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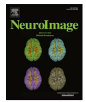
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Does higher sampling rate (multiband + SENSE) improve group statistics - An example from social neuroscience block design at 3T



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ABSTRACT

Multiband (MB) or Simultaneous multi-slice (SMS) acquisition schemes allow the acquisition of MRI signals from more than one spatial coordinate at a time. Commercial availability has brought this technique within the reach of many neuroscientists and psychologists. Most early evaluation of the performance of MB acquisition employed resting state fMRI or the most basic tasks. In this study, we tested whether the advantages of using MB acquisition schemes generalize to group analyses using a cognitive task more representative of typical cognitive neuroscience applications. Twenty-three subjects were scanned on a Philips 3 T scanner using five sequences, up to eight-fold acceleration with MB-factors 1 to 4, SENSE factors up to 2 and corresponding TRs of 2.45s down to 0.63s, while they viewed (i) movie blocks showing complex actions with hand object interactions and (ii) control movie blocks without hand object interaction. Data were processed using a widely used analysis pipeline implemented in SPM12 including the unified segmentation and canonical HRF modelling. Using random effects group-level, voxel-wise analysis we found that all sequences were able to detect the basic action observation network known to be recruited by our task. The highest t-values were found for sequences with MB4 acceleration. For the MB1 sequence, a 50% bigger voxel volume was needed to reach comparable t-statistics. The group-level t-values for resting state networks (RSNs) were also highest for MB4 sequences. Here the MB1 sequence with larger voxel size did not perform comparable to the MB4 sequence. Altogether, we can thus recommend the use of MB4 (and SENSE 1.5 or 2) on a Philips scanner when aiming to perform group-level analyses using cognitive block design fMRI tasks and voxel sizes in the range of cortical thickness (e.g. 2.7 mm isotropic). While results will not be dramatically changed by the use of multiband, our results suggest that MB will bring a moderate but significant benefit.

1. Introduction

Multiband (MB) or Simultaneous multi-slice (SMS) acquisition schemes allow the acquisition of magnetic resonance imaging (MRI) signals from more than one spatial coordinate at a time. Apart from the obvious advantage in the reduction of per volume acquisition times (Moeller et al., 2010), these sequences have been postulated to have several advantages for functional MRI (fMRI), especially, higher signal to noise per time unit, higher sampling rate resulting in higher statistical power and a better estimation of physiological noise (Barth et al., 2016; Feinberg and Setsompop, 2013; Olafsson et al., 2015). Most of the early testing of performance of MB acquisition employed resting state fMRI and showed decreased total scan times (X.-H. Liao et al., 2013), better noise estimation and spectral and spatial de-aliasing (Griffanti et al., 2014; Kalcher et al., 2014; Tong and Frederick, 2014a, 2014b; Tong et al., 2014) as well as better localization and estimation of functional networks (Koopmans et al., 2012). However, much less information is available about the benefits of MB acquisition for task based fMRI (Feinberg and Yacoub, 2012). Particularly, the utility of MB acquisition for group level statistics has not been evaluated yet. In this study we therefore performed comprehensive analyses to test the advantages of using MB acquisition schemes over single band EPI acquisition schemes

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in detecting task-based Blood Oxygenation Level Dependent (BOLD) responses.

The idea of acquiring data from different spatial locations simultaneously was first proposed in the 1980s (see Barth et al., 2016 for a historical review). However, the studies that brought widespread attention to the use of MB technique in fMRI were published in 2010 as part of the Human Connectome Project (Smith et al., 2013; Uğurbil et al., 2013). These two studies used multiplexed EPI combining simultaneous echo refocusing (SIR) and multiband radio frequency pulses and showed increased sensitivity to detect resting state networks (RSN) with 6 fold higher sampling rate at 3 T (Feinberg et al., 2010), and comparable activation between single band and MB acquisition using simple sensory tasks at 7 T (Moeller et al., 2010). Since then, MB sequences were improved to include multiple modalities (Cohen et al., 2017a), to reduce radiofrequency power deposition (Auerbach et al., 2013; Koopmans et al., 2012; Norris et al., 2011; Wu et al., 2013) and g-factor penalty due to the suboptimal reconstruction of the images (Setsompop et al., 2012), to ameliorate image reconstruction using controlled aliasing methods (Blaimer et al., 2013) such as radial CAIPIRINHA (Yutzy et al., 2011) and blipped CAIPI techniques (Setsompop et al., 2012), and by optimizing coil designs (Poser et al., 2014). Together with these technical advancements in MR pulse sequence development and image quality improvements, the commercial availability of compatible hardware and tailored acquisition and reconstruction software has brought the technique within the reach of many neuroscientists and psychologists, paving the way to systematic testing, validation and application to basic and clinical research settings (Chekroud et al., 2017; Cohen-Gilbert et al., 2017; Gabrielsen et al., 2018; Harenski et al., 2018; Kim et al., 2016; King et al., 2018; Kyathanahally et al., 2017; Provencher et al., 2018; Shah et al., 2016; Suri et al., 2017).

Since the publication of the first fMRI studies using MB acquisition in 2010 (Feinberg et al., 2010; Moeller et al., 2010), several studies have tested its benefits on resting and task based functional data. Using the keywords 'Multiband EPI', 'Multiband fMRI', 'Simultaneous Multislice EPI' and 'Simultaneous Multislice fMRI' in Pubmed on July 15, 2019, we identified 41 publications (including the two 2010 publications) that assessed the advantages of multiband acquisition for studying resting state and/or task based fMRI (Table 1). The data presented in these studies were collected on either 3 T or 7 T field strength scanners, generally using small to moderate sample sizes (median = 10; range = 3to 476), and different voxel sizes (ranging between 1.5 mm isotropic and 3.75 mm isotropic). Of these 40 studies, 20 studies used resting state acquisition not only to assess the performance of short repetition times (TRs) (Feinberg et al., 2010; Koopmans et al., 2012; X.-H. Liao et al., 2013; Preibisch et al., 2015) but also to harness the advantages of the short TRs in spectral de-aliasing of high frequency bands to study the BOLD specific as well as noise specific components in different frequency bands, and for ICA based de-noising methods (Boubela et al., 2013; Gohel and Biswal, 2015; Griffanti et al., 2014; Kalcher et al., 2014; Tong and Frederick, 2014a).

The main aim of the current study is to perform a detailed assessment of the effects of MB acceleration on random effects group level statistics for task-based fMRI, and identify the optimal acceleration for task-based fMRI. We therefore analysed the 21 studies that employed task based BOLD, in greater detail. Boyacioğlu et al. (2014) compared the performance of Gradient echo (GE) and Spin echo (SE) sequences in combination with MB factor of 3 and using a random effect analysis concluded that the performance of GE was superior in terms of the BOLD compared to SE. De Martino et al. (2015), ingeniously used longer silent periods between two TRs to present their auditory stimuli and showed a higher BOLD contrast with MB2 compared to single band, concluding that reducing the length of the scanner noise results in stronger auditory responses. Similarly, other studies that used MB accelerated sequences, but did not directly compare their performances for task related statistics were not analysed further (Shah et al., 2016; Vu et al., 2016). We identified three other studies that directly compared MB sequences for noise

amplification, slice leakage and serial autocorrelation matrices. These studies unequivocally showed that the noise is amplified as MB factor increases, leakage might occur at very high acceleration (>MB8) and that the conventional auto-correlation models might not be sufficient for data acquired with faster TRs (Bollmann et al., 2018; Boubela et al., 2014; Risk et al., 2018). Since group-level task-based BOLD was not the focus of these studies, we did not analyse them further. This left us with 13 studies which directly tested MB acquisition compared to the conventional single band acquisition or standard datasets to assess the benefits of higher sampling rate with short TRs on task-based fMRI. Collectively, these studies showed that using MB acceleration of 2-3 times might yield comparable or better statistics for task-based BOLD (Table 1). While these studies are valuable in enhancing our understanding of the effects of MB, there are several criteria, typical of modern cognitive neuroscience applications, not fulfilled in these studies and therefore warrants further investigation.

Up until now, the effects of MB acceleration have mostly been studied in basic functional designs using finger tapping and visual checkerboard (L. Chen et al., 2015; Chu and Noll, 2016; McDowell and Carmichael, 2018; Moeller et al., 2010; Sahib et al., 2018, 2016; Su et al., 2018; Todd et al., 2016). These tasks massively activate motor and visual cortices with strong contrasts between ON and OFF conditions. Since these are predominantly positioned in the cortex of the brain and are close to the MR receiver coils, higher acceleration factors are easily achieved. In contrast, modern cognitive neuroscience tasks often contrast tightly controlled conditions resulting in substantially smaller effect sizes than has been explored in past multiband studies. Further, cognitive tasks may accrue more within- and between-subject variability. Within-subject variability may increase because higher cognitive processing is less stimulus locked and may habituate more than early sensory or motor tasks. Between-subject variance may increase because complex cognitive tasks allow more individual variability in interpretation and depend more on past experiences. This will further reduce effect sizes. Whether MB provides advantages in the context of such smaller effects sizes and increased individual variability is difficult to predict. We thus feel that findings from MB-studies with basic sensory stimuli cannot be directly generalized to cognitive neuroscience applications. Moreover, while testing complex cognitive neuroscience applications, larger sample sizes are needed to reliably detect the BOLD as compared to basic stimuli where a sample size of <15 is enough for a reliable activation. A simple power calculation showed that with a sample of size 3 (smallest sample size reported in the literature reviewed in Table 1) one can expect detection of an effect with an effect size of 2.3, whereas using 14 subjects would allow us to detect an effect with effect size of 0.7. We included a sample of 23 subjects as recommended for cognitive tasks in fMRI by Thirion et al. (2007) which would allow us to observe the effects with effect size of ~0.5.

Furthermore, the majority of these 13 studies that tested the effects of higher sampling rate only report summary statistics such as mean t- or top 10% t-values within selected ROIs from subject level analysis (L. Chen et al., 2015; Chu and Noll, 2016; Kiss et al., 2018; McDowell and Carmichael, 2018; Sahib et al., 2018, 2016; Todd et al., 2017, 2016). Some studies additionally presented activation maps from representative subjects for visual comparisons (Moeller et al., 2010; Su et al., 2018), but only three studies presented voxel-wise, random-effects group-level, task-related BOLD statistics followed by a visual assessment of the performance of fast scanning.

In the current study, we add to our understanding of the effect of accelerated image acquisition and therefore higher sampling on task related BOLD statistics by addressing these gaps. We took advantage of a paradigm we have often used in our laboratory (Arnstein et al., 2011; Gazzola and Keysers, 2009; Gazzola et al., 2007; Valchev et al., 2016) and thus have reference data to systematically test the effect of MB acquisition. This paradigm aims to identify brain regions involved in the action observation network (Gazzola and Keysers, 2009) and involves observing complex goal directed actions (CA) and contrasting that activation

Summary of past studies exploring the benefits of MB acquisition and their main findings.

Technicus		Number of	Poet/Tealr (Main results
Technique		Number of subjects	Rest/Task (~scan time)	Main results
Field strength	MB factor(s)	Voxel size (mm ³)	TRs compared	
Feinberg et	al. (2010)			
SIR, MB		3	Rest (10 min)	Increased peak functional sensitivity.
3 T	1, 2, 3	$3 \times 3 \times 3$	2.0s, 0.8s, 0.4s	
-	et al. (2012)			
MB		6	Rest (15 min)	Exquisite localization to grey matter.
7 T	1, 4	1.5 imes1.5 imes 1.6	7.4s, 1.8s	
XH. Liao	et al., 2013)	1.0		
MB	,	11	Rest (6 min)	Validity of multiband rs-fMRI to reliably detect functional hubs.
3 T	1, 4	$3\times 3\times 3$	2.5s, 0.6s	
Boubela et	al. (2013)			
SIR, MB		10	Rest (6 min)	Resting-state networks like the default-mode network in frequencies above 0.25 Hz.
3 T	4	$2.4 \times 1.9 \times$	0.3s	
-	al (2014)	3.5		
Griffanti et MB	al. (2014)	76	Rest (10 min)	With optimal cleaning procedures, functional connectivity results from accelerated data were statistica
мв 3 Т	1, 6	$3 \times 3 \times 32 \times$	3.0s, 1.3s	comparable or significantly better.
	_, 0	$3 \times 3 \times 3 \times 2 \times 2 \times 2$	2.00, 2.00	
Tong et al.	(2014)			
MB		5	Rest (6 & 10 min)	Many voxels are highly correlated with pulsation regressors or its temporally shifted version.
3 T	6	$3\times 3\times 3$	0.4s	
Гong et al.	(2014b)			
MB		9	Rest (6 min)	Spatial distributions of different physiological processes are distinct.
3 T	6	$3 \times 3 \times 3$	0.4s	
Fong et al.	(2014a)	7	$Post(6 \ \ 10 \ \ min)$	Sustamia availations powered the POLD signal. Temporal traces evalue as the blood propagates theorem
MB 3 T	6	$3 \times 3 \times 3$	Rest (6 & 10 min) 0.4s	Systemic oscillations pervade the BOLD signal; Temporal traces evolve as the blood propagates though brain; They can be effectively extracted via a recursive procedure and used to derive the cerebral circulat
, 1	0	3 × 3 × 3	0.43	map.
Kalcher et a	al. (2014)			inspi
SIR, MB		20 + 20	Rest (6 min)	Correlations between resting-state signal fluctuations of distant brain regions even at high frequencies,
3 T	1, 4	$1.5\times1.5\times3$	1.8s, 0.3s	which can be measured using low-TR fMRI. In the high-TR data, loss of specificity of measured fluctuation
		$2.4 \times 1.9 \; \times$		leads to lower sensitivity in detecting functional connectivity.
		3.5		
Dlafsson et		10	Deat (10 min)	METCA identifies significantly more DOLD like components in the MECMC data as compound to data
	33-fold phase cceleration	12	Rest (10 min)	ME-ICA identifies significantly more BOLD-like components in the MESMS data as compared to data acquired with a conventional multi-echo single-slice acquisition.
3 T	1, 3	3.7 imes 3.7 imes 4	2.6s, 0.87s	acquired with a conventional multi-echo single-since acquisition.
	swal (2015)		2100, 01070	
MB		21	Rest (10 min)	Functional integration between brain regions at rest occurs over multiple frequency bands.
3 T	4	$3\times 3\times 3$	0.6s	
X. Liao et a	al., 2015)			
MB		11	Rest (10 min)	Economical, efficient, and flexible characteristics of dynamic functional coordination in large-scale hum
3 T	4	$3 \times 3 \times 3$	0.6s	brain networks during rest, and their relationship with underlying structural connectivity.
	t al. (2015)	20	Post (7 min)	MP factor of 2 only causes pacificials CNP decreases but reveals common PCN with increased consitivity
-	in-plane sensitivity acceleration	20	Rest (7 min)	MB factor of 2 only causes negligible SNR decrease but reveals common RSN with increased sensitivity a stability. Further MB factor increase produced random artifacts that may affect interpretation of RSNs un
	, 3, 4	3 imes 3 imes 3	2.0s, 1.0s, 0.7s,	common scanning conditions.
-, -,			0.5s	U ··· ···
Smith-Collin	ns et al. (2015)			
ИB		21	Rest (5 min)	Rapid rs-fMRI acquisition in neonates, and adoption of an extended frequency range for analysis allow
3 Т	1, 3	2.5 \times 2.5 \times	1.7s, 0.9s	identification of a substantial proportion of signal power residing above 0.2 Hz.
	1 (0015)	2.5		
Kalcher et a	ai. (2015)	15	Boot (7 min)	Craph disstoring based method for identifying yerrors usuals has a kish gravificity or 1 - 1 iting 1
MB 3 T	8	15 1.7 imes 1.7 imes 2	Rest (7 min) 0.3s	Graph clustering based method for identifying venous voxels has a high specificity and additional advantages of being computed in the same voxel grid as the fMRI dataset itself and not needing any
	-	10 0 10 0 4	5.05	additional data beyond what is usually acquired in standard fMRI experiments.
	et al. (2016)			
	d in-plane phase	24	Rest (15 min)	High resolution images acquired at 7 T provide increased functional contrast to noise ratios with
-	acceleration	0.0 0.0	07.10.10	significantly less partial volume effects and more distinct spatial features.
3 T vs 7 T	3, 5, 8	$0.9 \times 0.9 \times 0.9 \times 0.912 \times 12$	0.7s, 1.3s 1.9s,	
Т		$0.9\;1.2 imes 1.2 \ imes 1.2\;1.5 imes$	3.7	
		$^{\times}$ 1.2 1.5 $^{\times}$ 1.5 \times 1.5 1.6		
		\times 1.6 \times 1.6 2		
		$\times 2 \times 2$		
Reynaud, Jo	orge, Gruetter, Marqu		ag (2017)	
MB, 2D vs.		8	Rest (5 min)	After physiological noise correction, 2D- and 3D-accelerated sequences provide similar performances a
ΎΤ	1, 6, 8	$2\times 2\times 2$	3.3s, 0.6s, 0.4s	high fields, both in terms of tSNR and resting state network identification and characterization.
	. (2017b)			

Table 1 (continued)

Publicatior	1							
Technique		Number of subjects	Rest/Task (~scan time)	Main results				
Field strength	MB factor(s)	Voxel size (mm ³)	TRs compared					
MB 3 T	8	10 $2 \times 2 \times 2$ $3.5 \times 3.5 \times$ 3.5	Rest (7 min) 0.8s, 2.0s	Sensitivity and specificity increases and reproducibility either increases or does not change for the MB compared to the single band acquisitions. The MB scans also show improved grey matter/white matter contrast compared to the single band scans. The local functional connectivity density and global function connectivity density patterns remain similar across MB and single band scans and confined predominant to grey matter. A strong spatial correlation of functional connectivity density between MB and single ban scans is observed indicating the two acquisitions provide similar information.				
MB	et al. (2017)	12	Rest (12 min)	Physiological noise characteristics differ between SMS-EPI and regular EPI, with cardiac pulsatility contributing more to noise in regular EPI data but low-frequency heart rate variability contributing more t SMS-EPI. Signficant slice-group bias was observed in the functional connectivity density maps derived from SMS-EPI data. Making appropriate corrections for physiological noise is likely more important for SMS-EI than for regular EPI acquisitions.				
3 T	1, 3	$\begin{array}{l} 3.4\times3.4\times\\ 4.6\end{array}$	0.3s					
Smitha et a MB+2-fold accelerat	l in plane	9	Rest (7.4 min)	Negligible differences between the conventional-rsfMRI and MB rsfMRI acquisitions on the computed grap theoretic measures. MB-rsfMRI may be used as a time reducing acquisition technique that enables mappin of functional connectivity with similar outcome as conventional rs-fMRI in healthy subjects.				
3 T	1, 3	3.3 imes3.3 imes 3.2	3.0s, 0.9s					
MB	al. (2014)	10	Rest (7 min) & Task	Fast scanning may help to identify and eliminate physiologic components, increasing tSNR and functions contrast.				
	8 1 et al. (2015) -fold in plane	$1.7 \times 1.7 \times 2$	0.3s Rest (5 min) &	After noise correction, the detection of re-networks improves with more non-extension independent				
GRAPPA 7 T	-	$3.5 \times 3.5 \times$	Task 2.2s, 0.7s	After noise correction, the detection of rs-networks improves with more non-artefactual independent components being observed. Additional activation clusters for task data are discovered for MBME data (increased sensitivity) whereas existing rs-networks become more localized (improved spatial specificity				
Shah et al. SIR, MB	(2016)	3.5 476	Rest (15 min) &	Longer scan times are needed to acquire data on single subjects for information on connections between				
3 T	Not specified	2 imes 2 imes 2	Task 0.7s	specific ROIs. Longer scans may be facilitated by acquisition during task paradigms, which will systematically affect functional connectivity but may preserve individual differences in connectivity on to of task modulations.				
	et al. (2018) ne acceleration 2 1, 2, 3, 4, 6	$\begin{array}{c} 10+14\\ 3\times3\times3\end{array}$	Rest (6 min) & Task 2.0s, 1.0s, 0.7s, 0.5s, 0.3s	Strong benefits of the multiband protocols on results derived from resting-state data, but more varied effect on results from the task paradigms. Multiband protocols were superior when Multi-Voxel Pattern Analys was used to interrogate the faces/places data, but showed less benefit in conventional General Linear Mod analyses of the same data. In general, ROI-derived measures of statistical effects benefitted only modest from higher sampling resolution.				
	al. (2010) sampling factor of 4	3	Task	Task/stimulus-induced signal changes and temporal signal behavior under basal conditions were comparable for multiband and standard single-band excitation and longer pulse repetition times.				
7 T	4	$\begin{array}{c} 2\times2\times2\\ 1\times1\times2 \end{array}$	1.2s, 1.5s	comparable for manifolding and standard single band exercition and fonger parse repetition antes.				
MB (GE vs	tet al. (2014) SE), 3-fold in-plane tion, PINS 3	6 1.5 × 1.5 ×	Task 1.4s, 2.0s	GE-EPI shows higher efficiency and higher CNR in most brain areas; GE EPI was able to detect robust activation near air/tissue interfaces such due to reduced intra-voxel dephasing because of the thin slices used to be back in a loss area bick in a loss area bick.				
Chu & Nol		1.3 × 1.5 × 1.3	1.43, 2.03	used and high in-plane resolution.				
3 Т	ompression 1, 3	5 Slice thickness 3	Task 2.0s, 0.6s	Method to compress and reconstruct concentric ring SMS data improves preservation of functional activation over standard coil compression methods.				
ИВ 3 Т	o et al. (2015) 1, 2 t al., 2015)	$\begin{matrix} 6 \\ 2\times2\times2 \end{matrix}$	Task 3.0s	Reducing the length of the scanner noise results in stronger functional responses.				
SIR, MB 3 T	1, 2, 4, 6, 8, 10, 12, 14, 16	$7\\2.5\times2.5\times3$	Task 4.0s–0.2s	Low acceleration factors (N \leq 6), setting SIR = 1 and varying MB alone yielded the best results in all evaluation metrics, while at acceleration N = 8 the results were mixed using both S = 1 and S = 2 sequence				
Sahib et al. MB 3 T	1, 2, 4, 5, 8, 10	$\begin{array}{c} 15\\ 3\times3\times3\end{array}$	Task 2.6s–0.3s	Colored noise in event-related fMRI obtained at short TRs originates mainly from neural sources and cal for more sophisticated correction of serial autocorrelations which cannot be achieved with standard methods relying on AR(1)+w models with globally fixed AR coefficients.				
Fodd et al. MB, 2-fold accelerat	in-plane GRAPPA	10	Task	Imaging protocols using an acceleration factor of MB $2 \times$ GRAPPA 2 can be confidently used for high- resolution whole-brain imaging to improve BOLD sensitivity with very low probability for false-positive				
3 T	1, 2, 4, 6	1.5 imes1.5 imes 1.5	6.6s, 3.3s, 1.6s, 1.1s	activation due to slice leakage. Imaging protocols using higher acceleration factors (MB 3 or MB 4 \times				

(continued on next page)

Table 1 (continued)

Publication			
Technique	Number of subjects	Rest/Task (~scan time)	Main results
Field MB factor(s) strength	Voxel size (mm ³)	TRs compared	
			GRAPPA 2) can likely provide even greater gains in sensitivity but should be carefully optimized to minimize the possibility of false activations.
Kim et al. (2016)			
ME, MB, Thin-slice summatic 3 T 5	n 10 Slice thickness 4 vs. 1	Task 2.5s	The SMSME-thin imaging technique enhanced the temporal-signal-to-noise ratio and functional activation at high susceptibility regions of the brain.
Vu et al. (2016)		m 1	
MB 3 T & 7 6, 7 T	4 Slice thickness 3 at 3 T and 2.5 at 7 T	Task 0.5s	Substantial word timing information can be extracted using fast TRs, with diminishing benefits beyond TRs of 1000 ms.
Bollmann et al. (2018) MB, 2 or 3-fold GRAPPA	10	Task	Commonly used noise models, such as the AR(1) model, are inadequate for modelling serial correlations in
acceleration 7 T 4, 3	$2.5 \times 2.5 \times$ 2.5	0.6s, 2.0s	fMRI using sub-second TRs. Rather, physiological noise modelling in combination with advanced pre- whitening schemes enable valid inference in single-subject analysis using fast fMRI sequences.
	1.3 imes1.3 imes 1.3 $ imes$ 1.3		
Todd et al. (2017) MB, 12% in phase acceleratio	on 10	Task	Lower g-factor noise area of V1 shows significant improvements at higher SMS factors; the moderate-level g- factor noise area of the para-hippocampal place area shows only a trend of improvement; and the high g- factor noise area of the ventral-medial pre-frontal cortex shows a trend of declining t-scores at higher SMS factors. This spatial variability suggests that the optimal SMS factor for fMRI studies is region dependent. SMS accelerations of 4x (conservative) to 8x (aggressive) for most studies and a more conservative acceleration of 2x for studies interested in anterior midline regions is recommended.
3 T 1, 2, 4, 8	$3\times3\times2.5$	2.8s, 1.4s, 0.7s, 0.4s	
Kiss et al. (2018)			
MB 3 T 4, 6	$\begin{array}{c} 21 \\ 3\times3\times3 \end{array}$	Task 2.0s, 1.0s, 0.4s	\sim 4 min of the scan time with 1 Hz (TR = 1000 ms) sampling rate and \sim 2 min scanning at \sim 2.5 Hz (TR = 410 ms) sampling rate provide similar localization sensitivity and selectivity to that obtained with 11-min session at conventional, 0.5 Hz (TR = 2000 ms) sampling rate.
McDowell & Carmichael (20) MB	10	Task	Medeet TD reductions (to 1000 + 200 me) estimates increased event related BADI as formance in descendent.
мы 3 Т 1, 2, 3, 4	$2.5 \times 2.5 \times 2.5$	2.5s, 1.2s, 0.8s, 0.4s	Modest TR reductions (to 1000 \pm 200 ms) optimally improved event related fMRI performance independent of design frequency. Autoregressive models with a local as opposed to global fit performed better, while low order autoregressive models were sufficient at the optimal TR.
Sahib et al. (2018)			
MB 3 T 1, 2, 4, 8	$\begin{array}{c} 15\\ 3\times3\times3\end{array}$	Task 2.6s, 1.3s, 0.7s, 0.3s	At a conventional TR of 2.6 s, Functional Connectivity Degree (FCD) values were marginal compared to FCD values using sub-seconds TRs achievable with multiband (MB) fMRI.
Su et al. (2018)	20	Teels	Accelerated gradient echo (GRE) sequence combining simultaneous multislice excitation (SMS) with echo-
MB, 2-fold GRAPPA acceleration	20	Task	shifting technique for high spatial resolution BOLD fMRI has potential for high spatial resolution fMRI at
7 T 5, 1	1 imes 1 imes 2.5	3s	ultra-high field because of its sufficient BOLD sensitivity as well as improved acquisition speed over conventional GRE-based techniques.
Risk et al. (2018)	08	Tool	When date wars empothed we found avidance of alice lashene in some hut not all subject. We she found
MB 3 T 8 Corbin et al. (2018)	$\begin{array}{c} 98\\ 2\times 2\times 2\end{array}$	Task 0.7s	When data were smoothed, we found evidence of slice leakage in some, but not all, subjects. We also found evidence of SMS noise amplification in unprocessed task and processed resting-state HCP data.
MB+12% phase-over samplin	ng 10	Task	The "FAST" model implemented in SPM is used with a well-controlled number of parameters, it can
3 T 1, 2, 4, 8	$3 \times 3 \times 2.5$	2.8s, 1.4s, 0.7s, 0.35s	successfully prewhiten 80% of grey matter voxels even with volume repetition times as short as 0.35 s. Temporal signal-to-noise ratio can be augmented to account for the temporal correlations in the time series.
Current Study	22	Tools & Decude	(i) Sequences with different excelention feature are able to detect the brain notworks involved in tech
MB, 1.5 or 2-fold in-plane phase encoding acceleratio	23 n	Task & Pseudo resting state	(i) Sequences with different acceleration factors are able to detect the brain networks involved in task processing. (ii) Group level t-statistics improves with faster scanning. (iii) However it cannot compensate for
3 T 1, 2, 4	$\begin{array}{c} 2.7 \times 2.7 \\ \times 2.7 \\ 3 \times 3 \times \\ 3.3 \end{array}$	2.4s, 2.0s, 1.2s, 0.7s, 0.6s	the effects of larger voxel sizes, sample sizes or total scan duration. (iv) This is true for both task and resting- state analyses.

against complex control stimuli (CC) that include the same objects and the same hand, but without the hand manipulating the objects. This contrast is motivated by the fact that viewing objects and hands in motion would already activate a very broad network including early visual areas, attentional brain circuits, spatial representations in addition to regions specifically representing how the hand manipulates the object. To focus on the latter, we thus subtract stimuli that include the same basic visual elements and their spatial distribution and also include movement, but without the actual hand-object interaction (Gazzola and Keysers, 2009). This contrast is known to identify a broad network of brain regions (often referred to as the action observation network) encoding hand-object interactions in all participants via the CA-CC contrast including premotor, somatosensory, insula, inferior parietal, visual, and cerebellar regions (Gazzola and Keysers, 2009). This allowed us to assess the benefits of MB across a wide range of brain regions and their functional connectivity. Unlike most of the previous studies, we present voxel-wise group-level analysis with detailed post-hoc analyses to quantify the differences in the data acquired with different acceleration using a sample size representative of modern task-based designs (n = 23 subjects).

To derive a hypothesis for the expected outcomes, we looked at the underlying mechanisms that are inherent to the MB technology. On the one hand, we might expect multiband to improve second level randomeffect statistics. This is because multiband increases the number of samples that can be acquired in a given experimental time-window. More samples should increase the precision of the parameter estimates at the first level (Constable and Spencer, 2001). Because the second level residual variance is composed of the sum of 'true' between subject variance and first level errors in parameter estimates, this could directly improve statistics by reducing residual errors. On the other hand, other factors may curtail these benefits. First, measurement noise might increase due to imperfect separation of the aliased voxels (Golestani et al., 2017). Second, more samples only improve parameter estimates if the additional samples are independent, however the temporal auto-correlation increases as the samples are acquired closer in time (Bollmann et al., 2018; Corbin et al., 2018), limiting the independence of the samples and hence the benefits from acquiring more samples. Whether multiband acquisition leads to a net improvement of group-level statistics thus depends on the trade-off between these factors and is difficult to predict a priori. We therefore based it on the previous literature and hypothesized that the use of MB acquisition may show comparable if not higher t-values at a group level in response to task stimuli. Moreover, we look at individual factors such as subsampling the acquired volumes, tSNR and within and between subject variances to tease apart the contribution of these factors on our group level statistics. Finally, we performed a dual-regression analysis with the pseudo resting state data obtained by regressing out task-correlated activity to approximate the findings of the previous studies that showed an improvement of resting state statistics with MB acceleration. We hypothesized that on a group level, MB would outperform single-band acquisition in terms of the voxel-wise t-values.

2. Materials and methods

2.1. Subjects

Twenty-four (13 males, 11 females) right-handed (self-reported) healthy volunteers ($M_{age} = 25.5$ years, $SD_{age} = 3.6$, Range = 21 to 33) with no contraindications for MRI, and normal or corrected-to-normal vision participated in the study. After the screening procedure, subjects were familiarized with the MRI environment and were verbally informed about the observation task that they will watch during the session. The research was approved by the Amsterdam Medical Centre Ethics Committee Review Board, and the volunteers provided written consent at the start of the study. In what follows, sequences will be shorthanded as $MBxSy_{res}^{TR}$, where TR is the acquisition time of one volume, res the resolution in mm, and x and y, the multiband and SENSE factors respectively. One subject (all sessions) was removed from the analysis due to excessive head motion and the MB2S2^{1.22}_{2.7iso} session of another subject was removed due to technical error during the acquisition, resulting in a final sample size of N = 23 for $MB1S2^{2.45}_{2.7iso}$, $MB1S2^{2.00}_{3x3x3.3}$, $MB4S1.5^{0.70}_{2.7iso}$, $MB4S2_{2.7iso}^{0.63}$ and of N = 22 subjects for $MB2S2_{2.7iso}^{1.22}$ session.

2.2. Task and experimental procedure

Subjects were shown movie stimuli previously used in our lab to trigger robust activity in the action observation network (Gazzola and Keysers, 2009; Valchev et al., 2016). These movies show a human hand interacting with everyday objects. Examples of such interactions are shown in Fig. 1 and listed in Supplementary Table 1 (see also Valchev et al., 2016). Two types of stimuli were used: Complex actions (CA) showed the hand interacting with the object in typical, goal directed actions. For example, the hand of the actor reached for a lighter placed on the table, grasped it, and lit a candle with it. Complex controls (CC) stimuli had the exact same setting as the CA but the actor's hand did not interact with or manipulate the object on the table, instead, made aimless

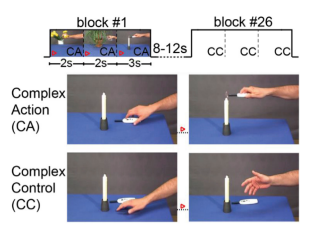


Fig. 1. Subjects observed 26 blocks of video stimuli per session (x 5 acquisition sequences) showing one of the two conditions: complex action (CA) or complex control (CC). Thirteen bocks per condition were presented and each block comprised of three clips of the same condition. The inter block interval was randomized between 8 and 12 s. The blocks and conditions were randomized between the five acquisition sequences per subject and between subjects. The top row schematically illustrates the structure of each run; the bottom two rows illustrate, for a randomly chosen clip, how it differs in the two conditions: a goal directed action in CA and a random movement in CC.

hand movements. A block was composed of three movies of the same category (CA or CC) and lasted 7s. Each fMRI session was composed of 13 blocks per stimulus category for a total of 26 blocks, presented in a randomized order. The inter-block-interval lasted between 8 and 12s and consisted of a fixation cross on a grey and blue background similar to the stimuli background. Five of these sessions were presented to each subject, showing the same blocks but in different order, with each session acquired with a different acquisition scheme, varying in MB factor, in-plane SENSE (S) acceleration and/or spatial resolution (Table 2). The order of acquisition was randomized between subjects. Importantly, the duration of the five sessions was similar (~8 min), but more functional volumes were acquired during sessions where the acquisition parameters lead to shorter TRs.

2.3. Image acquisition

Data were acquired on a 3 T Philips scanner, using a commercial version of Philips' MB implementation (based on software release version R5.4). A 32-channel head coil was used. The total length of the scanning session per subject was around 50 minutes and included the functional and anatomical scans. Functional data were acquired using five different acquisition sequences (Table 2). These sequences were chosen based on the following considerations. First, we wanted to acquire one of the most typically used non-multiband sequences as a meter of comparison, and therefore acquired a sequence with a TR = 2.00s and close to 3 mm isotropic resolution ($MB1S2^{2.00}_{3x3x3.3}$). Because multiband is typically used with slightly smaller voxels, we acquired all other sequences at a resolution of 2.7 mm isotropic, which is closer to the average grey matter thickness. At that resolution, we then measured an acquisition at MB1S2 $(MB1S2_{2.7iso}^{2.45})$, and increased MB to 2 and 4. At MB4, we additionally reduced SENSE acceleration to 1.5 as recommended by Philips, to mitigate potentially higher noise amplification. The SENSE reconstruction method has been characterized as an image domain "unfolding" algorithm. An acceleration (with reduction factor R) would result in a reduced FOV in every component coil image. Each pixel in the individual reduced FOV coil image will contain information from multiple (R), equidistantly distributed pixels in the desired full FOV image. Additionally, these pixels will be weighted with the coil sensitivity at the corresponding location in the full FOV (Blaimer et al., 2004). In our case,

Overview of the scanning parameters used for the five acquisition sequences and the reference study. Note (in red) that the sequence $MB1S2^{2,00}_{3x3x3.3}$ has a coarser spatial resolution compared to the other sequences. The sequence $MB4S1.5^{0.70}_{2.7iso}$ differs from $MB4S2^{0.63}_{2.7iso}$ in the SENSE acceleration and not in MB factor. Reference study was collected with a different set of subjects and a different scanner. It is used to compare some results from this study.

Sequence Acronym	MB1S2 ^{2.00} _{3x3x3.3}	MB1S2 ^{2.45} 2.7iso	MB2S2 ^{1.22} _{2.7iso}	MB4S1.5 ^{0.70} 2.7iso	MB4S2 ^{0.63} 2.7iso	Ref. Study
TR (seconds)	2.00	2.45	1.22	0.70	0.63	2.00
Multiband factor (MB)	none	none	2	4	4	none
SENSE acceleration (S)	2	2	2	1.5	2	2
Acquired voxel size (mm ³)	$3 \times 3 \times 3.3$	2.7 isotropic	2.7 isotropic	2.7 isotropic	2.7 isotropic	3.5 isotropic
Flip angle in degrees	75	79	64	51	50	70
Number of slices	36	44	44	44	44	41
Acquired volumes	245	200	400	700	780	345
Slice gap (mm)	0.33	0.27	0.27	0.27	0.27	0.00
Field of view (mm ³)	$240 \times 240 \times 130.3$	$216\times216\times130.4$	$216\times216\times130.4$	$216\times216\times130.4$	$216 \times 216 \times 130.4$	$224\times224\times143$

the reduction factor of 1.5 would result in reduction of the FOV in the phase-encoding direction by this factor. The resulting TRs were 2.45s, 2.00s, 1.22s, 0.70s and 0.63s for the sequences $MB1S2_{2.7iso}^{2.45}$, $MB1S2_{3x3x3.3}^{2.00}$, $MB2S2_{2.7iso}^{1.22}$, $MB4S1.5_{2.7iso}^{0.70}$ and $MB4S2_{2.7iso}^{0.63}$ respectively (Table 2). Slices were acquired in ascending order in transverse direction. TE was kept constant for all sequences at 30 ms. A default shift of 1/2 a FOV was used while acquiring with MB 2 and 4 to improve reconstruction quality. The flip angle was optimized to the Ernst angle of each TR. All functional images were reconstructed using the SENSE reconstruction algorithm (Setsompop et al., 2012) and this was done simultaneously with acquisition. Note that the sequence $MB1S2^{2.00}_{3x3x3.3}$ has a coarser spatial resolution compared to the other sequences, which would lead to differences that are not related to the total acceleration in acquisition. We therefore present the results of this sequence separately on figures rather than in the same line as the other sequences. In addition, the sequence $MB4S1.5^{0.70}_{2.7iso}$ differs from $MB4S2^{0.63}_{2.7iso}$ in the SENSE acceleration but not in MB factor. Any differences in these two sequences therefore will not reflect the effect of MB acceleration. A T1-weighted image was acquired at the end of the functional runs with a field of view of $240\times256\times250$ mm and the voxel size was 1 mm isotropic. Geometry factor (g-factor) maps (see supplementary analysis 1) that quantify the aliasing noise per voxel (Pruessmann, 1999) were obtained for sequences MB1S22.45 MB1S22.75a, $MB2S2^{1.22}_{2.7iso}$, $MB4S1.5^{0.70}_{2.7iso}$. Here the g-factor maps were computed on-scanner using the vendor's implementation and represent additional SNR penalty of multiband and in-plane SENSE acceleration. Note that the g-factor is computed from the coil sensitivity maps and the noise correlation matrix, both of which are measured in the preparation phase ahead of scanning (Pruessmann, 1999).

2.4. Reference study

As a reference for some of the analyses performed to test the effect of accelerated acquisition, we used a previously acquired dataset. Data from this previous study (Abdelgabar et al., 2018) consisted of fMRI data from an independent group of 31 subjects. Images included an anatomical scan and a functional run where the subjects viewed the same action observation paradigm used in this study (CA and CC blocks of 7s). In addition, they also saw thirteen 7s blocks of Static Control (SC) movies, which had the same elements as the CA and CC movies, but the hand lay still next to the objects without any movement. Data were acquired using a Philips Intera 3 T scanner (University Medical Centre Groningen, University of Groningen. Groningen, The Netherlands), using a 32-channel coil. First, a high-resolution, structural image (170 slices; scan resolution = 256 \times 256; field of view = 232×232 mm; voxel size: 1 mm isotropic) was acquired. Functional images were acquired using an echo planar T2*-weighted gradient sequence (See Table 2 for additional parameters). The aim of choosing this independent dataset as a reference instead of the average data acquired in the current study was to avoid circularity, which may result in inflated statistical estimates, and to use a study with an even larger sample size to further approximate the true population

activity that each multiband experiments aim to estimate. These data were acquired without any MB acceleration.

2.5. Preprocessing

Reconstructed whole brain functional data were preprocessed using SPM12 (Wellcome Trust Centre for Neuroimaging, UCL, UK) with Matlab version 8.4 (The MathWorks Inc., Natick, USA). Briefly, functional images were slice-time corrected and then realigned to the estimated average. Anatomical images were co-registered to the mean functional image (rigid body transformation, DOF = 6), and segmented. The normalization parameters that were generated during segmentation (unified segmentation) were used to bring all the images to the MNI space. The resampled voxel size for the functional images was $2 \times 2 \times 2$ mm and $1 \times 1 \times 1$ for the anatomical scans. We adjusted the SPM12 bounding box settings to [-90 -126 -72; 90 90 108] in order to include the cerebellum completely, as the default settings of [-78 -112 -70; 78 76 85] may result in omission of some of the cerebellar voxels (Abdelgabar et al., 2018; Gazzola and Keysers, 2009). Smoothing of 6 mm FWHM Gaussian kernel was applied to the functional data. The reference dataset was preprocessed similarly.

2.6. Subject level general linear models (GLM)

GLM with task predictors: Subject level GLM included CA and CC as two separate task predictors with each predictor having 13 blocks of 7s. The reference dataset additionally included SC as a predictor with 13 blocks of 7s. Boxcar functions were convolved with canonical haemodynamic response function implemented in SPM. Regressors included the six motion parameters estimated during realignment, first five principal components of cerebrospinal fluid (CSF) and five principal components of white matter (WM) (total 16 regressors). The CSF and WM principal components were extracted from the normalized but unsmoothed functional data for each subject using the average WM and CSF segments of the subjects' segmented and normalized anatomy, thresholded at 0.7 (arbitrary units) as masks (Behzadi et al., 2007). We used the principal component analysis based regressors as they have been shown to yield better results for physiological de-noising as compared to regressors generated based on cardiac and respiratory traces recorded with the respiration belt and pulse oximeter (Kirilina et al., 2016). FAST autocorrelation algorithm implemented in SPM12 was used to correct for temporal autocorrelations in the data which has been suggested to be superior to the default AR(1) models specially for the shorter TRs (Bollmann et al., 2018; Todd et al., 2016). All the other parameters, including the high-pass filter (128 s), were left as the default in SPM12. The residuals were saved and were used to calculate the effective tSNR (see supplementary analysis 2).

<u>GLM for resting state</u>: Previous studies with MB show that accelerating image acquisition may have benefits for detecting RSNs (Demetriou et al., 2018; Griffanti et al., 2014; Preibisch et al., 2015; Reynaud et al., 2017). To replicate these positive findings of MB acceleration on RSNs, in

this study we created a pseudo resting state dataset by regressing out the BOLD associated with task predictors CA, CC and the 16 nuisance regressors. Briefly, a subject level GLM was performed on unsmoothed functional data with the GLM described above (i.e. two task predictors, CA and CC, and the 16 regressors). The residuals were saved which were devoid of the BOLD that was detected as correlated to our action observation task. These residuals were smoothed with 6 mm FWHM Gaussian kernel and used in a spatial GLM in FSL (http://www.fmrib.ox.a c.uk/fsl/index.html) with 20 resting state networks (RSN20) as predictors (Smith et al., 2009). FSL was used as it has a well-implemented spatial GLM algorithm as a first step of the dual regression approach. The RSN20 maps classified by Smith et al. (2009) consist of 10 primary cortical networks, 3 cortical networks with partial spatial/functional overlap with primary cortical network(s), 3 networks with spatial location corresponding to subcortical or deep cortical areas and 4 artefactual components. This spatial GLM step generated 20 time-courses per subject per acquisition sequence, one for each RSN. These time courses were then used as predictors in a subject level GLM along with the 16 nuisance regressors for the normalized and smoothed data. Note that this analysis is very similar to the dual regression analysis implemented in FSL. We decided to perform the second step in SPM to keep the default processing (e.g. auto-regression) similar to those used in the task-based GLM which was performed in SPM, so as to avoid effects of using different packages and to enable qualitative comparisons between task based and resting state analyses.

2.7. Group statistics

Random effects group t-tests were performed separately for each acquisition sequence using the parameter estimates from the subject level GLMs (CA-CC contrast or RSNs). The significance of all group analyses was evaluated at q_{fdr}<0.05 at the voxel level and a cluster-size threshold of 50 voxels. We use FDR correction rather than FWE correction here to improve sensitivity (Lieberman and Cunningham, 2009; Lindquist and Mejia, 2015), but also provide histograms of t-values to enable scientists to appreciate the effect of MB at any threshold. When looking at the data at uncorrected p < 0.001 (see Supplementary Analysis 7), the FDRc cluster thresholds for different sequences ranged from 49 to 85. FDRc represents the minimum cluster size that limits the false discovery rate to 0.05. For visualization purposes, we therefore decided to use a common cluster threshold of 50 while evaluating the FDR corrected results.

Basic signal and noise characteristics of different acquisition sequence, which include g-factor, tSNR (predicted, effective and raw) and CNR are presented in supplementary materials (see Supplementary analysis). In the result section 3.1, we focus on the random-effect grouplevel task based BOLD outcomes for different acquisition sequences. We look at:

- (a) Voxel-wise t-statistics of individual sequences to see which sequences perform the best in terms of t-statistics. Next, using Receiver Operating Characteristics (ROC) on the resulting voxel-by-voxel t-values we analyse how similar/different the activation maps are between different acquisition sequences and the reference study. The ROC analysis allows us to look at the hit rate and false alarm rate in a non-threshold dependent manner. The intent of using the ROC measures is not to find a "winner sequence" but to see concordance between sequences. If a sequence would show poor concordance with all other sequences, it might warrant particular prudence. However, if all sequences show high concordance, this will point to the fact that although the t-statistics might differ between sequences, the sequence that one chooses would not greatly alter functional interpretation about the brain networks involved in the cognition of interest.
- (b) We then label the activated brain regions using the AAL atlas (Automated Anatomical Labeling, Tzourio-Mazoyer et al., 2002), and explore the similarity of conclusions at the brain-region level

using an ROC analysis (i.e. which AAL region are detected as active and which not).

- (c) To explain the small differences in the outcomes of different acquisition sequences we performed within and between subject variance analyses to see if there are systematic differences in the variance patterns between different sequences as that would directly impact the observed t-statistics.
- (d) Previous studies with resting state show that MB acceleration may enable us to shorten total acquisition times (Smitha et al., 2018). To test this theory for task-based studies, we truncated the total acquired data and looked if it would be possible to reduce the number of task trials/total acquisition time with less loss in accuracy at higher MB acceleration.

Section 3.2 reports the group statistics of the pseudo resting state data as replication for the widely reported positive effects of MB in the resting state literature.

3. Results

3.1. Random effect group level analysis

(a) <u>Voxel-wise analyses</u>: Fig. 2A and B presents the random-effect group activation t-maps for each sequence separately. As a validation, we also present the group maps of the reference study (Fig. 2C). Visual comparison of the group voxel-wise results look very similar across sequences, and consistent with what has been reported in Gazzola and Keysers (2009). Fig. 2D presents the histograms of the significant t-values for the sequences $MB1S2^{2,00}_{3x3x3.3}$ $MB1S2^{2.45}_{2.7iso}$, $MB2S2^{1.22}_{.27iso}$, $MB4S1.5^{0.70}_{2.7iso}$ and $MB4S2^{0.63}_{2.7iso}$. Both sequences with MB4 have higher t-values than sequences with MB 1 and 2 with same spatial resolution of 2.7 mm isotropic. Interestingly, this is the case despite the effective smoothness (especially average subject FWHM) being lower in the MB4 sequences compared to sequences with MB1 and 2 at the same acquired spatial resolution (Fig. 2A).

One notable difference in the visual inspection of the maps is that all of the sequences tested here activate fewer voxels than the independent study when visualized at the same threshold ($q_{fdr} < 0.05$). Fig. 2E shows that the independent study (black line) has much higher t-values than the two best sequences (pink and red lines in Fig. 2D) of the current study. This could potentially be due to three reasons: i) the independent study was conducted on 31 participants instead of the 23 participants included here, affording it more statistical power. ii) While in the reference study subjects viewed the stimuli only once, they saw them 5 times in our within-subject MB comparison, potentially leading to some form of repetition suppression. iii) The acquired voxel size in the reference study (3.5 mm isotropic) was larger than the resolution used here (2.7 mm isotropic or $3 \times 3 \times 3.3$ mm³). We therefore compared two more GLMs, one including just the first 23 subjects of the reference study and one with just the first view of the stimuli in the current study (Fig. 2C). Note that the "first view" GLM consists of data coming from different sequences. To assess the impact of voxel size, we compared MB1S2^{2,45}_{2,7iso} and $MB1S2^{2.00}_{3x3x3.3}$, as these differ only in the acquired voxel sizes.

Visual inspection of the maps confirmed the intuition that the sample size (N = 31 vs. N = 23) has an impact on the group BOLD levels, as although the networks look similar, the size of the clusters appears a bit smaller when fewer participants are included (see green arrows in Fig. 2C, and grey and black lines in Fig. 2E). Next, we considered the first view in the current study regardless of the acceleration with which it was acquired. Some clusters start to appear as significant (green boxes in Fig. 2C compared to maps in Fig. 2A and B), which were either not present or were smaller, when group analyses were done per sequence and therefore contained first to fifth view of the task. Fig. 2E shows that reference study (N = 23) vs. the first view in this study are very similar with slightly higher t-values for the reference study. Finally, the impact of

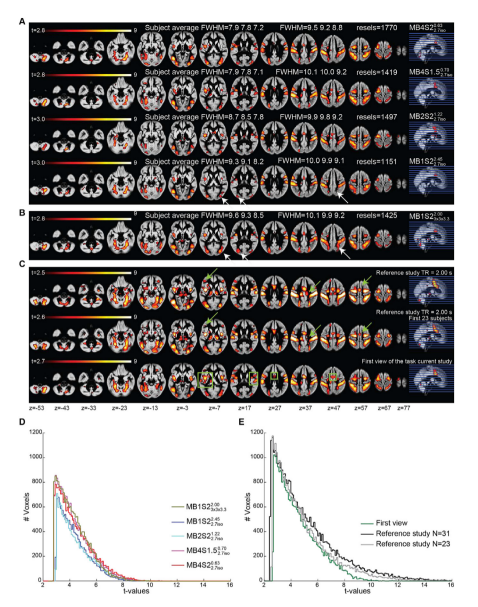


Fig. 2. All maps are overlaid on the mean grey matter segment of the group. q_{fdr}<0.05, cluster threshold 50 voxels. (A) Group maps showing the task correlated activity detected using the GLM predictors for the acquisition sequences $MB1S2_{2.7ico}^{2.45}$ $MB2S2_{2.7iso}^{1.22}$, $MB4S1.5_{2.7iso}^{0.70}$ and $MB4S2_{2.7iso}^{0.63}$. At the group level the effective smoothing is ~ 0.5 mm more in MB1S22.45 compared to MB4S22.063 and the average smoothness at the subject level is ~1.4 mm more for $MB1S2_{2.7iso}^{2.45}$ compared to $MB4S2_{2.7iso}^{0.63}$. (B) Group maps for the acquisition sequence $MB1S2_{3x3x3,3}^{2.00}$. White arrows represent the effect of voxel size on the BOLD outcomes. (C) Group maps from the reference study using the same task (N = 31subjects), maps from the reference study with a smaller sample of N = 23 subjects and from the current study looking at the first view of the task. Green arrows show how results change with the number of subjects. Green boxes represent the clusters that become bigger or more significant if we only consider the first view. (D) Histogram of the group t values for the CA-CC contrast in Fig. 2A and B. (E) Histogram of the group t values for the CA-CC contrast in Fig. 2C.

higher voxel size can be seen in sequences $MB1S2_{2.7iso}^{2.45}$ and $MB1S2_{3x3x3.3}^{2.00}$ which differ in the voxel size but not in the acceleration: white arrows in Fig. 2A and B show the areas with better BOLD for sequence with $3 \times 3 \times$ 3.3 mm³ voxels compared to sequence with 2.7 mm isotropic voxels (Fig. 2D and E). Although it has been argued that temporal SNR no longer depends on voxel volume beyond a voxel-size of 1.5 mm isotropic while using 32 channel coils (Triantafyllou et al., 2011), our results show that going from 2.7 mm isotropic to 3 \times 3 \times 3.3 mm 3 improves the tSNR (Supplementary analysis 2) as well as the t-values for task based BOLD (Fig. 2D). Taken together the findings of the random effect analysis suggest that in otherwise identical scanning parameters, MB4 shows higher t-values than lower acceleration with MB2 or no MB. Larger voxel sizes, larger group sizes and/or novelty of the task seems to show higher t-values. In our sample, MB1 sequence with larger voxels show t-values most comparable to MB4 sequences in terms of t-values, suggesting that MB4 can be used to obtain finer spatial resolution while keeping the t-sensitivity similar to a non-accelerated sequence with larger voxel sizes.

Since the GLM measures reported above are very threshold dependent, we used ROC measures to compare the conclusions one would

reach from different acceleration factors. Table 3 presents signal detection metrics (ROC, hit rate and false alarm rate) comparing the outcomes of these GLMs. The area under the ROC curve, which is less threshold dependent, is very high in all pairwise comparisons. It ranges from 85% to 90% when comparing the sequences against the reference study, and from 94% to 98% when comparing different sequences acquired here. The hit-rate was low when assessing how many of the voxels in the reference study were activated in each sequence (40% on average), and there was no systematic trend of this hit rate increasing or decreasing with the total acceleration. The hit rate was much higher when examining how much of the voxels from the current sequences were activated in the highly powered reference study (83-90%), and when comparing different accelerated sequences with each other (72% on average). The false alarm rate was consistently low, with on average only 2% of the voxels not activated in the reference study activated at any sequences, and only 3% of voxels not activated in one sequence activated in another. The sub-sample (N = 23) of the reference study showed high correspondence with the full sample (99%), had a high hit rate (81%) and a small false alarm rate (1%). The hit rate for the sequences against the subsample of the reference study was slightly higher (average 45%). The

ROC measures for assessing similarity/differences between the t-maps resulting from the group ANOVA of the different sequences. Ref = reference study. Ref 23 = reference study with first 23 subjects. FV = first view in the current study.

ROC		Reference								
		Ref	Ref 23	M1S2 ^{2.45} 2.7iso	MB1S2 ^{2.00} 3x3x3.3	MB2S2 ^{1.22} _{2.7iso}	MB4S1.5 ^{0.70} _{2.7iso}	MB4S2 ^{0.63} 2.7iso	FV	
Comparison	Ref		99%	95%	92%	95%	95%	94%	93%	
	Ref 23	99%		94%	92%	94%	94%	93%	92%	
	MB1S22.45 2.7iso	87%	88%		96%	95%	96%	96%	95%	
	$MB1S2^{2.00}_{3x3x3.3}$	88%	89%	98%		94%	97%	97%	95%	
	MB2S2 ^{1.22} _{2.7iso}	85%	86%	96%	94%		96%	94%	94%	
	MB4S1.5 ^{0.70} 2.7iso	90%	90%	97%	96%	96%		97%	97%	
	MB4S22.7iso	87%	88%	97%	97%	94%	97%		96%	
	FV	90%	90%	97%	96%	96%	98%	97%		
Hit Rate		Referenc	e							
		Ref	Ref 23	MB1S2 ^{2.45} 2.7iso	MB 1S2 ^{2.00} _{3x3x3.3}	MB2S2 ^{1.22} _{2.7iso}	MB4S1.5 ^{0.70} 2.7iso	MB4S2 ^{0.63} 2.7iso	FV	
Comparison	Ref		96%	90%	83%	89%	87%	85%	80%	
	Ref 23	81%		87%	77%	85%	80%	80%	73%	
	MB1S22.45 2.7iso	32%	37%		57%	68%	58%	60%	47%	
	MB1S2 ^{2.00} 3r3r3.3	45%	50%	87%		78%	75%	78%	63%	
	MB2S2 ^{1.22} _{2.7iso}	32%	37%	69%	52%		58%	55%	47%	
	MB4S1.5 ^{0.70} 2.7iso	46%	50%	86%	73%	84%		77%	68%	
	MB4S22.7iso	45%	50%	87%	75%	79%	76%		64%	
	FV	56%	61%	92%	81%	91%	90%	87%		
FA Rate		Reference								
		Ref	Ref 23	MB1S22.45 2.7iso	MB 1S2 ^{2.00} _{3x3x3.3}	MB2S2 ^{1.22} _{2.7iso}	MB4S1.5 ^{0.70} _{2.7iso}	MB4S2 ^{0.63} _{2.7iso}	FV	
Comparison	Ref		4%	14%	12%	14%	12%	12%	10%	
	Ref 23	1%		11%	9%	11%	9%	9%	7%	
	MB1S22.45 2.7iso	1%	1%		1%	2%	1%	1%	1%	
	MB1S2 ^{2.00} 3x3x3x3.3	2%	3%	5%		5%	3%	3%	2%	
	MB2S2 ^{1.22} _{2.7iso}	1%	1%	2%	2%		1%	2%	1%	
	MB4S1.5 ^{0.70} 2.7iso	2%	2%	5%	3%	5%		3%	1%	
	MB4S2 ^{0.63} _{2.7iso}	2%	2%	4%	3%	5%	3%		2%	
	FV	3%	5%	8%	6%	8%	5%	6%		

hit rate was also higher when only the first view (vs. the reference study) was considered. These matrices suggest that on average, the maps from different acquisition sequences have a high concordance.

(b) Region-wise analyses: Next, we labelled the brain regions using AAL (thresholded at q_{fdr} <0.05) to explore whether different sequences would lead to different conclusions about the brain network recruited by our task (Table 4). What is apparent from the table is that the core regions typically conceived of as part of the action observation network are found to be activated whatever parameters one uses (e.g. Postcentral, Precentral, Rolandic, and Supramarginal Gyrus). Such perfect agreement on the fact that a region is activated was true for 14 out of 62 brain regions in at least one hemisphere. There is also broad agreement on the fact that many regions are not activated (i.e. activated to less than 5%) in any of the sequences. This was true for 39 of the 62 brain regions in at least one hemisphere. In total, for 53 out of the tested 62 brain regions, scientists would thus arrive at the same conclusion whatever MB choice they would use. There was however a number of regions on which the different parameter choices did show substantial disagreement (highlighted in blue). It is notable, that regions of disagreement are often subcortical, including the cerebellum and basal ganglia.

While this provides a qualitative overview, to get a quantitative summary measure of how similar a conclusion would be drawn from such activation tables using different sequences, we examined ROC measures and correlation measured across the regions obtained using different sequences (Table 5). The ROC analysis revealed that the concordance has an area under the curve of at least 0.87 for the worst of the comparisons, and is very close to perfect (0.99) when comparing the highest acceleration ($MB4S2^{0.63}_{2.7iso}$) against the gold standard sequence without MB ($MB1S2^{2.00}_{3x3x3.3}$) that had voxels with 50% larger volume. Correlation measures on un-thresholded tables (in terms of percent activated) was at least 0.77 and on average 0.88, further supporting how similar a

conclusion would be reached. Binarizing the table in those regions activated (i.e. with at least 5% of the voxels in that region showing significant activation) or not, reduced the correlation to 0.74 on average.

Taken together these findings suggest that on a group level, with the exception of a few regions, one would reach very similar conclusions about the cortical networks that are involved in action observation, regardless of which acquisition sequence is used, with the highest acceleration allowing a voxel-volume reduction of 50% without changing conclusions.

(c) Between and within subject variances: The measurements with MB acceleration are noisier due to the imperfect separation of the simultaneously acquired voxels (Golestani et al., 2017). However, it also allows acquisition of many more samples which are thought to increase the temporal SNR per time unit (see supplementary analysis 1, 2). As the noise levels have a direct impact on the t-value estimation, we looked at the between and within subject variances to see if the marginally higher t-values in sequences with MB4 can be explained by systematic differences in the variances. One might expect that MB acceleration has benefits on within subject variance by providing more samples, but not on between subject variance, because we did not measure more participants at higher MB. We therefore performed a variance analysis as explained in Kirilina et al. (2016). The maps of the intra-subject variance (σ^2) were calculated using the following equation.

$$\sigma^2 = \frac{1}{N} \left(c \, X^{(1)-T} c^T \right) C^{(1)} \tag{1}$$

Here, N = number of subjects, c = contrast of interest (CA-CC), $X^{(1)}$ is the generalized inverse of the design matrix of the fixed effect analysis and $C^{(1)}$ is the residual variance maps (ResMS) generated while estimating the fixed effect ANOVA. Superscripted T signifies the transposition of the matrix. Since the random effect variance maps are a

AAL labelling of the brain regions activated for the CA-CC contrast at q < 0.05 level. Red colours are used to colour-code the level of activation in a region in terms of percentage activated. Blue cells are the brain region where there is a disagreement between different sequences, with some sequences showing more and some less than 5% activation (in both hemispheres). Note that for the vermis we show the joint activity of right and left hemisphere on the left side of the table. Values are % brain area active. Minimum red intensity is 5 and maximum is 50.

	Left hemisphere			Right hemisphere						
Brain region	MB1S2 ^{2,00} 3x3x3x333.3	$MB1S2^{2.45}_{2.7iso}$	$MB2S2_{2.7iso}^{1.22}$	$MB4S1 5_{27iso}^{070}$	MB4S2 ^{0.63} _{27iso}	$MB1S2^{2.00}_{3x3x33.3}$	$MB1S2^{2.45}_{2.7iso}$	MB2S2 ^{1.22} /2.7iso	MB2S2 ^{1.22} /2.7iso	$MB4S2_{27iso}^{0.63}$
Postcentral	30	26	33	32	30	33	32	33	39	39
Parietal Sup	45	29	27	37	39	29	14	16	25	20
Occipital Inf	39	34	30	34	42	24	13	11	13	18
Parietal Inf	35	29	27	34	36	25	17	14	20	23
SupraMarginal	30	28	21	39	34	23	18	14	26	22
Cerebelum 6	38	26	6	21	25	33	29	16	24	36
Occipital Mid	34	20	14	20	32	27	13	14	18	30
Fusiform	32	22	17	28	24	14	8	1	9	9
Precentral	17	20 3	26	27	25	5	7	8	9	7
Cerebelum 8 Rolandic Oper	5 18	3 11	0 17	8 17	11 24	28 6	23 4	16 17	23	25 8
Occipital Sup	26	6	6	17	24	25	4	0	16 9	11
Insula	19	16	12	26	19	5	4	6	4	4
Frontal Inf Oper	8	7	7	16	15	12	13	6	15	13
Vermis 8	28	0	3	5	18	 	10	-		10
Vermis 7	32	0	0	0	11					
Temporal Inf	9	6	5	8	7	10	7	4	9	9
Cingulum Mid	5	6	13	14	8	4	2	4	5	2
Cerebelum 4 5	5	3	4	6	4	9	6	3	9	10
Cerebelum 7b	2	1	9	2	3	8	8	0	12	11
Vermis 6	8	0	0	0	16					
Thalamus	1	0	4	14	0	0	0	7	0	0
Frontal Sup	6	5	7	12	9	5	5	7	8	7
Lingual	13	2	1	8	6	2	0	1	2	2
Cerebelum 9	0	0	0	0	0	13	16	0	0	8
Putamen	0	5	0	21	4	0	1	0	0	5
Frontal Sup Orb	0	0	0	0	0	0	0	0	0	0
ParaHippocampal	3	2	6	3	3	2	1	0 3	2	1
Supp Motor Area Pallidum	0 0	0 4	8 0	8 11	0 4	0 0	0 0	3 0	0 0	0 0
Hippocampus	2	4	6	0	4	0	0	0	0	0
Caudate	0	0	3	0	0	0	0	6	0	0
Heschl	1	0	5	0	3	 0	0	0	0	0
Temporal Mid	5	4	2	4	4	5	4	3	4	4
Cerebelum Crus1	2	1	0	0	1	0	1	0	0	3
Precuneus	3	0	1	4	4	1	0	0	1	1
Temporal Sup	2	2	2	2	4	0	0	1	1	0
Amygdala	0	4	4	0	0	0	0	0	0	0
Cingulum Ant	1	0	1	3	2	0	0	0	1	0
Vermis 10	4	0	0	0	0					
Frontal Mid Orb	0	0	0	1	0	0	0	0	0	0
Calcarine	2	0	0	0	0	1	0	0	0	1
Frontal Inf Orb	0	0	0	4	0	0	0	0	0	0
Angular Carabalum Crus2	0	0 0	0 0	0 0	0 1	4 1	0 0	0 0	1 0	0
Cerebelum Crus2 Frontal Mid	1 0	0	1	1	1	1	1	1	2	1 1
Cuneus	0	0	0	0	0	2	0	0	2	1
Paracentralobule	0	0	1	1	0	2	0	0	0	0
Frontal Inf Tri	0	0	0	0	0	0	0	0	0	0
Cerebelum 10	0	0	0	0	0	1	0	0	0	0
Vermis 1 2	0	0	0	0	0					
Vermis 3	0	0	0	0	0					
Vermis 4 5	0	0	0	0	0					
Vermis 9	0	0	0	0	0					
Cerebelum 3	0	0	0	0	0	0	0	0	0	0
Cingulum Post	0	0	0	0	0	0	0	0	0	0
Frontal Med Orb	0	0	0	0	0	0	0	0	0	0
Frontal Sup Medial	0	0	0	0	0	0	0	0	0	0
Olfactory	0	0	0	0	0	0	0	0	0	0
Rectus Tomporal Bala Mid	0	0	0	0	0	0	0	0	0	0
Temporal Pole Mid Temporal Pole Sup	0 0	0	0	0 0	0	0 0	0	0	0	0 0
remporal Pole Sup	U	0	0	U	0	U	0	0	0	U

combination of inter subject and intra subject variances, inter-subject variance maps (Σ^2) were calculated by subtracting σ^2 from the ResMS maps estimated during the random effect second level ANOVA. Five voxels with peak activities in the CA-CC contrast of the reference study (Fig. 2C) were selected as centers of 6 mm radius spheres (ROI1: -50, -24, 34 area PF of the inferior parietal lobe; ROI2: 40, -32, 44 Area 2 of the primary somatosensory cortex; ROI3: 30, -12, 60 superior frontal gyrus; ROI4: -40, -6, 4 insula lobe; ROI5: 38, -4, 10 area OP3 of the secondary somatosensory cortex; See inset in Fig. 3 for the location of each ROI). Fig. 3 presents the mean t-values for the fixed and random effect models, and within- and between-subject variances. As can be seen, some ROIs were dominated by within- (ROI4, 5) and some by betweensubject (ROI1, 2) variance, but in all cases, the MB4S1.5^{0.70}_{2.7iso} sequence outperforms the $MB1S2^{2.45}_{2.7iso}$ sequence in terms of t-values. Also, within subject variance was often but not always reduced in MB4S1.5^{0.70}_{2.7iso} vs. MB1S2^{2,45}_{2,7iso} sequences. There was thus no systematic relationship between benefits of MB against the dominant source of variance.

The inter-subject variance is a positively defined value. However, its estimation is obtained as a difference between the total variance and intra-subject variance in the second level analysis. In cases when inter-subject variance is much smaller than the intra-subject contribution, the difference might by chance become negative as can be seen in the case of ROI 5 and 6 where intra-subject variance dominates.

(d) GLM on first third of the data: With a specific total acquisition time, higher MB allows the acquisition of up to 4 times the number of volumes acquired with no MB. We thus explored whether the additional volumes per unit time at high MB can help when experimental time is limited. To test this, we repeated the subject level analysis using only the first third of each session for all five sequences. Group level t-maps were calculated using the CA-CC contrast image of each sequence separately and the resulting group t-map was correlated to the group t-map of the same contrast from the reference study. A similar correlation was also computed between the t-maps of the full GLM (see section 3.2) and the reference study.

As can be seen in the t-maps of the truncated dataset (one-third of the total scan) presented in Fig. 4A, many nodes that are a part of the action observation network can already be detected (see Fig. 2A-C for comparison). Moreover, looking at the histogram of these images, the overall performance pattern is similar to the results of the full dataset: higher acceleration result in higher t-values (Fig. 4B). $MB1S2^{2.00}_{3x3x3.3}$ which have bigger voxel size also shows higher t-values than the sequence with same acceleration but smaller voxel size (MB1S22.45). Looking at the correlation levels, we observed two phenomena. First, all of the truncated dataset were worse than the full dataset of the same sequence (Fig. 4C), suggesting that longer tasks are beneficial even at high multiband. Second, even high multiband acceleration failed to perform quite as well as the full dataset without MB acceleration (Fig. 4C comparing the solid blue point against the dashed green, cyan or red). Given that the dashed red point (i.e. 1/3 at $MB4S2^{0.63}_{2.7iso}$) actually has 1.33x as many functional volumes as the solid blue dot ($MB1S2_{2.7iso}^{2.45}$), this highlights that the number of dynamic volumes alone does not determine the reliability of activations. In summary, multiband is not a surrogate for total acquisition time.

3.2. GLM for resting state networks

Multiband has so far been mainly used and validated for resting state studies, where it has been suggested to improve the ability to detect RSNs. To explore whether we can replicate that finding in our data, we performed a pseudo resting state analysis with our data. The detailed description of the method can be found in section 2.6.

Fig. 5 shows a representative network (rsn06 in Smith et al., 2009) separately for each sequence. Overall, at the threshold of $q_{fdr} < 0.05$, the

ROC and correlations between the levels of activation labelled using AAL atlas. No threshold in Yellow and with 5% threshold in green.

ROC	Reference											
		MB1S2245 7iso	MB1S2 ^{2.00} 3x3x3.3	MB2S2 ^{1.22} _{2.7iso}	MB4S1 5 ^{0.70} 2.7iso	MB4S2 ^{0.63} 2.7iso						
Comparison	MB1S2 ^{2.45} 2.7iso		0.93	0.89	0.94	0.93						
	MB1S2 ^{2.00} 3x3x3.3	0.95		0.88	0.92	1.00						
	MB2S2 ^{1.22} _{2.7iso}	0.91	0.89		0.90	0.87						
0	MB4S1.5 ^{0.70} 2.7iso	0.96	0.92	0.92		0.93						
	MB4S2 ^{0.63} 2.7iso	0.97	0.99	0.89	0.94							

Correlation

	MB1S2 ^{2.45} 2.7iso	$MB1S2^{2.00}_{3x3x3.3}$	MB2S2 ^{1.22} 2.7iso	MB4S1 5 ^{0.70} 2.7iso	MB4S2 ^{0.63} 2.7iso
MB1S22 45 2 7 iso		0.87	0.87	0.92	0.93
$MB1S2^{2.00}_{3x3x3.3}$	0.77		0.77	0.84	0.94
MB2S2 ^{1.22} 2.7iso	0.64	0.62		0.90	0.86
$MB4S1.5^{0.70}_{2.7iso}$	0.78	0.77	0.63		0.92
MB4S2 ^{0.63} 2.7iso	0.81	0.92	0.62	0.81	

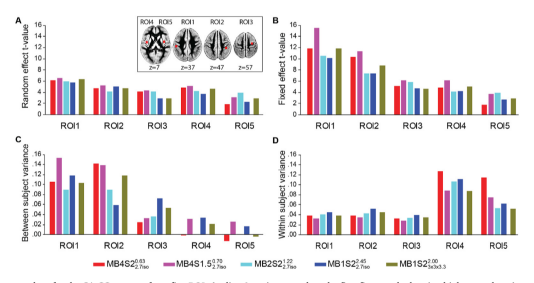


Fig. 3. Mean parameter values for the CA-CC contrast from five ROIs (radius 6 mm) centered on the first five voxels showing highest t values (at correction q_{fdr} <0.05) for the CA-CC contrast in the reference study. Figure A and B show the mean t-values from the random effect model and the fixed effect model, respectively. Figure C and D show the between-subject variance and the within-subject variance in these ROIs, respectively. Inset shows the location of the five ROIs.

network looks very similar for all the five sequences. Careful visual examination shows a small decrease in the cluster sizes as the TR increases (see white arrows and white boxes in Fig. 5 for examples).

To quantify this, we counted the total number of voxels that were significant at q_{fdr} <0.05 for all 20 networks. Fig. 6 present the average number (over all networks) of voxels as a function of acquisition sequence. An ANOVA was computed in SPSS with 20 components and 5 sequences and revealed a main effect of sequence (F(4,76) = 28.82, p < 0.001). Post-hoc pairwise comparisons show that the slowest acquisition ($MB1S2^{2,00}_{3x3x3,3}$, $MB1S2^{2,45}_{2,7iso}$) have fewer significant voxels than any of the MB accelerated sequences ($MB2S2^{1.22}_{2,7iso}$, $MB4S1.5^{0.70}_{2,7iso}$, $MB4S2^{0.63}_{2,7iso}$). On average across all RSNs, sequence with $MB2S2^{1.22}_{2,7iso}$ show a 15% increase in the number of significant voxels and sequence with $MB4S1.5^{0.70}_{2,7iso}$ and $MB4S2^{0.63}_{2,7iso}$ show 35% and 38% increase respectively, as compared to $MB1S2^{2,45}_{2,7iso}$, $MB1S2^{2,45}_{2,7iso}$, $MB1S2^{2,00}_{2,7iso}$ and the lowest number of voxels significant at

q_{fdr}<0.05.

To explore this phenomenon in a way that is less threshold dependent, and separately for each rsn, we plotted normalized cumulative histograms in Fig. 7. Using $q_{fdr} < 0.05$ for each rsn for $MB1S2_{2.7iso}^{2.45}$, we counted the number of significant voxels, N_{ref} . For any higher point along the x axis, we plot $N(t \ge x)/N_{ref}$. Accordingly, for $MB1S2_{2.7iso}^{2.45}$, the first point along the x axis has a value of 1, which then decreases as a higher threshold is used while moving to the right. Values of other x-coordinates of sequences can then be directly understood as the number of voxels surviving that threshold relative to N_{ref} . Plots framed in the orange outline show the component for which $MB2S2_{2.7iso}^{1.22}$ performs best, and plots framed in red are the ones where sequences with no MB show the highest number of supra-threshold voxels. All the others components showed that $MB4S1.5_{2.7iso}^{0.70}$ or $MB4S2_{2.7iso}^{0.63}$ indeed identify more voxels across a wide range of thresholds. The two components (in red) which do

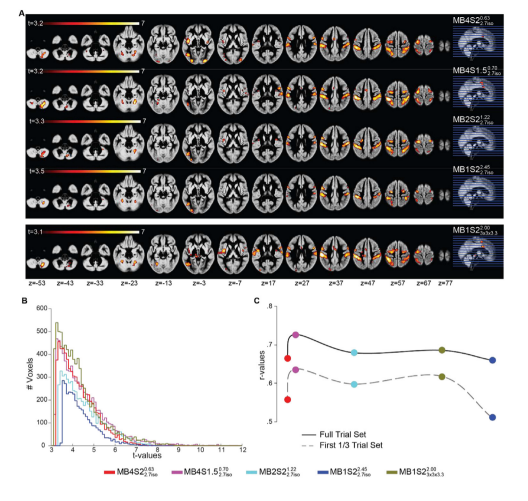


Fig. 4. (A) Group maps showing the task correlated activity detected using the task GLM predictors, but using only the first one third of the total acquisition per sequence. Overlaid on the mean grey matter segment of the group. $q_{\rm fdr}$ <0.05, cluster threshold 50 voxels. (B) Histogram of the t values for the CA-CC contrast for the one third of the total acquisition per sequence. (C) Correlation between t-maps of the CA-CC contrast per sequence and t-maps of the same contrast from the 31 subjects of the reference study.

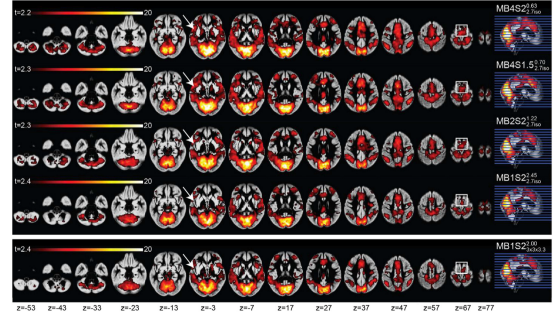


Fig. 5. Group maps per acquired sequence showing a representative RSN: network number 6, as described in Smith et al. (2009). White arrows and rectangles evidence areas with visually different cluster extension across different sequences. Maps overlaid on the mean grey matter segment of the group, and thresholded at $q_{fdr} < 0.05$ with a minimum cluster size of 50 voxels.

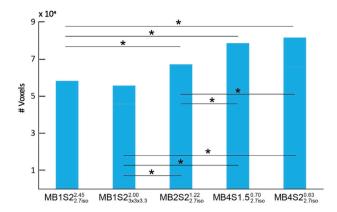


Fig. 6. Total number of voxels that are significant at $q_{fdr} < 0.05$ for each sequence. (*p < 0.001 for between sequence comparisons.)

not show an advantage of MB are classified by Smith et al. (2009) as belonging either to deep brain regions, or are artefactual components. However, since 17 out of 20 components show that using MB4 would yield higher t-values, this analysis recommends the use of MB 4 while studying RSNs at the group level, if the aim is to be inclusive. One unexpected finding of this analysis was that the $MB1S2^{2.00}_{3X3x3.3}$, which was excellent for task based fMRI (performing better than $MB1S2^{2.45}_{2.7iso}$ and similar to the $MB4S1.5^{0.70}_{2.7iso}$ and $MB4S2^{0.63}_{2.7iso}$), was one of the least sensitive sequence for the RSNs. The potential reason for this curious finding is discussed in the discussion section.

4. Discussion

We tested if acquisition with a shorter TR and therefore higher sampling rate – as afforded by MB – may improve our power to identify the neural substrates of cognitive functions. Based on our analysis of taskbased fMRI data, for group level analyses, the use of MB acceleration is beneficial to improve voxel-wise statistics. In our Philips

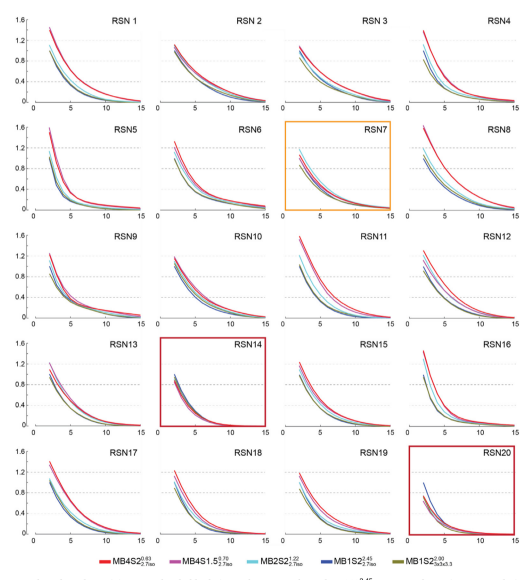


Fig. 7. Per component number of voxels surviving any t-threshold relative to the non-accelerated $MB1S2_{2.760}^{2.45}$ sequence for resting state analysis. The x-axis represents the voxel wise t-values, and the y-axis the number of voxels surviving that threshold relative to the number of voxels surviving t ≥ 2 at $MB1S2_{2.760}^{2.45}$. MB4 sequences are most inclusive, except for the plot framed in orange and red, where $MB2S2_{2.760}^{1.22}$ and $MB1S2_{2.760}^{2.45}$ sequences include the largest number of voxels.

implementation, sequences with MB4 acceleration (both S1.5 and S2) and 2.7 mmisotropic voxels, showed the highest t-statistics in our taskbased fMRI analysis. Interestingly, this was the case despite the MB4 sequence generating data with less effective smoothing than the sequences with lower MB acceleration. This is surprising, because more smoothing normally benefits t-values and reduces the correction for multiple comparisons, and would thus be expected to put our MB4 sequences at a disadvantage.

Other than the effect of MB acceleration, there were three main factors that influenced the group statistics. First, the impact of voxel size is well known (Robinson et al., 2008) and with this data we show that while for tSNR, MB2 is enough to compensate for 50% reduction in voxel volume (see Supplementary analysis, tSNR section, Supplementary Fig. 3), when it comes to the group level t-values, MB4 had t-values that were similar to the t-values seen with MB1 sequence with $3 \times 3 \times 3.3$ mm³ voxel size (Fig. 2B, E). Furthermore, higher t-values were observed when the group sizes were bigger (Turner et al., 2018) and the subjects viewed the stimuli only once (Larsson and Smith, 2011). While these are indirect observations from the data presented here, it can be clearly seen from Fig. 2 C-E that MB does not compensate for factors such as larger sample sizes and repetition suppression effect. While these contributions of voxel size, group size and multiple view is not a novel discovery, the use of the higher-powered reference study helps visualize the magnitude of t-value gains with MB acceleration in a new light. It suggests that the effect sizes of the gains are not big enough to nullify the effects of these factors.

It should be noted that all the sequences used in the current acquisition were able to identify the core regions that have been shown to be a part of the action observation network (Abdelgabar et al., 2018; Gazzola and Keysers, 2009). ROC analyses suggest a strong concordance between the outcomes of different sequences (Tables 3 and 5). So if one selects any of the sequences presented here, they would reach more or less similar conclusion about the brain regions implicated in the cognition of interest. Looking at the areas where we found disagreement between the sequences, we notice that they particularly include subcortical regions including the cerebellum and basal ganglia, which are detected more often in sequences with MB acceleration or with the sequences with bigger voxel sizes (see blue box in Table 4). This finding further supports the use of MB acceleration for detecting task correlated BOLD particularly if one is interested in these regions. Moreover, using dual regression analysis we show similar benefits of MB acceleration on RSNs (Figs. 5-7) supporting the evidence present in the literature that tests MB acquisition using resting state fMRI (Feinberg et al., 2010; Griffanti et al., 2014; X.-H. Liao et al., 2013).

We looked at inter and intra-subject variability in an attempt to shed light on how MB benefits group analyses. In our previous study where we tested standard single-shot 2D echo planar imaging (EPI) to three advanced EPI sequences, i.e., 2D multi-echo EPI, 3D high resolution EPI and 3D dual-echo fast EPI, inter-subject variability had a major impact in determining the sensitivity of the group-level analyses (Kirilina et al., 2016) in almost all brain regions except of prefrontal cortex, thereby limiting the potential for improving results by improving the measurement of each subject. However, in the current study, we found that the contribution of intra- and inter-subject variance to the random effect group level analysis was comparable. Therefore the dominant form of variance varies across regions, making the distinction of between- and within-subject variance less fruitful in understanding how MB benefits the group analysis (Fig. 3).

Looking at other image statistics such as G-factor and tSNR, we see that g-factor values increase with higher MB suggesting increase in noise due to poorer separation of the aliased voxels (Supplementary Fig. 1). However, correcting for the number of acquired volumes, the predicted tSNR and the measured effective tSNR (Supplementary Table 2, Fig. 2) both improve with higher MB factor. Since the variance shows no systematic differences (Fig. 3) and the noise measures increase with higher MB (Supplementary Analysis 1, 3), increasing the sampling rate is the most likely explanation for the better t-statistics with higher MB, and appears to have outweighed the disadvantages of MB (e.g. reduction in raw tSNR and increased autocorrelation of samples).

To explore whether the additional volumes acquired with higher MB can compensate for performing fewer repetitions of the task, we truncated the data and examined the first third of each experiment. With one third of the MB4 acquisition containing as many samples as the full MB1 acquisition, one might have expected the former to generate maps quite similar to the latter. However, we see that higher multiband did not show benefits in terms of localizing activations more reliably when fewer repetitions of a condition are available. This is contrary to what has been claimed for rs-fMRI (X.-H. Liao et al., 2013). This inability of the additional data points obtained using multiband to replace those obtained by a longer task is similar to the recent findings showing that temporal down sampling by randomly removing up to 50% of time points has little effects on BOLD reliability, while truncating the datasets is associated with decreased reliability (Shah et al., 2016), and may reflect the fact that temporal autocorrelation in fMRI data may make adjacent data-points acquired with short TR comparatively redundant.

Taking these factors together, our data suggests that on our Philips 3 T scanner, and implementation of multiband (based on scanner software release version R5.4), faster scanning offers modest but significant benefits for group-level voxel-wise task-correlated statistics and can be used as a better alternative to the single-band EPI, if other variables such as voxel size, scan duration and sample size are identical. Using MB factor of 4 with in plane SENSE factor of 1.5 or 2 and spatial resolution of 2.7 mm isotropic seems superior to the other sequences used here. However, when deciding whether to invest into multiband technology for cognitive neuroscience paradigms, our analyses suggests that for task-based studies, results similar to MB acceleration of factor 4 can be achieved by increasing the voxel size. This however might not be true for resting state studies. In our resting state analysis, sequence with MB1 and 3 \times 3×3.3 mm³voxel size showed the lowest t-values. The reason for the observed better connectivity estimates for smaller voxel sizes at MB1 is unclear. One potential reason could be a reduced intra-voxel dephasing due to reduced field inhomogeneity for smaller voxel sizes.

One important thing to note here is that our study focuses on comparing group-level statistics. This was chosen, because most cognitive neuroscience studies draw their conclusions from such group studies with ~20 participants. An interesting additional question is whether MB would alter the t-values obtained at the single subject level. This might be particularly interesting when studying rare patients, for instance, or performing pre-surgical scanning. Even though each of our subjects was scanned with all sequences, we found it difficult to answer this seemingly simple question. This is due to two reasons. First, the difference in TR across sequences substantially changes the white-matter/grey-matter contrast (Supplementary Figs. 6 and 7). As a result, the global mean normalization that is part of the traditional SPM pipeline applies different normalizing factors to images from different TR, and hence makes the beta maps non-comparable. Second, sequences with shorter TR acquire more samples and their single subject t-values thus need to be compared against different critical t-values due to the difference in degrees of freedom. This makes comparing the single subject t-values across sequences meaningless. We tried to compensate for this effect by z-transforming the p values (invnorm(p)) or comparing effect sizes (t/sqrt(df)), but found that doing so leads to counterintuitive results that suggest potential issues with the calculation of degrees of freedom. For a detailed analysis on this topic using the same data presented here, see (Bhandari et al., 2019).

To assess the task correlated BOLD, we convolved the boxcar with the canonical haemodynamic function. It assumes that the BOLD response peaks approximately 5 s after stimulation, and is followed by an undershoot. While this may be true in the primary sensory areas such as VI and M1, this pattern may vary in other brain regions and between subjects. Several alternative measures have been proposed for modelling the HRF (Badillo et al., 2013; Kumar and Penny, 2014; Quirós et al., 2010). A systematic study of some of the commonly used algorithms and

alternatives for modelling the haemodynamic response concluded that it is surprisingly difficult to accurately recover true task-evoked changes in BOLD signal and that there are substantial differences among models in terms of power, bias and parameter confusability (Lindquist et al., 2009). One should bear in mind the settings used here while interpreting the findings.

To briefly elaborate on this issue with our own data, we re-analysed our data using the additional time and dispersion derivatives that allow us to capture differences in the latency and the duration of the peak response, respectively. The results (Supplementary Analysis 6) show that the network recruited by the task remains largely unchanged. Moreover, the histogram of the t-values shows that overall, MB4 accelerated sequences still performs better than the sequences with no or lower MB accelerated sequences as seen while modelling only with the canonical HRF.

Our study has a number of strengths and limitations that should be considered. Limitations: Because we only used a block design task, our study cannot address whether event-related designs may benefit more from increased acquisition speed, and this should be investigated in future studies. Theoretically, if the contribution of the white noise (dependent on the subject and the MRI system electronics) in the data is not substantial, then one may expect a higher gain in GLM activation with increasing task frequency (characteristic of event related designs) with shorter TRs (Chen et al., 2019). Several studies have investigated the effect of MB acceleration on event-related designs (Demetriou et al., 2018; McDowell and Carmichael, 2018; Sahib et al., 2018, 2016). Together these studies showed that there might be benefits of using MB acceleration for task based paradigms while analysing functional connectivity and task-based BOLD contrasts. However, these effects might be more nuanced and may depend on more factors than just the task design, as Demetriou et al. (2018) specifically showed that while the ROI analysis were inconsistent, the benefits of using MB became apparent only when using multivariate pattern analysis. Also, whether cognitive neuroscience questions which can use MB acquisition without decreasing the TR and thus using the silent periods for presenting auditory stimuli (De Martino et al., 2015) or performing simultaneous EEG recordings (Uji et al., 2018) or other electrical recordings that may benefit from reduced interference from the radio-frequency pulses from fMRI, have not been tested here. Next, higher sampling with MB acceleration allows acquisition of high frequency components. Here we do not explore whether these higher frequencies, which are absent in the HRF convolved predictor, contain task-induced neural information. Moreover, shorter TRs offer advantages for spectral de-aliasing (Tong and Frederick, 2014a; Tong et al., 2014). Here we decided not to perform low-pass filtering because studies have shown the presence of BOLD like components in high frequency bands (Boubela et al., 2013). Here we employ 16 regressors including the six motion and ten principal components from CSF and WM (CompCorr, Behzadi et al., 2007) to de-noise the data. However, the effect of other noise cleaning procedures that have been proposed for cleaning MB data such as FIX (Boyacioğlu et al., 2015), have not been tested on this data and may be considered in the future exploration using this data. We performed our analysis using the algorithms implemented in SPM that are most commonly used while analysing the data in cognitive and social neurosciences. For instance to normalize the data, the unified segmentation was used. Alternative algorithms implemented in other software, such as AIR, ANIMAL, ART, Diffeomorphic Demons, FNIRT, IRTK, JRD-fluid, ROMEO, SICLE, SyN, DARTEL, or using surfaces (Coalson et al., 2018; Klein et al., 2009) might provide alternate ways to perform normalization. Future studies could use the data we make https://osf.io/ncm25/?view_only=1d15a45aa3444 available at a7caed90f3842d9f4e5 to examine the impact of these methods on the effect of MB in our dataset. An additional limitation of our study is that for the MB2 condition, we had 22 instead of 23 participants. Because t-values scale with the square route of the degrees of freedom, this means that we have underestimated the t-values for the MB2 by about 2%. The differences between MB2 and MB4 we report here are however larger than 2%, and we feel that our data therefore nevertheless supports

benefits of MB4 over MB2, and performing all analysis on N = 22 participants lead to similar results (See Supplementary Fig. 8). Finally, we report that the image grey-white CNR decreases as the TR becomes shorter (Supplementary Figs. 6 and 7). In this data where the volunteers were healthy adults and there was minimal subject movement, we did not encounter any issues with the realignment steps of pre-processing pipeline. However, in patient population where motion might be an issue, an additional loss of CNR may result in sub-optimal realignment, thereby affecting the final statistics. This concept can be explored using controlled head movements in the scanner and may be addressed in future studies. Strengths: While most of the previous studies looked the effect of MB on the summary statistics from the subject level analyses (Demetriou et al., 2018; Kiss et al., 2018; Sahib et al., 2016; Todd et al., 2016), the current study is one of the only 2 studies that look at the effect of MB acceleration on voxel-wise group-level task-related statistics (Boyacioğlu et al., 2015). While the subject-level statics are good indicators of the performance of MB, one should be careful while interpreting the findings coming from sequences that result in variable image CNR as well as have different degrees of freedom per sequence (Bhandari et al., 2019). While most previous studies looked at the effect of multiband on very simple sensory tasks and lenient contrasts, using very small sample sizes (median N = 10), our study used a task and sample size more representative of contemporary cognitive neuroscience studies to have the power to detect even moderate benefits of multiband and ensure that our results are reproducible and representative. That our group results are so similar at different MB and so similar to an independent study (Fig. 2) provides evidence for the robustness of our cognitive neuroscience task and results. Using the same participants on the same scanner at different multiband levels ensures that differences in multiband performance are not the result of differences across subject pools or scanners.

Finally, there may be additional considerations for a potential user. For example, the head coil must have a sufficient number of coil elements, with a minimum of 32 elements as used in this study. Commercial solutions require appropriate licensing that needs to be acquired. The size of the data increases linearly with the MB factor, requiring additional storage space and processing timings. Moreover, different noise sources may affect the data differently because of the reduced signal to-noise ratio (SNR) per time frame due to reduced longitudinal magnetization recovery (for a detailed analysis of this topic see J. E. Chen et al., 2019). For instance, the impact of head movement on respiratory artifacts is much more pronounced with short TRs than longer TRs. Therefore, with special populations where head movement is an issue, use of shorter TRs may not be ideal or would require implementation of special preprocessing steps (J. E. Chen et al., 2019).

In summary, we can thus recommend the use of MB4 on a Philips scanner when aiming to perform group-level analyses using cognitive block design fMRI tasks using voxel sizes in the range of cortical thickness (e.g. 2.7 mm isotropic). While results will not be dramatically changed by the use of multiband, our results suggest that MB will bring a moderate but significant benefit.

Declaration of competing interest

The authors report that M.W.A. Caan is shareholder of Nicolab Inc.

CRediT authorship contribution statement

Ritu Bhandari: Conceptualization, Funding acquisition, Formal analysis, Writing - original draft, Data curation, Writing - review & editing. Evgeniya Kirilina: Formal analysis. Matthan Caan: Formal analysis. Judith Suttrup: Data curation. Teresa De Sanctis: Data curation. Lorenzo De Angelis: Formal analysis. Christian Keysers: Conceptualization, Funding acquisition, Formal analysis, Writing - original draft, Data curation, Writing - review & editing. Valeria Gazzola: Conceptualization, Funding acquisition, Writing - original draft, Data curation, Formal analysis, Writing - review & editing.

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Appendix A. Supplementary data

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