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Composing inference algorithms as program transformations

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Abstract

Probabilistic inference procedures are usually coded painstakingly from scratch, for each target model and each inference algorithm. We reduce this effort by generating inference procedures from models automatically. We make this code generation modular by decomposing inference algorithms into reusable program-to-program transformations. These transformations perform exact inference as well as generate probabilistic programs that compute expectations, densities, and MCMC samples. The resulting inference procedures are about as accurate and fast as other probabilistic programming systems on real-world problems.

1 INTRODUCTION

Writing inference algorithms for probabilistic models is tedious and error-prone. Conceptually, these algorithms are combinations of simpler operations, such as computing the density of a distribution at a given point. So it is unfortunate that these algorithms are traditionally implemented from scratch. In this paper, we show how to describe these building blocks in code, so that they need not be rewritten for every new inference algorithm or model.

We contribute the first method for composing multiple inference algorithms over the same model, even exact and approximate ones over the same factor. Our approach is to express inference in terms of operations that transform one probabilistic program into another. We use probabilistic programs to represent distributions, though our approach is compatible with other representations such as factor graphs. The goal of our transformations is to turn a program that denotes a model into another program that, when interpreted to draw a weighted sample, is equivalent to the desired inference algorithm.

Because the output of an inference transformation is still a probabilistic program, we can apply further inference transformations to the program. In this way, we can subject a single model to multiple inference methods without coding them from scratch. We thus reduce the informal problem of combining inference methods to the formal and more automatable problem of composing program transformations. In particular, approximate inference methods can be composed with taking advantage of exact mathematical equivalences such as conjugacy.

2 MOTIVATION AND RELATED WORK

Developing inference algorithms that work on a variety of models has long been a goal of probabilistic inference, including graphical models and probabilistic programming. The composability of inference algorithms has unfortunately lagged behind the composability of models.

Many probabilistic programming systems allow a choice of inference methods, both exact and approximate. For example, the probabilistic language Church (Goodman et al. 2008) has many interpreters, each of which implements a different inference method. Systems such as Figaro, Factorie, Anglican and Wolfe (Pfeffer 2009; McCallum et al. 2008; Wood et al. 2014; Riedel et al. 2014) also allow adding inference methods, as new code in the host language where the systems are embedded. However, the end result of applying these inference methods is behavior or code in a different language, no longer a probabilistic program. Thus, it is difficult in these systems to apply one method to the result of another method.

More similar to our approach is the approach of Ścibior et al.'s (2015; 2016). Like us, they express and compose inference methods as transformations that produce probabilistic programs in the same language. Thus for example they reuse Sequential Monte Carlo to implement Particle Independent MH. But because their probabilistic

language reuses many primitives from the host language Haskell, their transformations cannot inspect most of the input code, notably deterministic computations and code under a binding. In contrast, we can perform exact inference (Section 5), we can compute densities and conditional distributions (Section 4.3) in the face of deterministic dependencies, and we can generate MH samplers (Section 4.5) using a variety of proposal distributions.

So in general, transformations need to take programs as input as well as produce them as output in order to support the variety of inference composition found in the literature. To illustrate the need and the variety, below we recall some patterns of inference composition where we want to *reuse* existing implementations in ways unsupported by existing systems such as those named above.

Sometimes, we apply approximate inference to a model, then post-process the results using exact inference. For example, a popular way to perform inference for latent Dirichlet allocation (LDA) is to use Gibbs sampling (Griffiths and Steyvers 2004) to infer topic markers for each word, then infer from these topic markers the exact distribution on words given each topic.

Other times, we apply exact inference to parts of our model, and use an approximate method for the rest. As an example, Hughes et al. (2015) develop an inference algorithm for hierarchical Dirichlet processes that samples the truncation dictating the number of topics then performs variational inference for the other model parameters. This inference combination requires that sampling a truncation still leave in place a representation on which we can perform variational inference.

Another composition pattern emerges from recent work on parallelizing an inference algorithm to run on multiple machines (Neiswanger et al. 2014; Xu et al. 2014; Gelman et al. 2014). The pattern is to transform a posterior distribution for a parameter given the data into a model that lets us infer noisy versions of the parameter given subsets of the data. We then combine these noisy parameter estimates to infer the underlying parameter.

Finally, given a linear sequential model, we often want to predict future states of the system and the dynamics that govern them. Given the dynamics, for systems like Kalman filters, we may use exact inference to derive the state transition functions in closed form. Learning the dynamics, on the other hand, is usually treated as an approximate inference problem where we sample different possible dynamics given some observed states. This joint learning and exact inference again composes two inference algorithms. Section 6.1 shows our inference composition at work in a tiny instance of this case. We illustrate our approach using this example in Section 3.

3 EXAMPLE OF INFERENCE COMPOSITION

We illustrate our program-transformation approach to inference composition using a simple linear dynamical system. Our model below defines a joint distribution:

```
\begin{aligned} & \text{noise}_{T} & \sim \text{Uniform}(3,8) \\ & \text{noise}_{E} & \sim \text{Uniform}(1,4) \\ & x_{1} \mid \text{noise}_{T} & \sim \text{Normal}(0, \text{noise}_{T}) \\ & m_{1} \mid x_{1}, \text{noise}_{E} & \sim \text{Normal}(x_{1}, \text{noise}_{E}) \\ & x_{2} \mid x_{1}, \text{noise}_{T} & \sim \text{Normal}(x_{1}, \text{noise}_{T}) \\ & m_{2} \mid x_{2}, \text{noise}_{E} & \sim \text{Normal}(x_{2}, \text{noise}_{E}) \end{aligned}
```

We would like to draw samples from the posterior distribution over noise_T and noise_E given observations m_1 and m_2 , using a Metropolis-Hastings (MH) sampler.

We start by representing the model in our language:

The use of Dirac at the bottom shows that this distribution ranges over pairs of pairs of reals.

We first apply the *disintegration* transformation to get another program. As detailed in Section 4.3, disintegration takes as input a joint distribution and produces a program representing a family of posterior distributions. The new program is a function from the observations to the posterior distribution. For example, disintegrating kalman produces the program kalman2 below. It takes as input (m1, m2) and returns the distribution over (noiseT, noiseE) given those values for m1 and m2.

The use of Lam at the top and Weight at the bottom shows that this is a function from pairs of reals (m1, m2) to measures over pairs of reals (noiseT, noiseE).

The next step is to apply the *simplification* transformation to kalman2 to get kalman3.

```
kalman3 = Lam((m1,m2),
noiseT <~ Uniform(3, 8);
noiseE <~ Uniform(1, 4);
Weight(P, (noiseT, noiseE)))</pre>
```

This program is equivalent to kalman2, except the simplification transformation has symbolically integrated out the Normal-distributed random variables $\times 1$ and $\times 2$ and replaced them by an observation likelihood in closed form, which we elide above as P.

We next apply to kalman3 another program transformation we call mh, which implements MH sampling. The mh transformation takes as input two programs. The first program represents a proposal distribution, or more precisely a function from the current sample to a distribution over proposed samples. Here we use a proposal distribution that with equal probability resamples one of noiseT and noiseE while keeping the other fixed:

The second input to the mh transformation represents the target distribution. In this example, it is the part of kalman3 above after the top line Lam((m1, m2),. From these inputs, the mh transformation computes a symbolic formula for the MH acceptance ratio and embeds it in a program representing a transition kernel. The new program, kalman4 below, is a function from the current sample to a distribution over pairs of proposed samples and acceptance ratios:

The elided part A is a symbolic formula that computes the acceptance ratio using the current sample (noiseT, noiseE) and the sample proposed by the Superpose. This acceptance ratio can then be used to decide whether to accept or reject the proposed.

We then perform further optimizations on kalman4, including algebraic simplifications and rewriting the program to use fewer <~s. We describe in more detail the kinds of optimizations we perform in Section 5. The resulting program, kalman5, has the following structure:

```
kalman5 = Lam((noiseT, noiseE),

Superpose(

(1/2, n <\sim Uniform(3, 8);

Dirac(((n, noiseE), A_T))),

(1/2, n <\sim Uniform(1, 4);

Dirac(((noiseT, n), A_E)))))
```

The elided parts A_T and A_E are algebraically simplified formulas for the acceptance ratio in each of the two cases.

Finally we feed this last program kalman5 to a sampler (Algorithm 1). Given an observation and a current sample, this sampler produces a proposed sample and the MH acceptance ratio of that sample.

In the command above, (0,1) is the observation, and (5,2) is the current sample. In the output above, (5,1.6811397) is the proposed sample, and 0.7924639 is its acceptance ratio.

4 INFERENCE METHODS AS PROGRAM TRANSFORMATIONS

To compose inference methods, we pose them as transformations of one probabilistic program into another. We then achieve the desired inference method for the former program by applying a simpler inference method, such as exact inference or weighted sampling, to the latter program. For example, in Section 3 we feed a program to disintegration (Section 4.3), then mh (Section 4.5), then simplification (Section 5), and finally sampling. Only in the final sampling step is any random choice made!

We first define our probabilistic language, then describe various program transformations that work in concert.

4.1 LANGUAGE DESCRIPTION

Below is our core grammar of probabilistic programs:

```
e ::= x | 1 | e - e | e < e | exp (e) | If (e, e, e) | ...
| Sum (e, e, x, e) | Int (e, e, x, e)
| Lam (x, e) | App (e, e) | (e, e) | e [0] | e [1]
| Uniform (e, e) | Normal (e, e)
| Gamma (e, e) | Weight (e, e)
| Categorical ((e, e), ...)
| Superpose ((e, e), ...) | x < ~ e; e</pre>
```

The first line of this grammar says that our language includes ordinary programming support for variables, math, and If. The second line adds primitives to represent Summation and Integration, used in Section 4.2. The third line adds functions and tuples.

The remainder of the grammar is what makes our language probabilistic: we add primitives that represent and compose measures. To start with, Uniform (1, 2) represents the uniform distribution over real numbers between 1 and 2, and Normal (3, 4) represents the normal distribution with mean 3 and standard deviation 4.

Weight (1,8) represents the probability distribution that assigns its entire probability mass 1 to the single outcome 8. We write Dirac(8) as syntactic sugar for it. In contrast, Weight (0.7,8) represents the measure, or unnormalized distribution, that assigns the probability 0.7 to the single outcome 8. This primitive lets our language represent (unnormalized) measures in general, not just (normalized) probability distributions. This expressivity lets us separately reuse a transformation that produces an unnormalized measure (Section 4.3) and a transformation that subsequently normalizes a measure (Section 4.4). Also, Weight lets us represent a distribution by combining a base measure and a density function.

Categorical represents the categorical distribution with a sequence of zero or more pairs. The first element of each pair is the probability of selecting the outcome that is the second element of the pair. If the first elements of the pairs do not sum to 1, they are normalized.

Superpose is like Categorical, except it does not normalize, so it can represent measures in general. We can define Superpose in terms of Categorical and Weight, but it is actually more convenient to define Weight and Categorical in terms of Superpose.

The final primitive <~ (pronounced "bind") composes two distributions e1 and e2. The second distribution e2 may depend on the outcome x of e1. The outcome of the composed distribution x<~e1; e2 is the outcome of e2. This primitive lets our language represent sequential and hierarchical models. A simple example is this model:

```
x \sim \text{Uniform}(0,2) y|x \sim \text{Uniform}(x,3)
```

We write the marginal distribution over y as

```
x \ll Uniform(0, 2); Uniform(x, 3) and the joint distribution over (x, y) as x \ll Uniform(0, 2); y \ll Uniform(x, 3); Dirac((x, y))
```

To make the semantics of our language more concrete, Algorithm 1 shows a sampler that takes a probabilistic program as input and returns a draw from the distribution it represents. It is our only operation that calls a random number generator. We apply it last in a sequence of transformations to perform approximate inference.

Like a typical interpreter, Algorithm 1 takes as input not only a program but also an environment, which is a table

```
Algorithm 1: Weighted sampler: sample(m, env = [])
```

```
Input: program representing a measure: m Input: environment: env
```

Output: pair of values (outcome, weight)

Examine m

if m has the form Weight (w_1, e_1) then

Evaluate e_1 in the environment env, obtaining v_1 Return (v_1, w_1)

else if m has the form Normal (e_1, e_2) then

Evaluate e_1 in the environment env, obtaining v_1 Evaluate e_2 in the environment env, obtaining v_2 Sample from the normal distribution with mean v_1 and standard deviation v_2 , obtaining v_3 Return $(v_3, 1)$

else if m has the form $x < \sim m_1$; m_2 then

Call Algorithm 1 on m_1 with the environment *env*, obtaining (v_1, w_1)

Let env' be the environment env extended with x having the value v_1

Call Algorithm 1 on m_2 with the environment env', obtaining (v_2, w_2)

Return $(v_2, w_1 \cdot w_2)$

else

The other cases are similar to above

end

mapping variable names to values. Also, because our language includes unnormalized measures, this sampler returns not only a draw but also an importance weight.

4.2 EXPECTATION TRANSFORMATION

The rest of this section describes various inference transformations that we apply to our probabilistic programs. Because we implement some transformations in terms of others, we describe the transformations not in the order we apply them but in the order we implement them.

Our expectation transformation turns any program that represents a distribution into another program that represents its expected value. This transformation is exact and simple even though the expected values of many distributions have no closed form, because our language represents integrals symbolically with Int. For example, the expectation transformation turns the program x < Uniform(0, 2); Uniform(x, 3) into

The latter program represents the integral $\frac{1}{2-0}\int_0^2 \frac{1}{3-x}\int_x^3 y\,dy\,dx$. To compute this integral in closed form is to perform exact inference on the given distribution. The expectation transformation itself does not do so; nor does it approximate the integral by sampling.

```
Input: program representing a measure: m
Input: program representing a function: f
Output: program representing a number
Examine m
if m has the form Weight (w_1, e_1) then
| \text{ Return } w_1 \cdot \text{App}(f, e_1) 
else if m has the form Normal (e_1, e_2) then
| \text{ Return Int } (-\infty, \infty, \times, e_3 \cdot \text{App}(f, \times)) 
where the program e_3 computes the density of the
| \text{ Normal}(e_1, e_2) \text{ distribution at } \times
else if m has the form x < \sim m_1; m_2 then
```

Algorithm 2: Expectation transformation: expect(m, f)

Call Algorithm 2 with m_2 and f obtaining e_3 Call Algorithm 2 with m_1 and Lam (x, e_3)

else

The other cases are similar to above **end**

Specified more generally, the expectation transformation turns any program that represents a measure, along with a function from the sample space to numbers, into another program that represents the integral of the given function with respect to the given measure. We show this transformation as Algorithm 2. It handles primitive distributions such as Normal by looking up their density from a table.

4.3 DENSITY AND DISINTEGRATION

Turning a distribution into its density function is naturally expressed as a program transformation (Bhat et al. 2012, 2013). More precisely, the density transformation takes as input a probabilistic program representing a distribution, and returns another program representing a function that maps each point in the sample space to the density at that point. Note that this transformation does not compute any density numerically. It only returns a program that computes densities when interpreted by our weighted sampler (Algorithm 1). For example, the density transformation turns the probabilistic program

```
x <~ Uniform(0, 2);
y <~ Uniform(x, 3); Dirac((x, y))
into the density function Lam((x,y), If(0<x<2,
If(x<y<3, 1/(3-x), 0)/(2-0), 0)).</pre>
```

We implement density in terms of another program transformation, *disintegration* (Shan and Ramsey 2017; Narayanan and Shan 2017). Disintegration is similar to conditioning in that it takes a probabilistic program representing a joint distribution $\Pr(X,Y)$ as input, but instead of returning a conditional distribution $\Pr(Y \mid X = x)$, disintegration returns an unnormalized slice $\Pr(Y, X = x)$ of the original distribution. More precisely, disintegration returns a program representing a

```
Algorithm 3: Density transformation: density(m, t)
```

```
Input: program representing a measure: m
Input: program representing value drawn from m: t
Output: program representing a nonnegative number
1. Disintegrate x<~m; Dirac((x,Unit)), obtaining e1</li>
2. Call Algorithm 2 on App (e1,t) and Lam(y,1)
```

```
Algorithm 4: Observation transformation: observe(m, t)
```

```
Input: program representing a measure: m
Input: program representing value drawn from m: t
Output: program representing a measure
Examine m
if m has the form Uniform(e_1,e_2) or

Normal (e_1,e_2) or Gamma(e_1,e_2) then
Let d be a program that computes the density of the distribution m
Return Weight (App (d,t), t)
else if m has the form x < m_1; m_2 then
Call Algorithm 4 recursively with m_2 and t
obtaining m_3
Return x < m_1; m_3
else
Raise an error about not being able to handle m
```

function from values of x to measures Pr(Y, X = x). Such a (measurable) function is also known as a *kernel*.

end

Taking advantage of the fact that disintegration does not normalize the measures it returns, we implement the density transformation in terms of disintegration and expectation. This implementation is shown in Algorithm 3. It invokes disintegration (letting Y be the space that consists of a single point Unit) then expectation (letting the integrand f be the function that maps Unit to 1).

Disintegration is useful independently of the density transformation. For example, Section 3 uses it to turn the prior kalman into the posterior kalman2.

We sketch how disintegration works in terms of a simpler program transformation, which we call *observation* (Algorithm 4). This transformation takes as input a measure m and a value t that could have been drawn from m, and returns a measure which only returns t, weighted by how likely that value was to be drawn from m. For example, the observation transformation turns the program

```
x <~ Uniform(0, 2); Uniform(x, 3)
and the variable y into the program
x <~ Uniform(0, 2);
Weight(If(x<y<3, 1/(3-x), 0), y)</pre>
```

Algorithm 5: Normalization transformation: normalize(m)

Input: program representing a measure: m

Output: program representing a probability distribution

- 1. Call Algorithm 2 on m and Lam (x, 1) obtaining the program e_1
- 2. Return x < m; Weight $(1/e_1, x)$

As indicated at the bottom in Algorithm 4, the observation transformation only handles a subset of our language. In particular, it does not handle Dirac, so it does not handle the typical program kalman in Section 3. In general, if the input program performs arithmetic or any other deterministic computation to produce the observation t, then we need to invert this deterministic computation and insert any Jacobian factors required. This inversion is what the disintegration provides over observation.

To relate observation and disintegration more precisely, suppose the program m represents a measure over X, the program e represents a value in Y, and observation turns e and e into e in

4.4 NORMALIZATION AND CONDITIONING

The presence of Weight in our language enables the observation and disintegration transformations to return measures that are typically unnormalized. To recover a probability distribution, we must reweight the measure. We define this *normalization* operation as a program transformation as well, shown as Algorithm 5.

Conditioning can now be defined by composition: it is just disintegration, followed by normalizing the measure.

4.5 MCMC SAMPLING TRANSFORMATIONS

A major contribution of this paper is to implement Markov chain Monte Carlo (MCMC) methods, such as MH sampling and Gibbs sampling, in a way that applies to a variety of target distributions and composes with other inference techniques. We express an MCMC method as a transformation from a program representing the target distribution to a program representing the transition kernel. Whereas the transformation itself makes no random choices, the latter program can be interpreted by our weighted sampler (Algorithm 1) to generate a random chain, or subject to simplification (Section 5).

Following this approach, our MCMC implementations closely resemble their textbook presentation. As shown in Algorithm 6, where the textbook presentation of the acceptance ratio refers to the target and proposal densi-

Algorithm 6: Metropolis-Hastings sampling transformation: mh(*proposal*, *target*)

Input: program representing the proposal kernel: *proposal*

Input: program representing the target distribution: *target*

Output: program representing MCMC transition kernel with acceptance ratio

- 1. Let old and new be fresh variable names
- 2. Call Algorithm 3 on *target* and old, obtaining p_{old}
- 3. Call Algorithm 3 on *target* and new, obtaining p_{new}
- 4. Call Algorithm 3 on App (proposal, new) and old, obtaining $q_{\rm old;new}$
- 5. Call Algorithm 3 on App (proposal, old) and new, obtaining $q_{new:old}$
- 6. Let e_1 be $(p_{\text{new}} \cdot q_{\text{old;new}})/(p_{\text{old}} \cdot q_{\text{new;old}})$
- 7. Return Lam(old, new<~App(proposal,old);
 Dirac((new, e₁)))

 Algorithm
 7:
 Gibbs sampling transformation:

 gibbs(target)

Input: program representing the n-dimensional target distribution: target

Output: program representing MCMC transition kernel Let \mathbf{x} be the set of the n variables in the *target* Initialize *choices* to the empty sequence []

For each $x_i \in \mathbf{x}$:

- 1. Let x_{-i} be the rest of the variables
- 2. Let e_1 be $\mathbf{x} < \sim target$; Dirac((x_{-i}, x_i))
- 3. Disintegrate e_1 , obtaining e_2
- 4. Let e_3 be App (e_2, x_{-i})
- 5. Call Algorithm 5 on e_3 , obtaining e_4
- 6. Let y be x except replacing x_i by new
- 7. Let e_5 be new<~ e_4 ; Dirac(\mathbf{y})
- 8. Add the pair $(1/n, e_5)$ to *choices*

Return Lam(x, Superpose(choices))

ties, our implementation invokes the density transformation (Algorithm 3) on two probabilistic programs, representing the target and proposal distributions. Using the fact that the density transformation symbolically handles free variables such as old and new, we perform the transformation just once (not once per sampler iteration) to generate a program that takes the current state as input.

Gibbs sampling is a special case of MH, where the proposal kernel combines the results of conditioning the target distribution along each dimension. The acceptance ratio is then always 1, so it need not be computed. To produce such a proposal kernel automatically, we implement Gibbs sampling as a separate transformation, Algorithm 7. The input is a program representing an n-di-

mensional distribution $\Pr(x_1, \ldots, x_n)$. For each random variable x_i , we condition (Section 4.4) on the other variables x_{-i} to get a program that resamples x_i . We then combine these n programs to form the proposal kernel.

5 SIMPLIFICATION

Because we express each inference technique as a transformation that produces a probabilistic program, rather than as an interpreter that makes immediate random choices, we can optimize and simplify the produced programs. To this end, we apply the optimizations discussed by Carette and Shan (2016). This *simplification* transformation does not change the measure represented by a program but tries to place the program in a form that, when interpreted by our weighted sampler (Algorithm 1), draws samples faster and with more uniform weights.

Based on computer algebra, the simplification transformation recognizes conjugacy relationships, integrates out latent variables, and performs algebraic simplifications. Like other transformations, simplification operates on a program before any variables receive their values, so in particular its efficacy is independent of data sizes. The rest of this section briefly describes these optimizations.

Conjugacy relationships Simplification recognizes when a density represented by Weight matches the density of a primitive distribution. A simple example arises from the joint distribution $\Pr(Y,X)$ represented below:

```
x <~ Normal(a, s);
y <~ Normal(x, t); Dirac((y, x))</pre>
```

Disintegrating this program (Section 4.3) produces

This latter program scales the measure Normal (a,s) with the density $\exp(\ldots)/t/\operatorname{sqrt}(2*\operatorname{pi})$ to represent the conditional distribution $\Pr(X\mid Y)$ up to a constant factor. Normalizing and simplifying it yields

```
Normal( (y*s^2+a*t^2)/(s^2+t^2), s*t/sqrt(s^2+t^2))
```

using the conjugacy relationship between Normals (assuming s and t are positive). This simplified program runs faster; it draws samples without weighting them.

This optimization is symbolic, in the sense that it works even when the initial program contains free variables such as a, s, and t, whose values are unknown.

This optimization is robust because it recognizes not words like Normal but the densities they denote. Thus it works even if we express Normal (0,1) by spelling

out its density, whether we expand the polynomial $-(y-x)^2$. All conjugacies among Normal, Gamma, and Beta thus fall out from recognizing their densities.

Integrating out a variable When a distribution is described using a latent random variable, it usually helps to eliminate the variable. Such latent variables include $\times 1$ and $\times 2$ in Section 3, as well as \times in

```
x <\sim Normal(0, 1); Normal(x, 1)
```

The simplification transformation eliminates these variables. In particular, it integrates out continuous latent variables symbolically. The density-recognition machinery just described then produces simpler, faster, and equivalent programs, such as Normal (0, sqrt (2)).

This integration is symbolic, again in the sense that it works even when the initial program contains free variables whose values are unknown. For example, the program x < Normal(a, s); Normal(x, t) simplifies to $Normal(a, sqrt(s^2+t^2))$.

Algebraic simplifications When we produce a program that calculates acceptance ratios, the numerator and denominator share many factors, which are usually canceled out by hand. The simplification transformation automates this optimization using computer algebra, so an expression like (a*b)/(a*c) becomes b/c.

6 EXPERIMENTAL RESULTS

To demonstrate that our approach is modular and practical, we apply multiple inference methods (MH, Gibbs) to a variety of models. We conduct three experiments using the Hakaru system (Narayanan et al. 2016).

Modular means we can re-use the components in Section 4 and Section 5 to produce all three samplers. In each experiment, a pipeline composed of reusable inference transformations turns a concise generative model into an executable MCMC sampler in seconds.

Practical means our approach can solve real-world problems by expressing popular models and inference methods discussed in the literature. The largest of our three experiments is the third, a document classification task using the 20 Newsgroups corpus.

We measure the accuracy and speed of our automatically generated samplers, showing they are in line with solutions from commonly used probabilistic programming languages. Our samplers are more accurate across the board because simplification eliminates all latent continuous variables, regardless of the dimensionality of the problem (that is, input and output array sizes).

Table 1: MH sampler run times for linear dynamics

Inference method	Run time (msecs)	
	Mean	SD
WebPPL	1078	16
Hakaru without simplificati	ons 1321	93
Hakaru with simplifications	269	10
Handwritten	207	4

Table 2: MH sampler ESS rates for linear dynamics

Inference method	ESS per sample	
	noise_T	noise_E
WebPPL	0.03	0.01
Hakaru	0.09	0.34

All measurements were produced on a quad-core Intel i5-2540M processor running 64-bit Ubuntu 16.04. Our samplers use Glasgow Haskell Compiler 8.0.1 -02.

6.1 MH SAMPLING FOR DYNAMICS

In our first experiment, we use MH to sample the random parameters of the linear dynamical system in Section 3. We compare our generated samplers with one produced by WebPPL, a state-of-the-art probabilistic programming system, and with one written by hand. The WebPPL sampler was compiled to JavaScript using Node 0.10.28.

Table 1 shows that our system generates a fast sampler, measured by using each sampler to draw 20,000 samples 10 times. Thanks to the simplifications that turn kalman4 into kalman5, the Hakaru sampler is 4 times as fast as the WebPPL sampler for the conditional distribution kalman2. (These times exclude the few seconds each system takes to compile the model into a sampler.)

Table 2 shows that our samplers generate good samples, quantified by the Effective Sample Size (ESS). Our ESS is higher per sample compared to WebPPL, because latent variables have been integrated out in kalman3.

6.2 GIBBS SAMPLING FOR CLASSIFICATION

In our second and third experiments, we generate Gibbs samplers and compare them to JAGS (v4.20), a probabilistic programming system widely considered practical for Gibbs sampling. We measure accuracy by how well the samplers recover true classifications, and speed by the time it takes to produce samples. This time consists of *initialization* time and time spent actually sampling. Initialization time is the time a system takes from receiving the model to generating the first sample: for JAGS to

load the model into memory, and for Hakaru to simplify the model and compile the result into machine code.

Our second experiment is to classify synthetic data using a Gaussian mixture model that has 3 components.

Figure 1 shows that Hakaru requires fewer sweeps than JAGS to achieve the same accuracy. Each curve plots the accuracy of one chain over the course of 15 sweeps on 250 data points. After just one sweep, all 20 Hakaru chains are >50% accurate, unlike the 20 JAGS chains, which take a few sweeps to catch up. The cause is that Hakaru's simplification transformation recovers a collapsed Gibbs sampler that computes the sample mean and variance of each mixture component in closed form.

Figure 2 shows Hakaru is about one order of magnitude slower than JAGS, measured by how long 6 sweeps take, varying data size from 500 to 2500 points. The top two curves represent two samplers generated by Hakaru with different lower-level optimizations: the second-from-top curve adds a *histogram* optimization to compute summary statistics such as the per-mixture-component sum $\sum_{i=1}^{N} \left\{ \begin{smallmatrix} t_i & \text{if } z_i = z^* \\ 0 & \text{otherwise} \end{smallmatrix} \right.$ in a single pass over the data for all components z^* . The bottom two curves show the run time of JAGS, with and without initialization. JAGS's speed advantage can be explained by Hakaru's current inability to reuse computation between updates during a sweep. Still, Hakaru is practical for this real-world task.

In our third experiment, Hakaru generates a classifier for the 20 Newsgroups corpus that is more accurate than JAGS and comparable in speed. We use the same Multinomial Naive Bayes model and 20 Newsgroups corpus as McCallum and Nigam (1998). We hold out 10% of the labels and use Gibbs sampling to infer them. We evaluate the samplers on data sizes ranging from 200 to all 19997 documents, evenly distributed among newsgroups.

Because the sampler generated by Hakaru is collapsed, it is more accurate than JAGS in two ways. First, Figure 3 shows Hakaru achieves better accuracy than JAGS after one sweep, and continues to for at least 1000 sweeps. Each curve there plots the accuracy (moving average with window size 20) of one chain on 400 documents. Second, Figure 4 shows Hakaru more accurate than JAGS across data sizes. We use 2 sweeps there since JAGS does not perform above chance with only 1 sweep.

Figure 5 shows Hakaru is as fast as JAGS, measured by how long 2 sweeps take, varying data size. JAGS's initialization time grows with the data size, while Hakaru's is constant. Whereas JAGS unrolls loops into a pointerbased stochastic graph whose size grows with the data, Hakaru generates tight loops over unboxed arrays irrespective of the data size. Even disregarding initialization time, JAGS is at best 4 times faster than Hakaru.

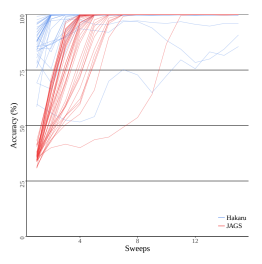


Figure 1: Gibbs sampler accuracy for Gaussian mixture

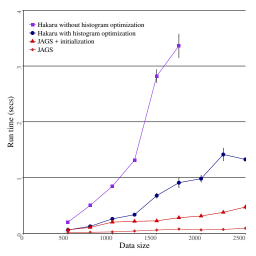


Figure 2: Gibbs sampler run times for Gaussian mixture

7 CONCLUSIONS

We express inference methods by composing program transformations such as disintegration and expectation. The resulting modular inference procedures perform comparably to other probabilistic programming systems and are usable for practical problems. This technique makes it easier and faster to create and test inference procedures and to explore novel inference methods.

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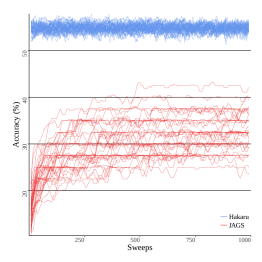


Figure 3: Document classification accuracy by sweeps

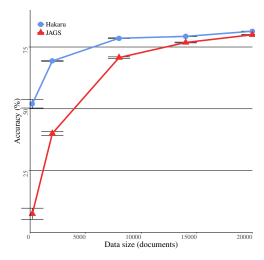


Figure 4: Document classification accuracy by data size

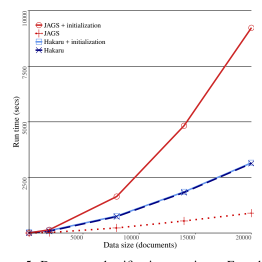


Figure 5: Document classification run times. Error bars were too small to display.

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Scaling Probabilistic Programming to Structured Spaces

1 Introduction

In this report, we introduce and review the necessary background material around probabilistic programming, the existing literature for performing inference in the models expressible with probabilistic programming languages, and propose mechanisms extending existing probabilistic programming systems so that models are easier to debug and may then be applied to more challenging probabilistic modelling tasks.

2 Literature Review

2.1 Probabilistic Programming

Probabilistic programming systems [Gordon et al., 2014] are specialised systems for specifying probabilistic models and efficiently performing inference with them. These models are usually expressed in a domain specific language (DSL) called a probabilistic programming language (PPL). These restricted languages are more amendable to program analysis and offer the potential of more easily generating efficient inference code for a particular model.

2.1.1 Taxonomy of Probabilistic Programming systems

Probabilistic programming languages can be partitioned into two designations: *first-order* PPLs and *higher-order* PPLs. First-order PPLs are ones where we can statically determine the number of latent variables of the model without performing any inference. The most popular systems are first-order as these restrictions make it possible to apply efficient inference algorithms to all models expressible

```
\begin{split} \langle e \rangle &:= \langle x \rangle \, | \, 1 \, | \, \langle e \rangle - \langle e \rangle \, | \, \langle e \rangle < \langle e \rangle \, | \, \exp(\langle e \rangle) \, | \, \operatorname{If} \left(\langle e \rangle_{}, \langle e \rangle_{}, \langle e \rangle_{}\right) \, | \, \ldots \\ &| \quad \operatorname{Sum}(\langle e \rangle_{}, \langle e \rangle_{}, \langle x \rangle_{}, \langle e \rangle_{}) \, | \, \operatorname{Int}(\langle e \rangle_{}, \langle e \rangle_{}, \langle x \rangle_{}, \langle e \rangle_{}) \\ &| \quad \operatorname{Lam}(\langle x \rangle_{}, \langle e \rangle_{}) \, | \, \operatorname{App}(\langle e \rangle_{}, \langle e \rangle_{}) \, | \, \langle e \rangle_{}, \langle e \rangle_{}) \, | \, \langle e \rangle_{} \, [0] \, | \, \langle e \rangle_{} \, [1] \\ &| \quad \operatorname{Uniform}(\langle e \rangle_{}, \langle e \rangle_{}) \, | \, \operatorname{Normal}(\langle e \rangle_{}, \langle e \rangle_{}) \\ &| \quad \operatorname{Gamma}(\langle e \rangle_{}, \langle e \rangle_{}) \, | \, \operatorname{Weight}(\langle e \rangle_{}, \langle e \rangle_{}) \\ &| \quad \operatorname{Categorical}((\langle e \rangle_{}, \langle e \rangle_{}), \, \ldots_{}) \\ &| \quad \operatorname{Superpose}((\langle e \rangle_{}, \langle e \rangle_{}), \, \ldots_{}) \, | \, \langle x \rangle < \sim \langle e \rangle_{}; \, \langle e \rangle_{} \end{split}
```

Figure 1: Grammar for the core of a probabilistic programming language

within them. First-order PPLs include BUGS [Lunn et al., 2000], Infer.NET [Minka et al., 2014], Stan [Carpenter et al., 2015], PyMC [Salvatier et al., 2016], as well as Edward [Tran et al., 2016].

Higher-order PPLs are languages which can express all the probabilistic models that first-order PPLs are unable to. This greater flexibility in specification comes with a greater challenge in devising generic inference algorithms that can be made to work in this larger model family. In the literature higher-order PPLs are sometimes referred to as universal probabilistic programming languages [Le et al., 2017, Yang et al., 2014] or languages with recursion.

It is deceptive to state that the defining characteristic of higher-order PPLs is that the languages have recursion as the recursion is only a problem if within recursion we make draws from a distribution. A fully deterministic recursive function is perfectly acceptable to use in a first-order PPL. This is indeed the case in the Stan language.

When our model has an unfixed number of latent variables, this introduces the additional challenge that we need to be able generate unique names for them as we sample from our model. This process of generating names is called an *addressing scheme*.

The addressing scheme is important for implementing MCMC algorithms where we need to maintain which of the latent variables in our proposed state are shared with variables in our current state.

2.1.2 Core language grammar

To keep things concrete we introduce a small language HAKARU. We define this grammar we more formally in Figure 1.

2.1.3 Importance sampler

The simplest inference algorithm to implement for higher-order PPLs is an importance sampling algorithm. This algorithm is also called likelihood-weighting.

This inference algorithm is first implemented in the BLOG probabilistic programming system [Milch et al., 2005].

2.1.4 Trace MH interpreter

Up until recently, the most common inference algorithm which came with probabilistic programming systems is a single-site Metropolis Hastings called Trace MH [Ścibior et al., 2018, Wingate et al., 2011]. In this algorithm, we generate run the program instrumented to keep track of all random variables we encountered. Then we re-execute the program in such a way that all but one of the random variables is kept fixed. This has the possibility of creating and removing random variables based on changes in control-flow.

2.1.5 SMC interpreter

Another popular to implement inference algorithm for probabilistic programming systems is the SMC interpreter [Wood et al., 2014]. In this setting, we generate a large population of particles and as we execute the trace we re-weight them based on which of the random variables we encounter are observed.

2.1.6 Variational and Amortised Inference

Variational inference encompasses a family of inference techniques centred around posing inference problems as optimisation problems. The idea being that we can construct a parameterised family of approximate distributions $q(\mathbf{x}; \phi)$ and then finding the ϕ closest to probability distribution we actually care about. The notion of closest is often some f-divergence and mostly commonly the reverse KL divergence.

$$D_{\mathrm{KL}}(q(\mathbf{x}; \phi) || p(\mathbf{x})) := \mathbb{E}_{q(\mathbf{x}; \phi)} \left[\log q(\mathbf{x}) - \log p(\mathbf{x}; \phi) \right] \tag{1}$$

This KL divergence can be then be rewritten as:

$$D_{\mathrm{KL}}(q(\mathbf{x} \mid \mathbf{y}; \phi || p(\mathbf{x} \mid \mathbf{y})) = \mathbb{E}_{q(\mathbf{x} \mid \mathbf{y}; \phi)}[\log p(\mathbf{x}, \mathbf{y}) - \log q(\mathbf{x} \mid \mathbf{y}; \phi)]$$
(2)

This is called the Evidence Lower Bound (ELBO).

The idea with amortised inference [Gershman and Goodman, 2014] is that we can share parameters ϕ for different choices of \mathbf{x} .

2.1.7 Variational Autoencoders

We can further generalise the amortised inference regime to allow us to jointly learn the model from the data as well the approximation of the posterior for the model.

2.2 Compiled Inference

Stochastic variational inference methods assume taking draws from an approximate distribution is easier than taking draws from the true distribution. But if the joint distribution of the true distribution is easy to draw samples from, there is an formulation of the KL divergence that we may use for amortised inference.

$$\mathcal{L}(\phi) = \mathbb{E}_{p(\mathbf{y})} [D_{\text{KL}} (p(\mathbf{x} \mid \mathbf{y}) || q(\mathbf{x} \mid \mathbf{y}; \phi)))]$$

$$= \mathbb{E}_{p(\mathbf{x}, \mathbf{y})} [-\log q(\mathbf{x} \mid \mathbf{y}; \phi)] + \text{const}$$
(3)

2.3 Implicit Models

For many of the models we can express in a probabilistic programming system, we are unable to efficiently compute the density of a trace associated with our model. For example, in large physics and biological simulators it can be computationally intractable to compute a density.

2.4 Model criticism

As part of defining and performing inference on our probabilistic models, we often want to evaluate our models. If our models when fitted can be used to make predictions, we can evaluate their accuracy on held-out labelled data, but this is inappropriate for many settings. More generally, if we have unlabelled data we may evaluate the test likelihood of it under our proposed model.

2.4.1 Two Sample Kernel Tests

Another approach, is to suppose we have our data Y and data generated after fitting Y'. We can ask if Y and Y' appear to come from the same distribution.

We can pose this statistic as a maximum mean discrepancy [Gretton et al., 2012] (MMD).

3 Proposal

The common thread that underlies this project is to push probabilistic programming to be applied in new settings that can take advantage of these new scalable inference algorithms as well as make the systems for usable for non-experts.

3.1 Debugging Probabilistic Programming

Often probabilistic modeling is an iterative process where a user tries out a model, evaluates it in some way, and then proposes some improvement. We can aid in the process by offering suggestions

automatically for ways to improve a probabilistic model.

The main idea is to define a neighbourhood $\mathcal{N}(p)$ which are the set of programs we can get to from our current program p by a set of known pre-defined rewrites.

An example of some possible rewrites could be:

- Replace $e_1 \sim \text{Normal}(e_2, e_3)$ with $e_1 \sim \text{Laplace}(e_2, e_3)$
- Replace $e_1 \sim e_2$ with $e_1 \sim \text{Mix}(e_2, \delta(0))$ where Mix is a mixture distribution

Each of these proposed programs can then be fitted and compared with the original program using a two-sample kernel test. We can then present the choices to the user interactively through some UI element in the PPS.

This work is most similar to work by Grosse et al. [2012]. The two main differences between this work and theirs is they restrict their search to a smaller space around a context-free grammar designed to mimic how humans heuristically choose to augment their probabilistic model, which means it can not discover. Additionally, their goal is to greedily discover the best structure while, we are more interesting in this as a debugging and diagnostic tool. In that sense, we are not doing a greedy search through the space of probabilistic programs as much as a quick exploration of the local neighbourhood of programs around the one the user has provided.

We mitigate the risks of this project by first restricting ourselves to a small subset of possible changes to the probabilistic model. This makes it more likely that we will find changes which improve the model while spending less computational time exploring the space of programs. I will further collaborate with my other supervisor Dino Sejinovic, who is an expert in using kernel methods like MMD for hypothesis testing.

3.2 Data Cleaning

While most research focuses on the model fitting and evaluation steps in data analysis, most of the time of practitioners is spent on acquiring and cleaning the datasets. Data cleaning can describe writing transformations to get data into a proper schema, correcting annotation errors, removing duplicates, as well correcting other internal inconsistencies. This project focuses on data which is already

in a tabular format, is likely to mostly fit into a schema unknown in advance, but may still have errors and misspellings that need to be corrected.

The approach we take is defining a generative model for tabular data where we define a prior on data schema, and a generative model for generating cells of data conditioned on this schema as well as previously generated cells.

We mitigate the risks of this project by initially restricting ourselves to synthetically-generated tabular datasets as well previous real-world noisy tabular data where a prior analysis has given us some ground truth to work with.

3.3 Automated Expectation Maximisation

Probabilistic programming systems can also be extended to do more than sample from a conditional distribution.

We mitigate the risk of this project in starting with a restricted model family of conditionally conjugate models where we know the exact rewrites our computer algebra system will need to do.

3.4 Proposed Schedule of Work

The timeline we are proposing for this study consists of submitting the preliminary work on debugging probabilistic programs and Bayesian metalearning to an appropriate NIPS workshop. If we are exceedingly lucky in obtaining results we will submit this work to AISTATS on October 4th. Otherwise, it is more likely that this work will be submitted to UAI in March. The data cleaning project is intended to be submitted to the KDD conference.

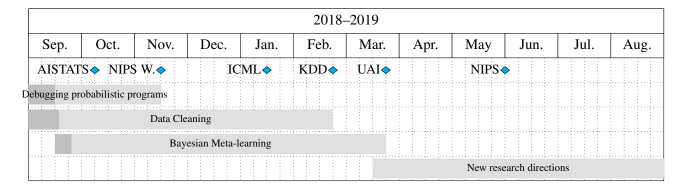


Figure 2: Proposed plan for the upcoming year. Conference deadlines are marked on the first row

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