pybedtools Documentation

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CHAPTER

ONE

OVERVIEW

pybedtools is a Python wrapper for Aaron Quinlan's BEDtools and is designed to leverage the "genome algebra" power of BEDtools from within Python scripts.

This documentation is written assuming you know how to use BEDTools and Python.

See full online documentation, including installation instructions, at http://packages.python.org/pybedtools/.

Created by Ryan Dale 2010

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pybedtools is released under the GPLv2 license; see LICENSE.txt for more info.

Note: pybedtools is still very much in progress. Please keep that in mind when assesing whether to use this package in production code.

The documentation is separated into 4 main parts, depending on the depth you'd like to cover:

- Lazy, or just want to jump in? Check out *Three brief examples* to get a feel for the package.
- Want a guided tour? Give the *Tutorial Contents* a shot.
- More advanced features are described in the *Topical Documentation* section.
- Finally, doctested module documentation can be found in *pybedtools Reference*.

CHAPTER

TWO

CONTENTS:

2.1 Installation

2.1.1 Requirements

First, make sure you have the required packages for installing pybedtools:

- 1. BEDTools. The version is not important, but later versions will have more features so it's a good idea to get the latest. Follow the instructions at https://github.com/arq5x/bedtools to install, and make sure the programs are on your path. That is, you should be able to call intersectBed from any directory
- 2. Python 2.5 or greater (Python 3 support is coming soon)
- 3. Cython, version 0.14.1 or greater
- 4. argparse if you are running Python < 2.7 (Python 2.7 comes with argparse already)

Both argparse and Cython can be installed with pip:

```
pip install cython argparse
or easy_install:
easy_install cython argparse
```

2.1.2 Installing pybedtools

For the latest stable version,

- 1. download the pybedtools source from http://pypi.python.org/pypi/pybedtools
- 2. Unzip the tarball
- 3. In the newly created directory, run:

```
python setup.py install
```

(you may need root privleges to do so)

Note: Due to poor support of Cython and C++ (which is what pybedtools uses), it is not currently possible to "easy_install" pybedtools all in one shot from the command line. As a result, users must download the source and install using the method above.

For the bleeding-edge development version, use the Git repository at http://github.com/daler/pybedtools.

2.1.3 Testing your installation

Quick test

A quick functional test is to create a new script with the following contents:

```
import pybedtools
a = pybedtools.example_bedtool('a.bed')
b = pybedtools.example_bedtool('b.bed')
print a.intersect(b)
```

If this script is called test.py, then running it with python test.py should print out:

```
chr1 155 200 feature2 0 + chr1 155 200 feature3 0 - chr1 900 901 feature4 0 +
```

Running the test suite

For more extensive testing, you can run the full test suite which requires nose and PyYAML to be installed (both easy_install-able). The test suite will re-compile the Cython extensions, run unit tests and doctests. To run the test suite, use:

```
sh test.sh $VERSION
```

where \$VERSION is the version of Python you'd like to run the tests with (e.g., 2.7).

If you have sphinx installed (e.g., via easy_install sphinx), you can run the doctests in this documentation by going to the docs directory and running:

make doctest

2.2 Three brief examples

Here are three examples to show typical usage of pybedtools. More info can be found in the docstrings of pybedtools methods and in the *Tutorial Contents*.

2.2.1 Example 1: Save a BED file of intersections, with track line

This example saves a new BED file of intersections between a . bed and b . bed, adding a track line to the output:

```
>>> import pybedtools
>>> a = pybedtools.BedTool('a.bed')
>>> a.intersect('b.bed').saveas('a-and-b.bed', trackline="track name='a and b' color=128,0,0")
```

2.2.2 Example 2: Intersections for a 3-way Venn diagram

This example gets values for a 3-way Venn diagram of overlaps. This demonstrates operator overloading of BedTool objects:

```
>>> import pybedtools
>>> # set up 3 different bedtools
>>> a = pybedtools.BedTool('a.bed')
>>> b = pybedtools.BedTool('b.bed')
>>> c = pybedtools.BedTool('c.bed')
>>> (a-b-c).count() # unique to a
>>> (a+b-c).count() # in a and b, not c
>>> (a+b+c).count() # common to all
>>> # ... and so on, for all the combinations.
```

For more, see the pybedtools.scripts.venn_mpl and pybedtools.scripts.venn_gchart scripts, which wrap this functionality in command-line scripts to create Venn diagrams using either matplotlib or Google Charts API respectively.

2.2.3 Example 3: Count reads in introns and exons, in parallel

This example shows how to count the number of reads in introns and exons in parallel. It is somewhat more involved, but illustrates several additional features of pybedtools such as:

- BAM file support (for more, see Working with BAM files)
- indexing into Interval objects (for more, see *Intervals*)
- filtering (for more, see *Filtering*)
- streaming (for more, see *Using BedTool objects as iterators/generators*)
- ability to use parallel processing

The first listing has many explanatory comments, and the second listing shows the same code with no comments to give more of a feel for pybedtools.

```
import sys
import multiprocessing
import pybedtools
# get example GFF and BAM filenames
gff = pybedtools.example_filename('gdc.gff')
bam = pybedtools.example_filename('gdc.bam')
# Some GFF files have invalid entries -- like chromosomes with negative coords
# or features of length = 0. This line removes them and saves the result in a
# tempfile
g = pybedtools.BedTool(gff).remove_invalid().saveas()
# Next, we create a function to pass only features for a particular
# featuretype. This is similar to a "grep" operation when applied to every
# feature in a BedTool
def featuretype_filter(feature, featuretype):
   if feature[2] == featuretype:
       return True
    return False
# This function will eventually be run in parallel, applying the filter above
# to several different BedTools simultaneously
```

```
def subset_featuretypes(featuretype):
   return g.filter(featuretype_filter, featuretype).saveas()
# This function performs the intersection of a BAM file with a GFF file and
# returns the total number of hits. It will eventually be run in parallel.
def count_reads_in_features(features):
    Callback function to count reads in features
    # This shows how to use BAM files by using the 'abam' kwarg and explicitly
    # specifying the 'b' kwarg as well.
    # In addition, we use stream=True so that no intermediate tempfile is
    # created, and bed=True so that the .count() method can iterate through the
    # resulting streamed BedTool.
    return features.intersect(abam=bam,
                             b=features.fn,
                             bed=True,
                             stream=True).count()
# Set up a pool of workers for parallel processing
pool = multiprocessing.Pool()
# Create separate files for introns and exons, using the function we defined
# above
featuretypes = ('intron', 'exon')
introns, exons = pool.map(subset_featuretypes, featuretypes)
# Perform some genome algebra to get unique and shared intron/exon regions
exon_only = exons.subtract(introns).merge().remove_invalid().saveas()
intron_only = introns.subtract(exons).merge().remove_invalid().saveas()
intron_and_exon = exons.intersect(introns).merge().remove_invalid().saveas()
# Do intersections with BAM file in parallel, using the other function we
# defined above
features = (exon_only, intron_only, intron_and_exon)
results = pool.map(count_reads_in_features, features)
# Print the results
labels = ('
               exon only:',
              intron only:',
          'intron and exon:')
for label, reads in zip(labels, results):
    sys.stdout.write(' \$s \$s \ n' \$ (label, reads))
# Clean up any tempfiles that were created
pybedtools.cleanup(verbose=False)
Here's the same code but with no comments:
import sys
import multiprocessing
import pybedtools
gff = pybedtools.example_filename('gdc.gff')
```

```
bam = pybedtools.example_filename('gdc.bam')
g = pybedtools.BedTool(gff).remove_invalid().saveas()
def featuretype_filter(feature, featuretype):
   if feature[2] == featuretype:
       return True
    return False
def subset_featuretypes(featuretype):
    return g.filter(featuretype_filter, featuretype).saveas()
def count_reads_in_features(features):
    Callback function to count reads in features
    return features.intersect(abam=bam,
                             b=features.fn,
                             bed=True,
                             stream=True).count()
pool = multiprocessing.Pool()
featuretypes = ('intron', 'exon')
introns, exons = pool.map(subset_featuretypes, featuretypes)
exon_only = exons.subtract(introns).merge().remove_invalid().saveas()
intron_only = introns.subtract(exons).merge().remove_invalid().saveas()
intron_and_exon = exons.intersect(introns).merge().remove_invalid().saveas()
features = (exon_only, intron_only, intron_and_exon)
results = pool.map(count_reads_in_features, features)
labels = ('
              exon only:',
              intron only:',
          'intron and exon:')
for label, reads in zip(labels, results):
    sys.stdout.write(' %s %s \n' % (label, reads))
pybedtools.cleanup(verbose=False)
```

For more on using pybedtools, continue on to the *Tutorial Contents* . . .

2.3 Tutorial Contents

2.3.1 Intro

This tutorial assumes that

- 1. You know how to use BEDTools (if not, check out the BEDTools documentation)
- 2. You know how to use Python (if not, check out some tutorials like Learn Python the Hard Way)

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A brief note on conventions

Throughout this documentation I've tried to use consistent typography, as follows:

- Python variables and arguments, as well as filenames look like this: s=True
- Methods, which are often linked to documentation look like this: BedTool.merge().
- Arguments that are passed to BEDTools programs, as if you were on the command line, look like this: -d.
- The ">>>" in the examples below indicates a Python interpreter prompt and means to type the code into an interactive Python interpreter like IPython or in a script. (don't type the >>>)

Onward!

2.3.2 Create a BedTool

First, follow the *Installation* instructions if you haven't already done so to install both BEDTools and pybedtools.

Then import the pybedtools module and make a new BedTool:

```
>>> import pybedtools
>>> # use a BED file that ships with pybedtools...
>>> a = pybedtools.example_bedtool('a.bed')
>>> # ...or use your own by passing a filename
>>> a = pybedtools.BedTool('peaks.bed')
```

This documentation uses example files that ship with pybedtools. To access these files from their installation location, we use the <code>example_bedtool()</code> function. This is convenient because if you copy-paste the examples, they will work. If you would rather learn using your own files, just pass the filename to a new <code>BedTool</code>, like the above example.

You can use any file that BEDTools supports – this includes BED, VCF, GFF, and gzipped versions of any of these. See *Creating a BedTool* for more on the different ways of creating a BedTool, including from iterators and directly from a string.

Now, let's see how to do a common task performed on BED files: intersections.

2.3.3 Intersections

One common use of BEDTools and pybedtools is to perform intersections.

First, let's create some example BedTool instances:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
```

Then do the intersection with the BedTool.intersect() method:

```
>>> a_and_b = a.intersect(b)
```

a_and_b is a new BedTool instance. It now points to a temp file on disk, which is stored in the attribute a_and_b.fn; this temp file contains the intersection of a and b.

We can either print the new BedTool (which will show ALL features – use with caution if you have huge files!) or use the BedTool.head() method to show up to the first N lines (10 by default). Here's what a, b, and a_and_b look like:

```
>>> a.head()
chr1 1 100 feature1
      100 200 feature2
                           0
chr1
chr1
       150 500 feature3
                           0
chr1
       900 950 feature4
                           0
>>> b.head()
chr1 155 200 feature5
                           0
chr1 800 901 feature6
                           \cap
>>> a_and_b.head()
chr1 155 200 feature2
chr1
       155 200 feature3
       900 901 feature4
chr1
```

The BedTool.intersect() method simply wraps the BEDTools program intersectBed. This means that we can pass BedTool.intersect() any arguments that intersectBed accepts. For example, if we want to use the intersectBed switch -u (which acts as a True/False switch to indicate that we want to see the features in a that overlapped something in b), then we can use the keyword argument u=True, like this:

```
>>> # Intersection using the -u switch
>>> a_with_b = a.intersect(b, u=True)
>>> a_with_b.head()
chr1  100 200 feature2  0 +
chr1  150 500 feature3  0 -
chr1  900 950 feature4  0 +
```

This time, a_with_b is another BedTool object that points to a different temp file whose name is stored in a_with_b.fn. You can read more about the use of temp files in *Principle 1: Temporary files are created automatically*. More on arguments that you can pass to BedTool objects in a moment, but first, some info about saving files.

2.3.4 Saving the results

If you want to save the results as a meaningful filename for later use, use the <code>BedTool.saveas()</code> method. This also lets you optionally specify a trackline for directly uploading to the UCSC Genome Browser, instead of opening up the files afterward and manually adding a trackline:

```
>>> c = a_with_b.saveas('intersection-of-a-and-b.bed', trackline='track name="a and b"')
>>> print c.fn
intersection-of-a-and-b.bed
>>> # opening the underlying file shows the track line
>>> print open(c.fn).read()
track name="a and b"
          1.5.5
                   200
                                            0
                                                    +
chr1
                            feature2
           155
                                            0
                   2.00
chr1
                            feature3
           900
                                            0
chr1
                   901
                            feature4
>>> # printing the BedTool object will not show non-feature lines
>>> print c
                                            \cap
chr1 155
                    2.00
                            feature2
                                                    +
          155
                   200
                                            0
chr1
                            feature3
          900
                   901
                            feature4
                                            0
```

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Note that the BedTool.saveas() method returns a new BedTool object which points to the newly created file on disk. This allows you to insert a BedTool.saveas() call in the middle of a chain of commands (described in another section below).

2.3.5 Default arguments

Recall that we passed the u=True argument to a.intersect():

```
>>> a_with_b = a.intersect(b, u=True)
```

While we're on the subject of arguments, note that we didn't have to specify -a or -b arguments, like you would need if calling intersectBed from the command line. That's because BedTool objects make some assumptions for convenience.

We could have supplied the arguments a=a.fn and b=b.fn:

```
>>> another_way = a.intersect(a=a.fn, b=b.fn, u=True)
>>> another_way == a_with_b
True
```

But since we're calling a method on the BedTool object a, pybedtools assumes that the file a points to (stored in the attribute a.fn) is the one we want to use as input. So by default, we don't need to explicitly give the keyword argument a=a.fn because the a.intersect() method does so automatically.

We're also calling a method that takes a second bed file as input — other such methods include BedTool.subtract() and BedTool.closest(), and others. For these methods, in addition to assuming —a is taken care of by the BedTool.fn attribute, pybedtools also assumes the first unnamed argument to these methods are the second file you want to operate on (and if you pass a BedTool, it'll automatically use the file in the fn attribute of that BedTool).

An example may help to illustrate: these different ways of calling BedTool.intersect() all have the same results, with the first version being the most compact (and probably most convenient):

```
>>> # these all have identical results
>>> x1 = a.intersect(b)
>>> x2 = a.intersect(a=a.fn, b=b.fn)
>>> x3 = a.intersect(b=b.fn)
>>> x4 = a.intersect(b, a=a.fn)
>>> x1 == x2 == x3 == x4
True
```

Note that a .intersect (a=a.fn, b) is not a valid Python expression, since non-keyword arguments must come before keyword arguments, but a .intersect (b, a=a.fn) works fine.

If you're ever unsure, the docstring for these methods indicates which, if any, arguments are used as default. For example, in the BedTool.intersect() help, it says:

OK, enough about arguments for now, but you can read more about them in *Principle 2: Names and arguments are as similar as possible to BEDTools*, *Principle 4: Sensible default args* and *Principal 5: Other arguments have no defaults*.

2.3.6 Chaining methods together (pipe)

One useful thing about BedTool methods is that they often return a new BedTool. In practice, this means that we can chain together multiple method calls all in one line, similar to piping on the command line.

For example, this intersect and merge can be combined into one command:

```
>>> # These two lines...
>>> x1 = a.intersect(b, u=True)
>>> x2 = x1.merge()
>>> # ...can be combined into one line:
>>> x3 = a.intersect(b, u=True).merge()
>>> x2 == x3
True
```

In general, methods that return BedTool objects have the following text in their docstring to indicate this:

```
.. note::

This method returns a new BedTool instance
```

A rule of thumb is that all methods that wrap BEDTools programs return BedTool objects, so you can chain these together. Many pybedtools-unique methods return BedTool objects too, just check the docs (according to *Principle 7: Check the help*). For example, as we saw in one of the examples above, the BedTool.saveas() method returns a BedTool object. That means we can sprinkle those commands within the example above to save the intermediate steps as meaningful filenames for later use. For example:

```
>>> x4 = a.intersect(b, u=True).saveas('a-with-b.bed').merge().saveas('a-with-b-merged.bed')
```

Now we have new files in the current directory called a-with-b.bed and a-with-b-merged.bed. Since BedTool.saveas() returns a BedTool object, x4 points to the a-with-b-merged.bed file.

2.3.7 Operator overloading

There's an even easier way to chain together commands.

I found myself doing intersections so much that I thought it would be useful to overload the + and – operators to do intersections. To illustrate, these two example commands do the same thing:

```
>>> x5 = a.intersect(b, u=True)
>>> x6 = a + b
>>> x5 == x6
True
```

Just as the + operator assumes intersectBed with the -u arg, the - operator assumes intersectBed with the -v arg.

```
>>> x7 = a.intersect(b, v=True)
>>> x8 = a - b
>>> x7 == x8
True
```

If you want to operating on the resulting BedTool that is returned by an addition or subtraction, you'll need to wrap the operation in parentheses. This is another way to do the chaining together of the intersection and merge example from above:

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```
>>> x9 = (a + b).merge()
```

And to double-check that all these methods return the same thing:

```
>>> x2 == x3 == x4 == x9
True
```

You can learn more about chaining in *Principle 6: Chaining together commands*.

2.3.8 Intervals

An Interval object is how pybedtools represents a line in a BED, GFF, GTF, or VCF file in a uniform fashion. This section will describe some useful features of Interval objects.

First, let's get a BedTool to work with:

```
>>> a = pybedtools.example_bedtool('a.bed')
```

We can access the Intervals of a several different ways. The most common use is probably by using the BedTool a as an iterator. For now though, let's look at a single Interval:

```
>>> feature = iter(a).next()
```

Common Interval attributes

Printing a feature converts it into the original line from the file:

All features have chrom, start, stop, name, score, and strand attributes. Note that start and stop are long integers, while everything else (including score) is a string.

```
>>> feature.chrom
'chr1'
>>> feature.start
1L
>>> feature.stop
100L
>>> feature.name
'feature1'
>>> feature.score
'0'
>>> feature.strand
'+'
```

Let's make another feature that only has chrom, start, and stop to see how pybedtools deals with missing attributes:

```
>>> feature2 = iter(pybedtools.BedTool('chrX 500 1000', from_string=True)).next()
>>> print feature2
chrX 500 1000
```

```
>>> feature2.chrom
'chrX'
>>> feature2.start
500L
>>> feature2.stop
1000L
>>> feature2.name
''
>>> feature2.score
''
>>> feature2.strand
```

This illustrates that default values are empty strings.

Indexing into Interval objects

Interval objects can also be indexed by position into the original line (like a list) or indexed by name of attribute (like a dictionary).

Fields

Interval objects have a Interval.fields attribute that contains the original line split into a list of strings. When an integer index is used on the Interval (for example, feature[3]), it is the fields attribute that is actually being indexed into.

```
>>> f = iter(pybedtools.BedTool('chr1 1 100 asdf 0 + a b c d', from_string=True)).next()
>>> f.fields
['chr1', '1', '100', 'asdf', '0', '+', 'a', 'b', 'c', 'd']
>>> len(f.fields)
10
```

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BED is 0-based, others are 1-based

One troublesome part about working with multiple formats is that BED files have a different coordinate system than GFF/GTF/VCF/ files.

BED files are 0-based (the first base of the chromosome is considered position 0) and the **feature does not include** the stop position.

GFF, GTF, and VCF files are 1-based (the first base of the chromosome is considered position 1) and the **feature** includes the stop position.

Note: pybedtools follows the following conventions:

- The value in Interval.start will always contain the 0-based start position, even if it came from a GFF or other 1-based feature.
- Getting the len() of an Interval will always return Interval.stop Interval.start, so no matter what format the original file was in, the length will be correct.
- The contents of Interval.fields will always be strings, which in turn always represent the original line in the file. This means that for a GFF feature, Interval.fields[3] or Interval[3], which is 1-based according to the file format, will always be one bp larger than Interval.start, which always contains the 0-based start position. However, Interval[3] will be a string and Interval.start will be a long.

To illustrate and confirm, let's create a GFF feature and a BED feature from scratch and compare them:

```
>>> # GFF Interval from scratch
>>> gff = ["chr1",
           "fake",
. . .
           "mRNA",
. . .
           "51",
                   # <- start is 1 greater than start for the BED feature below
. . .
           "300",
. . .
           ".",
           "+",
           ".",
. . .
           "ID=mRNA1; Parent=gene1; "]
>>> gff = pybedtools.create_interval_from_list(gff)
>>> print qff
chr1
           fake
                    mRNA
                                     300
                                                                       ID=mRNA1; Parent=gene1;
>>> # BED Interval from scratch
>>> bed = ["chr1",
           "50",
. . .
           "300",
. . .
           "mRNA1",
. . .
           ".",
           "+"]
>>> bed = pybedtools.create_interval_from_list(bed)
>>> print bed
chrl 50
                    300
                            mRNA1
>>> # confirm they are recognized as the right type
>>> gff.file_type
'gff'
>>> bed.file_type
'bed'
>>> # Start attributes should be identical
```

```
>>> bed.start == gff.start == 50
True
>>> bed.start
50L
>>> bed[1]
'50'
>>> # GFF .start is 1 less than the string value stored at index 3
>>> gff.start
50L
>>> gff[3]
'51'
>>> len(bed) == len(gff) == 250
True
```

GFF features have access to attributes

GFF and GTF files have lots of useful information in their attributes field (the last field in each line). These attributes can be accessed with the Interval.attrs attribute, which acts like a dictionary. For speed, the attributes are lazy – they are only parsed when you ask for them. BED files, which do not have an attributes field, will return an empty dictionary.

```
>>> # original feature
>>> print qff
                                     300
chr1
           fake
                 mRNA
                            51
                                                                     ID=mRNA1; Parent=gene1;
>>> # original attributes
>>> qff.attrs
{'ID': 'mRNA1', 'Parent': 'gene1'}
>>> # add some new attributes
>>> gff.attrs['Awesomeness'] = 99
>>> gff.attrs['ID'] = 'transcript1'
>>> # Changes in attributes are propagated to the printable feature
>>> print qff
chr1
           fake
                    mRNA
                            51
                                     300
                                                     +
                                                                     Awesomeness=99; ID=transcript1; Pa
```

Understanding Interval objects is important for using the powerful filtering and mapping facilities of BedTool objects, as described in the next section.

2.3.9 Filtering

The filter() method lets you pass in a function that accepts an Interval as its first argument and returns True for False. This allows you to perform "grep"-like operations on BedTool objects. For example, here's how to get a new BedTool containing features from a that are more than 100 bp long:

The filter() method will pass its *args and **kwargs to the function provided. So here is a more generic case, where the function is defined once and different arguments are passed in for filtering on different lengths:

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```
>>> def len_filter(feature, L):
... "Returns True if feature is longer than L"
... return len(feature) > L
```

Now we can pass different lengths without defining a new function for each length of interest, like this:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> print a.filter(len_filter, L=10)
chrl 1
               100 feature1
                                               +
          100
                 200
                                        0
chr1
                        feature2
          150
                                        0
                 500
chr1
                         feature3
          900
                 950
                                        0
chr1
                         feature4
>>> print a.filter(len_filter, L=99)
chr1 100 200 feature2
                                        0
chr1
          150
                 500
                         feature3
                                        0
>>> print a.filter(len_filter, L=200)
                 500
          150
```

See *Using BedTool objects as iterators/generators* for more advanced and space-efficient usage of filter() using iterators.

Fast filtering functions in Cython

The featurefuncs module contains some ready-made functions written in Cython that will be faster than pure Python equivalents. For example, there are greater_than() and less_than() functions, which are about 70% faster. In IPython:

```
>>> from pybedtools.featurefuncs import greater_than
>>> len(a)
310456
>>> def L(x,width=100):
... return len(x) > 100

>>> %timeit a.filter(greater_than, 100)
1 loops, best of 3: 1.74 s per loop
>>> %timeit a.filter(L, 100)
1 loops, best of 3: 2.96 s per loop
```

2.3.10 Each

Similar to BedTool.filter(), which applies a function to return True or False given an Interval, the BedTool.each() method applies a function to return a new, possibly modified Interval.

The <code>BedTool.each()</code> method applies a function to every feature. Like <code>BedTool.filter()</code>, you can use your own function or some pre-defined ones in the featurefuncs module. Also like filter(), *args and **kwargs are sent to the function.

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> # The results of an "intersect" with c=True will return features
>>> # with an additional field representing the counts.
>>> with_counts = a.intersect(b, c=True)
```

Let's define a function that will take the number of counts in each feature as calculated above and divide by the number of bases in that feature. We can also supply an optional scalar, like 0.001, to get the results in "number of intersections per kb". We then insert that value into the score field of the feature. Here's the function:

```
>>> def normalize_count(feature, scalar=0.001):
       assume feature's last field is the count
. . .
. . .
       counts = float(feature[-1])
       normalized = counts / len(feature) * scalar
       # need to convert back to string to insert into feature
       feature.score = str(normalized)
       return feature
And we apply it like this:
>>> normalized = with_counts.each(normalize_count)
>>> print normalized
chr1
          1
                  100
                           feature1
                                           0.0
                                                           0
           100
                   200
                                           1e-05 +
                                                           1
chr1
                          feature2
          150
                   500
                                           2.85714285714e-06
chr1
                           feature3
                                                                           1
                                           2e-05 +
           900
                   950
                           feature4
```

2.3.11 Using the history and tags

chr1

BEDTools makes it very easy to do rather complex genomic algebra. Sometimes when you're doing some exploratory work, you'd like to rewind back to a previous step, or clean up temporary files that have been left on disk over the course of some experimentation.

To assist this sort of workflow, BedTool instances keep track of their history in the BedTool.history attribute. Let's make an example BedTool, c, that has some history:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> c = a.intersect(b, u=True)
```

c now has a history which tells you all sorts of useful things (described in more detail below):

```
>>> print c.history
(HistoryStep> bedtool("/home/ryan/pybedtools/pybedtools/test/a.bed").intersect("/home/ryan/pybedtools/
```

There are several things to note here. First, the history describes the full commands, including all the names of the temp files and all the arguments that you would need to run in order to re-create it. Since BedTool objects are fundamentally file-based, the command refers to the underlying filenames (i.e., a.bed and b.bed) instead of the BedTool instances (i.e., a and b). A simple copy-paste of the command will be enough re-run the command. While this may be useful in some situations, be aware that if you do run the command again you'll get another temp file that has the same contents as c's temp file.

To avoid such cluttering of your temp dir, the history also reports tags. BedTool objects, when created, get a random tag assigned to them. You can get get the BedTool associated with tag with the pybedtools.find_tagged()

2.3. Tutorial Contents 17 function. These tags are used to keep track of instances during this session.

So in this case, we could get a reference to the a instance with:

```
>>> should_be_a = pybedtools.find_tagged('klkreuay')
```

Here's confirmation that the parent of the first step of c's history is a (note that HistoryStep objects have a HistoryStep.parent_tag and HistoryStep.result_tag):

```
>>> pybedtools.find_tagged(c.history[0].parent_tag) == a
True
```

Let's make something with a more complicated history:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> c = a.intersect(b)
>>> d = c.slop(g=pybedtools.chromsizes('hg19'), b=1)
>>> e = d.merge()
>>> # this step adds complexity!
>>> f = e.subtract(b)
```

Let's see what the history of f (the last BedTool created) looks like . . . note that here I'm formatting the results to make it easier to see:

```
>>> print f.history
Γ
[
    [
       |<HistoryStep> BedTool("/usr/local/lib/python2.6/dist-packages/pybedtools/test/data/a.bea
                                  "/usr/local/lib/python2.6/dist-packages/pybedtools/test/data/b.bed
        parent tag: rzrztxlw,
                                  result tag: ifbsanqk
    -
           ],
        |<HistoryStep> BedTool("/tmp/pybedtools.BgULVj.tmp").slop(
    b=1, genome="hg19"
    ),
    parent tag: ifbsangk,
    result tag: omfrkwjp
       |<HistoryStep> BedTool("/tmp/pybedtools.SFmbYc.tmp").merge(),
                         parent tag: omfrkwjp,
                          result tag: zlwqblvk
    -
   ],
|<HistoryStep> BedTool("/tmp/pybedtools.wlBiMo.tmp").subtract(
                      "/usr/local/lib/python2.6/dist-packages/pybedtools/test/data/b.bed",
                      ),
                      parent tag: zlwqblvk,
                      result tag: reztxhen
```

Those first three history steps correspond to c, d, and e respectively, as we can see by comparing the code snippet above with the commands in each history step. In other words, e can be described by the sequence of 3 commands in the first three history steps. In fact, if we checked e.history, we'd see exactly those same 3 steps.

When f was created above, it operated both on e, which had its own history, as well as b – note the nesting of the list. You can do arbitrarily complex "genome algebra" operations, and the history of the BEDTools will keep track of this. It may not be useful in every situation, but the ability to backtrack and have a record of what you've done can sometimes be helpful.

2.3.12 Deleting temp files specific to a single BedTool

You can delete temp files that have been created over the history of a <code>BedTool</code> with <code>BedTool.delete_temporary_history()</code>. This method will inspect the history, figure out which items point to files in the temp dir (which you can see with <code>get_tempdir()</code>), and prompt you for their deletion:

```
>>> f.delete_temporary_history()
Delete these files?
    /tmp/pybedtools..BgULVj.tmp
    /tmp/pybedtools.SFmbYc.tmp
    /tmp/pybedtools.wlBiMo.tmp
(y/N) y
```

Note that the file that f points to is left alone. To clarify, the <code>BedTool.delete_temporary_history()</code> will only delete temp files that match the pattern <TEMP_DIR>/pybedtools.*.tmp from the history of f, up to but not including the file for fitself. Any <code>BedTool</code> instances that do not match the pattern are left alone. Use the kwarg <code>ask=False</code> to disable the prompt.

2.4 Topical Documentation

This section contains additional documentation not covered in the tutorial.

2.4.1 Design principles

Hopefully, understanding (or just being aware of) these design principles will help in getting the most out of pybedtools and working efficiently.

Principle 1: Temporary files are created automatically

Using BedTool instances typically has the side effect of creating temporary files on disk. Even when using the iterator protocol of BedTool objects, temporary files may be created in order to run BEDTools programs (see **Iterators**_ for more on this latter topic).

Let's illustrate some of the design principles behind pybedtools by merging features in a . bed that are 100 bp or less apart (d=100) in a strand-specific way (s=True):

```
>>> from pybedtools import BedTool
>>> import pybedtools
>>> a = BedTool(pybedtools.example_filename('a.bed'))
>>> merged_a = a.merge(d=100, s=True)
```

Now merged a is a BedTool instance that contains the results of the merge.

BedTool objects must always point to a file on disk. So in the example above, merged_a is a BedTool, but what file does it point to? You can always check the BedTool.fn attribute to find out:

```
>>> # what file does 'merged_a' point to?
>>> merged_a.fn
'/tmp/pybedtools.MPPp5f.tmp'
```

Note that the specific filename will be different for you since it is a randomly chosen name (handled by Python's tempfile module). This shows one important aspect of pybedtools: every operation results in a new temporary file. Temporary files are stored in /tmp by default, and have the form /tmp/pybedtools.*.tmp.

When you are done using the pybedtools module, make sure to clean up all the temp files created with:

```
>>> # Don't do this yet if you're following the tutorial!
>>> pybedtools.cleanup()
```

If you forget to do this, from the command line you can always do a:

```
rm /tmp/pybedtools.*.tmp
```

to clean everything up. Alternatively, in this session or another session you can use:

```
>>> pybedtools.cleanup(remove_all=True)
```

to remove all files that match the pattern <tempdir>/pybedtools.*.tmp where <tempdir> is the current value of pybedtools.get_tempdir().

If you need to specify a different directory than that used by default by Python's tempdir module, then you can set it with:

```
>>> pybedtools.set_tempdir('/scratch')
```

You'll need write permissions to this directory, and it needs to already exist. All temp files will then be written to that directory, until the tempdir is changed again.

Principle 2: Names and arguments are as similar as possible to BEDTools

As much as possible, BEDTools programs and BedTool methods share the same names and arguments.

Returning again to this example:

```
>>> merged_a = a.merge(d=100, s=True)
```

This demonstrates that the <code>BedTool</code> methods that wrap BEDTools programs do the same thing and take the exact same arguments as the BEDTools program. Here we can pass <code>d=100</code> and <code>s=True</code> only because the underlying BEDTools program, <code>mergeBed</code>, can accept these arguments. Need to know what arguments <code>mergeBed</code> can take? See the docs for <code>BedTool.merge()</code>; for more on this see <code>Principle 7</code>: Check the help.

In general, remove the "Bed" from the end of the BEDTools program to get the corresponding BedTool method. So there's a BedTool.subtract() method for subtractBed, a BedTool.intersect() method for intersectBed, and so on.

Principle 3: Indifference to BEDTools version

Since BedTool methods just wrap BEDTools programs, they are as up-to-date as the version of BEDTools you have installed on disk. If you are using a cutting-edge version of BEDTools that has some hypothetical argument -z for intersectBed, then you can use a intersectBed (z=True).

pybedtools will also raise an exception if you try to use a method that relies on a more recent version of BEDTools than you have installed.

Principle 4: Sensible default args

If we were running the mergeBed program from the command line, we would would have to specify the input file with the mergeBed -i option.

pybedtools assumes that if we're calling the merge() method on the BedTool, a, we want to operate on the bed file that a points to.

In general, BEDTools programs that accept a single BED file as input (by convention typically specified with the -i option) the default behavior for pybedtools is to use the BedTool's file (indicated in the BedTool fin attribute) as input.

We can still pass a file using the i keyword argument if we wanted to be absolutely explicit. In fact, the following two versions produce the same output:

```
>>> # The default is to use existing file for input -- no need
>>> # to specify "i" . . .
>>> result1 = a.merge(d=100, s=True)
>>> # . . . but you can always be explicit if you'd like
>>> result2 = a.merge(i=a.fn, d=100, s=True)
>>> # Confirm that the output is identical
>>> result1 == result2
True
```

Methods that have this type of default behavior are indicated by the following text in their docstring:

```
.. note::

For convenience, the file this BedTool object points to is passed as "-i"
```

There are some BEDTools programs that accept two BED files as input, like intersectBed where the the first file is specified with -a and the second file with -b. The default behavior for pybedtools is to consider the BedTool's file as -a and the first non-keyword argument to the method as -b, like this:

```
>>> b = pybedtools.example_bedtool('b.bed')
>>> result3 = a.intersect(b)
```

This is exactly the same as passing the a and b arguments explicitly:

```
>>> result4 = a.intersect(a=a.fn, b=b.fn)
>>> result3 == result4
True
```

Furthermore, the first non-keyword argument used as -b can either be a filename *or* another BedTool object; that is, these commands also do the same thing:

```
>>> result5 = a.intersect(b=b.fn)
>>> result6 = a.intersect(b=b)
>>> str(result5) == str(result6)
True
```

Methods that accept either a filename or another BedTool instance as their first non-keyword argument are indicated by the following text in their docstring:

```
.. note::
    This method accepts either a BedTool or a file name as the first
    unnamed argument
```

Principal 5: Other arguments have no defaults

Only the BEDTools arguments that refer to BED (or other interval) files have defaults. In the current version of BEDTools, this means only the -i, -a, and -b arguments have defaults. All others have no defaults specified by pybedtools; they pass the buck to BEDTools programs. This means if you do not specify the d kwarg when calling BedTool.merge(), then it will use whatever the installed version of BEDTools uses for -d (currently, mergeBed's default for -d is 0).

-d is an option to BEDTools mergeBed that accepts a value, while -s is an option that acts as a switch. In pybedtools, simply pass a value (integer, float, whatever) for value-type options like -d, and boolean values (True or False) for the switch-type options like -s.

Here's another example using both types of keyword arguments; the BedTool object b (or it could be a string filename too) is implicitly passed to intersectBed as -b (see *Principle 4: Sensible default args* above):

```
>>> a.intersect(b, v=True, f=0.5)
```

Again, any option that can be passed to a BEDTools program can be passed to the corresonding BedTool method.

Principle 6: Chaining together commands

Most methods return new BedTool objects, allowing you to chain things together just like piping commands together on the command line. To give you a flavor of this, here is how you would get the merged regions of features shared between a.bed (as referred to by the BedTool a we made previously) and b.bed: (as referred to by the BedTool b):

```
>>> a.intersect(b).merge().saveas('shared_merged.bed')
<BedTool(shared_merged.bed)>
```

This is equivalent to the following BEDTools commands:

```
intersectBed -a a.bed -b b.bed | merge -i stdin > shared_merged.bed
```

Methods that return a new BedTool instance are indicated with the following text in their docstring:

```
.. note::
    This method returns a new BedTool instance
```

Principle 7: Check the help

If you're unsure of whether a method uses a default, or if you want to read about what options an underlying BED-Tools program accepts, check the help. Each pyBedTool method that wraps a BEDTools program also wraps the BEDTools program help string. There are often examples of how to use a method in the docstring as well. The documentation is also run through doctests, so the code you read here is guaranteed to work and be up-to-date.

2.4.2 Creating a BedTool

To create a BedTool, first you need to import the pybedtools module. For these examples, I'm assuming you have already done the following:

```
>>> import pybedtools
>>> from pybedtools import BedTool
```

Next, you need a BED file to work with. If you already have one, then great – move on to the next section. If not, pybedtools comes with some example bed files used for testing. You can take a look at the list of example files that ship with pybedtools with the list example files () function:

```
>>> # list the example bed files
>>> pybedtools.list_example_files()
['a.bed', 'b.bed', 'c.gff', 'd.gff', 'dm3-chr2L-5M-invalid.gff.gz', 'dm3-chr2L-5M.gff.gz', 'dmel-all-
```

Once you decide on a file to use, feed the your choice to the example_filename () function to get the full path:

```
>>> # get the full path to an example bed file
>>> bedfn = pybedtools.example_filename('a.bed')
```

The full path of bedfn will depend on your installation (this is similar to the data() function in \mathbf{R}_{-} , if you're familiar with that).

Now that you have a filename – either one of the example files or your own, you create a new BedTool simply by pointing it to that filename:

```
>>> # create a new BedTool from the example bed file
>>> myBedTool = BedTool(bedfn)
```

Alternatively, you can construct BED files from scratch by using the from_string keyword argument. However, all spaces will be converted to tabs using this method, so you'll have to be careful if you add "name" columns. This can be useful if you want to create *de novo* BED files on the fly:

```
>>> # an "inline" example:
>>> fromscratch1 = pybedtools.BedTool('chrX 1 100', from_string=True)
>>> print fromscratch1
chrX 1
           100
>>> # using a longer string to make a bed file. Note that
>>> # newlines don't matter, and one or more consecutive
>>> # spaces will be converted to a tab character.
>>> larger_string = """
... chrX 1 100
                  feature1 0 +
... chrX 50 350 feature2 0 -
... chr2 5000 10000 another_feature 0 +
>>> fromscratch2 = BedTool(larger_string, from_string=True)
>>> print fromscratch2
      1 100 feature1
                           0
chrX
       50 350 feature2
chrX
                           \cap
chr2
               10000 another_feature 0
```

Of course, you'll usually be using your own bed files that have some biological importance for your work that are saved in places convenient for you, for example:

```
>>> a = BedTool('/data/sample1/peaks.bed')
```

2.4.3 Using BedTool objects as iterators/generators

Typically, BedTool objects are used somewhat like handles to individual files on disk that contain BED lines. To save disk space, BedTool objects also have the ability to "stream", much like piping in Unix. That is, the data are created only one line at a time in memory, instead of either creating a list of all data in memory or writing all data to disk.

Warning: You'll need to be careful when using BedTool objects as generators, since any operation that reads all the features of a BedTool will consume the iterable.

To get a streaming BedTool, use the stream=True kwarg. This BedTool will act a little differently from a standard, file-based BedTool.

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> c = a.intersect(b, stream=True)
>>> # checking the length consumes the iterator
>>> len(c)
3
>>> # nothing left, so checking length again returns 0
>>> len(c)
0
```

In some cases, a stream may be "rendered" to a temp file. This is because BEDTools programs can only accept one input file as stdin. This is typically the first input (-i or -a), while the other input (-b) must be a file. Consider this example, where the second intersection needs to convert the streaming BedTool to a file before sending to intersectBed:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> # first we set up a streaming BedTool:
>>> c = a.intersect(b, stream=True)
>>> # But supplying a streaming BedTool as the first unnamed argument
>>> # means it is being passed as -b to intersectBed, and so must be a file.
>>> # In this case, 'c' is rendered to a tempfile before being passed.
>>> d = a.intersect(c, stream=True)
```

Creating a BedTool from an iterable

You can create a BedTool on the fly from a generator or iterator — in fact, this is what the BedTool.filter() method does for you:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> print a
          1
                   100
                                            0
chr1
                           feature1
          100
chr1
                   200
                           feature2
                                            0
                   500
                                            0
chr1
           150
                           feature3
           900
                   950
                            feature4
                                            0
chr1
>>> b = pybedtools.BedTool(f for f in a if f.start > 200)
>>> # this is the same as using filter:
>>> c = a.filter(lambda x: x.start > 200)
```

We need to "render" these BedTools to string before we can check equality – consuming them both – since they are both iterables for which == is not defined:

```
>>> b == c
Traceback (most recent call last):
```

NotImplementedError: Testing equality only supported for BedTools that point to a file
>>> str(b) == str(c)
True

Indexing a BedTool

In some cases it may be useful to index into a BedTool object. We can use standard list slice syntax, and get an iterable of Interval objects as a result. This iterable can in turn be used to create a new BedTool instance:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> a[2:4]
<itertools.islice object at 0x...>
>>> for i in a[2:4]:
... print i
                   500
                           feature3
chr1
          150
chr1
          900
                   950
                           feature4
>>> b = pybedtools.example_bedtool('b.bed')
>>> print pybedtools.BedTool(a[:3]).intersect(b)
                   200
chr1
           155
                         feature2
           155
                   200
chr1
                           feature3
                                           0
```

2.4.4 Low-level operations

We can use the $BedTool.as_intervalfile()$ method to return an IntervalFile instance. This class provides low-level support to the BEDTools C++ API.

The method IntervalFile.all_hits() takes a single Interval as the query and returns a list of all features in the IntervalFile that intersect:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> ivf = a.as_intervalfile()
>>> query = a[2]
>>> ivf.all_hits(query)
[Interval(chr1:100-200), Interval(chr1:150-500)]
```

Similarly, we can just return if there were *any* hits, a much faster operation:

```
>>> ivf.any_hits(query)
1
```

Or count how many hits:

```
>>> ivf.count_hits(query)
2
```

See the docstrings for IntervalFile.all_hits(), IntervalFile.any_hits(), and IntervalFile.count_hits() for more, including stranded hits and restricting hits to a specific overlap.

2.4.5 Working with BAM files

Some BEDTools programs, like intersecteBed, support BAM files as input. From the command line, you would need to specify the -abam argument to do so. However, pybedtools auto-detects BAM files and passes the abam argument automatically for you. That means if you create a BamTool out of a BAM file, like this:

```
x = pybedtools.example_bedtool('gdc.bam')
```

you can intersect it with a BED file without doing anything special:

```
b = pybedtools.example_bedtool('gdc.gff')
y = x.intersect(b)
```

The output of this operation follows the semantics of BEDTools. That is, for programs like intersectBed, if abam is used then the output will be BAM format as well. But if the -bed argument is passed, then the output will be BED format. Similarly, in pybedtools, if a BAM file is used to create the BedTool then the results will also be in BAM format. If the bed=True kwarg is passed, then the results be in BED format.

As an example, let's intersect a BAM file of reads with annotations using files that ship with pybedtools. First, we create the BedTool objects:

```
>>> a = pybedtools.example_bedtool('x.bam')
>>> b = pybedtools.example_bedtool('dm3-chr2L-5M.gff.gz')
```

The first call below will return BAM results, and the second will return BED results.

```
>>> bam_results = a.intersect(b)
>>> bed_results = a.intersect(b, bed=True)
```

We can iterate over BAM files to get Interval objects just like iterating over BED or GFF files. Indexing works, too:

```
>>> for i in bam_results[:2]:
       print i
HWUSI-NAME:2:69:512:1017#0 16
                                    chr2L
                                            9330
                                                    3
                                                            36M
                                                                             0
                                                                                     0
HWUSI-NAME:2:91:1201:1113#0 16
                                            10213
                                                    255
                                    chr2L
                                                            36M
                                                                                     0
>>> bam_results[0]
Interval (chr2L: 9329-9365)
>>> bam_results[:10]
<itertools.islice object at ...>
>>> cigar_string = i[5]
```

Note that pybedtools uses the convention that BAM features in plain text format are considered SAM features, so these SAM features are **one-based and include the stop coordinate** as illustrated below:

```
>>> bam_results[0].start
9329L
>>> bam_results[0][3]
'9330'
```

Currently, the stop coordinate is defined as the start coord plus the length of the sequence; eventually a more sophisticated, CIGAR-aware approach may be used. Similarly, the length is defined to be stop - start, again, not CIGAR-aware at the moment. For more sophisticated low-level manipulation of BAM features, you might want to consider using HTSeq.

TACAAATC'

TGTAGAAT

When we specified the bed=True kwarg above, the intersected BAM results are converted to BED format. We can use those like a normal BED file. Note that since we are viewing BED output, *the start and stops are 0-based*:

```
>>> d = a.intersect(b, bed=True)
>>> d.head(3)
                     9365
                             HWUSI-NAME:2:69:512:1017#0
                                                              3
chr2L
            9329
chr2L
            9329
                     9365
                             HWUSI-NAME:2:69:512:1017#0
                                                              3
chr2L
            9329
                     9365
                             HWUSI-NAME:2:69:512:1017#0
```

Consistent with BEDTools programs, BAM files are **not** supported as the second input argument. In other words, intersectBed does not have both -abam and -bbam arguments, so pybedtools will not not allow this either.

However, pybedtools does allow streaming BAM files to be the input of methods that allow BAM input as the first input. In this [trivial] example, we can stream the first intersection to save disk space, and then send that streaming BAM to the next BedTool.intersect() call. Since it's not streamed, the second intersection will be saved as a temp BAM file on disk:

```
>>> a.intersect(b, stream=True).intersect(b)
```

2.4.6 Notes on BAM file semantics

These are some implementation notes about how BAM files are handled by mod:pybedtools for those interested in the implementation.

The initial creation of a BedTool that points to a file will trigger a check on the first 15 bytes of a file to see if it's a BAM file. If so, then the BedTool's _isbam attribute is set to True. If the BedTool is a stream, then the check will not be made, and it is up to the creator (whether it's the user on the command line or a method or function) to set the BAM-streaming BedTool's ._isbam attribute to True. This is handled automatically for wrapped BEDTools programs (described below).

Some BEDTools programs natively handle BAM files. The <code>@_wraps</code> decorator that is used to wrap each method has a bam kwarg that specifies what input argument the wrapped tool will accept as BAM (for example, the wrapper for <code>intersectBed</code> has the kwarg <code>bam="abam"</code>).

If self._isbam == True, then self.fn is passed to the bam input arg instead of the default implicit input arg (so intersectBed, self.fn is passed as abam instead of -a).

Trying to call a method that does not have a bam kwarg registered will result in a ValueError, along with a message that says to use <code>BedTool.bam_to_bed()</code> first. For example, <code>subtractBed</code> currently doesn't accept BAM files as input, so this doesn't work:

```
>>> a = pybedtools.example_bedtool('gdc.bam')
>>> b = pybedtools.example_bedtool('gdc.gff')
>>> # doesn't work:
>>> c = a.subtract(b)
```

However, converting to a to BED format first (and setting stream=True to save on disk I/O) works fine:

```
>>> # works:
>>> c = a.bam_to_bed(stream=True).subtract(b)
```

Iterating over a file-based BedTool that points to a BAM will call samtools view and yields lines which sent to IntervalIterator, which splits the lines and passes them to create_interval_from_list which in turn decides on the fly whether it's gff, bed, or sam.

However, we can't easily check the first 15 bytes of a streaming BedTool, because that would consume those bytes. The @_wraps decorator needs to know some information about which arguments to a wrapped program result in BAM output and which result in non-BAM output.

```
Given a = BedTool('x.bam'):
```

- c = a.intersect(b) creates BAM output, so it returns a new BedTool with c._isbam = True.
- a.intersect (b, bed=True) returns BED output. @_wraps needs to know, if the input was BAM, which kwarg[s] disable BAM output. For example, if -bed is passed to intersectBed, the output will NOT be BAM. This is implemented with the nonbam kwarg for _wraps(). In this case, the resulting BED file is treated like any other BED file.
- c = a.intersect (b, stream=True) returns streaming BAM output. In this case, iterating over c will send the BAM stream to stdin of a samtools call

2.4.7 Specifying genomes

This section illustrates the use of genome files for use with BEDTools programs that need to know chromosome limits to prevent out-of-range coordinates.

Using BEDTools programs like slopBed or shuffleBed from the command line requires "genome" or "chrom-sizes" files. pybedtools comes with common genome assemblies already set up as a dictionary with chromosomes as keys and zero-based (start, stop) tuples as values:

```
>>> from pybedtools import genome_registry
>>> genome_registry.dm3['chr2L']
(0, 23011544)
```

The rules for specifying a genome for methods that require a genome are as follows (use whatever is most convenient):

- Use g to specify either a filename or a dictionary
- Use genome to specify either an assembly name or a dictionary

Below are examples of each.

As a file

This is the typical way of using BEDTools programs, by specifying an existing genome file with g:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = a.slop(b=100, g='hg19.genome')
```

As a string

This is probably the most convenient way of specifying a genome. If the genome exists in the genome registry it will be used directly; otherwise it will automatically be downloaded from UCSC. You must use the genome kwarg for this; if you use g a string will be interpreted as a filename:

```
>>> c = a.slop(b=100, genome='hg19')
```

As a dictionary

This is a good way of providing custom coordinates; either g or genome will accept a dictionary:

```
>>> d = a.slop(b=100, g={'chr1':(1, 10000)})
>>> e = a.slop(b=100, genome={'chr1':(1,100000)})
```

Make sure that all these different methods return the same results

```
>>> b == c == d == e
```

Converting to a file

Since **BEDTools**_ programs operate on files, the fastest choice will be to use an existing file. While the time to convert a dictionary to a file is extremely small, over 1000's of files (e.g., for Monte Carlo simulations), the time may add up. The function pybedtools.chromsizes_to_file() will create a file from a dictionary or string:

```
>>> # with no filename specified, a tempfile will be created
>>> pybedtools.chromsizes_to_file(pybedtools.chromsizes('dm3'), 'dm3.genome')
'dm3.genome'
>>> print open('dm3.genome').read()
chr2L 23011544
chr2LHet 368872
chr2R
         21146708
chr2RHet 3288761
         24543557
chr3L
chr3LHet 2555491
         27905053
chr3R
chr3RHet 2517507
          1351857
chr4
          19517
chrM
chrU
          10049037
chrUextra 29004656
        22422827
chrX
         204112
chrXHet
chrYHet
         347038
```

2.4.8 Randomization

pybedtools provides some basic functionality for assigning some significance value to the overlap between two BEDfiles.

The strategy is to randomly shuffle a file many times, each time doing an intersection with another file of interest and counting the number of intersections. Upon doing this many times, an empirical distribution is constructed, and the number of intersections between the original, un-shuffled file is compared to this empirical distribution to obtain a p-value, or compared to the median of the distribution to get a score.

There are two methods, pybedtools.BedTool.randomintersection() which does the brute force randomizations, and BedTool.randomstats() which compiles and reports the results from the former method.

Example workflow

As a somewhat trivial example, we'll intersect the example a .bed with b .bed, taking care to set some options that will let it run in a deterministic way so that these tests will run.

We will be shuffling a.bed, so we'll need to specify the limits of its chromosomes with BedTool.set_chromsizes(). Here, we set it to an artifically small chromosome size so that we can get some meaningful results in reasonable time. In practice, you would either supply your own dictionary or use a string assembly name (e.g., 'hg19', 'mm9', 'dm3', etc). The genome-handling code will find the chromsizes we've set, so there's no need to tell shuffleBed which genome file to use each time.

```
>>> chromsizes = {'chr1': (0, 1000)}
>>> a = pybedtools.example_bedtool('a.bed').set_chromsizes(chromsizes)
>>> b = pybedtools.example_bedtool('b.bed')
```

We have the option of specifying what kwargs to provide <code>BedTool.shuffle()</code> and <code>BedTool.intersect()</code>, which will be called each iteration. In this example, we'll tell <code>shuffleBed</code> to only shuffle within the chromsome just to illustrate the kwargs passing. We also need to specify how many iterations to perform. In practice, 1000 or 10000 are good numbers, but for the sake of this example we'll only do 100.

Last, setting debug=True means that the random seed will be set in a predictable manner so that we'll always get the same results for testing. In practice, make sure you use debug=False (the default) to ensure random results.

```
>>> results = a.randomintersection(b, iterations=100, shuffle_kwargs={'chrom': True}, debug=True)
```

results is a generator of intersection counts where each number is the number of times the shuffled a intersected with b. We need to convert it to a list in order to look at it:

```
>>> results = list(results)
>>> len(results)
100

>>> print results[:10]
[1, 1, 2, 2, 1, 2, 1, 0, 2, 3]
```

Running thousands of iterations on files with many features will of course result in more complex results. We could then take these results and plot them in matplotlib, or get some statistics on them.

The method BedTool.randomstats() does this for us, but requires NumPy and SciPy to be installed. This method also calls BedTool.randomintersection() for us, returning the summarized results in a dictionary.

```
BedTool.randomstats() takes the same arguments as BedTool.randomintersection():
>>> results_dict = a.randomstats(b, iterations=100, shuffle_kwargs={'chrom': True}, debug=True)
```

The keys to this results dictionary are as follows (some are redundant, I've found these keys useful for writing out to file):

iterations the number of iterations we specified

actual the number of intersections between then un-shuffled a and b

file_a the filename of a

file b the filename of b

<file_a> the key is actully the filename of a, and the value is the number of features in a

<file_b> the key is actually the filename of b and the value is the number of features in b

self number of features in a (or "self"; same value as for <file_a>)

other number of features in b (or "other"; same value as for <file_b>)

frac randomized above actual fraction of iterations that had counts above the actual count

frac randomized below actual fraction of iterations that had counts below the actual count

median randomized the median of the distribution of randomized intersections

normalized the actual count divided by the median; can be considered as a score

percentile the percentile of actual within the distribution of randomized intersections; can be considered an empirical p-value

upper 97.5th the 97.5th percentile of the randomized distribution

lower 2.5th the 2.5th percentile of the randomized distribution

For example:

```
>>> keys = ['self', 'other', 'actual', 'median randomized', 'normalized', 'percentile']
>>> for key in keys:
... print '%s: %s' % (key, results_dict[key])
self: 4
other: 2
actual: 3
median randomized: 2.0
normalized: 1.5
percentile: 92.0
```

Contributions toward improving this code or implementing other methods of statistical testing are very welcome!

2.4.9 Wrapping new tools

This section serves as a reference for wrapping new tools as they are added to BEDTools.

Example program description

Let's assume we would like to wrap a new program, appropriately named newProgramBed. Its signature from the command line is newProgramBed -a <infile> -b <other file> [options], and it accepts -a stdin to indicate data is being piped to it:

```
newProgramBed -a <BED/VCF/GFF> -b <BED/VCF/GFF> [options]
```

Method name

Generally, I've tried to keep method names as similar as possible to BEDTools programs while still being PEP8-compliant. The trailing 'Bed' is usually removed from the program name. So here the name would probably be new_program.

Define a method in BedTool

Define a method in BedTool . . . and *don't add any content to the function body*. This is because the decorator we're about to add will replace the method wholesale; anything that's in the function body will effectively be ignored.

```
def new_program(self):
    pass
```

Add the _wraps() decorator

This is where most of the work happens.

Since most of the work of wrapping BEDTools programs needs to happen every time a new program is wrapped, this work is abstracted out into the _wraps() function. The _wraps() docstring and source is the best place to learn the details on what it's doing; here we'll focus on using it.

Our hypothetical program, newProgramBed, takes -a as the first input. We'd like to have -a implicitly be passed as whatever our BedTool already points to, so we use the implicit='a' kwarg to _wraps() here. newProgramBed also takes a second input, -b. We describe that to the wrapper with the other='b' kwarg.

Any other keyword args that are used when calling the method will automatically be passed to the program. So if newProgramBed has an optional -s argument, we don't need to specify that here. When the user passes an s=True kwarg, it will be passed automatically to newProgramBed as the argument -s. If newProgramBed does not accept a -z argument but the user passes one anyway, we rely on the BEDTools program to do the error-checking of arguments and report any errors back to Python.

Here's what the new method looks like so far:

```
@_wraps (prog='newProgramBed', implicit='a', other='b')
def new_program(self):
    pass
```

For wrapped programs that expect a genome file or have more complex arguments, see the docstring and source for wrap ().

Add doctests

While the function body will be replaced wholesale by the decorator, the docstring will be copied to the new function. This is important because it means we can write meaningful documentation and, even more importantly, doctests for this method. Writing a doctest within the method's docstring means it will automatically be found by the test suite.

```
@_wraps (prog='newProgramBed', implicit='a', other='b')
def new_program(self):
    Converts all features to length of 1.
   Example usage:
   >>> a = pybedtools.example_bedtool('a.bed')
    >>> b = pybedtools.example_bedtool('b.bed')
    >>> c = a.new_program(b, s=True)
    >>> print c #+NORMALIZE_WHITESPACE
    chrl 1
                   2
          100
    chr1
                   101
           150
                   151
    chr1
   chr1 900
    <BLANKLINE>
```

Summary

That's it! We now have a method, BedTool.new_program(), that wraps a hypothetical newProgramBed BED-Tools program, will accept any optional args that newProgramBed does, will return a new BedTool containing the results, and it's tested.

This new method can be be chained with other BedTool instances, used as an iterator or generator, or anything else a normal BedTool can do . . . for example:

```
a = pybedtools.example_bed('a.bed')
b = pybedtools.example_bed('b.bed')
c = a.new_program(b, s=True).filter(lambda x: x.start < 125).saveas('t.bed', trackline='track name="orange")</pre>
```

2.5 Scripts

pybedtools comes with several scripts that illustrate common use-cases.

In Python 2.7, you can use:

python -m pybedtools

to get a list of scripts and their description.

2.5.1 Venn diagram scripts

There are two scripts for making Venn diagrams, depending on how you'd like the diagrams to look. Both simply take 3 BED files as input. venn_gchart.py uses the Google Chart API, while venn_mpl.py uses matplotlib if you have it installed.

Upon installing pybedtools, these scripts should be available on your path. Calling them with the -h option will print the help, and using the --test option will run a test, creating a new file out.png in the current working directory.

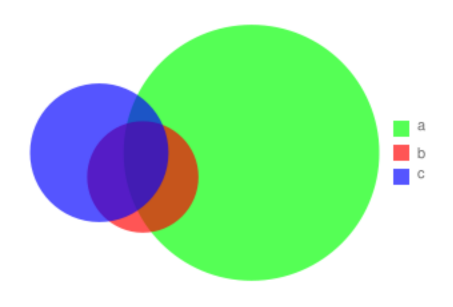


Figure 2.1: Above: using --test with venn_gchart.py results in this figure

2.5.2 Intron/exon classification

The script intron_exon_reads.py accepts a GFF file (with introns and exons annotated) and a BAM file. When complete, it prints out the number of exonic, intronic, and both intronic and exonic (i.e., from overlapping genes or isoforms). This script is also a good example of how to do use Python's multiprocessing for parallel computation.

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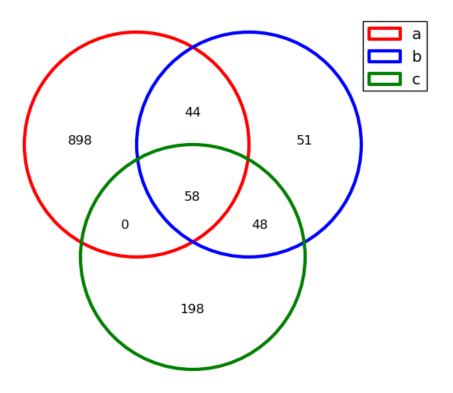


Figure 2.2: Above: Result of using --test with venn_mpl.py

2.5.3 Annotate.py

The annotate.py script extends closestBed by classifying features (intron, exon) that are a distance of 0 away from the query features.

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· Interval class

· IntervalFile class

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Chapter 2. Contents:

2.6 pybedtools Reference

This section is the module reference documentation, and includes the full docstrings for methods and functions in pybedtools. It is separated into module-level, BedTool methods that wrap BEDTools programs, and BedTool methods unique to pybedtools.

class pybedtools.BedTool (fn, from_string=False)

```
___init___(fn, from_string=False)
```

Wrapper around Aaron Quinlan's BEDtools suite of programs (https://github.com/arq5x/bedtools); also contains many useful methods for more detailed work with BED files.

fn is typically the name of a BED-like file, but can also be one of the following:

- •a string filename
- •another BedTool object
- •an iterable of Interval objects
- •an open file object
- •a "file contents" string (see below)

If *from_string* is True, then you can pass a string that contains the contents of the BedTool you want to create. This will treat all spaces as TABs and write to tempfile, treating whatever you pass as *fn* as the contents of the bed file. This also strips empty lines.

Typical usage is to point to an existing file:

```
a = BedTool('a.bed')
```

But you can also create one from scratch from a string:

```
>>> s = '''
... chrX 1 100
... chrX 25 800
... '''
>>> a = BedTool(s,from_string=True).saveas('a.bed')
```

Or use examples that come with pybedtools:

```
>>> example_files = pybedtools.list_example_files()
>>> assert example_files[0] == 'a.bed'
>>> a = pybedtools.example_bedtool('a.bed')
```

2.6.1 pybedtools module-level functions

Functions for working with example files

```
pybedtools.example_bedtool(fn)
```

Return a bedtool using a bed file from the pybedtools examples directory. Use <code>list_example_files()</code> to see a list of files that are included.

```
{\tt pybedtools.example\_filename}\ (fn)
```

Return a bed file from the pybedtools examples directory. Use <code>list_example_files()</code> to see a list of files that are included.

```
pybedtools.list_example_files()
```

Returns a list of files in the examples dir. Choose one and pass it to <code>example_filename()</code> to get the full path to an example file.

Example usage:

```
>>> choices = list_example_files()
>>> assert 'a.bed' in choices
>>> bedfn = example_filename('a.bed')
>>> mybedtool = BedTool(bedfn)
```

pybedtools.data_dir()

Returns the data directory that contains example files for tests and documentation.

Functions for specifying genome assemblies

```
pybedtools.chromsizes(genome)
```

Looks for a *genome* already included in the genome registry; if not found then it looks it up on UCSC. Returns the dictionary of chromsize tuples where each tuple has (start,stop).

Chromsizes are described as (start, stop) tuples to allow randomization within specified regions; e. g., you can make a chromsizes dictionary that represents the extent of a tiling array.

Example usage:

```
>>> dm3_chromsizes = chromsizes('dm3')
>>> for i in sorted(dm3_chromsizes.items()):
       print i
('chr2L', (0, 23011544))
('chr2LHet', (0, 368872))
('chr2R', (0, 21146708))
('chr2RHet', (0, 3288761))
('chr3L', (0, 24543557))
('chr3LHet', (0, 2555491))
('chr3R', (0, 27905053))
('chr3RHet', (0, 2517507))
('chr4', (0, 1351857))
('chrM', (0, 19517))
('chrU', (0, 10049037))
('chrUextra', (0, 29004656))
('chrX', (0, 22422827))
('chrXHet', (0, 204112))
('chrYHet', (0, 347038))
```

pybedtools.chromsizes_to_file (chromsizes, fn=None)

Converts a *chromsizes* dictionary to a file. If fn is None, then a tempfile is created (which can be deleted with pybedtools.cleanup()).

Returns the filename.

pybedtools.get_chromsizes_from_ucsc (genome, saveas=None, mysql='mysql', timeout=None)

Download chrom size info for genome from UCSC and returns the dictionary.

If you need the file, then specify a filename with *saveas* (the dictionary will still be returned as well).

If mysql is not on your path, specify where to find it with mysql=<path to mysql executable>.

timeout is how long to wait for a response; mostly used for testing.

```
>>> dm3_chromsizes = get_chromsizes_from_ucsc('dm3')
>>> for i in sorted(dm3_chromsizes.items()):
       print i
('chr2L', (0, 23011544))
('chr2LHet', (0, 368872))
('chr2R', (0, 21146708))
('chr2RHet', (0, 3288761))
('chr3L', (0, 24543557))
('chr3LHet', (0, 2555491))
('chr3R', (0, 27905053))
('chr3RHet', (0, 2517507))
('chr4', (0, 1351857))
('chrM', (0, 19517))
('chrU', (0, 10049037))
('chrUextra', (0, 29004656))
('chrX', (0, 22422827))
('chrXHet', (0, 204112))
('chrYHet', (0, 347038))
```

Setup

```
pybedtools.set_tempdir(tempdir)
```

Sets the directory for temp files. Useful for clusters that use a /scratch partition rather than a /tmp dir. Convenience function to simply set tempfile.tempdir.

```
pybedtools.get_tempdir()
```

Gets the current tempdir for the module.

```
pybedtools.set_bedtools_path(path='')
```

If BEDTools is not available on your system path, specify the path to the dir containing the BEDTools executables (intersectBed, subtractBed, etc) with this function.

To reset and use the default system path, call this function with no arguments or use path="".

Utilities

```
pybedtools.cleanup(verbose=False, remove all=False)
```

Deletes all temporary files in the BedTool.TEMPFILES class variable.

If verbose, reports what it's doing

If remove all, then ALL files matching "pybedtools.*.tmp" in the temp dir will be deleted.

```
pybedtools.IntervalIterator()
pybedtools.find_tagged(tag)
```

Returns the bedtool object with tagged with tag. Useful for tracking down bedtools you made previously.

Wrapping

```
\label{local_problem} $$ pybedtools.bedtool.\_wraps (prog=None, implicit=None, bam=None, other=None, uses\_genome=False, make\_tempfile\_for=None, check\_stderr=None, add\_to\_bedtool=None, nonbam=None, force\_bam=False) $$
```

Do-it-all wrapper, to be used as a decorator.

prog is the name of the BEDTools program that will be called. The help for this program will also be added to the decorated method's docstring.

implicit is the BEDTools program arg that should be filled in automatically.

bam will disable the implicit substitution if bam is in the kwargs. This is typically 'abam' or 'ibam' if the program accepts BAM input.

other is the BEDTools program arg that is passed in as the second input, if supported. Within the semantics of BEDTools, the typical case will be that if implicit='a' then other='b'; if implicit='i' then other=None.

uses_genome, if True, will check for 'g' and/or 'genome' args and retrieve the corresponding genome files as needed.

make_tempfile_for is used for the sequence methods and indicates which kwarg should have a tempfile made for it if it's not provided ('fo' for the sequence methods)

check_stderr, if not None, is a function that accepts a string (which will be anything written to stdout when calling the wrapped program). This function should return True if the string is OK, and False if it should truly be considered an error. This is needed for wrapping fastaFromBed, which will report to stderr that it's creating an index file.

add_to_bedtool is used for sequence methods. It is a dictionary mapping kwargs to attributes to be created in the resulting BedTool. Typically it is {'fo':'seqfn'} which will add the resulting sequence name to the BedTool's .seqfn attribute. If add_to_bedtool is not None, then the returned BedTool will be self with the added attribute.

nonbam is a kwarg that even if the input file was a BAM, the output will not be BAM format. For example, the -bed arg for intersectBed will cause the output to be in BED format, not BAM. If not None, this can be a string, a list of strings, or the special string "ALL", which means that the wrapped program will never return BAM output.

force_bam, if True, will force the output to be BAM. This is used for bedToBam.

2.6.2 BedTool methods that wrap BEDTools programs

"Genome algebra" methods

```
intersect() (wraps "intersectBed")
```

```
BedTool.intersect(*args, **kwargs)
```

Intersect with another BED file. If you want to use BAM as input, you need to specify *abam='filename.bam'*. Returns a new BedTool object.

Example usage:

Create new BedTool object

```
>>> a = pybedtools.example_bedtool('a.bed')
Get overlaps with b.bed:
>>> b = pybedtools.example_bedtool('b.bed')
>>> overlaps = a.intersect(b)
Use v=True to get the inverse - those unique to "a.bed":
>>> unique_to_a = a.intersect(b, v=True)
```

Program: intersectBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Report overlaps between two feature files.

Usage: intersectBed [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf>

Options:

-abam	The A input file is in BAM format. Output will be BAM as well.
-ubam	Write uncompressed BAM output. Default is to write compressed BAM.
-bed	When using BAM input (-abam), write output as BED. The default is to write output in BAM when using -abam.
-wa	Write the original entry in A for each overlap.
-wb	Write the original entry in B for each overlap Useful for knowing what A overlaps. Restricted by -f and -r.
-W0	Write the original A and B entries plus the number of base pairs of overlap between the two features Overlaps restricted by -f and -r.
	Only A features with overlap are reported.
-wao	Write the original A and B entries plus the number of base pairs of overlap between the two features Overlapping features restricted by -f and -r.
	However, A features w/o overlap are also reported with a NULL B feature and overlap = 0.
-u	Write the original A entry once if any overlaps found in B In other words, just report the fact >=1 hit was found Overlaps restricted by -f and -r.
-c	For each entry in A, report the number of overlaps with B Reports 0 for A entries that have no overlap with B Overlaps restricted by -f and -r.
- v	Only report those entries in A that have no overlaps with B Similar to "grep -v" (an homage).
-f	Minimum overlap required as a fraction of A Default is 1E-9 (i.e., 1bp) FLOAT (e.g. 0.50)
-r	Require that the fraction overlap be reciprocal for A and B In other words, if -f is 0.90 and -r is used, this requires
	that B overlap 90% of A and A also overlaps 90% of B.
-s	Force strandedness. That is, only report hits in B that overlap A on the same strand By default, overlaps are reported without respect to strand.
-split	Treat "split" BAM or BED12 entries as distinct BED intervals.

merge() (wraps "mergeBed")

```
BedTool.merge(*args, **kwargs)
```

Merge overlapping features together. Returns a new BedTool object.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
```

Merge:

```
>>> c = a.merge()
Allow merging of features 500 bp apart:
>>> c = a.merge(d=500)
Report number of merged features:
>>> c = a.merge(n=True)
Report names of merged features:
>>> c = a.merge(nms=True)
```

Program: mergeBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Merges overlapping BED/GFF/VCF entries into a single interval.

Usage: mergeBed [OPTIONS] -i <bed/gff/vcf>

Options:

-S	Force strandedness. That is, only merge features that are the same strand By default, merging is done without respect to strand.
-n	Report the number of BED entries that were merged Note: "1" is reported if no merging occurred.
-d	Maximum distance between features allowed for features to be merged Def. 0. That is, overlapping & book-ended features are merged (INTEGER)
-nms	Report the names of the merged features separated by semicolons.

-scores [STRING] Report the scores of the merged features. Specify one of

the following options for reporting scores: sum, min, max, mean, median, mode, antimode, collapse (i.e., print a semicolon-separated list),

```
subtract() (wraps "subtractBed")
```

```
BedTool.subtract(*args, **kwargs)
```

Subtracts from another BED file and returns a new BedTool object.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
Do a "stranded" subtraction:
>>> c = a.subtract(b, s=True)
Require 50% of features in a to overlap:
>>> c = a.subtract(b, f=0.5)
```

Program: subtractBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Removes the portion(s) of an interval that is overlapped

by another feature(s).

Usage: subtractBed [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf>

Options:

-f Minimum overlap required as a fraction of A. - Default is 1E-9 (i.e.,

1bp). - (FLOAT) (e.g. 0.50)

-s Force strandedness. That is, only report hits in B that overlap A on the same strand. - By default, overlaps are reported without respect

to strand.

closest() (wraps "closestBed")

```
BedTool.closest(*args, **kwargs)
```

Return a new BedTool object containing closest features in b. Note that the resulting file is no longer a valid BED format; use the special "_closest" methods to work with the resulting file.

Example usage:

```
a = BedTool('in.bed')
# get the closest feature in 'other.bed' on the same strand
b = a.closest('other.bed', s=True)
```

Program: closestBed (v2.11.2) Authors: Aaron Quinlan (aaronquinlan@gmail.com)

Erik Arner, Riken

Summary: For each feature in A, finds the closest feature (upstream or downstream) in B.

Usage: closestBed [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf>

Options:

-s Force strandedness. That is, find the closest feature in B that overlaps A on the same strand. - By default, overlaps are reported without respect to strand.

-d In addition to the closest feature in B, report its distance to A as an extra column. - The reported distance for overlapping features will

be 0.

How ties for closest feature are handled. This occurs when two features in B have exactly the same overlap with A. By default, all such features in B are reported. Here are all the options: - "all" Report all ties (default). - "first" Report the first tie that occurred in the B file. - "last" Report the last tie that occurred in the B file.

Notes: Reports "none" for chrom and "-1" for all other fields when a feature is not found in B on the same chromosome as the feature in A. E.g. none -1 -1

window() (wraps "windowBed")

BedTool.window(*args, **kwargs)

Intersect with a window.

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> print a.window(b, w=1000)
chr1
         1
                 100
                         feature1
                                                      chr1
                                                             155
                                                                    200
                                                                            feature5
          1
                                      0
chr1
                 100
                         feature1
                                                      chr1
                                                             800
                                                                    901
                                                                            feature6
                                      0
chr1
          100
                  200
                         feature2
                                                      chr1
                                                             155
                                                                    200
                                                                            feature5
chr1
          100
                 200
                         feature2
                                      0
                                      0
                                                             800
                                                                    901
                                                      chr1
                                                                            feature6
                        feature3
feature3
feature4
          150
chr1
                500
                                                      chr1
                                                             155
                                                                    200
                                                                           feature5
                500
          150
                                      0
                                                             800
                                                      chr1
                                                                    901
chr1
                                                                           feature6
                                      0
          900
                950
                                                             155
                                                                    200
chr1
                                                      chr1
                                                                           feature5
                                      0
          900
                 950
                                                             800
                                                                    901
chr1
                        feature4
                                                      chr1
                                                                            feature6
<BLANKLINE>
```

Program: windowBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Examines a "window" around each feature in A and

reports all features in B that overlap the window. For each overlap the entire entry in A and B are reported.

Usage: windowBed [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf>

Options:

-abam	The A input file is in BAM format. Output will be BAM as well.
-ubam	Write uncompressed BAM output. Default is to write compressed BAM.
-bed	When using BAM input (-abam), write output as BED. The default is to write output in BAM when using -abam.
-w	Base pairs added upstream and downstream of each entry in A when searching for overlaps in B Creates symterical "windows" around A Default is 1000 bp (INTEGER)
-1	Base pairs added upstream (left of) of each entry in A when searching for overlaps in B Allows one to define assymterical "windows" Default is 1000 bp (INTEGER)
-r	Base pairs added downstream (right of) of each entry in A when searching for overlaps in B Allows one to define assymterical "windows" Default is 1000 bp (INTEGER)
-sw	Define -1 and -r based on strand. For example if used, -1 500 for a negative-stranded feature will add 500 bp downstream Default = disabled.
-sm	Only report hits in B that overlap A on the same strand By default, overlaps are reported without respect to strand.
-u	Write the original A entry once if any overlaps found in B In other words, just report the fact >=1 hit was found.
-c	For each entry in A, report the number of overlaps with B Reports 0 for A entries that have no overlap with B Overlaps restricted by -f.
-v	Only report those entries in A that have no overlaps with B Similar to "grep -v."

sort() (wraps "sortBed")

```
BedTool.sort(*args, **kwargs)
```

Note that chromosomes are sorted lexograpically, so chr12 will come before chr9.

Example usage:

```
>>> a = pybedtools.BedTool('''
... chr9 300 400
... chr1 100 200
... chr1 1 50
... chr12 1 100
... chr9 500 600
... ''', from_string=True)
>>> print a.sort()
chr1
       1
       100
               200
chr1
chr12 1
               100
      300
               400
chr9
               600
chr9
     500
<BLANKLINE>
```

Program: sortBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Sorts a feature file in various and useful ways.

Usage: sortBed [OPTIONS] -i <bed/gff/vcf>

Options:

-sizeA	Sort by feature size in ascending order.
-sizeD	Sort by feature size in descending order.
-chrThenSizeA	Sort by chrom (asc), then feature size (asc).
-chrThenSizeD	Sort by chrom (asc), then feature size (desc).
-chrThenScoreA	Sort by chrom (asc), then score (asc).
-chrThenScoreD	Sort by chrom (asc), then score (desc).

slop() (wraps "slopBed")

```
BedTool.slop(*args, **kwargs)
```

Wraps slopBed, which adds bp to each feature. Returns a new BedTool object.

If g is a dictionary (for example, return values from pybedtools.chromsizes()) it will be converted to a temp file for use with slopBed. If it is a string, then it is assumed to be a filename.

Alternatively, use genome to indicate a pybedtools-created genome. Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
```

Increase the size of features by 100 bp in either direction. Note that you need to specify either a dictionary of chromsizes or a filename containing chromsizes for the genome that your bed file corresponds to:

```
>>> c = a.slop(g=pybedtools.chromsizes('hg19'), b=100)
```

Grow features by 10 bp upstream and 500 bp downstream, using a genome file you already have constructed called 'hg19.genome'

First, create the file:

```
>>> fout = open('hg19.genome','w')
>>> chromdict = pybedtools.get_chromsizes_from_ucsc('hg19')
>>> for chrom, size in chromdict.items():
...    fout.write("%s\t%s\n" % (chrom, size[1]))
>>> fout.close()

Then use it:
>>> c = a.slop(g='hg19.genome', l=10, r=500, s=True)

Clean up afterwards:
>>> os.unlink('hg19.genome')
```

Program: slopBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Add requested base pairs of "slop" to each feature.

Usage: slopBed [OPTIONS] -i <bed/gff/vcf> -g <genome> [-b <int> or (-l and -r)]

Options:

-b	Increase the BED/GFF/VCF entry by -b base pairs in each direction (Integer) or (Float, e.g. 0.1) if used with -pct.
-1	The number of base pairs to subtract from the start coordinate (Integer) or (Float, e.g. 0.1) if used with -pct.
-r	The number of base pairs to add to the end coordinate (Integer) or (Float, e.g. 0.1) if used with -pct.
-S	Define -l and -r based on strand. E.g. if used, -l 500 for a negative-stranded feature, it will add 500 bp downstream. Default = false.
-pct	Define -l and -r as a fraction of the feature's length. E.g. if used on a 1000bp feature, -l 0.50, will add 500 bp "upstream". Default = false.

Notes:

- 1. Starts will be set to 0 if options would force it below 0.
- (2) Ends will be set to the chromosome length if requested slop would force it above the max chrom length.
- (3) The genome file should tab delimited and structured as follows:

```
<chromName><TAB><chromSize>
```

For example, Human (hg19): chr1 249250621 chr2 243199373 ... chr18**gl000207**random 4262

Tips: One can use the UCSC Genome Browser's MySQL database to extract chromosome sizes. For example, H. sapiens:

mysql –user=genome –host=genome-mysql.cse.ucsc.edu -A -e / "select chrom, size from hg19.chromInfo" > hg19.genome

complementBed() (wraps "complementBed")

```
BedTool.complement(*args, **kwargs)
```

Program: complementBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Returns the base pair complement of a feature file.

Usage: complementBed [OPTIONS] -i <bed/gff/vcf> -g <genome>

Notes:

1. The genome file should tab delimited and structured as follows: <chromName><TAB><chromSize> For example, Human (hg19): chr1 249250621 chr2 243199373 ... chr18**gl000207**random 4262

Tips: One can use the UCSC Genome Browser's MySQL database to extract chromosome sizes. For example, H. sapiens:

mysql –user=genome –host=genome-mysql.cse.ucsc.edu -A -e / "select chrom, size from hg19.chromInfo" > hg19.genome

flank() (wraps "flankBed")

BedTool.flank(*args, **kwargs)

Create flanking intervals on either side of input BED.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> print a.flank(genome='hg19', b=100)
chr1
     0
             1
                   feature1
      100
             200
                                    0
chr1
                    feature1
     0
             100 feature2
chr1
                                    0
      200
chr1
              300
                     feature2
                                    0
      150
500 600
800 900
950 1050
INE>
chr1
                     feature3
                                    0
                                    0
chr1
                     feature3
chr1
                    feature4
                                    0
             1050
chr1
                     feature4
<BLANKLINE>
```

Program: flankBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Creates flanking interval(s) for each BED/GFF/VCF feature.

Usage: flankBed [OPTIONS] -i <bed/gff/vcf> -g <genome> [-b <int> or (-l and -r)]

Options:

-b	Create flanking intervak using -b base pairs in each direction (Integer) or (Float, e.g. 0.1) if used with -pct.
-1	The number of base pairs that a flank should start from orig. start coordinate (Integer) or (Float, e.g. 0.1) if used with -pct.
-r	The number of base pairs that a flank should end from orig. end coordinate (Integer) or (Float, e.g. 0.1) if used with -pct.

-S	Define -l and -r based on strand. E.g. if used, -l 500 for a negative-stranded feature, it will start the flank 500 bp downstream. Default = false.
-pct	Define -l and -r as a fraction of the feature's length. E.g. if used on a 1000bp feature, -l 0.50, will add 500 bp "upstream". Default = false.

Notes:

- 1. Starts will be set to 0 if options would force it below 0.
- (2) Ends will be set to the chromosome length if requested flank would force it above the max chrom length. (3) The genome file should tab delimited and structured as follows:

```
<chromName><TAB><chromSize>
```

For example, Human (hg19): chr1 249250621 chr2 243199373 ... chr18**gl000207**random 4262

Tips: One can use the UCSC Genome Browser's MySQL database to extract chromosome sizes. For example, H. sapiens:

mysql –user=genome –host=genome-mysql.cse.ucsc.edu -A -e / "select chrom, size from hg19.chromInfo" > hg19.genome

shuffle() (wraps "shuffleBed")

```
BedTool.shuffle(*args, **kwargs)
```

Shuffle coordinates.

Example usage:

Program: shuffleBed (v2.12.0) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Randomly permute the locations of a feature file among a genome.

Usage: shuffleBed [OPTIONS] -i <bed/gff/vcf> -g <genome>

Options:

-excl	A BED/GFF/VCF file of coordinates in which features in -i should not be placed (e.g. gaps.bed).
-incl	Instead of randomly placing features in a genome, the -incl options defines a BED/GFF/VCF file of coordinates in which features in -i should be randomly placed (e.g. genes.bed).
-chrom	Keep features in -i on the same chromosome By default, the chrom and position are randomly chosen.
-seed	Supply an integer seed for the shuffling By default, the seed is chosen automatically (INTEGER)

-f

Maximum overlap (as a fraction of the -i feature) with an -excl feature that is tolerated before searching for a new, randomized locus. For example, -f 0.10 allows up to 10% of a randomized feature to overlap with a given feature in the -excl file. **Cannot be used with -incl file.** - Default is 1E-9 (i.e., 1bp). - FLOAT (e.g. 0.50)

Notes:

1. The genome file should tab delimited and structured as follows: <chromName><TAB><chromSize> For example, Human (hg19): chr1 249250621 chr2 243199373 ... chr18**gl000207**random 4262

Tips: One can use the UCSC Genome Browser's MySQL database to extract chromosome sizes. For example, H. sapiens:

mysql –user=genome –host=genome-mysql.cse.ucsc.edu -A -e / "select chrom, size from hg19.chromInfo" > hg19.genome

annotate() (wraps "annotateBed")

BedTool.annotate(*args, **kwargs)

Annotate this BedTool with a list of other files. Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b_fn = pybedtools.example_filename('b.bed')
>>> print a.annotate(files=b_fn)
                                         0
chr1
       1
                100
                                                          0.000000
                        feature1
                200
                                         0
        100
                                                          0.450000
chr1
                        feature2
                500
                                         0
        150
                        feature3
                                                          0.128571
chr1
                950
                                         \cap
chr1
        900
                        feature4
                                                          0.020000
<BLANKLINE>
```

Program: annotateBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Annotates the depth & breadth of coverage of features from multiple files

on the intervals in -i.

Usage: annotateBed [OPTIONS] -i <bed/gff/vcf> -files FILE1 FILE2 .. FILEn

Options:

-names

A list of names (one / file) to describe each file in -i. These names will be printed as a header line.

-counts Report the count of features in each file that overlap -i.

• Default is to report the fraction of -i covered by each file.

-both Report the counts followed by the % coverage. - Default is to report the fraction of -i covered by each file.

-s Force strandedness. That is, only include hits in A that overlap B on the same strand. - By default, hits are included without respect to

strand.

coverage() (wraps "coverageBed")

```
BedTool.coverage(*args, **kwargs)
```

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> c = a.coverage(b)
>>> c.head(3)
                200
                                         0
      155
                         feature5
                                                                  45
                                                                           45
                                                                                   1.0000000
chr1
        800
                901
                        feature6
                                         0
                                                          1
                                                                  1
                                                                           101
                                                                                   0.0099010
```

Program: coverageBed (v2.12.0) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Returns the depth and breadth of coverage of features from A

on the intervals in B.

Usage: coverageBed [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf>

Options:

-abam	The A input file is in BAM format.
-s	Force strandedness. That is, only include hits in A that overlap B on the same strand By default, hits are included without respect to strand.
-hist	Report a histogram of coverage for each feature in B as well as a summary histogram for all features in B.
	Output (tab delimited) after each feature in B:
	1. depth
	2. # bases at depth
	3. size of B
	4. % of B at depth
-d	Report the depth at each position in each B feature. Positions reported are one based. Each position and depth follow the complete B feature.
-split	Treat "split" BAM or BED12 entries as distinct BED intervals. when computing coverage. For BAM files, this uses the CIGAR "N" and "D" operations to infer the blocks for computing coverage. For

BED12 files, this uses the BlockCount, BlockStarts, and BlockEnds

Default Output:

After each entry in B, reports:

1. The number of features in A that overlapped the B interval.

fields (i.e., columns 10,11,12).

- 2. The number of bases in B that had non-zero coverage.
- 3. The length of the entry in B.
- 4. The fraction of bases in B that had non-zero coverage.

genome_coverage() (wraps "genomeCoverageBed")

```
BedTool.genome_coverage(*args, **kwargs)
```

Calculates coverage at each position in the genome.

Use bg=True to have the resulting BedTool return valid BED-like features

```
>>> a = pybedtools.example_bedtool('x.bam')
>>> b = a.genome_coverage(genome='dm3', bg=True)
>>> b.head(3)
chr2L 9329 9365 1
chr2L 10212 10248 1
chr2L 10255 10291 1
```

Program: genomeCoverageBed (v2.12.0) Authors: Aaron Quinlan (aaronquinlan@gmail.com)

Assaf Gordon, CSHL

Summary: Compute the coverage of a feature file among a genome.

Usage: genomeCoverageBed [OPTIONS] -i <bed/gff/vcf> -g <genome>

Options:

-ibam	The input file is in BAM format. Note: BAM must be sorted by position
-d	Report the depth at each genome position. Default behavior is to report a histogram.
-bg	Report depth in BedGraph format. For details, see: genome.ucsc.edu/goldenPath/help/bedgraph.html
-bga	Report depth in BedGraph format, as above (-bg). However with this option, regions with zero coverage are also reported. This allows one to quickly extract all regions of a genome with 0 coverage by applying: "grep -w 0\$" to the output.
-split	Treat "split" BAM or BED12 entries as distinct BED intervals. when computing coverage. For BAM files, this uses the CIGAR "N" and "D" operations to infer the blocks for computing coverage. For BED12 files, this uses the BlockCount, BlockStarts, and BlockEnds fields (i.e., columns 10,11,12).

-strand Calculate coverage of intervals from a specific strand. With BED files, requires at least 6 columns (strand is column 6). - (STRING): can be + or -

-max Combine all positions with a depth >= max into a single bin in the histogram. Irrelevant for -d and -bedGraph - (INTEGER)

Notes:

- 1. The genome file should tab delimited and structured as follows: <chromName><TAB><chromSize> For example, Human (hg19): chr1 249250621 chr2 243199373 ... chr18**gl000207**random 4262
- 2. The input BED (-i) file must be grouped by chromosome. A simple "sort -k 1,1 <BED> > <BED>.sorted" will suffice.
- 3. The input BAM (-ibam) file must be sorted by position. A "samtools sort <BAM>" should suffice.

Tips: One can use the UCSC Genome Browser's MySQL database to extract chromosome sizes. For example, H. sapiens:

mysql –user=genome –host=genome-mysql.cse.ucsc.edu -A -e / "select chrom, size from hg19.chromInfo" > hg19.genome

overlap() (wraps "overlap") BedTool.overlap(*args, **kwargs) >>> a = pybedtools.example_bedtool('a.bed') >>> b = pybedtools.example_bedtool('b.bed') >>> c = a.window(b, w=10).overlap(cols=[2,3,8,9])>>> print c 100 200 feature2 150 500 feature3 0 155 200 chr1 100 chr1 feature5 0 155 200 chr1 chr1 feature5 950 chr1 900 feature4 0 chr1 800 901 feature6 <BLANKLINE>

Program: overlap (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Computes the amount of overlap (positive values)

or distance (negative values) between genome features and reports the result at the end of the same line.

Usage: overlap [OPTIONS] -i <input> -cols s1,e1,s2,e2

Options:

-i Input file. Use "stdin" for pipes.

-cols Specify the columns (1-based) for the starts and ends of the features for which you'd like to compute the overlap/distance. The columns

must be listed in the following order:

start1,end1,start2,end2

Example: \$ windowBed -a A.bed -b B.bed -w 10 chr1 10 20 A chr1 15 25 B chr1 10 20 C chr1 25 35 D

\$ windowBed -a A.bed -b B.bed -w 10 | overlap -i stdin -cols 2,3,6,7 chr1 10 20 A chr1 15 25 B 5 chr1 10 20 C chr1 25 35 D -5

groupby() (wraps "groupBy")

BedTool.groupby(*args, **kwargs)

```
>>> a = pybedtools.example_bedtool('gdc.gff')
>>> b = pybedtools.example_bedtool('gdc.bed')
>>> c = a.intersect(b, c=True)
>>> d = c.groupby(g=[1, 4, 5], c=10, ops=['sum'])
>>> print d
              70
                     0
chr2L 41
             130
chr2L 71
                     2
chr2L 71
chr2L 131
             170
                     4
chr2L 171
              200
chr2L 201
              220
                     1
chr2L 41
              130
chr2L 171
              220
                     1
     41
              220
                     7
chr2L
             230
chr2L
       161
                     6
chr2L
      41
              220
                     7
<BLANKLINE>
```

Program: groupBy (v1.1.0) Authors: Aaron Quinlan (aaronquinlan@gmail.com)

Assaf Gordon

Summary: Summarizes a dataset column based upon common column groupings. Akin to the SQL "group by" command.

Usage: groupBy -i [FILE] -g [group**column(s)] -c [op**column(s)] -o [ops] cat [FILE] | groupBy -g [group**column(s)] -c [op**column(s)] -o [ops]

Options:

- -i Input file. Assumes "stdin" if omitted.
- **-g -grp Specify the columns (1-based) for the grouping.** The columns must be comma separated. Default: 1,2,3
- -c -opCols Specify the column (1-based) that should be summarized.
 - · Required.
- -o -ops Specify the operation that should be applied to opCol.

Valid operations: sum, count, min, max, mean, median, mode, antimode, stdev, sstdev (sample standard dev.), collapse (i.e., print a comma separated list), freqdesc (i.e., print desc. list of values:freq) freqasc (i.e., print asc. list of values:freq)

• Default: sum

-full Print all columns from input file. Default: print only grouped

columns.

-inheader Input file has a header line - the first line will be ignored.

-outheader Print header line in the output, detailing the column names. If the

input file has headers (-inheader), the output file will use the input's column names. If the input file has no headers, the output file will use

"col**1", "col**2", etc. as the column names.

-header same as '-inheader -outheader'

-ignorecase Group values regardless of upper/lower case.

Examples: \$ cat ex1.out chr1 10 20 A chr1 15 25 B.1 1000 chr1 10 20 A chr1 25 35 B.2 10000

\$ groupBy -i ex1.out -g 1,2,3,4 -c 9 -o sum chr1 10 20 A 11000

\$ groupBy -i ex1.out -grp 1,2,3,4 -opCols 9,9 -ops sum,max chr1 10 20 A 11000 10000

\$ groupBy -i ex1.out -g 1,2,3,4 -c 8,9 -o collapse, mean chr1 10 20 A B.1,B.2, 5500

\$ cat ex1.out | groupBy -g 1,2,3,4 -c 8,9 -o collapse,mean chr1 10 20 A B.1,B.2, 5500

Notes:

- 1. The input file/stream should be sorted/grouped by the -grp. columns
- 2. If -i is unspecified, input is assumed to come from stdin.

pair_to_bed() (wraps "pairToBed")

BedTool.pair_to_bed(*args, **kwargs)

Program: pairToBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Report overlaps between a BEDPE file and a BED/GFF/VCF file.

Usage: pairToBed [OPTIONS] -a <bedpe> -b <bed/gff/vcf>

Options:

-abam The A input file is in BAM format. Output will be BAM as well. -

Requires BAM to be grouped or sorted by query.

-ubam Write uncompressed BAM output. Default is to write compressed

BAM.

is to write output in BAM when using -abam.

-bedpe When using BAM input (-abam), write output as BEDPE. The default

is to write output in BAM when using -abam.

-ed Use BAM total edit distance (NM tag) for BEDPE score. - Default

for BEDPE is to use the minimum of

of the two mapping qualities for the pair.

• When -ed is used the total edit distance from the two mates is

reported as the score.

-f Minimum overlap required as fraction of A (e.g. 0.05). Default is

1E-9 (effectively 1bp).

s Enforce strandedness when finding overlaps. Default is to ignore

stand. Not applicable with -type inspan or -type outspan.

-type Approach to reporting overlaps between BEDPE and BED.

either Report overlaps if either end of A overlaps B.

• Default.

neither Report A if neither end of A overlaps B. both Report overlaps if both ends of A overlap B. xor Report overlaps if one and only one end of A overlaps B. notboth Report overlaps if neither end or one and only one

end of A overlap B. That is, xor + neither.

ispan Report overlaps between [end1, start2] of A and B.

• Note: If chrom1 <> chrom2, entry is ignored.

ospan Report overlaps between [start1, end2] of A and B.

• Note: If chrom1 <> chrom2, entry is ignored.

notispan Report A if ispan of A doesn't overlap B.

• Note: If chrom1 <> chrom2, entry is ignored.

notospan Report A if ospan of A doesn't overlap B.

• Note: If chrom1 <> chrom2, entry is ignored.

Refer to the BEDTools manual for BEDPE format.

pair_to_pair() (wraps "pairToPair")

BedTool.pair_to_pair(*args, **kwargs)

Program: pairToPair (v2.12.0) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Report overlaps between two paired-end BED files (BEDPE).

Usage: pairToPair [OPTIONS] -a <BEDPE> -b <BEDPE>

Options:

-f Minimum overlap required as fraction of A (e.g. 0.05). Default is

1E-9 (effectively 1bp).

-type Approach to reporting overlaps between A and B.

neither Report overlaps if neither end of A overlaps B. either Report overlaps if either ends of A overlap B. both Report overlaps if both ends of A overlap B. notboth Report overlaps if one or neither of ends

of A overlap B. - Default = both.

-slop The amount of slop (in b.p.). to be added to each footprint. *Note*:

Slop is subtracted from start1 and start2 and added to end1 and end2.

-ss Add slop based to each BEDPE footprint based on strand. - If strand

is "+", slop is only added to the end coordinates. - If strand is "-", slop is only added to the start coordinates. - By default, slop is added

in both directions.

-is Ignore strands when searching for overlaps. - By default, strands are

enforced.

-rdn Require the hits to have different names (i.e. avoid self-hits). - By

default, same names are allowed.

Refer to the BEDTools manual for BEDPE format.

Methods for converting between formats

bed6() (wraps "Bed12To6")

BedTool.bed6(*args, **kwargs)

convert a BED12 to a BED6 file

Program: bed12ToBed6 (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Splits BED12 features into discrete BED6 features.

Usage: bed12ToBed6 [OPTIONS] -i <bed12>

Options:

-n Force the score to be the (1-based) block number from the BED12.

Methods for working with sequences

sequence() (wraps "fastaFromBed")

```
BedTool.sequence(*args, **kwargs)
```

Wraps fastaFromBed. fi is passed in by the user; bed is automatically passed in as the bedfile of this object; fo by default is a temp file. Use save_seqs() to save as a file.

The end result is that this BedTool will have an attribute, self.seqfn, that points to the new fasta file.

Example usage:

```
>>> a = pybedtools.BedTool("""
... chr1 1 10
... chr1 50 55""", from_string=True)
>>> fasta = pybedtools.example_filename('test.fa')
>>> a = a.sequence(fi=fasta)
>>> print open(a.seqfn).read()
>chr1:1-10
GATGAGTCT
>chr1:50-55
CCATC
<BLANKLINE>
```

Program: fastaFromBed (v2.12.0) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Extract DNA sequences into a fasta file based on feature coordinates.

Usage: fastaFromBed [OPTIONS] -fi <fasta> -bed <bed/gff/vcf> -fo <fasta>

Options:

-fi	Input FASTA file
-bed	BED/GFF/VCF file of ranges to extract from -fi
-fo	Output file (can be FASTA or TAB-delimited)
-name	Use the name field for the FASTA header
-tab	Write output in TAB delimited format Default is FASTA format.
-S	Force strandedness. If the feature occupies the antisense strand, the sequence will be reverse complemented By default, strand information is ignored.

mask_fasta() (wraps "maskFastaFromBed")

```
BedTool.mask_fasta(*args, **kwargs)
```

Masks a fasta file at the positions in a BED file and saves result as 'out' and stores the filename in seqfn.

```
>>> a = pybedtools.BedTool('chr1 100 110', from_string=True)
>>> fasta_fn = pybedtools.example_filename('test.fa')
>>> a = a.mask_fasta(fi=fasta_fn, fo='masked.fa.example')
>>> b = a.slop(b=2, genome='hg19')
>>> b = b.sequence(fi=a.seqfn)
>>> print open(b.seqfn).read()
>chr1:98-112
TTNNNNNNNNNAT

<BLANKLINE>
>>> os.unlink('masked.fa.example')
>>> if os.path.exists('masked.fa.example.fai'):
... os.unlink('masked.fa.example.fai')
```

Program: maskFastaFromBed (v2.11.2) Author: Aaron Quinlan (aaronquinlan@gmail.com) Summary: Mask a fasta file based on feature coordinates.

Usage: maskFastaFromBed [OPTIONS] -fi <fasta> -out <fasta> -bed <bed/gff/vcf>

Options:

-fi	Input FASTA file
-bed	BED/GFF/VCF file of ranges to mask in -fi
-fo	Output FASTA file
-soft	Enforce "soft" masking. That is, instead of masking with Ns, mask with lower-case bases.
-mc	Replace masking character. That is, instead of masking with Ns, use another character.

2.6.3 BedTool methods unique to pybedtools

Introspection

count()

```
BedTool.count()
```

Number of features in BED file. Does the same thing as len(self), which actually just calls this method.

Only counts the actual features. Ignores any track lines, browser lines, lines starting with a "#", or blank lines.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> a.count()
4
```

print_sequence()

```
BedTool.print_sequence()
```

Print the sequence that was retrieved by the BedTool.sequence() method.

See usage example in BedTool.sequence().

field_count()

```
BedTool.field_count (n=10)
```

Return the number of fields in the features this file contains. Checks the first n features.

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> a.field_count()
6
```

head()

BedTool.head(n=10)

Prints the first *n* lines

Saving

saveas()

```
BedTool.saveas(*args, **kwargs)
```

Save BED file as a new file, adding the optional trackline to the beginning.

Returns a new BedTool for the newly saved file.

A newline is automatically added to the trackline if it does not already have one.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = a.saveas('other.bed')
>>> b.fn
'other.bed'
>>> print b == a
True

>>> b = a.saveas('other.bed', trackline="name='test run' color=0,55,0")
>>> open(b.fn).readline()
    "name='test run' color=0,55,0\n"
save_seqs()
```

BedTool.save_seqs(fn)

Save sequences of features in this BedTool object as a fasta file fn.

In order to use this function, you need to have called the <code>BedTool.sequence()</code> method.

A new BedTool object is returned which references the newly saved file.

Example usage:

```
>>> a = pybedtools.BedTool('''
... chr1 1 10
... chr1 50 55''', from_string=True)
>>> fasta = pybedtools.example_filename('test.fa')
>>> a = a.sequence(fi=fasta)
>>> print open(a.seqfn).read()
>chr1:1-10
GATGAGTCT
>chr1:50-55
CCATC
<BLANKLINE>
>>> b = a.save_seqs('example.fa')
>>> assert open(b.fn).read() == open(a.fn).read()
```

Utilities

```
with_attrs()
```

```
BedTool.with_attrs(*args, **kwargs)
```

Given arbitrary keyword arguments, turns the keys and values into attributes. Useful for labeling BedTools at creation time.

.with_attrs(label=

.with_attrs(label=

Example usage:

```
>>> # add a "label" attribute to each BedTool
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> for i in [a, b]:
... print i.count(), 'features for', i.label
4 features for transcription factor 1
2 features for transcription factor 2
```

cat()

BedTool.cat(*args, **kwargs)

Concatenates two BedTool objects (or an object and a file) and does an optional post-merge of the features.

Use *postmerge=False* if you want to keep features separate.

TODO:

currently truncates at BED3 format!

kwargs are sent to BedTool.merge().

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> b = pybedtools.example_bedtool('b.bed')
>>> print a.cat(b)
chr1 1 500
chr1 800 950
<BLANKLINE>
```

total_coverage()

BedTool.total_coverage()

Returns the total number of bases covered by this BED file. Does a self.merge() first to remove potentially multiple-counting bases.

Example usage:

```
>>> a = pybedtools.example_bedtool('a.bed')
```

This does a merge() first, so this is what the total coverage is counting:

delete_temporary_history()

BedTool.delete_temporary_history(ask=True, raw_input_func=None)

Use at your own risk! This method will delete temp files. You will be prompted for deletion of files unless you specify *ask=False*.

Deletes all temporary files created during the history of this BedTool up to but not including the file this current BedTool points to.

Any filenames that are in the history and have the following pattern will be deleted:

```
<TEMP_DIR>/pybedtools.*.tmp
```

(where <TEMP_DIR> is the result from get_tempdir() and is by default "/tmp")

Any files that don't have this format will be left alone.

(raw_input_func is used for testing)

as_intervalfile()

```
BedTool.as_intervalfile()
```

Returns an IntervalFile of this BedTool, which provides a low-level interface

Feature-by-feature operations

each()

```
BedTool.each (func, *args, **kwargs)
```

Applies user-defined function *func* to each feature. *func* must accept an Interval as its first argument; *args and* **kwargs will be passed to *func.

func must return an Interval object.

filter()

```
BedTool.filter(func, *args, **kwargs)
```

Takes a function *func* that is called for each feature in the BedTool object and returns only those for which the function returns True.

args and **kwargs are passed directly to *func.

Returns a streaming BedTool; if you want the filename then use the .saveas() method.

```
>>> a = pybedtools.example_bedtool('a.bed')
>>> subset = a.filter(lambda b: b.chrom == 'chr1' and b.start < 150)
>>> len(a), len(subset)
(4, 2)
```

so it has extracted 2 records from the original 4.

cut()

```
BedTool.cut (indexes)
```

Similar to unix cut except indexes are 0-based, must be a list and the columns are returned in the order requested.

In addition, indexes can contain keys of the GFF/GTF attributes, in which case the values are returned. e.g. 'gene_name' will return the corresponding name from a GTF, or 'start' will return the start attribute of a BED Interval.

See .with_column() if you need to do more complex operations.

features()

```
BedTool.features()
```

Returns an iterator of feature objects.

Randomization helpers

randomintersection()

BedTool.randomintersection (other, iterations, intersect_kwargs=None, shuffle_kwargs=None, debug=False)

Performs iterations shufflings of self, each time intersecting with other.

Returns a generator of integers where each integer is the number of intersections of a shuffled file with *other*. This distribution can be used in downstream analysis for things like empirical p-values.

intersect_kwargs and *shuffle_kwargs* are passed to self.intersect() and self.shuffle() respectively. By default for intersect, u=True is specified – but s=True might be a useful option for strand-specific work.

Useful kwargs for *shuffle_kwargs* are chrom, excl, or incl. If you use the "seed" kwarg, that seed will be used *each* time shuffleBed is called – so all your randomization results will be identical for each iteration. To get around this and to allow for tests, debug=True will set the seed to the iteration number.

```
>>> chromsizes = {'chr1':(0, 1000)}
>>> a = pybedtools.example_bedtool('a.bed')
>>> a = a.set_chromsizes(chromsizes)
>>> b = pybedtools.example_bedtool('b.bed')
>>> results = a.randomintersection(b, 10, debug=True)
>>> print list(results)
[2, 2, 2, 0, 2, 3, 2, 1, 2, 3]
```

randomstats()

```
BedTool.randomstats (other, iterations, **kwargs)
```

Sends args and kwargs to BedTool.randomintersection() and compiles results into a dictionary with useful stats. Requires scipy and numpy.

This is one possible way of assigning significance to overlaps between two files. See, for example:

Negre N, Brown CD, Shah PK, Kheradpour P, Morrison CA, et al. 2010 A Comprehensive Map of Insulator Elements for the Drosophila Genome. PLoS Genet 6(1): e1000814. doi:10.1371/journal.pgen.1000814

Example usage:

Make chromsizes a very small genome for this example:

```
>>> chromsizes = {'chr1':(1,1000)}
>>> a = pybedtools.example_bedtool('a.bed').set_chromsizes(chromsizes)
>>> b = pybedtools.example_bedtool('b.bed')
>>> results = a.randomstats(b, 100, debug=True)
```

results is a dictionary that you can inspect. The actual overlap:

```
>>> print results['actual']
3
```

The median of all randomized overlaps:

```
>>> print results['median randomized']
2.0
```

The percentile of the actual overlap in the distribution of randomized overlaps, which can be used to get an empirical p-value:

```
>>> print results['percentile']
90.0
```

Interval class

class pybedtools.Interval

Constructor:

```
Interval(chrom, start, end, name=".", score=".", strand=".", otherfields=None)
```

Class to represent a genomic interval of any format. Requires at least 3 args: chrom (string), start (int), end (int).

start is *always* the 0-based start coordinate. If this Interval is to represent a GFF object (which uses a 1-based coordinate system), then subtract 1 from the 4th item in the line to get the start position in 0-based coords for this Interval. The 1-based GFF coord will still be available, albeit as a string, in fields[3].

otherfields is a list of fields that don't fit into the other kwargs, and will be stored in the fields attribute of the Interval.

All the items in otherfields must be strings for proper conversion to C++.

By convention, for BED files, otherfields is everything past the first 6 items in the line. This allows an Interval to represent composite features (e.g., a GFF line concatenated to the end of a BED line)

But for other formats (VCF, GFF, SAM), the entire line should be passed in as a list for otherfields so that we can always check the Interval.file_type and extract the fields we want, knowing that they'll be in the right order as passed in with otherfields.

```
>>> from pybedtools import Interval
     >>> i = Interval("chr1", 22, 44, strand='-')
     Interval(chr1:22-44)
     >>> i.start, i.end, i.strand, i.length
     (22L, 44L, '-', 22L)
     chrom
         the chromosome of the feature
     end
         The end of the feature
     file_type
         bed/vcf/gff
     length
         the length of the feature
     name
         >>> import pybedtools
         >>> vcf = pybedtools.example_bedtool('v.vcf')
         >>> [v.name for v in vcf]
         ['rs6054257', 'chr1:16', 'rs6040355', 'chr1:222', 'microsat1']
     start
         The 0-based start of the feature.
     stop
         the end of the feature
     strand
         the strand of the feature
IntervalFile class
class pybedtools. IntervalFile
     all hits
             Signature IntervalFile.all_hits(interval, same_strand=False,
                 overlap=0.0)
         Search for the Interval interval this file and return all overlaps as a list.
         same_strand, if True, will only consider hits on the same strand as interval.
         overlap can be used to specify the fraction of overlap between interval and each feature in the
         IntervalFile.
         Example usage:
         >>> fn = pybedtools.example_filename('a.bed')
```

```
>>> # create an Interval to query with
>>> i = pybedtools.Interval('chr1', 1, 10000, strand='+')

>>> # Create an IntervalFile out of a.bed
>>> intervalfile = pybedtools.IntervalFile(fn)

>>> # get stranded hits
>>> intervalfile.all_hits(i, same_strand=True)
[Interval(chr1:1-100), Interval(chr1:100-200), Interval(chr1:900-950)]

any_hits

Signature IntervalFile.any_hits(interval, same_strand=False, overlap=0.0)

Return 1 if the Interval interval had >=1 hit in this IntervalFile, 0 otherwise.
```

same_strand, if True, will only consider hits on the same strand as interval.

overlap can be used to specify the fraction of overlap between interval and each feature in the IntervalFile.

Example usage:

```
>>> fn = pybedtools.example_filename('a.bed')
>>> # create an Interval to query with
>>> i = pybedtools.Interval('chr1', 1, 10000, strand='+')
>>> # Create an IntervalFile out of a.bed
>>> intervalfile = pybedtools.IntervalFile(fn)
>>> # any stranded hits?
>>> intervalfile.any_hits(i, same_strand=True)
1
```

count hits

```
Signature IntervalFile.count_hits(interval, same_strand=False,
    overlap=0.0)
```

Return the number of overlaps of the Interval interval had with this IntervalFile.

same_strand, if True, will only consider hits on the same strand as interval.

overlap can be used to specify the fraction of overlap between interval and each feature in the IntervalFile.

```
>>> fn = pybedtools.example_filename('a.bed')
>>> # create an Interval to query with
>>> i = pybedtools.Interval('chr1', 1, 10000, strand='+')
>>> # Create an IntervalFile out of a.bed
>>> intervalfile = pybedtools.IntervalFile(fn)
>>> # get number of stranded hits
>>> intervalfile.count_hits(i, same_strand=True)
3
```

loadIntoMap

Prepares file for checking intersections. Used by other methods like all_hits()

next

x.next() -> the next value, or raise StopIteration

search

```
Signature IntervalFile.all_hits(interval, same_strand=False,
    overlap=0.0)
```

Search for the Interval interval this file and return all overlaps as a list.

same_strand, if True, will only consider hits on the same strand as interval.

overlap can be used to specify the fraction of overlap between interval and each feature in the IntervalFile.

```
>>> fn = pybedtools.example_filename('a.bed')
>>> # create an Interval to query with
>>> i = pybedtools.Interval('chr1', 1, 10000, strand='+')
>>> # Create an IntervalFile out of a.bed
>>> intervalfile = pybedtools.IntervalFile(fn)
>>> # get stranded hits
>>> intervalfile.all_hits(i, same_strand=True)
[Interval(chr1:1-100), Interval(chr1:100-200), Interval(chr1:900-950)]
```

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