

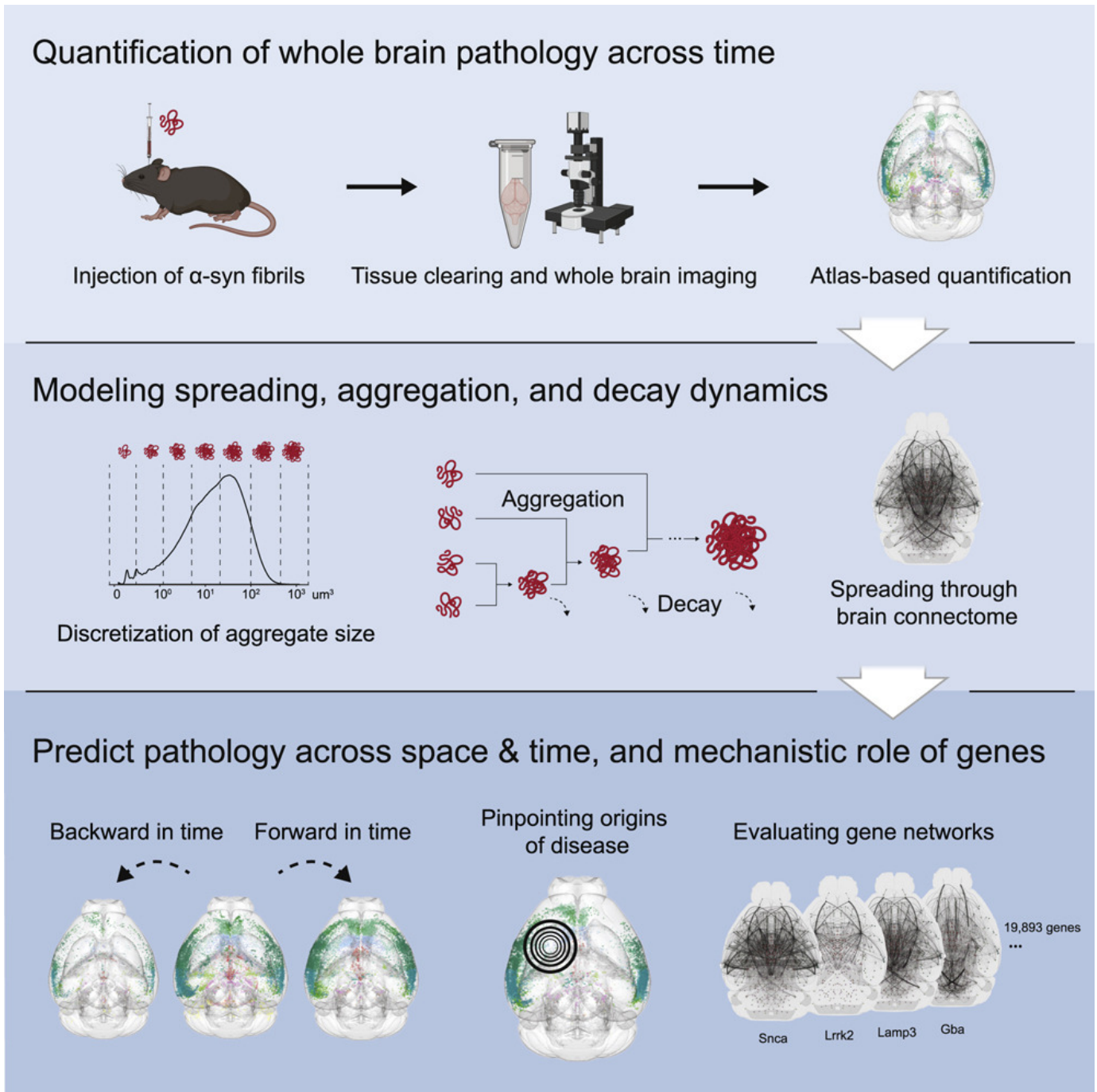
**Docket #:** S22-403

# **Solving brain circuit function and dysfunction with computational modeling and optogenetic fMRI**

Stanford researchers at the Lee Lab have developed a method to understand whole-brain circuit mechanisms underlying neurological disease and its application to predict the outcome of therapeutic interventions.

By combining optogenetic fMRI with computational modeling, cell type-specific, large-scale brain circuit function and dysfunction are now starting to be quantitatively parameterized. These new findings can pave the path for future therapeutics developments based on a systems engineering approach aimed at directly restoring brain function.

**Figure:**



**Figure description: Graphical Abstract**

Image credit: <https://doi.org/10.1016/j.celrep.2022.111631>

**Stage of Development**

- Proof of Concept

**Applications**

- **Neuromodulation therapy design** for neurodegenerative diseases
- The method can be applied to predict the optimal targets and parameters of the neuromodulation treatments for a number of neurological disorders, including but not limited to Parkinson's disease (PD), dystonia, and epilepsy

## Advantages

- **Systematically designed treatments for brain disorders**
- **Enables virtual neuromodulations** for the treatments of neurological disorders
- By implementing such virtual neuromodulations, rapid optimization and customization can be achieved that **saves the time and financial costs of carrying out *in vivo* experiments**
- **Incorporates cell-type information to modeling**

## Publications

- Lee, Jin Hyung, Qin Liu, and Ehsan Dadgar-Kiani. ["Solving brain circuit function and dysfunction with computational modeling and optogenetic fMRI."](#) *Science* 378, no. 6619 (2022): 493-499.
- Dadgar-Kiani, Ehsan, Gregor Bieri, Ronald Melki, Aaron D. Gitler, and Jin Hyung Lee. ["Mesoscale connections and gene expression empower whole-brain modeling of  \$\alpha\$ -synuclein spread, aggregation, and decay dynamics."](#) *Cell Reports* 41, no. 6 (2022): 111631.

## Patents

- Published Application: [WO2024073593](#)

## Innovators

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