Hemodynamic Deconvolution Demystified: Sparsity-Driven Regularization at Work

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ABSTRACT

Deconvolution of the hemodynamic response is an important step to access short timescales of brain activity recorded by functional magnetic resonance imaging (fMRI). Albeit conventional deconvolution algorithms have been around for a long time (e.g., Wiener deconvolution), recent state-of-the-art methods based on sparsity-pursuing regularization are attracting increasing interest to investigate brain dynamics and connectivity with fMRI. This technical note revisits the main concepts underlying two main methods, paradigm free mapping and total activation, in the most accessible way. Despite their apparent differences in the formulation, these methods are theoretically equivalent as they represent the synthesis and analysis sides of the same problem, respectively. We demonstrate this equivalence in practice with their best-available implementations using both simulations, with different signal-to-noise ratios, and experimental fMRI data acquired during a motor task and resting state. We evaluate the parameter settings that lead to equivalent results and showcase the potential of these algorithms compared to other common approaches. This note is useful for practitioners interested in gaining a better understanding of state-of-the-art hemodynamic deconvolution and aims to answer questions that practitioners often have regarding the differences between the two methods.

Keywords: fMRI deconvolution, paradigm free mapping, total activation, temporal regularization

INTRODUCTION

Functional magnetic resonance imaging (fMRI) data analysis is often directed to identify and disentangle the neural processes that occur in different brain regions during task or at rest. As the blood oxygenation level-dependent (BOLD) signal of fMRI is only a proxy for neuronal activity mediated through neurovascular coupling, an intermediate step that estimates the activity-inducing signal, at the timescale of fMRI, from the BOLD time-series can be useful. Conventional analysis of task fMRI data relies on the general linear models (GLM) to establish statistical parametric maps of brain activity by regression of the empirical timecourses against hypothetical ones built from the knowledge of the experimental paradigm. However, timing information of the paradigm can be unknown, inaccurate, or insufficient in some scenarios such as naturalistic stimuli, resting state, or clinically relevant assessments.

Deconvolution and methods alike are aiming to estimate neuronal activity by undoing the blurring effect of the hemodynamic response, characterized as a hemodynamic response function (HRF).\textsuperscript{1} Given the inherently ill-posed nature of hemodynamic deconvolution, due to the strong temporal low-pass characteristics of the HRF, the key is to introduce additional constraints in the estimation problem that are typically expressed as regularizers. For instance, the so-called Wiener deconvolution

\footnotesize{\textsuperscript{1} Note that the term deconvolution is also alternatively employed to refer to the estimation of the hemodynamic response shape assuming a known activity-inducing signal or neuronal activity (1–4).}
is expressing a “minimal energy” constraint on the deconvolved signal and has been used in the framework of psychophysiological interaction analysis to compute the interaction between a seed’s activity-inducing timecourse and an experimental modulation (5–9). Complementarily, the interest in deconvolution has increased to explore time-varying activity in resting-state fMRI data (10–13). In that case, the aim is to gain better insights of the neuronal signals that drive functional connectivity at short time scales, as well as learning about the spatiotemporal structure of functional components that dynamically construct resting-state networks and their interactions (14).

Deconvolution of the resting-state fMRI signal has illustrated the significance of transient, sparse spontaneous events (15,16) that refine the hierarchical clustering of functional networks (17) and reveal their temporal overlap based on their signal innovations not only in the human brain (18), but also in the spinal cord (19). Similar to task-related studies, deconvolution allows to investigate modulatory interactions within and between resting-state functional networks (20,21). In addition, decoding of the deconvolved spontaneous events allows to decipher the flow of spontaneous thoughts and actions across different cognitive and sensory domains while at rest (18,22,23). Beyond findings on healthy subjects, deconvolution techniques have also proven their utility in clinical conditions to characterize functional alterations of patients with a progressive stage of multiple sclerosis at rest (24), to find functional signatures of prodromal psychotic symptoms and anxiety at rest on patients suffering from schizophrenia (25), to detect the foci of interictal events in epilepsy patients without an EEG recording (17,26), or to study functional dissociations observed during nonrapid eye movement sleep (27).

The algorithms for hemodynamic deconvolution can be classified based on the assumed hemodynamic model and the optimization problem used to estimate the neuronal-related signal. Most approaches assume a linear time-invariant model for the hemodynamic response that is inverted by means of variational (regularized) least-squares estimators (5,6,28–36), logistic functions (37–39), probabilistic mixture models (40), convolutional autoencoders (41), or nonparametric homomorphic filtering (42). Alternatively, several methods have also been proposed to invert nonlinear models of the neuronal and hemodynamic coupling (43–49).

Among the variety of approaches, those based on regularized least-squares estimators have been employed more often due to their appropriate performance at small spatial scales (e.g., voxelwise). Relevant for this work, two different formulations can be established for the regularized least-squares deconvolution problem, either based on a synthesis- or analysis-based model (50,51). The rationale of the synthesis-based model is that we know or suspect that the true signal (here, the neurally driven BOLD component of the fMRI signal) can be represented as a linear combination of predefined patterns or dictionary atoms (for instance, the HRF). In contrast, the analysis-based approach considers that the true signal is analyzed by some relevant operator and the resulting signal is small (i.e., sparse).

As members of the groups that developed deconvolution-free mapping (PFM; synthesis based, solved with regularized least-squares estimators such as ridge regression (28) or LASSO (30)) and total activation (TA; analysis based, also solved with a regularized least-squares estimator using generalized total variation (52,33)) deconvolution methods for fMRI data analysis, we are often contacted by researchers who want to know about the similarities and differences between the two methods and which one is better. It depends—and to clarify this point, this note revisits synthesis- and analysis-based deconvolution methods for fMRI data and comprises four sections. First, we present the theory behind these two deconvolution approaches based on regularized least-squares estimators that promote sparsity: PFM (30)—available in AFNI as 3dPFM² and 3dMEPFM³ for single-echo and multi-echo data, respectively—and TA (33)—available as part of the iCAPs toolbox.⁴ We describe the similarities and differences in their analytical formulations, and how they can be related to each other. Next, we assess their performance controlling for a fair comparison on simulated and experimental data. Finally, we discuss their benefits and shortcomings and conclude with our vision on potential extensions and developments.

**THEORY**

**Notations and Definitions**

Matrices of size $N$ rows and $M$ columns are denoted by boldface capital letters, e.g., $X \in \mathbb{R}^{N \times M}$, whereas column vectors of length $N$ are denoted as boldface lowercase letters, e.g., $x \in \mathbb{R}^N$. Scalars are denoted by lowercase letters, e.g., $k$. Continuous functions are denoted by brackets, e.g., $h(t)$, while discrete functions are denoted by square brackets, e.g., $x[k]$. The Euclidean norm of a matrix $X$ is denoted as $||X||_2$, the $\ell_p$-norm is denoted by $||X||_p$, and the Frobenius norm is denoted by $||X||_F$.

The discrete integration ($L$) and difference ($D$) operators are defined as:

$$
L = \begin{bmatrix}
1 & 0 & \cdots \\
0 & 1 & 0 & \cdots \\
\vdots & \ddots & \ddots & \ddots \\
\end{bmatrix},
D = \begin{bmatrix}
1 & 0 & \cdots \\
-1 & 1 & 0 & \cdots \\
\vdots & \ddots & \ddots & \ddots \\
0 & \cdots & \cdots & -1 \\
\end{bmatrix}.
$$

⁴ https://c4science.ch/source/iCAPs/.
Conventional General Linear Model Analysis

Conventional general linear model (GLM) analysis puts forward a number of regressors incorporating the knowledge about the paradigm or behavior. For instance, the timing of epochs for a certain condition can be modeled as an indicator function \( p(t) \) (e.g., Dirac functions for event-related designs or box-car functions for block-designs) convolved with the HRF \( h(t) \) and sampled at TR resolution (53–56):

\[
x(t) = P \times h(t) \rightarrow x[k] = P \times h(k \cdot TR).
\]

The vector \( x = [x[k]]_{k=1,...,N} \in \mathbb{R}^N \) then constitutes the regressor modeling the hypothetical response, and several of them can be stacked as columns of the design matrix \( X = [x, \ldots, x] \in \mathbb{R}^{N \times L} \), leading to the well-known GLM formulation:

\[
y = X\beta + e,
\]

where the empirical timecourse \( y \in \mathbb{R}^N \) is explained by a linear combination of the regressors in \( X \) weighted by the parameters in \( \beta \in \mathbb{R}^L \) and corrupted by additive noise \( e \in \mathbb{R}^N \). Under independent and identically distributed Gaussian assumptions of the latter, the maximum likelihood estimate of the parameter weights reverts to choosing \( \hat{\beta} = (X^T X)^{-1} X^T y \), which is equivalent as fitting the measurements using the atoms of the columns of \( H \) in a least-squares sense, the optimization problem to solve is given by

\[
\hat{s} = \text{argmin}_s \frac{1}{2} \| y - Hs \|_2^2 + \Omega(s)
\]

(3)

The first term quantifies data fitness, which can be justified as the log-likelihood term derived from Gaussian noise assumptions, while the second term \( \Omega(s) \) brings in regularization and can be interpreted as a prior on the activity-inducing signal. For example, the \( \ell_1 \)-norm of \( s \) (i.e., \( \Omega(s) = \lambda_1 |s|_1 \)) is imposed for ridge regression or Wiener deconvolution, which introduces a tradeoff between the data fit term and “energy” of the estimates that is controlled by the regularization parameter \( \lambda \). Other well-known regularized terms are related to the elastic net (i.e., \( \Omega(x) = \lambda_1 |x|_1 + \lambda_2 |x|_2 \)) (58).

Paradigm Free Mapping

In PFM, the formulation of Eq. (3) was considered equivalently as fitting the measurements using the atoms of the HRF (i.e., columns of \( H \)) with corresponding weights (entries of \( s \)). This model corresponds to a synthesis formulation. In Gaudes et al. (30) a sparsity-pursuing regularization term was introduced on \( s \), which, in a strict way, reverts to choosing \( \Omega(s) = \lambda |s|_1 \) as the regularization term and solving the optimization problem (59). However, finding the optimal solution to the problem demands an exhaustive search across all possible combinations of the columns of \( H \). Hence, a pragmatic solution is to solve the convex-relaxed optimization problem for the \( \ell_1 \)-norm, commonly known as Basis Pursuit Denoising (60) or equivalently as the least absolute shrinkage and selection operator (LASSO) (61):

\[
\hat{s} = \text{argmin}_s \frac{1}{2} \| y - Hs \|_2^2 + \lambda |s|_1,
\]

(4)

which provides fast convergence to a global solution. Imposing sparsity on the activity-inducing signal implies that it is assumed to be well represented by a reduced subset of few nonzero coefficients at the fMRI timescale, which in turn trigger event-related BOLD responses. Hereinafter, we refer to this assumption as the spike model. However, even if PFM was developed as a spike model, its formulation in Eq. (4) can be extended to estimate the innovation signal, i.e., the derivative of the activity-inducing signal, as shown in Section 2.5.

Total Activation

Alternatively, deconvolution can be formulated as if the signal to be recovered directly fits the measurements and at the same time satisfies some suitable regularization, which leads to

\[
\hat{x} = \text{argmin}_x \frac{1}{2} \| y - x \|_2^2 + \Omega(x).
\]

(5)
Under this analysis formulation, total variation (TV), i.e., the $\ell_1$-norm of the derivative $\Omega(x) = \lambda \|Dx\|_1$, is a powerful regularizer since it favors recovery of piecewise-constant signals (62). Going beyond, the approach of generalized TV introduces an additional differential operator $D_n$ in the regularizer that can be tailored as the inverse operator of a linear system (52), that is, $\Omega(x) = \lambda \|DD_n x\|_1$. In the context of hemodynamic deconvolution, TA is proposed for which the discrete operator $D_n$ is derived from the inverse of the continuous-domain linearized Balloon-Windkessel model (63,64). The interested reader is referred to (33,52,65) for a detailed description of this derivation.

Therefore, the solution of the TA problem

$$\hat{x} = \text{argmin}_x \frac{1}{2} \| y - x \|^2 + \lambda \| DD_n x \|_1 \tag{6}$$

will yield the activity-related signal $x$ for which the activity-inducing signal $s = D_s x$ and the so-called innovation signal $u = D_s$, i.e., the derivative of the activity-inducing signal, will also be available, as they are required for the regularization. We refer to modeling the activity-inducing signal based on the innovation signal as the block model. Nevertheless, even if TA was originally developed as a block model, its formulation in Eq. (6) can be made equivalent to the spike model as shown in Section 2.5.

**Unifying Both Perspectives**

PFM and TA are based on the synthesis- and analysis-based formulation of the deconvolution problem, respectively. They are also tailored for the spike and block model, respectively. In the first case, the recovered deconvolved signal is synthesized to be matched to the measurements, while in the second case, the recovered signal is directly matched to the measurements but needs to satisfy its analysis in terms of deconvolution. This also corresponds to using the forward or backward model of the hemodynamic system, respectively. Hence, it is possible to make both approaches equivalent (50).\(^5\)

To start with, TA can be made equivalent to PFM by adapting it for the spike model; i.e., when removing the derivative operator $D$ of the regularizer in Eq. (6), it can be readily verified that replacing in that case $x = Hs$ leads to identical equations and thus both assume a spike model, since $H$ and $D_n$ will cancel out each other (52).\(^6\)

Conversely, the PFM spike model can also accommodate the TA block model by modifying Eq. (4) with the forward model $y = HL u + e$. Here, the activity-inducing signal $s$ is rewritten in terms of the innovation signal $u$ as $s = Lu$, where the matrix $L$ is the first-order integration operator (34,66). This way, PFM can estimate the innovation signal $u$ as follows,

$$\hat{u} = \text{argmin}_u \frac{1}{2} \| y - HL u \|^2 + \lambda \| u \|_1 \tag{7}$$

and becomes equivalent to TA by replacing $u = DD_n x$, and thus adopting the block model. Based on the previous Eqs. (4), (6), and (7), it is clear that both PFM and TA can operate under the spike and block models, providing a convenient signal model according to the different assumptions of the underlying neuronal-related signal. This work evaluates the core of the two techniques, i.e., the regularized least-squares problem with temporal regularization without considering the spatial regularization term originally incorporated in TA. For the remainder of this paper, we will use the PFM and TA formalisms with both spike and block models.

**Algorithms and Parameter Selection**

Despite their apparent resemblance, the practical implementations of the PFM and TA methods proposed different algorithms to solve the corresponding optimization problem and select an adequate regularization parameter $\lambda$ (30,33). The PFM implementation available in AFNI employs the least angle regression (LARS) (67), whereas the TA implementation uses the fast iterative shrinkage-thresholding algorithm (FISTA) (68). The blue box in Figure 1 provides a descriptive view of the iterative modus operandi of the two algorithms.

On the one hand, LARS is a homotopy approach that computes all the possible solutions to the optimization problem and their corresponding value of $\lambda$, i.e., the regularization path, and the solution according to the Bayesian Information Criterion (BIC) (69), was recommended as the most appropriate in the case of PFM approaches since Akaike Information Criteria (AIC) often tends to overfit the signal (30,31).

On the other hand, FISTA is an extension of the classical gradient algorithm that provides fast convergence for large-scale problems. In the case of FISTA though, the regularization parameter $\lambda$ must be selected prior to solving the problem but can be updated in every iteration so that the residuals of the data fit converge to an estimated noise level of the data:

$$\lambda^{n+1} = \frac{N\hat{\sigma}}{2 \| y - x^n \|^2} \lambda^n \tag{8}$$

where $x^n$ is the $n$th iteration estimate, $\lambda^n$ and $\lambda^{n+1}$ are the $n$th and $(n + 1)$th iteration values for the regularization parameter $\lambda$, and $N$ is the number of points in the timecourse. The preestimated noise level can be obtained as the median absolute deviation (MAD) of the fine-scale wavelet coefficients (Daubechies, order 3) of the fMRI timecourse. The MAD criterion has been adopted in TA (33). Of note, similar formulations based on the MAD estimate have also been applied in PFM formulations (29,70).

\(^5\) Without dwelling into technicalities, for total variation, this equivalence is correct up to the constant, which is in the null space of the derivative operator.

\(^6\) Again, this holds up to elements of the null space.
Fig. 1. Flowchart detailing the different steps of the fMRI signal and the deconvolution methods described. The orange arrows indicate the flow to estimate the innovation signals, i.e., the derivative of the activity-inducing signal. The blue box depicts the iterative modus operandi of the two algorithms used in this paper to solve the PFM and total activation (TA) deconvolution problems. The plot on the left shows the regularization path obtained with the least angle regression (LARS) algorithm, where the x-axis illustrates the different iterations of the algorithm, the y-axis represents points in time, and the color describes the amplitude of the estimated signal. The middle plot depicts the decreasing values of $\lambda$ for each iteration of LARS as the regularization path is computed. The green and red dashed lines in both plots illustrate the Bayesian Information Criterion (BIC) and median absolute deviation (MAD) solutions, respectively. Comparatively, the changes in $\lambda$ when the fast iterative shrinkage-thresholding algorithm (FISTA) method is made to converge to the MAD estimate of the noise are shown on the right. Likewise, the $\lambda$ corresponding to the BIC and MAD solutions are shown with dashed lines.

METHODS

Simulated Data

To compare the two methods while controlling for their correct performance, we created a simulation scenario that can be found in the GitHub repository shared in Section 6. For the sake of illustration, we describe here the simulations corresponding to a timecourse with a duration of 400 seconds (TR = 2 s) where the activity-inducing signal includes 5 events, which are convolved with the canonical HRF. Different noise sources (physiological, thermal, and motion related) were also added, and we simulated three different scenarios with varying signal-to-noise ratios (SNR = 20, 10, 3 dB) that represent high, medium, and low contrast-to-noise ratios as shown in Figure 2A. Noise was created following the procedure in (30) as the sum of uncorrelated Gaussian noise and sinusoidal signals to simulate a realistic noise model with thermal noise, cardiac and respiratory physiological fluctuations, respectively. The physiological signals were generated as

$$\sum_{i=1}^{2} \frac{1}{2} \left( \sin(2\pi f_t t + \phi) + \sin(2\pi f_r t + \phi) \right),$$

with up to second-order harmonics per cardiac ($f_c$) and respiratory ($f_r$) component that were randomly generated following normal distributions with variance 0.04 and mean $i f_c$ and $i f_r$ for $i = 1, 2$. We set the fundamental frequencies to $f_c = 0.3$ Hz for the respiratory component (71) and $f_r = 1.1$ Hz for the cardiac component (72).

The phases of each harmonic $\phi$ were randomly selected from a uniform distribution between 0 and $2\pi$ radians. To simulate physiological noise that is proportional to the change in BOLD signal, a variable ratio between the physiological ($\sigma_p$) and the thermal ($\sigma_0$) noise was modeled as $\sigma_p/\sigma_0 = a(tSNR)^b + c$, where $a = 5.01 \times 10^{-6}$, $b = 2.81$, and $c = 0.397$, following the experimental measures available in table 3 from (73).

Experimental Data

To compare the performance of the two approaches as well as illustrate their operation, we employ two representative experimental data sets.

Motor task data set: One healthy subject was scanned in a 3T MR scanner (Siemens) under a Basque Center on Cognition, Brain and Language Review Board-approved protocol. T2*-weighted multi-echo fMRI data was acquired with a simultaneous-multislice multi-echo gradient echo-planar imaging sequence, kindly provided by the Center of Magnetic Resonance Research (University of Minnesota, USA) (74–76), with the following parameters: 340 time frames, 52 slices, partial Fourier = 6/8, voxel size = 2.4 × 2.4 × 3 mm$^3$, TR = 1.5 s, TEs = 10.6/28.69/46.78/64.87/82.96 ms, flip angle = 70°, multi-band factor = 4, GRAPPA = 2.

During the fMRI acquisition, the subject performed a motor task consisting of five different movements (left-hand finger-tapping, right-hand finger-tapping, moving the left toes, moving the right toes, and moving the
tongue) that were visually cued through a mirror located on the head coil. These conditions were randomly intermixed every 16 seconds and were only repeated once the entire set of stimuli was presented. Data preprocessing consisted of, first, discarding the first 10 volumes of the functional data to achieve a steady state of magnetization. Then, image realignment to the skull-stripped single-band reference image (SBRef) was computed on the first echo, and the estimated rigid-body spatial transformation was applied to all other echoes (77,78). A brain mask obtained from the SBRef volume was applied to all the echoes, and the different echo time-series were optimally combined (OC) voxelwise by weighting each time-series contribution by its T2* value (79). AFNI (80) was employed for a detrending of up to fourth-order Legendre polynomials, within-brain spatial smoothing (3 mm FWHM) and voxelwise signal normalization to percentage change. Finally, distortion field correction was performed on the OC volume with Topup (81), using the pair of spin-echo EPI images with reversed phase encoding acquired before the ME-EPI acquisition (82).

Resting-state data sets: One healthy subject was scanned in a 3T MR scanner (Siemens) under a Basque Center on Cognition, Brain and Language Review Board-approved protocol. Two runs of T2*-weighted fMRI data were acquired during resting state, each with 10 min duration, with (1) a standard gradient-echo echo-planar imaging sequence (monoband) (TR = 2000 ms, TE = 29 ms, flip angle = 78°, matrix size = 64 × 64, voxel size = 3 × 3 × 3 mm3, 33 axial slices with interleaved acquisition, slice gap = 0.6 mm) and (2) a simultaneous-multislice gradient-echo echo-planar imaging sequence (multiband factor = 3, TR = 800 ms, TE = 29 ms, flip angle = 60°, matrix size = 64 × 64, voxel size = 3 × 3 × 3 mm3, 42 axial slices with interleaved acquisition, no slice gap). Single-band reference images were also collected in both resting-state acquisitions for head motion realignment. Field maps were also obtained to correct for field distortions.

During both acquisitions, participants were instructed to keep their eyes open, fixating a white cross that they saw through a mirror located on the head coil, and not to think about anything specific. The data were preprocessed using AFNI (80). First, volumes corresponding to the initial 10 seconds were removed to allow for a steady-state magnetization. Then, the voxel time-series

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**Fig. 2.** (A) Simulated signal with different SNRs (20, 10, and 3 dB) and ground truth given in signal percentage change (SPC). (B) Canonical HRF models typically used by PFM (solid line) and TA (dashed line) at TR = 0.5 s (blue), TR = 1 s (green), and TR = 2 s (black). Without loss of generality, the waveforms are scaled to unit amplitude for visualization. (C) Representation of shifted HRFs at TR = 2 s that build the design matrix for PFM when the HRF model has been matched to that in TA. The red line corresponds to one of the columns of the HRF matrix.
were despiked to reduce large-amplitude deviations and slice-time corrected. Inhomogeneities caused by magnetic susceptibility were corrected with FUGUE (FSL) using the field map images (77). Next, functional images were realigned to a base volume (monoband: volume with the lowest head motion; multiband: single-band reference image). Finally, a simultaneous nuisance regression step was performed comprising up to sixth-order Legendre polynomials, low-pass filtering with a cutoff frequency of 0.25 Hz (only on multiband data to match the frequency content of the monoband), 6 realignment parameters plus temporal derivatives, 5 principal components of white matter (WM), 5 principal components of lateral ventricle voxels (anatomical CompCor) (83), and 5 principal components of the brain’s edge voxels (84). WM, CSF, and brain’s edge-voxel masks were obtained from Freesurfer tissue and brain segmentations. In addition, scans with potential artifacts were identified and censored when the Euclidean norm of the temporal derivative of the realignment parameters (ENORM) was larger than 0.4 and the proportion of voxels adjusted in the despiking step exceeded 10%.

Selection of the Hemodynamic Response Function

In their original formulations, PFM and TA specify the discrete-time HRF in different ways. For PFM, the continuous-domain specification of the canonical double-gamma HRF (57) is sampled at the TR and then put as shifted impulse responses to build the matrix $H$. In the case of TA, however, the continuous-domain linearized version of the Balloon-Windkessel model is discretized to build the linear differential operator in $D_h$. While the TR only changes the resolution of the HRF shape for PFM, the impact of an equivalent impulse response of the discretized differential operator at different TR is more pronounced. As shown in Figure 2B, longer TR leads to equivalent impulse responses of TA that are shifted in time, provoking a lack of the initial baseline and rise of the response. We refer the reader to Figure S1 to see the differences in the estimation of the activity-inducing and innovation signals in terms of neuronal-related BOLD activity cannot be predicted. Apart from being able to explore individual maps of deconvolved activity (i.e., innovation signals, activity-inducing signals, or hemodynamic signals) at the temporal resolution of the acquisition (or deconvolution), here we calculate the average extreme points of the activity-inducing and innovation maps (given that these examples do not have a sufficient number of scans to perform a clustering step) and illustrate how popular approaches like coactivation patterns (CAPs) (85,86) and innovation-driven coactivation patterns (iCAPs) (18) can be applied on the deconvolved signals to reveal patterns of coordinated brain activity. To achieve this, we calculate the average time-series in a seed of 9 voxels located in the precuneus, supramarginal gyrus, and occipital gyri independently and solve the deconvolution problem to find the activity-inducing and innovation signals in the

Analyses in Experimental fMRI Data

Difference between approaches: To assess the discrepancies between both approaches when applied on experimental fMRI data, we calculate the square root of the sum of squares of the differences (RSSD) between the activity-inducing signals estimated with PFM and TA on the three experimental data sets as

$$\text{RSSD} = \sqrt{\frac{1}{N} \sum_{k=1}^{N} (\hat{s}_{PFM}[k] - \hat{s}_{TA}[k])^2},$$

where $N$ is the number of time-points of the acquisition. The RSSD of the innovation signals $\hat{u}$ was computed equally.

Task fMRI data: In the analysis of the motor task data, we evaluate the performance of PFM and TA in comparison with a conventional general linear model analysis (3dDeconvolve in AFNI) that takes advantage of the information about the duration and onsets of the motor trials. Given the block design of the motor task, we only make this comparison with the block model.

Resting-state fMRI data: We also illustrate the usefulness of deconvolution approaches in the analysis of resting-state data where information about the timings of neuronal-related BOLD activity cannot be predicted.
seeds. We then apply a 95th percentile threshold and average the maps of the time frames that survive the threshold. Finally, we apply the same procedure to the original—i.e., non-deconvolved—signal in the seed and compare the results with the widely used seed correlation approach.

RESULTS

Performance Based on the Regularization Parameter

Figure 3A shows the regularization paths of PFM and TA side by side obtained for the spike model of Eq. (4) for SNR = 3 dB. The solutions for all three SNR conditions are shown in Figures S2 and S3. Starting from the maximum $\lambda$ corresponding to a null estimate and for decreasing values of $\lambda$, LARS computes a new estimate at the value of $\lambda$ that reduces the sparsity promoted by the $\ell_1$-norm and causes a change in the active set of nonzero coefficients of the estimate (i.e., a zero coefficient becomes nonzero or vice versa) as shown in the horizontal axis of the heatmaps. Vertical dashed lines depict the selection of the regularization parameter based on the BIC, and thus, the colored coefficients indicated by these depict the estimated activity-inducing signal $\hat{s}$. Figure 3B illustrates the resulting estimates of the activity-inducing and activity-related hemodynamic signals when basing the selection of $\lambda$ on the BIC for SNR = 3 dB. Given that the regularization paths of both approaches are identical, it can be clearly observed that the BIC-based estimates are identical too for the corresponding $\lambda$. Thus, Figures 3A and B, S2, and S3 demonstrate that, regardless of the simulated SNR condition, the spike model of both deconvolution algorithms produces identical regularization paths when the same HRF and regularization parameters are applied, and hence, identical estimates of the activity-inducing signal $\hat{s}$ and neuronal-related hemodynamic signal $\hat{x}$. Likewise, Figure 3C demonstrates that the regularization paths for the block model defined in Eqs. (6) and (7) also yield virtually identical estimates of the innovation signals for both PFM and TA methods. Again, the BIC-based selection of $\lambda$ is identical for both PFM and TA. As illustrated in Figure 3D, the estimates of the innovation signal $u$ also show no distinguishable differences between the algorithms. Figure 3A–D demonstrates that both PFM and TA yield equivalent regularization paths and estimates of the innovation signal and activity-inducing signal regardless of the simulated SNR condition when applying the same HRF and regularization parameters with the block and spike models.

As for selecting $\lambda$ with the MAD criterion defined in Eq. (8), Figure 3E depicts the estimated activity-inducing and activity-related signals for the simulated low-SNR setting using the spike model, while Figure 3F shows the estimated signals corresponding to the block model. Both plots in Figure 3E and F depict nearly identical results between PFM and TA with both models. Given that the regularization paths of both techniques are identical, minor dissimilarities are owing to the slight differences in the selection of $\lambda$ due to the quantization of the values returned by LARS.

Performance on Experimental Data

Figure 4 depicts the RSSD maps revealing differences between PFM and TA estimates for the spike (Figure 4A and C) and block (Figure 4B and D) models when applied to the three experimental fMRI data sets. The RSSD values are virtually negligible (i.e., depicted in yellow) in most of the within-brain voxels and lower than the amplitude of the estimates of the activity-inducing and innovation signals. Based on the maximum value of the range shown in each image, we observe that the similarity between both approaches is more evident for the spike model (with both selection criteria) and the block model with the BIC selection. However, given the different approaches used for the selection of the regularization parameter $\lambda$ based on the MAD estimate of the noise (i.e., converging so that the residuals of FISTA are equal to the MAD estimate of the noise for TA vs. finding the LARS residual that is closest to the MAD estimate of the noise), 319 higher RSSD values can be observed with the largest differences occurring in gray matter voxels. 320 These areas also correspond to low values of $\lambda$ (see Figure S4) and MAD estimates of the noise 321 (see Figure S5), while the highest values are visible in regions with signal dropouts, ventricles, 322 and white matter. These differences that arise from the differing methods to find the optimal regularization parameter based on the MAD estimate of the noise can be clearly seen in the root sum of squares (RSS) of the estimates of the two methods (Figure S6). These differences are also observable in the activation time-series (ATS) calculated from estimates obtained with the MAD selection as shown in Figure S9. However, the identical regularization paths shown in Figure S7 demonstrate that both methods perform equivalently on experimental data (see estimates of innovation signal obtained with an identical selection of $\lambda$ in Figure S8). Hence, the higher RSSD values originate from the different methods to find the optimal regularization parameter based on the MAD estimate of the noise that yield different solutions as shown by the dashed vertical lines in Figure S7.

Figure 5 depicts the results of the analysis of the Motor data set with the PFM and TA algorithms using the BIC selection of $\lambda$ (see Figure S9 for results with MAD selection), as well as a conventional GLM approach. The ATS (top left), calculated as the sum of squares of all voxel amplitudes (positive vs. negative) for a given moment in time, obtained with PFM and TA show nearly identical patterns. These ATS help to summarize the four-dimensional information available in the results across
the spatial domain and identify instances of significant BOLD activity. The second to sixth rows show the voxel time-series and the corresponding activity-related, activity-inducing, and innovation signals obtained with PFM using the BIC criterion of representative voxels in the regions activated in each of the motor tasks. The TA-estimated time-series are not shown because they were virtually identical. The maps shown on the right correspond to statistical parametric maps obtained with the GLM for each motor condition ($p < 0.001$) as well as the maps of the PFM and TA estimates at the onsets of individual motor events (indicated with arrows in the timecourses). The estimated activity-related, activity-inducing, and innovation signals clearly reveal the activity patterns of each condition in the task, as they exhibit a BOLD response locked to the onset and duration of the conditions. Overall, activity maps of the innovation signal obtained with PFM and TA highly resemble those obtained with a GLM for individual events, with small differences arising from the distinct specificity of the GLM and deconvolution analyses. Notice that the differences observed with the different approaches to select $\lambda$ based on the MAD estimate shown in Figure 4 are reflected on the ATS shown in Figure S9 as well.
Fig. 4. Square root of the sum of squared differences (RSSD) between the estimates obtained with PFM and TA for (A) spike model (activity-inducing signal) and BIC selection of $\lambda$, (B) block model (innovation signal) and BIC selection, (C) spike model (activity-inducing signal) and MAD selection, (D) block model (innovation signal) and MAD selection. RSSD maps are shown for the three experimental fMRI data sets: the motor task (Motor), the monoband resting state (Mono), and the multiband resting state (Multi).

As an illustration of the insights that deconvolution methods can provide in the analysis of resting-state data, Figure 6 depicts the average activity-inducing and innovation maps of common resting-state networks obtained from thresholding and averaging the activity-inducing and innovation signals, respectively, estimated from the resting-state multiband data using PFM with a selection of $\lambda$ based on the BIC. The average activity-inducing maps obtained via deconvolution show spatial patterns of the default mode network (DMN), dorsal attention network (DAN), and visual network (VIS) that highly resemble the maps obtained with conventional seed correlation analysis using Pearson's correlation and the average maps of extreme points of the signal (i.e., with no deconvolution). With deconvolution, the average activity-inducing maps seem to depict more accurate spatial delineation (i.e., less smoothness) than those obtained from the original data, while maintaining the structure of the networks. The BIC-informed selection of $\lambda$ yields spatial patterns of average activity-inducing and innovation maps that are more sparse than those obtained with a selection of $\lambda$ based on the MAD estimate (see Figure S10). Furthermore, the spatial patterns of the average innovation maps based on the innovation signals using the block model yield complementary information to those obtained with the activity-inducing signal since iCAPs allow to reveal regions with synchronous innovations, i.e., with the same upregulating and downregulating events. For instance, it is interesting to observe that the structure of the visual network nearly disappears in its corresponding average innovation maps, suggesting the existence of different temporal neuronal patterns across voxels in the primary and secondary visual cortices.

DISCUSSION AND CONCLUSION

Hemodynamic deconvolution can be formulated using a synthesis- and analysis-based approach as proposed by PFM and TA, respectively. This work demonstrates that the theoretical equivalence of both approaches is confirmed in practice given virtually identical results when the same HRF model and equivalent regularization parameters are employed. Hence, we argue that previously observed differences in performance can be explained by specific settings, such as the HRF model and selection of the regularization parameter (as shown in Figures 4, S6, and S7), convergence thresholds, as well as the addition of a spatial regularization term in the spatiotemporal TA formulation (33). For instance, the use of PFM with the spike model in (23) was seen not to be ideal due to
the prolonged trials in the paradigm, which better fit the block model as described here (7). However, given the equivalence of the temporal deconvolution, incorporating extra spatial or temporal regularization terms in the optimization problem should not modify this equivalence provided convex operators are employed. For a convex optimization problem, with a unique global solution, iterative shrinkage-thresholding procedures alternating between the different regularization terms guarantee convergence, e.g., the generalized forward-backward splitting (87) algorithm originally employed for TA. Our findings are also in line with the equivalence of analysis and synthesis methods in underdetermined cases (N ≤ V) demonstrated in (50) and (51). Still, we have shown that a slight difference in the selection of the regularization parameter can lead to small differences in the estimated signals when employing the block model with the MAD selection of λ. However, since their regularization paths are equivalent, the algorithms can easily be forced to converge to the same selection of λ, thus resulting in identical estimated signals.

Nevertheless, the different formulations of analysis and synthesis deconvolution models bring along different kinds of flexibility. One notable advantage of PFM is
that it can readily incorporate any HRF as part of the synthesis operator (50), only requiring the sampled HRF at the desired temporal resolution, which is typically equal to the TR of the acquisition. Conversely, TA relies upon the specification of the discrete differential operator that inverts the HRF, which needs to be derived either by the inverse solution of the sampled HRF impulse response, or by discretizing a continuous-domain differential operator motivated by a biophysical model. The more versatile structure of PFM allows for instance an elegant extension of the algorithm to multi-echo fMRI data (31) where multiple measurements relate to a common underlying signal. Therefore, the one-to-many synthesis scenario (i.e., from activity-inducing to several activity-related signals) is more cumbersome to express using TA; i.e., a set of differential operators should be defined and the differences between their outputs constrained. Conversely, the one-to-many analysis scenario (i.e., from the measurements to several regularizing signals) is more convenient to be expressed by TA, e.g., combining spike and block regularizers. While the specification of the differential operator in TA only indirectly controls the HRF, the use of the derivative operator to enforce the block model, instead of the integrator in PFM, impacts positively the stability and rate of the convergence of the optimization algorithms. Moreover, analysis formulations can be more suitable for online applications that are still to be explored in fMRI data but are employed for calcium imaging deconvolution (88,89) and that have been applied for offline calcium deconvolution (90).

Deconvolution techniques can be used before more downstream analysis of brain activity in terms of functional network organization as they estimate interactions between voxels or brain regions that occur at the activity-inducing level and are thus less affected by the slowness of the hemodynamic response compared to when the BOLD signals are analyzed directly. In addition, deconvolution approaches hold a close parallelism to recent methodologies aiming to understand the dynamics of neuronal activations and interactions at short temporal resolution and that focus on extreme events of the fMRI signal (91). As an illustration, Figure 6 shows that the innovation- or activity-inducing CAPs computed from deconvolved events in a single resting-state fMRI data set closely resemble the conventional CAPs computed directly from extreme events of the fMRI signal (85,86,92–99). Similarly, we hypothesize that these extreme events will also show a close resemblance to intrinsic ignition events (100,101).

As shown in the maps, deconvolution approaches can offer a more straightforward interpretability of the activation events and resulting functional connectivity patterns. Here, CAPs were computed as the average of spatial maps corresponding to the events of a single data set. Beyond simple averaging, clustering algorithms (e.g., K-means and consensus clustering) can be employed to discern multiple CAPs or iCAPs at the whole-brain level.

Fig. 6. Average activity-inducing (left) and innovation (right) maps obtained from PFM-estimated activity-inducing and innovation signals, respectively, using a BIC-based selection of \( \lambda \). Time-points selected with a 95th percentile threshold (horizontal lines) are shown over the average time-series (blue) in the seed region (white cross) and the deconvolved signals, i.e., activity-inducing (left) and innovation (right) signals (orange). Average maps of extreme points and seed correlation maps are illustrated in the center.
for a large number of subjects. Previous findings based on iCAPs have for instance revealed organizational principles of brain function during rest (18) and sleep (102) in healthy controls, next to alterations in 22q11ds (103) and multiple sclerosis (104). Next to CAPs-inspired approaches, dynamic functional connectivity has recently been investigated with the use of cofluctuations and edge-centric techniques (105–109). The ATS shown in Figure 5 aim to provide equivalent information to the root of sum of squares timecourses used in edge-centric approaches, where timecourses with peaks delineate instances of significant brain activity. Future work could address which type of information is redundant or distinct across these frameworks. In summary, these examples illustrate that deconvolution techniques can be employed prior to other computational approaches and could serve as an effective way of denoising the fMRI data. We foresee an increase in the number of studies that take advantage of the potential benefits of using deconvolution methods prior to functional connectivity analyses.

In sum, hemodynamic deconvolution approaches using sparsity-driven regularization are valuable tools to complete the fMRI processing pipeline. Although the two approaches examined in detail here provide alternative representations of the BOLD signals in terms of innovation and activity-inducing signals, their current implementations have certain limitations, calling for further developments or more elaborate models, where some of them have been initially addressed in the literature. One relevant focus is to account for the variability in HRF that can be observed in different regions of the brain. First, variability in the temporal characteristics of the HRF can arise from differences in stimulus intensity and patterns, as well as with short interevent intervals like in fast cognitive processes or experimental designs (110–113). Similarly, the HRF shape at rest might differ from the canonical HRF commonly used for task-based fMRI data analysis. A wide variety of HRF patterns could be elicited across the whole brain and possible detected with sufficiently large signal-to-noise ratio; e.g., (114) showed two gamma-shaped responses at the onset and the end of the evoked trial, respectively. This unique HRF shape would be deconvolved as two separate events with the conventional deconvolution techniques. The impact of HRF variability could be reduced using structured regularization terms along with multiple basis functions (29) or procedures that estimate the HRF shape in an adaptive fashion in both analysis (115) and synthesis formulations (116).

Another avenue of research consists in leveraging spatial information by adopting multivariate deconvolution approaches that operate at the whole-brain level, instead of working voxelwise and beyond regional regularization terms (e.g., as proposed in 33). Operating at the whole-brain level would open the way for methods that consider shared neuronal activity using mixed norm regularization terms (117) or can capture long-range neuronal cofluctuations using low-rank decompositions (118). For example, multivariate deconvolution approaches could yield better localized activity patterns while reducing the effect of global fluctuations such as respiratory artifacts, which cannot be modeled at the voxel level (119).

Similar to solving other inverse problems by means of regularized estimators, the selection of the regularization parameter is critical to correctly estimate the neuronal-related signal. Hence, methods that take advantage of a more robust selection of the regularization parameter could considerably yield more reliable estimates of the neuronal-related signal. For instance, the stability selection (65,120) procedure could be included to the deconvolution problem to ensure that the estimated coefficients are obtained with high probability. Furthermore, an important issue of regularized estimation is that the estimates are biased with respect to the true value. In that sense, the use of nonconvex $\ell_p$-norm regularization terms (e.g., $p < 1$) can reduce this bias while maintaining the sparsity constraint, at the cost of potentially converging to a local minima of the regularized estimation problem. In practice, these approaches could avoid the optional debiasing step that overcomes the shrinkage of the estimates and obtain a more accurate and less biased fit of the fMRI signal (30,31). Finally, cutting-edge developments on physics-informed deep learning techniques for inverse problems (118,121–123) could be transferred for deconvolution by considering the biophysical model of the hemodynamic system and could potentially offer algorithms with reduced computational time and more flexibility.

CODE AND DATA AVAILABILITY

The code and materials used in this work can be found in the following GitHub repository: https://github.com/eurunuela/pfm_vs_ta. We encourage the reader to explore the parameters (e.g., SNR, varying HRF options and mismatch between algorithms, TR, number of events, onsets, and durations) in the provided Jupyter notebooks. Likewise, the data used to produce the figures can be found in https://osf.io/f3ryg/.

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REFERENCES


SUPPLEMENTARY MATERIAL FOR HEMODYNAMIC 850 DECONVOLUTION DEMYSTIFIED: SPARSITY-DRIVEN REGULARIZATION AT WORK

Fig. S1. Activity-inducing (A) and innovation (B) signals estimated with PFM (red) and TA (blue) using their built-in HRF as opposed to using the same. The black line depicts the simulated signal, while the green lines indicate the onsets of the simulated neuronal events. X-axis shows time in TRs.
Fig. S2. Spike model simulations. (Left) Heatmap of the regularization paths of the activity-inducing signal estimated with PFM and TA as a function of $\lambda$ (increasing number of iterations in x-axis), whereas each row in the y-axis shows one time-point. Vertical lines denote iterations corresponding to the Akaike and Bayesian Information Criteria (AIC and BIC) optima. (Right) Estimated activity-inducing (blue) and activity-related (green) signals when set based on BIC. All estimates are identical, regardless of SNR.
Fig. S3. Block model simulations. (Left) Heatmap of the regularization paths of the innovation signal estimated with PFM and TA as a function of $\lambda$ (increasing number of iterations in x-axis), whereas each row in the y-axis illustrates one time-point. Vertical lines denote iterations corresponding to the Akaike and Bayesian Information Criteria (AIC and BIC) optima. (Right) Estimated innovation (blue) and activity-related (green) signals when is set based on BIC. All the estimates are identical when compared between the PFM and TA cases, regardless of SNR.
Fig. S4. Values of $\lambda$ across the different voxels in the brain used to estimate (A) the activity-inducing signal (spike model) and (B) the innovation signal (block model) with the BIC selection, as well as (C) the activity-inducing signal (block model) and (D) the innovation signal (block model) with a MAD-based selection. The $\lambda$ maps are shown for the three experimental fMRI data sets: the motor task (Motor), the monoband resting state (Mono), and the multiband resting state (Multi).

Fig. S5. Values of the MAD estimate of standard deviation of the noise across the different voxels in the brain for the three experimental fMRI data sets: the motor task (Motor), the monoband resting state (Mono), and the multiband resting state (Multi).
Fig. S6. Root sum of squares (RSS) comparison between paradigm free mapping and total activation for the three experimental fMRI data sets: the motor task (Motor), the monoband resting state (Mono), and the multiband resting state (Multi). RSS maps are shown for the spike (left) and block (right) models solved with a selection of $\lambda$ based on the BIC (top) and MAD (bottom) criteria.
Fig. S7. Regularization paths of the innovation signal estimated with PFM and TA as a function of \( \lambda \) (increasing number of iterations in x-axis, whereas each row in the y-axis shows one time-point) for the representative voxels of the motor task shown in Figure 5. Vertical lines denote selections of \( \lambda \) corresponding to the BIC (black), MAD based on LARS residuals (blue), and MAD based on FISTA residuals (green) optima.
Fig. S8. Estimated innovation signal (blue) and activity-related signal (green) for the representative voxels of the motor task shown in Figure 5 with the MAD selection of $\lambda$ made by TA, i.e., employing the same $\lambda$ with both PFM and TA.
Fig. S9. Activity maps of the motor task using a selection of $\lambda$ based on the MAD estimate. Row 1: Activation time-series of the innovation signals estimated by PFM (in blue) or TA (in red) calculated as the sum of squares of all voxels at every time-point. Positive-valued and negative-valued contributions were separated into two distinct timecourses. Color bands indicate the onset and duration of each condition in the task (green: tongue, purple: left-hand finger-tapping, blue: right-hand finger-tapping, red: left-foot toes, orange: right-foot toes). Rows 2–6: Time-series of a representative voxel for each task with the PFM-estimated innovation (blue), PFM-estimated activity-inducing (green), and activity-related (i.e., fitted, orange) signals, with their corresponding GLM, PFM, and TA maps on the right. The maps shown on the right are sampled at the time-point labeled with the red arrows and display the innovation signals at that moment across the whole brain.
Fig. S10. Activity-inducing CAPs (left) and innovation CAPs (right) obtained with the PFM-estimated activity-inducing and innovation signals respectively, using a MAD-based selection of $\lambda$. Time-points selected with a 95th percentile threshold are shown over the average time-series (blue) in the seed region (white cross) and the deconvolved signal (orange). CAPs and seed correlation maps are illustrated in the center.