Updated Outlook of Cerebral Amyloid Angiopathy and Inflammatory Subtypes: Pathophysiology, Clinical Manifestations, Diagnosis and Management

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Abstract. Cerebral amyloid angiopathy (CAA) is a common untreatable cause of lobar hemorrhages and cognitive decline in the older population. Subset of patients present with its inflammatory subtype with rapid decline in cognitive functions and neurological deficits. Most commonly the underlying pathophysiology of this disease is deposition of insoluble amyloid protein into blood vessel walls which results in vessel fragility leading to local neurotoxicity which may eventually leads to lobar hemorrhages and cognitive decline. The term “Amyloid Spell” encompasses transient focal neurological deficits which is commonly misdiagnosed as seizures or transient ischemic attack in the emergency department. Radiologic findings in these patients may reveal microbleeds, cortical superficial siderosis, white matter hyperintensities, and cerebral edema which support the clinical diagnosis which could be otherwise challenging. CAA diagnostic criteria require CT (Edinburgh Criteria) or MRI imaging, or neuropathology. The diagnosis can be suspected without imaging or neuropathology but cannot be confirmed. This review article provides a critical outlook on different types of presentations, updated diagnostic criteria and management of CAA patients illustrating underlying mechanisms associated with neuronal injury secondary to amyloid deposition.

Keywords: Amyloid beta-related angiitis, amyloid spells, cerebral amyloid angiopathy, cerebral amyloid angiopathy-related inflammation, transient focal neurologic deficits

INTRODUCTION

Amyloidosis is defined as abnormal folding of soluble proteins into insoluble fibrillar protein forms associating as \( \beta \)-pleated sheets, a unique molecular conformation which is resistant to proteolytic degradation. The irreversible insoluble protein state and are responsible for binding affinity to Congo red stain. The main composition of amyloid protein responsible for insoluble properties are serum amyloid-\( \beta \) protein precursor (A\( \beta \)PP) and apolipoprotein E. The extracellular deposition of insoluble amyloid protein can be localized or systemic, which can be toxic to local cells. Amyloid-\( \beta \) protein (A\( \beta \)) is derived from A\( \beta \)PP...
and is localized to the central nervous system (CNS), its wild type is acquired, and its variant is hereditary [1]. Local deposition of Aβ into small to medium vessels of CNS and leptomeninges can lead to cerebral amyloid angiopathy (CAA), which is an untreatable disease mostly of the older population known for lobar cerebral hemorrhages and cognitive decline [2]. CAA is the second leading cause of spontaneous intracerebral hemorrhage (ICH) after hypertension [3]. Another long-term serious manifestation of this disease is disabling cognitive decline. Cerebral amyloid deposition is considered the overlapping cause of Alzheimer’s disease (AD) and cerebrovascular lesions [4]. AD primarily has amyloid deposition in the brain parenchyma in comparison to CAA, where amyloid deposition particularly involves vessel walls. The deposition of amyloid protein in aging brain vessels makes them fragile which contributes to microhemorrhages. CAA can also have acute on chronic presentation in a subset of patients with cerebral amyloid angiopathy-related inflammation (CAARI), or they may present with amyloid spells mimicking transient ischemic attack. Management of each patient depends on the severity and type of symptoms. The purpose of writing this updated review is to discuss cerebral amyloid angiopathy and its various types of presentation in the emergency department.

**EPIDEMIOLOGICAL FEATURES**

**Incidence**

Amyloid protein deposition is a common finding in the aging brain, and its incidence increases with advancing age. Block et al. reported the prevalence of CAA around 30% in the sixth decade, which increases to 50% by the seventh decade [5]. Due to the chronic course of the disease, patients may be asymptomatic until their first presentation with lobar hemorrhage or amyloid spell. CAA-related ICH is seen in 3.8% to 20% of non-traumatic hemorrhages, which makes CAA the second leading cause of spontaneous ICH [4, 6]. Some studies reported amyloid deposition in 50% of lobar hemorrhage [7]. In general, ICH is more common in men, but CAA-related cerebral hemorrhages are more prevalent in women [4, 8, 9]. When compared to AD, CAA is less prevalent in patients without AD [10, 11].

**Risk factors**

Apolipoprotein E is a major cholesterol carrier and helps in lipid transportation. APOE and its polymorphic alleles are the major determinants of AD. Genetic predilection of APOE ε4 is a significant risk factor of CAA [10, 12]. APOE ε4 carriers present with more severe symptoms in comparison to non-ε4 carriers [10]. However, the presence of APOE ε2 is associated with increased risk of ICH [13]. Various other mutated proteins such as prese-nilin 1, α1-antichymotrypsin, neprilysin, low-density lipoprotein-receptor related protein, and angiotensin-converting enzyme genes are associated with CAA [10, 14–16]. These mutated proteins may interact with AβPP to increase its enzymatic cleavage to form insoluble toxic β-pleated amyloid fibrils thus unbalance between the production and clearance of these proteins lead to disease progression. CAA can be present in patients without a history of hypertension, hyperlipidemia, or diabetes mellitus, which probably indicates that it has almost no relation to classic vascular risk factors [10]. However, the presence of hypertension markedly amplifies the risk of hemorrhage, and strict control of arterial blood pressure in patients with probable CAA may reduce the risk of hemorrhage by 77% [5].

**CLASSIFICATION OF CEREBRAL AMYLOID ANGIOPATHY**

There are seven different types of amyloid proteins reported to be associated with CAA. The most widely known amyloid protein associated with CAA is Aβ. Its wild type is seen in sporadic cases, and its variant form is considered hereditary. A missense mutation in the APP gene produces AβPP variants. Different types of variants have been reported, such as Italian, Iowa, and Piedmont variants. However, there are various other protein depositions that have been associated with CAA, such as mutated Cystatin C which is seen in Icelandic patients with familial autosomal dominant hemorrhagic strokes. In Hungarian kindreds, mutated transthyretin protein is reported with familial meningo-cerebrovascular amyloidosis. Mutated prion protein produces mutated glycosphatidylinositol anchor which forms an insoluble amyloid fibril in patients with prion disease.

In 1969, Finnish ophthalmologist Jouko Meretoja described a systemic disease known as Meretoja disease or familial gelsolin amyloidosis due to mutations of the gelsolin gene [17]. Gelsolin is a principal
actin-modulating protein involved in axonal transport, myelination, and neuroprotection. Its mutated variant presents as CAA with corneal lattice dystrophy, cutis laxa, polyneuropathy, and facial palsy [18]. CAA is also linked to a mutation of the *BRI-2* gene, which has been associated with hereditary conditions such as Familial British Dementia and Familial Danish Dementia, which presents with widespread amyloid deposition in the brain parenchyma and vessels [19]. In a Chinese prospective study, high serum levels of neurofilament light chain (NfL) were seen in patients with recurrent intraparenchymal hemorrhage secondary to CAA, and higher serum NfL levels were associated with severity and prognosis of CAA-related ICH. NfL are important for structural stability of axons and for achieving conduction of electrical impulses along the axons and might be released into cerebrospinal fluid (CSF) or blood during ongoing axonal damage. Higher than normal levels have been reported in many other conditions such as frontotemporal dementia, AD, and vascular dementia therefore elevated NfL are not used for diagnosis of any of these disorders (Table 1) [20].

**PATHOPHYSIOLOGY**

CAA involves deposition of insoluble Aβ fibrils in the capillaries, arterioles, small to medium-sized cortical vessels and leptomeninges, and less commonly veins resulting in degenerative vascular changes [21, 22]. The imbalance between excessive production or impaired clearance of amyloid protein plays an important role in disease pathogenesis [22]. The exact origin of Aβ in vessels is still not clear, but its origin from neural tissues and drainage into periarterial interstitial fluid has been proposed. Deposition of amyloid fibrils into periarterial spaces are associated with the disease process [23]. The process of amyloidogenic protein buildup within the brain parenchyma and vessel walls is an area of ongoing investigation.

The highly characteristic microscopic appearance of CAA shows acellular thickening of vessels by an amorphous, intensely eosinophilic material which gives a “smudged” appearance on the light microscope (Fig. 1) [24]. Amyloid deposition is commonly studied using Congo Red stain and viewing histologic sections under polarized light which gives apple-green birefringence [25]. Deposition of amyloid protein into cerebral vessels is a multistep process that includes infiltration of amyloid proteins into media and adventitia of vessels leading to effacement of the affected blood vessels with loss of smooth muscle cells [24]. Deposition of insoluble proteins into the affected blood vessels makes them fragile and is responsible for vasculopathies which include fibrinoid necrosis with onion skin appearance of vessel wall, development of microaneurysms, hyaline degeneration, and perivascular infiltration of lymphocytes [26].

| Type of amyloid protein | Clinical characteristic |
|------------------------|------------------------|
| CAA due to Aβ peptide deposition | Sporadic: associated with advanced age; APOE e4<br>Hereditary: missense mutations in APP gene; Italian (e693k), arctic (e693g), Iowa (d694n), and piedmont (I705v) variants; associated with Down syndrome [2]. |
| CAA due to mutated cystatin c in hereditary cerebral hemorrhage with amyloidosis of Icelandic-type (HCHWA-I) | Autosomal dominant: point mutation at codon 68 of the cystatin c gene located on chromosome 20, associated with fatal strokes in Icelandic patients with familial cerebral hemorrhage secondary to a form of autosomal dominant amyloidosis [3].<br>Mutations of the TTR gene, located on chromosome 18; most common neurological phenotype is familial amyloid sensorimotor polyneuropathy with or without associated autonomic neuropathy; associated with meningoencephalovascular amyloidosis, producing dementia, ataxia, and spasticity in Hungarian kindred [4, 5].<br>Mutations of stop codon 145 in PRNP gene; y145stop or the y163stop variants leads to loss of glycosylphosphatidylinositol (GPI) anchor. In normal cells, GPI interferes with the ability of PRP to form amyloid fibrils in the cerebrovascular system [2, 6].<br>G654a and the G654t mutations of the gelsolin gene located on chromosome 9; ophthalmological (lattice corneal dystrophy), dermatological and neurological symptoms and signs (mild generalized polyneuropathy as well as the involvement of vessels) [7]. |
| CAA in human prion diseases | Mutations of stop codon 145 in PRNP gene; y145stop or the y163stop variants leads to loss of glycosylphosphatidylinositol (GPI) anchor. In normal cells, GPI interferes with the ability of PRP to form amyloid fibrils in the cerebrovascular system [2, 6]. |
| Gelsolin related familial amyloidosis of the Finnish type/Meretoja disease | Mutations of stop codon 145 in PRNP gene; y145stop or the y163stop variants leads to loss of glycosylphosphatidylinositol (GPI) anchor. In normal cells, GPI interferes with the ability of PRP to form amyloid fibrils in the cerebrovascular system [2, 6].<br>G654a and the G654t mutations of the gelsolin gene located on chromosome 9; ophthalmological (lattice corneal dystrophy), dermatological and neurological symptoms and signs (mild generalized polyneuropathy as well as the involvement of vessels) [7]. |
| Hereditary CAA in familial British dementia (FBD) and familial Danish dementia (FDD)-bri2 gene related dementias | A point mutation (T to A) of the normal stop codon of the bri2 gene; mutated proteins deposit in the cerebrovascular system [8]. |
Fig. 1. Amyloid special stain showing vascular amyloid deposition in a cortical leptomeningeal vessel with preservation of some vascular smooth muscle cells, corresponding to CAA grade 1. Note the vascular microaneurysm.

Fig. 2. Leptomeningeal vascular biopsy in patient with suspected CAA showing complete replacement of vascular wall by homogenous eosinophilic amyloid categorized as CAA Grade 2. Confirmed by amyloid-β immunohistochemistry (not shown). Hematoxylin and Eosin, 400x.

CAA can be graded into mild to severe forms based on histopathological features. 1) Mild: amyloid protein is restricted to congophilic rim around smooth muscle fibers. 2) Moderate: tunica media is thicker than normal and is replaced by amyloid protein with no microscopic evidence of leakage of blood. 3) Severe: extensive deposition of amyloid protein with focal wall fragmentation [27]. Fibrinoid necrosis is...
seen more commonly in moderate to severe forms of CAA and has a strong correlation with cerebral hemorrhage (Figs. 1–3) [27, 28].

A subset of patients develops an inflammatory reaction to the amyloid protein known as CAA-associated inflammatory form (CAARI). The presence of perivascular inflammation in these cases may present in various clinical non-specific syndromes such as subacute cognitive decline, seizures, and leukoencephalopathy. Patients without CAARI more commonly presents with lobar hemorrhages [29].

Cerebral amyloid angiopathy and Alzheimer’s disease

The deposition of Aβ into parenchymal cells and vascular tissues occurs independently or may overlap (Fig. 4). Aβ can aggregate into toxic oligomers. Aβ42 has more tendency to aggregate into plaques, while Aβ40 is predominantly seen as amyloid deposition in the walls of bleed vessels. AD is the most common type of dementia seen in elderly people. Underlying pathology starts decades before the first clinical manifestation of AD. People at risk in the preclinical phase of the disease can have an early presence of CSF biomarkers which include low Aβ42 levels and high tau protein level. The risk of progression to clinical AD increases with age and in APOE ε4 carriers.

Various modifiable risk factors such as hypertension, diabetes mellitus, smoking, and physical inactivity have been associated with AD.

CLINICAL FEATURES

Intracerebral hemorrhage

Spontaneous ICH is the most common presentation of rupture of affected vessels in CAA [30]. Aβ deposition has a predilection for cortical and leptomeningeal vessels and rarely for deep structures such as basal ganglia, thalamus, and pons; therefore CAA-related hemorrhage occurs more commonly in peripheral cortical and subcortical lobar locations [11, 30]. Localization of cerebral hemorrhages radiologically may help in the differentiation of CAA-related hemorrhages from hypertensive hemorrhages. Clinical and radiological analysis in SMASH-U classification (Structural lesion, Medication, Amyloid angiopathy, Systemic/other disease, Hypertension, Undetermined) showed that cortical hemorrhages are strongly associated with amyloid deposition compared to hypertension. Deep supratentorial involvement is rare in CAA (5%) as compared to hypertension (83%). In-hospital mortality of CAA-related hemorrhages is reduced (13%) when compared to hypertensive hemorrhages (23%). Study also showed that initial presentation with CAA
is less severe with average NIH stoke scale (NIHSS) scoring 06 as compared to hypertension hemorrhages averaging NIHSS scores of 13. CAA-related hemorrhage is the second leading cause of spontaneous cerebral hemorrhage following chronic hypertension [31]. The risk of clinical signs of CAA increases with age, and it has shown female predominance, particularly in the range of 65 to 74 years of age [11]. The most common manifestations are motor paresis, change in mental status, abnormalities in higher brain functions, visual loss, seizures, sensory symptoms, speech disturbance, or ataxia [32]. Furthermore, an ICH may extend to the subarachnoid space leading to headache and meningeal signs [33]. CAA-associated hemorrhages tend to have less mortality however the recurrence rate is quite common and carries a higher risk of mortality [34].

**Secondary effects of cortical microhemorrhages**

Cortical surface microhemorrhages are commonly seen in the aging brain, and their incidence is associated with an increased risk of neurodegenerative conditions. The distribution of blood oozing from affected capillaries and small vessels is restricted to the immediate area surrounding the leaking vessel. The spatial limitation of blood is attributed to rapid clotting of the ruptured vessel. Anticoagulation increases the risk of extension of intracerebral hematoma by limiting the clotting in ruptured vessels; therefore, anticoagulants of warfarin, direct factor Xa, and direct thrombin inhibitors potentially may increase the severity of the disease [35]. Tissue compression by microhemorrhages is too low to cause ischemia; however, additional injury can be caused by sustained inflammatory response due to exposure of brain parenchyma to blood plasma components which may cause long-term neuronal cell dysregulation or death that underlies the cognitive decline in patients with CAA [35, 36].

**Cortical superficial siderosis**

Cortical superficial siderosis (cSS) is defined as the deposition of blood breakdown products on cortical sulci and subarachnoid spaces. Its clinical presentation is transient focal neurological deficits, often known as amyloid spells [37]. cSS has been reported as an independent risk factor for increased risk of recurrent ICH [38]. Advanced leptomeningeal amyloid disease is associated with cSS, likely due to the rupture of fragile vessels into the subarachnoid space with blood breakdown products eventually entering onto superficial cortical surfaces [39]. On magnetic resonance imaging (MRI), cSS is characterized by low signal intensity over cortical surfaces having a characteristic bi-linear track-like appearance [40].
Amyloid spells or transient focal neurological episodes (TFNE)

TFNE are defined as brief transient stereotypical neurological symptoms that mimic transient ischemic attacks (TIA) or may present as seizure-like activity [41]. TFNEs may occur from seconds to several minutes, often related to cortical superficial siderosis. The exact underlying mechanisms which may initiate these events are not yet clear. The proposed mechanisms include cortical spreading depression or possibly focal seizures resulting from cortical irritation of blood breakdown products which overly the cortical hemispheres [42]. For some patients, a TFNE will be the first clinical presentation of underlying CAA. It is not uncommon to have an initial diagnosis of unspecified seizure or TIA then, after undergoing a thorough clinical exam, assessment of cognitive function, and obtaining advanced radiographic imaging, be advised that the underlying concern is for a diagnosis of possible or probable CAA.

Cerebral amyloid angiopathy-related inflammation

CAARI is a rare form of the disease characterized by an inflammatory response to amyloid deposition in the vascular system. CAARI predominantly affects females with a mean age of 69 years [42]. Neuropathological findings include perivascular inflammatory changes with or without granulomatous formation. The most common clinical presentation in these patients is the subacute onset of cognitive impairment, acute encephalopathy, followed by headache, seizures, hemiparesis, aphasia, and visual symptoms [43]. CSF analysis predominantly shows elevated proteins, nucleated cells greater than 5, normal glucose levels, although pleocytosis is not seen in every patient [42]. However, CSF analysis is neither sensitive nor specific. MRI findings may show symmetric or asymmetric cortical vasogenic edema and subcortical hyperintense T2 lesions [44]. CAARI can be diagnosed from the initial clinical presentation with complimentary MRI findings which provide good sensitivity as well as specificity. Definitive diagnosis requires brain biopsy but that should not delay the treatment in patients suspected to have CAARI. The subset of patients who fail to respond to immunosuppressant therapy may require brain biopsy within 3 weeks [45].

Amyloid beta-related angiitis (ABRA)

ABRA is another rare form of CAA which shares clinical features with CNS vasculitis and is distinct from the inflammatory CAA subtype. This variant presents earlier in life compared to typical CAA, but at a later age in comparison to patients with CNS vasculitis. ABRA is characterized as a perivascular and transmural inflammatory response to amyloid protein, often with granuloma formation and CAARI is characterized by inflammatory reaction around the blood vessels without angiodestruction [46, 47]. Unlike CAA patients, ABRA presents more often with a rapid decline in mental status followed by cognitive impairment, focal neurological deficits, seizures, and behavioral symptoms [48]. CSF commonly shows elevated protein and pleocytosis, findings which are not always present and are not very sensitive. Imaging findings in ABRA may vary from asymmetric white matter changes to leukoencephalopathy [49]. ABRA is an angio-destructive condition which may present with microinfarcts in addition to microhemorrhages and cortical superficial siderosis. Like CAARI, ABRA can also be diagnosed from clinical and MRI findings, obviating the need for brain biopsy; however, brain biopsy is the gold standard that shows granulomatous inflammatory findings [50]. ABRA is noted to have a clear response to immunosuppression (Table 2) [48].

| Pathophysiology | Clinical manifestations |
|-----------------|------------------------|
| Peripheral cortical and subcortical hemorrhages | Most common manifestation is motor paresis, disturbance of consciousness, abnormalities in higher brain functions, visual loss, seizures, sensory symptoms, speech disturbances, or ataxia. |
| Sulcal subarachnoid hemorrhages leading to superficial cortical siderosis | Presents as transient focal neurological episodes (TFNE), aka amyloid spells. |
| Cerebral amyloid angiopathy-related inflammation | Presents as subacute onset of cognitive impairment followed by headache, seizures, hemiparesis, aphasia, and visual symptoms. |
| Amyloid beta-related angiitis (ABRA) | Presents as a rapid decline in mental status followed by focal neurological deficits, seizures, and behavioral symptoms. |
Fig. 5. Radiographic findings in 65-year-old-woman presented with TFNE secondary to CAARI with microhemorrhages. Notes: Axial non-contrast CT scan of the head (a), axial MRI T2-FLAIR (b), and SWI sequence (c) show a pattern of inflammatory CAA with diffuse vasogenic edema involving predominantly the frontal, parietal, and temporal lobes (white arrows) resulting in effacement of adjacent sulci; there are associated multiple foci of petechial microhemorrhages (double arrows) and acute subarachnoid hemorrhage in the left central sulcus (dotted arrow).

**DIAGNOSIS**

Clinical presentation and imaging findings together have good sensitivity and excellent specificity in the initial diagnosis of CAA and related subtypes; however, definitive diagnosis requires a brain biopsy. On initial computed tomography (CT) head, location and size of the hematoma should be noted. Sensitive MR imaging is more helpful than CT head to detect cerebral microhemorrhages. [51] Susceptibility weighted images (SWI) are superior to conventional gradient-echo techniques in diagnosing cerebral microhemorrhages and allow more precise analysis of the natural course of the disease. Brain imaging with SWI requires only additional few minutes and should be included in routine neuroimages (Fig. 5). Formal diagnostic subtracted angiography, or MR angiography, while potentially useful to exclude alternative causes of lobar hemorrhages such as vascular malformations, are not used to diagnose CAA because they do not visualize the small arteries and arterioles that are affected by it [52]. Definite CAA is diagnosed by postmortem autopsy. The modified Boston criteria version 2.0 is now the clinical standard to divide CAA into possible or probable disease by incorporating emerging MRI findings. The addition of multifocality of cSS is one of the main updates of the Boston Criteria Version 2.0. This modification increases sensitivity without lowering specificity (Table 3) [53]. Patients who are very sick in acute settings or in certain contraindications such as non-MRI compatible implants and in low income or middle-income countries where MRI brain might be unavailable, the Edinberg CT and genetic rule in and rule out diagnostic criteria can help the clinicians to make therapeutic decisions and inform prognosis by using three predictors: APOE ε4 possession, subarachnoid hemorrhage, and finger-like projections (Table 4) [54]. Additionally, CAA-associated inflammatory and ABRA need to be recognized, and their clinical presentations with associated radiological findings can help in those diagnoses. The presence of anti-Aβ antibodies in CSF has been reported in CAARI, which further supports the CSF analysis, but routine CSF analysis is not a specific test [55].
Modified Boston criteria Version 2.0 for diagnosis of CAA

| Criteria                                      | Definite CAA                                                                 | Probable CAA with supporting pathology | Probable CAA | Possible CAA                                                                 |
|-----------------------------------------------|------------------------------------------------------------------------------|----------------------------------------|--------------|------------------------------------------------------------------------------|
| Full postmortem examination demonstrating:    | • Lobar, cortical, or cortico-subcortical hemorrhage                         | Clinical data and pathological tissue (evacuated hematoma or cortical biopsy) demonstrating: |              | Clinical data and MRI or CT demonstrating:                                  |
|                                               | • Severe CAA with vasculopathy                                               | • Lobar, cortical, or cortico-subcortical hemorrhage                           |              | • Age ≥ 55 years                                                             |
|                                               | • Absence of another diagnostic lesion                                       | • Some degree of CAA in the specimen                                             |              | • Presentation of spontaneous intracerebral hemorrhage.                    |
|                                               |                                                                              | • Absence of another diagnostic lesion                                           |              | OR                                                                           |
| Probable CAA                                  |                                                                              |                                                        |              | • One lobar hemorrhage with one of the white matter lesions                 |
| Clinical data and pathological tissue (evacuated hematoma or cortical biopsy) demonstrating: |                                                                              |                                                        |              | (severe perivascular spaces in the central semioval or white matter        |
|                                               |                                                                              |                                                        |              | hyperintensities in a multisport pattern)                                   |
|                                               |                                                                              |                                                        |              | • Absence of deep hemorrhagic lesions                                       |
|                                               |                                                                              |                                                        |              | • Absence of other causes of hemorrhage or cSS                              |
| Possible CAA                                  |                                                                              |                                                        |              | OR                                                                           |
| Clinical data and MRI or CT demonstrating:    |                                                                              |                                                        |              | • One lobar hemorrhage (severe perivascular spaces in the central           |
|                                               |                                                                              |                                                        |              | semioval or white matter hyperintensities in a multisport pattern)         |
|                                               |                                                                              |                                                        |              | • Absence of deep hemorrhagic lesions                                       |
|                                               |                                                                              |                                                        |              | • Absence of other causes of hemorrhages                                    |

The Edinburgh CT and genetic diagnostic criteria for lobar intracerebral hemorrhage associated with cerebral amyloid angiopathy

| Probability of moderate to severe CAA | Low | Medium | Severe |
|--------------------------------------|-----|--------|--------|
| Subarachnoid hemorrhage              | –   | +      | –      |
| APOE4 possession                     | –   | –      | +      |
| Finger like projections on CT head   | –   | –      | –      |
| Diagnostic test accuracy             |    |        |        |
| Rule out sensitivity                 |    |        |        |
| 100% (CI 95%)                        |    |        |        |
| Rule in sensitivity                  |    |        |        |
| 96% (CI 95%)                         |    |        |        |

**MANAGEMENT**

*Evaluation and management in the emergency department*

Patients with CAA may present acutely to the emergency department with stroke-like symptoms involving deficits of motor, sensory, cranial nerve, language, speech, coordination, or gait dysfunction. They may present with first-ever or recurrent seizures, acute to subacute cognitive impairment, even encephalopathy or headache. When stroke-like deficits are noted emergency personnel must rapidly screen for possible acute stroke interventions. Initial evaluation with CT head imaging is required to evaluate for spontaneous ICH which, if found, may indicate CAA as CAA-related ICH commonly have lobar location. These unique hemorrhages may
also involve cortical surfaces allowing hemorrhaging blood to extend into the subarachnoid spaces. A thoughtful bedside history will include questions regarding anticoagulant use which raises the risk for rebleeding, and increases the severity of spontaneous hemorrhage, morbidity, and mortality, and requires rapid measures to reverse anticoagulation effect. Sudden neurological deterioration in the emergency department mandates immediate re-imaging to assess for hematoma expansion, with considerations for intubation, transfer to intensive care unit, and neurosurgical consultation. Another challenge faced by emergency physicians is identification of underlying disease in patients presenting with transient neurological deficits which resolves within minutes. When TIA is suspected we recommend CAA be included in the working differential, thus highlighting the importance of admission for a complete stroke workup prior to discharge.

**Intensive monitoring for intracranial spontaneous hemorrhage**

The risk of neurological deterioration from rebleeding is highest within the first 24 hours of ICH. Following initial stabilization, patients should be observed in the intensive care unit with hourly neurological assessments. Blood pressure management is deemed critical to prevent hematoma expansion and blood pressure monitoring with systemic arterial pressure should be considered. American Heart Association recommends lowering systolic blood pressures in the setting of acute ICH to 140 is safe while maintaining mean arterial pressures greater than 65 [56]. Furthermore, critical care management in the emergency department should include monitoring for any change in the neurological examination. This warrants immediate CT imaging of the head to rule out hematoma expansion, worsening cerebral edema with cerebral compression, brain herniation, and development of acute hydrocephalus which may require neurosurgical consultation.

**Management of long-term blood pressure**

Chronic CAA leads to cerebral blood vessel fragility. In the presence of chronic hypertension, there is increased risk for hemorrhage and hemorrhagic recurrence. Long-term management of blood pressure plays an important role in reducing the risk of CAA-related ICH [57]. In general, effective blood pressure control also reduces the risk of all types of ICH. Increases of blood pressure variability significantly above the average baseline has an association with the increased size of cerebral microhemorrhages, particularly in the deep and infratentorial regions [58]. Poor control of blood pressure is associated with increased progression of CAA disease and subsequent mortality.

**Antiplatelet use in patients with CAA**

Current ACC/AHA guidelines recommend against the use of antiplatelets for primary prevention of cardiovascular events in CAA patients due to potential risk for hemorrhage related to CAA disease [59]. However, The REstart or STop Antithrombotics Randomised Trial (RESTART) study has showed no significant difference in ICH incidence between the two groups, those taking antiplatelet therapy for secondary prevention of all major occlusive vascular events such as ischemic stroke, myocardial infarction, and peripheral artery occlusion and those not on antiplatelet therapy. Hence RESTART authors conclude the benefits of using antiplatelets for secondary prevention is greater than the risk of hemorrhage in patients with CAA. Interestingly, all-cause mortality was less in patients on antiplatelet therapy for secondary prevention [60].

**Anticoagulation use in patients with CAA**

**Non-valvular atrial fibrillation**

The use of anticoagulants increases the risk of hematoma expansion and mortality associated with ICH, and there have been no specific randomized controlled trials examining patients with CAA. This places the burden of the decision to use anticoagulants onto the treating physician and family when faced with conditions requiring anticoagulants. Because patients with CAA are at increased risk of recurrent hemorrhages, potential alternatives to anticoagulation should be discussed when anticoagulation is required such as for risk reduction of ischemic stroke secondary to atrial fibrillation. In general, anticoagulants should be avoided in CAA patients if possible; however, emerging FDA-approved alternatives to anticoagulants such as left atrial appendage closure have expanded the non-pharmacological strategies of atrial fibrillation in patients with increased risk of ICH [61].
Valvular heart disease

Patients with mechanical valves require lifelong anticoagulation because of the thrombogenicity of a mechanical valve. The benefit of using anticoagulants to prevent vascular occlusive disease, including in patients with CAA, outweighs the risk of ICH [49]. Patients with a previous history of ICH can be a candidate for bioprosthetic valves; however, the decision of starting anticoagulants in these patients (even short-term anticoagulation) should be a multidisciplinary consideration.

Antiepileptic use in patients with CAA

The CAA patient may present with amyloid spells/TFNE with transient neurological symptoms, or seizures. One of the widely accepted theories for amyloid spells/TFNE is cortical depolarization spreading secondary to hemosiderin following microhemorrhages. Cortical depolarization spreading is also associated with migraine pathophysiology, and various antiepileptics have proven to be beneficial in migraine patients but there is no randomized study examining the effect of antiepileptic medications for the management of TFNE.

Immunosuppression for patients with CAARI and ABRA

A subset of patients may present with an inflammatory form of CAA which is an extremely rare and potentially reversible condition. Treatment relies on high-dose intravenous (IV) pulse steroids followed by steroid tapering for 6 months. One retrospective study showed that the early course of disease and recurrence rate is reduced with early use of high dose immunosuppressant. [42] In this study the typical steroid dosage used was 1 gram IV daily for 3 to 5 days, followed by oral prednisone 60 mg daily for 6 months followed by subsequent tapering.

Cyclophosphamide, mycophenolate, and rituximab are acceptable alternatives, especially for patients with ABRA, as this is severe variant of CAA disease has strong similarities to CNS vasculitis. [62] Despite immunosuppressive therapies, relapse can happen in certain patients. This population may benefit from trials of alternative immunosuppressants.

CONCLUSION

This review article highlights the importance of understanding the different clinical manifestations of CAA in the emergency department, intensive care unit, and in patient clinic follow-up. Considering that TFNE can mimic TIA as well as seizures. Initial diagnosis of CAA can easily be missed, and the inflammatory subtype can be misdiagnosed as tumor if MRI brain with SWI sequencing is not performed early in evaluation. The characteristic MRI radiographic findings of CAA include evidence for prior multiple microhemorrhages, hemosiderosis, and even large lobar hemorrhages. Systolic blood pressure for active CAA-related hemorrhage is recommend by AHA to be 140 mmHg, and we recommend always maintaining mean arterial pressures above 65. Anti-platelet therapy for secondary stroke prevention appears to be safe whereas anticoagulation should be multidisciplinary and other alternatives should be considered if possible. A randomized trial is required to determine the use of anti-epileptics in treating TFNE presenting as seizures. Our current understanding of amyloid deposition into the walls of small to medium cortical vessels is not well-understood. New treatments need to be developed which prevent or decrease amyloid deposition in affected vessels. Studies of genetic factors associated with CAA deposition are ongoing. Factors leading to inflammatory CAA subtype also need further study.

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CONFLICT OF INTEREST

The authors have no conflict of interest to report.

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