One-way row column design

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One-way ANOVA & pairwise comparison post hoc tests in a resolvable row column design.

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	<pre># (install &) load packages pacman::p_load(conflicted, desplot, emmeans, ggtext, lme4, lmerTest, multcomp, multcompView, tidyverse)</pre>										
	<pre># handle function conflicts conflicts_prefer(dplyr::filter)</pre>										

```
conflicts_prefer(dplyr::select)
conflicts_prefer(lmerTest::lmer)
```

1 Data

This example is taken from Chapter "3.10 Analysis of a resolvable row-column design" of the course material "Mixed models for metric data (3402-451)" by Prof. Dr. Hans-Peter Piepho. It considers data published in Kempton, Fox, and Cerezo (1996) from a yield trial laid out as a resolvable row-column design. The trial had 35 genotypes (gen), 2 complete replicates (rep) with 5 rows (row) and 7 columns (col). Thus, a complete replicate is subdivided into incomplete rows and columns.

1.1 Import

The data is available as part of the {agridat} package:

```
dat <- as_tibble(agridat::kempton.rowcol)</pre>
dat
# A tibble: 68 x 5
            row
   rep
                   col gen
                              yield
   <fct> <int> <int> <fct> <dbl>
                     1 G20
                               3.77
 1 R1
              1
                     2 G04
 2 R1
              1
                               3.21
 3 R1
              1
                     3 G33
                               4.55
 4 R1
              1
                     4 G28
                               4.09
 5 R1
              1
                     5 G07
                               5.05
 6 R.1
              1
                     6 G12
                               4.19
                     7 G30
                               3.27
 7 R1
              1
8 R1
              2
                     1 G10
                               3.44
              2
                               4.3
9 R1
                     2 G14
              2
                     4 G21
                               3.86
10 R1
# i 58 more rows
```

1.2 Format

For our analysis, gen, row and col should be encoded as factors. However, the desplot() function needs row and col as formatted as integers. Therefore we create copies of these columns encoded as factors and named rowF and colF:

```
dat <- dat %>%
  mutate(
    colF = as.factor(col),
    rowF = as.factor(row)
)
```

1.3 Explore

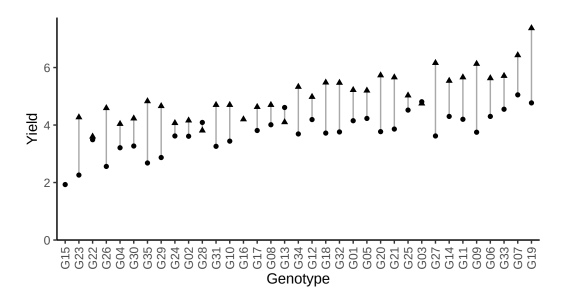
We make use of **dlookr::describe()** to conveniently obtain descriptive summary tables. Here, we get can summarize per block and per cultivar.

```
dat %>%
 group_by(gen) %>%
 dlookr::describe(yield) %>%
 select(2:sd) %>%
 arrange(desc(mean))
# A tibble: 35 \times 5
  gen
            n
                 na mean
                              sd
  <fct> <int> <int> <dbl> <dbl>
1 G19
            2
                  0 6.07 1.84
2 G07
            2
                  0 5.74 0.976
            2
3 G33
                  0 5.13 0.820
            2
                  0 4.96 0.940
 4 G06
            2
                  0 4.94 1.68
5 G09
            2
6 G11
                  0 4.93 1.03
7 G14
            2
                 0 4.92 0.877
            2
                 0 4.89 1.80
8 G27
9 G03
            2
                  0 4.78 0.0424
                  0 4.78 0.361
10 G25
            2
# i 25 more rows
```

Additionally, we can decide to plot our data.

```
# sort genotypes by mean yield
gen_order <- dat %>%
  group_by(gen) %>%
  summarise(mean = mean(yield, na.rm = TRUE)) %>%
  arrange(mean) %>%
  pull(gen) %>%
```

```
as.character()
ggplot(data = dat) +
  aes(
    y = yield,
    x = gen,
    shape = rep
  geom_line(
    aes(group = gen),
    color = "darkgrey"
  ) +
  geom_point() +
  scale_x_discrete(
    name = "Genotype",
    limits = gen_order
  ) +
  scale_y_continuous(
   name = "Yield",
   limits = c(0, NA),
   expand = expansion(mult = c(0, 0.05))
  ) +
  scale_shape_discrete(
   name = "Replicate"
  ) +
  guides(shape = guide_legend(nrow = 1)) +
  theme_classic() +
  theme(
    legend.position = "top",
    axis.text.x = element_text(angle = 90, vjust = 0.5)
  )
```



Finally, since this is an experiment that was laid with a certain experimental design (= a resolvable row column design) - it makes sense to also get a field plan. This can be done via desplot() from {desplot}. In this case it is worth noting that there is missing data, as yield values for two plots are not present in the data.

```
desplot(
  data = dat,
  form = gen ~ col + row | rep, # fill color per genotype, headers per replicate
  text = gen,
  cex = 0.7,
  shorten = "no",
  out1 = row, out1.gpar=list(col="black"), # lines between rows
  out2 = col, out2.gpar=list(col="black"), # lines between columns
  main = "Field layout",
  show.key = FALSE
)
```

Field layout

R1							R2						
G17	G09	G03	G34	G13	G35	G01	G01	G27	G16	G29	G14	G28	G22
G24	G25	G05	G32	G02	G27	G08	G33	G09	G17	G18	G32		G02
G22	G11	G19	G26	G29	G15	G23	G11	G07	G26	G05	G35	G10	G30
G10	G14		G21	G31	G06	G18	G24	G21	G12	G04	G23	G13	G03
G20	G04	G33	G28	G07	G12	G30	G31	G19	G25	G34	G20	G08	G06

2 Model

Finally, we can decide to fit a linear model with yield as the response variable and gen as fixed effects, since our goal is to compare them to each other. Since the trial was laid out in rows and columns, we also need rowF and colF effects in the model, but these can be taken either as a fixed or as random effects. Since our goal is to compare genotypes, we will determine which of the two models we prefer by comparing the average standard error of a difference (s.e.d.) for the comparisons between adjusted genotype means - the lower the s.e.d. the better.

As a result, we find that the model with random row and column effects has the slightly smaller

s.e.d. and is therefore more precise in terms of comparing genotypes.

⚠ Model assumptions met? (click to show)

It would be at this moment (i.e. after fitting the model and before running the ANOVA), that you should check whether the model assumptions are met. Find out more in the summary article "Model Diagnostics"

3 ANOVA

[1] 1.18948

Based on our model, we can then conduct an ANOVA:

```
ANOVA <- anova(mod.rrc)
ANOVA

Type III Analysis of Variance Table with Satterthwaite's method
Sum Sq Mean Sq NumDF DenDF F value Pr(>F)
gen 32.157 0.94578 34 33 0.8969 0.6233
```

Accordingly, the ANOVA's F-test did not find the cultivar effects to be statistically significant (p = 0.623).

4 Mean comparison

Besides an ANOVA, one may also want to compare adjusted yield means between cultivars via post hoc tests (t-test, Tukey test etc.).

```
mean_comp <- mod.rrc %>%
  emmeans(specs = ~ gen) %>% # adj. mean per genotype
  cld(adjust = "Tukey", Letters = letters) # compact letter display (CLD)
mean_comp
                    df lower.CL upper.CL .group
 gen emmean
               SE
 G15
       1.93 1.151 32.5
                         -2.072
                                    5.93 a
 G23
       3.27 0.804 32.5
                          0.470
                                    6.06 a
 G22
       3.54 0.794 32.5
                          0.784
                                    6.31 a
 G26
       3.58 0.842 24.6
                          0.561
                                    6.59 a
 G04
       3.62 0.796 32.5
                          0.857
                                    6.39 a
 G30
       3.75 0.851 23.9
                          0.692
                                    6.81 a
 G35
       3.75 0.800 32.5
                          0.975
                                    6.54 a
 G29
       3.77 0.800 32.5
                          0.985
                                    6.55 a
 G24
                                    6.91 a
       3.85 0.853 23.9
                          0.779
 G02
       3.88 0.847 24.6
                                    6.92 a
                          0.852
 G28
       3.95 0.811 32.5
                          1.132
                                    6.77 a
 G31
       3.98 0.802 32.5
                          1.192
                                    6.77
                                    6.87 a
 G10
       4.07 0.804 32.5
                          1.275
 G16
       4.20 1.163 32.5
                          0.155
                                    8.24 a
 G17
       4.22 0.794 32.5
                          1.459
                                    6.98 a
 G08
       4.36 0.796 32.5
                          1.587
                                    7.12 a
 G13
       4.36 0.804 32.5
                                    7.15 a
                          1.560
                                    7.60 a
 G34
       4.51 0.859 23.9
                          1.422
 G12
       4.58 0.802 32.5
                          1.797
                                    7.37
 G18
       4.60 0.804 32.5
                          1.805
                                    7.40 a
                                    7.65 a
 G32
       4.62 0.847 24.6
                          1.582
                                    7.76 a
 G01
       4.68 0.858 24.6
                          1.611
 G05
       4.71 0.798 32.5
                          1.939
                                    7.49 a
 G20
       4.75 0.872 24.6
                          1.629
                                    7.87 a
 G21
       4.76 0.858 24.6
                          1.686
                                    7.83 a
 G25
                                    7.54 a
       4.78 0.796 32.5
                          2.007
 G03
                                    7.58 a
       4.78 0.804 32.5
                          1.985
                                    7.67 a
 G27
       4.89 0.800 32.5
                          2.110
 G14
       4.92 0.804 32.5
                          2.125
                                    7.72 a
```

```
G11
      4.93 0.847 24.6
                         1.897
                                   7.96 a
 G09
      4.94 0.849 23.9
                         1.889
                                    7.99 a
                                   7.73 a
 G06
      4.96 0.796 32.5
                         2.197
 G33
      5.13 0.796 32.5
                         2.362
                                   7.90 a
 G07
      5.74 0.802 32.5
                          2.952
                                    8.53 a
 G19
     6.07 0.796 32.5
                          3.302
                                   8.84 a
Degrees-of-freedom method: kenward-roger
Confidence level used: 0.95
Conf-level adjustment: sidak method for 35 estimates
P value adjustment: tukey method for comparing a family of 35 estimates
significance level used: alpha = 0.05
NOTE: If two or more means share the same grouping symbol,
      then we cannot show them to be different.
      But we also did not show them to be the same.
```

It can be seen that while some genotypes have a higher yield than others, no differences are found to be statistically significant here. Thus, this is in agreement with the non-significant ANOVA results.

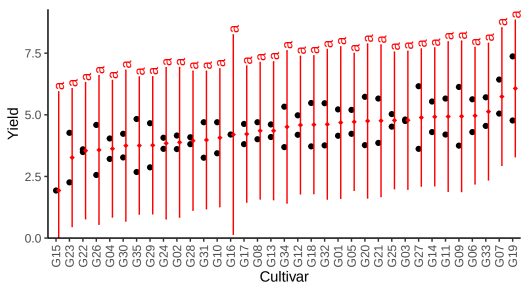
Note that if you would like to see the underlying individual contrasts/differences between adjusted means, simply add details = TRUE to the cld() statement. Furthermore, check out the Summary Article "Compact Letter Display".

Finally, we can create a plot that displays both the raw data and the results, i.e. the comparisons of the adjusted means that are based on the linear model.

```
# reorder genotype factor levels according to adjusted mean
my_caption <- "Black dots represent raw data. Red diamonds and error bars represent adjust

ggplot() +
    # green/red dots representing the raw data
    geom_point(
        data = dat,
        aes(y = yield, x = gen)
    ) +
    # red diamonds representing the adjusted means
    geom_point(
        data = mean_comp,
        aes(y = emmean, x = gen),
        shape = 18,
        color = "red",
        position = position_nudge(x = 0.2)</pre>
```

```
# red error bars representing the confidence limits of the adjusted means
geom_errorbar(
  data = mean_comp,
  aes(ymin = lower.CL, ymax = upper.CL, x = gen),
  color = "red",
 width = 0.1,
 position = position_nudge(x = 0.2)
) +
# red letters
geom_text(
  data = mean_comp,
  aes(y = upper.CL, x = gen, label = str_trim(.group)),
  color = "red",
  angle = 90,
 hjust = -0.2,
 position = position_nudge(x = 0.2)
) +
scale_x_discrete(
 name = "Cultivar",
 limits = as.character(mean_comp$gen)
) +
scale_y_continuous(
 name = "Yield",
  # limits = c(0, NA),
  expand = expansion(mult = c(0, 0.05))
coord_cartesian(ylim = c(0, NA)) +
labs(caption = my_caption) +
theme_classic() +
theme(plot.caption = element_textbox_simple(margin = margin(t = 5)),
      plot.caption.position = "plot",
      axis.text.x = element_text(angle = 90, vjust = 0.5))
```



Black dots represent raw data. Red diamonds and error bars represent adjusted means with 95% confidence limits per cultivar. Means followed by a common letter are not significantly different according to the Tukey–test.

Kempton, R. A., P. N. Fox, and M. Cerezo. 1996. Statistical Methods for Plant Variety Evaluation. Springer Netherlands. https://doi.org/10.1007/978-94-009-1503-9.