Susceptibility-Weighted Imaging as a Distinctive Imaging Technique for Providing Complementary Information for Precise Diagnosis of Neurologic Disorder

신경계 질환에 관한 정확한 진단을 위해 다양한 보완 정보를 제공하는 독특한 영상 기법으로서의 자기화율 강조 영상

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Various sequences have been developed for MRI to aid in the radiologic diagnosis. Among the various MR sequences, susceptibility-weighted imaging (SWI) is a high-spatial-resolution, three-dimensional gradient-echo MR sequence, which is very sensitive in detecting deoxyhemoglobin, ferritin, hemosiderin, and bone minerals through local magnetic field distortion. In this regard, SWI has been used for the diagnosis and treatment of various neurologic disorders, and the improved image quality has enabled to acquire more useful information for radiologists. Here, we explain the principle of various signals on SWI arising in neurological disorders and provide a retrospective review of many cases of clinically or pathologically proven disease or components with distinctive imaging features of various neurological diseases. Additionally, we outline a short and condensed overview of principles of SWI in relation to neurological disorders and describe various cases with characteristic imaging features on SWI. There are many different types diseases involving the brain parenchyma, and they have distinct SWI features. SWI is an effective imaging tool that provides complementary information for the diagnosis of various diseases.

Index terms Magnetic Resonance Imaging; Brain; Diagnostic Imaging
INTRODUCTION

Susceptibility-weighted imaging (SWI) is a high-spatial resolution 3D gradient-echo magnetic resonance (MR) sequence. This imaging sequence emphasizes paramagnetic, diamagnetic and ferromagnetic qualities and is very responsive to the detection of deoxyhemoglobin, ferritin, hemosiderin and bone minerals by distorted local magnetic field (1, 2). The SWI sequence is a further development of the gradient-echo sequence, which is an image generated by the pulse with the flip angle, and reflects the heterogeneity of the surrounding magnetic field. Hemoglobin, deoxyhemoglobin, iron, etc. are the substances that break down the homogeneity of the local magnetic field, which are frequently seen in neurological diseases. In this regard, SWI has been used for the diagnosis and treatment of various neurologic disorders, and the improved image quality has enabled it to provide more useful information to radiologists. In this paper, we will describe about brief information on various neurological disorders and explain the role of SWI sequence in providing additional imaging information for accurate diagnosis.

CONTENTS

INCREASED DEOXYHEMOGLOBIN

SWI can provide a lot of information about the detailed structure of cerebral venous systems and can occur in signal deficit of cortical veins due to magnetic field differences with surrounding structures depending on the concentration of deoxyhemoglobin present in the vein (3). In the normal physiological condition, the deoxyhemoglobin level is measured about 50%, which is observed as dark signal intensity in SWI. In addition, the oxygen supplied to the brain is cut off during brain death, causing the deoxyhemoglobin level to rise further, which can be observed with the engorgement of the cortical veins with dark signal intensity (4, 5). Despite of cortical vein with dark signal can observe in these normal or special conditions, we will take a closer look at some of the disease with the engorged cortical veins with signal deficit.

MIGRAINE

Symptoms of migraines usually include unilateral debilitating headaches associated with photophobia, phonophobia, nausea and vomiting (6). To comprehend the pathogenesis of migraine, it is important to be aware of change in cerebral perfusion when symptoms occur (7). SWI showed prominent hypointense signals of venous vasculature within the perfusion disorder area (Fig. 1). Decoupling the supply and demand of oxygen in hypoperfused cerebral lesion can lead to a relatively increase in deoxyhemoglobin levels in capillaries and veins and a decrease in oxyhemoglobin (8). Karaarslan et al. (7) reported that the area with more noticeable veins on SWI was consistent with the abnormal area on the perfusion MR.

MOYAMOYA DISEASE

Moyamoya disease (MMD) is a rare cerebrovascular disease, causing gradual vascular stenosis of the distal part of the bilateral internal carotid arteries, with formation of a marked reticular compensatory collateral artery called “moyamoya vessels” (9, 10). The disease can
induce cerebral ischemia in children, and causes intracerebral hemorrhage of abnormal blood vessels in adults (11). In MDD patients, multiple deep medullary veins (DMVs) were detected lateral side of the lateral ventricles (12). SWI has been recognized as an imaging tool to identify flow of deep vein in acute or chronic ischemia (Fig. 2) and to indicate increased oxygen extraction in focal ischemia (2). Horie et al. (12) reported that the increase in a number of dilated DMVs, known as “brush signs”, could predict the severity of MMD.

**Fig. 1. Migraine.**
A–C. The diffusion-weighted image (A) and apparent diffusion coefficient map (B) show non-specific findings in the brain parenchyma. The susceptibility-weighted image (C) shows more diffuse prominent venous structures (dark signal) in the left hemisphere than in the right hemisphere, suggesting elevated deoxyhemoglobin levels.

**MEDULLARY VENOUS ENGORGEMENT IN UNILATERAL MCA OCCLUSION**

Middle cerebral artery (MCA) is a branched vessel, originating from lateral aspect of the Circle of Willis, and it is the most commonly pathologically affected vessel in the brain. MCA occlusion is most often caused by embolism or a combination of thrombosis and atherosclerosis. Unilaterally dilated medullary veins observed on SWI has been reported in many patients diagnosed with acute ischemic stroke (Fig. 3) (13). The predominant presence of hypointense veins on SWI is explained by the hypothesis that there is a mismatch between oxygen supply and demand in hypoperfusion tissues, which is thought to be due to the relative increased in deoxyhemoglobin to oxyhemoglobin ratio (8). Payabvash et al. (14) reported that significant correlations between prominent hypointensed thick medullary veins on SWI with arterial occlusion and large infarct volume in acute ischemic stroke.

**ACUTE THROMBOSIS**

**ARTERIAL THROMBOSIS**

Thrombosis in cerebral vessels is one of the most common cause for acute cerebral infarction. In patients with cerebral infarction, recanalization of occluded cerebral arteries by rapid
thrombus removal within 6 hours can increase brain tissue reperfusion and save the ischemic penumbra, which is important for improving patient’s prognosis (15). Intravascular thrombus is seen as a hypointense susceptibility vessel sign (SVS) which has imaging features as a thickened and dark signal intense intravascular artifact (Fig. 4). SVS indicates an increase in deoxyhemoglobin due to a local accumulation in red blood cells (RBCs), which is manifested by the trapping of RBCs in occluded blood vessels (16-18).

VENOUS SYSTEM THROMBOSIS

Thrombosis that occurs in the cerebral venous system is a serious cerebrovascular disease that can cause serious brain injury, such as venous infarction and parenchymal bleeding (19). Cerebral vein thrombosis (CVT) occurs primarily in the venous sinus, which has been reported to be associated with cortical or deep cerebral venous thrombosis (20). Due to the high
sensitivity to the susceptible effects of SWI in the acute phase of thrombosis and cortical CVT, SWI provides an additional diagnostic clue for blood clot detection with conventional MR sequences and magnetic resonance venography (21). Thickened and prominent cerebral cortical veins with hypointensity on SWI, accompanied by hyperintense of nearby brain parenchyma can be diagnosed as presence of CVT with venous hypertension (Fig. 5) (22). In addition, hypointense thrombus is visible in T2-weighted images caused by flow voids in the presence of dural sinus thrombosis. When MRI images are obtained using gadolinium-containing contrast media, thrombi appear as filling defects in contrast-enhanced venous sinuses (empty delta sign) (Fig. 6) (23).

Fig. 3. Medullary venous engorgement in unilateral middle cerebral artery occlusion. A–D. Initial diffusion-weighted images (A, B) show focal, small, patchy, and increased signal intensity in the left posterior internal capsule and posterior frontal paraventricular white matter area (arrows). Susceptibility-weighted images (C, D) show localized prominently visible deep medullary veins in the left frontoparietal white matter area (arrows).
ACUTE HEMORRHAGE

DIFFUSE AXONAL INJURY

Diffuse axon injury (DAI) refers to axons and small blood vessel injuries caused by sudden

Fig. 4. Arterial thrombosis.
A, B. The diffusion-weighted (A) and fluid-attenuated inversion recovery (B) images show high signal intensity in the right frontoparietotemporal cortex white matter area (arrows). It suggests acute infarction in the right internal carotid arterial territory.
C. SWI shows focal thickened linear dark SWI signal intensity (arrow) in the right proximal M1 of middle cerebral artery.
D. Computed tomography angiography shows total occlusion in the right proximal internal carotid artery, (arrow) with poorly visible right internal carotid artery, middle cerebral artery, right anterior cerebral artery, and distal run-off.
SWI = susceptibility-weighted imaging
acceleration or deceleration of the head by external impacts and is one of the main causes of expire and post-accident sequelae (24). In patients with apparent hemorrhage have worse prognosis than without hemorrhage (24). The histological classification of DAI by the distribution of anatomical damages was first proposed by Adams in 1989. In grade I, there is axonal injuries in the white matter of the cerebrum, cerebellum and brainstem; in grade 2, there is localized injuries appear in the corpus callosum as well; and in grade 3, it shows localized injuries at the dorsolateral or rostral part of the brain stem (25). Image findings on post-DAI

Fig. 5. Venous system thrombosis. 
A–D. The susceptibility-weighted images (A–C) show multifocal curvilinear dark signal intensity in both inferior cerebellar folias, with multiple engorged superficial and deep cerebral veins in both posterior cerebellar hemispheres, (arrows) both superficial cerebral cortical veins, both high frontoparietal cortical veins, and deep medullary and internal cerebral veins. It suggests multifocal cerebellar superficial and deep cerebral venous thrombosis. Initial diffusion-weighted image (D) shows focal, small, patchy, high signal intensity in the right corpus callosum splenium, (arrow) suggesting a focal venous infarct.
hemosrrhage in SWI clearly showed the size and extent of the focal hemorrhagic point. Various sized spot-like or nodular, patchy low signal foci are seen scattered in the predilection site (Fig. 7) (26).

**REVERSIBLE CEREBRAL VASOCONSTRICTION SYNDROME WITH SAH**

Reversible cerebral vasoconstriction syndrome (RCVS) can cause severe headaches that can accompany acute neurological symptoms, which are accompanied by segmental contraction of the cerebral artery, which shows resolution within 3 months (27). Thunder-clap headaches occur for less than one minute, accompanied by small localized convexity subarachnoid hemorrhage (22–34%) and intracranial hemorrhage (6–20%) (28, 29). Brain MR findings in RCVS patients are normally seen, but diffuse vasocontractions are observed in cerebral angiography (27). In fluid attenuated inversion recovery images, it can be used to identify the presence of subarachnoid hemorrhage or cerebral edema, and SWI also helps to identify the presence of convexity subarachnoid hemorrhage and intracranial hemorrhage (Fig. 8) (30).
FERROMAGNETIC & PARAMAGNETIC COMPOUNDS

CEREBRAL AMYLOID ANGIOPATHY

CAA is well known for appearing as amyloid protein precipitates in small arteries (31). Although CAA and systemic amyloidosis are less relevant, it is linked to aging, dementia and Alzheimer disease (31). In SWI, multiple dark signal foci distributed in cortical-subcortical area (Fig. 9). This image finding is thought to be caused by two reasons: paramagnetic (hemosiderin deposition) and ferromagnetic (ferritin) compounds. Weakened vascular walls resulting in impaired elasticity by the accumulation of β-amyloid protein in elastic lamina of vessel walls become vulnerable to shock, which causes micro-hemorrhage (32). Studies have also reported that excessive iron increases amyloid and β-amyloid protein condensation and thus cause neurotoxicity. The accumulation of protein can be considered to have occurred signal void in the SWI sequence due to ferromagnetic artifact (33).

Fig. 7. Diffuse axonal injury. A–F. The fluid-attenuated inversion recovery (A) and T2-weighted (B) images show multifocal intermediate or high signal intensity, surrounding edema in both anteroinferior and high frontoparietal anterior temporal white matter cortex areas and in the body and splenium of the corpus callosum area. The susceptibility-weighted images (C–F) show multifocal dark and some curvilinear signal intensity in the anteroinferior high frontoparietal white matter cortex area (arrows).
Brain capillary telangiectasia (BCT) is mostly found incidentally during cerebral MR scans. It is generally considered benign entities that are non-symptomatic and no longer implicit in patient further evaluation. Telangiectasia is a commonly small lesion, usually found in pons in size from a few mm to a few cm, but is not well represented in conventional MR sequences (34). In SWI sequences, BCT is well observed by focal dark signal foci (Fig. 10). It can explain by decreased blood velocity in ectatic vein of a BCT, resulting in increasing the deoxyhemo-
globin concentration by decrease of oxygen saturation of hemoglobin, causing the observed susceptibility effect (35).

DEVELOPMENTAL VENOUS ANOMALY

Developmental venous anomaly (DVA), also known as cerebral venous angioma, is the most commonly found vascular malformation of cerebral veins. DVA is mostly discovered incidentally, and although it is not yet clear as to the cause of the outbreak, it is thought to occur due to arrested development of venous structures (36). The SWI sequence can be much more sensitive to detection of malformation of venous structures than conventional T2 contrast enhancement images (Fig. 11). The signal intensity of the vein is observed darkly in

Fig. 9. Cerebral amyloid angiopathy.
A–D. The susceptibility-weighted images show multifocal dark signal intensity in both cerebellar hemispheres, both basal ganglia, right thalamus, and both cerebral cortex white matter areas. The patient was diagnosed with Alzheimer dementia approximately 10 years ago.
SWI, but with varying signal intensities according to blood flow in venous structures, as relatively higher flow may reduce the amount of deoxyhemoglobin (37).

OTHERS

DUAL RIM SIGN

Dual rim sign refers to two concentric rims appears in the pyogenic brain abscess, surrounding the abscess cavity. Outer one of dual rim sign is hypointense, and the inner one relatively more hyperintense (Fig. 12) (38). The paramagnetic free radicals, metabolic products of macrophages have been suggested as forming the hypointense outer rim, and the hyperintense inner rim is formed by granulation tissues between the necrotic center and fibrocol-
lagenous capsule (38, 39).

CONCLUSION

The use of MR in sensitization diagnosis for various lesions has increased, with the recent development of various MR sequences by rapid technological development. Among the various kinds of MR sequences, the SWI provides additional radiologic clue to the radiologist’s diagnosis about various vascular diseases, hemorrhage, and infectious/inflammatory disease.

Fig. 11. Developmental venous anomaly. A–D. The magnetic resonance FLAIR (A) and susceptibility-weighted (B, C) images show focal, small, lobulated, central high signal intensity, internal septated, marginal dark signal lesion (arrows) with blooming artifact (arrowheads) in the right frontal paraventricular white matter area. It appears to be the accompanying cavernous malformation. Another finding on FLAIR image (A) is focal, small, high signal intensity adjacent to the developmental venous anomaly, with an enhancing vascular structure (arrow) (D). FLAIR = fluid attenuated inversion recovery
lesions. It is thought to be improved of the quality of SWI sequence, it will be necessary to be aware the characteristic radiologic findings for radiologists to make accurate diagnosis and provide proper treatment to the clinician.

**Author Contributions**

Conceptualization, J.B., Y.I.K.; data curation, all authors; investigation, J.B., Y.I.K.; methodology, all authors; project administration, J.B.; resources, all authors; supervision, Y.I.K.; visualization, all authors; writing—original draft, J.B., Y.I.K.; and writing—review & editing, J.B., Y.I.K.

**Conflicts of Interest**

The authors have no potential conflicts of interest to disclose.

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Fig. 12. Dual rim sign.
A–F. The T2-weighted (A) and fluid-attenuated inversion recovery (B) images show localized, relatively smooth marginated, central cavitory lesion [high diffusion-weighted imaging signal intensity (C), with diffusion restriction on apparent diffusion coefficient maps (D)], with marginal wall enhancement in the left occipital white matter and some cortex areas. And there are outer dark susceptibility-weighted imaging (E, F) signal rim (arrows), with slightly uneven thickened enhancing capsule (maximal thickness: approximately 5 mm in the anteromedial portion), surrounding moderate edema, and mass effect. The patient underwent tumor resection and was diagnosed with a brain abscess.
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신경계 질환에 관한 정확한 진단을 위해 다양한 보완 정보를 제공하는 독특한 영상 기법으로서의 자기화율 강조 영상

전병욱1·유인규1*·김태건1·김하연1·황승배2

 자기공명영상 기술의 개발에 따라 다양한 종류의 시퀀스가 개발되어 방사선 진단에 큰 도움이 되었다. 다양한 자기공명영상 시퀀스 중에서 자기화율 강조 영상은 고 공간 분해능 3차원 경사 에코 시퀀스를 발전시킨 것으로 국소 자기장 왜곡에 의한 디옥시헤모글로빈, 페리틴, 해모시데린 및 골, 괴물 검출에 매우 민감하다. 이러한 영상 특징으로 인해 자기화율 강조 영상은 다양한 신경 장애의 진단과 치료에 사용되어 왔으며, 영상 화질이 향상되어 방사선 전문의에게 보다 유용한 정보를 제공할 수 있게 되었다. 다양한 신경 장애에서 발생할 수 있는 자기화율 강조 영상에 나타날 수 있는 다양한 신호의 원리를 설명하고, 독특한 영상학적 특징을 가진 질환 혹은 물질에 대해 임상적 또는 병리학적으로 진단된 환자들에 관하여 각각의 질병에 맞추어 조사하였다. 또한 자기화율 강조 영상에서 각각의 신경 장애에서 보일 수 있는 영상학적 특징에 대해 질환의 전반적인 정보를 함께 요약하여 정리하였다. 뇌 실질 및 주변 조직에 생기는 다양한 신경계 질환들은 자기화율 강조 영상에서 두텁게 구분되는 다양한 영상학적 특징을 보인다. 이에 의해 자기화율 강조 영상은 다양한 보조적 정보를 통해 적절한 진단에 도움을 준다.

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