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ORIGINAL RESEARCH

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BACKGROUND AND PURPOSE: The detection of microbleeds differs strongly between studies, due to differences in scan protocol. This study aims to compare the visualization of microbleeds with 3D T2*-weighted imaging at 1.5T with 3D dual-echo T2*-weighted imaging at 7T.

MATERIALS AND METHODS: Thirty-four patients (29 male; mean age, 58 ± 12 years) with atherosclerotic disease from the Second Manifestations of ARTerial Disease study were included. 3D T2*-weighted imaging at 1.5T and dual-echo T2*-weighted imaging at 7T were done in all patients. The presence and number of definite microbleeds were recorded on minimal intensity projections. Interand intraobserver reliability was assessed with Cohen κ test and the ICC. The difference in presence and number of microbleeds was tested with the McNemar test and Wilcoxon signed rank test.

RESULTS: The interobserver ICC at 7T was 0.61 and the intraobserver ICC was 0.94, whereas at 1.5T the interobserver ICC was 0.50 and the intraobserver ICC was 0.59. Microbleeds were detected in significantly more patients on 7T (50%) than on 1.5T scans (21%) (P = .001). The number of microbleeds was also higher at 7T (median, 0.5; range, 0–5) than on 1.5T (median, 0.0; range, 0–6) (P = .002).

CONCLUSIONS: 3D dual-echo T2*-weighted imaging at 7T results in better and more reliable detection of microbleeds compared with 3D T2*-weighted imaging at 1.5T.

ABBREVIATIONS: GRE = gradient recalled-echo; ICC = intraclass correlation coefficient; MARS = Microbleed Anatomical Rating Scale; MIP = minimum intensity projection; PRESTO = principles of echo shifting with a train of observations; SMART = Second Manifestations of ARTerial disease; SNR = signal-to-noise ratio; SWI = susceptibility-weighted imaging; TE1 = first echo; TE2 = second echo

Recent studies show that cerebral microbleeds are commonly detected on MR imaging in the general population as well as in specific patient populations, such as patients with cerebral small vessel disease, Moyamoya disease, and other neurologic pathologies. ¹⁻⁸ Simultaneously, the evidence about the clinical relevance of microbleeds increases. Cerebral microbleeds seem to be direct markers of vascular disease, because they are associated with hypertensive vasculopathy and cerebral amyloid angiopathy and also with white matter lesions and lacunar infarcts. ^{1,8-10} Although still under debate, the presence of microbleeds could be of importance in patients with ischemic stroke receiving anticoagulation, because it might indicate a higher risk of future intracerebral hemorrhage. ¹¹⁻¹⁵ Furthermore, some studies have shown that microbleeds are associated with cognitive impairment, functional dependence, and death. ^{10,16,17}

The number of microbleeds detected differs strongly between studies, even between studies with similar study populations. In population-based studies, eg, prevalences range from 5% to 23%.^{3,7} Differences in lesion prevalence are probably not only related to the characteristics of the different study populations but also to differences in the MR protocol,

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such as differences in scan technique, TR, TE, flip angle, band width, section thickness, and spatial resolution. ^{18,19}

Microbleeds are paramagnetic, which induces a susceptibility effect on the MR imaging scan. This leads to a fast decay of local T2*-weighted MR imaging signal intensity because of a local inhomogeneity of the field induced by the microbleeds. This makes T2*-weighted GRE very sensitive to microbleeds. 9,10,20 One of the factors in the imaging protocol that influences the visualization of microbleeds is the TE. A longer TE gives more time for dephasing, which enhances the susceptibility effect. This so-called blooming effect causes the microbleeds to appear as hypointense spots that are larger than their actual size. It has been shown that prolonging the TE leads to an increase in diameter of these lesions and also to the detection of an increased number of microbleeds.²¹ However, microbleeds also can be obscured by overlapping structures with a high susceptibility effect, such as veins (deoxyhemoglobin) or the basal ganglia (ferritin deposition). With a longer TE, these structures also increase in size, making it harder to distinguish the microbleeds. Therefore, both a short TE and a long TE may have benefits for the visualization of microbleeds. Besides the TE, the spatial resolution also is important for the visualization of microbleeds. With a higher resolution, partial volume effects are minimized and smaller microbleeds can be visualized.²² Another factor that influences the visualization of microbleeds is magnetic field strength. So far, most studies investigating microbleeds used a 1.5T MR imaging scanner. However, studies comparing 1.5T and 3T show a higher number of microbleeds at 3T. 18,23 As the susceptibility effect scales with the magnetic field, scanning at higher field strengths improves the sensitivity to microbleeds. At ultrahigh field

strengths, such as 7T, the susceptibility effect further increases as does the SNR. This increase in SNR can be used for higher spatial resolution with increased conspicuity of small hemosiderin deposits that may be obscured by partial volume effects at lower field strengths operating at lower spatial resolution.²⁴ Furthermore, scanning at ultrahigh field strengths enables the use of dual-echo T2*-weighted imaging for the visualization of microbleeds, whereas at lower field strengths this is probably not feasible because the susceptibility effect may not be large enough to visualize the microbleeds at the short TEs. The use of a dual-echo sequence at 7T has been shown to combine the benefits of a short TE and a long TE for the visualization of microbleeds in one sequence.²⁵ It has not been established whether the use of a dual-echo sequence at 7T increases the number of detected microbleeds compared with the commonly used single-echo T2*-weighted imaging at lower field strengths. The purpose of our study was to compare the visualization of microbleeds with T2*-weighted imaging at 1.5T with dual-echo T2*-weighted imaging at 7T and assess the reliability of the detection of microbleeds with the 2 field strengths.

Materials and Methods

Patients

Patients from the SMART study,²⁶ without contraindications for 7T MR imaging, were consecutively included between July 2008 and December 2009. The objectives of the SMART study are to determine the prevalence of vascular risk factors and concomitant arterial disease and to study the incidence of future cardiovascular events and its predictors in patients newly referred to our hospital with atherosclerotic disease. The SMART study and the 7T imaging were approved by the medical ethics committee. Written informed consent was given by all patients. In total, 38 patients could be included in our study. The scans of 3 patients were used for a training session of the 2 observers and therefore were excluded from the final analysis. In one patient, >20 microbleeds were seen on the 7T scan. This was considerably more than in all other patients; the sum of all microbleeds in the remaining 34 patients on the 7T scan was 36. To prevent the results depending highly on the scan of this single patient with many microbleeds, this patient was considered as an outlier and excluded from the analysis. Therefore, the analysis was done on 34 patients.

MR Imaging

MR imaging was performed on a 1.5T whole-body system (Gyroscan ACS-NT; Philips Medical Systems, Best, the Netherlands) and a 7T whole-body system (Achieva; Philips Healthcare, Cleveland, Ohio). In all patients, a 3D T2*-weighted imaging scan at 1.5T and a 3D dual-echo T2*-weighted MR imaging scan at 7T were performed on the same day (MR imaging parameters shown in Table 1).

Because the dual-echo sequence at 7T was originally designed as combination of an arteriogram and a venogram, the TE1 was used to make an inflow arteriogram. To control for intravoxel dephasing of flowing blood, the TE1 had to be short, with full flow compensation and partial Fourier encoding. The fat had to be suppressed; therefore, water and fat should be out of phase. For 7T, this is possible at 0.5, 1.5, 2.5 ms, and so on. The minimal TE depends on the resolution and the gradient power, and in our case 2.5 ms is the optimum. Therefore, the TE1 was chosen at 2.5 ms.

Table 1: MR sequence parameters				
Sequence Parameter	1.5T	7T		
TR (ms)	25	20.0		
TE (ms)	35	2.5/15.0		
Flip angle (°)	10	16-24 (variation		
		over slab)		
Bandwidth (Hz/pixel)	100	203		
Matrix size	276×226	508×399		
Parallel imaging	Yes (acceleration	Yes (acceleration		
	factor = 1.8)	factor = 2.5)		
Flow compensation	No	Yes		
Section thickness (mm)	0.8	0.6		
Gap (mm)	None	None		
No. of sections	125	167		
Acquired voxel size (mm)	$0.8 \times 0.8 \times 0.8$	$0.35\times0.4\times0.6$		
Interpolated voxel size (mm)	$0.43 \times 0.43 \times 0.4$	$0.35 \times 0.35 \times 0.3$		
Acquisition time	6 min 56 s	8 min 50 s		

The TE2 was designed to make a venogram. Therefore, it was important to have a full echo and a flyback gradient, so the second readout gradient had the same polarity as the first readout gradient.²⁷ To obtain enough sensitivity, the TE had to be relatively long, preferably between 20 and 30 ms. A longer TE was not useful because the SNR would become too low. For the choice of the TE2, it is important that the TR is short enough to obtain good background suppression on the TE1 image (arteriogram). The flip angle was optimized for angiography in the TE1 and for venography in the TE2. Excitation pulses consisted of tilt-optimized nonsaturated excitation pulses with nominal flip angle variation of 16–24° in the feet-head direction. With this flip angle, the TR had to be approximately 20 ms.²⁸ Here, a balance had to be found between a long enough TE for the venogram and a short enough TR for good background suppression in the arteriogram. As a compromise, the TE2 was chosen at 15 ms.

The parameters of the T2*-weighted imaging at 1.5T were chosen to be comparable to the 7T scan within the possibilities of the 1.5T MR imaging. Therefore, a scan with a high spatial resolution was made, with an acquired voxel size of $0.8 \times 0.8 \times 0.8$ mm. A PRESTO technique with a TE of 35 ms and a TR of 25 ms was used. During this sequence, additional field gradients shift the refocused gradient echo into the subsequent TR period, which results in a TE longer than the TR. In this sequence, the flip angle is not optimized for maximum SNR with the given TR, but small flip angles are needed to attenuate the contribution of spin echoes and stimulated echoes.²⁹ With this technique, it is possible to obtain a longer TE with a relatively short TR, resulting in a shorter scan time. This makes it possible to obtain a high-resolution scan with a long TE within a reasonable scan time. The use of PRESTO has been shown to result in a higher contrast-tonoise ratio compared with 3D-GRE for the visualization of microbleeds.30

Postprocessing

The postprocessing of the data was performed on the console of the MR imaging system. The MIPs were reconstructed for transversal slabs (thickness, 3 mm; 2-mm overlap; 100 sections) for the 1.5T scan and for the TE1 and the TE2 of the 7T scan (Fig 1).

Rating of Microbleeds

All 1.5T and 7T scans were independently scored by 2 observers, both experienced neuroradiologists. The observers were trained in one session where they together evaluated scans of 3 patients with microbleeds. These 3 patients were excluded from the statistical analysis.

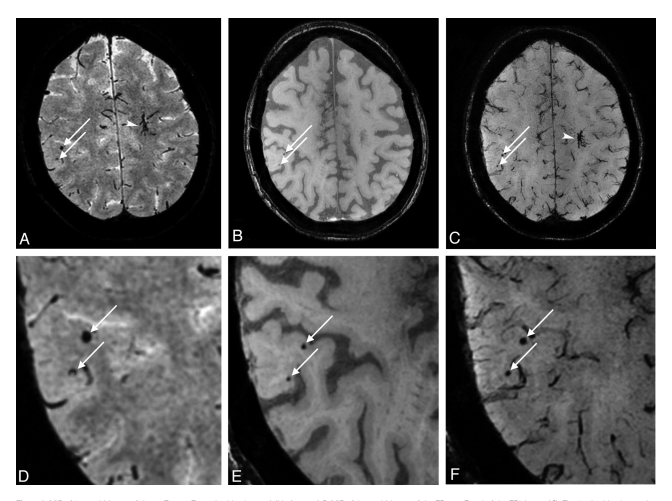


Fig 1. A, MIP of 3 mm thickness of the 1.5T scan. Two microbleeds are visible (arrows).B, MIP of 3 mm thickness of the TE1 at 7T and of the TE2 image (C). The 2 microbleeds are also visible on both the TE1 and TE2 images (arrows). Note the developmental venous anomaly visible on the 1.5T scan (arrowhead) and very clearly demarcated on the TE2 of the 7T scan. D-F, Magnifications of the microbleeds on the 1.5T scan, the TE1 of the 7T scan, and the TE2 of the 7T scan, respectively.

For the 7T scans, TE1 and TE2 images were both used to score the microbleeds. The observers were blinded to the scan on the other field strength and to all clinical information. The scans of 10 patients were scored twice by one of the observers with a gap of 1 week to assess intraobserver reliability. The scans that were scored differently by the 2 observers were evaluated in a consensus meeting to obtain a final score.

The 1.5T and 7T scans were evaluated in a random order. There was a minimum gap of 1 week between the evaluation of a 1.5T and 7T scan of the same patient. The rating of the microbleeds was based on the Microbleed Anatomical Rating Scale (MARS).³¹ In this rating scale, microbleeds are scored as "definite" or "possible" microbleeds. In our study, we focused on definite microbleeds that were scored when they were visible on the TE1 or on the TE2 or on both echo images at 7T. Definite microbleeds are defined as small, rounded or circular, well-defined hypointense lesions within brain parenchyma with clear margins ranging from 2 to 10 mm on GRE T2*-weighted images in the MARS. In this study, we slightly adjusted this definition by excluding the size criterion. Due to the blooming effect, microbleeds appear larger on GRE T2*-weighted scans than their actual size. The blooming effect depends among other things on the field strength and the TE and will therefore be different for the 1.5T and 7T scan, which will result in different sizes for the same microbleeds at the 2 field strengths. This makes the size criterion not appropriate for this comparison study. Symmetrical areas of calcification in the basal ganglia, choroid plexus, and pineal gland were excluded, as were signal intensity voids caused by sulcal vessels and low-signal intensity lesions thought to be signal intensity voids due to adjacent bone. ^{3,10,31} Afterward, dissimilarities between the 1.5T and the 7T scans were assessed in a side-by-side comparison.

Statistical Analysis

Inter- and intraobserver reliability for the presence and absence of microbleeds was tested with Cohen κ test. The κ -statistic was interpreted as poor (0–0.20), fair (0.21–0.40), moderate (0.41–0.60), good (0.61–0.80), or very good (0.8–1). The reliability of the number of microbleeds was quantified by the ICC. The difference in prevalence of microbleeds between 1.5T and 7T scans was tested with the McNemar test. For the analysis of the difference in number of visible microbleeds between 1.5T and 7T scans, the Wilcoxon signed rank test was used. This analysis was done for all patients and for a selection of patients whose scans showed microbleeds at both field strengths. The analyses were performed by using the statistical software package SPSS (Version 15.0 for Windows; SPSS, Chicago, Illinois).

Results

Baseline characteristics of the 34 patients are shown in Table 2.

Inter- and Intraobserver Reliability

Table 3 shows the number of patients with and without definite microbleeds scored by both raters on 1.5T, resulting in a

Table 2: Baseline characteristics of all patients			
	All Patients (n = 34)		
Age (yr) ^a	58 ± 12		
Male gender (%)	85		
Inclusion in SMART study with			
Vascular risk factors (%)	50		
Cerebrovascular disease (%)	26		
Peripheral artery disease (%)	15		
Cardiovascular disease (%)	9		

a Age is expressed as mean ± SD

Table 3: Number of patients with visible microbleeds at 1.5T for the 2 observers

	Observer 2		
1.5T	Microbleeds Absent	Microbleeds Present	Total
Observer 1			
Microbleeds absent	21	6	27
Microbleeds present	3	4	7
Total	24	10	34

Table 4: Number of patients with visible microbleeds at 7T for the 2 observers

	Observer 2		
7T	Microbleeds Absent	Microbleeds Present	Total
Observer 1			
Microbleeds absent	18	7	25
Microbleeds present	2	7	9
Total	20	14	34

Table 5: Number of patients with visible microbleeds at 1.5T and 7T after consensus between the 2 observers

	7T		
Consensus 2 observers	Microbleeds Absent	Microbleeds Present	Total
1.5T			
Microbleeds absent	17	9	26
Microbleeds present	0	8	8
Total	17	17	34

fair interobserver reliability ($\kappa=0.30$). At 7T, the interobserver reliability for definite microbleeds was moderate ($\kappa=0.42$) (Table 4). The intraobserver reliability was moderate at 1.5T ($\kappa=0.52$) and good at 7T ($\kappa=0.80$). For the number of microbleeds, moderate interobserver reliability was demonstrated for the 1.5T (ICC = 0.50) and good reliability for the 7T (ICC = 0.61) scans. The intraobserver reliability for the number of microbleeds was moderate at 1.5T (ICC = 0.59) and very good at 7T (ICC = 0.94).

Presence and Number of Microbleeds

Table 5 shows the number of patients with and without microbleeds at 1.5T and 7T after consensus. One or more microbleeds were found on 1.5T scans in 8 (23.5%) patients and on 7T scans in 17 (50%) patients. The number of patients with microbleeds at 7T was significantly higher than at 1.5T (P = .002). There were no patients in whom microbleeds were scored at 1.5T that were not scored at 7T.

In the 8 patients with microbleeds on the 1.5T scans, 15 microbleeds in total were scored (median number of microbleeds, 0.0; range, 0-6). Of these 15 microbleeds, 7 were located frontal, 3 parietal, 2 occipital, 1 temporal, 1 in the basal ganglia, and 1 in the deep and periventricular white matter. In the 17 patients with microbleeds on the 7T scans, 36 microbleeds in total were scored (median, 0.5; range, 0-5), a number that was significantly more than at 1.5T (P = .003). Of the 36 microbleeds scored on the 7T, 14 were located frontal, 7 parietal, 5 occipital, 5 temporal, 2 in the basal ganglia, 2 in the cerebellum, and 1 in the thalamus. In all 8 patients with microbleeds on 1.5T, microbleeds also were present on the 7T scan. The median number of microbleeds in these patients with visible microbleeds at both field strengths was 1.0 (range, 1-6) at 1.5T and 2.0 (range, 1-5) at 7T. However, this difference was not significant (P = .236).

Side-by-Side Comparison

Twenty-seven microbleeds (75%) out of the 36 microbleeds that were scored on 7T scans were not scored on 1.5T scans. Side-by-side comparison of 1.5T and 7T scans showed that 13 (48%) of the microbleeds that were scored at 7T and not on 1.5T were small microbleeds on the 7T scans, which were not visible on the 1.5T scan. Two of these microbleeds were not visible due to overlap of a venous structure. Five other microbleeds (19%) were visible at 1.5T, however they were located in the vicinity of a venous structure and therefore could not be reliably distinguished as a microbleed. Four microbleeds (15%) were visible as hypointense lesions, but not as a typical well-defined, round-shaped lesion, as illustrated in Fig 2. As shown in Fig 3, 3 microbleeds (11%) that were not scored on the 1.5T scans were in retrospect visible as very small lesions that could hardly be distinguished from the background noise. One microbleed (4%) was missed, which was located near the hypointense basal ganglia; on the 7T scan, this microbleed was only visible on the TE1 image. One other microbleed (4%) was just overlooked by both observers.

Of the total of 15 microbleeds that were scored on 1.5T scans, 5 microbleeds (33%) were not scored on 7T scans. In retrospect, these microbleeds were visible as very small lesions on the 7T scans and were missed by the observers, or they were defined as no definite microbleed during the consensus meeting. An example of one of those microbleeds is shown in Fig 4.

Discussion

In this study, we found a fair-to-moderate inter- and intraobserver reliability for the presence of microbleeds by using 3D T2*-weighted imaging at 1.5T and moderate-to-good reliability by using dual-echo T2*-weighted imaging at 7T. For the number of microbleeds, the reliability was better; it was moderate at 1.5T and good to very good at 7T. Overall, we found an increased reliability for both presence and number of microbleeds at 7T compared with 1.5T. Cerebral microbleeds were detected in more patients at 7T than at 1.5T. Furthermore, the number of microbleeds detected was higher at 7T.

Besides the detection of microbleeds in more patients and a higher number of microbleeds at 7T, the reliability of the detection also improves. Compared with 1.5T, both inter- and intraobserver reliability improved at 7T. This might be due to the higher resolution, increased SNR, and the use of the dual-

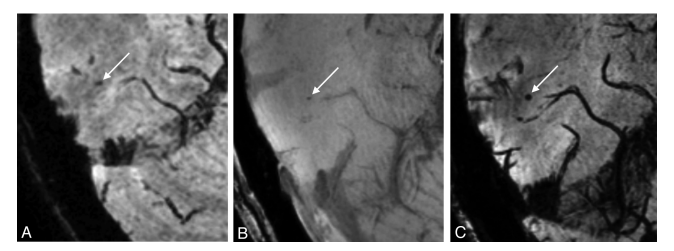


Fig 2. A, On this 1.5T scan, a hypointense lesion (arrow) is visible that is not a typical well-defined, round-shaped lesion as in the definition of a microbleed. This lesion was not scored as a microbleed on the 1.5T scan. In B and C, this same lesion is visible on the 7T scan. On the TE1 image (B), it is visible as a faint, round hypointensity and on the TE2 image (C), it is a clearly visible round hypointensity (arrow). On the 7T scan, both raters scored this lesion as a microbleed.

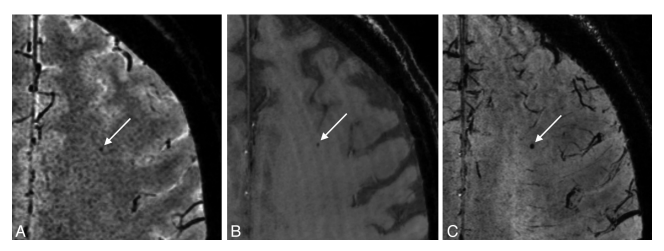


Fig 3. A, 1.5T scan shows a hypointense lesion (arrow) that can hardly be distinguished from noise and was not scored as a microbleed. On both the TE1 image (B) and TE2 image (C) of the 7T scan, this hypointense lesion is visible as a typical microbleed, showing enlargement at the TE2 due to the blooming effect. This lesion was scored as a microbleed by both observers.

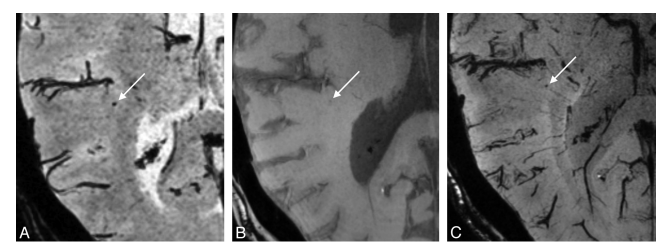


Fig 4. A, 1.5T scan shows a well-defined, round, hypointense lesion (arrow) that was scored as a microbleed. However, on the TE1 of the 7T scan (B), this lesion is only visible as a very faint hypointensity and on the TE2 image (C) it appears as a small microbleed. On the 7T scan, this lesion was not scored as a microbleed.

echo sequence at 7T. Recently, we showed that the TE1 image provides a good contrast of the dark microbleeds against a homogeneous, more hyperintense signal intensity of the brain

tissue and that 85% of all microbleeds are visible on the TE1 image.²⁵ If a microbleed is visible on the TE1 image and also shows a blooming effect on the TE2, it is more easily classified

as a definite microbleed, whereas the 1.5T scan provides only one single-echo image of the microbleed. In addition, the contrast between microbleeds and the background is less on the 1.5T scan and the resolution is slightly lower, making it more difficult to distinguish microbleeds from the background noise and from other hypointense structures compared with the 7T scan. The use of the TE1 image at 7T is helpful to distinguish microbleeds from deoxyhemoglobin in veins and ferritin in the basal ganglia, whereas microbleeds can be missed by overlap of veins or the hypointense appearance of ferritin-containing basal ganglia on the 1.5T scan or on the TE2 of the 7T scan.²⁵

Previous studies comparing the detection of microbleeds at 3T and 1.5T found significantly more microbleeds at 3T. In addition, the microbleed conspicuity improved at 3T. ^{18,23} These results are in line with our results of a better detection of microbleeds at 7T, indicating that the detection of microbleeds improves with increasing field strength. The higher resolution, higher SNR, and increased susceptibility effect at 7T increased the conspicuity of smaller microbleeds. This resulted in the detection of more and smaller microbleeds. However, a drawback is that very small microbleeds were hard to distinguish from background noise, and small microbleeds were easily missed by the observers.

Unexpectedly, some microbleeds were scored at 1.5T but not at 7T. A possible reason for this is that at 7T more and smaller veins are visible, making it harder to distinguish small microbleeds from small veins. This is illustrated by Fig 4, where the microbleed that was missed on the 7T scan was located near a very small vein and could therefore be mistaken as part of the vein.

Nowadays, SWI is increasingly used for the detection of microbleeds. ^{18,33} In SWI, the magnitude and phase images of the T2*-weighted scan are combined to create enhanced contrast between tissues with different susceptibilities. ³⁴ However, SWI depends strongly on the voxel aspect ratio, with an optimal voxel aspect ratio of 1:1:4; we used more isotropic voxels (isotropic voxels at 1.5T and an aspect ratio of 1:1.3:1.7 at 7T). ³⁵ The additional value of SWI will be minimal for scans with (near) isotropic voxels, and we therefore chose T2*-weighted imaging instead of SWI for the detection of microbleeds in our study.

A limitation of the study is the relatively low observer reliability at 1.5T. The study by Gregoire et al³¹ describing the MARS shows a very good observer reliability by using MARS at 1.5T. There are several possible explanations for this difference. First, Gregoire et al³¹ excluded patients with one definite microbleed from the analysis, which substantially improved their interobserver reliability; we did not exclude patients with only one microbleed. Besides that, in the study of Gregoire et al³¹ a section thickness of 5 mm with a gap of 1.5 mm was used, which is a large difference from our 0.8-mm sections. The use of thin sections results in many more sections and also more small microbleeds will become visible. Those small microbleeds are easily missed by the observers, especially when they have to evaluate many sections, inducing differences between the scores of observers. Furthermore, Gregoire et al³¹ did not score microbleeds <2 mm, whereas we did not use a minimum diameter in our study. In other studies at 1.5T, observer reliabilities vary from fair to very good;^{3,7,36} probably this is also due to differences in scan parameters and definition of microbleeds.

Another limitation of this study is that the scan parameters of the 1.5T T2*-weighted sequence were different from the 7T dual-echo sequence. This makes it difficult to evaluate the net effect of the increased field strength. However, when going to higher field strength, it is not feasible to maintain the same scan parameters. Therefore, we tried to optimize both scans as much as possible for the corresponding field strength within a reasonable scan time. The TE is the parameter that most affects sensitivity in T2*-weighted MR imaging. 10 The TE at 1.5T is 35 ms, whereas it is only 15 ms for the TE2 image at 7T. A longer TE gives more time for dephasing, resulting in an increase in diameter of microbleeds and also an increase in the number of microbleeds detected.¹⁹ However, within the dual-echo sequence at 7T, a balance had to be found between a TE, which gives enough dephasing and a short TR, which is necessary for good background suppression in the TE1 image. Therefore, the TE could not be equally long in the 7T sequence as in the 1.5T sequence. The use of a longer TE at 7T could result in the detection of more microbleeds. However, because other structures with a high susceptibility, such as veins, or iron deposition in the basal ganglia, will also become larger, this may lead to more overlap of microbleeds by these brain structures. Therefore, we think it is important to use 2 TEs to combine the benefits of a short and a long TE.

Although interest in cerebral microbleeds has greatly increased, their relevance in clinical practice remains unclear.³⁷ It is possible that the current studies, which are now mainly done at 1.5T, only detect the tip of the iceberg of all microbleeds present in the brain. Scanning at higher field strengths, such as 7T, might be necessary to improve the detection of microbleeds and the reliability of this detection. Because microbleeds can be detected in more patients and also the lesion load can be determined more accurately, this could result in more detailed and consistent information about the clinical significance of microbleeds. Furthermore, microbleeds are regarded as markers of vascular pathology and it can be interesting to study changes in patients with cerebral amyloid angiopathy, hypertensive vasculopathy, or Alzheimer disease. 10 The improved detection of microbleeds at 7T may be useful for follow-up studies with small changes in microbleed load over time. Furthermore, good detection methods for microbleeds are needed for future studies to determine whether microbleeds should affect clinical decision making, to identify their relation with other signs of cerebral small vessel disease, and to describe their independent contribution to cognitive and neurologic dysfunction.¹⁰

Conclusions

We found that presence and number of detected microbleeds are higher and also that the reliability of the detection of microbleeds improves with a 3D dual-echo T2*-weighted sequence at 7T compared with a 3D T2*-weighted sequence at 1.5T.

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