Update on GOUT
A Practical Approach......

Introduction

Gout is an ancient disease which dates back to the time of Babylon. Its name comes from the Latin word gutta, (meaning drop), which reflects an old belief that the disease is caused by a poison which falls into the joint drop by drop.

The disease is quite prevalent nowadays and is considered as the most common inflammatory type of arthritis in men affecting 1-2% of adults in Western countries, with male to female ratio of 3.6:1. It is rare in pre-menopausal women and its incidence and prevalence increases with age.

Gout is an inflammatory syndrome caused as a response to monosodium urate monohydrate crystals

(MSUM) formed in humans in the presence of having elevated serum urate concentration what is known as hyperuricemia. Hyperuricemia is defined as serum urate levels above 6.8mg/dl (≥ 400µmol/L) which is the level above which the physiological saturation threshold is exceeded.

Why there is increase in prevalence of GOUT?

The current increase in the prevalence of gout is related to overweight, of metabolic syndrome & change in our diet with high intake of meat, seafood, fructose sweetened beverages and beer, in addition to the increase in life expectancy. But, the main reason is related to the renal uric acid hypo-excretion, due to genetic and environmental factors such as diuretic use, low dose of aspirin and high alcohol consumption.

Stages of Gout

a. Asymptomatic hyperuricemia with elevated uric acid but no clinical gout.
b. Acute gout with acute inflammation and intervals between flares.
c. Advanced chronic gout with long-term gouty complications of uncontrolled hyperuricemia.

Serum uric acid is determined by

a. Dietary purine intake.
b. Cellular degradation of purines to urate
c. Urate excretion from body via intestines and kidneys.

Urate saturation above 6.8 mg/dl can lead to monosodium urate crystals

a. Induce painful inflammatory responses.
b. Grow into tophi.
c. Precipitate erosive joint damage over time (Table).

Diagnosis of gout

Score of 8 or more of ACR/EULAR criteria for classification of gout which consists of:

a. Clinical score
b. Laboratory: serum urates level score
c. Imaging score

Joint aspiration and synovial fluid analysis, gold standard compensated polarized light microscopy

Imaging urate crystal deposits and tophaceous gout: The crystals deposits can be clearly identified by the use of the dual energy CT imaging or by ultrasound which show double contours punctiform deposits in synovial membrane

a. Plain x-ray: insensitive in early disease
b. Ultrasound :sensitive in early disease but can be abnormal in asymptomatic hyperuricemia
c. CT Scan: sensitive but expensive
d. MRI :sensitive but expensive

Common sites of gout:

a. Olecranon bursa 
b. Elbow 
c. Wrist 
d. Fingers 
e. Knee 
f. Ankle 
g. Subtalar 
h. Mid-foot 
i. First Metatarsophalangeal joint 50% of initial attack & affecting 90% of patients

Table 1 2015 ACR/EULAR Criteria for Classification of Gout I

<table>
<thead>
<tr>
<th>Clinical</th>
<th>Score</th>
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<tbody>
<tr>
<td>Ankle or midfoot</td>
<td>1</td>
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<tr>
<td>First MTP joint</td>
<td>1</td>
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<tr>
<td>Characteristic episodes erythema, ulceration, unbearable to touch, difficulty walking</td>
<td>+1 up to+3</td>
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<tr>
<td>Episode time frame</td>
<td>One typical +1, Recurrent +2</td>
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Update on GOUT

Clinical Score
Tophus present 4
Laboratory Serum urate less than 4:-4, 6-8 mg/100ml:+3, Equal or more 10mg/dl:+4
Synovial fluid analysis MSU negative: -2
Imaging Presence of gout-related joint damage: +4

Differential diagnosis
a. Septic arthritis
b. Pseudogout
c. Cellulitis
d. Osteoarthritis first MTP joint or nodal osteoarthritis
e. Rheumatoid arthritis
f. Psoriatic arthritis, spondylarthropathy
g. Lyme disease

Co morbidity checklist for primary care
a. Obesity and diet including excessive alcohol intake.
b. Modifiable risk factors: hyperlipidemia, hypertension, metabolic syndrome, Type 2 diabetes mellitus.
c. Serum urate-elevating medications
d. History of urolithiasis
e. Chronic kidney disease
f. Genetic cause of uric acid overproduction
g. Acquired cause of uric acid overproduction: psoriasis, myeloproliferative or lymphoproliferative disease
h. Lead intoxication

2012 ACR Gout guidelines

Pharmacologic therapy for the acute flares:
a. Supplement with topical ice as needed
b. Initiate therapy within 24h of acute flare onset
c. Continue pharmacologic rate-lowering therapy during attacks

Mild / moderate pain 1-3 small joints or 1-2 large joints: Monotherapy:
a. NSAID: Cox 2 inhibitor has the same efficacy as conventional non-steroidal anti-inflammatory
drugs
b. Systemic corticosteroids : prednisolone greater or equal 0.5 mg/kg/day for 5-10 days
d. Colchicines: 1.2 mg initially and 0.6 mg 1 hour later

Polyparticular or multiple large joint with severe pain: Combination therapy:
a. Colchicine + NSAID
b. oral corticosteroid + colchicine
c. Intra-articular steroids with all other modalities

Patient education after successful outcome:

Diet and lifestyle triggers: avoid organ meats, high fructose corn syrup-sweetened soda and excessive alcohol (more than two drinks per day for men and more than one drink per day for women). Limit beef, lamb, high purine seafood (shellfish) & beer.
a. Prompt self-treatment of subsequent attacks
b. Urate-lowering therapy (Figure)

Indications for pharmacologic urate-lowering therapy:
a. Tophus or Tophi
b. Frequent attacks equal or greater than 2 attacks per year
c. Chronic kidney disease: stage 2 or worse
d. History of urolithiasis

Current recommendations for urate-lowering Therapy:
a. Usual serum urate target is less than 6 mg/dl
b. Serum urate levels less than 5 mg/dl may be needed to improve gout signs and symptoms.

Select first-line agent
a. Xanthine Oxidase inhibitor : Allopurinol or Febuxostat.
b. Probenecid: If xanthine oxidase inhibitor is contraindicated or not tolerated.

Acute Gout Prophylaxis

Allopurinol as first-line urate lowering therapy:
a. Effective in uric acid overproducers and underexcretors.
b. Starting dosage should be no greater than 100 mg/day for any patient, and start at 50 mg/day in stage 4 or worse CKD (evidence B). Gradually titrate maintenance dose upward every 2–5 weeks to appropriate maximum dose in order to treat to chosen SUA target (evidence C).
c. Dose can be raised above 300 mg daily, even with renal impairment, as long as it is accompanied by
   adequate patient education and monitoring for drug toxicity (e.g. pruritis, rash, elevated hepatic transaminases; evidence B).
f. Maximum approved dose 800mg/day (reduced in chronic kidney disease, CKD).
g. Intolerance in 5-10%, pruritic rash 2% & major allopurinol hypersensitivity syndrome 0.1-0.4%

Citation: El-Labban O. Update on GOUT. Biom Biostat Int J. 2016;4(3):113–116. DOI: 10.15406/bbij.2016.04.00098
Figure 1 Specific recommendations: general health, and life style measures for GOUT patients#

Urate lowering therapy: Febuxostat
a. Selective inhibitor of xanthine oxidase with nonpurine backbone.
b. The dose is 40 mg/day starting dose. Labeled use up to 80 mg/d (ACR guideline allow for doses up to 120 mg/d.

Advantages: more selective than allopurinol. There is lower renal excretion vs allopurinol active metabolite. No dose adjustment with mild / moderate renal or hepatic impairment.

Urate-lowering Therapy: Uricosurics Increase urate excretion by inhibiting urate reabsorption in the kidney.

Probenecid:

a. Alternative in those patients intolerant to at least 1 xanthine oxidase inhibitor.
b. Not recommended in persons with history of urolithiasis as it increases urolithiathis risk especially with acidic urine PH.

c. Not recommended if creatinine clearance less than 50ml/min.

Losartan, atorvastatin and fenofibrate are less potent: The use of losartan and fenofibrate alone or in combination with urate lowering therapies can have an extra value in the prevention and management of hyperuricaemia in patients who have both hypertension and dyslipidaemia.

Emerging options in development
a. Lesinurad: Selective uric acid reabsorption inhibitor.
b. Arhalofenate: Dual-acting anti-inflammatory and urate-lowering therapy.

Gout flare prophylaxis
Initiate with or just before initiating pharmacologic urate-lowering therapy:
a. First line: Low dose colchicine (0.5 or 0.6 mg once or twice
daily) or Low-dose NSAID (with proton pump inhibitor where indicated).

b. **Second-line**: low-dose prednisone or predinsolone (equal or less 10 mg/day).

**If Gout signs / symptoms appear**: evaluate gout symptoms with urate-lowering therapy

**If No gout signs symptoms appear**: Treat with Longest period among the following:

a. At least 6 months.

b. 3 month after achieving target serum urate level with no tophi.

c. 6 months after achieving target serum urate level with greater or equal 1 tophus.

**Long-term Management**

a. Treat to Target serum urate level no higher than less 6 mg/dl.

b. If serum urate level achieved then continue gout attack prophylaxis. Regularly monitor serum urate.

c. if serum level is not achieved, increase intensity of lowering therapy and reevaluate serum urate.

d. Diagnose, treat and prevent acute gout flare.

e. identify and manage comorbidities and causes of hyperuricemia.

**Take home messages**

a. Aspirate joint if diagnosis is unclear or septic arthritis is possible

b. Use ultrasound or advanced imaging particularly in early disease.

c. Initiate pharmacologic urate lowering therapy in all patients with Tophi, multiple attacks/year, CKD stage 2, or higher, or previous urolithiasis

d. Prescribe prophylactic anti-inflammatory therapy for at least 6 months hen initiating a pharmacologic urate lowering regimen

e. Assess adherence of patients to prescribed regimen.

**Acknowledgement**

None.

**Conflict of interest**

None.