
Copyright:

This is an author copy of a manuscript presented at the ISMRM-ESMRMB Joint Meeting 2014.

Date deposited:

15/09/2015
Brain oxygenation responses to an autonomic challenge: 
a quantitative fMRI investigation of the Valsalva manoeuver

Iwo Bohr, Claire McDonald, Jiabao He, Simon Kerr,
Julia L Newton, Andrew M Blamire

a Institute of Cellular Medicine & Newcastle Magnetic Resonance Centre, Campus for Ageing and Vitality, Newcastle University, Newcastle upon Tyne, NE4 5PL, UK,
b Newcastle University Institute for Ageing, Newcastle University, Newcastle upon Tyne, NE4 5PL, UK

c UK NIHR Biomedical Research Centre in Ageing, Newcastle upon Tyne NE4 5PL, UK
d Aberdeen Biomedical Imaging Centre, Lilian Sutton Building, Foresterhill, University of Aberdeen, Aberdeen, AB25 2ZD, Scotland, UK,
e Institute for Ageing and Health, Campus for Ageing and Vitality, Newcastle University, Newcastle upon Tyne, NE4 5PL, UK,

Corresponding author:
Dr Iwo Bohr, PhD,
Newcastle Magnetic Resonance Centre,
Campus for Ageing and Vitality,
Newcastle University, Newcastle upon Tyne,
NE4 5PL,
United Kingdom
Tel. +44 191 208 1160
corresponding author: iwo.bohr@newcastle.ac.uk

Acknowledgements
The research was funded by the National Institute for Health Research (NIHR) Newcastle Biomedical Research Centre based at Newcastle upon Tyne Hospitals NHS Foundation Trust, the British Geriatric Society and Research into Ageing Fund; a fund set up and managed by Age UK. The views expressed are those of the authors and not necessarily those of the NHS, the NIHR or the Department of Health. The authors are grateful to Dr Michael Firbank (Newcastle University) for providing us with his code for WMH segmentation and guidelines for its usage and Dr Andreas Finkelmeyer (Newcastle University) for his valuable comments and advice.
Abstract

In late age the autonomic nervous system (ANS) has diminished ability to maintain physiological homeostasis in the brain in response to challenges such as to systemic blood pressure changes caused by standing. We devised an fMRI experiment aiming to map the cerebral effects of an ANS challenge (Valsalva manoeuvre; VM). We used dual-echo fMRI to measure the effective transverse relaxation rate ($R_2^*$, which is inversely proportional to brain tissue oxygenation levels) in 45 elderly subjects (median age: 80 years old, total range: 75-89) during performance of the VM. In addition we collected Fluid Attenuation Inversion Recovery (FLAIR) data from which we quantified white matter hyperintensity (WMH) volumes. We conducted voxelwise analysis of the dynamic changes in $R_2^*$ during the VM to determine the distribution of oxygenation changes due to the autonomic stressor. In white matter we observed significant decreases in oxygenation levels. These effects were predominantly located in posterior white matter and to a lesser degree in the right anterior brain, both concentrated around the border zones (watershed) between cerebral perfusion territories. These areas are known to be particularly vulnerable to hypoxia and are prone to formation of white matter hyperintensities. Although we observed overlap between localization of WMH and triggered deoxygenation on the group level we did not find significant association between these independent variables using subject-wise statistics. This could suggest other than recurrent transient hypoxia mechanisms causing/contributing to the formation of WMH.

Keywords: autonomic nervous system; brain aging; Valsalva manoeuvre; hypoxia; watershed zones; fMRI; effective transverse relaxation rate ($R_2^*$)
Abbreviations: ACA, anterior artery perfusion territory; ANS, autonomic nervous system; BOLD, Blood Oxygenation Level Dependent; CBF, cerebral blood flow; EPI, echo planar imaging; GLM, General Linear Model; GM, gray matter; HWGFS, half-width of the gamma function smoothing of the input waveform; MCA, medial cerebral artery territory; R₂*: effective transverse relaxation rate; TE, echo time; TR, repetition time; VM, Valsalva manoeuvre; WM, white matter; WMH, white matter hyperintensities
1. Introduction

Brain functional and structural integrity depends heavily on maintaining energetic homeostasis, with a particular importance of preserving levels of cerebral perfusion. The extent of this dependence is emphasized by the fact that the human brain constitutes only about 2% of the whole body weight but consumes as much as 20% of the total oxygen and other resources supplied by blood (Kalaria 2010). The autonomic nervous system (ANS) plays a pivotal role in securing a stable environment in the brain in response to changes in peripheral blood pressure which may affect brain perfusion. In this work we applied an fMRI method to examine the transient changes in brain tissue oxygenation in response to an autonomic challenge in healthy elderly subjects.

In everyday life the ANS must continually compensate for varying blood pressure in response to stress such as standing. The pressure of blood entering the brain drops significantly as a result of taking an upright position from sitting and even more from lying, requiring swift adjustments to heart rate, stroke volume and peripheral vascular resistance in order to maintain perfusion pressure of the brain within the limits of tight cerebral autoregulation (Perlmuter et al. 2013). The capacity of the ANS to maintain physiological homeostasis in response to such autonomic challenge declines with age (Hotta and Uchida 2010) and this may have a direct impact on the integrity of the brain. In addition to autonomic control of arterial blood pressure and cardiac output, cerebral autoregulation acts within the brain to further dampen transient changes in pressure and maintain a constant level of tissue perfusion. However it should be borne in mind that age-related arteriosclerosis is another factor contributing to compromised blood supply the brain tissue (Kalaria 2010). Mandell et al (Mandell et al. 2008) have shown that gray matter (GM) is preferentially defended against deficits in perfusion under stressful conditions by “taking it away” from the white matter (WM) – a “steal phenomenon” – such that WM is potentially more susceptible to worsened homeostatic control. In addition, some features of the vasculature in the WM, including low number of anastomoses in the terminal type of vessels, increase its vulnerability by compromising blood supply in this tissue under conditions of a raised demand for blood flow (Mangla et al. 2011; Pantoni 2002). The brain tissue is also particularly vulnerable to perturbations in perfusion in the watershed zones at the borders of the major cerebral arterial perfusion territories. The watershed zones are characterized by a high susceptibility to
hypoxia (Caine and Watson 2000) and hypo-perfusion (Lanterna et al. 2011; Mandell et al. 2008; Mangla et al. 2011).

Frequently the ageing brain is characterised by areas of hyperintense signal known as white matter hyperintensities (WMH) seen in the WM on T2 weighted MRI scans, particularly using the Fluid Attenuation Inversion Recovery (FLAIR) sequence (Grueter and Schulz 2012; O'Sullivan 2008). The WMH load is associated with age and a number of comorbidities such as stroke and cognitive decline (Debette and Markus 2010). Ischemia is considered as a leading pathological feature of WMH (Fernando et al. 2006; O'Sullivan 2008) and a reduction in perfusion during hypercapnic challenge has been shown within tissue with a high density of WM lesions in elderly subjects, suggesting a possible causative link between selective hypoxia in watershed zones and white matter damage (Mandell et al. 2008). The presence of WMH has been associated with a range of conditions where formal laboratory testing reveals autonomic dysfunction (Galluzzi et al. 2009; Newton et al. 2008; Richardson et al. 2009; Waldstein et al. 2004) and one study has found strong association between altered heart rate variability (a measure of autonomic control) and WMH burden in patients with mild cognitive impairment (Galluzzi et al. 2009). Together these data suggest that autonomic dysfunction may be an important mechanism contributing to WMH formation.

The Valsalva Manoeuvre (VM) is a standard method used to induce changes in blood pressure and heart rate and to study the integrity of the ANS in the clinic. The test requires the subject to maintain an expirational pressure against a closed outlet of 40 mmHg, normally for about 16 s. The physiological response to the VM is divided into four phases (I-IV) characterised by changes in mean arterial blood pressure and cardiac output which reflect sympathetic and parasympathetic activity and which must stimulate cerebral autoregulation (Dawson et al. 1999). In the present study we used Blood Oxygenation Level Dependent (BOLD) functional MRI (fMRI) to non-invasively study the neuro-physiological effects of the VM as a challenge to the ANS via the sensitivity of BOLD to changes in brain tissue oxygenation. We specifically aimed to test the hypothesis that watershed perfusion areas, known to be vulnerable to white matter injury, would show significant tissue deoxygenation during autonomic challenge and that there would be a direct relationship between degree of deoxygenation and the present of WMH in our subjects.
2. Methods

2.1. Participants and Recruitment

The current work is part of a wider longitudinal study examining the prevalence and clinical associations of neurocardiovascular instability and autonomic nervous system function in older community dwelling people who have been studied since 2002 (for detail of the study see: Kerr et al. 2006). A subsample of 45 subjects from the cohort took part in the current study. Participants were aged 77 years old or over when they were scanned (2012). There were 24 male subjects, median age was 80.0 years, interquartile range=77.0 - 83.3 years. Data on medical history and medications taken presented in Table 1 were based on records collected at the initial recruitment wave and updated at the time of MRI scanning sessions based on participant interviews and general practitioner medical records. The study was approved by the local ethical committee and all subjects provided written informed consent. A community dwelling, healthy older age subject group was specifically chosen as they would be expected to have some levels of WMH.

5.2. Experimental procedure

Participants underwent MRI scanning including an fMRI scan during which they performed a VM protocol as a dynamic challenge to the ANS (Meyer et al. 1966; Taylor 1996) according to our previously reported procedure (He et al. 2013). In brief, during a single fMRI time series acquisition subjects performed four episodes of VM, each lasting 16s (time of exhaling strain). The interval between the onsets of each VM was 60s. The time course of the protocol is shown in Figure 2A. Participants’ performance was guided by instructions projected on a MR scanner display system. First, an instruction to prepare for the VM (to inhale) was presented (5s ahead of the manoeuvre) and then the instruction ‘start’ was displayed followed by the instruction ‘stop’ at 16s after ‘start’. During the strain (pressuring phase) a visual feedback of the instantaneous pressure was displayed to the subject as a bar chart graphic showing exhaled pressure relative to the target of 40 mmHg. The displayed content was controlled by an in-house designed controller device (programmed in Labview 7.0, National Instruments, USA) which also recorded the data of subject performance (He et al. 2013). An example of an exhaled pressure trace is shown in Figure 2B.
5.3. Imaging

Scanning was performed on a 3T Intera Achieva machine (Philips Healthcare Systems, Best, The Netherlands) using an 8-channel SENSE coil. For measurement of BOLD signal a customized dual gradient echo EPI sequence was employed (135 volumes were acquired with TR = 2000 ms, TE1/TE2 = 13.82/39.27 ms, flip angle = 90°, field of view = 240×240 mm, matrix = 112×112, with 20 slices of 4 mm thickness). The simultaneous collection of 2 echo times allows calculation of the effective transverse relaxation rate (R2*), which varies with tissue oxygenation level and allows separation of tissue oxygenation changes from inflow effects (Glover et al. 1996) associated with the changes in blood pressure, which are a major feature of the VM. For the purpose of displaying BOLD statistical maps, a 3D T1-weighted anatomical image was also acquired (TR = 8.3 ms, TE = 4.6 ms, flip angle = 8°, field of view = 216×239×180 mm, isometric 1 mm voxels). These images were averaged and registered to the MNI space, creating a study specific template reflecting age-related changes in anatomy.

For WMH segmentation, we acquired fluid attenuated inversion recovery (FLAIR) images (TR=11000 ms, TE = 125 ms, TIR=2800 ms; Turbo SE factor = 27; refocus angle = 120; field of view (FOV) = 240×240 mm, matrix size = 256×256, with 50 slices of 3mm thickness, SENSE factor = 1.5). Figure 1 shows examples of images from each modality employed in this study from one typical subject.

5.4. Image pre-processing and analysis

All pre-processing and statistical analysis steps were performed using the FSL package (version 1.4.9, FMRIB's Software Library, www.fmrib.ox.ac.uk/fsl) (Smith et al. 2004; Woolrich et al. 2009) and customized Matlab scripts (Matlab R2012a; Mathworks Inc., Natick, MA, USA; including make_nii and save_nii functions, part of the “Tools for NIfTI and ANALYZE image” package written by Jim Shen and freely available on Matlab file exchange website (Shen 2005).

Echo 1 (E1) images were pre-processed using motion correction (Jenkinson et al. 2002) followed by brain extraction (Smith 2002). The resultant transformation matrix and brain masks were then applied to the echo 2 (E2) images bringing all images into alignment. Images of R2* were created based on the pre-processed E1 and E2 images according to the standard formula:
\[ R_2^* = -\ln \frac{S_2}{S_1} \frac{T E_2 - T E_1}{S_2} \]

where \( S_1 \) and \( S_2 \) are the voxel intensities in the images with echo times TE$_1$ and TE$_2$ respectively.

Estimation of the effects of the VM on \( R_2^* \) was carried out using a mixed-effects General Linear Model (GLM) as implemented in the FEAT tool (FSL library, GUI version). The FEAT steps also included further standard fMRI pre-processing procedures: high pass filtering with a 100s cut-off, noise reduction (SUSAN tool) (Smith and Brady 1997), linear registration to the template and spatial smoothing with 5 mm isotropic Gaussian kernel.

To determine the basic form of the GLM model we first extracted the \( R_2^* \) response averaged across the whole brain and across all subjects as shown in Figure 1D. Neural tissue (gray matter + white matter) was segmented in each individuals anatomical image using the SPM8 package (Wellcome Department of Imaging Neuroscience Group, London, UK) and applied as a global mask to the \( R_2^* \) image series from which the signal was extracted and subsequently averaged, normalized and smoothed using a spline algorithm (Fig. 1B). Responses during the ‘strain’ or pressuring (exhaling air against a closed glottis) were modelled with a boxcar function convolved with a gamma function. The convolution parameters were selected heuristically in order to obtain the best possible match to the average extracted \( R_2^* \) signal as described above (Fig. 1E). They were as follows: duration of 16 s, a mean time lag of the gamma smoothing of the input waveform (mean lag) of 6s and half-width of the gamma function smoothing of the input waveform (HWGFS) set to 4s.

To examine the oxygenation changes during the VM we conducted a first level analysis contrasting signal change during strain against baseline in each individual subject. To perform groupwise statistical analysis the first level statistical maps were registered to the MNI space. The second level analysis was then performed with the FSL FLAME1 random effects tool. Reported are thresholded statistical maps \((z>2.3)\) forming significant clusters \((p<0.05, \text{Family Wise Error corrected})\).
In order to assess the localization of $R_2^*$ changes relative to the location of the vascular watershed zones we also created binary masks representing the major arterial perfusion territories in standard space. These were generated by manually defining the territory boundaries on the standard space T1-weighted image template using anatomical hallmarks as described in (Ferrer et al. 2008). The spatial distribution of significant $R_2^*$ changes found in the second level analysis were then visually compared against these masks.

To quantify WMH volumes we used a semi-automated threshold-based algorithm employing SPM8 (Wellcome Department of Imaging Neuroscience Group, London, UK; http://www.fil.ion.ucl.ac.uk/spm/) functions embedded in an in house Matlab (Matlab 8.1.0.604; R2013a, Mathworks, Inc., Natick, MA, USA) package as described previously (Firbank et al. 2003; Woolrich et al. 2009). Briefly, first WM, GM and CSF segments were defined using the subjects T1-weighted image and the SPM Segment toolbox (Ashburner and Friston 2005) followed by registering them to the acquisition space of the FLAIR images. Subsequently WMH were identified by applying a threshold to slice-specific intensity histograms of the tissue within the WM segment. Resulting masks were manually edited if necessary by two instructed raters using ITK-SNAP software (www.itksnap.org) to account for minor inaccuracies in the automated procedure.

To compare the volumes of WMH and the spatial extent of significant deoxygenation we registered individual WMH masks and $R_2^*$ change z-maps to the study template in the MNI space. For each subject we then quantified the volume of WMH in MNI space as a fraction of the total white matter and the volume of significantly deoxygenating white matter on $R_2^*$ maps. For each individual, we also calculated the proportion of WMH which was contained within the area of significant deoxygenating tissue ($z \geq 2.3$; corrected) and also the proportion of WMH which was in non-deoxygenating tissue. Further, to test for any association between the magnitude of transient deoxygenation and presence of WMH in individuals we ran a voxel-wise logistic regression analysis. The distribution of WMH across the whole group was first determined by averaging the individual WMH masks in MNI space. We then analysed all voxels where there was a non-zero probability of finding WMH within the group and carried out a binomial regression between voxel the z-values (degree of deoxygenation) and the presence or absence (binary) of WMH in the voxel. Analysis was conducted using the multinomial logistic regression framework as implemented in Matlab statistics toolbox (mnrfit function).
3. Results

Typical image data from one subject is shown in Figure 1 which illustrates the presence of WMH on the FLAIR scan, a basic fMRI scan and the calculated $R_2^*$ map at the same brain level.

3.1. $R_2^*$ data

Illustrations of the basic fMRI time series for each of the 2 echo times and the $R_2^*$ series derived from these data are shown in Figures 2C and 2D respectively. It is observed that during the first ~5s (equating to approximately phase I and II of the VM) there are almost identical positive signal changes in both of the E1 and E2 time series, indicating significant effects of inflow associated with the increased mean arterial blood pressure during these phases of the VM. This effect is not seen in the $R_2^*$ response where the combination of data from the two echoes has effectively separated the blood pressure changes and inflow effects from oxygenation changes (Glover et al. 1996). $R_2^*$ is observed to gradually increase up to and slightly beyond the end of the VM indicating rising deoxygenation of the tissue during this time, followed by a decline to baseline with a small undershoot on release of the strain.

The GLM analysis was defined to evaluate regional patterns of tissue deoxygenation during phases I-III of the VM (where higher $R_2^*$ equates to greater tissue deoxygenation) and the resulting statistical maps ($z>2.3$, cluster-corrected) are shown in Figure 3 and the characteristics of the significant clusters are summarized in Table 2. The $R_2^*$ changes were concentrated in the right posterior part of the WM including areas adjacent to the posterior horns of the lateral ventricles (Fig. 3). A similar distribution could be observed in the left hemisphere but with lower values of $z$-statistics and less spatial extent. In addition there was a significant cluster in the left anterior part of the brain. The deoxygenation maps showed the greatest effect within the territory of the middle cerebral artery (MCA) but also overlapped extensively with the watershed zone separating the MCA and posterior cerebral artery (PCA) territories in the posterior brain and between anterior CA (ACA) and MCA in the left anterior brain. Voxels characterized by higher $z$-statistics tended to concentrate near the territory boundaries (Fig. 3B and C).

3.2. Relationship between the presence of WMH and deoxygenation

The total volume of WHM measured as a percentage of the whole WM was (mean±SD): 1.52±1.73%, while the volume of tissue with statistically significant $R_2^*$ deoxygenation was
10.94±7.10%. To quantify the relationship between these measurements we calculated the degree of overlap between the WMH and deoxygenating tissue in each subject. Only 1.9±1.3% of the deoxygenating tissue was composed of pixels classified as WMH. It is clear from these values that the spatial extent of R₂* effects is considerably and significantly larger (p<<0.001 for two-sample t-test). Conversely, when we examined the location of the WMH only 38.1±19.8% of the WMH was found within the deoxygenating tissue. This is consistent with the fact that WMH are particularly located around the anterior horns of the ventricles and the superior regions of the corona radiate where the deoxygenation changes where smaller or non-significant. In line with these values a more refined, formal logistic regression analysis did not find any significant pixelwise association between the magnitude of transient deoxygenation and presence of WMH. Thus the R₂* changes are not simply reflective of the apparent WM pathology seen on the FLAIR images.

4. Discussion

In this study we analysed the changes in brain tissue oxygenation during an autonomic challenge. We found that deoxygenation changes during the VM strain were greatest in the WM border zones between the middle and posterior and to a lesser extent between anterior and middle cerebral arterial territories, while the GM showed no significant changes. The spatial extent of the areas of deoxygenation during VM was significantly greater than the areas of WMH, while conversely only a proportion of the observed WMH were located in deoxygenating white matter. We did not find any evidence of a direct association between the degree of transient deoxygenation and the presence of WMH.

3.1. Global hypoxic changes

Strong responses during the strain period were found only in the WM and reflected widespread reduction in WM tissue oxygenation. We did not find any areas of significant deoxygenation in the GM. Together these observations suggest that in the presence of reduced cardiac output during the VM, the autoregulatory system protects the GM at the expense of the WM. Importantly we found that the WM areas with highly significant tissue deoxygenation were strongly overlapping with the watershed zone separating the two major arterial territories in the WM: medial cerebral artery (MCA) and posterior cerebral artery (PCA) and also between anterior (ACA) and MCA in the left hemisphere (Fig. 3B). This finding provides support for the particular vulnerability of the WM to transient episodes of hypoxia and ischemia. Mandell and co-workers (Mandell et al. 2008) using fMRI and arterial
spin labelling found that this zone was prone to disturbances caused by another physiological stressful condition: hypercapnia, even in young healthy subjects. They described this as a “stealing phenomenon” whereby in conditions of a stress (increased blood CO₂ levels in their study) cerebral blood supplies are in the first instance directed to the GM, diverting them away from the WM. These processes were already detectable in young subjects (Mandell et al. 2008) and we postulate that they may increase during the life span and probably become more pronounced in the elderly. Although the localization of the WM lesions and transient deoxygenation triggered by VM was visually similar we did not find statistical association between these two measures. Thus, although WM lesions emerge in areas vulnerable to deoxygenation triggered by the autonomic challenge this vulnerability, although important, does not appear to be the only or leading cause of WM damage. Other causative factors have been identified such as leakage of potentially toxic substances through a disrupted Blood Brain Barrier (Pantoni 2002; Topakian et al. 2010; Wardlaw et al. 2003).

3.2. Laterality of global deoxygenation
Oxyge
nation changes during the VM strain in this cohort of subjects were right-dominant. This may be related to lateralization in cerebral perfusion at rest which has been reported in previous studies (Pfefferbaum et al. 2010; Rodriguez et al. 1991; Wentland et al. 2010). In particular Pfefferbaum and co-workers found left-right and anterio-posterior pattern of differences in cerebral blood flow in the GM measured by arterial spin labelling MRI. They detected higher flow in the right hemisphere in the anterior part of the brain whereas in the posterior part it was higher in the left hemisphere. We observed higher deoxygenation in response to the VM in the right posterior part of the WM (Fig. 3). This potentially could be explained by lower basal CBF in this part of the brain in the rest conditions as found by (Pfefferbaum et al. 2010), however it should be noted that the study of those authors was restricted to the GM (detection of white matter CBF is challenging using the arterial spin labelling they employed), whereas our findings were the strongest in the WM.

3.3. Comparison to other MRI studies of the VM
Several studies have previously used the VM as a physiological challenge combined with fMRI data acquisition (Henderson et al. 2002; Henderson et al. 2003). In particular, Henderson et al (2002) examined the BOLD response to study the brain regions recruited during the VM using region of interest analysis and correlation of the BOLD response to physiological measures. The authors demonstrated signal changes in key brain regions.
associated with autonomic control. It is interesting to note that in both the studies of Henderson et al and in the current study, changes in the WM were observed and that repeated performance of the VM led to increasing magnitude of BOLD signal change. However, it should be noted that in comparison to our approach Henderson et al used only a single echo time fMRI method which could not separate inflow effects from changes in oxygenation. Our data therefore extend the understanding of the brain response to VM by establishing that tissue deoxygenation does occur in the WM during the VM, at least in the elderly population.

3.5. Limitations

In our study there were no selection criteria other than being elderly, community dwelling subjects. There may be other pathologies involved in the response beyond the effect of ageing itself (see Table 1 for comorbidities and medications taken). It is known that hypertension is a major cardiovascular risk factor for WM damage and is associated with microvascular rarefaction and vessel remodelling (Feihl et al. 2008). In our group 55% of subjects were classified as being hypertensive according to their GP records and medication regime. Although hypertension was medication-controlled we undertook a further second level group analysis of the fMRI data (standard two-group FSL FEAT analysis) comparing those patients with a diagnosis of hypertension against the rest of the cohort, in order to determine whether hypertension was a driving factor in our findings of WM deoxygenation. We found no difference in deoxygenation between participants clustered as hypertensive and the rest of the sample. However an impact of blood pressure cannot be entirely ruled out considering our sample composition and the possible existence of a "pre-hypertensive" status in subjects which were not diagnosed as “hypertensive”.

Our data did not show any direct relationship between the presence of WMH and deoxygenation changes during autonomic challenge. However it must been noted that the variability and total volume of WMH seen in our subjects was relatively restricted and thus our ability to detect any correlation may therefore be limited. Since WMH are a known risk factor for stroke, dementia and death (Debette and Markus 2010) it is conceivable that the elderly nature and general good health status of our community dwelling cohort means our group may be biased towards individuals who are resistant to WMH formation and hence are not totally characteristic of the general population. Having a wider age range would allow for a more sensitive assessment of this potential relationship. It should be also noted that our study evaluated only one type of autonomic challenge (a voluntary increase of thoracic pressure) and the observations may not directly extend to other autonomic tasks.
Methodologically our data were collected using a gradient echo sequence which is sensitive to changes in oxygenation of the large and small vessels as well as the capillary bed. The precise level within the tissue where the deoxygenation takes place (for example in the small arteries and arterioles versus within the capillaries) cannot be conclusively determined from the current data. Further work is needed using spin-echo based perfusion measurements to explore this in more detail. Our dual-echo approach was used in order to map $R_2^*$ and attempt to separate the effects of blood pressure related signal changes from deoxygenation effects in the tissue. Some level of $T_1$ based inflow contrast may still be present at the 2s repetition time which was needed in order to obtain good temporal resolution during the VM performance. In support of our methods however, while the basic BOLD data show biphasic signal changes which closely resemble the blood pressure swings reported during the VM (Dawson et al. 1999), these features are not present in the $R_2^*$ time-series suggesting that $T_1$ inflow effects are not a major feature of the final data.

5. Conclusions
To conclude, this study reports the data on brain oxygenation changes in response to an ANS challenge pointing to watershed zones in the white matter as the areas most vulnerable to transient hypoxia. Our study provide further and novel evidence of the “stealing phenomenon” (Mandell et al. 2008) as a result of autonomic challenge. The data did not however support cumulative effects of autonomic dysfunction as a leading mechanism for formation of WMH.

6. References


Galluzzi S et al. (2009) Cardiac autonomic dysfunction is associated with white matter lesions in patients with mild cognitive impairment The journals of gerontology Series A, Biological sciences and medical sciences 64:1312-1315 doi:10.1093/gerona/glp105


Henderson LA et al. (2002) Brain responses associated with the Valsalva maneuver revealed by functional magnetic resonance imaging Journal of neurophysiology 88:3477-3486 doi:10.1152/jn.00107.2002


Rodriguez G et al. (1991) Regional cerebral blood flow asymmetries in a group of 189 normal subjects at rest Brain topography 4:57-63


10.1016/j.neuroimage.2004.07.051


TABLES

**Table 1** Demographic and clinical data of the participants

<table>
<thead>
<tr>
<th>males\females</th>
<th>24/21</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>median age = 80.0, interquartile range=77.0-83.3, full range=75-89</td>
</tr>
<tr>
<td><strong>MMSE</strong></td>
<td>mean MMSE=27.8±2.0 (SD); full range=23.0-30.0</td>
</tr>
<tr>
<td></td>
<td>median=29.0, interquartile range=26.5-29.0</td>
</tr>
<tr>
<td><strong>medical history</strong></td>
<td>number of occurrences(^a) (%)</td>
</tr>
<tr>
<td>ischemic heart disease</td>
<td>7 (15.56)</td>
</tr>
<tr>
<td>hypertension</td>
<td>24 (53.33)</td>
</tr>
<tr>
<td>stroke or TIA</td>
<td>4 (8.89)</td>
</tr>
<tr>
<td>cardiovascular or cerebrovascular disease</td>
<td>14 (31.11)</td>
</tr>
<tr>
<td>diabetes</td>
<td>7 (15.56)</td>
</tr>
<tr>
<td><strong>medications(^b)</strong></td>
<td>n(^a) (%)</td>
</tr>
<tr>
<td>cardioactive</td>
<td>30 (66.7)</td>
</tr>
<tr>
<td>antihypertensive</td>
<td>29 (64.4)</td>
</tr>
</tbody>
</table>

\(^a\) note there may be more than one disorder occurring (medication taken) for a participant

\(^b\) cardioactive: any antihypertensive, diuretic, antianginal, antiarrythmic, fludrocortiosone or midodrine; anti-hypertensive: any angiotensin converting enzyme (ACE) inhibitors, alpha blockers, beta blockers, calcium channel blockers, angiotensin type 2 receptor blockers and thiazides;
Table 2 Clusters (in the white matter) showing significant transient deoxygenation triggered by the Valsalva manoeuvre.

<table>
<thead>
<tr>
<th>cluster</th>
<th>number of voxels (% brain)</th>
<th>z max (z y z; MNI mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>21026 (2.3)</td>
<td>6.41 (42 -64 26)</td>
</tr>
<tr>
<td>2</td>
<td>2951 (0.3)</td>
<td>5.37 (-14 30 18)</td>
</tr>
</tbody>
</table>
Fig. 1 Examples of four modalities of images used in the study from the same subject (middle axial slice); A: T1 weighted, B: FLAIR (note the presence of WMH, esp. around the horns of the lateral ventricles); C: fMRI; echo 2; D: R₂*
**Fig. 2 A:** Diagram illustrating the timing of Valsalva manoeuvre (VM) performance within the fMRI time series acquisition. Gray bars indicate the 5s period during which subjects were instructed to prepare to perform the VM, while the solid line indicates the timing of each of the 16s periods of VM. **B:** Representative pressure trace of exhaled air during one of the VM periods. The dashed horizontal line depicts required level of pressure. **C:** Group averaged (n=45) time series of the BOLD signal change for the two echo times averaged across the 4 VM periods. Black line - Echo 1, Gray line – Echo 2. The gray box indicates the period of the VM. **D:** Time series of the mean normalized R2* signal (n=45) extracted from the whole brain (WM + GM, but excluding CSF); **E:** Time course of models of response for ‘strain’ during the VM.
Fig. 3 A: Group deoxygenation maps overlaid on study-specific template registered to the MNI space for contrast between ‘strain’ phase and baseline showing highly significant areas of tissue deoxygenation confined to posterior and left anterior white matter. Numbers beneath each slice denote z coordinates (mm). B: Deoxygenation map (z=16) overlaid on outlines of ideal major arterial territories; ACA: anterior cerebral artery perfusion territory, MCA: middle cerebral artery perfusion territory, PCA: posterior cerebral artery perfusion territory, note a strong overlap of deoxygenation maps and watersheds between arterial territories. C: Slices presenting theoretical outlines of perfusion territories - green ACA, red: MCA and blue: PCA (z-coordinates as on panel B). Note radiological orientation. Note radiological orientation.