ACEseqDocs Documentation

Release 1.2.8

Kortine Kleinheinz

Contents

	Se Control of the Con	1
Need h	nelp?	3
3.1	The Standard Way	5 5
Quicks	Start	11
5.1 1	Hardware	13 13 13
6.1 I 6.2 I 6.3 I	Run Without Control	15 15 15 16
6.4	Run with/without 5 v breakpoint incorporation	16
		16 17
Input 1	Parameters	
Input I	Parameters Evaluation	17
Input I	Parameters Evaluation ty check	17 21
Input Inp	Parameters Evaluation Ey check Output	17 21 27
	Install 3.1 3.2 Quick Requi 5.1 5.2 Altern 6.1 6.2	3.2 Prepackaged files (ACEseq 1.2.10 only) QuickStart Requirements 5.1 Hardware

12.8	Copy Number Estimation	38
12.9	Purity and ploidy estimation	39

			-4
CHA	PT	FR	

License

The license of the ACEseq code is MIT https://github.com/eilslabs/ACEseqWorkflow/blob/github/package_LICENSES.md licenses of packages used by the workflow.

2 Chapter 1. License

CHAP	TER	_

Need help?

In case of question pleae contact Kortine Kleinheinz (k.kleinheinz@dkfz-heidelberg.de)

Installation & Run instructions

To run the ACEseq-workflow multiple components are needed:

- · ACEseq workflow plugin
- The Roddy workflow management framework
- · Software stack
- Reference data
- COWorkflowsBasePlugin

The *The Standard Way* to install the workflow is described below and involves the installation of each of these components. For the older 1.2.10 release we currently also provide prepackaged files and a Docker container. See *Prepackaged files (ACEseq 1.2.10 only)* below for instructions.

3.1 The Standard Way

The standard way to install the workflow is the manual installation of all components.

- 1. Download the COWorkflowBasePlugin zip-archive from Github-Releases. The version to download can be found in the ACEseq buildinfo.txt.
- 2. Download the ACEseq zip-archive from Github-Releases. The archive already contains a Jar-archive with the compiled Java/Groovy code (JAR-file) for the given Roddy API version. No compilation of the plugin is therefore required.
- 3. The file ACEseq buildinfo.txt in also shows you the Roddy API version that you need for the chosen ACEseq workflow version.
- 4. Install the required Roddy version. Please see the Roddy repository for installation instructions for Roddy.
- 5. Install the software stack (see Software Stack (Conda) below) via Conda
- 6. Install the reference files (see *Reference files* below) via the preparation script.

3.1.1 Software Stack (Conda)

The workflow contains a description of a Conda environment. A number of Conda packages from BioConda are required. You should set up the Conda environment at a centralized position available from all compute hosts.

First install the BioConda channels:

conda config --add channels r

conda config --add channels defaults

conda config --add channels conda-forge

conda config --add channels bioconda

Then install the environment

The name of the Conda environment is arbitrary but needs to be consistent with the *condaEnvironmentName* variable. You can set the *condaEnvironmentName* variable in any of the loaded configuration files (see Roddy documentation) or even directly in your Roddy call via *-cvalues="condaEnvironmentName:\$value"*.

If you do not want to use Conda, you can get a complete list of all packages and package versions Conda would install from the \$PATH_TO_PLUGIN_DIRECTORY/resources/analysisTools/copyNumberEstimationWorkflow/environments/conda.yml.

3.1.2 Reference files

The workflow uses various files as reference files, such as a reference genome or annotation files. Depending on the contents of these files also the outcome of your analysis may change. We provide installation scripts in the *installation/* directory (currently only in the *github* branch of the repository). To download and prepare the reference files please check out the ACEseq repository and do

bash \$PATH_TO_PLUGIN_DIRECTORY/installation/downloadRefrences \$targetDirectory

with \$targetDirectory\$ being the directory into which you want to install the files. The variable baseDirectoryReference in your configurations needs to be set to the \$targetDirectory\$ path.

Note that the current plugin version is tuned to be run on the hg19 human assembly, but a liftover of all files should probably enable a run on GRch38.

3.2 Prepackaged files (ACEseq 1.2.10 only)

On http://bfg-nfs3.ipmb.uni-heidelberg.de you can find archives for the 1.2.10 plugin version. The prepackaged zip files contains a full Roddy / Plugin setup and include different scripts to install all necessary software and download the required reference files. Currently, we do not intent to update these prepackaged installation files or the Docker version. Note that the Roddy version packaged not capable of submitting to LSF.

Please see the standard way to install recent workflow versions.

3.2.1 Stand-alone Roddy for Execution on HTC Cluster

To run the Roddy-based version of ACEseq please download the pre-packed zip file from http://bfg-nfs3.ipmb. uni-heidelberg.de. Three steps are required to ensure running of ACEseq.

- 1. Run the "prepareRoddyInstallation.sh" script.
- 2. Download all reference files as specified in the section "Reference files" (below).
- 3. Set up the Conda environment or install the necessary software as specified in the section "Software" (below).

Before running ACEseq a few parameters need to be adjusted in the configuration files. The output directory is specified in \$PATH_TO_ACEseq_RODDY_VERSION/configurations/projectsACEseqTest.xml. Here the variables "baseDirectoryReference", "inputBaseDirectory", "outputBaseDirectory", "outputAnalysisBaseDirectory" need to be set. If no SVs should be included the following configuration values (cvalues) should be included:

```
<cvalue name='runWithSv' value='true' type="boolean"/>
<cvalue name='SV' value='yes' type="boolean"/>
```

Otherwise "svOutputDirectory" and the SV bedpe filename in the filenames section need to be set.

Technical specifications are set in the file \$PATH_TO_ACEseq_RODDY_VERSION/configurations/applicationProperties.ini. The path to the project.xml and the path to the plugins (\$PATH_TO_ACEseq_RODDY_VERSION/Roddy/dist/plugins/) need to be set under configurationDirectories and pluginDirectories. Finally the job manager and execution host need to be set.

Please have a look at the following default applicationProperties.ini file:

```
[COMMON]
useRoddyVersion=current
                                             # Use the most current version for tests
[DIRECTORIES]
configurationDirectories=[FOLDER_WITH_CONFIGURATION_FILES]
pluginDirectories=[FOLDER_WITH_PLUGINS]
[COMMANDS]
jobManagerClass=de.dkfz.roddy.execution.jobs.direct.synchronousexecution.
→ DirectSynchronous Execution Job Manager
#jobManagerClass=de.dkfz.roddy.execution.jobs.cluster.pbs.PBSJobManager
#jobManagerClass=de.dkfz.roddy.execution.jobs.cluster.sge.SGEJobManager
#jobManagerClass=de.dkfz.roddy.execution.jobs.cluster.slurm.SlurmJobManager
#jobManagerClass=de.dkfz.roddy.execution.jobs.cluster.lsf.rest.LSFRestJobManager
commandFactoryUpdateInterval=300
commandLogTruncate=80
                                             # Truncate logged commands to this length.
\hookrightarrow If <= 0, then no truncation.
[COMMANDLINE]
CLI.executionServiceUser=USERNAME
```

```
CLI.executionServiceClass=de.dkfz.roddy.execution.io.LocalExecutionService
#CLI.executionServiceClass=de.dkfz.roddy.execution.io.SSHExecutionService
CLI.executionServiceHost=[YOURHOST]
CLI.executionServiceAuth=keyfile
#CLI.executionServiceAuth=password
CLI.executionServicePasswd=
CLI.executionServiceStorePassword=false
CLI.executionServiceUseCompression=false
CLI.fileSystemInfoProviderClass=de.dkfz.roddy.execution.io.fs.FileSystemInfoProvider
```

To execute ACEseq run

```
sh $PATH_TO_ACEseq_RODDY_VERSION//Roddy/roddy.sh rerun ACEseq@copyNumberEstimation

→$pid \

--useconfig=$PATH_TO_ACEseq_RODDY_VERSION/configuration/applicationProperties.ini \

--cvalues="bamfile_list:$pathToControlBamFile;$pathToTumorBamFile,sample_list:control;

→tumor,possibleControlSampleNamePrefixes:control,

→possibleTumorSampleNamePrefixes:tumor"
```

More information on Roddy can be found here.

3.2.2 Docker version

- 1. Download all reference files as specified in the section below.
- 2. Download the Base and ACEseq Docker images from the website: http://bfg-nfs3.ipmb.uni-heidelberg.de
- 3. Import both files with (names might differ based on supplied version):

```
docker load < BaseDockerContainer.tar.gz

docker load < ACEseqDockerContainer.tar.gz
```

4. Download the control files archive and extract them. The directory contains the file "roddy.sh". Please call this script with: bash roddy.sh. You will see:

```
#!/bin/bash
# 1: Run mode, which might be "run" or "testrun"
# 2: Configuration identifier, normally "ACEseq"
# 3: Configuration directory
# 4: Dataset identifier / PID
# 5: Control bam file
# 6: Tumor bam file
# 7: Control bam sample name
# 8: Tumor bam sample name
# 9: Reference files path
# 10: Output folder
# 11: Optional: The SV file
```

An example call is:

```
bash roddy.sh run ACEseq ./config/ stds /home/roddy/someproject/control_MB99_merged. 
 \rightarrowmdup.bam /home/roddy/someproject/tumor_MB99_merged.mdup.bam control tumor /icgc/ngs_ \rightarrowshare/assemblies/hg19_GRCh37_1000genomes ./output
```

Here you tell roddy to run the ACEseq configuration using the config folder in the current directory with a control and tumor bam. Also you tell Roddy the samples for both files namely control and tumor. Finally, you supply the path to the reference files and the folder where you will store your output data.

QuickStart

To start ACEseq download package from here and install the reference files and conda package as described under *Installation & Run instructions*.

```
sh $PATH_TO_PLUGIN_DIRECTORY/Roddy/roddy.sh rerun ACEseq@copyNumberEstimation $pid \
--useconfig=$PATH_TO_PLUGIN_DIRECTORY/applicationProperties.ini \
--cvalues="bamfile_list:$pathToControlBamFile;$pathToTumorBamFile,sample_list:control;
--tumor,possibleControlSampleNamePrefixes:control,
--possibleTumorSampleNamePrefixes:tumor"
```

Following parameters should be changed in the project.xml:

- baseDirectoryReference
- · outputBaseDirectory
- outputFileGroup (in case all outputfiles should have different group than primary group)

Alternative running modes:

- runWithoutControl (in case it should be run without control)
- runwithFakeControl (in case the coverage should be taken from a different control)

Requirements

5.1 Hardware

ACEseq requires the execution of multiple jobs that are highly parallelized in the beginning but linearize towards the end of the workflow. It requires a maximum of 50g RAM in few of the Jobs. On a HPC cluster with multiple cores available it will usually finish within 24h (100-160 CPU h). The final output usually requires between 4 and 6g memory.

5.2 Software

The installation of all required software can be found under *Installation & Run instructions*.

Alternative Running Modes

Multiple alternative running modes are enabled with ACEseq.

6.1 Run Without Control

If no control sample is available, but ACEseq was already used to process other tumor sample pairs one of their control coverage profile can be used for normalization. In this case the patient's sex needs to be set with PATIENT-SEX="male|female|k|inefelter".

Please specify the path and prefix to a control coverage profile for a male (MALE_FAKE_CONTROL_PRE) and a female patient (FEMALE_FAKE_CONTROL_PRE) so it can be matched to the processed sample. To activate this option the value runWithout control needs to be set to 'true', either via the command line execution under cvalues or in the project.xml.

6.2 Run quality check only

In case you do not want to run the full ACEseq pipeline immediately, but would rather access the sample's quality first you can start ACEseq with the option "runQualityCheckOnly" set to "true".

6.3 Replace low quality control

If a control sample is very noisy and masks CNAs it can be replaced with the coverage profile from a different control of the same sex. For this run ACEseq with "runWithFakeControl" set to "true" and specify the values "FEMALE_FAKE_CONTROL_PRE" and "MALE_FAKE_CONTROL_PRE" as described in the section for analysis without matched control.

6.4 Run with/without SV breakpoint incorporation

To process samples with incorporation of SV breakpoints set the following in the project.xml:

If the bedpe file does not exist ACEseq will submit all steps until the bedpe file is required. A rerun once the SV file is generated will start the pipeline up from the point where SV breakpoints are incorporated.

To process a samples without SVs please set the following in the project.xml:

```
<cvalue name='runWithSv' value='false' type="boolean"/>
<cvalue name='SV' value='no' type="string"/>
```

$\mathsf{CHAPTER}\ 7$

Input Parameters

Multiple parameters can be set with ACEseq though not all are necessary to change. This table gives and overview and description for all available parameters

Table 7.1: "ACEseq parameters"

aceseqOstputDirectsputSisBaseDirectory}/ACEseq_\${tumorSample} svOutputDirectory pathsisBaseDirectory}/SV_calls crestOutputDirectory pathsisBaseDirectory}/crest cnvSnpOstputSiqOpathstDirectory}/cnv_snp impute OstputSiqOpathstDirectory}/cnv_snp impute OstputSiqOpathstDirectory}/cnv_snp impute OstputSiqOpathstDirectory}/plasing plotOutpst(Directory)/plasing plotOutpst(Directory)/plots runWithoutsControl boolean use control for analysis (falseltrue) integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer minimum coverage in control for SNP cnv_min5000crage integer minimum coverage for lkb windows to be considered for merging in 10kb windows mapping1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESS_F float float fparameter for R lowess function SCALE_GRCTOR float scale_factor for R lowess function COVER_AGEPLOTENALIMS ylims for Rplots in GC-bias plots FILEN_ASTRIGOTEGETYPESTRIGOPROSESTRIGOSTRIGORNICORNICORNICORNICORNICORNICORNICORNIC	name	value	type	description		
crestOutsution_analysisBaseDirectory]/crest cnvSnp OntpotRincopephitDirectory]/cnv_snp impute OstpatRincopephitDirectory]/phasing plotOutsution_analysis (faiseItrue) minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer cnv_min5000erage integer minimum coverage in control for SNP cnv_min5000erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping_1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_10180 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_10110 float f parameter for R lowess function SCALE_GENCTOR float scale_factor for R lowess function SCALE_GENCTOR float scale_factor for R lowess function SCALE_GENCTOR_float scale_factor for R lowess function FILENAMPIction_quarrectory_float_files_pload_scance/bard_ng/gfter correction GC_bias_gisbinakey string key in GC-bias json FILE_DENGERS(DENGERS)_float_files_pload_scance/bard_ng/gfter correction min_DD101000gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration	aceseq	O stputPire	Atquarky hsisBa	seDirectory}/ACEseq_\${tumorSample}		
cnvSnpOstpoetRinCeptabletDirectory}/cnv_snp impute Ostpatistic patabletDirectory}/phasing plotOutpstRincettOptpattDirectory}/plots runWithoutControl boolean use control for analysis (falseltrue) minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_minScoverage integer minimum coverage in control for SNP cnv_minSoOcrage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping_1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_10180 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_fattal float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESS_F float fparameter for R lowess function SCALE_GSACTOR float scale_factor for R lowess function COVER_AGEPLOTfloatIMS ylims for Rplots in GC-bias plots FILENAMFIGOURGABREGOTSUSSACESTACETION key for GC-bias json FILE_DENASERGOTSUSSACESTACETION integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
impute OstpartDirectory)/phasing plotOut pstDirectory option Directory)/plots runWithoutControl boolean use control for analysis (falseltrue) minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer minimum coverage in control for SNP cnv_min5000erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_winslows integer minimum number of 1kb windows fullfilling cnv_min_coverage and map- ping_quality to obtain merged 10kb windows min_X_r0x00 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_r0x10 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESO_F float f parameter for R lowess function SCALE_GEACTOR float scale_factor for R lowess function COVERAGEPLOTfl0x1x1MS ylims for Rplots in GC-bias plots FILEN_AMFl0x00_ganRREGGTx01x18xibis)ogdcstome/chardngs/gfter correction GC_biasgistomaskey string key in GC-bias json FILE_DSN&ESO(SPIRMATDirectory)/densityBeta.pdf min_DD10x00gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
plotOutp\${Ritesect_OptphiletDirectory}{plots} runWithdulscontrol boolean minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer minimum coverage in control for SNP cnv_min5e00erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping1q0dlity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and map- ping_quality to obtain merged 10kb windows min_X_r0x80 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_r0x10 float f parameter for R lowess function SCALE_0F9CTOR float scale_factor for R lowess function SCALE_0F9CTOR float scale_factor for R lowess function COVERAGEPLOTENSALIMS FILEN_AMF1C000_GOBRECGT_PINOLOGY_densityBeta.pdf min_DD1000gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
runWithoutControl boolean use control for analysis (falseltrue) minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer minimum coverage in control for SNP cnv_min5000erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_ranko float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_ranko float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESS_F float f parameter for R lowess function SCALE_FRACTOR float scale_factor for R lowess function COVER AGEPLOTENATIMS ylims for Rplots in GC-bias plots FILENAMFLOCOLOGORRECOTENATIMS key in GC-bias json FILE_DENAMERACOTENATION for DEL/DUP/INV to be considered for breakpoint integration						
minHT 5 integer minimum number of consecutive SNPs to be considered for haploblocks snp_min5coverage integer minimum coverage in control for SNP cnv_min5000erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows (maximum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_winflows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_0000 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_00102 float float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVER_AGEPLOTfloatIMS ylims for Rplots in GC-bias plots FILENAMFloatOn_GOURGERECOT_RSUM_blogdocome/chardnpy/gfter correction GC_biasgisonaskey string key in GC-bias json FILE_DISNASERSUBJECTOR float integer minimum length for DEL/DUP/INV to be considered for breakpoint integration	-		** *	¥ * ±		
snp_min5coverage integer minimum coverage in control for SNP			l boolean			
cnv_min5000erage integer minimum coverage for 1kb windows to be considered for merging in 10kb windows mapping1000lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_win500ws integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_ratso float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_ratio float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWES9_F float fparameter for R lowess function SCALE_GEACTOR float scale_factor for R lowess function COVERAGEPLOTfivalims ylims for Rplots in GC-bias plots FILENAMplotoutfiathrecology/station/logodecone/chardpy/gfter correction GC_biasgisonaskey string key in GC-bias json FILE_DISNASERAGEPIANTDirectory}/densityBeta.pdf min_DD1000gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration				•		
mapping 1000 lity integer minimum mapping quality for 1kb windows to be considered for merging in 10kb windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_ranso float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_ratio float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESS_F float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVERAGEPLOTENATIONS ylims for Rplots in GC-bias plots FILENAMELGOO_GRAREGER_SINGLEGER_						
windows (maximum mappability) min_windows integer minimum number of 1kb windows fullfilling cnv_min_coverage and mapping_quality to obtain merged 10kb windows min_X_ratso float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_ratso float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESS_F float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVERAGEPLOTflvatIMS ylims for Rplots in GC-bias plots FILENAMPLation_fare.com_fare.co			e integer			
min_winflows integer minimum number of 1kb windows fullfilling cnv_min_coverage and map- ping_quality to obtain merged 10kb windows min_X_rate float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_rate float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWES9_IF float f parameter for R lowess function SCALE_GENCTOR float scale_factor for R lowess function COVERAGEPLOTfloatIMS ylims for Rplots in GC-bias plots FILENAMFload float fight for C-bias json FILE_DENAMERATE float key string key in GC-bias json FILE_DENAMERATE float for C-bias json FILE_DENAMERATE float float for DEL/DUP/INV to be considered for breakpoint integration	mappin	g <u>1</u> q00lity	integer			
ping_quality to obtain merged 10kb windows min_X_ratso float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_ratio float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESD_F float f parameter for R lowess function SCALE_GENCTOR float scale_factor for R lowess function COVER_AGEPLOTfivatiMS ylims for Rplots in GC-bias plots FILEN_AMFLGGOUTGABREGGT_VARATION_Sylims_log_dence/charlng/after correction GC_biasgisbiaskey string key in GC-bias json FILE_DENCER_AGEPLOTIMALDirectory}/densityBeta.pdf min_DDIGOOgth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
min_X_rat80 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as female min_Y_rat12 float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWES9_F float f parameter for R lowess function SCALE_GEACTOR float scale_factor for R lowess function COVERAGEPLOTffvatIMS ylims for Rplots in GC-bias plots FILENAMFlc@OutgatRecotrys/Ratione/cherlng/gfter correction GC_biasgisbiaskey string key in GC-bias json FILE_DENAMERAGEPLOTffvatIDirectory}/densityBeta.pdf min_DD1000gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration	min_w	in 5 lows	integer			
over whole genome to be considered as female min_Y_ratio float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESD_IF float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVERAGEPLOTflvatIMS ylims for Rplots in GC-bias plots FILEN_AMFLGGOUGABREGGTVLAMADOLOGOU						
min_Y_ratia float minimum ratio for number of reads on chrY per base over number of reads per base over whole genome to be considered as male LOWESD_IF float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVERAGEPLOTfivatIMS ylims for Rplots in GC-bias plots FILENAMFICOUNTARECTIVELYMAD option correction GC_biasgisbiaskey string key in GC-bias json FILE_DISNAERAGEPLOTIVALUTIVECTORY density Beta.pdf min_DDIOWOOTH integer minimum length for DEL/DUP/INV to be considered for breakpoint integration	min_X	_1 0.18 0	float			
over whole genome to be considered as male LOWESD_IF float f parameter for R lowess function SCALE_GRACTOR float scale_factor for R lowess function COVERAGEPLOTfIVATIMS ylims for Rplots in GC-bias plots FILENAMFIGOU GRAREGO PLANT September September				•		
LOWESO_F float f parameter for R lowess function SCALE_GACTOR float scale_factor for R lowess function COVERAGEPLOTffvatIMS ylims for Rplots in GC-bias plots FILENAMFlc@outfatBRE@TryPlank* Ogbe@fone/cherlnp/gfter correction GC_biasgisbiaskey string key in GC-bias json FILE_DENAMFR@TryPlantDirectory}/densityBeta.pdf min_DDI@Ogth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration	min_Y	_rat1a	float			
SCALE_GACTOR float scale_factor for R lowess function COVERAGEPLOTflyatiMS ylims for Rplots in GC-bias plots FILENAMFLGGUIGGARREGTryPISMEDD og destone/cherlqpy/gfter correction GC_biasgisbiaskey string key in GC-bias json FILE_DISMESSAGEFUNDANT Directory }/densityBeta.pdf min_DDI_GOOgth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
COVER AGEPLOT floatIMS ylims for Rplots in GC-bias plots FILENA MFLOTOL from Record from State of Sta						
FILENAMFICOUNTABRECT Type ISOME of Jog Obst Connected Ingular Correction GC_biasgistic String key in GC-bias json FILE_DISNAMERA (Bip in the Correction) / density Beta.pdf min_DDIO Object integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
GC_biasgisbiaskey string key in GC-bias json FILE_DISNASERAGE Tribut Directory }/density Beta.pdf min_DDI@@gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration				• 1		
FILE_DISNAMERACE FROM the Directory density Beta.pdf min_DDI_000 gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
min_DDI_01000gth integer minimum length for DEL/DUP/INV to be considered for breakpoint integration						
	min_D	D1 <u>0</u> 10200 gth	integer	minimum length for DEL/DUP/INV to be considered for breakpoint integration		

Table 7.1 – continued from previous page

Table 7.1 – continued from previous page					
name		type	description		
	Co ewent Sco		column from bedpe file to be recored in \${pid}_sv_points.txt file		
	g <u>2</u> 0000th	integer	segmentByPairedPSCBS() minwidth parameter in PSCBS R package		
undo_S	SD24	integer	segmentByPairedPSCBS() undo.SD parameter in PSCBS R package		
	or u ne_heig		pruneByHClust() parameter h in PSCBS R package		
min_se	g íoh.6 nt_ma	p float	minimum average mappability over segment to be kept after segmentation		
min_se	g <u>9</u> 10000gth_p	rumteger	maximum of segment to be considered for merging to neighbouring segment prior		
			to clustering		
min_nu	ım <u>1.5</u> SNPs	integer	maximum number of SNPs in segment to be considered for merging to neighbouring segment prior to clustering		
clusteri	กพอร	string	should segments be clustered (yeslno), coerage and BAF will be estimated and as-		
Clusteri	11803	sumg	signed clusterwide		
min cl	uster_numl	peinteger	minimum number of clusters to be tried with BIC		
	en0d&ership		obsolete		
min_di		float	min_distance		
	co luppHideRr e		prefix for file with haplogroups per chromosome		
	o u ntFileSu		suffix for file with haplogroups per chromosome		
1 0			ectory}/\${haplogroupFilePrefix}		
	ngt 000000 0		minimum length of segments to be considered for tumor cell content and ploidy		
	g.u. <u>s</u> p.u.u.,	11110801	estimation		
min he	t SNOP s_pu	itinteger	minimum number of control heterozygous SNPs in segments to be considered for		
			tumor cell content and ploidy estimation		
dh_sto	n max	string			
	n glt0<u>0</u>0d0 00t				
dh_zer		string			
purity_		float	minimum tumor cell content allowed		
purity_		float	i		
ploidy_		float			
ploidy_		float			
			editothe / Salpedis. changed the value for the filename pattern MUST also be changed.		
			RPHIM alue must be converted to a string because of a bug.		
	Usfnipilikab.gz		- ,		
	1 0				
CHR 1	NIS (CHR I	PREFIX \\$	{PARM_CHR_INDEX}\${CHR_SUFFIX}		
	-	CiHRg INI			
		D KakkA rra			
	{122}				
)				
CREST	yes	string	include SV breakpoints in analysis (yeslno)		
mpileu	p <u>Q</u> qual	integer	quality used for parameter 'Q' in samtools mpileup		
	MPIAEURP_		options for mpileup to determine which bases/reads to use		
	-B -Q				
	\${mpileu	p_qual}			
	-q 1 -I				
	"				
	V@F_SUF	string	suffix for vcf files		
	Γ Χ ΧΤΕ_SUF	string	suffix for txt files		
	G photyd<u>e</u>s		prefix for phased genotypes file		
unphased@phosypeshila@refix prefix for unphased genotypes file					
phased G& Diypes Vees not suffix for phased genotypes file					
unphased GFIbFyp SFIIns unphased genotypes file					
Continued on payt page					

Table 7.1 – continued from previous page

		Table 7.1 – continued from previous page	
	ype	description	
BCFTOOLS NOPTS	string	beftools options for imputation	
"			
FAKE CONTROLLS	1 gage 11	suffix for chromosome wise 1kb coverage files used for fake control workflow	
PATIENTES S	string	patient sex used in case of no control workflow (malelfemalelklinefelter)	
CNV_ANNOn66 Fall	Hig zag	suffix for mappability annotated chromosome-wise 1kb coverage files	
CNV_\$\(\mathbb{U}\)ff\(\mathbb{F}\)f\(\mathbb{E}\) b.gz s	string	suffix chromosome-wise 1kb coverage files	
FILE_UNPHASTED	na PruHDire	ctory}/\${unphasedGenotypesFilePrefix}	
FILE_UNPHASTED	DAGHENDOTE	MtPfy}/unphased_genotype_chr	
FILE PHAISIFIDATED	Math utDire	ctory}/\${phasedGenotypesFilePrefix}	
		Etory}/phased_genotype_chr	
	string	J.F6 J.F	
FILE_INIFO SANGE			
	string		
FILE_HARS_COME			
FILE_\$UMMARY s			
FILE_WARMINGS			
	string		
		csany)/samfile_usext by imputation on X chromosome for females	
		Relatify/and_sraf/\$ {toid hackmosome-wise 1kb coverage file used for fake control work-	
MALE EXAMENTATION	Marrika di 13 Gr	flow for male patients	
EEN (A LIECEARZIE M	2015/L/T/T/DO		
FEMALIS PARKINAP	parmedice	supartification in the state of	
DI OTE DANE		flow for female patients	
PLOT_PRaceseqOp			
FILE_MOSTAJIMARG			
FILE_M@&T_IMP6			
FILE_\$BGAdEssqOptoBtDPRectory}/\${pid}			
FILE_\$EGMENT_\$	_	ST	
	string		
outputFile@coupGro		group for output files and directories	
output Access Rights,	,0-	access rights for written files	
rwx			
output A acess Rightsl	Fxont Direct	carives ss rights for written directories	
rwx			
possible@ontrolSanb	pakehNamae)	Ppesisible prefix of control bam if named \${prefix}_\${pid}_\$mergedBamSuffix	
blood)			
possible Tutunon Stamb	okesNv AvrreaR	reainesas possibleControlSampleNamePrefixes	
	Ĭ	-	
reference Continuens	Hatti Rfeaf	reference genome file	
REFERENCE En Concor			
dbSNP_I\$I[pEth]/00e		•	
All.SNV.vc			
	_	rgMapability Align100mer_chr.bedGraph.gz	
CHROMOSOM FCHp			
REPLICATION/RHATE Treetidation intering the 10KB.Rda			
		37_100genomes_gc_content_10kb.txt	
_ \ \1 \7_\2		binhtheta_NAME}_combined_b37.txt	
		IHRONEAMHS : integrated_phase1_v3.	
		ls_svs.genotypes.nomono.haplotypes.gz	
		HOLD MAN Lintegrated_phase1_v3.	
20101123.si	nps_inde	ls_svs.genotypes.nomono.legend.gz	
		Continued on next nego	

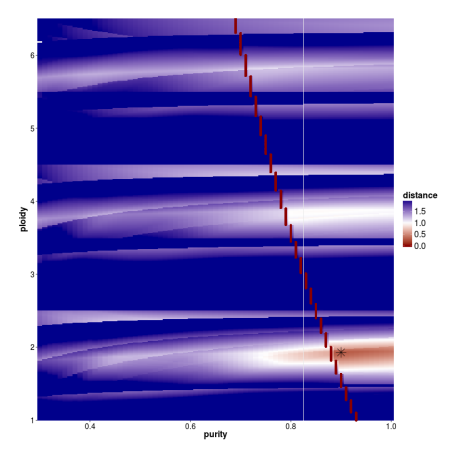
Table 7.1 – continued from previous page

10.0.100.0	مينامير	t	description
name	value	type	description
			pinhnXenfilmPAR_combined_b37.txt
KNOW	/N\$_{ H aAIP}_/@	ALEKTER KALEK	Ti_phpaXe Ifintes grated_v3_chrX_nonPAR_impute.hap.gz
KNOW	/N\$_{ [H a AI P]_/@	ALEKTER EXPLORED	Fighth Diffied grated_v3_chrX_nonPAR_impute.legend.gz
output	Ex&{paitbr}R	inectonfexe	c uptaid the Tion trues of the state of the
impute	B\$s{qrDihe}dt	o p ath	directory for impute files
merged	BanerSpetfin	ndatpiabgam	A list of all known suffixes for merged bam files. I.e. merged.dupmark.bam,
			merged.mdup.bam
merged	B&(mSerffex	lBistin S guffix	A list of all known suffixes for merged bam files. I.e. merged.dupmark.bam,
			merged.mdup.bam
default	Metgochea	HBáurfífg uffix	}The default suffix for merged bam files when they are created by Roddy.
libloc_	PSCBS	string	path to PSCBS library in R
libloc_	flexclust	string	path to felxclust library in R

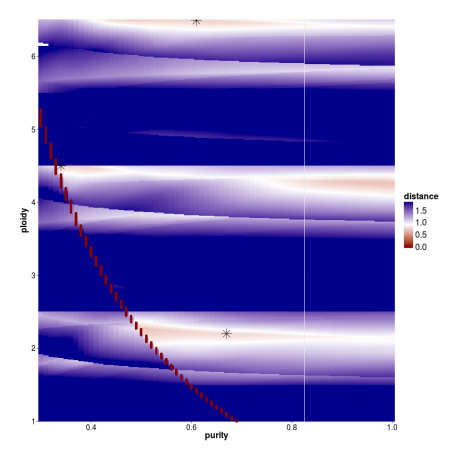
Purity Evaluation

Depending on the sample, ACEseq might return multiple possible solutions. These solutions can be found in the file \${PatientID}_ploidy_purity_2D.txt within the ACEseq results directory.

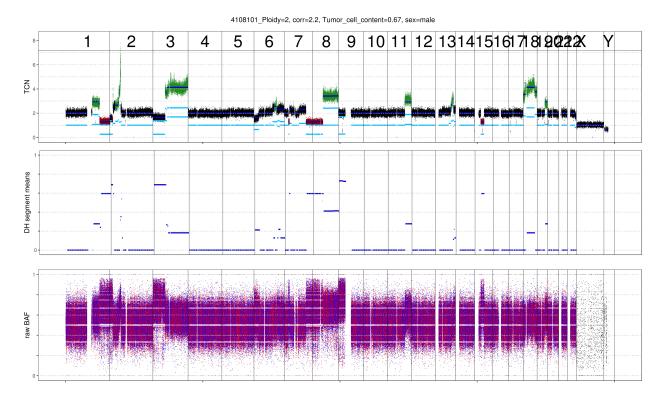
In addition to the table the plot ${\Phi}_{cm}$ addition to table the plot ${\Phi}_{cm}$ and ${\Phi}_{cm}$ addition to table the plot ${\Phi}_{cm}$ and ${\Phi}_{cm}$ and ${\Phi}_{cm}$ addition to table the plot ${\Phi}_{cm}$ and ${\Phi$



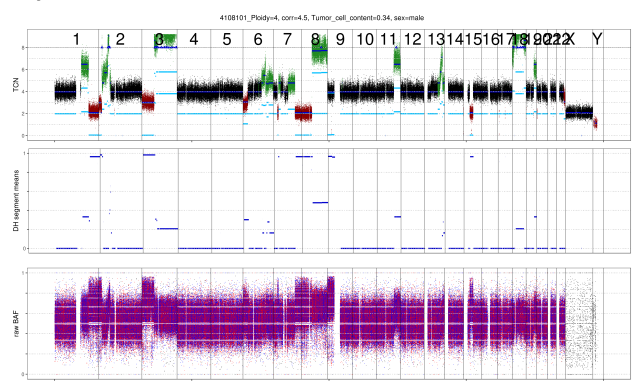
In case of multiple solutions, one can choose between the solution with the lowest distance or do as we suggest and choose the solution that is closest to diplot. It is recommended to make use of the prior knowledge about tumor biology as well as checking the final output as well as the mutant-allele frequency of a patient.



As can be seen below the benefit of increasing the ploidy of this sample to tetraploid leads to a clonal fit of multiple segments though many others remain subclonal (indicated by deviation from integer copy number states). This is often observed for heterogenous samples such as this shown lymphoma sample. Lymphoma tend to be diploid and heterogenous indicating that the diploid solution is correct. In addition we plotted the MAF distribution over balanced segments, that supports our assumption. Diploid solution:

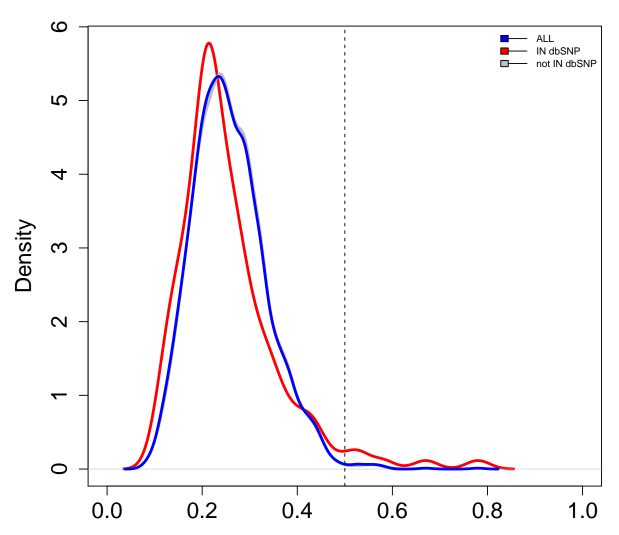


Tetraploid solution:



MAF distribution over balanced segments:

4108101 5.62 % in dbSNP



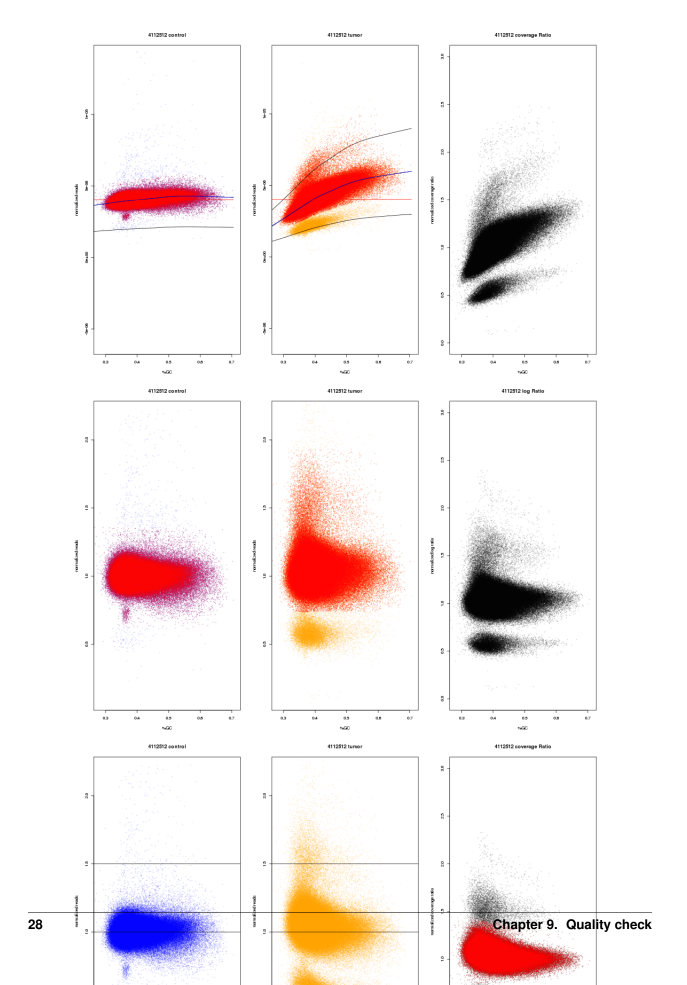
Mutant allele frequency from 2454 SNVs and 138 in dbSNP

					\cap
CH	JΛ	D_{\perp}		\Box	u
\cup r	1/4	г	ı⊏	П	J

Quality check

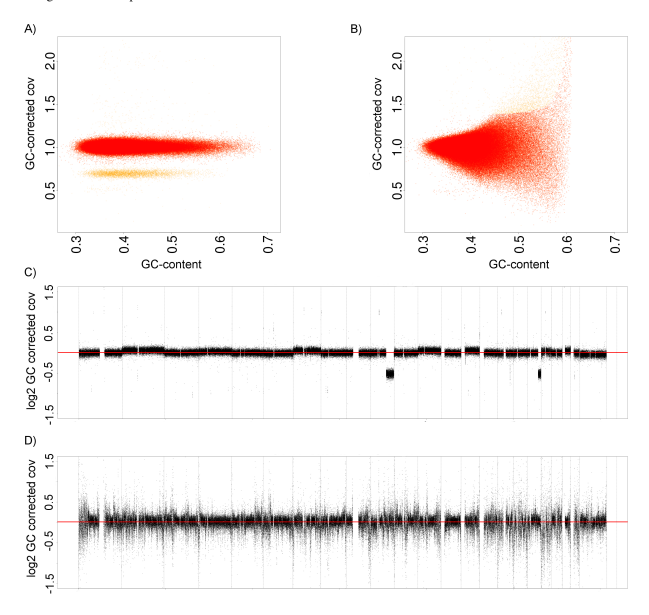
ACEseq provides a thorough quality check of the sample by investigation of the GC-bias: 1.differences in GC-bias between tumor and control 2.evenness of coverage in tumor and control

The following plot shows the raw GC bias for a healthy control (left), a corresponding tumor (middle) and the tumor/control ratio (right). The top row depicts raw data while the middle row indicates GC-bias corrected data and the bottom line indicates GC-bias and RT-bias corrected data.



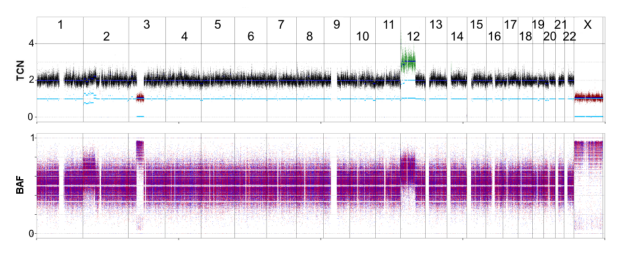
The file \${pid}_qc_gc_corrected.json provides information about slope, curvature and their differences between tumor and control. A strong diffrence between tumor and control can impact sensitivity and specificty of other variant calls.

The full width half maximum (FWHM) indicates the noise level within the majority copy number state in a sample. If it exceeds 0.205 in the control or 0.34 in the tumor a sample should be flagged with a warning. Yellow flagged tumors might have decreased specificity and sensitivity. Samples should be red flagged in case the FWHM exceed 0.29 in healthy controls of 0.54 in tumors. Red flagged tumor samples are very likely to accumulate artifacts due to a noisy coverage profile and should be excluded from further analysis Flagged controls can be rescued by rerunning the pipeline with "rerunWithFakeControl=true". The following plot shows a sample with low FWHM (A and C) and a sample with noisy coverage and thus high FWHM (B and D). No good copy number estimates can be obtained from the high FWHM sample.



Final Output

A graphical presentation of the final results is given for each tcc/ploidy solution. A general overview is give for the whole genome as shown here and per chromosome. Points represent raw SNP data points, colored by their copy number with regard to the majority copy number state (red:loss, black: neutral, green: gain). Segments are indicated by dark and light blue lines for total and allele-specific copy number respectively. Raw BAF are shown in the bottom panel which can be used to evaluate tcc and confirm allele-specific copy numbers.



Final results are provided in two formats. A reduced set of information is contained in the file ${\phi}_{most_info}_{poidy}_{purity}.txt$ while the full set is presented in ${\phi}_{most_info}_{poidy}_{purity}.txt$. A mapping of both headers and a corresponding description is given here.

Table 10.1: "Final output column headers"

most_important_info	comb_pro_extra	description
chromosome	chromosome	
start	start	start coordinate

Table 10.1 – continued from previous page

most_important_info	comb pro extra	description
end	end	end coordinate
SV.Type	crest	SV type connected to both or one
• •		breakpoint
length	length	length of segment
TCN	tcnMean	total copy number
NbrOfHetSNPs	tcnNbrOfHets	number of control heterozygous
		SNPs in segment
dhSNPs	dhMax	estimated DH
minStart	minStart	last SNP prior to segment start
maxStart	maxStart	first SNP after segment start
minEnd	minStop	last SNP prior to segment end
maxEnd	maxStop	first SNP after segment end
covRatio	tcnMeanRaw	bias corrected coverage ratio
dhEst	dhMean	raw DH
c1Mean	c1Mean	minor allele copy number
c2Mean	c2Mean	major allele copy number
genotype	genotype	ratio of rounded allele copy num-
		bers
CNA.type	CNA.type	DUP/DEL/LOH/TCNneutral
GNL	GNL	gain/loss/loh compared to diploid
	tcnNbrOfSNPs	number of SNPs per segment
	tcnNbrOfLoci	number of SNPs per segment
	dhNbrOfLoci	heterozygous SNPs per segment
	map	mappable/unmappable
	cluster	cluster assigned during merging
	neighbour	indicates whether neighbouring seg-
		ments exist on both sides
	distDH	distance to main cluster center DH
	errorSNP	error for DH direction
	distTcn	distance to main cluster center cov-
		erage ratio
	errorLength	error in coverage ratio direction
	totalError	sum of errorSNP and errorLength
	area	AUC ratio
	peaks	1 for balanced; 2 for imbalanced
	meanCovT	average total coverage per cluster
	meanCovB	average total coverage of B allele
	AF	allelic factor
	BAF	B-allele frequency
	A	rounded minor allele copy number
	В	rounded major allele copy number
	TCN	rounded copy number
	ploidy	majority copy number used as reference for CNA.type
	Sex	patient sex
	cytoband	cytoband

HRD, LST, TAI

HRD, LST and TAI scores are provided in the file \${pid}_HRDscore_\${ploidy}_\$tcc} for each solution.

HRD and LST are merged based on smoothed segments as suggested by Popova et al. (doi: 10.1158/0008-5472.CAN-12-1470).

TAI are based on unmerged segments. A combination of the three scores might be useful.

Methods - Theory

ACEseq can be used to estimate copy-numbers from WGS data using a tumor vs. control approach. Thus a pre-requesite is WGS data from healthy tissue and tumor tissue of the same patient with at least 30x coverage. Samtools [] mpileup is used to determine the coverage for tumor and control sample - position specific for each single nucleotide polymorphism (SNP) position recorded in dbSNP and per 1 kb window. To get chromosome specific allele frequencies, the genotypes of SNP positions are phased with Impute2 [] and A and B allele are assigned accordingly. Haploblocks are defined as regions with consecutively phased SNPs. Subsequently, B-allele frequencies (BAFs) are estimated for all SNP positions in tumor and control with sufficient coverage in the control:

$$BAF = \frac{cov_{SNP}^B}{cov_{SNP}^A + cov_{SNP}^B}$$

These can be converted to the decrease of heterozygosity a measure of the allelic state [Olshen et al.].

$$DH = 2 \times |BAF - 0.5|$$

12.1 Pre-processing

To estimate the coverage of each SNP position a general coverage of 10 kb windows was determined. 1 kb coverage windows are merged into 10 kb windows in case enough contributing windows with sufficient coverage and mappability are found in the corresponding region. The resulting coverage values are normalized with the sum of all 10 kb coverage windows for tumor and control respectively. These normalized estimates are subsequently corrected for a possible GC- and replication-timing bias.

12.2 GC-/Replication timing bias correction

12.2.1 Correction for GC bias

Correction for GC bias

As described in detail by Benjamini and Speed (REF) genomic regions with varying GC content may be sequenced at different depth due to selection bias or sequencing efficiency. Differing raw read counts in these regions even in the absence of copy number alterations can could lead to false positive calls. A GC-bias plot (Figure XY) can be used to visually inspect the bias of a sample. ACEseq first fits a curve to the data using LOWESS (locally weighted scatterplot smoothing, implemented in R) to identify the main copy number state first, which will be used to for a second fit. The second fit to the main copy number state is used for parameter assessment and correction of the data. This two-step fitting is necessary to compensate for large copy number changes that could lead to a misfit. The LOWESS fit as described above interpolates over all 10 kb windows. It thus averages over all different copy number states. If two states have their respective center of mass at different GC content, this first LOWESS fit might be distorted and not well suited for the correction. The full width half maximum (FWHM) of the density over all windows of the main copy number state is estimated for control and tumor. An usual large value here indicates quality issues with the sample.

12.2.2 Correction for replication time

Once the data is corrected for GC-bias the replication timing bias is considered. In general, if a fraction of the cells in the analyzed sample is cycling, early replicating regions would be expected to display higher coverage than late replicating regions, as a higher percentage of these would already have undergone replication in the S-phase [Zitat Koren et al.]. For a subtle analysis of copy number alterations, it would be beneficial to correct for this replication timing bias. Large fractions of the genome have common replication timing in different cell types or tissues, but there are regions of tissue or organ specificity [] [Zitat RepliSeq]. In the present work, a consensus replication timing score, the RepliSeq score as described by [] [Zitat RepliSeq], is attributed to every 10 kb window of the genome by averaging over the RepliSeq information from different cell lines. Replication timing bias plots can be generated analogously to the GC bias plots. A LOWESS fit on the already identified main cluster is carried out to correct for this bias (Figure?). This correction is performed on the GC-corrected data to obtain the final corrected coverage data, which will be used in the following.

The two bias correction steps described above are done sequentially. A simultaneous 2D LOWESS or LOESS correction would be desirable, but fails due to computational load (the clusters to be fitted have 106 points). Different parameters such as slope and curvature of the both LOWESS correction curves used are extracted. The GC curve parameters is used as quality measures to determine the suitability of the sample for further analysis whereas the replication timing curve parameters is used to infer the proliferation activity of the tumor. We could show a strong correlation between Ki-67 estimates and the slope of the fitted curve (Figure).

Once corrected a coverage ratio is calculated as the ratio of normalized tumor coverage over normalized control coverage:

$$covR = \frac{covT_{window}^{corrected}}{covN_{window}^{corrected}}$$

Finally SNP and coverage data are merged. Regions without coverage or SNP information are discarded.

12.3 Segmentation

Once data pre-processing is completed the genome is segmented with the PSCBS (parent specific circular binary segmentation) [] (Version!!!) algorithm. Prior to the actual segmentation, segment-boundaries due to a lack of coverage are determined. Single outliers among the coverage and very low coverage regions are determined using PSCBS functions. In addition to these, breakpoints that are indicated by previously called structural variations are taken into account. During the actual segmentation step the genome is segmented based on the pre-defined breakpoints, changes in the coverage ratio and DH. DH values are only considered in case the SNP position is heterozygous in the control.

12.4 Segment reliability

Homozygous deletions are called in segments that lack mapped reads. These deletions are only considered to be true in case the low read count is unlikely to be caused by a low mappability. Thus, the mappability is assessed for all segments. Regions with mappability below 60% are considered unmappable and not further considered for copy number estimation. Each SNP position is annotated with the new segment information and mappability.

12.5 Segment clustering and merging

In order to avoid over-segmentation short segments (default <9 kb) are attached to the closest neighboring segment according to the coverage ratio. Subsequently, segments from diploid chromosomes are clustered according to the log2 of the coverage ratio and DH. These values are scaled prior to clustering. The DH of a segment is defined as the most commonly found DH value among all SNPs in the segment that are heterozygous in the control. In a first step, c-means clustering is performed. The segments are weighted according to the log2 of their length. A minimum number of one clusters is required allowing up to 20 clusters and the optimal cluster number is determined with BIC clustering :raw-latex: \cite{}. The number is used to cluster the points with cmeans subsequently (with the R fpc package clusterboot function).

To avoid over-fitting a further downstream processing is applied. Firstly, the minimal accuracy defined by the FWHM is taken into account. Cluster with more than 85% of all points within these coverage limits are chosen. Of these the cluster with most segments is defined as main cluster. The other chosen clusters are merged with the main cluster if their the difference between their center and the main cluster center is not bigger than XX times the DH-MAD of the main clusters. Neighboring segments are merged before new cluster centers are determined. In a second step segments that are embedded within main cluster segments are considered for merging. The number of control heterozygous SNP positions and the length are considered here to establish two criteria. Segments with less than 5 heterozygous SNPs are merged with the main cluster if they lie between the FWHM boundaries. Additionally, error values defining the tolerable deviation from the main cluster center is defined both for DH and coverage value as follows:

$$error DH = \frac{1}{\sqrt{number of heterozygous SNPs}}$$

$$error Coverage = \frac{1}{log2(length)}$$

If the SNP error of a selected segment exceeds the distance in DH and the length error exceeds the coverage difference it is appointed to the main cluster. Again neighboring segments with identical clusters are merged. Finally,

12.3. Segmentation 37

a general cluster coverage is estimated from all relevant segments and assigned to the cluster members to further reduce noise in the data.

12.6 Allelic adjustment

To get better estimates of a segments allelic state as balanced or imbalanced the phasing and segmentation information are combined. Within an imbalanced segment the more prominent allele should be consistently assigned to the same allele across all haploblocks. For balanced segments a haploblock-wise swap of A- and B-allele should have no effect. Thus, the median tumor BAF is calculated haploblock-wise for all SNP positions that are heterozygous in the control. If it is below 0.5 A- and B-allele are swapped within the haploblock region to get consistency across the haploblocks of a segment. This procedure ensures a more accurate estimation of the allelic state of a region in the next step.

12.7 Calling of Allelic Balance and Imbalance

In order to be able to identify the allelic state of a segments, a first test to distinguish between allelic balance and imbalance of a segment independent from the degree of imbalance was implemented. Our method evaluates the area under the BAF density curve left and right of 0.5. Balanced segments should have an equal area and the allelic state of a segment can be defined by equation [eq:areaDiff], i.e. computing the absolute value of the relative difference between the left and right area.

$$diff A_{segment} = \frac{|A_{right} - A_{left}|}{A_{right} + A_{left}}$$

For balanced segments $diffA_{segment}$ should be close to zero, whereas this value should shift more towards one for imbalanced segments. Thus, a cut-off to differentiate between balanced and imbalanced segments is needed. In the following we propose a way to establish a dynamic and sample dependent cut-off. In case a sample has several segments that correspond to different states, e.g one balanced and one imbalanced state, these will be represented by different peaks in the density distribution of $diffA_{segment}$. Hence the minima between the peaks can be used as cut-off. Corresponding to the above reasoning peaks further left in the distribution are more likely to represent balanced states. The minimum that differentiates a balanced from an imbalanced state varies across different samples. Potentially this depends on the relative contribution of copy number states, tumor cell content, contamination, subpopulations and sequencing biases. Empirically the discrimination is optimal for cut-off values in the range of 0.25 and 0.35. The minimum value of the density function within this interval is chosen as cut-off. The allelic state is only evaluated for segments on diplod chromosomes that fullfill certain quality criteria in order to ensure confident calls. Once $diffA_{segment}$ was calculated for a segment and the overall cut-off determined segments that exceed the cut-off are classified imbalanced. Segments below the cut-off are classified as balanced.

12.8 Copy Number Estimation

Once the allelic state of a segment is determined it can be used for the computation of tumor cell content and ploidy of the main tumor cell population. The average observed tumor ploidy can be determined with equation [eq:averagePloidy].

$$D_t = p_t \times P_t + 2 \times (1 - p_t)$$

Where p_t is the tumor purity and P_t is the tumor ploidy. Using the observed tumor ploidy and the coverage ratio of a segment (covR:math:_{segment}), the total copy number of a segment can be estimated as follows:

$$TCN_{segment} = \frac{covR_{segment} \times D_t - 2 \times (1 - pt)}{p_t}$$

This can be used subsequently to obtain the real BAF value for each segment by converting the coverage data to a copy number. The allelic factor (AF) is introduced for this as a segment-wise conversion measure.

$$AF_{segment} = \frac{\frac{covT_{segment}^{norm}}{10000}}{p_t \times TCN_{segment} + 2 \times (1 - p_t)}$$

 $covT_{segment}^{norm}$ represents the observed tumor coverage of a segment. The factor $\frac{1}{10000}$ is introduced to get from the initial 10 kb window coverage to a per base pair coverage. The BAF value of a segment can be calculated as follows.

where $covT_{segment}^{B}$ is the observed tumor coverage of a segment. The BAF value can now be used to calculate the DH of a segment according to [eq:DH]. Finally the allele-specific copy numbers are estimated.

$$TCN_{segment}^{B} = \frac{1}{2} \times TCN_{segment} \times (1 - DH_{segment})$$

$$TCN_{segment}^{A} = TCN_{segment} - TCN_{segment}^{B}$$

12.9 Purity and ploidy estimation

To obtain actual copy numbers for each segment ploidy and tumor cell content of the tumor sample have to be inferred from the data. Information about the allelic state of a segment is combined with TCN, DH and allele-specific copy numbers calculations. The combination of ploidy and tumor cell content that can explain the observed data the best is to be found. Possible ploidies in the range from 1 to 6.5 in steps of 0.1 and possible tumor cell content from 30% to 100% in steps of 1% are tested. The evaluation is done based on the distance of all segments from their next plausible copy number state. Imbalanced segments are fitted to a positive integer value.

$$distance_{tcn_imbalanced} = abs(TCN_{segment} - round(TCN_{segment}))$$

In addition the allele specific copy number is estimated according to equation [eq:TCNb] and [eq:TCNa]. For each allele a distance is defined accordingly:

$$\begin{aligned} distance_{tcn_a_imbalanced} &= abs(TCN_{segment}^{A} - round(TCN_{segment}^{A})) \\ distance_{tcn_b_imbalanced} &= abs(TCN_{segment}^{B} - round(TCN_{segment}^{B})) \end{aligned}$$

The total distance as quality measure of a fit is defined as the sum of the distances.

$$distance_{segment_imbalanced} = distance_{tcn_imbalanced} + distance_{tcn_a_imbalanced} + distance_{tcn_b_imbalanced} + dist$$

Balanced segments can only be fitted to even total copy numbers. The distance is defined as follows:

$$\begin{aligned} distance_{tcn_balanced} &= \frac{TCN_{segment}}{2} - floor(\frac{TCN_{segment}}{2}) \\ &?identicalto \\ distance_{tcn_balanced} &= abs(\frac{TCN_{segment}}{2} - round(\frac{TCN_{segment}}{2})) \times 2 \end{aligned}$$

As both alleles are expected to be present in equal numbers the allele specific copy number as well as the total distance can be derived.

$$\begin{aligned} distance_{tcn_a_balanced} &= distance_{tcn_b_balanced} = \frac{distance_{tcn_balanced}}{2} \\ distance_{segment_balanced} &= distance_{tcn_balanced} + distance_{tcn_a_balanced} + distance_{tcn_b_balanced} \\ &= 2 \times distance_{tcn_balanced} \end{aligned}$$

For each ploidy and tumor cell content combination a mean distance is defined by using the segment length as weights:

$$meanDist(p_t, P_t) = \frac{\sum_{1:N_{segments}}^{i} (distance_{segment_i} * length_{segment_i})}{\sum_{1:N_{segments}}^{i} length_{segment_i}}$$

All segments on diploid chromosomes that exceed a pre-set length and contain a sufficient amount of heterozygous SNP positions are used for the estimation. The smaller the distance the more likely a combination is chosen as final solution. Combinations of ploidy and tumor cell content that lead to negative copy numbers or exceed the DH limits are excluded as solution and used to set a minimum limit.

12.9.1 Final output

Once the optimal ploidy and tumor cell content combinations are found the TCN and allele-specific CN will be estimated for all segments in the genome and classified (gain, loss, copy-neutral LOH, loss LOH, gain LOH, sub). If a segments TCN is further than 0.3 away from an integer value it is assumed to originate from subpopulations in the tumor sample that lead to gains or losses in part of the tumor cell population.

Index

```
C contents table of, 1

T table of contents, 1
```