iMKT loading data

iMKT PopFly and PopHuman data

PopFly and PopHuman are. The package include functions to download all genes statics of all populations located in these genome browsers. Here we will explain how to load this information in your working environment and how to access it in order to use it efficiently!

Loading the row data

Remember you are downloading gene information of 20ypico *Droshopila melanogaster* populations and 26 Human populations. This process could take a while!

```
library(iMKT)
dataPopfly<-loadPopFly()
#> Loading PopFly data into your workspace.
#> This process may take some seconds to complete, please be patient.
knitr::kable(head(dataPopfly))
```

Pop	Name	Start	End	Chr	p0	pi	d0	di	m0	mi	alpha	fisher_pval	DoS	ŀ
EA	FBgn0000008	18024473	18060339	2R	28	17	36	19	647	2449	-0.1504	0.8348	-0.0323	0.
EA	FBgn0000014	12632936	12655771	3R	5	0	19	0	155	643	NA	1.0000	0.0000	0.
EA	FBgn0000015	12752932	12797958	3R	6	1	4	0	150	630	-Inf	1.0000	-0.1429	0.
EA	FBgn0000017	16608966	16640982	3L	10	5	83	134	867	3263	0.6903	0.0530	0.2842	1.
EA	FBgn0000018	10973443	10975293	2L	12	8	18	18	233	1019	0.3333	0.5796	0.1000	1.
EA	FBgn0000022	264064	264980	X	1	0	17	73	63	345	1.0000	0.1978	0.8111	4.

The data is presented at the most compress form in a simple dataframe. Each gene were analyzed by populations and contains information about segrating, divergent and analyzed sites, in addition to the Standard MKT calculation, Direction of Selection, Ka/Ks, and the Derived Allele Frequency (P0). The same information is provided by selected sites (Pi). If you have some basics R knowledge, you can access manually to the diverse gene information manually, in order to perform the test individually, by recombination information or concatenating them.

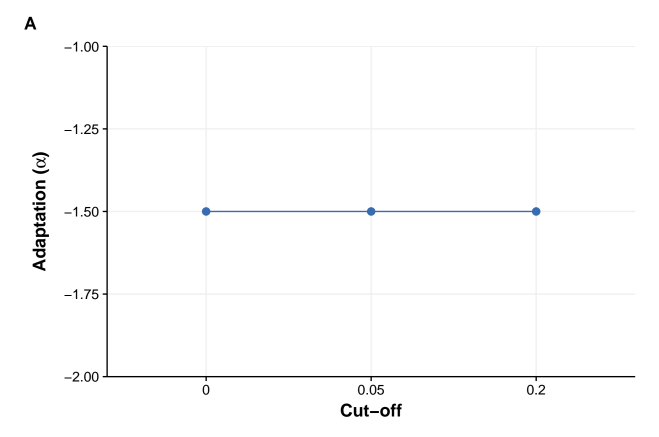
As we compress the Derived Allele Frequency in a column field, you will need to create the correct variables forms to pass them properly at the functions. It can be done in a few R basics lines!

```
## Preparing RAL Adh
adhRAL<-dataPopfly[dataPopfly$Name=='FBgn0000055' & dataPopfly$Pop=='RAL',]
adhRAL$DAF0f <- as.character(adhRAL$DAF0f); adhRAL$DAF4f <- as.character(adhRAL$DAF4f)
adhRAL0f<-unlist(strsplit(adhRAL$DAF0f, split=";"))
adhRAL4f <- unlist(strsplit(adhRAL$DAF4f, split=":"))
adhRAL0f <- as.numeric(adhRAL0f); adhRAL4f <- as.numeric(adhRAL0f)

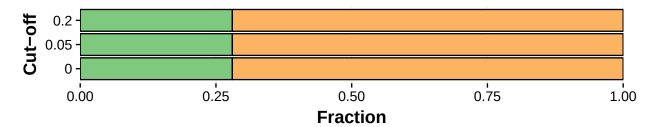
f <- seq(0.05,0.95,0.1)
mi <- adhRAL$mi; m0 <- adhRAL$m0
Di <- adhRAL$di; D0 <- adhRAL$d0

dafAdhRAL <- cbind(f, adhRAL0f, adhRAL4f); dafAdhRAL <- as.data.frame(dafAdhRAL)
names(dafAdhRAL) <- c("daf","Pi","P0")
divAdhRAL <- cbind(mi, Di, m0, D0); divAdhRAL <- as.data.frame(divAdhRAL)</pre>
```

```
names(divAdhRAL) <- c("mi", "Di", "m0", "D0")</pre>
## Preparing ZI Adh
adhZI<-dataPopfly[dataPopfly$Name=='FBgn0000055' & dataPopfly$Pop=='ZI',]
adhZI$DAFOf <- as.character(adhZI$DAFOf); adhZI$DAF4f <- as.character(adhZI$DAF4f)
adhZIOf<-unlist(strsplit(adhZI$DAFOf, split=";"))</pre>
adhZI4f <- unlist(strsplit(adhZI$DAF4f, split=":"))</pre>
adhZIOf <- as.numeric(adhZIOf); adhZI4f <- as.numeric(adhZIOf)
f \leftarrow seq(0.05, 0.95, 0.1)
mi <- adhZI$mi; m0 <- adhZI$m0
Di <- adhZI$di; DO <- adhZI$d0
dafAdhZI <- cbind(f, adhZIOf, adhZI4f); dafAdhZI <- as.data.frame(dafAdhZI)</pre>
names(dafAdhZI) <- c("daf","Pi","P0")</pre>
divAdhZI <- cbind(mi, Di, m0, D0); divAdhZI <- as.data.frame(divAdhZI)</pre>
names(divAdhZI) <- c("mi", "Di", "m0", "D0")</pre>
Once the data is in the proper form you can perform the test without problems
standard(daf = dafAdhRAL, divergence = divAdhRAL)
#> Warning in check_input(daf, divergence, 0, 1): Input daf file contains PO values = 0.
#> This can bias the function fitting and the estimation of alpha.
#> $alpha.symbol
#> [1] -1.5
#> $`Fishers exact test P-value`
#> [1] 0.6083916
#>
#> $`MKT table`
#>
#>
                   Polymorphism Divergence
                                           5
#> Neutral class
#> Selected class
#> $`Divergence metrics`
#>
#>
               Ks omega omegaA
#>
#> ------ -----
#> 0.00409 0.0364964 0.1120654 -0.1680982 0.2801636
DGRP(\underline{daf} = dafAdhRAL, divergence = divAdhRAL,plot = TRUE)
\#> Warning in check_input(daf, divergence, 0, 1): Input daf file contains PO values = 0.
#> This can bias the function fitting and the estimation of alpha.
#> $Results
#>
                alpha.symbol Fishers exact test P-value
\# Cutoff = 0
                  -1.5
                                 0.6083916
#> Cutoff = 0.05
                         -1.5
                                              0.6083916
#> Cutoff = 0.2
                        -1.5
                                              0.6083916
#>
#> $Graph
```





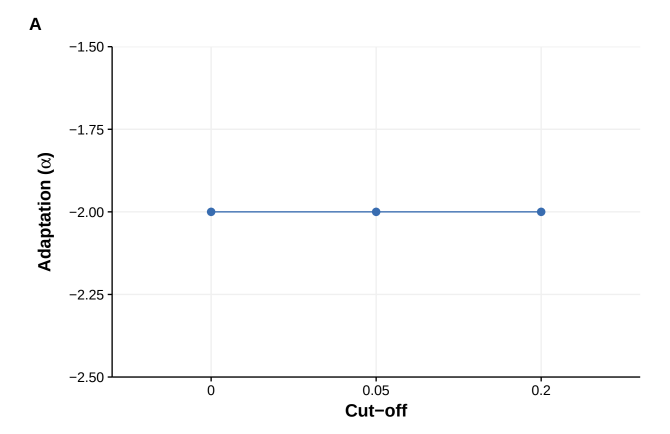


Fraction dffb

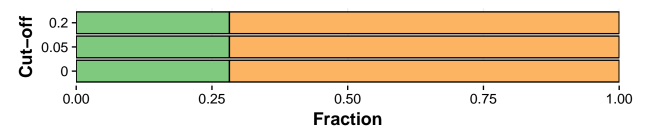
```
#>
#> $`Divergence metrics`
#> $`Divergence metrics`$`Global metrics`
                        Кs
#> 1 0.00408998 0.03649635 0.1120654
#> $`Divergence metrics`$`Estimates by cutoff`
                  omegaA.symbol omegaD.symbol
#>
\# Cutoff = 0
                     -0.1680982
                                    0.2801636
#> Cutoff = 0.05
                     -0.1680982
                                    0.2801636
#> Cutoff = 0.2
                     -0.1680982
                                    0.2801636
#>
#>
#> $`MKT tables`
```

```
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0`
#>
#>
#> Table: cutoff
#>
                   DAF.below.cutoff DAF.above.cutoff
                                0
#> Neutral class
#> Selected class
                                  0
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0.05`
#>
#>
#> Table: cutoff
#>
#>
                   DAF.below.cutoff DAF.above.cutoff
                                 3
#> Neutral class
#> Selected class
                                  3
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0.2`
#>
#> Table: cutoff
#>
                  DAF.below.cutoff DAF.above.cutoff
#> Neutral class
#> Selected class
#>
#> $`MKT tables`$`MKT standard table`
#>
#>
                  Polymorphism Divergence
#>
#> -----
                      4
#> Neutral class
#> Selected class
                            4
#>
#>
#> $Fractions
#> 0 0.05 0.2
#> d 0.7198364 0.7198364 0.7198364
#> f 0.2801636 0.2801636 0.2801636
#> b 0.0000000 0.0000000 0.0000000
standard(daf = dafAdhZI, divergence = divAdhZI)
\#> Warning in check_input \overline{(daf, divergence, 0, 1)}: Input daf file contains PO values = 0.
#> This can bias the function fitting and the estimation of alpha.
#> $alpha.symbol
#> [1] -2
#> $`Fishers exact test P-value`
#> [1] 0.3665158
#>
```

```
#> $`MKT table`
#>
#>
#>
                 Polymorphism Divergence
#> -----
#> Neutral class 5
#> Selected class 5
#>
#> $`Divergence metrics`
#>
#>
#> Ka Ks omega omegaA omegaD
#>-----
#> 0.00409 0.0434783 0.0940695 -0.1881391 0.2822086
DGRP(daf = dafAdhZI, divergence = divAdhZI,plot = TRUE)
\#> Warning in check_i nput(daf, divergence, <math>\overline{O, 1}): Input daf file contains PO values = 0.
#> This can bias the function fitting and the estimation of alpha.
#> $Results
#>
               alpha.symbol Fishers exact test P-value
\#> Cutoff = 0
                  -2
-2
                                           0.3665158
#> Cutoff = 0.05 -2
#> Cutoff = 0.2 -2
                                           0.3665158
                                           0.3665158
#>
#> $Graph
```







Fraction dffb

```
#>
#> $`Divergence metrics`
#> $`Divergence metrics`$`Global metrics`
             Ka
                        Ks
#> 1 0.00408998 0.04347826 0.09406953
#> $`Divergence metrics`$`Estimates by cutoff`
                  omegaA.symbol omegaD.symbol
#> Cutoff = 0
                     -0.1881391
                                    0.2822086
#> Cutoff = 0.05
                     -0.1881391
                                    0.2822086
#> Cutoff = 0.2
                     -0.1881391
                                    0.2822086
#>
#>
#> $`MKT tables`
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0`
```

```
#>
#>
#> Table: cutoff
#>
                 DAF.below.cutoff DAF.above.cutoff
#> ------
#> Neutral class
                              0
#> Selected class
#>
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0.05`
#>
#>
#> Table: cutoff
#>
#>
                 DAF.below.cutoff DAF.above.cutoff
                              5
#> Neutral class
                              5
#> Selected class
#> $`MKT tables`$`Number of segregating sites by DAF category - Cutoff = 0.2`
#>
#>
#> Table: cutoff
#>
#>
                DAF.below.cutoff DAF.above.cutoff
#> ----- ----
#> Neutral class
                              5
                              5
#> Selected class
#>
#> $`MKT tables`$`MKT standard table`
#>
#>
#>
                Polymorphism Divergence
                     5
5
#> Neutral class
                                      6
#> Selected class
#>
#>
#> $Fractions
   0 0.05 0.2
#>
#> d 0.7177914 0.7177914 0.7177914
#> f 0.2822086 0.2822086 0.2822086
#> b 0.0000000 0.0000000 0.0000000
```

This process could be tedious depending on the number of genes you want to analyze. To solve this, we just add the *PopFlyAnalisys()* function to the package. It allows you select a huge amount of genes and/or populations and retrieve the information easly! Remember to save the information in the most appropriate way to yours analysis or results!

```
PopFlyAnalysis(genes = c("FBgn0000055","FBgn0003016"),pops = c("RAL","ZI","FR"),recomb = F)
#> [1] "Population = FR"
#> Warning in check_input(daf, divergence, 0, 1): Input daf file contains PO values = O.
#> This can bias the function fitting and the estimation of alpha.
#> [1] "Population = RAL"
#> Warning in check_input(daf, divergence, 0, 1): Input daf file contains PO values = O.
```

```
#> This can bias the function fitting and the estimation of alpha.
#> [1] "Population = ZI"
#> $`Population = FR`
#> $`Population = FR`$alpha.symbol
#> [1] -0.3675214
#>
#> $`Population = FR`$`Fishers exact test P-value`
#> [1] 0.5274392
#>
#> $`Population = FR`$`MKT table`
#>
#>
                Polymorphism Divergence
#> -----
                               50
13
                45
16
#> Neutral class
                         16
#> Selected class
#>
#> $`Population = FR`$`Divergence metrics`
#>
#>
      Ka Ks omega omegaA omegaD
#>
#> 0.003767 0.058072 0.0648682 -0.0238404 0.0887086
#>
#>
#> $`Population = RAL`
#> $`Population = RAL`$alpha.symbol
#> [1] -1.197802
#>
#> $`Population = RAL`$`Fishers exact test P-value`
#> [1] 0.03712979
#>
#> $`Population = RAL`$`MKT table`
#>
#>
                Polymorphism Divergence
#> ------ -----
#> Neutral class
                  63
                                45
#> Selected class
                          40
                                     13
#>
#> $`Population = RAL`$`Divergence metrics`
#>
#>
#>
      Ka Ks omega omegaA omegaD
#> 0.003767 0.0522648 0.0720757 -0.0863325 0.1584082
#>
#>
#> $`Population = ZI`
\#> $`Population = ZI`$alpha.symbol
#> [1] -0.5279503
#> $`Population = ZI`$`Fishers exact test P-value`
#> [1] 0.2545795
```

```
#>
#> $`Population = ZI`$`MKT table`
#>
#>
                    Polymorphism
                                 Divergence
#> Neutral class
                             161
#> Selected class
                             82
                                          1.3
#>
#> $`Population = ZI`$`Divergence metrics`
#>
#>
          Ka
                     Ks
                              omega
                                          omeqaA
                                                      omeqaD
                          -----
#> 0.0037659
             0.0452436
                         0.0832368
                                      -0.0439449
                                                  0.1271817
```

Moreover PopFlyAnalisys() allows you to analyze a subset of genes by recombination bins, grouping them by the recombination values described by Comeron et al (ref). Remember you also can select the test to execute (for more information about the function check the manual or write ??PopFlyAnalysis() in your console)!

geneList <- c("FBgn0053196", "FBgn0086906", "FBgn0261836", "FBgn0031617", "FBgn0260965", "FBgn0028899",

```
PopFlyAnalysis(genes=geneList , pops=c("RAL","ZI"), recomb=T, bins=3, test="DGRP")
#> [1] "Population = RAL"
#> [1] "Recombination bin = 1"
#> [1] "Recombination bin = 2"
#> [1] "Recombination bin = 3"
#> [1] "Population = ZI"
#> [1] "Recombination bin = 1"
#> [1] "Recombination bin = 2"
#> [1] "Recombination bin = 3"
#> $`Population = RAL`
#> $`Population = RAL`$`Recombination bin = 1`
#> $`Population = RAL`$`Recombination bin = 1`$Results
#>
                 alpha.symbol Fishers exact test P-value
\# Cutoff = 0
                    0.3156666
                                            6.789846e-11
\# Cutoff = 0.05
                    0.4073286
                                            1.169402e-18
\# Cutoff = 0.2
                    0.3953727
                                            1.578535e-17
#>
#> $`Population = RAL`$`Recombination bin = 1`$`Divergence metrics`
#> $`Population = RAL`$`Recombination bin = 1`$`Divergence metrics`$`Global metrics`
          Ka
                  Ks
                             omega
#> 1 0.0417312 0.08882339 0.4698222
#>
#> $`Population = RAL`$`Recombination bin = 1`$`Divergence metrics`$`Estimates by cutoff`
                  omegaA.symbol omegaD.symbol
#>
\# Cutoff = 0
                    0.1483072
                                   0.3215150
\# Cutoff = 0.05
                     0.1913720
                                   0.2784502
\# Cutoff = 0.2
                    0.1857549
                                   0.2840674
#>
#>
#> $`Population = RAL`$`Recombination bin = 1`$`MKT tables`
\# \ `Population = RAL`$`Recombination bin = 1`$`MKT tables`$`Number of segregating sites by DAF categ
#>
```

```
#>
#> Table: cutoff
#>
#> Neutral class
                            0
                                          826
#> Selected class
                             0
                                         1202
\# ^* Population = RAL ^* Recombination bin = 1 ^* MKT tables ^* Number of segregating sites by DAF categ
#>
#>
#> Table: cutoff
               {\it DAF.below.cutoff} {\it DAF.above.cutoff}
#> ----- ----
                         375
#> Neutral class
                           707
#> Selected class
                                          495
#>
#> $`Population = RAL`$`Recombination bin = 1`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#> Table: cutoff
#>
                {\it DAF.below.cutoff} \quad {\it DAF.above.cutoff}
#> ------
                          464
#> Neutral class
#> Selected class
                          815
                                          387
#> $`Population = RAL`$`Recombination bin = 1`$`MKT tables`$`MKT standard table`
#>
#>
               Polymorphism Divergence
#> -----
#> Neutral class
                       826
                                 1115
#> Selected class 1202
                                 2371
#> $`Population = RAL`$`Recombination bin = 1`$Fractions
#> 0 0.05 0.2
#> d 0.678485 0.67840530 0.67854260
#> f 0.321515 0.27845021 0.28406737
#> b 0.000000 0.04314449 0.03739003
#> $`Population = RAL`$`Recombination bin = 1`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
#>
#>
\# $`Population = RAL`$`Recombination bin = 2`
#> $`Population = RAL`$`Recombination bin = 2`$Results
#> alpha.symbol Fishers exact test P-value
#> Cutoff = 0 0.02881113
                                    0.76379938
#> Cutoff = 0.05 0.14129004
                                     0.08157691
#> Cutoff = 0.2 0.13282389
                                     0.10747337
```

```
#>
#> $`Population = RAL`$`Recombination bin = 2`$`Divergence metrics`
#> $`Population = RAL`$`Recombination bin = 2`$`Divergence metrics`$`Global metrics`
#> Ka Ks
                           omega
#> 1 0.05678986 0.09626249 0.589948
#> $`Population = RAL`$`Recombination bin = 2`$`Divergence metrics`$`Estimates by cutoff`
#> omegaA.symbol omegaD.symbol #> Cutoff = 0 0.01699707 0.5729509
#> Cutoff = 0.05 0.08335377
                               0.5065942
#> Cutoff = 0.2 0.07835918 0.5115888
#>
#>
#> $`Population = RAL`$`Recombination bin = 2`$`MKT tables`
\# $`Population = RAL`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#>
#> Table: cutoff
#>
                  {\it DAF.below.cutoff} \quad {\it DAF.above.cutoff}
#> -----
#> Neutral class
                               0
                                0
#> Selected class
                                               803
#> $`Population = RAL`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#>
#> Table: cutoff
                 DAF.below.cutoff DAF.above.cutoff
#> ------
#> Neutral class
                              160
                              469
#> Selected class
                                               334
#>
\# $`Population = RAL`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#> Table: cutoff
#>
                 DAF.below.cutoff DAF.above.cutoff
#> -----
#> Neutral class
                             185
                                              157
                              520
#> Selected class
                                               283
#> $`Population = RAL`$`Recombination bin = 2`$`MKT tables`$`MKT standard table`
#>
#>
                 Polymorphism Divergence
#> Neutral class
                          342
                                      443
                       803
                                 1071
#> Selected class
#>
#>
#> $`Population = RAL`$`Recombination bin = 2`$Fractions
```

```
0 0.05 0.2
#> d 0.4270491 0.42681545 0.42731408
#> f 0.5729509 0.50659418 0.51158877
#> b 0.0000000 0.06659037 0.06109715
#> $`Population = RAL`$`Recombination bin = 2`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
        4 1.524576 2.018437 2.238379 3.392067
#>
#>
#> $`Population = RAL`$`Recombination bin = 3`
#> $`Population = RAL`$`Recombination bin = 3`$Results
     alpha.symbol Fishers exact test P-value
\# Cutoff = 0 0.4812584
                                 2.791715e-59
#> Cutoff = 0.05 0.5573293
                                       2.578578e-86
#> Cutoff = 0.2 0.5472239
                                        3.195443e-82
#>
#> $`Population = RAL`$`Recombination bin = 3`$`Divergence metrics`
#> $`Population = RAL`$`Recombination bin = 3`$`Divergence metrics`$`Global metrics`
                Ks
      Ka
                          omega
#> 1 0.0646586 0.1393272 0.4640773
#> $`Population = RAL`$`Recombination bin = 3`$`Divergence metrics`$`Estimates by cutoff`
               omegaA.symbol omegaD.symbol
#>
#> Cutoff = 0 0.2233411
                             0.2407362
#> Cutoff = 0.05 0.2586439 0.2054335
#> Cutoff = 0.2
                  0.2539542 0.2101231
#>
#>
#> $`Population = RAL`$`Recombination bin = 3`$`MKT tables`
\# $`Population = RAL`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#>
#> Table: cutoff
                  DAF.below.cutoff DAF.above.cutoff
#> ------
#> Neutral class
                               0
                                              1851
#> Selected class
                                0
                                              1848
#> $`Population = RAL`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#>
#> Table: cutoff
#>
                  DAF.below.cutoff DAF.above.cutoff
                              904
#> Neutral class
                                               947
#> Selected class
                             1174
                                                674
\# $`Population = RAL`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF categ
#>
#> Table: cutoff
```

```
#>
#>
                DAF.below.cutoff DAF.above.cutoff
#> ------
                                              764
#> Neutral class
                            1087
#> Selected class
                            1320
                                              528
#>
#> $`Population = RAL`$`Recombination bin = 3`$`MKT tables`$`MKT standard table`
#>
#>
#>
                 Polymorphism Divergence
#> -----
                        1851
                                    2746
#> Neutral class
#> Selected class
                        1848
                                    5285
#>
#>
#> $`Population = RAL`$`Recombination bin = 3`$Fractions
     0
                  0.05 0.2
#>
#> d 0.7592638 0.75920318 0.75929481
#> f 0.2407362 0.20543346 0.21012312
#> b 0.0000000 0.03536336 0.03058207
#>
#> $`Population = RAL`$`Recombination bin = 3`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
#>
#>
#>
#> $`Population = ZI`
#> $`Population = ZI`$`Recombination bin = 1`
#> $`Population = ZI`$`Recombination bin = 1`$Results
              alpha.symbol Fishers exact test P-value
#>
#> Cutoff = 0 -0.05192913
                                        0.4064121
#> Cutoff = 0.05 0.07538958
                                         0.1998081
#> Cutoff = 0.2 0.03676480
                                         0.5321975
#> $`Population = ZI`$`Recombination bin = 1`$`Divergence metrics`
#> $`Population = ZI`$`Recombination bin = 1`$`Divergence metrics`$`Global metrics`
#>
          Ka Ks
                           omega
#> 1 0.02203177 0.07151324 0.3080796
#>
#> $`Population = ZI`$`Recombination bin = 1`$`Divergence metrics`$`Estimates by cutoff`
               omegaA.symbol omegaD.symbol
\# Cutoff = 0
               -0.01599831 0.3240779
#> Cutoff = 0.05 0.02322599 0.2848536
#> Cutoff = 0.2 0.01132649
                            0.2967531
#>
#> $`Population = ZI`$`Recombination bin = 1`$`MKT tables`
\# \ `Population = ZI`$`Recombination bin = 1`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#> Table: cutoff
#>
                  DAF.below.cutoff DAF.above.cutoff
#>
```

```
#> Neutral class
                              0
                                             1575
                               0
#> Selected class
                                             2206
#>
\# ^* Population = ZI`$`Recombination bin = 1`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
#>
                 DAF.below.cutoff DAF.above.cutoff
#> -----
                                             522
#> Neutral class
                            1053
#> Selected class
                            1742
                                              464
\# $`Population = ZI`$`Recombination bin = 1`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
#>
                 {\it DAF.below.cutoff} {\it DAF.above.cutoff}
#> -----
#> Neutral class
                            1186
#> Selected class
                            1847
                                              359
#> $`Population = ZI`$`Recombination bin = 1`$`MKT tables`$`MKT standard table`
#>
#>
                 Polymorphism Divergence
#> Neutral class
                        1575
                                     724
#> Selected class
                         2206
                                     964
#>
#> $`Population = ZI`$`Recombination bin = 1`$Fractions
    0 0.05 0.2
#> d 0.6759221 0.6759028 0.67594459
#> f 0.3240779 0.2848536 0.29675310
#> b 0.0000000 0.0392436 0.02730231
#>
#> $`Population = ZI`$`Recombination bin = 1`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
#>
#>
#> $`Population = ZI`$`Recombination bin = 2`
#> $`Population = ZI`$`Recombination bin = 2`$Results
              alpha.symbol Fishers exact test P-value
#>
\# Cutoff = 0 0.1704018
                                      0.014492246
#> Cutoff = 0.05 0.2219296
                                        0.001046664
#> Cutoff = 0.2 0.2110515
                                        0.002012941
#> $`Population = ZI`$`Recombination bin = 2`$`Divergence metrics`
#> $`Population = ZI`$`Recombination bin = 2`$`Divergence metrics`$`Global metrics`
#>
        Ka
                Ks
                          omega
```

```
#> 1 0.05226333 0.08340555 0.626617
#>
#> $`Population = ZI`$`Recombination bin = 2`$`Divergence metrics`$`Estimates by cutoff`
#> omegaA.symbol omegaD.symbol
#> Cutoff = 0 0.1067766 0.5198403
#>
#>
#> $`Population = ZI`$`Recombination bin = 2`$`MKT tables`
\# $`Population = ZI`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
#>
                 DAF.below.cutoff DAF.above.cutoff
#> Neutral class
                              0
#> Selected class
                              0
                                            1449
#> $`Population = ZI`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
                DAF.below.cutoff DAF.above.cutoff
#> ------
                           446
                                            236
#> Neutral class
                           1038
#> Selected class
                                            411
#>
#> $`Population = ZI`$`Recombination bin = 2`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#> Table: cutoff
#>
                DAF.below.cutoff DAF.above.cutoff
#> ------
#> Neutral class
                           500
                                            182
#> Selected class
                           1133
                                            316
#>
#> $`Population = ZI`$`Recombination bin = 2`$`MKT tables`$`MKT standard table`
#>
#>
                Polymorphism Divergence
#> Neutral class
                        682
                                   385
                       1449
#> Selected class
                                    986
#>
#> $`Population = ZI`$`Recombination bin = 2`$Fractions
#> 0 0.05 0.2
#> d 0.4801597 0.48001131 0.48027328
#> f 0.5198403 0.48755212 0.49436853
#> b 0.0000000 0.03243656 0.02535819
```

```
#>
#> $`Population = ZI`$`Recombination bin = 2`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
#> 1 4 1.524576 2.018437 2.238379 3.392067
#>
#>
#> $`Population = ZI`$`Recombination bin = 3`
#> $`Population = ZI`$`Recombination bin = 3`$Results
              alpha.symbol Fishers exact test P-value
#>
#> Cutoff = 0 0.4217070
                                      3.382690e-61
#> Cutoff = 0.05 0.5070683
                                       7.771059e-98
#> Cutoff = 0.2 0.4848931
                                      5.523986e-87
#> $`Population = ZI`$`Recombination bin = 3`$`Divergence metrics`
#> $`Population = ZI`$`Recombination bin = 3`$`Divergence metrics`$`Global metrics`
#> Ka Ks omega
#> 1 0.06157707 0.1228686 0.501162
#> $`Population = ZI`$`Recombination bin = 3`$`Divergence metrics`$`Estimates by cutoff`
              omegaA.\mathit{symbol} omegaD.\mathit{symbol}
#> Cutoff = 0 0.2113435 0.2898185
#>
#>
#> $`Population = ZI`$`Recombination bin = 3`$`MKT tables`
\# $`Population = ZI`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
#>
                 {\it DAF.below.cutoff} {\it DAF.above.cutoff}
#> ------
                               0
#> Neutral class
                                              3611
#> Selected class
                               0
                                              4329
\# \ `Population = ZI`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
                 DAF.below.cutoff DAF.above.cutoff
                                  _____
#> -----
#> Neutral class
                            2275
                                             1336
                             3366
#> Selected class
                                               963
#>
#> $`Population = ZI`$`Recombination bin = 3`$`MKT tables`$`Number of segregating sites by DAF catego
#>
#>
#> Table: cutoff
#>
                DAF.below.cutoff DAF.above.cutoff
#> Neutral class
                             2600
                                              1011
```

```
#> Selected class
                          3590
                                          739
#>
#> $`Population = ZI`$`Recombination bin = 3`$`MKT tables`$`MKT standard table`
#>
          Polymorphism Divergence
#> ------
#> Neutral class 3611 2450
#> Selected class 4329 5079
#>
#>
#> $`Population = ZI`$`Recombination bin = 3`$Fractions
#> 0 0.05 0.2
#> d 0.7101815 0.71020522 0.71017991
#> f 0.2898185 0.24703862 0.25815201
#> b 0.0000000 0.04275616 0.03166808
#>
#> $`Population = ZI`$`Recombination bin = 3`$`Recombination bin Summary`
#> numGenes minRecomb medianRecomb meanRecomb maxRecomb
```