



May 08,  
2019

Working

## Protocol for a reproducible circRNA analysis using Docker4Circ

Version 3

Giulio Ferrero<sup>1</sup>, Nicola Licheri<sup>1</sup>, Lucia Coscujuela Tarrero<sup>1</sup>, Carlo De Intinis<sup>1</sup>, Valentina Miano<sup>1</sup>, Raffaele Adolfo Calogero<sup>1</sup>, Francesca Cordero<sup>1</sup>, Marco Beccuti<sup>1</sup>, Michele De Bortoli<sup>1</sup>

<sup>1</sup>University of Turin

[dx.doi.org/10.17504/protocols.io.zrrf556](https://doi.org/10.17504/protocols.io.zrrf556)

Q-Bio Turin



Giulio Ferrero  
University of Turin



### ABSTRACT

Despite many computational tools were developed to predict circular RNAs (circRNAs), a limited number of work-flows exists to fully analyse a circRNA set ensuring the computational reproducibility of the whole analysis.

For this purpose, we designed Docker4Circ, a computational work-flow for a comprehensive circRNAs analysis of a circRNAs composed of four modules: the circRNAs prediction (module 1), the circRNAs classification and annotation (module 2), the circRNAs sequence analysis (module 3), and circRNAs expression analysis (module 4).

To ensure reproducibility each function of Docker4Circ was embedded into a docker image following guideline provided by Reproducible Bioinformatics Project (RBP, <http://reproducible-bioinformatics.org/>). Each function is included in the Docker4Seq R package which already includes different solutions for reproducible bioinformatic analyses.

This protocol describes the use of each function of Docker4Circ to analyse the circRNAs predicted from a set of RNA-Seq experiments performed in normal colon and colorectal cancer cell lines. Furthermore, the description of the functions required to compute the expression level of the analysed circRNAs in a set of RNA-Seq experiments of colorectal primary tumors is provided.

### BEFORE STARTING

Before starting the analysis, the softwares R and Docker must be installed.

To install R follow the information reported in:

<https://www.r-project.org/>

To install docker follow the information reported in:

<https://docs.docker.com/engine/installation/>.

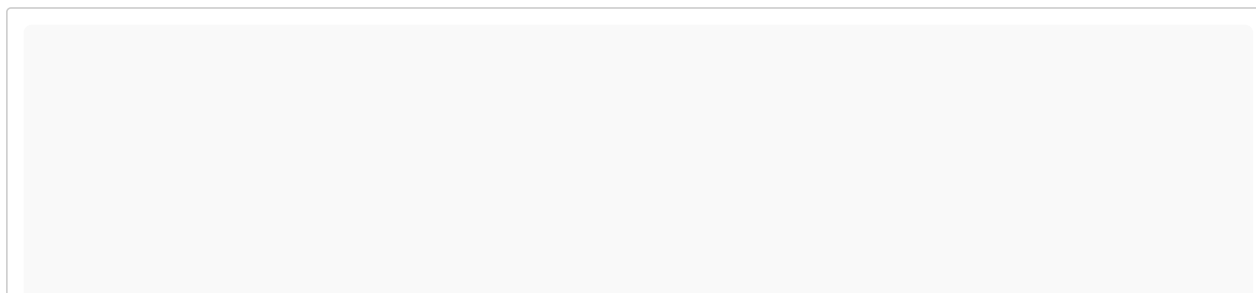
The pipeline is expected to run on 64 bits linux machine with at least 4 cores. 32 Gb RAM are required only if mapping will be done with STAR. A scratch folder should be present, e.g. /data/scratch and it should be writable from everybody.

Each Docker4Seq function has its specific set of parameters that are fully described in the documentation of each function. Access to the documentation using the "?" R operator followed by the name of the function (?ciri).

### Softwares and datasets preparation

#### 1 Installation of the docker4seq R package.

In the R environment digit the following command to install and run the docker4seq pipeline





### # Installation

```
install.packages("devtools")
library("devtools")
install_github("kendomaniac/docker4seq", ref="master")
```

### # Program execution

```
library("docker4seq")
docker4seq R package installation
```



## 2 Creation of the analysis folders.

Before running the analyses, create the following directories in the working folder "**data**" "**reference**" "**scratch**".

The "data" folder will be used to store all the input and output data, the "scratch" folder will be used to store the temporary data, and the "reference" folder will be used to store the reference genome and annotation data.

The folders can be created in your working directory using the command "**mkdir**".



### # Creation of the directories for the analysis

```
system("mkdir data reference scratch")
```

Creation of working folders



Based on the path of your working directory, the path to these folder will be different. In our example the working directory is "**/home**". Then, the path to these three folders will be "**/home/data**" "**/home/scratch**" "**/home/reference**".

You can obtain the path to your working directory using the `R getwd()` command.

## 3 Download of the input RNA-Seq data

The RNA-Seq raw reads for the circRNA prediction can be downloaded from **PRJNA393626**. This dataset includes a triplicate paired-end total and RNAse-R treated RNA-Seq performed on **NCM460** (normal colon cells), **SW480** (primary colorectal cancer cells), and **SW620** cell lines (metastatic colorectal cancer cells).



**PRJNA411984** [↗](#)

The detailed list of datasets needed for the analysis is the following:

SRA ID	ID
<a href="#">SRR5889371</a>	NCM460_rep1
<a href="#">SRR5889366</a>	NCM460_rep2
<a href="#">SRR5889367</a>	NCM460_rep3
<a href="#">SRR5889368</a>	SW480_rep1
<a href="#">SRR5889369</a>	SW480_rep2
<a href="#">SRR5889362</a>	SW480_rep3
<a href="#">SRR5889363</a>	SW620_rep1
<a href="#">SRR5889364</a>	SW620_rep2
<a href="#">SRR5889365</a>	SW620_rep3

The the .fastq files can be downloaded from the ENA page of the experiment (linked below). The links to these files are reported under the column "**FASTQ files (FTP)**".

Each dataset should be located in a specific directory, for example the experiment SRR5889371 should be put in the folder

/data/SRR5889371.

Use the following code to automatically create these folders:

```
# Define a list containing the dataset ids
ids = c("SRR5889362", "SRR5889363", "SRR5889364", "SRR5889365", "SRR5889366", "SRR5889367")

# Define a variable storing the path to data folder
data = paste0(getwd(), "/data")

# Create each dataset folder
for(i in 1:length(ids)){

  dir.create(paste0(data, "/", ids[i]))

}
```

Creation of the dataset folders

To download the files you can use the following R code.

```
# Define a variable storing the fastq url
url1 <- "ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR588/002/SRR5889362/SRR5889362_1.fastq.gz"
url2 <- "ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR588/002/SRR5889362/SRR5889362_2.fastq.gz"

# Call wget to download the dataset into a specific folder
system(paste("wget", url1, "-P", paste0(data, "/SRR5889362")))
system(paste("wget", url2, "-P", paste0(data, "/SRR5889362")))

Dataset download
```

## Module 1a: circRNAs prediction with CIRI2

### 4 Indexing of the reference genome

The first module of Docker4Circ was designed for the prediction of circRNAs from raw RNA-Seq reads. In this part of the protocol, the prediction performed using the **CIRI2** algorithm is described. The first step of the pipeline is the BWA indexing of a reference genome using the **bwaIndex** function.

```
# Define a variable storing the path to the reference folder
ref <- paste0(getwd(), "/reference")

# Define a variable storing the url to the reference genome sequence
genome <- "ftp://ftp.ensembl.org/pub/grch37/update/fasta/homo_sapiens/dna/Homo_sapiens.GRCh37.dna.toplevel.fa"

# BWA indexing of the genomic sequence
bwaIndex(group="docker", genome.folder=ref, genome.url=genome, mode="General")

Human genome BWA indexing
```

All the index files of the reference genome will be stored in the reference folder.

### 5 Read alignment with BWA

The second step of the pipeline will be the RNA-Seq read quality control using the *fastqc* function, the read alignment using the *bwa* function, and the circRNAs prediction using *ciri2* function. The sequential call of these three functions is implemented in the *wrapperCiri* function that can be used as follow:

```
# Define a variable storing the path to the scratch folder
scratch = paste0(getwd(), "/scratch")

# Define a variable storing the path to the reference genome sequence
gs <- paste0(ref, "/genome.fa")

# Download the reference gene annotations
url_gene <- "ftp://ftp.ensembl.org/pub/grch37/release-87/gtf/homo_sapiens/Homo_sapiens.GRCh37"

system(paste("wget", url_gene, "-F", ref))

# Define a variable storing the path to the reference gene annotations
gtf <- paste0(ref, "/gencode.v28lift37.annotation.gtf.gz")

# Repeat the BWA alignment for each dataset
for (i in 1:length(ids)){

  fqfolder <- paste(data, ids[i], sep="/")

  wrapperCiri(group = "docker", scratch.folder=scratch, data.folder=fqfolder,
    genome.file=gs, seq.type="pe", sample.id=ids[i], threads=1, annotation.file=gtf,
    max.span=200000, stringency.value="high", quality.threshold=10)

}
```

BWA RNA-Seq read alignment and circRNAs prediction with CIRI2

The read alignment quality control files, the BWA read alignment files and the set of predicted circRNAs (.ciri files) will be stored in the folder in which the input fastq files are located.

## 6 Merge of multiple CIRI2 predictions

To merge multiple CIRI2 circRNAs predictions into a common list of circRNAs, the function *ciri2MergePredictions* was implemented. Each .ciri files located in a specific dataset folder will be merged into a single file. Each sample name should be reported as element of the *samples.list* vector with corresponding sample type specified as element of the *covariate.list* vector.

```
# Copy each CIRI2 prediction in the data folder
system(paste0("cp ", data, "/*/*.ciri ", data))

# Define a list of sample groups
cov <- c("NCM460","NCM460","NCM460","SW480","SW480","SW480","SW620","SW620","SW620")
ids <- c("SRR5889366","SRR5889367","SRR5889371","SRR5889362","SRR5889368","SRR5889369")
cov_ord <- c("NCM460", "SW480", "SW620")

# Merge the CIRI2 outputs
ciri2MergePredictions(group="docker", scratch.folder=scratch, data.folder=data,
  samples.list=ids, covariates.list=cov, covariate.order=cov_ord, min_reads=2, min_reps=2, min_av=2)
```

Merge of different CIRI2 predictions

This function generates two files named *merged\_circs.crna* and *merged\_circs.crna\_count*. The first file reports the ids and coordinates of each circRNA while the second file is a matrix reporting the number of backsplicing reads supporting each circRNA in each

dataset.

## Module 1b: circRNAs prediction with STARChip

### 7 Indexing of the reference genome

In this part of the protocol, the prediction performed using the **STARChip** algorithm is described. The first step of the pipeline is the STAR indexing of a reference genome using the *rsemstarIndex* function.



```
# Call the function to index the reference genome
rsemstarIndex(group="docker", genome.folder=ref,
ensembl.urlgenome=genome, ensembl.urlgtf=url_gene, threads=40)
```

Reference genome indexing with STAR



The STAR index files of the reference genome will be stored in the **reference** folder.

### 8 Create the reference file for STARChip

The STARChip algorithm required an additional .bed file defined starting from the STAR-indexed reference genome. To generate this file, the function *starChipIndex* should be applied as follow:



```
# Creation of the STARChip reference file
starChipIndex(group="docker", genome.folder=ref)
```

Reference creation for STARChip



Two files called **genome.gtf.bed** and **genome.gtf.exon.bed** will be generated in the **reference** folder.

### 9 Running of STAR chimeric analysis

The STARChip circRNAs prediction require a run of STAR chimeric algorithm on the RNA-Seq reads. The STAR chimeric analysis is implemented in the *starChimeric* function that should be applied as follow:



```
# Run of the STAR chimeric analysis on each RNA-Seq dataset
for(i in 1:length(ids)){

  fqfolder = paste(data, ids[i] , sep="/")

  starChimeric(group="docker", fastq.folder=fqfolder, scratch.folder=scratch, genome.folder=ref,

}

STAR chimeric analysis
```



This function will generate each STAR chimeric output in each sample directory.

### 10 STARChip circRNAs prediction

The circRNA prediction will be performed using the *starchipCircle* function applied on the **data** folder containing each sample-specific folder



```
# Prediction of circRNAs using STARChip
starchipCircle_temp(group="docker", genome.folder=ref, scratch.folder=scratch,
samples.folder=data, reads.cutoff=1, min.subject.limit=2, threads=40,
do.splice=TRUE, cpm.cutoff=0, subjectCPM.cutoff=0, annotation=TRUE)
```

STARChip circRNAs prediction



## Module 2: circRNAs annotation and classification

### 11 Preparation of the reference files for the classification

To classify the circRNAs a reference of Ensembl exon and transcript annotations is required. To provide these data in the correct format, the *circrnaPrepareFiles* function should be applied as follow.



```
# Create the reference exon and gene isoforms files required for the circRNA classification analysis
circrnaPrepareFiles(group="docker", scratch.folder=scratch, data.folder=data,
assembly="hg19")
```

Preparation of the reference files for the classification



Two files called **exons\_reference** and **isoforms\_reference** reporting the exons and the gene isoform annotations respectively, will be generated in the data folder.

### 12 Classification of circRNAs based on their mapping location

To classify the circRNAs into distinct classes the function *circrnaClassification* should be applied as follow.



```
# Classification of a circRNA set
circrnaClassification(group="docker", scratch.folder=scratch,
circrna.data=paste0(data,"/merged_circs.crna"), exon.data=paste0(data,"/exons_reference",
isoform.data=paste0(data,"/isoforms_reference"), assembly="hg19"))
```

circRNAs classification



Two files called **circRNA\_classification** and **circRNA\_univocal\_classification** reporting the transcript- and the gene-level classification respectively, will be generated in the data folder.

### 13 Annotation of the circRNA based on databases information

To annotate the circRNAs based on the information stored in the circBase and TSCD database, the function *circAnnotation* should be applied as follow.



```
# Annotation of a circRNA set based on information from public databases
circAnnotations(group="docker", scratch.folder=scratch,
ciri.file=paste0(data,"/merged_circs.crna"), genome.version="hg19")
```

circRNAs annotation



Different files with suffix **.anno** will be generated in the data folder. Each of these files will report the information about the circRNAs found in a specific database (es. circbase.anno will report the circBase information)

## Module 3: circRNAs sequence analysis

### 14 Reconstruction of the circRNAs backsplicing junction sequences

To reconstruct the sequence of the backsplicing junction of each circRNA the function *circrnaBSJunctions* should be applied as follow



```
# Reconstruction of circRNA backsplicing junction sequence
circrnaBSJunctions(group="docker", scratch.folder=scratch,
circrna.data=paste0(data,"/merged_circs.crna"), exon.data=paste0(data,"/exons_reference"),
assembly="hg19")
```

Backsplicing sequence reconstruction



This function will generate a files called **circRNA\_backsplicing\_sequences.fasta** reporting the reconstructed sequence.

## 15 Prediction of alternative splicing events involving the circRNAs

To predict the set of alternative splicing events involving the circRNA, the **ciriAS** function can be applied as follow using each BWA read alignment file and CIRI2 output.

```
# Application of CIRI AS algorithm on each dataset
for(i in 1:length(ids)){

  fqfolder = paste(path, ids[i] , sep="")

  ciriAS(group="docker", scratch.folder=scratch, sam.file=paste0(fqfolder,"/aligned_reads.sam"),
  ciri.file=paste0(fqfolder, "/", ids[i], ".ciri"), genome.file=gs,
  annotation.file=gtf)

}
```

Detection of alternative splicing events involving the circRNAs

The output of the function will be a two different files called **structure.list** and **structure\_AS.list**.

### Module 4: circRNAs expression analysis

## 16 Download of the RNA-Seq data in which quantify the expression of a circRNA set

The RNA-Seq raw reads for the circRNAs quantification can be downloaded from **PRJNA-411984**. This dataset includes RNA-Seq data of three CRC samples with matched data from normal mucosa samples.

 **PRJNA411984** [↗](#)

The detailed list of datasets needed for the analysis is the following:

SRA ID	ID
<a href="#">SRR6067723</a>	CRC_1
<a href="#">SRR6067725</a>	CRC_2
<a href="#">SRR6067727</a>	CRC_3
<a href="#">SRR6067729</a>	Normal mucosa 1
<a href="#">SRR6067731</a>	Normal mucosa 2
<a href="#">SRR6067733</a>	Normal mucosa 3

For each dataset a specific folder in the data folder

The datasets can be downloaded using the following code applied using the url of the different datasets.

```
# Define a variable storing the fastq url
url1 <- "ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR606/003/SRR6067723/SRR6067723_1.fastq.gz"
url2 <- "ftp://ftp.sra.ebi.ac.uk/vol1/fastq/SRR606/003/SRR6067723/SRR6067723_2.fastq.gz"

# Call wget to download the dataset into a specific folder
system(paste("wget", url1, "-P", paste0(data, "/SRR6067723")))
system(paste("wget", url2, "-P", paste0(data, "/SRR6067723")))

Download dataset
```

## 17

The quantification of each circRNA in a specific RNA-Seq dataset is computed using the ***circrnaQuantification*** function

```
# Define a list containing the dataset ids
ids2 <- c("SRR6067723", "SRR6067725", "SRR6067727", "SRR6067729", "SRR6067731", "SRR6067733")

# Apply the quantification function on the fastq file stored in each sample folder
for(i in 1:length(ids2)){

    fqfolder = paste0(data, "/", ids2[i] , sep="")

    setwd(fqfolder)

    # For paired-end reads, the two fastq files must be joined into a single fastq file
    system(paste0("cat ", ids2[i], "_1.fastq ", ids2[i], "_2.fastq > ", ids2[i], ".fastq"))

    circrnaQuantification(group="docker", scratch.folder=scratch, rnaseq.data=paste0(fqfolder, "/", ids2[i], ".fastq"),
        backsplicing_junctions.data=paste0(data, "/circRNA_backsplicing_sequences.fasta"),
        hc.params=c(21, 40, 1000000, 1000000, 17, 30))

}
```

CircRNAs expression analysis

This function will generate a file with extension **.hashcirc** reporting for each dataset the number of reads supporting each circRNA.

## 18

Before the differential expression analysis of the circRNAs level detected by the `circrnaQuantification` function, the function ***mergeData*** should be applied in order to merge the result obtained for each dataset.

```
# Define a variable storing the list of sample classes
cov <- c("T", "T", "T", "N", "N", "N")
cov.order <- c("T", "N")

# Merge different file with extension hashcirc in the subfolders of data
mergeData(group="docker", scratch.folder=scratch, samples.ids=ids2, covariates=cov, covariate
extension="hashcirc", column_index=2)
```

Merge different data

The output of this function called **MergedData.tsv** is a table reporting the counts of each annotation for each dataset. Furthermore, the sample identifiers reported in the header will be report the sample id followed by the corresponding sample class. On this table the differential expression analysis can be performed using DESeq2 algorithm implemented in the *wrapperDeseq2* function.

```
# Differential expression analysis with DESeq2
wrapperDeseq2(group="docker", output.folder=data,
experiment.table=paste0(data, "/MergedData.tsv"), log2fc=0, fdr=0.05, ref.covar="N",
type="gene", batch=FALSE)
```

Differential expression analysis



