Project1

1/15/2017

```
setwd("~/Documents/Biostatistics/Collaboration II")
totalData <- readRDS("dataForABP_CBF_2017-01-11.rds")</pre>
library(Hmisc)
library(knitr)
crvdata <- totalData[totalData$asl.reac.usable == 1,</pre>
                      c(1:18, 61, grep("asl.reac", names(totalData)))]
names(crvdata) <- sub("asl.reac.", "", names(crvdata), fixed = TRUE)</pre>
names(crvdata) <- sub(".hct", "", names(crvdata), fixed = TRUE)</pre>
exclude <- totalData[totalData$asl.reac.usable == 0,</pre>
                      c(1:18, 61, grep("asl.reac", names(totalData)))]
names(exclude) <- sub("asl.reac.", "", names(exclude), fixed = TRUE)</pre>
names(exclude) <- sub(".hct", "", names(exclude), fixed = TRUE)</pre>
#totalData$asl.reac.usable == 1,
cats <- names(crvdata)[2:19]</pre>
comparison \leftarrow c(c(), c(), c(), c())
is <- length(crvdata$map.id)</pre>
xs <- length(exclude$map.id)</pre>
for (cat in cats){
  if (is.factor(crvdata[,cat])){
    chiData <- rbind(cbind(crvdata[,cat],rep("is", length(crvdata[,cat]))),</pre>
                      cbind(exclude[,cat],rep("xs", length(exclude[,cat]))))
    pp <- chisq.test(table(chiData[,1], chiData[,2]))$p.value</pre>
    comparison <- rbind(comparison, c(label(crvdata[,cat]),'','', round(pp,4)))</pre>
    for (lev in levels(crvdata[,cat])){
      comparison <- rbind(comparison, c(paste("--",lev ),</pre>
                       paste(s <- sum(crvdata[,cat]==lev, na.rm=T), " (", round(s*100/is),</pre>
                              "%)", sep=""),
                       paste(s <- sum(exclude[,cat]==lev, na.rm=T), " (", round(s*100/xs),</pre>
                              "%)", sep=""),""))
    }
    next
  }
  anovaData <- as.data.frame(rbind(cbind(crvdata[,cat],rep("is", length(crvdata[,cat]))),</pre>
                      cbind(exclude[,cat],rep("xs", length(exclude[,cat])))))
  anovaData[,1] <- as.numeric(as.character(anovaData[,1]))</pre>
  pp <- kruskal.test(anovaData[,1] ~ anovaData[,2])$p.value</pre>
  comparison <- rbind(c(label(crvdata[,cat]),</pre>
                 paste(round(mean(crvdata[,cat], na.rm=T),1)," (",
                       round(sd(crvdata[,cat],na.rm=T),1), ")",sep=""),
                 paste(round(mean(exclude[,cat], na.rm=T),1)," (",
                       round(sd(exclude[,cat],na.rm=T),1), ")",sep=""),
                 round(pp,4)), comparison)
}
comparison <- as.data.frame(comparison[,c(1,3,2,4)])</pre>
colnames(comparison) <- c("Variable", "Excluded", "Analyzed Data", "P-Value")</pre>
```

Table 1: Comparison of Demographics for Excluded & Included Data, w/ N=0 and N=0 respectively

Variable	Excluded	Analyzed Data	P-Value
Diff. in mean SBP, wake - sleep, self-reported periods	14.5 (10.5)	13.4 (9.4)	0.6075
systolic.post.wake.mean minus systolic.pre.wake.mean	$11.1\ (12.3)$	$12.3\ (12.2)$	0.4331
systolic.post.wake.1 minus systolic.pre.wake.1	8.6(14.2)	$8.4\ (13.6)$	0.8162
ICV (calculated)	1431.2 (146.7)	1360.3 (134.6)	1e-04
Education (years)	16.5(2.5)	15.5(2.7)	9e-04
Age at medhx.date, recalculated	$73.1\ (7.1)$	72.8(7.4)	0.5341
Consensus Decision for Diagnosis			0.3736
- Normal	55 (49%)	121~(54%)	
- MCI	46 (41%)	86 (39%)	
- Dementia	1 (1%)	$0 \ (0\%)$	
– Ambiguous At Risk	11 (10%)	16(7%)	
Sex	, ,	,	0
- Male	90 (80%)	109 (49%)	
- Female	23 (20%)	114 (51%)	
Two-level race/ethnicity	,	,	0.9012
- Non-Hispanic White	97 (86%)	194 (87%)	
- Other	16 (14%)	29 (13%)	
ApoE4+ (at least one E4 allele)	- (/ 0)	_= (_=, _)	0.7178
- Yes	41 (36%)	75 (34%)	
- No	72 (64%)	148 (66%)	
Consensus Decision for Diagnosis	.= (01/0)	110 (00/0)	0.3736
- Normal	55 (49%)	121 (54%)	0.0100
- MCI	46 (41%)	86 (39%)	
- Dementia	1 (1%)	0 (0%)	
- Ambiguous At Risk	11 (10%)	16 (7%)	
Taking at least 1 anti-hypertensive med	11 (10/0)	10 (170)	0.9461
- Yes	62 (55%)	120 (54%)	0.3401
- No	51 (45%)	103 (46%)	
Diabetic, determined by a1c, glucose, and/or rx	01 (40/0)	100 (4070)	0.4304
- Yes	24 (21%)	38 (17%)	0.4304
- Tes - No	'	, ,	
	89 (79%)	185 (83%)	0.2549
Current smoker (or quit in this or last calendar yr)	4 (407)	2 (107)	0.3542
- Yes	4 (4%)	3 (1%)	
$-N_0$	109 (96%)	220 (99%)	0.007
CVD, determined from variables in med hx	9 (907)	0 (407)	0.897
- Yes	3 (3%)	8 (4%)	
- No	$110 \ (97\%)$	215 (96%)	4
A-fib, determined by med hx and/or echo and/or cmr rhythm	a (F (V))	10 (004)	1
- Yes	6 (5%)	13 (6%)	
$-N_0$	106 (94%)	209 (94%)	
LV hypertrophy, determined by sex and scaled LV mass	= (.~)	(204)	1
- Yes	5 (4%)	11 (5%)	
- No	108 (96%)	211 (95%)	
At least 39 readings AND max gap btw readings is <3h			0.0525

Variable	Excluded	Analyzed Data	P-Value
- No	6 (5%)	30 (13%)	
- Yes	91 (81%)	174~(78%)	