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Enhancement on Postcontrast 7T 3D-FLAIR
MRI in Multiple Sclerosis**

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Longitudinal Persistence of Meningeal Enhancement on Postcontrast 7T 3D-FLAIR MRI in Multiple Sclerosis

S.N. Jonas, I. Izbudak, A.A. Frazier, and D.M. Harrison



ABSTRACT

BACKGROUND AND PURPOSE: Preliminary research has demonstrated that postgadolinium 3D-FLAIR MR imaging at 7T may be a valuable tool for detecting abnormal meningeal enhancement and inflammation in MS; however, researchers have not systematically investigated its longitudinal persistence. We hypothesized that persistence of meningeal enhancement in MS varies on the basis of pattern of enhancement as well as demographic and clinical factors such as treatment status, disease phenotype, and disability score.

MATERIALS AND METHODS: Thirty-one subjects with MS were prospectively scanned before and after intravenous contrast administration at 2 time points, approximately 1 year apart. Fifteen subjects in the cohort were scanned at another time approximately 1 year later. Foci of enhancement were categorized into 4 subtypes: subarachnoid spread/fill, subarachnoid nodular, vessel wall, and dural foci. We reviewed follow-up scans to determine whether foci changed between time points and then compared persistence with demographic and clinical variables.

RESULTS: Persistence ranged from 71% to 100% at 1 year and 73% to 100% at 2 years, depending on the enhancement pattern. Subarachnoid spread/fill and subarachnoid nodular subtypes persisted less often than vessel wall and dural foci. Persistence was not significantly different between those on/off treatment and those with progressive/nonprogressive disease phenotypes. The number of persisting foci was significantly different in subjects with/without increasing Expanded Disability Status Scale scores (median, 12 versus 7.5, $P = .04$).

CONCLUSIONS: Longitudinal persistence of meningeal enhancement on 3D-FLAIR at 7T in MS varies by pattern of enhancement and correlates with worsening disability; however, it is not significantly different in those on/off treatment or in those with progressive/nonprogressive disease phenotypes.

ABBREVIATIONS: EDSS = Expanded Disability Status Scale; MP2RAGE = magnetization-prepared rapid acquisition of 2 gradient echoes

MS is a chronic inflammatory demyelinating disorder classically affecting white matter within the brain and spinal cord. In the past few decades, an additional pathophysiologic mechanism—meningeal inflammation—has been elucidated in MS, which is now believed to directly contribute to cortical demyeli-

nation, cortical neuroaxonal loss, microglial activation, and oligodendrocyte dysfunction.¹⁻⁶ Visualization of meningeal inflammation on MR imaging has become an active, and somewhat controversial, area of recent investigation.⁷ Landmark studies have demonstrated that gadolinium-enhanced 3D-FLAIR sequences, which have long been useful for identifying meningeal infection and carcinomatosis, can also be used to image meningeal disease in the MS population. At a magnetic field strength of 3T, Absinta et al⁷ found that approximately 25% of patients with MS demonstrated leptomeningeal enhancement on gadolinium-enhanced FLAIR.

Protocols for imaging meningeal enhancement were improved by Zivadinov et al,⁸ who showed the benefit of both pre- and postcontrast acquisitions and generating subtraction images when assessing meningeal enhancement, because these techniques decrease false-positives and reduce interpretation time. Recent preliminary research has also suggested that 7T MR imaging may be more sensitive than 3T for detecting meningeal enhancement. Although no direct 3T-versus-7T comparisons have

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Table 1: MRI sequence parameters

Sequence	Resolution (mm)	TR	TI	TE	Parallel Imaging	Flip Angle	Time (min:sec)
MP2RAGE	0.7 × 0.7 × 0.7	TR _{volume} = 8.25 s TR _{TFE} = 6.9 ms	TI ₁ = 1 s TI ₂ = 3.3 s	1.97 ms	SENSE = 2 × 2	FA ₁ = 7° FA ₂ = 5°	9:46
MPFLAIR	0.7 × 0.7 × 0.7	8000 ms	2077 ms	400 ms	SENSE = 2 × 3	90°	10:48

Note:—SENSE indicates sensitivity encoding; MPFLAIR, magnetization-prepared FLAIR; FA, flip angle; TFE, turbo field echo.

been made in the same study population, up to 90% of patients with MS undergoing contrast-enhanced brain MR imaging at 7T demonstrated at least 1 enhancing focus.⁹ This result closely approximates the 89% of patients with MS reported to show some element of leptomeningeal inflammation at postmortem examination.^{10,11} Given this radiologic-pathologic concordance, it is conceivable that 7T 3D-FLAIR may soon provide a noninvasive in vivo method of detecting and accurately quantifying the extent of meningeal inflammation in patients with MS.

Meningeal enhancement was noted to be a persistent phenomenon in prior 3T studies⁷; however, at 7T, where sensitivity for meningeal enhancement in MS appears to be significantly higher, it remains unknown whether smaller, more subtle foci of enhancement wax and wane in a predictable pattern across time or whether they remain longitudinally stable. Also unknown is the degree to which enhancement persistence with time is associated with previously described enhancement shape and morphology, including subarachnoid spread/fill and subarachnoid nodular patterns.^{7,9} Because prior studies have shown that the prevalence of meningeal enhancement varies with enhancement morphology,^{7,9} in this study, we hypothesized that persistence of meningeal enhancement in MS would vary on the basis of the morphology of enhancement as well as demographic and clinical factors such as treatment status, disease phenotype, and disability scores. Greater understanding of the imaging and clinical characteristics of meningeal enhancement is necessary if these features are to aid in the diagnosis of and prognosis for patients with MS.

MATERIALS AND METHODS

Standard Protocol Approval and Informed Consent

The institutional review boards at the authors' institutions approved this Health Insurance Portability and Accountability Act-compliant, prospective study. Written, informed consent was obtained from all participants.

Participants

Thirty-one volunteers, 26–61 years of age, with diagnoses of relapsing-remitting MS, secondary-progressive MS, or primary-progressive MS according to the 2010 revised McDonald Criteria were recruited.¹² Exclusion criteria included contraindications to contrast-enhanced MR imaging.

MR Imaging Protocol

Study participants were prospectively scanned at 2 time points approximately 1 year apart on a 7T Achieva scanner (Philips Healthcare, Best, the Netherlands) with a volume-transmit/32-channel head coil (Nova Medical, Wilmington, Massachusetts). Fifteen patients in the cohort were scanned at an additional third visit approximately 1 year later. Scans were obtained between September 9, 2014, and August 21, 2017. Dielectric padding was used for improved image homogeneity.¹³ Scanning parameters are listed in Table 1. Images

were acquired before the administration of contrast and again after the intravenous administration of 0.1 mmol/kg of gadoteridol (ProHance; Bracco Diagnostics, Princeton, New Jersey). Magnetization-prepared rapid acquisition of 2 gradient echoes (MP2RAGE) images were initiated approximately 3 minutes after contrast administration, and magnetization-prepared FLAIR images were initiated approximately 20 minutes after contrast administration.

Image Processing and Analysis

MP2RAGE images were processed to create a T1-weighted image and a T1 map.¹⁴ Images were then manipulated using Medical Image Processing, Analysis, and Visualization (Version 7.2; <http://mipav.cit.nih.gov>). Magnetization-prepared FLAIR images underwent N4 inhomogeneity correction before analysis.¹⁵ Pre- and postcontrast magnetization-prepared FLAIR images were registered to the precontrast T1 map. A magnetization-prepared FLAIR subtraction image was created by direct subtraction of the registered pre- and postcontrast images.

The magnetization-prepared FLAIR subtraction image, alongside the pre- and postcontrast magnetization-prepared FLAIR images, was reviewed by 2 independent raters (a postgraduate year 4 radiology resident and an academic MS neurologist) who were blinded to subject identity, disease state, and treatment regimen. Hyperintensities noted on the subtraction image were located on anatomic images and demarcated, if present, in the meningeal space on postcontrast images only. All foci were localized in 3 orthogonal planes before notation. When needed, coregistered MP2RAGE T1-weighted images were used for confirmation of anatomic locations. The pattern of enhancement was categorized on the basis of location and morphology and stratified into 1 of 4 subtypes. Subarachnoid spread/fill foci were characterized by the presence of contrast in the subarachnoid space distributed in an amorphous manner (Fig 1A). Subarachnoid nodular foci were characterized by small, round areas of contrast, usually 1–2 voxels (0.7–1.4 mm) and were adherent to the pial surface (Fig 1B). Vessel wall enhancement was characterized by contrast outlining the outer margin of a large meningeal vessel with a signal void in the lumen of the vessel, often resulting in a characteristic tram-track appearance (Fig 1C). Dural foci were characterized by discrete regions of enhancement clearly situated along the dural surface without extension into the subarachnoid space (Fig 1D). Following both independent reviews, a consensus review was performed under the supervision of an expert third rater (an academic neuroradiologist with 12 years of experience). After consensus review, follow-up images underwent linear registration (with 9 *df*) to baseline images. Consensus regions of contrast enhancement on baseline images were reviewed for their presence or absence on follow-up scans. The total number of foci per subject that persisted between scans was compared among different morphologies of meningeal enhancement and correlated

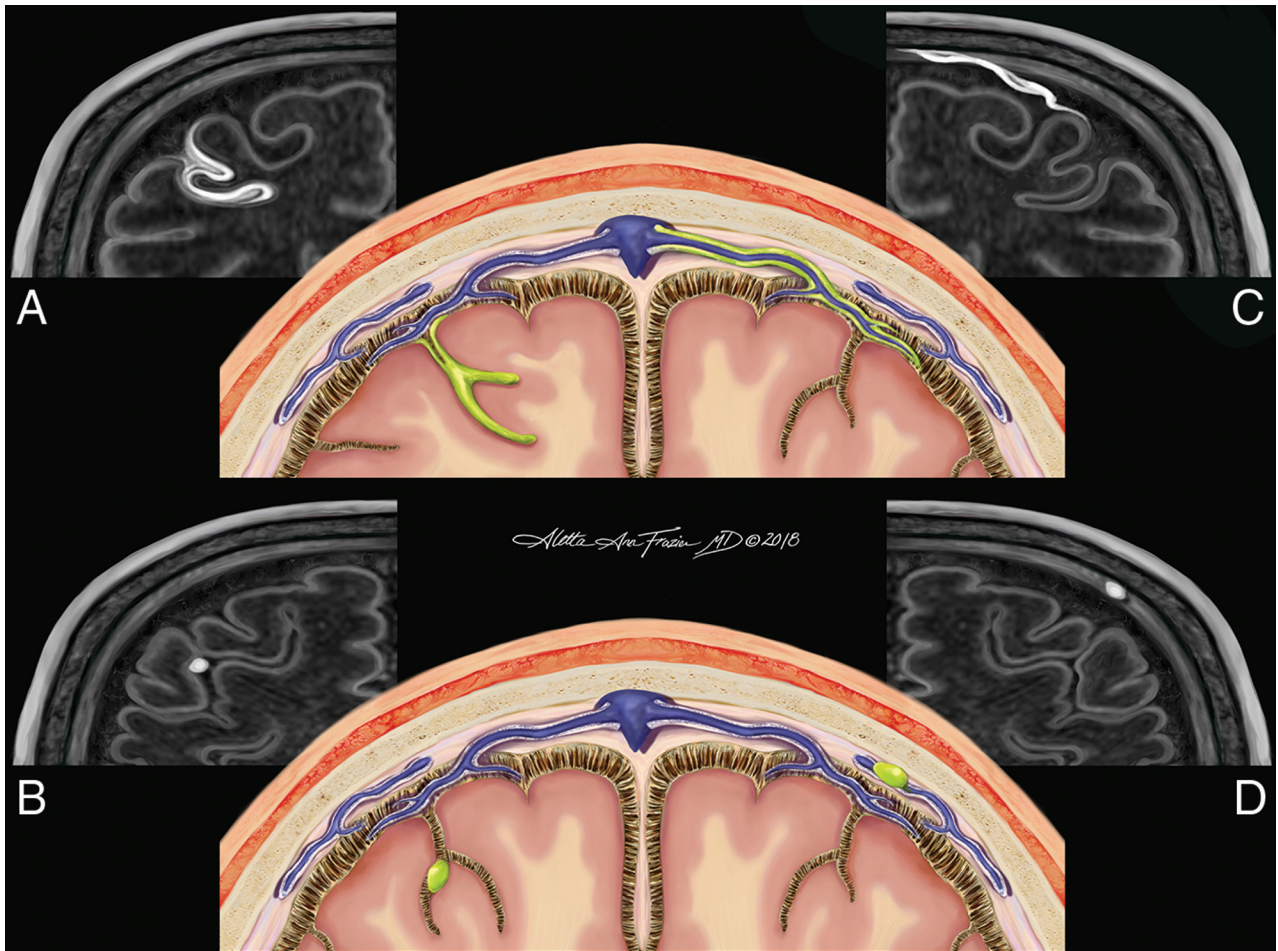


FIG 1. Original illustration depicting the 4 morphologies of meningeal enhancement seen in this analysis. Subarachnoid spread/fill pattern (represented by green in A) is an amorphous and ill-defined collection of contrast pooling within the cerebral sulci. The subarachnoid nodular pattern (B) is defined as a punctate, discrete site of meningeal enhancement located within the cerebral sulci abutting the pial surface. The vessel wall pattern (C) is characterized by extension of contrast along the outer margin of large meningeal vessels with a preserved internal flow void creating a characteristic tram-track appearance. The dural pattern (D) is a circumscribed, rounded focus of contrast situated along the dural margin without extension into the subarachnoid space. The perivascular, tubular white structures (seen in schematics A, B, and D) represent the recently discovered meningeal lymphatic system. Reaccumulation of leaked contrast from the CSF into these meningeal lymph channels is a potential mechanism for the venous rim pattern (C).

with demographic and clinical data. Additionally, the proportion of baseline foci per subject that persisted to follow-up scans was also compared with morphologic, demographic, and clinical factors.

Disability Measures

The Kurtzke Expanded Disability Status Scale (EDSS) was used to characterize disability.¹⁶ EDSS progression was defined as an increase of the EDSS score at follow-up of ≥ 1.0 if the baseline EDSS score was < 5.0 or an increase of ≥ 0.5 if the baseline EDSS score was ≥ 5.0 . The Modified Fatigue Impact Scale was used to assess MS-related fatigue.^{17,18} The Symbol Digit Modalities Test was used to assess cognitive functioning.¹⁹ These tests were administered at each study visit.

Statistical Analysis

Statistical analysis was performed in Stata 10.1 IC (StataCorp, College Station, Texas). Nonparametric testing was used due to the non-normal distribution of data. We performed group comparisons for

demographic and clinical variables using the Wilcoxon rank sum statistic. We computed the Spearman rank correlation for correlation testing. All statistical tests were performed with a significance threshold of $P < .05$. Due to the small sample size and exploratory nature of this study, adjustment for multiple comparisons was not performed.

RESULTS

We recruited 31 patients with MS; most had the relapsing-remitting MS phenotype ($n = 21$, 68%), though 7 subjects had secondary-progressive MS (23%) and 3 subjects had primary-progressive MS (10%) (Table 2). No subject had a comorbid neuroinflammatory disorder. Most subjects were on disease-modifying therapy ($n = 25$, 81%). This was a relatively stable and moderately disabled patient population with a median of 0 (range, 0–3) relapses in the year before enrollment and a median Expanded Disability Status Scale score of 3 (range, 1–6.5). Enhancing white matter lesions were seen in 3 subjects on review of

Table 2: Cohort baseline characteristics^a

Characteristics	
Age at enrollment (yr)	49 (26–61)
Sex	11/31 Men (35%), 20/31 women (65%)
Disease subtype at enrollment	21/31 (68%) RR, 7/31 (23%) SP, 3/31 (10%) PP
Disease duration at enrollment (mo)	109 (8–461)
Patients with new relapses in past 30 days	1/31 (3%)
No. of relapses in past year per subject	0 (0–3)
Modified Fatigue Impact Scale score at enrollment	43 (0–78)
Symbol Digit Modality Test at enrollment	50 (35–81)
Expanded Disability Status Scale score at enrollment	3 (1–6.5)
Immunomodulatory treatment status at baseline	
On treatment	25/31 (81%)
Not on treatment	6/31 (19%)
Treatment type at baseline	
Interferon	3/25 (12%)
Glatiramer	6/25 (24%)
Natalizumab	2/25 (8%)
Teriflunomide	1/25 (4%)
Fingolimod	4/25 (16%)
Dimethyl fumarate	9/25 (36%)
No. of subjects who switched between disease-modifying therapies from baseline to follow-up scans	10/31

Note:—RR indicates relapsing-remitting MS; SP, secondary-progressive MS; PP, primary-progressive MS.

^a Median values are shown with the range of observed values in parentheses.

Table 3: Anatomic distribution within the brain of enhancing meningeal foci at baseline

Brain Region	No. of Foci at Baseline	Percentage of Foci at Baseline
Right frontal	60	21.1
Left frontal	64	22.5
Right parietal	44	15.5
Left parietal	44	15.5
Right occipital	20	7.0
Left occipital	24	8.4
Right temporal	16	5.6
Left temporal	8	2.8
Right cerebellum	2	0.7
Left cerebellum	2	0.7

T1-weighted images, with 2 subjects having 1 enhancing lesion and 1 subject having 2 enhancing lesions.

At baseline, a total of 284 enhancing foci were identified across all 31 subjects. Table 3 lists the anatomic distribution of these foci within the brain. Most (>98%) of the observed foci were located supratentorially. Figure 2 shows the percentage of enhancing meningeal foci identified at baseline that persisted at later time points. Figures 3 and 4 provide examples of persisting and resolving enhancing meningeal foci from each group. Table 4 and Online Tables 1–3 compare the persistence of meningeal enhancement with demographic and clinical variables. We found no significant difference in the total number or proportion of longitudinally persistent enhancing meningeal foci between those on or off treatment or between those with progressive phenotypes (primary-progressive MS and secondary-progressive MS) versus a relapsing phenotype (relapsing-remitting MS). However, we did find significantly more ($P = .04$) persistent foci in Expanded Disability Status Scale progressors (median, 12; range, 1–15) compared with those who were not progressors (median, 7.5; range, 1–24). We also observed a nonsignificant trend toward a negative

association ($\rho = -0.31$, $P = .09$) between the proportion of persisting foci overall and the interval change in Symbol Digit Modalities Test scores at 1 year (On-line Table 3). Surprisingly and counterintuitively, we observed a positive correlation ($\rho = 0.45$, $P = .01$) between the proportion of enhancing meningeal foci that persisted at 1 year and baseline Symbol Digit Modalities Test scores (On-line Table 3). This association was driven by the correlation ($\rho = 0.48$, $P = .01$) between the proportion of subarachnoid spread/fill subtype that persisted at 1 year and baseline Symbol Digit Modalities Test scores. We also observed 15 foci of meningeal enhancement that developed in the interval between baseline and follow-up scans. The morphologies of these 15 foci were as follows: 6 subarachnoid spread/fill, 4 subarachnoid nodular, 2 vessel wall, and 3 dural foci.

DISCUSSION

In this study, we catalogued 2 enhancement patterns described in prior analyses (subarachnoid spread/fill and subarachnoid nodular)^{7,9} in addition to describing 2 new patterns of meningeal enhancement for the first time: vessel wall enhancement and dural foci. Previous studies without precontrast comparison sequences excluded from consideration regions of postcontrast hyperintensity in/near the dural sinuses, large subarachnoid veins, and the basal meninges to reduce false-positives because these structures often manifest precontrast T1 or FLAIR hyperintensity.^{7,9} Using techniques similar to those in the recent investigation by Zivadinov et al,⁸ we coregistered and subtracted pre- and postcontrast magnetization-prepared FLAIR sequences in all cases. Given this protocol, we did not have to exclude any structures a priori, and we were confident in our ability to differentiate true vessel wall and dural foci enhancement from intrinsically increased signal. Which anatomic/pathologic substrates are represented by vessel wall and dural foci is unknown, but most interesting, both closely match what was recently described for visualization of meningeal lymphatics by FLAIR MR imaging.^{20–22} Thus, these findings may represent gadolinium absorption by lymphatic structures after leakage into the CSF. The accumulation of gadolinium signal alongside the outer wall of vessels in the vessel wall pattern is also very reminiscent of the expected location and direction of drainage of solutes from brain parenchyma along the recently described glymphatic system.²³ Alternatively, it is also possible that vessel wall and dural foci could represent the reaccumulation, under hydrostatic pressure, of CSF-leaked gadolinium back into the venous system. They could also feasibly represent the actual sites of blood-brain and blood-CSF barrier disturbance secondary to ongoing inflammation.²⁴ Because age- and sex-matched healthy controls were not used in this study, the specificity of dural and vessel wall enhancements to MS is unknown. Future

work is needed to determine whether such findings are specific to MS, neuroinflammatory disease in general, or are seen in all patients.

We found no significant difference in the total number or proportion of persisting meningeal enhancement per subject between those on treatment and those off treatment. Lack of a significant difference between subjects on/off treatment may, in part, be explained by the low statistical power of our study because a relatively small number of untreated subjects were included.

However, the lack of difference is not surprising because prior studies have also failed to show differences in meningeal enhancement between those who are and are not taking disease-modifying medications.⁷ Our data reinforce the notion that current immunomodulatory medications may not adequately control meningeal inflammation. Of note, none of our subjects received a course of corticosteroids during the study. However, 10 of 31 subjects in this cohort switched between disease-modifying therapies from baseline to follow-up scans, including 2 subjects who changed to rituximab and 1 subject who switched to alemtuzumab—both monoclonal antibodies that impact B-cell function.

Despite such changes, most foci remained stable. Given the sample size of this report, we would not want to comment on the persistence (or lack thereof) of foci with individual therapy changes because conclusions from 1 or 2 examples would not be generalizable. Future comparative studies are needed to determine whether changes in any specific disease-modifying therapies or monoclonal regimens alter the longitudinal persistence of meningeal enhancement.

Surprisingly, we did not detect a significant difference in the persistence of meningeal enhancement between MS subjects with progressive phenotypes (primary-progressive MS and secondary-progressive MS) and those with re-

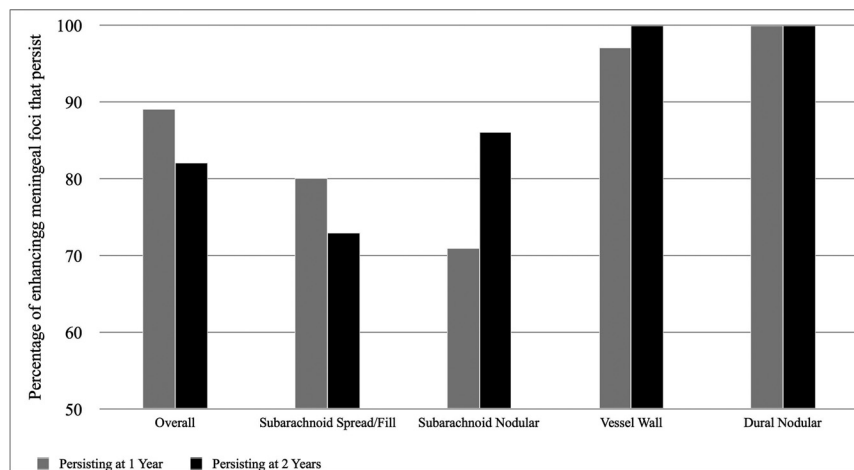


FIG 2. Graph displaying the percentages of baseline enhancing meningeal foci that persist 1 year later (gray bars) and 2 years later (black bars). All 31 participants were scanned at baseline and at 1 year, but 2-year data are limited to 15 participants. At 1 year, persistence was noted in 253/284 (89%) overall foci, 91/114 (80%) subarachnoid spread/fill foci, 10/14 (71%) subarachnoid nodular foci, 104/107 (97%) vessel wall foci, and 46/46 (100%) dural nodular foci. At 2 years, persistence was noted in 132/161 (82%) overall foci, 45/62 (73%) subarachnoid spread/fill foci, 6/7 (86%) subarachnoid nodular foci, 55/55 (100%) vessel wall foci, and 34/34 (100%) dural foci.

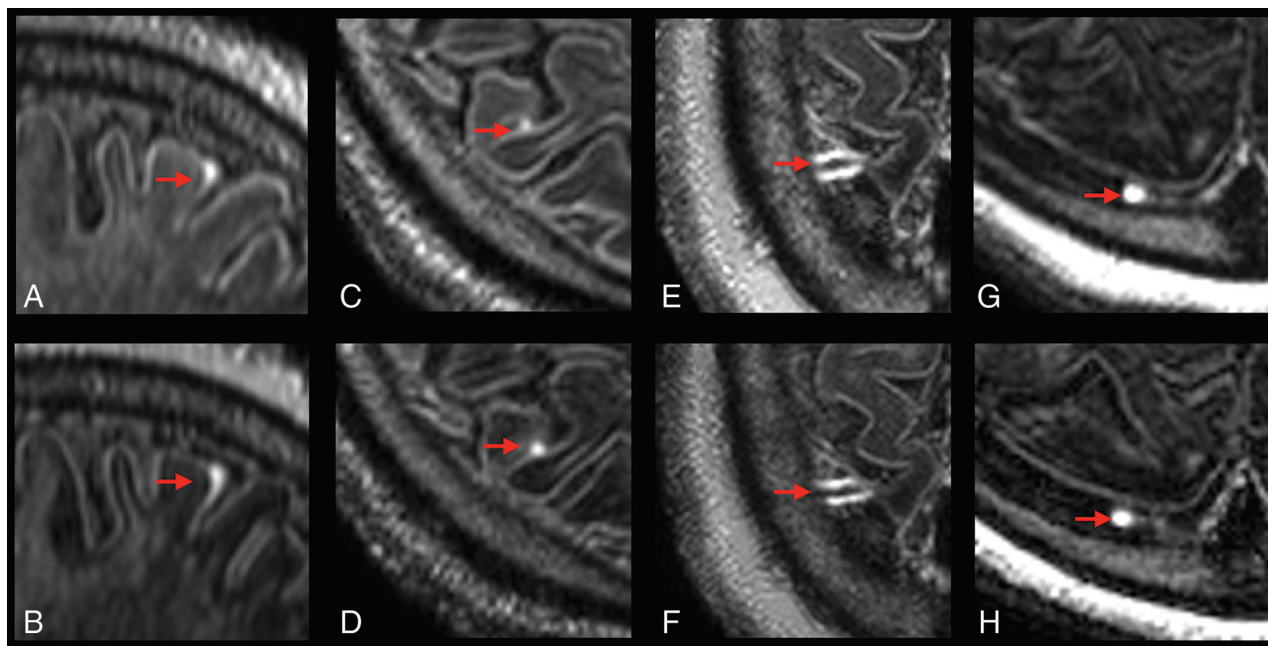


FIG 3. Examples of persisting foci of meningeal enhancement on delayed postcontrast FLAIR at 7T. Sagittal reformatted images show subarachnoid spread/fill enhancement that persists from December 15, 2015 (A), to March 3, 2017 (B), in a 58-year-old woman with relapsing-remitting MS. Axial images show subarachnoid nodular enhancement that persists from October 8, 2014 (C), to March 3, 2017 (D), in a 49-year-old man with relapsing-remitting MS. Axial images show vessel wall enhancement that persists from March 14, 2016 (E), to April 4, 2017 (F), in a 57-year-old man with primary-progressive MS. Axial images show dural enhancement that persists from May 9, 2016 (G), to May 31, 2017 (H), in a 44-year-old woman with secondary-progressive MS. Note that no intrinsic signal was observed in these locations on precontrast acquisitions (not shown).

lapsing-relapsing MS. This finding runs counter to the previously proposed theory that meningeal enhancement may be a substrate specific to progressive MS, with the associated cortical demyelination and volume loss representing a distinctly late marker of disease.^{1,7} Indeed, previous studies have shown that the presence of leptomeningeal enhancement at 3T was 1.7-fold higher in progressive MS compared with relapsing-relapsing MS, and postmortem findings of meningeal inflammation were more profound in those with secondary-progressive MS.^{1,7} However, our 7T data showed no difference in the frequency of longitudinal persistence of meningeal enhancement between patients with MS with progressive and relapsing phenotypes.

Although the persistence of enhancing foci was not related to clinical phenotype, we did detect a significant relationship between the persistence of foci (especially subarachnoid spread/fill

foci) at 1 year with disability progression (by the Expanded Disability Status Scale) during the same time period. If postcontrast meningeal enhancement on magnetization-prepared FLAIR is indeed representative of meningeal inflammation, this finding may indicate that persistent rather than transient meningeal inflammation is required to affect prognosis in patients with MS. This notion is supported by postmortem data showing that the development of structures that support ongoing inflammation, such as ectopic lymphoid follicular tissue, in the meninges of patients with MS is associated with earlier onset of disease, shorter diagnosis-death interval, and more severe cortical pathology.^{1,25} If this relationship can be confirmed in larger studies, perhaps the elimination of persistently enhancing meningeal foci can become a target outcome for patients with MS.

While we found a significant relationship between changes in Expanded Disability Status Scale scores and the persistence of

enhancement, we were unable to detect any similar association between imaging findings and MS-related fatigue or cognitive deficits (On-line Tables). This difference may be due to the small sample size of our study or the short follow-up duration. Still to be investigated is the rate at which foci of meningeal enhancement develop with time, whether the rate of development is associated with enhancement morphology, and whether the rate of development is associated with demographic and clinical parameters. This study focused on subjects with MS only; future work is needed to elucidate the rate of longitudinal persistence of meningeal enhancement in other neuroinflammatory diseases.²⁶ Finally, our study is limited by possible false discovery because we did not perform multiple-comparison correction, given the exploratory nature of this investigation. Therefore, our results will require replication before widespread acceptance of these conclusions. Despite these limitations, these preliminary results provide

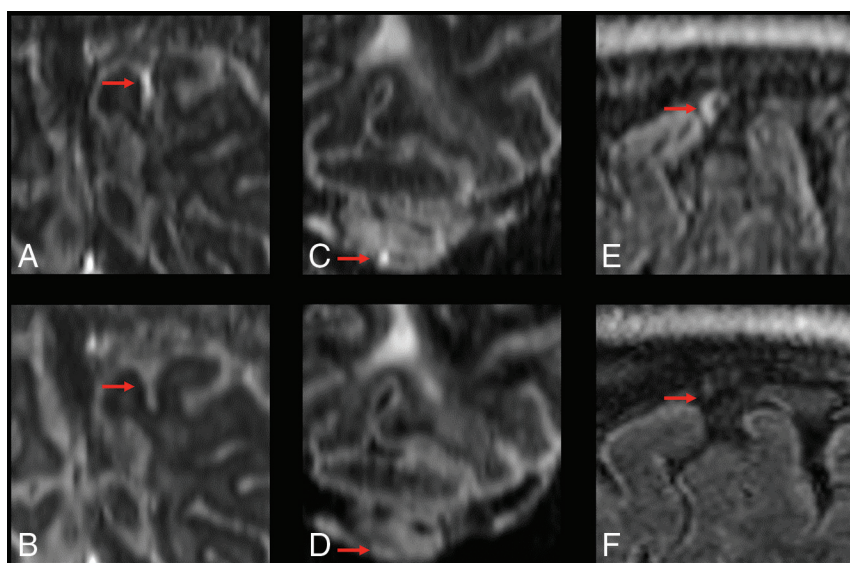


FIG 4. Examples of resolving foci of meningeal enhancement on delayed postcontrast FLAIR at 7T. Coronal reformatted images show subarachnoid spread/fill enhancement that resolves between October 23, 2014 (A), and February 26, 2016 (B), in a 49-year-old woman with relapsing-relapsing MS. Coronal reformatted images show subarachnoid nodular enhancement within the cerebellar folia that resolves between October 8, 2014 (C), and February 19, 2016 (D), in a 49-year-old man with relapsing-relapsing MS. Sagittal formatted images show vessel wall enhancement that resolves from May 9, 2016 (E), to May 31, 2017 (F), in a 44-year-old woman with secondary-progressive MS. Note that no foci of meningeal enhancement classified as a dural subtype resolved in this study.

Table 4: Wilcoxon rank sum test for longitudinal persistence of meningeal enhancement versus demographic and clinical factors^a

	On Treatment (n = 25)	Not on Treatment (n = 6)	Progressive MS (n = 10)	Nonprogressive MS (n = 21)	EDSS Progressor at 1 yr ^b (n = 7)	EDSS Nonprogressor at 1 yr ^b (n = 24)
Total No. of overall foci persisting at 1 yr per subject	9 (1–24)	9 (1–15)	8 (1–24)	9 (1–15)	12 (1–15)	7.5 (1–24) ^c
Total No. of subarachnoid spread/fill foci persisting at 1 yr per subject	2 (0–9)	2 (0–6)	2 (0–9)	2 (0–9)	5 (1–9)	2 (0–9)
Total No. of subarachnoid nodular foci persisting at 1 yr per subject	0 (0–2)	0 (0–1)	0 (0–0)	0 (0–2) ^c	0 (0–0)	0 (0–2)
Total No. of vessel wall foci persisting at 1 yr per subject	3 (0–11)	3.5 (1–6)	2 (0–6)	3 (0–11)	3 (0–11)	3 (0–7)
Total No. of dural foci persisting at 1 yr per subject	1 (0–9)	1 (0–6)	1 (0–9)	1 (0–6)	1 (0–6)	1 (0–9)

^a Median values are listed with the range of observed values in parentheses.

^b Criteria for EDSS progressor status is listed in the “Materials and Methods” section.

^c $P < .05$.

important new insight into the longitudinal activity of meningeal enhancement in MS.

CONCLUSIONS

Here we describe the results of a prospective, systematic investigation into the longitudinal persistence of meningeal enhancement in MS using 7T 3D-FLAIR. Given our pre- and postcontrast techniques, we are able to include, for the first time, vessel wall and dural foci subtypes, which persist most frequently; their appearance very closely matches recent descriptions of meningeal lymphatics or the glial lymphatics system.^{19–22} Longitudinal persistence of meningeal enhancement is not significantly different between those on or off immunomodulatory treatment, and there is not a significant difference in the rates of longitudinal persistence between those with progressive clinical phenotypes (primary-progressive MS and secondary-progressive MS) and those without a progressive clinical phenotype (relapsing-remitting MS). However, there is a significantly increased number of persistent foci in subjects who have worsening Expanded Disability Status Scale scores at 1 year compared with those who do not, suggesting that persistently enhancing meningeal foci may be an in vivo imaging marker for ongoing meningeal inflammation causing clinical progression.

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