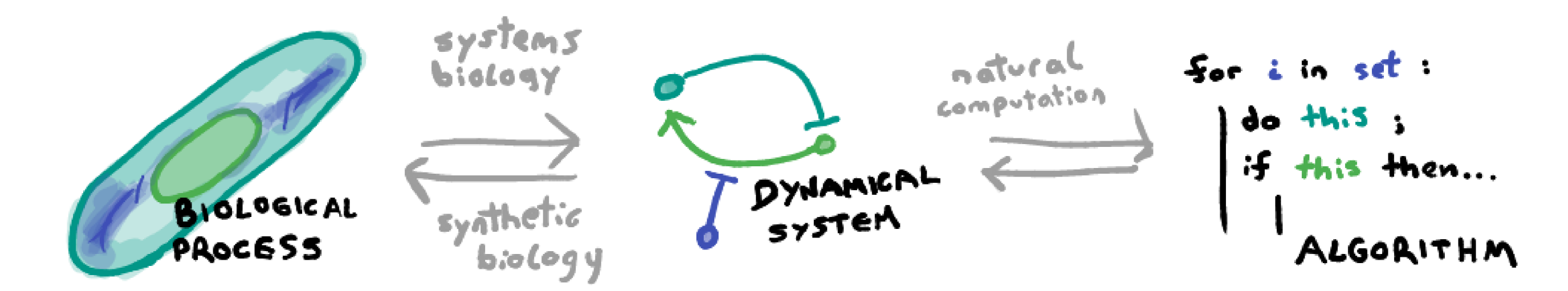


# Reverse-engineering natural computation

using reaction-diffusion approaches beyond linear stability



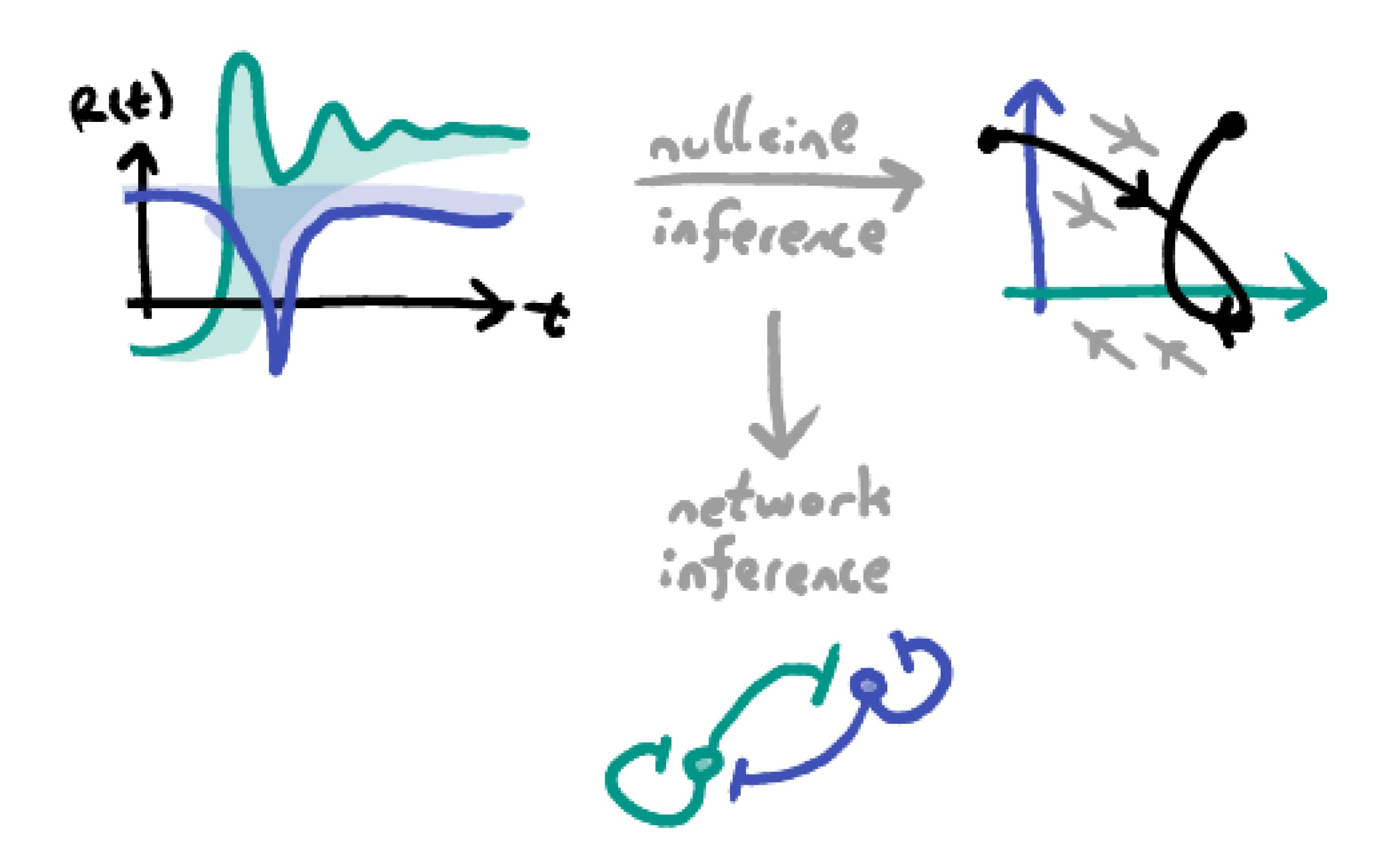
### Gregory Szép, Luca Cardelli, Attila Csikász-Nagy



## Evolution of response behaviour

Specific mappings have been explored between between algorithms, electrical engineering circuits and chemical reaction networks [1]. Understanding function of large biochemical networks from a computer science perspective guides experiments in synthetic biology and in-vitro reconstructive approaches [2].

How does one design the *least complex* chemical reaction network that obeys a given response function?

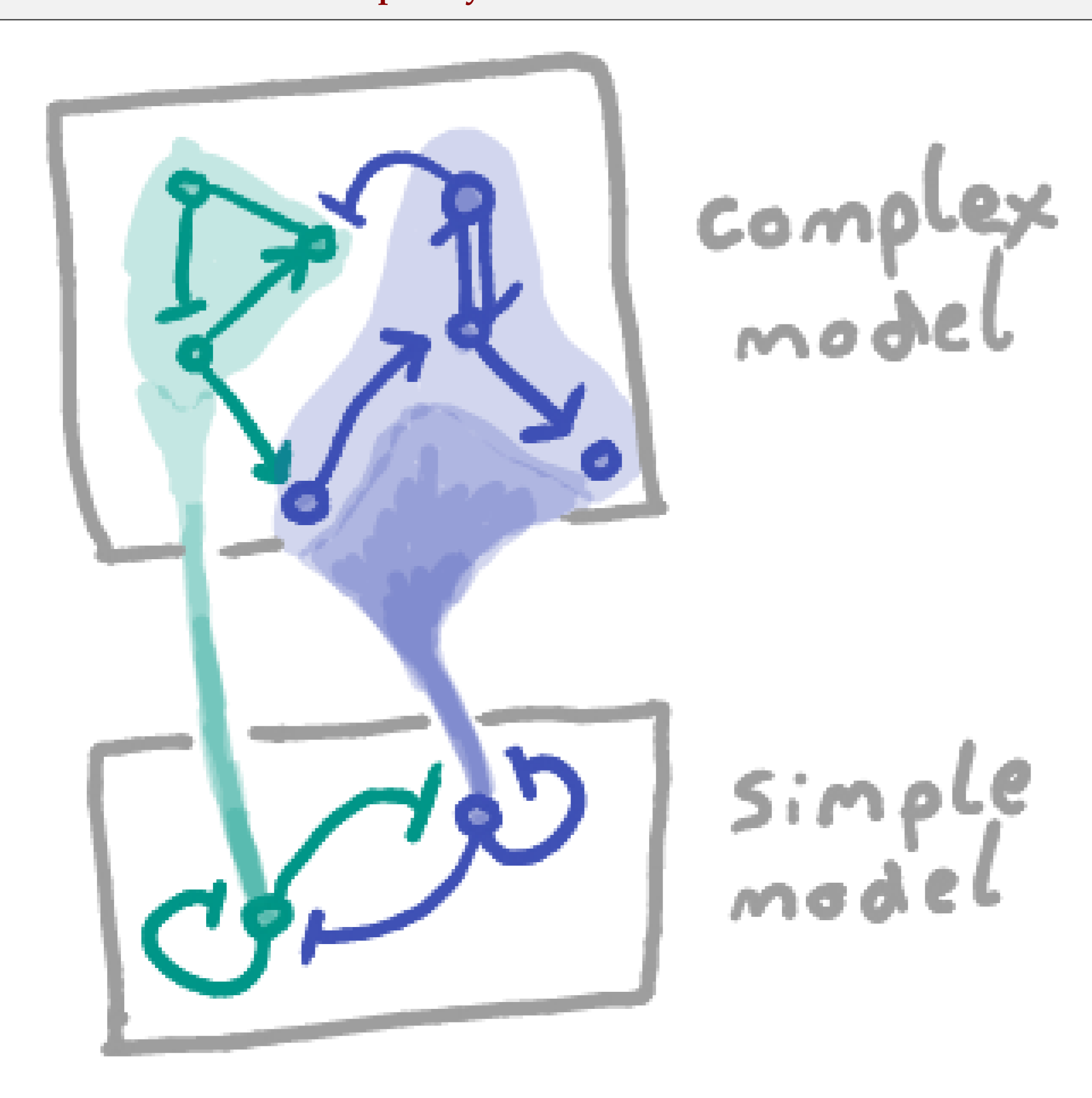


The function of known biochemical networks such as the MAPK pathway, circadian rhythms and cell-cycles can be understood in terms of simple response functions; decomposition of large networks into switches and clocks are proposed in literature.

The networks found in nature are far from the least complicated realisations of particular response functions. The additional complexity can be explained by molecular evolutionary

paths towards robust biological function [3].

Can model reduction methods [4] identify relevant components, parameters and reduce complexity in reaction networks?



Using measures of relative complexity between two given networks, can we construct evolutionary trees and understand how primitive switches and clocks evolved?

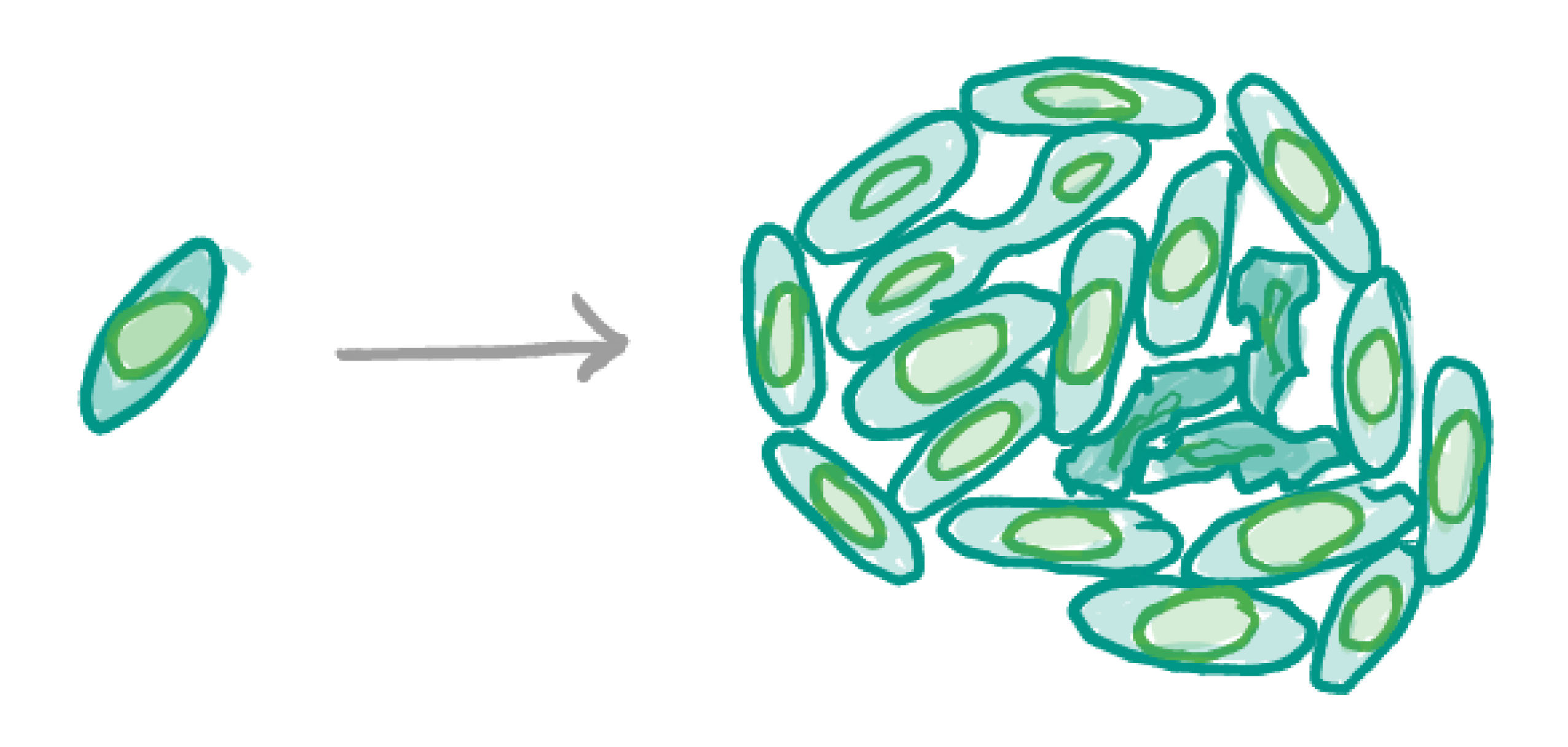
#### 2 Patterns in dynamic populations

Ever since Turing formulated the differential diffusion condition [] for pattern formation, whether a biological pattern is truly driven by a diffusion instability or not has been a matter of debate and speculation.

It is conceivable that the differential diffusion condition is satisfied by the time-scale separation between cytosolic, membrane and inter-cellular reactions.

Finite element reaction-diffusion simulations can take these effects into account explicity at an enourmous computational cost, which leads researches to resort to more abstract Kuramototype models.

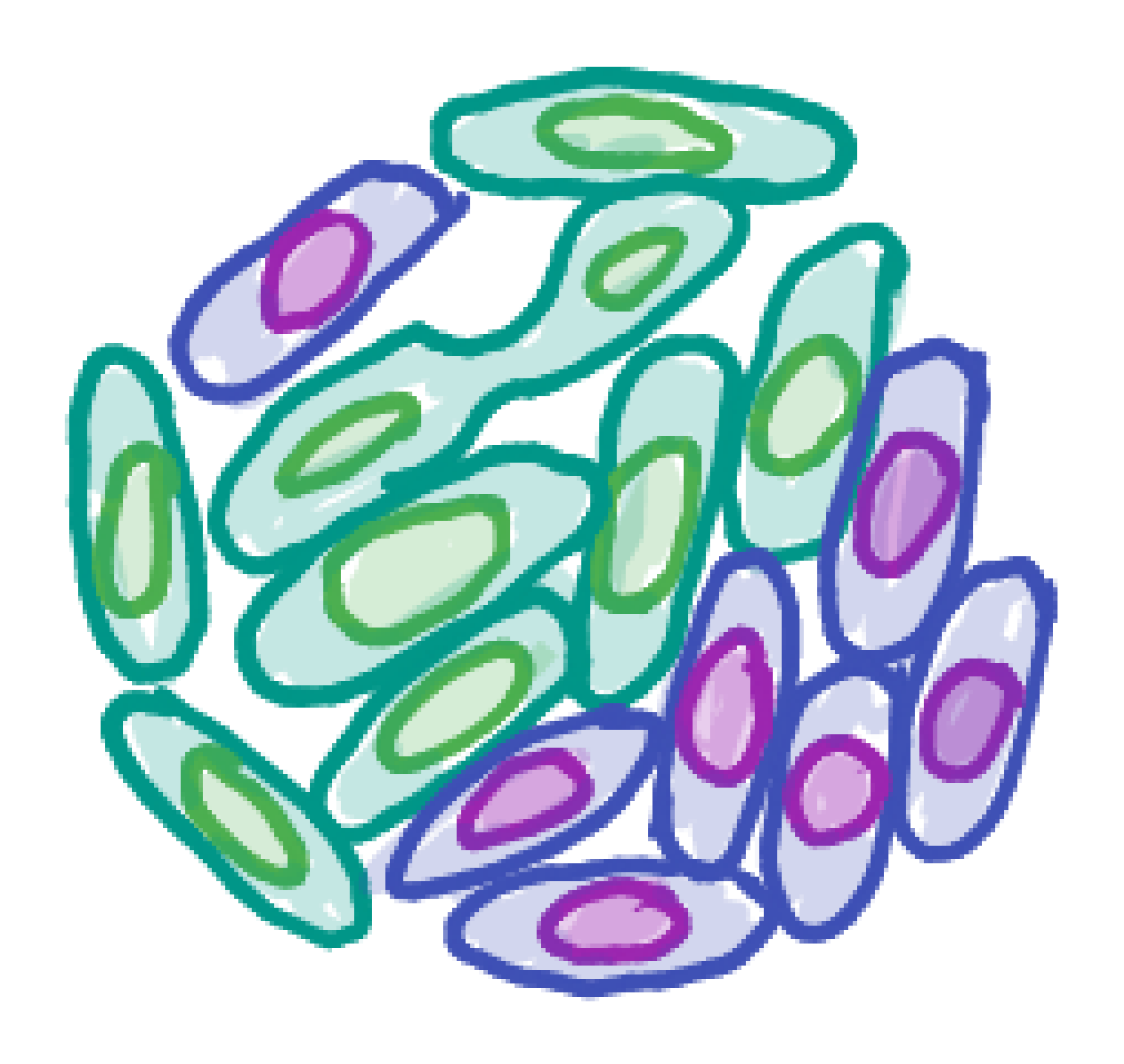
Can we construct a computationally tractable reaction-diffusion model that takes cell division and death into account?



$$\partial_t \rho = \partial_x^2(D\rho) + R(\rho) \qquad \partial_t D = ?$$

Evolution of biochemical chemical networks is not just driven by random mutation, but are subject to environmental pressures and competition. Snythetic approaches to bacterial competition experiments [] allow us to probe the fitness of different chemical networks.

How to we model competition between cell types that contain different reaction networks?



#### 4 Outlook

- Help synthetic biologists
- Help systems biologists
- Help evolutionary biologists
- Do theory; be happy