

Molecular Automata

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Cells store and process information, largely through regulatory networks etc. but overall this is not understood at the level of information processing. Considerable effort is also being put into devising synthetic biological systems which can process information, a field known as molecular computation. Both fields are focused on DNA being the memory which stores information with molecular machines operating on it. However, this is not necessarily the case, we look at the information storage capability of an out-of-equilibrium allosteric complex. Here we build a thermodynamically consistent kinetic model of a molecular complex made of identical subunits each in one of two states (e.g., structural conformations, phosphorylation). We then analyse the dynamics of the complex as we allow each subunit's state to be changed by driven enzymes which act conditionally based on the states of the subunit's neighbours. We identify a set of simplest such systems and identify a one-to-one mapping between those and elementary cellular automata rules.

Want to get accross: we look at the simplest/minimal systems of which there are a discrete count of, we find a one-to-one mapping to ECA rule (which we use to identify symmetries)

We identify a rich set of behaviours including multi-stability and dynamical steady states in various geometries. We further draw an analogy of each of the 256 possible minimal rules to a corresponding cellular automata rule and highlight the difference.

1 New Notes

- describe the model
 - Identical subunits
 - Out-of-equilibrium + kinetic model
 - Rates depend on nearest neighbors
 - Thermodynamically consistent, but maximally driven
- what we find
 - connection to cellular automata (classic model of computation)
 - rich set of behaviours
 - identify analyze attracting components (attractors) which include
 - * multistability and dynamical steady states
 - * individual states (with basins of attraction)
 - * loops (directed or not), single state or branching, of either travelling waves or transitioning between all 0s and all 1s
 - * more complex components, including plane/2D like structures
 - we look at
 - can provide insights on the "error correction" ideas of Crick and others - can have a rule that error corrects but nothing else and can put limits on it from the nearest neighbour limit (no global counting)

2 Old New Notes

2.1 Questions and things to discuss

- Title! Right now it is very clunky. Should it include "thermodynamically consistent"?
- Conclusion bit! Very much not sure about it as it is – is it too short? Should I specifically cite the paper I do there as that really proposes a similar but different model for it, or should I instead highlight the locality difference we add to it? Not sure
- In order to expand the conclusion I need to slightly cut down something.
- Should I reference the figure from the abstract?
- Little Things
 - Should I explicitly list what our futile cycle is? It's a bit long and perhaps obvious but I think it's nice to give a concrete, simple thing to visualize.
 - I should probably check that using "artificially" where I do is appropriate.
 - "this robustness" → "biological robustness"?
 - Is "We consider" acceptable language?

2.2 General

- So going for **Clocks, timers and cell cycle dynamics** topic
- Can I squeeze in "non-Hermitian topology" somewhere (I have some references for it, and it is relevant)?

3 Old Notes

Format For POL2025 (the one due on the 6th) it should be 250 words, include references and up to 1 figure.

Topic I need to choose one of a list of topics. Perhaps the closest might be: **"Biomolecular assemblies and condensates"** given that the main model is of an allosteric assembly? Others that may be relevant:

- "Patterns, waves, transport, collective phenomena, and microswimmers" – there's collective phenomena! but idk about microswimmers
- "Clocks, timers and cell cycle dynamics" – if we lean into KaiABC then maybe
- "Protein structure, dynamics and interactions" – cause I guess the polymer as I've been calling it would realistically be a protein?
- "Emerging Areas in the Physics of Life" – idk what this is but probably not

3.1 Questions

- Approach to talk about a project that has only just started?
- Should I be talking about allostery or not so much? It seems relevant and as an interesting topic but it's not really a core ingredient in it.
- I'm a bit worried that the only "result" is a bit obvious once you think about it. If we add a penalty for NN having different conformation then of course the ones with all the same conformation will be preferred. Then if all are in conformation 1 (tense) then they are ATP driven to bind ligands so they do so. After that they are by our choice of parameters not favoured to debind hence the only thing they can do is change conformation. The same sort of reasoning then completes the cycle.
- Is an ArXiv citation admissible here?

3.2 Key points to cover

- stochastic systems
- futile cycles
- non-equilibrium dynamics
- allostery
- system features
 - complex – high dimensional network, not pen-and-paper tractable
 - locality – unlike the previous models, have NNs, can model waves along the polymer
 - discrete configurational space
 - Thermodynamically consistent? idk if this is the right wording

- Search for topological states/patterns in realistic systems (this means starting from the ground up with (non-equilibrium) statphys, LDB, no arbitrary choices) with futile cycles
- the futile cycle is implemented via physical parameters by coupling to two physically reasonable asymmetrical processes