TEQC: Target Enrichment Quality Control

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1 Introduction

With whole genome sequencing it is still rather expensive to achieve sufficient read coverage for example for the detection of genomic variants. Further, in some cases one might be interested only in some fraction rather than the whole genome, for example linkage regions or the complete exome. Target capture (target enrichment, targeted sequencing) experiments are a suitable strategy in these situations. The genomic regions of interest are selected and enriched previous to next-generation sequencing. A frequently used application for the enrichment of the target sequences is based on hybridization with pre-designed probes, either on microarrays or in solution. The hybridized molecules are then captured (eluted from the microarrays or pulled-down from the solution, respectively), amplified and sequenced.

Besides quality control of the sequencing data, it is also crucial to assess whether the capture had been successful, i.e. if most of the sequenced reads actually fall on the target, if the targeted bases reach sufficient coverage, and so on. This package provides functionalities to address this issue. Quality measures comprise specificity and sensitivity of the capture, enrichment, per-target read coverage, coverage uniformity and reproducibility, and read duplicate analysis. The coverage can further be examined for its relation to target region length and GC content of the hybridization probes. The analyses can be based on either single reads or read pairs in case of paired-end sequence data. Results are given as values (e.g. enrichment), tables (e.g. per-target coverage), and several diagnostic plots. The package makes use of data structures and methods from the *IRanges* package, which makes dealing with large sequence data feasible.

TEQC does not include general sequencing data quality control (e.g. Phred quality plots), neither tools for sequence alignment. It also does not provide functionalities for follow-up analysis like SNP detection.

2 Automated html Report

The fastest way to run TEQC quality analysis is to launch the html report generation. The main arguments for TEQCreport are tables containing positions of sequenced reads and of genomic targets, or information about how to read the respective data files. There are several other options for customization of the report. In especially, you can choose whether or not to produce certain figures or coverage wiggle files and whether the data should be treated as single- or pairedend data. Calculation-intensive R objects can also be saved as workspace, for further usage. With option destDir you can specify the output folder, where all results will go and where you will find the index.html page of the html report. The parameters sampleName, targetsName and referenceName simply allow to include information about the sample and experiment in the report. For details about the different steps, options and results of a TEQC analysis, it is recommended to read also the following sections of this documentation.

```
> library(TEQC)
> exptPath <- system.file("extdata", package="TEQC")
> readsfile <- file.path(exptPath, "ExampleSet_Reads.bed")
> targetsfile <- file.path(exptPath, "ExampleSet_Targets.bed")
> TEQCreport(sampleName="Test Sample",
+ targetsName="Human Exome",
+ referenceName="Human Genome",
+ destDir="report",
+ reads=get.reads(readsfile, skip=0, idcol=4),
+ targets=get.targets(targetsfile, skip=0),
+ genome="hg19")
```

3 Load Reads and Targets Data

The (minimum) input needed for the quality control analysis are two files

• A bed file containing the genomic positions (chromosome, start, end) of the targeted regions, one genomic range per line. The targets might be custom designed or commercial solutions, e.g. for the capture of the whole human exome. The file does not have to be sorted with respect to genomic position.

• A bed or BAM file containing the genomic positions (chromosome, start, end) of sequenced reads aligned to a reference genome, one genomic range per line. In case of paired-end data an additional column with the read pair ID is suggested. The bed file format is very general and hence the QC analysis is not limited to any sequencing platform or alignment tool. The file does not have to be sorted with respect to genomic position. The BAM file format is now quite standard in NGS analysis. The usage of bam files has the advantage that reading the data is much faster (using functionality of the *ShortRead* package) than with bed files.

The package includes a small example data set. First we load the target positions. The input targetsfile does not need to have a fixed format. Just three columns containing chromosome (as string, e.g. chr1), start and end position of each target are required. The options chrcol, startcol and endcol specify in which columns of the file the respective information is found. The output of get.targets is of class RangedData from the IRanges package. Note that overlapping or adjacent targets are merged, such that the returned target regions are not-overlapping. Therefore, also a bed file containing information about the hybridization probes, which might be highly overlapping due to tiling, can be used as targetsfile.

```
> library(TEQC)
> exptPath <- system.file("extdata", package="TEQC")
> targets <- get.targets(targetsfile=paste(exptPath, "ExampleSet_Targets.bed", sep="/"),
+ chrcol=1, startcol=2, endcol=3, skip=0)

[1] "read 50 (non-overlapping) target regions"
> targets
```

RangedData with 50 rows and 0 value columns across 21 spaces

```
space
                              ranges
    <factor>
                           <IRanges>
        chr1 [ 11158025,
                           11158264]
1
2
        chr1 [ 25870174,
                           25870293]
3
        chr1 [ 65656333,
                           65656572]
4
        chr1 [ 68611504,
                           68611743]
5
        chr1 [ 70225862,
                           70226101]
6
        chr1 [112269528, 112270487]
7
        chr1 [160970804, 160970923]
8
        chr1 [182635998, 182636117]
9
        chr1 [186270664, 186270903]
42
        chr6 [152787105, 152787224]
43
        chr6 [170639528, 170639647]
        chr7 [144345876, 144345995]
44
45
        chr8 [ 1830798,
                            1830917]
```

```
46 chr9 [ 6241680, 6241799] | 47 chr9 [ 77277336, 77277575] | 48 chr9 [ 79827855, 79827974] | 49 chrX [ 47086386, 47086505] | 50 chrX [150891046, 150891285] |
```

NOTE: We assume that genomic positions in bed files follow the 0-based start / 1-based end coordinate system as defined by UCSC (http://genome.ucsc.edu/FAQ/FAQformat). In TEQC we need 1-based coordinates, so by default get.targets and get.reads (see later) shift all start positions forward by 1. If the coordinates in your files are already 1-based, set the parameter zerobased to FALSE in order to avoid the shifting.

We might ask what fraction of the genome is targeted. In the function fraction.target the corresponding genome can be specified by the option genome. At the moment only hg18 and hg19 are available. The corresponding genome sizes are taken from http://genomewiki.ucsc.edu/index.php/Genome_size_statistics. For any other case, you can specify the genome size manually with the option genomesize. In our little example the total targeted region and hence the fraction within the genome is very small.

```
> ft <- fraction.target(targets, genome="hg19")
> ft
```

[1] 4.158855e-06

Next, we load the genomic positions of the aligned reads. Depending on the number of reads, this can be quite time and memory consuming with real data. The function get.reads is quite similar to get.targets. However, overlapping or identical reads are not merged. Furthermore, a column containing read identifiers can be specified with option idcol. This is essential in case of paired-end data, when you want quality statistics to be done (also) on read pairs rather than on single reads. In this case the ID has to be the identifier of the read pair (i.e. the same unique ID for both reads of the pair). Our example data was derived by paired-end sequencing, so we keep the pair IDs in the resulting RangedData object. The genomic positions in our bedfile containing the reads are 1-based, so we avoid coordinate shifting by setting zerobased to FALSE.

```
> reads <- get.reads(paste(exptPath, "ExampleSet_Reads.bed", sep="/"),
+ chrcol=1, startcol=2, endcol=3, idcol=4, zerobased=F, skip=0)</pre>
```

[1] "read 19546 sequenced reads"

> reads

RangedData with 19546 rows and 1 value column across 24 spaces

	space	ranges	- [ID
	<factor></factor>	Ranges	- 1	<character></character>
1	chr1	[13328, 13381]	- [1_16_7090_2464
2	chr1	[13467, 13520]	- [1_16_7090_2464
3	chr1	[1420325, 1420378]	- 1	1_99_1631_6326
4	chr1	[1420402, 1420455]	- [1_99_1631_6326

```
5
                    [2321365, 2321418]
          chr1
                                               1_5_14614_17275
6
                    [2321479, 2321532]
          chr1
                                               1_5_14614_17275
7
          chr1
                    [2452643, 2452696]
                                               1_15_18642_6232
8
          chr1
                    [2452775, 2452828]
                                               1_15_18642_6232
9
          chr1
                    [2535217, 2535270]
                                                1_20_2015_5490
          chrX [154014582, 154014635]
                                                1_6_7123_16490
19538
19539
          chrX [154261641, 154261694]
                                                1_88_4216_1073
          chrX [154261777, 154261830]
19540
                                                1_88_4216_1073
          chrY [
                   3551497,
                                                1_87_3763_6007
19541
                               3551550]
19542
          chrY [
                   3551666
                               3551719]
                                                1_87_3763_6007
19543
          chrY [ 10028334,
                              10028387]
                                               1_63_1439_10606
19544
          chrY [ 10028434,
                              10028487]
                                               1_63_1439_10606
19545
          chrY [ 28425424,
                              28425477]
                                            1_105_17963_15521
19546
          chrY [ 28425590,
                              28425643]
                                           | 1_105_17963_15521
```

Whenever BAM files are available, it is recommended to use those instead of bed files as input for get.reads, while setting parameter filetype to "bam". In this case, get.reads makes use of the ShortRead function scanBam, with flag option is UnmappedQuery=FALSE. All other previously described options of get.reads are ignored, since genomic positions and read identifiers are taken automatically from the BAM file.

```
> reads <- get.reads("myBAM.bam", filetype="bam")</pre>
```

Paired-end data

When reads are paired, in order to perform statistics on pairs rather than on single reads, the read pairs have to be matched together using function reads2pairs. To run the function can be quite time consuming, depending on the number of reads. The output is a RangedData object whose ranges start at the first base of the first read within a read pair and end at the last base of the respective second read. This is equivalent to the positions of the DNA molecule that was actually sequenced from both ends. The reads table might also contain single reads (i.e. whose corresponding partners did not align to the reference). In this case a list of two RangedData tables will be returned, the first one containing the original positions of the single reads without partners (table singleReads), and the other one containing the merged pairs positions (table readpairs). The provided reads data might also contain cases where the two reads of a pair align to different chromosomes. Since for such read pairs a 'merging' does not make sense, they will be returned within the singleReads table. Further, for some pairs the respective reads might align very far apart within the same chromosome. If you wish to remove such reads, you can specify a value for option max.distance. Reads with a larger distance (from start position of first read to end position of second read) will be added to table singleReads.

```
> readpairs <- reads2pairs(reads)
> readpairs
```

```
RangedData with 9773 rows and 1 value column across 24 spaces space ranges | ID
```

```
<factor>
                             <IRanges>
                                                   <character>
1
         chr1
                      13328,
                                13520]
                                                1_16_7090_2464
2
         chr1
                   [1420325, 1420455]
                                                1_99_1631_6326
3
         chr1
                   [2321365, 2321532]
                                              1_5_14614_17275
4
         chr1
                   [2452643, 2452828]
                                              1_15_18642_6232
5
                   [2535217, 2535407]
                                                1_20_2015_5490
         chr1
6
                   [2814194, 2814357]
                                             1_36_17959_12842
         chr1
7
         chr1
                    [6488248, 6488458]
                                             1_58_10090_11513
8
         chr1
                   [6741034, 6741168]
                                                1_66_3980_7233
                   [7332529, 7332654]
9
         chr1
                                                1_85_5028_4821
           . . .
. . .
9765
         chrX [153700924, 153701112]
                                              1_26_10299_1105
               [153814666, 153814897]
                                             1_74_16848_10502
9766
         chrX
9767
         chrX [153851675, 153851897]
                                                1_14_5061_8480
         chrX [153997431, 153997581]
                                             1_100_7570_16280
9768
9769
         chrX [154014446, 154014635]
                                                1_6_7123_16490
         chrX [154261641, 154261830]
                                                1_88_4216_1073
9770
9771
                  3551497,
                              3551719]
                                                1_87_3763_6007
9772
         chrY [ 10028334,
                             100284877
                                              1_63_1439_10606
9773
         chrY [ 28425424.
                             284256431
                                            1_105_17963_15521
```

You may decide to use for all further analyses only "valid pairs", i.e. to exlude those reads that ended up in the singleReads table (in case there were any). This can be done e.g. in the following way

```
> reads <- reads[!(reads$ID %in% readpairs$singleReads$ID), , drop=TRUE]
```

Again, only for the case of paired-end data, we can visualize the read pair insert sizes, i.e. the distances from the start of read 1 to the end of read 2, respectively. For the function <code>insert.size.hist</code> we need the output of the previous call to <code>reads2pairs</code>. In our example, the alignment was done in a way that reads were only kept if the two reads aligned at a maximum distance of 250 bases. Therefore, the resulting histogram is truncated at 250, see figure 1. Note that also average and median insert sizes displayed in the graph are based on the truncated data, and hence are not statistics for the true read pair distribution. The insert sizes for all read pairs can also be returned using option returnInserts = TRUE.

> insert.size.hist(readpairs, breaks=10)

4 Specificity and Enrichment

One important component of quality control in target capture experiments is to check whether most of the sequenced reads actually fall on target regions. A barplot showing the numbers of reads aligning to each chromosome can give a first impression on that. When providing the function chrom.barplot only with the reads table, the resulting barplot will show absolute counts. There is also the option to give both the reads and the targets table, which will show fractions of reads and targets, respectively, falling on each chromosome. For

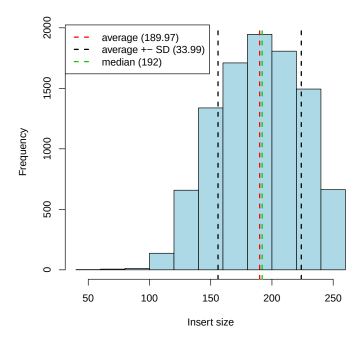


Figure 1: Histogram of read pair insert sizes.

the reads, this is the fraction within the total *number* of reads (since reads are usually expected to have all the same length). In contrast, for the targets, the fraction of targeted bases on each chromosome is calculated. Since targets might strongly vary in length it is reasonable to account for the actual target *sizes* instead of considering merely numbers of targets per chromosome. In this way you can compare directly if the amount of reads corresponds more or less to the amount of target on a certain chromosome (see figure 2).

> chrom.barplot(reads, targets)

A measure for the capture specificity is the fraction of aligned reads that overlap with any target region. It can be calculated by function fraction.reads.target. The function has an option mappingReads that can be set to TRUE in order to retrieve a reduced reads RangedData table containing only those reads overlapping target regions.

> fr <- fraction.reads.target(reads, targets)
> fr

[1] 0.3899007

In many of the functions within TEQC you can specify an "offset" that will enlarge every target on each side by the specified number of bases. Since usually the captured DNA molecules are longer than what is actually sequenced, it is

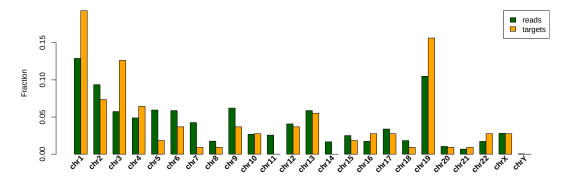


Figure 2: Fractions of reads (green) and target (orange) per chromosome.

expected to have many reads that do not overlap, but are close to the target. Considering e.g. the actual targets plus 100 bases on each side, we get a higher on-target fraction:

> fraction.reads.target(reads, targets, Offset=100)

[1] 0.4853679

With the fraction of on-target reads and the fraction of the targeted region within the reference genome, we can calculate the *enrichment*

$${\rm enrichment} = \frac{\# \ {\rm reads} \ {\rm on} \ {\rm target}/\# \ {\rm aligned} \ {\rm reads}}{{\rm target} \ {\rm size/genome} \ {\rm size}}$$

Since in our artificial example the total target size is unrealistically small, we achieve a huge enrichment.

> fr / ft

[1] 93751.94

Instead of considering single reads, we could also calculate the fraction of read pairs that are on-target. A read pair is counted as on-target if at least one of its reads overlaps with a target region or, in case of small targets, if the first read lies "left" and the second read "right" of the target and hence the corresponding sequenced molecule covered the target completely. For the specificity calculation the same function can be used, just the input changes from the table containing all reads to the table created above by reads2pairs.

> fraction.reads.target(readpairs, targets)

[1] 0.4934002

5 Coverage

Besides high capture specificity, it is of course important to check the read coverage within target regions, since it is crucial for follow-up analyses. The function coverage.target calculates read coverage for each base that is sequenced and/or located in a target region. It returns a list with the average (list element avgTargetCoverage), standard deviation (targetCoverageSD) and main quantiles (targetCoverageQuantiles) of the coverage over all targeted bases. When option perBase is set to TRUE, the returned list additionally has the two elements coverageAll and coverageTarget. The former one is a SimpleRleList containing coverages for all bases present in the reads table, the latter one contains coverages for all targeted bases. For some TEQC functions we need either one or the other. When option perTarget is set to TRUE, the returned list has the additional element targetCoverages. This is the input targets Ranged-Data table, now including as 'values' columns the average coverage (column avgCoverage) and standard deviation (column coverageSD) per target region. Coverage calculations might take a while, depending on the numbers of reads and targets.

```
> Coverage <- coverage.target(reads, targets, perTarget=T, perBase=T)
> Coverage
```

\$avgTargetCoverage
[1] 27.47927

\$targetCoverageSD
[1] 23.68119

\$targetCoverageQuantiles 0% 25% 50% 75% 100% 0 8 21 42 117

\$targetCoverages

RangedData with 50 rows and 2 value columns across 21 spaces space ranges avgCoverage coverageSD <factor> <IRanges> <numeric> <numeric> chr1 [11158025, 11158264] 44.12917 10.088934 1 2 chr1 [25870174, 25870293] 0.00000 0.000000 3 chr1 [65656333, 65656572] 31.75833 6.931217 4 chr1 [68611504, 686117437 28.79583 13.489341 5 chr1 [70225862, 70226101] 24.16667 8.049360 6 chr1 [112269528, 112270487] 10.14479 3.755132 7 chr1 [160970804, 160970923] 3.074689 16.25833 8 chr1 [182635998, 182636117] 33.85833 5.987797 9 chr1 [186270664, 186270903] 32.48333 7.509529 42 chr6 [152787105, 152787224] 45.0750000 13.6708993 43 chr6 [170639528, 170639647] 31.1916667 9.0834560 chr7 [144345876, 144345995] 44 78.3250000 7.9061326 45 chr8 [1830798, 1830917] 0.2166667 0.4137009 chr9 [6241680, 6241799] 42.5083333 46 5.7203688

47 48 49 50	chr9 chr9 chrX	[7982 [4708	2785 3638	5, 6,	7982 4708	7974 6505]	 	26 14	. 512 . 125 . 591 . 008	0000 6667	7 2	3.8° 2.8°	731: 269:	190 288					
\$coverageAll SimpleRleList of length 24 \$chr1																				
integer-		_	gth :	2487	3716	2 wi	th 3	378	5 rı	ıns										
Lengt		13327		54		85		į			185	5876	3		54		13	30		54
Value	s:	0		1		0			1			()		1			0		1
\$chr10 integer Lengtl	hs: 2	94227	gth	54		116			54			37	7		17 1		1	14		54
value	s:	0		1		0			1			2	2		1			0		1
\$chr11 integer		f leng 33012	gth	1340 54		1 wi 14				ıns	33!	5161	ı		54		8	35		54
Value		0		1		0		`			000)		1		Ì	0		1
	-	-		_		_									_					_
\$chr12																				
integer	-Rle o	f leng	gth	1337	4017	3 wi	th :	1352	2 rı	ıns										
Lengt	hs: 2	74541		54		37		í	54		56	3585	5		54		7	76		54
Value	s:	0		1		0			1			()		1			0		1
\$chr13 integer Lengtl Values	hs: 19		7		0781 54 1	O wi	th : 81 0		3 rı	54				54 1			92			54 1
<pre> <19 more elements></pre>																				
<pre>\$coverageTarget SimpleRleList of length 21 \$chr1</pre>																				
_	-Rle o hs: 1 s: 23	1 1	ĺ 1	2	2	1 2	1	1	1											1 33
	-Rle o hs: 8 s: 6	12 1	l 1	5	1	2 5	6	3												
\$chr12 integer- Lengtl	-Rle o	_	-						4		1	4	3	5	5	3	1	3	2	9

```
Values: 18 20 22 20 21 20 22 21 23 24 25 ... 3 4 5 6 7 8 7 8 10 9
```

\$chr13

integer-Rle of length 720 with 491 runs

Lengths: 1 1 1 2 1 5 2 1 1 1 3 ... 1 1 3 2 2 3 8 1 1 1 1 Values: 32 31 32 33 37 38 37 39 43 41 43 ... 25 23 21 22 25 24 23 22 24 25

\$chr15

integer-Rle of length 240 with 102 runs

Lengths: 9 1 1 9 1 6 2 5 3 3 5 ... 2 2 1 1 2 2 3 1 5 4 Values: 14 13 12 11 10 9 10 9 10 11 12 ... 19 18 17 15 16 17 16 15 16 17

. . .

<16 more elements>

> targets2 <- Coverage\$targetCoverages

Note that coverage should *not* be calculated using the output of function reads2pairs in place of *reads*, since the genomic ranges in that table span the whole region from the first to the second read of a pair. However, for the respective read insert there is no actual coverage, because only the ends of the original fragment were sequenced.

A different measure for the per-target coverage would be to count the numbers of reads overlapping with each target. This can be calculated by the function readsPerTarget. It returns again the input targets table while adding a 'values' column that gives the respective numbers of reads per target. If we provide the table targets2 from above, we will get both the average per-target coverages and the numbers of target-overlapping reads in the same table. In order to speed things up, instead of all reads we could also provide the mappingReads output of function fraction.reads.target, since for counting the reads per target it is enough to look at the reads from which we already know they are on-target. Just make sure that the Offset option was set the same in both fraction.reads.target and readsPerTarget.

> targets2 <- readsPerTarget(reads, targets2)</pre>

> targets2

RangedData with 50 rows and 3 value columns across 21 spaces

	space		ranges		avgCoverage	coverageSD	nReads
	<factor></factor>		<iranges></iranges>	ĺ	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>
1	chr1	[11158025,	11158264]	- 1	44.12917	10.088934	231
2	chr1	[25870174,	25870293]	- 1	0.00000	0.000000	0
3	chr1	[65656333,	65656572]	- 1	31.75833	6.931217	164
4	chr1	[68611504,	68611743]	- 1	28.79583	13.489341	152
5	chr1	[70225862,	70226101]	- 1	24.16667	8.049360	126
6	chr1	[112269528,	112270487]	- 1	10.14479	3.755132	185
7	chr1	[160970804,	160970923]	- 1	16.25833	3.074689	50
8	chr1	[182635998,	182636117]	- 1	33.85833	5.987797	90
9	chr1	[186270664,	186270903]	- 1	32.48333	7.509529	166

```
42
        chr6 [152787105, 152787224]
                                           45.0750000 13.6708993
                                                                        129
        chr6 [170639528, 170639647]
43
                                           31.1916667 9.0834560
                                                                         84
        chr7 [144345876, 144345995]
44
                                           78.3250000
                                                       7.9061326
                                                                        241
45
        chr8 [
               1830798,
                            1830917]
                                            0.2166667
                                                       0.4137009
                                                                          1
46
        chr9 [ 6241680,
                            6241799]
                                           42.5083333 5.7203688
                                                                        124
        chr9 [ 77277336,
                                           72.5125000 18.9239794
47
                          77277575]
                                                                        369
48
        chr9 [ 79827855.
                           798279741
                                           26.1250000
                                                                         84
                                                      3.8731190
49
        chrX [ 47086386,
                           47086505]
                                           14.5916667
                                                       2.8269288
                                                                         46
50
        chrX [150891046, 150891285]
                                           10.0083333
                                                      2.7244310
                                                                         51
```

The resulting *RangedData* table can be converted to a data frame, and as such easily be written to a file, e.g. by

```
> write.table(as.data.frame(targets2), file="target_coverage.txt",
sep="\t", row.names=F, quote=F)
```

Talking about coverage, it is interesting to ask which fraction of target bases reach a coverage of at least k (some value relevant for further analyses, e.g. SNP calling) or which fraction of target bases is covered at all by any read (sensitivity of the capture). The function covered.k calculates such fractions based on the coverageTarget output of coverage.target. Option k specifies the values for which to calculate the fraction of bases achieving the respective coverage.

```
> covered.k(Coverage$coverageTarget, k=c(1, 5, 10))
```

```
1 5 10
0.9520963 0.8444087 0.7033801
```

With coverage.hist we can visualize the coverage distribution, see figure 3. A line is added to the histogram that shows the cumulative fraction of target bases with a coverage of at least the corresponding x-axis value. The line represents the results of covered.k for all possible values of k. Additionally, you can highlight with dashed lines the base fraction achieving a coverage of at least a certain value by defining the option covthreshold.

> coverage.hist(Coverage\$coverageTarget, covthreshold=8)

A similar graph is the coverage uniformity plot, see figure 4. It corresponds more or less to the cumulative line in the coverage histogram. However, coverage.uniformity calculates normalized coverages, i.e. the per-base coverages divided by the average coverage over all target bases. Normalized coverages are not dependent on the absolute quantity of sequenced reads and are hence better comparable between different samples or even different experiments. By default, the x-axis in the figure is truncated at 1, which corresponds to the average normalized coverage. The steeper the curve is falling, the less uniform is the coverage.

> coverage.uniformity(Coverage)

There are more graphical functions concerning read coverage. For example you might be interested in whether large targets are covered by more reads, as

Coverage Distribution

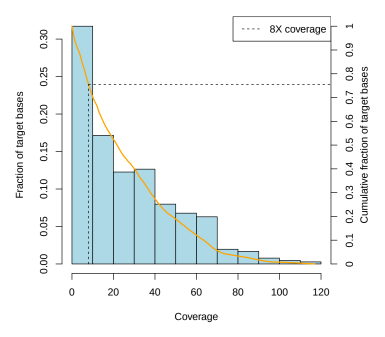


Figure 3: On-target coverage histogram. The orange line shows the cumulative fraction of target bases (right y-axis) with a read coverage of at least x. The dashed lines highlight the fraction of target bases covered by at least 8 reads.

expected, since for larger target regions there should also be more hybridization capture probes ("baits"). Or you might ask whether quite small targets have worse coverage, because the bait tiling might not be as good as for larger targets. Those questions can be addressed by the function coverage.targetlength.plot. As input a RangedData targets table has to be given that contains the relevant information in the 'values' column(s). The graphs are useful for targets of very different lengths. In our example, where all the targets are rather small, the figures are not very informative, see figure 5.

```
> par(mfrow=c(1,2))
> coverage.targetlength.plot(targets2, plotcolumn="nReads", pch=16,
cex=1.5)
> coverage.targetlength.plot(targets2, plotcolumn="avgCoverage", pch=16,
cex=1.5)
```

Another thing to check is the dependency between coverage and GC content of the hybridization capture probes ("baits"). For calculating the GC contents the bait sequences are needed. A file has to be created beforehand that contains the positions as well as the sequences of all baits. The file is loaded by function get.baits which is similar to get.targets, with the difference that overlapping or adjacent baits are not merged, and that a column seqcol has to be

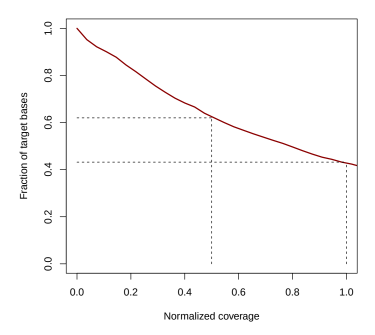


Figure 4: Fraction of targeted bases (y-axis) achieving a normalized coverage of at least x. Dashed lines indicate the fractions of bases achieving at least the average (= 1) or at least half the average coverage (= 0.5).

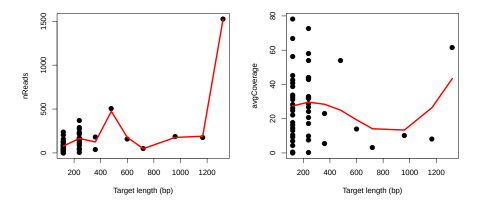


Figure 5: Scatter plots and smoothing splines of number of reads per target (left panel) or average coverage per target (right panel) versus respective target length.

specified that holds the bait sequences. Like get.targets and get.reads, also get.baits by default converts the bait start positions from 0-based to 1-based

coordinates. If the positions given in your baitsfile are already 1-based, set option zerobased to FALSE.

```
> baitsfile <- paste(exptPath, "ExampleSet_Baits.txt", sep="/")
> baits <- get.baits(baitsfile, chrcol=3, startcol=4, endcol=5, seqcol=2)</pre>
```

[1] "read 108 hybridization probes"

With the baits RangedData table and the coverageAll output of coverage.target the (normalized) coverage versus GC content plot can be created. This can take quite some time, since GC content and average (normalized) coverage have to be calculated for every bait. The bait coverages can be returned by setting option returnBaitValues = TRUE. You would expect the added smoothing spline to have an inverse U-shape, with a peak in coverage for baits with GC content around 40-50%. In our small example there are not enough baits with low GC content to encounter the expected shape, see figure 6.

> coverage.GC(Coverage\$coverageAll, baits, pch=16, cex=1.5)

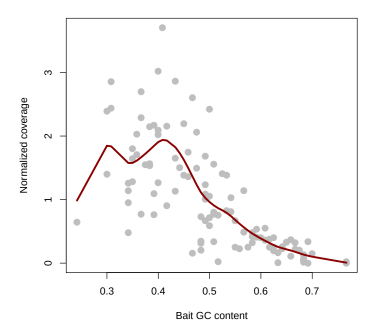


Figure 6: Scatter plot and smoothing spline of normalized average coverage per hybridization probe versus GC content of the respective bait.

There is also a function coverage.plot to visualize per-base coverages along chromosomal positions. The input has to be the coverageAll output of function coverage.target, since also coverages of off-target bases are needed. The

positions of target regions, potentially extended on both sides, can be highlighted as well by specifying options *targets* and *Offset* (see figure 7).

> coverage.plot(Coverage\$coverageAll, targets, Offset=100, chr="chr1", Start=11157524, End=11158764)

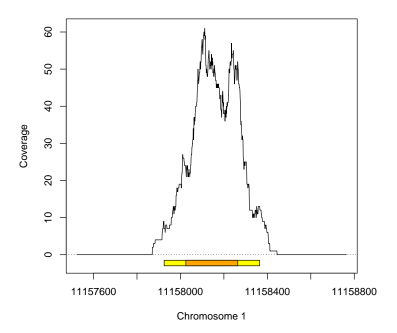


Figure 7: Per-base coverages along chromosomal positions. Target regions (plus addition of 100 bases on both sides) are highlighted in orange (yellow).

Of course, coverages can also be visualized by genome browsers. We provide the function make.wigfiles to create wiggle files that can then be uploaded e.g. to the UCSC genome browser. You can make wiggle files for all chromosomes on which there are reads or just for some selected chromosome(s) by specifying option *chroms*. With option *filename* the name and location where to save the files can be manipulated. Base positions are given as 1-based (i.e. the first base on a chromsome has position 1), as defined for wiggle files by UCSC (http://genome.ucsc.edu/goldenPath/help/wiggle.html).

> make.wigfiles(Coverage\$coverageAll)

6 Read Duplicates

A crucial issue in target capture experiments is read duplication. Usually, read duplicates, i.e. reads that have exact same start and end positions, are removed before follow-up analysis of sequencing data, since they are supposed to be PCR

artifacts. However, here we expect a probably substantial amount of "real" read duplication due to the enrichment process. "Real" read duplicates would be derived from actually separate input DNA molecules that by chance were fragmented at the same position. The duplicates.barplot shows which fraction of reads / read pairs is present in the data in what number of copies. Read multiplicity proportions are calculated and shown separately for on- and off-target reads / read pairs. Therefore, the plot gives an impression about the amount of "real" duplication (expected mostly in the target regions) versus artifactual duplication (expected both on- and off-target). With option returnDups = TRUE the absolute numbers (given on top of the bars, in millions) and percentages (bar heights) can be returned.

> duplicates.barplot(reads, targets)

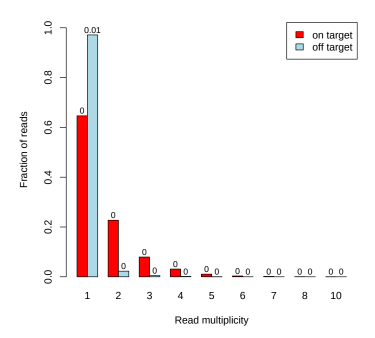


Figure 8: Duplicates barplot treating data as single-end. It shows fractions of onand off-target reads, respectively, that are unique, that are there in two copies, three copies, etc. (x-axis). The numbers on top of the bars are absolute counts in millions.

In figure 8 we see, firstly, that for some reads there are quite a lot of identical copies (x-axis up to 10). Secondly, the percentage of reads with multiple copies is much higher within on-target reads than within off-target reads (red bars are much higher than blue ones for x > 1). This suggest, as mentioned before, that there might be substantial amount of "real" read duplication.

In the case of paired-end data, the position information of both reads of a pair can be used. Reads only have to be considered duplicated if the positions

of both reads are found again in another pair. We can use duplicates.barplot with the table readpairs we created before to make the graph for read pairs instead of single reads. For read pairs the extent of duplication is by far not as high as for single reads, see figure 9.

> duplicates.barplot(readpairs, targets, ylab="Fraction of read pairs")

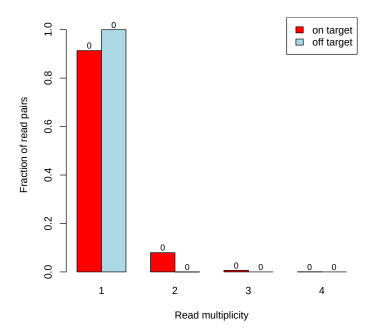


Figure 9: Duplicates barplot for read pairs.

Unfortunately, it is not possible to distinguish the artfactual duplicates from the naturally occurring ones. For that reason it might still be recommendable to remove duplication before further analysis by keeping each duplicated read position only once. (Since here we just deal with genomic positions and not with the actual reads that might differ in sequence and quality, it does not matter which copy of a duplicated read to keep.) You might also decide to do all the QC analysis on the "collapsed" data. Either a bed file is provided that includes only unique read positions, or the tables are collapsed within R, e.g. by the following code.

```
> dupfun <- function(x) duplicated(x$ranges)
> params <- RDApplyParams(rangedData=reads, applyFun=dupfun)
> dups <- unlist(rdapply(params))
> reads.collapsed <- reads[!dups,,drop=T]</pre>
```

In the case of paired-end data, as said before, reads only have to be considered duplicated if the positions of both reads are found again in another

pair. Removing duplicates can be done like follows, selecting non-duplicated reads from the **readpairs** table (and in case of duplication just keep one of the respective reads) and extracting those from the original **read** table.

```
> params2 <- RDApplyParams(rangedData=readpairs, applyFun=dupfun)
> dups2 <- unlist(rdapply(params2))
> ID.nondups <- readpairs$ID[!dups2]
> sel <- reads$ID %in% ID.nondups
> reads.collapsed.pairs <- reads[sel,,drop=T]</pre>
```

In single-end data it is probable to lose a large number of reads by removing duplicates. When treating our example data like single-end sequences, we are left with only 16417 out of originally 19546 reads. Hence, the actual coverage could decrease dramatically.

```
> coverage.target(reads.collapsed, targets, perBase=F, perTarget=F)
$avgTargetCoverage
[1] 17.43535
```

\$targetCoverageSD
[1] 11.7878

```
$targetCoverageQuantiles
0% 25% 50% 75% 100%
0 7 16 27 46
```

Before removing duplicates we calculated a coverage of 27.5.

When using position information of both reads per pair, much less reads are lost, e.g. in the example we can keep 18714 reads. Therefore, the coverage will not decrease that much.

```
> coverage.target(reads.collapsed.pairs, targets, perBase=F, perTarget=F)
$avgTargetCoverage
[1] 25.09604
$targetCoverageSD
[1] 21.22714
```

```
$targetCoverageQuantiles
0% 25% 50% 75% 100%
0 7 20 38 110
```

7 Reproducibility

For any new technology, reproducibility is an important issue. Here we base the reproducibility check on per-target-base coverages. Especially for technical replicates we should yield similar results. But the following graphs might also be useful to ensure homogeneity across biological replicates.

To give an example, we create an artificial new sample by removing randomly 10% of the reads from our data.

```
> r <- sample(nrow(reads), 0.1 * nrow(reads))
> reads2 <- reads[-r,,drop=T]
> Coverage2 <- coverage.target(reads2, targets, perBase=T)</pre>
```

With function coverage.density the coverage densities of several samples can be compared. With option normalized you can choose whether to plot original or normalized coverages. When plotting original values in the example, it is obvious that the second sample does not reach as high coverage for many bases as the first one, see figure 10, right panel. In contrast, normalized coverages are not dependent on the total amount of reads, and we observe a very similar coverage distribution for both samples (left panel), as expected since we just removed some reads randomly.

```
> covlist <- list(Coverage, Coverage2)
> par(mfrow=c(1,2))
> coverage.density(covlist)
> coverage.density(covlist, normalized=F)
```

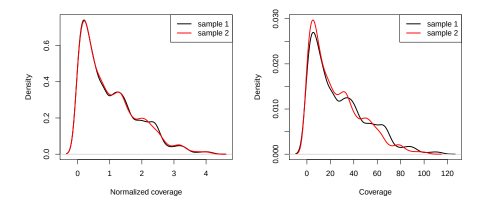


Figure 10: Target coverage densities of two samples, using normalized (left) or original (right) coverages.

Also the coverage uniformity and coverage along chromosome plots shown above (figures 4 and 7) can be produced for several samples within the same graph. The functions can be called repeatedly, while specifying option add=TRUE (see figures 11 and 12). As for the densities, the uniformity is almost identical for the two samples since normalized coverage values are used for this plot.

```
> coverage.uniformity(Coverage, addlines=F)
> coverage.uniformity(Coverage2, addlines=F, add=T, col="blue", lty=2)
> coverage.plot(Coverage$coverageAll, targets, Offset=100, chr="chr1", Start=11157524, End=11158764)
> coverage.plot(Coverage2$coverageAll, add=T, col.line=2, chr="chr1",
```

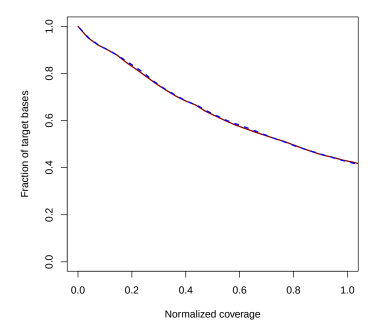


Figure 11: Target coverage uniformity of two samples.

Start=11157524, End=11158764)

With function coverage.correlation we can produce scatterplots between coverage values of pairs of replicate samples. Since a scatterplot for all pertarget-base coverages would be huge (e.g. as pdf graph), and moreover the difference between one million or ten million points in the graphic might not be visible at all, just some fraction of randomly selected values is displayed. The amount of points to plot can be controlled by option plotfrac. By default, 0.1% of all targeted bases are taken into account. In the example we set the fraction to 10%, see figure 13. Scatterplots are shown in the style of a pairs plot. In the lower panels the corresponding Pearson correlation coefficients are shown. Correlation calculations are always based on all coverage values, even if plotfrac < 1 is chosen. Like in coverage.density, by option normalized it can be chosen whether to plot normalized or original coverage values.

> coverage.correlation(covlist, plotfrac=0.1, cex.pch=4)

8 Acknowledgement

The example data used in this manual was taken from an exome sequencing data set of Raquel Rabionet and Xavier Estivill.

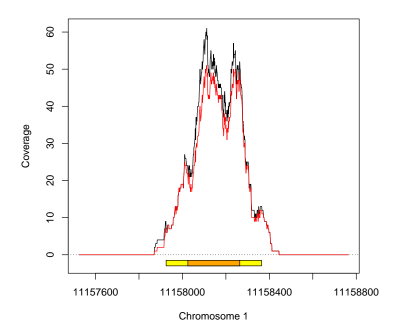


Figure 12: Coverage along chromosome plot for two samples.

9 Session Information

- > toLatex(sessionInfo())
 - R version 2.15.1 (2012-06-22), x86_64-unknown-linux-gnu
 - Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=C, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
 - Base packages: base, datasets, grDevices, graphics, methods, stats, utils
 - Other packages: BiocGenerics 0.4.0, Biostrings 2.26.0, GenomicRanges 1.10.0, IRanges 1.16.0, Rsamtools 1.10.0, TEQC 2.6.0, hwriter 1.3
 - Loaded via a namespace (and not attached): Biobase 2.18.0, bitops 1.0-4.1, parallel 2.15.1, stats4 2.15.1, tools 2.15.1, zlibbioc 1.4.0

10 References

Hummel M, Bonnin S, Lowy E, Roma G. TEQC: an R-package for quality control in target capture experiments. Bioinformatics 2011; doi: 10.1093/bioin-

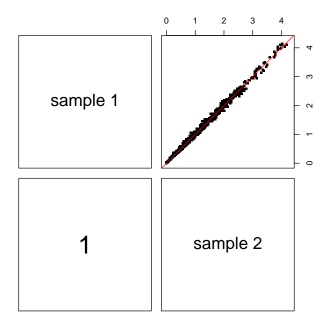


Figure 13: Correlation plots of normalized target coverage values of two samples.

formatics/btr122.

Bainbridge MN, Wang M, Burgess DL, Kovar C, Rodesch MJ, D'Ascenzo M, Kitzman J, Wu Y-Q, Newsham I, Richmond TA, Jeddeloh JA, Muzny D, Albert TJ, Gibbs RA. Whole exome capture in solution with 3 Gbp of data. Genome Biology 2010; 11:R62.

Gnirke A, Melnikov A, Maguire J, Rogov P, LeProust EM, Brockman W, Fennell T, Giannoukos G, Fisher S, Russ C, Gabriel S, Jaffe DB, Lander ES, Nusbaum C. Solution hybrid selection with ultra-long oligonucleotides for massively parallel targeted sequencing. Nat Biotechnol. 2009; 27(2): 182-9.

Tewhey R, Nakano M, Wang X, Pabon-Pena C, Novak B, Giuffre A, Lin E, Happe S, Roberts DN, LeProust EM, Topol EJ, Harismendy O, Frazer KA. Enrichment of sequencing targets from the human genome by solution hybridization. Genome Biol. 2009; 10(10): R116.