- 1 Reproducible, flexible and high throughput data extraction from primary
- 2 literature: The metaDigitise R package
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8 Abstract

- 9 Research synthesis, especially in the form of meta-analysis, requires data extraction
- 10 from primary studies. Meta-analysis synthesizes effect sizes, often calculated from
- 11 summary statistics of studies. However, exact values of such statistics are commonly
- 12 hidden in figures. The R package **metaDigitise** extracts descriptive statistics such as
- 13 means, standard deviations and, if applicable, correlations from the four types of plots:
- 14 1) mean and error plots (e.g. bar graphs with standard errors), 2) box plots, 3) scatter
- 15 plots and 4) histograms. The package interactively guides the user through data
- 16 extraction process. Notably, it enables a large-scale extraction using image files, letting
- 17 the user stop processing, edit and add to the resulting data fame at any point. Further,
- 18 it facilitates reproducible data extraction from plots with little inter-observer bias, thus,
- 19 allowing a group of people to participate the extraction of data collaboratively.
- 20 Keywords: meta-analysis, comparative analysis, data extraction, R, reproducibility,
- 21 figures, images, summary statistics

22 Introduction

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23
    In many different contexts, researchers need to make use of data presented in primary
    literature. Most notably, this includes meta-analysis, which is becoming increasingly
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25
    common in many research fields. Meta-analysis uses effect size estimates and their
    sampling variance, taken from many studies, to understand whether particular effects
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    are common across studies and to explain variation among these effects (Glass, 1976;
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    Borenstein et al., 2009; Koricheva, Gurevitch & Mengersen, 2013; Nakagawa et al.,
28
    2017). Meta-analysis therefore relies foremost on data extracted from primary
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    literature, and more specifically, descriptive statistics (e.g., means, standard deviations,
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31
    correlation coefficients) that have been reported in the text or tables of research papers.
    Descriptive statistics are also, however, frequently presented in figures and so need to be
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    manually extracted using digitising programs. While inferential statistics (e.g., t- and
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    F-statistics) are often presented along side descriptive statistics and can be used to
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    derive effect sizes, descriptive statistics are much more appropriate to use because
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    sources of non-independence in experimental designs can be dealt with more easily
36
    (Noble et al., 2017). Although there are several existing tools to perform tasks like this
37
    (e.g. DataThief (Tummers, 2006), GraphClick (Arizona-Software, 2008), WebPlotDigitizer
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    (Rohatgi, 2017)), these tools are not designed specifically for meta-analysis for three
    main reasons.
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    First, they typically only provide the user with calibrated x,y coordinates from
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    imported figures, and do not differentiate between common plot types that are used to
    present data. This means that a large amount of downstream data manipulation is
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    subsequently required, that is different across plots types. For example, data are
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    frequently presented in mean and error plots (Figure 1A), for which the user wants a
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    mean and error estimate for each group presented in the figure. With existing
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47
    programs, x, y coordinates of means and errors are returned, to which the user must
    manually discern between mean and error coordinates and assign points to groups. The
48
```

- 49 error then needs to be calculated as the deviation from the mean, and then transformed
- 50 to a standard deviation, depending on the type of error presented.
- 51 Second, digitising programs do not easily allow the integration of metadata at the time
- 52 of data extraction, such as experimental group or variable names, and sample sizes.
- 53 This makes the downstream calculations more laborious, as the information has to be
- 54 added later, in most cases using different software.
- 55 Finally, existing programs do not import a set of images and allow the user to
- 56 systematically work through them. Instead they require the user to manually import
- 57 images one by one, and export data into individual files, that need to be imported and
- 58 edited using different software. In essence, existing software does not provide an
- 59 optimized research pipeline to facilitate data extraction, editing and
- 60 reproducibility.
- 61 These are major issues because extracting from figures can be an incredibly
- 62 time-consuming process. Furthermore, although meta-analysis is an important tool in
- 63 consolidating the data from multiple studies, many of the processes involved in data
- 64 extraction are opaque and difficult to reproduce, making extending studies problematic.
- 65 Having a tool that facilitates reproducibility in meta-analyses will increase transparency
- and go a long way to resolving the reproducibility crises we are seeing in many fields
- 67 (Peng, Dominici & Zeger, 2006; Peng, 2011; Sandve et al., 2013; Parker et al., 2016; Ihle
- 68 et al., 2017).
- 69 Here, we present an interactive R package, **metaDigitise**, which is designed for large
- 70 scale data extraction from figures, specifically catering to the the needs of
- 71 meta-analysts. To this end, we provide tools specific to data extraction from common
- 72 plot types (mean and error plots, box plots, scatter plots and histograms, see Figure 1).
- 73 metaDigitise operates within the R environment making data extraction, analysis and
- 74 export more streamlined. It also provides users with options to conduct the necessary
- 75 calculations on processed data immediately after extraction so that comparable

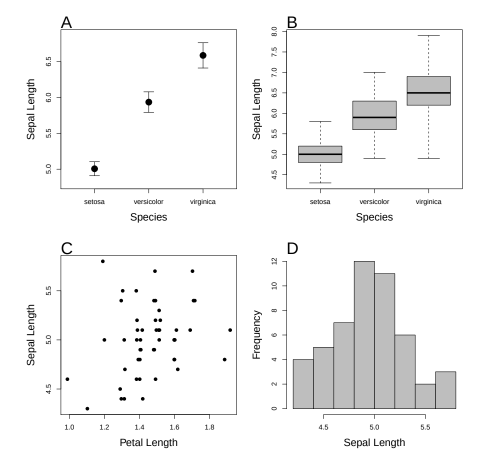


Figure 1: Four plot types that **metaDigitise** is designed to extract data from: A) mean and error plot, B) box plot, C) scatter plot and D) histogram. Data is taken from the iris dataset in R. A and B are plotted with the whole dataset, C and D are just the data for the species setosa.

summary statistics can be obtained quickly. **metaDigitise** condenses summary data extracted from multiple figures into a single data frame which can be can easily exported. Processed data can also be easily extracted and analysed in any way the user desires in downstream analysis within R. Conveniently, when needing to process many figures at different times **metaDigitise** will only import figures not already completed within a directory. This makes it easy to add new figures at any time. **metaDigitise** has also been built for reproducibility in mind. It has functions that allow users to redraw their digitisations on figures, make corrections and access the raw calibration data which is written automatically for each figure that is digitised into a special folder within the directory. This makes sharing figure digitisation and reproducing the work of others

87 Directory Structure, Image Processing and

88 Reproducibility

- The **metaDigitise** package is designed to be flexible, yet simple to use. There is one 89 main function in the package, metaDigitise(), which interactively takes the user 90 through the process of extracting data from figures. metaDigitise() was created with the idea that the user would likely have multiple images to extract from. It therefore 92 operates in the same way whether the user has one or multiple images. 93 metaDigitise() is designed to work on a directory containing images of figures copied 94 from primary literature, in .png, .jpg, .tiff, .pdf format. This directory is specified to 95 metaDigitise() through the dir argument. The user is free to set their own broad 96 directory structure (e.g. one directory for all images or one directory for each paper 97 extracted from). We would recommend having all files for one project in a single 98 directory with an informative and unambiguous naming scheme for images to make it easy to identify the paper and figure the data come from. This cuts out the need to 100 101 change directories constantly. For example the directory structure could look like:
 - * Main project directory
 - + FiguresToExtract/
 - + Paper1_Figure1_trait1.png
 - + Paper1_Figure2_trait2.png
 - + Paper1_Figure3_trait3.png
 - + Paper2_Figure1_trait1.png
 - + Paper2_Figure2_trait2.png
 - + Paper2_Figure3_trait3.png

102 It is important for the user to think about their directory structure early on in this 103 process (also more generally in the context of their entire project), especially if they 104 plan to share the extractions with collaborators or when publishing the project. 105 When metaDigitise() is run, it recognizes all the images in a directory and automatically imports them one by one, allowing the user to click and enter relevant 106 information about a figure as they go. This expedites digitising figures by preventing 107 users from having to constantly change directories and / or open new images. The data 108 109 from a completed image is automatically saved as a metaDigitise object in an .RDS file to a caldat directory that is created within the parent directory when first 110 executing the metaDigitise() function. These files enable re-plotting and editing of 111 images at a later point (see below). 112 A particularly powerful and flexible aspect of metaDigitise() is its ability to identify 113 114 images that have been previously digitised and only import images that have not been digitised in subsequent calls of the function. This means that all figures do not need to 115 be extracted at one time and that new figures can be added as the project develops. 116 After each image is extracted, the user is asked whether they wish to continue or quit 117 the extraction process. Upon rerunning metaDigitise(), previously digitised figures 118 119 are simply ignored during processing, but their data is re-integrated within the final 120 output after new files are completed automatically. After completing all images, or upon quitting, the processed data (in a form specified 121 by the user) is then returned. From all plot types, metaDigitise() summarises the 122 data from a figure as a mean, standard deviation and sample size, for each identified 123 124 group within the plot (should multiple groups exist). These are the descriptive statistics needed to create many of the relevant effect sizes and sampling error for a 125 meta-analysis. In the case of scatter plots, metaDigitise() also returns the correlation 126 coefficient between the points within each identified group. 127

128 Diverse Plot Types

129 metaDigitise recognises four main types of plot; Mean and error plots, box plots, scatter plots and histograms, shown in Figure 1. Each of these can be processed 130 131 together and integrated into a single output. Alternatively, users can keep like figures 132 together and process them separately. In order to correctly extract data from figures metaDigitise() always requires the user 133 to calibrate the axes in the figure. To do this, the user is required to click on two known 134 points on the axis in question, and then enter the value of those points in the figure. 135 Using this information, metaDigitise() then calculates the value of any clicked points 136 in terms of the figure axes. In the case of mean and error plots and box plots, it 137 calibrates only the y-axis (assuming the x-axis is redundant). For scatter plots and 138 histograms both axes are calibrated. 139

140 Mean and error plots

metaDigitise() prompts the user to enter group names and allows the user to enter 141 sample sizes (n), which are used in downstream processing. The user is then prompted 142 143 to click on an error bar followed by the mean. Error bars above or below the mean can be clicked - sometimes one is clearer than the other. metaDigitise() assumes that the 144 error bars are symmetrical. Where the user has clicked the error is displayed in a 145 different colour to the mean (Figure 2A). The user can subsequently add more groups, 146 147 edit groups or remove groups. Finally the user is asked what type of error was used in the figure: standard deviation (SD, σ), standard error (SE) or 95% confidence intervals 148 (CI95). Standard deviation is calculated from standard error as 149

$$\sigma = SE\sqrt{n} \tag{1}$$

150 and from 95% confidence intervals as

$$\sigma = \frac{CI}{1.96}\sqrt{n} \tag{2}$$

151 If the user does not enter a sample size at the time of data extraction (if, for example, 152 the information is not readily available) the SD is not calculated. This can be entered 153 at a later time, however (see below). A function, error_to_sd(), that converts the 154 different error types to SD is also available in the package.

155 Box plots

As with mean and error plots, metaDigitise() prompts the user to enter group names and allows the user to enter sample sizes (n), which are used in downstream processing. The user is then prompted to click on the maximum (b), upper quartile (q_3) , median (m), lower quartile (q_1) and minimum (a). metaDigitise() will check that the maximum is greater than the minimum, and return a warning if that is not the case. The user can subsequently add, edit or remove groups. From the extracted data, the mean (μ) and SD are calculated as

$$\mu = \frac{(n+3)(a+b) + 2(n-1)(q_1 + m + q_3)}{8n} \tag{3}$$

163 following Bland (2015) and

$$\sigma = \frac{b - a}{4\Phi^{-1}(\frac{n - 0.375}{n + 0.25})} + \frac{q_3 - q_1}{4\Phi^{-1}(\frac{0.75n - 0.125}{n + 0.25})} \tag{4}$$

where $\Phi^{-1}(z)$ is the upper zth percentile of the standard normal distribution, following Wan et al. (2014). As with mean and error plots, if the user does not enter a sample size at the time of data extraction the SD is not calculated. Two functions, rqm_to_mean() and rqm_to_sd(), that convert box plot data to mean and SD 68 respectively are also available in the package.

169 Scatter plots

metaDigitise() prompts the user to enter groups names and then to click on points.

Points added by mistake can be deleted. The user can subsequently add groups, edit

groups (add or remove points) or delete groups. Different groups are plotted in different

colours and shapes, with a legend at the bottom of the figure (Figure 2C). Mean, SD

and sample size are calculated from the clicked points, for each group. Where the

sample size from the clicked points does not match a known sample size (e.g. if there

are overlaid points), the user can enter an alternate sample size.

177 Histograms

metaDigitise() prompts the user to click on the top corners of each bar. Bars can subsequently be deleted. For each bar a midpoint (m; mean x coordinates) and a frequency (f; mean y coordinates, rounded to the nearest integer) is calculated. The sample size, mean and SD are calculated as:

$$n = \sum_{i=1}^{n} f_i \tag{5}$$

$$\mu = \frac{\sum_{i=1}^{n} m_i f_i}{n} \tag{6}$$

$$\sigma = \sqrt{\frac{\sum_{i=1}^{n} (m_i f_i - \mu f_i)^2}{n-1}}$$
 (7)

As with the scatterplots, if the sample size from the extracted data does not match a known sample size, the user can enter an alternate sample size.

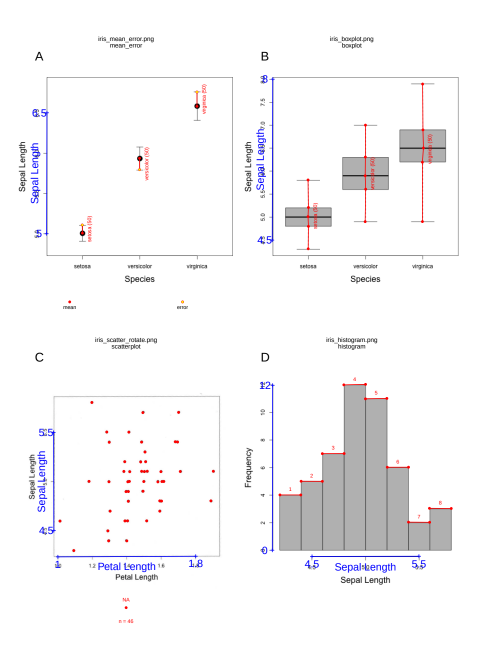


Figure 2: Demonstration of data extraction from different plot types

184 Extracting Data From Plots

- We will now demonstrate how metaDigitise() works using figures generated from the well known iris data set. Users can install the metaDigitise package from GitHub as follows:
 - R> install.packages("devtools")
 - R> devtools::install_github("daniel1noble/metaDigitise")

R> library(metaDigitise)

Assume that the user would like to extract descriptive statistics from studies measuring sepal length or width in iris species for a fictitious project. There are a few studies that only present these data in figures. As the user reads papers found from a systematic search, they add figures with relevant data to a "FiguresToExtract" folder as follows

*FiguresToExtract/

193

194

195

196

+ 001_Anderson_1935_Fig1.png

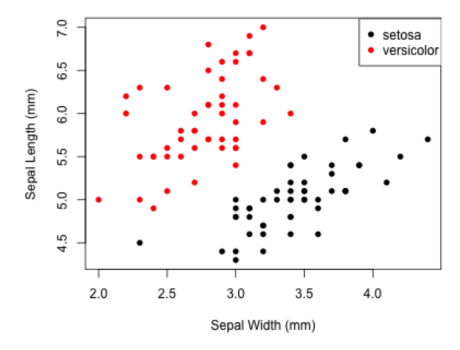


Figure 3: Example scatterplot (001_Anderson_1935_Fig1.png) of sepal length and width for two species of iris (setosa and versicolor)

Here, the naming of the files placed in the folder will contain the paper number, first author and the figure number to keep data uniquely associated with figures. At first there is one figure in the folder, shown in Figure 3. Running metaDigitise() brings up a series of prompts for the user using a main menu that provides access to a number of

its features ("..." here represents the user's path to the project directory):

R> digitised_data <- metaDigitise(".../FiguresToExtract", summary = TRUE)</pre>

Do you want to...

- 1: Process new images
- 2: Import existing data
- 3: Edit existing data

Selection:

198 The user simply enters in the numeric value that corresponds to what they would like to

199 do. In this case they want to "Process new images". The user is then asked whether

200 there are different types of plot(s) in the folder. This question is most relevant when

201 there are lots of different figures in the folder because it will then ask the user for the

202 type of figure as they are cycled through.

Are all plot types Different or the Same? (d/s)

203 metaDigitise() then asks the user whether the figure needs to be rotated or flipped.

204 This can be needed when box plots and mean and error plots are not orientated

205 correctly. In some cases, older papers can give slightly off angled images which can be

206 corrected by rotating. So, in this prompt the user has three options: f for "Flip", r for

207 "rotate" or c for "continue".

mean_error and boxplots should be vertically orientated

-| | I.E. o NOT |-o-| |

If they are not then chose flip to correct this.

If figures are wonky, chose rotate.

Otherwise chose continue

Flip, rotate or continue (f/r/c)

R > c

After this, metaDigitise() will ask the user to specify the plot type. Depending on the figure, the user can specify that it is a figure containing the mean and error (m), a box plot (b), a scatter plot (s) or a histogram (h). If the user has specified d instead of s in response to the question about whether the plot types are the same or different, this question will pop up for each plot, but will only be asked once if plots are all the same.

Please specify the plot_type as either:

m: Mean and error

b: Box plot

s: Scatter plot

h: Histogram

R> s

214 After selecting the figure type a new set of prompts will come up that will ask the user

215 first what the y and x-axis variables are. This is useful as users can keep track of the

216 different variables across figures and papers. Here, the user can just add this

217 information in to the R console. Once complete, details on how to calibrate the x and

218 y-axis appear, so that the relevant statistics / data can be correctly calculated. When

219 working with a plot of mean and standard errors, the x-axis is rather useless in terms of

```
What is the y variable?
R> Sepal Length (mm)
What is the x variable?
R> Sepal Width (mm)
On the Figure, click IN ORDER:
     y1, y2 , x1, x2
   Step 1 ----> Click on known value on y axis - y1
 у1
   Step 3 ----> Click on known value on x axis - x1
  |____x1_____
```

220 calibration so metaDigitise() just asks the user to calibrate the y-axis.

. . . .

The user can just follow the instructions on screen step-by-step (instructions above have been truncated by '...' to simplify), and in the order specified. Before moving on, the user is forced to check whether or not the calibration has been set up correctly. If n is chosen because something needs to be fixed then the user can re-calibrate.

What is the value of y1 ?

R> 4.5

What is the value of y2 ?

R> 7

What is the value of x1 ?

R> 2

What is the value of x2 ?

R> 4

Re-calibrate? (y/n)

R> n

Often, plots might contain multiple groups that the meta-analyst wants to extract from. 225 metaDigitise() handles this nicely by prompting the user to enter the group first, 226 227 followed by digitisation of this groups data. After digitising the first group, and having 228 exited, metaDigitise() will ask the user whether they would like to add another group. Users can continually add groups (a), delete groups (d), edit groups (e) or finish 229 a plot and continue to the next one (f - if another plot exists). The number of groups 230 are not really limited and users can just keep adding in groups to accommodate the 231 232 different numbers that may be presented across figures (although it can get complicated with too many). 233

If there are multiple groups, enter unique group identifiers (otherwise press enter) Group identifier:

R> setosa

Click on points you want to add.

If you want to remove a point, or are finished with a group, exit by clicking on red box in bottom left corner, then follow prompts

- 234 To finish selecting points, the user can exit by clicking on the red button that appears
- 235 when extracting points. The user is then asked if they want to add or delete points
- 236 from that group.

Add or Delete points to this group, or Continue? (a/d/c)

R> c

- 237 Once we are done digitising all the groups our plot will look something like Figure
- 238 4.
- When completed metaDigitise() will write the digitised data as a metaDigitise
- 240 object to a RDS file in the caldat directory, such that our new directory structure is as
- 241 follows
 - *FiguresToExtract/
 - + caldat/
 - + 001_Anderson_1935_Fig1
 - + 001_Anderson_1935_Fig1.png
- 242 Users can access the metaDigitise object created (001_Anderson_1935_Fig1) at any
- 243 time using the metaDigitse() function. In the R console, the summarised data for the
- 244 digitised figure can be printed on screen or even written to a .csv file:

R> digitised_data

filename group_id variable mean error error_type n r sd plot_type

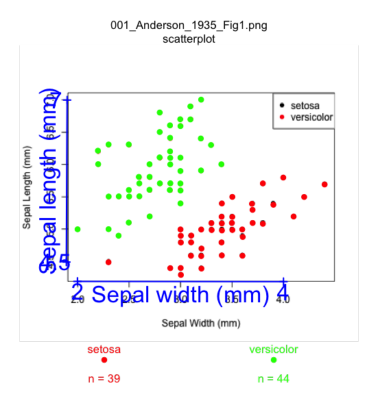


Figure 4: Digitisation of sepal length and width for two species of iris (setosa and versicolor). Names of the variables and calibration (in blue) are plotted alongside the digitised points (green = versicolor; red = setosa). The sample sizes for each group are provided on the lower part of the plot. All figures are clearly labelled at the top to remind users of the filename and plot type. This reduces errors throughout the digitisation process.

```
001_Anderson_1935_Fig1.png
                              setosa Sepal width (mm) 3.42 0.40
                                                                           39 0.75 0.40 scatterplot
001_Anderson_1935_Fig1.png
                              setosa Sepal length (mm) 5.00 0.38
                                                                              0.75 0.38 scatterplot
                                                                           39
001_Anderson_1935_Fig1.png versicolor Sepal width (mm) 2.77 0.32
                                                                           44 0.52 0.32 scatterplot
001_Anderson_1935_Fig1.png versicolor Sepal length (mm) 5.95 0.53 sd
                                                                           44 0.52 0.53 scatterplot
```

245 The mean for each of the two variables, along with the two species, are provided. Since 246 this is a scatterplot, the user also gets the Person's correlation coefficient between sepal 247 length and width for each species. These match reasonably well with the actual means of sepal length and width for each of the species in the full 'iris' dataset: 248

```
Species meanSL meanSW
              5.006
      setosa
                     3.428
2 versicolor
              5.936
                     2.770
```

1

One thing anyone with a familiarity with the iris dataset will notice is that the sample 249

250 sizes for each of these species (which are n = 50 each) are quite a bit lower. This is an example of some of the challenges when extracting data from scatter plots. Often data 251 points will overlap with each other making it impossible (without having the real data) 252 to know whether this is a problem. However, a meta-analyst will probably realise that 253 254 the sample sizes here conflict with what is reported in the paper. Hence, **metaDigitise** also provides the user with options to input the sample sizes directly (see Editing 255 section below), even for scatter plots and histograms where, strictly speaking, this 256 257 should not be necessary. Nonetheless, it is important to recognise the impact that overlapping points can have on summary statistics, particularly its effects on standard 258 deviation (SD) and standard error (SE). Here, the mean point estimates are nearly 259 exactly the same as the true values, but the SD's are slightly over-estimated: 260

Species meanSL meanSW

1 setosa 0.3524897 0.3790644

2 versicolor 0.5161711 0.3137983

261 Adding new figures

Users can add additional figures as new papers with relevant information are found.

Each figure should be in its own file with unique naming, even if a single paper has

multiple figures for extraction. For example, another paper on different populations

(and one new species) of iris contained two additional figures where important data

could be extracted. These figures can simply be named accordingly and added directly

to the same extraction folder:

*FiguresToExtract/

- + caldat/
 - + 001_Anderson_1935_Fig1
- + 001_Anderson_1935_Fig1.png
- + 002_Doe_2013_Fig1.png

+ 002_Doe_2013_Fig3.png

The user has already processed one figure (001_Anderson_1935_Fig1.png). We can tell this because the caldat folder has digitised data in it (caldat/001_Anderson_1935_Fig1).

Now the user has two new figures that have not yet been digitised. This example will nicely demonstrate how users can easily pick up from where they left off and how all previous data gets re-integrated. It will also demonstrate how different plot types are handled. All we have to do to begin, is again, provide the directory where all the figures are located:

R> digitised_data <- metaDigitise(".../FiguresToExtract", summary = TRUE)</pre>

The user gets the same set of prompts and simply chooses option one. This will permit users to digitise new figures, and will integrate previously completed digitisations along with newly digitised data together at the end of the session, or when the user decides to quit. This time, 001_Anderson_1935_Fig1.png is ignored and the new plots cycle on screen; first 002_Doe_2013_Fig1.png and then 002_Doe_2013_Fig3.png. Since there are a few different figure types, the user answers the first question in the R console as "d":

Are all plot types Different or the Same? (d/s)

R>d

275

276

277

278

279

280

281

**** NEW PLOT ****

mean_error and boxplots should be vertically orientated

| | I.E. o NOT |-o-| | If they are not then chose flip to correct this.

If figures are wonky, chose rotate.

Otherwise chose continue

Flip, rotate or continue (f/r/c)

R> c

Please specify the plot_type as either:

m: Mean and error

b: Box plot

s: Scatter plot

h: Histogram

R> m

Here, the user specifies the new plot type as m for 002_Doe_2013_Fig1.png because the

283 user has a plot of the mean and error of sepal length for each of the three species. The

284 user is then prompted a bit differently from our scatter plot as the x-axis is not needed

285 for calibration:

What is the y variable?

R> Sepal length

On the Figure, click IN ORDER:

y1, y2

```
Step 1 ----> Click on y1
  у1
   Step 2 ----> Click on y2
  у2
What is the value of y1 ?
R> 5
What is the value of y2 ?
R> 6.5
Re-calibrate? (y/n)
R> n
Do you know sample sizes? (y/n)
R> y
If there are multiple groups, enter unique group identifiers (otherwise press enter)
```

Group identifier:

R> setosa

Group sample size:

R> 50

Click on Error Bar, followed by the Mean

Add group, Edit Group, Delete group or Finish plot? (a/e/d/f)

R> a

Again, metaDigitise() will simply guide the user through digitising each of these figures describing to them exactly what needs to be done. At any point if mistakes are made the user can choose relevant options to edit or correct things before ending the figure. This process continues for each plot so long as the user would like to continue and after completing a single plot the user is always prompted as follows:

Do you want continue: 1 plots out of 2 plots remaining (y/n)

R> y

This continues until users have completed all non-digitised figures in the folder, at which point metaDigitise() concatenates the new data with previously digitised data in the object:

data

filename	group_id	variabl	e mean	error	error_typ	e n	r	sd	plot_type
001_Anderson_1935_Fig1.png	setosa	Sepal width (m	m) 3.42	0.40	sd	39	0.75	0.40	scatterplot
001_Anderson_1935_Fig1.png	setosa	Sepal length (mm) 5.00	0.38	sd	39	0.75	0.38	scatterplot
001_Anderson_1935_Fig1.png	versicolor	Sepal width (m	m) 2.77	0.32	sd	44	0.52	0.32	scatterplot
001_Anderson_1935_Fig1.png	versicolor	Sepal length (mm) 5.95	0.53	sd	44	0.52	0.53	scatterplot
002_Doe_2013_Fig1.png	setosa	Sepal length	5.00	0.11	se	50	NA	0.78	mean_error
002_Doe_2013_Fig1.png	viriginica	Sepal length	6.59	0.18	se	50	NA	1.26	mean_error
002_Doe_2013_Fig1.png	versicolor	Sepal length	5.94	0.14	se	50	NA	1.01	mean_error
003_Doe_2013_Fig3.png	catana	Sepal length	4.95	0.36	sd	50	NA	0.36	histogram

294 Re-importing, Editing and Plotting Previously

295 Digitised data

- 296 A particularly useful feature of **metaDigitise** is its ability to re-import, edit and re-plot
- 297 previously digitised figures. We can do this from the initial options from
- 298 metaDigitise()
 - R> digitised_data <- metaDigitise(".../FiguresToExtract")</pre>
 - Do you want to...
 - 1: Process new images
 - 2: Import existing data
 - 3: Edit existing data

Selection:

- 299 If the user chooses "Import existing data", they have the option of either 1) importing
- 300 data from all digitised images or 2) importing data from a single image that has been
- 301 digitised. If 2, then a list of files are provided to the user that they can select. Editing
- 302 existing data allows users to easily re-plot or edit information or digitisations that have
- 303 previously be done for any plot. This is accomplished by guiding the user through a
- 304 new set of options:

Choose how you want to edit files:

- 1: Cycle through images
- 2: Choose specific file to edit
- 3: Enter previously omitted sample sizes

Selection:

- 305 If the user is unsure about the name of the specific figure they need to edit or simply
- 306 want to just check the digitisations of figures they can choose "Cycle through images",
- 307 which will bring up each figure, one by one, overlaying the calibrations, group names (if

they exist), sample sizes (if they were entered) and the selected points. The user will then be given the choice to edit individual images. Alternatively, choosing option 2, will bring up a list of the completed files in the folder and the specific file can be chosen, at which point it will be replotted. Either of these options will cycle through a number of questions asking the user what they would like to edit:

Edit rotation? If yes, then the whole extraction will be redone (y/n)R> n Change plot type? If yes, then the whole extraction will be redone (y/n)R> n Variable entered as: R> Sepal length Rename Variables (y/n) R> n Edit calibration? (y/n)R> n Re-extract data (y/n) R > yChange group identifier? (y/n) R> n Add group, Delete group or Finish plot? (a/d/f) R > d

1: setosa

2: versicolor

3: viriginica

Selection:

R> 2

Add group, Delete group or Finish plot? (a/d/f)

R> a

- 313 A whole host of information can be edited including the rotation, plot type, the variable
- 314 name(s) that were provided, the calibration and even the digitisation of groups. When
- 315 editing the metaDigitise object is re-written to the caldat folder and the edits are
- immediately integrated into the existing object once complete.

317 Additional Features

318 Figure Rotation and Adjustment

- 319 Figures may have been extracted from old publications, for example from scanned
- 320 images, and so are not perfectly orientated on the image. This will make the calibration
- 321 of the points in the figure from the image problematic. metaDigitise() allows users to
- 322 rotate the image. By clicking two points on the x-axis, metaDigitse calculates the angle
- 323 needed to rotate the image so the x-axis is horizontal, and rotates it. (Figure
- 324 5A,B)
- 325 Furthermore, some figures, including mean and error, boxplots or histograms, may be
- 326 presented with horizontal bars. metaDigitise() assumes that the bars are vertical, but
- 327 allows the user to flip the image so that the bars are vertical if provided horizontally
- 328 (Figure 5C,D).

329 Obtaining Processed Data

While metaDigitise() provides users with the summary statistics by default, for all plot types, in many cases the user may actually be interested in obtaining the processed digitised data from scatter plots (i.e. calibrated points). This is very easy to do my changing the default summary argument from TRUE to FALSE in metaDigitise().

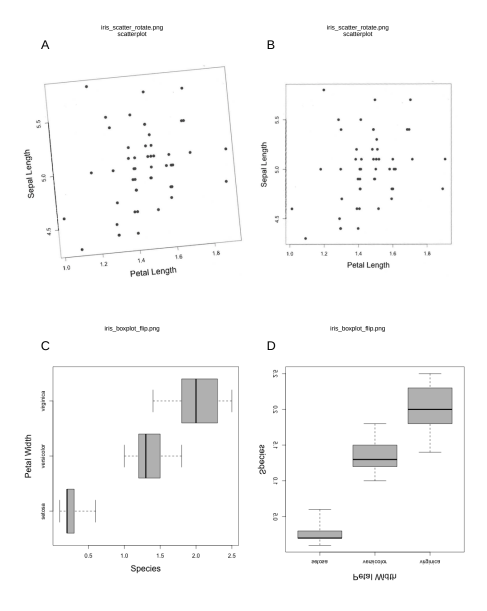


Figure 5: Figure rotation. A) and B) show how non-aligned images can be realigned through user defined rotation. C) and D) show how figures can be re-orientated so as to aid data input.

- Instead of providing the user with summary statistics it will return a list containing
- four slots for each of the figure types (mean error, box plot, histogram and scatter 335
- plots). An example of a data object returned from digitising figures is as follows:

>R str(data)

```
List of 3
 $ mean_error :List of 1
  ..$ 002_Doe_2013_Fig1.png:'data.frame': 3 obs. of 5 variables:
  .. ..$ id
                 : Factor w/ 3 levels "setosa", "versicolor", ...: 1 2 3
                 : num [1:3] 5 5.93 6.59
  .. ..$ mean
                 : num [1:3] 0.111 0.148 0.178
  .. ..$ error
  .. ..$ n
                 : num [1:3] 50 50 50
  .... $\square\text{variable: chr [1:3] "Sepal length" "Sepal length" "Sepal length"
 $ hist
              :List of 1
  ..$ 003_Doe_2013_Fig3.png:'data.frame': 8 obs. of 3 variables:
  ....$ midpoints: num [1:8] 4.3 4.5 4.7 4.9 5.1 ...
  ....$ frequency: num [1:8] 4 5 7 12 11 6 2 3
  ....$ variable : chr [1:8] "Sepal length" "Sepal length" ...
 $ scatterplot:List of 1
  ..$ 001_Anderson_1935_Fig1.png:'data.frame': 83 obs. of 8 variables:
  .. ..$ id
                   : Factor w/ 2 levels "setosa", "versicolor": 1 1 1 1 1 ...
  .. ..$ x
                   : num [1:83] 2.3 2.9 3 3 3 ...
  .. ..$ y
                   : num [1:83] 4.5 4.4 4.41 4.3 4.8 ...
                   : num [1:83] 1 1 1 1 1 1 1 1 1 1 ...
  .. ..$ group
  .. ..$ col
                   : Factor w/ 2 levels "red", "green": 1 1 1 1 1 1 1 1 1 ...
                   : num [1:83] 19 19 19 19 19 19 19 19 19 ...
  .. ..$ pch
  ....$ y_variable: chr [1:83] "Sepal length (mm)" "Sepal length (mm)"
```

....\$ x_variable: chr [1:83] "Sepal width (mm)" "Sepal width (mm)"

Here, the user can easily access the list of processed scatter plot data by simply extracting the scatter plot slot:

>R scatterplot <- data\$scatterplot

339 Adding sample sizes to previous Digitisations

In many cases important information, such as sample sizes, may not be readily available 340 or clear when digitising figures. In these circumstances users will have answered 'no' to 341 the question about whether they have sample sizes or not while digitising. To expedite 342 343 finding and adding in these sample sizes to do the necessary calculations (if for example a figure presented 95% CI's or standard errors), metaDigitise() has s specific edit 344 option that allows users to enter in previously omitted sample sizes. It works by first 345 346 identifying the missing sample sizes in the digitised output, re-plotting the relevant figure and then prompting the user to enter the sample sizes for the relevant groups in 347 348 the figure, one by one. As an example, assume that we were missing sample sizes for two groups in 002_Doe_2013_Fig1.png: 349

```
filename
                       group_id
                                        variable
                                                   mean error error_type n
                                                                                        plot_type
                                  Sepal length
002_Doe_2013_Fig1.png
                          setosa
                                                    5.00 0.11 se
                                                                         NA
                                                                                       mean_error
002_Doe_2013_Fig1.png viriginica Sepal length
                                                    6.59 0.18 se
                                                                         NA
                                                                            NA
                                                                                   NA mean_error
```

Here, we can see that we are missing the sample sizes for setosa and viriginica, and as a result, sd is not calculated because metaDigitise() needs this information to make the calculation. If the user found this information after contacting the authors for clarification then they can add these back in as follows:

```
R> digitised_data <- metaDigitise(".../FiguresToExtract")
Do you want to...</pre>
```

- 1: Process new images
- 2: Import existing data

3: Edit existing data Selection: R> 3 Choose how you want to edit files: 1: Cycle through images 2: Choose specific file to edit 3: Enter previously omitted sample sizes Selection: >R 3 metaDigitise() will replot the figure after this and list, only the groups missing data, for which the user can then update the data. This is then re-integrated back into the data automatically and the sd calculated. Group " setosa ": Enter sample size R> 50 Group " viriginica ": Enter sample size

355

356

R> 50

357 Inter-observer Variability and Validation

358 Inter-observer variability in digitisations

359 In order to evaluate the consistency of digitisation using **metaDigitise** between users, we 360 simulated a dataset of two traits with two different groups. These data were then used to construct plots of the four different types (scatterplot, mean and error, histogram 361 and boxplots). Each variable was plotted twice for each given plot type (figures were 362 modified slightly to give users a sense that they were digitising new data) generating a 363 total of 14 figures. 14 independent digitisers were provided with a directory with all 14 364 365 figures in a randomised order. Digitisers ran **metaDigistise** on their own computers, across different operating systems (including Mac, Windows and Linux). Digitisers 366 varied in their level of experience, from people with experience of meta-analyses or 367 comparative work to those without any science background. We asked users to digitise 368 all 14 figures and collected the mean, standard deviation and correlation coefficient (for 369 370 scatterplots) generated by metaDigitise() for every plot digitised. We transformed 371 these data to standardized differences as

$$\frac{\theta - \hat{\theta}}{\hat{\theta}} \tag{8}$$

where θ is the estimate value and $\hat{\theta}$ is the true value, meaning that deviations were 372 373 percentage differences from the true summary statistics. The correlation coefficient 374 deviation was not divided by the true value, as it is already on a standardised scale. This deviation can be seen as a measure of bias. The resulting data was used to assess 375 between- and within- user variability (i.e., the intra-class correlation coefficient) in the 376 377 data. This was done using linear mixed effect models with user identify as a random effect. Standardised mean, standard deviation and correlation coeficients were used as 378 response variables in seperate models. Sampling variance for ICC estiamtes was 379

- 380 generated based on 1000 parametric bootstraps of the model and the significance was
- 381 tested using liklihood ratio tests. These models were run using the lme4 (Bates et al.,
- 382 2015) and **rptR** (Stoffel, Nakagawa & Schielzeth, 2017) packages in R.
- 383 If digitisations were consistent across all users then we should find no significant
- 384 between user variability in the data. Indeed, across plot types we found no evidence for
- any inter-observer variability in digitisations for the mean (ICC = 0, 95% CI = 0 to
- 386 0.029, p = 1), standard deviation (ICC = 0, 95% CI = 0 to 0.033, p = 0.5) or
- 387 correlation coefficient (ICC = 0.053, 95% CI = 0 to 0.296, p = 0.377). There were was
- 388 little bias between digitised and true values, on average 1.63% (mean = 0.02%, SD =
- 389 4.9\%, r = -0.03%) and overall there were only small absolute differences between
- 390 digitised and true values, deviating, on average 2.18% (mean = 0.40%, SD = 5.81%, r
- 391 = 0.33%) across all three summary statistics.
- 392 SD estimates from digitisations are clearly more prone to error than means or
- 393 correlation coefficients. If the mean absoluate difference is calculated for each plot type,
- 394 we can see that this effect is driven mainly by extraction from boxplots and histgrams
- 395 (% difference):

- 396 This is because SD estimation from the summary statistics extracted from boxplots is
- 397 more error prone, especially at small sample sizes (Wan et al., 2014).

398 Testing the accuracy of digitisations

- 399 To test how accurate **metaDigitise** is at matching points to their true values, we
- 400 generated four random scatterplots, each with 20 data points, and digitised these with
- 401 metaDigitise(). This was done by one digitiser, as there is no detectable between user
- 402 variation. Data digitised using **metaDigitise** was essentially perfectly correlated with

403 the true simulated data for both the x-variable (Pearson's correlation; r = 0.9999915, t = 2137.4, df = 78, $p \neq 0.001$) and y-variable (r = 0.9999892, t = 1897.8, df = 78, $p \neq 0.001$).

406 Discussion and Conclusions

Although **metaDigitise** is already very flexible, and provides functionality not seen in 407 any other package (Table 1) it is clear that there are some functions that it does not 408 perform. A notable feature that **metaDigitise** lacks is automated point detection. Point 409 detection is available in several packages (Table 1). However, from our experience of 410 using these functions, manual digitising is more reliable and often equally as fast. 411 412Particularly given that calibration (for point detection) needs to be done for each plot 413 individually in any case. Additionally, auto-detection often misses many points which then subsequently need to be manually added. Based on tests of metaDigitise (see 414 above), figures can be extracted in around 1-2 minutes, including the entry of metadata. 415 As a result, we do not belive that current automated point detection provides 416417 substantial benefits in terms of time or accuracy. Another feature that **metaDigitise** (currently) lacks, is an ability to zoom in on plots. 418 Zooming may enable users to gain greater accuracy when clicking on points. However, 419420 from our own experience (and indeed from the results above), if you are using a 421 reasonably sized screen then the accuracy is already high from these programs, and there is not much gain to be had from zooming in on points in many 422 423 circumstances. In contrast to some other packages, **metaDigitise** currently also does not extract lines 424 from figures. In our own experience, line extraction is not particularly useful for 425meta-analysis, although we recognise that it may be useful in other fields. Should a user 426 like to extract lines with **metaDigitise**, we would recommend extracting data as a 427

Function	metaDigitise	$GraphClick^1$	$DataThief^2$	$DigitizeIt^3$	$WebPlotDigitizer^4$	$ m metagear^5$	$\operatorname{digitize}^6$
Scatterplots	>	>	>	>	>	7.	>
Mean and error plots	>	>	>	×	×	77	×
Boxplots	>	×	×	×	×	×	×
Histograms	>	×	×	×	~ 7	×	×
Graph rotation ⁸	>	>	>	>	>	×	×
Groups	>	>	×	>	>	×	×
Entry of metadata	>	×	×	×	×	×	×
Summarising data	>	×	×	×	×	×	×
Multiple image processing	>	×	×	×	×	×	×
${ m Reproducable}^9$	>	>	>	×	>	×	×
Automated point detection	×	>	×	>	>	>	×
Line extraction	×	>	>	>	>	×	×
Zoom	×	>	>	>	>	×	×
Log axis	×	>	>	>	>	×	×
Dates	×	×	>	×	>	×	×
Asymmetric error bars	×	×	>	×	×	×	×
Freeware	\checkmark 10	\checkmark 11	\checkmark^{11}	\times^{11}	\checkmark^{11}	\checkmark 10	\checkmark 10
1 A C. C. C. C (9000) 2 T (9000) 3 T	(900e) 3 D	: (9019) 4 D.L	1,5 (2017) 5 1	(9016)	2) 6 Delegat (2011)		

 $^{^{1}}$ Arizona-Software (2008) 2 Tummers (2006) 3 Bormann (2012) 4 Rohatgi (2017) 5 Lajeunesse (2016) 6 Poisot (2011)

Table 1: Comparison of functionality between different digitisation softwares.

 $^{^{7}}$ Only automated, no manual extraction.

 $^{^8}$ Or handles rotated graphs.

 $^{^{9}}$ Allows saving, re-plotting and editing of data extraction.

¹⁰ R package.

¹¹ Standalone software.

scatter plot, and clicking along the line in question. A model can then be fitted to these 428points (setting the argument "summary = FALSE" in **metaDigitise** - will provide access 429 to the processed data) to estimate the parameters needed. 430 Finally, metaDigitise currently does not allow for asymmetric error bars. At present 431 this is a deliberate omission, as it is not clear how best to derive SD from such data, 432 433 given also that such asymmetric error bars may represent different things in different 434 figures. 435 Descriptve statistics are usually the most robust sources of information for calculating effect size statistics (Noble et al., 2017). These are most often presented in figures. 436 Users may therefore also want to compare effect size estimates from inferential statistics 437 438 with those derived from descriptive statistics (obtained for example using metaDigitise) from a paper. Comparing these different effects sizes can be useful in identifying 439 uncertainties and problems within a paper. In the future, we hope to provide functions 440 to easily convert inferential statistics to standardised effect size estimates, which can 441 seamlessly be integrated with summary statistics from metaDigitise, to calculate 442 equivalent standardised effect size estimates and their sampling variance. 443 Increasing the reproducibility of figure extraction for meta-analysis and making this 444 laborious process more streamlined, flexible and integrated with existing statistical 445 software will go a long way in facilitating the production of high quality meta-analytic 446 studies that can be updated in the future. We believe that **metaDigitise** will improve 447 this research synthesis pipeline, and will hopefully become an integral package that can 448

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449

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