Copy number calling and SNV classification using targeted short read sequencing

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Abstract

PureCN [1] is a purity and ploidy aware copy number caller for cancer samples inspired by the *ABSOLUTE* algorithm [2]. It was designed for hybrid capture sequencing data, especially with medium-sized targeted gene panels without matching normal samples in mind (matched whole-exome data is of course supported).

It can be used to supplement existing normalization and segmentation algorithms, i.e. the software can start from BAM files, from target-level coverage data, from copy number log2-ratios or from already segmented data. After the correct purity and ploidy solution was identified, *PureCN* will accurately classify variants as germline vs. somatic or clonal vs. sub-clonal.

PureCN was further designed to integrate well with industry standard pipelines [3], but it is straightforward to generate input data from other pipelines.

Package

PureCN 2.11.0

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1 Introduction

This tutorial will demonstrate on a toy example how we recommend running *PureCN* on targeted sequencing data. To estimate tumor purity, we jointly utilize both target-level¹ coverage data and allelic fractions of single nucleotide variants (SNVs), inside - and optionally outside - the targeted regions. Knowledge of purity will in turn allow us to accurately (i) infer integer copy number and (ii) classify variants (somatic vs. germline, mono-clonal vs. sub-clonal, heterozygous vs. homozygous etc.).

¹The captured genomic regions, e.g. exons.

This requires 3 basic input files:

- 1. A VCF file containing germline SNPs and somatic mutations. Somatic status is not required in case the variant caller was run without matching normal sample.
- 2. The tumor BAM file.
- 3. At least one BAM file from a normal control sample, either matched or processmatched.

In addition, we need to know a little bit more about the assay. This is the annoying step since here the user needs to provide some information. Most importantly, we need to know the positions of all targets. Then we need to correct for GC-bias, for which we need GC-content for each target. Optionally, if gene-level calls are wanted, we also need for each target a gene symbol. We may also observe subtile differences in coverage of tumor compared to normal due to varying proliferation rates and we can provide replication timing data to check and correct for such a bias. To obtain best results, we can finally use a pool of normal samples to automatically learn more about the assay and its biases and common artifacts.

The next sections will show how to do all this with *PureCN* alone or with the help of *GATK* and/or existing copy number pipelines.

All the steps described in the following are available in easy to use command line scripts described in a separate vignette.

2 Basic input files

2.1 VCF

Germline SNPs and somatic mutations are expected in a single VCF file. At the bare minimum, this VCF should contain read depths of reference and alt alleles in an AD format field and a DB info flag for membership in germline databases².

Without DB flag, variant ids starting with **rs** are assumed to be in dbSNP. Population allele frequencies are expected in a POP_AF info field. These frequencies are currently only used to infer membership in germline databases when the DB flag is missing; in future versions they will be used calculate somatic prior probabilities more accurately.

If a matched normal is available, then somatic status information is currently expected in a SOMATIC info flag in the VCF. The *VariantAnnotation* package provides examples how to add info fields to a VCF in case the used variant caller does not add this flag. If the VCF contains a BQ format field containing base quality scores, *PureCN* can remove low quality calls.

²The name of the flag is customizable in runAb soluteCN VCF files generated by *MuTect* [4] should work well and in general require no post-processing. *PureCN* can handle *MuTect* VCF files generated in both tumor-only and matched normal mode. Beta support for *MuTect 2* and *FreeBayes* VCFs is available.

As alternative to VCF input, read counts of common SNPs are also supported in case classification of somatic variants is of no interest and those genotyped SNPs are already generated in an upstream pipeline. The function **readAllelicCountsFile** can be used to parse a *GATK4 CollectAllelicCounts* file into a VCF supported by *PureCN*³:

```
line is required for ex-
@HD VN:1.6
                                                                                        tracting the sample id
@SQ SN:1 LN:248956422
@SQ SN:2 LN:242193529
. . .
@RG ID:GATK SM:Sample1
CONTIG POSITION REF_COUNT ALT_COUNT REF_NUCLEOTIDE ALT_NUCLEOTIDE
1 114515871 177 189 G A
1 150044293 119 157 T G
ac.file <- system.file("extdata", "example_allelic_counts.tsv",</pre>
          package="PureCN")
vcf.ac <- readAllelicCountsFile(ac.file)</pre>
## INFO [2024-05-01 18:54:13] Found 20 variants in VCF file.
## INFO [2024-05-01 18:54:13] 20 (100.0%) variants annotated as likely germline (DB INFO flag).
## WARN [2024-05-01 18:54:13] Did not find base quality scores, will use global error rate of 0.0010 instead
```

2.2 Target information

For the default segmentation function provided by *PureCN*, the algorithm first needs to calculate log2-ratios of tumor vs. normal control coverage. To do this, we need to know the locations of the captured genomic regions (targets). These are provided by the manufacturer of your capture kit⁴. Please double check that the genome version of the target file matches the reference. Usually the manufacturer provides two files: the baits file containing the coordinates of the actual capture baits, and the target file containing the coordinates of the actual regions we want to capture. We recommend to use the baits file (and recognize the confusing nomenclature that we follow due to convention in established tools).

Default parameters assume that these targets do NOT include a "padding" to include flanking regions. *PureCN* will automatically include variants in the 50bp flanking regions if the variant caller was either run without interval file or with interval padding (See section 12.2).

PureCN will attempt to optimize the targets for copy number calling (similar to [5]):

- Large targets are split to obtain a higher resolution
- Targets in regions of low mappability are dropped
- Optionally, accessible regions in-between the target (off-target) regions are included so that available coverage information in on- and off-target reads can be used by the segmentation function. In the following, we will use *intervals* when something applies to both on-target and off-target regions and *targets* when it only applies to on-target.

It further annotates intervals by GC-content (how coverage is normalized is described later in Section 3).

PureCN provides the preprocessIntervals function:

⁴While *PureCN* can use a pool of normal samples to learn which intervals are reliable and which not, it is highly recommended to provide the correct intervals. Garbage in, garbage out.

³Only the RG header

4

```
reference.file <- system.file("extdata", "ex2_reference.fa",
    package = "PureCN", mustWork = TRUE)
bed.file <- system.file("extdata", "ex2_intervals.bed",
    package = "PureCN", mustWork = TRUE)
mappability.file <- system.file("extdata", "ex2_mappability.bigWig",
    package = "PureCN", mustWork = TRUE)
intervals <- import(bed.file)
mappability <- import(bed.file)
preprocessIntervals(intervals, reference.file,
    mappability = mappability, output.file = "ex2_gc_file.txt")
## WARN [2024-05-01 18:54:13] Found small target regions (< 100bp). Will resize them.
## INFO [2024-05-01 18:54:14] Splitting 5 large targets to an average width of 400.
## WARN [2024-05-01 18:54:14] No reptiming scores provided.
## INFO [2024-05-01 18:54:14] Calculating GC-content...
```

A command line script described in a separate vignette provides convenient access to this function and also attempts to annotate the targets with gene symbols using the annotate Targets function.

2.3 Coverage data

The calculateBamCoverageByInterval function can be used to generate the required coverage data from BAM files. All we need to do is providing the desired intervals (as generated by preprocessIntervals):

```
bam.file <- system.file("extdata", "ex1.bam", package="PureCN",
    mustWork = TRUE)
interval.file <- system.file("extdata", "ex1_intervals.txt",
    package = "PureCN", mustWork = TRUE)
```

```
calculateBamCoverageByInterval(bam.file = bam.file,
    interval.file = interval.file, output.file = "ex1_coverage.txt")
```

2.4 Third-party coverage tools

Calculating coverage from BAM files is a common task and your pipeline might already provide this information. As alternative to calculateBamCoverageByInterval, *PureCN* currently supports coverage files generated by *GATK3 DepthOfCoverage*, *GATK4 CollectFragmentCounts* and by *CNVkit*. By providing files with standard file extension, *PureCN* will automatically detect the correct format and all following steps are the same for all tools. You will, however, still need the interval file generated in Section 2.2 and the third-party tool must use the exact same intervals. See also FAQ Section 12.2 for recommended settings for *GATK3 DepthOfCoverage*.

2.5 Third-party segmentation tools

PureCN integrates well with existing copy number pipelines. Instead of coverage data, the user then needs to provide either already segmented data or a wrapper function. This is described in Section 10.1.

2.6 Example data

We now load a few example files that we will use throughout this tutorial:

```
library(PureCN)
normal.coverage.file <- system.file("extdata", "example_normal.txt.gz",
    package = "PureCN")
normal2.coverage.file <- system.file("extdata", "example_normal2.txt.gz",
    package = "PureCN")
normal.coverage.files <- c(normal.coverage.file, normal2.coverage.file)
tumor.coverage.file <- system.file("extdata", "example_tumor.txt.gz",
    package = "PureCN")
seg.file <- system.file("extdata", "example_seg.txt",
    package = "PureCN")
vcf.file <- system.file("extdata", "example.vcf.gz", package = "PureCN")
interval.file <- system.file("extdata", "example_intervals.txt",
    package = "PureCN")</pre>
```

3 Library-specific coverage bias

In coverage normalization, we distinguish between assay- and library-specific biases. Assayspecific biases, for example due to probe density, probe capture efficiency and read mappability, are best removed with a pool of normal samples (Section 4.1, [5]). In other words, by examining the coverage of particular intervals in a pool of normals, we can estimate how well this assay captures these intervals and will then adjust the tumor coverage accordingly.

Other biases are library-specific, meaning a patient sample captured in different libraries may display dramatically different coverage profiles across libraries. Data from great sequencing centers usually show relatively small technical variance nowadays, but some biases are not completely avoidable. The most important library-specific bias is due to GC-content, i.e. regions of high AT- or GC-content are not always captured with exactly the same efficiency in tumor and normals.

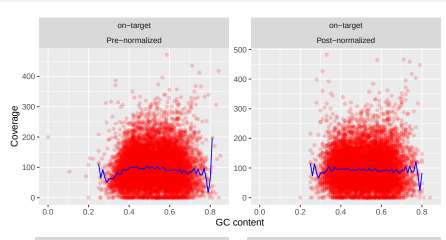
We usually also observe that early replicating regions have a slightly higher coverage than late replicating regions [6, 7]. Since there is often a significant difference in proliferation rates of tumor and normal, the pool of normals might also not completely adjust for this small bias.

A good panel of normal samples will remove both assay- and library-specific biases. We recommend sequencing the normals in multiple batches so that the coverage normalization has access to any expected day-to-day assay variance. This will greatly increase the normalization's capability in removing library-specific biases.

Sometimes it can be beneficial to remove library-specific biases explicitly, especially with smaller panels of normals that might show significant differences to tumors (FFPE vs blood, high vs low coverage, etc.).

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In the following example, we correct the raw coverage of all samples, tumor and normal, for the two major sources of library-specific coverage biases mentioned above (Figure 1). For GC-normalization, we use a 2-step loess normalization [8]. For the replication timing bias, a linear model of log-transformed coverage and provided replication timing score is used.



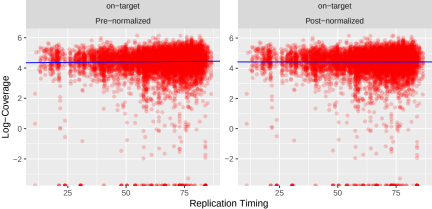


Figure 1: Coverage before and after normalization for GC-content and replication timing. This plot shows coverage as a function of on- and off-target GC-content and replication timing before and after normalization. Each dot is an interval. The example files are already GC-normalized; real data will show more dramatic differences.

The example coverage files are already GC-normalized. We provide a convenient command line script for generating normalized coverage data from BAM files or from *GATK* coverage files (see Quick vignette).

4 Pool of normals

4.1 Coverage normalization

For calculating copy number log2-ratios of tumor vs. normal, *PureCN* requires coverage from a process-matched normal sample. Using a normal that was sequenced using a similar, but not identical assay, rarely works. Differently covered genomic regions simply result in too many log2-ratio outliers. This section describes how to optimally normalize coverage against a pool of normals.

The createNormalDatabase function builds a database of coverage files (a command line script providing this functionality is described in a separate vignette):

```
normalDB <- createNormalDatabase(normal.coverage.files)</pre>
```

```
## INF0 [2024-05-01 18:54:25] 576 on-target bins with low coverage in all samples.
## WARN [2024-05-01 18:54:25] You are likely not using the correct baits file!
## WARN [2024-05-01 18:54:25] Allosome coverage missing, cannot determine sex.
## WARN [2024-05-01 18:54:25] Allosome coverage missing, cannot determine sex.
## INF0 [2024-05-01 18:54:25] Processing on-target regions...
## INF0 [2024-05-01 18:54:25] Removing 930 intervals with low coverage in normalDB.
## INF0 [2024-05-01 18:54:25] Removing 1 intervals with zero coverage in more than 3% of normalDB.
```

```
# serialize, so that we need to do this only once for each assay
saveRDS(normalDB, file = "normalDB.rds")
```

When using a GC-normalization in tumors, please make sure that all provided coverage files here were also GC-normalized prior to building the database (Section 3). Internally, creat eNormalDatabase determines the sex of the samples and trains a PCA that is later used for denoising tumor coverage using Tangent normalization [9]:

```
normalDB <- readRDS("normalDB.rds")
pool <- calculateTangentNormal(tumor.coverage.file, normalDB)</pre>
```

This createNormalDatabase function further automatically estimates copy number log2-ratio standard deviations. Assuming that all normal samples are in general diploid, a high variance in log2-ratio is indicative of an interval with either common germline alterations or frequent artifacts; high or low copy number log2-ratios in these intervals are unlikely measuring somatic copy number events. The segmentation function can use this information to skip over such noisy regions.

4.2 Artifact filtering

It is important to remove as many artifacts as possible, because low ploidy solutions are typically punished more by artifacts than high ploidy solutions. High ploidy solutions are complex and usually find ways of explaining artifacts reasonably well. The following steps in this section are optional, but recommended since they will reduce the number of samples requiring manual curation, especially when matching normal samples are not available.

4.2.1 VCF

We recommend running *MuTect* with a pool of normal samples to filter common sequencing errors and alignment artifacts from the VCF. *MuTect* requires a single VCF containing all normal samples, for example generated by the *GATK3 CombineVariants* tool (see Section 12.2). For *Mutect 2* in *GATK4*, please follow the somatic best practices including the generation of a *GenomicsDB* containing variants from the normal samples.

It is highly recommended to provide *PureCN* these variant pools of normals as well; it will help the software correcting non-reference read mapping biases. This is described in the setMappingBiasVcf documentation. To reduce memory usuage, the normal panel VCF can be reduced to contain only variants present in 2-3 or more samples. *GenomicsDBs* from *GATK4* can be provided as is, but require the installation of the *genomicsdb* R package from https://github.com/nalinigans/GenomicsDB-R.

Because these pools can become huge with increasing number of normal samples, we precompute the mapping bias:

This function will first fit beta-binomial distributions of allelic fractions from heterozygous variants (between fractions of 0.1 and 0.9) observed in sufficient number of samples; by default min.normals.betafit is set to 7. Under the assumption that there are only a limited number of error modes resulting in different distributions, we next cluster those position-specific distributions into num.betafit.clusters (default 9). Next, all variants observed heterozygous in less than min.normals.betafit but equal or more than min.normals.assign.betafit (default 3) are assigned the best fitting beta-distribution cluster.

For variants with at least min.normals (default 1) but less than min.normals.assign.betafit, the empirical Bayes estimate described in the original *PureCN* paper is used where the reference and alternate counts of an average high quality SNP is added to the observed counts, thus forcing the estimate to the average when position-specific information is limited.

Note that variants with low median coverage (min.median.coverage.betafit, default 5) are not attempted to fit to beta-binomal distributions and instead the empirical Bayes method is used.

4.3 Artifact filtering without a pool of normals

By default, *PureCN* will exclude targets with coverage below 15X from segmentation (with a pool of normals, targets are filtered based on the coverage and variance in normal database only). For variants in the provided VCF, the same 15X cutoff is applied. *MuTect* applies more sophisticated artifact tests and flags suspicious variants. If *MuTect* was run in matched normal mode, then both potential artifacts and germline variants are rejected, that means

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we cannot just filter by the PASS/REJECT *MuTect* flags. The filterVcfMuTect function optionally reads the *MuTect* 1.1.7 stats file and will keep germline variants, while removing potential artifacts. Without the stats file, *PureCN* will use only the filters based on read depths as defined in filterVcfBasic. Both functions are automatically called by *PureCN*, but can be easily modified and replaced if necessary.

We can also use a BED file to blacklist regions expected to be problematic, for example the simple repeats track from the UCSC:

```
# Instead of using a pool of normals to find low quality regions,
# we use suitable BED files, for example from the UCSC genome browser.
# We do not download these in this vignette to avoid build failures
# due to internet connectivity problems.
downloadFromUCSC <- FALSE
if (downloadFromUCSC) {
    library(rtracklayer)
    mySession <- browserSession("UCSC")
    genome(mySession) <- "hg19"
    simpleRepeats <- track(ucscTableQuery(mySession,
        track = "Simple Repeats", table = "simpleRepeat"))
    export(simpleRepeats, "hg19_simpleRepeats.bed")
}
```

```
snp.blacklist <- "hg19_simpleRepeats.bed"</pre>
```

5 Recommended run

Finally, we can run *PureCN* with all that information:

```
ret <-runAbsoluteCN(normal.coverage.file = pool,</pre>
    tumor.coverage.file = tumor.coverage.file, vcf.file = vcf.file,
    genome = "hg19", sampleid = "Sample1",
    interval.file = interval.file, normalDB = normalDB,
#
    args.setMappingBiasVcf=list(
        # mapping.bias.file = mapping.bias.file
#
    ),
    args.filterVcf=list(
        # snp.blacklist = snp.blacklist,
        # stats.file = mutect.stats.file
    ),
    args.filterIntervals = list(
        min.total.counts = 50
    ),
    post.optimize = FALSE, plot.cnv = FALSE, verbose = FALSE)
## WARN [2024-05-01 18:54:30] Allosome coverage missing, cannot determine sex.
## WARN [2024-05-01 18:54:30] Allosome coverage missing, cannot determine sex.
```

The normal.coverage.file argument points to a coverage file obtained from either a matched or a process-matched normal sample, but can be also a small pool of best normals (Section 4.1).

The normalDB argument (Section 4.1) provides a pool of normal samples and for example allows the segmentation function to skip targets with low coverage or common germline deletions in the pool of normals. If available, a VCF containing all variants from the normal samples should be provided via args.setMappingBiasVcf to correct read mapping biases. The files specified in args.filterVcf help *PureCN* filtering SNVs more efficiently for artifacts as described in Sections 4.2 and 4.3. The snp.blacklist is only necessary if neither a matched normal nor a large pool of normals is available. We set min.total.counts to a very low value to make this toy example use all simulated intervals.

The post.optimize flag will increase the runtime by about a factor of 2-5, but might return slightly more accurate purity estimates. For high quality whole-exome data, this is typically not necessary for copy number calling (but might be for variant classification, see Section 6.2.1). For smaller targeted panels, the runtime increase is typically marginal and post.optimize should be always set to TRUE.

The plot.cnv argument allows the segmentation function to generate

additional plots if set to TRUE. Finally, verbose outputs important and helpful information about all the steps performed and is therefore set to TRUE by default.

6 Output

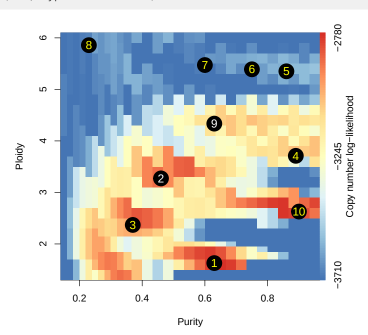
6.1 Plots

We now create a few output files:

```
file.rds <- "Sample1_PureCN.rds"</pre>
saveRDS(ret, file = file.rds)
pdf("Sample1_PureCN.pdf", width = 10, height = 11)
plotAbs(ret, type = "all")
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## Warning in par(par.mar): argument 1 does not name a graphical parameter
dev.off()
## pdf
##
    2
```

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The RDS file now contains the serialized return object of the runAbsoluteCN call. The PDF contains helpful plots for all local minima, sorted by likelihood. The first plot in the generated PDF is displayed in Figure 2 and shows the purity and ploidy local optima, sorted by final likelihood score after fitting both copy number and allelic fractions.



plotAbs(ret, type = "overview")

Figure 2: Overview. The colors visualize the copy number fitting score from low (blue) to high (red). The numbers indicate the ranks of the local optima. Yellow fonts indicate that the corresponding solutions were flagged, which does not necessarily mean the solutions are wrong. The correct solution (number 1) of this toy example was flagged due to large amount of LOH.

We now look at the main plots of the maximum likelihood solution in more detail.

```
plotAbs(ret, 1, type = "hist")
## Warning in par(par.mar): argument 1 does not name a graphical parameter
## NULL
```

Figure 3 displays a histogram of tumor vs. normal copy number log2-ratios for the maximum likelihood solution (number 1 in Figure 2). The height of a bar in this plot is proportional to the fraction of the genome falling into the particular log2-ratio copy number range. The vertical dotted lines and numbers visualize the, for the given purity/ploidy combination, expected log2-ratios for all integer copy numbers from 0 to 7. It can be seen that most of the log2-ratios of the maximum likelihood solution align well to expected values for copy numbers of 0, 1, 2 and 4.

plotAbs(ret, 1, type = "BAF")

Germline variant data are informative for calculating integer copy number because unbalanced maternal and paternal chromosome numbers in the tumor portion of the sample lead to unbalanced germline allelic fractions. Figure 4 shows the allelic fractions of predicted germline SNPs. The goodness of fit (GoF) is provided on an arbitrary scale in which 100% corresponds to a perfect fit and 0% to the worst possible fit. The latter is defined as a fit in which allelic

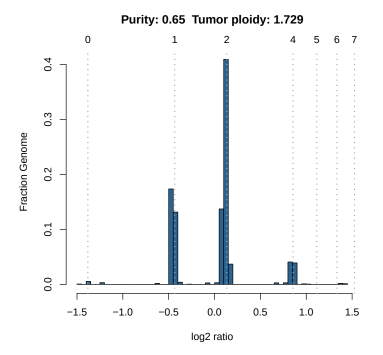


Figure 3: Log-ratio histogram.

fractions on average differ by 0.2 from their expected fractions. Note that this does not take purity into account and low purity samples are expected to have a better fit. In the middle panel, the corresponding copy number log2-ratios are shown. The lower panel displays the calculated integer copy numbers, corrected for purity and ploidy. We can zoom into particular chromosomes (Figure 5).

plotAbs(ret, 1, type = "BAF", chr = "chr19")
plotAbs(ret, 1, type = "AF")

Finally, Figure 6 provides more insight into how well the variants fit the expected values.

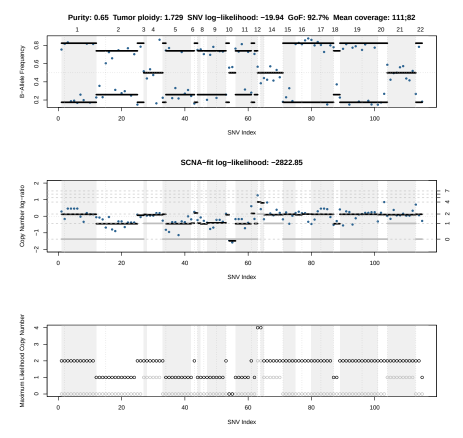


Figure 4: B-allele frequency plot. Each dot is a (predicted) germline SNP. The first panel shows the allelic fractions as provided in the VCF file. The alternating grey and white background colors visualize odd and even chromosome numbers, respectively. The black lines visualize the expected (not the average!) allelic fractions in the segment. These are calculated using the estimated purity and the total and minor segment copy numbers. These are visualized in black and grey, respectively, in the second and third panel. The second panel shows the copy number log2-ratios, the third panel the integer copy numbers.

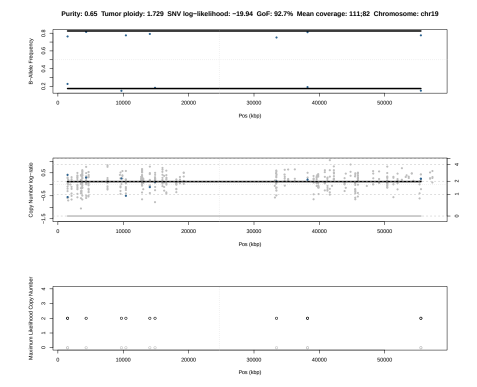


Figure 5: Chromosome plot. Similar to Figure 4, but zoomed into a particular chromosome. The grey dots in the middle panel visualize copy number log2-ratios of targets without heterozygous SNPs, which are omitted from the previous genome-wide plot. The x-axis now indicates genomic coordinates in kbps.

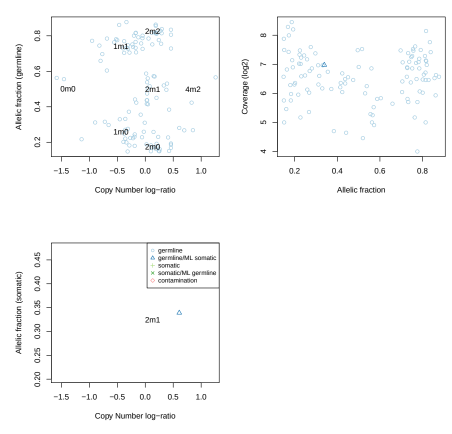


Figure 6: Allele fraction plots. Each dot is again a (predicted) germline SNP. The shapes visualize the different SNV groups based on prior and posterior probabilities. The labels show the expected values for all called states; 2m1 would be diploid, heterozygous, 2m2 diploid, homozygous. The relationship of allelic fraction and coverage is shown in the top right panel. This plot normally also shows somatic mutations in two additional panels, with the left panel showing the same plot as for germline SNPs and the bottom right panel a histogram of cellular fraction of predicted somatic mutations. This toy example contains only germline SNPs however.

6.2 Data structures

The R data file (file.rds) contains gene-level copy number calls, SNV status and LOH calls. The purity/ploidy combinations are sorted by likelihood and stored in ret\$results.

names(ret)
[1] "candidates" "results" "input" "version"

We provide convenient functions to extract information from this data structure and show their usage in the next sections. We recommend using these functions instead of accessing the data directly since data structures might change in future versions.

6.2.1 Prediction of somatic status and cellular fraction

To understand allelic fractions of particular SNVs, we must know the (i) somatic status, the (ii) tumor purity, the (iii) local copy number, as well as the (iv) number of chromosomes harboring the mutations or SNPs. One of *PureCN* main functions is to find the most likely combination of these four values. We further assign posterior probabilities to all possible combinations or states. Availability of matched normals reduces the search space by already providing somatic status.

The predictSomatic function provides access to these probabilities. For predicted somatic mutations, this function also provides cellular fraction estimates [10], i.e. the fraction of tumor cells with mutation. Fractions significantly below 1 indicate sub-clonality:

```
head(predictSomatic(ret), 3)
```

```
##
      chr
              start
                                             ID REF ALT
                                                            SOMATIC.M0
                           end
## 1 chr1 114515871 114515871 chr1114515871xxx
                                                       A 4.046059e-102
                                                  G
## 2 chr1 150044293 150044293 chr1150044293xxx
                                                       G
                                                          8.631774e-92
                                                  Т
## 3 chr1 158449835 158449835 chr1158449835xxx
                                                       G 1.248635e-147
                                                  А
##
       SOMATIC.M1
                    SOMATIC.M2
                                   SOMATIC.M3 SOMATIC.M4 SOMATIC.M5 SOMATIC.M6
## 1 8.420377e-39 2.107876e-07 5.330398e-310
                                                        0
                                                                   0
                                                                               0
## 2 4.835490e-39 2.452234e-10 4.617747e-312
                                                        0
                                                                   0
                                                                               0
                                                                   0
                                                                               0
## 3 2.177662e-62 5.337189e-15 5.337599e-314
                                                        0
                  GERMLINE.M0 GERMLINE.M1 GERMLINE.M2
##
     SOMATIC.M7
                                                           GERMLINE.M3
## 1
              0 1.607523e-69 6.946405e-16
                                              0.9999998 7.018564e-308
## 2
              0 3.623805e-64 1.081334e-18
                                              1.0000000 3.778064e-306
              0 1.172374e-105 5.100431e-30
                                              1.0000000 1.157132e-305
## 3
##
     GERMLINE.M4 GERMLINE.M5 GERMLINE.M6 GERMLINE.M7 GERMLINE.CONTHIGH
## 1
               0
                            0
                                        0
                                                     0
                                                            5.799092e-42
## 2
               0
                            0
                                        0
                                                     0
                                                            1.891829e-20
## 3
               0
                            0
                                        0
                                                     0
                                                            1.087505e-25
##
     GERMLINE.CONTLOW GERMLINE.HOMOZYGOUS ML.SOMATIC POSTERIOR.SOMATIC ML.M
## 1
        8.219408e-289
                                         0
                                                 FALSE
                                                            2.107876e-07
                                                                             2
## 2
        2.345594e-243
                                         0
                                                 FALSE
                                                            2.452234e-10
                                                                             2
         0.000000e+00
                                         0
                                                 FALSE
                                                                             2
## 3
                                                            5.337189e-15
##
     ML.C ML.M.SEGMENT M.SEGMENT.POSTERIOR M.SEGMENT.FLAGGED ML.AR
                                                                            AR
## 1
        2
                     0
                                          1
                                                         FALSE 0.825 0.755183
## 2
        2
                     0
                                          1
                                                         FALSE 0.825 0.817078
                                          1
                                                         FALSE 0.825 0.834266
## 3
        2
                     0
     AR.ADJUSTED MAPPING.BIAS ML.LOH CN.SUBCLONAL CELLFRACTION
##
## 1
       0.7775993
                    0.9711724
                                 TRUE
                                             FALSE
                                                              NA
```

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0.8413316 ## 2 0.9711724 TRUE FALSE NA ## 3 0.8590298 TRUE 0.9711724 FALSE NA ## CELLFRACTION.95.LOWER CELLFRACTION.95.UPPER ALLELIC.IMBALANCE FLAGGED ## 1 NA NA -17,46605 FALSE ## 2 NA NA -20.15393 FALSE ## 3 NA NA -33.34446 FALSE ## log.ratio depth prior.somatic prior.contamination on.target seg.id ## 1 0.2842136 184 9.9e-05 0.01 1 1 ## 2 -0.1686186 138 9.9e-05 0.01 1 1 ## 3 0.4596841 0.01 1 1 217 9.9e-05 ## pon.count gene.symbol ## 1 0 HIPK1 ## 2 0 VPS45 ## 3 0 <NA>

The output columns are explained in Table 1.

To annotate the input VCF file with these values:

vcf <- predictSomatic(ret, return.vcf = TRUE)
writeVcf(vcf, file = "Sample1_PureCN.vcf")</pre>

For optimal classification results:

- Set post.optimize = TRUE since small inaccuracies in purity can decrease the classification performance significantly
- Provide args.setMappingBias a pool of normal VCF to obtain position-specific mapping bias information
- Exclude variants in regions of low mappability
- Use a somatic posterior probability cutoff of 0.8 and 0.2 for somatic and germline variants, respectively. This appears to be a good compromise of call rate and accuracy. If the beta-binomial model was selected in the model argument of runAbsoluteCN with small panel of normals, these cutoffs might need to be relaxed to get acceptable call rates.
- Add a Cosmic.CNT info field to the VCF or provide a COSMIC VCF in runAbsoluteCN (see Section 10.2).

Note that the posterior probabilities assume that the purity and ploidy combination is correct. Before classifying variants, it is thus recommended to manually curate flagged samples.

6.2.2 Amplifications and deletions

To call amplifications, we recommend using a cutoff of 6 for focal amplifications and a cutoff of 7 otherwise. For homozygous deletions, a cutoff of 0.5 is useful to allow some heterogeneity in copy number.

For samples that failed *PureCN* calling we recommended using common log2-ratio cutoffs to call amplifications, for example 0.9.

This strategy is implemented in the callAlterations function:

Column name	Description
chr, start, end	Variant coordinates
ID	The variant ID as provided in the VCF
REF, ALT	The reference and alt alleles
SOMATIC.M*	Posterior probabilities for all somatic states. M0 to M7 are mul-
GERMLINE.M*	tiplicity values, i.e. the number of chromosomes harboring the mutation (e.g. 1 heterozygous, 2 homozygous if copy number C is 2). SOMATIC.M0 represents a sub-clonal state (somatic mutations by definition have a multiplicity larger than 0). Posterior probabilities for all heterozygous germline states
GERMLINE.CONTHIGH	Posterior probability for contamination. This state corresponds
GERMEINE . CONTINUIT	to homozygous germline SNPs that were not filtered out because reference alleles from another individual were sequenced, resulting in allelic fractions smaller than 1.
GERMLINE.CONTLOW	Posterior probability for contamination. This state corresponds to non-reference alleles only present in the contamination.
GERMLINE.HOMOZYGOUS	Posterior probability that SNP is homozygous in normal. Requires
	the model.homozygous option in runAbsoluteCN. See Section 8.
ML.SOMATIC	TRUE if the maximum likelihood state is a somatic state
POSTERIOR.SOMATIC	The posterior probability that the variant is somatic (sum of a
	somatic state posterior probabilities)
ML.M	Maximum likelihood multiplicity
ML.C	Maximum likelihood integer copy number
ML.M.SEGMENT	Maximum likelihood minor segment copy number
M.SEGMENT.POSTERIOR	Posterior probability of ML.M.SEGMENT
M.SEGMENT.FLAGGED	Segment flag indicating ML.M.SEGMENT is unreliable, eithe due to low posterior probability (< 0.5) or few variants ($<$
	min.variants.segment). Indels are always flagged.
ML.AR	Expected allelic fraction of the maximum likelihood state
AR	Observed allelic fraction (as provided in VCF)
AR.ADJUSTED	Observed allelic fraction adjusted for mapping bias TRUE if variant is most likely in LOH
ML.LOH CN.SUBCLONAL	TRUE if copy number segment is sub-clonal
CELLFRACTION	Fraction of tumor cells harboring the somatic mutation [10]
CELLFRACTION CELLFRACTION.95.LOWER	Lower 95% of confidence interval
CELLFRACTION.95.UPPER	Upper 95% of confidence interval
ALLELIC.IMBALANCE	Log-likelihood that variant is in allelic balance. Requires position
ALLELICITIDALANCE	specific mapping bias information to be reliable.
FLAGGED	Flag indicating that call is unreliable (currently only due to high mapping bias and high pool of normal counts)
log.ratio	The copy number log2-ratio (tumor vs. normal) for this variant
depth	The total sequencing depth at this position
prior.somatic	Prior probability of variant being somatic
prior.contamination	Prior probability that variant is contamination from another indi vidual
on.target	1 for variants within intervals, 2 for variants in flanking regions 0 for off-target variants
seg.id	Segment id
pon.count	Number of hits in the mapping.bias.file
gene.symbol	Gene symbol if available

 Table 1: predictSomatic
 output
 columns.

```
gene.calls <- callAlterations(ret)</pre>
head(gene.calls)
##
                                 end C C.flagged seg.mean seg.id
             chr
                     start
## EIF2A
                                                               5
            chr3 150270143 150301699 6
                                           FALSE
                                                   1.3859
                                                               5
## MIR548H2 chr3 151531951 151538241 6
                                           FALSE
                                                   1.3859
## AADAC
            chr3 151542451 151545961 6
                                                   1.3859
                                                               5
                                           FALSE
## GPNMB
            chr7 23286477 23313844 7
                                           FALSE
                                                   1.4462
                                                              19
## SH2D4B chr10 82329910 82403838 0
                                                              26
                                           FALSE -1.4863
## SLC35G1 chr10 95653791 95661248 0
                                           FALSE -1.2007
                                                              28
##
           number.targets gene.mean gene.min gene.max focal breakpoints
## EIF2A
                       13 1.5643189 0.7544062 2.1392167 TRUE
                                                                           0
## MIR548H2
                       3 0.8261594 0.4682628 1.2000253 TRUE
                                                                           0
                        2 0.7851630 0.6864438 0.8838822 TRUE
## AADAC
                                                                           1
## GPNMB
                       11 1.4462179 0.9397072 1.8200775 TRUE
                                                                           0
                                                                           0
## SH2D4B
                        6 -1.4862885 -1.5836964 -1.4113313 FALSE
## SLC35G1
                        3 -1.5817134 -1.8066816 -1.4493561 FALSE
                                                                           0
##
                    type num.snps M M.flagged
                                                loh
## EIF2A
           AMPLIFICATION
                                0 NA
                                            NA
                                                 NA
## MIR548H2 AMPLIFICATION
                                0 NA
                                                 NA
                                            NA
           AMPLIFICATION
## AADAC
                                0 NA
                                            NA
                                                 NA
## GPNMB
           AMPLIFICATION
                                0 NA
                                            NA
                                                 NA
## SH2D4B
                DELETION
                                2 0
                                          TRUE TRUE
## SLC35G1
                DELETION
                                0 NA
                                            NA
                                                 NA
```

It is also often useful to filter the list further by known biology, for example to exclude non-focal amplifications of tumor suppressor genes. The Sanger Cancer Gene Census [11] for example provides such a list.

The output columns of callAlterations are explained in Table 2.

6.2.3 Amplifications in low purity samples

The default callAlterations is not suited for samples of very low purity below 10% such as frequently observed in cfDNA samples. To call high level amplifications in samples between 4 to 10%, we recommend a simple z-score approach implemented in callAmplificationsIn LowPurity. This will calculate gene-level log2 copy number ratio standard deviations in all normal samples that are then used to obtain gene-level cutoffs for amplifications in tumor samples. The output of this function is nearly identical to callAlterations with additional columns explained in Table 3.

gene.calls.zscore <- callAmplificationsInLowPurity(ret, normalDB)</pre>

WARN [2024-05-01 18:55:46] Extensive noise in tumor compared to normals.

```
head(gene.calls.zscore)
```

##		chr	start	end	С	C.flagged	number.targets	gene.mean
##	EIF2A	chr3	150270143	150301699	NA	FALSE	13	1.5643189
##	MIR548H2	chr3	151531951	151538241	NA	FALSE	3	0.8261594
##	FTLP10	chr4	69056959	69078196	NA	FALSE	3	0.5852635
##	FGF5	chr4	81187979	81207827	NA	FALSE	3	1.0400133
##	SPP1	chr4	88898048	88904049	NA	FALSE	5	0.7200921
##	PPM1K	chr4	89183747	89199736	NA	FALSE	6	0.8253634

Column name	Description
chr, start, end	Gene coordinates
С	Segment integer copy number. See also seg.id.
C.flagged	Flagged when segment interval weights are low. Requires normalDB. In case multiple segments overlap, C.flagged will be TRUE when any segment is flagged.
seg.mean	Segment mean of copy number log2-ratios (not adjusted for pu- rity/ploidy). See also seg.id.
seg.id	Segment id. In case multiple segments overlap, seg.id will point to the smallest overlapping segment and the column breakpoints will be non-zero.
number.targets	Number of targets annotated with this symbol
gene.*	Gene copy number log2-ratios (not adjusted for purity/ploidy). gene.mean is weighted by interval weights when these are avail- able.
focal	TRUE for focal amplification, as defined by the <pre>fun.focal</pre> argument in <pre>runAbsoluteCN</pre>
breakpoints	Number of breakpoints between start and end
type	Amplification or deletion
num.snps	Number of SNPs in this segment informative for LOH detection (requires VCF)
Μ	minor copy number of segment (requires VCF)
M.flagged	flag indicating that M is unreliable (requires VCF)
loh	TRUE if segment is in LOH, FALSE if not and NA if number of SNPs is insufficient (requires VCF)

 Table 2: callAlterations
 output
 columns.

##		gene.min	gene.max	breakpoints	p.value	percentile.genome	
##	EIF2A	0.7544062	2.139217	0	Θ	100.00000	
##	MIR548H2	0.4682628	1.200025	0	Θ	95.05495	
##	FTLP10	0.4369763	1.262028	0	Θ	90.76923	
##	FGF5	0.8330433	1.249963	0	Θ	98.35165	
##	SPP1	0.4282286	1.061844	0	Θ	92.30769	
##	PPM1K	0.1361852	1.456415	0	Θ	94.94505	
##		percentile	e.chromoso	ome	type		
##	EIF2A		100.000	000 AMPLIFICA	ATION		
##	MIR548H2		95.652	217 AMPLIFICA	ATION		
##	FTLP10		77.500	000 AMPLIFICA	ATION		
##	FGF5		97.500	000 AMPLIFICA	ATION		
##	SPP1		82.500	000 AMPLIFICA	ATION		
##	PPM1K		92.500	000 AMPLIFIC	ATION		
	<pre># filter to known amplifications known.calls.zscore <- gene.calls.zscore[gene.calls.zscore\$gene.symbol %in</pre>						
5							

.n% c("AR", "BRAF", "CCND1", "CCNE1", "CDK4", "CDK6", "EGFR", "ERBB2", "FGFR1", "FGFR2", "KIT", "KRAS", "MCL1", "MDM2", "MDM4", "MET", "MYC", "PDGFRA", "PIK3CA", "RAF1"),]

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Since this algorithm is intended to rescue known high-level amplifications in very low tumor purities, we recommend using a liberal p-value cutoff, but restricting the search to a limited set of genes.

 Table 3: callAmplificationsInLowPurity
 output
 columns
 not
 already
 explained
 in
 Table 2.

Column name	Description
p.value	P-value of observing such a gene-level log2-ratio in normal samples
percentile.genome	Percentile of gene-level log2-ratio in the tumor sample. Useful for distinguishing focal amplifications from broad gains.
percentile.chromosome	Same as percentile.genome, but for the chromosome

6.2.4 Find genomic regions in LOH

The gene.calls data.frame described above provides gene-level LOH information. To find the corresponding genomic regions in LOH, we can use the callLOH function:

```
loh <- callLOH(ret)</pre>
head(loh)
##
     chr
             start
                        end arm C M
                                                         type
                                                                seg.mean
## 1 chr1 114515871 121535434 p 2 0 WHOLE ARM COPY-NEUTRAL LOH 0.1173894
## 2 chr1 124535434 247419499 q 2 0 WHOLE ARM COPY-NEUTRAL LOH 0.1173894
## 3 chr2 10262881 92326171 p 1 0
                                                WHOLE ARM LOH -0.4517019
## 4 chr2 95326171 231775678 q 1 0
                                                 WHOLE ARM LOH -0.4517019
## 5 chr2 236403331 239039169 q 2 0
                                             COPY-NEUTRAL LOH 0.0579000
## 6 chr3 11888043 90504854 p 2 1
                                                               0.1056000
##
    num.mark num.snps M.flagged maf.expected maf.observed
## 1
         886
                  11
                         FALSE
                                  0.1750000
                                               0.1767647
## 2
         886
                   11
                         FALSE
                                  0.1750000
                                               0.1767647
## 3
         657
                   13
                         FALSE
                                  0.2592593
                                               0.2751720
## 4
         657
                   13
                          FALSE
                                  0.2592593
                                               0.2751720
## 5
         86
                   2
                          TRUE
                                  0.1750000
                                               0.1719800
         403
                          TRUE
## 6
                   1
                                  0.5000000
                                               0.4685382
```

The output columns are explained in Table 4.

Column name	Description
chr, start, end	Segment coordinates
arm	Chromosome arm
С	Segment integer copy number
Μ	Minor integer copy number $(M + N = C, M \leq N)$
type	LOH type if $M = 0$
seg.mean	Segment mean of copy number log2-ratios (not adjusted for purity/ploidy)
num.mark	Number of intervals in this segment (following DNAcopy naming convention)
num.snps	Number of SNPs in this segment informative for LOH detection
M.flagged	flag indicating that M is unreliable (due to small number of variants defined by min.variants.segment or because of ambiguous M call)
<pre>maf.expected</pre>	Expected minor allele frequency of heterozygous SNPs in this seg- ment
maf.observed	Median of observed minor allele frequency of heterozygous SNPs in this segment

 Table 4: callLOH output columns.

7 Curation

For prediction of variant status (germline vs. somatic, sub-clonal vs. clonal, homozygous vs. heterozygous), it is important that both purity and ploidy are correct. We provide functionality for curating results:

createCurationFile(file.rds)

This will generate a CSV file in which the correct purity and ploidy values can be manually entered. It also contains a column "Curated", which should be set to TRUE, otherwise the file will be overwritten when re-run.

Then in R, the correct solution (closest to the combination in the CSV file) can be loaded with the readCurationFile function:

ret <- readCurationFile(file.rds)</pre>

This function has various handy features, but most importantly it will re-order the local optima so that the curated purity and ploidy combination is ranked first. This means plotAbs(ret,1,type="hist") would show the plot for the curated purity/ploidy combination, for example.

The default curation file will list the maximum likelihood solution:

```
read.csv("Sample1_PureCN.csv")
## Sampleid Purity Ploidy Sex Contamination Flagged Failed Curated
## 1 Sample1 0.65 1.728527 ? 0 TRUE FALSE FALSE
## Comment
## 1 EXCESSIVE LOSSES;EXCESSIVE LOH
```

PureCN currently only flags samples with warnings, it does not mark any samples as failed. The Failed column in the curation file can be used to manually flag samples for exclusion in downstream analyses. See Table 5 for an explanation of all flags.

Flag	Description
EXCESSIVE LOH	>50% of genome in LOH and ploidy $<$
	2.6
EXCESSIVE LOSSES	$\geq 1\%$ of genome deleted
HIGH AT- OR GC-DROPOUT	High GC-bias exceeding cutoff in
	max.dropout
HIGH PURITY	(when model.homozygous=FALSE). For
	very high purity samples, it is recom-
	mended to set model.homozygous=TRUE.
	See Section 8.
LOW PURITY	Purity < 30%
LOW BOOTSTRAP VALUE	bootstrapResults identified multiple
	plausible solutions
NOISY LOG-RATIO	Log-ratio standard deviation $>$
	max.logr.sdev
NOISY SEGMENTATION	More than max.segments
NON-ABERRANT	$\geq 99\%$ of genome has identical copy
	number and $\geq~0.5\%$ has second most
	common state
POLYGENOMIC	$\geq 0.75 \times$ max.non.clonal fraction of the
	genome in sub-clonal state
POOR GOF	GoF < min.gof
POTENTIAL SAMPLE CONTAMINATION	Significant portion of variants present in
	germline databases are potentially cross-
	contaminated
RARE KARYOTYPE	Ploidy < 1.5 or > 4.5

 Table 5: createCurationFile
 flags.

8 Cell lines

Default parameters assume some normal contamination. In 100% pure samples without matching normal samples such as cell lines, we cannot distinguish homozygous SNPs from LOH by looking at single allelic fractions alone. It is thus necessary to keep homozygous variants and include this uncertainty in the likelihood model. This is done by setting the runAbsoluteCN argument model.homozygous=TRUE. If matched normals are available, then variants homozygous in normal are automatically removed since they are uninformative.

For technical reasons, the maximum purity *PureCN* currently models is 0.99. We recommend setting test.purity=seq(0.9,0.99,by=0.01) in runAbsoluteCN for cell lines.

Please note that in order to detect homozygous deletions in 100% pure samples, you will need to provide a normalDB in runAbsoluteCN to filter low quality targets efficiently (Section 5).

9 Maximizing the number of heterozygous SNPs

It is possible to use SNPs in off-target reads in the variant fitting step by running *MuTect* without interval file and then setting the filterVcfBasic argument remove.off.target.snvs to FALSE. We recommend a large pool of normals in this case and then generating SNP blacklists as described in Sections 4.2 and 4.3. Remember to also run all the normals in *MuTect* without interval file.

An often better alternative to including all off-target reads is only including variants in the flanking regions of targets (between 50-100bp). This will usually significantly increase the number of heterozygous SNPs (see Section 12.2). These SNPs are automatically added if the variant caller was run without interval file or with interval padding.

10 Advanced usage

10.1 Custom normalization and segmentation

Copy number normalization and segmentation are crucial for obtaining good purity and ploidy estimates. If you have a well-tested pipeline that produces clean results for your data, you might want to use *PureCN* as add-on to your pipeline. By default, we will use *DNAcopy* [12] to segment normalized target-level coverage log2-ratios. It is straightforward to replace the default with other methods and the segmentationCBS function can serve as an example.

The next section describes how to replace the default segmentation. For the probably more uncommon case that only the coverage normalization is performed by third-party tools, see Section 10.1.2.

10.1.1 Custom segmentation

It is possible to provide already segmented data, which is especially recommended when third-party segmentation tools are not written in R. Otherwise it is usually however better to customize the default segmentation function, since the algorithm then has access to the raw log2-ratio distribution⁵. The expected file format for already segmented copy number data parsed by readSegmentationFile is⁶:

ID chrom loc.start loc.end num.mark seg.mean Sample1 1 61723 5773942 2681 0.125406444072723 Sample1 1 5774674 5785170 10 -0.756511807441712

Since its likelihood model is exon-based, *PureCN* currently still requires an interval file to generate simulated target-level log2-ratios from a segmentation file. For simplicity, this interval file is expected either via the tumor.coverage.file or via the interval.file argument (see Figure 7). Note that *PureCN* will re-segment the simulated log2-ratios using the default segmentationCBS function, in particular to identify regions of copy-number neutral LOH and to cluster segments with similar allelic imbalance and log2-ratio. The provided interval file should therefore cover all significant copy number alterations⁷. Please check that the log2-ratios are similar to the ones obtained by the default *PureCN* segmentation and normalization.

```
retSegmented <- runAbsoluteCN(seg.file = seg.file,
    interval.file = interval.file, vcf.file = vcf.file,
    max.candidate.solutions = 1, genome = "hg19",
    test.purity = seq(0.3,0.7,by = 0.05), verbose = FALSE,
```

⁵If the third-party tool provides target-level log2-ratios, then these can be provided via the log.ratio argument in addition to seg.file though. See also Section 10.1.2.

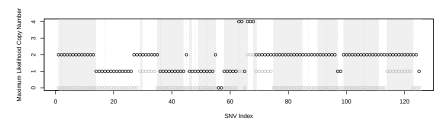
⁶This segmentation file can contain multiple samples, in which case the provided sampleid must match a sample in the column ID

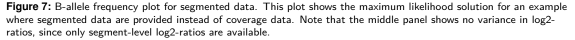
⁷If this behaviour is not wanted, because maybe the custom function already identifies CNNLOH reliably, segmentationCBS can be replaced with a minimal version. plotAbs(retSegmented, 1, type = "BAF")

plot.cnv = FALSE)
WARN [2024-05-01 18:56:22] Allosome coverage missing, cannot determine sex.
WARN [2024-05-01 18:56:22] Allosome coverage missing, cannot determine sex.

The max.candidate.solutions and test.purity arguments are set to non-default values to reduce the runtime of this vignette.

Purity: 0.65 Tumor ploidy: 1.74 SNV log-likelihood: -5.18 GoF: 93.4% Mean coverage: 109;82 10 11 12 22 689 14 15 17 18 0.8 B-Allele Frequency 0.6 0.4 0.2 60 100 120 SNV Index SCNA-fit log-likelihood: -274.5 1.0 Copy Number log-ratio 0.0 -1.0 0 2.0 20 40 60 100 120 80 SNV Index





10.1.2 Custom normalization

If third-party tools such as GATK4 are used to calculate target-level copy number log2-ratios, and PureCN should be used for segmentation and purity/ploidy inference only, it is possible to provide these log2-ratios:

```
retLogRatio <- runAbsoluteCN(log.ratio = log.ratio,
    interval.file = interval.file, vcf.file = vcf.file,
    max.candidate.solutions = 1, genome = "hg19",
    test.purity = seq(0.3, 0.7, by = 0.05), verbose = FALSE,
    normalDB = normalDB, plot.cnv = FALSE)
## WARN [2024-05-01 18:56:41] Allosome coverage missing, cannot determine sex.
## WARN [2024-05-01 18:56:41] Allosome coverage missing, cannot determine sex.
```

Again, the max.candidate.solutions and test.purity arguments are set to non-default values to reduce the runtime of this vignette. Note that this example uses a pool of normals to filter low quality targets. Interval coordinates are again expected in either a interval.file or a tumor.coverage.file.

The **readLogRatioFile** function provides support for parsing *GATK3*-style and *GATK4 DenoiseReadCounts* formats:

```
# GATK3-style
Target log_ratio
1:3682750-3683489 0.109473
1:3690585-3691264 0.126205
# GATK4
@HD VN:1.6
@SQ SN:1 LN:248956422
@SQ SN:2 LN:242193529
...
CONTIG START END LOG2_COPY_RATIO
1 3682750 3683489 0.109473
1 3690585 3691264 0.126205
```

It is highly recommended to compare the log2-ratios obtained by *PureCN* and the third-party tool, since some pipelines automatically adjust log2-ratios for a default purity value.

10.2 COSMIC annotation

If a matched normal is not available, it is also helpful to provide runAbsoluteCN the COSMIC database [13] via cosmic.vcf.file (or via a Cosmic.CNT INFO field in the VCF). While this has limited effect on purity and ploidy estimation due the sparsity of hotspot mutations, it often helps in the manual curation to compare how well high confidence germline (based on dbSNP membership or POPAF) vs. somatic (COSMIC) variants fit a particular purity/ploidy combination.

For variant classification (Section 6.2.1), providing COSMIC annotation also avoids that hotspot mutations in germline databases get assigned a very low prior probability of being somatic.

10.3 ExAC and gnomAD annotation

PureCN is not automatically annotating input VCFs with data from common germline databases such as ExAC. See Section 2.1 for ways to tell *PureCN* where to find either a summary binary flag (i.e. likely germline yes/no) or population allele frequencies.

10.4 Mutation burden

The predictSomatic function described in Section 6.2.1 can be used to efficiently remove private germline mutations. This in turn allows the calculation of mutation burden for unmatched tumor samples. A wrapper function for this specific task is included as callMuta tionBurden:

```
callableBed <- import(system.file("extdata", "example_callable.bed.gz",</pre>
    package = "PureCN"))
callMutationBurden(ret, callable = callableBed)
##
     somatic.ontarget somatic.all private.germline.ontarget
## 1
                    0
                                 0
                                                            0
##
     private.germline.all callable.bases.ontarget callable.bases.flanking
## 1
                                           1493367
                                                                    2194877
                        0
##
    callable.bases.all somatic.rate.ontarget somatic.rate.ontarget.95.lower
## 1
                3063762
                                             0
                                                                             0
##
     somatic.rate.ontarget.95.upper private.germline.rate.ontarget
## 1
                           2.470173
                                                                   0
##
     private.germline.rate.ontarget.95.lower
## 1
                                            0
##
     private.germline.rate.ontarget.95.upper
                                     2.470173
## 1
```

The callableBed file should be a file parsable by *rtracklayer*. This file can specify genomic regions that are callable, for example as obtained by *GATK3 CallableLoci*. This is optional, but if provided can be used to accurately calculate mutation rates per megabase. Variants outside the callable regions are not counted. Private germline rates should be fairly constant across samples; outliers here should be manually inspected.

The output columns are explained in Table 6.

10.5 Chromosomal Instability

Chromosomal Instability (CIN) is usually defined as the fraction of the genome that is altered. The callCIN function can be used to estimate this fraction.

Parameters define regions that are altered. First, allele.specific defines whether only total vs. both minor and major copy number define a state. Copy number neutral LOH would count as altered only when this parameter is set to TRUE. Second, reference.state defines the unaltered copy number state. This can be either normal for 2/1, or dominant for the most common state. While technically potentially wrong, the latter is robust to errors in ploidy and is thus recommended without careful manual curation. Similarly, setting allele.specific to FALSE makes this metric more robust to purity and ploidy errors, but usually to a much lesser extend.

It is recommended to test for a relationship of tumor purity and CIN and if necessary exclude low purity samples with uncertain CIN.

```
# All 4 possible ways to calculate fraction of genome altered
cin <- data.frame(
    cin = callCIN(ret, allele.specific = FALSE, reference.state = "normal"),
    cin.allele.specific = callCIN(ret, reference.state = "normal"),
```

Column name	Description
somatic.ontarget	Number of mutations in target regions
somatic.all	Total number of mutations. Depending on input VCF and runAbsoluteCN argu- ments, this might include calls in flanking regions and off-targets reads.
private.germline.ontarget	Number of private germline SNPs in tar- gets. For matched tumor/normal sam- ples, this is the number of variants an- notated as somatic, but classified as germline.
private.germline.all	Total number of private germline SNPs
callable.bases.ontarget	Number of callable on-target bases
callable.bases.flanking	Number of callable on-target and flanking bases
callable.bases.all	Total number of callable bases. With default parameters includes off-target re- gions that were ignored by runAbso luteCN.
<pre>somatic.rate.ontarget</pre>	Somatic mutations per megabase in tar- get regions
<pre>somatic.rate.ontarget.95.lower</pre>	Lower 95% of confidence interval
<pre>somatic.rate.ontarget.95.upper</pre>	Upper 95% of confidence interval
private.germline.rate.ontarget	Private germline mutations per megabase in target regions
<pre>private.germline.rate.ontarget.95.lower</pre>	Lower 95% of confidence interval
private.germline.rate.ontarget.95.upper	Upper 95% of confidence interval

 Table 6: callMutationBurden output columns.

```
cin.ploidy.robust = callCIN(ret, allele.specific = FALSE),
cin.allele.specific.ploidy.robust = callCIN(ret)
```

)

10.6 Detect cross-sample contamination

It is important to correctly handle heterozygous common SNPs that do not have an expected allelic fraction of 0.5 in normal samples. These can be SNPs in poor quality regions (as already described, see Section 4.2.1), but also SNPs from cross-sample contaminated DNA. Without matched normals, detection of those problematic SNPs is not trivial.

For cross-sample contamination, PureCN by default always tests for a 1% contamination and assigns common SNPs to a contamination state when allelic fractions are either close to 0 or close to 1 and when this cannot be explained by CNAs. The main purpose of these states is to provide a bin for common SNPs that for artifactual reasons do not fit any other state.

This tool applies a simple heuristic to flag samples for cross-contamination: Given the coverage and putative contamination rate based on allelic fractions of potentially contaminated SNPs, how many SNPs do we expect to detect based on our power to detect variants at that contamination rate? If the expected number is much higher than observed, then significant contamination is unlikely; observed SNPs close to 0 or 1 are more likely artifacts or the contamination rate is much lower than the minimum tested. Otherwise *PureCN* will perform a post-optimization in which contamination rate is optimized in additional variant fitting steps.

Cross-sample contamination can also result in increased observed heterozygosity on chrX for males, which in turn often results in a *PureCN* warning that sex inferred from coverage and VCF are in conflict.

By default, cross-contamination is tested in the range from 1 to 7.5%. Catastrophic failures with higher contamination might not get flagged.

10.7 Power to detect somatic mutations

As final quality control step, we can test if coverage and tumor purity are sufficent to detect mono-clonal or even sub-clonal somatic mutations. We strictly follow the power calculation by Carter et al. [2].

The following Figure 8 shows the power to detect mono-clonal somatic mutations as a function of tumor purity and sequencing coverage (reproduced from [2]):

```
purity <- c(0.1,0.15,0.2,0.25,0.4,0.6,1)
coverage <- seq(5,35,1)
power <- lapply(purity, function(p) sapply(coverage, function(cv)
        calculatePowerDetectSomatic(coverage = cv, purity = p, ploidy = 2,
        verbose = FALSE)$power))
# Figure S7b in Carter et al.
plot(coverage, power[[1]], col = 1, xlab = "Sequence coverage",
        ylab = "Detection power", ylim = c(0, 1), type = "l")
for (i in 2:length(power)) lines(coverage, power[[i]], col = i)
abline(h = 0.8, lty = 2, col = "grey")
legend("bottomright", legend = paste("Purity", purity),
        fill = seq_along(purity))</pre>
```

Figure 9 then shows the same plot for sub-clonal mutations present in 20% of all tumor cells:

```
coverage <- seq(5,350,1)
power <- lapply(purity, function(p) sapply(coverage, function(cv)
        calculatePowerDetectSomatic(coverage = cv, purity = p, ploidy = 2,
        cell.fraction = 0.2, verbose = FALSE)$power))
plot(coverage, power[[1]], col = 1, xlab = "Sequence coverage",
        ylab="Detection power", ylim = c(0, 1), type = "l")
for (i in 2:length(power)) lines(coverage, power[[i]], col = i)
abline(h = 0.8, lty = 2, col = "grey")
legend("bottomright", legend = paste("Purity", purity),
        fill = seq_along(purity))</pre>
```

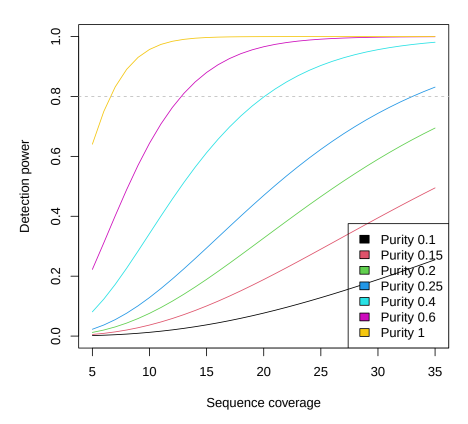


Figure 8: Power to detect mono-clonal somatic mutations. Reproduced from [2].

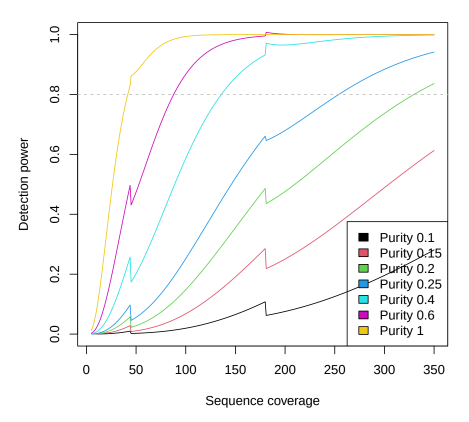


Figure 9: Power to detect sub-clonal somatic mutations present in 20% of all tumor cells. Reproduced from [2].

11 Limitations

PureCN currently assumes a completely diploid normal genome. For human samples, it tries to detect sex by calculating the coverage ratio of chromosomes X and Y and will then remove sex chromosomes in male samples⁸. For non-human samples, the user needs to manually remove all non-diploid chromosomes from the coverage data and specify sex="diploid" in the *PureCN* call.

While *PureCN* supports and models sub-clonal somatic copy number alterations, it currently assumes that the majority of alterations are mono-clonal. For most clinical samples, this is reasonable, but very heterogeneous samples are likely not possible to call without manual curation. Poly-genomic tumors are often called as high ploidy or low purity. The former usually happens when sub-clonal losses are called as 2 copies and mono-clonal losses correctly as 1 copy. The latter when sub-clonal losses are called mono-clonal, which only happens when there are far more sub-clonal than mono-clonal losses. Please note however that unless purities are very high, algorithms that model poly-genomic tumors do not necessarily have a higher call rate, since they tend to overfit noisy samples or similarly confuse true high-ploidy with poly-genomic tumors. Due to the lack of signal, manual curation is also recommended in low purity samples or very quiet genomes.

12 Support

If you encounter bugs or have problems running *PureCN*, please post them at

- https://support.bioconductor.org/p/new/post/?tag_val=PureCN or
- https://github.com/lima1/PureCN/issues.

If *PureCN* throws user errors, then there is likely a problem with the input files. If the error message is not self-explanatory, feel free to seek help at the support site.

In your report, please add the outputs of the runAbsoluteCN call (with verbose=TRUE, or the *.log file in PureCN.R) and sessionInfo(). Please also check that your problem is not already covered in the following sections.

For general feedback such as suggestions for improvements, feature requests, complaints, etc. please do not hesitate to send us an email.

12.1 Checklist

- Used the correct interval files provided by the manufacturer of the capture kit and the genome version of the interval file matches the reference. Ideally used the baits file, not the targets file (in Agilent data, the baits files are called "covered" and the targets are "regions").
- For hybrid capture data, included off-target reads in the coverage calculation
- BAM files were generated following established best practices and tools finished successfully.
- Checked standard QC metrics such as AT/GC dropout and duplication rates.
- Tumor and normal data were obtained using the same capture kit and pipeline.
- Coverage data of tumor and normal were GC-normalized.

⁸Loss of Y chromosome (LOY) can result in wrong female calls, especially in high purity samples or if LOY is in both tumor and contaminating normal cells. The software throws a warning in this case when germline SNPs are provided.

- The VCF file contains germline variants (i.e. not only somatic calls).
- Maximized the number of high coverage heterozygous SNPs, for example by running *MuTect* with a 50-75bp interval padding (Section 9). The runAbsoluteCN output lists the percentage of targets with variants and this should be around 10-15%. Ultradeep sequencing data can provide good SNP allelic fractions in the 100-200bp flanking regions.
- If a pool of normal samples is available, followed the steps in Section 4.2.
- Read the output of runAbsoluteCN with verbose = TRUE, fixed all warnings.
- If third-party segmentation tools are used, checked that normalized log2-ratios are not biased, i.e. very similar compared to *PureCN* log2-ratios (some pipelines already adjust for a default normal contamination).
- Followed the best practices described in the Quick vignette

12.2 FAQ

If the ploidy is frequently too high, please check:

- Does the log2-ratio histogram (Figure 3) look noisy? If yes, then
 - Is the coverage sufficient? Tumor coverages below 80X can be difficult, especially in low purity samples. Normal coverages below 50X might result in high variance of log2-ratios. See Section 4.1 for finding a good normal sample for log2-ratio calculation.
 - Is the coverage data of both tumor and normal GC-normalized? If not, see correct CoverageBias. It can be safe to skip the normalization, with large panels of normals potentially even slighly beneficial, but make sure that GC-normalization was either skipped or performed in both tumor sample and normal coverage database.
 - Is the quality of both tumor and normal sufficient? A high AT or GC-dropout might result in high variance of log2-ratios. Challenging FFPE samples also might need parameter tuning of the segmentation function. See segmentationCBS. A high expected tumor purity allows more aggressive segmentation parameters, such as prune.hclust.h=0.2 or higher.
 - Was the correct target interval file used (genome version and capture kit, see Section 2.4)? If unsure, ask the help desk of your sequencing center.
 - Were the normal samples run with the same assay and pipeline?
 - Did you provide runAbsoluteCN all the recommended files as described in Section 5?
 - For whole-genome data, you will get better results using a specialized third-party segmentation method as described in section 10.1, since our default is optimized for targeted sequencing. In general, you should probably start with tools optimized for WGS data, such as Battenberg [14], ABSOLUTE [2], ACEseq [7], or TitanCNA [8]. We are planning to incorporate proper support for WGS once high coverage diagnostic WGS becomes more common.
- Otherwise, if log2-ratio peaks are clean as in Figure 3:

- Was MuTect run without a matched normal? If yes, then make sure to provide either a pool of normal VCF or a SNP blacklist (if no pool of normal samples is available) as described in Sections 4.2 and 4.3.
- A high fraction of sub-clonal copy-number alterations might also result in a low ranking of correct low ploidy solutions (see Section 11).

If the ploidy is frequently too low:

- PureCN with default parameters is conservative in calling genome duplications.
- This should only affect low purity samples (< 35%), since in higher purity samples the duplication signal is usually strong enough to reliably detect it.
- In whole-exome data, it is usually safe to decrease the max.homozygous.loss default, since such large losses are rare.

Will PureCN work with my data?

- *PureCN* was designed for medium-sized (>2-3Mb) targeted panels. The more data, the better, best results are typically achieved in whole-exome data.
- The number of heterozygous SNPs is also important (>1000 per sample). Copy number baits enriched in SNPs are therefore very helpful (see Section 9).
- Some users got acceptable results with small (<1Mb) panels. Try to find a perfect off-target bin width (average.off.target.width in preprocessIntervals) and maximize the number of heterozygous SNPs by including as much padding as possible. Keep in mind that without tiling baits, you will only have a poor resolution to detect LOH.
- Coverages below 80X are difficult unless purities are high and coverages are even.
- PureCN also needs process-matched normal samples, again, the more the better.
- Samples with tumor purities below 15-20% usually cannot be analyzed with this algorithm and *PureCN* might return very wrong purity estimates. In high coverage samples with low duplication rates, this limit can be close to 10%.
- Whole-genome data is not officially supported and specialized tools will likely provide better results. Third-party segmentation tools designed for this data type would be again required.
- Amplicon sequencing data is also not officially supported. If the assay contains tiling
 probes (at least with 1Mb spacing) and uses a barcode protocol that reduces PCR
 bias of measured allelic fractions, then this method might produce acceptable results.
 Setting the model argument of runAbsoluteCN to betabin is recommended. Specialized
 segmentation tools might be again better than our default.
- This tool was originally designed for single sample pipelines. It becomes more common to have multiple biopsies per patient and sharing information across biopsies might be beneficial. Other tools worth checking out are lumosVar 2.0 [15] and SuperFreq [16].

If you have trouble generating input data PureCN accepts, please check:

- For problems related to generating valid coverage data, either consult the GATK manual for the DepthOfCoverage or CollectFragmentCounts tools or Section 2.3 for the equivalent function in PureCN. If you use DepthOfCoverage and off-target intervals as generated by IntervalFile.R (See Quick vignette), make sure to run it with parameters
 -omitDepthOutputAtEachBase and -interval_merging OVERLAPPING_ONLY.
- Currently only VCF files generated by *MuTect 1* are officially supported and well tested. A minimal example *MuTect* call would be:

```
$JAVA7 -Xmx6g -jar $MUTECT \
--analysis_type MuTect -R $REFERENCE \
--dbsnp $DBSNP_VCF \
--cosmic $COSMIC_VCF \
-I:normal $BAM_NORMAL \
-I:tumor $BAM_TUMOR \
-o $0UT/${ID}_bwa_mutect_stats.txt \
-vcf $0UT/${ID}_bwa_mutect.vcf
```

The normal needs to be matched; without matched normal, only provide the tumor BAM file (do NOT provide a process-matched normal here). The default output file is the stats or call-stats file; this can be provided in addition to the required VCF file via args.filterVcf in runAbsoluteCN. If provided, it may help *PureCN* filter artifacts. This requires *MuTect* in version 1.1.7. This version is currently available at https://software.broadinstitute.org/gatk/download/mutect and requires Java 1.7 (it does not work with Java 1.8).

Note that this *MuTect* VCF will contain variants in off-target reads. By default, *PureCN* will remove variants outside the provided targets and their 50bp flanking regions. We highly recommend finding good values for each assay. A good cutoff will maximize the number of heterozygous SNPs and keep only an acceptable number of lower quality calls. This cutoff is set via interval.padding in args.filterVcf. See Section 9.

- Support for GATK4 and Mutect2 is in beta. Version 4.1.7.0 and higher as well as specifying the -genotype-germline-sites flag in Mutect2 is required. Please follow the Broad Best Practices as closely as possible as we only test those workflows.
- For VCFs generated by other callers, the required dbSNP annotation can be added for example with *bcftools*:

```
bcftools annotate --annotation $DBSNP_VCF \
    --columns ID --output $OUT/${ID}_bwa_dbsnp.vcf.gz --output-type z \
    $OUT/${ID}_bwa.vcf.gz
```

- To generate a mappability file with the GEM library:
 - Calculate mappability, set kmer size to length of mapped reads.

```
REFERENCE="hg38.fa"
PREF=`basename $REFERENCE .fa`
THREADS=4
KMER=100
gem-indexer -T ${THREADS} -c dna -i ${REFERENCE} -o ${PREF}_index
gem-mappability -T ${THREADS} -I ${PREF}_index.gem -l ${KMER} \
        -0 ${PREF}_${KMER} -m 2 -e 2
gem-2-wig -I ${PREF}_index.gem -i ${PREF}_${KMER}.mappability \
```

```
-o ${PREF}_${KMER}
```

• Convert to bigWig format, for example using the UCSC wigToBigWig tool:

```
cut -f1,2 ${REFERENCE}.fai > ${PREF}.sizes
wigToBigWig ${PREF}_${KMER}.wig ${PREF}.sizes ${PREF}_${KMER}.bw
```

- To generate the normal panel VCF (normal.panel.vcf.file) with GATK3:
 - Run *MuTect* on the normal with -I:tumor \$BAM_NORMAL and optionally with the -artifact_detection_mode flag.
 - Remove the empty none sample from the VCF:

```
$JAVA -Xmx6g -jar $GATK3 \
--analysis_type SelectVariants -R $REFERENCE \
--exclude_sample_expressions none \
-V $OUT/${ID}_bwa_mutect_artifact_detection_mode.vcf \
-o $OUT/${ID}_bwa_mutect_artifact_detection_mode_no_none.vcf
```

Merge the VCFs:

```
CMD="java -Xmx12g -jar $GATK3 -T CombineVariants --minimumN 3 -R $REFERENCE"
for VCF in $OUT/*bwa_mutect_artifact_detection_mode_no_none.vcf;
do
        CMD="$CMD --variant $VCF"
done
CMD="$CMD -o $OUT/normals_merged_min3.vcf"
echo $CMD > $OUT/normals_min3.sh
bgzip $OUT/normals_merged_min3.vcf
tabix $OUT/normals_merged_min3.vcf
```

Questions related to the output.

- Where is the major and minor segment copy number? The minor copy number is ML.M.SEGMENT in predictSomatic (Table 1) and M in callAlterations (Table 2) and callLOH (Table 4). The major copy number is simply the total copy number (column C or ML.C in predictSomatic) minus the minor copy number.
- Why are there two different amplification output files? See Section 6.2.2. Genelevel p-values are not very useful for calling high-level amplifications in tissue samples with significant amount of tumor cells because even regions of single gains will have significant read pile-ups. These p-values can be however useful in low purity samples commonly observed in cfDNA samples. There is no corresponding file for homozygous deletions since detecting those is usually impossible in very low purity samples.

Questions related to pool of normals. As described in detail in Sections 4.1 and 4.2, a pool of normal samples is used in *PureCN* for coverage normalization (to adjust for target-specific capture biases) and for artifact filtering. A few recommendations:

 Matched normals are obviously perfect for identifying most alignment artifacts and mapping biases. While not critical, we still recommend generating a normal.panel.vcf.file for *MuTect* and calculateMappingBiasVcf using the available normals.

- Without matched normals, we need process-matched normals for coverage normalization. We recommend more than 5 from 2-3 different library preparation and sequencing batches. The more, the better.
- If you are able to add new normal samples over time capturing potential subtle changes in the lab, do so!
- These normals used for coverage normalization should be ideally sequenced to at least half the coverage as the tumor samples. This is completely different from matched normals for germline calling where 30-40X provide sufficient power to detect heterozygosity.
- For artifact removal, the more normals available, the more rare artifacts are removed. We recommend at least 10, 50 or more are ideal. The more artifacts are removed, the less likely *PureCN* output requires manual curation (Section 4.2).
- For position-specific mapping bias correction, the more normals are available, the more rare SNPs will have reliable mapping bias estimates. This requires again at least about 10 normals to be useful, 100 or more are ideal.
- With smaller pool of normals, we additionally recommend filtering SNPs from low quality regions (Section 4.3).
- It is worth trying the beta-binomial function instead of the default in the model argument
 of runAbsoluteCN. This will incorporate uncertainty of observed variant allelic fractions
 in the variant fitting step. The more normals, the more SNPs will have reliable estimates
 of allelic fraction overdispersion.
- Do I really need a pool of normals? I only have tumor samples. Unfortunately, yes. If you used a commercial capture kit, you might be able to obtain control samples from the vendor or the public domain. This is not optimal, but usually better than nothing.
- It is safe to include multiple normals from the same individual. Fewer common germline CNVs in createNormalDatabase and fewer SNPs in calculateMappingBiasVcf will be detected. But especially when the controls were sequenced in multiple batches, these replicates will still provide useful information for coverage normalization.

Questions related to manual curation. *PureCN*, like most other related tools, essentially finds the most simple explanation of the data. There are three major problems with this approach:

- First, hybrid capture data can be noisy and the algorithm must distinguish signal from noise; if the algorithm mistakes noise for signal, then this often results in wrong high ploidy calls (see Sections 4.2 and 4.3). If all steps in this vignette were followed, then *PureCN* should ignore common artifacts. Noisy samples thus often have outlier ploidy values and are often automatically flagged by *PureCN*. The correct solution is in most of these cases ranked second or third.
- The second problem is that signal can be sparse, i.e. when the tumor purity is very low or when there are only few somatic events. Manual curation is often easy in the latter case. For example when small losses are called as homozygous, but corresponding germline allele-frequencies are unbalanced (a complete loss would result in balanced germline allele frequencies, since only normal DNA is left). Future versions might improve calling in these cases by underweighting uninformative genomic regions.

• The third problem is that tumor evolution is fast and complex and very difficult to incorporate into general likelihood models. Sometimes multiple solutions explain the data equally well, but one solution is then often clearly more consistent with known biology, for example LOH in tumor suppressor genes such as *TP53*. A basic understanding of both the algorithm and the tumor biology of the particular cancer type are thus important for curation. Fortunately, in most cancer types, such ambiguity is rather rare. See also Section 11.

Questions related to matched normals.

- **Coverage normalization:** Even when matched normals are available, we recommend building a normal database for coverage normalization. This usually produces cleaner coverage profiles than the matched normal ([9]).
- **VCFs:** When matched normals are available, simply provide this information to the variant caller and make sure that germline SNPs are not filtered out. *PureCN* should automatically find the matched information.

Questions related to off-target reads. Hybrid capture assays usually provide a significant amount of reads mapping outside the captured regions. This means these assays provide shallow whole genome sequencing data for free. *PureCN* makes it straightforward to use that information, but it might not always make sense to do so:

- Does the assay contain a fairly large number of big gaps (>1Mb) between neighboring baits? For whole exome data or targeted panels with copy number tiling probes, the answer can be no.
- Off-target reads are usually less affected by library-specific biases (Section 3) than ontarget reads. In noisy data, off-target reads can thus provide relatively clean additional information.
- Very efficient assays with low off-target mapping might not provide a large number of reads in useful window sizes (usually below 0.2Mb), especially when total sequencing coverages are rather low. This can result in off-target data that are significantly noisier than on-target data.
- It is recommended to check the runAbsoluteCN output for the standard deviations of log2 copy number ratios for both on- and off-target reads. If the off-target noise is consistently larger than the on-target noise, you can try increasing the off-target bin width - and vice versa make it smaller when the on-target noise is larger.

Questions related to GC normalization.

- Explicit GC-normalization is useful whenever a significant difference in tumor and normal controls is suspected. One example would be FFPE tumors normalized with blood controls. Otherwise the normalization using the panel of normals should automatically remove any GC bias.
- Smaller panels of normals might also benefit from explicit GC-normalization.
- Otherwise it is safe skip the GC-normalization. In the command line tools, set -skipgc-norm in Coverage.R.

If all or most of the samples are flagged as:

- **Noisy segmentation:** The default of 300 for max.segments is calibrated for high quality and high coverage whole-exome data. For whole-genome data or lower coverage data, this value needs to be re-calibrated. In case the copy number data looks indeed noisy, please see the first FAQ. Please be aware that *PureCN* will apply more aggressive segmentation parameters when the number of segments exceeds this cutoff. If the high segment count is real, this might confound downstream analyses.
- **High AT/GC dropout:** If the data are GC-normalized, then there might be issues with either the target intervals or the provided GC content. Please double check that all files are correct and that all the coverage files are GC-normalized (Section 3).
- Sex mismatch of coverage and VCF: If the panel contains baits for chromosome Y, then the interval file was probably generated without mappability file (Section 2.2). Similarly when third-party tools were used for coverage normalization and segmentation, this usually means probes on chromosome Y were not filtered for mappability. Cross-sample contamination (Section 10.6) can also cause sex mismatches.

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Session Info

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- TZcode source: system (glibc)
- Running under: Ubuntu 22.04.4 LTS
- Matrix products: default
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- LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, stats4, utils
- Other packages: Biobase 2.65.0, BiocGenerics 0.51.0, BiocStyle 2.33.0, Biostrings 2.73.0, DNAcopy 1.79.0, GenomeInfoDb 1.41.0, GenomicRanges 1.57.0, IRanges 2.39.0, MatrixGenerics 1.17.0, PureCN 2.11.0, Rsamtools 2.21.0, S4Vectors 0.43.0, SummarizedExperiment 1.35.0, VariantAnnotation 1.51.0, XVector 0.45.0, matrixStats 1.3.0

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