Classification of Intracerebral Haemorrhages

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Abstract

Spontaneous intracerebral haemorrhage (ICH) is defined as a collection of blood in the cerebral parenchyma that is not caused by trauma. ICH is the second most frequent cause of stroke, accounting for 10–15 % of all cases in high-income countries and about 20 % in low- to middle-income countries. Despite an apparent stability of incidence over the past decades, the profile of ICH has changed: there are fewer deep ICHs associated with pre-stroke hypertension, whereas the increasing age of the population associated with a more extensive use of antithrombotic drugs leads to an increase of lobar ICH. Deep perforating angio-pathy remains the most important cause of ICH, followed by cerebral amyloid angiopathy, these two aetiologies account for nearly 70 % of all ICH cases. Recent scientific evidence has highlighted new aspects of the pathophysiology of such disorders; nevertheless, the morbidity and mortality of ICH remain extremely high. In the present article, the different causes of ICH will be reviewed.

Keywords

Intracerebral haemorrhage (ICH), deep perforating vasculopathy, cerebral amyloid angiopathy (CAA), cerebral microbleeds (CMBs), brain intracranial vascular malformations, haemorrhagic transformation

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Spontaneous intracerebral haemorrhage (ICH) is defined as a focal collection of blood within the brain parenchyma or ventricular system that is not caused by trauma. 1 It is a heterogeneous condition resulting from several distinct underlying vasculopathies. Several interacting and overlapping risk factors may play a role in the vessel rupture.

The overall incidence of ICH ranges from 15 to 40 per 100,000 person-years. 2–4 ICH accounts for 10–15 % of all strokes, but this proportion may be higher in Asian populations. 5 The risk of ICH increases with age, being 9.6-fold higher in people over 85 years old compared with those less than 45 years of age. 6 ICH incidence is higher in men, especially in Asian populations. 7 Despite a significant improvement in ischaemic stroke management, ICH treatment has not significantly changed and this condition remains associated with a high case fatality rate in the first month, ranging from 13 to 61 % of patients, with a median of 40 % across studies. 8

The clinical and epidemiological scenario of ICH has been changing in the last decades. 9–11 Despite an overall stable incidence of ICH, the incidence among people older than 75 years has increased and the incidence among people younger than 60 years has decreased, with a larger proportion of lobar haemorrhages, suggesting that vasculopathies more strongly associated with the elderly, particularly cerebral amyloid angiopathy (CAA), represent an increasing proportion within the aetiological distribution of ICH. 12 The poor prognosis of ICH may be partly due to our poor understanding of this heterogeneous disease. Herein, we review the different causes of ICH.

Anatomical Distribution (see Figure 1)

ICH location can be classified as deep, lobar and infratentorial (involving the cerebellum and/or the brainstem). The anatomical distribution of the haemorrhage and its extension to other compartments (subarachnoid, subdural, intraventricular) may bring clues to identify the underlying cause of the bleeding.

Our knowledge on the anatomical distribution remains imprecise because most estimates are based on hospital series, which suffer from bias (referral is less often considered for moribund patients or, at the other extreme, for patients with only mild deficits), and population-based studies, which are unbiased might contain a small proportion of haemorrhages precluding any further anatomical subdivision. 11,13 In population-based registries, deep ICH accounts for 60–65 % while lobar ICH accounts for 31–40 % of all ICH cases. 9,10 Multiple ICH accounts from 0.7–4.7 % of all ICH cases.

The pooled 1-year survival estimate in nine population-based studies was 46 % (95 % confidence interval [CI] 43 to 49), while when location is considered the 1-year survival was 45–59 % after lobar ICH, 45 to 59 % after deep ICH and 40 to 54 % after infratentorial ICH. 12

The most frequent cause of deep ICH is deep perforating vasculopathy that supervenes mostly in small perforating arterioles (50–700 µm in diameter) originating from the middle cerebral artery and from the basilary artery, thus explaining the classic location in the basal ganglia.
Figure 1. Deep Perforating Vasculopathy Magnetic Resonance Imaging Findings

(A) Axial slice, T2* Gradient echo, showing right thalamic haemorrhage (white arrow) with intraventricular extension (heads of arrows). (B) Axial slice, T2* Gradient echo showing another characteristic feature of deep perforating vasculopathy: deep cerebral microbleeds located in right external capsule (white arrow). (C) and (D) Axial slices, fluid-attenuated inversion recovery (FLAIR) sequence, showing extensive white matter damage involving subcortical regions, resulting also from deep perforating vasculopathy (white arrows).

and brainstem. Deep ICH may be restricted to brain parenchyma or may extend to the ventricules. Intraventricular haemorrhage (IVH) is a frequent complication occurring in nearly 50 % of ICH patients, and it is a predictor of poor outcome. The risk of bleeding in patients with deep perforating vasculopathy might be enhanced in patients receiving oral anticoagulants and among patients with heavy alcohol consumption history.

Hypertension was found to be twice as common a risk factor for patients with deep ICH than with lobar ICH, particularly in younger age groups (odds ratio [OR] 2.27, 95 % CI 1.94 to 2.66). These findings are, however, heavily influenced by studies with less robust methods. In the methodologically more rigorous studies, a smaller, but still statistically significant, excess of hypertension among patients with deep ICH was found (OR 1.50, 95 % CI 1.09 to 2.07). Further large, methodologically robust studies are needed to determine accurately the relative contribution of hypertension to deep and lobar ICH in different age groups. Although deep perforating vasculopathy is the most frequent cause, other disorders, such as intracranial vascular malformations (IVMs), may less frequently lead to deep ICH.

Lobar ICH can result from several distinct diseases. The most common is CAA. The pathological process seen in CAA occurs in small- to medium-sized leptomeningeal and cortical vessels (especially in the occipital and temporal regions), while vessels in deep areas (thalamus, basal ganglia as well as brainstem) are usually spared. A recent population-based study showed an important increase (~80%) in ICH incidence among people aged >75 years. This result was attributed to a twofold increase in lobar ICH, concomitantly with an observed rise in the premorbid use of antithrombotic drugs at this age. These results suggest that some bleeding-prone vasculopathies in the elderly are more likely to bleed when antithrombotic drugs are used, as illustrated by the rise in the incidence of lobar ICH, in which CAA may be strongly implicated. Intracranial vascular malformations, brain tumours, cerebral venous thrombosis (CVT), haemorrhagic transformation (HT), other vasculopathies and systemic diseases may also lead to lobar ICH.

Diagnostic Work-up

The importance of the diagnostic work-up relies on the identification of the ICH but also of other neuroradiological biomarkers that can give clues to detect the underlying cause of the haemorrhage.

At admission, after the first brain image demonstrating the ICH, it is not always clear whether, how and when to undertake further radiological investigation. In a survey of current practice in three European countries, younger patient age strongly influenced whether further investigation of ICH was performed, followed by the absence of prestroke hypertension and lobar ICH location. There is no general consensus concerning the choice and the timing of neuroradiological work-up for ICH. The American Heart Association/American Stroke Association recommended rapid brain computed tomography (CT) or magnetic resonance imaging (MRI) to distinguish ICH from ischaemic stroke and further vessel examination when structural underlying lesions are suspected. Besides vessel imaging, exploring brain parenchyma with MRI might bring additional clues to the underlying vessel disease.

Brain CT detects symptomatic ICH within minutes of symptom onset but may lack sensitivity if delayed for >1 week after ICH onset. Hyperacute CT angiography followed by a post-contrast scan may identify a ‘spot sign’ – one or more hyperintense spots in the haemorrhage – representing a contrast leak. Its presence might suggest a risk of haematoma expansion, poor outcome and mortality, even if the translation of its value into clinical practice remains controversial. CT-venography is useful to search for a venous thrombosis.

Gradient recalled echo (GRE) T2* sequences identify the ICH soon after onset and reliably detect chronic post-haemorrhagic iron deposits. Conventional sequences may give indirect clues for a diagnosis: old lobar haemorrhages in a context of an acute lobar ICH suggesting CAA; white matter hyperintensities and lacunes of presumed vascular origin in a context of deep ICH suggesting deep perforating vasculopathy; acute lesions in diffusion-weighted imaging (DWI) in other arterial territories suggesting a HT of an infarct. MR-venography is useful to search for a CVT. Intracranial magnetic resonance angiography (MRA) associated with dynamic sequences (4D-MRA) has been evaluated in small studies on arteriovenous malformation (AVM), arteriovenous fistula (AVF) and it is emerging as an alternative to digital subtraction angiography (DSA) in the diagnosis and follow-up of these lesions, even though it may not fully substitute DSA due to limitations in spatial and temporal resolution. In patients with ICH, conventional DSA remains the gold standard to detect underlying vascular lesions (AVM, AVF and arterial aneurysms). This should apply to all patients with ICH, provided they are fit to undergo DSA, or if no other cause is identified.

Imaging data may also be used as part of prognostic scores predicting clinical outcome after acute ICH. After the ascertainment of the haemorrhagic nature of the stroke, a swift diagnosis of the underlying cause can expedite management to improve outcome and/or prevent recurrent ICH.
### Classification of Intracerebral Haemorrhages

#### Causes of Intracerebral Haemorrhage (see Table 1)

There have been some attempts to propose an aetiological classification of ICH, but no consensus has yet been achieved. Of note, the recent Structural lesion, Medication, Amyloid angiopathy, Systemic/other disease, Hypertension, Undetermined (SMASH-U) classification describes categories that are mutually exclusive. In clinical practice, a single ICH may result from several factors. Clinicians should therefore distinguish: (i) predisposing risk factors, such as older age, hypertension, high alcohol intake; and (ii) precipitating factors, such as use of oral anticoagulants.

For example, in an elderly patient with past history of hypertension and myocardial infarction treated with clopidogrel, both hypertension and antithrombotic may contribute to the deep ICH eventually due to deep perforating vasculopathy. Therefore, it may be difficult to establish in a specific patient to which extent these elements interacted. Histological proof should remain the gold standard.

#### Deep Perforating Vasculopathy (see Figure 2)

Deep perforating vasculopathy accounts for nearly 50% of the ICH worldwide. Chronic arterial hypertension is the most frequent risk factor associated with deep perforating vasculopathy. The current understanding of the arterial pathology underlying deep haemorrhage is largely based on studies conducted in the 1960s and 1970s. The proposed pathophysiological basis includes a reactive hyperplasia of vessel wall components followed by microscopic degenerative changes with collagen deposition, leading to reduced vascular reactivity and enhanced vessel wall fragility of small- to-medium penetrating vessels. Deep perforating vasculopathy usually occurs in lenticulostrate arteries originating from middle cerebral artery and perforating arteries that arise from the basilar artery, this explains the classic location of deep ICH in the basal ganglia and brainstem. The hypertensive changes (also known as arteriosclerosis) affecting perforating vessels may lead to lacunar

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### Table 1: Most Frequent Causes of Non-traumatic Intracerebral Haemorrhage

<table>
<thead>
<tr>
<th>Aetiology</th>
<th>Main Features</th>
<th>Associated Radiological Findings</th>
</tr>
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</table>
| Deep perforating vasculopathy | - Hypertension is the most important risk factor  
- Most frequently associated with deep ICH | - Lacunes  
- White matter hyperintensities  
- Deep CMB  
- Recent small subcortical infarcts  
- Brain atrophy |
| CAA | - Age >55 years  
- Lobar ICH without any other detected cause  
- Classified according to the Boston criteria | - Lobar CMB  
- Superficial siderosis  
- White matter hyperintensities  
- Cortical microinfarcts  
- Enlarged perivascular spaces |
| Brain AVM | - Clinical history of seizures and focal neurological deficits may be present  
- DSA remains the gold standard for the diagnosis and follow-up | - CT angiography and MRI with dynamic sequences (4D-MRI) are sensitive diagnostic tools |
| Cerebral CM | - History of previous epilepsy may be present  
- Diagnosis is made by MRI | - Single or multiple cerebral CM  
- Occasionally calcified |
| Dural arteriovenous fistula | - History of pulsatile tinnitus  
- Often is an acquired lesion (thrombotic, traumatic or neoplastic occlusion of a major sinus) | - Meningeal artery supply draining into a sinus or meningeal vein  
- ICH is usually superficial and can be associated to SAH/SDH |
| Haemorrhagic transformation | - Occurs in 15% of patients with cerebral infarction  
- Risk factors of HT: early use/larger doses of anti-thrombotic agents, cardioembolic stroke | - Non-homogeneous haemorrhagic lesion  
- Bleeding within a larger territorial infarct on MRI  
- Location confined to a single arterial territory |
| CVT | - Usually associated with other conditions, such as prothrombotic disorders, cancer, haematological diseases, vasculitis and other inflammatory systemic disorders, pregnancy and puerperium, infections, as well as several local causes, such as brain tumours, AVM, head trauma, CNS infections and infections of the ear, sinus, mouth, face or neck | - Haemorrhage is usually preceded by other neurological signs and symptoms  
- Multiple haemorrhages  
- CT or MR venography are efficient to show thrombosed sinus/cortical vein but MR is preferred because of CT bone artefact |
| RCVS | - More frequent among women  
- Thunderclap headache  
- Differential diagnosis with subarachnoid haemorrhage | - Angiography shows areas of arterial constriction and dilatation (spontaneously reversible within 3 months) |
| Primary or systemic vasculitis | - Presence of laboratorial or radiological features suggestive of vasculitis  
- Increased CSF cell count  
- Diagnosis confirmation requires neuropathological evidence of cerebral vasculitis | - Multifocal white matter lesions  
- Angiographic multifocal segmental occlusion, collateral vessel formation, and prolonged circulation |
| Infective endocarditis | - Fever and changes in white blood cell count  
- Evidence of systemic embolism | - Associated ischaemic lesions  
- Multiple CMB  
- Angiographic evidence of mycotic aneurysm |
| Brain tumours and brain metastases | - Previous history of tumour and/or radiological evidence of other cerebral metastases | - Definitive diagnosis requires histological confirmation of the presence of a brain tumour with spatial relationship with the ICH  
- Contrast CT or MRI suggesting a tumour |

**AVM = arteriovenous malformation; CAA = cerebral amyloid angiopathy; CSF = cerebrospinal fluid; CM = cavernous malformations; CMB = cerebral microbleeds; CNS = central nervous system; CT = computed tomography; CVT = cerebral venous thrombosis; DSA = digital subtraction angiography; ICH = intracerebral haemorrhage; MRI = magnetic resonance imaging; RCVS = reversible cerebral vasoconstriction syndrome; SAH = subarachnoid haemorrhage; SDH = subdural haematoma.**
infarction in some circumstances and to deep ICH in others. Given the absence of anatomical demonstration of perforating arteries in the cerebellum, the implication of deep perforating vasculopathy in the cerebellar ICH remains controversial.

In the absence of prospectively validated diagnostic criteria, clinicians should keep in mind that the presence of risk factors is neither necessary nor sufficient to conclude that a deep ICH is due to arteriolosclerosis. Indeed, the prevalence of hypertension is also very high in patients with lobar ICH.10 Because this vasculopathy has two modalities of expression (haemorrhagic and occlusive), clinicians should search for other radiological markers of the disease, such as the presence of recent small subcortical infarcts, white matter hyperintensities, lacunes of presumed vascular origin, brain atrophy, cerebral microbleeds (CMB) in the basal ganglia and pons.10,36

**Cerebral Amyloid Angiopathy**

CAA-related ICH preferentially affects cortical-subcortical (lobar) regions, less commonly the cerebellum and, rarely, deep or brainstem structures, reflecting the distribution of the underlying microangiopathy.37 The predilection for the occipital lobes20 is not well understood but one hypothesis is that greater tortuosity of occipital small arteries impairs perivascular drainage.38 Clinicopathological studies suggest that CAA-related ICH account for at least 5–20 % of all spontaneous ICH.39 However, there are methodological challenges in attributing ICH to CAA as most pathological case control studies did not systematically control for potential confounding risk factors for CAA, including cognitive impairment, ethnicity or age.37 Advancing age is the strongest known risk factor for developing CAA.40 Apolipoprotein E ε4 and ε2 polymorphisms are both related with increased risk of CAA-related ICH.37 Besides ICH, CAA is also associated with ischaemic damage, such as cortical-subcortical microinfarcts, and white matter hyperintensities. These ischaemic lesions probably result from reduced perfusion by CAA-affected arteries.39

Neuroradiological manifestations of CAA are: strictly cortico-subcortical CMB, convexity subarachnoid haemorrhages (SAHs)/superficial siderosis (SS), white matter hyperintensities and silent small cortical acute ischaemic lesions (see Figure 3). White-matter dilated perivascular spaces could also be an interesting biomarker of CAA. Lobar CMB have been considered as a marker of CAA, as the histological characteristics of the vessels involved in lobar CMB are similar to those of CAA, suggesting that lobar CMB and CAA may be different expressions of the same pathological process.41 Recent data have implicated acute or chronic haemorrhage within or adjacent to the cortical sulci (often described as SS when chronic or as convexity SAH when acute) as another form of bleeding associated with CAA. SS seems also to have clinical meaning in CAA as a trigger of transient focal neurological symptoms and a possible marker of increased risk for future ICH. White matter hyperintensities of presumed vascular origin, visualised on T2-weighted or fluid-attenuated inversion recovery (FLAIR) MRI sequences, are a ubiquitous phenomenon of ageing, but occur with much greater volume in individuals diagnosed with CAA than in those with healthy ageing Alzheimer’s disease or mild cognitive impairment. Microinfarcts can be detected as clinically silent foci of restricted diffusion on DWI sequences.43 Such lesions have been reported in about 10–20 % of patients with CAA imaged either at the time of ICH or during follow-up, suggesting that they might represent a marker of ongoing brain injury related to CAA. Enlarged or dilated perivascular spaces, presumed to be due to the accumulation of interstitial fluid, have been linked to the presence and severity of cerebral small-vessel disease. CAA seems to be
Table 2: Boston Criteria for Diagnosis of Cerebral Amyloid Angiopathy in Patients Suffering a Lobar Intracerebral Haemorrhage

<table>
<thead>
<tr>
<th>Classification of Intracerebral Haemorrhages</th>
<th>Classic Boston Criteria</th>
<th>Modified Boston Criteria</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Definite CAA</td>
<td>Full post-mortem examination demonstrating:</td>
<td>No modification&lt;sup&gt;a&lt;/sup&gt;</td>
<td>The term “haemorrhage” referred, in the pathological validation, to lobar ICH. However, some authors suggest that multiple CMB in cortical or cortico-subcortical regions, without lobar ICH, may be considered as probable CAA.</td>
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<tr>
<td></td>
<td>• Lobar, cortical or cortico-subcortical haemorrhage</td>
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<tr>
<td></td>
<td>• Severe CAA with vasculopathy</td>
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<tr>
<td></td>
<td>• Absence of other diagnostic lesion*</td>
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<td></td>
</tr>
<tr>
<td>Probable CAA with supporting pathology</td>
<td>Clinical data and pathological tissue (evacuated haematoma or cortical biopsy) demonstrating:</td>
<td>No modification&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
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<tr>
<td></td>
<td>• Lobar, cortical or cortico-subcortical haemorrhage</td>
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<tr>
<td></td>
<td>• Some degree of CAA in specimen</td>
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<tr>
<td></td>
<td>• Absence of other cause of haemorrhage*</td>
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<tr>
<td>Probable CAA</td>
<td>Clinical data and MRI or CT demonstrating:</td>
<td>Clinical data and MRI or CT demonstrating:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Multiple haemorrhages restricted to lobar, cortical or cortico-subcortical regions (cerebellar haemorrhage allowed)</td>
<td>• Multiple haemorrhages restricted to lobar, cortical or cortico-subcortical regions (cerebellar haemorrhage allowed) or</td>
<td></td>
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<tr>
<td></td>
<td>• Age ≥55 years</td>
<td>• Single lobar, cortical or cortico-subcortical haemorrhage and focal&lt;sup&gt;b&lt;/sup&gt; or disseminated&lt;sup&gt;c&lt;/sup&gt; superficial siderosis</td>
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<tr>
<td></td>
<td>• Absence of other cause of haemorrhage*</td>
<td>• Age ≥55 years</td>
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<tr>
<td></td>
<td></td>
<td>• Absence of other cause of haemorrhage or superficial siderosis*</td>
<td></td>
</tr>
<tr>
<td>Possible CAA</td>
<td>Clinical data and MRI or CT demonstrating:</td>
<td>Clinical data and MRI or CT demonstrating:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Single lobar, cortical or cortico-subcortical haemorrhage</td>
<td>• Single lobar, cortical or cortico-subcortical haemorrhage or focal&lt;sup&gt;b&lt;/sup&gt; or disseminated&lt;sup&gt;c&lt;/sup&gt; superficial siderosis</td>
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<td></td>
<td>• Absence of other cause of haemorrhage*</td>
<td>• Absence of other cause of haemorrhage or superficial siderosis*</td>
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</table>

Adapted from Knudsen et al. 2001 and Linn et al. 2010.43,45 For absence of other causes/other diagnostic lesion, a complete diagnostic workup aimed at excluding in the acute phase all the causes of symptomatic lobar haemorrhage (cf Table 1), magnetic resonance imaging (MRI) at 6 months follow-up and in selected cases digital subtraction angiography (DSA) in the acute phase/follow-up are necessary.48 Other causes of intracerebral haemorrhage (ICH) that question the diagnosis of cerebral amyloid angiopathy (CAA) are: excessive warfarin dose (international normalised ratio [INR] >3.0) (INR 3.0 or other non-specific laboratory abnormalities are permitted for diagnosis of possible CAA); antecedent of head trauma or ischaemic stroke; haemorrhagic transformation of an ischaemic stroke; arteriovenous malformation; central nervous system (CNS) tumour, vascular malformation or vasculitis; and blood dyscrasia or coagulopathy. The value of 3 remains controversial.45 *No modification compared with the classic Boston criteria; *Siderosis restricted to three or fewer sulci; *Siderosis affecting at least four sulci. CT = computed tomography.

preferentially associated with high numbers of visible perivascular spaces in the centrum semiovale.44

The Boston criteria<sup>a</sup> (see Table 2) represented an effort to estimate the likelihood of the presence of CAA in patients with ICH with categories of probable and possible CAA based on the pattern and number of haemorrhagic lesions on neuroimaging.45 These criteria are widely used but a few limitations have to be highlighted.46 In the original version, the authors took into account only patients with symptomatic lobar ICH, not considering other CAA clinical manifestations, such as cognitive decline or focal transient neurological symptoms. Furthermore, they did not validate the impact of other MRI biomarkers of CAA, such as the presence of strictly cortico-subcortical CMB or SS. Recent scientific evidence suggests that taking into account strictly lobar CMB and/or SS together with the presence of symptomatic lobar ICH would allow to increase specificity and sensitivity of the Boston Criteria, thus identifying a higher number of patients suffering from CAA.47,48 The sensitivity and specificity of SS and lobar CMB for CAA have to be validated in larger prospective cohorts.

Intracranial Vascular Malformations

Several types of IVMs may cause ICH. Overall, patients with ICH due to vascular malformations have a better prognosis than patients with spontaneous ICH.49 In the Scottish population-based registry SIVMS, the crude detection rate (per 100,000 adults per year) was 2.27 for all IVMs, 1.12 for brain AVMs, 0.56 for cavernous malformations, 0.43 for venous malformations and 0.16 for dural AVMs.50

Arteriovenous Malformations

AVMs consist of a pial artery supply collected into a nidus and then on a draining vein. Half of the AVMs are revealed by an ICH.51 Unruptured AVMs are associated with an annual haemorrhage rate of 2.3–2.8 %.51,52 Initial haemorrhagic AVM presentation, female sex and increasing age are major risk of recurrent ICH. Some anatomical features, such as deep brain location or exclusive deep venous drainage, might also be considered as markers of haemorrhagic risk.51

Cerebral Cavernous Malformations

CCM are angiographically occult vascular malformations, consisting of blood vessels devoided of muscular and elastic tissue that are lined with endothelial cells without intervening brain tissue. They can be single or multiple and occasionally calcified.53 The risk of ICH is around 6 % per person-year.54 ICH due to CCM occurs at a younger age and tends to be less disabling at onset than those due to other cerebral AVMs. Furthermore, CCM tend to cause small ICHs thus explaining a mild severity. CCM-related ICHs rarely have an extension to other compartments (subarachnoid, subdural or intraventricular) unlike half of the ICHs due to AVM and AVF.55
Haemorrhagic Transformation

HT occurs in some patients with cerebral infarction (up to 15 % in the acute phase of stroke, depending on patient selection and radiological criteria). HT may be distinguished from a spontaneous ICH by the lack of homogeneity of the haemorrhagic lesion that lies within an area of infarction (appearing as an area of low density on CT or of hyperintensity in DWI sequences on MRI) with a typical location confined to a single arterial territory. Sometimes the haemorrhage is so dense that it would have been regarded as a so-called ‘primary’ ICH, had not an earlier CT scan or MRI in the acute phase shown an infarct (see Figure 4A). The diagnostic challenge comes from the fact that in some patients the first radiological exam already shows blood and the underlying ischaemic process might be difficult to disentangle (see Figure 4B). An important diagnostic tool in this case is the DWI sequence that enables concomitant ischaemic lesions in the same or in other arterial territories to be disclosed. It is also important to explore intracranial vessels to search for an arterial occlusion (see Figure 4D).

Systemic Disease and Brain Tumours/Metastases

ICH can result from systemic conditions, such as liver diseases with abnormal coagulation, haematological malignancies and other haematological conditions with reduced platelet count, systemic infections, infective endocarditis and systemic vasculitis. Brain tumours may potentially bleed and represent 4.4 % of all ICH. Glioblastoma multiforme, oligodendroglioma and brain metastases from melanoma, renal cell carcinoma, thyroid carcinoma and choriocarcinoma (tumours associated with the higher risk of tumour associated-ICH). Intra-tumour neovascularisation, fragile vessels within tumour tissue and immature cell-to-cell junctions in tumour vasculature are reported as the mechanisms of tumour-associated ICH.

Cerebral Venous Thrombosis

CVT is a rare type of cerebrovascular disease that accounts for 0.5 % of all cases of stroke. CVT presents with a wide spectrum of signs and modes of onset, thus mimicking numerous other disorders. The most common symptoms and signs are unusual headache (which occurs in nearly 80 % of the patients), seizures, focal neurological deficits, altered consciousness and papillary oedema, which can present in isolation or in association with other symptoms. In about 39 % of cases, the CVT is associated with an ICH. Haemorrhages result from venous infarcts.

Several disorders can cause or predispose patients to CVT. These disorders include all medical and surgical causes of deep vein thrombosis, genetic and acquired prothrombotic disorders, cancer, haematological diseases, vasculitis and other inflammatory systemic disorders, pregnancy and puerperium, infections, as well as several local causes – such as brain tumours, AVMs, head trauma, central nervous system (CNS) infections and infections of the ear, sinus, mouth, face or neck. Diagnostic and therapeutic procedures such as lumbar puncture, jugular catheter and some drugs – in particular oral contraceptives, hormonal replacement therapy, steroids and oncology treatments – can also cause or predispose people to CVT.

CVT is typically multifactorial, which means that the identification of a risk factor or even of a cause should not deter clinicians from looking for other risk factors, particularly congenital thrombophilia. The diagnosis can be suspected when the location of the ICH is: parasagittal (mono or bilateral) with thrombosis of the superior sagittal sinus, on the cerebral convexity (when the thrombosis involves a cortical vein) or in the temporal lobes with thrombosis of the lateral sinus. Extensive haemorrhages occur in only a minority of patients. CVT should be suspected especially when: (i) the bleeding is in a lobar location, (ii) the haemorrhage has been preceded by other signs and symptoms and (iii) the haemorrhage is multiple (mainly in the parasagittal or temporal regions).

In case of CVT, even in the presence of massive ICH, there is the indication of full-dose anticoagulation as first-line therapy.

Reversible Cerebral Vasoconstriction Syndrome

Reversible cerebral vasoconstriction syndrome (RCVS) is a rare condition that is characterised by the association of severe headaches (with or without additional neurological symptoms) and constriction of cerebral arteries that resolves spontaneously in 1 to 3 months. This disturbance may be spontaneous, while 25–60 % of cases are secondary, mostly to exposure to vasoactive sympathomimetic or serotoninergic substances, and/or to the postpartum state. The most common clinical feature is a severe acute headache (‘thunderclap’ headache) and the major complications are focal cortical SAHs (20–25 %) and ischaemic or haemorrhagic strokes (5–10 %). Parenchymal haemorrhages, which occurred in nearly 12 % of all RCVS patients, are of variable volume, more frequently single than multiple, and lobar than deep and more often associated with another type of stroke (convexity haemorrhage or infarction, or both) than isolated. They occur early in the course of RCVS and are revealed mostly by a persisting focal deficit concomitant with thunderclap headache. However, causal relationship remains controversial.

Conclusion

In 2014, ICH remains a devastating disorder. There is no such thing as a ‘primary’ ICH like there is no primary ischaemic stroke. Disentangling the underlying vessel disease that eventually led to the bleeding might help tailoring new treatment strategies and preventing recurrent events.