Homework 3

DATA604 Simulation and Modeling

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1

Starting with $X_0 = 1$, write down the entire cycle for $X_i = 11X_{i-1} \mod(16)$

```
fn1 <- function(x0)
{
    df <- data.frame(X=c(), R=c())
    x <- x0
    continue <- TRUE

while(continue)
{
    xi <- (11 * x) %% 16
    df <- rbind(df, data.frame(X=x, R=xi))
    x <- xi

    if(xi == x0)
    {
        break
    }
}

return(df)
}</pre>
```

X	R
1	11
11	9
9	3
3	1

 $\mathbf{2}$

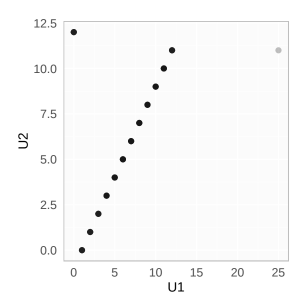
Using the LCG provided below: $X_i = (X_{i-1} + 12) mod(13)$, plot the pairs $(U_1.U_2), (U_2, U_3), ...$ and observe the lattice structure obtained. Discuss what you observe.

```
fn2 <- function(x0, max=100)
{
    df <- data.frame(U1=c(), U2=c())
    x <- x0

    for(i in 1:max)
    {
        xi <- (x + 12) %% 13
        df <- rbind(df, data.frame(U1=x, U2=xi))</pre>
```

```
x <- xi
}

return(df)
}
# Call the function starting at x0=1
res <- fn2(25)</pre>
```



The chart above suggests there are only 13 points, but actually the LCG cycle period is 13 and numbers are repeating.

U2
11
10
9
8
7
6
5
4
3
2
1
0
12
11

3

Implement the pseudo-random number generator:

$$X_i = 16807X_{i-1} \bmod (2^{31} - 1)$$

Using the seed 1234567, run the generator for 100,000 observations. Perform a chi-square goodness-of-fit test on the resulting PRN's. Use 20 equal-probability intervals and level $\alpha=0.05$. Now perform a runs up-and-down test with $\alpha=0.05$ on the observations to see if they are independent.

```
fnLCG3 <- function(seed = 1, n = 1)
{</pre>
```

```
rands <- rep(NA, n)
x <- seed
modVal <- (2^31 - 1)

for(i in 1:n)
{
    xi <- (16807 * x) %% (modVal)
    rands[i] <- xi
    x <- xi
}

return(rands)
}

n=100000
rn <- fnLCG3(1234567, n)</pre>
```

The first 6 generated numbers are shown below:

Chi-Square Test

```
intervals <- 20
maxRn <- max(rn)</pre>
minRn <- min(rn)
intWidth <- (maxRn - minRn) / intervals</pre>
lwr <- minRn</pre>
dfCounts <- data.frame(intID=c(), count=c())</pre>
# Bin the data ourselves, I'd quess there
# is an easier way, but this will do.
for(i in 1:intervals)
  upr <- lwr + intWidth</pre>
  inRange <- rn[lwr <= rn & rn < upr]</pre>
  dfCounts <- rbind(dfCounts, data.frame(intID=i, count=length(inRange)))</pre>
  # setup for next interval range
  lwr <- upr</pre>
}
# Do our own Chi-Squared test
Expected <- (100000 / intervals)
chi2 <- sum((dfCounts$count - Expected)^2 / Expected)</pre>
chi2
## [1] 14.7762
# Use built-in function to compare
chiTest <- chisq.test(dfCounts$count)</pre>
chiTest
```

```
## Chi-squared test for given probabilities
##
## data: dfCounts$count
## X-squared = 14.776, df = 19, p-value = 0.7367
```

The p-value = 0.7367029 is not less than $\alpha = 0.05$, therefore we don't reject the null hypothesis that the distribution is uniform.

intID	count
1	5069
2	5028
3	5044
4	5087
5	4948
6	4953
7	4937
8	4933
9	4900
10	4957
11	5088
12	4994
13	5076
14	5019
15	5002
16	5067
17	4981
18	4914
19	5062
20	4940

Runs Up-and-Down Test

Using the tseries package, we execute the Runs test (Trapletti and Hornik, 2015). First we have to construct the +/- vector. Here we simply convert to boolean.

```
s <- rep(NA, n - 1)
for(i in 1:n - 1)
{
    s[i] <- rn[i] < rn[i + 1]
}
runsTest <- runs.test(as.factor(s))
runsTest</pre>
```

```
##
## Runs Test
##
## data: as.factor(s)
## Standard Normal = 105.84, p-value < 2.2e-16
## alternative hypothesis: two.sided</pre>
```

Based on the p-value < 0.05, we reject the null hypothesis and conclude there is not evidence to support independence in the psuedo-random numbers.

4.

Give inverse-transforms, composition, and acceptance-rejection algorithms for generating from the following density:

$$f(x) = \begin{cases} \frac{3x^2}{2} & -1 \le x \le 1\\ 0 & otherwise \end{cases}$$

Inverse-Transforms

First find the indefinite integral of the probability density function:

$$F(x) = \int \frac{3x^2}{2} dx = \frac{x^3}{2}$$

Next set F(x) = R and solve for x in terms of R:

$$\frac{x^3}{2} = R$$

$$x^3 = 2R$$

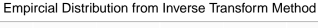
$$F^-1(x) = \sqrt[3]{2R}$$

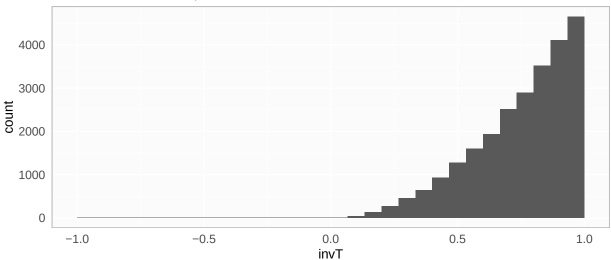
```
# Define a function of the F^-1(X)
invTfn4 <- function(r)
{
   vals <- (2 * r)^(1/3)
   return (vals)
}

# Generate the uniform psuedo-random vars
rVals <- runif(n, -1, 1)
# Convert to the desired distribution using the inverse transform method.
invTVals <- invTfn4(rVals)</pre>
```

Warning: Removed 74997 rows containing non-finite values (stat_bin).

Warning: Removed 2 rows containing missing values (geom_bar).

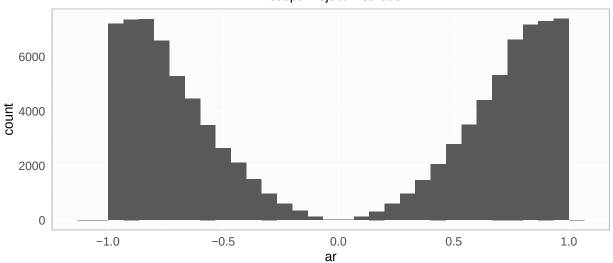




```
# Helper function for the Accept/Reject approach
myRARmethod <- function(fun, min, max)</pre>
{
  M <- 2
  accepted <- FALSE
  while(!accepted)
    # Get a random value from uniform distribtion (g(x) \text{ for } us)
    r <- runif(1, min, max)
    # Sample x from g(x) and u from U(0,1) (the uniform distribution over the unit interval)
    u <- runif(1, 0, 1)
    gx <- dunif(r, min, max)</pre>
    # Check whether or not u < f(x)/Mg(x).
    if(u < fun(r) / (M * gx))
    {
      accepted = TRUE
    }
  }
  return(r)
}
# Define a function for the PDF
Arfn4 <- function(x)</pre>
  if(-1 \le x \&\& x \le 1)
    val <- (3 * x^2) / 2
  }
  else
  {
    val = 0
  }
  return (val)
}
# Loop to generate the values
rarVals <- rep(NA, n)
for(i in 1:n)
  rarVals[i] <- myRARmethod(Arfn4, -1, 1)</pre>
}
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.





Composition

 $\operatorname{Hmm}.\dots$

5

Implement, test and compare different methods to generate from a N(0,1) distribution.

$$f(x) = \frac{1}{\sqrt{2\pi}}e^{\frac{-x^2}{2}}$$

Inverse Transform Method

$$F(X) = \int \frac{1}{\sqrt{2\pi}} e^{\frac{-x^2}{2}} dx$$

It turns out the qnorm function is the inverse normal CDF function, so we'll use it.

```
normrandit <- function()
{
    u <- runif(1)
    return (qnorm(u))
}

itstats <- function(n)
{
    vals <- rep(NA, n)
    for(i in 1:n)
    {
      vals[i] <- normrandit()
    }

    return(list(values=vals, mean=mean(vals), sd=sd(vals)))
}

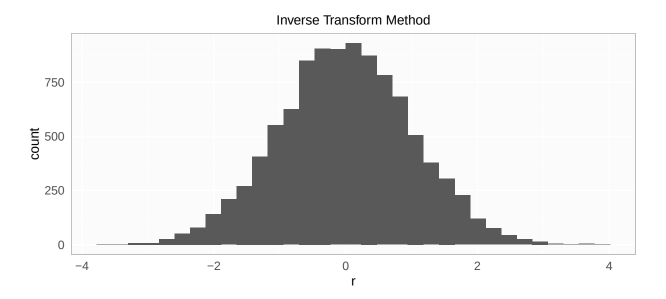
res <- itstats(10000)
res$mean</pre>
```

[1] -0.005543553

res\$sd

```
## [1] 1.003801
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



Box-Muller Method

```
normrandbm <- function()</pre>
  u2 <- runif(2)
  v1 \leftarrow (-2 * \log(u2[1]))^(1/2) * \cos(2 * pi * u2[2])
  v2 \leftarrow (-2 * \log(u2[1]))^(1/2) * \sin(2 * pi * u2[2])
  return (c(v1, v2))
}
bmstats <- function(n)</pre>
  vals \leftarrow rep(NA, n / 2)
  for(i in seq(1, n, by=2))
    rs <- normrandbm()
     vals[i] <- rs[1]</pre>
     vals[i + 1] \leftarrow rs[2]
  return(list(values=vals, mean=mean(vals), sd=sd(vals)))
}
resBm <- bmstats(10000)</pre>
length(resBm[[1]])
```

[1] 10000

```
# Mean
```

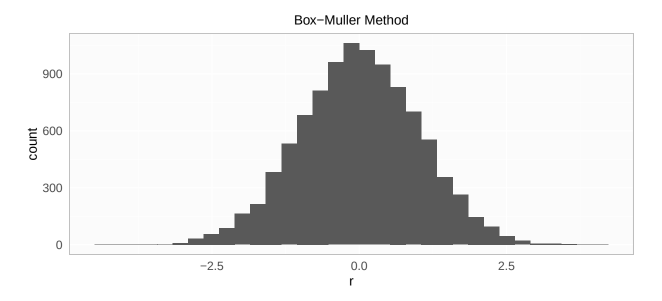
resBm\$mean

```
## [1] 0.00486827
```

SD

resBm\$sd

[1] 1.009415



Accept Reject Method

```
normrandar <- function()</pre>
{
  continue <- TRUE
  while(continue)
    u2 <- runif(2)
    eu2 <- - log(u2)
    if(eu2[2] >= ((eu2[1] - 1)^2 / 2))
    {
      break
  }
  sign <- runif(1)</pre>
  if(sign > 0.5)
    eu2[1] \leftarrow eu2[1] * -1
  }
  return(eu2[1])
}
arstats <- function(n)</pre>
```

```
vals <- rep(NA, n)
for(i in seq(1, n))
{
    rs <- normrandar()
    vals[i] <- rs
    #vals[i + 1] <- rs[2]
}
return(list(values=vals, mean=mean(vals), sd=sd(vals)))
}</pre>
```

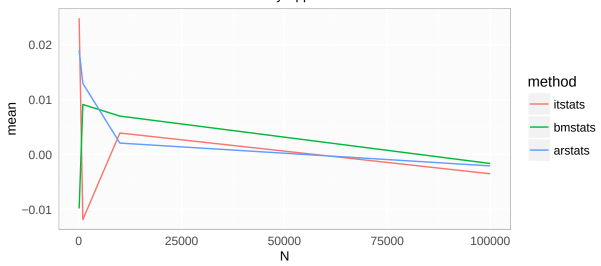
Compare

```
dfCompare <- data.frame(method=c(), N=c(), mean=c(), sd=c(), time=c())</pre>
perfTest <- function(fun)</pre>
{
  dfResult <- data.frame(method=c(), N=c(), mean=c(), sd=c(), time=c())</pre>
  Ns <- c(100, 1000, 10000, 100000)
  it <- 2
  for(n in Ns)
    m <- rep(NA, it)
    s <- rep(NA, it)
    t <- rep(NA, it)
    for(i in 1:it)
      st <- system.time({ret <- fun(n)})</pre>
      #print(st)
      m[i] <- ret$mean
      s[i] <- ret$sd
      t[i] <- st[[3]]
    }
    dfResult <- rbind(dfResult, data.frame(method=deparse(substitute(fun)),</pre>
                                              N=n,
                                              mean=mean(m),
                                              sd=mean(s),
                                              time=mean(t)))
  }
  return (dfResult)
}
# Force less use of scientific notation
\# http://stackoverflow.com/questions/9397664/force-r-not-to-use-exponential-notation-e-g-e10
options("scipen"=100, "digits"=5)
dfCompare <- rbind(dfCompare, perfTest(itstats))</pre>
dfCompare <- rbind(dfCompare, perfTest(bmstats))</pre>
dfCompare <- rbind(dfCompare, perfTest(arstats))</pre>
```

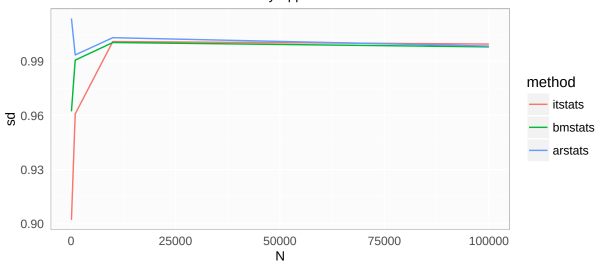
method	N	mean	sd	$_{ m time}$
itstats	100	0.02481	0.90215	0.000
itstats	1000	-0.01185	0.96086	0.020
itstats	10000	0.00391	1.00094	0.110
itstats	100000	-0.00350	0.99954	1.240

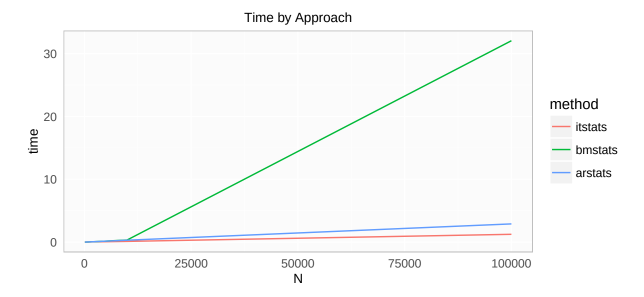
method	N	mean	sd	time
bmstats	100	-0.00982	0.96226	0.005
bmstats	1000	0.00911	0.99064	0.015
bmstats	10000	0.00700	1.00040	0.335
bmstats	100000	-0.00163	0.99798	32.060
arstats	100	0.01899	1.01370	0.000
arstats	1000	0.01295	0.99351	0.030
arstats	10000	0.00208	1.00308	0.300
arstats	100000	-0.00207	0.99842	2.895

Means by Approach



Stdev by Approach





For me, the Inverse Transform approach takes the least time, but that might be due to the qnorm optimizations.

Million Sample Histograms

```
m <- 1000000
itMil <- itstats(m)
bmMil <- bmstats(m)
arMil <- arstats(m)</pre>
```

References

Trapletti, A. and K. Hornik. tseries: Time Series Analysis and Computational Finance. R package version 0.10-34. 2015. URL: http://CRAN.R-project.org/package=tseries.