Reference-based 2bRAD scripts: a manual

October 14, 2014 - April 18, 2015

These are the printouts from scripts when run without arguments, in order of their appearance in the walkthrough.

```
ngs_concat.pl :
concatenates files by matching pattern in their names
arg1: common pattern for files
arg2: perl-like pattern in the filename to recognize,
       use brackets to specify the unique part, as in
      "FilenameTextImmediatelyBeforeSampleID(.+)FilenameTextImmediatelyAfterSampleID"
Example (to concatenate files names like Sample_Pop1_L1.fastq, Sample_Pop1_L2.fastq):
nqs_concat.pl 'Sample' 'Sample_(.+)_L'
2bRAD_trim_launch_dedup.pl :
Prints out list of calls to trim2bRAD_2barcodes_dedup.pl, one for
each fastq file, to to trim and deduplicate raw 2bRAD reads.
See help for trim2bRAD_2barcodes_dedup.pl for more details.
prints to STDOUT
Arguments:
arg1, required: glob to fastq files
site=[pattern] perl-style pattern of the restriction site to recognise, in single quotes.
           default is BcgI: \'.{12}CGA.{6}TGC.{12}\.{12}GCA.{6}TCG.{12}\'
           if you need AlfI, here is how to define it: \'.{12}GCA.{6}TGC.{12}\'
adaptor=[dna sequence] adaptor sequence to look for on the far end of the read.
           Default AGATC
sampleID=[integer] the position of name-deriving string in the file name
                 if separated by underscores, such as:
                 for input file Sample_RNA_2DVH_L002_R1.cat.fastq
                 specifying arg2 as \'3\' would create output
                 file with a name \'2DVH.trim'
barcode2=[integer] length of the in-line barcode immediately following
                    the restriction fragment. Default 0.
Example:
2bRAD_trim_launch_dedup.pl fastq sampleID=3 > trims
```

trim2bRAD_2barcodes_dedup.pl :

This script does three things:

- Filters 2bRAD fastq reads to leave only those with a 100% matching restriction site, degenerate 5'-leader, secondary 3'-barcode, and adaptor on the far 3'end, trims away everything except the IIb-fragment itself;
- Deduplicates: removes all but one read sharing the same 64-fold degenerate leader, the first 34 bases of the insert sequence, and secondary barcode (this results in 128-fold dynamic range: 64-fold degeneracy x 2 strand orientations);
- Splits reads into separate files according to secondary barcode.

Writes trimmed fastq files named according to the secondary barcodes detected.

Arguments:

input=[fastq filename]

site=[perl-style pattern for restrictase site]

Default is BcgI: \'.{12}CGA.{6}TGC.{12}|.{12}GCA.{6}TCG.{12}\'

bc2=[perls-style barcode pattern] in-line barcode that immediately follows the RAD fragment,

default \'[ATGC]{4}\'

adaptor=[far-end adaptor sequence] default AGATC

clip=[integer] number of bases to clip off the ends of the reads, default 0 minBCcount=[integer] minimum count per in-line barcode to output a separate file.

Default 100000.

sampleID=[integer] the position of name-deriving string in the file name

if separated by underscores, such as:

for input file Sample_RNA_2DVH_L002_R1.cat.fastq

specifying arg2 as \'3\' would create output

file with a name \'2DVH.trim'

Example:

trim2bRAD_2barcodes_dedup.pl input=Myproject_L8_G3.fastq sampleID=2

concatFasta.pl :

concatenates fasta records in a draft genome reference into \"pseudo-chromosomes\" (either a specified number of them or as many as needed to split the genome into chunks just over 100G bases in length)

Do this to your genotyping reference to improve the speed and memory usage of the UnifiedGenotyper (GATK)

Arguments:

fasta=[file name] : fasta file name

num=[integer] : number of pseudo-chromosomes, default 10

Output:

[fasta filename base]_cc.fasta : new fasta file

[fasta filename base]_cc.tab : table of original sequence IDs, their coordinates

in pseudo-chromosomes, and full original header lines

Example: concatFasta.pl fasta=mygenome.fasta

retabvcf.pl :

rewrites coordinates of variants in a VCF file according to a table of locations of original contigs in pseudo-chromosomes (created by concatFasta.pl)

Arguments:

vcf=[file name] : vcf file to re-coordinate

tab=[file name] : tab-delimited table of contig locations in the form

contig-chromosome-start-end

Output:

VCF with reformatted coordinates (printed to STDOUT)

Example:

retabvcf.pl vcf=gatk_after_vqsr.vcf tab=where/genome/is/mygenome_cc.tab > retab.vcf

replicatesMatch.pl : (version 0.5, September 11 2015)

Selects polymorphic variants that have identical genotypes among replicates.

Genotypes of two replicates should be identical or missing; the maximal fraction of genotype pairs with missing genotypes is controlled by missing argument.

Arguments:

matching=[float] required fraction of matching genotypes

(missing counts as match if there are other non-missing matching pairs),

default 1 (all must match)

missing=[float] allowed fraction of missing genotypes, default 0.25

allAlts= $\lceil 1/0 \rceil$ output all SNPs showing non-reference alleles (not necessarily

polymorphic among genotyped samples). Default ${\tt 0}$

hetMatch=2 minimal number of matching heterozygotes. Ignored if allAlts=1.

max.het=0.5 maximum fraction of heterozygotes among non-missing genotypes

(guards against lumped paralogous loci).

Example:

replicatesMatch.pl vcf=round2.vcf replicates=clonepairs.tab > vqsr.vcf

recalibrateSNPs_gatk.pl :

Non-parametric variant quality recalibration based on INFO fields in the set of variants that are reproducibly genotyped among replicates (output of replicatesMatch.pl)

This script is for reference-based pipeline. Use recalibrateSNPs.pl for de novo pipeline.

Three parameters (INFO field) are assessed for each SNP:

- fisher-strand (FS)
- mapping quality (MQ)
- quality by depth (QD)
- depth (DP)

The script computes the product of quantiles for different INFO fields, determines the quantiles of the result within the 'true' set (JOINT quantiles), and then computes the new quality scores in the main vcf file.

These scores are supposed to correspond to the probability (x100) that the SNPs comes from the same distribution as the 'true' SNPs.

Output:

- Recalibrated VCF (printed to STDOUT) with QUAL field replaced by the new score minus 0.1 (for easy filtering)
- a table (printed to STDERR) showing the \"gain\" at each recalibrated quality score setting, which is excess variants removed when filtering at this quality score. For example, if 40% of all variants are removed at the quality score 15, the gain is 40 15 = 35% (i.e., in addition to removing 15% of the 'true' variants, additional 35% of the dataset, likely corresponding to wrong variants, is removed). The optimal filtering score is the one giving maximum gain. Try different combinations of the four possible filters to find the one maximizing the gain.

Arguments:

vcf=[file name] : vcf file to be recalibrated

true=[file name] : vcf file that is subset of the above, with SNPs that are considered

true because they show matching and polymorphic genotypes in

replicates

(replicatesMatch.pl polyonly=1).

-nodp : do not use DP
-nofs : do not use FS
-nomq : do not use MQ
-noqd : do not use QD

Example:

recalibrateSNPs_gatk.pl vcf=round2.names.vcf true=vqsr.vcf > recal.vcf

filterStats_gatk.pl :

Calculates quantiles for three filtering criteria using VCF file:

- fisher-strand (FS)
- mapping quality (MQ)
- depth (DP)

```
For MQ, low values are bad and high values are good;
```

for FS, high values are bad;

for DP, both high and low values are bad

Arguments:

vcf=[file name] vcf file to analyze

0utput

Tables of quantiles (printed to STDOUT).

Example:

filterStats_gatk.pl vcf=round2.names.vcf

NOTE: only use this to select your filtering criteria if you don't have genotyping replicates.

repMatchStats.pl :

Summarizes genotypic match between replicates in a vcf file.

Arguments:

```
vcf=[file name] : input vcf file
```

replicates=[file name] : a two column tab-delimited table listing

pairs of samples that are replicates

Output:

A table (printed to STDOUT) of the following form:

```
pair gtyped match [ 00 01 11 ] HetMatch HomoHetMism HetNoCall HetsDiscRate K210:K212 7328 7169(97.8%) [ 78% 17% 5% ] 1200 135 5 0.94 K212:K213 7369 7117(96.6%) [ 78% 17% 5% ] 1202 179 4 0.93
```

The first four columns are self-evident;

the last four columns show how good is the match between heterozygote calls (HetNoCall being a heterozygote in one and missing data in another replicate). The last column is the most important: it is the heterozygote discovery rate, the fraction of all heterozygotes discovered in each replicate.

Example: repMatchStats.pl vcf=filt.recode.vcf replicates=clonepairs.tab

thinner.pl :

Leaves only one SNP per given interval, the one with the highest minor allele frequency.

Arguments:

vcf=[file name] : vcf file name

interval=[integer] : interval length, default 40 (for 2bRAD tags)

Output: thinned VCF, printed to STDOUT

Example: thinner.pl vcf=recal.vcf > thin.vcf

NOTE: do not thin your variants if you want to calculate Tajima's D! Also, to select variants for dadi analysis, use --thin option in vcftools instead.

vcf2genepop.pl :

converts VCF to multiallelic GENEPOP, preserves chromosome and position info

Arguments:

vcf=[filename] : vcf file to convert

pops=[list] : comma-separated perl patterns to determine population

affiliation of samples in the VCF file based on sample names

Output:

GENEPOP formatted genotypes, printed to STDOUT

Example:

vcf2genepop.pl vcf=filtered.vcf pops=0,K,M,S > filtered.gen

vcf2dadi.pl :

Converts VCF format to dadi data format (requires bioperl)

Usage: perl vcf2dadi.pl <genome file> <vcf file> <list file>

Genome file is in fasta format;

List file gives population designations, like this:

BEGIN

sample1 population1
sample2 population1
sample3 population2

END