STAT 242 Final Project: Adaptive Rejection Sampling

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In this project we implement the adaptive rejection sampling algorithm proposed by Gilks et al. (1995), a method for rejection sampling from any univariate log-concave probability density function f(x). This method is particularly useful in situations where the density function is computationally expensive to evaluate, as it uses successive evaluations of $h(x) = \log(f(x))$ to improve the approximation of a rejection envelope u(x) and squeezing function l(x), which are (theoretically) much simpler to compute and sample from than the original density function.

The R package, ars, containing our solution can be found using the following link:

https://github.com/jtmorrell/ars

Or, can be installed using devtools:

```
devtools::install_github('jtmorrell/ars')
```

Methodology

Our implementation consists of one function, ars, which takes as arguments the univariate log-concave density function f(x), the number of samples desired N, the initial points x_0 used to approximate $h(x) = \log(f(x))$ (default: c(-1, 1)), and optionally the bounds of the density function.

To demonstrate the implementation of ars we will go through the example of sampling 100000 points from the normal distribution.

```
## Normally inputs to ars()
f <- dnorm
N <- 100000
x0 <- c(-1.0, 1.0)
bounds <- c(-Inf, Inf)</pre>
```

First we check the inputs.

```
## Input sanity check
if (!is.function(f)) {
    stop('f is not a function.')
}

if (length(bounds) != 2) {
    stop('Length of bounds must be 2! (upper and lower bounds required)')
}

if (bounds[1] == bounds[2]) {
    warning('Same upper and lower bounds, replacing them by default: bounds=c(-Inf, Inf)')
    bounds <- c(-Inf, Inf)
}

if (!all((x0 > bounds[1]) & (x0 < bounds[2]))) {
    stop('Bad inputs: x0 must be inside bounds.')
}</pre>
```

Then we'll initialize some helper variables.

```
## define h(x) = log(f(x))
h <- function(x) {
   return (log(f(x)))
}

## helper variables
dx <- 1E-8  # mesh size for finite difference approximation to h'(x)
max_iters <- 10000  # prevent infinite loop
current_iter <- 0
bds_warn <- FALSE  # Flag for boundary warning in main while loop</pre>
```

Because the derivative of the log-density function h'(x) is required in our calculation, we need to approximate it using a finite-difference method. We'll use a forward differencing approximation, as we only need to evaluate h(x) one additional time to calculate h'(x). The forward difference method uses the following Taylor expansion

$$f(x_0 + h) = f(x_0) + f'(x_0)h + \frac{1}{2}f''(x_0)h^2 + \mathcal{O}(h^3)$$

to approximate the first derivative as

$$f'(x_0) \approx \frac{f(x_0 + h) - f(x_0)}{h}$$

where h is a small number, often called the mesh spacing in numerical methods.

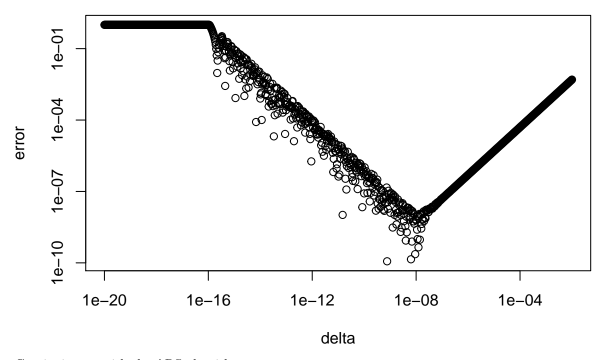
This approximation ignores higher order corrections, which means it will have first order truncation error:

$$\epsilon = \frac{1}{2}f''(x_0)h + \mathcal{O}(h^2)$$

Therefore, we want to choose our mesh spacing h as small as possible. However, because we are performing these computations on double-precision floating point computers there is a limit to how small we can make h before machine precision impacts the precision of this approximation. Looking at how we will compute the approximation of the derivative, we see that the numerator $f(x_0 + h) - f(x_0)$ requires the subtraction of two very similar numbers. Assuming $\mathcal{O}(f'(x)) \approx 1$, the difference will be $\approx \mathcal{O}(h)$. However the numbers we're subtracting have 16 digits of accuracy, so if h were 10^{-10} we the difference would only have 6 digits of accuracy! (The numerator and denominator are of similar order, so we expect only a small loss of precision in the division operation). Therefore we expect the mesh spacing with the highest accuracy to be around 10^{-8} .

If we consider the simple example of $f(x) = x^2$ we can see that this is approximately correct.

```
## vector of possible mesh spacings
delta <- 10**seq(-20, -2, length = 1000)
## example function x^2
quad <- function(x) x**2
## approximate derivative using forward difference
h_prime <- (quad(1.0 + delta) - quad(1.0))/delta
## Difference from true value of 2
error <- abs(h_prime - 2.0)/2.0
plot(delta, error, log = 'xy')</pre>
```



Continuing on with the ARS algorithm

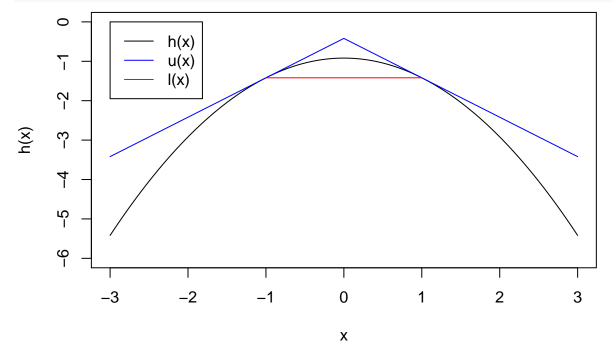
```
## initialize vectors
x0 <- sort(x0) # Ensures x0 is in ascending order</pre>
x_j \leftarrow c() # evaluated x values
h_j \leftarrow c() # evaluated h(x) values
dh_j <- c() # evaluated h'(x) values
x <- c() # accepted sample vector
## Evaluate h(x0)
h0 \leftarrow h(x0)
## Finite difference approximation to h'(x)
dh0 <- (h(x0 + dx) - h0)/dx
## Check for NaNs and infinities
isnum <- is.finite(h0)&is.finite(dh0)</pre>
x0 <- x0[isnum]
h0 <- h0[isnum]
dh0 <- dh0[isnum]
## Error if no good x0 values
if (length(h0) == 0) {
  stop('h(x0) either infinite or NaN.')
}
## ARS requires either finite bounds or at least
## one positive and one negative derivative
if (!(dh0[1] > 0 || is.finite(bounds[1]))) {
  stop('dh(x0)/dx must have at least one positive value, or left bound must be greater than -infinity.'
}
if (!(dh0[length(dh0)] < 0 || is.finite(bounds[2]))) {</pre>
```

```
stop('dh(x0)/dx must have at least one negative value, or right bound must be less than infinity.')
```

The first step specified by Wilks is to initialize our functions $u_k(x)$, $l_k(x)$, $s_k(x)$ and to pre-compute the intercepts of $u_k(x)$ (z_k). To do this we will write helper functions to calculate the coefficients of the piecewise linear functions as specified by Wilks.

```
##### HELPER FUNCTIONS #####
calc_z_j <- function (x, h, dh, bounds) {</pre>
  ## Calculate intercepts of piecewise u_j(x)
  ##
  ## Arguments
  ## x, h, dh: evaluated x, h(x) and h'(x)
  ## bounds: bounds of distribution h(x)
  ## Value
  ## intercepts (bounds) of piecewise u_j(x)
 L <- length(h)
  z \leftarrow (h[2:L] - h[1:L-1] - x[2:L]*dh[2:L] + x[1:L-1]*dh[1:L-1])/(dh[1:L-1] - dh[2:L])
 return (append(bounds[1], append(z, bounds[2])))
}
calc_u <- function (x, h, dh) {</pre>
  ## Calculate slope/intercept for piecewise u_j(x)
  ## Arguments
  ## x, h, dh: evaluated x, h(x), and h'(x)
  ## Value (list)
  ## m, b: slope/intercept of u(x)
 return (list(m = dh, b = h - x*dh))
}
calc_l <- function (x, h) {</pre>
  ## Calculate slope/intercept for piecewise l_j(x)
 ## Arguments
  ## x, h: evaluated x, and h(x)
  ##
  ## Value (list)
  ## m, b: slope/intercept of l(x)
 L <- length(h)
  m \leftarrow (h[2:L] - h[1:L-1])/(x[2:L] - x[1:L-1])
  b \leftarrow (x[2:L]*h[1:L-1] - x[1:L-1]*h[2:L])/(x[2:L] - x[1:L-1])
  return (list(m = m, b = b))
}
```

We see that already with only two points we have a good approximation to the rejection envelope $u_k(x)$ and squeezing function $l_k(x)$.



For the function

$$s_k(x) = \exp u_k(x) / \int_D \exp u(x') dx'$$

from which we'll sample x, instead of computing the function we will compute coefficients which will allow us to sample easily. To do this we'll re-define $s_k(x)$ as

$$s_k(x) = e^{u_k(x)} / \int_D e^{u(x')} dx' = \frac{e^{b_k} e^{m_k x}}{\sum_{j=1}^K \int_{z_j}^{z_{j+1}} e^{b_j} e^{m_j x'} dx'} = \frac{e^{b_k} e^{m_k x}}{\sum_{j=1}^K \frac{e^{b_k}}{m_k} (e^{m_k z_{j+1}} - e^{m_k z_j})}$$

```
calc_s_j <- function (m, b, z) {
    ## Calculate beta values and weights
    ## needed to sample from u(x)
    ##

## Arguments
## m, b: slope/offset for piecewise u_j(x)
## z: bounds z_j for piecewise function u_j(x)</pre>
```

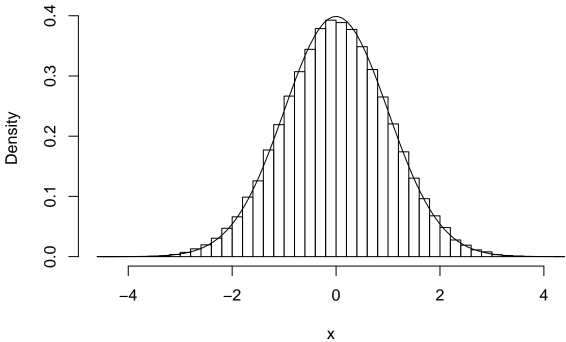
```
##
  ## Value (list)
  ## beta: amplitude of each piecewise segment in u(x)=beta*exp(m*x)
  ## w: area of each segment
  L <- length(z)
  eb <- exp(b) # un-normalized betas</pre>
  eb <- ifelse(is.finite(eb), eb, 0.0) # Inf/Nan -> 0
  ## treat m=0 case
  nz < - (m != 0.0)
  nz_sum \leftarrow sum((eb[nz]/m[nz])*(exp(m[nz]*z[2:L][nz]) - exp(m[nz]*z[1:L-1][nz])))
  z_{sum} \leftarrow sum(eb[!nz]*(z[2:L][!nz] - z[1:L-1]))
  ## Ensure integral of s(x) > 0
  if (nz_sum + z_sum <= 0.0) {</pre>
    stop('Area of s(x)=exp(u(x)) <= 0')
  ## Normalize beta
  beta <- eb/(nz_sum + z_sum)
  ## Calculate weights for both m!=0 and m=0 cases
  w \leftarrow ifelse(nz, (beta/m)*(exp(m*z[2:L]) - exp(m*z[1:L-1])), beta*(z[2:L] - z[1:L-1]))
 return (list(beta = beta, w = ifelse((w > 0) & is.finite(w), w, 0.0)))
}
draw_x <- function (N, beta, m, w, z) {</pre>
  ## Sample from distribution u(x), by selecting segment j
       based on segment areas (probabilities), and then sample
  ##
     from exponential distribution within segment using
  ##
      inverse CDF sampling.
  ##
  ## Arguments
  ## N: number of samples
  ## beta, m, w: parameters of u_j(x)
  ## z: intercepts (bounds) of u_j(x)
  ##
  ## Value (list)
  ## x: samples from u(x)
  ## J: segment index j of each sample
  J <- sample(length(w), N, replace = TRUE, prob = w) # choose random segments with probability w
  u <- runif(N) # uniform random samples
  ## Inverse CDF sampling of x
  x \leftarrow ifelse(m[J] != 0, (1.0/m[J])*log((m[J]*w[J]/beta[J])*u + exp(m[J]*z[J])), z[J] + w[J]*u/beta[J])
 return (list(x = x, J = J))
}
##### MAIN LOOP #####
## Loop until we have N samples (or until error)
while (length(x) < N) {
```

```
## Track iterations, 10000 is arbitrary upper bound
current_iter <- current_iter + 1</pre>
if (current iter > max iters) {
 stop('Maximum number of iterations reached.')
## Vectorized sampling in chunks
## Chunk size will grow as square of iteration,
## up until it is the size of the sample (N)
## This ensures small samples at first, while
## u(x) is a poor approximation of h(x), and
## large samples when the approximation improves.
chunk_size <- min(c(N, current_iter**2))</pre>
##### INITIALIZATION AND UPDATING STEP #####
## only re-initialize if there are new samples
if (length(x0) > 0) {
 ## Update with new values
 x_j \leftarrow append(x_j, x0)
 h_j \leftarrow append(h_j, h0)
 dh_j <- append(dh_j, dh0)
 x0 <- c()
 h0 \leftarrow c()
 dh0 < -c()
 ## Sort so that x_j's are in ascending order
 srtd <- sort(x_j, index.return = TRUE)</pre>
 x_j <- srtd$x
 h_j <- h_j[srtd$ix]</pre>
 dh_j <- dh_j[srtd$ix]</pre>
 L <- length(dh_j)
 ## Check for duplicates in x and h'(x)
 ## This prevents discontinuities when
 ## computing z_j's
 while (!all(((dh_j[1:L-1] - dh_j[2:L]) > dx) & ((x_j[2:L] - x_j[1:L-1]) > dx))) {
    ## Only keep values with dissimilar neighbors
    ## Always keep first index (one is always unique)
    dup \leftarrow append(TRUE, (((dh_j[1:L-1] - dh_j[2:L]) > dx)) & ((x_j[2:L] - x_j[1:L-1]) > dx)))
    x_j \leftarrow x_j[dup]
    h_j <- h_j[dup]
    dh_j <- dh_j[dup]
    L <- length(dh_j)
    if (L == 1) {
      break
    }
```

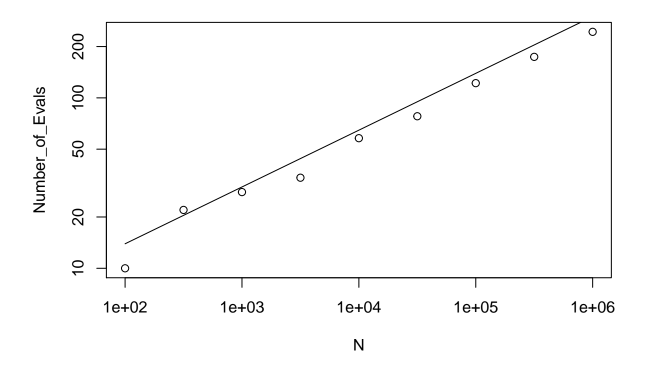
```
## Ensure log-concavity of function
  if(!all(dh_j[2:L] \leftarrow dh_j[1:L-1])) {
    stop('Input function f not log-concave.')
  ## pre-compute z_j, u_j(x), l_j(x), s_j(x)
  z_j \leftarrow calc_z_j(x_j, h_j, dh_j, bounds)
  u_j \leftarrow calc_u(x_j, h_j, dh_j)
  m_u <- u_j$m
  b_u <- u_j$b
  l_j \leftarrow calc_l(x_j, h_j)
  m_1 <- 1_j$m
  b_1 <- 1_j$b
  s_j \leftarrow calc_s_j(m_u, b_u, z_j)
  beta_j <- s_j$beta
  w_j <- s_j$w
}
##### SAMPLING STEP #####
## draw x from exp(u(x))
draws <- draw_x(chunk_size, beta_j, m_u, w_j, z_j)</pre>
x_s <- draws$x
J <- draws$J
## random uniform for rejection sampling
w <- runif(chunk_size)</pre>
## Warn if samples were outside of bounds
## This happens if bounds given don't reflect
## the actual bounds of the distribution
if (!all((x_s > bounds[1]) & (x_s < bounds[2]))) {
  ## Flag so warning only happens once
  if (!bds_warn) {
    warning('Sampled x* not inside bounds...please check bounds.')
    bds_warn <- TRUE
  ## Only keep x values within bounds
  ibd <- ((x_s > bounds[1]) & (x_s < bounds[2]))
  J \leftarrow J[ibd]
  x_s < x_s[ibd]
  w \leftarrow w[ibd]
## Index shift for l_j(x)
```

```
J_1 \leftarrow J - ifelse(x_s < x_j[J], 1, 0)
  ## only use x-values where l_j(x) > -Inf
  ibd <- (J_1 \ge 1) & (J_1 < length(x_j))
  ## Perform first rejection test on u(x)/l(x)
  y \leftarrow \exp(x_s[ibd]*(m_1[J_1[ibd]] - m_u[J[ibd]]) + b_1[J_1[ibd]] - b_u[J[ibd]])
  acc <- (w[ibd] <= y)
  ## Append accepted values to x
  x <- append(x, x_s[ibd][acc])</pre>
  ## Append rejected values to x0
  x0 <- append(x_s[!ibd], x_s[ibd][!acc])</pre>
  if (length(x0) > 0) {
    ## Evaluate h(x0)
    h0 \leftarrow h(x0)
    ## Finite difference approximation to h'(x)
    dh0 <- (h(x0 + dx) - h0)/dx
    ## Check for NaNs and infinities
    isnum <- is.finite(h0)&is.finite(dh0)</pre>
    x0 <- x0[isnum]
    h0 <- h0[isnum]
    dh0 <- dh0[isnum]</pre>
    w <- append(w[!ibd], w[ibd][!acc])[isnum]</pre>
    J <- append(J[!ibd], J[ibd][!acc])[isnum]</pre>
    ## Perform second rejection test on u(x)/h(x)
    acc \leftarrow (w \leftarrow exp(h0 - x0*m_u[J] - b_u[J]))
    ## Append accepted values to x
    x <- append(x, x0[acc])</pre>
  }
}
## Only return N samples (vectorized operations makes x sometimes larger)
x \leftarrow x[1:N]
hist(x, breaks = 50, freq = FALSE)
xr \leftarrow seq(min(x), max(x), length = 100)
lines(xr, dnorm(xr))
```

Histogram of x



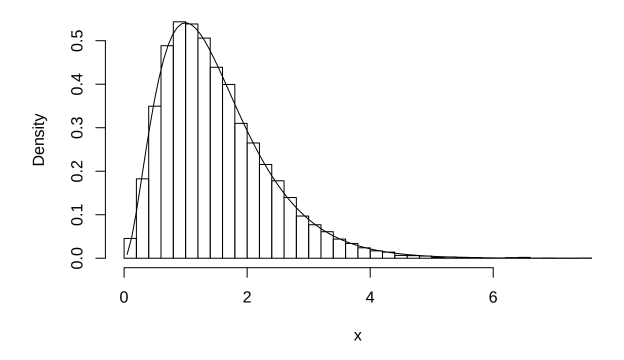
```
print(length(x_j))
## [1] 152
print(3*N**(1/3))
## [1] 139.2477
N_evals <- 0
f <- function (x) {
  N_{evals} << - N_{evals} + 1
  return (dnorm(x))
library('ars')
Number_of_Evals <- c()</pre>
N < -10**seq(2, 6, 0.5)
for (n in N) {
  x \leftarrow ars(f, n)
  Number_of_Evals <- append(Number_of_Evals, N_evals)</pre>
  N_evals <- 0
plot(N, Number_of_Evals, log = 'xy')
lines(N, 3*N**(1/3))
```



Example Usage

```
x <- ars(dgamma, 10000, x0 = c(0.1, 2.5), bounds = c(0.0, Inf), shape = 3, rate = 2)
hist(x, breaks = 50, freq = FALSE)
xr <- seq(min(x), max(x), length = 100)
lines(xr, dgamma(xr, shape = 3, rate = 2))</pre>
```

Histogram of x



Testing