New insights into immunological aspects of atherosclerosis

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Abstract: Although atherosclerosis was previously thought to be mainly a degenerative disease, it is now well ascertained that its pathogenesis is inflammatory. This review describes the history of a new atherogenetic concept, including the pivotal role of apoE-knockout mice in understanding the inflammatory background of atherosclerosis. There has been lack of unequivocal evidence of an important inflammatory component in atherogenesis. This evidence was delivered by a new technique – gene targeting, for the invention of which Mario R. Capecchi, Martin J. Evans and Oliver Smithies received in 2007 the Nobel Prize in Physiology or Medicine. The pivotal stage of atherogenesis is the antigen presentation by macrophages to T lymphocytes. This antigen could be a fragment of oxidized low-density lipoproteins “digested” by macrophage, heat shock protein 60, β2-glycoprotein I or fragments of bacterial antigens. For interaction between the immunological cells a presence of CD40 receptor on macrophages and its ligand CD40L on the surface of T lymphocytes are necessary. During the interaction between these cells an immunological type T helper 1 (Th1 – cellular) or T helper 2 (Th2 – humoral) response arises. Th1 response and its mediators: interferonγ, tumor necrosis factorα, interleukin-1, interleukin-12 and interleukin-18 enhance atherogenesis, whereas Th2 response and its mediators: interleukin-4, interleukin-5, interleukin-10 and interleukin-13 inhibit the development of atherosclerosis. Atherosclerosis is therefore a chronic inflammatory disease, in most cases initiated by hypercholesterolemia. Nowadays, hypercholesterolemia and inflammation are considered as “partners in crime”. The concept of atherosclerosis as inflammatory disease is fairly new, however, it is already considered as an undisputable achievement of science which have particular therapeutic consequences.

Key words: atherogenesis, atherosclerosis, immunity, inflammation

INTRODUCTION

Studies concerning the pathogenesis of atherosclerosis entered a new phase at the turn of the 21st century. The 20th century was the age of cholesterol and lipoproteins, which has been concluded in a number of clinical studies carried out on a large scale, and they demonstrated unequivocally that normalization of hypercholesterolemia significantly decreased the incidence and mortality of coronary artery disease[1,2]. Nearly to the end of the nineties, atherosclerosis had been assumed to develop as the so-called chronic response to injury (response-to-injury hypothesis) that resulted in the loss of endothelial cells which line the inner side of the vessels[3]. However, other studies showed that the endothelial cells covering the early atheromatous changes were in reality intact[4].

Thus, atherosclerosis had been considered first of all a degenerative disease[5-7]. However, approximately 20 years ago, the trials started to focus to a large extent on another pathogenetic mechanism of atherosclerosis, not considered so far – the inflammatory process.

The first indications

In 1986, with the use of monoclonal antibodies, the small cells with round nucleus present in the atheromatous plaque, known before as “small monocytes”, were demonstrated to be T lymphocytes[8]. Several years later it was shown that these lymphocytes “recognize” as antigens the oxidized molecules of low-density lipoproteins (LDL) – oxLDL[9].

Moreover, the correlation between atherosclerosis and the presence of at least two types of infectious microorganisms; Chlamydia pneumoniae and herpes simplex virus was observed[10,11]. It raised the question if the inflammatory process participate in atherosclerosis. Speculations of this kind were initially received with great scepticism because there was no spectacular and unequivocal evidence of a significant role of inflammation in atherosclerosis.
Additional evidence for the presence of inflammation in atherosclerosis

The newest model of atherosclerosis (described precisely at the end of the paper) enabled the investigators to create apoE-knockout mice, an ideal animal model to test the influence of singular proteins participating in the inflammatory response on the development of atherosclerosis. These studies showed, for example, that the absence of only one cytokine – interferon γ (IFN-γ), reduced atherosclerosis even by 60% [12]. The overexpression of adhesive molecules (vascular adhesion molecule 1 and intercellular adhesion molecule 1) at sites with atheromatous changes was also observed in apoE-knockout mice [13]. Monocyte chemotactic protein was shown to play an important part in the progression of atheromatous lesions [14,15]. Moreover, it was observed that interleukin-18 knockout decreased atherosclerosis by 35% [16,17].

Inhibition of CD40 signaling reduced atherosclerosis [18]. This was explained by the fact that ligation of CD40 molecule (tumor necrosis factor α [TNF-α] receptor superfamily member) – found in the atheromatous plaque, on endothelial cells, vascular smooth muscle cells, antigen-presenting cells, platelets – with CD40L activates a number of transcription factors: NF-κB, AP-1, STAT-1 or Egr-1. Therefore, it influences, for example, the endothelial cell, which, in consequence, acquires proinflammatory and proatherosclerotic phenotype leading to the expression of adhesive molecules and tissue factor on its surface. It creates new possibilities of therapeutic approach, consisting in inhibition of the CD40–CD40L pathway [19-21]. In mice the effect of CD40 is also antagonized by transforming growth factor β [22].

Finally, in apoE-knockout mice with severe combined immunodeficiency (SCID) atherosclerosis was reduced by 70% in comparison to the control group, due to a significantly lower number of lymphocytes in mice with SCID. It was demonstrated that transfer of T cells to these mice aggravated atherosclerosis even by 164% [23].

Atherosclerosis as an inflammatory process

These and other facts made the investigators realize unequivocally that inflammation was essential for atherogenesis. Therefore, in 1999, just before his death, Russell Ross (the author of the previous theory of atherosclerosis as a chronic response to injury) officially proclaimed that atherosclerosis was an inflammatory disease [24].

Whereas the deposition of atheromatous lipids and the accumulation of foam cells – macrophages filled with such lipids – in intima is the main morphological hallmark of atherosclerosis, the more subtle changes in the environment of the arterial wall, stimulated by the influx of inflammatory cells and local release of cytokines and other inflammatory mediators are currently recognized as the crucial causative factors of atherogenesis [25,26].

Inflammation occurs in response to a factor that destabilizes the local homeostasis. The factors that cause Toll-like receptor dependent macrophage activation in the arterial wall include oxLDL, heat shock protein 60 (HSP60) and bacterial toxins [27].

The first stage of atherogenesis consists in endothelial dysfunction [24]. It involves first of all the regions of arterial bifurcations where the blood flow is not laminar. Hence, these localizations are prone to develop atherosclerosis. In such places LDL is stored in the subendothelial space. Low-density lipoprotein accumulation is increased if serum LDL level is elevated. Low-density lipoprotein is transported by passive diffusion and its accumulation in the vascular wall seems to depend on the interaction between apolipoprotein B of the LDL molecule and proteoglycans of the matrix [28].

There is evidence that unchanged LDL are “collected” by the macrophages too slowly to activate their transformation into foam cells. Therefore, it has been suggested that LDL molecule is “modified” in the vascular wall. The most significant modification is lipid oxidation, resulting in the formation of so-called “minimally oxidized” LDL [29]. The generation of these “aliens” for the body molecules leads to the development of inflammatory response, with participation of monocytes and lymphocytes in the first place [30,31].

The inflammation is triggered by accumulation of the minimally oxidized LDLs in the subendothelial space, thus stimulating the endothelial cells to produce a number of proinflammatory molecules [32].

Before the “minimally oxidized” LDL have been phagocytized by the macrophages, they have to be modified into “highly oxidized” LDL. The scavenger receptors are responsible for the rapid uptake of the modified LDL [33].

During the following phase macrophages “present the antigen” to T lymphocytes. This antigen may be a fragment of oxidized LDL “digested” by the macrophages (Fig.), HSP60, β₂-glycoprotein I or the fragments of bacterial antigens [34].

The interaction between the immunological cells requires the presence of CD40 receptor on the surface of macrophages and its ligand CD40L on the surface of T lymphocytes [35,36]. The reciprocal action of these cells produces the immunological response of type T helper 1 (Th1 – cellular) or of type T helper 2 (Th2 – humoral). It is currently believed that the immunological response of Th1 type and its mediators: IFN-γ, TNF-α, interleukin-1, interleukin-12 as well as interleukin-18 accelerate atherosclerosis, whereas the response of Th2 type and its mediators: interleukin-4, interleukin-5, interleukin-10 as well as interleukin-13 inhibit the development of atherosclerosis [37-39]. Therefore, there has arisen an idea of vaccination as a future treatment against atherogenesis [40].

The next phase of atherogenesis is the development of fibrous atheroma. The deposition of extracellular cholesterol
and its esters is then intensified as well as the migration of smooth muscle cells from media to intima, proliferation of these cells and finally production of the extracellular matrix by the smooth muscles cells.

A stable atheromatous plaque is most commonly covered with a fairly thick fibrous layer, protecting the lipid nucleus from contact with the blood. In an unstable plaque there is a big lipid nucleus with a fairly thin fibrous layer. In atheromatous plaque, changed as described above, the proinflammatory factors produced by T lymphocytes (such as IFN-γ) seem to play a crucial role. They decrease production of the extracellular matrix by smooth muscles and at the same time increase production of the metalloproteinases by macrophages [41].

Is atherosclerosis an autoimmunological disease?

The role of HSP60 as an initiator of atherogenesis is currently intensively investigated. Its “molecular mimicry” with HSP of *Chlamydia* has been observed [42]. Moreover, the anti-oxLDL antibodies resemble antiphospholipid antibodies, therefore the concept of atherosclerosis as an autoimmunological disease has been established [34,43,44]. The investigators also emphasize a high pathogenetic similarity of atherosclerosis to rheumatoid arthritis [45].

The newest experimental model of atherosclerosis

Since 1992 the mouse has become an excellent object for the studies on atherosclerosis, replacing the previous animal models [46–48]. Then, the first line of mice with a switched-off gene for apolipoprotein E (apoE-knockout) was developed almost contemporaneously in two laboratories in the United States [49,50]. These mice were soon described as “reliable and useful, the best animal model of atherosclerosis in present times” [51].

During the generation of apoE-knockout mice (known also as apoE null or apoE deficient mice) the normal gene coding apolipoprotein E is replaced by a mutated gene which does not produce this molecule. Such mice are called apoE-knockout because they have a knockout, switched-off, null or inactivated gene coding apolipoprotein E. For clarity, in the following sections of this paper we will use the most popular name: apoE-knockout mice.

The year 1992, in which apoE-knockout mice were invented by a homological exchange of genes, was a real breakthrough year in the studies on the pathogenesis of atherosclerosis [52].

The apoE-knockout mice were formed by homological recombination of embryonic stem cells. The changed cells were implanted into the blastocyst of a mouse of C57BL/6J strain (wild type), which led to the formation of apoE-knockout, homozygous mice in the second generation [53]. The inactivation of the gene coding apoE resulted in the formation of mice with a phenotype with a complete suppression of apoE, but with preservation of fertility and vitality [54]. The apoE-knockout mice, in contrast to all of other animal models, develop atherosclerosis spontaneously, without high-cholesterol diet [55].

The generation of such a model changed the nature of the studies on the pathogenesis of atherosclerosis and enabled the investigators to formulate a new definition of atherosclerosis as a chronic inflammatory disease, in most cases initiated and aggravated by hypercholesterolemia. In the review published in *Nature Medicine* hypercholesterolemia and inflammation were described as “partners in crime” [56].

The inflammatory concept of atherosclerosis has been formulated just in the recent years. However, it is currently an unquestionable achievement of science which also have specific therapeutic implications [57–62].
REFERENCES


