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João Loula, Gaël Varoquaux, Bertrand Thirion. Decoding fMRI activity in the time domain improves classification performance. NeuroImage, 2017, 10.1016/j.neuroimage.2017.08.018 . hal-01576641

## HAL Id: hal-01576641 https://inria.hal.science/hal-01576641

Submitted on 25 Aug 2017  $\,$ 

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# Decoding fMRI activity in the time domain improves classification performance

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#### Abstract

Most current functional Magnetic Resonance Imaging (fMRI) decoding analyses rely on statistical summaries of the data resulting from a deconvolution approach: each stimulation event is associated with a brain response. This standard approach leads to simple learning procedures, yet it is ill-suited for decoding events with short inter-stimulus intervals. In order to overcome this issue, we propose a novel framework that separates the spatial and temporal components of the prediction by decoding the fMRI time-series continuously, i.e. scan-by-scan. The stimulation events can then be identified through a deconvolution of the reconstructed time series. We show that this model performs as well as or better than standard approaches across several datasets, most notably in regimes with small inter-stimuli intervals (3 to 5s), while also offering predictions that are highly interpretable in the time domain. This opens the way toward analyzing datasets not normally thought of as suitable for decoding and makes it possible to run decoding on studies with reduced scan time.

*Keywords:* Functional magnetic resonance imaging, Classification analysis, MVPA, Decoding, Rapid event-related design

#### 1. Introduction

The application of multivariate analysis techniques to fMRI datasets, aka *decoding*, has become a popular approach to probe the relationships between stimuli and brain activity [20, 24, 14]. The very nature of fMRI data makes it a challenging problem: relatively few samples (events or blocks corresponding to stimulus presentation) are avail-

- able, in comparison with the high dimensionality -number of voxels- of each observation. This mismatch leads to the so-called *curse of dimensionality*: learning distributed patterns from few samples is a hard problem. The power of high-dimensional regression methods is thus needed to achieve high
- <sup>15</sup> accuracy and return an interpretable discriminative pattern (see e.g. [5]). However, the sluggishness of the Blood-Oxygen-Level-dependent (BOLD) response observed in fMRI implies that the occurrence of brain activity is not synchronous with the

presentation of stimuli, but delayed by approximately 6s and smooth in time [9]. For the sake of statistical analysis, a preliminary regression step is thus typically performed, so that pairs of stimulus events and associated brain response can be considered. This prior regression is simply carried out by the traditional General Linear Model (GLM) used in standard statistical analyses of fMRI [8].

Although it is the standard solution used by nearly all practitioners, this two-step approach is not optimal; in particular, the intermediate eventrelated brain response estimates are very noisy, limiting decoding accuracy. The reason is that, unlike traditional brain mapping settings in which all events from one condition end up being one single regressor, for decoding purpose, events are split into different regressors, resulting in a loss in design efficiency and high-variance estimates. This approach is also bound to perform poorly on event-related tasks using small inter-stimuli inter-

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- <sup>40</sup> vals (ISIs): the overlap in the hemodynamic response functions (HRFs) coupled with the acquisition noise lead to ill-estimated event-related brain responses and harm subsequent classification accuracy. In this work, we investigate a novel inference
- 45 scheme that swaps the two steps: we propose to perform the challenging and expensive estimation of the discriminative pattern in the time frame of the slow BOLD response. For this we substitute the standard classification problem with one where the
- <sup>50</sup> prediction target is a model of BOLD activity that includes the hemodynamic delay and blurring: we call this approach *time-domain fMRI decoding*. By nature, this approach uses the fMRI data to build a predictor of the convolved stimulus function; then
- a second step is necessary to go from conditionspecific time courses to the identification of events. Given the estimated time course of several conditions during an acquisition, we use a second predictive model to decide which stimulus was presented
- <sup>60</sup> at a given time. The key point is that this learning problem is *easy*, as it boils down to selecting a deconvolution filter and applying a relatively simple selection mechanism.
  - The expected benefit of this approach is to remain tractable whenever the ISI is short (3s to 5s).
- The promise of decoding with shorter ISIs is to optimize scanning efficiency: it allows for either reducing scan duration, leading to cheaper acquisitions and less taxing on subject's attention, or alternatively for keeping scan durations unchanged and
- acquiring more data per experiment.

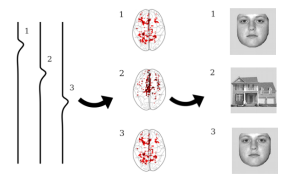


Figure 1: Schema of the GLM approach to decoding. A design matrix representing ideal task responses is used to derive event-specific activation estimates. These estimates are then used in a classification setting.

In the sequel, we describe the so-called *time-domain decoding* framework, and present experiments to compare it with state-of-the-art alternatives: the standard GLM-based regression, the so-called separate GLM approach and spatio-temporal analysis schemes. The fMRI datasets used for validation were chosen so as to represent a wide range of experimental settings, with block and event-related designs, the latter with ISIs ranging from 1.6 to 11.5s. We illustrate accuracy gains in these different settings. Before describing in detail the time-domain decoding approach and our experiments, we review state-of-the-art solutions.

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#### 2. Prior Work

Most decoding studies today are done fitting a first-level GLM regression: a design matrix  $\mathbf{X}$  is created having as columns the timing of the experimental events, convolved with a canonical HRF model, and possibly additional columns to capture nuisance effects. Such an approach is illustrated in Figure 1. A crucial fact is that events corresponding to the same condition are disseminated into different columns, leading to poor (high-variance) pertrial activation estimates.

The activation coefficients are then estimated by solving the  $\mathbf{X}\boldsymbol{\beta} = \mathbf{Y}$  regression problem, where  $\mathbf{Y}$ are the BOLD data, written as a (time, voxel) array. The resulting least-squares estimates  $\hat{\boldsymbol{\beta}}$  for activation coefficients have one value per voxel, hence 100 they make up brain images, one image per event. Data classification is then performed by fitting a classifier to these activation maps: each activation image  $\hat{\boldsymbol{\beta}}_i$  is associated with a label  $l_i$ , that indexes the cognitive condition corresponding to this event. 105 Multivariate inference typically proceeds by estimating a function that predicts l given  $\hat{\beta}$ : this function is a classifier (support vector machine, or logistic regression model) when the labels  $(l_i)_{i=1..I}$  are discrete, or a regression function when the  $(l_i)_{i=1..I}$ 110 are continuous.

To summarize, this approach entails three estimation challenges:

- The one-event-per-column design is statistically inefficient [22].
- Curse of dimensionality: decoding is performed on brain-wide maps estimated based on a limited number of samples.

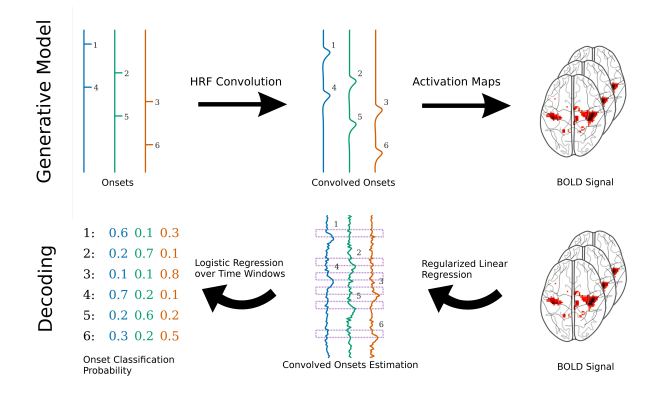


Figure 2: Schema of the time-domain decoding model. Straight arrows represent generative steps, while curved ones represent estimation steps.

• When trying to decode events with small ISIs, the regressors  $(\mathbf{X}_i)_{i=1..I}$  used in the initial regression become highly correlated, thus rendering the first-level estimation ill-posed.

The so-called separate GLM or GLMs approach [19] has been proposed as a means of tackling the first and third issues: it is analogous to the GLM, only a 125 separate design matrix  $\mathbf{X}^{(i)}$  is built for deconvolving each trial i. These separate design matrices only have two columns (besides confounds): one for the stimulus regressor and one for the sum of all other regressors. The activation map for event iis obtained by solving  $\mathbf{X}^{(i)}\boldsymbol{\beta}_i = \mathbf{Y}$ , and the classification is then done through logistic regression on

One thing that the GLM and GLMs approaches have in common is that they proceed by isolat-135 ing temporal features in order to create activation maps. This is typically done by assuming a standard or canonical HRF model. Yet, extensions to data-driven approaches have been proposed in that

 $(\boldsymbol{\beta}_i, l_i)_{i=1..I}$  as in the usual setting.

framework [23][21], using finite impulse response fil-140 ters. It should however be noted that these datadriven models require lots of stimulus occurrences, as they need to estimate regression coefficients in each voxel. Again, this types of model err on the large-variance side, given that a great number of coefficients are estimated per voxel.

Instead of isolating temporal features and decoding over activation maps, a different approach is to perform classification in the time domain. To capture the information from fMRI time series di-150 rectly, the so-called Spatiotemporal SVM approach has been proposed [18], in which, for each event i, a vector  $\mathbf{Y}_i$  is created by concatenating BOLD scans in a time-window following the stimulus onset. Classification is then performed by fitting a 155 linear SVM over these concatenated vectors. While this approach nicely bypasses the prior specification of an HRF, it makes the problem worse regarding the "curse of dimensionality" (second issue outlined above): the number of features in the BOLD sig-160

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nal is multiplied by the length of the time window, thus rendering the classifier fitting problem even more ill-posed.

#### 3. Time-domain decoding: a two-step approach

#### 3.1. Motivation

One way of overcoming the dimensionality and efficiency issues with decoding procedures is to swap the spatial and the temporal estimation prob-

170 lems, by applying the time-lagged analysis in a lowdimensional space. This is the basis for the method proposed here.

#### 3.2. Model

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The Time-domain decoding method (3.2) first recovers the class-specific BOLD time courses to then assigns a class label to events. It comprises two steps:

- 1. Regression of the class-by-class convolved events time-series;
- Classification of the stimulus occurrences based on time windows extracted from the regressed time-series.

More formally, let  $\mathbf{X}$  be the design matrix of size (number of scans  $\times$  number of stimulus classes), containing in each column the time-series for each stimulus class already convolved by an HRF model. Importantly, all the events of any given class are gathered into a single regressor. Let us note the BOLD signal matrix by  $\mathbf{Y}$ . It is assumed that highpass filtering and motion parameter regression have been performed as a preprocessing step on Y. The data can be divided (across sessions) into train and test subsets  $\mathbf{Y}_{train}$  and  $\mathbf{Y}_{test}$ , the corresponding design matrices being  $\mathbf{X}_{train}$  and  $\mathbf{X}_{test}$ ; the spatial step consists in solving a regularized regression problem (written here with a Ridge penalty):

$$\hat{\mathbf{B}} = \underset{\mathbf{B}}{\operatorname{argmin}} \left\{ \left\| \mathbf{X}_{train} - \mathbf{Y}_{train} \mathbf{B} \right\|^2 + \lambda \left\| \mathbf{B} \right\|^2 \right\},\$$
$$\widetilde{\mathbf{X}}_{test} = \mathbf{Y}_{test} \hat{\mathbf{B}},$$
(1)

where  $\lambda$  is a positive scalar to be specified; in the experiments described in this paper, it is set by nested cross-validation (for more details, see Annex

A). Note that **B** is a  $(n_{voxels} \times n_{conditions})$  matrix similar to a standard parameter matrix.

Once an estimate  $\mathbf{X}_{test}$  is obtained, the temporal step classifies each onset using a time window of tscans. Thus, if we denote by  $\widetilde{\mathbf{x}}_{test[i:i+t]}$  the vector formed by concatenating the rows from i to i + tof  $\widetilde{\mathbf{X}}_{test}$ , the temporal step consists in determining, for each onset time i, its corresponding class label  $l_i$ . This is done by multiclass logistic regression:

$$\boldsymbol{l}_{i} = \operatorname*{argmax}_{\boldsymbol{c}_{j}, j \in [\![1, n_{classes}]\!]} \left\{ \operatorname{logit}(\langle \hat{\boldsymbol{w}}_{c_{j}}, \widetilde{\boldsymbol{x}}_{test[i:i+t]} \rangle + \hat{\boldsymbol{b}}_{c_{j}}) \right\},$$
(2)

where  $\hat{\boldsymbol{w}}_{c_j}$  and  $\hat{\boldsymbol{b}}_{c_j}$  are computed for each class  $c_j$  <sup>195</sup> by maximum likelihood estimation. Note that the number of weight coefficients estimated in this step is only t times the number  $n_{classes}$  of classes.

This can be seen in the following manner: the first step (eq. 1) handles the decoding problem as a regression task, in which the strength of the hemodynamic response to each class is fitted by a regularized linear regression model over the BOLD signal on a scan-by-scan basis. Note that this first step relies on a fixed HRF model. In practice we chose the canonical double-gamma function of SPM.

The second, temporal step (see eq. 2 and figure 3) extracts from all classes' predicted time-series a time-window for each onset, and feeds them to a logistic regression model that works as a deconvolution filter for classifying the onset. The time-window length is defined heuristically using the canonical time dynamics of the HRF.

#### 4. Experiments

#### 4.1. Models

We compared the following classification methods: GLM, GLMs, Spatiotemporal SVM and Timedomain decoding described in Sections 2 and 3. Comparisons are presented on 4 datasets. We provide also simulations in appendix (section 7) that reproduce the results on fMRI datasets.

For the time-based models (Spatiotemporal SVM, Time-domain decoding), the time windows were chosen as the closest possible interval to the 2-8s range for event-related design datasets to ensure a good fit of the peak of the canonical HRF, and as the length of the block for datasets with a block design; we present in the annex some data to

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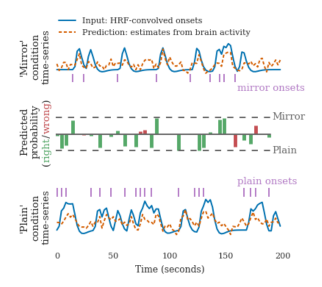


Figure 3: Illustration of time-domain decoding on real data using the *Mirror-reversed text* dataset (see Section 4.3): starting from the BOLD signal, the time-series are estimated for the two classes ('Plain' and 'Mirror') that are the two main conditions. Next, at each stimulus onset, a logistic regression is applied to a time-window in order to classify it. The class probabilities for each onset are shown in the middle graph, with a baseline of 50% at chance; the direction shows which class is predicted as being more likely; the length of the bars encodes the decision confidence and their color represents the correctness of that prediction.

discuss the choice of time-window in model perfor-<sup>230</sup> mance in a post-hoc experiment (Fig. 11).

#### 4.2. Simulation study

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We performed a simulation study to assess the impact of ISI on model performance in a controlled setting. We generated data using a model of the form  $\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon}$ :

• The design matrix Χ, of shape (number of scans  $\times$  2), was created by convolving stimuli randomly assigned to one of two classes, separated by the ISI chosen for the session. As in the real data studies, we use the HRF model of [9]. We chose a number of scans of 1000 for the ISI=5s condition (as an approximation for the concatenation of the number of scans across all runs in a real experiment), and then adjusted for the other classes so as to have balanced number

of stimuli for each ISI length (thus yielding 800 scans for ISI=4s and 600 for ISI=3s). We can note that, given a TR of 2s, this would correspond to 33 minutes of scanning time for the 5s ISI, 26 minutes for the 4s ISI and 20 minutes for the 3s one (not counting resting intervals).

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- For the two classes  $\beta$ , we created the activation maps, a matrix of shape  $(2 \times 10000)$ , by drawing their coefficients from a multivariate normal distribution  $\mathcal{N}(3, \sigma_{\beta}^2 I)$ , where  $\sigma_{\beta}$  was set to .5. We chose both the mean and the standard deviation as in the simulation study in [19]. The number 10000 for the features was chosen so as to reflect the number of features chosen by ANOVA variable selection in the real data studies.
- We generated the noise  $\epsilon$ , a matrix of shape (number of scans × 10000), from an i.i.d. normal distribution  $\mathcal{N}(0, \sigma_{\epsilon}^2)$ , where  $\sigma_{\epsilon}$  was set to 1.6, again as in [19]. We then smoothed the noise both temporally and spatially using a Gaussian filter with unit standard deviation, which led to an average standard deviation of approximately 0.72 for  $\epsilon$  across simulations.

We set the TR to 2 seconds and test ISIs of 3, 4, and 5 seconds. Results for activation map correlations of 0, 0.3 and 0.6 are also shown in 7. For each ISI and correlation, we run 100 simulations, in <sup>275</sup> which we generate both a train and a test set with the procedure described above.

#### 4.3. Real data

In order to probe different timing intervals and decoding complexity levels, we considered 4 280 datasets:

• The Haxby dataset [13] yields a study of face and object representations in human ventral temporal cortex, with 6 subjects and 12 runs per subject. Stimuli consisted of greyscale images from eight different classes: faces, cats, houses, chairs, scissors, shoes, bottles and scrambled images, and we considered the 8class classification problem. Images for each class were shown during 24s, followed by 12s 290 of rest; TR=2.5s, ISI=36s; • The *Mirror-reversed text* [15] dataset yields a study of the neural basis of task-switching, with 14 subjects and 6 runs per subject. Stimuli were words shown in either plain or mirrorreversed fashion, coupled with a semantic classification task. The design is event-related, with TR=2s and ISI=4-11.5s.

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- The *Textures* dataset [6] yields a study of responses to textures along different regions of 300 interest in the brain, with 4 subjects undergoing a total of 7 acquisitions (3 subjects having gone through two acquisitions), with 6 runs per acquisition. Stimuli were greyscale texture images from 6 different classes in the 305 UIUC texture database [17], appearing during three flashes of 200ms, separated by 200ms grey screen, so that an event duration is 1s. These images were shown in pairs separated by 4s followed by a probe after which the sub-310 ject had to decide which of the first two the third image shown was extracted from. The design is event-related design, with TR=2.4s and ISI=4-8s.
- The Temporal tuning dataset [10] yields a study of rate-dependence of neural responses, with 11 subjects and 12 runs per subject. Greyscale images of faces and houses were shown in alternating fashion during 20s blocks, followed by 10s of rest. The design is event-related, with TR=1.5s and ISI=1.6, 3.2 or 4.8s.

Performance was analyzed in a within-subject setting. The cross-validation method used was Leave-one-session-out for *Mirror-reversed text* and *Textures*, and Leave-two-sessions-out for *Haxby* and *Temporal tuning* (based on the heuristic of having approximately 20% of the data in the test set). Cross-validation on the *Temporal tuning* dataset also followed a class-rebalancing scheme described in detail in section 7. We used classification accuracy of the grants as the evaluation metric. One

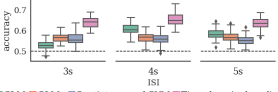
racy of the events as the evaluation metric. One score was computed per cross-validation step, and significance of mean accuracy difference between methods was tested using paired t-tests.

### 335 4.4. Implementation

The analyses were performed in Python using the module Nilearn version 0.2.6, with Scikit-learn version 0.17.1 and Numpy version 1.11.3. Statistical analyses were performed using Nistats version 0.1.0, using the SPM model for the HRF. Plots were created using Matplotlib version 1.5.1 and Seaborn version 0.7.1. An implementation of the analysis for the public *Haxby* dataset can be found at https: //github.com/joaoloula/time\_decoding.

#### 5. Results

Fig. 4 shows the results of the simulation study with no correlation between activation maps. The simulation suggests greater performance of the Time-domain decoding method over alternatives for this controlled environment, particularly for small ISIs.



GLM GLMs Spatiotemporal SVM Time-domain decoding

Figure 4: Simulation study showing the prediction accuracy for varying ISIs. The dotted line represents the chance level (50%).

For the real data, the relative accuracies for all methods on the *Haxby*, *Mirror-reversed text* and *Texture decoding* datasets are presented in Figure 5.

- On the Haxby dataset, Time-domain decoding outperforms all other methods ( $p < 10^{-10}$ , uncorrected), showing that it does well in traditional block designs. We can also see that this is the dataset in which Spatiotemporal SVM has its worst relative performance : th is is most likely an effect of the curse of dimensionality, given that the time-window is largest in this dataset. The (across methods) mean accuracy is 49% and the chance level is 12.5%;
- On the *Mirror-reversed text* dataset, Timedomain decoding outperforms GLM ( $p < 10^{-9}$ , uncorrected), and is outperformed by Spatiotemporal SVM ( $p < 10^{-3}$ , uncorrected).

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- The mean accuracy is 76% and the chance level 370 is 50%.
  - On the *Texture decoding* dataset, Timedomain decoding outperforms GLM (p  $\,<\,$  $10^{-8}$ , uncorrected) and Spatiotemporal SVM  $(p < 10^{-7}, \text{ uncorrected})$  and is outperformed by GLMs (p < 0.05, uncorrected). The mean accuracy is 43% and the chance level is 16.7%.

In Figure 6, we give results on the *Temporal tun*ing dataset: the accuracies for each method, obtained through the cross-validation procedure de-380 scribed in 7, are shown separately according to the test-set ISI, which can be of 1.6, 3.2 or 4.8 seconds.

- When ISI=1.6 seconds, no method performs significantly better than chance;
- When ISI=3.2 seconds, Time-domain decoding 385 outperforms GLMs ( $p < 10^{-6}$ , uncorrected);
  - When ISI=4.8 seconds, Time-domain decoding significantly outperforms GLM (p < 0.05, uncorrected), GLMs ( $p < 10^{-4}$ , uncorrected) and Spatiotemporal SVM ( $p < 10^{-6}$ , uncorrected).

The fact that GLMs is outperformed by GLM for small ISIs in this dataset is most likely a consequence of high inter-trial variability (see [1]). The simulations results presented in section 7 confirm the superiority of the time-domain decoding method for ISIs of 3s to 5s.

Finally, Figure 7 shows the activation maps obtained for all methods on the Face vs. House task 400 on the *Haxby* dataset. We can see that the maps for the four methods are highly similar: this indicates that the prediction problem as posed by the Time-domain decoding method still yields meaningful brain maps.

#### 6. Discussion 405

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Our experiments on the simulated data and four different real datasets establish clearly that the Time-domain decoding method performs as well as or better than state-of-the-art approaches. It does so in spite of the differences between datasets with respect to their timing characteristics. Put differently, this means that this approach is a safe default choice for the sake of decoding performance.

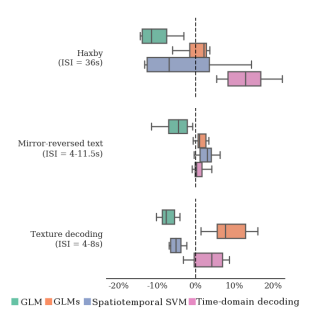


Figure 5: Subject-by-subject accuracy comparison between GLM, GLMs, Spatiotemporal SVM and Time-domain decoding across all datasets. Only the perfold accuracy difference between methods is plotted in these figures: the dotted line represents the per-fold mean performance across methods. The mean accuracies (chance levels) are respectively: 49% (12.5%) on Haxby, 76% (50%) on Mirror-reversed text and 43% (16.7%) on Textures.

The results on the Temporal Tuning dataset are of particular interest: though extremely small ISIs 415 degrade the performance of all methods to chance level, with an ISII of 3.2s, Time-domain decoding outperforms GLMs and Spatiotemporal SVM, and at 4.8s it outperforms all other methods. With respect to the Spatiotemporal SVM, in particular, we 420 confirm that Time-domain decoding does not suffer from the additional ill-posedness inherent to the strategy that augments the dimension of the input space. The direct implication of this result is that decoding becomes usable for ISIs as low as 3-4s, 425 without jeopardizing too much prediction accuracy. This is thus a useful contribution toward cheaper, less demanding experiments for participants and opens the possibility to re-analyze existing datasets that have not been designed for decoding purposes.

More in detail, the first step of the Time-domain decoding uses a pre-defined HRF at learning time (to form the time courses used to train the spatial decoder), while the second step does not rely on a

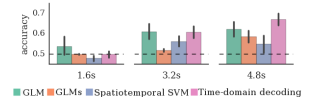


Figure 6: Accuracy comparison between GLM, GLMs, Spatiotemporal SVM and Time-domain decoding across different ISIs on the *Temporal tuning* dataset.

- temporal model. Our experience is that the procedure is robust to the choice of convolution model amongst canonical options e.g. using the canonical SPM response [7]. On the other hand, the deconvolution is a very sensitive step: in particular we
- <sup>440</sup> have considered using model-based deconvolution instead of temporal decoding as in Eq. 2 –actually inverting a canonical generative model of the data– but this systematically lead to poorer predictions. The Time-domain decoding is also of a different na-
- ture than other spatio-temporal methods such as the one presented in [16], that tackle the question of decoding without timing information as opposed to that of separating stimuli in rapid succession. Notably, the *total activation* approach leverages a
  prior on neural events timing (minimization of the

discontinuities) that is not used here.

It is worth noting that the Time-domain decoding method is distinct from HRF estimation approaches, as it does not aim at recovering the actual HRF that couples neural reponses with BOLD signals (an HRF model is used only in the first step to set  $X_{train}$ ): no physiological model is estimated, and instead, the convolution filter is handled in the temporal step as a nuisance factor that simply needs to be inverted. In this respect, the presented approach is a discriminative rather than a generative model. In particular, while the HRF is likely to vary across brain regions [4, 21, 2], the temporal deconvolution performed in this work is

<sup>465</sup> an abstract, location-free filter estimate. Critically, it may not correspond to the local signal model of any brain region, although it can be interpreted as an inverse filter of the average HRF.

Aside from performance, two advantages of the <sup>470</sup> Time-domain decoding model are worth pointing out:

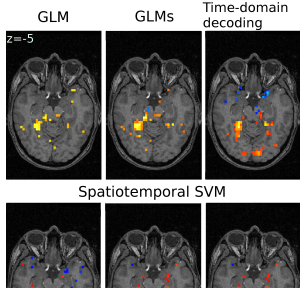


Figure 7: Activation maps for Face vs. House classes on the *Haxby* dataset for subject 2 for all models, shown at a 99.5% percentage threshold of signal strength. Since Spatiotemporal SVM produces a map for each scan in the time window (10 in the case of the *Haxby* dataset), we present only 3 maps corresponding to the timepoints of 2, 4 and 6s. We note the similarity between the maps, namely in the activation of the Fusiform Face Area: this seems to indicate that the regression problem posed by Logistic Regression yields meaningful maps.

- The decoupling of the spatial and temporal steps makes the method modular, and therefore particularly well-adapted to the substitution of richer models at each step. Possibilities include performing the spatial regression step using a spatially-regularized Graph-Net [12] or TV-L1 model [3] [11], or performing the lowdimensional deconvolution with another classifier such as Random Forest.
- The introduction of time-series for each class as an intermediary prediction step provides useful time-domain interpretability: a misclassification can be traced back to the time-series (see for example Figure 3). Notably, brain activations for each class can be tracked throughout scan times, allowing one to observe the ef-

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fects of rivaling class-specific time courses and signal strength on decoding performance.

- Finally, it should be noted that the use of Time-490 domain decoding can enhance the consequence of bad experimental design: if there exists a timedomain dependency between the occurrence of the different classes, it is possible that the model will
- capture those characteristics rather than meaning-495 ful cognitive features. Its use can therefore only be recommended on datasets with properly randomized events.

#### 7. Conclusion

- We presented the Time-domain decoding method 500 for multivariate fMRI data decoding, which allows for efficient decoding with smaller ISIs than the state of the art, and is flexible to HRF variations. By design, it avoids the computational burden associated with time embedding approach used so far
- in the so-called spatio-temporal hrf model. The method is modular in nature, with weakly coupled spatial and temporal steps, and offers interpretability in the time domain. It has been shown
- to perform robustly on four different datasets, and 510 to outperform alternatives in a short-ISI dataset. Code implementing the method as well as examples on the *Haxby* dataset can be found at https: //github.com/Joaoloula/time\_decoding.
- Acknowledgements. We are grateful to Baptiste 515 Gauthier for allowing us to use the Temporal Tuning dataset in the experiments presented here. This project has received funding from the European Union's Horizon 2020 Framework Programme for Research and Innovation under Grant Agreement 520
- No 720270 (Human Brain Project SGA1).

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#### Annex A: Cross-validation scheme for the Temporal tuning dataset

The Temporal tuning dataset contains stimuli
with ISIs of 4.8, 3,2 and 1.6 seconds. The experiment design, however, makes it so that these stimuli are not balanced: trials occur in blocks in which
"Face" and "House" conditions are alternated with
a given rate, making small ISI trials more numerous than larger ISI ones, as seen in Figure 8. To avoid biases introduced by the different number of samples, we subsampled the most frequent categories in order to achieve a balance in the number of examples for each ISI.

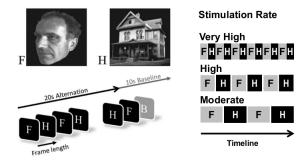


Figure 8: Event structure of the Temporal tuning dataset. Images of faces and houses were shown in alternation during continuous 20s blocks separated by 10s of rest. The ISI for each of these blocks was set to either 4.8, 3.2 or 1.6 seconds. It should be observed that this leads to an overabundance of trials with small ISI with respect to larger ISI ones: this makes rebalancing in the cross-validation necessary (see main text). Adapted from [10].

With these considerations, the cross-validation procedure performed was the following:

- 1. Different subsets of the 1.6s and 3.2s ISI stimuli are created, each containing the same number of examples as the set of 4.8s stimuli (which is the smallest group of the three, with 4 stimuli per block);
- 2. For each of these subsets, 20% of the data are left out in the validation set; the rest constitutes the decoding set. On the decoding set, a nested cross-validation loop is used to define the Ridge regression constant for the Logistic Regression model, as visualized in Figure 9.

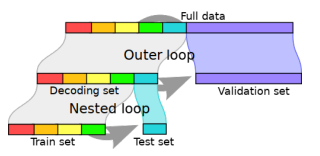


Figure 9: Nested cross-validation procedure for the Logistic Regression model. For each of the balanced subsets of the data, an inner CV loop is used to determine the Ridge hyperparameter while an outer one is used to assess model performance for each ISI.

#### Annex B: Simulation Study

In the context of the simulation study described in section 4.2, we also checked the influence of the correlation between activation maps on model performance. The setting is the same as in section 4.2, but now the distribution from which the activation maps are drawn has a non-trivial correlation structure  $\sigma_{beta}^2(I+Cor)$ , where  $Cor_{i,j} = c$  if the features *i* and *j* correspond to the same voxel across the two maps and  $Cor_{i,j} = 0$  otherwise. The value of *c* was made to vary between 0, 0.3 and 0.6, and for each ISI-correlation pair 100 simulations were run. The results are shown in fig 10.

We observe that, while higher correlation decreases performance across all methods (as expected), it does so in a non-homogeneous way: notably, while tests with low correlation show that Time-domain decoding outperforms all other methods, with a correlation of 0.6 the performance is mostly uniform across the four models. This indicates that Time-domain decoding does not address the issue of ill-posed spatial pattern estimation.

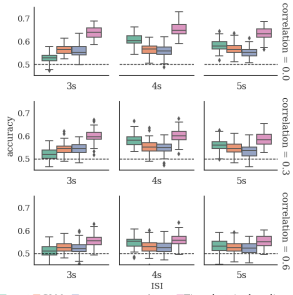
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#### Annex C: Time window length analysis

In order to study the influence of the timewindow length parameter on the performance of the temporal step of the Time-domain decoding method (as described in section 3) we performed tests on the Haxby dataset using 5 different window lengths: 2.5, 5, 7.5, 10 and 20 seconds, the first 4 centered around the canonical peak for the HRF (around 5 to 6 seconds after the stimulus onset),



GLM GLMs Spatiotemporal SVM Time-domain decoding

Figure 10: **Prediction accuracy for varying ISIs and between-map correlations (simulated data)**. The dotted line represents the chance level (50%). Correlations degrade the accuracy obtained, but perserves the relative performance of the methods.

- <sup>715</sup> and the 20s window beginning at the stimulus onset. These are equivalent respectively to one, two, three, four and eight scans: for the case with one scan, we performed classification by simply taking the label to be the one with maximum activation for
- <sup>720</sup> that scan. The performance was measured across all 8 classes on the Haxby dataset, and the crossvalidation procedure used was the same as the one in the previous experiments.
- The accuracy metric across these different window lengths gives an indication of how variations in the time-window size affect decoding performance: as we can see, though performance is high across all windows, there is a high increase in the 7.5s window relative to the smaller ones. In particular, we notice
- that the decoder that only uses the maximum activation at one timestep for classification (the 2.5s window decoder) fails to achieve the performance of the time-domain decoding methods for longer timewindows, attesting the utility of the logistic regres-
- r35 sion as a time-domain deconvolution step. Given the notable ISI on the Haxby dataset, we see steady increase in accuracy with time-window length satu-

rating around 10s, while doubling the length to 20s has almost no impact on performance.

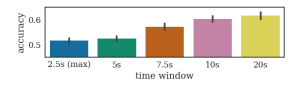


Figure 11: Impact of the time-window length on the performance of the Time-domain decoding method in the Haxby dataset. One can see that the jump from 5s to 7.5s lenght (2 to 3 scans) yields great improvement in performance, and that there is generally steady increase in accuracy with time-window length, saturating at around 10s. Recall that chance is 1/8 = .125.