**INTRODUCTION**

Blood oxygen level-dependent (BOLD) contrast in functional magnetic resonance imaging (fMRI) has been widely applied to investigate brain activity (Bandettini, Wong, Hinks, Tikofsky, & Hyde, 1992; Ogawa et al., 1992). However, BOLD fMRI does not measure neuronal activation directly. It reflects neuronal activation through the neurovascular coupling. In addition to neuronal activity, physiological processes which make alterations in cerebral blood flow (CBF), cerebral blood volume (CBV), and cerebral blood oxygen consumption (CMRO₂) can also affect BOLD signals (Buxton, Uludag, Dubowitz, & Liu, 2004; D’Esposito, Deouell, & Gazzaley, 2003; Hoge et al., 1999). Thus, some features of BOLD signals can be used to assess systemic physiology. This additional physiological information does not require extra scans. More importantly, it can offer concurrent and complement physiological information to the neuronal activation. The combined information can shed light on the interactions between neuronal activation and non-neuronal physiological processes.

**KEYWORDS**

blood vessels, cerebral blood flow, cerebral cortex, magnetic resonance imaging
Significance
The blood oxygen level-dependent (BOLD) signal in functional magnetic resonance imaging can be also affected by non-neuronal cerebral perfusion fluctuation. We developed a method to extract perfusion information from resting-state BOLD data without extra scans. Using this method, we studied the vascular effects of caffeine from resting state data and found caffeine would increase cerebral blood velocity as result of vasoconstriction. Our method offers concurrent free perfusion information in addition to the functional information, which can expose the interactions between perfusion and neural activity.

Among all the features of resting-state (RS) BOLD signals, it has been demonstrated that the time delay and amplitude of low-frequency fluctuations (ALFFs) can be used to assess the cerebral blood velocity and cerebral blood density, respectively (Tong, 2010; Vigneau-Roy, Bernier, Descoteaux, & Whittingstall, 2014). Systemic low-frequency oscillations (sLFOs) are the physiological RS BOLD signals with the frequency range from 0.01 to 0.1 Hz, which overlaps with the frequency range of RS BOLD signals (i.e., neuronal). However, the RS BOLD signals are more regional, leading to different RS networks. On the other hand, sLFOs were traveling global signals with likely extracerebral origins (Hocke, Tong, Lindsey, & Frederick, 2016), which might be related to CO2 fluctuation in the blood, vasomotion, Mayer wave, etc. (Golestani, Kwinta, Strother, Khatamian, & Chen, 2016; Julien, 2006; Wise, Ide, Poulin, & Tracey, 2004). The reason that they are called “systemic” is that the same sLFOs, which are found in RS BOLD signals in the brain (especially large veins), can be detected concurrently by optical method at peripheral sites (fingertip and toe) as slow changes in oxy- and deoxyhemoglobin concentration with time delays (Tong, Hocke, Nickerson, Licata, & Lindsey, 2013). This study demonstrated that the sLFO is a blood-borne oscillation which travels to different parts of the body with blood. Recently, we were able to identify these sLFOs in large arteries and veins (i.e., the internal carotid artery (ICA), superior sagittal sinus (SSS) and internal jugular vein) in the neck from the RS Myconnectome data (Tong, Yao, Chen, & Frederick, 2018). Time delays of approximately 7 s were found between these two highly correlated signals, which is consistent with the cerebral circulation time (CCT) normally measured by ultrasound (Strangman, Culver, Thompson, & Boas, 2002). Another feature is ALFFs (ALFF was determined by standard deviation (SD) of LFO time series), which also provides physiological cerebral vascular properties, such as CBV, vascular density, etc. (Yan, Zhuo, Wang, & Wang, 2011). The ALFFs has been shown to contain a vascular component related to hypercapnic index (Kannurpatti, Biswal, Kim, & Rosen, 2008). Vigneau-Roy et al. (2014) illustrated that the ALFFs of RS BOLD signals were related to vascular density (Vigneau-Roy et al., 2014). The LFO amplitude has also been used to indicate vasoconstriction (Rack-Gomer, Liau, & Liu, 2009).

In this study, we exploited these parameters to derive the non-neuronal physiological parameters from RS fMRI studies and combine the results with the neuronal findings.

Caffeine, as an adenosine A2 receptor antagonist, has both neuronal and cerebral vascular effect. For neuronal effects, it is well known that caffeine can increase the alertness level (Hindmarch et al., 2000). It has also been found that visual BOLD response is faster after ingestion of caffeine (Behzadi & Liu, 2006; Liu, Perthen, & Liu, 2008; Liu et al., 2004). Many studies on caffeine effects in the brain focus on these neuronal activities, for example, investigating brain network connectivity (Rack-Gomer et al., 2009; Tal et al., 2013; Wu, Lien, Chang, & Yang, 2014). However, caffeine also has profound vascular effects. After 250-mg caffeine intake, the resulting vasoconstriction can decrease the CBF about 27% (Addicott et al., 2009; Griffeth, Perthen, & Buxton, 2011; Mathew & Wilson, 1985). These global vascular effects also alter RS BOLD signals as we discussed previously. So far, few fMRI studies have sought to understand caffeine's vascular effect on RS fMRI, and more importantly, understand the interaction with neuronal activations. In this work, we will study vascular effects of caffeine using (a) time delays between sLFOs to map CCT and cerebral blood arrival; (b) LFOs’ SDs to infer the blood density or the size of large cerebral blood vessel. The results will be compared with those from brain network analyses (published separately) and paint a thorough picture of the effects of caffeine.

2 | METHOD

2.1 | Data selection

The publicly available Myconnectome data set (http://myconnectome.org/wp) was chosen to investigate the cerebral vascular effects of caffeine. This data set is about a 2-year study of an individual subject (Male, aged 45). We included the RS scans from session 13 to session 104 (no RS scan in session 52 and session 90), and excluded the pilot sessions and follow-up sessions. There are a number of advantages to using this data set: (a) the Myconnectome data set recruited only one single subject, which eliminates between-subject variation. (b) The data contain 90 high-quality RS fMRI scans, with the equal number of caffeinated scans (scans were taken 90 min after caffeine ingestion) and uncaffeinated scans (i.e., 45 each). (c) Structural images included T1 and T2 weighted scans with high spatial resolution, which made identification of large blood vessels (arteries and veins) easier.

2.2 | Image acquisition

Detailed scan parameters can be found in Poldrack et al. (2015). In short, most of the 10-min RS fMRI scans with eyes closed were acquired using multi-band EPI sequence (TR = 1.16 s, TE = 30 ms, flip angle = 63° (the Ernst angle for gray matter), voxel size = 2.4 × 2.4 × 2 mm, distance factor = 20 (percentage sign), 68 slices,
oriented 30° back from AC/PC, 96 × 96 matrix, 230 mm FOV, MB factor = 4, 10:00 scan length) (Poldrack et al., 2015).

2.3 | Data processing

Figure 1 shows the flow chart of data processing. Data were processed using FSL (FMRIB Expert Analysis Tool, v6.01, http://www.fmrib.ox.ac.uk/fsl, Oxford University, UK (Jenkinson, Beckmann, Behrens, Woolrich, & Smith, 2012)) and a locally developed Matlab program (MATLAB 2017a, The MathWorks Inc., Natick, MA, 2000). RS fMRI scans were preprocessed with following steps recommended by Power et al. (2014). This included: (a) motion correction (FSL mcflirt), (b) linear detrending (MATLAB detrend; “linear”), (c) Nuisance signals removal (six motion parameters and their derivatives estimated by mcflirt) using general linear model (FSL fsl_glm), and (d) spatial smoothing with a Full-Width Half-Maximum of 5-mm isotropic Gaussian kernel (Power et al., 2014).

2.4 | Vascular measures

We assessed the caffeinated/uncaffeinated difference in the following parameters: (a) CCT from the time delays between sLFOs in fMRI signals from ICA and SSS; (b) delay maps (i.e., cerebral blood arrival map) obtained by calculating the voxel-wise time delays between sLFOs in fMRI signals from each voxel and SSS; and (c) cerebral blood density from the voxel-wise SDs (ALFFs) of LFO in fMRI signals. Lastly, we compared our results with previous research of RS networks connectivity on the same data set (Poldrack et al., 2015).

2.4.1 | Cerebral circulation

In order to calculate the delays of the sLFOs between ICA and SSS in caffeinated and uncaffeinated scans, we applied the same method in Tong et al. (2018) to extract the fMRI signals in these large blood vessels. In short, left and right ICAs and SSS were identified from structural images (T1- and T2-weighted images) (Tong et al., 2018). Time series within these identified blood vessels were extracted and averaged (FSL fslmeans) from the RS fMRI (after registering high-resolution vessel masks onto the low-resolution fMRI images). To calculate the time delays between the sLFOs of two vessels, we conducted the following steps. (a) A band-pass zero-lag Butterworth filter (0.01–0.1 Hz) was applied to the time series from SSS, global signal (GS) and ICA to extract the sLFOs, which were then oversampled to increase the temporal resolution (from 1.16 to 0.116 s). (b) The principal component analysis was then applied on the signals from left and right ICAs to obtain the principal signal of the low-frequency component to improve the signal-to-noise ratio. (c) Time delays between the sLFOs from GS, ICA, and SSS were calculated using cross-correlation (xcorr in Matlab), which found the absolute maximum cross-correlation coefficient (MCCC) between the signals within a ± 15 s window. The MCCC threshold was set at 0.28, which was the significance threshold based on previous research (Hocke et al., 2016). In short, voxel-wise delay maps were generated based on the method developed previously (Tong & Frederick, 2014). In short, the time delay between every voxel’s RS BOLD signal and the SSS signal was calculated to represent the blood arrival time at that voxel. The delays calculated between large vessels, as well as the whole brain delay maps, were later compared between caffeinated and uncaffeinated scans to expose the caffeine effects on the circulation.

![FIGURE 1](image1) The flowchart presents the procedure of data processing. First, 90 resting-state scans were sorted into caffeinated and uncaffeinated groups. Second, vascular information was extracted from scans separately, which included (a) Time delays between systemic low-frequency oscillations (sLFOs, 0.01–0.1 Hz) of superior sagittal sinus (SSS) and brain voxels, (b) Cerebral circulation time (CCT) was measured by time delay between sLFOs of SSS and internal carotid artery (ICA), (c) LFOs amplitude in brain voxels were measured. Lastly, the above vascular effect of caffeine was compared with the neuronal effect of caffeine from previous research.
2.4.2 Cerebral blood density (volume)

Previous research reported that the ALFFs in the RS BOLD signal were positively correlated with blood density in the voxel (Vigneau-Roy et al., 2014). Here, we calculated the SD in RS BOLD signal to represent the cerebral blood density (volume). Voxel-wise SD maps were calculated with the following steps: First, a band-pass zero-lag Butterworth filter (0.01–0.1 Hz) was applied to the time series of each voxel in the brain. Second, SDs of the low-frequency time series were calculated in each voxel. Finally, the SD differences (in percentage) between caffeinated and uncaffeinated scans were obtained by the subtraction.

2.4.3 Statistical analysis

The distributions differences of time delays between GS, ICA, and SSS in caffeinated and uncaffeinated scans were tested by the Kolmogorov–Smirnov test (Daniel, 1990).

To assess the statistical significance of the difference between caffeinated and uncaffeinated scans, we conducted the randomization procedure. In detail, 90 delay maps (or SD maps) were randomly placed into two groups (45 each). Each group would have roughly same numbers of caffeinated scans as well as uncaffeinated scans. The averaged delay maps (or SD maps) were calculated from each group and the subtraction between these two groups was obtained. Since there is no difference in the subtraction (because equal numbers of caffeinated and uncaffeinated scans are included in each group), the subtraction result should be close to 0 and its distribution could be used to extract the threshold of the null hypothesis to assess the real effect. The processes of generating random groups and the subtraction between these groups were repeated for 10,000 trials. The threshold was selected as $p < 0.01$ or $p > 0.99$ ($p$ indicated the probability of having spurious results), from the averaged distribution of the subtractions. Also, we applied two-sample $t$ test to further validate the differences in the delay and SD maps between caffeinated and uncaffeinated scans (SPM (https://www.fil.ion.ucl.ac.uk/spm/)). The resulting SPM(T) map was thresholded at $p < 0.05$, corrected for multiple comparisons using the false discovery rate (FDR)-criterion (Genovese, Lazar, & Nichols, 2002).

3 RESULTS

3.1 Comparison of caffeinated and uncaffeinated results

3.1.1 CCT difference

Figure 2 shows the time delay differences between SSS and GS, GS and ICA, ICA and GS. Time delays are significantly shorter between SSS and ICA in the caffeinated scans ($5.03 \pm 0.617$ s) than those from the uncaffeinated scans ($5.96 \pm 0.659$ s) ($p < 0.001$), indicating the shorter CCTs in caffeinated scans. This result is consistent with the following observations: (a) time delays are significantly shorter between SSS and GS in the caffeinated scans ($2.87 \pm 0.237$ s) than the uncaffeinated scans ($2.74 \pm 0.303$ s) ($p < 0.001$). (b) Time delays are significantly shorter between GS and ICA in caffeinated scans ($2.87 \pm 0.767$ s) than uncaffeinated scans ($3.38 \pm 0.783$ s) ($p < 0.05$).

3.1.2 Delay map difference

Figure 3a shows the average delay map from 90 RS scans (from a single subject). It demonstrates that the delay map calculated from this data largely matches the cerebral blood circulation. Red indicates the early arrival of the sLFOs (i.e., blood), while blue denotes late arrival. Since the voxel-wise delay value was calculated between the sLFO from each voxel and the SSS, the arrival times were relevant to that of SSS. Figure 3a shows that the brain is dominated by "Red," indicating that blood reached most of the voxels before it arrived at SSS (which is expected, as the SSS is a major blood drainage vessel from the brain). Interestingly, the ICAs were automatically identified as the regions where blood arrives very early (see the arrows in Figure 3a). This phenomenon confirmed that sLFOs were pumped into the brain through arteries, such as ICAs. The detailed discussion about the sLFOs in ICA can be found in our previous publication (Tong et al., 2018). The fact that the delay map can automatically locate ICA out of RS fMRI data with correct delay values showed the robustness of the sLFO in fMRI signal as well as our method. Delay map difference obtained by subtracting the delay

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**FIGURE 2** Comparison of time delay distribution between 45 caffeinated RS scans (brown) and 45 uncaffeinated RS scans (gray). Time delays were calculated between (a) superior sagittal sinus (SSS) and global signal (GS), (b) GS and internal carotid artery (ICA) and (c) SSS and ICA. Results with absolute maximum cross-correlation coefficient less than 0.28 were excluded to avoid spurious correlations. Time delay distributions between caffeinated and uncaffeinated scans were compared by Kolmogorov–Smirnov test.
map of 45 caffeinated scans from that of 45 uncaffeinated scans is shown in Figure 3b. It shows that the subtracting result of the delay maps is dominated by "Blue," indicating that blood flow speed increased in most of the brain regions under the effect of caffeine.

Figure 4a shows the distribution of randomized subtraction (from 10,000 random combinations) in blue and the that of real caffeinated versus uncaffeinated subtraction in red (see Section 2.4.3). From the distribution of randomized subtraction in Figure 4a, we found the results were clustered tightly around 0, which was expected due to no caffeine effects in the randomized data. However, the distribution of real caffeinated versus uncaffeinated subtraction was much broader. This further indicates that the differences calculated between caffeinated and uncaffeinated scans are significant. We obtained the thresholds on delay differences to be \( >0.037 \) s or \( <-0.042 \) s (\( p < 0.01 \)), which were used on Figure 5a.1 and upper panel of Figure 8 to produce statistically significant delay maps differences. Also, Figure 5a.2 showed that the delay differences between caffeinated and uncaffeinated scans were significant (i.e., two-sample t test) at a low statistical threshold (\( p < 0.05 \), FDR corrected).

### 3.1.3 Standard deviation difference

Figure 6a shows that voxels with higher LFO amplitude (i.e., SD) in yellow, projected onto the subject’s own structural scan (T2-weighted image). The yellow regions overlapped precisely with the visible blood vessels (marked by squares in T2-weighted image Figure 6b). This is in accordance with the findings in Vigneau-Roy et al. (2014), which reported that the ALFFs of RS BOLD signal were positively correlated with blood density in the voxel (Vigneau-Roy et al., 2014). We used this feature to assess blood vessel changes due to caffeine effect. The rows/columns of voxels (shown by arrows) in Figure 7a(1–2) pass through areas with high blood density (or large blood vessels). The corresponding profile (of RS BOLD signals’ SDs along the white arrow) is shown in Figure 7b(1–2), which presented the averaged results of uncaffeinated scans and caffeinated scans. From Figure 7a(1–2), visible decreases in the yellow pattern were found in caffeinated results, which is confirmed by Figure 7b(1–2). It shows the intensity profile of the SDs of LFOs was “constricted” and decreased after coffee. This observation could be interpreted as vasoconstriction due to the caffeine. To assess the vessel constriction effect in the whole brain, we compared the SD

![Figure 3](image)

**FIGURE 3** (a) Averaged delay map from 90 RS scans. Time delays were calculated between systemic low-frequency oscillations of superior sagittal sinus (SSS) and all the voxels in the brain. The red-yellow color showed early arrival time of blood and blue color showed late arrival time with reference to SSS. (b) The difference between caffeinated and uncaffeinated delay maps. Blue color showed time delays between the voxels and SSS were shorter in the caffeinated scans than the uncaffeinated scans.

![Figure 4](image)

**FIGURE 4** Original results of time delay difference (a) and SD difference (b) between caffeinated and uncaffeinated scans were presented as red distribution. Randomization results of time delay difference (a) and SD difference (b) between caffeinated and uncaffeinated scans were presented as blue distribution.
difference in percentage ((caffeinated–uncaffeinated)/caffeinated) between caffeinated and uncaffeinated scans. To assess the result in the statistical point of view, we also conducted the “randomization” procedure used before, with similar findings (shown in Figure 4b). The distribution of real caffeinated versus uncaffeinated subtraction was much broader than the results from randomized subtractions, indicating the difference calculated between caffeinated versus uncaffeinated is not spurious but robust. We obtained the thresholds on percentage SD difference to be >1.01% or <−0.12% (p < 0.01), which were applied on Figure 5b.1 and lower panel of Figure 8 to produce the map of statistically significant SD differences. Also, Figure 5b.2 showed that the SD differences between caffeinated and uncaffeinated scans were significant (i.e., two-sample t test) at a low statistical threshold (p < 0.05, FDR corrected).

3.2 | Neuronal and vascular effects

As we know, caffeine has both vascular and neural effects. It is of great interest to understand the spatial correlation between vascular and neuronal effect (from previous studies). In detail, we conducted following steps: first, significant whole brain time delay differences (Figure 5a) and SD percentage differences (Figure 5b) were parcellated according to nine functionally coherent RS networks (Gordon et al., 2014). These well-known RS networks have been studied with the Myconnectome data set previously to assess the neuronal effect of caffeine (Laumann et al., 2015; Poldrack et al., 2015). Subsequently, the average of time delay differences and

FIGURE 5 (a.1) Thresholded (from randomization) subtraction of delay map (caffeinated–uncaffeinated). (a.2) Subtraction of delay map corrected by false discovery rate (FDR) (p < 0.05) (caffeinated–uncaffeinated). (b.1) Thresholded (from randomization) SD percentage difference ((caffeinated–uncaffeinated)/caffeinated). (b.2) SD percentage difference corrected by FDR (p < 0.05) ((caffeinated–uncaffeinated)/caffeinated)

FIGURE 6 (a) Averaged SD map from 90 RS scans. (b) T2-weighted image. Brain region with high systemic low-frequency oscillation amplitude matched the locations of big blood vessels (marked by squares in (b))
The significant time delay differences (upper blue panel) and SD differences (lower red panel) between caffeinated and uncaffeinated scans were showed for 9 RS networks. (DMN: default mode network, VIS: visual network, FP: fronto-parietal network, DA: dorsal attention network, VA: ventral attention network, SAL: salience network, CO: cingulo-opercular network, SM: somatomotor network, AUD: auditory network)

**FIGURE 8** Vascular effect of caffeine on 9 RS networks. The significant time delay differences (upper blue panel) and SD differences (lower red panel) between caffeinated and uncaffeinated scans were showed for 9 RS networks. (DMN: default mode network, VIS: visual network, FP: fronto-parietal network, DA: dorsal attention network, VA: ventral attention network, SAL: salience network, CO: cingulo-opercular network, SM: somatomotor network, AUD: auditory network)

SD percentage differences were obtained from each network. The results were showed in Figure 8. Negative values of blue boxplot indicated shorter time delays, while negative values of red boxplot showed smaller vessel size after caffeine intake.

In Figure 8, significantly shorter time delays and smaller SDs can be found in all nine RS networks in caffeinated scans. Top four RS networks with the highest time delay differences (i.e., auditory, somatomotor, visual, and ventral attention networks), are also the top four networks with the highest SD percentage differences. Moreover, we found high linear correlation between relative fluctuations (represented by blue and red lines in Figure 8) in averaged differences of time delays and SDs among different networks ($r = 0.6$, $p = 0.09$, high $p$-value reflected few data points). This feature indicates that network-specific flow velocity changes well correlated with the vessel size changes under the influence of caffeine.

Interestingly, Poldrack et al. found significantly lower connectivity within and between somatomotor, dorsal attention, and visual networks in caffeinated scans. Moreover, similar decreasing connectivity was found within motor cortex after caffeine intake (Rack-Gomer et al., 2009). These RS networks (i.e., motor, visual) with decreasing connectivity were the same networks which have the largest time delays and SDs changes after caffeine intake (except dorsal attention network). It shows that caffeine has profound vascular effects (e.g., vessel constriction) in the brain regions where neuronal connectivities were also weakened. This is the first study to expose simultaneously the vascular and neuronal effect of a drug from the same RS fMRI data. The interaction between regional perfusion and neuronal activity, exposed by the method, are crucial for the understanding of the effects of drugs, brain diseases, etc. More qualitative studies large populations are needed to fully explore the effects.
4 | DISCUSSION

4.1 | Caffeinated scans have the faster speed of blood flow

Figure 2 showed statistically significant shorter time delays of sLFOs between SSS and ICA, in caffeinated scans than in uncaffeinated scans (~0.9 s), which could be interpreted as decreases of CCTs. However, studies from positron emission tomography (PET) and arterial spin labeling (ASL) reported that decreasing CBF was detected after caffeine ingestion (Addicott et al., 2009; Cameron, Modell, & Hariharan, 1990; Rack-Gomer et al., 2009; Wong, Olafsson, Tal, & Liu, 2012). Decreasing CBF would, on the surface, seem to imply an increase of the CCT. This seems to contradict the finding in this study. One possible explanation may lay in the fact that CCT is not necessarily inversely correlated with CBF. CCT reflects the time for a bolus or red blood cells to travel from the ICA to the SSS (speed). However, CBF is measured in the amount of blood (ml) passing through a certain mass of tissue (100 g) in unit time (e.g., min). If vessel size stays the same, an increase in CBF will decrease CCT. However, in the situation of caffeine, pressure induced by vessels constriction (coupled with elevated cerebral blood (James, 2004)) leads to the increased blood flow speed (see Section 4.2 in the discussion). The effects of vessel constriction on CBF are twofold: (a) elevated blood speed will increase CBF; (b) vessel constriction will decrease CBF. Since the latter effect is dominant (~r², r is vessel diameter), vessel constriction will lead to a decrease in CBF. This model explains how caffeine related vessel constriction can increase the blood flow speed inside the vessels, (leading to shorter CCT) while decreasing the CBF. Nevertheless, caffeinated study using transcranial Doppler to measure the velocity of the middle cerebral artery (MCA) after 250 mg of caffeine ingestion showed the significant reduction of the velocity by 13% and reduction of MCA diameter by 4.3% (Lunt, Ragab, Birch, Schley, & Jenkinson, 2004). This does not contradict our model (see in next paragraph) since the RS BOLD signals we studied (e.g., SSS and global mean) are mostly vein signals. The perfusion parameters we derived, including time delay and SD, are biased/more sensitive toward the venules and veins. Therefore, we think the velocity change and vessel constriction measured by RS BOLD signals mostly happens in the capillary, venules, and veins (see discussion in Section 4.3).

Delay map differences between uncaffeinated scans and caffeinated scans are shown in Figure 5a. The regions in blue are the areas where the time delays are shortened referred to SSS (i.e., increase CBF speed). Figure 5a showed that in most of the brain the CBF speed did increase mildly (within 1 s) after caffeine intake. However, there are also regions where CBF speed reduced (red regions: longer time delays referred to SSS, especially in Figure 5a). These regions are concentrated in the lower part of the brain overlapping with MCA. This observation matched the previous publication of transcranial Doppler studies of caffeine (Lunt et al., 2004).

4.2 | Caffeinated scans have lower LFO amplitude

It has been known that areas of high SDs in RS fMRI correspond very well with large vessels in the flow-weighted image (Kim, Hendrich, Hu, Merkle, & Uğurbil, 1994). It was even suggested to use these SD maps to identify the location of large vessels. A recent study has investigated the link between regional variations of vascular density (VAD) and the ALFFs of RS BOLD signals. They found the positive correlation between the VAD (from the venous vascular tree in susceptibility-weighted imaging) and the ALFFs. These findings were similar to the results showed in Figure 6. As result, vessel constriction will decrease the VAD signal, which results in smaller ALFFs. We can see in Figure 7 that the SD profile curves are narrower and shorter in the caffeinated scans, which indicated that the effect of caffeine is largely vessel constriction. This is more obvious in the case of large veins. In the top left graphs of Figure 7, SSS were identified with RS BOLD signals of high SD. Its diameter became smaller after caffeine intake. The profile curves (shown on the right) confirmed the observation, which was consistent with the previous findings of the reduction of BOLD amplitude after caffeine intake (Mulderink, Gitelman, Mesulam, & Parrish, 2002; Rack-Gomer et al., 2009)). Moreover, a similar effect was also found in the global LFO (Wong et al., 2012). To validate the global effect of caffeine, the SD of global LFO was also calculated in this study. As result in supplement material Figure S1, we compared SDs of global LFO between uncaffeinated scans and caffeinated scans. The result indicated that the SDs of LFOs were significantly larger in uncaffeinated scans than in caffeinated state (p < 0.001), which was consistent with previous findings (Wong et al., 2012).

Whole brain SD difference between uncaffeinated scans and caffeinated scans was showed in Figure 5b. Blue regions indicated the vessel constriction after caffeine intake, while the red regions showed the opposite effects. We can see that vessel constrictions are commonly found in the regions of high blood density (e.g., gray matter), while the opposite was found in the white matter and lower part of the brain. Our results are consistent with previous ASL findings, which reported that decreasing CBF were widespread in the gray matter after caffeine intake (Buch, Ye, & Haacke, 2017; Merola et al., 2017). Compared to the delay map difference shown in Figure 5a, we found that the regions of vessel constriction after caffeine intake overlap largely with the regions where the blood flow speed increases. This result is consistent with our model proposed in Section 4.1. In Section 4.3, we will further discuss the spatial correlations of these two vascular effects.

4.3 | Possible confounds

4.3.1 | Caffeine effect

In this study, we want to understand the effects of caffeine on the vasculature. However, there are three limitations. First, our study was on a single subject. The studies with more subjects are needed to
To the best of our knowledge, this is the first study using dynamic features of sLFOs to extract physiological perfusion information from RS fMRI drug studies (i.e., caffeine). We have illustrated that caffeine decreases the vessels size (i.e., the lower dynamic SD of LFO) and increased the blood flow speed in those vessels (i.e., the shorter time delays), using only fMRI data. These findings were supported by the spatial overlapping of these two vasoconstriction effects of caffeine. Furthermore, we found these vascular effects are also spatially correlated with the neuronal effect of caffeine obtained from previous studies. In conclusion, we demonstrate that vascular features can be extracted from fMRI data which may offer valuable complementary information to the neuronal activation. This is extremely useful in drug studies, where vascular and neuronal effects are intertwined and understanding the individual effect as well as the interactions is crucial. In the future, we will apply our method to assess the vascular information from drug-related fMRI studies and compared it with the neuronal activations quantitatively.

Declaration of Transparency

The authors, reviewers and editors affirm that in accordance to the policies set by the Journal of Neuroscience Research, this manuscript presents an accurate and transparent account of the study being reported and that all critical details describing the methods and results are present.

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Conflict of Interest

The authors have no conflict of interest to declare.

Authors Contributions


Data Accessibility

The publicly available Myconnectome data set can be found on http://myconnectome.org/wp. The corresponding codes used in section 2.4.1 can be found on the following website (https://github.com/TonglabPurdue/Systematic-low-frequency-oscillation-in-fMRI/tree/master/Delay_Map_fMRI).

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Figure S1.** Global signal’s SD difference between caffeinated scans (brown) and uncaffeinated scans (gray).

**Figure S2.** Framewise displacement (FD) difference between caffeinated scans (brown) and uncaffeinated scans (gray).

Transparent Science Questionnaire for Authors.

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