## Math 504 HW11

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2

 $\mathbf{a}$ 

$$S(x) = \sum_{j=1}^{D} \alpha_{j} h_{j}(x) = \alpha_{1} h_{1}(x) + \alpha_{2} h_{2}(x) + \dots + \alpha_{D} h_{D}(x)$$

which create a linear function space F, We have previously shown that

$$\min_{f \in F} \sum_{i=1}^{N} (y_i - f(x^{(i)}))^2 = \min_{\alpha \in \mathbb{R}^k} ||y - B\alpha||^2$$

where B is an  $N \times K$  matrix of functions of  $b_i(x)$ , where entry  $B_{kl} = b_l(x_k)$  in a linear function space F, and  $\alpha$  is the coefficients. So with S(x) being the f(x),  $h_j(x)$  is analogous to  $b_j(x)$ , and the  $\alpha$ 's fill the same roll. Then

$$\min_{S \in F} \sum_{i=1}^{N} (y_i - S(x_i))^2 = \min_{\alpha \in \mathbb{R}^k} ||y - B\alpha||^2 \to \alpha = (B^T B)^{-1} B^T y$$

where as mentioned  $\alpha$  is the vector of  $\alpha_j$ 's, and each  $B_{kl} = h_l(x_k)$ 

b

$$S(x) = \begin{cases} S_0(x) & \text{if } x < \xi_1 \\ S_1(x) & \text{if } x \in [\xi_1, \xi_2) \\ S_2(x) & \text{if } x \ge \xi_2, \end{cases} = \alpha_0 + \alpha_1 x + \alpha_2 x^2 +_3 x^3$$

 $\mathbf{c}$ 

 $\mathbf{d}$ 

```
bone_mass <- read_delim("BoneMassData.txt", delim = " ")

females <- bone_mass %>% filter(gender == "female")

y <- females$spnbmd
x <- females$age</pre>
```

```
h_5 <- function(x, zeta_1 = 15) {ifelse((x - zeta_1) > 0, (x - zeta_1)^3, 0) }
h_6 <- function(x, zeta_2 = 20) {ifelse((x - zeta_2) > 0, (x - zeta_2)^3, 0) }

B <- matrix(
    c(rep(1, length(x)),
    x,
    x^2,
    x^3,
    h_5(x),
    h_6(x)),
    ncol = 6
)

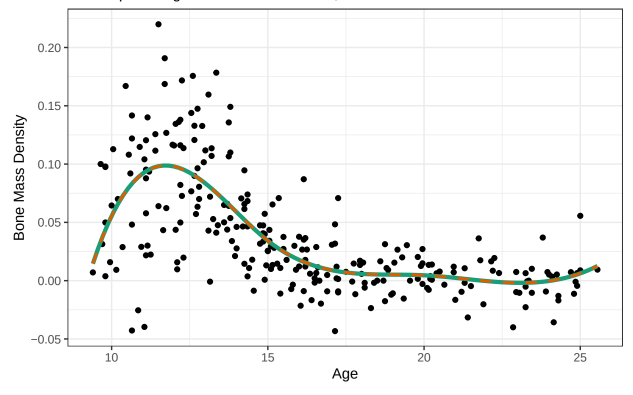
alpha <- solve(t(B) %*% B) %*% t(B) %*% y</pre>
```

So  $S(x) = -4.2488263 + 0.9762416x + -0.0716499x^2 + 0.0017069x^3 + -0.0021305[x - 15]_+^3 + 7.2804957 \times 10^{-4}[x - 20]_+^3$ 

```
S_x \leftarrow function(x, zeta_1 = 15, zeta_2 = 20, A = alpha) {
 H <- matrix(
   c(rep(1, length(x)),
    х,
   x^2,
   x^3,
   h_5(x),
   h 6(x)),
   ncol = 6
  Sx <- apply(H, 1, function(x) {sum(A*x)})</pre>
  return (Sx)
ggplot(data = females, aes(x = age, y = spnbmd)) +
 geom_point() +
 theme_bw() +
  stat_function(fun = S_x, color = "#1B9E77", size = 1.5) +
  geom\_smooth (method = "lm", formula = y ~ bs(x, knots=c(15,20)),
              alpha = 0, color = \#D95F02", linetype = 2, size = 1) +
  labs(x = "Age", y = "Bone Mass Density",
       title = "Bone Mass Density by Age",
       subtitle = "Cubic Spline Regression with knots at 15, 20")
```

## Bone Mass Density by Age

Cubic Spline Regression with knots at 15, 20



Here the solid teal line is our S(x) plotted over the data, and the dashed orange line is the spline regression formed using the bs function to compute the spline with knots at 15 and 20. As you can see, the regressions match up perfectly.

3

$$f(x) = e^x \to f'(x) = e^x \to f'(0) = 1$$

```
options(digits = 16)
i <- seq(-20,0,1)
h <- 10^i

finite_difference_1 <- function(x,h) {
   fx <- (exp(x + h) - exp(x)) / h

   return(fx)
}

finite_difference_2 <- function(x,h) {
   fx <- (exp(x + h) - exp(x - h)) / (2 * h)

   return(fx)
}

differences_1 <- sapply(h, finite_difference_1, x = 0)
differences_2 <- sapply(h, finite_difference_2, x = 0)</pre>
```

```
results <- data.frame(
  h = format(10^i, scientific = T, digits = 2),
  fin_diff_1 = differences_1,
  fin_diff_2 = differences_2
)

print(xtable::xtable(results, digits = 16), comment = F)</pre>
```

	h	fin_diff_1	fin_diff_2
1	1e-20	0.00000000000000000	0.00000000000000000
2	1e-19	0.000000000000000000	0.00000000000000000
3	1e-18	0.000000000000000000	0.00000000000000000
4	1e-17	0.00000000000000000	0.00000000000000000
5	1e-16	0.00000000000000000	0.5551115123125783
6	1e-15	1.1102230246251565	1.0547118733938987
7	1e-14	0.9992007221626409	0.9992007221626409
8	1e-13	0.9992007221626409	0.9997558336749535
9	1e-12	1.0000889005823410	1.0000333894311098
10	1e-11	1.0000000827403710	1.0000000827403710
11	1e-10	1.0000000827403710	1.0000000827403710
12	1e-09	1.0000000827403710	1.0000000272292198
13	1e-08	0.9999999939225290	0.9999999939225290
14	1e-07	1.0000000494336803	0.999999994736442
15	1e-06	1.0000004999621837	0.9999999999732445
16	1e-05	1.0000050000069649	1.0000000000121023
17	1e-04	1.0000500016671410	1.0000000016668897
18	1e-03	1.0005001667083846	1.0000001666666813
19	1e-02	1.0050167084167949	1.0000166667499921
20	1e-01	1.0517091807564771	1.0016675001984410
_21_	1e+00	1.7182818284590451	1.1752011936438014

## 4

In general, the cdf F(x) of a distribution is  $P(X \le x) \to P(-\infty < X \le x)$ . However here we have defined F(X) with a lower bound of 0. Since we know the standard normal distribution is centered at 0, with  $P(X \le 0) = 0.5$  and P(X > 0) = 0.5, then

$$F(x) = \int_0^x dz \frac{1}{\sqrt{2\pi}} e^{-z^2/2} \to F(\infty) = 0.5$$

Using R's built in integrate function, we see that an upper limit of 7.5 gives a value of 0.4999999999999883, accurate to 12 decimal places, so we set the upper limit for our subdivisions at 7.5. Increasing the upperbound beyond this will give us minimal added area and increase our error. However, we make this bound adjustable to the function's user.

```
norm_cdf <- function(z){1 / (sqrt(2 * pi)) * exp(-z^2 / 2)}
integrate(norm_cdf, 0, 7.5, subdivisions = 100)</pre>
```

0.499999999999883 with absolute error < 3.2e-07

```
Fapprox <- function(n, method = "reimann", upperlimit = 7.5) {</pre>
  `%notin%` = Negate(`%in%`)
  if (method %notin% c("reimann", "trapezoid", "useR")) {
    return("Please enter a valid integration method")
  }
  h <- upperlimit / n
  rectangle <- function(z) {
    height <-1 / sqrt(2 * pi) * exp(-(z^2) / 2)
    width <- h
    area <- height ★ width
    return (area)
  trapezoid <- function(z) {</pre>
    height_1 <- 1 / sqrt(2 * pi) * exp(-(z^2) / 2)
    height_2 <- 1 / sqrt(2 * pi) * exp(-((z + h)^2) / 2)
    area <- (height_1 + height_2) / 2 * h
  }
  if (method == "reimann") {
    cuts <- seq(0, upperlimit, h)
    norm_cdf <- sum(sapply(cuts[-n], rectangle))</pre>
  }else if (method == "trapezoid") {
    cuts <- seq(0, upperlimit, h)
    norm_cdf <- sum(sapply(cuts[-n], trapezoid))</pre>
  }else{
    norm\_cdf\_func \leftarrow function(z) \{1 / (sqrt(2 * pi)) * exp(-z^2 / 2)\}
    norm_cdf <- integrate(norm_cdf_func, 0,</pre>
                            upperlimit,
                            subdivisions = n)$value
  return (norm_cdf)
}
n_s < -c(10, 100, 1000, 10000)
evaluations <- data.frame(</pre>
  n = format(n_s, digits = 0),
  Riemann = sapply(n_s, Fapprox),
  Trapezoid = sapply(n_s, Fapprox, method = "trapezoid"),
  useR = sapply(n_s, Fapprox, method = "useR")
)
print(xtable::xtable(evaluations, digits = 16), comment = F)
```

Looking at the table of our results, we see that the rectangle method consistently overestimates the probability by a good amount, only reaching 3 digits of accuracy after we bump n to 10000. This is what we would expect as the normal curve is a decreasing function so we would consistently be adding additional area at each step. The trapezoid method is much more accurate, achieving 10 digits of accuracy at nust n=10, and almost reaching the same accuracy as R at n=10000. The accuracy of R's built in integral function did

	n	Riemann	Trapezoid	useR
1	10	0.6496033551123254	0.4999999999808031	0.499999999999883
2	100	0.5149603355149982	0.499999999999565	0.499999999999883
3	1000	0.5014960335514724	0.4999999999999680	0.499999999999883
4	10000	0.5001496033551186	0.4999999999999881	0.499999999999883

not change with varying values of n.