Package CompSign

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CompSign is a package for yadayada... overlooked that mutational signatures are compositional in nature yadayada. The reference manual can be found here.

```
knitr::opts_chunk$set(cache = FALSE)
```

```
## This chunk was last ran in
timestamp()
## ##---- Tue Nov 20 14:37:32 2018 ----##
## install latest version
library(devtools)
#devtools::install_github("lm687/CompSign")
library(CompSign)
library(compositions)
## Loading required package: tensorA
## Attaching package: 'tensorA'
## The following object is masked from 'package:base':
##
##
      norm
## Loading required package: robustbase
## Loading required package:
## Loading required package: bayesm
## Welcome to compositions, a package for compositional data analysis.
## Find an intro with "? compositions"
## Attaching package: 'compositions'
## The following objects are masked from 'package:stats':
##
      cor, cov, dist, var
##
## The following objects are masked from 'package:base':
##
      %*%, scale, scale.default
##
```

1 Summarise the signature matrix

```
## This chunk was last ran in
timestamp()
## ##---- Tue Nov 20 14:37:33 2018 ----##
add_together_matrix(sign_dummy)
## An object of class "sign"
## Slot "id":
## [1] "input_dummy"
##
## Slot "id_samples":
## [1] "sam1" "sam2" "sam3" "sam4"
##
## Slot "id_signatures":
## [1] "s1" "s2" "s3" "s4" "s5" "s6" "s7" "s8" "s9" "s10" "s11"
## [12] "s12" "s13" "s14" "s15" "s16" "s17" "s18" "s19" "s20" "s21" "s22"
## [23] "s23" "s24" "s25"
##
## Slot "count_matrix":
                        s2
                                  s3
## sam1 0.4727530 0.7201406 0.8626873 3.136111e-01 0.9059268 0.4715935
## sam2 0.4826859 0.9948616 0.8773814 3.650784e-05 0.5019962 0.7576524
## sam3 0.2523774 0.9881778 0.3274175 2.680551e-01 0.5750464 0.1806003
```

```
## sam4 0.3274017 0.9455770 0.8501642 7.948695e-01 0.3600513 0.9096839
               s7
                           s8
                                     s9
                                              s10
## sam1 0.6978941 0.872510572 0.1561251 0.6240480 0.73754948 0.3603802
## sam2 0.6557791 0.032059090 0.2069292 0.6067926 0.02658617 0.6702366
## sam3 0.1816618 0.005489897 0.4276585 0.5450189 0.06333866 0.3877280
## sam4 0.4866828 0.230437925 0.6783273 0.7602374 0.56992840 0.2220025
##
                         s14
                                              s16
              s13
                                    s15
                                                          s17
## sam1 0.2048943 0.24460129 0.75364351 0.2974315 0.80752923 0.4098196
## sam2 0.8481758 0.64424103 0.65221243 0.4387976 0.18614973 0.4897189
## sam3 0.8716103 0.17252595 0.50023542 0.0416386 0.08797126 0.3429558
## sam4 0.9638690 0.07959214 0.05546313 0.6837526 0.06263779 0.1150085
                       s20
                                  s21
                                            s22
## sam1 0.2600145 0.9672693 0.2792357 0.4716439 0.6954225 0.9449028
## sam2 0.4587714 0.3095924 0.9354094 0.3902753 0.2834248 0.4680484
## sam3 0.1881336 0.0212424 0.8787618 0.7476690 0.7910700 0.9811331
## sam4 0.3040355 0.4355955 0.4403698 0.1664639 0.8771529 0.6486360
##
               s25
## sam1 0.27511797
## sam2 0.80813636
## sam3 0.19301647
## sam4 0.07873073
##
## Slot "modified":
## [1] TRUE
results_sumarise <- summarise(add_together_matrix(sign_dummy))</pre>
results_sumarise$General
## [1] "Object of class sign"
```

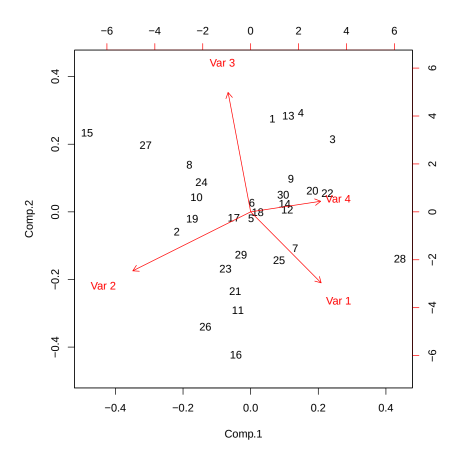
2 Linear model for numerical predictors

```
## [[1]]
## Response Y1 :
##
## Call:
## lm(formula = Y1 ~ as.matrix((x@df)[, indices_predictor]))
##
## Residuals:
      Min
               1Q Median
                                3Q
                                       Max
## -4.0614 -0.8163 -0.1171 1.0808 2.2565
##
## Coefficients: (1 not defined because of singularities)
                                             Estimate Std. Error t value
##
## (Intercept)
                                           -1.056e-02 5.052e-01 -0.021
## as.matrix((x@df)[, indices_predictor])a 1.538e-05 8.813e-05
                                                                   0.174
## as.matrix((x@df)[, indices_predictor])b
                                                 NA
                                                             NA
                                                                      NA
##
                                           Pr(>|t|)
## (Intercept)
                                              0.983
## as.matrix((x@df)[, indices_predictor])a
                                              0.863
## as.matrix((x@df)[, indices_predictor])b
##
## Residual standard error: 1.379 on 28 degrees of freedom
## Multiple R-squared: 0.001086, Adjusted R-squared: -0.03459
## F-statistic: 0.03044 on 1 and 28 DF, p-value: 0.8628
##
##
## Response Y2 :
##
## Call:
## lm(formula = Y2 ~ as.matrix((x@df)[, indices_predictor]))
##
## Residuals:
      Min
               1Q Median
                                3Q
                                       Max
## -3.0846 -0.9442 -0.1559 1.0345 2.3290
##
## Coefficients: (1 not defined because of singularities)
##
                                             Estimate Std. Error t value
## (Intercept)
                                           -8.977e-01 5.109e-01 -1.757
## as.matrix((x@df)[, indices_predictor])a 1.912e-04 8.913e-05
## as.matrix((x@df)[, indices_predictor])b
                                                   NA
                                                             NA
                                                                      NA
                                           Pr(>|t|)
## (Intercept)
                                             0.0899 .
## as.matrix((x@df)[, indices_predictor])a
                                             0.0408 *
## as.matrix((x@df)[, indices_predictor])b
                                                 NA
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
##
## Residual standard error: 1.394 on 28 degrees of freedom
## Multiple R-squared: 0.1411, Adjusted R-squared: 0.1105
## F-statistic: 4.601 on 1 and 28 DF, p-value: 0.04076
##
##
## Response Y3:
##
## Call:
## lm(formula = Y3 ~ as.matrix((x@df)[, indices_predictor]))
##
## Residuals:
               1Q Median
     Min
                               3Q
                                      Max
## -3.7103 -0.3316 0.1997 1.1198 2.3917
##
## Coefficients: (1 not defined because of singularities)
##
                                            Estimate Std. Error t value
## (Intercept)
                                           -4.082e-01 5.577e-01 -0.732
## as.matrix((x@df)[, indices_predictor])a 4.784e-05 9.728e-05 0.492
## as.matrix((x@df)[, indices_predictor])b
                                                 NA
                                                             NA
                                           Pr(>|t|)
##
## (Intercept)
                                              0.470
                                              0.627
## as.matrix((x@df)[, indices_predictor])a
## as.matrix((x@df)[, indices_predictor])b
                                                 NA
##
## Residual standard error: 1.522 on 28 degrees of freedom
## Multiple R-squared: 0.008561, Adjusted R-squared: -0.02685
## F-statistic: 0.2418 on 1 and 28 DF, p-value: 0.6268
```

3 Importing data

```
## This chunk was last ran in
timestamp()
## ##----- Tue Nov 20 14:37:35 2018 -----##
biplot(princomp(acomp(MCMCpack::rdirichlet(30, rep(1, 4)))))
```



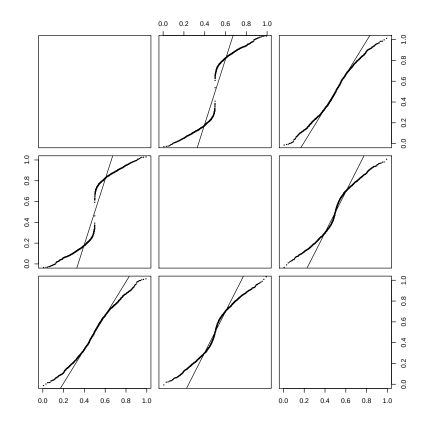
4 Other

1. Test for normality as follows:

```
## This chunk was last ran in
timestamp()

## ##----- Tue Nov 20 14:37:35 2018 -----##

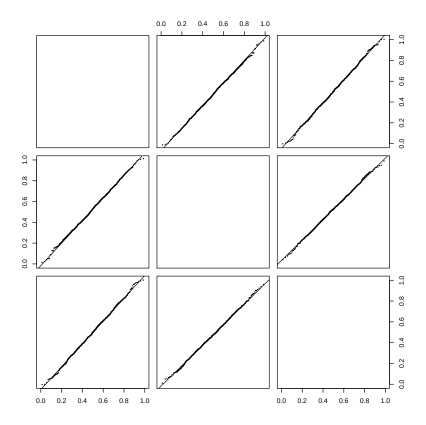
data(two_normal_pops)
par(mfrow=c(1,2))
qqnorm.acomp(acomp(two_normal_pops@count_matrix), pch=19, cex=0.2)
```



qqnorm.acomp(acomp(two_normal_pops@count_matrix[1:1000,]), pch=19, cex=0.2, plot.it=FAI
Warning in plot.window(...): "plot.it" is not a graphical
parameter
Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
Warning in title(...): "plot.it" is not a graphical parameter
Warning in plot.window(...): "plot.it" is not a graphical
parameter
Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
Warning in title(...): "plot.it" is not a graphical parameter
Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter

```
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
## Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
## Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
## Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter
```

```
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in plot.xy(xy.coords(x, y), type = type, ...): "plot.it"
is not a graphical parameter
## Warning in plot.window(...): "plot.it" is not a graphical
parameter
## Warning in plot.xy(xy, type, ...): "plot.it" is not a graphical
parameter
## Warning in title(...): "plot.it" is not a graphical parameter
## Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
## Warning in axis(side = side, at = at, labels = labels, ...):
"plot.it" is not a graphical parameter
```



5 Clustering of samples

5.1 Testing hypotheses about two populations

We might have our samples split into two categories; e.g. sex. As in Aithison 1986, I follow a hierarchy of alternative hypotheses, from least to most complex.

Our first question is whether two populations have the same covariance and structure and center (i.e. if there is any distributional difference)

```
## This chunk was last ran in
timestamp()
## ##----- Tue Nov 20 14:37:35 2018 -----##
##TODO!!
```

The next is whether the populations have a different center:

```
## This chunk was last ran in
timestamp()
## ##----- Tue Nov 20 14:37:35 2018 -----##
## This dataset includes the two components above, as well as four others
## (a total of seven)
data("two_normal_pops_extended")
## Data from the Landscape... paper
data("Breast560")
wrapper_compare_populations <- function(predictors, response, ...){</pre>
  if(length(unique(response)) == 2){
    tmp <- compare_populations(predictors, response, ...)</pre>
    tmp <- tmp$info[1:2]</pre>
    tmp
x <- do.call('rbind', lapply(1:ncol(metadata(Breast560)),</pre>
       function(k){
         wrapper_compare_populations(predictors = count_matrix(Breast560),
                                      response = metadata(Breast560)[,k])
       ))
## Loading required package: Compositional
##
## Attaching package: 'Compositional'
## The following object is masked from 'package:compositions':
##
##
      alr
```

```
## test p-value
## [1,] 215.35615 1.224333e-24
## [2,] 161.50152 3.522373e-18
## [3,] 235.90667 5.324172e-43
## [4,] 77.27718 1.042179e-11
```

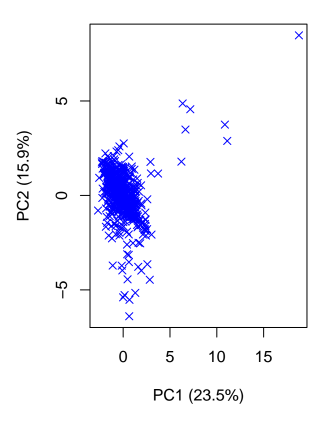
6 Data for 560 breast cancer patients

Data from 560 breast cancer patients is available as part of the document as well:

```
data("Breast560")
metadata(Breast560)[1:4,1:5]
##
           donor_gender donor_age_at_diagnosis donor_age_at_last_follow.up
## PD10010
                female
                                             56
                                                           no_data_supplied
## PD10011
                 female
                                             75
                                                           no_data_supplied
## PD10014
                 female
                                             64
                                                           no_data_supplied
## PD11326
                 female
                                             38
                                                                          47
            specimen_type donor_vital_status
## PD10010 tumour_primary
## PD10011 tumour_primary
                                        alive
## PD10014 tumour_primary
                                    deceased
## PD11326 tumour_primary
                                        alive
count_matrix(Breast560)[1:4,1:5]
##
           Signature.1 Signature.2 Signature.3 Signature.5 Signature.6
## PD10010 0.22386831 0.04197531
                                     0.2181070
                                                                      0
                                                  0.3374486
## PD10011
           0.09840426
                        0.00000000
                                      0.1791586
                                                  0.4514023
                                                                      0
                                                                      0
## PD10014
            0.03290722
                        0.00000000
                                      0.2859381
                                                  0.2812371
## PD11326 0.04040299 0.02596593
                                     0.3579144
                                                  0.1606772
```

Not sure if this is correct

```
source("../../CDA_in_Cancer/code/functions/basic_functions.R")
plotPCA(ilr(count_matrix(Breast560)), pch=4, col='blue')
## Loading required package: ggplot2
```



6.1 (ongoing) test for equality

```
## Sample 1 1.442203 1.255236 1.216817 0.9930453 1.012318
## Sample 2 1.535435 1.119812 1.252353 1.0633045 1.044466
##
## $info
## test p-value correction
## 2.790161e+02 9.100067e-51 1.046438e+00
## corrected.critical
## 2.058881e+01
```

7 Data for 12k TCGA samples, with ovarian cancer-derived CNA signatures

```
timestamp()
## ##----- Tue Nov 20 14:37:37 2018 -----##
data("CNA_12K_TCGA")
dim(metadata(CNA_12K_TCGA))
## [1] 10899 37
dim(count_matrix(CNA_12K_TCGA))
## [1] 10899 7
```

7.1 Logistic regression

Based on ACDWR pg $200\,$

```
#setwd("~/Documents/PhD/CompSign/vignette_knitr/")
load("../data/two_normal_pops_extended.rda")
load("../data/CNA_12K_TCGA.rda")

## avoid perfect separation
L <- length((as.numeric(metadata(two_normal_pops_extended)[,1])))
scramble <- sample(1:L, floor(L*0.05), replace = FALSE)

scrambled_labels <- (as.numeric(metadata(two_normal_pops_extended)[,1]))
scrambled_labels[scramble] <- 1-scrambled_labels[scramble]

auxcomp <- scale(ilr(count_matrix(two_normal_pops_extended)),</pre>
```

```
center = TRUE, scale = FALSE)
summary(glm(formula = scrambled_labels ~ ilr(acomp(count_matrix(two_normal_pops_extended)))
            family = binomial(link = "logit")))
##
## Call:
## glm(formula = scrambled_labels ~ ilr(acomp(count_matrix(two_normal_pops_extended))),
       family = binomial(link = "logit"))
##
## Deviance Residuals:
      Min
##
                1Q Median
                                   3Q
                                           Max
  -3.1008 -0.3067 -0.1343 0.3337
                                        3.1135
##
## Coefficients:
##
                                                       Estimate Std. Error
## (Intercept)
                                                        0.17669
                                                                 0.29572
## ilr(acomp(count_matrix(two_normal_pops_extended)))1 2.16946
                                                                   0.08293
## ilr(acomp(count_matrix(two_normal_pops_extended)))2  0.17045
                                                                   0.20491
## ilr(acomp(count_matrix(two_normal_pops_extended)))3 -0.04711
                                                                   0.13539
## ilr(acomp(count_matrix(two_normal_pops_extended)))4 -0.04876
                                                                   0.11905
## ilr(acomp(count_matrix(two_normal_pops_extended)))5 -0.15034
                                                                   0.10905
## ilr(acomp(count_matrix(two_normal_pops_extended)))6 -0.08281
                                                                   0.10105
## ilr(acomp(count_matrix(two_normal_pops_extended)))7 -0.02220
                                                                   0.09876
##
                                                       z value Pr(>|z|)
## (Intercept)
                                                         0.597
                                                                  0.550
## ilr(acomp(count_matrix(two_normal_pops_extended)))1 26.160
                                                                 <2e-16 ***
## ilr(acomp(count_matrix(two_normal_pops_extended)))2
                                                         0.832
                                                                  0.406
## ilr(acomp(count_matrix(two_normal_pops_extended)))3
                                                       -0.348
                                                                  0.728
## ilr(acomp(count_matrix(two_normal_pops_extended)))4
                                                       -0.410
                                                                  0.682
## ilr(acomp(count_matrix(two_normal_pops_extended)))5 -1.379
                                                                  0.168
## ilr(acomp(count_matrix(two_normal_pops_extended)))6 -0.819
                                                                  0.413
## ilr(acomp(count_matrix(two_normal_pops_extended)))7 -0.225
                                                                  0.822
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 2772.46 on 1999 degrees of freedom
## Residual deviance: 845.77 on 1992 degrees of freedom
## AIC: 861.77
##
## Number of Fisher Scoring iterations: 6
res <- comp_logistic(count_matrix(two_normal_pops), scrambled_labels)</pre>
```

Loading required package: nnet

```
## # weights: 5 (4 variable)
## initial value 1386.294361
## iter 10 value 436.460676
## iter 10 value 436.460676
## final value 436.460676
## converged
res
## $coefTransformed
              X1
                               X2
## (Intercept)
   0.2155614 -7.3456904 7.9497205 -0.3884687
##
## $summary
## Call:
## multinom(formula = as.formula(frm), data = dat)
##
## Coefficients:
                  Values Std. Err.
## (Intercept) 0.2155614 0.1370176
             -7.3456904 0.4188397
## X1
## X2
              7.9497205 0.3840895
## X3
              -0.3884687 0.4370199
##
## Residual Deviance: 872.9214
## AIC: 878.9214
##
## $res
## Call:
## multinom(formula = as.formula(frm), data = dat)
##
## Coefficients:
                     X1
## (Intercept)
                                X2
                                             Х3
##
   0.2155614 -7.3456904 7.9497205 -0.3884687
##
## Residual Deviance: 872.9214
## AIC: 878.9214
##
## [[4]]
##
   FP_FN_table
##
     0 1
   0 954 54
##
##
   1 46 946
resCNA_gender <- comp_logistic(count_matrix(cleanObject(CNA_12K_TCGA, 'gender')),</pre>
                 metadata(cleanObject(CNA_12K_TCGA, 'gender'))[,'gender'])
```

```
## # weights: 9 (8 variable)
## initial value 7535.896147
## iter 10 value 7477.896938
## final value 7477.896342
## converged
resCNA_gender
## $coefTransformed
## (Intercept) s1 s2 s3 s4
                                                               s5
## -0.2991671 0.2381913 -1.5952762 -0.7137485 -0.3424139
                                                         1.1287220
   s6 s7
##
## 1.1463550 -0.1609969
##
## $summary
## Call:
## multinom(formula = as.formula(frm), data = dat)
##
## Coefficients:
##
                Values Std. Err.
## (Intercept) -0.2991671 0.04634220
       0.2381913 0.07069156
## s1
## s2
            -1.5952762 0.34187887
## s3
            -0.7137485 0.15918132
## s4
            -0.3424139 0.22213187
## s5
             1.1287220 0.15577367
## s6
             1.1463550 0.25201363
## s7
            -0.1609969 0.19798834
## Residual Deviance: 14955.79
## AIC: 14969.79
##
## $res
## Call:
## multinom(formula = as.formula(frm), data = dat)
## Coefficients:
## (Intercept)
                s1 s2
                                        s3
                                               s4
## -0.2991671 0.2381913 -1.5952762 -0.7137485 -0.3424139 1.1287220
   s6 s7
   1.1463550 -0.1609969
##
##
## Residual Deviance: 14955.79
## AIC: 14969.79
##
## [[4]]
```

```
FP_FN_table
##
##
              0
##
     female 4268 1385
##
    male
          3520 1699
## below: incorrect
resCNA_race <- comp_logistic(count_matrix(cleanObject(CNA_12K_TCGA, 'race')),</pre>
                              metadata(cleanObject(CNA_12K_TCGA, 'race'))[,'race'], relax_b;
## # weights: 54 (40 variable)
## initial value 19480.008949
## iter 10 value 9749.868694
## iter 20 value 9542.043736
## iter 30 value 9517.674549
## iter 40 value 9516.466317
## final value 9516.429451
## converged
resCNA_race
## $coefTransformed
##
                                             (Intercept)
                                                                 s1
## asian
                                                2.675822 0.9362561 -3.734532
                                                3.126298 0.7315326 -2.226914
## black or african american
## native hawaiian or other pacific islander -2.905191 -0.9673919 -6.062152
## not reported
                                                3.312094 0.6140444 -1.617130
                                                4.865439 1.4609214 -2.578245
## white
##
                                                   s3
                                                              s4
                                                                        s5
## asian
                                             1.690197 -0.5707642 -1.252844
## black or african american
                                             2.138134 0.9178396 -2.071713
## native hawaiian or other pacific islander 2.202370 8.4638717 4.265265
                                             2.954000 0.9466145 -1.448352
## not reported
## white
                                             1.957366 0.1287871 -1.119340
##
                                                      s6
                                                               s7
## asian
                                               1.6057156 4.001795
## black or african american
                                              -0.0760912 3.713510
## native hawaiian or other pacific islander -12.1163351 1.309181
## not reported
                                              -1.9068323 3.769750
## white
                                               1.0245549 3.991395
##
## $summary
## Call:
## multinom(formula = as.formula(frm), data = dat)
## Coefficients:
                                             (Intercept)
##
                                                                            s2
```

```
## asian
                                                2.675822 0.9362561 -3.734532
## black or african american
                                                3.126298 0.7315326 -2.226914
## native hawaiian or other pacific islander
                                               -2.905191 -0.9673919 -6.062152
## not reported
                                                3.312094 0.6140444 -1.617130
## white
                                                4.865439 1.4609214 -2.578245
##
                                                   s3
                                                              s4
                                                                        s5
                                             1.690197 -0.5707642 -1.252844
## asian
## black or african american
                                             2.138134 0.9178396 -2.071713
## native hawaiian or other pacific islander 2.202370 8.4638717 4.265265
## not reported
                                             2.954000 0.9466145 -1.448352
## white
                                             1.957366 0.1287871 -1.119340
##
                                                      s6
                                                               s7
                                               1.6057156 4.001795
## asian
## black or african american
                                              -0.0760912 3.713510
## native hawaiian or other pacific islander -12.1163351 1.309181
## not reported
                                              -1.9068323 3.769750
## white
                                               1.0245549 3.991395
## Std. Errors:
##
                                             (Intercept)
                                                                s1
## asian
                                               0.3708243 0.7298643 2.416836
## black or african american
                                               0.3649411 0.7246340 2.359150
## native hawaiian or other pacific islander 1.0000360 2.0428540 5.431924
## not reported
                                               0.3629791 0.7220111 2.338804
## white
                                               0.3589692 0.7159120 2.311800
##
                                                            s4
                                                   s3
## asian
                                             1.478489 2.095677 1.280650
## black or african american
                                             1.467926 2.074927 1.272645
## native hawaiian or other pacific islander 2.601987 3.041292 2.183757
                                             1.459629 2.068598 1.261350
## not reported
## white
                                             1.446293 2.050308 1.247049
##
                                                   56
                                                            s7
## asian
                                             2.320487 1.965550
## black or african american
                                             2.311944 1.954391
## native hawaiian or other pacific islander 5.819587 3.151494
## not reported
                                             2.311425 1.946600
## white
                                             2.278335 1.929741
##
## Residual Deviance: 19032.86
## AIC: 19102.86
##
## $res
## Call:
## multinom(formula = as.formula(frm), data = dat)
```

```
## Coefficients:
##
                                             (Intercept)
                                                               s1
## asian
                                               2.675822 0.9362561 -3.734532
                                               3.126298 0.7315326 -2.226914
## black or african american
## native hawaiian or other pacific islander -2.905191 -0.9673919 -6.062152
## not reported
                                               3.312094 0.6140444 -1.617130
## white
                                               4.865439 1.4609214 -2.578245
##
                                                  s3
                                                             s4
                                                                       s5
                                             1.690197 -0.5707642 -1.252844
## asian
                                            2.138134 0.9178396 -2.071713
## black or african american
## native hawaiian or other pacific islander 2.202370 8.4638717 4.265265
## not reported
                                            2.954000 0.9466145 -1.448352
## white
                                             1.957366 0.1287871 -1.119340
##
                                                     s6
                                                              s7
## asian
                                              1.6057156 4.001795
## black or african american
                                             -0.0760912 3.713510
## native hawaiian or other pacific islander -12.1163351 1.309181
## not reported
                                             -1.9068323 3.769750
## white
                                              1.0245549 3.991395
##
## Residual Deviance: 19032.86
## AIC: 19102.86
##
## [[4]]
##
                                              FP_FN_table
##
                                                 0
##
    american indian or alaska native
                                                 27
##
    asian
                                               665
## black or african american
                                                910
## native hawaiian or other pacific islander
                                               13
##
   not reported
                                               1296
                                               7961
## white
```