Probabilistic Programming for Stochastic Dynamical Systems (ProPPA)

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Outline

1. Stochastic Process Algebras

2. Modelling Biological Processes
   - Bio-PEPA
   - Approaches to system biology modelling

3. ProPPA

4. Inference

5. Results

6. Conclusions
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1 Stochastic Process Algebras

2 Modelling Biological Processes
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5 Results

6 Conclusions
Models are constructed from components which engage in activities.

\[(\alpha, r).P\]

- **Action type or name**
- **Activity rate** (parameter of an exponential distribution)
- **Component/derivative**
Stochastic Process Algebra

- Models are constructed from components which engage in activities.

\[(\alpha, r).P\]

- The language is used to generate a Continuous Time Markov Chain (CTMC) for performance modelling.

```
SPA MODEL  SOS rules  LABELLED MULTITRANSITION SYSTEM  state transition diagram  CTMC Q
```
Benefits of integration

- Properties of the underlying mathematical structure may be deduced by the construction at the process algebra level.
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- Formal reasoning techniques such as equivalence relations and model checking can be used to manipulate or interrogate models.

- For example the congruence Markovian bisimulation, allows exact model reduction to be carried out compositionally.

- Stochastic model checking based on the Continuous Stochastic Logic (CSL) allows automatic evaluation of quantified properties of the behaviour of the system.
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In some cases there have been new languages developed to support particular features of the application domain. These have included stochastic process algebras for modelling hybrid systems (e.g. HYPE, HyPA), spatial temporal collective adaptive systems (e.g. PALOMA, CARMA) and ecological processes (e.g. PALPS, MELA).
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This is most noticeable in the arena of systems biology, which is often focussed on biomolecular processing systems, for example Bio-PEPA.
### Molecular processes as concurrent computations

<table>
<thead>
<tr>
<th>Concurrency</th>
<th>Molecular Biology</th>
<th>Metabolism</th>
<th>Signal Transduction</th>
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<td>Concurrent computational processes</td>
<td>Molecules</td>
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<td>Binding and catalysis</td>
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<tr>
<td>Transition or mobility</td>
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Each “species” (biochemical component) $i$ is described by a species component $C_i$. \

The Bio-PEPA abstraction
The Bio-PEPA abstraction

- Each "species" (biochemical component) $i$ is described by a species component $C_i$.
- Each reaction $j$ is associated with an action type $\alpha_j$ and its dynamics is described by a specific function $f_{\alpha_j}$. 
The Bio-PEPA abstraction

- Each “species” (biochemical component) $i$ is described by a species component $C_i$

- Each reaction $j$ is associated with an action type $\alpha_j$ and its dynamics is described by a specific function $f_{\alpha_j}$

The species components are then composed together to describe the behaviour of the system.
Bio-PEPA modelling

- The state of the system at any time consists of the local states of each of its sequential/species components.
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Bio-PEPA modelling

- The **state of the system** at any time consists of the **local states** of each of its sequential/species components.

- The local states of components are **quantitative** rather than functional, i.e. biological changes to species are represented as distinct components.

- A component varying its state corresponds to it varying its **amount**.

- This is captured by an integer parameter associated with the species and the effect of a reaction is to vary that parameter by a number corresponding to the **stoichiometry** of this species in the reaction.
The syntax

Sequential component (species component)

\[ S ::= (\alpha, \kappa) \text{ op } S \mid S + S \mid C \quad \text{where } \text{op} = \downarrow \mid \uparrow \mid \oplus \mid \ominus \mid \odot \]
The syntax

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Model component

\[ P ::= P \otimes P \mid S(I) \]
The syntax

Sequential component (species component)

\[ S ::= (\alpha, \kappa) \ op \ S | S + S | C \]

where \( \op = \downarrow | \uparrow | \oplus | \ominus | \odot \)

Model component

\[ P ::= P \boxdot_{L} P | S(I) \]
The syntax

Sequential component (species component)

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Model component

\[ P ::= P \boxdot_{\mathcal{L}} P \mid S(l) \]
The syntax

Sequential component (species component)

\[ S ::= (\alpha, \kappa) \ op \ S | S + S | C \quad \text{where } \ op = \downarrow | \uparrow | \oplus | \ominus | \bigcirc \]

Model component

\[ P ::= P \boxtimes P | S(I) \]

Each action \( \alpha_j \) is associated with a rate \( f_{\alpha_j} \)
The semantics is defined by two transition relations:

- First, a **capability relation** — is a transition possible?

- Second, a **stochastic relation** — gives rate of a transition, derived from the parameters of the model.
Example

\[ I \xrightarrow{\text{spread}} S \xrightarrow{\text{stop}1} R \]

\[ I = (\text{spread},1) \]
\[ S = (\text{spread},1) \uparrow + (\text{stop}1,1) \downarrow + (\text{stop}2,1) \downarrow \]

\[ R = (\text{stop}1,1) \uparrow + (\text{stop}2,1) \uparrow \]

\[ I[10] \times S[5] \times R[0] \]

\[ k_s = 0.5; \]
\[ k_r = 0.1; \]

\[ \text{kineticLawOf spread : } k_s \times I \times S; \]
\[ \text{kineticLawOf stop1 : } k_r \times S \times S; \]
\[ \text{kineticLawOf stop2 : } k_r \times S \times R; \]
Example

\[
\begin{align*}
I & \xrightarrow{\text{spread}} S \\
S & \xrightarrow{\text{stop1}} S \xrightarrow{\text{stop2}} R \\
I & \xrightarrow{\text{spread}} S \\
S & \xrightarrow{\text{stop1}} S \xrightarrow{\text{stop2}} R
\end{align*}
\]

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\]

\[
I[10] \times S[5] \times R[0]
\]
Formal modelling in systems biology

- Formal languages like Bio-PEPA provide a convenient interface for describing complex systems, reflecting what is known about the biological components and their behaviour.
- High-level abstraction eases writing and manipulating models.
- They are compiled into executable models which can be run to deepen understanding of the model.
- Formal nature lends itself to automatic, rigorous methods for analysis and verification.
- Executing the model generates data that can be compared with biological data.

... but what if parts of the system are unknown?

Usual process of parameterising a model is iterative and manual.
Model creation is data-driven
Machine Learning: Bayesian statistics

Represent belief and uncertainty as probability distributions (prior, posterior). Treat parameters and unobserved variables similarly. Bayesian Theorem:

$$P(\theta|D) = \frac{P(\theta) \cdot P(D|\theta)}{P(D)}$$

posterior $\propto$ prior $\cdot$ likelihood
Machine Learning: Bayesian statistics

- Represent belief and uncertainty as probability distributions (prior, posterior).
- Treat parameters and unobserved variables similarly.
- Bayes’ Theorem:

\[
P(\theta \mid D) = \frac{P(\theta) \cdot P(D \mid \theta)}{P(D)}
\]

posterior \propto prior \cdot likelihood
Bayesian statistics
Bayesian statistics
Bayesian statistics

After 40 data points

[Graph showing a Bayesian curve with data points]
Bayesian statistics
Thus there are two approaches to model construction:

**Machine Learning:** extracting a model from the data generated by the system, or refining a model based on system behaviour using statistical techniques.

**Mechanistic Modelling:** starting from a description or hypothesis, construct a formal model that algorithmically mimics the behaviour of the system, validated against data.
Comparing the techniques

Data-driven modelling:

+ rigorous handling of parameter uncertainty
- limited or no treatment of stochasticity
- in many cases bespoke solutions are required which can limit the size of system which can be handled
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Data-driven modelling:

+ rigorous handling of parameter uncertainty
- limited or no treatment of stochasticity
- in many cases bespoke solutions are required which can limit the size of system which can be handled

Mechanistic modelling:

+ general execution "engine" (deterministic or stochastic) can be reused for many models
+ models can be used speculatively to investigate roles of parameters, or alternative hypotheses
- parameters are assumed to be known and fixed, or costly approaches must be used to seek appropriate parameterisation
Developing a probabilistic programming approach

What if we could...

- include information about uncertainty in the model?
- automatically use observations to refine this uncertainty?
- do all this in a formal context?

Starting from the existing process algebra (Bio-PEPA), we have developed a new language ProPPA that addresses these issues.

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Probabilistic programming

- A programming paradigm for describing incomplete knowledge scenarios, and resolving the uncertainty.
- Programs probabilistic models in a high level language, like software code.
- Offers automated inference without the need to write bespoke solutions.
- Platforms: IBAL, Church, Infer.NET, Fun, Anglican, Stan, WebPPL, ....
- Key actions: specify a distribution, specify observations, infer posterior distribution.
Describe how the data is generated in syntax like a conventional programming language, but leaving some variables uncertain.
Probabilistic programming workflow

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- Specify observations, which impose constraints on acceptable outputs of the program.
Probabilistic programming workflow

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- **Specify observations**, which impose constraints on acceptable outputs of the program.

- **Run program forwards**: Generate data consistent with observations.
Describe how the data is generated in syntax like a conventional programming language, but leaving some variables uncertain.

Specify observations, which impose constraints on acceptable outputs of the program.

Run program forwards: Generate data consistent with observations.

Run program backwards: Find values for the uncertain variables which make the output match the observations.
A Probabilistic Programming Process Algebra: ProPPA

The objective of ProPPA is to retain the features of the stochastic process algebra:

- simple model description in terms of components
- rigorous semantics giving an executable version of the model...
The objective of ProPPA is to retain the features of the stochastic process algebra:

- simple model description in terms of components
- rigorous semantics giving an executable version of the model...

... whilst also incorporating features of a probabilistic programming language:

- recording uncertainty in the parameters
- ability to incorporate observations into models
- access to inference to update uncertainty based on observations
Example Revisited

\[ k_s = 0.5; \]
\[ k_r = 0.1; \]

kineticLawOf spread : \( k_s \times I \times S \);
kineticLawOf stop1 : \( k_r \times S \times S \);
kineticLawOf stop2 : \( k_r \times S \times R \);

\[ I = (\text{spread},1) \downarrow ; \]
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Declaring uncertain parameters:
- \( k_s = \text{Uniform}(0,1); \)
- \( k_t = \text{Uniform}(0,1); \)

Providing observations:
- `observe('trace')`

Specifying inference approach:
- `infer('ABC')`
k_s = Uniform(0,1);
k_r = Uniform(0,1);

kineticLawOf spread : k_s * I * S;
kineticLawOf stop1 : k_r * S * S;
kineticLawOf stop2 : k_r * S * R;

I = (spread,1) ↓ ;
S = (spread,1) ↑ + (stop1,1) ↓ + (stop2,1) ↓ ;
R = (stop1,1) ↑ + (stop2,1) ↑ ;


observe('trace')
infer('ABC') //Approximate Bayesian Computation
As with PEPA, a Bio-PEPA model can be interpreted as a CTMC;

However, CTMCs cannot capture uncertainty in the rates (every transition must have a concrete rate).

ProPPA models include uncertainty in the parameters, which translates into uncertainty in the transition rates.

A ProPPA model should be mapped to something like a distribution over CTMCs.
$k = 2$

model

CTMC
$k \in [0,5]$

Parameter model

Set of CTMCs
k \sim p

model

\mu
distribution over CTMCs
Constraint Markov Chains (CMCs) are a generalization of DTMCs, in which the transition probabilities are not concrete, but can take any value satisfying some constraints.

**Constraint Markov Chain**

A CMC is a tuple \( \langle S, o, A, V, \phi \rangle \), where:

- \( S \) is the set of states, of cardinality \( k \).
- \( o \in S \) is the initial state.
- \( A \) is a set of atomic propositions.
- \( V : S \to 2^A \) gives a set of acceptable labellings for each state.
- \( \phi : S \times [0, 1]^k \to \{0, 1\} \) is the constraint function.

In a CMC, arbitrary constraints are permitted, expressed through the function $\phi$: $\phi(s, \vec{p}) = 1$ iff $\vec{p}$ is an acceptable vector of transition probabilities from state $s$.

However,

- CMCs are defined only for the discrete-time case, and
- this does not say anything about how likely a value is to be chosen, only about whether it is acceptable.

To address these shortcomings, we define **Probabilistic Constraint Markov Chains**.
A Probabilistic Constraint Markov Chain is a tuple $\langle S, o, A, V, \phi \rangle$, where:

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- $o \in S$ is the initial state.
- $A$ is a set of atomic propositions.
- $V : S \rightarrow 2^A$ gives a set of acceptable labellings for each state.
- $\phi : S \times [0, \infty)^k \rightarrow [0, \infty)$ is the constraint function.

- This is applicable to continuous-time systems.
- $\phi(s, \cdot)$ is now a probability density function on the transition rates from state $s$. 
Semantics of ProPPA

The semantics definition follows that of Bio-PEPA, which is defined using two transition relations:

- Capability relation — is a transition possible?
- Stochastic relation — gives distribution of the rate of a transition

The distribution over the parameter values induces a distribution over transition rates.
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The distribution over the parameter values induces a distribution over transition rates.

This gives rise the underlying PCMC.
Simulating Probabilistic Constraint Markov Chains

Probabilistic Constraint Markov Chains are open to two alternative dynamic interpretations:

1. **Uncertain Markov Chains**: For each trajectory, for each uncertain transition rate, sample once at the start of the run and use that value throughout;

2. **Imprecise Markov Chains**: During each trajectory, each time a transition with an uncertain rate is encountered, sample a value but then discard it and re-sample whenever this transition is visited again.
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Current work on ProPPA is focused on the **Uncertain Markov Chain** case.
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Inference

Parameter model

\[ k \sim p \]

Distribution over CTMCs

\[ \mu \]
Inference

Inference parameter model

\( k \sim p \)

distribution over CTMCs

\( \mu \)

observations

inference

posterior distribution

\( \mu^* \)
Inference

\[ P(\theta \mid D) \propto P(\theta)P(D \mid \theta) \]

- Exact inference is impossible, as we cannot calculate the likelihood.
- We must use approximate algorithms or approximations of the system.
- The ProPPA semantics does not define a single inference algorithm, allowing for a modular approach.
- Different algorithms can act on different input (time-series vs properties), return different results or in different forms.
Approximate Bayesian Computation is a simple simulation-based solution:

- Approximates posterior distribution over parameters as a set of samples
- Likelihood of parameters is hard to compute in CTMCs, approximates that with a notion of distance.
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\[ \sum (x_i - y_i)^2 > \varepsilon \]

rejected
Approximate Bayesian Computation is a simple simulation-based solution:

- Approximates posterior distribution over parameters as a set of samples
- Likelihood of parameters is hard to compute in CTMCs, approximates that with a notion of distance.

\[ \sum (x_i - y_i)^2 < \varepsilon \]

accepted
Approximate Bayesian Computation

ABC algorithm

1. Sample a parameter set from the prior distribution.
2. Simulate the system using these parameters.
3. Compare the simulation trace obtained with the observations.
4. If distance \(<\epsilon\), accept, otherwise reject.

This results in an approximate to the posterior distribution. As \(\epsilon \to 0\), set of samples converges to true posterior.

We use a more elaborate version based on Markov Chain Monte Carlo sampling.
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Example model

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\[ k_r = \text{Uniform}(0,1); \]

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\[ S = (\text{spread},1) \uparrow + (\text{stop1},1) \downarrow + (\text{stop2},1) \downarrow; \]
\[ R = (\text{stop1},1) \uparrow + (\text{stop2},1) \uparrow; \]

\[ I[10] \otimes S[5] \otimes R[0] \]

\[ \text{observe('trace')} \]
\[ \text{infer('ABC')} //\text{Approximate Bayesian Computation} \]
Results

Tested on the rumour-spreading example, with uniform priors for the two parameters $k_s$ and $k_r$.

- Approximate Bayesian Computation
- Returns posterior as a set of points (samples)
- Observations: sparse time-series (single simulation)
Results: ABC
Results: ABC

![Histograms of $k_r$ and $k_s$]
Genetic Toggle Switch

- Two mutually-repressing genes: promoters (unobserved) and their protein products
- Bistable behaviour: switching induced by environmental changes
- Synthesised in *E. coli*
  

- Stochastic variant where switching is induced by noise
  
Genetic Toggle Switch

G1, on \rightarrow \text{participates} \rightarrow P1 \rightarrow G2, off

\text{accelerates} \leftarrow P2 \leftarrow G2, on

G1, off \rightarrow G1, on
Toggle switch model: species

\[ G_1 = \text{activ1} \uparrow + \text{deact1} \downarrow + \text{expr1} \oplus ; \]
\[ G_2 = \text{activ2} \uparrow + \text{deact2} \downarrow + \text{expr2} \oplus ; \]
\[ P_1 = \text{expr1} \uparrow + \text{degr1} \downarrow + \text{deact2} \oplus ; \]
\[ P_2 = \text{expr2} \uparrow + \text{degr2} \downarrow + \text{deact1} \oplus \]
\[ G_1[1] \leftrightarrow G_2[0] \leftrightarrow P_1[20] \leftrightarrow P_2[0] \]
\[ \text{observe(toggle_obs);} \]
\[ \text{infer(rouletteGibbs);} \]
\[ \theta_1 = \text{Gamma}(3, 5); //etc... \]

kineticLawOf expr1 : \( \theta_1 \times G1 \);
kineticLawOf expr2 : \( \theta_2 \times G2 \);
kineticLawOf degr1 : \( \theta_3 \times P1 \);
kineticLawOf degr2 : \( \theta_4 \times P2 \);

kineticLawOf activ1 : \( \theta_5 \times (1 - G1) \);
kineticLawOf activ2 : \( \theta_6 \times (1 - G2) \);
kineticLawOf deact1 : \( \theta_7 \times \exp(r \times P2) \times G1 \);
kineticLawOf deact2 : \( \theta_8 \times \exp(r \times P1) \times G2 \);

G1 = activ1 \uparrow + deact1 \downarrow + expr1 \oplus ;
G2 = activ2 \uparrow + deact2 \downarrow + expr2 \oplus ;
P1 = expr1 \uparrow + degr1 \downarrow + deact2 \oplus ;
P2 = expr2 \uparrow + degr2 \downarrow + deact1 \oplus 

G1[1] \leftrightarrow G2[0] \leftrightarrow P1[20] \leftrightarrow P2[0]

observe(toggle_obs);
infer(rouletteGibbs);
Experiment

- Simulated observations
- Gamma priors on all parameters (required by algorithm)
- Goal: learn posterior of 8 parameters
- 5000 samples taken using the Gibbs-like random truncation algorithm
Promoters
Observations used
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The ProPPA software framework contains a number of different inference engines, suitable for models with different characteristics, including infinite state space.

Integrating CTMCs with process algebras has brought a wide-range of fruitful application domains for stochastic process algebras.

ProPPA is a process algebra that incorporates uncertainty and observations directly in the model, influenced by probabilistic programming.

Syntax remains similar to Bio-PEPA whilst the semantics defined in terms of an extension of Constraint Markov Chains.

Observations can be either time-series or logical properties.

Embedding inference within a formal language, in the style of probabilistic programming brings many possibilities.
Challenges and Future Directions

- The value of observations

Can we reason about the "distance" between $\mu$ and $\mu^*$?
Challenges and Future Directions

- Heterogeneous populations

What if we are seeking the “optimal mix” rather than the best individual representative?
Thanks!

- **Bio-PEPA** was joint work with *Federica Ciocchetta*, funded by EPSRC.

- **ProPPA** was joint work with *Anastasis Georgoulas* and *Guido Sanguinetti*, funded by Microsoft Research, ERC and CEC FP7 FET Proactive programme FoCAS.