# Deriving a Probability Density Calculator (Functional Pearl)

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## Abstract

Given an expression that denotes a probability distribution, often we want a corresponding *density* function, to use in probabilistic inference. Fortunately, the task of finding a density has been automated. It turns out that we can *derive* a compositional procedure for finding a density, by equational reasoning about integrals, starting with the mathematical specification of what a density is. Moreover, the density found can be run as an estimation algorithm, as well as simplified as an exact formula to improve the estimate.

*Categories and Subject Descriptors* G.3 [*Probability and Statistics*]: distribution functions; F.3.1 [*Logics and Meanings of Programs*]: Specifying and Verifying and Reasoning about Programs—specification techniques; D.1.1 [*Programming Techniques*]: Applicative (Functional) Programming

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## 1. Introduction

A popular way to handle uncertainty in AI, statistics, and science is to compute with probability distributions. Typically, we define a distribution then answer questions about it such as "what is its expected value?" and "what does its histogram look like?". Over a century, practitioners of this approach have identified many patterns in how to define distributions (that is, *modeling*) and how to answer questions about them (called *inference*). These patterns constitute the beginning of a *combinator library* (Hughes 1995).

Unfortunately, models and inference procedures do not compose in tandem: as illustrated in Sections 3.1 and 8.2, often we rejoice that a large distribution we're interested in can be expressed naturally by composing smaller distributions, but then despair that many questions we want to pose about the overall distribution cannot be answered using answers to corresponding questions about the constituent distributions. In other words, the natural compositional structure of models and of inference procedures are not the same. This mismatch is disappointing because it makes it harder for us to automate the labor-intensive process of turning a distribution that models the world into a program that answers relevant questions about it. This difficulty is the bane of declarative programming. It's like trying to build a SAT solver that generates assignments satisfying a compound expression  $e_1 \wedge e_2$  by combining assignments satisfying the subexpressions  $e_1$  and  $e_2$ .

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				Var v : a	ı
		$x \in \mathbb{R}$	e : a	e' : b	
StdRando	om : <i>Real</i>	it x : Real	Let v	e e' : b	
e : Real	e : Real	e : F	Real	<i>e</i> : <i>B</i>	ool
Neg e : Real	Exp e : Real	Log e	: Real	Not e :	Bool
$e_1$ : Real $e_2$ : R	eal e <sub>1</sub> : Real	$e_2$ : Real	e : Bool	$e_1$ : $a$	$e_2: a$
Add $e_1 e_2$ : Red	al Less $e_1$	$e_2$ : Bool	lf e	$e e_1 e_2$ :	а

**Figure 1.** The type system of our language of distributions

Still, there's hope to answer more inference questions while following the natural compositional structure of models, if only we could figure out how to generalize the questions as if strengthening an induction hypothesis or adding an accumulator argument. This paper tells one such success story. We answer the questions

- 1. "What is the expected value of this distribution?"
- 2. "What is a density function of this distribution?"

by generalizing them to compositional interpreters. Our interpreters are compilers in the sense that their output can be simplified using a computer algebra system or executed as a randomized algorithm. We derive these compilers by equational reasoning from a semantic specification. Our derivation appeals to  $\lambda$ -calculus equations alongside integral-calculus equations.

## 2. A Language of Generative Stories

To be concrete, we define a small language of distributions:

$e ::= StdRandom \mid Lit x \mid Var v \mid Let v e e$
$ \operatorname{Neg} e   \operatorname{Exp} e   \operatorname{Log} e   \operatorname{Not} e$
Add <i>e e</i>   Less <i>e e</i>   If <i>e e e</i>
v
x

Figure 1 shows our type system. To keep things simple, we include only two types in this language, *Real* and *Bool*. As usual, the typing judgment e : a means that the expression e has the type a. In Let v e e', the bound variable v takes scope over e' and not e.

Each expression says how to generate a random outcome. For example, the atomic expression StdRandom says to choose a random real number uniformly between 0 and 1. That's why its type is *Real*. To take another example, the compound expression

#### Add StdRandom StdRandom

says to choose two random real numbers independently, each uniformly between 0 and 1, then sum them. The sum is again a real number, so this expression's type is also Real. These descriptions of how to generate a random outcome are called generative stories by applied statisticians, and also called generators in QuickCheck (Claessen and Hughes 2000). The intuitive meaning of a generative story is a distribution over its outcomes, such as over reals. Because generative stories are intuitive to tell, and because they make it easy to detect dependencies among random choices (Pearl 1988), it's popular to express probability distributions by composing generative stories-such as using Add. The syntax of our language thus embodies the "natural compositional structure of models" referred to in the introduction above.

We define a Haskell data type Expr to encode the expressions of our language. Actually, using the GADT (generalized algebraic data type) extension, let's define two Haskell types Expr Real and *Expr Bool* at the same time, to distinguish our types *Real* and *Bool*.

## data Expr a where

StdRandor	n ::	Expr Real
Lit	:: Rational $\rightarrow$	Expr Real
Var	:: Var $a \rightarrow$	Expr a
Let	:: Var $a \rightarrow Expr \ a \rightarrow Expr \ b \rightarrow$	Expr b
Neg, Exp,		
Log	:: Expr Real $\rightarrow$	Expr Real
Not	$:: Expr Bool \rightarrow$	Expr Bool
Add	:: Expr Real $ ightarrow$ Expr Real $ ightarrow$	Expr Real
Less	:: Expr Real $ ightarrow$ Expr Real $ ightarrow$	Expr Bool
lf	$:: Expr Bool \rightarrow Expr a \rightarrow Expr a -$	$\rightarrow Expr a$

(Although we can easily construct an infinite Expr value by writing a recursive Haskell program, we avoid it because it would not correspond to any expression of our language.) We also define the types Var Real and Var Bool for variable names.

data Var a where Real :: String  $\rightarrow$  Var Real  $Bool :: String \rightarrow Var Bool$ 

But for brevity, we elide the constructors Real and Bool applied to literal strings in examples.

To interpret expressions in our language as generative stories, we write a function sample, which takes an expression and an environment as input and returns an IO action. To express that the type of the expression matches the outcome of the action, let's take the convenient shortcut of defining Real as a type synonym for Double, so that the Haskell type IO Real makes sense. The code for *sample* is straightforward:

#### type Real = Double

sample :: Expr  $a \rightarrow Env \rightarrow IO a$ *sample* StdRandom \_ = *getStdRandom random sample* (Lit x)  $_{-} = return (from Rational x)$ sample (Var v)  $\rho = return (lookupEnv \rho v)$ sample (Let v e e')  $\rho = \mathbf{do} x \leftarrow sample e \rho$ sample e' (extendEnv v x  $\rho$ ) sample (Neg e)  $\rho = liftM$  negate (sample  $e \rho$ ) sample (Exp e) $\rho = liftM exp$  $(sample \ e \ \rho)$ sample (Log e) $\rho = liftM \log$  $(sample \ e \ \rho)$ *sample* (Not *e*)  $\rho = liftM$  not  $(sample \ e \ \rho)$ sample (Add  $e_1 e_2$ )  $\rho = liftM2 (+)$ (sample  $e_1 \rho$ )  $(sample e_2 \rho)$ *sample* (Less  $e_1 e_2$ )  $\rho = liftM2$  (<) (sample  $e_1 \rho$ )  $(sample e_2 \rho)$ sample (If  $e e_1 e_2$ )  $\rho = \mathbf{do} b \leftarrow sample e \rho$ sample (if b then  $e_1$  else  $e_2$ )  $\rho$ 

As is typical of an interpreter, this sample function uses a type Env of environments (mapping variable names to values), along



Figure 2. Histograms of two distributions over real numbers. Each histogram is produced by generating 1000 samples (as shown at the end of Section 2) and putting them into 20 equally spaced bins.

with the functions lookupEnv and extendEnv for querying and extending environments. For concision, here we opt to represent environments as functions. All this code is standard:

**type** 
$$Env = \forall a.Var \ a \rightarrow a$$
  
 $lookupEnv :: Env \rightarrow Var \ a \rightarrow a$   
 $lookupEnv \ \rho = \rho$   
 $emptyEnv :: Env$   
 $emptyEnv v = error$  "Unbound"  
 $extendEnv :: Var \ a \rightarrow a \rightarrow Env \rightarrow Env$   
 $extendEnv$  (Real v)  $x_{-}$  (Real v')  $| v \equiv v' = x$   
 $extendEnv$  (Bool v)  $x_{-}$  (Bool v')  $| v \equiv v' = x$   
 $extendEnv_{-} - \rho v' = \rho$ 

 $-\rho v'$ 

We can now run our programs to get random outcomes:

 $= \rho v'$ 

> sample (Add StdRandom StdRandom) emptyEnv 0.8422448686660571 > sample (Add StdRandom StdRandom) emptyEnv 1.25881932199967 > sample (Let "x" StdRandom (Add (Var "x") (Var "x"))) emptyEnv 0.23258391029872305 > sample (Let "x" StdRandom (Add (Var "x") (Var "x"))) emptyEnv 1.1712041724765878

Your outcomes may vary, of course.

For more of a bird's-eye view of the distributions, we can take many independent samples then make a histogram out of each distribution. Two such histograms are shown in Figure 2. As those histograms indicate, the generative story of

Add StdRandom StdRandom

is different from the generative story of

Let "x" StdRandom (Add (Var "x") (Var "x"))

even though both expressions have the type Real. The latter expression means to choose just one random real number uniformly between 0 and 1, then double it. In general, Let means to choose an outcome once then use it any number of times. Thus, this is a call-by-value language whose side effect is random choice.

Although this language is small, our development below is wholly compatible with adding more types (such as Integer, products, and sums), arithmetic operations (such as division and sine), and distributions (such as normal, gamma, and Poisson). Besides, this language already lets us express many distributions, such as the exponential distribution, which ranges over the positive reals:

exponential :: Expr Real exponential = Neg (Log StdRandom)

## **3.** Composing Expectation Functionals

Although the *sample* interpreter is easy to write and intuitive to use, we shouldn't think that the *IO* action it returns is exactly the meaning of an expression. By "meaning" here, we mean what inference should preserve. We often want to reduce e to some other expression e' to speed up inference. The problem with treating *sample* e as the meaning of e is that such reduction would usually change the meaning of e, because *sample* e' makes different and fewer random choices than *sample* e.

For example, the expression Let "x" StdRandom (Lit 3) always produces the outcome 3, so we should be allowed to reduce it to just Lit 3, and an inference procedure shouldn't be obliged to consume any random seed before generating the 3. In other words, inference shouldn't be obliged to distinguish Lit 3 from Let "x" StdRandom (Lit 3), so we should assign these expressions the same meaning. They draw outcomes from the same distribution.

A less trivial example is that the definition of *sample* above specifies that, in an expression of the form Add  $e_1 e_2$  or Less  $e_1 e_2$ , the random choices in  $e_1$  must be made before the random choices in  $e_2$ , even though the order doesn't matter. Since addition is commutative, we should assign Add StdRandom (Neg StdRandom) and Add (Neg StdRandom) StdRandom the same meaning. They draw outcomes from the same random seed gives different outcomes.

Thus, the meaning equivalence relation produced by the *sample* semantics is too fine-grained. To coarsen the equivalence properly, measure theory informs us to consider the *expected values* of distributions. Given an expression of type *Real*, its expected value, or *mean*, is basically what the average of many samples approaches as the number of samples approaches infinity. For example, if we run

#### > sample (Add StdRandom StdRandom) emptyEnv

many times and average the results, the average will approach 1 as we take more samples. In the examples above, the expressions Let "x" StdRandom (Lit 3) and Lit 3 both have the expected value 3, and the expressions Add  $e_1 e_2$  and Add  $e_2 e_1$  always have the same expected value.

#### 3.1 The Expectation Interpreter

If the only question we ever ask about a distribution is "what is its expected value?", then it would be adequate for the meaning of each expression to equal its expected value. Unfortunately, there are other questions we ask whose answers differ on expressions with the same expected value. For example, given an expression of type *Real*, we might ask "what is the probability for its outcome to be less than 1/2?"—perhaps to decide how to bet on it. The two distributions sampled in Figure 2 both have expected value 1, but the probability of being less than 1/2 is 1/8 for the first distribution and 1/4 in the second distribution. Put differently, even though the two distributions have the same expected value, plugging them into the same *context* 

If (Less ... (Lit (1/2))) (Lit 1) (Lit 0)

gives two distributions with different expected values  $(1/8 \neq 1/4)$ . Even if we know the expected value of an expression *e*, we don't necessarily know the expected value of the larger expression

If 
$$(\text{Less } e (\text{Lit} (1/2))) (\text{Lit } 1) (\text{Lit } 0)$$

containing e.

Another way to phrase this complaint is to say that the expectedvalue interpretation is not compositional. That is, if we were to define a Haskell function

 $mean :: Expr Real \rightarrow Env \rightarrow Real$ 

then it wouldn't be straightforward the way *sample* is. For example, as the counterexamples in Figure 2 show, there's no way to define

*mean* (If (Less *e* (Lit 
$$(1/2)$$
)) (Lit 1) (Lit 0))  $\rho = \cdots$ 

in terms of mean e. In particular, it's incorrect to define

$$mean (If (Less e (Lit (1/2))) (Lit 1) (Lit 0)) \mu$$
  
= if mean  $e < 1/2$  then 1 else 0

because it would give the result 0 on the distributions in Figure 2, not 1/8 and 1/4 as it should! People building a compiler for distributions, including the present authors, want compositionality in order to achieve separate compilation.

To make *mean* compositional, we add an argument to it to represent the context (Hughes 1995; Hinze 2000) that an expression is plugged into before its expected value is observed. The new function *expect* has the type signature

 $expect :: Expr \ a \to Env \to (a \to Real) \to Real$ 

where the third argument may or may not be the identity function. In other words, the question that *expect e*  $\rho$  *c* asks is "what is the expected value of the distribution *e* in the environment  $\rho$  after its outcomes are transformed by the function *c*?". This value is also called the *expectation* of *c* with respect to the distribution. (To keep our derivation mathematically rigorous, we maintain the invariant that *c* is always measurable and non-negative (Pollard 2001, §2.4), but don't worry if you are not familiar with these side conditions.)

This generalization of *mean* is compositional: we can define *expect* on an expression in terms of *expect* on its subexpressions. Here is the definition:

```
expect StdRandom _ c = \int_0^1 \lambda x. c x
expect (Lit x) _ c = c (fromRational x)
expect (Var v)
                           \rho c = c (lookupEnv \rho v)
expect (Let v e e') \rho c = expect e \rho (\lambda x.
                                   expect e' (extendEnv v x \rho) c)
expect (Neg e)
                           \rho c = expect \ e \ \rho \ (\lambda x. c \ (negate \ x)))
                           \rho c = expect e \rho (\lambda x. c (exp))
expect (Exp e)
                                                                     (x))
expect (Log e)
                          \rho c = expect e \rho (\lambda x. c (log
                                                                     x))
                          \rho c = expect e \rho (\lambda x. c (not))
expect (Not e)
                                                                     x))
expect (Add e_1 e_2) \rho c = expect e_1 \rho (\lambda x.
                                   expect e_2 \rho (\lambda y.c (x+y)))
expect (Less e_1 e_2) \rho c = expect e_1 \rho (\lambda x.
                                   expect e_2 \rho (\lambda y. c (x < y)))
expect (If e e_1 e_2) \rho c = expect e \rho (\lambda b.
                                   expect (if b then e_1 else e_2) \rho c)
```

#### 3.2 Integrals Denoted by Random Choices

To understand this definition, let's start at the top.

The expected value of StdRandom is 1/2, but that's not the only question that *expect* StdRandom needs to answer. Given any function *c* from reals to reals (and any environment  $\rho$ ), *expect* StdRandom  $\rho$  *c* is supposed to be the expected value of choosing a random number uniformly between 0 and 1 then applying *c* to it. That expected value is the integral of *c* from 0 to 1, which in conventional mathematical notation is written as  $\int_0^1 c(x) dx$ . (More precisely, we mean the Lebesgue integral of *c* with respect to the Lebesgue measure from 0 to 1.) In this paper, we blend Haskell and mathematical notation by writing this integral as  $\int_0^1 \lambda x. c x$ , as if there's a function

$$\int_{\cdot}^{\cdot} \cdot :: Real \to Real \to (Real \to Real) \to Real$$

already defined. If we want to actually implement such a function, it could perform numerical integration. Or it could perform symbolic integration, or just print formulas containing  $\int$ , if we redefine the



**Figure 3.** The expectation of 3 different functions with respect to the same distribution StdRandom, defined in terms of integration from 0 to 1 (the shaded areas)

*Real* type and overload the *Num* class to produce text or syntax trees. (We also notate multiplication by  $\times$ , so c x always means applying c to x, as in Haskell, and not multiplying c by x.)

So for example, the expected value of *squaring* a uniform random number between 0 and 1 is

expect StdRandom emptyEnv 
$$(\lambda x.x \times x)$$
  
=  $\int_0^1 \lambda x.x \times x$   
=  $1/3$ 

which is less than 1/2 because squaring a number between 0 and 1 makes it smaller. And the probability that a uniform random number between 0 and 1 is less than 1/2 is

expect StdRandom emptyEnv ( $\lambda x$ . if x < 1/2 then 1 else 0) =  $\int_0^1 \lambda x$ . if x < 1/2 then 1 else 0 = 1/2

Figure 3 depicts these integrals.

=

=

The rest of the definition of *expect* is in *continuation-passing style* (Fischer 1993; Reynolds 1972; Strachey and Wadsworth 1974). The continuation c is the function whose expectation we want. The Lit and Var cases are deterministic (that is, they don't make any random choices), so the expectation of c with respect to those distributions simply applies c to one value. The unary operators (Neg, Exp, Log, Not) each compose the continuation with a mathematical function.

The remaining cases of *expect* involve multiple subexpressions and produce nested integrals if these subexpressions each yield integrals. For example, it follows from the definition that

expect (Add StdRandom (Neg StdRandom)) emptyEnv c = expect StdRandom emptyEnv ( $\lambda x$ . expect StdRandom emptyEnv ( $\lambda y$ . c (x + negate y))) =  $\int_0^1 \lambda x$ .  $\int_0^1 \lambda y$ . c (x - y)

The nesting order on the last line doesn't matter, as Tonelli's theorem (Pollard 2001, §4.4) assures it's equal to  $\int_0^1 \lambda y \cdot \int_0^1 \lambda x \cdot c \ (x-y)$ . In general, Tonelli's theorem lets us exchange nested integrals as long as the integrand (here  $\lambda x y \cdot c \ (x-y)$ ) is measurable and nonnegative. Thus we could just as well define equivalently

$$expect (Add e_1 e_2) \rho c = expect e_2 \rho (\lambda y.$$

$$expect e_1 \rho (\lambda x. c (x+y)))$$

$$expect (Less e_1 e_2) \rho c = expect e_2 \rho (\lambda y.$$

$$expect e_1 \rho (\lambda x. c (x < y)))$$

Besides compositionality, another benefit of generalizing *expect* is that it subsumes every question we can ask about a distribution. After all, a distribution is completely determined by the probability it assigns to each set of outcomes, and the probability of a set is the expectation of the set's characteristic function—the function

that maps outcomes in the set to 1 and outcomes not in the set to 0. For example, if e is a closed expression of type *Real*, then the probability that the outcome of e is less than 1/2 is

expect e emptyEnv (
$$\lambda x$$
. if  $x < 1/2$  then 1 else 0)

Not only can we express the expected value of the distribution e as

mean 
$$e \rho = expect e \rho id$$

but we can also express all the other *moments* of *e*, such as the *variance* (the expected squared difference from the mean):

*variance* :: *Expr Real*  $\rightarrow$  *Env*  $\rightarrow$  *Real variance e*  $\rho$  = *expect e*  $\rho$  ( $\lambda x. (x - mean e \rho)^2$ )

To take another example, the ideal height of each histogram bar in Figure 2 is

expect e emptyEnv ( $\lambda x$ . if  $lo < x \le hi$  then 1000 else 0)

where *lo* and *hi* are the bounds of the bin and 1000 is the total number of samples. (We abbreviate  $lo < x \land x \le hi$  to  $lo < x \le hi$ .)

Mathematically speaking, every distribution corresponds to a *functional*, which is a function—sort of a generalized integrator—that takes as argument another function, namely the integrand *c*. This correspondence is expressed by *expect*, and it's injective. (In fact, it's bijective between measures and "increasing linear functionals with the Monotone Convergence property" (Pollard 2001, page 27).) Therefore, if *expect e*  $\rho$  and *expect e'*  $\rho$  are equal (in other words, if *expect e*  $\rho$  *c* and *expect e'*  $\rho$  *c* are equal for all *c*), then *e* and *e'* are equivalent and we can feel free to reduce *e* to *e'*.

In short, we define the meaning of the expression e in the environment  $\rho$  to be the functional *expect e*  $\rho$ . Returning to Figure 2, it follows from this definition that the meaning of

Add StdRandom StdRandom

in the empty environment is

$$= \lambda c. \int_0^1 \lambda x. \int_0^1 \lambda y. c (x+y)$$

and the meaning of

Let "x" StdRandom (Add (Var "x") (Var "x"))

in the empty environment is

$$= \lambda c. \int_0^1 \lambda x. c (x+x)$$

The two expressions are not equivalent, because the two functionals are not equal: applied to the function  $\lambda x.$  if x < 1/2 then 1 else 0, the first functional returns 1/8 whereas the second functional returns 1/4. In this way, *expect* defines the semantics of our distribution language. Therefore, it naturally enters our specification of a probability density calculator, which we present next.

#### 4. Specifying Probability Densities

Some distributions enjoy the existence of a *density function*. If the distribution is over the type a, then the density function maps from a to reals. Density functions are very useful in probabilistic inference: they underpin many concepts and techniques, including *maximum-likelihood estimation*, *conditioning*, and *Monte Carlo sampling* (MacKay 1998; Tierney 1998). We illustrate these applications in Section 7 below.

Intuitively, a density function is the outline of a histogram as the bin size approaches zero. For example, the two distributions in Figure 2 have the respective density functions shown in Figure 4. The shapes in the two figures are similar, but the histograms are



Figure 4. Density functions of the two distributions in Figure 2

randomly generated as this paper is typeset, whereas each density is a fixed mathematical function.

The precise definition of when a given function qualifies as a density for a given distribution depends on a *base* or *dominating measure*. When the distribution is over reals, the base measure is typically the Lebesgue measure over reals, in which case the definition amounts to the following.

**Definition 1.** A function  $d :: Real \rightarrow Real$  is a *density* (with respect to the Lebesgue measure) for an expression e :: Expr Real in an environment  $\rho :: Env$  if and only if

expect 
$$e \rho c = \int_{-\infty}^{\infty} \lambda t d t \times c t$$

for all continuations  $c :: Real \rightarrow Real$ .

And when the distribution is over booleans, the base measure is typically the counting measure over booleans, in which case the definition amounts to the following.

**Definition 2.** A function  $d::Bool \rightarrow Real$  is a *density* (with respect to the counting measure) for an expression e::Expr Bool in an environment  $\rho::Env$  if and only if

expect 
$$e \rho c = sum [d t \times c t | t \leftarrow [True, False]]$$

for all continuations  $c :: Bool \rightarrow Real$ .

One way to gain intuition for these definitions is to let c be the characteristic function of a set. For example, in Definition 2, if

$$c = \lambda t$$
. if t then 1 else (

then *expect e*  $\rho$  *c* is the probability of *True* according to *e* in  $\rho$ , and the equation requires *d True* to equal it. In Definition 1, if

$$c = \lambda t$$
. if  $lo < t \le hi$  then 1 else 0

where *lo* and *hi* are the bounds of a histogram bin, then *expect e*  $\rho$  *c* is the probability of that bin according to *e* in  $\rho$ , and the equation requires  $\int_{l_{\rho}}^{h_{i}} d$  to equal the ideal proportion of that histogram bar.

These definitions use e and  $\rho$  just to represent the functional *expect*  $e \rho$ . Given e and  $\rho$ , because densities are useful, our goal is to find some function d that satisfies the specification above.

#### 4.1 Examples of Densities and Their Non-existence

To illustrate these definitions, let's check that the functions in Figure 4 are indeed densities of their respective distributions. First let's consider the function on the right of Figure 4, which is supposed to be a density for

$$e = \text{Let "x" StdRandom (Add (Var "x") (Var "x"))}$$

in *emptyEnv*. We start with the functional

 $m = expect \ e \ empty Env = \lambda c. \int_0^1 \lambda x. c \ (x+x)$ 

and reason equationally by univariate calculus. We extend the domain of integration from the interval (0,1) to the entire real line:

$$m = \lambda c. \int_{-\infty}^{\infty} \lambda x.$$
 (if  $0 < x < 1$  then 1 else  $0) \times c (x + x)$ 

Then we change the integration variable from *x* to t = x + x:

$$m = \lambda c. \int_{-\infty}^{\infty} \lambda t. (1/2) \times (\text{if } 0 < t/2 < 1 \text{ then } 1 \text{ else } 0) \times c t$$

(The factor 1/2 is the (absolute value of the) derivative of x = t/2 with respect to *t*.) Matching this equation against Definition 1 shows that

$$\lambda t. (1/2) \times (\mathbf{if} \ 0 < t/2 < 1 \ \mathbf{then} \ 1 \ \mathbf{else} \ 0)$$
  
=  $\lambda t. \mathbf{if} \ 0 < t < 2 \ \mathbf{then} \ 1/2 \ \mathbf{else} \ 0$ 

is a density as desired. By the way, because changing the value of the integrand at a few points does not affect the integral, functions such as

$$\lambda t$$
. if  $t \equiv 1 \lor t \equiv 3$  then 42 else if  $0 < t \le 2$  then  $1/2$  else 0

are densities just as well.

Turning to the function on the left of Figure 4, we want to check that it is a density for

e = Add StdRandom StdRandom

in emptyEnv. Again we start with the functional

 $m = expect \ e \ emptyEnv = \lambda c. \int_0^1 \lambda x. \int_0^1 \lambda y. c \ (x+y)$ 

and do calculus. We extend the inner domain of integration from the interval (0, 1) to the entire real line:

$$n = \lambda c. \int_0^1 \lambda x. \int_{-\infty}^\infty \lambda y.$$
 (if  $0 < y < 1$  then 1 else  $0) \times c (x+y)$ 

Then we change the inner integration variable from *y* to t = x + y:

$$m = \lambda c. \int_0^1 \lambda x. \int_{-\infty}^\infty \lambda t.$$
 (if  $0 < t - x < 1$  then 1 else 0)  $\times c t$ 

(No factor is required because the (partial) derivative of y = t - x with respect to t is 1.) Tonelli's theorem lets us exchange the nested integrals:

$$m = \lambda c. \int_{-\infty}^{\infty} \lambda t. \int_{0}^{1} \lambda x.$$
 (if  $0 < t - x < 1$  then 1 else 0)  $\times c t$ 

Finally, because the inner integration variable *x* does not appear in the factor *c t*, we can pull *c t* out (in other words, we can use the linearity of  $\int_0^1 \cdot$ ):

$$m = \lambda c. \int_{-\infty}^{\infty} \lambda t. \left( \int_{0}^{1} \lambda x. \text{ if } 0 < t - x < 1 \text{ then } 1 \text{ else } 0 \right) \times c t$$

Matching this last equation against Definition 1 shows that

$$\lambda t \cdot \int_0^1 \lambda x \cdot \mathbf{if} \ 0 < t - x < 1$$
 then 1 else 0

is a density. This formula can be further simplified to the desired closed form in the lower-left corner of Figure 4, either by hand or using a computer algebra system.

So far we have seen two examples of distributions and their densities. But not all distributions have a density. For example, when e = Lit 3, we have

$$n = expect \ e \ \rho = \lambda c.c.3$$

so a density function d would have to satisfy

$$c 3 = \int_{-\infty}^{\infty} \lambda t d t \times c t$$

for all  $c::Real \rightarrow Real$ . But if  $c = \lambda t$ . if  $t \equiv 3$  then 1 else 0, then the left-hand-side is 1 whereas the right-hand-side is

$$\int_{-\infty}^{\infty} \lambda t$$
. if  $t \equiv 3$  then  $d t$  else 0

which is 0 no matter what  $d::Real \rightarrow Real$  is. So there's no such d.

It may surprise the reader that we say Lit 3 has no density, because sometimes Lit 3 is said to have a density, easily expressed in terms of the *Dirac delta function*. However, the Dirac delta is not a function in the ordinary sense but a *generalized function*, which only makes sense under integration. For example, the Dirac delta doesn't map any number to anything, but rather integrates an ordinary function c to yield c(0). Whereas an ordinary density function has the type  $a \rightarrow Real$ , a generalized function has the type  $(a \rightarrow Real) \rightarrow Real$ , same as produced by *expect*. So, generalized functions are essentially distributions, which is indeed what many people call them. In this paper we seek ordinary density functions, which are much more useful. For example, generalized functions cannot be multiplied together, which precludes their use in Monte Carlo sampling techniques such as *importance sampling* (illustrated in Section 7) and *Metropolis-Hastings sampling* (MacKay 1998; Tierney 1998).

#### 4.2 Relation to Prior Density Calculators

Because density functions are useful, we want a program that automatically computes density functions from distribution expressions. Such a program can "compute functions" in two different senses of the phrase. Pfeffer's (2009, §5.2) density calculator is a random algorithm that produces a number. By running the algorithm many times and averaging the results, we can approximate the density of a distribution at a given point. In contrast, Bhat et al.'s (2012, 2013) density calculator deterministically produces an exact mathematical formula (which may contain integrals). As they suggest, we can then feed the formula to a computer algebra system or inference procedure to be analyzed or executed.

In the rest of this paper, we use equational reasoning to *derive* a compositional density calculator for the first time. Even though Definitions 1 and 2 seem to require conjuring a function out of thin air, the semantic specification above turns out to pave the way for the equational derivation below. The resulting calculator produces a density function that can be treated either as an exact formula (not necessarily closed form) or as an approximation algorithm, depending on whether  $\int$  is treated as symbolic or numerical integration. It thus unites the density calculators of Pfeffer (2009) and Bhat et al. (2012, 2013). In particular, the variety of program constructs that those techniques and ours can handle appear to be the same.

### 5. Calculating Probability Densities

To recap, our goal in the rest of this paper to derive a program that, given *e* and  $\rho$ , finds a function *d* that satisfies Definitions 1 and 2.

We just saw that not every distribution has a density. Moreover, not every density can be represented using the operations on *Real* available to us. And even if a density can be represented, discovering it may require mathematical reasoning too advanced to be automated compositionally. For all these reasons, we have to relax our goal: let's write a program

*density* :: *Expr*  $a \rightarrow [Env \rightarrow a \rightarrow Real]$ 

that maps each distribution expression *e* to a *list* of successes (Wadler 1985). For every element  $\delta$  of the list, and for every environment  $\rho$  that binds all the free variables in *e*, we require that the function  $\delta \rho$  be a density for *e* in  $\rho$ . That is, depending on if *e* has type *Expr Real* or *Expr Bool*, we want one of the two equations

expect 
$$e \rho c = \int_{-\infty}^{\infty} \lambda t. \delta \rho t \times c t$$
  
expect  $e \rho c = sum [\delta \rho t \times c t | t \leftarrow [True, False]$ 

to hold for all  $\delta$ ,  $\rho$ , and *c*.

The list returned by *density* might be empty, but we'll do our best to keep it non-empty. For example, as shown in Section 4, we regret that *density* (Lit 3) must be the empty list, but

density (Let "x" StdRandom (Add (Var "x") (Var "x")))

and

density (Add StdRandom StdRandom)

can be the non-empty lists

 $[\lambda \rho t. \mathbf{if } 0 < t < 2 \mathbf{ then } 1/2 \mathbf{ else } 0] \\ [\lambda \rho t. \int_0^1 \lambda x. \mathbf{if } 0 < t - x < 1 \mathbf{ then } 1 \mathbf{ else } 0]$ 

respectively. Our density calculator ends up missing the first density (see Section 8.1), but it does find the second as soon as we introduce nondeterminism by concatenating lists (see Section 5.5).

The fact that not every distribution has a density holds another lesson for us. It turns out that *density* is not compositional. In other words, *density* on an expression cannot be defined in terms of *density* on its subexpressions, for the following reason. On one hand, Lit 3 and Lit 4 have no density, so *density* must map them both to the empty list. On the other hand, the larger expressions Add (Lit 3) StdRandom and Add (Lit 4) StdRandom have densities but different ones, so we want *density* to map them to different non-empty lists. Thus, *density e* does not determine *density* (Add *e* StdRandom). Instead, it will be in terms of *expect e* that we define *density* (Add *e* StdRandom). That is, although *density* is not compositional, the interpreter  $\lambda e$ . (*density e*, *expect e*) is compositional (but see Section 8.2).

We define *density e* by structural induction on *e*.

#### 5.1 Real Base Cases

An important base case is when e = StdRandom: we define

density StdRandom =  $[\lambda \rho t. if 0 < t \land t < 1$  then 1 else 0]

expressing the characteristic function of the unit interval. This clause satisfies Definition 1 because

expect StdRandom  $\rho$  c = -- definition of expect  $\int_0^1 \lambda t. c t$ = -- extending the domain of integration  $\int_{-\infty}^{\infty} \lambda t. (\mathbf{if} \ 0 < t \land t < 1 \mathbf{ then } 1 \mathbf{ else } 0) \times c t$ 

For the other base cases of type *Real*, we must fail, as discussed in Section 4.1.

 $density (Lit _) = []$  $density (Var (Real _)) = []$ 

## 5.2 Boolean Cases

In a countable type such as *Bool* (in contrast to *Real*), every distribution has a density. In other words, there always exists a function *d* that satisfies Definition 2. We can derive it as follows:

 $expect \ e \ \rho \ c$   $= \ - \ \eta \ expansion$   $expect \ e \ \rho \ (\lambda x. c \ x)$   $= \ - \ case \ analysis \ on \ x$   $expect \ e \ \rho \ (\lambda x. sum \ [ \ (if \ t \equiv x \ then \ 1 \ else \ 0) \times c \ t$   $| \ t \leftarrow [True, False]])$   $= \ - \ Tonelli's \ theorem, \ or \ just \ linearity \ of \ m$   $sum \ [ \ expect \ e \ \rho \ (\lambda x. if \ t \equiv x \ then \ 1 \ else \ 0) \times c \ t$   $| \ t \leftarrow [True, False]]$ 

In the right-hand-side above, *expect e*  $\rho$  ( $\lambda x$ . **if**  $t \equiv x$  **then** 1 **else** 0) is just the probability of the boolean value *t*. Matching the right-hand-side against Definition 2 shows that the function *prob e*  $\rho$  defined by

*prob* :: *Expr Bool*  $\rightarrow$  *Env*  $\rightarrow$  *Bool*  $\rightarrow$  *Real prob*  $e \rho t = expect e \rho (\lambda x. \text{ if } t \equiv x \text{ then } 1 \text{ else } 0)$ 

is a density for e in  $\rho$ . Accordingly, we define

density (Var (Bool v)) = [prob (Var (Bool v))]density (Not e) = [prob (Not e) ] $density (Less e_1 e_2) = [prob (Less e_1 e_2) ]$ 



**Figure 5.** A density of Neg StdRandom (top) results from transforming a density of StdRandom (bottom)

#### 5.3 Unary Cases

Things get more interesting in the other recursive cases. Take the case *density* (Neg *e*) for example. Suppose that the recursive call *density e* returns the successful result  $\delta$ , so the induction hypothesis is that the equation

expect 
$$e \rho c = \int_{-\infty}^{\infty} \lambda t \cdot \delta \rho t \times c t$$

holds for all  $\rho$  and c. We seek some  $\delta'$  such that the equation

expect (Neg e)  $\rho c = \int_{-\infty}^{\infty} \lambda t \, \delta' \rho t \times c t$ 

holds for all  $\rho$  and c. Starting with the left-hand-side, we calculate

 $expect (Neg e) \rho c$  = -- definition of expect

expect  $e \rho (\lambda x. c (-x))$ 

= --- induction hypothesis, substituting  $\lambda x. c (-x)$  for  $c \int_{-\infty}^{\infty} \lambda x. \delta \rho x \times c (-x)$ 

= -- changing the integration variable from x to t = -x $\int_{-\infty}^{\infty} \lambda t . \delta \rho (-t) \times c t$ 

Therefore, to match the goal, we define

density (Neg e) =  $[\lambda \rho t. \delta \rho (-t) | \delta \leftarrow density e]$ 

This clause says that the density of Neg *e* at *t* is the density of *e* at -t. This makes sense because the histogram of Neg *e* is the horizontal mirror image of the histogram of *e*. For example, Figure 5 depicts how the density of Neg StdRandom at *t* is the density of StdRandom at -t: when a sample from Neg StdRandom occurs between -0.5 and -0.4 (shaded above), it's because a sample from StdRandom occurred between 0.4 and 0.5 (shaded below).

A slightly more advanced case is *density* ( $E \times p e$ ). Again, we assume the induction hypothesis

expect 
$$e \rho c = \int_{-\infty}^{\infty} \lambda t \cdot \delta \rho t \times c t$$

and seek some  $\delta'$  satisfying

expect (Exp e) 
$$\rho c = \int_{-\infty}^{\infty} \lambda t. \, \delta' \, \rho \, t \times c \, t$$

Starting with the left-hand-side, we calculate

*expect* (Exp e)  $\rho$  c

- = -- definition of *expect*
- expect  $e \rho (\lambda x. c (exp x))$
- = -- induction hypothesis, substituting  $\lambda x. c (exp x)$  for  $c \int_{-\infty}^{\infty} \lambda x. \delta \rho x \times c (exp x)$
- = -- changing the integration variable from x to t = exp x $\int_0^\infty \lambda t. (\delta \rho (\log t)/t) \times c t$
- = -- extending the domain of integration  $\int_{-\infty}^{\infty} \lambda t. (\mathbf{if} \ 0 < t \ \mathbf{then} \ \delta \ \rho \ (\log t)/t \ \mathbf{else} \ 0) \times c \ t$



**Figure 6.** A density of Exp StdRandom (top) results from transforming a density of StdRandom (bottom)

Compared to the Neg case above, this calculation illustrates two complications:

- 1. The second-to-last step introduces the factor 1/t, which is the (absolute value of the) derivative of x = log t with respect to t. This factor makes sense because the histogram of Exp e is a *distorted* image of the histogram of e. For example, Figure 6 depicts how the density of Exp StdRandom at t is the density of StdRandom at log t, multiplied by 1/t because the interval  $(e^{0.4}, e^{0.5})$  (shaded above) is *wider* than the interval (0.4, 0.5) (shaded below) (Freedman et al. 2007, Chapter 3). After all, when a sample from Exp StdRandom occurs between  $e^{0.4}$  and  $e^{0.5}$ , it's because a sample from StdRandom occured between 0.4 and 0.5, so the two shaded areas in Figure 6 should be equal.
- 2. The last step introduces a conditional to account for the fact that the result of exponentiation is never negative.

In the end, to match the goal, we define

density (Exp e) = 
$$[\lambda \rho t. \mathbf{if} 0 < t \mathbf{then} \delta \rho (\log t)/t \mathbf{else} 0$$
  
 $|\delta \leftarrow density e]$ 

The case *density* (Log e) can be handled similarly, so we omit the derivation:

density 
$$(\text{Log } e) = [\lambda \rho \ t. \delta \rho \ (exp \ t) \times exp \ t \mid \delta \leftarrow density \ e]$$

We can add other unary operators as well, such as reciprocal. (Alternatively, we can express reciprocal in terms of Exp, Neg, and Log, like with a slide rule.)

All these unary operators have in common that their densities *invert* their usual interpretation as functions: the density of Exp e at *t* uses the density of *e* at *log t*; the density of Neg *e* at *t* uses the density of *e* at -t; and so on. This pattern underlies the intuition that density calculation is backward sampling.

#### 5.4 Conditional

For the case *density* (If  $e e_1 e_2$ ), suppose that the recursive calls *density*  $e_1$  and *density*  $e_2$  return the successful results  $\delta_1$  and  $\delta_2$ . (It turns out that we don't need a density for the subexpression *e*.) We seek some  $\delta'$  such that the equation

expect (If  $e e_1 e_2$ )  $\rho c = \int \lambda t \cdot \delta' \rho t \times c t$ 

holds for all  $\rho$  and *c*. Here the linear functional  $\int \cdot$  is either  $\int_{-\infty}^{\infty} \cdot$  (if  $e_1$  and  $e_2$  have type *Expr Real*) or  $\lambda c. sum (map c [True, False])$  (if  $e_1$  and  $e_2$  have type *Expr Bool*). Starting with the left-hand-side, we calculate

$$expect (If e e_1 e_2) \rho c$$

$$= -- definition of expect$$

$$expect e \rho (\lambda b. expect (if b then e_1 else e_2) \rho c)$$

$$= -- induction hypotheses$$

$$expect e \rho (\lambda b. \int \lambda t. (if b then \delta_1 else \delta_2) \rho t \times c t)$$

$$= -- Tonelli's theorem, exchanging the integrals$$

$$-- expect e \rho (\lambda b...) and \int \lambda t... \times c t$$

$$\int \lambda t. expect e \rho (\lambda b. (if b then \delta_1 else \delta_2) \rho t) \times c t$$

Therefore, to match the goal, we define

density (If 
$$e e_1 e_2$$
) = [ $\lambda \rho t$ . expect  $e \rho (\lambda b$ .  
(if  $b$  then  $\delta_1$  else  $\delta_2$ )  $\rho t$ )  
|  $\delta_1 \leftarrow$  density  $e_1, \delta_2 \leftarrow$  density  $e_2$ ]

Using the fact that  $\lambda b...$  above is just a function from *Bool* to *Real* (essentially the *Real* pair ( $\delta_1 \rho t, \delta_2 \rho t$ )), we can rewrite this definition more intuitively:

density (If 
$$e e_1 e_2$$
) = [ $\lambda \rho t$ . prob  $e \rho$  True  $\times \delta_1 \rho t$   
+ prob  $e \rho$  False  $\times \delta_2 \rho t$   
|  $\delta_1 \leftarrow$  density  $e_1, \delta_2 \leftarrow$  density  $e_2$ 

For example, if *e* is Less StdRandom (Lit (1/2)), then

prob e 
$$\rho$$
 True = prob e  $\rho$  False =  $1/2$ 

and to sample from If  $e e_1 e_2$  is to flip a fair coin to decide whether to sample from  $e_1$  or from  $e_2$ . Accordingly, the density of If  $e e_1 e_2$ is just the average of the densities of  $e_1$  and  $e_2$ .

#### 5.5 Binary Operators

Recall from Section 5.3 that the density of a unary operator f(x) inverts f as a function of its operand x. The density of a binary operator f(x,y) can invert f as a function of either x or y, treating the other operand as fixed. This choice brings a new twist to our derivation, namely that our density calculator can be *nondeterministic*: it can try multiple strategies for finding a density. If multiple strategies succeed, the resulting density functions are equivalent, in that they disagree only on a zero-probability set of outcomes. (But their subsequent performance may differ, so we keep them all.)

Take Add  $e_1 e_2$  for example. The distribution denoted by Add  $e_1 e_2$  is the convolution of the distributions denoted by  $e_1$  and  $e_2$ . What we seek is some  $\delta'$  such that the equation

expect (Add 
$$e_1 e_2$$
)  $\rho c = \int_{-\infty}^{\infty} \lambda t \cdot \delta' \rho t \times c t$ 

holds for all  $\rho$  and c.

Again starting with the left-hand-side, we calculate

```
expect (Add e_1 e_2) \rho c
= -- definition of expect
expect e_1 \rho (\lambda x. expect e_2 \rho (\lambda y. c (x+y)))
```

If the recursive call *density*  $e_2$  returns the successful result  $\delta_2$ , then the induction hypothesis lets us continue calculating as follows:

= -- induction hypothesis  

$$expect e_1 \rho (\lambda x. \int_{-\infty}^{\infty} \lambda y. \delta_2 \rho y \times c (x+y))$$
= -- changing the integration variable from y to  $t = x+y$   

$$expect e_1 \rho (\lambda x. \int_{-\infty}^{\infty} \lambda t. \delta_2 \rho (t-x) \times c t)$$
= -- Tonelli's theorem  

$$\int_{-\infty}^{\infty} \lambda t. expect e_1 \rho (\lambda x. \delta_2 \rho (t-x)) \times c t$$

Therefore, having solved for *y* in t = x + y, we can define

density (Add 
$$e_1 e_2$$
) = [ $\lambda \rho$  t. expect  $e_1 \rho$  ( $\lambda x. \delta_2 \rho$  (t-x))  
|  $\delta_2 \leftarrow$  density  $e_2$ ]

For example, when  $e_1$  is Lit 3 and  $e_2$  is StdRandom, this definition amounts to solving for y in t = 3 + y. The density calculator finds y = t - 3 and thus returns  $[\lambda \rho t. \mathbf{if} 0 < t - 3 < 1 \mathbf{then} 1 \mathbf{else} 0]$ 

expressing the characteristic function of the interval (3,4). We can also handle the example on the left of Figure 4 now: when  $e_1$  and  $e_2$ are both StdRandom, the density calculator returns

$$\left[\lambda\rho t, \int_0^1 \lambda x, \text{ if } 0 < t - x < 1 \text{ then } 1 \text{ else } 0\right]$$

because *expect*  $e_1 \rho$  produces the integral and  $\delta_2 \rho (t-x)$  produces the conditional.

By analogous reasoning, we can also solve for x and define

density (Add 
$$e_1 e_2$$
) = [ $\lambda \rho t. expect e_2 \rho (\lambda y. \delta_1 \rho (t-y))$   
|  $\delta_1 \leftarrow density e_1$ ]

Although solving for *y* and for *x* can produce overlapping lists (like when  $e_1$  and  $e_2$  are both StdRandom), the two lists do not subsume each other. For example, because Lit 3 has no density, only the first definition handles Add (Lit 3) StdRandom and only the second definition handles Add StdRandom (Lit 3). In the end, we define

$$density (\mathsf{Add} \ e_1 \ e_2) = [\lambda \rho \ t. expect \ e_1 \ \rho \ (\lambda x. \ \delta_2 \ \rho \ (t-x)) \\ | \ \delta_2 \leftarrow density \ e_2] \\ ++ [\lambda \rho \ t. expect \ e_2 \ \rho \ (\lambda y. \ \delta_1 \ \rho \ (t-y)) \\ | \ \delta_1 \leftarrow density \ e_1]$$

We can add other binary operators, such as multiplication, to our language and handle them similarly. (Alternatively, we can express multiplication in terms of Exp, Add, and Log, like with a slide rule.)

## 5.6 Variable Binding and Sharing

As with Add, an expression Let v e e' may have a density even if one of its subexpressions e and e' doesn't. We call v the bound variable, e the *right-hand-side* (Landin 1964; Peyton Jones 2003), and e' the *body* of the Let. There are two strategies for handling Let.

First, if the body e' has a density  $\delta'$ , then a density of the Let is the expectation of  $\delta'$  with respect to the right-hand-side e. That is, if the recursive call *density* e' returns the successful result  $\delta'$ , then we calculate

expect (Let 
$$v e e'$$
)  $\rho c$   
= -- definition of expect  
expect  $e \rho (\lambda x. expect e' (extendEnv v x \rho) c)$   
= -- induction hypothesis  
expect  $e \rho (\lambda x. \int \lambda t. \delta' (extendEnv v x \rho) t \times c t)$   
= -- Tonelli's theorem  
 $\int \lambda t. (expect e \rho (\lambda x. \delta' (extendEnv v x \rho) t)) \times c t$ 

Therefore, we can define

density (Let 
$$v e e'$$
)  
=  $[\lambda \rho t. expect e \rho (\lambda x. \delta' (extendEnv v x \rho) t)$   
 $|\delta' \leftarrow density e']$ 

This strategy handles Let expressions that use the bound variable as a parameter. The right-hand-side can be deterministic, as in

or random, as in

Let "x" StdRandom

(Add (Add (Var "x") (Var "x")) StdRandom)

These examples show that Var "x" can be used multiple times. That is, the outcome of the right-hand-side can be *shared*. We need Let in the language to introduce such sharing. However, this strategy fails on Let expressions whose bodies are deterministic, such as

Let "x" (Neg StdRandom) 
$$(Exp (Var "x"))$$

These Let expressions have densities only because their right-handsides are random. Hence we introduce another strategy for handling Let: check if the body of the Let uses the bound variable at most once. If so, we can inline the right-hand-side into the body. That is, we can replace Let v e e' by the result of substituting e for vin e', which we write as  $e'\{v \mapsto e\}$ . (This substitution operation sometimes needs to rename variables in e' to avoid capture.) This replacement preserves the meaning of the Let expression even if the body is random. For example, we can handle the expression

$$e_1 = \text{Let "x" (Neg StdRandom)}$$
  
(Add StdRandom (Exp (Var "x")))

by turning it into the equivalent expression

 $e_2 = \text{Add StdRandom}(\text{Exp}(\text{Neg StdRandom}))$ 

To see this equivalence, apply the definition of *expect* to  $e_1$  and  $e_2$ :

expect 
$$e_1 \rho c = \int_0^1 \lambda x. \int_0^1 \lambda t. c (t + exp(-x))$$
  
expect  $e_2 \rho c = \int_0^1 \lambda t. \int_0^1 \lambda x. c (t + exp(-x))$ 

Then use Tonelli's theorem to move inward the outer integral  $\int_0^1 \lambda x$  in *expect e*<sub>1</sub>  $\rho$  *c*, which corresponds to the random choice made in Neg StdRandom. If we think of random choice as a side effect, then Tonelli's theorem lets us delay evaluating the right-hand-side Neg StdRandom until the body Add StdRandom (Exp (Var "x")) actually uses the bound variable "x".

In general, Tonelli's theorem tells us that delayed evaluation preserves the expectation semantics of the expression Let  $v \ e \ e'$ when the body e' uses the bound variable v exactly once. Moreover, in the case where e' never uses v, delayed evaluation also preserves the expectation semantics, but for a different reason. If e' never uses v, then *expect* e' (*extendEnv*  $v \ x \rho$ )  $c = expect \ e' \rho \ c$ , so

$$expect (Let v e e') \rho c$$

$$= -- definition of expect$$

$$expect e \rho (\lambda x. expect e' (extendEnv v x \rho) c)$$

$$= -- e' never uses v$$

$$expect e \rho (\lambda x. expect e' \rho c)$$

$$= -- pull expect e' \rho c out of the integral expect e \rho$$

$$expect e \rho (\lambda x. 1) \times expect e' \rho c$$

Finally, a simple induction on *e* shows that *expect e*  $\rho$  ( $\lambda x.1$ ) always equals 1 in our language. In other words, the distributions denoted in our language are all *probability* distributions.

Backed by this reasoning, we put the two strategies together to define

$$\begin{array}{l} density \ (\text{Let } v \ e \ e') \\ = \left[ \lambda \rho \ t. expect \ e \ \rho \ (\lambda x. \ \delta' \ (extendEnv \ v \ x \ \rho) \ t) \\ & | \ \delta' \leftarrow density \ e'| & -- \ \text{first strategy} \\ ++ \left[ \delta' \ | \ usage \ e' \ v \leqslant AtMostOnce \\ & , \ \delta' \leftarrow density \ (e'\{v \mapsto e\}) \right] & -- \ \text{second strategy} \end{array}$$

The condition *usage*  $e' v \leq AtMostOnce$  above tests conservatively whether the expression e' uses the variable v at most once—in other words, whether v is *not shared* in e'. This test serves the purpose of Bhat et al.'s (2012) active variables and independence test. We relegate its implementation to Appendix A.

## 6. Approximating Probability Densities

We are done deriving our density calculator! It produces output rife with integrals. The definition of *density* itself does not contain integrals, but *expect* StdRandom contains an integral, and *density* invokes *expect* in the boolean, If, Add, and Let cases. For example, Section 5.5 shows one success of our density calculator:

density (Add StdRandom StdRandom)

- $= \begin{bmatrix} \lambda \rho \ t. \ expect \ \mathsf{StdRandom} \ \rho \ (\lambda x. \ \delta_2 \ \rho \ (t-x)) \\ \mid \delta_2 \leftarrow density \ \mathsf{StdRandom} \end{bmatrix} + \cdots$
- =  $[\lambda \rho t] \int_0^1 \lambda x$ . if 0 < t x < 1 then 1 else 0] ++ ...

One way to use *density* is to feed its output to a symbolic integrator, as Bhat et al. (2012) suggest. If we're lucky, we might get a closed form that can be run as an exact deterministic algorithm. For example, Maxima, Maple, and Mathematica can each simplify the successful result above to the closed form in the lower-left corner of Figure 4. To produce output that those systems can parse, we would redefine the *Real* type and overload the *Num* class, which is not difficult to do.

Even without computer algebra or without eliminating all integrals, we can execute the density found as a *randomized* algorithm whose *expected* output is the density at the given point. All it takes is interpreting each call from *density* to *expect* as sampling randomly from a distribution. This interpretation is a form of numerical integration, carried out by Pfeffer's (2009, §5.2) approximate algorithm for density estimation. For example, we can interpret the successful result above, as is, as the following randomized (and embarrassingly parallel) algorithm:

Given  $\rho$  and *t*, choose a random real number *x* uniformly between 0 and 1, then compute if 0 < t - x < 1 then 1 else 0.

When time is about to run out, we average the results from repeated independent runs of this algorithm.

## 7. Applications of Our Density Calculator

Returning to our motivation, let us briefly demonstrate two ways to use density functions in probabilistic inference.

#### 7.1 Computing and Comparing Likelihoods

Imagine that we are adjudicating between two competing scientific hypotheses, which we model by two generative stories  $e_1$  and  $e_2$ . Using their density functions, we can update our belief in light of empirical evidence.

Concretely, suppose we conduct many independent trials of the following experiment:

- 1. We use a known device to draw a quantity from the *exponential* distribution (defined at the end of Section 2). We cannot observe this quantity directly, but it can cause effects that we can observe. So we give it a name *x*.
- 2. We use an unknown device to transform x to another quantity, which we do observe. The goal of the experiment is to find out what the unknown device does. Suppose we know that either the unknown device produces x as is, or it produces  $e^x 1$ .

We can model our two hypotheses about the unknown device by two generative stories whose outcomes are possible observations:

 $\begin{array}{l} e_1, e_2 :: Expr \ Real \\ e_1 = {\sf Let} "x" \ exponential \ ({\sf Var} "x") \\ e_2 = {\sf Let} "x" \ exponential \ ({\sf Add} \ ({\sf Exp} \ ({\sf Var} "x")) \ ({\sf Lit} \ (-1))) \end{array}$ 

Initially, say that we judge the two hypotheses to be equally probable. Then we start the experiment and the observations roll in:

obs :: [Real]obs = [3.07, 0.74, 2.23]

What do we believe now?

Our calculator finds the following densities for  $e_1$  and  $e_2$ :

$$d_1, d_2 :: Real \to Real$$
  

$$d_1 t = (if \ 0 < exp \ (-t) \land exp \ (-t) < 1 \text{ then } 1 \text{ else } 0)$$
  

$$\times exp \ (-t)$$

 $(\lambda x...)$ 



Figure 7. Likelihoods for two competing hypotheses. The crosses on the horizontal axis mark the *obs*erved outcomes.

$$d_2 t = if 0 < t - (-1) then d_1 (log (t - (-1)))/(t - (-1))$$
  
else 0

Figure 7 plots these densities. They are called *likelihoods* because they are densities of distributions over observations.

The likelihood  $d_1 t$  measures how likely it would be for us to observe *t* in a trial if the hypothesis  $e_1$  were true. Moreover, because the trials are independent, the likelihood of multiple observations is just the product of their likelihoods. Thus *product* (*map*  $d_1$  *obs*) measures how likely our *obs*ervations would be if the hypothesis  $e_1$  were true, and similarly for  $e_2$ . To compare the hypotheses against each other, we calculate the ratio of the likelihoods:

> product (map d<sub>1</sub> obs)/product (map d<sub>2</sub> obs) 1.2461022752116167

The likelihood ratio is above 1, which means the evidence favors our first hypothesis—that is, the unknown device produces x as is. We see this faintly in Figure 7: above where the crosses mark the *obs*ervations, the value of  $d_1$  tends to be greater than the value of  $d_2$ .

Whenever we have two or more hypotheses to choose from, the one with the greatest observation likelihood is called the *maximum-likelihood estimate* (MLE). So the MLE between  $e_1$  and  $e_2$  above is  $e_1$ . But we can also choose from an infinite family of hypotheses. In the experiment above for example, we can hypothesize instead that we observe  $a \times x$  in each trial, where a is an unobserved positive parameter that describes the unknown device. We can model this infinite family of hypotheses by an expression with a free variable "a":

$$e_{3} :: Expr Real$$

$$e_{3} = \text{Let "x" exponential (mul (Var "a") (Var "x"))}$$

$$mul :: Expr Real \rightarrow Expr Real \rightarrow Expr Real$$

$$mul a x = \text{Exp } (\text{Add } (\text{Log } a) (\text{Log } x)) \quad --a > 0 \land x > 0$$

Our calculator finds a density for  $e_3$  that simplifies algebraically to

$$d_3 :: Real \rightarrow Real \rightarrow Real$$
  
 $d_3 a = \lambda t$ . if  $t > 0$  then  $exp(-t/a)/a$  else 0

Figure 8 plots three members of this family of densities.

Using a bit of differential calculus, we can find the exact value of *a* that maximizes the likelihood *product* (*map* ( $d_3 a$ ) *obs*). That value is the MLE of the parameter *a*. Somewhat intuitively, it is the average of *obs*, represented by the solid line in Figure 8:

> **let** *a*<sub>MLE</sub> = *sum obs/fromIntegral* (*length obs*) > *a*<sub>MLE</sub> 2.013333333333333333

#### 7.2 Importance Sampling

We can also use densities for *importance sampling* (MacKay 1998). Suppose we have a density function *d* (called the *target*), and we



**Figure 8.** Likelihoods for three members of an infinite family  $d_3$  of competing hypotheses. The solid line is the MLE likelihood. The crosses on the horizontal axis mark the *observed* outcomes.

wish to sample repeatedly from the distribution with density d, perhaps to estimate the expectation of some function c with respect to the target distribution. Unfortunately, we don't know how to sample from the target distribution; in other words, we don't know any generative story with density d. We can pick a generative story  $e_1$  we do know (called the *proposal*), find a density  $d_1$  for  $e_1$ , and sample from  $e_1$  repeatedly instead. To correct for the difference between the target and proposal densities, we pair each outcome t from  $e_1$  with the *importance weight*  $d t/d_1 t$ .

importance\_sample :: 
$$(Real \rightarrow Real) \rightarrow IO (Real, Real)$$
  
importance\_sample  $d = \mathbf{do} \ t \leftarrow sample \ e_1 \ emptyEnv$   
return  $(t, d \ t/d_1 \ t)$ 

In particular, to estimate the expectation of the function c with respect to the target distribution, we compute the *weighted* average of c t where t is drawn from the proposal distribution.

 $\begin{array}{l} estimate\_expectation :: (Real \rightarrow Real) \rightarrow (Real \rightarrow Real) \\ \rightarrow IO \ Real \\ estimate\_expectation \ d \ c = \textbf{do} \\ samples \leftarrow replicateM \ 10000 \ (importance\_sample \ d) \\ return \ (sum \ [c \ t \times w \mid (t, w) \leftarrow samples] \\ \ /sum \ [w \mid (t, w) \leftarrow samples]) \end{array}$ 

For example, suppose we don't know how to sample from the target distribution with density

$$d:: Real \to Real$$
$$d \ t = exp \ (-t^3)$$

but we would like to estimate the expectation of *sin* with respect to it. We can use the definitions above to estimate the expectation:

> estimate\_expectation d sin 0.4508234334172205

# 8. Properties of Our Density Calculator

As explained at the beginning of Section 5, we want our density calculator to succeed as often as possible and to be compositional. Unfortunately, *density* does not succeed as often as we want. However, it can be made compositional.

#### 8.1 Incompleteness

As shown in Section 4, the distribution

Let "x" StdRandom (Add (Var "x") (Var "x"))

has a density function. In particular, it would be correct if

were to return the non-empty list

 $[\lambda \rho t. if 0 < t < 2 then 1/2 else 0]$ 

Nevertheless, our *density* function returns the empty list, because

$$usage (Add (Var "x") (Var "x")) "x" = Unknown density [Add (Var "x") (Var "x")] = []$$

and our code does not know  $x + x = 2 \times x$ .

This example shows there is room for our code to improve by succeeding more often. Transformations that reduce the usage of variables (for example, rewriting x + x to  $2 \times x$ ) would help, as would computer-algebra facilities for inverting a function that is expressed using non-invertible primitives (such as  $x^3 + x$ ). Unfortunately, those improvements would make it harder to keep a density calculator compositional and equationally derived.

Another source of incompleteness is demonstrated by the following example. Suppose *e* is some distribution expression that generates both positive and negative reals. For example, *e* could be Add *exponential* (Lit (-42)). We can express taking the absolute value of the outcome of *e*:

$$\begin{array}{l} e' = \mathsf{Let} "\mathtt{x}" \; e \; (\mathsf{If} \; (\mathsf{Less} \; (\mathsf{Lit} \; 0) \; (\mathsf{Var} \; "\mathtt{x}")) \\ (\mathsf{Var} \; "\mathtt{x}") \\ (\mathsf{Neg} \; (\mathsf{Var} \; "\mathtt{x}"))) \end{array}$$

If *e* has a density *d*, then e' has a fairly intuitive density d':

 $\begin{aligned} &d'::Real \rightarrow Real \\ &d' t = \mathbf{if} \ t > 0 \ \mathbf{then} \ d \ t + d \ (-t) \ \mathbf{else} \ 0 \end{aligned}$ 

But our calculator cannot find d', because the branches Var "x" and Neg (Var "x") have no density separately from the Let.

To handle these cases, it turns out that we can generalize our density calculator so that, when applied to Let "x" e (Var "x") or Let "x" e (Neg (Var "x")), it not only returns a density function but also *updates the environment*, mapping "x" to t or to -t respectively. The condition Less (Lit 0) (Var "x") can then be evaluated in the updated environment. In other words, it helps to generalize the environment to the heap of a lazy evaluator (Launchbury 1993), and to delay evaluating the condition.

## 8.2 Compositionality

As promised above Section 5.1, our definition of *density e* necessarily uses not only the *density* of the subexpressions of *e*, but also *expect*. But to handle Let, we strayed even further from perfect compositionality: our definition depends on substitution and *usage*, two more functions defined by structural induction on expressions. Can we still express *density* as a special case of a compositional and more general function, just as *mean* is a special case of the compositional and more general function *expect*? The answer turns out to be yes—we just need to rearrange the code already derived above. This is good news for people building a compiler from distributions to densities, including the present authors, because compositionality enables separate compilation.

If we had only used *expect* and *usage* to define *density*, it would have been straightforward to generalize *density* to a compositional function: just specify the omnibus interpretation *generalDensity* by

**data** GeneralDensity a = GD {  $gdExpect :: Env \rightarrow (a \rightarrow Real) \rightarrow Real,$   $gdUsage :: \forall b.Var b \rightarrow Usage,$   $gdDensity :: [Env \rightarrow a \rightarrow Real]$  }  $generalDensity :: Expr a \rightarrow GeneralDensity a$  generalDensity e = GD {gdExpect = expect e, gdUsage = usage e,gdDensity = density e }

and fuse it with our clauses defining *expect*, *usage*, and *density*, so as to define *generalDensity* e purely by structural induction

on *e*. For example, the new clause defining *generalDensity* on Add expressions would read

$$\begin{array}{l} generalDensity (\mathsf{Add} \ e_1 \ e_2) = GD \ \{\\ gdExpect \ = \ \lambda\rho \ c. \ gdExpect \ gd_1 \ \rho \ (\lambda x. \\ gdExpect \ gd_2 \ \rho \ (\lambda y. \ c \ (x+y))), \\ gdUsage \ = \ \lambda v. \ gdUsage \ gd_1 \ v \oplus \ gdUsage \ gd_2 \ v, \\ gdDensity \ = \ \left[\lambda\rho \ t. \ gdExpect \ gd_1 \ \rho \ (\lambda x. \ \delta_2 \ \rho \ (t-x))\right] \\ \ | \ \delta_2 \leftarrow \ gdDensity \ gd_2 ] \\ ++ \left[\lambda\rho \ t. \ gdExpect \ gd_2 \ \rho \ (\lambda y. \ \delta_1 \ \rho \ (t-y))\right] \\ \ | \ \delta_1 \leftarrow \ gdDensity \ gd_1 ] \} \\ \mathbf{where} \ gd_1 = \ generalDensity \ e_1 \\ \ gd_2 = \ generalDensity \ e_2 \end{array}$$

collecting the definition of *expect* (Add  $e_1 e_2$ ) in Section 3.1, the definition of *usage* (Add  $e_1 e_2$ ) in Appendix A, and the definition of *density* (Add  $e_1 e_2$ ) in Section 5.5. This is the *tupling transformation* (Pettorossi 1984; Bird 1980) applied to our *dependent interpretations* (Gibbons and Wu 2014, §4.2).

Our use of *density*  $(e' \{v \mapsto e\})$  to define *density* (Let v e e') in Section 5.6 complicates our quest for compositionality, because the recursive argument  $e' \{v \mapsto e\}$  is not necessarily a subexpression of Let v e e'. Instead of substituting e for v, we need the semantic analogue: some map, which we call *SEnv* for "static environment", that associates the variable v to the *expect* and *density* interpretations of e. We group these interpretations into a record type *General*. And instead of storing values in *Env* and renaming variables to avoid capture, we need the semantic analogue: storing values in lists, which we call *DEnv* for "dynamic environment", and allocating a fresh position in the lists for each variable. The type *General a* below contrasts with the type *GeneralDensity a* above.

$$\begin{array}{ll} \textbf{data } SEnv = SEnv \left\{ & \\ freshReal, freshBool :: Int, & \\ lookupSEnv & :: \forall a. Var \ a \rightarrow General \ a \\ \textbf{gExpect} & :: DEnv \rightarrow (a \rightarrow Real) \rightarrow Real, \\ gDensity & :: [DEnv \rightarrow a \rightarrow Real] \\ \end{array} \right\}$$
$$\begin{array}{ll} \textbf{data } DEnv = DEnv \left\{ & \\ lookupReal & :: [Real], \\ lookupBool & :: [Bool] \\ \end{array} \right\}$$

We call our omnibus interpretation *general*. It maps each distribution expression to its *usage* alongside a function from static environments to *expect* and *density* interpretations. The definition of *general* is mostly rearranging the code in Sections 3.1 and 5, so we relegate it to Appendix B.

general :: Expr 
$$a \rightarrow (\forall b. Var \ b \rightarrow Usage, SEnv \rightarrow General a)$$

At the top-level scope where the processing of a closed distribution expression commences, the static environment maps every variable name to an error and begins allocation at list position 0, matching the initially empty dynamic environment.

$$emptySEnv :: SEnv$$

$$emptySEnv = SEnv \{ freshReal = 0, freshBool = 0, lookupSEnv = \lambda v. error "Unbound" \}$$

$$emptyDEnv :: DEnv$$

$$emptyDEnv = DEnv \{ lookupReal = [], lookupBool = [] \}$$

We can finally define our density calculator as a special case of the compositional function *general*:

$$\begin{aligned} runDensity :: Expr \ a \to [a \to Real] \\ runDensity \ e &= [\delta \ emptyDEnv \\ & | \ \delta \leftarrow gDensity \ (snd \ (general \ e) \ emptySEnv)] \end{aligned}$$

# 9. Conclusion

We have turned a specification of density functions in terms of expectation functionals into a syntax-directed implementation that supports separate compilation. Our equational derivation draws from algebra, integral calculus, and  $\lambda$ -calculus. It suggests that program calculation and transformation are powerful ways to turn expressive probabilistic models into effective inference procedures. We are investigating this hypothesis in ongoing work.

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# A. Usage Testing

Section 5.6 uses the function

 $usage :: Expr \ a \rightarrow Var \ b \rightarrow Usage$ 

to test how many times an expression uses a variable. The return type *Usage* offers just three possibilities:

**data** Usage = Never | AtMostOnce | Unknown **deriving** (Eq, Ord)

For example, we want

usage (If (Var "b") (Var "x") (Var "x")) "x" = AtMostOnceusage (Add (Var "x") (Var "x")) "x" = Unknown

This contrast between If and Add indicates that we need two algebraic structures on the type *Usage*.

First, some Usage values entail others as propositions. For example, if v is never used, then v is used at most once. This entailment relation just happens to be a total order, so we define the operator  $\leq$  to mean entailment, by **deriving** Ord above.

Second, when two subexpressions together produce a final outcome, the counts of how many times they use v add up, and our knowledge of the counts forms a commutative monoid. For example, suppose  $e' = \text{Add } e'_1 e'_2$ , and we know that  $e'_1$  never uses v and  $e'_2$  uses v at most once. Then we know that e' uses v at most once. If instead we only know that  $e'_1$  and  $e'_2$  each use v at most once, then all we know about e' is it uses v at most twice. That's not useful knowledge about e', so we might as well represent it as *Unknown*. We define the operator  $\oplus$  to add up our knowledge in this way:

#### instance Monoid Usage where

тетр	ty	= Never		
Never	$\oplus u$	=	и	
и	$\oplus$ Never	=	и	
_	⊕_	=	Unknown	

Armed with these two instances, we can define the *usage* function. It amounts to an abstract interpretation of expressions:

usage StdRandom	$_{-} = Never$
$usage$ (Lit _)	$_{-} = Never$
usage (Var v)	v' = <b>if</b> eq $v v'$ <b>then</b> AtMostOnce
	else Never
usage (Let $v e e'$ )	$v' = usage \ e \ v' \oplus \mathbf{if} \ eq \ v \ v' \ \mathbf{then} \ Never$
	else usage e' v'
usage (Neg $e$ )	$v = usage \ e \ v$
usage (Exp e)	$v = usage \ e \ v$
usage (Log e)	$v = usage \ e \ v$
usage (Not $e$ )	$v = usage \ e \ v$
usage (Add $e_1 e_2$ )	$v = usage e_1 v \oplus usage e_2 v$

 $usage (Less e_1 e_2) v = usage e_1 v \oplus usage e_2 v$  $usage (If e e_1 e_2) v = usage e v \oplus max (usage e_1 v)$  $(usage e_2 v)$ 

The Var and Let cases above use the function *eq* to test the equality of two *Vars* whose types might differ:

 $\begin{array}{l} eq :: Var \ a \to Var \ b \to Bool \\ eq \ (\mathsf{Real} \ v) \ (\mathsf{Real} \ w) = v \equiv w \\ eq \ (\mathsf{Bool} \ v) \ (\mathsf{Bool} \ w) = v \equiv w \\ eq \ - \qquad = False \end{array}$ 

To fit Haskell's type system better, we distinguish variables whose names are the same *String* if their types differ. For example, *extendEnv* in Section 2 treats Real "x" and Bool "x" as different variables that do not shadow each other's bindings. In other words, *Real* and *Bool* variables in our language reside in separate namespaces. Hence *eq* (Real "x") (Bool "x") = *False*.

## **B.** Compositional Density Calculator

```
extendSEnv :: Var a \rightarrow General a \rightarrow SEnv \rightarrow SEnv
extendSEnv v x \sigma = \sigma {
   lookupSEnv = extendSEnv' v x (lookupSEnv \sigma)
extendSEnv' :: Var a \rightarrow General a \rightarrow (\forall b. Var b \rightarrow General b)
                                              \rightarrow (\forall b. Var \ b \rightarrow General \ b)
extendSEnv' (Real v) x = (\text{Real } v') | v \equiv v' = x
extendSEnv' (Bool v) x = (Bool v') | v \equiv v' = x
extendSEnv' _
                             -\sigma v'
                                                           = \sigma v'
extendList :: Int \rightarrow a \rightarrow [a] \rightarrow [a]
extendList i x xs
    |i \equiv length xs = xs + [x]
    | otherwise = error ("Expected length " ++ show i ++
                                    ", got " ++ show (length xs))
generalReal :: (DEnv \rightarrow Real) \rightarrow GeneralReal
generalReal f = General \{
   gExpect = \lambda \rho c.c (f \rho),
   gDensity = [] \}
generalBool :: (DEnv \rightarrow (Bool \rightarrow Real) \rightarrow Real) \rightarrow GeneralBool
generalBool \ e = General \{
   gExpect = e,
   gDensity = [\lambda \rho \ t. e \ \rho \ (\lambda x. \text{ if } t \equiv x \text{ then } 1 \text{ else } 0)] \}
allocate :: Var a \rightarrow SEnv \rightarrow (SEnv, a \rightarrow DEnv \rightarrow DEnv)
allocate v@(Real _) \sigma =
   let i = freshReal \sigma
   in (extendSEnv v (generalReal (\lambda \rho.lookupReal \rho !! i))
                        \sigma {freshReal = i+1},
        \lambda x \rho . \rho \{ lookupReal = extendList i x (lookupReal \rho) \} \}
allocate v@(Bool_) \sigma =
   let i = freshBool \sigma
   in (extendSEnv v (generalBool (\lambda \rho c.c (lookupBool \rho !! i)))
                        \sigma \{ freshBool = i+1 \},\
        \lambda x \rho. \rho \{ lookupBool = extendList i x (lookupBool \rho) \} \}
general StdRandom = (\lambda_{-}.Never,
   \lambda_{-}. General {
   gExpect = \lambda_{-}c.\int_{0}^{1}\lambda x.c x,
   gDensity = [\lambda_t \cdot \mathbf{if} \ 0 < t \land t < 1 \ \mathbf{then} \ 1 \ \mathbf{else} \ 0] \}
general (Lit x) = (\lambda_...Never,
```

 $\lambda_{-}$ .generalReal ( $\lambda_{-}$ .fromRational x))

```
\lambda \sigma. General {
   gExpect = \lambda \rho c. gExpect (g \sigma) \rho (\lambda x. c (log x)),
   gDensity = [\lambda \rho t. \delta \rho (exp t) \times exp t | \delta \leftarrow gDensity (g \sigma)] \})
   where (u,g) = general e
general (Not e) = (u,
   \lambda \sigma. generalBool (\lambda \rho c. gExpect (g \sigma) \rho (\lambda x. c (not x))))
   where (u,g) = general e
general (Add e_1 e_2) = (\lambda v. u_1 v \oplus u_2 v,
   \lambda \sigma. General {
   gExpect = \lambda \rho c. gExpect (g_1 \sigma) \rho (\lambda x.
                              gExpect (g_2 \sigma) \rho (\lambda y. c (x+y))),
   gDensity = [\lambda \rho t. gExpect (g_1 \sigma) \rho (\lambda x. \delta_2 \rho (t-x))]
                       |\delta_2 \leftarrow gDensity(g_2 \sigma)|
                 ++ [\lambda \rho t. gExpect (g_2 \sigma) \rho (\lambda y. \delta_1 \rho (t-y))]
                       |\delta_1 \leftarrow gDensity(g_1 \sigma)]
   where (u_1, g_1) = general e_1
             (u_2, g_2) = general e_2
general (Less e_1 e_2) = (\lambda v. u_1 v \oplus u_2 v,
   \lambda \sigma. generalBool (\lambda \rho c. gExpect (g_1 \sigma) \rho (\lambda x.
                                      gExpect (g_2 \sigma) \rho (\lambda y. c (x < y))))
   where (u_1, g_1) = general e_1
              (u_2, g_2) = general e_2
general (If e e_1 e_2) = (\lambda v. u v \oplus max (u_1 v) (u_2 v),
   \lambda \sigma. General {
   gExpect = \lambda \rho c. gExpect (g \sigma) \rho (\lambda b.
                              gExpect ((if b then g_1 else g_2) \sigma) \rho c),
   gDensity = [\lambda \rho \ t. gExpect \ (g \ \sigma) \ \rho \ (\lambda b.
                               (if b then \delta_1 else \delta_2) \rho t)
                       |\delta_1 \leftarrow gDensity(g_1 \sigma), \delta_2 \leftarrow gDensity(g_2 \sigma)]\})
   where (u, g) = general e
              (u_1,g_1) = general e_1
              (u_2, g_2) = general e_2
```

general (Var v) =  $(\lambda v')$  if eq v v' then AtMostOnce else Never,

general (Let v e e') =  $(\lambda v' . u v' \oplus if eq v v'$  then Never else u' v',

gExpect  $(g' \sigma') (\varepsilon x \rho) c)$ ,

 $\sigma'' = extendSEnv v (g \sigma) \sigma$  in General {

 $gDensity = [\lambda \rho t. gExpect (g \sigma) \rho (\lambda x. \delta' (\varepsilon x \rho) t)]$ 

 $++ [\delta' | u' v \leq AtMostOnce$ 

 $gExpect = \lambda \rho \ c. gExpect \ (g \ \sigma) \ \rho \ (\lambda x. c \ (-x)),$  $gDensity = [\lambda \rho \ t. \delta \ \rho \ (-t) \ | \ \delta \leftarrow gDensity \ (g \ \sigma)] \})$ 

gExpect =  $\lambda \rho c.gExpect (g \sigma) \rho (\lambda x.c (exp x)),$ 

 $gDensity = [\lambda \rho t. if 0 < t then \delta \rho (log t)/t else 0$ 

 $|\delta \leftarrow gDensity(g\sigma)]\})$ 

 $|\delta' \leftarrow gDensity(g'\sigma')|$ 

,  $\delta' \leftarrow gDensity(g' \sigma'')$ })

 $\lambda \sigma$ . lookupSEnv  $\sigma v$ )

 $\lambda \sigma$ . let  $(\sigma', \varepsilon) = allocate \ v \ \sigma$ 

where (u, g) = general e

where (u,g) = general egeneral (Exp e) = (u,

where (u,g) = general e

general (Neg e) = (u,

 $\lambda \sigma$ . General {

 $\lambda \sigma$ . General {

general (Log e) = (u,

(u',g') = general e'

gExpect =  $\lambda \rho c.gExpect (g \sigma) \rho (\lambda x.$