6.1 Conventional MRI and Stroke

6.1.1 Hyperacute Infarct

In the hyperacute stage of infarct, there is occlusion or slow flow in the vessels supplying the area of infarcted tissue. Within minutes of the infarct, the signal flow void on T2-weighted images is lost (Fig. 6.1). FLAIR (fluid attenuated inversion recovery), an inversion recovery sequence that suppresses the CSF signal, can show high intravascular signal against the surrounding low-signal subarachnoid space [1] (Fig. 6.2). In one study, 65% of infarcts <6 h old showed a FLAIR high signal within vessels, and, in some cases, the finding of a FLAIR high signal in vessels preceded changes in the diffusion-weighted images [2].

Gradient recalled echo (GRE) T2*-weighted images can detect an intraluminal thrombus (deoxyhemoglobin) in hyperacute infarcts as a linear low signal region of magnetic susceptibility (Fig. 6.2). In one

6.1.2 Acute Infarct

6.1.3 Subacute Infarct

6.1.4 Chronic Infarcts

6.1.5 Hemorrhagic Transformation

6.1.6 Conclusion

6.2 MR Angiogram and Stroke

6.2.1 Noncontrast MRA

6.2.1.1 TOF MRA

6.2.1.2 Phase-Contrast MRA

6.2.2 Contrast-Enhanced MRA

6.2.3 Image Processing

6.2.4 Extracranial Atherosclerosis and Occlusions

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6.2.7.1 Moya Moya

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6.2.7.3 Fibromuscular Dysplasia

6.2.8 Venous Infarct

6.2.9 Conclusion

References
study of 30 patients with MCA thrombus, gradient echo had 83% sensitivity in detecting the thrombus, compared with 52% sensitivity for noncontrast CT in detecting a dense MCA sign [3].

Contrast-enhanced T1-weighted images show arterial enhancement in 50% of hyperacute strokes [1] (Fig. 6.3). This arterial enhancement is thought to be secondary to slow flow, collateral flow or hyperperfusion following early recanalization. It may be detected as early as 2 h after stroke onset and can persist for up to 7 days. During this period, there is usually no parenchymal abnormality because inadequate collateral circulation prevents contrast from reaching the infarcted tissue. Rarely, early parenchymal enhancement may occur when there is early reperfusion or good collateralization.
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In the hyperacute period (first 6 h), there is shift of water from the extracellular to the intracellular space but there may be little increase in overall tissue water. Therefore, while the development of altered signal intensity on FLAIR- or T2-weighted images may occur as early as 2–3 h after stroke onset, conventional MRI is not sensitive enough for evaluation of infarcts in the hyperacute stage; one study found an 18% sensitivity of T2-weighted images in detecting infarct in the first 6 h and a false-negative rate of 30–50% [4]. FLAIR-weighted sequences are slightly more sensitive than T2-weighted images in the detection of parenchymal changes in acute infarcts but have an estimated sensitivity of only 29% in the first 6 h [5].

6.1.2 Acute Infarct

By 24 h, as the overall tissue water content increases due to vasogenic edema following blood–brain barrier disruption, conventional MRI becomes more sensitive for the detection of parenchymal infarcts. Signal changes during the first 24 h are best appreciated in the cortical and deep gray matter. Infarcts in the acute stage usually demonstrate focal or confluent areas of T2 and FLAIR hyperintensity with sulcal effacement. During this time, the white matter may be hyperintense, but also may show no abnormality or demonstrate hypointensity. Proposed etiologies for the subcortical white matter hypointensity are free radicals, sludging of deoxygenated red blood cells, and iron deposition [6]. Because cerebrospinal fluid (CSF) is hypointense, FLAIR has improved detection of small infarctions in brain parenchyma, such as cortex and periventricular white matter, adjacent to CSF. By 24 h, T2-weighted and FLAIR-weighted images detect 90% of infarcts [1]. An increase in tissue water also leads to hypointensity on T1-weighted images. However, in the acute period, T1-weighted images are relatively insensitive at detecting parenchymal changes compared with T2-weighted images. At 24 h, sensitivity is still only approximately 50%.

6.1.3 Subacute Infarct

In the subacute phase of infarct (1 day to 2 weeks), the increase in vasogenic edema results in increased T2 and FLAIR hyperintensity, increased T1 hypointensity, and better definition of the infarction and swelling (Fig. 6.4). The brain swelling is manifest as gyral thickening, effacement of sulci and cisterns, effacement of adjacent ventricles, midline shift, and brain herniation. The swelling reaches a maximum at about 3 days and resolves by 7–10 days [7]. There is increased T2 and FLAIR signal within the first week that usually persists but there may be “MR fogging” [8]. MR fogging occurs when the infarcted tissue becomes difficult to see because it has developed a signal intensity similar to that of normal tissue. This is thought to result from infiltration of the infarcted tissue by inflammatory cells. One study of 7- to 10-day-old strokes identified 88% of subacute strokes on T2-weighted images, and another determined that T2- and FLAIR-weighted images were equally sensitive at detecting 10-day-old infarcts [7].

In the subacute phase, arterial enhancement peaks at 1–3 days. Large infarcts will also demonstrate meningeal enhancement that may represent reactive
hyperemia, which peaks at 2–6 days. Arterial and meningeal enhancement both typically resolve by 1 week [9]. In addition, parenchymal enhancement occurs during this phase. Gray matter enhancement can appear band-like or gyriform (Fig. 6.5). This is secondary to disruption of the blood–brain barrier and restored tissue perfusion from a recanalized occlusion or collateral flow. This parenchymal enhancement may be visible at 2–3 days but is consistently present at 6 days and persists for 6–8 weeks [9]. Some infarcts, such as watershed and noncortical infarcts, may enhance earlier.

6.1.4 Chronic Infarcts

After 2 weeks, the mass effect and edema within infarcts decrease and the parenchyma develops tissue loss and gliosis. During this time, parenchymal enhancement peaks at 1–4 weeks and then gradually fades [9]. The chronic stage of infarction is well es-

**Fig. 6.4 a, b**

Cortical edema in a subacute infarct. **a** The axial FLAIR-weighted image shows high signal, gyral swelling, and sulcal effacement. **b** There is subtle low signal and gyral swelling (arrow) seen on the T1-weighted sagittal image.
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6.1.5 Hemorrhagic Transformation

Hemorrhagic transformation (HT) of brain infarction represents secondary bleeding into ischemic tissue, varying from small petechiae to parenchymal hematoma. It has a natural incidence of 15% to 26% during the first 2 weeks and up to 43% over the first month after cerebral infarction [12, 13]. Predisposing factors include stroke etiology (HT is more frequent with embolic strokes), reperfusion, good collateral circulation, hypertension, anticoagulant therapy, and thrombolytic therapy. In patients treated with intraarterial (i.a.) thrombolytic therapy, a higher National Institutes of Health Stroke Scale (NIHSS) score, longer time to recanalization, lower platelet count and a higher glucose level are associated with HT [14].

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Wallerian degeneration. Coronal T2-weighted image shows encephalomalacia of the right frontal and temporal lobes and T2 high signal extending into the right cerebral peduncle (arrow) from Wallerian degeneration.

Laminar necrosis. This sagittal noncontrast T1-weighted image shows gyriform T1 high signal in a chronic left MCA infarct. Mild enlargement of the sulci is consistent with encephalomalacia.

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Established by 6 weeks. At this point, necrotic tissue and edema are resorbed, the gliotic reaction is complete, the blood–brain barrier is intact, and reperfusion is established [9]. There is no longer parenchymal, meningeal or vascular enhancement, and the vessels are no longer hyperintense on FLAIR images. There is tissue loss with ventricular, sulcal, and cisternal enlargement. There is increased T2 hyperintensity and T1 hypointensity due to increased water content associated with cystic cavitation. With large middle cerebral artery (MCA) territory infarctions, there is Wallerian degeneration, characterized by T2 hyperintensity and tissue loss, of the ipsilateral cortical spinal tract [10] (Fig. 6.6).

Chronic infarcts can demonstrate peripheral gyriform T1 high signal from petechial hemorrhage or from laminar necrosis [11] (Fig. 6.7).
Because T1-, T2-, and FLAIR-weighted images are insensitive at detecting acute blood products (deoxyhemoglobin), GRE T2* sequences should be used to detect hemorrhage in the acute stroke setting (Fig. 6.8). GRE T2* sequences have increased sensitivity to blood breakdown products due to their paramagnetic properties. One study demonstrated GRE images to be as sensitive as CT at detecting parenchymal hemorrhage in acute strokes [15]. Another study observed that in detecting acute blood products, GRE T2*-weighted images were more sensitive than T2-, FLAIR-, or echo planar T2-weighted images [16]. Some data suggest that microbleeds detected by susceptibility predict symptomatic hemorrhage following tissue plasminogen activator (t-PA) treatment [17].

As blood products evolve into methemoglobin, T1-weighted sequences become more sensitive at detecting blood products [18]. Chronic hemorrhages are best detected on GRE T2* images as areas of susceptibility (Fig. 6.9).

6.1.6 Conclusion

Conventional MRI can diagnose infarcts at all stages of temporal evolution but are most sensitive after the hyperacute stage (see Table 6.1). During the hyperacute stage, the predominant findings are loss of flow voids on T2-weighted images, FLAIR hyperintensity in affected vessels, and vascular enhancement. In the acute to chronic stage, FLAIR and T2 parenchymal abnormalities are evident and Wallerian degeneration develops. The subacute stage is also marked by parenchymal swelling followed by parenchymal enhancement. The detection of acute hemorrhagic infarct requires the use of a T2* GRE susceptibility sequence since other conventional MRI sequences are not sensitive enough for acute bleeds.
6.2 MR Angiogram and Stroke

Magnetic resonance angiography (MRA) is a set of vascular imaging techniques capable of depicting the extracranial and intracranial circulation. In the setting of acute stroke, these techniques are useful for determining stroke etiology and assessing vascular flow dynamics. Specifically, they are used to evaluate the severity of stenosis or occlusion as well as collateral flow. A typical stroke protocol includes two-dimensional (2D) and/or three-dimensional (3D) time-of-flight (TOF) and contrast-enhanced MRA images through the neck and 3D TOF MRA images through the Circle of Willis. For dissection, a fat saturated pre-gadolinium axial T1 sequence through the neck is

Table 6.1 The appearance of arterial infarcts on conventional MRI

<table>
<thead>
<tr>
<th>Stage</th>
<th>Conventional MR appearance</th>
<th>Evolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyperacute (0–6 h)</td>
<td>T2 shows loss of signal flow void</td>
<td>Occurs within minutes of the infarct</td>
</tr>
<tr>
<td></td>
<td>FLAIR shows vessel high signal</td>
<td>Occurs within minutes of the infarct</td>
</tr>
<tr>
<td></td>
<td>GRE T2* shows blooming susceptibility</td>
<td>Occurs within minutes of the infarct</td>
</tr>
<tr>
<td></td>
<td>T1 post-contrast shows arterial enhancement</td>
<td>Occurs at 2 hours and can last 7 days</td>
</tr>
<tr>
<td></td>
<td>There are no reliable parenchymal findings for infarct at this stage</td>
<td></td>
</tr>
<tr>
<td>Acute (6–24 hours)</td>
<td>Vascular abnormalities from the hyperacute stage persist</td>
<td>Appears by 24 h (90% sensitivity)</td>
</tr>
<tr>
<td></td>
<td>T2 and FLAIR show gyriform high signal and sulcal effacement</td>
<td></td>
</tr>
<tr>
<td></td>
<td>T2 can show subcortical low signal</td>
<td>Can occur at any time in the acute to subacute stage</td>
</tr>
<tr>
<td></td>
<td>GRE T2* show gyriform susceptibility from petechial bleed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No parenchymal enhancement is present</td>
<td></td>
</tr>
<tr>
<td>Subacute (1 day to 2 weeks)</td>
<td>T2 and FLAIR show gyriform high signal and sulcal effacement from mass effect</td>
<td>Reaches a maximum at 3–5 days and decreases by 1 week. Rarely, “MR fogging” appears at 2 weeks</td>
</tr>
<tr>
<td></td>
<td>T1 can show gyriform high signal from petechial bleed</td>
<td>Appears once methemoglobin develops</td>
</tr>
<tr>
<td></td>
<td>T1 post-contrast shows arterial enhancement</td>
<td>Peaks at 1–3 days and resolves by 1 week</td>
</tr>
<tr>
<td></td>
<td>T1 post-contrast shows meningeal enhancement</td>
<td>Peaks at 2–6 days and resolves by 1 week</td>
</tr>
<tr>
<td></td>
<td>T1 post-contrast show parenchymal enhancement as vessels recanalize</td>
<td>Can be seen at 2–3 days, consistently seen at 6 days and persists for 6–8 weeks</td>
</tr>
<tr>
<td>Chronic</td>
<td>T2 shows high signal from gliosis and Wallerian degeneration</td>
<td>Persists indefinitely</td>
</tr>
<tr>
<td></td>
<td>Volume loss is present</td>
<td>Persists indefinitely</td>
</tr>
<tr>
<td></td>
<td>T1 shows low signal from cavitation</td>
<td>Persists indefinitely</td>
</tr>
<tr>
<td></td>
<td>GRE T2* shows low signal hemosiderin from petechial bleeds</td>
<td>Persists indefinitely</td>
</tr>
</tbody>
</table>
added. This section discusses the major MRA techniques, including the advantages and disadvantages of each, as well as how MRA is used to evaluate the specific disease processes that lead to stroke.

### 6.2.1 Noncontrast MRA

MRA is broadly divided into noncontrast and contrast-enhanced techniques. Noncontrast MRA can be acquired with phase contrast (PC) or TOF techniques, and both can be acquired as 2D slabs or 3D volumes.

#### 6.2.1.1 TOF MRA

TOF MRA is a gradient echo sequence that depicts vascular flow by repeatedly applying a radio frequency (RF) pulse to a volume of tissue, followed by dephasing and rephasing gradients. Stationary tissues in this volume become saturated by the repeated RF pulses and have relatively low signal. By contrast, flowing blood in vessels has relatively increased signal because it continuously carries unsaturated spins into the imaging volume. The vessel contrast or flow-related enhancement is proportional to the velocity. Signal from venous blood is minimized by placing a saturation band above the imaging volume [19].

Two-dimensional TOF MRA is typically performed in the neck region with a relatively large flip angle (60°). Multiple sequential, 1-mm-thick axial slices are obtained. Blood flowing perpendicular to the multiple thin slices is well imaged because it is not exposed to enough RF pulses to become saturated. However, blood flowing in the imaging plane (in-plane flow) is exposed to more RF pulses and may become saturated and lose signal; this artifact can often be seen at the horizontal turns of the vertebral arteries as well as the petrous segments of the internal carotid arteries (Fig. 6.10) [19].

Three-dimensional TOF MRA is typically performed in the head region with a smaller flip angle (20°). A volume of tissue covering the skull base to the Circle of Willis is obtained and then divided into 1-mm-thick slices using an additional phase encoding step [20]. While the smaller flip angle reduces saturation artifacts, any flowing blood that remains in the imaging volume for long enough is exposed to multiple RF pulses and can artifactually lose signal. This signal loss is usually seen in the distal Circle of Willis vessels (Fig. 6.11). The smaller flip angle also decreases the background saturation. Typically, a ramped flip angle is used to minimize vascular saturation effects while maximizing the suppression of the background signal. Magnetization transfer is additionally applied in 3D TOF MRA to further decrease the background signal.

Compared with 2D TOF, the 3D TOF technique has better spatial resolution, a better signal-to-noise ratio and less intravoxel dephasing, but it is more limited by the vascular saturation artifact and therefore can cover only a small volume. A hybrid technique between 2D and 3D TOF known as MOTSA (multiple overlapping thin slab acquisition) acquires partially overlapping thin 3D volumes. The ends of the 3D volume have saturation artifact and are discarded while...
central portions are assembled into a single MRA. MOTSA has higher spatial resolution than 2D TOF while covering a larger area than 3D TOF MRA, because it is less susceptible to saturation artifact.

TOF MRA, especially 2D TOF, is also vulnerable to artifactual signal loss from flow turbulence. This causes phase dispersion so that the rephasing gradient is unable to generate a strong echo. This artifact can be seen at vessel bifurcation points or distal to stenoses and can result in overestimates of the degree or length of vascular stenoses (Fig. 6.12).
TOF MRA can have artifact from poor background suppression. Signal from stationary tissues are in theory suppressed with repeated RF pulses, but substances with a short T1, such as fat or methemoglobin in hematomas, are not usually fully saturated. Consequently, these substances demonstrate high signal on TOF MRA and can mimic areas of flow or obscure vessels (Fig. 6.13). This artifact can sometimes be overcome by segmenting out areas of high signal, and high fat signal can be reduced by setting the echo time (TE) to 2.3 or 6.9 ms, to place fat and water out of phase.

6.2.1.2 Phase-Contrast MRA

Phase-contrast MRA (PC MRA) is a gradient echo sequence that depicts blood flow by quantifying differences in the transverse magnetization between stationary and moving tissue [19]. Following a RF pulse, pairs of symmetric but opposed phase encoding gradients are applied in one direction within the imaging volume; the first gradient dephases and the second rephases the transverse magnetization. Stationary tissues have no net change in phase because they experience equal but opposite magnetic field environments during the dephasing and then the rephasing gradients. Moving blood, however, experiences different magnetic field environments as each gradient is applied. The spins in moving blood acquire a phase shift during the dephasing pulse, which is not completely reversed during the rephasing pulse. The net phase shift, either positive or negative, determines direction of flow, and the amount of phase shift (in degrees) is proportional to the velocity or magnitude of blood flow.

PC MRA is acquired in three orthogonal directions, and the direction of flow is depicted as a relatively high or low signal against a gray background (Fig. 6.14). Orthogonal maps can also be combined to form an overall flow-related enhancement map without directionality. Like TOF MRA, PC MRA can be obtained with 2D or 3D techniques. Clinically, PC MRA is used to evaluate intracranial collateral flow distal to a stenosis.

An artifact unique to PC MRA is that phase shifts exceeding 180° are interpreted as slow flow in the opposite direction. This aliasing artifact leads to incorrect determination of flow direction. In order to avoid this, a velocity encoding parameter (VENC) is selected which represents the maximum expected flow velocity in the imaging volume. This value adjusts the strength of the bipolar gradients to prevent phase shifts from exceeding 180°.
PC MRA has several advantages over TOF MRA. As mentioned above, PC MRA can demonstrate flow direction. Also, since PC MRA shows only moving tissues, stationary tissues with short T1 such as fat or methemoglobin do not demonstrate high signal. Another advantage is that PC MRA can image very slowly moving blood since it does not suffer the saturation effects of TOF imaging. Finally, PC MRA can be obtained after i.v. gadolinium administration without image degradation because PC MRA does not rely on T1 values to generate the MRA image. Following contrast, TOF MRA is usually limited because gadolinium shortens the T1 and veins become hyper-intense.

The major disadvantage of PC MRA is that it uses a longer TE than TOF MRA. This results in increased intravoxel dephasing and signal loss around stenoses and areas of turbulence. Also, 3D PC MRA has similar spatial resolution but is a longer sequence compared to 3D TOF MRA and is therefore more susceptible to motion artifacts. Consequently, 3D TOF MRA is usually employed to image the head. However, if there is concern that a subacute clot may mimic flow-related enhancement, 3D PC MRA should be performed. Also, because 3D TOF MRA cannot determine flow direction, 2D PC MRA (a much shorter sequence compared to 3D PC MRA) is frequently used to assess collateral retrograde flow in the anterior or posterior communicating arteries in association with severe internal carotid artery (ICA) stenosis or to assess retrograde flow in the basilar artery due to severe stenosis.

In the neck, TOF techniques are preferred over PC techniques due to the latter’s longer scan times, which are needed to provide the same coverage and spatial resolution. Two-dimensional TOF MRA provides superior flow-related enhancement and allows coverage of the entire neck. Compared to 2D TOF MRA, 3D TOF MRA provides superior spatial resolution and is less susceptible to phase dispersion artifacts, but it is more susceptible to saturation effects and cannot cover a large area. Three-dimensional TOF techniques are therefore used to delineate the bifurcation only. Two-dimensional PC techniques are used to evaluate flow direction in the vertebral arteries when subclavian steal is suspected. Also, since
compared to TOF, 2D PC techniques are more sensitive to the detection of very slow flow, they can be used to differentiate high-grade stenosis with a string sign from occlusion.

### 6.2.2 Contrast-Enhanced MRA

Contrast-enhanced MRA (CE MRA) is performed with a rapid, short repetition time (TR, 10 ms) gradient echo sequence following an i.v. bolus of gadolinium. The gadolinium shortens the T1 to less than 10 ms so that opacified vessels are hyperintense. Background tissues, including normally T1-hyperintense structures such as fat and methemoglobin in hematomas, have low signal because they have intrinsic T1 relaxation times of much greater than 10 ms [19].

CE MRA is usually obtained from the arch to the skull base in the coronal plane and is often obtained with a first-pass technique (Fig. 6.10). This requires obtaining the MRA during peak arterial enhancement to avoid venous enhancement. The timing of this arterial phase can be determined by a test bolus or by automatic bolus detection. k-space is filled during peak arterial enhancement in order to maximize image contrast. Another technique known as time-resolved CE MRA acquires the MRA and fills k-space before, during, and after the arterial bolus. This does not require synchronization of the MRA with the injection. In theory, this technique can depict flow dynamics but is limited by the trade-off between temporal and spatial resolution [19].

CE MRA is a reliable modality to image neck vessels but can have poor signal-to-noise ratio at the edge of the imaging volume or have respiratory motion artifact. Respiratory motion artifact limits adequate visualization of the major vessel origins off the arch as well as the origins of the right common, right subclavian, and bilateral vertebral arteries.

CE MRA has several advantages over noncontrast MRA (i.e., PC and TOF MRA): CE MRA can cover a much larger area of anatomy (from the arch to the skull base) in a much shorter acquisition time and is less susceptible to patient motion. CE MRA also has a greater signal-to-noise ratio and less dephasing from turbulence and does not suffer signal loss from saturation effects. CE MRA images the contrast within a vessel lumen and is therefore a more anatomic evaluation, while noncontrast MRA depicts physiology, and anatomy must be inferred from blood flow. This can be misleading when a vessel is not seen on TOF techniques due to reversal of flow or very slow flow (Fig. 6.15).

The disadvantages of CE MRA compared to noncontrast MRA is that the CE MRA data must be obtained during the narrow time window of arterial enhancement and cannot be repeated until the intravascular gadolinium agent is cleared. Thus improper timing (scanning too early or late) results in poor arterial enhancement and an inadequate study that cannot immediately be repeated. Also, the spatial resolution of CE techniques is inferior to that of TOF techniques, and contrast MRA is also minimally invasive as it typically requires a 20-ml power injection of gadolinium contrast at 2 ml/s.

In clinical practice, CE MRA is routinely used to image the great vessel origins and the neck. In general, it does not overestimate carotid bifurcation and other stenoses as much as TOF techniques because it is less susceptible to dephasing from turbulence and does not suffer signal loss from saturation effects. However, many institutions continue to image the neck with both CE and TOF techniques. Due to poorer spatial resolution, CE MRA may underestimate carotid stenosis. TOF techniques are also useful when the arterial bolus is timed incorrectly and when there is unsuspected reversal of flow. For example, in Fig. 6.15, the left vertebral artery appears normal on the CE image and one would assume antegrade flow. However, the vessel is not seen on the 2D TOF flight images, suggesting retrograde flow with saturation of spins due to a superior saturation pulse.

A comparison of the MRA techniques and their typical clinical applications is presented in Table 6.2.

### 6.2.3 Image Processing

Noncontrast and contrast MRA can be postprocessed as a maximum intensity projection (MIP) image. This technique first creates a 3D model of the vessels from the MRA raw data. A set of parallel rays is then drawn from the model and the highest inten-
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sity along each ray is projected to form a 2D MIP image. Multiple projections of the model are created from different vantage points so that viewing sequential MIP images gives the illusion of observing a rotating 3D MRA model.

6.2.4 Extracranial Atherosclerosis and Occlusions

MRA of neck vessels is important in stroke management because extracranial atherosclerosis causes an estimated 20–30% of strokes. The North American Symptomatic Carotid Endarterectomy Trial (NASCET) trial demonstrated that carotid endarterectomy improves survival in symptomatic patients with carotid stenosis of 70–99% [21]. The Asymptomatic Carotid Atherosclerosis Study also suggested that asymptomatic patients with a stenosis of 60% could benefit from endarterectomy [22]. Since then, multiple studies have evaluated the ability of CE and noncontrast MRA to distinguish between nonsurgical (<70%) and surgical (70–99%) stenoses.

CE MRA is generally accurate in evaluation of stenoses. A 2003 study comparing CE MRA and digital subtraction angiography (DSA) showed a sensitivity of 97% and specificity of 95% for stratifying nonsurgical from surgical stenoses [23]. However, a 2004 prospective study also comparing DSA and CE MRA showed a sensitivity of 93% and specificity of 81% for detecting severe stenosis [24]. This poorer performance of CE MRA was attributed to interobserver variability, and this study noted that using CE

Fig. 6.15 a–c

Slow flow or subclavian steal syndrome. Two-dimensional TOF MRA source image (a) and MIP image of the TOF MRA (b) show absent flow and possible occlusion of the left vertebral artery (arrows in b). CE MRA (c), however, shows flow in this vessel (arrows). This discrepancy can occur from slow flow leading to signal saturation on 2D TOF or from reversed flow in subclavian steal syndrome.
<table>
<thead>
<tr>
<th>Technique</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Clinical applications</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2D TOF</strong></td>
<td>Is noninvasive</td>
<td>Typically overestimates vessel stenosis</td>
<td>Is used routinely as the “back up” MRA of the neck in case the CE MRA is suboptimal</td>
</tr>
<tr>
<td></td>
<td>Can image slow flow</td>
<td>Has in-plane artifactual signal loss</td>
<td>Is used routinely in MR venography</td>
</tr>
<tr>
<td></td>
<td>Can image a large volume of tissue</td>
<td>Is susceptible to signal loss from flow turbulence</td>
<td>Can sometimes suggest subclavian steal if combined with CE MRA</td>
</tr>
<tr>
<td></td>
<td>Can be repeated if suboptimal</td>
<td>Has low spatial resolution</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Can be obtained after contrast administration but is slightly degraded by venous signal</td>
<td>Has misregistration artifact on MIP reconstructions</td>
<td></td>
</tr>
<tr>
<td><strong>3D TOF</strong></td>
<td>Is noninvasive</td>
<td>Can image only a small volume due to saturation artifact</td>
<td>Is used routinely to evaluate the circle of Willis to detect intracranial stenoses and occlusions</td>
</tr>
<tr>
<td></td>
<td>Has high spatial resolution</td>
<td>Has poor background suppression; fat or blood may appear bright on the MRA</td>
<td>Can estimate carotid bifurcation stenoses</td>
</tr>
<tr>
<td></td>
<td>Shows complex vascular flow</td>
<td>Cannot image slow flow because of saturation effects</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Is less susceptible to intravoxel dephasing</td>
<td>Is time consuming and susceptible to patient motion</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Can be repeated if suboptimal</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Can be obtained after contrast administration but is slightly degraded by venous signal</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>2D PC MRA</strong></td>
<td>Is able to show the direction and magnitude of flow</td>
<td>Has low spatial resolution</td>
<td>Is used occasionally to determine collateral flow around the circle of Willis</td>
</tr>
<tr>
<td></td>
<td>Does not show high signal artifact from fat or blood</td>
<td>Is more susceptible to turbulent dephasing than TOF MRA</td>
<td>Is used occasionally to determines subclavian steal and abnormal flow direction in the neck</td>
</tr>
<tr>
<td></td>
<td>Can show very slow moving blood and helps differentiate occlusion from near occlusion</td>
<td>Can have aliasing artifact if an incorrect VENC is used</td>
<td>Can detect slow flow if near-occlusion is suspected</td>
</tr>
<tr>
<td></td>
<td>Can be obtained after gadolinium administration</td>
<td>2D PC of the neck takes longer than 2D TOF of the neck with similar coverage</td>
<td>Is used occasionally in MR venography</td>
</tr>
<tr>
<td></td>
<td>Can be repeated if suboptimal</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>2D PC of the circle of Willis is faster than 3D TOF or PC MRA</td>
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</tbody>
</table>
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MRA alone would have misclassified 15\% of cases and would have altered clinical decision-making in 6.0\% of cases [24]. The authors concluded that this was a sufficiently low error rate to support use of CE MRA, but noted that catheter angiography was still the gold standard.

A 2003 study observed that stenoses measured by CE MRA and DSA are tightly correlated by a linear regression analysis ($r=0.967$ with a 95\% confidence interval of 2.8\%) [25]. While this was true for the study overall, the study warned that stenosis measurements by a single CE MRA exam for an individual patient have a larger confidence interval and may not reliably discriminate between small increments of stenosis (e.g., between a 69\% and 71\% stenosis). Nevertheless, in most patients, CE MRA is accurate for stratifying stenoses into the broad categories of surgical versus nonsurgical lesions.

Some reports suggest that noncontrast MRA of the neck is less accurate than CE MRA in discriminating surgical from nonsurgical stenoses. One 2002 study comparing 2D and 3D TOF MRA to DSA reported that 23\% of their patients would have received unindicated endarterectomies while another 33\% would have been improperly denied endarterectomies had noncontrast MRA results been used alone [26]. Several studies from 1992 to 1994 also reported that 2D TOF MRA overestimated up to 48\% of moderate stenoses, erroneously categorizing them as surgical lesions [27] and one study demonstrated that 2D TOF MRA has a sensitivity of 84\% and specificity of 75\% in differentiating surgical from nonsurgical stenoses [28]. However, a number of studies suggest that 3D TOF MRA has a sensitivity and specificity of grading carotid artery stenosis similar to those of CE MRA. One study demonstrated a sensitivity of 94\%
and specificity of 85% for 3D TOF MRA in distinguishing surgical from nonsurgical stenoses [28]. Another study demonstrated Pearson correlation coefficients of 0.94 for CE MRA versus DSA, 0.95 for 3D TOF MRA versus DSA and 0.94 for CE MRA versus 3D TOF MRA [29]. Also, a 2003 meta-analysis combining 4 contrast and 17 noncontrast MRA studies published from 1994 to 2001 reported a pooled sensitivity of 95% and pooled specificity of 90% for MRA’s ability to discriminate between stenoses greater than or equal to 70% [30].

In general, both noncontrast and CE MRA overestimate stenoses when compared with DSA, and this leads to a decrease in specificity. For noncontrast MRA, this is primarily related to signal loss from dephasing and saturation artifacts. Pixel exclusion on MIP images and motion degradation can also artifactually exaggerate the degree of stenosis on both noncontrast and CE MRA [24]. Nearly all studies evaluating MRA rely on DSA as a gold standard since this was used in the NASCET and ACAS trials. Some papers, however, question the accuracy of DSA in grading stenoses, pointing to underestimation of stenosis by DSA compared with 3D rotational catheter angiography [31, 32]. Some have suggested that the specificity measured on MRA studies may be falsely low from using a suboptimal gold standard [31].

Discriminating near occlusion from total occlusion is critical, as the former can indicate urgent surgery while the latter contraindicates surgical treatment. Several studies show that MRA has high sensitivity and specificity for making this differentiation [30, 33, 34]. El-Saden et al. [33], in a retrospective study using noncontrast and CE MRA together, reported a 92% sensitivity for detecting 37 total occlusions and a 100% sensitivity for detecting 21 near-occlusions [33]. A meta-analysis of both CE MRA and noncontrast MRA studies reported a pooled sensitivity of 98% and pooled specificity of 100% in differentiating high-grade stenosis from occlusion [30]. Other studies, however, do not report such high accuracies [35] and many practices still rely on DSA to differentiate definitively occlusion from near-occlusion.

6.2.5 Intracranial Atherosclerosis and Occlusions

In the setting of acute stroke, intracranial MRA can detect areas of stenosis and occlusion as well as determine collateral flow (Fig. 6.16). Defining the location of intracranial vessel pathology is clinically important since an estimated 38% of patients with acute strokes have arterial occlusion seen on DSA [36] (and distal clots are more likely than proximal clots to recanalize following tissue plasminogen activator (t-PA) [37]. As a result, proximal clots are treated more aggressively, sometimes using intra-arterial techniques. In the acute setting, localizing intracranial occlusions is often performed by CT angiogram, but MRA can also depict these findings (Fig. 6.17).

Several studies report that intracranial MRA has a variable reliability of detecting occlusion and stenoses in the acute stroke setting. A 1994 study of TOF MRA compared to DSA in stroke reported 100% sensitivity and 95% specificity for detecting intracranial occlusion [38], and another demonstrated 88% sensitivity and 97% specificity for detecting MCA lesions compared with DSA [39]. A 2002 study, however, showed that, with the addition of contrast to TOF MRA, 21% of vessels initially thought to be occluded on noncontrast TOF MRA were actually patent on CE TOF MRA [40]. Few studies have determined the clinical significance of these MRA findings in stroke, but a recent 2004 study using phase-contrast MRA in acute stroke showed that absent flow in the M1 segment can help predict infarct growth [41].

The degree of collateral flow seen on DSA is an independent radiologic predictor of favorable outcome following thrombolytic treatment [42]. TOF MRA, however, is limited in evaluation of collateral flow. One study found a negative predictive value as low as 53% for TOF MRA’s ability to detect collateral flow when compared with transcranial Doppler [43]. A 2004 study showed that, on TOF MRA, prominence of distal PCA vessels ipsilateral to an M1 occlusion represents collateral blood flow via leptomeningeal vessels [44]. The significance of this finding remains uncertain.
6.2.6 Dissection

Vascular dissection is an important etiology of acute infarction, causing up to 20% of infarcts in young patients and an estimated 2.5% of infarcts in the overall population [45]. Dissection occurs when blood extends into the wall of a vessel through an intimal tear. This may occur in the extracranial or intracranial vessels, the carotid or vertebral arteries, and may be spontaneous or post-traumatic in etiology [46, 47]. Dissections most frequently occur in the carotid artery as it enters the skull base and in the vertebral artery segment from C2 to the foramen magnum. Dissections cause stroke primarily through embolization rather than through flow limitation [48].

Acute dissections show luminal narrowing on MRA and a flap can occasionally appear as a linear low signal defect on MRA. The signal of the intramural blood follows that of parenchymal hematomas, but hemosiderin deposition is not typically seen. Once methemoglobin develops, the wall of the vessel appears hyperintense on fat-saturated T1-weighted images [49] (Fig. 6.18). Chronically, the vessel can occlude, recanalize, show pseudoaneurysms, or become dilated [50].
In a study of 19 internal carotid and five vertebral artery dissections, the MR appearance alone had an estimated 84% sensitivity and 99% specificity for diagnosing carotid dissections and a 60% sensitivity and 98% specificity for diagnosing vertebral artery dissection [51]. Noncontrast MRA (TOF) has low sensitivity (20%) for detecting vertebral artery dissection, but preliminary data suggest that CE MRA may improve the evaluation of vertebral artery dissection [52]. Recently, case reports of false-negatives have been reported for CE MRA in diagnosing dissection [53].

6.2.7 Other Infarct Etiologies

MRA can help determine other etiologies of arterial infarct, including moya moya disease, vasculitis, and fibromuscular dysplasia.

6.2.7.1 Moya Moya

The term moya moya refers to primary moya moya disease and moya moya pattern, associated with an underlying disease such as atherosclerosis or radiation therapy. There is an increased incidence of primary moya moya disease in Asians and in patients with neurofibromatosis or sickle cell disease. Pathologically, there is a progressive occlusive vasculopathy of the supraclinoid internal carotid artery with extension into the proximal anterior and middle cerebral arteries associated with characteristic dilated prominent collateral vessels. Pediatric patients with moya moya disease tend to develop symptoms from acute infarction while adults with moya moya disease more frequently present with symptoms from intracranial hemorrhage into the deep gray nuclei. MRA can depict stenoses and occlusion of the internal carotid, middle cerebral, and anterior cerebral arteries (Fig. 6.19) [54, 55], and the estimated sensitivity and specificity of MRA in diagnosing moya moya disease in one study of 26 patients were 73% and 100% respectively [56]. Furthermore, one recent study reported that MRA depiction of moya moya collaterals in patients with sickle cell anemia was correlated with future cerebrovascular events [57]. MRA is also frequently used in planning vascular by-pass surgery and following response to treatment.
6.2.7.2 Vasculitis

Vasculitis affecting the central nervous system (CNS) represents a heterogeneous group of inflammatory diseases that may be idiopathic or associated with autoimmune diseases, infections, drug exposure, radiation or cancer. Vessel walls are infiltrated by inflammatory cells, and there is increased vasomotor reactivity related to the release of neuropeptides. These properties lead to vessel narrowing. There is also loss of normal endothelial anticoagulant properties and vessels have increased susceptibility to thrombosis. Consequently, patients with vasculitis develop ischemic and thrombotic infarctions. There is also altered wall competence, which can result in dissection or vessel wall disruption with intracranial hemorrhage. MRA is clinically used to screen for vasculitis, but is less sensitive than DSA (Fig. 6.20). One study of 14 patients with suspected vasculitis reported that MRA could detect distal stenoses in vasculitis with a sensitivity of 62–79% and a specificity of 83–87% when compared with a DSA gold standard [58].

6.2.7.3 Fibromuscular Dysplasia

Fibromuscular dysplasia (FMD) is an uncommon idiopathic vasculopathy causing stenoses most often in the renal and internal carotid arteries, and patients with FMD of the neck vessels can present with infarcts or transient ischemic attacks. MRA in FMD can show alternating areas of stenosis (Fig. 6.21) but one study noted that 2D TOF MRA is limited in evaluation of FMD, as slice misregistration artifacts can mimic alternating stenoses [59]. In practice, MRA is useful in distinguishing FMD from vessel dissection or hypoplasia [60].
6.2.8 Venous Infarct

Venous occlusion can lead to infarct through a reduction in cerebral blood flow. Venous infarcts are under-recognized and tend to develop into a hemorrhage [61]. Parenchymal findings on MR include imaging cerebral swelling, venous infarctions, and intracranial hemorrhage. The MR appearance of intravascular clot is variable depending on the age of thrombus and the degree of residual flow. In general, methemoglobin demonstrates hyperintensity on T1-weighted images and there is usually the absence of a flow void on T2-weighted images.

MR venography greatly aids in diagnosing cerebral venous thrombosis and in determining the extent of thrombosis. Typically, a 2D TOF sequence is obtained in the coronal plane [62]. If flow-related enhancement is not seen within a sinus, there should be a high suspicion of sinus thrombosis (Fig. 6.22). However, MRV must be interpreted alongside standard MR sequences because the absence of flow-related enhancement can also be seen in atretic sinuses, in regions of complex flow due to complex geometry or where there is in-plane flow [63]. In addition, T1 hyperintense clot can be confused with flow-related enhancement on TOF techniques but the flow-related enhancement usually has higher signal intensity. Preliminary data also support the use of first-pass contrast-enhanced venography [64].

6.2.9 Conclusion

In the setting of acute stroke, MRA is useful for determining the severity of stenosis, vascular occlusion, and collateral flow. CE MRA and 3D TOF techniques have relatively high sensitivity and specificity in differentiating surgical from nonsurgical carotid stenoses. Three-dimensional TOF MRA is quite sensitive and specific for the evaluation of intracranial proximal stenoses and occlusions. Two-dimensional PC MRA is useful for determining collateral flow patterns in the circle of Willis. MRA is also useful in the determination of stroke etiologies such as dissection, fibromuscular dysplasia, vasculitis, and moyamoya. Currently, MRA is relatively insensitive to the detection of stenoses in distal intracranial vessels but this detection will improve with new MR hardware and software.
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References


Fig. 6.22 a–c
Venous thrombus and venous infarct. 2D TOF MRV (a) and MIP image of the MR venogram (b) show absent flow in the left transverse sinus corresponding to thrombus (arrow). c FLAIR image shows high and low signal at the posterior left temporal lobe corresponding to hemorrhagic infarct