Magnetic resonance susceptibility weighted in evaluation of cerebrovascular diseases

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Abstract

Background: Cerebrovascular diseases are considered a very hard burden as they may lead to poor outcome, and they are considered the second most common cause of morbidity and mortality after coronary artery disease. They include wide variety of diseases that affect vascularity of brain tissue with the most common one is stroke—either ischemic or hemorrhagic. The aim of the current study was to assess the role of susceptibility weighted imaging (SWI) in imaging of different cerebrovascular diseases and what would be added by SWI to different routine magnetic resonance imaging (MRI) sequences.

Results: Fifty-five patients enrolled in this study, 14 patients had lesions with calcifications, 13 patients had cavernoma, 10 patients had diffuse axonal injury, 11 patients with infarction, 2 patients with AVM, 2 patients with chronic microbleed, 2 patients with hemorrhage, and 1 patient with hemorrhagic tumor, and the result showed that SWI has sensitivity 100%, specificity 60%, and accuracy 91.9% in regard to diagnosis of cavernoma while sensitivity 91.7%, specificity 50%, and accuracy 85.7% in regard to diagnosis of calcification and regarding diagnosis of diffuse axonal injury; SWI has 98.3% sensitivity, 100% specificity, and 98.4% accuracy; finally, in regard to diagnosis of hemorrhagic lesions, SWI has 96.1% sensitivity, 66.7% specificity, and 93.1% accuracy.

Conclusion: SWI is very sensitive in the diagnosis and detection of actual number of vascular malformation like cavernomas than conventional MRI. SWI adds significant diagnostic value to routine MRI sequences in regard to calcification that was nearly limited in its diagnosis by CT. Diagnosis of microbleeds becomes easier and accurate with SWI. Diffuse axonal injury was and still considered a clinical diagnosis, but SWI becomes the gold standard in its imaging diagnosis confirming the clinical one.

Keywords: Susceptibility weighted imaging, Cerebrovascular diseases, Vascular malformation

Background

Cerebrovascular disease includes a variety of medical conditions that affect the blood vessels of the brain and the cerebral circulation. It is considered the second leading cause of morbidity and mortality after coronary artery diseases. There are many causes of cerebrovascular diseases like traumatic brain injury and stroke which are the most common and the medical emergency that requires prompt treatment to minimize brain damage and potential complications, and less common and less severe diseases like cavernous malformation, arteriovenous fistula, amyloid angiopathy, developmental venous anomaly, etc., that are usually asymptomatic until an acute event such as a stroke occurs. The diagnosis of cerebrovascular diseases is clinical with assistance from imaging techniques like computed tomography (CT) scan and magnetic resonance imaging (MRI) [1–3].

MRI uses different sequences for imaging like T1-weighted images (T1W), T2-weighted images (T2W), diffusion weighted images (DWI), proton density images (PDI), and fluid attenuation inversion recovery (FLAIR) in addition to new sequences exploiting higher field strength like arterial spin labelling (ASL) and susceptibility weighted imaging (SWI) [3, 4].
Susceptibility weighted imaging (SWI) is one of the most important of these sequences. It is a fully flow compensated, three-dimensional, high spatial resolution, and gradient echo MR imaging technique. The basic concept of this technique is maintaining phase information into the final image, discarding phase artifacts, and keeping just the local phase of interest. Thus, it utilizes the phase information and signal loss to reveal the susceptibility differences between various tissues and substances like blood products, iron, and calcification [5, 6].

SWI is used as a complimentary tool to conventional MR imaging sequences, and its importance has been demonstrated in various neurological disorders as it is sensitive to both diamagnetic and paramagnetic substances like deoxygenated blood and intracranial mineral deposition which generate different phase shift in MRI data. SWI images can be displayed as a minimum intensity projection that provides high resolution delineation of the cerebral venous architecture, a feature that is not available in other MRI techniques [7–9].

The aim of the current study was to assess the role of SWI in imaging of different cerebrovascular diseases and what would be added by SWI to different routine MRI sequences in the form of comparison between them by various statistical tests like chi-square test, sensitivity, specificity, and accuracy and also by the assessment of the detection rate to finally evaluate some of the clinical applications of SWI.

**Methods**

**Study design and population**

This prospective study was carried on 55 patients (32 males, 23 females) with their age ranged from 1 year and 4 months to 64 years old, at the period between June 2018 and June 2019; the patients had wide variety of cerebrovascular diseases and were examined by conventional MRI sequences and SWI. CT was done in some patients especially with calcification. Ethics committee approved, and informed consent were obtained for all patients or their guardians. Privacy and confidentiality of all patients’ data were guaranteed; all data provision were monitored and used for security purpose only.

**Inclusion criteria**

A. Patients with suspected symptoms of stroke or vascular malformations are as follows:

- Confusion, including trouble with speaking and understanding.
- Headache, possibly with altered consciousness or vomiting.
- Numbness or inability to move parts of the face, arm, or leg, particularly on one side of the body.
- Walking troubles, dizziness, and lack of coordination.
- Vision loss.
- Severe unsteadiness.

B. Patients presented with trauma and non-contrast CT were free or revealed few small hyperdense foci.

C. Patients with past history of brain tumor are presented with new symptoms suspecting complications like hemorrhagic transformation.

**Exclusion criteria**

A. Patients with metallic medical implants (intraocular metallic foreign body, cardiac pace makers, MR non compatible intracranial clips of arterial brain aneurysms).

B. Unfit patients for examination.

- Patient who presented with impaired cognition and uncontrolled movement.
- Patients with some mental disorders.

C. Patients who refused to do the examination.

**Preparation and protocol**

All patients in this study were submitted to complete history taking, clinical examination (by neuropsychiatry physicians), and patient preparation including removing of any detachable metallic implants in patient body or clothes like keys and teeth prosthesis prior to entrance to magnetic area.

The patients were placed in the supine position and the head in the neutral position with the use of a circular polarized head-array coil and ultra gradients. All MRI scans were performed by a MRI 1.5 T unit (GE Signa Explorer) closed magnet right-handed system using a head coil in radiodiagnosis and medical imaging department, with magnetic resonance imaging protocol included Axial T1WI (TR/TE = 400–600/10–20 m/s and FOV = 180 mm), Axial T2WI (TR/TE = 2000–4000/100–120 m/s, and FOV = 180 mm), coronal and/or axial FLAIR images (TR/TE/inversion time (TI) = 4000–6000/140/1200 and FOV = 180 mm), and diffusion weighted imaging (using b value of 0 and 1000) with apparent diffusion coefficient calculation that were obtained by using an axial echo-planar SE sequence and susceptibility weighted imaging (SWI).
Susceptibility weighted imaging (SWI)

Susceptibility weighted images were generated from gradient-echo (GRE) pulse sequences with typical imaging parameters (TR/TE = 6400/30 ms, FOV = 280 mm, acquisition matrix = 256 × 256, slice thickness = 0.2 mm, gap = 2 mm, and scan time = 6 min and 10 s with MIP reconstruction for the image).

A key feature of SWI was that magnitude and phase information were independently processed/displayed as well as combined for diagnostic purposes.

The magnitude image was saved for diagnostic purposes, displaying background tissue with spin-density-like contrast while the raw phase image was dominated by large but slowly changing “macro”-susceptibility gradients due to generalized field in homogeneities as well as distortions due to air and bone at the skull base. Digital high-pass filtering could be used to remove these low-frequency fluctuations, and additional local phase correction algorithms may be employed to reduce artifacts at the skull base. The end result is a filtered phase image that was saved and used for diagnostic purposes.

A phase mask was next created that scales data from the filtered phase images over a 0–1 range to accentuate tissues with different susceptibilities. The magnitude image was digitally multiplied by this phase mask several times until the desired mix of phase information was imparted. The end result was a susceptibility weighted (SW) image, which simultaneously contained both magnitude and phase information and was used for clinical diagnosis. Minimum intensity projections (mIP) of the SW image were typically displayed, which vary in thickness depending on how many of the original 3D source images were stacked together.

A comparison between the SW image and the filter phase was made for interpretation, but firstly we make sure that venous structures display hypointense signal in filter phase, if it is hyperintense, then reverse the window first for correct interpretation.

Both bleeding and calcification displayed hypointense signal in SW image and would be differentiated on the filter phase. Bleeding displayed hypointense signal in filter phase, and calcification displayed hyperintense signal in filter phase; these results were reversed in left handed MR system.

The interpretation of the images was done by two expert radiologist who had experience in neuroradiology 18 and 10 years and in functional radiology for more than 6 years. All cases reviewed blindly, and the decision was taken together.

Statistical analysis

All the data were entered into Excel sheet Microsoft Office Excel 2007 and analyzed statistically using the SPSS Statistical software (version 20.0.0: IBM Corp: Armonk, NY) and XL-STAT (Addinsoft, NY, USA).

Results

This study included 55 patients; the age of the patients ranged from 1 year and 4 months to 64 years old with a mean of 34 ± 14.9 years and median of 43.5 years; fourteen patients (25.4%) had calcifications at different sites with different pathologies (either in calcified oligodendroglioma, teratoma, basal ganglia calcification, calcified granuloma or sporadically), 13 patients (23.6%) had cavernomas, 11 patients (20%) had infarction (9 patients had hemorrhagic infarction and 2 patients had non-hemorrhagic infarction), 10 patients (18.1%) had diffuse axonal injury (DAI), 2 patients (3.6%) had AVM, 2 patients (3.6%) had microbleeds, 2 patients (3.6%) had hemorrhage, and 1 patient (1.8%) had hemorrhagic tumor (Table 1).

Regarding the diagnosis of cavernoma (Fig. 1)

Among the 55 patients examined in this study, 13 patients were diagnosed with cavernoma(either sporadic or inherited) on the basis of specific signals on different MRI sequences with no obvious common risk factor between the patients, but with positive family history of the same pathology in 9 patients. As cavernoma is commonly being multiple in number especially in inherited type so in the current study, the total number of cavernomas detected in 13 patients was 33 lesions.

The Pearson chi-square value for SWI in comparison to conventional MRI sequences (T1WI + T2WI) was 34.803, and the $P$ value was 0.004 while the chi-square value for T1WI + T2WI in comparison to SWI was 47.95, and the $P$ value was 0.086.

Regarding to the final outcome of the reviewed 33 cavernoma lesions depending on the comparison with other conventional MRI sequences and CT, SWI sequence defined 30 lesions (90.9%) as positive and 3 lesions (9.1%) as negative. There were 28 true positive, 2 false positive which were misdiagnosed and confirmed to be calcification by CT, 3 true negative, and no false negative studies. On the other hand, T1WI defined 14
lesions as positive and 19 (57.57%) as negative. There were 12 true positive, 2 false positive, 1 true negative, and 18 false negative studies; also, T2WI defined 21 (63.63%) lesions as positive and 12 (36.36%) as negative. There were 19 true positive, 2 false positive, 1 true negative, and 11 false negative studies with sensitivity, specificity, and accuracy as shown in Table 2.

Regarding the detection of calcification (Fig. 2) Among the 55 patients examined in this study, 14 patients were diagnosed with different pathologies containing calcification, either sporadically or in calcified tumor like teratoma, oligodendroglioma and meningioma, or calcified granuloma; conventional MRI cannot accurately diagnose calcification, and there are usually diverse signals or no signal at calcification sites in T1WI and T2WI. In this study, just detection of altered signal intensity in conventional MRI regarding calcification sites was done and compared with SWI and computed tomography (CT) taken as the control.

The Pearson chi-square value for SWI in comparison to T1WI was 42.507, and the \( P \) value was 0.032. So SWI performed significantly better in detection of calcification lesions than T1WI, and the Pearson chi-square value for SWI in comparison to T2WI was 47.2, and the \( P \) value was 0.078 which is statistically insignificant probably due to decreased number of tested patients.

Regarding the final outcome of the reviewed 14 calcification lesions depending on the comparison with CT, SWI sequence defined 12 lesions (85.7%) as positive and 2 lesions (14.2%) as negative. There were 11 true positive, 1 false positive which were misdiagnosed and confirmed not to be calcification by CT, 1 true negative, and 1 false negative study with sensitivity, specificity, and accuracy as shown in Table 3.

Regarding the diagnosis of diffuse axonal injury (DAI) (Fig. 3) Among the 55 patients examined in this study, 10 patients were diagnosed with DAI with its different grading based on location of scattered microbleeds. The total number of microbleeds in the 10 patients was 62 lesions detected.

The Pearson chi-square value for SWI in comparison to conventional MRI sequences (T1WI + T2WI) was 37.8, and the \( P \) value was 0.0067 while the chi-square value for (T1WI + T2WI) in comparison with SWI was 52.06, and the \( P \) value was 0.2. So SWI performed significantly better in detecting DAI lesions (microbleeds) than conventional MRI.

Regarding the final outcome of the reviewed 62 DAI lesions (microbleeds) depending on the comparison with other conventional MRI sequences and CT, SWI sequence defined 59 lesions (95.2%) as positive and 3 lesions (4.8%) as negative. There were 59 true positive, no false positive studies, 2 (3.2%) true negative, and 1 (1.6%) false negative studies. On the other hand, T1WI defined 36 lesions (58%) as positive and 26 lesions (41.9%) as negative. There were 34 (54.9%) true positive, 2 (3.2%) false positive, 2 (3.1%) true negative, and 24 (38.7%) false negative studies; also, T2WI defined 32 lesions (51.6%) as positive and 30 (48.3%) as negative. There were 30 (48.3%) true positive, 2 (3.1%) false positive, 2 (3.1%) true negative, and 28 (45.1%) false negative studies with sensitivity, specificity, and accuracy as shown in Table 4.

Regarding the detection of hemorrhage (Figs. 4 and 5) In the current study, 16 patients diagnosed with hemorrhage due to several causes like bleeding AVM, hemorrhagic tumor, hemorrhagic infarction, chronic microbleeds, and

| Table 1 Types of pathological lesions included in the current study |
|---------------------------------------------------------------|
| Pathological lesion                          | Number of patients | Types of lesions       | Percentage (%) |
|---------------------------------------------------------------|-------------------|------------------------|----------------|
| Calcification                                              | 14                | 2 Calcified oligodendroglioma | 25.4         |
| 2 Teratoma                                                   |                   |                        |                |
| 3 Basal ganglia calcification                                |                   |                        |                |
| 1 Calcified granuloma                                        |                   |                        |                |
| 6 Sporadic calcification                                    |                   |                        |                |
| Cavernoma                                                  | 13                |                        | 23.6          |
| Infarction                                                  | 11                | 9 Hemorrhagic infarction | 20            |
| 2 Non-hemorrhagic infarction                                 |                   |                        |                |
| Diffuse axonal injury                                       | 10                |                        | 18.1          |
| AVM                                                        | 2                 |                        | 3.6           |
| Chronic microbleeds                                         | 2                 |                        | 3.6           |
| Hemorrhage                                                  | 2                 |                        | 3.6           |
| Hemorrhagic tumor                                           | 1                 |                        | 1.8           |
Fig. 1 Female patient aged 45 years old presented with severe headache and confusion. MRI brain was done and showed right upper frontal cavernoma together with multiple microbleeds that were inconspicuous in T1WI, T2WI, and FLAIR sequences (a, b, and c) but are evident in SWI (d) confirmed by low signal intensity in filter phase (e). Lower level in the same patient showed another right much smaller cerebellar cavernoma that was overlooked by conventional MRI sequences, faint hyperintense in T2WI (f), and faint hypointense in FLAIR (g), and no lesion could be detected in T1 WI (h) but becomes obvious in SWI and filter phase (i and j).
intraparenchymal hemorrhage were examined with routine MRI sequences and SWI.

The Pearson chi-square value for SWI in comparison to conventional MRI sequences (T1WI + T2WI) was 43.06, and the \( P \) value was 0.028 while the chi-square value for (T1WI + T2WI) in comparison to SWI was 48.03, and the \( P \) value was 0.094. So SWI performed significantly better in detecting hemorrhagic lesions than conventional MRI.

Regarding to the final outcome of the reviewed 29 hemorrhagic lesions depending on the comparison with other conventional MRI sequences and CT, SWI

| MRI sequence | Number of lesions | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|--------------|------------------|----------------|----------------|-------------|
|              | Isointense | Hypointense | Hyperintense |              |              |              |
| T1WI         | 19        | 0            | 14          | 40           | 33.3        | 39.4        |
| T2WI         | 12        | 0            | 21          | 63.3         | 33.3        | 60.6        |
| SWI          | 3         | 30           | 0           | 100          | 60          | 93.9        |

\( P \) value 0.004* for SWI and 0.086* for (T1WI + T2WI)

*Chi-square test, \( P \) value < 0.05 is considered statistically significant

Fig. 2 Male patient aged 39 years old, presented with repeated headache and drowsiness with past history of oligodendroglioma that was surgically removed. Current MRI showed right tempo-parietal space occupying lesion with non-specific signal intensity in T1WI, T2WI, and FLAIR sequences (a, b, and c) showing large intervening low signal intensity in SWI (d), high signal intensity in filter phase (e), and corresponding large hyperdense area in CT (f) denoting calcification mostly recurrent calcified oligodendroglioma
Table 3 T1WI, T2WI, and SWI lesion signal intensity of the calcification (number of patients = 14)

| MRI sequence | Number of lesions | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|--------------|-------------------|----------------|-----------------|--------------|
|              | Isointense | Hypointense | Hyperintense | | |
| T1WI         | 7         | 3           | 4              |               | |
| T2WI         | 5         | 4           | 5              |               | |
| SWI          | 2         | 12          | 0              | 91.7          | 50           | 85.7          |

*Chi-square test, *P* value < 0.05 is considered statistically significant

Fig. 3 Male patient aged 28 years old had road traffic accident and presented with disturbed conscious level and Glasgow coma scale = 6. MRI brain showed bilateral subdural hematomas together with multiple microhemorrhage at grey white matter interface, corpus callosum, and around the lateral ventricles denoting diffuse axonal injury that were inconspicuous in T1WI (a, b, and c) but more evident in SWI (d, e, and f) and confirmed to be microhemorrhage by filter phase (g, h, and i) displaying hypointense SI
sequence defined 26 (89.6%) lesions as positive and 3 (10.3%) as negative. There were 25 (86.2%) true positive, 1 (3.4%) false positive study, 2 (6.8%) true negative, and 1 (3.4%) false negative study. On the other hand, T1WI defined 22 (75.8%) lesions as positive and 7 (24.1%) as negative. There were 18 (62%) true positive, 4 (13.7%) false positive, 4 (13.7%) true negative, and 3 (10.3%) false negative studies; also, T2WI defined 21 (72.4%) lesions as positive and 8 (27.5%) as negative. There were 16 (55.1%) true positive, 5 (17.2%) false positive, 6 (20.6%) true negative, and 2 (6.8%) false negative studies with sensitivity, specificity, and accuracy as shown in Table 5.

**Discussion**

Cerebral cavernous malformations can be sporadic or inherited, the latter characterized by multiple lesions. SWI is more sensitive in detecting the accurate number of lesions in inherited type of cerebral cavernous malformations than other sequences. In our study, 13 patients diagnosed with cavernoma, 9 of them have the inherited type with multiple lesions, and 4 have sporadic type with total number of lesions in both types = 33. SWI could detect 30 lesions while T1WI could detect 14 lesions, and T2WI detected 21 lesions with SWI sensitivity 100%, specificity 60%, accuracy 93.9%, and P value 0.004 which was matched with Tan et al. [10] who examined 27 patients with total number of cavernomas 41 lesions stating that SWI was the highest sequence for detection of cavernomas (36 lesions, P value = 0.0023, and sensitivity 98%). Similarly, Sparacia et al. [11] made his research on 42 patients with inherited type CCMs with total number of lesions 57 stating that 51 lesions were detectable by SWI (P value 0.0095 and sensitivity 96.5%). Also, because the incidence of hemorrhage from occult cerebrovascular malformations depends on presence of any previous or old microbleeds or not, so using SWI to detect microbleeds helps to predict any future risks [12].

In regard to calcification, the current study revealed 14 patients diagnosed with different pathologies including calcification with total number of lesions equals 14 interpreted with SWI and other routine MRI sequences and showed that SWI could detect 12 out of 14 lesions (85.7%). T1WI could detect 7 out of 14 lesions (50%), and T2WI could detect 9 out of 14 lesions (64.2) with P value of SWI in detection of calcification 0.032 compared with T1WI and 0.078 compared with T2WI (that made SWI statistically insignificant in detection of calcification compared with T2WI) with sensitivity 91.7%, specificity 50%, and accuracy 85.7%, and this matched with Malhotra et al. [13] who examined 50 patients with 76 focal calcifications and showed that SWI could detect 69 out of 76 lesions (90.7%). T1WI could detect 40 out of 76 lesions (52.6%), and T2WI could detect 53 out of 76 lesions (69.7%) with P value of SWI compared with T1WI that was 0.01 and mismatched with P value of SWI compared with T2WI that was 0.04, and this might be due to increased total number of patients Malhotra examined in his study.

The results of the current study were better than the study conducted by Berberat et al. [14] in which the sensitivity for detection of calcification by magnitude SWI was 86%, however, and in a study conducted by Wu et al. [15], the detection rate of calcification by magnitude SWI was better than our study with detection rate 90.4%, and also the location and size of calcifications perfectly matched with that of CT. In regard to number of lesions that could be detected in DAI patients, SWI was the highest sequence in detection of the accurate number, size, and site of the lesions; as in the current study, the total number of lesions were 62, and SWI could detect 59 lesions (95.2%) with P value = 0.0067, sensitivity = 98.3%, specificity = 100%, and accuracy = 98.4%, which was in close relation to Sinha et al. [16] who showed SWI detection rate = 98.4% with P value = 0.001, sensitivity = 95%, specificity = 98.5%, and accuracy = 96%; also, Babikian et al. [17] demonstrated that SWI is 3–6 times more sensitive than T2 × GE sequences in terms of detecting the number, size, volume, and distribution of hemorrhagic lesions seen in DAI cases. Similarly, Hamdeh et al. [18] emphasized the role of SWI in detecting lesions showing that it was better than conventional MRI sequences and also more helpful in predicting the outcome.

In our study, there were 16 patients presented with hemorrhagic lesions; some of them had multiple lesions, so the total number of lesions was 29. SWI could detect 26 lesions (89.6%) with P value 0.028, sensitivity 96.1%, specificity 66.7%, and accuracy 93.1%, and this in close relation to Lawrence et al. [19]. This study encountered

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**Table 4** T1WI, T2WI, and SWI lesion signal intensity of diffuse axonal injury (number of patients = 10 and number of lesions = 62)

| MRI sequence | Number of lesions | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|--------------|-------------------|----------------|----------------|--------------|
|              | Isointense | Hypointense | Hyperintense |              |              |              |              |              |
| T1WI         | 26      | 0          | 36           | 58.6         | 50           | 58.1         |              |              |
| T2WI         | 30      | 22         | 10           | 51.7         | 50           | 51.6         |              |              |
| SWI          | 3       | 59         | 0            | 98.3         | 100          | 98.4         |              |              |

*Chi-square test, P value < 0.05 is considered statistically significant*
13 patients with total lesion number 30 and SWI detection rate 92% with P value 0.03 and sensitivity 91%.

In regard to the detection of chronic cerebral microbleeds, the current study revealed that 2 patients with chronic microbleeds; SWI could detect the largest number of lesions by 10 lesions while T1WI detected only 4 out of 10 lesions, and T2WI detected 6 out of 10 lesions, so SWI confers greater reliability as well as greater sensitivity for cerebral microbleed detection compared with other MRI sequences, and this matched with Cheng et al. [20] who made a study on 38 patients with chronic microbleeds and proved that SWI should be the

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**Fig. 4** Female patient aged 64 years old presented with hemiparesis and difficulty in speech. MRI brain showed left parietal acute infarction (short arrow) showing low signal intensity in T1WI (a) and high signal intensity in T2WI and FLAIR sequences (b, c) with restricted diffusion, hyperintense SI in diffusion (d), and hypointense SI in ADC (e), and no blooming effect in SWI (f) denoting non-hemorrhagic infarction together with another left basal ganglion (body of caudate) (long arrow) acute infarction with small dots of microhemorrhage seen inside by SWI (f) confirmed by low signal intensity in filter phase (g) denoting hemorrhagic infarction that was overlooked by T1WI.
Fig. 5 Male patient aged 57 years old, presented with seizures and medical history of hypertension. MRI brain was done and showed faint hyperintense curvilinear lesion seen in the right external capsule in T1WI (a) and displayed hypointense signal in T2WI and FLAIR sequences (b, c). SWI showed hypointense signal with blooming effect (d), and filter phase showed hypointense signal (e) denoting right external capsule hematoma.

Table 5 T1WI, T2WI, and SWI lesion signal intensity of the hemorrhage (number of patients = 16 and number of lesions = 29)

| MRI sequence | Number of lesions | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|--------------|-------------------|----------------|-----------------|--------------|
|              | Isointense        | Hypointense    | Hyperintense    |              |
| T1WI         | 7                 | 5              | 17              | 85.7         |
| T2WI         | 8                 | 9              | 12              | 88.9         |
| SWI          | 3                 | 26             | 0               | 96.1         |

*Chi-square test, P value < 0.05 is considered statistically significant
preferred sequence for quantifying cerebral microbleed counts as it detected the largest number of lesions by 136 lesions.

Detection of cerebral microbleeds especially that associated with stroke patients appeared to be important biomarkers for risk of intracranial hemorrhage in case of thrombolytic therapy as presence of more than 5 microbleeds in such patients carry high risk for hemorrhagic transformation as proven by multiple studies like Hale-foglu et al. [2], Chia et al. [22], and Chris et al. [23].

Limitation of the study
Our study shows some limitation. Firstly, all the patients were examined by 1.5 tesla MRI machine; further studies with 3 T MRI machine are recommended for better results. Secondly, the relatively long time of SWI sequence which was obstacle in few patients. Another drawback of this study was the inability to follow some patients after treatment; also, lack of CT study in few patients with calcified lesions decrease our ability to confirm the diagnosis of calcification.

Conclusion
SWI is very sensitive in the diagnosis and detection of actual number of microbleed and hemorrhage in any pathological condition like infarction, diffuse axonal injury, and vascular malformation like cavernoma and arteriovenous malformation; also, calcium with its different compounds are very susceptible substances that make SWI also very sensitive in diagnosis of calcification in various pathological lesions that were difficult to be diagnosed with conventional MRI sequences.

We recommend adding the SWI to the routine MRI in patients suspecting to have hemorrhage, vascular malformation, or calcified lesions.

Abbreviations
SWI: Susceptibility weighted imaging; FOV: Field of view; GRE: Gradient echo; TE: Echo time; TR: Repetition time; CWM: Cerebrovascular malformation; CCM: Cerebral cavernous malformation; DAI: Diffuse axonal injury; ADC: Apparent diffusion coefficient

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Authors’ contributions
WY suggested the research idea, ensured the original figures and data in the work, minimized the obstacles to the team of work, correlated the study concept and design, and had the major role in analysis. RE supervised the study with significant contribution to design the methodology, manuscript revision, and preparation. HA correlated the clinical data of patient and matched it with the findings, and drafted and revised the work. TG collected data in all stages of manuscript and performed data analysis. All authors read and approved the final manuscript for submission.

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Availability of data and materials
The author’s confirm that all data supporting the finding of the study are available within the article, and the raw data and data supporting the findings were generated and available at the corresponding author on request.

Ethics approval and consent to participate
Informed written consents taken from the patients and healthy volunteers, the study was approved by ethical committee of Tanta university hospital, faculty of medicine (32271/04/18).

Consent for publication
All participants included in the research gave written consent to publish the data included in the study. Authors accepted to publish the paper.

Competing interests
The authors declare that they have no competing of interests.

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