## FMRI Connectivity Analysis in AFNI

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## Structure of this lecture

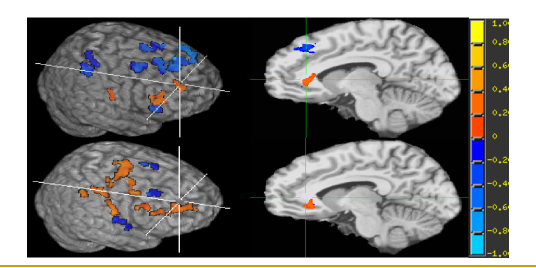
- Overview
- Correlation analysis
  - Simple correlation
  - Context-dependent correlation (PPI)
- Structural equation modeling (SEM)
  - Model validation
  - Model search
- Granger causality (GC)
  - □ Bivariate: exploratory ROI search
  - Multivariate: validating path strength among pre-selected ROIs

## Overview: FMRI connectivity analysis

- All about FMRI
  - Not for DTI
  - Some methodologies may work for MEG, EEG-ERP
- Information we have
  - Anatomical structures
    - Seed-based: A seed region in a network, or
    - Network-based: A network with all relevant regions known
  - □ Brain output (BOLD signal): regional time series
- What can we say about inter-regional communications?
  - Inverse problem: make inference about intra-cerebral neural processes from extra-cerebral/vascular signal
  - Based on response similarity (and sequence)

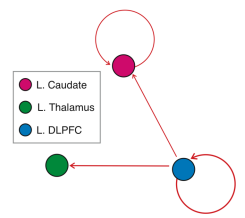
## Approach I: seed-based; ROI search

- Regions involved in a network are unknown
  - □ Bi-regional (seed vs. whole brain) (3d\*): brain volume as input
  - Mainly for ROI search
  - □ Popular name: functional connectivity
  - Basic, coarse, exploratory with weak assumptions
  - □ Methodologies: simple correlation, PPI, bivariate GC
  - □ Weak interpretation: may or may not indicate directionality/causality



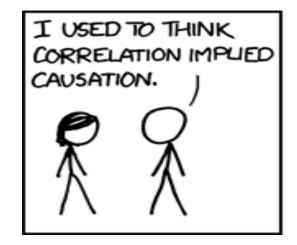
## Approach II: network-based

- Regions in a network are known
  - Multi-regional (1d\*): ROI data as input
  - Model validation, connectivity strength testing
  - Popular name: effective or structural connectivity
  - Strong assumptions: specific, but with high risk
  - □ Methodologies: SEM, multivariate GC, DCM
  - Directionality, causality (?)

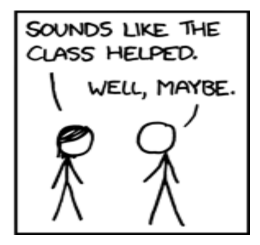


### Interpretation Trap: Correlation vs. Causation!

- Some analyses require fine time resolution we usually lack
- Path from (or correlation btw) A to (and) B doesn't necessarily mean causation
  - Bi-regional approach simply ignores the possibility of other regions involved
  - Analysis invalid if a relevant region is missing in a multi-regional model
- Robust: connectivity analysis < regression analysis</li>
- Determinism in academics and in life
  - Linguistic determinism: Sapir-Whorf hypothesis







## Preparatory Steps

- Warp brain to standard space
  - adwarp, @auto-tlrc, align\_epi\_anat.py
- Create ROI
  - □ Sphere around a peak activation voxel: **3dUndump** —**master** ... —**srad** ...
  - □ Activation cluster-based (biased unless from independent data?): localizer
  - Anatomical database
  - Manual drawing
- Extract ROI time series
  - □ Average over ROI: 3dmaskave —mask, or 3dR0Istats —mask
  - Principal component among voxels within ROI: 3dmaskdump, then 1dsvd
  - □ Seed voxel with peak activation: **3dmaskdump** -noijk -dbox
- Remove effects of no interest
  - 3dSynthesize and 3dcalc
  - □ 3dDetrend —polort
  - □ RETROICORR/RetroTS.m
  - 3dBandpass

# Simple Correlation Analysis

- Seed vs. rest of brain
- ROI search based on response similarity
  - Looking for regions with similar signal to seed
- Correlation at individual subject level
  - □ Usually have to control for effects of no interest: drift, head motion, physiological variables, censored time points, tasks of no interest, etc.
- Applying to experiment types
  - □ Straightforward for resting state experiment: default mode network (DMN)
  - With tasks: correlation under specific condition(s) or resting state?
- Program: 3dfim+ or 3dDeconvolve
  - ightharpoonup r: not general, but **linear**, relation; slope for standardized Y and X
  - ullet eta: slope, amount of linear change in Y when X increases by 1 unit
- Website: http://afni.nimh.nih.gov/sscc/gangc/SimCorrAna.html
- Interactive tools in AFNI and SUMA: InstaCor, GroupInstaCor

# Simple Correlation Analysis

### Group analysis

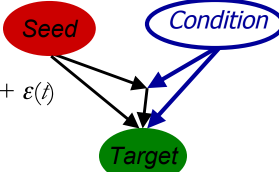
- Run Fisher-transformation of r to Z-score and t-test: **3dttest**
- $\square$  Take  $\beta$  and run *t*-test (pseudo random-effects analysis): **3dttest**
- □ Take  $\beta$  + t-statistic and run random-effects model: **3dMEMA**

### **Caution**: don't over-interpret

- Not proof for anatomical connectivity
- No golden standard procedure and so many versions in analysis: seed region selection, covariates,  $r(Z)/\beta$ , bandpass filtering, ...
- □ Information limited if other regions present in network
- Be careful with group comparison (normal vs. disease): assuming within-group homogeneity, can we claim
  - No between-group difference → same correlation/connectivity across groups?
  - o Between-group difference → different correlation/connectivity across groups?

## Context-Dependent Correlation

- Popularized name: Psycho-Physiological Interaction (PPI)
- 3 explanatory variables
  - $\Box$  Condition (or contrast) effect: C(t)
  - $\Box$  Seed effect on rest of brain: S(t)
  - □ Interaction between seed and condition (or contrast): I(C(t), S(t))
    - Directionality here!
- Model for each subject
  - Original GLM:  $y = [C(t) \text{ Others}] \beta + \varepsilon(t)$
  - New model:  $y = [C(t) S(t) I(C(t), S(t)) Others] \beta + \varepsilon(t)$
  - 2 more regressors than original model
  - Others NOT included in SPM
  - □ What we care for: r or  $\beta$  for I(C(t), S(t))



# Context-Dependent Correlation

- How to formulate I(C(t), S(t))?
  - Interaction occurs at neuronal, not BOLD (an indirect measure) level
  - □ **Deconvolution**: derive "neuronal response" at seed based on BOLD response
    - **3dTfitter:** Impulse ⊗ Neuronal events = BOLD response
  - A difficult and an inaccurate process!
  - Deconvolution matters more for event-related than block experiments
  - □ Useful tool: **timing\_tool.py** can convert stimulus timing into 0s and 1s
- If stimuli were presented in a resolution finer than TR
  - Use **1dUpsample n**: interpolate time series  $n \times$  finer before deconvolution **3dTffiter**
  - Downsample interaction regressor back to original TR with **1dcat** with selector '{0..\$(n)}'
- Group analysis
  - $\square$  Run Fisher-transformation of r to Z-score and t-test: **3dttest**
  - $\square$  Take  $\beta$  and run t-test (pseudo random-effects analysis): **3dttest**
  - $\Box$  Take  $\beta$  and t-statistic and run random-effects model: **3dMEMA**
- Website: http://afni.nimh.nih.gov/sscc/gangc/CD-CorrAna.html

# PPI Caution: avoid over-interpretation

- Not proof for anatomical connectivity
- Information limited if other regions involved in the network
- Neuronal response is hard to decode: Deconvolution is very far from reliable, plus we have to assume a shape-fixed HRF (same shape regardless of condition or regions in the brain)
- Doesn't say anything about interaction between seed and target on seed
- Doesn't differentiate whether modulation is
  - Condition on neuronal connectivity from seed to target, or
  - □ Neuronal connectivity from seed to target on condition effect
- Be careful with group comparison (normal vs. disease group): assuming within-group homogeneity, can we claim
  - No between-group difference => same correlation/connectivity across groups?
  - □ Between-group difference => different correlation/connectivity across groups?

## Context-Dependent Correlation: hands-on

#### Data

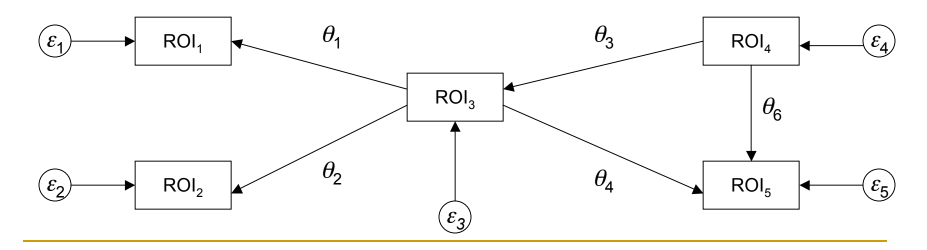
- □ Downloaded from <a href="http://www.fil.ion.ucl.ac.uk/spm/data/attention/">http://www.fil.ion.ucl.ac.uk/spm/data/attention/</a>
- Event-related attention to visual motion experiment
- □ 4 conditions: fixation, stationary, attention motion (att), no attention motion (natt)
- $\square$  TR=3.22s, 360 time points = 90 TR's/run × 4 runs, seed ROI = V2
- □ All steps coded in commands.txt: tcsh –x commands.txt (~5 minutes)

#### Should effects of no interest be included in PPI model?

Compare results between AFNI and SPM

### Structural Equation Modeling (SEM) or Path Analysis

- All possible regions involved in network are included
- All regions are treated equally as endogenous (dependent) variable
- Residuals (unexplained) are exogenous (independent) variables
- Analysis based on summarized data (not original ROI times series) with model specification, covariance/correlation matrix,
   DF and residual error variances (?) as input



# SEM: theory

- Hypothetical model  $X = KX + \varepsilon$ 
  - $\square$  X: *i*-th row  $x_i(t)$  is *i*-th ROI time series
  - $\Box$  K: matrix of path coefficients  $\theta$ 's whose diagonals are all 0's
  - $\bullet$   $\varepsilon$ : *i*-th row  $\varepsilon_i(t)$  is residual time series of *i*-th ROI
- Predicted (theoretical) covariance

$$\Sigma(\theta) = (I-K)^{-1}E[\varepsilon(t)\varepsilon(t)^T][(I-K)^{-1}]^T$$
 as  $X = (I-K)^{-1}\varepsilon$ 

■ ML discrepancy/cost/objective function btw predicted and estimated covariance (*P*: # of ROIs)

$$F(\theta) = \ln \sum_{n} (\theta) \left| + tr[C\sum_{n} (\theta)] - \ln |C| - P \right|$$

- □ Input: model specification; covariance (correlation?) matrix C; DF (calculating model fit statistic chi-square); residual error variances?
- □ Usually we're interested in a network under resting state or specific condition

## SEM: 1st approach - validation

- Knowing directional connectivity btw ROIs, data support model?
- Null hypothesis  $H_0$ : It's a good model
- If  $H_0$  is **not** rejected, what are the path strengths, plus fit indices?
- Analysis for whole network, path strength estimates by-product
- 2 programs
  - □ **1dSEM** in C
    - Residual error variances as input (DF was a big concern due to limited number of time points)
    - Group level only; no CI and p value for path strength
    - Based on <u>Bullmore et al.</u>, How Good is Good Enough in Path Analysis of fMRI Data? NeuroImage 11, 289-301 (2000)
  - □ **1dSEMr.R** in R
    - Residual error variances not used as input
    - CI and *p* value for path strength
    - Individual and group level

## SEM: 2nd approach - search

- All possible ROIs known with some or all paths are uncertain
- Estimate unknown path strengths
- Start with a minimum model (can be empty)
- Grow (add) one path at a time that lowers cost
- How to add a path?
  - □ Tree growth: branching out from previous generation
  - □ Forest growth: whatever lowers the cost no inheritance
- Program 1dSEM: only at group level
- Various fit indices other than cost and chi-square:
  - □ AIC (Akaike's information criterion)
  - □ RMSEA (root mean square error of approximation)
  - CFI (comparative fit index)
  - □ GFI (goodness fit index)

## SEM: caution I

- Correlation or covariance: What's the big deal?
  - □ Almost **ALL** publications in FMRI use correlation as input
  - $\Box$  A path connecting from region A to B with strength  $\theta$ 
    - Not correlation coefficient
    - o If A increases by one SD from its mean, B would be expected to increase by  $\theta$  units (or decrease if  $\theta$  is negative) of its own SD from its own mean while holding all other relevant regional connections constant
    - With correlation as input
      - May end up with different connection and/or path sign
      - Results are not interpretable
      - o Difficult to compare path strength across models/groups/studies,...
  - □ Scale ROI time series to 1 (instead of 100 as usual)
- □ ROI selection very important
  - □ If one ROI is left out, whole analysis (and interpretation) would be invalid

### SEM: caution II

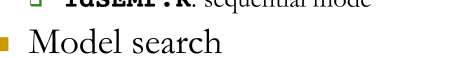
#### Validation

- □ It's validation, not proof, when not rejecting null hypothesis
- □ Different network might be equally valid, or even with lower cost: model comparison possible if nested
- Search: How much faith can we put into final 'optimal' model?
  - □ Model comparison only meaningful when nested (tree > forest?)
  - □ Is cost everything considering noisy FMRI data? (forest > tree?)
  - □ Fundamentally SEM is about validation, not discovery
- Only model regional relationship at current moment
  - $\Box X = KX + \varepsilon$
  - No time delays

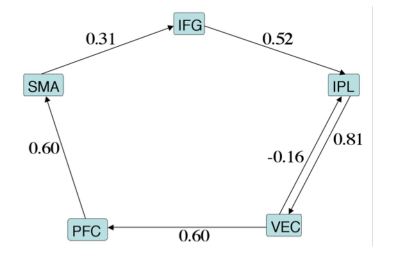
### SEM: hands-on

#### Model validation

- □ Data: Bullmore *et al.* (2000)
- Correlation as input
- □ Residual error variances as input
- □ **SEMscript.csh** maybe useful
- □ **1dSEM**: tcsh —x commands.txt
- □ **1dSEMr.R**: sequential mode



- Data courtesy: Ruben Alvarez (MAP/NIMH/NIH)
- □ 6 ROIs: PHC, HIP, AMG, OFC, SAC, INS
- □ Tree growth
- □ Covariance as input for **1dSEM**
- □ Shell script **SEMscript.csh** taking subject ROI time series and minimum model as input: tcsh –x commands.txt (~10 minutes)



# Granger Causality: introduction

- Classical univariate autoregressive model AR(p)

  - □ Current state depends linearly on immediate past ones with a random error
  - □ Why called autoregressive?
    - Special multiple regression model (on past *p* values)
    - Dependent and independent variable are the same
- What we typically deal with in GLM
  - $y = X\beta + \varepsilon$ ,  $\varepsilon \sim N(0, \sigma^2 V)$ ,  $\sigma^2$  varies spatially (across voxels)
  - Difficulty: V has some structure (e.g., ARMA(1,1) in **3dREMLfit**) and may vary spatially
  - $\square$  We handle autocorrelation structure in noise  $\varepsilon$
  - Sometimes called time series regression

## Rationale for Causality in FMRI

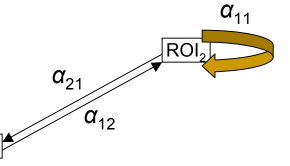
- Networks in brain should leave some signature (e.g, latency) in fine texture of BOLD signal because of dynamic interaction among ROIs
- Response to stimuli does not occur simultaneously across brain: latency
- Reverse engineering: signature may reveal network structure
- Problem: latency might be due to neurovascular differences!

## Start simple: bivariate AR model

- Granger causality: A Granger causes B if
  - the time series at A provides statistically significant information about the time series at B at some time delays (order)
- 2 ROI time series,  $y_1(t)$  and  $y_2(t)$ , with a VAR(1) model

$$y_{1}(t) = \alpha_{10} + \alpha_{11}y_{1}(t-1) + \alpha_{12}y_{2}(t-1) + \varepsilon_{1}(t)$$

$$y_{2}(t) = \alpha_{20} + \alpha_{21}y_{1}(t-1) + \alpha_{21}y_{2}(t-1) + \varepsilon_{2}(t)$$



### Assumptions

Linearity

- Stationarity/invariance: mean, variance, and autocovariance
- □ White noise, positive definite contemporaneous covariance matrix, and no serial correlation in individual residual time series
- Matrix form:  $Y(t) = \alpha + AY(t-1) + \epsilon(t)$ , where

$$Y(t) = \begin{bmatrix} y_1(t) \\ y_2(t) \end{bmatrix} \qquad \alpha = \begin{bmatrix} \alpha_{10} \\ \alpha_{20} \end{bmatrix} \qquad A = \begin{bmatrix} \alpha_{11} & \alpha_{12} \\ \alpha_{21} & \alpha_{22} \end{bmatrix} \quad \varepsilon(t) = \begin{bmatrix} \varepsilon_1(t) \\ \varepsilon_2(t) \end{bmatrix}$$

## Multivariate AR model

• n ROI time series,  $y_1(t), \dots, y_n(t)$ , with VAR(p) model

$$y_{1}(t) = \alpha_{10} + \sum_{k=1}^{p} \alpha_{11k} y_{1}(t-k) + \dots + \sum_{k=1}^{p} \alpha_{1nk} y_{n}(t-k) + \varepsilon_{1}(t)$$

$$\vdots$$

$$y_{n}(t) = \alpha_{20} + \sum_{k=1}^{p} \alpha_{n1k} y_{1}(t-k) + \dots + \sum_{k=1}^{p} \alpha_{nnk} y_{n}(t-k) + \varepsilon_{n}(t)$$

• Hide ROIs:  $Y(t) = \alpha + A_1 Y(t-1) + \dots + A_p Y(t-p) + \varepsilon(t)$ ,

$$Y(t) = \alpha + \sum_{i=1}^{p} A_{i}Y(t-i) + \varepsilon(t) \quad \alpha = \begin{bmatrix} \alpha_{10} \\ \vdots \\ \alpha_{n0} \end{bmatrix} Y(t) = \begin{bmatrix} y_{1}(t) \\ \vdots \\ y_{n}(t) \end{bmatrix} \quad A_{i} = \begin{bmatrix} \alpha_{11i} & \cdots & \alpha_{1ni} \\ \vdots & \ddots & \vdots \\ \alpha_{n1i} & \cdots & \alpha_{n1i} \end{bmatrix} \varepsilon(t) = \begin{bmatrix} \varepsilon_{1}(t) \\ \vdots \\ \varepsilon_{n}(t) \end{bmatrix}$$

### VAR: convenient forms

- Matrix form (hide ROIs)  $Y(t) = \alpha + A_1 Y(t-1) + ... + A_p Y(t-p) + \varepsilon(t)$
- Nice VAR(1) form (hide ROIs and lags): Z(t) = v + BZ(t-1) + u(t)

$$Z(t) = \begin{bmatrix} Y(t) \\ Y(t-1) \\ \vdots \\ Y(t-p+1) \end{bmatrix} \quad v = \begin{bmatrix} \alpha \\ 0 \\ \vdots \\ 0 \end{bmatrix} \quad B = \begin{bmatrix} A_1 & \cdots & A_{p-1} & A_p \\ I_n & \cdots & 0 & 0 \\ \vdots & \ddots & \vdots & \vdots \\ 0 & \cdots & I_n & 0 \end{bmatrix} \quad u(t) = \begin{bmatrix} \varepsilon(t) \\ 0 \\ \vdots \\ 0 \end{bmatrix}$$

• Even neater form (hide ROIs, lags and time): Y=BZ+U

$$Y = [Y(p+1), \dots, Y(T)], \qquad B = [\alpha, A_1, \dots, A_p], \qquad U = [\varepsilon(p+1), \dots, \varepsilon(T)],$$

$$Z = \begin{bmatrix} 1 & 1 & \dots & 1 \\ Y(p) & Y(p+1) & \dots & Y(T-1) \\ \vdots & \vdots & \vdots & \vdots \\ Y(1) & Y(2) & \dots & Y(T-p) \end{bmatrix}$$

■ Solve it with OLS:

$$\hat{B} = YZ^{+} = YZ^{t} (ZZ^{t})^{-1}$$

### VAR extended with covariates

- Standard VAR(p)  $Y(t) = \alpha + A_1Y(t-1) + ... + A_pY(t-p) + ε(t)$
- Covariates are all over the place!
  - □ Trend, tasks/conditions of no interest, head motion, time breaks (due to multiple runs), censored time points, physiological noises, etc.
- Extended VAR(*p*)

$$Y(t) = \alpha + A_1 Y(t-1) + \dots + A_p Y(t-p) + BZ_1(t) + \dots + B_q Z_q(t) + \mathbf{E}(t),$$
  
where  $Z_1, \dots, Z_q$  are covariates

- Endogenous (dependent: ROI time series)
- Exogenous (independent: covariates) variables
- □ Path strength significance: *t*-statistic (*F* in BrainVoyager)

# Model quality check

- Order selection: 4 criteria (1st two tend to overestimate)
  - AIC: Akaike Information Criterion
  - FPE: Final Prediction Error
  - □ HQ: Hannan-Quinn
  - □ SC: Schwartz Criterion
- Stationarity: VAR(p)  $Y(t) = \alpha + A_1 Y(t-1) + ... + A_p Y(t-p) + \varepsilon(t)$ 
  - □ Check characteristic polynomial  $\det(I_n A_1 z ... A_p z^p) \neq 0$  for  $|z| \leq 1$
- Residuals normality test
  - □ Gaussian process: Jarque-Bera test (dependent on variable order)
  - □ Skewness (symmetric or tilted?)
  - □ Kurtosis (leptokurtic or spread-out?)

## Model quality check (continued)

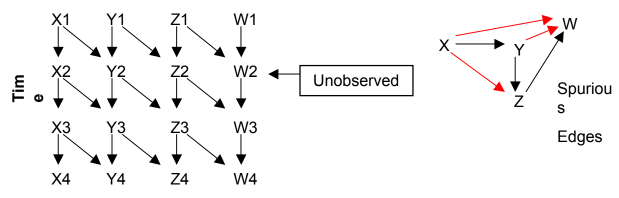
- Residual autocorrelation
  - Portmanteau test (asymptotic and adjusted)
  - □ Breusch-Godfrey LM test
  - Edgerton-Shukur *F* test
- Autoregressive conditional heteroskedasticity (ARCH)
  - □ Time-varying volatility
- Structural stability/stationarity detection
  - Is there any structural change in the data?
  - Based on residuals or path coefficients

# GC applied to FMRI

- Resting state
  - Ideal situation: no cut and paste involved
  - □ Physiological data maybe essential?
- Block experiments
  - □ Duration  $\geq$  5 seconds?
  - Extraction via cut and paste
    - Important especially when handling confounding effects
    - o Tricky: where to cut especially when blocks not well-separated?
- Event-related design
  - With rapid event-related, might not need to cut and paste (at least impractical)
  - Other tasks/conditions as confounding effects

### GC: caveats

- Assumptions (stationarity, linearity, Gaussian residuals, no serial correlations in residuals, etc.)
- Accurate ROI selection
- Sensitive to lags
- o Interpretation of path coefficient: slope, like classical regression
- Confounding latency due to vascular effects
- No transitive relationship: If  $Y_3(t)$  Granger causes  $Y_2(t)$ , and  $Y_2(t)$  Granger causes  $Y_1(t)$ , it does not necessarily follow that  $Y_3(t)$  Granger causes  $Y_1(t)$ .
- Time resolution? Not so serious a problem? Not neuronal signal, but blurred through IRF
   Regions



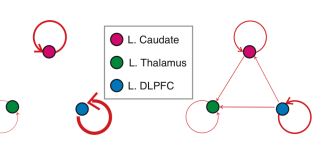
### GC in AFNI

- Exploratory: ROI searching with 3dGC
  - □ Seed vs. rest of brain
  - Bivariate model
  - □ 3 paths: seed to target, target to seed, and self-effect
  - □ Group analysis with **3dMEMA** or **3dttest**
- Path strength significance testing in network: 1dGC
  - Pre-selected ROIs
  - Multivariate model
  - Multiple comparisons issue
  - Group analysis
    - path coefficients only
    - path coefficients + standard error
    - F-statistic (BrainVoyager)

## GC: hands-on

Exploratory: ROI searching with 3dGC

- Seed: sACC
- Sequential and batch mode (~5 minutes)
- Data courtesy: Paul Hamilton (Stanford)
- Path strength significance testing in network: 1dGC
  - Data courtesy: Paul Hamilton (Stanford)
  - Individual subject
    - 3 pre-selected ROIs: left caudate, left thalamus, left DLPFC
    - 8 covariates: 6 head motion parameters, 2 physiological datasets
  - Group analysis
    - path coefficients only
    - path coefficients + standard errors

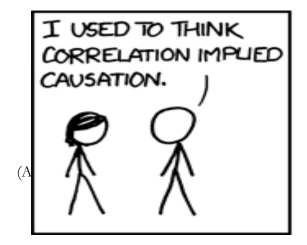


## Summary: connectivity analysis

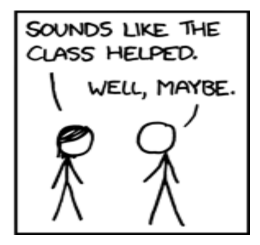
- 2 basic categories
  - Seed-based method for ROI searching
  - Network-based for network validation
- 3 approaches
  - Correlation analysis
  - Structural equal modeling
  - Granger causality
- A lot of interpretation traps
  - Over-interpretation seems everywhere
  - I may have sounded too negative about connectivity analysis
- Causality regarding the class: Has it helped you somehow?
  - □ Well, maybe?

### Interpretation Trap: Correlation vs. Causation!

- Some analyses require fine time resolution we usually lack
- Path from (or correlation btw) A to (and) B doesn't necessarily mean causation
  - □ Bi-regional approach simply ignores the possibility of other regions involved
  - Analysis invalid if a relevant region is missing in a multi-regional model
- Robust: connectivity analysis < regression analysis</li>
- Determinism in academics and in life
  - □ Linguistic determinism: Sapir-Whorf hypothesis







# Other approaches

- Multivariate (data-driven)
  - □ Techniques from machine learning, pattern recognition
  - □ Training + prediction
  - PCA/ICA
  - □ SVM: 3dsvm, plug-in
  - Kernel methods