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CanICA: Model-based extraction of reproducible group-level ICA patterns from fMRI time series

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Abstract. Spatial Independent Component Analysis (ICA) is an increasingly used data-driven method to analyze functional Magnetic Resonance Imaging (fMRI) data. To date, it has been used to extract meaningful patterns without prior information. However, ICA is not robust to mild data variation and remains a parameter-sensitive algorithm. The validity of the extracted patterns is hard to establish, as well as the significance of differences between patterns extracted from different groups of subjects. We start from a generative model of the fMRI group data to introduce a probabilistic ICA pattern-extraction algorithm, called CanICA (Canonical ICA). Thanks to an explicit noise model and canonical correlation analysis, our method is auto-calibrated and identifies the group-reproducible data subspace before performing ICA. We compare our method to state-of-the-art multi-subject fMRI ICA methods and show that the features extracted are more reproducible.

1 Introduction

Resting-state fMRI is expected to give insight into the intrinsic structure of the brain and its networks. In addition, such protocols can be easily applied to impaired subjects and can thus yield useful biomarkers to understand the mechanisms of brain diseases and for diagnosis. Spatial ICA has been the most successful method for identifying meaningful patterns in resting-state fMRI data without prior knowledge. The use of the resulting patterns is widespread in cognitive neuroscience, as they are usually well-contrasted, separate different underlying physiological, physical, and cognitive processes, and bring into light relevant long-range cognitive networks.

However, validation of the resulting individual patterns suffers from the lack of testable hypothesis. As a result, cognitive studies seldom rely on automatic analysis, and relevant ICA maps are cherry picked by eye to separate them from noise-induced patterns. Probabilistic ICA models have been used to provide pattern-level noise-rejection criteria [1] or likelihood for the model [2], but have yet to provide adequate auto-calibration and pattern-significance testing. The lack of reproducibility is detrimental to group analysis: various, often non-overlapping, patterns have been published [3,4] and the statistical frameworks for comparison or inference on ICA maps have to be further developed.

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To allow for group analysis, it is important to extract from a group of subjects ICA maps that are well-represented in the group. Various strategies have been adopted: group ICA \( \text{gICA} \) concatenates individual time series; tensor ICA \( \text{tICA} \) estimates ICA maps across subjects with different loadings per subject; NEDICA \( \text{NEDICA} \) merges ICA maps by hierarchical clustering.

In this paper, we present a novel model and a method, that we dub CanICA, to extract only the reproducible ICA maps from group data. The strength of this method lies in the elimination of the components non-reproducible across subjects using model-informed statistical testing and canonical correlation analysis (CCA). We compare the reproducibility of features extracted by our method to features extracted using tensor ICA and group ICA, but could not compare to NEDICA since it suppresses in some way between-subject variability.

2 Methods

2.1 Generative model: from group-level patterns to observations

At the group level, we describe intrinsic brain activity by a set of spatial patterns \( \mathbf{B} \) corresponding to networks common to the group. We give a generative model to account for inter-subject variability and observation noise.

The activity recorded on each subject \( s \) can be described by a set of subject-specific spatial patterns \( \mathbf{P}_s \), which are a combination of the group-level patterns \( \mathbf{B} \) with loadings given by \( \mathbf{\Lambda}_s \) and additional subject-variability denoted as a residual matrix \( \mathbf{R}_s \). If we write the spatial patterns \( \mathbf{B}, \mathbf{P}_s \) and \( \mathbf{R}_s \) as \( n_{\text{patterns}} \times n_{\text{voxels}} \) matrices, for each subject \( s \), \( \mathbf{P}_s = \mathbf{\Lambda}_s \mathbf{B} + \mathbf{R}_s \). In other words, at the group level, considering the group of patterns (vertically concatenated matrices) \( \mathbf{P} = \{ \mathbf{P}_s \}, \mathbf{R} = \{ \mathbf{R}_s \}, \) and \( \mathbf{\Lambda} = \{ \mathbf{\Lambda}_s \}, s = 1 \ldots S \),

\[
\mathbf{P} = \mathbf{\Lambda} \mathbf{B} + \mathbf{R}.
\]

For paradigm-free acquisitions, there is no specific time course set by an external stimulus or task, and for each acquisition-frame time point a mixture of different processes, described by different patterns, is observed. The observed fMRI data is a mixture of these patterns confounded by observation noise: let \( \mathbf{Y}_s \) be the resulting spatial images in BOLD MRI sequences for subject \( s \) (an \( n_{\text{frames}} \times n_{\text{voxels}} \) matrix), \( \mathbf{E}_s \) the observation noise, and \( \mathbf{W}_s \) a loading matrix such that:

\[
\mathbf{Y}_s = \mathbf{W}_s \mathbf{P}_s + \mathbf{E}_s.
\]

2.2 Estimating group-reproducible patterns from fMRI data

Noise rejection using the generative model. Starting from fMRI image sequences, \( \{ \mathbf{Y}_s, s = 1 \ldots S \} \), time-sliced interpolated and registered to the MNI152 template, we separate reproducible patterns from noise by estimating successively each step of the above hierarchical model.
First, we separate observation noise $E_s$ from subject-specific patterns $P_s$ (Eq. 2) through principal component analysis (PCA). The principal components explaining most of the variance for a given subject’s data set form the patterns of interest, while the tail of the spectrum is considered as observation noise. Using a singular value decomposition (SVD), $Y_s = U_s \Sigma_s V_s^T$. The $n$ first columns of $V_s^T$ constitute the “whitened” patterns $P_s$ that we retain: $\hat{P}_s = (V_s)_{1...n}$, and the residual constitutes the observation noise: $\hat{E}_s = Y_s - (U_s \Sigma_s V_s^T)_{1...n}$.

As PCA can be interpreted as latent-variable separation under the assumption of normally-distributed individual random variates, we model observation noise as normally-distributed. Following [7], we set the number $n$ of significant PCA components retained by drawing a sample null-hypothesis dataset using a random normal matrix and comparing the bootstrap stability of PCA patterns for the measured signal and the null-hypothesis sample. Unlike information-based criteria used in previous methods [1,5] for order selection, such as approximations of model evidence or BIC, the selected number of significant components does not increase when adding artificially number of noise sources [7], and thus does not diverge with long fMRI time series. This is important to avoid extracting group-reproducible patterns from observation noise.

To identify a stable-component subspace across subjects, estimating Eq. (1), we use a generalization of canonical correlation analysis (CCA). CCA is used to identify common subspaces between two different datasets. While there is no unique generalization to multiple datasets, an SVD of the various whitened and concatenated datasets can be used and is equivalent to standard CCA in the two-datasets case [8]. Given $\hat{P} = \{\hat{P}_s\}$, SVD yields $\hat{P} = \hat{Y} Z \Theta^T$ where $\Theta^T$ forms the canonical variables, and $Z$ the canonical correlations, which yield a measure of between-subject reproducibility. Estimation of the inter-subject reproducible components $\hat{B}$ is given by the vectors of $\Theta^T$ for which the corresponding canonical correlation $Z$ is above significance threshold. $\hat{A}$ is identified as the corresponding loading vectors of $\hat{Y} Z$. For a given number of selected components, this estimator minimizes the sum of squares for the residual $\hat{R} = \hat{P} - \hat{A} \hat{B}$.

The significance threshold on the canonical correlations is set by sampling a bootstrap distribution of the maximum canonical correlation using $\hat{E}_s$, the subject observation noise identified previously, instead of $\hat{P}_s$. Selected canonical variables have a probability $p < 0.05$ of being generated by the noise.

Identifying independent features in the group-level patterns. The selected group-level components $B$ form reproducible patterns of task-free activation, but they represent a mixture of various processes and are difficult to interpret for lack of distinguishable shape standing out. We perform source separation using spatial ICA on this subspace. From the patterns $B$ we estimate a mixing matrix $M$ and group-level independent components $A$ using the FASTICA algorithm [9]: $B = MA$. FASTICA is an optimization algorithm which successively extracts patterns with maximally non-Gaussian marginal distributions. This separation corresponds to identifying the maximum-contrast patterns in the subspace of interest. These spatial patterns correspond to minimally-
dependant processes and contain identifiable physiological, physical, or neuronal components.

Consistent with the FASTICA model, the main regions that form the nodes of the functional networks within the resulting patterns are the regions with values corresponding to the non-Gaussian tails of the histogram. Following [10], we model the non-interesting voxels as normally-distributed and estimate the null distribution from the central part of the histogram. The voxels of interest are selected using an uncorrected p-value of $10^{-3}$. We use a specificity criterion rather than a false discovery rate as it yields more stable results, especially on the very-long-tailed distributions that we encounter. As the total number of voxels in the brain is about 40,000, we expect no more than 40 false positives, which corresponds to a small false-discovery rate.

2.3 Model validation for inter-subject generalization

The validation criteria for an ICA decomposition are unclear, as this algorithm is not based on a testable hypothesis. The use of ICA is motivated by the fact that the patterns extracted from the fMRI data display meaningful features in relation to our knowledge of functional neuroanatomy. These features should be comparable between subjects, and thus generalize to new subjects.

To test the reproducibility of the results across subjects, we split our group of subjects in two and learn ICA maps from each sub-group: this yields $A_1$ and $A_2$. We compare the overlap of thresholded maps and reorder one set to match maps by maximum overlap. Reproducibility can be quantified by studying the cross-correlation matrix $C = A_1^T A_2$. For unit-normed components, $C_{i,j}$ is 1 if and only if $(A_1)_i$ and $(A_2)_j$ are identical.

We define two measures for overall stability and reproducibility of the maps. First, a measure of the overlap of the subspaces selected in both groups is given by the energy of the matrix: $E = \text{tr} (C^T C)$. To compare this quantity for different subspace sizes, we normalize it by the minimum dimension of the subspaces, $d = \min(\text{rank } A_1, \text{rank } A_2)$: $e = \frac{1}{2} \text{tr} (C^T C)$. $e$ quantifies the reproducibility of the subspace spanned by the maps. For $e = 1$, the two groups of maps span the same subspace, although individual independent components may differ. Second, we use an overall measure of reproducibility for the maps: the normalized trace of the reordered cross-correlation matrix $C$, $t = \frac{1}{d} \text{tr} (C)$. Indeed, after $A_2$ has been reordered to maximize matching with $A_1$, the diagonal coefficients of $C$ give the overlap between matched components. Finally, the maximum value of each row and column of $C$ expresses the best match each component learned in one group on the set learned in the other. We plot its histogram. This indicator accounts for components of one group matching multiple components of the other.

3 Results

3.1 Extracted group-level patterns of interest

Twelve healthy volunteers were scanned after giving informed consent. 820 EPI volumes were acquired every $TR = 1.5$ s at a 3mm isotropic resolution, during a
CanICA identified 50 non-observation-noise principal components at the subject level (Eq. 2) and a subspace of 42 reproducible patterns at the group level (Eq. 1), which matches numbers commonly hand-selected by users in current ICA studies. On these long sequences, model-evidence-based methods such as those used in [1] select more than 300 components. Extracted maps can be classified by eye in neuronal components, cardio-spinal-fluid (CSF) induced fluctuations, and movement-related patterns (Fig. 1). The empirical-null-based thresholding yields best results for neuronal components (see Fig. 2). An interesting side result of these maps is that measurement artifacts such as movement or CSF noise form reproducible patterns between different subjects.

### 3.2 Inter-group reproducibility

We performed 38 analyses on paired groups of 6 different subjects. Out of the 76 groups, 2 yielded 20 stable components, 19 yielded 21 components, 36 yielded 22 components, 17 yielded 23 components, and the last 2 yielded 24 components. We compare our method with tensor ICA [6], running analysis using the MELODIC software on each group, and group-ICA, using the GIFT ICA toolbox (http://icatb.sourceforge.net/). To avoid bias from the selected-subspace dimension, we run the tensor ICA and group ICA analysis specifying 23 components.

We do cross-correlation analysis on the non-thresholded ICA maps, but also use each implementation’s thresholding algorithm to separate the features of interest. For Group-ICA, the thresholding is done on the $t$ statistics maps generated by the algorithm. As these maps have low amplitude, thresholding on $|t| > 3$ leaves very few selected voxels; we use $|t| > 2$, which yields the same number of selected voxels as those selected by the two other methods.

On non-thresholded maps, CanICA and Group ICA perform similarly whereas tensor ICA selects a slightly less stable subspace, and thus yields less reproducible ICA maps (see Fig. 3 and Tab. 1). Thresholding the maps does not significantly change the subspace stability ($e$) and map reproducibility ($t$) for CanICA, but decreases performance for tensor ICA and drastically affects stability and reproducibility for group ICA.
Fig. 2. (a) A neuronal ICA map and (b) its histogram. The null distribution, shown in red on the histogram, is estimated from the center of the histogram. (c) Corresponding thresholded map; only voxels with $p < 5 \times 10^{-4}$ (uncorrected, two-sided test) are kept.

Fig. 3. (a) Typical cross-correlation matrices between non-thresholded ICA maps learned on two half-splits of the total 12 subjects, for group ICA, tensor ICA, and CanICA. (b) Histogram of the maximal overlap for each component from one half-split to a component from the other half. (c) and (d) cross-correlation matrices and histogram for thresholded ICA maps.
4 Discussion

**Importance of capturing inter-subject variability** The close correspondence between the overlap of the selected subspace \((e)\) and independent-component matching quality \((t)\) suggests that identification of the reproducible signal is a key step to identifying stable independent components. Our method explains only a small fraction of the total signal variance—less than 50% for all 12 subjects. This fraction is selected both on noise-rejection criteria, with well-specified noise models, and to best take in account inter-subject variability, by CCA. Indeed, CCA selects linear combinations of subject patterns that have highest canonical correlations. As the individual subject components are whitened, each one can contribute no more than 1 to the canonical correlation. Thus high canonical correlations ensure representation of a large fraction of the group. In addition, the algorithm minimizes the sum of squares of the total subject-variability residual, \(R\). The stability of the subspace selected by the GIFT implementation of group ICA is explained by similar reasons.

We believe that subject variability is not accounted for as well by tensor ICA because the variability noise is estimated during the tensorial ICA step, for which statistical significance is hard to establish. The combination of subject-specific components present in the final estimated independent components is not guaranteed to reflect multiple-subject contributions and in practice the corresponding subject-loading vectors are often unbalanced across subjects.

**Residual ICA-patterns instability** ICA extracts maps by successive optimizations of linear combinations to maximize negentropy. It is very flexible because, unlike PCA, it does not require orthogonality of the corresponding time courses. However, even with the careful noise reduction performed by CanICA, small signal perturbation can lead to different patterns. As an example, CanICA run on two eleven-subjects groups, differing only by one subject, can yield patterns in which the plausible neuronal activation clusters differ (Fig. 4). One should thus be careful when inferring cognitive networks using ICA and, when possible, do significance testing using other criteria, such as seed-voxel correlation analysis.

![Fig. 4](image-url). Two ICA patterns corresponding to Fig. 2 estimated from two subgroups of 11 subjects, with 10 common subjects. On the left, (a), an activation cluster in the left superior parietal lobe is clearly visible; whereas on the right, (b), the corresponding cluster does not stand out from noise.
5 Conclusion

We have presented a novel blind pattern-extraction algorithm for fMRI data. Our method, CanICA, is auto-calibrated and extracts the significant patterns according to a noise model. From these patterns, reproducible and meaningful features could be extracted. An important aspect of our method, specifically designed to perform group analysis, is that the features selected are more reproducible than other group-level ICA methods because it identifies a significantly-reproducible signal subspace and extracts localized features with a criteria consistent with the ICA algorithm.

CanICA is numerically efficient, as it relies solely on well-optimized linear algebra routines and performs the ICA optimization loop on a small number of components. Performance is important to scale to long fMRI time series, high-resolution data, or large groups. In addition, as the group-level pattern extraction (CCA and ICA) is very fast, cross-validation is feasible. ICA is an unstable algorithm with no intrinsic significance testing, but we have shown that cross-validation can be used to establish validity of group-level maps.

References


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3 a few minutes for our data on a 2 GHz Intel core Duo
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<tr>
<th></th>
<th>Non-thresholded maps</th>
<th>Thresholded maps</th>
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<tr>
<td></td>
<td>Group ICA</td>
<td>Tensor ICA</td>
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<tr>
<td>$e$ (standard deviation)</td>
<td>0.58 (0.04)</td>
<td>0.47 (0.06)</td>
</tr>
<tr>
<td>$t$  (standard deviation)</td>
<td>0.53 (0.04)</td>
<td>0.36 (0.03)</td>
</tr>
</tbody>
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Table 1. Reproducibility measures $e$ and $t$ for Group ICA, Tensor ICA and CanICA calculated on the half-split cross-correlation matrices, both for non-thresholded and thresholded maps. Numbers in parenthesis give the standard deviation of the mean.