

Chemistry *and Beyond*

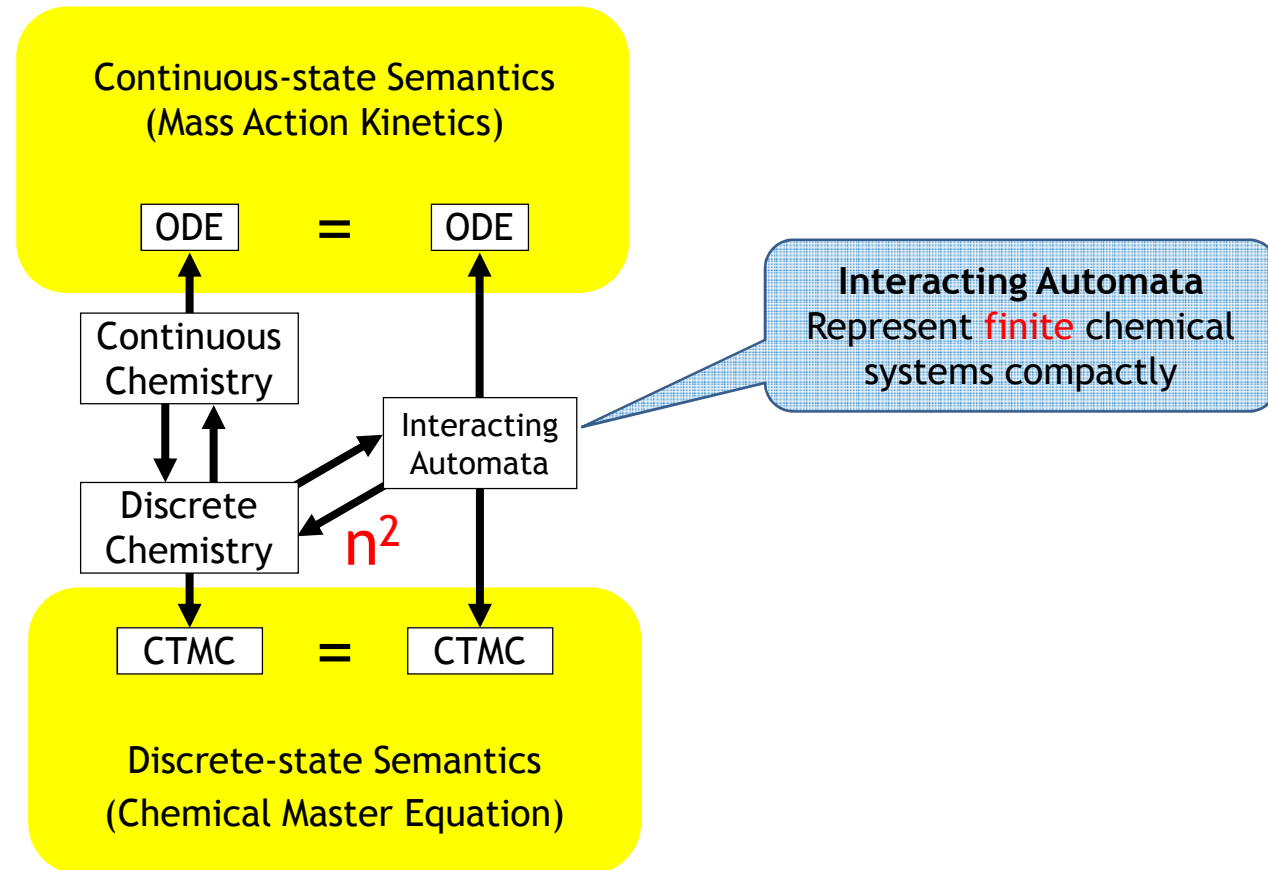
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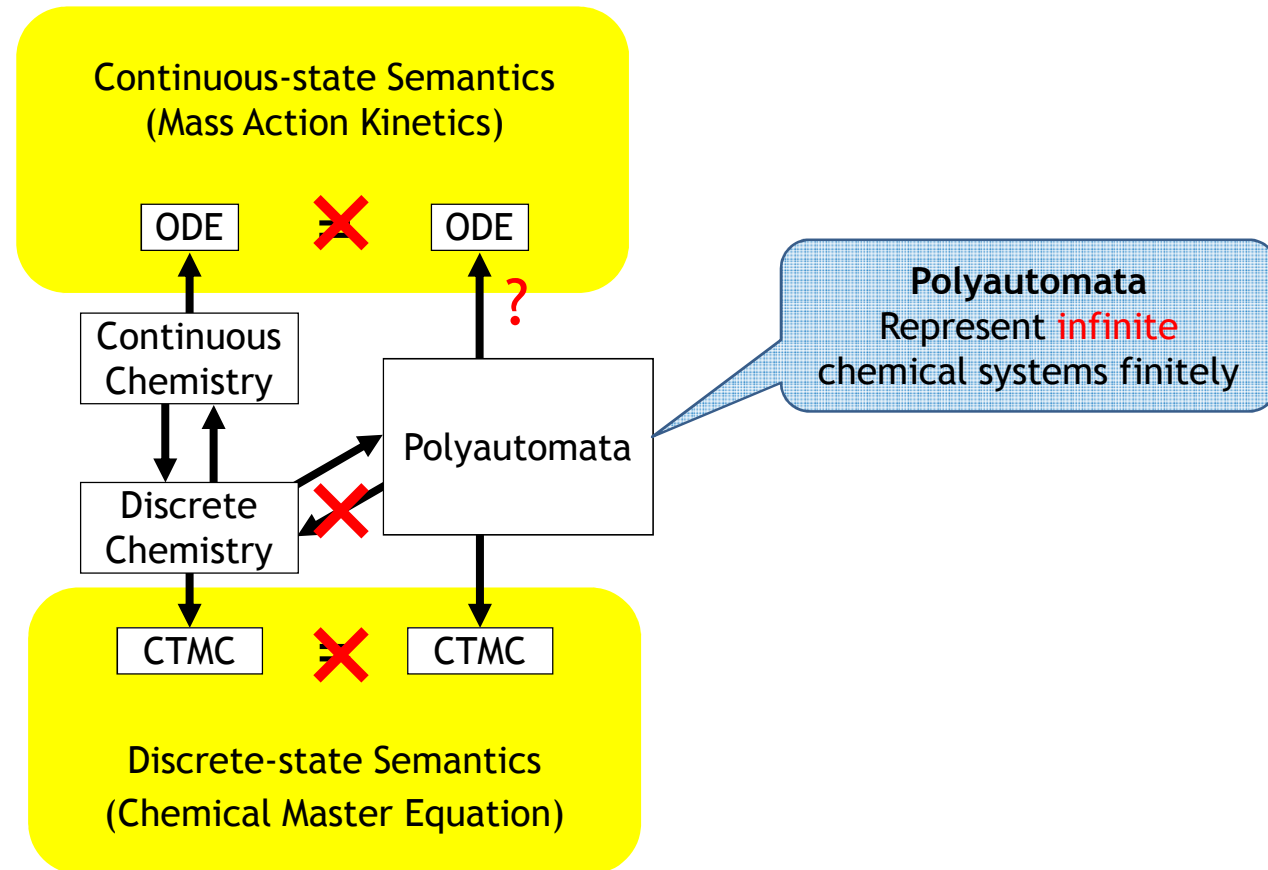
Open Lectures for PhD Students in Computer Science
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<http://lucacardelli.name>

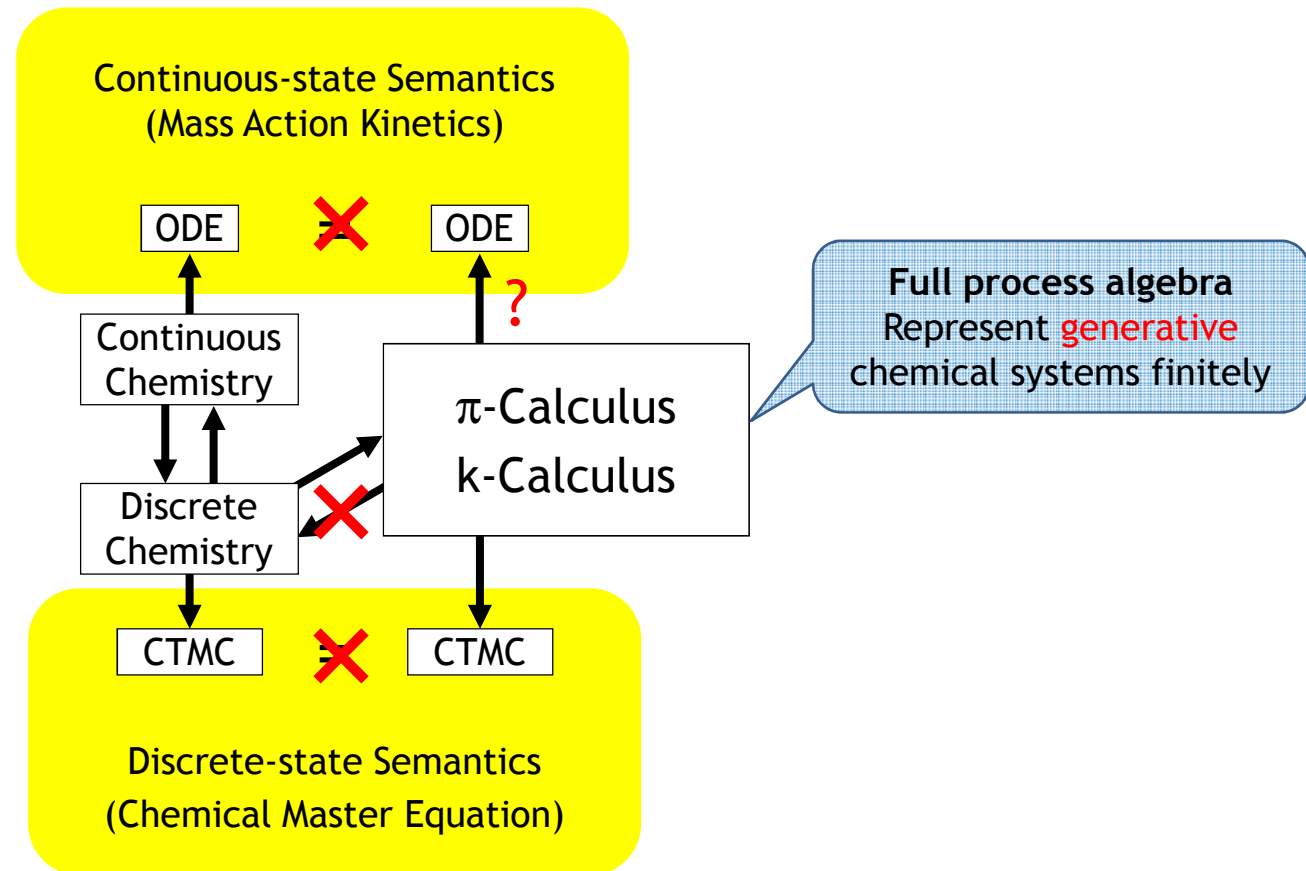
Process Algebra is 'Bigger' than Chemistry



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On the Computational Power of Biochemistry

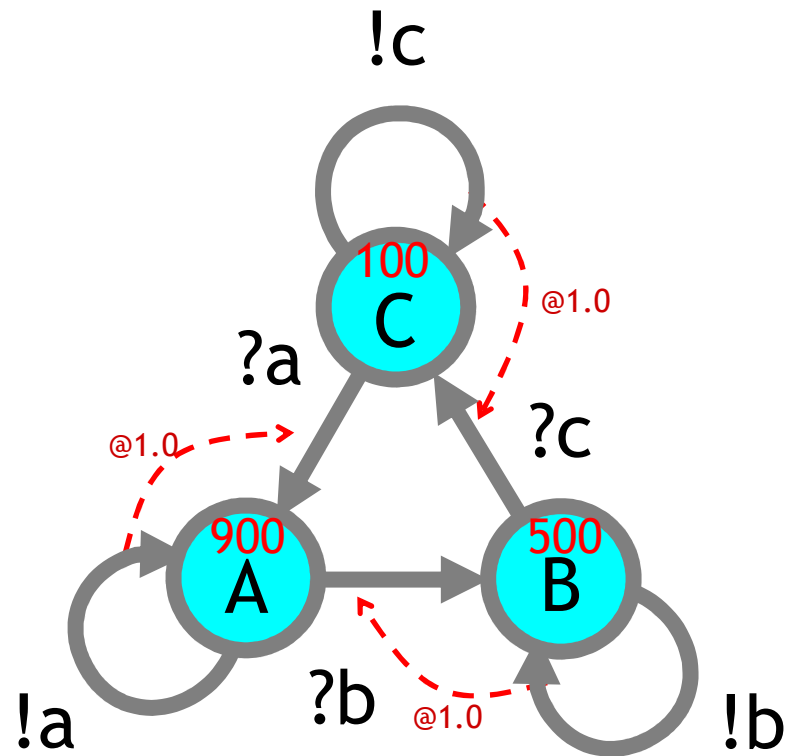
joint work with

Gianluigi Zavattaro

University of Bologna

in: Algebraic Biology '08

Can this program terminate?



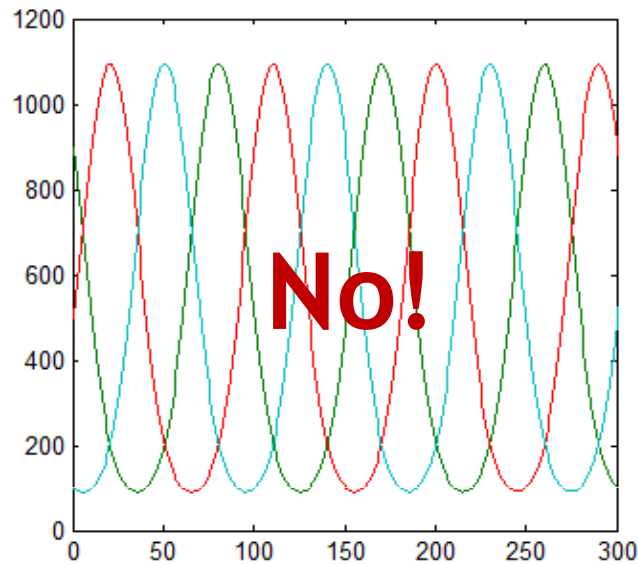
b: $A+B \rightarrow B+B$

c: $B+C \rightarrow C+C$

a: $C+A \rightarrow A+A$

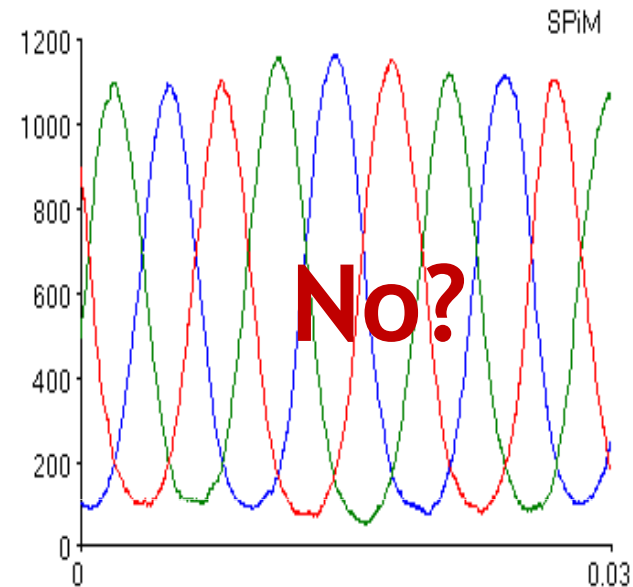
900A + 500B + 100C

“Experimental evidence”



Continuous-State Simulation

```
interval/step [0:0.0001:0.03]
(A) dx1/dt = - x1*x2 + x3*x1 900.0
(B) dx2/dt = - x2*x3 + x1*x2 500.0
(C) dx3/dt = - x3*x1 + x2*x3 100.0
```



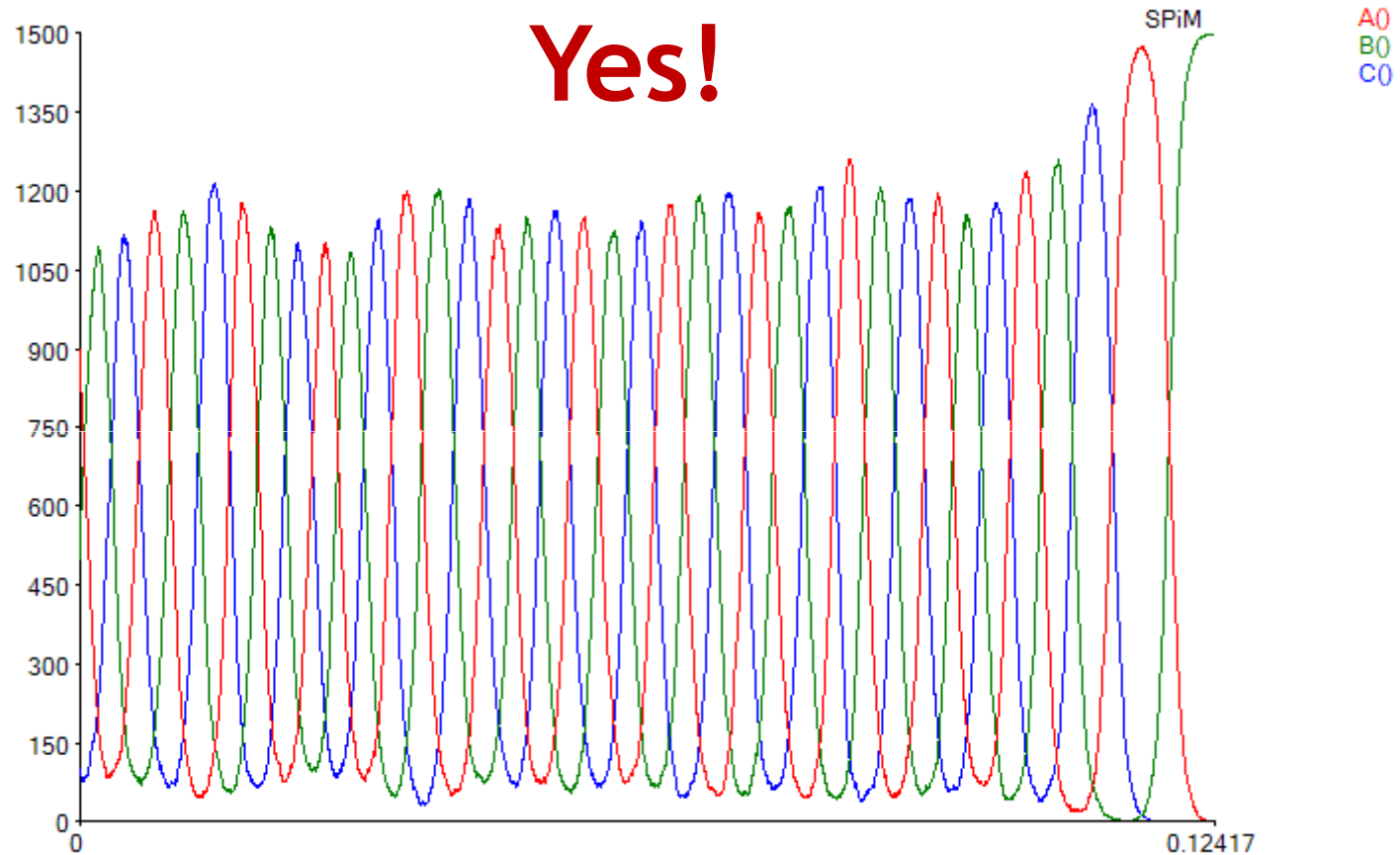
Discrete-State Simulation

```
directive sample 0.03 1000
directive plot A(); B(); C()

new a@1.0:chan new b@1.0:chan new
c@1.0:chan
let A() = do !a;A() or ?b; B()
and B() = do !b;B() or ?c; C()
and C() = do !c;C() or ?a; A()

run (900 of A() | 500 of B() | 100 of C())
```

But in a longer simulation...



Discrete-State
Simulation

$0A + 1500B + 0C$

Is termination (possible death) decidable in Chemistry?

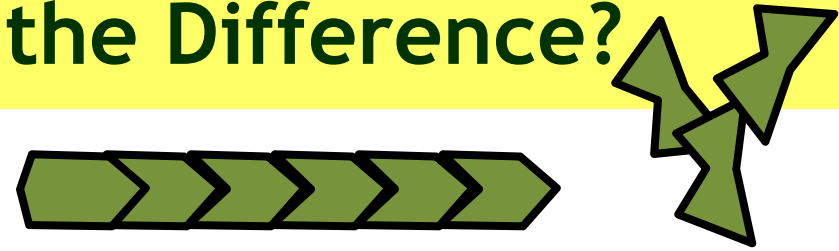
- Termination in Chemistry is at least *subtle*. Is it decidable?
- Three equivalent definitions of “basic chemistry”:
 - FSRN: Finite Stochastic Reaction Networks (finite systems of stochastic chemical reactions)
 - CGF (Interacting Automata): our process algebra.
 - Place-Transition (stochastic) Petri nets.
- Surprising answer: termination in basic chemistry is *decidable!*
 - (Soloveichik et al. *Computation with Finite Stochastic Chemical Reaction Networks*. In Nat. Computing. 2008) by reduction to a decidable problem in Petri Nets (reachability).
- Hence, basic chemistry **cannot compute!**
 - By Turing’s theorem, termination for a universal computer is undecidable.
 - Hence basic chemistry is not Turing-complete.
 - (Although the full story is more subtle and interesting: stochastic chemistry can *approximate* Turing machines to arbitrary precision.)

Can Biochemistry Compute?

- Chemistry cannot compute; is that true of Biochemistry? Not necessarily.
- Although Chemistry (FSRNs) encompasses huge complexity (e.g. chaotic systems), it is in fact unable to express (finitely) virtually *any* biological system of interest!! (and many non-biological ones)
 - So, how have people managed so far? By manipulating awkward infinite collections of chemical reactions or ODEs.
- The language of Biochemistry is intrinsically more powerful than the language of Chemistry: it can represent finitely systems that Chemistry can't. Since it is more powerful it can be Turing complete (and it is).
- What is the language of Biochemistry? Until recently, there wasn't one. Historically the first language used in that sense has been stochastic π -calculus, then (a bit more appropriately) k-calculus.
- The most elementary such language is “polyautomata”.

C.vs. BioC. What's the Difference?

Consider linear polymerization:



The “**chemical program**” for polymerization:



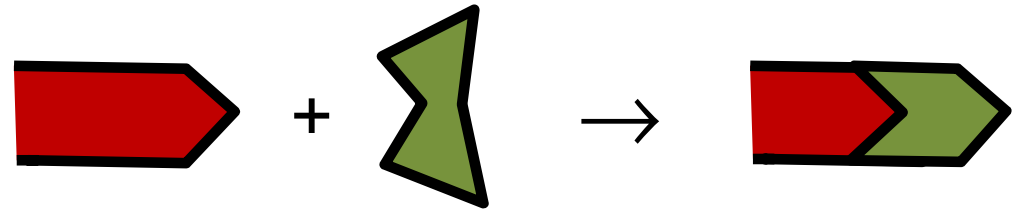
....

- an infinite (non-)program
- an infinite set of species
- an infinite set of ODEs



Such specificity is unreal.

But “**nature's program**” for polymerization **has to fit in the genome**, so it cannot be infinite! Clearly, nature must be using a different “language” than basic chemistry:



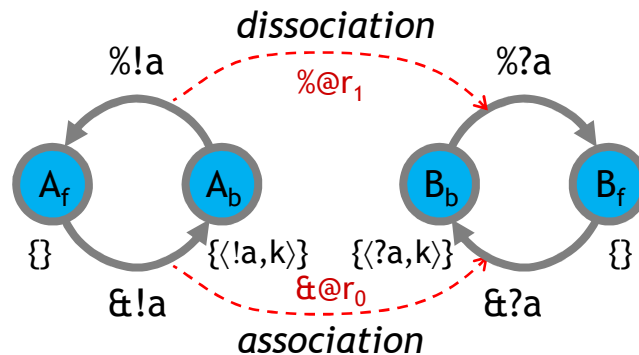
molecule with convex patch + molecule with concave patch → molecule with convex patch

- a finite program
- a local rule

Biochemistry = Collision + Complexation



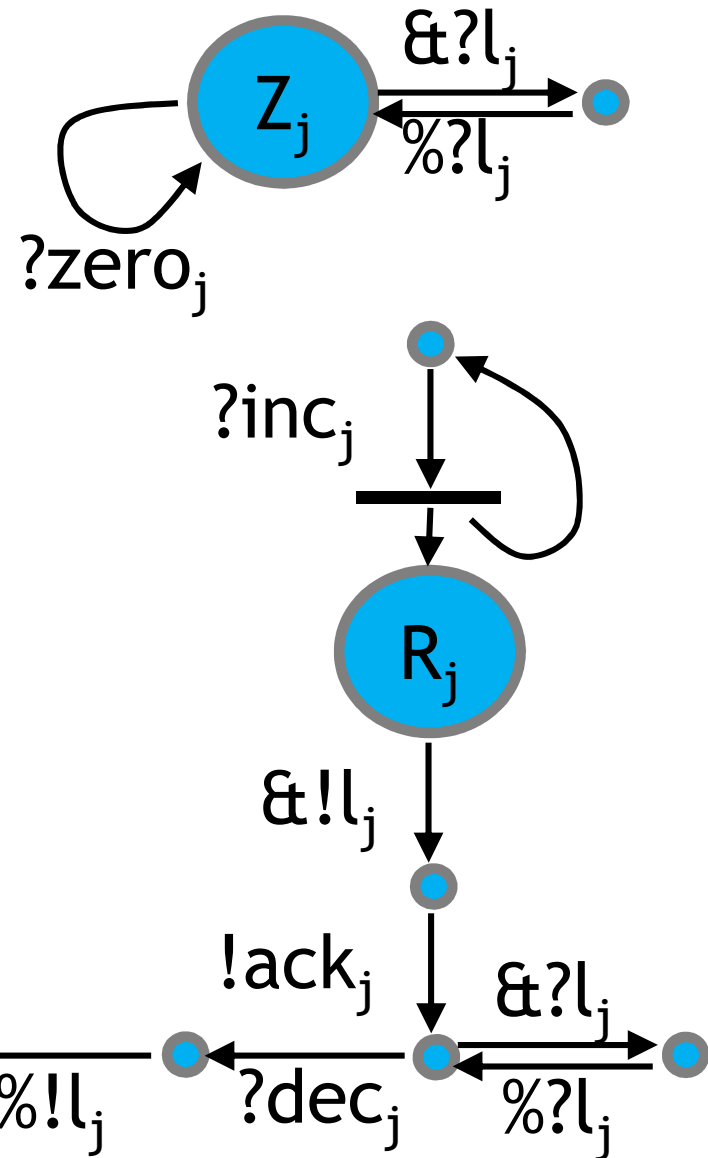
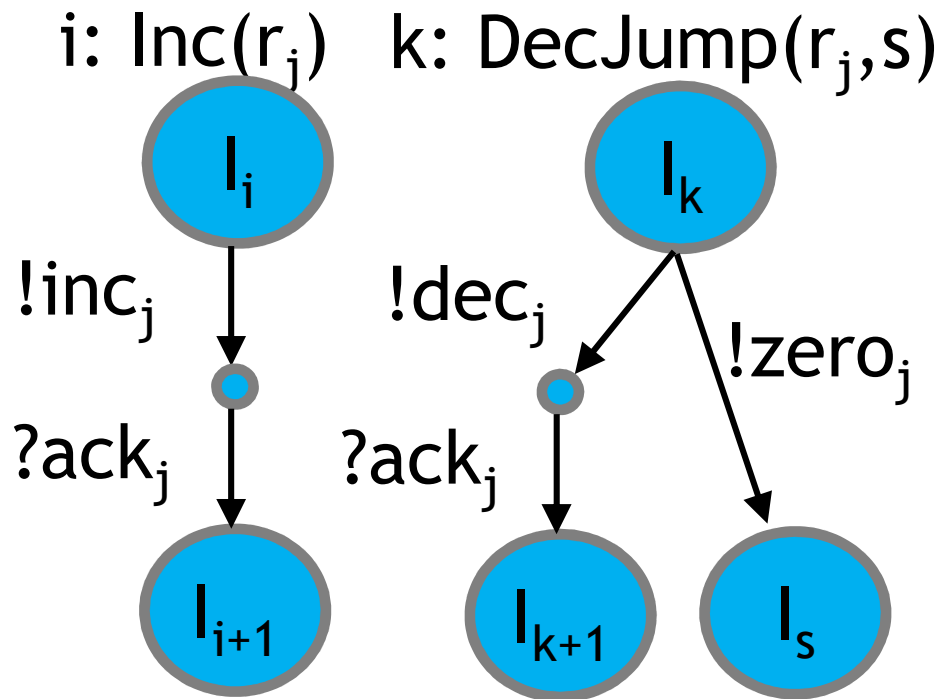
- Complexation is what proteins “do”, in contrast to simpler chemicals.



Polyautomata
(polymerizing automata)

- Leading to a process algebra that we call the **Biochemical Ground Form (BGF)**.

RAM encoding in BGF



Expressiveness of Biochemistry

- Basic chemistry (FSRN, or CGF) **is not** Turing-complete
 - By reduction to Petri Net reachability [Soleveichik&al.].
- Biochemistry (FSRN + complexation, or BGF) **is** Turing-complete.
 - By an encoding of Random Access Machines, using polymers for registers.
- A relatively simple extension of our CGF automata
 - **But it is not as easy to find a corresponding extension of chemistry!**
- More powerful process algebras of course *are* Turing complete
 - They (e.g. π -calculus) include BGF, but they also have mechanisms that are not directly biologically justifiable.
 - In BGF we have in a sense the minimal biologically-inspired extension of FSRN, and it is already Turing-complete.
- **Intrinsic to biochemistry (but not to simple chemistry) is a Turing-complete mechanism.**