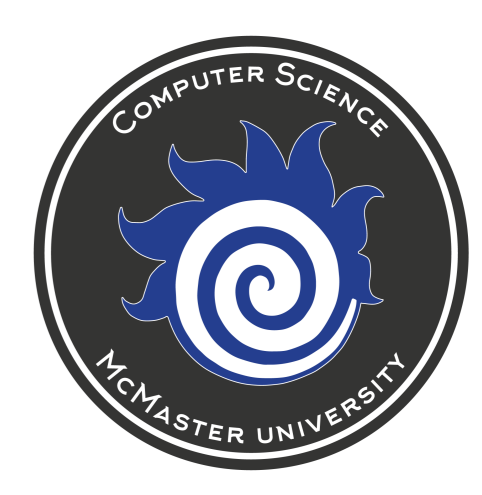


# Hakaru Language: Standard Library Implementation and Language Validation Testing

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## Introduction

Real world phenomena often behave according to a probability distribution. By letting us sample from distributions, Hakaru enables us to model real world phenomena that include uncertainty. Hakaru can then perform transformations on these models to turn them into conditional probabilities that let us make inferences based on a priori knowledge. This, for example, is useful in machine learning applications where the goal is often to predict or infer information about the model based on the past outcomes.

#### Mathematical Background

The mathematical functions that we use to describe these probabilities are called probability distributions and the quantity that denotes the outcome of the experiment is called a random variable. A random variable can take on continuous or discrete values depending on the application. Continuous random variables are formally defined by a probability density function (PDF), which maps the outcomes of the random variable to real numbers that represent a probability. Similarly, a discrete random variable is described by a probability mass function (PMF). As an example, let us consider one of the most ubiquitous and naturally occurring distributions, the normal distribution. We can say that a random variable, X, is normally distributed with the following notation.



Here, we can see that the normal distribution is parametrized by  $\mu$ , the population mean, and  $\sigma$ , the standard deviation. A plot the PDF is shown below. Remember that the PDF describes the liklihood that a value sampled from this distribution will equal x.

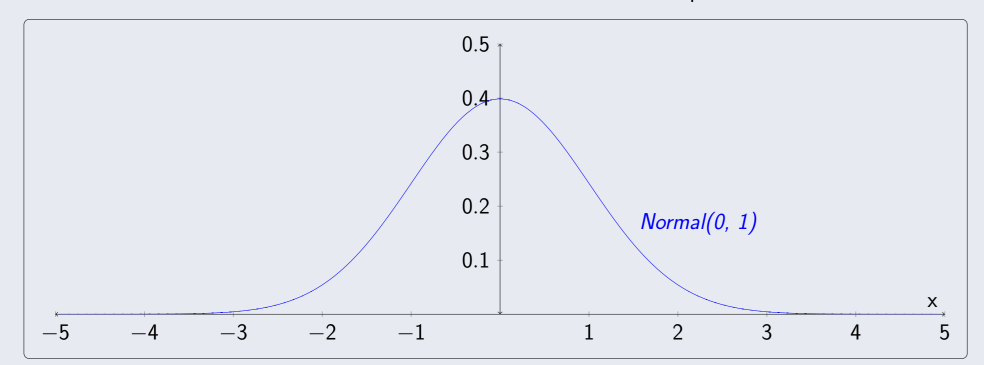


Figure 1: PDF of the normal distribution  $^{[1]}$ , with  $\mu=0$  and  $\sigma=1$ 

#### Language Guide

Hakaru is an experimental **probabilistic programming language**, which aims to simplify the way users implement statistical distributions.

- Niche application means that Hakaru is a small language with limited features.
- Running a hakaru program generates a stream of random numbers distributed according the statistical distribution modeled.
- Hakaru code can be compiled to C and Haskell so that models can be exported for use in larger applications.
- Terminology: implementation of statistical distribution  $\iff$  probabilistic model  $\iff$  model.

'hk-maple' is a provided inference algorithm that uses Maple to perform algebraic transformations on hakaru programs. In its default mode (Simplify), it returns an equivalent hakaru program with greater sampling efficiency. In applications requiring billions of samples (e.g. machine learning) this has the potential to save significant processing time!

# Key Concepts

- $\star$  Hakaru program  $\iff$  implementation of statistical distribution  $\iff$  probabalistic model  $\iff$  model.
- ★ Values pulled from a distribution are called measures. Measures ⇔ samples.
- $\star$  ' $\sim$ ' vs ' $<\sim$ ':
  - $X \sim Normal(0,1)$  means that the random variable, X, is distributed according to the standard normal distribution (statistics literature). -  $x <\sim normal(0,1)$  means to pull a random variable from the standard normal distribution and bind it to the variable, x (Hakaru code).
- \* PDF: Probability Density Function (continuous random variables).
- \* PMF: Probability Mass Function (discrete random variables).
- \* UDR chart: Univariate Distribution Relationship chart (describes relationships and transformations between different distributions).

## Motivation

### **Objectives**

- Increase language accessibility: add new language features such as primivite mathematical functions (e.g. log, choose), new distributions, error handling, etc.
- Test language validity: use relationships between distributions to test the validity of program transformations like 'hk-maple'.

#### **Endeavours**

- Standard library development: the UDR is used to branch out and implement new distributions.
- Syntax-highlighting-for-hakaru package for Sublime Text: this package makes developing Hakaru code much more convenient and streamlined (Mahmoud).
- Test case writing: we focus mainly on comprehensively testing only a select few distributions (e.g. chi-square).

## Standard Library Development

In order to make Hakaru more accessible to users, the standard library should implement commonly used probabilistic models. To this this end, we have implemented the distributions described in the UDR, a portion of which can be seen below. In general, we have used the following principles to guide the development of the standard library:

- Whenever possible, implement distributions as transformations on pre-existing models.
- Reference the UDR chart for possible ways to implement each distribution.
- In the case of multiple possible implementations, take the shortest possible path from a primitive distribution on the UDR.

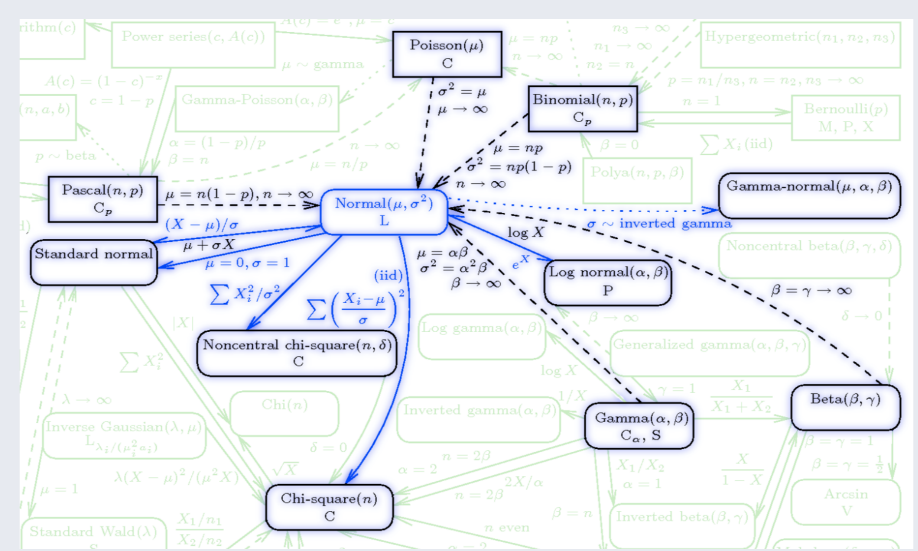


Figure 2: A snapshot of the UDR [1] shows how the normal distribution can be transformed into a multitude of other distributions.

Unfortunately, Hakaru does not allow us to do algebraic operations on the PDF of a given distribution. The only operator we can use on a model is bind ( $<\sim$ ), to pull a sample from it. However, we are able to transform samples however we like. Therefore, we are interested in implementing transformations of the following form.

 $R(p,q) \Rightarrow X \sim A(p) \Rightarrow f(X) \sim B(q)$ 

In the equation above, p parameterizes the distribution A, q parameterizes the distribution B, R(p, q) is a set of relationships between p and q that must be satisfied, and f is a function that applies a transformation to a sample. We can expand this definition to include transformations defined in terms of an aggregation of multiple independent samples. For example, the standard chi-square distribution is defined as the sum of the squares of n normal random variables (see Figure 3).

Hakaru also lends itself very well to Bayesian transformations where we want to pull a sample from a distribution which is parameterized with a value sampled from another distribution. These transformations take the following form. Here, we are saying that if X is parameterized by p, then the distribution C, which is parameterized by p and p0, is equal to another distribution, p1, which is parameterized by p2 and p3. The gamma-poisson distribution can be described by such a transformation (see Figure 4).

 $X \sim A(p) \Rightarrow Y \sim B(q,X) = C(p,q)$ 

def chiSq(means array(real), stdevs array(prob) ):
 q <~ plate \_ of size(means): normal(means[\_], stdevs[\_])
 return summate i from 0 to size(q):
 ((q[i]-means[i])/stdevs[i])^2</pre>

Figure 3: Our implementation of the chi-square distribution.

def gammaPoisson(shape prob, scale prob) measure(nat):
 mu <~ gamma(shape, scale)
 X <~ poisson(mu)
 return X</pre>

Figure 4: Our implementation of the gamma-poisson distribution.

Some distributions on the UDR are unreachable from Hakaru's primitive distributions using transformations like the 2 described above. In these cases, we have to implement the model in terms of the PMF for a discrete distribution or in terms of the PDF for a continuous distribution.

# Testing Relationships Between Distributions

Recall that when developing the standard library, the UDR chart often implied multiple possible implementations for a given distribution. Naturally, we would expect alternative implementations for the same distribution to result in equivalent probabilistic models. This line of thought is the basis for the kinds of test cases we have implemented. More generally, we have the following hypothesis.

Assume we know a proven relationship between 2 statistical distributions, A and B, which allows us to transform A and B into distributions that are equivalent to each other. Further assume this transformation takes on one of the forms discussed in the Standard Library Development section. We hypothesize that by applying the appropriate transformations to implementations of A and B, we can create two Hakaru programs whose 'hk-maple' outputs will be equivalent to each other. Test cases that prove our hypothesis true indicate the validity of the Hakaru language implementation. Test cases that prove our hypothesis false indicate an underlying bug in the language definition which is to be passed back to the language developers.

# Conclusions & Future Work

This project was aimed at increasing language accessibility and testing the validity of Hakaru. A lot of ground was covered in both of these aspects in this project, but compared to other popular languages, Hakaru is still under development and has a lot of room for improvement. Here are just a few examples of where major development can still be done.

- Languages features: Hakaru can be expanded further with new features like import statements and error/exception handling.
- Standard library development: there are still many types of distributions left to be implemented, such as multivariate distributions.
- Testing: There are a plethora of relationships between distributions that have yet to be investigated

## References

[1] L. Leemis, "Univariate Distribution Relationship Chart", Math.wm.edu, 2018. [Online]. Available:

 $http://www.math.wm.edu/\ leemis/chart/UDR/UDR.html.$ 

[2] P. Narayanan, J. Carette, W. Romano, C. Shan and R. Zinkov, Probabilistic Inference by Program Transformation in Hakaru (System Description), Functional and Logic Programming, pp. 62-79, 2016.