Using RcppTN in R and C++

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This brief document shows simple usage of the function rtn() provided by the R package RcppTN for drawing from an arbitrary sequence of truncated Normal distributions. Much of the value added by the RcppTN package comes from providing a C++-level API to call in development of other Rcpp-based C++ codes. Use of this API is also demonstrated. While no other R packages currently provide this functionality in an API, some existing implementations for drawing from a truncated Normal distribution at the R-level include truncnorm and msm.

1 Installation

Currently, there is no CRAN version of the package, so the simplest installation mechanism is using the install_github() function from the devtools package.

```
library(devtools)
install_github("RcppTN", username = "olmjo", subdir = "pkg")
```

2 R-level Usage

Usage of the rtn() function in R is straightforward (albeit not feature-rich). Without any options, we get a single draw from the standard Normal distribution. And, this draw respects R's RNG state so the stream of output is reproducible.

```
library(RcppTN)

## Loading required package: Rcpp

set.seed(1)
rtn()

## [1] -0.6265

set.seed(1)
rtn()

## [1] -0.6265
```

Under this implementation of the Robert (1995) algorithm, a request for a single draw from a Standard Normal distribution truncated from $-\infty$ to ∞ — the default behavior of the function when called without any arguments — results in the same return value as a single draw from a Standard Normal distribution using rnorm(). This is just a by-product of the implementation and holds no practical significance.¹

```
set.seed(1)
rtn()

## [1] -0.6265

set.seed(1)
rtn(.mean = 0, .sd = 1, .low = -Inf, .high = Inf)

## [1] -0.6265

set.seed(1)
rtn()

## [1] -0.6265

set.seed(1)
rnorm(1)

## [1] -0.6265
```

Of course, rtn()'s behavior given RNG seeds is exactly as you would expect for any other generator in **R**.

```
set.seed(11)
rtn()
## [1] -0.591
rtn()
## [1] 0.02659
set.seed(1)
rtn()
## [1] -0.6265
rtn()
## [1] 0.1836
set.seed(11)
rtn()
```

¹See the R package documentation for the citation to the algorithm.

```
## [1] -0.591

rtn()

## [1] 0.02659
```

In practice, this R-level function will likely be used in one of two ways:

- 1. drawing many values from the same truncated Normal distribution
- 2. drawing many values from different truncated Normal distributions

For the rtn() function, these two uses look very similar. The function accepts a .mean argument, an .sd argument, a .low argument, and a .high argument. Each should be a vector of length K corresponding to the K distributions of interest. The function does not handle value recycling for the user, so the construction of these vectors must be done before or during the call of the rtn() function. Importantly, this function quietly returns an NA value for invalid input parameters. NA-inducing input parameters don't interfere with other valid parameters and a vector of the requested length is returned.

For example,

```
rtn(0, -1, 0, 1)

## Warning: NAs returned. Check for invalid parameters.

## [1] NA

rtn(0, 1, 0, -1)

## Warning: NAs returned. Check for invalid parameters.

## [1] NA

rtn(c(0,0), c(1,1), c(0,0), c(-Inf,Inf))

## Warning: NAs returned. Check for invalid parameters.

## [1] NA 1.178
```

2.1 Multiple Draws from a Single Distribution

Multiple draws from the same distribution may be requested with a function call like the following:

```
## [1] 1000

mean(output)

## [1] 1.389
```

Here, we are generating 1,000 draws, with each draw, x, coming from N(0,1) truncated below at 1 and above at 2. The population mean of this distribution is

$$E[x] = \mu + \frac{\phi(\frac{a-\mu}{\sigma}) - \phi(\frac{b-\mu}{\sigma})}{\Phi(\frac{b-\mu}{\sigma}) - \Phi(\frac{a-\mu}{\sigma})} \cdot \sigma,$$

where $\mu = 0$, $\sigma = 1$, ϕ denotes the pdf of the standard Normal distribution, Φ denotes the standard cdf of the standard Normal distribution, and a and b are the lower and upper bounds of truncation, respectively. So, for the above parameter values we have

$$E[x] = \mu + \frac{\phi(\frac{a-\mu}{\sigma}) - \phi(\frac{b-\mu}{\sigma})}{\Phi(\frac{b-\mu}{\sigma}) - \Phi(\frac{a-\mu}{\sigma})} \cdot \sigma$$
$$= 0 + \frac{.242 - .054}{.977 - .841} \cdot 1$$
$$\approx 1.383$$

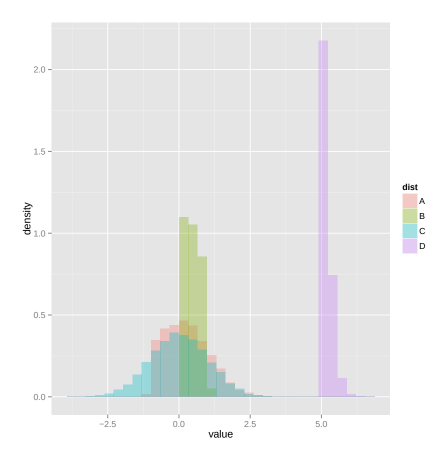
Our sample mean for the 1,000 draws (1.389) is close to the population mean (1.383). To get a better sense of how dispersed the sampling distribution for the mean of a sample of 1,000 draws from this distribution is, we can simulate it.

```
bigoutput <- rep(NA, 1000)
for (i in 1:length(bigoutput)) {
    bigoutput[i] <- mean(rtn(.mean = rep(0, 1000),</pre>
                               .sd = rep(1, 1000),
                               .low = rep(1, 1000),
                               .high = rep(2, 1000)
                          )
summary(bigoutput)
##
      Min. 1st Qu.
                     Median
                               Mean 3rd Qu.
                                                Max.
##
           1.38
                       1.38
                               1.38
                                     1.39
                                                1.41
```

Looking at the summary of the sample means, we see that the sampling distribution of sample means is centered directly on the population mean.

As shown above, valid input for rtn() includes -Inf and Inf. Below are histograms for four different truncated Normal distributions. The rtn() function works perfectly well in simulating draws from regions that have a low (read nearly 0) density in a non-truncated Normal distribution. Distribution "D" is an example of this.

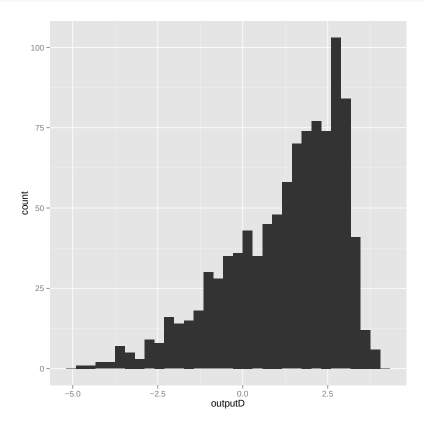
```
library(ggplot2)
outputA \leftarrow rtn(.mean = rep(0, 5000),
               .sd = rep(1, 5000),
                .low = rep(-1, 5000),
                .high = rep(Inf, 5000)
outputB \leftarrow rtn(.mean = rep(0, 5000),
               .sd = rep(1, 5000),
                .low = rep(0, 5000),
                .high = rep(1, 5000)
outputC \leftarrow rtn(.mean = rep(0, 5000),
                .sd = rep(1, 5000),
                .1ow = rep(-Inf, 5000),
                .high = rep(Inf, 5000)
outputD <- rtn(.mean = rep(0, 5000),
                .sd = rep(1, 5000),
                .low = rep(5, 5000),
                .high = rep(Inf, 5000)
dfOutput <- rbind(data.frame(value = outputA, dist = "A"),</pre>
                   data.frame(value = outputB, dist = "B"),
                   data.frame(value = outputC, dist = "C"),
                   data.frame(value = outputD, dist = "D")
ggplot(dfOutput) +
    geom_histogram(aes(x=value,
                        y=..density..,
                        fill = dist
                        ),
                    alpha = 1/3,
                    position = "identity"
```



2.2 Multiple Draws from Different Distributions

Taking multiple draws from different distributions proceeds in a similar way, though the construction of the arguments passed to rtn() changes a bit. If we were interested in characterizing a distribution of draws from a truncated Normal distribution where one (or more) of the parameters was, itself, stochastic, rtn() could easily be put to use. Here, the vector of lower bounds and upper bounds are each the result of an rtn() function call (notice that a < b by construction).

Then, we cam sample 1,000 draws from this truncated Normal distribution of interest where the mean and standard deviation are fixed, but the bounds of truncation, themselves, are taken from a distribution (in this case, a truncated Normal distribution).



This sampling distribution is non-standard and the easiest way to characterize it would be through a simulation like the above.

3 C++-level Usage

This section documents how to use the C++-level functionality in subsequent C++ development. Specifically, using the RcppTN C++ API via sourceCpp() and an Rcpp-based R package are shown. Presently, only two functions are exposed at the C++ level: rtn1() and etn1().

```
rtn1
```

```
double rtn1(double mean, double sd, double low, double high);
etn1
double etn1(double mean, double sd, double low, double high);
```

Caveats. The R-level function ultimately calls these C++-level functions. So, all of the features of the R-level function apply here (e.g., respecting R' RNG state). However, as is true in **Rcpp**, this is left to the user to enforce. No checking or error handling is provided with these functions. These functions live in the RcppTN namespace.

3.1 Examples

Via sourceCpp(). In non-package R code, use is very straightforward due to the mechanisms provided by Rcpp. Include the appropriate header file as you would for Rcpp. In addition, use the depends pseudo-attribute with "// [[Rcpp::depends(RcppTN)]]" to ensure that linker finds the symbols. From there, use is as you would expect.

```
library(Rcpp)
sourceCpp(code = "
#include <Rcpp.h>
#include <RcppTN.h>
// [[Rcpp::depends(RcppTN)]]
using namespace Rcpp;
// [[Rcpp::export]]
List rcpp_hello_world() {
 double a = RcppTN::rtn1(0.0, 1.0, 3.5, 3.7);
 double b = RcppTN::etn1(0.0, 1.0, 3.5, 3.7);
 NumericVector y = NumericVector::create( a, b ) ;
 List z = List::create( y ) ;
 return(z);
}
H
rcpp_hello_world()
## [[1]]
## [1] 3.570 3.588
```

Via an Rcpp-based package. In R, use Rcpp.package.skeleton() from Rcpp to create an empty, but functional, R package.

```
library(Rcpp)
Rcpp.package.skeleton(path = "~/Desktop")
```

Navigate inside the newly created anRpackage directory and edit the DESCRIPTION file. Add RcppTN to the Depends: and LinkingTo: lines of the file as in

```
Depends: Rcpp (>= 0.10.5), RcppTN LinkingTo: Rcpp, RcppTN
```

Now, edit the C++ function rcpp_hello_world() in anRpackage/src/rcpp_hello_world.cpp to read

```
#include <Rcpp.h>
#include <RcppTN.h>

using namespace Rcpp;

// [[Rcpp::export]]
List rcpp_hello_world() {
  double a = RcppTN::rtn1(0.0, 1.0, 3.5, 3.7);
  double b = RcppTN::etn1(0.0, 1.0, 3.5, 3.7);
  NumericVector y = NumericVector::create(a, b);
  List z = List::create(y);
  return(z);
}
```

To see the effect of this, install the "anRpackage" package and the load it in R. From there, make subsequent calls to the rcpp_hello_world() function. With a similar approach, the rtn1() function can be called in a more useful way within other C++-level codes without the need for re-coding the wheel. The only difference between this approach and the sourceCpp() approach is that the depends pseudo-attribute is no longer needed and is replaced by the modification to the LinkingTo: field of the DESCRIPTION file.