### statgen\_sibs

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```
library(tidyverse)
## Warning: package 'tidyverse' was built under R version 4.1.3
## -- Attaching packages ------ tidyverse 1.3.1 --
## v ggplot2 3.3.5 v purrr 0.3.4
## v tibble 3.1.4 v dplyr 1.0.7
## v tidyr 1.1.3 v stringr 1.4.0
## v readr 2.0.1 v forcats 0.5.1
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag() masks stats::lag()
library(arsenal)
## Warning: package 'arsenal' was built under R version 4.1.3
# reading phenotype data
phenotype <- read.csv("T2D-GENES_P1_Hispanic_phenotypes.txt", sep="")</pre>
head(phenotype)
##
                        IID DBP SBP BPMED SMOKE EXAM_YEAR AGE SEX
            FID
## 1 SAMPLE_0001 SAMPLE_0001 51 103 0 NA 2000 57 1
## 2 SAMPLE_0002 SAMPLE_0002 109 184 1 NA
## 3 SAMPLE_0003 SAMPLE_0003 92 182 1 NA
## 4 SAMPLE_0004 SAMPLE_0004 78 141 1 NA
                                                    1999 60
                                                    1999 50
                                                     2000 59
                                                               2
## 5 SAMPLE_0005 SAMPLE_0005 89 150 1 NA
                                                    1999 41
                                                               2
## 6 SAMPLE_0006 SAMPLE_0006 57 100 0 NA
                                                    1997 75
# reading genetic data
pcs <- read.csv("all_chr_eigensoft_PCs.csv")</pre>
head(pcs)
                    PC1
                            PC2
                                    PC3
                                            PC4
                                                    PC5
                                                            PC6
                                                                            PC8
##
             ID
                                                                    PC7
## 1 SAMPLE_0001 -0.0613 -0.0033 -0.0087 0.0194 0.0182 -0.0194 -0.0244 0.0149
## 2 SAMPLE_0002 -0.0144 -0.0126  0.0248 -0.0131 -0.0207 -0.0316  0.0166  0.0361
```

```
## 3 SAMPLE_0003 -0.0217 -0.0124 0.0176 -0.0160 -0.0186 -0.0101 0.0191 0.0103
## 4 SAMPLE_0004 -0.0130 -0.0079 -0.0013 0.0284 0.0304 -0.0025 0.0061 0.0117
## 5 SAMPLE 0005 -0.0136 -0.0163 -0.0433 -0.0469 -0.0417 0.0141 0.0006 -0.0175
## 6 SAMPLE_0006 -0.0185 -0.0139 0.0091 0.0035 0.0029 -0.0132 -0.0029 0.0035
        PC9
               PC10
## 1 0.0180 -0.0323
## 2 -0.0100 0.0269
## 3 -0.0093 -0.0054
## 4 -0.0001 -0.0201
## 5 0.0310 0.0236
## 6 -0.0091 -0.0102
phenotype <- subset(phenotype, select = -SMOKE) #removing smoke variable - all NAs
phenotype$HTN <- ifelse(phenotype$BP>130, 1, ifelse(is.null(phenotype$BPMED), 1, ifelse(phenotype$BPME
phenotype <- rename(phenotype, "ID" = "IID") # renaming IID in phenotype data to ID to merge with genet
phenotype <- merge(phenotype, pcs, by="ID") # merging genetic data by id var
head(phenotype)
                        FID DBP SBP BPMED EXAM_YEAR AGE SEX HTN
                                                                            PC2
## 1 SAMPLE_0001 SAMPLE_0001 51 103
                                               2000 57
                                                              0 -0.0613 -0.0033
                                        0
## 2 SAMPLE_0002 SAMPLE_0002 109 184
                                        1
                                               1999 60
                                                          1
                                                              1 -0.0144 -0.0126
## 3 SAMPLE_0003 SAMPLE_0003 92 182
                                               1999 50
                                                          2
                                                              1 -0.0217 -0.0124
## 4 SAMPLE_0004 SAMPLE_0004 78 141
                                               2000 59
                                                         2 1 -0.0130 -0.0079
                                        1
## 5 SAMPLE_0005 SAMPLE_0005 89 150
                                                          2
                                        1
                                               1999 41
                                                             1 -0.0136 -0.0163
## 6 SAMPLE_0006 SAMPLE_0006 57 100
                                        0
                                               1997 75
                                                         2
                                                              0 -0.0185 -0.0139
                PC4
##
        PC3
                        PC5
                                PC6
                                        PC7
                                                PC8
                                                        PC9
                                                               PC10
## 1 -0.0087 0.0194 0.0182 -0.0194 -0.0244 0.0149 0.0180 -0.0323
## 2 0.0248 -0.0131 -0.0207 -0.0316 0.0166 0.0361 -0.0100 0.0269
## 3 0.0176 -0.0160 -0.0186 -0.0101 0.0191 0.0103 -0.0093 -0.0054
## 4 -0.0013 0.0284 0.0304 -0.0025 0.0061 0.0117 -0.0001 -0.0201
## 5 -0.0433 -0.0469 -0.0417 0.0141 0.0006 -0.0175 0.0310 0.0236
## 6 0.0091 0.0035 0.0029 -0.0132 -0.0029 0.0035 -0.0091 -0.0102
colnames(phenotype)
  [1] "ID"
                    "FID"
                               "DBP"
                                           "SBP"
                                                       "BPMED"
                                                                   "EXAM_YEAR"
## [7] "AGE"
                    "SEX"
                               "HTN"
                                           "PC1"
                                                       "PC2"
                                                                   "PC3"
                                           "PC7"
                                                       "PC8"
                                                                   "PC9"
## [13] "PC4"
                   "PC5"
                               "PC6"
## [19] "PC10"
# Controls for summary stat tables
my_controls <- tableby.control(</pre>
 test = T,
 numeric.test="kwt", cat.test="chisq", #Specify Kruskal-Wallis test for numeric vars and chi square fo
 numeric.stats = c("meansd", "medianq1q3", "range", "Nmiss2"), #Display mean, sd, median, q1 and q3, r
 cat.stats = c("countpct", "Nmiss2"),
 stats.labels=list(
   meansd = "Mean (SD)",
   mediang1g3 = "Median (Q1, Q3)",
  range = "Min - Max",
```

```
Nmiss2 = "Missing"
  )
)
# Creating labels for variables in table
my_labels <- list(</pre>
 DBP = "Diastolic Blood Pressure",
  SBP = "Systolic Blood Pressure",
 AGE = "Age",
 SEX = "Sex",
  BPMED = "Blood Pressure Medication Status",
 HTN = "Hypertension Status"
)
# Creating a list of the PC variables
pc_list <- paste("PC", 1:10, sep="")</pre>
htn_fmla <- as.formula(paste("HTN ~ + DBP + SBP + AGE +", paste(pc_list, collapse= "+")))</pre>
sex_fmla <- as.formula(paste("SEX ~ + DBP + SBP + AGE +", paste(pc_list, collapse= "+")))</pre>
bpmed_fmla <- as.formula(paste("BPMED ~ + DBP + SBP + AGE +", paste(pc_list, collapse= "+")))</pre>
htn_table <- tableby(htn_fmla, data = phenotype, control=my_controls)</pre>
summary(htn_table, labelTranslations = my_labels, title = "Summary Statistics for Phenotypes By Hyperter
```

Table 1: Summary Statistics for Phenotypes By Hypertension

				p
	0 (N=1212)	1 (N=639)	Total $(N=1851)$	value
Diastolic Blood				<
Pressure				0.001
Mean (SD)	$70.144 \ (8.953)$	$79.776 \ (11.802)$	73.469 (11.022)	
Median (Q1, Q3)	70.500 (64.000,	79.000 (73.000,	73.000 (66.000,	
	76.000)	86.000)	80.000)	
Min - Max	32.000 - 98.000	37.000 - 123.000	32.000 - 123.000	
Missing	0	0	0	
Systolic Blood				<
Pressure				0.001
Mean (SD)	$112.889 \ (10.623)$	148.099 (16.249)	125.044 (21.103)	
Median (Q1, Q3)	114.000 (105.000,	143.000 (136.000,	122.000 (110.000,	
	121.000)	154.500)	137.000)	
Min - Max	66.000 - 130.000	130.500 - 213.000	66.000 - 213.000	
Missing	0	0	0	
Age				<
				0.001
Mean (SD)	$45.418\ (13.395)$	$57.025\ (12.589)$	$49.425 \ (14.233)$	
Median (Q1, Q3)	43.000 (35.000,	56.590 (49.000,	48.000 (38.000,	
	53.105)	66.000)	59.789)	
Min - Max	20.000 - 92.000	19.000 - 91.970	19.000 - 92.000	
Missing	0	0	0	
PC1				<
				0.001
Mean (SD)	$0.002 \ (0.022)$	$-0.002 \ (0.023)$	$0.001 \ (0.023)$	
Median (Q1, Q3)	0.005 (-0.015, 0.021)	-0.002 (-0.021, 0.018)	$0.003 \ (-0.017, \ 0.020)$	
Min - Max	-0.070 - 0.046	-0.053 - 0.044	-0.070 - 0.046	
Missing	0	0	0	

	0 (N=1212)	1 (N=639)	Total~(N=1851)	l valu
PC2				<
Mean (SD)	0.002 (0.024)	-0.002 (0.021)	0.001 (0.023)	0.00
Median (Q1, Q3)	-0.003 (-0.016, 0.014)	-0.002 (0.021)	-0.005 (-0.016, 0.012)	
Min - Max	-0.035 - 0.109	-0.008 (-0.010, 0.007)	-0.005 (-0.010, 0.012)	
Missing	-0.033 - 0.109	-0.034 - 0.097	-0.033 - 0.109	
PC3	U	U	U	0.090
Mean (SD)	-0.001 (0.023)	0.001 (0.023)	-0.000 (0.023)	0.03
Median (Q1, Q3)	0.001 (-0.016, 0.015)	0.001 (0.023)	0.001 (-0.015, 0.016)	
Min - Max	-0.073 - 0.065	-0.073 - 0.069	-0.073 - 0.069	
Missing	0.073 - 0.003	0.075 - 0.005	0.003	
PC4	O	U	U	0.15'
Mean (SD)	0.001 (0.023)	-0.001 (0.023)	0.000 (0.023)	0.10
Median (Q1, Q3)	0.002 (-0.014, 0.016)	-0.001 (-0.016, 0.015)	0.000 (0.025)	
Min - Max	-0.082 - 0.066	-0.069 - 0.063	-0.082 - 0.066	
Missing	0.002 - 0.000	0.003 - 0.003	0.002 - 0.000	
PC5	O	U	U	0.46'
Mean (SD)	0.001 (0.024)	-0.001 (0.021)	0.000 (0.023)	0.40
Median (Q1, Q3)	-0.000 (-0.014, 0.015)	0.001 (-0.014, 0.014)	0.000 (-0.014, 0.014)	
Min - Max	-0.080 - 0.098	-0.066 - 0.069	-0.080 - 0.098	
Missing	0.000 - 0.030	0.000 - 0.003	0.000 - 0.000	
PC6	O	U	U	0.714
Mean (SD)	-0.001 (0.024)	0.001 (0.020)	-0.000 (0.023)	0.11
Median (Q1, Q3)	0.002 (-0.012, 0.015)	0.002 (-0.010, 0.013)	0.002 (-0.011, 0.014)	
Min - Max	-0.122 - 0.057	-0.112 - 0.051	-0.122 - 0.057	
Missing	0.122 0.001	0.112 0.001	0.122 0.001	
PC7	O	v	V	0.55
Mean (SD)	0.000 (0.022)	-0.001 (0.024)	0.000 (0.023)	0.00
Median (Q1, Q3)	0.004 (-0.008, 0.014)	0.004 (-0.009, 0.014)	0.004 (-0.009, 0.014)	
Min - Max	-0.115 - 0.052	-0.121 - 0.053	-0.121 - 0.053	
Missing	0	0.121 0.000	0.121 0.000	
PC8	· ·	v	v	0.82
Mean (SD)	0.000 (0.023)	-0.000 (0.023)	-0.000 (0.023)	0.0_
Median (Q1, Q3)	0.001 (-0.015, 0.016)	0.000 (-0.016, 0.015)	$0.000 \ (-0.015, \ 0.016)$	
Min - Max	,	-0.071 - 0.075		
Missing	0	0	0	
PC9	Ç .	, and the second	·	0.13
Mean (SD)	0.000 (0.023)	-0.001 (0.023)	0.000 (0.023)	
Median (Q1, Q3)	0.000 (-0.014, 0.015)	-0.002 (-0.015, 0.012)	-0.001 (-0.014, 0.014)	
Min - Max	-0.072 - 0.106	-0.071 - 0.075	-0.072 - 0.106	
Missing	0	0	0	
PC10				0.729
Mean (SD)	-0.000 (0.022)	0.000(0.023)	-0.000 (0.023)	
Median (Q1, Q3)	-0.000 (-0.016, 0.015)	0.000 (-0.016, 0.016)	0.000 (-0.016, 0.016)	
Min - Max	-0.077 - 0.077	-0.075 - 0.072	-0.077 - 0.077	
Missing	0	0	0	

sex\_table <- tableby(sex\_fmla, data = phenotype, control=my\_controls)
summary(sex\_table, labelTranslations = my\_labels, title = "Summary Statistics for Phenotype Data By Sex</pre>

Table 2: Summary Statistics for Phenotype Data By Sex

	1 (N=702)	2 (N=1241)	Total (N=1943)	p value
Diastolic Blood				<
Pressure				0.001
Mean (SD)	$76.619 \ (11.939)$	$71.720 \ (10.067)$	$73.469 \ (11.022)$	
Median $(Q1, Q3)$	76.000 (69.000,	71.000 (65.000,	73.000 (66.000,	
	83.000)	78.000)	80.000)	
Min - Max	42.000 - 123.000	32.000 - 115.000	32.000 - 123.000	
Missing	41	51	92	
Systolic Blood				<
Pressure				0.001
Mean (SD)	127.914 (19.799)	123.450 (21.637)	125.044 (21.103)	
Median (Q1, Q3)	125.000 (114.000,	120.000 (107.000,	122.000 (110.000,	
	139.000)	136.000)	137.000)	
Min - Max	66.000 - 203.000	79.000 - 213.000	66.000 - 213.000	
Missing	41	51	92	
Age		-	-	<
6-				0.001
Mean (SD)	50.898 (14.243)	48.700 (14.188)	49.485 (14.243)	0.00-
Median (Q1, Q3)	50.110 (40.000,	47.000 (38.000,	48.000 (38.000,	
(4, 4,	61.000)	59.000)	60.000)	
Min - Max	19.000 - 87.000	22.190 - 92.000	19.000 - 92.000	
Missing	37	44	81	
PC1	51	11	O1	0.829
Mean (SD)	-0.000 (0.023)	0.000 (0.022)	-0.000 (0.023)	0.020
Median (Q1, Q3)	0.001 (-0.018, 0.020)	0.000 (0.022)	0.002 (-0.018, 0.019)	
Min - Max	-0.061 - 0.045	-0.070 - 0.046	-0.070 - 0.046	
Missing	0.001 - 0.049	0.070 - 0.040	0.070 - 0.040	
PC2	0	U	U	0.018
Mean (SD)	-0.001 (0.023)	0.001 (0.023)	0.000 (0.023)	0.010
Median (Q1, Q3)	-0.008 (-0.017, 0.009)	-0.004 (-0.016, 0.011)	-0.006 (-0.016, 0.010)	
Min - Max	-0.035 - 0.109	-0.004 (-0.010, 0.011)	-0.035 - 0.109	
Missing	0.035 - 0.105	0.034 - 0.033	0.033 - 0.103	
PC3	U	Ü	U	0.080
	0.001 (0.022)	0.001 (0.022)	0.000 (0.022)	0.060
Mean (SD)	0.001 (0.022) 0.002 (-0.013, 0.016)	-0.001 (0.023)	-0.000 (0.023)	
Median (Q1, Q3)	, ,	0.001 (-0.016, 0.015)	0.001 (-0.015, 0.016) -0.073 - 0.069	
Min - Max	-0.073 - 0.069	-0.073 - 0.065 0		
Missing PC4	0	U	0	0.367
	0.001 (0.022)	0.000 (0.022)	0.000 (0.022)	0.307
Mean (SD)	-0.001 (0.022)	0.000 (0.023)	0.000 (0.023)	
Median (Q1, Q3)	-0.001 (-0.015, 0.015)	0.001 (-0.015, 0.016)	0.000 (-0.015, 0.015)	
Min - Max	-0.069 - 0.054	-0.082 - 0.066	-0.082 - 0.066	
Missing	0	0	0	0.000
PC5	0.001 (0.000)	0.001 (0.000)	0.000 (0.000)	0.088
Mean (SD)	-0.001 (0.022)	0.001 (0.023)	-0.000 (0.023)	
Median (Q1, Q3)	-0.001 (-0.016, 0.013)	0.000 (-0.014, 0.016)	-0.000 (-0.014, 0.014)	
Min - Max	-0.080 - 0.070	-0.080 - 0.098	-0.080 - 0.098	
Missing	0	0	0	
PC6		0.051.75	0.0 ( )	0.224
Mean (SD)	0.001 (0.021)	-0.001 (0.023)	0.000 (0.023)	
Median $(Q1, Q3)$	$0.002 \ (-0.010, \ 0.015)$	0.002 (-0.012, 0.014)	0.002 (-0.011, 0.014)	
Min - Max	-0.112 - 0.054	-0.122 - 0.057	-0.122 - 0.057	

	1 (N=702)	2 (N=1241)	Total (N=1943)	p value
Missing	0	0	0	
PC7				0.208
Mean (SD)	$0.001 \ (0.023)$	-0.000 (0.022)	0.000 (0.023)	
Median (Q1, Q3)	0.004 (-0.008, 0.015)	0.003 (-0.009, 0.014)	0.004 (-0.009, 0.014)	
Min - Max	-0.121 - 0.053	-0.115 - 0.052	-0.121 - 0.053	
Missing	0	0	0	
PC8				0.787
Mean (SD)	$0.000 \ (0.023)$	-0.000 (0.023)	$0.000 \ (0.023)$	
Median (Q1, Q3)	0.001 (-0.015, 0.016)	0.001 (-0.015, 0.015)	0.001 (-0.015, 0.016)	
Min - Max	-0.072 - 0.075	-0.071 - 0.076	-0.072 - 0.076	
Missing	0	0	0	
PC9				0.540
Mean (SD)	-0.000 (0.023)	$0.000 \ (0.023)$	-0.000 (0.023)	
Median (Q1, Q3)	-0.001 (-0.015, 0.015)	-0.000 (-0.013, 0.013)	-0.001 (-0.014, 0.014)	
Min - Max	-0.068 - 0.106	-0.072 - 0.086	-0.072 - 0.106	
Missing	0	0	0	
PC10				0.491
Mean (SD)	-0.000 (0.023)	$0.000 \ (0.023)$	-0.000 (0.023)	
Median (Q1, Q3)	-0.000 (-0.015, 0.016)	$0.001 \ (-0.015, \ 0.016)$	$0.000 \ (-0.015, \ 0.016)$	
Min - Max	-0.075 - 0.072	-0.077 - 0.077	-0.077 - 0.077	
Missing	0	0	0	

bpmed\_table <- tableby(bpmed\_fmla, data = phenotype, control=my\_controls)
summary(bpmed\_table, labelTranslations = my\_labels, title = "Summary Statistics for Phenotype Data By B</pre>

Table 3: Summary Statistics for Phenotype Data By Blood Pressure Medication Status  $\,$ 

	0 (N 900)	1 (NT 147)	TD 1 (NI 407)	p
	0 (N=260)	1 (N=147)	Total (N=407)	value
Diastolic Blood				0.061
Pressure				
Mean (SD)	73.303 (11.161)	74.711 (11.126)	73.811 (11.155)	
Median (Q1, Q3)	73.000 (66.000,	76.000 (67.500,	74.000 (67.000,	
	80.625)	83.000)	81.750)	
Min - Max	49.000 - 115.000	42.000 - 109.000	42.000 - 115.000	
Missing	0	0	0	
Systolic Blood				<
Pressure				0.001
Mean (SD)	126.839 (18.159)	145.367 (21.082)	133.531 (21.203)	
Median (Q1, Q3)	125.000 (114.000,	143.000 (132.750,	131.000 (118.000,	
	136.125)	156.000)	145.000)	
Min - Max	91.000 - 197.000	100.000 - 212.000	91.000 - 212.000	
Missing	0	0	0	
$\mathbf{Age}$				<
				0.001
Mean (SD)	50.476 (13.539)	$61.490 \ (12.521)$	54.454 (14.191)	
Median (Q1, Q3)	48.990 (42.000,	62.248 (52.000,	53.000 (44.270,	
	59.000)	71.647)	64.703)	
Min - Max	19.000 - 88.630	35.000 - 91.970	19.000 - 91.970	

	0 (N=260)	1 (N=147)	Total $(N=407)$	value
Missing	0	0	0	
PC1				0.966
Mean (SD)	-0.014 (0.019)	-0.014 (0.018)	-0.014 (0.019)	
Median (Q1, Q3)	-0.015 (-0.027,	-0.015 (-0.026,	-0.015 (-0.026,	
	-0.003)	-0.004)	-0.003)	
Min - Max	-0.061 - 0.045	-0.052 - 0.040	-0.061 - 0.045	
Missing	0	0	0	
PC2				0.917
Mean (SD)	-0.014 (0.006)	-0.014 (0.007)	-0.014 (0.006)	
Median (Q1, Q3)	-0.015 (-0.018,	-0.015 (-0.018,	-0.015 (-0.018,	
1.1041411 (421, 469)	-0.011)	-0.011)	-0.011)	
Min - Max	-0.032 - 0.005	-0.029 - 0.015	-0.032 - 0.015	
Missing	0.002	0	0.002	
PC3	O	O	O	0.659
Mean (SD)	0.003 (0.021)	0.003 (0.022)	0.003 (0.022)	0.059
` ,			` ,	
Median (Q1, Q3)	0.004 (-0.011, 0.018)	0.003 (-0.012, 0.017)	0.003 (-0.012, 0.018)	
Min - Max	-0.056 - 0.055	-0.059 - 0.059	-0.059 - 0.059	
Missing	0	0	0	
PC4				<
				0.001
Mean (SD)	$0.004 \ (0.022)$	$-0.003 \ (0.022)$	$0.002 \ (0.022)$	
Median (Q1, Q3)	$0.006 \ (-0.009, \ 0.021)$	-0.004 (-0.017, 0.013)	0.003 (-0.013, 0.018)	
Min - Max	-0.061 - 0.053	-0.055 - 0.054	-0.061 - 0.054	
Missing	0	0	0	
PC5				0.416
Mean (SD)	$0.001 \ (0.021)$	-0.001 (0.022)	0.000(0.021)	
Median (Q1, Q3)	0.001 (-0.012, 0.016)	0.002 (-0.016, 0.014)	0.001 (-0.013, 0.015)	
Min - Max	-0.072 - 0.049	-0.060 - 0.056	-0.072 - 0.056	
Missing	0	0	0	
PC6	Ŭ	Ŭ	Ü	0.715
Mean (SD)	0.002 (0.017)	0.002 (0.018)	0.002 (0.017)	0.110
Median (Q1, Q3)	0.001 (-0.009, 0.012)	0.003 (-0.011, 0.014)	0.002 (0.017)	
Min - Max	-0.041 - 0.051	-0.033 - 0.042	-0.041 - 0.051	
	0.041 - 0.051	0.033 - 0.042	-0.041 - 0.031	
Missing	0	U	U	0.407
PC7	0.000 (0.010)	0.002 (0.001)	0.005 (0.010)	0.407
Mean (SD)	0.006 (0.018)	0.003 (0.021)	0.005 (0.019)	
Median (Q1, Q3)	0.007 (-0.004, 0.018)	0.006 (-0.007, 0.016)	0.007 (-0.004, 0.017)	
Min - Max	-0.063 - 0.053	-0.081 - 0.046	-0.081 - 0.053	
Missing	0	0	0	
PC8				0.307
Mean (SD)	$-0.000 \ (0.022)$	$-0.003 \ (0.023)$	$-0.001 \ (0.023)$	
Median (Q1, Q3)	0.000 (-0.016, 0.014)	-0.002 (-0.015, 0.011)	-0.001 (-0.016, 0.013)	
Min - Max	-0.061 - 0.076	-0.071 - 0.055	-0.071 - 0.076	
Missing	0	0	0	
PC9				0.219
Mean (SD)	0.000(0.019)	-0.003 (0.022)	-0.001 (0.020)	
Median (Q1, Q3)	0.000 (-0.013, 0.013)	-0.003 (-0.016, 0.009)	-0.001 (-0.014, 0.012)	
Min - Max	-0.046 - 0.066	-0.056 - 0.048	-0.056 - 0.066	
Missing	0	0	0	
PC10	V	V	U	0.922
Mean (SD)	0.003 (0.022)	0.003 (0.021)	0.003 (0.022)	0.344
	` ,	0.003 (0.021)		
Median (Q1, Q3)	0.003 (-0.012, 0.019)	[] [][]/[ [ 11 11 11 11 11 11 11 11 11 11 11 11	0.004 (-0.012, 0.018)	

	0 (N=260)	1 (N=147)	Total (N=407)	p value
Min - Max	-0.060 - 0.077	-0.054 - 0.052	-0.060 - 0.077	
Missing	0	0	0	

gen\_table <- tableby(~DBP + SBP + AGE, data=phenotype, control=my\_controls)
summary(gen\_table, labelTranslations = my\_labels, title="Summary Statistics for Phenotype Data")</pre>

Table 4: Summary Statistics for Phenotype Data

	Overall (N=1943)
Diastolic Blood Pressure	
Mean (SD)	73.469 (11.022)
Median (Q1, Q3)	73.000 (66.000, 80.000)
Min - Max	32.000 - 123.000
Missing	92
Systolic Blood Pressure	
Mean (SD)	125.044 (21.103)
Median (Q1, Q3)	122.000 (110.000, 137.000)
Min - Max	66.000 - 213.000
Missing	92
Age	
Mean (SD)	49.485 (14.243)
Median (Q1, Q3)	48.000 (38.000, 60.000)
Min - Max	19.000 - 92.000
Missing	81

summary(freqlist(~SEX, data=phenotype), title="Frequency Table By Sex for Phenotype Data", labelTransla

Table 5: Frequency Table By Sex for Phenotype Data

Sex	Freq	Cumulative Freq	Percent	Cumulative Percent
1	702	702	36.13	36.13
2	1241	1943	63.87	100.00

summary(freqlist(~BPMED, data=phenotype), title="Frequency Table By Blood Pressure Medication Status for

Table 6: Frequency Table By Blood Pressure Medication Status for Phenotype Data

Blood Pressure Medication Status	Freq	Cumulative Freq	Percent	Cumulative Percent
0	260	260	13.38	13.38
1	147	407	7.57	20.95
NA	1536	1943	79.05	100.00

Table 7: Frequency Table By Hypertension Status for Phenotype Data

Hypertension Status	Freq	Cumulative Freq	Percent	Cumulative Percent
0	1212	1212	62.38	62.38
1	639	1851	32.89	95.27
NA	92	1943	4.73	100.00

Hypertension has significant associations with DBP, SBP, AGE, PC1, PC2.

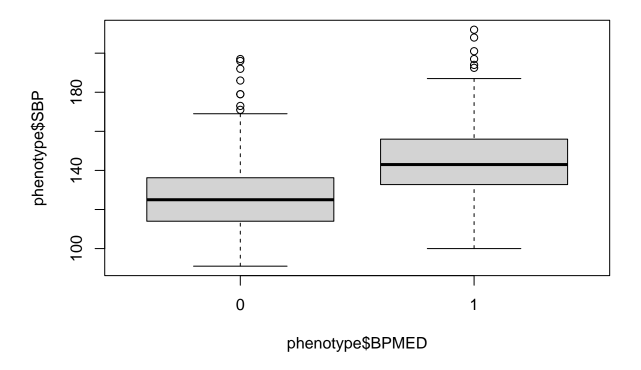
Sex has significant associations with DBP, SBP, AGE, and PC2

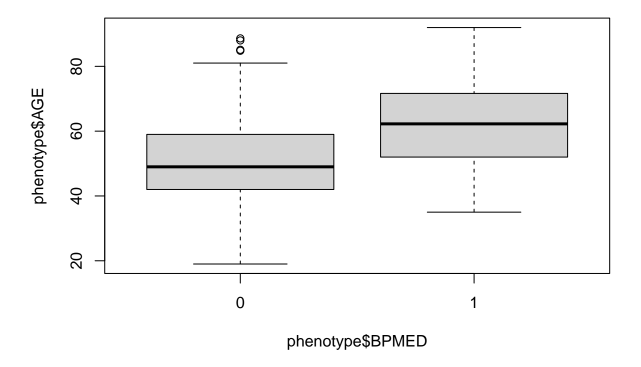
Blood Pressure Medication Status has significant associations with SBP, AGE, and PC4 (most significant PC yet). It makes sense for it not to have an association with DBP since high blood pressure is due to high SBP. It is interesting to note that hypertension does have a significant association with DBP. It appears that those with hypertension have a higher likelihood of having a high DBP.

There were only 3 signficant PCs in this analysis - PC1, PC2, and PC4.

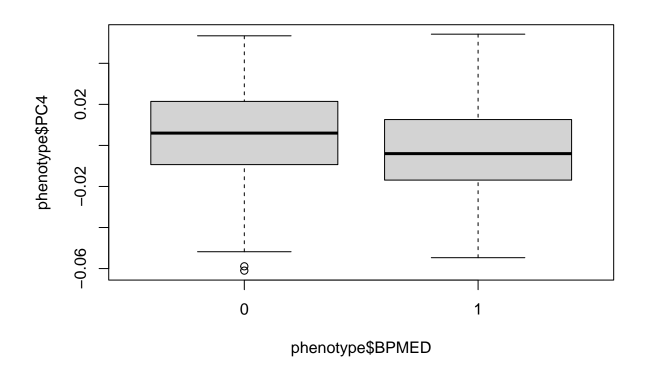
There are many more missing BPMED instances (1536) compared to missing SBP and DBP (92). This does not impact hypertension as it was coded to still code depending on SBP if BPMED was null.

#### boxplot(phenotype\$SBP~phenotype\$BPMED)



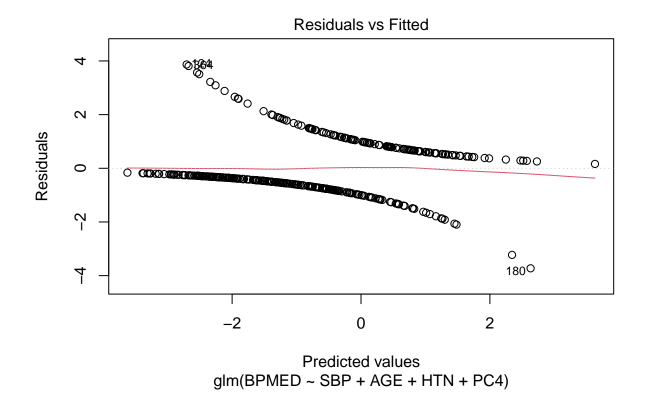


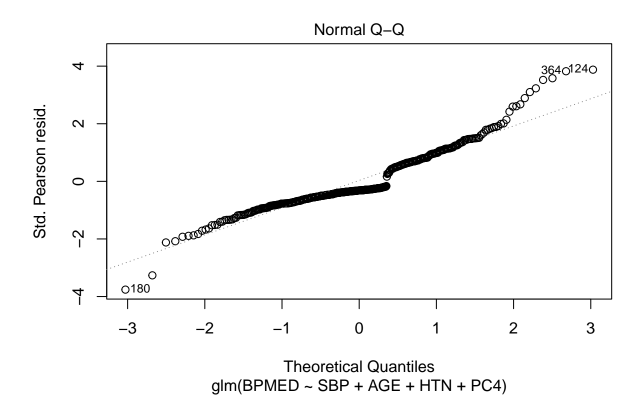
boxplot(phenotype\$PC4~phenotype\$BPMED)

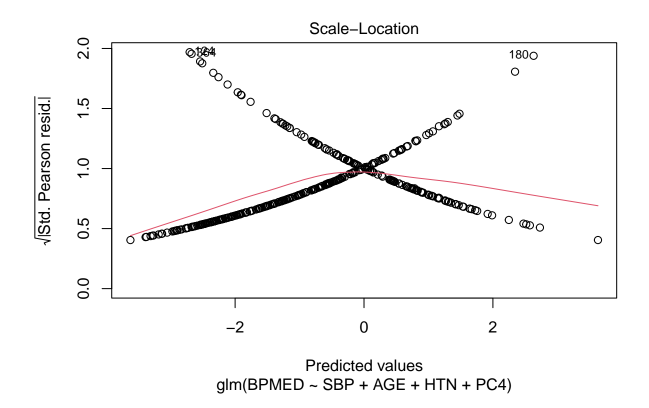


```
bpmed_fmla <- as.formula(paste("BPMED ~ SBP + AGE + SEX + BPMED + HTN + + ", paste(pc_list, collapse="+</pre>
bpmed_model <- glm(bpmed_fmla,family=binomial(link='logit'), data=phenotype)</pre>
## Warning in model.matrix.default(mt, mf, contrasts): the response appeared on the
## right-hand side and was dropped
## Warning in model.matrix.default(mt, mf, contrasts): problem with term 4 in
## model.matrix: no columns are assigned
#summary(bpmed_model)
bpmed_model_reduc <- glm(BPMED~SBP+AGE+HTN+PC4, family=binomial(link='logit'), data=phenotype)</pre>
summary(bpmed_model_reduc)
##
## Call:
## glm(formula = BPMED ~ SBP + AGE + HTN + PC4, family = binomial(link = "logit"),
##
       data = phenotype)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -2.3242 -0.7858 -0.4329
                               0.8566
                                         2.3539
##
## Coefficients:
```

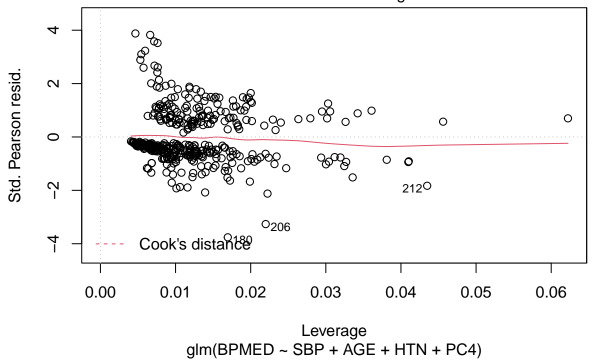
```
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -6.713073
                          1.206632 -5.563 2.64e-08 ***
                0.020317
                           0.009419
## SBP
                                      2.157 0.03100 *
                           0.009596
## AGE
                0.050556
                                      5.269 1.37e-07 ***
## HTN
                0.997735
                           0.373270
                                      2.673 0.00752 **
## PC4
              -15.366247
                           5.613926 -2.737 0.00620 **
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 532.43 on 406 degrees of freedom
## Residual deviance: 412.57 on 402 degrees of freedom
     (1536 observations deleted due to missingness)
## AIC: 422.57
##
## Number of Fisher Scoring iterations: 4
anova(bpmed_model_reduc, bpmed_model, test="Chisq")
## Analysis of Deviance Table
##
## Model 1: BPMED ~ SBP + AGE + HTN + PC4
## Model 2: BPMED ~ SBP + AGE + SEX + BPMED + HTN + +PC1 + PC2 + PC3 + PC4 +
      PC5 + PC6 + PC7 + PC8 + PC9 + PC10
##
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          402
                  412.57
## 2
          392
                  407.59 10
                              4.9798
                                       0.8925
exp(coef(bpmed_model_reduc))
## (Intercept)
                        SBP
                                      AGE
                                                   HTN
                                                                PC4
## 1.214925e-03 1.020525e+00 1.051856e+00 2.712132e+00 2.120918e-07
plot(bpmed_model_reduc) # Little shaky on assumptions of linearity, deviation from normality present, p
```







### Residuals vs Leverage



deviance(bpmed\_model\_reduc)/df.residual(bpmed\_model\_reduc)

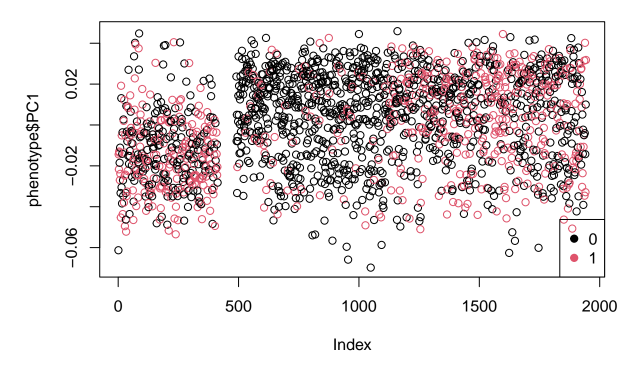
#### ## [1] 1.02629

The reduced model contains the variables SBP, AGE, HTN, and PC4 and appears to be a better fit model than the model with all variables as seen by the p-value of .8925 in the analysis of deviance table. Each unit increase in systolic blood pressure increases the odds of being on blood pressure medication by 1.02 (2%). Each year increase in age increases the odds of being on blood pressure medication by 1.05 (5%). Having hypertension increases the odds of being on blood pressure medication by 2.71 (171%)

It is important to note that logistic regression assumptions may not be met such as homoskedasticity as seen in the scale-location plot and normality as seen in the normal q-q plot. There are no influential outliers present as seen in the residuals vs leverage plot.

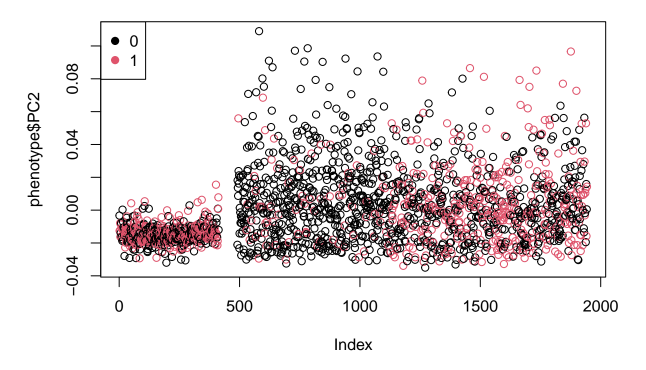
```
#Plotting significant PCs and dichotomous outcomes
plot(phenotype$PC1,col=factor(phenotype$HTN), main="PC1 grouped by hypertension")
legend("bottomright", legend=levels(factor(phenotype$HTN)), pch=19, col=factor(levels(factor(phenotype$.
```

## PC1 grouped by hypertension



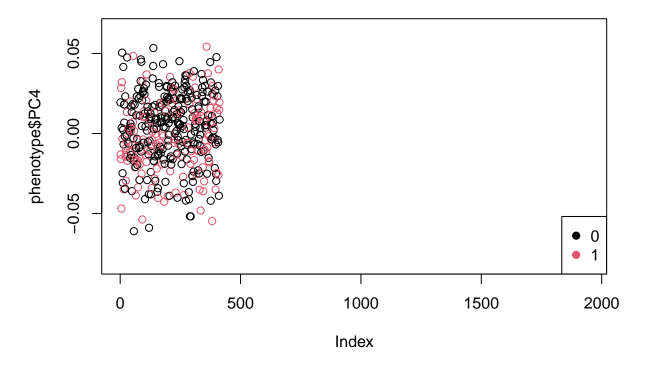
plot(phenotype\$PC2, col=factor(phenotype\$HTN), main="PC2 grouped by hypertension")
legend("topleft", legend=levels(factor(phenotype\$HTN)), pch=19, col=factor(levels(factor(phenotype\$HTN))

## PC2 grouped by hypertension



plot(phenotype\$PC4, col=factor(phenotype\$BPMED), main="PC4 grouped by BPMED")
legend("bottomright", legend=levels(factor(phenotype\$BPMED)), pch=19, col=factor(levels(factor(phenotype))

## PC4 grouped by BPMED



plot(phenotype\$PC2, col=factor(phenotype\$SEX), main="PC2 grouped by sex")
legend("topleft", legend=levels(factor(phenotype\$SEX)), pch=19, col=factor(levels(factor(phenotype\$SEX))

# PC2 grouped by sex

