

# Package ‘primaryTranscriptAnnotation’

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**Type** Package

**Title** Data-driven gene coordinate inference and assignment of annotations to transcriptional units identified de novo

**Version** 0.1.0

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**Description** This package was developed for two purposes: (1) to generate data-driven transcript annotations based on nascent transcript sequencing data, and (2) to annotate de novo identified transcriptional units (e.g., genes and enhancers) based on existing annotations such as those from (1). The code for this package was employed in analyses early adipogenesis.

**License** NA

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**Imports** bigWig, pracma, dplyr, RColorBrewer, NMF

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adjacent.coords.plot    *Plot adjacent gene pair coordinates*

---

## Description

Plot the coordinates, along with read data, for overlapping genes that were identified as an adjacent pair, with coordinates determined using adjacent.gene.coords().

## Usage

```
adjacent.coords.plot(adjacent.coords = NULL, pair = NULL,
  bw.plus = NULL, bw.minus = NULL, bp.bin = 5, xper = 0.2,
  yper = 0.3)
```

## Arguments

adjacent.coords	bed6 frame with coordinates for adjacent genes
pair	Frame with one row, upstream and downstream genes
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
xper	Fraction of the x-axis before the gene starts
yper	Fraction of the y-axis with open space above the read data

## Value

A plot with read data and gene coordinate annotations.

## Examples

```
# visualize coordinates for a specific pair
adjacent.coords.plot(adjacent.coords=adjacent.coords,
                     pair=fix.genes[4,],
                     bw.plus=bw.plus, bw.minus=bw.minus)
```

---

adjacent.gene.coords    *Function to identify coordinates for adjacent gene pairs*

---

## Description

Adjacent gene pairs can be identified based on manual analyses, despite overlaps in the largest interval coordinates. We empirically define pause sites for these genes by binning the region spanned by both genes, fitting smooth spline curves to the binned read counts, and identifying the two largest peaks separated by a specified distance. We set a bin size and apply the constraint that the identified pause peaks must be a given distance apart. For the spline fits, we set the number of knots to the number of bins divided by specified setting. We identify the TSSs as the exon 1 coordinates closest to the pause site peaks.

## Usage

```
adjacent.gene.coords(fix.genes = NULL, bed.long = NULL, exon1 = NULL,
                    bw.plus = NULL, bw.minus = NULL, bp.bin = NULL, shift.up = NULL,
                    dist.from.start = 50, delta.tss = NULL, knot.div = 4,
                    knot.thresh = 5, diff.tss = 1000, pause.bp = 120,
                    fname = "adjacentSplines.pdf")
```

## Arguments

<code>fix.genes</code>	A frame with upstream and downstream genes
<code>bed.long</code>	Long gene annotations, see <code>get.largest.interval()</code>
<code>exon1</code>	A bed6 frame with first exon gene annotations
<code>bw.plus</code>	Plus strand bigWig data
<code>bw.minus</code>	Minus strand bigWig data
<code>bp.bin</code>	The interval will be separated into adjacent bins of this size
<code>shift.up</code>	Look upstream of the long gene start by this amount to search for a viable TSS
<code>dist.from.start</code>	Distance between the TTS of the upstream gene and the TSS of the downstream gene
<code>delta.tss</code>	Amount by which to shift the identified TSS upstream of the identified interval
<code>knot.div</code>	The number of knots for the spline fit is defined as the number of bins covering the gene end divided by this number
<code>knot.thresh</code>	Minimum number of knots

diff.tss	Minimum distance between TSSs
pause.bp	Number of bp by which a pause site should be considered downstream of a TSS
fname	File name (.pdf) for output plots

### Value

A frame with the adjusted coordinates for the input gene set, along with a plot. The titles for the plots indicate the upstream gene, the downstream gene, and the strand. For genes on the plus strand, the upstream gene is on the left. For the minus strand, the upstream gene is on the right. The red line denotes the spline fit and the vertical lines indicate the pause peaks. The plot is intended for diagnostic purposes.

### Examples

```
# get the start sites for adjacent gene pairs
fix.genes = rbind(fix.genes.id, fix.genes.ov)
bp.bin = 5
knot.div = 40
shift.up = 100
delta.tss = 50
diff.tss = 1000
dist.from.start = 50
adjacent.coords = adjacent.gene.coords(fix.genes=fix.genes, bed.long=TSS.gene,
                                       exon1=gencode.firstExon,
                                       bw.plus=bw.plus, bw.minus=bw.minus,
                                       knot.div=knot.div, bp.bin=bp.bin,
                                       shift.up=shift.up, delta.tss=delta.tss,
                                       dist.from.start=dist.from.start,
                                       diff.tss=diff.tss, fname="adjacentSplines.pdf")
```

---

apply.TSS.coords	<i>Apply previously identified TSSs to an existing gene annotation</i>
------------------	--

---

### Description

This function will apply transcription start site (TSS) coordinates to a bed6 frame. This function is embedded within get.TSS().

### Usage

```
apply.TSS.coords(bed = NULL, tss = NULL)
```

### Arguments

bed	A bed6 frame with gene annotations
tss	TSS estimates, from inside get.TSS(). Note that the TSS genes must be a subset of the genes in the bed6 data.

## Value

A bed6 frame with estimated TSSs incorporated

## Examples

```
out = apply.TSS.coords(bed=bed.out, tss=tss)
```

---

assess.final.coords      *Function to assessing the final identified coordinates*

---

## Description

This function compares the final coordinates to coordinates to inferred coordinates, `tu.gene.overlaps()` versus `get.TTS()`.

## Usage

```
assess.final.coords(bed1 = NULL, bed2 = NULL)
```

## Arguments

bed1	A reference bed frame from <code>get TTS()</code>
bed2	A final bed frame from <code>tu.gene.overlaps()</code>

## Value

A list of metrics: `dtss` is a vector of tss differences, `dtts` is a vector of tts differences, `dtss.mean` is the average tss differences, `dtss.uneq` is the number of tss's that are not identical `dtts.mean` is the mean for tts differences, and `dtts.uneq` is the number of non-identical tts's

## Examples

```
# evaluate the final annotation
metrics = assess.final.coords(bed1=bed.tss.tts, bed2=res)
metrics$dtss.uneq
metrics$dtts.uneq
```

---

chrom.size.filter	<i>Filter identified gene coordinates for chromosome sizes.</i>
-------------------	---

---

### Description

First acquire a chrom.sizes file (e.g., <http://hgdownload.cse.ucsc.edu/goldenPath/mm10/bigZips/mm10.chrom.sizes>; see vignette). If any ends are greater than the species chromosome size, reduce the end to the size limit. Any negative starts will be set to 0.

### Usage

```
chrom.size.filter(coords = NULL, chrom.sizes = NULL)
```

### Arguments

coords	A bed6 frame with inferred coordinates (from get.TTS())
chrom.sizes	A two column frame with chromosomes and their respective sizes in bp

### Value

A list with a bed6 frame with corrected coordinates (bed) and a log documenting which corrections, if any, were made (log).

### Examples

```
coords.filt = chrom.size.filter(coords=coords, chrom.sizes=chrom.sizes)
head(coords.filt$bed)
coords.filt$log
```

---

eval.tss	<i>Evaluate TSS inference performance</i>
----------	---

---

### Description

This function was designed to evaluate the results of our TSS identification analysis. The user should input inferred and largest interval coordinates (bed1 and bed2, respectively). We specify a region centered on the TSS. We obtain read counts in bins that span this window. We sort the genes based on the bin with the maximal reads and we scale the data to the interval (0,1) for visualization. We compute the distances between the TSS and the bin with the max reads within the specified window. Note that this function calls TSS.count.dist() for the TSS distance analysis.

### Usage

```
eval.tss.bed1 = NULL, bed2 = NULL, bw.plus = NULL, bw.minus = NULL,
  window = NULL, bp.bin = NULL, fname = NULL, bin.thresh = 10)
```

**Arguments**

bed1	A bed frame with inferred TSSs
bed2	A bed frame with reference TSSs
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
window	Region size, centered on the TSS for analysis
bp.bin	The interval will be separated into adjacent bins of this size
fname	A .pdf file name for output plots (if NULL, no plot)
bin.thresh	Minimal value of reads in the bin with the max count, below this value the transcript will be removed

**Value**

A list of two lists and a plot (optional, specify fname for the plot to be printed to file). Each list contains three vector elements: raw, scaled, and dist. \$tss.dists.inf cooresponds to the TSS data (bed1) and \$tss.dists.lng cooresponds to the largest interval data (bed2). All outputs are organized based on TSS position (left-upstream). raw = binned raw counts. scaled = binned scaled counts (0,1), with genes sorted based on the distance between the TSS and the max read interval. dist = upper bound distances (  $\min(|\text{dists}|) = \text{bp.bin}$  ). See examples and vignette for more details. The first row of plots includes distributions of distances between the TSS and the maximal reads for the inferred TSSs (left) and largest interval TSSs (right). The second row of plots shows the relationship between the distances from TSSs to max reads and the read depth in the region of read count evaluation (left inferred, right largest interval). The heatmaps show read profiles (horizontal axis, distances centered in the middle of the window; vertical axis, distinct genes). The genes are sorted based on the inferred TSSs to the max reads (left), and the largest interval distances (right) are organized based on the gene order for the inferred TSSs.

**Examples**

```
#
```

---

gene.end.plot	<i>Plot the results of transcription termination site identification</i>
---------------	--

---

**Description**

This function can be used to evaluate the performance of get.TTS(). We also recommend visualizing the results of data-driven gene annotations using a genome browser.

**Usage**

```
gene.end.plot(
  bed = NULL, gene = NULL, bw.plus = NULL,
  bw.minus = NULL, bp.bin = NULL, pk.thresh = 0.1, knot.div = 40,
  knot.thresh = 5, cnt.thresh = 5, add.to.end = 50000,
  tau.dist = 10000, frac.max = 1, frac.min = 0.2)
```

**Arguments**

bed	Bed6 gene coordinates with gene end intervals for estimation of the TTS
gene	A specific gene for plotting
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
bp.bin	The interval at the gene end will be separated into adjacent bins of this size
pk.thresh	The TTS is defined as pk.thresh percent of max peak of the spline fit
knot.div	The number of knots for the spline fit is defined as the number of bins covering the gene end divided by this number; increasing this parameter results and a smoother curve
knot.thresh	Minimum number of knots, if knot.div gives a lower number, use this
cnt.thresh	Read count number below which the TTS is not evaluated
add.to.end	The maximal length of the search region (bp)
tau.dist	Distance constant for the exponential defining the region for peak detection
frac.max	Maximal fraction of gene end region for peak detection
frac.min	Minimal fraction of gene end region for peak detection

**Value**

A single plot is generated with the binned reads (circles), a fitted curve (red), and the location of the identified TTS (vertical line).

**Examples**

```
gene.end.plot(bed=bed.for.tts.eval, gene="Pparg",
              bw.plus=bw.plus, bw.minus=bw.minus,
              bp.bin=bp.bin, add.to.end=add.to.end, knot.div=knot.div,
              pk.thresh=pk.thresh, knot.thresh=knot.thresh,
              cnt.thresh=cnt.thresh, tau.dist=tau.dist,
              frac.max=frac.max, frac.min=frac.min)
```

---

gene.overlaps

*Documentation of gene overlaps*

---

**Description**

This function identifies gene overlaps.

**Usage**

```
gene.overlaps(bed = NULL)
```

**Arguments**

bed	A bed6 frame with processed gene annotations
-----	--



**Value**

A list with `has.start.inside`, `is.a.start.inside`, and `cases`. The object `has.start.inside` is a `bed6` frame that documents genes in which there are starts of other genes. The object `is.a.start.inside` is a `bed6` frame that documents genes that start inside other genes. The object `cases` documents loci in which 2 or more genes overlap (col 7 of a `bed` format frame).

**Examples**

```
# run overlap analysis
overlap.data = gene.overlaps( bed = TSS.gene.filtered1 )
has.start.inside = overlap.data$has.start.inside
is.a.start.inside = overlap.data$is.a.start.inside
head(has.start.inside)
head(is.a.start.inside)
head(overlap.data$cases)
length(unique(overlap.data$cases$gene))
```

---

get.dups

---

*Get duplicates in which start sites match or end sites match*


---

**Description**

This function identifies an extreme form of gene overlaps in which multiple identifiers correspond to identical start or end coordinates.

**Usage**

```
get.dups(bed = NULL)
```

**Arguments**

`bed`                      A `bed` frame

**Value**

A `bed` frame with duplicate entries. Col 7 indicates ordered overlap loci.

**Examples**

```
dups0 = get.dups(bed = TSS.gene)
```

---

get.end.intervals	<i>Get intervals at the gene ends for transcription termination site (TTS) estimation</i>
-------------------	---

---

## Description

We define regions at the gene ends to examine read counts for TTS identification. Note that transcription frequently extends beyond the poly-A site of a gene. To capture the end of transcription, it is critical to examine regions beyond annotated gene boundaries. We evaluate evidence of transcriptional termination in regions extending from a 3' subset of the gene to a selected number of base pairs downstream of the most distal annotated gene end. We initiate the search region with the end of the largest gene annotation (see `get.largest.interval()`). We extend the search region up to a given distance past the annotated gene end. We also apply the constraint that a TTS cannot be identified closer than a specified distance to the previously identified TSS of a downstream gene. This analysis also incorporates the constraint that a gene end region identified cannot cross the TSS of a downstream gene, thereby preventing gene overlaps on a given strand. Thus, we clip the amount of bases added on to the gene end as necessary to avoid overlaps. The output of this function is the input for `get.TTS()`.

## Usage

```
get.end.intervals(bed = NULL, add.to.end = NULL, fraction.end = NULL,
  dist.from.start = NULL)
```

## Arguments

bed	Processed bed6 frame with one entry per gene and identified TSSs
add.to.end	Number of bases to add to the end of each gene to define the search region
fraction.end	Fraction of the gene annotation to consider at the end of the gene
dist.from.start	The maximal allowable distance between the end of one gene and the start of the next

## Value

A bed6 frame for TTS evaluation, see `get.TTS()`.

## Examples

```
# get intervals for TTS evaluation
add.to.end = 100000
fraction.end = 0.2
dist.from.start = 50
bed.for.tts.eval = get.end.intervals(bed=bed.long.filtered4.tss,
  add.to.end=add.to.end,
  fraction.end=fraction.end,
  dist.from.start=dist.from.start)
```

get.gene.end

*Estimate the transcription termination sites (TTSs)***Description**

This function estimates the TTSs. This function is called by get.TTS() and is not designed to be run on its own. TTSs that cannot be estimated are assigned zeros.

**Usage**

```
get.gene.end(bed = NULL, bw = NULL, bp.bin = NULL,
             pk.thresh = NULL, knot.div = NULL, cnt.thresh = NULL,
             knot.thresh = NULL, add.to.end = NULL, tau.dist = NULL,
             frac.max = NULL, frac.min = NULL, sum.thresh = NULL)
```

**Arguments**

bed	A bed6 gene coordinates with gene end intervals for estimation of the TTS
bw	Plus or minus strand bigWig data
bp.bin	The interval at the gene end will be separated into adjacent bins of this size
pk.thresh	The TTS is defined as pk.thresh percent of max peak of the spline fit (identified within the peak search region)
knot.div	The number of knots for the spline fit is defined as the number of bins covering the gene end divided by this number
cnt.thresh	Read count number below which the TTS is not evaluated
knot.thresh	Minimum number of knots, if knot.div gives a lower number, use this
add.to.end	The maximal length of the search region (bp)
tau.dist	Distance constant for the exponential function defining the region for peak detection
frac.max	Maximal fraction of gene end region for peak detection
frac.min	Minimal fraction of gene end region for peak detection
sum.thresh	Threshold for the min sum of read counts

**Value**

A vector of TTS coordinates

**Examples**

See get.TTS()

---

get.largest.interval	<i>Get the largest interval for each gene, given multiple TSS and TTS annotations</i>
----------------------	---

---

### Description

The input bed6 file can be derived from a gencode annotation file, as described in the vignette

### Usage

```
get.largest.interval(bed = NULL)
```

### Arguments

bed	A bed6 frame with comprehensive gene annotations, defaults to NULL
-----	--

### Value

A bed6 frame with the largest annotation for each gene

### Examples

```
# get intervals for furthest TSS and TTS +/- interval
bed.long = get.largest.interval(bed=dat0)
```

---

get.TSS	<i>Select an optimal transcription start site (TSS) for each gene</i>
---------	---

---

### Description

We set a range around each annotated exon 1 within which to search for a read density. For each gene, we select the upstream coordinate of the first exon with the greatest read density, in the specified region (bp.range), out of all first exons annotated for the given gene. Note that input and output genes must be matched (bed.in and bed.out).

### Usage

```
get.TSS(bed.in = NULL, bed.out = NULL, bw.plus = NULL,
        bw.minus = NULL, bp.range = NULL, cnt.thresh = NULL,
        low.read = FALSE, by = "cnt")
```

**Arguments**

bed.in	A bed6 frame with first exon gene annotations
bed.out	A bed6 frame for output (same genes as bed.in)
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
bp.range	The range in bp for identifying the exon 1 with the greatest density (e.g., c(20,120))
cnt.thresh	Read count number below which the TSS is not evaluated, by default the gene is removed from the analysis for max(counts) < cnt.thresh (low.read=FALSE). Indicate low.read=TRUE to use the upstream-most TSS for max(counts) < cnt.thresh.
low.read	Logical for whether to pick the upstream-most TSS if max(counts) < cnt.thresh (default FALSE)
by	indicate whether to select the TSS based on max read count (by="cnt", default) or density (cnt="den")

**Value**

A list with a bed6 frame with inferred TSSs (bed) and a list of problematic genes (issues). Problems documented include start > end and start < 0. For start > end, the original input coordinates are retained (bed.out). For start < 0, start = 0 is applied.

**Examples**

```
# select the TSS for each gene and incorporate these TSSs into the largest interval coordinates
bp.range = c(20,120)
cnt.thresh = 5
bed.out = largest.interval.expr.bed
bed.in = gencode.firstExon[gencode.firstExon$gene %in% bed.out$gene,]
TSS.gene = get.TSS(bed.in=bed.in, bed.out=bed.out,
                  bw.plus=bw.plus, bw.minus=bw.minus,
                  bp.range=bp.range, cnt.thresh=cnt.thresh)
```

get.TTS

*Defining gene ends***Description**

Given regions within which to search for TSSs, we operationally define the TSSs by binning the TTS search regions, counting reads within the bins, fitting smooth spline curves to the bin counts, and detecting points at which the curves peak and then decay towards zero. We applied the constraint that there must be a specified number of bases in the TTS search region, otherwise the TTS analysis is not applied. Similarly, if the number of knots identified is too low, then we set the number of knots to a specified threshold. For this analysis, we set a peak search region at the beginning of the TTS search region and identify the maximal peak from the spline fit. Then we identify the point at which the spline fit decays to a threshold relative to the peak. We reasoned that the peak search region should be largest, as a fraction of the TTS search region, for genes with the

greatest numbers of clipped bases. Such cases occur when the conventional gene ends are proximal to identified TSSs, and we should include these entire regions for analysis of the TTS. Similarly, we reasoned that for genes with substantially less clipped bases, and correspondingly larger TTS search regions with greater potential for observing enhancers or divergent transcripts, the peak search regions should be smaller fractions of the TTS search regions. We use an exponential model to define the peak search regions. The user should use the output of `get.end.intervals()` as an input to this function. Note that gene ends will be set to zeros if there are no counts or if the interval is too small. The gene ends in `tss` will be the default if there are problems with estimating the TTS. This function calls `get.gene.end()`.

## Usage

```
get.TTS(bed = NULL, tss = NULL, bw.plus = NULL, bw.minus = NULL,
        bp.bin = NULL, pk.thresh = 0.1, knot.div = 40, knot.thresh = 5,
        cnt.thresh = 5, add.to.end = 50000, sum.thresh = 20,
        tau.dist = 10000, frac.max = 1, frac.min = 0.2)
```

## Arguments

<code>bed</code>	A <code>bed6</code> gene coordinates with gene end intervals for estimation of the TTS
<code>tss</code>	A <code>bed6</code> gene coordinates containing estimated TSSs and largest interval gene ends
<code>bw.plus</code>	Plus strand bigWig data
<code>bw.minus</code>	Minus strand bigWig data
<code>bp.bin</code>	The interval at the gene end will be separated into adjacent bins of this size
<code>pk.thresh</code>	The TTS is defined as <code>pk.thresh</code> percent of max peak of the spline fit
<code>knot.div</code>	the number of knots for the spline fit is defined as the number of bins covering the gene end divided by this number; increasing this parameter results and a smoother curve
<code>knot.thresh</code>	Minimum number of knots, if <code>knot.div</code> gives a lower number, use this
<code>cnt.thresh</code>	Read count number below which the TTS is not evaluated
<code>add.to.end</code>	The maximal length of the search region (bp)
<code>sum.thresh</code>	Threshold for the min sum of read counts, below this default to the input TTS (default 20)
<code>tau.dist</code>	Distance constant for the exponential defining the region for peak detection
<code>frac.max</code>	Maximal fraction of gene end region for peak detection
<code>frac.min</code>	Minimal fraction of gene end region for peak detection

## Value

A `bed6` file with TTS estimates incorporated, original TTSs are present if the TTS could not be estimated. The output is a list including the `bed` frame along with several metrics (see example below and vignette).

## Examples

```
# identify gene ends
add.to.end = max(bed.for.tts.eval$xy)
knot.div = 40
pk.thresh = 0.05
bp.bin = 50
knot.thresh = 5
cnt.thresh = 5
tau.dist = 10000
frac.max = 1
frac.min = 0.2
inferred.coords = get.TTS(bed=bed.for.tts.eval, tss=TSS.gene.filtered3,
                          bw.plus=bw.plus, bw.minus=bw.minus,
                          bp.bin=bp.bin, add.to.end=add.to.end,
                          knot.div=knot.div,
                          pk.thresh=pk.thresh, knot.thresh=knot.thresh,
                          cnt.thresh=cnt.thresh, tau.dist=tau.dist,
                          frac.max=frac.max, frac.min=frac.min)
```

---

get.tu.gene.coords	<i>Integrate inferred gene coordinates with transcriptional units</i>
--------------------	---

---

## Description

Here we annotate all of the regions in the TU frame based on overlaps with genes. The analysis details are handled by `single.overlaps()` and `multi.overlaps()`. See the vignette for formatting the inputs to this function.

## Usage

```
get.tu.gene.coords(hmm.ann.overlap = NULL, gene.frame = NULL,
                  tss.thresh = NULL, delta.tss = NULL, delta.tts = NULL)
```

## Arguments

<code>hmm.ann.overlap</code>	A frame with coordinates from intersecting tus with inferred gene annotations
<code>gene.frame</code>	A bed6 frame with the full set of inferred annotations, the input to the bedtools overlap analysis
<code>tss.thresh</code>	Number of bp a TU beginning can be off from an annotation in order to be assigned that annotation
<code>delta.tss</code>	Max distance between an upstream gene end and downstream gene start
<code>delta.tts</code>	Max distance between and annotated gene end and the start of a downstream gene before an intermediate TU id is assigned

**Value**

A bed frame with gene/tu coordinates.

**Examples**

```
tss.thresh = 200
delta.tss = 50
delta.tts = 1000
res = get.tu.gene.coords(hmm.ann.overlap=hmm.ann.overlap,
                        tss.thresh=tss.thresh,
                        delta.tss=delta.tss,
                        delta.tts=delta.tts)
```

---

inside.starts

*Identify genes with multiple starts inside*


---

**Description**

This function identifies genes within which >1 start is observed. That is, relatively 'large' genes in which 2 or more genes originate. Data generated from gene.overlaps() serve as an input to this function.

**Usage**

```
inside.starts(vec = NULL)
```

**Arguments**

vec                      Input the xy column from the \$is.a.start.inside output of gene.overlaps()

**Value**

A vector of genes within which multiple starts are observed.

**Examples**

```
overlap.data = gene.overlaps( bed = TSS.gene.filtered1 )
is.a.start.inside = overlap.data$is.a.start.inside
mult.inside.starts = inside.starts(vec = is.a.start.inside$xy)
```



---

`multi.overlap.assign`     *Function to assign identifiers for class5 overlap TUs*

---

### Description

This function is called inside of `multiple.overlaps()` and is not recommended to be used on its own.

### Usage

```
multi.overlap.assign(fr = NULL, tss.thresh = NULL, delta.tss = NULL,
  delta.tts = NULL, cl = NULL)
```

### Arguments

<code>fr</code>	A frame with full bedtools overlap data for a class5 TU with multiple internal annotations
<code>tss.thresh</code>	Number of bp a TU beginning can be off from an annotation in order to be assigned that annotation
<code>delta.tss</code>	Max distance between an upstream gene end and downstream gene start <code>cl</code> = multi-overlap class
<code>delta.tts</code>	Max difference distance between and annotated gene end and the start of a downstream gene before an intermediate TU id is assigned
<code>cl</code>	Multi-overlap class

### Value

A bed data with `chr`, `start`, `end`, `id`, `class`, and `strand`. Note that `id` = 0 for regions of large TUs that do no match input annotations.

### Examples

See `multiple.overlaps()`

---

`multi.overlaps`     *Assign identifiers to TUs with multiple gene overlaps*

---

### Description

This function will assign identifiers to TUs that overlapped with multiple genes. Trusted gene annotations are considered in terms of their degree of overlap with TUs identified in an unbiased manner. Marginal regions of TUs outside of gene overlaps are given generic identifiers.

### Usage

```
multi.overlaps(overlaps = NULL, tss.thresh = NULL, delta.tss = NULL,
  delta.tts = NULL)
```

**Arguments**

overlaps	A frame with bed intersect of inferred TUs (col1-6) with annotated TUs (col7-12) and overlapping bps (col14)
tss.thresh	Number of bp a TU beginning can be off from an annotation in order to be assigned that annotation
delta.tss	Max distance between an upstream gene end and downstream gene start
delta.tts	Max difference distance between and annotated gene end and the start of a downstream gene before an intermediate TU id is assigned

**Value**

A list with bed data (bed) and counts for each class (cnt5-8). bed = data with chr, start, end, id, class, and strand. cnt5 = count of TU from class 5 overlaps

**Examples**

```
dup.hmm.rows = overlap.tu[duplicated(overlap.tu[,c(1:3,6)]) |
  duplicated(overlap.tu[,c(1:3,6)], fromLast=TRUE),]
dup.full = dup.hmm.rows
names(dup.full)[1:6] = c("infr.chr", "infr.start", "infr.end",
  "infr.gene", "infr.xy", "infr.strand")
nrow(dup.full[,1:3] %>% unique)
nrow(dup.full %>% select(ann.gene) %>% unique)
tss.thresh = 200
delta.tss = 50
delta.tts = 1000
class5678 = multi.overlaps(overlaps=dup.full, tss.thresh=tss.thresh,
  delta.tss=delta.tss, delta.tts=delta.tts)
new.ann.mult = class5678$bed
```

---

read.count.body	<i>Count gene body reads and compute density</i>
-----------------	--

---

**Description**

Input coordinates for evaluation of counts and densities. This function works with input that includes one annotation per gene.

**Usage**

```
read.count.body(bed = NULL, bw.plus = NULL, bw.minus = NULL,
  bp.start = 0)
```

**Arguments**

bed	A bed6 frame
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
bp.start	Optional number of bases to omit from the upstream gene boundary

**Value**

A list with counts and desity. counts = vector with end of gene counts for each gene. density = vector with end of gene densities for each gene.

**Examples**

```
# get read counts and densities at the body of each annotated gene

body.reads = read.count.body(bed=largest.interval.bed, bw.plus=bw.plus,
bw.minus=bw.minus, bp.start=bp.start)
hist(log(body.reads$density), breaks=200, col="black",xlab="log read density",main="")
hist(log(body.reads$counts), breaks=200, col="black",xlab="log read count",main="")
```

---

read.count.end	<i>Select a regions containing the gene ends</i>
----------------	--

---

**Description**

Select a fraction of the annotated gene end and consider an additional number of base pairs beyond the gene end within which to count reads. This function works with input that includes one annotation per gene.

**Usage**

```
read.count.end(bed = NULL, bw.plus = NULL, bw.minus = NULL,
fraction.end = NULL, add.to.end = NULL)
```

**Arguments**

bed	A bed6 frame
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
fraction.end	Fraction of the annotated gene end (0,1)
add.to.end	Number of base pairs beyond the gene end within which to count reads

**Value**

A list with counts and desity. counts = vector with end of gene counts for each gene. density = vector with end of gene densities for each gene.

**Examples**

```
# get read counts and densities at the end of each annotated gene
fraction.end = 0.1
add.to.end = 0
end.reads = read.count.end(bed=bed.long, bw.plus=bw.plus, bw.minus=bw.minus,
                           fraction.end=fraction.end, add.to.end=add.to.end)
hist(log(end.reads$density))
hist(log(end.reads$counts))
```

---

`read.count.transcript` *Evaluate reads and compute densities for each transcript of each gene.*

---

### Description

Input coordinates for evaluation of counts and densities. Return the maximal values across all transcripts of a given gene. For `by="cnt"` (default), we return the density associated with the max count for a specific gene. For `by="den"`, we return the density associated with the max count for a specific gene.

### Usage

```
read.count.transcript(bed = NULL, bw.plus = NULL, bw.minus = NULL,
  by = "cnt")
```

### Arguments

<code>bed</code>	A bed6 frame
<code>bw.plus</code>	Plus strand bigWig data
<code>bw.minus</code>	Minus strand bigWig data
<code>by</code>	Either "cnt" (default) or "den" for selecting transcripts based on either count or density

### Value

A list with counts and density. `counts` = vector with end of gene counts for each gene. `density` = vector with end of gene densities for each gene.

### Examples

```
# get read counts and densities for the max transcript of each annotated gene
transcript.reads = read.count.transcript(bed=gencode.transcript, bw.plus=bw.plus, bw.minus=bw.minus)
hist(log(transcript.reads$density), breaks=200, col="black", xlab="log read density", main="")
hist(log(transcript.reads$counts), breaks=200, col="black", xlab="log read count", main="")
```

---

`remove.overlaps` *Filter genes with overlaps*

---

### Description

For a given case in which multiple genes overlap, this function evaluates read density for all transcripts coresponding to each gene. The gene with the largest read count (`by="cnt"`, default) or density (`by="den"`) across all transcripts is retained in the input annotation. The related functions `get.dups()` and `gene.overlaps()` can be used to identify overlap cases.

**Usage**

```
remove.overlaps(bed = NULL, overlaps = NULL, transcripts = NULL,
  bw.plus = NULL, bw.minus = NULL, by = "den")
```

**Arguments**

bed	A bed6 frame with gene annotations
overlaps	Frame with 6 columns in the bed6 format and a 7th column indicating integer overlap cases
by	Indicate whether to select transcripts based on read counts (by="cnt") or densities (by="den",default)

**Value**

A bed6 frame with overlaps removed. That is, a modified version of the input bed.

**Examples**

```
# run overlap analysis
overlap.data = gene.overlaps( bed = bed.long.filtered2.tss )
has.start.inside = overlap.data$has.start.inside
is.a.start.inside = overlap.data$is.a.start.inside
dim(has.start.inside)
dim(is.a.start.inside)
```

---

`single.overlap.assign` *Function to assign identifiers for class1 overlap TUs*

---

**Description**

This function is called inside of `single.overlaps()` and is not recommended to be used on its own.

**Usage**

```
single.overlap.assign(fr = NULL, tss.thresh = NULL, delta.tss = NULL,
  delta.tts = NULL, cl = NULL)
```

**Arguments**

fr	A frame with full bedtools overlap data for a class1
tss.thresh	Number of bp a TU beginning can be off from an annotation in order to be assigned that annotation
delta.tss	Max distance between an upstream gene end and downstream gene start
cl	Overlap class

**Value**

A bed data frame with chr, start, end, id, class, and strand. Note that id = 0 for regions of large TUs that do no match input annotations.

**Examples**

See `single.overlaps()`

---

<code>single.overlaps</code>	<i>Assign identifiers to TUs with single gene overlaps</i>
------------------------------	--

---

**Description**

This function will assign identifiers to TUs that overlapped with single genes. Trusted gene annotations are considered in terms of their degree of overlap with TUs identified in an unbiased manner. Marginal regions of TUs outside of gene overlaps are given generic identifiers.

**Usage**

```
single.overlaps(overlaps = NULL, tss.thresh = NULL, delta.tss = NULL,
  delta.tts = NULL)
```

**Arguments**

<code>overlaps</code>	A frame with bed intersect of inferred TUs (col1-6) with annotated TUs (col7-12) and overlapping bps (col14)
<code>tss.thresh</code>	The number of bp that a TU beginning can be off from an annotation in order to be assigned that annotation
<code>delta.tss</code>	Max distance between an upstream gene end and downstream gene start
<code>delta.tts</code>	Max difference distance between and annotated gene end and the start of a downstream gene before an intermediate TU id is assigned

**Value**

A list with bed data (bed) and counts for each class (cnt1-4). bed = data with chr, start, end, id, class, and strand. cnt1 = count of TU from class 1 overlaps

**Examples**

```
sing.hmm.rows = overlap.tu[!duplicated(overlap.tu[,c(1:3,6)]) &
  !duplicated(overlap.tu[,c(1:3,6)], fromLast=TRUE),]
overlaps = sing.hmm.rows
names(overlaps)[1:6] = c("infr.chr", "infr.start", "infr.end",
  "infr.gene", "infr.xy", "infr.strand")

tss.thresh = 200
delta.tss = 50
delta.tts = 1000
```

```
class1234 = single.overlaps(overlaps=overlaps, tss.thresh=tss.thresh,
                           delta.tss=delta.tss, delta.tts=delta.tts)
new.ann.sing = class1234$bed
```

---

TSS.count.dist	<i>Get distances from identified transcription start sites (TSSs) to the nearest regions with peak reads</i>
----------------	--

---

## Description

This function was designed to evaluate the results of our TSS identification analysis. We specify a region centered on the TSS. We obtain read counts in bins that span this window. We sort the genes based on the bin with the maximal reads and we scale the data to the interval (0,1) for visualization. We compute the distances between the TSS and the bin with the max reads within the specified window.

## Usage

```
TSS.count.dist(bed = NULL, bw.plus = NULL, bw.minus = NULL,
               window = NULL, bp.bin = NULL, bin.thresh = NULL)
```

## Arguments

bed	A bed frame for TSS evaluation
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
window	Region size, centered on the TSS for analysis
bp.bin	The interval will be separated into adjacent bins of this size
bin.thresh	Minimal value of reads in the bin with the max count, below this value the transcript will be removed

## Value

A list with three vector elements: raw, scaled, and dist. All outputs are organized based on TSS position (left-upstream). raw = binned raw counts. scaled = binned scaled counts (0,1). dist = upper bound distances ( min(ldistsl) = bp.bin ). See examples and vignette for more details.

## Examples

```
see eval.tss()
```

---

TTS.boundary.clip	<i>Look at whether TTSs identified at the search boundry were clipped.</i>
-------------------	--

---

### Description

This function considers information in both coords and bed.for.tts.eval (see vignette) to identify the fractions of search boundary TTSs that were clipped and unclipped.

### Usage

```
TTS.boundary.clip(coords = NULL, bed.for.tts.eval = NULL)
```

### Arguments

coords	A bed6 frame with inferred coordinates (from get.TTS())
bed.for.tts.eval	A bed6 frame with TTS search regions (from get.end.intervals())

### Value

A list with two fractions [0,1], frac.clip and frac.noclip, and a vector of clip distances (clip.tts) for genes with boundary TTSs.

### Examples

```
# look at whether TTSs identified at the search boundry had clipped boundaries
frac.bound.clip = TTS.boundary.clip(coords=coords, bed.for.tts.eval=bed.for.tts.eval)
frac.bound.clip$frac.clip
frac.bound.clip$frac.noclip

# plot didtribution of clip distances for genes with boundary TTSs
hist(frac.bound.clip$clip.tts,main="",xlab="bp clipped for boundry genes")
```

---

TTS.boundary.match	<i>Check for the fraction of identified TTSs that match the search boundry</i>
--------------------	--

---

### Description

This function considers information in both coords and bed.for.tts.eval (see vignette) to identify the fraction of TTSs that match the boundary of the search region.

### Usage

```
TTS.boundary.match(coords = NULL, bed.for.tts.eval = NULL)
```



**Arguments**

coords                    A bed6 frame with inferred coordinates (from get.TTS())  
 bed.for.tts.eval                    A bed6 frame with TTS search regions (from get.end.intervals())

**Value**

A fraction [0,1].

**Examples**

```
frac.match = TTS.boundary.match(coords=coords, bed.for.tts.eval=bed.for.tts.eval)
```

---

TTS.count.dist	<i>Get read counts around transcription termination sites (TTSs)</i>
----------------	--

---

**Description**

We set a window around the TTS and segment the window into bins. To sort the read data, we compute cumulative counts of reads, and we sort based on the bin at which a specified percentage of the reads are found. We also take a ratio of gene counts downstream / upstream of the TTS with regions within an interval.

**Usage**

```
TTS.count.dist(bed = NULL, bw.plus = NULL, bw.minus = NULL,  
  window = NULL, bp.bin = NULL, frac.max = NULL,  
  ratio.region = NULL)
```

**Arguments**

bed                    A bed frame for TSS evaluation  
 bw.plus                Plus strand bigWig data  
 bw.minus              Minus strand bigWig data  
 window                Region size, centered on the TTS for analysis  
 bp.bin                The interval will be separated into adjacent bins of this size  
 frac.max              Fraction of cumulative distribution for sorting entries are sorted by indices  $\min(\text{cumsum}(x)/\text{sum}(x)) < \text{frac.max}$   
 ratio.region          Number of bp on either side of the TTS to evaluate the ratio entries

**Value**

A list with three elements: raw, scaled, and ratio. All outputs are organized based on TTS position (left-upstream, see frac.max). raw = binned raw counts. scaled = binned scaled counts (0,1). ratio = downstream(ratio.region) / upstream(ratio.region).

## Examples

```
# look at read distribution around identified TTSs
window = 1000
bp.bin = 10
frac.max = 0.8
ratio.region = 300
tts.dists = TTS.count.dist(bed=coords, bw.plus=bw.plus, bw.minus=bw.minus,
    window=window, bp.bin=bp.bin, frac.max=frac.max,
    ratio.region=ratio.region)
```

---

tts.plot

---

*Plot inferred and largest interval coordinates*


---

## Description

This function plots the inferred and largest interval coordinates along with read data. The plot annotates the peak search region and the total search region.

## Usage

```
tts.plot(coords = NULL, gene.end = NULL, long.gene = NULL,
    bw.plus = NULL, bw.minus = NULL, bp.bin = 5, gene = NULL,
    xper = 0.2, yper = 1.3, add.to.end = NULL, tau.dist = NULL,
    frac.max = NULL, frac.min = NULL)
```

## Arguments

coords	A bed6 frame with inferred coordinates
gene.end	A bed6 frame with peak search regions
long.gene	A bed6 frame with largest interval coordinates
bw.plus	Plus strand bigWig data
bw.minus	Minus strand bigWig data
gene	A gene for plotting
xper	Fraction of the x-axis before the gene starts
yper	Fraction of the y-axis with open space above the read data
add.to.end	The maximal length of the search region (bp)
tau.dist	Distance constant for the exponential defining the region for peak detection
frac.max	Maximal fraction of gene end region for peak detection
frac.min	Minimal fraction of gene end region for peak detection

## Value

A plot with reads, the interred annotation (black), the largest interval annotation (gray), the TTS search region (solid, vertical), and the TTS peak region (dashed, vertical).

**Examples**

```
# first run inferred.coords = get.TTS(...)
coords = inferred.coords$bed
gene.end = bed.for.tts.eval
long.gene = largest.interval.bed
tts.plot(coords=coords, gene.end=gene.end, long.gene=long.gene,
         gene = "Pparg", xper=0.2, yper=0.3,
         frac.min=frac.min, frac.max=frac.max,
         bw.plus=bw.plus, bw.minus=bw.minus, bp.bin=5,
         add.to.end=add.to.end, tau.dist=tau.dist)
```

tu.gene.overlaps

*Assign identifiers to TUs with single gene overlaps***Description**

Summarize gene/tu counts after intersecting the respective annotations. See the vignette for analysis details and instructions.

**Usage**

```
tu.gene.overlaps(hmm.ann.overlap = NULL)
```

**Arguments**

`hmm.ann.overlap`

A frame with coordinates from intersecting tus with inferred gene annotations

**Value**

A list of metrics including the total number of TUs (`n.tus`), the number of TUs without gene overlaps (`n.tu.no.gene.overlap`), the number of TUs with overlapping genes (`n.tu.overlap.gene`), the number of genes with overlapping TUs (`n.gene.overlap.tu`), the number of TUs overlapping single genes (`n.tu.overlap.single.genes`), and the number of TUs with multiple gene overlaps (`n.tu.overlap.multiple.genes`). Note that `n.tu.overlap.gene = n.tu.overlap.single.genes + n.tu.overlap.multiple.genes`.

**Examples**

```
tg.overlaps = tu.gene.overlaps(hmm.ann.overlap=hmm.ann.overlap)
```

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