The ChIPpeakAnno user's guide

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September 7, 2011

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1 Introduction

Chromatin immunoprecipitation (ChIP) followed by high-throughput tag sequencing (ChIP-seq) and ChIP followed by genome tiling array analysis (ChIP-chip) become more and more prevalent high throughput technologies for identifying the binding sites of DNA-binding proteins in a genome-wide bases. A number of algorithms have been published to facilitate the identification of the binding sites of the DNA-binding proteins of interest. The identified binding sites in the list of peaks are usually converted to BED or WIG file format to be loaded to UCSC genome browser as custom tracks for investigators to view the proximity to various genomic features such as genes, exons and conserved elements. However, clicking through the genome browser could be a daunting task for the biologist if the number of peaks gets large or the peaks spread widely across the genome. Here we have developed a

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Bioconducor package called ChIPpeakAnno to facilitate the batch annotation of the peaks identified from either ChIP-seq or ChIP-chip experiments. We have implemented functionality to find the nearest gene, exon, miRNA or custom features supplied by users such as most conserved elements and other transcription factor binding sites leveraging IRanges. Since the genome annotation gets updated from time to time, we have leveraged the biomaRt package from Bioconductor to retrieve the annotation data on the fly if the annotation of interest is available via the biomaRt package. The users also have the flexibility to pass their own annotation data as RangedData or pass in annotation data from GenomicFeatures. We have also leveraged BSgenome and biomaRt package on implementing functions to retrieve the sequences around the peak identified for peak validation. To understand whether the identified peaks are enriched around genes with certain GO terms, we have implemented GO enrichment test in ChIPpeakAnno package leveraging the hypergeometric test phyper in stats package and integrated with Gene Ontology (GO) annotation from GO.db package and multiplicity adjustment functions from multtest package.

2 Examples of using ChIPpeakAnno

2.1 Task 1: Find the nearest feature such as gene and the distance to the feature such as the transcription start site (TSS) of the nearest gene

We have a list of peaks identified from ChIP-seq or ChIP-chip experiments and we would like to retrieve the nearest gene and distance to the corresponding gene transcription start site. We have retrieved all the genomic locations of the genes for human genome as TSS.human.NCBI36 data package for repeated use with function getAnnotation, now we just pass the annotation to the annotatePeakInBatch function.

```
> library(ChIPpeakAnno)
> data(myPeakList)
> data(TSS.human.NCBI36)
> annotatedPeak = annotatePeakInBatch(myPeakList[1:6, ], AnnotationData = TSS.human.NCBI36)
> as.data.frame(annotatedPeak)
                                                                   peak strand
 space
         start
                    end width
                                                     names
     1 703885 703985 101 1_12_703729 ENSG00000197049 1_12_703729
      1 559774 559874
                         101 1_41_559455 ENSG00000212678 1_41_559455
2
3
        556660 556760
                         101 1_93_556427 ENSG00000212875 1_93_556427
                         101 1_11_1041174 ENSG00000131591 1_11_1041174
     1 1041646 1041746
                         101 1_14_1269014 ENSG00000107404 1_14_1269014
5
     1 1270239 1270339
      1 926058 926158 101 1_20_925025 ENSG00000188290 1_20_925025
         feature start_position end_position insideFeature distancetoFeature
1 ENSG00000197049
                         711183
                                       712376
                                                   upstream
                                                                        -7298
2 ENSG00000212678
                          559619
                                       560165
                                                    inside
                                                                          155
3 ENSG00000212875
                         556317
                                      557859
                                                     inside
                                                                          343
4 ENSG00000131591
                         1007061
                                      1041341
                                                                         -305
                                                   upstream
5 ENSG00000107404
                         1260522
                                      1274623
                                                    inside
                                                                         4384
6 ENSG00000188290
                          924208
                                      925333
                                                   upstream
                                                                         -725
 \verb|shortestDistance| from Overlapping Or Nearest|
              7198
                              NearestStart
              155
                               NearestStart
```

3	343	NearestStart
4	305	NearestStart
5	4284	NearestStart
6	725	NearestStart

To annotate the peaks with other genomic feature, you will need to call function getAnnotation with featureType, e.g., "Exon" for finding the nearest exon, and "miRNA" for finding the nearest miRNA, "5utr" or '3utr"for finding the overlapping 5 prime UTR or 3 prime UTR. Please refer to getAnnotation function for more details.

We have presented the examples using human genome as annotation source. To annotate your data with other species, you will need to pass to the function getAnnotation the appropriate dataset for example, drerio_gene_ensembl for zebrafish genome, mmusculus_gene_ensembl for mouse genome and rnorvegicus_gene_ensembl for rat genome. For a list of available biomart and dataset, please refer to the biomaRt package documentation (Durinck S. et al., 2005). For fast access, in addition to TSS.human.NCBI36, TSS.mouse.NCBIM37, TSS.rat.RGSC3.4 and TSS.zebrafish.Zv8 are included as annotation data packages.

You could also pass your own annotation data into the function annotatePeakInBatch. For example, if you have a list of transcription factor biding sites from literature and are interested in obtaining the nearest binding site of the transcription factor and distance to it for the list of peaks.

```
> myPeak1 = RangedData(IRanges(start = c(967654, 2010897, 2496704,
      3075869, 3123260, 3857501, 201089, 1543200, 1557200, 1563000,
      1569800, 167889600), end = c(967754, 2010997, 2496804, 3075969,
      3123360, 3857601, 201089, 1555199, 1560599, 1565199, 1573799,
      167893599), names = c("Site1", "Site2", "Site3", "Site4",
      "Site5", "Site6", "Site7", "Site8", "Site9", "Site10", "Site11",
      "Site12")), space = c("1", "2", "3", "4", "5", "6", "2",
      "6", "6", "6", "6", "5"))
> TFbindingSites = RangedData(IRanges(start = c(967659, 2010898,
      2496700, 3075866, 3123260, 3857500, 96765, 201089, 249670,
      307586, 312326, 385750, 1549800, 1554400, 1565000, 1569400,
      167888600), end = c(967869, 2011108, 2496920, 3076166, 3123470,
      3857780, 96985, 201299, 249890, 307796, 312586, 385960, 1550599,
      1560799, 1565399, 1571199, 167888999), names = c("t1", "t2", "t2")
      "t3", "t4", "t5", "t6", "t7", "t8", "t9", "t10", "t11", "t12"
      "t13", "t14", "t15", "t16", "t17")), space = c("1", "2", "3", "4", "5", "6", "1", "2", "3", "4", "5", "6", "6", "6",
          "6", "5"), strand = c(1, 1, 1, 1, 1, 1, -1, -1, -1,
      -1, -1, -1, 1, 1, 1, 1)
> annotatedPeak2 = annotatePeakInBatch(myPeak1, AnnotationData = TFbindingSites)
> pie(table(as.data.frame(annotatedPeak2)$insideFeature))
> as.data.frame(annotatedPeak2)
                         end width
   space
             start
                                        names
                                                 peak strand feature
                      967754 101
            967654
                                     Site1 t1 Site1
      1
2
           2010897
                     2010997
                               101
                                      Site2 t2
                                                Site2
                                     Site7 t8 Site7
3
           201089
                     201089
                                1
                                                                   t8
4
           2496704
                     2496804
                              101
                                     Site3 t3 Site3
                                                                   t3
5
          3075869
                     3075969
                              101
                                     Site4 t4 Site4
                                                                  t4
       5 167889600 167893599 4000 Site12 t17 Site12
                                                                  t17
6
          3123260
                     3123360
                               101
       5
                                      Site5 t5 Site5
                              2200 Site10 t15 Site10
8
       6
          1563000
                     1565199
                                                                  t15
          1569800
                     1573799
                              4000 Site11 t16 Site11
                                                                  t16
       6
10
      6
           3857501
                     3857601
                              101
                                    Site6 t6 Site6
                                                                   t6
          1543200
                     1555199 12000 Site8 t13 Site8
                                                                  t13
```

12	6 1557200	1560599	3400 Site9 t14	Site9 +	t14
	start_position e	nd_position	insideFeature	distancetoFeature	
1	967659	967869	overlapStart	-5	
2	2010898	2011108	overlapStart	-1	
3	201089	201299	inside	210	
4	2496700	2496920	inside	4	
5	3075866	3076166	inside	3	
6	167888600	167888999	downstream	1000	
7	3123260	3123470	inside	0	
8	1565000	1565399	overlapStart	-2000	
9	1569400	1571199	overlapEnd	400	
10	3857500	3857780	inside	1	
11	1549800	1550599	includeFeature	-6600	
12	1554400	1560799	inside	2800	
	shortestDistance	fromOverlap	ppingOrNearest		
1	5		NearestStart		
2	1		NearestStart		
3	0		NearestStart		
4	4		NearestStart		
5	3		NearestStart		
6	601		NearestStart		
7	0		NearestStart		
8	199		NearestStart		
9	400		NearestStart		
10	1		NearestStart		
11	4600		NearestStart		
12	200		NearestStart		

Both BED format and GFF format are common file format that provides a flexible way to define the peaks and annotations as the data lines. Therefore, conversion functions RfunctionBED2RangedData and RfunctionGFF2RangedData were implemented for converting these data format to RangedData before calling annotatePeakInBatch

Once you annotated the peak list, you can plot the distance to nearest feature such as TSS.

2.2 Task 2: Obtain overlapping peaks for potential transcription factor complex and determine the significance of the overlapping and generate Venn Diagram

Here is an example of obtaining overlapping peaks with maximum gap 1kb for two peak ranges.

```
+ "t11", "t12", "t13", "t14", "t15", "t16", "t17")), space = c("1", 
+ "2", "3", "4", "5", "6", "1", "2", "3", "4", "5", "6", "6", 
+ "6", "6", "6", "5"), strand = c(1, 1, 1, 1, 1, 1, -1, -1, 
+ -1, -1, -1, -1, 1, 1, 1, 1))
> t1 = findOverlappingPeaks(peaks1, peaks2, maxgap = 1000, multiple = F, 
+ NameOfPeaks1 = "TF1", NameOfPeaks2 = "TF2")
```

Here is a list of overlapping peaks with maximum gap 1kb and a pie graph describing the distribution of relative position of peaks1 to peaks2 for overlapping peaks.

```
> overlappingPeaks = t1$OverlappingPeaks
> overlappingPeaks
     TF1 chr TF2 TF2_start
                        TF2_end strand TF1_start
                                                TF1_end strand1
   Site1 1 t1
                 967659
                          967869 + 967654
                                                 967754
1
   Site2
         2 t2
                2010898
                         2011108
                                       2010897
                                                 2010997
10 Site7 2 t8
                201089
                         201299 -
                                        201089
                                                201089
   Site3 3 t3 2496700 2496920 + 2496704
                                                 2496804
  Site4 4 t4 3075866 3076166 + 3075869
                                                3075969
  Site12
         5 t17 167888600 167888999
                                    + 167889600 167893599
  Site5 5 t5 3123260 3123470
                                   + 3123260
8
                                                3123360
2 Site10 6 t15 1565000
                        1565399
                                    + 1563000
                                                1565199
3 Site11 6 t16 1569400
                        1571199
                                     + 1569800
                                                1573799
                        3857780
                                       3857501
  Site6 6 t6 3857500
                                                 3857601
9
         6 t13
                1549800
                         1550599
                                        1543200
                                                 1555199
12 Site9 6 t14 1554400 1560799
                                    + 1557200
                                                1560599
  overlapFeature shortestDistance
   overlapStart
5
   overlapStart
                            1
10
                            0
         inside
6
         inside
                            4
7
        inside
4
     downstream
                          601
8
                           0
       inside
2
    overlapStart
                          199
3
    overlapEnd
                          400
        inside
                          4600
11 includeFeature
```

200

> pie(table(overlappingPeaks\$overlapFeature))

Here is the merged overlapping peaks, which can be used to obtain overlapping peaks with another TF binding sites from a protein complex.

> as.data.frame(t1\$MergedPeaks)

inside

12

	space	start	end	${\tt width}$	names
1	1	967654	967869	216	TF1-Site1-TF2-t1
2	2	2010897	2011108	212	TF1-Site2-TF2-t2
3	2	201089	201299	211	TF1-Site7-TF2-t8
4	3	2496700	2496920	221	TF1-Site3-TF2-t3
5	4	3075866	3076166	301	TF1-Site4-TF2-t4
6	5	167888600	167893599	5000	TF1-Site12-TF2-t17
7	5	3123260	3123470	211	TF1-Site5-TF2-t5
8	6	1563000	1565399	2400	TF1-Site10-TF2-t15
9	6	1569400	1573799	4400	TF1-Site11-TF2-t16
10	6	3857500	3857780	281	TF1-Site6-TF2-t6
11	6	1543200	1555199	12000	TF1-Site8-TF2-t13
12	6	1554400	1560799	6400	TF1-Site9-TF2-t14

Here is the peaks in peaks1 that overlaps with peaks in peaks2

> as.data.frame(t1\$Peaks1withOverlaps)

```
start
                      end width names strand
  space
1
      1
          967654
                    967754 101 Site1
2
      2
         2010897
                   2010997
                            101 Site2
3
          201089
                   201089
                            1 Site7
4
      3
         2496704
                   2496804 101 Site3
                   3075969 101 Site4
         3075869
6
      5 167889600 167893599 4000 Site12
7
      5
         3123260
                   3123360
                            101 Site5
8
      6
         1563000
                   1565199 2200 Site10
         1569800
9
                   1573799
                           4000 Site11
      6
         3857501
                   3857601
                            101 Site6
      6
         1543200
                   1555199 12000 Site8
11
12
      6
          1557200
                   1560599 3400
                                 Site9
```

Here is the peaks in peaks2 that overlap with peaks in peaks1

> as.data.frame(t1\$Peaks2with0verlaps)

```
space
            start
                       end width names strand
           967659
                    967869
                             211
                                   t1
1
      1
2
      2
          2010898
                   2011108
                             211
                                    t2
                                   t8
          201089
                   201299
                             211
3
      2
      3 2496700
                   2496920
                   3076166
                             301
                                   t.4
5
      4 3075866
6
      5 167888600 167888999
                             400
                                  t17
7
      5 3123260
                   3123470
                             211
                                   t5
8
      6
        1565000
                   1565399 400
                                   t15
        1569400
                   1571199 1800
                                   t16
10
      6
          3857500
                   3857780
                             281
                                   t6
11
      6
          1549800
                   1550599
                            800
                                   t13
12
          1554400
                   1560799 6400
                                   t14
```

The findOVerlappingPeaks function can be repeatedly called to obtain for example, the peaks in peaks1 that overlap with peaks in both peaks2 and peaks3.

Venn Diagram can be generated by the following function call with p-value that indicates whether the extent of overlapping is significant.

2.3 Task 3: Obtain sequences surrounding the peaks for PCR validation or motif discovery

Here is an example of obtaining sequences surrounding the peak intervals including 20 bp upstream and downstream sequence.

```
> peaks = RangedData(IRanges(start = c(100, 500), end = c(300,
+ 600), names = c("peak1", "peak2")), space = c("NC_008253",
+ "NC_010468"))
> library(BSgenome.Ecoli.NCBI.20080805)
> peaksWithSequences = getAllPeakSequence(peaks, upstream = 20,
+ downstream = 20, genome = Ecoli)
```

You can easily convert the obtained sequences into fasta format for motif discovery by calling the function write2FASTA.

2.4 Task 3: Obtain enriched gene ontology (GO) terms near the peaks

Once you have obtained the annotated peak data from the example above, you can also use the function **getEnrichedGO** to obtain a list of enriched gene ontology (GO) terms using hypergeometric test.

```
library(org.Hs.eg.db) 
 enrichedGO = getEnrichedGO (annotatedPeak, orgAnn = "org.Hs.eg.db", maxP = 0.01, multiAdj = TRUE, minGOterm = 10, multiAdjMethod = "BH")
```

Please note that org.Hs.eg.db is the GO gene mapping for Human, for other organisms, please refer to http://www.bioconductor.org/packages/release/data/annotation/ for additional org.xx.eg.db packages.

> data(enrichedGO)

Here is a list of enriched GO biological process for myPeakList dataset.

```
> enrichedGO$bp[1:6, ]
       go.id
1 GO:0000187
2 GO:0002573
3 GO:0002702
4 GO:0002761
5 GO:0002763
6 GD:0006213
                                                                         go.term
1
                                                   activation of MAPK activity
2
                                             myeloid leukocyte differentiation
3 positive regulation of production of molecular mediator of immune response
4
                              regulation of myeloid leukocyte differentiation
                     positive regulation of myeloid leukocyte differentiation
5
                                       pyrimidine nucleoside metabolic process
6
1
2
3
4
5
6 The chemical reactions and pathways involving any pyrimidine nucleoside, one of a fami
  Ontology count.InDataset count.InGenome
                                                 pvalue totaltermInDataset
        BP
                         17
                                         65 0.001673400
1
                                                                      85892
                                         81 0.004192510
2
        BP
                         19
                                                                      85892
3
        BP
                          4
                                         10 0.005921074
                                                                      85892
                                         50 0.004712934
4
        BP
                         13
                                                                      85892
5
        BP
                          8
                                         22 0.001277580
                                                                      85892
        ΒP
                                         10 0.005921074
                                                                      85892
6
  totaltermInGenome
             644151
1
2
             644151
3
             644151
4
             644151
5
             644151
6
             644151
```

Here is a list of enriched GO molecular functions for myPeakList dataset.

> enrichedGO\$mf[1:6,]

	go.id				go.t	erm
1	GD:0003702		RNA polymer	case II trans	cription factor activ	ity
2	GO:0003705 RNA	polymerase II	transcripti	ion factor act	tivity, enhancer bind	ing
3	GO:0004112		cyclic-r	nucleotide pho	osphodiesterase activ	ity
4	GO:0004114	3',	5'-cyclic-r	nucleotide pho	osphodiesterase activ	ity
5	GO:0004659			pre	enyltransferase activ	ity
6	GD:0004896			су	tokine receptor activ	rity
			_			_
1					itiate or regulate RN	- •
		•	-	v	transcription by bind	•
3		•			de cyclic phosphate +	
4		•			'-cyclic phosphate +	
5		alysis of the t			up from one compound	
6				•	h a cytokine to initi	ate a change
	••	.InDataset cour			totaltermInDataset	
1		39		0.0065818928		
2		11		0.0001003699	29657	
3		9		0.0007622170	29657	
4		9		0.0005282939	29657	
5		9		0.0002346785	29657	
6	MF	16	66	0.0027160003	29657	
	totaltermInGen					
1						
2						
3						
4						
5						
6	235	991				

Heres is a list of enriched GO cellular components for myPeakList dataset.

> enrichedGO\$cc

	go.id	go.term
1	GO:0005811	lipid particle
2	GO:0005942	phosphoinositide 3-kinase complex
3	GO:0016363	nuclear matrix
4	GD:0034399	nuclear periphery

Any particle of coalesced lipids in the cytopl

2 A complex containing a heterodimer of a catalytic subunit and a regulatory (adaptor) s

3					The dense fibrillar network
4					The portion of the nuclea
	Ontology	<pre>count.InDataset</pre>	$\verb"count.InGenome"$	pvalue	totaltermInDataset
1	CC	5	15	0.006685158	45317
2	CC	4	11	0.007074546	45317
3	CC	12	49	0.005607016	45317
4	CC	12	52	0.009516449	45317
	totalterm	InGenome			
1		365523			
2		365523			
3		365523			
4		365523			

3 References

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4 Session Info

> sessionInfo()

R version 2.13.1 (2011-07-08)

Platform: x86_64-unknown-linux-gnu (64-bit)

locale:

- [1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8 LC_COLLATE=C
- [5] LC_MONETARY=C LC_MESSAGES=en_US.UTF-8
- [7] LC_PAPER=en_US.UTF-8 LC_NAME=C
- [9] LC_ADDRESS=C LC_TELEPHONE=C
- [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

- [1] ChIPpeakAnno_1.9.0 limma_3.8.3 [3] org.Hs.eg.db_2.5.0 GO.db_2.5.0 [5] RSQLite_0.9-4 DBI_0.2-5
- [7] AnnotationDbi_1.14.1 BSgenome.Ecoli.NCBI.20080805_1.3.17
- [9] BSgenome_1.20.0 GenomicRanges_1.4.8
- [15] biomaRt_2.8.1

loaded via a namespace (and not attached):

- [1] MASS_7.3-14 RCurl_1.6-10 XML_3.4-2 splines_2.13.1
- [5] survival_2.36-9 tools_2.13.1