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Diffusion-weighted MRI in acute spinal cord ischaemia

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Abstract Acute spinal cord ischaemia is often undetectable with conventional MRI. Diffusion-weighted MRI (DWI) has been difficult to use in the spine because of susceptibility artefacts. We assessed the diagnostic value of echoplanar DWI for early confirmation of spinal cord ischaemia. We performed conventional MRI and DWI in two men and three women, aged 54–75 years with clinically suspected acute spinal cord ischaemia. Imaging was performed 9–46 h after the onset of symptoms, and 2–9 days later to assess the extent of ischaemic signal change. Spatial resolution of DWI within the spine using standard equipment was poor, but in all patients, early DWI revealed areas of high signal indicating decreased diffusion,

confirmed by measurement of apparent diffusion coefficients. Follow-up MRI showed high signal on T2-weighted images and contrast enhancement at the expected levels. Neurological deficits corresponded with radiological findings in four patients: various syndromes, including isolated bilateral weakness or sensory change and combined deficits, were found. Echoplanar DWI may be helpful for confirmation of spinal cord ischaemia in the acute stage, but follow-up T2-weighted images have superior spatial resolution and correlation with clinical findings and lesion extent.

Keywords Spinal cord infarcts · Diffusion-weighted magnetic resonance imaging

Introduction

In the absence of well-established diagnostic methods, radiological confirmation of acute stroke has until recently been difficult. Conventional methods such as CT and T2-weighted MRI frequently cannot provide a definitive diagnosis in the very acute stage. Diffusion-weighted MRI (DWI), sensitive to water mobility [1,2], is an established method for investigating acute cerebral ischaemia [3,4]. Since the advent of echoplanar imaging (EPI), DWI can be used in clinical practice, but is difficult to apply to the spine with standard EPI techniques, since it is sensitive to changes in the local magnetic field: bone and tissue interfaces cause susceptibility artefacts. Few studies have dealt with the use

of DWI for spinal cord ischaemia [5, 6, 7, 8, 9,10]. We report five consecutive patients with acute spinal cord ischaemia who underwent EPI DWI in the acute stage with follow-up imaging. Our aim was to assess the diagnostic value of DWI in the acute stage and clinical correlations with follow-up T2-weighted imaging.

Materials and methods

We studied five consecutive patients, aged 54–75 years, with clinically suspected acute spinal cord ischaemia, using conventional MRI and DWI. We used a commercial 1.5 tesla imager capable of EPI. The first study was performed 9, 38, 10, 46 and 12 h after the onset of symptoms in patients 1–5, respectively. Follow-up imaging was performed 2–9 days later using conventional sagittal and axial

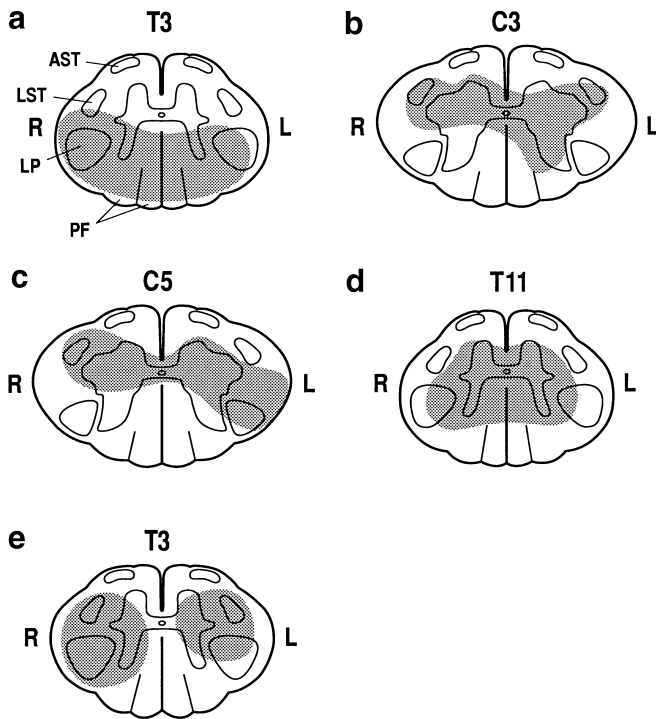
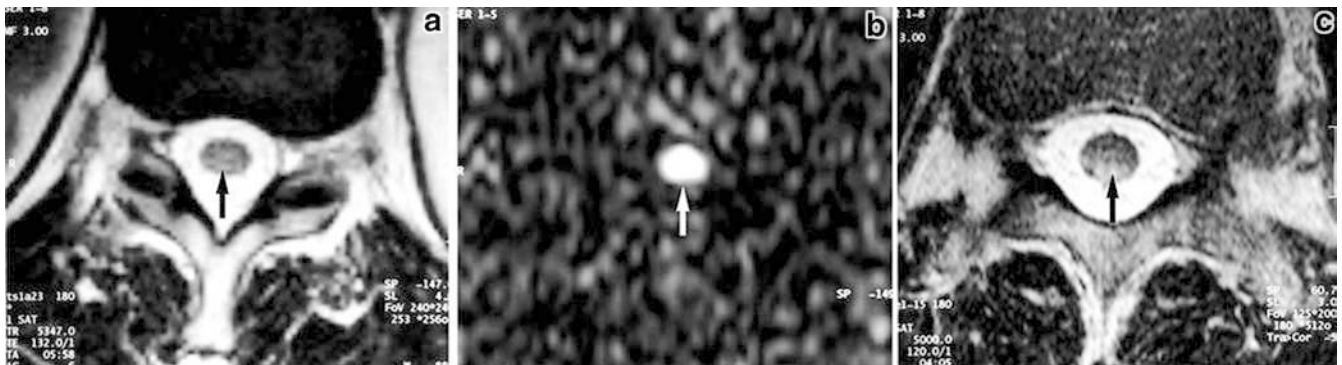


Fig. 1a–e Diagrams of levels with maximum lesion extent as seen on follow-up axial T2-weighted images in each patients. *AST* anterior spinothalamic tract (light touch); *LP* lateral pyramidal tract (motor); *LST* lateral spinothalamic tract (pain and temperature); *PF* posterior funiculus, including gracile and cuneate fasciculi (posture and vibration sense)

T1- and T2-weighted images with contrast enhancement to assess the extent of ischaemic signal change. We used a spine coil, with a standard head isotropic EPI single-shot spin-echo DWI sequence (TR 5700, TE 139.0 ms, slice thickness 5 mm, one acquisition, field of view 240×240 mm, 96×200 pixels, acquisition time 22 s) with b values of 50, 500 and 1000 s/mm². We also obtained T2-weighted images in axial (TR 4888 TE 130 ms, slice thickness 4 mm, field

Fig. 2a–c Patient 1. **a** Axial T2-weighted image 9 h after onset of symptoms shows slightly high signal postero-centrally in the spinal cord (*arrow*). **b** Axial diffusion-weighted image (DWI) at the same time and level, shows high signal in the area corresponding to the spinal cord T12 (*arrow*). **c** Axial T2-weighted image 48 h later more definite high signal at the same site (*arrow*)



of view 130 × 260 mm, 180 × 512 pixels) and sagittal (TR 3500 TE 120 ms, slice thickness 4 mm, field of view 190 × 380, 130 × 512 pixels) planes. Precise localisation in the horizontal plane was performed by overlaying a grid using the imager console software. Apparent diffusion coefficient (ADC) maps were generated from the DWI on a pixel-by-pixel basis with software supplied by the manufacturer. Regions of interest were defined in areas giving higher signal than unaffected portions of the spinal cord, and ADC measured in them on the console. All patients underwent lumbar puncture to exclude haemorrhagic or inflammatory disorders, and had extensive blood analyses, 12-lead ECG, CT of thorax and abdomen, and transthoracic echocardiography in addition to repeated neurological examination.

Results

On DWI and T2-weighted images the spinal cord lesion was thoracic in three patients and cervical in two (Fig. 1). In all patients, the early DWI showed high signal in the spinal cord (Figs. 2, 3) corresponding to decreased diffusion as shown by the ADC. Because of the EPI acquisition technique, severe susceptibility artefacts around the spinal cord caused loss of detail of surrounding structures, especially the bones. The initial ADC were: 0.70, 0.83, 0.87, 0.84 and 0.90 × 10⁻³ mm²/s in patients 1–5, respectively. Three patients had also high signal on T2-weighted images at 9 (patient 1, Fig. 2a), 38 (patient 2), and 12 h (patient 5, Fig. 3a) after the onset of symptoms. We saw no abnormality on DWI (Fig. 2b, 3b) or ADC maps (Fig. 3c, d) adjacent to the ischaemic zones, in areas which proved to be unaffected on follow-up imaging. For the latter, we took care to position the slice at the same level (Fig. 4). After 2–9 days, T2-weighted images (Fig. 2c) showed high signal in all patients contrast enhancement on T1-weighted images at the relevant sites (Fig. 3d); signal changes were still present on DWI, but less marked.

Detailed clinical information is presented in Table 1. The spinal stroke was preceded by transient ischaemic attacks in three patients, in the form of intermittent attacks of thoracic belt-like pain in patient 1 and intermittent painless leg weakness in patients 3 and 4.

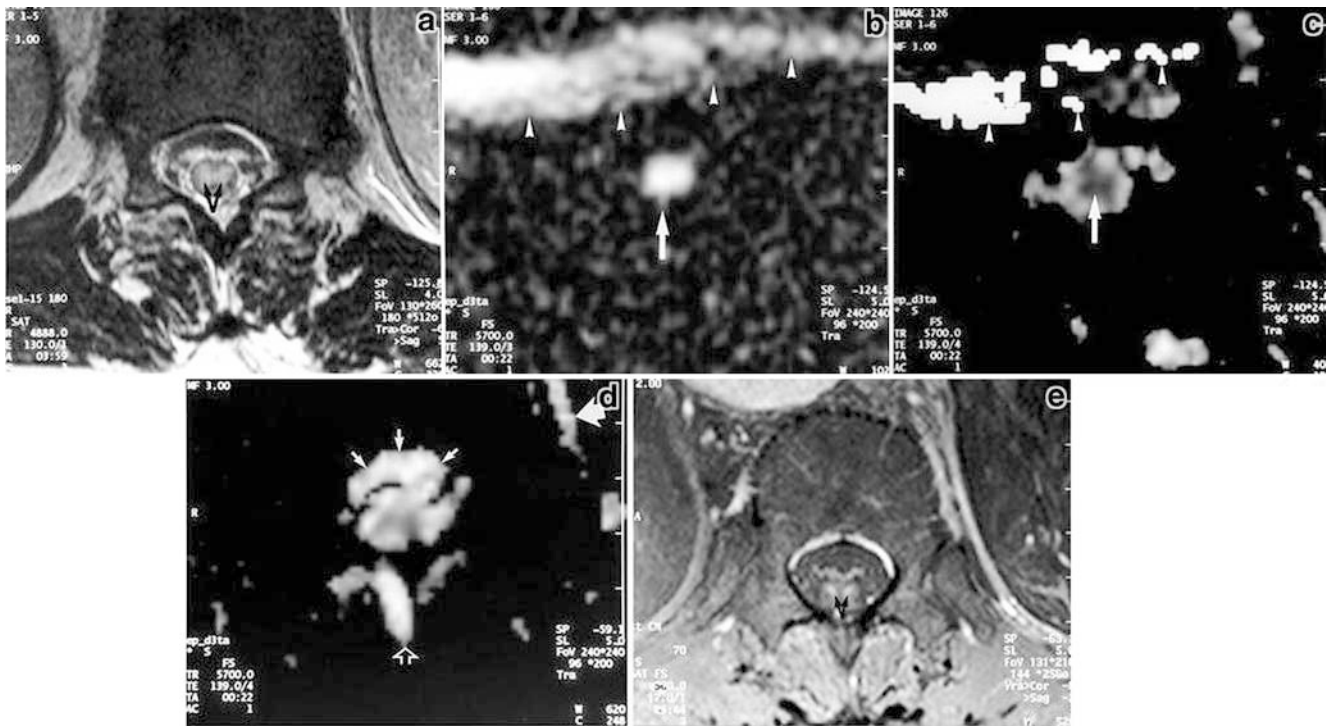


Fig. 3 a–e Patient 5. **a** Axial T2-weighted image at T11 12 h after the onset of symptoms onset shows a large area of high signal centrally in the spinal cord (arrows). **b** Axial DWI at the same time and level shows high signal (arrow). Note strong chemical-shift artefact above spinal cord (arrowheads). **c** Axial ADC (apparent diffusion coefficient) map at the same time and level shows a dark area in the spinal cord (arrow). Note susceptibility-related artefactual signal loss (arrowheads). **d** Axial ADC map 5 days later still shows still low ADC, but less marked. One can also see contour of the left side of the vertebral body (large arrows) and spinous process (open arrow) and the ventral contour of the spinal canal (small arrows). **e** T1-weighted fat-saturated image after 5 days shows contrast enhancement of the central spinal cord, predominantly in the ventral horns (the “snake bite” pattern) (arrows)

The symptoms of spinal stroke developed within from a few minutes to 12 h. In patients 1, 2 and 4, the initial symptom, before the development of sensory and motor deficits, was transient, lancinating, local or radicular back and/or neck pain. In patient 2, atrial fibrillation was the presumed cause, while in patient 5, the spinal stroke occurred after grafting of a thoraco-abdominal aortic aneurysm. In patients 1, 3 and 4, investigations revealed no specific cause; they all, however, had vascular risk factors: patients 3 and 4 had arterial hypertension, patients 1 and 4 were heavy smokers and patient 3 suffered from hypertriglyceridaemia. Neither cerebrospinal fluid (CSF) and extensive blood tests nor other radiological examinations showed inflammatory, neoplastic or other metabolic disturbances other than the vascular risk factors.

Discussion

We have demonstrated the use of EPI DWI in acute spinal cord ischaemia. While recent reports have shown that DWI of the spinal cord is feasible and of potential interest, few have addressed its utility in acute spinal cord infarcts. We showed high signal on DWI in all five patients admitted with acute myelopathy. The diagnosis of spinal cord ischaemia could be confirmed on the basis of the history, initial symptoms, CSF analysis and follow-up MRI.

The ADC were reduced in the areas of high signal on DWI. The time course of DWI changes was similar to that reported in the brain [4], and the ADC seem to correlate with those reported in animal experiments [11] or those obtained with interleaved EPI DWI sequences [12]. Murphy et al. [5] found ADC of $1.2 \pm 0.1 \mu\text{m}^2/\text{ms}$ in preterm infants, which are higher than those in adults. None of our patients had a vertebral body infarct [13,14]. Despite the limited number of patients studied so far, DWI appears to be a sensitive method for early diagnosis of spinal cord ischaemia. However, clinical entities mimicking spinal stroke, such as inflammatory or neoplastic medullary lesions have not, to our knowledge, been fully investigated with DWI. Thus, the specificity of our findings remains to be confirmed. It is probable that, as in the brain, spinal DWI will produce a number of false positives and negatives. In a recent study, Clark et al. [9] found higher diffusion in spinal lesions of multiple sclerosis than in normal volunteers.

Fig. 4a–d Demonstration of axial slice positioning for two examinations. *Left* First examination *right* second. **a, b** Show precise selection of axial slices **c, d** which show the same positioning on scout images and relative to the kidneys and aorta

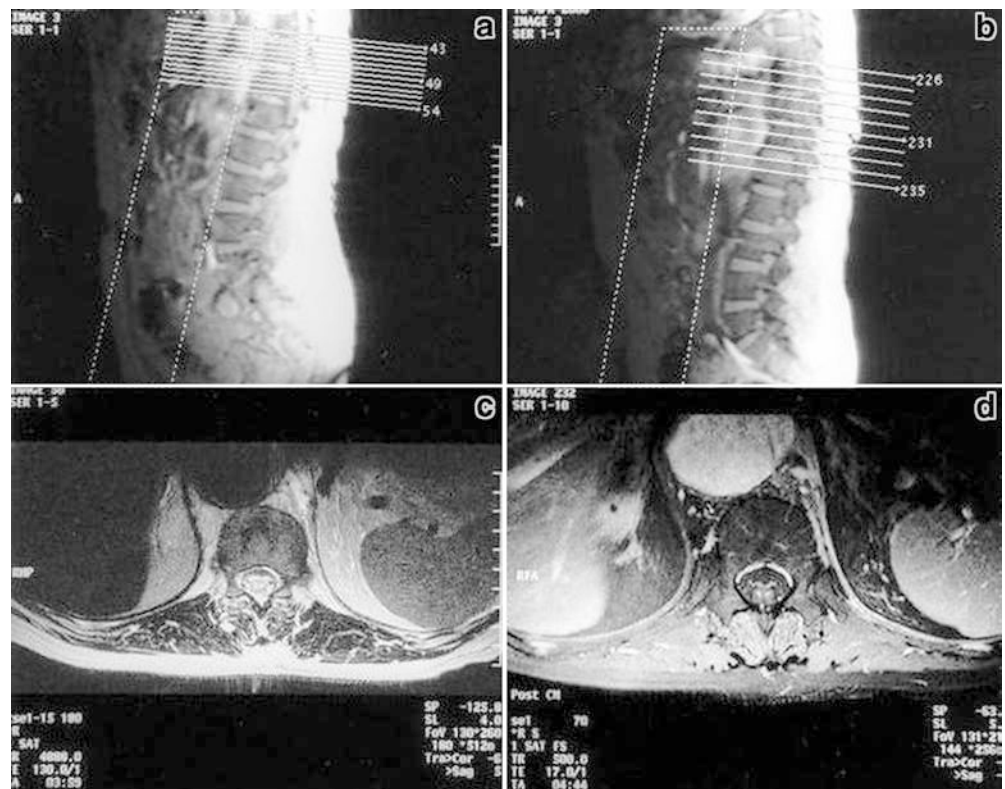


Table 1 Patient characteristics and clinical findings

Patient, age (years)/sex	Extent and site of lesion on axial T2-weighted images	Weakness Site	Grade (right/left)	Sensory level	Sensory deficits	Hyperreflexia	Babinski sign	Bladder dysfunction
1. 60/M	T12, bilateral posterior	Legs Proximal Distal	3/4 4/4	T12	Bilateral loss of vibration, pain and temperature; loss of touch on the right	Right > left	Right	None
2. 75/F	C5, left anterior and right posterior	Arms Proximal Distal Legs Proximal Distal	5/3 4/1 5/4 5/4	T4	Loss of pain and temperature on right	None	Left	None
3. 74/F	T3, bilateral intermediate	Legs Proximal Distal	2/2 2/2	T6	Bilateral loss of pain and temperature	Bilateral	Bilateral	None
4. 58/F	C3, bilateral intermediate	None		C4	Bilateral loss of pain and temperature; loss of postural sensation on left	None	None	Yes
5. 54/M	T11, bilateral central	Legs Proximal Distal	0/0 4/4	None	None	Bilateral	None	None

A drawback of the EPI technique is very strong susceptibility artefacts, which destroy most information concerning neighboring structures such as the vertebrae. In order to image stroke, it is necessary to use high values of b , to retain sensitivity to ischaemia [15]. This can be done using susceptibility-insensitive methods such as

line-scan or HASTE-based DWI [16]. We used a trace image automatically generated by our imager [17]. This has been validated for human cerebral stroke but is more crucial in the spinal cord where tracts are oriented strongly craniocaudally. Even in the brain orientation-dependent ADC measurements can induce errors [18,19].

We confirmed that spinal cord ischaemia can present with a variety of syndromes including isolated bilateral motor or sensory or combined deficits (Table 1). The clinical findings in four patients correlated with maximum axial lesion extent on follow-up T2-weighted images (Fig 1). In patient 1, deficits in vibration sense were consistent with a posterior lesion (Fig 1a) including the lateral pyramidal tract and correlated with the motor deficits. In patients 2 and 4, with atypical presentations, we found a good correlation between clinical and radiological findings (Fig 1b, d). Patient 3, presenting with an anterior spinal artery syndrome showed an anterocentral lesion on axial images (Fig 1c). Patient 5, however, had pure motor deficits whereas MRI showed a central stroke including both dorsal horns (Fig 1e). The lesion on T2-weighted images was up to seven vertebrae longer than indicated by the sensory levels, as has been reported previously [20]. The lesions did not all fit with textbook anterior or posterior spinal artery strokes but, given the known extremely variation in individual blood supply to the spinal cord, this is not surprising.

Early DWI may therefore be especially helpful in clinically atypical syndromes.

Despite extensive investigation, we were unable to determine the aetiology of spinal cord ischaemia in three patients. Aetiological studies suggest that most cases of spinal cord ischaemia are secondary to aortic disease (rupture, dissection, aneurysms, malformations), surgery to the aorta, or severe arterial hypotension [21].

DWI may become important for early diagnosis of spinal cord ischaemia. Further studies of larger numbers of patients are needed to determine its sensitivity and DWI should be performed in other causes of myelopathy, such as inflammation, to establish its specificity. Development of sequences less subject to susceptibility effects will increase image quality and thereby spatial resolution. Perfusion imaging of the spine remains to be implemented, but the demand imposed on MRI gradient systems to obtain high-resolution fast multislice perfusion imaging may render it difficult.

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