Targeting inflammation in atherosclerosis — from experimental insights to the clinic

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Abstract | Atherosclerosis, a dominant and growing cause of death and disability worldwide, involves inflammation from its inception to the emergence of complications. Targeting inflammatory pathways could therefore provide a promising new avenue to prevent and treat atherosclerosis. Indeed, clinical studies have now demonstrated unequivocally that modulation of inflammation can forestall the clinical complications of atherosclerosis. This progress pinpoints the need for preclinical investigations to refine strategies for combatting inflammation in the human disease. In this Review, we consider a gamut of attractive possibilities for modifying inflammation in atherosclerosis, including targeting pivotal inflammatory pathways such as the inflammasomes, inhibiting cytokines, manipulating adaptive immunity and promoting pro-resolution mechanisms. Along with lifestyle measures, pharmacological interventions to mute inflammation could complement traditional targets, such as lipids and hypertension, to make new inroads into the management of atherosclerotic risk.

Atherosclerosis now comprises the major contributor to a global epidemic of cardiovascular disease, which has overtaken communicable diseases to become the leading cause of death and disability worldwide1. The spread of obesity and attendant diabetes has aggravated the risk of atherosclerosis both in high-income regions and, alarmingly, in the developing world as well. Atherosclerosis is not only the main underlying cause in coronary heart disease but also causes many strokes and affects peripheral arteries. This lipid-driven disease arises from the accumulation of low-density lipoprotein (LDL) and remnant lipoprotein particles in focal areas of medium and large arteries. Predilection sites for lipid entry and retention in the subendothelial space localize in regions of disturbed, non-laminar flow at arterial branch points. Although lipids undoubtedly contribute causally to atherosclerosis, the ensuing inflammatory response orchestrates the progression and outcome of the disease.

Until recently, after lifestyle measures, non-invasive therapeutic intervention for atherosclerosis focused primarily on pharmacologically limiting risk factors such as arterial hypertension and hypercholesterolaemia. However, inflammation provides a set of pathways that link traditional risk factors to the altered behaviour of cells of the artery wall and the recruitment of leukocytes, mechanisms that promote the disease and its complications. The recent Canakinumab Anti-inflammatory Thrombosis Outcomes Study (CANTOS) moved targeting inflammation in atherosclerosis from conjecture to clinical reality2. This study targeted a specific pro-inflammatory cytokine, IL-1β, which has been implicated in atherogenesis through decades of experimental work. CANTOS showed that administration of an anti-IL-1β antibody to patients who were cardiovascularly stable after a myocardial infarction and were being treated according to current guidelines (including with statins) reduced recurrent major adverse cardiovascular events. Subsequent clinical trials with colchicine, another anti-inflammatory agent, have likewise demonstrated clinical benefit in patients with recent or temporally remote acute coronary syndromes (ACS)3,4.

This Review summarizes our current understanding of inflammatory processes and cellular participants in atherosclerosis. We then discuss how current clinical trials have started to target some of these processes and which experimental strategies could quell inflammation in the clinic. We also discuss means to promote inflammation resolution. However, both inhibiting inflammation and stimulating its resolution may cause unwanted effects. Hence, we elaborate on means to limit adverse effects, including chronopharmacology and targeted drug delivery with nanoparticles. Beyond pharmacological intervention, we also consider lifestyle changes and how their beneficial effects may arise, in part, from alleviating arterial inflammation. Taming inflammation in atherosclerosis could substantially benefit human health, adding to established targets such as LDL and hypertension, to stem the growing global burden of cardiovascular disease.
Key cells in arterial inflammation

Atherosclerotic lesions evolve from an orderly structured arterial wall to form a complex conglomerate of resident and newly recruited cells. Plaques, also known as atheromas, form in the inner layer of arteries (the tunica intima). Mature atheromas typically contain a lipid-rich central core underneath a fibrous cap (Fig. 1). A monolayer of endothelial cells rests on the surface of plaques, providing the crucial interface with the blood. In this section, we discuss three key cell types that contribute to arterial inflammation: smooth muscle cells (SMCs), macrophages and neutrophils. For other cell types, including endothelial cells and cells of the lymphoid lineage, we refer the reader to other excellent review articles.\(^1,2\)

SMCs: more than just stability?

The majority of cardiovascular complications result from plaque rupture, an event whose likelihood correlates inversely with lesional SMC content. Hence, the prevailing view of SMCs in advanced atherosclerosis posits that these cells exert predominantly beneficial roles as they furnish the stabilizing fibrous cap and generate the extracellular matrix. However, the use of fate-mapping and lineage-tracing approaches, in combination with single-cell RNA sequencing, have unveiled a much more diverse picture of the SMC population throughout the various stages of atherosclerosis\(^3\) (BOX 1). At early stages of atherosclerosis, lipids accumulate in the subendothelial region, where they become trapped as a consequence of the interaction between positively charged apolipoproteins and negatively charged proteoglycan side chains and components of the extracellular matrix produced by SMCs\(^4\). SMCs exposed to modified lipids or pro-inflammatory cytokines then release chemoattractants, including C-C motif chemokine 2 (CCL2) and CCL5 (REFS\(^5,6,11\)) which promote the recruitment of monocytes\(^7\) (Fig. 1a).

During lesion evolution, macrophages and SMCs die and contribute to the formation of the necrotic core (Fig. 1b). This highly inflammatory milieu evokes a healing response that initially fosters formation of the plaque’s protective fibrous cap. This structure contains abundant extracellular matrix and SMCs that contain α-smooth muscle actin (αSMA), which is a marker of contractile SMCs. In mouse models of atherosclerosis, fibrous cap SMCs arise from the tunica media\(^11\). These cells migrate towards the fibrous cap in a process that depends on the transcription factor OCT4 (also known as POU5F1)\(^8\), where they undergo clonal expansion\(^9\). Somatic mutations drive clonal expansion in non-malignant tissues during ageing, and clonal expansion in myeloid cells has been linked to an increased risk for cardiovascular events\(^10\). Similar mechanisms may occur in SMCs in atherosclerotic lesions. Indeed, early observations made in humans showed that lesional SMCs are monotypic\(^12\).

Once in the fibrous cap, SMCs are major products of the extracellular matrix. Although definitive evidence is missing, data from mice lacking COL15A1, which encodes a collagen chain, specifically in SMCs\(^13,14\) and secretome proteomics of lipid-laden SMCs\(^15\) firmly suggest that SMCs contribute to fibrous cap formation. However, lipid uptake and changes in the interactions between SMCs and the extracellular matrix alter the SMC phenotype, increasing the expression of markers typically ascribed to macrophages\(^16,17\). In advanced atherosclerotic plaques in mice, 30–70% of cells with macrophage markers\(^16,18\) (also known as foam cells) are SMCs; similar observations have been made in human atherosclerotic plaques\(^12,14\). The detrimental consequences of this phenotypic switch towards macrophage-like SMCs derives support from a study in which the transcription factor Krueppel-like factor 4 (KLF4), a transcription factor earlier reported to control the phenotypic transition of SMCs during development\(^19\), was deleted specifically in SMCs. These mice show reduced SMC switching, a clear-cut increase in fibrous cap thickness and increased αSMA+ cell content of the fibrous cap\(^20\).

The SMC transition to macrophage-like cells is likely pro-inflammatory and contributes to plaque vulnerability, but SMCs can also transition to fibroblast cells, which have previously been thought to have the opposite effects. A recent study combining single-cell RNA sequencing and fate-mapping technologies revealed that lesional SMCs predominantly become fibroblast-like cells (sometimes called fibromyocytes)\(^21\). This transition depends on the transcription factor TCF21, deletion of which reduced the transition to fibromyocytes and, consequently, thinned fibrous caps. Single-cell RNA sequencing analyses of human atherosclerotic lesions and data sets from genome-wide association studies support a key role for TCF21: carriers of single-nucleotide polymorphisms that are associated with lower TCF21 expression have more coronary events\(^22,23\). These findings could accelerate SMC-targeted therapeutics, as both fibroblast-like and macrophage-like cells derived from SMCs could have roles in progressing lesions.

Macrophages: the central regulators

Macrophages are the most abundant leukocyte subset in atherosclerotic lesions. In healthy arteries, macrophages reside in the adventitia where they contribute to steady-state functions such as regulating the blood vessel diameter by interacting with SMCs\(^24\). In certain areas of the aortic tree, such as the lesser curvature of the aortic arch and at the branch points of large arteries, macrophages localize underneath the endothelium where they contribute to lipid retention\(^25\). Resident arterial macrophages in mice derive largely from embryogenic progenitors and can renew themselves\(^26,27\). In response to endothelial activation, namely as a consequence of hypertension, hyperglycaemia or hypercholesterolaemia, classical monocytes are recruited to the intima in a process involving integrins (such as α4β1; also known as VLA4) and chemokine receptors (including C-C chemokine receptor type 2 (CCR2) and CCR5)\(^28,29\) (Fig. 1a). Targeting monocyte integrins or chemokine receptors consistently reduced monocyte and macrophage accumulation and lesion burden in atherosclerotic mice, suggesting that recruitment is a main driver of lesional macrophage accumulation\(^30,31\). This is in agreement with a recent study that used a lineage-tracing approach to study lesional macrophage turnover\(^32\). Previous work has established that
Macrophages also proliferate locally, predominantly in advanced atherosclerotic plaques; the mechanisms and relative importance of lesional macrophage proliferation remain unclear. Scavenger receptor A signalling and LXRs phosphorylation, which control macrophage cholesterol content, seem to alter lesional macrophage proliferation, and administration of a statin decreased macrophage proliferation and lesion inflammation. In addition to contributing to endothelial cell activation along the arterial tree, metabolic risk factors also activate blood cell production in the bone marrow in a myeloid cell-biased fashion, thereby boosting the monocyte and neutrophil supply for delivery to the developing atheroma.

Activation of atheroma macrophages decisively determines the inflammatory environment in the plaque. Cholesterol crystals in the plaque co-activate the NACHT, LRR and PYD domains-containing protein 3 (NLRP3)-containing inflammasome, which has received considerable attention in the context of atherosclerosis. Inhibition of NLRP3 or its genetic deletion reduces atherosclerosis in mice. The output of the NLRP3 inflammasome requires two hits — priming and activation — thus unleashing caspase 1, which cleaves IL-1β and IL-18 to their mature forms. Oxidized LDL can prime and activate the NLRP3 inflammasome in atherosclerotic mice. Priming by oxidized LDL depends on the lipid binding to a complex containing CD36, Toll-like receptor 4 (TLR4) and TLR6; activation of NLRP3 follows lysosomal damage after internalization of the oxidized LDL. Inhibition of cholesterol efflux in myeloid cells increases inflammasome priming and activation, whereas induction of cholesterol efflux has the opposite effects. NLRP3 inflammasome activation can also be induced by cholesterol crystals through lysosomal damage. In summary, modified lipids activate the NLRP3 inflammasome in macrophages and cholesterol depletion from macrophages may mute the ensuing inflammatory response.

Macrophage function and phenotype can vary widely depending on several factors, including the microenvironment, the macrophage origin and the stage of the disease. Such determinants define macrophages in their global appearance and functionality but also define their heterogeneity on a single-cell level, and different subsets can coexist within one tissue. Although single-cell data from human plaques have become available, we await detailed analysis of the macrophage compartment and alignment with data from mouse studies.

Research in the past 10 years indicates that macrophage programming in response to numerous stimuli can persist even after the trigger has subsided. In fact, macrophages can acquire characteristics of immunological memory that were initially identified after brief exposure to microorganisms. This phenomenon, denoted ‘trained immunity’, involves a heightened cytokine response to a second stimulus, a process that reflects long-term cellular reprogramming. Although the initial concept of trained immunity focused on pathogens as the first stimuli, we now recognize that non-microbial endogenous stimuli, such as lipoproteins, can also initiate trained immunity. In mice, for example, a cholesterol-rich diet leads to a profound inflammatory transcriptional and epigenetic rewiring of monocytes in circulation and their bone marrow progenitors. Feeding-induced hypercholesterolaemia in these mice elicited a heightened inflammatory response to subsequent inflammatory stimuli, a response maintained even weeks after switching them to a normal chow diet.

Support for the clinical importance of such studies stems from observations made in patients with familial hypercholesterolaemia. Monocytes from these patients have a higher capacity to produce cytokines and have epigenetic marks — namely enrichment of histone H3 trimethylated at lysine 4 (H3K4me3), along with a decrease of H3K9me3 — in the promoter region of at least one of those cytokines, tumour necrosis factor (TNF). In vitro studies suggested that statins can reverse such training, the in vivo response remained heightened even after statin treatment. As IL-1β occupies a central role in inflammatory signalling, it is not surprising that this cytokine also contributes to innate immune training. IL-1β drives epigenetic reprogramming in monocytes reminiscent of the programme acquired by monocytes trained by β-glucan, the Bacillus Calmette-Guérin (BCG) vaccine or oxidized LDL. In addition, IL-1β exposure can elicit the metabolic changes seen during trained immunity in monocytes. Indeed, pharmacological inhibition of IL-1 using an anti-IL-1 receptor antibody, anakinra, annulled the metabolic switch, myeloid skewing and cell cycle modulation that occurred in haematopoietic stem cells following treatment with β-glucan.

Granulocytes: unusual suspects

Much attention has focused on macrophages as important regulators of atherosclerosis. Neutrophils, the most abundant circulating white blood cells in humans, have, until recently, received far less notice. Epidemiological evidence, however, indicates that counts of circulating neutrophils predict future cardiovascular events in humans, suggesting an association between neutrophils and atherosclerosis. Blood neutrophil counts in hypercholesterolaemic mice correlate with lesion sizes. Depletion of neutrophils during atherogenesis in hyperlipidaemic mice reduces lesional macrophage accumulation, indicating that neutrophils, in part, control macrophage migration. Indeed, neutrophils secrete several proteins with chemotactic activity for monocytes including CCL2, cathelicidin, cathepsin G and α-defensins. The latter forms heterodimers with platelet-derived CCL5, and therapeutic disruption of this complex reduces monocyte recruitment in mouse models of vascular inflammation.

Neutrophil secretory products not only attract but also activate macrophages. Neutrophil extracellular traps (NETs) prime macrophages in atherosclerotic mice to produce IL-1β by stimulating the NLRP3 inflammasome. In addition, macrophages may sense NETs through absent in melanoma 2 (AIM2), a cytosolic DNA-recognizing component of the inflammasome. NLRP3-containing and AIM2-containing inflammasomes have similar outcomes, through the activation of...
Endotoxaemia: The presence of bacterially produced endotoxins in the blood. Endotoxaemia can progress to septic shock.

NETosis: Neutrophil extracellular trap activation and release.

ST-elevation myocardial infarction (STEMI): A subtype of myocardial infarction, along with non-ST-elevation myocardial infarction, defined by changes in electrocardiogram recordings.

Risk factors in arterial inflammation

Plaque characteristics

The most dreaded and dramatic consequences of atherosclerosis involve thrombotic complications. A great deal of interest over many decades has focused on the rupture of a plaque's protective fibrous cap as a thrombosis trigger. Indeed, the term 'vulnerable plaque' has entered common parlance and refers to a lipid-rich lesion with a thin (<60 μm) fibrous cap overlying a necrotic core packed with macrophage foam cells. When the weak fibrous cap bursts open, coagulation proteins in the blood gain access to the thrombogenic mediator tissue factor in the lipid core, thus triggering thrombus formation.

A second type of plaque disruption involves erosion of the superficial layer of the intima without a fracture of the fibrous cap. Superficial erosion can occur in plaques that are extracellular matrix-rich and relatively lipid-poor. Numerous observational studies have shown that, coincident with the advent of effective lipid lowering and other preventive therapies, there has been a shift from ST-elevation myocardial infarction (STEMI) to more cases of non-ST-elevation myocardial infarction (NSTEMI). Contemporary data suggest that plaque erosion — rather than accounting for a fifth of ACS, as described previously — causes approximately a third of ACS cases. The application of optical coherence tomography, an intravascular imaging technique, to patients with ACS has shown that plaque erosion associates with NSTEMI whereas plaque rupture more commonly causes STEMI. As plaque rupture becomes less common, in part because of better control of classical...
Box 1 | Single-cell technologies in the arterial immune cell compartment

Single-cell RNA sequencing has contributed to characterizing the cellular heterogeneity of macrophages and smooth muscle cells in human and murine atherosclerotic lesions as well as of neutrophils in the circulation or in other tissues\(^{485,486}\). Although single-cell RNA sequencing is a great discovery tool for immune cell heterogeneity, several inherent limitations may preclude this technique from being ideal for drug target discovery. Single-cell RNA sequencing preferentially detects abundant genes, and genes expressed at low levels or periodically may not be detected. In addition, single-cell RNA sequencing requires generation of a single cell suspension; in the context of atherosclerotic lesions, time-consuming enzymatic digestion is unavoidable, which leads to uncertain yields of single cells and exposure to potential inducers of gene expression in the dissociation mixtures. Consequent artefactual expression of immediate early genes may skew the pattern of gene expression. In addition, mRNA expression for many proteins, including cell surface markers, poorly correlates with protein expression. For surface proteins, this is because of the complex translation and post-translational modification that cell surface proteins undergo, including glycosylation, cleavage of the signal peptide and vesicular trafficking. Thus, combining single-cell RNA sequencing and mass cytometry analyses may help to relate cell clusters identified by surface markers to transcriptional cell states. In addition, cellular indexing of transcriptomes and epitopes by sequencing (CITE-seq) strategies, which link cell surface phenotypes to transcriptomes by using oligonucleotide-tagged antibodies, thus combining single-cell RNA sequencing with quantitative measurements of cell surface markers, will help to obviate these issues. However, the major limitation for all of the methods discussed here is that information on the anatomical location and spatial relationship of cell types is lost. In recent years, several techniques have been developed to study the transcriptome and proteome in tissue in detail. Multiplexed ion beam imaging by time of flight (MIBI-TOF)\(^{487}\) and co-detection by indexing (CODEX)\(^{488}\) are techniques based on the use of a large number of antibodies tagged with isotopically pure elemental metal reporters or unique oligonucleotide sequences, respectively. Subsequent optical imaging allows for subcellular resolution of structures labelled in this way. Currently, no data sets employing these techniques are available from human or mouse atherosclerosis specimens. MIBI-TOF and CODEX provide spatial information on a limited number of antibody-detected proteins, and additional techniques have evolved that allow visualization of the transcriptome in tissue sections. Spatial transcriptomics using a spatially encoded barcoded bead array at a resolution of as little as 2 µm has been reported to yield high-definition transcriptomics\(^{489}\). Super-resolution imaging and multiplexing of up to 10,000 genes in a single cell can be achieved by RNA sequential fluorescence in situ hybridization (seqFISH)\(^{490}\). However, all of these methodologies confront the limitation that protein and RNA are not detected simultaneously. Recently developed deterministic barcoding in tissue for spatial omics sequencing (DBit-seq) for co-mapping of mRNAs and proteins in a tissue section has enabled the first protocol that can detect both protein and RNA\(^{491}\). Such techniques, and others in development, will doubtlessly be applied to arterial inflammation. Extension of these methods from experimental to human atherosclerosis promises to provide important information on the spatial relationship of ligand–receptor pairs and cell subsets, and to enrich the discovery and validation of drug targets.

risk factors, the proportion of ACS caused by superficial erosion and, thus, NSTEMI may be on the rise. LDL elevation may be permissive to atheroma formation, and likely contributes to all causes of plaque disruption. The decrease in the lipid content and the relative increase in fibrous tissue in the plaques of patients who have undergone intensive LDL lowering, which renders these lesions relatively lipid-poor, raises the possibility that erosion might strongly contribute to the residual burden of ACS events in these individuals, despite highly effective lipid control\(^{28,29}\).

Attack of the clones

Cells in many tissues acquire somatic mutations over time. Most such somatic mutations will never manifest as disease. However, acquired mutations in a small subset of known driver genes for leukaemia can give rise to clones of myeloid cells that circulate in peripheral blood\(^{20}\). Exome sequencing of blood has shown that up to 10% of septuagenarians have such clones of mutant leukocytes\(^{20,40}\). The transition to a haematological malignancy requires the serial mutation of three or more leukaemia driver genes. Individuals with a single acquired mutation, which can lead to clonal haematopoesis of indeterminate potential (CHIP), transition to acute leukaemia at a rate of 1% per year or lower. However, individuals with CHIP have a strikingly increased risk for cardiovascular events that far outstrips their tendency to develop leukaemia\(^{41}\). Fully adjusted for all traditional risk factors, having clonal haematopoesis confers an almost twofold increase in the risk of acute myocardial infarction or stroke\(^{40,42}\). Individuals with CHIP also have worse outcomes when affected by heart failure or following percutaneous aortic valve replacement. The two most commonly mutated genes in CHIP regulate the methylation of DNA. The augmented cardiovascular risk may thus derive from epigenetic changes in gene expression. Interestingly, the cause and consequence may also be inverted in some cases; traits of atherosclerosis can cause CHIP through continuous stimulation of stem cell proliferation\(^{43}\). In agreement with previous observations\(^{44}\), a combination of risk factors of atherosclerosis accelerated stem cell cycling and, hence, increased the expansion of clones bearing the somatic mutations that cause CHIP.

Convergent data from several sources implicate altered inflammatory signalling as a key consequence of clonal haematopoesis due to mutations in DNMT3A, TET2 and JAK2 [\(\text{REFS}^{45,46,47}\)]. Recent work identified increased macrophage proliferation and prominent necrotic core formation in atherosclerotic lesions of mice expressing JAK2\(^{48,49}\), a gain-of-function mutation that increases JAK–STAT signalling and associates with a substantial risk of premature coronary heart disease\(^{50}\). Deletion of the essential inflammasome components caspase 1 and caspase 11, the pyroptosis executioner gasdermin D or the double-stranded DNA-sensing inflammasome AIM2 (but not NLRP3) reversed these adverse changes\(^{51}\). CHIP — a potent, common, age-related, independent and newly recognized risk factor for adverse cardiovascular outcomes — could be a novel link between inflammation and cardiovascular disease that was completely unsuspected just a few years ago. These recent observations provide a fertile field for further exploration of the roles of inflammation in cardiovascular disease. Curiously, CHIP associates inconsistently with increases in the widely used biomarker of inflammation, C-reactive protein (CRP)\(^{52}\). This apparent paradox implies that there are aspects of cardiovascular inflammation beyond those captured by elevations in CRP.

Acute infections

Excess mortality from cardiovascular disease during influenza epidemics was initially recognized early in the twentieth century, and recent reports indicate that COVID-19 may be associated with a similar increase\(^{53,54}\). In a recent study, the risk for an acute coronary event within 1 week after laboratory-confirmed infection with respiratory syncytial virus or influenza virus was four
or six times higher, respectively, than the risk during the previous year8. Several-fold increases in ACS risk were also found for acute bacterial pneumonia, urinary tract infection and bacteremia with one of several bacterial strains9,10. Of note, the increased risk peaks within the first couple of weeks after infection and relates to the severity of the infection. Thus, acute viral or bacterial infection can augment a pre-existing cardiovascular risk. A connection between infection and ACS pertains to a variety of pathogens, and as severity of infection correlates with ACS risk, it is likely that the host response to the infection is a major determinant. Despite this overwhelming epidemiological evidence, there is surprisingly little experimental understanding of how an acute infection accelerates atherosclerosis. As one obvious mechanism, circulating cytokines, a consequence of acute infection, may activate lesion-resident inflammatory cells11,12. This mechanism doubtlessly contributes to advanced COVID-19, which is characterized by excessive cytokine production and multi-organ system failure13. Polymicrobial sepsis in mice aggravated atherosclerosis in as few as 24 h, and atherosclerosis increased with time14. Mechanistically, abdominal sepsis induced the expansion of the lesional macrophage population, likely as a consequence of increased recruitment. In addition, increases in lesional TNF, IL-6 and CCL2 levels indicate induction of the local arterial cytokine and chemokine network. At the intimal surface, increases in platelet activity associate with myocardial infarction in patients with pneumonia15. This observation may relate to the direct prothrombotic state evoked by activated platelets, the ability of platelets to induce NETs16 or their ability to team up with neutrophils to promote arterial monocyte recruitment17,18. Treating the inciting infection and providing haemodynamic and respiratory support are part of the established tools to fight the complications of sepsis. A suite of clinical trials currently underway will evaluate the efficacy of various anti-thrombotic and anticoagulant regimens in severe COVID-19 infections.

**Clinical studies**

Thousands of publications describe experimental studies of inflammation in atherosclerosis, and numerous biomarker studies have probed the relationship between inflammation and cardiovascular events. Yet, until recently, we lacked direct evidence from properly powered and well-controlled clinical trials that an anti-inflammatory intervention could improve cardiovascular outcomes. We have now entered an era when clinical investigations have begun to close this gap between experiment and observation, and clinical practice (Table 1).

**Targeting innate immunity**

CANTOS was the first trial to validate the concept that targeting inflammation could be therapeutically relevant in human atherosclerosis19. CANTOS enrolled more than 10,000 individuals who had sustained an acute myocardial infarction but were in the stable phase at least 1 month after the qualifying event20. The enrollees all received standard care: guideline-directed medical therapy including statin administration. Indeed, the baseline LDL concentration of approximately 2 mM (81 mg/dl) established excellent treatment of this classical risk factor. CANTOS selected individuals who had CRP levels, measured with a high sensitivity CRP assay (hsCRP), greater than 2 mg/l; this cut-off value is approximately the median for a general population. Thus, CANTOS preselected individuals deemed likely to benefit from anti-inflammatory intervention by showing evidence of residual inflammatory burden despite standard therapy. CANTOS met its primary end point and showed a 15% reduction in major adverse cardiovascular events, including myocardial infarction, stroke or cardiovascular death, in individuals treated with the anti-IL-1β antibody canakinumab (150 mg four times per year). In addition, in an exploratory analysis, individuals treated with canakinumab had a reduction in incident cancer and a striking decrease in mortality from lung cancer21.

A secondary analysis of CANTOS stratified outcomes based on the achieved hsCRP levels 3 months...
Overview of selected clinical studies targeting inflammatory pathways in cardiovascular disease

| Trial | Study population | Study design | Outcome | Ref. |
|-------|------------------|--------------|---------|------|
| ASSAIL-MI | First-time STEMI presenting within 6 h of the onset of chest pain | Single dose of tocilizumab (IL-6 antibody) vs placebo | Improved myocardial salvage in patients assigned to tocilizumab | NCT03004703 |
| CANTOS | Stable CAD, persistent elevation of hsCRP (>2 mg/l) | Canakinumab (IL-1β antibody) subcutaneously vs placebo | Canakinumab lowered plasma CRP, IL-1 and IL-6 | NCT01327846 (REF.7) |
| CIRT | Stable CAD and persistent evidence of inflammation, type 2 diabetes or metabolic syndrome | Low-dose (15–20 mg) methotrexate (a purine metabolism inhibitor) once per week vs placebo | Halted prematurely for futility | NCT02576067 (REF.25) |
| COLCOT | Recent myocardial infarction (<30 days) | Low-dose (0.5 mg/day) colchicine (a tubulin disrupter) vs placebo | Reduction in cardiovascular death and cardiovascular events | NCT02551094 (REF.7) |
| CLEAR-Synergy | STEMI with primary PCI | SYNERGY bioabsorbable polymer drug eluting stent plus colchicine and spironolactone or placebo | Increase in pneumonia | NCT03048825 |
| CONVINCE | Adults >40 years of age with an ischaemic stroke or TIA not caused by cardiac embolism | Low-dose (0.5 mg/day) colchicine plus usual care or standard care alone | Ongoing, estimated completion in early 2025 | NCT02898610 |
| LATITUDE-TIMI 60 | Patients hospitalized with acute myocardial infarction | Losmapimod (a selective inhibitor of p38α/β mitogen-activated protein kinases) twice per day vs placebo | No reduction in major ischaemic cardiovascular events | NCT02145468 (REF.20) |
| LoDoCo | Stable CAD | Low-dose (0.5 mg/day) colchicine plus usual care or standard care alone | Significant reduction in ACS | 64 |
| LoDoCo2 | Chronic coronary disease | Low-dose (0.5 mg/day) colchicine plus usual care or standard care plus placebo | Reduction in cardiovascular events | ACTRN12614000093684 (REF.4) |
| LILACS | Stable ischaemic heart disease and ACS | Low-dose IL-2 (0.3–3 × 10^6 I.U./day) | Phase I/II ongoing | NCT03113773 |
| FUTURE 1 | Psoriatic arthritis (prospective randomized) | Secukinumab (IL-17A antibody) vs placebo | Improved arthritis score | NCT01392326 (REF.209) |
| Tocilizumab in NSTEMI | NSTEMI | Single dose of tocilizumab (IL-6 receptor antibody) vs placebo | Reduction in hsCRP and troponin T release | NCT01491074 (REF.205) |

ACS, acute coronary syndromes; CAD, coronary artery disease; CANTOS, Canakinumab Anti-inflammatory Thrombosis Outcomes Study; CIRT, Cardiovascular Inflammation Reduction Trial; COLCOT, Colchicine Cardiovascular Outcomes Trial; CRP, C-reactive protein; hsCRP, CRP measured with a highly sensitive assay; MACE, major adverse cardiac event; NSTEMI, non-ST-elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST-elevation myocardial infarction; TIA, transient ischaemic attack.

after initiation of therapy. Participants who responded to the anti-inflammatory drug by a greater than median reduction in hsCRP levels showed a 26% decrease in the prespecified primary end point of myocardial infarction, stroke or cardiovascular death, and a reduction in total mortality as well. Gout attacks decreased by more than half6, and patients treated with canakinumab were hospitalized or died from heart failure less often8 and had less incident anaemia7 than those treated with placebo.

These benefits of canakinumab therapy in this population came at a price. There was a small but statistically significant increase in infections, including fatal infections, in those receiving canakinumab. These infections were from common viral and bacterial agents; there was no increase in infections with opportunistic pathogens or tuberculosis5. These findings can inform the design of future trials and guide the safer deployment of anti-inflammatory therapies in the future, and indicate the need for further anti-inflammatory therapies to optimize the benefit–risk ratio.

Smaller trials blocked both IL-1α and IL-1β with the receptor antagonist anakinra. The MRC–ILA study enrolled patients with NSTEMI and showed a decrease in the area under the curve of the inflammatory biomarker CRP18. Likewise, Virginia Commonwealth University Anakinra Remodeling Trial 3 (VCUART3)
A second anti-inflammatory trial addressed the use of low-dose weekly methotrexate in individuals who had had or were at high risk for coronary events. Methotrexate, a disease-modifying antirheumatic drug, has transformed the practice of rheumatology and has also benefited patients with psoriasis. Observations of cardiovascular events in individuals enrolled in the studies of low-dose methotrexate for rheumatological or dermatological indications showed strong evidence for a cardiovascular benefit. The Cardiovascular Inflammation Reduction Trial (CIRT) tested this hypothesis prospectively in a randomized controlled study. However, CIRT halted prematurely on the advice of the data safety monitoring committee because of futility. The enrolled population in CIRT, which was not selected a priori on the basis of elevated inflammation as indicated by hsCRP, had a much lower inflammatory burden than individuals enrolled in CANTOS. The mean baseline CRP in CANTOS was 4.2 mg/l, compared with 1.5 mg/l in CIRT. Moreover, although canakinumab handily decreased CRP levels in participants of CANTOS, biomarkers of inflammation, including hsCRP, IL-1β and IL-6, did not fall in individuals receiving methotrexate. In contrast with CANTOS, which showed a reduction in cancer, CIRT showed a significant increase in cutaneous cancer, raising another potential limitation of anti-inflammatory therapy: interference with immune surveillance for malignancy.

In sum, the results of CIRT may indicate that, despite the observational data, methotrexate does not reduce cardiovascular events. Alternatively, the enrolled population in CIRT may have already exhibited tamed inflammation, as indicated by the normal baseline hsCRP levels, and stood to benefit little from additional anti-inflammatory therapy. The negative result from CIRT underscores the vital importance of conducting rigorous randomized controlled clinical trials in this arena as the results of CIRT differed dramatically from the observational analyses, which were not designed to evaluate and did not duly adjudicate cardiovascular events.

Colchicine, a natural product, has been used for millennia for inflammatory diseases. In the cardiovascular field, colchicine has become the mainstay for management of pericarditis, particularly recurrent pericarditis. The low-dose colchicine (LoDoCo) study enrolled 532 individuals in an open-label, non-placebo-controlled study with 3 years of follow up. There was a remarkable reduction of more than two-thirds in the primary end point of major adverse cardiovascular events. Despite the small size of this study, which only had 55 primary end point events, the LoDoCo results spawned two large clinical trials, the Colchicine Cardiovascular Outcomes Trial (COLCOT) and LoDoCo2. COLCOT enrolled individuals within 30 days following an ACS event. Nearly 5,000 participants received 0.5 mg of colchicine daily or a placebo. The participants treated with colchicine, after almost 2 years of follow up, showed a statistically significant reduction in a primary end point that combined “hard” events such as myocardial infarction and stroke with other end points, notably hospitalization for angina leading to coronary revascularization. The clinical benefit was a 23% reduction in the primary end point, driven primarily by a reduction in urgent hospitalizations for angina requiring revascularization. LoDoCo2, which was of similar size to COLCOT, focused on individuals more than 6 months after their last coronary event or revascularization. After a 30-day open-label run-in with 0.5 mg of colchicine daily, those who tolerated the therapy were randomly allocated to either continue on colchicine or receive placebo. This second large study met its primary, event-driven end point, which included cardiovascular death, non-fatal ACS or non-fatal stroke.

COLCOT and LoDoCo2 neatly bracket two important populations: those with recent ACS, and those in the stable phase of coronary artery disease. The information from these two trials could well change practice and make anti-inflammatory therapy a mainstay of care for individuals post ACS. In COLCOT, colchicine showed unexpectedly little gastrointestinal intolerance. As in CANTOS, colchicine treatment was associated with a small but significant increase in infections, notably pneumonia.

In the wake of COLCOT and LoDoCo2, a raft of trials employing colchicine in various scenarios, ranging from acute myocardial infarction to post-percutaneous intervention, have been performed, are underway or are planned. The colchicine in patients with STEMI/synergy stent registry (NCT03048823) is evaluating, using a factorial design, the combination of colchicine and spironolactone on end points including heart failure. Thus, the upcoming results from a number of studies will inform us regarding the clinical utility of colchicine in various forms of cardiovascular disease.

Beyond IL-1β and colchicine, a number of other targets appear worthy of investigation in the realm of innate immunity. Like methotrexate, agents that antagonize TNF have proven highly effective in diseases such as rheumatoid arthritis and inflammatory bowel disease. TNF antagonism, however, yielded a safety concern in one of the trials that used this strategy in patients with heart failure. Herein lies another important message: despite robust experimental evidence in animals and smaller pilot studies in humans that showed a benefit, the large-scale randomized trial not only showed no benefit but suggested possible harm at a higher dose. For these reasons, anti-TNF therapy has not engendered enthusiasm in patients at risk for heart failure, including those with coronary artery disease after ACS.

IL-6 is strongly induced by IL-1 (REFS 96,97), and likely participates causally in human cardiovascular events as shown by concordant Mendelian randomization studies98,99. Thus, agents that neutralize this cytokine also merit evaluation for their capacity to reduce cardiovascular events. However, tocilizumab, a readily available therapeutic that targets the IL-6 receptor (IL-6R), consistently causes an increase in triglycerides. This finding has diminished enthusiasm for long-term studies with this agent in individuals with coronary artery disease. Abundant human genetic evidence points to triglyceride-rich lipoproteins as causal factors in
human cardiovascular events. Nonetheless, studies on the short-term administration of an IL-6R antagonist during acute myocardial infarction have yielded encouraging signals for benefit based on cardiac magnetic resonance imaging\(^{110}\). A larger clinical trial is now underway. A phase II study in patients with chronic kidney disease treated with the monoclonal antibody ziltivekimab, which neutralizes IL-6, will be completed in 2021 and could serve as a prelude to a large phase III trial.

Targeting other cytokines may also be valuable. IL-17 isoforms have a controversial role in experimental atherosclerosis\(^{111}\). Although antibodies that neutralize IL-17A benefit patients with certain inflammatory diseases, no rigorous trial has tested their use in patients with atherosclerosis. IL-23, another potential adaptive immune target, has been considered in atherosclerosis but in some studies may produce cardiovascular harm, limiting the enthusiasm for anti-atherosclerotic therapeutics that target IL-23 (REF.\(^{112}\)).

### Targeting adaptive immunity

The adaptive immune system has also received attention as a potential target in atherosclerotic cardiovascular disease. Seemingly paradoxically, immunization with oxidized forms of LDL can mitigate experimental atherosclerosis\(^{113,114}\). These observations led to a number of attempts to modulate atherosclerosis using vaccination. Although preclinical data were promising, despite considerable effort, none of these vaccination strategies have yielded clinical data that substantiate cardiovascular event reduction in patients\(^{115}\).

Strong preclinical evidence supports the operation of regulatory T cells (T\(_{\text{reg}}\) cells) in experimental atherosclerosis. Mitigation of inflammation by transforming growth factor-\(\beta\) (TGF-\(\beta\)) and other mediators released by T\(_{\text{reg}}\) cells can mute experimental atherogenesis\(^{116}\). This observation has led to an innovative clinical trial using low-dose IL-2 to skew the T cell balance towards T\(_{\text{reg}}\) cells\(^{117}\). Biomarker studies in humans are substantiating the importance of T\(_{\text{reg}}\) cells\(^{118}\). We look forward to a large-scale outcomes-based trial that would substantiate the approach of modulating adaptive immunity in patients with atherosclerosis.

### The importance of biomarkers

As shown by contrasting the results of CANTOS and those of CIRT, the use of biomarkers to select individuals who are particularly likely to benefit from a therapy provides a potential pathway to success. This kind of patient stratification has been used for therapies that alter LDL levels and blood pressure for generations. In the oncology field, the use of genetic markers to target therapies has become routine. Today in the cardiovascular arena, save perhaps for transthyretin mutations, we have no therapies targeted on the basis of genetic testing.

Harnessing the emerging field of genetic markers of cardiovascular risk might provide a path to precisely target particular anti-inflammatory agents to certain patients. For example, the proposal that individuals with CHIP due to loss-of-function mutations in DNMT3A or TET2 might benefit from an anti-IL-6 strategy warrants further investigation. Likewise, those with clonal haematopoiesis due to mutations in JAK2 might benefit from administration of a JAK1/2 inhibitor such as ruxolitinib\(^{119,120}\). The use of biomarkers, including genetic variants, as a strategy for precision targeting of anti-inflammatory therapies could permit smaller, quicker and more economical end point studies that are ultimately needed to establish benefit and receive approval from regulatory authorities.

### Preclinical strategies

#### Limiting arterial inflammation

With the success of CANTOS, several strategies have emerged to interfere with IL-1\(\beta\) or its production, including inhibition of inflammasome activation. In addition, there are currently several promising avenues of research that have not yet entered the clinical trial phase but are in active development. In this section, we focus on selected aspects of controlling arterial inflammation and direct the reader to recent overview articles for broader information\(^{121,122}\).

Closing the gates for myeloid cells. A group of small chemotactic cytokines known as chemokines orchestrate immune cell trafficking. Upon binding to G-protein-coupled receptors, chemokines regulate immune cell movement in steady state as well as during inflammation. Given the importance of inflammasome activation during atheroprophosis, antagonizing chemokine–receptor interactions may be a promising therapeutic avenue.

Activated endothelial cells can release C-X-C motif chemokine 1 (CXCL1), which interacts with C-X-C chemokine receptor type 2 (CXCR2) on myeloid cells, thereby promoting mobilization from the bone marrow and recruitment to sites of inflammation, including the atherosclerotic lesion. Thus, genetic deletion of CXCR2 from bone marrow cells or antibody-mediated neutralization of CXCL1 reduces the atherosclerotic burden and lesional macrophage accumulation in mice\(^{12}\). Mobilization of classical monocytes from the bone marrow is under control of the CCR2–CCL2 axis\(^{12}\) and genetic deletion of Ccr2 greatly reduces atherosclerotic lesion sizes, likely due to monocytopenia in this model\(^{112,123}\). In myocardial infarction in mice, small interfering RNA (siRNA)-mediated silencing of CCR2 or delivery of a non-agonistic CCL2-competing mutant protein that exhibits strong proteoglycan binding lowered monocyte recruitment, ventricular remodelling and ischaemia–reperfusion injury\(^{121,124}\) (FIG. 2a). In humans, higher plasma CCL2 levels are associated with a higher risk of cardiovascular events and higher lesional CCL2 associates with features of plaque destabilization\(^{125,126}\). Blocking CCR2 with an antibody that blocks CCL2 binding lowers CRP levels in patients at risk of cardiovascular disease\(^{121}\). Upon ligation of the chemokine receptor, a signalling cascade ensues that, in many cases, leads to integrin activation and, consequently, to cell adhesion. In atherosclerosis, platelets are a prominent source of CCL5, which, when immobilized on arterial endothelium, promotes monocyte adhesion and recruitment\(^{121}\). Consequently, lack of CCR5, a receptor of CCL5, or inhibition of CCR5 with maraviroc,
a FDA-approved inhibitor of HIV entry, lowers the degree of atherosclerosis and lesion macrophage content in hypercholesterolaemic mice \(^{115,119}\). In a clinical setting, treatment of patients with HIV with maraviroc reduces atheroprotection \(^{121}\).

An intriguing aspect of chemokines is their ability to form oligomers either with themselves or with another partner. In this context, CCL5 exhibits a high degree of promiscuity by forming complexes with chemokines CXCL4 or CCL17, the neutrophil-derived peptide neutrophil defensin 1 (DEFA1; also known as HNP1) as well as galectin 3 (REFS \(^{53,122–124}\)). Functionally, CCL5–CXCL4 complexes as well as CCL5–DEFA1 complexes increase inflammatory monocyte recruitment, and selective disruption with cyclical peptides decreases atherosclerosis formation or improves recovery after myocardial infarction, respectively (FIG. 2a).

**Multiple ways to target NETs.** The recognition of the importance of NETs in arterial thrombosis, vascular inflammation and vascular injury during the past decade demonstrates the potential of NET release pathways,

**Fig. 2 | Preclinical strategies to limit cardiovascular inflammation and stimulate its resolution.** a | Reducing monocyte recruitment. Silencing of C-C chemokine receptor 2 (CCR2) or timed inhibition of CCR2 signalling reduces monocyte adhesion. Overriding chemokine receptor signalling, for example with the small molecule Ac2-26, reduces integrin activation and monocyte arrest. Heterodimers of C-C motif chemokine 5 (CCL5) and C-X-C motif chemokine 4 (CXCL4) as well as of CCL5 and neutrophil defensin 1 (DEFA1) promote monocyte adhesion. Small peptides that disrupt these interactions reduce monocyte adhesion during cardiovascular inflammation. b | Inhibiting neutrophil extracellular traps (NETs). Inhibition of protein-arginine deiminase type 4 (PAD4) halts NET release. DNase I cleaves DNA strands in NETs. Neutralization of gasdermin D prevents NET discharge. Neutralization of NET-resident proteins by antibodies or small-molecule inhibitors reduces NET-driven inflammation. c | Examples of strategies to inhibit immune checkpoints. Neutralization of CD80/86 can reduce T cell and dendritic cell responses. The CD40–CD40 ligand (CD40L) interaction activates macrophages via intracellular TNF receptor-associated factor 6 (TRAF6) signalling, a cascade that can be inhibited with small molecules. d | Increasing inflammation resolution. Putrescine improves the ability of macrophages to engulf dead cells. Resolving N-formyl peptide receptor 2 (FPR2) agonists (Ac2-26, lipoxin A4, resolvin D1) and chemokine-like receptor 1 (CMKLR1) agonists (chemerin C15, resolvin E1) lower the production of inflammatory cytokines and improve the efferocytosis capacity, NF-κB, nuclear factor-κB.
the NET structure and components of NETs as therapeutic targets. Numerous experimental studies have utilized an inhibitor of protein-arginine deiminase type 4 (PAD4), chloramidine, to inhibit histone citrullination and NET release\(^{60,65,125}\) [Fig. 2b]. PAD4 is an enzyme enriched in the cytosol of haematopoietic cells. Upon translocation to the nucleus, PAD4 citrullinates transcription factors, thereby participating in epigenetic regulation of gene expression and stem cell differentiation\(^{126,127}\). The citrullination of arginine residues in histones is an important step in chromatin decondensation during NET release\(^{128}\). Although it is useful as a preclinical agent, chloramidine inhibits several isoforms of PAD, and hence lacks the desired specificity of a clinical therapeutic. A therapeutic anti-citrullinated protein antibody (tAPCA) can inhibit NET release and improves ical therapeutic. A therapeutic anti-citrullinated protein antibody (tAPCA) can inhibit NET release and improves signs of inflammation in a mouse model of arthritis\(^{129}\).

Although the mechanism through which this antibody inhibits NET formation is unknown, these observations made in mice are encouraging. Beyond limiting NET release, dissolving NETs provides an alternative therapeutic approach. DNase I, for example, breaks up double-stranded DNA and, hence, dissolves NETs. DNase is a FDA-approved drug used in patients with cystic fibrosis to clear bronchial mucous, which is rich in neutrophil products. DNase administration has proven useful in murine atherosclerosis\(^{130}\), and there is also early clinical evidence that it reduces myocardial infarction following ischaemia–reperfusion injury\(^{130,131}\). Gasdermin D is also critically involved in NET release\(^{32}\). Gasdermin D itself is cleaved by neutrophil elastase during NETosis and, in turn, activates neutrophil elastase, furnishing a feed-forward loop\(^{132}\). Given the importance of gasdermin D during NET formation, inhibitors of gasdermin D, such as necrosulfonamide, may harbour therapeutic potential during cardiovascular inflammation\(^{133}\).

Targeting proteins embedded within the NETs could also be therapeutically useful. For example, inhibitors of myeloperoxidase might quell one of the NET-associated generators of reactive oxygen species. Neutralization of NET-associated neutrophil elastase and cathepsin G may prevent NET-driven coagulation\(^{134}\). In addition, NET-bound H4 can trigger death of intimal SMCs and accelerate plaque destabilization in mice\(^{135}\). Therapeutic interference using either antibodies or a cyclical histone interference peptide that neutralizes the positive charge at the N terminus of H4 reduced SMC death and conferred plaque stability. A similar approach was chosen to target H2a during endotoxaemia. In this experiment, neutralization of H2a with an antibody or a small peptide lowered myeloid cell adhesion to the carotid artery and the consequent lesion expansion promoted by NETs\(^{136}\).

**Titrating the immune response.** Immune checkpoint regulators comprise central control elements in inflammation and can accelerate or halt leukocyte activation upon interaction with antigen-presenting cells. Upon T cell receptor ligation, co-stimulatory molecules induce T cell activation. Co-inhibitory receptor–ligand interactions, in turn, dampen T cell responses and thereby contribute to tolerance towards self-antigens. The potential for therapeutic interference in immune checkpoint modulation has been demonstrated in the field of oncology, in which agents targeting PD1 or CTLA4 are now used routinely\(^{137}\). By contrast, this knowledge is just now being applied to vascular inflammation. Mice lacking functional PD1 and PD2 exhibit augmented atherosclerosis, supporting the notion that checkpoint inhibition operates in this disease\(^{138}\).

The interaction between CD40 and CD40 ligand (CD40L) promotes T cell activation but also stimulates macrophages and dendritic cells. Deletion or antibody-assisted inhibition of either CD40 or CD40L lowers plaque burden and induces plaque stability in mouse models of atherosclerosis\(^{137,140}\). Clinical observations indicate that the CD40–CD40L axis is also important in human cardiovascular disease. Levels of soluble CD40L, for example, associate with primary risk factors such as hypercholesterolaemia and diabetes and can predict future cardiovascular disease\(^{141}\). Expression of CD40 and CD40L in human plaque specimens associates with features of instability. CD40 signals through TNF receptor-associated factors (TRAFs), and activation of TRAF6 but not TRAF2, TRAF3 or TRAF5 augmented atherosclerosis in mice\(^{142}\). This finding stimulated the generation of small-molecule inhibitors that selectively target the interaction between CD40 and TRAF6. These so-called TRAF-STOPs can reduce atherogenesis and prevent atheroprogession in hypercholesterolaemic mice\(^{143}\) [Fig. 2c]. Because these molecules do not interfere with TRAF2, TRAF3 or TRAF5, they should impair host defences less than inhibiting CD40–CD40L interactions does. Nanodelivery techniques might further alleviate such unwanted effects. Indeed, encapsulation of TRAF-STOPs into recombinant high-density lipoprotein (rHDL) nanoparticles (rHDL-6877002), delivered into mice with pre-established lesions, reduced macrophage accumulation\(^{144,145}\), rHDL-6877002 was safe, and had a favourable biodistribution in non-human primates\(^{146}\).

The interactions between the immune checkpoint proteins CD80/86 and CD28 or CTLA4 have received considerable attention in cardiovascular inflammation. In human lesions, the expression of CD80/86 correlates with lesion vulnerability\(^{147}\). In turn, mice lacking CD80/86 develop less atherosclerosis with lower effector T cell activity, suggesting a connection between CD80/86 and plaque development\(^{148}\). Overexpression of CTLA4 in mice likewise reduced T cell activation and plaque development\(^{149}\). Pharmacological inhibition of the interaction between CD28 and CD80/86, using a CTLA4 fusion protein, lowered atherosclerosis development in hypercholesterolaemic mice\(^{150}\), supporting the overall viability of therapeutically targeting CD80/86 co-stimulatory signalling in cardiovascular inflammation.

**Harnessing trained immunity.** As described above, trained immunity involves immune cell reprogramming upon an initial inflammatory stimulus that elicits a hyper-responsive state to a secondary stimulus; this phenomenon’s underlying molecular mechanisms provide an innovative targeting framework (reviewed elsewhere\(^{151}\)). Although only at its infancy, therapeutic
targeting of trained immunity could have relevance to cardiovascular inflammation. In the context of blockade of cholesterol synthesis, fluvastatin blunts trained immunity induction by β-glucan in vitro, and hence delivery of statins to macrophages, in the form of HDL nanoparticles, may reduce vascular inflammation, in part, by reducing immune training. IL-1β and granulocyte–macrophage colony-stimulating factor (GM-CSF) are key regulators of trained immunity in bone marrow progenitors, so antibody-assisted neutralization of these two factors may also inhibit trained immunity. Antibodies to IL-1β tamed cardiovascular inflammation as shown in CANTOS, and antibodies to GM-CSF are at an advanced phase of clinical development for various inflammatory conditions including asthma and rheumatoid arthritis. Finally, trained immunity involves epigenetic regulation, so inhibitors of histone or DNA methylation can modulate this process. A recent article provides details and a broader overview of targeting of trained immunity.

Inflammasomes and their output. In the wake of CANTOS, several approaches have emerged that therapeutically interfere with inflammasome priming and activation as well as the main product of inflammasome activation, mature IL-1β. Inhibition of nuclear factor-κB (NF-κB) signalling has been the primary goal for targeting inflammasome priming, as NF-κB transcriptionally regulates NLRP3, and several small-molecule drugs targeting the NF-κB pathway have been successfully employed in preclinical models of cardiovascular inflammation. However, inhibition of NF-κB signalling will have broad effects beyond the expression of components of the inflammasome complex — non-specific immunosuppression and impaired host defence may result.

The most thoroughly characterized inhibitor of the NLRP3 inflammasome is MCC950 (also known as CP-456,773). This compound potently inhibits NLRP3 activation, preventing ATP-triggered, NLRP3-driven IL-1β release. In mice, MCC950 reduced the lesion burden and macrophage accumulation in hypercholesterolaemia and hyperglycaemia-induced atherosclerosis, MCC950 was also tested in phase II clinical trials for rheumatoid arthritis. However, the compound was not tested further owing to liver toxicity. Key reports on the importance of NLRP3 in atherosclerosis and other manifestations of chronic inflammation have engendered intense interest in the development of specific antagonists of the NLRP3-containing inflammasome. Recent work indicates that the AIM2 inflammasome is also important hypercholesterolaemia-driven and CHIP-accelerated atherosclerosis, and consequently AIM2 may be an alternative target in age-related atherosclerosis.

Downstream of inflammasome activation, inhibiting caspase 1 could reduce the release of active IL-1β. The small-molecule inhibitors VX-740 and VX-765 are caspase 1 inhibitor prodrugs that are intracellularly activated by esterases. VX-765 can mitigate atherosclerosis in mice. However, as in the case of MCC950, clinical studies did not proceed beyond phase II clinical trials due to concerns regarding hepatotoxicity. We have provided a few illustrative examples of inflammasome targeting in this section, and refer the reader to comprehensive reviews of this topic.

Stimulating inflammation resolution
A typical acute inflammatory response is self-limiting: a resolution phase follows the acute inflammatory reaction to an injury, limiting inflammation and evoking reparative processes. Such organized responses promote the return to tissue integrity and function. Pro-resolving pathways and mediators have been mapped. Whereas agents neutralizing pro-inflammatory mediators have long been investigated as potential drugs, transferring the concepts of inflammation resolution from bench to bedside requires a paradigm shift, from antagonism towards agonism. The main advantage of immuno-resolvants is that they would limit several core processes of atherosclerosis simultaneously: reducing leukocyte infiltration and the production of pro-inflammatory mediators, and increasing the containment and phagocytosis of cellular debris and apoptotic cells. Therapies that actively promote resolution may also enhance innate immune responses to bacterial infections, whereas established anti-inflammatory therapies, including TNF and IL-1β neutralizing strategies, may compromise host defence. For example, in CANTOS, IL-1β neutralization led to a small yet statistically significant increase in lethal infections. Inflammation resolution is controlled by various classes of mediators including peptides, lipids and gases (including NO and H2S). Numerous cell types, such as Treg cells and myeloid-derived suppressor cells, regulate resolution as well as vernal innervation. In this section, we focus on peptides and lipids, as these have been extensively studied in the context of atherosclerosis. Inflammation resolution has been reviewed elsewhere.

Inflammatory and resolving mediators converge at FPR2 and CMKLR1. N-Formyl peptide receptor 2 (FPR2) is a G-protein-coupled receptor expressed on the cell membrane of many cell types relevant to atherosclerosis, including monocytes, macrophages and neutrophils as well as SMCs. The FPR gene family has a complex evolutionary history and, as a consequence, these proteins sense not just formylated peptides of prokaryotic and eukaryotic origin but also an array of endogenous pro-inflammatory and pro-resolving peptides and lipids. In addition to formylated peptides, serum amyloid A and cathelicidin trigger inflammatory responses via FPR2. In the context of atherosclerosis, cathelicidin serves as a chemoattractant for monocytes and, hence, promotes the early stages of atherosclerosis. With regard to resolving ligands, lipoxin A4, aspirin-triggered lipoxin, resolvin D1, resolvin D3 and annexin A1 all act via FPR2. Functionally, these ligands limit leukocyte recruitment, promote effectorcytosis and stimulate leukocyte egress from sites of inflammation. With such a heterogeneous array of ligands and functional consequences, it is not surprising that genetic deletion of Fpr2 in hyperlipidaemic mice has yielded variable outcomes in the context of atherosclerosis. In intermediate...
and advanced stages of atherosclerosis generated in Apoe<sup>−/−</sup> or Ldlr<sup>−/−</sup> mice, lack of FPR2 yielded either protective effects<sup>162,163</sup> or no effects on lesion size<sup>160</sup>. By contrast, mice lacking FPR2 during the early stages of lesion development generated larger lesions with a higher macrophage and neutrophil content<sup>160</sup>. A possible explanation for such apparent stage-dependent differences may be the imbalance between pro-inflammatory and pro-resolving FPR2 agonists that are present at different stages of lesion formation. As an example, resolvin D1 levels decrease substantially during the advanced stages of atherosclerosis<sup>160</sup>. Chemokine-like receptor 1 (CMKLR1; also known as ChemR23), a class A G-protein-coupled receptor, has structural similarity with the FPRLs. It is widely expressed with higher expression in dendritic cells, monocytes and macrophages. Endothelial cells, vascular SMCs and adipocytes also have high CMKLR1 expression. The CMKLR1 ligand, chemerin, is a chemotactic factor that is secreted as a propeptide. Successive proteolytic cleavages can produce chemerin variants with pro-inflammatory or anti-inflammatory activity, so the effects of chemerin depend on the class of proteases that dominate the microenvironment. Of note, the C15 isoform of chemerin can repolarize macrophages and dampen inflammation<sup>167–169</sup>. Resolvin E1, an anti-inflammatory lipid mediator derived from the ω-3 fatty acid eicosapentaenoic acid (EPA), resolves inflammation by limiting neutrophil recruitment, producing inhibitory cytokines in macrophages and increasing phagocytosis<sup>166–172</sup> through CMKLR1. Two recent studies aimed to understand the role of CMKLR1 in atherosclerosis. Both studies were performed in Apoe<sup>−/−</sup> mice and animals received a diet with a similar fat content for comparable periods of time. However, the study investigating plaque development at an earlier stage reported accelerated atherosclerosis in mice lacking CMKLR1<sup>173</sup>, a finding that contrasts with those in mice developing advanced lesions<sup>173</sup>. As for FPR2, such dichotomous reports on the role of CMKLR1 may result from the relative abundance of pro-inflammatory versus anti-inflammatory CMKLR1 ligands.

Attempts have emerged to increase levels of pro-resolving mediators by exogenous administration<sup>174</sup>. Intraperitoneal administration of resolvin D1 restores lesional resolvin D1 levels, improves lesion stability and inhibits progression of atherosclerosis<sup>166</sup>. Delivery of lipoxin A4 had similar effects<sup>165</sup>. Non-lipid FPR2 activators, such as the N-terminal annexin A1 fragment Ac2-26, can also reduce early atherosclerosis by inhibiting chemokine-mediated leukocyte activation<sup>165</sup>. However, such restoration strategies have limitations, including the unfavourable pharmacokinetics of these agonists as well as their potential side effects when delivered systemically. Ac2-26 encapsulated in collagen-targeted nanoparticles can revert advanced atherosclerosis, and these particles accumulate specifically in the atherosclerotic lesion<sup>164</sup>. To stimulate CMKLR1-driven inflammation resolution, resolvin E1 has been applied topically in the oral cavity of hypercholesterolemic rabbits or via oral gavage in hypercholesterolemic mice. Both treatments limited experimental atherosclerosis development<sup>174,175</sup>. Finally, treatment of mice with maresin 1 and resolvin D2 reduced atheroprogression, halting the expansion of the necrotic core and increasing the thickness of the fibrous cap<sup>176</sup>. These encouraging effects in animal studies raise the question of whether supplementation with ω-3 fatty acids, a precursor of resolving lipid mediators, could potentially increase the generation of resolving lipid mediators and hence counterbalance their decline in advanced lesions. The REDUCE-IT trial showed that a high dose of purified pharmaceutical-grade EPA improved cardiovascular outcomes in patients with high cardiovascular risk and high levels of triglycerides (150–499 mg/dl) treated with statins. Studies with lower doses of EPA or mixtures of ω-3 fatty acids have not yielded positive results<sup>177–180</sup>.

**Efferocytosis: greenkeeping the lesion.** The clearance of dying cells can limit inflammatory responses and maintain tissue homeostasis. Accordingly, efferocytosis — the clearance of dead cells by macrophages — comprises a critical cellular arm of the inflammation resolution machinery. In the context of atherosclerosis, cells that die during the early stages of the disease are swiftly cleared, but the efferocytosis capacity diminishes as lesion development progresses. Macrophages recognize apoptotic cells via receptors and their ligation triggers engulfment. In the context of atherosclerosis, MER proto-oncogene tyrosine kinase (MERTK) has received much attention. In advanced atherosclerosis in humans and mice, the macrophage surface expression of MERTK declines, and the expression of disintegrin and metalloproteinase domain-containing protein 17 (ADAM17), a protease that cleaves MERTK, increases, thus providing one mechanism of impaired efferocytosis in advanced atherosclerosis<sup>181</sup>. Further support for this notion stems from mice expressing cleavage-resistant MERTK, which have lesions with smaller necrotic cores and macrophages with higher efferocytosis capacity<sup>182</sup>. The enzyme calcium/calmodulin-dependent protein...
kinase IIy (CaMKIIy) regulates MERTK levels via a pathway that includes LXRa. Mice lacking CaMKIIy in myeloid cells show increased MERTK expression in lesional macrophages, increased efferocytosis and smaller necrotic cores. Efferocytosis, however, is not just impaired because of changes in the efferocyte — lesional apoptotic cells also acquire strategies to escape clearance. As an example, expression of the ‘do not eat me’ signal CD47 is inappropriately increased on lesional apoptotic cells, rendering them resistant to efferocytosis. Antibody-assisted neutralization of CD47 is one possible way to revert this misdirected inflammatory response.

Numerous nanoparticle-based strategies have emerged to alter the capacity for efferocytosis (details in section Targeted delivery). Alternative strategies may derive from an improved understanding of the metabolic regulation of continued efferocytosis. For example, the metabolism of apoptotic cell-derived arginine and ornithine to putrescine by macrophage arginase 1 and ornithine decarboxylase promotes continued efferocytosis. Consequently, treatment with putrescine promotes atherosclerosis resolution in mice.

Optimizing therapy in time and space
Targeting inflammation systemically may come at the cost of impairing host defences. Optimizing drug delivery in time and space furnishes two options to reduce such unwanted actions.

Chronopharmacology
In circadian medicine, two general strategies may be applicable: targeting the molecular clock or exploiting its rhythmic outputs. Sleep control, exercise and feeding can modulate the molecular clock to influence cardiovascular health. Exploiting circadian outputs for atherosclerosis treatment dates back to the recognition that a critical enzyme that regulates cholesterol synthesis oscillates within a 24-h period. Two decades later, when simvastatin, a short-acting statin, was approved for the treatment of hyperlipidaemia, it was prescribed to be taken "one time per day in the evenings", thus marking the first clinical application of chronopharmacological treatment.

In the past 10 years we have learnt that levels of certain drug targets oscillate over the day and that such oscillations are phase-shifted between organs. Indeed, a recent comprehensive analysis of a high-resolution time series in 64 tissues revealed that 82% of all protein-coding genes oscillate in primates, thus offering possibilities for time-optimized treatment strategies. Indeed, a recent study using the cyclic ordering by periodic structure (CYCLOPS) algorithm revealed that half of the protein-coding genes cycled in at least one of 13 analysed tissues in humans. One thousand of these oscillating genes encode proteins that are drug targets themselves or are involved in drug metabolism (Fig. 3a). Of note, oscillating protein-coding genes include several chemokines, cytokines and adhesion molecules with reported relevance in cardiovascular biology, such as vascular cell adhesion protein 1 (VCAM1), IL-1β, IL-6 and CCL2 (Ref. 186). With regard to CCL2, a recent study has shown that discrete diurnal invasion of the arterial wall driven by the CCL2–CCR2 axis could sustain atherogenic growth.

Indeed, recruitment of neutrophils and monocytes into atherosclerotic lesions oscillates with a peak during the transition from the active to the resting phase, whereas recruitment to microvascular beds peaks 12 h later. Thus, time-optimized pharmacological CCR2 neutralization limited macrovascular myeloid cell recruitment and atheroprogession without disturbing microvascular recruitment. Beyond chemokines, arterial endothelial cell adhesion molecules were also found to oscillate in mice: intercellular adhesion molecule 1 (ICAM1) and VCAM1 peak during the transition from the active to the resting phase. This rhythmic expression depends on oscillatory sympathetic innervation throughout the day; local denervation or systemic administration of a β2-adrenergic receptor antagonist depleted the morning peak of myeloid cell influx into arteries. Thus, two preclinical studies centred on arterial myeloid cell recruitment illustrate the potential of chronopharmacological intervention to alleviate the atherosclerosis burden.

Apart from the oscillation of the target, the pharmacokinetics of the drug can be modulated with chronopharmacology. A recent meta-analysis of 106 clinical trials evaluating time-of-day administration of drugs revealed a time-of-day impact in efficacy and/or toxicity in 75% of the studies. Because of pharmacokinetics, only drugs with short half-lives (typically less than 6 h) should show time-of-day dependency. Yet, in the above-mentioned meta-analysis, a time-of-day dependency was also found for most drugs with plasma half-lives between 8 and 15 h, thus including most drugs used in cardiovascular medicine. These observations demonstrate the enormous potential for timed intervention in cardiovascular diseases.

Targeted delivery
In addition to time-optimized delivery, the delivery of a therapeutic to specific sites may limit potential systemic side effects. Nanoparticle formulations are typically pursued to improve the pharmacokinetics and pharmacodynamics of medicines, for example to increase the plasma half-lives and improve the delivery of drugs with low water solubility. Although nanomedicines are still scarce in the cardiovascular field, several studies have examined nanomedicines in clinical trials, including a second-generation lipid nanoparticle that was used to deliver siRNA that targeted proprotein convertase subtilisin/kexin type 9 (PCSK9) and prednisolone-loaded liposomes. However, nanomedicine’s importance goes beyond improving the pharmacology. Its full potential resides in the potential to couple systemic delivery with homing to specific sites. In the context of atherosclerosis, preclinical strategies have emerged to direct nanoparticles to the fibrous cap, to endothelial cells and to macrophages. As an example, in a study centred on resolving chronic vascular inflammation, nanoparticles smaller than 100 nm were generated to access the inflamed vascular wall by enhanced permeability and retention. The surface of these nanoparticles was decorated with a collagen IV-binding heptapeptide, which was identified from a phage display library. Particles generated in this way accumulated in the lesion shoulder, and using them to deliver Ac2-26 stabilized atherosclerotic...
lesions\(^{164}\) (Fig. 3b). Similar effects were found when these copolymers were loaded with IL-10 (REF.\(^{148}\)). Very recently, the same targeting strategy served to successfully deliver PAD4 inhibitors to eroded arteries in mice, thus inhibiting NET release and reducing vascular damage\(^{189}\). Another recent study used a much simpler approach, employing an employing an immunocytokine to deliver vascular endothelial growth factor C (VEGF-C) to the fibrous cap. In this strategy, VEGF-C was fused to an antibody that recognizes the alternatively spliced extra domain A of fibronectin, a molecular motif expressed in the basement membrane of inflamed vessels\(^{196}\) (Fig. 3b). VEGF-C deposited at the fibrous cap reduced cholesterol overload-induced cell death of SMCs, and hence stabilized atherosclerotic plaques.

Beyond targeting the fibrous cap, strategies have emerged that enable targeting of lesional macrophages. To this end, a combinatorial library of nanoparticles with distinct physical properties and immune cell specificities has been generated and screened in vivo\(^{166}\). This nanoparticle library was based on endogenous HDL particles that incorporate the LXR agonist GW3965, and sought to permit lesional macrophage targeting whilst minimizing known hepatotoxic side effects. In a more recent study, PEGLyated single-walled carbon nanotubes were delivered to atherosclerotic mice and uptake of these nanotubes was reported to occur specifically in lesional macrophages\(^{199}\). Loading these nanotubes with a chemical inhibitor of the antiopagocytic CD47–SIRPα signalling axis reduced plaque burden without the toxic side effects that are typically associated with off-target cell clearance, such as anaemia. Incorporating stabilin 2-targeting peptides in nanoparticles could also target lesional macrophages\(^{199}\). Loading such particles with siRNA to CaMKII\(\gamma\) improved lesional efferocytosis and limited the size of necrotic cores\(^{199}\). Alternatively, HDL-like particles reconstituted from human apolipoprotein A-I and phospholipids can target lesional macrophages. These 22 nm nanoparticles accumulate in lesional macrophages. Loading these cells with inhibitors of CD40 signalling reduced macrophage–driven inflammatory cascades in advanced atherosclerotic lesions\(^{12,14}\). For targeting endothelial cells, αvβ3 integrin targeted nanoparticles loaded with fumagilin were designed to specifically home to neovessels, thus reducing plaque neoangiogenesis\(^{160}\). These studies are some

**Box 3 | Lifestyle modifications**

**Keep moving!** A sedentary lifestyle is associated with augmented cardiovascular risk, whereas regular, moderate physical activity accompanies reduced risk for a future cardiovascular event\(^{101}\). Potential protective effects are multifaceted and likely include metabolic benefits such as improved glucose handling and lowered plasma lipid levels. However, exercise also has distinct anti-inflammatory effects. Regular physical activity lowered circulating counts of mononuclear cells as well as their ability to produce pro-inflammatory cytokines\(^{102}\). Mechanistically, physical activity was associated with decreased intracellular oxygen consumption, thus connecting activity with cellular metabolism. Regular exercise also decreases the number of circulating leukocytes in both mice and humans\(^{125,126}\), an important observation given the stringent connection between circulating leukocyte counts and the risk of future cardiovascular events\(^{131,134}\). Indeed, a recent study shed light on how voluntary exercise dampens haemapoiesis\(^{157}\). Mechanistically, physical activity diminished the production of leptin in adipose tissue, thereby augmenting quiescence-promoting haemapoietic niche factors in leptin receptor-positive stromal bone marrow cells. Importantly, although withdrawal of the running wheel restored leptin levels, leukocyte production remained low and the stem cell transcriptome and epigenome were maintained. Of note, decreased myeloid cell production in this setting did not impact emergency haemapoiesis, which is required to fight an acute bacterial infection.

**You are what you eat!** Reduced calorie intake and improved dietary composition has the potential to prevent primary and secondary cardiovascular events. Current guidelines recommend diets high in fruits, vegetables, whole grains, nuts and legumes. Such food composition closely resembles a Mediterranean diet, and interventional studies, such as the PREDIMED trial (including the revised publication of those results), suggest that intake of a Mediterranean diet can improve cardiovascular health outcomes, including relevant reductions in ischaemic stroke and coronary heart disease\(^{103,104}\). By contrast, sweetened beverages, excessively salty foods (daily salt intake >5 g) and red meat should be avoided\(^{105}\). Consumption of red meat is associated with the progression of atherosclerosis. Mechanistically, bacterial metabolism of dietary L-carnitine, a trimethylamine abundant in red meats, produces trimethylamine N-oxide, a metabolite that raises inflammatory monocyte counts and impairs reverse cholesterol transport\(^{169-176}\).

Although the importance of well-balanced nutrition is established, other forms of nutrition modulation are gaining popularity. Intermittent fasting-related approaches, such as alternate day fasting and time-restricted feeding, have received much attention for their benefits on cardiovascular health\(^{171}\). Accumulating evidence from human studies shows that intermittent fasting generates a beneficial lipid profile\(^{142,194}\) reduces arterial blood pressure\(^{145,146}\) and lowers blood glucose\(^{147,148}\). Intermittent fasting also reduces markers of systemic inflammation and oxidative stress that are typically associated with atherosclerosis\(^{119,124}\). A recent study aimed at connecting short-term fasting and inflammatory responses\(^{126}\). In both humans and mice, short-term fasting induced a drastic drop in circulating monocytes and blunted their inflammatory activity. During the fasting period, monocytes were retained in the bone marrow compartment. Importantly, mice lacking hepatic 5′-AMP-activated protein kinase (AMPK) or peroxisome proliferator-activated receptor-α (PPARα), both of which are important nutrient sensors, failed to respond to fasting in the same way. In addition, short-term fasting drastically reduced plasma levels of C motif chemokine 2 (CCL2), a chemokine known to mobilize monocytes from the bone marrow and recruit them to sites of inflammation\(^{172}\). Thus, this work provides a fascinating link between nutrition, liver metabolism and inflammatory cell trafficking.

Haematopoietic stem and progenitor cells (HSPCs) are functionally different in patients with cardiovascular risk\(^{131}\), and hypercholesterolaemia can imprint persistent changes in bone marrow HSPCs. In competitive bone marrow transplant studies, HSPCs from bone marrow of donors with hypercholesterolaemia exhibited epigenetic modifications in comparison with cells from donors who were normocholesterolemic. Epigenetic differences in donors with hypercholesterolaemia were associated with increased expansion upon bone marrow transplantation\(^{132,133}\). A more recent study confirmed these findings and added a mechanistic angle centred on trained immunity\(^{171}\). In this study, hypercholesterolaemia in Ldlr-deficient mice induced long-lasting epigenetic changes in myeloid progenitors that altered their functionality to be more inflammatory even after termination of high-fat diet feeding. Importantly, such training was lost in mice deficient for NACHT, LRR and PYD domains-containing protein 3 (NLRP3), thus placing NLRP3 as a master switch between metabolic disturbance and myeloid cell output in inflammation\(^{194}\).
of the examples of how nanotechnology can be used to alleviate atherosclerosis. For an overview, we would like to redirect the reader to recent reviews.  

Lifestyle interventions

Although it is generally accepted that a healthy lifestyle reduces cardiovascular risk, its therapeutic application is not as easy as taking a tablet. Large cohort studies have consistently shown that adhering to a healthy lifestyle is associated with reduced cardiovascular risk. As an example, two prospective observational studies revealed an almost 80% reduction in the incidence of myocardial infarction when adhering to a healthy lifestyle. In both studies, a healthy lifestyle was defined as limited alcohol consumption, no smoking, a healthy diet and moderate physical activity. Although the results from these studies are striking, implementation may be difficult on an individual level and underlying mechanisms are not well defined. Moreover, lifestyle interventions can modify genetically determined cardiovascular risk. Of note, many of the lifestyle modifications that are known to alter cardiovascular risk target inflammatory processes (Box 3).

Conclusion and outlook

Only a few years ago, many viewed atherosclerosis as a lipid storage disease, because cholesterol accumulates in plaques. Indeed, a popular formulation of the pathophysiology of atherosclerosis in the 1970s showed bland proliferative lesions containing SMCs and extracellular matrix but not immune cells. The ensuing decades have shone a bright light on inflammation as a common mediator of many risk factors, including levels of LDL and triglyceride-rich lipoproteins, and the altered behaviour of artery wall cells, which beckon leukocytes. These immune cells accumulate within complicated lesions, propagate the disease and produce its complications. The enormous advances in dissecting the cellular and molecular mechanisms of innate and adaptive immunity in atherosclerosis have moved from theory and promise to clinical practice, as evidenced by the success of recent trials in which anti-inflammatory intervention reduced recurrent events, even in patients with excellent control of LDL.

The perception that inflammation replaces traditional risk factors poses a false dichotomy. Rather than displacing classic risk factors such as LDL, the concept of inflammation as a driver of atherosclerosis provides a rich set of pathways by which the traditional drivers can give rise to the disease and its clinical expression. The ample preclinical implications of inflammation in atherosclerosis have opened the door to many new therapeutic targets. The recent demonstration that anti-inflammatory interventions can forestall atherosclerotic complications merely scratches the surface of the potential for developing novel therapeutics. The success of targeting IL-1β highlights the inflammamome pathway as a promising pathway for further therapeutic interventions. For example, the NLRP3 inflammasome as well as the downstream cytokines IL-1β, IL-18 and IL-6 are attractive candidate targets for intervention. Selective neutralization of certain chemokines also merits consideration in the context. The challenge remains to optimize net benefit, as interference with inflammatory pathways can impair host defences.

Promoting resolution without impairing defences furnishes one possible avenue to optimize benefit over risk in novel therapeutics to combat inflammation. Another opportunity lies in targeting the most appropriate therapeutic agent to particular patients. Achieving the goal of personalized intervention or precision therapeutics will require aligning biomarkers to identify susceptible individuals for particular interventions. It is likely that the advent of multi-omic technologies, machine learning and artificial intelligence will help solve the multidimensional problem presented by the myriad mediators and plentiful pathways that are susceptible to therapeutic manipulation. Thus, although we have achieved much, much more remains to be done in harnessing the concept of inflammation in atherosclerosis to ameliorate the disease and forestall its clinical complications. Mining this exciting area promises to unearth further fundamental discoveries, and hasten the translation of research to the clinic to confront the rising worldwide tide of cardiovascular disease.

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**Author contributions**

The authors contributed equally to all aspects of the article.

**Competing interests**

O.S. holds two patents on targeting histones in inflammation and one on disrupting CCL5-HNP1 heteromers, and also receives funding from Novo Nordisk to study chronopharmacological treatment strategies in cardiovascular disease. P.L. is a member of the scientific advisory board for Amgen, Corvidia Therapeutics, DaiCor Pharmaceuticals, Kowa Pharmaceuticals, Olatec Therapeutics, Medimmune, Novartis and XBiotech, Inc. P.L.’s laboratory has received research funding in the past 2 years from Novartis. P.L. is on the Board of Directors of XBiotech, Inc. P.L. has a financial interest in XBiotech, a company developing therapeutic human antibodies. P.L.’s interests were reviewed and are managed by Brigham and Women’s Hospital and Partners HealthCare in accordance with their conflict of interest policies.

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