Evolutionary construction of dynamic biochemical systems performing mathematical calculations

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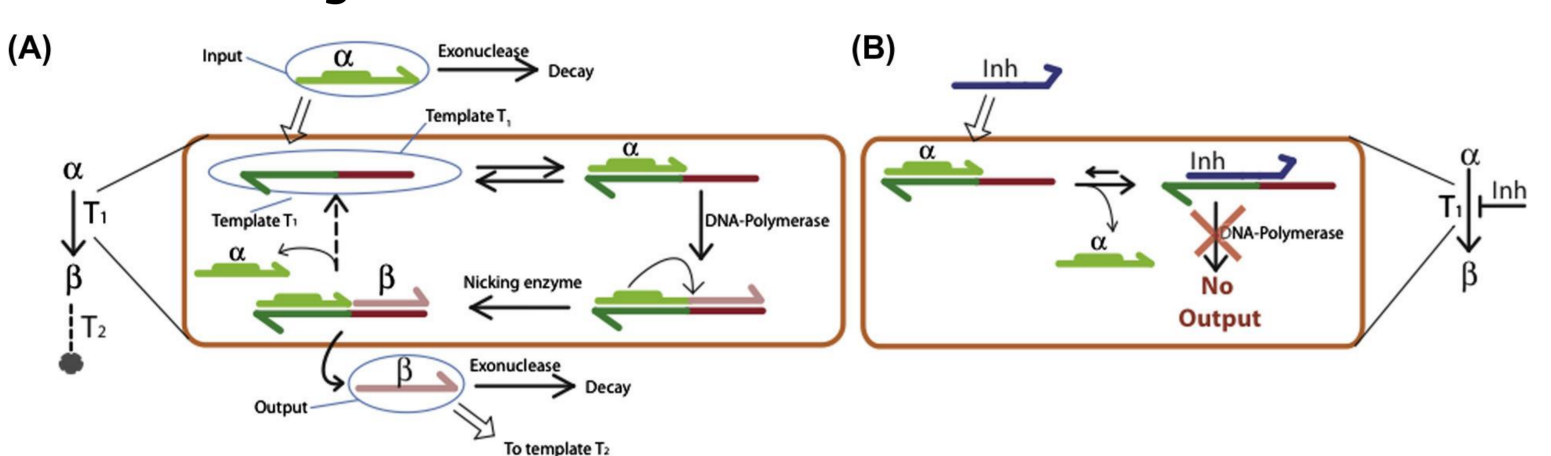


1. Introduction

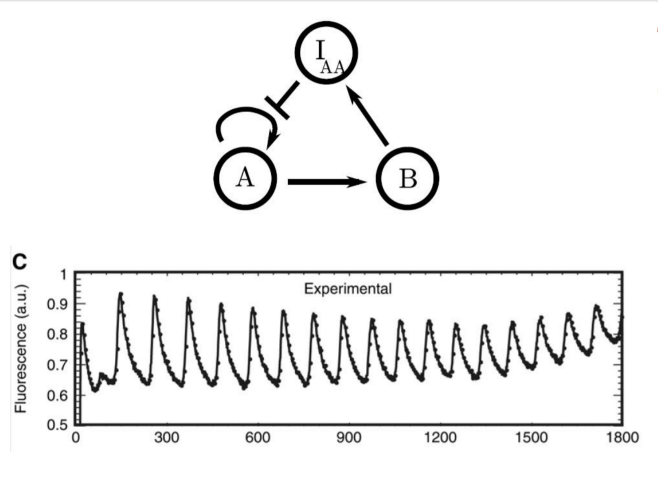
- Goal: use Evolutionary Computations to design analog dynamic biochemical systems, starting from the desired behavior to the actual sequence and concentration of each chemical compound. Apply to cases where there is no clues about the relevant network structure.
- Generally, trial-and-error procedures are applied in parallel with the use of mathematical models \rightarrow not efficient for optimization and implementation of complex systems.
- The DNA Toolbox [1] an experimental molecular programming scheme coupled with quantitative mathematical models could theoretically permit the automation of the full design of biochemical circuits.
- This work uses the combination of an effective technique called ERNe [2] and Differential Evolution to discover feasible and robust solutions to a challenging information processing problem: mathematical calculations which could be used as filters or integrators in molecular circuits.
- This work also tests the possibility of completely automated function-to-test tube design process.

2. Models and Methods

2.1 Building molecular networks: the PEN DNA Toolbox



(A) Activation link between two species (B) Inhibition of activation links



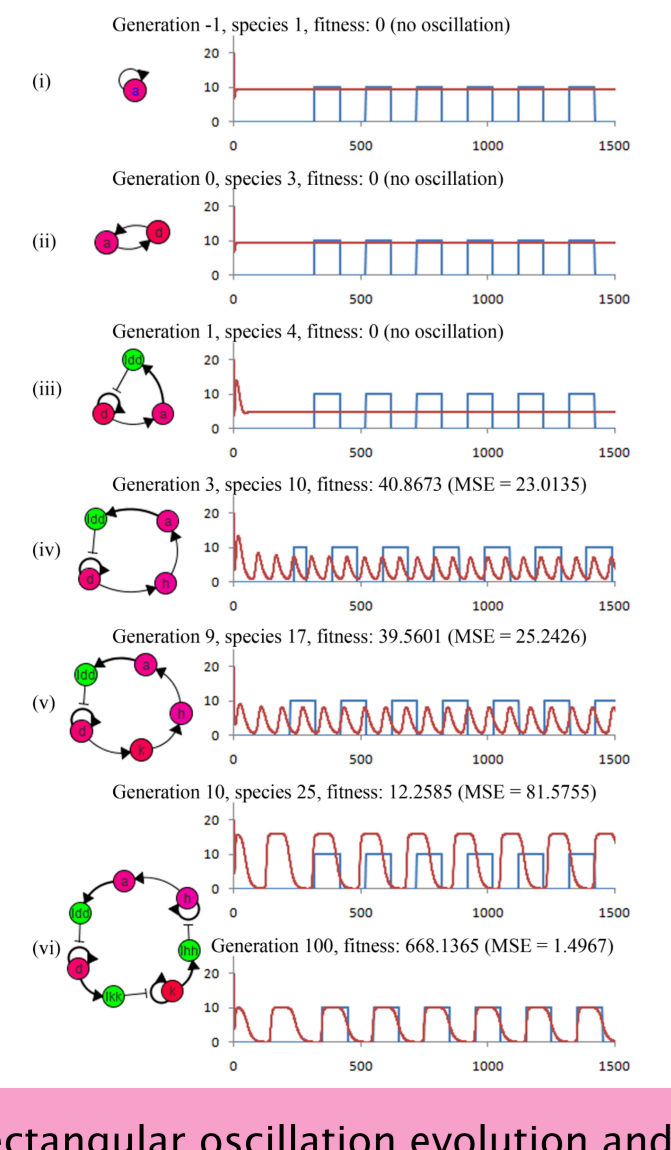
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Oscillator's graphical representation and its equivalent equations [1]

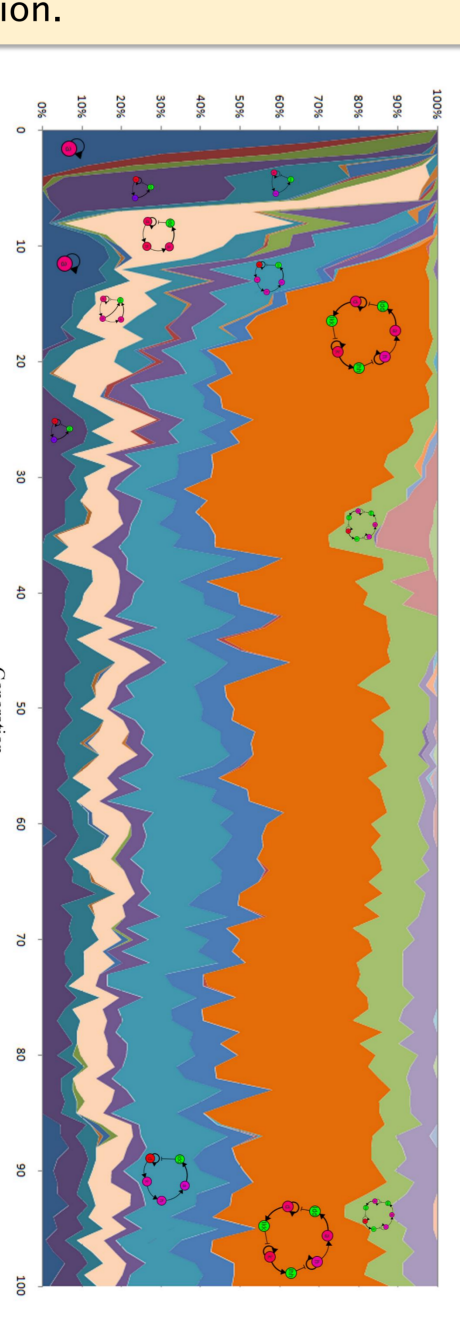
2.2 Searching through topology/parameter space efficiently – ERNe Algorithm

An efficient derivative of the NeuroEvolution of Augmenting Topologies algorithm [3] directed at the evolution of biochemical systems or molecular programs.

- meaningful crossovers between two chemical reaction networks of different topologies.
- preserving topological innovations through speciation.



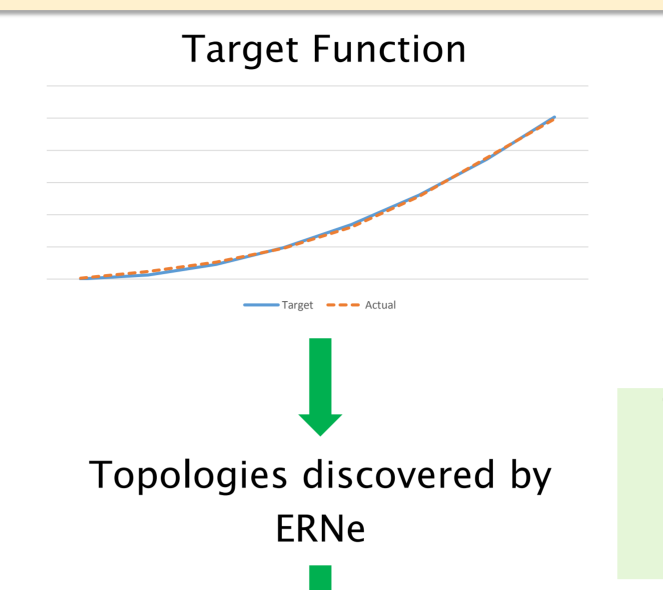
Rectangular oscillation evolution and its species visualization.



2.3 Construction of mathematical calculator

Information Encoding:

- Input: Input sequence's initial concentration.
- Output: Output sequence's concentration at stable state.



- Initial attempt: Square function
- $output ext{ \pi} input^2$ − Input range: from 1µM to 50µM.
- Fitness measured by MSE.

The discovered topologies clearly shows that there must be an autocatalysis inhibiting the activation from input node to output node.

Minimal topologies obtained by manual pruning

by manual pruning

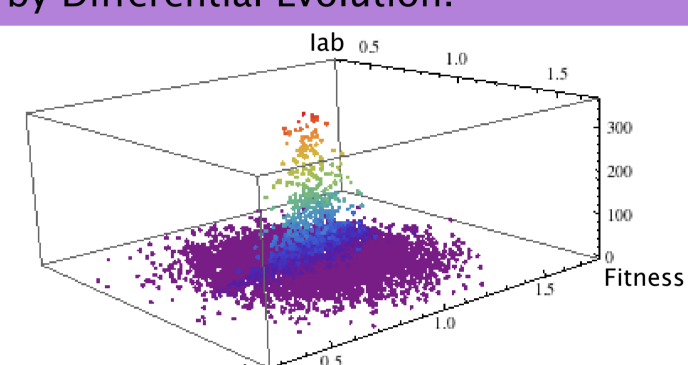
Parameters optimized by Differential Evolution

Robustness Evaluation
5000 tests:
 multiplicative noise is
 given to sequences'
 kinetic parameters.

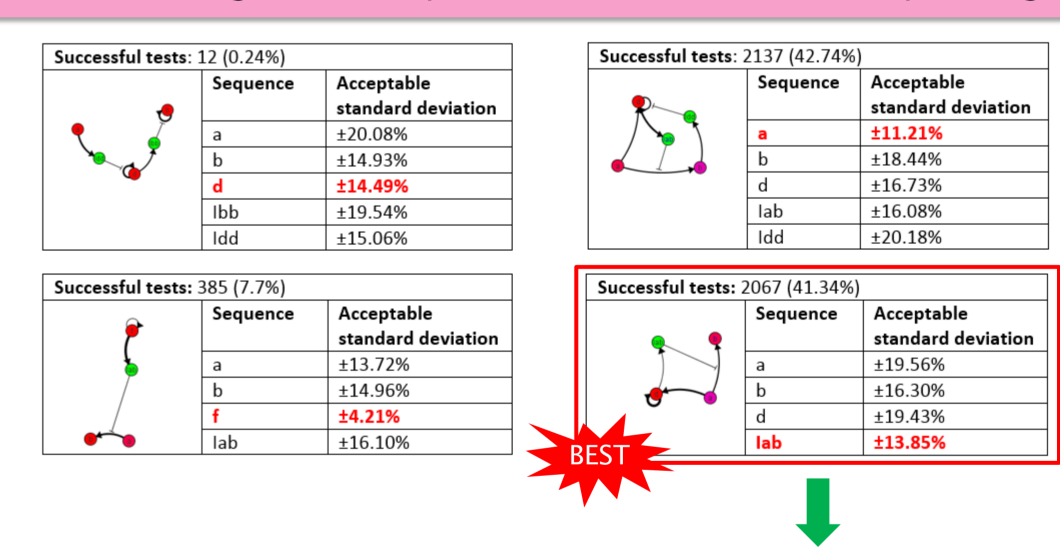
Pruning process:

- Remove irrelevant parts.
- Simplify indirect connections.

Usually, after pruning, fitness reduces significantly. We keep the topologies, and have their parameters optimized by Differential Evolution.



Plot of multiplicative noise given to sequence b and lab, and corresponding fitness

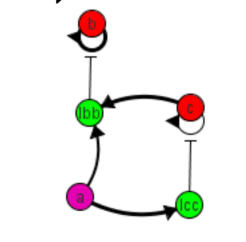


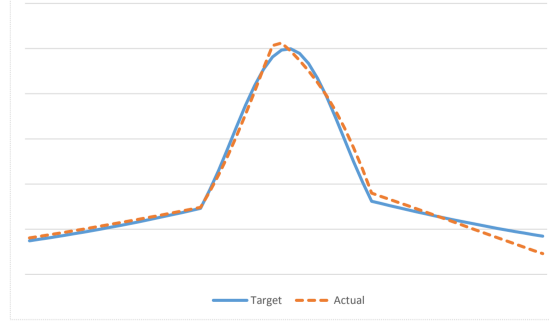
Take the sequences to test tube (on progress)

3. Ongoing works

Gaussian function (log scale)

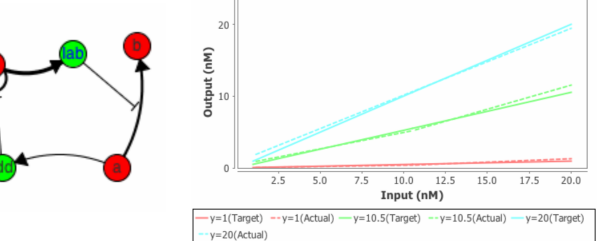
output $\propto e^{-(\log(input)-2)^2/1}$



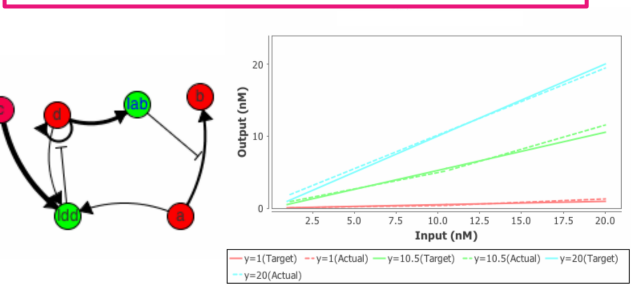


Two-input functions

 $output \propto input1 \times input2$



2.5 20.0 (Target)



 $output \propto input 1/input 2$

References

- [1] Montagne, Kevin, et al. "Programming an in vitro DNA oscillator using a molecular networking strategy." Molecular systems biology, 7(466), 2011.
- [2] Quang Huy, Dinh, *et al.* "An Effective Method for Evolving Reaction Networks in Synthetic Biochemical Systems." IEEE Transactions on Evolutionary Computations. (in press).
- [3] Kenneth O., Stanley and Risto Milkkulainen. "Evolving neural networks through augmenting topologies." Evolutionary Computation, 10(2), 2002.