

# Design and Analysis of Experiments

## 11 - Blocking

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*“Science doesn’t always go forwards. It’s a bit like doing a Rubik’s cube. You sometimes have to make more of a mess with a Rubik’s cube before you can get it to go right.”*

Jocelyn Bell Burnell  
1943 -  
Irish astrophysicist



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Image:

<http://www.topbritishinnovations.org/PastInnovations/DiscoveryofPulsars.aspx>

# Blocking

## Nuisance factors

In the completely randomized design (CRD) introduced in the last chapter, the observations are assigned - as the name implies - in a completely random manner.

Implicitly, we assume that the experimental conditions are homogeneous, i.e., that no other important effects are present besides those of the experimental factor.

In some situations, however, the experiment may present other factors that are not of immediate interest, but that can influence the response variable. This was already discussed when we introduced the concept of *pairing* in Chapter 7.

# Blocking

## Nuisance factors

The generalization of the *paired design* for an arbitrary number of factor levels is called *blocking*, which is an elegant way to configure the data collection in order to enable the modeling and exclusion of the effects of nuisance factors.

In fact, it can be argued that the systematic use of blocking is beneficial even in the absence of known nuisance factors, as it helps boost statistical power by excluding the between-replicates variability from the residual term.

*“Block what you can,  
randomize what you cannot.”*

George E.P. Box (1919–2013)  
British Statistician



# Blocking

## Blocking *versus* Randomizing

The idea behind randomization is trying to prevent unknown factors to bias our observations;

Blocking comes into play whenever we know from the beginning that certain factors can influence our response variable, but for some reason we are not interested in their effects<sup>a</sup>, for instance:

- the effect of different batches of raw material when comparing the performance of chemical reactions;
- the effect of different benchmark problems when comparing the performance of computer algorithms.

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<sup>a</sup> If we're interested in their effects, they are no longer nuisance factors, but additional experimental factors. We'll deal with factorial designs in the next chapter.

# Example: algorithm comparison

## Problem definition

A Ph.D. student decides to compare a standard optimization algorithm with six modified versions for the solution of a certain family of Vehicle Routing Problems (VRPs). His intention is to verify whether any of the modified versions is systematically better than the standard one.

The algorithms are applied for the solution of 180 problem instances, divided in 36 groups of five homogeneous instances. The algorithms are run 30 times on each instance, and the cost value found after each run is recorded.

The student wants an experiment with  $\alpha = 0.05$  and power  $\pi = 1 - \beta = 0.8$  for a minimally interesting effect  $\delta^* = 50$ .

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Image: PhD Comics by Jorge Cham:

<http://www.phdcomics.com/comics/archive.php?comicid=1139>



# Example: algorithm comparison

## Nuisance factor

Besides the effect of the experimental factor (*Algorithm*), we have a possible nuisance effect due to the variability between experimental units (instance groups).

If we employ a CRD, the residual term would contain all the between-instances variability, which would almost certainly mask any between-algorithms effect of interest.

Since we can control the allocation of algorithm runs within each instance, one possibility is to do it in a systematic way so that we can model the instance effects (in addition to the factor effect).

# Example: algorithm comparison

## Randomized complete block design

Within each *block*, the order of execution should in principle be randomized, so as to prevent other unknown factors from affecting our analysis<sup>b</sup>.

This experimental setup is known as a *randomized complete block design* (RCBD) with one blocking factor.

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<sup>b</sup> For this kind of computer experiment the within-block randomization is unnecessary. It is very important, however, when dealing with systems that are less controllable.



# Randomized complete block design

## Assumptions

The RCBD assumes:

- *one replicate per block*<sup>c</sup>;
- independent blocks;
- independent within-block randomization.

Failure to account for dependency structures (i.e., using blocks that are not independent) can result in pseudoreplication, i.e., the use of the incorrect number of degrees-of-freedom in the test statistics, which usually results in an inflation of the type-I error rate.

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<sup>c</sup>For multiple within-block replicates the appropriate design is known as *Generalized randomized block design* (GRBD), which is analyzed using a factorial model.

# Example: algorithm comparison

RCBD

To comply with these requirements, we consider as our response variable the average performance of an algorithm on a particular *instance group*.

The performance of each algorithm will be averaged within each individual instance (mean of 30 runs), and then within each group (mean of 5 instances), so that each algorithm has 36 independent observations.

# Example: algorithm comparison

## Data loading and preprocessing

```
# Load data
> data <- read.table("../data files/algo.csv", header = TRUE)
# Aggregate data (algorithm means by instance group)
> aggdata <- with(data, aggregate(x = Result,
                                by = list(Algorithm, Group),
                                FUN = mean))

# Rename columns and coerce factor variables
> names(aggdata) <- c("Algorithm", "Instance_Group", "Y")
> for (i in 1:2) aggdata[, i] <- as.factor(aggdata[, i])

> levels(aggdata$Algorithm) <- c("Original",
                                unlist(lapply(X = "Mod",
                                                FUN = paste0, 1:6)))

> summary(aggdata)
```

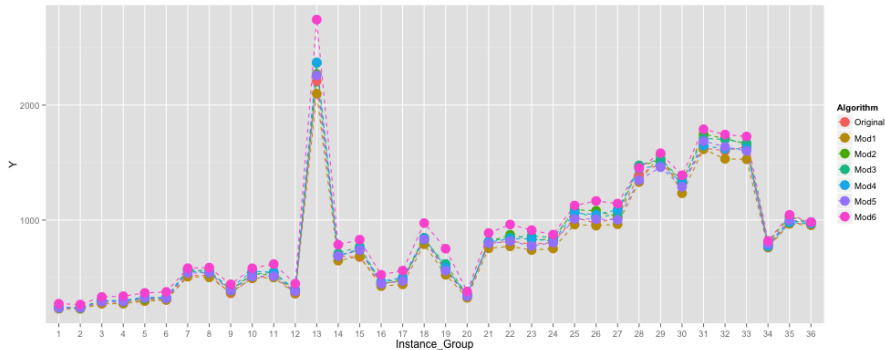
Algorithm	Instance_Group	Y
Original:36	1 : 7	Min. : 227.3
Mod1 :36	2 : 7	1st Qu.: 440.0
Mod2 :36	3 : 7	Median : 756.8
Mod3 :36	4 : 7	Mean : 821.0
Mod4 :36	5 : 7	3rd Qu.:1020.5
Mod5 :36	6 : 7	Max. :2743.6
Mod6 :36	(Other):210\$	

# Example: algorithm comparison

## Exploratory data analysis

```
library(ggplot2)
> p <- ggplot(agdata, aes(x = Instance_Group, y = Y,
                           group = Algorithm, colour = Algorithm))

> p + geom_line(linetype=2) + geom_point(size=5)
```



# Analysis of the RCBD

## Statistical model

In the general case, we have  $a$  levels of the experimental factor, and  $b$  levels for the blocking factor. The statistical model is given as:

$$y_{ij} = \mu + \tau_i + \beta_j + \epsilon_{ij} \begin{cases} i = 1, \dots, a \\ j = 1, \dots, b \end{cases}, \text{ assuming } \epsilon_{ij} \sim \mathcal{N}(0, \sigma^2)$$

Similarly to the CRD we have, by construction:

$$\sum_{i=1}^a \tau_i = 0 \qquad \sum_{j=1}^b \beta_j = 0$$

# Analysis of the RCBD

## Statistical model

In the RCBD, we are interested only on the effect of the experimental factor. Consequently, the test hypotheses refer only to its coefficients:

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

The partition of the total sample variability is given by:

$$\begin{aligned} SS_T &= \sum_{i=1}^a \sum_{j=1}^b (y_{ij} - \bar{y}_{..})^2 \\ &= \underbrace{b \sum_{i=1}^a (\bar{y}_{i.} - \bar{y}_{..})^2}_{SS_{\text{levels}}} + \underbrace{a \sum_{j=1}^b (\bar{y}_{.j} - \bar{y}_{..})^2}_{SS_{\text{blocks}}} + \underbrace{\sum_{i=1}^a \sum_{j=1}^b (y_{ij} - \bar{y}_{i.} - \bar{y}_{.j} + \bar{y}_{..})^2}_{SS_E} \end{aligned}$$

# Analysis of the RCBD

## Statistical model

The mean squares are given by:

$$MS_{\text{levels}} = \frac{SS_{\text{levels}}}{a - 1} \quad MS_{\text{blocks}} = \frac{SS_{\text{blocks}}}{b - 1} \quad MS_E = \frac{SS_E}{(a - 1)(b - 1)}$$

and their expected values are:

$$E[MS_{\text{levels}}] = \sigma^2 + \frac{b \sum_{i=1}^a \tau_i^2}{a - 1} \quad E[MS_{\text{blocks}}] = \sigma^2 + \frac{a \sum_{j=1}^b \beta_j^2}{b - 1}$$

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

# Analysis of the RCBD

## Statistical model

$$E[MS_{\text{levels}}] = \sigma^2 + \frac{b \sum_{i=1}^a \tau_i^2}{a-1} \qquad E[MS_E] = \sigma^2$$

Under the null hypotheses we have that the test statistic

$$F_0 = \frac{MS_{\text{levels}}}{MS_E}$$

is distributed according to an  $F$  distribution with  $(a-1)$  numerator degrees-of-freedom and  $(a-1)(b-1)$  denominator DoFs.

The critical region for the test is therefore given by

$$f_0 > F_{\alpha; (a-1), (a-1)(b-1)}$$



# Analysis of the RCBD

## Statistical model

```
# First model
> model <- aov(Y~Algorithm+Instance_Group, data=aggdata)

> summary(model)
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
Algorithm	6	359949	59991	41.75	<2e-16 ***
Instance_Group	35	60438639	1726818	1201.86	<2e-16 ***
Residuals	210	301725	1437		

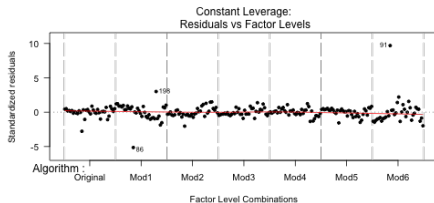
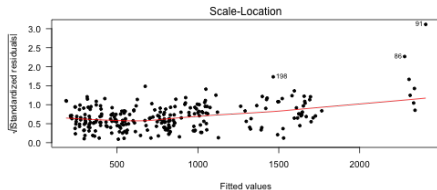
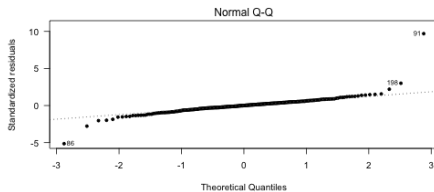
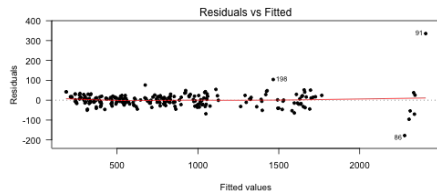
```
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

> summary.lm(model)$r.squared
[1] 0.9950618
```

# Analysis of the RCBD

## Statistical model - residuals

```
> par(mfrow = c(2, 2))  
> plot(model, pch = 20, las = 1)
```



# Analysis of the RCBD

## Statistical model - data transformation

```
# Trying with log-transformed response variable
> model2 <- aov(log(Y)~Algorithm+Instance_Group, data=aggdata)

> summary(model2)
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)	
Algorithm	6	0.60	0.1002	120.4	<2e-16	***
Instance_Group	35	89.92	2.5690	3089.0	<2e-16	***
Residuals	210	0.17	0.0008			

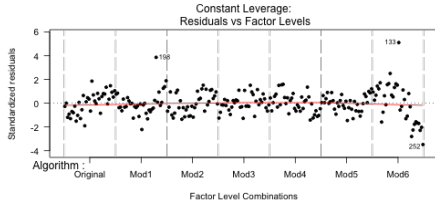
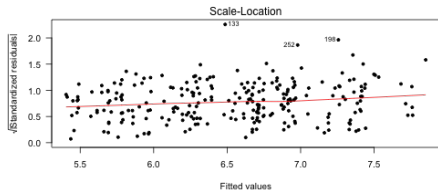
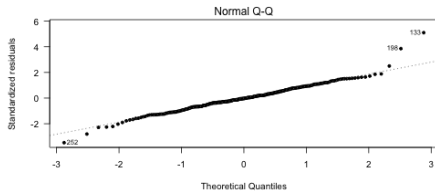
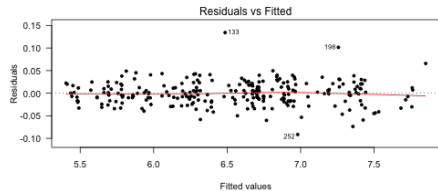
```
---
```

```
> summary.lm(model2)$r.squared
[1] 0.9980742
```

# Analysis of the RCBD

## Statistical model - residuals

```
> par(mfrow = c(2, 2))  
> plot(model2, pch = 20, las = 1)
```



# Analysis of the RCBD

## Multiple blocking factors

If there is more than one factor that need to be blocked out of the analysis, it is possible to simply add new terms to the statistical model. The sum of squares for this new term is similar to the ones already shown for the single block case;

# Analysis of the RCBD

## Blocking efficiency

After an experiment is performed and the ANOVA model is fit to the data, it is possible to calculate the *relative blocking efficiency* ( $E$ ), which quantifies how much larger a CRD would have to be in order to attain the same power as the corresponding RCBD.

The relative efficiency can be calculated from the mean squares returned in the ANOVA table of the RCBD as:

$$E = \frac{(b - 1)MS_{\text{Blocks}} + b(a - 1)MS_E}{(ab - 1)MS_E}$$

A value of 1.3, for instance, would indicate that the CRD would have required 30% more observations to achieve the same power. For the running example, this value can be calculated from the summary table:

$$E = 431.6$$

# Analysis of the RCBD

## Sample size calculations

The determination of the required sample size for an RCBD follows the same procedures as the CRD: the number of replicates calculated using the CRD formulas represent the number of blocks required (since we have one replicate per block).

The within-group variability term necessary for calculating the required sample size corresponds to the estimated residual variance, i.e., the unexplained variability after accounting for the experimental and blocking factors, which can be somewhat challenging to estimate without a information on previous studies (or a pilot experiment).

# Analysis of the RCBD

## Multiple comparisons

For the post-hoc multiple comparison procedures, we also follow the same general guidelines used for the CRD, with some minor differences.

Paired t-tests are used for the pairwise comparisons, with  $df_E = (a - 1)(b - 1)$  as the number of degrees of freedom used for the reference distributions;

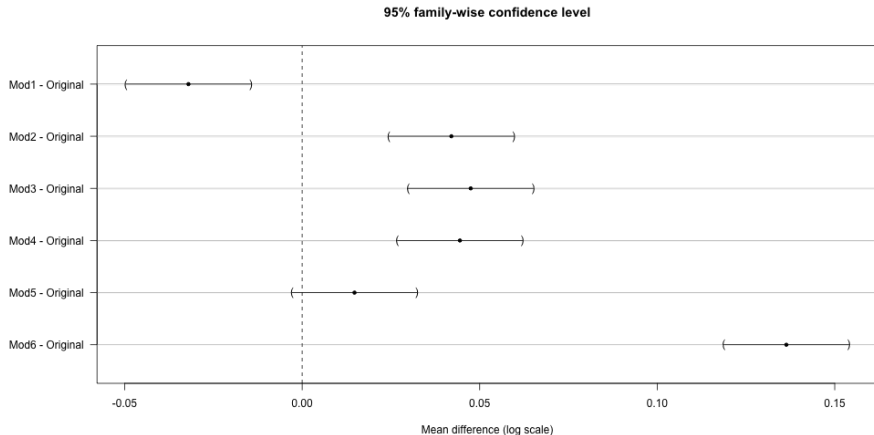
As with the CRD, it may be convenient to use the post-hoc pairwise comparison procedures as the basis for sample size calculations, particularly in cases where the cost of observations is relatively low.



# Analysis of the RCBD

## Multiple comparisons

```
> library(multcomp)
> duntest      <- glht(model2, linfct = mcp(Algorithm = "Dunnett"))
> duntestCI    <- confint(duntest)
> par(mar = c(5, 8, 4, 2), las = 1)
> plot(duntestCI, xlab = "Mean difference (log scale)")
```



# Randomized complete block designs

## Final considerations

The RCBD is a generalization of the pairing concept used with t-tests, and provides an elegant way to remove nuisance effects from the analysis;

The consequence of using a blocking design is an increased sensitivity of the test, which means smaller sample sizes.

In practice, running an experiment as a RCBD results in little experimental overhead, and should be preferred over the CRD whenever possible.

The design and analysis of incomplete block designs (e.g., with missing observations) are also relatively simple<sup>e</sup>.

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<sup>e</sup> See Montgomery (2005) for details.

# Bibliography

## Required reading

- 1 D.C. Montgomery, *Design and Analysis of Experiments*, Ch. 4-5, 5th ed., Wiley, 2005;
- 2 M. Christman, R. Littell, *STA166 - Statistical Methods Research I: RCBD ANOVA Notes II* - <http://goo.gl/KrpbKL>

## Recommended reading

- 1 R. Dawkins, *Unweaving the Rainbow: Science, Delusion and the Appetite for Wonder*, Mariner Books, 2000.
- 2 J. Morgan and C. Albon, *Partially Derivative* (podcast) - <http://www.partiallyderivative.com>

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