Endothelial Dysfunction Induced by Chronic Psychological Stress: A Risk Factor for Atherosclerosis

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Abstract

The exposure to psychological stress can increase the risk of cardiovascular disease due to impaired endothelial function. Mediators such as glucocorticoids, catecholamines, angiotensin II and/or pro-inflammatory cytokines, induced by stress, can contribute to endothelial dysfunction due to increased levels of oxidative stress. The endothelial dysfunction induces reduced expression and/or endothelial nitric oxide synthase enzyme functionality, as well as the impairment of the actions triggered by its metabolite, nitric oxide. Chronic psychological stress leads to atherosclerosis development, which has an endothelial dysfunction at early stages. The oxidative damage and inflammatory mediators, induced by chronic psychological stress, play a key role in this process. Furthermore, chronic psychological stress can contribute to the formation of unstable atherosclerotic lesions as a result of immune system cells accumulation and molecules adhesion, leading to thrombosis and cardiac complications. According to what was stated above, we aim to discuss about the endothelial function impairment mediated by psychological stress, and the involvement of mediators such as glucocorticoids, catecholamines, angiotensin II and pro-inflammatory cytokines in this response. This review covers current advancements to understand how chronic psychological stress could lead to atherosclerosis development.

Keywords: Chronic psychological stress; Oxidative stress; Inflammatory factors; Endothelial dysfunction; Atherosclerosis

Introduction

Chronic exposure to stressful situations is associated with increased risk of cardiovascular diseases [1], including acute myocardial infarction [2], coronary artery disease [3-5], myocardial ischemia [6] and atherosclerosis [7,8]. The odds for acute myocardial infarction have increased more than twice in individuals who were exposed to psychosocial factors associated with exposure to multiple risk factors, such as hypertension, smoking, diabetes mellitus and obesity, in comparison to those who were not exposed to psychosocial factors [2]. The increased risk to acute myocardial infarction triggered by exposure to psychosocial factors was independent of gender, age or geographic distribution [2]. The greatest severity of stress-triggered depressive symptoms was related to reduce long-term survival after myocardial infarction [9]. It has been studied the relation between panic disorder, considered as part of the multifactorial clinical picture of mental stress, and coronary artery disease [10]. Soares-Filho et al. [11] discussed about the endothelial dysfunction and microvascular disease present in mental stress response and suggest that mechanisms from the perspective of endothelial dysfunction and microvascular disease might be involved in mental stress. All these findings suggest that the pathophysiologic effects of psychological stress may trigger the development of cardiac events.

Psychological Stress and Endothelial Dysfunction

Stress is a state of disharmony or threat to homeostasis, caused by psychological, environmental or physiological stressors, which lead to specific or generalized adaptive responses [12]. The biological systems respond to stressful stimuli and attempt to develop an adaptation process to these alterations induced by stressors [13]. An imbalance between stressors and response against it leads to maladaptation, which can be a promoter of various diseases, including cardiovascular diseases [13].

The responses to stressors are initiated by the sensory system stimulation, leading to the activation of several adaptive mechanisms, such as hypothalamic-pituitary-adrenal (HPA) axis and the sympathetic nervous system [14]. Mental stress stimulates the paraventricular nucleus of the hypothalamus, where corticotropin-releasing factor is produced and released, which acts on pituitary in order to release the adrenocorticotropin hormone, resulting in the releasing of glucocorticoids from the adrenal glands cortex [15]. Increased serum levels of glucocorticoids can lead to biological alterations in the body, including the development of some psychological disorders and systemic diseases, such as inflammatory, metabolic and autoimmune diseases [16-19]. Abnormalities in HPA axis functionality have been associated with cardiovascular risk factors [20,21]. Additionally, the sympathetic nervous system activation triggers increased vascular tone, myocardial oxygen consumption, platelets activation, as well as activation of renin-angiotensin system, which results in endothelial dysfunction, and hence, the development of cardiovascular diseases [13].

Endothelial dysfunction is mediated by impaired nitric oxide (NO). The endothelium-derived NO plays a main role in maintaining vascular homeostasis, and it is an important mediator of vasodilation. The biosynthesis of NO results from the oxidation from L-arginine to L-citrulline, catalyzed by the nitric oxide synthase enzyme (NOS),...
in presence of tetrahydrobiopterin cofactor (BH$_4$). There are three
isoforms of NOS, one induced by immunological stimuli (iNOS) and
two of them constitutive, endothelial NOS (eNOS) and neuronal
NOS (nNOS) [22]. After it is produced, NO diffuses from the endothelium
into the vascular smooth muscle and lead to vascular relaxation. This
fact occurs because NO binds to the soluble guanylyl cyclase heme
(sGC) enzyme and produces cyclic guanosine monophosphate (cGMP),
which activates protein kinase (PK) G, triggering a cellular signaling
cascade, which decreases the concentration of cytosolic calcium, and
hence, the vascular relaxation [23].

Considering the importance of NO on vascular homeostasis
regulation, an imbalance between reduced NO bioavailability and/or
increased endothelium-derived contractile factor could lead to
endothelial dysfunction [24]. The lower NO bioavailability can result
from the consumption of NO by reactive oxygen species (ROS),
reduced eNOS expression [25] and lack of eNOS substrate or cofactors
[26]. The latter plays an important role in the enzymatic decoupling,
resulting in ROS production instead of NO.

Oxidative stress is defined as an imbalance between antioxidant
defenses, such as superoxide dismutase (SOD) and catalase, and ROS,
such as superoxide anion (O$_2^-$) and hydrogen peroxide (H$_2$O$_2$) [27].
Many enzymatic systems can produce O$_2^-$, as decoupled NOS [28],
cytochrome oxidase (COX) [29] and nicotinamide-adenine-dinucleotide-
phosphate enzymatic complex (NADPH oxidase). The latter is the
biggest source of ROS in vascular cells [30,31].

Recent studies suggest that psychological stress triggers endothelial
dysfunction [32]. Patients with mild symptoms of depression or stress,
showed high depression or stress score associated to reduced brachial
artery flow-mediated dilation, which suggests the relationship between
stress and endothelial dysfunction [33]. Mice exposed to chronic
unpredictable stress, for 8 weeks, showed impaired endothelium-
dependent relaxation in aorta [34]. The relaxation response, induced
by acetylcholine (ACh), was also significantly reduced in thoracic
aortas from rats exposed to chronic unpredictable stress for 3 weeks
[35]. These results suggest reduced production and/or bioavailability
of NO after stress. Additionally, chronic unpredictable stress increased
the potency of phenylephrine (PE) in thoracic arterial rings from rats,
suggesting that the possible lack of NO bioavailability could contribute
to hyper reactivity of agonists and impair the vascular function [35,36].

According to the previously described findings, the acute restraint
stress reduced vascular and plasma levels of NO metabolites, nitrate
and nitrite, and increased O$_2^-$ levels in rats’ aorta, suggesting endothelial
dysfunction as a result of oxidative stress, induced by stressful
situations [37]. Acute restraint stress also increased the reactivity to
PE in rats’ aorta, and pre-incubation with L-NAME non-selective
NOS inhibitor did not affect this response, suggesting that increased
contraction induced PE in aorta from stressed rats may be due to
impaired modulation of NO [37].

The mechanism in which stress induces endothelial dysfunction
may involve important mediators, including glucocorticoids,
catecholamines, angiotensin (Ang) II and pro-inflammatory cytokines.
Treatment with antagonists of glucocorticoid receptors, mifepristone,
increased the endothelium-dependent relaxation and the eNOS gene
expression in mice’s aorta, both impaired by chronic stress [38].
Mifepristone also reduced ROS vascular levels, which were higher after
stress. These findings suggest that glucocorticoid generation, induced
by exposure to stress, impair endothelial function [38]. The exposure to
dual challenge (physical and psychological stress) increased the levels
of noradrenaline and oxidative stress, as well as the noradrenaline area-
under-the-curve was positively correlated with peak oxidative stress
marker 8-isoprostane, suggesting that the increased noradrenaline
levels may be involved with increased oxidative stress after physical
and psychological stress. This fact could contribute to endothelial
dysfunction [39].

The increased sympathetic activity, induced by stress, contributes to
the activation of the renin-angiotensin system [40,13]. Stress-induced
sympathetic activation of beta-adrenergic receptors in the kidneys
increases renin formation and subsequently, the generation and release
of Ang II into circulation [41]. Plasma renin activity and/or Ang II levels
are increased in subjects exposed to situations of fear, novelty [42] and
to hot temperatures [43]. In rats, the activation of renin-angiotensin
system also increased in response to chronic restraint stress [44,45].
Chies et al. [46] demonstrated that rats exposed to swimming stress
presented reduction in the contractile response induced by Ang II in
thoracic aortas and mesenteric arteries, suggesting that the exposure to
stress alters the vascular reactivity to Ang II.

The increase in Ang II levels, induced by psychological stress,
influences endothelium-dependent relaxation. The chronic restraint stress reduced
endothelium-dependent relaxation, plasma levels of stable NO
metabolite (nitrate and nitrite) and vascular eNOS gene expression,
as well as increased oxidative stress [45]. The treatment with AT$_1$
receptor antagonist, losartan, or angiotensin-converting enzyme inhibitor,
ramipril, blunted these responses, suggesting that the involvement of
AT$_1$ receptors and Ang II in the endothelial dysfunction, induced by
chronic restraint stress [45].

The inflammatory process can also trigger endothelial dysfunction.
Treatment with etanercept, an anti-TNF-$
\alpha$, improved the endothelium-
dependent relaxation and the eNOS protein expression, both impaired
in aorta from rats, after exposure to chronic unpredictable stress [47].
These results show that TNF-$\alpha$ plays an important pathophysiological
role in the endothelial dysfunction induced by stress.

Thus, abnormalities in functionality of HPA axis or increased
sympathetic activity, and hence, increased levels of glucocorticoids
and catecholamines, lead to higher levels of oxidative stress and
inflammation, which are considered risk factors for endothelial
dysfunction. Consequently, these factors may increase the risk of
cardiovascular diseases development, including atherosclerosis [48].

Psychological Stress as a Risk Factor for Atherosclerosis

Atherosclerosis is a chronic disease characterized by formation
of atheroma in the medium and large artery wall. The development
and progression of atherosclerotic lesions are triggered by endothelial
dysfunction, followed by recruitment of leucocytes to the vascular
wall, which modifies the structure of low density lipoproteins (LDL)
deposited on the endothelial layer. Among the LDL modification, the
oxidation induced by ROS results in formation of oxidized LDL and
hence, these modified LDL suffer phagocytosis by macrophages and
accumulate in vascular cells. These alterations lead to formation of
foam cells, which can initiate atherosclerotic lesions [49].

The development of atherosclerosis associated with psychological
stress has been studied since 1983. Chronic psychosocial stress
increased area occupied by intimal lesion in monkeys’ coronary
artery, in the absence of other risk factors for atherosclerosis, such as
alterations in lipid levels, blood pressure or glucose. These findings
suggest that psychosocial stress causes atherosclerotic lesions in non-
human primates without any traditional risk factor [7].

The impaired endothelial function induced by stress triggers
pro-atherogenic effects. Concentration-effect curves showed greater PE potency in endothelium-intact aorta from rats, after exposure to chronic unpredictable stress [36]. The pre-incubation with NOS inhibitor, L-NAME, did not alter this response, suggesting that, the increased PE potency induced by stress can be related to decreased production and/or bioavailability of endothelial NO [36]. These alterations were associated with an increase in the total intima-media thickness of the carotid artery [36], a parameter that indicates a risk factor for cardiovascular disease [50].

Chronic restraint stress has increased the levels of oxidative stress in aortas from apolipoprotein (Apo) E-deficient mice, resulting in increased expression of LOX-1, a receptor to the modified LDL [51]. Treatment with EUK-8, a mimic of SOD and catalase enzymes, reversed the increased LOX-1 expression, suggesting that chronic restraint stress upregulates vascular LOX-1 expression, through a mechanism that involves increased oxygen free radicals generation, which may contribute to the development of pro-atherosclerotic conditions [51]. In addition, mice exposure to chronic restraint stress increased oxidative stress levels in the liver, kidney and heart, and this response was associated with reduced activity of liver antioxidant enzymes [52]. These changes were accompanied by the formation of foam cells and accumulation of lipids in the mice’s aorta, suggesting that chronic stress leads to oxidative stress and also to initiate atherosclerotic lesions [52].

Atherosclerotic lesions are characterized by sub-endothelial retention of inflammatory cells, whose adhesion is mediated by adhesion molecules, such as vascular (VCAM-1) and intercellular cell adhesion molecules (ICAM-1). On the other hand, the leukocyte migration is induced by chemokine expression in the vascular wall, such as monocyte chemotactic protein (MCP-1). The pro-inflammatory cytokines, including TNF-α, interleukin (IL) -1 and IL-6, secreted by activate inflammatory cells, maintain an inflammatory vascular environment and thus, perpetuate atherosclerosis [53].

Chronic stress plays a critical role in the induction of vascular inflammation and progression of atherosclerotic lesions [54-57]. The chronic unpredictable stress increased plasma TNF-α levels and its expression in rabbits’ aorta, as well as infiltration of macrophage and lipid accumulation in the sub-endothelial space. Such characteristics are similar to initial stages of atherosclerosis [57]. Vascular smooth muscle cells treated with serum from stressed rabbits showed increase in the expression of MCP-1, ICAM-1, phosphorylation of mitogen-activated protein kinase (MAPKs), MAPKp38 and c-jun N-terminal kinase (JNK) [57]. The MAPKs signaling pathway regulates the processes of cell differentiation and proliferation, protein expression to leukocyte adhesion and it is involved in the atherosclerotic process. Pro-inflammatory effects induced by chronic stress were partially blocked by anti-TNF-α antibody, MAPKp38 and JNK inhibitors, suggesting that TNF-α, MAPKp38 and JNK activation can play a critical role in the pro-inflammatory and pro-atherogenic effects after chronic stress [57].

The rupture of atherosclerotic plaque is considered one of the main causes of cardiovascular complications [53]. A plate with large lipids pool, a thin fibrous cap and many inflammatory cells is prone to suffer rupture when an external stimulus occurs, such as psychological stress. Chronic unpredictable stress led to a plaque rupture and increased plaque size in brachiocephalic arteries from Apo E-deficient mice, as well as accumulation of inflammatory cells and adhesion molecule in the atherosclerotic lesions. These data suggest that the alterations, mediated by chronic stress on plate composition, can promote instability and contribute to its rupture [54].

The participation of adrenergic receptors in the inflammation, induced by stress-associated atherosclerosis, has been studied. Apo E-deficient mice exposed to social disruption stress showed a correlation between stress-score and median aortic plaque area, as well as increased levels of IL-6 pro-inflammatory cytokine. These observations suggest that psychological stress is involved with the atherosclerosis progression, and the release of IL-6 may be a potential mechanism for the atherosclerosis development [56]. The selective beta1-adrenoceptor blockade with metoprolol prevented the increased IL-6 levels in plasma, induced by social disruption stress, suggesting that the sympathetic nervous system may mediate the release of IL-6 and lead to a progression of atherosclerotic lesions [56]. The inhibition of alpha/beta-adrenergic receptors reduced the lipid deposits, induced by chronic restraint stress, in aortas from Apo E-deficient mice, suggesting that the stress-induced atherosclerotic lesions are associated with activation of adrenergic receptors [58].

Increased activity of the sympathetic nervous system reduced the CXCL12 chemokine expression in the bone marrow from mice, after exposure to chronic unpredictable stress [59]. One of the main functions of this chemokine is to inhibit the migration and proliferation of hematopoietic stem cell, and hold the neutrophils inside the bone marrow. Thus, the reduced expression of CXCL12 led to proliferation of hematopoietic stem cells, and increased production of neutrophils and monocytes. These events resulted in increased release of inflammatory leukocytes into circulation, and hence, the development of inflammatory plaque [59]. The treatment of stressed mice with beta3-selective receptor antagonist, SR 59230A, has limited the progression of atherosclerotic lesions, which supports the idea that the sympathetic nervous system, signaling via beta 3-adrenergic receptors activation, has targeted the CXCL12 chemokine in the bone marrow, and it could be explored as potential therapeutic approach in the stress-induced atherosclerosis [59].

Final Considerations

Psychological stress activates the sympathetic nervous system, renin-angiotensin system and the HPA axis, triggering increased levels of catecholamines, glucocorticoids and Ang II, respectively. All these mediators can promote endothelial dysfunction due to the increase in oxidative stress and circulating inflammatory mediators, as well as reduced NO production or bioavailability. These alterations lead to vascular inflammation and, consequently, the development of atherosclerosis. Furthermore, TNF-alpha and MAP kinases signaling pathway played a potential role in the inflammatory process, as well as the adrenergic receptors and CXCL12 chemokine that are involved with hematopoietic activation and the development of atherosclerotic lesions.

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