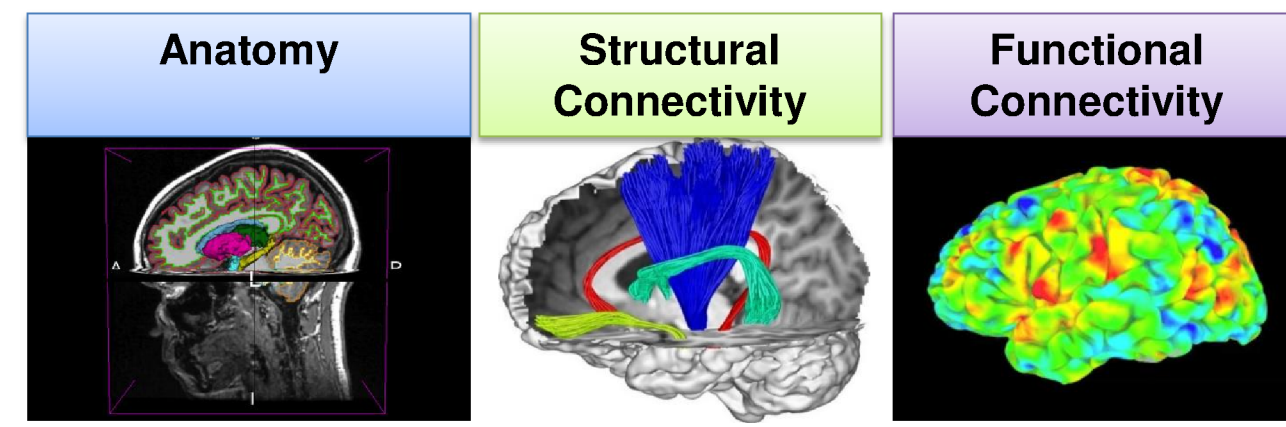


Impact of brain parcellation on parameter optimization of the whole-brain dynamical models

Introduction

- Resting-state (RS) functional connectivity (FC) analysis has brought new insights to the inter-individual variability and the pathophysiology of brain disorders [1,2].
- We constructed model networks based on the empirical structural connectivity (SC) and the simulation results are compared with empirical functional data.
- We considered two brain atlases and brain parcellations and evaluated their impact on the dynamics of the whole-brain models.



The computational model: Kuramoto

We use a computational Kuramoto model of coupled phase oscillators to simulate the dynamics of the resting-state (RS) brain networks [5]. The phase θ_n of node n at time t , obeys the following dynamical equation [6]:

$$\frac{d\theta_i}{dt} = \omega_i + K \sum_{j=1}^N C_{ij} \sin[\theta_j(t - \tau_{ij}) - \theta_i] + \eta_i(t), \quad i = 1, \dots, N.$$

Model variables	Description	Model variables	Description
θ_i	phase of node i at time t	$\eta_i(t)$	noise term
K	global coupling strength	$L_{ij}, (L)$	relative, mean fiber length
C_{ij}	relative coupling strength from node j to node i	V	conduction speed
$\tau_{ij} = \frac{L_{ij}}{V} = \langle \tau \rangle L_{ij} / (L)$	time delay between node j to node i	$\langle \tau \rangle$	mean time delay
$f_i = \omega_i / 2\pi$	intrinsic frequency of node i on its limit cycle ($f = 10$ or 60 Hz)	$r_i = \sin[\theta_i(t)]$	neural activity

Structural, Diffusion & Functional data preprocessing

We used 50 healthy subjects from the **Human Connectome Project** [3] database



Magnetic Resonance Imaging protocol:

- Filtered blood-oxygen-level dependent (BOLD) time series are extracted from the FIX denoised RS data in MNI152 template space
- Parcellation-based empirical FC matrices: from the mean BOLD signals extracted for each brain regions after 5mm and without spatial smoothing (mean and 1st eigenvariate time series)

Structural pipeline [4]:

- Parcellation: Schaefer & Harvard-Oxford brain atlases
- Software method: FreeSurfer
- Motion/eddy correction: ✓
- Intensity normalization: ✓
- Tractography: Probabilistic (MRtrix 3.0)
- SC Metric: Voxel pairs connected with 10⁶ streamlines, ROI volume corrected



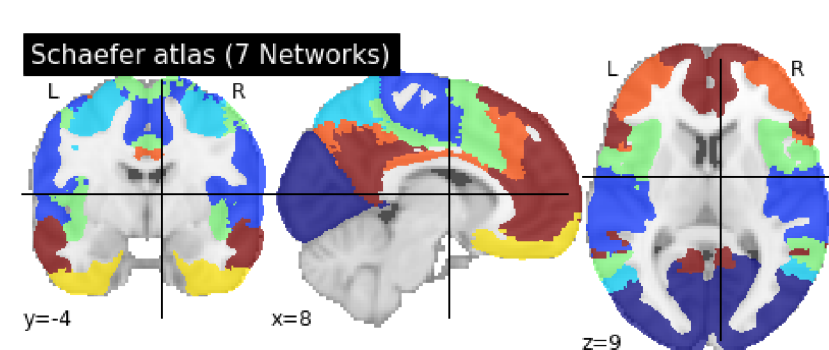
Parameter Sweep Exploration

- Essential step in order to fit simulations with empirical data.
- Performed using **tvb-hpc** (CUDA) [7].
- TVB model kernels optimized for HPC on hybrid architectures.
- GPU code allows thousands of parameters to be explored in parallel: each parameter is assigned to a thread in the GPU.
- Global coupling (K) and mean time delay (τ) are varied to maximize FC/SC correlation with simulated FC.
- Runs are performed on the JURECA GPU partition (Research Centre Jülich).

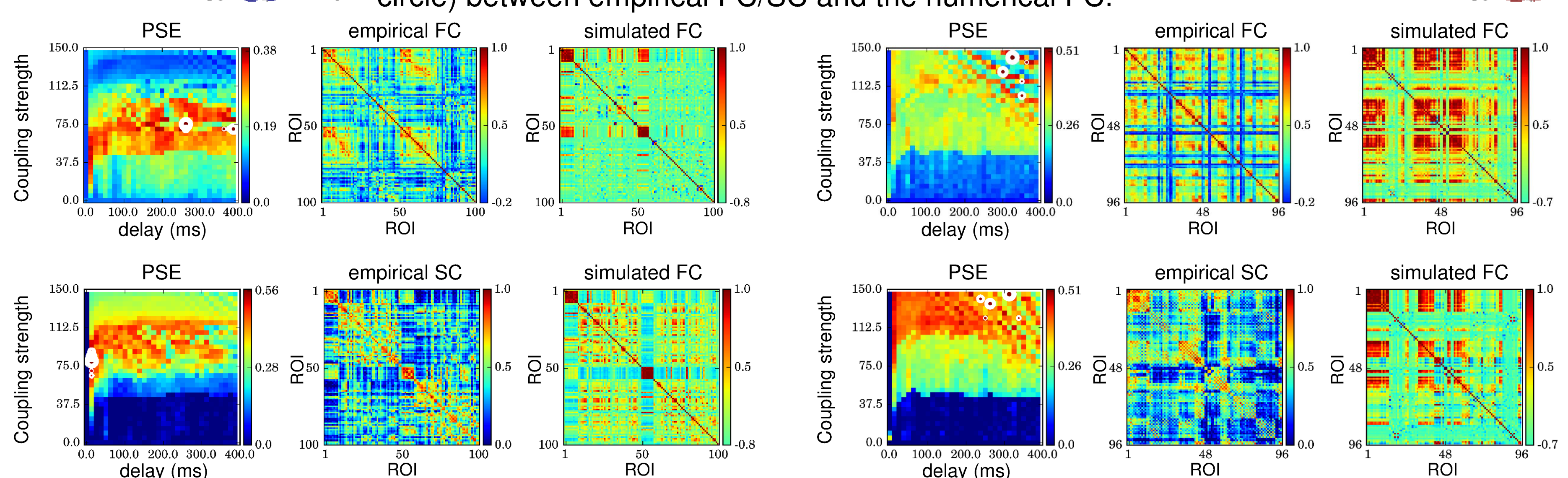
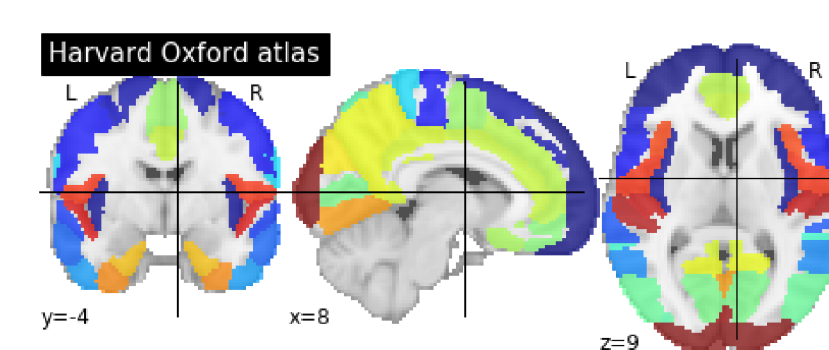


Results: numerical simulations

Parameter sweep exploration (PSE) for Schaefer and Harvard-Oxford atlases ($f_i = 60$ Hz – FC : mean over region – a case example)

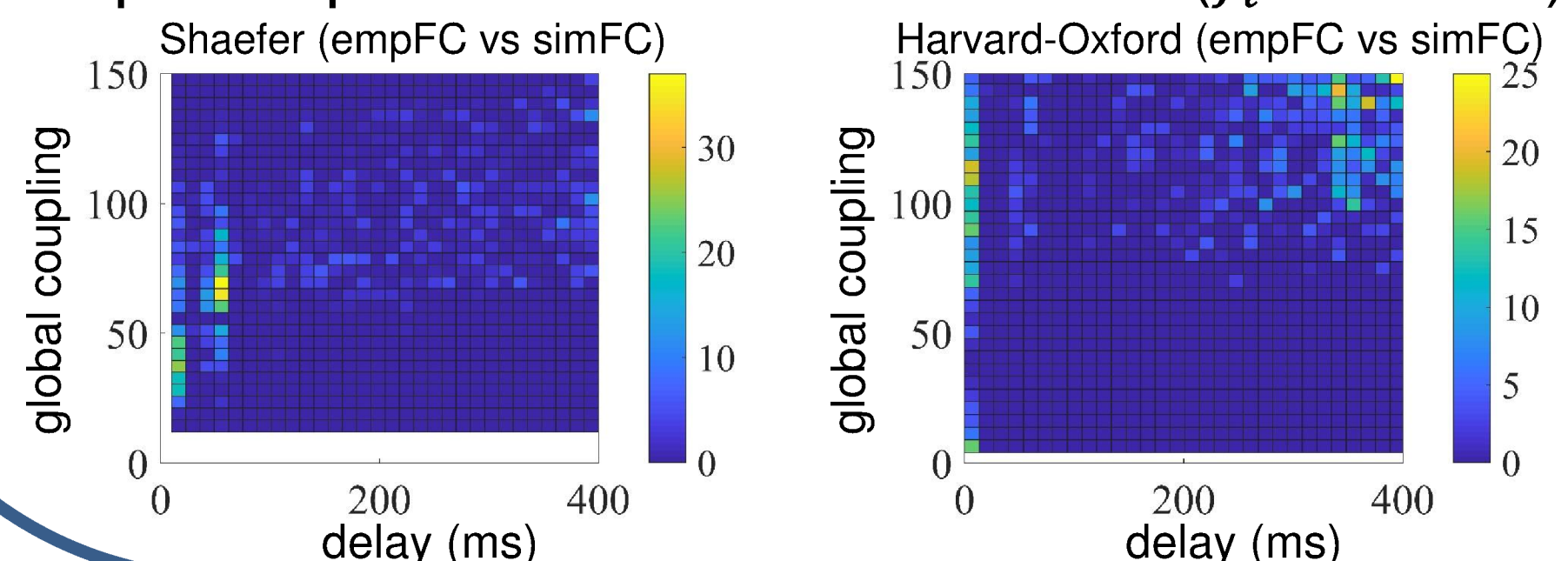


The depicted simulated FC matrices correspond to those models with coupling strength and delay parameter values with maximal Pearson's correlation coefficient value (larger white circle) between empirical FC/SC and the numerical FC.

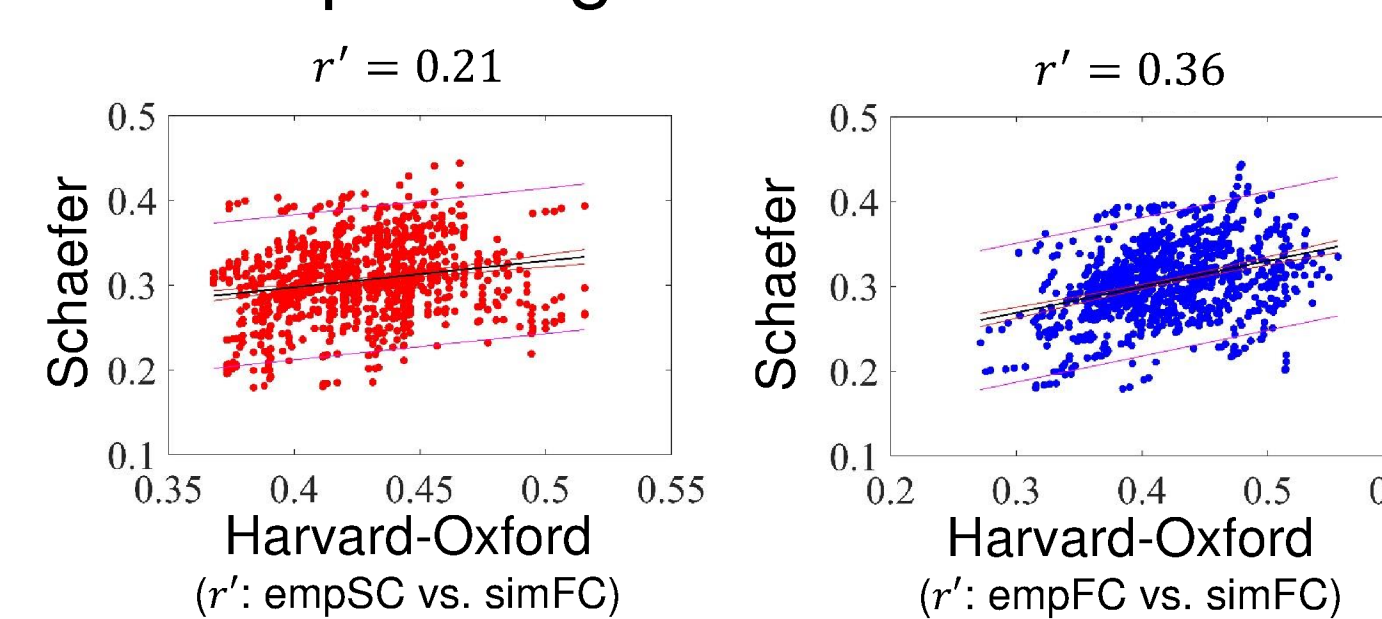


(white circles: 5 maximum Pearson's correlation coefficient values / 30s of simulated electrical activity)

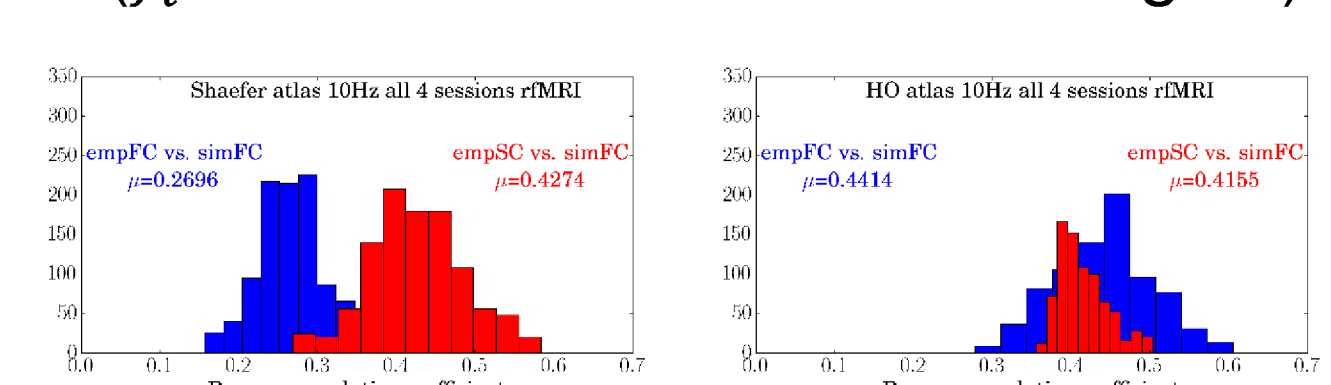
Optimal parameter 2D distributions ($f_i = 60$ Hz)



Corresponding correlation coefficients (r')

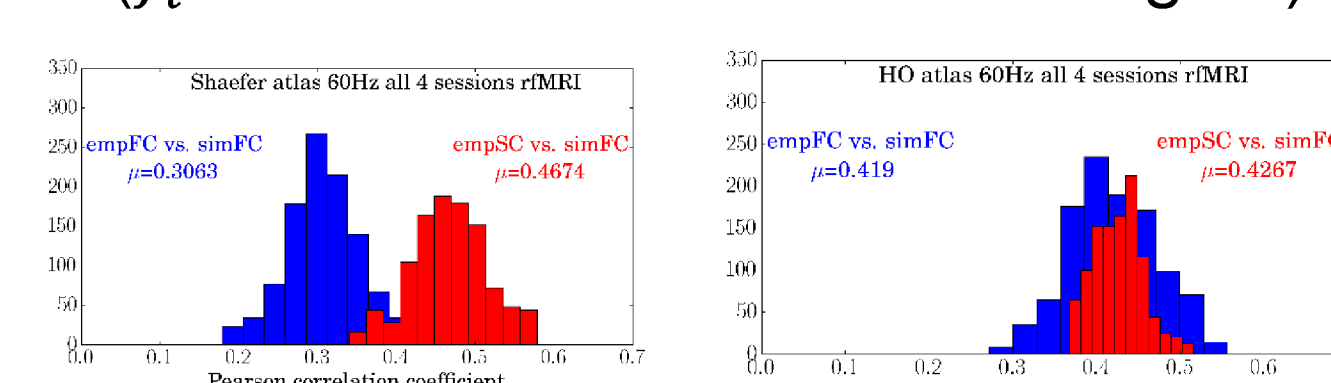


Schaefer vs Harvard-Oxford ($f_i = 10$ Hz – FC: mean over region)



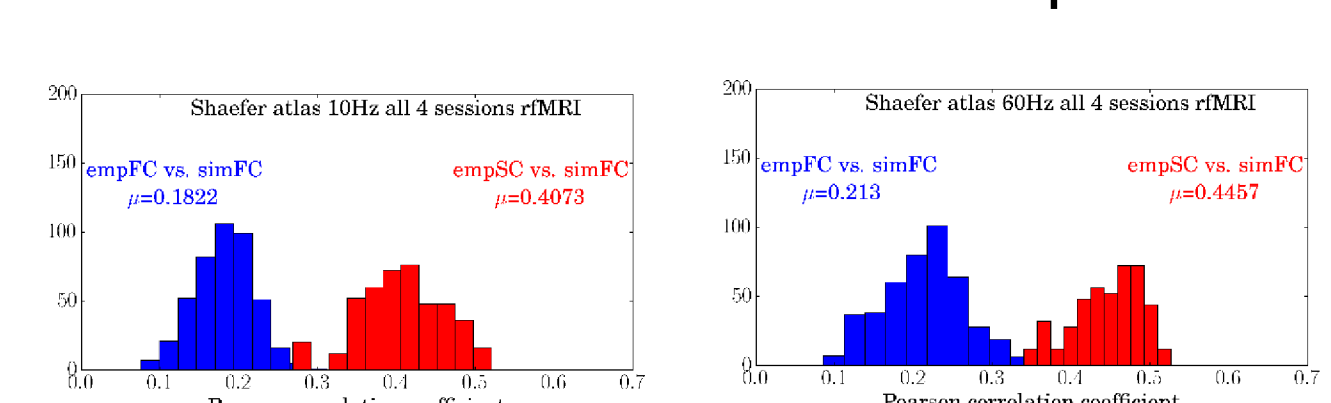
of subjects: 50
of sessions: 4

($f_i = 60$ Hz – FC: mean over region)



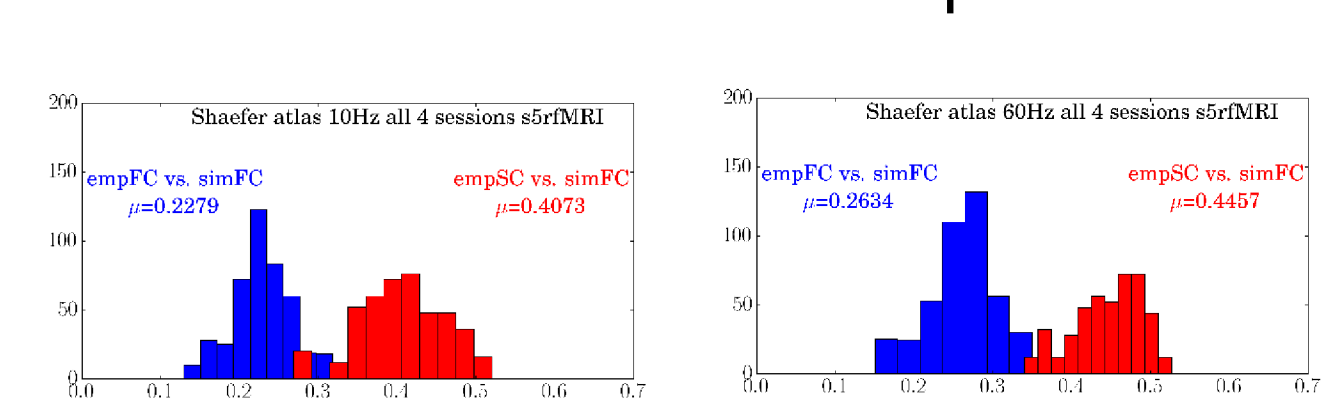
Using the 5 maximal Pearson's correlation coefficient values (4 sessions per subject) from the parameter sweep exploration analysis (white circles in the PSE figures)

Schaefer ($f_i = 10, 60$ Hz – FC: 1st eigenvariate without spatial smoothing)



of subjects: 23
of sessions: 4

($f_i = 10, 60$ Hz – FC: 1st eigenvariate with spatial smoothing)



Summary

- We calculated the SC matrices and FC matrices (mean and 1st eigenvariate over region with/without spatial smoothing) for 50 HCP subjects.
- We simulated resting-state network dynamics (mean node electrical activity) using the Kuramoto model (with $f_i = 10, 60$ Hz).
- We performed parameter sweep exploration for global coupling and mean time delay to maximize empirical FC/SC correlation with simulated FC.
- No observable advantage was found when using 1st eigenvariates instead of mean value from the BOLD time series.
- We produced 2D distributions for the optimal parameters for both atlases.
- We found relatively strong correlations ($r \geq 0.35$) between emp. FC and sim. FC matrices, whereas the correspondence between emp. SC and sim. FC matrices is, however, weaker ($r \geq 0.20$) for both atlases.

Outlook ➤ Add more subjects, atlases, models, refine parameter space intervals, explore different empirical to simulated measurements

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Acknowledgments

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