

# Analysis of Functional Magnetic Resonance Imaging in Python

The authors describe a package for analyzing magnetic resonance imaging (MRI) and functional MRI (fMRI) data, which is part of the Neuroimaging in Python (NIPY) project. An international group of leading statisticians, physicists, programmers, and neuroimaging methodologists are developing NIPY for wider use.

agnetic resonance imaging (MRI) measures induced magnetic properties of tissue. It has long been the chosen technique for creating high-resolution anatomical images of the human brain. Over the past decade, a new technique called functional MRI (fMRI) has become a powerful and widely used method for studying human brain function. fMRI measures regional blood flow changes in the brain, which can help researchers identify the most active brain areas during mental tasks such as memory and language.

#### **Functional MRI Analysis**

The first step of an fMRI analysis—image reconstruction—takes raw data from the scanner and performs a highly customized inverse Fourier transform to create a time series of 3D functional images. A typical next step is to estimate the movement between scans via an automated image-matching algorithm and then use that estimate to remove artifacts due to

motion. Researchers commonly relate the activity detected in the low-resolution functional images to a high-resolution structural image of the same subject. However, to compare between subjects, the functional or structural data must be warped to match some standard brain, a process that requires sophisticated models of brain anatomy. Finally, statistical techniques are used to determine which brain regions are related to certain tasks or activities.

Clearly, fMRI data analysis comes with several challenges. First, it has a wide variety of computationally intensive spatial and statistical processing steps. Thus, an analysis software package must cover the range from file system and network interaction through complex image processing to advanced statistical inference and 3D visualization. Second, such analysis involves a massive volume of data, often reaching several hundred gigabytes.

The most common software package in use today is SPM (www.fil.ion.ucl.ac.uk/spm/), which is written in Matlab. Other common packages include FSL (www.fmrib.ox.ac.uk/fsl/) and AFNI (http://afni.nimh.nih.gov/afni/), written in C and C++, respectively. Although these packages are well-designed and supported, an increasing number of imaging scientists have found that Matlab is not powerful enough to support the industrial level of code size and complexity that neuroimaging requires, and that C and C++ are too low-level for rapid development.

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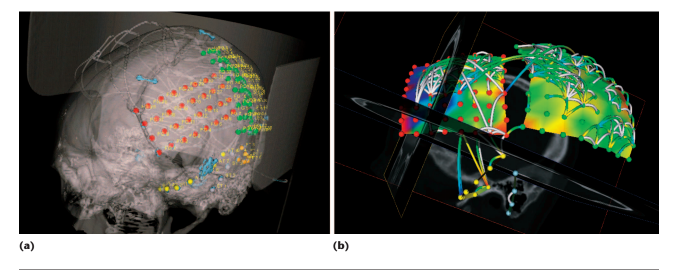


Figure 1. Python visualization widgets. Images from PBrain showing, (a) the position of the electrode arrays on the surface of the brain superimposed with a surface reconstruction of the skull from a CT scan and (b) measures of frequency and coherence of electrical activity overlaid on an image of the electrode positions.

## **Neuroimaging and Python**

Python has become a natural choice for neuroimaging because it is high level, object-oriented, and interactive. All these features make it particularly well suited to scientific programming. It also has robust libraries for system interaction, image processing, matrix algebra, and visualization. Moreover, Python has very good tools for providing scripting support to software written in other languages. Finally, because of Python's strong integration with C and C++, it is often used as a type of high-level language glue for calling routines in a huge array of high-quality C/C++ libraries.

Accordingly, several significant neuroimaging packages have already been written in Python. Led by Daniel Sheltraw at Berkeley, researchers recently developed a set of Python tools for MRI reconstruction (https://cirl.berkeley.edu/view/BIC/ReconTools). John Hunter, the author of the matplotlib Python plotting library, developed PBrain, a sophisticated application for analyzing and visualizing the data from electrical recordings on the brain surface of epileptic patients (see Figure 1 and p. 90). Finally, BrainVISA is a comprehensive pipeline-analysis tool for anatomical data, developed by a team of researchers in France (www.brainvisa.info).

# Integrating Python Development in Neuroimaging

In this article, our main focus is on NIPY (http://neuroimaging.scipy.org), a new initiative to create a unified, open source, and open development environment for the analysis of neuroimaging data.<sup>2</sup>

In particular, we focus on the fMRI component of NIPY, which is based on the BrainSTAT package that Jonathan Taylor wrote at Stanford University.<sup>3</sup> We are also working with Hunter and researchers at the University of Chicago to better integrate PBrain into the NIPY framework.

#### **Image Model**

An MRI scanner produces images that represent slices of brain tissue, and several of these slices together constitute a 3D whole-brain scan. The values in each voxel (volume element) in a 3D image slice represent measurements from a small volume of the brain.

A major problem in neuroimaging is that different analysis packages and scanners use different output image formats that are not readily compatible. To address this, NIPY provides read and write capabilities for all the popular image formats in neuroimaging, as well as access to binary data images and Python arrays in memory. Images can be compressed upon read or write, loaded from an arbitrary URL (with local caching), and managed by memory mapping where possible. Images also have iterators for reading data in slices and other subsets; Python iterators offer an elegant mechanism for constructing lazily instantiated sequential data structures—a perfect abstraction for intuitively representing sequential data (such as time series, spatial slices, or generally (n-1) - d data bricks from an n - d data set) without sacrificing memory efficiency.

Spatial transforms on images are fundamental to neuroimaging—for movement correction, warping to templates, and many other analysis steps. NIPY

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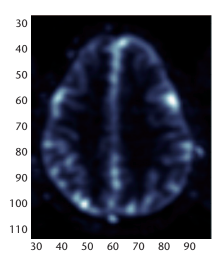


Figure 2. Image of standard deviation across time points. By summarizing data, the standard deviation image can highlight problems of background noise and artifacts that vary over time.

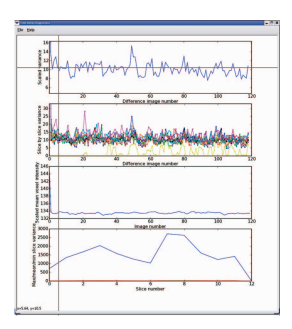


Figure 3. Time-series diagnostics. Researchers can use these four plots to diagnose potential problems in the time-series data.

offers a general model of image spatial transforms that allow arbitrary combinations of linear and nonlinear transforms, volume-to-surface mappings, and flexible levels of image interpolation. A common analysis technique is to examine the signal from a restricted region of the brain, or *region of interest* (ROI). NIPY has several ROI objects and functions, including discrete (point-level) and con-

tinuous (sphere, ellipse) region definitions, region combination algebra, and region data extraction.

#### **Data Diagnostics**

Because the scanner's signal quality can vary from day to day, special attention is needed to ensure good data. Currently, NIPY offers several diagnostic tools to let researchers discover problems with their data before they analyze it.

Images that summarize data across a time series provide a straightforward way to examine fMRI data. Taking an image of the standard deviation of signal intensity across time, for example, can highlight problems in the time series acquisition that occur with subject movement or instabilities of the data acquisition over time (see Figure 2).

Figure 3 shows four plots that can help diagnose potential problems in the time series. The top plot displays the scaled variance from image to image; the second plot shows the scaled variance per slice; the third plot shows the scaled mean voxel intensity for each image; and the bottom one plots the maximum, mean, and minimum scaled slice variances per slice.

Another powerful technique for data diagnosis is principal components analysis (PCA), which is particularly useful for detecting outlying time points or unexpected sources of spatially coherent noise. PCA on an image time series takes the time points as rows and voxels as columns, and decomposes the data into components that can efficiently express the source of variance in the data. Each component consists of a characteristic time series and the voxel weights contributing to that component, and these weights are viewable as an image. Figure 4 shows images of the weights for the first four principal components of a functional data set.

#### Statistical Processing

Subjects in a typical fMRI experiment perform some task or receive stimuli while being scanned; thus, areas activated by the task or stimulus exhibit voxel time series correlated with the experimental design. Unfortunately, noise in fMRI analysis can come from multiple sources, and the signal is relatively low. This problem with signal detection has led to several standard and more complex statistical methods. Partly for historical reasons, current analysis packages use nonstandard or low-level statistical terminology and interfaces, making them less accessible to scientists with general training in statistics.

At Stanford, Taylor developed a general set of statistical model objects in Python that use standard statistical terminology and allow flexible high-level statistical designs. Similar to the R statistical language, these objects implement a general series

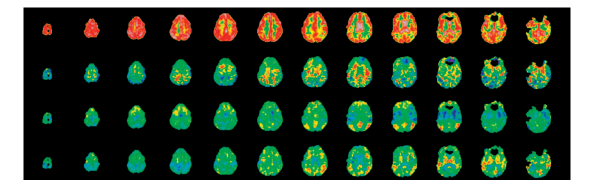


Figure 4. Principal components analysis (PCA). This type of analysis is particularly useful for detecting outlying time points or unexpected sources of spatially coherent noise.

of statistical models and contrasts for data, including functional images. These models form the basis of NIPY statistical analysis and are now maintained as part of the SciPy distribution; we hope these will attract further development from researchers outside brain imaging.

ython's power and generality mean that we have many fruitful directions in which to take NIPY. At the most basic, its high-level language features make it much easier to refactor the code into a wellpatterned, high-level interface that scientific developers can quickly pick up. Because Python has such good integration with C/C++, we also have ready access to very powerful visualization and image-processing libraries. Two very important examples are the Visualization Toolkit (VTK) and the Insight Toolkit (ITK). VTK provides high-quality 3D graphical display and includes Python wrappers as part of its standard distribution, whereas ITK is a library of image registration and segmentation routines originally developed for the Visible Human Project. Like VTK, Python wrappers are part of the standard ITK distribution.

We intend NIPY to become the standard analysis library in neuroimaging in the medium term, which means we will need to provide the ability to call routines in other packages that are more familiar to researchers. Thankfully, this is a much easier task in Python than in many other languages because of its ability to interact with languages such as C.

Python's language features can also help us tackle the problem of provenance tracking. Because imaging analyses and data sets are very diverse, researchers use a variety of analysis packages and rarely record all their data and analysis parameters, making it very difficult for other people to reproduce published analyses. Fortunately, Python has excellent support for metaprogramming techniques (including metaclasses and function decorators) that can transparently change object behavior. We can thus use these techniques with Python's object introspection to capture the data's nature and processing in a way that can be closely wedded to the analysis. Ultimately, this means that the analysis can become self-documenting, making it far easier to reproduce.

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