EXTENDED REPORT

Obesity and the risk of psoriatic arthritis: a population-based study

Thorvardur Jon Love,1 Yanyan Zhu,2 Yuqing Zhang,2 Lindsay Wall-Burns,3,4 Alexis Ogdie,5 Joel M Gelfand,6 Hyon K Choi7

1Department of Science, Education, and Innovation, Landspitali University Hospital, Fossjavogur, Reykjavik, Iceland
2The Clinical Epidemiology Unit, Department of Medicine Boston University School of Medicine, Boston, Massachusetts, USA
3Arthritis Research Centre of Canada, Vancouver, Canada
4Department of Dermatology and Skin Science, University of British Columbia, Vancouver, Canada
5Department of Medicine/Rheumatology, University of Pennsylvania, Philadelphia, Pennsylvania, USA
6Department of Dermatology, University of Pennsylvania, Philadelphia, Pennsylvania, USA
7Section of Rheumatology and the Clinical Epidemiology Unit, Boston University School of Medicine, Boston, Massachusetts, USA

Correspondence to
Hyon K Choi, Section of Rheumatology and the Clinical Epidemiology Unit, Boston University School of Medicine, 650 Albany Street, Suite 200, Boston, MA 02118, USA; hchoius@bu.edu

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ABSTRACT

Background Obesity is associated with an increased risk of psoriasis; however, its potential impact on the risk of psoriatic arthritis (PsA) remains unclear. Objectives To evaluate the association between body mass index (BMI) and the risk of PsA among patients with psoriasis from the general population. Methods The authors conducted a cohort study using data from The Health Improvement Network, an electronic medical records database representative of the UK general population, collected between 1995 and 2010. The exposure of interest was the first BMI measured after psoriasis diagnosis and endpoints were incident cases of physician-diagnosed PsA. The authors estimated the RR of PsA after adjusting for age, sex, and histories of trauma, smoking and alcohol consumption. Results Among 75 395 individuals with psoriasis (43% male, mean follow-up of 5 years, and mean age of 52 years), 976 developed PsA (incidence rate, 26.5 per 10 000 person-years). The PsA incidence rates increased with increasing BMI. Compared with psoriasis patients with BMI <25 kg/m², the RRs for developing PsA were 1.09 (0.93–1.28) for BMIs from 25.0 to 29.9, 1.22 (1.02–1.47) for BMIs from 30.0 to 34.9 and 1.48 (1.20–1.81) for BMIs ≥35.0. In our secondary analysis among all individuals, regardless of psoriasis (~2 million), the corresponding multivariate RRs tended to be stronger (1.0, 1.17, 1.57, 1.96; p for trend <0.001).

Conclusions This general population study suggests that obesity is associated with an increased risk of incident PsA and supports the importance of weight reduction among psoriasis patients who often suffer from the metabolic syndrome and obesity.

PsA is common in patients with psoriasis, affecting about 6–10% of psoriasis patients overall and substantially more patients with more extensive skin disease (up to 40%).1–3 In most patients with PsA, the symptoms do not develop until years after the onset of cutaneous psoriasis. As a result, patients with psoriasis represent a unique opportunity to study individuals at very high risk of developing a chronic inflammatory arthropathy (ie, PsA).1 To determine which patients with psoriasis are at the greatest risk of developing PsA, it is essential that risk factors be identified using robust epidemiological approaches.1

The adipose tissue in obese individuals is a site of overproduction of important inflammatory cytokines such as tumour necrosis factor α, interleukin 1, IL-6 and IL-8,9 10 and the resulting abnormal inflammatory milieu may contribute to the pathophysiology of psoriatic conditions.10 Obesity has been found to be associated with an increased risk of psoriasis,11 but its link to the risk of PsA remains less clear. Recent cross-sectional studies have found that PsA patients have a higher body mass index (BMI) than healthy controls12,13 and patients with PsA have a 61% higher odds of having obesity than psoriasis patients.13 A case series study (n=197 for PsA) reported that recalled BMI at 18 years of age was associated with an increased risk of PsA, whereas current BMI was not.14

In this study, we examined the relation between BMI and the risk of incident PsA among 75 395 individuals with psoriasis from the general population.

METHODS

Study population

The Health Improvement Network (THIN) is a computerised medical record database entered by general practitioners (GP) in the UK. Data on approximately 7.3 million patients are systematically recorded and sent anonymously to THIN. Because the National Health Service in the UK requires every individual to be registered with a GP regardless of health status, THIN is a population-based cohort representative of the UK general population. The computerised information includes demographics, details from GPs’ visits, diagnoses from specialists’ referrals and hospital admissions, results of laboratory tests, and additional health information recorded systematically including height, weight, blood pressure, smoking and vaccinations. The Read classification is used to code specific diagnoses, and a drug dictionary based on data
from the Multlex classification is used to code drugs. Health information is recorded on site at each practice using a computerised system with quality control procedures to maintain high data completion rates and accuracy. Thus, THIN data represent routine medical practice in a population-based setting.

Our primary study population consisted of psoriasis patients aged 20–89 years and enrolled in THIN at any time between January 1995 and May 2010 (n=75 955). Our secondary study populations included the cohort of individuals with or without psoriasis at baseline (N=2 026 139) and that limited to patients with incident psoriasis (n=27 980). Study cohort members had to be free of PsA before entering the study and were required to have two or more years of enrolment with the GP to help ensure exclusion of prevalent PsA. An individual’s start (index) date corresponded to the date of the first recorded BMI after all these eligibility criteria were met. Members of this cohort were followed until the date of one of the following endpoints: PsA diagnosis, 90th birthday, death or end of study period, whichever came first.

**Ascertainment of psoriasis and PsA**

Validity of THIN Read codes for psoriasis has been documented based on a sample of 4900 psoriasis patients. With a 95% response rate in a GP survey for clinical confirmation of psoriasis, the Read codes for psoriasis in THIN showed a positive predictive value (PPV) of 90% of clinical confirmation. Therefore, we defined psoriasis based on its Read code in THIN.

Although validity of PsA Read codes has not been assessed in THIN, the positive predictive value (PPV) of a single code for PsA has been reported to be over 90% in another electronic medical record database. The use of diagnostic codes plus disease-modifying antirheumatic drug data definitions only slightly and significantly improved the specificity for each diagnosis, but at the cost of a dramatic reduction in sensitivity. Therefore, we defined PsA based on its Read diagnostic code in this study.

**Assessment of exposure and covariates**

Demographics, lifestyle factors and history of trauma were collected from the database before the index date. THIN lifestyle exposure variables have been collected prospectively and successfully used in previous analyses by confirming anticipated associations such as those among alcohol, BMI and gout; smoking and lung cancer; smoking, obesity and myocardial infarction; and alcohol and paroxysmal atrial fibrillation. Our exposure of interest was BMI, defined as the patient’s body mass in kilograms divided by the square of his or her height in metres. We analysed the impact of the first recorded BMI after all these eligibility criteria were met (eg, the first BMI recorded after psoriasis diagnosis in our primary study cohort). We repeated our analysis in the THIN population by additionally including those without psoriasis status (n=1 950 744). This allowed us to examine the potential impact of psoriasis duration on varying risks of PsA. For all RRs, we calculated 95% CIs. All p values are two-sided. All statistical analyses were performed using SAS software, V9.2 (SAS Institute, Cary, North Carolina, USA).

**RESULTS**

Our primary analysis included 75 395 psoriasis patients for a combined follow-up time of 368 302 person-years, during which 976 cases of incident PsA were diagnosed. The incidence rate of PsA in this population was 26.5 per 10 000 person-years.

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**Statistical analysis**

We calculated the eligible person-time of follow-up and incidence rates according to four BMI categories: <25, 25–29.9, 30–34.9 and ≥35 kg/m², corresponding to the WHO’s classification of normal weight, overweight, obese class I and obese class II or III, respectively. We used a Cox proportional hazards model to compute the multivariate RR for incident PsA adjusting for sex, age, smoking status (never/past/current), alcohol use (six categories) and history of trauma.

We repeated our analysis in the THIN population by additionally including those without psoriasis status (n=1 950 744). This allowed us to examine the potential impact of psoriasis duration on varying risks of PsA. For all RRs, we calculated 95% CIs. All p values are two-sided. All statistical analyses were performed using SAS software, V9.2 (SAS Institute, Cary, North Carolina, USA).

**RESULTS**

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Table 1 Baseline characteristics according to body mass index for 75 395 patients with psoriasis

<table>
<thead>
<tr>
<th>Body mass index (kg/m²)</th>
<th>All</th>
<th>&lt;25.0</th>
<th>25.0–29.9</th>
<th>30.0–34.9</th>
<th>≥35.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>75 395</td>
<td>26 263</td>
<td>27 147</td>
<td>14 088</td>
<td>7897</td>
</tr>
<tr>
<td>Age (mean, years)</td>
<td>52.2</td>
<td>50.0</td>
<td>54.2</td>
<td>53.5</td>
<td>50.1</td>
</tr>
<tr>
<td>Male</td>
<td>43.4</td>
<td>35.0</td>
<td>51.6</td>
<td>48.3</td>
<td>34.8</td>
</tr>
<tr>
<td>Smoking status</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>43.4</td>
<td>42.5</td>
<td>43.8</td>
<td>43.1</td>
<td>45.7</td>
</tr>
<tr>
<td>Previous</td>
<td>26.0</td>
<td>20.3</td>
<td>28.2</td>
<td>31.1</td>
<td>28.6</td>
</tr>
<tr>
<td>Current</td>
<td>28.8</td>
<td>35.6</td>
<td>26.4</td>
<td>23.7</td>
<td>23.4</td>
</tr>
<tr>
<td>Missing</td>
<td>1.79</td>
<td>1.66</td>
<td>1.64</td>
<td>2.04</td>
<td>2.27</td>
</tr>
<tr>
<td>Alcohol intake, no. of servings per week (mean)</td>
<td>7.85</td>
<td>7.60</td>
<td>8.55</td>
<td>7.88</td>
<td>6.08</td>
</tr>
</tbody>
</table>

Values are percentages unless otherwise noted.
Our primary objective was to evaluate the association between BMI and the risk of PsA among patients with psoriasis from the general population. We found that increasing BMI among patients with psoriasis was associated with an increasing risk of incident PsA, demonstrating a dose-response relation. When we examined the risk in all eligible individuals regardless of psoriasis status, the magnitude of association became larger. These associations were independent of other potential confounders such as age, sex, trauma, smoking and alcohol intake. These data provide the first general population-based evidence that obesity is associated with an increased risk of developing PsA among psoriasis patients as well as in the general population.

A recent cross-sectional study reported that PsA patients have a higher BMI than psoriasis patients. Another study based on psoriasis patients recruited consecutively from the dermatology clinic of the University of Utah reported an independent association with recalled BMI at age 18, but no such associations with current BMI among psoriasis patients. However, as the current BMI data in that study were measured after the study recruitment when PsA status was determined, they reflect adiposity levels after the onset of PsA (as opposed to BMI before the onset of PsA). Furthermore, because the recalled BMI data at age 18 do not take into account BMI changes that occurred from age 18 leading up to and following the development and diagnosis of psoriasis, its utility for predicting the future risk of PsA based on adiposity status of psoriasis patients is unclear. Our results filled this knowledge gap by examining the relation between BMI after psoriasis diagnosis (and before PsA onset) and the risk of PsA.

Potential explanations for this increased risk for PsA among obese psoriasis patients include the inflammatory load associated with adiposity, contributing further to that associated with their underlying psoriatic condition. Obesity is associated with an overproduction of inflammatory cytokines including IL-1, IL-6, IL-8 and tumour necrosis factor α, and there is a correlation between the level of adiposity and the measured peripheral levels of associated inflammatory cytokines. Randomised trial data showing beneficial additive effects of weight loss and case reports of psoriasis clearing with reducing obesity suggest that this inflammatory milieu provided by the overabundance of adipose tissue may be important in the pathogenesis of psoriatic conditions. An alternative speculation for the increased risk observed here could be that mechanical wear, secondary to obesity, could trigger PsA in load-bearing joints, with subsequent spread of the inflammatory arthritis to other joints.

Our findings may have practical implications for the management of psoriasis patients, as PsA is common in these patients, affecting about 6–10% of psoriasis patients overall and substantially more patients with extensive skin disease (up to 40%). Thus, a modifiable risk factor, such as obesity, requires a relatively small number needed to treat among psoriasis patients compared with the general population context with a lower background risk of developing PsA. Although a randomised trial would be needed to definitively establish the usefulness of weight reduction in reducing the risk of PsA, this type of study may be difficult to perform, given the large number of psoriasis patients and trial durations necessary for sufficient cases of PsA to occur, the high costs and anticipated ethical issues. Meanwhile, as weight reduction in obese individuals has multiple health benefits, including reduced cardiovascular and mortality risks, these benefits appear all the more relevant for obese psoriasis patients to help prevent...
their excess risk of PsA along with co-morbidity and mortality complications.

The strengths and limitations of our study deserve comment. This study was performed using a large UK general practice database; therefore, findings are likely to be applicable to the general population. While the designs of previous studies on the topic made it difficult to establish the temporal relations between BMI and the risk of PsA, our study design, including the time-point of exposure measurement and the incident case outcome analysis, enabled us to establish temporal relations between these factors. Uncertainty surrounding diagnostic accuracy is a potential concern in studies that identify cases from administrative databases; therefore, findings are likely to be applicable to the general population. While the designs of previous studies on the topic made it difficult to establish the temporal relations between BMI and the risk of PsA, our study design, including the time-point of exposure measurement and the incident case outcome analysis, enabled us to establish temporal relations between these factors.

Table 3  RR for incident psoriatic arthritis by body mass index in the general population

<table>
<thead>
<tr>
<th>Body mass index (kg/m²)</th>
<th>&lt;25.0</th>
<th>25.0–29.9</th>
<th>30.0–34.9</th>
<th>≥35.0</th>
<th>p For trend</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cases of psoriatic arthritis, N</td>
<td>551</td>
<td>591</td>
<td>363</td>
<td>231</td>
<td></td>
</tr>
<tr>
<td>Incidence rate/10 000 person-years</td>
<td>1.23 (1.13–1.34)</td>
<td>1.47 (1.35–1.59)</td>
<td>2.05 (1.84–2.27)</td>
<td>2.67 (2.34–3.04)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Age- and sex-adjusted RR (95% CI)*</td>
<td>1.00 (reference)</td>
<td>1.17 (1.04–1.32)</td>
<td>1.60 (1.40–1.83)</td>
<td>2.01 (1.72–2.34)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Multivariate RR (95% CI)*</td>
<td>1.00 (reference)</td>
<td>1.17 (1.04–1.31)</td>
<td>1.57 (1.38–1.80)</td>
<td>1.96 (1.68–2.29)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*The multivariate adjusted model includes age, sex, smoking status, alcohol consumption and history of trauma.

In conclusion, this general population study suggests that obesity is associated with an increased risk of incident PsA and supports the importance of weight reduction among psoriasis patients who often suffer from the metabolic syndrome and obesity.

Contributors All authors participated in the conception, design and analyses of the study. TJL and HKC drafted the manuscript and are guarantors. All authors contributed to interpretation of the results.

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Competing interests None.

Ethical approval The current study was approved by the NHS Research Ethics Committee (09/H0305/75).

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