Optimising 7T-fMRI for imaging regions of magnetic susceptibility

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Abstract The temporal signal-to-noise ratio (tSNR) of functional magnetic resonance imaging (fMRI) is particularly poor in ventral anterior temporal and orbitofrontal regions because of magnetic field inhomogeneity, a problem that is exacerbated at higher field strengths. In this 7T-fMRI study we compared three methods of improving sensitivity in these areas: parallel transmit, which uses multiple transmit elements, controlled independently, to homogenise the flip angle experienced by the tissue; multi-echo, which entails collection of multiple volumes at different echo times following a single radiofrequency pulse; and multiband, in which multiple slices are acquired simultaneously. We found that parallel transmit and multi-echo increased the magnitude of the BOLD signal change, but only multi-echo increased BOLD magnitude in areas prone to susceptibility artefacts. Multiband and denoising of multi-echo data with independent components analysis (ICA) both improved precision of GLM fit. Exploratory results suggested that multi-echo and ICA denoising can both benefit multivariate analyses. In conclusion, a multi-echo, multiband sequence improved fMRI quality in areas prone to susceptibility artefacts while maintaining sensitivity across the whole brain. We recommend this approach for studies investigating the functional roles of ventral temporal and orbitofrontal regions with 7T fMRI. Keywords: 7T-fMRI, parallel transmit, multi-echo, multiband, temporal lobe, orbitofrontal cortex

1. Introduction

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The temporal signal-to-noise ratio (tSNR) of functional magnetic resonance imaging (fMRI) varies across the brain. The ventral anterior temporal cortex and orbitofrontal cortex, for example, are located next to air-filled sinuses and so are affected by magnetic field inhomogeneity that causes signal dropout and geometric distortions (Devlin et al., 2000; Halai et al., 2014, 2015, 2024). This makes it challenging to use fMRI to investigate the roles that these regions may play in a myriad of cognitive processes – including vision (Devereux et al., 2018), language (Borghesani et al., 2016), multimodal semantic cognition (Lambon Ralph et al., 2017), emotion (Fernandez et al., 2017), social cognition (Binney et al., 2016; Zahn et al., 2007), theory of mind (DuPre et al., 2016), and executive function (Duncan, 2010). Signal dropout and geometric distortions are more severe at higher field strengths, implying that it may be especially difficult to measure task-related activity in susceptible regions with ultra-high-field fMRI (e.g., 7T-fMRI). However, 7T-fMRI also has many advantages in regions unaffected by magnetic susceptibility artefacts: 7T-fMRI offers improved tSNR relative to 3T-fMRI (Morris et al., 2019), which can be used to reduce voxel size and enable applications such as laminar fMRI (Koopmans et al., 2011) or to reduce acquisition times and enable shorter scan times for special populations such as patients with neurodegenerative diseases (Cope et al., 2023). 7T-fMRI also benefits from improved spatial specificity relative to 3T-fMRI because the signal from cortical microvasculature is enhanced while the signal from large veins is reduced (Marques & Norris, 2018). Improving signal homogeneity would allow researchers studying the whole brain (or focusing on regions prone to susceptibility artefacts) to take full advantage of 7T-fMRI; therefore, in this study we compared three methods of doing so. One possible method for counteracting signal dropout is parallel transmit (pTx), which uses multiple transmit elements, controlled independently, to homogenise the flip angle pattern experienced by the tissue (Deniz et al., 2019). A recent study used pTx to counteract signal dropout in ventral anterior temporal regions for echo-planar imaging (EPI) 7T fMRI (Ding et al., 2022). pTx improved tSNR across the brain compared to a standard sequence, particularly in the temporal lobes. However, there was no improvement in functional contrast during a semantic association task that is known to recruit the anterior temporal lobes in 3T-fMRI studies (Jung et al., 2017).

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A second method of recovering signal in these regions is multi-echo (ME) imaging (Kundu et al., 2017; Poser et al., 2006; Posse, 2012). T₂* is known to vary across the brain (Hagberg et al., 2002); in areas prone to magnetic susceptibility artefacts, T₂* is particularly short due to increased intravoxel dephasing. A single echo provides sensitivity to a narrow range of T₂* values; the echo time (TE) is therefore selected to provide the best compromise of sensitivity to T₂* across the whole brain. Combining data from multiple echoes increases the range of T_2^* that can be imaged with high fidelity. ME has been shown to improve functional contrast (Poser & Norris, 2009) and spatial specificity (Boyacioğlu et al., 2015) at 7T and 3T (Fernandez et al., 2017; Halai et al., 2024; Kirilina et al., 2016; Lynch et al., 2020). Having multiple echoes also facilitates the separation of signal and noise because signals decay in a well-characterised way across echoes, whereas noise does not. This principle underpins multi-echo independent components analysis (ME-ICA), via which ICA components that are TE-independent, and thus are likely to be noise rather than bloodoxygen-level-dependent (BOLD) signal, can be removed (Dipasquale et al., 2017; Kundu et al., 2011, 2013, 2015, 2017). This method may enhance signal detection in areas prone to susceptibility artefacts on top of the advantage offered by ME alone (e.g. Lombardo et al., 2016). ME sequences have some potential disadvantages. For example, multi-echo can lengthen repetition time (TR). In-plane acceleration is frequently needed to achieve a sufficiently short first TE, which reduces tSNR (Yun & Shah, 2017). In turn, a short first TE, combined with hardware constraints, often limits the minimum voxel size (Koopmans et al., 2011). Critically, in many previous studies examining the benefits of ME sequences compared to single-echo (SE), the "single-echo" data were extracted from the ME dataset (Amemiya et al., 2019; Bhavsar et al., 2014; Caballero-Gaudes et al., 2019; Cohen et al., 2017, 2017, 2018; Dipasquale et al., 2017; Evans et al., 2015; Fernandez et al., 2017; Gilmore et al., 2022; Kovářová et al., 2022). This means that the "single-echo" data will inherit suboptimal parameters that are ME-specific, making the comparison unfair. A third strategy for improving acquisition is multiband (MB) imaging, also known as simultaneous multi-slice, in which multiple slices are acquired simultaneously (Barth et al., 2016; Moeller et al., 2010; Setsompop et al., 2012). Typically MB has been applied to increase temporal resolution – multiple studies have shown that MB can reduce noise aliasing, increase statistical power and counteract the increase in TR associated with ME (Feinberg et al., 2010; Griffanti et al., 2014; Halai et al., 2024; Puckett et al., 2018; Smith et

al., 2013). Benefits specific to task fMRI have been less clear (Demetriou et al., 2018; Todd et al., 2016) since some noise features are unlikely to be correlated with task regressors and can be removed with high-pass filtering. Other possible disadvantages of MB include a reduction in tSNR due to increases in g-factor effects (Demetriou et al., 2018; Risk et al., 2021; Setsompop et al., 2012) and leakage of signal into the simultaneously-excited slices (Todd et al., 2016).

There is therefore a need to evaluate 7T-fMRI sequences to determine which parameters are important for counteracting signal dropout and improving sensitivity in regions prone to susceptibility artefacts without compromising signal quality elsewhere. We tested five possible sequences. These consisted of pTx, plus a 2 x 2 factorial design varying number of echoes and multiband factor: single-echo single band (SESB), single-echo multiband (SEMB), multi-echo single band (MESB), and multi-echo multiband (MEMB). We used a semantic judgment task that is known to evoke activity across the semantic network, including areas severely affected and those relatively unaffected by susceptibility artefacts (Jung et al., 2017).

We had two univariate effects of interest – activation magnitude and activation precision (Halai et al., 2024). Activation magnitude is the magnitude of the BOLD signal change, operationalised as the 1st-level beta values extracted from each voxel. We hypothesised that both pTx and ME would recover signal and hence increase activation magnitude relative to the SESB sequence (which we used as a baseline). Activation precision is the reliability of the BOLD signal change, analogous to the functional contrast-to-noise ratio (fCNR) and operationalised as the 1st-level t-values extracted from each voxel. We hypothesised that MB sequences, with greater effective degrees of freedom, would increase activation precision relative to single band (SB) sequences. We also had two secondary hypotheses: first, since ME-ICA denoising removes TE-independent noise, greater activation precision would be observed in multi-echo denoised data (MEdn) relative to ME data without denoising; second, that any MB advantage would be due to the increase in the number of volumes.

Our study focused primarily on univariate effects. However, multivariate analysis techniques, which exploit variance and covariance between voxels and can accommodate participant-specific differences in activation patterns (Coutanche, 2013; Davis et al., 2014; Davis & Poldrack, 2013) are rapidly gaining popularity. These methods frequently rely on the

assumption of good-quality signal across the whole brain (Frisby et al., 2023) and so we conducted an exploratory analysis, following the method of Haxby et al. (2001), to decode task condition from the data acquired with each sequence. Finally, we tested for the presence of slice leakage artefacts in our MB data.

To summarise, our study aimed to compare pTx, ME and MB as methods for improving sensitivity in ventral temporal and orbitofrontal regions while maintaining image quality across the rest of the brain.

2. Methods

2.1. Participants

20 healthy native speakers of British English (age range 18-50, mean age 33.45 years, 12 female, 8 male) participated in the study. All were right-handed, had normal or corrected-to-normal vision, and had no neurological or sensory disorders. All participants gave written informed consent. The research was approved by a local National Health Service (NHS) ethics committee (04/Q105/66).

2.2. Stimuli and task

All participants performed a semantic association task and a visual pattern matching task (hereafter called the "control task") adapted from a previous study (Jung et al., 2017). Each stimulus consisted of three pictures presented simultaneously (Figure 1). Some pictures were line drawings taken from the Pyramids and Palm Trees Test (Howard & Patterson, 1992) and some were colour cartoons or photographs taken from the Camel and Cactus Test (Bozeat et al., 2000). In the semantic task, participants were instructed to indicate which of the two pictures at the bottom of the screen had the closest semantic relationship to the picture at the top (hereafter the "probe picture"). In the control task, participants were instructed to indicate which of two scrambled pictures (generated from the pictures used in the semantic task) at the bottom of the screen was identical to a scrambled probe. There were 248 unique picture triplets, so some stimuli were repeated between runs, but no stimulus was repeated within a run. E-Prime software (Psychology Software Tools Inc., Pittsburgh, USA) was used to display stimuli and record responses. Stimuli were rearprojected onto a screen at the back of the MRI scanner, and observers viewed stimuli through a mirror mounted to the head coil directly above the eyes.

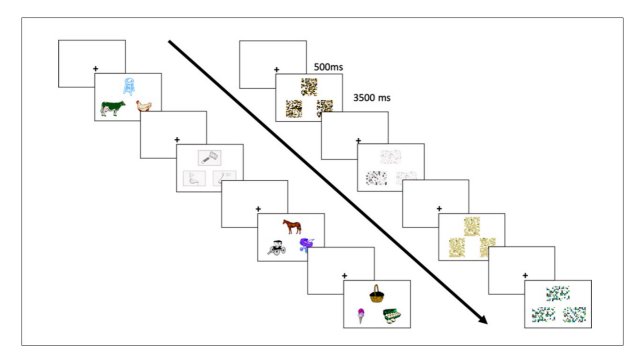


Figure 1: One block of the semantic association task (left of the arrow) and one block of the control task (right of the arrow). Each trial consisted of a fixation cross (500 ms) followed by three pictures presented simultaneously. Participants had to select which of the two pictures at the bottom was more semantically similar to (in the semantic task) or visually identical to (in the control task) the picture at the top (the probe picture). Each block lasted 16 s in total. Reprinted with permission from (Halai et al., 2024).

The study had a block design with three types of block: semantic, control and rest. Each task block consisted of four trials. Each trial consisted of a fixation cross presented for 500 ms followed by a stimulus presented for 3500 ms. Each rest block consisted of a fixation cross presented for 16 s. Each run began 16 s after the start of the MR sequence and then consisted of 30 blocks presented in the order: semantic, control, semantic, control, rest. There were five runs per participant, collected in a single session. Accuracy and reaction time were measured for each trial. Since reaction time is not normally distributed, both metrics were compared across tasks using Wilcoxon's signed-rank tests and across sequences using Friedman's nonparametric ANOVA.

2.3. Image acquisition

All images were acquired on a whole-body MAGNETOM Terra 7T MRI (Siemens Healthcare, Germany). An 8Tx32Rx head coil (Nova Medical, USA) was used to run all structural and functional imaging sequences.

An MP2RAGE anatomical scan was acquired with the following parameters: 224 sagittal slices (interleaved acquisition), FOV 240 x 225.12 x 240 mm³, voxel size 0.75 mm isotropic, TR 4300 ms, inversion times 840 and 2370 ms, TE 1.99 ms, nominal flip angles 5° and 6°, GRAPPA acceleration factor 3, and duration 8 minutes 50 seconds.

Next, manual B_0 -shimming was performed over the volume to be used for EPI. The aim was to reduce the water linewidth, defined as full width of the spectrum at half height, to below 40 Hz. However, for some participants the adjustments proved time-consuming and so adjustment time was capped at 30 minutes from the start of scanning after which the best shim parameters were adopted. The actual average water linewidth was 42 Hz (standard deviation = 9 Hz; missing data for 4 participants). Then a dummy pTx-EPI scan of one volume was acquired to trigger the acquisition of subject-specific B_0 and per-channel B_1 ⁺ field maps. The brain was divided into 5 slabs along the slice direction and slab-specific 2-spoke pTx excitation pulses were designed offline. Variable-rate selective excitation (VERSE; Hargreaves et al., 2004) was applied to reduce specific absorption rate (SAR) for the pTx sequence only.

There were five functional runs of EPI, one run of each sequence. Key parameters are given in Table 1. The order of sequences was counterbalanced across participants. The following parameters were held constant across sequences: 48 axial slices (interleaved acquisition), FOV 210 x 210 x 210 mm³ to cover the whole brain in most participants (visual inspection ensured that the ventral anterior temporal lobe was included in the FOV for all participants and the FOV was tilted up at the nose to avoid ghosting of the eyes into the temporal lobe), voxel size 2.5 mm isotropic (no gap), and A-P phase encoding direction. After each run, 5 further volumes were acquired with the phase encoding direction changed to P-A to facilitate distortion correction during preprocessing.

	SESB	SEMB	MESB	MEMB	рТх
No. echoes	1	1	3	3	1
Multiband factor	1	2	1	2	1
TR (ms)	3020	1510	3020	1510	3000
TE ₁ (ms)	25.00	25.00	11.80	11.80	25.00
TE_2 (ms)	-	-	27.05	27.05	-
TE ₃ (ms)	-	-	42.30	42.30	-
iPAT type	Off	Off	GRAPPA	GRAPPA	GRAPPA
iPAT factor	Off	Off	3	3	2
Phase partial Fourier	7/8	7/8	7/8	7/8	Off

Nominal flip angle (°)	50	50	78	63	40
Bandwidth (Hz/Px)	2204	2204	2204	2204	1804
Number of volumes acquired	171	340	171	340	172
Number of pulses	1	1	1	1	5 (VERSE 2- spoke pTx pulses)

Table 1: parameters of each sequence.

2.4. Data analysis

All analysis code is available at https://github.com/slfrisby/7TOptimisation/.

2.4.1. Preprocessing

For ease of data sharing we converted all DICOMs to BIDS format (Gorgolewski et al., 2016) using *heudiconv* v1.0.0 (Halchenko et al., 2024).

Standard reproducible preprocessing pipelines designed for 3T-fMRI, such as *fMRIprep* (Esteban et al., 2019; Gorgolewski et al., 2011; Markiewicz et al., 2024) perform poorly on 7T-fMRI EPI data. Therefore, the analysis pipeline was split into stages using different software packages.

Since it is notoriously difficult to perform a good-quality brain extraction on MP2RAGE data (because of salt-and-pepper noise in the background and cavities), two pipelines were used. The MP2RAGE T1w (combined) image first had its background noise removed using O'Brien regularisation (O'Brien et al., 2014) and was then submitted to the CAT12 pipeline for segmentation (Gaser et al., 2023; in SPM12; https://www.fil.ion.ucl.ac.uk/spm/). The bias- and global-intensity corrected T1w image produced was provided to as input to the anatomy pipeline in fMRIPrep 21.0.1. TheT1w image was skull-stripped with a Nipype implementation of the antsBrainExtraction.sh workflow (ANTs 2.3.3; Avants et al., 2009, 2011; https://github.com/ANTsX/ANTs/). Volume-based spatial normalisation to standard space (MNI152NLin2009cAsym) was performed through nonlinear registration with antsRegistration.sh (ANTs).

Functional preprocessing was performed using in-house code, composed of functions from AFNI (v.18.3.03; Cox, 1996; Cox & Hyde, 1997; https://afni.nimh.nih.gov/), FSL (v.5.0; Andersson et al., 2003; Jenkinson et al., 2012; Smith, 2002; Smith et al., 2004;

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https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/), tedana (v.23.0.1; DuPre et al., 2021; Kundu et al., 2011, 2013; The tedana Community et al., 2023; https://tedana.readthedocs.io/en/stable/index.html) and ANTS (v. 2.2.0; Avants et al., 2009, 2011; https://github.com/ANTsX/ANTs/). EPIs were despiked using 3dDespike (AFNI), slice timing was corrected to the middle slice using 3dTshift (AFNI), motion was corrected with 3dvolreg and 3dAllineate (AFNI) using the first volume of each run as a reference (for ME datasets, the TE₁ image was aligned and the resulting transform was applied to the TE₂ and TE₃ images), and skull-stripped using BET (FSL) to create a participant-specific brain mask. For ME datasets only, tedana was used to create two timeseries – one in which the echoes were optimally combined based on T₂* weighting (Posse, 2012), and one in which the T₂* optimally-combined data were denoised using ICA. tedana conducts denoising by decomposing data using PCA and ICA, classifying components according to whether the signal scales linearly with TE (as the BOLD signal does), and reconstructing the data using only BOLD-like components. The brain mask created with BET was used as the mask for this stage. Next, for all datasets, unwarping was conducted using topup and applytopup (FSL). Field displacement maps were calculated using ten volumes (five with A-P phase encoding direction, extracted from the start of each functional run, and five with P-A phase encoding direction, collected separately after each run) and the resulting correction was applied to all images. Finally, the mean EPI for each run was coregistered to the skull-stripped native structural image using a rigid-body registration with AntsRegistrationSyN.sh (ANTS). EPIs were then transformed into standard space (MNI152NLin2009cAsym) by combining the transforms from native EPI to native T1 and the transforms from native T1 to standard space and applying those transforms to the EPIs using antsApplyTransforms (ANTS). Images were smoothed with a 6 mm FWHM Gaussian filter in SPM12 (https://www.fil.ion.ucl.ac.uk/spm/) for GLM analysis. A separate functional preprocessing pipeline was used to create images for slice leakage analysis. This pipeline differed from the main pipeline in the following ways. Despiking was omitted to avoid the removal of noise. For ME datasets only, rather than run the full tedana workflow, we conducted optimal combination of data from multiple echoes (but no denoising) using the t2smap command using all voxels in the volume (i.e. no brain

mask). All coregistration steps were omitted (the images remained in native EPI space) and no smoothing was applied.

2.4.2. 1st-level (within-participant) GLM

Data were analysed using the general linear model (GLM) approach implemented in SPM12 in MATLAB r2019a. We had 5 timeseries of primary interest – standard single-echo single band (SESB), parallel transmit (pTx), single-echo multiband (SEMB), multi-echo single band (MESB) and multi-echo multiband (MEMB). For the latter two sequences, data from all echoes were optimally combined but were not ICA-denoised. We also generated two timeseries with ME-ICA denoising - multi-echo single band with ICA denoising (MESBdn) and multi-echo multiband with ICA denoising (MEMBdn) - and two downsampled MB timeseries created by extracting odd-numbered volumes to match the number of volumes in the single band timeseries - odd-numbered volumes of single-echo multiband (SEMBodd) and odd-volumes of multi-echo multiband (MEMBodd).

At the individual subject level, each block of the semantic and control task was modelled as a boxcar function (resting blocks were modelled implicitly) and these boxcar functions were subsequently convolved with SPM's difference of gammas haemodynamic response function. The six motion parameters extracted during preprocessing were used as regressors of no interest. The micro-time resolution was set as the number of slices (n = 48), the micro-time onset was set as the reference slice for slice-time correction (n = 24), and the high-pass filter cutoff was 128 seconds. The same MNI template used during preprocessing was used as a mask for the analysis. The parameter estimation method was restricted maximum likelihood estimation (ReML) and serial correlations were accounted for using an autoregressive AR(1) model during estimation. For univariate analyses the contrast of interest was greater activation for the semantic task than the control task (S > C). For exploratory multivariate pattern analysis (MVPA) each block (12 semantic and 12 control) was modelled individually in order to obtain one beta image per block.

Finally, for the slice leakage analysis, the modelling was rerun on the minimally-preprocessed timeseries without any brain mask. We obtained both univariate contrasts of interest (S > C) and beta images per block (for MVPA).

2.4.3. 2nd-level (across-participant) GLM

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2.4.3.1. Region-of-interest (ROI) analysis Regions of interest (ROIs) were defined based on a large-scale distortion-corrected 3T-fMRI study of the semantic network (Humphreys et al., 2015). For each comparison, ROIs were analysed only if they overlapped by at least one voxel with the whole-brain contrast of interest (S > C) summed over all sequences in the comparison. 2.4.3.1.1. Univariate analysis Activation magnitude (1st-level beta values) and activation precision (1st-level tvalues) were extracted from each ROI using a publicly-available script, roi extract.m (https://github.com/MRC-CBU/riksneurotools/blob/master/Util/roi extract.m). There were two planned t-tests for activation magnitude (pTx > SESB, ME > SE) and four planned t-tests for activation precision (MB > SB, MEdn > ME, MBodd > SB, SB > MBodd). For each planned t-test, results were Bonferroni-corrected for the number of ROIs included. 2.4.3.1.2. Exploratory multivariate pattern analysis (MVPA) The input to this analysis was the activation magnitude values extracted from each block individually (12 semantic and 12 control). For each block and each ROI, a vector of beta values within that ROI was created. The cosine dissimilarity between every possible pair of blocks was calculated. MVPA performance was operationalised as the mean between-task dissimilarity minus the mean within-task dissimilarity and paired t-tests were used to compare the metric across sequences (all planned t-tests described in the univariate ROI analysis were conducted; Haxby et al., 2001). 2.4.3.2. Whole-brain analysis All the above contrasts were assessed at the whole-brain level using t-tests (for comparing pTx and SESB) or using random-effects ANOVAs with one-sample t-tests on the summary statistic (for the three factorial designs: varying echo and band (SESB, SEMB, MESB and MEMB); varying denoising and band (MESBdn, MEMBdn, MESB and MEMB); and varying echo and downsampled band (SEMBodd, MEMBodd, SEMB and MEMB)). The ANOVAs were conducted using a publicly-available script, batch spm anova.m (https://github.com/MRC-CBU/riksneurotools/blob/master/SPM/batch_spm_anova.m). The group t-maps were assessed for significance by using a voxel-height threshold of p < 0.001

to define clusters and then a cluster-defining family-wise-error corrected threshold of p < 0.05 for statistical inference.

2.4.3.3. Slice leakage analysis

The group-level, whole-brain contrast of interest (S > C) in standard space was inspected and the coordinates of the peak t-values of the top 5 clusters were identified. These coordinates were back-projected to obtain 5 sets of corresponding coordinates in each participant's native EPI space (hereafter "seeds", labelled A). Next, voxels to which signal might be warped were identified as artefact locations based on phase shift (FOV/2, labelled B) and, in the MEMB data, GRAPPA (labelled Ag) and phase shift plus GRAPPA (labelled Bg). A spherical ROI, 4 voxels in radius, was defined around each seed location and possible artefact location using a modified version of the scripts developed for McNabb et al. (2020; https://github.com/DrMichaelLindner/MAP4SL/; our version available at https://github.com/slfrisby/7TOptimisation/).

Both univariate (McNabb et al., 2020) and multivariate (Halai et al., 2024) slice leakage tests were conducted. Activation magnitude was extracted from minimally-preprocessed data, and, for the multivariate analysis, the difference between mean within-task similarity and mean between-task similarity was calculated as in the ROI analysis. Paired t-tests were conducted for each seed and artefact location between each sequence of interest (SEMB and MEMB) and a control sequence. For artefact locations based on phase shift (B), the control sequence was the corresponding SB sequence (SESB for SEMB and MESB for MEMB). For artefact locations based on GRAPPA, the control sequence was the corresponding SE sequence, because SE sequences were collected without GRAPPA (SEMB for MEMB). Results were Bonferroni-corrected for the number of peaks (n = 5) and the number of possible artefact regions (n = 1 for SEMB and n = 3 for MEMB).

3. Results

3.1. Excluded participants

Two participants were excluded because of excessive head motion (this was defined by calculating, for each participant, the percentage of volumes per run with absolute translation values of over 2 mm or absolute rotation values over 1°, averaging these percentages over runs, and excluding any participant whose mean percentage was greater

than 2 standard deviations above the mean percentage across participants). One participant was excluded because of technical problems during data acquisition which meant that the pTx run failed. All subsequent analyses were conducted on the remaining 17 participants.

3.2. Behavioural results

 The 17 participants had good performance on both tasks and, importantly, there were no significant differences between the 5 sequences in terms of accuracy (Friedman's χ^2 = 5.26, p = 0.26) or reaction time (Friedman's χ^2 = 7.20, p = 0.16). The semantic and control tasks did not differ reliably in accuracy (Wilcoxon's z = 0; p = 0.06) or reaction time (Wilcoxon's z = 7; p = 1.00).

3.3. Region-of-interest analysis

ROIs that overlapped with the whole-brain contrast of interest (S > C) for at least one of the five sequences are shown in Figure 2.

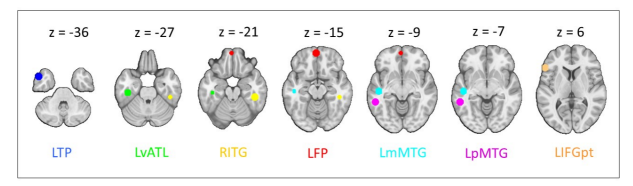


Figure 2: Regions of interest, taken from a meta-analysis of semantic tasks by (Humphreys et al., 2015). All spheres are 8 mm in radius. Regions of interest are overlaid on the MNI152NLin2009cAsym template. LTP = left temporal pole; LvATL = left ventral anterior temporal lobe; RITG = right inferior temporal gyrus; LFP = left frontal pole; LmMTG = left medial middle temporal gyrus; LpMTG = left posterior middle temporal gyrus; LIFGpt = left inferior temporal gyrus pars triangularis.

	LTP	LvATL	RITG	LFP	LmMTG	LpMTG	LIFGpt		
Activation magnitude									
pTx > SESB	-	0.2488	0.0240*	-	-	0.0042**	0.4114		
ME > SE	0.0502	0.0001**	0.0333*	0.2951	0.0092*	0.0003*	0.4155		
Activation precision									
MB > SB	0.4569	0.0007**	0.0019**	0.0989	0.2187	0.0266*	0.0016**		
MEdn > ME	0.3117	<0.0001**	<0.0001**	0.0014**	0.1363	<0.0001**	<0.0001**		
MBodd > SB	0.4225	0.1057	0.0481*	-	0.5697	0.1879	0.3300		

MVPA							
pTx > SESB	-	0.8569	0.3280	-	-	0.1529	0.9179
ME > SE	0.0096*	0.0063**	0.1612	0.0655	0.0465*	0.0024**	0.2508
MB > SB	0.2455	0.1417	0.0667	0.3021	0.3949	0.2693	0.5207
MEdn > ME	0.0027**	<0.0001**	<0.0001**	0.0023**	0.0001**	0.0012**	0.0007**
SB > MB	0.7416	0.4570	0.7222	-	0.3803	0.6716	0.3242
MBodd > SB	0.2584	0.5430	0.2778	-	0.6197	0.3284	0.6758

Table 2: p-values for all t-tests within regions of interest (ROIs) based on the semantic network. * = p < 0.05, ** = p < 0.05 (Bonferroni-corrected for the number of ROIs), - = ROI does not overlap by at least one voxel with the whole-brain contrast of interest (S > C) summed over all sequences in the comparison, LTP = left temporal pole, LvATL = left ventral anterior temporal lobe, RITG = right inferior temporal gyrus, LFP = left frontal pole, LmMTG = left medial middle temporal gyrus, LpMTG = left posterior middle temporal gyrus, LIFGpt = left inferior temporal gyrus pars triangularis.

3.3.1. Univariate analysis

Table 2 shows the p-values for all planned t-contrasts. pTx provided significantly better activation magnitude than SESB in the left posterior middle temporal gyrus (LpMTG). ME sequences provided significantly better activation magnitude than SE sequences in the left ventral anterior temporal lobe (LvATL).

MB sequences offered significantly better activation precision than SB sequences in the LvATL, right inferior temporal gyrus (RITG) and left inferior frontal gyrus pars triangularis (LIFGpt). MEdn sequences offered significantly better activation precision than ME in the LvATL, RITG, LpMTG, LIFGpt and left frontal pole (LFP). There was no significant difference in either direction between MBodd sequences and SB sequences (see Supplementary Table 1 for detailed results, plus reverse contrasts). We also found no significant interaction between multi-echo and multi-band.

3.3.2. Exploratory MVPA

Table 2 also shows the p-values for t-tests comparing our MVPA dissimilarity metric between sequences. The ME sequences produced significantly better performance than SE sequences in the LvATL and LpMTG. MEdn sequences produced better performance than ME sequences in every ROI. All other comparisons failed to reach significance.

3.4. Whole-brain analysis

Figure 3 shows the results for activation magnitude. Figure 3A shows a single cluster, within the right fusiform gyrus and lateral inferior occipital cortex, that showed greater activation magnitude with pTx than with SESB (pTx > SESB). Figure 3B shows the main effect of ME over SE, featuring clusters in the fusiform and inferior temporal gyri bilaterally plus the left orbitofrontal cortex (cluster and peak information is provided in Supplementary Table 2).

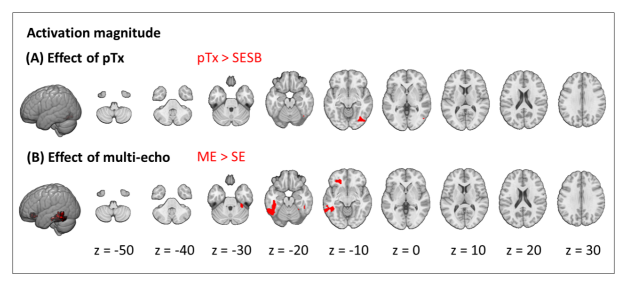


Figure 3: Effects on activation magnitude: (A) effect of parallel transmit (pTx > SESB); (B) effect of echo (ME > SE). Results are cluster-corrected at p < 0.05 based on an uncorrected voxel threshold of p < 0.001 and are overlaid on the MNI152NLin2009cAsym template.

Figure 4 shows the results for activation precision. Both the main effect of MB over SB and the main effect of ME-ICA denoising over ME without ICA denoising extended down the temporal lobe and included frontal regions (Supplementary Table 2). There was no significant difference between downsampled MB sequences and SB sequences in either direction.

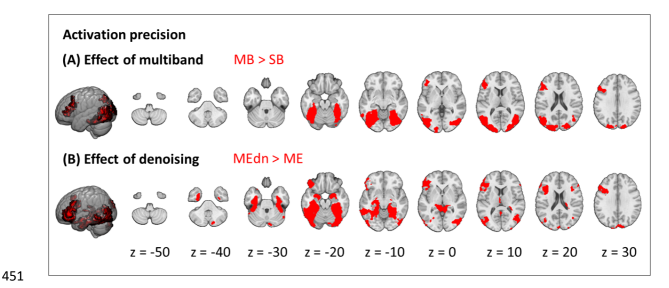


Figure 4: Effects on activation precision: (A) effect of multiband (MB > SB); (B) effect of ME-ICA denoising (MEdn > ME). Results are cluster-corrected at p < 0.05 based on an uncorrected voxel threshold of p < 0.001 and are overlaid on the MNI152NLin2009cAsym template.

To summarise, these results aligned with results from our ROI analyses – ME increased activation magnitude and both MB and ME-ICA denoising increased activation precision (see Supplementary Figures 1-6 for detailed results, plus reverse contrasts).

3.5. Slice leakage analysis

Seed and possible artefact locations for an example participant are shown in Figure 5A. Figure 5B shows activation magnitude for an example seed and its corresponding possible artefact locations for all participants in each MB sequence and its corresponding sequences. Figure 5C shows MVPA performance. Violin plots for all other peaks are shown in Supplementary Figure 7. t-tests were conducted at each location, but no comparison reached statistical significance (p-values are shown in Supplementary Table 3). We therefore concluded that there was no evidence for slice leakage in either of our MB datasets.

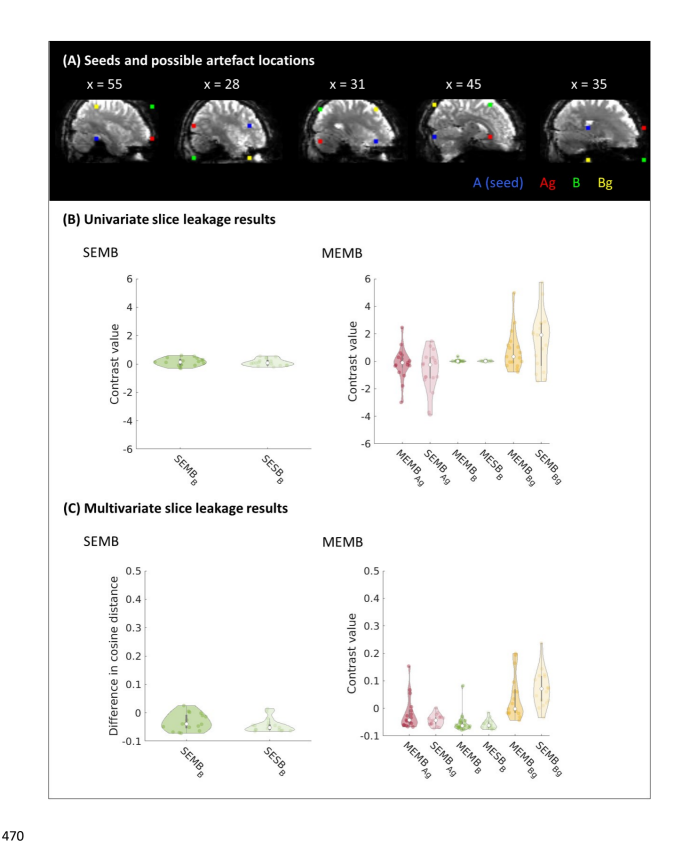


Figure 5: Slice leakage analysis. (A) Seed and possible artefact locations for a single participant. The ROIs (radius 4 voxels) indicate the seed location (blue), possible artefact location based on phase shift (green), possible artefact location based on GRAPPA (red), and possible artefact location based on phase shift and GRAPPA (yellow) in native EPI space. Informed consent was obtained from the participant for this image to be published. (B) Mean activation magnitude within each sphere for each MB sequence and the

corresponding control sequences for the first seed location (plots for other seeds are shown in Supplementary Figure 7; statistics for all seeds are shown in Supplementary Table 3). (C) Mean MVPA dissimilarity within each sphere for each MB sequence and the corresponding control sequences.

4. Discussion

The tSNR of fMRI varies across the brain. This is especially evident in ventral anterior temporal and orbitofrontal regions, which are located next to the air-filled sinuses and are therefore affected by signal dropout and geometric distortions (Devlin et al., 2000; Halai et al., 2014, 2015, 2024). This study is the first 7T-fMRI study systematically comparing pTx, ME and MB as methods for improving sensitivity in these regions while maintaining sensitivity across the brain. We found that pTx improved activation magnitude in posterior temporal and occipital regions. ME, however, resulted in improved activation magnitude extending down the temporal lobe and including inferior frontal regions. Both MB and ME-ICA denoising resulted in improved activation precision in the same areas. In an exploratory analysis we found that ME and ME-ICA improved MVPA performance but MB did not. No slice leakage artefacts were associated with our multiband sequences.

Although parallel transmit produced better activation magnitude in posterior temporal and occipital regions than the baseline (SESB) sequence without compromising activation magnitude elsewhere, activation magnitude in anterior temporal areas was comparable for both sequences. These results replicated those of Ding et al. (2022) who failed to find improved task contrast in anterior temporal regions with pTx in spite of improved tSNR in the resting state. Note that, despite visible signal dropout on the EPIs, even the baseline sequence was able to identify semantic activity extending into ventral anterior temporal regions (Figure 6). It is possible that this finding reflects the increased sensitivity of 7T-fMRI compared to 3T-fMRI, where semantic activity is rarely observed without using a method such as multi-echo (Halai et al., 2014, 2015, 2024) or spin-echo (Binney et al., 2010; Embleton et al., 2010). That a standard sequence can detect signal in these regions might come as a surprise to many neuroimaging researchers as 7T-fMRI is not only associated with increased sensitivity, but also with exacerbated magnetic susceptibility artefacts. This finding should therefore embolden researchers to use 7T fMRI for experimental designs that are difficult to conduct at a lower field strength - for example, studies of special populations, who may benefit from shorter scan times (Cope et al., 2023),

or studies requiring high spatial specificity (Marques & Norris, 2018). We do acknowledge that the way we have utilised 7T fMRI in this study will not be suitable for, for example, studies that require ultra-high resolution (in our case, scanner hardware constrained voxel size when combined with a short first echo time).

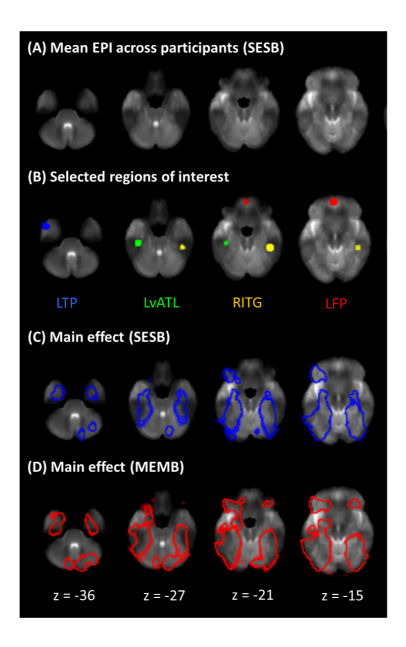


Figure 6: Contrast despite signal dropout and geometric distortions. (A) Mean EPI across all subjects for the baseline sequence (SESB) registered to MNI152NLin2009cAsym space. (B) Selected regions of interest overlaid on the mean EPI image. (C) Main effect of activation magnitude for the contrast of interest (S > C) for the baseline sequence. (D) Main effect of activation magnitude for the contrast of interest (S > C) for the MEMB sequence. Results are cluster-corrected at p < 0.05 based on an uncorrected voxel threshold of p < 0.001.

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The factorial design enabled us to disentangle the effects of ME and MB. As hypothesised, ME improved activation magnitude and these effects were localised to inferior temporal and orbitofrontal areas. Adverse effects on activation magnitude were negligible - although ME reduced activation precision in two very small clusters (Supplementary Figure 4), these were not within areas that the semantic task is known to recruit (Jung et al., 2017). These results demonstrated that having one very short echo is a successful method of increasing activation magnitude in areas where T₂* is particularly short (Halai et al., 2014, 2015, 2024; Jung et al., 2017). ME also opens up the opportunity for sophisticated denoising such as tedana (DuPre et al., 2021; Kundu et al., 2011, 2013; The tedana Community et al., 2023), which improved activation precision in agreement with previous findings (Amemiya et al., 2019; Gonzalez-Castillo et al., 2016). Small clusters in which ME-ICA increased activation magnitude (Supplementary Figure 5) or decreased activation precision (Supplementary Figure 6) were not within regions recruited by the task (Jung et al., 2017). As hypothesised, multiband improved activation precision in areas of inhomogeneity and in agreement with other work (Puckett et al., 2018). Supplementary Figure 4 shows two very small clusters in which multiband reduced activation precision, but which lie outside regions recruited by the task (Jung et al., 2017). After downsampling the MB timeseries to match the number of volumes in the SB timeseries, there was no longer any difference between the SB and MB sequences; this suggests that it is the increase in the number of volumes that accounts for the benefits of multiband. Additionally, there was no evidence for signal leakage into the simultaneously-acquired slice location (McNabb et al., 2020; Todd et al., 2016), at least for our modest multiband factor of 2. We note that MB and ME seem to have independent effects as the interaction was not significant within any ROI. However, ME seems to offer larger activation magnitude (Figure 6) and MB offers better activation precision. MB can help reduce the longer TR associated with ME and may decrease noise aliasing in studies using short TRs (~1s). Although this study focused primarily on univariate effects, we also identified improvements to multivariate metrics that will be of interest to researchers using increasingly-popular MVPA tools (Frisby et al., 2023). ME was found to improve MVPA performance compared to SE sequences and, in turn, MEdn sequences performed

significantly better than ME sequences without denoising in every region of interest (with

no detrimental effects). Future studies employing multivariate methods should strongly consider using both ME acquisitions and ME-ICA preprocessing.

This study provides evidence supporting use of ME and/or MB sequences to detect signal in regions prone to susceptibility artefacts. However, it is important to note that individual 7T MRI scanners have their own hardware and software limitations; this means that each site must optimise and test feasible parameters locally. For example, our system did not allow us to take advantage of the higher spatial resolution offered by 7T-fMRI while retaining an adequately short first echo for our multi-echo sequences. Puckett et al. (2018) used comparable voxel sizes but had a shorter first TE than us (9.9 ms vs our 11.8 ms); Miletić et al. (2020) had higher resolution (1.6 mm vs our 2.5 mm isotropic) and a shorter first echo of 9.66 ms. Additionally, our 7T Terra pTx system imposes very conservative SAR limits (1W per channel, 8W total) for pTx scans but permits 20W total (i.e. 2.5W per channel) for circularly polarised mode (CP-mode or "TrueForm") scans (equivalent to single transmit). We therefore had to use VERSE-modified pTx pulses which are known to be more sensitive to B₀ inhomogeneity. We also acknowledge that pTx can be combined with multiecho or multiband (Ding et al., 2023; Wu et al., 2016). Although we used the pTx head coil for all scans, the combination of sequences was not implemented on our scanner at the time of running this study and therefore needs to be tested empirically in future. Thus, rather than identifying the limits of what is possible, we offer an accessible framework that will enable researchers to leverage 7T-fMRI for investigating regions affected by susceptibility artefacts despite the hardware and software constraints of individual scanners.

5. Conclusion

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In this study we compared pTx, ME and MB as methods of improving sensitivity in temporal and frontal areas prone to magnetic susceptibility artefacts. Both pTx and ME improved activation magnitude, but only ME showed improvements in artefact-prone regions. MB and ME-ICA improved activation precision in these areas. Exploratory results suggested that both ME and ME-ICA may also benefit MVPA. We demonstrated that a multiecho, multiband sequence running on the 8Tx32Rx pTx head coil (in CP mode) can detect signal in susceptible regions while maintaining sensitivity across the whole brain and is therefore a versatile choice for future studies using high field strength and investigating the

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functional roles of ventral temporal and/or orbitofrontal cortex (Binney et al., 2016; Borghesani et al., 2016; Devereux et al., 2018; Duncan, 2010; DuPre et al., 2016; Fernandez et al., 2017; Lambon Ralph et al., 2017; Zahn et al., 2007). Data and code availability: Data will be made publicly available upon peer review and acceptance. Code is publicly available at: https://github.com/slfrisby/7TOptimisation/. **CRediT statement:** Saskia L. Frisby: Conceptualisation, Methodology, Investigation, Formal Analysis, Writing – Original Draft, Writing – Review & Editing, Visualisation; Marta M. Correia: Conceptualisation, Methodology, Writing – Review & Editing, Funding Acquisition; Minghao Zhang: Methodology, Writing – Review & Editing; Christopher T. Rodgers: Methodology, Writing - Review & Editing, Supervision; Timothy T. Rogers: Writing - Review & Editing, Supervision; Matthew A. Lambon Ralph: Writing – Review & Editing, Supervision, Funding Acquisition; Ajay D. Halai: Conceptualisation, Methodology, Formal Analysis, Writing – Review & Editing, Supervision, Funding Acquisition. Funding: This work was supported by an MRC Unit grant (SUAG/019 G116768) to M.M.C., an MRC PhD studentship (MR N013433-1) to M.Z., an MRC programme grant (MR/R023883/1) and intramural funding (MC UU 00005/18) to M.A.L.R, and an MRC Career Development Award (MR/V031481/1) to A.D.H. This work was also supported by the NIHR Cambridge Biomedical Research Centre (NIHR203312) and an MRC Clinical Research Infrastructure Award for 7T (MR/M008983/1). The views expressed are those of the authors and not necessarily those of the NIHR or the Department of Health and Social Care. **Competing interests:** C.T.R. receives research support from Siemens. Acknowledgements: We thank the participants and the Wolfson Brain Imaging Centre radiographers for their assistance. References

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