Usage of SWI (susceptibility weighted imaging) acquired at 7T for qualitative evaluation of temporal lobe epilepsy patients with histopathological and clinical correlation: An initial pilot study.

Benjamin Y M Kwan  
*Department of Medical Imaging, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada*

Fateme Salehi  
*Department of Medical Imaging, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada*

Pavlo Ohorodnyk  
*Department of Medical Imaging, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada*

Donald H Lee  
*Department of Medical Imaging, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada*

Jorge G Burneo  
*Epilepsy Program, Department of Clinical Neurological Sciences, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada*

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Authors
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Benjamin Y.M. Kwan b, Fateme Salehi b, Pavlo Ohorodnyk b, Donald H. Lee b, Jorge G. Burneo d, Seyed M. Mirsattari d, David Steven d, Robert Hammond e, Terry M. Peters a,b,c, Ali R. Khan a,b,c,*

a Imaging Research Laboratories, Robarts Research Institute, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada
b Department of Medical Imaging, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada
c Department of Medical Biophysics, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada
d Epilepsy Program, Department of Clinical Neurological Sciences, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada
e Department of Pathology and Laboratory Medicine, Schulich School of Medicine and Dentistry, Western University, 1151 Richmond St. North, London, Ontario, N6A 5B7, Canada

Abstract

Objectives: Ultra high field MRI at 7 T is able to provide much improved spatial and contrast resolution which may aid in the diagnosis of hippocampal abnormalities. This paper presents a preliminary experience on qualitative evaluation of 7 T MRI in temporal lobe epilepsy patients with a focus on comparison to histopathology.

Methods: 7 T ultra high field MRI data, using T1-weighted, T2*-weighted and susceptibility-weighted images (SWI), were acquired for 13 patients with drug resistant temporal lobe epilepsy (TLE) during evaluation for potential epilepsy surgery. Qualitative evaluation of the imaging data for scan quality and presence of hippocampal and temporal lobe abnormalities were scored while blinded to the clinical data. Correlation of imaging findings with the clinical data was performed. Blinded evaluation of 1.5 T scans was also performed.

Results: On the 7 T MRI findings, eight out of 13 cases demonstrated concordance with the clinically suspected TLE. Among these concordant cases, three exhibited supportive abnormal 7 T MRI findings which were not detected by the clinical 1.5 T MRI. Of the ten cases that progressed to epilepsy surgery, seven showed concordance between 7 T MRI findings and histopathology; of these, four cases had hippocampal sclerosis. SWI had the highest concordance with the clinical and histopathological findings. Similar clinical and histopathological concordance was found with 1.5 T MRI.

Conclusions: There was moderate and high concordance between the 7 T imaging findings with the clinical data and histopathology respectively.

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using 7 T MR imaging have shown its superior diagnostic benefits in various pathologies including multiple sclerosis, cerebrovascular diseases, aneurysms, cavernous malformations, polymicrogyria, and brain tumors [11,12]. T2*-weighted MR images acquired at 7 T have previously been shown to depict hippocampal subfield structures as small as 100 μm [13]. However there are drawbacks to higher field strength imaging, including greater signal dropout in areas of local susceptibility differences (sinuses), and a higher level of inhomogeneities in the magnetic fields, which can cause distortions and signal loss artifacts.

The aim of the current study was to 1) determine the ability of 7 T MR imaging (and particularly susceptibility weighted images (SWI)) to detect hippocampal and mesial temporal lobe abnormalities in patients before temporal lobe resection and 2) evaluate concordance of 7 T MRI findings with post-operative histopathological results which can serve as a basis for validation of imaging findings.

2. Methods

Ethics approval was obtained through our institutional review board including participation of patients under 18 years of age. All human and animal studies have been approved by the appropriate ethics committee and have therefore been performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. Patients (male or female) with a history of drug-resistant TLE aged 16–65 years old were recruited for this study. Exclusion criteria included patients with severe coexisting or terminal systemic disease and those unsuitable for MRI evaluation. Thirteen patients undergoing evaluation for epilepsy were scanned using the 7 T MRI scanner. Each patient also underwent standard clinical 1.5 T MRI, prolonged video-EEG, and neuropsychological testing, as part of the clinical epilepsy evaluation at an academic tertiary care center. Seven patients also underwent invasive EEG electrode placement. Clinical data were collected for each patient, including age at surgery, gender, suspected epilepsy type, clinical 1.5 T MRI reports, EEG reports, subdural electrode recording reports, clinical follow-up notes post resection and corresponding histopathology reports if applicable.

Ultra high field data were acquired on a 7 T neuroimaging optimized MRI scanner (Agilent, Santa Clara, CA, USA/Siemens, Erlangen, Germany) using a 16-channel transmit-receive head coil array constructed in-house. The imaging sequences used for this study were a multi-echo gradient-echo sequence with six echoes acquired with a 0.5 mm in-plane resolution (TR = 40 ms, TE = 4.57 ms, echo spacing = 4.89 ms, flip angle = 13 degrees, NEX = 1, matrix = 256 × 360, 80 slices, slice thickness = 1.5 mm, in-plane resolution 0.5 mm, FOV = 128 × 180 × 120 mm, acquisition time = 12 min), with slices acquired perpendicular to the long axis of the hippocampus in a coronal oblique orientation, and a T1-weighted MPRAGE sequence (matrix = 256 × 512 × 172, resolution = 0.58 × 0.43 × 1 mm, acquisition time = 5:42 min). For the multi-echo images, the magnitude images for each of the six echoes were averaged to produce a T2*-weighted image. SWI were generated from the multi-echo images using a frequency mask derived from the unwrapped and filtered phase images, using a process described in detail previously [14]. These volumes (T1, T2*, SWI), three for each patient, had randomized identification numbers assigned and were converted to DICOM format for blinded review by a fellowship trained neuroradiologist with over 30 years of experience. Additionally, each volume was processed to remove shading artifacts using a non-uniformity intensity normalization filter (N4) [15], which we refer to henceforth as N4, to assess the impact of viewing images with and without shading artifact removal.

Clinical MRI data was acquired on a 1.5 T MRI scanner (GE, USA). The imaging protocol for epilepsy imaging at this institution included T2 axial (2 mm contiguous slices, matrix = 256 × 256 mm, NEX = 1, TR = 12,666 ms, TE = 49 ms, FOV = 22 cm), FSEIR coronal (3.5/1, matrix = 288 × 256 Zipped to 512 × 512, TR = 5116 ms, TE = 15 ms, TI = 150 ms, FOV = 22 cm), gradient echo coronal (5/1, matrix = 256 × 192 mm, NEX = 2, FOV = 20 cm), 3D FLAIR coronal (with 1.8 mm partitions, reconstructed at 0.9 mm intervals, TR = 6000 ms, TE = 122 ms, TI = 1872 ms, FOV = 24 cm), 3D T1 axial (with 2 mm partitions, reconstructed at 1 mm intervals, FOV = 24 cm, matrix = 256 × 256 mm) and diffusion axial ((6 directions, 80, B1000), 5 mm contiguous, matrix = 128 × 192, FOV = 24 cm, TR = 8100 ms, TE = 88.8 ms).

Each of the patient’s 7 T MRI data was reviewed by the neuroradiologist using a qualitative grading scale while blinded to clinical information. Volumes were graded for scan quality and the presence of artifact. Qualitative grading of the mesial temporal lobe internal architecture (including note of signal abnormality and/or architectural abnormalities), mesial temporal lobe size and temporal neocortical architecture was performed (Table 1). The rest of the brain architecture was briefly evaluated for any gross abnormalities. The results of the neuroradiologist were converted into a single rating for each category by taking the most abnormal grade assigned for each category among the six volumes evaluated. Additionally, the grades of ‘normal’ or ‘possibly normal’ were merged into a single rating value of “normal” and the grades of ‘possibly abnormal’ or ‘definitely abnormal’ were merged into a single rating value of “abnormal”. A second reader (a senior radiology resident) also evaluated the 7 T MRI scans using the qualitative grading scale and assigned a single rating for each category while taking into account all the volumes. A Cohen’s kappa was calculated between the two readers (GraphPad Software, La Jolla, CA, USA and StatsToDo, Australia).

The neuroradiologist also blindly evaluated the 1.5 T MRI data in a similar fashion. Resected specimens were processed according to established protocol. Briefly, specimens were fixed in 10% buffered formalin. Fixed

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Qualitative rating scale.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meshal temporal lobe internal architecture (including signal abnormalities)</td>
<td></td>
</tr>
<tr>
<td>Meshal temporal lobe size</td>
<td></td>
</tr>
<tr>
<td>Temporal neocortical architecture</td>
<td></td>
</tr>
<tr>
<td>Rest of brain architecture</td>
<td></td>
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</tbody>
</table>
material was serially blocked, sectioned and examined by routine histological stains (H&E) and by immunoperoxidase for the expression of neuronal (NeuN) and glial (GFAP) antigens. Stained sections were examined by a neuropathologist and digitized.

Comparison of 7 T MRI findings (including location of abnormal findings) as interpreted by the neuroradiologist to the clinical data (including clinical history, multidisciplinary epilepsy group decisions, EEG reports and subdural electrode recording reports for seizure activity localization). Concordant correlation to histopathology was defined as any identified abnormal imaging finding with a corresponding abnormal histopathological finding on the ipsilateral side. Concordant HS was defined as abnormal mesial temporal lobe internal architecture and abnormal mesial temporal lobe size with corresponding MTS on histopathology. Clinical follow-up data was collected in regards to seizure free status if the patient underwent epilepsy surgery. Additionally, a similar comparison was made as detailed above but individually using each of the 6 volumes of data alone (T1, T2*, SWI, and their corresponding non-uniformity corrected data). Comparison of 1.5 T MRI findings was also performed in a similar fashion.

3. Results

Thirteen patients were included in the study (7 males, 6 females). Ten patients underwent epilepsy surgery (5 males, 5 females, average age of 33.4 years; median age 31.5, range 18–50) and clinical follow-up was performed for an average of 25.6 months. Seven of ten cases undergoing epilepsy surgery had concordance between 1.5 T MRI findings with histopathology. Nine out of 13 cases had concordance of suspected clinical epilepsy syndrome with 1.5 T MRI findings.

Interrater agreement between the neuroradiologist and the second reader (for the 7 T MRI findings) calculated using the Cohen’s Kappa was 0.6094; SE 0.0971 (95% CI 0.4191 to 0.7998) with the percentage of observed agreements at 86.5%.

3.1. Quality of scans and presence of artifacts

Of the 78 volumes inspected (six volumes per patient), three were rendered non-diagnostic, due to the predominant artifacts of susceptibility and shading. Susceptibility artifacts were observed, particularly involving signal drop out affecting the inferior temporal lobe. Shading artifacts were partially mitigated on N4 correction, although ten of 37 corrected volumes were still deemed to be affected. Motion artifact affected three of the 13 cases.

3.2. Concordance of clinical findings to 7 T MRI findings

Eight of the 13 cases demonstrated concordance with the suspected clinical epilepsy syndrome as determined by the clinical management team (including data from neurological presentation, EEG and/or subdural electrode recordings) with the 7 T MRI findings. Among these concordant cases, three were deemed abnormal by the 7 T MRI findings that were supported by the clinical data, but not detected by the routine 1.5 T clinical MRI (cases 10, 11 and 12) (Fig. 2). Subset analysis of clinical concordance per volume revealed that the SWI volumes had the highest clinical concordance with eight of 12 cases (case 7 had a non-diagnostic SWI volume). T2* and T2* N4 volumes had the second highest clinical concordance with seven of 13 cases.

3.3. Concordance of surgical pathology to 7 T MRI findings

Ten subjects underwent epilepsy surgery, of which seven had concordant histopathology with 7 T MRI findings and of these four cases had concordant HS (Fig. 3). In one out of the ten cases (case 2), while there was no structural cause for the TLE in the histopathology of the resected specimen, there was nevertheless complete concordance of 7 T MRI with the histopathology in this case. Three cases were discordant with 7 T MRI findings (cases 4, 7, 8). The results of each case are discussed in Table 2. Analysis of histopathological concordance per volume revealed that the SWI volumes had the highest histopathological concordance with seven of nine cases (case 7 had a non-diagnostic SWI volume). T2* N4 volumes had the second highest histopathological concordance with seven of ten cases.

4. Discussion

Previous studies have shown that 7 T MRI allows higher resolution depiction of the complex hippocampal anatomy [8,9,16–19]. Breyer et al. and Henry et al. [9,19] applied imaging at 7 T in patients with clinical diagnosis of TLE and corresponding hippocampal abnormality on clinical 1.5 T or 3 T imaging. Breyer et al. had shown confirmation of HS (that was diagnosed at 1.5 T) in all 6 of their patients on 7 T MRI [9]. However no pathological correlation was available for confirmation of the imaging findings in these two studies [9,19]. Coras et al. and Zucca et al. [20,21] have performed studies focusing on 7 T MRI imaging of resected specimens in hippocampal sclerosis and focal cortical dysplasia which reveal the potential of high correlation between ex vivo imaging findings and histopathology of the resected specimens. Our study aimed to compare in vivo 7 T MRI imaging findings with histopathology.
The overall concordance of 7 T MRI findings and histopathology was high. There was complete concordance in patients with pathological proven HS suggesting that 7 T MRI can consistently detect this abnormality. This work can serve as a basis for validation of imaging findings at 7 T MRI. Discordant cases (cases 4, 7 and 8) had histopathology consisting of cortical dysplasia/gliosis, mild dentate fascia dispersion/gliosis and focal cortical dysplasia type Ia/gliosis suggesting that 7 T MRI may still not have enough resolution to routinely detect these subtle findings. It is interesting to note that an abnormality was detected on 7 T MRI in these discordant cases but on the contralateral side (on both non-uniformity corrected and uniformity corrected sequences). These three patients have had good control of their seizures following resection which suggests that the 7 T MRI findings are false positives. Shading or susceptibility artifact (even after uniformity correction) may have played a role in this potential misinterpretation; this raises caution that imaging abnormalities should be interpreted in the clinical context. Additionally, as the 7 T MRI were interpreted while blind to clinical data, directed search with clinical suspicion may have a role in increasing detection rates of subtle abnormalities. SWI and T2*-weighted non-uniformity-corrected volumes had the best concordance with histopathology; it is interesting to note that Breyer et al. also found that T2*-weighted imaging at 7 T was valuable for anatomical delineation. [9].

Another point to note is that 1.5 T MRI performed similarly in both clinical and histopathological concordance to 7 T MRI. There was a slight apparent edge to 1.5 T MRI on clinical concordance, however this is likely a false positive coincidence as the suspected abnormality yielded normal histopathology (case 2). Overall, no clear advantage of 7 T MRI is noted for qualitative evaluation as compared to 1.5 T. Subjectively, the image resolution at 7 T was felt to be of higher quality.

The limitations of this study include the small sample size and lack of an age and gender matched control group. More false positives may have been introduced since histopathological concordance was defined as any abnormal 7 T imaging finding with a corresponding ipsilateral abnormal histological finding, however this decision was made as the experience with 7 T MRI is relatively new and in this initial study a higher sensitivity approach was opted for to maximize the potential detection capabilities of 7 T MRI. Another potential limitation was that there may have been recall during blinded evaluation of the 1.5 T and 7 T MRI scans, however this was minimized as the 1.5 T scans were evaluated approximately 12 months after the 7 T MRI scans. Directed interpretation with clinical data may enhance the detection abilities at 7 T. Additionally, a potential limitation is that a 3 T MRI comparison group was not available, as 3 T MRI was not available for clinical use at our institution during this study. Although it is important to note that 1.5 T MRI may still be more frequently used than 3 T MRI at many clinical centers.

5. Conclusion

7 T MRI in patients presenting with epilepsy for pre-surgical workup may reveal findings not detected on lower field imaging. In patients undergoing epilepsy surgery, 7 T MRI findings had high correlation in our small series with the histopathology of the resected specimen. Pathologically proven hippocampal sclerosis was detected by 7 T MRI in all 4 patients which suggests it has the ability to reliably detect this finding. Discordance involving subtle findings such as focal cortical dysplasia and gliosis suggests that 7 T MRI may not reliably detect these findings and directed search with clinical information could be helpful. This experience also raises a cautionary flag to interpret 7 T MRI findings within the clinical context as false positive findings were suggested in the discordant cases; more experience with high field artifacts is needed to ensure correct interpretation. SWI and T2*-weighted volumes showed the highest clinical and histopathological concordance lending further credence that these sequences have superior anatomical depiction at high field strengths; it may be beneficial to consider the addition of these sequences for 7 T MRI in epilepsy workup. Qualitative hippocampal subfield analysis may be a potential application. Although a small study, no clear advantage of 7 T over 1.5 T MRI was observed for clinical and histopathological concordance. Future work with larger numbers and accompanying histopathology will be needed to determine the utility of 7 T MRI for epilepsy imaging and verifying imaging findings.

Authors' contributions

Kwan: Protocol/project development, Data collection or management, Data analysis, Manuscript writing/editing
Salehi: Protocol/project development, Data collection or management, Data analysis, Manuscript writing/editing
Ohorodnyk: Protocol/project development, Data collection or management, Data analysis, Manuscript writing/editing
Lee: Protocol/project development, Data collection or management, Data analysis, Manuscript writing/editing
Burneo: Data analysis, Manuscript writing/editing
Mirsattari: Data analysis, Manuscript writing/editing
Steven: Data analysis, Manuscript writing/editing
Hammond: Data analysis, Manuscript writing/editing
Table 2
Summary of findings in patient group (MTL: mesial temporal lobe; MTS: mesial temporal sclerosis; NA: not applicable; ATR: anterior temporal resection; AHC: amygdalohippocampectomy).

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Age at surgery</th>
<th>1.5 T MR findings</th>
<th>7 T MR findings</th>
<th>Scalp EEG localization</th>
<th>Subdural electrode findings</th>
<th>Surgery performed</th>
<th>Pathology findings</th>
<th>Follow up duration (months)</th>
<th>Engel classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>42</td>
<td>Right and left mesial temporal lobe internal architecture abnormality</td>
<td>Right mesial temporal lobe internal architecture abnormality, right fornix atrophy</td>
<td>Right temporal</td>
<td>Right MTL</td>
<td>Right ATR, AHC</td>
<td>Gliosis (temporal neocortex, hippocampus and amygdala)</td>
<td>17</td>
<td>I</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>24</td>
<td>Right MTS, Left temporal neocortical architecture abnormality</td>
<td>Normal</td>
<td>Left temporal</td>
<td>Left MTL</td>
<td>Left ATR, AHC</td>
<td>Normal</td>
<td>7</td>
<td>I</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>45</td>
<td>Right MTS</td>
<td>Right MTS</td>
<td>Multifocal</td>
<td>Right MTL</td>
<td>Right ATR, AHC</td>
<td>Gliosis (temporal lobe), <strong>mesial temporal sclerosis</strong> Mild cortical dysplasia and mild gliosis (temporal neocortex), normal hippocampus</td>
<td>39</td>
<td>I</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>31</td>
<td>Right mesial temporal internal architecture abnormality</td>
<td>Left MTS</td>
<td>Bitemporal</td>
<td>NA</td>
<td>Right ATR, AHC</td>
<td>Glialosis (temporal neocortex)</td>
<td>18</td>
<td>I</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>32</td>
<td>Left MTS</td>
<td>Left MTS</td>
<td>Left temporal</td>
<td>NA</td>
<td>Left ATR, AHC</td>
<td>Glialosis (temporal neocortex), <strong>mesial temporal sclerosis</strong></td>
<td>27</td>
<td>I</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>NA</td>
<td>Normal temporal lobes</td>
<td>Right MTS</td>
<td>Left temporal, parietal</td>
<td>Right temporal</td>
<td>NA</td>
<td>No resection</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>18</td>
<td>Left temporal neocortical architecture abnormality</td>
<td>Left MTS</td>
<td>Left MTL</td>
<td>Left ATR, AHC</td>
<td>Glialosis (temporal neocortex, hippocampus and amygdala); mild dentate fascia dispersion</td>
<td>36</td>
<td>I</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>M</td>
<td>23</td>
<td>Normal temporal lobes</td>
<td>Right mesial temporal lobe internal architecture abnormality</td>
<td>Left frontotemporal</td>
<td>Left MTL</td>
<td>Left ATR, AHC</td>
<td>Glialosis (temporal neocortex, hippocampus and amygdala), focal cortical dysplasia type la (temporal neocortex)</td>
<td>18</td>
<td>I</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>49</td>
<td>Right mesial temporal lobe internal architecture abnormality, Left periventricular heterotopia</td>
<td>Right MTS, left mesial temporal lobe size abnormality</td>
<td>Multifocal</td>
<td>Right MTL</td>
<td>Right ATR, AHC</td>
<td>Glialosis (temporal neocortex, hippocampus and amygdala)</td>
<td>23</td>
<td>II</td>
</tr>
<tr>
<td>10</td>
<td>M</td>
<td>NA</td>
<td>Left mesial temporal internal architecture abnormality, Left periventricular heterotopia</td>
<td>Right mesial temporal lobe internal architecture abnormality</td>
<td>Multifocal</td>
<td>NA</td>
<td>No resection</td>
<td>NA</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>F</td>
<td>50</td>
<td>Left mesial temporal internal architecture abnormality, left temporal neocortical architecture abnormality</td>
<td>Left mesial temporal lobe size abnormality</td>
<td>Left temporal</td>
<td>NA</td>
<td>Left ATR, AHC</td>
<td>Glialosis (temporal neocortex, amygdala and hippocampus)</td>
<td>36</td>
<td>IV</td>
</tr>
<tr>
<td>12</td>
<td>F</td>
<td>NA</td>
<td>Left mesial temporal lobe size abnormality</td>
<td>Left mesial temporal lobe size abnormality</td>
<td>Bitemporal, greater on right</td>
<td>Bitemporal</td>
<td>No resection</td>
<td>NA</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>M</td>
<td>20</td>
<td>Left MTS, left occipital lobe encephalomalacia (remote)</td>
<td>Left MTS, left occipital lobe encephalomalacia</td>
<td>Left temporal, occipital</td>
<td>Left MTL</td>
<td>Left ATR, AHC</td>
<td>Glialosis (temporal neocortex and amygdala), <strong>mesial temporal sclerosis</strong></td>
<td>35</td>
<td>I</td>
</tr>
</tbody>
</table>

**References**


