Giant-Cell Arteritis: Immunopathogenic Mechanisms Involved in Vascular Inflammation and Remodeling

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Abstract

Giant-cell arteritis (GCA) is a large-vessel granulomatous vasculitis in which aging, gender and genetics likely play a significant role. The association with polymorphisms in the major histocompatibility complex suggests that GCA may be an antigen-driven disease. Immunopathology studies performed with temporal artery biopsies from patients with GCA have generated relevant clues regarding to pathogenesis by indicating participation of Th1 and Th17-mediated pathways, a prominent role for macrophages in tissue injury, and the relevance of vascular response to inflammation. Vascular wall elements, especially endothelial cells and vascular smooth muscle cells are not passive bystanders. Through expression of chemokines and adhesion molecules vascular cells contribute to the continuous recruitment of inflammatory cells that are able to enter the artery wall through newly formed neovessels. Inflammatory cell products, as well as vascular injury, trigger a vascular remodelling process. This eventually leads to the development of intimal hyperplasia and vascular lumen obliteration, source of ischemic complications.

Key words

Vasculitis; Giant-cell arteritis; Genetic polymorphisms; Inflammation; Cytokines; Growth factors; Angiogenesis; Vascular remodelling.

Introduction

Giant-cell arteritis (GCA) is a granulomatous vasculitis affecting large and medium-sized vessels with a special tropism for the carotid and vertebral arteries [1]. Involvement of the superficial temporal artery is very frequent and performance of temporal artery biopsies is a common diagnostic procedure which facilitates histopathological confirmation of GCA (Figure 1) [2]. Temporal artery biopsies are also a source of valuable tissue not only for diagnostic purposes but also for immunopathology studies which have provided important pathogenetic clues. In fact, the current pathogenesis model is essentially based on the demonstration of particular cell types and subsets in involved tissue or peripheral blood, the expression of activation and differentiation markers by these cells and the production of certain inflammatory molecules in lesions. The role of infiltrating cells and their products is assumed from their known biologic functions and from correlation with relevant histopathological features, clinical phenotypes or disease outcomes [3,4]. However, the pathogenesis of GCA is incompletely understood due to the limited availability of functional studies where the participation of specific pathways can be mechanistically proved.

Predisposing background

Epidemiology surveys indicate that genetic substrate, senescence and gender contribute to GCA pathogenesis [5]. GCA selectively targets aged individuals and predominates in women. However, the role of aging and gender remains virtually unexplored in GCA. Both senescence of the immune system and the arterial tree may be relevant. Recently, senescent T cells (CD4+ CD28−) have been identified in GCA-involved temporal arteries and are found in increased numbers in peripheral blood from patients with several chronic inflammatory diseases, including GCA [6]. Some of these cells express NKG2D
receptors and their ligation induces expression of IFNγ and TNFα, which are relevant cytokines in GCA. Therefore, senescent T cells may have pro-inflammatory responses, facilitating subsequent inflammatory cascades.

GCA largely predominates in Caucasians, particularly in those from Northern European regions or ancestry. This observation, along with the occasionally reported familiar clustering, supports genetic predisposition to develop GCA [7,8]. Polymorphisms in a variety of genes encoding for molecules participating in immune, inflammatory, and vascular responses (i.e., class II major histocompatibility complex (MHC), PTNP22, NOS2, VEGF) are associated with increased GCA risk [8-10].

GCA has been consistently found to be associated with the MHC particularly with class II HLA-DRB1*04 alleles (generally DRB1*0401, but also DRB1*0404) [8,11]. A large-scale genetic survey based on international multi-centre collaboration has recently confirmed that the strongest association identified occurs with variants in the class II MHC and that the resulting risk amino acids are located in the antigen-binding cavity of the HLA molecule [12]. This finding reinforces the role of adaptive immunity and supports the concept that GCA may be an antigen-driven disease.

Initial events: T cell activation and functional differentiation

The nature of the triggering agent or agents has not been consistently identified. Although various microorganisms or viruses have been proposed as causal agents, no definitive causal relationship with a particular microorganism or virus has been demonstrated [13,14]. A variety of microbe and viral sequences have been detected in temporal arteries but none has consistently been associated with GCA [15]. These sequences might contribute to activate pathogen sensing receptors since both innate and adaptive immune mechanisms appear to contribute to GCA [4]. Dendritic cells have been detected in normal or early inflamed large and medium-sized arteries [16-19] and can be activated through toll-like receptors (TLR) to produce chemokines that attract and retain additional dendritic cells [17,18,20]. Activated dendritic cells are able to process and present antigens and express co-stimulatory molecules (CD83 and CD86) for T –cell activation [17,18]. The participation of antigen-specific adaptive immune responses is supported by the demonstration of oligoclonal T cell expansion in lesions [21]. The role of the immunological synapsis and accessory molecules in T –cell activation is supported by interesting observations. On one hand there are several case reports of patients with metastatic melanoma who have developed GCA shortly after receiving ipilimumab (anti –CTLA-4), a form of immunostimulatory treatment favouring CD28-mediated T cell stimulation by neutralizing the inhibitory molecule CTLA-4 [22,23]. Moreover, preliminary results indicate that abatacept (recombinant chimeric Ig-CTLA-4) may be useful for sustaining remission in patients with GCA [24]. After antigen recognition, both Th1 and Th17 differentiation pathways seem to be crucial to the pathogenesis of GCA. IFNγ is markedly and selectively expressed in GCA-involved arteries [25,26] and its functional impact is supported by the expression of many interleukin-induced products in lesions including class II MHC antigens [16], endothelial adhesion molecules [27], inducible nitric oxide synthase [28] and chemokines [26,29,30]. One of the most relevant functions of IFNγ is macrophage activation, granuloma formation and differentiation into giant-cells, all characteristically found in GCA lesions (Figures 1 and 2). Activated macrophages subsequently orchestrate a variety of inflammation amplifying cascades. These are seminal to the development of full-blown transmural inflammatory infiltrates, vascular wall injury and remodelling, which configure the pathologic substrate of the clinical symptoms and complications of GCA [2-4,16,26,28]. In recent years it has become apparent that Th17-mediated mechanisms also contribute to GCA [31-33]. CD161 positive CD4T lymphocytes which are precursors of Th1 and Th17 functional subsets can be identified in inflammatory infiltrates [34]. Cytokines promoting Th17 differentiation such as IL-1, IL-21, TGα, and IL-6 are produced in GCA [25,28,35,36]. Consequently, IL-17A is remarkably expressed in lesions [32]. IL-17A is a highly pro-inflammatory cytokine with pleiotropic effects on a variety of cells including macrophages, neutrophils, endothelial cells and fibroblasts and actively contributes to inflammatory cascades [37]. Both Th1 and Th17 lymphocyte numbers are increased in peripheral blood from patients with GCA where a Th1- Th17 double positive cell population can be also detected [34,35,38]. IL-17A expression rapidly and remarkably decreases with glucocorticoid treatment suggesting that IL-17A suppression may contribute to the dramatic symptomatic improvement that most patients with GCA experience with high dose glucocorticoids [32]. Patients with prominent Th17 response seem to respond better to glucocorticoids and experience less relapses [32]. Consistent with these findings, genes related to T-cell activation and Th1 and Th17 differentiation are hypomethylated in GCA lesions [39].

Regulatory T cells, limiting immune activation and the accompanying inflammatory response are also present in vascular lesions and are decreased in peripheral blood from patients with GCA [32,34]. According to the well-recognized plasticity among T cell subsets, in a strong inflammatory microenvironment, such as GCA lesions, regulatory T cells may not be suppressive and may produce IL-17A [32].

Other cell types contributing to GCA pathogenesis

The role of B cells in GCA has been neglected for years. However, B lymphocytes are present in vascular inflammatory lesions [16,40,41] and may have an even more important role in lymphoid organs. Decreased concentrations of circulating B lymphocytes have been observed in patients with active GCA, which recover with glucocorticoid treatment [40]. Although GCA has been primarily considered a T-cell mediated disease, B-lymphocytes are crucial to T-cell activation. Consistent with the potential participation of B cells, a few reports refer improvement of relapsing patients with B-cell depletion therapy with rituximab [42,43]. A variety of auto-antibodies have been detected in sera from patients with GCA. These include anti-ferritin antibodies, anti-endothelial or anti-vacular smooth muscle cell antibodies recognizing different antigens (i.e., vinculin, annexin V, among others) [44,45]. However, disease specificity, and the consequent diagnostic performance of these antibodies has not been widely validated. More recently, circulating auto-antibodies against 14-3-3, a pleiotropic protein in nucleated cells, have been detected in patients with aortic aneurysm due to large-vessel vasculitis [46]. It is likely that many of these antibodies are generated as a consequence of inflammation and tissue injury rather than having a primary pathogenic role. The biologic significance and diagnostic performance of these auto-antibodies needs further investigation.

Neutrophils are scarce in GCA transmural inflammatory infiltrates but are clearly present in small vessels surrounding the temporal artery and in early inflammatory infiltrates surrounding vasa vasorum [47,48]. Recently, changes in the phenotype of circulating neutrophils have been detected in patients with GCA. In patients treated with high
dose glucocorticoids, neutrophils have lower membrane expression of integrin CD11b, are less adhesive to endothelial cells and are able to suppress T cell proliferation. These abnormalities revert when glucocorticoids are tapered along with the rebound in elevated serum concentrations of IL-8, IL-6 and IL-17 which are able to induce a pro-inflammatory phenotype in neutrophils in vitro [49].

**Amplification cascades**

Following these initiating events, magnifying loops are crucial in the development and progression of transmural inflammatory infiltrates in GCA. Pro-inflammatory macrophages produce cytokines with prominent local and systemic effects and with a strong impact in disease manifestations and outcome. TNFa, IL-1β, IL-6, and IL-33 are expressed in GCA lesions and their expression correlates with the intensity of the systemic inflammatory response, characteristic of GCA [35,50]. Moreover, tissue expression of TNFa and circulating TNFa and IL-6 correlate with relapses and disease persistence [35,51]. Chemokines, endothelial adhesion molecules and colony-stimulating factors are induced or increased in lesions and reinforce inflammatory loops by continuously recruiting and expanding the half-life of additional inflammatory cells [25-27,29]. Angiogenic factors such as VEGF, FGF-2, and PDGFs, among others, are expressed in vascular lesions of GCA and promote new vessel formation [3,52,53]. Acute phase proteins, typically increased in GCA patients, may also be angiogenic [54,55]. Neovessels express endothelial adhesion molecules providing new vascular entries for infiltrating leukocytes and may sustain the active metabolic demands of the inflammatory process [3,27] (Figure 3). The role of neovessels in recruiting phagocytes is supported by the strong expression of MRP8 and MRP12 in surrounding leukocytes. MRP8 is expressed by circulating and freshly recruited phagocytes but expression is lost in tissue macrophages [48]. In addition to favouring the progression of inflammation, angiogenesis may have a protective role by compensating for ischemia at distal sites, and a strong angiogenic response in lesions is associated with lower frequency of neuro-ophthalmic ischemic complications [56,57].

**Arterial damage**

Activated macrophages produce reactive oxygen species which contribute to oxidative damage and vessel wall injury [28]. Proteases may also have an important destructive role. Matrix metalloproteases MMP-9 and MMP-2 have elastolytic activity and are up-regulated in GCA lesions whereas their natural inhibitors TIMP-1 and TIMP-2 are down-regulated yielding an increase in proteolytic balance [28,58]. Increased MMP9/MMP2 proteolytic activity has been demonstrated in lesions and likely contributes to disruption of elastic fibres and abnormal vascular remodelling [58,59] (Figure 2). Disruption of elastic fibres may favour aortic dilatation, an increasingly recognized delayed complication in patients with GCA [59-62].

Currently, the treatment of GCA mainly relies on glucocorticoids which elicit a rapid relief of symptoms but fail to induce sustained remission in 60-70% of patients [63]. Moreover, glucocorticoids are unable to avoid aortic dilatation in 22-30% of affected individuals [59,60]. Understanding the pathogenic mechanisms leading to GCA may foster the identification of better therapeutic agents. The association between increased expression of TNFa and persistent disease activity observed in various studies [35,51] provided support to the performance of clinical trials blocking TNF with infliximab, etanercept or adalimumab which, unfortunately, have proven to be insufficient to abrogate disease activity and maintain remission, presumably due to redundancy in concomitantly activated inflammatory pathways [64-66]. IL-6 is a multifunctional cytokine involved not only in inducing the acute phase response and its systemic manifestations but also in inducing Th17 differentiation and promoting B cell functions. Currently, blocking the IL-6 receptor with tocilizumab is being tested in large international multicenter trials [67-68]. Recently communicated results indicate that tocilizumab may be more effective than placebo in reducing relapses in patients with short exposure to glucocorticoids [68]. IL-1β and IL-17A are additional potential targets. As mentioned, both are potent pro-inflammatory cytokines profusely expressed in GCA lesions. Moreover, IL-1 receptor antagonist deficiency results in large vessel arthritis in a mouse model [69]. Mice deficient in IRF-4 binding protein leading to increased expression of IL-21 and IL-17A develop large-vessel vasculitides as well [70]. Proof of concept pilot studies testing IL-1α and IL-17 antagonism are ongoing (www.clinicaltrials.gov). As mentioned,
interfering with CD28-mediated T-cell co-stimulation with CTLA-4-Ig (abatacept), has been recently reported to reduce relapse rate in patients with GCA in a randomized controlled trial [24]. Therefore, the results from clinical trials with targeted therapies, in addition to expanding the currently reduced therapeutic armamentarium in GCA, provide some proof of concept about the potentially involved immunopathogenic pathways (summarized in Figure 4).

**Figure 4: Summary representation of immunopathogenic mechanisms involved in vascular inflammation and remodelling in giant-cell arteritis.**

**Vascular remodelling and occlusion**

Growth factors produced by activated macrophages or by injured vascular smooth muscle cells trigger a vascular remodelling process leading to myofibroblast differentiation of vascular smooth muscle cells, migration towards the intimal layer and deposition of extracellular matrix proteins (Figures 4 and 5). This results in intimal hyperplasia and vessel occlusion, source of symptoms of vascular insufficiency and ischemic complications in patients with GCA. Several factors including PDGFs, TGFβ and endothelin-1, able to induce myofibroblast activation and production of matrix proteins, are expressed in lesions and might participate in vascular remodelling in GCA [25,71-73]. Blocking PDGF receptor signalling with imatinib mesylate results in reduced myointimal cell outgrowth from cultured temporal arteries from patients with GCA, supporting PDGF contribution to intimal hyperplasia [52]. Increased circulating concentrations of endothelin-1 can be detected in patients with neuro-ophthalmic ischemic complications pointing out a potential role in vasospasm or vascular occlusion [72]. Recently, neurotrophins NGF and BDNF have been shown to be expressed in GCA lesions and promote proliferation and migration of vascular smooth muscle cells. Consequently, their participation in the generation of intimal hyperplasia has been proposed [74]. A number of micro-RNAs regulating vascular smooth muscle cell functions are up-regulated in GCA lesions further supporting their involvement in the generation of intimal hyperplasia [75]. The expression of many of the above mentioned vascular remodelling factors in lesions is not down-regulated by glucocorticoids, suggesting that modulation of their
potential impact in vessel stenosis and occlusion may require a specific approach in large-vessel vasculitis [26,33].

**Figure 5:** PDGF and other growth factors regulate myocardial cell outgrowth from temporal arteries from patients with GCA cultured on tri-dimensional matrix as in reference 52. This may be a useful model to explore mechanisms of vascular remodelling in GCA.

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