

HHS Public Access

Author manuscript *Neuroimage*. Author manuscript; available in PMC 2021 August 09.

Published in final edited form as:

Neuroimage. 2021 June ; 233: 117914. doi:10.1016/j.neuroimage.2021.117914.

ICA-based denoising strategies in breath-hold induced cerebrovascular reactivity mapping with multi echo BOLD fMRI

Stefano Moia^{a,b,*}, Maite Termenon^a, Eneko Uruñuela^{a,b}, Gang Chen^c, Rachael C. Stickland^d, Molly G. Bright^{d,e}, César Caballero-Gaudes^{a,*}

^aBasque Center on Cognition, Brain and Language, Donostia, Spain

^bUniversity of the Basque Country UPV/EHU, Donostia, Spain

°Scientific and Statistical Computing Core, NIMH/NIH/HHS, Bethesda, MD, United States

^dPhysical Therapy and Human Movement Sciences, Feinberg School of Medicine, Northwestern University, Chicago, IL, United States

^eBiomedical Engineering, McCormick School of Engineering, Northwestern University, Evanston, IL, United States

Abstract

Performing a BOLD functional MRI (fMRI) acquisition during breath-hold (BH) tasks is a non-invasive, robust method to estimate cerebrovascular reactivity (CVR). However, movement and breathing-related artefacts caused by the BH can substantially hinder CVR estimates due to their high temporal collinearity with the effect of interest, and attention has to be paid when choosing which analysis model should be applied to the data. In this study, we evaluate the performance of multiple analysis strategies based on lagged general linear models applied on multi-echo BOLD fMRI data, acquired in ten subjects performing a BH task during ten sessions, to obtain subject-specific CVR and haemodynamic lag estimates. The evaluated approaches range from conventional regression models, i.e. including drifts and motion timecourses as nuisance regressors, applied on single-echo or optimally-combined data, to more complex models including regressors obtained from multi-echo independent component analysis with different grades of orthogonalization in order to preserve the effect of interest, i.e. the CVR. We compare these models in terms of their ability to make signal intensity changes independent from motion, as well as the reliability as measured by voxelwise intraclass correlation coefficients of both CVR and lag maps over time. Our results reveal that a conservative independent component analysis model applied on the optimally-combined multi-echo fMRI signal offers the largest reduction of motion-related effects in the signal, while yielding reliable CVR amplitude and

Supplementary materials

This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/) *Corresponding authors. s.moia@bcbl.eu (S. Moia), c.caballero@bcbl.eu (C. Caballero-Gaudes). CRediT

Stefano Moia: Conceptualisation, Methodology, Software, Formal Analysis, Investigation, Data Curation, Writing (OD), Writing (RE), Visualisation, Funding acquisition; Maite Termenon: Methodology, Supervision, Writing (RE); Eneko Uruñuela: Investigation, Writing (RE); Rachael C. Stickland: Methodology, Writing (RE); Gang Chen: Methodology, Formal Analysis, Writing (RE); Molly G. Bright: Methodology, Supervision, Resources, Writing (RE); César Caballero-Gaudes: Conceptualisation, Methodology, Investigation, Supervision, Resources, Writing (RE), Funding acquisition.

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2021.117914.

lag estimates, although a conventional regression model applied on the optimally-combined data results in similar estimates. This work demonstrates the usefulness of multi-echo based fMRI acquisitions and independent component analysis denoising for precision mapping of CVR in single subjects based on BH paradigms, fostering its potential as a clinically-viable neuroimaging tool for individual patients. It also proves that the way in which data-driven regressors should be incorporated in the analysis model is not straight-forward due to their complex interaction with the BH-induced BOLD response.

Keywords

Cerebrovascular reactivity; Breath-hold; Multi-echo fMRI; Independent component analysis; Denoising; Precision functional mapping

1. Introduction

Cerebrovascular reactivity (CVR) is a physiological response of the cerebral vessels to vasodilatory or vasoconstrictive stimuli. Mapping of the CVR response provides an important indicator of cerebrovascular health. In recent years, functional magnetic resonance imaging (fMRI), either based on the blood oxygenation level-dependant (BOLD) contrast, arterial spin labelling, or a mixture of both, has demonstrated its effectiveness to assess CVR. As a result, its use is spreading into clinical practice, where its potential as a diagnostic measure is being ascertained in different diseases, spanning from vascular diseases (Hartkamp et al., 2017; Markus and Cullinane, 2001; Webster et al., 1995; Ziyeh et al., 2005), to stroke and aphasia (Krainik et al., 2005; Van Oers et al., 2018), brain tumors (Fierstra et al., 2018; Zacà et al., 2014), neurodegenerative diseases (Camargo et al., 2015; Glodzik et al., 2013; Marshall et al., 2014), hypertension (Iadecola and Davisson, 2008; Leoni et al., 2011; Tchistiakova et al., 2014), lifestyle habits (Friedman et al., 2008; Gonzales et al., 2014), sleep apnoea (Buterbaugh et al., 2015; Prilipko et al., 2014), and traumatic brain injury or concussions (Churchill et al., 2020; Markus and Cullinane, 2001).

CVR measurements are obtained by evoking a vasodilatory response during imaging. This is typically done by injecting intravenous acetazolamide, or by exposing the subject to gas challenges with computerised dynamic deployment of CO_2 and O_2 (Liu et al., 2018). However, acetazolamide is an invasive technique not indicated for vulnerable subjects (e.g. elderly or children), while gas challenges require dedicated setups and can also cause discomfort in some subjects, which might potentially bias CVR measurement (Urback et al., 2017). Alternatively, CO_2 changes in the blood due to breathing tasks, such as paced deep breathing or breath-hold (BH) tasks (Bright et al., 2009; Kastrup et al., 1998; Pinto et al., 2021), can elicit a CVR response that is equivalent to that of inhaled CO_2 (Kastrup et al., 2001; Tancredi and Hoge, 2013). A BH task can be successfully implemented in young children and elderly subjects (Handwerker et al., 2007; Thomason et al., 2005), and it is a robust measurement even if subjects are not able to hold their breath for as long as instructed (Bright and Murphy, 2013a). Moreover, BH-induced CVR is reliable across different sessions, both in the short (same day) and long term (Peng et al., 2019), in terms of spatial reliability (i.e. comparing variability of voxels across multiple sessions in one

subject) and general reliability (i.e. average CVR value across sessions and within subjects) (Lipp et al., 2015; Magon et al., 2009). Both short and long term reliability of BH-induced CVR were found to be comparable to that of other non-invasive means of estimating CVR, such as resting state fMRI (Liu et al., 2017), inhaled gas challenges (Dengel et al., 2017; Evanoff et al., 2020; Leung et al., 2016), Fourier modelling of a BH task (Pinto et al., 2016), and a paced deep breathing task (Sousa et al., 2014).

However, BOLD fMRI data exhibit signal variation arising from different sources, most of which corresponds to hardware-related artefacts and drifts, head motion, confounding physiological fluctuations, and other sources of noise (Bianciardi et al., 2009; Jorge et al., 2013). It is important that the signal variance associated with these confounding signals is accounted for and minimized during preprocessing or data analyses (Caballero-Gaudes and Reynolds, 2017; Liu, 2016). Head motion is a particularly problematic source of noise for task-based fMRI experiments, mainly in block designs (Johnstone et al., 2006) and in particular experimental paradigms, such as in overt speech production (Barch et al., 1999; Soltysik and Hyde, 2006; Xu et al., 2014). This concern with task-induced movement artefacts extends to respiration tasks: the experimental design is similar to that of block designs, but the amount of motion associated with paced breathing, deep breaths, or "recovery" breaths following a BH task can be very prominent and concur with the pattern of the task. Moreover, respiration can perturb the B0 field due to the change of air in the lungs (Raj et al., 2001) and introduce aliasing artefacts or pseudo-movement effects in the signal (Gratton et al., 2020; Pais-Roldán et al., 2018; Power et al., 2019).

There are different ways to account for motion effects on task-based fMRI data analysis. For instance, such effects can be reduced during acquisition by implementing an event-related task paradigm (Birn et al., 1999, 2004). However, in a BH task the periods of apnoea are typically between 10 and 20 s in duration to achieve a robust and repro-ducible vasodilatory response (Bright and Murphy, 2013a; Magon et al., 2009), and are not readily adapted to a brief event-related design. The most straight-forward approach is then to include the realignment parameters, as well as their derivatives and non-linear expansions, in the analysis model to account for part of the motion-related variance of the signal (Friston et al., 1996). In addition, fMRI data decomposition, for example with Principal Component Analysis or Independent Component Analysis (ICA), can be used to identify and remove components that are mostly related to motion or other sources of noise (Behzadi et al., 2007; Griffanti et al., 2014; Muschelli et al., 2014; Pruim et al., 2015a, 2015b; Salimi-Khorshidi et al., 2011).

Alternatively, noise in fMRI can be reduced by using multi-echo (ME) acquisitions that sample the data at multiple successive echo times (TE). A weighted combination of the multiple echoes (Poser et al., 2006; Posse et al., 1999) can smear out random noise and enhance the sensitivity to the BOLD contrast. Compared with single-echo data, this optimal combination can improve the mapping of neuronal activity at 3T (Fernandez et al., 2017) and 7T (Puckett et al., 2018). Optimal combination of multiple echo volumes can also improve BH-induced CVR mapping sensitivity, specificity, repeatability and reliability (Cohen and Wang, 2019).

Furthermore, assuming a monoexponential decay model, the voxelwise fMRI signal (in terms of signal percentage change) can be disentangled into BOLD-related fluctuations that depend linearly on the echo time (TE), and non-BOLD fluctuations related to changes in the net magnetization (Kundu et al., 2012). This can be used for denoising purposes. For example, in a dual-echo acquisition with a sufficiently short first TE, the first echo signal mainly captures changes in the net magnetization. It is then possible to perform nuisance regression from the second echo signal acquired at a longer TE with appropriate BOLD contrast (Bright and Murphy, 2013b). Collecting more echoes opens up the possibility of applying ICA and classifying independent components into BOLD-related or noise, an approach known as multi-echo independent component analysis (ME-ICA) (Kundu et al., 2013, 2012, 2017). Compared to single-echo data denoising, ME-ICA can improve the mapping of task-induced activation (Amemiya et al., 2019; DuPre et al., 2016; Evans et al., 2015; Gonzalez-Castillo et al., 2016; Lombardo et al., 2016). It also outperforms singleecho ICA-based denoising of resting-state fMRI data (Dipasquale et al., 2017), which is particularly beneficial to obtain more reliable functional connectivity mapping in individual subjects (Lynch et al., 2020) and in brain regions with reduced signal-to-noise ratio, such as the basal forebrain (Markello et al., 2018). Furthermore, ME-ICA also enhances the deconvolution of neuronal-related signal changes (Caballero-Gaudes et al., 2019).

However, up to now, the operation of ME-ICA has not been evaluated thoroughly in experimental paradigms with unavoidable task-correlated artefacts. Under such scenarios, one open question is how to obtain the right trade-off between removing as much noise as possible while saving the signal of interest (Bright and Murphy, 2015; Griffanti et al., 2014). In this study, we acquire ME-fMRI data during a BH task in 10 subjects acquired weekly, i.e. adopting a similar framework to precision functional mapping experiments (Gordon et al., 2017; Greene et al., 2020; Laumann et al., 2015; Lynch et al., 2020; Lynch and Liston, 2020; Marek et al., 2018), and assess the efficiency of different nuisance regression models to remove artefacts that are highly correlated with the effect of interest, i.e. the CVR response. In particular, we compare traditional nuisance regression approaches, applied to single- or multi-echo data, and three different ME-ICA denoising approaches ranging from aggressive to conservative. For each denoising strategy, we assess the correlation of the cleaned signal with measures of motion, and evaluate the amplitude and lag of the CVR signal response in terms of their physiological interpretability and inter-session reliability.

2. Material and methods

2.1. Participants

Ten healthy subjects with no record of psychiatric or neurological disorders (5F, age range 24–40 y at the start of the study) underwent ten MRI sessions in a 3T Siemens PrismaFit scanner with a 64-channel head coil. Each session took place one week apart, on the same day of the week and at the same time of the day.

All participants had to meet several further requirements, i.e. being non-smokers and refrain from smoking for the whole duration of the experiment, and not suffering from respiratory or cardiac health issues. They were also instructed to refrain from consuming caffeinated

drinks for two hours before the session. Informed consent was obtained before each session, and the study was approved by the local ethics committee.

2.2. Data acquisition and MRI session

Within the MRI session, subjects performed a BH task while T2*-weighted ME-fMRI data was acquired with the simultaneous multislice (a.k.a. multiband, MB) gradient-echo planar imaging sequence provided by the Centre for Magnetic Resonance Research (CMRR, Minnesota) (Moeller et al., 2010; Setsompop et al., 2012) with the following parameters: 340 scans, TR = 1.5 s, TEs = 10.6/28.69/46.78/64.87/82.96 ms, flip angle = 70° , MB acceleration factor = 4, GRAPPA = 2 with gradient-echo reference scan, 52 slices with interleaved acquisition, Partial Fourier = 6/8, FoV = 211×211 mm², voxel size = 2.4×2.4 \times 3 mm³, Phase Encoding = AP, bandwidth = 2470 Hz/px, LeakBlock kernel reconstruction (Cauley et al., 2014) and SENSE coil combination (Sotiropoulos et al., 2013). Single-band reference (SBRef) images were also acquired for each TE. The BH task was preceded by 64 min of ME-fMRI scanning, consisting of three task-based and four 10-minute resting state acquisitions, which are not part of the current study. The BH task always followed a resting state run. A pair of spin echo echo planar images (EPI) with opposite phase-encoding (AP or PA) directions and identical volume layout (TR = 2920 ms, TE = 28.6 ms, flip angle $= 70^{\circ}$) were also acquired before each functional run in order to estimate field distortions, similarly to the Human Connectome Project protocol (Glasser et al., 2016). A T1-weighted MP2RAGE image (Marques et al., 2009) (TR = 5 s, TE = 2.98 ms, TI1 = 700 ms, TI2 = 2.5 s, flip angle $1 = 4^{\circ}$, flip angle $2 = 5^{\circ}$, GRAPPA = 3, 176 slices, FoV read = 256 mm, voxel size = $1 \times 1 \times 1$ mm³, TA = 662 s) and a T2-weighted Turbo Spin Echo image (Hennig et al., 1986) (TR = 3.39 s, TE = 389 ms, GRAPPA = 2, 176 slices, FoV read = 256 mm, voxel size $= 1 \times 1 \times 1 \text{ mm}^3$, TA = 300 s) were also collected at the end and at the beginning of each MRI session, respectively.

During the fMRI acquisition runs exhaled CO_2 and O_2 levels were monitored and recorded using a nasal cannula (Intersurgical) with an ADInstruments ML206 gas analyser unit and transferred to a BIOPAC MP150 physiological monitoring system where scan triggers were simultaneously recorded. Photoplethismography and respiration effort data were also measured via the BIOPAC system, but these physiological signals were not used in the current study. All signals were sampled at 10 kHz. The physiological recordings started before and lasted longer than the ME-fMRI data recording to enable the shifting of physiological regressors.

2.3. Breath-hold task

Following Bright and Murphy (2013a), the BH paradigm consisted of eight repetitions of a BH trial composed of four paced breathing cycles of 6 s each, an apnoea (BH) of 20 s, an exhalation of 3 s, and 11 s of "recovery" breathing (unpaced) (i.e. total trial duration of 58 s) (Fig. 1). The BH paradigm was padded with a 15 s resting period to ensure that shifted physiological regressors would always match the BH paradigm period. Subjects were instructed prior to scanning about the importance of the exhalations preceding and following the apnoea (Pinto et al., 2021). Without these exhalations providing CO_2 measurements, the change in systemic CO_2 levels achieved by each BH cannot be robustly estimated; as a

result, CVR (%BOLD/mmHg CO₂ change) cannot be estimated quantitatively. Participants were instructed textually throughout the task through a mirror screen located in the head coil.

2.4. MRI data preprocessing

The DICOM files of the MRI data were transformed into nifti files with dcm2nii (Li et al., 2016) and formatted into Brain Imaging Data Structure (Gorgolewski et al., 2016) with heudiconv (Halchenko et al., 2019).

MRI data were minimally preprocessed with custom scripts based mainly in FSL (Jenkinson et al., 2012), AFNI (Cox, 1996), and ANTs (Tustison et al., 2014). In brief, the T2-weighted image was skull-stripped and co-registered to the MP2RAGE image along with the brain mask. The latter was applied to the MP2RAGE image, that was then segmented into grey matter (GM), white matter (WM) and cerebrospinal fluid tissues (Avants et al., 2011). The MP2RAGE image was normalised to an asymmetric version of the MNI152 6th generation template at 1 mm resolution (Grabner et al., 2006), while the T2-weighted volume was co-registered to the skull-stripped single-band reference image (SBRef) of the first echo. The first 10 volumes of the functional data were discarded to allow the signal to achieve a steady state of magnetisation. Image realignment to the SBRef was computed on the first echo, and the estimated rigid-body spatial transformation was then applied to all other echoes (Jenkinson et al., 2002; Jenkinson and Smith, 2001). A brain mask obtained from the SBRef volume was applied to all the echoes. The different echo timeseries were optimally combined (OC) voxelwise by weighting each timeseries contribution by its T_2^* value (Posse et al., 1999). Next, ME-ICA decomposition was performed on each run independently with tedana (DuPre et al., 2019) using the minimum description length criterion for estimation of the number of components (Harris, 1978; Li et al., 2016). The independent components (ICs) were then manually classified by SM and CCG into two categories (rejected or

accepted components) based on temporal, spatial, spectral and TE-dependence features of each component (Griffanti et al., 2017). The manual classifications are available in the data repository. A distortion field correction was performed on the OC volume with Topup (Andersson et al., 2003), using the pair of spin-echo EPI images with reversed phase encoding acquired before the ME-EPI acquisition (Glasser et al., 2016). Finally, the BOLD timeseries was converted in signal percentage change. For comparison, the dataset acquired at the second echo time (TE₂ = 28.6 ms) was used as a surrogate for standard single-echo (SE) acquisitions. This volume followed the same preprocessing steps as the OC volume, except for the optimal combination and the ICA decomposition.

2.5. CO₂ trace processing and CVR estimation

The files exported from the AcqKnowledge software were transformed and formatted into BIDS with phys2bids (The phys2bids developers et al., 2019).

The CO_2 timecourse was processed using custom scripts in Python 3.6.7. Briefly, the CO_2 timecourse was downsampled to 40 Hz to reduce computational costs. The end-tidal peaks were automatically and manually individuated. The amplitude envelope was obtained by linearly interpolating between the end-tidal peaks and it was then demeaned and convolved

with a canonical HRF to obtain the $P_{ET}CO_2hrf$ trace. In order to account for measurement delay, the $P_{ET}CO_2hrf$ trace was shifted to maximise the cross-correlation with the average timecourse of an eroded version of the GM mask (bulk shift) (Yezhuvath et al., 2009). This step was performed on both OC and the SE data (see Supplementary figure 1).

A lagged general linear model (GLM) approach was adopted in this study for CVR estimation (Moia, Stickland, et al., 2020) in order to model temporal offsets between the $P_{ET}CO_2$ recording and the CVR response across voxels that occur due to measurement and physiological delays (Donahue et al., 2016; Geranmayeh et al., 2015; Murphy et al., 2011; Sousa et al., 2014; Tong et al., 2011). Sixty shifted versions of the $P_{ET}CO_2hrf$ trace were created, ranging between ±9 s from the bulk shift, with a shift increment of 0.3 s (fine shift). This temporal range was based on previous literature, which rarely reports haemodynamic lags over ±8 s in healthy individuals (Bright et al., 2009; Donahue et al., 2016; Sousa et al., 2014). For each shift, a lagged GLM was defined with a design matrix comprised of the shifted $P_{ET}CO_2hrf$ timecourse as the regressor of interest, and different combinations of nuisance regressors (see below) in order to examine their efficiency in modelling artefactual signals of the voxel timeseries that might degrade CVR estimates. The simultaneous fitting of the nuisance regressors and the regressor of interest (i.e. the shifted $P_{ET}CO_2hrf$ trace) is preferable, rather than denoising via nuisance regression prior to the analysis (Jo et al., 2013; Lindquist et al., 2019; Moia et al., 2020).

Five different modelling strategies were evaluated, varying which nuisance regressors were included in the design matrix or how they were derived from ME-ICA:

 A lagged GLM model on the SE data where the design matrix includes the motion parameters and their temporal derivatives (denoted as *Mot*), Legendre polynomials of up to the fourth order (denoted as *Poly*), together with the P_{ET}CO₂*hrf* trace (SE-MPR):

$$Y_{SE} = P_{ET}CO_2hrf + Mot + Poly + n \tag{1}$$

2. The same model applied on the OC data (OC-MPR):

$$Y_{OC} = P_{ET}CO_2hrf + Mot + Poly + n \tag{2}$$

3. An *aggressive* model applied on the OC data (ME-AGG) in which the design matrix also includes the timecourses of the ME-ICA rejected components (denoted as IC_{rej}) added to the design matrix of the lagged GLM, orthogonalised with respect to the motion parameters, their temporal derivatives, and Legendre polynomials of up to the fourth order. This orthogonalisation step was performed to maintain a low Variance Inflation Factor in this model, and thus not bias the CVR estimation, without altering the relative variance explained by the original nuisance regressors and the regressor of interest (Mumford et al., 2015):

$$Y_{OC} = P_{ET}CO_2hrf + Mot + Poly + \left[IC_{rej} \perp (Mot, Poly)\right] + n$$
(3)

4. A moderate model applied on the OC data (ME-MOD) in which the timecourses of the ME-ICA rejected components are also orthogonalised with respect to the P_{ET}CO₂*hrf* trace (i.e. the regressor of interest describing the CVR response):

$$Y_{OC} = P_{ET}CO_2hrf + Mot + Poly + \left| IC_{rej} \perp (P_{ET}CO_2hrf, Mot, Poly) \right| + n$$
(4)

5. A conservative model applied on the OC data (ME-CON) in which the timeseries of the ME-ICA rejected components are orthogonalised with respect to the PETCO2hrf trace and the ME-ICA accepted components (denoted as ICacc):

$$Y_{OC} = P_{ET}CO_2hrf + Mot + Poly + [IC_{rej} \perp (P_{ET}CO_2hrf, IC_{acc}, Mot, Poly)] + n$$
⁽⁵⁾

In the models above, Y_{SE} and Y_{OC} are the SE and OC voxel timeseries respectively and n denotes the random noise.

For each modelling strategy and each of the sixty shifted PETCO2hrf traces, the corresponding lagged GLM was fitted via orthogonal least squares using AFNI. Then, for each voxel, the beta coefficient (i.e. weight) of the best fine-shifted P_{FT}CO₂hrf trace, corresponding to the lagged GLM model with maximum coefficient of determination (\mathbb{R}^2) , was selected. Finally, the beta coefficients expressed in BOLD signal percentage change over Volts (BOLD_{SPC}/V) were rescaled to be expressed in BOLD percentage over millimetres of mercury (%BOLD/mmHg) as indicated by the gas analyser manufacturer.¹

In this way, a lag-optimised CVR map and a t-value map were obtained, together with the associated lag map representing the voxelwise delay from the bulk shift, for each analysis pipeline. To account for sixty comparisons computed in the lagged GLM approach (one per regressor), the CVR and lag maps were thresholded at p < 0.05 adjusted with the Šidák correction (Bright et al., 2017; Šidák, 1967), and the voxels that were not statistically significant were excluded. The maps were further thresholded on the basis of the lag: those voxels in which the optimal lag was at or adjacent to the boundary (i.e. |lag| = 8.7 s) were considered not truly optimised and not readily physiologically plausible in healthy subjects and therefore masked in all maps (Moia et al., 2020).

2.6. Evaluation of motion removal across denoising strategies

For each type of lagged GLM analysis, 4-D volumes representing the modelled noise variance were reconstructed by multiplying the optimised beta coefficient maps of the nuisance regressors by their time-series using 3dSynthesize in AFNI. Then, they were subtracted from the OC or the SE data to obtain five different denoised datasets. DVARS, the root of the spatial mean square of the first derivative of the signal (Smyser et al., 2010), was computed on each denoised dataset as:

¹https://www.adinstruments.com/support/knowledge-base/it-possible-measure-expired-gasses-partial-pressure-mmhg-rather-

percentage. The adopted formula was $CO_2[mmHg] = (P_{atm} - P_{Vap})[mmHg] \cdot 10 \cdot CO_2[V]/100[V]$, where $CO_2[V]$ is the original CO₂ timeseries, P_{atm} is the atmospheric pressure in the laboratory at the moment of acquisition, and P_{Vap} is the water vapour pressure associated with expired air. The values of $P_{atm} = 759$ and $P_{Vap} = 47$ were used for all the sessions.

Neuroimage. Author manuscript; available in PMC 2021 August 09.

$$DVARS_{t} = \sqrt{\left\langle \left[I_{t}(x) - I_{t-1}(x)\right]^{2}\right\rangle},\tag{6}$$

where $I_t(x)$ is the image intensity of voxel x and at time t and $\langle \rangle$ indicates the spatial average over the whole brain. These DVARS timeseries were compared with the Framewise Displacement (FD) time courses (Power et al., 2012), computed using the realignment parameters estimated during preprocessing using the fsl_motion_outliers tool as:

$$FD_t = |\Delta d_x| + |\Delta d_y| + |\Delta d_z| + |\Delta \alpha| + |\Delta \beta| + |\Delta \gamma|, \tag{7}$$

where *t* denotes the time, d_x , d_y , d_z are the translational displacements along the three axes, α , β , γ are the rotational displacements of pitch, yaw, and roll, and $d_x = d_{x,t-1} - d_{x,t}$ (and similarly for the other parameters). DVARS was also computed on the SE volume before preprocessing (SE-PRE) to serve as a reference, as its relationship with FD should be at its maximum prior to the effects of motion being removed.

In order to test the moderating effect of each analysis on the relationship between DVARS and FD, a Linear Mixed Effects (LME) model was set up using the lme4 and lmer packages (Bates et al., 2015; Kuznetsova et al., 2017) in R (R Core Team, 2020), computing the p value with Satterthwaite's method (Satterthwaite, 1946), and accounting for the random effect of subject and session. The model was formulated as following in R equation notation:

$$DVARS \sim FD * model + (1|subject) + (1|session)$$
 (8)

Then, the same model was also used to assess differences in motion removal between pairs of denoising strategies. The results were thresholded at p = 0.05 corrected with the Šidák correction (Šidák, 1967).

In order to visualise the CVR responses to a BH trial, the average timeseries within GM was extracted from each denoised dataset from each model SE-MPR, OC-MPR, ME-AGG, ME-MOD, ME-CON, as well as from SE-PRE. These timeseries were transformed to BOLD percentage signal change, then the response to individual BH trials from each session were extracted using the timing of the third paced breathing cycle as a reference onset, and averaged together for each subject. The DVARS and FD timeseries followed the same process, except that the FD time-series were not expressed in percentage.

Finally, the amount of BH trials necessary to achieve a robust estimation of the BH response was computed for each denoising approach. The Manhattan distance from a pool of a gradually increasing number of trials to the average BOLD response over all BH trials (across the ten sessions, 80 trials in total) was also computed for each analysis model and subject.

2.7. Comparison of CVR and lag estimation and reliability across denoising strategies

For each denoising strategy, the average CVR and lag values across the significant voxels in GM and WM was computed for all subjects and all sessions, in addition to the amount of statistically significant voxels in the thresholded CVR maps.

In order to compare the results of the different denoising strategies, the thresholded CVR and lag maps of each session were normalised with a nearest neighbour interpolation to the MNI152 template (Grabner et al., 2006). Then, a LME model was computed voxelwise using 3dLMEr (Chen et al., 2013), considering the effect of subjects and sessions as random effects. The model was formulated as following in R equation notation:

$$X \sim model + (1|subject) + (1|session) \tag{9}$$

where X represents either the CVR or the lag value of each voxel. The same model was used to perform pairwise comparisons between the different strategies.

After normalising the t-value maps, the normalised CVR and lag maps were used to compute the intraclass correlation coefficient (ICC). ICC was computed voxelwise using a regularized multilevel mixed effect model in 3dICC (AFNI) in order to take into account the standard error of CVR and lag for each session in the ICC estimation (Chen et al., 2018). ICC assesses the reliability of a metric by comparing the intersubject, intrasubject, and total variability of that metric, which is equivalent to:

$$ICC(2,1) \simeq \widehat{\rho_2} = \frac{MS_{subj} - MS_n}{\frac{k}{n}(MS_{sess} - MS_n) + MS_{subj} + (k-1)MS_n}$$
(10)

where MS_{subj} , MS_{sess} , and MS_n are the mean squares of the effects of subjects, sessions, and residuals respectively, k = 10 is the number of sessions, and n = 7 the number of subjects (Chen et al., 2018; Mcgraw and Wong, 1996; Shrout and Fleiss, 1979). ICC(2,1) was chosen since both subjects and sessions were considered random effects. High ICC scores indicate high reliability, where the intrasubject variability is lower than the intersubject variability. Note that, since 3dICC uses the t-statistic map associated with the estimation of the CVR, CVR and lag maps used in this computation were thresholded only on the basis of the lag and not on the basis of the t-statistic.

3. Results

Three subjects were excluded due poor performance of the BH task in part of the sessions, mainly due to inadequate execution of the exhalations preceding and following the apnoea that prevented accurate determination of the $P_{ET}CO_2hrf$ traces. These traces are shown in red in Fig. 2 that plots the PetCO2hrf trace for all subjects and sessions. Hence, only the seven subjects that had all ten session were used for subsequent analyses (4F, age 25–40y).

3.1. Evaluation of motion removal across denoising strategies

Fig. 3 a illustrates the relationship between FD and DVARS in the raw data (SE-PRE) and after removing the reconstructed noise of each analysis model from the SE or OC volume for a representative subject; each point represents a timepoint and each line represents the linear regression between both timeseries in one session. The corresponding figures for the remaining subjects are available as Supplementary Material (Supplementary figure 2). Fig. 3b shows the same plot considering all the subjects and sessions. The modulating effect of the denoising approaches on the relation between DVARS and FD was tested

with a LME model that was found to be significant (F(6,161,181) = 34,597, p < 0.001). To further investigate the significant differences between analysis strategies, Table 1 reports the results of the same LME model considering pairwise combinations of all of the denoising approaches. From both Fig. 3 and Table 1, it can be seen that the optimal combination (OC-MPR) of ME data reduces DVARS compared to single-echo (SE-MPR). Although a similar relationship is observed between DVARS and FD in both approaches, OC-MPR significantly reduces the impact of FD compared to SE-MPR (β =715.10, CI₉₅ [710.17, 720.04], p < 0.001). This relationship is even more mitigated in the moderate (ME-MOD) (β=145.40, CI₉₅ [141.92, 148.88], p<0.001) and conservative (ME-CON) (β=146.69, CI₉₅ [143.05, 150.33], p<0.001) denoising approaches, which show similar modulatory effects on it. Note that this similarity is common, but not the same for all the subjects; for instance, ME-MOD showed larger reduction of motion related effects than ME-CON for two subjects (subject 003 and 007), while the opposite pattern was clearly observed in two other subjects (subject 004 and 009), and there was no apparent difference in the remaining subjects (see Supplementary figure 2). Considering all subjects and all sessions together, the difference between the ME-MOD and ME-CON approaches is statistically not significant (β =1.29, CI₉₅ [-0.70, 3.27], p>0.5). Compared to OC-MPR, both ME-MOD and ME-CON reduce the impact of FD on DVARS (β=130.84, CI₉₅ [127.89, 133.79], p<0.001 and β=132.13, CI₉₅ [128.99, 135.27], p<0.001 respectively). The aggressive strategy (ME-AGG) is the most successful in reducing motion-related effects described by FD on DVARS of all approaches.

Fig. 4 a plots the average percentage DVARS (left column) and average GM percentage BOLD response (central column) of all the BH trials across all of the sessions of a representative subject. The FD trace features a clear peak right after the end of the apnoea (highlighted in grey), likely associated with large head movement caused by the recovery breaths following the apnoea period. The percentage DVARS curves of the SE-PRE, SE-MPR and OC-MPR denoised timeseries reflect this peak in FD, which is absent in the ME-ICA based denoising timeseries, indicating a strong influence of movement on the signal intensity changes. All DVARS curves present a peak at a later time (between timepoints 25 and 30) that, as DVARS is akin to the first derivative of the BOLD signal changes, may agree with the return to the baseline seen in the BOLD response. The percentage BOLD signal change curves feature a delayed peak compared to the FD trace, reflecting a delayed CVR response compared to instantaneous head movements associated with respiration. However, they also feature a modulation in the BOLD signal change in correspondence with the peak in the FD trace, with the exception of ME-MOD and ME-AGG. The flattened DVARS and BOLD responses seen for ME-AGG indicate that the inclusion of the ME-ICA rejected components substantially removes part of the true CVR response, compared with the OC-MPR time courses. The average percentage DVARS and percentage BOLD response of the other subjects can be found in the Supplementary Material (Supplementary figure 3).

Fig. 4 b plots the Manhattan distance between the average of *N* trials and the average of all 80 BH trials as *N* increases from 1 to 80. ME-AGG tends to be more similar to the total average compared to all the other timeseries. For most of the subjects, SE-MPR, OC-MPR and ME-MOD have a similar behaviour and need more trials than SE-PRE, ME-CON and ME-AGG to converge to the total average. Note that the convergence to the analysis-specific 'ground truth' BH response is not monotonic and fluctuates across trials of the same session

and across sessions, indicating that the convergence does not depend only on the number of BH trials, but also on their quality and possible physiological variability in the CVR response across trials and sessions.

3.2. Cerebrovascular reactivity and lag maps

Fig. 5 and 6 show CVR and lag maps respectively, for all analysis strategies and all sessions of a representative subject (subject 002). The CVR and lag maps of other subjects are available in the Supplementary Material (Supplementary figure 4 and 5). The CVR maps were masked to exclude the voxels that were not statistically significant or whose lag is at the boundary of the explored range and might not have been truly optimised or physiologically plausible. Across all subjects, SE-MPR features more spatial variation and speckled noise in CVR and lag estimates of voxels within the same brain region compared to ME approaches like OC-MPR or ME-CON. In general, the ME-AGG and ME-MOD approaches do not yield CVR maps with as much clear distinction between brain tissues or delineation of the cortical folding and subcortical structures (e.g. see putamen and caudate nucleus) as obtained with the OC-MPR and ME-CON models. Amongst the ICA-based approaches, the adoption of an aggressive (ME-AGG) or moderate (ME-MOD) modelling strategy results in lag maps without anatomically defined patterns, as well as a higher rate of voxels with a lag estimation that is not within physiologically plausible range, and in CVR maps with lower responses and fewer significant voxels. ME-AGG also produces CVR maps with a higher percentage of negative values than any other analysis model, and a reduced CVR response in voxels near the posterior part of the superior sagittal and transverse sinuses.

Fig. 7 shows the distribution of the average values of CVR, lag, and the percentage of significant voxels for all subjects and sessions, and across all denoising strategies after thresholding. Considering the summaries within GM, although SE-MPR shows higher average CVR compared to the other approaches, it also features lower percentage of significant voxels compared to OC-MPR, ME-MOD and ME-CON. ME-AGG shows the lowest CVR value of all strategies, the most variable average of lag values, as well as the lowest percentage of significant voxels. ME-MOD features a lower percentage of significant voxels than SE-MPR, OC-MPR, and ME-CON. The same considerations can be extended to the WM. Table 2 reports the subject average CVR, lag, and the percentage of significant voxels across all denoising strategies after thresholding for GM only. The same table for WM can be found in the Supplementary Material (Supplementary table 1). For all models, the average CVR in the GM in the group and in each subject are comparable or higher than the BH-induced CVR (in%BOLD/mmHg) reported in previous literature (cfr. Bright et al., 2011; Bright and Murphy, 2013a; Lipp et al., 2015; Pinto et al., 2016).

3.3. Comparison of CVR and lag estimation and reliability across denoising strategies

Fig. 8 shows the results of comparing the CVR and lag maps across all of the denoising strategies. The top row shows the thresholded χ score of the contrast between SE-MPR and all other denoising strategies, while the other maps depict the pairwise comparison between all of the denoising strategies. Amongst the most interesting comparisons, all of the strategies based on ME have lower CVR and an anticipated response in areas vascularised

by big vessels (indicated by an arrow in the figure), where the blood transit time is usually faster compared to the rest of the brain. This could indicate that the response shown in SE-MPR could be overestimated due to the misestimation of its lag. Compared to SE-MPR, ME-MOD shows lower CVR and a delayed response in subcortical areas, while OC-MPR and ME-CON show higher CVR and an anticipated response in the insula, frontal, and parietal areas. OC-MPR shows no statistically significant differences with ME-CON, but a general higher CVR and an anticipated response compared to ME-MOD and ME-AGG, with the exception of the cerebellum, where it shows a delayed response. This difference could be related to the different local impact of motion artefacts, especially on the cerebellum. Between the three approaches based on ME-ICA, ME-AGG features generally lower CVR compared to the other two, and a generally anticipated response compared to ME-MOD and a delayed response to ME-CON.

In order to assess the reliability of each model, we also computed voxelwise ICC(2,1) maps for both CVR and haemodynamic lag. Fig. 9 depicts the ICC(2,1) maps for all analysis strategies for both CVR and lag maps, as well as their distributions. High ICC scores indicate that the intra-subject variability is lower than the inter-subject variability, hence the estimations of CVR or haemodynamic lag can be considered consistent across sessions. Conversely, low ICC scores indicate that the inter-subject variability is low compared to the intra-subject variability, hence the estimations of CVR and haemodynamic lag cannot be considered consistent across sessions. Following the classification given by (Cicchetti, 2001), an ICC score lower than 0.4 is considered poor, lower than 0.6 fair, lower than 0.75 good, and equal or higher than 0.75 excellent.

In terms of whole brain CVR reliability, the ME-CON demonstrated excellent reliability (spatial average across the whole brain of 0.86 ± 0.16) as well as the highest ICC values amongst all methods tested, closely followed by the OC-MPR (excellent, 0.85 ± 0.16), SE-MPR (excellent, 0.81 ± 0.19), and ME-MOD (excellent, 0.79 ± 0.19), while ME-AGG had a fair reliability (0.46 ± 0.22). If only voxels in GM are considered, the ICC of all approaches increases slightly $(0.88 \pm 0.14, 0.87 \pm 0.15, 0.85 \pm 0.17, 0.82 \pm 0.17, and 0.49 \pm 0.19)$ 0.22 for ME-CON, OC-MPR, SE-MPR, ME-MOD, and ME-AGG respectively). Despite the average fair reliability observed for ME-AGG, it can be observed that this approach exhibits a considerable number of voxels with poor reliability (ICC below 0.4). These voxels are mostly located in white matter, which also exhibit lower ICC values in the other analyses. In terms of whole-brain lag reliability, OC-MPR performed the best (good reliability, 0.67 \pm 0.21), closely followed by ME-CON (good reliability, 0.66 \pm 0.21). SE-MPR, ME-MOD, and ME-AGG demonstrated fair lag reliability (0.6 ± 0.22 and 0.42 ± 0.19 , 0.41 ± 0.20 , respectively). Considering only GM voxels, the reliability of all the approaches increases minimally $(0.68 \pm 0.21, 0.67 \pm 0.21, 0.61 \pm 0.21, 0.43 \pm 0.19, 0.42 \pm 0.20, \text{ for OC-MPR},$ ME-CON, SE-MPR, ME-MOD, and ME-AGG respectively). The reliability of CVR lag estimates was lower than that of CVR amplitude estimates, even though certain cortical regions, such as the visual and motor cortices, also show excellent ICC values for the OC-MPR and ME-CON denoising approaches. Interestingly, it can be observed that ME-MOD offers excellent ICC values for the CVR response amplitude in grey matter voxels, whereas they are poor for the lag estimates.

4. Discussion

In this study, we compared five different analysis strategies based on a lagged GLM model (Moia, Stickland, et al., 2020) to simultaneously remove motion-related effects and non-BOLD artefacts in the BOLD fMRI signal while estimating CVR and haemodynamic lag in order to identify the best modelling approach for BH paradigms in which prominent task-correlated artefacts coexist with the effect of interest. The lagged GLM model adopted in this study is similar to other models for CVR estimation that take into account local variations in the haemodynamic lag (Donahue et al., 2016; Geranmayeh et al., 2015; Murphy et al., 2011; Sousa et al., 2014; Tong et al., 2011). The main difference with such models is that, in this lagged GLM approach, after a first bulk shift that matches the average GM response with the PETCO2*hrf* regressor, the denoising and the voxelwise optimised response estimation take place simultaneously. This ensures that the interaction between regressors is properly taken into account and that the degrees of freedom of the model are properly estimated in the computation of the statistics. Amongst all possible modelling strategies, the five presented here were included in our analysis for different reasons. The optimal combination of ME fMRI data, with subsequent motion and Legendre polynomial regression (MPR), was expected to remove more noise and improve reliability of the CVR estimation due to its increased BOLD sensitivity compared to MPR on single-echo data, which is the standard approach for BH CVR estimation (Cohen and Wang, 2019). However, while optimal combination of ME volumes alone can partially reduce the noise present in the data, it still cannot remove motion artefacts, as illustrated in Fig. 3, in which SE-MPR and OC-MPR exhibit the same dependence of signal changes (DVARS) with motion (FD). For this reason, we further adopted three different ME-ICA based approaches, ranging from a conservative to an aggressive motion removal. ICA-based approaches are known to outperform traditional MPR in typical denoising fMRI data, possibly because they can identify and separate artefactual sources in the data in a data-driven and non-linear manner (Griffanti et al., 2014; Pruim et al., 2015a, 2015b; Salimi-Khorshidi et al., 2014). We did not apply ICA to single-echo data because it has already been demonstrated that ICA-based denoising applied to OC data outperforms ICA denoising applied to single-echo data (Dipasquale et al., 2017) and the ICs estimated from OC data might not have matched the ICs obtained from single-echo data, making such comparison less straightforward than the one based on MPR.

Spatial ICA decomposition is applied to fMRI data more often than temporal ICA decomposition, as the latter requires many more samples in time than normally available. Having many sessions for each subject, temporal ICA could have been leveraged in this study. In fact, temporal ICA could be more appropriate than spatial ICA to estimate a proper decomposition of timeseries sources (Smith et al., 2012), improving the modelling of temporal noise (Glasser et al., 2018), and potentially leading to better disentanglement of noise from CVR effects. However, we decided to apply spatial ICA in order to maintain the independence of each session, both to simulate a more common denoising approach to fMRI data, and to be able to capture session-specific noise contributions that could have been missed otherwise. Further studies could compare temporal and spatial ICA denoising for CVR mapping when many temporal samples have been collected in the same session,

for instance reducing the TR by acquiring fewer echoes. Here, our decision to acquire five echoes, instead of conventional multi-echo protocols with three or four echoes, was made to facilitate and improve the classification of the ICs based on their TE-dependence (Kundu et al., 2013).

The choice of comparing different levels of orthogonalisation of only the ICA-based nuisance regressors compared to regressors of interest might seem in contrast with previous literature, that suggests that orthogonalisation of collinear confounding factors could lead to misinterpreted results (Mumford et al., 2015). Our results clearly demonstrated that using the original (e.g., non orthogonalised) rejected ICs as nuisance regressors in the analysis (ME-AGG) removes the CVR effect of interest (see Figs. 4, 5, 6 and 9). To decide which regressors should be orthogonalised, and with respect to what, we considered the different origin of the nuisance regressors. While Legendre polynomials and motion parameters can be considered adequate models of noise sources in the data, intrinsic data-driven regressors may well contain variance related to the effect of interest, especially as spatial ICA was adopted and because of the high collinearity between the PETCO2hrf, motion, physiological adaptations to vascular dilation (e.g. cerebrospinal fluid flows), or changes in the magnetisation related to breathing (Raj et al., 2001). In these scenarios, it becomes more important to understand how to properly implement ICA denoising in order to preserve the effect of interest. For these reasons, three different ICA-based approaches were selected, from an aggressive strategy to a conservative approach, to assess if they preserved the BOLD effects related to the CVR response happening at different lags.

As hypothesized, all of the ME-based solutions outperformed the SE-MPR model in their ability to account for the effect of motion, summarized in terms of FD, on the fMRI signal intensity changes, described in terms of DVARS (see Fig. 3). Furthermore, all of the ICA-based strategies outperformed traditional MPR, and within ICA-based strategies, the aggressive one (ME-AGG) showed the best performance to remove these motionrelated effects in the signal. However, observing the average DVARS and BOLD response timecourses (Fig. 4) and the CVR and lag maps (Figs. 5 and 6) it becomes evident how the aggressive and moderate approaches result in lower estimates of CVR responses, even compared to the SE-MPR approach. Similarly, these two approaches result in the estimated haemodynamic lag hitting the boundaries of a physiologically plausible lag range in healthy adults. The substantial reduction in the CVR estimates in the aggressive approach (Figs. 4 and 5) occurs because the effect of interest can also be explained as a linear combination of the timecourses of rejected ICs related to motion, vascular effects or large susceptibility changes due to chest expansions and contractions while performing the BH task (Caballero-Gaudes and Reynolds, 2017; Griffanti et al., 2017). As for the moderate approach, the lower estimates of CVR could be due to the fact that orthogonalising data-driven nuisance regressors with respect to the PETCO2hrf trace per sé is not sufficient to save all the variance associated to real CVR. The $P_{ET}CO_2$ trace can only be estimated during exhalations, hence it is unable to capture local dynamic signal changes that are captured by ICs timeseries. Furthermore, CVR has a sigmoidal non-linear relation with the P_{FT}CO₂hrf trace (Bhogal et al., 2014), and the local BH-induced BOLD response has a complex shape, in terms of response amplitude and temporal delays, due to multiple physiological factors (Magon et al., 2009) that must be accounted for in order to improve its estimation. Our results illustrate

that these local complexities might be adequately captured by the linear combination of the accepted ICs timecourses, and not removing this variance from the rejected ICs when they are included as nuisance regressors in the model is detrimental (as observed with the ME-MOD and ME-AGG approaches). In other words, only a conservative approach (ME-CON) that preserves the BOLD variance associated with local CVR responses performs well, while also reducing motion-related effects more than conventional MPR models.

To further explore the benefit of different modelling strategies, we assessed the reliability of the resulting CVR and haemodynamic lag maps over the course of two and a half months (ten sessions) using ICC(2,1). To our knowledge, this was the first time that CVR reliability was tested over the course of ten sessions in individual subjects, and the first time that intersession haemodynamic lag reliability was tested. The ME-CON and OC-MPR strategies featured the greatest reliability for CVR and lag estimation, while the ME-AGG and ME-MOD approaches produced lower reliability values than even the simple SE-MPR model.

The lag maps are computed as the temporal offset related to the bulk shift, which is obtained by aligning the average GM BOLD response with the $P_{ET}CO_2hrf$ trace. If the bulk shift computation is misestimated this would create a systematic bias in the estimated lag maps, potentially reducing the apparent intersession reliability. While the CVR reliability should not be affected by this issue, due to the use of a lagged GLM approach that can overcome bulk shift misestimation (see session 4 of subject 007 in Supplementary figure 4 and 5), the true lag map reliability might be higher than reported here.

Regarding CVR reliability, the whole-brain average reliability of SE-MPR was comparable to long-term reliability (days or weeks apart) found in previous studies of CVR induced by BH (Peng et al., 2019), by paced deep breathing (Sousa et al., 2014), or by gas challenges (Leung et al., 2016), and higher than that reported in other studies on BH induced CVR estimated with a non-lagged optimized PETCO2hrf trace (Lipp et al., 2015) or with Fourier modelling (Pinto et al., 2016), and by gas challenges (Dengel et al., 2017; Evanoff et al., 2020). Consequently, the reliability of CVR estimates obtained with the optimal combination dataset and conservative ME-ICA modelling approaches were found higher than those previously reported in the literature. However, all strategies produced a reliability that was lower than the short-term (within-session) reliability reported in BH induced CVR (Peng et al., 2019), resting state based CVR (P. Liu et al., 2017), and gas challenge induced CVR (Leung et al., 2016), although lower intersession reliability in gas challenges has also been reported (Dengel et al., 2017; Evanoff et al., 2020). Note that the reliability observed in this study seems to be globally higher and spatially less variable than that reported in previous studies (Lipp et al., 2015; Sousa et al., 2014). However, discrepancies in the reliability measurements might be related to the different methods used to compute the CVR maps and the ICC score itself.

Using ICC to test reliability has the drawback that higher scores might be related to the presence of residual task-correlated motion effects that artificially stabilise the CVR estimation and reduce intra-subject variability compared to intersubject variability. In fact, recent studies have shown that individuals have particular movement patterns during fMRI

sessions that may be a stable characteristic of a person (Bolton et al., 2020) related to stable physical characteristics, such as body mass index (Ekhtiari et al., 2019) and could even be a heritable characteristic (Couvy-Duchesne et al., 2014; Hodgson et al., 2017). If subjects have similar motion patterns across the 10 repeated sessions, fMRI responses might appear more similar than they truly are, and the ICC might be inflated by such effects. Moreover, higher spatial reliability does not necessarily mean higher accuracy: a denoising strategy might be systematically misestimating CVR or haemodynamic lag. The fact that both optimal combination with traditional nuisance regression and the conservative ME-ICA denoising approaches resulted in similar CVR and lag spatial patterns and exhibited higher reliability than the single-echo model, while at the same time reduced the apparent effect of motion on the data variance, suggests that such drawbacks are mitigated in our data. However, further studies could compare different BH analysis strategies with a CVR estimation based on an independent computerised gas delivery protocol.

Another possibility would be to assess CVR in resting state fMRI, either measuring resting fluctuations in exhaled CO₂ levels (Golestani et al., 2016; Lipp et al., 2015), or by using a band of the power spectrum of the global signal as a regressor of interest (Liu et al., 2017, 2020). Such method might be more robust to motion collinearity, as the amount of movement in each breath is less pronounced and not consistently time-locked to the paradigm cues. At the same time, the lower amplitude of intrinsic CO_2 fluctuations relative to BH CO₂ change might also make this approach more susceptible to general motion effects and other sources of variance (e.g. neural or artefactual) unrelated to CO2. Moreover, previous work has shown that the optimal temporal shift between BOLD and P_{FT}CO₂ is hard to reliably identify in resting state data alone, in contrast to BH datasets where the temporal shift can be reliably identified (Bright et al., 2017; Stickland et al., 2021). Current resting state fMRI methods for CVR mapping may therefore be inappropriate to use with the lagged GLM approach that we have applied here. Either way, the analyses presented in this study can be easily implemented in other CVR assessment pipelines to mitigate the dependence of the response on motion. Beyond BH-based CVR studies, similar conclusions might be applicable to other experimental paradigms that present high collinearity between the expected task induced activity and artefactual sources, such as in overt speech production with long trial durations (Birn et al., 1999, 2004; Gracco et al., 2005), and that aim to use (ME-) ICA-based nuisance regressors as part of the model.

Note that MPR and ICA denoising are not the only viable options to reduce motion effects on fMRI and BH-induced CVR in particular: advanced setups can be used to reduce motion during the acquisition itself. For instance, subject specific moulded head casts can be used to reduce head motion (Power et al., 2019). Mounting an MRI compatible camera or tracker in the scanner enables prospective motion correction techniques (Faraji-Dana et al., 2016; Maziero et al., 2020; Parkes et al., 2018; Schulz et al., 2014; Zaitsev et al., 2017) or concurrent field monitoring enables the dynamic correction of field distortions dynamically (Vannesjo et al., 2015; Wilm et al., 2015) in order to effectively reduce effects of motion and magnetic field susceptibility changes. However, such advanced setups are not always available.

A limitation of this study is that the results are influenced by the manual classification of the ICA components performed by two of the authors. Despite being based on the automatic classification made by *tedana* (DuPre et al., 2019), we adopted a manual approach because often multiple ICs clearly exhibiting CVR-related timeseries and spatial maps were misclassified as noise. This manual classification was made with a cautious approach: if an IC seemed to be temporally and spatially related to the CVR response, it was accepted. Manual classification is still considered the gold standard for the classification of ICA components when performed by experts, despite the introduction of automatic classification algorithms (Griffanti et al., 2017), calling for further improvements in the automatic classification of (ME-)ICA components for BH tasks.

Another limitation is the lack of a CO_2 automated delivery protocol. The choice not to include one was driven by the necessity to reduce the discomfort of the participants during the imaging sessions, however further studies should compare denoised CVR maps to a CVR estimation based on independent computerised gas delivery protocols. This would also help estimating the accuracy of the denoised results on top of the reliability analysis featured in the present study.

Moreover, despite the fact that a BH task can be a valid alternative to gas delivery protocols for CVR estimation and its easy implementation, not all the subjects in this study could perform the task during all of the sessions. In total, 86% of the sessions were completed successfully by the subjects, although three subjects had to be excluded due to poor performance or non-compliance to the task in a subset of the sessions (four in two subjects and six in the third, see Fig. 2).

Finally, it is worth noticing that the adoption of ME imaging requires an increase in TR or a decrease in the spatial resolution. A way to cope for this issue is the adoption of simultaneous multislice (a.k.a. multi-band) acquisition, and despite the fact that this choice might introduce additional slice-leaking artefacts, a ME-ICA based denoising approach can successfully deal with their removal (Olafsson et al., 2015). Note that in this study we adopted one of the echo volumes as an approximation of a SE acquisition. Further studies could evaluate if this solution improves the estimation of CVR compared to SE imaging with higher spatial or temporal resolution.

Conclusion

Breath Holding (BH) is a non-invasive, robust way to estimate cerebrovascular reactivity (CVR). However, due to the task-correlated movement introduced by the BH task, attention has to be paid when choosing an appropriate modelling strategy to remove movement-related effects while preserving the effect of interest (P_{ET}CO₂). We compared different multi-echo (ME) independent component analysis (ICA) based denoising strategies to the standard data acquisition and analysis procedure, i.e. single-echo motion parameters regression. We found that a conservative ICA-based approach, but not an aggressive or moderate ICA approach, best removes motion-related effects while obtaining reliable CVR and lag responses, although a simple optimal combination of ME data with motion parameters regression provides similar CVR and lag estimations, and both ME-based approaches offer improvements in reliability compared with single-echo data acquisition.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

The authors would like to thank Vicente Ferrer for collaborating in data acquisition and two anonymous reviewers for helping improving the quality of the paper.

This research was supported by the European Union's Horizon 2020 research and innovation program (Marie Skłodowska-Curie grant agreement No. 713673), a fellowship from La Caixa Foundation (ID 100010434, fellowship code LCF/BQ/IN17/11620063), the Spanish Ministry of Economy and Competitiveness (Ramon y Cajal Fellowship, RYC-2017- 21845), the Spanish State Research Agency (BCBL "Severo Ochoa" excellence accreditation, SEV- 2015-490), the Basque Government (BERC 2018-2021 and PIBA_2019_104), the Spanish Ministry of Science, Innovation and Universities (MICINN; PID2019-105520GB-100 and FJCI-2017-31814), and the Eunice Kennedy Shriver National Institute of Child Health and Human Development of the National Institutes of Health under award number K12HD073945.

Data and code availability statement

In order to guarantee the replication of methods and results, all the code has been prepared to be run in a Singularity container based on a Ubuntu 18.04 Neurodebian OS. The container is publicly available at https://git.bcbl.eu/smoia/euskalibur_container, the methods pipeline is available at https://github.com/smoia/EuskalIBUR_dataproc, while all of the MRI images, physiological data, and manual classification used in this study is available in OpenNeuro (Moia et al., 2020).

References

- Amemiya S, Yamashita H, Takao H, Abe O, 2019. Integrated multi-echo denoising strategy improves identification of inherent language laterality. Magn. Reson. Med 81 (5), 3262–3271. doi: 10.1002/ mrm.27620. [PubMed: 30561807]
- Andersson JLR, Skare S, Ashburner J, 2003. How to correct susceptibility distortions in spin-echo echo-planar images: application to diffusion tensor imaging. Neuroimage 20 (2), 870–888. doi: 10.1016/S1053-8119(03)00336-7. [PubMed: 14568458]
- Avants BB, Tustison NJ, Wu J, Cook PA, Gee JC, 2011. An open source multivariate framework for N-tissue segmentation with evaluation on public data. Neuroinformatics 9 (4), 381–400. doi: 10.1007/s12021-011-9109-y. [PubMed: 21373993]
- Barch DMDM, Sabb FWFW, Carter CSCS, Braver TSTS, Noll DCDC, Cohen JDJD, 1999. Overt verbal responding during fMRI scanning: empirical investigations of problems and potential solutions. Neuroimage 10 (6), 642–657. [PubMed: 10600410]
- Bates D, Mächler M, Bolker BM, Walker SC, 2015. Fitting linear mixed-effects models using lme4. J. Stat. Softw 67 (1). doi: 10.18637/jss.v067.i01.
- Behzadi Y, Restom K, Liau J, Liu TT, 2007. A component based noise correction method (CompCor) for BOLD and perfusion based fMRI. Neuroimage 37 (1), 90–101. doi: 10.1016/j.neuroimage.2007.04.042. [PubMed: 17560126]
- Bhogal AA, Siero JCW, Fisher JA, Froeling M, Luijten P, Philippens M, Hoogduin H, 2014. Investigating the non-linearity of the BOLD cerebrovascular reactivity response to targeted hypo/ hypercapnia at 7T. Neuroimage 98, 296–305. doi: 10.1016/j.neuroimage.2014.05.006. [PubMed: 24830840]
- Bianciardi M, Fukunaga M, van Gelderen P, Horovitz SG, de Zwart JA, Shmueli K, Duyn JH, 2009. Sources of functional magnetic resonance imaging signal fluctuations in the human brain at rest: a 7 T study. Magn. Reson. Imaging 27 (8), 1019–1029. doi: 10.1016/j.mri.2009.02.004. [PubMed: 19375260]

- Birn RM, Bandettini PA, Cox RW, Shaker R, 1999. Event-related fMRI of tasks involving brief motion. Hum. Brain Mapp 7 (2), 106–114. doi: 10.1002/(SICI)1097-0193(1999)7:2<106::AID-HBM4>3.0.CO;2-O. [PubMed: 9950068]
- Birn RM, Cox RW, Bandettini PA, 2004. Experimental designs and processing strategies for fMRI studies involving overt verbal responses. Neuroimage 23 (3), 1046–1058. doi: 10.1016/ j.neuroimage.2004.07.039. [PubMed: 15528105]
- Bolton TAW, Kebets V, Glerean E, Zöller D, Li J, Yeo BTT, Caballero-Gaudes C, Van De Ville D, 2020. Agito ergo sum: correlates of spatio-temporal motion characteristics during fMRI. Neuroimage 209 (June 2019). doi: 10.1016/j.neuroimage.2019.116433.
- Bright MG, Bulte DP, Jezzard P, Duyn JH, 2009. Characterization of regional heterogeneity in cerebrovascular reactivity dynamics using novel hypocapnia task and BOLD fMRI. Neuroimage 48 (1), 166–175. doi: 10.1016/j.neuroimage.2009.05.026. [PubMed: 19450694]
- Bright MG, Donahue MJ, Duyn JH, Jezzard P, Bulte DP, 2011. The effect of basal vasodilation on hypercapnic and hypocapnic reactivity measured using magnetic resonance imaging. J. Cereb. Blood Flow Metab 31 (2), 426–438. doi: 10.1038/jcbfm.2010.187. [PubMed: 20959855]
- Bright MG, Murphy K, 2013a. Reliable quantification of BOLD fMRI cerebrovascular reactivity despite poor breath-hold performance. Neuroimage 83, 559–568. doi: 10.1016/ j.neuroimage.2013.07.007. [PubMed: 23845426]
- Bright MG, Murphy K, 2013b. Removing motion and physiological artifacts from intrinsic BOLD fluctuations using short echo data. Neuroimage 64 (1), 526–537. doi: 10.1016/ j.neuroimage.2012.09.043. [PubMed: 23006803]
- Bright MG, Murphy K, 2015. Is fMRI "noise" really noise? Resting state nuisance regressors remove variance with network structure. Neuroimage 114, 158–169. doi: 10.1016/ j.neuroimage.2015.03.070. [PubMed: 25862264]
- Bright MG, Tench CR, Murphy K, 2017. Potential pitfalls when denoising resting state fMRI data using nuisance regression. Neuroimage 154 (December 2016), 159–168. doi: 10.1016/j.neuroimage.2016.12.027. [PubMed: 28025128]
- Buterbaugh J, Wynstra C, Provencio N, Combs D, Gilbert M, Parthasarathy S, 2015. Cerebrovascular reactivity in Young subjects with sleep apnea. Sleep 38 (2), 241–250. doi: 10.5665/sleep.4406. [PubMed: 25409111]
- Caballero-Gaudes C, Moia S, Panwar P, Bandettini PA, Gonzalez-Castillo J, 2019. A deconvolution algorithm for multi-echo functional MRI: multi-echo Sparse paradigm free mapping. Neuroimage 202 (August), 1–15. doi: 10.1016/j.neuroimage.2019.116081.
- Caballero-Gaudes C, Reynolds RC, 2017. Methods for cleaning the BOLD fMRI signal. Neuroimage 154 (December 2016), 128–149. doi: 10.1016/j.neuroimage.2016.12.018. [PubMed: 27956209]
- Camargo CHF, Martins EA, Lange MC, Hoffmann HA, Luciano JJ, Young Blood MR, Schafranski MD, Ferro MM, Miyoshi E, 2015. Abnormal cerebrovascular reactivity in patients with Parkinson's disease. Parkinson's Dis. 2015. doi: 10.1155/2015/523041.
- Cauley SF, Polimeni JR, Bhat H, Wald LL, Setsompop K, 2014. Interslice leakage artifact reduction technique for simultaneous multislice acquisitions. Magn. Reson. Med 72 (1), 93–102. doi: 10.1002/mrm.24898. [PubMed: 23963964]
- Chen G, Saad ZS, Britton JC, Pine DS, Cox RW, 2013. Linear mixed-effects modeling approach to FMRI group analysis. Neuroimage 73, 176–190. doi: 10.1016/j.neuroimage.2013.01.047. [PubMed: 23376789]
- Chen G, Taylor PA, Haller SP, Kircanski K, Stoddard J, Pine DS, Leibenluft E, Brotman MA, Cox RW, 2018. Intraclass correlation: improved modeling approaches and applications for neuroimaging. Hum. Brain Mapp 39 (3), 1187–1206. doi: 10.1002/hbm.23909. [PubMed: 29218829]
- Churchill NW, Hutchison MG, Graham SJ, Schweizer TA, 2020. Cerebrovascular reactivity after sport concussion: from acute injury to 1 year after medical clearance. Front. Neurol 11 (July), 1–11. doi: 10.3389/fneur.2020.00558. [PubMed: 32116995]
- Cicchetti DV, 2001. The precision of reliability and validity estimates re-visited: distinguishing between clinical and statistical significance of sample size requirements. J. Clin. Exp. Neuropsychol 23 (5), 695–700. doi: 10.1076/jcen.23.5.695.1249. [PubMed: 11778646]

- Cohen AD, Wang Y, 2019. Improving the assessment of breath-holding induced cerebral vascular reactivity using a multiband multi-echo ASL/BOLD sequence. Sci. Rep 9 (1), 1–12. doi: 10.1038/ s41598-019-41199-w. [PubMed: 30626917]
- Couvy-Duchesne B, Blokland GAM, Hickie IB, Thompson PM, Martin NG, de Zubicaray GI, McMahon KL, Wright MJ, 2014. Heritability of head motion during resting state functional MRI in 462 healthy twins. Neuroimage 102 (P2), 424–434. doi: 10.1016/j.neuroimage.2014.08.010. [PubMed: 25132021]
- Cox RW, 1996. AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. Comput. Biomed. Res 29 (29), 162–173. doi: 10.1006/cbmr.1996.0014. [PubMed: 8812068]
- Dengel DR, Evanoff NG, Marlatt KL, Geijer JR, Mueller BA, Lim KO, 2017. Reproducibility of blood oxygen level-dependent signal changes with end-tidal carbon dioxide alterations. Clin. Physiol. Funct. Imaging 37 (6), 794–798. doi: 10.1111/cpf.12358. [PubMed: 26934185]
- Dipasquale O, Sethi A, Marcella Laganà M, Baglio F, Baselli G, Kundu P, Harrison NA, Cercignani M, 2017. Comparing resting state fMRI de-noising approaches using multi- and single-echo acquisitions. Clin. Imaging Sci. Centre, Brighton Sussex Med. School 3. doi: 10.1371/ journal.pone.0173289.
- Donahue MJ, Strother MK, Lindsey KP, Hocke LM, Tong Y, Frederick BDB, 2016. Time delay processing of hypercapnic fMRI allows quantitative parameterization of cerebrovascular reactivity and blood flow delays. J. Cereb. Blood Flow Metab 36 (10), 1767–1779. doi: 10.1177/0271678X15608643. [PubMed: 26661192]
- DuPre E, Luh WM, Spreng RN, 2016. Multi-echo fMRI replication sample of autobiographical memory, prospection and theory of mind reasoning tasks. Sci Data 3 (October), 1–9. doi: 10.1038/ sdata.2016.116.
- DuPre E, Salo T, Markello R, Kundu P, Whitaker K, & Handwerker D (2019). ME-ICA/tedana: 0.0.6. 10.5281/ZENODO.2558498.
- Ekhtiari H, Kuplicki R, Yeh H.wen, Paulus MP, 2019. Physical characteristics not psychological state or trait characteristics predict motion during resting state fMRI. Sci. Rep 9 (1), 1–8. doi: 10.1038/ s41598-018-36699-0. [PubMed: 30626917]
- Evanoff NG, Mueller BA, Marlatt KL, Geijer JR, Lim KO, Dengel DR, 2020. Reproducibility of a ramping protocol to measure cerebral vascular reactivity using functional magnetic resonance imaging. Clin. Physiol. Funct. Imaging 40 (3), 183–189. doi: 10.1111/cpf.12621. [PubMed: 31984617]
- Evans JW, Kundu P, Horovitz SG, Bandettini PA, 2015. Separating slow BOLD from non-BOLD baseline drifts using multi-echo fMRI. Neuroimage 105, 189–197. doi: 10.1016/ j.neuroimage.2014.10.051. [PubMed: 25449746]
- Faraji-Dana Z, Tam F, Chen JJ, Graham SJ, 2016. A robust method for suppressing motion-induced coil sensitivity variations during prospective correction of head motion in fMRI. Magn. Reson. Imaging 34 (8), 1206–1219. doi: 10.1016/j.mri.2016.06.005. [PubMed: 27451407]
- Fernandez B, Leuchs L, Sämann PG, Czisch M, Spoormaker VI, 2017. Multi-echo EPI of human fear conditioning reveals improved BOLD detection in ventromedial prefrontal cortex. Neuroimage 156 (May), 65–77. doi: 10.1016/j.neuroimage.2017.05.005. [PubMed: 28483719]
- Fierstra J, van Niftrik C, Piccirelli M, Bozinov O, Pangalu A, Krayenbühl N, Valavanis A, Weller M, Regli L, 2018. Diffuse gliomas exhibit whole brain impaired cerebrovascular reactivity. Magn. Reson. Imaging 45 (September 2017), 78–83. doi: 10.1016/j.mri.2017.09.017. [PubMed: 28986176]
- Friedman L, Turner JA, Stern H, Mathalon DH, Trondsen LC, Potkin SG, 2008. Chronic smoking and the BOLD response to a visual activation task and a breath hold task in patients with schizophrenia and healthy controls. Neuroimage 40 (3), 1181–1194. doi: 10.1016/j.neuroimage.2007.12.040. [PubMed: 18289881]
- Friston KJ, Williams S, Howard R, Frackowiak RSJ, Turner R, 1996. Movement-related effects in fMRI time-series. Magn. Reson. Med 35 (3), 346–355. doi: 10.1002/mrm.1910350312. [PubMed: 8699946]

- Geranmayeh F, Wise RJS, Leech R, Murphy K, 2015. Measuring vascular reactivity with breathholds after stroke: a method to aid interpretation of group-level BOLD signal changes in longitudinal fMRI studies. Hum. Brain Mapp 36 (5), 1755–1771. doi: 10.1002/hbm.22735. [PubMed: 25727648]
- Glasser MF, Coalson TS, Bijsterbosch JD, Harrison SJ, Harms MP, Anticevic A, Van Essen DC, Smith SM, 2018. Using temporal ICA to selectively remove global noise while preserving global signal in functional MRI data. Neuroimage 181 (December 2017), 692–717. doi: 10.1016/ j.neuroimage.2018.04.076. [PubMed: 29753843]
- Glasser MF, Smith SM, Marcus DS, Andersson JLRR, Auerbach EJ, Behrens TEJJ, Coalson TS, Harms MP, Jenkinson M, Moeller S, Robinson EC, Sotiropoulos SN, Xu J, Yacoub E, Ugurbil K, Van Essen DC, 2016. The human connectome project's neuroimaging approach. Nat. Neurosci 19 (9), 1175–1187. doi: 10.1038/nn.4361. [PubMed: 27571196]
- Glodzik L, Randall C, Rusinek H, de Leon MJ, 2013. Cerebrovascular reactivity to carbon dioxide in Alzheimer's disease. J. Alzheimer's Dis 35 (3), 427–440. doi: 10.3233/JAD-122011. [PubMed: 23478306]
- Golestani AM, Wei LL, Chen JJ, 2016. Quantitative mapping of cerebrovascular reactivity using resting-state BOLD fMRI: validation in healthy adults. Neuroimage 138, 147–163. doi: 10.1016/ j.neuroimage.2016.05.025. [PubMed: 27177763]
- Gonzales MM, Tarumi T, Mumford JA, Ellis RC, Hungate JR, Pyron M, Tanaka H, Haley AP, 2014. Greater BOLD response to working memory in endurance-trained adults revealed by breath-hold calibration. Hum. Brain Mapp 35 (7), 2898–2910. doi: 10.1002/hbm.22372. [PubMed: 24038949]
- Gonzalez-Castillo J, Panwar P, Buchanan LC, Caballero-Gaudes C, Handwerker DA, Jangraw DC, Zachariou V, Inati S, Roopchansingh V, Derbyshire JA, Bandettini PA, 2016. Evaluation of multiecho ICA denoising for task based fMRI studies: block designs, rapid event-related designs, and cardiac-gated fMRI. Neuroimage 141, 452–468. doi: 10.1016/j.neuroimage.2016.07.049. [PubMed: 27475290]
- Gordon EM, Laumann TO, Gilmore AW, Newbold DJ, Greene DJ, Berg JJ, Ortega M, Hoyt-Drazen C, Gratton C, Sun H, Hampton JM, Coalson RS, Nguyen AL, McDermott KB, Shimony JS, Snyder AZ, Schlaggar BL, Petersen SE, Nelson SM, Dosenbach NUF, 2017. Precision functional mapping of individual human brains. Neuron 95 (4), 791–807.e7. doi: 10.1016/j.neuron.2017.07.011. [PubMed: 28757305]
- Gorgolewski KJ, Auer T, Calhoun VD, Craddock RC, Das S, Duff EP, Flandin G, Ghosh SS, Glatard T, Halchenko YO, Handwerker DA, Hanke M, Keator D, Li X, Michael Z, Maumet C, Nichols BN, Nichols TE, Pellman J, Poldrack RA, 2016. The brain imaging data structure, a format for organizing and describing outputs of neuroimaging experiments. Sci. Data 3 (1). doi: 10.1038/ sdata.2016.44.
- Grabner G, Janke AL, Budge MM, Smith D, Pruessner J, Collins DL, 2006. Symmetric atlasing and model based segmentation: an application to the hippocampus in older adults. In: Larsen R, Nielsen M, Sporring J (Eds.), Medical Image Computing and Computer-Assisted Intervention MICCAI 2006: 9th International Conference, Copenhagen, Denmark, October 1–6, 2006. Proceedings, Part II. Springer Berlin Heidelberg, pp. 58–66. doi: 10.1007/11866763_8.
- Gracco VL, Tremblay P, Pike B, 2005. Imaging speech production using fMRI. Neuroimage 26 (1), 294–301. doi: 10.1016/j.neuroimage.2005.01.033. [PubMed: 15862230]
- Gratton C, Dworetsky A, Coalson RS, Adeyemo B, Laumann TO, Wig GS, Kong TS, Gratton G, Fabiani M, Barch DM, Tranel D, Miranda-Dominguez O, Fair DA, Dosenbach NUF, Snyder AZ, Perlmutter JS, Petersen SE, Camp-bell MC, 2020. Removal of high frequency contamination from motion estimates in single-band fMRI saves data without biasing functional connectivity. Neuroimage 217 (April), 116866. doi: 10.1016/j.neuroimage.2020.116866. [PubMed: 32325210]
- Greene DJ, Marek S, Gordon EM, Siegel JS, Gratton C, Laumann TO, Gilmore AW, Berg JJ, Nguyen AL, Dierker D, Van AN, Ortega M, New-bold DJ, Hampton JM, Nielsen AN, McDermott KB, Roland JL, Norris SA, Nelson SM, ... Dosenbach NUF, 2020. Integrative and network-specific connectivity of the Basal Ganglia and Thalamus defined in individuals. Neuron 105 (4), 742–758.e6. doi: 10.1016/j.neuron.2019.11.012. [PubMed: 31836321]
- Griffanti L, Douaud G, Bijsterbosch J, Evangelisti S, Alfaro-Almagro F, Glasser MF, Duff EP, Fitzgibbon S, Westphal R, Carone D, Beckmann CF, Smith SM, 2017. Hand classification

of fMRI ICA noise components. Neuroimage 154 (June 2016), 188–205. doi: 10.1016/ j.neuroimage.2016.12.036. [PubMed: 27989777]

- Griffanti L, Salimi-Khorshidi G, Beckmann CF, Auerbach EJ, Douaud G, Sexton CE, Zsoldos E, Ebmeier KP, Filippini N, Mackay CE, Moeller S, Xu J, Yacoub E, Baselli G, Ugurbil K, Miller KL, Smith SM, 2014. ICA-based artefact removal and accelerated fMRI acquisition for improved resting state network imaging. Neuroimage 95, 232–247. doi: 10.1016/j.neuroimage.2014.03.034. [PubMed: 24657355]
- Halchenko Y, Goncalves M, di Oleggio Castello MV, Ghosh S, Hanke M, Dae, Salo T, Kent J, Amlien I, Brett M, Tilley S, Markiewicz C, Gorgolewski C, pvelasco, Kim S, Stadler J, Kaczmarzyk J, Lukas DC, john lee, & Kahn A, 2019. nipy/heudiconv v0.6.0. Zenodo. 10.5281/zenodo.3579455.
- Handwerker DA, Gazzaley A, Inglis BA, D'Esposito M, 2007. Reducing vascular variability of fMRI data across aging populations using a breathholding task. Hum. Brain Mapp 28 (9), 846–859. doi: 10.1002/hbm.20307. [PubMed: 17094119]
- Harris FJ, 1978. On the use of windows for harmonic analysis with the discrete Fourier transform. Proc. IEEE 66 (1), 51–83. doi: 10.1109/PROC.1978.10837.
- Hartkamp NS, Bokkers RPH, van Osch MJP, de Borst GJ, Hendrikse J, 2017. Cerebrovascular reactivity in the caudate nucleus, lentiform nucleus and thalamus in patients with carotid artery disease. J. Neuroradiol 44 (2), 143–150. doi: 10.1016/j.neurad.2016.07.003. [PubMed: 27743788]
- Hennig J, Nauerth A, Friedburg H, 1986. RARE imaging: a fast imaging method for clinical MR. Magn. Reson. Med 3 (6), 823–833. doi: 10.1002/mrm.1910030602. [PubMed: 3821461]
- Hodgson K, Poldrack RA, Curran JE, Knowles EE, Mathias S, Göring HHH, Yao N, Olvera RL, Fox PT, Almasy L, Duggirala R, Barch DM, Blangero J, Glahn DC, 2017. Shared genetic factors influence head motion during MRI and body mass index. Cereb. Cortex (New York, N.Y. : 1991) 27 (12), 5539–5546. doi: 10.1093/cercor/bhw321.
- Iadecola C, Davisson RL, 2008. Hypertension and cerebrovascular dysfunction. Cell Metab. 7 (6), 476–484. doi: 10.1016/j.cmet.2008.03.010. [PubMed: 18522829]
- Jenkinson M, Bannister P, Brady M, Smith S, 2002. Improved optimization for the robust and accurate linear registration and motion correction of brain images. Neuroimage 17 (2), 825–841. doi: 10.1006/nimg.2002.1132. [PubMed: 12377157]
- Jenkinson M, Beckmann CF, Behrens TEJ, Woolrich MW, Smith SM, 2012. FSL. Neuroimage 62 (2), 782–790. doi: 10.1016/j.neuroimage.2011.09.015. [PubMed: 21979382]
- Jenkinson M, Smith S, 2001. A global optimisation method for robust affine registration of brain images. Med. Image Anal 5 (2), 143–156. doi: 10.1016/S1361-8415(01)00036-6. [PubMed: 11516708]
- Jo HJ, Gotts SJ, Reynolds RC, Bandettini PA, Martin A, Cox RW, & Saad ZS, 2013. Effective preprocessing procedures virtually eliminate distance-dependent motion artifacts in resting state fMRI. Journal of Applied Mathematics 2013. doi:10.1155/2013/935154. 10.1155/2013/935154.
- Johnstone T, Ores Walsh KS, Greischar LL, Alexander AL, Fox AS, Davidson RJ, Oakes TR, 2006. Motion correction and the use of motion covariates in multiple-subject fMRI analysis. Hum. Brain Mapp 27 (10), 779–788. doi: 10.1002/hbm.20219. [PubMed: 16456818]
- Jorge J, Figueiredo P, van der Zwaag W, Marques JP, 2013. Signal fluctuations in fMRI data acquired with 2D-EPI and 3D-EPI at 7 Tesla. Magn. Reson. Imaging 31 (2), 212–220. doi: 10.1016/ j.mri.2012.07.001. [PubMed: 22921734]
- Kastrup A, Krüger G, Neumann-Haefelin T, Moseley ME, 2001. Assessment of cerebrovascular reactivity with functional magnetic resonance imaging: comparison of CO2 and breath holding. Magn. Reson. Imaging 19 (1), 13–20. doi: 10.1016/S0730-725X(01)00227-2. [PubMed: 11295341]
- Kastrup A, Li T, Takahashi A, Glover GH, Moseley ME, 1998. Functional magnetic resonance imaging of regional cerebral blood oxygenation changes during breath holding. Stroke 29 (12), 2641–2645. doi: 10.1161/01.STR.29.12.2641. [PubMed: 9836778]
- Krainik A, Hund-Georgiadis M, Zysset S, Von Cramon DY, 2005. Regional impairment of cerebrovascular reactivity and BOLD signal in adults after stroke. Stroke 36 (6), 1146–1152. doi: 10.1161/01.STR.0000166178.40973.a7. [PubMed: 15879326]

- Kundu P, Brenowitz ND, Voon V, Worbe Y, Vertes PE, Inati SJ, Saad ZS, Bandettini PA, Bullmore ET, 2013. Integrated strategy for improving functional connectivity mapping using multiecho fMRI. Proc. Natl. Acad. Sci 110 (40), 16187–16192. doi: 10.1073/pnas.1301725110. [PubMed: 24038744]
- Kundu P, Inati SJ, Evans JW, Luh W–M, Bandettini PA, 2012. Differentiating BOLD and non-BOLD signals in fMRI time series using multi-echo EPI. Neuroimage 60, 1759–1770. doi: 10.1016/ j.neuroimage.2011.12.028. [PubMed: 22209809]
- Kundu P, Voon V, Balchandani P, Lombardo MV, Poser BA, Bandettini PA, 2017. Multi-echo fMRI: a review of applications in fMRI denoising and analysis of BOLD signals. Neuroimage 154 (March), 59–80. doi: 10.1016/j.neuroimage.2017.03.033. [PubMed: 28363836]
- Kuznetsova A, Brockhoff PB, Christensen RHB, 2017. ImerTest package: tests in linear mixed effects models. J. Stat. Softw 82 (13). doi: 10.18637/jss.v082.i13.
- Laumann TO, Gordon EM, Adeyemo B, Snyder AZ, Joo SJ, Chen MY, Gilmore AW, McDermott KB, Nelson SM, Dosenbach NUF, Schlaggar BL, Mumford JA, Poldrack RA, Petersen SE, 2015. Functional system and areal organization of a highly sampled individual human brain. Neuron 87 (3), 658–671. doi: 10.1016/j.neuron.2015.06.037.
- Leoni RF, Paiva FF, Henning EC, Nascimento GC, Tannús A, De Araujo DB, Silva AC, 2011. Magnetic resonance imaging quantification of regional cerebral blood flow and cerebrovascular reactivity to carbon dioxide in normotensive and hypertensive rats. Neuroimage 58 (1), 75–81. doi: 10.1016/j.neuroimage.2011.06.030. [PubMed: 21708273]
- Leung J, Kim JA, Kassner A, 2016. Reproducibility of cerebrovascular reactivity measures in children using BOLD MRI. J. Magn. Reson. Imaging 43 (5), 1191–1195. doi: 10.1002/jmri.25063. [PubMed: 26435493]
- Li X, Morgan PS, Ashburner J, Smith J, Rorden C, 2016. The first step for neuroimaging data analysis: DICOM to NIfTI conversion. J. Neurosci. Methods 264, 47–56. doi: 10.1016/ j.jneumeth.2016.03.001. [PubMed: 26945974]
- Lindquist MA, Geuter S, Wager TD, Caffo BS, 2019. Modular preprocessing pipelines can reintroduce artifacts into fMRI data. Hum. Brain Mapp 40 (8), 2358–2376. doi: 10.1002/hbm.24528. [PubMed: 30666750]
- Lipp I, Murphy K, Caseras X, Wise RG, 2015. Agreement and repeatability of vascular reactivity estimates based on a breath-hold task and a resting state scan. Neuroimage 113, 387–396. doi: 10.1016/j.neuroimage.2015.03.004. [PubMed: 25795342]
- Liu P, De Vis JB, Lu H, De Vis JB, Lu H, 2018. Cerebrovascular reactivity (CVR) MRI with CO2 challenge: a technical review. Neuroimage 187 (February), 104–115. doi: 10.1016/ j.neuroimage.2018.03.047. [PubMed: 29574034]
- Liu P, Li Y, Pinho M, Park DC, Welch BG, Lu H, 2017. Cerebrovascular reactivity mapping without gas challenges. Neuroimage 146 (November 2016), 320–326. doi: 10.1016/ j.neuroimage.2016.11.054. [PubMed: 27888058]
- Liu P, Xu C, Lin Z, Sur S, Li Y, Yasar S, Rosenberg P, Albert M, Lu H, 2020. Cerebrovascular reactivity mapping using intermittent breath modulation. Neuroimage 215 (March), 116787. doi: 10.1016/j.neuroimage.2020.116787. [PubMed: 32278094]
- Liu TT, 2016. Noise contributions to the fMRI signal: an overview. Neuroimage 143, 141–151. doi: 10.1016/j.neuroimage.2016.09.008. [PubMed: 27612646]
- Lombardo MV, Auyeung B, Holt RJ, Waldman J, Ruigrok ANV, Mooney N, Bullmore ET, Baron-Cohen S, Kundu P, 2016. Improving effect size estimation and statistical power with multi-echo fMRI and its impact on understanding the neural systems supporting mentalizing. Neuroimage 142, 55–66. doi: 10.1016/j.neuroimage.2016.07.022. [PubMed: 27417345]
- Lynch CJ, Liston C, 2020. Precision functional mapping of corticostriatal and corticothalamic circuits: parallel processing reconsidered. Neuron 105 (4), 595–597. doi: 10.1016/j.neuron.2020.01.025. [PubMed: 32078793]
- Lynch CJ, Power JD, Scult MA, Dubin M, Gunning FM, Liston C, 2020. Rapid precision functional mapping of individuals using multi-echo fMRI. Cell Rep. 33 (12), 108540. doi: 10.1016/ j.celrep.2020.108540. [PubMed: 33357444]

- Magon S, Basso G, Farace P, Ricciardi GK, Beltramello A, Sbarbati A, 2009. Reproducibility of BOLD signal change induced by breath holding. Neuroimage 45, 702–712. doi: 10.1016/ j.neuroimage.2008.12.059. [PubMed: 19211035]
- Marek S, Siegel JS, Gordon EM, Raut RV, Gratton C, Newbold DJ, Ortega M, Laumann TO, Adeyemo B, Miller DB, Zheng A, Lopez KC, Berg JJ, Coalson RS, Nguyen AL, Dierker D, Van AN, Hoyt CR, McDermott KB, Dosenbach NUF, 2018. Spatial and temporal organization of the individual human cerebellum. Neuron 100 (4), 977–993.e7. doi: 10.1016/j.neuron.2018.10.010. [PubMed: 30473014]
- Markello RD, Spreng RN, Luh WM, Anderson AK, De Rosa E, 2018. Segregation of the human basal forebrain using resting state functional MRI. Neuroimage 173 (October 2017), 287–297. doi: 10.1016/j.neuroimage.2018.02.042. [PubMed: 29496614]
- Markus H, Cullinane M, 2001. Severely impaired cerebrovascular reactivity predicts stroke and TIA risk in patients with carotid artery stenosis and occlusion. Brain 124 (3), 457–467. doi: 10.1093/ brain/124.3.457. [PubMed: 11222446]
- Marques JP, Kober T, Krueger G, Van Der Zwaag W, Van De Moortele P-F, Gruetter R, 2009. MP2RAGE, a self bias-field corrected sequence for improved segmentation and T 1 -mapping at high field. Neuroimage 49, 1271–1281. doi: 10.1016/j.neuroimage.2009.10.002. [PubMed: 19819338]
- Marshall O, Lu H, Brisset JC, Xu F, Liu P, Herbert J, Grossman RI, Ge Y, 2014. Impaired cerebrovascular reactivity in multiple sclerosis. JAMA Neurol. 71 (10), 1275–1281. doi: 10.1001/ jamaneurol.2014.1668. [PubMed: 25133874]
- Maziero D, Rondinoni C, Marins T, Stenger VA, Ernst T, 2020. Prospective motion correction of fMRI: improving the quality of resting state data affected by large head motion. Neuroimage 212 (August 2019), 116594. doi: 10.1016/j.neuroimage.2020.116594. [PubMed: 32044436]
- Mcgraw KO, Wong SP, 1996. Forming inferences about some intraclass correlation coefficients. Psychol. Methods 1 (1), 30–46. doi: 10.1037/1082-989X.1.1.30.
- Moeller S, Yacoub E, Olman CA, Auerbach E, Strupp J, Harel N, Uğurbil K, 2010. Multiband multislice GE-EPI at 7 tesla, with 16-fold acceleration using partial parallel imaging with application to high spatial and temporal whole-brain FMRI. Magn. Reson. Med 63 (5), 1144– 1153. doi: 10.1002/mrm.22361. [PubMed: 20432285]
- Moia S, Stickland RC, Ayyagari A, Termenon M, Caballero-Gaudes C, Bright MG, 2020a. Voxelwise optimization of hemodynamic lags to improve regional CVR estimates in breath-hold fMRI.
 In: 2020 42nd Annual International Conference of the IEEE Engineering in Medicine Biology Society (EMBC), pp. 1489–1492. doi: 10.1109/EMBC44109.2020.9176225.
- Moia S, Uruñuela E, Ferrer V, Caballero-Gaudes C, 2020b. EuskalIBUR. OpenNeuro doi: 10.18112/ openneuro.ds003192.v1.0.1.
- Mumford JA, Poline JB, Poldrack RA, 2015. Orthogonalization of regressors in fMRI models. PLoS ONE 10 (4), 1–11. doi: 10.1371/journal.pone.0126255.
- Murphy K, Harris AD, Wise RG, 2011. Robustly measuring vascular reactivity differences with breath-hold: normalising stimulus-evoked and resting state BOLD fMRI data. Neuroimage 54 (1), 369–379. doi: 10.1016/j.neuroimage.2010.07.059. [PubMed: 20682354]
- Muschelli J, Nebel MB, Caffo BS, Barber AD, Pekar JJ, Mostofsky SH, 2014. Reduction of motionrelated artifacts in resting state fMRI using aCompCor. Neuroimage 96, 22–35. doi: 10.1016/ j.neuroimage.2014.03.028. [PubMed: 24657780]
- Olafsson V, Kundu P, Wong EC, Bandettini PA, Liu TT, 2015. Enhanced identification of BOLD-like components with multi-echo simultaneous multi-slice (MESMS) fMRI and multi-echo ICA. Neuroimage 112, 43–51. doi: 10.1016/j.neuroimage.2015.02.052. [PubMed: 25743045]
- Pais-Roldán P, Biswal B, Scheffler K, Yu X, 2018. Identifying respiration-related aliasing artifacts in the rodent resting-state fMRI. Front. Neurosci 12 (NOV), 1–14. doi: 10.3389/fnins.2018.00788. [PubMed: 29403346]
- Parkes L, Fulcher B, Yücel M, Fornito A, 2018. An Evaluation of the Efficacy, Reliability, and Sensitivity of Motion Correction Strategies for Resting-State Functional MRI doi: 10.1016/ j.neuroimage.2017.12.073.

- Peng S–L, Yang H–C, Chen C–M, Shih C–T, 2019. Short- and long-term reproducibility of BOLD signal change induced by breath-holding at 1.5 and 3 T. NMR Biomed. doi: 10.1002/nbm.4195.
- Pinto J, Bright MG, Bulte DP, Figueiredo P, 2021. Cerebrovascular reactivity mapping without gas challenges: a methodological guide. Front. Physiol 11, 1711. doi: 10.3389/fphys.2020.608475, https://www.frontiersin.org/article/10.3389/fphys.2020.608475.
- Pinto J, Jorge J, Sousa I, Vilela P, Figueiredo P, 2016. Fourier modeling of the BOLD response to a breath-hold task: optimization and reproducibility. Neuroimage 135, 223–231. doi: 10.1016/ j.neuroimage.2016.02.037. [PubMed: 26908316]
- Poser BA, Versluis MJ, Hoogduin JM, Norris DG, 2006. BOLD contrast sensitivity enhancement and artifact reduction with multiecho EPI: parallel-acquired inhomogeneity-desensitized fMRI. Magn. Reson. Med 55 (6), 1227–1235. doi: 10.1002/mrm.20900. [PubMed: 16680688]
- Posse S, Wiese S, Gembris D, Mathiak K, Kessler C, Grosse-Ruyken ML, Elghahwagi B, Richards T, Dager SR, Kiselev VG, 1999. Enhancement of BOLD-contrast sensitivity by single-shot multi-echo functional MR imaging. Magn. Reson. Med 42 (1), 87–97. doi: 10.1002/ (SICI)1522-2594(199907)42:1<87::AID-MRM13>3.0.CO;2-O. [PubMed: 10398954]
- Power JD, Barnes KA, Snyder AZ, Schlaggar BL, Petersen SE, 2012. Spurious but systematic correlations in functional connectivity MRI networks arise from subject motion. Neuroimage 59 (3), 2142–2154. doi: 10.1016/j.neuroimage.2011.10.018. [PubMed: 22019881]
- Power JD, Lynch CJ, Silver BM, Dubin MJ, Martin A, Jones RM, 2019a. Distinctions among real and apparent respiratory motions in human fMRI data. Neuroimage 201 (July), 116041. doi: 10.1016/j.neuroimage.2019.116041. [PubMed: 31344484]
- Power JD, Silver BM, Silverman MR, Ajodan EL, Bos DJ, Jones RM, 2019b. Customized head molds reduce motion during resting state fMRI scans. Neuroimage 189 (October 2018), 141–149. doi: 10.1016/j.neuroimage.2019.01.016. [PubMed: 30639840]
- Prilipko O, Huynh N, Thomason ME, Kushida CA, Guilleminault C, 2014. An fMRI study of cerebrovascular reactivity and perfusion in obstructive sleep apnea patients before and after CPAP treatment. Sleep Med. 15 (8), 892–898. doi: 10.1016/j.sleep.2014.04.004. [PubMed: 24916094]
- Pruim RHR, Mennes M, Buitelaar JK, Beckmann CF, 2015a. Evaluation of ICA-AROMA and alternative strategies for motion artifact removal in resting state fMRI. Neuroimage 112, 278– 287. doi: 10.1016/j.neuroimage.2015.02.063. [PubMed: 25770990]
- Pruim RHRR, Mennes M, Van Rooij D, Llera A, Buitelaar JK, Beckmann CF, van Rooij D, Llera A, Buitelaar JK, Beckmann CF, 2015b. ICA-AROMA: a robust ICA-based strategy for removing motion artifacts from fMRI data. Neuroimage 112, 267–277. doi: 10.1016/ j.neuroimage.2015.02.064. [PubMed: 25770991]
- Puckett AM, Bollmann S, Poser BA, Palmer J, Barth M, Cunnington R, 2018. Using multi-echo simultaneous multi-slice (SMS) EPI to improve functional MRI of the subcortical nuclei of the basal ganglia at ultra-high field (7T). Neuroimage 172 (December 2017), 886–895. doi: 10.1016/ j.neuroimage.2017.12.005. [PubMed: 29208571]
- R Core Team. (2020). R: A Language and Environment for Statistical Computing (3.6.3). https:// www.r-project.org/
- Raj D, Anderson AW, Gore JC, 2001. Respiratory effects in human functional magnetic resonance imaging due to bulk susceptibility changes. Phys. Med. Biol 46 (12), 3331–3340. doi: 10.1088/0031-9155/46/12/318. [PubMed: 11768509]
- Salimi-Khorshidi G, Douaud G, Beckmann CF, Glasser MF, Griffanti L, Smith SM, 2014. Automatic denoising of functional MRI data: combining independent component analysis and hierarchical fusion of classifiers. Neuroimage 90, 449–468. doi: 10.1016/j.neuroimage.2013.11.046. [PubMed: 24389422]
- Salimi-Khorshidi G, Smith SM, Nichols TE, 2011. Adjusting the effect of non-stationarity in cluster-based and TFCE inference. Neuroimage 54 (3), 2006–2019. doi: 10.1016/j.neuroimage.2010.09.088. [PubMed: 20955803]
- Satterthwaite FE, 1946. An approximate distribution of estimates of variance components. Biometr. Bull 2 (6), 110–114. [PubMed: 20287815]

- Schulz J, Siegert T, Bazin PL, Maclaren J, Herbst M, Zaitsev M, Turner R, 2014. Prospective slice-byslice motion correction reduces false positive activations in fMRI with task-correlated motion. Neuroimage 84, 124–132. doi: 10.1016/j.neuroimage.2013.08.006. [PubMed: 23954484]
- Setsompop K, Gagoski BA, Polimeni JR, Witzel T, Wedeen VJ, Wald LL, 2012. Blipped-controlled aliasing in parallel imaging for simultaneous multislice echo planar imaging with reduced g-factor penalty. Magn. Reson. Med 67 (5), 1210–1224. doi: 10.1002/mrm.23097. [PubMed: 21858868]
- Shrout PE, Fleiss JL, 1979. Intraclass correlations: uses in assessing rater reliability. Psychol. Bull 86 (2), 420–428. doi: 10.1037/0033-2909.86.2.420. [PubMed: 18839484]
- Šidák Z, 1967. Rectangular confidence regions for the means of multivariate normal distributions. J. Am. Statist. Assoc 62 (318), 626–633.
- Smith SM, Miller KL, Moeller S, Xu J, Auerbach EJ, Woolrich MW, Beckmann CF, Jenkinson M, Andersson J, Glasser MF, Van Essen DC, Feinberg DA, Yacoubb ES, Ugurbil K, 2012.
 Temporally-independent functional modes of spontaneous brain activity. PNAS 109 (8), 3131–3136. doi: 10.1073/pnas.1121329109. [PubMed: 22323591]
- Smyser CD, Inder TE, Shimony JS, Hill JE, Degnan AJ, Snyder AZ, Neil JJ, 2010. Longitudinal analysis of neural network development in preterm infants. Cereb. Cortex 20 (12), 2852–2862. doi: 10.1093/cercor/bhq035. [PubMed: 20237243]
- Soltysik DA, Hyde JS, 2006. Strategies for block-design fMRI experiments during task-related motion of structures of the oral cavity. Neuroimage 29 (4), 1260–1271. doi: 10.1016/j.neuroimage.2005.08.063. [PubMed: 16275020]
- Sotiropoulos SN, Moeller S, Jbabdi S, Xu J, Andersson JL, Auerbach EJ, Yacoub E, Feinberg D, Setsompop K, Wald LL, Behrens TEJ, Ugurbil K, Lenglet C, 2013. Effects of image reconstruction on fiber orientation mapping from multichannel diffusion MRI: reducing the noise floor using SENSE. Magn. Reson. Med 70 (6), 1682–1689. doi: 10.1002/mrm.24623. [PubMed: 23401137]
- Sousa I, Vilela P, Figueiredo P, 2014. Reproducibility of hypocapnic cerebrovascular reactivity measurements using BOLD fMRI in combination with a paced deep breathing task. Neuroimage 98, 31–41. doi: 10.1016/j.neuroimage.2014.04.049. [PubMed: 24769177]
- Stickland Rachael C., Zvolanek Kristina M., Moia Stefano, Ayyagari Apoorva, Caballero-Gaudes César, Bright Molly G., et al., 2021. A practical modification to a resting state fMRI protocol for improved characterization of cerebrovascular function. bioRxiv doi: 10.1101/2021.02.15.431289. https://www.biorxiv.org/content/early/2021/02/16/2021.02.15.431289.1.
- Tancredi FB, Hoge RD, 2013. Comparison of cerebral vascular reactivity measures obtained using breath-holding and CO₂ inhalation. J. Cereb. Blood Flow Metab 33 (7), 1066–1074. doi: 10.1038/jcbfm.2013.48. [PubMed: 23571282]
- Tchistiakova E, Anderson ND, Greenwood CE, Macintosh BJ, 2014. Combined effects of type 2 diabetes and hypertension associated with cortical thinning and impaired cerebrovascular reactivity relative to hypertension alone in older adults. NeuroImage: Clin. 5, 36–41. doi: 10.1016/j.nicl.2014.05.020. [PubMed: 24967157]
- The phys2bids developers, Alcalá D, Ayyagari A, Bright M, Ferrer V, Caballero-Gaudes C, Hayashi S, Markello R, Moia S, Stickland R, Uruñuela E, Zvolanek K, 2019. physiopy/phys2bids: BIDS formatting of physiological recordings. Zenodo doi: 10.5281/zenodo.3586045.
- Thomason ME, Burrows BE, Gabrieli JDE, Glover GH, 2005. Breath holding reveals differences in fMRI BOLD signal in children and adults. Neuroimage 25 (3), 824–837. doi: 10.1016/ j.neuroimage.2004.12.026. [PubMed: 15808983]
- Tong Y, Bergethon PR, Frederick B.de B., 2011. An improved method for mapping cerebrovascular reserve using concurrent fMRI and near-infrared spectroscopy with Regressor Interpolation at Progressive Time Delays (RIPTiDe). Neuroimage 56 (4), 2047–2057. doi: 10.1016/j.neuroimage.2011.03.071. [PubMed: 21459147]
- Tustison NJ, Cook PA, Klein A, Song G, Das SR, Duda JT, Kandel BM, van Strien N, Stone JR, Gee JC, Avants BB, 2014. Large-scale evaluation of ANTs and FreeSurfer cortical thickness measurements. Neuroimage 99, 166–179. doi: 10.1016/j.neuroimage.2014.05.044. [PubMed: 24879923]

- Urback AL, MacIntosh BJ, Goldstein BI, 2017. Cerebrovascular reactivity measured by functional magnetic resonance imaging during breath-hold challenge: a systematic review. Neurosci. Biobehav. Rev 79 (April), 27–47. doi: 10.1016/j.neubiorev.2017.05.003. [PubMed: 28487157]
- Van Oers CAMM, Van Der Worp HB, Kappelle LJ, Raemaekers MAH, Otte WM, Dijkhuizen RM, 2018. Etiology of language network changes during recovery of aphasia after stroke. Sci. Rep 8 (1), 1–12. doi: 10.1038/s41598-018-19302-4. [PubMed: 29311619]
- Vannesjo SJ, Wilm BJ, Duerst Y, Gross S, Brunner DO, Dietrich BE, Schmid T, Barmet C, Pruessmann KP, 2015. Retrospective correction of physiological field fluctuations in high-field brain MRI using concurrent field monitoring. Magn. Reson. Med 73 (5), 1833–1843. doi: 10.1002/mrm.25303. [PubMed: 24903278]
- Webster MW, Makaroun MS, Steed DL, Smith HA, Johnson DW, Yonas H, 1995. Compromised cerebral blood flow reactivity is a predictor of stroke in patients with symptomatic carotid artery occlusive disease. J. Vasc. Surg 21 (2), 338–345. doi: 10.1016/S0741-5214(95)70274-1. [PubMed: 7853605]
- Wilm BJ, Nagy Z, Barmet C, Vannesjo SJ, Kasper L, Haeberlin M, Gross S, Dietrich BE, Brunner DO, Schmid T, Pruessmann KP, 2015. Diffusion MRI with concurrent magnetic field monitoring. Magn. Reson. Med 74 (4), 925–933. doi: 10.1002/mrm.25827. [PubMed: 26183218]
- Xu Y, Tong Y, Liu S, Chow HM, AbdulSabur NY, Mattay GS, Braun AR, 2014. Denoising the speaking brain: toward a robust technique for correcting artifact-contaminated fMRI data under severe motion. Neuroimage 103, 33–47. doi: 10.1016/j.neuroimage.2014.09.013. [PubMed: 25225001]
- Yezhuvath US, Lewis-Amezcua K, Varghese R, Xiao G, Lu H, 2009. On the assessment of cerebrovascular reactivity using hypercapnia BOLD MRI. NMR Biomed. 22 (7), 779–786. doi: 10.1002/nbm.1392. [PubMed: 19388006]
- Zacà D, Jovicich J, Nadar SR, Voyvodic JT, Pillai JJ, 2014. Cerebrovascular reactivity mapping in patients with low grade gliomas undergoing presurgical sensorimotor mapping with BOLD fMRI. J. Magn. Reson. Imaging 40 (2), 383–390. doi: 10.1002/jmri.24406. [PubMed: 24338845]
- Zaitsev M, Akin B, LeVan P, Knowles BR, 2017. Prospective motion correction in functional MRI. Neuroimage 154 (November 2016), 33–42. doi: 10.1016/j.neuroimage.2016.11.014. [PubMed: 27845256]
- Ziyeh S, Rick J, Reinhard M, Hetzel A, Mader I, Speck O, 2005. Blood oxygen level-dependent MRI of cerebral CO2 reactivity in severe carotid stenosis and occlusion. Stroke 36 (4), 751–756. doi: 10.1161/01.STR.0000157593.03470.3d. [PubMed: 15705935]





Schematic of Breath-Hold trial. Apnoea was preceded and followed by exhalations.



Fig. 2.

 $P_{ET}CO_2hrf$ trace for all subjects and all sessions. Rejected sessions are plotted in red. Rejection was based on having less than three proper $P_{ET}CO_2$ increases after breathholds or having more $P_{ET}CO_2$ decreases than increases after breathholds. Note that the first session of subject 10 was lost due to a software malfunction during acquisition.



Fig. 3.

Unthresholded CVR map obtained with the different lagged-GLM analysis for all of the sessions of a representative subject (subject 002). The maps are not thresholded for statistical significance, but the voxels with lag that is higher than 8.4 (or lower than -8.4) seconds are removed (see text). Note the low CVR response in ME-AGG, depicting numerous voxels with a negative values, as well as the increased amount of masked voxels in SE-MPR, ME-AGG and ME-MOD due to high lag value (see text). SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative. The CVR maps of other subjects are available in the Supplementary Material. The thresholded version of this figure is available at https://zenodo.org/record/ 4739486.



Fig. 4.

(A) Average GM %DVARS and %BOLD response of all BH trials across ten sessions for the same representative subject. The apnoea period is highlighted in grey. Each transparent line is a trial, the solid line is the average across all the trials. (B) Manhattan distance between the average of N trials and the average of all 80 BH trials as N increases from 1 to 80 for each subject. Each vertical line divides the number of trials in each session. SE-PRE: raw data; SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative. The average %DVARS and %BOLD response of the other subjects can be found in the Supplementary Material (Supplementary figure 2).



Fig. 5.

Thresholded CVR map obtained with the different lagged-GLM analysis for all the sessions of a representative subject (subject 002). Note the low CVR response in ME-AGG, depicting numerous voxels with a negative values, as well as the increased amount of masked voxels in SE-MPR, ME-AGG and ME-MOD. SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative. The CVR maps of other subjects are available in the Supplementary Material.



Fig. 6.

Unthresholded lag map obtained with the different lagged-GLM analysis, for all the sessions of a representative subject (same as Fig. 5). These lag maps represent the delay between the best shifted version of the $P_{ET}CO_2hrf$ trace and the bulk shift (i.e. the best match between average grey matter signal and $P_{ET}CO_2hrf$ trace). The scale from -5 to +5 represents earlier to later haemodynamic responses. Note the lack of anatomically informative patterns in ME-MOD and ME-AGG. SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative. The lag maps of other subjects are available in the Supplementary Material.



Fig. 7.

Average values of CVR, lag, and percentage of significant voxels, for voxels in the grey and white matter tissues separately, for all denoising strategies. The dots correspond to a singular session of a singular subject considered an outlier in the distribution. Note that all maps were thresholded before plotting. SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative.



Fig. 8.

Top row: Thresholded χ value of the LME model used for the comparison of CVR and lag maps across all denoising strategies. Other rows: pairwise comparison between denoising strategies. Arrows indicate areas vascularised by big vessels. SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative.



Fig. 9.

ICC(2,1) maps of CVR (left) and haemodynamic lag (right) for each analysis pipeline. The maps are thresholded at 0.4 since scores lower than it indicate poor reliability. A high ICC score indicates that the inter-subject variability is higher than the intra-session variability, while a low ICC score suggest that the variability across sessions is the same as the one across subjects. Following the classification given by Cicchetti (2001), an ICC score lower than 0.4 is considered poor, lower than 0.6 fair, lower than 0.75 good, and equal or higher than 0.75 excellent. The bottom rows depict the whole brain distribution of ICC scores across voxels. Note how OC-MPR and ME-CON have generally higher ICC scores than the other approaches, and are very similar to each other, while ME-AGG has the lowest ICC scores for both CVR and lag maps. SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative. The distribution of ICC scores across grey matter voxels only is available in the Supplementary Material (Supplementary figure 6).

~
\sim
~
<u> </u>
t
5
0
>
\leq
Ma
Mar
Man
Manu
Manus
Manusc
Manusci
Manuscri
Manuscrip

Author Manuscript

Table 1

Comparisons of motion dependance in image intensity and general noise between different denoising approaches.

SF.MPR	OC-MPR	ME-CON	ME-MOD	ME-AGG	
β =512.65 [*] , CI ₉₅ [506.88, 518.42]	<i>β</i> =527.21 [*] , Cl ₉₅ [521.74, 532.68]	<i>β</i> =659.34 [*] , CI ₉₅ [654.27, 664.41]	<i>β</i> =658.05 [*] , CI ₉₅ [653.09, 663.01]	β =715.10 [*] , CI ₉₅ [710.17, 720.04]	SE-PRE
	β =715.10 [*] , Cl ₉₅ [710.17, 720.04]	β =146.69 [*] , CI ₉₅ [143.05, 150.33]	<i>β</i> =145.40 [*] , CI ₉₅ [141.92, 148.88]	<i>β</i> =202.45 [*] , CI ₉₅ [199.02, 205.88]	SE-MPR
		<i>β</i> =132.13 [*] , CI ₉₅ [128.99, 135.27]	<i>β</i> =130.84 [*] , CI ₉₅ [127.89, 133.79]	β =187.90 [*] , CI ₉₅ [185.01, 190.78]	OC-MPR
			<i>β</i> =1.29, CI ₉₅ [-0.70, 3.27]	β =55.77 *, CI ₉₅ [53.91, 57.63]	ME-CON
				<i>β</i> =57.05 [*] , CI ₉₅ [55.59, 58.52]	ME-MOD
*			:		

significant for p<0.001; all p values are computed with Satterthwaite's method, and they are the equivalent of the p value after Šidák correction for multiple comparisons. SE-PRE: raw data; SE-MPR: single-echo; OC-MPR: optimally combined; ME-AGG: aggressive; ME-MOD: moderate; ME-CON: conservative.

Author Manuscript

Moia et al.

Subject	Average value	SE-MPR	OC-MPR	ME-AGG		
001	CVR [%BOLD/mmHg]	0.54	0.5	0.37	0.43	0.49
	Lag [s]	-0.54	-0.49	0.4	-0.11	-0.4
	% significant voxels	9.79	10.44	3.33	8.1	10.75
002	CVR [%BOLD/mmHg]	0.38	0.35	0.24	0.3	0.35
	Lag [s]	-0.38	-0.43	0.58	0	-0.42
	% significant voxels	10.67	11.65	3.29	8.61	11.92
003	CVR [%BOLD/mmHg]	0.4	0.34	0.18	0.31	0.33
	Lag [s]	-0.4	-0.28	-0.73	-0.74	-0.26
	% significant voxels	7.42	8.04	3.62	6.55	8.28
004	CVR [%BOLD/mmHg]	0.44	0.38	0.08	0.32	0.37
	Lag [s]	-1	-1.12	-0.57	-0.87	-1.1
	% significant voxels	8.46	9.27	2.87	6.57	9.56
007	CVR [%BOLD/mmHg]	0.33	0.29	0.17	0.28	0.29
	Lag [s]	-0.7	-0.61	0.95	-0.1	-0.53
	% significant voxels	7.98	9.19	2.6	6.31	9.44
008	CVR [%BOLD/mmHg]	0.34	0.14	-0.03	0.26	0.14
	Lag [s]	-0.98	-1.19	-0.28	-0.6	-1.18
	% significant voxels	6.34	6.89	1.5	4.9	7.19
600	CVR [%BOLD/mmHg]	0.44	0.38	-0.18	0.31	0.37
	Lag [s]	-1.75	-1.75	0.91	-0.11	-1.69
	% significant voxels	7.52	9.16	2.25	6.42	9.55
Total	CVR [%BOLD/mmHg]	0.41	0.34	0.12	0.32	0.33
	Lag [s]	-0.82	-0.84	0.18	-0.36	-0.80
	% significant voxels	8.31	9.23	2.78	6.78	9.53