

SCIENTIFIC REPORTS



OPEN

Influence of Resting Venous Blood Volume Fraction on Dynamic Causal Modeling and System Identifiability

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Received: 08 February 2016

Accepted: 17 June 2016

Published: 08 July 2016

Changes in BOLD signals are sensitive to the regional blood content associated with the vasculature, which is known as V_0 in hemodynamic models. In previous studies involving dynamic causal modeling (DCM) which embodies the hemodynamic model to invert the functional magnetic resonance imaging signals into neuronal activity, V_0 was arbitrarily set to a physiologically plausible value to overcome the ill-posedness of the inverse problem. It is interesting to investigate how the V_0 value influences DCM. In this study we addressed this issue by using both synthetic and real experiments. The results show that the ability of DCM analysis to reveal information about brain causality depends critically on the assumed V_0 value used in the analysis procedure. The choice of V_0 value not only directly affects the strength of system connections, but more importantly also affects the inferences about the network architecture. Our analyses speak to a possible refinement of how the hemodynamic process is parameterized (i.e., by making V_0 a free parameter); however, the conditional dependencies induced by a more complex model may create more problems than they solve. Obtaining more realistic V_0 information in DCM can improve the identifiability of the system and would provide more reliable inferences about the properties of brain connectivity.

Dynamic causal modeling (DCM) is now widely employed to explore causality between neural systems based on neuroimaging data obtained using methods such as electroencephalography (EEG), magnetoencephalography (MEG), and functional magnetic resonance imaging (fMRI)^{1–3}. DCM is based on a neurobiology-based model of neuronal population dynamics. A typical application of this technique is to select a set of specific task-dependent “nodes” (active loci) within regions of interest (ROI) on the basis of conventional activation analysis, then to infer the causal links among these distributed brain regions from the candidate models^{4–6}; this constitutes a classic inverse problem. For the case of fMRI data, because the hemodynamic response is a secondary effect induced by neuronal activity, the method also incorporates a hemodynamic model (Balloon model) to convert the underlying neuronal activity into the observed fMRI response in each brain region of interest^{7,8} (Fig. 1). This hemodynamic step in data generation can result in complications when attempting to make statistical inferences about the connectivity architecture.

DCM reflects the increasing interest in functional integration among brain regions, as measured using neuroimaging. Figure 1 shows a schematic representation of DCM. In contrast to conventional model analysis of region-specific activations only allowing model inputs of experimental factors (εu in Fig. 1), in DCM the inputs from several regions (represented by a neuronal state vector z with one state per region) embrace not only the direct efficacy of experimental inputs (the C matrices in Fig. 1) but also the efficacy of neuronal input from other regions (the A matrices in Fig. 1)^{9,10}. In this sense, DCM can be regarded as a collection of hemodynamic models, one for each area, in which the experimental inputs are supplemented with neural activity from other areas¹. The causal correlation is then parameterized in terms of coupling among the neuronal activities in different regions. The objective is to correctly estimate these parameters on the basis of all of the observed hemodynamic responses.

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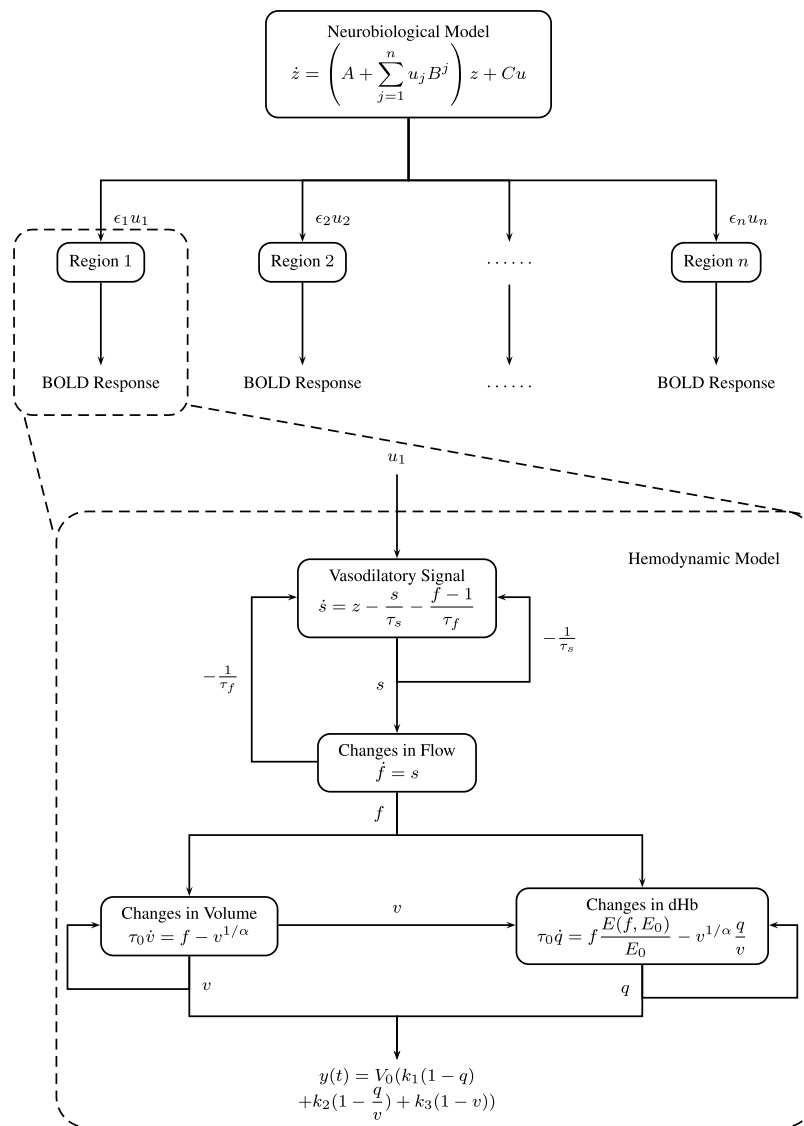


Figure 1. Schematic illustration of DCM. DCM expresses brain causality at the level of neuronal dynamics. The neurobiological model z describes neural dynamics and connectivity among brain regions. Matrix A represents the fixed connectivity among the regions in the absence of input, matrix B encodes the change in connectivity induced by an input, and matrix C embodies the strength of the direct influences of inputs. The B parameters correspond to condition or context sensitive changes in the A parameters. These mediate the influence of experimental or exogenous inputs u on the (linear) connectivity encoded by the A parameters. It is usually these (B parameters) that are of interest because they report the effects of changes in experimental condition. The hemodynamic model with four state variables $s, f, v,$ and q converts the neural dynamics into the observed BOLD signals. The model parameters comprise inputs u , neuronal efficacy ϵ , the time constant for vasodilatory signal decay τ_f , the feedback autoregulation time constant τ_s , transit time τ_0 , stiffness parameter α , resting oxygen extraction fraction E_0 , and resting blood volume fraction V_0 . Abbreviation: dHb, deoxyhemoglobin.

The nonlinearities inherent in hemodynamic response, and the conditional dependence between hemodynamic and the neuronal model parameters make model inversion and subsequent inference a challenging problem. The output of the hemodynamic model is the product of the resting blood volume fraction V_0 and a nonlinear function of the state variables (Equation $y(t)$ in Fig. 1). This means that it is not possible to simultaneously estimate V_0 and other parameters without applying additional constraints; instead, only their product can be estimated¹¹. Moreover, the use of a complicated nonlinear model means that the effects of some parameters on the output interfere with those of other parameters^{11,12}. The influence of changing one parameter could be compensated by adjusting other parameters to produce exactly the same output¹¹. On the other hand, changes in BOLD signals are sensitive to the regional venous blood volume fraction, which is represented as the physiological parameter V_0 in the hemodynamic model¹³. This can result in the active domain of the BOLD signal often being overly influenced

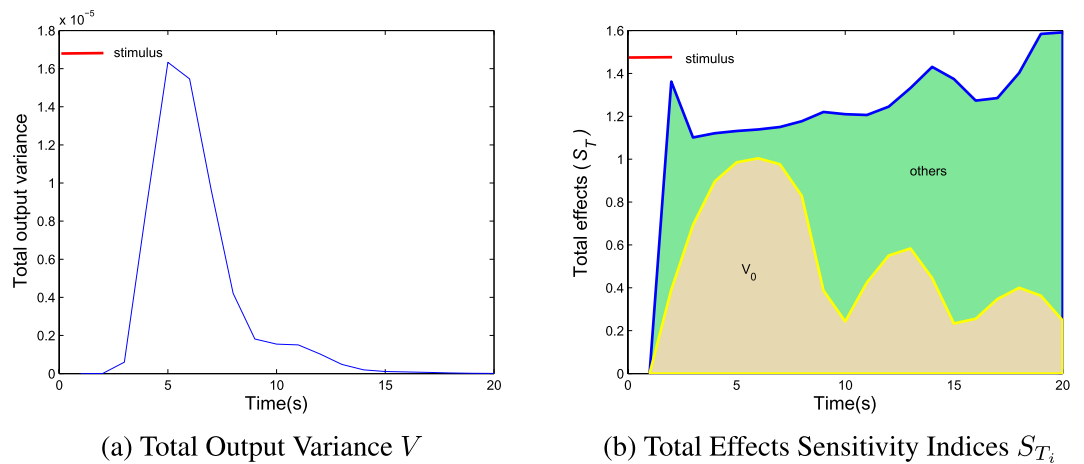


Figure 2. Time course of the total output variance (a) and total effects sensitivity indices S_{T_i} of V_0 and other parameters (b) in an event-related paradigm for a 2s stimulus in the hemodynamic model. (a) The general uncertainty of the model output for all model variables is similar to the known BOLD responses. It starts to deviate from the baseline at approximately 2s after the onset of the stimulus, peaks at about 5s, and takes about 13s to return to the baseline after the stimulus is turned off. (b) S_{T_i} indicates the relative contribution of model output associated with variables X_i . It is clear that V_0 play a leading role in driving the hemodynamic model output, especially during a positive BOLD response, indicating that calibrating V_0 is an important parameter for raising the identifiability of hemodynamic system. The red line indicates the duration of the external stimulus.

by regions with large blood contents¹⁴; in these regions V_0 might exceed 0.6¹⁵. Many pathological conditions also might be associated with an abnormal V_0 ^{16,17}.

To assess the sensitivity of the inference to all parameters, we computed the variance in measurement space (fMRI time-series) induced by changes in V_0 (and other parameters). If a parameter change causes a large change in measurement space, then this usually means it can be identified efficiently. Figure 2a shows the results of this analysis. Because the measurements space is a time-series, we can express this sensitivity as a function of peristimulus time using a short input (2s). It is clear that the general variance of model output produced by all parameters is similar to the known BOLD responses. The relative contribution of output variance associated with V_0 is shown in Fig. 2b. The results show that V_0 plays a leading role in driving the uncertainty of the hemodynamic model output, especially during a positive BOLD response. Calibrating this parameter is therefore of great importance for increasing the system identifiability¹⁸. Despite its importance, as reflected by numerous attempts to design models and suggest improvements to them^{3,4,19}, in most studies this parameter is arbitrarily assigned a physiologically plausible value (e.g., $V_0 = 0.04$), owing to the ill-posedness of the inverse problem^{11,12,19}.

While the DCM approach is often applied to the analysis of activation areas, such an arbitrary assumption about V_0 is not always appropriate, and it might lead to inaccurate estimation of the model behavior. It is therefore necessary to consider the vascular content of areas showing significant BOLD activation in DCM studies. In this study we investigated how V_0 influence DCM using both synthetic and real experiments. We approached the problem from a purely mathematical and modeling perspective, and did not consider the physiological mechanism underlying the BOLD contrast.

Experiments

Synthetic Data. Since ground-truth fMRI data are not available, a simulated architecture was chosen in this study. The model architecture, which is depicted in Fig. 3, comprises three regions: A1, A2, and A3. The extrinsic input is applied to A1, while A1 drives A2 and A3 directly, and gets feedback from them. We designed three artificial BOLD responses with three distinct V_0 values for the three regions: $V_0^{A1} = 0.04$, $V_0^{A2} = 0.02$, and $V_0^{A3} = 0.04$. The experimental condition of the synthetic dataset involved successive blocks alternating between rest and task, starting with task. All connectivity parameters are set as 1. The prior expectations for the biophysical parameters follow the defaults set in SPM software¹². The analysis was performed over 120 time points (repeat time, TR = 2s) in blocks of 6, giving 12 20-s blocks. The external stimulus had value of 1 and 0 when the stimulus was on and off, respectively^{11,20}. We then added white Gaussian noise at a signal-to-noise ratio of 3 to the clear BOLD signal³. The addition of white Gaussian noise is a slight simplification, which violates the prior assumptions of DCM. This is because DCM expects serial correlations in observation noise that are, in part, mediated neuronally. However, this mild violation does not affect the point that we want to make using our simulations. Uncorrelated or white thermal noise is present in fMRI time-series; however, this is generally averaged away when taking region of interest summaries. Except where stated otherwise, all data manipulation (simulation and processing) performed in this study was conducted using SPM software.

We specified 13 competing models in the model space. For all of these models, the exogenous stimulation was applied over area A1. Figure 4 shows the results of DCM analysis applied to the synthetic data when V_0 was presupposed to be correctly known ($V_0^{A1} = 0.04$, $V_0^{A2} = 0.02$, and $V_0^{A3} = 0.04$) in the analysis procedure. The values in brackets are V_0 in these regions, and the coupling parameters are shown alongside the corresponding

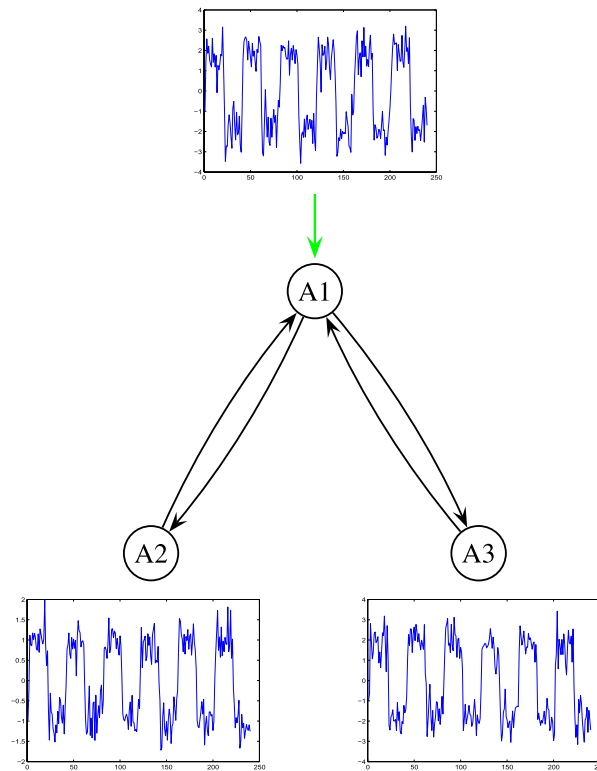


Figure 3. Simulated model. The value of all connectivity parameters are 1.

connections. Not surprisingly, DCM analysis can reliably and accurately identify the model architecture in this case (Fig. 4). However, when another set of unrealistic V_0 value was introduced in the DCM analysis ($V_0^{A1} = 0.04$, $V_0^{A2} = 0.04$ and $V_0^{A3} = 0.02$), this produced a completely different (and wrong) inferences about the architecture (Fig. 5). The most plausible model now had a false connection from A2 to A3, whereas the backward link from A2 to A1 was missing. Figure 6 is another example for a different set of V_0 values: $V_0^{A1} = 0.02$, $V_0^{A2} = 0.04$, and $V_0^{A3} = 0.02$. The results suggest that the choice of V_0 value directly affects not only the estimated connection strength but also even the underlying architecture.

Figure 7 shows the results of combining all 13 models under the three different sets of prior assumptions about V_0 . We see here that the correct model under the correct prior assumptions is superior.

Real Data. Real fMRI data were collected in a visual attention experiment, which can be downloaded freely from the SPM website (<http://www.fil.ion.ucl.ac.uk/spm/>). Details of the experimental paradigm and acquisition parameters are available elsewhere²¹. Four regions were selected for DCM analysis following a conventional SPM analysis. The entire model comprised four regions: V1, V5, PPC, and PFC. Figure 8 shows the locations of the regions and their time series that were entered into the DCM.

In this study we specified 17 competing models in the model space. In all of these competing models, the extrinsic influences of the inputs (e.g., a photic stimulus, indicated in green in Figs 9 and 10) on neuronal activity and the modulation of motion by the experimental manipulations (blue in Figs 9 and 10) were the same, but the attentional modulation of the response (yellow in Figs 9 and 10) could switch between functionally specialized sensory areas and the backward connection from area PFC to area PPC. Figure 9 shows the most reasonable model based on the present observations selected from the models space when V_0 values of areas V1, V5, PPC, and PFC were assigned as 0.02, 0.02, 0.04, and 0.04 respectively. We then changed the V_0 value in area V5, V_0^{V5} , from 0.02 to 0.04, while other conditions remained unchanged. In this case the most reasonable model did not contain the backward connection from area PFC to area PPC, and attentional modulation was imposed on the link from area PPC to area V5 (9). Its structure was completely different from the one in Figs 9 and 10 presents the model structure and results obtained comparisons of the DCMs.

Discussion

This study used DCM to address how the resting venous blood volume fraction (i.e., V_0) influences effective connectivity. The employment of an arbitrary V_0 is universal in DCM analysis due to the ill-posedness of the inverse problem^{11,22}. However, for the BOLD modality, the signal intensity is sensitive to the regional V_0 . This can result in the active domains that are subject to DCM analysis being overly influenced by those areas with large blood contents¹⁴. In this situation the use of incorrect V_0 values might lead to inaccurate causality inference between brain areas. In this study we have addressed this concern by using both synthetic and real experiments. The

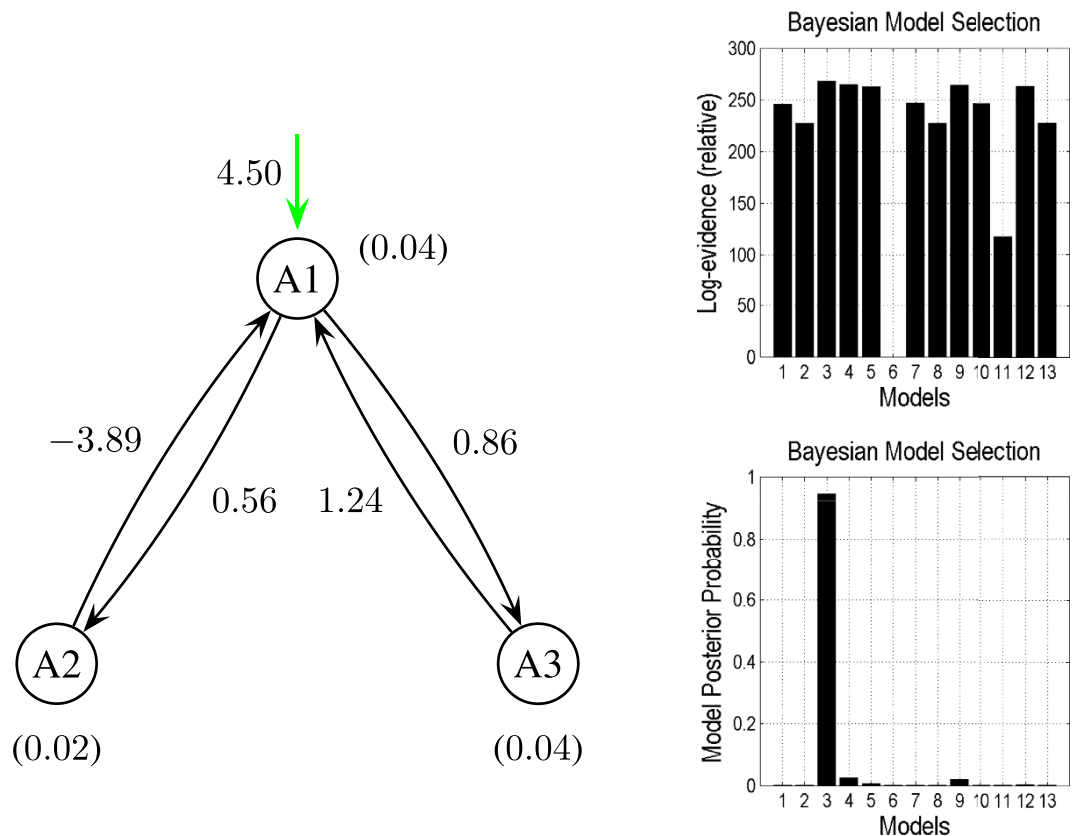


Figure 4. The most reasonable model based on the present observations with $V_0^{A1} = 0.04$, $V_0^{A2} = 0.02$, and $V_0^{A3} = 0.04$, selected from 13 competing models. The green line represents the extrinsic input. The coupling strength is shown alongside each corresponding connection. The values in brackets are the V_0 values in the corresponding regions. The right graphs provide the results obtained in comparisons of the DCMs, illustrating that Model 3 is superior to the other models.

obtained results show that the ability of DCM analysis to discover the effective connectivity depends critically on the assumed V_0 value used in the analysis procedure. The choice of V_0 value not only directly affects the estimated connection strengths, but more importantly might also affect the inferred connection structure. Making inappropriate assumptions about V_0 could interfere with the accurate identification of causal correlations²³. We therefore argue that prior knowledge about V_0 should be introduced in DCM analysis, given that we have demonstrated that the use of the physiologically realistic V_0 can improve the identifiability of the system. The incorporation of more accurate information (i.e., V_0 values) in the analysis procedure can be expected to yield more accurate information about brain connectivity.

In illustrating the importance of valid prior expectations, in terms of inferences about models and their parameters, we have used what is called a ‘Bayesian illusion’. In other words, we have deliberately generated data in a way that violates prior expectations to show that Bayesian inversion produces posterior beliefs that are wrong. This is the basis of most perceptual illusions, in which stimuli are generated in an implausible way to produce illusory percepts in subjects. Indeed, the psychophysics of illusions are a powerful way to demonstrate a subject’s prior beliefs. An interesting question here is which is the best belief? For example, in Fig. 6, if we did not know the true parameters or architecture generating the data, then is this architecture ‘wrong’? As shown by the Bayesian model comparison, this model is a better explanation for the data in the sense that it provides an accurate but more parsimonious explanation than the model used to generate the data. Clearly, this is a philosophical question but has practical relevance when dealing with real world analyses.

In presenting these analyses, we are not implying that DCM is invalid. We are trying to illustrate the importance - and potential usefulness - of using more informed prior expectations. As noted above, we are effectively reporting a Bayesian illusion by generating data that violates prior expectations to show that posterior beliefs can deviate from ground truth. However, to illustrate this failure we had to use data that was very implausible: although a change in V_0 from 0.04 to 0.02 may seem small, it corresponds a large departure from prior expectations. This is because it entails a doubling of the hemodynamic sensitivity to neuronal inputs. In other words, we have effectively multiplied the amplitude of one regional time-series in our simulations by 100%. When we repeated the analysis using a more plausible violation (of 10%), DCM was able to recover the correct model in most instances²⁴.

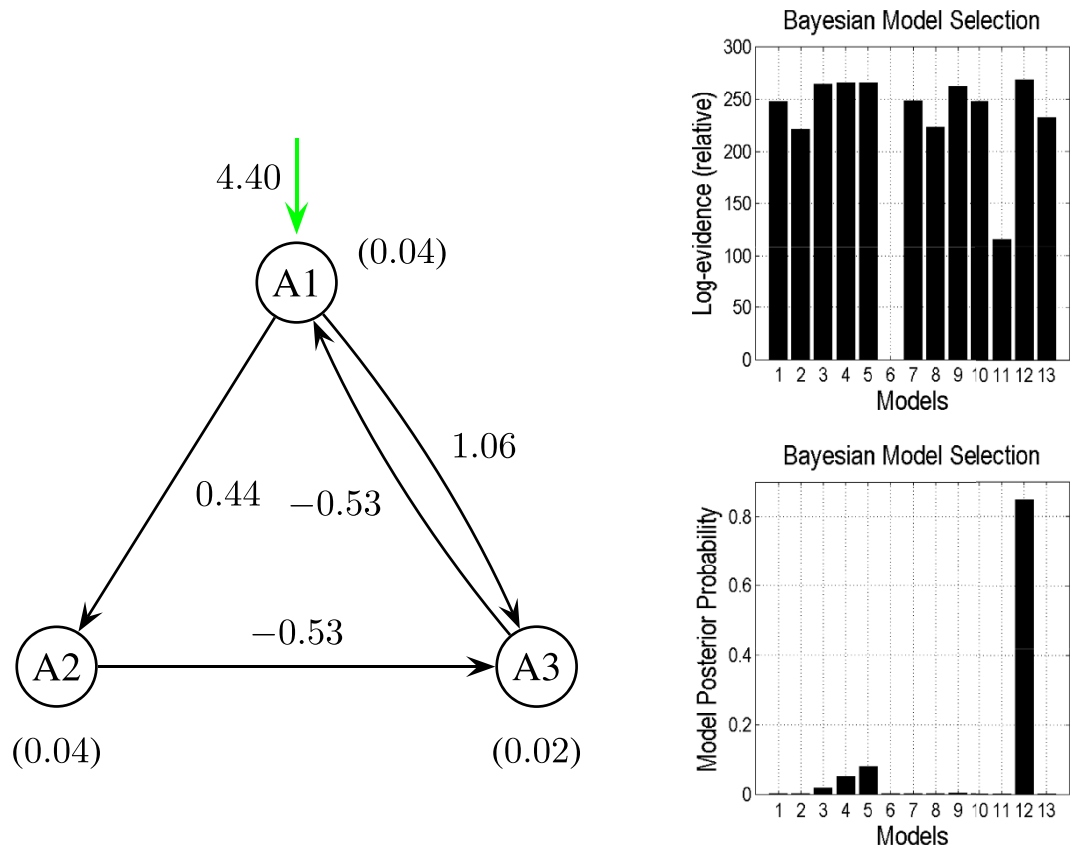


Figure 5. Same as Fig. 4 but with $V_0^{A1} = 0.04$, $V_0^{A2} = 0.04$, and $V_0^{A3} = 0.02$ illustrate that Model 12 is superior to the other models.

The key issue that we have highlighted rests upon the conditional dependence between uninteresting parameters with tight priors (e.g., V_0) and interesting parameters such as effective connectivity (that have relatively uninformative priors). The conditional dependency between posterior estimates means that improperly specified prior beliefs about V_0 can be expressed in terms of connectivity estimates. This behaviour is well-known in DCM and its parameterisation has been designed to attenuate conditional independencies. For example, most Bayesian model comparisons deal with experimental effects mediated by the coefficients of the B matrix in Fig. 1. Our simulations exploited the conditional dependency between V_0 and A , which is much less pronounced between V_0 and B . When we repeated the above simulations using a full connectivity (A) matrix but modelling an increase in the forward connection from A1 to A2 in the B matrix, DCM was able to recover the true model. This is because changing the V_0 parameter cannot reproduce the effects of changing a condition-specific connection (B parameter)¹.

We have used our results to illustrate the putative failure of DCM when violating its prior expectations (in terms of V_0). However, DCM can be used to infer the values of V_0 (either by treating it as a free parameter or using Bayesian model comparison). For example, if we combine the free energies across all three sets of 13 models in Figs 4–6, we can include different beliefs about V_0 in our model space. Figure 7, shows the result of this Bayesian model comparison (noting that the previous figures expressed the log evidence or free energy relative to the worst model within each set). Here we see that the winning model corresponds to the correct model under the correct values of V_0 . The more interesting application of Bayesian model comparison, in this context, is the use of family-wise comparisons to compare the three sets of V_0 parameters. In other words, we can use DCM to test different prior beliefs about V_0 and assess whether empirically informed priors are better than the uniform priors currently adopted. This device has been used to assess the improvement in model evidence afforded by priors on connectivity from probabilistic tractography data¹⁰. In principle, exactly the same approach can be used to test the improvement in model evidence provided by empirical estimates of V_0 . A promising development in MRI physics that may provide such measures (VasA) uses the spectral behaviour of residuals from any fMRI timeseries to estimate V_0 in a way that does not require any extra scans or data²⁵.

In the ideal situation, perfect prior knowledge of V_0 will make it possible for DCM analysis to reliably reconstruct the connections between brain areas. However, in practice, the V_0 value associated with the BOLD signal in a specific tissue region can not be determined. This has resulted in previous studies instead adopting an arbitrary V_0 value. We suggest that more useful information could be obtained using two other techniques that are also based on magnetic resonance: magnetic resonance angiography (MRA) imaging and cerebral blood volume (CBV) imaging. In the hemodynamic model, V_0 is the venous volume of blood present in a voxel; it represents the ratio of occupied vessels with sizes ranging from capillaries to large veins that all contribute to the fMRI

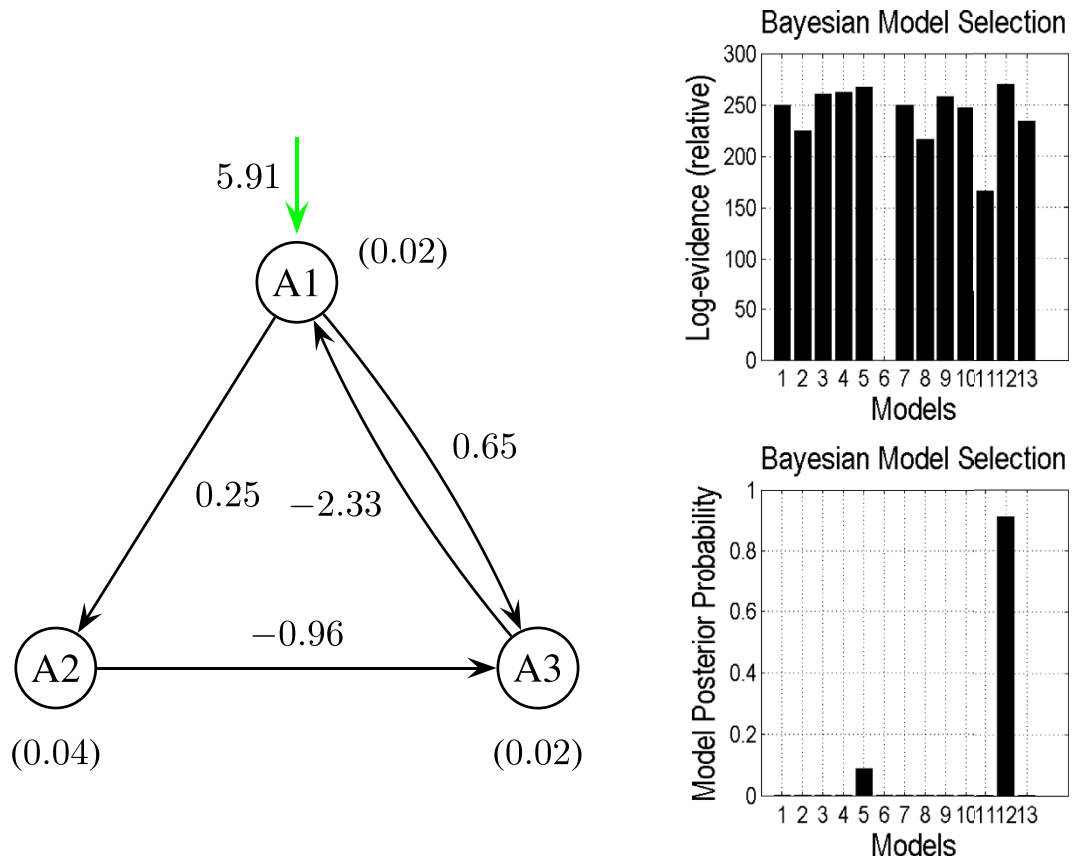


Figure 6. Same as Fig. 4 but with $V_0^{A1} = 0.02$, $V_0^{A2} = 0.04$, and $V_0^{A3} = 0.02$ illustrate that Model 12 is superior to the other models.

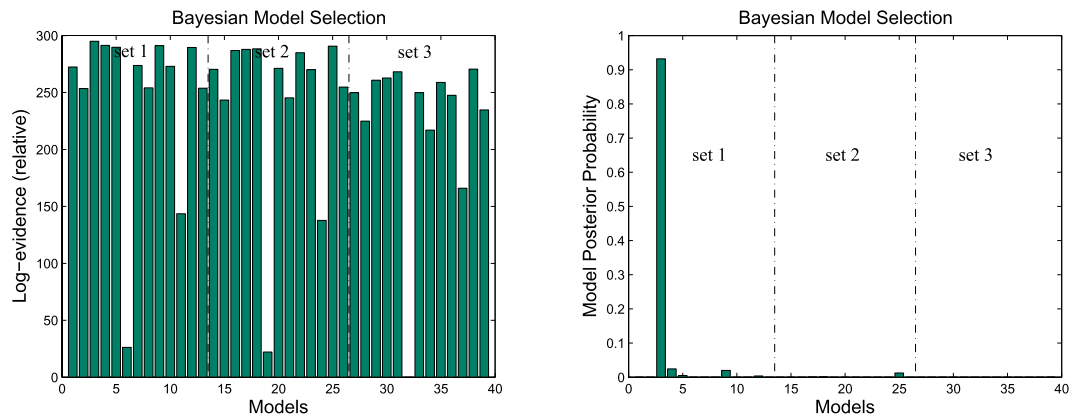


Figure 7. The comparisons of all $3 \times 13 = 39$ models for each of the 13 architectures under the three different sets of prior assumptions about V_0 . The correct model (Model 3) under the correct prior assumptions is superior. Set 1: $V_0^{A1} = 0.04$, $V_0^{A2} = 0.02$, $V_0^{A3} = 0.04$; Set 2: $V_0^{A1} = 0.04$, $V_0^{A2} = 0.04$, $V_0^{A3} = 0.02$; Set 3: $V_0^{A1} = 0.02$, $V_0^{A2} = 0.04$, $V_0^{A3} = 0.02$.

measurements in that area^{7,8}. MRA images (each of which has a spatial resolution of $0.9375 \times 0.9375 \times 1.5 \text{ mm}^3$ in a typical MRA protocol) provide much finer detail than fMRI images (which have a typical resolution of $3.75 \times 3.75 \times 6 \text{ mm}^3$). MRA may therefore provide additional information about large blood vessels (veins and venules with a radius $> \mu\text{m}$, depending on the magnetic-field intensity) that can be used to refine the V_0 value²². On the other hand, CBV imaging measures the volume of blood across arteries, capillaries, and veins in a more direct manner, and it provides a slightly overestimated value of V_0 due to the inclusion of arterial blood components^{26–28}. However, although the proposed methods have potential physiological meaning and attractive robustness, it is indeed difficult to determine if the combinative approach really present an improvement for

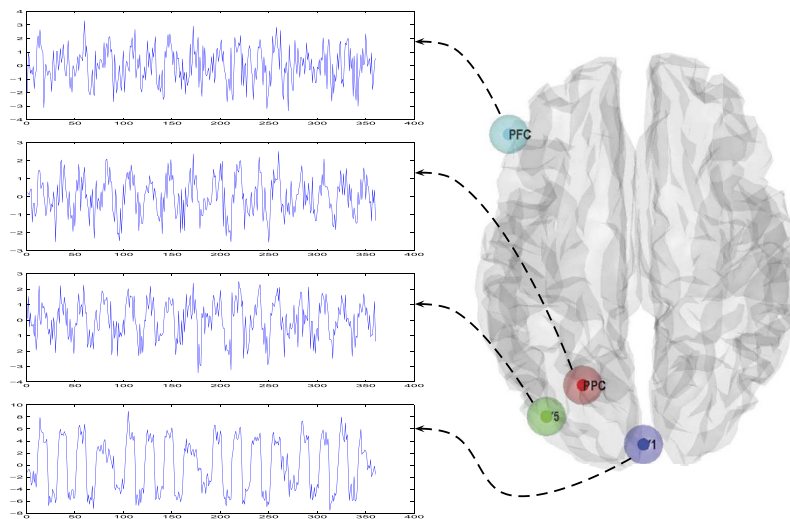


Figure 8. Location of ROI and their time series.

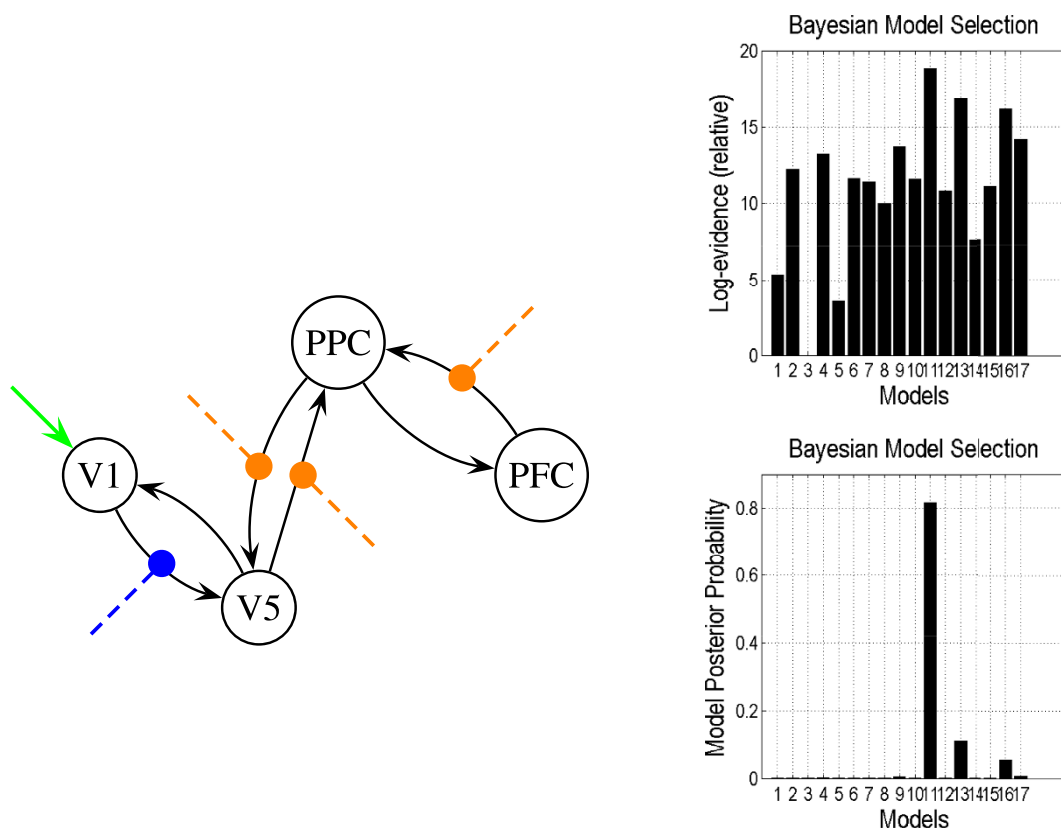


Figure 9. The most reasonable model based on the present observations with $V_0^{V1} = 0.02$, $V_0^{V5} = 0.02$, $V_0^{PPC} = 0.04$, and $V_0^{PFC} = 0.04$, selected from 17 competing models. The green line represents the extrinsic input, the blue line represents the modulation of motion by experimental manipulation, and the yellow line represents attentional modulation. The right graphs provide the results obtained in comparisons of the DCMs, illustrating that Model 11 is superior to the other models.

DCM analysis, because simultaneous fMRI and electrophysiological measurement are unavailable. In fact, such a concern exists in all effective connectivity studies (includes DCM and Granger causal analysis, GCA). It is a logical argument what obtaining more realistic V_0 information in DCM analysis would provide more reliable inferences about the properties of brain connectivity¹⁸. In our knowledge, there is the only study that compares causal connectivity determined from fMRI time series with actual neural coupling estimated from iEEG in a rat

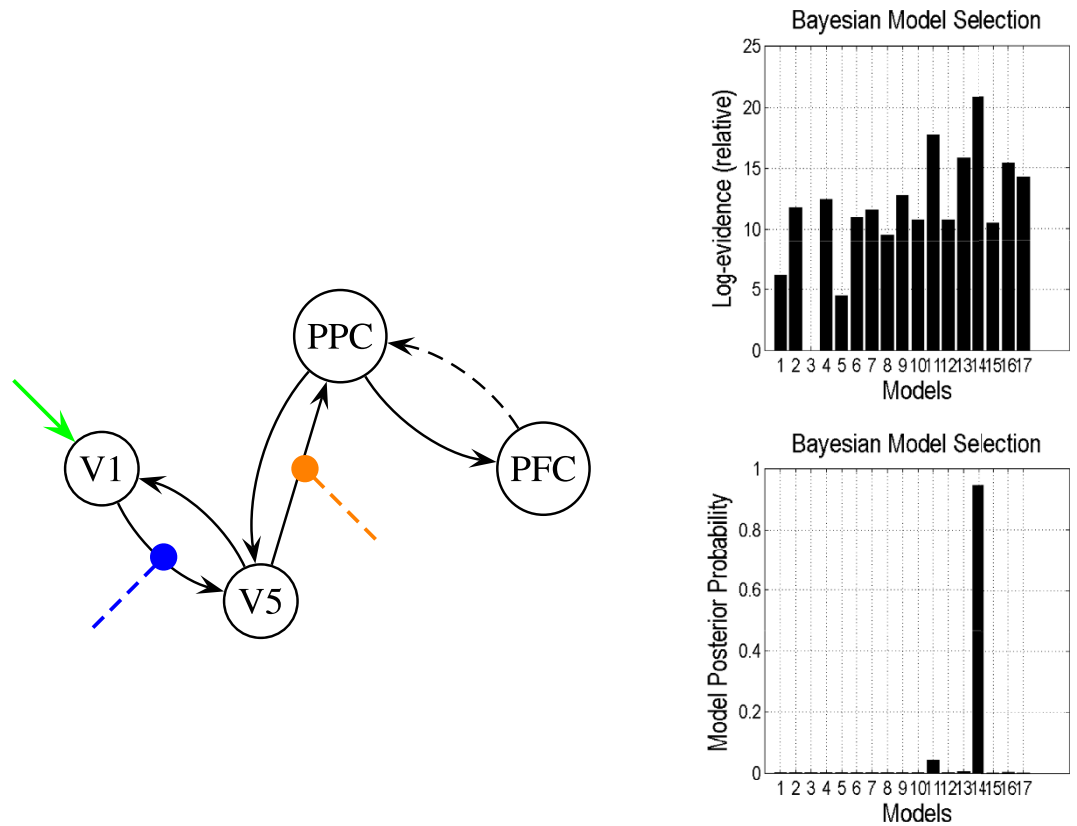


Figure 10. Same as Fig. 9, but with $V_0^{V1} = 0.02$, $V_0^{V5} = 0.04$, $V_0^{PPC} = 0.04$, and $V_0^{PFC} = 0.04$. The right graphs illustrate that Model 14 is superior to the other models.

model of absence epilepsy, where fMRI and iEEG measurements are recorded neither at the same time nor in the same subject²³. Currently, we are trying to design the psychological experiment to partly validate the effectiveness of proposed approach.

Recently there have been fairly extensive discussions on effective connectivity studies using neuroimaging methods^{29–33}. Although there is still controversy about which technique is the most suitable for addressing the causal correlation of neural system (e.g., DCM versus Granger causality analysis, GCA), there is broad consensus that hemodynamic model/deconvolution and model inference/selection are crucial to identifying the connections among distributed brain areas³⁴. DCM marries neurobiological and hemodynamic models and expresses causality at the level of neuronal dynamics that we are actually concerned with, whereas GCM operates at the level of the measured signals. The observed hemodynamic responses in the BOLD modality represent a secondary effect induced by neuronal activity. Different brain regions may exhibit marked differences in neurovascular coupling, and the associated difference in latencies, undershoots, and other features could lead to false inferences about connectivity. In this sense, DCM is superior to GCM. However, the use of nonlinear physiological modeling generally increases the modeling complexity. Our results suggest that the hemodynamic model parameters may be helpful for eliminating imaging bias but it may impair the overall system identifiability. In other words, the benefit of introducing the hemodynamic model parameters may be partly offset by difficulties in parameter estimation and conditional dependencies.

It also should be noted that the relationship between neuronal activity and BOLD signals is still being debated³⁵, with in particular it being unclear how neuronal activity in the hemodynamic model corresponds to physical mechanisms underlying the measured electro-physiological signals, such as obtained using EEG and MEG^{36,37}. This might be another factor in uncovering causality using the DCM analysis²³.

These findings could lead to future directions for DCM developments²⁹. Moreover, we would like to highlight the relevance of system identifiability to any causal inference about neural systems (not just DCM) using neuroimaging techniques. Although nonlinear modeling is indispensable for neural signal analysis, the nonlinearity of the signal does not always imply the validity of the application of a nonlinear model³⁸. For a low-time-resolution BOLD signal, nonlinear GCA only can provide a predictive performance comparable to that obtained by linear GCA³⁹, suggesting that it would be more efficient to apply a linear model when describing such real-world data sets. This is because nonlinear models are more complex and require a greater number of parameters than linear models, since the model efficiency depends on a trade-off between the complexity of the model and the accuracy of data fitting⁴⁰. In addition, from a pure mathematical perspective, DCM, GCA, and indeed scientific method used for making evidence-based inferences involve solving an inverse problem. They use the results of actual observations to infer model information (e.g., parameters, and architecture) characterizing the system under investigation. The internal mechanisms of the system always have a substantially higher order than the external

observations⁴¹. Although the modeling approach is closely linked to a reductionistic world view, the description of the system still requires a large and complex model⁴², and such a model is often poorly identifiable. In this situation it is hardly reasonable to expect that it will be possible to identify true information from the experimentally measured data^{43,44}. Instead, the task is to find biologically reasonable parameter values and modeling structure that together describe the data adequately, rather than providing true information⁴⁵. The goal of identification is to obtain some insight into the system. Therefore, despite these limitations, we still believe that applying the DCM technique to neuroimaging data can provide useful insight at least on a rational basis into the understanding of brain connectivity.

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Acknowledgements

This work is supported in part by the National Basic Research Program of China under Grant 2013CB329501, in part by the National High Technology Research and Development Program of China under Grant 2012AA011600, in part by the National Natural Science Foundation of China under Grant 81271645, in part by the Public Projects of Science Technology Department of Zhejiang Province under Grant 2013C33162, in part by the Zhejiang Provincial Natural Science Foundation of China under Grant LY12H18004, and in part by the Customer Experience Transformation Center (CETC), Huawei Technology Corporation Limited.

Author Contributions

Z.H. conceived of the study, guided its design and coordination, and drafted the manuscript. P.N. performed the data analysis and prepared Figures 3–6, 8–10. Q.W. and Y.Z. assisted with data analysis. P.S. and Q.L. assisted with results analysis and the drafting of the manuscript. All authors reviewed the manuscript.

Additional Information

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Hu, Z. *et al.* Influence of Resting Venous Blood Volume Fraction on Dynamic Causal Modeling and System Identifiability. *Sci. Rep.* **6**, 29426; doi: 10.1038/srep29426 (2016).



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