Contribution of blood platelets to vascular pathology in Alzheimer’s disease

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Abstract: Cerebral amyloid angiopathy (CAA) is a critical factor in the pathogenesis of Alzheimer’s disease (AD). In the clinical setting, nearly 98% AD patients have CAA, and 75% of these patients are rated as severe CAA. It is characterized by the deposition of the β-amyloid peptide (mainly Aβ40) in the walls of cerebral vessels, which induces the degeneration of vessel wall components, reduces cerebral blood flow, and aggravates cognitive decline. Platelets are anuclear cell fragments from bone marrow megakaryocytes and their function in hemostasis and thrombosis has long been recognized. Recently, increasing evidence suggests that platelet activation can also mediate the onset and development of CAA. First, platelet activation and adhesion to a vessel wall is the initial step of vascular injury. Activated platelets contribute to more than 90% circulating Aβ (mainly Aβ1-40), which in turn activates platelets and results in the vicious cycle of Aβ overproduction in damaged vessel. Second, the uncontrolled activation of platelets leads to a chronic inflammatory reaction by secretion of chemokines (e.g., platelet factor 4 [PF4], regulated upon activation normal T-cell expressed and presumably secreted [RANTES], and macrophage inflammatory protein [MIP-1α], interleukins [IL-1β, IL-7, and IL-8], prostaglandins, and CD40 ligand [CD40L]). The interaction of these biological response modulators with platelets, endothelial cells, and leukocytes establishes a localized inflammatory response that contributes to CAA formation. Finally, activated platelets are the upholder of fibrin clots, which are structurally abnormal and resistant to degradation in the presence of Aβ42. Thus, opinion has emerged that targeting blood platelets may provide a new avenue for anti-AD therapy.

Keywords: cerebral amyloid angiopathy, Aβ40, chronic inflammatory, cerebral vessel

Introduction

Platelets are anuclear cell fragments from bone marrow megakaryocytes and usually survive in the bloodstream of humans for 7–10 days.1 Besides their normal function in hemostasis, platelets play a central role in pathological thrombus formation, which is an important risk factor for Alzheimer’s disease (AD) occurrence.2,3 In addition, many studies have indicated that AD patients have altered platelet function.4 Recent studies have demonstrated a close association between the degree of platelet activation and the progress of AD.4,5 The purpose of this review is to provide new insights that review an association between blood platelets and AD vasculopathy.

Platelet activation and beta-amyloid (Aβ)1-40 overproduction

The accumulation of Aβ peptides (mainly Aβ1-40) as amyloid plaque in cerebral vessels plays an important role in the severity of AD pathology. Platelet activation
and adhesion to a vascular wall is the first step of vascular damage. There are many receptors, enzymes, and signaling molecules involved in this process (Table 1). Among them, thromboxane A₂ (TXA₂) synthesis and thrombin-mediated signaling pathway are regarded as the major events involved in Aβ1-40 overproduction. The initial activation of platelets triggers cyclooxygenase 1 (COX-1)-induced arachidonic acid (AA) metabolism, the result is subsequently converted to TXA₂, which is a potent platelet activator.⁶ Thromboxane receptor α (TPα) is a G-protein-coupled-receptor (GPCR) that is coupled to Gq and G₁₂/₁₃. Binding of TXA₂ with TPα may activate a number of intracellular pathways which enhance primary platelet activation through thrombin (the most potent known platelet activator) or collagen. Protease activated receptor 1 (PAR1) is the major human platelet receptor through which thrombin facilitates the cellular effects of platelet activation without interfering with thrombin-induced cleavage of fibrinogen.⁷

Aβ is a 36–43 amino acid peptide. It is cleaved from the integral membrane amyloid precursor protein (APP) by β and γ-secretases to yield Aβ fragments of various lengths and a smaller C-terminal fragment (CTFγ). Overproduction of Aβ peptides, as well as the failure of their degradation by enzymes, such as neprilysin and insulin degrading enzyme, lead to their oligomerization and aggregation over time to produce senile plaques that are the main neuropathological features of AD. Unlike Aβ1-42 deposited in senile plaques, the circulating Aβ form contributing to perivascular amyloid plaques seen in AD is primarily composed of Aβ1-40, which accounts for 90% of total Aβ.⁶⁻¹¹ APP is found in platelets, and human platelets express all of the enzymes which are required to process APP into Aβ peptide. Platelets are therefore regarded as the main source of circulating Aβ. In human platelets, the major APP isoforms are APP770 and APP75,¹² and they can be hydrolyzed by either α-secretases (non-amyloidogenic pathway) or β-secretases (amyloidogenic pathway) to produce secreted sAPPα and Aβ (mainly Aβ1-40).¹²,¹³ Both sAPPα and Aβ1-40 can be stored in platelet α-granules and released upon platelet activation by thrombin or collagen.¹⁴⁻¹⁷ Once Aβ1-40 is released from the activated platelets, it can in turn activate platelets.¹⁸ As shown in Figure 1A, Aβ1-40-induced platelet activation has been linked to a specific signaling pathway that initiates with the activation of the thrombin receptor PAR1 by Aβ and leads to subsequent activation of p38MAPK (mitogen-activated protein kinases) pathway, which results in the stimulation of cytosolic phospholipase A₂ (cPLA₂) that catalyzes the release of AA for TXA₂ synthesis.¹⁹ TXA₂ augments the activation of platelets and the consequent secretion of Aβ in AD. Platelet activation and Aβ1-40 overproduction may represent a mechanism whereby Aβ1-40 deposition in the walls of cerebral vessels leads to angiopathy occurring in AD.

### Platelet activation and proinflammatory mediator release

Inflammatory reaction plays an important role in the pathogenesis of AD. Platelet activation and consequent degranulation can result in the secretion of numerous biological mediators that are mainly stored in platelet α-granules (Table 2). These mediators include chemokines, such as connective tissue-activating peptide III (CTAP-III), platelet factor 4 (PF4), regulated upon activation normal T-cell expressed and presumably secreted (RANTES), and macrophage inflammatory protein (MIP)-1α, interleukins (IL-1β, IL-7, and IL-8), prostaglandins, and CD40 ligand (CD40L).²⁰ As shown in Figure 1B, platelet-derived mediators enhance leukocyte adherence and endothelial vasoconstriction release of proinflammatory cytokines, which induces cerebral

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Table 1: Major molecules involved in platelet activation

| Platelet ligand | Source | Platelet receptor | Function |
|-----------------|--------|------------------|----------|
| vWF             | Plasma | Gpib-IX-V        | Mediates initial platelet adhesion to damaged vessel walls at high shear flow (>500 s⁻¹) for platelet activation |
| Collagen        |        | Gpvi             | Initiate platelet activation |
| ADP             | Platelet dense granules | P2Y1 or P2Y12 | Provide important positive feedback loop for platelet activation |
| SHT             | Platelet dense granules | SHT2A | |
| TXA₂¹     | COX-1-dependent signaling pathway | TPα | |
| Thrombin² | Coagulation cascade or platelet α-granules | PAR1,4 | |
| Fibrinogen or vWF | Plasma | CD1b3 | Platelet-to-platelet aggregation |

Note: Major pathway involved in Aβ1-40 overproduction.

Abbreviations: ADP, adenosine diphosphate; vWF, von Willebrand factor; GP, glycoprotein; P2Y, purinoceptor; SHT, 5-hydroxytryptamine (also known as serotonin); COX, cyclooxygenase; TXA₂, thromboxane A₂; TPα, thromboxane receptor α; PAR, protease activated receptor; Aβ, beta-amyloid.

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Figure 1 Contribution of blood platelets to vascular pathology in AD. (A) Platelet-originated Aβ40 vicious cycle; (B) inflammatory events initiated by activated platelets; (C) Aβ42-platelet-fibrin clot-induced thrombus formation.

Abbreviations: AD, Alzheimer’s disease; CAA, cerebral amyloid angiopathy; WBC, white blood cell; EC, endothelial cell; APP, amyloid precursor protein; PAR, protease activated receptor; cPLA, cytosolic phospholipase A; TXA, thromboxane A; Aβ, beta-amyloid; TXA2R, thromboxane A2 receptor.

Table 2 Key platelet biological mediators underlying Alzheimer’s disease

| Bioactive mediators | Source | Category | Target cell | Function | Study |
|---------------------|--------|----------|-------------|----------|-------|
| CTAP-III (CXCL7)    | α-granule | Chemokines | Leukocytes | • Enhances neutrophil and monocyte adhesion | Gleissner et al.24 |
|                     |         |          |             | • Promotes neutrophil transendothelial migration | Brandt et al.25 |
|                     |         |          |             | • Cooperates with other cytokines to promote leukocyte adhesion and monocyte differentiation | Hundelshausen et al.26 |
| PF4 (CXCL4)         | α-granule | Chemokines | Leukocytes | • Promotes monocytes adhesion to the endothelial cell | Von Hundelshausen et al.27 |
|                     |         |          |             | • Enhances chemokine synthesis | Baltus et al.28 |
|                     |         |          |             | • Recruits and activates polymorphonuclear leukocytes | Weyrich et al.29 |
| RANTES (CCL5)       | α-granule | Chemokines | Leukocytes and endothelial cells | • Upregulation of leukocyte adhesion molecules (ICAM-1, αvβ3, and MCP-1) | Reichel et al.30 |
| MIP-1 (CCL3)        | α-granule | Chemokines | Leukocytes | • Stimulation of endothelial release of proinflammatory cytokines | Hawrylowicz et al.31 |
| IL-1β                | α-granule | Cytokines | Endothelial cells | • Upregulation of leukocyte adhesion molecules (ICAM-1, VCAM-1, and E- and P-selectin) | Kaplaniski et al.32 |
| CD40L (CD154)       | α-granule and platelet membrane | Cytokines | Endothelial cell CD40 | • Enhances endothelial release of proinflammatory cytokines (IL-1 and TF) | Antoniades et al.33 |
|                     |         |          |             | • Enhances endothelial release of proinflammatory cytokines (IL-1 and TF) | Krocsek et al.34 |
|                     |         |          |             | • Enhances endothelial release of proinflammatory cytokines (IL-1 and TF) | Henn et al.35 |
|                     |         |          |             | • Enhances endothelial release of proinflammatory cytokines (IL-1 and TF) | Heeschen et al.36 |

Abbreviations: CTAP-III, connective tissue-activating peptide III; PF4, platelet factor 4; RANTES, regulated upon activation, normal T-cell expressed and secreted; MIP-1, macrophage inflammatory protein 1; IL-1, interleukin 1; CD40L, CD40 ligand; ICAM-1, intercellular adhesion molecule 1; MCP-1, monocyte chemoattractant protein 1; VCAM-1, vascular cell adhesion molecule 1; CXCL, chemokine (C-X-C motif) ligand; TF, tissue factor; CCL, chemokine (C-C motif) ligand.
amyloid angiopathy (CAA)-related perivascular inflammation or angitis and aggravates the reduction of cerebral blood flow and cognitive decline.21–23

Chemokines
CTAP-III (also known as CXCL7) and PF4 (CXCL4) are chemokines with C-X-C motif, which are the two most abundant CXC chemokines stored in platelet α-granules. CTAP-III can be hydrolyzed by the neutrophil membrane-associated serine-protease cathepsin G and converts into active neutrophil-activating protein (NAP)-2 that induces neutrophil and monocyte adhesion to the endothelium and monocyte transendothelial migration.24–26 Unlike CTAP-III, PF4 needs to synergize with other cytokines, such as RANTES, to promote leukocyte adhesion and monocyte differentiation.27 RANTES is also known as chemokine (C-C motif) ligand 5 (CCL5). It is released from the α-granules of activated platelets and deposited on activated endothelium that promotes monocytes adhesion to the endothelium and chemokine synthesis.28–30 MIP-1α (or CCL3) is another chemokine found in platelet α-granules that involves the recruitment and activation of polymorphonuclear leukocytes.31

Interleukins
IL-1β is a platelet-derived cytokine, playing a key role in platelet-induced endothelial activation.32,33 It induces endothelial cells to secrete IL-6 and IL-8 that results in the upregulation of intercellular adhesion molecule (ICAM)-1, α,β, and monocyte chemotactic protein (MCP)-1, enhancing monocyte and neutrophil adhesion to the endothelium.33–35 Patel et al have characterized the cytokine expression profile in the brain of two transgenic mouse models of AD (TgAPPsw and PS1/APPsw).36 Compared with control littermates, transgenic mice showed a significant increase in the following proinflammatory cytokines: tumor necrosis factor (TNF)-α, IL-6, IL-12p40, IL-1β, IL-1α, and granulocyte-macrophage colony stimulating factor (GM-CSF). The concentrations of these inflammatory cytokines, which are likely derived from activated microglia, correlate with the level of soluble (Aβ1-40) and insoluble (Aβ1-42) forms of Aβ present in the brain. This suggests that pathological accumulation of Aβ is a key driver of the neuroinflammatory response.

CD40L
CD40L, also called CD154, is an important platelet-derived mediator with structural homology to the TNF superfamily.37 Although initially expressed on activated CD4+ T-cells, platelet α-granules are a rich source of CD40L and contribute to more than 95% circulating CD40L.38–40 The binding of CD40L with its endothelium surface receptor CD40 can result in upregulation of leukocyte adhesion molecules (eg, intercellular adhesion molecule-1 [ICAM-1], vascular cell adhesion molecule-1 [VCAM-1], and E- and P-selectin) and stimulation of endothelial release of proinflammatory cytokines (IL-6 and tissue factor).41 Of note, CD40–CD40L interaction is also crucial in pathogenesis of AD. Studies have demonstrated that in human embryonic kidney (HEK)/APPsw, CD40wt, and the CD40-mutant cells, CD40L can increase levels of Aβ (1-40), Aβ (1-42), sAPPβ, sAPPα, and CTFβ. Furthermore, results from CD40L treatment of a neoblastoma cell line overexpressing the C-99 APP fragment suggest that CD40L can increase gamma-secretase activity independently of tumor necrosis factor receptor associated factor (TRAF) signaling. Thus, CD40–CD40L interaction modulates APP processing.

Enhanced platelet activation in AD patients
Platelet activation is essential in hemostasis by forming a hemostatic plug to halt hemorrhage after vascular damage. Recently, increased platelet activation has been found in AD patients (Table 3). Sevush et al reported that platelets of patients with AD exhibit greater unstimulated activation than those of controls. Potential causes of such activation include possible stimulation of platelets by damaged cerebral endothelial cells or platelet activation induced by membrane abnormalities present in platelets of patients with AD. A recent work by Ciabattoni et al showed a continuing potentiation of platelet activation in AD patients, which is relevant to increased lipid peroxidation associated with inadequate levels of vitamin E. Bermejo et al reported an increased platelet level of COX-2 in AD and mild cognitive impairment patients compared with elderly controls, indicating that platelet inflammatory pathways are activated, and that this could be considered an early event in AD development. Iarlori et al reported that higher levels of RANTES were detected in peripheral blood mononuclear cells of AD patients compared with control subjects and AD patients treated with donepezil. Casoli et al found high concentrations of MIP-1α in T-cells and brain microvessels of AD patients. Coated-platelets are a recently described subset of platelets that originate upon dual stimulation of platelets with collagen and thrombin, which represents a highly pro-coagulant subset of activated platelets.46–48 A more recent study showed that elevated coated-platelet levels in patients with amnestic
mild cognitive impairment are associated with increased risk for progression to AD.\textsuperscript{57}

**Epidemiology relevance of AD and stroke**

Platelets have a central role in thrombus formation. At the cellular level, thrombosis is initiated by platelets tethering to subendothelial von Willebrand factor (vWF) via the glycoprotein Ib (GPIb).\textsuperscript{49–52} The adherent platelets become activated and co-aggregate with fibrinogen and vWF via GPIIb-IIIa.\textsuperscript{53–56} At the same time, activated platelets act as a catalytic surface for thrombin generation from its plasma pro-enzymes.\textsuperscript{57} This leads to thrombus stabilization by insoluble fibrin intermeshed within and around the platelet thrombus. The three-dimensional platelet plugs under pathophysiological conditions can obstruct circulatory system patency leading to ischemic heart disease (myocardial infarction and unstable angina), ischemic stroke, and related conditions.

A number of studies suggest that AD patients may have an enhanced potential for thrombosis in the circulation. Purandare et al reported that asymptomatic spontaneous cerebral emboli (SCE) were associated with the concurrent presence of clinically relevant depressive symptoms and the future rapid cognitive decline.\textsuperscript{58,59} Brundel et al found that microinfarcts detected by conventional magnetic resonance imaging are more common in AD patients compared with non-demented controls.\textsuperscript{60} Schenider et al found that subcortical infarcts had an interaction with AD pathology to further worsen working memory.\textsuperscript{61} Thus, stroke may increase the risk of developing dementia.\textsuperscript{62} AD patients in turn demonstrate a greater risk for stroke.\textsuperscript{63} Activated platelets are the upholder of fibrin clots. In vitro and in vivo experiments have demonstrated that fibrin clots are more difficult to degrade in the presence of Aβ\textsuperscript{42,64,65} suggesting a mechanism by which platelets providing common adherent surface of fibrin and Aβ may contribute to enhanced thrombosis (Figure 1C).

**Antiplatelet therapy for AD**

Antiplatelet agents are well established as treatments that can help to prevent strokes.\textsuperscript{66} Aspirin, the most widely used antiplatelet agent, irreversibly inhibits platelet COX-1 activity, leading to reduced synthesis of prostaglandin and TXA2.\textsuperscript{67,68} Long-term aspirin therapy brings about a 20%–25% reduction in the odds of subsequent myocardial infarction, stroke, or vascular death among intermediate- or high-risk cardiovascular disease patients.\textsuperscript{69,70} Recent studies have shown that it is also an effective treatment for AD patients.\textsuperscript{71} In human studies, users of high-dose aspirin had significantly lower prevalence of AD and better-maintained cognitive function than nonusers.\textsuperscript{72} However, aspirin needs to be taken before the symptoms of AD occur. It had no effect on AD at a later stage when the brain damage is severe. These results suggest that repurposing existing antiplatelet drugs for the treatment of AD may be beneficial.

**Conclusion**

Conquering AD remains a major challenge in today's medical research due to the lack of good targets and a limited
understanding of its pathogenesis. This review has highlighted that blood platelets have an important role in AD and CAA. However, there are other mechanisms, apart from platelet activation, that are emerging as important for AD progress. For example, the perivascular drainage hypothesis (ie, blockage of lymphatic drainage of the brain by CAA appears to be a significant factor in the pathogenesis of AD and other dementias and is now widely accepted as an AD risk factor). Nevertheless, the current review has established the concept of developing a different approach to combat AD by targeting blood platelets.

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Disclosure
The authors report no conflicts of interest in this work.

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