Variational Physiologically Informed Solution to Hemodynamic and Perfusion Response Estimation from ASL fMRI Data

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Abstract—Functional Arterial Spin Labeling (fASL) MRI can provide a quantitative measurement of cerebral blood flow. A joint detection-estimation (JDE) framework has been considered to extract task-related perfusion and hemodynamic responses not restricted to canonical response function shapes. In this work, we provide a variational expectation-maximization (VEM) algorithm for hemodynamic and perfusion responses estimation. This approach provides a lower computational load compared to previous attempts, and facilitates the incorporation of prior knowledge and constraints in the estimation. Validation on simulated and real data sets has been performed.

I. INTRODUCTION

Functional Arterial Spin Labeling (fASL) is a functional MRI modality that is able to provide a quantitative measurement of cerebral blood flow (CBF) and its variations elicited by specific tasks. Although ASL [1] is mainly used as a static probe of cerebral perfusion, it has also been used in functional MRI as an alternative to the standard blood-oxygen-level-dependent (BOLD) [2] modality, as it can give more specific information about brain function. FASL data consists of alternating pairs of control and magnetically tagged (“tag”) images. Local CBF or perfusion changes can be measured by doing the “control-tag” difference, so as to get rid of the hemodynamic component (BOLD effect) of the signal contained in both control and tag images. Many pairs (> 50) of control-tag images need to be acquired to compensate for the low Signal-to-Noise Ratio (SNR) of this difference.

The standard approach for fASL analysis is general linear model (GLM) [3], [4] using the stimulus-induced canonical hemodynamic response function (HRF) as regressor for both components of the signal: hemodynamic and perfusion components. The latter one encodes “control-tag” differences. Although the canonical HRF is generally used, it has been just calibrated on BOLD experiments and it reflects simultaneous variations of CBF, cerebral blood volume (CBV) and cerebral oxygen consumption (CMRO2). In contrast, the perfusion response function (PRF) only reflects the variation in CBF and thus departs from the canonical shape. Moreover, there has been evidence that this response can vary between regions and across subjects so that assuming a constant response shape might be misleading. Some flexibility in the response shape has already been considered in BOLD GLM analysis by using time/dispersion derivatives [5] or finite impulse response models.

In a joint detection-estimation (JDE) framework both task-related perfusion and hemodynamic responses can be estimated as well as perfusion-related and BOLD-related maps of evoked activity. JDE has been originally developed for BOLD data analysis [6], [7] and has been extended in [8] for fASL data analysis, considering Monte Carlo Markov Chains (MCMC) methods. In this work, following the spirit of [7], we provide an alternative solution based on the variational expectation-maximization (VEM) algorithm. This framework is more convenient to deal with constraints (e.g., normalization or positivity) in the M-step. JDE-VEM provides comparable results to JDE-MCMC for a much lower computational load. One of the novelties is the introduction of prior knowledge as in [9] through the relationship between perfusion and hemodynamic responses derived from physiological models [10], [11]. This relationship allows us to inform the PRF estimation from the HRF one, as the hemodynamic component has a higher SNR than the perfusion one due to the acquisition procedure. Experiments on simulated and real data show the good performance of the method, with a fast convergence of the parameter estimates.

II. JOINT DETECTION ESTIMATION MODEL FOR FASL

In fASL the magnetically tagged image reflects a perfusion effect besides the hemodynamic effect, that can be subtracted by comparison with the control image. The control image is a noisy BOLD signal because of the effect of the magnetic tagging. The different ASL signal components can be explained with the ASL JDE model [8], [12], a region-based approach that considers functionally homogeneous regions. In a region \( P \) comprising \( J \) voxels, the generative model for \( j \in J \):

\[
y_j = \sum_{m=1}^{M} a_{jm} X_{jm} h + \sum_{m} W_{j} X_{jm} g + \sum_{j} P_{j} + a_{j} w + b_{j} \quad (a) \quad (b) \quad (c) \quad (d) \quad (e)
\]

in the presence of \( M \) experimental conditions. The data \( y_j \in \mathbb{R}^N \) can be decomposed into (a) task-related hemodynamic...
and (b) perfusion components; (c) a drift component \( P \ell_j \); (d) a perfusion baseline term \( \alpha_j w_1 \) which completes the modelling of the perfusion component and (e) a noise term, assumed white Gaussian of variance \( \sigma_n^2 \). The control/tag effect is implicit in the ASL JDE model given the use of \( W = \text{diag}(w) \). Vectors \( h \) and \( g \) represent the \( D \)-dimensional \( (D < N) \) unknown HRF and PRF shapes, constant within \( \mathcal{P} \). The magnitudes of activation or response levels for hemodynamic and perfusion components are \( a = \{ a_m \} \) and \( c = \{ c_m \} \) and denoted as BRLs and PRLs hereafter. \( \mathbf{X} \in \mathbb{R}^{N \times D} \) is a binary matrix that encodes the lagged onset stimuli. The response levels are assumed to follow different spatial Gaussian mixture models but governed by \( M \) common binary hidden Markov random fields \( q_m \) with \( q_{m} = \{ q_{mj} \in \mathcal{P} \} \) encoding voxels’ activation states for each experimental condition \( m \). BRLs and PRLs are assumed independent conditionally to \( q = \{ q_m, m = 1 : M \} \). For further detail, please refer to [8].

The perfusion component in the ASL signal has a very low SNR owing to its small size captured by the “control-tag” subtraction. To address this issue, a conditional prior has been used to infer the PRF estimation from the less noisy HRF recovery during inference. To link these two responses, a relationship \( g = \Omega h \) was derived in [9] from physiological models. In our model, we consider that HRF and PRF shapes follow prior Gaussian distributions \( h \sim \mathcal{N}(0, \nu_h \Sigma_h) \) and \( g | h \sim \mathcal{N}(\Omega h, \nu_g \Sigma_g) \), with covariance matrices \( \Sigma_h \) and \( \Sigma_g \) encoding a constraint on the second order derivative so as to account for temporal smoothness. We also consider constraints on the response functions to enforce their \( L_2 \)-norm to be 1, i.e. \( h, g \in S^2(\mathbb{R}^{D+1}), S^2(\mathbb{R}^{D+1}) \) being the \( L_2 \) unit ball of \( \mathbb{R}^{D+1} \).

III. VARIATIONAL EM

In our fASL model, there are missing parameters that need to be estimated: \( a \in \mathcal{A}, h \in \mathcal{H}, c \in \mathcal{C}, g \in \mathcal{G}, q \in \mathcal{Q} \). In this work, an Expectation-Maximization (EM) framework is proposed to perform the estimation. EM can be viewed [13] as an alternating maximization procedure of a function \( F \) such that for any \( \tilde{p} \in \mathcal{D} \), being \( \mathcal{D} \) the set of all probability distributions on \( A \times H \times C \times G \times Q \),

\[
F(\tilde{p}, \theta) = E_{\tilde{p}} \left[ \log p(y, a, h, c, g, q; \theta) \right] + I(\tilde{p}).
\]

Here \( I(\tilde{p}) = -E_{\tilde{p}}[\log \tilde{p}(a, h, c, g, q)] \) is the entropy of \( \tilde{p} \), and \( E_{\tilde{p}}[\cdot] \) denotes the expectation with respect to \( \tilde{p} \). Maximizing function \( F \) is equivalent to minimizing the Kullback-Leibler divergence \( \text{KL}(\tilde{p}, p(a, h, c, g, q)) \), that gives a measure of the distance between two distributions, in this case the approximation \( \tilde{p} \) and the true \( p(a, h, c, g, q) \) distributions. This view of EM has led to a number of variants in which the E-step is solved over a restricted class of probability distributions, \( \mathcal{D} \). The variational approach corresponds to \( \mathcal{D} \) chosen as the set of distributions that factorize as \( \tilde{p}(a, h, c, g, q) = \tilde{p}_a(a) \tilde{p}_h(h) \tilde{p}_c(c) \tilde{p}_g(g) \tilde{p}_q(q) \) where \( \tilde{p}_a \in \mathcal{D}_A, \tilde{p}_h \in \mathcal{D}_H, \tilde{p}_c \in \mathcal{D}_C, \tilde{p}_g \in \mathcal{D}_G \) and \( \tilde{p}_q \in \mathcal{D}_Q \), the sets of probability distributions on \( A, H, C, G, Q \) respectively. Note that the dependence between random variables, as in MCMC, is translated into dependence between statistical moments in VEM. The E-step becomes an approximate E-step that can be further decomposed into five stages updating the different variables in turn. At iteration \( (r) \), with current estimates denoted by \( \tilde{p}_r \), \( \tilde{p}_h^{(r-1)}, \tilde{p}_c^{(r-1)}, \tilde{p}_g^{(r-1)}, \tilde{p}_q^{(r-1)} \) and \( \theta^{(r)} \), the updating formulae are of the form:

**E-H-step:**

\[
\tilde{p}_h^{(r)} = \arg \max_{\tilde{p}_h \in \mathcal{D}_H} F(\tilde{p}_a^{(r-1)}, \tilde{p}_h^{(r-1)}, \tilde{p}_c^{(r-1)}, \tilde{p}_g^{(r-1)}; \theta^{(r)}) \tag{2}
\]

**E-G-step:**

\[
\tilde{p}_g^{(r)} = \arg \max_{\tilde{p}_g \in \mathcal{D}_G} F(\tilde{p}_a^{(r-1)}, \tilde{p}_h^{(r)}, \tilde{p}_c^{(r-1)}, \tilde{p}_g^{(r-1)}; \theta^{(r)}) \tag{3}
\]

with similar expressions for the other steps obtained by permuting the roles of the variables. Hereafter, for the ease of presentation, the \( (r) \) and \( (r - 1) \) superscripts are omitted. We also consider normalization constraints on \( h \) and \( g \) so that our solution has unit \( L_2 \)-norm. For that matter, we modify the seek variational approximation to \( \tilde{p} = \tilde{p}_a \delta_h \tilde{p}_c \delta_g \tilde{p}_q \), where the probabilities on \( h \) and \( g \) are replaced by Dirac functions. This reduces the search to pointwise estimates \( \hat{h} \) and \( \hat{g} \). The E-H and E-G steps in Eqs. (2)-(3) yield then maximizations problems which are easily constrained to account for normalization:

**E-H:**

\[
\hat{h} = \arg \max_{h} E_{\tilde{p}_a, \hat{g}, \hat{c}} \left[ \log p(h | y, a, c, \hat{g}, q; \theta) \right] \tag{4}
\]

**E-G:**

\[
\hat{g} = \arg \max_{g} E_{\tilde{p}_a, \hat{h}_r, \hat{c}} \left[ \log p(g | y, a, \hat{h}, c, q; \theta) \right] \tag{5}
\]

It is straightforward to see that (4) and (5) amount to minimizing a quadratic function under a quadratic constraint, namely \( |h|^2 = 1 \) and \( |g|^2 = 1 \) respectively. The other E-steps can be derived from standard expressions replacing expectations over \( h \) and \( g \) by \( \hat{h} \) and \( \hat{g} \), e.g.:

**E-Q:**

\[
\tilde{p}_q(q) \propto \exp \left( E_{\tilde{p}_a, \hat{h}, \hat{c}, \hat{g}} \left[ \log p(q | y, a, \hat{h}, \hat{c}, \hat{g}; \theta) \right] \right),
\]

with similar expressions for the E-A and E-C steps obtained by permutations of the variables. BRLs and PRLs have Gaussian distributions as the likelihood and the priors are Gaussian too. Regarding the labels, a further factorization is needed:

\[
\tilde{p}_{q_{mj}}(q_{mj}) = \prod_{j=1}^{M} \tilde{p}_{q_{mj}}(q_{mj}).
\]

The labels depend on the BRLs and PRLs as well as on the binary activation states \( q_{mj} \).

The corresponding M-step is given by:

**M:**

\[
\theta = \arg \max_{\theta \in \Theta} \left[ E_{\tilde{p}_a, \hat{h}, \hat{g}} \left[ \log p(y | a, \hat{h}, c, \hat{g}; \alpha, \ell, \sigma^2) \right] \right. \]  
+ \left. \log p(h; \nu_h) + \log p(g; \nu_g) \right. \]  
+ \left. E_{\tilde{p}_a, \hat{h}} \left[ \log p(a | q; \mu_a, \sigma_a) \right] \right. \]  
+ \left. E_{\tilde{p}_a, \hat{g}} \left[ \log p(c | q; \mu_c, \sigma_c) \right] \right. \]  
+ \left. E_{\tilde{p}_a, \hat{g}} \left[ \log p(g; \beta) \right] \right]
\]

where \( \theta = \{ \alpha, \ell, \sigma^2, \mu_a, \sigma_a, \mu_c, \sigma_c, v_h, v_g, \beta \} \). Given the separability of the prior probability density functions, the M-step can be divided into separate M-steps, as in [7]. Note here
that a mean-field approximation is used to compute the MRF parameter $\beta$. For further details on these aspects, the reader is invited to refer to [7].

IV. RESULTS

Different data sets have been analysed to test the performance of this algorithm. First, some artificial data has been generated with the ASL JDE generative model. Then, real data acquired on different individuals from the AINSI initiative\(^2\) have been analysed to validate the proposed approach.

A. Artificial data

$N = 288$ ASL artificial images (i.e. 144 control/tag pairs) have been simulated using a realistic low SNR according to the observation model in Eq. (1). Different levels of SNR have been used, in order to show the performance of the method depending on the noise level. To emulate the slow sampling rate of ASL images, Eq. (1) was synthesized at $\Delta t = 0.5\text{s}$ and then down-sampled at a certain time of repetition (TR), which means that the temporal resolution of rows (TR) and columns ($\Delta t$) of $X^m$ is different. Here, we considered a fast event-related paradigm comprising two conditions ($M = 2$), with mean ISI = 5s. TR = 3s. is considered as a realistic ASL experiment, compared to the TR = 1s. that could be used for a realistic experiment when using BOLD signal.

In the experiments, $h$ and $g$ are generated as depicted in Fig 2(a)-(b) by dashed lines. $P$ is a polynomial basis of order $O = 4$. Drift coefficients and noise realizations were drawn according to $\ell_j \sim \mathcal{N}(0,10I_D)$ and $b_j \sim \mathcal{N}(0,2I_N)$, respectively. BRLs were sampled from $(a_j^m|q_j^m = 1) \sim \mathcal{N}(2.2,0.3)$ (for activating voxels) and from $(a_j^m|q_j^m = 0) \sim \mathcal{N}(0,0.3)$ (for inactivating voxels). PRLs were generated with a lower contrast than BRLs: $(c_j^m|q_j^m = 1) \sim \mathcal{N}(1.6,0.3)$ and $(c_j^m|q_j^m = 0) \sim \mathcal{N}(0,0.3)$. PRLs and BRLs were chosen so as to make this synthetic setting realistic: PRLs lower than BRLs, and activating/non-activating voxels distribution means close. Activation states (assignment variables $Q$) were set by a hand-drawn map.

\(\text{Fig. 2. Artificial data: Ground-truth response curves (black dashed lines) and estimated perfusion (a) and hemodynamic (b) response functions. The first and second rows correspond to the response functions for TR = 3s and noise variances 1 (i.e. SNR = 2.4 dB, top) and 6 (SNR = 0.5 dB, bottom).}\)

\(\text{Fig. 3. Results on artificial data for response levels. Top row: ground-truth maps. Center and bottom rows: estimated perfusion (a) and hemodynamic (b) response functions. (a) Hemodynamic response levels. (b) Perfusion response levels. (c) Assignment variables (activation states).}\)

Fig. 1 shows the root mean squared error (RMSE) of the estimated PRFs and HRFs for experiments generated with noise variances from 1 to 6, which corresponds to decreasing the SNR from 2 down to 0.5 dB. Curves are depicted for TR = 1s and TR = 3s, so as to show the impact of increasing TR on the performance of the method and thus mimicking part of the increased difficulty when moving from BOLD to ASL fMRI data. For TR = 1s, we observed as expected good HRF and PRF estimates with an accurate recovery of both peaks (Fig. 2 top), as well as good BRLs and PRLs map estimates (Fig. 3 center). In terms of RMSE (Fig. 1), the values are pretty close to each other although slightly larger for the PRF. However, in the case of TR = 3s, we recovered a much better PRF estimate owing to a large undershoot in the HRF shapes for very low SNR scenarios (Fig. 2 bottom right). For the worse SNR scenario, we obtained over-smoothed shapes, with peaks displaced compared to the ground truth.

\(\text{http://thalie.ujf-grenoble.fr/ainsi}\)
This impacts the recovery of the hemodynamic and perfusion activation maps too (see Fig. 3 bottom vs center), much worse in this case. Figs. 1-2 show that despite the dependence on the SNR level, our shape estimates are pretty reliable.

**B. Real data**

Real ASL data were recorded during an experiment designed to map auditory and visual brain functions, which consisted of $N = 291$ scans lasting $TR = 3$ s, with $TE = 18$ ms, FoV 192 mm, each yielding a 3-D volume composed of $64 \times 64 \times 22$ voxels (resolution of $3 \times 3 \times 7$ mm$^3$). The tagging scheme used was PICORE Q2T, with TI $= 700$ ms, $T12 = 1700$ ms. The paradigm was a fast event-related design (mean $ISI = 5.1$ s) comprising sixty auditory and visual stimuli. Ward parcellations were computed in auditory and visual cortices for regionally analysing the evoked activity elicited by auditory and visual stimuli, respectively.

The estimation results are depicted in Fig. 4. First, it is worth noting that as expected we recovered BOLD response levels with larger magnitude compared to that of the perfusion response levels (Fig. 4(a)-(b)). Interestingly, in response to auditory stimuli, we retrieved bilateral evoked activity in the auditory cortices for the BOLD and perfusion components, although not exactly in the same parcels. Subsequent thresholding of individual posterior probability maps should allow us to derive reliable statistical maps. Besides, we also recovered plausible shapes for BRF and PRF estimates, as illustrated in Fig. 4(c), where we selected a parcel corresponding to high BRL values. As expected, the HRF shape in the primary auditory cortex (left hemisphere) is well captured by the canonical shape. The physiological prior helps the parameter estimation, in the perfusion component and enforces the temporal precedence of the PRF estimate over the HRF one.

**V. DISCUSSION**

We proposed a variational Expectation-Maximization algorithm to address the issue of jointly detecting activity and estimating hemodynamic and perfusion responses from functional ASL data. The variational approach has the advantage to provide estimations in analytic form for each variable of interest. It facilitates the inclusion of additional information from physiological models and the incorporation of constraints on the responses to favor stability in the estimations. In particular, we considered a physiologically informed link between normalized hemodynamic and perfusion responses so as to compensate the low signal-to-noise ratio of the perfusion component. The results obtained on simulated and real data indicate a good performance of the approach and suggest the variational approach as a potential robust, fast and pragmatic method to analyse challenging ASL data. Addressing perfusion quantification is then the main perspective of this work as providing a good estimation of perfusion modulation effects might translate into more accurate quantification of perfusion.

**REFERENCES**


