Computational Statistics Summary

$Maximilian\ Pfundstein$ 2019-03-12

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1 Handling Computational Errors

```
x1 = 1/3
x2 = 1/4

if (all.equal(x1-x2, 1/12)) {
   print("Substraction is correct.")
} else {
   print("Substraction is wrong.")
}
```

2 Difference Quotient

$$f'(x) = \frac{f(x+\epsilon) - f(x)}{\epsilon}$$

```
f_prime = function(x, epsilon = 10^(-5)) {
  return( (f(x + epsilon) - f(x)) / epsilon)
}
```

3 Variance Estimators

Krzysztof:

$$\operatorname{Var}(\vec{x}) = \frac{1}{n-1} \left(\sum_{i=1}^{n} x_i^2 - \frac{1}{n} \left(\sum_{i=1}^{n} x_i \right)^2 \right)$$

My:

$$s = \frac{\sum_{i=1}^{n} (x_i - \overline{x})^2}{n-1}$$

```
custom_variance = function(x) {
  diff_mean = x - mean(x)
  return(sum(diff_mean^2 / (length(x) - 1)))
}
```

4 Optimization

4.1 optimize()

```
myMSE = function(lambda, pars) {
  model = loess(Y ~ X, data=pars, enp.target = lambda)
  prediction = predict(model, newdata = pars$Xtest)
  mse = sum((prediction - pars$Ytest)^2)/length(pars$Ytest)
  return(mse)
}

# parameters for the myMSE function -------
pars = list(X = train$Day, Y = train$LMR, Xtest = test$Day, Ytest = test$LMR)
lambdas = seq(from = 0.1, to = 40, by = 0.1)
# applying the myMSE function to all lambdas ------
mses = sapply(X = lambdas, FUN = myMSE, pars = pars)

o = optimize(myMSE, tol = 0.01, interval = c(0.1, 40), pars = pars)
o$minimum
o$objective
```

Plotting a function with a minimum:

```
lambdas[which.min(mses)]
length(lambdas)

df = data.frame(lambdas, mses)

ggplot(df) +
    geom_line(aes(x = lambdas, y = mses), color = "#C70039") +
    geom_point(aes(x = seq(0.1, 40, by = 0.1)[which.min(mses)],
    y = mses[which.min(mses)], color = "min MSE"),
    colour = "blue") +
    labs(title = "Lambdas VS MSEs", y = "MSE", x = "Lambda") +
    theme_minimal()
```

4.2 optim()

General usage:

```
optim(35, myMSE, method = "BFGS", pars = pars, control = list(fnscale = 1))
```

For optimizing likelihood:

```
# c(mu, sigma)
neg_llik_norm = function(par) {
    n = nrow(as.matrix(data))
    p1 = (n/2)*log(2*pi)
    p2 = (n/2)*log(par[2]^2)
    sum = sum((data - par[1])^2)
    p3 = 1/(2*par[2]^2) * sum
    return(p1+p2+p3)
}
```

```
# c(mu, sigma)
neg_llik_norm_prime = function(par) {
    n = nrow(as.matrix(data))
    mu_prime = -1/(n*par[2]^2) * sum(data-par[1])
    sigma_prime = 1/(2*par[2]^2) * (n - (1/(par[2]^2)) * sum((data-par[1])^2))

return(c(mu_prime, sigma_prime))
}

optim(c(0, 1), neg_llik_norm, method = "CG")
    optim(c(0, 1), neg_llik_norm, method = "CG", gr = neg_llik_norm_prime)
    optim(c(0, 1), neg_llik_norm, method = "BFGS")
    optim(c(0, 1), neg_llik_norm, method = "BFGS")
```

Answer: The negative log-likelihood function for the normal distribution is defined by:

$$\mathcal{L}(\mu, \sigma^2, x_1, ..., x_{100}) = \frac{n}{2} ln(2\pi) + \frac{n}{2} ln(\sigma^2) + \frac{1}{2\sigma^2} \sum_{j=1}^{n} (x_j - \mu)^2$$

The estimators are:

$$\hat{\mu}_n = \frac{1}{n} \sum_{j=1}^n x_j$$

and

$$\hat{\sigma}_n^2 = \frac{1}{n} \sum_{j=1}^n (x_j - \hat{\mu})^2$$

Answer: The partial derivates for the negative log-likelihood are given by:

$$\frac{\partial \mathcal{L}(\mu, \sigma^2, x_1, \dots, x_{100})}{\partial \mu} = -\frac{1}{n\sigma^2} \sum_{j=1}^n (x_j - \mu)$$

$$\frac{\partial \mathcal{L}(\mu, \sigma^2, x_1, \dots, x_{100})}{\partial \sigma^2} = \frac{1}{2\sigma^2} \left(n - \frac{1}{\sigma^2} \sum_{j=1}^n (x_j - \mu)^2 \right)$$

5 Sampling Based on Size

Task: Use a uniform random number generator to create a function that selects 1 city from the whole list by the probability scheme offered above (do not use standard sampling functions present in R).

```
get_city_by_urn_wo = function(city_pool) {

# We take the cumulative sum and then runif from 1 to max(cumulative sum).

# This way we respect the proportions. As we need every intermediate result,

# we use a loop

cumulative_pop_sum = 0
```

```
for (i in 1:nrow(city_pool)) {
    cumulative_pop_sum = cumulative_pop_sum + city_pool$Population[i]
    city_pool$CumSum[i] = cumulative_pop_sum
}

# Now we get a random value between 1 to max(cumulative sum). As larger muni-
# cipalities have larger ranges, this works as expected
selection =
    floor(runif(n = 1, min = 1, max = city_pool$CumSum[nrow(city_pool)]))

# Return the first city which has a greater CumSum than the selection
return(city_pool[city_pool$CumSum > selection,][1, c(1, 2)])
}
```

Task: Use the function you have created in step 2 as follows:

- a. Apply it to the list of all cities and select one city
- b. Remove this city from the list
- c. Apply this function again to the updated list of the cities
- d. Remove this city from the list
- e. ... and so on until you get exactly 20 cities.

Answer: We will combine all of these steps in one function. We're lazy.

```
get_n_cities = function(data, n) {

# Create a copy to not touch the original data.
city_pool = data
selected_cities = data.frame()

# As long as we don't have n samples, get one and remove it from the pool,
# as we sample without replacement
while(nrow(selected_cities) < n) {
    selected_city = get_city_by_urn_wo(city_pool)
    selected_cities = rbind(selected_cities, selected_city)
    city_pool = city_pool[!rownames(city_pool) %in% rownames(selected_cities),]
}

return(selected_cities)
}
sample = get_n_cities(data, 20)</pre>
```

6 Inverse CDF Method

The double exponential (Laplace) distribution is given by formula

$$DE(\mu, \alpha) = \frac{\alpha}{2} e^{-\alpha|x-\mu|}$$

Task: Write a code generating double exponential distribution DE(0,1) from Unif(0,1) by using the inverse CDF method. Explain how you obtained that code step by step. Generate 10000 random numbers from this distribution, plot the histogram and comment whether the result looks reasonable.

- 1. Derive the CDF from the PDF (dfunc())by taking the integral $\int_{-\infty}^{x} \text{CDF} dx$. This function is the pfunc() (CDF!, cumulative).
- 2. Swap x and y to receive the quantile function qfunc().
- 3. Combine both functions to create the rfunc().

This can look like this:

```
# double exponential (Laplace) distribution
# PDF
ddel = function(x = 1, mu = 0, b = 1) {
return(1/(2*b) * exp(-abs(x-mu)/(b)))
}
# CDF
pdel = function(x = 1, mu = 0, b = 1) {
  return(1/2 + 1/2 * sgn(x-mu) * (1 - exp(-abs(x-mu)/b)))
# Quantile
qdel = function(p, mu = 0, b = 1) {
  if (p < 0 \mid p > 1) stop("p must be in range (0, 1)")
  if (p \le 0.5) return(mu + b * log(2 * p))
  return (mu - b * log(2 - 2 * p))
}
# Random
rdel = function(n = 1, mu = 0, b = 1) {
  quantiles = runif(n = n, min = 0, max = 1)
  rdels = sapply(X = quantiles, FUN = qdel, mu = mu, b = b)
  return(rdels)
```

To look how the distribution looks like with different parameters, use the following code and plots:

```
sample_rdel_0_1 = rdel(10000, mu = 0, b = 1)
sample_rdel_0_2 = rdel(10000, mu = 0, b = 2)
sample_rdel_0_4 = rdel(10000, mu = 0, b = 4)
sample_rdel_m5_4 = rdel(10000, mu = -5, b = 4)

df = data.frame(sample_rdel_0_1, sample_rdel_0_2, sample_rdel_0_4, sample_rdel_m5_4)

p1 = ggplot(df) +
    geom_histogram(aes(x = sample_rdel_0_1), color = "#FFC300", fill = "#FFC300", binwidth = 0.01) +
    xlim(-10, 10) +
    ylim(0, 60) +
    ggtitle("RD(0, 1)") +
    theme_minimal()

# p2, p3, p4

grid.arrange(p1, p2, p3, p4, nrow = 2)
```

7 Acceptance / Rejection Method

For plotting the PDF and CDF to compare for instance two of them:

```
sequence = seq(from = -10, to = 10, by = 0.01)
dnorm_samples = sapply(X = sequence, FUN = dnorm)
ddel_samples = sapply(X = sequence, FUN = ddel)
pnorm_samples = sapply(X = sequence, FUN = pnorm)
pdel_samples = sapply(X = sequence, FUN = pdel)
df = data.frame(dnorm_samples, ddel_samples, pnorm_samples, pdel_samples)
ggplot(df) +
  geom_line(aes(x = sequence, y = dnorm_samples,
                colour = "Normal Distribution (PDF)")) +
  geom_line(aes(x = sequence, y = ddel_samples,
                colour = "Double Exponential Distribution (PDF)")) +
  labs(title = "dnorm() and ddel()", y = "Density",
  x = "X", color = "Legend") +
  scale_color_manual(values = c("#17202A", "#C70039")) +
  theme_minimal()
ggplot(df) +
  geom_line(aes(x = sequence, y = pnorm_samples,
                colour = "Normal Distribution (CDF)")) +
  geom_line(aes(x = sequence, y = pdel_samples,
               colour = "Double Exponential Distribution (CDF)")) +
  labs(title = "pnorm() and pdel()", y = "Cumulative Density",
  x = "X", color = "Legend") +
  scale_color_manual(values = c("#17202A", "#C70039")) +
  theme minimal()
```

For calculating the C value, use this code:

```
c = max(dnorm_samples / ddel_samples)
```

Sometimes we cannot do this this easily or have to derive the optimal C by hand. This can be done as follows:

$$C \ge \frac{\text{target}}{\text{proposal}} = \frac{f(x)}{g(x)}$$

We then calculate:

$$\max_{x} \frac{f(x)}{g(x)}$$

Which is taking the derivative and setting it to 0. Then we solve for x and plug in x into

$$\frac{f(x)}{g(x)} = C$$

Note: Sometimes C = M.

For creating nice plots, use this:

```
df$scaled_envelop = c * df$ddel_samples
ggplot(df) +
 geom_line(aes(x = sequence, y = dnorm_samples,
                colour = "Normal Distribution (PDF)")) +
  geom_line(aes(x = sequence, y = ddel_samples,
                colour = "Double Exponential Distribution (PDF)")) +
 geom_line(aes(x = sequence, y = scaled_envelop,
               colour = "Scaled Double Exponential Distribution (PDF)")) +
 labs(title = "Envelope", y = "Density",
  x = "X", color = "Legend") +
  scale_color_manual(values = c("#17202A", "#C70039", "#581845")) +
  theme_minimal()
ggplot(df) +
  geom_ribbon(aes(x = sequence, ymin = df$dnorm_samples, ymax = df$scaled_envelop),
              alpha = 0.8, fill = "#C70039", color = "#C70039") +
  geom_ribbon(aes(x = sequence, ymin = 0, ymax = df$dnorm_samples),
              alpha = 0.8, fill = "#DAF7A6", color = "#DAF7A6") +
 labs(title = "Acceptance and Rejection Regions", y = "Density",
  x = "X", color = "Legend") +
  scale_color_manual(values = c("#17202A", "#C70039", "#581845")) +
  theme minimal()
```

And for actually using it, use:

```
rs = c()
rs_rejected = c()

while (length(rs) < 2000) {
    # Take a random sample from our proposal (x-axis)
    z = rdel(n = 1, mu = 0, b = 1)

# Take a uniform, thus a random y value
    u = runif(n = 1, min = 0, max = c * ddel(z))

# Check in which region this on lies
    if (u <= dnorm(z)) {
        rs = c(rs, z)
    }
    else {
        rs_rejected = c(rs_rejected, z)
    }
}</pre>
```

Plot the drawn samples:

```
theme_minimal()

Expected rejection rate:
1 - 1/c

Observed rejection rate:
length(rs_rejected) / (length(rs)+length(rs_rejected))
```

8 Metropolis-Hastings Algorithm

We have given a target function (probably only a proportional one).

```
# Target function with original scaling
f = function(x) {
  return(120 * x^5 * exp(-x))
}
# Target function
df = function(x) {
 x = ifelse(x \le 0, 0.000001, x)
 return(x^5 * exp(-x))
sequence = seq(from = 0.01, to = 20, by = 0.01)
real_f = f(sequence)
plotdf = data.frame(sequence, real_f)
ggplot(plotdf) +
  geom_line(aes(x = sequence, y = real_f), color = "#6091ec") +
  labs(title = "Target Density Function", y = "Density",
  x = "X", color = "Legend") +
 theme_minimal()
```

The Metropolis-Hastings algorithm is implemented by:

```
# Samples from proposel from the random walk (MC)
xt = x 0
xt_1 = x_0
while (length(samples) < n) {</pre>
  # Generate proposal state
  if (proposal == "lnorm") {
    x_star = rlnorm(n = 1, meanlog = log(xt), sdlog = 1)
    # Calculate correction factor C
    c = dlnorm(xt_1, meanlog = log(xt), sdlog = 1) /
    dlnorm(x_star, meanlog = log(xt), sdlog = 1)
  else if (proposal == "chisquared") {
   x_star = rchisq(n = 1, df = floor(xt + 1))
    # Calculate correction factor C
    c = dchisq(x = xt_1, df = floor(xt + 1)) /
    dchisq(x_star, df = floor(xt + 1)) }
  else {
    stop("Invalid proposal.")
  # Calculate acceptance probability alpha
  if (df(xt_1) <= 0) {</pre>
    # We need this to avoid troublesome areas where the density is so low that
    # it's O from a computiationally point of view.
   alpha = 0
  }
  else {
    alpha = min(1, df(x_star)/df(xt_1) * c)
  # Generate u from uniform
  u = runif(n = 1, min = 0, max = 1)
  # Decide if to accept or reject the proposal
  if (u <= alpha) { # Accept</pre>
   xt_1 = xt
   xt = x_star
    samples = c(samples, x_star)
  }
  else {
   # Reject
   xt = xt_1
# Return samples
if (keep_burnin) return(samples) {
  return(samples[b+1:length(samples)])
```

```
}
```

Create a fancy plot of the burn-in period using:

And to plot the distribution of the samples, use:

9 Gelman-Rubin Factor

The custom implementation looks like this:

```
gelman_rubin_factor = function(sequence_matrix) {
    k = nrow(sequence_matrix)
    n = ncol(sequence_matrix)

    B = n/(k-1) * sum((rowMeans(sequence_matrix) - mean(sequence_matrix))^2)
    S_squared = rowSums((sequence_matrix - rowMeans(sequence_matrix))^2 / (n - 1))
    W = sum(S_squared / k)
    V = ((n - 1) / n * W) + (1/n * B)
    R = sqrt(V / W)
    return(R)
}
```

And calling it with the previously defined Metropolis-Hastings algorithm, use:

```
k = 10 # row
n = 1000 # col
```

There is also a built-in implementation which can be invoked with the following code:

```
library(coda)
results5 = list()
results10 = list()
results50 = list()
k = 10
for (i in 1:k) {
  results5[[i]] = mcmc(metropolis_hastings(5, x_0 = i, b = 0,
                                   proposal = "chisquared", keep_burnin = TRUE))
}
for (i in 1:k) {
  results10[[i]] = mcmc(metropolis_hastings(10, x_0 = i, b = 0,
                                   proposal = "chisquared", keep_burnin = TRUE))
}
for (i in 1:k) {
  results50[[i]] = mcmc(metropolis_hastings(50, x_0 = i, b = 0,
                                   proposal = "chisquared", keep_burnin = TRUE))
}
mcmc list5 = mcmc.list(results5)
gelman.diag(mcmc_list5)
mcmc_list10 = mcmc.list(results10)
gelman.diag(mcmc_list10)
mcmc_list50 = mcmc.list(results50)
gelman.diag(mcmc_list50)
```

10 Monte Carlo Integration

Let's say we want to integrate the following integral:

$$\int_0^\infty x^4 e^{-2x} dx$$

so

$$f(x) = x^4 e^{-2x}$$

For estimating this integral with MCMC we need to split f(x) into d(x)g(x) which can be chosen in any way, as long as the following conditions for g(x) are met:

- $g(x) \ge 0$ for the drawn interval.
- The integral inside the interval is finite, so $\int_a^b g(x) = C < \infty$

g(x) must drawn from the whole integral range, so from a to b, and exactly the range. If one of the limits its ∞ , g(x) must have the same limit (like a gamma distribution for instance for 0 to ∞).

Then draw samples from g(x), x_i .

Then calculate:

$$\frac{1}{n} \sum_{i=1}^{n} d(x_i)$$

Which is the mean of the $d(x_i)$ values.

In this example this looks like this:

The sampling function is given by:

$$d(x, \alpha = 2, \beta = 2) = 4xe^{-2x}$$

Note that $Gamma(\alpha) = (\alpha - 1)!$

We then identify:

$$g(x) = \frac{1}{4}x^3$$

So we simply put our drawn samples into g(x) and calculate the mean.

```
g = function(x) return(1/4 * x^3)
mean(g(samples))
```

Note: When the limits do not include infinity, we have to scale the results. So we take the result times C where C = b - a. This might be wrong, but that's what I got from different examples.

11 Gibbs Sampling

For Gibbs Sampling it is mandatory to find the Marginals. For doing that, do the following:

- First write down the prior and the likelihood. The is no general way for this, it depends on the problem.
- Then write down the posterior which is the product of both.
- For finding the marginals, drop every factor that is not dependend on the marginal variable.

If we have to combine two normal distributions, it can be done using the following rules:

The product of two normal distributions is given by (see https://www.johndcook.com/blog/2012/10/29/product-of-normal-pdfs/ as a recap if not present):

$$\sigma_{new}^2 = \frac{\sigma_1^2 \sigma_2^2}{\sigma_1^2 + \sigma_2^2}$$

$$\mu_{new} = \frac{\sigma_1^{-2}\mu_1 + \sigma_2^{-2}\mu_2}{\sigma_1^{-2} + \sigma_2^{-2}}$$

For implementing the Gibbs Sampler we write one function for getting the marginals and one main function for doing the actual sampling.

```
get_mu = function(i, sigma = 0.2, mu, Y) {
  if (i == 1) {
    # First Marginal
   return(rnorm(n = 1, mean = (mu[2] + Y[1])/2, sd = sqrt(1/2 * sigma)))
  else if (i == 50) {
    # Last Marginal
   return(rnorm(n = 1, mean = (2 * mu[i-1] - Y[i-1] + 2 * Y[i])/3,
                 sd = sqrt(2/3 * sigma)))
  }
  # General Marginals
   return(rnorm(n = 1, mean = (2 * mu[i-1] + 2 * mu[i+1] + 2 * Y[i] - Y[i-1])/5,
                 sd = sqrt(2/5 * sigma)))
}
gibbs_sample = function(n, Y, include_matrix = FALSE) {
  # Vectors to store the samples in
  mu matrix = matrix(0, ncol = length(Y), nrow = n)
  mu = rep(0, length(Y))
  for(j in 1:n) {
   for (i in 1:length(Y)) {
      mu[i] = get_mu(i, sigma = 0.2, mu = mu, Y = Y)
    mu_matrix[j,] = mu
  if (include_matrix) {
   return(list(means = colMeans(mu_matrix), matrix = mu_matrix))
  }
```

```
return(colMeans(mu_matrix))
}
```

Then this function can be used like this. This also includes a plot of the random walk.

```
gibbs_sample_result = gibbs_sample(1000, Y, TRUE)

data$gibbs_samples = gibbs_sample_result$means

ggplot(data) +
   geom_line(aes(x = X, y = Y), color = "#C70039") +
   geom_line(aes(x = X, y = gibbs_samples), color = "#581845") +
   labs(title = "Dependence from Concentration on Day of Measurement", y =
        "Concentration", x = "Day of Measurement", color = "Legend") +
   theme_minimal()
```

If a trace plot is needed, use this:

12 Hypothesis Testing

12.1 Defining a Test-Statistics

Pay attention how **exactly** this on is defined, as here for instance Y and \hat{Y} mixed.

$$T = \frac{\hat{Y}(X_b) - \hat{Y}(X_a)}{X_b - X_a}, \text{ where } X_b = \operatorname{argmax}_X Y(X), X_a = \operatorname{argmin}_X Y(X)$$

```
data = data.frame(X = lottery$Day_of_year, Y = lottery$Draft_No)

test_statistics = function(X, Y, Y_hat) {
  b_index = which.max(Y)
  a_index = which.min(Y)

return((Y_hat[b_index] - Y_hat[a_index]) / (X[b_index] - X[a_index]))
}
```

12.2 Non-Parametric Bootstrap

```
f = function(data, ind) {
  data1 = data[ind,]
```

```
model = loess(Draft_No ~ Day_of_year, data1)

T_value =
    test_statistics(data1$Day_of_year, data1$Draft_No, Y_hat = model$fitted)

return(T_value)
}

# T(D) for the original data
data$Y_hat = loess(Draft_No ~ Day_of_year, lottery)$fitted
T_value_original = test_statistics(data$X, data$Y, data$Y_hat)

# T for the bootstrapped samples
nonparam_bootstrap =
    boot(lottery, statistic = f, R = 2000, parallel = "multicore")
p_value_original = mean(nonparam_bootstrap$t > 0)
```

12.3 Density of non-parametric T-Values

12.4 Defining the Hypothesis Test (Permutation Test)

12.5 Crude Estimate of the Power

Question: Make a crude estimate of the power of the test constructed in Step 4:

- a) Generate (an obviously non-random) dataset with n=366 observations by using same X as in the original data set and $Y(x)=max(0,min(\alpha x+\beta,366))$, where $\alpha=0.1$ and $\beta\sim N(183,sd=10)$.
- b) Plug these data into the permutation test with B = 200 and note whether it was rejected.
- c) Repeat Steps 5a-5b for $\alpha = 0.2, 0.3, ..., 10$.

What can you say about the quality of your test statistics considering the value of the power?

```
simulate_data = function(X, Y_hat, hypothesis = test_hypothesis,
                         statistics = test_statistics, alpha = 0.1,
                         beta_mean = 183, beta_sd = 10, b = 200, limit = 366) {
  artificial = function(X, alpha) {
   beta = rnorm(n = nrow(lottery), mean = beta_mean, sd = beta_sd)
   return(max(0, min(alpha * X + beta, limit)))
  }
  X dataframe = X
  Y_dateframe = sapply(X, artificial, alpha)
  Y_hat = loess(Y_dateframe ~ X_dataframe)$fitted
  Y_hat_dataframe = Y_hat
  data_artificial = data.frame(X_dataframe, Y_dateframe, Y_hat_dataframe)
  colnames(data_artificial) = c("X", "Y", "Y_hat")
  T_val = test_statistics(data_artificial$X, data_artificial$Y, data_artificial$Y_hat)
 return(test_hypothesis(data_artificial, statistics, b, T_val))
alphas = seq(from = 0.1, to = 10.0, by = 0.1)
no_cores = detectCores()
cl = makeCluster(no cores)
clusterExport(cl, list("simulate_data", "lottery", "test_hypothesis",
                       "test_statistics", "data"))
simulated_p_values =
  parSapply(cl, alphas, FUN = function(alpha) {
    simulate_data(alpha = alpha, X = lottery$Day_of_year, Y_hat = data$Y_hat)
  })
stopCluster(cl)
```

Plot for p-values with respect to α and p-value line:

```
df = data.frame(alphas, simulated_p_values)

ggplot(df)+
  geom_point(aes(x = alphas, y = simulated_p_values), color = "black",
```

```
fill = "#dedede", shape = 21) +
geom_hline(aes(yintercept = 0.05), color = "orange") +
labs(title = "p-values depending on alpha",
    y = "p-value", x = "Alpha", color = "Legend") +
theme_minimal()
```

Calculating the actual power is done by:

```
1 - mean(simulated_p_values > 0.05)
```

13 Boostrap with Bias-Correction

Doing a normal bootstrap.

The estimate of the mean price is:

```
bootstrap_mean_price = mean(house_bootstrap$t)
print(bootstrap_mean_price)
```

The bootstrap-bias-correction is given by the following values. The first one is the bias-correction and the second one is the bias corrected mean.

```
prices_mean = mean(prices$Price)
print(prices_mean - bootstrap_mean_price)
print(2 * prices_mean - bootstrap_mean_price)
```

The variance of the mean is given by:

```
bootstrap_variance_price = as.numeric(var(house_bootstrap$t))
# bootstrap_mean_price = 1 / (B-1) * sum((house_bootstrap$t -
# mean(house_bootstrap$t))^2)
print(bootstrap_variance_price)
```

Now we will create the 95% confidence intervals.

```
confidence_interval = boot.ci(house_bootstrap)
print(confidence_interval)
```

As this output does not include the BCa, we will print the intervals manually. Note that the last to values in each row represent the confidence interval.

```
confidence_interval$percent
confidence_interval$bca
confidence_interval$normal
```

14 Variance Estmation using Jackknife

```
f_prices_jackknife = function(ind, data) {
  data1 = data[-ind,]
  return(mean(data1$Price))
}
# First create the statistics using jackknife
n = length(prices$Price)
indices = seq(from = 1, to = n, by = 1)
jackknife_statistics = sapply(indices, f_prices_jackknife, prices)
# For the variance estimate we will first calculate Ti_star
Ti_star = sapply(jackknife_statistics, FUN = function(tdi, n, prices_mean) {
 return(n * prices mean - ((n - 1) * tdi))
}, n = n, prices_mean = prices_mean)
# Now we calculate J_T
J_T = mean(Ti_star)
# And now we can calculate the Variance
variance_jackknife = 1 / (n * (n - 1)) * sum((Ti_star - J_T)^2)
print(variance_jackknife)
```

15 Confidence Intervals with Respect to Length and Location

```
boostrap_estimated_mean_corrected = (2 * prices_mean - bootstrap_mean_price)
The length of the CI is:
print(confidence_interval$percent[5] - confidence_interval$percent[4])
The mean is located the following "percent" of the CI range.
(boostrap_estimated_mean_corrected - confidence_interval$percent[4]) /
  (confidence_interval$percent[5] - confidence_interval$percent[4]) * 100
ggplot(prices) +
  geom_histogram(aes(x = prices$Price),
                 color = "#000000", fill = "#dedede") +
  annotate("rect", xmin=confidence_interval$percent[4],
           xmax=confidence_interval$percent[5], ymin=0, ymax=Inf,
           alpha=0.5, fill="#86b4ff") +
  geom_vline(aes(xintercept = boostrap_estimated_mean_corrected),
             color = "orange") +
  labs(title = "Histrogram of Prices",
       y = "Frequency",
  x = "Prices", color = "Legend") +
  theme_minimal()
```

16 Genetic Algorithms

First define the function to minimize or maximize.

```
f = function(x) {
  left = x^2/exp(x)
  exponent = (-9*sin(x))/(x^2+x+1)
  right = 2*exp(exponent)
  return(left-right)
}
```

Plotting this function:

```
sequence = seq(from = 0, to = 30, by = 0.1)
f.sequence = f(sequence)

df = data.frame(sequence, f.sequence)

ggplot(df) +
    geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
    labs(title = "f(x)", y = "f(x)", x = "x") +
    theme_minimal()
```

Max value at x.

```
print(sequence[which.max(f.sequence)])
```

And y or respect f(x) for that:

```
print(max(f.sequence))
```

Then the crossover function, might be different though.

```
crossover = function(x, y) return((x+y)/2)
```

Then the mutation.

```
mutate = function(x) return((x^2)%%30)
```

Initial Population.

```
X = seq(from = 0, to = 30, by = 5)
```

Corresponding values.

```
Values = f(X)
```

And then the algorithm itself. Keep in mind that this one does not save the overall best individual.

```
genetic = function(X, Values, maxiter = 100, mutprob = 0.05) {
  best_individual = NaN

for (i in 1:maxiter) {
    # 1)
    parents = sample(1:length(X), size = 2)

# 2) I don't know why we should order here!?
    victim = which.min(Values)
```

```
# 3)
    child = crossover(X[parents][1], X[parents][2])
    if (mutprob > runif(n = 1, min = 0, max = 1)) {
        child= (mutate(child))
    }

# 4)
    X[victim] = child
    Values[victim] = f(child)

# 5)
    best_index = which.max(Values)
    best_individual = list(cx = X[best_index], cy = Values[best_index])
}
return(list(best = best_individual, population = X))
}
best_individual = genetic(X, Values, 100, 0.05)$best
```

Plot the best individual.

Plot the initial distribution.

Six grid plot with different settings.

```
genetic1 = genetic(X, Values, 10, 0.1)
genetic2 = genetic(X, Values, 10, 0.5)
genetic3 = genetic(X, Values, 10, 0.9)
genetic4 = genetic(X, Values, 100, 0.1)
genetic5 = genetic(X, Values, 100, 0.5)
genetic6 = genetic(X, Values, 100, 0.9)
population1 = data.frame(genetic1$population, f(genetic1$population))
best1 = as.data.frame(genetic1$best)
```

```
population2 = data.frame(genetic2$population, f(genetic2$population))
best2 = as.data.frame(genetic2$best)
population3 = data.frame(genetic3$population, f(genetic3$population))
best3 = as.data.frame(genetic3$best)
population4 = data.frame(genetic4$population, f(genetic4$population))
best4 = as.data.frame(genetic4$best)
population5 = data.frame(genetic5$population, f(genetic5$population))
best5 = as.data.frame(genetic5$best)
population6 = data.frame(genetic6$population, f(genetic6$population))
best6 = as.data.frame(genetic6$best)
p1 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom_point(aes(x = genetic1.population, y = f.genetic1.population.),
             data = population1, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best1, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
  labs(title = "maxiter = 10, mutprob = 0.1", y = "f(x)", x = "x") +
 theme_minimal()
p2 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom_point(aes(x = genetic2.population, y = f.genetic2.population.),
             data = population2, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best2, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
 labs(title = "maxiter = 10, mutprob = 0.5", y = "f(x)", x = "x") +
  theme_minimal()
p3 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom point(aes(x = genetic3.population, y = f.genetic3.population.),
             data = population3, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best3, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
  labs(title = "maxiter = 10, mutprob = 0.9", y = "f(x)", x = "x") +
  theme minimal()
p4 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom_point(aes(x = genetic4.population, y = f.genetic4.population.),
             data = population4, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best4, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
  labs(title = "maxiter = 100, mutprob = 0.1", y = "f(x)", x = "x") +
```

```
theme_minimal()
p5 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom_point(aes(x = genetic5.population, y = f.genetic5.population.),
             data = population5, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best5, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
  labs(title = "maxiter = 100, mutprob = 0.5", y = "f(x)", x = "x") +
  theme_minimal()
p6 = ggplot(df) +
  geom_line(aes(x = sequence, y = f.sequence), color = "#C70039") +
  geom_point(aes(x = genetic6.population, y = f.genetic6.population.),
             data = population6, color = "black",
            fill = "#FFC300", shape = 21, size = 2, stroke = 1) +
  geom_point(aes(x = cx, y = cy), data = best6, color = "black",
            fill = "#7BA9FF", shape = 21, size = 2, stroke = 1) +
  labs(title = "maxiter = 100, mutprob = 0.9", y = "f(x)", x = "x") +
  theme_minimal()
grid.arrange(p1, p2, p3, p4, p5, p6, nrow = 3)
```

17 Expectation-Maximization Algorithm (EM)

Plot for two lines in one graph.

```
ggplot(data) +
  geom_line(aes(x = X, y = Z, color = "Z")) +
  geom_line(aes(x = X, y = Y, color = "Y")) +
  labs(title = "Dependence of Z and Y versus X", y = "Z/Y", x = "X") +
  scale_color_manual(values = c("#C70039", "#581845")) +
  theme_minimal()
```

In general, perform:

- E-Step (Expectation)
- M-Step (Maximazation)

17.1 Expectation-Step (E-Step)

For the E-Step, we write down:

$$Q(\theta, \theta^k) = E[\mathcal{L}(\theta|Y, Z)|\theta^k, Y]$$

where θ is the unknown parameter (for instance λ). Then we take the log likelihood and estimate the latent parameters, for instance by using the expected value. This depends on the distribution. The log likelihood is usually the log of the product of the given distributions.

It could look like this:

$$\mathcal{L}(\lambda|Y,Z) \sim \ln \left(\prod_{i=1}^{n} \frac{X_i}{\lambda} e^{-X_i/\lambda} * \frac{X_i}{2\lambda} e^{-X_i/2\lambda} \right)$$

And then taking the expected value for the missing variables.

$$E[\mathcal{L}(\lambda|Y,Z)] = -n * ln(2\lambda^2) + 2\sum_{i=1}^{n} ln(X_i) - \sum_{i=1}^{n} \frac{X_i Y_i}{\lambda} - E\left[\sum_{i=1}^{n} \frac{X_i Z_i}{2\lambda}\right]$$

So the E-Step concludes with:

$$Q(\lambda, \lambda^k) = E[\mathcal{L}(\lambda|Y, Z)] = -n * ln(2\lambda^2) + 2\sum_{i=1}^n ln(X_i) - \sum_{i=1}^n \frac{X_i Y_i}{\lambda} - \sum_{i=1}^\beta \frac{X_i Z_i}{2\lambda} - (n-\beta)\frac{\lambda_k}{\lambda}$$

17.2 Maximazation-Step (M-Step)

Usually this can be done by taking the derivative with respect to the parameters to estimate. There might be different ways to do the maximization though.

$$\frac{\partial Q(\lambda, \lambda^k)}{\partial \lambda} = -\frac{2n}{\lambda} + \sum_{i=1}^n \frac{X_i Y_i}{\lambda^2} + \frac{1}{2} \sum_{i=1}^\beta \frac{X_i Z_i}{\lambda^2} + (n - \beta) \frac{\lambda_k}{\lambda^2}$$

And then simply solve for the parameter that maximizes.

$$\lambda = \frac{\sum_{i=1}^{n} X_{i} Y_{i} + \frac{1}{2} \sum_{i=1}^{\beta} X_{i} Z_{i} + (n - \beta) \lambda_{k}}{2n}$$

17.3 EM-Implementation

The implementatio can look like this:

```
# To keep it simple we will not pass the data to the function.
lambda_estimate_em = function (input_iterations = 100, input_threshold = 0.0001) {
   iterations_max = input_iterations
   iterations = 0
    threshold = input_threshold

   n = nrow(data)
   lambdas = c()
   lambda = NaN
   lambda_k = 100

   Z = data$Z[!is.na(data$Z)]
   A = data$Z[is.na(data$Z)]

   Z_index = which(!is.na(data$Z))
   A_index = which(is.na(data$Z))

   X = data$X[Z_index]
```

If we have to plot the expected values, take the formula from that from the distribution, for instance for the exponential one it would look like this.

$$E[Y] = \frac{\lambda}{X_i}, \quad E[Y] = \frac{2\lambda}{X_i}$$

```
data$Y_E = (lambda_result)/data$X
data$Z_E = (2*lambda_result)/data$X

ggplot(data) +
    geom_line(aes(x = X, y = Z, color = "Z")) +
    geom_line(aes(x = X, y = Y, color = "Y")) +
        geom_line(aes(x = X, y = Y, color = "Y")) +
        geom_line(aes(x = X, y = Z_E, color = "E[Z]")) +
        geom_line(aes(x = X, y = Y_E, color = "E[Y]")) +
        labs(title = "Dependence of Z, Y, E[Z] and E[Y] versus X", y = "Z/Y", x = "X") +
        scale_color_manual(values = c("#581845", "#FFC300", "#C70039", "#6091ec")) +
        theme_minimal()
```

18 Miscellaneous Plots

18.1 Histrogram

General rool for n datapoints: \sqrt{n} bars.

18.2 Histrogram with Mean

18.3 Simple X/Y Plot

```
ggplot(data) +
  geom_line(aes(x = X, y = Y), color = "#c70039") +
  labs(title = "Dependence from Concentration on Day of Measurement",
      y = "Concentration", x = "Day of Measurement", color = "Legend") +
  theme_minimal()
```

18.4 Variance Plot

```
for (i in 1:length(v)) {
   Xi = v[1:i]
   X$vec_customvar[[i]] = custom_variance(as.vector(Xi))
}

ggplot(X[2:nrow(X),]) +
   geom_point(aes(x = index, y = value, colour = "Difference")) +
   geom_point(aes(x = index, y = vec_myvar, colour = "my_var()")) +
   geom_point(aes(x = index, y = vec_var, colour = "var()")) +
   geom_point(aes(x = index, y = vec_customvar, colour = "custom_variance()")) +
   labs(title = "Difference in Variance", y = "Variance",
   x = "Sequence", color = "Legend") +
   scale_color_manual(values = c("#17202A", "#C70039", "#407AFF", "#FFC300")) +
   scale_x_log10() +
   theme_minimal()
```

18.5 Scatterplot With Geom Smoother

19 Useful Code Snippets

19.1 RMarkdown Setup

```
# ---
#title: "Computational Statistics Summary"
#author: "Maximilian Pfundstein"
#date: "`r Sys.Date()`"
#output:
# html_document:
# df_print: paged
# toc: true
  toc_float: true
#
# number_sections: true
# pdf_document:
#
  toc: true
  toc_depth: 3
#
  number\_sections:\ true
```

19.2 Knitr options

```
#{r setup, include=FALSE}
#knitr::opts_chunk$set(echo = TRUE, cache = TRUE, include = TRUE, eval = FALSE,
# fig.pos = 'H')
```

19.3 Including Source Code

```
#{r, ref.label=knitr::all_labels(), echo = TRUE, eval = FALSE, results = 'show'}
```