Anti-inflammatory therapy in ischaemic heart disease: from canakinumab to colchicine

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Four large trials have recently evaluated the effects of anti-inflammatory drugs in the secondary prevention of major cardiovascular events (MACE) in over 25 000 patients followed for 1.9–3.7 years. CANTOS tested subcutaneous canakinumab [an anti-interleukin (IL) 1β antibody] 300 mg every 3 months against placebo in patients with a history of myocardial infarction (MI) and serum C-reactive protein (CRP) >2 mg/L, demonstrating efficacy in preventing MACE but increased rates of fatal infections. COLCOT (in patients with recent MI) and LoDoCo2 (in patients with chronic coronary syndromes) tested oral colchicine (an NLRP3 inflammasome inhibitor) 0.5 mg daily vs. placebo, demonstrating prevention of MACE with a slightly increased risk of pneumonia in COLCOT (0.9 vs. 0.4%) but not in LoDoCo2. CIRT tested oral methotrexate (an anti-rheumatic anti-nuclear factor-kB) 15-20 mg per week against placebo in ischaemic heart disease patients with diabetes or metabolic syndrome, without significant reduction in MACE rates or in circulating IL6 or CRP levels, and with increased risk of skin cancers. In summary, canakinumab and colchicine have shown efficacy in preventing MACE in ischaemic heart disease patients, but only colchicine has acceptable safety (and cost) for use in secondary cardiovascular prevention. Clinical results are expected with the anti-IL6 ziltivekimab.

Introduction

Four large trials conducted in recent years have tested the hypothesis that anti-inflammatory drugs, such as canakinumab and colchicine, can reduce the incidence of major cardiovascular events (MACE) in patients with previous myocardial infarction (MI) or chronic coronary syndromes. To optimize the understanding of these studies we briefly illustrate the mechanisms of action of the tested drugs, the multiple aspects that relate inflammation to ischaemic heart disease, and some preliminary results with aspirin and ziltivekimab; the main findings of the large placebo-controlled trials with canakinumab, colchicine, and methotrexate are presented and their clinical implications briefly discussed.

Anti-inflammatory drugs and ischaemic heart disease

The anti-inflammatory effects of low-dose aspirin (100-300 mg/day), although less well known than its antiplatelet effects, are documented in various clinical settings and may contribute to aspirin’s benefits in the secondary prevention of MACE; the exact molecular mechanisms of such effects are under investigation. Canakinumab is a recombinant human monoclonal G1k immunoglobulin against
interleukin (IL) 1β that neutralizes the signals induced by IL1β on lymphoid, myeloid, endothelial, and other cell types; canakinumab administration reduces circulating IL6, fibrinogen, and C-reactive protein (CRP) levels compared to placebo.1,5 Colchicine is a plant alkaloid that inhibits tubulin polymerization and the nod-like receptor pyrin domain containing protein-3 (NLRP3) inflammasome within monocytes and other cell types; in vitro, colchicine is reported to reduce platelet aggregation, superoxide production, neutrophil recruitment and adhesion, mast cell degranulation, monocyte chemotaxis, endothelial pyroptosis by cholesterol crystals, and endothelial activation by oxidized LDL cholesterol1; in rats, colchicine has been found to inhibit hepatic secretion of fibrinogen causing fibrinogen accumulation within the endoplasmic reticulum and Golgi apparatus6; rabbits subjected to hypercholesterolaemic diets treated with colchicine show reduced atherosclerotic development and reduced circulating fibrinogen levels2; in rodents, colchicine was found to protect against acute cerebral ischaemia by inhibiting cell chemotaxis and exocytosis.1 Methotrexate—used to treat rheumatic diseases and cancer—has multiple effects including inhibition of the nuclear transcription factor-kB, of DNA/RNA synthesis and of dihydrofolate reductase1; in patients with previous MI, methotrexate did not reduce circulating IL6 levels.5 Ziltivekimab is an anti-IL6 human monoclonal antibody; unlike antibodies directed against the IL6 receptor, ziltivekimab may act at lower concentrations causing fewer adverse events.5 The actions of the aforementioned drugs are schematically illustrated in Figure 1.

### Inflammation and ischaemic heart disease

The close links between inflammation, atherothrombosis, and coronary syndromes have long been known, stemming from pathological, mechanistic, and clinical data (Table 1). Atherosclerosis itself has been defined as a chronic vascular inflammatory process.10 Unstable plaques have specific characteristics, with an increased share of macrophages and neutrophils compared to stable plaques.11 Coronary thrombi contain leucocytes, in addition to platelets, fibrin, and red cells.12 Various prothrombotic factors, such as fibrinogen (the precursor of fibrin and bridging molecule among aggregated platelets), plasminogen activator inhibitor-1 (rapid inhibitor of endogenous fibrinolysis and profibrotic factor), and von Willebrand factor (platelet adhesion molecule), are acute phase proteins.13,14 As stated, in animal models, colchicine can slow the development of atherosclerosis and hepatic fibrinogen secretion.5,6 In humans, inhibition of IL1β or IL6 reduces circulating levels not only of other inflammatory cytokines and CRP but also of prothrombotic factors such as fibrinogen.5,9 The main traditional cardiovascular risk factors (hypertension, diabetes, dyslipidaemia, smoking, obesity, sedentary lifestyle) are associated with systemic low-grade inflammation: for each additional factor, an approximate increase of 1 mg/L of circulating CRP is observed.15 Subclinical low-grade inflammation increases the risk of MACE among healthy subjects: elevated fibrinogen concentrations16 and white blood cell counts,7 even within the normal range, are predictors of MACE. Common inflammatory markers predict the risk of MACE in patients with both acute18 and chronic9 coronary syndromes. In patients with chronic coronary syndromes, low-dose aspirin reduces the circulating levels of inflammatory biomarkers and the incidence of inducible ischaemia.2 Finally, acute MI stimulates an acute-phase and prothrombotic response, with peak values and duration of response directly proportional to the extent of MI.12,13 Based on this extensive collection of data, phase II studies and recent large-scale trials have evaluated the effects of anti-
Table 1: Multiple interrelations between inflammation and ischaemic heart disease

**Histopathology**
- Atherosclerosis is a chronic vascular inflammatory process.
- Unstable plaques contain a higher percentage of macrophages and neutrophils than stable plaques.
- Arterial thrombi contain platelets, fibrin, erythrocytes, and leucocytes.

**Molecular and experimental data**
- Prothrombotic factors (fibrinogen, PAI-1, and VWF) are acute phase proteins.
- Inhibition of IL1β or IL6 reduces circulating fibrinogen levels in humans.
- Colchicine slows the development of atherosclerosis and hepatic secretion of fibrinogen in experimental models.

**Pathophysiology and prognostic data**
- Traditional CV risk factors induce low grade systemic inflammation.
- Low grade inflammation increases the risk of major CV events.
- Inflammatory biomarkers predict the risk of CV events in patients with acute or chronic coronary syndromes.
- Aspirin reduces the levels of inflammatory biomarkers and the risk of major CV events.
- Acute MI stimulates an acute phase response in proportion to the extent of necrosis.

**Large randomized trials**
- Canakinumab prevents CV events in patients with previous MI and elevated CRP but increases the risk of fatal infections.
- Colchicine prevents CV events in patients with acute or chronic coronary syndromes without excess of adverse events.

**Inflammatory drugs for the secondary prevention of atherothrombotic events.**

**Selected phase II studies**

**RESCEU** (Trial to Evaluate Reduction in Inflammation in Patients With Advanced Chronic Renal Disease Utilizing Antibody-Mediated IL-6 Inhibition) is a recent, double-blind, randomized US study that tested increasing doses of the recombinant antibody antibody (CIRT8) and low-dose colchicine (COLCOT and LoDoCo2).

**CIRT:** Cardiovascular Inflammation Reduction Trial

CIRT is a double blind, randomized study sponsored by the National Institutes of Health. It compared low-dose methotrexate (target 15-30 mg once weekly) vs. placebo in 4786 patients with previous MI or multivessel epicardial artery disease plus diabetes or metabolic syndrome followed for a median of 2.3 years. The primary endpoint was the classical triple composite of MACE, i.e. non-fatal MI, non-fatal stroke or cardiovascular death; subsequently, for a lower than expected event rate, hospitalization for revascularization from unstable angina was added. The results were
disappointing: low-dose methotrexate did not reduce circulating levels of IL1β, IL6, or CRP, nor cardiovascular events vs. placebo (201 vs. 207; hazard ratio 0.96, CI 0.79–1.16). With methotrexate vs. placebo, there was an increased incidence of oral lesions, leucopenia, unwanted weight loss, transaminase elevation, and cancer (mostly non-basal cell skin cancers; 52 vs. 30, \( P = 0.02 \)). For these reasons, the trial was terminated prematurely.

**COLCOT: COLchicine Cardiovascular Outcome Trial**

COLCOT\(^2\) is an independent, multinational study funded by the Canadian government. It randomized double-blindly 4745 patients with recent MI (within 30 days), regardless of CRP values, to colchicine (0.5 mg daily) or placebo. Patients were treated with optimal medical therapy and followed for a median of 2.3 years. The primary endpoint was the combination of cardiovascular death, resuscitated cardiac arrest, non-fatal MI, non-fatal stroke, and revascularization for unstable angina. The primary endpoint was documented in 5.5% of patients treated with colchicine and in 7.1% treated with placebo, a significant relative reduction of 23% (hazard ratio 0.77, CI 0.61–0.96; \( P = 0.02 \)). Treatment with colchicine had favourable effects on each individual component of the composite primary endpoint. All-cause mortality was not different in the two arms (43 deaths with colchicine vs. 44 with placebo). A higher rate of pneumonia was observed with colchicine vs. placebo, although with a low incidence (0.9% vs. 0.5%, \( P = 0.03 \)), while the frequency of diarrhoea (9.7% vs. 8.9%) did not differ significantly in the two arms. Study limitations include the relatively short follow-up and the relatively small sample size, providing reliable answers for the entire population but not for specific subgroups.

COLCOT has confirmed the crucial role of inflammation in the progression of ischaemic heart disease by providing an effective, low cost, reasonably safe preventive treatment with a drug already known to the medical community for the treatment of gout, familial Mediterranean fever and pericarditis. The COLCOT results, however, cannot be generalized to all ischaemic heart disease patients, being limited to patients with recent MI, in whom the intensity of inflammation may be more relevant than in stable patients with either an old MI or no previous history of acute coronary syndromes.

**LoDoCo2: Low-Dose Colchicine 2 Trial**

LoDoCo2\(^2\) is a double blind, randomized study that tested low-dose colchicine (0.5 mg daily) vs. placebo in 5500 patients with documented obstructive epicardial artery disease, stable for at least 6 months, followed for a median of 2.4 years. It was conducted in Australia and the Netherlands, and funded by public and private foundations and a consortium of pharmaceutical companies. The primary endpoint was a composite of cardiovascular death, spontaneous (non-procedural) MI, ischaemic stroke, and ischaemia-driven coronary revascularization. The main secondary endpoint was the classical triple composite of cardiovascular death, MI, or stroke. The results were, once again, favourable. The primary endpoint occurred in 6.8% of patients treated with colchicine and in 9.6% of patients treated with placebo, representing a highly significant relative reduction of 31% (hazard ratio 0.69, CI 0.57–0.83; \( P < 0.001 \)). The primary secondary endpoint showed a
relative reduction of 28% (absolute rate 4.2% with colchicine vs. 5.7% with placebo; hazard ratio 0.72, CI 0.57–0.92; P = 0.007), i.e. a significant reduction not only from a statistical but also from a clinical standpoint. Regarding safety, there were no significant differences in the rates of expected adverse events, such as pneumonia or gastrointestinal disturbances, between treatment arms. The incidence of non-cardiovascular death was numerically (but not statistically) higher with colchicine vs. placebo (0.7 vs. 0.5 events per 100 person-years; hazard ratio 1.51, CI 0.99–2.31). Subgroup analyses showed homogeneous effects in all analysed subgroups. An important study limitation was the lack of information on circulating levels of inflammatory indices before and after treatment.

Thus, LoDoCo2 and COLCOT appear to close the circle, confirming that inflammation is a determinant of ischaemic heart disease progression and atherothrombosis, and that anti-inflammatory drugs can prevent MACES in patients with recent MI (COLCOT) or chronic coronary syndrome (LoDoCo2).

Conclusions and perspectives

Recent large-scale placebo-controlled trials have confirmed the role of inflammation in the pathogenesis of atherothrombotic events by demonstrating that specific anti-inflammatory drugs are able to prevent MACE in patients with ischaemic heart disease. As a result, the therapeutic options for secondary prevention of cardiovascular events are expanding. CANTOS demonstrated a significant involvement of IL1β in atherothrombosis, and the colchicine trials have confirmed the benefits of anti-inflammatory therapy in patients with recent MI or chronic coronary syndromes using a well-known, reasonably safe and economical treatment. Inflammation can therefore be added to the three traditional therapeutic targets of atherothrombotic diseases (thrombosis, dyslipidaemia, neuroendocrine activation) (Figure 2). Forthcoming international guidelines will very likely provide indications on the use of colchicine for the secondary prevention of MACE. The addition of a new drug to existing ones will trigger the acute stress-induced increase in circulating levels of interleukin-6: a randomized, double-blind, placebo-controlled study. Brain Behav Immun 2008; 22:150–157.

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