# An Introduction to Guitar2 Package

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## 1 Quick Start with Guitar2

This is a manual for Guitar2 package. The Guitar2 package is aimed for RNA landmark-guided <u>transcriptomic analysis</u> of RNA-reated genomic featu<u>res</u>.

The Guitar2 package enables the comparison of multiple genomic features, which need to be stored in a name list. Please see the following example, which will read 1000 RNA m6A methylation sites into R, and then splitted them into 3 groups to be examined. Of course, in real data analysis, the 3 groups of features are likely to be from different resources.

```
> library(Guitar2)
> narrowPeak <- system.file(
+    "extdata", "m6A_hg19_1000peaks_macs2.narrowPeak",
+    package="Guitar2")
> # genomic features imported into named list
> m6A_Bcell <- narrowPeaktoGRanges(narrowPeak)
> m6A_Bcell_1 <- m6A_Bcell[1:300]
> m6A_Bcell_2 <- m6A_Bcell[301:600]
> m6A_Bcell_3 <- m6A_Bcell[601:900]
> feature_hg19 <- list(m6A_Bcell_1, m6A_Bcell_2, m6A_Bcell_3)
> names(feature_hg19) <- c("Group_1", "Group_2", "Group_3")</pre>
```

With the following script, we may generate the transcriptomic distribution of genomic features to be tested, and the result will be automatically saved into a PDF file under the working directory with prefix "example". With the Guitar2Plot function, the gene annotation can be downloaded from internet automatically with a genome assembly number provided; however, this feature requires working internet and might take a longer time. The toy Guitar2 coordinates generated internally should never be re-used in other real data analysis.

```
> count <- Guitar2Plot(feature_hg19,
+ genome="hg19",
+ saveToPDFprefix = "example")</pre>
```

In a more efficient protocol, in order to re-use the gene annotation and Guitar2 coordinates, you will have to build Guitar2 Coordinates from a TxDb object

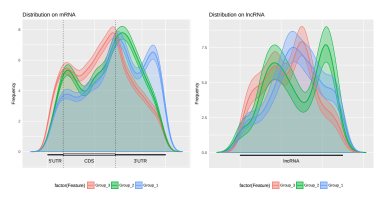
in a separate step. The transcriptDb contains the gene annotation information and can be obtained in a number of ways, .e.g, with command  $\mathtt{makeTxDbFro-mUCSC}$  from GenomicFeatures package to download the complete gene annotation of species from UCSC automatically, which might takes a few minutes. In the following analysis, we load the TxDb object from a toy dataset provided with the Guitar2 package. Please note that this is only a very small part of the complete hg19 transcriptome, and the TxDb object provided with Guitar2 package should not be used in real data analysis.

With a *TxDb* object that contains gene annotation information, we in the next build *Guitar2 coordiantes*, which is essentially a bridge connects the transcriptomic landmarks and genomic coordinates. The parameter noBins=20 defines the resolution of your analysis. Higher resolution will lead to finer details revealed and increased computation time and memory usage.

```
> gc_txdb <- makeGuitarCoordsFromTxDb(txdb, noBins=100)
```

```
[1] "total 1932 transcripts extracted ..."
[1] "total 1043 transcripts left after ambiguity filter ..."
[1] "total 471 mRNAs left after component length filter ..."
[1] "total 148 ncRNAs left after ncRNA length filter ..."
[1] "Building Guitar Coordinates. It may take a few minutes ..."
[1] "Guitar Coordinates Built ..."
```

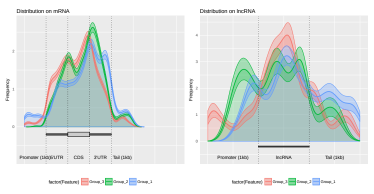
You may now generate the Guitar2 plot from the named list of genome-based features and the prebuilt *Guitar2 coordinates*.



The figure generated reflects the true size of RNA components (5'UTR, CDS and 3'UTR); alternatively, you may cancel the component rescaling, and treat each component equally with option rescaleComponent=FALSE. The figure generated in this way looks clearer, but the relative size of the components will be missing. In this way, 5'UTR, although is mucher smaller, shows up to be of the same size as CDS and 3'UTR.

```
> Guitar2Plot(gfeatures = feature_hg19,
+ GuitarCoordsFromTxDb = gc_txdb,
+ rescaleComponent=FALSE)
```

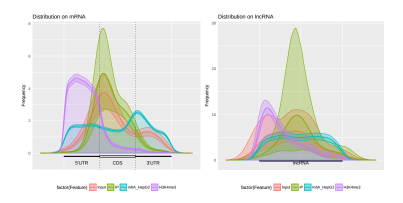
Alternatively, you may also optionally include the promoter DNA region and its complementary region on the 3' side of a transcript in the plot with parameter includeNeighborDNA =TRUE.



## 2 Supported Data Format

Besides BED12 format, *Guitar2* package also supports BAM/SAM and bed3 via GRangesList, GRanges and GAlignments data structures. Please see the following examples.

```
> # import different data formats into a named list object.
> # These genomic features are using mm10 genome assembly
> bed3=system.file("extdata", "H3K4me3_mm10_1000peaks.bed", package="Guitar2")
> bed12=system.file("extdata", "m6A_mm10_exomePeak_1000peaks_bed12.bed", package="Guitar2")
> bam1=system.file("extdata", "866991_part.bam", package="Guitar2")
> bam2=system.file("extdata", "866992_part.bam", package="Guitar2")
> H3K4me3 <- import.bed(bed3) # bed3 imported as GRanges
> m6A_HepG2 <- BED12toGRangesList(bed12) # bed12 imported as GRangesList
[1] "Converting BED12 to GRangesList"
[1] "It may take a few minutes"
> MeRIP_IP <- readGAlignments(bam1) # bam imported as GAlignments
> MeRIP_Input <- readGAlignments(bam2) # bam imported as GAlignments
> feature_mm10 <- list(H3K4me3,m6A_HepG2,MeRIP_IP,MeRIP_Input)</pre>
> names(feature_mm10) <- c("H3K4me3", "m6A_HepG2", "IP", "Input")
> # Build Guitar2 Coordinates
> txdb_file <- system.file("extdata", "mm10_toy.sqlite",</pre>
                           package="Guitar2")
> mm10_toy_txdb <- loadDb(txdb_file)</pre>
> gc_mm10_txdb <- makeGuitarCoordsFromTxDb(mm10_toy_txdb, noBins=30)
[1] "total 2946 transcripts extracted ..."
[1] "total 2272 transcripts left after ambiguity filter ..."
[1] "total 1038 mRNAs left after component length filter ..."
[1] "total 353 ncRNAs left after ncRNA length filter ..."
[1] "Building Guitar Coordinates. It may take a few minutes ..."
[1] "Guitar Coordinates Built ..."
> # Guitar2 Plot
> Guitar2Plot(gfeatures = feature_mm10,
             GuitarCoordsFromTxDb = gc_mm10_txdb,
             rescaleComponent=FALSE)
[1] "Using provided Guitar Coordinates"
[1] "resolving ambiguious features ..."
[1] "no figure saved ..."
```



## 3 Analysis of BS-Seq data report from Bismark

Here is one more example for analysis of DNA m5C methylation using Bismark software from BS-Seq data.

# 4 Guitar2 Coordinates - Transcriptomic Landmarks Projected on Genome

The GuitarCoordsFromTxDb object contains the genome-projected transcriptome coordinates, which can be valuable for evaluating transcriptomic information related applications, such as checking the quality of MeRIP-Seq data. The Guitar2 coordinates are essentially the genomic projection of standardized transcript-based coordinates, making a viable bridge beween the landmarks on transcript and genome-based coordinates. The referred "transcript id", standardized position, the interval between two adjacent check points on that components, component name and type are assessible with mcols function.

## > head(gc\_txdb)

GRanges object with 6 ranges and 5 metadata columns:

	seqnames		ranges	strand	1
	<rle></rle>	<ii< td=""><td>Ranges&gt;</td><td><rle></rle></td><td>1</td></ii<>	Ranges>	<rle></rle>	1
uc010nxq.1	chr1 [	10854,	10904]	+	1
uc009vjk.2	chr1 [	321017,	321067]	+	1
uc001aau.3	chr1 [	322872,	322922]	+	1
uc021oeh.1	chr1 [	323268, 3	323318]	+	1
uc021oei.1	chr1 [	326526, 3	326576]	+	1
uc001acv.3	chr1 [	1071377, 10	071427]	+	1
	txid	pos	interv	<i>r</i> al	comp
	<factor></factor>	<numeric></numeric>	<numeri< td=""><td>ic&gt; <fa< td=""><td>ctor&gt;</td></fa<></td></numeri<>	ic> <fa< td=""><td>ctor&gt;</td></fa<>	ctor>
uc010nxq.1	uc010nxq.1	0.005		10	Front
uc009vjk.2	uc009vjk.2	0.005		10	Front
uc001aau.3	uc001aau.3	0.005		10	Front
uc021oeh.1	uc021oeh.1	0.005		10	Front
uc021oei.1	uc021oei.1	0.005		10	Front
uc001acv.3	uc001acv.3	0.005		10	Front
	category				
	<factor></factor>				
uc010nxq.1	mRNA				
uc009vjk.2	mRNA				
uc001aau.3					
uc021oei.1	mRNA				
uc001acv.3	mRNA				

seqinfo: 1 sequence from an unspecified genome; no seqlengths

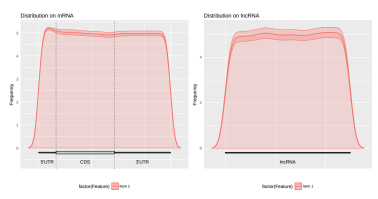
# > mcols(gc\_txdb)

DataFrame with 279900 rows and 5 columns

	txid	pos	interval	comp	category
	<factor></factor>	<numeric></numeric>	<numeric></numeric>	<factor></factor>	<factor></factor>
1	uc010nxq.1	0.005	10	Front	mRNA
2	uc009vjk.2	0.005	10	Front	mRNA
3	uc001aau.3	0.005	10	Front	mRNA
4	uc021oeh.1	0.005	10	Front	mRNA
5	uc021oei.1	0.005	10	Front	mRNA
279896	uc001byp.3	0.995	10	Back	lncRNA
279897	uc001byq.3	0.995	10	Back	lncRNA
279898	uc010ohw.2	0.995	10	Back	lncRNA
279899	uc021olm.1	0.995	10	Back	lncRNA
279900	uc010oid.3	0.995	10	Back	lncRNA

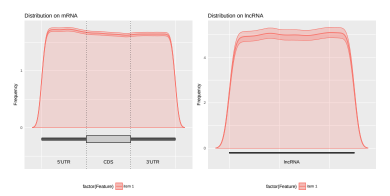
# 5 Check the Overlapping between Different Components

We can also check the distribution of the Guitar2 coordinates built.



or, check without rescaling

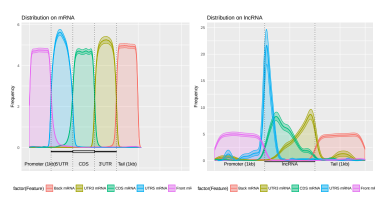
- [1] "Using provided Guitar Coordinates"
- [1] "resolving ambiguious features ..."
- [1] "no figure saved ..."



Alternatively, we can extract the RNA components

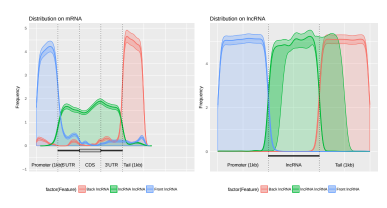
```
> GuitarCoords <- gc_txdb
> type <- paste(mcols(GuitarCoords)$comp,mcols(GuitarCoords)$category)
> key <- unique(type)
> landmark <- list(1,2,3,4,5,6,7,8)
> names(landmark) <- key
> for (i in 1:length(key)) {
+ landmark[[i]] <- GuitarCoords[type==key[i]]
+ }</pre>
```

Check the distribution of mRNA components in the transcriptome



Check the distribution of lncRNA components in the transcriptome

- > Guitar2Plot(gfeatures=landmark[6:8],
- + GuitarCoordsFromTxDb = gc\_txdb,
- + includeNeighborDNA =TRUE,
- + rescaleComponent=FALSE)
- [1] "Using provided Guitar Coordinates"
- [1] "resolving ambiguious features ..."
- [1] "no figure saved ..."



## 6 Session Information

### > sessionInfo()

R version 3.4.1 (2017-06-30)

Platform: x86\_64-pc-linux-gnu (64-bit) Running under: Ubuntu 16.04.3 LTS

Matrix products: default

BLAS: /usr/lib/libblas/libblas.so.3.6.0 LAPACK: /usr/lib/lapack/liblapack.so.3.6.0

#### locale:

[1]	LC_CTYPE=en_US.UTF-8	LC_NUMERIC=C
[3]	LC_TIME=zh_CN.UTF-8	LC_COLLATE=C

[5] LC\_MONETARY=zh\_CN.UTF-8 LC\_MESSAGES=en\_US.UTF-8

[7] LC\_PAPER=zh\_CN.UTF-8 LC\_NAME=C
[9] LC\_ADDRESS=C LC\_TELEPHONE=C
[11] LC\_MEASUREMENT=zh\_CN.UTF-8 LC\_IDENTIFICATION=C

### attached base packages:

[1] grid parallel stats4 stats graphics [6] grDevices utils datasets methods base

## other attached packages:

[1]	Guitar2_0.99.1	ggplot2_2.2.1
[3]	GenomicAlignments_1.6.3	SummarizedExperiment_1.0.2
[5]	rtracklayer_1.30.4	<pre>GenomicFeatures_1.22.13</pre>
[7]	AnnotationDbi_1.32.3	Biobase_2.30.0
[9]	Rsamtools_1.22.0	Biostrings_2.38.4
[11]	XVector_0.10.0	GenomicRanges_1.22.4
Г137	GenomeInfoDb 1.6.3	TRanges 2.4.8

[13] GenomeInfoDb\_1.6.3

## loaded via a name space (and not attached):

Rcpp_0.12.13	plyr_1.8.4
compiler_3.4.1	futile.logger_1.4.3
bitops_1.0-6	<pre>futile.options_1.0.0</pre>
tools_3.4.1	zlibbioc_1.16.0
biomaRt_2.26.1	digest_0.6.12
bit_1.1-12	RSQLite_2.0
memoise_1.1.0	tibble_1.3.4
gtable_0.2.0	pkgconfig_2.0.1
rlang_0.1.2	DBI_0.7
bit64_0.9-7	XML_3.98-1.9
BiocParallel_1.4.3	lambda.r_1.2
blob_1.1.0	scales_0.5.0
colorspace_1.3-2	labeling_0.3
lazyeval_0.2.0	RCurl_1.95-4.8
	compiler_3.4.1 bitops_1.0-6 tools_3.4.1 biomaRt_2.26.1 bit_1.1-12 memoise_1.1.0 gtable_0.2.0 rlang_0.1.2 bit64_0.9-7 BiocParallel_1.4.3 blob_1.1.0 colorspace_1.3-2

[27] lazyeval\_0.2.0 [29] munsell\_0.4.3