# Using PACViR

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#### Contents

1	Introduction	1
2	Requirements	1
3	Executing PACViR via R interpreter	2
4	Executing PACViR via BASH shell	3
5	sessionInfo	4

### 1 Introduction

PACViR is a user-friendly software tool to visualize the coverage depth of a complete plastid genome as well as the equality of its IR regions while simultaneously accounting for the circular, quadripartite structure of the genome. This vignette provides two examples of executing PACViR on empirical data via the R function PACViR.complete() from within the R interpreter and from the BASH commandline shell.

## 2 Requirements

To run PACViR, several dependencies have to be installed.

```
# Mosdepth (a general dependency)
system("conda install -y mosdepth")
# CRAN packages
if (!require("pacman"))
    install.packages("pacman")
pacman::p_load("RCircos", "optparse", install=TRUE)
# Bioconductor packages
if (!requireNamespace("BiocManager", quietly=TRUE))
    install.packages("BiocManager")
if (!requireNamespace("genbankr", quietly=TRUE))
    BiocManager::install("genbankr")
```

## 3 Executing PACViR via R interpreter

PACViR can be executed via the R function PACViR.complete() from within the R interpreter.

#### Nuphar japonica isolate NY692 chloroplast, complete genome.

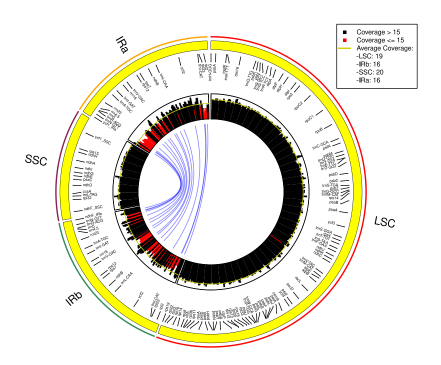


Figure 1: File MH161174\_AssemblyCoverage\_viz.svg as generated via PACViR.complete()

## 4 Executing PACViR via BASH shell

PACViR can be executed via the BASH shell command PACViR\_Rscript.R from the BASH commandline shell.

```
# MH899017
Rscript ./inst/extdata/PACViR_Rscript.R \
   -k ./inst/extdata/MH899017.gb \
   -b ./inst/extdata/MH899017_PlastomeReadsOnly.sorted.bam \
   -o ./MH899017_AssemblyCoverage_viz.svg
```

### Dasyphyllum excelsum isolate DAS01 chloroplast, complete genome.

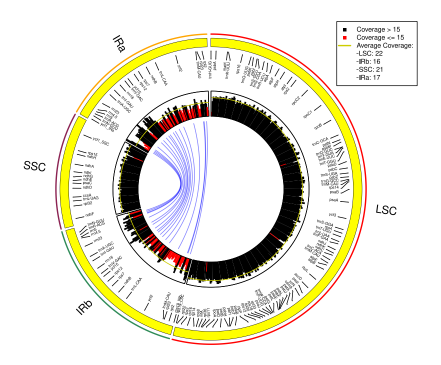


Figure 2: File MH899017\_AssemblyCoverage\_viz.svg as generated via PACViR\_Rscript.R

## 5 sessionInfo

> sessionInfo()

R version 3.3.3 (2017-03-06)

Platform: x86\_64-pc-linux-gnu (64-bit)
Running under: Debian GNU/Linux 9 (stretch)

#### locale:

[1] LC\_CTYPE=de\_DE.UTF-8 LC\_NUMERIC=C

[3] LC\_TIME=de\_DE.UTF-8 LC\_COLLATE=de\_DE.UTF-8
[5] LC\_MONETARY=de\_DE.UTF-8 LC\_MESSAGES=de\_DE.UTF-8

[7] LC\_PAPER=de\_DE.UTF-8 LC\_NAME=C
[9] LC\_ADDRESS=C LC\_TELEPHONE=C
[11] LC\_MEASUREMENT=de\_DE.UTF-8 LC\_IDENTIFICATION=C

#### attached base packages:

[1] stats graphics grDevices utils datasets

[6] methods base

loaded via a namespace (and not attached):

[1] tools\_3.3.3