acorn example in Rstudio

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Abstract

This document contains documentation for using acorn.

call up acorn

```
library('acorn')
```

check out acorn functions

```
lsf.str("package:acorn")

## annotateCpG : function (dnv0bject, CpGannotFile)
## calculateDeletionInsertionratio : function (x)
## calculateDeletionlengths : function (x)
## calculateMNVlengths : function (x)
## calculateTiNotatio : function (x)
## countsPerIndividual : function (x)
## countsPerIndividual : function (x)
## extractAutosomes : function (x)
## extractINDELs : function (x)
## extractINDELs : function (x)
## extractSNVs : function (x)
## extractSNVs : function (x)
## extractSNVs : function (x)
## extractX : function (x)
## extractX : function (x)
## extractY : function (x)
## parentalAge : function (parentalAgeObject)
## motherAge : function (parentalAgeObject)
## parentalAgeObject : function (counts, parentalData)
## readDNV : function (x)
```

load in test data

```
input <- \ readDNV("\sim/Documents/Github/acorn/inst/extdata/dnms\_from\_Ng\_et\_al\_2022\_Human\_Mutation\_paper.txt.gz") \\ head(input)
```

```
SAMPLE CHROM POS_B38 REFERENCE
                                                              ALTERNATE
## 1 HG01928 chr1 913941
## 2 HG03915 chr1 917676
## 3 HG03915 chr1 918783
## 4 HG00450 chr1 1216505
## 5 HG02257 chr1 1217502
## 6 HG00465 chr1 1366883
                                     G GGTGTGAATTGGTGTAGTGTGAATGAGT
##
                                                   ID
                                     chr1_913941_G_T
## 1
## 2
                                     chr1_917676_C_A
## 3
                                     chr1_918783_G_C
## 4
                                    chr1_1216505_A_G
                                    chr1_1217502_G_A
## 6 chr1_1366883_G_GGTGTGAATTGGTGTAGTGTGAATGAGT
```

```
## 'data.frame': 9741 obs. of 6 variables:

## $ SAMPLE : chr "HG01928" "HG03915" "HG090450" ...

## $ POS_B38 : int 913941 917676 918783 1216505 1217502 1366883 1765426 2332062 2645102 3355666 ...

## $ REFERENCE: chr "G" "C" "G" "A" ...

## $ ALTERNATE: chr "T" "A" "C" "G" ...

## $ 10 : chr "chr1_913941_6_T" "chr1_917676_C_A" "chr1_918783_6_C" "chr1_1216505_A_G" ...
```

extract SNVs

```
snvs <- extractSNVs(input)
nrow(snvs)
## [1] 8558</pre>
```

extract indels

```
indels <- extractINDELs(input)
nrow(indels)
## [1] 1183</pre>
```

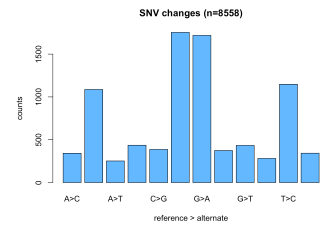
extract MNVs (there are none in the test set)

```
mnvs <- extractMNVs(input)
nrow(mnvs)
## [1] 0</pre>
```

calculate the Transition/Transversion ratio

```
calculateTiTvratio(input)

## number of transitions (A>G, C>T, G>A, T>C): 5708
## number of transversions (A>C, A>T, C>A, C>G, G>C, G>T, T>A, T>G: 2850
## Ti/Tv ratio: 2.00280701754386
## Plot of different nucleotide changes:
```

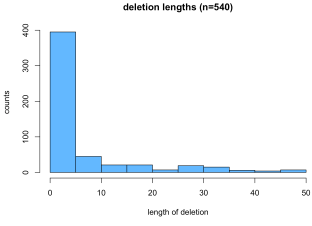


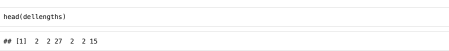
calculate deletion/insertion ratio

```
## number of deletions 540
## number of insertions 643
## deletion/insertion ratio: 0.839813374805599
```

calculate deletion lengths

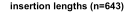
dellengths <- calculateDeletionlengths(input)</pre>

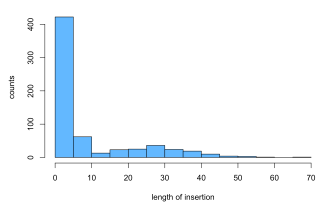




calculate insertion lengths

inslengths <- calculateInsertionLengths(input)</pre>





```
head(inslengths)
## [1] 28 3 37 37 41 5
```

keep only the autosomes

```
## [1] 9262

table(aut[,2])

##

## chrl chrl0 chrl1 chrl2 chrl3 chrl4 chrl5 chrl6 chrl7 chrl8 chrl9 chr2 chr20

## 707 408 477 408 338 479 251 273 233 232 206 833 257

## chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9

## 143 154 696 642 614 551 482 501 377
```

keep only the X chromosome

```
X <- extractX(input)
nrow(X)

## [1] 479

table(X[,2])

##
## chrX
## 479</pre>
```

keep only the Y chromosome (there are none on the Y in the test dataset)

```
Y <- extractY(input)
nrow(Y)
## [1] 0</pre>
```

annotate the CpG (note it takes ~45 seconds to even read the CpG annotation file so this function is a little slow)

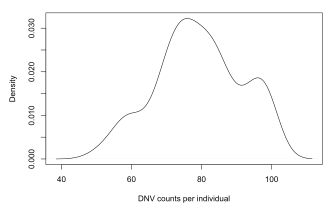
```
cpgResult <- annotateCpG(input, "~/Documents/Github/accessory_files/for_acorn/cpg_b38.rda")</pre>
## The number of DNVs at CpG sites is: 1566
## The total number of SNVs is: 8558
## The percent of DNV that are SNVs at CpG sites is: 18.2986679130638
head(cpgResult)
        SAMPLE CHROM POS_B38 REFERENCE ALTERNATE
                                                                            ID
                                                                                      matcher cpg
## 5 HG02257 chr1 1217502
                                                        A chr1_1217502_G_A chr1_1217502 yes
## 27 HG00526 chr1 6066908
## 29 HG02726 chr1 6280056
                                                        A chr1_6066908_G_A chr1_6066908 yes
A chr1_6280056_G_A chr1_6280056 yes
T chr1_6588427_G_T chr1_6588427 yes
                                            G
## 32 HG02135 chr1 6588427
                                                        C chr1_7354889_G_C chr1_7354889 yes
A chr1_7574692_G_A chr1_7574692 yes
## 34 HG02056 chr1 7354889
## 35 HG00465 chr1 7574692
```

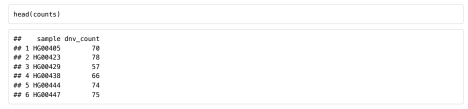
calculate counts per individual

```
counts <- countsPerIndividual(input)</pre>
```

```
## mean of the counts per individual:
## 79.19512
## standard deviation of the counts per individual:
## 12.20431
## Plot generating of the density of the DNV counts.
```

density of DNV counts per individual





load in example data for parental age analyses

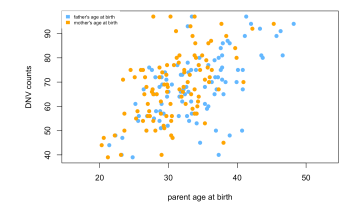
```
countExample <- read.delim("~/Documents/Github/acorn/inst/extdata/dnm_count_example.txt")
parentExample <- read.delim("~/Documents/Github/acorn/inst/extdata/parental_age_example.txt")</pre>
```

make parental age object

```
parents <- parentalAgeObject(countExample, parentExample)</pre>
```

run parental age analyses including both mother and father

parentalAge(parents)



```
## $`summary of linear model for father's and mother's age at birth and DNV counts
## Call:
## lm(formula = dnm counts ~ fatherAge + motherAge, data = parentalAgeObject)
## Residuals:
                1Q Median
                                  30
       Min
## -30.190 -6.831 -0.477 6.975 31.700
## Coefficients:
## Estimate Std. Error t value Pr(>|t|) ## (Intercept) 10.8374 7.7691 1.395 0.166220
## fatherAge
                  1.0331
                              0.2669
                                      3.871 0.000196 ***
## motherAge
                0.7179
                            0.3174 2.262 0.025955 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 11.14 on 97 degrees of freedom
## Multiple R-squared: 0.3732, Adjusted R-squared: 0.3603
## F-statistic: 28.87 on 2 and 97 DF, p-value: 1.45e-10
##
## $`confidence interval of linear model for father's and mother's age at birth and DNV counts` ## 2.5 \% 97.5 \%
                      2.5 %
## (Intercept) -4.58207432 26.256945
```

run parental age analyses for father age only

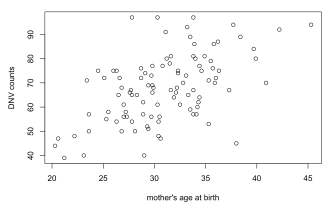
```
fatherAge(parents)
```



```
## $`correlation of father's age at birth and DNV counts`
## Pearson's product-moment correlation
## data: parentalAgeObject[, 3] and parentalAgeObject[, 2]
## t = 7.1073, df = 98, p-value = 1.927e-10
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.4368243 0.6994934
## sample estimates:
## 0.5832034
##
## $`summary of linear model for father's age at birth and DNV counts`
## Call:
## lm(formula = dnm_counts ~ fatherAge, data = parentalAgeObject)
## Residuals:
                 10 Median
                                   30
## -33.027 -6.690 -0.768 7.624 29.579
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 19.4131
                            6.9218 2.805 0.00608 **
0.2022 7.107 1.93e-10 ***
## fatherAge
                 1.4374
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 11.37 on 98 degrees of freedom
## Multiple R-squared: 0.3401, Adjusted R-squared: 0.3334
## F-statistic: 50.51 on 1 and 98 DF, p-value: 1.927e-10
## $`confidence interval of linear model for father's age at birth and DNV counts
## (Intercept) 5.676962 33.149157
## fatherAge 1.036030 1.838704
```

run parental age analyses for mother age only

motherAge(parents)



```
## $`correlation of mother's age at birth and DNV counts`
## Pearson's product-moment correlation
## rearSon's product-moment correctation

##

## data: parentalAgeObject[, 4] and parentalAgeObject[, 2]

## t = 6.1173, df = 98, p-value = 1.955e-08

## alternative hypothesis: true correlation is not equal to 0

## 95 percent confidence interval:

## 0.3671661 0.6545122
## sample estimates:
## cor
## 0.5256736
##
##
## $`summary of linear model for mother's age at birth and DNV counts`
##
## Call:
## \operatorname{Im}(\operatorname{formula} = \operatorname{dnm}_{\operatorname{counts}} \sim \operatorname{motherAge}, \operatorname{data} = \operatorname{parentalAgeObject}) ##
## Residuals:
## Min 10 Median 30 Max
## -33.793 -7.578 -0.467 7.825 33.927
## Coefficients:
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 20.2284 7.8898 2.564 0.0119 *
## motherAge
## ---
                          1.5412
                                             0.2519
                                                           6.117 1.96e-08 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1 ##
## Residual standard error: 11.91 on 98 degrees of freedom
## Multiple R-squared: 0.2763, Adjusted R-squared: 0.2689
## F-statistic: 37.42 on 1 and 98 DF, p-value: 1.955e-08
## $`confidence interval of linear model for mother's age at birth and DNV counts`
## 2.5 % 97.5 %
## (Intercept) 4.571389 35.88550
## motherAge 1.041211 2.04113
```

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