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1. Synopsis

SSBT is a program that will allow you to index a set of short-read sequencing experiments and then query them quickly for a given sequence. The first step to use SSBT is to create such an index. This is done via the "hashes", "count", "build", and "compress" commands. Then you can query that index for any sequence to find the files in which that sequence likely appears.

If you use SBT, please cite:

• <pre-release>

2. Analysis Pipeline Overview

To build and query a dataset using SSBT, you should follow this general pipeline:

- 1. You first initialize the hash functions by running "hashes" command.
- 2. Convert each fasta file in the database to a bloom filter bit vector using the "count" command.
- 3. Compile a list of all the bit vectors generated and build the SSBT for that set using the "build" command.
- 4. Compress the bit vectors that make up the entire tree using the built in "compress" command.
- 5. Save your input queries as a line-separated text file and run the "query" command with the desired threshold and using the compressed SSBT file.

More details are given below.

3. Analysis Pipeline Details

You first initialize the hash functions by running "hashes" command. The parameter hashfile is the hashfile output by the "hashes" command, the parameter -k sets the kmer index size (default 20), and the parameter nb_hashes sets the number of hash functions to use. A typical value for this is 1:

```
bt hashes [-k 20] hashfile nb_hashes
```

Next, you count the kmers in each of your input fastq/fasta files using the "count" subcommand; this will produce a bloom filter (with the name filter_out.bf.bv) for the given uncompressed file fasta_in.fasta:

```
bt count [--cutoff 3] [--threads 16] hashfile bf_size fasta_in.fasta filter_out.bf.bv
```

Although we recommend using the default values for cutoff and counting threads, it is possible to adjust the minimum count required for a k-mer to be added to the bloom filter with the --cutoff parameter (the default is to include any element with a count of 3 or greater) or to adjust the number of threads used by the Jellyfish library with the --threads parameter. The parameter hashfile is output from the previous "hashes" command and the bf_size gives the bloom filter size. Although bf_size can be set arbitrarily

large, we suggest setting the bloom filter size to the total count of unique kmers being inserted into the SSBT. We have provided a Jellyfish script (get_bfsize.sh) which takes in a list of fasta.gz files and compiles the total count of unique canonical kmers by their frequency in the total file set. Regardless of the tool used or number selected, it must be the SAME for all files that you are putting into the same SSBT. If you have lots of files on which to build a SSBT, you should write a little script that uses the above command to create a bloom filter for each. Something like:

```
for f in *.fasta ; do
    bt count hashfile bf_size $f 'basename $f .fasta'.bf.bv
done
```

Now, create a list of the .bf.bv files that you just created as text file with one filename per line (using, say, ls *.bf.bv > listoffiles.txt) and call the "build" subcommand:

```
bt build [--sim-type 0] hashfile filterlistfile bloomtreefile
```

Here, filterlistfile is a list of the .bf.bv files. sim-type is an advanced command that changes the rule for where the files are inserted into the tree. After building (which may take some time), bloomtreefile will contain the data about the tree, and there will be many more .bf.bv files created.

Finally, you must compress the tree to query it:

```
bt compress bloomtreefile compressedbloomtreefile
```

This will create a compressed version of the tree (compressed bloomtreefile) which you can then query.

You can issue a query by putting 1 query sequence per line in a query text file and using the "query" subcommand:

```
bt query [--max-filters\ 1]\ [-t\ 0.8]\ [--leaf-only\ 0] [--weighted\ weightfile\ ]\ bloomtreefile\ queryfile\ outfile
```

Here, -t gives the sensitivity threshold: the number of kmers that must be present for a query to be found. Values closer to 1 reduce the number of false positives. You can also weight the individual kmers in the query file by providing a complete array of weights for each kmer in a space separated weightfile.

The "check", "draw", and "sim" subcommands can be used to carry out more specialized tasks. See details below.

4. Command Descriptions

4.1 Hashes

bt hashes [-k 20] hashfile nb_hashes

- k is an optional parameter that sets the k-mer size used in every step of the SSBT pipeline
- hashfile is the location of the file being written
- nb_hashes is an integer that sets the number of hashes generated for the bloom filters

Usage:

To build a set of conserved hash functions for the bloom filters, use a command like:

```
bt hashes myhashfile.hh 1
```

This will write a file 'myhashfile.hh' which stores the necessary information for the Jellyfish library's bloom filter functions.

4.2 Count

bt count [-cutoff 3] [-threads 16] hashfile bf_size fasta_in filter_out.bf.bv

- cutoff is an optional parameter that sets the minimum count required required for a unique k-mer to be added to the bloom filter
- threads is an optional parameter that sets the number of threads the Jellyfish library uses when counting k-mers
- hashfile is the location of the hashfile written using the "hashes" function
- **bf_size** is the number of expected k-mers in the bloom filter.
- fasta_in is the location of the input fasta being counted
- filter_out.bf.bv is the location of the bloom filter being written

Usage:

To convert a fasta short-read file to a SSBT bit vector, use a command like:

bt count myhashfile.hh 2000000000 SRR001.fasta SRR0001.bf.bv

This will count and hash the k-mers associated with 'SRR001.fasta' using the settings defined by 'my-hashfile.hh' and store all k-mers in 'SRR0001.bt'

A note on setting the bf_size: As every filter must have a uniform size, bf_size should be set to an approximate count of the unique k-mers in your complete data set. This avoids saturation in the highest levels of the SSBT and is the extra space is largely factored out through the compression step. If space is a concern, it is also possible to set this value to be the size of the largest leaf filters, as overall accuracy is only affected by the false positive rate of the leaf filters. This will however greatly increase the run-time of SSBT queries.

4.3 Build

bt build [-sim-type 2] hashfile filterlistfile bloomtreefile

- sim-type is an option that defines the similarity metric used. (0) uses the default Hamming distance between the union of bit vector's similarity and remainder filters while (2) first counts the total number of matching 1's in the similarity filter and then uses Hamming distance on the remainder filter when there is no similarity between vectors.
- hashfile is the location of the hashfile written using the "hashes" function
- filterlistfile is the location of a plaintext file containing the paths to all the bit vectors generated by the "count" function
- bloomtreefile is the location of the SSBT structure file being written

Usage:

To build the bloomtree from a list of SSBT bit vectors, use a command like:

 $bt\ build\ myhashfile.hh\ mybitvector list.txt\ mySSBT.bloomtree$

This will build the SSBT through single-threaded insertions of each element in 'mybitvectorlist.txt' and write the union filters to the same directory as the leaves. Once the tree is completely built, the edge-relationships that define the tree will be saved to 'mySSBT.bloomtree'.

4.4 Compress

 $bt\ compress\ bloomtree file\ compressed bloomtree file$

- bloomtreefile is the location of the SSBT structure file written by the "build" function
- compressedbloomtreefile is the location of the [compressed] SSBT structure file being written

Usage:

To compress the bloomtree from bit vectors to rrr compressed vectors, use a command like:

 $bt\ compress\ mySSBT.bloomtree\ myCompressedSSBT.bloomtree$

This will compress every file in the original SSBT and write a new bloomtree using the same edgerelationships but the rrr compressed files.

4.5 Query

bt query [-max-filters 100] [-t 0.8] [-leaf-only 0] [-weighted weightfile] bloomtreefile queryfile outfile

- max-filters is an option that defines the total number of filters that can be loaded at one time into memory. The default is set to 100 filters in memory.
- threshold (t) is a float between 0 and 1 that defines the proportion of query k-mers that must be present in any bloom filter to define a "hit". The default value assumes a valid hit contains 80% of exact-matching k-mers.
- leaf-only has two possible values. (0) is the default value and searches the entire SSBT while (1) ignores the tree structure and queries just the leaf nodes of the tree in a naive search.
- weighted is an optional text file that contains space-separated floats which define in-order weights on the kmer starting at that index in the queryfile. For a length n query, only n-k weights must be provided.
- bloomtreefile is the location of the SSBT structure file written by the "build" function or the compressed SSBT structure file written by the "compressed" function. Using the "compressed" file results in a substantially faster query time.
- queryfile is the location of a text file containing line-separated full-length sequences.
- outfile is the location of the [compressed] SSBT structure file being written

Usage:

To query the SSBT for an arbitrary set of sequences, use a command like:

bt query -t 0.8 mySSBT.bloomtree myQueryFile.txt myOutFile.txt

This will batch query the bloom tree encoded by 'mySSBT.bloomtree' for every line-separated sequence in 'myQueryFile.txt' at a query k-mer threshold of 0.8. If your query of interest is a housekeeping gene or is known to be expressed in the majority of files, it may be beneficial to set the 'leaf_only' option to 1 and ignore the tree structure by querying only the tree leaves.

4.6 Sim

bt sim [-sim-type 0] hashfile bvfile1 bvfile2

- **sim-type** is an option that defines the similarity metric used. (0) uses the default Hamming distance between two bit vectors while (1) uses a Jaccard index metric.
- hashfile is the location of the hashfile written using the "hashes" function

• bvfile1, bvfile2 are any combination of two SSBT bit vectors constructed using the "count" function.

Usage:

To test the raw bit similarity between two bloom filters encoded by the SSBT, use a command like:

 $bt\ sim\ hashfile.hh\ SRR0001.bf.bv\ SRR0002.bf.bv$

This will return the similarity using either the default Hamming (0) or Jaccard (1) metrics.