Aerobic fitness is associated with greater hippocampal cerebral blood flow in children

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ARTICLE INFO

Article history:
Received 26 August 2015
Received in revised form 30 June 2016
Accepted 1 July 2016
Available online 4 July 2016

Keywords:
Arterial spin labeling
Childhood
Development
Hippocampus
Perfusion
Physical activity

ABSTRACT

The present study is the first to investigate whether cerebral blood flow in the hippocampus relates to aerobic fitness in children. In particular, we used arterial spin labeling (ASL) perfusion MRI to provide a quantitative measure of blood flow in the hippocampus in 73 7- to 9-year-old preadolescent children. Indeed, aerobic fitness was found to relate to greater perfusion in the hippocampus, independent of age, sex, and hippocampal volume. Such results suggest improved microcirculation and cerebral vasculature in preadolescent children with higher levels of aerobic fitness. Further, aerobic fitness may influence how the brain regulates its metabolic demands via blood flow in a region of the brain important for learning and memory. To add specificity to the relationship of fitness to the hippocampus, we demonstrate no significant association between aerobic fitness and cerebral blood flow in the brainstem. Our results reinforce the importance of aerobic fitness during a critical period of child development.

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1. Introduction

Participation in physical activity and higher levels of aerobic fitness are associated with superior scholastic achievement, cognitive control, and memory in children (Buck et al., 2008; Castelli et al., 2007; Chaddock et al., 2010a,b, 2011; Chomitz et al., 2009; Hillman et al., 2009; Pontifex et al., 2011; Voss et al., 2011). Still, little is known about the neural mechanisms by which aerobic fitness influences the developing brain during childhood. Volumetric and functional magnetic resonance imaging (fMRI) techniques provide some clues, such that higher fit children show larger brain volumes in the hippocampus and basal ganglia (Chaddock et al., 2010a,b), as well as differences in blood oxygenation level dependent (BOLD) fMRI brain activation in areas of frontal and parietal cortex (Chaddock et al., 2012; Voss et al., 2011), relative to their lower fit peers. Non-human animal work raises the possibility that, mechanistically, children with higher levels of aerobic fitness may have increased growth and expansion of neural tissue and/or increased vasculature (see Voss et al., 2013 for a review).

The present study is the first to investigate whether increased cerebral blood flow (CBF) in the hippocampus is associated with aerobic fitness during childhood. This hypothesis cannot be directly tested with traditional BOLD techniques, given that BOLD can change depending on a number of factors related to local metabolism and neural function, including blood volume, perfusion, blood velocity, and cerebral metabolic rate of oxygen (Hoge et al., 1999). Hence, here we use arterial spin labeling (ASL) perfusion MRI to provide a quantitative measure of blood flow and a more direct link to local neuronal activity (Alsop and Detre, 1996). Specifically, an ASL signal arises from the delivery of magnetically tagged arterial water into an imaging slice of interest, where the blood water exchanges in the tissue. The output measure of CBF, or the blood supply to a brain area in a given time (mL/100 g/min), is known to provide information regarding how the brain meets and regulates its metabolic demands via the delivery of metabolites, oxygen and nutrients to activated neurons (Hales et al., 2014; Sokoloff et al., 1977) (see ASL Scan Acquisition section in the Method for more information about ASL).

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http://dx.doi.org/10.1016/j.dcn.2016.07.001
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We specifically focused on CBF in the hippocampus, in view of converging evidence that demonstrates positive physical activity-related brain changes in the hippocampus in rodents and humans across the lifespan (Bugg and Head, 2011; Burdette et al., 2010; Chaddock et al., 2010a; Erickson et al., 2009, 2011; Honig et al., 2009; Pereira et al., 2007; van Praag et al., 1999b). For example, voluntary wheel running in rodents has been found to enhance learning and memory (van Praag et al., 2005) as well as induce angiogenesis and increased vascular density (Black et al., 1990; Clark et al., 2009; Kleim et al., 2002; Rhyu et al., 2010), and the growth of new neurons in the hippocampus (van Praag et al., 1999a). In humans, physical activity and aerobic fitness are associated with a greater number of small-caliber vessels (Bullitt et al., 2009), increased cerebral blood volume in the hippocampus in middle-aged adults (Pereira et al., 2007; age 21–45) and increased hippocampal blood flow in older adults (Burdette et al., 2010).

Cognitively, increased hippocampal CBF has been linked to higher task performance on a spatial memory task in middle-aged and older adults (Heo et al., 2010; Pereira et al., 2007). It is possible that these findings extend to children, such that aerobic fitness relates to greater perfusion in the hippocampus, which may suggest improved microcirculation, cerebral vasculature, and function. We hypothesized that aerobic fitness in 7- to 9-year-old preadolescent children would be associated with increased resting CBF in the hippocampus. We explored anterior and posterior subsections of the hippocampus to examine whether aerobic fitness had selective effects on hippocampal blood flow, and to investigate distinct contributions of different anatomical regions within the hippocampus given that functional distinctions have been described along the anterior/posterior axis of the hippocampus (i.e., spatial versus relational processing) (Giovanello et al., 2009; Sperling et al., 2003). To provide additional specificity to a fitness–CBF relationship, we also measured CBF in the brainstem as a control region. Like the hippocampus, the brainstem is a subcortical structure in the midbrain included in our ASL slice acquisition, yet this region has not been found to relate to aerobic fitness. Thus, we did not predict an association between aerobic fitness and brainstem CBF. Given these hypotheses, the present study will provide insight into a potential cerebrovascular mechanism by which aerobic fitness enhances brain health in children.

91 children were eligible for the study. Eighteen children were excluded from the analysis due to excessive head motion during the ASL scan. Seventy-three children (41 girls, 32 boys), ages 7–9 years (M = 8.63 years, SD = 0.54) were included in the ASL analysis.

### 2.2. Aerobic fitness testing

Children completed a VO2max test to assess aerobic fitness. The aerobic fitness of each child was measured as maximal oxygen consumption (VO2max) during a graded exercise test (GXT). The GXT employed a modified Balke Protocol and was administered on a LifeFitness 92T motor-driven treadmill (LifeFitness, Schiller Park, IL) with expired gases analyzed using a TrueOne2400 Metabolic Measurement System (ParMedics, Sandy, Utah). Children walked and/or ran on a treadmill at a constant speed with increasing grade increments of 2.5% every 2 min until volitional exhaustion occurred.

Oxygen consumption was measured using a computerized indirect calorimetry system (ParvoMedics True Max 2400) with averages for VO2 and respiratory exchange ratio (RER) assessed every 20 s. A polar heart rate (HR) monitor (Polar WearLink+ 31; Polar Electro, Finland) was used to measure HR throughout the test, and ratings of perceived exertion (RPE) were assessed every 2 min using the children’s OMNI scale (Utter et al., 2002). Maximal oxygen consumption was expressed in mL/100 g/min and VO2max was based upon maximal effort as evidenced by (1) a plateau in oxygen consumption corresponding to an increase of less than 2 mL/kg/min despite an increase in workload; (2) a peak HR > 185 beats per minute (American College of Sports Medicine, 2014) and an HR plateau (Freedson and Goodman, 1993); (3) RER > 1.0 (Bar-Or, 1983); and/or (4) a score on the children’s OMNI ratings of perceived exertion (RPE) scale ≥ 8 (Utter et al., 2002). Our sample consisted of relatively lower fit children (average VO2max percentile = 33.5%).

### 2.3. Arterial spin labeling (ASL) scan acquisition

Quantitative resting CBF in the child sample was measured using multi-slice pseudo-continuous arterial spin labeling (pCASL) (Wu et al., 2007). A number of studies have reported successful use of the ASL technique in children (Helton et al., 2009; Thomason et al., 2009; van den Tweel et al., 2009; Wang et al., 2003). In general, during an ASL scan, one or more radiofrequency (RF) pulses excite water molecules in arterial blood water in upstream blood (i.e., below the slice or region of interest), thus “labeling” or “tagging” the blood. Following a period of time, the labeled blood enters the imaging plane and alters the signal in the image, which is referred to as the “tag image.” Then, this acquisition is repeated, without the RF labeling, and a “control image” is created without the added signal contribution from tagged arterial blood. The difference between the tag image and the control image is the perfusion image, which reflects only blood flow, or CBF. Therefore, the perfusion image quantifies the amount of arterial blood delivered to each voxel in the slice within the post-label delay. This is affected by the arterial transit time (ATT), which is the time it takes for blood to travel from labeling plane to image voxel.

During the pCASL scan, the slices were oriented axially, perpendicular to the vertebral arteries. The top slice was positioned superior to the corpus callosum so that the slices covered the temporal lobes of the brain, including the hippocampus. Prior to acquisition, shimming was performed over a region that extended from the imaging slices to the tagging plane. The acquisition parameters included: multi-slice, gradient-echo echoplanar imaging (EPI) sequence, repetition time (TR) = 4000 ms, echo time (TE) = 19 ms, field-of-view (FOV) = 220 mm × 220 mm, matrix = 64 × 64, in-plane resolution = 3.4 mm, 16 slices, slice
thickness = 4 mm, slice gap = 1 mm, bandwidth = 3004 Hz/Pixel, flip angle = 90°, no background suppression, ascending slice acquisition. The acquisition also included a fat saturation pulse to remove signal contamination from subcutaneous fat. Spins were tagged with a series of 1640 RF pulses for a total tagging duration of 1500 ms, with a tagging efficiency of ~85% (Wu et al., 2007). The offset between the center of the imaging slices and the labeling plane was 70 mm and a post-label delay of 700 ms was used, although this delay has since been considered short for perfusion studies with pCASL (Alsop et al., 2015; see limitations). A series of 60 images (30 tag and 30 control) was acquired for a total scan duration of 4 min.

To assist with registration and anatomical identification for the ASL scan, a high-resolution T1-weighted image was acquired over the entire brain using a 3D MPRAE (Magnetization Prepared Rapid Gradient Echo) protocol with 0.9 mm isotropic resolution (1900/2.3/900 ms repetition/echo/inversion times). All images were collected on a Siemens Magnetom Trio 3T whole-body MRI scanner with 12-channel receiver head coil (Siemens Medical Solutions; Erlangen, Germany).

2.4. ASL analysis

To analyze the ASL data, we used BASIL, a collection of tools in FMRIB’s Software Library (FSL) version 5.0.1 that creates quantitative CBF images from ASL data (www.fmrib.ox.ac.uk/fsl/basil) (Jenkinson et al., 2012). First, the ASL data were reconstructed and motion corrected via a rigid body algorithm in MCFLIRT (Jenkinson et al., 2002) and participants with excessive head motion (relative motion >4.0 mm) were excluded as it was expected data quality would be compromised. Next, the Oxford ASL (oxford_asl) tool within BASIL was applied in order to quantify CBF (Chappell et al., 2009). The tool performed a tag-control subtraction to remove the static tissue contribution, and the resulting time series was used to calculate relative CBF by inversion of the standard model for delivery of the ASL label (Butson et al., 1998) using a Bayesian model inversion technique (Chappell et al., 2009) and an ATT of 700 ms, T1 relaxation time of arterial blood and tissue of 1.6 and 1.3 s, respectively, and a whole brain blood-tissue partition coefficient of 0.9 (Herscovitch and Raichle, 1985). Resulting CBF maps were visually inspected for data quality and clearly corrupted images were excluded.

Relative CBF (in scanner native units) was then converted to absolute physiological units (mL/100 g/min) by estimation of the equilibrium magnetization of arterial blood. This was estimated in a separate calibration process using the unlabeled (control) images from the data by measuring the equilibrium magnetization in the CSF in the ventricles and converting to the equivalent blood value accounting for differences in T1 relaxation time and proton densities (MacIntosh et al., 2008; Wong et al., 2006). The ventricles were automatically masked in the ASL image space by first segmenting the MPRAGE image using FMRIB’s Automated Segmentation Tool (FAST) (Zhang et al., 2001). The resulting tissue-type partial volume estimate was masked using a lateral ventricle template in the MNI152 standard brain image that was linearly transformed into the same space as each participant’s MPRAGE image. The resulting ventricle CSF mask was then transformed into the low-resolution ASL image space and finally thresholded at 90% to identify pure CSF voxels in the ventricles (the resulting mask was inspected for correctness in each subject). The resulting mask for each participant was used to calculate equilibrium magnetization in the ventricles. We also corrected for spatial intensity variations (or bias field or RF inhomogeneities) by calculating the bias field via FAST and scaling the CBF images accordingly. To account for inversion efficiency, we scaled the final CBF values by 1/0.85.

Segmentation and volumetric analysis of the hippocampus and brainstem were performed using a semi-automated, model-based subcortical tool (FIRST; FMRIB’s Integrated Registration and Segmentation Tool) (Patenaude et al., 2011). A two-stage affine registration to a standard space template (MNI space) with 1 mm resolution using 12° of freedom and a subcortical mask to exclude voxels outside the subcortical regions was first performed on each participant’s MPRAGE scan. Next, the hippocampus and brainstem were segmented with 30 and 40 modes of variation, respectively. To achieve accurate segmentation, the FIRST methodology models 317 manually segmented and labeled T1-weighted brain images from normal children, adults, and pathological populations (obtained from the Center for Morphometric Analysis, Massachusetts General Hospital, Boston) as a point distribution model with the geometry and variation of the shape of each structure submitted as priors. Volumetric labels are parameterized by a 3D deformation of a surface model based on multivariate Gaussian assumptions. FIRST searches through linear combinations of shape modes of variation for the most probable shape (i.e., brain structure) given the intensity distribution in the T1-weighted image, and specific brain regions are extracted (see Patenaude et al., 2011 for further description of the method). Modes of variation are optimized based on leave-one-out cross-validation on the training set, and they increase the robustness and reliability of the results (Patenaude et al., 2011).

Anterior and posterior sections of the hippocampus were calculated by determining the center of gravity for both the left and right hippocampus for each participant. The y coordinate from the center-of-gravity calculation was used to divide the region into anterior and posterior sections (Erickson et al., 2011). All segmentations were visually checked for errors. The volume of each participant’s left and right hippocampus (average, anterior, posterior) was measured in cubic millimeters.

The hippocampus and brainstem segmentations were converted into ASL space via FMRIB’s Linear Image Registration Tool (FLIRT) affine linear registration with trilinear interpolation (Jenkinson and Smith, 2001; Jenkinson et al., 2002). The transformation was used in the final calculation of CBF in absolute units. See Fig. 1 for sample CBF maps and anterior/posterior hippocampal segmentations.

2.5. Statistical analysis

Because there were significant and robust correlations between hemispheres (all r > 0.55, p < 0.0001), and no differences between left and right hippocampal CBF or left and right hippocampal volume (t < 1, p > 0.3), we averaged hippocampal CBF values across left and right hemispheres and averaged hippocampal volumes across left and right hemispheres. Average hippocampal CBF and average hippocampal volume were used in subsequent analyses.

Bivariate correlation analyses were conducted using Pearson product-moment correlation coefficients between the descriptive variables (age, sex, SES), aerobic fitness (VO2max), hippocampal CBF, hippocampal volume, brainstem CBF, and brainstem volume to determine appropriate covariates. Linear regressions were employed to test associations between aerobic fitness and hippocampal CBF (average, anterior, posterior) and brainstem CBF, when controlling for age, sex, and volume. T-scores and standardized betas (β) are presented. The alpha level for all tests was set at p < 0.05.

3. Results

Younger age was associated with greater CBF in the hippocampus (r = −0.393; p = 0.01) and brainstem (r = −0.289; p = 0.013), and aerobic fitness was associated with sex such that females had lower fitness scores than males (r = 0.311, p = 0.007). Thus, we included age and sex as covariates in the regression model. Despite no significant correlations between hippocampal CBF and hippocampal...
volume ($r = 0.004$, $p > 0.05$), or brainstem CBF and brainstem volume ($r = -0.015$, $p > 0.05$), we also included volume as a covariate to confirm that blood flow effects were independent of the size of the brain region.

Higher aerobic fitness predicted greater CBF in the hippocampus ($\beta = 0.235$, $t = 2.085$, $p = 0.041$) when controlling for age, sex, and average hippocampal volume (Fig. 2). In particular, higher aerobic fitness predicted greater CBF in the posterior hippocampus ($\beta = 0.241$, $t = 2.163$, $p = 0.034$), when controlling for age, sex, and posterior hippocampal volume (Fig. 2). As hypothesized, aerobic fitness did not predict CBF in the brainstem ($\beta = 0.017$, $t = 0.136$, $p = 0.892$), when controlling for age, sex, and brainstem volume (Fig. 2).

4. Discussion

Consistent with our predictions, aerobic fitness was associated with greater cerebral blood flow in children. Specifically, using ASL perfusion techniques, we demonstrate that higher levels of aerobic fitness are associated with increased cerebral blood flow in the microvasculature of the hippocampus in 7- to 9-year-old children, independent of age, sex, and hippocampal volume. We also explored specificity of the relationship of fitness to the hippocampus and demonstrated, as predicted, no significant association between aerobic fitness and brainstem CBF.

Our results raise the possibility that aerobic fitness plays a role in vascularization of the hippocampus during childhood. Studies suggest a positive benefit of aerobic exercise on brain vasculature in animals (Black et al., 1990; Clark et al., 2009; Kleim et al., 2002; Rhyu et al., 2010) as well as cerebral blood flow measures in middle-aged and older humans (Bullitt et al., 2009; Burdette et al., 2010; Pereira et al., 2007). Here we suggest that these associations may extend to a child population during a critical period of maturation. In fact, angiogenesis has been directly coupled with cerebral blood volume (Dunn et al., 2004; Jiang et al., 2005; Lin et al., 2002; Maia et al., 2005; Sugahara et al., 1998). Although we measured cerebral blood flow rather than cerebral blood volume, we expect perfusion and blood volume to be closely related via the Central Volume Theorem (Newman et al., 2006; Stewart, 1893). A tissue with higher perfusion likely has higher blood volume to sustain the perfusion; however, flow also depends on how quickly the blood passes through the tissue, usually quantified in terms of mean transit time. We postulate that increased blood water delivery and availability in the hippocampus, as a function of higher aerobic fitness, may be due to more blood vessels in this region.

Yet as a variety of molecular and cellular cascades accompany hippocampal changes with aerobic exercise, we can only speculate about the biological mechanisms underlying increased perfusion. For example, in addition to changes in vasculature, aerobic exercise is known to increase cell proliferation and cell survival (Cotman and Berchtold, 2002; Ding et al., 2006), dendritic structure (Redila and Christie, 2006), growth factors (Neuper et al., 1996), and gliogenesis (Uda et al., 2006) in the hippocampus. In fact, angiogenesis and neurogenesis are tightly linked (Louissaint et al., 2002; Palmer et al., 2000). For instance, blocking the secretion of vascular endothelial growth factor (VEGF), a neurotrophic molecule involved in blood vessel growth (Lopez-Lopez et al., 2004), has been found to abolish exercise-induced neurogenesis (Fabel et al., 2003). Further, measures of cerebral blood volume have been said to provide an in vivo correlate of neurogenesis (Pereira et al., 2007). It is possible that some fitness-related differences in cerebral blood flow may be mediated, in part, by neurogenesis.

We are the first to explore the plasticity of perfusion in the anterior and posterior hippocampus in children. Our results do not suggest compelling specificity of fitness on anterior or posterior perfusion (independent of volume), with a significant positive association between fitness and posterior hippocampal CBF and a marginal positive relationship between fitness and anterior hippocampus. Given functional distinctions along the anterior/posterior axis of the hippocampus (Giovanello et al., 2009; Sperling et al., 2003), future studies should integrate a relational
It is interesting that the relationship between aerobic fitness and hippocampal CBF was independent of hippocampal volume, and resting CBF in the hippocampus was not related to the volume of the hippocampus. This lack of CBF–volume association is also supported by a study (Mozolic et al., 2010) that reported significant increases in resting cerebral blood flow with participation in a cognitive training program, but no associations among cognitive training, CBF, and changes in brain structure (via VBM). Together, the data raise the possibility that mechanisms underlying plasticity of blood flow and brain volume in humans are partially independent, and increased blood flow is not solely driven by a larger size of the brain region.

Additionally, our data raise the possibility that extreme group differences in aerobic fitness (e.g., 70th percentile versus 30th percentile VO₂max) may be needed to demonstrate hippocampal volume effects (Chaddock et al., 2010a), whereas CBF differences may be a more sensitive marker to understand how small relative differences in aerobic fitness influence brain health, particularly in terms of microcirculation, during development. We did not observe a significant relationship between aerobic fitness and hippocampal volume in the present study, which included a relatively lower fit sample of children (average VO₂max percentile = 33.5%). However, our previous work demonstrates that higher fit children (>70th percentile VO₂max) show larger hippocampal volumes compared to their lower fit peers (<30th percentile VO₂max) (Shvartz and Reibold, 1990). Thus, different child fitness spectrums in each study may lead to different results and outcomes. Whereas the present study provides a first step in understanding the predictive power of aerobic fitness in hippocampal perfusion during child development, questions still remain regarding the associations among individual differences in aerobic fitness, hippocampal structure and function, and performance on specific memory tasks (e.g., relational, spatial) which require additional research.

Despite our result that aerobic fitness is related to increased hippocampal blood flow in children, we acknowledge limitations of the study, including the choice of an ATT of 700 ms used in post-processing CBF quantification. This value is used in the FSL software based on experience with adult ASL studies (Chappell et al., 2009), although it may not be appropriate for the study of a pediatric population. However, choice of ATT simply provides a scaling factor in CBF calculations and would not affect the relationships presented here. Future work may look to incorporate a multi-delay labeling scheme to simultaneously estimate ATT and CBF, thus providing a more quantitatively accurate measure. We also note that the parameters of the ASL sequence in this study used a short post-label delay for the tagging scheme employed. Specifically, we used a pCASL acquisition as it is recognized to offer the highest signal to noise ratio of all ASL labeling schemes, but with a post-label delay of 700 ms. Subsequent to the present study, the consensus recommendation in the field of ASL acquisition was a longer post-labeling delay for routine pCASL perfusion studies of 1500 ms for children (Alsop et al., 2015). A likely consequence of a shorter post-label delay is that some labeled blood may remain in the arterial vasculature, thereby not being delivered to the tissue at the time of imaging. At the voxel level, the image intensity may not be a pure measure of perfusion, but also include a contribution from arterial blood volume (Chappell et al., 2010a). Thus, it is possible that the region of interest measurements are a mixture of CBF and arterial blood volume, with the amount of arterial blood in the signal dependent on the ATT and post-label delay. In addition, as we did not account for partial volume effects in the hippocampus, this may

Fig. 2. Scatterplots of the associations among aerobic fitness, anterior and posterior hippocampal CBF, and brainstem CBF.
have contributed to the fitness associations we report. A goal for future research will be to use partial volume correction methods on ASL data derived from the hippocampus to determine whether the effects change after accounting for this factor.

This research has important implications, as physical activity is decreasing in and out of the school environment (Troiano et al., 2008), and children are becoming increasingly unfit (Centers for Disease Control and Prevention, 2009). We provide additional evidence to suggest that the developing brain may be plastic and sensitive to individual differences in aerobic fitness levels. Specifically, aerobic fitness may influence how the brain regulates its metabolic demands via blood flow in a region of the brain important for learning and memory.

Acknowledgements

The study was supported by the National Institute of Child Health and Human Development (NICHD) RO1 grant to Dr. Kramer and Dr. Hillman. We would like to thank the Biomedical Imaging Center for their help with data collection and the design of the ASL imaging protocol, in particular Ryan Larsen, Brad Sutton, Holly Tracy, and Nancy Dodge.

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