figure.s3

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2025-05-20

```
#Figure S3
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
             1.1.4
## v dplyr
                        v readr
                                   2.1.5
## v forcats 1.0.0
                        v stringr
                                   1.5.1
## v ggplot2 3.5.2
                        v tibble
                                   3.2.1
## v lubridate 1.9.4
                     v tidyr
                                   1.3.1
## v purrr
              1.0.4
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become
library(reshape2)
##
## 载入程序包: 'reshape2'
## The following object is masked from 'package:tidyr':
##
##
      smiths
library(data.table)
##
## 载入程序包: 'data.table'
##
## The following objects are masked from 'package:reshape2':
##
##
      dcast, melt
```

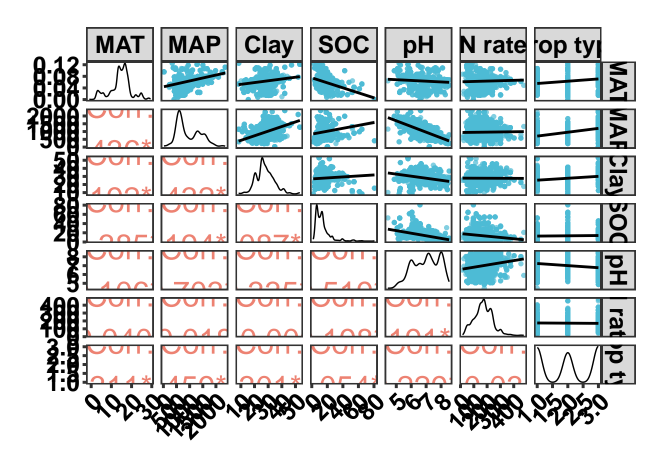
```
##
## The following objects are masked from 'package:lubridate':
##
      hour, isoweek, mday, minute, month, quarter, second, wday, week,
##
##
       yday, year
##
## The following objects are masked from 'package:dplyr':
##
##
       between, first, last
##
## The following object is masked from 'package:purrr':
##
##
      transpose
library(metafor)
## 载入需要的程序包: Matrix
##
## 载入程序包: 'Matrix'
##
## The following objects are masked from 'package:tidyr':
##
##
       expand, pack, unpack
##
## 载入需要的程序包: metadat
## 载入需要的程序包: numDeriv
##
## Loading the 'metafor' package (version 4.8-0). For an
## introduction to the package please type: help(metafor)
library(metagear)
## ** metagear 0.7, for installing/troubleshooting help see:
         http://lajeunesse.myweb.usf.edu/metagear/metagear_basic_vignette.html
## **** External dependencies check:
## **** setup supports GUIs [ TRUE ]
## **** setup supports data extraction from plots/figures [ FALSE ]
              NOTE: EBImage package (Bioconductor) will be installed only
## ****
## ****
                    when a figure_* function is used.
library(GGally)
```

Registered S3 method overwritten by 'GGally':

```
##
    method from
           ggplot2
##
    +.gg
# read data
d1 <- readxl::read_xlsx('D:/date/homework/Source Data.xlsx',sheet = "FigureS3")</pre>
d1 <- as.data.table(d1)</pre>
d2 < -d1
CV_nuet_bar<-mean(d2\$nuet_sd[is.na(d2\$nuet_sd)==FALSE]/d2\$nuet_mean[is.na(d2\$nuet_sd)==FALSE])
d2\$nuet_sd[is.na(d2\$nuet_sd)==TRUE] <-d2\$nuet_mean[is.na(d2\$nuet_sd)==TRUE] *1.25*CV_nuet_bar
d2\$nuec\_sd[is.na(d2\$nuec\_sd) == TRUE] < -d2\$nuec\_mean[is.na(d2\$nuec\_sd) == TRUE] *1.25*CV\_nuec\_bar
# clean up column names
d2 <- as.data.table(d2)</pre>
setnames(d2,gsub('\\/','_',gsub(' |\\(|\\)','',colnames(d2))))
setnames(d2,tolower(colnames(d2)))
#Supplement and update the missing values for n_dose and p_dose
d2[is.na(n_dose), n_dose := median(d2$n_dose,na.rm=TRUE)]
# update the database (g_crop_type)
d2[g_crop_type=='maize', g_crop_type := 1]
d2[g_crop_type=='wheat', g_crop_type := 2]
d2[g_crop_type=='rice', g_crop_type := 3]
#Conversion of factored data to numeric data
d2$g_crop_type <- as.numeric(d2$g_crop_type)</pre>
str(d2)
## Classes 'data.table' and 'data.frame':
                                          2436 obs. of 15 variables:
                : num 1 2 2 2 2 3 3 4 4 4 ...
## $ studyid
## $ mat
                : num 14.2 14.2 14.2 14.2 14.2 ...
                : num 495 495 495 495 ...
## $ map
                : num 26.7 26.7 26.7 26.7 26.7 ...
## $ clay
## $ soc
                : num 11.5 11.5 11.5 11.5 11.5 ...
                : num 7.79 7.79 7.79 7.79 ...
## $ ph
## $ g_crop_type: num 2 1 1 1 1 1 1 1 1 ...
## $ management : chr "EE" "CF" "CF" "RES" ...
                : num 120 150 150 150 150 337 349 150 150 150 ...
## $ n_dose
```

```
## $ replication: num 2 3 3 3 4 4 3 3 3 ...
                 : chr "REN" "REN" "REN" "REN" ...
## $ nue_type
## $ nuet_mean : num 78.3 35.7 25.3 35.7 43.9 ...
                 : num 10.17 4.64 3.29 4.64 5.7 ...
## $ nuet sd
## $ nuec_mean : num 72.5 30.6 30.6 30.6 30.6 ...
## $ nuec_sd
                 : num 9.89 4.17 4.17 4.17 4.17 ...
## - attr(*, ".internal.selfref")=<externalptr>
## - attr(*, "index")= int(0)
# update the database (tillage)
d2[management=='ROT', management := 7]
d2[management=='CC', management := 8]
d2[management=='RES', management := 9]
d2[management=='RFR', management := 10]
d2[management=='RFP', management := 11]
d2[management=='RFT', management := 12]
d2[management=='CF', management := 13]
d2[management=='OF', management := 14]
d2[management=='RT', management := 15]
d2[management=='NT', management := 16]
d2[management=='EE', management := 17]
#Conversion of factored data to numeric data
d2$management <- as.numeric(d2$management)</pre>
# Estimate meta-analytical response measure (ROM Method)
# calculate effect size (NUE)
es21 <- escalc(measure = "ROM", data = d2,
               m1i = nuet_mean, sd1i = nuet_sd, n1i = replication,
               m2i = nuec_mean, sd2i = nuec_sd, n2i = replication )
# make forest plots per group treatments
# convert to data.tables
d02 <- as.data.table(es21)</pre>
d3 <- d02[,c("mat", "map", "clay", "soc", "ph", "n_dose", "g_crop_type")] #Extraction correlation of
#Standardize data to prevent large gaps in values
```

```
df=scale(d3[,1:7],center=TRUE,scale=TRUE)
#Change column name
names(d3)<-c("MAT", "MAP", "Clay", "SOC", "pH", "N rate", "Crop type")</pre>
head(d3)
##
           TAM
                                       SOC
                                                 pH N rate Crop type
                     MAP
                            Clay
##
         <num>
                   <num>
                            <num>
                                     <num>
                                                    <num>
                                                               <num>
## 1: 14.23171 494.9222 26.72143 11.50714 7.792857
                                                                   2
## 2: 14.23171 494.9222 26.72143 11.50714 7.792857
                                                       150
                                                                   1
## 3: 14.23171 494.9222 26.72143 11.50714 7.792857
                                                       150
                                                                   1
## 4: 14.23171 494.9222 26.72143 11.50714 7.792857
                                                       150
                                                                   1
## 5: 14.23171 494.9222 26.72143 11.50714 7.792857
                                                       150
                                                                   1
## 6: 17.87691 1281.7686 34.41429 9.05000 5.700000
                                                       337
                                                                   1
str(d3)
## Classes 'data.table' and 'data.frame': 2436 obs. of 7 variables:
   $ MAT
               : num 14.2 14.2 14.2 14.2 14.2 ...
##
   $ MAP
               : num 495 495 495 495 ...
               : num 26.7 26.7 26.7 26.7 26.7 ...
## $ Clay
## $ SOC
               : num 11.5 11.5 11.5 11.5 11.5 ...
## $ pH
               : num 7.79 7.79 7.79 7.79 ...
## $ N rate
               : num 120 150 150 150 150 337 349 150 150 150 ...
## $ Crop type: num 2 1 1 1 1 1 1 1 1 ...
## - attr(*, "digits")= Named num [1:9] 4 4 4 4 4 4 4 4 4
   ..- attr(*, "names")= chr [1:9] "est" "se" "test" "pval" ...
##
## - attr(*, "yi.names")= chr "yi"
## - attr(*, "vi.names")= chr "vi"
## - attr(*, ".internal.selfref")=<externalptr>
#Calculate correlation coefficients and scatter plots and fit linearity
ggpairs(d3, lower = list(continuous = wrap("cor", size = 8,color="#E64B35B2")),
        upper = list(continuous = wrap("smooth", size =1.2,color="#4DBBD5B2")))+
 theme_bw(base_line_size = 1.05,base_rect_size = 1.05)+
 theme(panel.grid.major=element_blank()),panel.grid.minor=element_blank())+
 theme(axis.text=element_text(colour='black',size=18, face="bold"), strip.text = element_text(col
 theme(axis.text.x = element_text(angle = 45, hjust = 1, vjust = 1))
```



ggsave(file = "D:/date/homework/picture/Figure_S3.png", width = 410, height = 297, units = "mm")

R Markdown

This is an R Markdown document. Markdown is a simple formatting syntax for authoring HTML, PDF, and MS Word documents. For more details on using R Markdown see http://rmarkdown.rstudio.com.

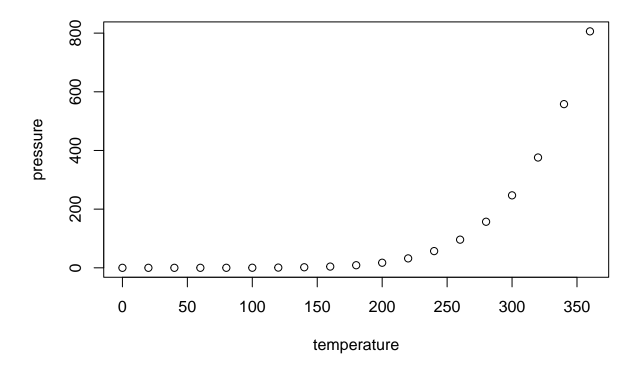
When you click the **Knit** button a document will be generated that includes both content as well as the output of any embedded R code chunks within the document. You can embed an R code chunk like this:

summary(cars)

```
##
        speed
                         dist
            : 4.0
                               2.00
    Min.
                    Min.
##
                    1st Qu.: 26.00
    1st Qu.:12.0
##
##
    Median:15.0
                    Median : 36.00
            :15.4
                            : 42.98
##
    Mean
                    Mean
    3rd Qu.:19.0
                    3rd Qu.: 56.00
##
##
    Max.
            :25.0
                            :120.00
                    Max.
```

Including Plots

You can also embed plots, for example:



Note that the echo = FALSE parameter was added to the code chunk to prevent printing of the R code that generated the plot.