Diagnostic report of high-resolution magnetic susceptibility weighted imaging for cerebral vascular amyloid angiopathy-related microbleeds

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Abstract: Objective: To determine the value of susceptibility weighted imaging (SWI) in the differential diagnosis of cerebral amyloid angiopathy (CAA)-related microbleeds. Methods: The clinical data of 33 patients with highly suspected CAA-related microbleeds were retrospectively analyzed. All subjects underwent conventional T1-weighted spin-echo imaging, T2-weighted fast spin-echo imaging and SWI. The investigators evaluated the diagnostic value of different magnetic resonance imaging (MRI) sequences by assessing the imaging features of CAA-related microbleeds. Results: The microbleeds of the subjects were mainly located in the cerebral cortex and subcortical regions. Fourteen cases were also found to have chronic hematoma remote hemorrhage in the cerebellar hemisphere. Seven cases had subacute or remote hemorrhage in the basal ganglia, 3 cases in the thalamus and another 3 cases in the brain lobe. In the conventional MRI sequences, CAA-related microbleeds manifested as low or intermediate signals in T1 weighted imaging (T1WI); in some cases short T1 signals could be seen. In T2 weighted imaging (T2WI), the lesions were heterogeneous pattern with mixed areas of high and low signal intensity. In the SWI sequence, more hemorrhagic foci were seen, showing low signal areas with clear margins. And punctate or patchy high signals were seen in the center of some lesions. The count of microbleeds showed by SWI was significantly higher than that of T1WI or T2WI (P<0.01). The Kappa consistency test showed that the SWI sequence was consistent with T2WI and T1WI sequences, and the Kappa values were 0.63 and 0.54, respectively. The P values were less than 0.01. Conclusion: For patients with cerebral vascular amyloid microbleeds, high-resolution SWI can achieve better diagnostic results, showing clear lesions with higher detection sensitivity.

Keywords: High-resolution susceptibility weighted imaging, cerebral amyloid angiopathy, microbleeds, magnetic resonance imaging

Introduction

Susceptibility weighted imaging (SWI) is a high-resolution three-dimensional technique that uses not only the magnitude information of magnetic resonance imaging (MRI) signals, but also the phase information compared with the traditional T2* weighted gradient-recalled echo imaging (GRE) [1]. The phase and magnitude data are combined in subsequent processing steps by minimum intensity projection technique to increase the sensitivity or contrast of paramagnetic tissue, which includes vein structure and cerebral microbleeds (CMB) [2].

Cerebral amyloid angiopathy (CAA) is associated with perivascular rupture, which is caused by deposition of amyloid β-protein in blood vessels. Previous autopsies have shown that the prevalence of CAA in Alzheimer’s disease is 70% to 90% [3]. CAA is an important cause of intracerebral hemorrhage (ICH) in the elderly, accounting for 10-20% of spontaneous ICH. With the increase of life expectancy, it’s expected that the prevalence of CAA-related ICH will follow [4]. This study evaluates the value of SWI in the diagnosis of CAA-related microbleeds.

Materials and methods

Patients

The clinical data of 33 patients with highly suspected CAA-related microbleeds were retrospectively analyzed. The patients were admitted to Linyi Central Hospital from July 2015 to
SWI for the diagnosis of cerebral microbleeds

**Table 1. Modified Boston diagnostic criteria**

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Definite CAA</td>
<td>Full post-mortem examination reveals lobar, cortical, or cortical/subcortical hemorrhage and pathological evidence of severe cerebral amyloid angiopathy</td>
</tr>
<tr>
<td>Probable CAA with supporting pathological evidence</td>
<td>Clinical data and pathological tissues (evacuated hematoma or cortical biopsy specimens) demonstrate a hemorrhage as mentioned above and some degree of vascular amyloid deposition</td>
</tr>
<tr>
<td>Probable CAA</td>
<td>A. Patient is 55 years or older&lt;br&gt;B. Appropriate clinical history&lt;br&gt;C. MRI or CT findings demonstrate: Multiple hemorrhages restricted to the lobar, cortical, or cortico-subcortical regions without another cause, or a single lobar, cortical, or cortico-subcortical hemorrhage and focal or disseminated cortical superficial siderosis without another cause</td>
</tr>
<tr>
<td>Possible CAA</td>
<td>A. Patient is 55 years or older&lt;br&gt;B. Appropriate clinical history&lt;br&gt;C. MRI or CT findings demonstrate: A single lobar, cortical, or cortico-subcortical hemorrhage without another cause, or focal or disseminated cortical superficial siderosis without another cause</td>
</tr>
</tbody>
</table>

Note: CAA, Cerebral amyloid angiopathy.

Figure 1. Cerebral microbleeds. A: T1WI, T1 weighted imaging; B: T2WI, T2 weighted imaging; C: SWI, susceptibility weighted imaging.

July 2018. This study was approved by the ethics committee of Linyi Central Hospital. Inclusion criteria: Patients were diagnosed with CAA according to Modified Boston Criteria (see Table 1) [5]; patients had provided written informed consent; patients underwent MRI examination including T1WI, T2WI and SWI; patients were at their first attack (Figure 1). Exclusion criteria: Patients had intracranial space-occupying lesions; patients had ICH caused by other reasons such as hypertension, trauma, and vascular malformation.

**Methods**

The Achieva 3.0 T-type superconducting MRI system produced by Philips was used to perform high-resolution SWI and conventional MRI sequences, including T1-weighted spin-echo imaging and T2-weighted fast spin-echo imaging. The patient’s head was fixed with a sponge pad before examination to avoid motion artifacts. The SWI parameters were as follows: TR = 34 ms, TE = 49 ms, slice thickness = 6 mm, slice gap = 1 mm, FOV = 24 cm × 24 cm, matrix = 256 × 256. The acquired images were transmitted to the Achieva MR System Release 2.6.3.6 workstation for analysis. The phase mask and magnitude images were generated simultaneously, and the SWI images were acquired through low-pass filtering, phase masking, multiple constant multiplications and minimum intensity projection.

**Observation indicators**

The images of this study were evaluated by two experienced MRI diagnosticians. The evalua-
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The data were analyzed using STATA 14.0 software. Enumeration data are expressed as number (n); quantitative values were expressed as mean ± sd. Differences between groups were compared using paired t-tests, χ² tests or Kruskal-Wallis tests; and the diagnostic consistency was determined by Kappa tests. The statistical significance level α = 0.05 with a correction for multiple comparisons (α/3 = 0.017).

Results

General information

The age range of the 33 patients is 59-87 years with an average of 67.9±7.6 years. There are 17 males and 16 females. The microbleeds were mainly located in the cerebral cortex and subcortical areas. Except the microbleed lesions, 14 cases were also found to have remote hemorrhage in the cerebellum. Seven cases had subacute or remote hemorrhage in the basal ganglia, 3 cases in the thalamus and another 3 cases in the brain lobe.

Image quality

In the conventional MRI sequence, CAA-related microbleeds manifested as low or intermediate signals in T1 weighted imaging (T1WI); in some cases short T1 signals could be seen. In T2 weighted imaging (T2WI), the lesions were a heterogeneous pattern with mixed areas of high and low signal intensity. The count of microbleeds showed by SWI was significantly higher than that of T1WI or T2WI, presenting a typical spotted or patchy low signal area.

The number of cases of each sequence based on the Likert scale is shown in Table 2. Statistical analysis showed χ² = 2.375 and P = 0.3049; the difference among the three sequences was not significant. The proportion of images with image quality equal to or greater than 3 points were 57.58%, 60.61% and 72.73% for the three sequences, respectively. There was no significant difference among the three groups (χ² = 1.8333, P = 0.40).

Distribution and count of microbleed lesions

The distribution and count of microbleed lesions on qualified T1WI, T2WI and SWI images are shown in Table 3. The microbleeds were mostly distributed in the cortical and subcortical areas. There was no statistically significant difference in distribution of the microbleeds (χ² = 1.95, P = 0.98). The mean number of microbleeds shown by T1WI, T2WI and SWI were 14.95±3.26, 16.60±4.11 and 22.42±4.72, respectively. The paired t-test showed significant differences between the three sequences (all P<0.01).

Diagnostic consistency

Seventeen patients with both qualified SWI and T2WI images were selected. The diagnostic results of the two sequences were examined using the Kappa consistency test. Statistics showed that Kappa = 0.63 and P<0.01, indicating the consistency of SWI and T2WI was good. Similarly, 15 patients with both qualified SWI and T1WI images were subject to the Kappa consistency test. The results showed Kappa = 0.54 and P<0.01, indicating the consistency was good. See Tables 4 and 5.

Discussion

CMB are rounded micro-hemosiderin deposits around blood vessels that are common in elderly patients and associated with neuropsychiatric disorders. CAA involves cerebral vascular amyloid deposition, which is classified into several types according to the amyloid involved, among which sporadic amyloid β-protein type in

Table 2. Number of cases of each sequence based on the Likert scale (n)

<table>
<thead>
<tr>
<th>Sequence</th>
<th>5 points</th>
<th>4 points</th>
<th>3 points</th>
<th>2 points</th>
<th>1 point</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1WI</td>
<td>33</td>
<td>4</td>
<td>8</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>T2WI</td>
<td>33</td>
<td>5</td>
<td>7</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>SWI</td>
<td>33</td>
<td>7</td>
<td>9</td>
<td>8</td>
<td>7</td>
</tr>
</tbody>
</table>

Note: T1WI, T1 weighted imaging; T2WI, T2 weighted imaging; SWI, susceptibility weighted imaging.
SWI for the diagnosis of cerebral microbleeds

CAA is most common in elderly and Alzheimer’s disease patients [6]. CAA can cause hemorrhagic lesions, ischemic lesions, and subacute leukoencephalopathy.

The causes of CMB are complex, which mainly include hypertension, cerebral ischemia, and CAA. CMB also may be caused by infective endocarditis (IE), thrombotic thrombocytopenic purpura, traumatic brain injury, subcortical infarction, and leukoencephalopathy [7-9]. Sparacía et al. found 91 CMB lesions from 15 patients with end-stage organ failure, 59 (64.84%) were supratentorial lobar distribution, 17 (18.68%) were supratentorial non-lobar distribution, and the remaining 15 (16.48%) were distributed subtentorially [10]. Malhotra et al. studied 66 IE patients who had CAA at the same time, he found that in 11 (16.67%) patients, CMB was the only neuroimaging abnormality, and most CMBs measured between 1 and 3mm. Within that size range, many CMBs can be detected by SWI but not by GRE T2* [11].

SWI is very sensitive to paramagnetic substances such as deoxygenated blood, blood products, iron and calcium. SWI obtains a three-dimensional reconstruction image of intracerebral vein, which can detect bleeding as early as 6 hours, and can reliably detect acute intraparenchymal hemorrhage and subarachnoid hemorrhage. SWI can also detect early hemorrhagic signs in the infarction, and help understand the cerebral hemodynamics after stroke, which is helpful for diagnosis of cerebral venous thrombosis. This sequence is useful for detecting microbleeds in various situations.

| Table 3. Distribution and count of microbleed lesions on qualified T1WI, T2WI and SWI images (n) |
|-----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| Sequence       | Number of cases | Cortico-subcortical | Basal ganglia | Thalamus | Cerebellum | Brain stem | Total |
| T1WI           | 19              | 268              | 3             | 4        | 6          | 284         |
| T2WI           | 20              | 309              | 5             | 6        | 8          | 332         |
| SWI            | 24              | 495              | 7             | 8        | 11         | 538         |

Note: T1WI, T1 weighted imaging; T2WI, T2 weighted imaging; SWI, susceptibility weighted imaging.

| Table 4. Display of brain microbleeds by three sequences (n) |
|-----------------|----------------|----------------|----------------|----------------|
| Sequence       | T1WI | T2WI |
| SWI Positive   | 223   | 279 |
| SWI Negative   | 25    | 8   |

Note: T1WI, T1 weighted imaging; T2WI, T2 weighted imaging; SWI, susceptibility weighted imaging.

| Table 5. Kappa consistency test |
|-----------------|----------------|----------------|----------------|----------------|
| Sequences       | Consistency    | Kappa | Standard error | Z   | P    |
| SWI/T1WI        | 77.93% | 0.54 | 0.05 | 11.46 | <0.01 |
| SWI/T2WI        | 82.23% | 0.63 | 0.04 | 14.89 | <0.01 |

Note: T1WI, T1 weighted imaging; T2WI, T2 weighted imaging.

296 microbleeds were observed in 54 patients, of which 38 (70.37%) showed lobar distribution, 13 (24.07%) were non-lobar distribution, and the remaining 3 (5.56%) were mixed distribution; suggesting CMBs in patients with Alzheimer’s disease were mainly distributed in the brain lobe, and the number of CMBs in the brain was positively correlated with age, course of disease and cognitive functional declines [12].

Conventional MRI can show non-specific white matter lesions, multifocal cerebral infarction, or brain atrophy. However, CMBs associated with white matter lesions are difficult to differentiate. Although positron emission tomography (PET) can detect CAA through amyloid-binding ligands, such as Pittsburgh Compound B, it does not distinguish between vascular and parenchymal amyloid deposition. Gradient echo (GE) imaging is considered to be a method of detecting hemorrhagic changes in CAA [13]. However, nearly 25% of CAA patients do not show CMBs in T2* GE sequence [14]. Malhotra et al. studied 66 IE patients who had CAA at the same time, he found that in 11 (16.67%) patients, CMB was the only neuroimaging abnormality, and most CMBs measured between 1 and 3mm. Within that size range, many CMBs can be detected by SWI but not by GRE T2* [11].

such as vasculitis, autosomal dominant arterial disease, subacute infarction and leukoencephalopathy, amyloid angiopathy, andBinswanger’s disease. This sequence also helps to diagnose vascular malformations and perinatal cerebral vascular damage. CMBs appear as <5 mm low signal points in the SWI due to the enhanced $T_2^*$ effect of paramagnetic substances such as hemosiderin or deoxyhemoglobin [15]. CAA-related inflammation (CAA-ri) is a rare and treatable variant of CAA that lacks specific imaging and clinical features, previously requiring an invasive brain biopsy to confirm. Kusakable et al. reported a CAA-ri patient with CT showing low-density lesions in the right occipital cortex. MRI showed a low signal on T1WI, a high signal on T2WI, and no enhancement on contrast-enhanced T1WI. However, T2*-GRE and SWI showed extensive cortical microbleeds, which were confirmed by biopsy [16]. Some scholars examined 9 patients with CAA; GRE detected 1146 CMBs, SWI detected 1432 CMBs. CMB scorers have better reliability with SWI (intraclass correlation coefficient, 0.87), but only moderate reliability with GRE (inter-class correlation coefficient, 0.52) [17]. SWI can not only detect CMBs, quantitatively detect iron changes, but also distinguish it from calcification [18]. In elderly patients with lacunar infarction, the SWI showed a higher positive rate for CMBs, followed by GRE-T2*WI and other conventional sequences like T1WI, T2WI and T2FLAIR. To evaluate the effect of the modified GRE, Nandigam studied the relationship between the CMB detection rate and different sequences, section thickness and magnetic field strength in patients with CAA. With the other parameters unchanged, the positive rate of CMB is moderately increased if using SWI, having thinner sections and higher field strength. SWI is more likely to find CMBs with smaller diameters. In a regular thick slice, GRE only recognizes 33% of CMBs seen in thin slices by SWI [19, 20].

In SWI, artificial recognition of CMBs takes a long time, with limited reliability and reproducibility, which is prone to misdiagnosis. In order to solve that problem, researchers modified the software to improve the efficiency of diagnosis. Fazlollahi et al. proposed a computer-aided CMB detection technology based on machine learning with a new multi-scale Laplacian of Gaussian filtering approach. In 66 CMB patients, the CMBs were manually divided into two levels, “likely” or “confirmed”, to verify the effectiveness of the method. Among the total population, the technology achieved 87% sensitivity and an average false detection rate of 27.1%. The sensitivity of the “confirmed” group was 93% and the false detection rate was 10% [21]. This is similar to another technique of automatic CMB detection based on independent subspace analysis and clustering [22]. SWI’s automatic CMB detection technology improves reliability, reduces internal score variation and evaluation time, which is superior to existing technology and expected to improve expert screening efficiency.

In conclusion, for patients with CAA-related microbleeds, high-resolution SWI can achieve better diagnostic results, showing clear lesions and higher detection sensitivity.

Disclosure of conflict of interest
None.

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References


