The pathogenesis of gout

ABSTRACT
An elevated serum urate level, together with local factors, can result in the deposition of urate crystals into the joints. Once crystals are deposited into a joint, they can be released into the joint space and initiate an inflammatory cascade causing acute gouty arthritis. These acute flares resolve, but the crystals remain in the joint. The way to ultimately correct the underlying metabolic problem of hyperuricemia and the crystal deposition is to lower the serum urate level and dissolve the crystal deposits. This will stop both the acute attacks and the progressive joint damage.

KEY POINTS
A serum urate level of approximately 6.8 mg/dL is the concentration at which urate crystals begin to precipitate. The higher the urate level, the more likely that crystals will deposit into joints.

Crystal deposition and the development of gout
Asymptomatic hyperuricemia is not a disease but rather is the underlying factor that can predispose to gout. A serum urate level of approximately 6.8 mg/dL is the concentration at which monosodium urate crystals begin to precipitate.1,2 Although this level is based on in vitro studies, it suggests a reasonable biological threshold for clinicians assessing patients for hyperuricemia. It should be noted that there are often no manifestations of gout during an extended period of hyperuricemia even though urate crystals are beginning to deposit into joints. The higher the serum urate level, the more likely that crystals will deposit into joints.

Predisposition is not causation
In the Normative Aging Study, 22% of men who had serum urate levels greater than 9 mg/dL developed gout during a 5-year period—a much higher rate than among men with serum urate levels less than 9 mg/dL.3 Nevertheless, a full 78% of the men in this study with serum urate levels greater than 9 mg/dL did not develop gout over the 5-year period, illustrating that while hyperuricemia predisposes to gout, it does not automatically cause gout.

Contributing factors beyond serum urate
Other factors, when combined with hyperuricemia, contribute to crystal deposition and the development of gout. Trauma or irritation. Patients with hyperuricemia tend to have monosodium urate crystal deposition at these crystals into the joint space, where they may initiate an acute inflammatory reaction recognized as acute gouty arthritis. The acute attack is self-limited, but crystals remain in the joint and low-grade, often subclinical, inflammation persists even between acute attacks. Although acute attacks can be treated with anti-inflammatory medications, the underlying cause of the disease can be treated only by lowering the serum urate level.
sites of trauma or irritation. The first metatarsophalangeal joint is often affected, at least in part because it is a site of mechanical stress. Likewise, mechanical irritation from leaning on the elbow may cause crystals to deposit in the olecranon bursa.

Lower temperatures favor crystal deposition, which may explain why the helix of the ear and the foot are often sites of crystal deposition and tophus development. Both temperature and mechanical effects probably play a role in crystal deposition, however, as gouty attacks tend to occur at the first metatarsophalangeal joint, not at the interphalangeal joints of the foot, which are at a lower temperature.

Previous disease. Crystals also deposit with an increased incidence in previously diseased joints. The Heberden node is a good example. A patient with osteoarthritis in the fingers may experience dramatically increased pain and swelling because of a gout flare superimposed on an osteoarthritic joint.

■ ACUTE GOUTY ARTHRITIS

In some patients, the deposited monosodium urate crystals will be released into the joint space and cause the dramatic acute inflammatory response of acute gouty arthritis. Crystals are believed to be released either by some metabolic change, such as an increase or decrease in serum urate level, or by mechanical trauma. In the joint space, synovial lining cells appear to be the first to phagocytize the crystals. This sets into motion the formation of a complex called the inflammasome, which releases IL-1 beta, one of the most important mediators of the acute attack. It stimulates the release of chemokines, other cytokines, prostaglandins, and a variety of other proinflammatory molecules. The chemokines attract neutrophils into the synovial tissue and the synovial fluid. Neutrophil influx into the joint is a key feature of an acute attack of gout (Figure 1).

Gout flares may resolve spontaneously

Clinicians should be aware that gout attacks initially subside spontaneously. Because acute attacks of gout typically resolve with or without treatment, especially early in the course of the disease, it can be difficult to evaluate which treatments actually are effective against acute attacks.

A number of factors have been identified to explain how inflammation in acute attacks can be spontaneously suppressed. Crystals may dissolve or become sequestered in the tissue. Monocytes mature into macrophages, changing their responsiveness to urate crystals, and can begin to produce anti-inflammatory cytokines. In addition, some proteins that exude into the joint space with the attack, such as apolipoprotein B, can coat the crystals and reduce their inflammatory properties.

Crystals persist during intercritical periods

Following an acute attack, the symptoms of gouty arthritis may be gone, but the crystals are still present in the joint. Therefore, the patient remains at risk for continued flares and progressive disease. The crystals that remain in the joint are often associated with a low-grade persistent inflammation. It is not known why these crystals that remain in the joint fluid between attacks, some of which are phagocytized by white cells, do not initiate the whole cascade of inflammation. The reason may be related to the number of crystals present, their protein coating, or the nature of the resident synovial cells. Crystals may also persist as micro-tophi in the synovium (Figure 2).
key point is that low-grade inflammation persists and crystals remain in the joint, which can lead to progressive disease.14

■ ADVANCED GOUT

Clinicians treating patients with gout need to prevent the development of chronic, destructive arthritis and the overt, large tophaceous deposits of advanced gout. Over time, even in the absence of flares, deposited crystals and inflammation can lead to the development of clinically evident joint damage and erosions that can be seen on radiographs (Figure 3) or magnetic resonance imaging.15

■ INTERVENTIONS MUST NORMALIZE URATE LEVEL

Acute gout attacks can be treated with anti-inflammatory drugs, but the disease can and often will continue to progress unless the serum urate level is normalized. Two studies of patients whose serum urate levels were successfully reduced to less than 6 mg/dL showed that crystals began to be depleted from the patients’ joint fluid, which should ultimately prevent the risk of progressive gouty arthritis.12,16 Perez-Ruiz and colleagues have shown that tophi can be dissolved by decreasing the serum urate level.17 When tophi are present, aiming for even lower levels of serum urate, such as 4 to 5 mg/dL, may help to promote more rapid dissolution of crystals.17

REFERENCES


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FIGURE 3. Large, cystic joint erosions producing an overhanging edge (circled area) characteristic of chronic gouty arthritis.