Cerebrovascular reactivity deficits in cognitively unimpaired older adults: vasodilatory versus vasoconstrictive responses

Belinda Yew a,1, Jung Yun Jang b,1, Shubir Dutt a,1, Yanrong Li b, Isabel J. Sible a, Aimée Gaubert b, Jean K. Ho b, Anna E. Blanken a, Anisa Marshall b, Xingfeng Shao c, Danny J.J. Wang c, Daniel A. Nation b,d,∗

a Department of Psychology, University of Southern California, Los Angeles, CA, USA
b Institute for Memory Impairments and Neurological Disorders, University of California, Irvine, Irvine, CA, USA
c Stevens Neuroimaging and Informatics Institute, University of Southern California, Los Angeles, CA, USA
d Department of Psychological Science, University of California, Irvine, Irvine, CA, USA

ABSTRACT

Cerebrovascular reactivity (CVR) deficits may index vulnerability to vascular brain injury and cognitive impairment, but findings on age-related changes in CVR have been mixed, and no studies to date have directly compared age-related changes in CVR to hypercapnia versus hypocapnia. The present study compared CVR in 31 cognitively unimpaired older adults (ages 55–87) and 30 healthy younger adults (ages 18–28). Breath control tasks induced CVR to hypercapnia (0.1 Hz paced breathing) and hypercapnia (15 s breath holds) during pseudo-continuous arterial spin labeling MRI. Relative to younger adults, cognitively unimpaired older adults displayed lower levels of global CVR under both hypcapnia and hypercapnia. In region-of-interest analyses, older adults exhibited attenuated CVR to hypcapnia in select frontal and temporal regions, and lower CVR to hypercapnia in all cortical, limbic, and subcortical regions examined, relative to younger adults. Results indicate age-related deficits in CVR are detectible even in cognitively unimpaired older adults and are disproportionately related to vasodilatory (hypcapnia) responses relative to vasoconstrictive (hypercapnia) responses. Findings may offer means for early detection of cerebrovascular dysfunction.

© 2022 The Author(s). Published by Elsevier Inc.

This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/)

1. Introduction

Vascular dysfunction has been increasingly implicated in age-related cognitive decline and dementia (Arvanitakis et al., 2016; Qiu and Fratiglioni, 2015; Snyder et al., 2015; Toth et al., 2017). Despite advances in neuroimaging approaches to detection of cerebrovascular lesions (Wardlaw et al., 2019), further studies are needed to establish cerebrovascular markers that may predate irreversible lesion formation and cognitive impairment. Cerebrovascular reactivity (CVR) is one candidate marker that represents the ability of cerebral vessels to modulate cerebral blood flow (CBF) through dilation or constriction in response to vasoactive stimuli (e.g., CO2) (Liu et al., 2019; Urbach et al., 2017). In the largest study of CVR to date, Sur and colleagues evaluated CVR to hypercapnia using blood oxygen level-dependent (BOLD) signal response to CO2 gas inhalation in 72 older adults with normal or impaired cognition (Sur et al., 2020). Findings indicated significant attenuation of CVR in older adults with cognitive impairment relative to those without cognitive impairment. There was also a positive association between CVR and cognition, after accounting for covariates such as white matter lesion burden. Despite these promising findings regarding CVR as a potentially valuable marker of cognitive dysfunction, it remains unclear whether CVR deficits are observable during the preclinical period, when aging individuals are cognitively unimpaired. Thus, studies of age-related CVR deficits in cognitively unimpaired individuals may inform preclinical CVR assessment. Further characterization of age-related changes in CVR may also provide relevant data for investigations of various disorders, such as Moyamoya disease and carotid artery stenosis, as impairments in CVR have significant implications in clinical outcomes (Markus and Cullinane, 2001; Han et al., 2011).
Previous research on age-related alterations in CBF response to CO₂ have yielded mixed results. Both BOLD-fMRI and transcranial doppler (TCD) investigations instigating hypercapnia via gas challenge and rebreathing have shown both decreased (McKeton et al., 2018; Peng et al., 2018) and increased (Tomoto et al., 2019; Zhu et al., 2013) CVR with aging. Apparent “improvements” in CBF response may, however, reflect a rightward shift in the reactivity curve prompted by general decreases in resting CBF with aging (Zhu et al., 2013). Notably, a number of studies have failed to detect any CBF changes in response to CO₂ manipulation with aging. Null findings in studies of age differences in CVR may be due to differential effects of hypercapnic stimuli in older versus younger adults (Ances et al., 2009), underscoring the importance of capnographic monitoring during CVR experiments. Nevertheless, the pattern of mixed findings from prior studies leaves open the question of whether aging itself may impact CVR, even in cognitively unimpaired older adults.

It is also unclear whether aging may differentially impact vasodilatory versus vasoconstrictive responses, as the overwhelming majority of reactivity studies have focused on hypercapnia exclusively (Liu et al., 2019). In the few studies that examined age-related changes in hypocapnic response, there has been a similarly mixed pattern of findings. For example, TCD investigations have variously found decreased (Tomoto et al., 2019; Zhu et al., 2013), increased (Stefanidis et al., 2019), and equivalent (Minhas et al., 2019) CBF response to hyperventilation-induced hypocapnia in older adults, relative to younger adult controls.

The literature on age-related changes in hypocapnia responses is also methodologically limited, as most studies have relied on TCD to measure CBF. Although this approach offers excellent temporal precision, it lacks the spatial resolution of MRI-based methods and is limited to assessment of flow through larger intracranial arteries in neuroanatomically superficial regions (van Beek et al., 2008). Arterial spin labeling MRI provides a measure directly proportional to cerebral perfusion in every voxel in the brain and offers distinct information to complement findings from TCD or BOLD fMRI studies (Liu et al., 2019). Given that many neurovascular changes underpinning age-related declines in cerebrovascular autoregulation appear to occur in smaller vascular compartments (Iadecola, 2004), pseudo-continuous arterial spin labeling (pCASL) perfusion MRI may be particularly adept at assessing age-related changes in cerebrovascular function.

The present study evaluated age-related differences in CVR to gain further insight into the potential for CVR as a preclinical marker. CVR comparisons were made between older adults screened for cognitive impairment and healthy younger adults to address whether age-related CVR changes are detectible even in cognitively unimpaired older adults. CVR experiments utilized pCASL MRI to observe cerebral perfusion responses to both hypocapnia and hypercapnia in order to compare age-related changes in vasoconstrictive and vasodilatory responses. Capnography was employed during all experiments to ensure compliance with breathing manipulations and equivalence of end-tidal CO₂ (etCO₂) responses across age groups, and to calculate CVR maps indicative of perfusion changes per unit changes in etCO₂. Regional CVR values were examined in cortical and limbic regions previously shown to have age-related changes in resting CBF and susceptibility to dementia (Zhang et al., 2017).

2. Material and methods

2.1. Participants

Participants were drawn from the Vascular Senescence and Cognition (VaSC) Study at the University of Southern California (USC). The VaSC Study enrolled individuals from the community via word-of-mouth, flyers, and hosted events through the USC School of Gerontology. Participants provided informed consent to the VaSC study protocol, approved by the USC Institutional Review Board, and were financially compensated for their participation. Exclusion criteria included diagnosis of cognitive impairment or dementia, known family history of a genetic mutation that produces dementia, insulin-dependent diabetes, MRI contraindication(s) (e.g., weight exceeding 270 pounds), current organ failure or major systemic illness, history of stroke, myocardial infarction, major neurologic or psychiatric disorder impacting cognition, head injury with loss of consciousness exceeding 15 minutes, substance abuse resulting in hospitalization, B12 deficiency, and/or hypothyroidism. Based on self-reported medical history, no individual in the current sample had a diagnosis of vascular or pulmonary diseases known to have a major impact on CVR (e.g., carotid artery stenosis, Moyamoya angiopathy). One participant had a history of pulmonary embolism but was able to complete breathing exercises, with their average resting etCO₂ within normal range (40.20 mmHg).

2.2. Neuropsychological screening

Older adults underwent comprehensive neuropsychological testing to screen for mild cognitive impairment by sensitive neuropsychological criteria (Bondi et al., 2014). Briefly, participants were excluded if they exhibited impaired performance on two or more tests within a cognitive domain or three or more tests across cognitive domains. Thus, all participants were determined to be cognitively unimpaired at the time of the evaluation. Specifically, the following cognitive domains and tests were considered for diagnosis of mild cognitive impairment (Bondi et al., 2014): Attention and executive function (Trail Making Test: A and B), memory (Rey Auditory Verbal Learning Test: Long Delay Recall and Recognition), and language (Verbal Fluency Test: Animals and Boston Naming Test/Multilingual Naming Test).

2.3. Breathing paradigms

Participants completed paced breathing and breath hold exercises during separate pCASL acquisitions. All participants engaged in a training protocol for breathing exercises before entering the scanner. Each breathing exercise was preceded by verbal and written instructions. Participants were also provided with in-scanner visual stimuli to guide breath control and ensure accurate completion of each exercise.

2.3.1. Paced breathing (0.1 Hz): hypocapnia

Participants were presented with a circle that filled with color over a 5-second interval. The color of the filling circle alternated between yellow and blue. Participants were instructed to “inhale while the circle is yellow” and “exhale while the circle is blue.” Perfusion images were acquired continuously throughout the 0.1 Hz paced breathing modulation at two exhale-inhale cycle intervals (Fig. 1).

2.3.2. Breath hold: hypercapnia

Participants were shown a circle that filled with color (green) over a 25-second interval. Once full, the circle “restarted” and began filling with another color (red) over a 15-second interval. Participants were instructed to, “breathe normally while the circle is green” and “hold your breath while the circle is red.” Participants were also instructed to exhale prior to each breath hold to equate initial lung volume across participants and to maximize hypercapnic effect given the breath hold duration (Unback et al., 2017). At the end of each breath hold, participants were instructed to exhale between each breath, filling the circle alternated between yellow and blue. Participants were instructed to “inhale while the circle is yellow” and “exhale while the circle is blue.” Perfusion images were acquired continuously throughout the 0.1 Hz paced breathing modulation at two exhale-inhale cycle intervals (Fig. 1).

2.3.2. Breath hold: hypercapnia

Participants were shown a circle that filled with color (green) over a 25-second interval. Once full, the circle “restarted” and began filling with another color (red) over a 15-second interval. Participants were instructed to, “breathe normally while the circle is green” and “hold your breath while the circle is red.” Participants were also instructed to exhale prior to each breath hold to equate initial lung volume across participants and to maximize hypercapnic effect given the breath hold duration (Unback et al., 2017). At the end of each breath hold, participants were instructed to exhale
in order to synchronize capnographic sampling across participants and to measure etCO$_2$ approximating increased CO$_2$ level induced by breath hold (Murphy et al., 2011). Each cycle of normal breathing and breath hold was synchronized with pCASL acquisition such that cerebral perfusion sampling occurred at the end of normal breathing period/beginning of breath hold and at 5s after the participant stopped the breath hold and was breathing normally again (i.e., during peak perfusion response) (Fig. 1).

2.4. Neuroimaging and physiological measures

All MRI scans were conducted on a 3T scanner (Siemens Magnetom Prisma System) using a 20-channel head coil at the USC Dana and David Dornsife Cognitive Neuroimaging Center. Anatomical images were collected using a 3D high-resolution T1-weighted magnetization-prepared rapid gradient-echo (MPRAGE) scan (TR = 2300 ms; TE = 2.98 ms; TI = 900 ms; resolution = 1.0 × 1.0 × 1.2 mm$^3$). Cerebral perfusion was measured by a 3D gradient and spin-echo (GRASE) pCASL scan with background suppression and the following sequence parameters: TR = 5000 ms; TE = 36.3 ms; FOV = 240 mm; resolution = 2.5 × 2.5 × 3.4 mm$^3$; slice thickness = 3.42 mm; number of slices = 24; labeling duration = 1517 ms; post-labeling delay = 2000 ms; 32 total acquisitions (1 M0 image + 1 dummy image + 30 alternating tag and control images); total scan time = 5 minutes 25 seconds. Altogether, each pCASL scan for 0.1 Hz paced breathing and breath hold yielded 15 tag-control pair images.

The pCASL scans were pre-processed using the ASLtbx pipeline, implemented in SPM12 within MATLAB (Wang et al., 2008; Wang, 2012). Pre-processing steps for pCASL scans included motion correction, co-registration to individual subject’s structural T1-weighted image, spatial smoothing with a 6 mm full-width at half-maximum Gaussian kernel, and tag-control subtraction resulting in 15 tag-control pairs for each subject with values for absolute CBF (mL/100g tissue/min). All CBF images were thresholded below 10 or above 150mL/100g/min to exclude CBF outside the expected physiological range of gray matter (Nation et al., 2013; Clark et al., 2021). Tag-control pairs were warped to MNI space and averaged to create mean CBF maps for each subject. Resulting mean CBF maps were visually inspected for quality and gross abnormalities (i.e., large signal dropout). Partial volume correction was performed by applying subject-specific gray matter masks derived from the gray matter tissue class segmentation of T1-weighted structural images (Petr et al., 2018). Segmented gray matter maps were thresholded at 0.3, binarized, and multiplied by the mean CBF maps to ensure CBF was limited to gray matter.

Capnography indexed etCO$_2$ during MRI acquisition using an M3015A sidestream carbon dioxide extension module (Philips Medical Systems) connected to a nasal cannula into which participants breathed. To correct for sampling tubing latency, etCO$_2$ time series were shifted by a pre-calibrated duration of time (i.e., 10 seconds in our set-up). For 0.1 Hz paced breathing exercise (hypocapnia), etCO$_2$ was extracted from the raw time series in accordance with the breathing rate (i.e., at every expiration). For breath hold exercise (hypercapnia), maximum etCO$_2$ was sampled for each pCASL image (i.e., maximum positive peak across the acquisition interval for each tag-control pair) from the raw time series, to effectively capture increases in etCO$_2$ induced by breath hold. For each breathing paradigm, participants were excluded from analysis if they failed to adhere to breathing instructions (e.g., lack of positive peaks in raw data – participant might be breathing through the mouth).

2.5. CVR maps

CVR was conceptualized as the percent change in CBF per unit change in etCO$_2$ (Liu et al., 2019; Marshall et al., 2014). Specific computational approach was adapted from the method used by Marshall et al. (2014), which evaluated CVR in hypercapnia via CO$_2$ gas administration. In the present study, participant’s individual CVR maps were generated for each breathing paradigm by calculating the following (1) in every voxel:

$$\text{CVR (}\%\text{CBF change/mm Hg)} = \frac{100 \times (\text{CBF}_{\text{maximum}} - \text{CBF}_{\text{minimum}})}{\text{etCO}_2\text{maximum} - \text{etCO}_2\text{minimum}}$$ (1)

Regional mean CVR values were extracted for our regions of interest (ROI), including inferior frontal gyrus, orbitofrontal cortex, anterior cingulate cortex, amygdala, hippocampus, parahippocampal gyrus, entorhinal cortex, perirhinal cortex, medial temporal lobe, inferior temporal gyrus, inferior parietal cortex, posterior cingulate cortex, precuneus, caudate, and thalamus (Zhang et al., 2017).

2.6. Statistical analyses

Analyses were performed using SAS 9.4 (Cary, NC). Age (young vs. old) comparisons of etCO$_2$ responses (i.e., etCO$_2\text{maximum} - \text{etCO}_2\text{minimum}$) for each breathing paradigm were performed using independent samples t-tests. Chi-square tests were conducted to compare sex distribution between younger and older adults. Analysis of covariance (ANCOVA) models tested the difference between younger and older adults in their regional CVR values, adjusting for sex. The Benjamini-Hochberg method was used to correct for multiple comparisons, with false discovery rate controlled at $p < 0.05$.

3. Results

3.1. Sample characteristics and effects of breathing manipulations

The overall sample consisted of 30 younger (age 18–28) and 31 older (age 55–87) adults. From this overall sample, participants successfully completing at least one of the breathing exercises were included in the analysis. For 0.1 Hz paced breathing (hypocapnia), participants were excluded due to non-adherence to breathing instructions ($n = 2$), no pCASL scan ($n = 3$), and physio equipment failure ($n = 1$). Thus, in the final analytic sample, 29
younger (mean age = 21.79, SD = 2.64; 27.6% female) and 26 older (mean age = 70.27, SD = 8.61; 26.9% female) adults were included. For breath hold (hypocapnia), participants were excluded due to non-adherence to breathing instructions (n = 5) and physio equipment failure (n = 1). Thus, in the final analytic sample, 26 younger (mean age = 22.00, SD = 2.67; 30.8% female) and 29 older (mean age = 69.62, SD = 8.38; 27.6% female) adults were included. Chi-square tests revealed no difference in sex distributions between younger and older adults in hypocapnia ($\chi^2 = 0.00, p = 0.96$) or hypercapnia ($\chi^2 = 0.07, p = 0.80$) conditions.

Older adults did not differ from younger adults in their etCO2 responses to either 0.1 Hz paced breathing (hypocapnia: younger adults mean = 6.07 mmHg, SD = 2.57; older adults mean = 6.16 mmHg, SD = 2.29; t = 0.14, p = 0.89) or breath hold (hypocapnia: younger adults mean = 9.36 mmHg, SD = 4.23; older adults mean = 10.43 mmHg, SD = 2.11; t = 1.21, p = 0.23) Fig. 2. shows representative raw capnography recordings and etCO2 time series data for 0.1 Hz paced breathing (hypocapnia) and breath hold (hypercapnia) exercises.

3.2. Age differences in CVR

Visual comparison of mean CVR maps of younger and older adults in 0.1 Hz paced breathing (hypocapnia) and breath hold (hypercapnia) conditions indicate attenuated CVR responses in older adults relative to younger adults (Fig. 3). Global CVR was attenuated in older adults during 0.1 Hz paced breathing (hypocapnia) and breath hold (hypercapnia) relative to younger adults (Table 1). Older adults demonstrated reduced CVR to hypocapnia in the orbitofrontal cortex, anterior cingulate cortex, and inferior temporal gyrus relative to younger adults, as well as reduced CVR to hypercapnia in all regions examined (Table 1).

4. Discussion

The present study found that cognitively unimpaired older adults exhibit attenuated whole-brain CVR in response to both hypocapnia and hypercapnia relative to younger adults. These findings are consistent with some prior studies utilizing a variety of methodologies to characterize age group differences in CVR (Mcketton et al., 2018; Peng et al., 2018), but are contrary to other studies suggesting either no age group difference (Ances et al., 2009) or an increase in CVR in older adults (Tomoto et al., 2019; Zhu et al., 2013). Several methodological differences across studies, including CVR paradigm, method of perfusion estimation, and monitoring and incorporation of etCO2, may contribute to variability across studies. For the present study, examination of both hypocapnia and hypercapnia, utilization of pCASL-MRI markers of cerebral perfusion, and calculation of CVR normalized to etCO2, all support clear interpretation of the findings indicating attenuated cerebrovascular function with aging.

The majority of prior efforts have examined vasmotor effects of hypercapnia, while few have focused on hypocapnia. For MRI-based studies of CVR, non-invasive induction of hypocapnia through hyperventilation can cause participant discomfort and movement artifacts that obscure interpretation of experimental results (Pinto et al., 2021). The present study’s novel use of a paced breathing paradigm reduced etCO2 and consequently induced vasocostrictive CVR responses with minimal reported participant discomfort or observed motion artifact. Utilization of short 15s interval breath holds similarly allowed for increased etCO2 levels with minimal reported discomfort in our older adult participants. Examination of CVR to both hypocapnia and hypercapnia allowed for comparison of vasoconstrictive and vasodilatory CVR deficits in older adults in the current study. Findings indicated a clear dif-
Fig. 3. Average whole-brain gray matter cerebrovascular reactivity (CVR) maps contrasting younger and older adults in 0.1 Hz paced breathing (top: hypocapnia) and breath hold (bottom: hypercapnia) conditions. CVR was computed as the percent change in CBF per mmHg change in etCO₂. Warm colors indicate higher CVR values, while cold colors show lower CVR values. It is easily noticeable that older adults demonstrated attenuated CVR in both conditions, compared with younger adults. Abbreviations: CBF, cerebral blood flow; etCO₂, end-tidal CO₂.
ference in the magnitude and anatomical distribution of aging effects. Older adults displayed significant global CVR deficits in vasocostric (hypocapnic stimuli), with specific regional vasocostric deficits in the frontal and inferior temporal regions. Age-related CVR deficits in vasodilatation (hypercapnic stimuli) were of large effect size and showed statistically significant effects in both the global CVR analysis and in all ROIs, including frontal, temporal, parietal, and limbic regions. These findings suggest that vasodilatory capacity may be disproportionately impacted by aging, with more subtle and circumscribed changes in vasocostric capacity also being detectible. This pattern of age-related CVR differences is consistent with a net vasocostric phenotype suggested by studies of age-related changes in the cerebrovasculature (Yew and Nation, 2017; Kislcr et al., 2017). It is possible these age-related changes in vasocostric versus vasocostric capacities may convey vulnerability to cerebral hypoperfusion, but further longitudinal studies are needed.

Deficits in CVR to hypercapnia have been linked to cognitive impairment (Canet al., 2011; Mcketten el al., 2018; Peng al., 2018; Sur et al., 2020) and dementia risk (Silvestrini et al., 2006). However, few studies have focused on cognitively unimpaired older adults, confirmed by comprehensive neuropsychological testing. A key goal of CVR research is to develop preclinical markers of vascular brain injury and risk for future cognitive decline, but achieving this goal will require improved understanding of age-related changes in CVR. Thus, the present study findings add to the existing literature on age-related CVR changes by establishing CVR deficits in cognitively unimpaired older adults relative to younger adults. Interestingly, very large aging effects were observed in response to hypercapnic stimuli despite the fact that participants were cognitively unimpaired. These findings suggest that decline in vasodilatory reserve may begin to attenuate at an earlier age than the age range of the present study (55–87 years). This is consistent with one of the few longitudinal studies of CVR change over the lifespan, which found that vasodilatory CVR declined most rapidly during midlife (Peng et al., 2018). This is also consistent with prior studies suggesting the importance of midlife (vs. late-life) vascular risk factors with respect to neurocognitive decline (Armstrong et al., 2019).

To our knowledge, no study to date has examined the longitudinal trajectory of vasocostric CVR. Although the present study is cross-sectional, our findings provide further insight into age-related changes in vasocostric capacity, as our comparisons of younger versus older adults detected milder, anatomically circumscribed age-related differences in vasocostric responses than those of the more well-studied vasodilatory responses. These findings suggest that vasocostric responses may be more preserved than vasodilatory responses into late life, or that significant declines in vasocostric responses may only occur in the context of cerebrovascular disease. Evidently, more data from longitudinal and clinical studies are needed to confirm these hypotheses. Yet, the present study underscores a value in evaluating dynamic changes in vasocostric responses in the older adult population, where vasodilatory changes may already be substantial.

The ROI analysis in the present study indicated widespread age group differences in vasodilatory response in all cortical, limbic, and subcortical regions with no clear anatomical pattern. In contrast, age group differences in vasocostric response involved specific frontal and temporal regions, including the orbitalfrontal, anterior cingulate, and inferior temporal cortices. It remains unclear why these particular regions would show age group differences in vasocostric to hypcapnia. However, the temporal lobe finding is consistent with prior work indicating that temporal regions are the earliest to show declines in CVR (Peng et al., 2018). Age-related declines in CVR to hypcapnia in frontal and anterior limbic regions may explain relatively increased perfusion found in these anterior regions in cognitively normal older adults (Lee et al., 2009). Researchers have posited that the frontal lobes and associated circuitry are particularly vulnerable to the effects of aging (“frontal lobe hypothesis”; Cabeza and Dennis, 2013), to explain functional deficits mediated by the prefrontal regions and compensatory mechanisms. However, our findings also indicated age-related deficits in temporal regions. Longitudinal studies of CVR to hypcapnia will provide further clarity regarding the spatiotemporal pattern of age-related changes in cerebral vasocostric. Moreover, future studies may investigate into possible hemispheric differences in CVR, about which little is known in the literature.

Study strengths include investigation of a well-tolerated, non-invasive approach to indexing relative cerebrovascular responses to both hypcapnia and hypercapnia in older and younger adults, correction for etCO₂ changes, and focus on a rigorously screened sam-

Table 1

| HYPOCAPNIA | YOUNGER | OLDER |
|-----------|---------|-------|
| **Whole brain** | 14.48 (1.41) | 16.90 (2.45) |
| **IFG** | 19.16 (2.77) | 20.60 (2.77) |
| **OFC** | 26.64 (3.35) | 28.20 (3.35) |
| **ACC** | 21.85 (2.37) | 23.45 (2.37) |
| **Amygdala** | 17.47 (3.99) | 19.05 (3.99) |
| **Hippocampus** | 13.61 (2.50) | 15.15 (2.50) |
| **PHG** | 13.69 (2.59) | 15.20 (2.59) |
| **EC** | 9.43 (2.36) | 11.12 (2.36) |
| **PCG** | 10.95 (2.40) | 12.56 (2.40) |
| **Precuneus** | 11.48 (2.64) | 13.12 (2.64) |
| **Caudate** | 20.79 (4.38) | 22.32 (4.38) |
| **Thalamus** | 18.31 (4.29) | 20.23 (4.29) |

**HYPERCAPNIA**

| YOUNGER | OLDER |
|---------|-------|
| **Whole brain** | 17.78 (1.41) | 19.30 (1.41) |
| **IFG** | 26.02 (2.08) | 27.58 (2.08) |
| **OFC** | 27.75 (2.58) | 29.35 (2.58) |
| **ACC** | 26.64 (2.42) | 28.20 (2.42) |
| **Amygdala** | 32.65 (3.46) | 34.20 (3.46) |
| **Hippocampus** | 25.89 (2.03) | 27.45 (2.03) |
| **PHG** | 24.76 (1.85) | 26.30 (1.85) |
| **EC** | 23.70 (2.11) | 25.25 (2.11) |
| **PCG** | 20.68 (1.88) | 22.23 (1.88) |
| **Precuneus** | 21.88 (1.68) | 23.43 (1.68) |
| **Caudate** | 19.10 (2.23) | 20.65 (2.23) |
| **Thalamus** | 21.74 (2.32) | 23.20 (2.32) |

Key: ACC, anterior cingulate cortex; CVR, cerebrovascular reactivity; EC, entorhinal cortex; IFG, inferior frontal gyrus; IPC, inferior parietal cortex; IFG, inferior temporal gyrus; MTL, medial temporal lobe; OFC, orbitofrontal cortex; PCG, posterior cingulate cortex; PHG, parahippocampal gyrus; PRB, perirhinal cortex.

Values in bold indicate statistically significant results at false discovery rate (FDR) p < 0.05.
ple of cognitively unimpaired older adults. Study limitations include the relatively small sample size. The use of pCASL acquisition represents a strength, in that pCASL captures perfusion changes in response to CO₂ levels more directly than BOLD-fMRI signals, which are influenced by other factors including cerebral blood volume and metabolic rate of oxygen consumption (Liu et al., 2019). However, the low signal-to-noise ratio of individual tag-control pairs, low temporal resolution, and limited ability to assess white matter perfusion represent widely recognized limitations to pCASL studies. Tradeoffs exist in terms of the strengths and limitations of any given approach to CVR studies, which warrants the continued investigation of multiple alternative approaches that may yield different information about cerebrovascular function in the older adult population, such as investigation of perfusion and CVR in the white matter. Additionally, our methods were able to detect patterns of age-related differences in CVR to hypocapnia and hypercapnia despite limitations inherent to pCASL methodology, such as differences in labeling efficiency during hypocapnia and hypercapnia (Aslan et al., 2010).

In sum, our findings and methodological approach present important clinical benefits. Firstly, the disproportionate pattern of results contrasting age-related CVR deficits in vasodilatation versus vasoconstriction provides implications for models of age-related diseases, such as Alzheimer’s disease, which has been characterized by researchers as a “hypercontractile” phenotype (Kim et al., 2013; Kisler et al., 2017). Secondly, utilization of CVR as a clinical biomarker of disease in older populations benefits from more refined understanding of age-related changes in cerebral hemodynamics. For instance, vasoconstrictive CVR might be expected to be normal if an older patient is presumed healthy, but the same may not hold true of vasodilatory CVR. Lastly, the present study provides a paradigm for clinical studies that is well-tolerated by older adults and allows evaluation of both vasodilatory and vasoconstrictive capacity, particularly under circumstances where gas inhalation is contraindicated or unavailable.

5. Conclusions

Our findings indicate that cognitively unimpaired older adults exhibit attenuated CVR to hypocapnia and hypercapnia, compared with younger adults. Further, the effects of age-related CVR deficits are greater for hypercapnia across a range of cortical, limbic, and subcortical regions than for hypocapnia, circumscribed to specific frontal and temporal regions. Aging may have a disproportionate impact on vasodilatory versus vasoconstrictive CVR, and this pattern of changes may support the utility of CVR assessment in estimating the risk for cerebrovascular injury and cognitive impairment in older adults. Further, considerations of aging effects may facilitate better characterization of CVR changes as a predictive and treatment-monitoring tool for clinical populations, such as individuals with reduced vascular reserve (e.g., Moyamoya angiopathy).

Submission declaration and verification

All authors have reviewed and approved the submission of this manuscript. This manuscript or any of its elements has not been published previously and is not under consideration for publication elsewhere. If accepted, the manuscript will not be published elsewhere in the same form, in English or in any other language, including electronically, without the written consent of the copyright-holder.

Disclosure statement

The authors declare that there is no conflict of interest.

CRediT authorship contribution statement

Belinda Yew: Conceptualization, Formal analysis, Writing – original draft, Visualization. Jung Yun Jung: Conceptualization, Formal analysis, Writing – original draft, Visualization. Shubir Dutt: Conceptualization, Formal analysis, Writing – original draft, Visualization. Yanrong Li: Data curation, Project administration, Writing – review & editing. Isabel J. Sible: Writing – review & editing. Aimée Gaubert: Writing – review & editing. Jean K. Ho: Writing – review & editing. Anna E. Blanken: Writing – review & editing. Anisa Marshall: Writing – review & editing. Xingfeng Shao: Writing – review & editing. Danny J.J. Wang: Writing – review & editing. Daniel A. Nation: Conceptualization, Formal analysis, Funding acquisition, Supervision, Writing – review & editing, Visualization.

Acknowledgements

This work was supported by the National Institutes of Health (grant numbers R01AG064228, R01AG060049, P50AG16573, P01AG052350); the National Science Foundation (grant number DGE1418060); and the Alzheimer’s Association (grant number AARG-17-532905).

References

Ance, B.M., Liang, C.L., Leonstev, O., Perthien, J.E., Fleisher, A.S., Lansing, A.E., Buxton, R.B., 2009. Effects of aging on cerebral blood flow, oxygen metabolism, and blood oxygenation level-dependent responses to visual stimulation. Hum. Brain. Mapp. 30, 1120–1132.

Armstrong, N.M., Bangen, K.J., Au, R., Gross, A.L., 2019. Associations between midlife (but not late-life) elevated coronary heart disease risk and lower cognitive performance: results from the Framingham Offspring Study. Am. J. Epidemiol. 188, 2175–2187.

Aranvistake, Z., Capuano, A.W., Leurgans, S.E., Bennett, D.A., Schneider, J.A., 2016. Relation of cerebral vessel disease to Alzheimer’s disease dementia and cognitive function in elderly people: a cross-sectional study. Lancet Neurol 15, 934–943.

Aslan, S., Xu, F., Wang, P.L., Uh, J., Yezbouch, U.S., van Osch, M., Lu, H., 2010. Estimation of labeling efficiency in pseudocontinuous arterial spin labeling. Magn. Reson Med 63, 765–771.

Bondi, M.W., Edmonds, E.C., Jak, A.J., Clark, L.R., Delano-Wood, L., McDonald, C.R., Nation, D.A., Libon, D.J.; Au, R.; Galasko, D.; Salmon, D.F., 2014. Neuropsychological criteria for mild cognitive impairment improves diagnostic precision, biomarker associations, and progression rates. J. Alzheimer’s Dis. 42, 275–289.

Cabeza, R., Dennies, N.A., 2013. Frontal lobes and aging: deterioration and compensation. In: Principles of Frontal Lobe Function. Oxford University Press, New York, pp. 628–652.

Cantin, S., Villien, M., Moreaud, O., Tropres, I., Keignart, S., Chipon, E., Le Bas, J.F., Warnking, J., Kainik, A., 2011. Impaired cerebral vasoreactivity to CO₂ in Alzheimer’s disease using BOLD fMRI. NeuroImage 58, 579–587.

Clark, A.L., Weigand, A.J., Bangen, K.J., Merritt, V.C., Bondi, M.W., Delano-Wood, L., 2021. Repetitive mTBI is associated with age-related reductions in cerebral blood flow but not cortical thickness. J. Cereb. Blood Flow Metab. 41, 431–444.

Han, J.S., Abou-Hamed, A., Mandell, D.M., Poublanc, J., Crawley, A.P., Fisher, J.A., Mikulis, D.J., Tynianni, M., 2011. Impact of extracranial-intracranial bypass on cerebrovascular reactivity and clinical outcome in patients with symptomatic moyamoya vasculopathy. Stroke 42, 3047–3054.

Iadecola, C., 2004. Neurovascular regulation in the normal brain and in Alzheimer’s disease. Nat. Rev. Neurosci. 5, 347–360.

Kim, S.M., Kim, M.J., Rhee, H.Y., Ryu, C-W., Kim, E.J., Petersen, E.T., Jahng, G-H., 2013. Regional cerebral perfusion in patients with Alzheimer’s disease and mild cognitive impairment: Effect of APOE Epsilon4 allele. Neuropsychology 55, 25–34.

Kisler, K., Nelson, A.R., Montagne, A., Zlokovic, B.V., 2017. Cerebral blood flow regulation and neurovascular dysfunction in Alzheimer disease. Nat. Rev. Neurosci. 18, 419–434.

Lee, C., Lopez, O.L., Becker, J.T., Raji, C., Dai, W., Muller, L.H., Gach, H.M., 2009. Imaging cerebral blood flow in the cognitively normal aging brain with arterial spin labeling: implications for imaging of neurodegenerative disease. J. Neuroimag. 19, 344–352.

Liu, P., De Vis, J.B., Lu, H., 2019. Cerebrovascular reactivity (CVR) MRI with O₂ challenge: a technical review. NeuroImage 187, 104–115.

Markus, H., Cullinane, M., 2001. Severely impaired cerebrovascular reactivity predicts stroke and TIA risk in patients with carotid artery stenosis and occlusion. Brain 124, 457–467.

Marshall, O., Lu, H., Brisset, J.C., Xu, F., Liu, P., Herbert, J., Grossman, R.J., Ge, Y., 2014. Impaired cerebrovascular reactivity in multiple sclerosis. JAMA Neurol 71, 1275–1281.
