

Introduction to AFNI-FATCAT

Tractography for data exploration and complementing functional connectivity

Paul A. Taylor^{1,2} & Ziad S. Saad³

¹Medical Imaging Research Unit, University of Cape Town, South Africa

²African Institute for Mathematical Sciences, Muizenberg, Western Cape, South Africa

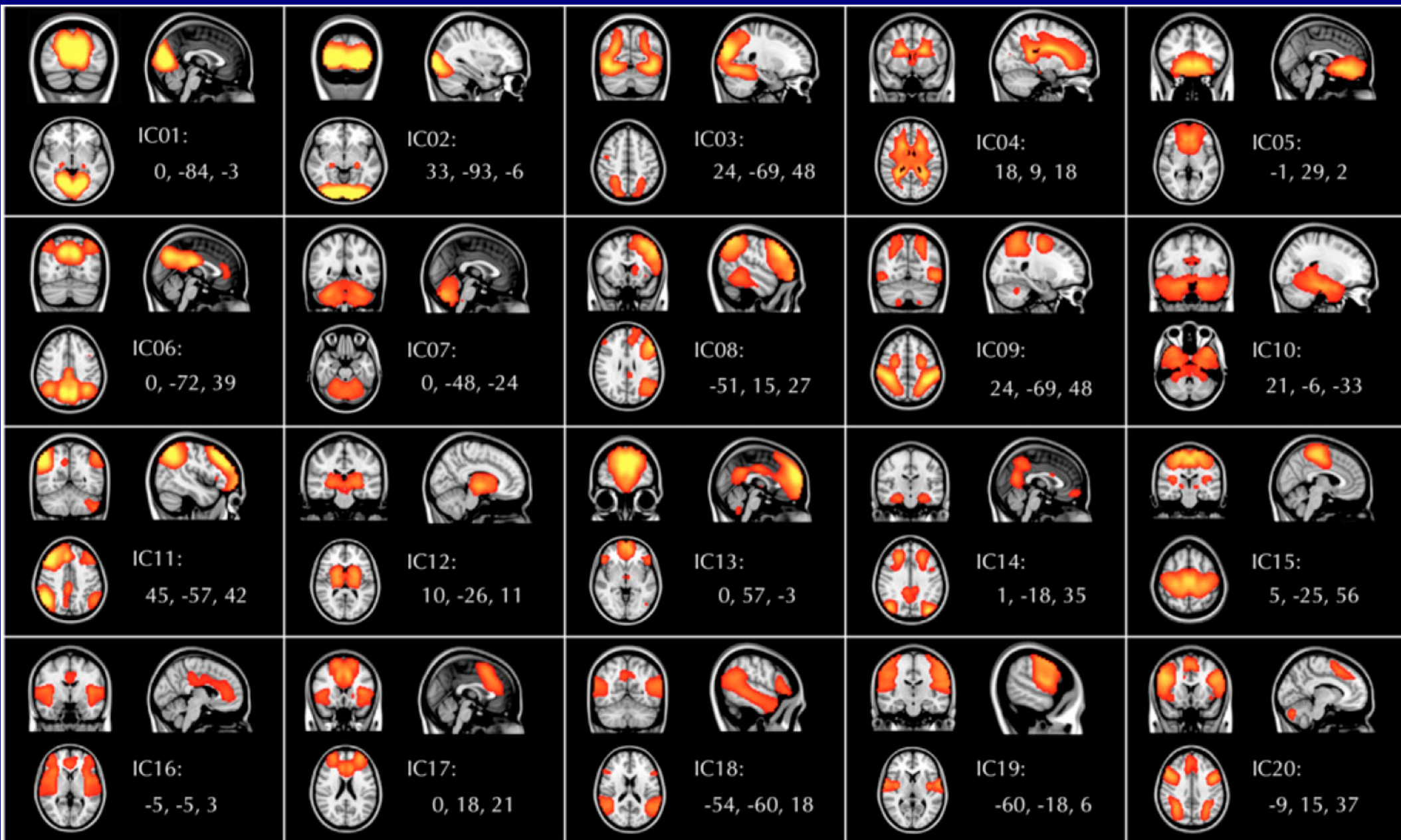
³Scientific and Statistical Computing Core, NIH, Bethesda, MD, USA



Outline

- + Why Function+Structure
- + DWI and DTI (very brief, following morning session)
 - Diffusion imaging basics and parameters
- + Using tractography to estimate WM connections
 - Making targets from functional data
 - Deterministic, probabilistic (or both?)
 - using WM region properties for quantitative comparison
- + Brief example – newborn alcohol exposure study
- + Further FATCAT applications:
 - HARDI tracking, Connectome studies

FMRI: GM Networks



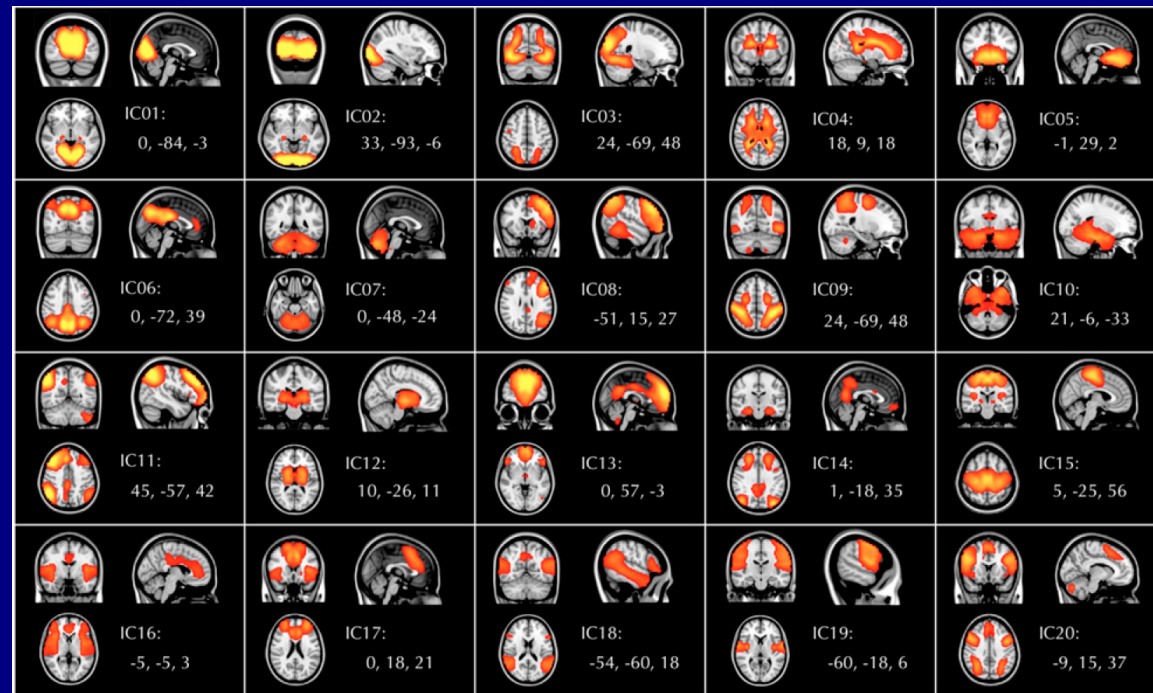
(Biswal et al., 2010 PNAS)

FMRI: GM Networks

Functional connectivity networks of distinct GM regions, from BOLD time series during task or rest/no task.

+ Quantify GM properties: ALFF, fALFF, RSFA, σ , ReHo, GMV, etc.

+ Quantify network props: seedbased correlation, ICA, graph theoretical measures, etc.

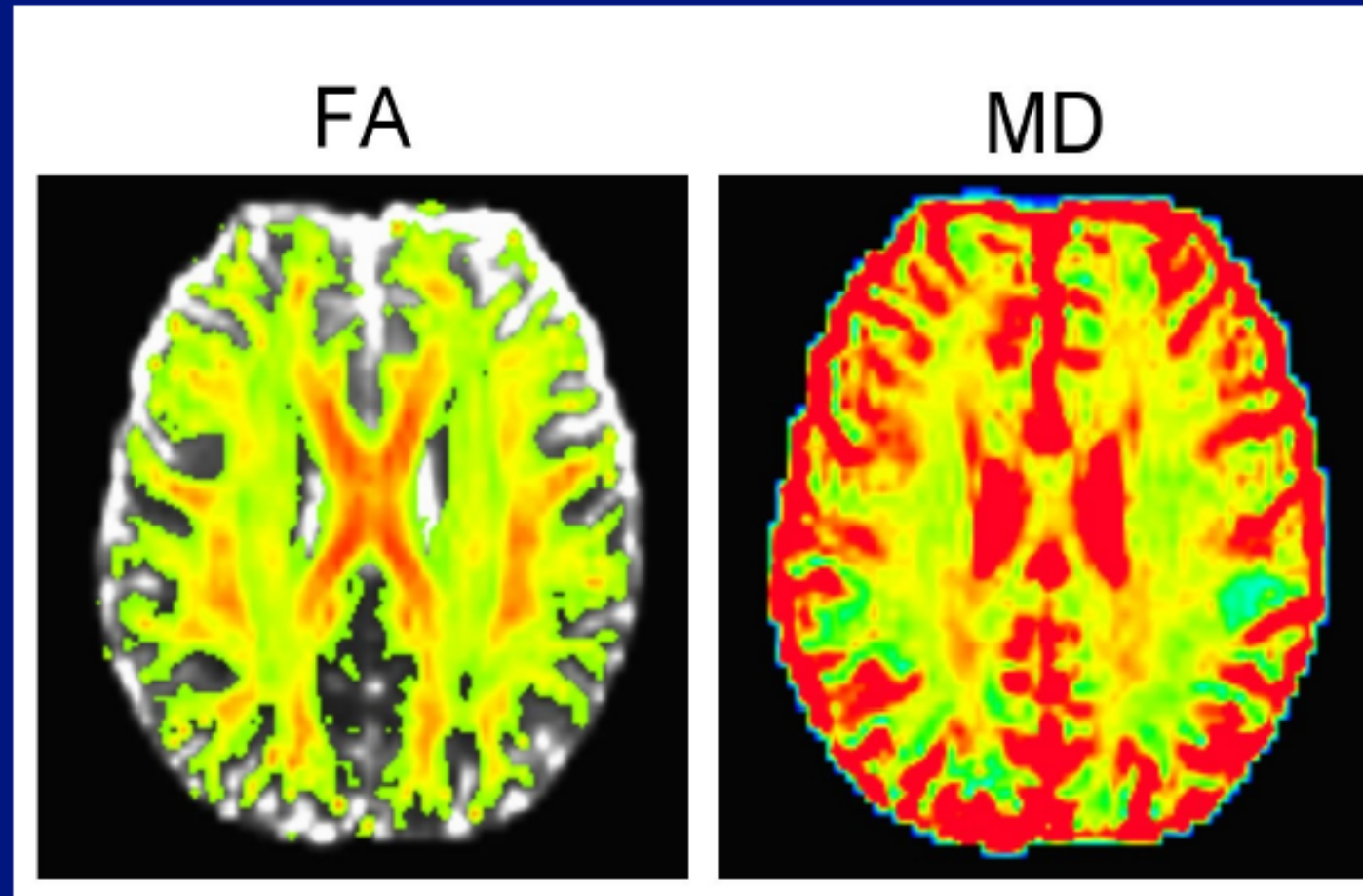


Structural (WM)

DTI-based parameters characterize some local properties, and also show presence of spatially-extended WM structures

Can investigate and quantify WM properties with:
FA, MD, RD, L1, etc.

Can investigate (and quantify?) network relations with:
tractography

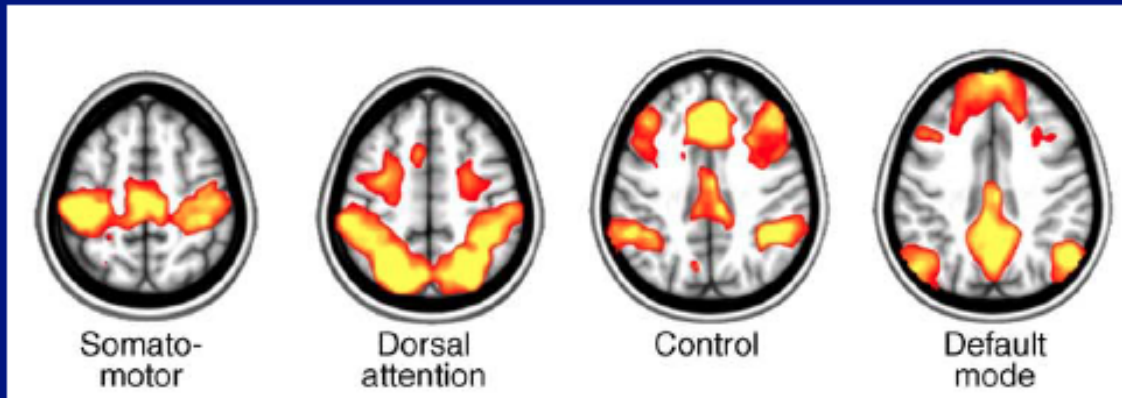


(FA>0.2)

Structure + Function

Simple example:

GM ROIs
network:

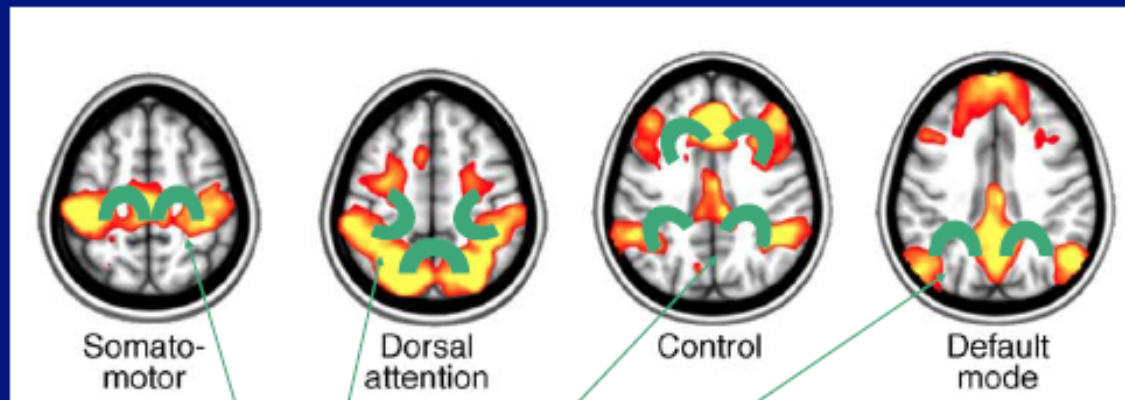


Raichle (2010, TICS)

Structure + Function

Simple example:

GM ROIs
network:



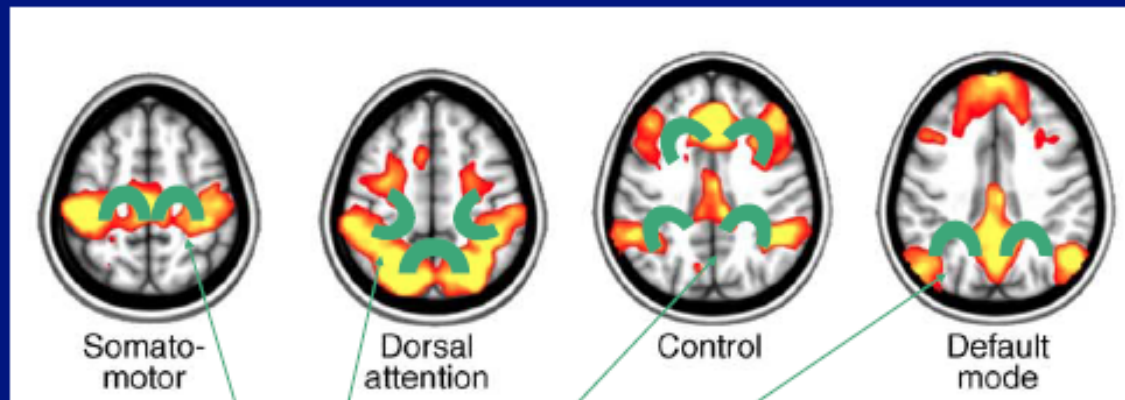
Raichle (2010, TICS)

Associated WM ROIs

Structure + Function

Simple example:

GM ROIs
network:



Raichle (2010, TICS)

Associated WM ROIs

Our goal for tractography->

*estimate likely/probable locations of WM associated with GM,
and relate ROI quantities with functional/GM properties*

Combining FC and SC

- + How to combine *quantitatively*?
 - fMRI has measures of functional **connectivity** and **'strength'** (e.g., correlation, network parameters)

Combining FC and SC

- + How to combine *quantitatively*?
 - fMRI has measures of functional connectivity and 'strength' (e.g., correlation, network parameters)

- DTI tracking between GM ROIs-- we can have 'structural connectivity' strength, e.g., in terms of # of fibers?
 - > will discuss more, but think this is *not* good road to be on

Combining FC and SC

- + How to combine *quantitatively*?
 - fMRI has measures of functional connectivity and 'strength' (e.g., correlation, network parameters)
- DTI tracking between GM ROIs-- we can have 'structural connectivity' strength, e.g., in terms of # of fibers?
 - > will discuss more, but think this is *not* good road to be on
 - how about:
 - find **likely areas** where WM is connecting GM regions, and **quantify properties** in those regions (FA, MD, proton density from structural images...)

Combining FC and SC

- + How to combine *quantitatively*?
 - FMRI has measures of functional connectivity and 'strength' (e.g., correlation, network parameters)
- DTI tracking between GM ROIs-- we can have 'structural connectivity' strength, e.g., in terms of # of fibers?
 - > will discuss more, but think this is *not* good road to be on
 - how about:
 - find likely areas where WM is connecting GM regions, and quantify properties in those regions (FA, MD, proton density from structural images...)

→ FC+SC provides sets of complementary quantities to describe a network, and can be further combined with behavioral/other measures (statistical modeling).

Tools for combining FC and SC:

Combining functional and tractographic connectivity will require:

- + determining networks from fMRI data;
- + finding correlations and local properties of functional networks;
- + turning GM ROIs into targets for tractography;
- + doing reasonable tractography to find WM ROIs;
- + estimating stats on WM ROIs...

Tools for combining FC and SC:

Combining functional and tractographic connectivity will require:

- + determining networks from fMRI data;
- + finding correlations and local properties of functional networks;
- + turning GM ROIs into targets for tractography;
- + doing reasonable tractography to find WM ROIs;
- + estimating stats on WM ROIs...

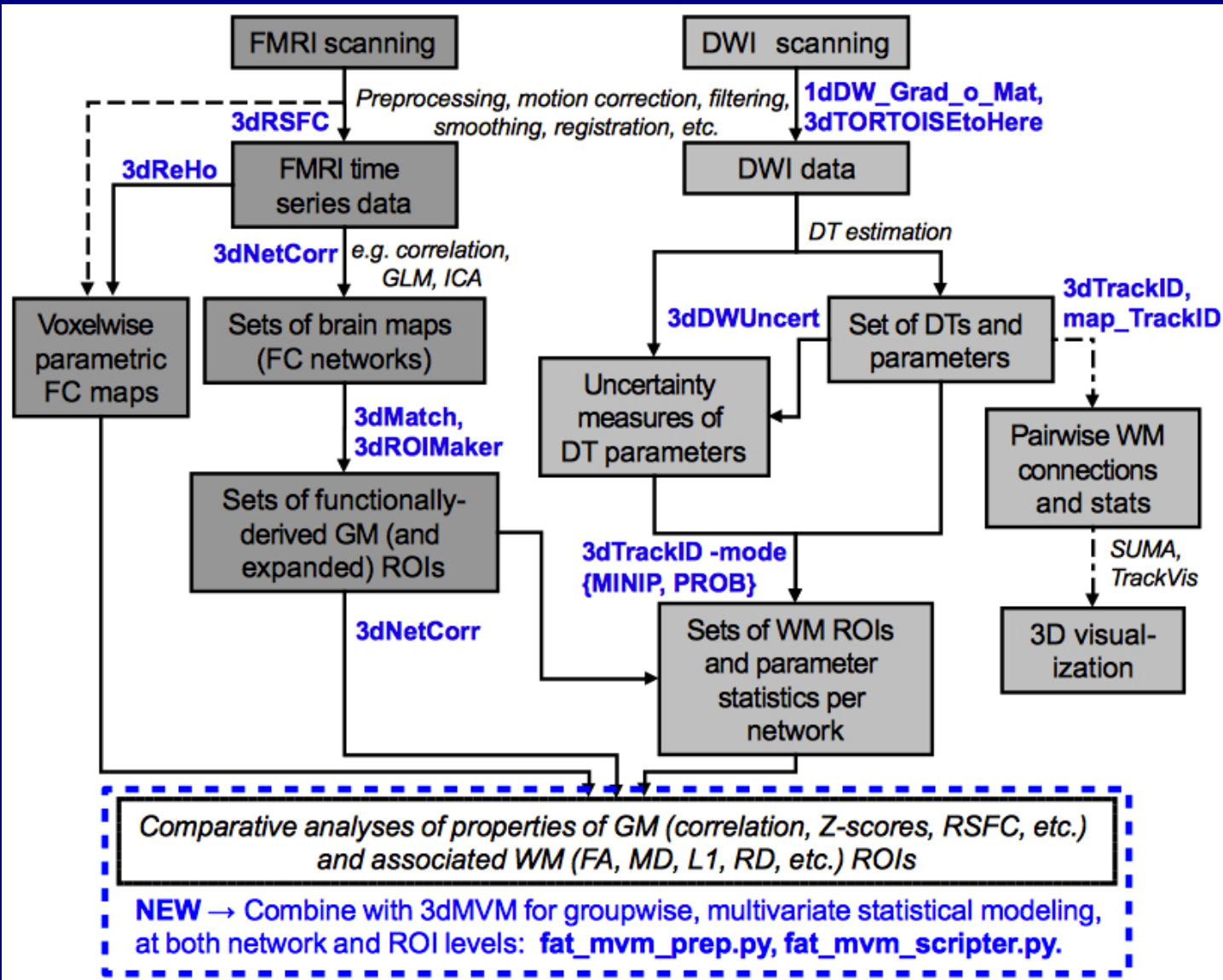
FATCAT: Functional And Tractographic Connectivity Analysis Toolbox (Taylor & Saad, 2013), now available in AFNI with demo data.



*picture from google search, not from/of either author

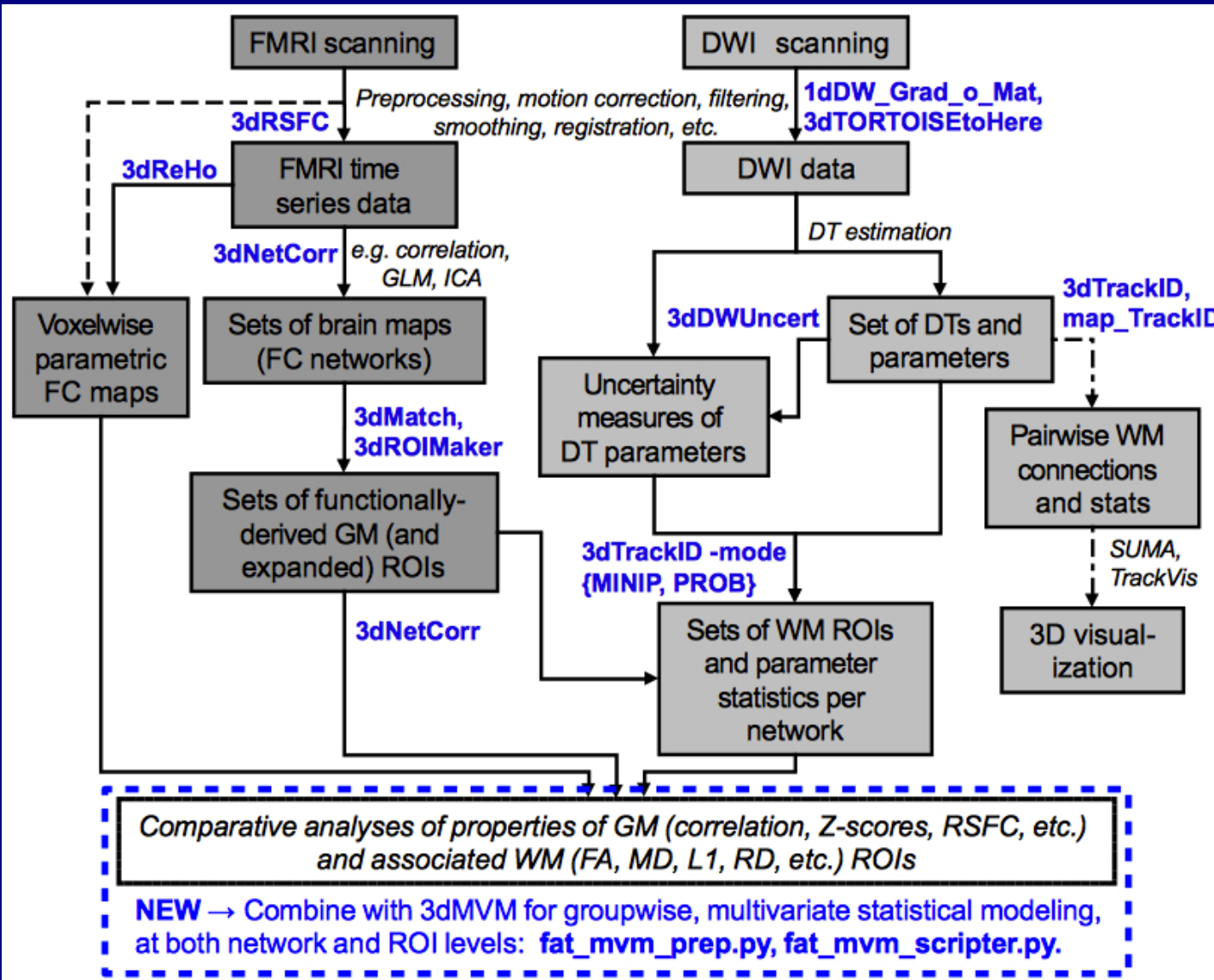
Functional and structural processing

Schematic for combining FMRI and DTI-tractography via FATCAT:



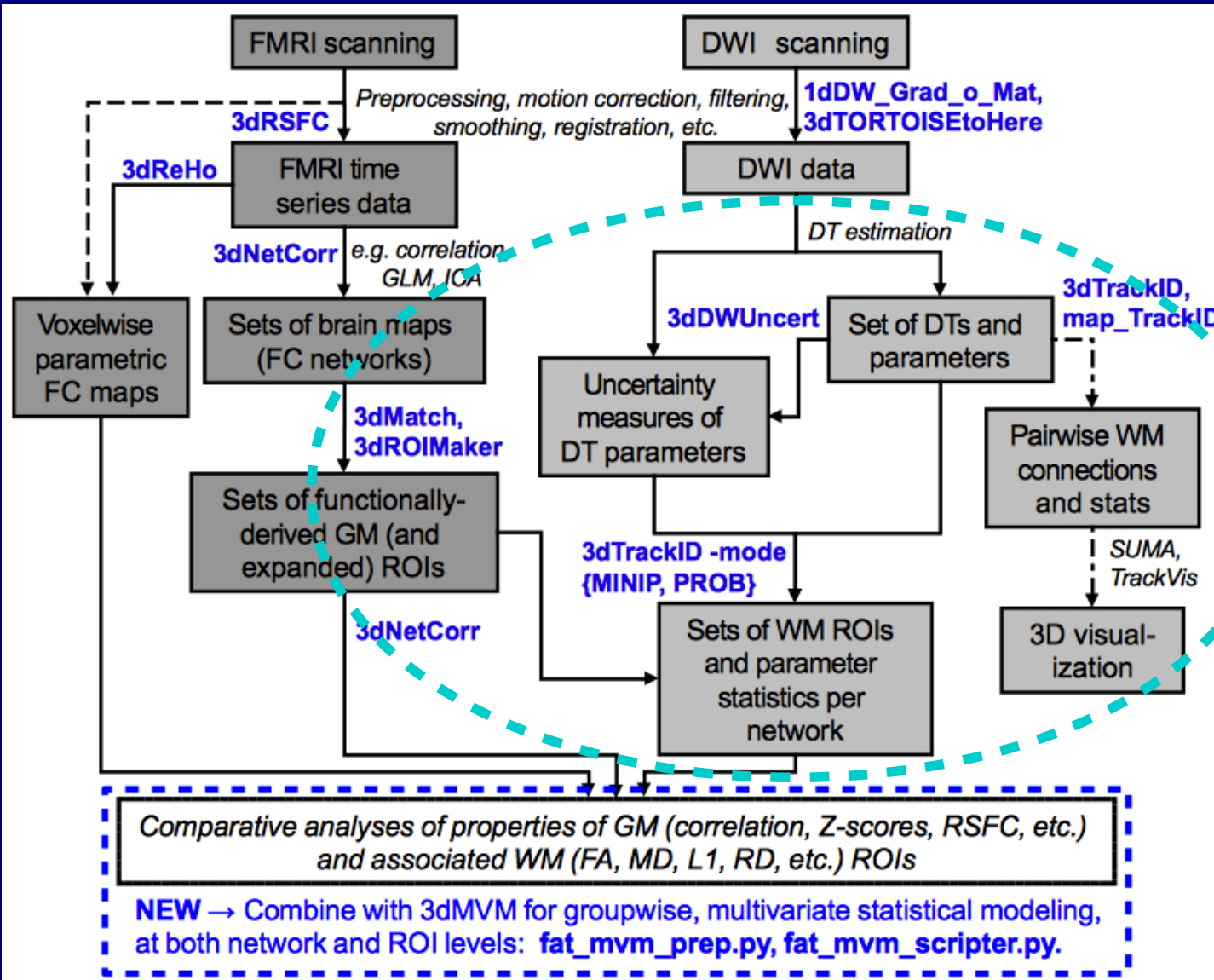
Functional and structural processing

Schematic for combining FMRI and DTI-tractography via FATCAT:



Functional and structural processing

Schematic for combining fMRI and DTI-tractography via FATCAT:



FATCAT goals:

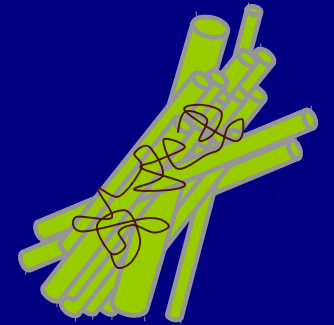
- + do useful tasks
- + integrate with existing pipelines/software
- + derive/use information from the data itself
- + be simple to implement
- + be efficient
- + be flexible and able to grow

Main focus today on DTI-tractography, including making ROIs from fMRI

Local Structure via Diffusion MRI

(In brief)

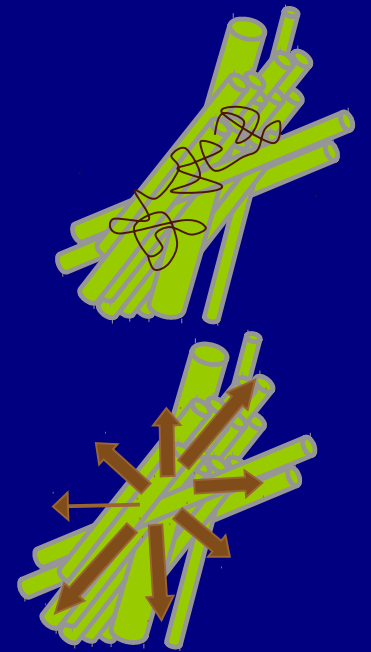
1) Random motion of molecules affected by local structures



Local Structure via Diffusion MRI

(In brief)

- 1) Random motion of molecules affected by local structures
- 2) Statistical motion measured using diffusion weighted MRI



Local Structure via Diffusion MRI

(In brief)

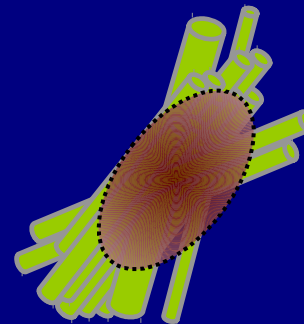
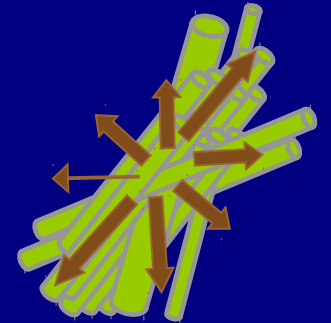
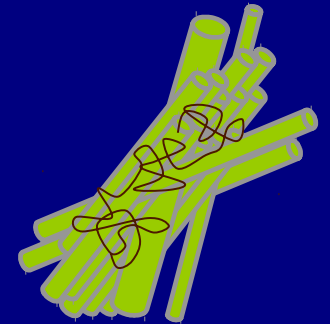
1) Random motion of molecules affected by local structures

2) Statistical motion measured using diffusion weighted MRI

3) Bulk features of local structure approximated with various reconstruction models, mainly grouped by number of major structure directions/voxel:

+ one direction:

DTI (Diffusion Tensor Imaging)



Local Structure via Diffusion MRI

(In brief)

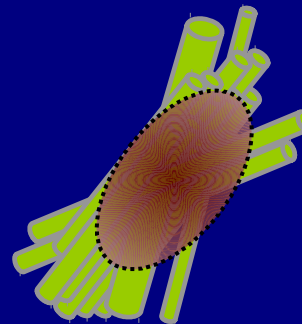
1) Random motion of molecules affected by local structures

2) Statistical motion measured using diffusion weighted MRI

3) Bulk features of local structure approximated with various reconstruction models, mainly grouped by number of major structure directions/voxel:

+ one direction:

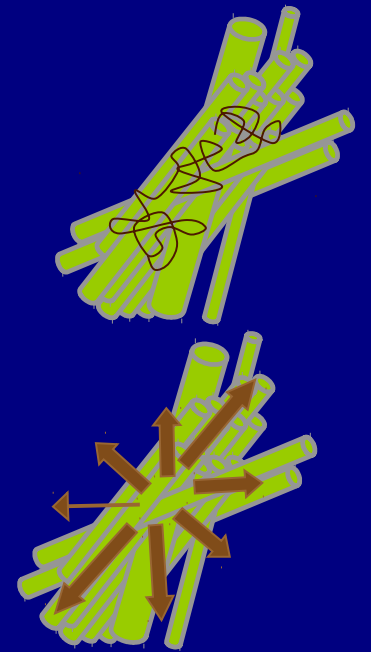
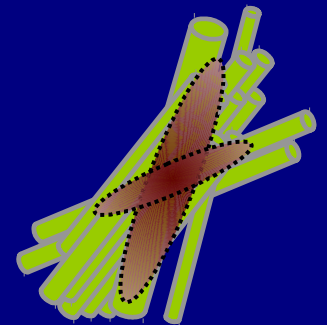
DTI (Diffusion Tensor Imaging)



+ ≥ 1 direction:

HARDI (High Angular Resolution Diffusion Imaging)

Qball, DSI, ODFs, ball-and-stick, multi-tensor, CSD, ...



DWI → Diffusion Tensors (DTs)

Mathematically, the properties of the diffusion tensor:

$$\mathbf{D} = \begin{pmatrix} D_{11} & D_{12} & D_{13} \\ D_{21} & D_{22} & D_{23} \\ D_{31} & D_{32} & D_{33} \end{pmatrix}$$

Having: 3 eigenvectors: \mathbf{e}_i
3 eigenvalues: λ_i

- Real-valued
- Positive definite ($\mathbf{r}^T \mathbf{D} \mathbf{r} > 0$)
 $\mathbf{D} \mathbf{e}_i = \lambda_i \mathbf{e}_i, \quad \lambda_i > 0$
- Symmetric ($D_{12} = D_{21}$, etc),
6 independent values

DWI → Diffusion Tensors (DTs)

Mathematically, the properties of the diffusion tensor:

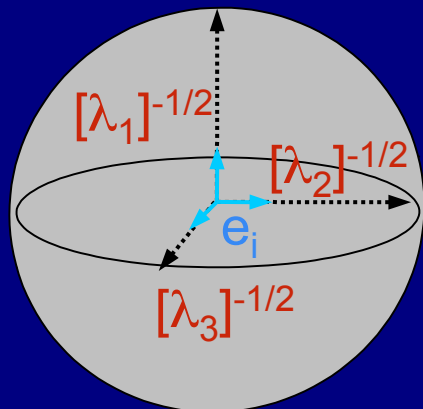
$$\mathbf{D} = \begin{pmatrix} D_{11} & D_{12} & D_{13} \\ D_{21} & D_{22} & D_{23} \\ D_{31} & D_{32} & D_{33} \end{pmatrix}$$

Having: 3 eigenvectors: \mathbf{e}_i
3 eigenvalues: λ_i

- Real-valued
- Positive definite ($\mathbf{r}^T \mathbf{D} \mathbf{r} > 0$)
 $\mathbf{D} \mathbf{e}_i = \lambda_i \mathbf{e}_i, \quad \lambda_i > 0$
- Symmetric ($D_{12} = D_{21}$, etc),
6 independent values

Geometrically, this describes ellipsoid surface, with $\mathbf{r} = (x, y, z)$:

$$C = \mathbf{r}^T \mathbf{D} \mathbf{r} = D_{11}x^2 + D_{22}y^2 + D_{33}z^2 + 2(D_{12}xy + D_{13}xz + D_{23}yz)$$



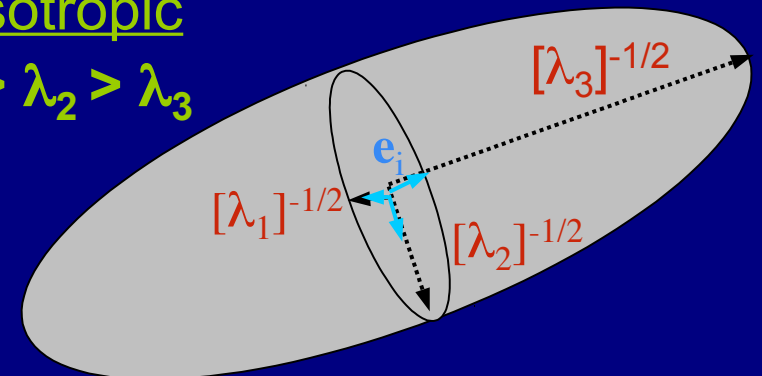
isotropic

$$\lambda_1 = \lambda_2 = \lambda_3$$

anisotropic

$$\lambda_1 > \lambda_2 > \lambda_3$$

**'Diffusion measure'
surfaces**



λ_i describe length of semiaxes; \mathbf{e}_i are spatial orientation of semiaxes

DWI → Diffusion Tensors (DTs)

diffusion tensor

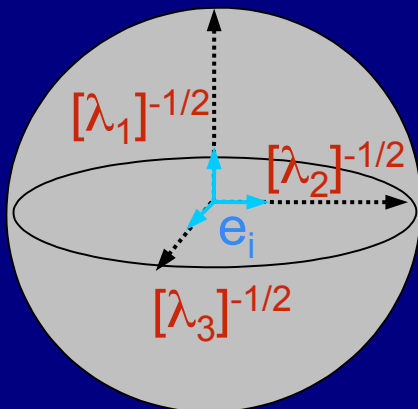
Orientation and magnitude

Having: 3 eigenvectors: e_i
3 eigenvalues: λ_i

Minimum number of measures

6 independent values

ellipsoid surface



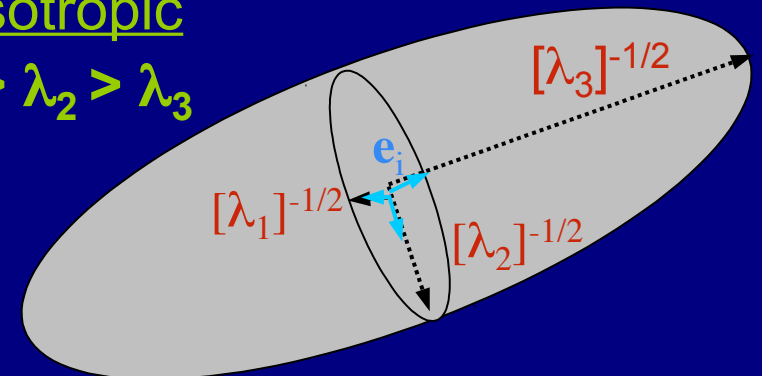
isotropic

$$\lambda_1 = \lambda_2 = \lambda_3$$

anisotropic

$$\lambda_1 > \lambda_2 > \lambda_3$$

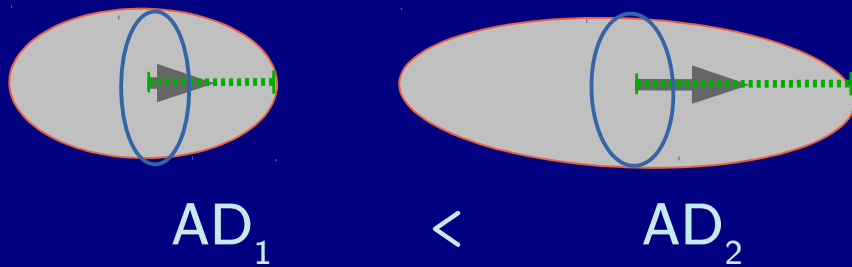
'Diffusion measure'
surfaces



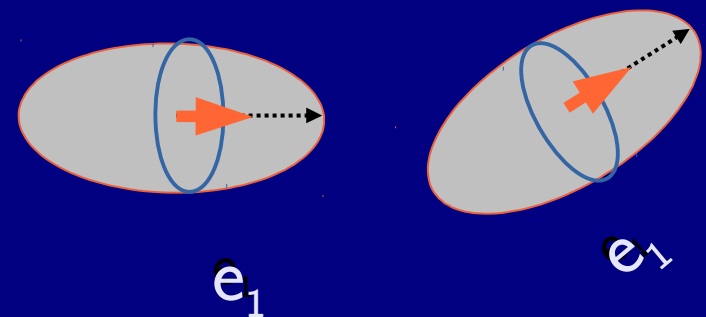
“Big 5” DTI ellipsoid parameters

Main quantities of diffusion (motion) surface

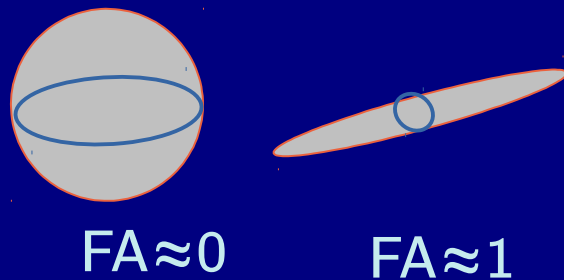
first eigenvalue, L_1
 $\equiv \lambda_1$, parallel/axial diffusivity, AD



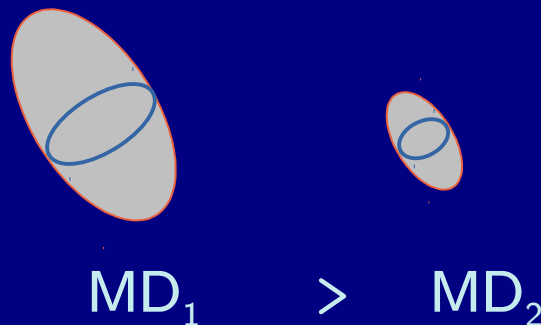
first eigenvector, e_1



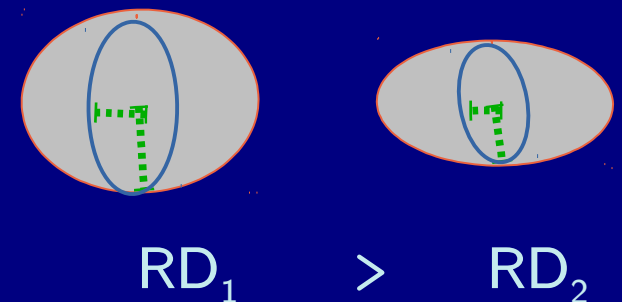
Fractional anisotropy, FA



Mean diffusivity, MD



Radial diffusivity, RD



Interpreting DTI parameters

General literature:

FA: measure of fiber bundle coherence and myelination

- in adults, $FA > 0.2$ is proxy for WM (strong segment. overlap)

MD, RD, L1: local density of structure

e_1 : orientation of major bundles

Interpreting DTI parameters

General literature:

FA: measure of fiber bundle coherence and myelination

- in adults, $FA > 0.2$ is proxy for WM (strong segment. overlap)

MD, RD, L1: local density of structure

e_1 : orientation of major bundles

Cautionary notes:

+ Degeneracies of structural interpretations

+ Changes in myelination may have small effects on FA

+ WM bundle diameter \ll voxel size

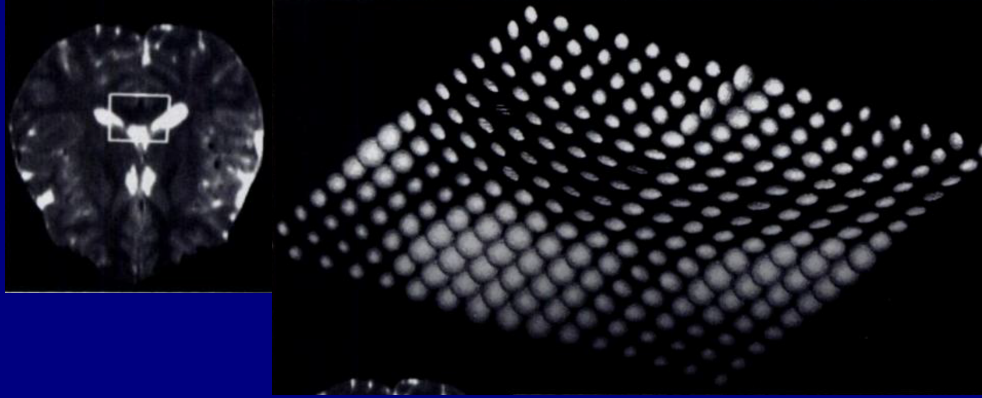
- don't know location/multiplicity of underlying structures

+ More to diffusion than just structure-- i.e., fluid properties

+ Noise, distortions, etc. in measures

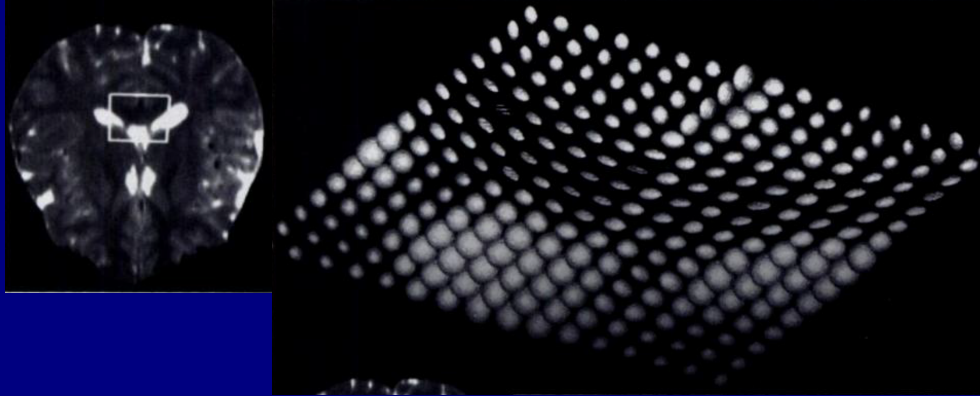
Local DTs \rightarrow Extended Tracts

Field of local diffusion parameters



Local DTs → Extended Tracts

Field of local diffusion parameters

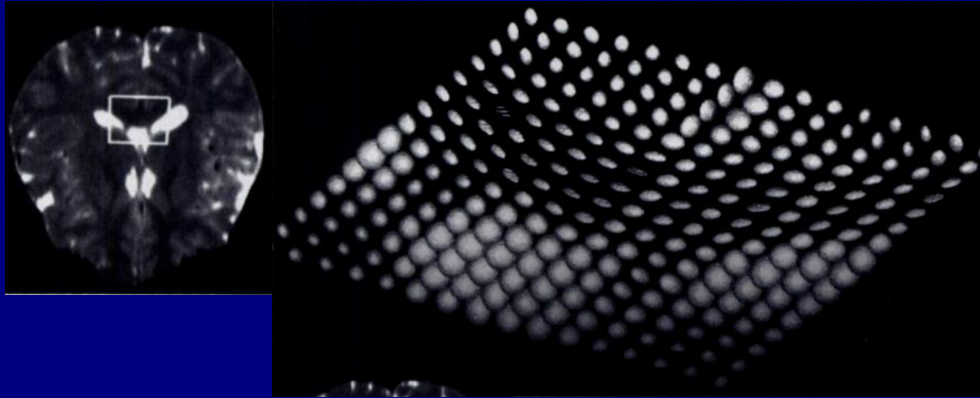


→ individual ellipsoids

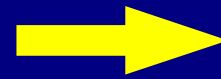
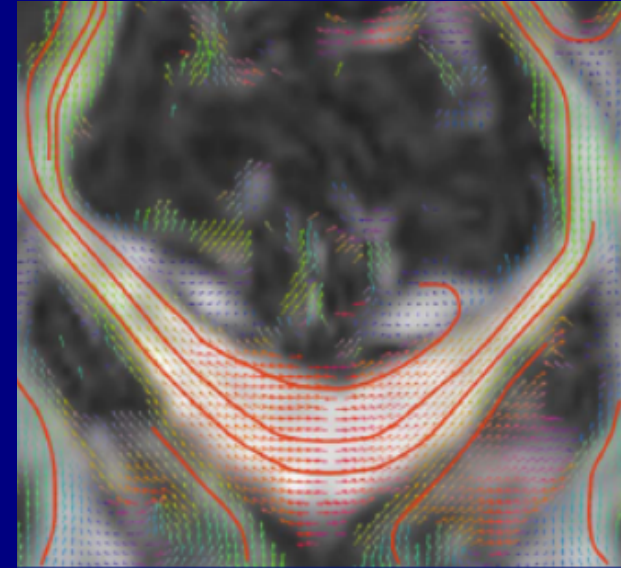


Local DTs → Extended Tracts

Field of local diffusion parameters



Connect to form extended tracts



→ individual ellipsoids

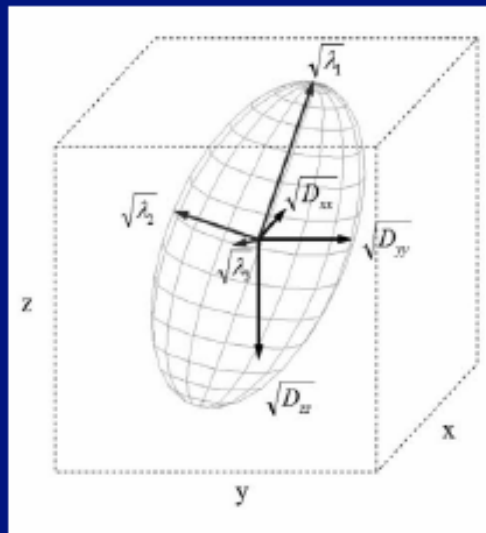


→ linked structures

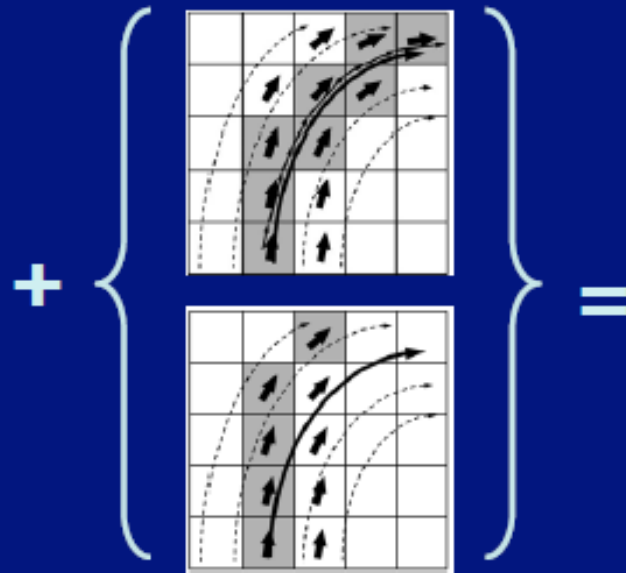


Tractography

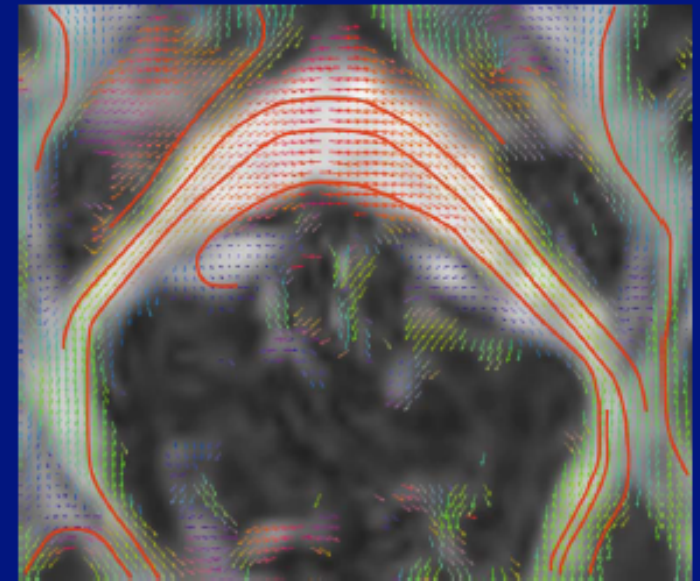
Estimate WM structure (fiber tract locations)



ellipsoid measures
(~smoothing of
real structures)



some kind of algorithm
for connecting



estimate spatial
extents of WM 'tracts'
in vivo

Diversity in tractography

Series of (mostly) logical, simple rules for estimating tracts

→ many methods/algorithms and kinds of parameters to choose:
(Mori et al., 1999; Conturo et al. 1999; Weinstein et al. 1999;
Basser et al. 2000; Poupon et al. 2001; Mangin et al. 2002;
Lazar et al. 2003;)

Propagation via, e.g.:

smoothing diffusion vectors and solving differential equations;
deflecting propagating tracts; allowing tracts themselves to
'diffuse'; solving for global minimum energy of connections...

To date, no single 'best' algorithm, work continues:

- histology can't give perfect answers.
- some test models (phantoms) exist, but not brain-complex

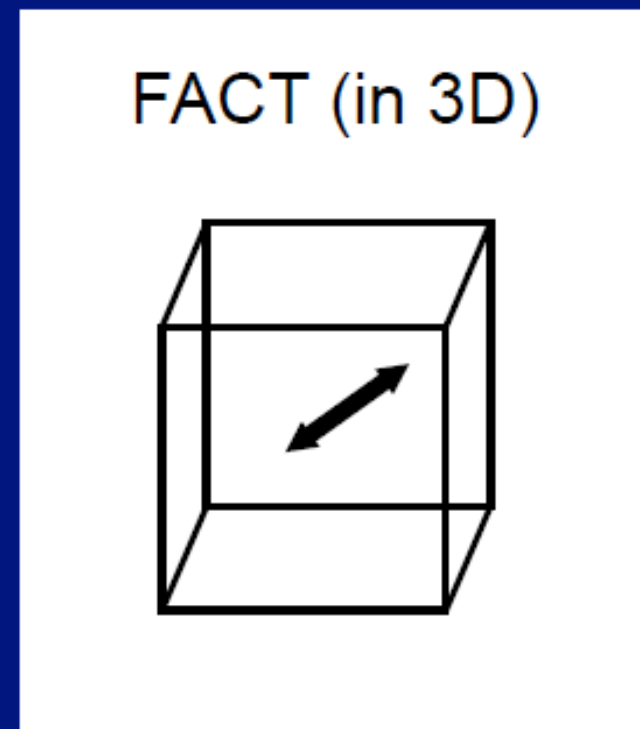
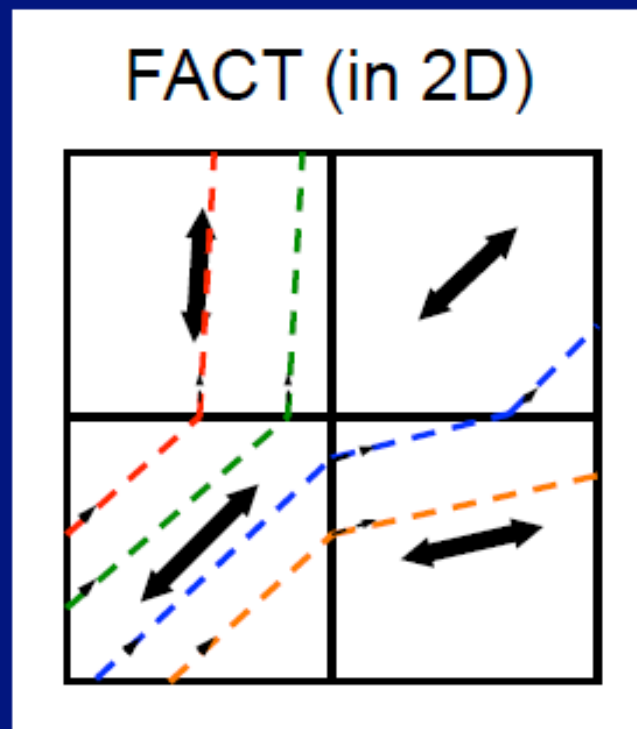
So, first question for using tractography in a study:

Which algorithm to choose?

Popular technique: FACT

- FACT = Fiber Assessment by Continuous Tracking (Mori et al. 1999) [used more than 200 times in past 1.5 yrs]
 - Start in voxel with $FA > 0.2$ (proxy definition for WM)
 - Follow 1st eigenvector/greatest diffusion direction to next voxel
 - Continue if $FA > 0.2$ and angle between e_1 s is < 45 deg

Ex.:



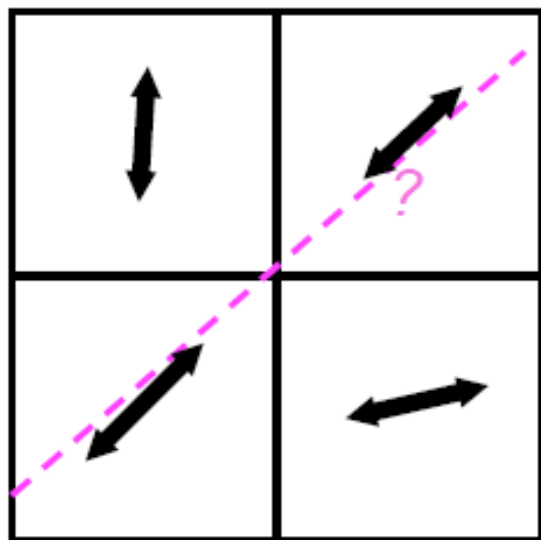
Very simple, but actually, gives some decent results, e.g. many known tracts

Popular technique: FACT

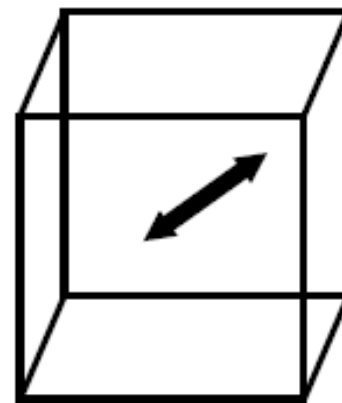
- FACT = Fiber Assessment by Continuous Tracking (Mori et al. 1999) [used more than 200 times in past 1.5 yrs]
 - Start in voxel with $FA > 0.2$ (proxy definition for WM)
 - Follow 1st eigenvector/greatest diffusion direction to next voxel
 - Continue if $FA > 0.2$ and angle between e_1 s is < 45 deg

Ex.:

FACT (in 2D)



FACT (in 3D)

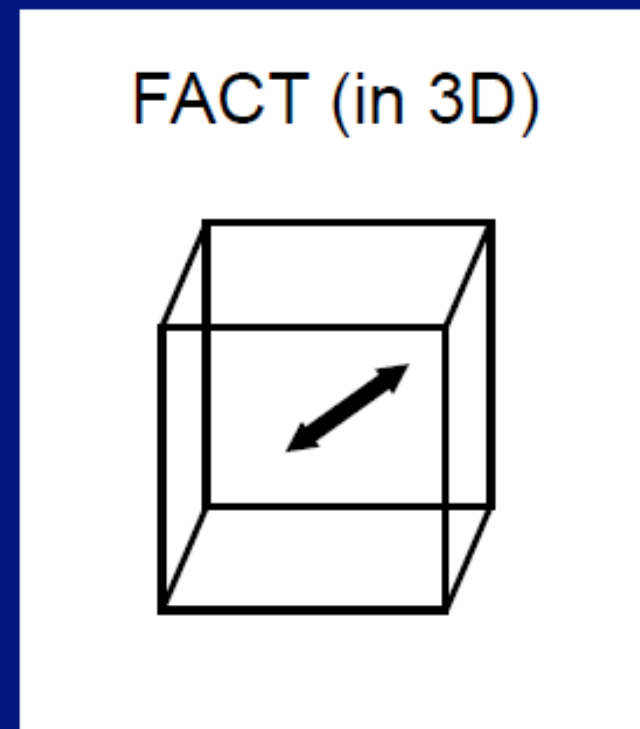
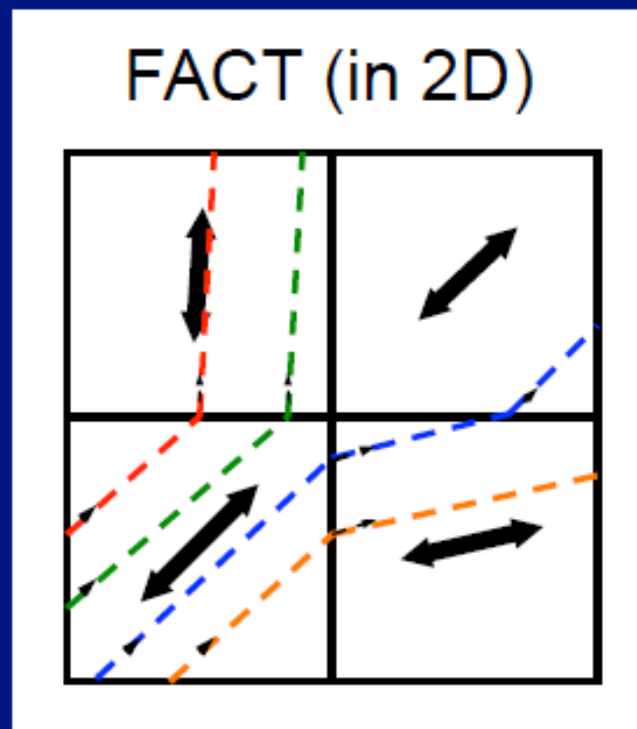


Very simple, but actually, gives some decent results, e.g. many known tracts **however... e.g. bias?*

Popular technique: FACT

- FACT = Fiber Assessment by Continuous Tracking (Mori et al. 1999) [used more than 200 times in past 1.5 yrs]
 - Start in voxel with $FA > 0.2$ (proxy definition for WM)
 - Follow 1st eigenvector/greatest diffusion direction to next voxel
 - Continue if $FA > 0.2$ and angle between e_1 s is < 45 deg

Ex.:

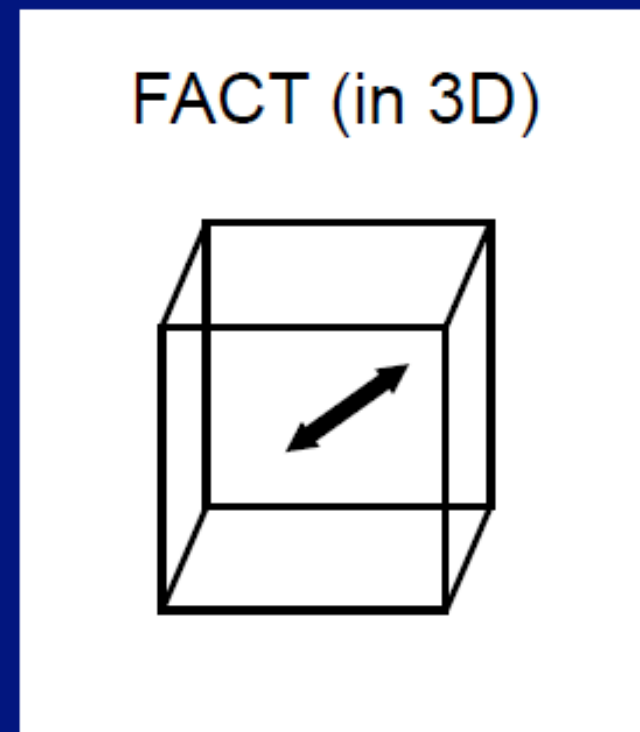
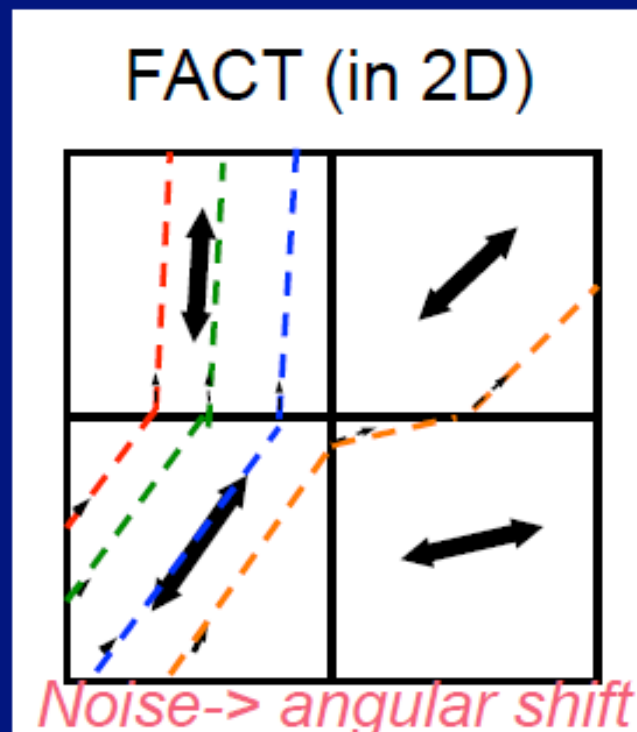


Very simple, but actually, gives some decent results, e.g. many known tracts **however... e.g. bias? noise dependence?*

Popular technique: FACT

- FACT = Fiber Assessment by Continuous Tracking (Mori et al. 1999) [used more than 200 times in past 1.5 yrs]
 - Start in voxel with $FA > 0.2$ (proxy definition for WM)
 - Follow 1st eigenvector/greatest diffusion direction to next voxel
 - Continue if $FA > 0.2$ and angle between e_1 s is < 45 deg

Ex.:



Very simple, but actually, gives some decent results, e.g. many known tracts **however... e.g. bias? noise dependence?*

Improving FACT->

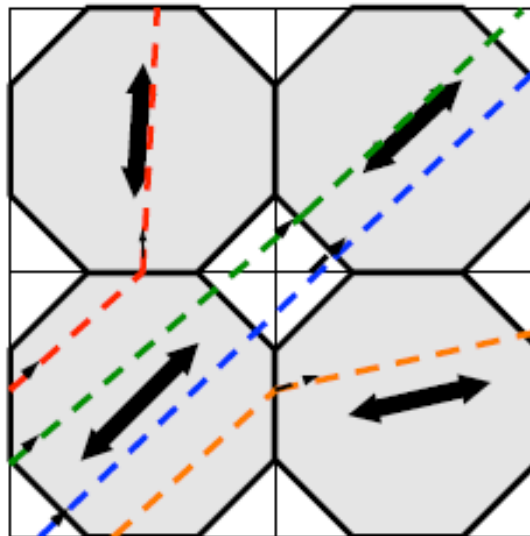
- Start by thinking: what properties a 'good' algorithm should have?
 - 1) Should be independent of coordinate axes (i.e., results invariant to rotation of data set)
 - 2) Should improve with spatial resolution (convergence in resolution)
e.g., like in calculus, diagonals are better approximated with small grid steps
 - 3) Should improve with SNR (converge in SNR)
 - 4) Should not have strong instability with or dependence on noise

Improving FACT->

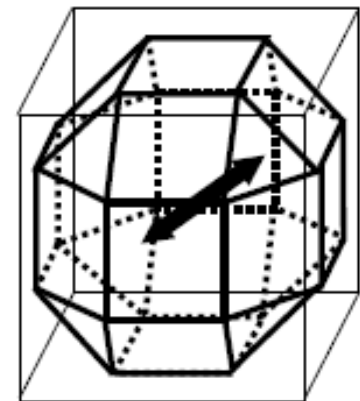
- Start by thinking: what properties a 'good' algorithm should have?
 - 1) Should be independent of coordinate axes (i.e., results invariant to rotation of data set)
 - 2) Should improve with spatial resolution (convergence in resolution)
e.g., like in calculus, diagonals are better approximated with small grid steps
 - 3) Should improve with SNR (converge in SNR)
 - 4) Should not have strong instability with or dependence on noise

Posit: including diagonal (ID) propagation helps 1 and 4, check about other props.

FACTID (in 2D)



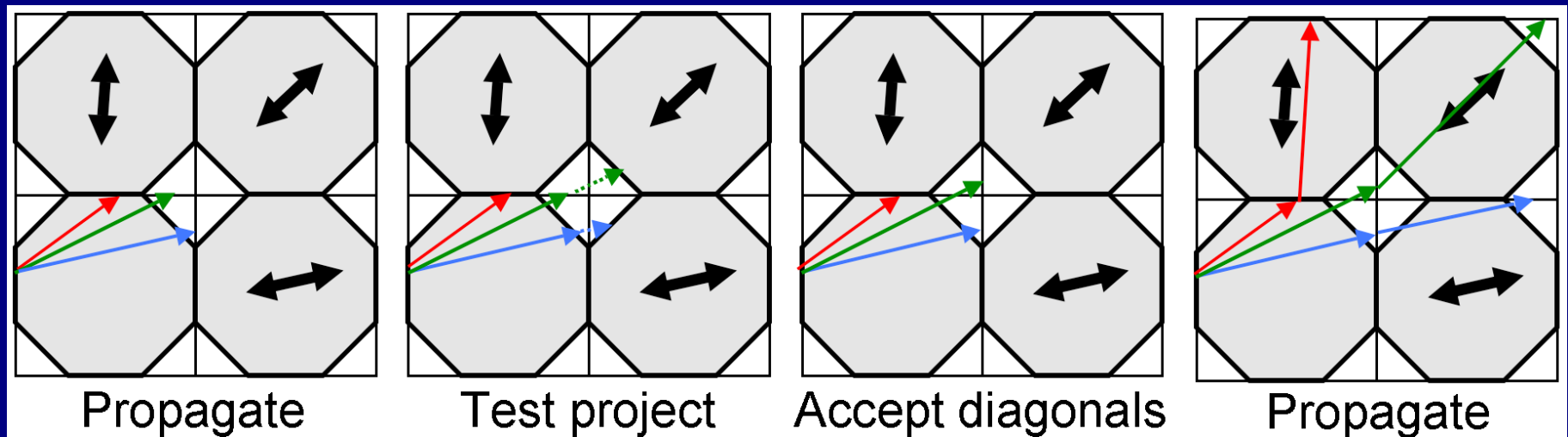
FACTID (in 3D)



FACTID (FACT Including Diagonals):

+ Utilize simple check for diagonals.

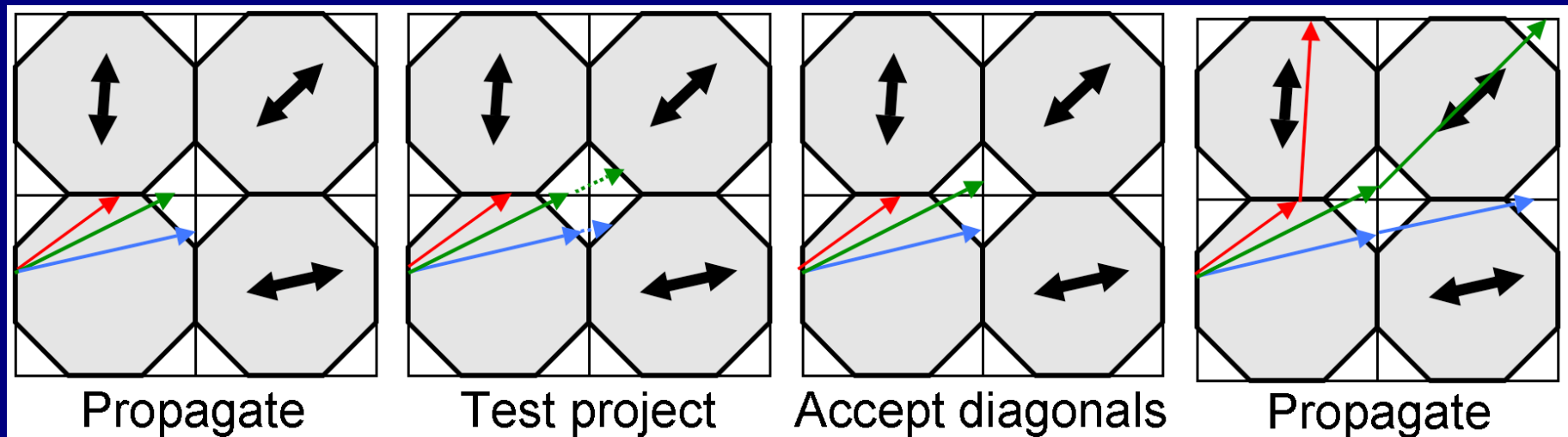
(2D) Schematic:



FACTID (FACT Including Diagonals):

+ Utilize simple check for diagonals.

(2D) Schematic:

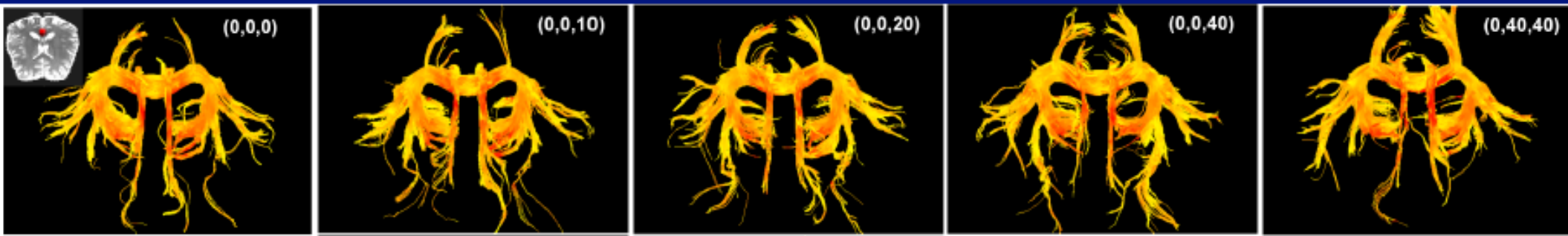


NB that in (3D) FACT, a single voxel has 6 neighbors for propagation, while in FACTID, a voxel has 26 neighbors propagation.

Test 1: Rotational invariance

A test for consistency of results when axes of data have been rotated; here, using data from a real subject (scan axes rotated)

FACTID

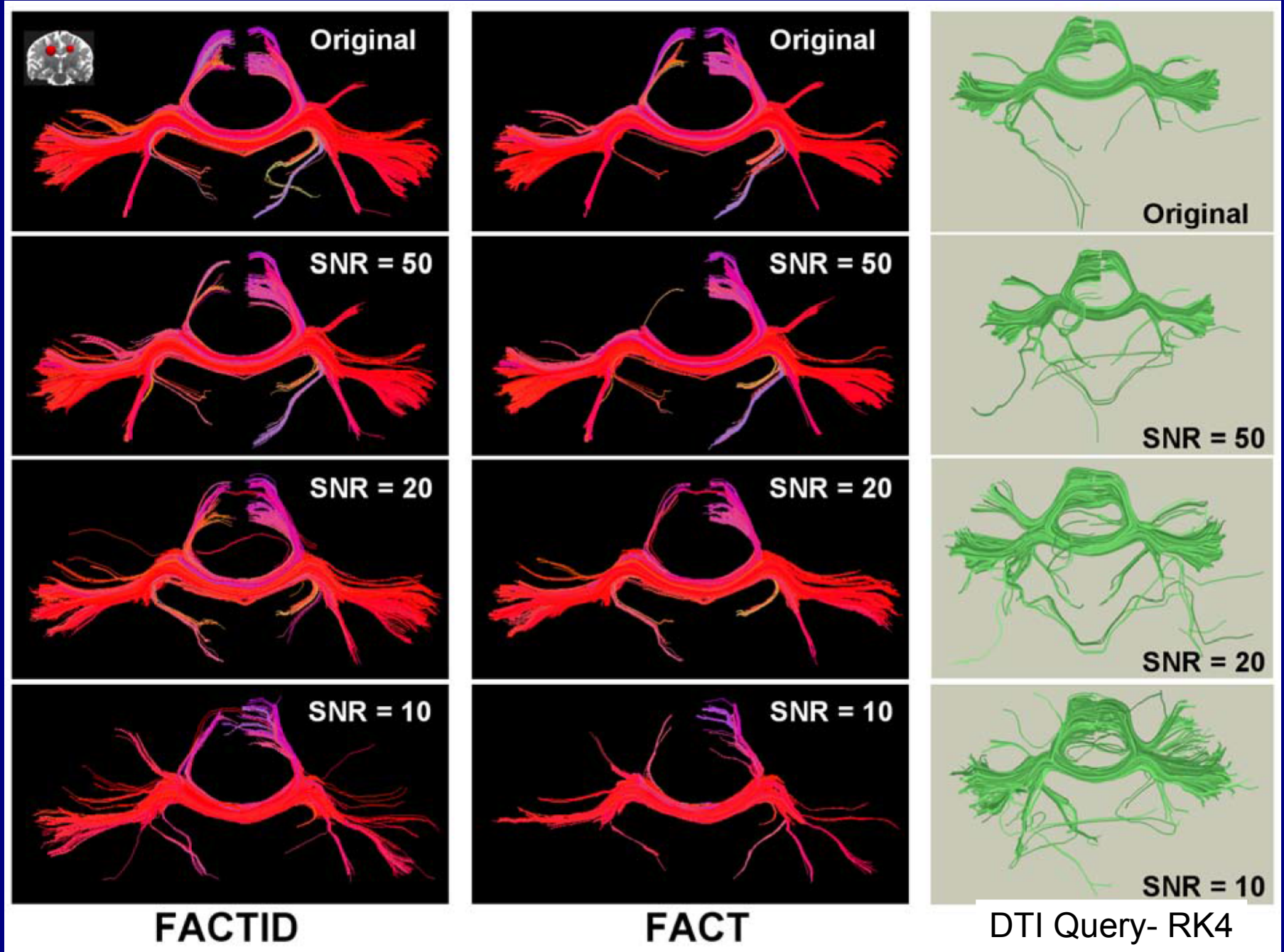


FACT



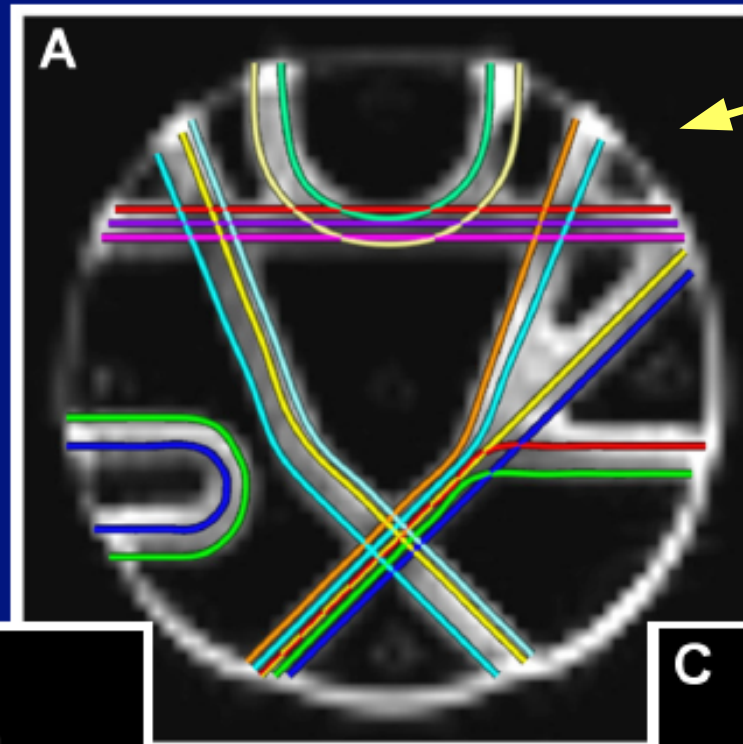
(Taylor, Cho, Lin & Biswal, 2012)

Test 3: Noise sensitivity



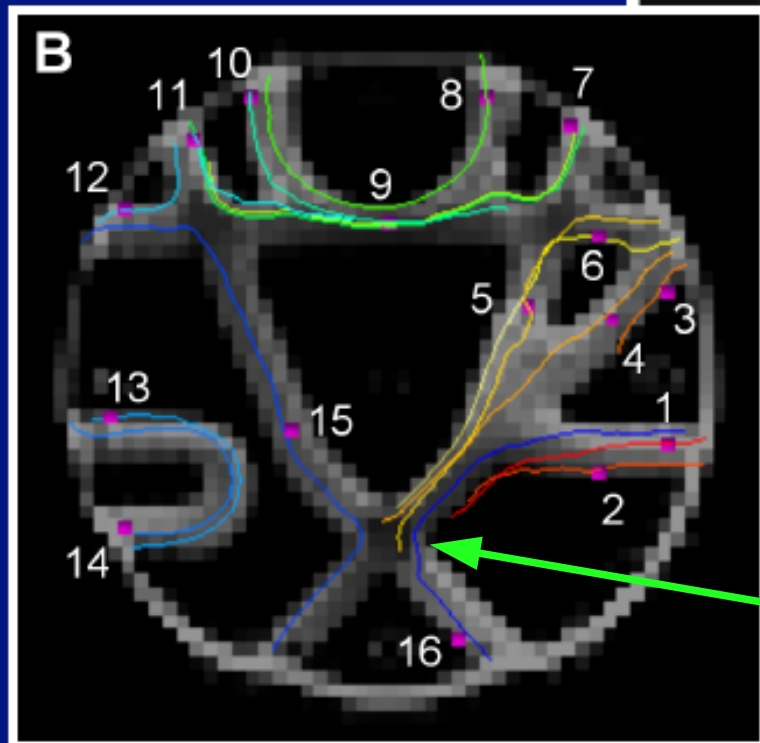
Test 5: Phantom Set

Fillard et al.
(2011, NI)
test phantom



FACT

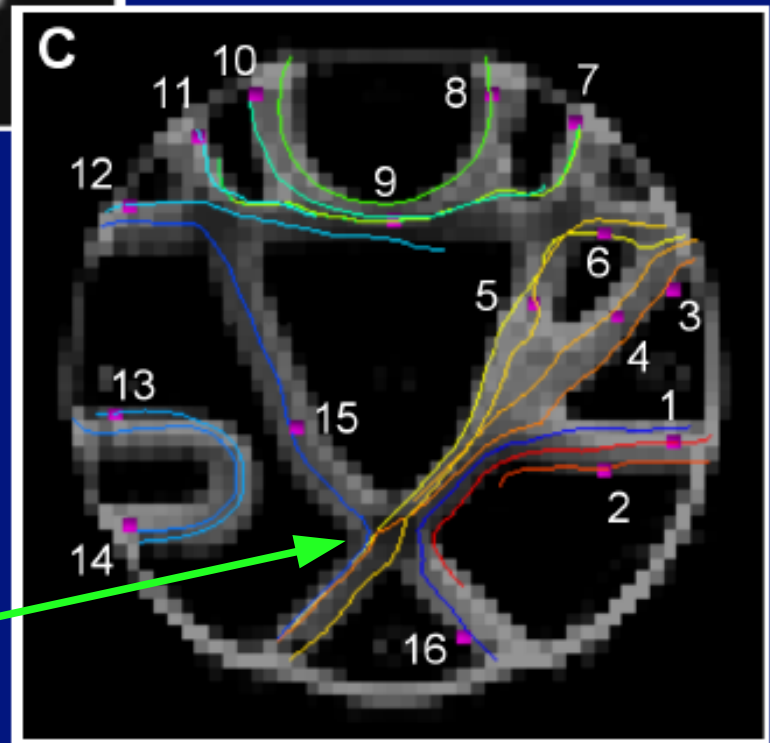
FACTID



(Taylor, Cho, Lin
& Biswal, 2012)

e.g. compare

This text is centered between images B and C. A green arrow points from the text to the vessels in image B, and another green arrow points from the text to the vessels in image C.



Importance of being processed (in earnest)

NB words of wisdom from wikipedia GIGO entry:

On two occasions I have been asked, "Pray, Mr. Babbage, if you put into the machine wrong figures, will the right answers come out?" ... I am not able rightly to apprehend the kind of confusion of ideas that could provoke such a question.

—Charles Babbage, Passages from the Life of a Philosopher

Importance of being processed (in earnest)

NB words of wisdom from wikipedia GIGO entry:

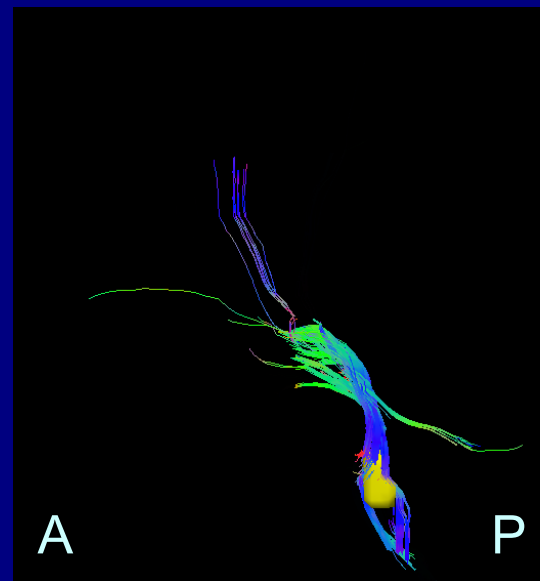
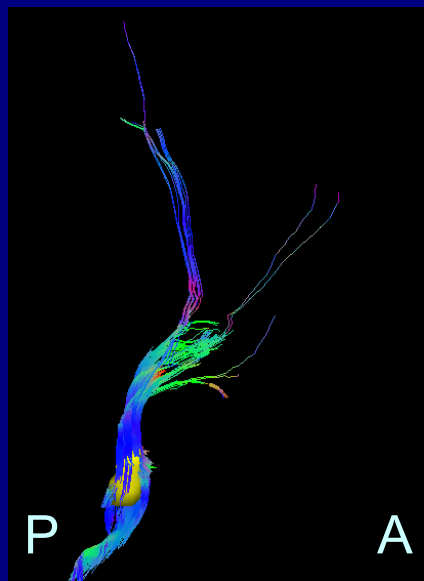
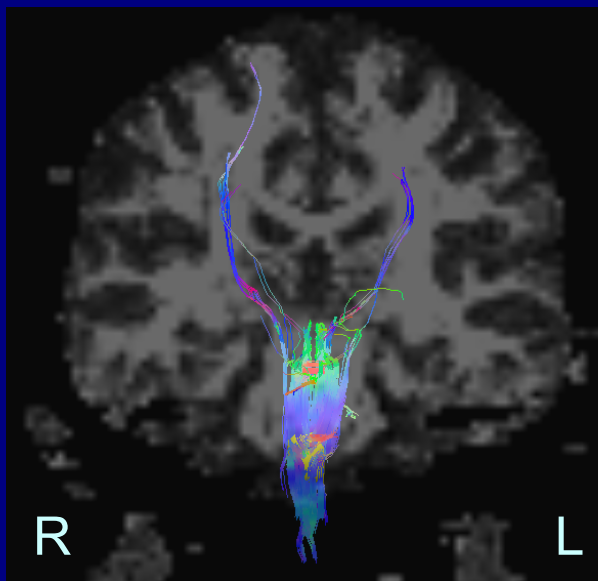
On two occasions I have been asked, "Pray, Mr. Babbage, if you put into the machine wrong figures, will the right answers come out?" ... I am not able rightly to apprehend the kind of confusion of ideas that could provoke such a question.

—Charles Babbage, Passages from the Life of a Philosopher

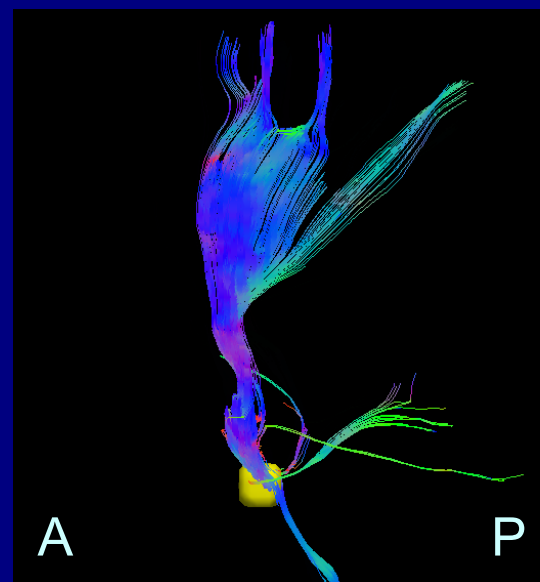
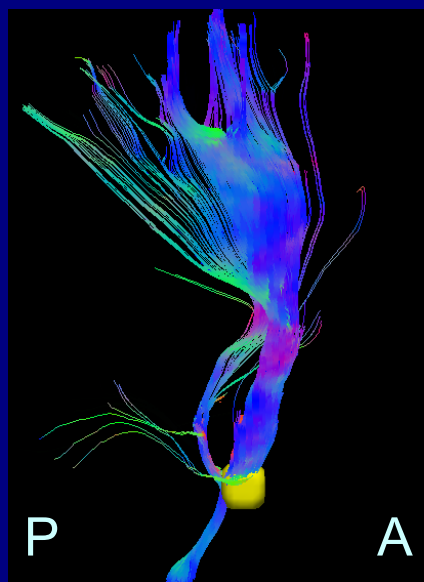
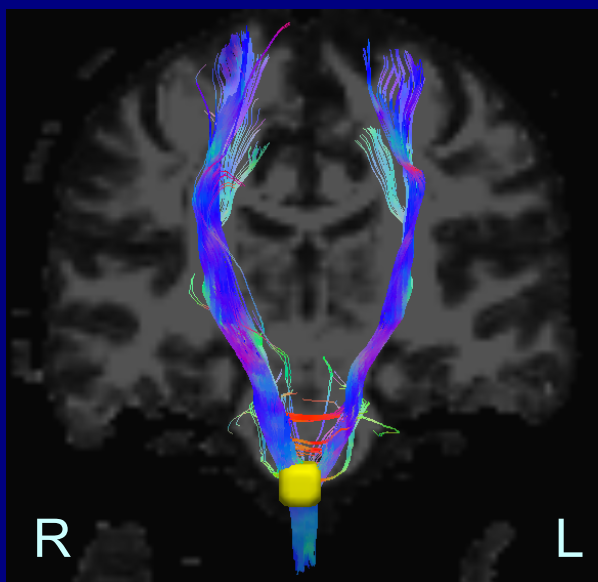
→ ** In addition to the tracking algorithm, the quality of data acquisition and preparation matter quite a bit (as seen in morning TORTOISE session). **

Importance of being processed (in earnest)

unprocessed



TORTOISED



Data from the morning session, same target ROI in brainstem.
Consider reach of tracks, symmetry, physiology, etc.

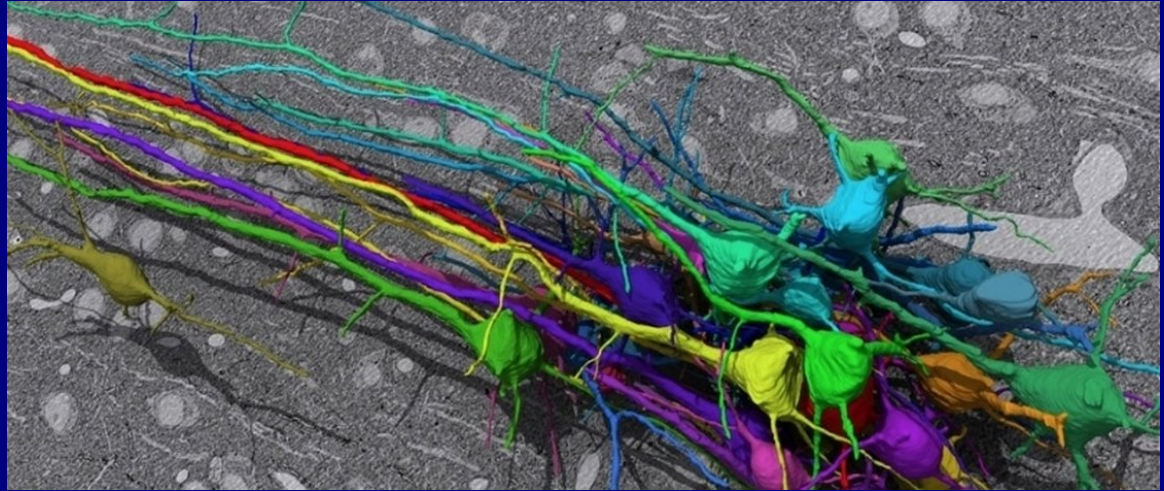
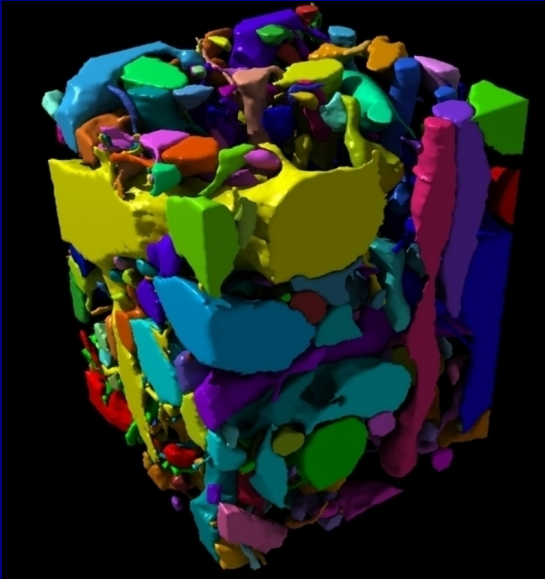
Cinematic side note:

La Belle et la Bête of tractography



Known Challenges for Tracking

- + Axon diameters are of order a few micrometers
- + MRI voxel size is of order millimeters

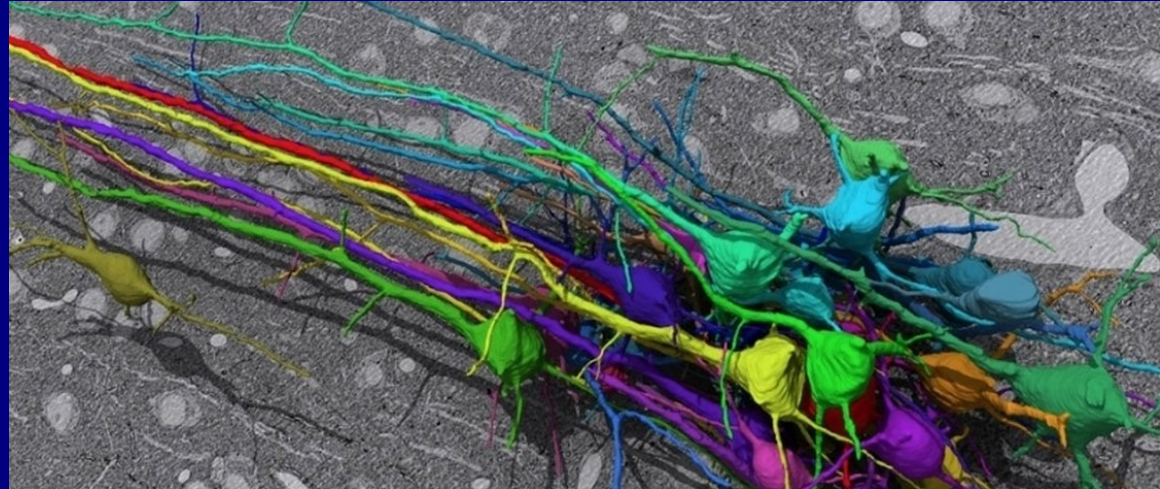
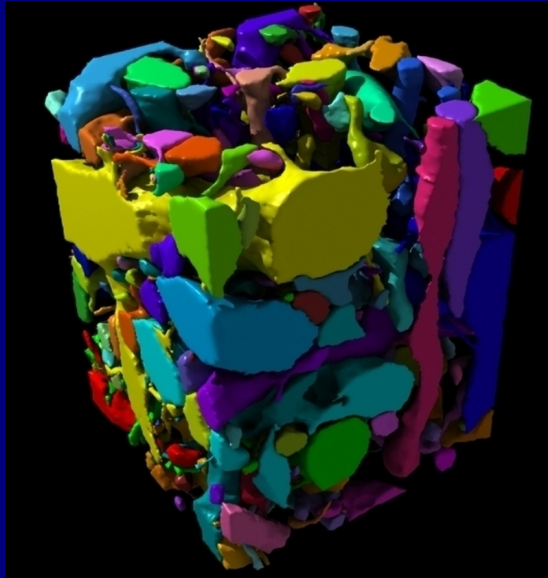


(images of Eyewire data via NPR website)

Known Challenges for Tracking

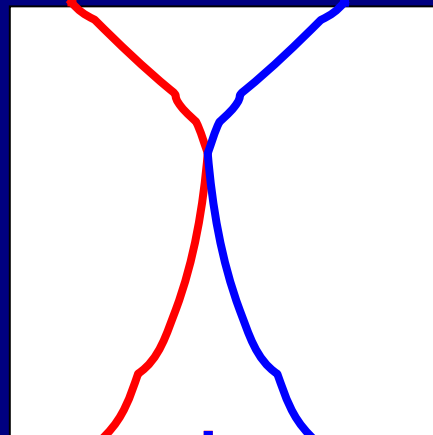
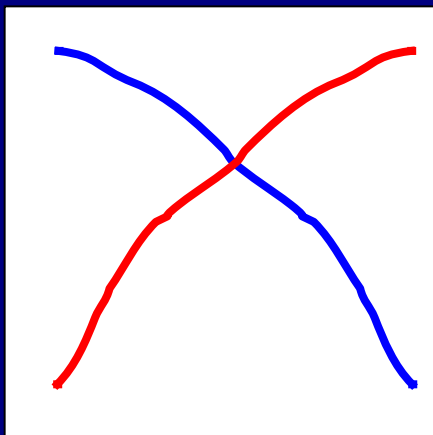


- + Axon diameters are of order a few micrometers
- + MRI voxel size is of order millimeters



(images of Eyewire data via NPR website)

- + WM regions are tightly packed, with many connections and potentially complicated sub-voxel scale structure



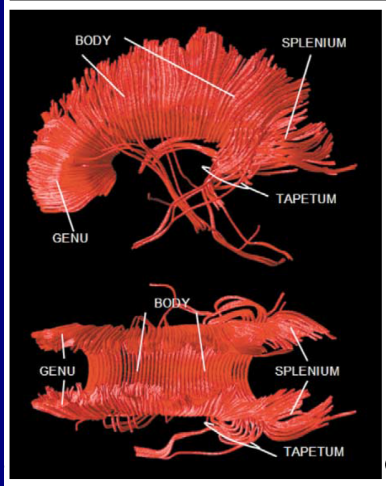
Crossing/kissing fibers can:

- Lower FA (stop tracking)
- Redirect (or *not*) tracking incorrectly.

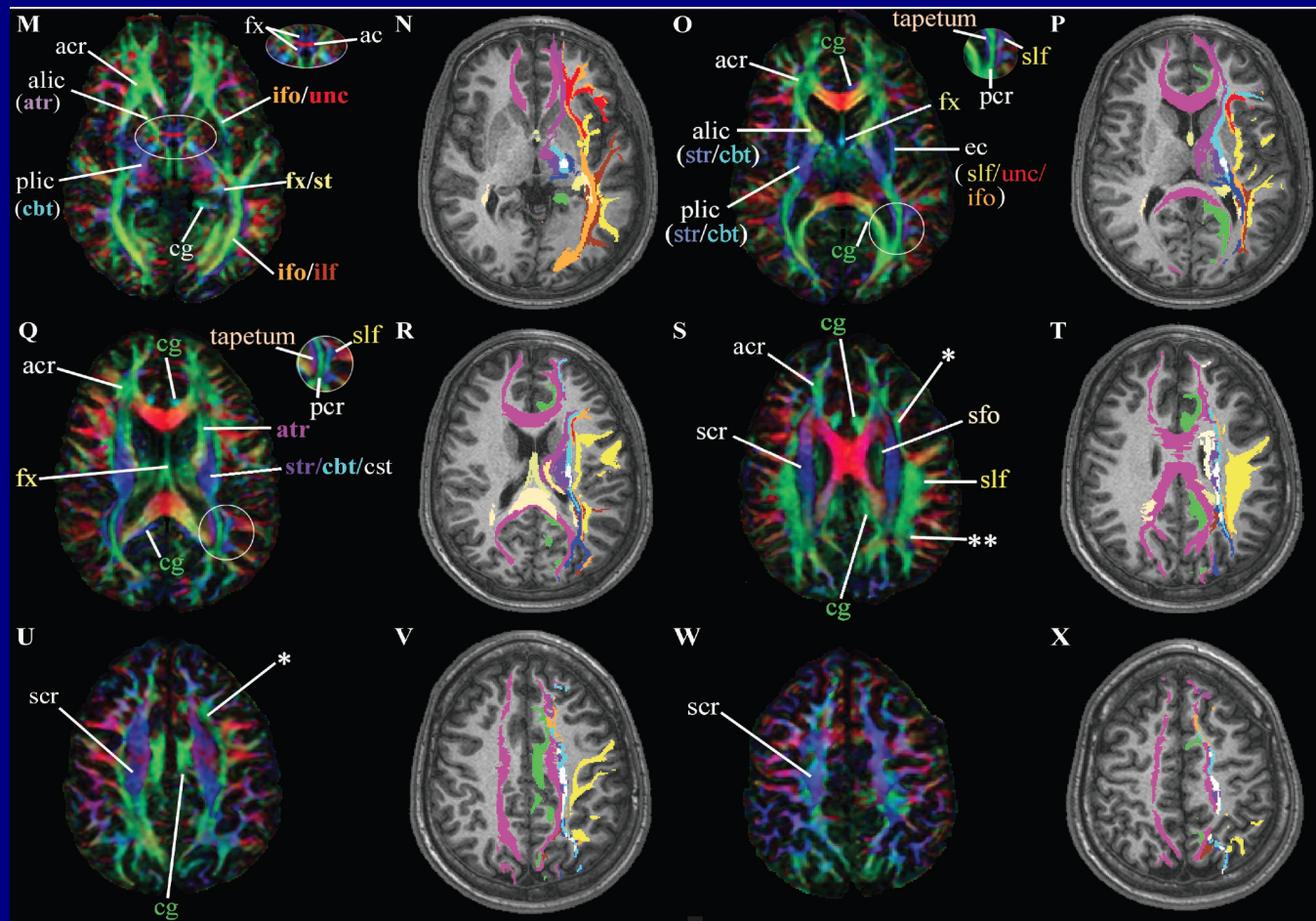
Achievements of Tracking



- + Reproduction of many known pathways
- + In vivo vs post-mortem information



(Bammer et al., 2003)



(Wakana et al., 2004)

Light at the end of the tunnel?



Application of tractography seems useful and logically consistent as follows:

- + GM ROIs *are* connected by WM skeleton.
- + Tractography can act to parcellate the WM skeleton based on subject's own data.
- + Avoid interpreting reconstructed tracks to represent literal, underlying fibers.
- + Use tracking to estimate and highlight WM likely to be associated with GM ROIs.
- + One can then use diffusion parameters in those 'WM ROIs' for quantitative comparisons (or use ROIs as masks for other data).

Next question for doing tractography:

where does one go to get the ROIs to try to connect?

Next question for doing tractography:

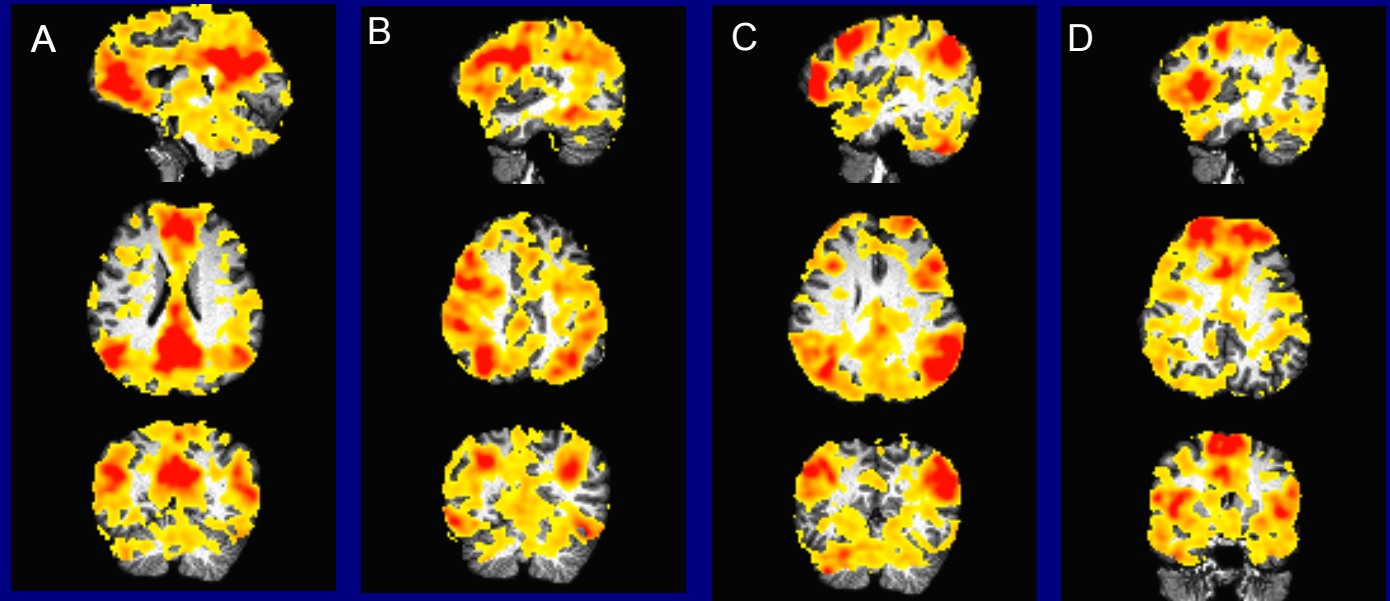
where does one go to get the ROIs to try to connect?

-> could go to atlases and standard maps,
or to exploratory spheres dotted around,

FMRI measures -> networks of ROIs

+ For example, one can perform ICA on a resting state study, resulting in several functional networks:

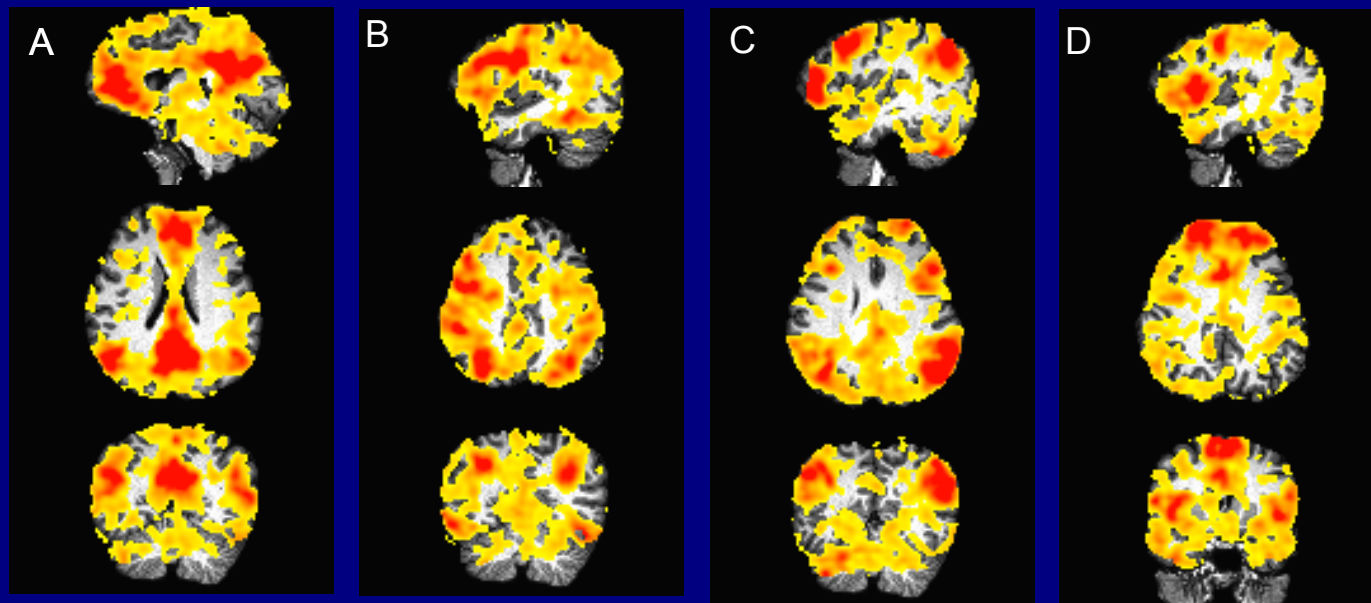
(each IC is map of Z-scores; here, shown for $Z > 0$)



FMRI measures -> networks of ROIs

- + For example, one can perform ICA on a resting state study, resulting in several functional networks:

(each IC is map of Z-scores; here, shown for $Z > 0$)

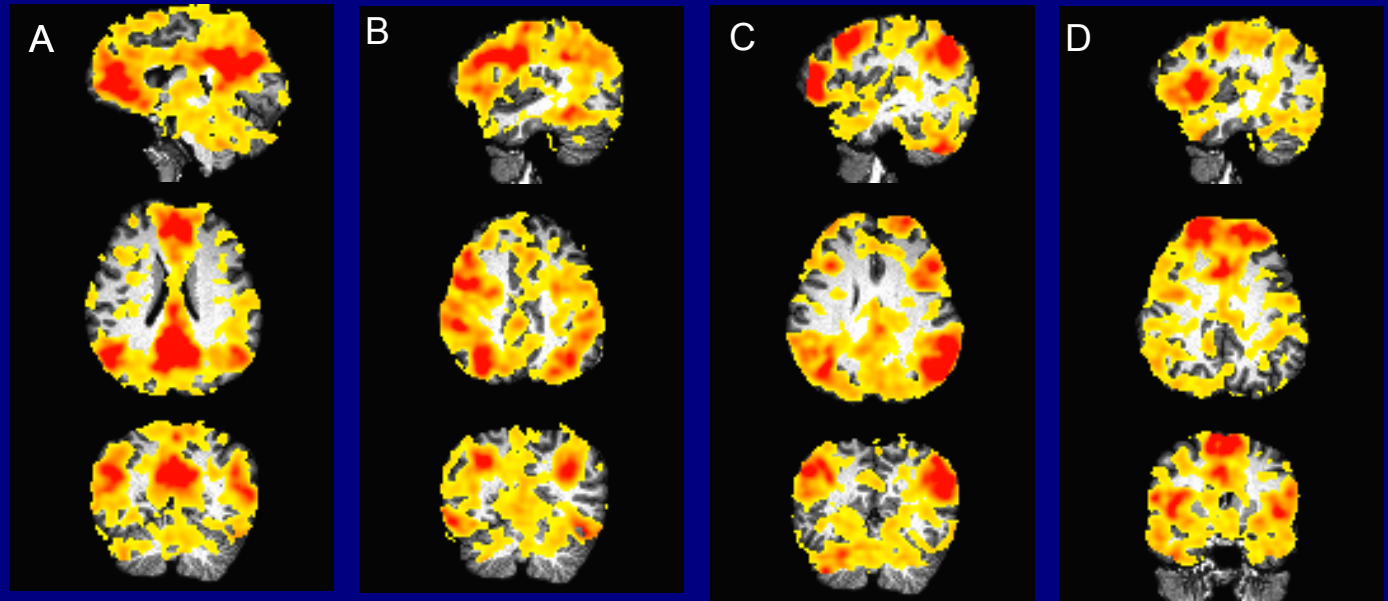


- + want to **isolate GM** ROIs, and then to **expand each** to make sure that they are at least touching nearby (*associated?*) WM voxels to have any hope to connect tracts

FMRI measures -> networks of ROIs

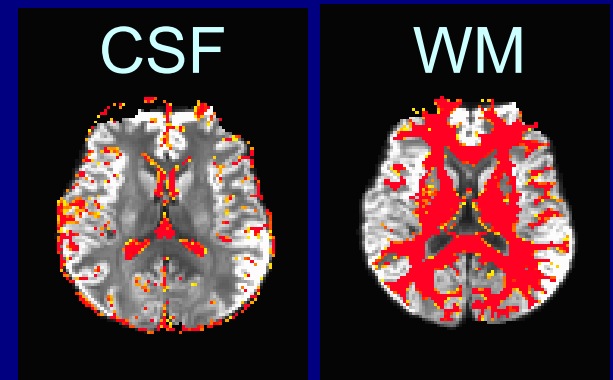
- + For example, one can perform ICA on a resting state study, resulting in several functional networks:

(each IC is map of Z-scores; here, shown for $Z > 0$)



- + **3dROIMaker** can parcellate into GM ROIs based on:

- thresholding **voxel values**
- thresholding **cluster size**
- subtract away CSF and WM voxels from segmentation maps
- **expand** each GM ROI to location of WM (don't want to *overexpand* unphysically)



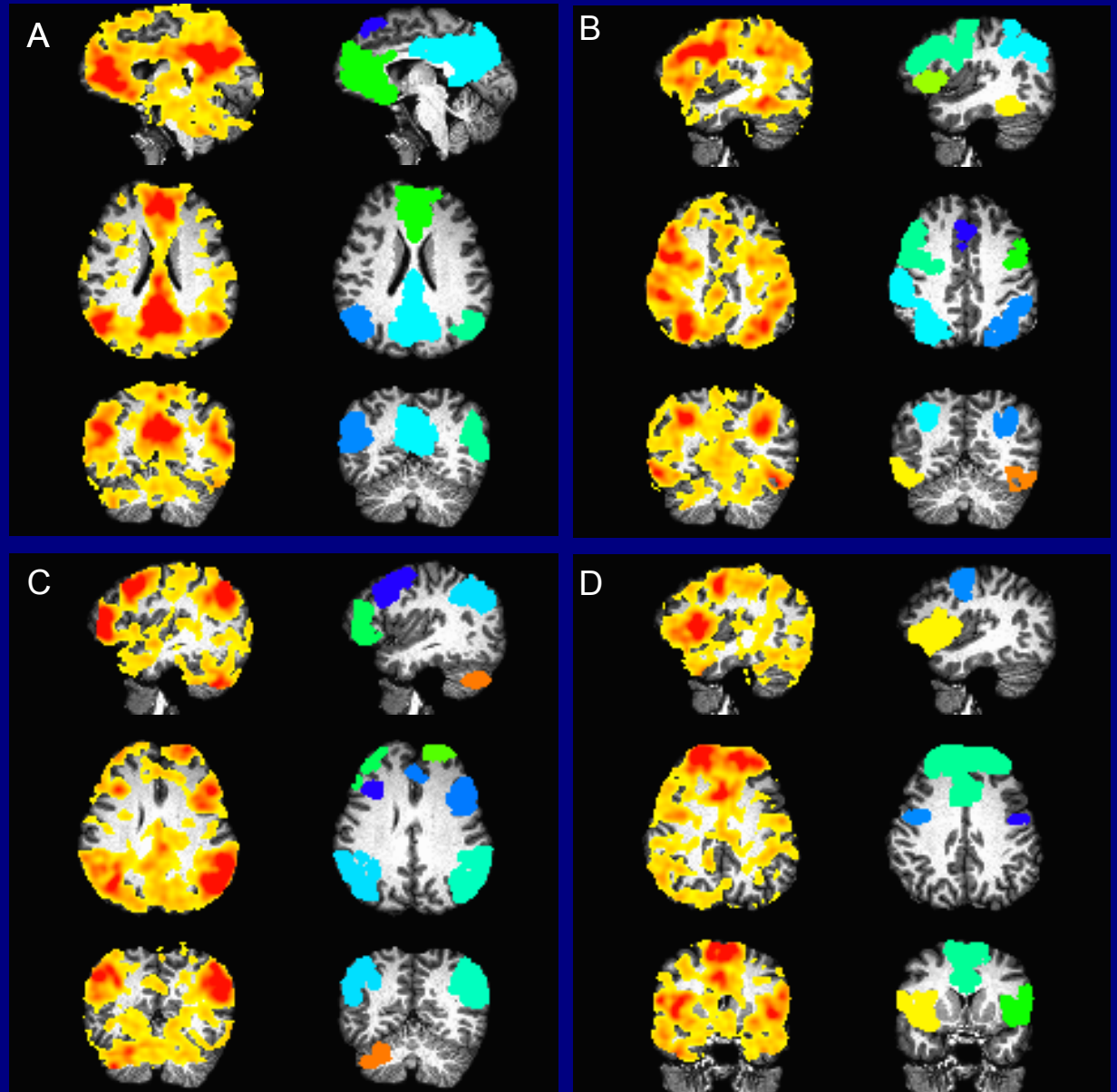
FMRI measures -> networks of ROIs

Example case for ICA networks:

Left col: ICA map (visualized at $Z > 0$, for clarity).

*Right col: ROIMaker ROI map, thresholded $Z > 3.0$
cluster volume > 130 voxels
expand clusters +2 voxels
limit expansion with $FA > 0.2$ info.
(An unexpanded set of maps is also made and saved.)*

Sidenote: this involved mapping FMRI data of ICs and T1 tissue segmentation results into DWI space; used 3dAllineate.



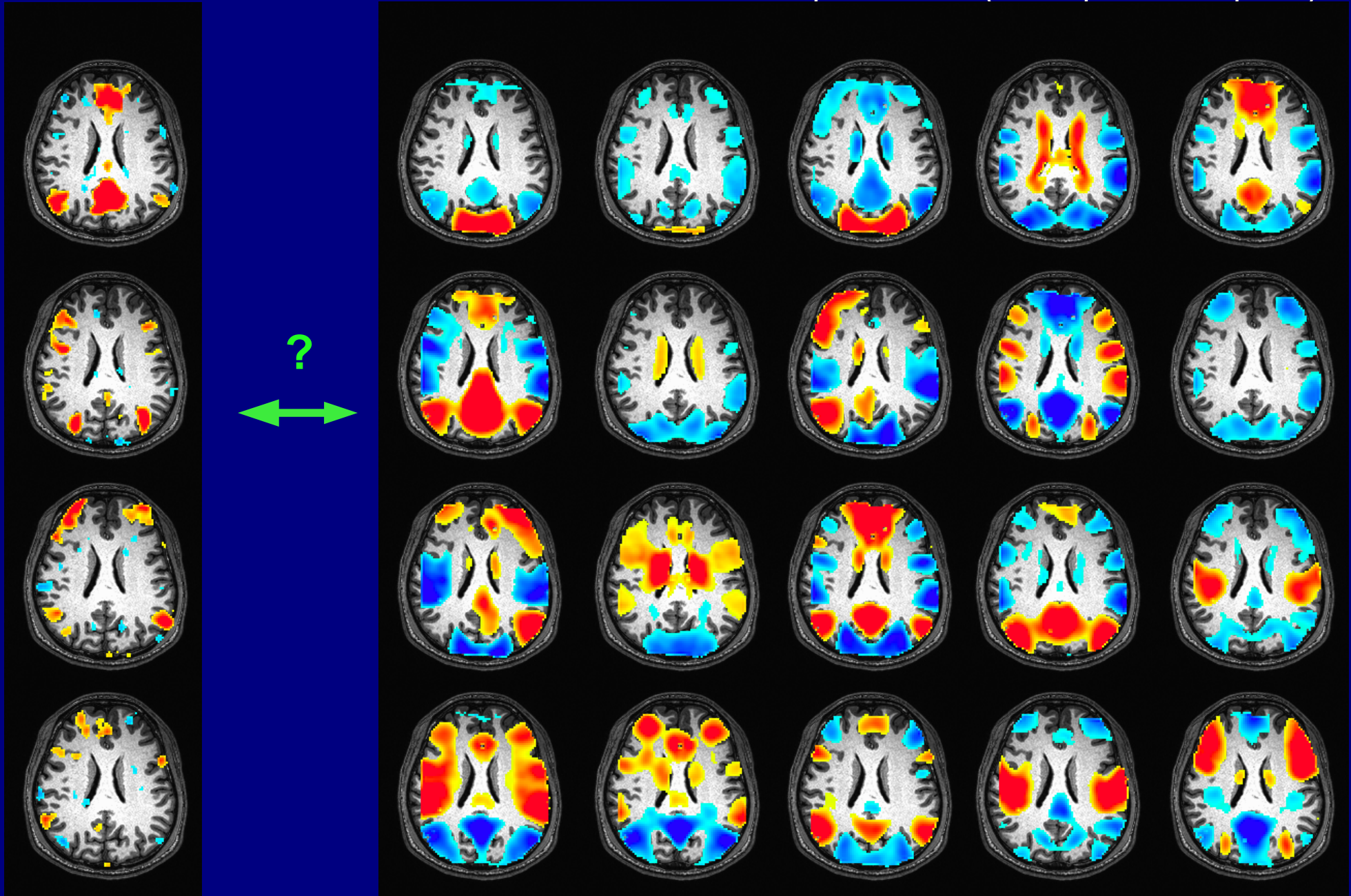
Sidenote:

***How to identify network maps, or
match them with reference/group set?***

Matching Network maps

Some Z-score

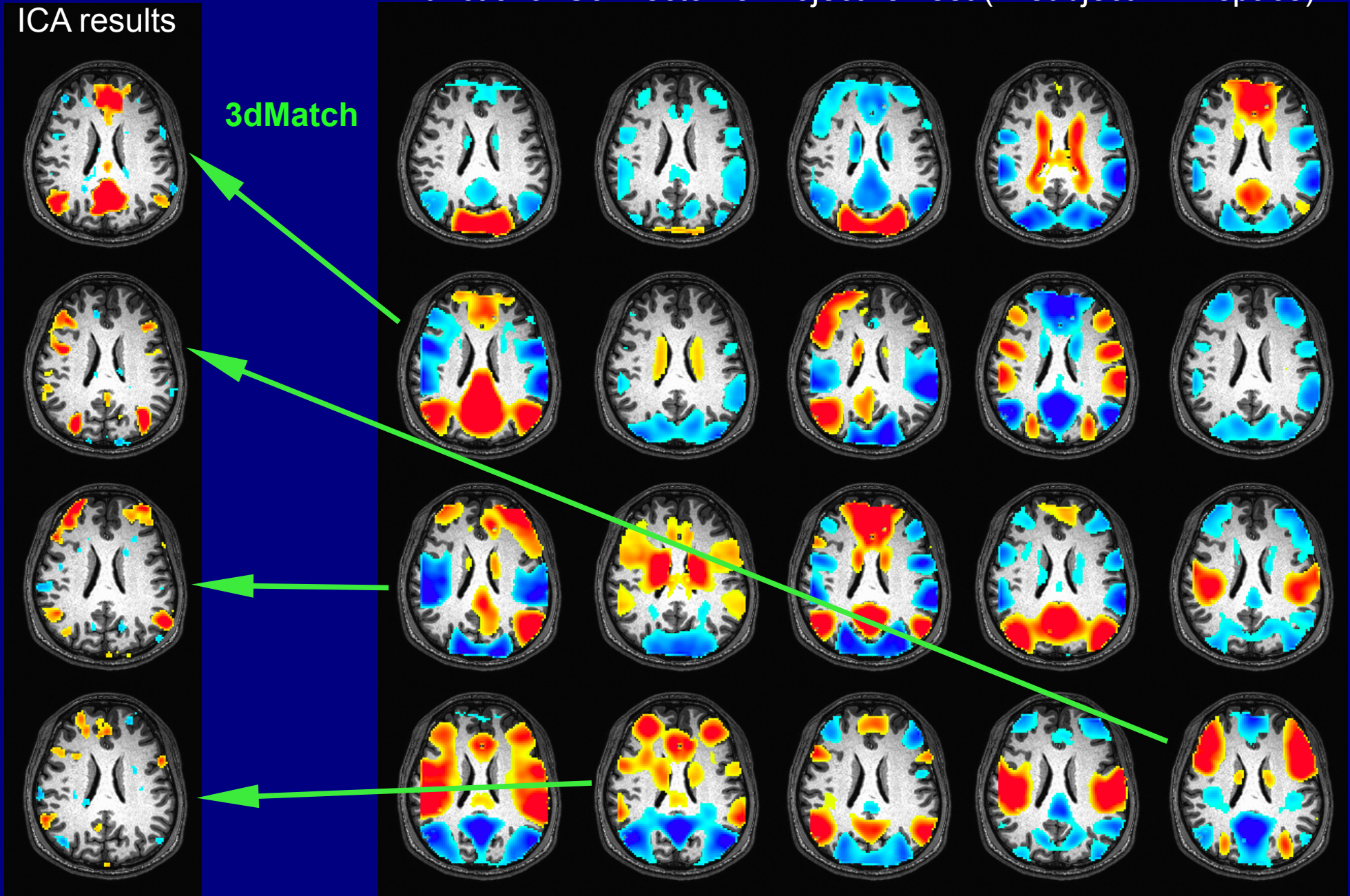
Functional Connectome Project ref. set (in subject DWI space)



Matching Network maps

Some Z-score
ICA results

Functional Connectome Project ref. set (in subject DWI space)



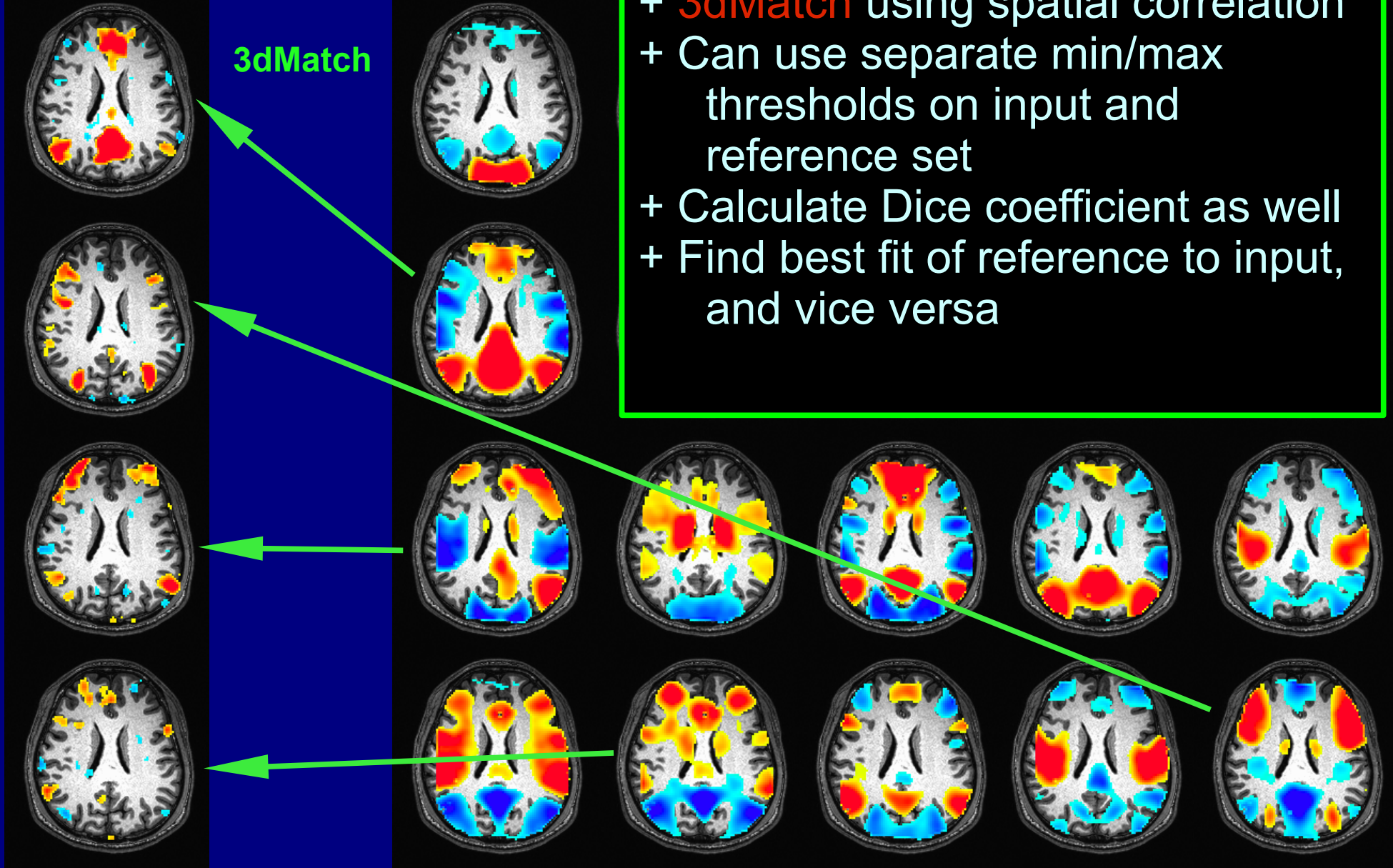
Matching Network maps

Some Z-score
ICA results

Functional Connectome Project ref. set (in subject DWI space)

3dMatch

- + 3dMatch using spatial correlation
- + Can use separate min/max thresholds on input and reference set
- + Calculate Dice coefficient as well
- + Find best fit of reference to input, and vice versa



Deterministic tractography

+ 3dTrackID -mode DET -logic { OR | AND }

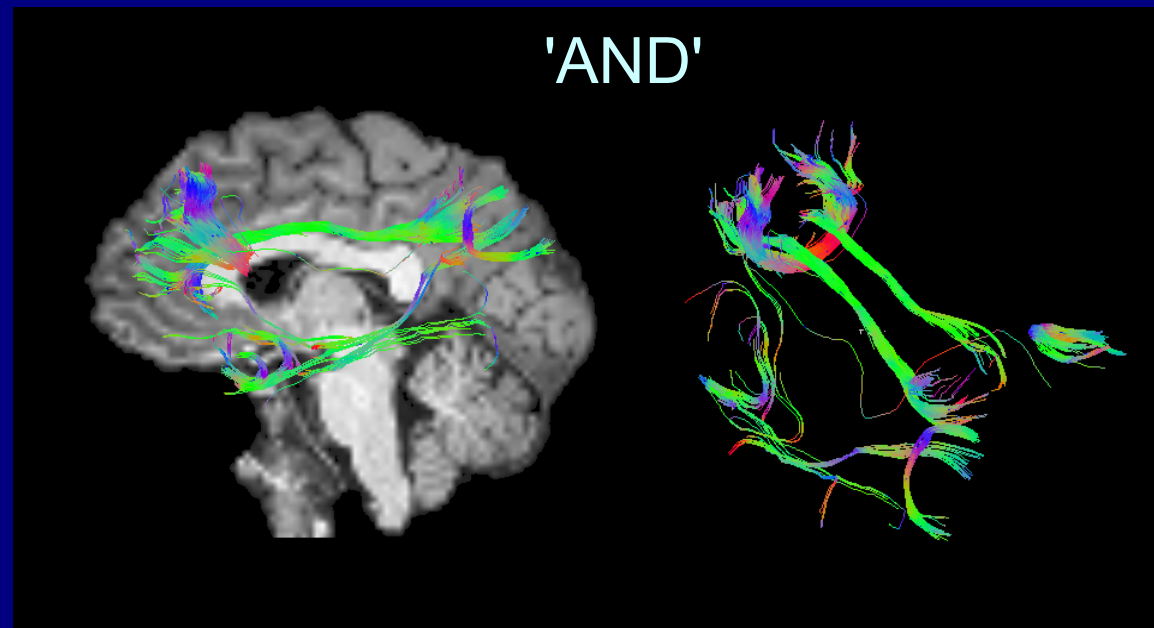
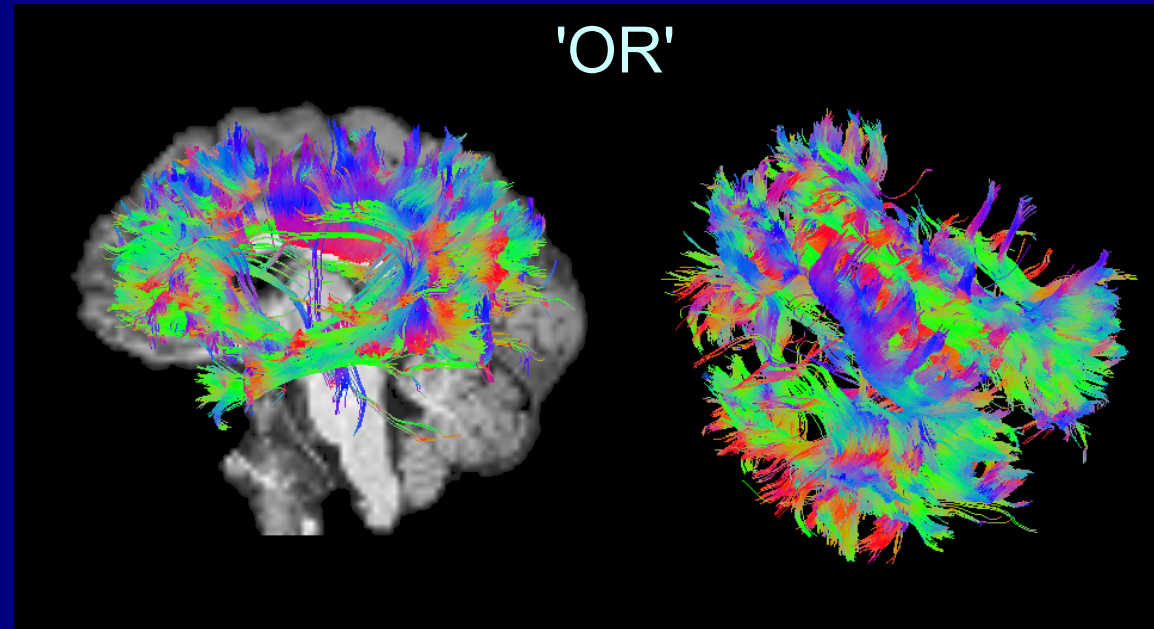
+ uses FACTID

+ good for exploratory
analysis and visualization
of results

ex.: DMN network tractography
results using ROIs from

3dROIMaker

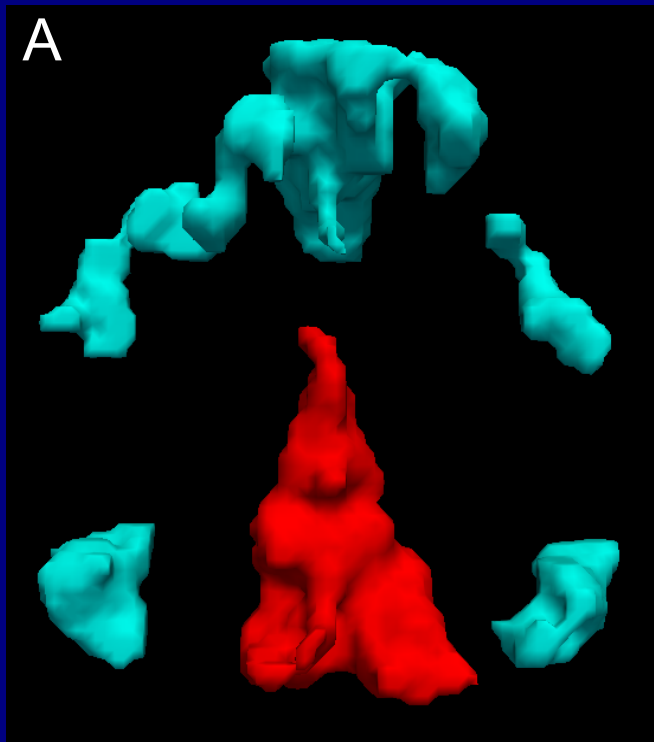
($FA > 0.2$; max angle 60deg;
8 seeds/voxel)



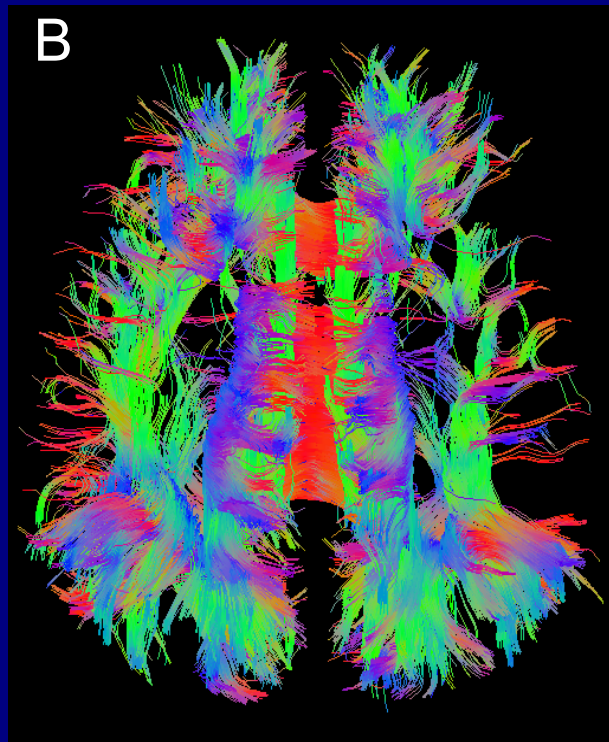
Deterministic tractography

+ 3dTrackID -mode DET -logic { OR | AND }

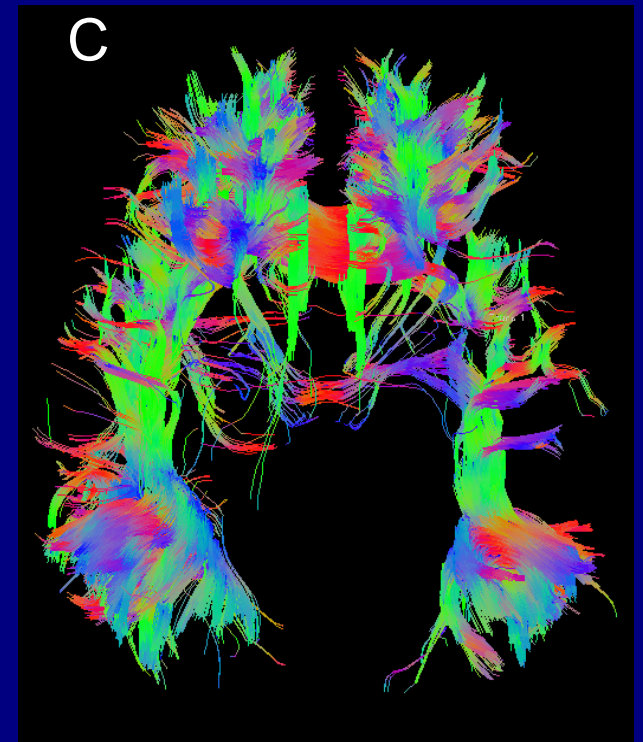
Control track propagation with `anti-mask' regions,
simply defined by voxels =-1:



ROIs: blue >0 , red <0



results when:
all ROIs >0
(no anti-mask)



results when:
blue >0 , red <0
(using anti-masks)

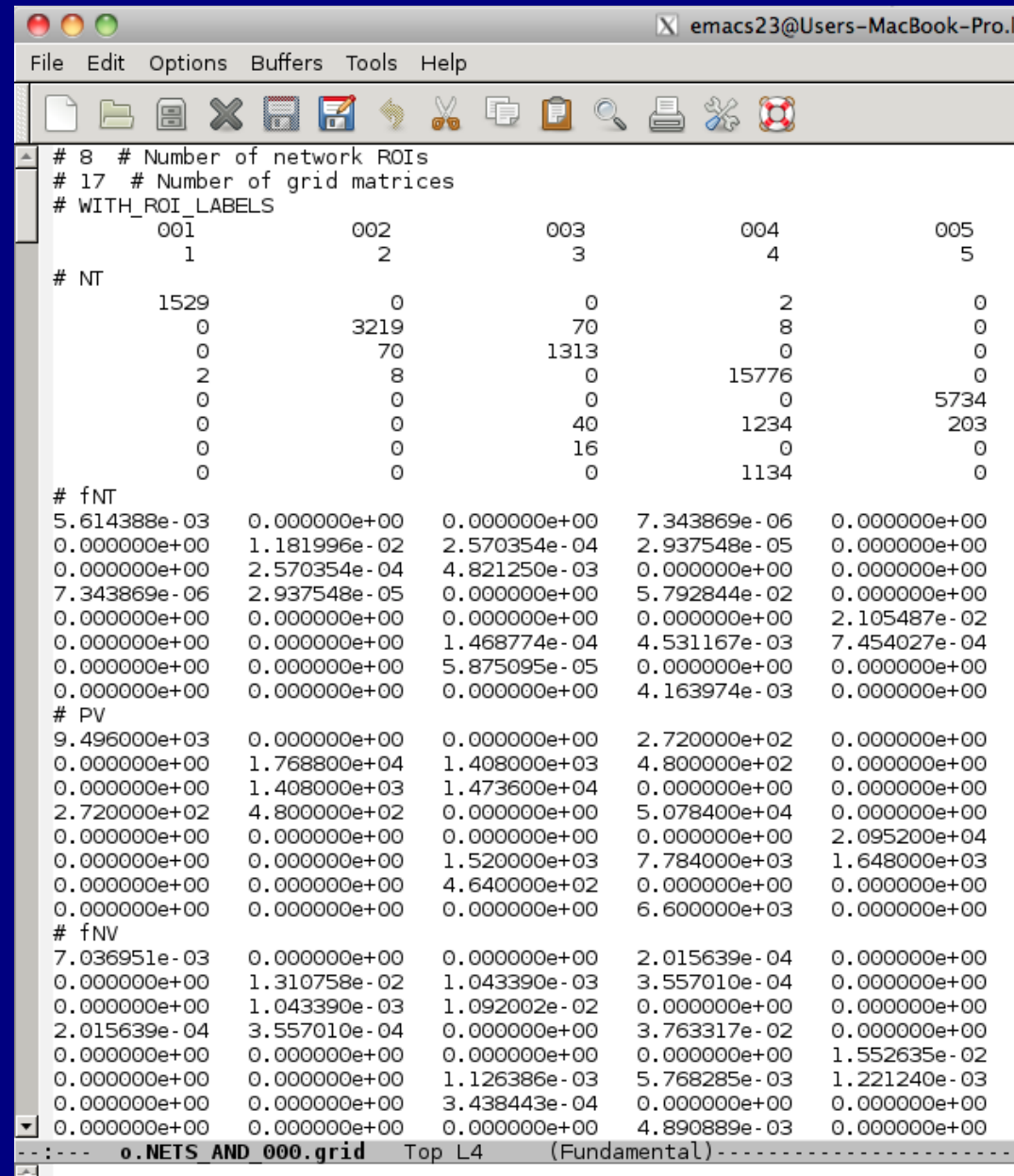
Deterministic tractography

+ 3dTrackID -mode DET -logic { OR | AND }

+ Automatic quantification per network in produced PREFIX.grid files.

Matrices of per-connection parameters such as:
mean/std of FA, MD, RD, L1, numbers of tracts, volume of tracts (and options for scaling tract-stats by ROI volumes)

+ possible to load in other files for automatic statistics, also.



```
# 8 # Number of network ROIs
# 17 # Number of grid matrices
# WITH_ROI_LABELS
      001      002      003      004      005
      1      2      3      4      5
# NT
      1529      0      0      2      0
      0      3219      70      8      0
      0      70      1313      0      0
      2      8      0      15776      0
      0      0      0      0      5734
      0      0      40      1234      203
      0      0      16      0      0
      0      0      0      1134      0
# fNT
5.614388e-03 0.000000e+00 0.000000e+00 7.343869e-06 0.000000e+00
0.000000e+00 1.181996e-02 2.570354e-04 2.937548e-05 0.000000e+00
0.000000e+00 2.570354e-04 4.821250e-03 0.000000e+00 0.000000e+00
7.343869e-06 2.937548e-05 0.000000e+00 5.792844e-02 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 2.105487e-02
0.000000e+00 0.000000e+00 1.468774e-04 4.531167e-03 7.454027e-04
0.000000e+00 0.000000e+00 5.875095e-05 0.000000e+00 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 4.163974e-03 0.000000e+00
# PV
9.496000e+03 0.000000e+00 0.000000e+00 2.720000e+02 0.000000e+00
0.000000e+00 1.768800e+04 1.408000e+03 4.800000e+02 0.000000e+00
0.000000e+00 1.408000e+03 1.473600e+04 0.000000e+00 0.000000e+00
2.720000e+02 4.800000e+02 0.000000e+00 5.078400e+04 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 2.095200e+04
0.000000e+00 0.000000e+00 1.520000e+03 7.784000e+03 1.648000e+03
0.000000e+00 0.000000e+00 4.640000e+02 0.000000e+00 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 6.600000e+03 0.000000e+00
# fNV
7.036951e-03 0.000000e+00 0.000000e+00 2.015639e-04 0.000000e+00
0.000000e+00 1.310758e-02 1.043390e-03 3.557010e-04 0.000000e+00
0.000000e+00 1.043390e-03 1.092002e-02 0.000000e+00 0.000000e+00
2.015639e-04 3.557010e-04 0.000000e+00 3.763317e-02 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 1.552635e-02
0.000000e+00 0.000000e+00 1.126386e-03 5.768285e-03 1.221240e-03
0.000000e+00 0.000000e+00 3.438443e-04 0.000000e+00 0.000000e+00
0.000000e+00 0.000000e+00 0.000000e+00 4.890889e-03 0.000000e+00
--:--- o.NETS AND 000.grid Top L4 (Fundamental)-----
```


Deterministic tractography

+ 3dTrackID -mode DET -logic { OR | AND }

+ uses FACTID

+ good for exploratory analysis and visualization of results

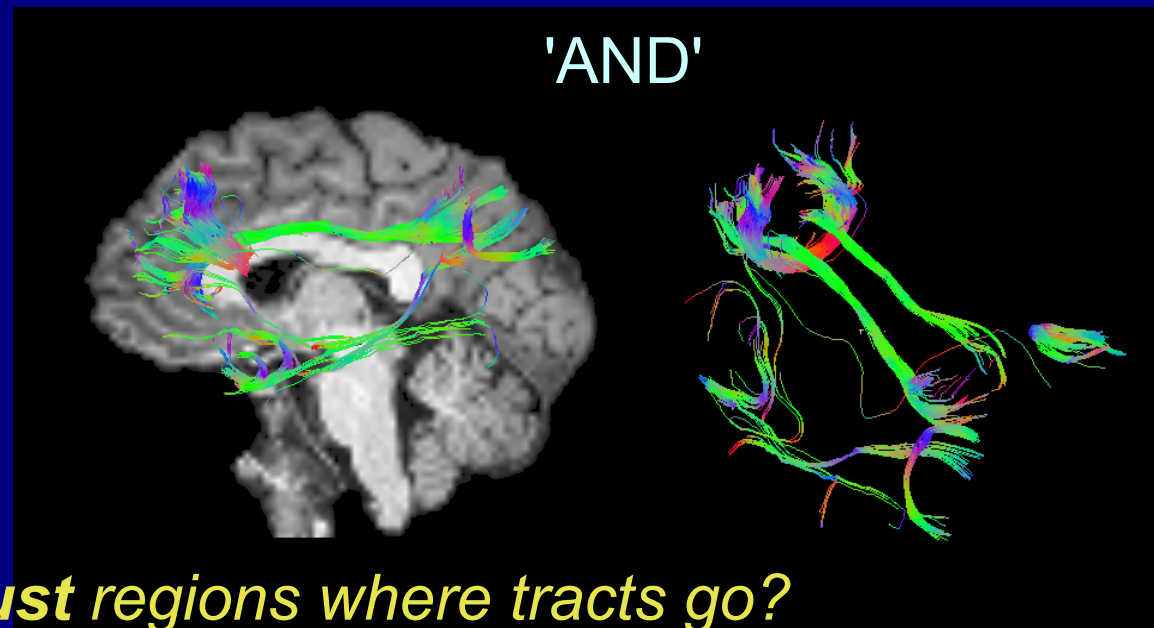
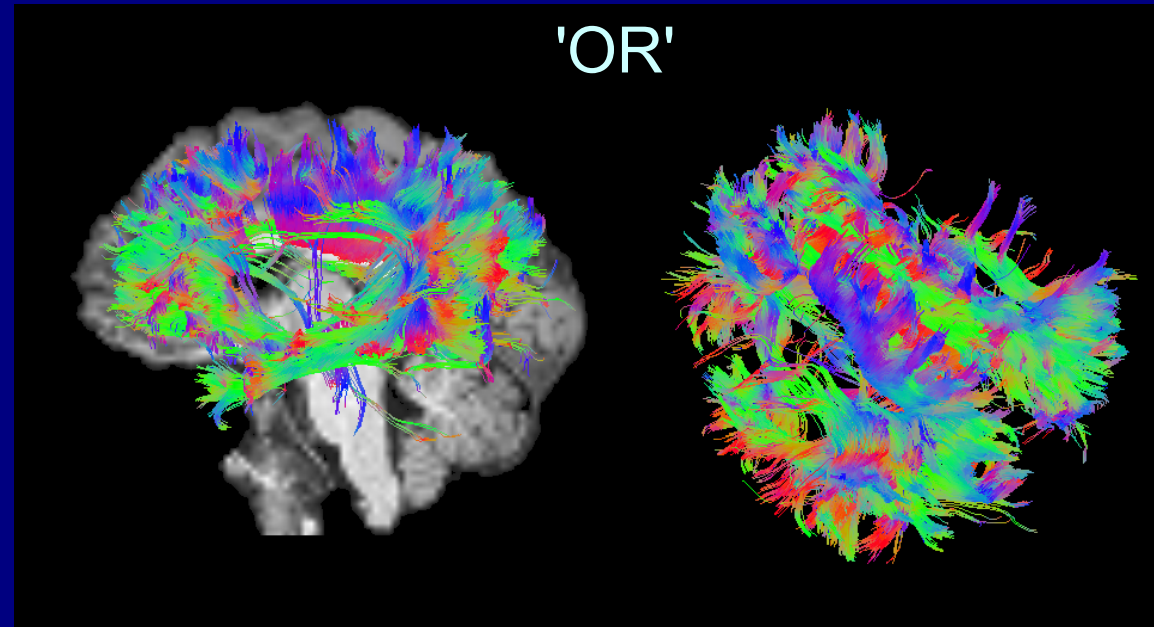
ex.: DMN network tractography results using ROIs from

3dROIMaker

($FA > 0.2$; max angle 60deg; 8 seeds/voxel)

*Tract results may seem 'fine', but is **noise** affecting them?*

*Are these the **most likely/robust** regions where tracts go?*



Brings up next question for doing tractography:

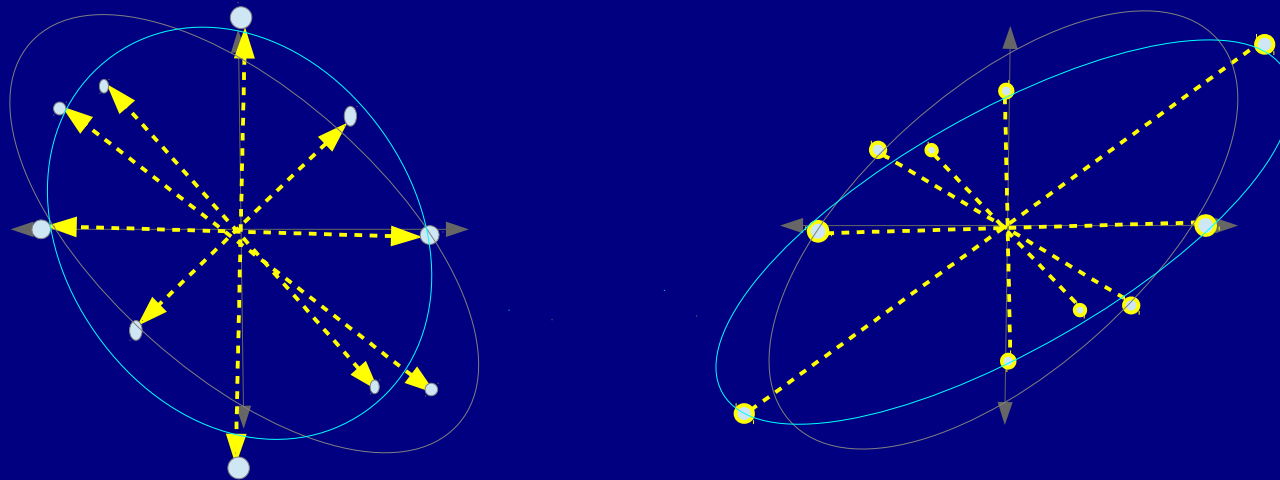
***How do we estimate tensor parameter
noise/uncertainty?***

Noise in DW signals

MRI signals have additive noise

$$S_i = S_0 e^{-b \mathbf{g}_i^T \mathbf{D} \mathbf{g}_i} + \varepsilon,$$

where ε is (Rician) noise, with the effect of leading to errors in surface fit, equivalent to *rotations* and *rescalings* of ellipsoids:



'Un-noisy' vs perturbed/noisy fit

EPI distortions, subject motion, et al. also warp ellipsoids.

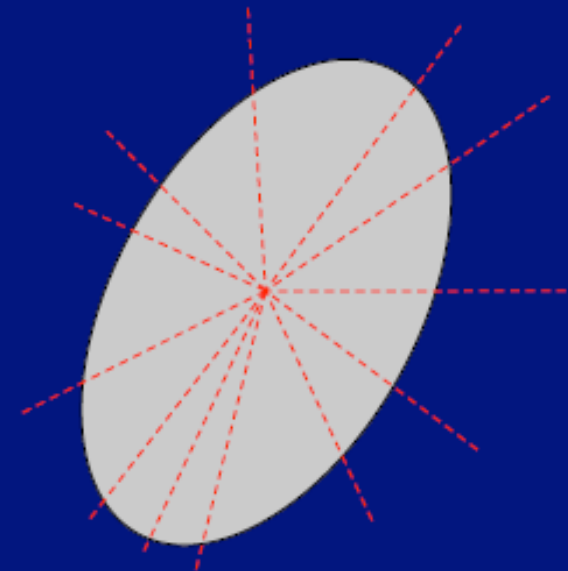
DTI Uncertainty

- We use jackknife resampling (e.g., Efron 1982)
 - Other studies have used bootstrapping (e.g., Jones 2003), or theoretical estimates (Jeong & Anderson 2008)
 - Jackknifing is efficient (just need one data set unlike bootstrap), simpler than theory, since, e.g., SNR is likely not constant across voxels

Jackknifing

- Basically, take M acquisitions

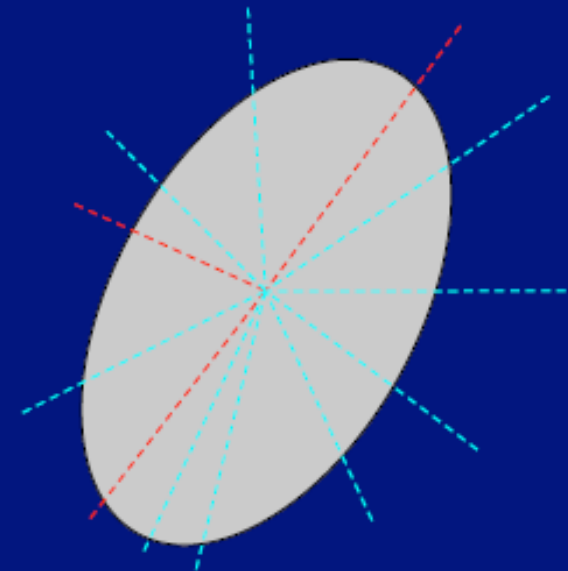
e.g., $M=12$



Jackknifing

- Basically, take M acquisitions
- Randomly select $M_J < M$ to use to calculate quantity of interest
 - standard nonlinear fits

e.g., $M=12$
 $M_J=9$

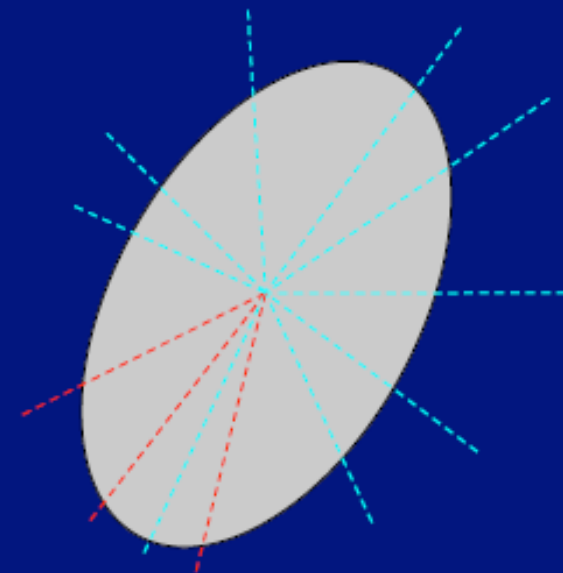


$$[D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] = \dots$$

Jackknifing

- Basically, take M acquisitions
- Randomly select $M_J < M$ to use to calculate quantity of interest
 - standard nonlinear fits
- Repeatedly subsample large number ($\sim 10^3$ - 10^4 times)

e.g., $M=12$
 $M_J=9$

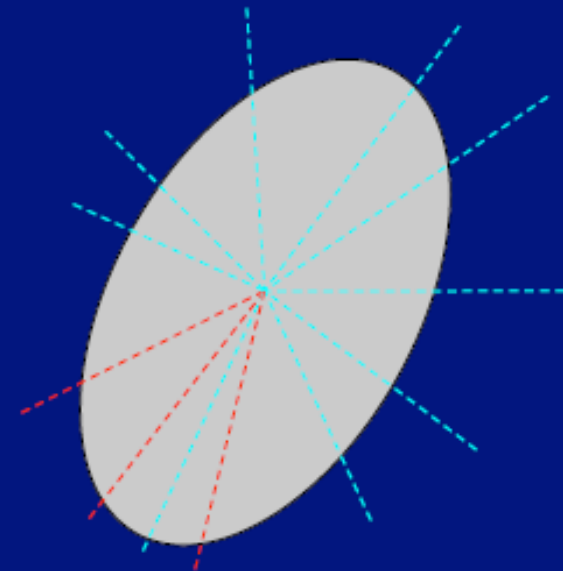


$$\begin{aligned} [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ &\dots \end{aligned}$$

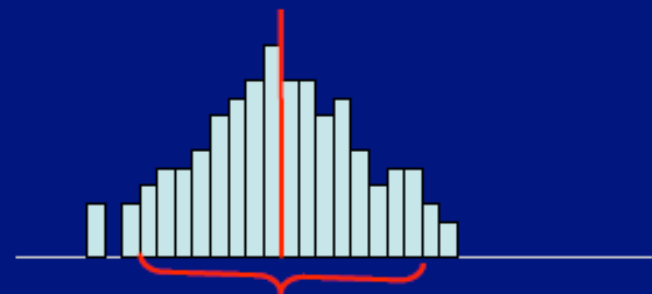
Jackknifing

- Basically, take M acquisitions
- Randomly select $M_J < M$ to use to calculate quantity of interest
 - standard nonlinear fits
- Repeatedly subsample large number ($\sim 10^3$ - 10^4 times)
- Analyze distribution of values for estimator (mean) and confidence interval
 - sort/%iles
 - (not so efficient)
 - if Gaussian, e.g. $\mu \pm 2\sigma$
 - simple

e.g., $M=12$
 $M_J=9$

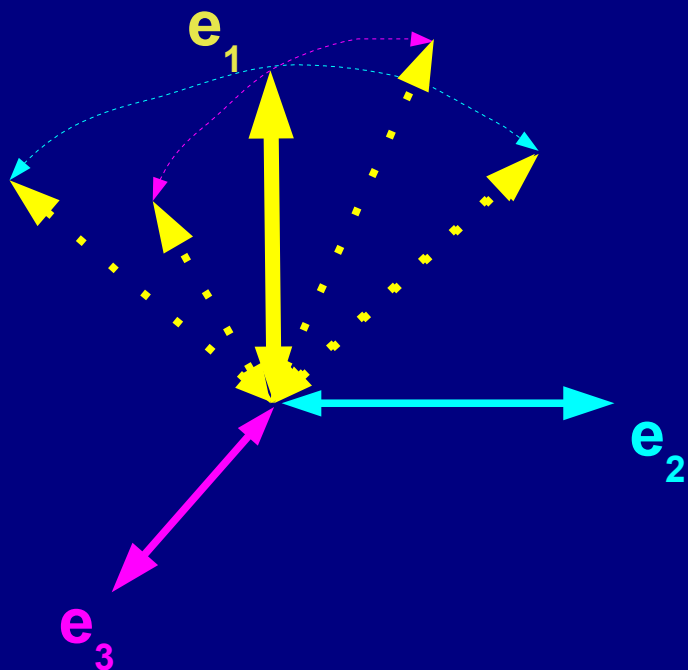


$$\begin{aligned} [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ [D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] &= \dots \\ &\dots \end{aligned}$$



Uncertainty estimation

+ **3dDWUncert** estimates bias and σ of first eigenvector \mathbf{e}_1 (main direction of diffusion), based on how much it could tip toward either \mathbf{e}_2 or \mathbf{e}_3 :



.... and the bias and σ of FA

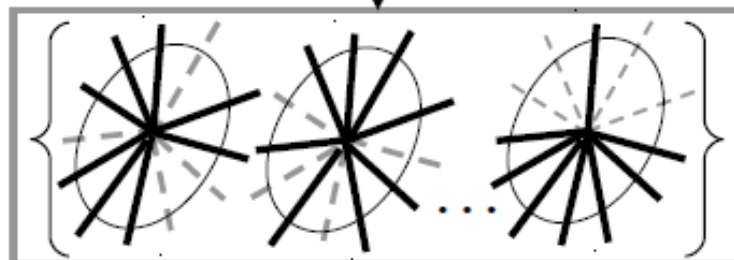
1) Obtain M DWIs.



1b) Estimate DT and parameters from M DWIs.

$\hat{\mathbf{D}}, \hat{\mathbf{F}}\mathbf{A}, \dots$

2) Make N_j subsets of M_j DWIs.



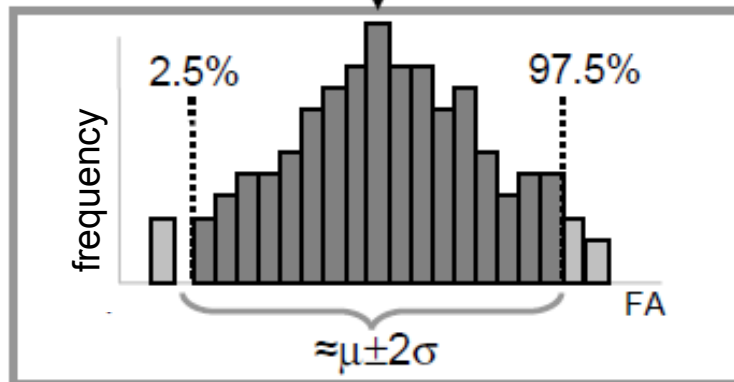
3) Estimate N_j DTs.

$\mathbf{D}_1^* \quad \mathbf{D}_2^* \quad \dots \quad \mathbf{D}_{N_j}^*$

4) Estimate set of N_j parameters.

$\{\mathbf{F}\mathbf{A}_1^*, \mathbf{F}\mathbf{A}_2^*, \dots, \mathbf{F}\mathbf{A}_{N_j}^*\}, \{(\Delta\mathbf{e}_{1,2}^*)_i\}, \dots$

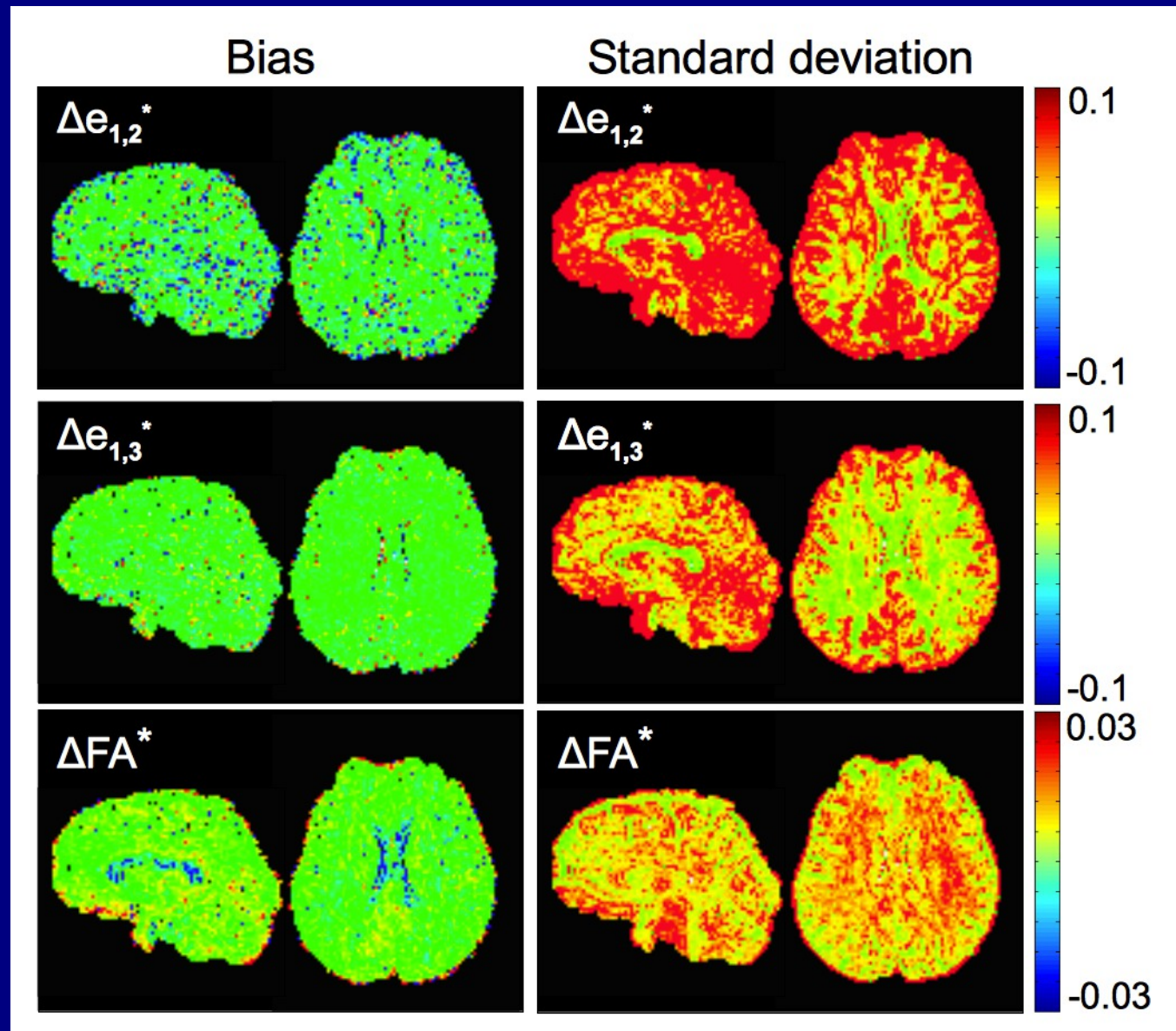
5) Find confidence intervals.



Uncertainty example

+ Can see difference in e_1 uncertainty along e_2 and e_3

+ Tissue-dependent differences in FA uncertainty



Next question for doing tractography:

***How do we take into account
noise/uncertainty during tracking?***

Probabilistic Tractography

- We know that estimates of DTI ellipsoids are not exactly representing tracts/bundles
 - Size scale differences between voxel/tracts, multiple tracts, complex structure, signal noise, eddy currents, nonlinear fits, etc.
- How to include errors/uncertainty in interpretation and usage?

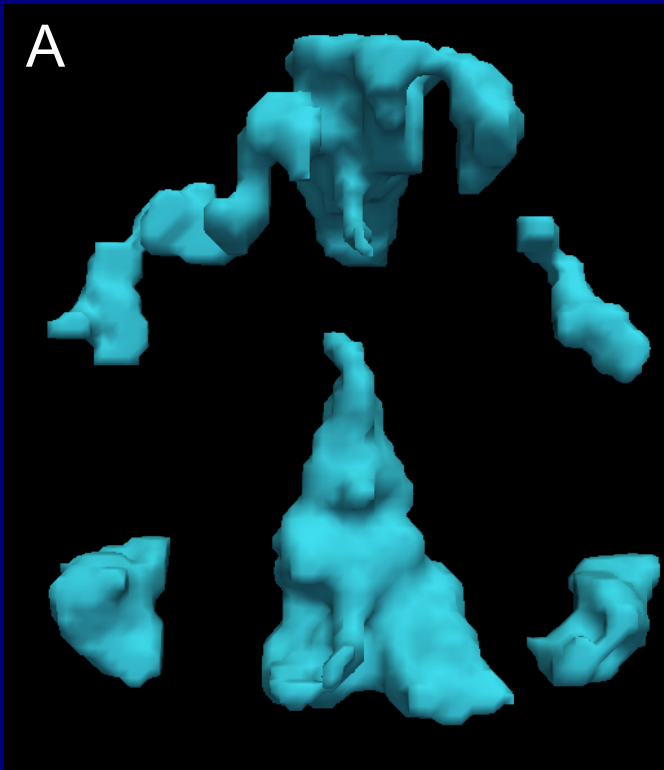
Probabilistic Tractography

- We know that estimates of DTI ellipsoids are not exactly representing tracts/bundles
 - Size scale differences between voxel/tracts, multiple tracts, complex structure, signal noise, eddy currents, nonlinear fits, etc.
- How to include errors/uncertainty in interpretation and usage?
- Probabilistic tractography: use uncertainty in ellipsoid measures with Monte Carlo-esque simulations and build up large ~population of possible trajectories
 - E.g., Parker et al. (2003); Behrens et al. (2003)
 - Do DTI estimates; do whole brain tractography; keep track of number of tracks through relevant voxels; perturb DTI voxel estimates based on uncertainty values; do whole brain tract... [repeat many ~1000 times] ... find voxels which had lots of traffic, define relative 'connectivity' based on traffic

(Side note before continuing with
'full' probabilistic tracking)

Mini-Probabilistic Tracking

- + Full probabilistic methods generate voxelwise brain maps without linear track structure
- + 'Mini-probabilistic' tracking performs a few extra iterations of deterministic tracking on uncertainty-perturbed data sets
 - track structure is retained,
 - results generally exhibit more robust tracks and fewer false negatives than deterministic tracking alone
 - false positives tend to be isolated and visually apparent.



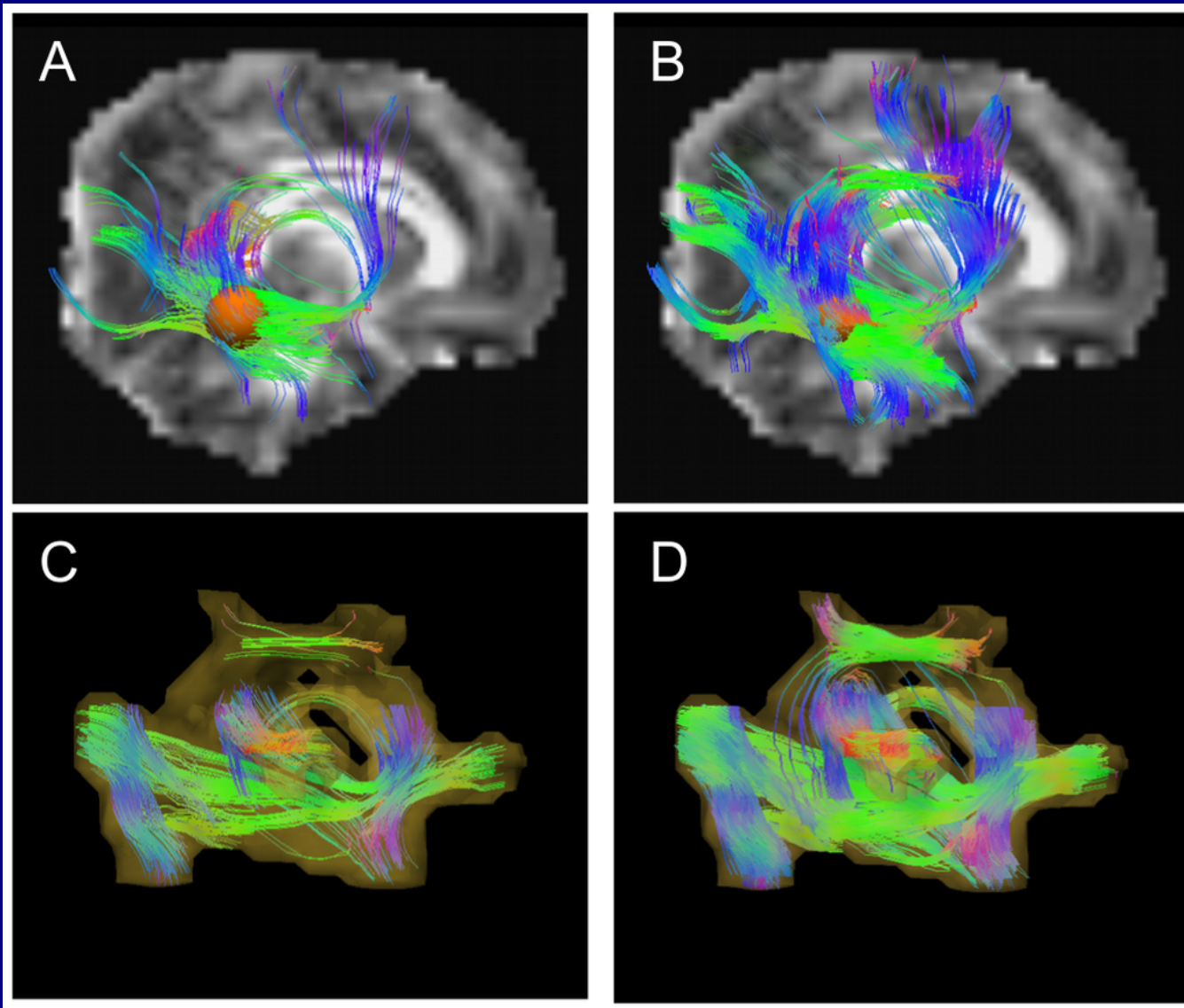
Deterministic (AND)

with '-mini_prob 7'

Mini-Probabilistic Tracking

Deterministic vs mini-Probabilistic

Through
single ROI



AND logic
through
network, cf
with full-prob
results

(Taylor et al., 2014)

Probabilistic Tractography

- Note on interpretation: most reports define a parameter to be the probability of connection between voxels A and X:
 $\Psi(X,A)=\mu(X,A)/N$
 - **N**: number of iterations
 - **μ** : number of tracts through voxel X which either start from or pass through A

Probabilistic Tractography

- Note on interpretation: most reports define a parameter to be the probability of connection between voxels A and X:
$$\Psi(X,A)=\mu(X,A)/N$$
 - N: number of iterations
 - μ : number of tracts through voxel X which either start from or pass through A
- While this quantity is somehow relevant in representing what relative 'connectivity' which can be estimated, exact interpretation as 'probability of connectivity' is tricky

Probabilistic Tractography

- Note on interpretation: most reports define a parameter to be the probability of connection between voxels A and X:
$$\Psi(X,A)=\mu(X,A)/N$$
 - N: number of iterations
 - μ : number of tracts through voxel X which either start from or pass through A
- While this quantity is somehow relevant in representing what relative 'connectivity' which can be estimated, exact interpretation as 'probability of connectivity' is tricky
 - > for example, how literally can one equate a numerically-constructed tract through a ~2x2x2mm voxel with a fiber bundle with **orders-of-magnitude** smaller diameter?
 - > or how can one compare this 'connectivity' between **ROIs of different sizes** on equal footing?

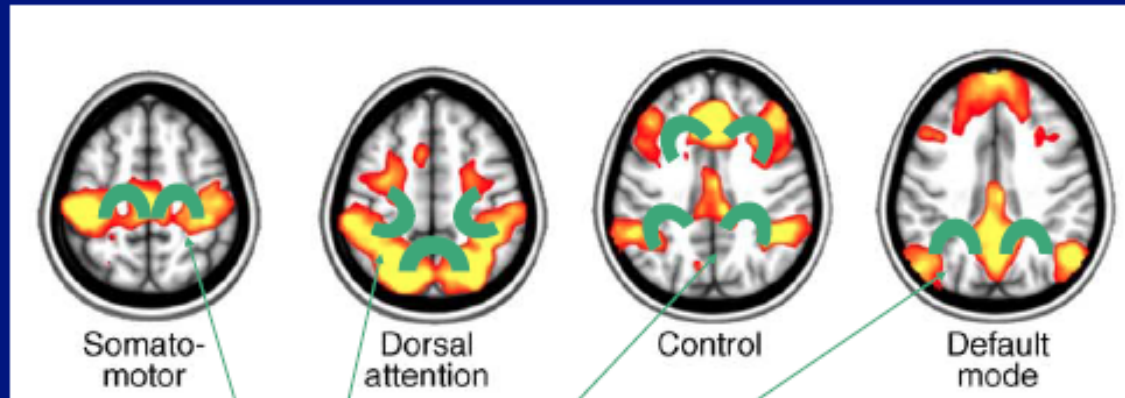
Probabilistic Tractography

- Note on interpretation: most reports define a parameter to be the probability of connection between voxels A and X:
$$\Psi(X,A)=\mu(X,A)/N$$
 - N: number of iterations
 - μ : number of tracts through voxel X which either start from or pass through A
- While this quantity is somehow relevant in representing what relative 'connectivity' which can be estimated, exact interpretation as 'probability of connectivity' is tricky
- Prefer to think of Ψ more loosely as a probability of that voxel being a part of WM volume related to the two ROI-voxels.
 - Not probability of *connectivity* of A and X, but more *likelihood of a voxel being part of associated WM*

Probabilistic Tractography

- This interpretation more useful for working with GM networks. Recall interest:

GM ROIs network:

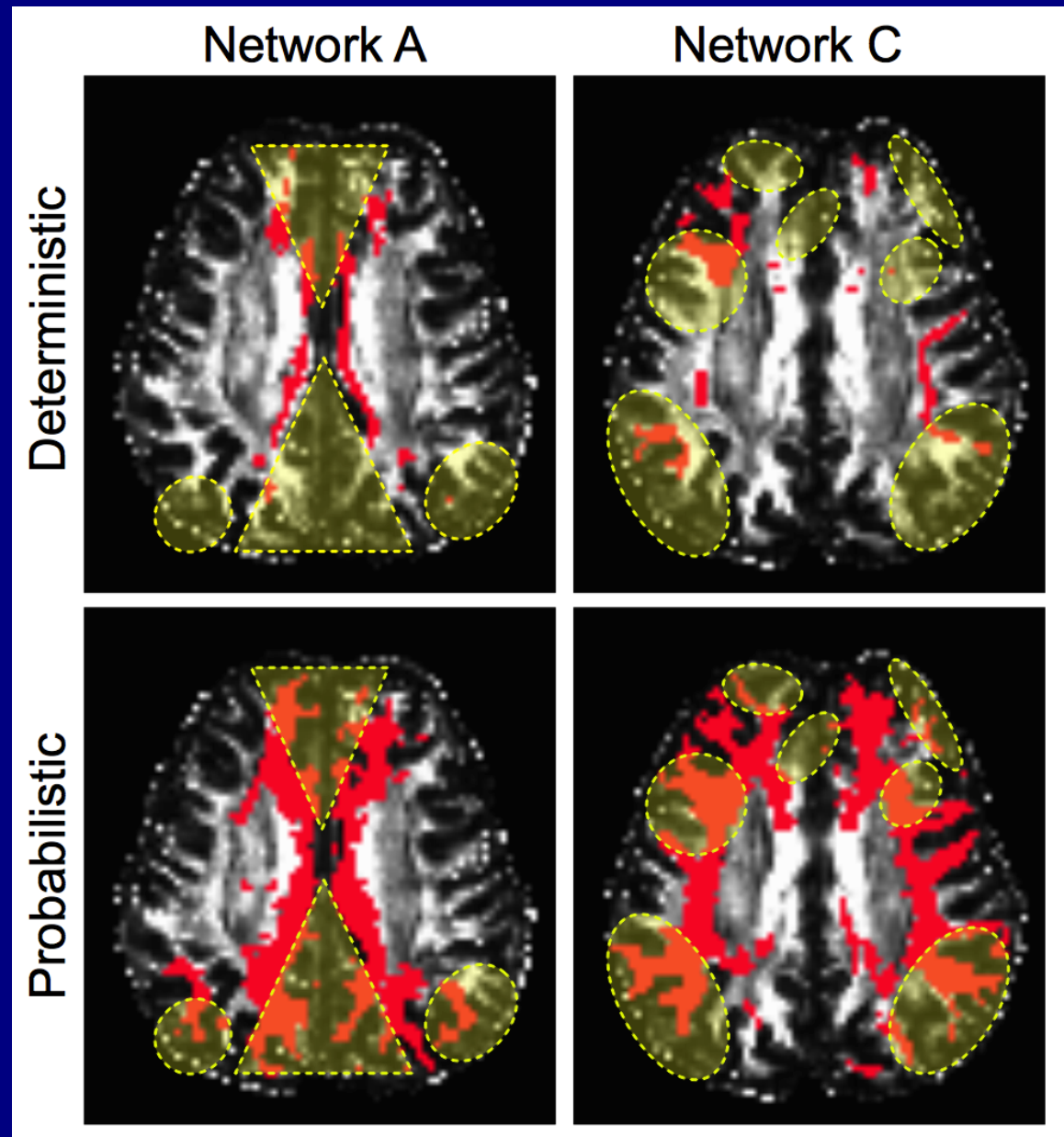


Associated WM ROIs

- Threshold Ψ per voxel after probabilistic tracking, use to define WM ROI between GM ROIs

Deterministic vs Probabilistic

- + NB: coverage and connectivity differences between tractography types
- + Deterministic can be useful for initial investigations, but is more susceptible to noise/errors and truncation



Probabilistic tractography

- + with networks of ROIs from **3dROIMaker** and uncertainty from **3dDWUncert** (as well as tensor estimates from, e.g., 3dDWItoDT), can finally do probabilistic tractography
- + **3dTrackID -mode PROB**
 - does lots of **Monte Carlo simulations**: wholebrain tractography -> perturb FA & e1 based on uncertainty -> wholebrain tracking -> perturb -> wholebrain tracking -> etc.

Probabilistic tractography

- + with networks of ROIs from **3dROIMaker** and uncertainty from **3dDWUncert** (as well as tensor estimates from, e.g., 3dDWItoDT), can finally do probabilistic tractography
- + **3dTrackID -mode PROB**
 - does lots of Monte Carlo simulations: wholebrain tractography -> perturb FA & e1 based on uncertainty -> wholebrain tracking -> perturb -> wholebrain tracking -> etc.
 - at each iteration, checks for **connections** between any pair of ROIs
 - can **trim** saved tracts to only keep voxels *between* 2 ROIs (i.e., no overrunners in the 'connection' ROIs)

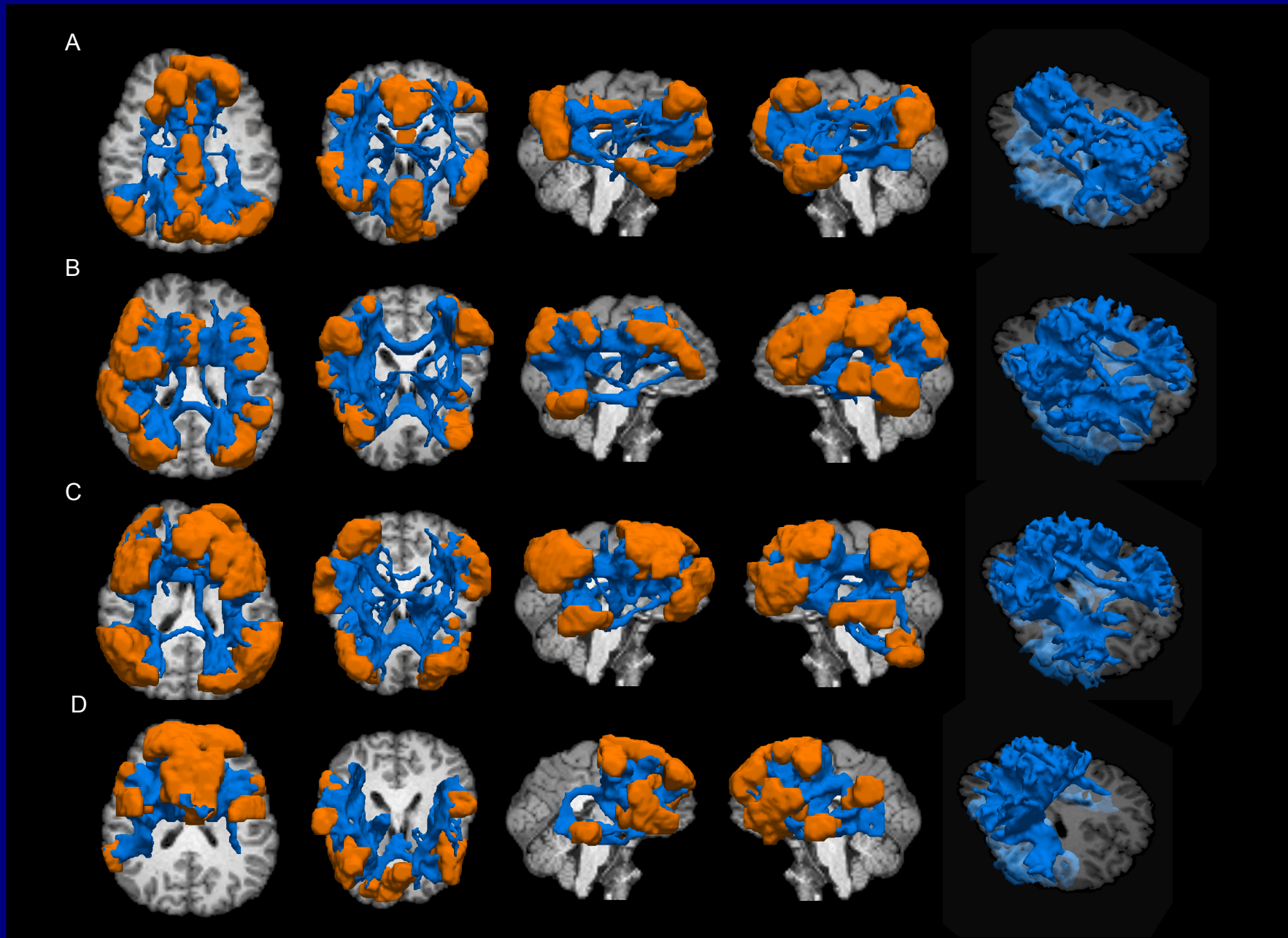
Probabilistic tractography

- + with networks of ROIs from **3dROIMaker** and uncertainty from **3dDWUncert** (as well as tensor estimates from, e.g., 3dDWItoDT), can finally do probabilistic tractography
- + **3dTrackID -mode PROB**
 - does lots of Monte Carlo simulations: wholebrain tractography -> perturb FA & e1 based on uncertainty -> wholebrain tracking -> perturb -> wholebrain tracking -> etc.
 - at each iteration, checks for connections between any pair of ROIs
 - can trim saved tracts to only keep voxels *between* 2 ROIs (i.e., no overrunners in the 'connection' ROIs)
 - also finds tracts through each individual ROI
 - to find WM region connecting, say, ROI 1 and 2: keep voxels through which Ntracks which intersected both ROI1 and ROI2 is greater than a user-defined threshold

Probabilistic tractography

- + with networks of ROIs from **3dROIMaker** and uncertainty from **3dDWUncert** (as well as tensor estimates from, e.g., 3dDWItoDT), can finally do probabilistic tractography
- + **3dTrackID -mode PROB**
 - does lots of Monte Carlo simulations: wholebrain tractography -> perturb FA & e1 based on uncertainty -> wholebrain tracking -> perturb -> wholebrain tracking -> etc.
 - at each iteration, checks for connections between any pair of ROIs
 - can trim saved tracts to only keep voxels *between* 2 ROIs (i.e., no overrunners in the 'connection' ROIs)
 - also finds tracts through each individual ROI
 - to find WM region connecting, say, ROI 1 and 2:
 - keep voxels through which Ntracks which intersected both ROI1 and ROI2 is greater than a user-defined threshold
 - calculate stats on final WM ROIs found
 - analyze multiple networks **simultaneously** for efficiency (i.e., very little extra cost)

3dTrackID: Probabilistic tractography

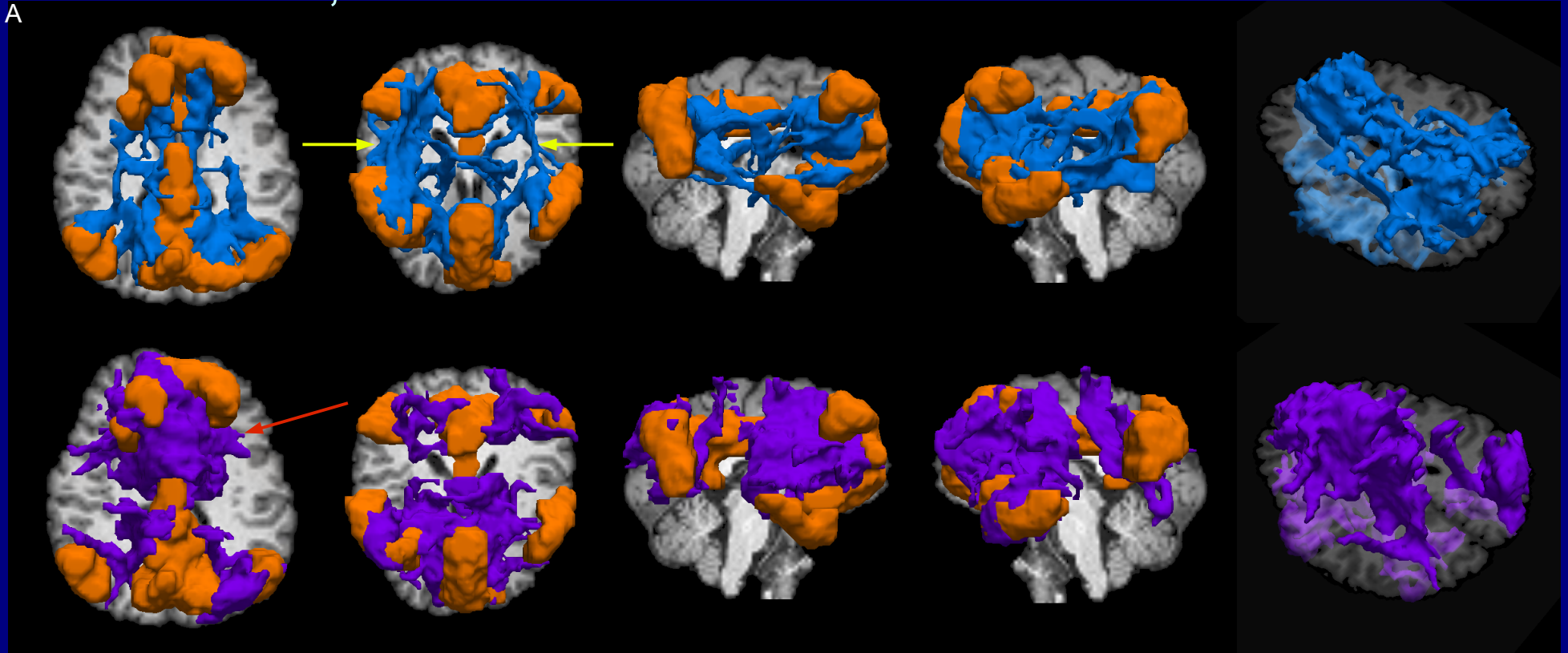


(orange is ROI; blue is set of WM regions with tracts connecting)

3dTrackID: Probabilistic tractography

+ compare with existing algorithms:

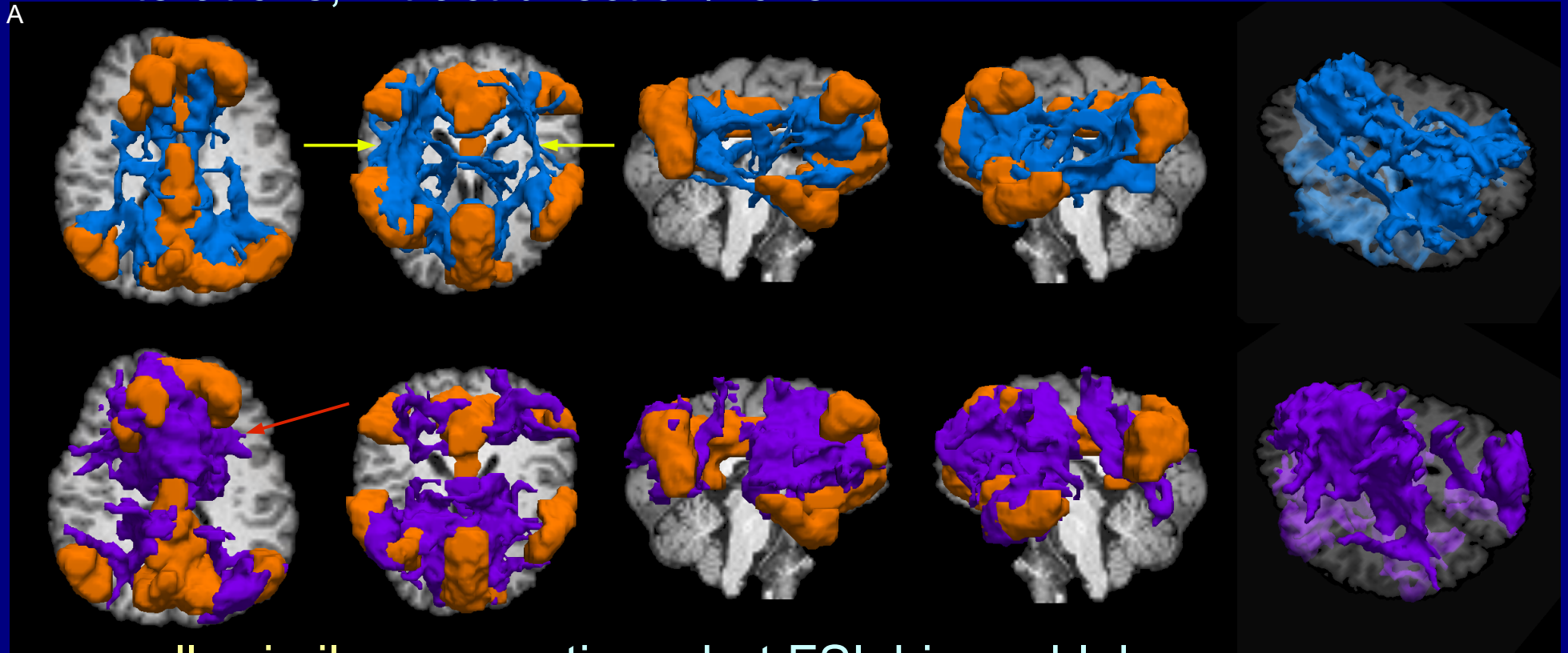
- purple: FSL-probtrackX (and FSL-bedpostX for uncertainty)
- same parameters: $FA > 0.2$, max angle 60deg, 5000 Monte Carlo iterations; 1 tract direction/voxel



3dTrackID: Probabilistic tractography

+ compare with existing algorithms:

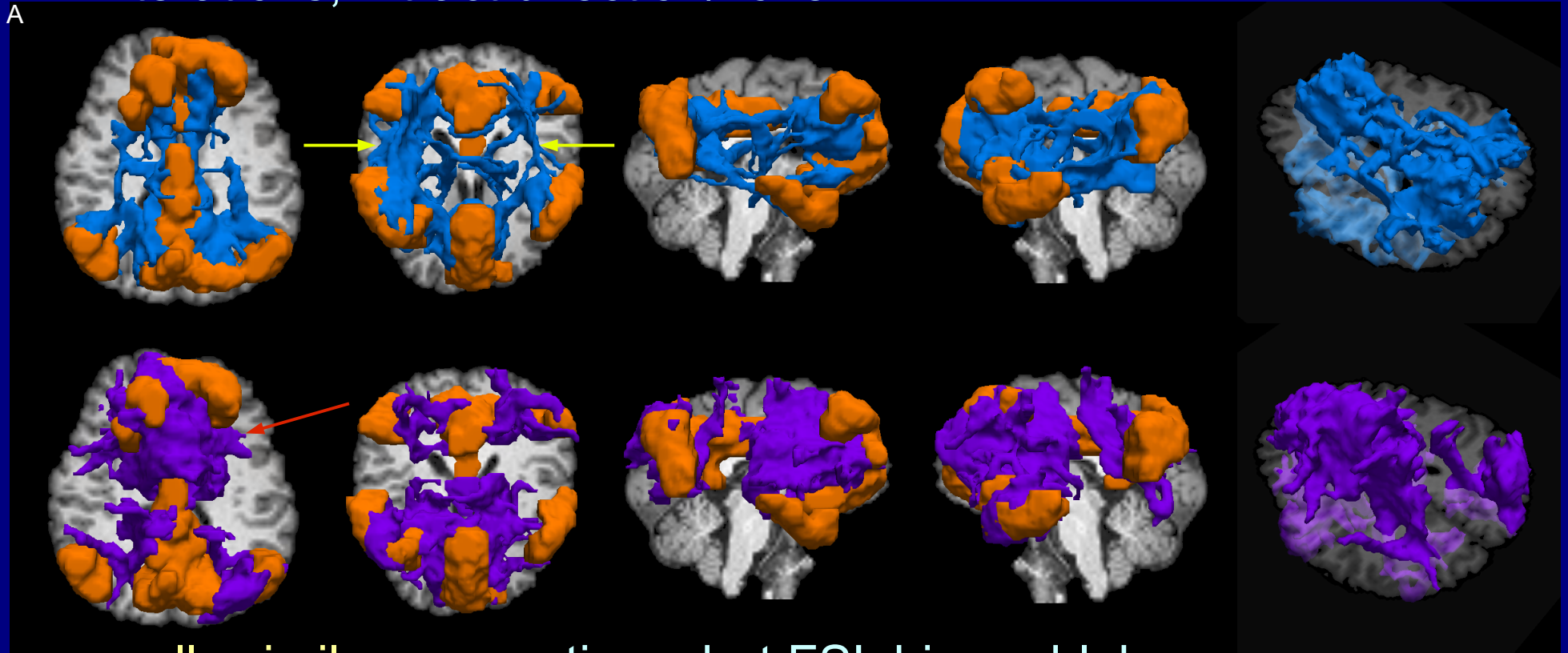
- purple: FSL-probtrackX (and FSL-bedpostX for uncertainty)
- same parameters: $FA > 0.2$, max angle 60deg, 5000 Monte Carlo iterations; 1 tract direction/voxel



+ generally similar connections, but FSL bigger blobs

3dTrackID: Probabilistic tractography

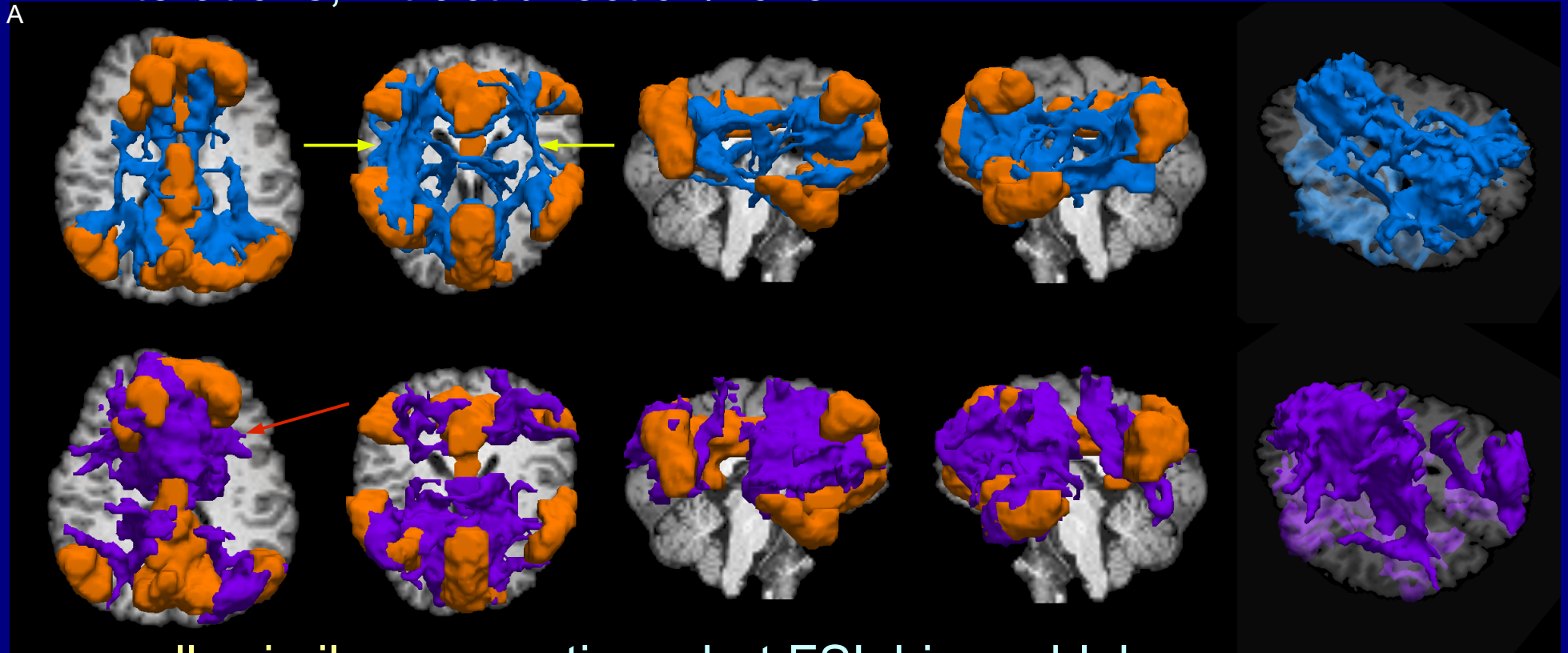
- + compare with existing algorithms:
 - purple: FSL-probtrackX (and FSL-bedpostX for uncertainty)
 - same parameters: $FA > 0.2$, max angle 60deg, 5000 Monte Carlo iterations; 1 tract direction/voxel



- + generally similar connections, but FSL bigger blobs
- + FSL took **several hours** for uncertainty, and then **>24 hours** for tracking this single network (and had to run 4 for this study)

3dTrackID: Probabilistic tractography

- + compare with existing algorithms:
 - purple: FSL-probtrackX (and FSL-bedpostX for uncertainty)
 - same parameters: $FA > 0.2$, max angle 60deg, 5000 Monte Carlo iterations; 1 tract direction/voxel

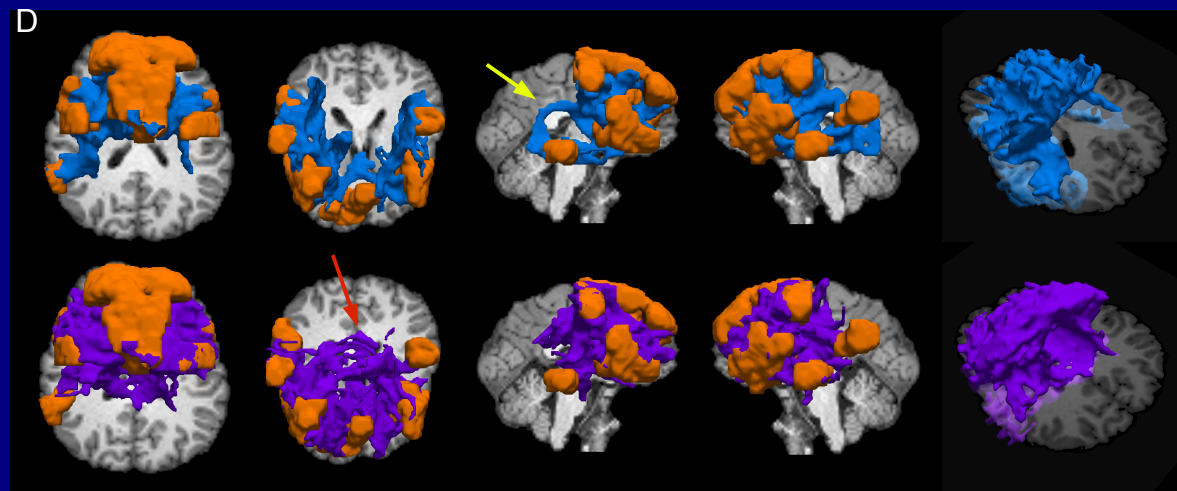
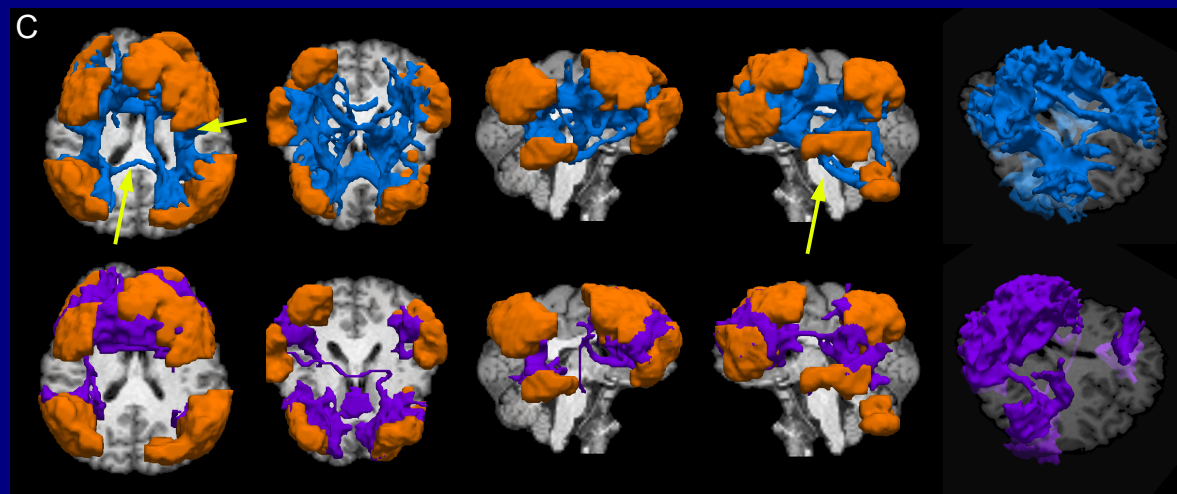
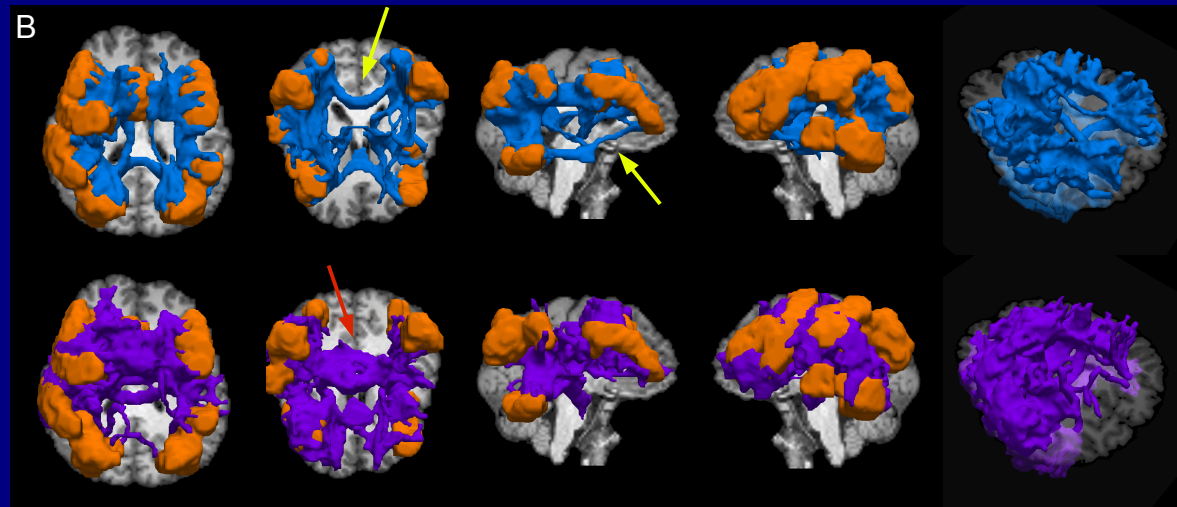


- + generally similar connections, but FSL bigger blobs
- + FSL took **several hours** for uncertainty, and then **>24 hours** for tracking this single network (and had to run 4 for this study)
- + **3dDWUncert** took **7min**; **3dTrackID** took **25mins** total for 4 netw.

3dTrackID:

(other networks show similar results in terms of:

- narrow/wide regions of tracts;
- broadly similar locations;
- each program shows some tracks which the other doesn't)

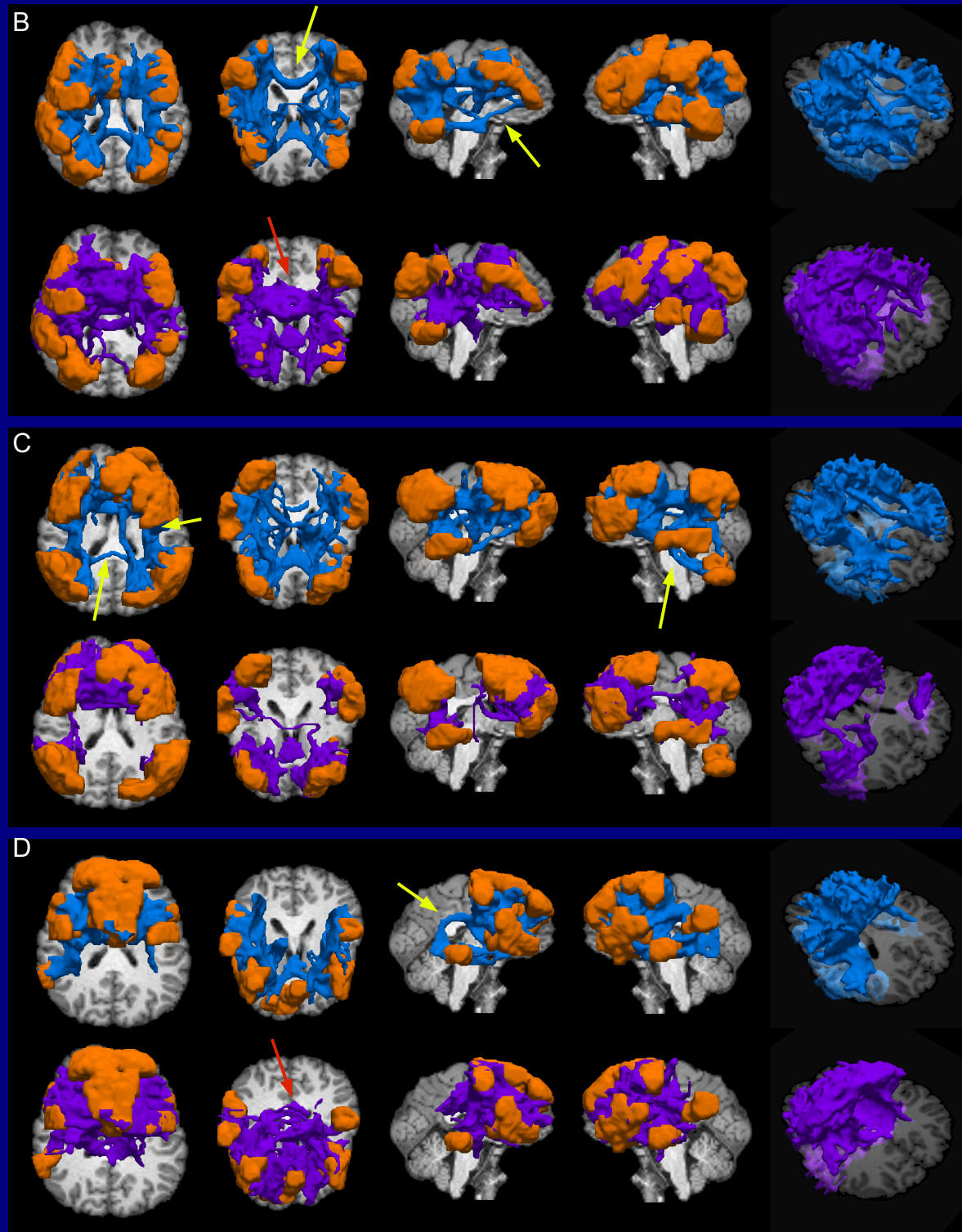


3dTrackID:

(other networks show similar results in terms of:

- narrow/wide regions of tracts;
- broadly similar locations;
- each program shows some tracks which the other doesn't)

(3dTrackID automatically creates *.grid files for probabilistic files, as well.)

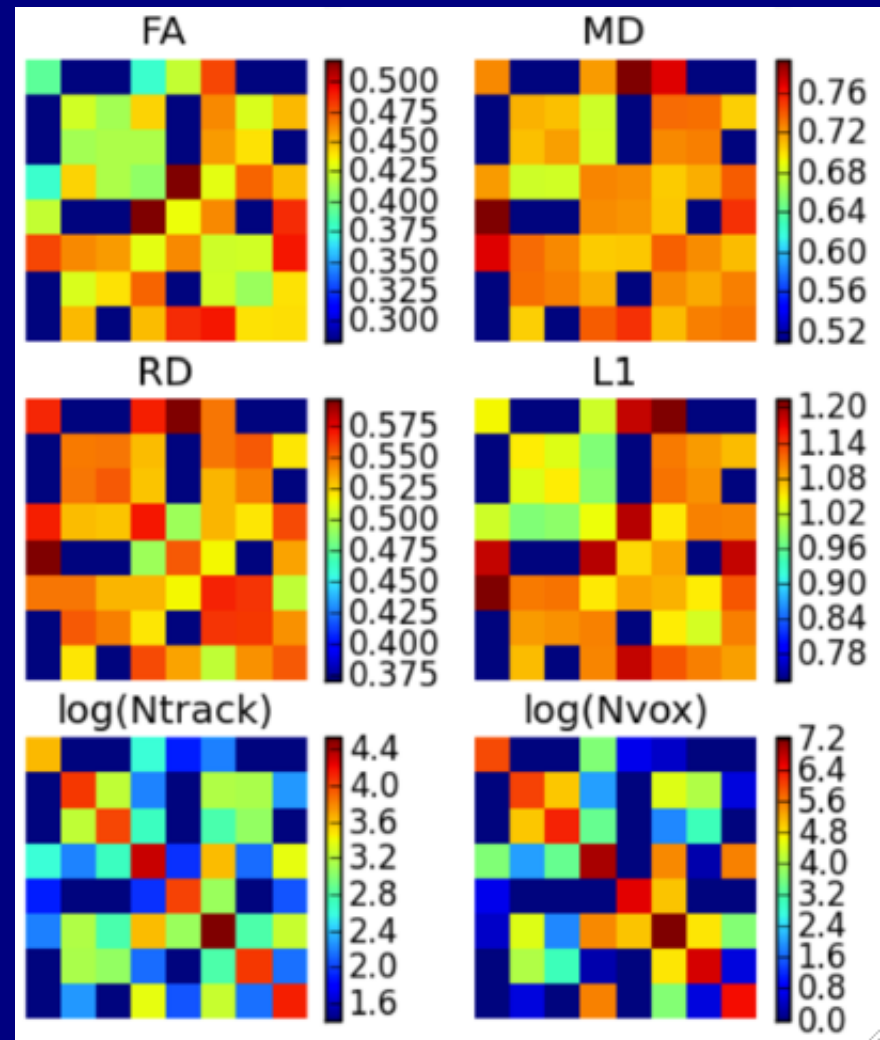


WM (ROI) Quantities

For connected pairs of GM ROIs in a network, have an average WM property (or can map to T1, PD...) →

Have produced sets of localized structural/anatomical quantities for comparison with functional values or behavioral scores, genetics, etc.

Can use for group or individual comparisons/regressions.



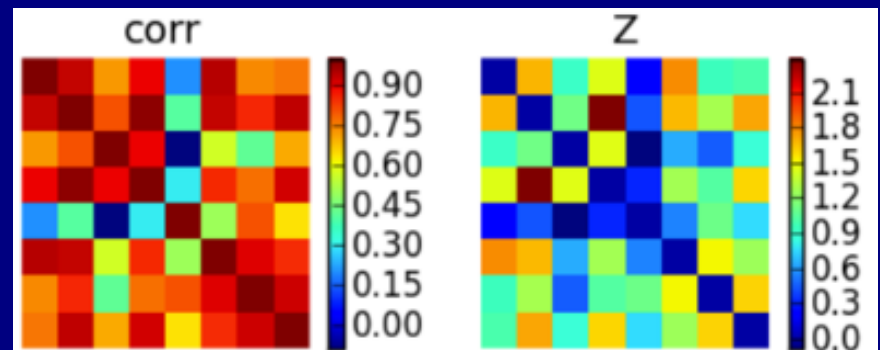
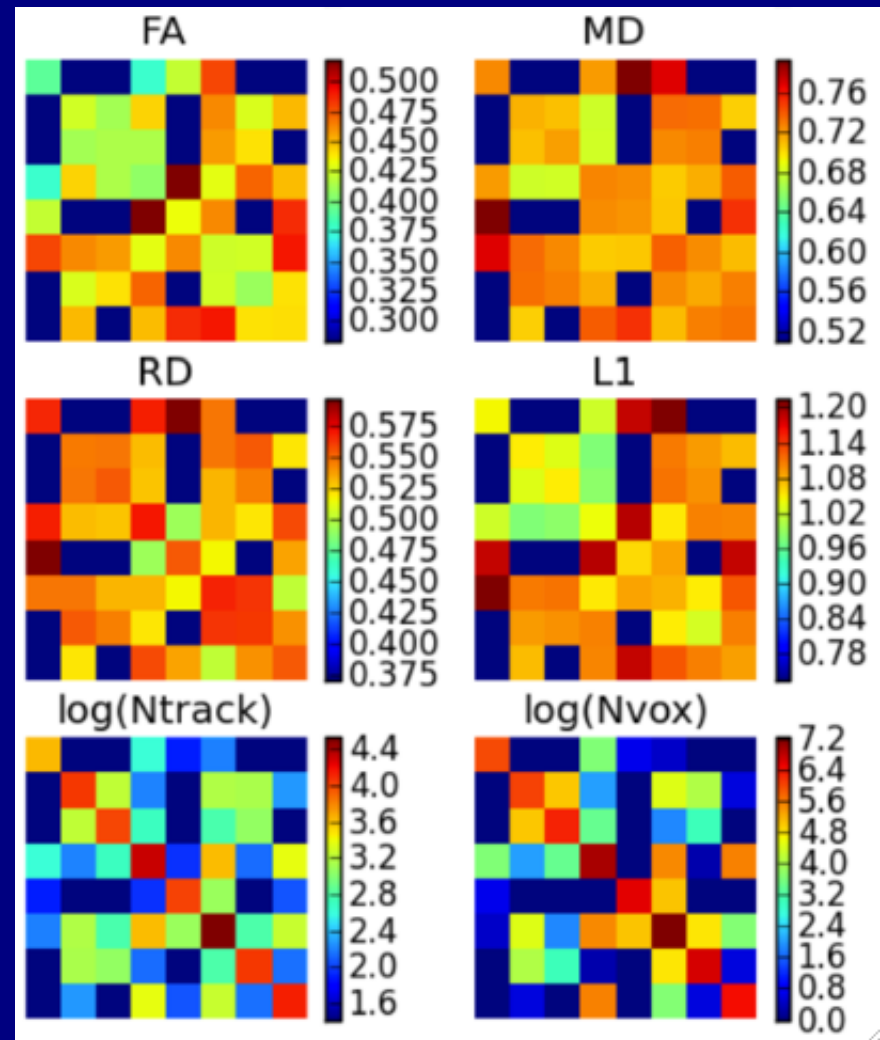
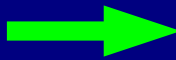
WM (ROI) Quantities

For connected pairs of GM ROIs in a network, have an average WM property (or can map to T1, PD...) →

Have produced sets of localized structural/anatomical quantities for comparison with functional values or behavioral scores, genetics, etc.

Can use for group or individual comparisons/regressions.

3dNetCorr: correlation matrices
Of average time series in ROIs
(e.g., uninflated GM ROIs from 3dROIMaker)



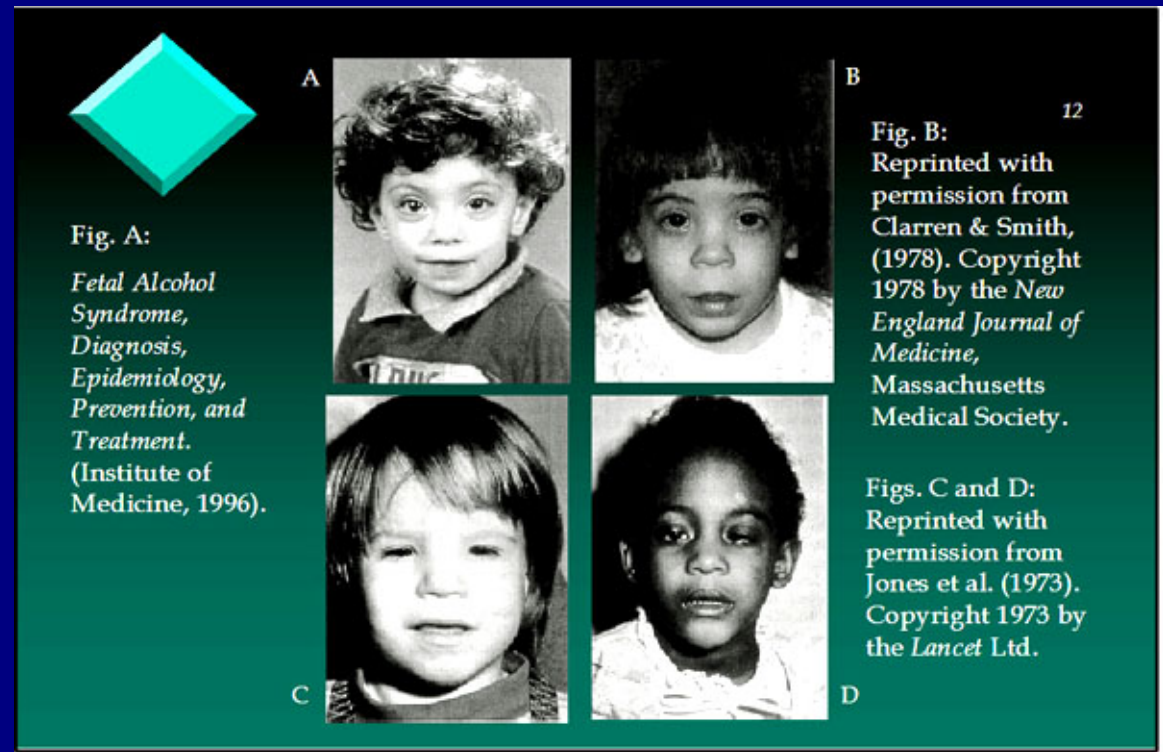
Example:
Group analysis with tracking output
using multivariate statistics

from study:

*A DTI-Based Tractography Study of Effects
on Brain Structure Associated with
Prenatal Alcohol Exposure in Newborns,*
*Taylor, Jacobson, van der Kouwe, Molteno, Chen,
Wintermark, Alhamud, Jacobson, Meintjes (2014)*

Prenatal alcohol exposure (PAE)

- Alcohol is a teratogen, disrupting healthy embryonic and fetal development.
 - leads to various **Fetal Alcohol Spectrum Disorders (FASD)**
- FASD occurs in children whose pregnant mothers binge drank
 - e.g., **≥4 drinks/occasion and/or ≥14 drinks/wk**
- Results in *poor*:
 - academic performance
 - language/math skills
 - impulse control
 - abstract reasoning
 - memory, attention and facial and skeletal dysmorphology



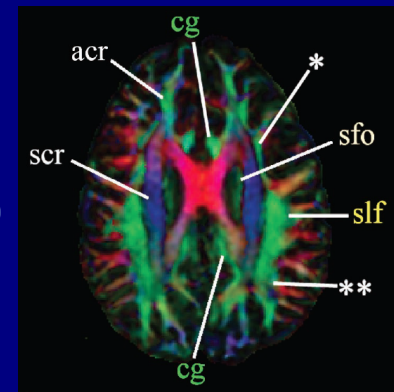
Goals of this study

To:

- 1) Use neuroimaging to compare structural brain development in newborns with PAE to that of HC newborns.
- 2) Quantitatively examine WM properties across the brain
- 3) Relate changes in (localized) WM properties with PAE, controlling for several confounding effects
→ examine several, and see which is/are (most) significant

Tools: diffusion tensor imaging (DTI) + tractography

- A) delineate similar WM ROIs across all subjects
- B) quantify structural properties (FA, MD, T1, ...)
- C) statistical modeling for comparisons
- *at whole brain, network and ROI levels*



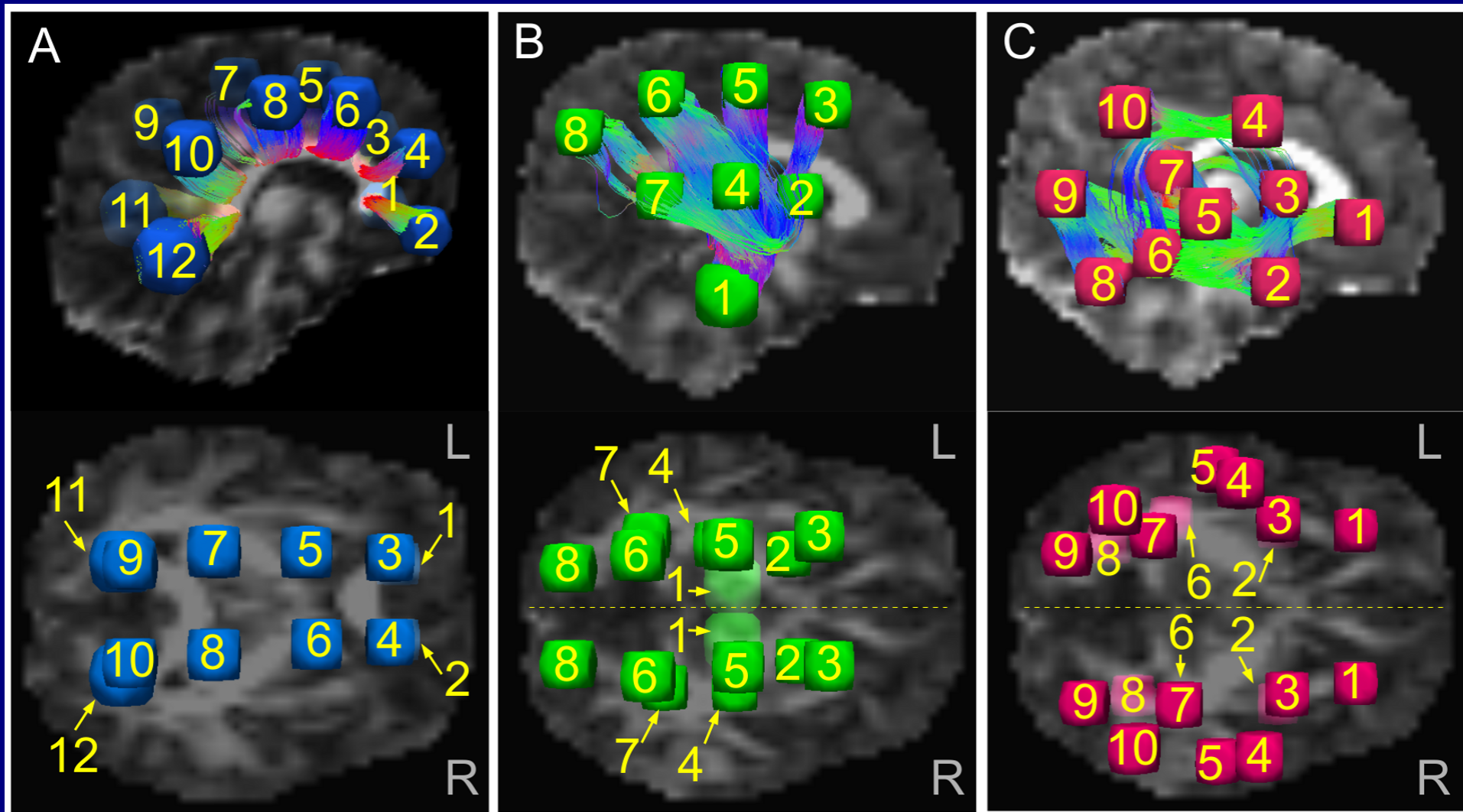
Setting up DTI-tractography

Location of targets for tractography: 5 WM networks.

CC and Cor. Rad.
(CCCR)

Projection
(L/R-PROJ)

Association
(L/R-ASSOC)



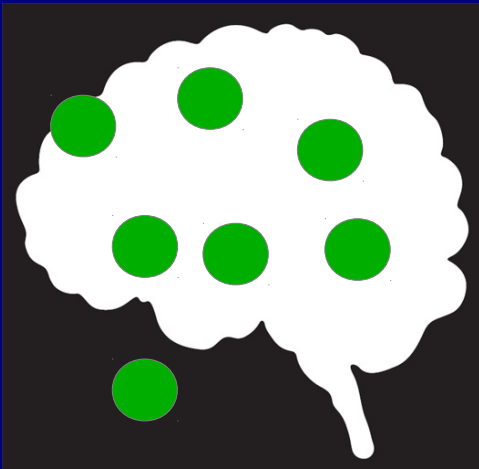
Analysis Steps

1) Place network targets

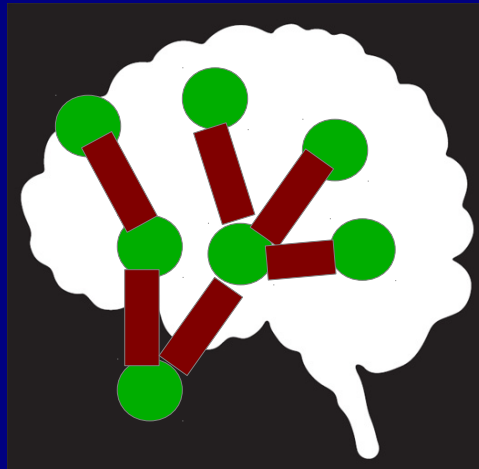


Analysis Steps

1) Place network targets

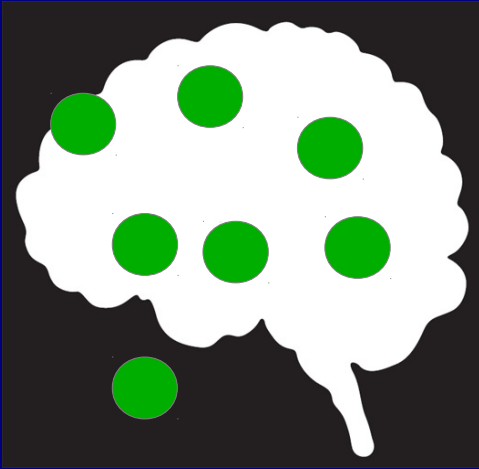


2) Probabilistic tracking

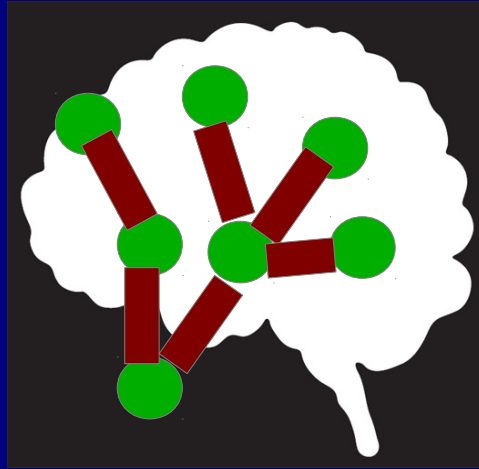


Analysis Steps

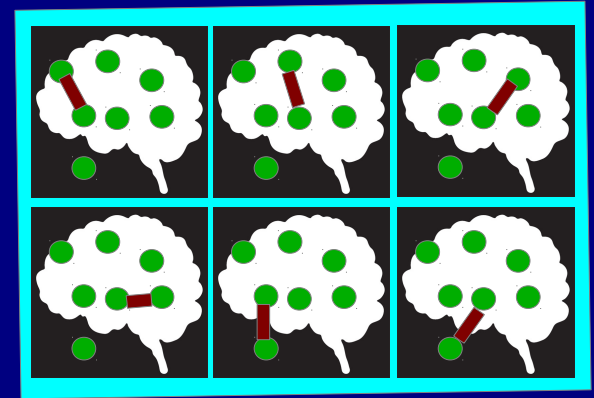
1) Place network targets



2) Probabilistic tracking

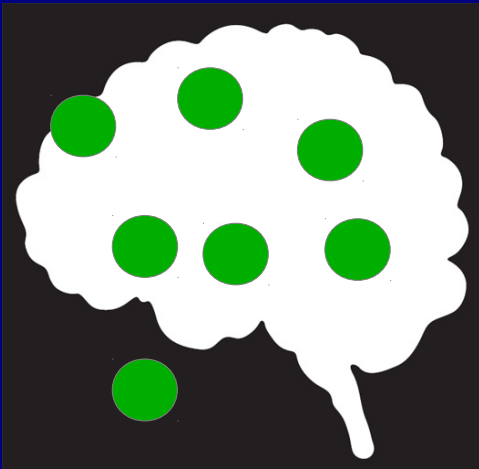


3) set of WM ROIs \rightarrow set of repeated measures

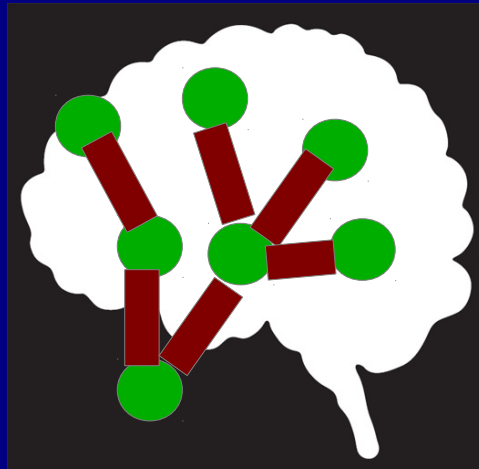


Analysis Steps

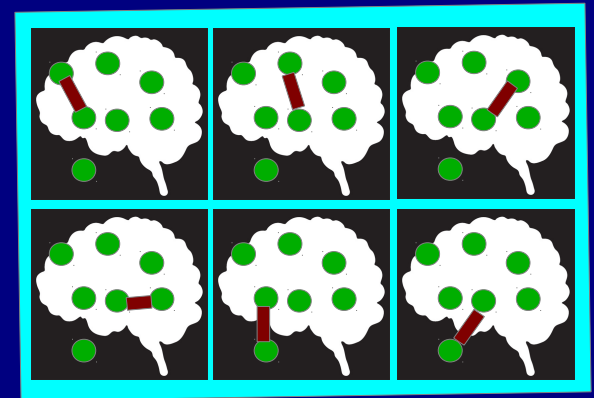
1) Place network targets



2) Probabilistic tracking



3) set of WM ROIs → set of repeated measures



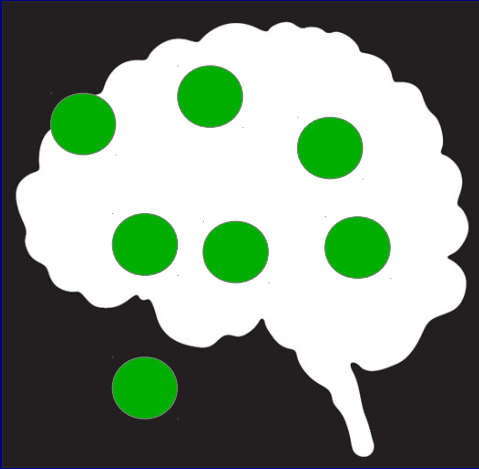
4) Multivariate model

- $\{FA_1, FA_2, FA_3, \dots\}$
- alc
- infant age
- infant sex
- maternal age
- maternal cig/day

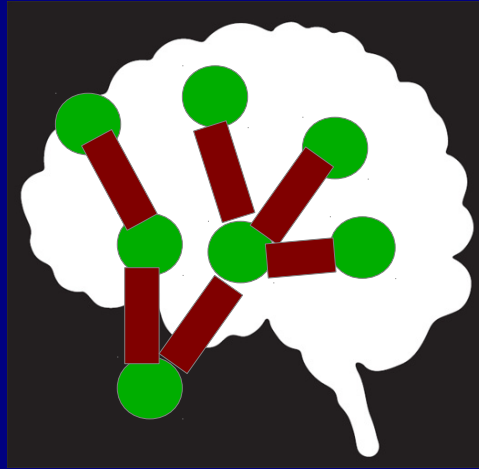
➔ AFNI's 3dMVM, written by G. Chen

Analysis Steps

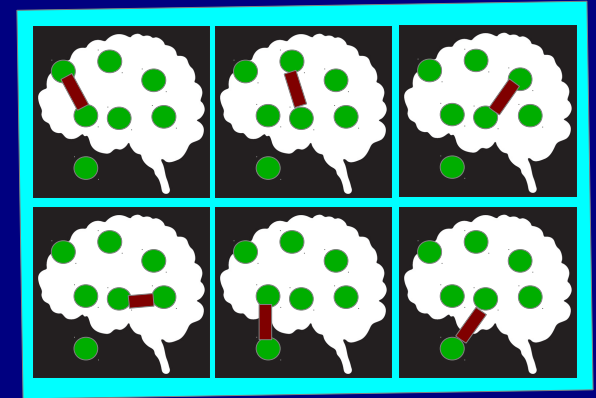
1) Place network targets



2) Probabilistic tracking



3) set of WM ROIs → set of repeated measures

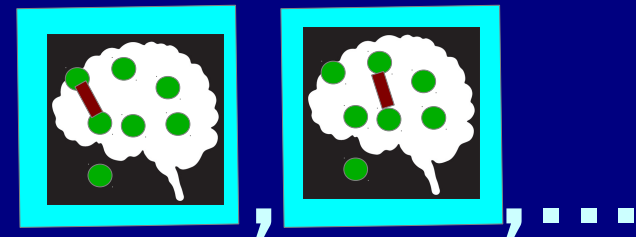


4) Multivariate model

- $\{FA_1, FA_2, FA_3, \dots\}$
- alc
- infant age
- infant sex
- maternal age
- maternal cig/day

5) Follow-up GLM for each WM ROI

- FA
- alc
- infant age
- infant sex
- maternal age
- maternal cig/day



➔ AFNI's 3dMVM, written by G. Chen

In Summary

- + Have motivated ways of combining FC and SC analyses
 - fMRI to define networks of GM ROIs
 - find locations of connections within/across networks -> WM ROIs
 - calculate stats of DTI/anatomical properties there
 - combine structural quantities of, e.g., mean FA, with fMRI connectivity matrices; behavioral measures; genetic values, etc.
- + Diffusion-based tractography is useful complement to fMRI
 - probabilistic tractography is more robust than deterministic
 - different types of quantities than fMRI, not necessarily 'strengths'
- + Still room to improve, tools to add.
 - Suggestions are quite welcome!

Analysis Steps

fat_mvm_prep.py

- + make a data table combining:
 - a CSV file of subject data with
 - a set of *.grid¹ files from 3dTrackID;
- + automatically selects tracked connections found across all groups (future version may have LME modeling that allows missing data)

fat_mvm_scripter.py

- + define a statistical model of variables from CSV file + DTI data
- + build a 3dMVM script to test the model using entire networks, and
- + construct follow-up GLTs to investigate individual regions.

¹Also works with *.netcc files from 3dNetCorr.

II) Results: network level

The questions:

- 1) which WM networks are affected by PAE?
- 2) which parameters show effects most strongly?

Answer using:

- (for each network) a multivariate GLM for
 - set of DTI parameters
 - alcohol (frequency: binge/wk)
 - infant age (wks since conception)
 - infant sex (M/F)
 - maternal age (yrs)
 - maternal cigarette smoking (cig/day).

II) Results: network level

The questions:

- 1) which WM networks are affected by PAE?
- 2) which parameters show effects most strongly?

Parameters showing at least trends ($p < 0.1$) →

← Networks

Network	FA				MD				AD				PD			
	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p
CCCR					alc	-0.70	8.6 (1, 14)	0.011*	alc	-0.72	14.0 (1, 14)	0.002**	cig	0.47	3.5 (1, 14)	0.083
					mat_age	0.56	5.5 (1, 14)	0.034*	mat_age	0.53	6.3 (1, 14)	0.025*				
L-PROJ	cig	0.12	4.2 (11, 4)	0.091	alc	-0.41	3.9 (10, 140)	0.000***	alc	-0.52	4.1 (10, 140)	0.000***	cig	0.52	4.0 (1, 14)	0.066
					mat_age	0.37	4.4 (1, 14)	0.056	mat_age	0.44	6.5 (1, 14)	0.023*				
R-PROJ					alc	-0.41	1.9 (12, 168)	0.035*	alc	-0.45	2.7 (12, 168)	0.002**	cig	0.48	3.4 (1, 14)	0.085
	age	0.33	8.6 (13, 2)	0.109	age	-0.41	5.8 (1, 14)	0.031*	age	-0.39	5.3 (1, 14)	0.038*				
	mat_age	-0.16	9.2 (13, 2)	0.103	sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
L-ASSOC					alc	-0.65	6.0 (7, 8)	0.011*	alc	-0.66	8.1 (1, 14)	0.013*	cig	0.49	3.6 (1, 14)	0.080
									age	-0.16	2.5 (6, 84)	0.030*				
					mat_age	0.44	3.8 (1, 14)	0.071	mat_age	0.43	4.7 (1, 14)	0.048*				
R-ASSOC	alc	0.23	1.8 (7, 98)	0.090	alc	-0.62	10.2 (1, 14)	0.007**	alc	-0.67	14.1 (1, 14)	0.002**	cig	0.5	3.5 (1, 14)	0.082
									cig	-0.29	3.9 (1, 14)	0.068				

* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$.

II) Results: network level

The questions:

- 1) which WM networks are affected by PAE?
- 2) which parameters show effects most strongly?

Parameters showing at least trends ($p < 0.1$) →

← Networks

Network	FA				MD				AD				PD			
	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p
CCCR					alc	-0.70	8.6 (1, 14)	0.011*	alc	-0.72	14.0 (1, 14)	0.002**	cig	0.47	3.5 (1, 14)	0.083
L-PROJ	cig	0.12	4.2 (11, 4)	0.091	mat_age	0.56	5.5 (1, 14)	0.034*	alc	-0.52	4.1 (10, 140)	0.000***	mat_age	0.53	6.3 (1, 14)	0.025*
R-PROJ					alc	-0.41	3.9 (10, 140)	0.000***	alc	-0.45	2.7 (12, 168)	0.002**	cig	0.52	4.0 (1, 14)	0.066
	age	0.33	8.6 (13, 2)	0.109	mat_age	0.37	4.4 (1, 14)	0.056	mat_age	0.44	6.5 (1, 14)	0.023*				
	mat_age	-0.16	9.2 (13, 2)	0.103	alc	-0.41	1.9 (12, 168)	0.035*	alc	-0.45	2.7 (12, 168)	0.002**	cig	0.48	3.4 (1, 14)	0.085
L-ASSOC					age	-0.41	5.8 (1, 14)	0.031*	age	-0.39	5.3 (1, 14)	0.038*				
					sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
					alc	-0.65	6.0 (7, 8)	0.011*	alc	-0.66	8.1 (1, 14)	0.013*	cig	0.49	3.6 (1, 14)	0.080
					mat_age	0.44	3.8 (1, 14)	0.071	age	-0.16	2.5 (6, 84)	0.030*				
									mat_age	0.43	4.7 (1, 14)	0.048*				
R-ASSOC	alc	0.23	1.8 (7, 98)	0.090	alc	-0.62	10.2 (1, 14)	0.007**	alc	-0.67	14.1 (1, 14)	0.002**	cig	0.5	3.5 (1, 14)	0.082
									cig	-0.29	3.9 (1, 14)	0.068				

* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$.

→ Statistically significant alcohol exposure associations in ~ every WM network

II) Results: network level

The questions:

- 1) which WM networks are affected by PAE?
- 2) which parameters show effects most strongly?

Parameters showing at least trends ($p < 0.1$) →

← Networks

Network	FA				MD				AD				PD			
	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p	var.	β_{med}	$F(df_N, df_D)$	p
CCCR					alc	-0.70	8.6 (1, 14)	0.011*	alc	-0.72	14.0 (1, 14)	0.002**	cig	0.47	3.5 (1, 14)	0.083
					mat_age	0.56	5.5 (1, 14)	0.034*	mat_age	0.53	6.3 (1, 14)	0.025*				
L-PROJ	cig	0.12	4.2 (11, 4)	0.091	alc	-0.41	3.9 (10, 140)	0.000***	alc	-0.52	4.1 (10, 140)	0.000***	cig	0.52	4.0 (1, 14)	0.066
					mat_age	0.37	4.4 (1, 14)	0.056	mat_age	0.44	6.5 (1, 14)	0.023*				
R-PROJ					alc	-0.41	1.9 (12, 168)	0.035*	alc	-0.45	2.7 (12, 168)	0.002**	cig	0.48	3.4 (1, 14)	0.085
	age	0.33	8.6 (13, 2)	0.109	age	-0.41	5.8 (1, 14)	0.031*	age	-0.39	5.3 (1, 14)	0.038*				
	mat_age	-0.16	9.2 (13, 2)	0.103	sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
L-ASSOC					alc	-0.65	6.0 (7, 8)	0.011*	alc	-0.66	8.1 (1, 14)	0.013*	cig	0.49	3.6 (1, 14)	0.080
									age	-0.16	2.5 (6, 84)	0.030*				
					mat_age	0.44	3.8 (1, 14)	0.071	mat_age	0.43	4.7 (1, 14)	0.048*				
R-ASSOC	alc	0.23	1.8 (7, 98)	0.090	alc	-0.62	10.2 (1, 14)	0.007**	alc	-0.67	14.1 (1, 14)	0.002**	cig	0.5	3.5 (1, 14)	0.082
									cig	-0.29	3.9 (1, 14)	0.068				

* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$.

→ Increased alcohol exposure:
decreased AD
(and decreased MD)






III) Results: ROI level

The question:

1) where are most significant AD-alcohol relations in each network?

Answer using:

- (for each ROI) a GLM for
 - single DTI parameter
 - alcohol (frequency: binge/wk)
 - infant age (wks since conception)
 - infant sex (M/F)
 - maternal age (yrs)
 - maternal cigarette smoking (cig/day).

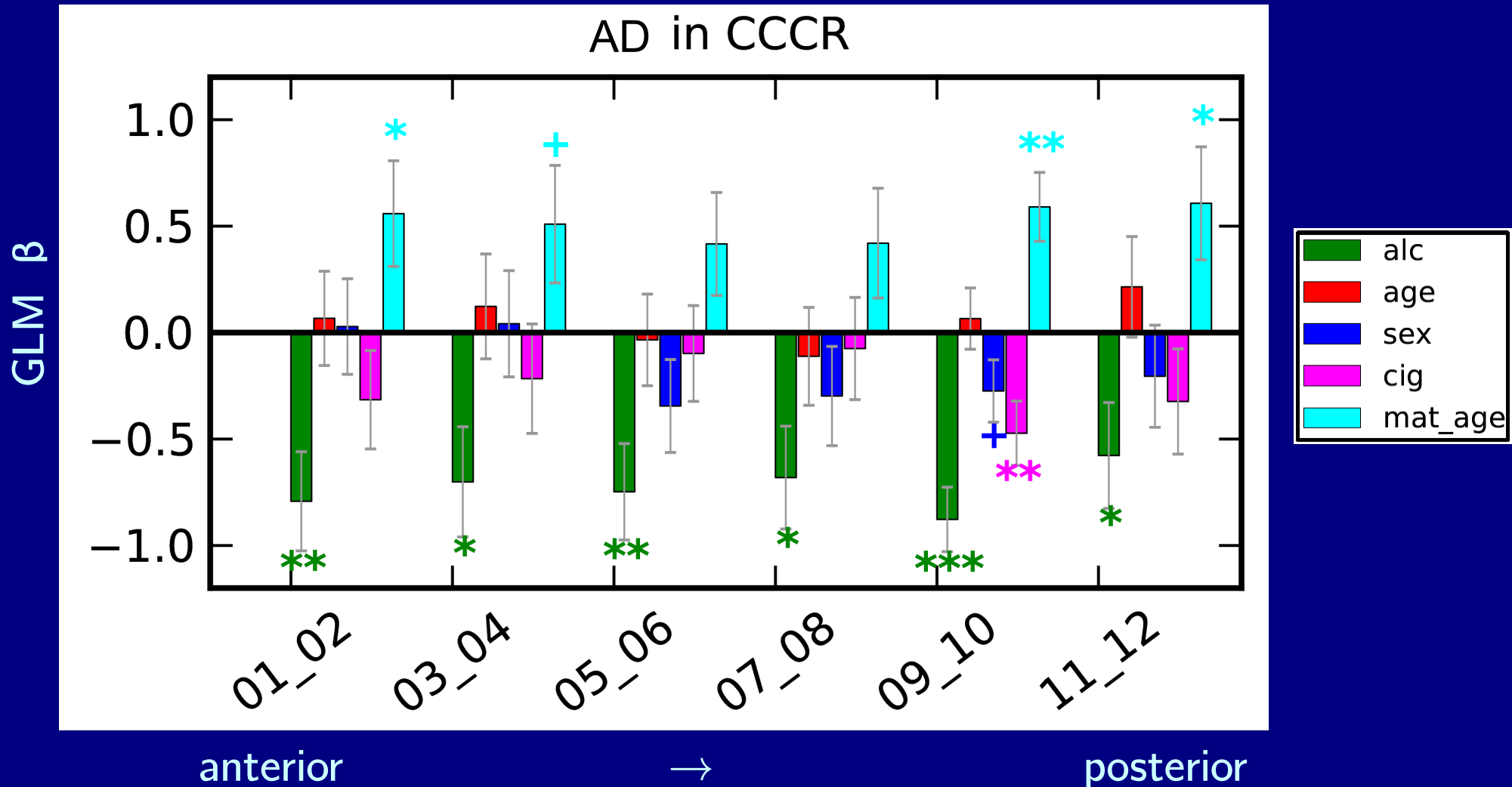
	alc
	age
	sex
	cig
	mat_age

III) Results: ROI level

The question:

1) where are most significant AD-alcohol relations in each network?

Transcallosal (CC and corona radiata)

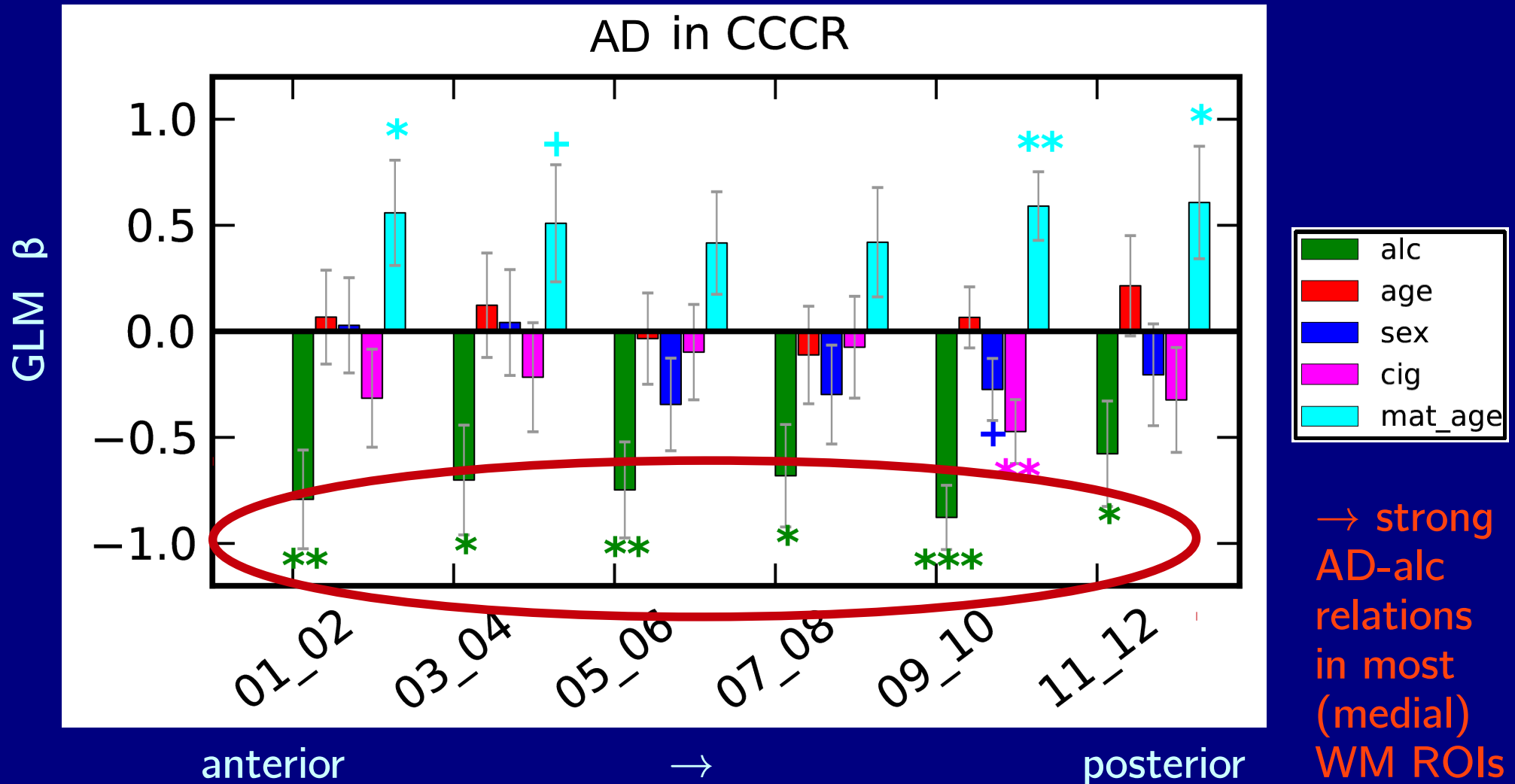


III) Results: ROI level

The question:

1) where are most significant AD-alcohol relations in each network?

Transcallosal (CC and corona radiata)

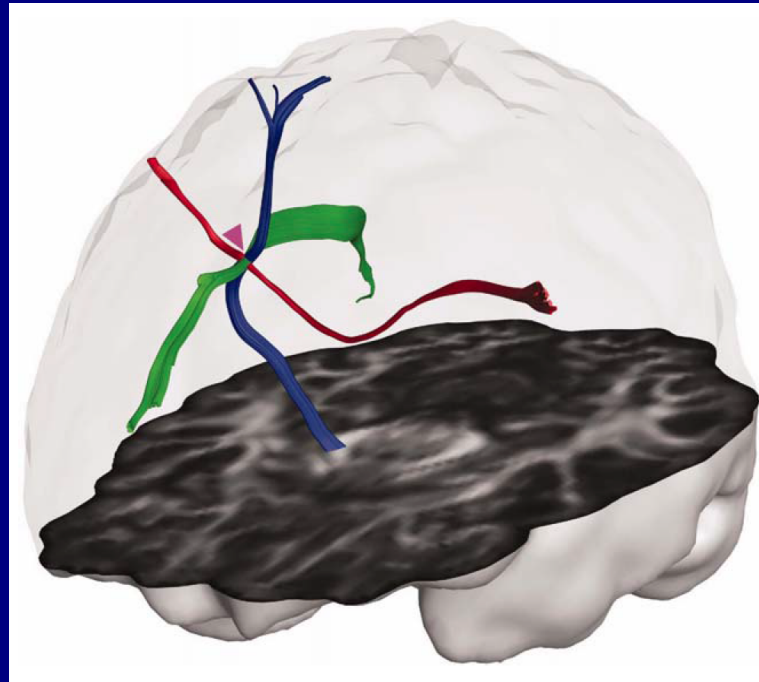


Example:
HARDI tracking

Higher order models

DTI tractography:

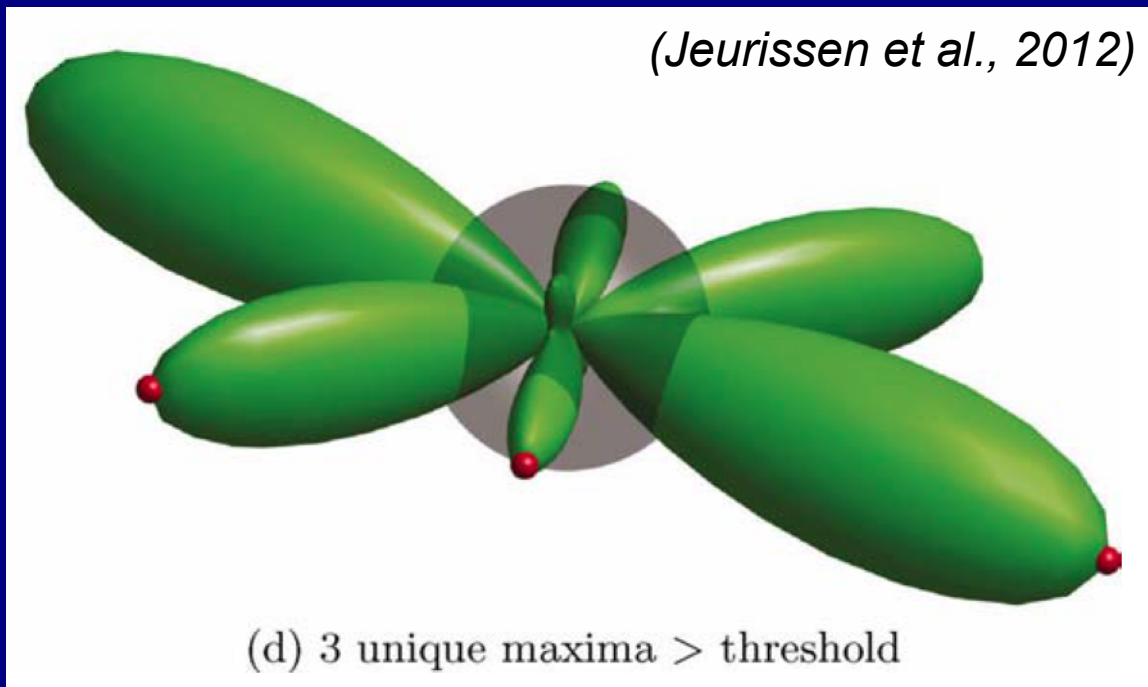
- + susceptible to false negatives, difficulty with long range tracts (noise/error accumulation)
- + Major diffusion can be average of multiple paths
- + Voxels can have low FA from several WM paths, false ending
- + Can't resolve complex underlying architecture
 - Jeurissen et al. (2012, HBM): 60-90% of WM voxels estimated to have multiple fibers



(Jeurissen et al., 2012)

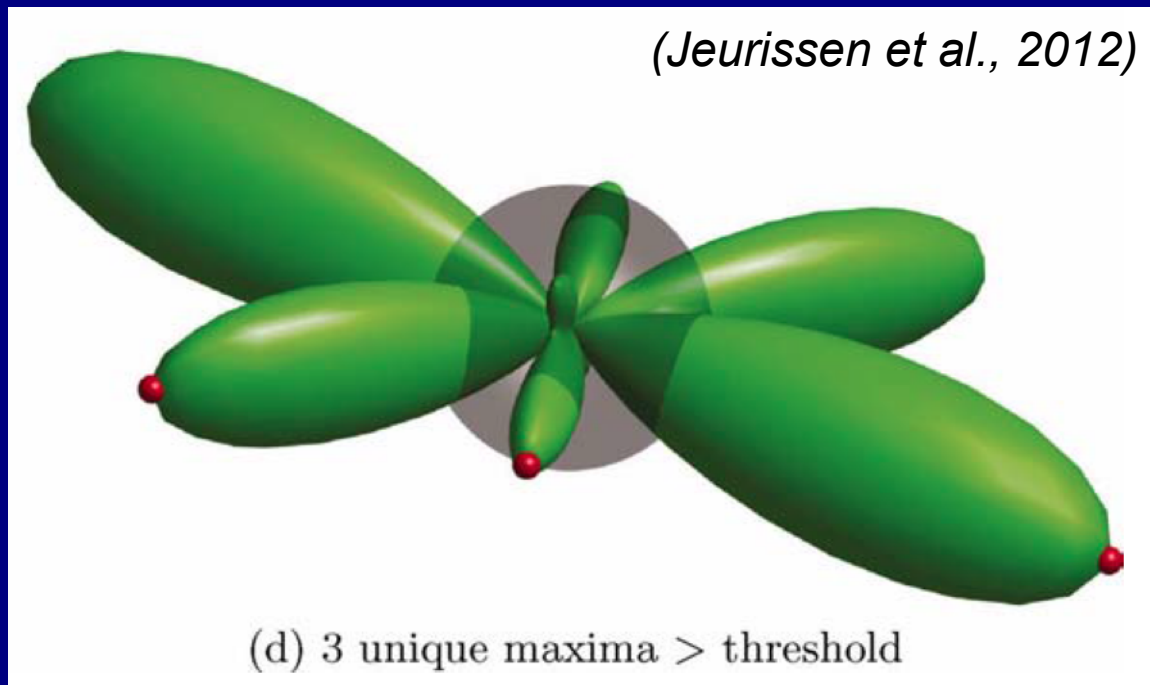
HARDI

- + High Angular Resolution Diffusion Imaging:
 - DSI, ODF, Qball, FOD...
 - model multiple fiber bundle directions per voxel
 - generally need more scan time and acquisitions and computational power, much higher b-values
 - still can't resolve intravoxel tract behavior (which of multiple paths?)
 - higher DW \rightarrow lower signal, so susceptible to noise



HARDI

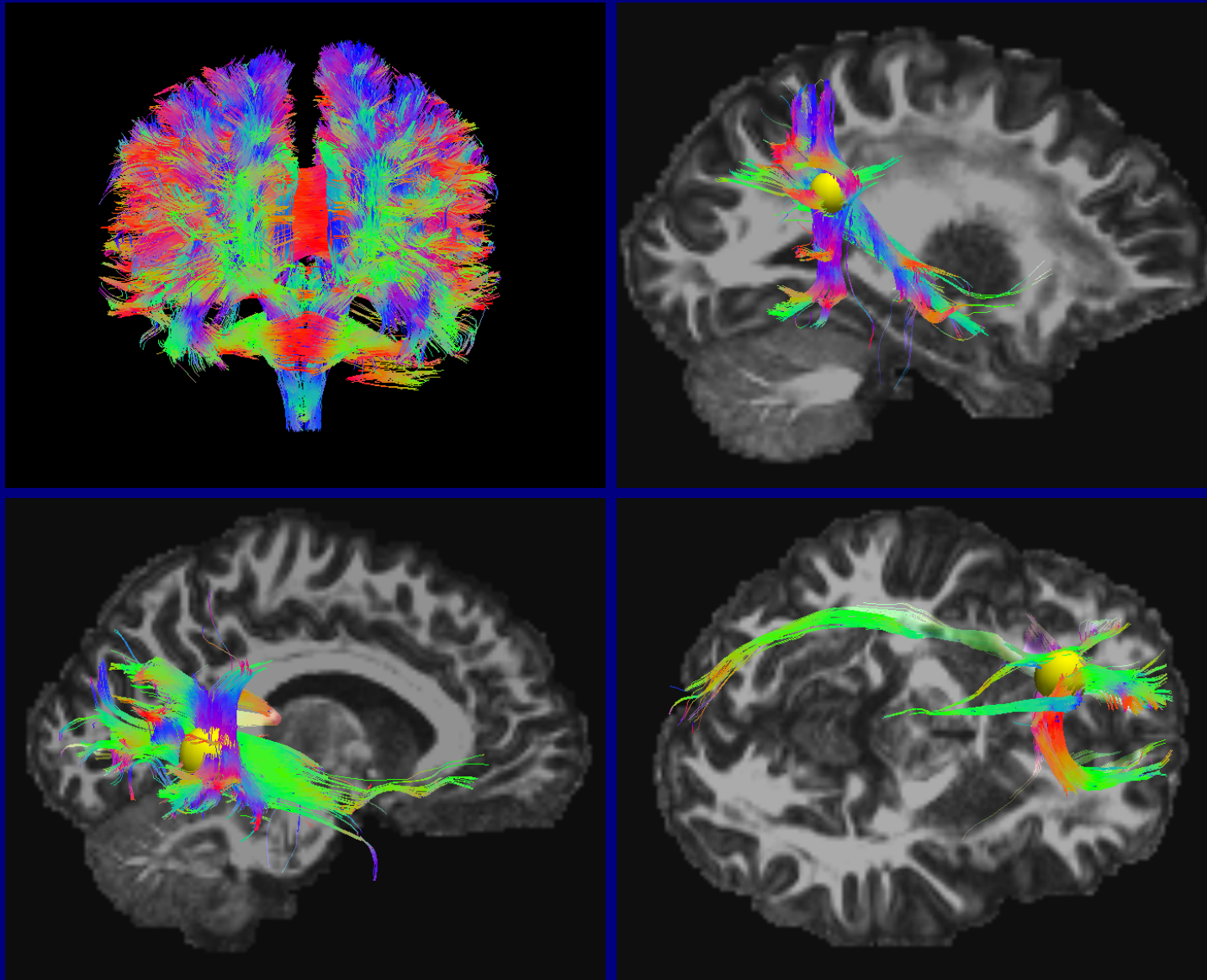
- + High Angular Resolution Diffusion Imaging:
 - DSI, ODF, Qball, FOD...
 - model multiple fiber bundle directions per voxel
 - generally need more scan time and acquisitions and computational power, much higher b-values
 - still can't resolve intravoxel tract behavior (which of multiple paths?)
 - higher DW \rightarrow lower signal, so susceptible to noise



FATCAT can now track through HARDI data
 \rightarrow HARDI reconstruction done outside AFNI (e.g., DSI-Studio, Diffusion Toolkit, FSL), and outputs tracked in FATCAT.

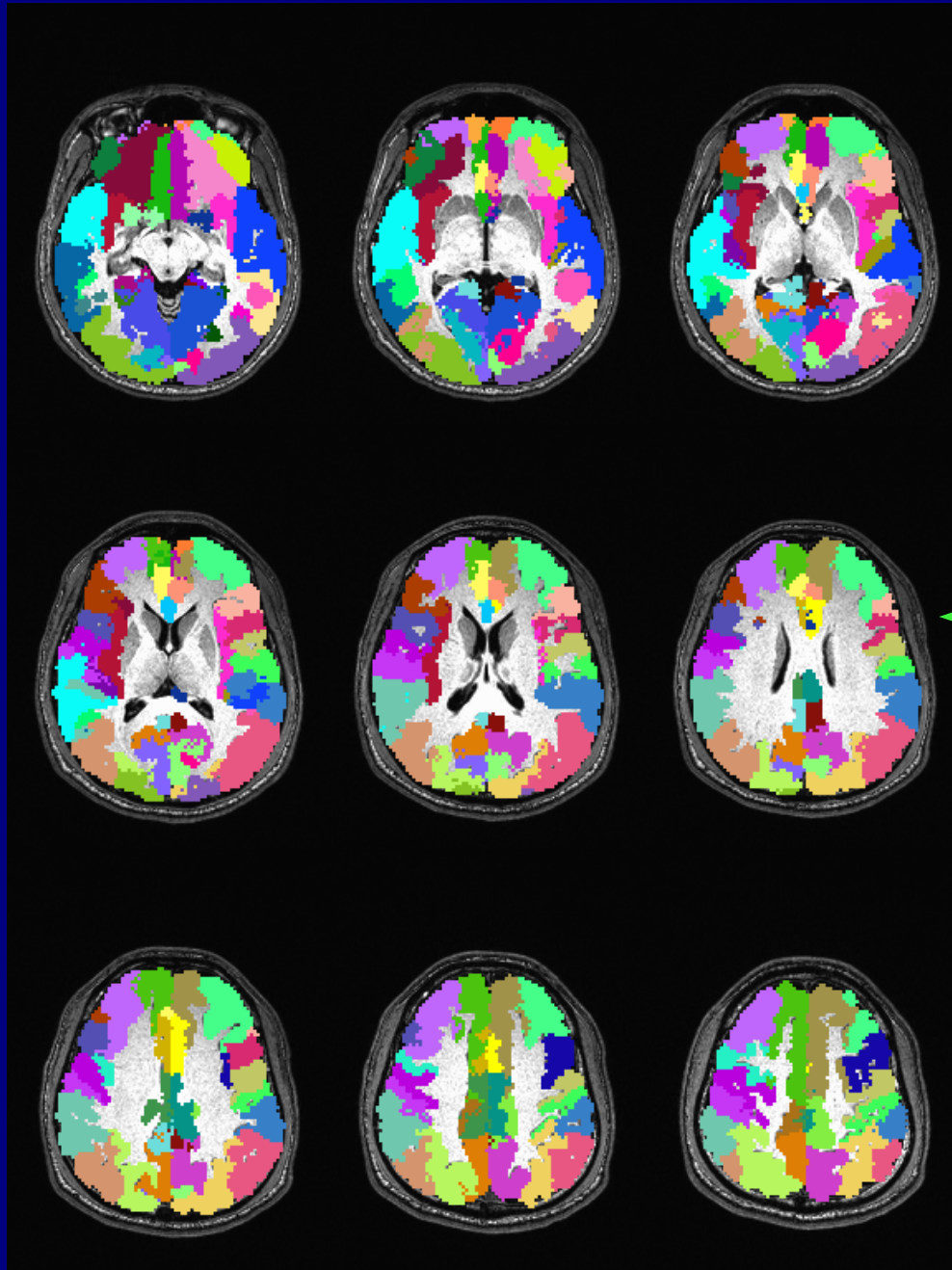
Example: 3dTrackID on HARDI data

*Ex: Human Connectome Project subject, 288 grads,
HARDI reconstructed with GQI in DSI-Studio.*



Example:
'Connectome'-type tracking

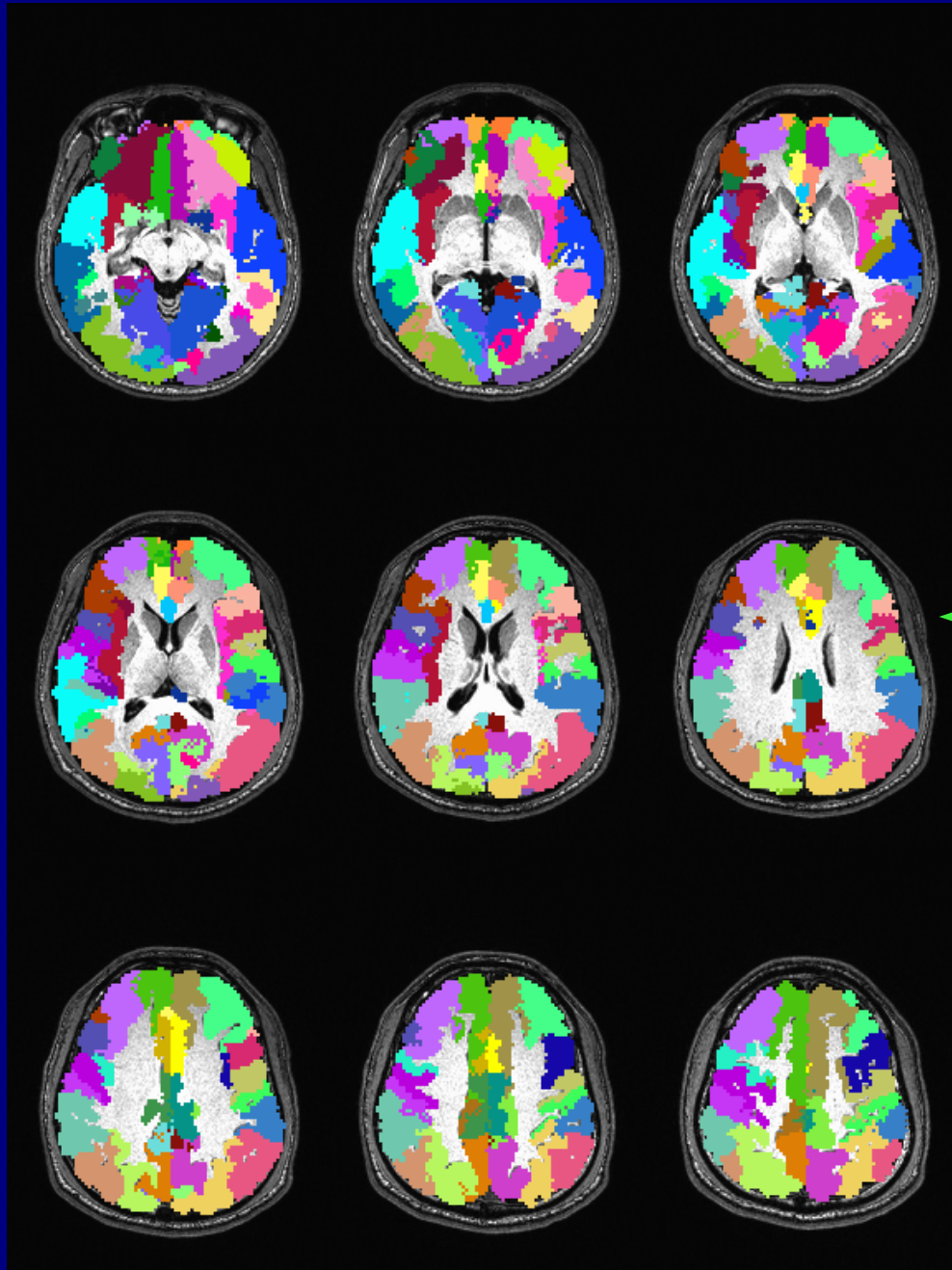
“Connectome”: parcellation of GM



Example (script available in FATCAT_DEMO):

- + Freesurfer parcellation into >112 ROIs.
- + Selected 80 cortical GM ROIs.
- + Used 3dROIMaker to inflate
← by 1 voxel, up to $FA > 0.2$.
(+ *NEW: keep labeltable labels and use them in output.*)
- + '3dTrackID -mode DET' among the regions

“Connectome”: parcellation of GM



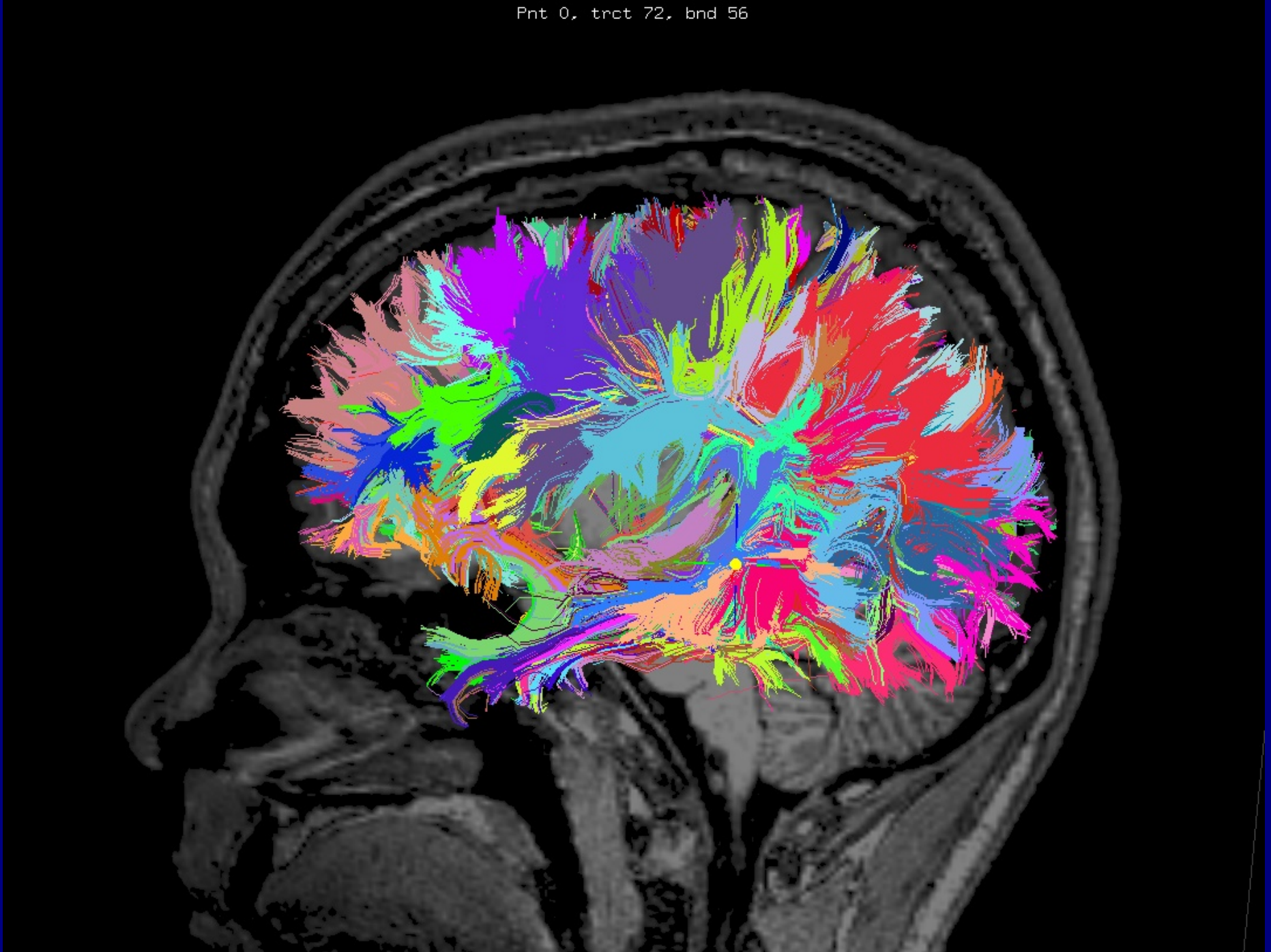
Example (script available in
FATCAT_DEMO):

- + Freesurfer parcellation into >112 ROIs.
 - + Selected 80 cortical GM ROIs.
 - + Used 3dROIMaker to inflate
- ← by 1 voxel, up to $FA > 0.2$.
(+ *NEW: keep labeltable labels and use them in output.*)
- + '3dTrackID -mode DET' among the regions

and a few seconds later... →

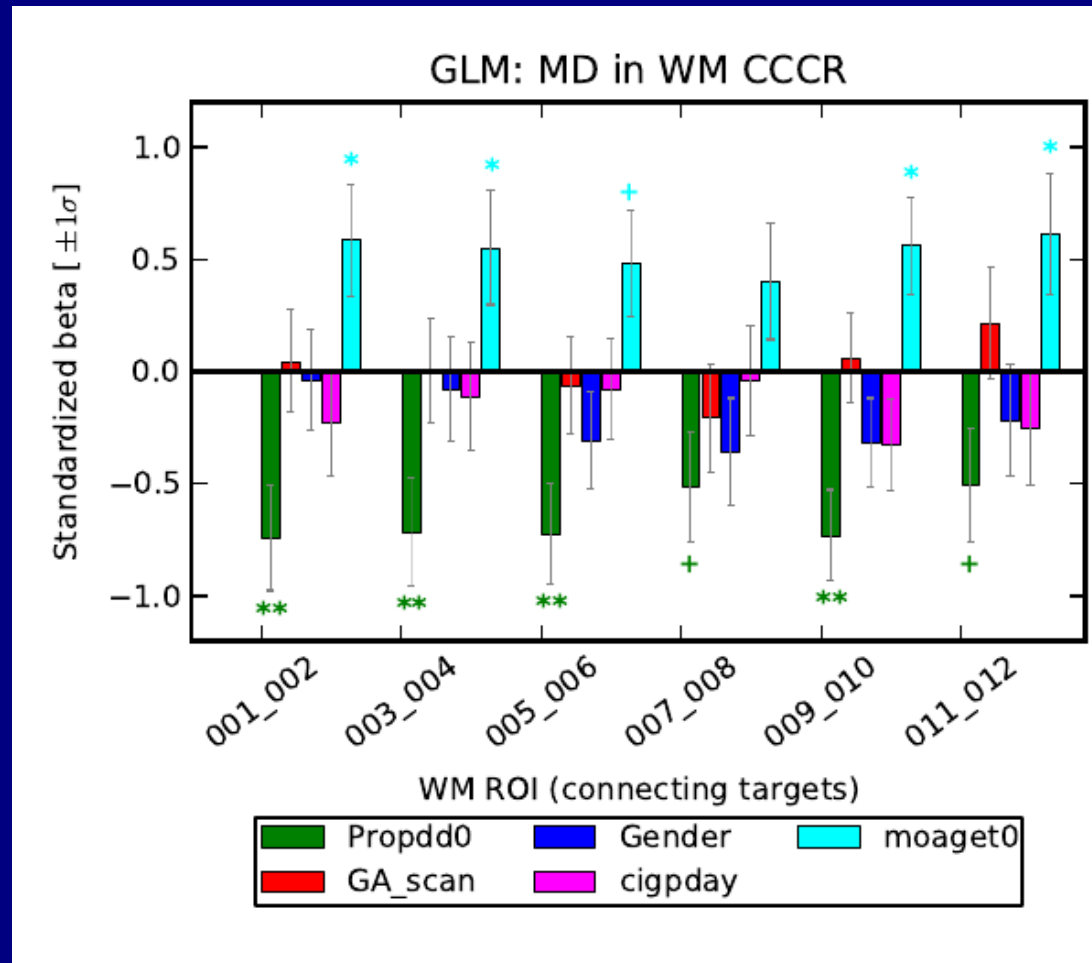
“Connectome”: tracking

Pnt 0, tract 72, bnd 56



A brief example for statistical analysis

- + Combining tractography, quantitative DTI and subject measures with GLM to find structure-alcohol consumption relation:

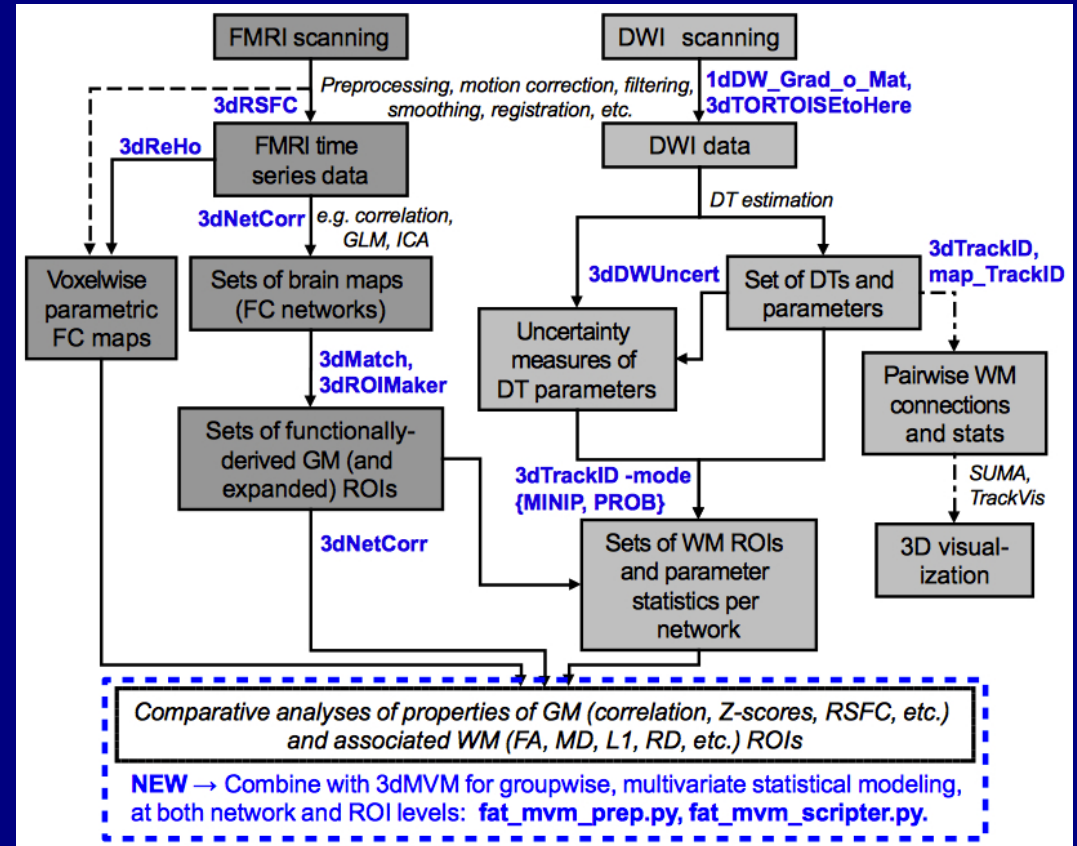


Significant (* $p < 0.05$; ** $p < 0.01$) explanation of DTI measures MD in specific WM regions of CC by alcohol measure (Propdd0) in GLMs which controlled for several other factors.

In Summary

We have discussed capabilities and benefits of:

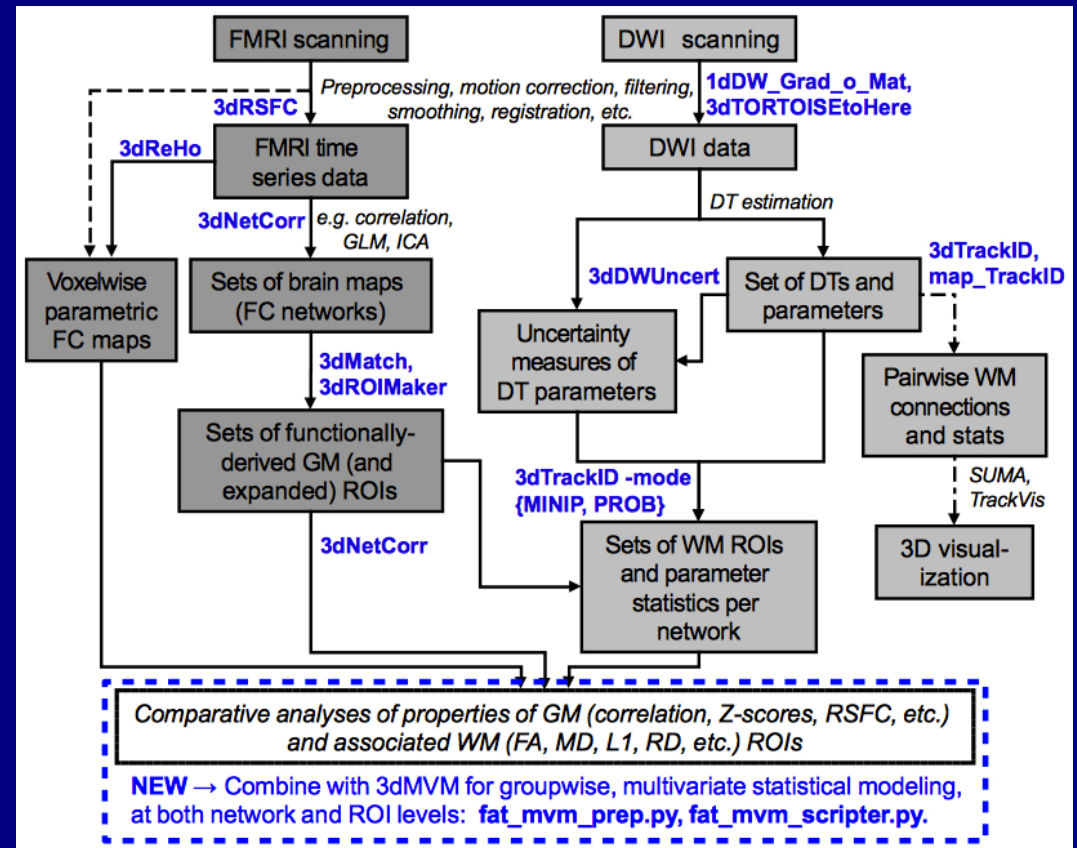
Combining multimodal data: FC+SC+...



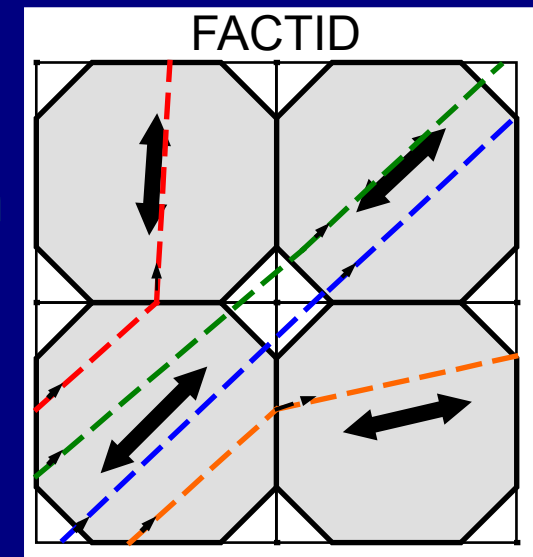
In Summary

We have discussed capabilities and benefits of:

Combining multimodal data: FC+SC+...



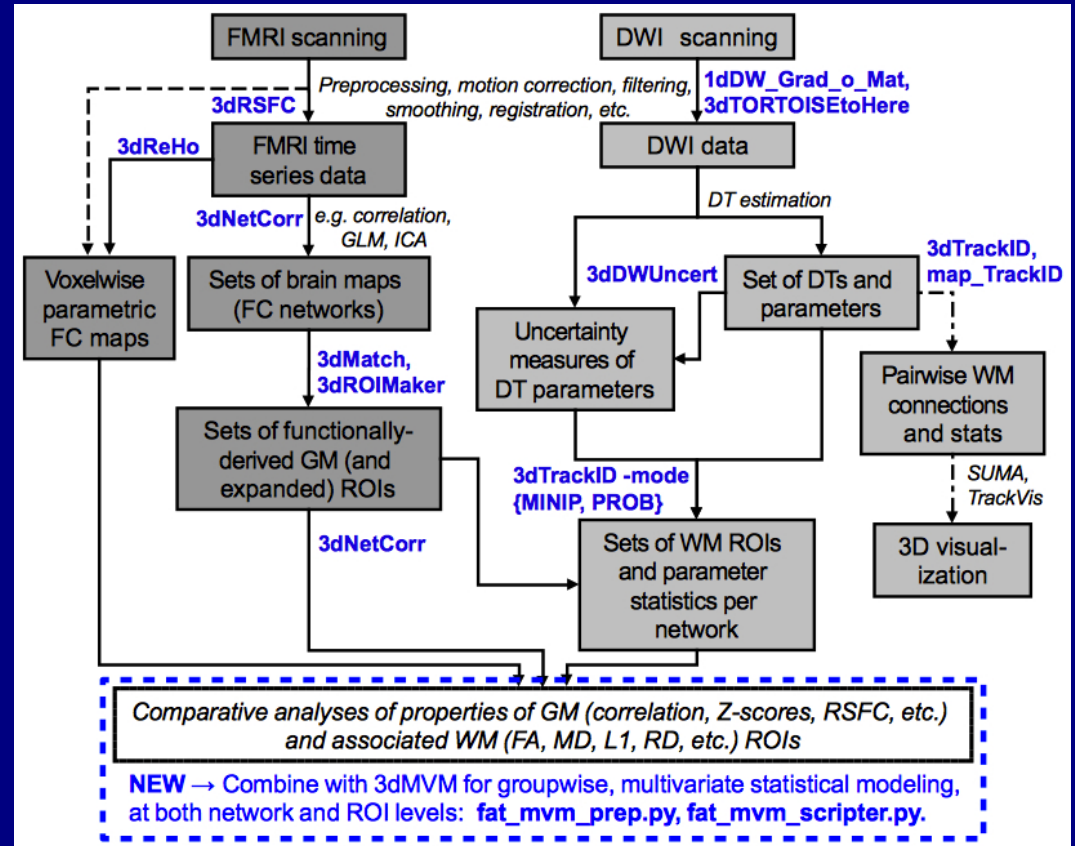
Using an efficient algorithm, reduced bias of propagation



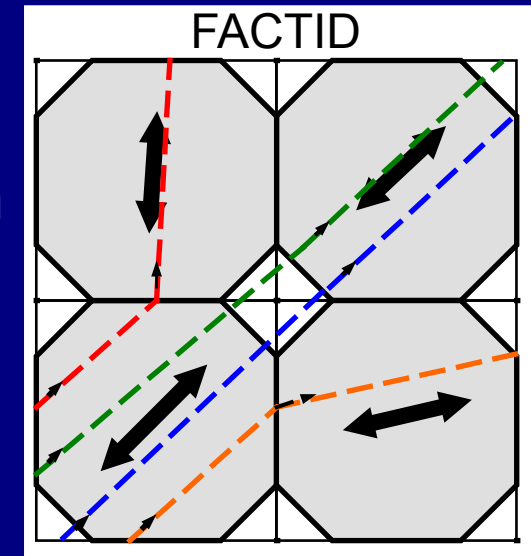
In Summary

We have discussed capabilities and benefits of:

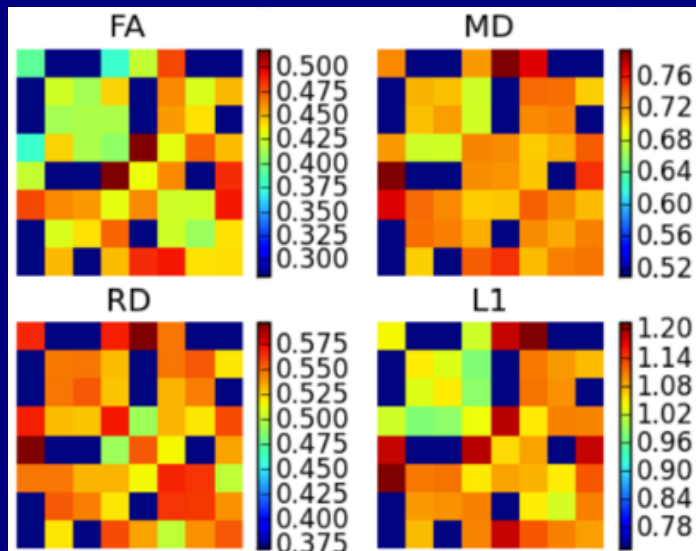
Combining multimodal data: FC+SC+...



Using an efficient algorithm, reduced bias of propagation



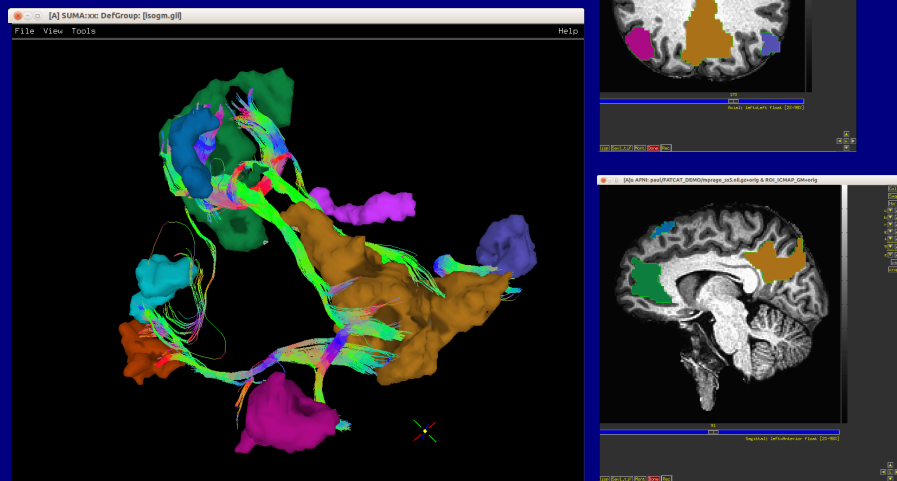
Tracking to define and quantify WM ROIs (with uncertainty/probabilistic)



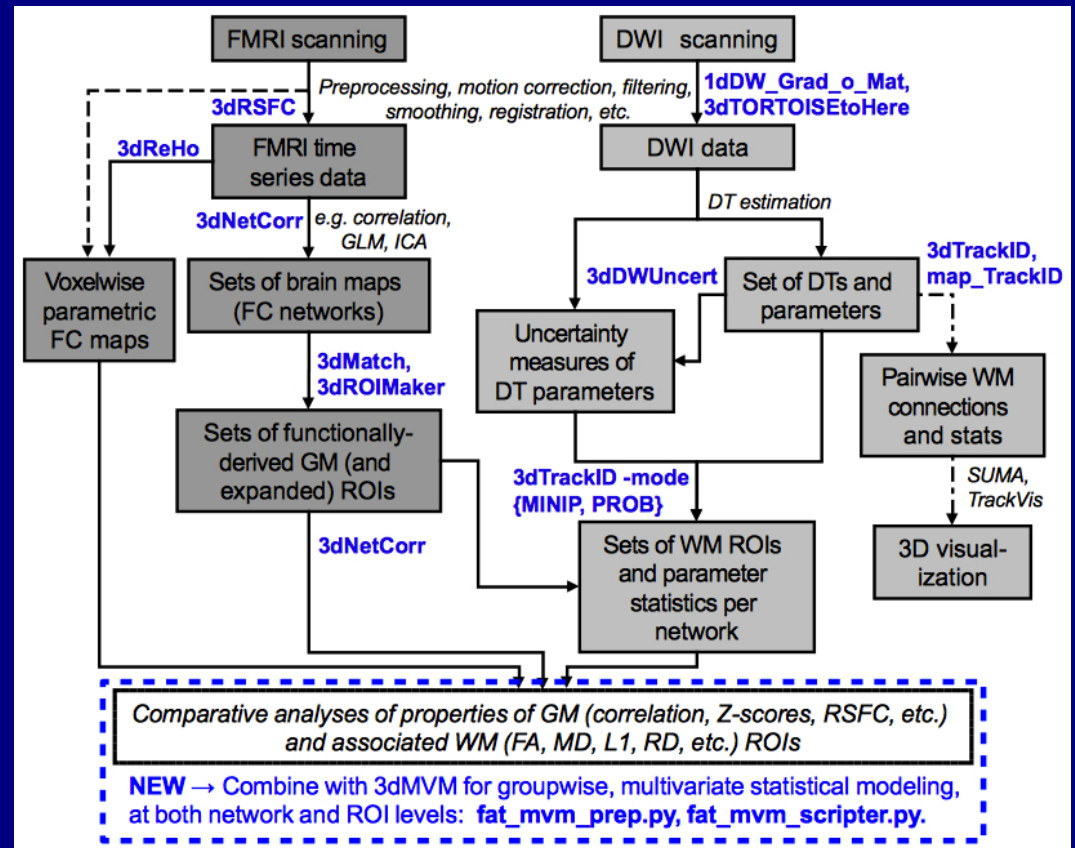
In Summary

We have discussed capabilities and benefits of:

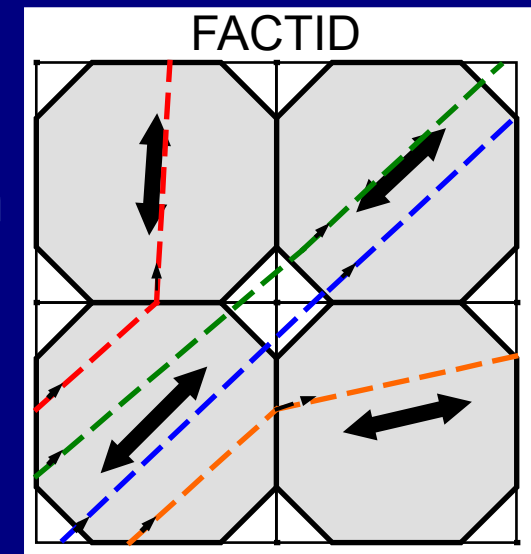
Integrating AFNI-SUMA visualization



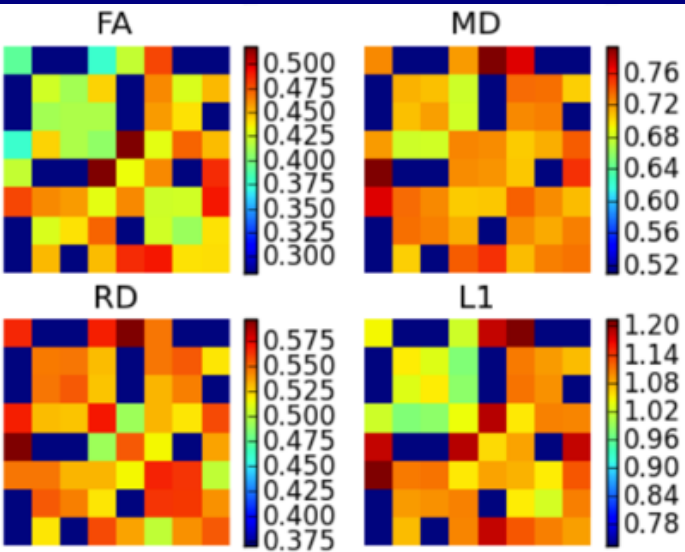
Combining multimodal data: FC+SC+...



Using an efficient algorithm, reduced bias of propagation



Tracking to define and quantify WM ROIs (with uncertainty/probabilistic)



Thanks

And thanks to collaborators:

UMDNJ/NJIT:

Bharat Biswal
Suril Gohel
Xin Di

NIMH/NIH:

Ziad Saad
Rick Reynolds
Gang Chen
Bob Cox

Emory:

Helen Mayberg
Justin Rajendra
Ki Sueng Choi

UCT:

Ernesta M. Meintjes
Alkathafi Alhamud
Chris Molteno
Fleur Warton
Mwape Mofya

CTLFASD Study:

Sandra W. Jacobson (Wayne St.)
Joseph L. Jacobson (Wayne St.)
Andre van der Kouwe (Harvard/MGH)
Pia Wintermark (Montreal Children's)

AIMS:

Johan de Villiers