Package 'STITCH'

February 7, 2017

Type Package

Title STITCH - Sequencing To Imputation Through Constructing Haplotypes

Version 1.2.7

Date 2017-02-07

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Description STITCH performs imputation of individuals sequenced to low coverage in a read aware fashion without a reference panel.

Installation To install, first install dependencies, then run the install.packages command, pointing to the downloaded tarball (STITCH.tar.gz)

Getting started A minimum run requires the following options to be set: the chromosome being run (chr); a path to a file with a set of bi-allelic SNP sites (posfile); a choice of K, the number of internally modelled haplotypes (K); a path to an output directory (outputdir); a path to a temporary directory, ideally on fast disks or a RAM disk (tempdir); a list of bam files (bamlist); and the number of generations since founding (nGen), which can be approximated from a choice of K for wild populations from 4 * Ne / K. Additional useful options relate to what region to impute (regionStart, regionEnd, buffer), whether to use validation data to benchmark imputation (genfile), the number of cores to use (nCores), whether imputation is run on a server or cluster (environment), the number of EM iterations (niterations), whether to run in diploid or pseudoHaploid mode (method), and if run in pseudoHaploid mode, what iteration to switch from pseudoHaploid to diploid (switchModelIteration).

Depends parallel, Rsamtools

Imports Rcpp

LinkingTo Rcpp, RcppArmadillo

RoxygenNote 5.0.1

License GPL | file LICENSE

SystemRequirements C++11

NeedsCompilation yes

Suggests testthat

R topics documented:

STITCH

Sequencing To Imputation Through Constructing Haplotypes

Usage

```
STITCH(chr, nGen = "", posfile, K, outputdir, tempdir, bamlist = "",
 cramlist = "", reference = "", genfile = "", method = "diploid",
 outputInputInVCFFormat = FALSE, downsampleToCov = 50,
 downsampleFraction = 1, readAware = TRUE, chrStart = NA, chrEnd = NA,
 regionStart = NA, regionEnd = NA, buffer = NA,
 maxDifferenceBetweenReads = 1000, alphaMatThreshold = 1e-04,
 emissionThreshold = 1e-04, iSizeUpperLimit = as.integer(600),
 bqFilter = as.integer(17), niterations = 40,
 shuffleHaplotypeIterations = c(4, 8, 12, 16), splitReadIterations = 25,
 nCores = 1, expRate = 0.5, maxRate = 100, minRate = 0.1,
  Jmax = 1000, regenerateInput = TRUE, originalRegionName = NA,
 keepInterimFiles = FALSE, keepTempDir = FALSE, environment = "server",
 pseudoHaploidModel = 9, outputHaplotypeProbabilities = FALSE,
  switchModelIteration = NA, generateInputOnly = FALSE,
 restartIterations = NA, refillIterations = c(6, 10, 14, 18),
 downsampleSamples = 1, downsampleSamplesKeepList = NA,
  subsetSNPsfile = NA, useSoftClippedBases = FALSE,
 outputBlockSize = 1000, inputBundleBlockSize = NA,
 reference_haplotype_file = "", reference_legend_file = "",
 reference_sample_file = "", reference_populations = NA,
 reference_phred = 20, reference_iterations = 10, vcf_output_name = NULL,
  initial_min_hapProb = 0.4, initial_max_hapProb = 0.6,
 regenerateInputWithDefaultValues = FALSE,
 plotHapSumDuringIterations = FALSE)
```

Arguments

| chr | What chromosome to run. Should match BAM headers |
|-----------|---|
| nGen | Number of generations since founding or mixing. Note that the algorithm is relatively robust to this. Use $nGen = 4 * Ne / K$ if unsure |
| posfile | Where to find file with positions to run. File is tab seperated with no header, one row per SNP, with col 1 = chromosome, col 2 = physical position (sorted from smallest to largest), col 3 = reference base, col 4 = alternate base. Bases are capitalized. Example first row: 1 <tab>1000<tab>A<tab>G<tab></tab></tab></tab></tab> |
| K | How many founder / mosaic haplotypes to use |
| outputdir | What output directory to use |
| tempdir | What directory to use as temporary directory. If possible, use ramdisk, like /dev/shm/ |

Path to file with bam file locations. File is one row per entry, path to bam files.

Bam index files should exist in same directory as for each bam, suffixed either

.bam.bai or .bai

cramlist Path to file with cram file locations. File is one row per entry, path to cram files.

cram files are converted to bam files on the fly for parsing into STITCH

reference Path to reference fasta used for making cram files. Only required if cramlist is

defined

genfile Path to gen file with high coverage results. Empty for no genfile. File has a

header row with a name for each sample, matching what is found in the bam file. Each subject is then a tab separated column, with 0 = hom ref, 1 = het, 2 = hom alt and NA indicating missing genotype, with rows corresponding to rows of the posfile. Note therefore this file has one more row than posfile which has

no header

method How to run imputation - either diploid or pseudoHaploid, the former being the

original method quadratic in K, the later being linear in K

outputInputInVCFFormat

Whether to output the input in vcf format

downsampleToCov

What coverage to downsample individual sites to. This ensures no floating point

errors at sites with really high coverage

downsampleFraction

Downsample BAMs by choosing a fraction of reads to retain. Must be value

0<downsampleFraction<1

readAware Whether to run the algorithm is read aware mode. If false, then reads are split

into new reads, one per SNP per read

chrStart When loading from BAM, some start position, before SNPs occur. Default NA

will infer this from either regionStart, regionEnd and buffer, or posfile

chrEnd When loading from BAM, some end position, after SNPs occur. Default NA

will infer this from either regionStart, regionEnd and buffer, or posfile

regionStart When running imputation, where to start from. The 1-based position x is kept if

 $regionStart \le x \le regionEnd$

regionEnd When running imputation, where to stop

buffer Buffer of region to perform imputation over. So imputation is run form regionStart-

buffer to regionEnd+buffer, and reported for regionStart to regionEnd, including

the bases of regionStart and regionEnd

 ${\tt maxDifferenceBetweenReads}$

How much of a difference to allow the reads to make in the forward backward probability calculation. For example, if P(read | state 1)=1 and P(read | state 2)=1e-6, re-scale so that their ratio is this value. This helps prevent any individual read as having too much of an influence on state changes, helping prevent

against influence by false positive SNPs

alphaMatThreshold

Minimum (maximum is 1 minus this) state switching into probabilities

emissionThreshold

Emission probability bounds. emission Threshold < P(alt read | state k) < (1-emission Threshold)

iSizeUpperLimit

Do not use reads with an insert size of more than this value

Minimum BQ for a SNP in a read. Also, the algorithm uses bq<=mq, so if bqFilter

mapping quality is less than this, the read isnt used

Number of EM iterations. niterations

shuffleHaplotypeIterations

Iterations on which to perform heuristic attempt to shuffle founder haplotypes for better fit. To disable set to NA.

splitReadIterations

Iterations to try and split reads which may span recombination breakpoints for a

better fit

nCores How many cores to use

expRate Expected recombination rate in cM/Mb

maxRate Maximum recomb rate cM/Mb minRate Minimum recomb rate cM/Mb Maximum number of SNPs on a read

regenerateInput

Jmax

Whether to regenerate input files

originalRegionName

If regenerateInput is FALSE (i.e. using existing data), this is the name of the original region name (chr.regionStart.regionEnd). This is necessary to load past variables

keepInterimFiles

Whether to keep interim parameter estimates

keepTempDir Whether to keep files in temporary directory

environment Whether to use server or cluster multicore options

pseudoHaploidModel

How to model read probabilities in pseudo diploid model (shouldn't be changed)

outputHaplotypeProbabilities

Whether to output haplotype probabilities in files

switchModelIteration

Whether to switch from pseudoHaploid to diploid and at what iteration (NA for no switching)

generateInputOnly

Whether to just generate input data then quit

restartIterations

In pseudoHaploid method, which iterations to look for collapsed haplotype prnobabilities to resolve

refillIterations

When to try and refill some of the less frequently used haplotypes

downsampleSamples

What fraction of samples to retain. Useful for checking effect of N on imputation. Not meant for general use

 ${\tt downsampleSamplesKeepList}$

When downsampling samples, specify a numeric list of samples to keep

subsetSNPsfile If input data has already been made for a region, then subset down to a new set of SNPs, as given by this file. Not meant for general use

useSoftClippedBases

Whether to use (TRUE) or not use (FALSE) bases in soft clipped portions of

outputBlockSize

How many samples to write out to disk at the same time when making temporary VCFs that are later pasted together at the end to make the final VCF. Smaller means lower RAM footprint, larger means faster write.

inputBundleBlockSize

If NA, disable bundling of input files. If not NA, bundle together input files in sets of <= inputBundleBlockSize together

reference_haplotype_file

Path to reference haplotype file in IMPUTE format (file with no header and no rownames, one row per SNP, one column per reference haplotype, space separated, values must be 0 or 1)

reference_legend_file

Path to reference haplotype legend file in IMPUTE format (file with one row per SNP, and a header including position for the physical position in 1 based coordinates, a0 for the reference allele, and a1 for the alternate allele)

reference_sample_file

Path to reference sample file (file with header, one must be POP, corresponding to populations that can be specified using reference_populations)

reference_populations

Vector with character populations to include from reference_sample_file e.g. CHB, CHS

reference_phred

Phred scaled likelihood or an error of reference haplotype. Higher means more confidence in reference haplotype genotypes, lower means less confidence

reference_iterations

When using reference haplotypes, how many iterations to use to train the starting data

vcf_output_name

Override the default VCF output name with this given file name. Please note that this does not change the names of inputs or outputs (e.g. RData, plots), so if outputdir is unchanged and if multiple STITCH runs are processing on the same region then they may over-write each others inputs and outputs

initial_min_hapProb

Initial lower bound for probability read comes from haplotype. Double bounded between 0 and 1

initial_max_hapProb

Initial upper bound for probability read comes from haplotype. Double bounded between $\boldsymbol{0}$ and $\boldsymbol{1}$

regenerate Input With Default Values

If regenerateInput is FALSE and the original input data was made using region-Start, regionEnd and buffer as default values, set this equal to TRUE

plotHapSumDuringIterations

Boolean TRUE/FALSE about whether to make a plot that shows the relative number of individuals using each ancestral haplotype in each iteration

Value

Results in properly formatted version

Author(s)

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Index

STITCH, 2