# LOG & EXTENDED README

## Henrik Christiansen

June 30, 2021

# Project: radseq pilot

RADseq optimization pilot experiment with several Antarctic species.

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#### Log:

30/06/2021: updated readme

29/06/2021: added coverage calculation in script 05\_marker\_density.R

24/06/2021: changed and rerun empirical digestion plots, set alpha to 1 and changed plotting order to avoid

overlays

23/06/2021: added digestion with a standard 1000 mb genome

10-14/06/2021: added empirical loci statistics and updated marker density calculations

05-06/2021: added new functions and code to run marker density calculations, new script: 05\_marker\_density.R

24/04/2021: added extensive documentation

29/03/2021: rerun 01\_digests.R again

24/03/2021: updated and rerun 01 digests.R for some additional size windows

22/03/2021: updated amphipod species ID

01/03/2021: updated bird plots in 02\_empirical\_digests.R

 $23/02/2021: \quad \text{updated double digest calculations in } \text{recto} \underline{\text{REs}}\underline{\text{and}}\underline{\text{functions.R}} \quad \text{and} \quad 02\underline{\text{-empiri-like}}$ 

 $cal\_digests.R$ 

22/02/2021: updated and rerun 02 empirical digests.R, added input files for coverage calculations

21/02/2021: fixed some git issue with large .RDataTmp file

18/02/2021: updated and rerun 02\_empirical\_digests.R

17/02/2021: updated and consolidated code and files locally and on github, downdgraded bioanalyzer to

v0.5.1 again, the other version throws an error

22/01/2021: fixed xml encoding issues

20/01/2021: added new xml file for run 7, updated bioanalyzer to v0.6.2 and updated 02\_empiri-

cal\_digests.R

20/01/2021: added the project to a github repository

16/12/2020: added a modified read.bioanalyzer function to read in xml file with a missing lower marker in one sample

14/12/2020: small, mostly cosmetic updates in most scripts; updated project and input/output description; finalized 02\_empirical\_digests.R

11/12/2020: added customized plot saving function in recto\_REs\_and\_functions.R

10/12/2020: updated 02\_empirical\_digests.R, added a customized ggplot plotting function in recto\_REs\_and\_functions.R added in silico functions for comparison in recto\_REs\_and\_functions.R

12/11/2020: updated 02\_empirical\_digests.R & added a customized plotting function in recto\_REs\_and\_functions.R

11/11/2020: updated to run with R v4.0.3

07-10/2020: added script 02\_empirical\_digests.R to read results from bioanalyzer and analyze them in R, instead of cumbersome "manual" analysis in spreadsheets

20/07/2019: updated plot ratio scripts

10/07/2019: updated digest scripts

09/07/2019: updated the plots for test library 2

02/07/2019: updated digest scripts

28/06/2019: finalized coverage plot

27/06/2019: re-orderd scripts, included scripts for in silico digestion (previously stored in different R project)

23-27/06/2019: updated parameter plots (now based on correct de novo runs, with PE concatenating); included & updated coverage plot

16-22/06/2019: updated parameter plots; started script to plot targeted and realized coverage

01-14/06/2019: included scripts to plot n\_loci and n\_snps\_per\_locus from de novo parameter optimization series

04/12/2018: rerun due to updated input data (Trematomus mainly)

29/11/2018: created script 01\_plotting.R and ratio plots

28/11/2018: created R project & input data

#### Input:

Various reference genomes from target species, or related species in fasta format under:

• ../refgenomes (note: these are not included in the online version on github and zenodo as they are too big; if you want to replicate these analyses you need to download the refgenomes from genbank)

Output files from the Agilent Bioanalyzer Software in XML format under:

• data/bioanalyzer\_results

CSV file with all metadata related to the Bioanalyzer runs, created manually:

• data/bioanalyzer\_results/run\_overview.csv

CSV/TSV files in data/test\_libraries with data from 5 test libraries:

- coverage\_stats.csv: number of loci and coverage for each library under different M values as reported in gstacks.log.distribs files (note: the original gstacks.log.distribgs files are included as well, but actually not needed as long as you have this summary csv file)
- density\_stats.csv: additional statistics for each library as reported in populations.log files (note: the
  original populations.log files are included as well, but actually not needed as long as you have this
  summary csv file)
- $\bullet$  various n\_snps\_per\_locus.tsv: files for each library/species and different r/M values with information about the loci and SNPs

R script that contains restriction enzyme information and various functions for analysis and plotting as used in the analyses scripts:

• scripts/recto REs and functions.R

Additional R script to export plots consistently:

• scripts/printfig.R

# Analyses:

To be run sequentially, all listed under /scripts.

- $\bullet~01\_digests.R:$  script to perform in silico digestions for different target organisms
- 02\_empirical\_digests.R: script to read bioanalyzer results and plot output in comparison with in silico results
- 03\_plot\_n\_loci.R: script based on Rochette & Catchen 2017 to plot the number of (polymorphic) loci for different M parameters
- 04\_plot\_n\_snps\_per\_locus.R: script based on Rochette & Catchen 2017 to plot the number of SNPs per locus for different M parameters
- 05 marker density.R: script to calculate how many snps are sequenced across the genome
- 06 plot coverage.R: script to plot target and realized coverage

## **Output:**

In /data/in\_silico\_results:

various tables.csv containing in silico digest results per target taxa

In /data:

• marker\_density\_results.csv containing the output from the 05\_marker\_density.R script

In /figures:

- genome digestion curves, empirical and in silico, created with 02 empirical digests.R
- various n\_loci\_Mn plots per library, created with 03\_plot\_n\_loci.R
- various n\_snps per library/species, created with 04\_n\_snps\_per\_locus.R
- coverage plots, created by 06\_plot\_coverage.R

## Session info:

Package citations:

```
## Warning in FUN(X[[i]], ...): no date field in DESCRIPTION file of package
## 'bioanalyzeR'
## [[1]]
##
## To cite package 'here' in publications use:
##
     Kirill Müller (2020). here: A Simpler Way to Find Your Files. R
     package version 1.0.1. https://CRAN.R-project.org/package=here
##
##
## A BibTeX entry for LaTeX users is
##
##
     @Manual{,
##
       title = {here: A Simpler Way to Find Your Files},
##
       author = {Kirill Müller},
##
       year = \{2020\},\
       note = {R package version 1.0.1},
##
##
       url = {https://CRAN.R-project.org/package=here},
##
##
##
## [[2]]
##
## To cite package 'SimRAD' in publications use:
##
##
     Olivier Lepais and Jason Weir (2016). SimRAD: Simulations to Predict
##
     the Number of RAD and GBS Loci. R package version 0.96.
     https://CRAN.R-project.org/package=SimRAD
##
##
## A BibTeX entry for LaTeX users is
##
##
     @Manual{,
       title = {SimRAD: Simulations to Predict the Number of RAD and GBS Loci},
##
       author = {Olivier Lepais and Jason Weir},
##
       year = \{2016\},\
##
##
       note = {R package version 0.96},
##
       url = {https://CRAN.R-project.org/package=SimRAD},
##
     }
##
##
## [[3]]
## To cite seginr in publications use:
##
##
     Charif, D. and Lobry, J.R. (2007)
##
## A BibTeX entry for LaTeX users is
##
##
     @InCollection{,
       author = {D. Charif and J.R. Lobry},
##
```

```
##
       title = {Seqin{R} 1.0-2: a contributed package to the {R} project for statistical computing devo
##
       booktitle = {Structural approaches to sequence evolution: Molecules, networks, populations},
##
       year = \{2007\},\
       editor = {U. Bastolla and M. Porto and H.E. Roman and M. Vendruscolo},
##
##
       series = {Biological and Medical Physics, Biomedical Engineering},
##
       pages = \{207-232\},
       address = {New York},
##
       publisher = {Springer Verlag},
##
##
       note = \{\{ISBN :\} 978-3-540-35305-8\},
##
##
##
## [[4]]
##
## To cite package 'bioanalyzeR' in publications use:
##
##
     Joseph Foley (2020). bioanalyzeR: Analysis of Agilent electrophoresis
##
     data. R package version 0.5.1.
##
## A BibTeX entry for LaTeX users is
##
##
     @Manual{,
       title = {bioanalyzeR: Analysis of Agilent electrophoresis data},
##
       author = {Joseph Foley},
##
##
       year = \{2020\},\
##
       note = {R package version 0.5.1},
##
##
##
## [[5]]
##
##
     Wickham et al., (2019). Welcome to the tidyverse. Journal of Open
##
     Source Software, 4(43), 1686, https://doi.org/10.21105/joss.01686
##
## A BibTeX entry for LaTeX users is
##
##
     @Article{,
##
       title = {Welcome to the {tidyverse}},
       author = {Hadley Wickham and Mara Averick and Jennifer Bryan and Winston Chang and Lucy D'Agosti:
##
##
       year = {2019},
       journal = {Journal of Open Source Software},
       volume = \{4\},
##
       number = \{43\},
##
##
       pages = \{1686\},
       doi = {10.21105/joss.01686},
##
##
##
##
## [[6]]
## To cite package 'ggsci' in publications use:
##
##
     Nan Xiao (2018). ggsci: Scientific Journal and Sci-Fi Themed Color
    Palettes for 'ggplot2'. R package version 2.9.
```

```
##
     https://CRAN.R-project.org/package=ggsci
##
## A BibTeX entry for LaTeX users is
##
##
     @Manual{,
       title = {ggsci: Scientific Journal and Sci-Fi Themed Color Palettes for
##
   'ggplot2'},
       author = {Nan Xiao},
##
##
       year = {2018},
##
       note = {R package version 2.9},
##
       url = {https://CRAN.R-project.org/package=ggsci},
##
##
##
## [[7]]
##
## To cite package 'gridExtra' in publications use:
##
##
     Baptiste Auguie (2017). gridExtra: Miscellaneous Functions for "Grid"
##
     Graphics. R package version 2.3.
##
    https://CRAN.R-project.org/package=gridExtra
## A BibTeX entry for LaTeX users is
##
##
     @Manual{,
##
       title = {gridExtra: Miscellaneous Functions for "Grid" Graphics},
##
       author = {Baptiste Auguie},
       year = {2017},
##
##
       note = {R package version 2.3},
       url = {https://CRAN.R-project.org/package=gridExtra},
##
     }
##
R Session:
## R version 4.1.0 (2021-05-18)
## Platform: x86 64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 18.04.5 LTS
## Matrix products: default
           /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.7.1
## LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.7.1
##
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                   LC_NUMERIC=C
## [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=en_US.UTF-8
## [5] LC_MONETARY=en_US.UTF-8
                                   LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
## [9] LC_ADDRESS=C
                                   LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
```

```
## other attached packages:
  [1] gridExtra_2.3
                                     ggsci_2.9
  [3] forcats 0.5.1
                                     stringr 1.4.0
                                     purrr_0.3.4
## [5] dplyr_1.0.6
   [7] readr_1.4.0
                                     tidyr_1.1.3
## [9] tibble 3.1.2
                                     ggplot2 3.3.3
## [11] tidyverse 1.3.1
                                     bioanalyzeR 0.5.1
## [13] seginr 4.2-5
                                     SimRAD 0.96
## [15] zlibbioc 1.38.0
                                     ShortRead 1.50.0
## [17] GenomicAlignments_1.28.0
                                     SummarizedExperiment_1.22.0
                                     MatrixGenerics_1.4.0
## [19] Biobase_2.52.0
## [21] matrixStats_0.59.0
                                     Rsamtools_2.8.0
## [23] GenomicRanges_1.44.0
                                     BiocParallel_1.26.0
## [25] Biostrings_2.60.0
                                     GenomeInfoDb_1.28.0
## [27] XVector_0.32.0
                                     IRanges_2.26.0
## [29] S4Vectors_0.30.0
                                     BiocGenerics_0.38.0
## [31] here_1.0.1
##
## loaded via a namespace (and not attached):
  [1] bitops 1.0-7
                                fs 1.5.0
                                                       lubridate 1.7.10
##
   [4] RColorBrewer_1.1-2
                               progress_1.2.2
                                                       httr_1.4.2
## [7] rprojroot_2.0.2
                                tools 4.1.0
                                                       backports_1.2.1
## [10] utf8_1.2.1
                               R6_2.5.0
                                                       DBI_1.1.1
## [13] colorspace 2.0-1
                                ade4 1.7-16
                                                       withr 2.4.2
                                                       compiler_4.1.0
## [16] tidyselect_1.1.1
                                prettyunits_1.1.1
## [19] cli 2.5.0
                               rvest_1.0.0
                                                       xml2 1.3.2
## [22] DelayedArray_0.18.0
                                scales_1.1.1
                                                       digest_0.6.27
## [25] rmarkdown_2.8
                                base64enc_0.1-3
                                                       jpeg_0.1-8.1
## [28] pkgconfig_2.0.3
                               htmltools_0.5.1.1
                                                       dbplyr_2.1.1
## [31] rlang_0.4.11
                               readxl_1.3.1
                                                       rstudioapi_0.13
## [34] generics_0.1.0
                                hwriter_1.3.2
                                                       jsonlite_1.7.2
## [37] RCurl_1.98-1.3
                               magrittr_2.0.1
                                                       GenomeInfoDbData_1.2.6
## [40] Matrix_1.3-4
                                Rcpp_1.0.6
                                                       munsell_0.5.0
## [43] fansi_0.5.0
                                lifecycle_1.0.0
                                                       stringi_1.6.2
## [46] vaml 2.2.1
                               MASS 7.3-54
                                                       grid 4.1.0
## [49] crayon_1.4.1
                               lattice_0.20-44
                                                       haven_2.4.1
## [52] hms 1.1.0
                               knitr 1.33
                                                       pillar 1.6.1
## [55] reprex_2.0.0
                               XML_3.99-0.6
                                                       glue_1.4.2
## [58] evaluate 0.14
                                latticeExtra_0.6-29
                                                       modelr_0.1.8
## [61] png_0.1-7
                                vctrs_0.3.8
                                                       cellranger_1.1.0
## [64] gtable 0.3.0
                                assertthat 0.2.1
                                                       xfun 0.23
## [67] broom_0.7.6
                                ellipsis_0.3.2
```