SOM - Read and data plotting from the manuscript Paixao et al. 2021 - JASr

Paixão et al.2021 - JARs

28/09/2020

Brief description of the script

This R markdown document reads, summarizes and plots data for the manuscript Paixão et al. 2021 Using Mechanical experiments to study Ground Stone Tool use: exploring the formation of percussive and grinding wear traces on Limestone tools. Journal of Archaeological Science: Reports

The document contains:

- 1. Manuscript tables
- 2. Manuscript figures (data analysis)
- 3. Supplementary material, including extra tables and figures (data analysis)

This R project and respective scripts follow the procedures described by Marwick et al. 2017.

To compile this markdown document do not delete or move files from their original folders. Please note that most of the tables and figures in this file do not match the numbering in the original manuscript.

For any questions, comments and inputs, please contact:

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Load data into R project

```
Imported files are in: '../analysis/raw_data'
Figures are saved in: '../analysis/plots'
Tables are saved in: '../analysis/derived_data'
```

Load libraries

-- Attaching packages ------ tidyverse 1.3.0 -## v ggplot2 3.3.3 v purrr 0.3.4 ## v tibble 3.0.6 v dplyr 1.0.4 ## v tidyr 1.1.2 v stringr 1.4.0 ## v readr 1.4.0 v forcats 0.5.1

```
## -- Conflicts -----
                                          ## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
library(utils)
library(knitr)
library(janitor)
##
## Attaching package: 'janitor'
## The following objects are masked from 'package:stats':
##
##
      chisq.test, fisher.test
library(kableExtra)
##
## Attaching package: 'kableExtra'
## The following object is masked from 'package:dplyr':
##
##
      group_rows
library(GGally)
## Registered S3 method overwritten by 'GGally':
    method from
##
    +.gg
           ggplot2
library(doBy)
##
## Attaching package: 'doBy'
## The following object is masked from 'package:dplyr':
##
##
      order_by
library(ggpubr)
library(ggfortify)
library(tools)
```

Import datasets

```
gisdata <- read_csv("../raw_data/gisdata.csv")
```

```
##
## -- Column specification ---------
    sample = col_character(),
##
##
    cycle = col_character(),
    parameter = col character(),
##
    motion = col character(),
    material = col_character(),
##
##
    id = col_double(),
##
    elev_min = col_double(),
    elev_max = col_double(),
##
    nparts = col_double(),
##
    npoints = col_double();
##
    perimeter = col_double(),
##
    area = col_double()
## )
confocaldata <- read.csv("../raw_data/confocaldata.csv", na.strings = "*****", encoding = "UTF-8")
data_file <- list.files("../raw_data/", pattern = "\\.csv$", full.names = TRUE)</pre>
md5_in <- md5sum(data_file)</pre>
info_in <- data.frame(file = basename(names(md5_in)), checksum = md5_in, row.names = NULL)
```

In this study two datasets are used:

1) $\mathbf{gisdata.csv}$: dataset for the QGIS analysis

```
str(gisdata)
```

```
## spec_tbl_df [355 x 12] (S3: spec_tbl_df/tbl_df/tbl/data.frame)
## $ sample : chr [1:355] "id2-5" "id2-5" "id3-3" "id3-3" ...
## $ cycle : chr [1:355] "before" "before" "before" "before" ...
## $ parameter: chr [1:355] "tri" "tri" "tri" "tri" ...
## $ motion : chr [1:355] "Impact" "Impact" "Impact" "Impact" ...
## $ material : chr [1:355] "Flint" "Flint" "Flint" "Flint" ...
             : num [1:355] 0 1 0 1 2 3 4 0 1 0 ...
   $ elev_min : num [1:355] 0 0.01 0 0.01 0.02 0.03 0.04 0 0.01 0 ...
## $ elev_max : num [1:355] 0.01 0.02 0.01 0.02 0.03 0.04 0.05 0.01 0.02 0.01 ...
             : num [1:355] 1007 1010 131 165 47 ...
## $ npoints : num [1:355] 67487 59648 14961 16143 3142 ...
   $ perimeter: num [1:355] 485.2 446 160.6 177.5 32.8 ...
              : num [1:355] 204.49 17.48 83.65 38.12 1.83 ...
   - attr(*, "spec")=
##
    .. cols(
##
    . .
         sample = col_character(),
##
    .. cycle = col_character(),
    .. parameter = col_character(),
##
         motion = col_character(),
##
##
    .. material = col_character(),
##
    .. id = col_double(),
##
    .. elev_min = col_double(),
##
    .. elev_max = col_double(),
```

```
## .. nparts = col_double(),
## .. npoints = col_double(),
## .. perimeter = col_double(),
## .. area = col_double()
## .. )
```

2) confocaldata.csv: dataset for the Confocal microscopy surface texture analysis analysis

str(confocaldata)

```
## 'data.frame':
                   25 obs. of 54 variables:
## $ Name
                                               "Lime2-5_LSM_50x075_suf1_Topo > Leveled (LS-plane) > For
## $ Created.on
                                               "6/24/2020 12:03:05 PM" "6/24/2020 12:21:59 PM" "6/24/2
                                        : chr
   $ sample
                                               "id2-5" "id2-5" "id3-3" ...
                                        : chr
                                               "impact" "impact" "impact" ...
## $ motion
                                        : chr
## $ workedmaterial
                                               "flint" "flint" "flint" "...
                                        : chr
                                               "Surface" "Surface" "Surface" ...
## $ Studiable.type
                                        : chr
   $ Axis.name...X
                                        : chr
                                               "X" "X" "X" "X" ...
## $ Axis.length...X
                                               255 255 255 255 255 ...
                                        : num
                                               3000 3000 3000 1024 1024 3000 3000 1024 3000 3000 ...
## $ Axis.size...X
                                        : int
## $ Axis.spacing...X
                                               85.2 85.2 85.2 249.6 249.6 ...
                                        : num
                                               "Y" "Y" "Y" "Y" ...
## $ Axis.name...Y
                                        : chr
## $ Axis.length...Y
                                       : num
                                               255 255 255 255 ...
                                               3000 3000 3000 1024 1024 3000 3000 1024 3000 3000 ...
## $ Axis.size...Y
                                        : int
## $ Axis.spacing...Y
                                        : num
                                               85.2 85.2 85.2 249.6 249.6 ...
                                               "Z" "Z" "Z" "Z" ...
## $ Axis.name...Z
                                       : chr
                                               "Topography" "Topography" "Topography" "Topography" ...
## $ Layer.type...Z
                                       : chr
## $ Axis.length...Z
                                       : num 8.3 30.6 14.9 48.6 23 ...
##
   $ Axis.size...Z
                                               128164 333950 130813 193492 260079 181249 180484 428496
                                        : int
                                       : num 0.0647 0.0916 0.1139 0.2511 0.0884 ...
## $ Axis.spacing...Z
## $ NM.points.ratio...Z
                                       : int 0000000000...
                                              0.836 4.619 2.321 5.348 3.491 ...
## $ Sq
                                        : num
## $ Ssk
                                        : num
                                              0.997 0.107 0.142 -0.491 0.201 ...
## $ Sku
                                        : num 8.63 4.48 3.11 5.97 3.46 ...
## $ Sp
                                        : num 4.8 15.02 7.46 23 11.88 ...
## $ Sv
                                               3.49 15.58 7.45 25.58 11.11 ...
                                        : num
## $ Sz
                                        : num 8.3 30.6 14.9 48.6 23 ...
## $ Sa
                                        : num 0.572 3.244 1.819 3.887 2.699 ...
## $ Smr
                                        : num 0.464 0.497 0.448 0.18 0.354 ...
                                              0.775 5.691 2.944 5.799 4.436 ...
## $ Smc
## $ Sxp
                                        : num 1.53 10.61 4.3 11.65 6.74 ...
                                        : num 13.5 18.9 20.1 18.7 22.9 ...
## $ Sal
## $ Str
                                        : num 0.371 0.416 0.592 0.468 0.803 ...
## $ Std
                                               149 65 150 51 124 ...
## $ Sdq
                                        : num 0.328 1.153 0.688 1.126 0.897 ...
## $ Sdr
                                        : num 4.36 20.02 16.47 31.23 24.6 ...
## $ Vm
                                        : num   0.0866   0.3378   0.1331   0.3133   0.2114   ...
## $ Vv
                                        : num 0.861 6.029 3.078 6.113 4.648 ...
## $ Vmp
                                        : num    0.0866    0.3378    0.1331    0.3133    0.2114    ...
## $ Vmc
                                        : num 0.528 3.111 2.097 3.932 3.146 ...
## $ Vvc
                                        : num 0.769 5.335 2.824 5.33 4.303 ...
##
   $ Vvv
                                        : num 0.0923 0.694 0.2534 0.7826 0.3444 ...
## $ Maximum.depth.of.furrows
                                        : num 4.56 20.63 7.65 25.68 13.88 ...
```

```
## $ Mean.depth.of.furrows
                                     : num 0.962 4.63 2.49 5.112 3.932 ...
## $ Mean.density.of.furrows
                                     : num 4523 3830 4509 2286 2201 ...
## $ First.direction
                                      : num 1.50e+02 6.36e+01 2.66e-03 9.00e+01 9.00e+01 ...
                                      : num 180 45 154 45 135 ...
## $ Second.direction
## $ Third.direction
                                      : num 141.3 56.2 63.5 51.2 123.7 ...
## $ Isotropy
                                      : num 23.5 26.2 77.8 33.1 73.8 ...
## $ Lengthscale.anisotropy.Sfrax.epLsar: num NA NA NA 0.000493 0.001218 ...
## $ Lengthscale.anisotropy.NewEplsar : num NA NA NA 0.0177 0.0172 ...
## $ Fractal.complexity.Asfc
                                       : num 8.66 23.18 30.55 37.79 44.11 ...
## $ Scale.of.max.complexity.Smfc
                                     : num 1.71e+06 1.81e+08 2.71e+06 1.17e+01 1.52e+01 ...
## $ HAsfc9
                                       : num 0.449 2.496 0.388 0.544 0.254 ...
## $ HAsfc81
                                       : num 0.659 3.446 0.481 0.701 0.487 ...
```

GIS analysis, Terrain analysis for Slope and TRI based on the 3D surface point clouds

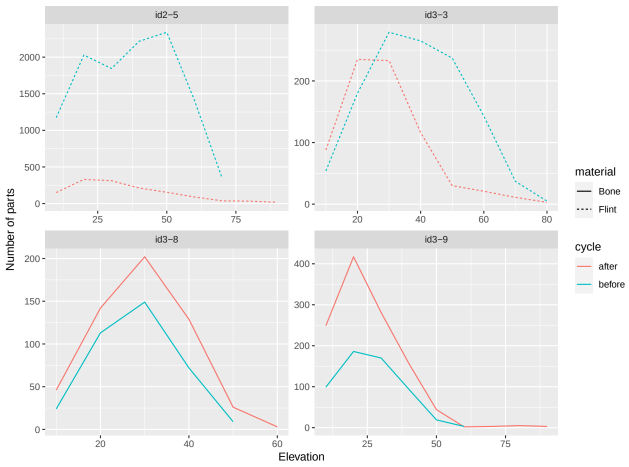
Slope

```
# Compute proportions for perimeter and area grouped by sample and GIS parameter
slope <- filter(gisdata, parameter == "slope")</pre>
slopebefore <- filter(slope, cycle == "before")</pre>
slopeafter <- filter(slope, cycle == "after")</pre>
# before experimental cycles (i.e. natural surfaces)
id2.5before <- filter(slopebefore, sample == "id2-5")</pre>
id3.3before <- filter(slopebefore, sample == "id3-3")
id3.8before <- filter(slopebefore, sample == "id3-8")
id3.9before <- filter(slopebefore, sample == "id3-9")
id6.1before <- filter(slopebefore, sample == "id6-1")</pre>
id6.3before <- filter(slopebefore, sample == "id6-3")</pre>
id6.6before <- filter(slopebefore, sample == "id6-6")</pre>
id6.7before <- filter(slopebefore, sample == "id6-7")</pre>
id2.5before <- id2.5before %>%
 group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.3before <- id3.3before %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.8before <- id3.8before %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.9before <- id3.9before %>%
  group_by(sample) %>%
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.1before <- id6.1before %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
```

```
id6.3before <- id6.3before %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.6before <- id6.6before %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.7before <- id6.7before %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
# after experimental cycles
id2.5after <- filter(slopeafter, sample == "id2-5")
id3.3after <- filter(slopeafter, sample == "id3-3")</pre>
id3.8after <- filter(slopeafter, sample == "id3-8")</pre>
id3.9after <- filter(slopeafter, sample == "id3-9")</pre>
id6.1after <- filter(slopeafter, sample == "id6-1")
id6.3after <- filter(slopeafter, sample == "id6-3")</pre>
id6.6after <- filter(slopeafter, sample == "id6-6")</pre>
id6.7after <- filter(slopeafter, sample == "id6-7")</pre>
id2.5after <- id2.5after %>%
 group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.3after <- id3.3after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.8after <- id3.8after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.9after <- id3.9after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
```

```
id6.1after <- id6.1after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.3after <- id6.3after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.6after <- id6.6after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.7after <- id6.7after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
newslope <- do.call("rbind", list(id2.5before, id3.3before, id3.8before, id3.9before, id6.1before, id6.
# save outputs
write_csv(newslope,"../derived_data/newslope.csv")
# Plot data
# Number of parts
impactdf <- filter(newslope, motion == "Impact")</pre>
grinding <- filter(newslope, motion == "Grinding")</pre>
slopepartsexp_impac <- ggplot(impactdf, aes(x = elev_max, y = nparts, colour = cycle)) +</pre>
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("Slope impact experiment, number of parts") +
    ylab("Number of parts") +
  xlab("Elevation")
slopepartsexp_impac
```

Slope impact experiment, number of parts

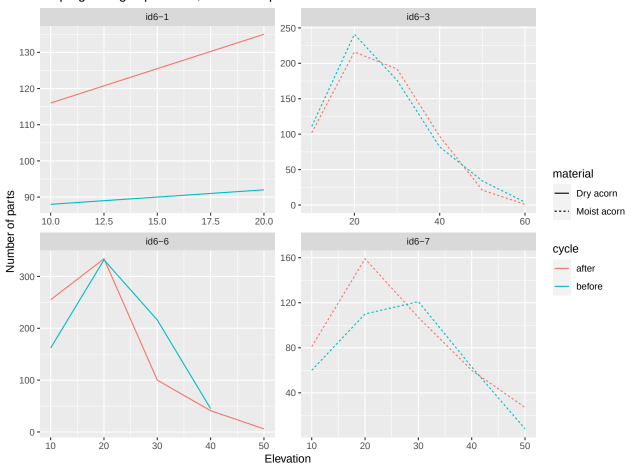


ggsave("../plots/slopepartsexp_impac.png")

```
slopepartsexp_grind <- ggplot(grinding, aes(x = elev_max, y = nparts, colour = cycle)) +
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("Slope grinding experiment, number of parts") +
    ylab("Number of parts") +
    xlab("Elevation")

slopepartsexp_grind</pre>
```

Slope grinding experiment, number of parts



ggsave("../plots/slopepartsexp_grind.png")

Saving 8.5 x 6.5 in image

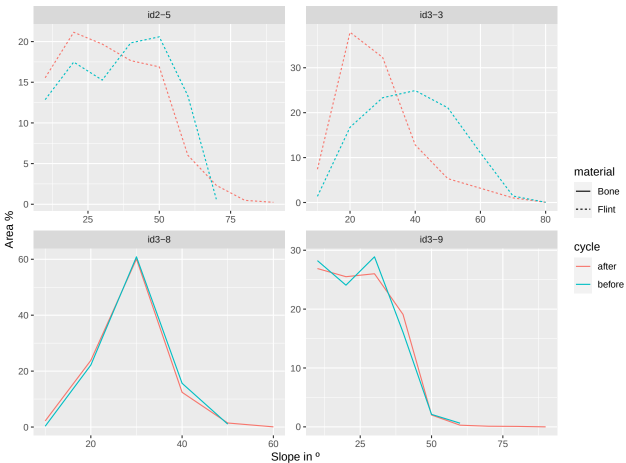
```
# Area %

impactdf <- filter(newslope, motion == "Impact")
grinding <- filter(newslope, motion == "Grinding")

areaimpact <- ggplot(impactdf, aes(x = elev_max, y = areaperc, colour = cycle)) +
    geom_line(aes(linetype = material)) +
    facet_wrap(~sample, scale = "free") +
    ggtitle("Slope analysis, impact") +
    ylab("Area %") +
    xlab("Slope in 0")

areaimpact</pre>
```

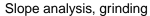
Slope analysis, impact

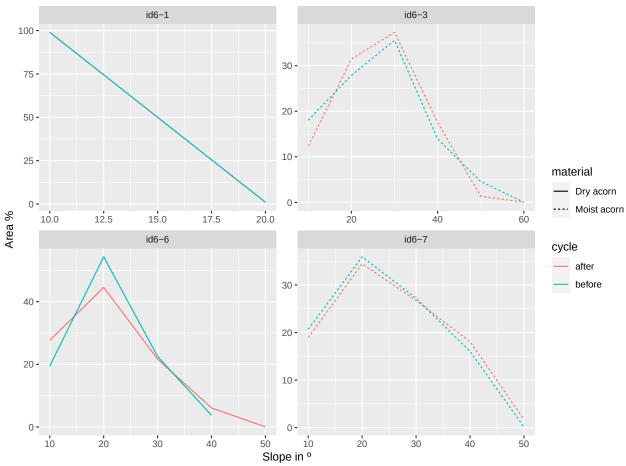


ggsave("../plots/slopeareaimpact.png")

```
areagrinding <- ggplot(grinding, aes(x = elev_max, y = areaperc, colour = cycle)) +
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("Slope analysis, grinding") +
  ylab("Area %") +
  xlab("Slope in º")

areagrinding</pre>
```



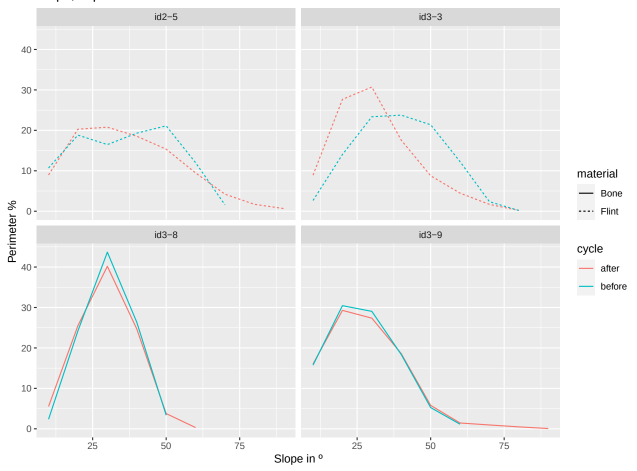


ggsave("../plots/slopeareagrinding.png")

```
# Perimeter %

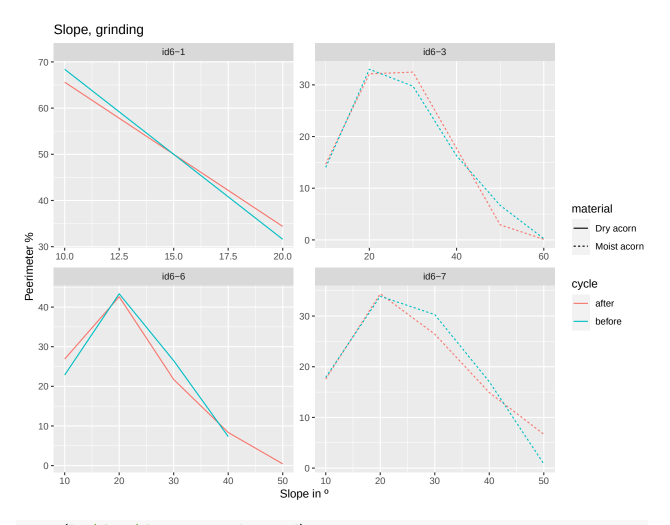
perimimpact <- ggplot(impactdf, aes(x = elev_max, y = perimperc, colour = cycle)) +
    geom_line(aes(linetype = material)) +
    facet_wrap(~sample) +
    ggtitle("Slope, impact") +
    ylab("Perimeter %") +
    xlab("Slope in 0")</pre>
```

Slope, impact



ggsave("../plots/slopeperimimpact.png")

```
perimgrinding <- ggplot(grinding, aes(x = elev_max, y = perimperc, colour = cycle)) +
   geom_line(aes(linetype = material)) +
   facet_wrap(~sample, scale = "free") +
   ggtitle("Slope, grinding") +
   ylab("Peerimeter %") +
   xlab("Slope in 0")</pre>
```



ggsave("../plots/slopeperimgrinding.png")

TRI (Terrain roughness index)

```
tri <- filter(gisdata, parameter == "tri")
tribefore <- filter(tri, cycle == "before")
triafter <- filter(tri, cycle =="after")

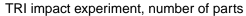
# before experimental cycles (i.e. natural surfaces)

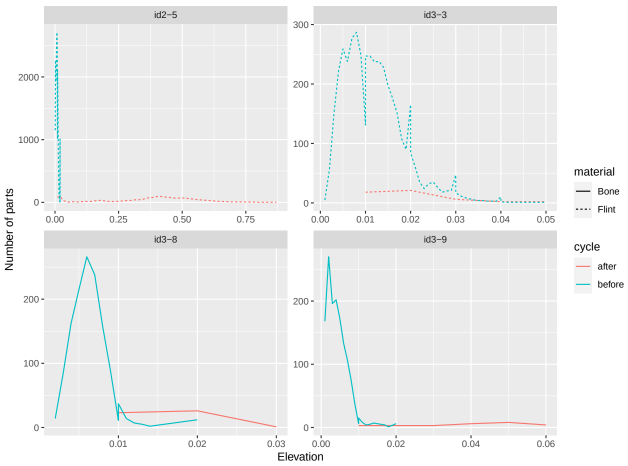
id2.5before <- filter(tribefore, sample == "id2-5")
id3.3before <- filter(tribefore, sample == "id3-3")
id3.8before <- filter(tribefore, sample == "id3-8")
id3.9before <- filter(tribefore, sample == "id3-9")
id6.1before <- filter(tribefore, sample == "id6-1")
id6.3before <- filter(tribefore, sample == "id6-3")
id6.6before <- filter(tribefore, sample == "id6-6")</pre>
```

```
id6.7before <- filter(tribefore, sample == "id6-7")</pre>
id2.5before <- id2.5before %>%
  group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id3.3before <- id3.3before %>%
  group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id3.8before <- id3.8before %>%
  group_by(sample) %>%
  mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id3.9before <- id3.9before %>%
  group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id6.1before <- id6.1before %>%
  group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id6.3before <- id6.3before %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id6.6before <- id6.6before %>%
  group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
id6.7before <- id6.7before %>%
 group_by(sample) %>%
 mutate(
   areaperc = area / sum(area) * 100,
   perimperc = perimeter / sum(perimeter) * 100)
# after experimental cycles
```

```
id2.5after <- filter(triafter, sample == "id2-5")</pre>
id3.3after <- filter(triafter, sample == "id3-3")
id3.8after <- filter(triafter, sample == "id3-8")</pre>
id3.9after <- filter(triafter, sample == "id3-9")
id6.1after <- filter(triafter, sample == "id6-1")</pre>
id6.3after <- filter(triafter, sample == "id6-3")</pre>
id6.6after <- filter(triafter, sample == "id6-6")</pre>
id6.7after <- filter(triafter, sample == "id6-7")</pre>
id2.5after <- id2.5after %>%
 group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.3after <- id3.3after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.8after <- id3.8after %>%
 group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id3.9after <- id3.9after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.1after <- id6.1after %>%
  group_by(sample) %>%
  mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.3after <- id6.3after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.6after <- id6.6after %>%
  group_by(sample) %>%
 mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
id6.7after <- id6.7after %>%
  group_by(sample) %>%
```

```
mutate(
    areaperc = area / sum(area) * 100,
    perimperc = perimeter / sum(perimeter) * 100)
newtri <- do.call("rbind", list(id2.5before, id3.3before, id3.8before, id3.9before, id6.1before, id6.3b
# save outputs
write_csv(newtri,"../derived_data/newtri.csv")
# Plot data
# Number of parts
# Motion
impactdf <- filter(newtri, motion == "Impact")</pre>
grinding <- filter(newtri, motion == "Grinding")</pre>
imapact_parts <- ggplot(impactdf, aes(x = elev_max, y = nparts, colour = cycle)) +</pre>
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("TRI impact experiment, number of parts") +
    ylab("Number of parts") +
  xlab("Elevation")
imapact_parts
```

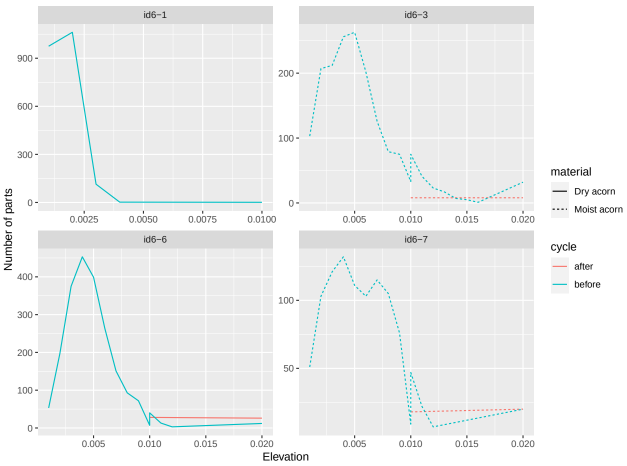




ggsave("../plots/tripartsexp_impac.png")

```
grinding_parts <- ggplot(grinding, aes(x = elev_max, y = nparts, colour = cycle)) +
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("TRI grinding experiment, number of parts") +
    ylab("Number of parts") +
    xlab("Elevation")</pre>
grinding_parts
```

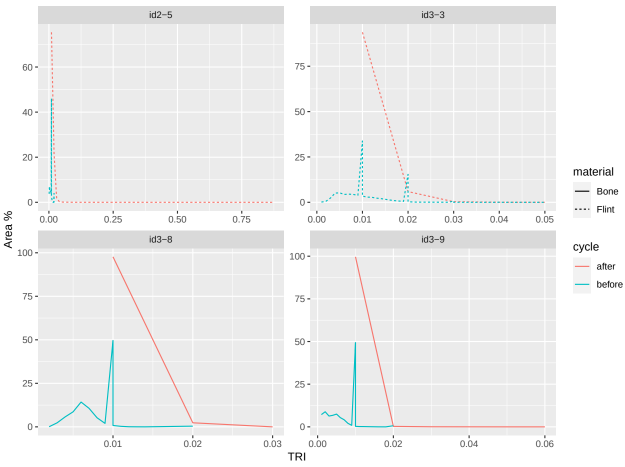




ggsave("../plots/tripartsexp_grind.png")

```
# Area %
areaimpact <- ggplot(impactdf, aes(x = elev_max, y = areaperc, colour = cycle)) +
    geom_line(aes(linetype = material)) +
    facet_wrap(~sample, scale = "free") +
    ggtitle("TRI analysis, impact") +
    ylab("Area %") +
    xlab("TRI")
areaimpact</pre>
```

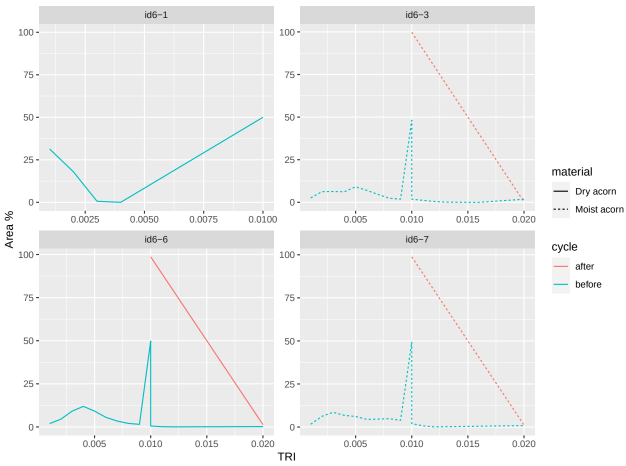
TRI analysis, impact



ggsave("../plots/triareaimpact.png")

```
areagrinding <- ggplot(grinding, aes(x = elev_max, y = areaperc, colour = cycle)) +
  geom_line(aes(linetype = material)) +
  facet_wrap(~sample, scale = "free") +
  ggtitle("TRI analysis, grinding") +
  ylab("Area %") +
  xlab("TRI")</pre>
areagrinding
```



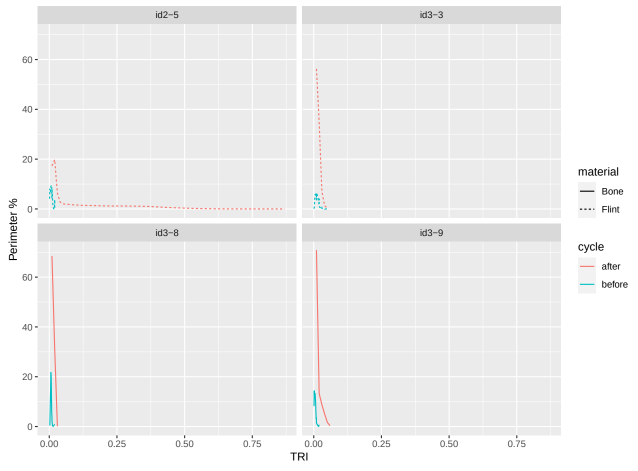


ggsave("../plots/triareagrinding.png")

```
# Perimeter %

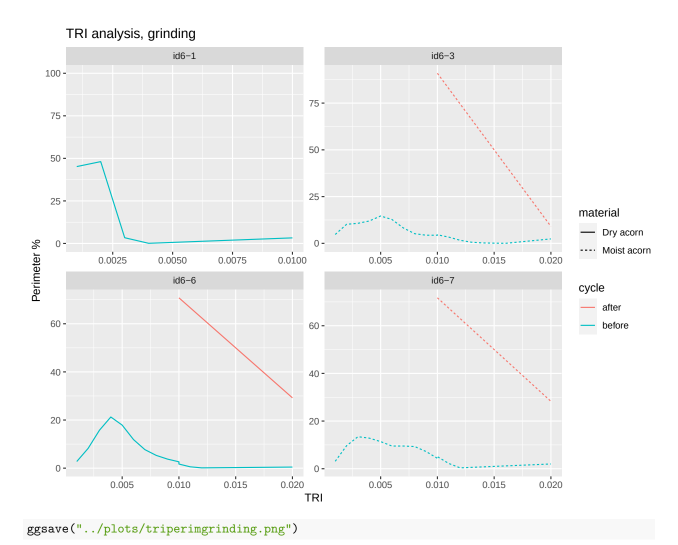
perimimpact <- ggplot(impactdf, aes(x = elev_max, y = perimperc, colour = cycle)) +
    geom_line(aes(linetype = material)) +
    facet_wrap(~sample) +
    ggtitle("TRI analysis, impact") +
    ylab("Perimeter %") +
    xlab("TRI")</pre>
```

TRI analysis, impact



ggsave("../plots/triperimimpact.png")

```
perimgrinding <- ggplot(grinding, aes(x = elev_max, y = perimperc, colour = cycle)) +
   geom_line(aes(linetype = material)) +
   facet_wrap(~sample, scale = "free") +
   ggtitle("TRI analysis, grinding") +
   ylab("Perimeter %") +
   xlab("TRI")</pre>
```



Saving 8.5 x 6.5 in image

Confocal micro surface texture data

Import and summarize data

```
# compute descriptive statistics

nminmaxmeanmedsd <- function(x){
    y <- x[!is.na(x)]
    n_test <- length(y)
    min_test <- min(y)
    max_test <- max(y)
    mean_test <- mean(y)
    med_test <- median(y)
    sd_test <- sd(y)
    out <- c(n_test, min_test, max_test, mean_test, med_test, sd_test)
    names(out) <- c("n", "min", "max", "mean", "median", "sd")</pre>
```

```
return(out)
}
num.var <- 21:length(confocaldata)

confostatsexp <- summaryBy(.~sample + motion + workedmaterial, data=confocaldata[c("sample", "motion","

## Warning in min(y): no non-missing arguments to min; returning Inf

## Warning in max(y): no non-missing arguments to max; returning -Inf

## Warning in min(y): no non-missing arguments to min; returning Inf

## Warning in max(y): no non-missing arguments to max; returning -Inf

## Warning in min(y): no non-missing arguments to min; returning Inf

## Warning in max(y): no non-missing arguments to max; returning -Inf

## Warning in min(y): no non-missing arguments to min; returning Inf

## Warning in max(y): no non-missing arguments to max; returning -Inf

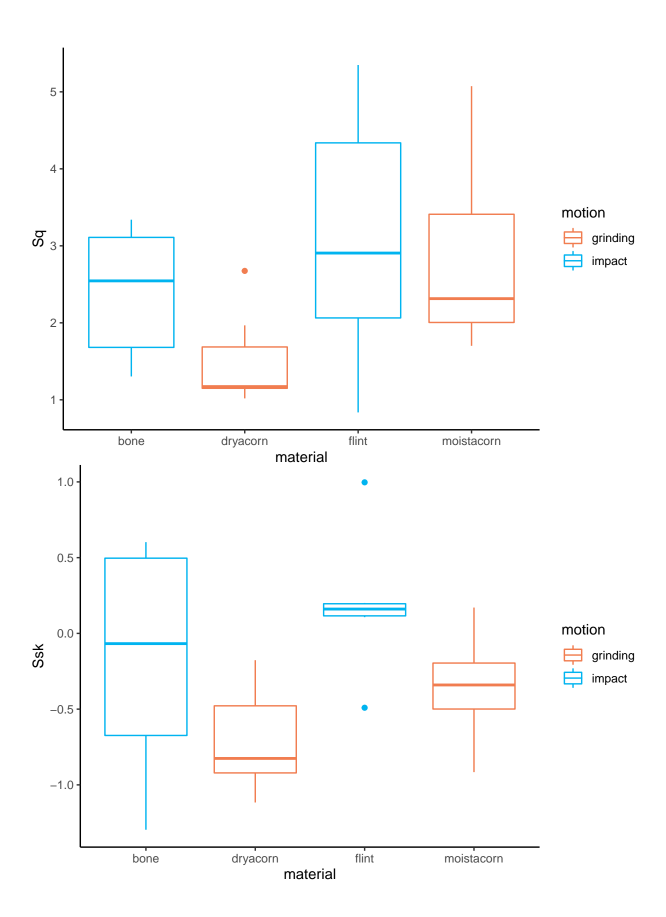
## Warning in max(y): no non-missing arguments to max; returning -Inf

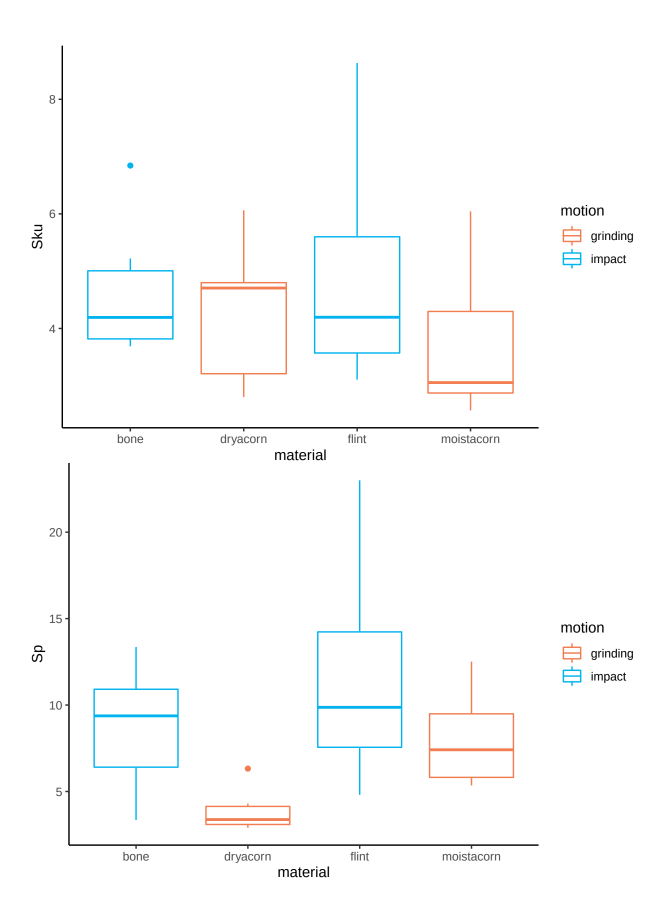
write_csv(confostatsexp, "../derived_data/confostats.csv")
```

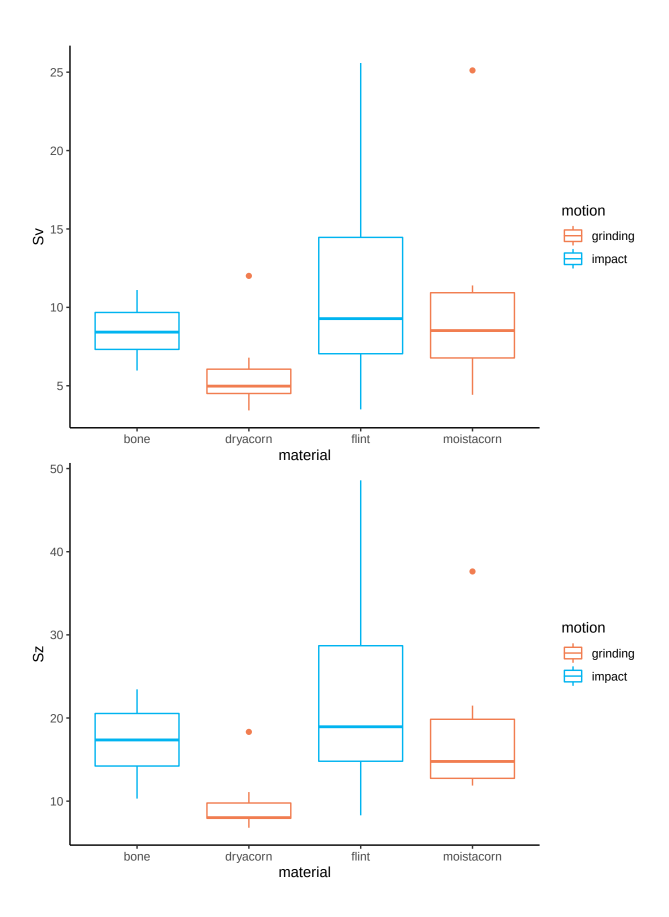
Plot all paramaters

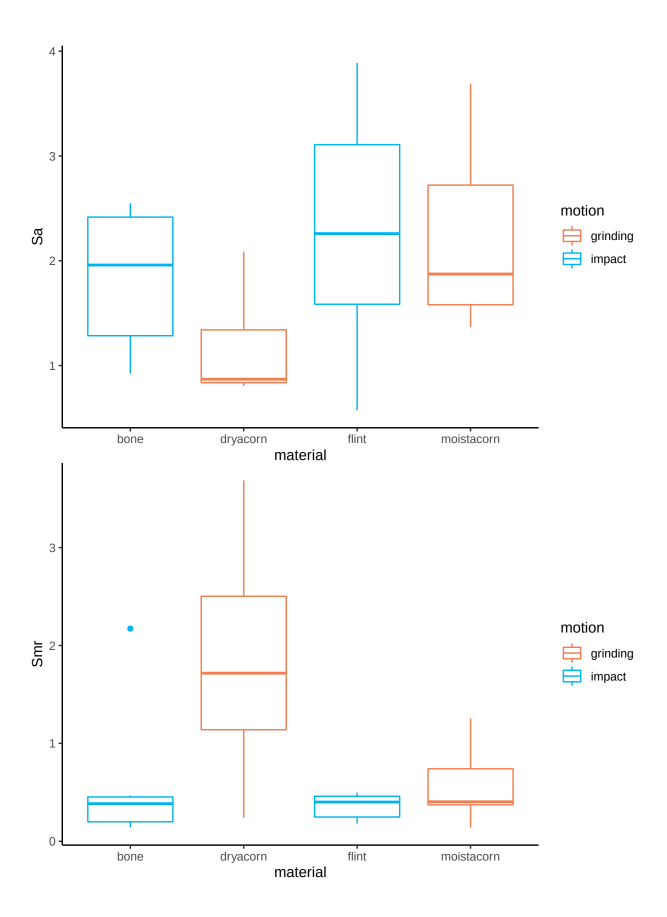
```
# Loop for plotting all surface texture parameters
for (i in num.var) cat("[",i,"] ", names(confocaldata)[i], "\n", sep = "")
## [21] Sq
## [22] Ssk
## [23] Sku
## [24] Sp
## [25] Sv
## [26] Sz
## [27] Sa
## [28] Smr
## [29] Smc
## [30] Sxp
## [31] Sal
## [32] Str
## [33] Std
## [34] Sdq
## [35] Sdr
## [36] Vm
## [37] Vv
```

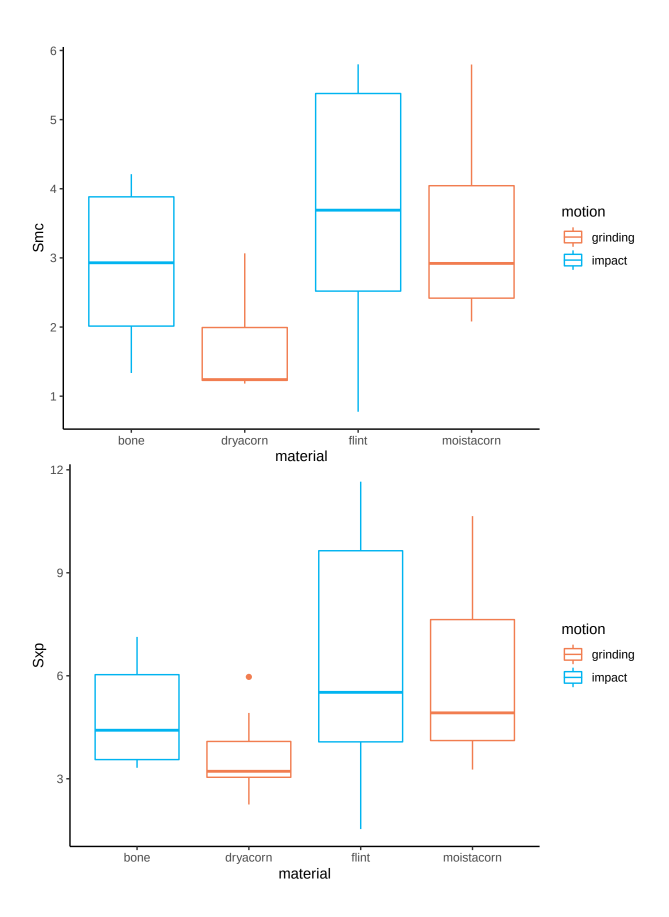
```
## [38] Vmp
## [39] Vmc
## [40] Vvc
## [41] Vvv
## [42] Maximum.depth.of.furrows
## [43] Mean.depth.of.furrows
## [44] Mean.density.of.furrows
## [45] First.direction
## [46] Second.direction
## [47] Third.direction
## [48] Isotropy
## [49] Lengthscale.anisotropy.Sfrax.epLsar
## [50] Lengthscale.anisotropy.NewEplsar
## [51] Fractal.complexity.Asfc
## [52] Scale.of.max.complexity.Smfc
## [53] HAsfc9
## [54] HAsfc81
for (i in num.var) {
  p <- ggplot(data = confocaldata, aes_string(x = "workedmaterial", y = names(confocaldata)[i],</pre>
                                             colour = "motion")) +
         geom_boxplot() +
         # geom_line(aes(group = motion)) +
         theme_classic() +
         labs(colour = "motion") +
         # facet_wrap(~ sample) +
         labs(x = "material", y = gsub("\\.", " ", names(confocaldata)[i])) +
         scale_colour_hue(h = c(25,225), limits = levels(confocaldata[["motion"]]))
  print(p)
  # saves the plots
  file_out <- pasteO(file_path_sans_ext(info_in[["file"]]), "_plot_",</pre>
                       names(confocaldata)[i], ".pdf")
    ggsave(filename = file_out, plot = p, path = "../plots", device = "pdf", width = 26,
           height = 21, units = "cm" )
}
```

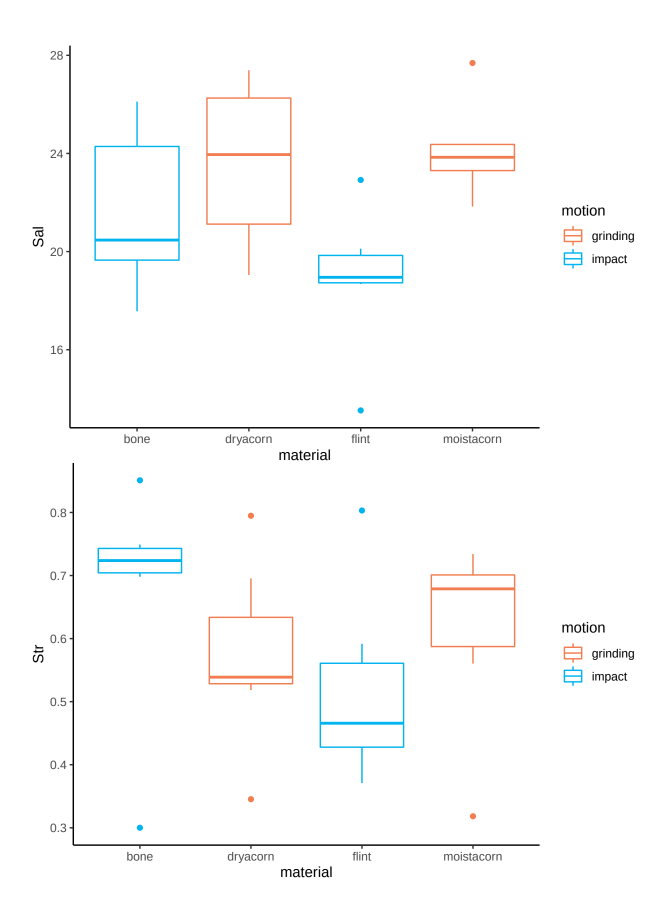


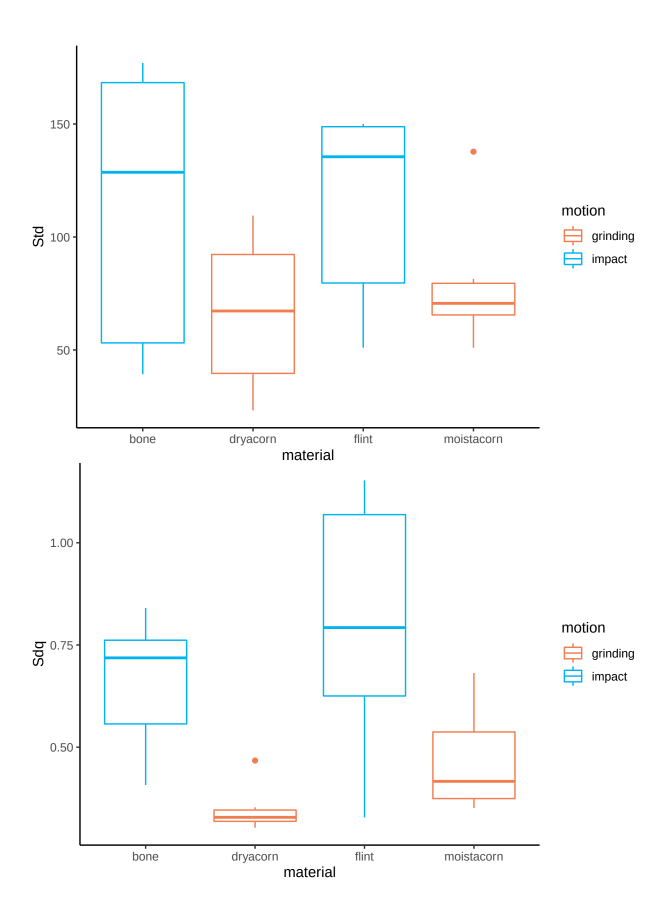


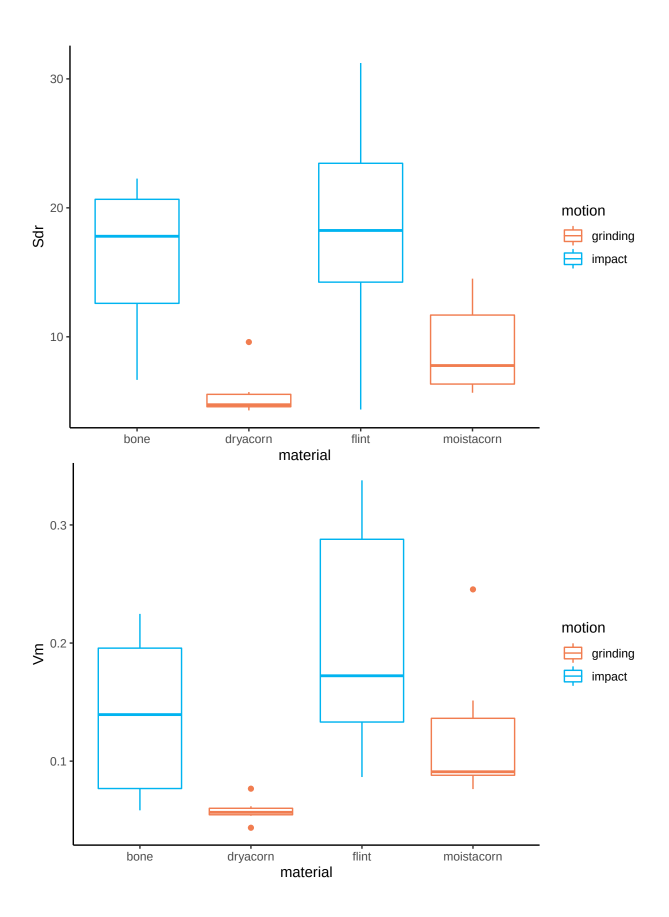


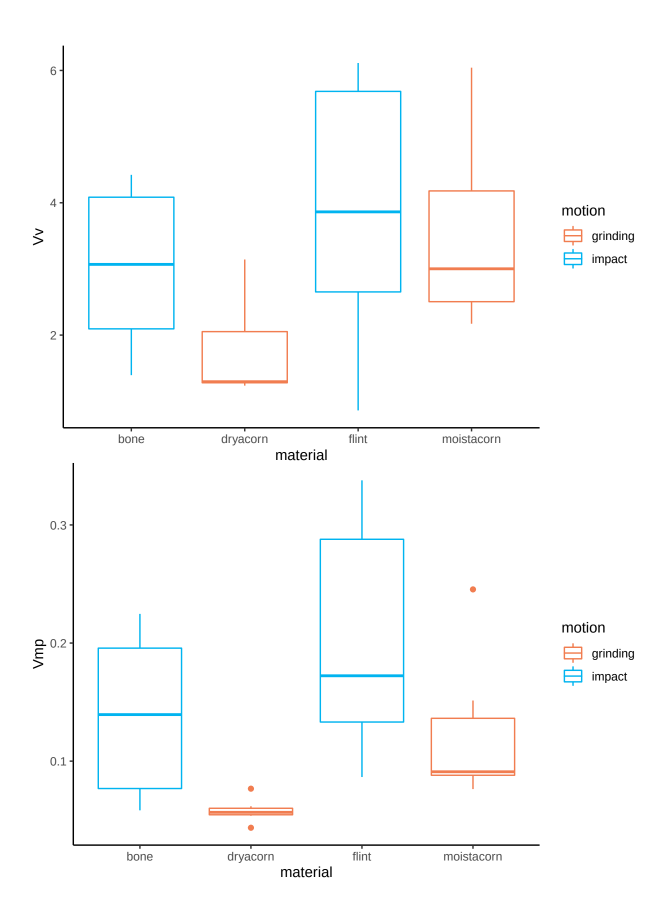


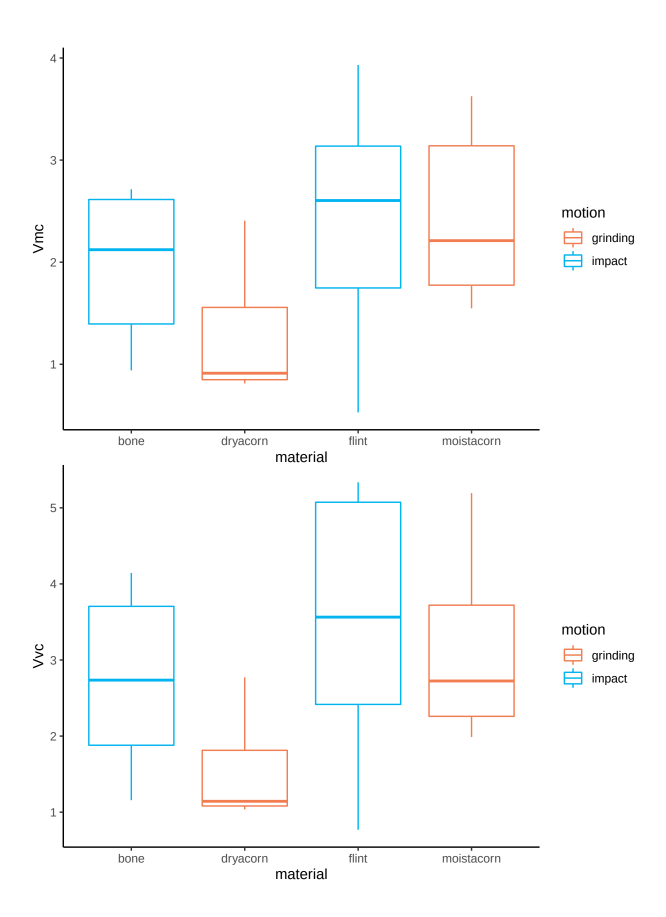


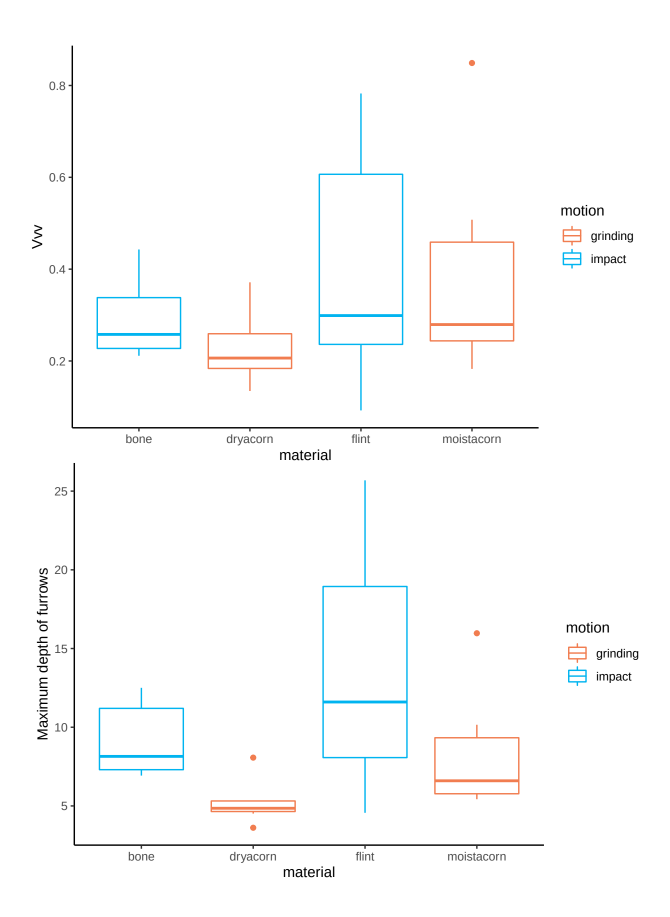


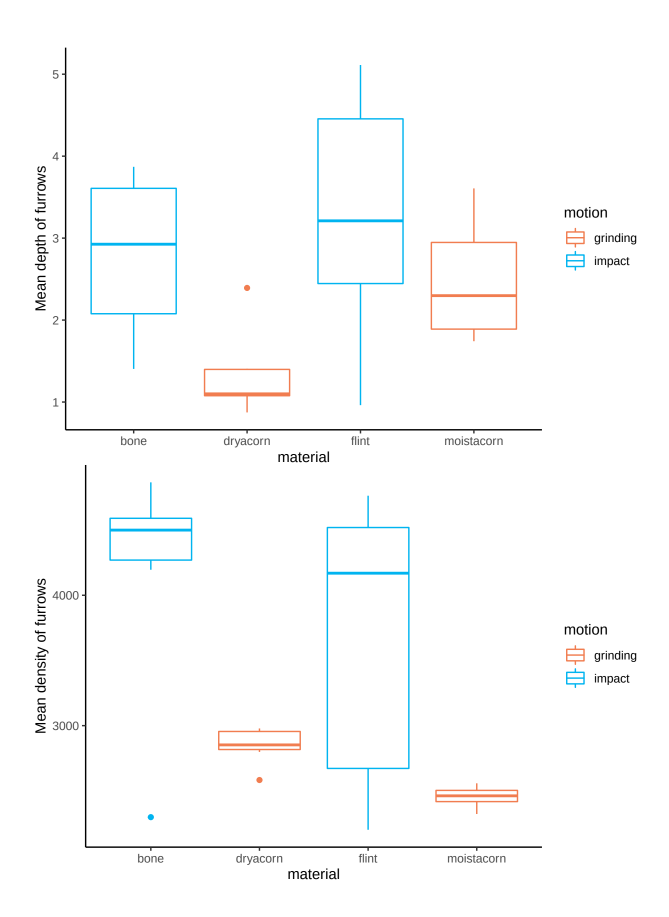


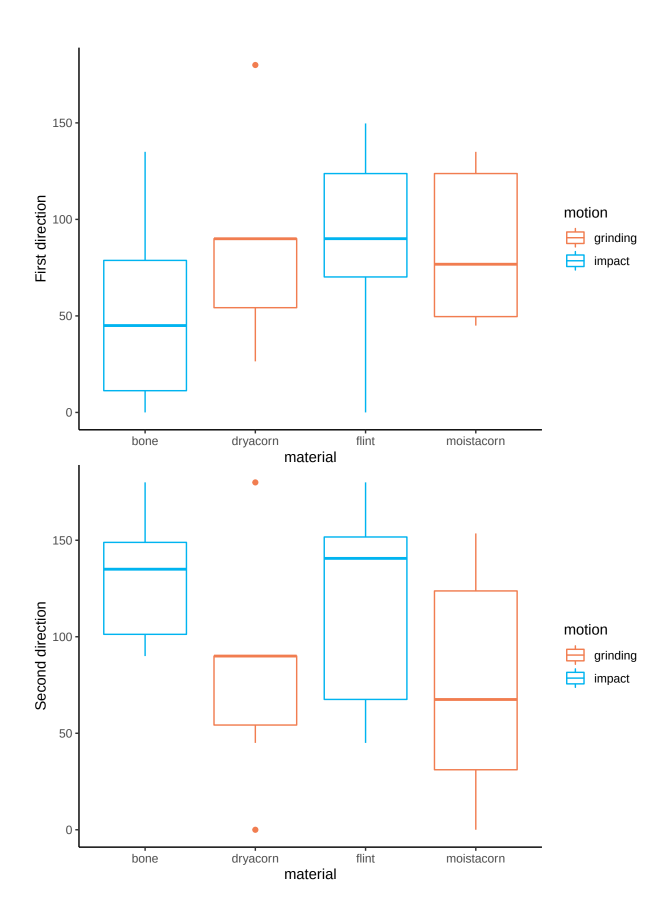


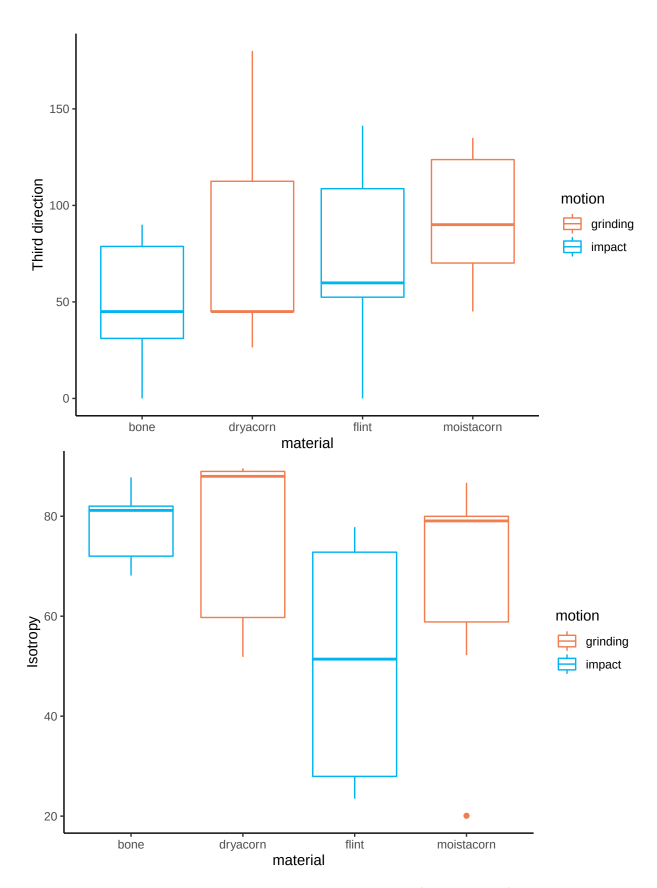






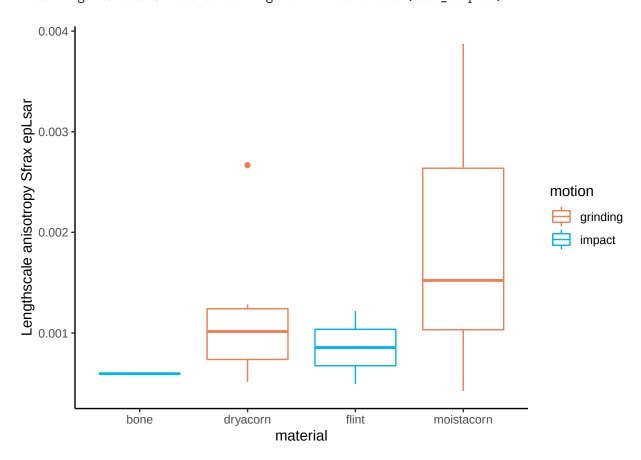




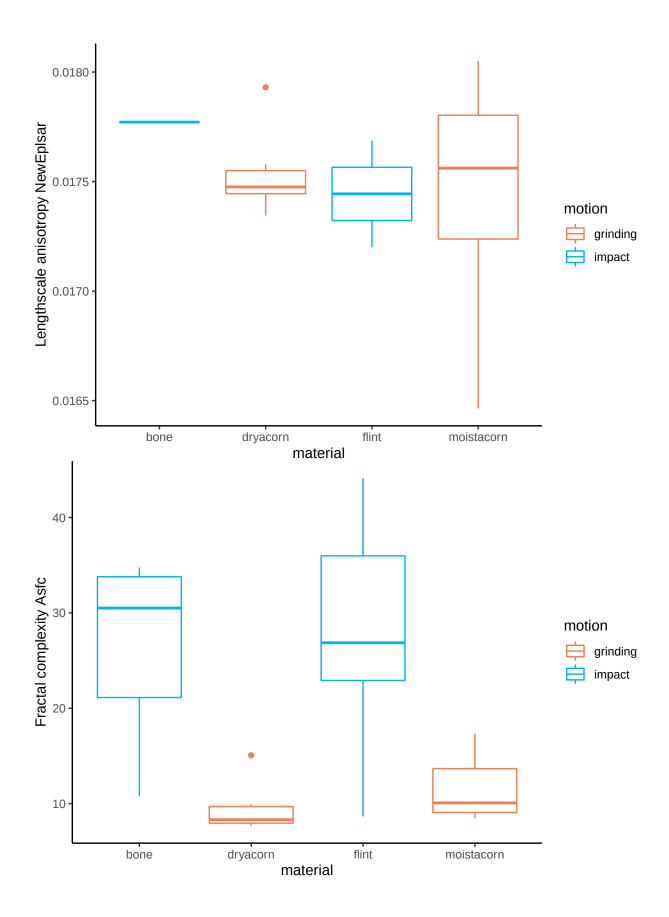


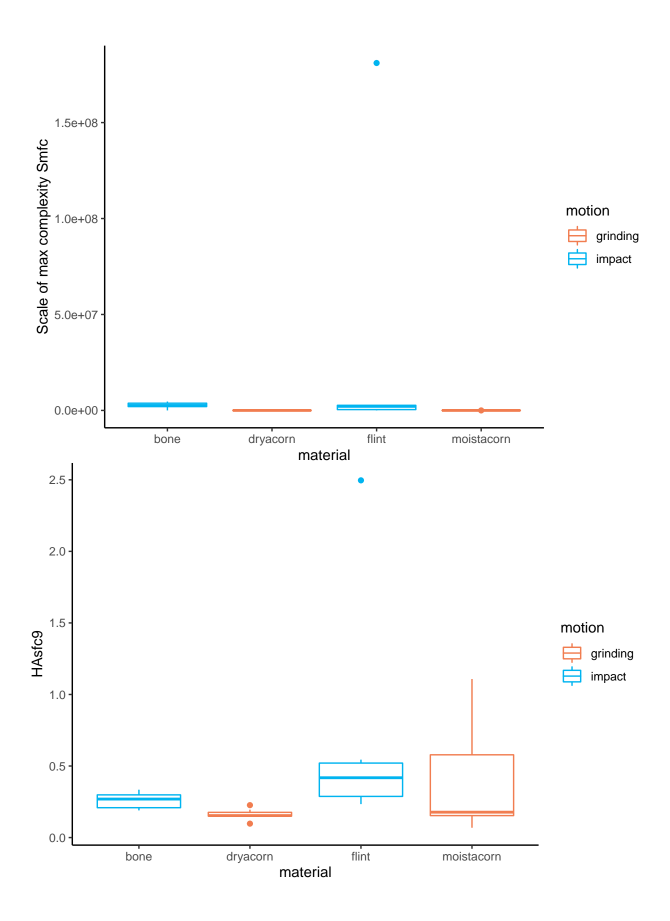
Warning: Removed 9 rows containing non-finite values (stat_boxplot).

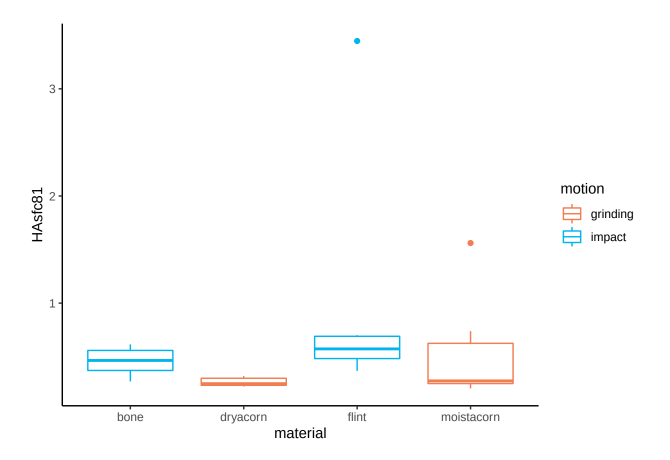
Warning: Removed 9 rows containing non-finite values (stat_boxplot).



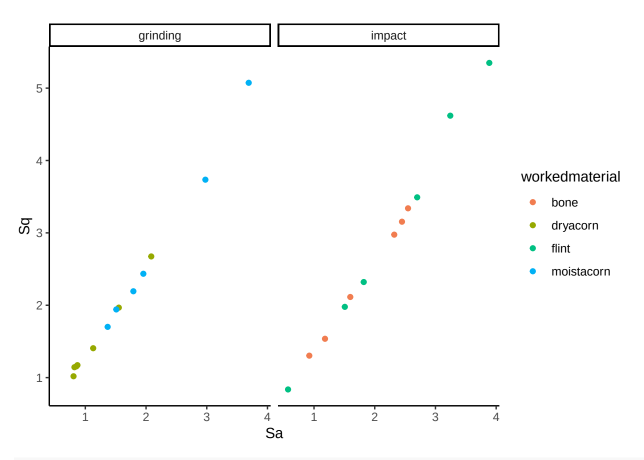
- ## Warning: Removed 9 rows containing non-finite values (stat_boxplot).
- ## Warning: Removed 9 rows containing non-finite values (stat_boxplot).





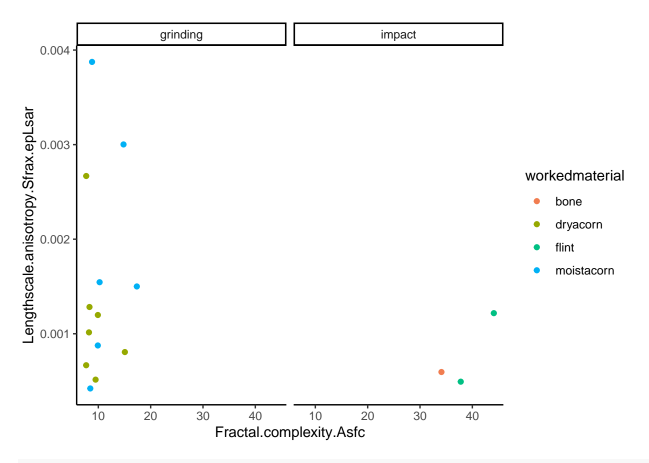


Scatterplots of selected variables combined by "Worked material" and "Motion"



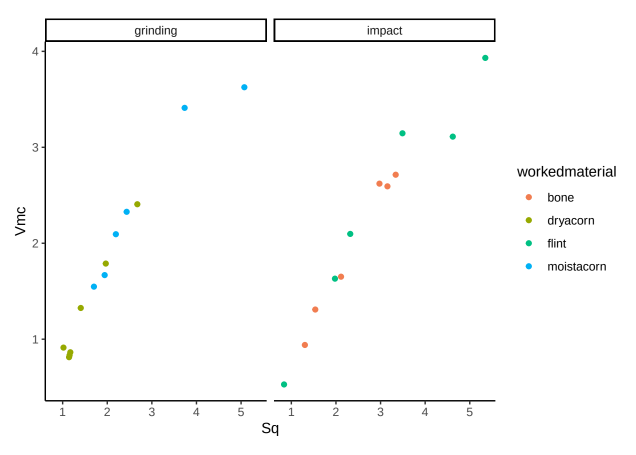
```
file_out <- paste0(file_path_sans_ext(info_in[["file"]]), "_scatterplot_Sa-Sq", ".pdf")
ggsave(filename = file_out, plot = Sa_Sq, path = "../plots", device = "pdf")</pre>
```

Warning: Removed 9 rows containing missing values (geom_point).



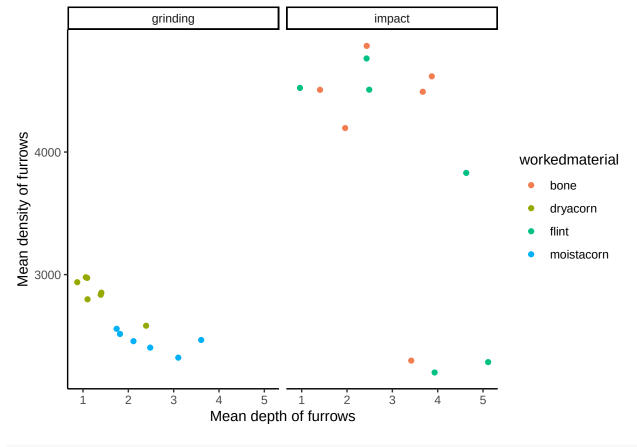
```
file_out <- paste0(file_path_sans_ext(info_in[["file"]]), "_scatterplot_Asfc-epLsar", ".pdf")
ggsave(filename = file_out, plot = ep_As, path = "../plots", device = "pdf")</pre>
```

Warning: Removed 9 rows containing missing values (geom_point).



```
file_out <- paste0(file_path_sans_ext(info_in[["file"]]), "_scatterplot_Sq-Vmc", ".pdf")
ggsave(filename = file_out, plot = Sq_Vmc, path = "../plots", device = "pdf")</pre>
```

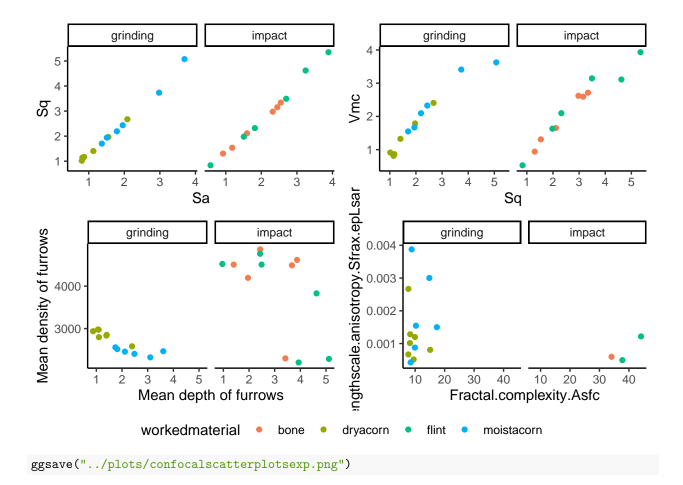
Saving 6.5 x 4.5 in image



```
file_out <- paste0(file_path_sans_ext(info_in[["file"]]), "_scatterplot_furrows", ".pdf")
ggsave(filename = file_out, plot = furrows, path = "../plots", device = "pdf")</pre>
```

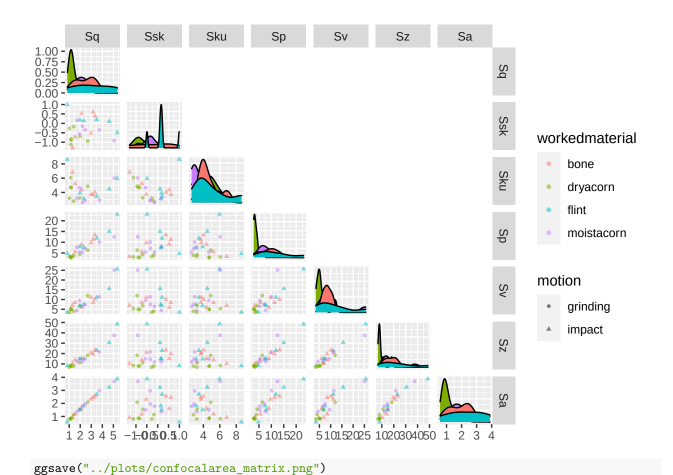
```
# combine all in a single image
ggarrange(Sa_Sq, Sq_Vmc, furrows, ep_As, common.legend = TRUE, legend = "bottom")
```

Warning: Removed 9 rows containing missing values (geom_point).

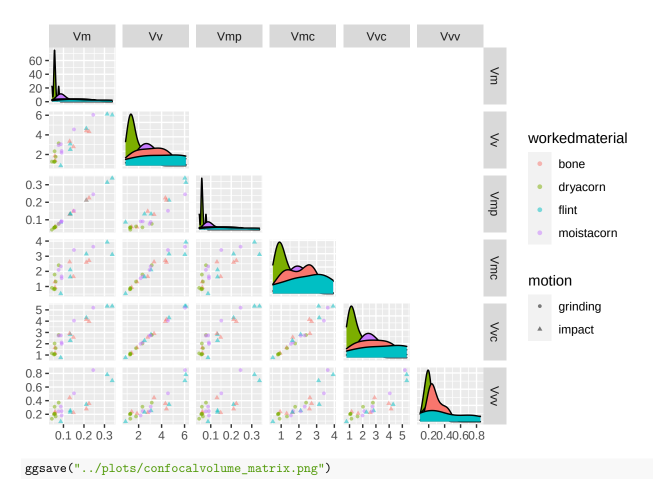


Saving 6.5 x 4.5 in image

Scatterplot matrix for the ISO 25178 Area scale, Height and volume parameters



Saving 6.5 x 4.5 in image



Saving 6.5 x 4.5 in image

Plot confostats for the ISO 25178 Area-scale, Height and volume parameters

```
p4 <- ggplot(heightconfostatsexp, aes(x=workedmaterial, y=Sp.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p5 <- ggplot(heightconfostatsexp, aes(x=workedmaterial, y=Sv.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p6 <- ggplot(heightconfostatsexp, aes(x=workedmaterial, y=Sz.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p7 <- ggplot(heightconfostatsexp, aes(x=workedmaterial, y=Sa.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
ggarrange(p1, p2, p3, p4, p5, p6, p7, common.legend = TRUE, font.label = list(size=8), legend="bottom")
                                   Ssk.mean
   3
                                       0.0 -
                                                                         3.5 -
                                      -1.0
       bone dryacorn flintmoistacorn
                                            bonedryacorn flintnoistacorn
                                                                              bonedryacorn flintmoistacorn
   12.5
Sp.mean
                                                                     Sz.mean
                                      12.5 -
   10.0
                                      10.0
    7.5
                                       7.5
    5.0
                                       5.0 -
    2.5
         bonedryacorn flintmoistacorn
                                            bonedryacorn flintmoistacorn
                                                                             bonedryacorn flintmoistacorn
  2.5
   2.0
   1.0
        bonedryacorn flintmoistacorn
                   Micro polish \rightleftharpoons bone \rightleftharpoons dryacorn \rightleftharpoons flint \rightleftharpoons
ggsave("../plots/confostatsarea_boxplots.png")
## Saving 6.5 \times 4.5 in image
```

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volumeconfostatsexp <- select(confostatsexp,sample,workedmaterial, Vm.mean,Vv.mean,Vmc.mean,Vvc.mean,V

Now, compute Volume parameters

```
p8 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vm.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p9 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vv.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p10 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vmp.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p11 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vmc.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p12 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vvc.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
p13 <- ggplot(volumeconfostatsexp, aes(x=workedmaterial, y=Vvv.mean, colour=workedmaterial)) +
  geom_boxplot() +
  labs(x="", colour="Micro polish")
ggarrange(p8, p9, p10, p11, p12, p13, common.legend = TRUE, font.label = list(size=8), legend="bottom")
   0.20
                                                                       0.20
                                                                    Vmp.mean
Vm.mean
                                  /v.mean
   0.15
                                                                       0.15 -
  0.10
                                                                       0.10
   0.05 -
                                                                       0.05 -
         bonedryacorn flintmoistacorn
                                         bone dryacorn flintmoistacorn
                                                                             bonedryacorn flintmoistacorn
   3.0 -
   2.5
                                  /vc.mean
                                                                    /vv.mean
   2.0
   1.5
   1.0 -
                                                                       0.2 -
        bonedryacorn flintmoistacorn
                                        bone dryacorn flintmoistacorn
                                                                            bonedryacorn flintmoistacorn
                  Micro polish \rightleftharpoons bone \rightleftharpoons dryacorn \rightleftharpoons flint \rightleftharpoons moistacorn
```

```
ggsave("../plots/confostatsvolume_boxplots.png")
```

Show plot files information

End and Session info

sessionInfo()

```
## R version 4.0.4 (2021-02-15)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Catalina 10.15.7
## Matrix products: default
          /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_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
## [1] tools
                 stats
                           graphics grDevices utils
                                                         datasets methods
## [8] base
##
## other attached packages:
## [1] ggfortify_0.4.11 ggpubr_0.4.0
                                          doBy_4.6.8
                                                           GGally_2.1.0
  [5] kableExtra_1.3.1 janitor_2.1.0
                                          knitr_1.31
                                                           forcats_0.5.1
  [9] stringr_1.4.0
                         dplyr_1.0.4
                                          purrr_0.3.4
                                                           readr_1.4.0
## [13] tidyr_1.1.2
                         tibble_3.0.6
                                          ggplot2_3.3.3
                                                           tidyverse_1.3.0
## loaded via a namespace (and not attached):
## [1] httr_1.4.2
                           jsonlite_1.7.2
                                              viridisLite_0.3.0 carData_3.0-4
   [5] modelr_0.1.8
                           assertthat_0.2.1
                                              highr_0.8
                                                                 cellranger_1.1.0
## [9] yaml_2.2.1
                           pillar_1.4.7
                                              backports_1.2.1
                                                                 lattice 0.20-41
## [13] glue_1.4.2
                           digest_0.6.27
                                              RColorBrewer_1.1-2 ggsignif_0.6.0
## [17] rvest_0.3.6
                           snakecase_0.11.0
                                              colorspace_2.0-0
                                                                 cowplot_1.1.1
## [21] htmltools_0.5.1.1 Matrix_1.3-2
                                              plyr_1.8.6
                                                                 pkgconfig_2.0.3
## [25] broom_0.7.4
                          haven_2.3.1
                                              scales_1.1.1
                                                                 webshot_0.5.2
## [29] openxlsx 4.2.3
                          rio 0.5.16
                                              farver 2.0.3
                                                                 generics 0.1.0
                                              withr_2.4.1
## [33] car_3.0-10
                           ellipsis_0.3.1
                                                                 cli_2.3.0
## [37] magrittr 2.0.1
                           crayon_1.4.0
                                              readxl_1.3.1
                                                                 evaluate 0.14
## [41] fs_1.5.0
                           MASS_7.3-53
                                              rstatix_0.6.0
                                                                 xm12_1.3.2
## [45] foreign_0.8-81
                           data.table_1.13.6 hms_1.0.0
                                                                 lifecycle_0.2.0
```

##	[49] munsell_0.5.0	reprex_1.0.0	zip_2.1.1	Deriv_4.1.2
##	[53] compiler_4.0.4	rlang_0.4.10	grid_4.0.4	rstudioapi_0.13
##	[57] labeling_0.4.2	rmarkdown_2.6	gtable_0.3.0	$abind_1.4-5$
##	[61] DBI_1.1.1	reshape_0.8.8	curl_4.3	R6_2.5.0
##	[65] gridExtra_2.3	<pre>lubridate_1.7.9.2</pre>	stringi_1.5.3	Rcpp_1.0.6
##	[69] vctrs_0.3.6	dbplyr_2.1.0	tidyselect_1.1.0	$xfun_0.20$