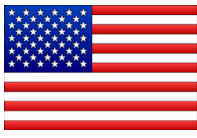


AFNI & FMRI

Introduction, Concepts, Principles



Analysis of Functional NeuroImages

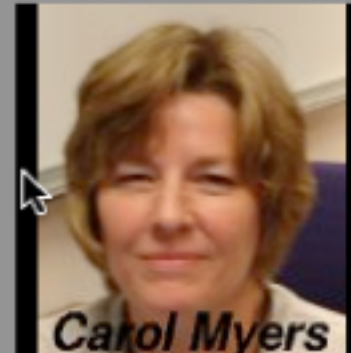
by

Robert W Cox, PhD

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Public License Version 2 (GPL)
[or any later GPL version]

AFNI is a research tool.

*Clinical uses are **not** supported or advised.*



Carol Myers

AFNI User



<http://afni.nimh.nih.gov/afni>

AFNI = Analysis of Functional NeuroImages

- Developed to provide an environment for FMRI data analyses
 - And a platform for development of new software
- **AFNI** refers to both the program of that name and the entire package of external programs and plugins (more than 200)
- Important principles in the development of AFNI:
 - Allow user to stay close to the data and view it in many different ways
 - Give users the power to assemble pieces in different ways to make customized analyses
 - “With great power comes great responsibility”
 - **to understand the analyses and the tools**
 - “Provide mechanism, not policy”
 - Allow other programmers to add features that can interact with the rest of the package

Principles (and Caveats) We* Live By

- Fix significant bugs as soon as possible
 - But, we define “significant”
- Nothing is secret or hidden (AFNI is open source)
 - But, possibly not very well documented or advertised
- Release early and often
 - All users are beta-testers for life
- Help the user (message board; consulting with NIH users)
 - Until our patience expires
- Try to anticipate users’ future needs
 - What we think you will need may not be what you actually end up needing

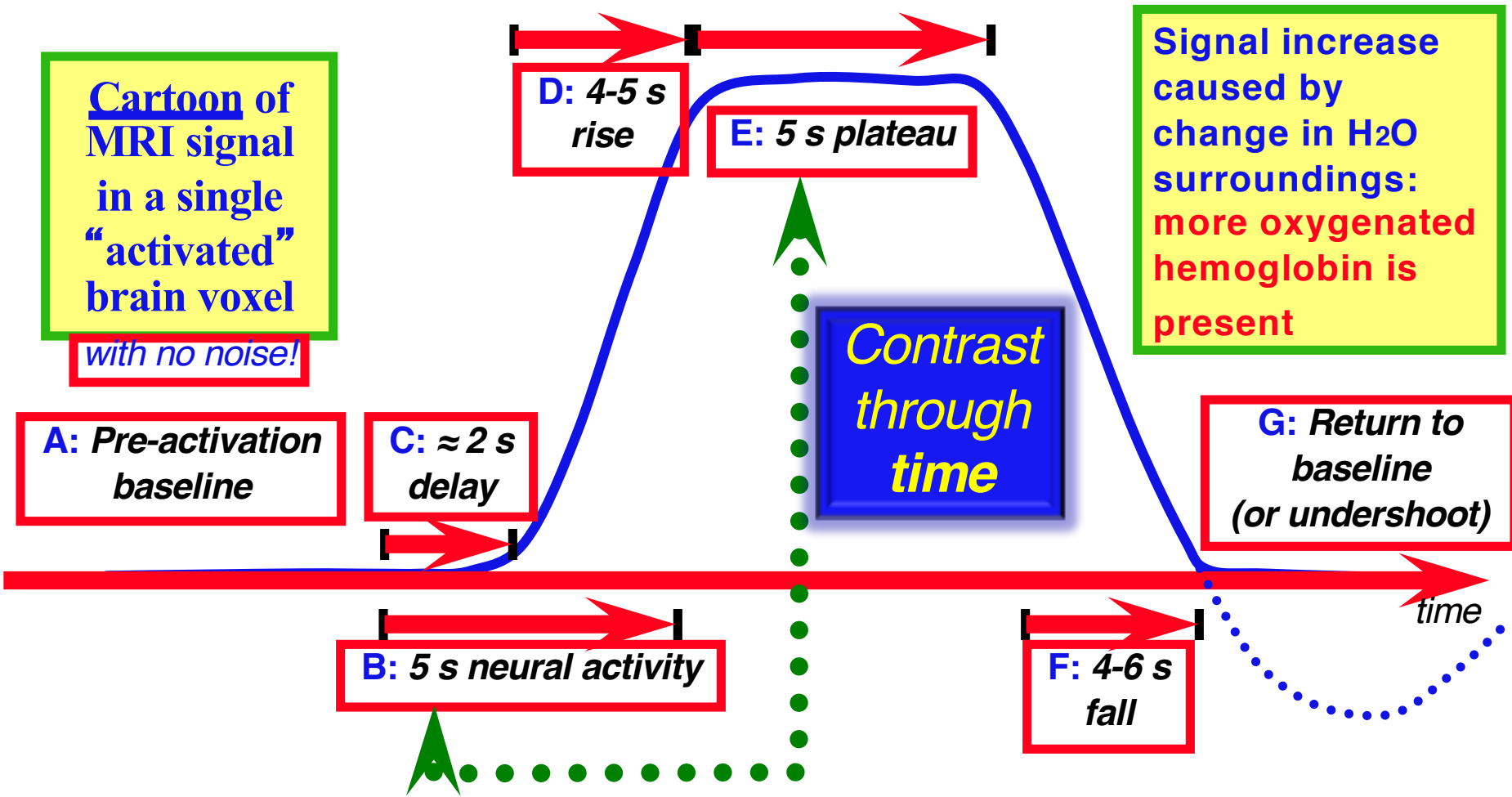


Before We Really Start

- AFNI has many programs and they have many options
- Assembling the programs to do something useful and good seems confusing (OK, *is* confusing) when you start
- To help overcome this problem, we have “super-scripts” that carry out important tasks
 - Each script runs multiple AFNI programs
 - We recommend using these as the basis for FMRI work
 - When you need help, it will make things simpler for us *and* for you if you are using these scripts
- **afni_proc.py** = Single subject FMRI pre-processing and time series analysis for functional activation
 - **uber_subject.py** = GUI for **afni_proc.py**
- **align_epi_anat.py** = Image alignment (registration), including anatomical-EPI, anatomical-anatomical, EPI-EPI, and alignment to atlas space (Talairach/MNI)

What is Functional MRI?

- 1991: Discovery that MRI-measurable signal increases a few % *locally* in the brain subsequent to increases in neuronal activity (Kwong, *et al.*)



How fMRI Experiments Are Done

- Alternate subject's neural state between 2 (or more) conditions using sensory stimuli, tasks to perform, ...
 - Can only measure relative signals, so must look for *changes* in the signal between the conditions

- Acquire MR images repeatedly during this process

- Search for voxels whose NMR signal time series (up-and-down) matches the stimulus time series pattern (on-and-off)
 - ➔ ▪ fMRI data analysis is basically pattern matching *in time*

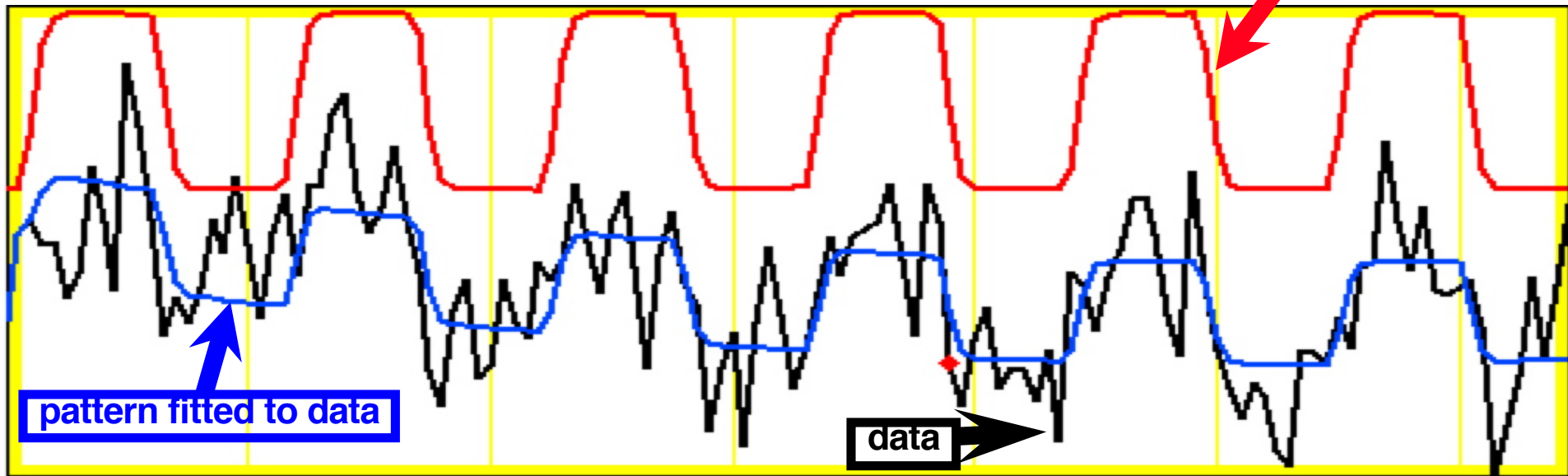
- Signal changes due to neural activity are small
 - Need 500 or so images in time series (in each slice) ➔ takes 30 min or so to get reliable activation maps
 - Usually break image acquisition into shorter "runs" to give the subject and scanner some break time
 - Other small effects can corrupt the results ➔ post-process the data to reduce these effects & *be vigilant*

- Lengthy computations for image recon and temporal pattern matching ➔ data analysis usually done offline

Sample Data Time Series

- 64×64 matrix (TR=2.5 s; 130 time points per imaging run)
- Somatosensory task: 27 s “on”, 27 s “rest”
- Note that this is *really* good data

pattern of expected BOLD signal



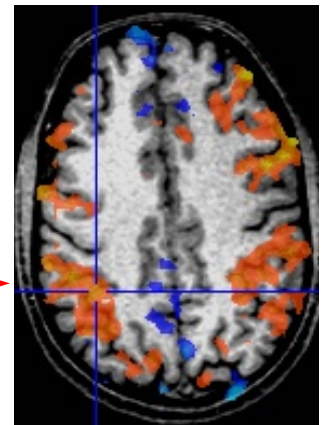
pattern fitted to data

data



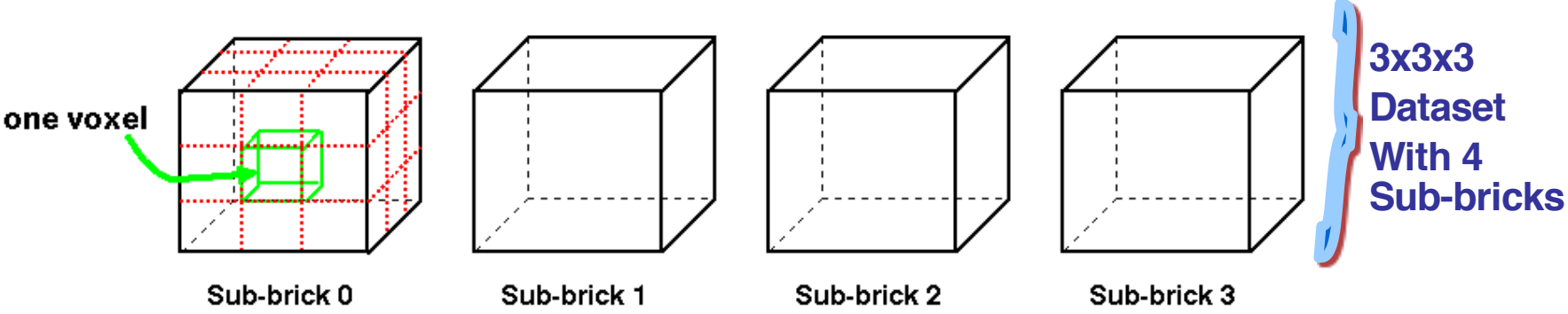
One echo-planar image

One anatomical image, with voxels that match the pattern given a color overlay



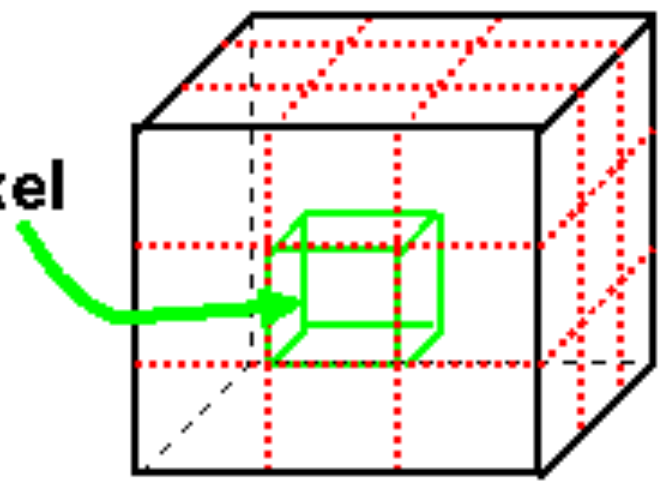
Fundamental AFNI Concepts

- Basic unit of data in AFNI is the dataset ← **Jargon!**
 - A collection of 1 or more 3D arrays of numbers
 - Each entry in the array is in a particular spatial location in a 3D grid (a voxel = 3D pixel)
 - Image datasets: each array holds a collection of slices from the scanner
 - Each number is the signal intensity for that particular voxel
 - Derived datasets: each number is computed from other dataset(s)
 - e.g., each voxel value is a *t*-statistic reporting “activation” significance from an fMRI time series dataset, for that voxel
 - Each 3D array in a dataset is called a sub-brick ← **Jargon!**
 - There is one number in each voxel in each sub-brick

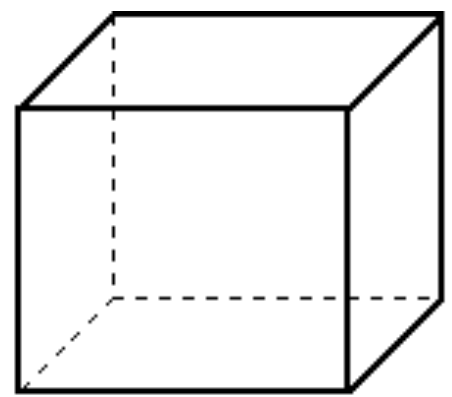


A Little Bit Bigger

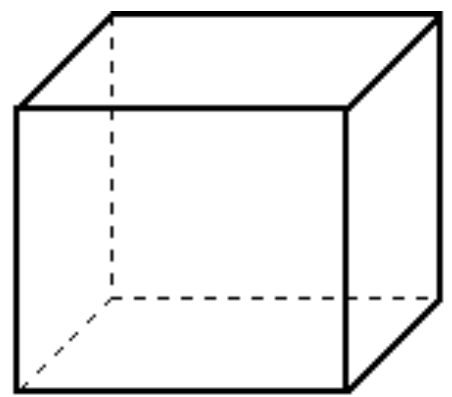
one voxel



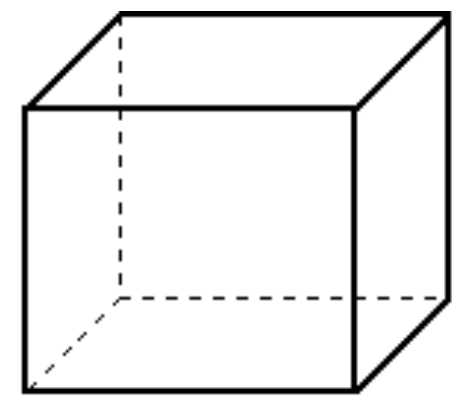
Sub-brick 0



Sub-brick 1



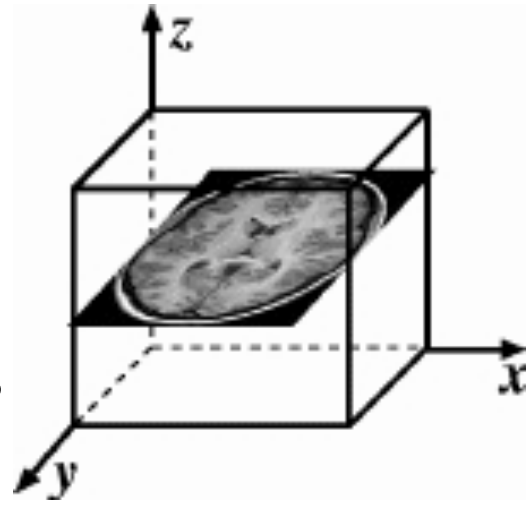
Sub-brick 2



Sub-brick 3

What's in a Dataset: Header Stuff


- Besides the voxel numerical values, a dataset also contains auxiliary information, including (some of which is optional):
 - xyz dimensions of each voxel (in mm)
 - Orientation of dataset axes;
for example, x-axis=R-L, y-axis=A-P, z-axis=I-S
= axial slices (we call this orientation “RAI”)
 - Location of dataset in scanner coordinates
 - Needed to overlay one dataset onto another
 - Very important to get right in FMRI, since we deal with many datasets
 - Time between sub-bricks, for 3D+time datasets ← **Jargon!**
 - Such datasets are the basic unit of FMRI data (one per imaging run)
 - Statistical parameters associated with each sub-brick
 - e.g., a *t*-statistic sub-brick has degrees-of-freedom parameter stored
 - e.g., an *F*-statistic sub-brick has 2 DOF parameters stored
 - Et cetera, et cetera, et cetera ...



AFNI Dataset Files - 1

- AFNI formatted datasets are stored in 2 files
 - The .HEAD file holds all the auxiliary information
 - The .BRIK file holds all the numbers in all the sub-bricks
- Datasets can be in one of ~~3~~ 2 coordinate systems (“views”)
 - Original data or +orig view: from the scanner
 - ~~AC-PC aligned or +acpc view:~~
 - ~~◦ Dataset rotated/shifted so that the anterior commissure and posterior commissure are horizontally axis, the AC is at (0,0,0) and the transverse commissure is vertical z-axis)~~
 - Talairach or +tlrc view:
 - Dataset has also been rescaled to conform to the Talairach-Tournoux atlas dimensions (RL=136 mm; AP=172 mm; IS=116 mm)
 - AKA Talairach or Stererotaxic coordinates
 - Not quite the same as MNI coordinates, but very close
 - **All** datasets scaled+aligned to some atlas are labeled +tlrc
 - Header can contain name of actual atlas “space” (e.g., MNI)

AFNI Dataset Files - 2

- AFNI dataset filenames consist of 3 parts
 - The user-selected prefix (almost anything) 
 - The view (one of +orig, ~~+axpc~~, or +tlrc)
 - The suffix (one of .HEAD or .BRIK)
 - Example: **BillGates+tlrc.HEAD** and **BillGates+tlrc.BRIK**
 - When creating a dataset with an AFNI program, you supply the prefix; the program supplies the rest

- AFNI programs can *read* datasets stored in several formats
 - ANALYZE (.hdr/.img file pairs); i.e., from SPM, FSL
 - MINC-1 (.mnc); i.e., from mnitools [but not MINC-2]
 - CTF (.mri, .svl) MEG analysis volumes
 - ASCII text (.1D) — numbers arranged into columns
 - Have conversion programs to write out MINC-1, ANALYZE, ASCII, and NIfTI-1.1 files from AFNI datasets, if desired

NIfTI Dataset Files

- NIfTI-1 ([.nii](#) or [.nii.gz](#)) is a standard format that AFNI, SPM, FSL, BrainVoyager, et al., have agreed upon
 - Adaptation and extension of the old ANALYZE 7.5 format
 - Goal: easier interoperability of tools from various packages
- All data is stored in 1 file (cf. <http://nifti.nimh.nih.gov/>)
 - 348 byte header (extensions allowed; AFNI uses this feature)
 - Followed by the image binary numerical values
 - Allows 1D–5D datasets of diverse numerical types
 - [.nii.gz](#) suffix means file is compressed (with gzip)
- AFNI now reads and writes NIfTI-1 (and NIfTI-2) datasets
 - **To write:** when you give the [prefix](#) for the output filename, end it in “[.nii](#)” or “[.nii.gz](#)”, and all AFNI programs will automatically write NIfTI-1.1 format instead of [.HEAD/.BRIK](#)
 - **To read:** just give the full filename ending in “[.nii](#)” or “[.nii.gz](#)”

Getting and Installing AFNI

- AFNI runs on Unix systems: Linux, Sun, Mac OS X
 - Can run under Windows with Cygwin Unix emulator
 - This option is really just for trying it out — not for production use!
- You can download precompiled binaries from our Website
 - <http://afni.nimh.nih.gov/afni>
 - Also: documentation, message board, humor, data, class materials, ...
- You can download source code and compile it
 - Also from GitHub: <https://github.com/afni/AFNI>
- AFNI is updated fairly frequently, so it is important to update occasionally -- [@update.afni.binaries](#)
 - We can't help you with outdated versions!
 - ***Please check for updates every 6 months (or less)***

AFNI at the NIH Scanners

- AFNI can take 2D images in “realtime” from an external program and assemble them into 3D+time datasets slice-by-slice
- FMRI Facility scanners at the NIH (GE and Siemens) are set up to start AFNI on a remote Linux computer automatically when EPI acquisition starts, and then the Dimon program is used to send images into AFNI as they are reconstructed:
 - For immediate display (images and graphs of time series)
 - **Plus**: graphs of estimated subject head movement
- Goal is to let you see image data as they are acquired, so that if there are any big problems, you can fix them right away
 - Sample problem: someone typed in the imaging field-of-view (FOV) size wrong (240 cm instead of 24 cm), and so got garbage data, ***but only realized this too late*** (after scanning 8 subjects this way) — ***D’oh!***

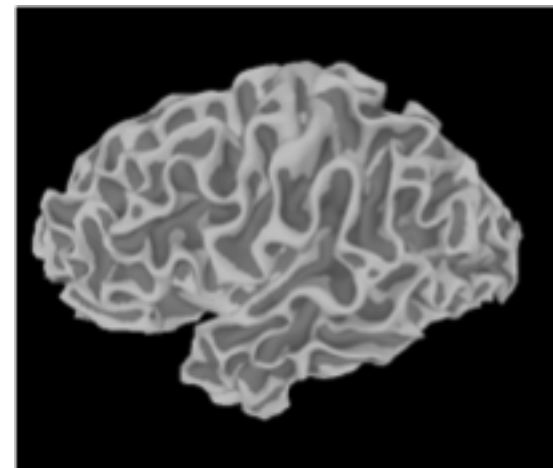
Other Parts of AFNI

- Batch mode programs and scripts
 - Are run by typing commands directly to computer, or by putting commands into a text file (script) and later executing them
- Good points about batch mode
 - Can process new datasets exactly the same as old ones
 - Can link together a sequence of programs to make a customized analysis (a personalized pipeline)
 - Some analyses take a long time (are not interactive)
- Bad points about batch mode
 - Learning curve is “all at once” rather than gradual
 - If you are, like, under age 35, you may not know how to, like, type commands into a computer to make it do things
 - But we don't make you use punched cards or paper tape (yet)

AFNI Batch Programs

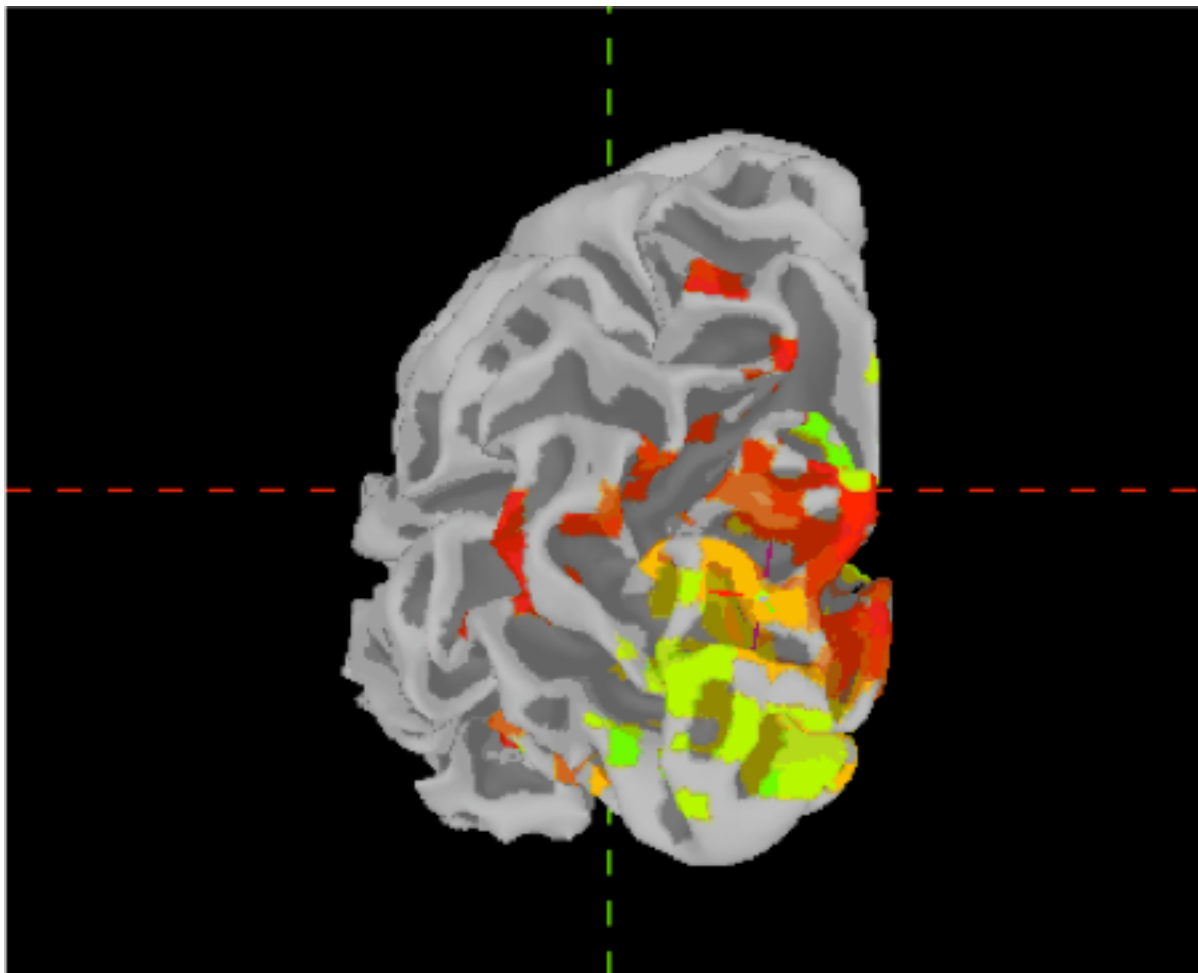
- Many many important capabilities in AFNI are **only** available in batch programs
 - A few examples (of more than 100, from trivial to complex)
- **3dDeconvolve** + **3dREMLfit** = multiple *linear* regression on 3D+time datasets; fits each voxel's time series to activation model, tests these fits for significance (**3dNLfim** = nonlinear fitting)
- **3dvolreg** = 3D+time dataset registration, to correct for small subject head movements, and for inter-day head positioning
- **3dANOVA** + **3dLME** = 1-, 2-, 3-, and 4- way ANOVA/LME layouts: combining & contrasting datasets in Talairach space
- **3dcalc** = general purpose voxel-wise calculator (very useful)
- **3dsvm** = SVM multi-voxel pattern analysis program
- **3dresample** = re-orient and/or re-size dataset voxel grid
- **3dSkullStrip** = remove “skull” from anatomical dataset
- **3dDWItoDT** = compute diffusion tensor from DWI (nonlinearly)

SUMA, et alii



- **SUMA** is the AFNI surface mapper
 - For displaying surface models of cortex
 - Surfaces from **FreeSurfer** (MGH) or **Caret** (Wash U) or **BrainVoyager** (Brain Innovation)
 - Can display functional activations mapped from 3D volumes to the cortical surface
 - Can draw ROIs directly on the cortical surface
 - vs. AFNI: ROIs are drawn into the 3D volume
- SUMA is a separate program from AFNI, but can “talk” with AFNI (like a plugout) so that volume & surface viewing are linked
 - Click in AFNI or SUMA to change focus point, and the other program jumps to that location at the same time
 - Functional (color) overlay in AFNI can be sent to SUMA for simultaneous display
- And much more — stayed tuned for the SUMA talks to come!

SUMA Teaser Movie



Color from AFNI, Images from SUMA
Images captured with the 'R' recorder function,
then saved as animation with [Save:aGif](#) control

Other Educational Presentations

- ~~• How to get images into AFNI or NIFTI format (program **to3d**)~~
- Detailed hands-on with using AFNI for data viewing (**fun**)
- Signal modeling & analysis: theory & hands-on (**3dDeconvolve et al.**)
- Image registration (**3dvolreg et al.**)
- ~~• Volume rendering hands-on (**fun level=high**)~~
- ~~• ROI drawing hands-on (**fun level=extreme**)~~
- ~~• Transformation to Talairach hands-on (**fun level=low**)~~
- Group analysis: theory and hands-on (**3dANOVAX** *and beyond*)
- ~~• Experiment design~~
- FMRI analysis from start to end (the “soup to nuts” hands-on)
- SUMA hands-on (**fun level=pretty good**)
- Surface-based analysis
- Connectivity (resting state, white matter tracts)
- AFNI “Jazzercise” (practice sessions & directed exercises)