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**New Indication** 

# canakinumab for gout attacks

Too risky

• There are no clinical trials of *canakinumab* in patients with gout attacks refractory to several prior treatments. Adverse effects mainly include infections, hypersensitivity reactions and dizziness.



Gout attacks are due to accumulation of uric acid crystals in the joints (1). If left untreated, the associ-

ated inflammation and pain subside after a few days. When ice, *paracetamol*, nonsteroidal anti-inflammatory drugs (NSAIDs) such as *ibuprofen*, and *colchicine* are not sufficiently effective or cannot be used, systemic corticosteroids are another option (1). Adding an opioid analgesic can help to relieve severe pain that is unresponsive to other treatments (2).

*Canakinumab* (Ilaris°, Novartis Pharma) is a monoclonal antibody targeting interleukin-1 beta, a cytokine involved in immune and inflammatory processes. After having been authorised in the European Union for serious cryopyrinassociated periodic syndrome, *canakinumab* is also approved for the treatment of gout when NSAIDs and *colchicine* are ineffective and corticosteroids are inappropriate (**a**)(3).

### canakinumab powder for

solution for SC injection

#### **LARIS**°

• **150 mg** canakinumab per vial

#### immunosuppressive drug; interleukin-1 beta antagonist

■ New indication: "(...) symptomatic treatment of adult patients with frequent gouty arthritis attacks (at least 3 attacks in the previous 12 months) in whom non-steroidal anti-inflammatory drugs (NSAIDs) and colchicine are contraindicated, are not tolerated, or do not provide an adequate response, and in whom repeated courses of corticosteroids are not appropriate (...)". [EU marketing authorisation, centralised procedure]

Little evaluation after multiple treatment failure. Clinical evaluation of canakinumab in this setting is based on two randomised, double-blind, doubledummy trials comparing subcutaneous canakinumab (150 mg) with intramuscular triamcinolone (40 mg), a longacting corticosteroid, in 454 patients in whom NSAIDs and/or colchicine had failed (2-5). Both NSAIDs and colchicine were ineffective, not tolerated or contraindicated in only about one-third of patients. Patients had experienced at least three gout attacks in the year prior to enrolment, and 42% were taking a preventive treatment (mainly allopurinol).

At baseline, pain intensity in the most painful joint averaged 74 mm on a visual analogue scale ranging from 0 mm to 100 mm. In a pooled analysis of the two trials, 72 hours after the injection, pain intensity was 25 mm with canakinumab versus 36 mm with triamcinolone (p < 0.0001) (2-5). The clinical relevance of this difference is uncertain. Full pain relief was achieved in respectively 46% and 37% of patients in the canakinumab and triamcinolone groups, a non-significant difference. Recourse to paracetamol, codeine or corticosteroids was less frequent with canakinumab than with triamcinolone (37% versus 55%, p < 0.0001).

Infections, hypersensitivity reactions, dizziness. The immunosuppressive effect of *canakinumab*, and its long elimination half-life (nearly a month), create a risk of infections and possibly cancer (6).

In the two trials, 19% of patients receiving canakinumab developed an infection, versus 13% of patients receiving triamcinolone. The infections mainly included nasopharyngitis, upper respiratory tract infections, urinary tract infections, and bronchitis (2,3). Four patients receiving canakinumab and none treated with triamcinolone developed serious infections, while respectively 3.2% and 2.4% of patients developed opportunistic infections. It is also noteworthy that two patients treated with canakinumab, one for juvenile idiopathic arthritis and one for periodic syndrome, died of septic shock and disseminated tuberculosis (3).

Overall, 63% of patients treated with *canakinumab* versus 51% of those receiving *triamcinolone* experienced at least one adverse event (severe in respectively 7% and 3% of patients) (2, 3). The most common adverse effects associated with *canakinumab* were hypersensitivity reactions (including angioedema), dizziness, haematological disorders (especially neutropenia), and hepatic disorders (2,3). Elevation of serum uric acid levels was more frequent with *canakinumab* than with *triamcinolone*.

The *canakinumab* risk management plan calls for special monitoring of infections and malignancies.

In practice. Given its uncertain efficacy on gout-related pain, *canakinumab* is too risky to use for treatment of gout attacks. Its price is also exorbitant. It is better to stick with existing symptomatic treatments, which should be initiated as early as possible.

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*a*- Scientists consulted by the US Food and Drug Administration (FDA) voted unanimously against authorising canakinumab for the treatment of gout (ref 4).

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## Selected references from Prescrire's literature search

In response to our request for information, Novartis Pharma provided us with no documentation on its product.

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4- US FDA "Ilaris (canakinumab)-Summary Minutes and Transcript of the Arthritis Advisory Committee Meeting" 21 June 2011: 347 pages. 5- Schlesinger N et al. "Canakinumab for acute

**5-** Schlesinger N et al. "Canakinumab for acute gouty arthritis in patients with limited treatment options: results from two randomised, multicentre, active-controlled, double-blind trials and their initial extensions" *Ann Rheum Dis* 2012; **71**: 1839-1848 + supplementary appendix (29 pages).

**6-** Prescrire Rédaction "canakinumab-llaris". Pour quelques formes graves de syndrome périodique associé à la cryopyrine" *Rev Prescrire* 2010; **30** (324): 726-729.

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