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Introduction

Molecular biology enzymes have nucleic acid preferences for their substrates; the preference of an enzyme is typically dictated by the sequence at or near the active site of the enzyme. This bias may result in spurious read count patterns when used to interpret high-resolution molecular genomics data. The <code>seqOutBias</code> (https://github.com/guertinlab/seqOutBias) program aims to correct this issue by scaling the aligned read counts by the ratio of genome-wide observed read counts to the expected sequence based counts for each k-mer. The sequence based k-mer counts take into account mappability at a given read length using Genome Tools' Tallymer program. The <code>seqOutBias</code> program allows for flexibility in specifying the k-mer, including varying the k-mer size, k-mer information spacing, and specifying strand-specific offsets for the start of the sequence reads. Due to the large size of some datasets, <code>seqOutBias</code> reads compressed files (FASTA, mappability information, and BAM files), and reuses intermediate results as much as possible.

Requirements

- Platform: OS X or Linux
- Compiler: rust >= 1.5.0 + cargo (<u>http://www.rust-lang.org</u>)
- Genome tools (http://www.genometools.org)
- wigToBigWig (http://genome.ucsc.edu/admin/git.html or http://hgdownload.cse.ucsc.edu/admin/exe/)

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seqOutBias is written in Rust, so it requires Rust and Cargo to compile it (www.rust-lang.org). seqOutBias is known to run on OS X and Linux. It may be possible to compile it on other platforms as long as supporting Rust libraries can be made to compile, specifically https://crates.io/crates/flate2 and https://crates.io/crates/rust-htslib are likely to be the dependencies that are most troublesome to compile.

At runtime, the step that computes mappability requires Genome Tools' Tallymer program to be in the PATH. It is possible to replicate the steps taken by *seqOutBias* to compute the mappability file independently, or run that part in a different machine, and supply the resulting file to *seqOutBias* (see the tallymer sub-command for more details).

The step that generates a bigWig file requires the UCSC executable wigToBigWig in the PATH.

Setup

Compilation

To install Rust and Cargo visit www.rust-lang.org, *seqOutBias* should compile with Rust 1.5.0 or later. Alternatively, on OS X, if you have Homebrew (http://brew.sh/) installed and up to date, you can use brew to install rust as so:

```
brew install rust
```

Afterwards, uncompress the source and build:

```
tar xzf guertinlab-seqOutBias-v1.0.2.tar.gz

cd guertinlab-seqOutBias-v1.0.2

cargo build --release
```

Installation

After compilation, copy the *seqOutBias* binary (seqOutBias_1.0.1/target/release/seqOutBias) to a folder in your PATH, for example /usr/local/bin.

You'll also need to install genome tools (http://www.genometools.org). Again, on OS X, if you have Homebrew (http://brew.sh) installed you can install genome tools as so:

```
brew install homebrew/science/genometools
```

You will need to build the wigToBigWig utility from the UCSC Genome Browser source code (http://genome.ucsc.edu/admin/git.html) and move the binary into your PATH, following the instructions included with the Genome Browser source code. This command is required to obtain bigWig files after scaling. ALternativey, you can download the platform-specific binary from UCSC (http://hgdownload.cse.ucsc.edu/admin/exe/).

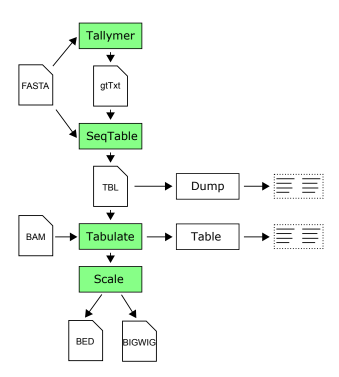
Quick Start

With *genometools, wigToBigWig* and *seqOutBias* installed and in the PATH, we can start processing files by simply invoking the command as:

where **ref.fa** and **reads.bam** are, respectively, the reference sequence FASTA file and the **sorted** aligned reads BAM file¹. This will run through the process of computing the mappability information, parsing the reference sequence to compute k-mer indexes, tallying the k-mer counts in both the sequence and the aligned reads, and finally producing the scaled BED and bigWig read pile-ups. Each intermediate step corresponds to a *seqOutBias* subcommand, allowing them to be run individually and even on different machines (see the next sections for more details).

When ran with no arguments, this will assume default values for read length (36), k-mer size (4) and the cut-site position (middle of the k-mer, i.e., plus and minus offset = 2). Run seqOutBias -h to see the full set of options and subcommands.

Overview



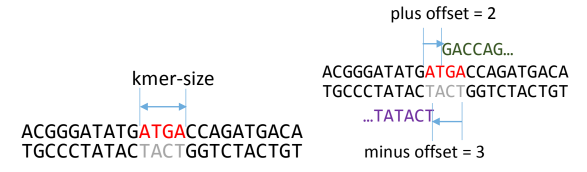
The figure above gives a high-level overview of the inputs (left side "documents"), intermediate files (mid-flow "documents") and output (bottom) of the *seqOutBias* program. Furthermore, it illustrates the various computation steps that the program performs:

¹ It's possible to supply more than one BAM file, for example, to combine the data from multiple replicates.

- Tallymer Indexes the reference sequence (FASTA) and computes mappability for the given read length;
- SeqTable Parses the reference sequence (FASTA) together with the mappability information to
 compute the k-mer that corresponds to each possible read alignment position; the resulting
 binary file stores this information in a compressed form that's easy to use for subsequent
 computation steps, as well as storing the corresponding parameters (read length, k-mer size,
 and cut-site offsets);
- Tabulate Tallies the k-mer counts across the selected regions (or full genome), as well as the k-mers corresponding to observed aligned reads (if a BAM file is supplied);
- Scale Compute the genome-wide aligned read pile-ups, scaling them by the expected/observed cut frequency;

When executed as described in the "Quick Start" section, all the main computation steps (marked in green) are executed, however, different subcommands can be used to run specific computation steps (see "Subcommands" section), or obtain text versions of the intermediate states ("dump" to display the SeqTable output, "Table" to obtain the normalization table data).

kmer-size definition



In *seqOutBias*, the sequence recognized by the enzyme to confer specificity, the k-mer, is characterized by four parameters (illustrated above): kmer-size, read size, and a pair of offsets for the plus and minus strands. This enables the use of *seqOutBias* with enzymes that have distinct recognition site lengths.

Note: Only non-negative value offsets are permitted.

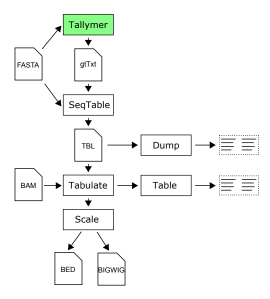
Spaced k-mers

For situations where some bases surrounding the first sequenced base do not contribute to site recognition, it is possible to specify a kmer-mask, using the "--kmer-mask" parameter. Positions that should be ignored are represented by an 'X' and regular positions by an 'N'. For example, a possible 8-mer that spans 16 bp could be represented as NNXXNNXXXXNNXXNN.

This parameter also provides an alternative way to specify the position intervening between the base directly upstream the first base sequenced and the first base sequence by inserting a 'C' in the mask string. For example, "NNCNN", would represent a recognition site with size equal to four, plus-offset = 2 and minus-offset = 2.

Note: When the kmer-mask is present, it takes precedence over the "--kmer-size" parameter. If a 'C' is used to indicate the cut-site position, it takes precedence over the offset parameters.

Subcommands tallymer



seqOutBias tallymer <fasta-file> <read-size> [--parts=<n>]

This subcommand creates the mappability file for a given read length. This process consists of three parts: 1) creating a suffix tree; 2) creating a genome index; 3) creating the mappability file.

It corresponds to running the following genome tools' commands (replacing the bolded parts with the *seqOutBias* arguments):

```
gt suffixerator -dna -pl -tis -suf -lcp -v -parts <n> -db <fasta-file>
    -indexname <fasta-file>.sft

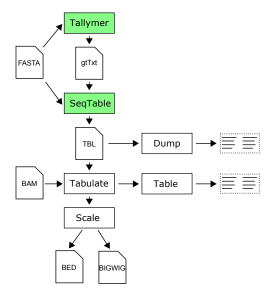
gt tallymer mkindex -mersize <read-size> -minocc 2 -indexname <fasta-
file>.tal_<read-size> -counts -pl -esa <fasta-file>.sft

gt "tallymer search -output qseqnum qpos -strand fp -tyr <fasta-
file>.tal_<read-size> -q <fasta-file> > <fasta-file>.tal_<read-
size>.gtTxt
```

Intermediate files are created in the current working directory. Furthermore, *seqOutBias* will recognize the existence of intermediate files and avoid recomputing those portions. So, if seqOutBias tallymer is executed for different read sizes, but keeping the same FASTA file, then the first suffix-tree portion is reused across invocations.

If the input FASTA file is compressed, then so will be the output file.

seqtable



seqOutBias seqtable <fasta-file> [options]

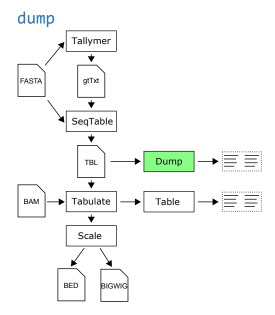
```
--kmer-size=<n>
                    kmer size [default: 4].
--tallymer=<file>
                    Unmappable positions file produced by tallymer (seq, pos).
--plus-offset=
                    Offset on plus strand [default: 2]. Eg, p=2 AA[A]A.
                   Offset on minus strand [default: 2]. Eg, m=2 A[A]AA.
--minus-offset=<m>
--kmer-mask=<str>
                    String indicating relevant kmer positions, eg.
NNXXNNCXXXXNNXXNN.
                    Read length [default: 36].
--read-size=<r>
                    Split suffix tree generation into n parts [default: 4].
--parts=<n>
--out=<outfile>
                    Output seqtable filename (defaults to fasta file basename with
.tbl extension).
```

This subcommand creates an intermediate table that combines mappability, read length, and offsets to map k-mer indexes to the aligned read positions. This intermediate file reduces the amount of computation needed when processing aligned read files and provides an intermediate file that decouples the reference sequence processing from the remaining steps.

Note: If no tallymer output file is supplied (via the --tallymer=<file> option), then it will invoke the tallymer subcommand.

Note: When the k-mer mask is present, it takes precedence over the "--kmer-size" parameter. If a 'C' is used in the "--kmer-mask" parameter to indicate the position relative to the first base sequenced, it takes precedence over the offset parameters.

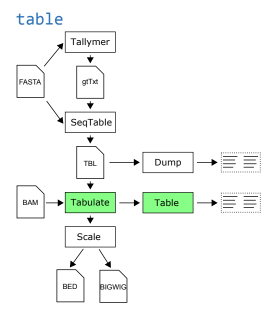
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seqOutBias dump <seqtbl-file> [<seqrange>]

This subcommand enables the inspection the seqtable file (or parts of it) by outputting the file contents in plain text. It takes an optional sequence range in the form of "chrom:start-end" that restricts the output to that region. The output is preceded by a list of the input parameters (read-size, kmer-size, etc.) that were used to build the sequence table file.

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```
seqOutBias table <seqtbl-file> [<bam-file>...] [--qual=<q>] [--
regions=<bedfile>] [--pdist=<min:max>] [--only-paired]
```

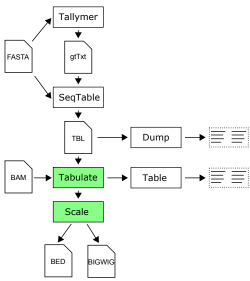
This subcommand produces an k-mer count table based on the sequence information (via the seqtbl file) and the optional BAM files. If more than one file is given, then the counts are pooled across files. **K-mers are indexed by their plus strand sequence** (for example, AAAA = 1, AAAC = 2, AAAG = 3, etc), but counts are reported independently for each strand. It's done this way because mappability can differ between strands. By default counts correspond to the entire genome, but can be constrained to specific regions by supplying a BED file with the "--regions" option.

When no BAM file is supplied, the output will have four columns: k-mer index, k-mer string, plus strand count, and minus strand count. If a BAM file is supplied, the output will have two additional columns with the plus and minus strand counts of observed aligned reads. For example:

```
1
      AAAA
             40584084
                          39080470
2
      AAAC
             14531729
                          14684467
3
      AAAG
             19041196
                          19266837
4
      AAAT
             26793796
                          27032258
5
      AACA
             15881340
                          16097718
6
                          8098141
      AACC
            7943846
7
      AACG
            1562521
                          1571244
8
      AACT
            12186297
                          12417246
9
      AAGA
             17771087
                          18024000
```

If the --exact-length flag was set, aligned reads are filtered by read length; only the reads that match the read length will be used to produce the sequence table are used. The user can optionally filter quality score of the aligned reads. For paired reads, it is also possible to filter by their distance (furthest edge-to-edge) and to discard singleton reads. Unaligned reads are discarded.

scale



seqOutBias scale <seqtbl-file> <bam-file>... [options]

--qual=<q> Minimum read quality [default: 0].

--regions=<bedfile> Count only cut-sites inside the regions indicated in the BED file.

--bed=<bedfile> Output scaled BED filename (defaults to BAM file basename with '_scaled.bed' extension).

--skip-bed Skip creating the BED file output.

--bw=<bigwigfile> Output scaled bigWig filename (defaults to BAM file basename with .bw extension).

--skip-bw Skip creating the bigWig file output.

--stranded Output per strand counts when writing scaled values.

--shift-counts Shift minus strand counts.

--no-scale Skip actual scaling in 'scale' command.

--pdist=<min:max> Distance range for included paired reads.

--only-paired Only accept aligned reads that have a mapped pair.

--tail-edge Use tail edge of reads (3') instead of start edge (5').

This subcommand produces the scaled aligned read pile-ups, both in BED and bigWig form. Listed above are the options to control the output. Aside from the filtering options that apply to the "tabulate" computation, there are a few options to tweak the results, namely:

- Scaling can be turned off using the "--no-scale" option;
- Minus strand pile-up position can be shifted to align with the plus strand pile-up ("--shift-counts" option), making reads from both sides of a cleavage site count to the same position;

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- Output per strand, using the "--stranded" option will result in one bigWig file per strand, as well as per strand counts inside the BED file (plus strand with a positive sign, minus strand with a negative sign);
- Output type selection, using either the "--skip-bed" or the "--skip-bw" option will omit the production of the BED or bigWig file respectively.
- For PRO-seq and similar protocols, the "--tail-edge" option will use the 3' end of the read.
- Output file name, using the "--bed" or "--bw" flags;