

EXTENDED REPORT

Comorbidities in patients with gout prior to and following diagnosis: case-control study

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ABSTRACT

Objectives To determine the burden of comorbidities in patients with gout at diagnosis and the risk of developing new comorbidities post diagnosis.

Methods There were 39 111 patients with incident gout and 39 111 matched controls identified from the UK Clinical Practice Research Data-link. The risk of comorbidity before (ORs) and after the diagnosis of gout (HRs) were estimated, adjusted for age, sex, diagnosis year, body mass index, smoking and alcohol consumption.

Results Gout was associated with adjusted ORs (95%) Cls) of 1.39 (1.34 to 1.45), 1.89 (1.76 to 2.03) and 2.51 (2.19 to 2.86) for the Charlson index of 1–2, 3–4 and \geq 5, respectively. Cardiovascular and genitourinary diseases, in addition to hyperlipidaemia, hypothyroidism, anaemia, psoriasis, chronic pulmonary diseases, osteoarthritis and depression, were associated with a higher risk for gout. Gout was also associated with an adjusted HR (95% CI) of 1.41 (1.34 to 1.48) for having a Charlson index \geq 1. Median time to first comorbidity was 43 months in cases and 111 months in controls. Risks for incident comorbidity were higher in cardiovascular, genitourinary, metabolic/ endocrine and musculoskeletal diseases, in addition to liver diseases, hemiplegia, depression, anaemia and psoriasis in patients with gout. After additionally adjusting for all comorbidities at diagnosis, gout was associated with a HR (95% CI) for all-cause mortality of 1.13 (1.08 to 1.18; p<0.001).

Conclusions The majority of patients with gout have worse pre-existing health status at diagnosis and the risk of incident comorbidity continues to rise following diagnosis. The range of associated comorbidities is broader than previously recognised and merits further evaluation.

INTRODUCTION

Gout is the most common inflammatory arthritis worldwide with a recent estimated prevalence of 2.49% in the UK.1 Initial presentation is usually with acute arthritis but chronic joint damage, subcutaneous tophi and periarticular inflammation are other common manifestations.² Since gout results from chronic hyperuricaemia and the deposition of urate crystals, urate-lowering treatment (ULT) capable of preventing crystal formation and dissolving existing crystal deposits may potentially 'cure' gout. Gout is also associated with comorbidities that may impair well-being and reduce longevity. 4-6 It is generally accepted that gout is highly associated with the features of metabolic syndrome, nephrolithiasis and chronic renal impairment. 10 However, suggested associations with other cardiovascular disease 11-14 is controversial, 15 and there is only limited evidence

for possible associations with cancer, 16 17 hypothyroidism¹⁸ and anaemia.²⁰

Current clinical guidelines and recommendations endorsed by the European League against Rheumatism, ²¹ American College of Rheumatology²² and British Society for Rheumatology/British Health Professionals in Rheumatology²³ agree on the importance of comorbidity on gout management but only discuss diseases that are recognised to be pathophysiologically related to gout. In addition, ULT is generally recommended only for patients with high urate load or with comorbidities such as poor kidney function. Although a focus on comorbidities that are causally related to or are seen as complications of gout is understandable, many patients with gout are 'complicated' by multiple comorbidities that may or may not be related directly to gout and management decisions in such cases are often difficult since recommendations and guidelines give no specific guidance in these scenarios.

Although comorbidity is considered to be epidemic in gout²⁴⁻²⁶ previous studies have only included limited categories of comorbidities. In addition, the timing of comorbidity occurrence relative to gout diagnosis has not been addressed. Therefore, using data representative of the general population of UK from the Clinical Practice Research Data-link (CPRD), this study aimed to examine the burden of comorbidity at the time of first diagnosis of gout compared with matched controls. We further followed patients with incident gout and their matched controls after diagnosis to compare their subsequent accumulation comorbidities and all-cause mortality.

METHODS

This was a case-control study using retrospective data (prior to diagnosis) and subsequent follow-up data (after diagnosis) within the CPRD. We hypothesised that gout has a higher burden of comorbidity at the time of first diagnosis and that the risk of comorbidity would continue to rise after the diagnosis of the disease. The study was approved by the Independent Scientific Advisory Committee for Medicines and Healthcare Products Regulatory Agency (MHRA) Database Research (protocol 11-021R).

Data source



The CPRD is a database containing prospectively collected medical records of research standard from around 12 million individuals in the UK. The database has been described previously²⁷ and has been

validated for many diagnoses, including gout and comorbid conditions under scrutiny in this study. ^{28–32} The database contains comprehensive information on patient demographics, lifestyle factors, medical diagnoses, results of investigations and examinations and prescribed medications. The CPRD is also linked to additional data sources including secondary care, Office for National Statistics cause of death data and information from specific disease registries.

Definition of incident gout

We used Read codes (a standard clinical terminology system used in general practice in the UK originally developed by Dr James Read) to identify patients with incident gout from the CPRD between 1997 and 2005. To be eligible as a patient with incident gout, participants had to have no evidence of gout and no prescription for ULT prior to the time of diagnosis (index date) and to have at least a 3-year continuous registration prior to the index date. The case definition was based on physician-diagnosis using 18 Read codes indicative of incident gout. The diagnosis of gout in the CPRD has been validated previously by a review of medical records of 10 patients with confirmed and 28 patients with probable gout showing that 10 out of 10 confirmed cases and 24 out of 28 probable cases were true gout (overall ascertainment rate 90%). The diagnosis of the confirmed cases are true gout (overall ascertainment rate 90%).

Selection of controls

From CPRD participants who had an active registration from 1997 to 2005, we identified at random one control per case who had no record of gout and no prescriptions of ULT in the CPRD. Control patients were frequency-matched in a 1:1 ratio to patients with incident gout by year of birth (±2 years), gender, general practice and year of first continuous registration (±2 years) at this CPRD practice. The same index date was assigned to each of the matched controls. As with patients with incident gout the matched controls had at least 3 years of registration prior to the index date.

Study period

We defined three periods to assess comorbidity: (A) the 10 years prior to diagnosis; (B) the 1 year prior to diagnosis; and (C) from diagnosis to the earliest date of occurrence of comorbidity, death, transfer out or end of study (31 December 2013), whichever came first. Cumulative probability of comorbidity was estimated at the time of diagnosis (index date) as well as 1 year, 2 years, 5 years and 10 years following diagnosis.

Comorbidity of interest and mortality

We defined gout as the index disease in this study and used Feinstein's definition for comorbidity, specifically: "Any distinct additional entity that has existed or may occur during the clinical course of a patient who has the index disease under study."34 General health status was measured by the Charlson comorbidity index. The index summarises 17 diagnostic categories (myocardial infarction, congestive heart failure, peripheral vascular disease, cerebrovascular disease, dementia, chronic pulmonary disease, rheumatological disease, peptic ulcer disease, mild liver disease, moderate or severe liver disease, diabetes mellitus (DM), DM with chronic complications, renal diseases, any malignancy (including leukaemia and lymphoma), metastatic solid tumour and HIV infection) to represent one's health status and has been a useful predictor for mortality.³⁵ Devo et al³⁶ produced a validated version for use with the International Classification of Diseases, Ninth Revision (ICD-9) based database, which was the basis of the validated Read code

version.³⁷ We calculated the Charlson score for each study person at baseline by adding scores assigned to each specific diagnosis. In addition, we also incorporated other comorbidities potentially associated with gout, including anaemia, cardiac arrhythmias, depression, hypertension, hypothyroidism, multiple sclerosis, neurological diseases, osteoarthritis, psoriasis, psychosis, urolithiasis and valvular heart disease (please refer to supplement for the rationale in selection of these comorbidities). All comorbidities were further categorised into eight categories according to systems. The definitions of these conditions were also based on physician diagnoses recorded as Read codes. We identified patients with these comorbidities from records within each of the three study periods as stated above. The Charlson index at diagnosis was based on results from the 10-year period. We categorised Charlson index as (1) 0, 1-2, 3-4 and ≥ 5 scores and (2) dichotomised it as 0 and ≥ 1 score. Mortality was based on the recording in the main database. We undertook a validation study and found the sensitivity, specificity, positive predictive value and negative predictive value of a CPRD recorded death were 0.99, 0.99, 0.93 and 1.00, respectively, suggesting that the recording of death in the CPRD is generally consistent with death registration (details in online supplement and table S1).

Statistical analysis

The prevalence of a specific comorbidity was calculated using the number of people ever diagnosed with a given comorbidity during the past 10-year period or 1-year period before the index date as the numerator and the number of patients with incident gout or matched controls as denominators. ORs and 95% CIs were used to estimate the association between gout and each coexisting medical condition. Conditional logistic regression was used to adjust for age, gender, index year, body mass index (BMI) category, smoking status and alcohol consumption. Missing data for BMI, smoking and alcohol status were coded as 'unknown'. Kaplan-Meier plots were used to estimate the cumulative probability of each comorbidity and allcause mortality in people with incident gout and those without. For calculation of cumulative probabilities, the first percentage (at baseline), indicates the percentage of people with a diagnosis before the index date. Only people at risk for a given comorbidity (not having such comorbidity at index date) were considered to estimate HRs of a specific comorbidity. HRs with 95% CIs were calculated for a Charlson index ≥ 1 , for each comorbidity separately and all-cause mortality using a Cox proportional hazards model. The HRs were adjusted by age, gender, index year, BMI categories, smoking status and alcohol consumption. For all-cause mortality we also adjusted for all categories of comorbidities at baseline. All statistical analyses were performed using SAS statistical software, V.9.3.

RESULTS

We identified 39 111 patients with incident gout during 1997 to 2005 with a mean age of 62.2±15.1 years and 72.49% being men. The median observation periods (interquartile range; IQR) before and after index date were 15 (9–27) and 9 (5–11) years, respectively. They were 1:1 matched to 39 111 controls with the same age and sex structure and observation periods before and after index date (table 1). The odds of gout increased significantly with BMI, alcohol consumption and smoking.

Retrospective observation

At the time of index date, the proportion of people having at least one comorbidity listed in the Charlson index was

Table 1 Characteristics of patients with incident gout and controls. Values are numbers (percentages) unless stated otherwise

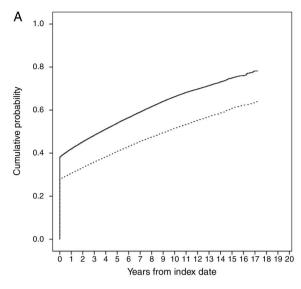
	Incident gout (n=39 111)	Controls (n=39 111)	Adjusted ORs (95% CI)	
Age (years)				
<50	8661 (22.14)	8709 (22.27)	N/A	
50-59	7815 (19.98)	7822 (20.00)	N/A	
60–69	8586 (21.95)	8608 (22.01)	N/A	
≥70	14 049 (35.92)	13 972 (35.72)	N/A	
Gender				
Men	28 350 (72.49)	28 350 (72.49)	N/A	
Women	10 761 (27.51)	10 761 (27.51)	N/A	
Median observation yea	ars (IQR)			
Prior to index date	15 (9–27)	15 (9–27)	N/A	
After index date	9 (5–11)	9 (5–11)	N/A	
BMI (kg/m²)				
<18.5	261 (0.67)	564 (1.44)	0.66 (0.56-0.77)	
18.5-24.9	7894 (20.18)	11 734 (30.00)	Reference	
25.0-29.9	15 206 (38.88)	12 815 (32.77)	1.81 (1.74–1.88)	
≥30	12 199 (31.19)	6076 (15.54)	3.15 (3.01–3.30)	
Unknown	3551 (9.08)	7922 (20.26)	0.60 (0.57-0.63)	
Smoking				
Non-smoker	10 315 (26.37)	8835 (22.59)	Reference	
Current smoker	5783 (14.79)	4233 (10.82)	1.22 (1.16–1.29)	
Ex-smoker	18 269 (46.71)	17 136 (43.81)	0.87 (0.83-0.90)	
Unknown	4744 (12.13)	8907 (22.77)	0.39 (0.37-0.41)	
Alcohol consumption (u	ınits/week)			
Never/Ex-drinker	4051 (10.36)	4338 (11.09)	Reference	
Current 1–9	17 410 (44.51)	17 123 (43.78)	1.14 (1.09–1.20)	
Current ≥10	10 285 (26.30)	5801 (14.83)	2.14 (2.02–2.27)	
Unknown	7365 (18.83)	11 849 (30.30)	0.63 (0.60-0.67)	

significantly higher in patients with incident gout (32.25%) than in controls (27.97%; p<0.001). In an adjusted logistic regression model in the 10-year period before the index date, ORs (95% CIs) were 1.39 (1.34 to 1.45), 1.89 (1.76 to 2.03) and 2.51 (2.19 to 2.86) for Charlson scores 1–2, 3–4 and ≥ 5 (p_{trend}<0.001), respectively. The analysis restricted to diagnoses recorded in the 1-year period prior to the index date gave a similar magnitude of effects (see online supplementary table S2).

All cardiovascular and genitourinary diseases were associated with a higher risk for incident gout, the highest risk being for renal diseases (OR, 5.96) and congestive heart disease (OR, 4.37) (see online supplementary table S2). In addition, hyperlipidaemia (OR 1.71), hypothyroidism (OR 1.50), anaemia (OR 1.40), psoriasis (OR 1.32), chronic pulmonary diseases (OR 1.30), osteoarthritis (OR 1.27) and depression (OR 1.09) also positively associated with incident gout in adjusted models in the 10-year and 1-year periods, while dementia (OR 0.54), multiple sclerosis (OR 0.61) and other neurological diseases (OR 0.83) were negatively associated with incident gout. Interestingly, uncomplicated DM and complicated DM were not significant risk factors for incident gout, even showing a slight negative association in logistic regression models.

Follow-up data after index date

Overall, median time to first comorbidity in the Charlson index was 43 months (95% CI 41 months to 45 months) in patients



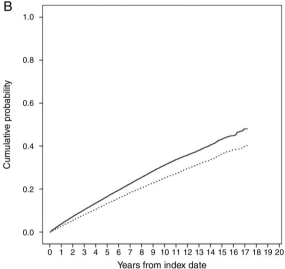


Figure 1 Cumulative probability of (A) having the first comorbidity in the categories of Charlson index (B) an increase in the score of Charlson index in patients with incident gout (solid line) and matched controls (dotted line).

with incident gout and 111 months (95% CI 108 months to 115 months) in controls (log-rank test p<0.001). The cumulative probability of having the first comorbidity in the Charlson index at 1 year, 2 years, 5 years and 10 years from index date were, respectively, 42.09%, 45.39%, 53.99% and 66.28% in patients with incident gout and 30.78%, 33.32%, 40.92% and 51.54% in controls (figure 1A). In addition, the cumulative probability of any increase in Charlson score was significantly greater in patients with gout than in controls at all time points after diagnosis (log-rank test, p<0.001; figure 1B). Table 2 shows cumulative probabilities for specific comorbidities (including existing and new ones) in patients with incident gout and matched controls at index date and 1 year, 2 years, 5 years and 10 years after index date. It is apparent that cumulative risks of most categories of comorbidity after index date were higher in patients with gout than in matched controls.

Next we investigated relative risk of incident comorbidity. Among people with a Charlson score of zero at index date, gout associated with adjusted HRs (95% CI) of 1.41 (1.34 to 1.48) for having a Charlson index ≥1. Furthermore, risk of

Table 2 Cumulative probability of comorbidity after diagnosis of gout

	Patients with incident gout Controls										
	At diagnosis	1 year	2 years	5 years	10 years	At diagnosis	1 year	2 years	5 years	10 years	p Value
Charlson index ≥1	38.25	42.09	45.39	53.99	66.27	27.97	30.78	33.32	40.91	51.54	< 0.001
Any increase in Charlson index	0	3.88	7.30	16.60	31.19	0	3.80	6.95	14.54	23.24	< 0.001
Neoplasms											
Solid malignancy, leukaemia and lymphoma	5.35	6.47	7.76	11.84	19.44	4.49	5.54	6.56	10.27	16.76	< 0.001
Metastatic solid tumours	0.14	0.30	0.51	1.05	2.08	0.16	0.33	0.49	0.97	1.88	0.266
Cardiovascular diseases											
Hypertension	34.81	38.46	41.58	49.62	59.64	18.70	21.28	23.85	30.50	39.19	< 0.00
Cardiac arrhythmias	8.80	9.93	11.00	14.01	19.10	3.72	4.30	5.02	6.95	10.64	<0.00
Cerebrovascular disease	5.99	7.02	7.95	10.46	14.48	4.13	4.92	5.57	7.65	10.74	< 0.00
Congestive heart failure	8.54	9.49	10.30	12.04	14.59	2.35	2.87	3.28	4.37	6.03	< 0.00
Myocardial infarction	5.47	6.12	6.63	8.15	10.54	2.76	3.23	3.57	4.86	6.87	< 0.00
Peripheral vascular disease	3.85	4.43	4.93	6.23	8.35	2.36	2.65	2.92	3.92	5.49	< 0.00
Valvular heart disease	2.27	2.60	2.96	3.94	5.71	0.96	1.12	1.29	1.78	2.87	< 0.00
Genitourinary diseases											
Urolithiasis	0.94	1.05	1.14	1.42	1.99	0.66	0.73	0.82	1.07	1.42	< 0.00
Renal diseases	3.05	4.20	5.30	8.50	12.51	0.54	0.76	1.07	2.22	4.04	<0.00
Metabolic and endocrine diseases											
Uncomplicated diabetes mellitus	7.08	8.37	9.66	13.87	20.87	5.83	6.46	7.11	9.21	12.50	< 0.00
Diabetes mellitus with complications	1.30	1.64	2.03	3.60	13.92	1.13	1.43	1.71	2.79	8.41	< 0.00
Hyperlipidaemia	10.11	11.75	13.23	17.10	22.05	5.37	6.30	7.28	10.04	13.94	< 0.00
Hypothyroidism	3.61	4.15	4.68	6.22	8.29	2.23	2.60	2.95	4.03	5.58	< 0.00
Gastrointestinal and hepatic diseases											
Peptic ulcer disease	2.02	2.33	2.54	3.10	3.88	1.78	1.96	2.11	2.50	3.14	< 0.00
Mild liver disease	0.39	0.52	0.62	1.11	1.99	0.29	0.35	0.40	0.60	0.94	< 0.00
Moderate to severe liver disease	0.08	0.10	0.11	0.21	0.43	0.07	0.08	0.10	0.18	0.30	0.002
Chronic pulmonary diseases	13.59	14.70	15.62	18.20	22.02	9.83	10.57	1.44	13.66	17.15	< 0.00
Musculoskeletal and connective tissue diseases											
Osteoarthritis	10.91	12.71	14.24	18.30	24.00	7.80	8.83	9.77	12.47	16.55	< 0.00
Rheumatological disease	7.12	8.46	9.72	12.94	18.78	6.12	6.77	7.37	9.22	12.31	< 0.00
Neurological and mental disorders											
Dementia	0.30	0.46	0.65	1.29	2.97	0.56	0.79	1.02	1.79	3.76	< 0.00
Hemiplegia	0.39	0.48	0.55	0.6	1.03	0.37	0.44	0.46	0.64	0.80	0.00
Multiple sclerosis	0.08	0.09	0.09	0.11	0.13	0.15	0.16	0.17	0.19	0.23	0.00
Other neurological diseases	1.77	2.17	2.53	3.62	5.30	2.17	2.51	2.84	3.84	5.65	0.07
Psychosis	0.40	0.49	0.55	0.75	1.16	0.24	0.61	0.68	0.93	1.27	0.23
Depression	12.10	13.60	14.97	18.90	24.74	10.39	11.45	12.59	15.76	20.59	< 0.00
Other comorbidities											
Anaemia	8.84	1.45	11.75	16.11	23.43	5.86	6.68	7.55	10.43	15.55	< 0.00
Psoriasis	3.20	3.53	3.81	4.63	5.81	2.24	2.40	2.61	3.12	3.85	<0.00
HIV infection	0.03	0.03	0.04	0.04	_	<0.001		_	0.03	_	0.02

developing incident comorbidity in the categories of cardiovascular, genitourinary, metabolic/endocrine and musculoskeletal/connective tissue diseases was higher in patients with gout (table 3). For example, patients with gout were three times more likely to develop renal diseases (HR 3.18, 95% CI 2.88 to 3.50). In addition, risks of liver diseases, hemiplegia, depression, anaemia and psoriasis were also higher in patients with gout. In contrast, patients with gout were less likely (HR 0.82, 95% CI 0.73 to 0.92) to have dementia.

After index date, mortality at 5 years and 10 years (95% CIs) from any cause was 14.43% and 26.98% in patients with incident gout, compared with 11.14% and 21.66% in matched controls (log-rank test, p<0.001; see online supplementary table S3). In men and women estimates of the effect of a gout diagnosis on mortality were attenuated after adjustment for demographic and lifestyle variables (model 2), and attenuated even

further after adjustment for comorbidity status at index date (model 3) (see online supplementary table S4). After the adjustment for age, sex, index year, BMI class, smoking status, alcohol consumption and all individual comorbidities at diagnosis (model 3), gout associated with a HR (95% CI) for all-cause mortality of 1.13 (1.08 to 1.18; p<0.001).

DISCUSSION

This study used a large primary care database in the UK to compare the burden of existing comorbidity at diagnosis in patients with gout with matched controls and to estimate the risks of developing new comorbidities following diagnosis. Overall, a third of patients with newly diagnosed gout already had a Charlson score ≥ 1 , compared with a quarter of matched controls. The prevalence of hypertension, cardiac arrhythmias, valvular heart disease, urolithiasis, hyperlipidaemia,

 Table 3
 Risk of incident comorbidity after diagnosis of gout

	At-risk cases	At-risk controls	Unadjusted HRs	Adjusted HRs†
All-cause mortality	39 111 (25.40)	39 111 (20.20)	1.34 (1.29–1.39)*	1.36 (1.31–1.42)
Charlson index ≥1	24 152 (41.31)	28 171 (29.40)	1.62 (1.55-1.70)*	1.41 (1.34–1.48)
Any increase in Charlson index	39 111 (25.51)	39 111 (21.18)	1.31 (1.26-1.36)*	1.20 (1.16–1.26)
Neoplasms				
Solid malignancy, leukaemia and lymphoma	37 040 (12.52)	37 375 (11.04)	1.17 (1.11–1.23)*	1.10 (1.04–1.16)
Metastatic solid tumours	39 057 (1.57)	39 047 (1.47)	1.06 (0.93-1.21)	1.04 (0.91-1.20)
Cardiovascular diseases				
Hypertension	25 496 (31.47)	31 798 (21.21)	1.83 (1.74-1.92)*	1.51 (1.43-1.58)
Cardiac arrhythmias	35 669 (9.60)	37 656 (6.18)	1.74 (1.62-1.86)*	1.59 (1.48–1.70)
Cerebrovascular disease	36 767 (7.74)	37 494 (5.99)	1.38 (1.29-1.47)*	1.29 (1.62–1.86)
Congestive heart failure	35 772 (5.78)	38 192 (3.25)	2.01 (1.84-2.18)*	1.81 (1.65–1.98)
Myocardial infarction	36 973 (4.39)	38 032 (3.52)	1.25 (1.14–1.37)*	1.16 (1.05–1.28)
Peripheral vascular disease	37 605 (3.90)	38 189 (2.62)	1.58 (1.44–1.74)*	1.56 (1.40–1.72)
Valvular heart disease	38 223 (2.93)	38 735 (1.64)	1.96 (1.76–2.19)*	1.80 (1.60–2.04)
Genitourinary diseases	, ,	` ,	, ,	
Urolithiasis	38 743 (0.84)	38 855 (0.63)	1.39 (1.16–1.67)*	1.26 (1.02–1.55)
Renal diseases	37 919 (8.00)	38 899 (2.81)	3.43 (3.13–3.76)*	3.18 (2.88–3.50)
Metabolic and endocrine diseases	, ,	` ,	, ,	` '
Uncomplicated diabetes mellitus	36 342 (12.07)	36 832 (5.94)	2.15 (2.02-2.29)*	1.65 (1.54–1.77)
Diabetes mellitus with complications	38 604 (10.82)	38 669 (6.29)	1.80 (1.70–1.92)*	1.35 (1.26–1.45)
Hyperlipidaemia	35 156 (10.69)	37 011 (7.30)	1.64 (1.54–1.75)*	1.40 (1.31–1.50)
Hypothyroidism	37 701 (4.00)	38 237 (2.81)	1.53 (1.40–1.68)*	1.46 (1.32–1.61)
Gastrointestinal and hepatic diseases	,	, ,	,	,
Peptic ulcer disease	39 079 (0.29)	39 084 (0.19)	1.53 (1.09–2.16)*	1.37 (0.95–2.00)
Mild liver disease	38 321 (1.64)	38 416 (1.12)	1.52 (1.32–1.75)*	1.47 (1.26–1.72)
Moderate to severe liver disease	38 960 (1.26)	38 997 (0.53)	2.39 (2.00–2.85)*	1.97 (1.61–2.41)
Chronic pulmonary diseases	33 795 (8.11)	35 267 (6.81)	1.20 (1.12–1.28)*	1.10 (1.02–1.18)
Musculoskeletal and connective tissue diseases	,	,	,	,
Osteoarthritis	34 843 (12.10)	36 062 (7.84)	1.71 (1.61–1.82)*	1.45 (1.35–1.54)
Rheumatological disease	36 325 (10.43)	36 719 (5.54)	1.96 (1.84–2.10)*	1.57 (1.46–1.69)
Neurological and mental disorders	((,	(
Dementia	38 994 (2.22)	38 892 (2.63)	0.83 (0.75-0.93)*	0.82 (0.73-0.92)
Hemiplegia	38 959 (0.51)	38 968 (0.36)	1.50 (1.19–1.89)*	1.48 (1.16–1.90)
Multiple sclerosis	39 081 (0.05)	39 053 (0.06)	0.70 (0.37–1.30)	0.70 (0.22–2.18)
Other neurological diseases	38 419 (3.01)	38 261 (2.94)	1.08 (0.98–1.19)	1.06 (0.96–1.17)
Psychosis	38 953 (0.63)	38 901 (0.60)	1.00 (0.81–1.23)	1.04 (0.82–1.32)
Depression	34 379 (11.93)	35 047 (9.42)	1.28 (1.21–1.35)*	1.19 (1.12–1.26)
Other comorbidities	3.3.3 (11.33)	55 0 (5. 1 <u>L</u>)	(1.55)	5 (11.12 11.20)
Anaemia	35 727 (13.54)	36 854 (8.63)	1.74 (1.65–1.84)*	1.58 (1.49–1.68)
Psoriasis	37 858 (2.17)	38 233 (1.35)	1.62 (1.44–1.82)*	1.53 (1.34–1.74)
HIV infection	39 099 (0.02)	39 108 (0.02)	2.00 (0.60–6.64)	1.84 (0.62–5.44)

Figures are number of at-risk population (percentage of at-risk population who developed incident comorbidity) and HRs (95% CIs).

BMI, body mass inde

hypothyroidism, chronic pulmonary disease, osteoarthritis, depression, anaemia and psoriasis were all significantly higher in patients with incident gout than controls. Furthermore, approximately half of patients with gout had a Charlson index ≥1 within 5 years from initial diagnosis, whereas it took 10 years for half the controls to develop a Charlson index ≥ 1 . Patients with incident gout without a prior history of these comorbidities were more likely than controls to develop cancer, cardiovascular diseases, urolithiasis, renal diseases, hyperlipidaemia, hypothyroidism, peptic ulcer disease, mild liver disease, osteoarthritis, rheumatological disease, depression, anaemia and psoriasis than controls. Despite other diseases which affect longevity, gout itself carried an additional 13% of deaths in the affected patients. Overall, the burden of

comorbidity is very high at diagnosis of gout and the risk of developing new comorbidity is also higher in patients with incident gout than in the general population.

The associations between gout and several categories of comorbidities are well known. For instance, the association between gout and the components of metabolic syndrome is consistently reported^{38–40} and is explained by known inter-relationship between these factors, in part mediated through effects on urate production, renal angiotensin secretion and renal urate excretion.⁴¹ ⁴² However, the association between gout and type 2 DM is more complex.^{43–45} Gout is a risk factor for DM but the risk of DM was not increased in patients with gout at diagnosis. Gout is a known risk factor for cardiovascular diseases such as cerebrovascular disease,⁴⁶

^{*}p<0.05.†Adjusted for age, gender, index year, BMI class, smoking status and alcohol consumption.

congestive heart failure⁴⁷ and myocardial infarction,⁶ 12-14 48 collectively leading to an increased risk for all-cause and cardio-vascular mortality in patients with gout.¹⁵ ⁴⁹ This association may be mediated via endothelial dysfunction caused by high uric acid and low grade crystal-induced inflammation.⁵⁰ Links between hyperuricaemia and kidney disease are well-established^{51–53} although convincing data for an aetiological link between gout and chronic kidney disease are sparse.¹⁰ Although a higher risk of urolithiasis was reported half a century ago^{54–55} solid evidence for this emerged only recently⁸ and studies using CT identified renal calculi in a third of patients with gout⁵⁶ suggesting underestimation of the magnitude of this risk in studies based solely on a history of symptomatic calculi.

The associations between gout and some or the other comorbidities identified in this study have received less attention. An association between gout and hypothyroidism has been suggested previously in older case series and studies of patients not representative of the general population. ¹⁸ ¹⁹ ⁵⁷ A possible explanation for this association is the ability of thyroid hormone to influence serum urate levels through regulation of glomerular filtration rate.⁵⁸ There is no clear explanation for the association between gout and anaemia, 20 although hyperuricaemia could result from the increased synthesis of nucleic acids resulting from an erythropoietic response⁵⁹ and conversely greater consumption of non-steroidal anti-inflammatory drugs (NSAID) for joint pain could cause anaemia through gastrointestinal bleeding. Co-localisation of gout and osteoarthritis has been reported previously⁶⁰ and possible explanations for this include reduced viability of chrondrocytes⁶¹ and mechanical and inflammatory effects of urate crystals on cartilage, and the heightened propensity of damaged cartilage to facilitate monosodium urate crystal nucleation. 62 Gout is also associated with cancer 16 17 and psoriasis, 63 both diseases being associated with increased cell turnover that can cause hyperuricaemia. An association between gout and liver disease is difficult to explain, although a relationship between gout and non-alcoholic fatty liver disease has been suggested previously,⁶⁴ both of which are related to insulin resistance.

In contrast to the positive associations above, neurodegenerative conditions such as dementia and multiple sclerosis seem to have an inverse relationship with incident gout. The association is more apparent prior to presentation with gout than during follow-up analysis. Uric acid has antioxidant effects that particularly may afford a beneficial effect on central neural tissue degeneration⁶⁵ and ageing,⁶⁶ though any such benefit is probably largely outweighed in other tissues due to the deleterious effects of uric acid on vascular endothelium.⁶⁷ Only a few studies have reported that lower urate associates with an elevated risk of multiple sclerosis⁶⁸ but our study supports this. Such negative associations with hyperuricaemia and gout are important in informing the debate concerning the optimum level of urate that we should target in the long term to prevent further crystal formation while avoiding neurological problems from taking the level too low.⁶⁹

The other associations identified in this study between gout and other comorbidities are harder to explain. Previous case reports have suggested that polycythaemia secondary to chronic lung disease might contribute to gout. In addition, tophi have been found in lung tissue of a patient with interstitial inflammation and pleural effusion. Patients with HIV infection seem to have a higher risk of gout and this potentially could result from use of antiretroviral agents such as ritonavir which may cause lipodystrophy, metabolic disturbance and hyperuricaemia.

The existence of comorbidity greatly complicates the management of gout and requires special attention in managing patients with gout. Currently, European, 21 British 23 and American guidelines²² have considered comorbidity as part of initial assessment of patients with gout and the recommended management of chronic hyperuricaemia also partly depends on the existence of comorbidities. However, as our data show, comorbidities with higher absolute and relative risks were not limited to the recommended 'checklist'. Hypothyroidism and anaemia are two examples. Yet screening for thyroid function is not mentioned in any of the guidelines and only the American College of Rheumatology guideline recommends a complete blood cell count for patients with gout, 22 which is mainly to look for myeloproliferative disease rather than anaemia. Therefore, this study supports the case for a comprehensive investigation for comorbidities, including but not limited to 'checklists', as an integral part of initial assessment for patients with gout at diagnosis. Furthermore, given that half of patients with gout are expected to have one or more comorbidities within 5 years from diagnosis, close observation by annual assessment for the occurrence of incident comorbidity may be warranted.

There are several limitations to this study. First, there may be some misclassification bias since the identification of patients with gout was based on physician diagnosis, rather than according to classification criteria⁷³ ⁷⁴ or to the 'gold standard' of urate crystal identification. However, the validity of gout diagnosis in the CPRD has been investigated and found to be high.³² Similarly, there may have been some misclassification of comorbidities though again, most of the comorbidities in this study have been validated previously.²⁸ In addition, there is no reason to suspect a differential misclassification between patients with gout and those without. Second, differential ascertainment bias between patients with incident gout and controls cannot be excluded entirely, even though the estimates based on different lengths of previous observations produced similar results. Third, the models did not adjust for all potential risk factors for gouty arthritis, such as diet and physical activity, both of which are not routinely recorded in the CPRD. Fourth, the problem associated with multiple testing may arise in our study. However, this study aimed to provide estimates of disease burden of patients with gout at diagnosis and subsequent to diagnosis, rather than to define causes of gout and the main focus of this analysis was to identify exposures and outcomes that are more closely associated with gout. Also, most of the results were interpreted in the light of existing research so the conclusions are unlikely to have been affected unduly by false-positive findings. In addition, the sample size of this study is about 62 000 people. Even when a Bonferroni correction is applied, which is very conservative when the multiple outcome measures under consideration are not independent, the majority of previously significant results remained.

In conclusion, the majority of patients with gout in the UK already have comorbidity at diagnosis and the risk of incident comorbidity continues to rise after diagnosis. All-cause mortality was higher in patients with gout and the increased risk was contributed to by gout itself and by comorbidities to approximately the same degree. This study suggests that a thorough search for a broad range of comorbidity and subsequent vigilant observation should be considered for all patients with gout from the date of first diagnosis.

Contributors C-FK had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. C-FK, WZ and MD conceived and designed the study. C-FK and WZ obtained the

funding and acquired the data. C-FK, MJG and WZ performed and supervised the statistical analysis. C-FK, MJG, CM, WZ and MD analysed and interpreted the data. C-FK and WZ drafted the manuscript. All authors contributed to the critical revision of the manuscript for important intellectual content. WZ, MJG and MD supervised the study. MD is the guarantor.

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