

Review

Gout

Novel therapies for treatment of gout and hyperuricemia

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Abstract

In the past few decades, gout has increased not only in prevalence, but also in clinical complexity, the latter accentuated in part by a dearth of novel advances in treatments for hyperuricemia and gouty arthritis. Fortunately, recent research reviewed here, much of it founded on elegant translational studies of the past decade, highlights how gout can be better managed with cost-effective, well-established therapies. In addition, the advent of both new urate-lowering and anti-inflammatory drugs, also reviewed here, promises for improved management of refractory gout, including in subjects with co-morbidities such as chronic kidney disease. Effectively delivering improved management of hyperuricemia and gout will require a frame shift in practice patterns, including increased recognition of the implications of refractory disease and frequent noncompliance of patients with gout, and understanding the evidence basis for therapeutic targets in serum urate-lowering and gouty inflammation.

Introduction

In the past few decades in the USA and elsewhere, gout has markedly increased in incidence and prevalence [1-3]. This includes a marked increase in gout in patients over the age of 65, and even more so in patients over 75 years of age, in lockstep with high prevalence of conditions linked with hyperuricemia (chronic kidney disease (CKD), hypertension, metabolic syndrome and diabetes, and congestive heart failure) and rampant use of diuretics and low dose acetylsalicylic acid [1-3]. Gout patients in this day and age are more clinically complex than in past memory, due to various combinations of advanced age, co-morbidities, potential drug-drug interactions, and refractory tophaceous disease [1]. In this light, clinicians are increasingly faced with patients with refractory gout, classic features of which are summarized in Table 1. Until recently, a lack of an innovative pipeline of emerging therapies for hyperuricemia and gouty inflammation has compounded this situation. This review frames what we

have recently learned regarding how the current scope of therapeutics for gout and hyperuricemia can be employed more effectively, and in particular for refractory gouty inflammation and hyperuricemia, focusing on new urate-lowering drugs (febuxostat and uricases) and biologic approaches to gouty inflammation via IL-1 inhibition.

Gout therapy: how the current armamentarium is actually employed in the 'real world'

Table 2 summarizes recent assessment of the scope of application of existing therapies for gout in the USA [4], and also highlights that primary care practitioners are, by far, prescribing the most gout therapies. Given that there are currently estimated to be at least approximately 3 million people with active gout, and 3 to 6 million subjects with a history of gout in the USA [5], the numbers summarized in Table 1 suggest that many gout patients receive inadequate therapy. In this context, there appears to be a shortfall in meeting practice guidelines [6,7] for prescribing of prophylactic colchicine relative to the allopurinol prescription numbers. Overall, the estimated colchicine utilization rate was only 4.6% in office visits for those with gout, versus 8.9% for prednisone and 18% for NSAIDs [4]. As it is elsewhere in the world, allopurinol is the first line choice for serum urate-lowering in the great majority of subjects in the USA. However, there appear to be large differences in prescribing patterns for allopurinol in Caucasians relative to both African-Americans and Asians, suggesting under-treatment of gout in the latter two subgroups.

Advances in treatment of gouty arthritis by better use of the current drug armamentarium

Acute gouty arthritis is mediated by the capacity of monosodium urate crystals to activate multiple pro-inflammatory

ABCG2 = ATP-binding cassette sub-family G member 2; CKD = chronic kidney disease; EULAR = European League Against Rheumatism; FDA = Food and Drug Administration; GLUT = glucose transporter; IL = interleukin; NSAID = nonsteroidal anti-inflammatory drug; PEG = polyethylene glycol; URAT, urate transporter.

Table 1

Common features of ‘treatment-refractory’ gout that complicate management

Polyarticular gout, uncontrolled flare activity, and/or chronic synovitis
Destructive tophi
Advanced age
Co-morbidities (for example, chronic kidney disease, cardiovascular disease, obesity, metabolic syndrome or diabetes, alcohol abuse)
Polypharmacy and drug interactions (for example, statins, macrolide antibiotics, oral anticoagulants)
Contra-indications or refractoriness to NSAIDs, colchicines, and/or glucocorticosteroids
Allopurinol intolerance or hypersensitivity and inability to employ uricosurics
Failure to adequately lower serum urate on appropriate doses of urate-lowering drugs

NSAID, nonsteroidal anti-inflammatory drug.

Table 2

Overview of recent treatment patterns of gout in the USA

Total ambulatory visits, and visits to primary care versus specialists	
Total number of ambulatory care visits	973 million
Number of visits for gout	3.9 million (0.4% of total)
Percentage of total visits for gout to:	
Primary care	69%
Cardiologists	10%
‘Other specialists or unknown’	<16%
Rheumatologists	<2%
Number of gout patient-specific anti-inflammatory prescriptions (absolute number of prescriptions/year)	
Colchicine	~381,000
NSAIDs	~700,000
Prednisone	~341,000
Number of gout patient-specific urate-lowering prescriptions (absolute number of prescriptions/year)	
Allopurinol	2.8 million
Probenecid	8,000
Demographics of allopurinol prescribing: percentage of gout patients that are:	
Caucasian	80%
African Americans	49%
Asians	19%

Data from the 2002 calendar year extracted from the work of Krishnan *et al.* [4]. NSAID, nonsteroidal anti-inflammatory drug.

pathways in the joint, culminating in early activation of resident macrophages, and neutrophil adhesion, migration into the joint, and activation in the synovium and joint space that drive gouty inflammation [8,9]. Current primary options for anti-inflammatory management of acute gout (nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, and colchicine) bluntly dampen these inflammatory mechanisms in a cost-effective manner, though are limited by broad drug toxicities, particularly in subjects with significant co-morbidities [8-13]. Moreover, the evidence basis for some of these treatments has been limited by inadequate assessment in randomized, controlled, double-blind clinical trials, an issue due to the intrinsic self-limitation of the acute gout flare.

The recent definition of etoricoxib as an effective COX-2-selective inhibitor in acute gout [14] has opened up a new therapeutic approach, but the cardiovascular safety of COX2 inhibitors remains under review. The establishment, in the past 2 years, of the evidence basis for oral glucocorticosteroid treatment of acute gout is also particularly significant, for example, for subjects with CKD. Specifically, prednisolone 35 mg daily for 5 days and naproxen 500 mg twice daily for 5 days have been demonstrated to be comparable in efficacy and tolerance in a recent trial of acute gout treatment [11]. Prednisolone (6 doses of 30 mg over 5 days) was also comparable in efficacy to indomethacin and better tolerated in an acute gout trial [12].

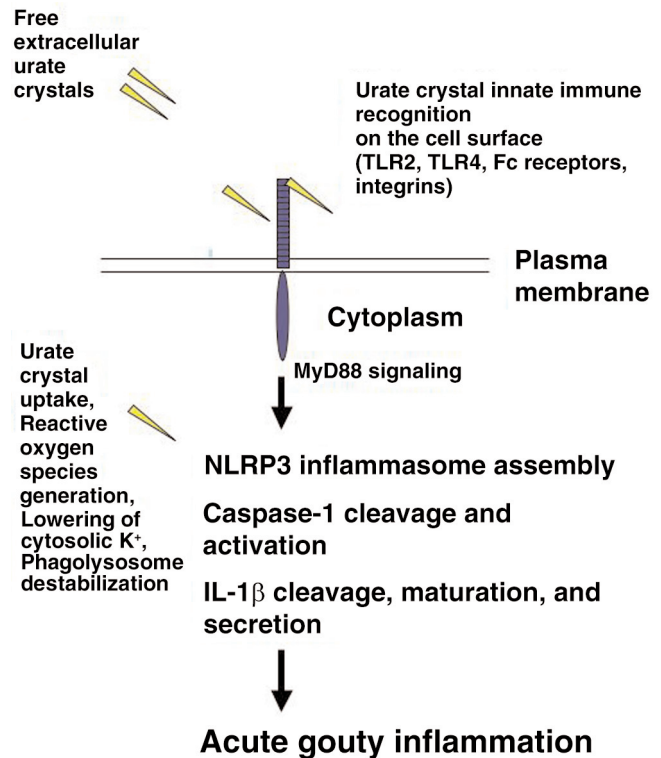
Dosing guidelines and the evidence basis for colchicine in acute gout treatment have also advanced in the past few years. In older oral colchicine regimens where the drug was given every 1 to 2 hours repeatedly for multiple doses, gastrointestinal toxicity, including severe diarrhea, was limiting, and occurred before a 50% reduction in pain was achieved in most subjects [13]. Intense colchicine regimens have justifiably fallen out of favor. As an example, European League Against Rheumatism (EULAR) expert consensus guidelines for oral colchicine in acute gout are for a maximum of three colchicine 0.5 mg tablets per 24 hour period [6]. Furthermore, in a large, randomized, controlled multicenter trial comparing low dose and extended dose colchicine regimens, results strongly supported the lower dose colchicine regimen for acute gout [15]. In this study, within 12 hours of onset of acute gout symptoms patients self-administered 'high dose' colchicine (1.2 mg followed by 0.6 mg every hour for 6 hours (4.8 mg total)) or 'low dose' colchicine (1.2 mg followed by 0.6 mg in 1 hour (1.8 mg total)), or placebo. The 'low dose' colchicine was comparable to 'high dose' colchicine in efficacy, but did not differ from placebo with respect to diarrhea or other gastrointestinal side effects.

Advanced anti-inflammatory therapies for gout

The typical response of acute gout to NSAIDs and COX2 selective inhibition therapy, systemic glucocorticosteroids, and colchicine is rapid but incomplete (for example, approximately 50% pain reduction achieved within 2 to 3 days [11,12,14,15]). This has left substantial room for improvement, particularly since a potent alternative, intravenous colchicine, was justifiably withdrawn from active marketing in the USA in 2008 due to serious safety considerations. Among selective targets or strategies for advanced anti-inflammatories for gouty inflammation identified in recent years are the complement C5b-9 membrane attack complex, agonism of phagocyte melanocortin receptor 3 (shown to be a direct peripheral target of adrenocorticotrophic hormone), the chemokines CXCL1 and CXCL8, tumor necrosis factor- α , and the NLRP3 (NLR family, pyrin domain containing 3) inflammasome (Figure 1), which, via caspase-1 activation, drives IL-1 β endoproteolysis and consequent IL-1 β maturation and secretion [8,9].

Though anecdotal reports have suggested tumor necrosis factor- α antagonism to be beneficial in some cases of refractory human gouty inflammation [16], IL-1 β appears to be far more central than tumor necrosis factor- α in experimental urate crystal-induced inflammation in mice [17]. Concordantly, the most investigated biologic drug strategy in humans for gouty inflammation has been neutralization of IL-1, with promising results [9,17]. A pilot study of ten patients with chronic refractory gouty inflammation given the soluble IL-1 receptor antagonist anakinra (100 mg daily subcutaneously for 3 days) suggested good overall responses [17], though results of larger, randomized, controlled studies of IL-1 inhibition for gouty arthritis are awaited.

Figure 1



The NLRP3 inflammasome and IL-1 β processing and secretion in crystal-induced inflammation. The figure shows monosodium urate crystal interaction with phagocytes, with crystal recognition at the macrophage surface mediated by innate immune mechanisms, in part employing Toll-like receptor (TLR)2 and TLR4 and associated MyD88 signaling, Fc receptors, and integrins. Crystal uptake with consequent phagolysosome destabilization, and reactive oxygen species generation and lowering of cytosolic K⁺ all appear to promote activation of the NLRP3 (cryopyrin) inflammasome. Consequent endoproteolytic activation of caspase-1, which drives pro-IL-1 β maturation, and consequent secretion of mature IL-1 β is a major mechanism stimulating experimental gouty inflammation, and appears to be implicated in human gouty arthritis, as discussed in the text.

Options in the treatment of hyperuricemia: recent establishment of the evidence basis for <6 mg/dL as the serum urate target level in gout

Pharmacologic urate-lowering approaches can employ primary, potent uricosurics (probenecid or benzbromarone), xanthine oxidase inhibitors to inhibit uric acid generation (allopurinol and the recently approved drug febuxostat), or experimental uricase treatment (with Rasburicase™ or pegloticase) to degrade urate [1,10]. Since the solubility of urate in physiologic solutions is exceeded at approximately 6.7 to 7.0 mg/dL, current guidelines for inhibiting ongoing urate crystal deposition, reduction of total body urate stores, and resolution of macroscopic tophi are for continuing (lifelong) reduction of serum urate concentration to <6 mg/dL

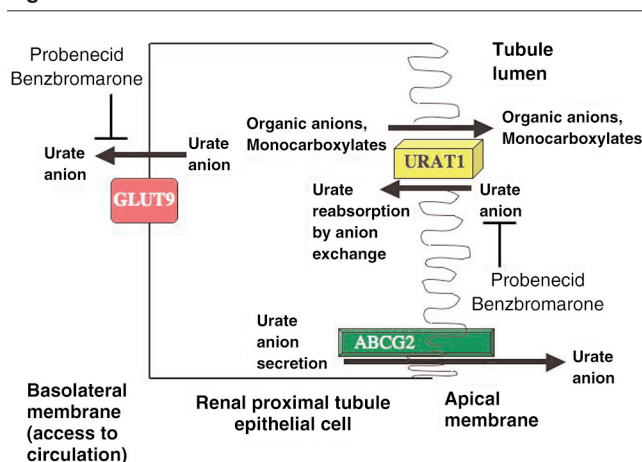
(approximately 360 mmol/l), and ideally in the 5 to 6 mg/dL range [18]. As summarized in a detailed, recent review [18], achieving this target level in gout patients ultimately is associated with fewer gout flares, and it also may have direct and indirect beneficial effects on renal function [19,20]. A more aggressive serum urate-lowering target such as 3 to 5 mg/dL appears appropriate for more rapid tophus debulking in those tophaceous gout patients with a body burden of urate assessed to be particularly large [21].

Advances in understanding renal urate handling and uricosuric therapy

Uricosurics act primarily by inhibiting proximal renal tubule epithelial cell reabsorption of urate anion, thereby enhancing renal uric acid excretion. This remains a compelling approach in some aspects, since reduction of miscible total body urate stores is initiated rapidly, and the velocity of tophus size reduction is comparable to that using allopurinol when similar degrees of serum reduction are achieved [22]. Moreover, uricosurics target the underlying basis for hyperuricemia in the majority of patients. Recent advances in understanding renal disposition of urate include identification of the anion exchanger URAT1 (urate transporter 1; SLC22A12) as a mediator of urate anion reabsorption from the lumen at the apical membrane of the proximal tubule epithelial cell [23], with the electrogenic hexose transporter GLUT9 (glucose transporter 9; SLC2A9) mediating urate anion reabsorption into the peritubular interstitium (and ultimately into the circulation) at the basolateral membrane [24-28]. Probenecid and benzbromarone both inhibit urate anion movement transduced by URAT1 and GLUT9 [24] (Figure 2). The findings related to GLUT9 also raise compelling questions about the relationships between hyperglycemia and increased fructose intake and hyperuricemia [24-29].

A major advance that also could point to new and potentially genomics-customized uricosuric strategies is the identification of ATP-binding cassette sub-family G member 2 (ABCG2) as one of the functional urate anion secretory transporters at the apical membrane of the renal proximal tubule epithelial cell (Figure 2) [30]. Moreover, genome-wide association studies have linked common URAT1, GLUT9, and now ABCG2 haplotypes or single nucleotide polymorphisms with altered susceptibility to gout [23-28,30]. For example, the common ABCG2 rs2231142 single nucleotide polymorphism encoding the Q141K mutation in the nucleotide-binding domain of ABCG2 suppresses ABCG2 urate transport rates by approximately 50% *in vitro*, and in a large, population-based study, rs2231142 was strongly associated with serum urate levels in whites, who have a minor allele frequency of 0.11 [30]. The adjusted odds ratio for gout of 1.68 per risk allele in whites and blacks argues that approximately 10% of all gout cases in whites may be attributable to ABCG2 rs2231142, and the risk allele also is highly prevalent in Asians, who have a higher gout prevalence than whites [30].

Figure 2



Effects of URAT1, GLUT9, and ABCG2 on urate anion disposition by the renal proximal tubule epithelial cell and inhibitory effects of the uricosurics probenecid and benzbromarone on renal urate reabsorption by inhibition of both URAT1 and GLUT9. The schematic summarizes the effects of the uricosurics probenecid and benzbromarone on urate handling in the renal proximal tubule epithelial cell by the URAT1 (SLC22A12) and GLUT9 (SLC2A9) transporters identified as linked with serum urate levels and gout susceptibility in genetic studies, including recent genome-wide association studies. Urate reabsorption at the apical membrane, which interfaces with the tubule lumen, is mediated in large part by the anion exchange function of URAT1. At the basolateral membrane, the hexose transport facilitator GLUT9 electrogenically transports urate anion into the peritubular interstitium, where urate is reabsorbed into the circulation. Recent genome-wide association studies and functional genomics analyses have also uncovered a substantial role for ABCG2 in secretion of urate into the proximal tubule lumen. The depicted model is a simplification, since other molecules that affect urate disposition in the proximal tubule and distally in the nephron are not depicted here, and effects of certain other drugs on renal urate disposition by inhibiting URAT1 or GLUT9 or other transporters are not represented. ABCG, ATP binding cassette sub-family G; GLUT, glucose transporter; URAT1, urate transporter 1.

Significantly, in current clinical practice, the most available primary uricosuric, probenecid, requires more than once daily dosing and increases the risk of urolithiasis, particularly in acid urine [31]. More selective and potent uricosurics ideally would have a once daily dosing profile and could be designed such that urolithiasis risk is not unduly elevated. All uricosurics also become less effective and ultimately ineffective with progressively lower glomerular filtration rate [10,31]. This may limit the role of combining uricosurics with xanthine oxidase inhibition in the treatment of refractory hyperuricemia in gout patients since xanthine oxidase inhibition lowers urinary uric acid clearance by excretion. Such a combination approach can normalize serum urate in a substantial fraction of patients on submaximal allopurinol [32]. An approach of this nature, using certain drugs with wider availability than benzbromarone (for example, losartan, and fenofibrate) [33,34] but with less potent uricosuric action than primary uricosurics such as probenecid, has to date

been, at best, only moderately successful, when studied in only small numbers of subjects, as a potential strategy to further lower serum urate where there is suboptimal control with allopurinol. It appears likely that such combination strategies will be particularly constrained in effectiveness in those with stage 3 CKD or worse (creatinine clearance <60 calculated by Cockcroft-Gault equation and adjusted for ideal body weight).

Advances in understanding allopurinol treatment failure

Given the limitations of uricosuric therapy highlighted above, the first line of pharmacologic therapy to lower serum urate for most gout patients is suppression of xanthine oxidase using allopurinol, which, when effective and well-tolerated, is a cost-effective option [6,10]. Allopurinol is US Food and Drug Administration (FDA) approved for doses up to 800 mg daily [35]. Recent expert consensus EULAR guidelines have reinforced FDA dosing guidelines for allopurinol in patients with preserved renal function [6,35], specifically to initiate allopurinol at 100 mg daily, and then to increase the dose by 100 mg every 1 to 4 weeks until a target serum urate level (<6 mg/dL) is achieved or the maximum appropriate allopurinol dose is reached. FDA dosing guidelines have also advocated 200 to 300 mg allopurinol daily as adequate for most patients with mild gout, and an average dose of 400 to 600 mg allopurinol daily as the expected amount to control hyperuricemia in patients with moderately severe tophaceous gout [35]. In small studies of gout patients, the mean daily dose of allopurinol needed to normalize serum urate was 372 mg [36], and allopurinol dose increases from 300 mg to 600 mg daily markedly increased serum urate-lowering efficiency in patients without stage 3 or worse CKD [37]. Data from recent, large, randomized, controlled clinical trials indicated that allopurinol 300 mg daily lowered serum urate by approximately 33% in a population of gout patients where approximately 25 to 30% had detectable tophi, serum urate was approximately 9.5 to 10 mg/dL, and renal function was largely intact [38,39].

In clinical practice, noncompliance with allopurinol has recently been elucidated to be a problem in approximately 50% of subjects in the first year of therapy [40]. Moreover, it appears that allopurinol is widely under-dosed overall in clinical practice, since the vast majority of allopurinol prescriptions are for 300 mg daily or less [41]. This circumstance reflects influential maintenance dosing guidelines for allopurinol in CKD dating from the 1980s and calibrated for serum levels (in relationship to estimated glomerular filtration rate) of oxypurinol, which is the major, long-lived active allopurinol metabolite and is primarily excreted by the kidney [35]. The intent of the older guidelines was to lessen the incidence of allopurinol hypersensitivity syndrome, particularly with CKD [35]. These guidelines are now recognized not to be based on evidence, to fail to adequately treat hyperuricemia, and also to fail to prevent allopurinol hypersensitivity

syndrome in all patients, including those with CKD [35,42]. Though HLA-B58 is a newly identified risk factor for severe cutaneous adverse reactions to allopurinol (that is, Stevens-Johnson syndrome or toxic epidermal necrolysis) [43-45], there remains no reliable way to identify whether an individual patient will develop such toxicity on allopurinol [35,42].

FDA and more recent EULAR dosing guidelines for allopurinol have suggested the use of reduced doses in renal failure in order to lessen the risk of drug toxicity [6,35]. For example, the FDA-recommended maximum allopurinol dose is 200 mg daily with a creatinine clearance of 10 to 20 ml/min, and 100 mg daily with a creatinine clearance of <10 ml/min. More recently, dose reduction of allopurinol in moderate CKD was supported via retrospective analysis of renal function-adjusted dosing of allopurinol in relation to drug toxicities [46]. The lack of a definition of safety and tolerability of allopurinol maintenance doses above those previously calibrated for serum oxypurinol levels related to creatinine clearance [46] needs to be considered when weighing the decision to employ more advanced serum urate lowering therapeutic options.

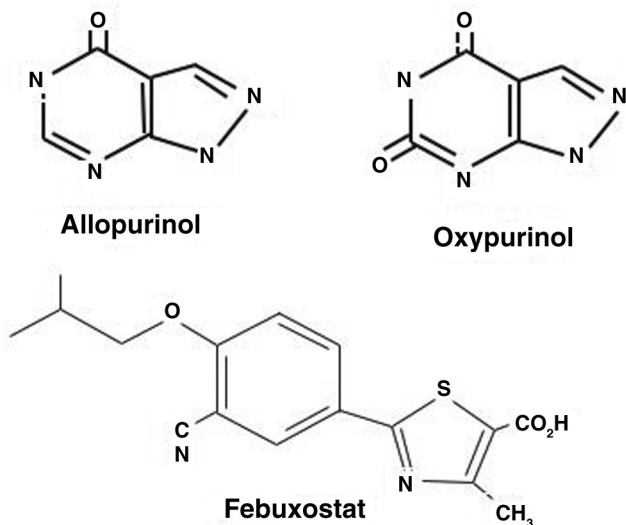
Advanced options for treatment-refractory hyperuricemia in gout: febuxostat

The xanthine oxidase inhibitor febuxostat, now approved in Europe and the USA, is an appropriate choice in circumstances of allopurinol hypersensitivity or intolerance, or failure of allopurinol (at a maximal dose appropriate for the individual patient) to normalize serum urate and, ultimately, improve physical function and quality of life parameters. Febuxostat is a particularly appropriate second line option to allopurinol where uricosuric therapy is contra-indicated, as in stage 3 or worse CKD, and in patients with a history of urolithiasis, an inability to adequately increase hydration, or with identified uric acid overproduction [21].

Febuxostat is a selective inhibitor of xanthine oxidase, the drug sitting in the access channel to the molybdenum-pterin active site of the enzyme [47]. Febuxostat does not have a purine-like backbone, unlike allopurinol and oxypurinol (Figure 3). Significantly, febuxostat is primarily metabolized by oxidation and glucuronidation in the liver and renal elimination plays a minor role in febuxostat pharmacokinetics, as opposed to allopurinol pharmacology. Febuxostat also does not directly regulate pyrimidine metabolism and it is not reincorporated into nucleotides, in contrast to allopurinol, where such properties have the potential to contribute to certain drug toxicities.

Febuxostat 40 to 120 mg daily (and a safety dose trial of 240 mg daily) has now been analyzed in large, randomized, clinical trials in which tophi were seen in approximately 25 to 30% of subjects, with a maximum dose of 300 mg allopurinol employed in comparison groups [38,39,48,49]. Results of all of these trials unequivocally established the failure of 300 mg

Figure 3

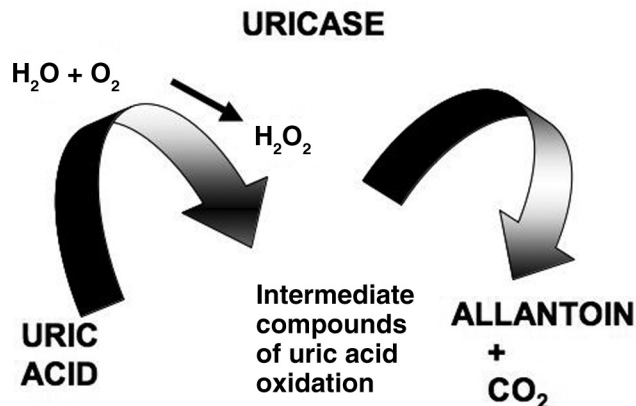


Comparison of allopurinol, oxypurinol, and febuxostat structures. Allopurinol and its long-lived major active metabolite oxypurinol (both pictured) inhibit xanthine oxidase, as does febuxostat (pictured), which, in contrast to the other two agents, does not have a purine-like backbone.

allopurinol daily to achieve a serum urate target level of <6 mg/dL in a substantial majority of the patient population studied. In a 52-week trial, febuxostat 80 and 120 mg both achieved the target level of serum urate <6 mg/dL in the majority of subjects, though gout flare rates at 52 weeks were comparable to those in subjects randomized to allopurinol 300 mg daily [38]. In a second, large phase 3 trial, febuxostat 40 mg daily demonstrated serum urate-lowering to target of <6 mg/dL roughly equivalent to allopurinol 300 mg daily in those with intact renal function, and 80 mg febuxostat daily was superior to allopurinol 300 mg or febuxostat 40 mg daily in achieving a serum urate target level of <6 mg/dL, with comparable drug tolerance [48]. In a subset of patients with stage 2 to 3 CKD, febuxostat 40 and 80 mg daily were also superior in achieving the serum urate target level in comparison to renally dose-adjusted allopurinol (200 to 300 mg daily) [48].

Comparison of early gouty arthritis flares, triggered by serum urate-lowering and putative remodeling, was instructive in these studies. The early flares occurred in association with the most intense serum urate-lowering effect in both febuxostat and allopurinol recipients, and early flares were a greater problem when prophylactic colchicine was stopped at 8 weeks as opposed to 6 months into urate-lowering treatment, but gout flares tapered off later in this study [38,48]. For these reasons, the European Medicines Agency (EMA) astutely recommended gout flare prophylaxis for a 6 month period when febuxostat is initiated.

Figure 4



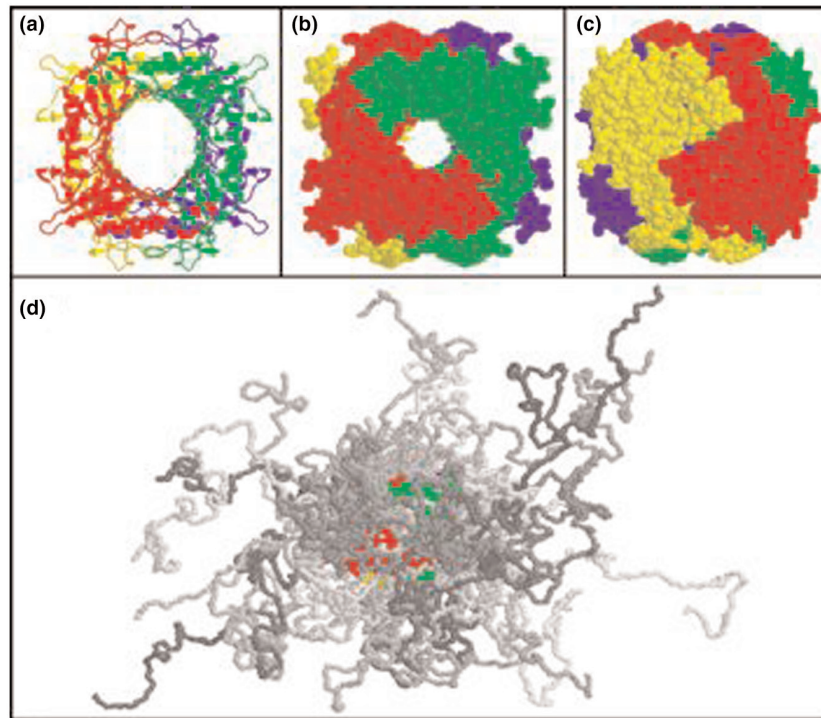
Enzymatic activity of uricase (uric acid oxidase). Uricase oxidizes uric acid, which is sparingly soluble, to the highly soluble end product allantoin, which is readily excreted in the urine. In doing so, uricase generates not only intermediate forms of uric acid that are subject to further metabolism (including 5-hydroxyisourate), but also the oxidant hydrogen peroxide as a byproduct of the enzymatic reaction. During evolution, humans and higher primates lost expression of not only uricase, but also enzymes that rapidly degrade intermediate forms of uric acid generated by uric acid oxidation.

Tophus size is reduced by 50 to 80% after 1 year of either febuxostat or allopurinol treatment, with the greatest tophus and gout flare reduction linked to the greatest degree of serum urate-lowering irrespective of drug. A small, open-label extension study in which patients failing initial therapy on allopurinol were switched to febuxostat to achieve serum urate <6 mg/dL suggested that approximately half of febuxostat-treated patients with tophi can achieve elimination of tophi by 2 years, and approximately 70% by 5 years [49]. Quality of life parameters have been favorably impacted by extended febuxostat treatment in uncontrolled studies [49].

Febuxostat is approved for use in European countries at 80 and 120 mg daily. The FDA approved febuxostat for use in the USA in February, 2009. The USA label is for a dose of 40 mg daily, followed by dose increase to 80 mg daily if serum urate is not normalized after at least 2 weeks. Side effects of febuxostat include rash in <2% of subjects, and elevation of hepatic enzymes, diarrhea, and arthralgia may also occur. As is the case for allopurinol, xanthine oxidase inhibition by febuxostat carries the potential for major drug interactions with azathioprine, 6-mercaptopurine and theophylline [50].

Uricase therapy: an experimental 'biologic' option for serum urate lowering

Uricases oxidatively degrade uric acid, thereby catalyzing conversion to soluble allantoin, which is much more soluble than uric acid [51]. Uricases also generate 1 mole of the oxidant hydrogen peroxide for each mole of uric acid degraded (Figure 4). Uricase expression was lost in humans

Figure 5

Molecular models of the uricase tetramer and of the PEGylated uricase pegloticase containing strands of 10 kDa polyethylene glycol (PEG) linked to each uricase tetramer. **(a)** Schematic model of the uricase tetramer, based on the crystal structure of *Aspergillus flavus* uricase. Each subunit is shown in a different color (red, blue, green, or yellow). **(b)** Space-filling model of the *A. flavus* uricase tetramer, showing the characteristic tunnel (or barrel) structure of the native enzyme tetramer. **(c)** Space-filling model of *A. flavus* uricase tetramer, rotated around the vertical axis so that the tunnel is not visible. **(d)** Space-filling model of the uricase tetramer in the same orientation as in (b) but to which nine strands of 10 kDa PEG per uricase subunit are attached. The structures of the PEG strands (shown in various shades of gray) were generated as described in [54]. The scale of (d) is about half that of (a-c). Figure 5 and the legend are reprinted with permission from [54].

and higher primates during the course of evolution [1]. Illustrating the huge role uricase plays in uric acid homeostasis in mammals, normal serum urate in rodents is approximately 1 mg/dL, whereas it is approximately 10 mg/dL in uricase knockout mice. Moreover, untreated hyperuricemia in uricase knockout mice leads to death by renal failure due to severe uric acid urolithiasis.

Various uricase therapies for hyperuricemia have been attempted experimentally for several decades [52]. For example, recent, limited reports or pilot studies have evaluated the off-label use in severe, chronic gout of the non-PEGylated recombinant fungal enzyme rasburicase [52,53], which is FDA-approved for single course therapy in pediatric tumor lysis syndrome. Unfortunately, rasburicase is both highly antigenic and has a plasma half-life of 18 to 24 hours [52]. Efficacy, tolerability, and sustainability of rasburicase treatment beyond 6 to 12 months appear to be poor for treatment of refractory hyperuricemia in gout [52,53].

A recent advance has been seen in clinical trials of recombinant porcine-baboon uricase (pegloticase); these

trials have evaluated the potential advantages for sustained management of refractory hyperuricemia in gout of PEGylation of this enzyme (Figure 5) to reduce immunogenicity as well as increase circulating half-life [51,54,55]. For refractory tophaceous disease, results to date indicate that intravenous PEGylated uricase treatment has the potential to rapidly decrease the pool size of miscible urate, and also to de-bulk tophi in weeks to months [56] rather than the months to years seen to date with therapy with xanthine oxidase inhibitors at conventional doses. Specifically, in a phase 2 and a pivotal placebo-controlled, randomized, 6-month phase 3 trial with open-label extension (approximately 40 and 200 patients, respectively), intravenous administration of pegloticase (up to 8 mg every 2 weeks) induced profound initial reductions of serum urate [55,57]. In the pivotal phase 3 trial of pegloticase, which assessed a patient population with severe gout overall (and approximately 70% with visible tophi) [57], pre-infusion of fexofenadine, acetaminophen, and hydrocortisone (200 mg) were employed in an attempt to limit infusion reactions [57]. The frequency of responders - subjects who reached a target serum urate level of <6 mg/dL at 6 months - was approximately 42% on 8 mg pegloticase every 2 weeks

in the intent to treat analysis [57]. Moreover, de-bulking of tophi in this study was notably rapid in the subset of patients on pegloticase 8 mg every 2 weeks, with complete resolution of tophi in 20% by 13 weeks and approximately 40% by 25 weeks [56].

Frequent early acute gout flares (up to approximately 80%) in the first few months of pegloticase therapy [55] tapered off with more prolonged therapy in responders. Infusion reactions were moderate to severe in approximately 8 to 11% of subjects, and included flushing, urticaria, and hypotension, and, by undefined mechanisms, noncardiac chest pain or muscle cramping [55,57]. Anaphylaxis was uncommon (approximately 2%) in the phase 3 pegloticase study [57]. However, high titer antibodies to pegloticase emerged in many patients as treatment evolved over a few months, including IgM and IgG antibodies that did not directly neutralize the enzyme but appeared to adversely alter both its pharmacokinetics and pharmacodynamics [58]. High titer anti-pegloticase antibodies were also strongly linked with infusion reactions and were rare in serum urate responders (as assessed at the 6-month time-point) [58]. Hence, the dense polyethylene glycol (PEG) multimers linked to pegloticase [54] (Figure 5) do not prevent antigenicity, and also have been suggested to independently modulate the immune response to pegloticase in some subjects [58].

All uricase therapies have the potential to induce oxidative stress, since degradation of the high micromolar plasma concentrations of urate in gout patients by uricases has the capacity to generate substantial amounts of hydrogen peroxide [1,59,60]. Whether increased nitric oxide bioavailability [61,62] and the profound, rapid decrement in the serum antioxidant activity normally exerted by serum urate [1] contribute to oxidative challenge by uricase therapy is not yet clear. Circulating oxidative stress triggered by hydrogen peroxide generation alone is subject to marked dampening by the normal abundance of catalase on erythrocytes [51,59,60] and potentially by other plasma antioxidant defenses. Yet methemoglobinemia and/or hemolysis have been unequivocal indicators of uricase-induced oxidative stress [1,59,60]. Importantly, with Rasburicase™ therapy, methemoglobinemia and hemolysis (fortunately <1% in incidence) were linked to glucose-6-phosphate dehydrogenase deficiency in some but not all affected subjects [59,60]; subsequently, this deficiency has become an exclusion criterion for any uricase therapy. It has been suggested that assessment for erythrocyte catalase activity should be done prior to uricase therapy [59,60]. In my opinion, monitoring for treatment-induced subclinical methemoglobinemia also could ultimately be informative.

Uricases, by oxidizing urate (Figure 4), generate the intermediate form 5-hydroxyisourate, and subsequent hydrolysis of this produces 2-oxo-4-hydroxy-4-carboxy-5-ureidoimidazole, which is decarboxylated to S-(+)-allantoin [63]. The enzymes carrying out rapid degradation of these urate

oxidation intermediates were lost in human evolution along with uricase [63]. It has been suggested that addition of these enzymes to uricase therapy would be useful if the aforementioned uric acid oxidation intermediates are found to have noxious biologic properties [63].

Overall, it is not yet known if there is significant subclinical oxidative stress at the tissue level, rather than simply at the erythrocyte level [59,60], with uricase treatment in gout patients. Because of this issue, close monitoring of uricase-treated gout patients appears in order. Whether potential concomitant oxidative stress due to selected co-medications, congestive heart failure, anemia, hyperlipidemia, and CKD influences uricase safety remains to be defined.

At the time of writing this review, uricase therapy for tophaceous gout for which treatment has failed remains an unapproved, experimental approach that will be substantially more expensive than oral therapies, and consensus, evidence-based therapeutic guidelines are needed, whereas only draft guidelines have been proposed for uricase [52]. Tophus debulking is impressively rapid (months) in responders, but uricase therapies tested to date have all been substantially limited by drug immunogenicity. The safety of this particular 'biologic' approach, especially beyond a term of 6 to 12 months, will require further investigation.

In my opinion, any form of uricase therapy (over a finite term) is appropriate only for carefully selected patients that would benefit from accelerated, tophus de-bulking to address incapacitating tophi linked with active synovitis, and where other serum urate lowering therapies have failed or cannot achieve this objective [52]. As an 'induction therapy', uricase could ultimately be replaced by less intensive maintenance oral urate-lowering therapy with other agents, once evidence of normalization of body urate stores, including resolution of clinically detectable tophi and gross synovitis, is achieved.

Do cardiovascular safety signals in urate-lowering trials in gout reflect the influence of inflammation?

There have been death signals in both congestive heart failure patients on experimental oxypurinol therapy [64] and febuxostat-treated gout patients [48]. Moreover, there is higher cardiovascular mortality in hyperuricemia gout patients, related in part to co-morbidities in gout, and also possibly to independent effects of hyperuricemia on the vasculature [65,66].

Clinical trials to date in which death signals have arisen with serum urate-lowering strategies all have limitations in interpretability due to small numbers of events and subjects and relatively short durations of treatment. Hence, statistical significance may not be in lockstep with clinical and biological significance in such studies to this point. One constant associated with the more intense serum urate

lowering achievable in recent trials of emerging antihyperuricemics is increased risk for acute gout flares in the first few months of therapy [38,55]. In my opinion, the known association of atrial and ventricular arrhythmias and quantifiable altered heart rate variability with systemic inflammation (putatively mediated by specific cytokines markedly up-regulated in acute gouty inflammation, such as IL-6 and CXCL8) [67-69] deserves direct investigation as a potential factor in cardiovascular morbidity and mortality in gout patients undergoing serum-urate lowering therapy.

Challenges in translating novel gout and hyperuricemia therapies to better clinical practice

Compliance of gout patients with therapy appears lower than that for therapy of a variety of other common medical conditions, including hypertension, diabetes, osteoporosis, and hyperlipidemia [70]. Younger gout patients with fewer co-morbidities and fewer office visits are the least compliant gout patients, and we need to address systematic failures in both physician and patient education in gout treatment. Physicians appear to underestimate the impact of gout on quality of life and physical function [71-74]. Gout patients have more co-morbidities, poorer quality of life and physical function, increased health care costs, and increased adverse cardiovascular outcomes than controls [65,71-75].

Not only patient education, but also quality of care in gout treatment have significant room for improvement [76-78]. The identification of certain improved outcomes with sustained serum urate lowering below 6 mg/dL has ushered in a new era of gout therapy, where practitioners 'treat to target' in lowering serum urate [18]. Now the true definition of 'treatment-refractory' gout and gout-specific quality of life and disability will need careful assessment and direct attention in clinical practice. Such efforts would be timely, since 'treatment-refractory' gout, associated with an overall decrease in quality of life [79], has been proposed as a specific indication for aggressive urate-lowering strategies and possibly for initially lower serum urate targets than the widely used metric of <6 mg/dL [21].

The future of gout treatment is intriguing. For example, promising genomics and imaging technologies have the potential to improve prevention, diagnosis, and therapy by identifying disease earlier and tailoring treatment strategies. Examples include single nucleotide polymorphism and

haplotype identification for renal urate transporters in patients with hyperuricemia [80]. Dual energy computed tomography, which is highly sensitive and specific in visualizing tissue stores of monosodium urate crystals as well as renal uric acid urolithiasis [81,82], has the potential, for example, to assist in diagnosis of gout in patients with hyperuricemia or joint pain, and to better quantify tophus dissolution in therapy.

Whether using well-established or newer and emerging approaches and agents for gout and hyperuricemia management, the bottom line will remain that gout and hyperuricemia treatments need to be better translated into a collective of favorable outcomes for both control of gouty inflammation and management of hyperuricemia, as well as improved outcomes of gout-related quality of life and co-morbidities. This will require careful attention to both drug safety and cost-effectiveness of established versus emerging therapies, relative to quantifiable patient-centered outcomes, in a financially challenging era.

Competing interests

RT serves as consultant for Takeda, Savient, BioCryst, ARDEA, Altus, Novartis, Regeneron, Pfizer, URL Pharma, and UCB, and is also the past recipient of research grant support from Takeda.

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