# Using the DNaseI hypersensitivity data from encode in R

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#### 1 Introduction

Annotation tracks from UCSC hg18 can be used with Bioconductor to help establish genomic contexts of events or alterations. The CD4-based hypersensitivity assays are collected in the structure rawCD4 in package encoDnaseI:

```
> library(encoDnaseI)
> data(rawCD4)
> rawCD4
hg18track (storageMode: lockedEnvironment)
assayData: 382713 features, 1 samples
  element names: dataVals
protocolData: none
phenoData: none
featureData
  featureNames: 1 2 ... 382713 (382713 total)
  fvarLabels: bin chrom chromStart chromEnd
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
  pubMedIds: 16791207
Annotation:
   At present, we can subset the data by casting a chromosome number:
> c19g = rawCD4[ chrnum(19) ]
> c19g
hg18track (storageMode: lockedEnvironment)
assayData: 11158 features, 1 samples
  element names: dataVals
```

protocolData: none
phenoData: none
featureData

featureNames: 129572 129573 ... 140729 (11158 total)

fvarLabels: bin chrom chromStart chromEnd

fvarMetadata: labelDescription

experimentData: use 'experimentData(object)'

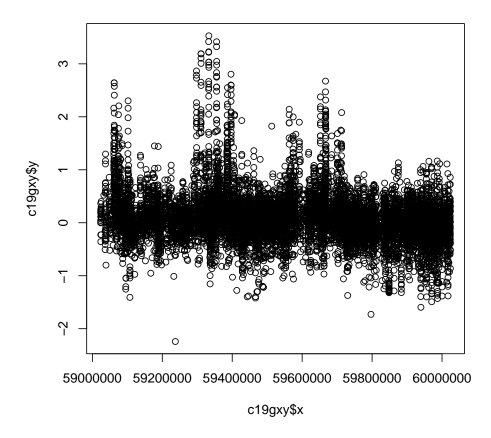
pubMedIds: 16791207

Annotation:

And we can get a trace of values along the chromosome:

> c19gxy = getTrkXY(c19g)

> plot(c19gxy)



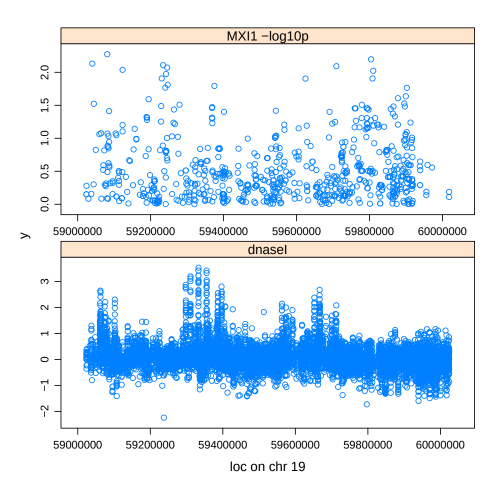
## 2 Coupling the DnaseI series to genetics of gene expression

We would like to subset a racExSet from GGdata and look at snps that are in regions of high DNaseI sensitivity. Some infrastructure to help with this is:

```
> clipSnps = function( sms, chrn, lo, hi ) {
+ allp = getSnpLocs(sms)
+ allp = allp-allp[1] # relative
+ ok = allp >= lo & allp <= hi
+ thesm = smList(sms)[[1]]
+ rsn = colnames(thesm)
+ rid = rsn[which(ok)]
+ thesm = thesm[, rid, drop=FALSE]
+ nn = new.env()
+ tmp = list(thesm)
+ names(tmp) = as.character(chrn)
+ assign("smList", tmp, nn)
+ sms@smlEnv = nn
+ sms@activeSnpInds=which(ok)
+ }
> rangeX = function(htrk) {
+ range(getTrkXY(htrk)$x)
+ }
  So we get the information on expression and SNPs in chr19g and filter:
> library(GGtools)
> library(GGdata)
> h19 = getSS("GGdata", "19")
> rs19g = rangeX(c19g)
> library(SNPlocs.Hsapiens.dbSNP.20090506)
> c191 = getSNPlocs("chr19")
> h19locs = rbind(rsid=as.numeric(c191[,"RefSNP_id"]), loc=as.numeric(c191[,"loc"]))
> #h19locs = getSnpLocs(hmceuB36[chrnum(19),])[[1]]
> goodlocs = which(h19locs[2,] >= rs19g[1] & h19locs[2,] <= rs19g[2])
> h19rsn = paste("rs", h19locs[1,goodlocs], sep="")
> h19trim = h19[rsid(h19rsn),]
> #ok
> #c19gf = clipSnps( hmceuB36[chrnum(19),], chrnum(19), rs19g[1], rs19g[2] )
> #c19gf
```

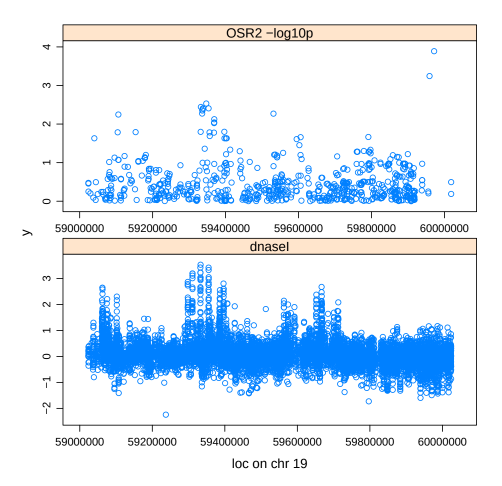
A gene-specific screen can be computed as follows:

```
> oo = options()  # don't take warnings on multiple probes... caveat emptor
> options(warn=0)
> library(GGtools)
> showMethods("gwSnpTests")
Function: gwSnpTests (package GGtools)
sym="formula", sms="smlSet"
> smxi1 = gwSnpTests(genesym("MXI1")~1-1, h19trim, chrnum(19))
[1] "GI_18641367-A" "GI_18641367-I" "GI_18641369-I"
> smxi1
gwSnpScreenResult for gene MXI1 [probe GI_18641367-A]
> #plot(smxi1)
> options(oo)
    We'd like to look at the SNP screen results juxtaposed with the DnaseI results.
> print(juxtaPlot( c19g, smxi1, h19locs ))
```



### Another example:

```
> oo = options()
> options(warn=0)
> sOSR2 = gwSnpTests(genesym("OSR2")~1-1, h19trim, chrnum(19))
> print(juxtaPlot( c19g, sOSR2, h19locs ))
> options(oo)
```



With these scores, we can find gene-snp combinations for which association is at least partly synchronized with DHS. Algorithms for systematically assessing synchronicity are in development.