Abstract interpretation of protein-protein interactions networks

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Tuesday, the 25th of June, 2019
Joint-work with...

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Tuesday, the 25th of June, 2019
Signalling Pathways

EGF, TGF-alpha, etc

EGFR

PI3-K

AKT

phosphorylation

mTOR

STAT

GRB2

SOS

RAS

RAF

ERK

MEK

Gene transcription
Cell cycle progression

Cell proliferation
Inhibition of apoptosis
Angiogenesis
Migration, Adhesion, Invasion

Eikuch, 2007
Bridging the gap between...
Site-graphs rewriting

- a language close to knowledge representation;
- rules are easy to update;
- a compact description of models.
Choices of semantics

interaction map

Markov chain

ordinary differential equations

\[
\begin{align*}
\frac{dx_1}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_2}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_3}{dt} &= k_1 \cdot x_1 \cdot x_2 - k_{-1} \cdot x_3 + 2 \cdot k_2 \cdot x_3 \cdot x_3 - k_{-2} \cdot x_4 \\
\frac{dx_4}{dt} &= k_2 \cdot x_3^2 - k_2 \cdot x_4 + \frac{x_4 x_5}{p_4 + x_5} - k_3 \cdot x_4 - k_{-3} \cdot x_5 \\
\frac{dx_5}{dt} &= \cdots \\
\frac{dx_n}{dt} &= -k_1 \cdot x_1 \cdot c_2 + k_{-1} \cdot x_3
\end{align*}
\]
Complexity walls

- **Combinatorial wall**
- **Event wall**

- **Deterministic differential equations**
- **Stochastic master equations**
- **Agent/rule-based**

- **Number of instances per molecular species**
- **Number of molecular species**

- **Early EGF**
- **EGF to ERK**
- **EGF to ERK and AKT**
- **EGF with receptor network, to ERK and AKT**

- **$10^6$**
- **1000**
- **100**

- **Tuesday, the 25th of June, 2019**
Abstractions offer different perspectives on models

concrete semantics

causal traces

information flow

exact projection of the ODE semantics
Static analysis of reachable species (I/II)

We capture the relationships between the states of the sites of each agent.
Static analysis of reachable species (I/II)

We capture the relationships between the states of the sites of each agent.
Static analysis of reachable species (II/II)

Applications:

1. check the consistency of a model [ICCMSE’07]
2. compute the properties to allow fast simulation [APLAS’07]
3. simplify models,
4. compute independent fragments of chemical species [PNAS’09, LICS’10, Chaos’10]

The analysis is complete (no false positif) for a significatif kernel of Kappa [VMCAI’08].
Model reduction

The ground differential system uses one variable per chemical species; we directly compute its exact projection over independent fragments of chemical species. With a small model, 356 chemical species are reduced into 38 fragments:

On a bigger model, $10^{19}$ chemical species are reduced into 180 000 fragments. [PNAS’09, LICS’10, Chaos’10]
Reachability Analysis of Rule-based Models

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Tuesday, the 25th of June, 2019
In this talk...

We illustrate the following concepts:

- **Galois connections:**
  - the upper closure operator $\gamma \circ \alpha$,
  - the lower closure operator $\alpha \circ \gamma$;

- **soundness:**
  - the abstraction forgets no behavior;

- **completeness:**
  - sufficient conditions that ensure the absence of false positive;

on an abstraction of the reachable connected components in a site-graph rewriting language.
Joint-work with...

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Jean Krivine  
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Overview

1. Introduction
2. Language: Kappa
3. Abstraction: Local views
4. Completeness: false positives?
5. Local fragment of Kappa
6. Decontextualization
7. Conclusion
Signaling Pathways

- EGF, TGF-alpha, etc
- EGFR
- PI3-K
- AKT
- phosphorylation
- mTOR
- STAT
- GRB2
- SOS
- RAS
- RAF
- MEK
- ERK
- Gene transcription
  - Cell cycle progression
  - Cell proliferation
  - Inhibition of apoptosis
  - Angiogenesis
  - Migration, Adhesion, Invasion

Eikuch, 2007
Causal traces
ODE semantics

EGF pathway (reduced ODEs)

Concentration

Time
What will happen if more Shc(s) is put in the system?
ODE semantics
Crowding effect

EGF pathway (reduced ODEs)

EGF pathway (reduced ODEs / with 10 times more of Shc(s))
Overview

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A chemical species

\[ \text{EGF(r!1), EGFR(l!1,r!2), EGFR(r!2,l!3), EGF(r!3)} \]
A Unbinding/Binding Rule

\[
\text{EGF(r), EGFR(l,r)} \leftrightarrow \text{EGF(r!1), EGFR(l!1,r)}
\]
Internal state

\[ \text{EGFR(Y48} \sim u? , l!1), \text{ EGF(r}!1) \leftrightarrow \text{EGFR(Y48} \sim p? , l!1), \text{ EGF(r}!1) \]
Don’t care, Don’t write
A contextual rule

\[ \text{EGFR}(Y48 \sim u, r! \_ ) \rightarrow \text{EGFR}(Y48 \sim p, r) \]
Creation/Suppression

\[ R(r) \rightarrow R(r!1), \, R(r!1, l, Y48 \sim u) \]

\[ R(r!1), \, R(r!1) \rightarrow R(r) \]
Early EGF example

egf rules 1

protein shorthands: E:=egf, R:=egfr, So:=Sos,Sh:=Sh,G:=grb2

* Ligand-receptor binding, receptor dimerisation, rtk x-phosph, & de-phosph
  01: R(l,r), E(r) <-> R(l¹,r), E(r¹)
  02: R(l¹,r), R(l²,r) <-> R(l¹,r³), R(l²,r³)
  03: R(r¹,Y68) -> R(r¹,Y68p)
    R(Y68p) -> R(Y68)
  04: R(r¹,Y48) -> R(r¹,Y48p)
    R(Y48p) -> R(Y48)

* Sh x-phosph & de-phosph
  14: R(r²,Y48p¹), Sh(π¹,Y7) -> R(r²,Y48p¹), Sh(π¹,Y7p)
  ??: Sh(π¹,Y7p) -> Sh(π¹,Y7)
  16: Sh(π,Y7p) -> Sh(π,Y7)
  refined from
  Sh(Y7p) -> Sh(Y7)

* Y68-G binding
  09: R(Y68p), G(a,b) <-> R(Y68p¹)+G(a¹,b)
  11: R(Y68p), G(a,b²) <-> R(Y68p¹)+G(a¹,b²)
  refined from
  R(Y68p¹)+G(a) <-> R(Y68p¹)+G(a¹)
Early EGF example

egf rules 2

refined from
So(d)+G(b)<->So(d)+G(b1)

interface note: highlight
the interacting parts

G-So binding
10: R(Y68, G(a1,b), So(d) <-> R(Y68, G(a1,b2), So(d2))
12: G(a,b), So(d)   <->  G(a,b1), So(d1)
22: Sh(π,Y7, G(a2,b), So(d) <-> Sh(π,Y7, G(a2,b1), S(d1))
19: Sh(π,Y7, G(a2,b), So(d)   <->  Sh(π,Y7, G(a2,b1), S(d1))

Y48-Sh binding
13: R(Y48, Sh(π,Y7) <-> R(Y48, Sh(π,Y7))
15: R(Y48, Sh(π,Y7) <-> R(Y48, Sh(π,Y7))
18: R(Y48, Sh(π,Y7), G(a1,b) <-> R(Y48, Sh(π,Y7), G(a1,b))
20: R(Y48, Sh(π,Y7), G(a1,b3), S(d3) <-> R(Y48, Sh(π,Y7), G(a1,b3), S(d3))

Sh-G binding
17: R(Y48, Sh(π,Y7), G(a,b) <-> R(Y48, Sh(π,Y7) G(a), G(b))
21: Sh(π,Y7), G(a,b) <-> Sh(π,Y7), G(a1,b)
23: Sh(π,Y7), G(a,b2) <-> Sh(π,Y7), G(a1,b2)
24: R(Y48, Sh(π,Y7), G(a,b3), S(d3) <-> R(Y48, Sh(π,Y7), G(a2, b3), S(d3))

refined from
Sh(π), G(a)<->Sh(π), G(a1)

refined from
R(Y48)+Sh(π)<->R(Y48)+Sh(π1)

why not simply G(b3)??

refined from
So(d)+G(b)<->So(d1)+G(b1)

interface note: highlight
the interacting parts
Properties of interest

1. Show the absence of modeling errors:
   - detect dead rules;
   - detect overlapping rules;
   - detect non exhaustive interactions;
   - detect rules with ambiguous molecularity.

2. Get idiomatic description of the networks:
   - capture causality;
   - capture potential interactions;
   - capture relationships between site states;
   - simplify rules.

3. Allow fast simulation:
   - capture accurate approximation of the wake-up relation.
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Concrete semantics

A rule is a symbolic representation of a multi-set of reactions.

For instance, the rule:

\[
\begin{array}{c}
/C8/CB/CU/D6/CP/CV \\
/D6/CT/D4/D0/CP \\
/CT/D1/CT/D2 \\
/D8/D7
\end{array}
\]

\[k_d\]

within a model with the following signature:

\[
\begin{array}{c}
/C8/CB/CU/D6/CP/CV \\
/D6/CT/D4/D0/CP \\
/CT/D1/CT/D2 \\
/D8/D7
\end{array}
\]

\[k_d\]

\[
\begin{array}{c}
/C8/CB/CU/D6/CP/CV \\
/D6/CT/D4/D0/CP \\
/CT/D1/CT/D2 \\
/D8/D7
\end{array}
\]

\[k_d\]

\[
\begin{array}{c}
/C8/CB/CU/D6/CP/CV \\
/D6/CT/D4/D0/CP \\
/CT/D1/CT/D2 \\
/D8/D7
\end{array}
\]

\[k_d\]

denotes the following two reactions:
Set of reachable chemical species

Let $\mathcal{R} = \{R_i\}$ be a set of rules.
Let $\textit{Species}$ be the set of all chemical species $(C, c_1, c'_1, \ldots, c_k, c'_k, \ldots \in \textit{Species})$.
Let $\textit{Species}_0$ be the set of initial .

We are interested in $\textit{Species}_\omega$, the set of all chemical species that can be constructed in one or several applications of the reactions induced by the rules in $\mathcal{R}$, starting from the set $\textit{Species}_0$ of initial chemical species.

(We do not care about the number of occurrences of each chemical species).
Inductive definition

We define the mapping $F$ as follows:

$$F : \begin{cases} \varphi(\text{Species}) & \rightarrow \varphi(\text{Species}) \\ X & \mapsto X \cup \left\{ c'_j \left| \exists R_k \in \mathcal{R}, c_1, \ldots, c_m \in X, c_1, \ldots, c_m \xrightarrow{R_k} c'_1, \ldots, c'_n \right. \right\} \right. \end{cases}$$

The set $\varphi(\text{Species})$ is a complete lattice.

The mapping $F$ is an extensive $\cup$-complete morphism.

We define the set of reachable chemical species as follows:

$$\text{Species}_\omega = \bigcup \{ F^n(\text{Species}_0) \mid n \in \mathbb{N} \}.$$
Local views

\[ \alpha(\{R(Y1 \sim u,l!1), E(r!1)\}) = \{R(Y1 \sim u,l!r.E); E(r!l.R)\}. \]
Galois connection

Let \( \text{Local\_view} \) be the set of all local views.

Let \( \alpha \in \wp(\text{Species}) \rightarrow \wp(\text{Local\_view}) \) be the function that maps any set of chemical species into the set of their local views.

The set \( \wp(\text{Local\_view}) \) is a complete lattice. The function \( \alpha \) is a \( \bigcup \)-complete morphism.

Thus, it defines a Galois connection:

\[
\varphi(\text{Species}) \xleftarrow{\gamma} \varphi(\text{Local\_view}).
\]

(The function \( \gamma \) maps a set of local views into the set of complexes that can be built with these local views).
\( \gamma \circ \alpha \) is an upper closure operator: it abstracts away some information.

Guess the image of the following set of chemical species?

\[
\{ R, \ l, \ r \} 
\]
\( \alpha \circ \gamma \)

\( \alpha \circ \gamma \) is a lower closure operator: it simplifies (or reduces) constraints.

Guess the image of the following set of local views?

\[
\begin{cases}
R & \text{and} & S
\end{cases}
\]
One more question

$\alpha \circ \gamma$ is a lower closure operator: it simplifies (or reduces) constraints.

Guess the image of the following set of local views ?

\[
\begin{cases}
\{ \begin{array}{c}
R \\
R.l \\
R.r \\
R.l
\end{array} \} & , \\
\{ \begin{array}{c}
R \\
R.r \\
R.l \\
R.l
\end{array} \}
\end{cases}
\]
Abstract reactions

EGFR \rightarrow EGFR

EGFR_{Y48} \rightarrow EGFR_{Y48}
Abstract counterpart to $\mathcal{F}$

We define $\mathcal{F}^\#$ as:

$$\mathcal{F}^\#: \begin{cases} 
\wp(\text{Local\_view}) & \rightarrow \wp(\text{Local\_view}) \\
\forall \mathcal{Y} \rightarrow \mathcal{Y} \cup \{ \mathcal{I}_j' \mid \exists R_k \in \mathcal{R}, \mathcal{I}_{v_1}, \ldots, \mathcal{I}_{v_m} \in \mathcal{Y}, \mathcal{I}_{v_1}, \ldots, \mathcal{I}_{v_m} \rightarrow_{R_k}^\# \mathcal{I}_{v_1}', \ldots, \mathcal{I}_{v_n}' \} 
\end{cases}.$$

We have:

- $\mathcal{F}^\#$ is extensive;
- $\mathcal{F}^\#$ is monotonic;
- $\mathcal{F} \circ \gamma \subseteq \gamma \circ \mathcal{F}^\#$;
- $\mathcal{F}^\# \circ \alpha = \alpha \circ \mathcal{F} \circ \gamma \circ \alpha$ (we will see later why).
Soundness

Theorem 1  Let:

1. \((D, \subseteq, \cup)\) and \((D^\#, \subseteq, \cup)\) be chain-complete partial orders;
2. \(D \xleftarrow{\gamma} \xrightarrow{\alpha} D^\#\) be a Galois connection;
3. \(F \in D \to D\) and \(F^\# \in D^\# \to D^\#\) be monotonic mappings such that:
   \[F \circ \gamma \subseteq \gamma \circ F^\#;\]
4. \(X_0 \in D\) be an element such that: \(X_0 \subseteq F(X_0)\);

Then:

1. both \(lfp_{X_0} F\) and \(lfp_{\alpha(X_0)} F^\#\) exist,
2. \(lfp_{X_0} F \subseteq \gamma(lfp_{\alpha(X_0)} F^\#).\)
Overview

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For any $X \in \wp(\text{Local\_view})$, $\gamma(X)$ is given by a rewrite system:

For any $lv \in X$, we add the following rules:

$I$ and semi-links are non-terminal.
$I$ is the initial symbol.
Pumping lemma

• We use this rewrite system to enumerate the chemical species of $\gamma(X)$.
• There are two cases:
  1. either there is a finite number of rewrite sequences;
  2. or we encounter cyclic derivations
     i.e. an open chemical species with a cycle of the following form:
     $$R.l-r.E \ldots R.l-r.E$$
     can be built.
• We only enumerate chemical species that are reached through an acyclic
  rewriting computation.
• It turns out that: if $X \in \alpha(\varrho(Species))$ then each rewrite sequence is the
  prefix of a terminating rewrite sequence.
  (So there is an unbounded number of species if, and only if,
  there is an unbounded number of rewrite sequences.)
Examples

1. Make the demo for egf
2. Make the demo for fgf
3. Make the demo for Global invariants
Counting chemical species

Given a set of local views $X$, we can easily count the number of species in $\gamma(X)$ by using the following lemmas:

**Lemma 1 (rigidity)** An embedding between two connected components is fully characterized by the image of one agent.

**Lemma 2 (automorphism)** If $\gamma(X)$ is finite, then for any $C \in \gamma(X)$:
- $C$ has at most two automorphisms;
- if $C$ has two automorphisms, then $C$ has a bond of the form $R.r - r.R$. Moreover one automorphism swaps the two $R$ of this bond.

**Lemma 3 (Euler)** If a chemical species has no cycle, then it has an agent with only one site.

sketch the algorithm
Which information is abstracted away?

Our analysis is exact (no false positive):

- for EGF cascade (356 chemical species);
- for FGF cascade (79080 chemical species);
- for SBF cascade (around $10^{19}$ chemical species).

We know how to build systems with false positives... 
...but they seem to be biologically meaningless.

This raises the following issues:

- Can we characterize which information is abstracted away?
- Which is the form of the systems, for which we have no false positive?
- Do we learn something about the biological systems that we describe?
Which information is abstracted away?

**Theorem 2** We suppose that:

1. \((D, \subseteq)\) be a partial order;
2. \((D^\#, \subseteq, \cup)\) be chain-complete partial order;
3. \(D \times \frac{\gamma}{\alpha} D^\#\) be a Galois connection;
4. \(F \in D \to D\) and \(F^\# \in D^\# \to D^\#\) are monotonic;
5. \(F \circ \gamma \subseteq \gamma \circ F^\#\);
6. \(X_0, \text{inv} \in D\) such that:
   - \(X_0 \subseteq F(X_0) \subseteq F(\text{inv}) \subseteq \text{inv}\),
   - \(\text{inv} = \gamma(\alpha(\text{inv}))\),
   - and \(\alpha(F(\text{inv})) = F^\#(\alpha(\text{inv}))\);

Then, \(\text{lfp}_{\alpha(X_0)F^\#}\) exists and \(\gamma(\text{lfp}_{\alpha(X_0)F^\#}) \subseteq \text{inv}\).
Proof I/III

We have already seen (previous lectures) that:

1. $\text{lfp}_{\alpha(X_0)} F^\#$ exists;

2. there exists an ordinal $\delta$ such that $\text{lfp}_{\alpha(X_0)} F^\# = F^\#(\alpha(X_0))$. 
Proof II/III

Let us show that $\gamma(\text{lfp}_\alpha(X_0)\prod^\#) \subseteq \text{inv}$.

Let us prove instead by induction over $\delta$ that $\prod^\#(\alpha(X_0)) \subseteq \alpha(\text{inv})$.

- If $Y \in D^\#$ is an element such that $Y \subseteq \alpha(\text{inv})$,
  $\prod^\#(Y) \subseteq \prod^\#(\alpha(\text{inv}))$ (as $\prod^\#$ is mon)
  $\prod^\#(\alpha(\text{inv})) = \alpha(\prod(\text{inv}))$ (assumption)
  $\alpha(\prod(\text{inv})) \subseteq \alpha(\text{inv})$. (as $\alpha$ is mon and $\text{inv}$ is a post)

  Thus: $\prod^\#(Y) \subseteq \alpha(\text{inv})$

- If $Y_i \in D^{\#I}$ is a chain of elements such that $Y_i \subseteq \alpha(\text{inv})$ for any $i \in I$,
  then, $\sqcup Y_i \subseteq \alpha(\text{inv})$ (lub).

So: $\prod^{\#\delta}(\alpha(X_0)) \subseteq \alpha(\text{inv})$. 
We have:

\[ \mathbb{F}^\# \delta(\alpha(X_0)) \subseteq \alpha(\text{inv}). \]

Since \( \gamma \) is monotonic:

\[ \gamma(\mathbb{F}^\# \delta(\alpha(X_0))) \subseteq \gamma(\alpha(\text{inv})). \]

But, by assumption, \( \gamma(\alpha(\text{inv})) = \text{inv} \).

Thus,

\[ \gamma(\mathbb{F}^\# \delta(\alpha(X_0))) \subseteq \text{inv}. \]
When is there no false positive?

**Theorem 3** We suppose that:

1. \((D, \subseteq, \cup)\) and \((D^\#, \subseteq, \sqcup)\) are chain-complete partial orders;
2. \((D, \subseteq) \xrightarrow{\gamma} (D^\#, \sqsubseteq)\) is a Galois connection;
3. \(F : D \rightarrow D\) is a monotonic map;
4. \(X_0\) is a concrete element such that \(X_0 \subseteq F(X_0)\);
5. \(F \circ \gamma \subseteq \gamma \circ F^\#\);
6. \(F^\# \circ \alpha = \alpha \circ F \circ \gamma \circ \alpha\).

Then:

- \(\text{lfp}_{X_0} F\) and \(\text{lfp}_{\alpha(X_0)} F^\#\) exist;
- \(\text{lfp}_{X_0} F = \gamma(\alpha(\text{lfp}_{X_0} F)) \iff \text{lfp}_{X_0} F = \gamma(\text{lfp}_{\alpha(X_0)} F^\#)).\)

We need to understand under which assumptions \(\text{lfp}_{X_0} F = \gamma(\alpha(\text{lfp}_{X_0} F)).\)
Local set of chemical species

Definition 1 We say that a set $X \in \wp(Species)$ of chemical species is local if and only if $X \in \gamma(\wp(Local\_view))$.

(ie. a set $X$ is local if and only if $X$ is exactly the set of all the species that are generated by a given set of local views.)
We define the binary relation \( \sim^{\text{SWAP}} \) among tuples \( \text{Species}^* \) of chemical species. We say that \( (C_1, \ldots, C_m) \sim^{\text{SWAP}} (D_1, \ldots, D_n) \) if and only if:

\[
(C_1, \ldots, C_m) \text{ matches with } (C_1, \ldots, C_m)
\]

\[
\text{while } (D_1, \ldots, D_n) \text{ matches with } (D_1, \ldots, D_n)
\]
Swapping closure

**Theorem 4** Let \( X \in \varphi(Species) \) be a set of chemical species.

The two following assertions are equivalent:

1. \( X = \gamma(\alpha(X)) \);
2. for any tuples \((C_i), (D_j) \in Species^*\) such that:
   - \((C_i) \in X^*\),
   - and \((C_i) \xrightarrow{\text{SWAP}} (D_j)\);
we have \((D_j) \in X^*\).
Proof (easier implication way)

If:
• $X = \gamma(\alpha(X))$,
• $(C_i)_{i \in I} \in X^*$,
• and $(C_i)_{i \in I} \overset{SWAP}{\sim} (D_j)_{j \in J}$;

Then:
we have $\alpha(\{C_i \mid i \in I\}) = \alpha(\{D_j \mid j \in J\})$ (because $(C_i) \overset{SWAP}{\sim} (D_j)$)
and $\alpha(\{C_i \mid i \in I\}) \subseteq \alpha(X)$ (because $(C_i) \in X^*$ and $\alpha$ mon);
so $\alpha(\{D_j \mid j \in J\}) \subseteq \alpha(X)$;
so $\{D_j \mid j \in J\} \subseteq \gamma(\alpha(X))$ (by def. of Galois connections);
so $\{D_j \mid j \in J\} \subseteq X$ (since $X = \gamma(\alpha(X))$);
so $(D_j)_{j \in J} \in X^*$. 
Proof: more difficult implication way

For any $X \in \wp(\text{Local\_view})$, $\gamma(X)$ is given by a rewrite system:
For any $lv \in X$, we add the following rules:

$I$ and semi-links are non-terminal.
$I$ is the initial symbol.
Proof (more difficult implication way)

We suppose that $X$ is close with respect to $\sim_{\text{SWAP}}$. We want to prove that $\gamma(\alpha(X)) \subseteq X$.

We prove, by induction, that any open complex that can be built by gathering the views of $\alpha(X)$, can be embedded in a complex in $X$:

- By def. of $\alpha$, this is satisfied for any local view in $\alpha(X)$;
- This remains satisfied after unfolding a semi-link with a local view;
- This remains satisfied after binding two semi-links.
Initialization

\[ C \in X \quad \text{(since } l\nu \in \alpha(X)) \]
Unfolding a semi-link

open partial species

$C \in X$

$C' \in X$

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Unfolding a semi-link

open partial species

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Binding two semi-links

\[ C \in X \quad (\text{SWAP}) \]
Consequences

Let $Y \in \wp(Local\_view)$ be a set of local views such that $\alpha(\gamma(Y)) = Y$.

1. Each open complex $C$ built with the local views in $Y$ is a sub-complex of a close complex $C'$ in $\gamma(Y)$.

2. When considering the rewrite system that computes $\gamma(Y)$, any partial rewriting sequence can be completed in a successful one.

Thus:

(a) $\gamma(Y)$ is finite if and only if the grammar has a finite set of prefixes (and the latter is decidable);

(b) We have $F^\sharp \circ \alpha = \alpha \circ F \circ \gamma \circ \alpha$. 
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We have proved that:

- if the set $\text{Species}_\omega$ of reachable chemical species is close with respect to swapping $\sim$,
- then the reachability analysis is exact (i.e. $\text{Species}_\omega = \gamma(\text{lpf}_\alpha(\text{Species}_0)^{\text{F}^d})$).

Now we give some sufficient conditions that ensure this property.
Sufficient conditions

Whenever the following assumptions:

1. initial agents are not bound;
2. rules are atomic;
3. rules are local:
   - only agents that interact are tested,
   - no cyclic patterns (neither in lhs, nor in rhs);
4. binding rules do not interfere i.e. if both:
   - \( A(a\sim m,S),B(b\sim n,T) \rightarrow A(a\sim m!1,S),B(b\sim n!1,T) \)
   - and \( A(a\sim m',S'),B(b\sim n',T') \rightarrow A(a\sim m'!1,S'),B(b\sim n'!1,T') \),
   then:
     - \( A(a\sim m,S),B(b\sim n',T') \rightarrow A(a\sim m!1,S),B(b\sim n'!1,T') \);
5. chemical species in \( \gamma(\alpha(Species_\omega)) \) are acyclic,
   are satisfied, the set of reachable chemical species is local.
Proof outline

We sketch a proof in order to discover sufficient conditions that ensure this property:

- We consider tuples of complexes in which the same kind of links occur twice.
- We want to swap these links.
- We introduce the history of their computation.
- There are several cases...
First case (I/V)
First case (II/V)

just before the links are made

$C \in \text{Species}_{\omega}^{*}$

$C' \in \text{Species}_{\omega}^{*}$
First case (III/V)

we suppose we can swap the links

$C \in \text{Species}_\omega^*$
First case (IV/V)

Then, we ensure that further computation steps:

- are always possible;
- have the same effect on local views;
- commute with the swapping relation $\sim$.

\[
\begin{array}{c}
C_n \xrightarrow{\text{SWAP}, \sigma} C'_n \\
\downarrow R, \phi \\
C_{n+1} \xrightarrow{\text{SWAP}, \sigma} C'_{n+1}
\end{array}
\]

\[
\begin{array}{c}
\downarrow R, \phi \\
C_n \xrightarrow{\text{SWAP}, \sigma} C'_{n+1}
\end{array}
\]
First case (V/V)

\[ C \in \text{Species}_{\omega}^* \]
Second case (I/II)

\[ C \in \text{Species}_\omega \]

we assume that the chemical species \( C \) is acyclic
Second case (II/II)
Sufficient conditions

Whenever the following assumptions:

1. initial agents are not bound;
2. rules are atomic;
3. rules are local:
   - only agents that interact are tested,
   - no cyclic patterns (neither in lhs, nor in rhs);
4. binding rules do not interfere i.e. if both:
   - \( A(a \sim m, S), B(b \sim n, T) \rightarrow A(a \sim m!1, S), B(b \sim n!1, T) \)
   - and \( A(a \sim m', S'), B(b \sim n', T') \rightarrow A(a \sim m'!1, S'), B(b \sim n'!1, T') \),
   then:
   - \( A(a \sim m, S), B(b \sim n', T') \rightarrow A(a \sim m!1, S), B(b \sim n'!1, T') \);
5. chemical species in \( \gamma(\alpha(\text{Species}_\omega)) \) are acyclic,

are satisfied, the set of reachable chemical species is local.
Third case (I/III)
Third case (II/III)

\[ C \in \text{Species}_\omega^* \]
Third case (II/III)

\[ C \in \text{Species}_\omega^* \]
Dangerous sites

A site is dangerous if it may occur in a cycle within a complex ($\in \gamma(\alpha(Species_\omega))$).

We would weaken the fifth requirement into:

- The binding state of a dangerous site is never tested, unless for binding or unbinding this site.
- When we bind dangerous sites, we only test that these sites are free.

Then, we prove that:

1. we can build any complex with free dangerous sites,
2. then, we can bind them as much as we like.
Non local systems

\( \text{Species}_0 \triangleq R(a \sim u) \)

\( \text{Rules} \triangleq \left\{ \begin{array}{l}
R(a \sim u) \iff R(a \sim p) \\
R(a \sim u), R(a \sim u) \rightarrow R(a \sim u!1), R(a \sim u!1) \\
R(a \sim p), R(a \sim u) \rightarrow R(a \sim p!1), R(a \sim p!1) \\
R(a \sim p), R(a \sim p) \rightarrow R(a \sim p!1), R(a \sim p!1)
\end{array} \right\} \)

\( R(a \sim u!1), R(a \sim u!1) \in \text{Species}_\omega \)
\( R(a \sim p!1), R(a \sim p!1) \in \text{Species}_\omega \)
But \( R(a \sim u!1), R(a \sim p!1) \not\in \text{Species}_\omega \).
Non local systems

\[ \text{Species}_0 \quad \triangleq \quad A(a \sim u), B(a \sim u) \]
\[ \text{Rules} \quad \triangleq \quad \begin{cases} 
A(a \sim u), B(a \sim u) \rightarrow A(a \sim u!1), B(a \sim u!1) \\
A(a \sim u!1), B(a \sim u!1) \rightarrow A(a \sim p!1), B(a \sim u!1) \\
A(a \sim u!1), B(a \sim u!1) \rightarrow A(a \sim u!1), B(a \sim p!1)
\end{cases} \]

A(a \sim u!1), B(a \sim p!1) \in \text{Species}_\omega

A(a \sim p!1), B(a \sim u!1) \in \text{Species}_\omega

But A(a \sim p!1), B(a \sim p!1) \notin \text{Species}_\omega.
Non local systems

\[ Species_0 \triangleq A(a \sim u) \]

\[ Rules \triangleq \begin{cases} 
A(a \sim u) \leftrightarrow A(a \sim p) \\
A(a \sim u), A(a \sim p) \rightarrow A(a \sim u!1), A(a \sim p!1) 
\end{cases} \]

\( A(a \sim u!1), A(a \sim p!1) \in Species_\omega \)

But \( A(a \sim p!1), A(a \sim p!1) \notin Species_\omega \).
Non local systems

\[
\begin{align*}
Species_0 & \triangleq R(a,b) \\
Rules & \triangleq \{ R(a,b), R(a) \to R(a,b!1), R(a!1) \}
\end{align*}
\]

\[\begin{align*}
R(a,b!2), R(a!2,b!1), R(a!1,b) & \in Species_\omega \\
\text{But } R(a!1,b!1) & \not\in Species_\omega.
\end{align*}\]
Overview

1. Introduction
2. Language: Kappa
3. Abstraction: Local views
4. Completeness: false positives?
5. Local fragment of Kappa
6. Decontextualization
7. Conclusion
Outline

- we have a syntactic criterion in order to ensure that the set of reachable chemical species of a kappa system is local;
- we now design program transformations to help systems satisfying this criterion;
  1. decontextualization
     - is fully automatic;
     - preserves the transition system;
     - simplifies rules thanks to reachability analysis.
  2. conjugation
     - manual;
     - preserves the set of reachable chemical species;
     - uses backtrack to add new rules.
Example

Initial rule:

\[ R_{2}(l!_{2},r), R_{1}(l!_{1},r), E_{2}(r!_{1}), E_{1}(r!_{2}) \rightarrow R_{2}(l!_{3},r!_{1}), R_{1}(l!_{2},r!_{1}), E_{2}(r!_{2}), E_{1}(r!_{3}) \]

Decontextualized rule:

\[ R_{2}(l!_{\_},r), R_{1}(l!_{\_},r) \rightarrow R_{2}(l!_{\_},r!_{1}), R_{1}(l!_{\_},r!_{1}) \]

We can remove redundant tests.
Example

Initial rules:

\[
\begin{align*}
&\text{Sh}(Y7 \sim p!2, pi!1), G(a!2, b), R(Y48 \sim p!1) \rightarrow \text{Sh}(Y7 \sim p, pi!1), G(a, b), R(Y48 \sim p!1) \\
&\text{Sh}(Y7 \sim p!3, pi!1), G(a!3, b!2), S(d!2), R(Y48 \sim p!1) \rightarrow \text{Sh}(Y7 \sim p, pi!1), G(a, b!2), S(d!2), R(Y48 \sim p!1) \\
&\quad \text{Sh}(Y7 \sim p!1, pi), G(a!1, b) \rightarrow \text{Sh}(Y7 \sim p, pi), G(a, b) \\
&\quad \text{Sh}(Y7 \sim p!1, pi), G(a!1, b!), \rightarrow \text{Sh}(Y7 \sim p, pi), G(a, b!,) \\
\end{align*}
\]

Decontextualized rule:

\[
\text{Sh}(Y7!1), G(a!1) \rightarrow \text{Sh}(Y7), G(a)
\]

We can remove exhaustive enumerations.
How does it work?

To remove a test, we prove that:

- this test is satisfied whenever the other tests are satisfied;
- or each complex that passes all tests but this one also matches with the left hand side of another rule that performs the same action.
More formally

More formally:

- Each rule $R$ is associated with the set $S(R)$ of open chemical species that can match its lhs;
- Rules are gathered in equivalence classes according to the actions they perform;
- For each class $[R]$, we compute:

$$G([R]) = \cup\{S(R') \mid R' \in [R]\}.$$

- For each class $[R]$, $Reach([R])$ is an over approximation of the set of open chemical species that may match the lhs of a rule $R' \in [R]$.

A rule $R$ may be decontextualized in a rule $R'$ if:

$$S(R') \cap Reach([R]) \subseteq G([R]).$$

Decontextualization is more efficient, if the reachability analysis is accurate.
An undecontextualizable rule

Initial rule:

\[ \text{Sh}(Y7\sim u, pi!1), R(Y48\sim p!1, r!_) \rightarrow \text{Sh}(Y7\sim p, pi!1), R(Y48\sim p!1, r!_) \]

Decontextualized rule:

\[ \text{Sh}(Y7\sim u, pi!1), R(Y48!1, r!_) \rightarrow \text{Sh}(Y7\sim p, pi!1), R(Y48!1, r!_) \]
Conjugation

If a rule $R'$ is equivalent to a rule in the transitive closure of the system. Then it may be included in the system without modifying reachable states. To remove the context $C$ of a rule, we try to apply it for another context $C'$ by:

1. removing the context $C'$ (backtrack) ;
2. building the context $C$ ;
3. applying the initial rule ;
4. removing the context $C$ (backtrack) ;
5. building the context $C'$.

This is proved manually.
Overview

1. Introduction
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Conclusion

• A scalable static analysis to abstract the reachable chemical species.
• A class of models for which the abstraction is complete.
• Many applications:
  – idiomatic description of reachable chemical species;
  – dead rule detection;
  – rule decontextualization;
  – computer-driven kinetic refinement.
• It can also help simulation algorithms:
  – wake up/inhibition map (agent-based simulation);
  – flat rule system generation (for bounded set of chemical species);
  – on the fly flat rule generation (for large/unbounded set)
4ième École Thématique
MODÉLISATION FORMELLE DE RÉSEAUX DE RÉGULATION BIOLOGIQUE

Reduction of models of intra-cellular signalling pathways

Jérôme Feret
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Tuesday, the 25th of June, 2019
On the menu today

1. Context and motivations
2. Case studies
3. Reduction of ordinary differential equations
4. Abstraction of the information flow
5. Model reduction
6. Conclusion
Intra-cellular signalling pathways

EGF, TGF-alpha, etc

EGFR

PI3-K

AKT

mTOR

phosphorylation

STAT

GRB2

SOS

RAS

RAF

MEK

ERK

Gene transcription
Cell cycle progression

Cell proliferation
Inhibition of apoptosis
Angiogenesis
Migration, Adhesion, Invasion

Eikuch, 2007
Interaction maps

Oda et al, 2005
Models of the behaviour of the system

\[
\begin{aligned}
\frac{dx_1}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_2}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_3}{dt} &= k_1 \cdot x_1 \cdot x_2 - k_{-1} \cdot x_3 + 2 \cdot k_2 \cdot x_3 \cdot x_3 - k_{-2} \cdot x_4 \\
\frac{dx_4}{dt} &= k_2 \cdot x_3^2 - k_2 \cdot x_4 + \frac{v_4 \cdot x_5}{p_4 + x_5} - k_3 \cdot x_4 - k_{-3} \cdot x_5 \\
\frac{dx_5}{dt} &= \ldots \\
\vdots \\
\frac{dx_{n \ell}}{dt} &= -k_1 \cdot x_1 \cdot c_2 + k_{-1} \cdot x_3 
\end{aligned}
\]
Bridge the gap between...
Site-graphs rewriting

- a language close to knowledge representation;
- rules are easy to update;
- a compact description of models.
Choices of semantics

interaction map

Markov chain

ordinary differential equations

\begin{align*}
\frac{dx_1}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_2}{dt} &= -k_1 \cdot x_1 \cdot x_2 + k_{-1} \cdot x_3 \\
\frac{dx_3}{dt} &= k_1 \cdot x_1 \cdot x_2 - k_{-1} \cdot x_3 + 2 \cdot k_2 \cdot x_3 \cdot x_3 - k_{-2} \cdot x_4 \\
\frac{dx_4}{dt} &= k_2 \cdot x_3^2 - k_2 \cdot x_4 + \frac{y_4 \cdot x_5}{p_{4}x_5} - k_3 \cdot x_4 - k_{-3} \cdot x_5 \\
\frac{dx_5}{dt} &= \cdots \\
\frac{dx_n}{dt} &= -k_1 \cdot x_1 \cdot c_2 + k_{-1} \cdot x_3
\end{align*}
Abstractions offer different perspectives on models

concrete semantics

causal traces

information flow

exact projection of the ODE semantics
Causal traces
ODE semantics

EGF pathway (reduced ODEs)

Concentration

Time

Jérôme Feret
Causal traces
Combinatorial wall
A potential breach
A potential breach
On the menu today

1. Context and motivations
2. Case studies
3. Reduction of ordinary differential equations
4. Abstraction of the information flow
5. Model reduction
6. Conclusion
Case study
Case study

[Diagram of chemical reactions]

Jérôme Feret
18
Tuesday, the 25th of June, 2019
Law of mass action

We consider that chemical species are elementary particles without any volume, and that they are diffusing in an infinite, perfectly fluid and homogeneous medium without borders.

Let \( \mathcal{X} \) be a set of chemical species.

A reaction network is a set of reactions \( \mathcal{R} \).

Each reaction \( r \) is defined by:

1. \( \alpha_r \), a function from \( \mathcal{X} \) to \( \mathbb{N} \) (the reactants);
2. \( \beta_r \), a function from \( \mathcal{X} \) to \( \mathbb{N} \) (the products);
3. \( k_r \), a non negative real number (the kinetic rate).

With these notations, the law of mass action defines the behaviour of the concentration \( [X] \) of each chemical species \( X \):

\[
\frac{d[X]}{dt} = \sum_{r \in \mathcal{R}} k_r \cdot (\beta_r(X) - \alpha_r(X)) \cdot \prod_{X' \in \mathcal{X}} [X']^{\alpha_r(X')}. 
\]
Case study

\[ \begin{align*}
    \frac{d[(u,u,u)]}{dt} &= -k_c \cdot [(u,u,u)] \\
    \frac{d[(u,p,u)]}{dt} &= k_c \cdot [(u,u,u)] \\
\end{align*} \]
Case study

\[
\begin{align*}
\frac{d[(u,u,u)]}{dt} &= -k_c \cdot [(u,u,u)] \\
\frac{d[(u,p,u)]}{dt} &= -k_g \cdot [(u,p,u)] + k_c \cdot [(u,u,u)] - k_d \cdot [(u,p,u)] \\
\frac{d[(u,p,p)]}{dt} &= -k_g \cdot [(u,p,p)] + k_d \cdot [(u,p,u)] \\
\frac{d[(p,p,u)]}{dt} &= k_g \cdot [(u,p,u)] - k_d \cdot [(p,p,u)] \\
\frac{d[(p,p,p)]}{dt} &= k_g \cdot [(u,p,p)] + k_d \cdot [(p,p,u)]
\end{align*}
\]
Case study
Case study
Case study

\[
\begin{align*}
[(u,u,u)] &= [(u,u,u)] \\
[(u,p,?)] &\overset{\Delta}{=} [(u,p,u)] + [(u,p,p)] \\
[(p,p,?)] &\overset{\Delta}{=} [(p,p,u)] + [(p,p,p)]
\end{align*}
\]

\[
\begin{align*}
\frac{d[(u,u,u)]}{dt} &= -k_c \cdot [(u,u,u)] \\
\frac{d[(u,p,?)]}{dt} &= -k_g \cdot [(u,p,?)] + k_c \cdot [(u,u,u)] \\
\frac{d[(p,p,?)]}{dt} &= k_g \cdot [(u,p,?)]
\end{align*}
\]
What we have learned so far:

We can use the absence of information flow to detect useless correlations between the states of sites in chemical species. We can use this to cut chemical species into fragments.

This transformation loses some information: we cannot compute the concentration of each chemical species anymore.
On the menu today

1. Context and motivations
2. Case studies
3. Reduction of ordinary differential equations
4. Abstraction of the information flow
5. Model reduction
6. Conclusion
Differential semantics

A system of ordinary differential equations is a pair \((\mathcal{V}, F)\) where:

- \(\mathcal{V}\) is a finite set of variables,
- \(F\) is a continuous function from \(\mathcal{V} \rightarrow \mathbb{R}^+\) to \(\mathcal{V} \rightarrow \mathbb{R}\).

Elements of \(\mathcal{V} \rightarrow \mathbb{R}^+\) are called states.

The differential semantics maps each initial state \(X_0 \in \mathcal{V} \rightarrow \mathbb{R}^+\) to the solution \(X_{X_0} \in \mathbb{R}^+\) of the following equation:

\[
X_{X_0}(T) = X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot dt.
\]

that is defined over the widest time interval as possible.
Back to the case study

1. $\mathcal{N} \triangleq \{[(u,u,u)], [(u,p,u)], [(p,p,u)], [(u,p,p)], [(p,p,p)]\},$

2. $\mathbf{F}(\rho) \triangleq \left\{ \begin{array}{c}
[(u,u,u)] \mapsto -k_c \cdot \rho([(u,u,u)]) \\
[(u,p,u)] \mapsto -k_g \cdot \rho([(u,p,u)]) + k_c \cdot \rho([(u,u,u)]) - k_d \cdot \rho([(u,p,u)]) \\
[(u,p,p)] \mapsto -k_g \cdot \rho([(u,p,p)]) + k_d \cdot \rho([(u,p,u)]) \\
[(p,p,u)] \mapsto k_g \cdot \rho([(u,p,u)]) - k_d \cdot \rho([(p,p,u)]) \\
[(p,p,p)] \mapsto k_g \cdot \rho([(u,p,p)]) + k_d \cdot \rho([(p,p,u)]). \end{array} \right.$
Abstraction

An abstraction is a 5-uple \((\mathcal{V}, \mathcal{F}, \mathcal{V}^\#, \psi, \mathcal{F}^\#)\), where:

- \((\mathcal{V}, \mathcal{F})\) is a system of ordinary equations,
- \(\mathcal{V}^\#\) is a finite set of observables,
- \(\psi\) is a function from the set \(\mathcal{V} \to \mathbb{R}\) into the set \(\mathcal{V}^\# \to \mathbb{R}\),
- \(\mathcal{F}^\#\) is a function \(\mathcal{C}^\infty\) from the set \(\mathcal{V}^\# \to \mathbb{R}^+\) into the set \(\mathcal{V}^\# \to \mathbb{R}\);

such that:

- \(\psi\) is linear with positive coefficients only and such that each variable \(v \in \mathcal{V}\) occurs in the image of at least one observable \(v^\# \in \mathcal{V}^\#\) with a non-zero coefficient;
- the following diagram commutes:

\[
\begin{array}{ccc}
(\mathcal{V} \to \mathbb{R}^+) & \xrightarrow{\mathcal{F}} & (\mathcal{V} \to \mathbb{R}) \\
\downarrow \psi & & \downarrow \psi \\
(\mathcal{V}^\# \to \mathbb{R}^+) & \xrightarrow{\mathcal{F}^\#} & (\mathcal{V}^\# \to \mathbb{R})
\end{array}
\]

that is to say that \(\psi \circ \mathcal{F} = \mathcal{F}^\# \circ \psi\).
Back to the case study

1. $\mathcal{V} \overset{\Delta}{=} \{(u,u,u), (u,p,u), (p,p,u), (u,p,p), (p,p,p)\}$

   $\begin{cases} 
   (u,u,u) \mapsto -k_c \cdot \rho([(u,u,u)]) \\
   (u,p,u) \mapsto -k_g \cdot \rho([(u,p,u)]) + k_c \cdot \rho([(u,u,u)]) - k_d \cdot \rho([(u,p,u)]) \\
   (u,p,p) \mapsto -k_g \cdot \rho([(u,p,p)]) + k_c \cdot \rho([(u,u,u)]) - k_d \cdot \rho([(u,p,u)]) \\
   \ldots
   \end{cases}$

2. $\mathbb{F}(\rho) \overset{\Delta}{=} \{ 
   (u,p,u) \mapsto -k_g \cdot \rho([(u,p,u)]) + k_c \cdot \rho([(u,u,u)]) - k_d \cdot \rho([(u,p,u)]) \\
   (u,p,p) \mapsto -k_g \cdot \rho([(u,p,p)]) + k_d \cdot \rho([(u,p,u)]) \\
   \ldots
   \}$

3. $\mathcal{V}^\sharp \overset{\Delta}{=} \{(u,u,u), (? ,p ,u), (? ,p ,p), (u ,p ,?), (p ,p ,?)\}$

   $\begin{cases} 
   (u,u,u) \mapsto \rho([(u,u,u)]) \\
   (?,p,u) \mapsto \rho([(u,p,u)]) + \rho([(p,p,u)]) \\
   (?,p,p) \mapsto \rho([(u,p,p)]) + \rho([(p,p,p)]) \\
   \ldots
   \end{cases}$

4. $\psi(\rho) \overset{\Delta}{=} \{ 
   (u,p,u) \mapsto \rho([(u,p,u)]) + \rho([(p,p,u)]) \\
   (u,p,p) \mapsto \rho([(u,p,p)]) + \rho([(p,p,p)]) \\
   \ldots
   \}$

5. $\mathbb{F}^\sharp(\rho^\sharp) \overset{\Delta}{=} \{ 
   (u,u,u) \mapsto -k_c \cdot \rho^\sharp([(u,u,u)]) \\
   (?,p,u) \mapsto -k_d \cdot \rho^\sharp([(?,p,u)]) + k_c \cdot \rho^\sharp([(u,u,u)]) \\
   (?,p,p) \mapsto k_d \cdot \rho^\sharp([(?,p,u)]) \\
   \ldots
   \}$
Let us apply the abstraction function

Let:

1. \((V, F, \nu, \psi, \nu')\) be an abstraction,
2. and \(X_0 \in V \rightarrow \mathbb{R}^+\) be an initial state.

We have, at any time \(T\) within the time interval \([0, T_{X_0}^{\text{max}}]\):

\[ X_{X_0}(T) = X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot \text{d}t. \]

So:

\[ \psi(X_{X_0}(T)) = \psi \left( X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot \text{d}t \right). \]
Let us push $\psi$ towards the right

Let:

1. $(\mathcal{V}, \mathcal{F}, \mathcal{V}^\sharp, \psi, \mathcal{F}^\sharp)$ be an abstraction,
2. and $X_0 \in \mathcal{V} \to \mathbb{R}^+$ be an initial state.

We have, at any time $T$ within the time interval $[0, T_{X_0}^{\text{max}}]$:

$$X_{X_0}(T) = X_0 + \int_{t=0}^{T} \mathcal{F}(X_{X_0}(t)) \cdot dt.$$ 

So:

$$\psi(X_{X_0}(T)) = \psi(X_0) + \psi \left( \int_{t=0}^{T} \mathcal{F}(X_{X_0}(t)) \cdot dt \right).$$
Let us push $\psi$ towards the right

Let:

1. $(\mathcal{V}, F, \mathcal{V}^\sharp, \psi, F^\sharp)$ be an abstraction,
2. and $X_0 \in \mathcal{V} \to \mathbb{R}^+$ be an initial state.

We have, at any time $T$ within the time interval $[0, T_{X_0}^{\max}]$:

$$X_{X_0}(T) = X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot dt.$$ 

So:

$$\psi(X_{X_0}(T)) = \psi(X_0) + \int_{t=0}^{T} [\psi \circ F](X_{X_0}(t)) \cdot dt.$$
Let us push $\psi$ towards the right

Let:

1. $(\mathcal{V}, F, \mathcal{V}^\sharp, \psi, F^\sharp)$ be an abstraction,
2. and $X_0 \in \mathcal{V} \rightarrow \mathbb{R}^+$ be an initial state.

We have, at any time $T$ within the time interval $[0, T_{\max}^X]$: 

$$X_{X_0}(T) = X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot dt.$$ 

So:

$$\psi(X_{X_0}(T)) = \psi(X_0) + \int_{t=0}^{T} [F^\sharp \circ \psi](X_{X_0}(t)) \cdot dt.$$
Let us push $\psi$ towards the right

Let:

1. $(\mathcal{V}, F, \mathcal{V}^\#, \psi, F^\#)$ be an abstraction,
2. and $X_0 \in \mathcal{V} \to \mathbb{R}^+$ be an initial state.

We have, at any time $T$ within the time interval $[0, T_{\text{max}}^{X_0}[$:

$$X_{X_0}(T) = X_0 + \int_{t=0}^{T} F(X_{X_0}(t)) \cdot dt.$$

So:

$$\psi(X_{X_0}(T)) = \psi(X_0) + \int_{t=0}^{T} F^\#(\psi(X_{X_0}(t))) \cdot dt.$$

Jérôme Feret 28 Tuesday, the 25th of June, 2019
Abstract semantics

Let \((\mathcal{V}, \mathcal{F}, \mathcal{V}^\#, \psi, \mathcal{F}^\#)\) be an abstraction.
The couple \((\mathcal{V}^\#, \mathcal{F}^\#)\) is a system of differential equations.
Let us denote by \(Y\) its semantics.
For each state \(Y_0 \in \mathcal{V}^\# \rightarrow \mathbb{R}^+\), we denote by \([0, T_{\max}^{\# Y_0}]\) the domain of the function \(Y_{Y_0}\). We have, at any time \(T^\# \in [0, T_{\max}^{\# X_0}],\)
\[
Y_{Y_0}(T^\#) = Y_0 + \int_{t=0}^{T^\#} \mathcal{F}^\#(Y_{Y_0}(t)) \cdot dt.
\]

**Theorem 1** For each initial state \(X_0 \in \mathcal{V} \rightarrow \mathbb{R}^+\), we have:

1. \(T_{\max}^{\# \psi(X_0)} = T_{\max}^{\# X_0}\);
2. at any time \(T \in [0, T_{\max}^{\# X_0}], \psi(X_{X_0}(T)) = Y_{\psi(X_0)}(T)\).

That is to say that the abstract semantics is the image of the concrete semantics by the abstraction function.
Abstract trajectories

\[ Y(t) \]

\[ t \]
Concrete trajectories

Y(t)
X(t)
t
On the menu today

1. Context and motivations
2. Case studies
3. Reduction of ordinary differential equations
4. Abstraction of the information flow
5. Model reduction
6. Conclusion
Concrete semantics

A rule is a symbolic representation of a multi-set of reactions.

For instance, the rule:

\[
/CT/D1/CT/D2 /D8/D7
\]

\[k_d\]

denotes the following two rules:

\[
/CT/D1/CT/D2 /D8/D7
\]

\[k_d\]

\[
/CT/D1/CT/D2 /D8/D7
\]

\[k_d\]

The semantics of a set of rules is the semantics of the underlying multi-set of reactions.
Flow of information (in the concrete)

Does the state of a given site influence the capability to modify another site?
Flow of information (in the concrete)
Flow of information (in the concrete)

If there exists a soup of chemical species in which the activation rate of the site of $\text{ShC}$ is different in these two contexts, then there may be a flow of information.
Discrimination by a rule

In this case, there exists a rule which makes a difference between these two contexts, for instance the following one:
Flow of information due to a rule
Flow of information due to a rule
Flow of information due to a rule
Flow of information due to a rule
Flow of information due to a rule
Projection on the contact map
Projection on the contact map
Projection on the contact map
Projection on the contact map

[Diagram of protein interactions involving EGF, EGFR, ShC, pi, Y7, Grb2, and Sos]
Projection on the contact map
Direct computation
Direct computation
Direct computation
Direct computation

Diagram of molecular interactions involving EGFR and Grb2 pathways.
Direct computation
On the menu today

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6. Conclusion
Which patterns shall we keep?
Which patterns shall we keep?
Pattern annotation
Definition 1 (prefragment) A pattern is a prefragment if, in its annotated form, there exists a site that it is reachable from every site (following the flow of information).
Definition 2 (fragment) A fragment is a prefragment that cannot be embedded in any bigger prefragment.
Examples

Which patterns are fragments?

- EGF
- EGFR
- Y48
- Y68
Examples: annotated map

EGF → EGFR

ShC → pi

EGF → EGFR

Grb2 → Sos

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Examples: pattern annotation
Examples
Which patterns are prefragments?
Examples
Prefragments

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Examples
Which patterns are fragments?
Examples
Fragments

EGF

EGF

EGFR

EGF

EGFR

Y48

EGF

EGFR

Y68

EGF

EGFR

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Examples: fragments
Almost done…

We are left to express the consumption and the production (in concentration) of each fragment as expressions of the concentration of fragments.

Firstly, we notice that the concentration of each prefragment can be expressed as a linear combination of the concentration of the fragments.
Whenever there is an overlap between a fragment and a connected component in the left-hand side of a rule such that the common region contains a site that is modified by the rule, then the connected component embeds in the fragment.
Fragments consumption

Whenever there is an overlap between a fragment and a connected component in the left hand side of a rule such that the common region contains a site that is modified by the rule, then the connected component embeds in the fragment.
Fragments consumption

For each fragment $F$, for each rule:

$$r: C_1, \ldots, C_n \rightarrow \text{rhs} \quad k$$

and for each occurrence of a connected component $C_j$ that is modified by the rule, in the fragment $F$, we have the following contribution:

$$\frac{d[F]}{dt} = \frac{k \cdot [F] \cdot \prod_{i \neq j} [C_i]}{\text{SYM}[C_1, \ldots, C_n] \cdot \text{SYM}[F]}.$$
Fragments production

Whenever there is an overlap between a fragment and the right hand side of a rule, such that the common region contains a site that is modified by the rule.

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Whenever there is an overlap between a fragment and the right hand side of a rule, such that the common region contains a site that is modified by the rule...
Whenever there is an overlap between a fragment and the right hand side of a rule such that the common region contains a site that is modified by the rule, each connected component in the left hand side of the refined rule, is a prefragment.
Fragment production

For each overlap $ch$ between a fragment and the right hand side of a rule, such that the common region contains a site that is modified by the rule:

$$r : C_1, \ldots, C_m \rightarrow \text{right hand side} \quad k,$$

we have the following contribution:

$$\frac{d[F]}{dt} = k \cdot \prod_i [C'_i] \cdot \text{SYM}[F].$$

where $C'_1, \ldots, C'_n$ is the left hand side of the refined rule.
On the menu today

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2. Case studies
3. Reduction of ordinary differential equations
4. Abstraction of the information flow
5. Model reduction
6. Conclusion
## Benchmark

<table>
<thead>
<tr>
<th>Model</th>
<th>early EGF</th>
<th>EGF/Insulin</th>
<th>SFB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of molecular species</td>
<td>356</td>
<td>2899</td>
<td>$\sim 2.10^{19}$</td>
</tr>
<tr>
<td>Number of fragments (ODEs semantics)</td>
<td>38</td>
<td>208</td>
<td>$\sim 2.10^5$</td>
</tr>
<tr>
<td>Number of fragments (CTMC semantics)</td>
<td>356</td>
<td>618</td>
<td>$\sim 2.10^{19}$</td>
</tr>
</tbody>
</table>
In short
Abstraction of the information flow
Abstraction of the information flow
Patterns of interest
Patterns of interest

EGF\n\nEGFR\n\nY48\n
EGF\n\nEGFR\n\nY68

ShC\npi\Y7
Grb2\na\b Sos

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Related topics and acknowledgements

- Model reduction (ODEs semantics)
  Vincent Danos, Walter Fontana, Russ Harmer, Jean Krivine
- Context-sensitive abstraction of information flow
  Ferdinanda Camporesi
- Model reduction (CTMC semantics)
  Tatjana Petrov, Heinz Koeppl, Tom Henzinger
- Bisimulations metrics
  Norm Ferns.