Imaging of intracranial haemorrhage

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Intracranial haemorrhage can be a devastating disorder that requires rapid diagnosis and management. Neuroimaging studies are not only required for diagnosis but also provide important insights into the type of haemorrhage, the underlying aetiology, and the accompanying pathophysiology. Historically, CT has been the diagnostic imaging study of choice; however, there is a growing body of data that suggest that MRI is at least as sensitive as CT to detect haemorrhage in the hyperacute setting, and superior to CT in the subacute and chronic settings. Blood has characteristic appearances on both imaging modalities at each stage (acute, subacute, and chronic) and it is important that physicians are familiar with the appearance of various types of intracranial haemorrhage on CT and MRI and their clinical implications. In addition, new imaging applications, such as magnetic resonance spectroscopy and diffusion tensor imaging, are promising research techniques that have the potential to enhance our understanding of the tissue injury and recovery that result from intracranial haemorrhage.

Introduction

Intracranial haemorrhage is a devastating disorder that has poor prognoses and high mortality rates. Advances in neuroimaging techniques have improved our diagnostic capabilities, increased our understanding of the underlying pathophysiology and aetiology of intracranial haemorrhage, and helped to establish prognoses. For example, rapid multimodal CT and MRI can now be done in real-time in the acute clinical setting, which enables prompt diagnosis of haemorrhage on parenchymal imaging and can also show vessel malformations and other underlying structural lesions on additional sequences. In particular, the introduction of blood-sensitive gradient echo sequences has enabled MRI to become a clinically useful technique to detect acute haemorrhage. A classification of the major types of adult intracranial haemorrhage is provided (panel 1).

First, we review the neuroimaging characteristics and development of signal change in response to blood breakdown products seen on CT and MRI at various stages. Then, the appearance of haemorrhage on CT and MRI sequences will be reviewed by individual haemorrhage types (intraparenchymal haemorrhage, subarachnoid haemorrhage, subdural haematoma, and epidural haematoma), with a discussion of the relevant clinical implications. Finally, we discuss other promising research techniques and their relevance. The reader is referred to other publications for more detailed discussions of the epidemiology, clinical characteristics, and management of these disorders.

Physical principles

A basic knowledge of the pathophysiology and development of intracranial haemorrhage is essential to understand the sequential imaging changes seen on CT and MRI. Intracranial haemorrhages are typically divided into five distinct stages on the basis of blood breakdown products (table and webfigure 1): hyperacute (<12 h); acute (12 h to 2 days); early subacute (2–7 days); late subacute (8 days to 1 month); and chronic (>1 month to years).

In the hyperacute stage, the intracranial haemorrhage is a liquid composed of 99% intracellular oxygenated haemoglobin. During the next few hours, a heterogeneous clot develops that comprises blood cells, platelets, and protein-rich serum. As the clot retracts, the expelled serum and the reactive vasogenic oedema in the adjacent brain parenchyma surround it. During the acute stage, the intracranial haemoglobin is progressively deoxygenated, and in the subsequent early subacute phase, there is oxidative denaturation of haemoglobin, and deoxyhaemoglobin is gradually converted to methaemoglobin. During the late subacute stage, lysis of the red blood cells results in the release of methaemoglobin into the extracellular space, and the surrounding vasogenic oedema slowly subsides. During the chronic stage, macrophages and astroglial cells surround and slowly phagocytise the haematoma. Extracellular methaemoglobin is stored in the macrophages and is converted into haemosiderin and ferritin. Eventually, the haematoma will resolve completely, leaving a fluid-filled cyst or collapsed brain defect as a sequela.4 Time-dependent changes in intracranial haemorrhage are influenced by local alterations in haematoctrit, pH, and oxygen tension.

The CT imaging characteristics of intracranial haemorrhage are determined by the degree that the X-rays are attenuated by the blood products. The haematoctrit, degree of blood clot retraction, and haemoglobin content determine the density of the lesion seen on CT. The appearance of an intracranial haemorrhage on CT parallels the temporal evolution of the haemorrhage. In the hyperacute and acute phases the clot is hyperdense compared with the adjacent healthy brain parenchyma. During the first few hours post-haemorrhage, the progressive retraction of the clot and extrusion of the serum increases the concentration of haemoglobin, and the density of the haematoma increases; in patients with severe anaemia, reduced concentrations of haemoglobin result in the acute haematoma being less hyperdense than expected. During the late subacute phase, progressive lysis of the
red blood cells and proteolysis of the haemoglobin result in decreasing CT density of the haemorrhage, which becomes isodense to the adjacent brain parenchyma. In the chronic phase, the hypodense haematoma is progressively resorbed and typically leaves a small hypodense brain defect.4,5

The magnetic resonance signal characteristics of intracranial haemorrhage are determined by the paramagnetic effects of the breakdown products of haemoglobin, the magnetic field strength, and the pulse sequence used (table).3 The breakdown products of haemoglobin are characterised by the different oxidation states of iron and, as such, have different magnetic properties. In the hyperacute stage, because oxygenated haemoglobin has no unpaired electrons and is diamagnetic, the haematoma is isointense or mildly hyperintense on T1-weighted MRI and hyperintense on T2-weighted MRI, owing to the haemoglobin content. Deoxyhaemoglobin has few unpaired electrons and is paramagnetic; consequently, in the acute stage the haemorrhage is isointense or hypointense on T1-weighted MRI and hypointense on T2-weighted MRI.4,5 In the early subacute phase, methaemoglobin is paramagnetic and confined to the intracellular compartment, with a magnetic gradient between the intracellular and the extracellular compartments; the haemorrhage is hyperintense on T1-weighted MRI and hypointense on T2-weighted MRI. In the late subacute stage, methaemoglobin is released by lysis of the red blood cells, which cancels the magnetic gradient between the intracellular and the extracellular compartments; the haemorrhage is hyperintense on T1-weighted and T2-weighted MRI. Finally, in the chronic stage, because haemosiderin has many unpaired electrons and is superparamagnetic, the haemorrhage is hypointense on T1-weighted and T2-weighted MRI.4,5 Fluid-attenuated inversion recovery (FLAIR) MRI intensity changes for intraparenchymal haemorrhages are the same as those for other T2-weighted MRI sequences.

Until the development of T2*-weighted gradient echo (GRE) MRI sequences, T1-weighted and T2-weighted MRI sequences were used to estimate the age of an intracranial haemorrhage. In the 1990s, it was shown that GRE sequences are highly sensitive to the susceptibility effects of the paramagnetic and superparamagnetic breakdown products of haemoglobin;3,11 this increases the number of haemorrhagic lesions that can be identified.11,12 Echo-planar T2*-weighted imaging can be done with acquisition times in the seconds, which is a substantial advantage in patients with acute intracranial haemorrhage, who are unable to cooperate; conventional spin-echo images of such patients are often degraded by motion artifacts. The appearance of blood on GRE MRI can also be described by stage. The hallmark of hyperacute haemorrhage on GRE MRI sequences is a rim of hypointense signal that surrounds an isointense core. In the acute and subacute stages, the haematoma becomes diffusely hypointense, whereas in the chronic stage it appears as a slit-like signal with a hypointense or isointense core that has a rim of hypointensity. In some cases (eg, small haemorrhages in which the blood becomes rapidly deoxygenated and, therefore, the haemorrhage quickly appears hypointense throughout), GRE MRI sequences cannot help to estimate accurately the age of the haematomas.12

The value of diffusion-weighted imaging in the evaluation of intracranial haemorrhage is still under investigation.10,11,13 Diffusion-weighted imaging has promise in the differentiation of primary intracerebral haemorrhage from haemorrhagic cerebral ischaemia.13

**Intraparenchymal haemorrhage**

**Primary (spontaneous) intracerebral haemorrhage**

The most common risk factors associated with adult primary intracerebral haemorrhage are hypertension and cerebral amyloid angiopathy, and the probable underlying pathophysiology varies by haemorrhage location.13 Primary intracerebral haemorrhage associated with hypertension most commonly occurs in deep brain structures (eg, putamen, thalamus, cerebellum, and pons). By contrast, primary intracerebral haemorrhages that occur in lobar regions, particularly in elderly patients, are most commonly related to cerebral amyloid angiopathy but might also be associated with hypertension.14

Until recently, non-contrast CT was the gold standard for the diagnosis of intracerebral haemorrhage, and at many centres it is still the imaging modality of choice.
for the assessment of intracerebral haemorrhage, owing to its widespread availability and rapid acquisition time. Conventional T1-weighted and T2-weighted MRI pulse sequences are not sensitive to blood in the hyperacute stage; however, several recent studies have shown that GRE MRI sequences are as accurate as CT for the detection of intraparenchymal haemorrhage and far superior to CT for the detection of chronic haemorrhage (figure 1 and webfigures 2, 3, and 4). In some cases, MRI might actually detect haemorrhages that are missed on CT, and MRI is better than CT at identifying underlying structural lesions and for quantifying perihematomal oedema. However, up to 20% of patients with acute stroke are unsuitable for MRI (eg, they have pacemakers or metal implants or are unable to lie flat or still for the duration of the scan).

For both MRI and CT, baseline and serial studies can be used to identify patients who might benefit from acute interventions (eg, hydrocephalus that requires cerebrospinal fluid drainage or large or expanding cerebellar haematomas that require posterior fossa decompression). Both modalities can include postcontrast studies and vessel angiography to rule out underlying structural lesions (eg, arteriovenous malformation, aneurysm, or tumour). In general, contrast studies and catheter angiography are indicated in patients without a clear underlying aetiology or in patients with haemorrhages in unusual locations. One recent study suggested a high yield for angiography, even in patients with putaminal haemorrhage, unless the patient is older than 55 years and has hypertension.

Neuroimaging provides considerable information with regard to prognosis. Early CT studies in patients with intracerebral haemorrhage have shown that more than a third of patients have substantial (>33%) haematoma growth when imaged within 3 hours of onset. Haematoma expansion is associated with larger baseline haematoma volumes. In turn, haematoma expansion and the presence of intraventricular haemorrhage (webfigure 5) are predictors of poor outcome. Cortical intracerebral haemorrhage can be associated with better functional outcome; however, the rate of long-term recurrence is increased by a factor of 3–8. Several recent studies have reported that contrast extravasation seen on CT angiography might also be an early predictor of haematoma expansion (figure 2) and poor outcome. Therefore, CT angiography could have important applications for the selection of patients for acute therapies (eg, blood pressure management or haemostatic therapy).

Imaging techniques also provide insights into the underlying pathophysiology of intracerebral haemorrhage. Interest in these imaging approaches is driven by the belief that poor outcome in primary intracerebral haemorrhage results, in part, from ongoing secondary neuronal injury in the perihematomal region. Some of the studies that use PET, single-photon emission computed tomography (SPECT), and, more recently, perfusion MR and perfusion CT have shown perihematomal regions of hypoperfusion and bioenergetic compromise. The results of MRI studies have suggested that approximately a third of patients who are imaged in the acute phase will have reduced perihematomal apparent diffusion coefficient abnormalities on diffusion-weighted imaging, and further studies have characterised the timecourse for early perihematomal changes. Apparent diffusion coefficient values were low during the first day and then gradually increased, which probably matches the development of perihematomal oedema.

Alternatively, the results of studies with various imaging modalities have suggested that frank perihematomal hypoperfusion that reaches classic ischaemic thresholds does not occur. Specifically, the results of PET studies have not shown evidence of hypoxia or ischaemia (ie, normal oxygen extraction fraction and 18F-fluoromisonidazole uptake) surrounding intracerebral haemorrhage; therefore, the most prevalent hypothesis to account for perihematomal hypoperfusion is that it is due to secondary metabolic disturbances. The timecourse of these changes might explain the conflicting findings among studies. Analysis of the studies performed to date shows that there might be a subset of patients with a rim of perihematomal hypoperfusion and possibly ischaemia in the hyperacute phase; however, this region rapidly disappears in the subacute phase as oedema and inflammation develop. The development of oedema and toxicity from blood breakdown products are the most significant contributors to ongoing perihematomal injury, and imaging studies have the potential to monitor the effect on recovery and the influences of putative interventions on these parameters.

Table: Appearance of blood on CT and MRI by stage

<table>
<thead>
<tr>
<th>Stage</th>
<th>CT T1-weighted MRI</th>
<th>T2-weighted MRI</th>
<th>MRI FLAIR</th>
<th>GRE MRI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyperacute</td>
<td>Hyperdense</td>
<td>Hypointense or mildly hyperintense</td>
<td>Hyperintense</td>
<td>Hyperintense or Hypointense rim</td>
</tr>
<tr>
<td>(12 h to 2 d)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute</td>
<td>Hyperdense</td>
<td>Hypointense or hypointense</td>
<td>Hypointense</td>
<td>Hypointense</td>
</tr>
<tr>
<td>(2–7 d)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early acute</td>
<td>Hyperdense</td>
<td>Hyperintense</td>
<td>Hypointense</td>
<td>Hypointense</td>
</tr>
<tr>
<td>(8 d to 1 m)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chronic</td>
<td>Hypodense</td>
<td>Hypointense or hypointense</td>
<td>Hypointense</td>
<td>Silt-like hyperintense or isointense core that is surrounded by a hypointense rim</td>
</tr>
<tr>
<td>(&gt;1 m)</td>
<td></td>
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Microbleeds

GRE MRI has the ability to not only detect acute and chronic haematomas but also old, clinically silent cerebral microbleeds that are not seen on CT. Microbleeds are generally defined as punctate, homogeneous, rounded, hypointense lesions in the parenchyma that are smaller than 5–10 mm (figure 1 and webfigures 2 and 4). Pathological studies have shown that microbleeds seen with GRE MRI usually correspond to haemosiderin-laden macrophages adjacent to small vessels and are indicative of previous extravasation of blood.41

Microbleeds have been seen in up to 80% of patients with primary intracerebral haemorrhage, 21–26% of patients with ischaemic stroke, and 5–6% of asymptomatic or healthy elderly individuals.43–50 Hypertension, cerebral amyloid angiopathy, getting older, and, less commonly, cerebral autosomal dominant arteriopathy with silent infarcts and leukoaraiosis (CADASIL) have been identified as important risk factors for microbleeds.49,51–53 Recently, a mutation in COL4A1, the gene that encodes the basement membrane protein procollagen type IV alpha 1, has been reported in a French family; this mutation leads to small vessel cerebral disease, intracerebral haemorrhage, and microbleeds.54, 55

Microbleeds have been suggested to be markers of a bleeding-prone angiopathy.56–58 The results of several case reports and small series suggest that patients with microbleeds might be at increased risk of haemorrhage after antithrombotic or thrombolytic therapies. By contrast, the results of two large studies did not show an increased risk of haemorrhage in patients with microbleeds who were treated with intravenous tissue plasminogen activator.59,60 However, in both studies, few patients with large microbleed burdens were included and, therefore, the risk of haemorrhage in patients with a large number of microbleeds (>5) is unknown.

Microbleeds have important prognostic significance and are a common occurrence in patients with cerebral amyloid angiopathy; they are most commonly found in lobar regions (figure 1).56 By contrast, in patients with intracerebral haemorrhage due to hypertensive disease, microbleeds are most commonly found in deep and infratentorial regions (webfigure 4), although hypertension can also contribute to lobar microbleeds.59

A pattern of multiple haemorrhages without an underlying cause and restricted to lobar regions in an elderly patient is highly indicative of a diagnosis of cerebral amyloid angiopathy with the Boston Criteria.60 A particularly noteworthy finding is that the total number of microbleeds predict the risk of future symptomatic intracerebral haemorrhage in patients with lobar haemorrhage and probable cerebral amyloid angiopathy.60 Moreover, in this population, new microbleeds seen on repeat MRI also predict increased risk of subsequent symptomatic intracerebral haemorrhage, and microbleed burden and the rate of their accumulation predicts cognitive decline and poor neurological or functional outcome.62 The presence and lesion burden of microbleeds are notable prognostic factors in hypertensive haemorrhage. Several studies have shown a correlation between microbleeds and new vascular events.52,63

Secondary intracerebral haemorrhage

Intracerebral haemorrhage can be secondary to a large number of underlying pathologies (panel 1), many of which have distinct imaging characteristics. Multimodal
MRI or CT can help to identify the underlying pathology, although in some cases this might be obscured by the acute haematoma.

Underlying vascular lesions are more common in patients with intracerebral haemorrhages located in lobar lesions, and larger haematomas are more commonly associated with arteriovenous malformations. With an underlying arteriovenous malformation, flow voids can be seen, particularly on MRI. CT angiography or MR angiography might reveal the underlying vascular lesion; however, in some cases, catheter angiography is required and might need to be repeated if the results were initially negative owing to the mass effect of the haematoma. The findings from imaging that might be deemed indications for catheter angiography include abnormal calcifications, the presence of subarachnoid blood, vessel abnormalities, or unusual haemorrhage locations.7 Cavernous malformations are best seen on GRE MRI sequences and appear as a hypointense ring (due to haemosiderin from a previous bleeding event) around the malformation (webfigure 3). Isolated cavernomas can be difficult to diagnose in the acute phase because their appearance can resemble that of a primary haemorrhage.

Imaging can also provide clues to non-vascular aetiologies of intracerebral haemorrhage. Traumatic intracerebral haemorrhage is commonly accompanied by evidence of a scalp haematoma and a coup–contrecoup location. Contusions have a marked predilection for the orbitofrontal regions and the anterior temporal lobes. The presence of a fluid-blood level (ie, the horizontal interface between hypodense bloody serum layered above hyperdense settled blood) is highly suggestive of a traumatic aetiology or an underlying coagulopathy.64,65 In one study, the presence of a fluid-blood level had 59% sensitivity and 98% specificity for abnormal prothrombin time or partial thromboplastin time (ie, the time it takes for the blood to clot).65 Haemorrhagic lesions associated with diffuse axonal injury from traumatic shearing lesions can be seen on MRI, particularly with high-spatial-resolution, susceptibility-weighted techniques.66 Haemorrhages due to sinus occlusions are typically found in the parasagittal region if the thrombosis is superficial, or they are bithalamic if due to occlusion of the deep sinuses. An underlying vascular lesion (aneurysm or arteriovenous malformation)67 or a vasculopathic pattern on invasive or non-invasive vessel imaging studies can be seen in intracerebral haemorrhages that accompany drug use. Tumours are typically enhanced on postcontrast studies and have been shown to have substantially greater amounts of vasogenic oedema compared with intracerebral haemorrhages due to other aetiologies.68 Haemorrhages due to underlying infections commonly have lesion enhancement.

Haemorrhagic transformation

Haemorrhagic transformation of an ischaemic infarct occurs commonly as part of the natural history of the disease and in the setting of reperfusion therapies. The most commonly used radiological classification system for rating the type and severity of haemorrhagic transformation on head CT scans divides the haemorrhage into two major categories: haemorrhagic infarct (figure 3) and parenchymal haematoma (webfigure 6); each division is subdivided into two subcategories on the basis of severity (panel 2).69

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Figure 2: Imaging of a 69-year-old man 2 hours after the onset of right-sided paralysis
(A) Non-contrast CT taken on admission shows left thalamic haematoma with extension of the haemorrhage into the third ventricle. (B) Contrast-enhanced CT taken on admission shows active extravasation within the haematoma. (C) Follow-up non-contrast CT taken 12 hours after admission shows notable growth of the haematoma, with haemorrhage in both lateral ventricles and severe hydrocephalus. Active contrast extravasation in intraparenchymal haematoma as seen with CT angiography and postcontrast CT predicts mortality and haematoma growth.65

Figure 3: Example of haemorrhagic transformation of an infarct in a branch of the left middle cerebral artery
The infarct is seen with (A) diffusion-weighted imaging, (B) CT, and (C) GRE MRI.

See Online for webfigure 6

Panel 2: CT classification of haemorrhagic transformation

- **Haemorrhagic infarction type 1 (HI-1)**: Small petechiae along the margins of the infarct
- **Haemorrhagic infarction type 2 (HI-2)**: More confluent petechiae within the infarcted area but without space-occupying effect
- **Parenchymal haematoma type 1 (PH-1)**: Haematoma in up to 30% of infarcted area with some space-occupying effect
- **Parenchymal haematoma type 2 (PH-2)**: Dense haematoma in more than 30% of infarcted area with substantial space-occupying effect or any haemorrhagic lesion outside the infarcted area
The application of this classification system to MRI has not been systematically assessed and compared with CT. Moreover, studies have shown that GRE MRI sequences are more sensitive to haemorrhagic transformation than is CT (figure 3), which makes the standard classification across modalities problematic. In one study of 25 patients, GRE MRI sequences detected haemorrhagic transformation in 12 patients compared with 7 patients detected with CT.

Imaging information can help to distinguish the haemorrhagic transformation of an ischaemic infarct from a primary haematoma. Most haemorrhagic transformations are smaller than the field of the ischaemic infarct. Thus, FLAIR MRI sequences, diffusion-weighted imaging, and the apparent diffusion coefficient sequence can provide information from the non-haemorrhagic regions and show that the blood is within a larger ischaemic infarct. Primary haematomas also tend to be rounder and commonly have a greater amount of surrounding oedema than would be seen in an ischaemic stroke. Furthermore, haematomas do not necessarily respect vascular territories.

A growing number of studies have assessed the clinical and radiological predictors of haemorrhagic transformation in the setting of thrombolytic therapy. In the NINDS trial of intravenous tissue plasminogen activator given within 3 hours of onset, the presence of signs of early infarct was associated with an increased risk of subsequent haemorrhagic transformation in the treated group; however, this association was not independent of baseline variables, which included the National Institutes of Health Stroke Scale score to measure the neurological deficit, and these patients fared better with tissue plasminogen activator treatment. In the European Cooperative Acute Stroke Study I (ECASS I), patients enrolled in the trial who had signs of major (involving more than a third of middle cerebral artery territory) early infarct on CT (protocol violators) had an increased rate of haemorrhagic transformation compared with patients without signs of extended early infarct on CT; the patients with signs of major infarct did not benefit from thrombolytic therapy. An analysis of the subsequent ECASS II results revealed that the extent of hypodensity at baseline was an independent risk factor for severe haemorrhagic transformation. Thus, on the basis of the results of these trials, CT changes detected within 3 hours that involve more than a third of the middle cerebral artery territory do not constitute a contraindication to tissue plasminogen activator treatment, whereas beyond 3 hours they would. Of note, the results of secondary analyses from these two ECASS studies have suggested that only parenchymal haematoma type 2 (webpanel 2) lesions are associated with clinical deterioration and poor prognosis, although this is controversial.

On both CT and MRI, contrast extravasation is a marker of blood–brain barrier disruption and a predictor of haemorrhagic transformation. In addition, a growing number of MRI studies have identified additional imaging characteristics that predict haemorrhagic transformation. The results of several of these studies have suggested that large baseline diffusion-weighted imaging lesions and low apparent diffusion coefficient values are independent predictors of symptomatic haemorrhage. These are important areas for future research and provide a potential biomarker to identify high-risk patients and test therapies designed to minimise the risk of blood–brain barrier opening and subsequent haemorrhagic transformation.

Subarachnoid haemorrhage

Immediately after a subarachnoid haemorrhage, the subarachnoid space is filled with erythrocytes suspended in the CSF. The erythrocytes are trapped within the arachnoid villi and trabeculae and are progressively cleared from the CSF into the vascular system or phagocytically removed by leptomeningeal macrophages while red blood cell lysis releases haemoglobin in the CSF.

Subarachnoid haemorrhage can be of traumatic or non-traumatic origin, with trauma being the most common aetiology. Subarachnoid haemorrhage can occur in head injury, as a result of laceration of the cortical veins or arteries that course through the subarachnoid space but can also occur as a result of cortical contusions and lacerations with extravasation of blood into the
subarachnoid space. Typical locations for subarachnoid haemorrhage after head trauma are the interpeduncular cistern, Sylvian fissure, or over the cerebral convexity. Spontaneous subarachnoid haemorrhage usually results from ruptured aneurysms (older patients), arteriovenous vascular malformations (younger patients), and can be less commonly due to intracranial vessel dissection or vasculitis. Perimesencephalic (or pretruncal) non-aneurysmal subarachnoid haemorrhage is a benign variant of unknown (probably venous) cause, with the centre of bleeding most commonly located in the prepontine cistern. Subarachnoid haemorrhage can also be present in Call–Fleming syndrome—a generally benign and reversible condition due to segmental cerebral vasoconstriction that presents with severe recurrent headaches and transient or fluctuating neurological abnormalities.

CT is sensitive in detecting blood in the CSF space and is usually more easily available than MRI in the acute setting of subarachnoid haemorrhage (figure 4). Typically, in patients with acute subarachnoid haemorrhages, hyperdense blood is seen within the basal cisterns and subarachnoid spaces. In about 10% of cases, the CT will be negative, particularly when obtained more than 12 hours after the onset of symptoms, and a lumbar puncture is required to confirm the diagnosis. The location and distribution of the subarachnoid haemorrhage can indicate the site of the vascular malformation, and contrast-enhanced CT angiography can show the exact location and shape of the aneurysm.

Acute subarachnoid haemorrhage is difficult to detect with conventional T1-weighted and T2-weighted MRI sequences. Subarachnoid haemorrhage differs from intraparenchymal haemorrhage in that the mixing of blood with high oxygen tension CSF delays the generation of paramagnetic deoxyhaemoglobin, and oxyhaemoglobin persists for longer than in intraparenchymal haemorrhage. Variability in the appearance of subarachnoid haemorrhage after 48 hours most probably relates to haemoglobin degradation, which adds further complexity to the MRI signal intensity.

FLAIR MRI is sensitive enough to detect acute subarachnoid haemorrhage because suppression of the CSF signal enhances the conspicuity of the haemorrhage (webfigure 7). The FLAIR MRI techniques consist of an inversion recovery pulse to null the signal from the CSF and a long echo time to produce a heavily T2-weighted sequence. When disease occurs within the subarachnoid space, the relaxation time of the CSF is altered, which results in a change in the T1-weighted signal; this change translates into reduced CSF signal nulling and consequent hyperintensity of the CSF or subarachnoid space during the FLAIR MRI sequence. Such findings have been well described in a wide range of pathological conditions, including subarachnoid haemorrhage, meningitis, and the leptomeningeal spread of malignant disease. Other, less common, causes of subarachnoid FLAIR MRI hyperintensity include slow flow downstream from vascular stenosis or occlusion, artifacts caused by supplemental oxygen, and sulcal enhancement following contrast administration when the blood–brain barrier is disrupted.

The results of in-vivo and in-vitro studies have suggested that FLAIR imaging (figure 4) is at least as sensitive as CT for the evaluation of acute subarachnoid haemorrhage; however, compared with the findings from lumbar puncture, the findings on FLAIR MRI are not definitive in excluding acute subarachnoid haemorrhage. The FLAIR MRI sequence is particularly useful to image acute subarachnoid haemorrhage in areas where CT might be limited because of beam-hardening artifacts. Few studies have systematically assessed GRE MRI sequences in patients with subarachnoid haemorrhage or compared the accuracy of the sequences with FLAIR MRI, although one recent study suggested good accuracy for GRE MRI sequences when assessed on a 3-Tesla scanner.

Subdural haematoma

Subdural haematomas can be diagnosed in the acute, subacute, or chronic stages on the bases of location.
severity, and various clinical presentations. They can occur after head trauma, coagulopathy, antithrombotic therapy, or spontaneously. They are most commonly seen along the cerebral convexities, the falk cerebri, and the tentorium cerebelli. If caused by trauma, a scalp haematoma or bone fracture might also be seen.

The appearance of subdural haematomas on CT and MRI depends on the stage and location (figure 5). In the acute stage, subdural haematoma typically appears as a hyperdense crescent along the cerebral convexity, a linear hyperdensity between the hemispheres, or a thin hyperdense layering along the tentorium cerebelli. In the subacute stage, subdural haematomas can be difficult to detect on CT because they become isodense to grey matter and might only be apparent because of the mass effect. Chronic subdural haematomas are hypodense compared with grey matter. Recurrent bleeding into the subdural space can lead to a bilayered or mixed pattern, with blood breakdown products at various stages.

The appearance of subdural haemorrhage on MRI depends not only on the stage of the lesion but also on the sequence used. The development of signal changes might not directly correlate with that of intraparenchymal haematomas.302 The results of a systematic study that compared signal changes in patients who had intracerebral haemorrhage with patients who had subdural haemorrhage showed that, in the acute stage, the subdural haematoma lesions appear as hyperintense on FLAIR MRI and T2-weighted sequences. On GRE MRI sequences, subdural haematomas can be isointense or hypointense in the acute stage, depending on the stage of blood breakdown products. In the subacute stage, the breakdown products appear hypointense, whereas at chronic timepoints they appear as either isointense or mixed signals, depending on the degree of blood reabsorption.

**Epidural haematoma**

Epidural haematomas usually occur after acute head traumas, and their clinical presentation might be delayed (eg, with a symptom-free or lucid interval). Arterial epidural haematomas usually result from laceration of branches of the meningeal arteries, whereas venous epidural haematomas result from a tear in the dural venous sinuses. Venous epidural haematomas are more commonly seen in children, and their clinical presentation might be delayed (eg, with a symptom-free or lucid interval). Arterial epidural haematomas usually result from laceration of branches of the meningeal arteries, whereas venous epidural haematomas result from a tear in the dural venous sinuses. Venous epidural haematomas are more commonly seen in children, and their clinical presentation might be delayed (eg, with a symptom-free or lucid interval). Arterial epidural haematomas usually result from laceration of branches of the meningeal arteries, whereas venous epidural haematomas result from a tear in the dural venous sinuses. Venous epidural haematomas are more commonly seen in children, and their clinical presentation might be delayed (eg, with a symptom-free or lucid interval).

The epidural haematoma or bone fracture might also be seen. The tight adherence of epidural haematomas. No dilution with CSF occurs because an epidural haematoma is separated from the CSF by the dura mater.

**New imaging applications**

PET and SPECT techniques, although not practical in the acute clinical setting, have provided important information with regard to changes in blood flow and their timecourse, particularly in patients with primary intracerebral haemorrhage. PET imaging with the [11C] Pittsburgh compound-B has been used to detect cerebrovascular β-amyloid in patients with intracerebral haemorrhage that is secondary to amyloid angiopathy.304

Several additional techniques, including near-infrared spectroscopy, magnetic resonance spectroscopy, and diffusion tensor imaging, promise to provide new insights into the diagnosis and pathophysiology of haemorrhage, the timecourse of cerebral injury, recovery, and neuroplasticity, and the response to putative therapies. Near-infrared spectroscopy shows potential as a diagnostic means for the rapid identification of subdural and epidural haematomas in the field or at the bedside in patients with head trauma.305,306

Several studies have used magnetic resonance spectroscopy to assess perihaematominal metabolic and haemodynamic changes in patients with intracerebral haemorrhage. Two small studies have reported metabolic changes in higher cortical structures in patients with deep intracerebral haemorrhage, which suggests that these changes might correlate with long-term motor deficit and clinical outcome.307,308

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Figure 6: Examples of epidural haematomas

(A) Non-contrast CT of a 20-year-old male patient who was involved in a motor vehicle accident shows a right frontal epidural haematoma, with typical biconvex shape and the non-crossing of sutures. This epidural haematoma resulted from an arterial bleed due to a right frontal calvarial fracture. (B,C) Epidural haematoma that displaced and compressed the white matter tracts of the left frontal lobe in a 22-year-old woman who was imaged 6 days after mild traumatic brain injury (Glasgow coma score of 15). (B) GRE MRI sequence from 3T MRI shows the large lentiform epidural haematoma as a dark signal, space-occupying, extra-axial mass in the left frontal lobe. There is medial displacement and compression of white matter in the left frontal lobe and midline shift to the right, which signifies subfalcine herniation. (C) Directionally encoded colour anisotropy map from 3T diffusion tensor imaging shows white matter tracts with left–right fibre orientation in red, anteroposterior in green, and craniocaudal in blue. 3D fibre tractography of the commissural tracts of the genu of the corpus callosum (bright yellow). Medial deviation of the commissural fibres of the left frontal lobe is evident.
Further studies of more patients with serial scanning are needed to better characterise the dynamic nature of perihematological metabolic changes.109

Another promising approach is diffusion tensor imaging (figure 6). Diffusion tensor imaging can be used to visualise white matter fibres and map fibre tracts (eg, the corticospinal tract). This technique might be particularly useful to understand the process and potential of motor recovery by assessment of the integrity of the corticospinal tract in the acute, subacute, and chronic stages of intracerebral haemorrhage.110 In patients with intracerebral haemorrhage secondary to cerebral amyloid angiopathy, diffusion tensor imaging shows a characteristic pattern of regional tissue degeneration.111

Future approaches to neuroimaging research are likely to be available for clinical studies of haemorrhage and might include high-strength magnets (eg, 8 Tesla) to detect microhaemorrhages and cerebrovascular amyloid, in addition to tissue probes and imaging biomarkers of inflammation, oedema, and excitotoxicity. An important future goal of neuroimaging techniques will include imaging-guided therapy that targets specific stages of injury (eg, blood–brain barrier disruption, oedema, or inflammation).

Conclusions

Neuroimaging sequences have a crucial role in the diagnosis of intracranial haemorrhage, and advances in neuroimaging techniques will continue to provide future insights into the mechanisms of injury. One of the most promising future applications of imaging is to validate markers of increased risk of haemorrhagic transformation in patients with ischaemic stroke and to use this information to guide treatment and develop therapies that reduce risk (eg, drugs that stabilise the blood–brain barrier). An important area for future research is to determine whether neuroimaging findings can be used as a surrogate outcome measure to test acute therapies for primary intraparenchymal haemorrhages. In addition, these imaging techniques have the potential to provide further insights into the timecourse and probable dynamic pathophysiology of injury in patients with intracerebral haemorrhage. New techniques, including magnetic resonance spectroscopy and diffusion tensor imaging, might have important applications in the understanding of injury due to and recovery from haemorrhage.

Contributors

The authors contributed equally to all parts of this Review.

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Conflicts of interest

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Review


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