Introduction to AFNI-FATCAT

Tractography for data exploration and complementing functional connectivity

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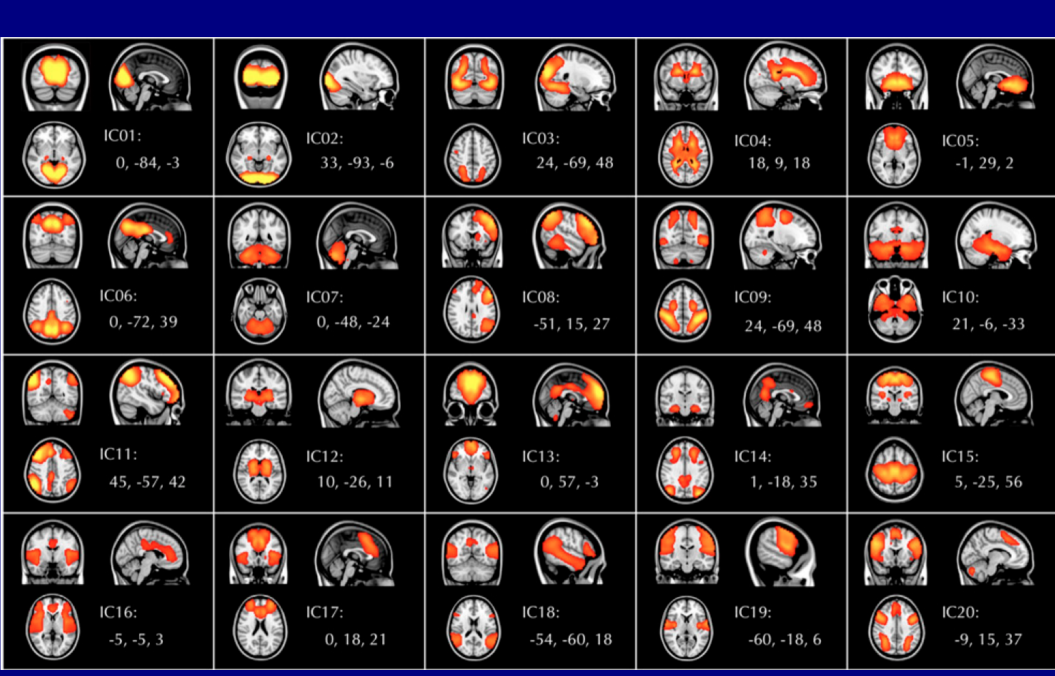




Outline

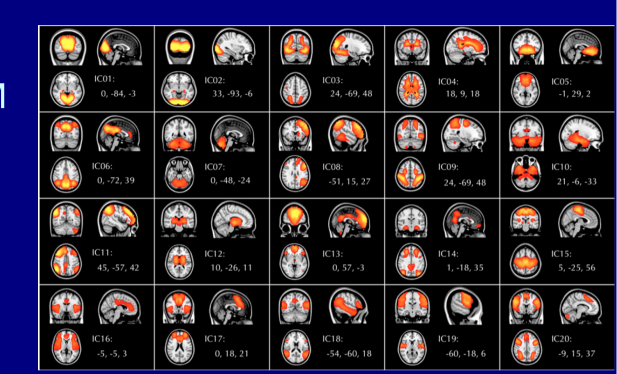
- + Why Function+Structure
- + DWI and DTI (very brief, following morning session)
 - Diffusion imaging basics and parameters
- + Using tractography to estimate WM connections
 - Making targets from functional data
 - Deterministic, probabilistic (or both?)
 - using WM region properties for quantitative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

FMRI: GM Networks



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.

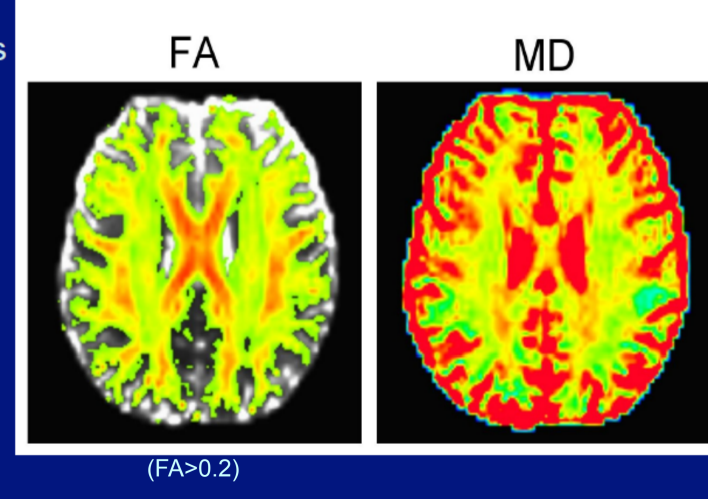


Structural (WM)

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.

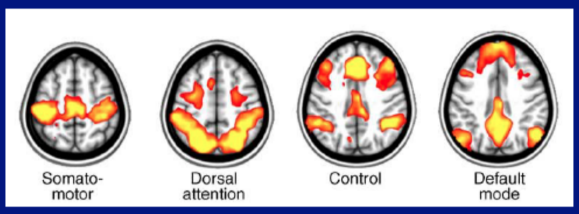
Can investigate (and quantify?) network relations with: tractography



Structure + Function

Simple example:

GM ROIs network:

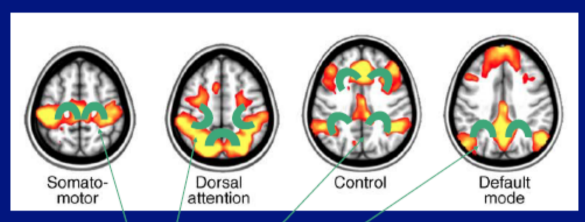


Raichle (2010, TiCS)

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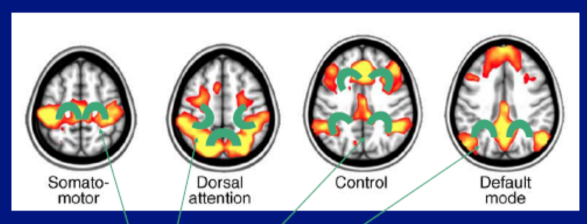
Raichle (2010, TiCS)

Associated WM ROIs

Structure + Function

Simple example:

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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

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 - 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...

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

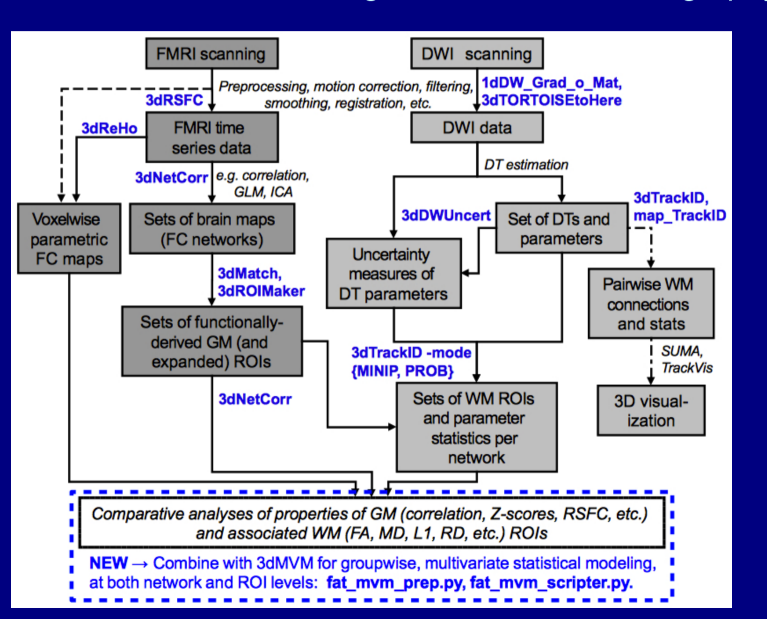




*picture from google search, not from/of either author

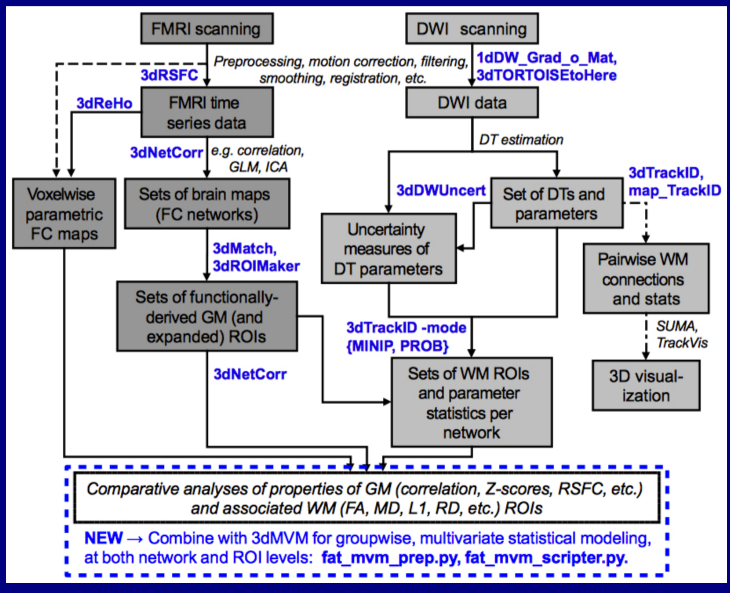
Functional and structural processing

Schematic for combining FMRI and DTI-tractography via FATCAT:



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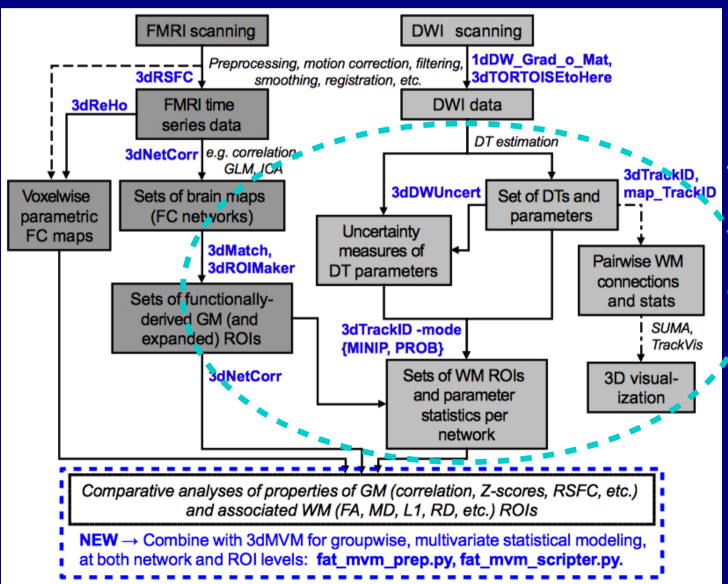


FATCAT goals:

- + 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

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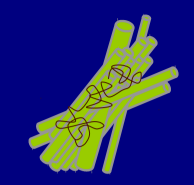
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Main focus today on DTItractography, including making ROIs from FMRI

(In brief)

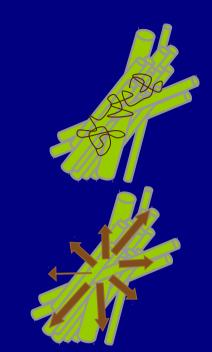
1) Random motion of molecules affected by local structures



(In brief)

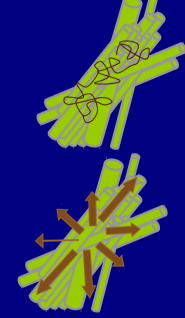
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2) Statistical motion measured using diffusion weighted MRI

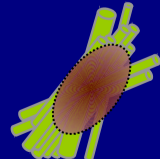


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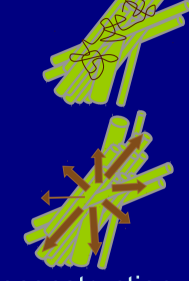


- 3) Bulk features of local structure approximated with various reconstruction models, mainly grouped by number of major structure directions/voxel:
 - + one direction:DTI (Diffusion Tensor Imaging)



(In brief)

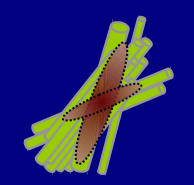
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- 3) Bulk features of local structure approximated with various reconstruction models, mainly grouped by number of major structure directions/voxel:
 - + one direction:DTI (Diffusion Tensor Imaging)
 - + >=1 direction:

 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: **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

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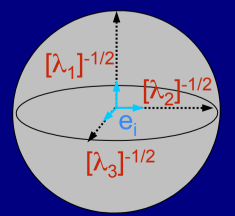
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6 independent values

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

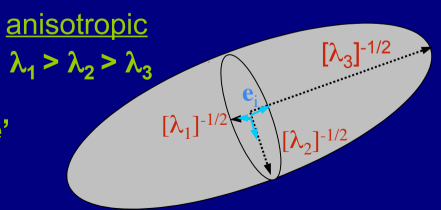
$$C = \mathbf{r}^{\mathsf{T}} \mathbf{D} \mathbf{r} = D_{11} x^2 + D_{22} y^2 + D_{33} z^2 + 2(D_{12} xy + D_{13} xz + D_{23} yz)$$



<u>isotropic</u>

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

`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

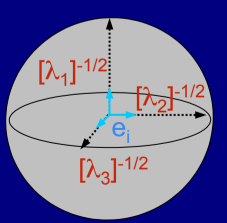
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Minimum number of measures

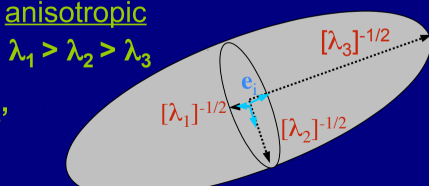
6 independent values

ellipsoid surface



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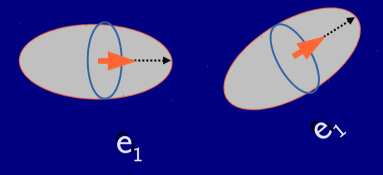
"Big 5" DTI ellipsoid parameters

Main quantities of diffusion (motion) surface

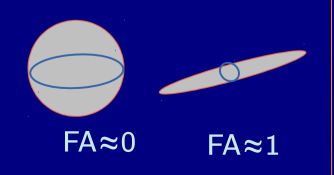
first eigenvalue, L1 = λ_1 , parallel/axial diffusivity, AD



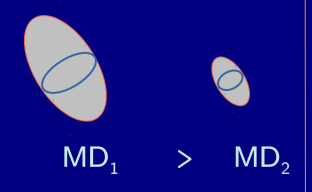
first eigenvector, e₁



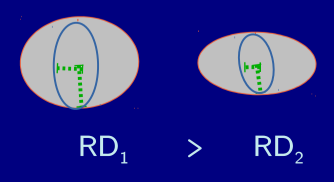
Fractional anisotropy, FA



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

e₁: orientation of major bundles

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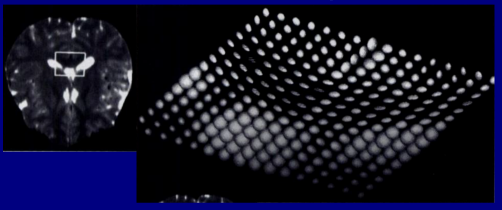
e₁: orientation of major bundles

Cautionary notes:

- + Degeneracies of structural interpretations
- + Changes in myelination may have small effects on FA
- + WM bundle diameter << 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

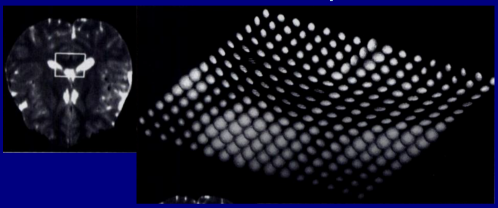
Local DTs → **Extended Tracts**

Field of local diffusion parameters



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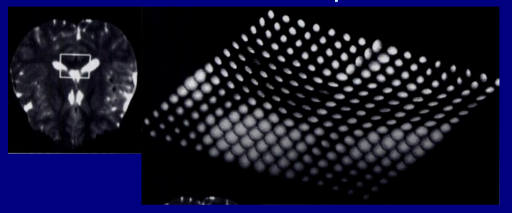


→ individual ellipsoids

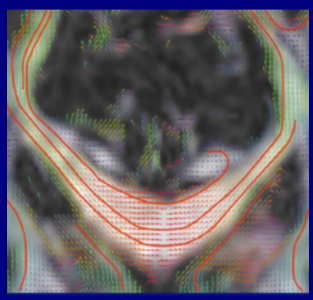


Local DTs → Extended Tracts

Field of local diffusion parameters



Connect to form extended tracts



→ individual ellipsoids

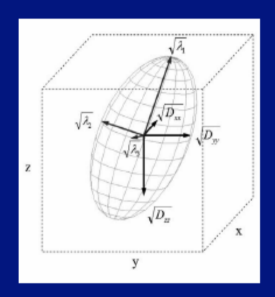


→ linked structures

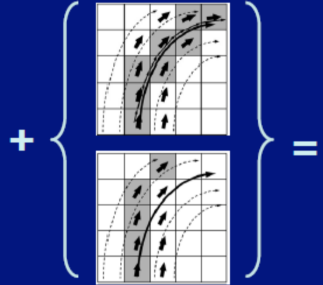


Tractography

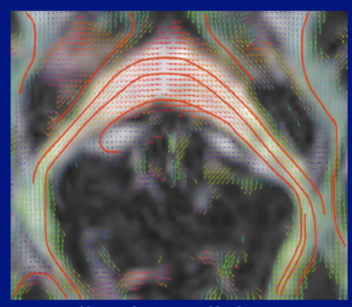
Estimate WM structure (fiber tract locations)



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

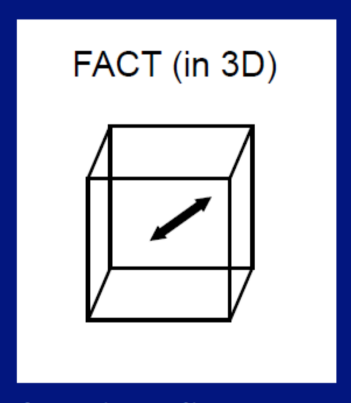
So, first question for using tractography in a study:

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 stays>0.2 and angle between e₁s is <45 deg

FACT (in 2D)



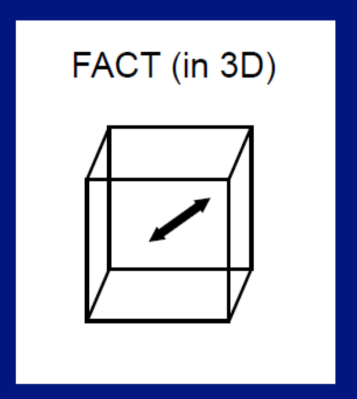
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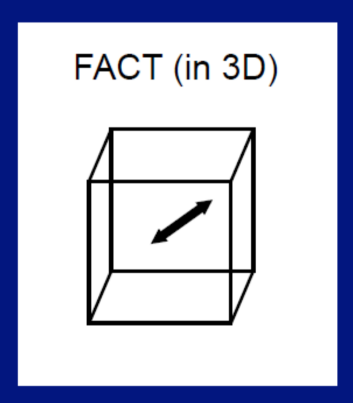


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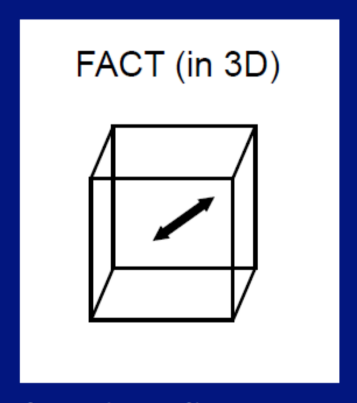
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FACT (in 2D)

Noise-> angular shift

Ex.:



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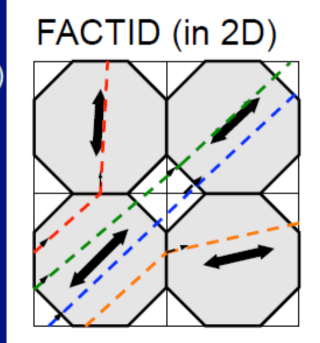
Improving FACT->

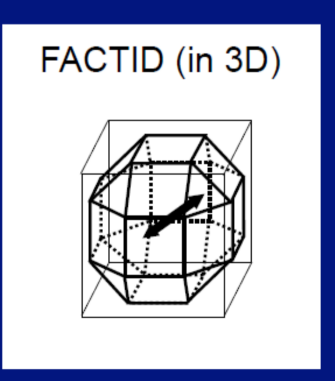
- Start by thinking: what properties a 'good' algorithm should have?
 - Should be independent of coordinate axes (i.e., results invariant to rotation of data set)
 - Should improve with spatial resolution (convergence in resolution)
 e.g., like in calculus, diagonals are better approximated with small grid steps
 - 3) Should improve with SNR (converge in SNR)
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Posit: including diagonal (ID) propagation helps 1 and 4, check about other props.

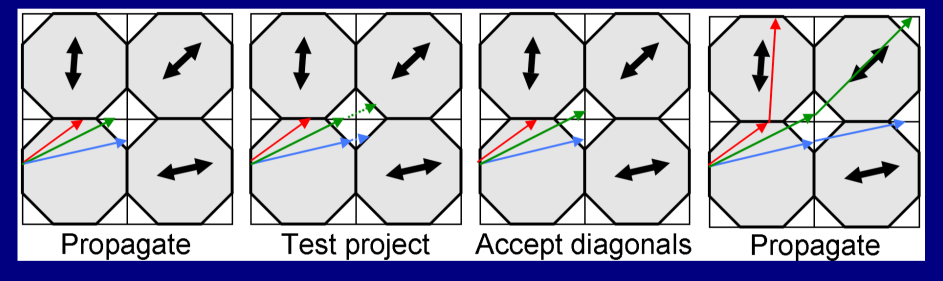




FACTID (FACT Including Diagonals):

+ Utilize simple check for diagonals.

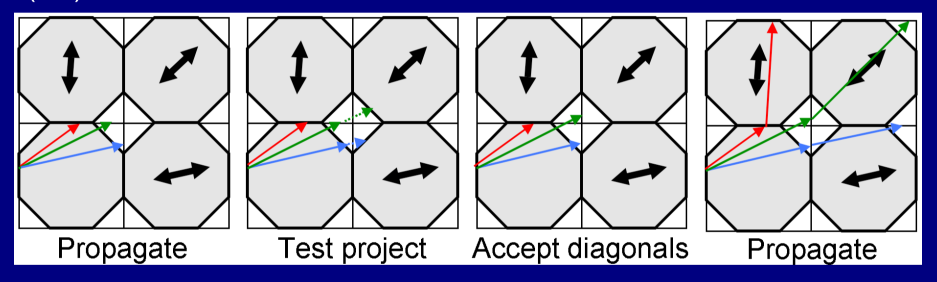
(2D) Schematic:



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(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)

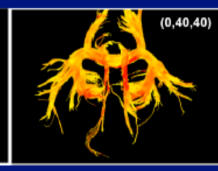
FACTID



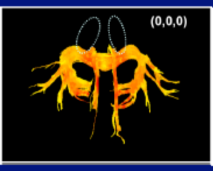






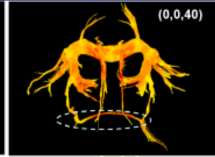


FACT



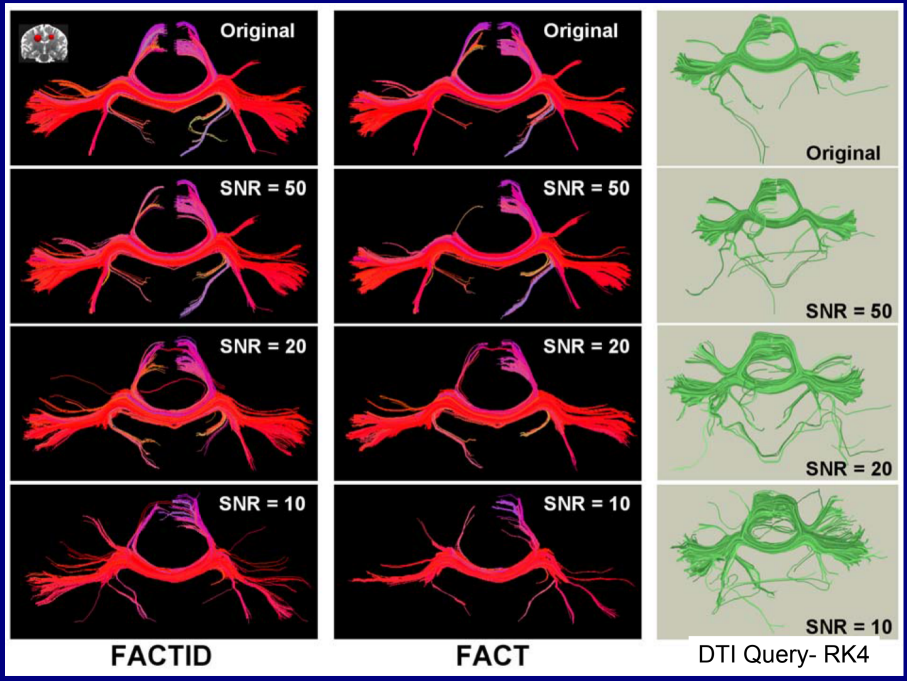








Test 3: Noise sensitivity



Test 5: Phantom Set

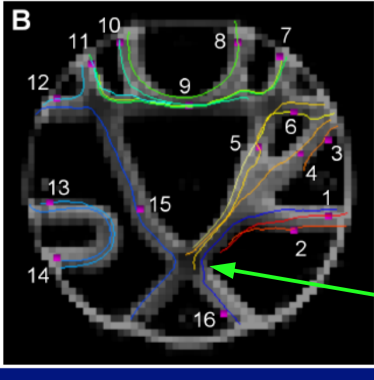
Fillard et al. (2011, NI) test phantom

FACT



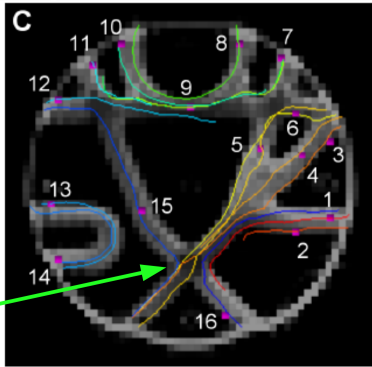
FACTID

"ANSWER"



(Taylor, Cho, Lin & Biswal, 2012)

e.g. compare



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.

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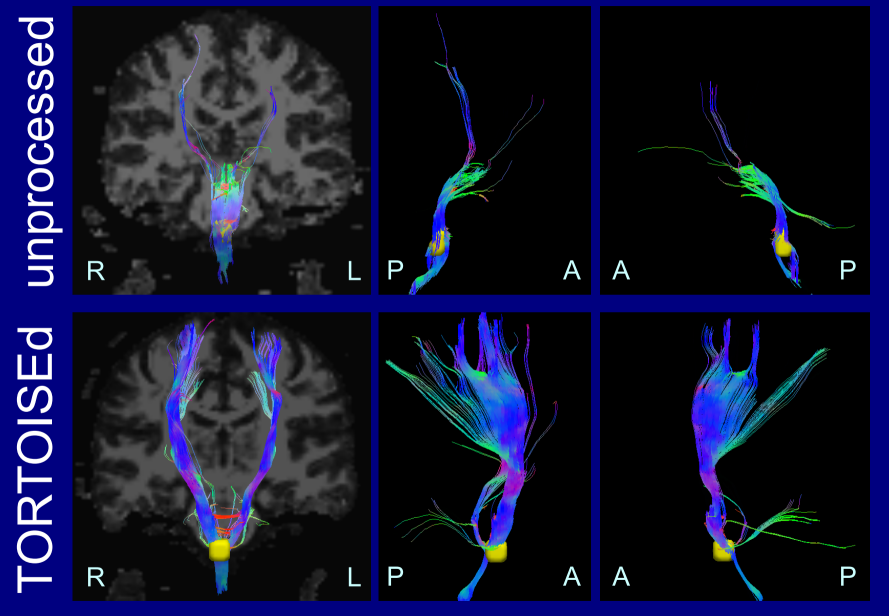
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→ ** In addition to the tracking algorithm, the quality of data acquisition and preparation matter quite a bit (as seen in morning TORTOISE session). **

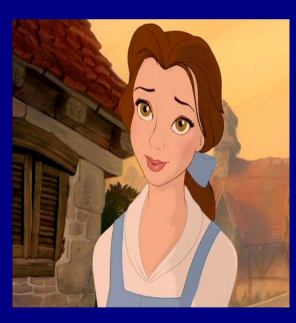
Importance of being processed (in earnest)



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

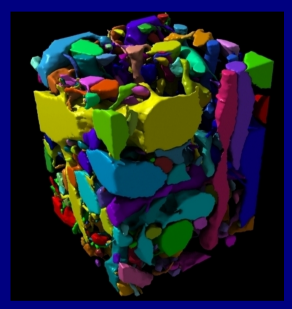


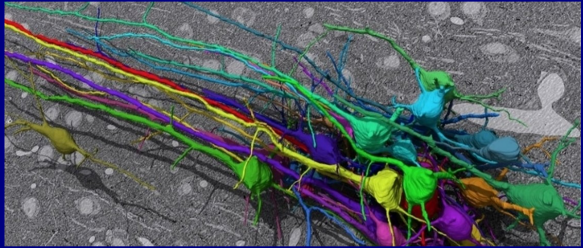


Known Challenges for Tracking

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





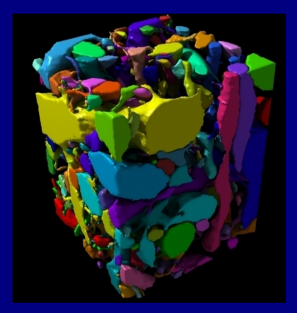


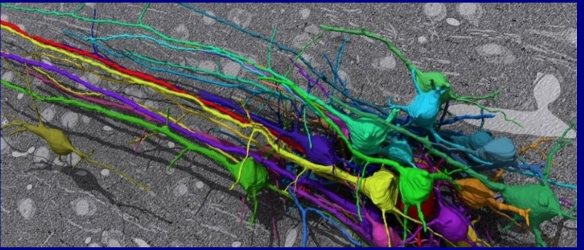
(images of Eyewire data via NPR website)

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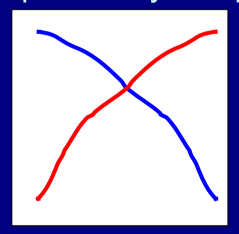


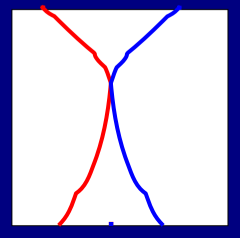




(images of Eyewire data via NPR website)

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





Crossing/kissing fibers can:

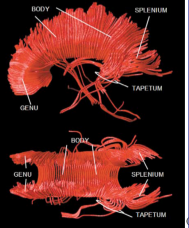
- Lower FA (stop tracking)
- Redirect (or *not*) tracking incorrectly.

Achievements of Tracking

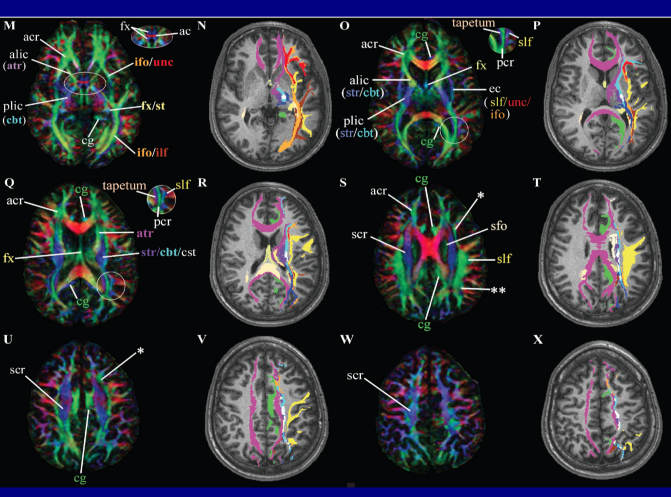


- + Reproduction of many known pathways
- + In vivo vs post-mortem information





(Bammer et al., 2003)



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?

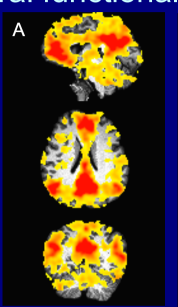
Next question for doing tractography:

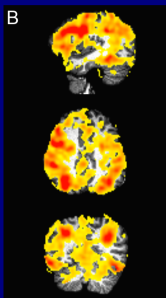
where does one go to get the ROIs to try to connect?

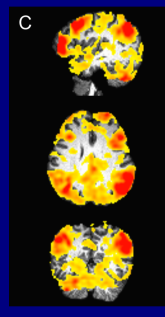
-> could go to atlases and standard maps, or to exploratory spheres dotted around,

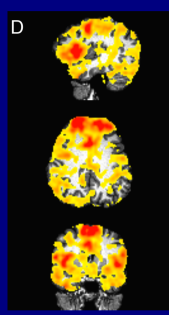
+ For example, one can perform ICA on a resting state study, resulting in several functional networks:

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





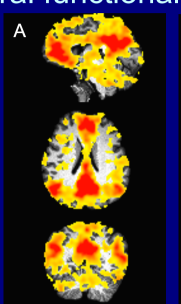


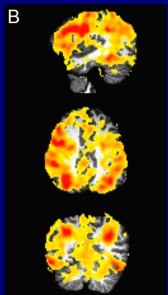


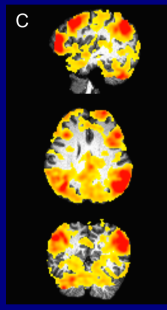
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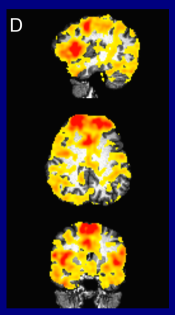
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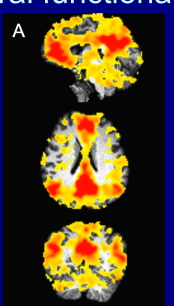


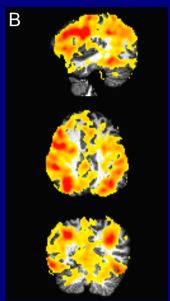


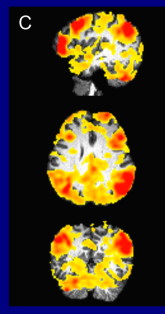
+ want to isolate GM ROIs, and then to expand each to make sure that they are at least touching nearby (associated?) WM voxels to have any hope to connect tracts

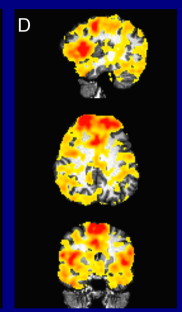
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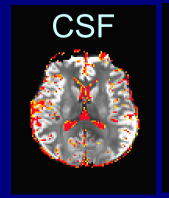


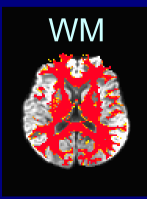






- + 3dROlMaker can parcellate into GM ROIs based on:
 - thresholding voxel values
 - thresholding cluster size
 - subtract away CSF and WM voxels from segmentation maps
 - expand each GM ROI to location of WM (don't want to overexpand unphysically)



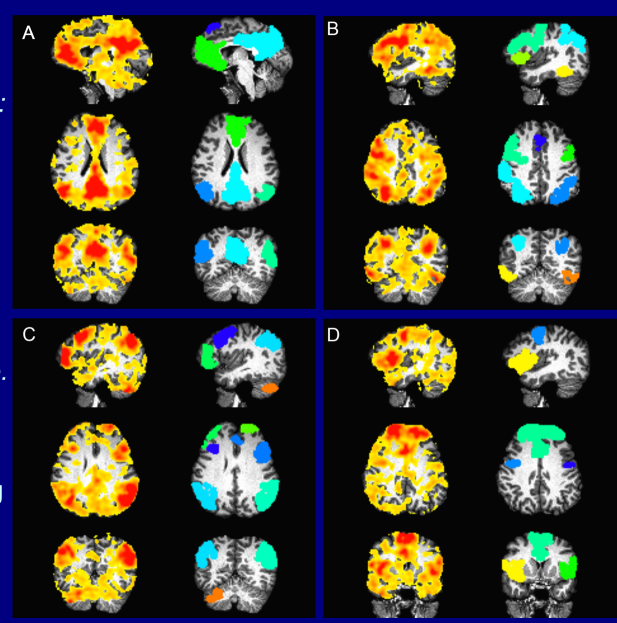


Example case for ICA networks:

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.



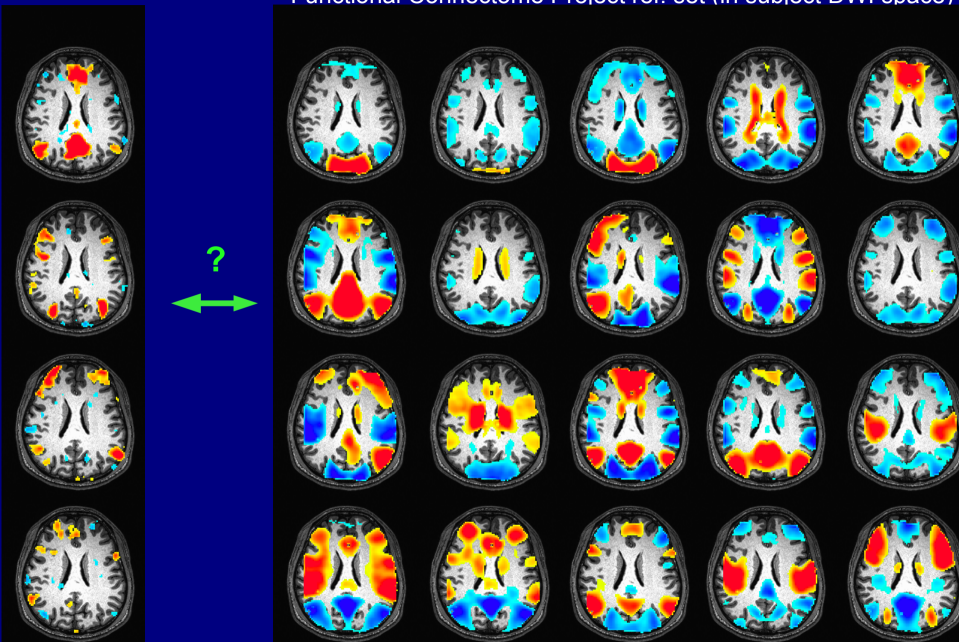
Sidenote:

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

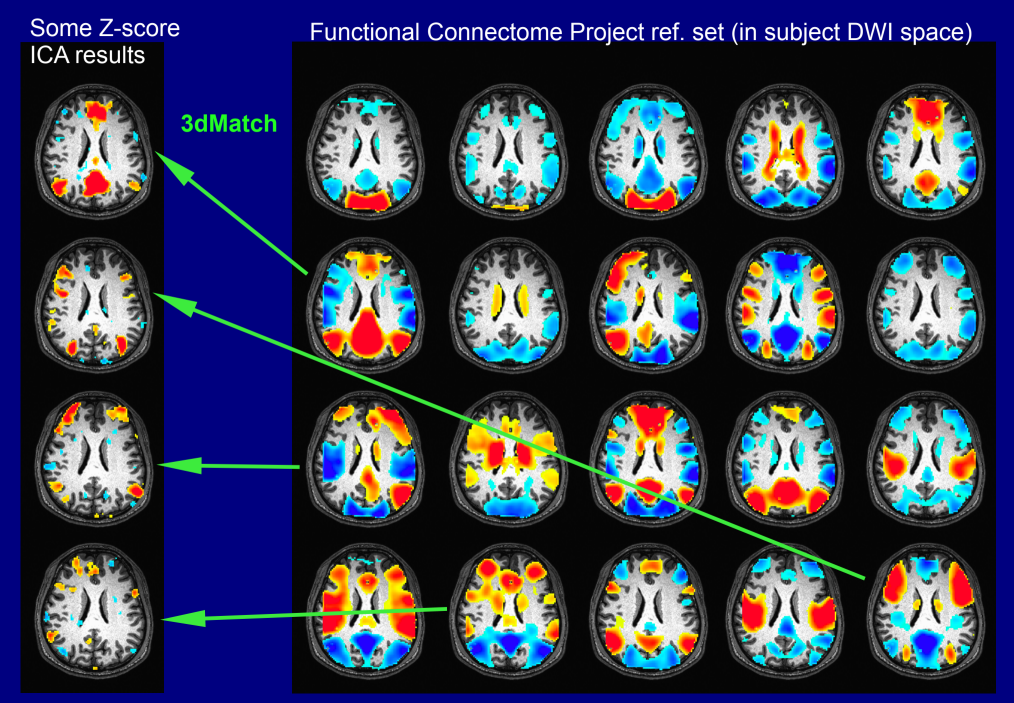
Matching Network maps

Some Z-score

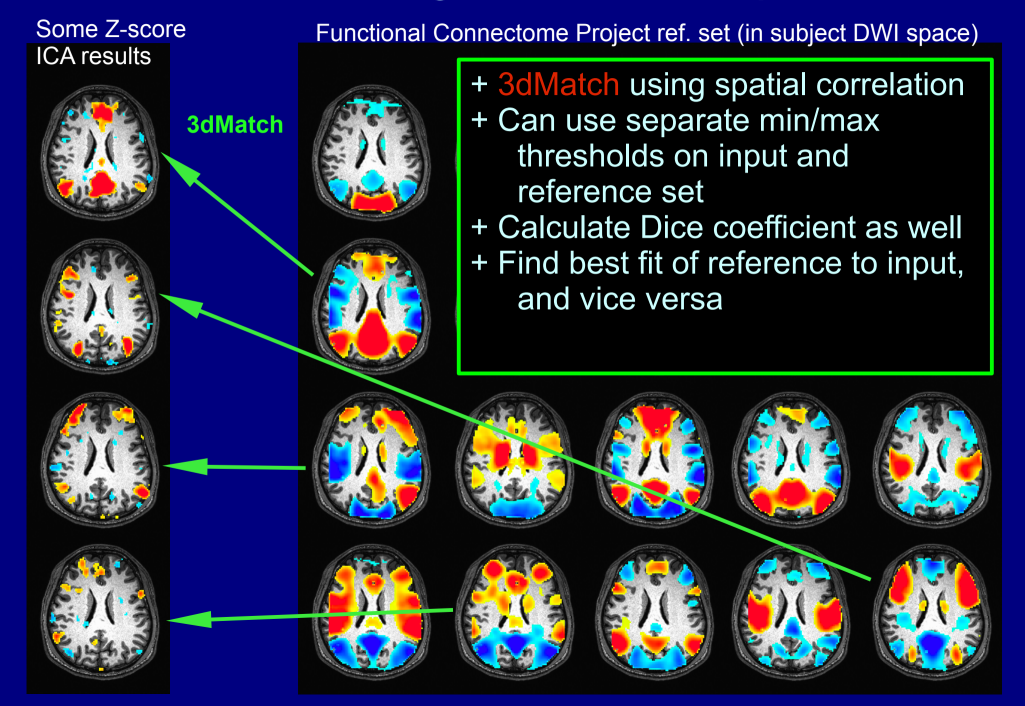
Functional Connectome Project ref. set (in subject DWI space)



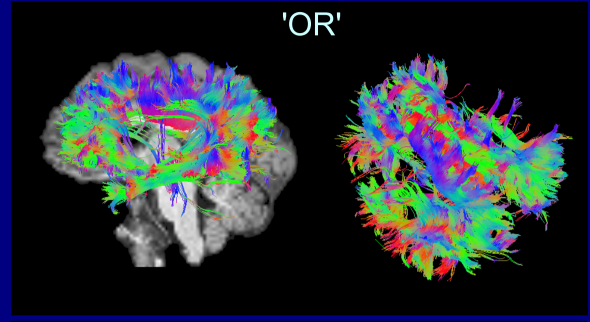
Matching Network maps



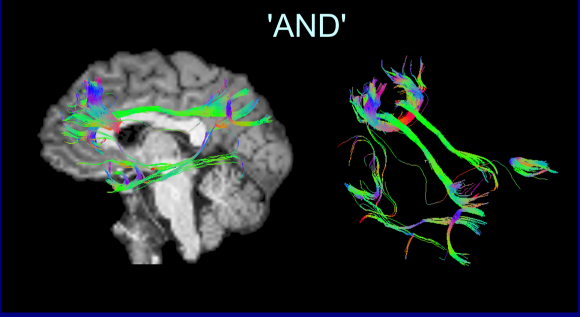
Matching Network maps



- + 3dTrackID -mode DET -logic { OR | AND }
- + uses FACTID
- + good for exploratory analysis and visualization of results

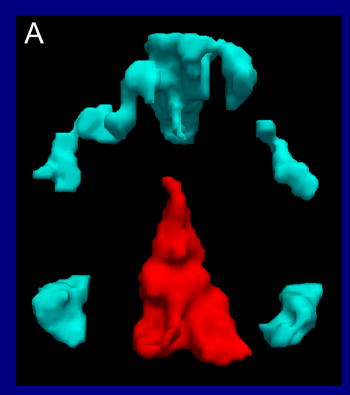


ex.: DMN network tractography results using ROIs from 3dROIMaker (FA>0.2; max angle 60deg; 8 seeds/voxel)



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

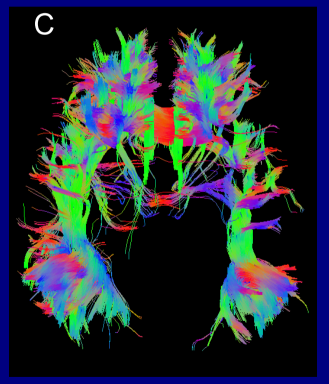
Control track propagation with `anti-mask' regions, simply defined by voxels =-1:



ROIs: blue>0, red<0



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

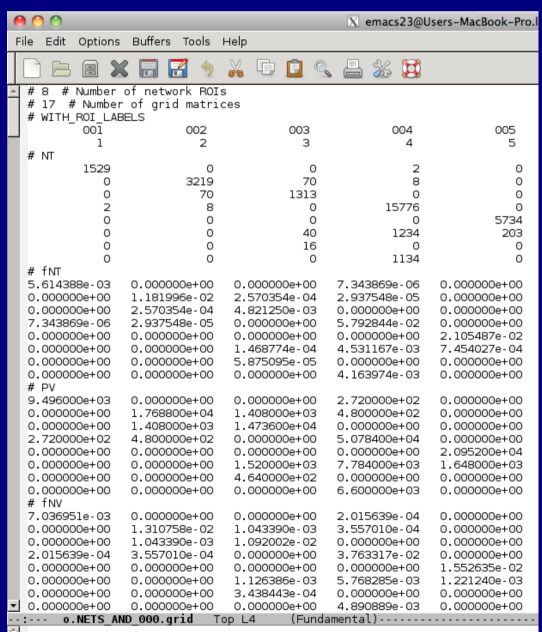


results when: blue>0, red<0 (using anti-masks)

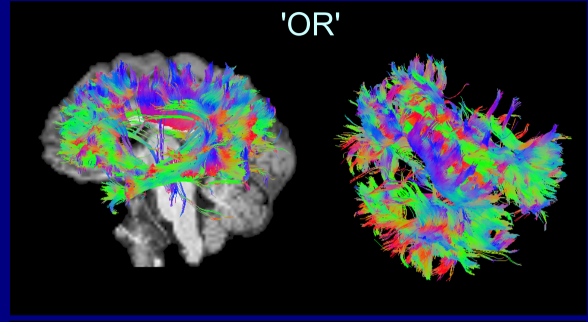
- + 3dTrackID -mode DET -logic { OR | AND }
- + Automatic quantification per network in produced PREFIX.grid files.

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.



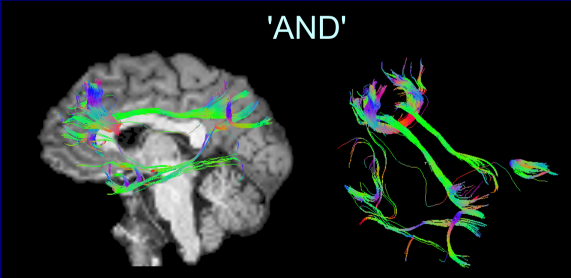
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ex.: DMN network tractography results using ROIs from 3dROIMaker

(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:

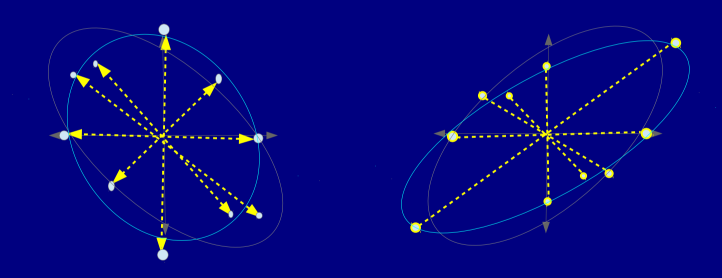
How do we estimate tensor parameter noise/uncertainty?

Noise in DW signals

MRI signals have additive noise

$$S_{i} = S_{0} e^{-b g_{i}^{\mathsf{T}} \mathsf{D} g_{i}} + \varepsilon,$$

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

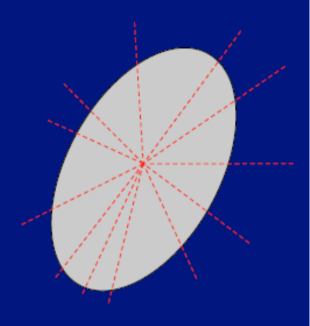
EPI distortions, subject motion, et al. also warp ellipsoids.

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

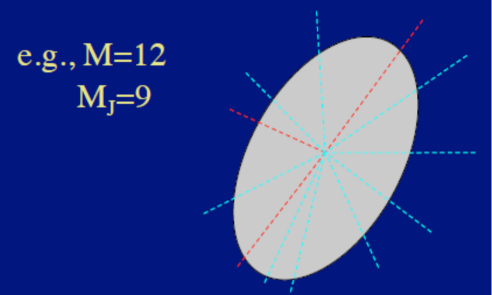
Jackknifing

• Basically, take M acquisitions



Jackknifing

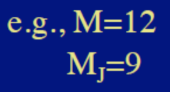
- Basically, take M acquisitions
- Randomly select M_J < M to use to calculate quantity of interest
 - standard nonlinear fits

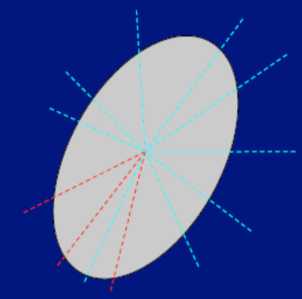


$$[D_{11} \ D_{22} \ D_{33} \ D_{12} \ D_{13} \ D_{23}] = \dots$$

Jackknifing

- Basically, take M acquisitions
- Randomly select M_J < M to use to calculate quantity of interest
 - standard nonlinear fits
- Repeatedly subsample large number (~10³-10⁴ times)



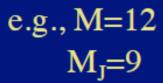


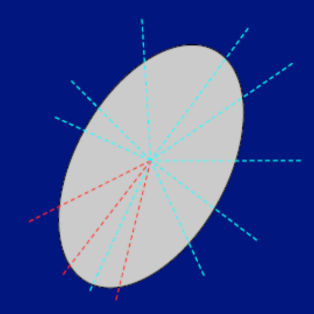
$$\begin{aligned} [D_{11} & D_{22} & D_{33} & D_{12} & D_{13} & D_{23}] = \dots \\ [D_{11} & D_{22} & D_{33} & D_{12} & D_{13} & D_{23}] = \dots \\ [D_{11} & D_{22} & D_{33} & D_{12} & D_{13} & D_{23}] = \dots \end{aligned}$$

• • • •

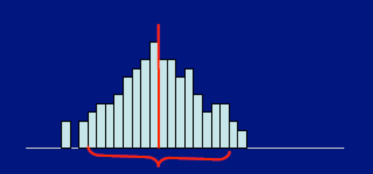
Jackknifing

- Basically, take M acquisitions
- Randomly select M_J < M to use to calculate quantity of interest
 - standard nonlinear fits
- Repeatedly subsample large number (~10³-10⁴ times)
- Analyze distribution of values for estimator (mean) and confidence interval
 - sort/%iles
 - (not so efficient)
 - if Gaussian, e.g. μ±2σ
 - simple



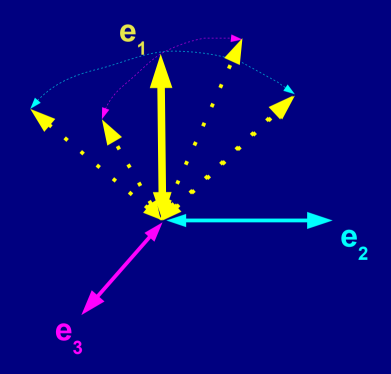


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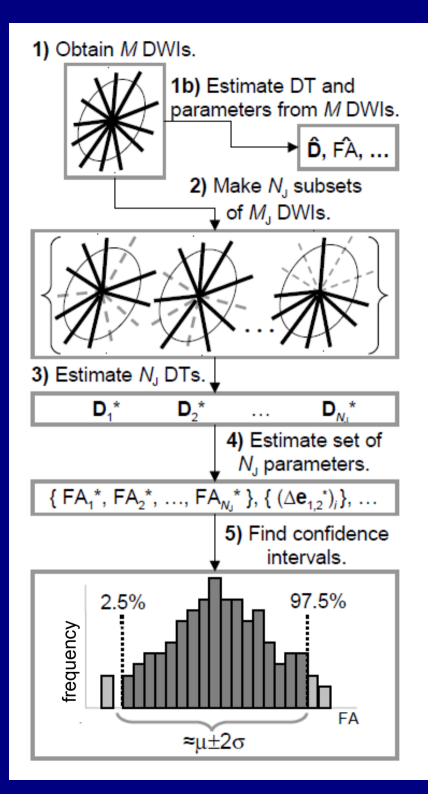


Uncertainty estimation

+ 3dDWUncert estimates bias and σ of first eigenvector e₁ (main direction of diffusion), based on how much it could tip toward either e₂ or e₃:

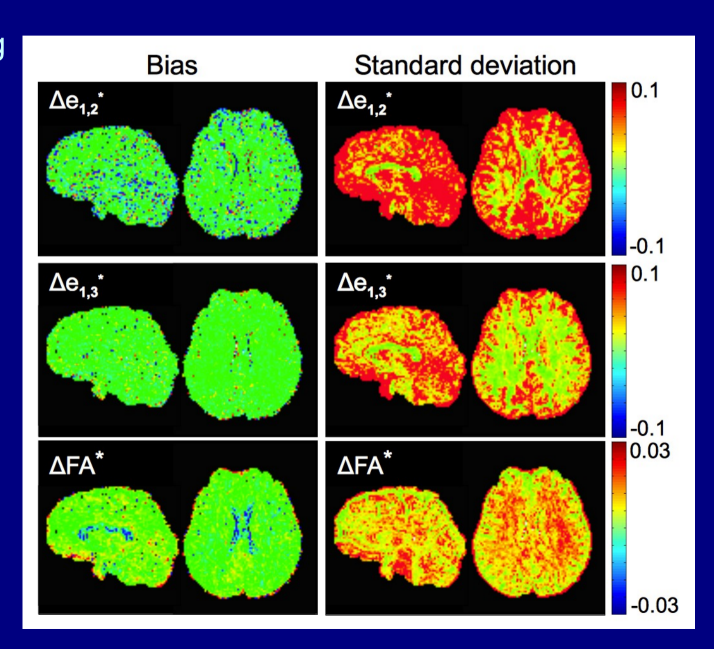


.... and the bias and σ of FA



Uncertainty example

- + Can see difference in e1 uncertainty along e2 and e3
- + Tissue-dependent differences in FA uncertainty



Next question for doing tractography:

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

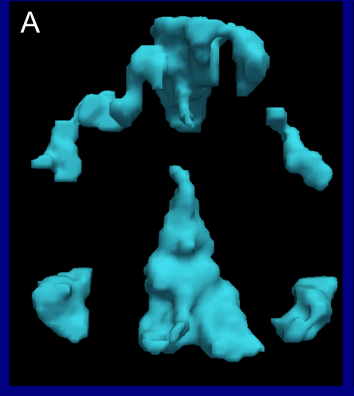
- We know that estimates of DTI ellipsoids are not exactly representing tracts/bundles
 - Size scale differences between voxel/tracts, multiple tracts, complex structure, signal noise, eddy currents, nonlinear fits, etc.
- How to include errors/uncertainty in interpretation and usage?

- 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)

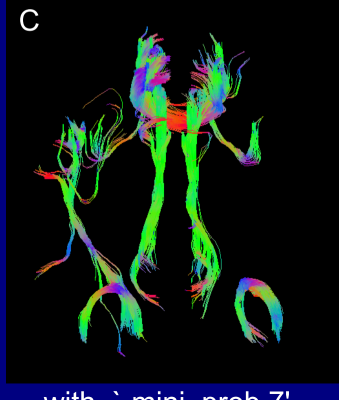
Mini-Probabilistic Tracking

- + 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.









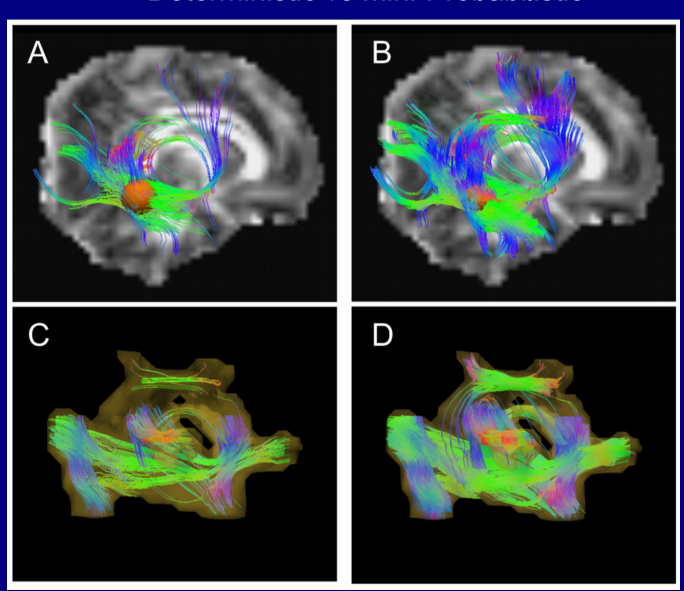
with `-mini_prob 7'

Mini-Probabilistic Tracking

Deterministic vs mini-Probabilistic

Through single ROI

AND logic through network, cf with full-prob results



(Taylor et al., 2014)

 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

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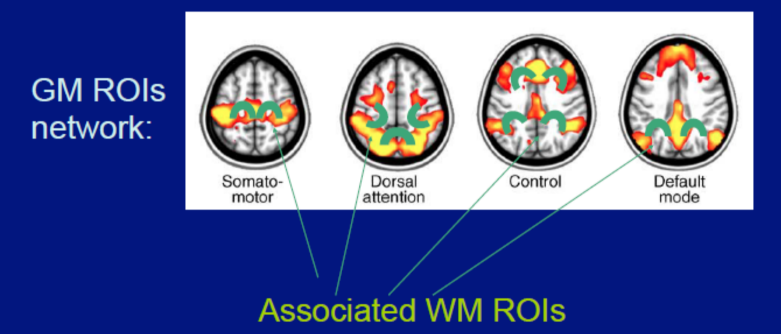
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- While this quantity is somehow relevant in representing what relative 'connectivity' which can be estimated, exact interpretation as 'probability of connectivity' is tricky
 - -> for example, how literally can one equate a numerically-constructed 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?

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- 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 ROIvoxels.
 - Not probability of connectivity of A and X, but more likelihood of a voxel being part of associated WM

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

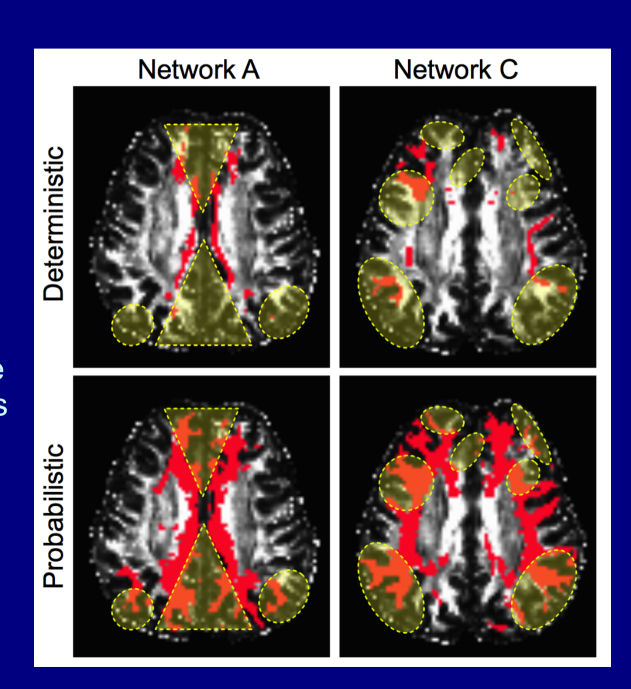


 Threshold Ψ per voxel after probabilistic tracking, use to define WM ROI between GM ROIs

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

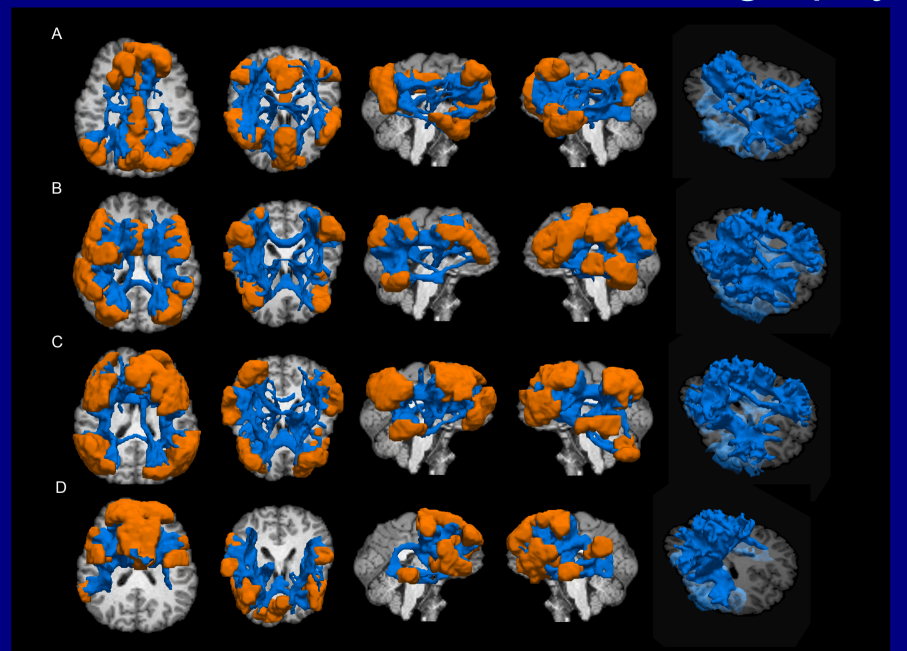


- + with networks of ROIs from 3dROIMaker and uncertainty from 3dDWUncert (as well as tensor estimates from, e.g., 3dDWItoDT), can finally do probabilistic tractography
- + 3dTrackID -mode PROB
 - does lots of Monte Carlo simulations: wholebrain tractography -> perturb FA & e1 based on uncertainty ->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

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 - can trim saved tracts to only keep voxels between 2 ROIs (i.e., no overrunners in the 'connection' ROIs)

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 - 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

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 - 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)



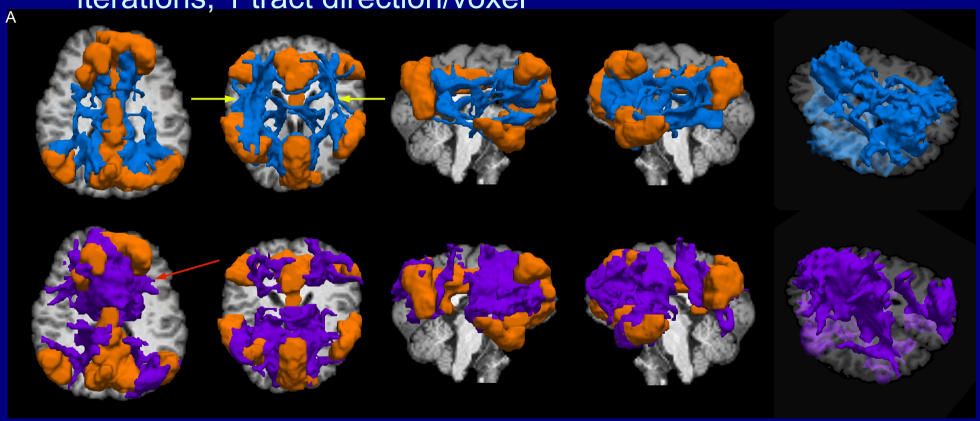
(orange is ROI; blue is set of WM regions with tracts connecting)

+ 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

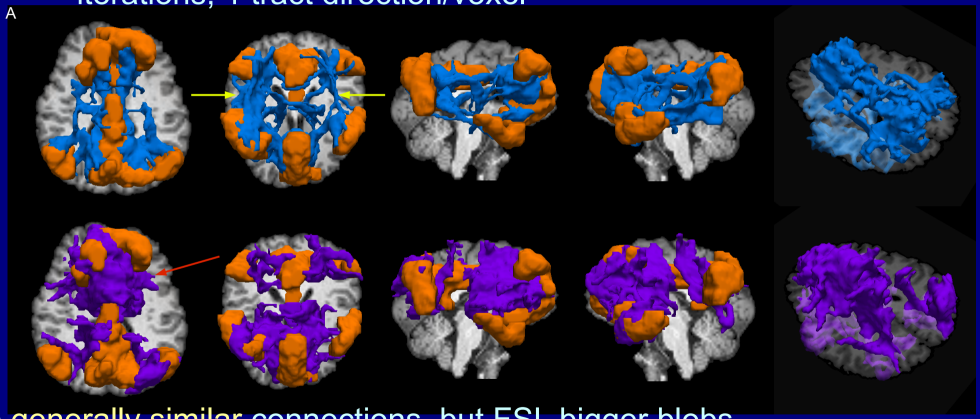


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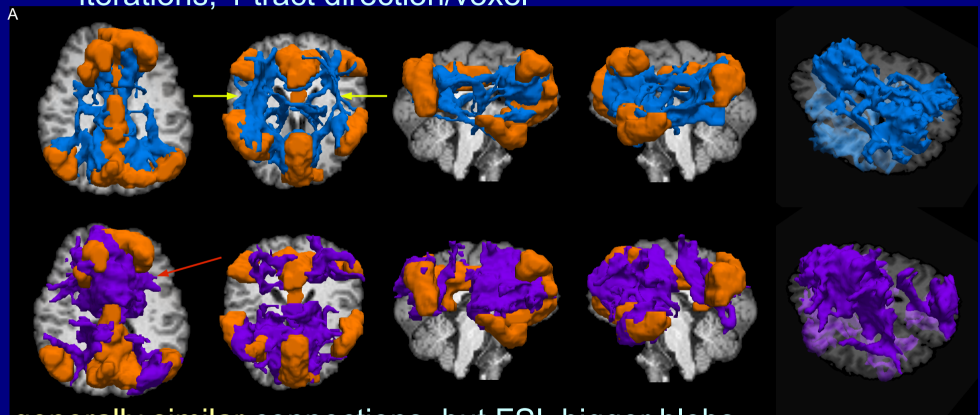
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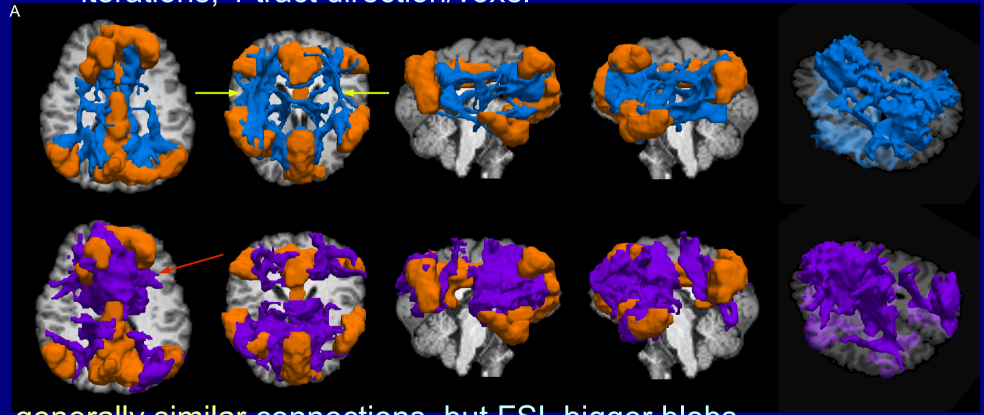
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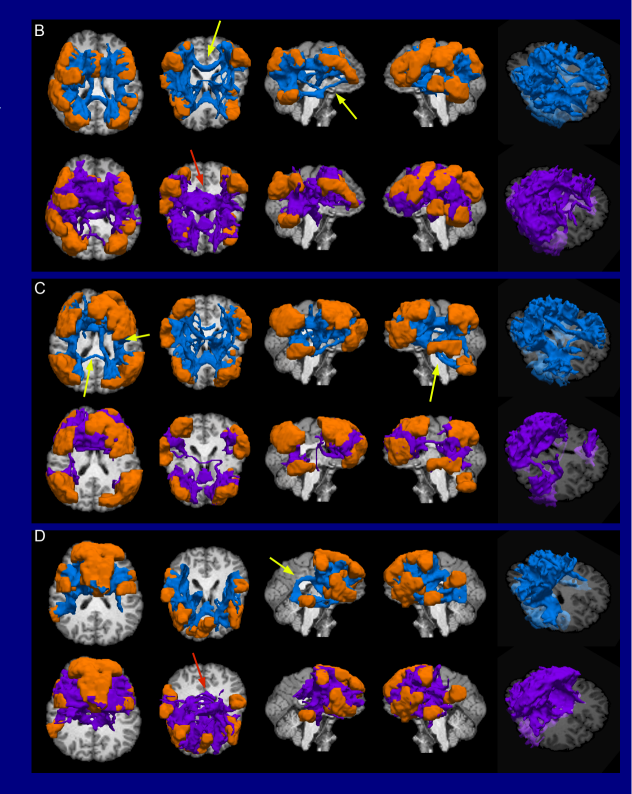


- + generally similar connections, but FSL bigger blobs
- + FSL took several hours for uncertainty, and then >24 hours for tracking this single network (and had to run 4 for this study)
- + 3dDWUncert took 7min; 3dTrackID took 25mins total for 4 netw.

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)

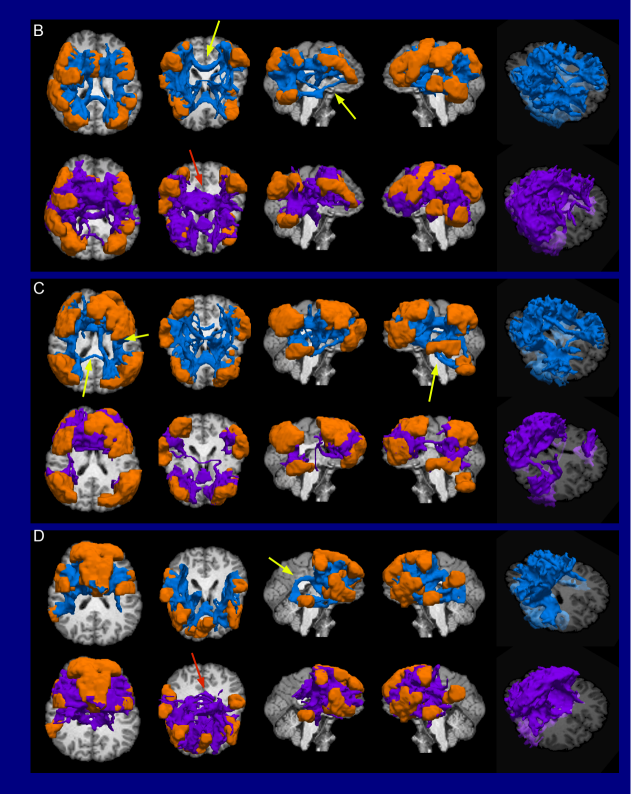


3dTrackID:

(other networks show similar results in terms of:

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

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

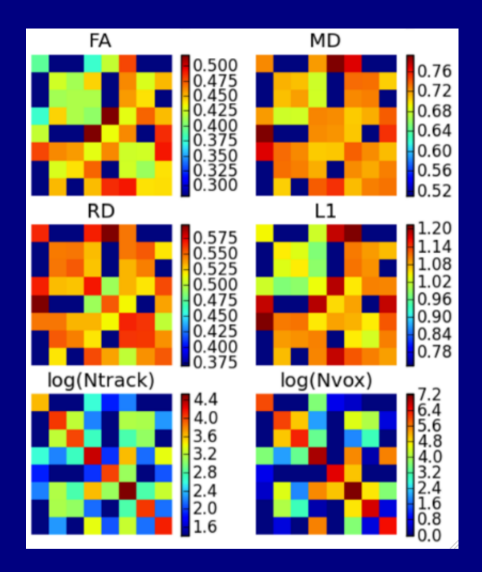


WM (ROI) Quantities

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.



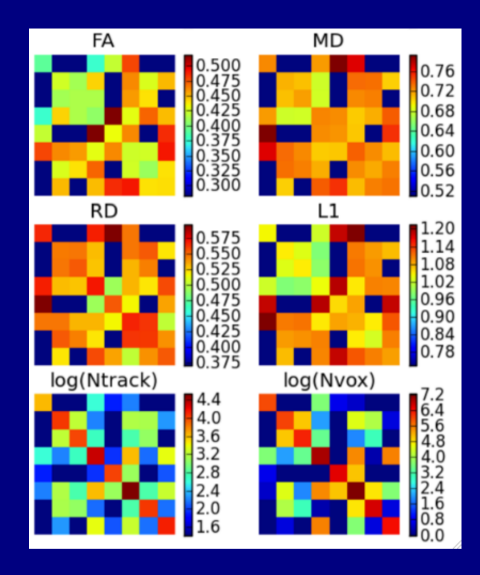
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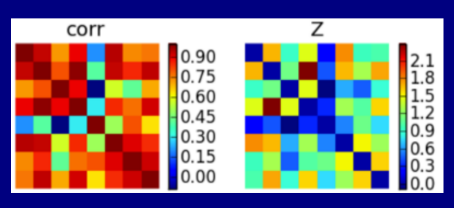
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3dNetCorr: correlation matrices Of average time series in ROIs (e.g., uninflated GM ROIs from 3dROIMaker)





Example: Group analysis with tracking output using multivariate statistics

from study:

<u>A DTI-Based Tractography Study of Effects</u>

<u>on Brain Structure Associated with</u>

<u>Prenatal Alcohol Exposure in Newborns,</u>

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



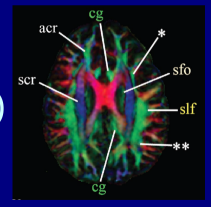
Goals of this study

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



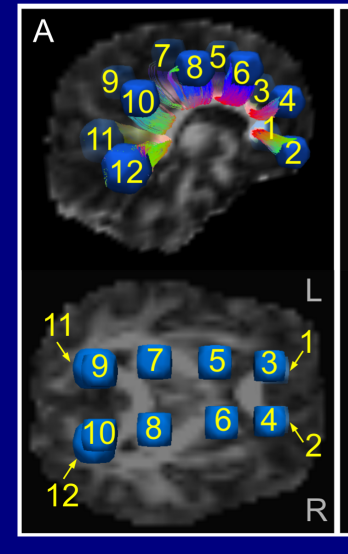
Setting up DTI-tractography

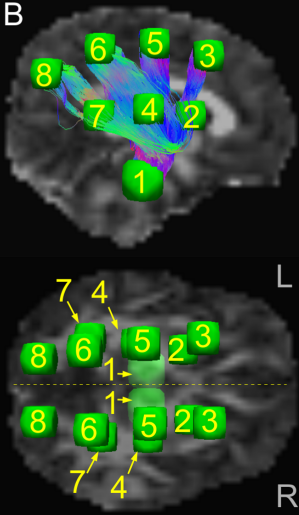
Location of targets for tractography: 5 WM networks.

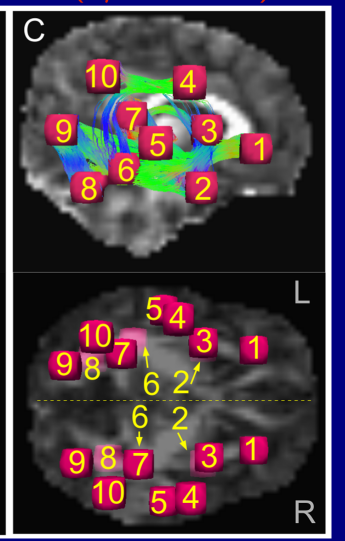
CC and Cor. Rad. (CCCR)

Projection (L/R-PROJ)

<u>Association</u> (L/R-ASSOC)



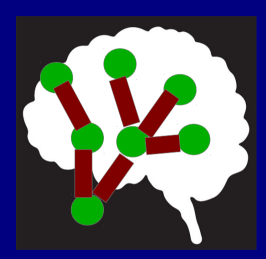




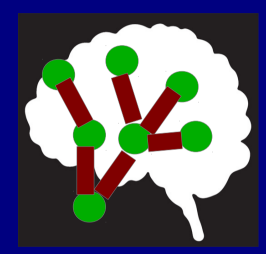
1) Place network targets



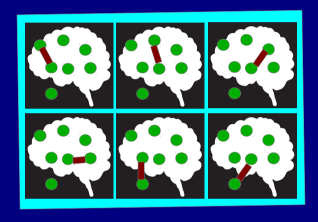
- 1) Place network targets
- 2) Probabilistic tracking



- 1) Place network targets
- 2) Probabilistic tracking

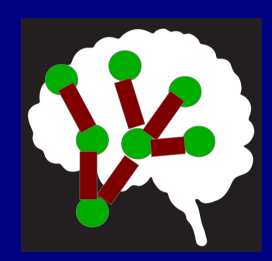


3) set of WM ROIs → set of repeated measures



- 1) Place network targets
- 2) Probabilistic tracking
- 3) set of WM ROIs \rightarrow set of repeated measures







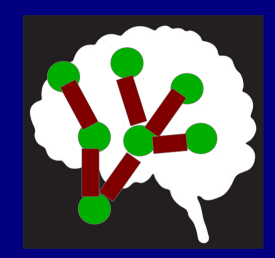
- 4) Multivariate model
 - {FA₁, FA₂, FA₃, ...}
 - alc
 - infant age
 - infant sex
 - maternal age
 - maternal cig/day

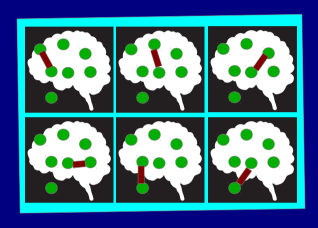


Analysis Steps

- 1) Place network targets
- 2) Probabilistic tracking
- 3) set of WM ROIs \rightarrow set of repeated measures







- 4) Multivariate model
 - {FA₁, FA₂, FA₃, ...}
 - 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

- + 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.

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).

The questions:

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

Parameters showing at least trends $(p<0.1) \rightarrow$

		FA				MD				AD				PD		
Network	var.	$\boldsymbol{\beta}_{med}$	$F(df_N, df_D)$	р	var.	β_{med}	F (df _N , df _D)	р	var.	β_{med}	F (df _N , df _D)	р	var.	β_{med}	$F(df_N, df_D)$	p
CCCR					alc	-0.70	8.6 (1, 14)		alc	-0.72	14.0 (1, 14)	0.002**				,
									cig	-0.27	2.5 (6, 9)	0.101	cig	0.47	3.5 (1, 14)	0.083
					mat_age	0.56	5.5 (1, 14)		mat_age	0.53	6.3 (1, 14)	0.025*				
L-PROJ					alc	-0.41	3.9 (10, 140)	0.000***	alc	-0.52	4.1 (10, 140)	0.000***				
	cig	0.12	4.2 (11, 4)	0.091									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*				
					-1-	0.44	4.0 (40, 400)	0.005*	-1-	0.45	0.7 (40, 400)	0.000++				
R-PROJ					alc	-0.41	1.9 (12, 168)	0.035*	alc	-0.45	2.7 (12, 168)	0.002**	oia	0.49	2.4.(4.4.4)	0.005
		0.00	0.0 (40.0)	0.400		0.44	F O (4, 44)	0.004*		0.00	F O (4 44)	0.000*	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)		age	-0.39	5.3 (1, 14)	0.038*				
	mat ana	0.46	0.2 (42.2)	0.402	sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
	mat_age	-0.16	9.2 (13, 2)	0.103	-1-	0.05	0.0 (7.0)	0.044*	-1-	0.00	0.4 (4.44)	0.040*				
L-ASSOC					alc	-0.65	6.0 (7, 8)	0.011*	alc	-0.66	8.1 (1, 14)	0.013*	oia	0.40	26 (4 44)	0.000
										0.40	0.5 (0.04)	0.000*	cig	0.49	3.6 (1, 14)	0.080
							00(4.44)		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*				
D 46606	ala	0.22	1 9 /7 09)	0.000	ala	0.60	10.0 (4. 14)	0.007**	ala	0.67	111(111)	0.000**				
R-ASSOC	alc	0.23	1.8 (7, 98)	0.090	alc	-0.62	10.2 (1, 14)		alc	-0.67	14.1 (1, 14)	0.002**	cia	0.5	3 5 (1 14)	0.082
	l								cig	-0.29	3.9 (1, 14)	0.068	cig	0.5	3.5 (1, 14)	0.082

^{*} p<0.05; ** p<0.01; *** p<0.001.

The questions:

- 1) which WM networks are affected by PAE?
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Parameters showing at least trends $(p<0.1) \rightarrow$

•			FA				MD				AD				PD		
Y	Network	var.	$oldsymbol{eta}_{med}$	F (df _N , df _D)	р	var.	β_{med}	F (df _N , df _D)	р	var.	β_{med}	F (df _N , df _D)	р	var.	β_{med}	$F(df_N, df_D)$	р
5(CCCR					alc	-0.70	8.6 (1, 14)	0.011*	alc	-0.72	14.0 (1, 14)	0.002**				
Z										cig	-0.27	2.5 (6, 9)	0.101	cig	0.47	3.5 (1, 14)	0.083
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บ	L-PROJ					alc	-0.41	3.9 (10, 140)	0.000***	alc	-0.52	4.1 (10, 140)	0.000***				
2		cig	0.12	4.2 (11, 4)	0.091									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*				
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				0.0 (40.0)				5 6 (4 4 4 4)						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*				
			0.40	0.0 (40.0)		sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
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														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	-0.29	3.9 (1, 14)	0.068	cig	0.5	3.5 (1, 14)	0.082

^{*} p<0.05; ** p<0.01; *** p<0.001.

 → Statistically significant alcohol exposure associations in ~every
 WM network

Networks

The questions:

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

Parameters showing at least trends $(p<0.1) \rightarrow$

		FA				MD				AD				PD		
Network	var.	FA 2	$F(df_N, df_D)$	р	var.	β _{med}	F (df _N , df _D)	р	var.	β_{med}	F (df _N , df _D)	р	var.	β_{med}	$F(df_N, df_D)$	р
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					_				_							
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													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*				
					sex	-0.20	4.3 (1, 14)	0.056	sex	-0.39	5.9 (1, 14)	0.029*				
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									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*				
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	<u> </u>								cig	-0.29	3.9 (1, 14)	0.068	cig	0.5	3.5 (1, 14)	0.082

^{*} p<0.05; ** p<0.01; *** p<0.001.

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

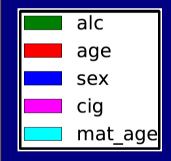
III) Results: ROI level

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).

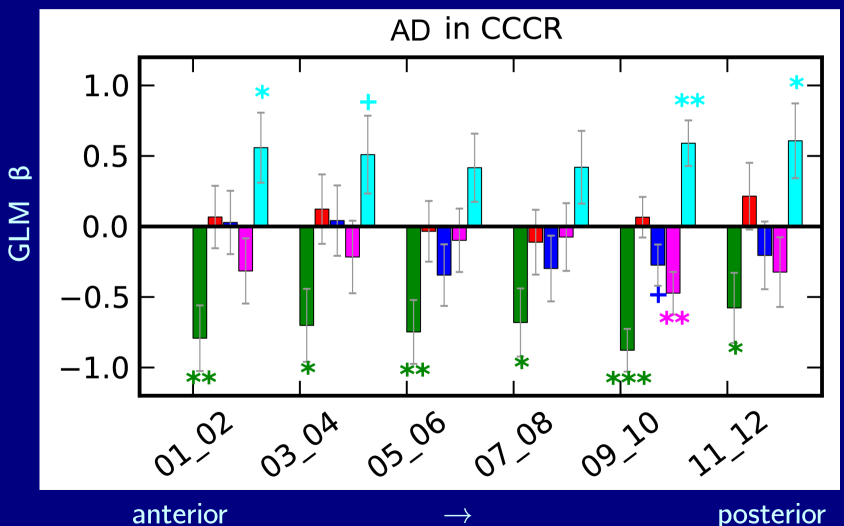


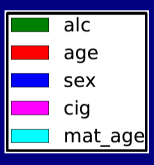
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Transcallosal (CC and corona radiata)





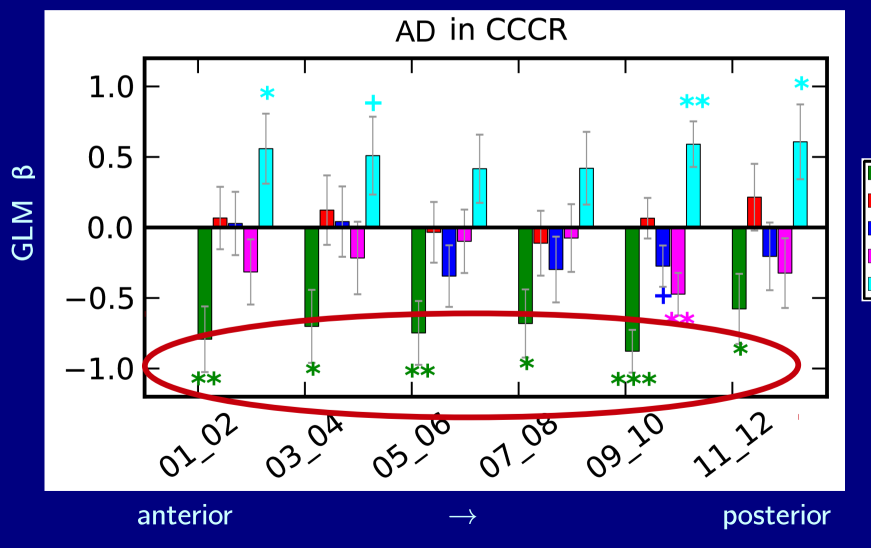
posterior

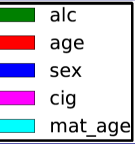
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The question:

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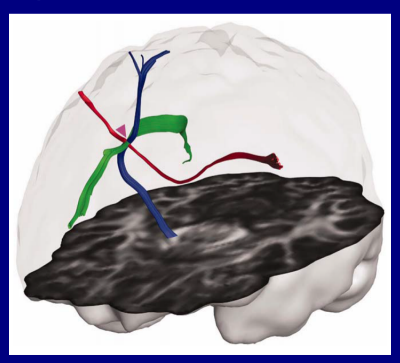


→ strong AD-alc relations in most (medial) WM ROIs Example: HARDI tracking

Higher order models

DTI tractography:

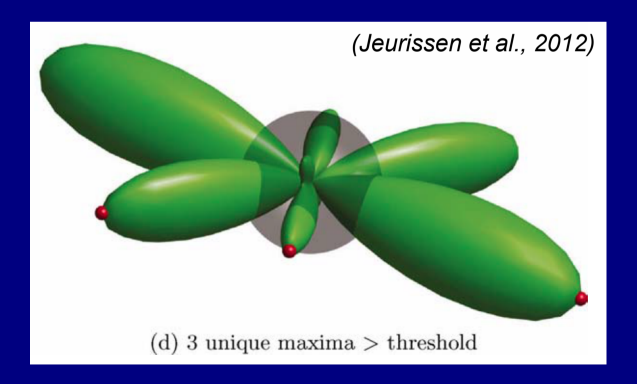
- + susceptible to false negatives, difficulty with long range tracts (noise/error accumulation)
- + Major diffusion can be average of multiple paths
- + Voxels can have low FA from several WM paths, false ending
- + Can't resolve complex underlying architecture
 - Jeurissen et al. (2012, HBM): 60-90%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



(Jeurissen et al., 2012)

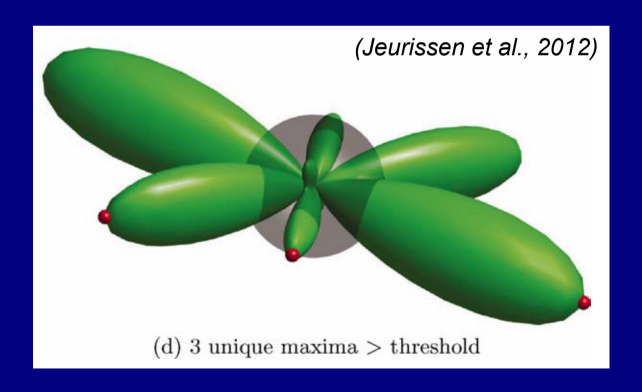
HARDI

- + 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 → lower signal, so susceptible to noise



HARDI

- + High Angular Resolution Diffusion Imaging:
 - DSI, ODF, Qball, FOD...
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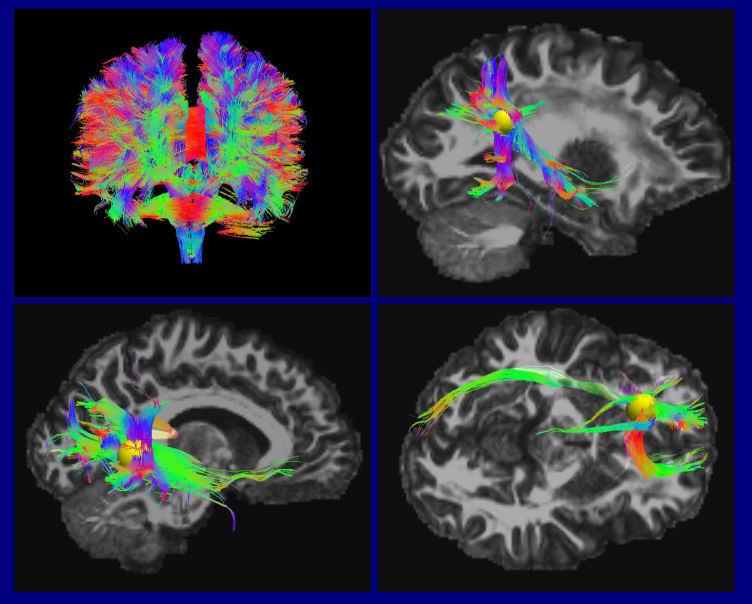


FATCAT can now track through HARDI data

 → HARDI reconstruction done outside AFNI (e.g., DSI-Studio, Diffusion Toolkit, FSL), and outputs tracked in FATCAT.

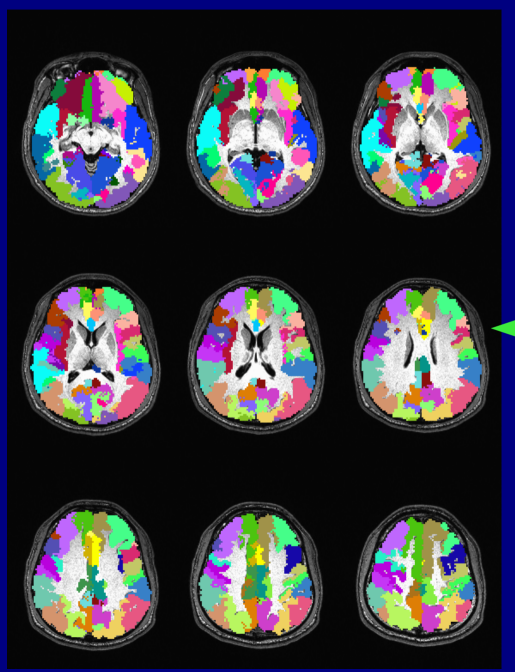
Example: 3dTrackID on HARDI data

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



Example: 'Connectome'-type tracking

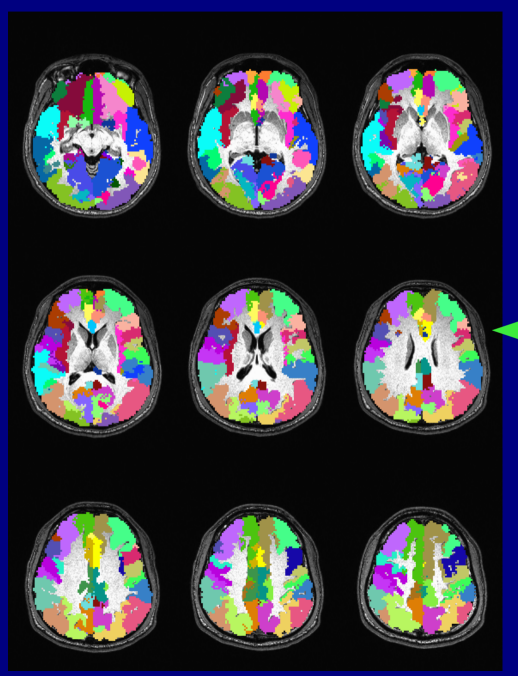
"Connectome": parcellation of GM



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

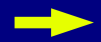
"Connectome": parcellation of GM



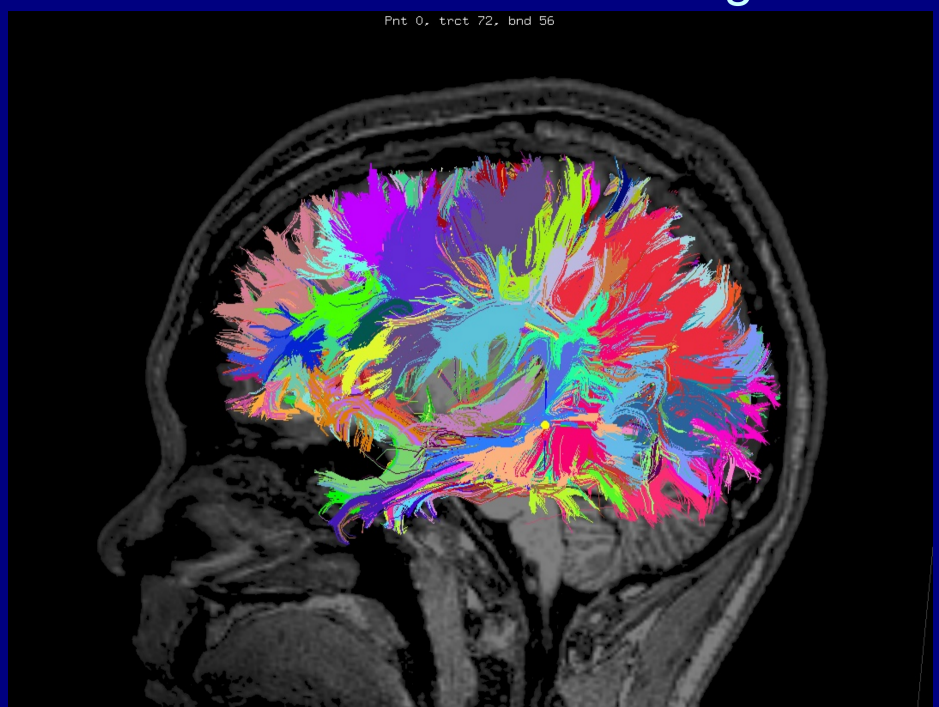
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- by 1 voxel, up to FA>0.2.
- (+ NEW: keep labeltable labels and use them in output.)
- + '3dTrackID -mode DET' among the regions

and a few seconds later...

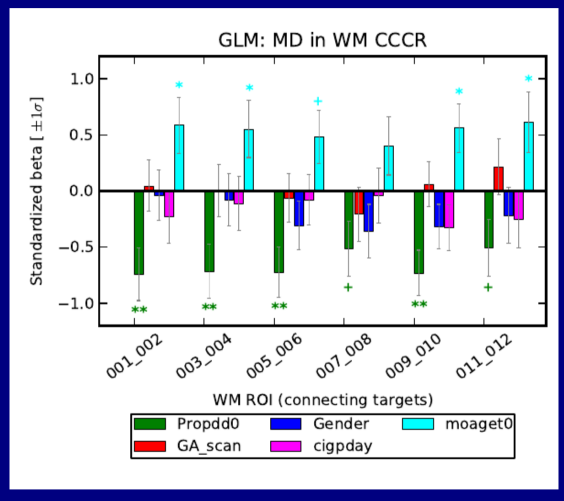


"Connectome": tracking



A brief example for statistical analysis

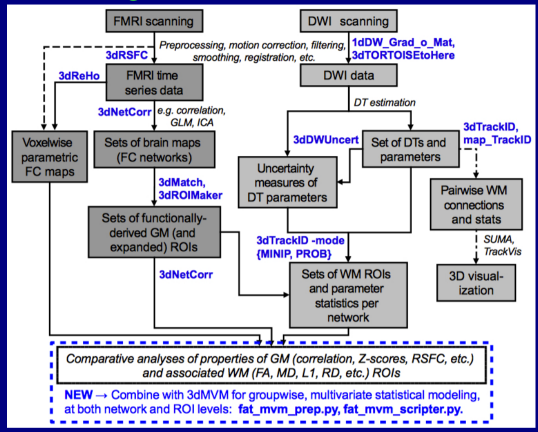
+ Combining tractography, quantitative DTI and subject measures with GLM to find structure-alcohol consumption relation:



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.

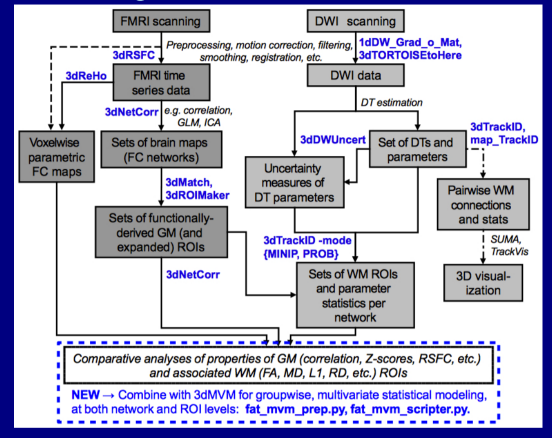
We have discussed capabilities and benefits of:

Combining multimodal data: FC+SC+..

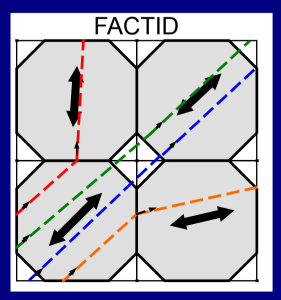


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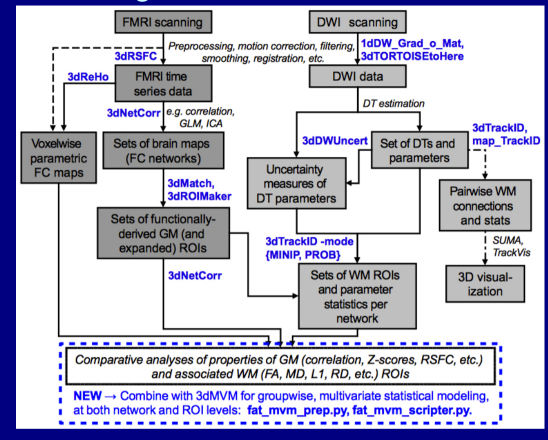


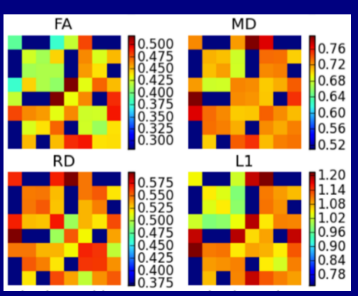
Using an efficient algorithm, reduced bias of propagation



We have discussed capabilities and benefits of:

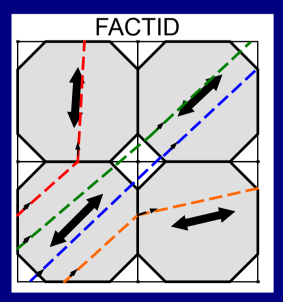
Combining multimodal data: FC+SC+...





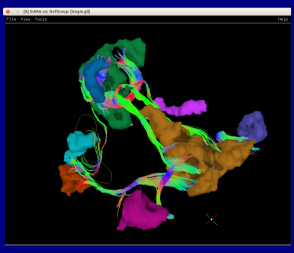
Using an efficient algorithm, reduced bias of propagation

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



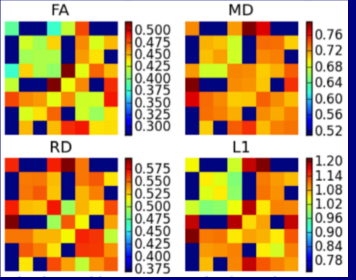
We have discussed capabilities and benefits of:

Integrating AFNI-SUMA visualization

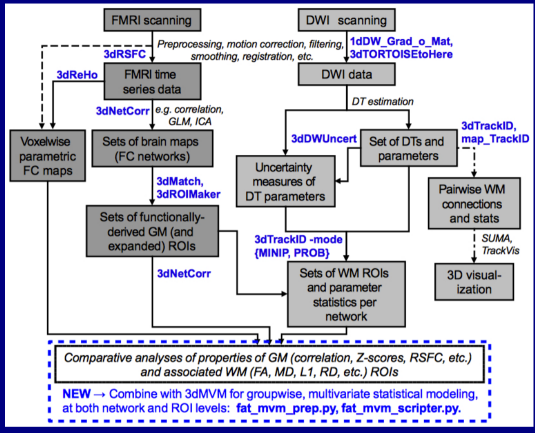






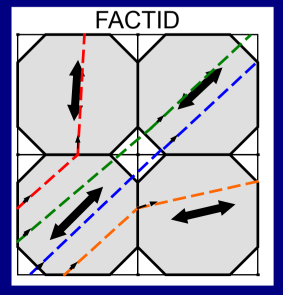


Combining multimodal data: FC+SC+...



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