

Design and Analysis of Experiments 10 - Analysis of Variance

Version 2.11

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"Nothing in life is to be feared, it is only to be understood. Now is the time to understand more, so that we may fear less."

Marie Sklodowska Curie 1867–1934 Polish-French physicist and chemist.



Comparison of multiple means

Introduction

In the previous sections, we have (hopefully) developed a solid understanding of the main concepts associated with comparing the means of two groups;

There are many cases, however, in which one may want to perform inferences about differences of the means of multiple populations;

We will develop the main concepts and ideas related with this kind of test by examining a simple example, related to a paper manufacturing operation.

Image: http://goo.gl/xYVW0M

Problem definition

Tensile strength (TS) is an important characteristic for certain types of paper for industrial use;

A reasonable conjecture is that this characteristic is influenced by the kind of hardwood used in the manufacturing process.

The process engineer wants to investigate whether four different hardwoods result in papers with relevant differences of TS, using a pilot plant as his experimental unit.

Problem definition

The total budget allocated for the experiment allows the execution of six production runs for each kind of hardwood.

Under these specifications, the experiment has a single experimental factor (Hardwood type) with a=4 levels (hardwood types A, B, C and D) and n=6 replicates at each level.

The response variable will be the tensile strength of paper (measured in kPa). The engineering team is interested in finding out whether any hardwood leads to an increase in the mean TS value of the paper.

The minimum difference of practical meaning is defined as 5kPa, and a reasonable upper estimate for the standard deviation of this process is $\hat{\sigma}=6kPa$. Desired error levels are defined as $\alpha=0.1$ and $\beta=0.2$.

Exploratory data analysis

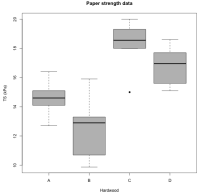
```
<- read.table(file = "../data files/paper_strength.csv",
 paper
                        header = TRUE.
                        sep = ",")
                                                    Paper strength data
 summary(paper)
Hardwood
             TS_kPa
A:6 Min. : 9.88
B:6 1st Ou.:13.90
C:6 Median :15.35
D:6
         Mean :15.56
                                     (kPa)
         3rd Ou.:17.77
         Max. :20.00
>
 boxplot (TS kPa~Hardwood,
         data = paper,
          xlab = "Hardwood",
         vlab = "TS (kPa)",
         main = "Paper strength data",
         pch = 16.
                                                       Hardwood
         col = "gray")
```

Exploratory data analysis

The boxplot suggests the existence of differences among the factor levels;

Besides, we can also observe a small variability in the spread of different levels; some suggestion of asymmetry in level B; and a possible outlier in level C.

These characteristics will need to be taken into account during the analysis.



This data can be described by a linear statistical model of the form:

$$y_{ij} = \underbrace{\mu_i + \epsilon_{ij}}_{\text{Means model}} = \underbrace{\mu + \tau_i + \epsilon_{ij}}_{\text{Effects model}} \begin{cases} i = 1, \dots, a \\ j = 1, \dots, n \end{cases}$$

where μ is the overall mean, τ_i represents the effect of the *i*-th level, and ϵ_{ij} is the residual (random error, or unmodeled variability);

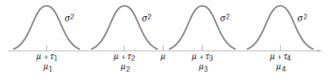
In the derivation of the statistical test for the existence of differences in the group means, we will employ the effects model, and initially consider a few assumptions about the residuals:

$$y_{ij} = \mu + \tau_i + \epsilon_{ij} \begin{cases} i = 1, \dots, a \\ j = 1, \dots, n \end{cases}$$
, with $\epsilon_{ij} \stackrel{\text{i.i.d.}}{\sim} \mathcal{N}\left(0, \sigma^2\right)$

Example: tensile strength

Statistical model

If these assumptions are correct, the populations are expected to be distributed as:



Since we are interested in testing our data for differences in the mean values of each population, the test hypotheses can be described as:

$$\begin{cases} H_0: \tau_i = 0, & \forall i \in \{1, 2, \dots, a\} \\ H_1: \exists \ \tau_i \neq 0 \end{cases}$$

If data collection is performed in random order, under constant experimental conditions, we have a *completely randomized design*.

This approach to modeling the mean effects of specific factor levels is known as the *fixed effects model*;

This approach is appropriate to testing hypotheses in situations when factor levels are arbitrarily defined by the experimenter;

For these cases, the inference is made over the mean values for each level, and cannot be extended to similar levels that were not tested (e.g., other types of hardwood);

Other situations may require different kinds of modeling, such as *random* or *mixed effects models*, but these will not be explored here.

Development

As mentioned earlier, we will use the *effects model* for describing the development of the statistical test:

$$y_{ij} = \mu + \tau_i + \epsilon_{ij}$$

$$\begin{cases} i = 1, \dots, a \\ j = 1, \dots, n \end{cases}$$

where treatment effects are seen as deviations from the grand mean μ .

By construction, we have that:

$$\sum_{i=1}^{a} \tau_i = 0;$$

Development

The total variability of the data can be expressed by the *total sum of squares*, which represents the sum of the squared deviations between each observation and the overall sample mean:

$$SS_T = \sum_{i=1}^a \sum_{j=1}^n (y_{ij} - \bar{y}_{\bullet \bullet})^2$$

With some relatively simple algebra, the SS_T can be divided into two terms, representing the within-group and the between-group variability:

$$SS_{T} = \sum_{i=1}^{a} \sum_{j=1}^{n} (y_{ij} - \bar{y}_{\bullet \bullet})^{2} = \underbrace{n \sum_{i=1}^{a} (\bar{y}_{i \bullet} - \bar{y}_{\bullet \bullet})^{2}}_{SS_{Levels}} + \underbrace{\sum_{i=1}^{a} \sum_{j=1}^{n} (y_{ij} - \bar{y}_{i \bullet})^{2}}_{SS_{E}}$$

where • indicates the summation over an index, and ⁻ indicates an averaging operation.

Development

Dividing the sums of squares by their respective number of degrees of freedom yields quantities known as *mean squares*.

The relevant means squares for our test will be the *levels mean square* and the *residual mean square*:

$$MS_E = rac{SS_E}{a(n-1)}$$
 $MS_{Levels} = rac{SS_{Levels}}{a-1}$

The expected values of these quantities are:

$$E[MS_E] = \sigma^2$$

$$E[MS_{Levels}] = \sigma^2 + \frac{n\sum_{i=1}^a \tau_i^2}{a-1}$$

Development

$$E[MS_E] = \sigma^2$$

$$E[MS_{Levels}] = \sigma^2 + \frac{n\sum_{i=1}^a \tau_i^2}{a-1}$$

Notice that MS_E is an unbiased estimator for the common variance of the residuals, while MS_{Levels} is biased by a term that is proportional to the squared values of the τ_i coefficients.

However, under H_0 we have that $\tau_i = 0$ for all i, that is, $E[MS_{Levels}] = E[MS_E] = \sigma^2$. But only if the null hypothesis is true.

Development

It can be shown that, if H_0 is true, the statistic

$$F_0 = \frac{MS_{Levels}}{MS_E}$$

is distributed according to an F distribution with a-1 degrees of freedom for the numerator and a(n-1) for the denominator. The usual notation is $F_{(a-1),a(n-1)}$

If H_0 is false, the expected value of MS_{Levels} is larger than that of MS_E , which results in larger values of F_0 and defines the critical region for our test:

Reject H_0 at the α significance level if $f_0 > F_{\alpha;(a-1),a(n-1)}$

Computational analysis

The *ANOVA table* provides information on the sources of variation, together with their corresponding degrees of freedom, sums of squares and mean square values. The table also informs the values of the test statistic and the corresponding p-value of the test (Pr(>F)).

In this case, the low p-value ($p=4.56\times 10^{-5}$) suggests the rejection of the null hypothesis in favor of the alternative. But what does that mean?

Computational analysis

Recall the null and alternative hypotheses for the ANOVA:

$$\begin{cases} H_0: \tau_i = 0, \ \forall i \\ H_1: \exists \ \tau_i \neq 0 \end{cases}$$

The rejection of the null hypothesis leads to the conclusion that *there is* at least one level with an effect significantly different from zero. But which one?

For this analysis to be complete, we still need to answer two questions:

- Can we verify the assumptions of the test?
- Which means are different from which, and by how much?

Assumptions

Model validation

As mentioned earlier, the ANOVA model is based on three assumptions on the behavior of the residuals:

- Independence;
- Homoscedasticity, i.e., equality of variances across groups;
- Normality;

The residuals of the model can be easily obtained as:

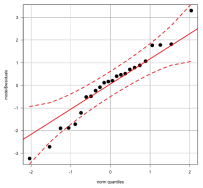
$$e_{ij} = y_{ij} - \hat{y}_{ij} = y_{ij} - (\hat{\mu} + \hat{\tau}_i) = y_{ij} - \bar{y}_{i\bullet}$$

Assumptions

Model validation

The normality assumption can be tested using the Shapiro-Wilk test coupled with a normal QQ plot of the residuals:

```
> shapiro.test(model$residuals)
Shapiro-Wilk normality test
data: model$residuals
W = 0.9722, p-value = 0.7225
> library(car)
> qqPlot(model$residuals,
pch = 16,
lwd = 3,
cex = 2,
las = 1)
```



The ANOVA is relatively robust to moderate violations of normality, as long as the other assumptions are verified or the sample size is large enough.

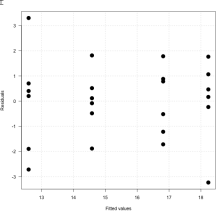
Assumptions

Model validation

The homoscedasticity assumption can be verified by the Fligner-Killeen test, together with plots of residuals by fitted values:

```
> fligner.test(TS_kPa~Hardwood, data = paper)
Fligner-Killeen test of homogeneity of
variances
data: TS_kPa by Hardwood
Fligner-Killeen:
med chi-squared = 1.0622, df = 3,
p-value = 0.7862
>
> plot(x = model$fitted.values,
+ y = model$residuals,
+ ...)
```

ANOVA is relatively robust to modest violations of homoscedasticity, as far as the sample is *balanced*.



Assumptions Model validation

As usual, the independence assumption should be guaranteed (to the best of the experimenter's knowledge) on the design phase;

To test for serial correlations, we can use the Durbin-Watson test:

```
> durbinWatsonTest(model)
 lag Autocorrelation D-W Statistic p-value
        -0.008996302
                         1.872801
                                     0.868
Alternative hypothesis: rho != 0
           = seq_along(model$residuals)
 plot(x
           = model$residuals,
      type = "1", ...)
 points(x
             = seq_along(model$residuals),
             = model$residuals,
        tvpe = "p",
        col
             = as.numeric(paper[,1]), ...)
```

The ANOVA is sensitive to violations of independence. Randomization and attention to possibly influential factors can help avoiding violations of this assumption.

The need for multiple comparisons

If the ANOVA assumptions are verified (i.e., if we have solid grounds for trusting the result of the test), we usually need to determine *which* levels of the factor are significantly different^a;

Whenever possible, the planning of which comparisons will be after an analysis of variance procedure should be defined *a priori*. Post-hoc definition of hypotheses (a.k.a. HARKing^b) are a common entry point for researcher biases into the analysis, and should be dealt with very carefully.

^aOf course this is only necessary if we rejected H_0 in the original test. For more on how to proceed with nonsignificant results, see Ellis(2010).

^bHypothesizing After the Results are Known. See Kerr(1998).

Types of comparisons

The planning of multiple comparisons must be guided by the technical question underlying the experiment.

Whenever possible, the researcher should opt to perform the smallest number of comparisons needed to adequately answer his or her question. This will require the smallest sample size, or result in the largest power for a given experimental setup.

Usual questions involve (but are not limited to):

- How does one level compare to the others?
- How does each level compare to the grand mean?
- How do the levels compare to each other (all vs. all)?

MHT considerations

The multiple comparisons performed after an ANOVA are essentially composed of a series of t-tests for the difference between two population means, with some slight modifications;

If the assumptions of the ANOVA are verified, we already have some information about the data: we know, for instance, that the groups are homoscedastic, and that their common variance is estimated by MS_E (with a(n-1) degrees of freedom);

We also know that, if we are going to perform multiple tests on the same data set, that the probability of a type-I error on each test is α . If we want to to keep our overall error rate controlled at a given level, we will need to correct the α value used for each test.

MHT corrections

There are a number of ways of adjusting the α value of the pairwise comparisons in order to maintain the *familywise error rate* (FWER) at a controlled level^c.

Two of the most common (and most conservative) are the Bonferroni and the Šidák corrections.

Assuming ${\it K}$ planned comparisons, the Bonferroni method tests each individual hypothesis with:

$$lpha_{\mathit{adj}} = rac{lpha_{\mathit{family}}}{\mathit{K}}$$

while the Šidák correction uses:

$$\alpha_{adj} = 1 - \left(1 - \alpha_{family}\right)^{1/K}$$

^CThe methods presented here work well for a relatively small number of comparison. For more on MHT, see Schaffer(1995)'s discussion on controlling the False Discovery Rate.

All vs. all

Pairwise comparisons of the *all vs. all* type appear whenever we are simply interested in detecting which levels are significantly different from which, without any prior information or special interest in one specific level or ordering.

In these cases, the number of comparisons is K = a(a-1)/2, where a is the number of levels.

The sample size calculations for this case follow the same equations used for the t test for two independent samples, but with the α value corrected for multiple hypotheses and the number of degrees of freedom of the reference distribution equal to those of the residual term in the ANOVA, i.e., a(n-1).

All vs. all

For performing *all vs. all* multiple comparisons, a common alternative is to use Tukey's *Honest Significant Difference* (HSD) approach. This method is generally chosen because it provides a slightly higher power when compared to the Bonferroni correction^d.

The usual approach is to calculate the sample size using Bonferroni (or Šidák)-corrected α -values (for simplicity), and performing the tests using Tukey's HSD corrections (for increased power):

$$n = 2\left(\frac{\left(t_{(lpha_{adj}/2)} + t_{eta}\right)\hat{\sigma}}{\delta^*}\right)^2$$

with a(n-1) degrees of freedom for the t variables.

^dThe difference is due to Tukey's approach using a modified value for the t_{β} term in the power calculations. See Mathews (2011) and Montgomery (2010) for details.

> library(multcomp)

All vs. all

Suppose that was the case for our hardwood example:

> paper tukey <- glht(model, linfct = mcp(Hardwood = "Tukey"))</pre>

```
> paper tukey CI <- confint(paper tukey, level = 0.95)
Simultaneous Confidence Intervals
Multiple Comparisons of Means: Tukey Contrasts
Fit: aov(formula = TS kPa ~ Hardwood, data = paper)
                                                       95% family-wise confidence level
Ouantile = 2.7989
95% family-wise confidence level
Linear Hypotheses:
           Estimate lwr
                            upr
B - A == 0 -1.9867 -4.6478 0.6745
C - A == 0 3.6500 0.9889 6.3111
D - A == 0 2.2333 -0.4278 4.8945
C - B == 0 5.6367 2.9755 8.2978
                                              C - B
D - B == 0 4.2200 1.5589 6.8811
D - C == 0 -1.4167
                    -4.0778 1.2445
> plot(paper tukev CI, ...)
```

. Hardwood data .

All vs. one

Pairwise comparisons of the *all vs. one* type usually emerge in the context of comparing levels against a control:

- Comparison of a proposed approach vs. existing ones;
- Comparison of different approaches vs. a standard one (or a placebo-like group);

In these cases, the number of comparisons is K = a - 1, where a is the number of levels. Each test can again be performed using the t_0 test statistic:

$$t_0^i = \frac{\bar{y}_i - \bar{y}_0}{\hat{\sigma}\sqrt{\left(\frac{1}{n_i} + \frac{1}{n_0}\right)}}$$

All vs. one

There are two main approaches to calculating sample size for *all vs. one* comparisons:

- Balanced design;
- Optimal allocation of units.

With a balanced design (that is, all levels have the same number of observations), the calculation of n follows the same approach as the *all vs. all* comparisons, but correcting α for only a-1 comparisons.

For the optimal allocation of units, an unbalanced design is used.

All vs. one - optimal allocation

As several levels will be compared against the single control group, the relative importance of the latter is greater and therefore it should have a larger sample size.

To maximize the power of this multiple comparisons procedure, the sample size of the control group should be:

$$n_0 = n_i \sqrt{K}$$

where n_i is the common sample size for the non-control levels:

$$n_i = \left(1 + \frac{1}{\sqrt{K}}\right) \left(\frac{(t_{(\alpha_{adj}/2)} + t_{\beta})\hat{\sigma}}{\delta^*}\right)^2$$

A good free software for doing sample size calculations and power analysis in nontrivial contexts such as this one is G*Power 3, http://www.gpower.hhu.de/. It is also not difficult to implement these calculations in R.

All vs. one - Dunnett's test

As in the case of *all vs. all* comparisons, there is a test that is usually employed for its superior sensitivity: Dunnett's test.

The control group sample size n_0 calculated assuming that Bonferroni-corrected t-tests will be used is slightly overestimated in relation to the required n_0 for Dunnett's test, but in practice the differences are small enough not to matter;

All vs. one

Assuming that in our example the *B* level is the standard one, against which the other ones are to be compared:

> paper\$Hardwood <- relevel(paper\$Hardwood, ref = "B")</pre>

```
> model2
            <- aov(TS kPa~Hardwood, data = paper)
> paper_dunnett <- glht (model2, linfct = mcp(Hardwood = "Dunnett"))</pre>
> paper dunnett CI <- confint(paper dunnett, level = 0.95)
Simultaneous Confidence Intervals
                                                        95% family-wise confidence level
Multiple Comparisons of Means: Dunnett Contrasts
Fit: aov(formula = TS kPa ~ Hardwood,
         data = paper)
Ouantile = 2.5394
95% family-wise confidence level
Linear Hypotheses:
           Estimate lwr upr
                                               C-B
A - B == 0 1.9867 -0.4277 4.4011
C - B == 0 5.6367 3.2223 8.0511
D - B == 0 4.2200 1.8056 6.6344
> plot(paper_dunnett_CI, ...)
```

Some final considerations

The kind of comparisons that are to be performed after an ANOVA should be planned in advance, as it influences your data collection and sample size calculations. There are of course sample size formulas for the pure ANOVA, but these are usually of limited use since researchers frequently want to know where the detected differences lie.

There are a myriad of approaches for post-ANOVA multiple comparisons^e, but in general the formulas for sample size calculation will follow he ideas outlined above: correct the α value to account for type-I error inflation and calculate n based on formulas for two-sample t tests.

e Check Hothorn et al. (2008) for an idea on how varied this can get.

Sample size formulas for ANOVA

If one is interested in calculating the required sample size for the ANOVA procedure (without worrying about the eventual post-hoc testing), the formulas are almost as simple as those used for the t tests.

Essentially, the power/sample size calculations for the ANOVA boil down to the equality:

$$F_{(1-lpha)} = F_{eta;\phi}$$

with both F distributions having (a-1) degrees of freedom in the numerator and a(n-1) in the denominator. The noncentrality parameter ϕ is given by:

$$\phi = \frac{n\sum_{i=1}^{a} \tau_i^2}{\hat{\sigma}^2}$$

Sample size formulas for ANOVA

To illustrate the sample size calculation procedure, imagine an experimental design with $a=4,~\alpha=0.05,~\hat{\sigma}=7$, and suppose that the researcher wants to be able to detect whether any two means present differences of magnitude $\delta^*=12$ with power $(1-\beta)=0.8$.

Under these conditions, two scenarios tend to be of interest: the first is if we have two levels biased symmetrically about the grand mean, and all the others equal to zero:

$$\tau = \left\{-\frac{\delta^*}{2}, \frac{\delta^*}{2}, 0, 0\right\}$$

and the second is if we have one level biased in relation to all others:

$$\tau = \left\{ -\frac{(a-1)\delta^*}{a}, \frac{\delta^*}{a}, \frac{\delta^*}{a}, \frac{\delta^*}{a} \right\}$$

Sample size formulas for ANOVA

For the first case we have a noncentrality parameter of:

$$\phi = \frac{4\left(6^2 + 6^2 + 0 + 0\right)}{7^2} = 5.88$$

Which allows us to calculate the required sample size by iterating on *n* until:

$$F_{(1-\alpha)} \leq F_{\beta;\phi}$$

Sample size formulas for ANOVA

Doing it manually:

```
> a      <- 4
> alpha      <- 0.05
> sigma      <- 7
> delta      <- 12
> beta      <- 0.2
>
> tau <- c(-delta/2, delta/2, rep(0, a - 2)) # define tau vector
> n      <- 2
> while (qf(1 - alpha, a - 1, a*(n - 1)) >
+            qf(beta, a - 1, a*(n - 1), n*sum(tau^2)/sigma^2)) n <- n + 1
> print(n)
[1] 9
```

Using power.anova.test():

Sample size formulas for ANOVA

The second case (one level biased in relation to all others) is also quite easy to calculate manually, but lets keep it simple:

```
> tau <- c(-delta*(a - 1)/a, rep(delta/a, a - 1))
> vartau <- var(tau)
> power.anova.test(groups = 4, between.var = vartau,
+ within.var = sigma^2, sig.level = alpha,
+ power = 1 - beta)$n
[1] 6.018937
```

It is important to remember that these are the sample sizes required for the ANOVA only - any multiple comparisons procedure executed afterwards to pinpoint the significant differences will have smaller power for same-sized effects (unless more observations are added). This is one reason why it is common to design experiments calculating the sample sizes based on the multiple comparisons procedure, instead of using the ANOVA formulas.

Bibliography

Required reading

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Recommended reading

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- P. Mathews, Sample Size Calculations: Practical Methods for Engineers and Scientists. Ch. 8, 1st ed., MMB, 2010.
- T. Hothorn, F. Bretz, P. Westfall, Simultaneous Inference in General Parametric Models. Biometrical Journal 50(3), 346–363, 2008.

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