Burden

Partition enhancers based on distance to TSS based on strict overlap. Basically, her pipeline takes intersections between enhancers and distance ranges, and remove bases of enhancers that are not overlapped. ### NOTICE

getwd() ## [1] "/media/yuwen/F/ASD/analysis" knitr::opts_chunk\$set(warning=FALSE, message=FALSE) read in data with predicted mutation count and observed mutation count load in functions and datastes that make the functions work Set the scaling factor for nonysn to 0.32 for now (which is actually used for Polyphen-2 Mis3 rate) ASD manuscript data, stringent ASD genes predicted ## [1] 3.26384 observed ## [1] 3 ppois(observed-1, predicted,lower.tail = FALSE) ## [1] 0.63326 ASD manuscript data, relaxed ASD genes predicted ## [1] 20.93894 observed ## [1] 27

```
ppois(observed-1, predicted,lower.tail = FALSE)
## [1] 0.1147154
control\_model\_693\_mutrate\ data,\ stringent\ ASD\ genes
predicted
## [1] 5.474641
{\tt observed}
## [1] 7
ppois(observed-1, predicted,lower.tail = FALSE)
## [1] 0.3099843
control\_model\_693\_mutrate, relaxed ASD genes
predicted
## [1] 35.07132
observed
## [1] 46
ppois(observed-1, predicted,lower.tail = FALSE)
## [1] 0.04363395
```