

2018 updated European League Against Rheumatism evidence-based recommendations for the diagnosis of gout

Pascal Richette,^{1,2} Michael Doherty,³ Eliseo Pascual,⁴ Victoria Barskova,⁵ Fabio Becce,⁶ Johann castaneda,⁷ Malcolm Coyfish,⁸ Sylvie Guillo,⁹ Tim Jansen,¹⁰ Hein Janssens,¹¹ Frédéric Lioté,^{12,13} Christian D Mallen,¹⁴ George Nuki,¹⁵ Fernando Perez-Ruiz,¹⁶ José Pimentao,¹⁷ Leonardo Punzi,¹⁸ Anthony Pywell,⁸ Alexander K So,¹⁹ Anne-Kathrin Tausche,²⁰ Till Uhlig,²¹ Jakub Zavada,²² Weiya Zhang,²³ Florence Tubach,²⁴ Thomas Bardin²⁵

Handling editor Josef S Smolen

► Additional material is published online only. To view please visit the journal online (<http://dx.doi.org/10.1136/annrheumdis-2019-215315>).

For numbered affiliations see end of article.

Correspondence to

Professor Pascal Richette, Service de Rhumatologie, Hopital Lariboisiere Centre Viggo Petersen, Paris 75010, France; pascal.richette@aphp.fr

Received 9 March 2019

Revised 24 April 2019

Accepted 25 April 2019

ABSTRACT

Although gout is the most common inflammatory arthritis, it is still frequently misdiagnosed. New data on imaging and clinical diagnosis have become available since the first EULAR recommendations for the diagnosis of gout in 2006. This prompted a systematic review and update of the 2006 recommendations. A systematic review of the literature concerning all aspects of gout diagnosis was performed. Recommendations were formulated using a Delphi consensus approach. Eight key recommendations were generated. A search for crystals in synovial fluid or tophus aspirates is recommended in every person with suspected gout, because demonstration of monosodium urate (MSU) crystals allows a definite diagnosis of gout. There was consensus that a number of suggestive clinical features support a clinical diagnosis of gout. These are monoarticular involvement of a foot or ankle joint (especially the first metatarsophalangeal joint); previous episodes of similar acute arthritis; rapid onset of severe pain and swelling; erythema; male gender and associated cardiovascular diseases and hyperuricaemia. When crystal identification is not possible, it is recommended that any atypical presentation should be investigated by imaging, in particular with ultrasound to seek features suggestive of MSU crystal deposition (double contour sign and tophi). There was consensus that a diagnosis of gout should not be based on the presence of hyperuricaemia alone. There was also a strong recommendation that all people with gout should be systematically assessed for presence of associated comorbidities and risk factors for cardiovascular disease, as well as for risk factors for chronic hyperuricaemia. Eight updated, evidence-based, expert consensus recommendations for the diagnosis of gout are proposed.

INTRODUCTION

Gout is caused by prolonged hyperuricaemia which leads to the formation of monosodium urate (MSU) crystals that accumulate in joints and other tissues.¹ It is recognised as the most common form of inflammatory arthritis,² with a prevalence of 0.9% to 2.5% in Europe,^{3,4} 3.9% in the USA⁵ and over 6% in some Oceanic-Pacific ethnic groups.^{6,7}

The natural history of MSU deposits evolves through a number of stages: asymptomatic MSU crystal deposition during which people have MSU crystal deposition in the absence of gout; gout defined by MSU crystal deposition and clinical disease elements such as gout flare, chronic gouty arthritis and tophi. If present, recurrent gout flares are separated by asymptomatic intervals named intercritical gout.^{8,9}

Despite effective treatments, gout is still often misdiagnosed and its management remains suboptimal.^{3,10,11} This may explain why the premature mortality among patients with gout remains unimproved over the last decade.¹²

In 2006, the EULAR produced its first evidence-based recommendations for the diagnosis of gout.¹³ The 2006 task force agreed that detection of MSU crystals in synovial fluid (SF) was the gold standard for the diagnosis of gout. Since then, a number of studies have explored the diagnostic value of clinical algorithms and of imaging modalities such as ultrasound (US) or dual-energy CT (DECT). This prompted a revision of the 2006 recommendations following an updated systematic literature review (SLR) and a Delphi process to achieve consensus.

METHODS

With the approval of the EULAR executive committee, the convenor (TB) along with two coconvenors of the 2006 task force (MD and EP), an epidemiologist (FT) and an academic rheumatologist (PR) formed a steering group to update the 2006 EULAR recommendations for the diagnosis of gout.

This EULAR task force comprised 15 rheumatologists, 1 musculoskeletal radiologist, 2 general practitioners (GPs), 1 research fellow, 2 patients and 3 experts in epidemiology/methodology from 12 European countries. The recommendations were developed according to the standardised operating procedures for the elaboration, evaluation dissemination and implementation of recommendations endorsed by EULAR.¹⁴

The task force used the same methodology as that used for developing the 2016 EULAR recommendations for the management of gout.¹⁵ Briefly, the first step was to determine whether the 10



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To cite: Richette P, Doherty M, Pascual E, et al. *Ann Rheum Dis* Epub ahead of print: [please include Day Month Year]. doi:10.1136/annrheumdis-2019-215315

Recommendation

former EULAR recommendations (2006) for the diagnosis of gout should be retained, modified or abandoned. Subsequently, one research fellow (JC) with the help of an expert in systematic review methodology (SG) performed an SLR by searching for literature published since 1 January 2005 in MEDLINE, EMBASE and Cochrane Library databases (1996) up to June 2013. The quality of evidence and grades of recommendation were determined by PR according to the standards of the Oxford Centre for Evidence-Based Medicine¹⁴ (see online supplementary material 1). Next, the task force members attended a 2-day meeting during which results of the SLR were presented in an aggregated form. The task force debated and evaluated the evidence presented and formulated a preliminary set of new recommendations. Consensus for eight updated recommendations was then reached following three Delphi rounds undertaken by email after the meeting. Each participant was asked to rate their level of agreement (LoA) with each final recommendation again using a 9-point numerical rating scale (1, totally disagree; 9, fully agree) and could propose a reformulation of the recommendation. Because the delay between the SLR and agreement of final guideline was longer than expected, the SLR was updated up to July 2018 (see online supplementary material 1). The steering group discussed the results of this additional SLR and agreed that a fourth Delphi round was required to gain full support from all participants. Finally, the task force agreed that this additional SLR did not impact the overall content of the recommendations.

RESULTS

The task force voted unanimously for changes in all items of the 2006 recommendations (see online supplementary material 1). The literature search yielded 1173 records, of which 83 references were analysed (see flow chart, online supplementary material 1). All the previous recommendations were amended to reflect the updated evidence from the SLR. Four Delphi rounds by email were undertaken to establish the final set of eight recommendations (table 1).

Ranking details for level of evidence and grade of recommendation: see online supplementary material 1. The LoA

(mean±SD) for each recommendation was graded by the task force from 1 (totally disagree) to 9 (fully agree).

RECOMMENDATIONS

(1) *Search for crystals in SF or tophus aspirates is recommended in every person with suspected gout, because demonstration of MSU crystals allows a definitive diagnosis of gout.*

Clinical features of gout are due to host defence mechanisms reacting to MSU crystals. The task force agreed that the gold standard for a diagnosis of gout still relies on the demonstration of MSU crystals in fluid or tophus aspirates, as it has 100% specificity. Crystals can be detected by polarised light microscopy in SF aspirated from both symptomatic and asymptomatic joints, particularly the first metatarsophalangeal (MTP) joint and joints previously inflamed. This means that a diagnosis of gout can be established even during the asymptomatic intercritical period after, or between gout flares, so called intercritical gout^{16 17} (figure 1). This recommendation is also determined by the absence of an alternative validated diagnostic test and a good safety profile.¹⁸ The recommendation also implies that patients with suspected gout should be referred if needed to a physician with expertise in aspirating and analysing SF.¹⁹

(2) *Gout should be considered in the diagnosis of any acute arthritis in an adult. When SF analysis is not feasible, a clinical diagnosis of gout is supported by the following suggestive features: monoarticular involvement of a foot (especially the first MTP) or ankle joint; previous similar acute arthritis episodes; rapid onset of severe pain and swelling (at its worst in <24 hours); erythema; male gender and associated cardiovascular diseases and hyperuricaemia. These features are highly suggestive but not specific for gout.*

The task force recognised that identification of MSU crystals in SF can be challenging because joint aspiration and SF analysis require skills and facilities that are not always present in the primary care setting or in the emergency departments. Moreover, handling and storage of SF in certain cases can affect the reliability of analysis,^{20 21} and urate lowering therapy (ULT), by dissolving crystals, can impact the sensitivity of MSU detection.¹⁷ In the previous EULAR recommendations,¹³ the first two

Table 1 Final set of eight recommendations for the diagnosis of gout

Recommendations	Level of evidence	Grade of recommendation	Level of agreement
1 Search for crystals in synovial fluid or tophus aspirates is recommended in every person with suspected gout, because demonstration of MSU crystals allows a definitive diagnosis of gout.	2b	B	8.6±1.0
2 Gout should be considered in the diagnosis of any acute arthritis in an adult. When synovial fluid analysis is not feasible, a clinical diagnosis of gout is supported by the following suggestive features: monoarticular involvement of a foot (especially the first MTP) or ankle joint; previous similar acute arthritis episodes; rapid onset of severe pain and swelling (at its worst in <24 hours); erythema; male gender and associated cardiovascular diseases and hyperuricaemia. These features are highly suggestive but not specific for gout.	2b	B	8.6±0.8
3 It is strongly recommended that synovial fluid aspiration and examination for crystals is undertaken in any patient with undiagnosed inflammatory arthritis.	3	C	8.8±0.3
4 The diagnosis of gout should not be made on the presence of hyperuricaemia alone.	2a	B	8.9±0.2
5 When a clinical diagnosis of gout is uncertain and crystal identification is not possible, patients should be investigated by imaging to search for MSU crystal deposition and features of any alternative diagnosis.	1b	A	8.5±1.0
6 Plain radiographs are indicated to search for imaging evidence of MSU crystal deposition but have limited value for the diagnosis of gout flare. Ultrasound scanning can be more helpful in establishing a diagnosis in patients with suspected gout flare or chronic gouty arthritis by detection of tophi not evident on clinical examination, or a double contour sign at cartilage surfaces, which is highly specific for urate deposits in joints.	1b	A	8.2±0.9
7 Risk factors for chronic hyperuricaemia should be searched for in every person with gout, specifically: chronic kidney disease; overweight, medications (including diuretics, low-dose aspirin, cyclosporine, tacrolimus); consumption of excess alcohol (particularly beer and spirits), non-diet sodas, meat and shellfish.	1a	A	8.2±1.3
8 Systematic assessment for the presence of associated comorbidities in people with gout is recommended, including obesity, renal impairment, hypertension, ischaemic heart disease, heart failure, diabetes and dyslipidaemia.	1a	A	8.7±0.6

MSU, monosodium urate; MTP, metatarsophalangeal.

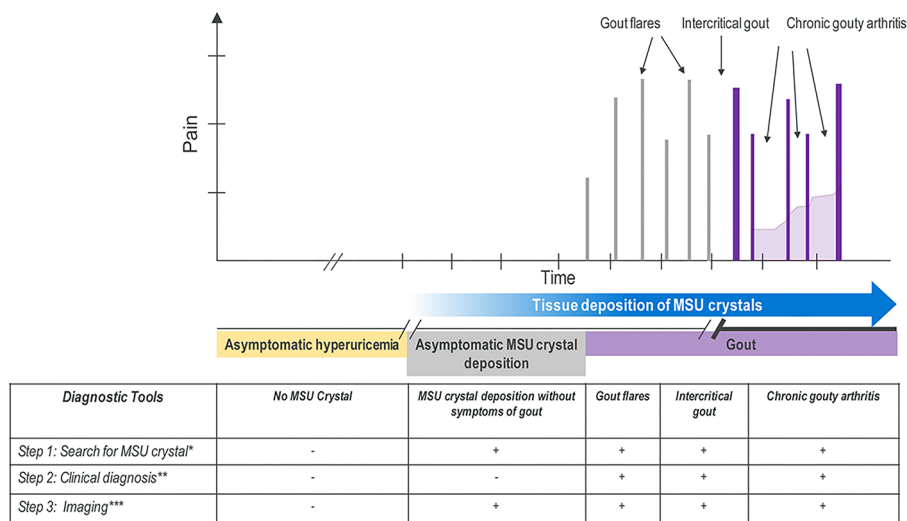


Figure 1 Recommended diagnostic modalities according to the disease states of gout. The figure shows the continuum from preclinical states (asymptomatic hyperuricaemia and then asymptomatic MSU crystal deposition) to gout (clinical states). The EULAR recommends a three-step approach for the diagnosis of gout. *The first step relies on MSU crystal identification in synovial fluid or tophus aspirates; **If not feasible, the second step relies on a clinical diagnosis (based on the presence of hyperuricaemia and associated clinical features of gout); ***The last step recommends imaging, particularly US or DECT, to search for imaging evidence of MSU crystal deposition when a clinical diagnosis of gout is uncertain and crystal identification is not possible. DECT, dual-energy CT; MSU, monosodium urate; US, ultrasound.

recommendations stated that in the absence of SF analysis, a clinical diagnosis of gout can be reasonably made for typical presentations, particularly when patients present with podagra, that is, a gout flare at the first MTP joint. Since then, the predictive values (individually and combined) of typical clinical features of gout have been determined^{22–25} and new classification criteria and algorithms based solely on clinical signs and symptoms have been produced, in patients experiencing^{26, 27} or not experiencing acute arthritis.^{28–30} Their external validity when compared with SF analysis has also been determined. Overall, both the sensitivities and specificities of these algorithms were greater than 80%

Table 2 Sensitivity and specificity of clinical algorithms published since 2006

	Sensitivity, %	Specificity, %	Demonstration of MSU crystals in cases
Mexico, 2010 ³⁰			
Vazquez-Mellado <i>et al</i> ⁶⁶	0.97	0.95	Yes
Taylor <i>et al</i> ⁶⁷			
Less than 2 years	0.87	0.66	Yes
More than 2 years	0.98	0.34	Yes
Jatuworapruk <i>et al</i>⁶⁷			
Less than 2 years	0.88	0.81	Yes
More than 2 years	0.99	0.3	Yes
Netherlands, 2010 ²⁶			
Taylor <i>et al</i> ⁶⁷			Yes
Less than 2 years	0.87	0.75	Yes
More than 2 years	0.96	0.47	Yes
Jatuworapruk <i>et al</i>⁶⁷			
Less than 2 years	0.73	0.85	Yes
More than 2 years	0.91	0.5	Yes
Paris, 2015 ²⁹			
	0.88	0.93	Yes
ACR/Eular, 2015 ²⁸			
Clinical only	0.85	0.78	Yes
Full set (with imaging)	0.92	0.89	Yes

MSU, monosodium urate.

for diagnosis as compared with the gold standard of MSU demonstration in SF (table 2). It must be emphasised, however, that most of these criteria except one²⁶ are classification criteria^{28–30} and not diagnostic criteria, precluding their use to guide the care of individual patients.³¹ Consequently, the task force included in this recommendation the most discriminating clinical and laboratory features for the diagnosis of gout, based on the results of these studies, apart from imaging, which is considered separately in the fifth and sixth recommendations. However, the task force considered that their specificity was not high enough to replace demonstration of MSU crystals in SF for the diagnosis of gout.

(3) *It is strongly recommended that SF aspiration and examination for crystals is undertaken in any patient with undiagnosed inflammatory arthritis.*

The rising prevalence of gout makes it the most frequent cause of inflammatory arthritis. Its crude prevalence ranges approximately from 1% to 4% in Europe and USA.² As atypical presentations of gout are not rare, and crystal identification allows a definite diagnosis of gout, the task force emphasised the need for SF examination in all cases of undiagnosed inflammatory arthritis.

(4) *The diagnosis of gout should not be made on the presence of hyperuricaemia alone.*

Epidemiological studies show a relationship between serum uric acid (SUA) levels and incident gout, and that not all hyperuricaemic patients have or will develop gout.^{32–35} For instance, only 22% of asymptomatic patients with SUA levels above 9 mg/dL developed incident gout over a 5-year period.³⁵ With MSU crystals identification as reference standard, the specificity of hyperuricaemia for the diagnosis of gout is low, ranging from 53% to 61%.^{23, 26} Therefore, hyperuricaemia alone should be considered solely as a strong risk factor for incident gout³⁶ and not as a surrogate marker for its diagnosis. Conversely, absence of hyperuricaemia has a markedly low negative likelihood ratio,²² indicating that after an episode of acute arthritis has settled, the absence of hyperuricaemia does not completely exclude a diagnosis of gout but makes the diagnosis very unlikely. A study found that roughly 10% of people with gout have SUA levels

Table 3 Sensitivity and specificity of DECT for the diagnosis of gout

	Sensitivity	Specificity
Manger <i>et al</i> 2012 ⁴⁹	0.78	NA
Wu <i>et al</i> 2014 ⁹⁸	0.97	0.87
HJ <i>et al</i> 2015 ⁹⁹	0.91	0.85
Huppertz <i>et al</i> 2014 ⁷⁷	0.78	0.93
Dalbeth <i>et al</i> 2015 ⁴⁰		
Early disease (≤3 years)	0.79	NA
Late disease (>3 years)	0.84	NA
Ahmad <i>et al</i> 2016 ¹⁰⁰	0.82	0.89
Kiefer <i>et al</i> 2016 ¹⁰¹	0.71	0.95
Ogdie <i>et al</i> 2015 ⁴³ MA*	0.87 (0.79–0.93)	0.84 (0.75–0.90)
Lee and Song 2017 ¹⁰² MA†	0.84 (0.81–0.87)	0.93 (0.93–0.96)

Diagnosis of gout, clinical classification criteria only^{98–101}; MSU crystal identification only^{40 50 103 104}; clinical criteria and/or MSU crystal identification.^{49 77}

*Pooled data from: ^{50 103 104}

†^{50 77 98–101 103 104}

DECT, dual-energy CT; MA, meta-analysis; MSU, monosodium urate; NA, not applicable.

below 6 mg/dL during gout flares.³⁷ Thus, the SUA levels have a limited diagnostic value, especially during a gout flare and should be preferably determined at distance from a gout flare. Of note, 15%–25% of people with asymptomatic hyperuricaemia have asymptomatic MSU crystal deposition,^{38–41} a finding which supports the concept that there is, in some people, a continuum from asymptomatic hyperuricaemia to gout^{9 42} (figure 1). Of note, the definition of hyperuricaemia varies widely across published studies, ranging from 6 to 7 mg/dL.⁹

(5) *When a clinical diagnosis of gout is uncertain and crystal identification is not possible, patients should be investigated by imaging to search for MSU crystal deposition and features of any alternative diagnosis.*

For patients with atypical clinical features and in whom crystal identification is not feasible, the task force recommends the use of conventional and/or advanced imaging techniques to help the physician diagnose gout. Since the last EULAR recommendations, major advances in imaging of gout have been made, particularly with regard to US, DECT, conventional CT and MRI.⁴³ All can detect urate deposition, tophi and bone erosion, but there are still uncertainties about the best imaging modality for diagnosing gout. The task force agreed that although all have their strengths

and weaknesses, overall US offers the best potential for diagnosing gout (see the sixth recommendation). DECT is promising and has the advantage of differentiating MSU crystal deposition from connective tissues and from calcium containing mineral deposits by their specific X-ray attenuation properties.^{43 44} DECT can also quantify the MSU crystal deposition burden in and around joints,^{45–47} and explore ‘deep-seated’ anatomical structures/regions (eg, spine). The metrological properties of DECT for the diagnosis of gout have been assessed in several studies summarised in table 3. It should be noted that many of these studies have included patients at an advanced stage of the disease, that is, patients often with tophaceous and/or erosive gout with chronic gouty arthritis. Sensitivity of DECT in patients with early disease or without tophi tends to be lower.^{40 48–50}

(6) *Plain radiographs are indicated to search for imaging evidence of MSU crystal deposition but have limited value for the diagnosis of gout flare. US scanning can be more helpful in establishing a diagnosis in patients with suspected gout flare or chronic gouty arthritis by detection of tophi not evident on clinical examination, or a double contour (DC) sign at cartilage surfaces, which is highly specific for urate deposits in joints.* Typical radiographic features include: bone erosions with overhanging edges and a sclerotic rim; bone proliferation; joint space narrowing, which occur late in the disease course; and soft-tissue masses, sometimes calcified, corresponding to soft-tissue tophi. Plain radiographic changes take several years to develop, so they may be helpful in supporting a diagnosis of gout in the later stages of the disease. In patients with 4 years duration of disease, sensitivity and specificity for erosions were 0.12 and 0.96.²⁴

US is of major interest for the diagnosis of gout because of its low cost, widespread availability and absence of radiation exposure. Its diagnostic performance with MSU crystal detection as reference test has been assessed in several studies.^{51–58} MSU crystal deposition can be detected in different ways: at the surface of the articular cartilage as a hyperechoic enhancement—the DC sign—within the joint space as floating hyperechoic foci with the appearance of a snowstorm; and within the joint or along tendons as hyperechoic aggregates suggestive of tophi. Sensitivity and specificity of these US features are summarised in table 4. Data from the largest US study showed that these features have high specificity (84%) and that the DC sign and

Table 4 Sensitivity and specificity of US features for the diagnosis of gout

	Sensitivity	Specificity	PPV	NPV
Tophus				
Ogdie <i>et al</i> 2015 ⁴³ MA*	0.65 (0.34–0.87)	0.80 (0.38–0.96)		
Ogdie <i>et al</i> 2017 ⁵¹				
Early disease (<2 years)	0.33 (0.25–0.43)	0.95 (0.91–0.97)	0.80 (0.65–0.90)	0.72 (0.66–0.77)
Late disease (≥2 years)	0.50 (0.44–0.56)	0.95 (0.91–0.97)	0.93 (0.88–0.97)	0.57 (0.51–0.62)
Double contour sign				
Ogdie <i>et al</i> 2015 ⁴³ MA†	0.83 (0.72–0.91)	0.76 (0.68–0.83)		
Ogdie <i>et al</i> 2017 ⁵¹				
Early disease (<2 years)	0.50 (0.41–0.60)	0.92 (0.87–0.95)	0.78 (0.67–0.87)	0.77 (0.71–0.82)
Late disease (≥2 years)	0.63 (0.57–0.68)	0.91 (0.86–0.94)	0.91 (0.86–0.94)	0.63 (0.57–0.68)
Snowstorm appearance				
Ogdie <i>et al</i> 2017 ⁵¹				
Early disease (<2 years)	0.32 (0.23–0.42)	0.90 (0.85–0.94)	0.68 (0.58–0.77)	0.64 (0.60–0.68)
Late disease (≥2 years)	0.29 (0.24–0.35)	0.92 (0.89–0.94)	0.88 (0.78–0.94)	0.11 (0.57–0.19)

Diagnosis of gout was based on MSU crystal identification in all referenced studies.

*Pooled data from: ^{53 55–57 105}

†^{53–56}

MA, meta-analysis; MSU, monosodium urate; NPV, negative predictive value; PPV, positive predictive value.

US imaging of tophi perform better than the snowstorm appearance.⁵¹ Overall, sensitivity of the US features is lower in early versus late disease⁵¹ (table 4). The US DC sign, the identification of MSU crystal deposition by DECT and imaging evidence of gout-related joint damage with radiographs are all now included in the American College of Rheumatology (ACR)/EULAR 2015 gout classification criteria.²⁸

(7) *Risk factors for chronic hyperuricaemia should be searched for in every person with gout, specifically: chronic kidney disease (CKD); overweight, medications (including diuretics, low-dose aspirin, cyclosporine and tacrolimus); consumption of excess alcohol (particularly beer and spirits), non-diet sodas, meat and shellfish.*

The task force emphasises that once a diagnosis of gout is made, identification of person-specific risk factors for hyperuricaemia is crucial,^{59–61} as some of these are modifiable. For instance, there is evidence that gradual weight loss in obese patients lowers SUA levels and reduces the likelihood of gout flare.^{62–63} However, as previously emphasised in the EULAR recommendation for the management of gout,¹⁵ the level of evidence to support the effect of lifestyle modification alone on SUA levels is low.^{64–65} Nevertheless, because of the high prevalence of cardiovascular comorbidities in patients with gout,⁶⁶ implementations of lifestyle modifications are recommended for cardiovascular disease prevention, while prevention of KD and excess alcohol consumption also need to be addressed separately.

(8) *Systematic assessment for the presence of associated comorbidities in people with gout is recommended including obesity, renal impairment, hypertension, ischaemic heart disease, heart failure, diabetes and dyslipidaemia.*

This recommendation underlines the importance of screening and managing comorbidities frequently associated with gout, as previously emphasised in the EULAR recommendations for the management of gout.¹⁵ The identification of some of these comorbidities, particularly CKD and cardiovascular diseases is crucial as it has therapeutic implications.¹⁵

DISCUSSION

This paper provides eight key recommendations for the diagnosis of gout to all physicians, including GPs, on the basis of an SLR and a Delphi consensus involving both experts and patients.

The task force recommends a three-step approach for the diagnosis of gout (figure 1). The first step relies on MSU crystal identification when SF analysis is feasible. If not possible, the second step relies on a clinical diagnosis based on suggestive and associated clinical features of gout and presence of hyperuricaemia. When a clinical diagnosis of gout is uncertain and crystal identification is not possible, the third step recommends imaging, particularly US, to search for imaging evidence of MSU crystal deposition.

In these updated EULAR recommendations, the identification of crystals using polarising microscopy remains the gold standard for the diagnosis of gout owing to its 100% specificity. It is a single sufficient criterion for gout classification according to the 2015 ACR/EULAR gout classification criteria.²⁸ However, the task force acknowledges that this may have some limitations in a primary care setting where most patients with gout are diagnosed and treated. Indeed, microscopic SF analysis requires both expertise and equipment that are not readily accessible for all physicians. Another barrier is the required expertise in joint puncture, and the challenge of

aspirating SF, without patient discomfort, from small joints or from certain anatomical regions such as the midfoot and wrist.

In patients suffering from acute arthritis and in whom SF analysis is not feasible, the task force recommends that the diagnosis of gout flare should be based both on certain suggestive clinical features and the SUA level. The task force considered that the level of evidence to support the use of any of the published algorithms^{26–28–30} was not sufficient for the diagnosis of gout in patients suffering from an acute arthritis. Apart from the Janssens' criteria which were developed for use in clinical practice,²⁶ the other recent algorithms^{28–30} were developed to classify patients and not to make a diagnosis at the individual level. In addition, for some of them,^{26–30} it has been shown that disease duration impacts on their performance, with lower specificity in established gout.⁶⁷ The six clinical features selected in the second recommendation are derived from several algorithms, particularly the Janssens' rule and the SUGAR study, because these had the best metrological performance among all the assessed variables when compared with crystal identification as reference.^{24–26}

In the second recommendation, the task force draws attention to the value of SUA levels for the clinical diagnosis of gout. Although there is no accepted definition of hyperuricaemia,⁹ the 6 mg/dL (360 µM) threshold has been proposed because the lifelong risk of gout increases above this level.⁶⁸ In the SUGAR study, the OR of having gout versus not having gout was close to 6 for SUA levels between 6 and 8 mg/dL, while this OR rose to 39 for SUA levels above 10 mg/dL.²⁴ However, as highlighted in the fourth recommendation, hyperuricaemia alone should not be used to diagnose gout, and should only be considered when there are suggestive clinical features for the diagnosis of gout. In general, crystallisation of MSU occurs when the SUA level exceeds its saturation point. This is not precisely known but it seems close to 6 mg/dL. However, nucleation and deposition of MSU crystals are very slow processes depending on multiple genetic and environmental factors including tissue nucleators and inhibitors. Among these, persistently high SUA levels are crucial.⁶⁹ Thus, hyperuricaemia is a strong predictor of incident gout but not all patients with asymptomatic hyperuricaemia will develop gout. For instance, a recent study found that only half of patients with SUA levels above 10 mg/dL will develop gout over 15 years.³²

The last decade has brought major advances in our understanding of the natural history of gout. In particular, the identification of a continuum between a preclinical state defined by asymptomatic MSU crystal deposition within joints and tendons, and occurrence of the first gout flare (figure 1), has been facilitated by the use of novel imaging such as US⁷⁰ and DECT.⁴⁰ This new knowledge has prompted the proposal of a novel staging for gout, which allows a diagnosis during the so-called intercritical period.^{8–9–42}

Among imaging modalities, US has been the most investigated, particularly in the SUGAR study⁵¹ and by Outcome Measures in Rheumatology (OMERACT).^{71–73} US features, notably the DC sign, have high specificity and good sensitivity,^{71–73} although the specificity is not so high in early gout. One retrospective study found that the DC sign cannot reliably distinguish gout from calcium pyrophosphate deposition disease.⁷⁴ However, these findings were not subsequently confirmed by Ogdie *et al*, who found that the DC sign still had a high specificity 92.9% (85.8–97.1) when compared with subjects with CPDD.⁵¹ Since the advantages of US include low cost, lack of radiation exposure, ease of use and increasing

availability in clinical practices, the task force prioritised US over other imaging modalities. In addition, US can identify associated inflammation using the Doppler mode.⁵⁸ Since the sites for scanning varied across studies and because of a lack of standardisation, the task force recommends screening affected joints and at least both first MTP joints and the knees, which are common sites for MSU crystal deposition. US can also facilitate SF aspiration and MSU crystal identification from joints with US evidence for urate deposits, but without clinical effusion or inflammation.⁷⁵

DECT also allows non-invasive detection and characterisation of MSU crystal deposition in joints and soft-tissues.⁷⁶ This technique may be helpful particularly in cases where US is not feasible or technically complicated (eg, spinal gout). However, it is not widely available, and in addition to being expensive and involving some radiation exposure, its use is often restricted to secondary and tertiary care centres. Its diagnostic performance, with MSU crystal identification as reference test, seems comparable to US, with a potential superiority for MSU crystal deposition detection in direct comparison with US.^{77–80} As observed with US, sensitivity for the diagnosis of gout is influenced by the duration of the disease, being lower in the early stage of the disease.^{40 48–50} Size and density of tophi also seem to influence the sensitivity for MSU crystal deposition detection.^{81 82} Lastly, reading and interpretation of images from DECT require skill and expertise, and artefacts that could lead to false positivity have been reported.^{81 82}

Both US^{83 84} and DECT⁸⁵ could be useful to assess tophus resolution in response to ULT.

MRI and conventional CT both have the ability to identify MSU crystal deposition. However, their diagnostic performance has been less studied than US and DECT. MRI provides information with regard to the size of tophi, crystal-induced inflammation such as synovitis, and joint damage including bone erosion.^{86–90} CT can also identify urate deposits but is more efficient in visualising bone damage.^{91–93} Therefore, the task force agreed that CT and MRI have limited utility for the diagnosis of gout in clinical practice, as compared with US or DECT.

As in the EULAR recommendations for the treatment of gout,¹⁵ the task force has emphasised in its two last recommendations the need to search for risk factors for hyperuricaemia once gout is diagnosed. Importantly, some risks factors, notably obesity,^{63 94} medications (diuretics, low-dose aspirin, cyclosporine, tacrolimus) and diet are potentially modifiable.¹⁵ Lastly, the task force underlines the importance of screening for several comorbidities, in particular obesity, CKD, cardiovascular diseases and components of the metabolic syndrome, which frequently coexist in patients with gout, but for which causality remains controversial.⁹⁵

In conclusion, since the EULAR recommendations for the diagnosis of gout were published in 2006, major advances have been made with regard to imaging, clinical diagnosis and understanding of the natural history of the disease. The EULAR recommendations have therefore been revised and updated in the light of these advances in order to better assist the physicians in diagnosing gout.

Author affiliations

- ¹Service de Rhumatologie, Hopital Lariboisiere Centre Viggo Petersen, Paris, France
- ²Inserm UMR1132 Bioscar, Université Paris Diderot UFR de Medecine, Paris, France
- ³Academic Rheumatology, University of Nottingham, Nottingham, UK
- ⁴Rheumatology, Hospital General Universitario de Alicante, Alicante, Spain
- ⁵Institute of Rheumatology, RAMS, Moscow, Russian
- ⁶Radiology, Lausanne University Hospital, Lausanne, Switzerland

- ⁷AP-HP, Hôpital Pitié-Salpêtrière, Département Biostatistique Santé Publique et Information Médicale, Centre de Pharmacopépidémiologie (Cephepi), INSERM, UMR 1123 ECEVE, CIC-1421, Paris, France, Paris, France
- ⁸Nottingham, UK
- ⁹Département d'Epidémiologie et Recherche Clinique, Paris, France
- ¹⁰Rheumatology, VieCuri, Venlo, Netherlands
- ¹¹Department of Primary and Community Care, Radboud University Medical Center, Radboud Institute for Health Sciences, Nijmegen, The Netherlands
- ¹²Department of Rhumatologie, Hôpital Lariboisière, Paris, France
- ¹³INSERM UMR-1132 and Université Paris Diderot, Paris, France
- ¹⁴Arthritis Research UK Primary Care Centre, Keele University, Keele, UK
- ¹⁵Centre Molecular Medicine, University of Edinburgh, Edinburgh, , UK
- ¹⁶Servicio de Reumatología, Hospital de Cruces, Baracaldo, Spain
- ¹⁷Rheumatology Unit, Clínica Coração de Jesus, Lisbon, Portugal
- ¹⁸Department of Medicine, University of Padua, Padua, Italy
- ¹⁹Musculoskeletal Medicine, Service de RMR, Lausanne, Switzerland
- ²⁰Department of Internal Medicine, Section of Rheumatology, University Clinic Carl Gustav Carus, Dresden, , Germany
- ²¹Rheumatology, Diakonhjemmet Hospital, Oslo, Norway
- ²²Institute of Rheumatology, Prague, , Czech Republic
- ²³Academic Rheumatology, Nottingham University, Nottingham, UK
- ²⁴Biostatistics and epidemiology, APHP, Hopital Pitié Salpêtrière, Paris, France
- ²⁵Rheumatology, Assistance Publique - Hopitaux de Paris, Paris, France

Acknowledgements This paper is dedicated to the memory of VB. The task force thanks EULAR for financial and logistic support. Part of this paper was presented as a poster during the Eular Congress (Paris, 2014) (10.1136/annrheumdis-2014-eular.5546).

Collaborators Augustin Latourte.

Contributors JC, SG and PR performed the systematic literature review under the supervision of FT. All authors were part of the Task Force, participated in the Delphi rounds and attended the face-to-face meeting. TB organised and chaired the meetings, along with MD and EP. PR wrote the manuscript, with contribution and approval of all coauthors.

Funding This project was funded by the European League Against Rheumatism.

Disclaimer The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health and Social Care.

Competing interests PR has received honoraria from Ipsen/Menarini, AstraZeneca, Savient and Grünenthal. MD has received honoraria for ad hoc advisory boards on gout or osteoarthritis from Ardea Biosciences, AstraZeneca, Nordic Biosciences and Roche and was CI for a Nottingham University Investigator-led non-drug study on gout funded by AstraZeneca. EP received fees from Ipsen/Menarini and Astra-Zeneca. JC and SG are employed by the Centre de Pharmacopépidémiologie (Cephepi) of the Assistance Publique – Hôpitaux de Paris that has received research funding, grants and fees for consultant activities from a large number of pharmaceutical companies, which have contributed indiscriminately to the salaries of its employees. TJ received fees for lectures and/or advisory boards from Ardea Biosciences, Astra/Zeneca global, Abbvie, BMS, Celgene, Grünenthal, Janssen, Lilly, Menarini International, Novartis, Pfizer, Roche, UCB. FL received fees for advisory boards: Ardea BioSciences, Astra-Zeneca, Ipsen Pharma, Menarini, Novartis, Savient, Mayoly-Spindler. He also received unrestricted grants for organising the European Crystal Network Workshops (Convenor Frédéric Lioté, France& Alexander So, Switzerland) since 2010: Ardea BioSciences, Astra-Zeneca, Ipsen Pharma, Mayoly-Spindler, Menarini, Novartis, Savient, SOBI. He received fees for lectures from Ardea BioSciences, Grünenthal, Ipsen Pharma, Menarini France, Novartis Global. CDM is funded by the NIHR Collaborations for Leadership in Applied Health Research and Care West Midlands. CDM is also funded by the NIHR School for Primary Care Research and an NIHR Research Professorship in General Practice (NIHR-RP-2014-04-026). NEF, an NIHR Senior Investigator, was funded by an NIHR Research Professorship (NIHR-RP-2011-015). GN has received fees for advisory boards and consultations from Savient, Ipsen, Menarini and Grünenthal and indirectly from Ardea Biosciences and AstraZeneca for work on the IDMC's for trials of lesinurad. FP-R has received fees from AstraZeneca, Grünenthal, Horizon, Menarini, Dyve Bioscience, Japan Tobacco, Logarithm, Astellas. LP has received consulting or speaker fees from Menarini, Fidia, Grünenthal, BMS. AKS has served as consultant to Astra-Zeneca and SOBI in regard to the treatment of gout. A-KT received fees from Berlin Chemie-Menarini, Novartis, AstraZeneca/Ardea Biosciences and Grünenthal Pharma. TU received fees from AstraZeneca/Ardea, Grünenthal Pharma and Novartis JZ received fees from Berlin Chemie-Menarini and Novartis. WZ received honorarium from Savient, AstraZeneca and Grünenthal. FT is head of the Centre de Pharmacopépidémiologie (Cephepi) of the Assistance Publique – Hôpitaux de Paris that has received research funding, grants and fees for consultant activities from a large number of pharmaceutical companies, that have contributed indiscriminately to the salaries of its employees. She did not receive any personal remuneration from these companies. TB has received consulting fees, speaker fees or grants from Ipsen Pharma, Menarini, AstraZeneca, Novartis, Sobi, Savient, Grünenthal and Cymabay.

Patient and public involvement statement Two patients with gout were involved in the elaboration of these recommendations. They participated in the face-to-face meeting, in the Delphi rounds and approved the manuscript.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement The data analysed in the current study are available from the corresponding author on request.

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