 770 (28.3)</td>
<td>122 223 (21)</td>
<td>&lt;0.001</td>
<td>2798 (25.1)</td>
<td>20 521 (18.6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Current</td>
<td>493 (11.7)</td>
<td>6968 (16.7)</td>
<td>&lt;0.001</td>
<td>7837 (13.2)</td>
<td>112 375 (19.3)</td>
<td>&lt;0.001</td>
<td>1279 (11.5)</td>
<td>17 642 (16)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Alcohol, n (%)</td>
<td>521 (12.4)</td>
<td>4670 (11.2)</td>
<td>&lt;0.001</td>
<td>7041 (11.9)</td>
<td>67 642 (11.6)</td>
<td>&lt;0.001</td>
<td>1870 (16.8)</td>
<td>15 363 (13.9)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Never</td>
<td>49 (1.2)</td>
<td>398 (1.0)</td>
<td>&lt;0.001</td>
<td>726 (1.2)</td>
<td>5376 (0.9)</td>
<td>&lt;0.001</td>
<td>175 (1.6)</td>
<td>1182 (1.1)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Past</td>
<td>2756 (65.7)</td>
<td>25 806 (61.7)</td>
<td>&lt;0.001</td>
<td>37 147 (62.7)</td>
<td>335 973 (57.8)</td>
<td>&lt;0.001</td>
<td>6770 (80.7)</td>
<td>66 374 (80.1)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unknown</td>
<td>870 (20.7)</td>
<td>10 920 (26.1)</td>
<td>&lt;0.001</td>
<td>14 367 (24.2)</td>
<td>172 257 (29.6)</td>
<td>&lt;0.001</td>
<td>2343 (21.0)</td>
<td>27 456 (24.9)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Oral glucocorticoid use, n (%)</td>
<td>343 (8.2)</td>
<td>1096 (2.6)</td>
<td>&lt;0.001</td>
<td>2525 (4.3)</td>
<td>15 354 (2.6)</td>
<td>&lt;0.001</td>
<td>2470 (22.1)</td>
<td>3707 (3.4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Topical glucocorticoid use, n (%)</td>
<td>1475 (35.2)</td>
<td>1808 (4.3)</td>
<td>&lt;0.001</td>
<td>14 844 (25)</td>
<td>23 956 (4.1)</td>
<td>&lt;0.001</td>
<td>900 (8.1)</td>
<td>5493 (5)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Charlson comorbidity index, mean (S.D.)</td>
<td>0.13 (0.38)</td>
<td>0.07 (0.34)</td>
<td>&lt;0.001</td>
<td>0.1 (0.39)</td>
<td>0.08 (0.37)</td>
<td>&lt;0.001</td>
<td>0.21 (0.51)</td>
<td>0.1 (0.41)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

All characteristics were assessed prior to study entry. Unknown refers to subjects with missing values for the variable indicated. P-values were calculated by t-test or Wilcoxon rank-sum test for continuous variables or chi-square test for indicator or categorical variables.
smoking, oral and topical glucocorticoids use and mean comorbidity indices. Patients with PsO and PsA had higher frequencies of current alcohol use than their respective comparison cohorts.

The cumulative incidence of diabetes in each cohort is depicted in Fig. 1. New diagnoses of diabetes occurred among 179 persons during 24,625 person-years (PY) in the PsA cohort, among 2202 persons during 345,507 PY in the PsO cohort and among 383 persons during 61,087 PY in the RA cohort, corresponding to incidence rates of 7.3, 6.4 and 6.3 cases per 1000 PY, respectively (Table 2). As shown in Fig. 1, the risk of diabetes was higher among PsA, PsO and RA subjects than their counterparts, respectively. Relative to the comparison cohorts, the age-, sex- and entry time-matched HRs for diabetes were 1.72 (95% CI 1.46, 2.02) for PsA, 1.39 (95% CI 1.32, 1.45) for PsO and 1.12 (95% CI 1.01, 1.25) for RA. After further adjustment for baseline BMI, smoking and alcohol use, these HRs were attenuated but remained significant in the PsA and PsO cohorts (HRs 1.43 and 1.24, respectively), but not in the RA cohort (HR 1.00). With further adjustment for baseline glucocorticoid use and co-morbidities, the HRs became 1.33 (95% CI 1.09, 1.61) in PsA, 1.21 (95% CI 1.15, 1.27) in PsO and 0.94 (95% CI 0.84, 1.06) in RA.

Sensitivity analyses did not materially change the results, including (i) the use of two Read code definitions for PsA (multivariable HR 1.50, 95% CI 1.10, 2.06), PsO (multivariable HR 1.27, 95% CI 1.17, 1.39) and RA (multivariable HR 0.97, 95% CI 0.85, 1.11) (see supplementary Table S1, available at Rheumatology Online); (ii) a single Read code definition for RA (multivariable HR 1.02, 95% CI 0.93, 1.11) (supplementary Table S1, available at Rheumatology Online); (iii) restriction to only those subjects without missing values for covariates (supplementary Table S2, available at Rheumatology Online); (iv) exclusion of subjects who were ever prescribed oral steroids (supplementary Table S3, available at Rheumatology Online); (v) exclusion of subjects ever prescribed oral or topical steroids (supplementary Table S4, available at Rheumatology Online) and (vi) restriction of RA cases to those after 1995, to evaluate diabetes risk in a more recent context (multivariable HR 0.96, 95% CI 0.84, 1.00; data not shown).

**Discussion**

In this large general practice cohort representative of the UK population, we found that diabetes risk was 72% higher among individuals with PsA than age- and sex-matched individuals without PsA. When we took baseline BMI, smoking, alcohol use and other covariates into account, the association was attenuated, but remained significant (33% increase over the comparison cohort). In contrast, the overall risk of diabetes among individuals with RA was only 12% higher than age- and sex-matched individuals without RA, and this increased risk was nullified after adjustment for lifestyle risk factors. This finding was replicated using alternate definitions of RA and other sensitivity analyses. Taken together, our findings provide the first general population evidence of increased diabetes risk specifically in PsA and the lack of an increased risk in RA after adjustment for obesity and lifestyle factors. These findings suggest that obesity and lifestyle factors partially explain the increased risk of diabetes in PsA and PsO patients. Indeed, in the general population up to 91% of type 2 diabetes is attributable to obesity, smoking and alcohol [17]. However, these three factors have also been associated with the risk of incident PsO, and obesity has been associated with the risk of PsA, which leads to the potential for confounding and the need to adjust for such factors in estimating diabetes risk [18]. These risk estimates were similar to those from a large prospective cohort study (Nurses’ Health Study) in which the risk of diabetes among individuals with PsO was attenuated after

![Fig. 1 Cumulative incidence of diabetes in the PsA, psoriasis and RA cohorts and age-, sex- and cohort entry-matched comparison cohorts.](https://academic.oup.com/rheumatology/article-abstract/53/2/346/1836698)
adjustment for obesity and lifestyle factors (from 2.08 to
1.63) [4].

Similar confounding likely exists in assessing the impact of RA on diabetes risk, because development of RA is more common among smokers and obese individuals [19–21]. Although our unadjusted HR (1.12) of diabetes in RA patients was directionally consistent with the previous Canadian study (crude HR 1.5) [2], risk in our cohort disappeared after adjustment for obesity and lifestyle factors (HR 1.01), whereas the increased risk in the Canadian study did not change after adjustment for available covariates in the insurance claim database (BMI and lifestyle variables unavailable). The HR for diabetes for RA patients in our study was lower than in the Canadian study due to the lower diabetes incidence rate in our RA population (6.3 vs 8.6 cases per 1000 PY), not due to different incidence rates in the comparison populations (5.7 vs 5.8 cases per 1000 PY). Further supporting the absence of any increased risk, the Olmsted County RA cohort reported a crude diabetes incidence rate ratio of 0.78 [1]. Finally, our consistent results from multiple sensitivity analyses provide no support for an independent impact of RA on the risk of diabetes.

The differential risks of diabetes in RA and PsA, despite both being progressive arthritic conditions associated with systemic inflammation, may indicate different mechanisms for the development of diabetes. Although obesity is common in both populations, PsA patients tend to have higher BMIs than RA patients (29.6 vs 27.9), and differences in body composition (central obesity or fat mass) may potentially be associated with differential effects of adipocytokines [5, 22, 23]. Notably, systemic inflammation has also been associated with insulin resistance and diabetes, suggesting an inflammatory basis for diabetes [24]. The arthritic component of PsA may further add to the inflammatory mechanisms due to PsO. However, it remains unclear which, if any, inflammatory factors specific to psoriatic conditions underlie the increased diabetes risk. Therefore, in addition to obesity-associated metabolic factors, immune-mediated inflammatory processes specific to psoriatic conditions could provide the potential links between PsA and diabetes.

The strengths and limitations of our study deserve comment. This study was performed using a large UK general practice database and therefore findings are likely to be generalizable to the general population. Our database is derived from the electronic medical records used for patient care, thus the overall diagnostic accuracy is expected to be high [13, 14, 25], but remains a potential concern. For example, the PPV estimate for PsA based on a relatively small sample size was 85% [12]. However, any non-differential misclassification would have biased the study results toward the null, and our results persisted in sensitivity analyses that used alternate definitions of PsA, PsO and RA. We attempted to define populations with incident PsA, PsO and RA. However, it is possible that some subjects did not truly have incident disease, in which case our diabetes risk estimate would reflect diabetes risk among those with new-onset and long-standing

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Incidence rates and HRs for diabetes among the PsA, PsO, RA and comparison cohorts</th>
</tr>
</thead>
<tbody>
<tr>
<td>PsA</td>
<td>No PsA</td>
</tr>
<tr>
<td>---------</td>
<td>--------</td>
</tr>
<tr>
<td>Diabetes cases, n</td>
<td>258</td>
</tr>
<tr>
<td>Total follow-up time, PY</td>
<td>15,539</td>
</tr>
<tr>
<td>Mean follow-up time, years</td>
<td>5.9</td>
</tr>
<tr>
<td>Incidence rate, per 1000 PY</td>
<td>7.3</td>
</tr>
<tr>
<td>Age-, sex- and entry time-adjusted HR (95% CI)</td>
<td>1.72 (1.46, 2.02)</td>
</tr>
<tr>
<td>+ BMI-, smoking- and alcohol-adjusted HR (95% CI)</td>
<td>1.43 (1.2, 1.7)</td>
</tr>
<tr>
<td>+ Oral and topical glucocorticoids and Charlson index-adjusted HR (95% CI)</td>
<td>1.33 (1.09, 1.61)</td>
</tr>
</tbody>
</table>
PsA, PsO or RA. While there were missing values for covariates in some subjects, it was reassuring that our primary results with imputation closely agreed with those restricted to only those subjects without missing values. Finally, as our study assessed the main links between PsA, PsO, RA and the risk of diabetes, future studies may identify more specific risk factors for diabetes related to these rheumatic conditions.

In conclusion, this large general population study suggests that the overall risk of diabetes is increased in PsA, which is partially explained by obesity and lifestyle factors. Nevertheless, PsA (and PsO) remained associated with the risk of diabetes even after taking these factors into account. In contrast, we found that the risk of diabetes among patients with RA was entirely due to obesity and smoking. Overall, these findings suggest that diabetes risk should not be ascribed only to the presence of inflammatory disease. Future diabetes prevention efforts may be directed at weight loss and smoking cessation, in addition to treatment of the underlying disease, in PsA and PsO patients.

### Acknowledgements

Thanks to Christine Peloquin for statistical assistance.

**Funding:** This work was supported in part by grants from the NIAMS (P60AR047785), National Heart, Blood and Lung Institute (R01 HL089744 to J.M.G.), NIH (K24-AR064310) and Boston University School of Medicine. The funding sources had no role in the design, conduct or reporting of the study or in the decision to submit the manuscript for publication.

**Disclosure statement:** J.G. serves as a consultant to Amgen, Abbott, Centocor, Celgene, Novartis, Merck and Pfizer and has received honoraria. J.G. has received grants from Amgen, Abbott, Pfizer, Novartis and Genentech. All other authors have declared no conflicts of interest.

### Supplementary data

Supplementary data are available at *Rheumatology* Online.

### References


