Insights Into Computational Reproducibility in Plant Pathology and a Way Forward

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Open research practices have been highlighted extensively during the last ten years in many fields of study as essential standards needed to promote transparency and reproducibility of scientific results. Scientific claims can only be evaluated based on how protocols, materials, equipment and methods were described; data were collected and prepared, and; analyses conducted. Openly sharing protocols, data and computational code is central for current scholarly dissemination and communication, but in many fields, including plant pathology, adoption of these practices has been slow. We randomly selected 300 articles published from 2012 to 2018 across 21 journals representative of the pathology discipline and assigned them scores reflecting their openness and reproducibility. We found that most of the articles were not very open, failing to share data or code in a meaningful way. We propose that using open source tools for producing open and reproducible work and analysis is advantageous, benefiting not just readers, but the authors as well and provide ideas and tools to promote open, reproducible research practices among plant pathologists.

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# Introduction

Modern plant pathological research has many facets given the array of disciplines and sub-disciplines currently involved. Collectively, they contribute to increasing our basic and applied knowledge of several aspects of pathogen biology and disease development to ultimately improve plant disease management. Scientific research in the field varies from the purely observational or descriptive nature to inferential, based on experimental or simulation-derived data sets. Whatever the case, research findings are verifiable based on how much of the research materials, processes and outcomes are made available beyond what is reported in the scientific article and the ability of others to make use of the methods and results. These research findings include biological materials (isolates or strains), nucleic/protein sequences, experimental and simulated raw data annotations, drawings and photographs and statistical analysis code among other materials and data generated as a course of conducting research.

Recently, open science has been highlighted with many donors expecting data to be available (Government of Canada 2016; Noorden 2017; Australian Research Council 2018), journals in the field promoting the sharing of data (Tropical Plant Pathology), and other scientists interested in sharing and collaborating more widely (Wald 2010).

While open science and reproducibility are not the same, they are linked. By proactively practising open science, the work becomes more reproducible through the availability of data and code. That is, open science leads to reproducibility and replicability.

## Definitions

For us to easily discuss the topic, we first must define what we mean so that we may clearly communicate. Many of the terms used in this area have varying definitions that may or may not agree with each other. For instance, reproducible research is recently highlighted by many authors (Fidler and Gordon 2013; Stodden et al. 2013; Fitzjohn et al. 2014; Ioannidis 2014; Brunsdon 2015; Sweedler 2015; Baker 2016; Iqbal et al. 2016; Patil et al. 2016; Weissgerber et al. 2016; Editors at Nature 2016) as an important issue. The International Union of Pure and Applied Chemistry provides definitions of repeatability in the “Compendium of Chemical Terminology – Gold Book” as:

“The closeness of agreement between independent results obtained with the same method on identical test material, under the same conditions (same operator, same apparatus, same laboratory and after short intervals of time). The measure of repeatability is the standard deviation qualified with the term: `repeatability’ as repeatability standard deviation. In some contexts repeatability may be defined as the value below which the absolute difference between two single test results obtained under the above conditions, may be expected to lie with a specified probability,” (McNaught and Wilkinson 1997).

and reproducibility is defined as:

“The closeness of agreement between independent results obtained with the same method on identical test material but under different conditions (different operators, different apparatus, different laboratories and/or after different intervals of time). The measure of reproducibility is the standard deviation qualified with the term ‘reproducibility’ as reproducibility standard deviation. In some contexts reproducibility may be defined as the value below which the absolute difference between two single test results on identical material obtained under the above conditions, may be expected to lie with a specified probability. Note that a complete statement of reproducibility requires specification of the experimental conditions which differ,” (McNaught and Wilkinson 1997).

However, while these definitions are clear and well accepted in physical sciences, in biological sciences it is not always possible to use identical test material or perhaps the the time or resources are not available for full reproduction, *e.g.*, field trials that span years and locations or complex glasshouse experiments. Therefore we will follow Peng’s (2009) definition that provides clear guidelines a minimum standard of “reproducible research”:

“The replication of scientific findings using independent investigators, methods, data, equipment, and protocols has long been, and will continue to be, the standard by which scientific claims are evaluated. However, in many fields of study there are examples of scientific investigations that cannot be fully replicated because of a lack of time or resources. In such a situation, there is a need for a minimum standard that can fill the void between full replication and nothing. One candidate for this minimum standard is ‘reproducible research,’ which requires that data sets and computer code be made available to others for verifying published results and conducting alternative analyses.” Peng, R. (2009). *Reproducible research and Biostatistics*. Biostatistics, 10 (3): 405-408.

Therefore our definition of reproducibility will be that the computer code and data are made freely available to others for verification and conducting alternate analyses or for use in instructional purposes. And that the software used are also easily obtained and preferably open source to avoid licensing or other issues related to accessibility for end-users related to costs or non-standard file formats, etc.

In an effort to understand where as a discipline we stand as plant pathologists in regards to open science and reproducible research, we surveyed a broad selection of articles to represent a broad swathe of publications to evaluate our status.

# Materials and methods

## Article Selection

We randomly selected 300 articles published from 2012 to 2018 across 21 journals (Table ) and assigned them scores reflecting their openness and reproducibility. The journals represented both fundamental and applied areas as well as pathogen or other group-specific publications. A list of randomly selected pages was assigned to a randomized list of the 21 journals where the page number fell within an article for the given journal. In cases where an article was not suitable, *e.g.*, a review or otherwise not related to plant pathology, the next article was selected until a suitable article was found. Notes regarding the selection of articles can be found in the file, XXXX, available in this paper’s repository. The pages list was numbered from page one and went to page 150. This was done since some journals restart their numbering with each issue and also ensures that the journal is more likely to have a page number corresponding to the randomly generated value. This also assumes that there is no effect or bias on reproducibility based on the time of year that an article was published since most journals start with page number one at the beginning of the year. The list of journals was saved from Google’s sheets as a comma-separated-value (CSV) file for later use and distribution in a research compendium.

## Scoring criteria

Four individuals were assigned to rate a randomised list of journal articles.

Each journal was classified as to whether they were completely open (TRUE), behind a paywall (FALSE) or a combination (BOTH). The five-year impact factor for 2018 for each journal was retrieved from InCites Journal Citation Reports, Clarivate Analytics and entered in a separate sheet in the Google sheets file. This was downloaded, saved as a CSV file and is left-joined with the article notes using a left\_join() from the ‘dplyr’ package (Wickham et al. 2020) for analysis when the import\_notes() command is executed.

Articles were classified as fundamental or applied and a note was made whether the article’s primary focus was using or developing molecular techniques (TRUE/FALSE). This classification was based on the evaluator’s judgement of the article.

Where possible, the software used in the preparation of the publication was recorded in the notes if it was cited or otherwise specified in the article text.

Articles were scored in four areas for reproducibility on a scale of 0 to 3:

* **computational methods available**
  + 0 - Not available or not mentioned in the publication
  + 1 - Uses expensive proprietary software that only institutions would typically purchase (*i.e.*, ArcGIS standard is $7000)
  + 2 - Uses proprietary software that most individuals can afford (*e.g.*, Excel, Matlab? ($500))
  + 3 - Uses entirely open source and free software (*e.g.*, R, Julia, Python)
  + NA - No software was used in the research that can be determined as the article is written
* **software available**
  + 0 - Not available or not mentioned in the publication
  + 1 - Uses expensive proprietary software that only institutions would typically purchase (*i.e.*, ArcGIS standard is $7000US)
  + 2 - Uses proprietary software that most individuals can afford (*e.g.*, Excel) or SAS that is now available free for use with a university login
  + 3 - Uses entirely open source and free software (*e.g.*, R, Julia, Python).
  + NA - No software was used in the research that can be determined as the article is written
* **software cited**
  + 0 - not mentioned
  + 1 - Software mentioned by name only
  + 2 - Software cited with version number
  + 3 - All software components (SAS PROCs, R, Julia or Python packages) etc., cited
  + NA - No software was used in the research that can be determined as the article is written.
* **data available**
  + 0 - Not available or not mentioned in the publication
  + 1 - Available upon request to the author
  + 2 - Online, but inconvenient or non-permanent (*e.g.*, login needed, paywall, FTP server, personal lab website that may disappear, or may have already disappeared)
  + 3 - Freely available online to anonymous users for foreseeable future (*e.g.*, archived using Zenodo, dataverse or university library or some other proper archiving system)

Values for software packages were checked for spelling consistency and corrections were made manually where necessary. When working with these data in R, all strings of software character values were converted to fully uppercase to standardise the capitalisation and alleviate any issues with capitalisation used between evaluators. A custom R function, Reproducibility.in.Plant.Pathology::import\_notes(), was written to import the data, format the columns properly and calculate the overall reproducibility score in R (R Core Team 2020).

A total reproducibility score was calculated as the per cent of total possible for a given paper, *e.g.* some papers did not use specialized software therefore they were not scored for this category.

We recognise that the scoring methods were developed by the authors specifically for this exercise and are not entirely subjective. We have based our scoring decisions on our own experiences in the field of plant pathology. While many software packages may be widely used or “freely” available if one has an institutional licence, these types of software often hinder partners and interested parties that do not have these luxuries. Thus, the emphasis was given to using open source alternatives where possible over commercially available, non-free software packages. The view of the authors is one that this is something to be encouraged as it allows the broadest possible audience to benefit from the work at the least cost.

## Statistical analysis

Bayesian mixed-effect models were fit using the contributed package ‘brms’ (Bürkner 2018) in the R environment (R Core Team 2020) to evaluate the fixed effects of the publication (journal title), article class, molecular focus, five-year journal impact factor, journal open access policy on the reproducibility index and year to test for changes over time in the reproducibility index.

Relatively flat priors were selected with a mean 0 and a standard deviation 10 for the fixed effects.

The random effects were year and assignee for all models excepting the model which tested for year effects. In this model, the random effects were the publication title and assignee.

Models were fit using the hurdle\_gamma() family function due to zero-inflated continuous data. Each model was run for 3000 iterations in four chains.

The adapt\_delta() and max\_treedepth() values were adjusted as necessary on a case-by-case basis for each of the models to ensure that the chains mixed well.

Model finesses were evaluated using model summaries and diagnostic plots from ‘brms’ (Bürkner 2018) and posterior fits using pp\_check() from the contributed R package ‘bayesplot’ (Gabry et al. 2019).

Probability of direction or maximum probability of effect was plotted, pd(), and a test for practical equivalence, equivalence\_test(), was performed for each model using the contributed R package ‘bayestestR’ (Makowski et al. 2019).

# Results

Forty-eight articles scored 0% out of a possible 100% for reproducibility. The mean value was 31.2% and the median value was 33.3%. One article scored 100% or was classed as fully reproducible.

No journals surveyed were completely closed, all offered at least an option for open access but some were completely open.

The majority of the articles were classed as fundamental, 160, with the remaining 138 classed as applied.

Most articles did not make any computational methods available in any fashion with two, 0.007%, classing as “3,” which was the highest score available. Two hundred eighty-seven articles, 96%, were classed as “0” and nine articles, 0.03%, appeared to not use any computational methods.

More articles did a good job of using software that was reasonably available to anyone with 206 scoring “2” or “3” (Table ).

Software citations were also reasonably well reported in the literature with 120 scoring “2” where the main software was cited with version numbers and 57 scoring “3” where full citations including R packages or the SAS PROC used were cited where appropriate (Table ).

The most frequently cited software was SAS. The remainder were mostly specialised software except for Microsoft Excel, tied for 9th (Table ).

# Discussion

Unfortunately using our scoring method, most articles scored poorly for reproducibility or openness. Most journals in our discipline fail to mention the availability of the computational methods to support the article as published.

The software availability score was better as was the citing of software and versions used. SAS remains firmly entrenched in the discipline as the choice of software for statistical analysis followed by SPSS. It is interesting to note however, that with the popularity of Python that it does not even appear in our survey results whereas R was the fourth most commonly used software.

While the articles that were surveyed did not score well, we are aware of and would like to highlight that there are several good examples from the plant pathology discipline that can be used as exemplars.

There are different ways to make research more open and reproducible, which can be thought of as levels of reproducibility. The first level is including tables of raw data or code with the paper as a supplemental file. This is suitable for studies that may have a small data set or simple analysis or for demonstrations purposes as Madden et al. (2015) demonstrate in their discussion regarding the use of values in statistical analyses where they supply an *e-****X****tra*\* with reproducible examples for readers to refer to. This allows the reader to quickly view the extra materials and a DOI is assigned as a part of the article itself.

At the second level, Lehner et al. (2017) used GitHub to host a code repository of their research compendium website with data and a reproducible report that explains in detail all steps of the analysis and the R code for conducting a meta-analysis for assessing heterogeneity in the relationship between white mold incidence and soybean yield and between incidence and soybean tied (<https://emdelponte.github.io/paper-white-mold-meta-analysis/>). The website clearly demonstrates the analysis for readers and uses R so that anyone can easily replicate the work. Likewise, Duku et al. (2016) provide scripts for models, data and code for graphs, via a website hosted by GitHub, <http://adamhsparks.github.io/MICCORDEA/>, that can be used to replicate their study modelling the effects of climate change on rice bacterial blight and rice leaf blast in Tanzania.

The third level is the use of proper data repositories such as FigShare, Zenodo or OSF.io among others, allow for the deposition and updating of code, figures, data preprints or any other materials that support the article itself while providing a DOI and citation for a standalone citation. Sparks et al. (2011, 2014) used FigShare to provide models, data and code, (Sparks 2016) necessary to replicate model development and the subsequent study on the effects of climate change on potato late blight.

The fourth level includes packaging the analysis as a software package, *e.g.* R Julia or Python package at a minimum, with the use of continuous integration and possibly a software container like Docker as Khaliq et al. (Khaliq et al. 2020) have done. Their research compendium provides a Docker instance, DOI and full R package that enable readers to fully replicate their analysis of chickpea ascochyta conida dispersal using the data collected and stepping through other points where weather data were investigated and various models were fit before deciding on the best fit and recreate any figures as published in the article (Khaliq et al.).

Miorini et al. (2019)

* Example from Grünwald lab:
  + paper <http://apsjournals.apsnet.org/doi/full/10.1094/PHYTO-12-14-0350-FI>
  + github repo <https://github.com/grunwaldlab/Sudden_Oak_Death_in_Oregon_Forests>

## A general workflow

A general work flow for producing academic research involves clearly defining a research question, obtaining data for testing the hypothesis, summarizing/analyzing and presenting data and results, and writing the manuscript. Here we defined three levels of reproducibility which are also related to the evolution of computational methods and reproducible practices (Figure ).

The first level of reproducibility involves openly available research materials such as isolates or strains and/or nucleic acid sequences in public collections and citations for methods used. A second level involves providing raw data and code as binary files (PDF or other non-text files) in supplemental materials, which do not allow prompt access to the data and running the code because of use of expensive commercial software or a pay-wall. The highest level includes efforts to annotate structured raw data (FAIR – Findable, Accessible, Interoperable and Re-usable) (Wilkinson et al. 2016) and fully document the analysis using open source code which are deposited in public repositories and can be run by anyone following the download of data and code. The first level, as reported, is an essential step that is not substituted by the other practices and eventually researchers fail to provide sufficient description or correct citations. In the next section we present standards and tools that can be used to ensure reproducibility.

When making your science more open and reproducible, methods, software used, this includes items such as R, Julia or Python packages that were directly used in the analysis or production of the paper, etc. should be cited properly. Deposit and annotate biological materials with an herbarium or other repository. Provide a full description of all equipment used, *e.g.* a Spectrum Technologies Watchdog 2700 weather station was used to record wind speed, direction, rainfall, temperature and relative humidity at one-hour intervals. This allows end-users to identify what was used and identify the methods used more accurately. Just as importantly, this acknowledges the contributions of others whose works were instrumental in your research. This also helps ensure that you as a researcher can reconstruct what you have done since you will have good notes and documentation and be able to identify if something changes, *e.g.* a package version, what effect it had on your research.

The use of programming or scripting languages such as R, Julia, SAS or Python enables you as a scientist to keep very detailed records of what was computationally performed. This is as opposed to using software such as spreadsheet programs like Excel, Google Sheets, Numbers, Calc or others which can be used for simple statistical analyses and visualization (but should not be) or other point-and-click software packages that do not enable you to keep an accurate record of the steps taken to import, format, visualise and analyse data.

### Data

Prefer text files for saving small sets of data. Data that are saved in binary formats such as PDF files are difficult to reuse because they are not easily machine readable. In many cases, data sets are small enough and curated in spreadsheets, which should be saved as a plain text file, *e.g.* comma separated (CSV) files. This also helps ensure that the data are reusable. Larger data sets may warrant the use of a proper database like MariaDB or PostgreSQL which provide users with several benefits but two important ones to mention here are data redundancy, no duplicated records; and data consistency, all of your records are sure to be recorded in the same way for every observation. While databases may offer many advantages, the trade-off is that they are more complex to set up and administer, especially for a small data set that might be only a few rows and columns in a database.

Ideally, once the data are complete, best practices for keeping your data as you perform your work include treating the raw data as read-only and using file permissions to prevent changes to the raw data files. It should be noted, that the use of a database management system also allows for both of these at the expense of added complexity. Saving files in proprietary formats such as .xls(x) can also lead to issues in the future when opening using newer (or older) software versions. Unexpected changes to values in the data (Ziemann et al. 2016) may also occur when using proprietary formats.

If steps are followed to make the data FAIR then it will be readable by humans and machines alike this will help support discoveries and support further research. In turn, sharing data will lead to new citations for your work as others discover and use it.

To make your data the most widely usable, ensure that it has a persistent identifier. A digital object identifier (DOI) is the most common (<https://www.doi.org/>) but handle is also an option (<https://handle.net/>). There are different options for generating a DOI for your data and other materials. FigShare, Zenodo and OSF all offer long-term archival along with a service to generate a DOI for your materials. The use of a persistent identifier works to ensure that even if the data are moved, they can still be located using that unique identifier. For more on FAIR data, visit Go-Fair <https://www.go-fair.org/fair-principles/>.

### Sharing your research

Once you have determined how to best manage your source code for analysis and the data sets the next step is to consider how to share your data. Providers such as FigShare, Dataverse, OSF.io (the Open Science Framework) and Zenodo allow for you to deposit your data and generate a DOI for sharing your project once you are finished with it.

Other providers exist that allow for you to not only track changes but also to share the data openly, these include GitHub, Gitlab and Bitbucket. GitHub is arguably the most popular and widely used software development platform currently. Data that is encoded in CSV or other plain text formats can easily be deposited in a repository along with code for analysis to enable changes to be tracked and other users to download and replicate the work.

We would advise against the practice of depositing data on a laboratory website or a site such as GitHub only though. Doing either of these leaves the work in an unstable state where future users may be unable to access the work. It is a best practice to always ensure that you have deposited the data with a provider such as Zenodo, FigShare or OSF and generated a DOI for the materials to help ensure continued accessibility. Many of these providers provide rather easy ways to link the project with a software development repository to help ensure that the data are available in perpetuity.

Readers should also consult with their local librarians about local resources. Most universities provide a facility for staff to deposit papers and other academic materials, but this may extend to software development repositories as well.

Readers are encouraged to avoid using lab websites and other personal pages for sharing projects over the longer term as they are fraught with link-rot and are not an optimal way to share your projects.

# Authors’ contributions

# Data availability statement

# Acknowledgements

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# Tables

Table : Journal titles selected for inclusion and the number (n) of articles from each journal that were evaluated.

| journal | n |
| --- | --- |
| Australasian Plant Pathology | 8 |
| Canadian Journal of Plant Pathology | 13 |
| Crop Protection | 14 |
| European Journal of Plant Pathology | 11 |
| Forest Pathology | 14 |
| Journal of General Plant Pathology | 17 |
| Journal of Phytopathology | 14 |
| Journal of Plant Pathology | 15 |
| Molecular Plant Pathology | 18 |
| Molecular Plant-Microbe Interactions | 16 |
| Nematology | 11 |
| Physiological and Molecular Plant Pathology | 18 |
| Phytoparasitica | 19 |
| Phytopathologia Mediterranea | 9 |
| Phytopathology | 19 |
| Plant Disease | 15 |
| Plant Health Progress | 10 |
| Plant Pathology | 24 |
| Revista Mexicana de Fitopatología | 10 |
| Tropical Plant Pathology | 9 |
| Virology Journal (Plant Viruses Section) | 16 |

Table : Article ratings for software availability where ‘0,’ not available or not mentioned in publication to ‘3,’ preferably open source and freely available.

| software availability | n | percent |
| --- | --- | --- |
| 0 | 57 | 19.7% |
| 1 | 27 | 9.3% |
| 2 | 145 | 50.0% |
| 3 | 61 | 21.0% |
|  | 10 | - |

Table : Article ratings for software citations where ‘0,’ not specified or mentioned in publication to ‘3,’ fully cited with proper version numbers and all procedures detailed.

| software citation | n | percent |
| --- | --- | --- |
| 0 | 58 | 19.9% |
| 1 | 57 | 19.5% |
| 2 | 120 | 41.1% |
| 3 | 57 | 19.5% |
|  | 8 | - |

Table : ally of reproducibility scores where 0 is not reproducible and 12 is fully reproducible, *e.g.* all software cited, software used are freely available to anyone, computational methods (statistical analysis or other computational code) are freely available for download and use, and raw data are available for download and use.

| software used | n |
| --- | --- |
| SAS | 53 |
| MEGA | 42 |
| SPSS | 26 |
| R | 19 |
| BLAST | 16 |
| STATISTICA | 11 |
| GENSTAT | 10 |
| CLUSTAL W | 9 |
| CLUSTAL X | 7 |
| EXCEL | 7 |
| MRBAYES | 7 |
| BIOEDIT | 5 |
| PAUP | 5 |

Table : Tally of reproducibility scores where 0 is not reproducible and 12 is fully reproducible, *e.g.* all software cited, software used are freely available to anyone, computational methods (statistical analysis or other computational code) are freely available for download and use, and raw data are available for download and use.

| score | n | percent |
| --- | --- | --- |
| 0.0 | 53 | 17.7% |
| 8.3 | 5 | 1.7% |
| 11.1 | 2 | 0.7% |
| 16.7 | 7 | 2.3% |
| 25.0 | 44 | 14.7% |
| 33.3 | 83 | 27.7% |
| 41.7 | 52 | 17.3% |
| 44.4 | 1 | 0.3% |
| 50.0 | 22 | 7.3% |
| 55.6 | 1 | 0.3% |
| 58.3 | 14 | 4.7% |
| 66.7 | 12 | 4.0% |
| 75.0 | 2 | 0.7% |
| 91.7 | 1 | 0.3% |
| 100.0 | 1 | 0.3% |

# Figures

![](data:None;base64,)

Figure : An open and reproducible research workflow.

# Colophon

This report was generated on 2020-12-07 15:48:49 using the following computational environment and dependencies:

#> R version 4.0.3 (2020-10-10)  
#> Platform: x86\_64-apple-darwin17.0 (64-bit)  
#> Running under: macOS Catalina 10.15.7  
#>   
#> Matrix products: default  
#> BLAS: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib  
#> LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib  
#>   
#> locale:  
#> [1] en\_AU.UTF-8/en\_AU.UTF-8/en\_AU.UTF-8/C/en\_AU.UTF-8/en\_AU.UTF-8  
#>   
#> attached base packages:  
#> [1] stats graphics grDevices utils datasets methods base   
#>   
#> other attached packages:  
#> [1] rsvg\_2.1   
#> [2] DiagrammeRsvg\_0.1   
#> [3] DiagrammeR\_1.0.6.1   
#> [4] tidyr\_1.1.2   
#> [5] officer\_0.3.15   
#> [6] flextable\_0.6.0   
#> [7] janitor\_2.0.1   
#> [8] dplyr\_1.0.2   
#> [9] Reproducibility.in.Plant.Pathology\_0.0.1-1  
#>   
#> loaded via a namespace (and not attached):  
#> [1] zip\_2.1.1 Rcpp\_1.0.5 RColorBrewer\_1.1-2   
#> [4] compiler\_4.0.3 pillar\_1.4.7 base64enc\_0.1-3   
#> [7] tools\_4.0.3 digest\_0.6.27 uuid\_0.1-4   
#> [10] jsonlite\_1.7.1 lubridate\_1.7.9.9001 evaluate\_0.14   
#> [13] memoise\_1.1.0 lifecycle\_0.2.0 tibble\_3.0.4   
#> [16] pkgconfig\_2.0.3 rlang\_0.4.9 rstudioapi\_0.13   
#> [19] cli\_2.2.0 curl\_4.3 yaml\_2.2.1   
#> [22] officedown\_0.2.0 xfun\_0.19 stringr\_1.4.0   
#> [25] xml2\_1.3.2 knitr\_1.30 htmlwidgets\_1.5.2   
#> [28] hms\_0.5.3 gdtools\_0.2.2 generics\_0.1.0   
#> [31] vctrs\_0.3.5 systemfonts\_0.3.2 tidyselect\_1.1.0   
#> [34] data.table\_1.13.3 glue\_1.4.2 snakecase\_0.11.0   
#> [37] R6\_2.5.0 rvg\_0.2.5 fansi\_0.4.1   
#> [40] rmarkdown\_2.5 readr\_1.4.0 purrr\_0.3.4   
#> [43] magrittr\_2.0.1 htmltools\_0.5.0 ellipsis\_0.3.1   
#> [46] assertthat\_0.2.1 V8\_3.4.0 stringi\_1.5.3   
#> [49] visNetwork\_2.0.9 crayon\_1.3.4.9000

The current Git commit details are:

#> Local: main /Users/adamsparks/Sources/GitHub/Analysis/Reproducibility\_in\_Plant\_Pathology  
#> Remote: main @ origin (git@github.com:openplantpathology/Reproducibility\_in\_Plant\_Pathology.git)  
#> Head: [d21ddb5] 2020-12-07: Fleshing out the discussion section