Dyson SIS Code Tutorial and Reproducibility Notes

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1 Overview

This document accompanies the *dyson-code* repository and explains how to reproduce the figures and explore some model variations. The code implements a Susceptible-Infected-Susceptible (SIS) model on a graph and computes a Dyson similarity transformation that approximately symmetrizes the generator matrix. The primary entry points are in src/dyson.jl and runnable scripts in scripts/ that generate the paper figures into results/figures/.

Key components

- default_initialisation() prepares a default complete graph with 6 nodes, default rates, and either loads or computes the Dyson transform.
- get_SIS_H(A, beta, gamma; incl_disease_free) builds the SIS generator given adjacency matrix A and rates.
- find_hermitian(H; tol, kmax) iteratively constructs h, η with reduced asymmetry.
- simulate(H, eta; symmetrize, tspan, saveat) integrates x'(t) = -H'x with optional similarity transform.
- Multiple functions to provide observables and statistics of the system.

2 Quick start

2.1 Prerequisites

• Julia 1.11.7 with the packages pinned by Project.toml

2.2 Reproducing all figures

From the repository root:

\$ bash run_all.sh

This runs every script in scripts/ and writes figures to results/figures/.

3 Core API and Usage

3.1 API Overview

The core functionality lives in src/dyson.jl. Below is a narrative overview of the exported functions used throughout the scripts, how to apply them in your own work, and when to prefer each. For a technical summary of arguments and return values, refer to the in-code documentation (docstrings) in src/dyson.jl.

default_initialisation() Loads cached defaults or generates them if missing: a complete graph with six nodes, SIS generator H, Hermitized matrix h, transformation η , rates β, γ , a flag incl_disease_free, and the convergence errors. Use this at the top of a script to get a reproducible baseline without waiting for recomputation. Artifacts are cached under results/intermediate/. If you want to repeat the experiments for an other system (dynamics, or network), this function serves as a great jumping-off point.

disease_free_initialisation() Alternative initializer identical to default_initialisation() but with the disease-free state included (incl_disease_free=true). This is useful for exploring regimes where the absorbing disease-free state meaningfully affects statistics or when you want to visualize transitions that cross the absorbing state. It caches its own intermediate file under a different name (full-6-wdf). Scripts such as scripts/transition.jl can adopt this initializer to produce ..._transition.pdf for the disease-free-inclusive model.

```
name, g, A, H, h, eta, beta, gamma, incl_disease_free, errors = default_initialisation()
```

get_SIS_H(A, beta, gamma; incl_disease_free=false) Constructs the SIS generator consistent with the chosen graph (represented by adjacency matrix A) and rates. Rows sum to zero by construction, and the disease-free state can be excluded, such that only a single steady-state exists. Use this when sweeping β/γ (see scripts/transition.jl) or when you want to try a different graph.

```
H = get_SIS_H(A, beta, gamma; incl_disease_free=false)
```

find_hermitian(H; tol, kmax, verbose=false) Iteratively finds a similarity transform that reduces the asymmetry of H, returning the transformed matrix h, the transform η , and an error history. Use when you need a near-Hermitian representation for analysis or optimization (cf. scripts/steady_state.jl).

```
h, eta, errors = find_hermitian(Matrix(H); kmax=5_000)
```

SIS_initial_state(n) Provides the default initial state used in simulations: one infected individual mapped by η . Override this if you need a different starting distribution (e.g., multiple infections or a distribution over states).

```
u0 = SIS_initial_state(size(H, 1))
```

simulate(H, eta; symmetrize, tspan, saveat, initial_func) Integrates the dynamics x'(t) = -H'x under the similarity transform η . Set symmetrize=true to simulate the symmetrized transformed matrix. This changes the system depending on how non-Hermitian H is. However it ensures that the dynamics are reversible. Supply a custom initial_condition to explore different initial states.

```
sol = simulate(H, I; tspan=(0.0, 5.0), saveat=0.05)  # original system
sol_h = simulate(H, eta; symmetrize=true)  # transformed
sol_custom = simulate(H, eta; initial_condition=(H,eta)->(eta*rand(size(H,1))))
```

get_steady_state(H; eta=I, symmetrize=false) Computes a steady state by forward integration. In the Hermitized frame, combine with η for the transformed steady state as in scripts/steady_state.jl.

```
P_ss = get_steady_state(H)
phi_ss = get_steady_state(h; symmetrize=false) # pass eta via keyword in your code
```

infectious_proportion_0(n; incl_disease_free=false) Returns the observable measuring the infected fraction. Use with observe on either the original state distribution p or the transformed state ϕ with metric Ω .

```
0 = infectious_proportion_0(n; incl_disease_free=false)
I_mean = observe(0, p)
Omega = inv(eta*eta')
I_mean_transformed = observe(eta*0*inv(eta), phi, Omega)
```

observe(0, p) and observe(0, phi, 0mega) Two observation modes: apply 0 to the probability vector p, or to a transformed state ϕ with the appropriate metric Ω . See scripts/statistics.jl for mean/variance time series in both frames.

shannon_entropy(p) and renyi_entropy(p) Entropy measures used for the entropy figures and transition curves. Apply directly to state vectors from simulations or steady states. For details (e.g., numerical safeguards near zeros), see the function docstrings.

3.2 Workflow

In practice, the workflow is: build or load H, obtain h, η via find_hermitian, simulate either original or transformed dynamics using simulate, and compute observables with observe and entropies with shannon_entropy/renyi_entropy. For precise signatures and edge cases, consult the code documentation in src/dyson.jl.

4 Scripts and outputs

Each script starts by including dyson-setup.jl, which activates the project, brings Dyson into scope, seeds RNG for reproducibility, and ensures results/figures/ exists. Every script is written to reproduce the figures of the paper, using the functionality of Dyson.

4.1 Matrix structure

Script: scripts/H_figure.jl

- Builds a color-coded visualization of the SIS generator's sparsity and transition types.
- Saves SIS_H_heatmap.pdf to results/figures/.

Adjustable parameters (Figure 2B)

- Color scaling and palette: edit the color ranges in the script for infections/healings/diagonals.
- Graph size/structure: modify default_initialisation() to change complete graph(6).

4.2 Entropy figures (Figure 1)

Script: scripts/entropy_figures.jl

- generate_homotopy(eta, sim_f, agg_f; steps) creates a homotopy $\eta(\alpha) = (1-\alpha)I + \alpha \eta$ and aggregates an observable (by default Rényi entropy).
- create_heatmap saves <name>_entropy_heatmap.pdf and create_entropies_figure saves <name>_entropy_graphs.pdf.

Adjustable parameters

- sim_f: adjust to perform a homotopy over other systems than the SIS one.
- agg_f: change to aggregate other observables, e.g., mean_0 or var_0.
- create_entropies_figure: modify to plot other statistics or change figure layout.
- create_heatmap: change the look of the figure, e.g., colormap or axis limits.

4.3 Statistics over time (Figure 3)

Script: scripts/statistics.jl

- Computes mean infection proportion and variance over time for original and transformed systems.
- Saves <name>_statistics_full.pdf.

Adjustable parameters

- tspan, saveat: change integration horizon and sampling density in simulate calls.
- symmetrize: switch between raw and symmetrized dynamics in the transformed system.
- Observable: replace infectious_proportion_0 with other operators to track different quantities.

4.4 Transition curves (Figure 4)

Script: scripts/transition.jl

- Sweeps ratios β/γ and records steady-state means, variances, and entropies (original and transformed).
- Saves <name>_transition.pdf and a JLD2 data file under results/intermediate/.

Adjustable parameters

- beta_range: adjust the logarithmic range of β/γ values.
- kmax, tol: control depth/precision of find_hermitian during the sweep.

4.5 Steady-state comparison (Figure 5)

Script: scripts/steady_state.jl

- Compares steady-states from Dyson-optimized distribution vs. dynamic evolution; annotates KS distance.
- Saves multiple files like <name>_SIS_ss_b001.pdf, ..._b2.pdf, ..._b200.pdf.

Adjustable parameters

- Rates: set specific β values relative to γ to probe different regimes.
- kmax: number of Hermitization iterations taken from default_initialisation().
- Plot aesthetics: toggle legend, change colors, layout, and bar alignment.

4.6 Algorithm convergence (Figure 6)

Script: scripts/convergence.jl

- Plots the error trajectory from find_hermitian in log scale.
- Saves <name>_convergence.pdf.

Adjustable parameters

- kmax: increase/decrease the number of iterations to control plot length.
- Styling: modify line width, colors, and axis labels.

5 Figure Gallery

Below we embed the figures produced by the scripts. If a file is missing, a placeholder box appears.

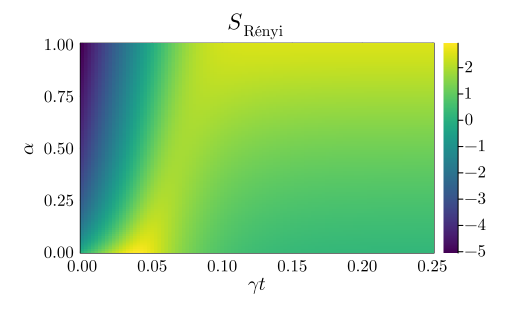


Figure 1: Entropy heatmap (Figure 1).

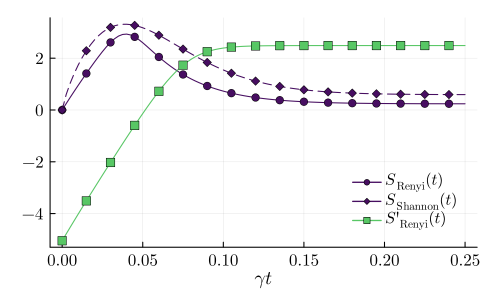


Figure 2: Entropy time series (Figure 1).

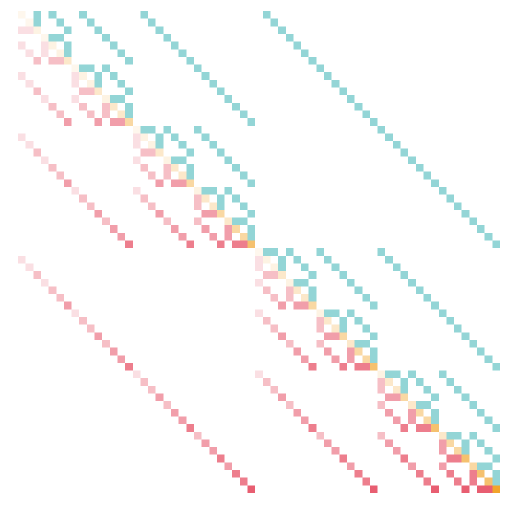


Figure 3: Structure of the SIS generator (Figure 2B).

6 Produced Figures

The scripts in scripts/ produce the following files in results/figures/:

- full-6-wodf_entropy_heatmap.pdf Entropy heatmap (Figure 1).
- full-6-wodf_entropy_graphs.pdf Entropy time series (Figure 2).
- SIS_H_heatmap.pdf Structure of the SIS generator (Figure 3).
- full-6-wodf_statistics_full.pdf Mean and standard deviation over time (Figure 4).
- full-6-wdf_transition.pdf (or full-6-wodf_transition.pdf if excluding the disease-free state) Transition curves over β/γ (Figure 5).
- full-6-wodf_SIS_ss_b001.pdf Steady-state bar plot at low β (Figure 6).
- full-6-wodf_SIS_ss_b2.pdf Steady-state bar plot at intermediate β (Figure 7).
- full-6-wodf_SIS_ss_b200.pdf Steady-state bar plot at high β (Figure 8).
- full-6-wodf_convergence.pdf Convergence of the Hermitization algorithm (Figure 9).

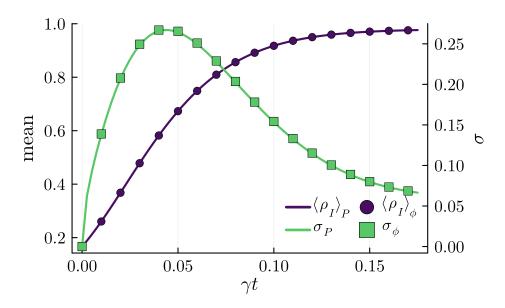


Figure 4: Mean and standard deviation over time (Figure 3).

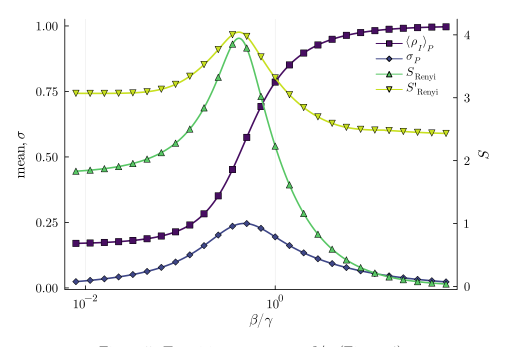


Figure 5: Transition curves over β/γ (Figure 4).

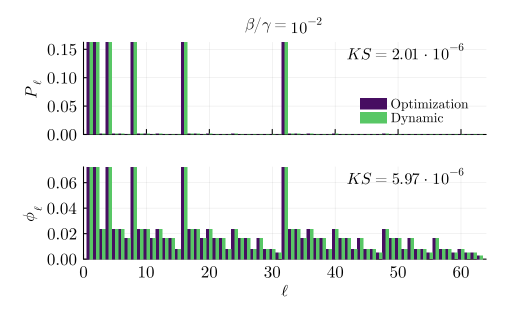


Figure 6: Steady-state comparison at low β (Figure 5A).

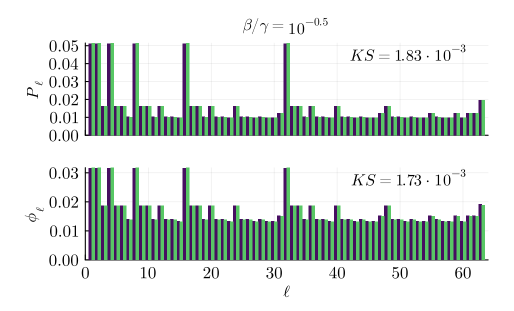


Figure 7: Steady-state comparison at intermediate β (Figure 5B).

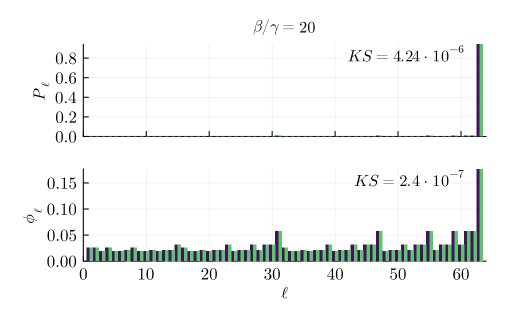


Figure 8: Steady-state comparison at high β (Figure 5C).

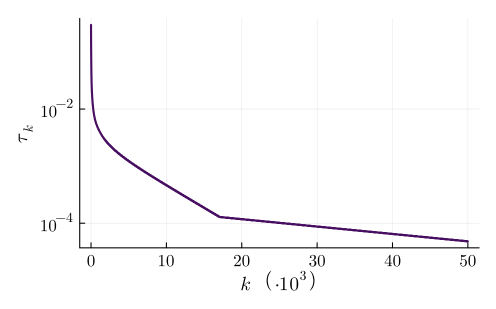


Figure 9: Convergence of the Hermitization algorithm (Figure 6).