geneXtendeR

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1 Introduction

This vignette describes geneXtendeR, an R/Bioconductor package for optimized functional annotation of ChIP-seq data. This software is designed for optimized annotation of genomic features (primarily peaks called from a ChIP-seq experiment, but any coverage island regions would work) with the nearest gene. "Extending" refers to performing gene-feature overlaps after adding to the gene-span a user-specified region upstream of the start of the gene model and a fixed (500 bp) region downstream of the gene, resulting in assigning to a gene the features that do not physically overlap with it but are sufficiently close. This facilitates the process of deciphering which differentially enriched peaks are dysregulating which specific genes which, in turn, aids experimental follow-up and validation in designing primers for a set of prospective genes during qPCR (Barbier et al. 2016).

1.1 Brief Background

With an abundance of Bioconductor software currently available for peak annotation to nearby features (e.g., ChIPpeakAnno (Zhu et al. 2010)) as well as the existence of various command line tools (e.g., BEDTools *closest* function (Quinlan and Hall, 2010), HOMER (Heinz et al. 2010)), what makes geneXtendeR different? Let's look at a concrete example presented as a case-study:

1.2 Case-study

1.2.1 R/Bioconductor package installation

Here we use geneXtendeR to analyze ChIP-seq data from a cardiac ischemia study published in *Journal of the American Heart Association* (Gidlöf et al. 2016). To follow along with the analysis steps of this workflow, please download

the latest version of geneXtendeR directly from Github, since Bioconductor is on a bi-annual release cycle (and thus may not be fully up-to-date with the latest package features). To download the latest version of the geneXtendeR package:

```
> install.packages("devtools")
> library(devtools)
> install_github("Bohdan-Khomtchouk/geneXtendeR")
> library(geneXtendeR)
```

Otherwise, to install directly from Bioconductor (may not be fully up-to-date), do:

```
> ## try http:// if https:// URLs are not supported
> source("https://bioconductor.org/biocLite.R")
> biocLite("geneXtendeR")
> library(geneXtendeR)
```

1.2.2 Quick setup

Per Gidlöf et al. 2016, please load in a mouse GTF file:

Note: Please make sure the above command is pre-formatted to fit on one line (as opposed to three separate lines like displayed above for page margin purposes).

Next, load in the ChIP-seq peak coordinates produced by the bioinformatics pipeline used by Gidlöf et al. 2016 (comes pre-bundled with the geneXtendeR package for convenience):

The structure of this peak coordinates file is explained in Section 2.1. For reference, these genomic coordinates were called using the SICER peak calling algorithm (Zang et al. 2009), and can be recreated by the user from the original sequencing files (deposited in the Gene Expression Omnibus), as specified in the instructions given in the Gidlöf et al. 2016 publication.

1.2.3 Gene-centric functional annotation

1.2.3.1 Mapping peaks to genes with the gene_annotate() function Type in the following command to annotate the peaks file produced in the previous section with a GTF file (the R object mouse above) whose genes have been extended 2000 bp upstream of their first exon (and, by default, 500 bp downstream of their last exon):

>	head(gene_a	annotate(mou	use, 2000),			
	Chromosome	Gene-Start	Gene-End	Gene-I	Gene-N	ame
1	8	119908841	124346222	ENSMUSG0000009232	9 Gm20	388
2	7	130161951	133125350	ENSMUSG0000003084) Fg	fr2
3	4	150916822	151863876	ENSMUSG0000001459	2 Cam	ta1
4	18	38599534	39376784	ENSMUSG0000003645	2 Arhga	p26
5	1	73909731	74126449	ENSMUSG0000005532	2 T	ns1
6	17	86165785	86658419	ENSMUSG0000004503	3 Pr	kce
	Peaks-on-G	ene-Body Mea	an-Distance	e-of-Gene-to-Neare	st-Peaks	sd
1		781			0.00000	0.0000
2		165			0.00000	0.0000
3		77			0.00000	0.0000
4		62			0.00000	0.0000
5		57		9	95.39655	726.5185
6		57		(55.29310	497.2575
	Number-of-	Peaks-Associ	iated-with	-Gene		
1				781		
2				165		
3				77		
4		62				
5				58		
6				58		

Clearly, the peaks file has now been functionally annotated with the content of the mouse genome (mm10 build). Specifically, each individual row of peak coordinates in the input file (chromosome, start position of peak, end position of peak) has been annotated with relevant gene information and collapsed into a tabular summary format. This output labels each individual gene and matches

it with the number of peaks that overlap its gene-body (2000 bp upstream and 500 bp downstream in the example above) and that are "first away" from its gene-body (i.e., closest/nearest but not overlapping). Distance is calculated between the 5-prime end of a gene and 3-prime end of a peak (or 3-prime end of a gene and 5-prime end of a peak, whichever is smallest). The table is sorted by number of peaks on gene body (i.e., Peaks-on-Gene-Body, which is the number of peaks that directly overlap the gene body) and include extra information such as mean and standard deviation (sd) for extra validation. Typically, a user would be looking for genes that have a high number of Peaks-on-Gene-Body to followup on for experimental validation. Genes that have peaks that reside close (but not overlapping) to the chosen gene-body (i.e., low mean) and that are clustered together spatially (i.e., low standard deviation) may also be good targets for follow-up analysis. Number-of-Peaks-Associated-with-Gene represents the number of peaks that directly overlap the gene body + the number of peaks that are directly adjacent to the gene body (first nearest/closest). Therefore, it should be noted that mean = 0 (i.e., Mean-Distance-of-Gene-to-Nearest-Peaks = 0) denotes cases where all peaks are overlapping a given gene body (with no nearest/closest peaks).

The table above shows that the top 3 genes (in terms of total number of peaks overlapping their gene body) are Gm20388, Fgfr2, and Camta1 — which have 781, 165, and 77 peaks (respectively). Although little is currently known in the literature about Gm20388 (since it is a predicted gene), the gene Fgfr2 plays a well-known role in cardiac ischemia (House et al. 2016). In addition, Gidlöf et al. 2016 reports that the gene Camta1 is significantly downregulated in ischemic heart tissue enriched in H3K9me2 (Table S2, Gidlöf et al. 2016), as quantified by p-value and fold change information acquired from microarray. Therefore, geneXtendeR has successfully shown at least 2 out of 3 top genes to play a role in ischemia.

1.2.3.2 Mapping genes to peaks with the gene_lookup() function Similarly, the gene_lookup() function looks up all peaks surrounding a specific gene or list of genes across all chromosomes and reports these peaks. This method is extremely useful when paired with gene_annotate() to check genes that may be used in a follow-up. Here, we examine the mTOR gene, which was also experimentally validated in Gidlöf et al. 2016:

```
4:
                148457000 148460199
                                                        148446611 148558183 Mtor
5:
             4
                148457400 148460399
                                                        148446611 148558183 Mtor
 6:
             4
               148458000 148461399
                                                     0
                                                        148446611 148558183 Mtor
7:
             4
                148462200 148463999
                                                        148446611 148558183 Mtor
                148469200 148474199
                                                        148446611 148558183 Mtor
8:
             4
9:
                148490200 148494799
                                                        148446611 148558183 Mtor
             4
10:
             4
                148491000 148493199
                                                     0
                                                        148446611 148558183 Mtor
                148507600 148510799
                                                        148446611 148558183 Mtor
11:
12:
             4
                148511400 148513199
                                                     0
                                                        148446611 148558183 Mtor
13:
             4
                148522200 148524599
                                                        148446611 148558183 Mtor
14:
             4
                148524800 148529199
                                                        148446611 148558183 Mtor
                148525600 148528799
15:
                                                        148446611 148558183 Mtor
             4
                148536000 148539799
                                                        148446611 148558183 Mtor
16:
             4
                148536000 148537999
17:
                                                     0
                                                        148446611 148558183 Mtor
             4
18:
                148537200 148542399
                                                        148446611 148558183 Mtor
             4
19:
             4
                148553200 148554399
                                                        148446611 148558183 Mtor
20:
             4
                148442000 148444399
                                                  2212
                                                        148446611 148558183 Mtor
21:
                148563000 148564799
                                                 4817
                                                        148446611 148558183 Mtor
             4
22:
                148569200 148571599
                                                11017
                                                        148446611 148558183 Mtor
    Chromosome Peak-Start Peak-End Distance-to-Gene Gene-Start Gene-End Gene
```

This output shows the sheer quantity of peaks that overlap the mTOR gene body (19 peaks!). It is no surprise that, with this many peaks directly on top of the mTOR gene, experimental validation was indeed successful in this case (Gidlöf et al. 2016). Using geneXtendeR can suggest such opportunities for wet-lab follow-up, especially when combined with biological knowledge/domain expertise. For instance, it is known that mTOR is involved in the regulation of autophagy, and as the cardioprotective effect of ischemic preconditioning is strongly linked with autophagy, mTOR was an interesting gene to follow up on in this study. The hypothesis was that ischemic preconditioning (IPC) leads to enrichment of H3K9me2 throughout the mTOR gene, transcriptional repression, induction of autophagy and ultimately, cardioprotection. This hypothesis was successfully validated. For instance, it was confirmed that mTOR is indeed downregulated in IPC-hearts compared with qPCR (Figure 3a from Gidlöf et al. 2016). Therefore, knowing these genomic peak coordinates facilitated the design of PCR primers. Likewise, Figures 4-6 validated the other points of the hypothesis.

In gene_lookup(organism, gene_name, n, extension), n represents the number of nearest (and overlapping) peaks to a given gene. We see that in the case of mTOR there are quite a number of nearest and overlapping peaks to the gene, where the gene_lookup() function displays their location as well as their dis-

tance from the gene. Thus, this function is motivated by the need of biologists to accurately design primers for specific genomic loci in order to experimentally validate the existence (realness) of a peak.

1.2.3.3 N-dimensional annotation with the annotate_n() function geneX tendeR also provides a function that combines both gene_lookup() and gene_annotate() called: annotate_n(). Instead of simply annotating a peak to a single closest gene (and reporting any overlapping peaks on gene bodies), this function annotates each peak to the closest, the second-closest, ..., to the nth-closest genes to provide the user an expanded picture of the gene neighborhood around each individual peak. When called, this function looks like:

> /	head(annotate_n(mous	se	, 2000, n =	3), 9)		
	Peak-Num Chromosom	e I	Peak-Start	Peak-End	Gene-Start	Gene-End
1:	1	1	4586400	4588199	4582629	4588252
2:	1	1	4586400	4588199	4608471	4611906
3:	1	1	4586400	4588199	4534337	4537286
4:	2	1	4769000	4770999	4769131	4772699
5:	2	1	4769000	4770999	4772706	4787739
6:	2	1	4769000	4770999	4777563	4781212
7:	3	1	5071800	5073199	4909076	5072285
8:	3	1	5071800	5073199	4938076	4942710
9:	3	1	5071800	5073199	4926528	4929299
	Gene-ID	G	ene-Name ra	nk Minimu	um-Distance	-to-Gene
1:	ENSMUSG00000104328		Gm37323	1		0
2:	ENSMUSG00000102735		Gm7369	2		20272
3:	ENSMUSG00000103003		Gm38076	3		49114
4:	ENSMUSG00000103922		Gm6123	1		0
5:	ENSMUSG00000033845		Mrpl15	2		1707
6:	ENSMUSG00000102275		Gm37144	3		6564
7:	ENSMUSG00000002459		Rgs20	1		0
8:	ENSMUSG00000102653		Gm37079	2		129090
9:	ENSMUSG00000091305		Gm17100	3		142501

Since n=3 in the example above, each peak is annotated thrice — once for closest gene, once for second-closest gene, and once for third-closest gene. This function is the most versatile (and compute-intensive) of the annotation functions provided and is designed for the purpose of providing peak-to-gene associations and follow-up information that goes beyond just a simple "closest/nearest" genomic distance criterion. Future work in this direction could also address three-dimensional genome interactions (when coupled with methods like Hi-C), and we encourage the reader to explore this integrative frontier

further. When moving away from the traditional "first closest gene to a peak" approach, this method opens up many more possibilities as to which peaks may play a role in biologically influencing which genes. It increases the scope of the individual peaks to reduce the chance that a peak that influences any particular gene is missed or misattributed to the wrong gene. It also informs follow-up wet-lab strategy, for example, in the table above, rows 1-3 clearly suggest that the peak on chromosome 1, start position 4586400, and end position 4588199 overlaps gene Gm37323 but is too far from Gm7369 to be biologically relevant (20272 base pairs away). In contrast, rows 4-6 show that the peak located on chromosome 1, start position 4769000, and end position 4770999 overlaps gene Gm37323 yet is only 1707 bp away from a known gene (Mrpl15). Given the choice, such proximity suggests that it would be wiser to design PCR primers specific for the second-closest gene (Mrpl15), given that Gm37323 is a predicted gene while Mrpl15 is known to be linked with hypertension (Ong et al. 2013) and therefore may play a putative role in ischemia as well.

1.3 Making functional annotation more robust

1.3.1 Peak variability

It is well-known that peak coordinates (peak start position, peak end position) exhibit a considerable degree of variance depending on the peak caller used (e.g., SICER (Zang et al. 2009), MACS2 (Zhang et al. 2008), etc.), both in terms of length distribution of peaks as well as the total number of peaks called, even when run at identical default parameter values (Koohy et al. 2014; Thomas et al. 2017). Tuning algorithm-specific parameters produces even greater variance amongst peak callers, thereby complicating the issue further. This variance becomes a factor when annotating peak lists genome-wide with their nearest genes as, depending on the peak caller, peaks can be either shifted in genomic position (towards 5' or 3' end) or be of different lengths. Thus, there is often considerable variability of peak overlap with cis-regulatory elements and proximal-promoter regions. As such, geneXtendeR represents a first step towards making functional annotation more robust and consistent, by allowing users to experiment with multiple different gene body dimensions. This allows the user to tailor the functional annotation of a ChIP-seq peak dataset according to the details of the peak coordinates (chromosome number, peak start position, peak end position).

1.3.2 Gene ontology optimization

The primary focus of geneXtendeR is to optimize the process of functional annotation of a ChIP-seq peak list whereby instead of just annotating peaks with their nearest genomic features (as statically defined by a given genome build's coordinates), geneXtendeR investigates how peaks may align to various userspecified gene extensions (e.g., 500 bp upstream extensions, 2000 bp upstream extensions, etc. for all genes in the genome). This shows where peaks localize across the genome with respect to their nearest gene, as well as what gene ontologies (BP, CC, and MF) are impacted at these various extension levels (e.g., assuming one is interested only in investigating peaks that directly overlap genes, i.e., distance = 0). This, in turn, informs the user what gene extensions ideally capture the GO terms involved in the biology of their experiment. For example, if a user's study is investigating the role of epigenetic enzymes in alcohol addiction and dependence, then functionally annotating a peak list using gene extensions that maximize the number of brain-related ontologies (for both BP, CC, and MF categories) makes sense. This will be explored more in depth in Section 2.5.

1.3.3 Summary

With regards to histone modification ChIP-seq analysis, geneXtendeR computes optimal gene extensions tailored to the broadness of the specific epigenetic mark (e.g., H3K9me1, H3K27me3), as determined by a user-supplied ChIP-seq peak input file. This will be explored in the next section. To accomplish this level of custom-tailored data analysis, geneXtendeR first optimally extends the boundaries of every gene in a genome by some genomic distance (in DNA base pairs) for the purpose of flexibly incorporating cis-regulatory elements, such as promoter regions, as well as downstream elements that are important to the function of the gene relative to an epigenetic histone modification ChIPseq dataset. This action effectively transforms genes into "gene-spheres", a new term that we coin to emphasize the 3D-nature of heterochromatin (and, more importantly, to subliminally remind our users that annotate_n may be their friend – as alluded to in Section 1.2.3.3). A gene-sphere is composed of cis-regulatory elements (e.g., proximal promoters $+/-\approx 3$ kb from TSS), distal regulatory elements (e.g., enhancers), transcription start/end sites (TSS/TES), exons, introns, and downstream elements of a gene. As such, geneXtendeR maximizes the signal-to-noise ratio of locating genes closest to and directly under peaks. By performing a computational expansion of this nature, ChIPseq reads that would initially not map strictly to a specific gene can now be optimally mapped to the regulatory regions of the gene, thereby implicating the gene as a potential candidate, and thereby making the ChIP-seq analysis more successful. Such an approach becomes particularly important when working with epigenetic histone modifications that have inherently broad peaks with a diffuse range of signal enrichment (e.g., H3K9me1, H3K27me3).

2 Sample workflow

2.1 Quick start

If you have not done so already (Section 1.2.1), please install the geneXtendeR package via Github or Bioconductor:

```
> ## try http:// if https:// URLs are not supported
> source("https://bioconductor.org/biocLite.R")
> biocLite("geneXtendeR")
> library(geneXtendeR)
```

This automatically loads the rtracklayer R package, which contains the read GFF() command used to retrieve GTF files of any model organism. As such, load in a GTF file into your R environment, e.g.:

```
> rat <- readGFF("ftp://ftp.ensembl.org/pub/release-84/gtf/
+ rattus_norvegicus/
+ Rattus_norvegicus.Rnor_6.0.84.chr.gtf.gz")</pre>
```

URLs may be obtained as direct links from: http://useast.ensembl.org/info/data/ftp/index.html. Click on the "GTF" link under the "Gene sets" column for a particular species and then right-click (or command-click on Mac OS X) the name of the file containing the species name/version number and file extension chr.gtf.gz, Homo_sapiens.GRCh38.84.chr.gtf.gz, Mus_musculus.GRCm38.84.chr.gtf.gz, etc.), and copy the link address. Then, paste the link address into the read GFF() as shown above. Alternatively, you can download the GTF file and place it directly in your local working directory. The command above will create an R dataframe object containing the respective GTF file.

2.2 Loading and preprocessing ChIP-seq peak data

Next, the user must input their peak data from a peak caller (e.g., SICER, MACS2, etc.). The peak data must contain only three tab-delimited columns (chromosome number, peak start, and peak end) and a header containing: "chr", "start", and "end". See <code>?samplepeaksinput</code> for an example. Once the peak input data (e.g., "somepeaksfile.txt") has been assembled properly (i.e., to contain only the three tab-delimited columns and header above), it must be properly formatted prior to the execution of downstream analyses.

First, the user must set their working directory to point to the location of their peak data file. Then type the following command:

```
> peaksInput("somepeaksfile.txt")
```

This command properly formats the user's peaks file in preparation for subsequent analyses, producing a resultant "peaks.txt" file in the user's working directory¹.

To see how the above command works using a built-in example, the geneXten deR package provides a peak input dataset² called "somepeaksfile.txt", which can be loaded into memory like this:

This creates a properly formatted (i.e., properly sorted) "peaks.txt" file in the user's working directory.

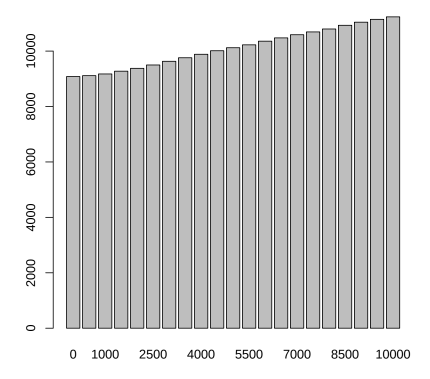
Now, we may use the R object that we created with readGFF() earlier to create a bar chart visualization showing the number of peaks that are sitting directly on top of genes across a series of upstream extensions (of each gene in a genome):

¹Similarly, users can transform their peaks file into a file of merged peaks (see peaksMerge()) and use the resultant "peaks.txt" file instead for the subsequent analysis.

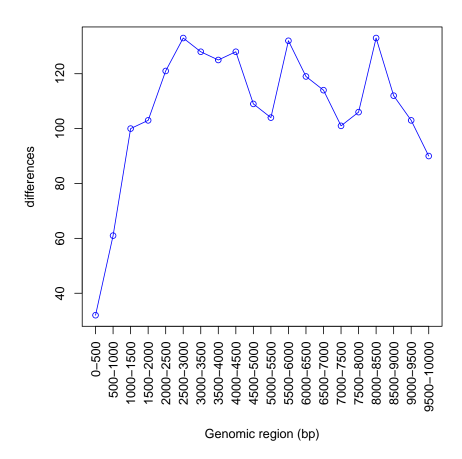
²This peaks dataset comes from a ChIP-seq investigation of brain tissue (pre-frontal cortex) in alcohol addiction and dependence (Barbier et al. 2016), see References section for details.

2.3 Charting ChIP-seq peaks into visualizations and tidy tables

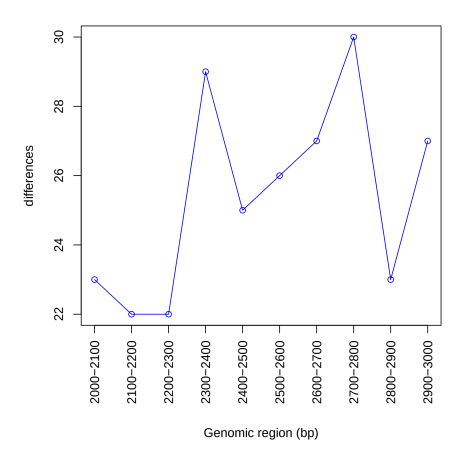
> barChart(rat, 0, 10000, 500)



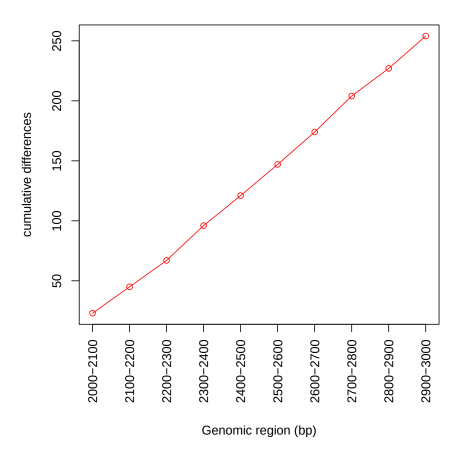
This command first generates 21 individual whole-genome files: 0, 500, 1000, ..., and 10000 bp upstream extension files for the rat (*Rattus norvegicus*) genome, each having an automatic 500 bp downstream extension. In other words, each gene in the rat genome is extended upstream and downstream by some user-specified distance, thereby creating a "gene-sphere." As such, this bar chart command visualizes the raw count of the number of peaks that are sitting on top of genes at each individual upstream cutoff. Clearly, the wider the gene-sphere, the more peaks-on-top-of-genes are found throughout the genome. However, the law of diminishing returns begins to kick in at increasing upstream extension levels (see linePlot() for a visual representation):



In this line plot, there is a sharp rise in the number of peaks-on-top-of-genes from a 0 bp upstream extension to a 1500 bp upstream extension, and from a 2000 bp upstream extension to a 3000 bp upstream extension. This steady rise up until 3000 bp is followed by a steady decline at subsequent extension levels followed by some noisy fluctuations. It may be interesting to investigate what is going on in the interval from 2000 bp to 3000 bp:

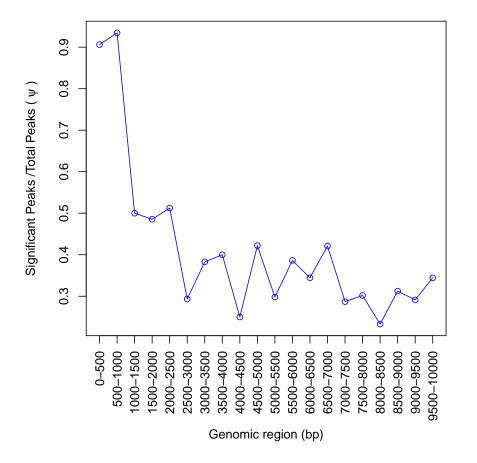


Visually, there is a relative spike in the number of peaks-on-top-of-genes at the 2400 bp upstream extension (as compared to the 2300 bp extension). This spike then drops back down at subsequent extension levels and fluctuates in a noisy manner. However, a cumulative line plot shows that this "spike" is more of a visual effect than anything else, since the graph is almost perfectly linear:



Hence, one very useful function in <code>geneXtendeR</code> is called <code>hotspotPlot()</code>, which allows users to examine the ratio of statistically significant peaks³ to the total number of peaks at each genomic interval (e.g., 0-500 bp upstream of every gene in the genome, 500-1000 bp upstream of every gene in the genome, etc.).

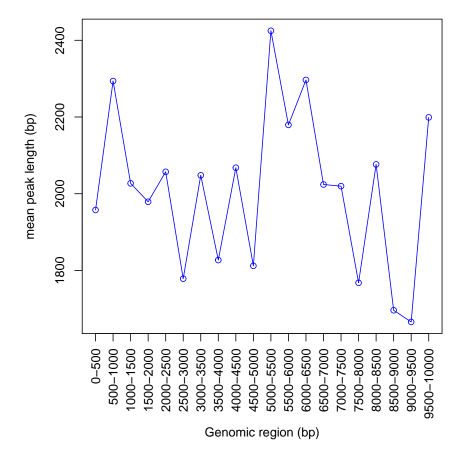
³Note that statistical significance is set apriori by the user at the peak calling stage (prior to geneXtendeR) to give the user the freedom to choose how to filter out peak coordinates that only pass specific p-value and FDR cutoffs from a peak caller. Peak caller output (e.g., from SICER) gives both p-value and FDR measures for each peak, thereby making it easy to extract only the peak coordinates that pass a specific set of statistical cutoff criteria.



This line plot shows that the concentration of significant peaks in this dataset (Barbier et al. 2016) is highest between 0 and 1000 bp upstream of a gene, with over 90% of peaks in these regions being statistically significant. In contrast, between 1000 bp and 2500 bp, only about half of the total peaks contained in these intervals are significant. Statistical significance then fluctuates noisly at further upstream genomic intervals, but with at least a quarter (25%) of the total peaks in these further upstream regions being statistically significant. As such, the take-home message is that genomic regions within the first 1000 bp upstream of their respective genes are most likely to contain significant peaks (relative to the total peak count in these regions) and are therefore hotspots, but regions beyond this also contain a fair share of statistically significant peaks.

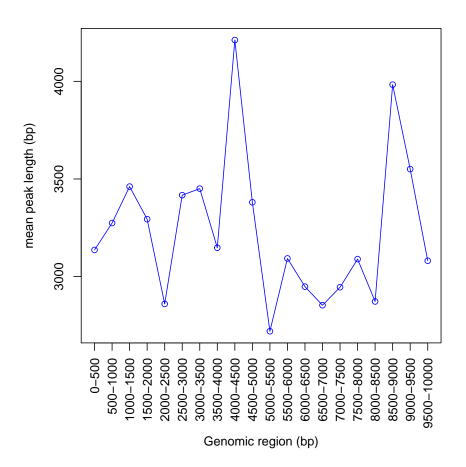
One interesting area to investigate is the variance in the broadness of significant (or total) peaks across different genomic intervals⁴. In other words, asking questions like "are statistically significant peaks that are located very close to their nearest gene (e.g., 0-500 bp away) wider or narrower than peaks located 500-1000 bp away from their nearest gene?". To answer this question we can do:

⁴One can either observe the global distribution of peak lengths within specific genomic intervals (see ?peak LengthBoxplot()), or observe the global distribution of peak lengths across all intervals (see ?allPeak Lengths()).

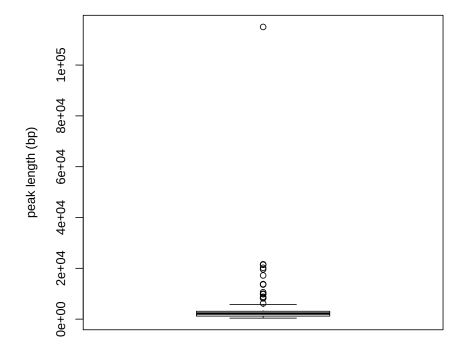


This line plot displays the mean (average) length of all significant peaks found within each genomic interval. Clearly, the "average peak" is slightly narrower in 0-500 bp intervals than in 500-1000 bp intervals yet, overall, peak lengths tend to fluctuate more or less stochastically at various intervals. To get the exact peak length, we can do:

So the mean peak length in the interval 0-500 bp is approximately 1958 bp. Although we see that there is no specific interval with peaks of extraordinary average lengths, it is still possible to see peak length outliers in certain cases (especially when looking at total peak sets):



We see that the 4000-4500 bp and 8500-9000 bp intervals both look quite different in terms of their mean peak lengths relative to the other intervals. To see if the mean might be influenced by a strong outlier(s), we can do:



This box-and-whisker plot shows a clear outlier, which is an example of a very broad peak. We can find the exact length of this outlier peak using:

```
> peak_lengths <- peakLengthBoxplot(rat, 4000, 4500)
> max(peak_lengths)
[1] 114999
```

So this outlier peak measures 114999 bp in total length, therefore making it an extremely broad peak. To see what nearest gene it resides to, we can first extract the peak's index by:

```
> peak_lengths <- peakLengthBoxplot(rat, 4000, 4500)
> match(114999, peak_lengths)
[1] 126
```

which returns the index of where this peak length is found. Then the following command finds all unique peaks that reside between 4000 and 4500 bp upstream of their nearest gene:

> dis	stinct(rat,	4000,	4500))			
	Chromosome	Peak-S	Start	Peak-End	Gene-Chr	Gene-Start	Gene-End
1:	1	1952	26200	19526799	1	19520708	19526671
2:	1	6163	30800	61631999	1	61624941	61630954
3:	1	7134	16800	71347999	1	71334629	71347133
4:	1	9838	35400	98394199	1	98394160	98403468
5:	1	10109	99600	101101399	1	101086377	101100094
124:	18	6000	06800	60007199	18	59985860	60007069
125:	19	4549	99400	45499799	19	45499420	45507827
126:	19	5487	77400	54992399	19	54871853	54877469
127:	20	3061	L0800	30620799	20	30606026	30611101
128:	100	7301	L7400	73018799	100	73018667	73024598
		Gene-1	[D	Gene-Nan	ne Distand	ce	
1:	ENSRN0G0000	9003079	96 AAE	3R07000595.	1	0	
2:	ENSRN0G0000	9002594	19	Vom1r2	22	0	
3:	ENSRN0G0000	9004901	L4 L	_OC10091226	53	0	
4:	ENSRN0G0000	9003733	31	Cd3	33	0	
5:	ENSRN0G0000	9002058	33	Fcgr	^t	0	
124:	ENSRN0G0000	9001785	52	Nar	^S	0	
125:	ENSRN0G0000	9005355	51 AAE	BR07043877.	1	0	
126:	ENSRN0G0000	9002857	78 AAE	BR07044065.	1	0	
127:	ENSRN0G0000	9004916	57 AAE	BR07044988.	1	0	
128:	ENSRN0G0000	9002798	BO AAE	BR07039245.	1	0	

where we see that index 126 belongs to gene AABR07044065.1⁵. Checking the arithmetic difference between column 3 and column 2 for this specific row verifies 114999, as these two columns represent the peak start position and peak

⁵This peak may not be statistically significant, but how could it be if it's so huge? In situations like this, it may be a good idea to check what is known about the gene already: http: //www.pantherdb. org/genes/ gene.do?acc= RAT%7CEnsembl% 3DENSRNOG00000028578% 7CUniProtKB% 3DA0A0G2K0W2. Clearly, not much is known yet.

end positions. Now let's identify what the other columns represent by running the distinct() function again (but this time on a smaller interval to have less output printed to the screen):

```
> fpath <- system.file("extdata", "somepeaksfile.txt",
                        package="geneXtendeR")
> peaksInput(fpath)
> distinct(rat, 2300, 2400)
    Chromosome Peak-Start
                            Peak-End Gene-Chr Gene-Start
                                                           Gene-End
1:
                  79718600
                            79725199
                                                  79725197
                                                            79728613
 2:
             1
                188715600 188716999
                                                 188688243 188715680
 3:
             1
                 214368800 214373199
                                                 214373115 214386385
 4:
                221669800 221671199
                                                 221671190 221694018
             1
                                             1
 5:
             1
                236532800 236534799
                                             1
                                                 236529431 236532885
             3
                  82239000
                            82242199
                                             3
                                                  82096568 82239064
 6:
 7:
             3
                            82784599
                                             3
                                                  82762362 82780214
                  82780200
 8:
             3
                146409600 146412399
                                             3
                                                 146376328 146409652
 9:
             3
                165702800 165706799
                                             3
                                                 165678807 165702889
10:
             4
                  84850400
                            84851999
                                                  84851986 84872257
11:
                118157000 118157799
                                             4
                                                 118157747 118166562
             4
12:
             4
                171955800 171956999
                                             4
                                                 171956961 171961084
13:
             4
                 180237200 180239199
                                             4
                                                 180231882 180237204
14:
             5
                  36437600
                            36438199
                                             5
                                                  36433358
                                                            36437694
                                                  69035218 69038218
15:
             5
                  69038200
                            69039399
                                             5
16:
             5
                121456000 121457199
                                                 121451803 121456072
              5
                153628200 153630199
                                             5
                                                 153568245 153628269
17:
18:
             7
                  14586000
                            14587199
                                             7
                                                  14587120 14615369
19:
             7
                                             7
                  75225000
                            75225799
                                                  75225775
                                                            75249569
                133130600 133133199
20:
             8
                                             8
                                                 133126720 133130690
21:
            10
                   1830200
                             1832199
                                            10
                                                   1832118
                                                             1841132
22:
                            80316799
                                                  80316777
            11
                  80315400
                                            11
                                                            80332099
23:
            14
                            76654999
                                            14
                                                  76654911
                  76654000
                                                            76833661
                 103716400 103719199
                                                 103711769 103716440
24:
            14
                                            14
                    631200
25:
            16
                              642399
                                                    517332
                                            16
                                                               631224
26:
            16
                   9020200
                             9020999
                                            16
                                                   9020987
                                                             9055164
27:
            16
                  75363800
                            75364599
                                            16
                                                  75364529
                                                            75368406
28:
            20
                   1747000
                             1747399
                                            20
                                                   1747316
                                                             1751142
29:
            20
                  22423400
                            22426199
                                            20
                                                  22420251
                                                            22423425
    Chromosome Peak-Start
                            Peak-End Gene-Chr Gene-Start
                                                            Gene-End
                Gene-ID
                             Gene-Name Distance
 1: ENSRN0G00000026891
                            AC093995.1
                                                0
 2: ENSRN0G00000016013
                                 Gprc5b
                                                0
```

3:	ENSRN0G00000018367	Taldo1	0
4:	ENSRN0G00000027456	Cdc42bpg	0
5:	ENSRN0G00000022308	L0C103691298	0
6:	ENSRN0G00000008758	Tspan18	0
7:	ENSRN0G00000042533	Accsl	0
8:	ENSRN0G00000006795	Apmap	0
9:	ENSRN0G00000042101	Zfp93	0
10:	ENSRN0G00000010205	Mturn	0
11:	ENSRN0G00000016273	Fam136a	0
12:	ENSRN0G00000057540	AABR07062363.1	Θ
13:	ENSRN0G00000048961	Bhlhe41	Θ
14:	ENSRN0G00000055329	AABR07047528.1	Θ
15:	ENSRN0G00000060997	U6	0
16:	ENSRN0G00000045614	L0C102552337	0
17:	ENSRN0G00000018109	Clic4	0
18:	ENSRN0G00000048450	Cyp4f37	0
19:	ENSRN0G00000061463	AABR07057510.3	0
20:	ENSRN0G00000006730	Ccr1l1	0
21:	ENSRN0G00000040121	RGD1565158	0
22:	ENSRN0G00000022160	Rtp2	0
23:	ENSRN0G00000051169	Clnk	0
24:	ENSRN0G00000054704	AABR07016558.1	Θ
25:	ENSRN0G00000061982	AABR07024473.2	Θ
26:	ENSRN0G00000042628	RGD1561145	Θ
27:	ENSRN0G00000029462	Defal1	Θ
28:	ENSRN0G00000050043	0lr1735	Θ
29:	ENSRN0G00000057124	AABR07044824.1	Θ
	Gene-ID	Gene-Name	Distance

This data table shows 29 separate entries sorted by chromosome and start position. Gene-ID refers to the Ensembl ID and the other columns named accordingly. It should be noted that the X chromosome is designated by the integer 100, the Y chromosome by the integer 200, and the mitochondrial chromosome by the integer 300. This is done for sorting purposes (see ?peaksIn put for details). In short, the distinct() command finds what peaks-on-top-of-genes would be missed if a 2300 bp upstream extension is used instead of a 2400 bp extension. In other words, these 29 genes all reside between 2300-2400 bp upstream of their nearest gene.

Once the user has chosen the specific upstream extension to be used, the peak file is ready to be fully annotated:

> annot	tate(rat, 24	400)				
	Chromosome	Peak-Start	Peak-End	Gene-Start	Gene-End	Gene-ID
1:	1	48800	51199	394300	410176	ENSRN0G00000046319
2:	1	53000	53799	394300	410176	ENSRN0G00000046319
3:	1	265600	266999	394300	410176	ENSRN0G00000046319
4:	1	506600	507999	394300	410176	ENSRN0G00000046319
5:	1	669400	672199	697013	708565	ENSRN0G00000047964
25085:	100	159818600	159820599	159723366	159843472	ENSRN0G00000000869
25086:	100	159821400	159823199	159723366	159843472	ENSRN0G00000000869
25087:	100	159898400	159899599	159889343	159892315	ENSRN0G00000054559
25088:	100	159913800	159915199	159889343	159892315	ENSRN0G00000054559
25089:	100	159947000	159948599	159889343	159892315	ENSRN0G00000054559
	Gene - Nar	ne Distance	-of-Gene-to	o-Nearest-Pe	eak	
1:	Vom2			3613	377	
2:	Vom2	r3		357	177	
3:	Vom2			1445	577	
4:	Vom2				425	
5:	L0C10090960	98		248	815	
25085:	Arhge ⁻				0	
25086:	Arhge ⁻				0	
25087:	SNORD				986	
25088:	SNORD				486	
25089:	SNORD	51		546	586	

which generates a fully annotated peaks outfile (in the user's working directory) containing various genomic features and labeled headers. An example of which is above.

2.4 Exploring functional annotation in depth

If a user is looking for a more gene-centric approach to annotation (as briefly outlined in Section 1.2.3), they may use either the <code>gene_lookup()</code> or <code>gene_annotate()</code> functions. The <code>gene_annotate()</code> function builds off of the <code>annotate()</code> function, but reorganizes and groups the output based on relevant gene information. If you (the reader) are just joining us now in the vignette and have not yet run any of the command on preceding pages, first just run the following commands:

Now do:

```
> head(gene_annotate(rat, 2400))
  Chromosome Gene-Start Gene-End
                                               Gene-ID Gene-Name
1
          12
               14448510 15101186 ENSRN0G00000001103
                                                             Sdk1
2
             168141047 168736696 ENSRN0G00000018602
                                                           Camta1
3
           8 127268889 127573488 ENSRN0G00000043167
                                                            Itga9
4
             106749225 107427829 ENSRN0G00000003738
                                                            Ush2a
5
               18557628 18944940 ENSRN0G00000005365
          10
                                                           Kcnip1
6
               51385263 51705130 ENSRN0G00000032590
          12
                                                            Ttc28
  Peaks-on-Gene-Body Mean-Distance-of-Gene-to-Nearest-Peaks
                                                                     sd
1
                  32
                                                     6290.222 19899.71
2
                  21
                                                         0.000
                                                                   0.00
3
                  20
                                                         0.000
                                                                   0.00
4
                   20
                                                         0.000
                                                                   0.00
5
                  19
                                                         0.000
                                                                   0.00
6
                   19
                                                         0.000
                                                                   0.00
  Number-of-Peaks-Associated-with-Gene
1
                                      36
2
                                      21
3
                                      20
4
                                      20
5
                                      19
6
                                      19
```

This output labels each gene and matches it with the number of peaks that overlap it and are "first away" from its gene-body (i.e., closest/nearest but not overlapping). Distance is calculated between 5-prime end of gene and 3-prime end of peak (or 3-prime end of gene and 5-prime end of peak, whichever is smallest). The table is sorted by number of peaks on gene body (i.e., number of peaks that directly overlap the gene body) and include extra information such as mean and standard deviation (sd) for extra validation. Typically, a user would be looking for genes that have a high number of Peaks-on-Gene-Body to follow-up on for experimental validation. Genes that have peaks that reside

close (but not overlapping) to the chosen gene-body (i.e., low mean) and that are clustered together spatially (i.e., low standard deviation) may also be good targets for follow-up analysis.

An example of how the <code>gene_annotate()</code> function is intended to be used is below, where we highlight three specific rows to highlight key points of the discussion:

```
> gene_annotate(rat, 2400)[c(1, 7, 11),]
   Chromosome Gene-Start Gene-End
                                               Gene-ID Gene-Name
1
           12
                14448510 15101186 ENSRNOG00000001103
                                                            Sdk1
7
            8
                52984813 53149353 ENSRN0G00000029980
                                                          Zbtb16
           19
11
                20144637 20406503 ENSRN0G00000014658
                                                          Zfp423
   Peaks-on-Gene-Body Mean-Distance-of-Gene-to-Nearest-Peaks
                                                                       sd
1
                    32
                                                     6290.2222 19899.710
7
                    17
                                                      740.1579 2336.913
11
                    13
                                                    19803.6500 33367.643
   Number-of-Peaks-Associated-with-Gene
1
                                       36
7
                                       19
11
                                       20
```

These three genes exemplify three of the four different scenarios that may occur in this table. The difference between the mean and the standard deviation of the peaks located closest to a specific gene can be used to judge the distribution of those peaks, thereby indicating what may or may not be worth following up on in the wet-lab experiments.

- 1. The first gene has 32 peaks on the gene-body of "Sdk1" (i.e., 32 peaks that overlap a 2400 bp upstream and 500 bp downstream extension of "Sdk1"), with a total of 36 genes annotated to the "Sdk1" gene body in total. The high SD and mean (relative to the fact that 32/36 of these genes reside on the gene-body itself) indicate that the other 4 peaks that do not reside on gene-body, also do not reside near enough to the gene to warrant biological meaning. In other words, focus on the 32 peaks on the gene-body itself and not the other 4.
- 2. The second gene has both a low mean distance as well as a relatively low standard deviation, which indicates that peaks not residing on the extended gene-body are still quite close to it and clustered together spatially at approximately the same genomic location (possibly a proximal-promoter region). "Zbtb16" is definitely a good gene to follow-up on

because the peaks are close enough to the gene body to be considered biologically important (e.g., might reside in important proximal-promoter regions of the gene).

- 3. The third gene showcases the default case, in which both the mean and sd are relatively high. The peaks that do not reside on "Zfp423" are not close or clustered together either, based on the spread of the mean and standard deviation, so the 7 additional peaks are probably unnecessary for use in a follow-up of that gene. The reason why the geneXtendeR package registers these 7 peaks in the first place (even though their mean distance is 19804 bp from their nearest genes) is because these peaks are located in intergenic regions where "Zfp423" just so happens to be the closest gene.
- 4. The final case is the rarest case, when the mean is high but the standard deviation is low. This indicates that the peaks are grouped, but located far away from the closest gene-body. This may be another case worth following up on, especially in the context of long-range interactions (e.g., trans-regulatory elements).

It should be noted that mean = 0 (i.e, Mean-Distance-of-Gene-to-Nearest-Peaks = 0) denotes cases where all peaks are overlapping a given gene body.

The <code>gene_lookup()</code> function looks up all peaks surrounding a specific gene or list of genes across all chromosomes and reports these peaks. This method is extremely useful when paired with <code>gene_annotate()</code> to check genes that may be used in a follow-up.

 $> gene_lookup(rat, c("Zbtb16"), n = 19, extension = 2400)$ Chromosome Peak-Start Peak-End Distance-to-Gene Gene-Start Gene-End Gene 1: 52983400 52986999 52984813 53149353 Zbtb16 2: 8 52988000 52988999 0 52984813 53149353 Zbtb16 8 52984813 53149353 Zbtb16 3: 52989600 52992199 4: 8 52993000 52995799 0 52984813 53149353 Zbtb16 5: 8 52998400 53004399 52984813 53149353 Zbtb16 6: 8 53006200 53009399 0 52984813 53149353 Zbtb16 7: 8 53024400 53031999 52984813 53149353 Zbtb16 8: 8 53038200 53040799 0 52984813 53149353 Zbtb16 9: 8 53044000 53046399 0 52984813 53149353 Zbtb16 52984813 53149353 Zbtb16 10: 8 53084800 53085799 0 8 11: 53090800 53094999 0 52984813 53149353 Zbtb16 12: 8 53096200 53099999 52984813 53149353 Zbtb16 0 53101000 53105799 52984813 53149353 Zbtb16 13: 8 0 14: 53106600 53110399 52984813 53149353 Zbtb16

15:	8	53119600	53132199	0	52984813	53149353	Zbtb16
16:	8	53132800	53135799	0	52984813	53149353	Zbtb16
17:	8	53138000	53152999	0	52984813	53149353	Zbtb16
18:	8	52946600	52979999	4814	52984813	53149353	Zbtb16
19:	8	53158600	53163799	9247	52984813	53149353	Zbtb16

This output shows all the peaks nearest to "Zbtb16" and their respective distances. Knowing these genomic peak coordinates facilitates the design of PCR primers. Although 17/19 of the peaks reside on the extended gene-body (2400 bp upstream extension, 500 bp downstream extension), the two additional peaks are still close enough to be considered for analysis. Out of all the genes on that specific chromosome, these two nearby peaks are located closest to the gene "Zbtb16".

In gene_lookup(organism, gene_name, n, extension), n represents the number of nearest (and overlapping) peaks to a given gene. We saw from gene_annotate() that, in the case of "Zbtb16," there are 19 nearest (and overlapping) peaks to the gene and gene_lookup() displays their location as well as their distance from the gene. This function is motivated by the need of biologists to accurately design primers for specific genomic loci in order to experimentally validate the existence (realness) of a peak.

For a much more in-depth analysis, a function that combines both <code>gene_lookup()</code> and <code>gene_annotate()</code> has been provided as <code>annotate_n()</code>. Instead of simply annotating a peak to a single closest gene (and reporting any overlapping peaks on gene bodies), this function annotates each peak to the closest, the second-closest, ..., to the nth-closest genes to provide the user an expanded picture of the peaks layout for further analysis. Called, this function looks like:

> annotate_n(rat, 3500, n = 3)

	Peak-Num	Chromosome	Peak-Start	Peak-End	Gene-Start	Gene-End
1:	1	1	48800	51199	393200	410176
2:	1	1	48800	51199	695913	708565
3:	1	1	48800	51199	744116	759145
4:	2	1	53000	53799	393200	410176
5:	2	1	53000	53799	695913	708565
75263:	25088	100	159913800	159915199	159889343	159893415
75264:	25088	100	159913800	159915199	159723366	159844572
75265:	25089	100	159947000	159948599	159884385	159894826
75266:	25089	100	159947000	159948599	159889343	159893415
75267:	25089	100	159947000	159948599	159723366	159844572
		Gene-ID	Gene-Name	e rank Min	imum-Distand	ce-to-Gene

1:	ENSRN0G00000046319	Vom2r3	1	342001
2:	ENSRN0G00000047964	L0C100909608	2	644714
3:	ENSRN0G00000050370	Vom2r6	3	692917
4:	ENSRN0G00000046319	Vom2r3	1	339401
5:	ENSRN0G00000047964	L0C100909608	2	642114
75263:	ENSRN0G00000054559	SNORD61	2	20385
75264:	ENSRN0G00000000869	Arhgef6	3	69228
75265:	ENSRN0G00000000866	Rbmx	1	52174
75266:	ENSRN0G00000054559	SNORD61	2	53585
75267:	ENSRN0G00000000869	Arhgef6	3	102428

This function is the most versatile of the annotation functions provided and is designed for the purpose of providing peak-to-gene associations and follow-up information that goes beyond just a simple closest genomic distance criterion. Future work in this direction can address three-dimensional genome interactions (when coupled with methods like Hi-C). When moving away from the traditional "first closest gene" to a peak, this method opens up many more possibilities as to which peaks influence which genes. It increases the scope of the individual peaks to reduce the chance that a peak that influences any particular gene is missed or misattributed to the wrong gene.

2.5 Gene Ontology functions

It may be of interest to note the differential gene ontologies between the following two upstream extensions:

```
> library(org.Rn.eg.db)
> library(G0.db)
> x <- diffGO(rat, 2300, 2400, BP, org.Rn.eg.db)
> head(x, 20)
                      GOID
   gene$SYMB0L
1
        Gprc5b G0:0001934
2
        Gprc5b G0:0007186
3
        Gprc5b G0:0007626
4
        Gprc5b G0:0010976
5
        Gprc5b G0:0032147
6
        Gprc5b G0:0042593
7
        Gprc5b G0:0043123
        Gprc5b G0:0045666
```

```
9
        Gprc5b G0:0045860
10
        Gprc5b G0:0050729
11
        Gprc5b G0:0060907
12
        Gprc5b G0:0061098
        Gprc5b G0:0090263
13
14
        Taldo1 G0:0005975
15
        Taldo1 G0:0006002
        Taldo1 G0:0006098
16
17
        Taldo1 G0:0009052
18
        Taldo1 G0:0019682
19
      Cdc42bpg G0:0006468
      Cdc42bpg G0:0031532
20
                                                           TERM
1
               positive regulation of protein phosphorylation
2
                 G-protein coupled receptor signaling pathway
3
                                           locomotory behavior
         positive regulation of neuron projection development
4
5
                         activation of protein kinase activity
6
                                           glucose homeostasis
7
   positive regulation of I-kappaB kinase/NF-kappaB signaling
8
                positive regulation of neuron differentiation
9
               positive regulation of protein kinase activity
10
                 positive regulation of inflammatory response
        positive regulation of macrophage cytokine production
11
12
      positive regulation of protein tyrosine kinase activity
13
       positive regulation of canonical Wnt signaling pathway
14
                                carbohydrate metabolic process
15
                       fructose 6-phosphate metabolic process
16
                                       pentose-phosphate shunt
17
                pentose-phosphate shunt, non-oxidative branch
18
                 glyceraldehyde-3-phosphate metabolic process
19
                                       protein phosphorylation
20
                             actin cytoskeleton reorganization
```

This dataframe shows the first 20 unique gene ontology terms, their IDs, and respective gene symbols. Clearly, gene name *Gprc5b* has several BP ontologies related explicitly to the brain, while *Taldo1* does not. Considering that the ChIP-seq peaks dataset used as input into geneXtendeR comes from a ChIP-seq study investigating the prefrontal cortex, this suggests that a 2400 bp extension may be more suitable for this brain dataset. However, such decisions are left entirely to the discretion and judgment of the user in deciding the relative importance

of specific genes and their respective GO terms (BP, CC, or MF) to the goals of the computational analysis (as well as plans for experimental follow-up and validation). See Discussion section for details.

It is also critical to note that the diffGO() function returns ALL known gene ontologies, NOT a gene ontology enrichment analysis (more about this in Discussion section). The goal is to provide users with knowledge regarding all possible known roles of any given gene. For example, by knowing that a potential gene candidate has previously been linked with known brain-related ontologies, a user may be prompted to look more closely into the relevant literature behind this gene and its implications to the biological question under study (before embarking on making a decision about its potential impact and suitability as a good candidate for experimental validation).

Furthermore, a user may plot the differential gene ontology results as an interactive network:

```
> library(networkD3)
```

- > library(org.Rn.eg.db)
- > library(dplyr)
- > makeNetwork(rat, 2300, 2400, BP, org.Rn.eg.db)

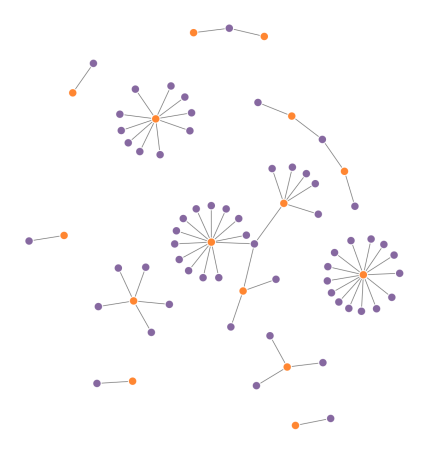


Figure 1: Orange color denotes gene names, purple color denotes GO terms A user can hover the mouse cursor over any given node to display its respective label directly within R Studio. Likewise, users can dynamically drag and re-organize the spatial orientation of nodes, as well as zoom in and out of them for visual effect.

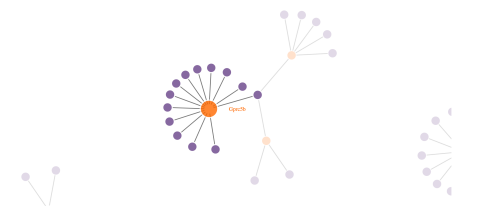


Figure 2: Orange color denotes gene names, purple color denotes GO terms A user can hover the mouse cursor over any given node to display its respective label directly within R Studio. Likewise, users can dynamically drag and reorganize the spatial orientation of nodes, as well as zoom in and out of them for visual effect.

In addition, users can generate word clouds comprised from words present in their GO terms:

- > library(tm)
- > library(SnowballC)
- > library(wordcloud)
- > library(RColorBrewer)
- > makeWordCloud(rat, 2300, 2400, BP, org.Rn.eg.db)



Figure 3: Word cloud generated from words comprising gene ontology terms of category BP

This word cloud shows the words that are used within BP gene ontology terms of peaks found to be present between 2300 and 2400 bp upstream of their nearest genes.

It may also be of interest to visually examine the most frequently used words found within GO terms:

- > library(tm)
- > library(SnowballC)
- > library(wordcloud)
- > library(RColorBrewer)
- > plotWordFreq(rat, 2300, 2400, BP, org.Rn.eg.db, 10)

Most frequent words found within GO terms

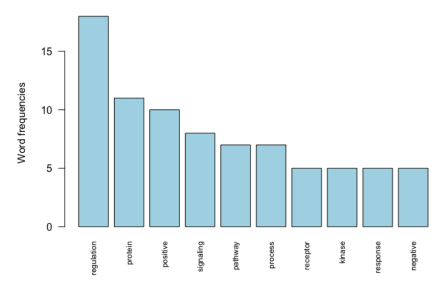
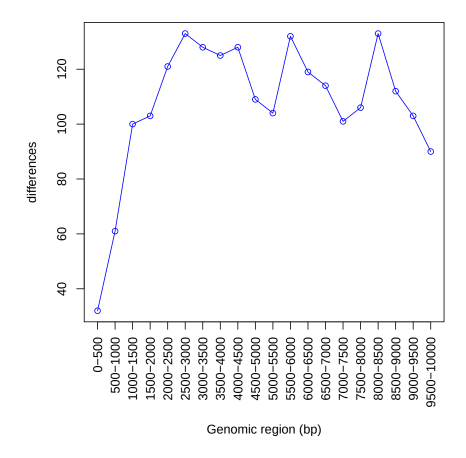


Figure 4: This barplot shows the top 10 words used within gene ontology terms (specific to BP) of peaks found to be present between 2300 and 2400 bp upstream of their nearest genes

3 Discussion

Even though geneXtendeR is designed to compute (and analyze/display) optimal gene extensions tailored to the characteristics of a specific peak input file, geneXtendeR will not explicitly impose on the user the optimal extension to select, since this information is highly study-dependent and, as such, is ultimately reserved to the user's discretion. For example, a user may choose a conservatively lower upstream extension (e.g., for studies investigating narrow peaks such as H3K4me3 or H3K9ac that exhibit a compact and localized enrichment pattern, where high upstream extensions may begin to lose biological relevance). An example of such a user-driven decision would be the selection of a 1500 bp upstream extension instead of a 3500 bp extension in situations like this:



This line plot is derived from the input peak dataset used from the H3K9me1 study examined earlier (Barbier et al. 2016). If the study had examined a narrower chromatin mark (e.g., H3K4me3) then the decision process for choosing an optimal extension may have been different.

In certain cases, additional extensions are unlikely to add significant value to the annotation of the peak file. Taking the example of the 0-10000 bp line plot, an upstream extension beyond 3500 bp globally across every gene in a genome would most likely not accurately reflect the biology of the peak input file (since such large global upstream extensions are likely to reach considerably beyond known proximal promoter elements, especially for relatively narrow histone marks or transcription factors). Such assumptions may be validated directly by the user by investigating the p-value and FDR of specific peaks using a combination of HT-seq (to count the reads) and edgeR/DESeq2 (to assess statistical significance). As such, geneXtendeR is designed to be used as part of a biological workflow involving subsequent statistical analysis:

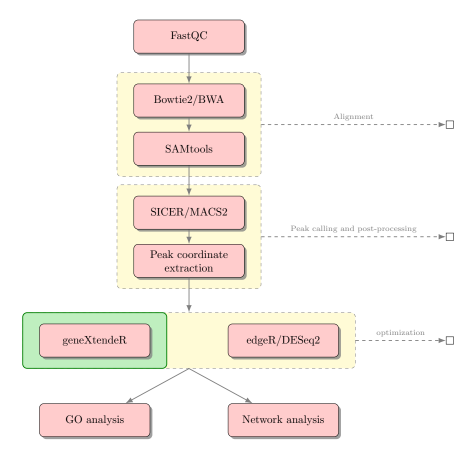
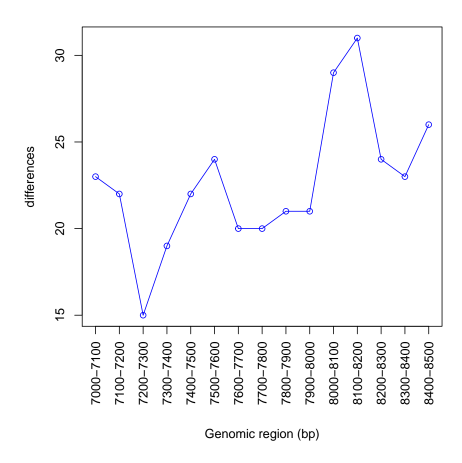


Figure 5: Sample biological workflow using geneXtendeR in combination with existing statistical software to analyze peak significance

Subsequent gene ontology enrichment or network analysis may be conducted on genes associated with statistically significant peaks.

It is entirely possible (and probable) for significant peaks to be present at relatively high upstream extension levels (i.e., large gene-spheres), albeit these significant peaks may be associated with biology not directly relevant to the study at-hand, due mainly to the sheer magnitude of the distance of the peak from traditional gene boundaries (where traditional gene boundaries may be loosely defined as $+/-\approx 3$ kb from TSS and $+/-\approx 0.5$ kb from TES). Consequently, it is likely for peaks-on-top-of-genes to exhibit higher levels of noise at higher upstream extension levels. Nevertheless, this does not mean that potential enhancer activity should be discounted. For instance, it is not uncommon to see a steady rise or even a surge in the number of peaks-on-top-of-genes at higher upstream extension levels:



This line plot shows that there are over 30 peaks in this dataset (across the rat genome) that reside between 8100 and 8200 bp upstream of their nearest gene. In far-out cases like this, it is particularly recommended to examine the statistical significance of peaks to get a sense for the possibility of potential enhancer activity/regulation. Of course, such computational findings would require experimental follow-up and/or database mining for known motifs. Assessment of such statistical significance values is beyond the scope of geneXtendeR, in order to allow the user freedom to choose the most appropriate statistical package/technique for their analysis. As before, first use the distinct() function to create a table of unique genes located under peaks between the two upstream extension levels:

> distinct(rat, 8100, 8200)

Then, assess the statistical significance of these peaks using a combination of HT-seq (Anders et al. 2015) and edgeR (Robinson et al. 2010), or HT-seq and DESeq2 (Love et al. 2014), or some other appropriate combination of existing software tools. Genes associated with the resultant statistically significant peaks may then be further assessed with gene ontology enrichment analysis to help answer a variety of interesting research questions. It should once again be noted that the diffGO() function does NOT perform gene ontology enrichment analysis. Instead, it returns all known gene ontologies for each gene. The purpose and utility of this is described in the previous section.

Moreover, DNA sequences under peaks may be checked for the presence of known regulatory motifs (e.g., using TRANSFAC (Matys et al. 2006) or MEME/JAS PAR (Sandelin et al. 2004, Bailey et al. 2009)), or for the presence of biological repeats (e.g., using RepeatMasker (Smit et al. 2015)). Pending a prospective GO enrichment and network analysis, functional validation may be followed up in the lab to test any potential regulatory sites or prospective enhancer elements, thereby bringing the computational analysis pipeline back to the bench.

In addition to the computational workflows discussed above, geneXtendeR's wide array of functions makes it possible to conduct some rather interesting and creative combinations of genomic analysis. Let's say, for example, that a user wants to explore all known ontological differences across specific disparate sectors of the genome (e.g., 0-500 bp vs. 2000-3000 bp, but removing 501-1999 bp from consideration). In other words, look at all peaks (across the entire genome) that reside between 0-500 bp upstream of their nearest gene (and 2000-3000 bp upstream of their nearest gene), and extract unique gene ontologies that differ between these two variable-length sectors (where one is 500 bp long and the other is 1000 bp in length). This can be accomplished rather conveniently using dplyr:

```
> library(dplyr)
> library(org.Rn.eg.db)
> library(G0.db)
> a <- diffGO(rat, 0, 500, BP, org.Rn.eg.db)
> b <- diffGO(rat, 2000, 3000, BP, org.Rn.eg.db)
> dplyr::filter(b, TERM %in% a$TERM)
   gene$SYMB0L
                     GOID
                                                                           TERM
1
          Sod2 G0:0001889
                                                              liver development
          Sod2 G0:0007507
2
                                                              heart development
3
          Sod2 G0:0008285
                                     negative regulation of cell proliferation
4
          Sod2 G0:0042311
                                                                   vasodilation
          Sod2 G0:0042493
5
                                                               response to drug
          Sod2 G0:0043066
                                      negative regulation of apoptotic process
```

7	Dll1	GO:0001757	somite specification
8	Dll1	G0:0008284	positive regulation of cell proliferation
9	Dll1	G0:0008285	negative regulation of cell proliferation
10	Dll1	G0:0045596	negative regulation of cell differentiation
11	0lr40	G0:0007186	G-protein coupled receptor signaling pathway
12	0lr139	G0:0007186	G-protein coupled receptor signaling pathway
13	0lr282	G0:0007186	G-protein coupled receptor signaling pathway
14	Gprc5b	G0:0007186	G-protein coupled receptor signaling pathway
15	8qpA	G0:0055085	transmembrane transport
16	8qpA	G0:0071320	cellular response to cAMP
17	Cdc42bpg	G0:0006468	protein phosphorylation
18	Dusp5	G0:0045892	$negative\ regulation\ of\ transcription,\ DNA\text{-}templated$
19	Adgrl2	G0:0007166	cell surface receptor signaling pathway
20	Adgrl2	G0:0007186	G-protein coupled receptor signaling pathway
21	Nfe2l2	G0:0016567	protein ubiquitination
22	Nfe2l2	G0:0071456	cellular response to hypoxia
23	0lr559	G0:0007186	G-protein coupled receptor signaling pathway
24	Tspan18	G0:0007166	cell surface receptor signaling pathway
25	Kcnq2	G0:0060081	membrane hyperpolarization
26	Reg3b	G0:0008284	positive regulation of cell proliferation
27	Reg3b	G0:0043066	negative regulation of apoptotic process
28	0lr828	G0:0007186	G-protein coupled receptor signaling pathway
29	Tspan9	G0:0007166	cell surface receptor signaling pathway
30	Bhlhe41	G0:0045892	negative regulation of transcription, DNA-templated
31	Aptx	G0:0006974	cellular response to DNA damage stimulus
32	Ccl21	G0:0007186	G-protein coupled receptor signaling pathway
33	Aldob	G0:0001889	liver development
34	Aldob	G0:0042493	response to drug
35	Clic4	G0:1902476	chloride transmembrane transport
36	Htr1d	G0:0042310	vasoconstriction
37	Nlrc4	G0:0016567	protein ubiquitination
38	Nlrc4	G0:0090307	mitotic spindle assembly
39	Alk	G0:0043066	negative regulation of apoptotic process
40	Esyt1	G0:0006869	lipid transport
41	Sbno2	G0:0045892	negative regulation of transcription, DNA-templated
42	0lr1085	G0:0007186	G-protein coupled receptor signaling pathway
43	Fbxo7	G0:0016567	protein ubiquitination
44	Dnmt1	G0:0042493	response to drug
45	Dnmt1	G0:0045892	negative regulation of transcription, DNA-templated
46	Xcr1	G0:0007186	G-protein coupled receptor signaling pathway
47	Ccr1l1	G0:0007186	G-protein coupled receptor signaling pathway
48	Clcn7	GO:1902476	chloride transmembrane transport

49	L0C684471	GO:0007186	G-protein coupled receptor signaling pathway
50	Il3	G0:0008284	positive regulation of cell proliferation
51	Il3	G0:0043066	negative regulation of apoptotic process
52	0lr1501	G0:0007186	G-protein coupled receptor signaling pathway
53	Socs3	G0:0016567	protein ubiquitination
54	Socs3	G0:0042493	response to drug
55	Socs3	G0:0043066	negative regulation of apoptotic process
56	Fbxw8	G0:0016567	protein ubiquitination
57	Fcgr2b	G0:0007166	cell surface receptor signaling pathway
58	Arhgef10	G0:0090307	mitotic spindle assembly
59	Eef1e1	G0:0008285	negative regulation of cell proliferation
60	F13a1	G0:0007596	blood coagulation
61	Tubb6	G0:0007010	cytoskeleton organization
62	Csnk2a2	G0:0006468	protein phosphorylation
63	Csnk2a2	G0:0051726	regulation of cell cycle
64	0lr1735	G0:0007186	G-protein coupled receptor signaling pathway
>			

This displays all biological process (BP) ontologies present in b that are not present in a. Similarly, one can look at all BP, CC, or MF ontologies present in a that are not present in b.

4 Concluding remarks

geneXtendeR is continually evolving, so any suggestions or new feature requests are always appreciated. Likewise, any bug reports may be posted to https://github.com/Bohdan-Khomtchouk/geneXtendeR/issues or emailed to the package maintainer directly.

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