# Package Vignette for Genomic Interactions: ChIA-PET data

## ChIA-PET

Chromatin interaction analysis with paired-end tag sequencing (ChIA-PET) is a recent method to study protein-mediated interactions at a genome-wide scale. Like most techniques for studying chromatin interaction it is based on chromosome conformation capture technology. Unlike 3C, 4C and 5C, however, it can detect interactions genome-wide, and includes a ChIP step to purify interactions involving a protein of interest.

The raw data from ChIA-PET is in the form of paired-end reads attached to one of two linker sequences. Reads with chimeric linkers are removed, and the data is aligned to the reference genome. The **ChIA-PET tool** can then be used to find pairs of regions ("anchors") which have a significant number of reads mapping between them and therefore represent biologically meaningful chromatin interactions in the sample.

#### Imports

First we need to load the GenomicInteractions package, and the mm9 reference genome:

```
library(GenomicInteractions)
library(GenomicRanges)
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
##
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
##
       clusterExport, clusterMap, parApply, parCapply, parLapply,
##
       parLapplyLB, parRapply, parSapply, parSapplyLB
##
  The following objects are masked from 'package:stats':
##
##
       IQR, mad, xtabs
## The following objects are masked from 'package:base':
##
##
       Filter, Find, Map, Position, Reduce, anyDuplicated, append,
##
       as.data.frame, as.vector, cbind, colnames, do.call,
##
       duplicated, eval, evalq, get, grep, grepl, intersect,
##
       is.unsorted, lapply, lengths, mapply, match, mget, order,
##
       paste, pmax, pmax.int, pmin, pmin.int, rank, rbind, rownames,
##
       sapply, setdiff, sort, table, tapply, union, unique, unlist,
##
       unsplit
```

```
## Loading required package: S4Vectors
## Loading required package: stats4
## Loading required package: IRanges
## Loading required package: GenomeInfoDb
```

#### Data

We can then read in our data directly from the output of the **ChIA-PET tool**. At this stage we can also provide information about the cell type and a description tag for the experiment. The data is taken from Li et al., 2012, published in **Cell**. They have used antibodies against the initiation form of Pol II, which you would expect to find at active promoters, and we are looking at data from the K562 myelogenous leukemia cell line. The data should therefore give us an insight into the processes which regulate genes that are being actively transcribed.

This loads the data into a GenomicInteractions object, which consists of two linked GenomicRanges objects containing the anchors in each interaction, as well as the p-value, FDR and the number of reads supporting each interaction.

#### **GenomicInteractions Objects**

The metadata we have added can easily be accessed, and edited:

```
name(k562.rep1)
```

```
## [1] "k562"
```

description(k562.rep1) = "PolII-8wg16 Chia-PET for K562"

As can the data from the ChIA-PET experiment:

head(interactionCounts(k562.rep1))

```
## [1] 3 562 3 3 3 3
```

```
head((k562.rep1)$fdr)
```

## [1] 1.25703e-10 0.00000e+00 1.17148e-06 4.86859e-08 2.76777e-08 3.97019e-08



# Histogram of -log10(k562.rep1\$p.value)

-log10(k562.rep1\$p.value)

The two linked **GRanges** objects can be returned, but not altered in-place:

```
anchorOne(k562.rep1)
```

##	GRanges	object w:	ith 64565	ranges	and $0$	meta	data colu	ımns	3:
##		seqname	es		rang	ges s	trand		
##		<rle< th=""><th>e&gt;</th><th>&lt;</th><th>IRange</th><th>es&gt;</th><th><rle></rle></th><th></th><th></th></rle<>	e>	<	IRange	es>	<rle></rle>		
##	[1	] chi	r1	[569922,	57142	22]	*		
##	[2	] chi	r1	[832761,	90548	32]	*		
##	[3	] chi	r1	[839092,	84232	25]	*		
##	[4	] chi	r1	[839393,	84179	92]	*		
##	[5	] chi	r1	[852731,	85523	34]	*		
##						• • •	•••		
##	[64561	] chi	rX [15443	2946, 15	6443572	28]	*		
##	[64562	] chi	rX [15443	6728, 15	443987	76]	*		
##	[64563	] chi	rX [15443	9789, 15	6444230	06]	*		
##	[64564	] chi	rX [15445	9648, 15	6446203	31]	*		
##	[64565	] chi	rX [15483	9050, 15	484394	19]	*		
##		-							
##	seqinf	o: 25 sec	quences f	rom an u	inspeci	ified	genome;	no	seqlengths

anchorTwo(k562.rep1)

## GRanges object with 64565 ranges and 0 metadata columns:
## seqnames ranges strand

## <Rle> <IRanges> <Rle> ## [1] [ 8342, 10675] chrM ## [2] chr1 [838470, 920603] [3] [935528, 939051] ## chr1 \* ## [4] chr1 [955081, 956755] ## [5] chr1 [933685, 937006] ## . . . . . . . . . chrX [154442294, 154446983] ## [64561] \* ## [64562] chrX [154442540, 154445105] ## [64563] chrX [154448371, 154451728] \* ## [64564] chrX [154469339, 154471852] chrX [154843728, 154848393] ## [64565] \* ## ## seqinfo: 25 sequences from an unspecified genome; no seqlengths

GenomicInteractions objects can easily handle interactions detected between chromosomes, known as *trans*-chromosomal interactions, since the anchors can be at any point along the genome. is.trans returns a logical vector; likewise is.cis is the opposite of this function.

#### ## [1] "Percentage of trans-chromosomal interactions 1.00"

The length of each interaction is not stored as metadata, but we can calculate the distance of each interaction using either the inner edge, outer edge or midpoints of the anchors. This is undefined for inter-chromosomal interactions, so NA is returned, so it is important to exclude these interactions from some analyses.

head(calculateDistances(k562.rep1, method="midpoint"))

#### ## [1] NA 10414 96580 115324 81362 79097

GenomicRanges objects can be subsetted by either integer or logical vectors like most R objects, and also BioConductor Rle objects.

k562.rep1[1:10] # first interactions in the dataset

##	Genomi	cInteractions object with 10 interactions and	1 2 meta	da	ata columns:
##	Name:	k562			
##	Descr	iption: PolII-8wg16 Chia-PET for K562			
##	Sum of	f interactions: 624			
##	Annota	ated: no			
##	Intera	actions:			
##		Anchor One Anchor Two	Counts	Ι	p.value
##	[1]	chr1:569922571422 chrM:834210675	3	T	1.6214e-12
##	[2]	chr1:832761905482 chr1:838470920603	562	T	0
##	[3]	chr1:839092842325 chr1:935528939051	3	T	4.21364e-08
##	[4]	chr1:839393841792 chr1:955081956755	3	T	1.45938e-09
##	[5]	chr1:852731855234 chr1:933685937006	3	T	7.85539e-10
##	[6]	chr1:855856858861 chr1:935669937245	3	Ι	1.16802e-09
##	[7]	chr1:874165879175 chr1:933340938306	10		1.23139e-25

```
##
      [8] chr1:874190..877867 --- chr1:955674..959630
                                                             5 | 6.63691e-15
##
      [9] chr1:889676..896594 --- chr1:933897..938982
                                                             13 | 4.91311e-36
     [10] chr1:898753..907581 --- chr1:931133..939571
##
                                                             19 |
                                                                            0
##
                  fdr
##
      [1] 1.25703e-10
##
      [2]
                    0
##
      [3] 1.17148e-06
      [4] 4.86859e-08
##
##
      [5] 2.76777e-08
      [6] 3.97019e-08
##
##
      [7] 3.58932e-23
##
      [8] 6.98795e-13
      [9] 2.33753e-33
##
     [10]
                    0
##
##
##
##
     seqinfo: 25 sequences from an unspecified genome; no seqlengths
# k562.rep1[sample(length(k562.rep1), 100)] # 100 interactions subsample
```

```
The length of each interaction is not stored as metadata, but we can calculate the distance of each interaction
```

using either the inner edge, outer edge or midpoints of the anchors. Since this is undefinable for *trans*chromosomal interactions it is best to first subset only *cis* interactions before calling calculateDistances, otherwise NAs will be present in the returned vector.

head(calculateDistances(k562.cis, method="midpoint"))

**##** [1] 10414 96580 115324 81362 79097 59152

k562.cis = k562.rep1[is.cis(k562.rep1)]

```
k562.short = k562.cis[calculateDistances(k562.cis) < 1e6] # subset shorter interactions
hist(calculateDistances(k562.short))</pre>
```



# Histogram of calculateDistances(k562.short)

calculateDistances(k562.short)

We can also subset based on the properties of the linked GRanges objects.

```
chrom = c("chr17", "chr18")
sub = as.vector(seqnames(anchorOne(k562.rep1)) %in% chrom & seqnames(anchorTwo(k562.rep1)) %in% chrom)
k562.rep1 = k562.rep1[sub]
```

## Annotation

Genomic Interaction data is often used to look at the interactions between different elements in the genome, which are believed to have different functional roles. Interactions between promoters and their transcription termination sites, for example, are thought to be a by-product of the transcription process, whereas long-range interactions with enhancers play a role in gene regulation.

Since GenomicInteractions is based on GenomicRanges, it is very easy to interrogate GenomicInteractions objects using GenomicRanges data. In the example, we want to annotate interactions that overlap the promoters, transcription termination sites or the body of any gene. Since this can be a time-consuming and data-heavy process, this example runs the analysis for only chromosomes 17 & 18.

First we need the list of RefSeq transcripts:

```
library(GenomicFeatures)
```

```
hg19.refseq.db <- makeTxDbFromUCSC(genome="hg19", table="refGene")
refseq.genes = genes(hg19.refseq.db)
refseq.transcripts = transcriptsBy(hg19.refseq.db, by="gene")
non_pseudogene = names(refseq.transcripts) %in% unlist(refseq.genes$gene_id)
refseq.transcripts = refseq.transcripts[non_pseudogene]</pre>
```

Rather than downloading the whole Refseq database, these are provided for chromosomes 17 & 18:

```
data("hg19.refseq.transcripts")
refseq.transcripts = hg19.refseq.transcripts
```

We can then use functions from GenomicRanges to call promoters and terminators for these transcripts. We have taken promoter regions to be within 2.5kb of an annotated TSS and terminators to be within 1kb of the end of an annotated transcript. Since genes can have multiple transcripts, they can also have multiple promoters/terminators, so these are GRangesList objects, which makes handling these objects slightly more complicated.

```
refseq.promoters = promoters(refseq.transcripts, upstream=2500, downstream=2500)
# unlist object so "strand" is one vector
refseq.transcripts.ul = unlist(refseq.transcripts)
# terminators can be called as promoters with the strand reversed
strand(refseq.transcripts.ul) = ifelse(strand(refseq.transcripts.ul) == "+", "-", "+")
refseq.terminators.ul = promoters(refseq.transcripts.ul, upstream=1000, downstream=1000)
# change back to original strand
strand(refseq.terminators.ul) = ifelse(strand(refseq.terminators.ul) == "+", "-", "+")
# `relist' maintains the original names and structure of the list
refseq.terminators = relist(refseq.terminators.ul, refseq.transcripts)
```

These can be used to subset a GenomicInteractions object directly from GRanges using the GenomicRanges overlaps methods. findOverlaps called on a GenomicInteractions object will return a list containing Hits objects for both anchors.

We can finds any interactions involving a RefSeq promoter:

```
subsetByFeatures(k562.rep1, refseq.promoters)
```

```
## GenomicInteractions object with 2907 interactions and 2 metadata columns:
  Name: k562
##
   Description: PolII-8wg16 Chia-PET for K562
##
   Sum of interactions: 58468
##
##
   Annotated: no
##
    Interactions:
##
                           Anchor One
                                                         Anchor Two Counts
##
        [1] chr18:32867581..32873274 --- chr18:32922822..32925514
                                                                          7
                                                                               [2] chr18:32868753..32872112 --- chr18:32951673..32954977
##
                                                                          4
                                                                               [3] chr18:32869486..32873870 --- chr18:32874778..32879603
##
                                                                              13
        [4] chr18:32869839..32873068 --- chr18:32879912..32884536
##
                                                                          4
                                                                               T
##
        [5] chr18:47003048..47019610 --- chr18:47008665..47025005
                                                                         85
                                                                               ##
                                  . . . . . . .
                                                                 . . .
        . . .
                                                                         . . . . . . .
     [2903] chr17:65222692..65227179 --- chr17:65239229..65241334
##
                                                                          4
                                                                               ##
     [2904] chr17:65231015..65237268 --- chr17:65235900..65244024
                                                                         12
                                                                               Ι
##
     [2905] chr17:45248915..45264031 --- chr17:45257291..45268317
                                                                         43
                                                                               T
##
     [2906] chr17:60130322..60137699 --- chr17:60137079..60144580
                                                                         21
                                                                               1
##
     [2907]
                chr18:227929..232043 ---
                                               chr18:265870..268355
                                                                          3
                                                                               1
##
                p.value
                                 fdr
##
        [1] 2.51251e-23 6.28044e-21
        [2] 4.69817e-16 5.97214e-14
##
##
        [3]
                      0
                                   0
##
        [4] 3.61832e-16 4.67152e-14
##
                      0
        [5]
                                   0
##
        . . .
                     . . .
                                 . . .
```

```
##
     [2903] 3.21725e-18 5.20359e-16
##
                       0
                                    0
     [2904]
##
     [2905]
                       0
                                    0
                       0
                                    0
##
     [2906]
##
     [2907] 1.57957e-11 9.55328e-10
##
##
##
     seqinfo: 25 sequences from an unspecified genome; no seqlengths
```

However, one of the most powerful features in the GenomicInteractions package is the ability to annotate each anchor with a list of genomic regions and then summarise interactions according to these features. This annotation is implemented as metadata columns for the anchors in the GenomicInteractions object and so is fast, and facilitates more complex analyses.

The order in which we annotate the anchors is important, since each anchor can only have one node.class. The first listed take precedence. Any regions not overlapping ranges in annotation.features will be labelled as distal.

```
## Annotating with promoter ...
## Annotating with terminator ...
## Annotating with gene.body ...
annotationFeatures(k562.rep1)
```

## [1] "gene.body" "promoter" "distal" "terminator"

We can now find interactions involving promoters using the annotated node.class for each anchor:

```
p.one = anchorOne(k562.rep1)$node.class == "promoter"
p.two = anchorTwo(k562.rep1)$node.class == "promoter"
k562.rep1[p.one|p.two]
```

```
## GenomicInteractions object with 2907 interactions and 2 metadata columns:
## Name: k562
   Description: PolII-8wg16 Chia-PET for K562
##
    Sum of interactions: 58468
##
##
    Annotated: yes
##
         Annotated with: promoter, gene.body, distal, terminator
##
    Interactions:
##
                           Anchor One
                                                         Anchor Two Counts
                                                                              I
                                              chr17:620668..626263
##
        [1]
                chr17:616579..621961 ---
                                                                         7
                                                                             T
##
        [2]
                chr17:632527..638035 ---
                                              chr17:636589..641349
                                                                         9
                                                                             T
##
        [3]
                chr17:634119..651606 ---
                                              chr17:642299..659172
                                                                        55
                                                                             ##
        [4]
                chr17:654892..657597 ---
                                              chr17:683191..687275
                                                                             T
                                                                         6
```

##	[5]	chr17:65	56002658841		chr17:679595682692	3	
##							
##	[2903]	chr18:777811	15177783476		chr18:7779296877795855	3	l
##	[2904]	chr18:777845	59077787797		chr18:7779214877795822	6	l
##	[2905]	chr18:777920	9377797983		chr18:7779745577803127	13	
##	[2906]	chr18:777933	36577797939		chr18:7786488977868321	8	
##	[2907]	chr18:778639	99277870413		chr18:7786829477877151	18	l
##		p.value	fdr				
##	[1]	2.06358e-32	8.50563e-30				
##	[2]	2.395e-36	1.1518e-33				
##	[3]	0	0				
##	[4]	4.86283e-24	1.27856e-21				
##	[5]	6.23098e-14	5.72161e-12				
##							
##	[2903]	8.97548e-14	8.09366e-12				
##	[2904]	4.16341e-26	1.24576e-23				
##	[2905]	0	0				
##	[2906]	6.61618e-30	2.41717e-27				
##	[2907]	0	0				
##							
##							
##	seqinfo	o: 25 sequenc	ces from an un	ispec	cified genome; no seqlengths		

This information can be used to categorise interactions into promoter-distal, promoter-terminator etc. A table of interaction types can be generated with categoriseInteractions:

categoriseInteractions(k562.rep1)

##		category	$\operatorname{count}$
##	1	gene.body-gene.body	795
##	2	gene.body-promoter	917
##	3	gene.body-distal	76
##	4	gene.body-terminator	164
##	5	promoter-promoter	1187
##	6	promoter-distal	519
##	7	promoter-terminator	284
##	8	distal-distal	396
##	9	distal-terminator	101
##	10	terminator-terminator	70

Alternatively, we can subset the object based on interaction type:

k562.rep1[isInteractionType(k562.rep1, "terminator", "gene.body")]

```
## GenomicInteractions object with 164 interactions and 2 metadata columns:
## Name: k562
## Description: PolII-8wg16 Chia-PET for K562
## Sum of interactions: 930
## Annotated: yes
## Annotated with: terminator, gene.body
## Interactions:
## Anchor One Anchor Two Counts |
```

## [1] chr17:1471460..1474306 --chr17:1476212..1479585 4 ## [2] chr17:1636657..1642967 12 chr17:1632603..1638741 --chr17:3845975..3849573 ---## [3] chr17:3908008..3910817 5 [4] ## chr17:4055645..4058706 --chr17:4063341..4068158 4 ## [5] chr17:4443889..4451615 --chr17:4446814..4454998 11 ## . . . . . . . . . . . . . . . . ## [160] chr18:32873043..32876330 --- chr18:32911004..32914907 4 ## [161] chr18:33566139..33569380 --- chr18:33571111..33574360 3 [162] chr18:33689982..33692774 --- chr18:33695495..33699363 ## 4 I [163] chr18:42638745..42641928 --- chr18:42647110..42649707 3 ## ## [164] chr18:77914125..77916781 --- chr18:77919050..77922761 3 ## p.value fdr 1.1438e-18 1.92712e-16 ## [1] ## [2] 0 4.06377e-44 ## [3] 1.46527e-19 2.69952e-17 ## [4] 4.96495e-19 8.66918e-17 ## [5] 7.2293e-42 4.17953e-39 ## . . . . . . . . . ## [160] 1.37802e-15 1.62547e-13 ## [161] 5.37957e-15 5.74333e-13 ## [162] 1.35703e-19 2.5096e-17 ## [163] 1.03659e-15 1.24713e-13 ## [164] 2.9097e-15 3.24878e-13 ## ## \_\_\_\_\_ ## seqinfo: 25 sequences from an unspecified genome; no seqlengths

The 3 most common node.class values have short functions defined for convenience (see ?is.pp for a complete list):

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```
k562.rep1[is.pp(k562.rep1)] # promoter-promoter interactions
k562.rep1[is.dd(k562.rep1)] # distal-distal interactions
k562.rep1[is.pt(k562.rep1)] # promoter-terminator interactions
```

Summary plots of interactions classes can easily be produced to get an overall feel for the data:

```
plotInteractionAnnotations(k562.rep1, other=5)
```

# **Interaction Classes**



viewpoints will only take those interactions with a certain node.class:

plotInteractionAnnotations(k562.rep1, other=5, viewpoints="promoter")



# **Interaction Classes**

These are also combined in the function plotSummaryStats.

## Feature Summaries

The summariseByFeatures allows us to look in more detail at interactions involving a specific set of loci. In this example we use all RefSeq promoters, which we already have loaded in a GRangesList object.

It is however possible to use any dataset which can be represented as a named **GRanges** object, for example transcription-factor ChIP data, predicted cis-regulatory sites or certain categories of genes.

The categories are generated automatically from the annotated node.class values in the object.

colnames(k562.rep1.promoter.annotation)

##	[1]	"Promoter.id"
##	[2]	"numberOfPromoterInteractions"
##	[3]	"numberOfPromoterUniqueInteractions"
##	[4]	"numberOfPromoterInterChromosomalInteractions"
##	[5]	"numberOfPromoterUniqueInterChromosomalInteractions"
##	[6]	"numberOfPromoterGene.bodyInteractions"
##	[7]	"numberOfPromoterPromoterInteractions"
##	[8]	"numberOfPromoterDistalInteractions"
##	[9]	"numberOfPromoterTerminatorInteractions"
##	[10]	"numberOfUniquePromoterGene.bodyInteractions"
##	[11]	"numberOfUniquePromoterPromoterInteractions"
##	[12]	"numberOfUniquePromoterDistalInteractions"
##	[13]	"numberOfUniquePromoterTerminatorInteractions"
##	[14]	"PromoterDistanceMedian"
##	[15]	"PromoterDistanceMean"
##	[16]	"PromoterDistanceMinimum"
##	[17]	"PromoterDistanceMaximum"
##	[18]	"PromoterDistanceWeightedMedian"
##	[19]	"numberOfSelfPromoterGene.bodyInteractions"
##	[20]	"numberOfSelfPromoterPromoterInteractions"
##	[21]	"numberOfSelfPromoterTerminatorInteractions"
##	[22]	"numberOfSelfUniquePromoterGene.bodyInteractions"
##	[23]	"numberOfSelfUniquePromoterPromoterInteractions"
##	[24]	"numberOfSelfUniquePromoterTerminatorInteractions"

This allows us to very quickly generate summaries of the data and provides a quick method to isolate genes of interest. In this case we produce lists of RefSeq IDs, which can easily be converted to EntrezIDs or gene symbols through existing BioConductor packages (in this case org.Hs.eg.db provides bimaps between common human genome annotations).

Which promoters have the strongest Promoter-Promoter interactions based on PET-counts?

Which promoters are contacting the largest number of distal elements?

## [1] "10140" "400604" "7050" "100130581" "100616277" ## [6] "26118" "100874261" "101927666" "140735" "5366"

What percentage of promoters are in contact with transcription termination sites?

```
total = sum(k562.rep1.promoter.annotation$numberOfPromoterTerminatorInteractions > 0)
sprintf("%.2f%% of promoters have P-T interactions", 100*total/nrow(k562.rep1.promoter.annotation))
```

## [1] "16.43% of promoters have P-T interactions"

### References

- 1. Li, Guoliang, et al. "Software ChIA-PET tool for comprehensive chromatin interaction analysis with paired-end tag sequencing." Genome Biol 11 (2010): R22.
- 2. Li, Guoliang, et al. "Extensive promoter-centered chromatin interactions provide a topological basis for transcription regulation." Cell 148.1 (2012): 84-98