

The Superpower R Package: Capabilities and Validation

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This is a compilation of documents for Superpower R package written in **Markdown** and compiled by **Bookdown**.

Chapter 1

Introduction

The goal of **Superpower** is to easily simulate factorial designs and empirically calculate power using a simulation approach. The R package is intended to be utilized for prospective (a priori) power analysis. Please don't be silly and compute post hoc power.

This package, and book, expect readers to have some familiarity with R. However, we have created two Shiny apps (one for `ANOVA_power` and one for `ANOVA`) to help use these functions if you are not familiar with R.

In this book we will display a variety of ways the **Superpower** package can be used for power analysis and sample size planning for factorial experimental designs. We also included various examples of the performance of **Superpower** against other R packages (e.g., `pwr2ppl` and `pwr`) and statistical programs (such as G*Power). All uses of the `ANOVA_power` function have been run with 10000 iterations (`nsims = 10000`). If you have any issues using **Superpower** or want to expand its capabilities please raise the issue on our [GitHub](#) page.

Chapter 2

The Experimental Design

This section introduces how **Superpower** sets up the design of each power analysis. Currently, this package only provides power analysis for ANOVA and for repeated measures MANOVA.

2.1 ANOVA__design function

Currently the ANOVA__design function can create designs up three factors, for both within, between, and mixed designs. It requires the following input: design, n, mu, sd, r, and optionally allows you to set labelnames.

1. design: string that specifies the design (see below).
2. n: the sample size for each between subject condition.
3. mu: a vector with the means for each condition.
4. sd: the population standard deviation. Assumes homogeneity of variances (only one standard deviation can be provided).
5. r: the correlation for within designs (or 0 for between designs).
6. labelnames: This is an optional vector of words that indicates factor names and level names (see below).
7. A final optional setting is to specify if you want to automatically print a plot or not (plot = TRUE or FALSE)

2.1.1 Specifying the design using ‘design’

‘design’ is used to specify the design. Every factor is specified with a number, indicating the number of levels of the factor, and a letter, b or w, to indicate whether the factor is manipulated between or within participants. For example, a 2b design has two between-participant groups. A 12w design has one factor with 12 levels, all manipulated within-participants. A "2b*3w" is a design with two factors (a 2b factor and a 3w factor), the first of which has 2 between participant levels (2b), and the second of which has 3 within participants levels (3w). **If there are multiple factors (the functions take up to three different factors) separate factors with a * (asterisk).** An example of a 2b*3w design is a group of people in one condition who get a drug, and a group of people in another condition who get a placebo, and we measure their health before they take the pill, one day after they take the pill, and a week after they take the pill.

2.1.2 Specifying the means using ‘mu’

Note that for each cell in the design, a mean must be provided. Thus, for a "2b*3w" design, 6 means need to be entered.

Means need to be entered in the correct order. ANOVA_design outputs a plot so you can check if you entered all means as you intended. Always carefully check if the plot that is generated matches your expectations.

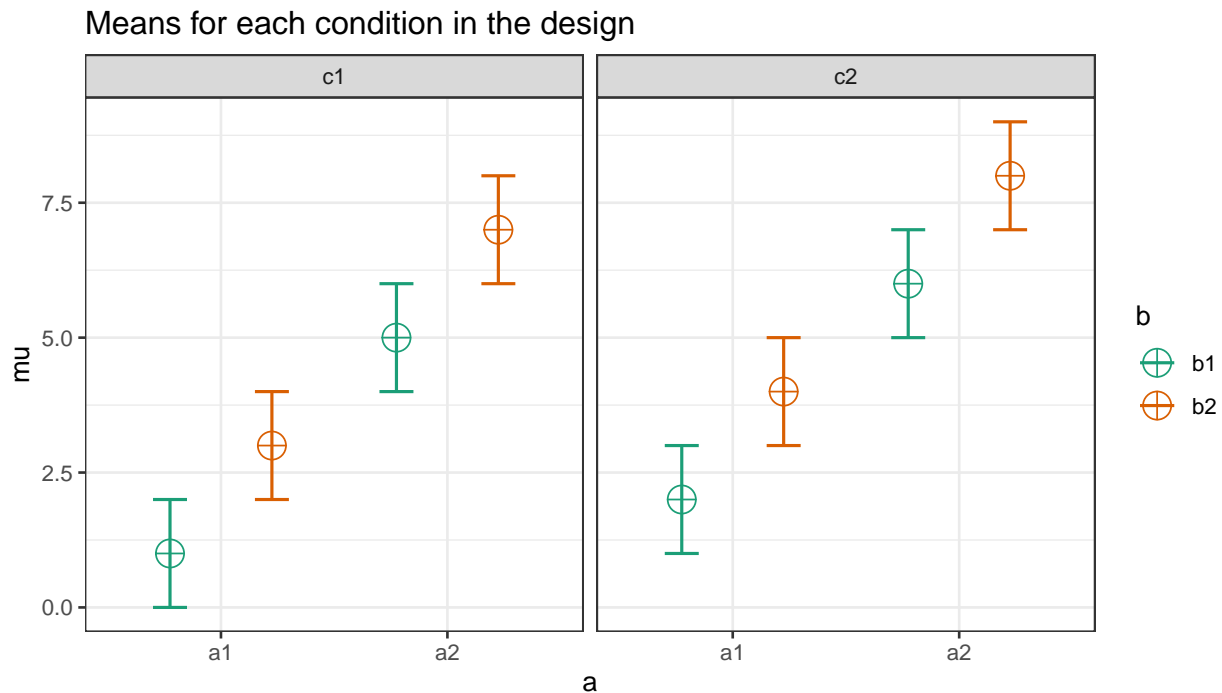
The general principle is that the code generates factors, indicated by the factor names you entered in the `labelnames` variable, (i.e., *condition* and *time*). Levels are indicated by factor names and levels (e.g., `control_time1`, `control_time2`, `control_time3`, etc).

If your design has just one factor, just enter the means in the same order as the `labelnames` (see below). For more factors, note the general pattern in the example below. Means are entered in the following order for a 3 factors design (each with 2 levels):

1. a1 b1 c1
2. a1 b1 c2
3. a1 b2 c1
4. a1 b2 c2
5. a2 b1 c1
6. a2 b1 c2
7. a2 b2 c1
8. a2 b2 c2

So if you enter the means 1, 2, 3, 4, 5, 6, 7, 8 the first 4 means correspond to level 1 of factor 1, the second 4 means correspond to level 2 of factor 1. Within the first 4 means, the first 2 correspond to level 1 of factor 2, and within those 2 means, the first corresponds to level 1 of factor 3.

The plot below visualizes means from 1 to 8 being entered in a vector: `mu = c(1, 2, 3, 4, 5, 6, 7, 8)` so you can see how the basic ordering works.



2.1.3 Specifying label names

To make sure the plots and tables with simulation results are easy to interpret, it really helps to name all factors and levels. You can enter the labels in the `'labelnames'` variable. You can also choose not to specify names. Then all factors are indicated by letters (a, b, c) and all levels by numbers (a1, a2, a3).

For the 2x3 design we have been using as an example, where there are 2 factors (condition and time of measurement), the first with 2 levels (placebo vs. medicine) and the second with three levels (time1, time2, and time3) we would enter the labels as follows:

```
c("condition", "placebo", "medicine", "time", "time1", "time2", "time3")
```

As you can see, you follow the order of the design (2b*3w), and first write the **FACTOR** label (condition) followed by the levels of that factor (placebo and medicine). Then you write the second factor name (time) followed by the three labels for each **LEVEL** (time1, time2, time3). **Do not use spaces in the names (so not "time 1" but "time1" or "time_1")**.

Some examples:

1. One within factor (time with 2 levels), 2w:

- `c("time", "morning", "evening")`

2. Two between factors (time and group, each with 2 levels), 2b*2b:

- `c("time", "morning", "evening", "group", "control", "experimental")`

3. Two between factors (time and group, first with 4 levels, second with 2 levels), 4b*2b:

- `c("time", "morning", "afternoon", "evening", "night", "group", "control", "experimental")`

2.1.4 Specifying the correlation

Depending on whether factors are manipulated within or between, variables are correlated, or not. You can set the correlation for within-participant factors. You can either assume all factors have the same correlation (e.g., $r = 0.7$), or enter the correlations for each pair of observations separately by specifying a correlation matrix.

In a 2x2 design, with factors A and B, each with 2 levels, there are 6 possible comparisons that can be made.

1. A1 vs. A2
2. A1 vs. B1
3. A1 vs. B2
4. A2 vs. B1
5. A2 vs. B2
6. B1 vs. B2

The number of possible comparisons is the product of the levels of all factors squared minus the product of all factors, divided by two. For a 2x2 design where each factor has two levels, this is:

```
((2 * 2) ^ 2 - (2 * 2))/2
```

```
## [1] 6
```

The number of possible comparisons increases rapidly when adding factors and levels for each factor. For example, for a 2x2x4 design it is:

```
((2 * 2 * 4) ^ 2) - (2 * 2 * 4))/2
```

```
## [1] 120
```

Each of these comparisons can have their own correlation if the factor is manipulated within subjects (if the factor is manipulated between subjects the correlation is 0). These correlations determine the covariance matrix. Potvin and Schutz (2000) surveyed statistical tools for power analysis and conclude that most software packages are limited to one factor repeated measure designs and do not provide power calculations for within designs with multiple factor (which is still true for software such as G*Power). Furthermore, software solutions which were available at the time (DATASIM by Bradley, Russel, & Reeve, 1996) required researchers to assume correlations were of the same magnitude for all within factors, which is not always realistic. If you do not want to assume equal correlations for all paired comparisons, you can specify the correlation for each possible comparison.

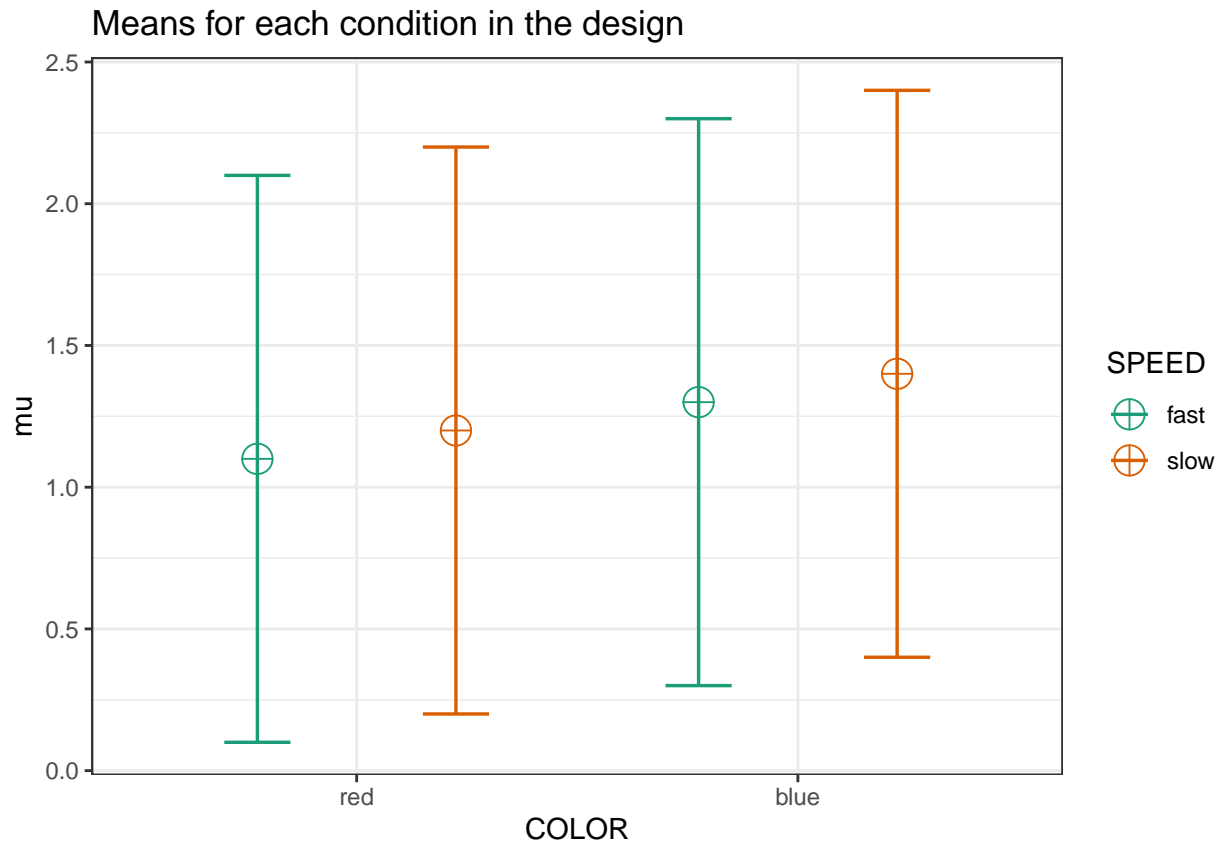
The order in which the correlations are entered in the vector should match the covariance matrix. The order for a 2x2 design is given in the 6 item list above. The general pattern is that the matrix is filled from top to bottom, and left to right, illustrated by the increasing correlations in the table below.

	a1_b1	a1_b2	a2_b1	a2_b2
a1_b1	1.00	0.91	0.92	0.93
a1_b2	0.91	1.00	0.94	0.95
a2_b1	0.92	0.94	1.00	0.96
a2_b2	0.93	0.95	0.96	1.00

The diagonal is generated dynamically (based on the standard deviation).

We would enter this correlation matrix as:

```
design_result <- ANOVA_design(design = "2w*2w",
                             n = 80,
                             mu = c(1.1, 1.2,
                                     1.3, 1.4),
                             sd = 1,
                             r <- c(0.91, 0.92,
                                     0.93, 0.94,
                                     0.95, 0.96),
                             labelnames = c("COLOR",
                                              "red", "blue",
                                              "SPEED",
                                              "fast", "slow"),
                             plot = TRUE)
```



We can check the correlation matrix by asking for it from the `design_result` object to check if it was entered the way we wanted:

```
design_result$cor_mat
```

```
##           red_fast red_slow blue_fast blue_slow
## red_fast      1.00    0.91    0.92    0.93
## red_slow      0.91    1.00    0.94    0.95
## blue_fast      0.92    0.94    1.00    0.96
## blue_slow      0.93    0.95    0.96    1.00
```

We should also check the covariance-variance matrix to ensure the `ANOVA_design` function working properly. The variance should be the diagonal element while the off-diagonal elements should be equal to `covariance` = `correlation*variance` or $cov_{x,y} = \frac{\sum_{i=1}^N (x_i - \bar{x})(y_i - \bar{y})}{N-1}$. In this case, it is identical to the correlation matrix because the variance is equal to 1.

```
design_result$sigmatrix
```

```
##           red_fast red_slow blue_fast blue_slow
## red_fast      1.00    0.91    0.92    0.93
## red_slow      0.91    1.00    0.94    0.95
## blue_fast      0.92    0.94    1.00    0.96
## blue_slow      0.93    0.95    0.96    1.00
```

2.1.5 Specifying the sample size

You can set the sample size **per condition** by setting a value for **n**. The assumption is that you will collect equal sample sizes in all conditions [expanding Superpower to allow different sample sizes in each group is a planned future option].

This means for the 2w*2w design above there is a total of 80 participants/subjects with a total of 320 observations.

2.1.6 Specifying the standard deviation

You can set the standard deviation by setting a value of **sd**. Currently, **Superpower** allows you to **violate the assumption of homogeneity of variance**. This will affect the type I error rate if the differences between conditions are extreme. Note that there is always some uncertainty in which values you can expect in the study you are planning. It is therefore useful to perform sensitivity analyses (e.g., running the simulation with the expected standard deviation, but also with more conservative or even worst-case-scenario values).

Chapter 3

One-way ANOVA

3.1 Part 1

Using the formula also used in Albers & Lakens (2018), we can determine the means that should yield a specified effect sizes (expressed in Cohen's f). Eta-squared (identical to partial eta-squared for One-Way ANOVA's) has benchmarks of .0099, .0588, and .1379 for small, medium, and large effect sizes (Cohen, 1988). Although these benchmarks are quite random, and researchers should only use such benchmarks for power analyses as a last resort, we will demonstrate a-priori power analysis for these values.

3.1.1 Two conditions

Imagine we aim to design a study to test the hypothesis that giving people a pet to take care of will increase their life satisfaction. We have a control condition, and a condition where people get a pet, and randomly assign participants to either condition. We can simulate a One-Way ANOVA with a specified alpha, sample size, and effect size, on see the statistical power we would have for the ANOVA and the follow-up comparisons. We expect pets to increase life-satisfaction compared to the control condition. Based on work by Pavot and Diener (1993) we believe that we can expect responses on the life-satisfaction scale to have a mean of approximately 24 in our population, with a standard deviation of 6.4. We expect having a pet increases life satisfaction with approximately 2.2 scale points for participants who get a pet. 200 participants in total, with 100 participants in each condition. But before we proceed with the data collection, we examine the statistical power our design would have to detect the differences we predict.

```
string <- "2b"
n <- 100
# We are thinking of running 50 people in each condition
mu <- c(24, 26.2)
# Enter means in the order that matches the labels below.
# In this case, control, cat, dog.
sd <- 6.4
labelnames <- c("condition", "control", "pet") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05
```

```
# You should think carefully about how to justify your alpha level.
# We will give some examples later, but for now, use 0.05.
```

```
simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)
```

```
knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
\begin{table}[t]
```

```
\caption{(\#tab:start_one_way)Simulated ANOVA Result}
```

	power	effect_size
anova_condition	67.91	0.0334874

```
\end{table}
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)
```

```
knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

```
\begin{table}[t]
```

```
\caption{(\#tab:start_one_way)Exact ANOVA Result}
```

	power	partial_eta_squared	cohen_f	non centrality
condition	67.69	0.029	0.1727	5.9082

```
\end{table}
```

The result shows that we have exactly the same power for the ANOVA, as we have for the t -test. This is because when there are only two groups, these tests are mathematically identical. In a study with 100 participants, we would have quite low power (around 67.7%). An ANOVA with 2 groups is identical to a t -test. For our example, Cohen's d (the standardized mean difference) is $2.2/6.4$, or $d = 0.34375$ for the difference between the control condition and pets, which we can use to easily compute the expected power for these simple comparisons using the `pwr` package.

```
pwr.t.test(d = 2.2/6.4,
            n = 100,
            sig.level = 0.05,
            type = "two.sample",
            alternative = "two.sided")$power
```

```
## [1] 0.6768572
```

We can also directly compute Cohen's f from Cohen's d for two groups, as Cohen (1988) describes, because $f = 1/2d$. So $f = 0.5 \cdot 0.34375 = 0.171875$. And indeed, power analysis using the `pwr` package yields the same result using the `pwr.anova.test` as the `power.t.test`.

```
K <- 2
n <- 100
f <- 0.171875
pwr.anova.test(n = n,
               k = K,
               f = f,
               sig.level = alpha_level)$power
```

```
## [1] 0.6768572
```

This analysis tells us that running the study with 100 participants in each condition is too likely to *not* yield a significant test result, even if our expected pattern of differences is true. This is not optimal.

Let's mathematically explore which pattern of means we would need to expect to have 90% power for the ANOVA with 50 participants in each group. We can use the pwr package in R to compute a sensitivity analysis that tells us the effect size, in Cohen's f , that we are able to detect with 3 groups and 50 participants in each group, in order to achieve 90% power with an alpha level of 5%.

```
K <- 2
n <- 100
sd <- 6.4
r <- 0
#Calculate f when running simulation
f <- pwr.anova.test(n = n,
                  k = K,
                  power = 0.9,
                  sig.level = alpha_level)$f
f
```

```
## [1] 0.2303587
```

This sensitivity analysis shows we have 90% power in our planned design to detect effects of Cohen's f of 0.2303587. Benchmarks by Cohen (1988) for small, medium, and large Cohen's f values are 0.1, 0.25, and 0.4, which correspond to eta-squared values of small (.0099), medium (.0588), and large (.1379), in line with $d = .2$, $.5$, or $.8$. So, at least based on these benchmarks, we have 90% power to detect effects that are slightly below a medium effect benchmark.

```
f2 <- f^2
ES <- f2 / (f2 + 1)
ES
```

```
## [1] 0.0503911
```

Expressed in eta-squared, we can detect values of eta-squared = 0.05 or larger.

```
mu <- mu_from_ES(K = K, ES = ES)
mu <- mu * sd
mu
```

```
## [1] -1.474295 1.474295
```

Table 3.1: Simulated ANOVA Result

	power	effect_size
anova_condition	90.14	0.0556411

Table 3.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
condition	90.01	0.0509	0.2315	10.613

We can compute a pattern of means, given a standard deviation of 6.4, that would give us an effect size of $f = 0.23$, or eta-squared of 0.05. We should be able to accomplish this if the means are -1.474295 and 1.474295. We can use these values to confirm the ANOVA has 90% power.

```
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)

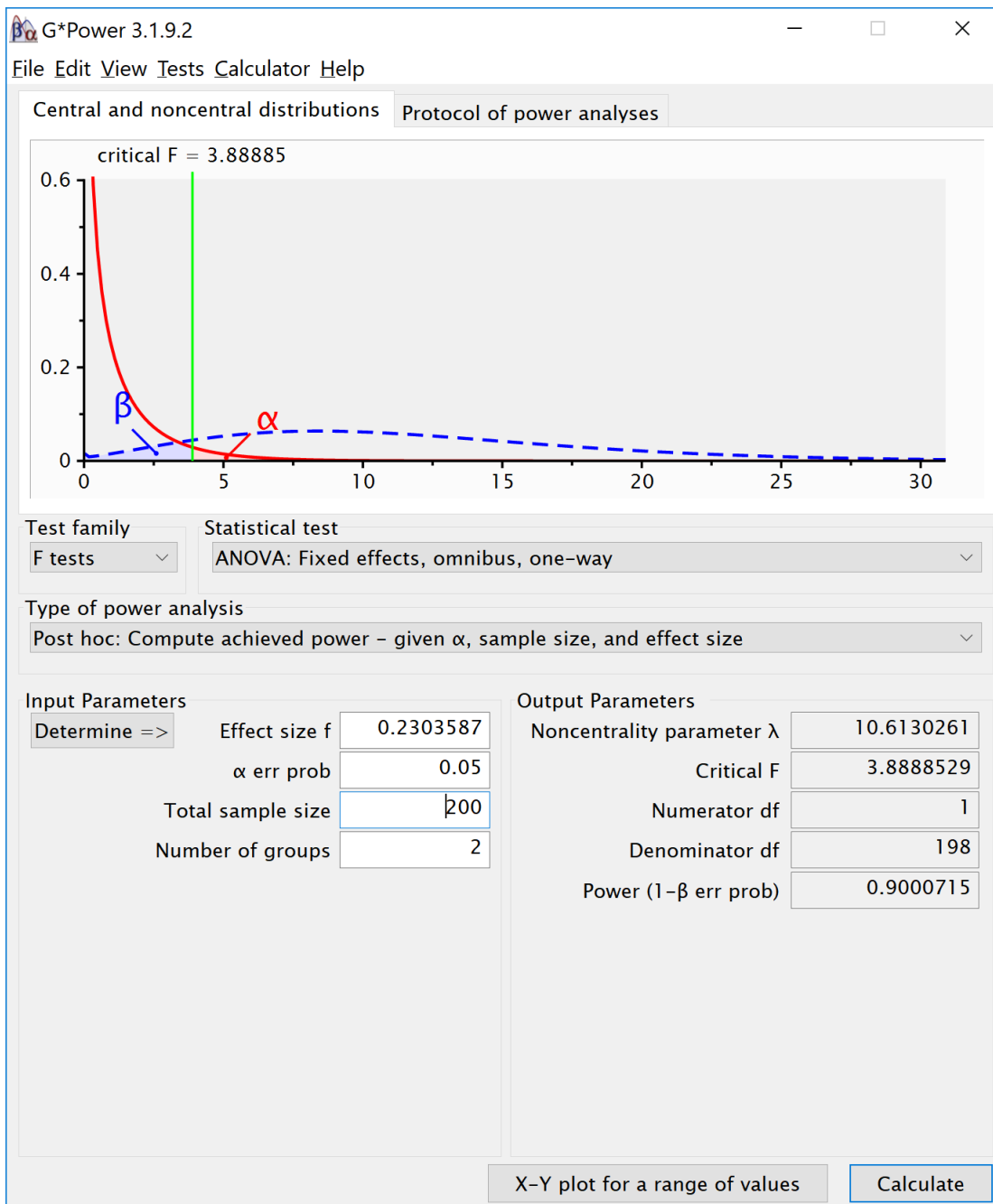
simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")
```

The simulation confirms that for the F -test for the ANOVA we have 90% power. This is also what g^* power tells us what would happen based on a post-hoc power analysis with an f of 0.2303587, 2 groups, 200 participants in total (100 in each between subject condition), and an alpha of 5%.



If we return to our expected means, how many participants do we need for sufficient power? Given the expected difference and standard deviation, $d = 0.34375$, and $f = 0.171875$. We can perform an a-priori power analysis for this simple case, which tells us we need 179 participants in each group (we can't split people in parts, and thus always round a power analysis upward), or 358 in total.

```
K <- 2
power <- 0.9
f <- 0.171875
pwr.anova.test(power = power,
               k = K,
               f = f,
               sig.level = alpha_level)
```

Table 3.3: Simulated ANOVA Result

	power	effect_size
anova_condition	89.65	0.0313263

```
##
##      Balanced one-way analysis of variance power calculation
##
##              k = 2
##              n = 178.8104
##              f = 0.171875
##      sig.level = 0.05
##              power = 0.9
##
## NOTE: n is number in each group
```

If we re-run the simulation with this sample size, we indeed have 90% power.

```
string <- "2b"
n <- 179
mu <- c(24, 26.2)
# Enter means in the order that matches the labels below.
# In this case, control, pet.
sd <- 6.4
labelnames <- c("condition", "control", "pet") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)
alpha_level <- 0.05

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

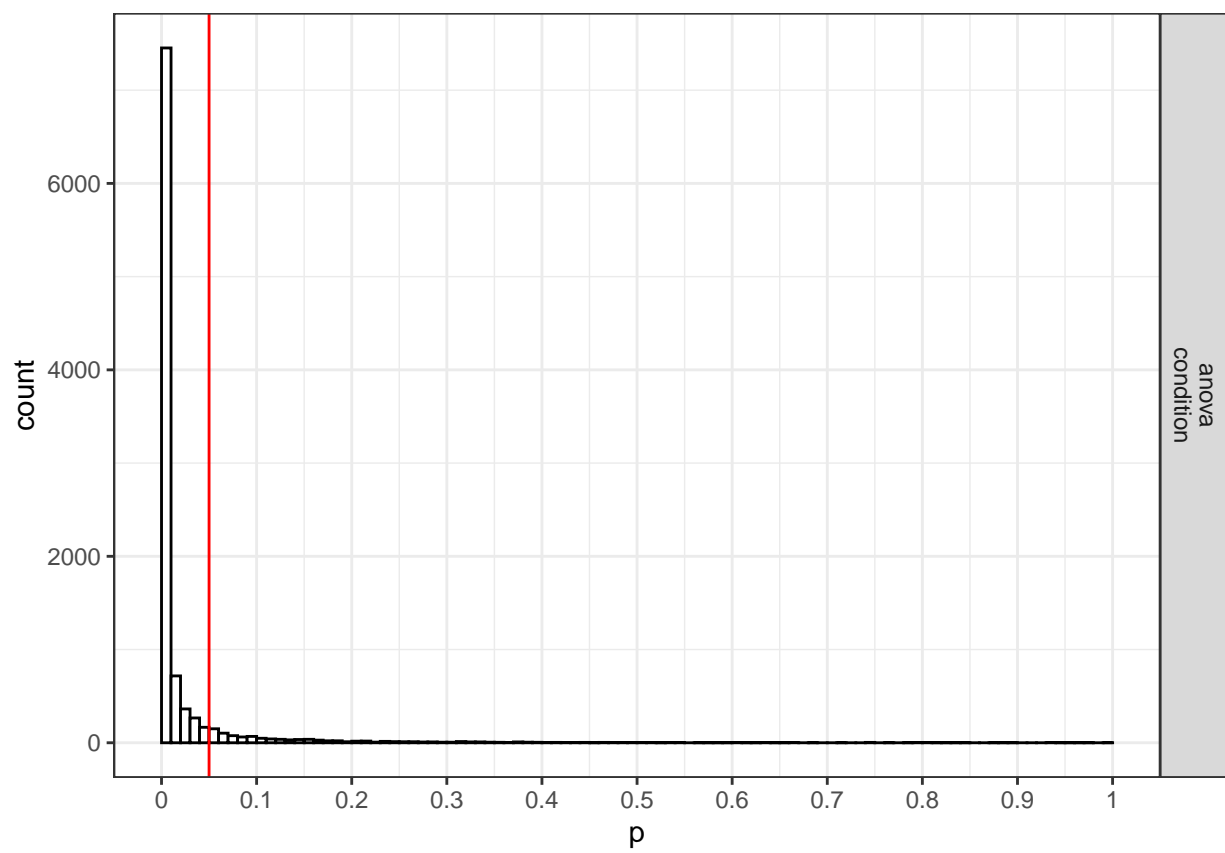
```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

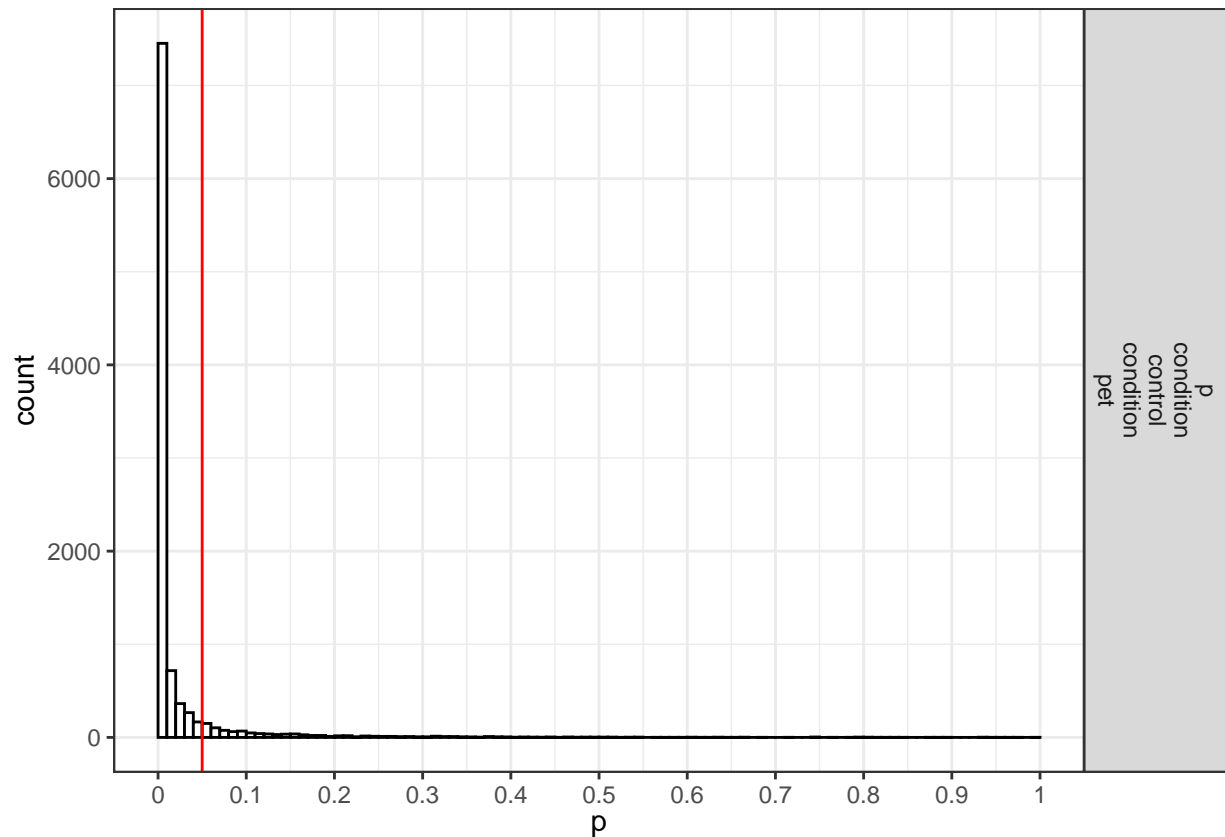
We stored the result from the power analysis in an object. This allows us to request plots (which are not printed automatically) showing the p -value distribution. If we request `simulation_result$plot1` we get the p -value distribution for the ANOVA:

Table 3.4: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
condition	90.03	0.0288	0.1724	10.5757



If we request `simulation_result$plot2` we get the p-value distribution for the paired comparisons (in this case only one):



3.2 Part 2

3.2.1 Three conditions

Imagine we aim to design a study to test the hypothesis that giving people a pet to take care of will increase their life satisfaction. We have a control condition, a ‘cat’ pet condition, and a ‘dog’ pet condition.

We can simulate a One-Way ANOVA with a specified alpha, sample size, and effect size, on see the statistical power we would have for the ANOVA and the follow-up comparisons. We expect all pets to increase life-satisfaction compared to the control condition. Obviously, we also expect the people who are in the ‘dog’ pet condition to have even greater life-satisfaction than people in the ‘cat’ pet condition. Based on work by Pavot and Diener (1993) we believe that we can expect responses on the life-satisfaction scale to have a mean of approximately 24 in our population, with a standard deviation of 6.4. We expect having a pet increases life satisfaction with approximately 2.2 scale points for participants who get a cat, and 2.6 scale points for participants who get a dog. We initially consider collecting data from 150 participants in total, with 50 participants in each condition. But before we proceed with the data collection, we examine the statistical power our design would have to detect the differences we predict.

```
string <- "3b"
n <- 50
# We are thinking of running 50 people in each condition
mu <- c(24, 26.2, 26.6)
# Enter means in the order that matches the labels below.
# In this case, control, cat, dog.
sd <- 6.4
```

Table 3.5: Simulated ANOVA Result

	power	effect_size
anova_condition	47.86	0.0440171

Table 3.6: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
condition	47.69	0.0315	0.1804	4.7852

```

labelnames <- c("condition", "control", "cat", "dog") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)
alpha_level <- 0.05
# You should think carefully about how to justify your alpha level.
# We will give some examples later, but for now, use 0.05.
simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

```

```

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

The result shows that you would have quite low power with 50 participants, both for the overall ANOVA (just around 50% power), as for the follow up comparisons (approximately 40% power for the control vs cat condition, around 50% for the control vs dogs condition, and a really low power (around 6%, just above the Type 1 error rate of 5%) for the expected difference between cats and dogs.

3.2.2 Power for simple effects

We are typically not just interested in the ANOVA, but also in follow up comparisons. In this case, we would perform a *t*-test comparing the control condition against the cat and dog condition, and we would compare the cat and dog conditions against each other, in independent *t*-tests.

For our example, Cohen's *d* (the standardized mean difference) is 2.2/6.4, or $d = 0.34375$ for the difference between the control condition and cats, 2.6/6.4 of $d = 0.40625$ for the difference between the control condition and dogs, and 0.4/6.4 or $d = 0.0625$ for the difference between cats and dogs as pets.

We can easily compute the expected power for these simple comparisons using the *pwr* package.

```
pwr.t.test(
  d = 2.2 / 6.4,
  n = 50,
  sig.level = 0.05,
  type = "two.sample",
  alternative = "two.sided"
)$power
```

```
## [1] 0.3983064
```

```
pwr.t.test(
  d = 2.6 / 6.4,
  n = 50,
  sig.level = 0.05,
  type = "two.sample",
  alternative = "two.sided"
)$power
```

```
## [1] 0.5205162
```

```
pwr.t.test(
  d = 0.4 / 6.4,
  n = 50,
  sig.level = 0.05,
  type = "two.sample",
  alternative = "two.sided"
)$power
```

```
## [1] 0.06104044
```

This analysis tells us that running the study with 50 participants in each condition is more likely to *not* yield a significant test result, even if our expected pattern of differences is true, than that we will observe a p -value smaller than our alpha level. This is not optimal.

Let's mathematically explore which pattern of means we would need to expect to have 90% power for the ANOVA with 50 participants in each group. We can use the `pwr` package in R to compute a sensitivity analysis that tells us the effect size, in Cohen's f , that we are able to detect with 3 groups and 50 participants in each group, in order to achieve 90% power with an alpha level of 5%.

```
K <- 3
n <- 50
sd <- 6.4
r <- 0
#Calculate f when running simulation
f <- pwr.anova.test(n = n,
  k = K,
  power = 0.9,
  sig.level = alpha_level)$f
f
```

```
## [1] 0.2934417
```

Table 3.7: Simulated ANOVA Result

	power	effect_size
anova_condition	89.41	0.0908705

This sensitivity analysis shows we have 90% power in our planned design to detect effects of Cohen's f of 0.2934417. Benchmarks by Cohen (1988) for small, medium, and large Cohen's f values are 0.1, 0.25, and 0.4, which correspond to eta-squared values of small (.0099), medium (.0588), and large (.1379), in line with $d = .2$, $.5$, or $.8$. So, at least based on these benchmarks, we have 90% power to detect effects that are somewhat sizeable.

```
f2 <- f^2
ES <- f2 / (f2 + 1)
ES
```

```
## [1] 0.07928127
```

Expressed in eta-squared, we can detect values of eta-squared = 0.0793 or larger.

```
mu <- mu_from_ES(K = K, ES = ES)
mu <- mu * sd
mu
```

```
## [1] -2.300104 0.000000 2.300104
```

We can compute a pattern of means, given a standard deviation of 6.4, that would give us an effect size of $f = 0.2934$, or eta-squared of 0.0793. We should be able to accomplish this if the means are -2.300104, 0.000000, and 2.300104. We can use these values to confirm the ANOVA has 90% power.

```
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")
```

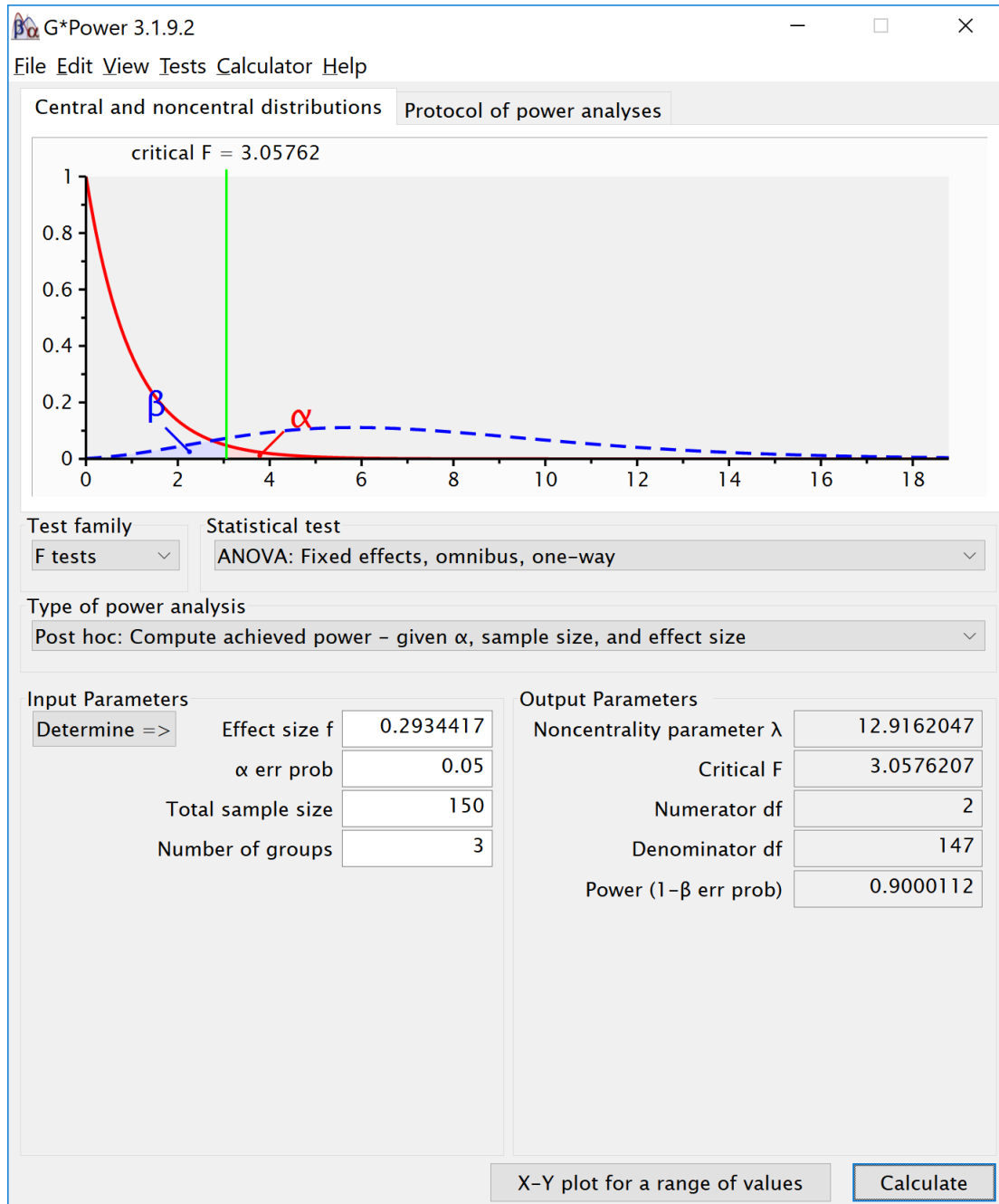
```
exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")
```

Table 3.8: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
condition	90	0.0808	0.2964	12.9162

The simulation confirms that for the F -test for the ANOVA we have 90% power. This is also what g*power tells us what would happen based on a post-hoc power analysis with an f of 0.2934417, 3 groups, 150 participants in total (50 in each between subject condition), and an alpha of 5%.



We can also compute the power for the ANOVA and simple effects in R with the pwr package. The calculated effect sizes and power match those from the simulation.


```
K <- 3
n <- 50
sd <- 6.4
f <- 0.2934417
pwr.anova.test(
  n = n,
  k = K,
  f = f,
  sig.level = alpha_level
)$power
```

```
## [1] 0.9000112
```

```
d <- 2.300104 / 6.4
d
```

```
## [1] 0.3593912
```

```
pwr.t.test(
  d = 2.300104 / 6.4,
  n = 50,
  sig.level = 0.05,
  type = "two.sample",
  alternative = "two.sided"
)$power
```

```
## [1] 0.4284243
```

```
d <- 2 * 2.300104 / 6.4
d
```

```
## [1] 0.7187825
```

```
pwr.t.test(
  d = d,
  n = 50,
  sig.level = 0.05,
  type = "two.sample",
  alternative = "two.sided"
)$power
```

```
## [1] 0.9450353
```

We can also compare the results against the analytic solution by Aberson (2019).

First, load the function for a 3-way ANOVA from the `pwr2pp1` package.

Then we use the function to calculate power.

```
#Initial example, low power
```

```
anova1f_3(
m1 = 24,
m2 = 26.2,
m3 = 26.6,
s1 = 6.4,
s2 = 6.4,
s3 = 6.4,
n1 = 50,
n2 = 50,
n3 = 50,
alpha = .05
)
```

```
## Sample size overall = 150
```

```
## Power = 0.4769 for eta-squared = 0.0315
```

```
#From: Aberson, Christopher L. Applied Power Analysis for the Behavioral Sciences, 2nd Edition.
```

```
# $Power [1] 0.4769468
```

```
#Later example, based on larger mean difference
```

```
anova1f_3(
m1 = -2.300104,
m2 = 0,
m3 = 2.300104,
s1 = 6.4,
s2 = 6.4,
s3 = 6.4,
n1 = 50,
n2 = 50,
n3 = 50,
alpha = .05
)
```

```
## Sample size overall = 150
```

```
## Power = 0.9 for eta-squared = 0.0808
```

```
# $Power [1] 0.9000112
```

3.3 Part 3

We first repeat the simulation by Brysbaert:

```
# Simulations to estimate the power of an ANOVA with three unrelated groups
# the effect between the two extreme groups is set to d = .4,
# the effect for the third group is d = .4 (see below for other situations)
# we use the built-in aov-test command
# give sample sizes (all samples sizes are equal)
N = 90
```

```

# give effect size d
d1 = .4 # difference between the extremes
d2 = .4 # third condition goes with the highest extreme
# give number of simulations
nSim = nsims
# give alpha levels
# alpha level for the omnibus ANOVA
alpha1 = .05
# alpha level for three post hoc one-tailed t-tests Bonferroni correction
alpha2 = .05

# create vectors to store p-values
p1 <- numeric(nSim) #p-value omnibus ANOVA
p2 <- numeric(nSim) #p-value first post hoc test
p3 <- numeric(nSim) #p-value second post hoc test
p4 <- numeric(nSim) #p-value third post hoc test
pes1 <- numeric(nSim) #partial eta-squared
pes2 <- numeric(nSim) #partial eta-squared two extreme conditions
library(lsr)
for (i in 1:nSim) {
  #for each simulated experiment

  x <- rnorm(n = N, mean = 0, sd = 1)
  y <- rnorm(n = N, mean = d1, sd = 1)
  z <- rnorm(n = N, mean = d2, sd = 1)
  data = c(x, y, z)
  groups = factor(rep(letters[24:26], each = N))
  test <- aov(data ~ groups)
  pes1[i] <- etaSquared(test)[1, 2]
  p1[i] <- summary(test)[[1]][["Pr(>F)"]][[1]]
  p2[i] <- t.test(x, y)$p.value
  p3[i] <- t.test(x, z)$p.value
  p4[i] <- t.test(y, z)$p.value
  data = c(x, y)
  groups = factor(rep(letters[24:25], each = N))
  test <- aov(data ~ groups)
  pes2[i] <- etaSquared(test)[1, 2]
}

# results are as predicted when omnibus ANOVA is significant,
# t-tests are significant between x and y plus x and z;
# not significant between y and z
# printing all unique tests (adjusted code by DL)
sum(p1 < alpha1) / nSim

```

```
## [1] 0.7915
```

```
sum(p2 < alpha2) / nSim
```

```
## [1] 0.7539
```

Table 3.9: Simulated ANOVA Result

	power	effect_size
anova_factor1	78.98	0.0411885

```
sum(p3 < alpha2) / nSim
```

```
## [1] 0.7601
```

```
sum(p4 < alpha2) / nSim
```

```
## [1] 0.0489
```

```
mean(pes1)
```

```
## [1] 0.04141889
```

```
mean(pes2)
```

```
## [1] 0.04353862
```

3.3.1 Three conditions replication

```
K <- 3
mu <- c(0, 0.4, 0.4)
n <- 90
sd <- 1
r <- 0
design = paste(K, "b", sep = "")
```

```
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = c("factor1", "level1", "level2", "level3")
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")
```

Table 3.10: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
factor1	79.37	0.0347	0.1896	9.6

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

3.3.2 Variation 1

```
# give sample sizes (all samples sizes are equal)
N = 145
# give effect size d
d1 = .4 #difference between the extremes
d2 = .0 #third condition goes with the highest extreme
# give number of simulations
nSim = nsims
# give alpha levels
#alpha level for the omnibus ANOVA
alpha1 = .05
#alpha level for three post hoc one-tailed t-tests Bonferroni correction
alpha2 = .05

# create vectors to store p-values
p1 <- numeric(nSim) #p-value omnibus ANOVA
p2 <- numeric(nSim) #p-value first post hoc test
p3 <- numeric(nSim) #p-value second post hoc test
p4 <- numeric(nSim) #p-value third post hoc test
pes1 <- numeric(nSim) #partial eta-squared
pes2 <- numeric(nSim) #partial eta-squared two extreme conditions
library(lsr)
for (i in 1:nSim) {
  #for each simulated experiment

  x <- rnorm(n = N, mean = 0, sd = 1)
  y <- rnorm(n = N, mean = d1, sd = 1)
  z <- rnorm(n = N, mean = d2, sd = 1)
  data = c(x, y, z)
  groups = factor(rep(letters[24:26], each = N))
  test <- aov(data ~ groups)
  pes1[i] <- etaSquared(test)[1, 2]
  p1[i] <- summary(test)[[1]][["Pr(>F)"]][[1]]
  p2[i] <- t.test(x, y)$p.value
  p3[i] <- t.test(x, z)$p.value
  p4[i] <- t.test(y, z)$p.value
  data = c(x, y)
```

```
groups = factor(rep(letters[24:25], each = N))
test <- aov(data ~ groups)
pes2[i] <- etaSquared(test)[1, 2]
}

# results are as predicted when omnibus ANOVA is significant,
# t-tests are significant between x and y plus x and z;
# not significant between y and z
# printing all unique tests (adjusted code by DL)
sum(p1 < alpha1) / nSim
```

```
## [1] 0.9492
```

```
sum(p2 < alpha2) / nSim
```

```
## [1] 0.9232
```

```
sum(p3 < alpha2) / nSim
```

```
## [1] 0.0511
```

```
sum(p4 < alpha2) / nSim
```

```
## [1] 0.9218
```

```
mean(pes1)
```

```
## [1] 0.03875397
```

```
mean(pes2)
```

```
## [1] 0.04195249
```

3.3.3 Three conditions replication

```
K <- 3
mu <- c(0, 0.4, 0.0)
n <- 145
sd <- 1
r <- 0
design = paste(K, "b", sep = "")
```

```
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
```

Table 3.11: Simulated ANOVA Result

	power	effect_size
anova_factor1	94.97	0.0389566

Table 3.12: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centralilty
factor1	94.89	0.0346	0.1892	15.4667

```
sd = sd,
labelnames = c("factor1", "level1", "level2", "level3")
)

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

3.3.4 Variation 2

```
# give sample sizes (all samples sizes are equal)
N = 82
# give effect size d
d1 = .4 #difference between the extremes
d2 = .2 #third condition goes with the highest extreme
# give number of simulations
nSim = nsims
# give alpha levels
#alpha level for the omnibus ANOVA
alpha1 = .05
#alpha level for three post hoc one-tailed t-tests Bonferroni correction
alpha2 = .05

# create vectors to store p-values
p1 <- numeric(nSim) #p-value omnibus ANOVA
p2 <- numeric(nSim) #p-value first post hoc test
p3 <- numeric(nSim) #p-value second post hoc test
p4 <- numeric(nSim) #p-value third post hoc test
pes1 <- numeric(nSim) #partial eta-squared
```

```

pes2 <- numeric(nSim) #partial eta-squared two extreme conditions
library(lsr)
for (i in 1:nSim) {
  #for each simulated experiment

  x <- rnorm(n = N, mean = 0, sd = 1)
  y <- rnorm(n = N, mean = d1, sd = 1)
  z <- rnorm(n = N, mean = d2, sd = 1)
  data = c(x, y, z)
  groups = factor(rep(letters[24:26], each = N))
  test <- aov(data ~ groups)
  pes1[i] <- etaSquared(test)[1, 2]
  p1[i] <- summary(test)[[1]][["Pr(>F)"]][[1]]
  p2[i] <- t.test(x, y)$p.value
  p3[i] <- t.test(x, z)$p.value
  p4[i] <- t.test(y, z)$p.value
  data = c(x, y)
  groups = factor(rep(letters[24:25], each = N))
  test <- aov(data ~ groups)
  pes2[i] <- etaSquared(test)[1, 2]
}

sum(p1 < alpha1) / nSim

```

```
## [1] 0.616
```

```
sum(p2 < alpha2) / nSim
```

```
## [1] 0.7188
```

```
sum(p3 < alpha2) / nSim
```

```
## [1] 0.2396
```

```
sum(p4 < alpha2) / nSim
```

```
## [1] 0.2488
```

```
mean(pes1)
```

```
## [1] 0.0337234
```

```
mean(pes2)
```

```
## [1] 0.04407838
```

3.3.5 Three conditions replication

Table 3.13: Simulated ANOVA Result

	power	effect_size
anova_factor1	61.15	0.0336575

Table 3.14: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
factor1	61.94	0.0263	0.1643	6.56

```
K <- 3
mu <- c(0, 0.4, 0.2)
n <- 82
sd <- 1
design = paste(K, "b", sep = "")
```

```
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = c("factor1", "level1", "level2", "level3")
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")
```

3.4 Effect Size Estimates for One-Way ANOVA

Using the formulas below, we can calculate the means for between designs with one factor (one-way ANOVA). Using the formula also used in Albers & Lakens (2018), we can determine the means that should yield a specified effect sizes (expressed in Cohen's f).

Eta-squared (identical to partial eta-squared for One-Way ANOVA's) has benchmarks of .0099, .0588, and .1379 for small, medium, and large effect sizes (Cohen, 1988).

Table 3.15: Simulated ANOVA Result

	power	effect_size
anova_factor1	100	0.0100534

Table 3.16: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
factor1	100	0.0099	0.1	149.9849

3.4.1 Three conditions, small effect size

We can simulate a one-factor anova setting means to achieve a certain effect size. Eta-squared is biased. Thus, the eta-squared we calculate based on the observed data overestimates the population effect size. This bias is largest for smaller sample sizes. Thus, to test whether the simulation yields the expected effect size, we use extreme large sample sizes in each between subject condition ($n = 5000$). This simulation should yield a small effect size (0.099)

```
K <- 3
ES <- .0099
mu <- mu_from_ES(K = K, ES = ES)
n <- 5000
sd <- 1
r <- 0
string = paste(K,"b",sep = "")

design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = c("factor1", "level1", "level2", "level3")
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")

exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")
```

The resulting effect size estimate from the simulation is very close to 0.0099

Table 3.17: Simulated ANOVA Result

	power	effect_size
anova_factor1	100	0.0589183

Table 3.18: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
factor1	100	0.0588	0.25	1249.469

3.4.2 Four conditions, medium effect size

This simulation should yield a medium effect size (0.588) across four independent conditions.

```
K <- 4
ES <- .0588
mu <- mu_from_ES(K = K, ES = ES)
n <- 5000
sd <- 1
r <- 0
string = paste(K,"b",sep = "")

design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = c("factor1", "level1", "level2", "level3", "level4")
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")

exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")
```

Results are very close to 0.588.

3.4.3 Two conditions, large effect size

We can simulate a one-factor anova that should yield a large effect size (0.1379) across two conditions.

Table 3.19: Simulated ANOVA Result

	power	effect_size
anova_factor1	100	0.1379042

Table 3.20: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centralilty
factor1	100	0.1379	0.4	1599.582

```
K <- 2
ES <- .1379
mu <- mu_from_ES(K = K, ES = ES)
n <- 5000
sd <- 1
r <- 0
string = paste(K,"b",sep = "")
```

```
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = c("factor1", "level1", "level2"))

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

The results are very close to is simulation should yield a small effect size (0.1379).

Chapter 4

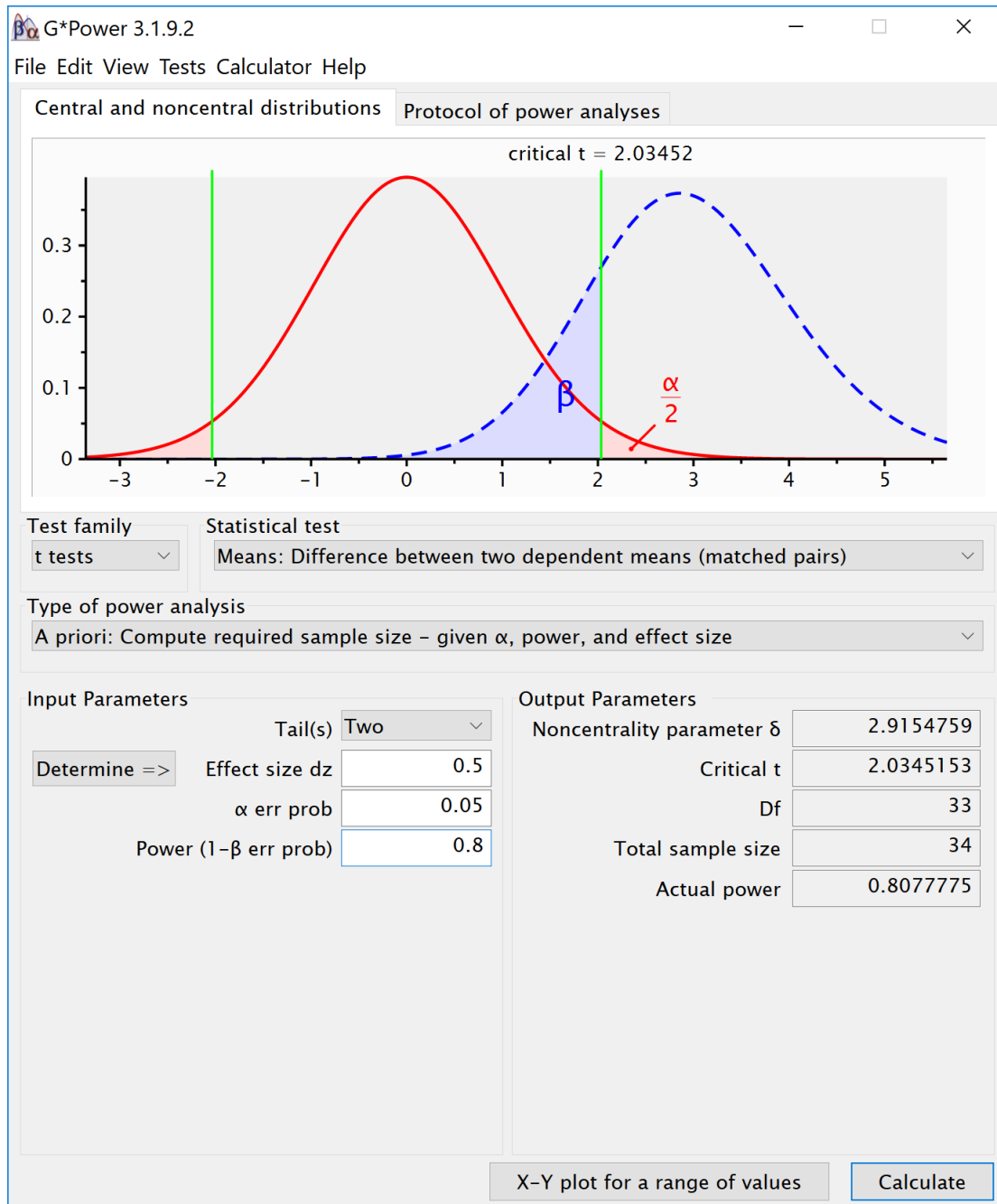
Repeated Measures ANOVA

4.1 Part 1

In a repeated measures design multiple observations are collected from the same participants. In the simplest case, where there are two repeated observations, a repeated measures ANOVA equals a dependent or paired t -test. The difference compared to a between subject design is that repeated measures can be correlated, and in psychology, they often are. Let's first explore the impact of this correlation on the power of a repeated measures ANOVA.

4.1.1 Two conditions, medium effect size

To illustrate the effect of correlated observations, we start by simulating data for a medium effect size for a dependent (or paired, or within-subject) t -test. Let's first look at G*power. If we want to perform an a-priori power analysis, we are asked to fill in the effect size d_z . As Cohen (1988) writes, "The Z subscript is used to emphasize the fact that our raw score unit is no longer X or Y, but Z", where Z are the difference scores of X-Y.



Within designs can have greater power to detect differences than between designs because the values are correlated, and a within design requires less participants because each participant provides multiple observations. One difference between an independent t -test and a dependent t -test is that an independent t -test has $2(n-1)$ degrees of freedom, while a dependent t -test has $(n-1)$ degrees of freedom. The sample size needed in a two-group within-design (NW) relative to the sample needed in two-group between-designs (NB), assuming normal distributions, and ignoring the difference in degrees of freedom between the two types of tests, is (from Maxwell & Delaney, 2004, p. 561, formula 45):

$$N_W = \frac{N_B(1-\rho)}{2}$$

The division by 2 in the equation is due to the fact that in a two-condition within design every participant provides two data-points. The extent to which this reduces the sample size compared to a between-subject design depends on the correlation (r) between the two dependent variables, as indicated by the $1-r$ part of

the equation. If the correlation is 0, a within-subject design needs half as many participants as a between-subject design (e.g., 64 instead 128 participants), simply because every participant provides 2 datapoints. The higher the correlation, the larger the relative benefit of within designs, and whenever the correlation is negative (up to -1) the relative benefit disappears.

Whereas in an independent t -test the two observations are uncorrelated, in a within design the observations are correlated. This has an effect on the standard deviation of the difference scores. In turn, because the standardized effect size is the mean difference divided by the standard deviation of the difference scores, the correlation has an effect on the standardized mean difference in a within design, Cohen's d_z . The relation, as Cohen (1988, formula 2.3.7) explains, is:

$$\sigma_z = \sigma \sqrt{2(1 - \rho)}$$

Therefore, the relation between d_z and d is $\sqrt{2(1 - \rho)}$. As Cohen (1988) writes: "In other words, a given difference between population means for matched (dependent) samples is standardized by a value which is $\sqrt{2(1 - \rho)}$ as large as would be the case were they independent. If we enter a correlation of 0.5 in the formula, we get $\sqrt{2(0.5)} = 1$. In other words, when the correlation is 0.5, $d = d_z$. When there is a strong correlation between dependent variables, for example $r = 0.9$, we get $d = d_z \sqrt{2(1 - 0.9)}$, and a d_z of 1 would be a $d = 0.45$. Reversely, $d_z = \frac{d}{\sqrt{2(1-r)}}$, so with a $r = 0.9$, a d of 1 would be a $d_z = 2.24$. Some consider this increase in d_z compared to d when observations are strongly correlated an 'inflation' when estimating effect sizes, but since the reduction in the standard deviation of the difference scores due to the correlation makes it easier to distinguish signal from noise in a hypothesis test, it leads to a clear power benefit.

```
# Check sample size formula Maxwell
# Power is pretty similar with n/2, same d (assuming r = 0.5).
# Small differences due to df = 2(n-1) vs df = n-1
pwr.t.test(d = 0.05,
           n = c(2000, 4000, 8000),
           sig.level = 0.05,
           type = "two.sample",
           alternative = "two.sided")
```

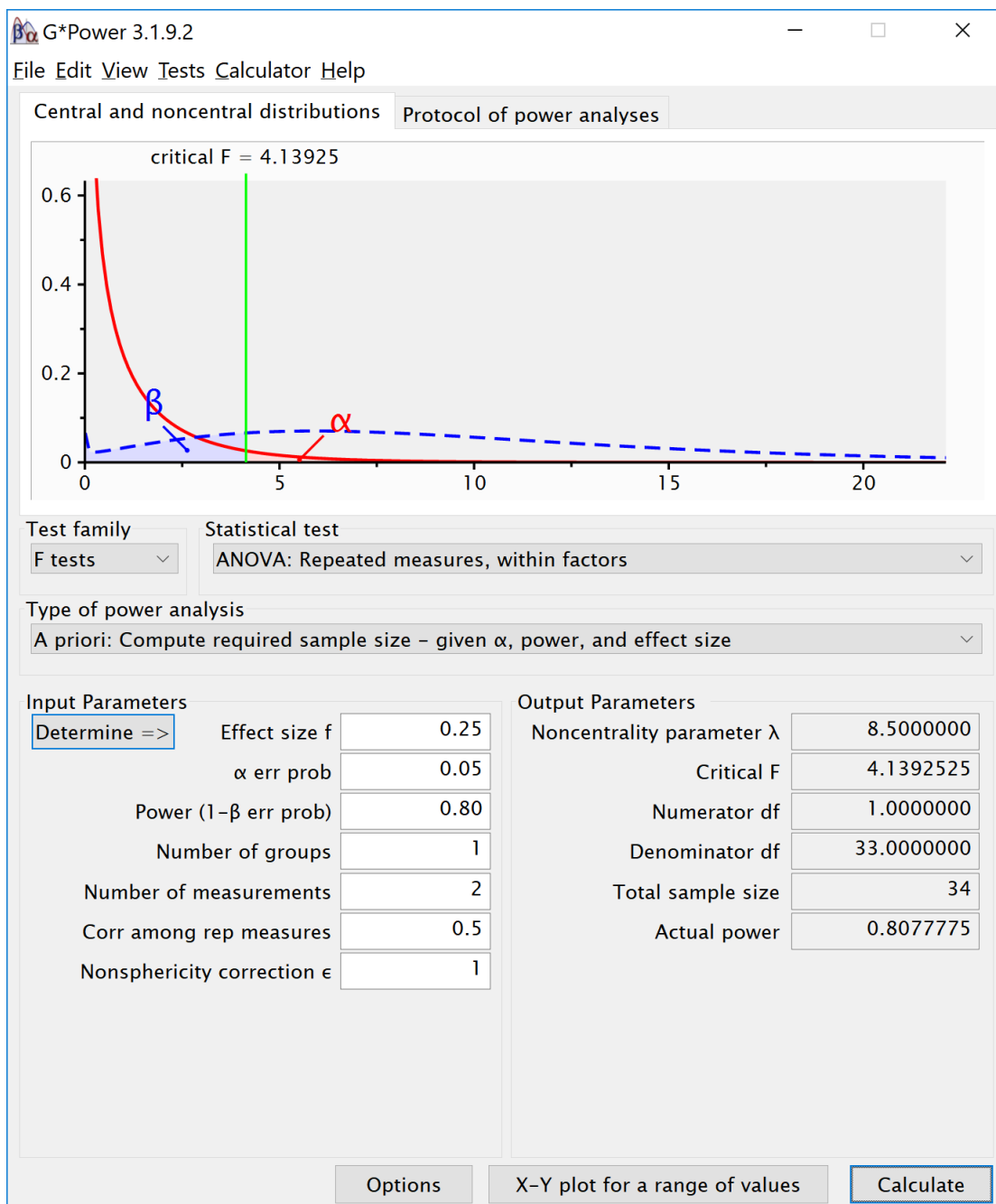
```
##
##      Two-sample t test power calculation
##
##              n = 2000, 4000, 8000
##              d = 0.05
##      sig.level = 0.05
##      power = 0.3524674, 0.6086764, 0.8853424
##      alternative = two.sided
##
## NOTE: n is number in *each* group
```

```
pwr.t.test(d = 0.05,
           n = c(1000, 2000, 4000),
           sig.level = 0.05,
           type = "paired",
           alternative = "two.sided")
```

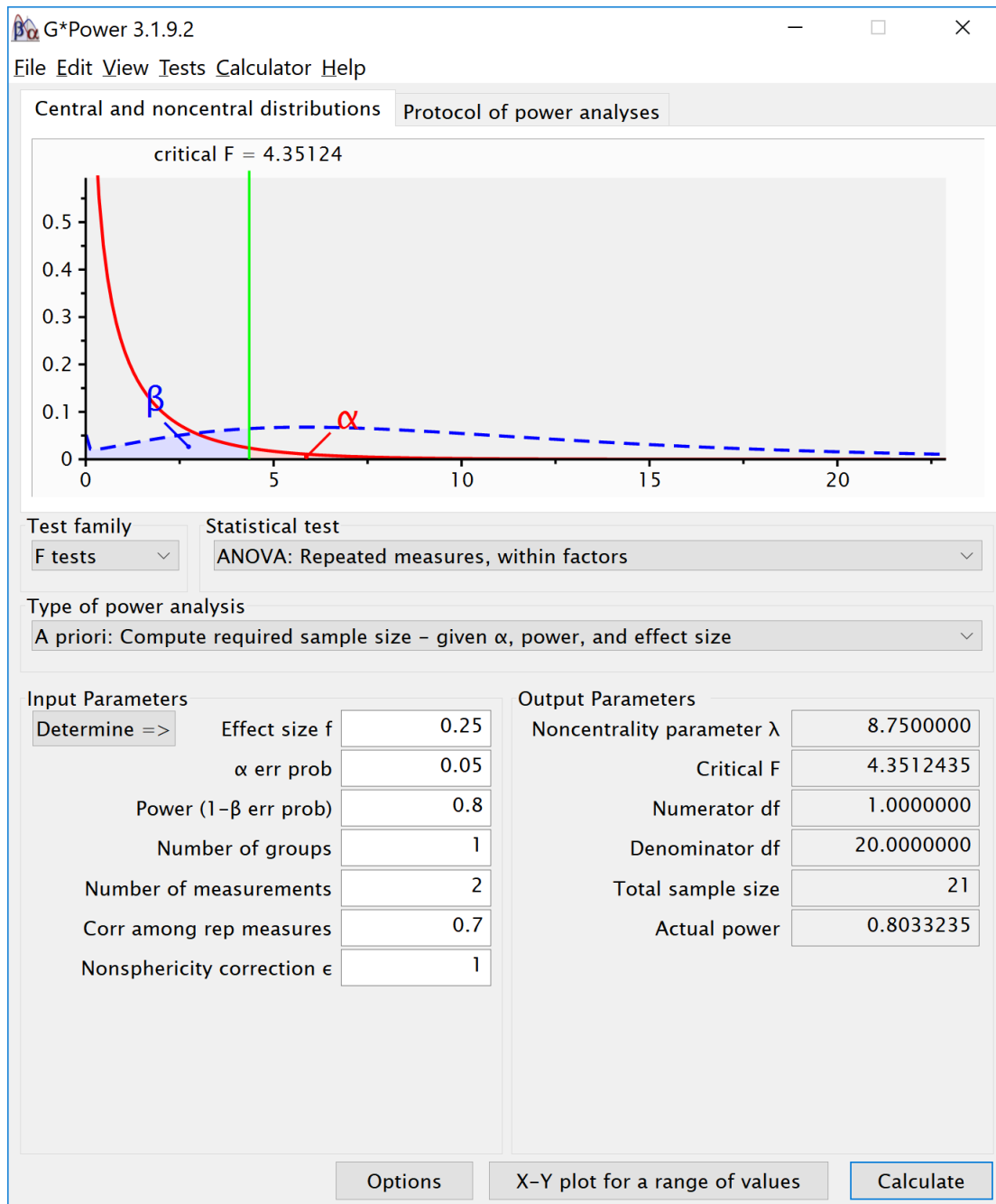
```
##
##      Paired t test power calculation
##
##              n = 1000, 2000, 4000
```

```
##          d = 0.05
##      sig.level = 0.05
##      power = 0.3520450, 0.6083669, 0.8852320
##      alternative = two.sided
##
## NOTE: n is number of *pairs*
```

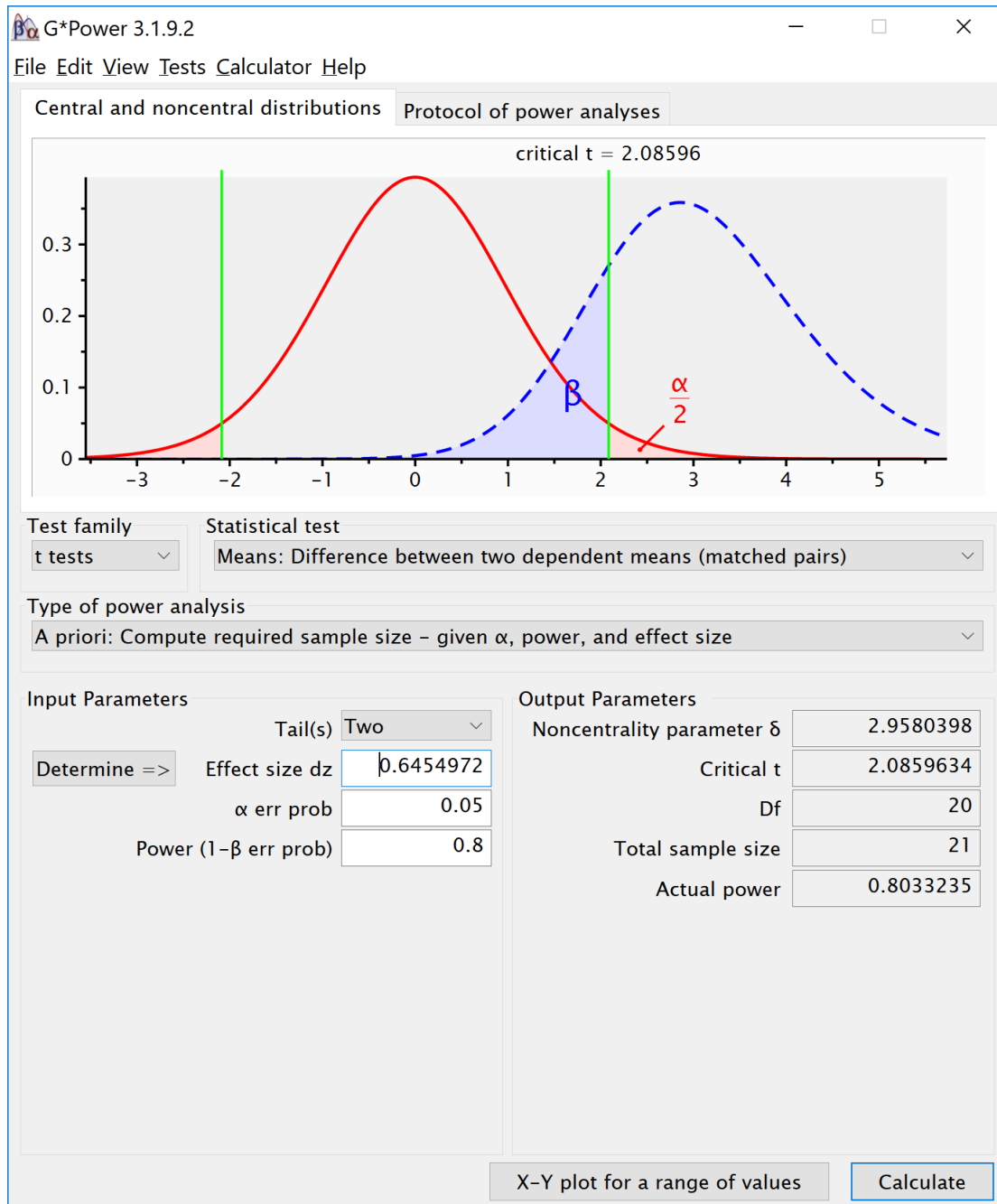
There is no equivalent f_z for Cohen's f for a within subject ANOVA. For two groups, we can directly compute Cohen's f from Cohen's d for two groups, as Cohen (1988) describes, because $f = 1/2d$. For a $d = 0.5$, $f = 0.25$. In Gpower we can run a 2 group within-subject power analysis for ANOVA. We plan for 80% power, and reproduce the analysis above for the dependent t -test. This works because the correlation is set to 0.5, when $d = d_z$, and thus the transformation of $f=1/2d$ works.



If we change the correlation to 0.7 and keep all other settings the same, the repeated measure a-priori power analysis yields a sample of 21. The correlation increases the power for the test.



To reproduce this analysis in Gpower with a dependent t -test we need to change d_z following the formula above, $d_z = \frac{0.5}{\sqrt{2(1-0.7)}}$, which yields $d_z = 0.6454972$. If we enter this value in Gpower for an a-priori power analysis, we get the exact same results (as we should, since an repeated measures ANOVA with 2 groups equals a dependent t -test). This example illustrates that the correlation between dependent variables always factors into a power analysis, both for a dependent t -test, and for a repeated measures ANOVA. Because a dependent t -test uses d_z the correlation might be less visible, but given the relation between d and d_z , the correlation is always taken into account and can greatly improve power for within designs compared to between designs.



We can perform both these power analyses using simulations as well. We set groups to 2 for the simulation, $n = 34$ (which should give 80.777 power, according to the g*power program), a correlation among repeated measures of 0.5, and an alpha of 0.05. In this case, we simulate data with means -0.25 and 0.25, and set the sd to 1. This means we have a mean difference of 0.5, and a Cohen's d of $0.5/1 = 0.5$. In the first example, we set the correlation to 0.5, and the result should be 80.77% power, and an effect size estimate of 0.5 for the simple effect. We also calculate partial eta-squared for the ANOVA, which equals $\frac{f^2}{f^2+1}$, or 0.05882353.

```
K <- 2
n <- 34
sd <- 1
r <- 0.5
```

Table 4.1: Simulated ANOVA Result

	power	effect_size
anova_speed	80.37	0.2156948

Table 4.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non centrality
speed	80.78	0.2048	0.5075	8.5

```
alpha = 0.05
f <- 0.25
f2 <- f^2
ES <- f2/(f2 + 1)
ES
```

```
## [1] 0.05882353
```

```
mu <- mu_from_ES(K = K, ES = ES)
design = paste(K,"w",sep = "")
labelnames <- c("speed", "fast", "slow")

design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

alpha_level <- 0.05

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

The results of the simulation are indeed very close to 80.777%. Note that the simulation calculates Cohen's d_z effect sizes for paired comparisons - which here given the correlation of 0.5 is also 0.5 for a medium effect size.

We should see a larger d_z if we increase the correlation, keeping the sample size the same, following the example in Gpower above. We repeat the simulation, and the only difference is a correlation between dependent variables of 0.7. This should yield an effect size $d_z = 0.6454972$.

Table 4.3: Simulated ANOVA Result

	power	effect_size
anova_speed	79.89	0.315253

```

K <- 2
n <- 21
sd <- 1
r <- 0.7
alpha = 0.05
f <- 0.25
f2 <- f^2
ES <- f2/(f2 + 1)
ES

```

```
## [1] 0.05882353
```

```

mu <- mu_from_ES(K = K, ES = ES)
design = paste(K,"w",sep = "")
labelnames <- c("speed", "fast", "slow")
design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

```

```
alpha_level <- 0.05
```

```
design_result$sigmatrix
```

```

##      fast slow
## fast  1.0  0.7
## slow  0.7  1.0

```

```

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

```

```

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

```

```

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

```

```

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

Table 4.4: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
speed	80.33	0.3043	0.6614	8.75

```
#relation dz and f for within designs
f <- 0.5*0.6454972
#
```

Entering this f in G*power, with a correlation of 0.5, yields the same as entering f = 0.25 and correlation = 0.7.

4.2 Repeated Measures-ANOVA Part 2

Here, we will examine a repeated measures experiment with 3 within-subject conditions, to illustrate how a repeated measures ANOVA extends a dependent *t*-test with 3 groups.

In the example for a two-group within design we provided a specific formula for the sample size benefit for two groups. The sample size needed in within-designs (NW) with more than 2 conditions, relative to the sample needed in between-designs (NB), assuming normal distributions and compound symmetry, and ignoring the difference in degrees of freedom between the two types of tests, is (from Maxwell & Delaney, 2004, p. 562, formula 47):

$$N_W = \frac{N_B(1-\rho)}{a}$$

Where *a* is the number of within-subject levels.

4.2.1 The relation between Cohen's f and Cohen's d

Whereas in the case of a repeated measures ANOVA with 2 groups we could explain the principles of a power analysis by comparing our test against a *t*-test and Cohen's *d*, this becomes more difficult when we have more than 2 groups. It is more useful to explain how to directly calculate Cohen's *f*, the effect size used in power analyses for ANOVA. Cohen's *f* is calculated following Cohen, 1988, formula 8.2.1 and 8.2.2:

$$f = \sqrt{\frac{\sum (\mu - \bar{\mu})^2}{\frac{N}{\sigma^2}}}$$

Imagine we have a within-subject experiment with 3 conditions. We ask people what they mood is when their alarm clock wakes them up, when they wake up naturally on a week day, and when they wake up naturally on a weekend day. Based on pilot data, we expect the means (on a 7 point validated mood scale) are 3.8, 4.2, and 4.3. The standard deviation is 0.9, and the correlation between the dependent measurements is 0.7. We can calculate Cohen's *f* for the ANOVA, and Cohen's *dz* for the contrasts:

```
mu <- c(3.8, 4.2, 4.3)
sd <- 0.9
f <- sqrt(sum((mu - mean(mu)) ^ 2) / length(mu)) / sd
#Cohen, 1988, formula 8.2.1 and 8.2.2
f
```

```
## [1] 0.2400274
```

```
r <- 0.7
(4.2 - 3.8) / 0.9 / sqrt(2 * (1 - r))
```

```
## [1] 0.5737753
```

```
(4.3 - 3.8) / 0.9 / sqrt(2 * (1 - r))
```

```
## [1] 0.7172191
```

```
(4.3 - 4.2) / 0.9 / sqrt(2 * (1 - r))
```

```
## [1] 0.1434438
```

The relation between Cohen's d or d_z and Cohen's f becomes more difficult when there are multiple groups, because the relationship depends on the pattern of the means. Cohen (1988) presents calculations for three patterns, minimal variability (for example, for 5 means: -0.25, 0, 0, 0, 0.25), medium variability (for example, for 5 means: -0.25, -0.25, 0.25, 0.25, 0.25 or -0.25, -0.25, -0.25, 0.25, 0.25). For these three patterns, formula's are available that compute Cohen's f from Cohen's d , where d is the effect size calculated for the difference between the largest and smallest mean (if the largest mean is 0.25 and the smallest mean is -0.25, $0.25 - -0.25 = 0.5$, so d is 0.5 divided by the standard deviation of 0.9). In our example, d would be $(4.3-3.8)/0.9 = 0.5555556$. If we divide this value by $\sqrt{2*(1-r)}$ we have $d_z = 0.5555556/0.7745967 = 0.7172191$.

I have created a custom function that will calculate f from d , based on a specification of one of the three patterns of means. Our pattern is most similar (but not identical) to a maximum variability pattern (two means are high, one is lower). So we could attempt to calculate f from d (0.5555556), by calculating d from the largest and smallest mean:

```
source("https://raw.githubusercontent.com/Lakens/ANOVA_power_simulation/master/calc_f_d_eta.R")
res <- calc_f_d_eta(mu = mu, sd = sd, variability = "maximum")
res$f
```

```
## [1] 0.2618914
```

```
res$d
```

```
## [1] 0.5555556
```

We see the Cohen's f value is 0.2618914 and $d = 0.5555556$. The Cohen's f is not perfectly accurate - it is assuming the pattern of means is 3.8, 4.3, 4.3, and not 3.8, 4.2, 4.3. If the means and sd is known, it is best to calculate Cohen's f directly from these values.

4.2.2 Three within conditions, medium effect size

We can perform power analyses for within designs using simulations. We set groups to 3 for the simulation, $n = 20$, and the correlation between dependent variables to 0.8. If the true effect size is $f = 0.25$, and the alpha level is 0.05, the power is 96.6%.

In this case, we simulate data with means -0.3061862, 0.0000000, and 0.3061862, and set the sd to 1.

Table 4.5: Simulated ANOVA Result

	power	effect_size
anova_speed	97.04	0.3457287

```

K <- 3
n <- 20
sd <- 1
r <- 0.8
alpha = 0.05
f <- 0.25
f2 <- f^2
ES <- f2 / (f2 + 1)
ES

## [1] 0.05882353

mu <- mu_from_ES(K = K, ES = ES)

#Cohen, 1988, formula 8.2.1 and 8.2.2
sqrt(sum((mu - mean(mu)) ^ 2) / length(mu)) / sd

## [1] 0.25

design = paste(K, "w", sep = "")
labelnames <- c("speed", "fast", "medium", "slow")
design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

alpha_level <- 0.05

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

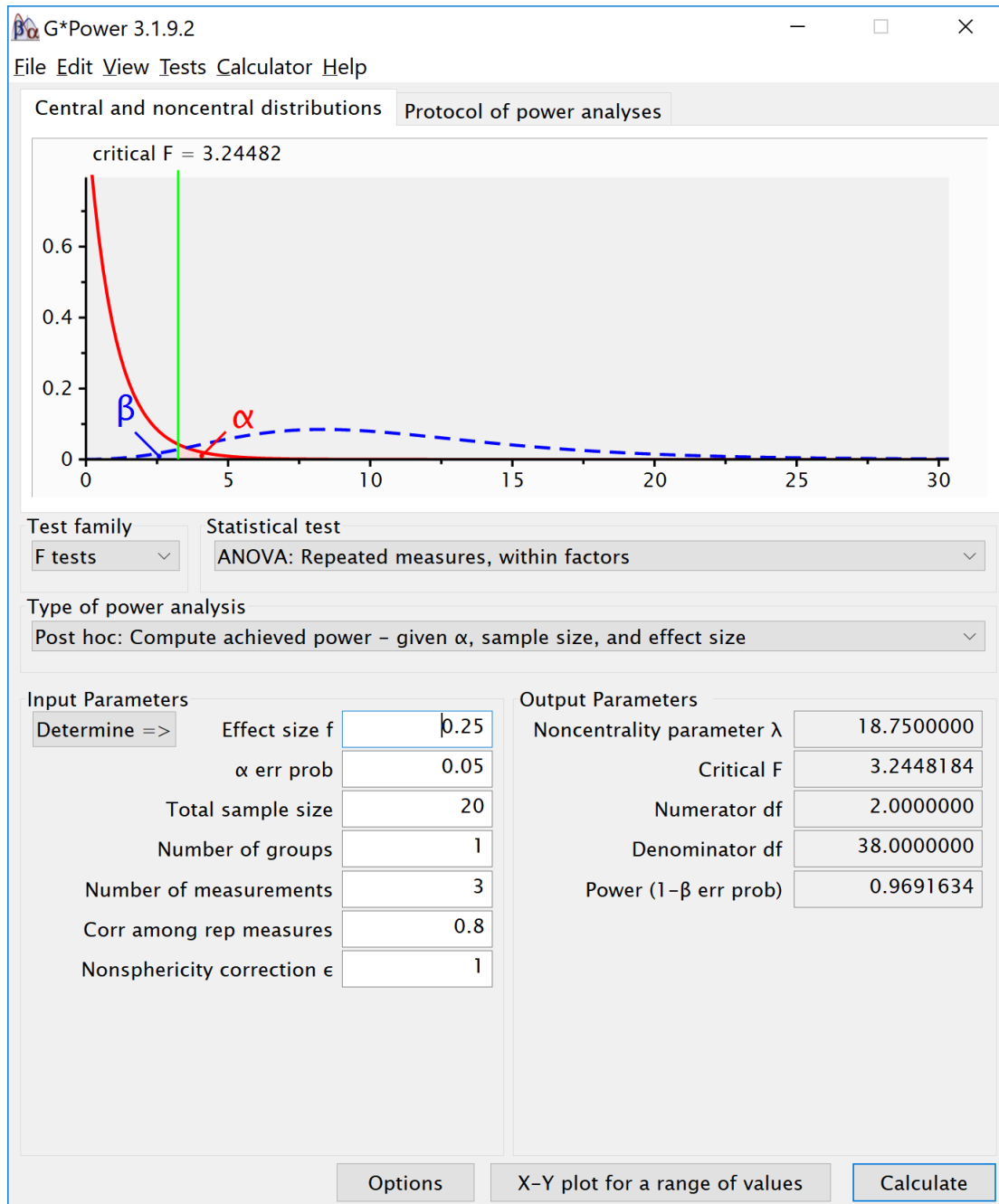
knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

The results of the simulation are indeed very close to 96.9%. We can see this is in line with the power estimate from Gpower:

Table 4.6: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
speed	96.92	0.3304	0.7024	18.75



We can also validate this by creating the code to do a power analysis in R from scratch:

```
K <- 3 #three groups
n <- 20
sd <- 1
r <- 0.8
```



```
alpha = 0.05
f <- 0.25
f2 <- f^2
ES <- f2 / (f2 + 1)
ES
```

```
## [1] 0.05882353
```

```
mu <- mu_from_ES(K = K, ES = ES)
design = paste(K, "w", sep = "")
labelnames <- c("speed", "fast", "medium", "slow")
design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

power_oneway_within(design_result)$power
```

```
## [1] 96.91634
```

```
power_oneway_within(design_result)$eta_p_2
```

```
## [1] 0.05882353
```

```
power_oneway_within(design_result)$eta_p_2_SPSS
```

```
## [1] 0.3303965
```

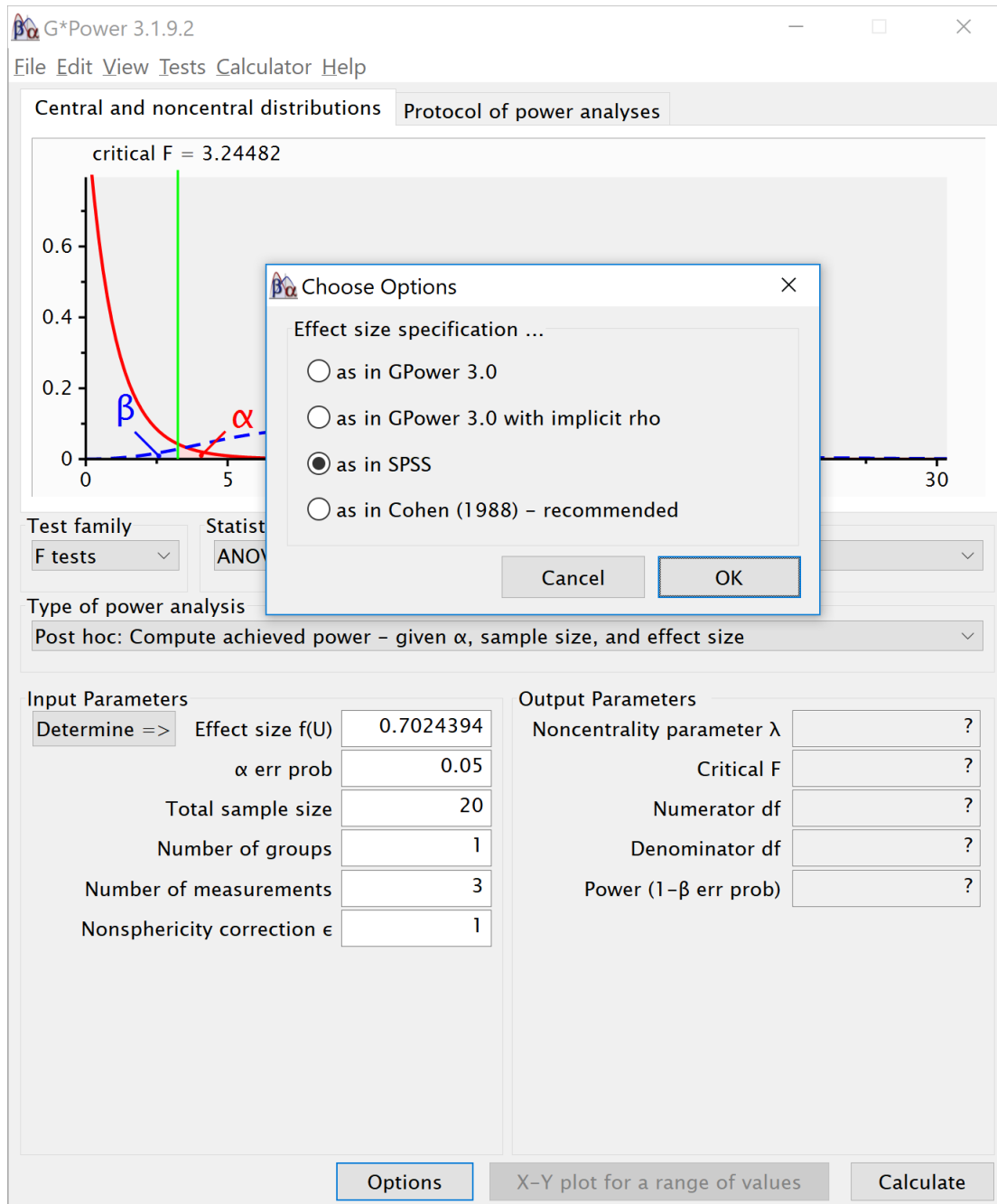
```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.25
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.7024394
```

We can even check the calculation of Cohen's f SPSS style in GPower. We take the GPower settings as illustrated above. We click the 'Options' button, and check the radiobutton next to 'As in SPSS'. Click ok, and you will notice that the 'Corr among rep measures' field has disappeared. The correlation does not need to be entered separately, but is incorporated in Cohen's f . The value of Cohen's f , which was 0.25, has changed into 0.7024394. This is the SPSS equivalent. The value is much larger. This value, and its corresponding partial eta-squared, incorporate the correlation between observations.



4.3 Part 3

We first repeat the simulation by Brysbaert:

```
# give sample size
N = 75
# give effect size d
d1 = .4 #difference between the extremes
d2 = .4 #third condition goes with the highest extreme
# give the correlation between the conditions
```

```

r = .5
# give number of simulations
nSim = nsims
# give alpha levels
alpha1 = .05 #alpha level for the omnibus ANOVA
alpha2 = .05 #also adjusted from original by DL
# create progress bar in case it takes a while
#pb <- winProgressBar(title = "progress bar", min = 0, max = nSim, width = 300)
# create vectors to store p-values
p1 <- numeric(nSim) #p-value omnibus ANOVA
p2 <- numeric(nSim) #p-value first post hoc test
p3 <- numeric(nSim) #p-value second post hoc test
p4 <- numeric(nSim) #p-value third post hoc test
# open library MASS
library('MASS')
# define correlation matrix
rho <- cbind(c(1, r, r), c(r, 1, r), c(r, r, 1))
# define participant codes
part <- paste("part", seq(1:N))
for (i in 1:nSim) {
  #for each simulated experiment
  # setWinProgressBar(pb, i, title=paste(round(i/nSim*100, 1), "% done"))
  data = mvrnorm(n = N,
    mu = c(0, 0, 0),
    Sigma = rho)
  data[, 2] = data[, 2] + d1
  data[, 3] = data[, 3] + d2
  datalong = c(data[, 1], data[, 2], data[, 3])
  conds = factor(rep(letters[24:26], each = N))
  partID = factor(rep(part, times = 3))
  output <- data.frame(partID, conds, datalong)
  test <- aov(datalong ~ conds + Error(partID / conds), data = output)
  tests <- (summary(test))
  p1[i] <- tests['Error: partID:conds'[[1]]$'Pr(>F)'[[1]]
  p2[i] <- t.test(data[, 1], data[, 2], paired = TRUE)$p.value
  p3[i] <- t.test(data[, 1], data[, 3], paired = TRUE)$p.value
  p4[i] <- t.test(data[, 2], data[, 3], paired = TRUE)$p.value
}
#close(pb)#close progress bar
#printing all unique tests (adjusted code by DL)
sum(p1 < alpha1) / nSim

```

```
## [1] 0.955
```

```
sum(p2 < alpha2) / nSim
```

```
## [1] 0.9281
```

```
sum(p3 < alpha2) / nSim
```

```
## [1] 0.9249
```

Table 4.7: Simulated ANOVA Result

	power	effect_size
anova_speed	95.1	0.1073421

Table 4.8: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
speed	95.29	0.0976	0.3288	16

```
sum(p4 < alpha2) / nSim
```

```
## [1] 0.0473
```

4.4 Reproducing Brysbaert

We can reproduce the same results as Brysbaert finds with his code:

```
design <- "3w"
n <- 75
mu <- c(0, 0.4, 0.4)
sd <- 1
r <- 0.5
labelnames <- c("speed", "fast", "medium", "slow")
```

We create the within design, and run the simulation

```
design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

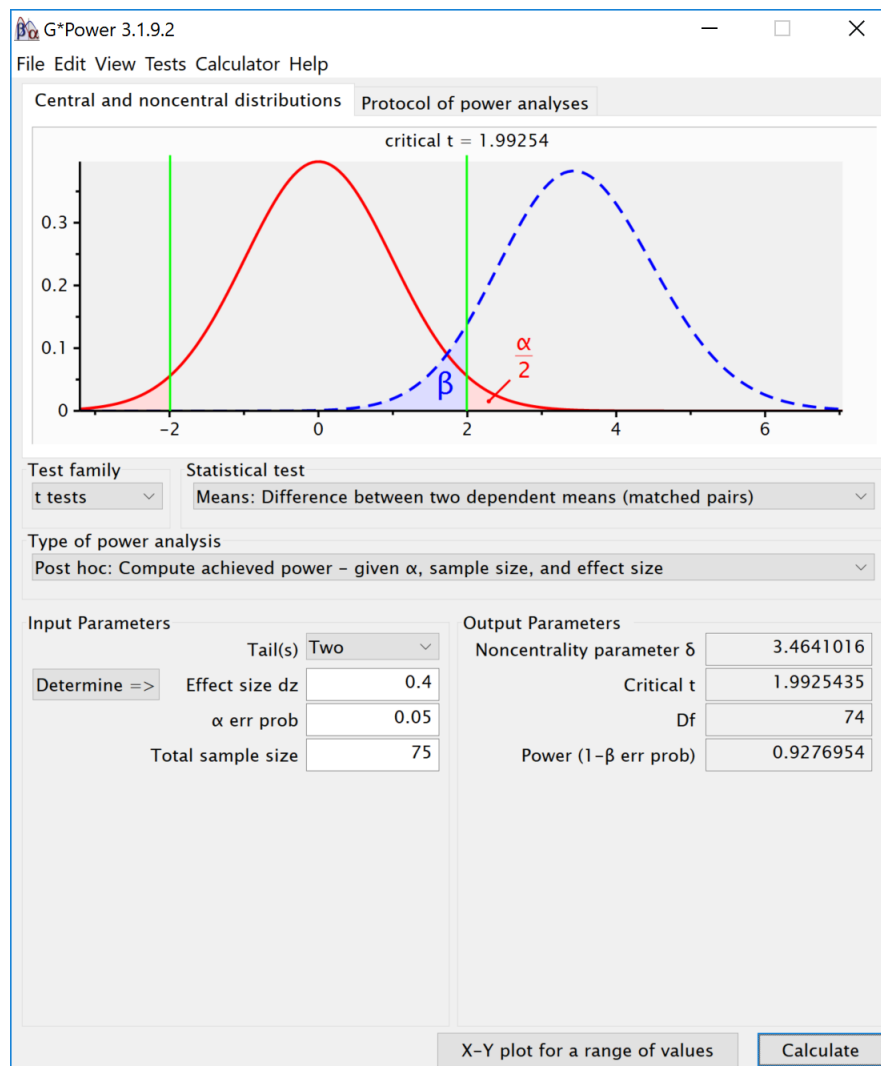
knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

4.4.1 Results

The results of the simulation are very similar. Power for the ANOVA F -test is around 95.2%. For the three paired t -tests, power is around 92.7. This is in line with the a-priori power analysis when using g^* power:



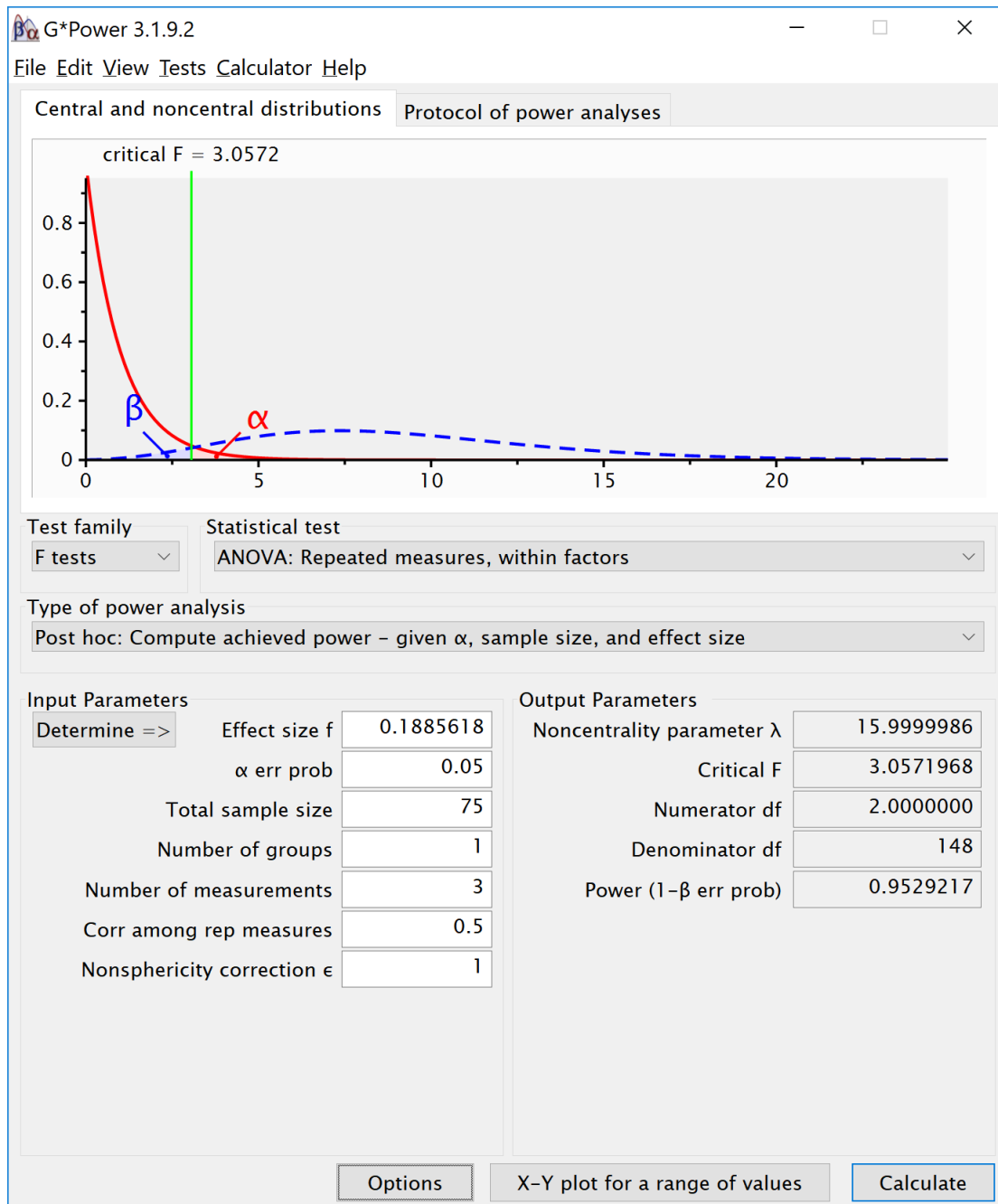
We can perform an post-hoc power analysis in G*power. We can calculate Cohen's f based on the means and sd , using our own custom formula.

```
# Our simulation is based on the following means and sd:
# mu <- c(0, 0.4, 0.4)
# sd <- 1
# Cohen, 1988, formula 8.2.1 and 8.2.2
f <- sqrt(sum((mu - mean(mu)) ^ 2) / length(mu)) / sd

# We can see why f = 0.5*d.
# Imagine 2 group, mu = 1 and 2
# Grand mean is 1.5, we have sqrt(sum(0.5^2 + 0.5^2)/2), or sqrt(0.5/2), = 0.5.
# For Cohen's d we use the difference, 2-1 = 1.
```

The Cohen's f is 0.1885618. We can enter the f (using the default 'as in G*Power 3.0' in the option

window) and enter a sample size of 75, number of groups as 1, number of measurements as 3, correlation as 0.5. This yields:



4.4.2 Reproducing Brysbaert Variation 1: Changing Correlation

```
# give sample size
N = 75
# give effect size d
d1 = .4 #difference between the extremes
d2 = .4 #third condition goes with the highest extreme
```

```

# give the correlation between the conditions
r = .6 #increased correlation
# give number of simulations
nSim = nsims
# give alpha levels
alpha1 = .05 #alpha level for the omnibus ANOVA
alpha2 = .05 #also adjusted from original by DL

# create vectors to store p-values
p1 <- numeric(nSim) #p-value omnibus ANOVA
p2 <- numeric(nSim) #p-value first post hoc test
p3 <- numeric(nSim) #p-value second post hoc test
p4 <- numeric(nSim) #p-value third post hoc test
# open library MASS
library('MASS')
# define correlation matrix
rho <- cbind(c(1, r, r), c(r, 1, r), c(r, r, 1))
# define participant codes
part <- paste("part", seq(1:N))
for (i in 1:nSim) {

  #for each simulated experiment
  # setWinProgressBar(pb, i, title=paste(round(i/nSim*100, 1), "% done"))
  data = mvrnorm(n = N,
    mu = c(0, 0, 0),
    Sigma = rho)
  data[, 2] = data[, 2] + d1
  data[, 3] = data[, 3] + d2
  datalong = c(data[, 1], data[, 2], data[, 3])
  conds = factor(rep(letters[24:26], each = N))
  partID = factor(rep(part, times = 3))
  output <- data.frame(partID, conds, datalong)
  test <- aov(datalong ~ conds + Error(partID / conds), data = output)
  tests <- (summary(test))
  p1[i] <- tests$'Error: partID:conds'[[1]]$'Pr(>F)'[[1]]
  p2[i] <- t.test(data[, 1], data[, 2], paired = TRUE)$p.value
  p3[i] <- t.test(data[, 1], data[, 3], paired = TRUE)$p.value
  p4[i] <- t.test(data[, 2], data[, 3], paired = TRUE)$p.value
}

sum(p1 < alpha1) / nSim

```

```
## [1] 0.9832
```

```
sum(p2 < alpha2) / nSim
```

```
## [1] 0.9673
```

```
sum(p3 < alpha2) / nSim
```

```
## [1] 0.9681
```

Table 4.9: Simulated ANOVA Result

	power	effect_size
anova_speed	98.37	0.1288113

Table 4.10: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
speed	98.35	0.119	0.3676	20

```
sum(p4 < alpha2) / nSim
```

```
## [1] 0.0511
```

```
design <- "3w"
n <- 75
mu <- c(0, 0.4, 0.4)
sd <- 1
r <- 0.6
labelnames <- c("speed", "fast", "medium", "slow")
```

We create the within design, and run the simulation.

```
design_result <- ANOVA_design(design = design,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

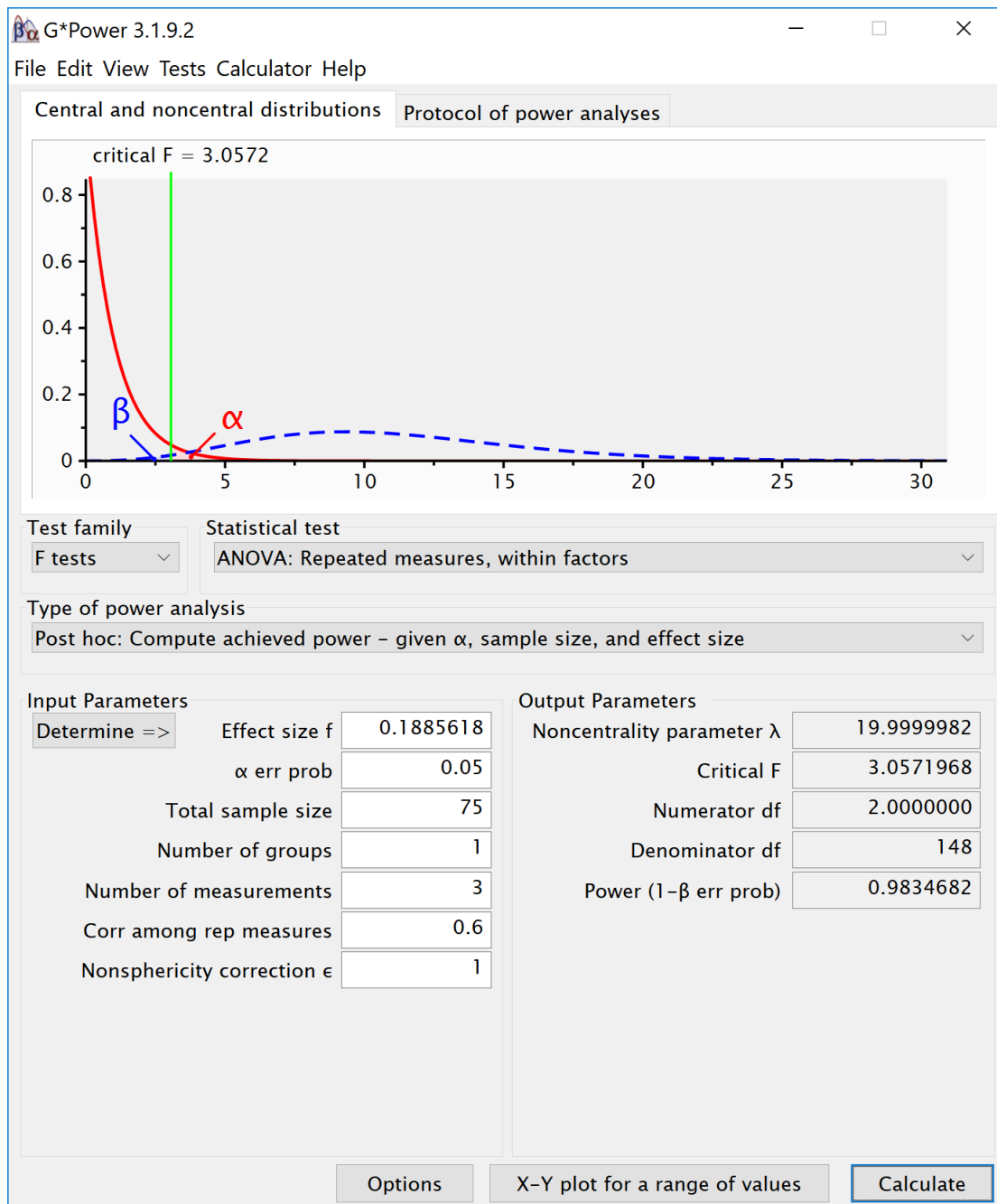
simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

Again, this is similar to G*Power for the ANOVA:



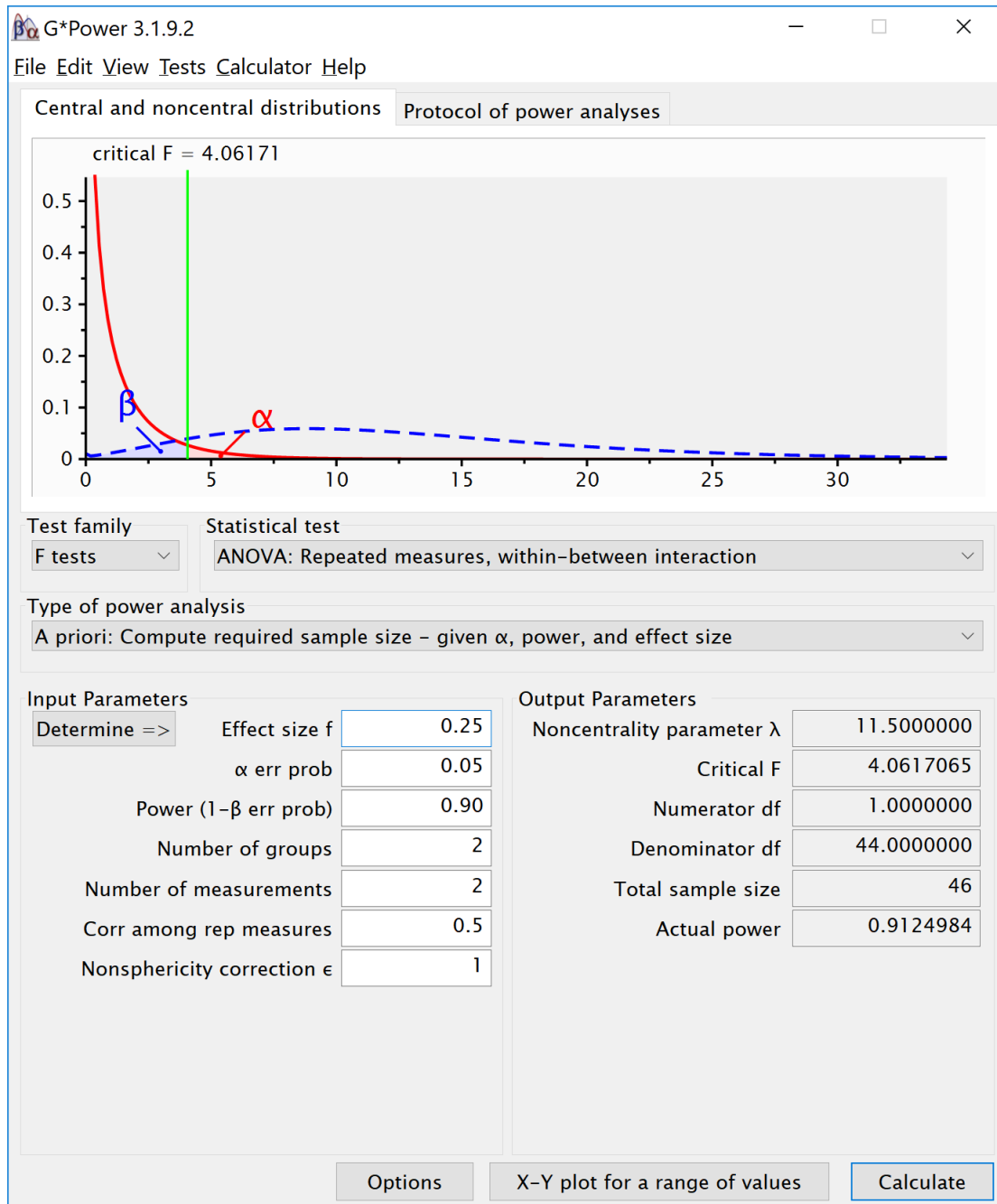
Chapter 5

Mixed ANOVA

5.1 Part 1

5.1.1 Two by two ANOVA, within-between design

We can simulate a Two-Way ANOVA with a specific alpha, sample size and effect size, to achieve a specified statistical power. We will try to reproduce the power analysis by g*power for an F-test, ANOVA: Repeated measures, within-between interaction.



For the 2-way interaction, the result should be a power of 91.25% if we have a total sample size of 46. Since we have 2 groups in the between factor that means the sample size per group is 2 (and both these groups collect 2 repeated measures).

```
mu <- c(-0.25, 0.25, 0.25, -0.25)
n <- 23
sd <- 1
r <- 0.5
string = "2w*2b"
alpha_level <- 0.05
labelnames = c("age", "old", "young", "color", "blue", "red")
```

```

design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)

simulation_result <- ANOVA_power(design_result,
  alpha_level = alpha_level,
  nsims = nsims,
  verbose = FALSE)

knitr::kable(simulation_result$main_results,
  caption = "Simulated ANOVA Result")

```

```

\begin{table}[t]

\caption{(\#tab:start_mixedANOVA)Simulated ANOVA Result}

```

	power	effect_size
anova_color	5.11	0.0226130
anova_age	5.01	0.0224623
anova_color:age	91.09	0.2154405

```
\end{table}
```

```

exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")

```

```

\begin{table}[t]

\caption{(\#tab:start_mixedANOVA)Exact ANOVA Result}

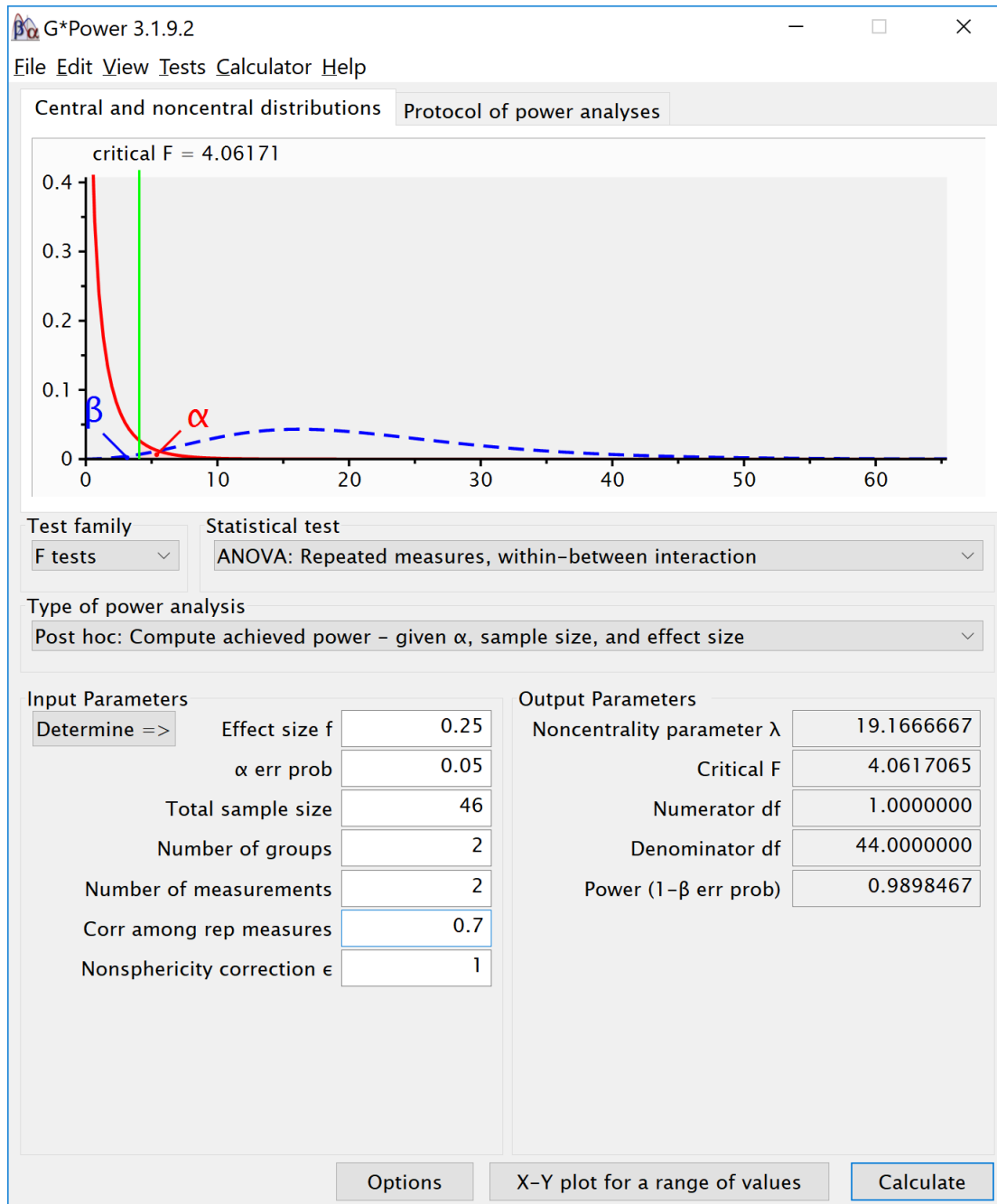
```

	power	partial_eta_squared	cohen_f	non_central
color	5.00	0.0000	0.0000	0.0
age	5.00	0.0000	0.0000	0.0
color:age	91.25	0.2072	0.5112	11.5

```
\end{table}
```

5.1.2 Two by two ANOVA, within-between design Variation 1

We can simulate the same two-way ANOVA increasing the correlation to 0.7.



```
mu <- c(-0.25, 0.25, 0.25, -0.25)
n <- 23
sd <- 1
r <- 0.7
string = "2w*2b"
alpha_level <- 0.05
labelnames = c("age", "old", "young", "color", "blue", "red")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
```

Table 5.1: Simulated ANOVA Result

	power	effect_size
anova_color	5.15	0.0223934
anova_age	5.09	0.0222605
anova_color:age	98.89	0.3076345

Table 5.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centralilty
color	5.00	0.0000	0.00	0.0000
age	5.00	0.0000	0.00	0.0000
color:age	98.98	0.3034	0.66	19.1667

```

      r = r,
      labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

```

```

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

5.2 Part 2

5.2.1 Two by two ANOVA, within-within design

We can simulate a 2x2 ANOVA, both factors manipulated within participants, with a specific sample size and effect size, to achieve a desired statistical power.

As Potvin & Schutz (2000) explain, analytic procedures for a two-factor repeated measures ANOVA do not seem to exist. The main problem is quantifying the error variance (the denominator when calculating lambda or Cohen's f). Simulation based approaches provide a solution.

We can reproduce the simulation coded by Ben Amsel

```

knitr::opts_chunk$set(echo = TRUE, warning = FALSE, message = FALSE)

# define the parameters
# true effects (in this case, a double dissociation)
mu = c(700, 670, 670, 700)
sigma = 150 # population standard deviation

```

```

rho = 0.75 # correlation between repeated measures
nsubs = 25 # how many subjects?
nsims = nsims # how many simulation replicates?

# create 2 factors representing the 2 independent variables
cond = data.frame(X1 = rep(factor(letters[1:2]), nsubs * 2),
                  X2 = rep(factor(letters[1:2]), nsubs, each = 2))

# create a subjects factor
subject = factor(sort(rep(1:nsubs, 4)))

# combine above into the design matrix
dm = data.frame(subject, cond)

```

Build Sigma: the population variance-covariance matrix

```

# create k x k matrix populated with sigma
sigma.mat <- rep(sigma, 4)
S <-
  matrix(sigma.mat,
        ncol = length(sigma.mat),
        nrow = length(sigma.mat))

# compute covariance between measures
Sigma <- t(S) * S * rho

# put the variances on the diagonal
diag(Sigma) <- sigma^2

```

Run the simulation

```

# stack 'nsims' individual data frames into one large data frame
df = dm[rep(seq_len(nrow(dm)), nsims), ]

# add an index column to track the simulation run
df$simID = sort(rep(seq_len(nsims), nrow(dm)))

# sample the observed data from a multivariate normal distribution
# using MASS::mvrnorm with the parameters mu and Sigma created earlier
# and bind to the existing df

make.y = expression(as.vector(t(mvrnorm(nsubs, mu, Sigma))))
df$y = as.vector(replicate(nsims, eval(make.y)))

# use do(), the general purpose complement to the specialized data
# manipulation functions available in dplyr, to run the ANOVA on
# each section of the grouped data frame created by group_by

mods <- df %>%
  group_by(simID) %>%
  do(model = aov(y ~ X1 * X2 + Error(subject / (X1 * X2)),

```



```

qr = FALSE, data = .))

# extract p-values for each effect and store in a data frame
p = data.frame(
  mods %>% do(as.data.frame(tidy(. $model[[3]])$p.value[1])),
  mods %>% do(as.data.frame(tidy(. $model[[4]])$p.value[1])),
  mods %>% do(as.data.frame(tidy(. $model[[5]])$p.value[1]))
colnames(p) = c('X1', 'X2', 'Interaction')

```

The empirical power is easy to compute, it's just the proportion of simulation runs where $p < .05$.

```

power.res = apply(as.matrix(p), 2,
  function(x) round(mean(ifelse(x < .05, 1, 0) * 100), 2))
power.res

```

```

##           X1           X2 Interaction
##          5.22          4.78          48.09

```

Visualize the distributions of p-values

```

# plot the known effects

means = data.frame(cond[1:4,], mu, SE = sigma / sqrt(nsubs))
plt1 = ggplot(means, aes(y = mu, x = X1, fill = X2)) +
  geom_bar(position = position_dodge(), stat = "identity") +
  geom_errorbar(
    aes(ymin = mu - SE, ymax = mu + SE),
    position = position_dodge(width = 0.9),
    size = .6,
    width = .3
  ) +
  coord_cartesian(ylim = c((.7 * min(mu)), 1.2 * max(mu))) +
  theme_bw()

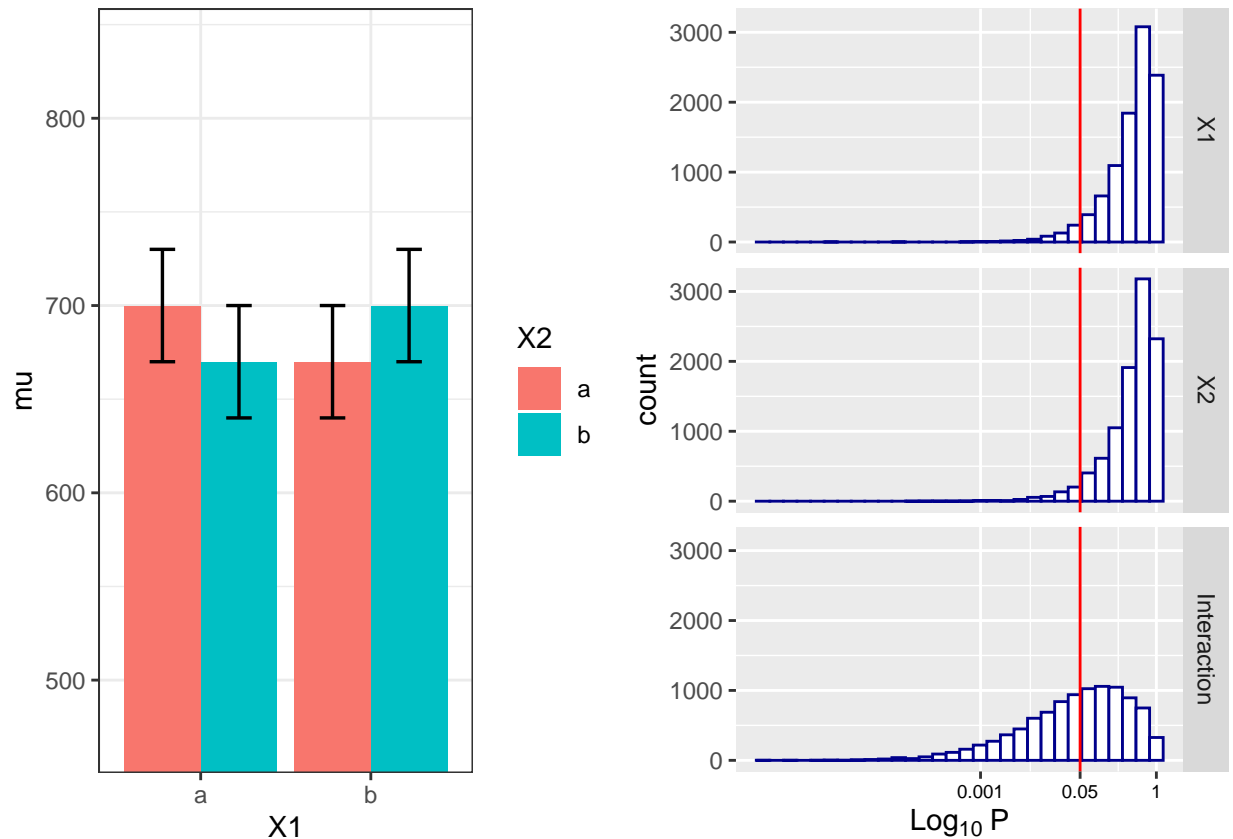
# melt the data into a ggplot friendly 'long' format

plotData <- melt(p, value.name = 'p')

# plot each of the p-value distributions on a log scale
options(scipen = 999) # 'turn off' scientific notation
plt2 = ggplot(plotData, aes(x = p)) +
  scale_x_log10(breaks = c(1, 0.05, 0.001),
    labels = c(1, 0.05, 0.001)) +
  geom_histogram(colour = "darkblue", fill = "white") +
  geom_vline(xintercept = 0.05, colour = 'red') +
  facet_grid(variable ~ .) +
  labs(x = expression(Log[10] ~ P)) +
  theme(axis.text.x = element_text(color = 'black', size = 7))

# arrange plots side by side and print
grid.arrange(plt1, plt2, nrow = 1)

```



We can reproduce this simulation:

```
mu = c(700, 670, 670, 700) # true effects (in this case, a double dissociation)
sigma = 150 # population standard deviation
n <- 25
sd <- 150
r <- 0.75
string = "2w*2w"
alpha_level <- 0.05
labelnames = c("age", "old", "young", "color", "blue", "red")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

Table 5.3: Simulated ANOVA Result

	power	effect_size
anova_age	5.14	0.0401014
anova_color	4.79	0.0393124
anova_age:color	48.34	0.1638188

Table 5.4: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non centrality
age	5.0	0.0000	0.0000	0
color	5.0	0.0000	0.0000	0
age:color	48.4	0.1429	0.4082	4

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

The simulations yield closely matching results.

5.2.2 Examine variation of means and correlation

```
# define the parameters
mu = c(700, 670, 690, 750) # true effects (in this case, a double dissociation)
sigma = 150 # population standard deviation
rho = 0.4 # correlation between repeated measures
nsubs = 25 # how many subjects?
nsims = nsims # how many simulation replicates?

# create 2 factors representing the 2 independent variables
cond = data.frame(X1 = rep(factor(letters[1:2]), nsubs * 2),
                  X2 = rep(factor(letters[1:2]), nsubs, each = 2))

# create a subjects factor
subject = factor(sort(rep(1:nsubs, 4)))

# combine above into the design matrix
dm = data.frame(subject, cond)
```

Build Sigma: the population variance-covariance matrix

```
# create k x k matrix populated with sigma
sigma.mat <- rep(sigma, 4)
S <-
matrix(sigma.mat,
       ncol = length(sigma.mat),
       nrow = length(sigma.mat))
```

```
# compute covariance between measures
Sigma <- t(S) * S * rho

# put the variances on the diagonal
diag(Sigma) <- sigma ^ 2
```

Run the simulation

```
# stack 'nsims' individual data frames into one large data frame
df = dm[rep(seq_len(nrow(dm)), nsims), ]

# add an index column to track the simulation run
df$simID = sort(rep(seq_len(nsims), nrow(dm)))

# sample the observed data from a multivariate normal distribution
# using MASS::mvrnorm with the parameters mu and Sigma created earlier
# and bind to the existing df

make.y = expression(as.vector(t(mvrnorm(nsubs, mu, Sigma))))
df$y = as.vector(replicate(nsims, eval(make.y)))

# use do(), the general purpose complement to the specialized data
# manipulation functions available in dplyr, to run the ANOVA on
# each section of the grouped data frame created by group_by

mods <- df %>%
  group_by(simID) %>%
  do(model = aov(y ~ X1 * X2 + Error(subject / (X1 * X2)),
    qr = FALSE, data = .))

# extract p-values for each effect and store in a data frame
p = data.frame(mods %>% do(as.data.frame(tidy(. $model[[3]])$p.value[1])),
  mods %>% do(as.data.frame(tidy(. $model[[4]])$p.value[1])),
  mods %>% do(as.data.frame(tidy(. $model[[5]])$p.value[1])))
colnames(p) = c('X1', 'X2', 'Interaction')
```

The empirical power is easy to compute, it's just the proportion of simulation runs where $p < .05$.

```
power.res = apply(as.matrix(p), 2,
  function(x) round(mean(ifelse(x < .05, 1, 0) * 100), 2))
power.res
```

```
##          X1          X2 Interaction
##        9.10        30.43         45.55
```

Visualize the distributions of p-values

```
means = data.frame(cond[1:4,], mu, SE = sigma / sqrt(nsubs))
plt1 = ggplot(means, aes(y = mu, x = X1, fill = X2)) +
  geom_bar(position = position_dodge(), stat = "identity") +
```

```

geom_errorbar(
  aes(ymin = mu - SE, ymax = mu + SE),
  position = position_dodge(width = 0.9),
  size = .6,
  width = .3
) +
coord_cartesian(ylim = c((.7 * min(mu)), 1.2 * max(mu))) +
theme_bw()

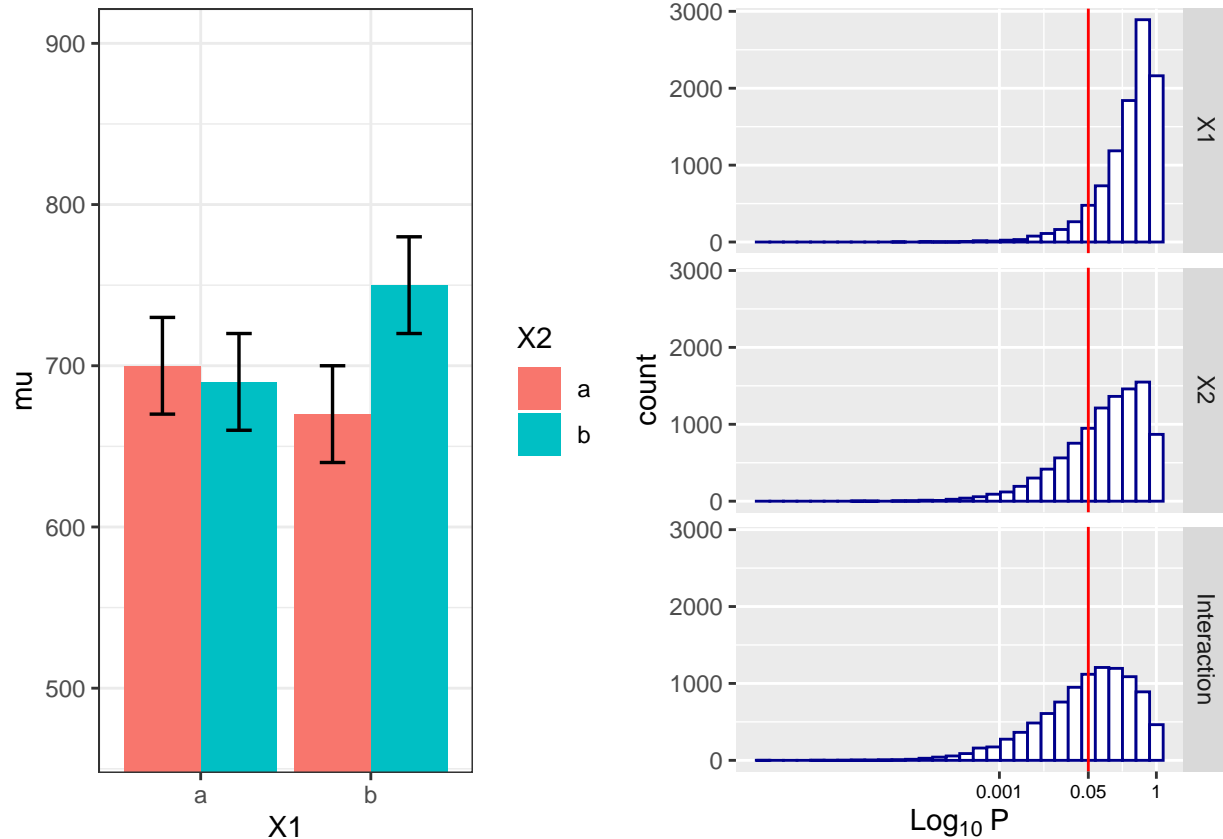
# melt the data into a ggplot friendly 'long' format

plotData <- melt(p, value.name = 'p')

# plot each of the p-value distributions on a log scale
options(scipen = 999) # 'turn off' scientific notation
plt2 = ggplot(plotData, aes(x = p)) +
  scale_x_log10(breaks = c(1, 0.05, 0.001),
  labels = c(1, 0.05, 0.001)) +
  geom_histogram(colour = "darkblue", fill = "white") +
  geom_vline(xintercept = 0.05, colour = 'red') +
  facet_grid(variable ~ .) +
  labs(x = expression(Log[10] ~ P)) +
  theme(axis.text.x = element_text(color = 'black', size = 7))

# arrange plots side by side and print
grid.arrange(plt1, plt2, nrow = 1)

```



We can reproduce this simulation:

```
mu = c(700, 670, 690, 750) # true effects (in this case, a double dissociation)
sigma = 150 # population standard deviation
n <- 25
sd <- 150
r <- 0.4
string = "2w*2w"
alpha_level <- 0.05
labelnames = c("age", "old", "young", "color", "blue", "red")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = 0.05,
                                 nsims = nsims)
```

```
## Power and Effect sizes for ANOVA tests
##           power effect_size
## anova_age      31.48    0.11655
## anova_color     9.99    0.05498
## anova_age:color 45.81    0.15789
```

```
##
## Power and Effect sizes for contrasts
##
##           power effect_size
## p_age_old_color_blue_age_old_color_red    14.34    -0.1882
## p_age_old_color_blue_age_young_color_blue    5.74    -0.0615
## p_age_old_color_blue_age_young_color_red    31.24     0.3157
## p_age_old_color_red_age_young_color_blue     9.38     0.1276
## p_age_old_color_red_age_young_color_red    64.49     0.5050
## p_age_young_color_blue_age_young_color_red  41.90     0.3768
##
## Within-Subject Factors Included: Check MANOVA Results
```

```
exact_result <- ANOVA_exact(design_result, alpha_level = alpha_level)
```

```
## Power and Effect sizes for ANOVA tests
##           power partial_eta_squared cohen_f non centrality
## age       30.40           0.0864  0.3074         2.2685
## color      9.51           0.0171  0.1318         0.4167
## age:color  45.98           0.1351  0.3953         3.7500
##
## Power and Effect sizes for contrasts
##
##           power effect_size
## p_age_old_color_blue_age_old_color_red    14.16    -0.1826
## p_age_old_color_blue_age_young_color_blue    5.98    -0.0609
## p_age_old_color_blue_age_young_color_red    30.91     0.3043
## p_age_old_color_red_age_young_color_blue     9.00     0.1217
## p_age_old_color_red_age_young_color_red    64.66     0.4869
## p_age_young_color_blue_age_young_color_red  41.80     0.3651
```

5.3 Part 3

Two by two ANOVA, within design

Potvin & Schutz (2000) simulate a wide range of repeated measure designs. They give an example of a 3x3 design, with the following correlation matrix:

Example

$\rho_A = 0.4$			$\rho_B = 0.8$			$\rho_{AB} = 0.4$		
A ₁			A ₂			A ₃		
B ₁	B ₂	B ₃	B ₁	B ₂	B ₃	B ₁	B ₂	B ₃
A ₁	B ₁	1.0	0.8	0.8	0.4	0.4	0.4	0.4
	B ₂		1.0	0.8	0.4	0.4	0.4	0.4
	B ₃			1.0	0.4	0.4	0.4	0.4
A ₂	B ₁			1.0	0.8	0.8	0.4	0.4
	B ₂				1.0	0.8	0.4	0.4
	B ₃					1.0	0.4	0.4
A ₃	B ₁						1.0	0.8
	B ₂							1.0
	B ₃							

Figure 1. Representation of a correlation matrix for a 3 (A) × 3 (B) RM ANOVA: General form and numeric example. ρ_A and ρ_B represent the average correlation among the A and B (pooled) trials, respectively, and ρ_{AB} represents the average correlation among the AB coefficients having dissimilar levels.

Variances were set to 1 (so all covariance matrices in their simulations were identical). In this specific example, the white fields are related to the correlation for the A main effect (these cells have the same level for B, but different levels of A). The grey cells are related to the main effect of B (the cells have the same level of A, but different levels of B). Finally, the black cells are related to the AxB interaction (they have different levels of A and B). The diagonal (all 1) relate to cells with the same levels of A and B.

Potvin & Schulz (2000) examine power for 2x2 within ANOVA designs and develop approximations of the error variance. For a design with 2 within factors (A and B) these are:

$$\text{For the main effect of A: } \sigma_e^2 = \sigma^2(1 - \bar{\rho}_A) + \sigma^2(q - 1)(\bar{\rho}_B - \bar{\rho}_{AB})$$

$$\text{For the main effect of B: } \sigma_e^2 = \sigma^2(1 - \bar{\rho}_B) + \sigma^2(p - 1)(\bar{\rho}_A - \bar{\rho}_{AB})$$

$$\text{For the interaction between A and B: } \sigma_e^2 = \sigma^2(1 - \rho_{\max}) - \sigma^2(\bar{\rho}_{\min} - \bar{\rho}_{AB})$$

5.4 Simple example: 2x2 within design

It is difficult to just come up with a positive definite covariance matrix. The best way to achieve this is to get the correlations from a pilot study. Indeed, it should be rather difficult to know which correlations to fill in without some pilot data.

We try to get the formulas in Potvin and Schutz (2000) working. **Below, I manage for the main effects, but not for the interaction.**

Table 5.5: Simulated ANOVA Result

	power	effect_size
anova_A	25.37	0.1223527
anova_B	81.45	0.3305097
anova_A:B	16.02	0.0915181

Table 5.6: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
A	24.71	0.0865	0.3078	1.8
B	81.21	0.3214	0.6882	9.0
A:B	15.81	0.0500	0.2294	1.0

```

mu = c(2,1,4,2)
n <- 20
sd <- 5
r <- c(
  0.8, 0.4, 0.4,
    0.4, 0.4,
      0.8
)
string = "2w*2w"
alpha_level <- 0.05
labelnames = c("A", "a1", "a2", "B", "b1", "b2")

design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

We can try to use the formula in Potvin & Schutz (2000).

```

k <- 1 #one group (because all factors are within)

```

```

rho_A <- 0.5 #mean r for factor A
rho_B <- 0.8 #mean r for factor B
rho_AB <- 0.4 #mean r for factor AB
alpha <- 0.05
sigma <- sd
m_A <- 2 #levels factor A

variance_e_A <- sigma^2 * (1 - rho_A) + sigma^2 * (m_A - 1) * (rho_B - rho_AB)
#Variance A
variance_e_A

```

```
## [1] 22.5
```

```

m_B <- 2 #levels factor B

variance_e_B <- sigma^2 * (1 - rho_B) + sigma^2 * (m_B - 1) * (rho_A - rho_AB)
#Variance B
variance_e_B

```

```
## [1] 7.5
```

```

variance_e_AB <-
  (sigma ^ 2 * (1 - max(rho_A, rho_B)) -
   sigma ^ 2 * (min(rho_A, rho_B) - rho_AB))
#Variance AB
variance_e_AB

```

```
## [1] 2.5
```

```

#Create a mean matrix
mean_mat <- t(matrix(mu, nrow = m_B, ncol = m_A))
mean_mat

```

```

##      [,1] [,2]
## [1,]    2    1
## [2,]    4    2

```

```

# Potvin & Schutz, 2000, formula 2, p. 348
# For main effect A
lambda_A <-
  n * m_A * sum((rowMeans(mean_mat) -
                  mean(rowMeans(mean_mat))) ^ 2) / variance_e_A
lambda_A

```

```
## [1] 2
```

```

#calculate degrees of freedom 1 - ignoring the * e sphericity correction
df1 <- (m_A - 1)

df2 <- (n - k) * (m_A - 1) #calculate degrees of freedom 2

F_critical <- qf(alpha, # critical F-value
                df1,
                df2,
                lower.tail = FALSE)

pow_A <- pf(qf(alpha, #power
                df1,
                df2,
                lower.tail = FALSE),
            df1,
            df2,
            lambda_A,
            lower.tail = FALSE)

lambda_B <-
  n * m_B * sum((colMeans(mean_mat) -
                  mean(colMeans(mean_mat))) ^ 2) / variance_e_B
lambda_B

```

```
## [1] 6
```

```

df1 <- (m_B - 1) #calculate degrees of freedom 1

df2 <- (n - k) * (m_B - 1) #calculate degrees of freedom 2

F_critical <- qf(alpha, # critical F-value
                df1,
                df2,
                lower.tail = FALSE)

pow_B <- pf(qf(alpha, #power
                df1,
                df2,
                lower.tail = FALSE),
            df1,
            df2,
            lambda_B,
            lower.tail = FALSE)

pow_A

```

```
## [1] 0.2691752
```

```
pow_B
```

```
## [1] 0.6422587
```

We see the 26.9 and 64.2 correspond to the results of the simulation quite closely.

```
#This (or the variance calculation above) does not work.
lambda_AB <- n * sum((
mean_mat - rowMeans(mean_mat) - colMeans(mean_mat) + mean(mean_mat)
) ^ 2) / variance_e_AB
lambda_AB
```

```
## [1] 38
```

```
df1 <- (m_A - 1) * (m_B - 1) #calculate degrees of freedom 1
df2 <-
(n - k) * (m_A - 1) * (m_B - 1) #calculate degrees of freedom 2
F_critical <- qf(alpha, # critical F-value
df1,
df2,
lower.tail = FALSE)

pow <- pf(qf(alpha, #power
df1,
df2,
lower.tail = FALSE),
df1,
df2,
lambda_AB,
lower.tail = FALSE)

pow
```

```
## [1] 0.9999458
```

Maybe the simulation is not correct for the interaction, or the formula is not correctly programmed.

Chapter 6

Analytic Power for Three-way Interactions

There are almost no software solutions that allow researchers to perform power analysis for more complex designs. Through simulation, it is relatively straightforward to examine the power for designs with multiple factors with many levels.

Let's start with a 2x2x2 between subjects design. We collect 50 participants in each between participant condition (so 400 participants in total - 50x2x2x2).

```
# With 2x2x2 designs, the names for paired comparisons can become very long.
# So here I abbreviate terms: Size, Color, and Cognitive Load, have values:
# b = big, s = small, g = green, r = red, pres = present, abs = absent.
labelnames <- c("Size", "b", "s", "Color", "g", "r",
               "Load", "pres", "abs") #
design_result <- ANOVA_design(design = "2b*2b*2b", #describe the design
                             n = 50, #sample size per group
                             mu = c(2, 2, 6, 1, 6, 6, 1, 8), #pattern of means
                             sd = 10, #standard deviation
                             labelnames = labelnames) #names of labels

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

\begin{table}[t]

\caption{(#tab:start_threewayinteraction)Simulated ANOVA Result}

	power	effect_size
anova_Size	70.35	0.0180484
anova_Color	5.40	0.0025813
anova_Load	8.19	0.0032585
anova_Size:Color	31.87	0.0082036
anova_Size:Load	84.67	0.0248491
anova_Color:Load	7.79	0.0031371
anova_Size:Color:Load	84.89	0.0247389

\end{table}

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

\begin{table}[t]

\caption{(#tab:start_threewayinteraction)Exact ANOVA Result}

	power	partial_eta_squared	cohen_f	non centrality
Size	70.33	0.0157	0.1263	6.25
Color	5.00	0.0000	0.0000	0.00
Load	7.90	0.0006	0.0253	0.25
Size:Color	32.17	0.0057	0.0758	2.25
Size:Load	84.91	0.0224	0.1515	9.00
Color:Load	7.90	0.0006	0.0253	0.25
Size:Color:Load	84.91	0.0224	0.1515	9.00

\end{table}

```
#Analytical power calculation
power_analytic <- power_threeway_between(design_result)
power_analytic$power_A
```

```
## [1] 70.33333
```

```
power_analytic$power_B
```

```
## [1] 5
```

```
power_analytic$power_C
```

```
## [1] 7.895539
```

```
power_analytic$power_AB
```

```
## [1] 32.17471
```

```
power_analytic$power_AC
```

```
## [1] 84.91491
```

```
power_analytic$power_BC
```

```
## [1] 7.895539
```

```
power_analytic$power_ABC
```

```
## [1] 84.91491
```

```
power_analytic$eta_p_2_A
```

```
## [1] 0.01538462
```

```
power_analytic$eta_p_2_B
```

```
## [1] 0
```

```
power_analytic$eta_p_2_C
```

```
## [1] 0.0006246096
```

```
power_analytic$eta_p_2_AB
```

```
## [1] 0.005593536
```

```
power_analytic$eta_p_2_AC
```

```
## [1] 0.02200489
```

```
power_analytic$eta_p_2_BC
```

```
## [1] 0.0006246096
```

```
power_analytic$eta_p_2_ABC
```

```
## [1] 0.02200489
```

We can also confirm the power analysis in GPower. GPower allows you to compute the power for a three-way interaction - if you know the Cohen's f value to enter. Cohen's f is calculated based on the means for the interaction, the sum of squares of the effect, and the sum of squares of the errors. This is quite a challenge by hand, but we can simulate the results, or use the analytical solution we programmed to get Cohen's f for the pattern of means that we specified.

```
# The power for the AC interaction (Size x Load) is 0.873535.  
power_analytic$power_AC
```

```
## [1] 84.91491
```

```
# We can enter the Cohen's f for this interaction.  
power_analytic$Cohen_f_AC
```

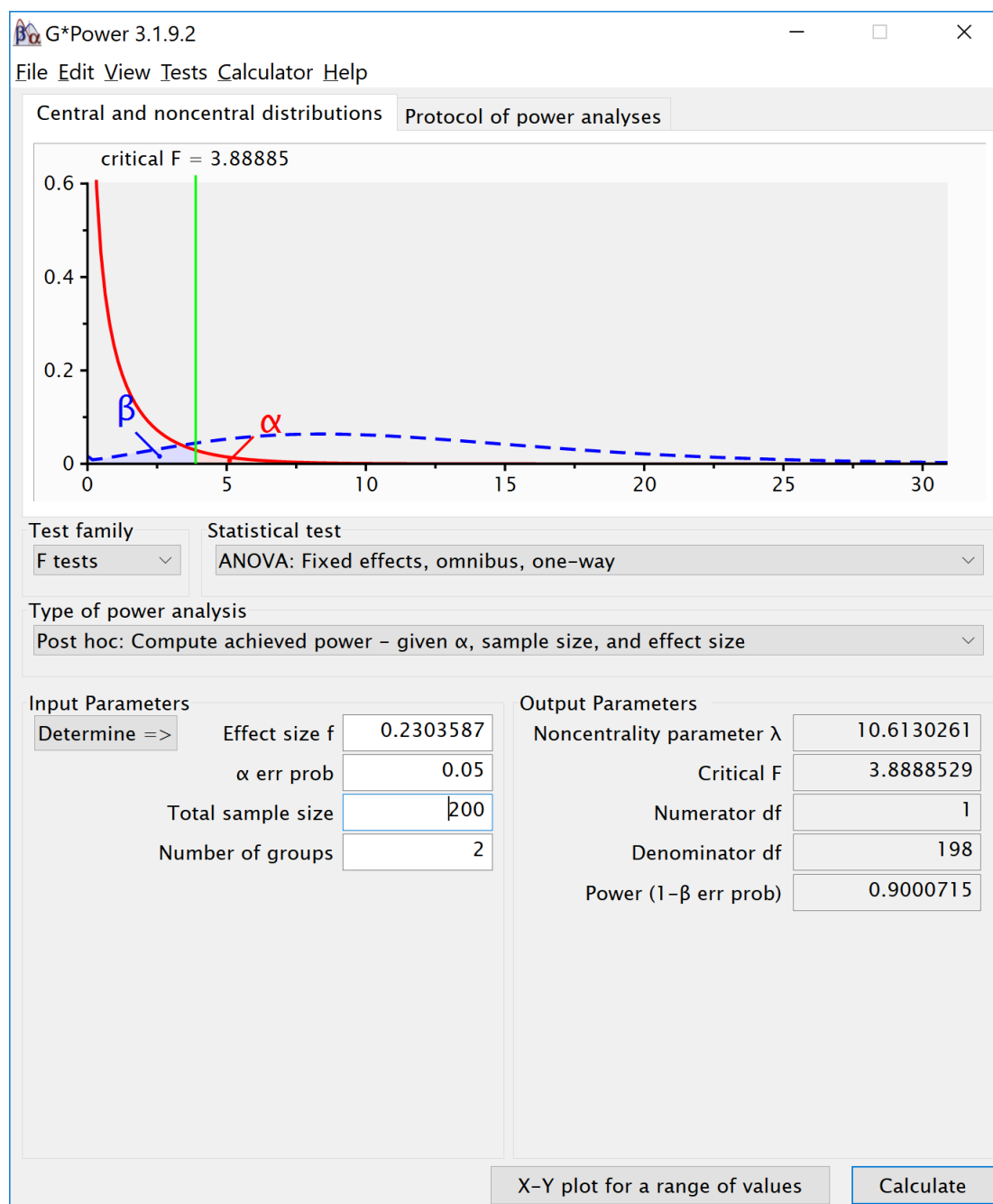
```
## [1] 0.15
```

```
# We can double check the calculated lambda  
power_analytic$lambda_AC
```

```
## [1] 9
```

```
# We can double check the critical F value  
power_analytic$F_critical_AC
```

```
## [1] 3.864929
```

A Three-Way ANOVA builds on the same principles as a One_Way ANOVA. We look at whether the differences between groups are large, compared to the standard deviation. For the main effects we simply have 2 groups of 200 participants, and 2 means. If the population standard deviations are identical across groups, this is not in any way different from a One-Way ANOVA. Indeed, we can show this by simulating a One-Way ANOVA, where instead of 8 conditions, we have two conditions, and we average over the 4 groups of the other two factors. For example, for the main effect of size above can be computed analytically. There might be a small difference in the degrees of freedom of the two tests, or it is just random variation (And it will disappear when repeating the simulation 1000.000 times instead of 100.000.

```
string <- "2b"
n <- 200
mu <- c(mean(c(2, 2, 6, 1)), mean(c(6, 6, 1, 8)))
```

Table 6.1: Simulated ANOVA Result

	power	effect_size
anova_Size	70.3	0.0177696

Table 6.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Size	70.33	0.0155	0.1253	6.25

```
sd <- 10
labelnames <- c("Size", "big", "small")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```
exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

```
# Power based on analytical solution
power_oneway_between(design_result)$power #using default alpha level of .05
```

```
## [1] 70.33333
```

Similarly, we can create a 2 factor design where we average over the third factor, and recreate the power analysis for the Two-Way interaction. For example, we can group over the Cognitive Load condition, and look at the Size by Color Interaction:

```
string <- "2b*2b"
n <- 100
mu <- c(mean(c(1, 1)), mean(c(6, 1)), mean(c(6, 6)), mean(c(1, 6)))
sd <- 10
labelnames <- c("Size", "big", "small", "Color", "green", "red")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
```

Table 6.3: Simulated ANOVA Result

	power	effect_size
anova_Size	70.37	0.0179101
anova_Color	5.14	0.0025735
anova_Size:Color	70.25	0.0180505

Table 6.4: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non centrality
Size	70.33	0.0155	0.1256	6.25
Color	5.00	0.0000	0.0000	0.00
Size:Color	70.33	0.0155	0.1256	6.25

```

      labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                nsims = nsims,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

# Power based on analytical solution
power_res <- power_twoway_between(design_result) #using default alpha level of .05
power_res$power_A

## [1] 70.33228

power_res$power_B

## [1] 5

power_res$power_AB

## [1] 70.33228

string <- "2b*2b*2b"
n <- 50
mu <- c(5, 3, 2, 6, 1, 4, 3, 1)
sd <- 10

```

Table 6.5: Simulated ANOVA Result

	power	effect_size
anova_Size	41.95	0.0102725
anova_Color	5.37	0.0026697
anova_CognitiveLoad	11.72	0.0039672
anova_Size:Color	5.66	0.0026893
anova_Size:CognitiveLoad	5.80	0.0026676
anova_Color:CognitiveLoad	6.06	0.0027339
anova_Size:Color:CognitiveLoad	77.22	0.0210871

Table 6.6: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Size	41.53	0.0078	0.0884	3.0625
Color	5.72	0.0002	0.0126	0.0625
CognitiveLoad	11.62	0.0014	0.0379	0.5625
Size:Color	5.72	0.0002	0.0126	0.0625
Size:CognitiveLoad	5.72	0.0002	0.0126	0.0625
Color:CognitiveLoad	5.72	0.0002	0.0126	0.0625
Size:Color:CognitiveLoad	78.33	0.0189	0.1389	7.5625

```

r <- 0.0
labelnames <- c("Size", "big", "small", "Color", "green", "red",
                "CognitiveLoad", "present", "absent") #
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")

```

```

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

```

#Analytical power calculation
power_analytic <- power_threeway_between(design_result)
power_analytic$power_A

```

```
## [1] 41.5306
```

```
power_analytic$power_B
```

```
## [1] 5.715533
```

```
power_analytic$power_C
```

```
## [1] 11.61827
```

```
power_analytic$power_AB
```

```
## [1] 5.715533
```

```
power_analytic$power_AC
```

```
## [1] 5.715533
```

```
power_analytic$power_BC
```

```
## [1] 5.715533
```

```
power_analytic$power_ABC
```

```
## [1] 78.33036
```

```
power_analytic$eta_p_2_A
```

```
## [1] 0.007598077
```

```
power_analytic$eta_p_2_B
```

```
## [1] 0.0001562256
```

```
power_analytic$eta_p_2_C
```

```
## [1] 0.001404275
```

```
power_analytic$eta_p_2_AB
```

```
## [1] 0.0001562256
```

```
power_analytic$eta_p_2_AC
```

```
## [1] 0.0001562256
```

```
power_analytic$eta_p_2_BC
```

```
## [1] 0.0001562256
```

```
power_analytic$eta_p_2_ABC
```

```
## [1] 0.01855544
```

The power for interactions depends on Cohen's f , the alpha level, the sample size, and the degrees of freedom.

```
# With 2x2x2 designs, the names for paired comparisons can become very long.
# So here the sample size abbreviate terms
# Size, Color, and Cognitive Load, have values:
# b = big, s = small, g = green, r = red, pres = present, abs = absent.
labelnames <- c("Size", "b", "s", "x", "Color", "g", "r",
               "Load", "pres", "abs") #
design_result <- ANOVA_design(design = "3b*2b*2b", #describe the design
                             n = 15, #sample size per group
                             mu = c(20, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 20),

                             sd = 20, #
                             labelnames = labelnames)

# Power based on exact simulations
exact_result <- ANOVA_exact(design_result)
```

```
## Power and Effect sizes for ANOVA tests
```

```
##               power partial_eta_squared cohen_f non centrality
## Size           26.93                0.0147  0.1220           2.5
## Color           5.00                0.0000  0.0000           0.0
## Load           5.00                0.0000  0.0000           0.0
## Size:Color      67.93                0.0427  0.2113           7.5
## Size:Load       67.93                0.0427  0.2113           7.5
## Color:Load      60.39                0.0289  0.1725           5.0
## Size:Color:Load 26.93                0.0147  0.1220           2.5
##
```

```
## Power and Effect sizes for contrasts
```

```
##               power effect_size
## p_Size_b_Color_g_Load_pres_Size_b_Color_g_Load_abs 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_b_Color_r_Load_pres 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_b_Color_r_Load_abs 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_s_Color_g_Load_pres 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_s_Color_g_Load_abs 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_s_Color_r_Load_pres 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_s_Color_r_Load_abs 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_x_Color_g_Load_pres 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_x_Color_g_Load_abs 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_x_Color_r_Load_pres 75.29      -1
## p_Size_b_Color_g_Load_pres_Size_x_Color_r_Load_abs 5.00        0
## p_Size_b_Color_g_Load_abs_Size_b_Color_r_Load_pres 5.00        0
```

[illegible]

```
#Analytical power calculation
power_analytic <- power_threeway_between(design_result)
power_analytic$power_A
```

```
## [1] 5
```

```
power_analytic$power_B
```

```
## [1] 5
```

```
power_analytic$power_C
```

```
## [1] 48.6496
```

```
power_analytic$power_AB
```

```
## [1] 34.7961
```

```
power_analytic$power_AC
```

```
## [1] 67.97466
```

```
power_analytic$power_BC
```

```
## [1] 91.55713
```

```
power_analytic$power_ABC
```

```
## [1] NaN
```

```
power_analytic$eta_p_2_A
```

```
## [1] 0
```

```
power_analytic$Cohen_f_A
```

```
## [1] 0
```

We see that a pattern of means of 0, 0, 0, 0, 0, 0, 0, 20 for a 2x2x2 interaction equals a Cohen's f of 0.25.

```
labelnames <- c("Size", "b", "s", "Color", "g", "r")
design_result <- ANOVA_design(design = "2b*2b", #describe the design
                             n = 10, #sample size per group
                             mu = c(0, 0, 0, 10), #pattern of means
                             sd = 10, #standard deviation
                             labelnames = labelnames) #names of labels

# Power based on exact simulations
exact_result <- ANOVA_exact(design_result)
```



```
## Power and Effect sizes for ANOVA tests
##           power partial_eta_squared cohen_f non centrality
## Size      33.71                0.0649  0.2635           2.5
## Color      33.71                0.0649  0.2635           2.5
## Size:Color 33.71                0.0649  0.2635           2.5
##
## Power and Effect sizes for contrasts
##                               power effect_size
## p_Size_b_Color_g_Size_b_Color_r    5.0          0
## p_Size_b_Color_g_Size_s_Color_g    5.0          0
## p_Size_b_Color_g_Size_s_Color_r   56.2          1
## p_Size_b_Color_r_Size_s_Color_g    5.0          0
## p_Size_b_Color_r_Size_s_Color_r   56.2          1
## p_Size_s_Color_g_Size_s_Color_r   56.2          1
```

```
#Analytical power calculation
```

```
power_analytic <- power_twoway_between(design_result)
power_analytic$power_A
```

```
## [1] 33.71329
```

```
power_analytic$eta_p_2_A
```

```
## [1] 0.05882353
```

```
power_analytic$Cohen_f_A
```

```
## [1] 0.25
```

Cohen's f is twice as large for a 2x2 design with the same mean value in one of four cells. In a 2 factor between design.

```
labelnames <- c("Size", "b", "s")
design_result <- ANOVA_design(design = "2b", #describe the design
                             n = 10, #sample size per group
                             mu = c(0, 5), #pattern of means
                             sd = 10, #standard deviation
                             labelnames = labelnames) #names of labels

# Power based on exact simulations
exact_result <- ANOVA_exact(design_result)
```

```
## Power and Effect sizes for ANOVA tests
##           power partial_eta_squared cohen_f non centrality
## Size 18.51                0.0649  0.2635           1.25
##
## Power and Effect sizes for contrasts
##                               power effect_size
## p_Size_b_Size_s 18.51                0.5
```

```
#Analytical power calculation  
power_analytic <- power_oneway_between(design_result)  
power_analytic$power
```

```
## [1] 18.50957
```

```
power_analytic$eta_p_2
```

```
## [1] 0.05882353
```

```
power_analytic$Cohen_f
```

```
## [1] 0.25
```

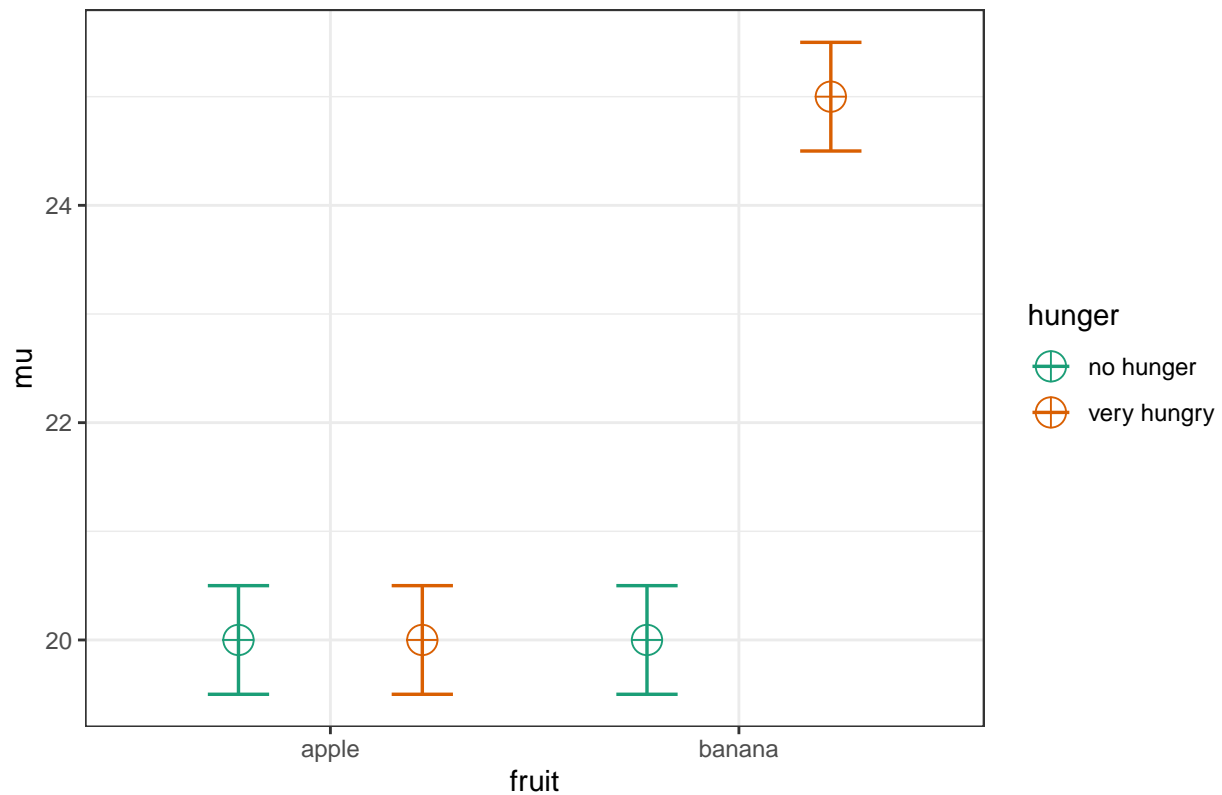
Chapter 7

Power for Interactions

In the 17th Data Colada blog post titled No-way Interactions Uri Simonsohn discusses how a moderated interaction (the effect is there in one condition, but disappears in another condition) requires at least twice as many subjects per cell as a study that simply aims to show the simple effect. For example, see the plot below. Assume the score on the vertical axis is desire for fruit, as a function of the fruit that is available (an apple or a banana) and how hungry people are (not, or very). We see there is a difference between the participants desire for a banana compared to an apple, but only for participants who are very hungry. The point that is made is that you need twice as many participants in each cell to have power for the interaction, as you need for the simple effect.

```
string <- "2b*2b"
n <- 20
# All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
mu <- c(20, 20, 20, 25)
sd <- 0.5
labelnames <- c("fruit", "apple", "banana",
                "hunger", "no hunger", "very hungry") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames,
                             plot = TRUE)
```

Means for each condition in the design



We can reproduce the simulations in the Data Colada blog post, using the original code.

```
#R-Code
#
#Written by Uri Simonsohn, March 2014
#
#
#In DataColada[17]
# I propose that 2x2 interaction studies need 2x the sample size
#http://datacolada.org/2014/03/10/17-no-way-interactions
#In a companion ,pdf I show the simple math behind it
#
#
#Simulations are often more persuasive than math, so here it goes.
#I run simulations that compute power for 2 and 4 cell design,
# the latter testing the interaction
#####
#Create function that computes power of Studies 1 and 2,
# where Study 1 has 2 cells and tests a simple effect
#and Study 2 has 4 cells and tests the interaction
colada17 = function(d1,d2,n1,n2,simt0t)
{
  #n1: sample size, per cell, study 1
  #n2: sample size, per cell, study 2
  #d1: simple effect M1-M2
  #d2: moderated effect M3-M4,
```

```

# full elimination of effect implies d2=0
#simtoto: how many simulations to run
#Here we will store results
  p1 = c()      #p-values for Study 1
  p2 = c()      #p-values for Study 2
for (i in 1:simtoto) {
  #draw data 4 samples
  y1 = rnorm(n = max(n1, n2), mean = d1)
  y2 = rnorm(n = max(n1, n2))
  y3 = rnorm(n = max(n1, n2), mean = d2)
  y4 = rnorm(n = max(n1, n2))

  #GET DATA READY FOR ANOVA
  y = c(y1, y2, y3, y4)      #the d.v.
  nrep = rep(n2, 4)
  A = rep(c(1, 1, 0, 0), times = nrep)
  B = rep(c(1, 0, 1, 0), times = nrep)

  #STUDY 1
  #Do a t-test on the first n1 observations
  p1.k = t.test(y1[1:n1], y2[1:n1], var.equal = TRUE)$p.value

  #STUDY 2
  #Do anova, keep p-value of the interaction
  p2.k = anova(lm(y ~ A * B))["A:B", "Pr(>F)"]

  #Store the results
  p1 = c(p1, p1.k)
  p2 = c(p2, p2.k)

}

#What share off comparisons are significant
#Simple test using estimate of variance from 2 cells only
power1 = sum(p1 <= .05) / simtoto
#Interaction
power2 = sum(p2 <= .05) / simtoto

cat("\nStudy 1 is powered to:",round(power1,2))
cat("\nStudy 2 is powered to:",round(power2,2))

}

#Same power for 2n regardless of n and d
colada17(simtoto = 2000, n1 = 20, n2 = 40, d1 = 1, d2 = 0)

##
## Study 1 is powered to: 0.86
## Study 2 is powered to: 0.88

colada17(simtoto = 2000, n1 = 50, n2 = 100, d1 = .3, d2 = 0)

```

```
##
## Study 1 is powered to: 0.32
## Study 2 is powered to: 0.36
```

```
colada17(simtot = 2000, n1 = 150, n2 = 300, d1 = .25, d2 = 0)
```

```
##
## Study 1 is powered to: 0.58
## Study 2 is powered to: 0.58
```

```
#Need 4n if effect is 70% attenuated
colada17(simtot = 2000, n1 = 25, n2 = 100, d1 = .5, d2 = .3 * .5)
```

```
##
## Study 1 is powered to: 0.39
## Study 2 is powered to: 0.42
```

```
colada17(simtot = 2000, n1 = 50, n2 = 200, d1 = .5, d2 = .3 * .5)
```

```
##
## Study 1 is powered to: 0.69
## Study 2 is powered to: 0.69
```

```
colada17(simtot = 2000, n1 = 22, n2 = 88, d1 = .41, d2 = .3 * .41)
```

```
##
## Study 1 is powered to: 0.27
## Study 2 is powered to: 0.26
```

```
#underpowered if run with the same n
colada17(simtot = nsims, n1 = 20, n2 = 20, d1 = 1, d2 = 0)
```

```
##
## Study 1 is powered to: 0.87
## Study 2 is powered to: 0.6
```

And we can reproduce the results using the ANOVA_exact function.

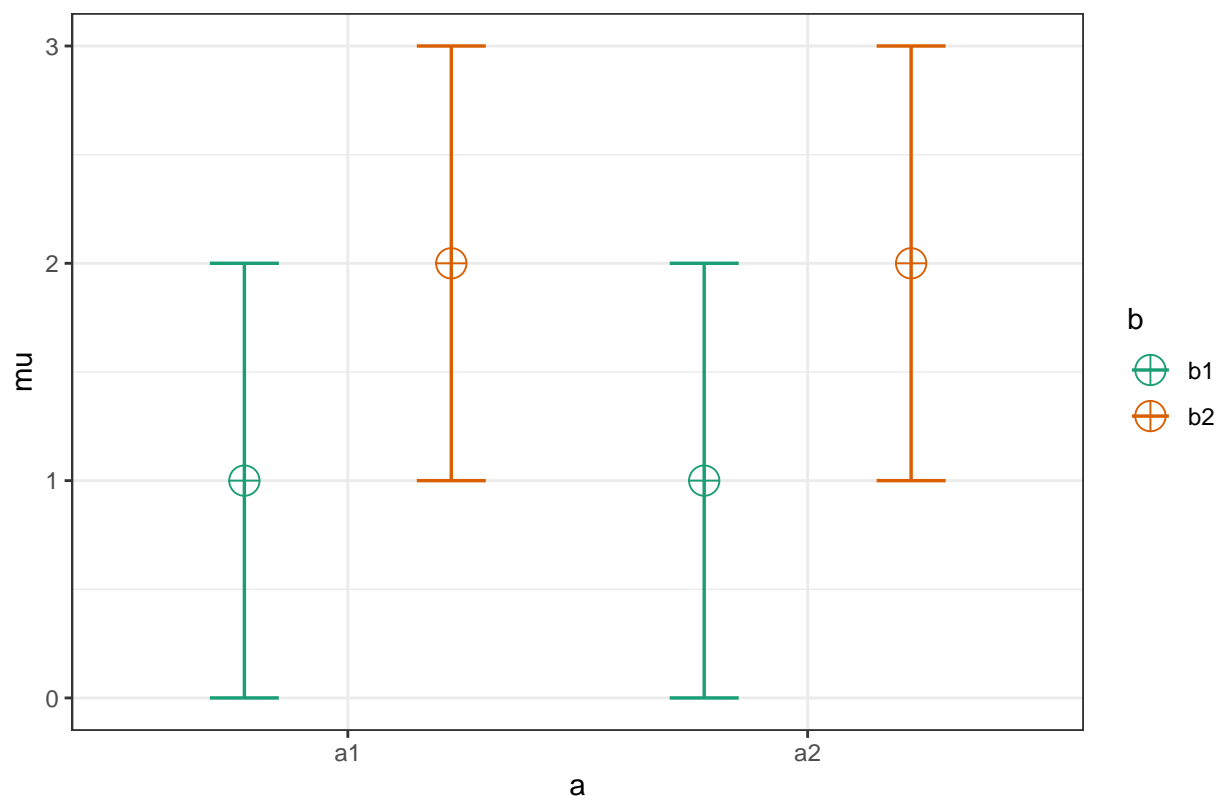
```
#Study 1
string <- "2b*2b"
n <- 10
# All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
mu <- c(1, 2, 1, 2)
sd <- 1

# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             plot = TRUE)
```

Table 7.1: ANOVA Results

	power	partial_eta_squared	cohen_f	non centrality
b	86.8	0.2174	0.527	10

Means for each condition in the design



```
alpha_level <- 0.05 #We set the alpha level at 0.05.
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)
```

```
knitr::kable(exact_result$main_results[2,],
              caption = "ANOVA Results")
```

```
#Study 2
```

```
string <- "2b*2b"
```

```
n <- 20
```

```
# All means are equal - so there is no real difference.
```

```
# Enter means in the order that matches the labels below.
```

```
mu <- c(1, 2, 1, 3)
```

```
sd <- 1
```

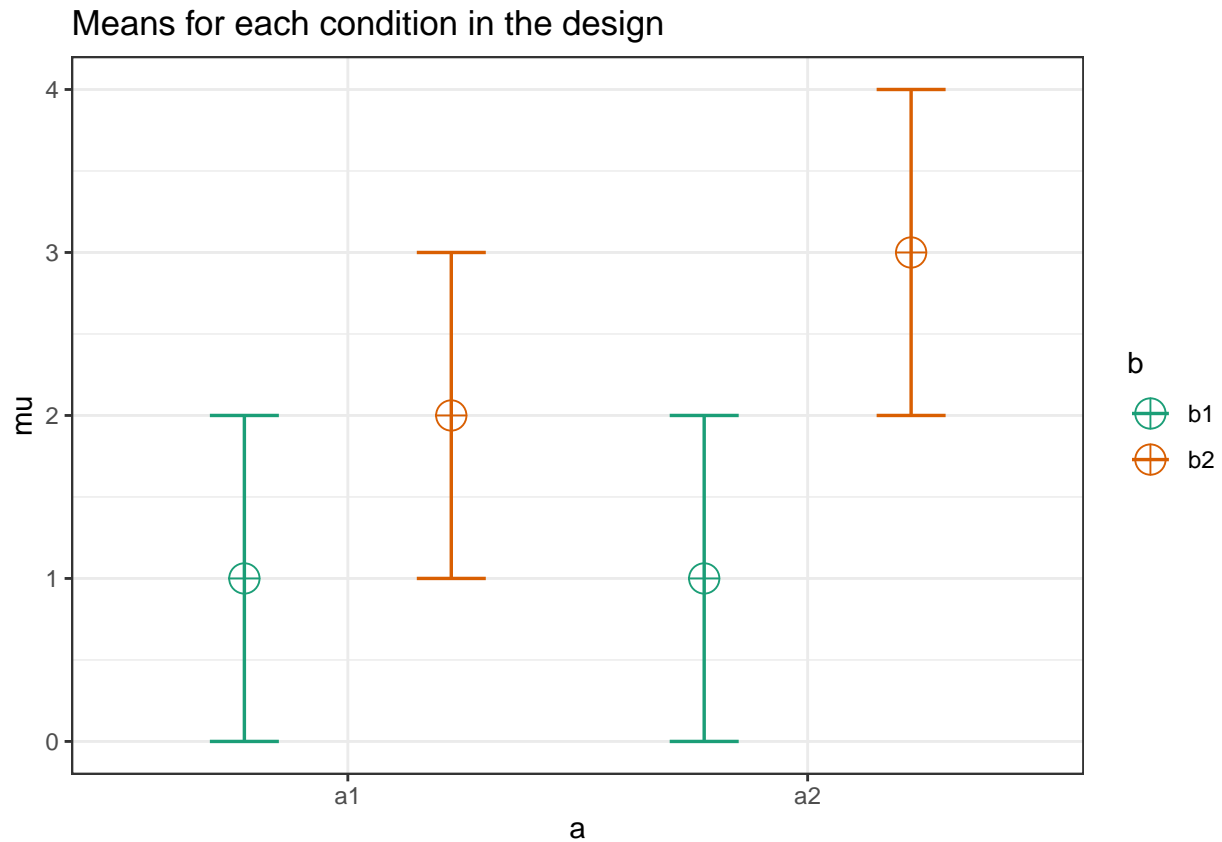
```
# the label names should be in the order of the means specified above.
```

```
design_result <- ANOVA_design(design = string,
                              n = n,
```

Table 7.2: ANOVA Results

	power	partial_eta_squared	cohen_f	non_centrality
a:b	59.79	0.0617	0.2565	5

```
mu = mu,
sd = sd,
plot = TRUE)
```



```
alpha_level <- 0.05 #We set the alpha level at 0.05.

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results[3,],
              caption = "ANOVA Results")
```

We see we get the same power for the `anova_fruit:hunger` interaction and for the simple effect `p_fruit_apple_hunger_very hungry_fruit_banana_hunger_very hungry` as the simulations by Uri Simonsohn in his blog post.

```
#Same power for 2n regardless of n and d
colada17(simtot = 10000, n1 = 20, n2 = 40, d1 = 1, d2 = 0)
```


Table 7.3: ANOVA Results

	power	partial_eta_squared	cohen_f	non_centralilty
hunger	33.33	0.0039	0.0627	2.3438

```
##
## Study 1 is powered to: 0.87
## Study 2 is powered to: 0.88
```

```
colada17(simtot = 10000, n1 = 50, n2 = 100, d1 = .3, d2 = 0)
```

```
##
## Study 1 is powered to: 0.31
## Study 2 is powered to: 0.32
```

```
colada17(simtot = 10000, n1 = 150, n2 = 300, d1 = .25, d2 = 0)
```

```
##
## Study 1 is powered to: 0.58
## Study 2 is powered to: 0.58
```

We can also reproduce the last example by adjusting the means and standard deviation. With 150 people, and a Cohen's d of 0.25 (the difference is 5, the sd 20, so $5/20 = 0.25$) we should reproduce the power for the simple effect.

```
string <- "2b*2b"
n <- 150
mu <- c(20, 20, 20, 25) #All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 20
labelnames <- c("fruit", "apple", "banana",
               "hunger", "no hunger", "very hungry") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05 #We set the alpha level at 0.05.

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results[2,],
              caption = "ANOVA Results")
```

And changing the sample size to 300 should reproduce the power for the interaction in the ANOVA.

```
string <- "2b*2b"
n <- 300
```

Table 7.4: ANOVA Results

	power	partial_eta_squared	cohen_f	non_centrality
fruit:hunger	58.06	0.0039	0.0626	4.6875

```

mu <- c(20, 20, 20, 25) #All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 20
labelnames <- c("fruit", "apple", "banana",
               "hunger", "no hunger", "very hungry") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05 #We set the alpha level at 0.05.

exact_result <- ANOVA_exact(design_result, alpha_level = alpha_level)

```

```

## Power and Effect sizes for ANOVA tests
##           power partial_eta_squared cohen_f non_centrality
## fruit      58.06             0.0039  0.0626         4.6875
## hunger      58.06             0.0039  0.0626         4.6875
## fruit:hunger 58.06             0.0039  0.0626         4.6875
##
## Power and Effect sizes for contrasts
##
## p_fruit_apple_hunger_no hunger_fruit_apple_hunger_very hungry    power
## p_fruit_apple_hunger_no hunger_fruit_banana_hunger_no hunger      5.00
## p_fruit_apple_hunger_no hunger_fruit_banana_hunger_very hungry    86.37
## p_fruit_apple_hunger_very hungry_fruit_banana_hunger_no hunger     5.00
## p_fruit_apple_hunger_very hungry_fruit_banana_hunger_very hungry    86.37
## p_fruit_banana_hunger_no hunger_fruit_banana_hunger_very hungry    86.37
##
## effect_size
## p_fruit_apple_hunger_no hunger_fruit_apple_hunger_very hungry      0.00
## p_fruit_apple_hunger_no hunger_fruit_banana_hunger_no hunger        0.00
## p_fruit_apple_hunger_no hunger_fruit_banana_hunger_very hungry      0.25
## p_fruit_apple_hunger_very hungry_fruit_banana_hunger_no hunger       0.00
## p_fruit_apple_hunger_very hungry_fruit_banana_hunger_very hungry     0.25
## p_fruit_banana_hunger_no hunger_fruit_banana_hunger_very hungry     0.25

```

```

knitr::kable(exact_result$main_results[3,],
             caption = "ANOVA Results")

```

Now if we look at the power analysis table for the last simulation, we see that the power for the ANOVA is the same for the main effect of fruit, the main effect of hunger, and the main effect of the interaction. All the effect sizes are equal as well. We can understand why if we look at the means in a 2x2 table:

```

mean_mat <- t(matrix(mu,
                    nrow = 2,
                    ncol = 2)) #Create a mean matrix

```

```
rownames(mean_mat) <- c("apple", "banana")
colnames(mean_mat) <- c("no hunger", "very hungry")
mean_mat
```

```
##      no hunger very hungry
## apple      20      20
## banana     20      25
```

The first main effect tests the marginal means if we sum over rows, 22.5 vs 20.

```
rowMeans(mean_mat)
```

```
##  apple banana
##  20.0   22.5
```

The second main effect tests the marginal means over the rows, which is also 22.5 vs 20.

```
colMeans(mean_mat)
```

```
##  no hunger very hungry
##    20.0      22.5
```

The interaction tests whether the average effect of hunger on liking fruit differs in the presence of bananas.

In the presence of bananas the effect of hunger on the desirability of fruit is 5 scalepoints. The average effect (that we get from the marginal means) of hunger on fruit desirability is 2.5 (22.5-20). In other words, the interaction tests whether the difference effect between hunger and no hunger is different in the presence of an apple versus in the presence of a banana.

Mathematically the interaction effect is computed as the difference between a cell mean and the grand mean, the marginal mean in row i and the grand mean, and the marginal mean in column j and grand mean. For example, for the very hungry-banana condition this is 25 (the value in the cell) - (21.25 [the grand mean] + 1.25 [the marginal mean in row 2, 22.5, minus the grand mean of 21.25] + 1.25 [the marginal mean in column 2, 22.5, minus the grand mean of 21.25]). $25 - (21.25 + (22.5-21.25) + (22.5-21.25)) = 1.25$.

We can repeat this for every cell, and get for no hunger-apple: $20 - (21.25 + (20-21.25) + (20-21.25)) = 1.25$, for very hungry apple: $20 - (21.25 + (22.5-21.25) + (20-21.25)) = 1.25$, and no hunger-banana: $20 - (21.25 + (20-21.25) + (22.5-21.25)) = 1.25$. These values are used to calculate the sum of squares.

```
a1 <- mean_mat[1,1] - (mean(mean_mat) +
                      (mean(mean_mat[1,]) - mean(mean_mat)) +
                      (mean(mean_mat[,1]) - mean(mean_mat)))
a2 <- mean_mat[1,2] - (mean(mean_mat) +
                      (mean(mean_mat[1,]) - mean(mean_mat)) +
                      (mean(mean_mat[,2]) - mean(mean_mat)))
b1 <- mean_mat[2,1] - (mean(mean_mat) +
                      (mean(mean_mat[2,]) - mean(mean_mat)) +
                      (mean(mean_mat[,1]) - mean(mean_mat)))
b2 <- mean_mat[2,2] - (mean(mean_mat) +
                      (mean(mean_mat[2,]) - mean(mean_mat)) +
                      (mean(mean_mat[,2]) - mean(mean_mat)))

SS_ab <- n * sum(c(a1, a2, b1, b2)^2)
```

The sum of squares is dependent on the sample size, as can be seen in the code above. The larger the sample size, the larger the sum of squares, and therefore (all else equal) the larger the F -statistic, and the smaller the p -value. We see from the simulations that all three tests have the same effect size, and therefore the same power.

Interactions can have more power than main effects if the effect size of the interaction is larger than the effect size of the main effects. An example of this is a cross-over interaction. For example, let's take a 2x2 matrix of means with a crossover interaction:

```
mu <- c(25, 20, 20, 25)
mean_mat <- t(matrix(mu,
                     nrow = 2,
                     ncol = 2)) #Create a mean matrix
rownames(mean_mat) <- c("apple", "banana")
colnames(mean_mat) <- c("no hunger", "very hungry")
mean_mat
```

```
##      no hunger very hungry
## apple      25      20
## banana     20      25
```

Neither of the main effects is now significant, as the marginal means are 22.5 vs 22.5 for both main effects.

The interaction is much stronger, however. We are testing whether the average effect of hunger on the desirability of fruit is different in the presence of bananas. Since the average effect is 0, and the effect of hunger on the desirability of bananas is 5, so the effect size is now twice as large.

```
string <- "2b*2b"
n <- 300
mu <- c(25, 20, 20, 25) #All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 20
labelnames <- c("fruit", "apple", "banana",
               "hunger", "no hunger", "very hungry") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05 #We set the alpha level at 0.05.

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
             caption = "ANOVA Results")
```

We can also reproduce the power analysis using the analytic function:

```
power_analytic <- power_twoway_between(design_result)
power_analytic$power_A
```

Table 7.5: ANOVA Results

	power	partial_eta_squared	cohen_f	non_centrality
fruit	5.0	0.0000	0.0000	0.00
hunger	5.0	0.0000	0.0000	0.00
fruit:hunger	99.1	0.0154	0.1252	18.75

```
## [1] 5
```

```
power_analytic$power_B
```

```
## [1] 5
```

```
power_analytic$power_AB
```

```
## [1] 99.10259
```


Chapter 8

Effect of Varying Designs on Power

Researchers might consider what the effects on the statistical power of their design is, when they add participants. Participants can be added to an additional condition, or to the existing design.

In a One-Way ANOVA adding a condition means, for example, going from a 1x2 to a 1x3 design. For example, in addition to a control and intensive training condition, we add a light training condition.

```
string <- "2b"
n <- 50
mu <- c(80, 86) #All means are equal - so there is no real difference.
sd <- 10
labelnames <- c("Condition", "control", "intensive_training")
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
# Power for the given N in the design_result
power_oneway_between(design_result)$power
```

```
## [1] 84.38754
```

```
power_oneway_between(design_result)$Cohen_f
```

```
## [1] 0.3
```

```
power_oneway_between(design_result)$eta_p_2
```

```
## [1] 0.08256881
```

```
simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 nsims = nsims,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_results,
              caption = "Simulated ANOVA Result")
```

```

\begin{table}[t]
\caption{(\#tab:variations_in_design)Simulated ANOVA Result}


|                 | power | effect_size |
|-----------------|-------|-------------|
| anova_Condition | 84.72 | 0.0917023   |


\end{table}

```

```

exact_result <- ANOVA_exact(design_result,
  alpha_level = alpha_level,
  verbose = FALSE)

knitr::kable(exact_result$main_results,
  caption = "Exact ANOVA Result")

```

```

\begin{table}[t]
\caption{(\#tab:variations_in_design)Exact ANOVA Result}


|           | power | partial_eta_squared | cohen_f | non centrality |
|-----------|-------|---------------------|---------|----------------|
| Condition | 84.39 | 0.0841              | 0.303   | 9              |


\end{table}

```

We now add a condition. Let's assume the 'light training' condition falls in between the other two means.

And we can see power across sample sizes

```

# Plot power curve (from 5 to 100)
plot_power(design_result, max_n = 100)

```

```

string <- "3b"
n <- 50
mu <-
c(80, 83, 86) #All means are equal - so there is no real difference.
sd <- 10
labelnames <-
c("Condition",
  "control",
  "light_training",
  "intensive_training")
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)
# Power for the given N in the design_result
power_oneway_between(design_result)$power

```

```
## [1] 76.16545
```

```
power_oneway_between(design_result)$Cohen_f
```


Table 8.1: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Condition	76.17	0.0577	0.2474	9

```
## [1] 0.244949
```

```
power_oneway_between(design_result)$eta_p_2
```

```
## [1] 0.05660377
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

We see that adding a condition that falls between the other two means reduces our power. Let's instead assume that the 'light training' condition is not different from the control condition. In other words, the mean we add is as extreme as one of the existing means.

```
# Plot power curve (from 5 to 100)
plot_power(design_result, max_n = 100)
```

```
string <- "3b"
n <- 50
mu <-
c(80, 80, 86) #All means are equal - so there is no real difference.
sd <- 10
labelnames <-
c("Condition",
  "control",
  "light_training",
  "intensive_training")
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)

# Power for the given N in the design_result
power_oneway_between(design_result)$power
```

```
## [1] 87.62941
```

```
power_oneway_between(design_result)$Cohen_f
```

```
## [1] 0.2828427
```

```
power_oneway_between(design_result)$eta_p_2
```

```
## [1] 0.07407407
```

Now power has increased. This is not always true. The power is a function of many factors in the design, including the effect size (Cohen's f) and the total sample size (and the degrees of freedom and number of groups). But as we will see below, as we keep adding conditions, the power will reduce, even if initially, the power might increase.

```
# Plot power curve (from 5 to 100)
plot_power(design_result,max_n = 100)
```

It helps to think of these different designs in terms of either partial eta-squared, or Cohen's f (the one can easily be converted into the other).

```
#Two groups
mu <- c(80, 86)
sd = 10
n <- 50 #sample size per condition
mean_mat <- t(matrix(mu,
nrow = 2,
ncol = 1)) #Create a mean matrix
# Using the sweep function to remove rowmeans from the matrix
mean_mat_res <- sweep(mean_mat, 2, rowMeans(mean_mat))
mean_mat_res
```

```
##      [,1] [,2]
## [1,]   -3    3
```

```
MS_a <- n * (sum(mean_mat_res ^ 2) / (2 - 1))
MS_a
```

```
## [1] 900
```

```
SS_A <- n * sum(mean_mat_res ^ 2)
SS_A
```

```
## [1] 900
```

```
MS_error <- sd ^ 2
MS_error
```

```
## [1] 100
```

```
SS_error <- MS_error * (n * 2)
SS_error
```

```
## [1] 10000
```

```
eta_p_2 <- SS_A / (SS_A + SS_error)
eta_p_2
```

```
## [1] 0.08256881
```

```
f_2 <- eta_p_2 / (1 - eta_p_2)
f_2
```

```
## [1] 0.09
```

```
Cohen_f <- sqrt(f_2)
Cohen_f
```

```
## [1] 0.3
```

```
#Three groups
mu <- c(80, 83, 86)
sd = 10
n <- 50
mean_mat <- t(matrix(mu,
nrow = 3,
ncol = 1)) #Create a mean matrix
# Using the sweep function to remove rowmeans from the matrix
mean_mat_res <- sweep(mean_mat, 2, rowMeans(mean_mat))
mean_mat_res
```

```
##      [,1] [,2] [,3]
## [1,]   -3    0    3
```

```
MS_a <- n * (sum(mean_mat_res ^ 2) / (3 - 1))
MS_a
```

```
## [1] 450
```

```
SS_A <- n * sum(mean_mat_res ^ 2)
SS_A
```

```
## [1] 900
```

```
MS_error <- sd ^ 2
MS_error
```

```
## [1] 100
```

```
SS_error <- MS_error * (n * 3)
SS_error
```

```
## [1] 15000
```

```
eta_p_2 <- SS_A / (SS_A + SS_error)
eta_p_2
```

```
## [1] 0.05660377
```

```
f_2 <- eta_p_2 / (1 - eta_p_2)
f_2
```

```
## [1] 0.06
```

```
Cohen_f <- sqrt(f_2)
Cohen_f
```

```
## [1] 0.244949
```

The SS_A or the sum of squares for the main effect, is 900 for two groups, and the SS_error for the error term is 10000. When we add a group, SS_A is 900, and the SS_error is 15000. Because the added condition falls exactly on the grand mean (83), the sum of squared for this extra group is 0. In other words, it does nothing to increase the signal that there is a difference between groups. However, the sum of squares for the error, which is a function of the total sample size, is increased, which reduces the effect size. So, adding a condition that falls on the grand mean reduces the power for the main effect of the ANOVA. Obviously, adding such a group has other benefits, such as being able to compare the two means to a new third condition.

We already saw that adding a condition that has a mean as extreme as one of the existing groups also reduces the power. Let's again do the calculations step by step when the extra group has a mean as extreme as one of the two original conditions.

```
#Three groups
mu <- c(80, 80, 86)
sd = 10
n <- 50
mean_mat <- t(matrix(mu,
  nrow = 3,
  ncol = 1)) #Create a mean matrix
# Using the sweep function to remove rowmeans from the matrix
mean_mat_res <- sweep(mean_mat, 2, rowMeans(mean_mat))
mean_mat_res
```

```
##      [,1] [,2] [,3]
## [1,]   -2   -2    4
```

```
MS_a <- n * (sum(mean_mat_res ^ 2) / (3 - 1))
MS_a
```

```
## [1] 600
```

```
SS_A <- n * sum(mean_mat_res ^ 2)
SS_A
```

```
## [1] 1200
```

```

MS_error <- sd ^ 2
MS_error

## [1] 100

SS_error <- MS_error * (n * 3)
SS_error

## [1] 15000

eta_p_2 <- SS_A / (SS_A + SS_error)
eta_p_2

## [1] 0.07407407

f_2 <- eta_p_2 / (1 - eta_p_2)
f_2

## [1] 0.08

Cohen_f <- sqrt(f_2)
Cohen_f

## [1] 0.2828427

```

We see the sum of squares of the error stays the same - 15000 - because it is only determined by the standard error and the sample size, but not by the differences in the means. This is an increase of 5000 compared to the 2 group design. The sum of squares (the second component that determines the size of partial eta-squared) increases, which increases Cohen's f .

8.1 Within Designs

Now imagine our design described above was a within design. The means and sd remain the same. We collect 50 participants (instead of 100, or 50 per group, for the between design). Let's first assume the two samples are completely uncorrelated.

```

string <- "2w"
n <- 50
mu <-
c(80, 86) #All means are equal - so there is no real difference.
sd <- 10
labelnames <- c("Condition", "control", "intensive_training") #
design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)

power_oneway_within(design_result)$power

```

Table 8.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Condition	83.66	0.1552	0.4286	9

```
## [1] 83.66436
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

We see power is ever so slightly less than for the between subject design. This is due to the loss in degrees of freedom, which is $2(n-1)$ for between designs, and $n-1$ for within designs. But as the correlation increases, the power advantage of within designs becomes stronger.

```
string <- "3w"
n <- 50
mu <-
c(80, 83, 86) #All means are equal - so there is no real difference.
sd <- 10
labelnames <-
c("Condition",
  "control",
  "light_training",
  "intensive_training") #

design_result <- ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  labelnames = labelnames
)

power_oneway_within(design_result)$power
```

```
## [1] 75.70841
```

```
exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

When we add a condition in a within design where we expect the mean to be identical to the grand mean, we again see that the power decreases. This similarly shows that adding a condition that equals the grand mean to a within subject design does not come for free, but has a power cost.

Table 8.3: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Condition	75.71	0.0841	0.303	9

```

n <- 30
sd <- 10
r <- 0.5
string <- "2w"
mu <- c(0, 5) #All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2") #
design_result <-
ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)

power_oneway_within(design_result)$power

```

```
## [1] 75.39647
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.25
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.5085476
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 7.5
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 4.182964
```

```

string <- "3w"
mu <-
c(0, 0, 5) #All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2", "a3")

design_result <-
ANOVA_design(
  design = string,

```

```

n = n,
mu = mu,
sd = sd,
r = r,
labelnames = labelnames
)

power_oneway_within(design_result)$power

```

```
## [1] 79.37037
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.2357023
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.4152274
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 10
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 3.155932
```

```

string <- "4w"
mu <-
c(0, 0, 0, 5) #All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2", "a3", "a4") #
design_result <-
ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)

power_oneway_within(design_result)$power

```

```
## [1] 79.40126
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.2165064
```



```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.3595975
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 11.25
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 2.709402
```

```
string <- "5w"
mu <-
c(0, 0, 0, 0, 5)
#All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2", "a3", "a4", "a5")
```

```
design_result <-
ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)
power_oneway_within(design_result)$power
```

```
## [1] 78.38682
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.2
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.3216338
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 12
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 2.44988
```

```

string <- "6w"
mu <- c(0, 0, 0, 0, 0, 5)
#All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2", "a3", "a4", "a5", "a6") #
design_result <-
ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)
power_oneway_within(design_result)$power

```

```
## [1] 76.99592
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.186339
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.2936101
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 12.5
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 2.276603
```

```

string <- "7w"
mu <- c(0, 0, 0, 0, 0, 0, 5)
#All means are equal - so there is no real difference.
labelnames <- c("Factor_A", "a1", "a2", "a3",
               "a4", "a5", "a6", "a7")
design_result <-
ANOVA_design(
  design = string,
  n = n,
  mu = mu,
  sd = sd,
  r = r,
  labelnames = labelnames
)

power_oneway_within(design_result)$power

```

```
## [1] 75.4601
```

```
power_oneway_within(design_result)$Cohen_f
```

```
## [1] 0.1749636
```

```
power_oneway_within(design_result)$Cohen_f_SPSS
```

```
## [1] 0.2718301
```

```
power_oneway_within(design_result)$lambda
```

```
## [1] 12.85714
```

```
power_oneway_within(design_result)$F_critical
```

```
## [1] 2.151016
```

This set of designs where we increase the number of conditions demonstrates a common pattern where the power initially increases, but then starts to decrease. Again, the exact pattern (and when the power starts to decrease) depends on the effect size and sample size. Note also that the effect size (Cohen's f) decreases as we add conditions, but the increased sample size compensates for this when calculating power. When using power analysis software such as GPower, this is important to realize. You can't just power for a medium effect size, and then keep adding conditions under the assumption that the increased power you see in the program will become a reality. Increasing the number of conditions will reduce the effect size, and therefore, adding conditions will not automatically increase power (and might even decrease it).

Overall, the effect of adding conditions with an effect close to the grand mean reduces power quite strongly, and adding conditions with means close to the extreme of the current conditions will either slightly increase or decrease power.

Chapter 9

Error Control in Exploratory ANOVA

In a $2 \times 2 \times 2$ design, an ANOVA will give the test results for three main effects, three two-way interactions, and one three-way interaction. That's 7 statistical tests. The probability of making at least one Type 1 error in a single $2 \times 2 \times 2$ ANOVA is $1-(0.95)^7 = 30\%$.

```
string <- "2b*2b*2b"
n <- 50
mu <- c(20, 20, 20, 20, 20, 20, 20, 20, 20)
# All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 5
p_adjust = "none"
# "none" means we do not correct for multiple comparisons
labelnames <- c("condition1", "a", "b",
                "condition2", "c", "d",
                "condition3", "e", "f") #
# The label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05
#We set the alpha level at 0.05.

simulation_result <- ANOVA_power(design_result,
                                 alpha_level = alpha_level,
                                 verbose = FALSE)

knitr::kable(simulation_result$main_result)
```

	power	effect_size
anova_condition1	3.5	0.0023044
anova_condition2	4.3	0.0024821
anova_condition3	6.2	0.0027568
anova_condition1:condition2	4.0	0.0024001
anova_condition1:condition3	4.0	0.0023874
anova_condition2:condition3	4.7	0.0024557
anova_condition1:condition2:condition3	4.9	0.0025069

When there is no true effect, we formally do not have ‘power’ (which is defined as the probability of finding $p < \alpha$ if there is a true effect to be found) so the power column should be read as the ‘Type 1 error rate’. Because we have saved the power simulation in the ‘simulation_result’ object, we can perform calculations on the ‘sim_data’ dataframe that is stored. This dataframe contains the results for the nsims simulations (e.g., 10000 rows if you ran 10000 simulations) and stores the p-values and effect size estimates for each ANOVA. The first 7 columns are the p-values for the ANOVA, first the main effects of condition 1, 2, and 3, then three two-way interactions, and finally the threeway interaction.

We can calculate the number of significant results for each test (which should be 5%) by counting the number of significant p-values in each of the 7 rows:

```
apply(as.matrix(simulation_result$sim_data[(1:7)]), 2,
      function(x) round(mean(ifelse(x < alpha_level, 1, 0) * 100),4))
```

```
##              anova_condition1
##              3.5
##              anova_condition2
##              4.3
##              anova_condition3
##              6.2
##      anova_condition1:condition2
##              4.0
##      anova_condition1:condition3
##              4.0
##      anova_condition2:condition3
##              4.7
## anova_condition1:condition2:condition3
##              4.9
```

This is the Type 1 error rate for each test. When we talk about error rate inflation due to multiple comparisons, we are talking about the probability that you conclude there is an effect, when there is actually no effect, when there is a significant effect for the main effect of condition 1, or condition 2, or condition 3, or for the two-way interaction between condition 1 and 2, or condition 1 and 3, or condition 2 and 3, or in the threeway interaction.

To calculate this error rate we do not just add the 7 error rates (so $7 * 5\% = 35\%$). Instead, we calculate the probability that there will be at least one significant result in an ANOVA we perform. Some ANOVA results will have multiple significant results, just due to the Type 1 error rate (e.g., a significant result for the three-way interaction, and for the main effect of condition 1) but such an ANOVA is counted only once. If we calculate this percentage from our simulations, we see the number is indeed very close to $1 - (0.95)^7 = 30\%$.

```
sum(apply(as.matrix(simulation_result$sim_data[(1:7)]), 1,
          function(x)
            round(mean(ifelse(x < alpha_level, 1, 0) * 100),4)) > 0)/nsims*100
```

```
## [1] 2.81
```

The question is what we should do about this alpha inflation. It is undesirable if you perform exploratory ANOVA’s and are fooled too often by Type 1 errors, which will not replicate if you try to build on them. Therefore, you need to control the Type 1 error rate.

In the simulation code, which relies on the afex package, there is the option to set p_adjust. In the simulation above, p_adjust was set to “none”. This means no adjustment is made to which p-values are considered to be significant, and the alpha level is used as it is set in the simulation (above this was 0.05).

Afex relies on the `p.adjust` function in the `stats` package in R (more information is available [here](#)). From the package details:

The adjustment methods include the Bonferroni correction (“bonferroni”) in which the p-values are multiplied by the number of comparisons. Less conservative corrections are also included by Holm (1979) (“holm”), Hochberg (1988) (“hochberg”), Hommel (1988) (“hommel”), Benjamini & Hochberg (1995) (“BH” or its alias “fdr”), and Benjamini & Yekutieli (2001) (“BY”), respectively. A pass-through option (“none”) is also included. The first four methods are designed to give strong control of the family-wise error rate. There seems no reason to use the unmodified Bonferroni correction because it is dominated by Holm’s method, which is also valid under arbitrary assumptions.

Hochberg’s and Hommel’s methods are valid when the hypothesis tests are independent or when they are non-negatively associated (Sarkar, 1998; Sarkar and Chang, 1997). Hommel’s method is more powerful than Hochberg’s, but the difference is usually small and the Hochberg p-values are faster to compute.

The “BH” (aka “fdr”) and “BY” method of Benjamini, Hochberg, and Yekutieli control the false discovery rate, the expected proportion of false discoveries amongst the rejected hypotheses. The false discovery rate is a less stringent condition than the family-wise error rate, so these methods are more powerful than the others.

Let’s re-run the simulation twith the Holm-Bonferroni correction, which is simple and require no assumptions.

```
string <- "2b*2b*2b"
n <- 50
mu <- c(20, 20, 20, 20, 20, 20, 20, 20, 20)
#All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 5
p_adjust = "holm"
# Changed to Holm-Bonferroni
labelnames <- c("condition1", "a", "b",
                "condition2", "c", "d",
                "condition3", "e", "f") #
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
alpha_level <- 0.05

simulation_result <- ANOVA_power(design_result,
                                alpha_level = alpha_level,
                                p_adjust = p_adjust,
                                verbose = FALSE)

knitr::kable(simulation_result$main_results, caption = "ANOVA Results")
```

Table 9.1: ANOVA Results

	power	effect_size
anova_condition1	0.4	0.0023999
anova_condition2	0.9	0.0026417
anova_condition3	0.5	0.0024140
anova_condition1:condition2	0.9	0.0025337
anova_condition1:condition3	0.8	0.0025789
anova_condition2:condition3	0.2	0.0025044
anova_condition1:condition2:condition3	0.8	0.0024475

```
knitr::kable(simulation_result$pc_results, caption = "Pairwise Results")
```

```
sum(apply(as.matrix(simulation_result$sim_data[(1:7)]), 1,
  function(x)
    round(mean(ifelse(x < alpha_level, 1, 0) * 100),4)) > 0)/nsims*100
```

```
## [1] 0.42
```

We see it is close to 5%. Note that error rates have variation, and even in a few thousand simulations, the error rate in the sample of studies can easily be half a percentage point higher or lower. But *in the long run* the error rate should equal the alpha level. Furthermore, note that the Holm-Bonferroni method is slightly more powerful than the Bonferroni procedure (which is simply α divided by the number of tests). There are more powerful procedures to control the Type 1 error rate, which require more assumptions. For a small number of tests, the Holm-Bonferroni procedure works well. Alternative procedures to control error rates can be found in the multcomp R package.

Table 9.2: Pairwise Results

	power	effect_size
p_condition1_a_condition2_c_condition3_e_condition1_a_condition2_c_condition3_f	0.1	-0.0068067
p_condition1_a_condition2_c_condition3_e_condition1_a_condition2_d_condition3_e	0.2	-0.0028161
p_condition1_a_condition2_c_condition3_e_condition1_a_condition2_d_condition3_f	0.4	-0.0096595
p_condition1_a_condition2_c_condition3_e_condition1_b_condition2_c_condition3_e	0.0	-0.0037777
p_condition1_a_condition2_c_condition3_e_condition1_b_condition2_c_condition3_f	0.0	-0.0096226
p_condition1_a_condition2_c_condition3_e_condition1_b_condition2_d_condition3_e	0.3	-0.0068070
p_condition1_a_condition2_c_condition3_e_condition1_b_condition2_d_condition3_f	0.2	-0.0060517
p_condition1_a_condition2_c_condition3_f_condition1_a_condition2_d_condition3_e	0.4	0.0038510
p_condition1_a_condition2_c_condition3_f_condition1_a_condition2_d_condition3_f	0.1	-0.0039822
p_condition1_a_condition2_c_condition3_f_condition1_b_condition2_c_condition3_e	0.3	0.0029050
p_condition1_a_condition2_c_condition3_f_condition1_b_condition2_c_condition3_f	0.2	-0.0035766
p_condition1_a_condition2_c_condition3_f_condition1_b_condition2_d_condition3_e	0.2	-0.0006991
p_condition1_a_condition2_c_condition3_f_condition1_b_condition2_d_condition3_f	0.4	0.0008182
p_condition1_a_condition2_d_condition3_e_condition1_a_condition2_d_condition3_f	0.0	-0.0072246
p_condition1_a_condition2_d_condition3_e_condition1_b_condition2_c_condition3_e	0.1	-0.0008379
p_condition1_a_condition2_d_condition3_e_condition1_b_condition2_c_condition3_f	0.0	-0.0069519
p_condition1_a_condition2_d_condition3_e_condition1_b_condition2_d_condition3_e	0.3	-0.0040988
p_condition1_a_condition2_d_condition3_e_condition1_b_condition2_d_condition3_f	0.2	-0.0034044
p_condition1_a_condition2_d_condition3_f_condition1_b_condition2_c_condition3_e	0.4	0.0065698
p_condition1_a_condition2_d_condition3_f_condition1_b_condition2_c_condition3_f	0.0	0.0005416
p_condition1_a_condition2_d_condition3_f_condition1_b_condition2_d_condition3_e	0.1	0.0035931
p_condition1_a_condition2_d_condition3_f_condition1_b_condition2_d_condition3_f	0.2	0.0042557
p_condition1_b_condition2_c_condition3_e_condition1_b_condition2_c_condition3_f	0.0	-0.0064965
p_condition1_b_condition2_c_condition3_e_condition1_b_condition2_d_condition3_e	0.2	-0.0039178
p_condition1_b_condition2_c_condition3_e_condition1_b_condition2_d_condition3_f	0.1	-0.0024063
p_condition1_b_condition2_c_condition3_f_condition1_b_condition2_d_condition3_e	0.1	0.0029019
p_condition1_b_condition2_c_condition3_f_condition1_b_condition2_d_condition3_f	0.1	0.0034800
p_condition1_b_condition2_d_condition3_e_condition1_b_condition2_d_condition3_f	0.1	0.0013388

Chapter 10

Analytic Power Functions

For some designs it is possible to calculate power analytically, using closed functions.

10.1 One-Way Between Subject ANOVA

```
string <- "4b"
n <- 60
mu <- c(80, 82, 82, 86)
#All means are equal - so there is no real difference.
# Enter means in the order that matches the labels below.
sd <- 10
labelnames <- c("Factor_A", "a1", "a2", "a3", "a4")
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

We can also calculate power analytically with a `Superpower` function.

```
#using default alpha level of .05
power_oneway_between(design_result)$power
```

```
## [1] 81.21291
```

This is a generalized function for One-Way ANOVA's for any number of groups. It is in part based on code provided with the excellent book by Aberson (2019) *Applied Power Analysis for the Behavioral Sciences* (but Aberson's code allows for different *n* per condition, and different *sd* per condition).

Table 10.1: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
Factor_A	81.21	0.0461	0.2198	11.4

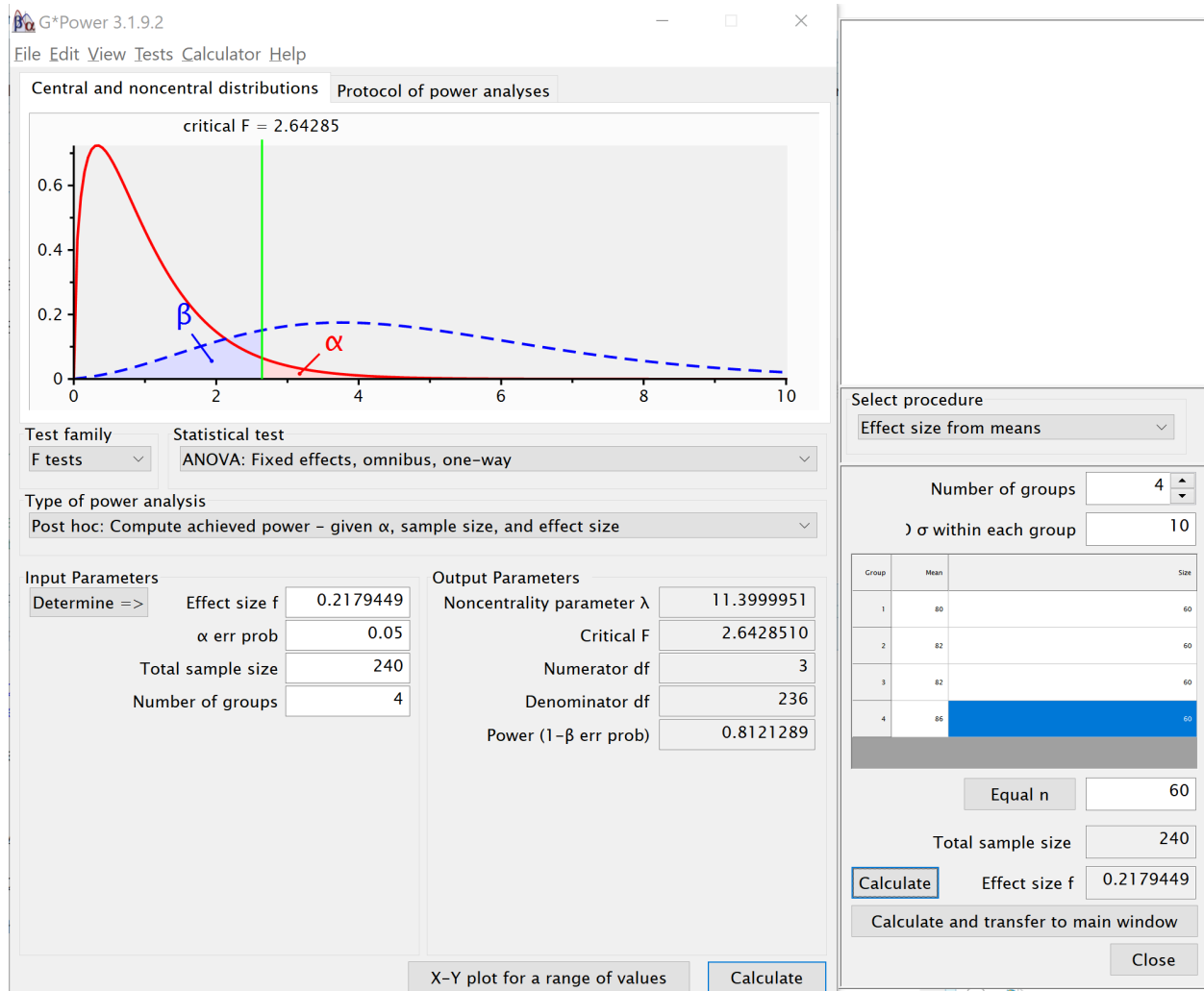
```
anova1f_4(m1 = 80, m2 = 82, m3 = 82, m4 = 86,
          s1 = 10, s2 = 10, s3 = 10, s4 = 10,
          n1 = 60, n2 = 60, n3 = 60, n4 = 60,
          alpha = .05)
```

We can also use the function in the `pwr` package. Note that we need to calculate f to use this function, which is based on the means and sd , as illustrated in the formulas above.

```
pwr.anova.test(n = 60,
               k = 4,
               f = 0.2179449,
               sig.level = 0.05)
```

```
##
##      Balanced one-way analysis of variance power calculation
##
##              k = 4
##              n = 60
##              f = 0.2179449
##      sig.level = 0.05
##      power = 0.8121289
##
## NOTE: n is number in each group
```

Finally, G*Power provides the option to calculate f from the means, sd and n for the cells. It can then be used to calculate power.



10.2 Two-way Between Subject Interaction

```
string <- "2b*2b"
n <- 20
mu <- c(20, 20, 20, 25)
# Enter means in the order that matches the labels below.
sd <- 5
labelnames <- c("A", "a1", "a2", "B", "b1", "b2")
# the label names should be in the order of the means specified above.

design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)
```

Table 10.2: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centralilty
A	59.79	0.0617	0.2565	5
B	59.79	0.0617	0.2565	5
A:B	59.79	0.0617	0.2565	5

```
exact_result <- ANOVA_exact(design_result,
                           alpha_level = alpha_level,
                           verbose = FALSE)
```

```
knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

```
#using default alpha level of .05
power_res <- power_twoway_between(design_result)
power_res$power_A
```

```
## [1] 59.78655
```

```
power_res$power_B
```

```
## [1] 59.78655
```

```
power_res$power_AB
```

```
## [1] 59.78655
```

We can use the function by Aberson, 2019, as well.

```
anova2x2(m1.1 = 20,
         m1.2 = 20,
         m2.1 = 20,
         m2.2 = 25,
         s1.1 = 5,
         s1.2 = 5,
         s2.1 = 5,
         s2.2 = 5,
         n1.1 = 20,
         n1.2 = 20,
         n2.1 = 20,
         n2.2 = 20,
         alpha = .05,
         all = "OFF")
```

10.3 3x3 Between Subject ANOVA

Table 10.3: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centralilty
Factor_A	44.86	0.0253	0.1612	4.4444
Factor_B	44.86	0.0253	0.1612	4.4444
Factor_A:Factor_B	64.34	0.0494	0.2280	8.8889

```

string <- "3b*3b"
n <- 20
#All means are equal - so there is no real difference.
mu <- c(20, 20, 20, 20, 20, 20, 20, 20, 20, 25)
# Enter means in the order that matches the labels below.
sd <- 5
labelnames <- c("Factor_A", "a1", "a2", "a3", "Factor_B", "b1", "b2", "b3")
# the label names should be in the order of the means specified above.
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             labelnames = labelnames)

exact_result <- ANOVA_exact(design_result,
                            alpha_level = alpha_level,
                            verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")

```

```

#using default alpha level of .05
power_res <- power_twoway_between(design_result)
power_res$power_A

```

```
## [1] 44.86306
```

```
power_res$power_B
```

```
## [1] 44.86306
```

```
power_res$power_AB
```

```
## [1] 64.34127
```

10.4 Two by two ANOVA, within design

Potvin & Schutz (2000) simulate a wide range of repeated measure designs. They give an example of a 3x3 design, with the following correlation matrix:

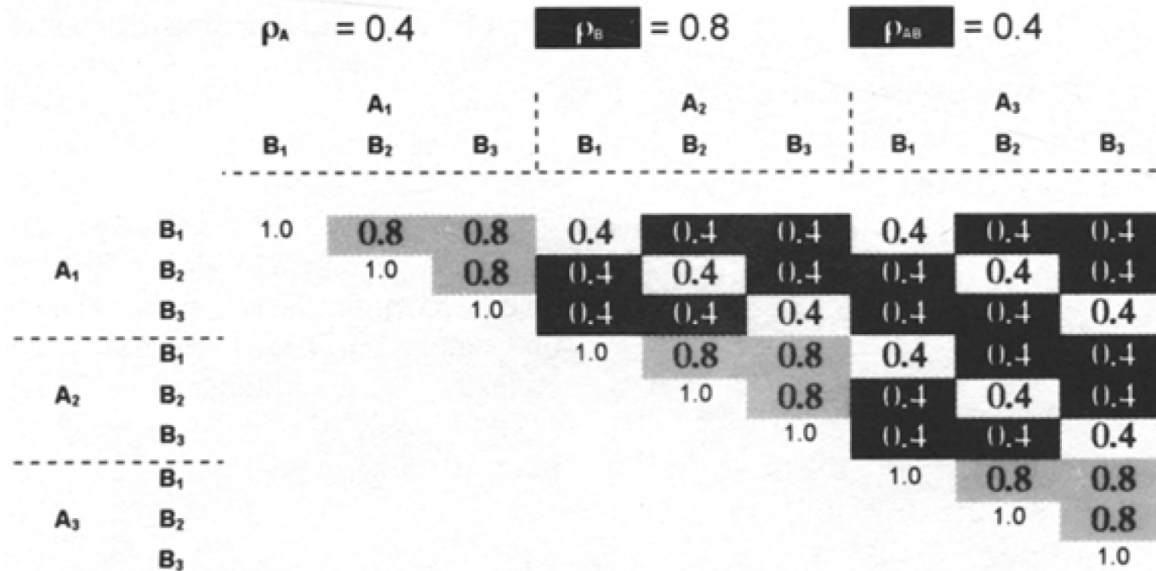
Example

Figure 1. Representation of a correlation matrix for a 3 (A) \times 3 (B) RM ANOVA: General form and numeric example. ρ_A and ρ_B represent the average correlation among the A and B (pooled) trials, respectively, and ρ_{AB} represents the average correlation among the AB coefficients having dissimilar levels.

Variances were set to 1 (so all covariance matrices in their simulations were identical). In this specific example, the white fields are related to the correlation for the A main effect (these cells have the same level for B, but different levels of A). The grey cells are related to the main effect of B (the cells have the same level of A, but different levels of B). Finally, the black cells are related to the AxB interaction (they have different levels of A and B). The diagonal (all 1) relate to cells with the same levels of A and B.

Potvin & Schulz (2000) examine power for 2x2 within ANOVA designs and develop approximations of the error variance. For a design with 2 within factors (A and B) these are:

$$\text{For the main effect of A: } \sigma_e^2 = \sigma^2(1 - \bar{\rho}_A) + \sigma^2(q - 1)(\bar{\rho}_B - \bar{\rho}_{AB})$$

$$\text{For the main effect of B: } \sigma_e^2 = \sigma^2(1 - \bar{\rho}_B) + \sigma^2(p - 1)(\bar{\rho}_A - \bar{\rho}_{AB})$$

$$\text{For the interaction between A and B: } \sigma_e^2 = \sigma^2(1 - \rho_{\max}) - \sigma^2(\bar{\rho}_{\min} - \bar{\rho}_{AB})$$

We first simulate a within subjects 2x2 ANOVA design.

```
mu = c(2,1,4,2)
n <- 20
sd <- 5
r <- c(
  0.8, 0.5, 0.4,
    0.4, 0.5,
      0.8
)
string = "2w*2w"
labelnames = c("A", "a1", "a2", "B", "b1", "b2")
```


Table 10.4: Exact ANOVA Result

	power	partial_eta_squared	cohen_f	non_centrality
A	26.92	0.0952	0.3244	2
B	64.23	0.2400	0.5620	6
A:B	26.92	0.0952	0.3244	2

```
design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

We can use the `ANOVA_exact` function to evaluate this design since there is not a analytic solution in Superpower.

```
power_res <- ANOVA_exact(design_result = design_result)
```

```
## Power and Effect sizes for ANOVA tests
##      power partial_eta_squared cohen_f non_centrality
## A   26.92           0.0952  0.3244           2
## B   64.23           0.2400  0.5620           6
## A:B 26.92           0.0952  0.3244           2
##
## Power and Effect sizes for contrasts
##      power effect_size
## p_A_a1_B_b1_A_a1_B_b2 26.92    -0.3162
## p_A_a1_B_b1_A_a2_B_b1 39.70     0.4000
## p_A_a1_B_b1_A_a2_B_b2  5.00     0.0000
## p_A_a1_B_b2_A_a2_B_b1 64.23     0.5477
## p_A_a1_B_b2_A_a2_B_b2 13.60     0.2000
## p_A_a2_B_b1_A_a2_B_b2 76.52    -0.6325
```

```
power_res$main_results
```

```
##      power partial_eta_squared cohen_f non_centrality
## A   26.92           0.0952  0.3244           2
## B   64.23           0.2400  0.5620           6
## A:B 26.92           0.0952  0.3244           2
```

We can use `pwr2ppl` by Aberson (2019) to produce the same results.

```
win2F(  
  m1.1 = 2,  
  m2.1 = 1,  
  m1.2 = 4,  
  m2.2 = 2,  
  s1.1 = 5,  
  s2.1 = 5,  
  s1.2 = 5,  
  s2.2 = 5,  
  r12 = 0.8,  
  r13 = 0.5,  
  r14 = 0.4,  
  r23 = 0.4,  
  r24 = 0.5,  
  r34 = 0.8,  
  n = 20  
)
```

Chapter 11

Power Curve

Power is calculated for a specific value of an effect size, alpha level, and sample size. Because you often do not know the true effect size, it often makes more sense to think of the power curve as a function of the size of the effect. Although power curves can be calculated based on simulations for any design, we will use the analytic solution to calculate the power of ANOVA designs because these calculations are much faster. The basic approach is to calculate power for a specific pattern of means, a specific effect size, a given alpha level, and a specific pattern of correlations. This is one example:

```
#2x2 design
string = "2w*2w"
mu = c(0,0,0,0.5)
n <- 20
sd <- 1
r <- 0.5
labelnames = c("A", "a1", "a2", "B", "b1", "b2")

design_result <- ANOVA_design(design = string,
                             n = n,
                             mu = mu,
                             sd = sd,
                             r = r,
                             labelnames = labelnames)

exact_result <- ANOVA_exact(design_result,
                             alpha_level = alpha_level,
                             verbose = FALSE)

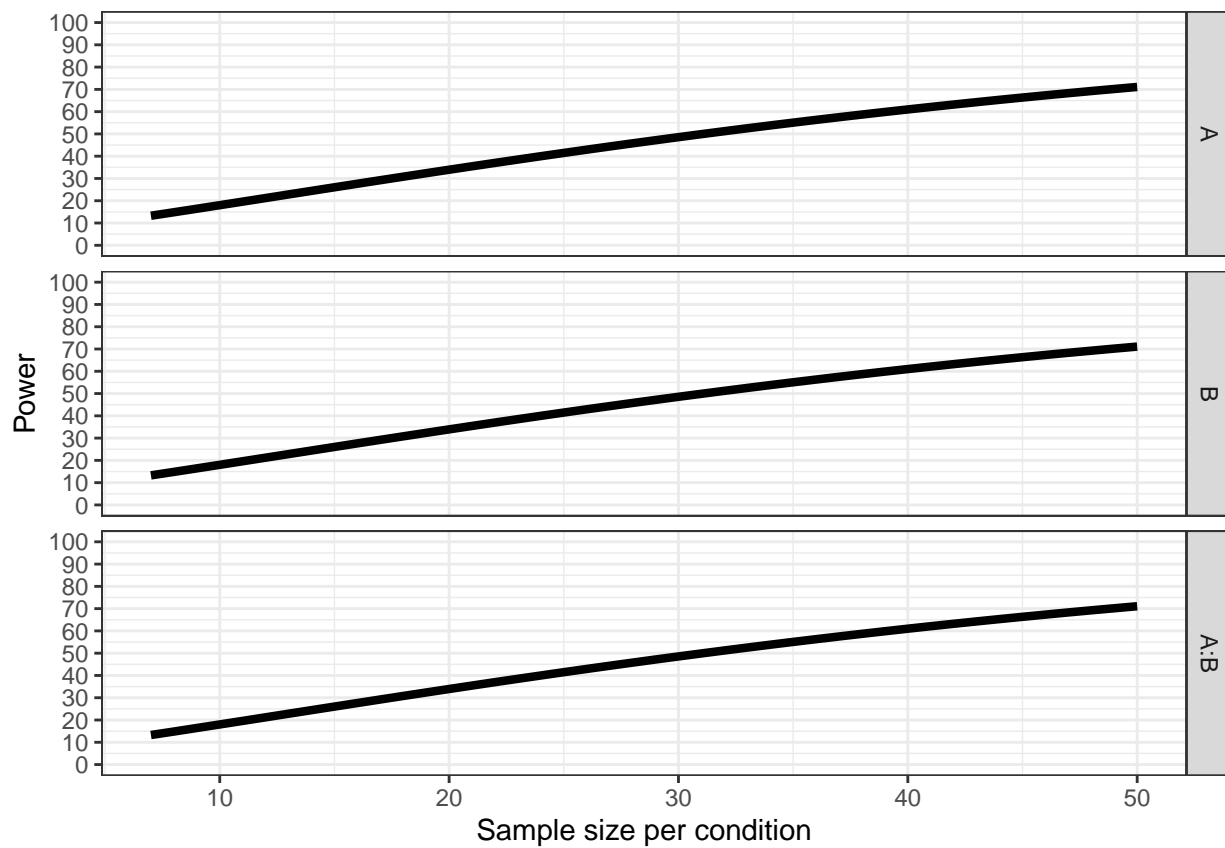
knitr::kable(exact_result$main_results,
              caption = "Exact ANOVA Result")
```

We can make these calculations for a range of sample sizes, to get a power curve. We created a simple function that performs these calculations across a range of sample sizes (from $n = 3$ to max_ , a variable you can specify in the function).

```
p_a <- plot_power(design_result,
                  max_n = 50)
p_a$plot_ANOVA
```

Table 11.1: Exact ANOVA Result

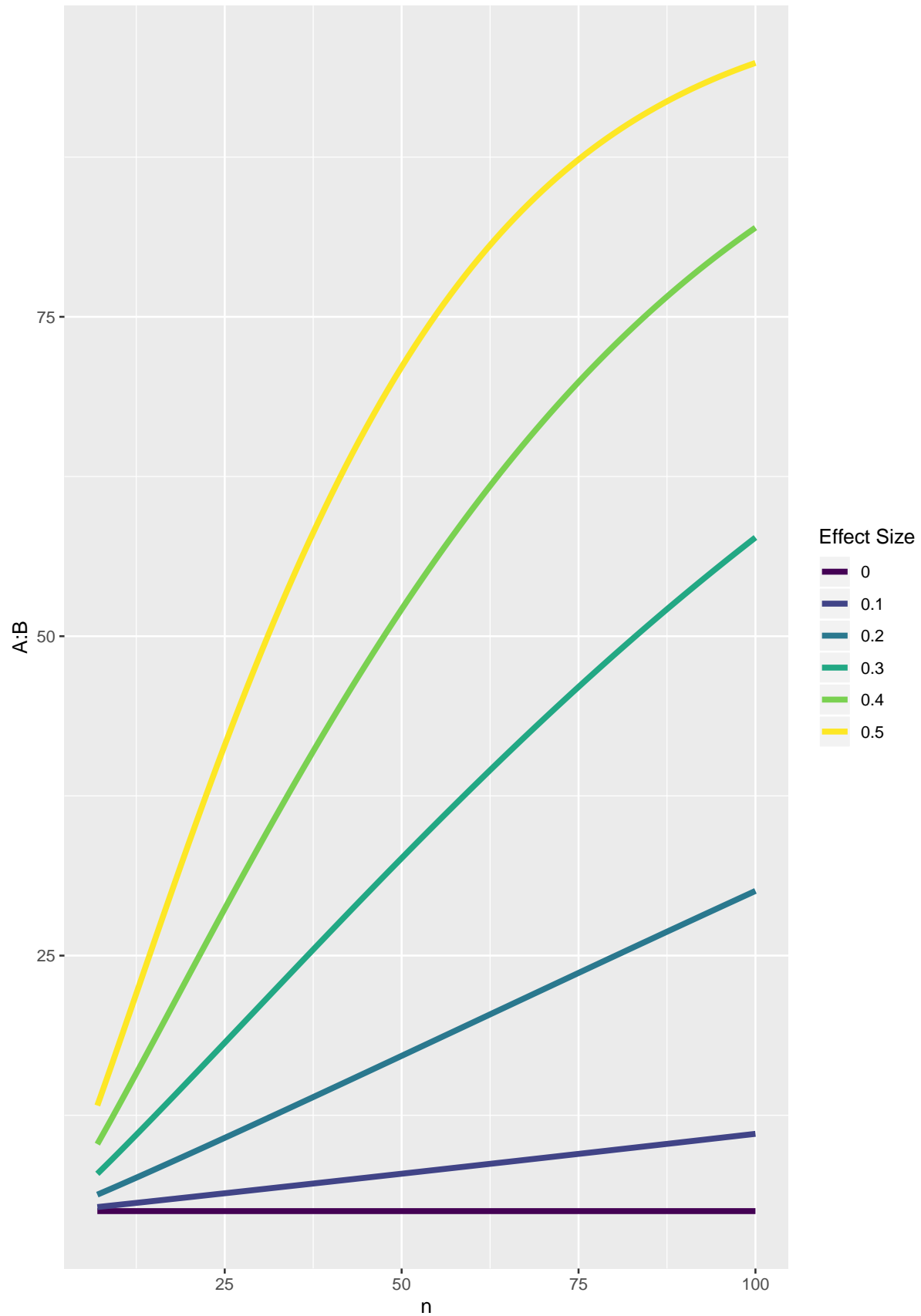
	power	partial_eta_squared	cohen_f	non_centralilty
A	32.36	0.1163	0.3627	2.5
B	32.36	0.1163	0.3627	2.5
A:B	32.36	0.1163	0.3627	2.5



Chapter 12

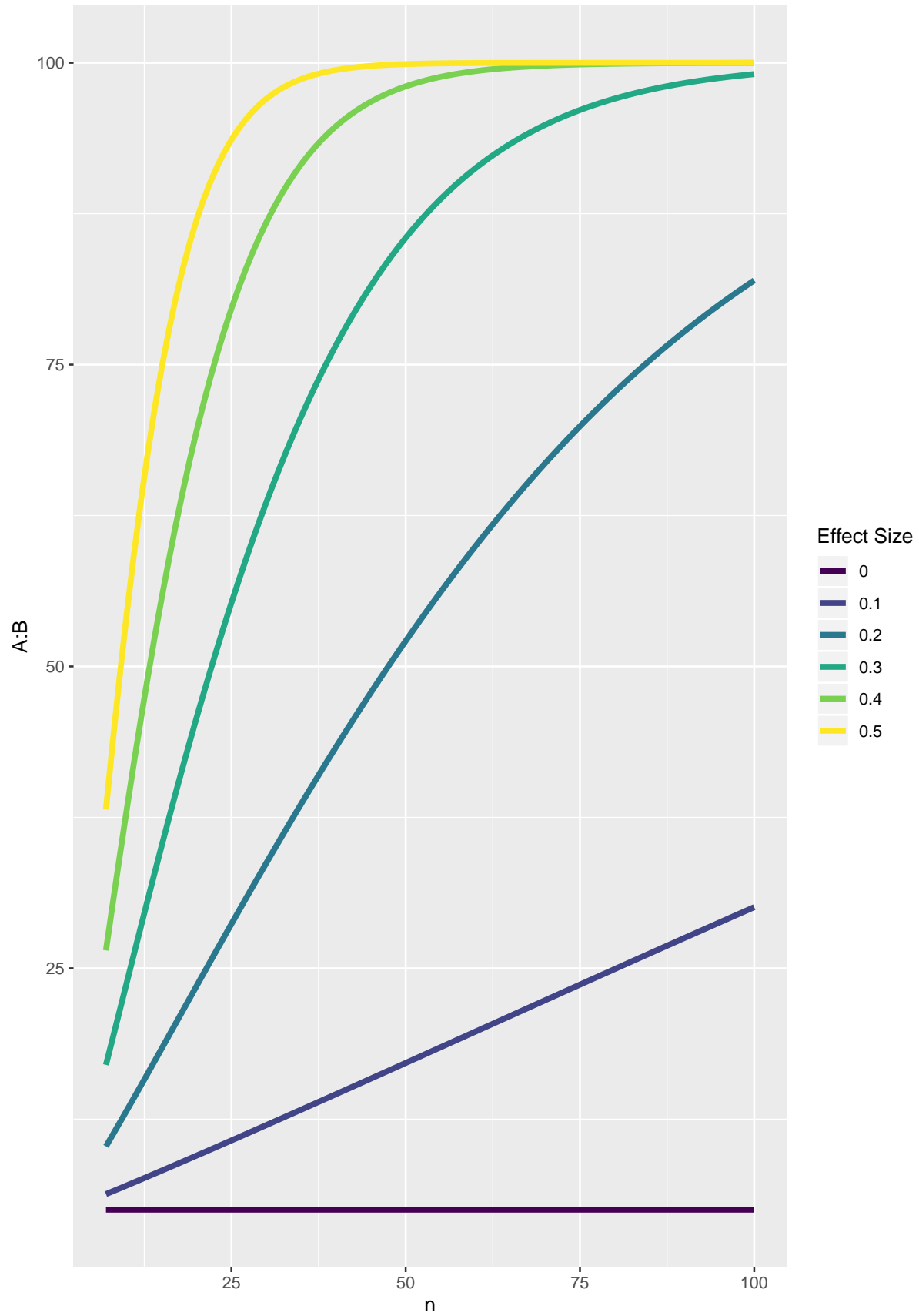
Explore increase in effect size for moderated interactions.

The design has means 0, 0, 0, 0, with one cell increasing by 0.1, up to 0, 0, 0, 0.5. The standard deviation is set to 1. The correlation between all variables is 0.5.



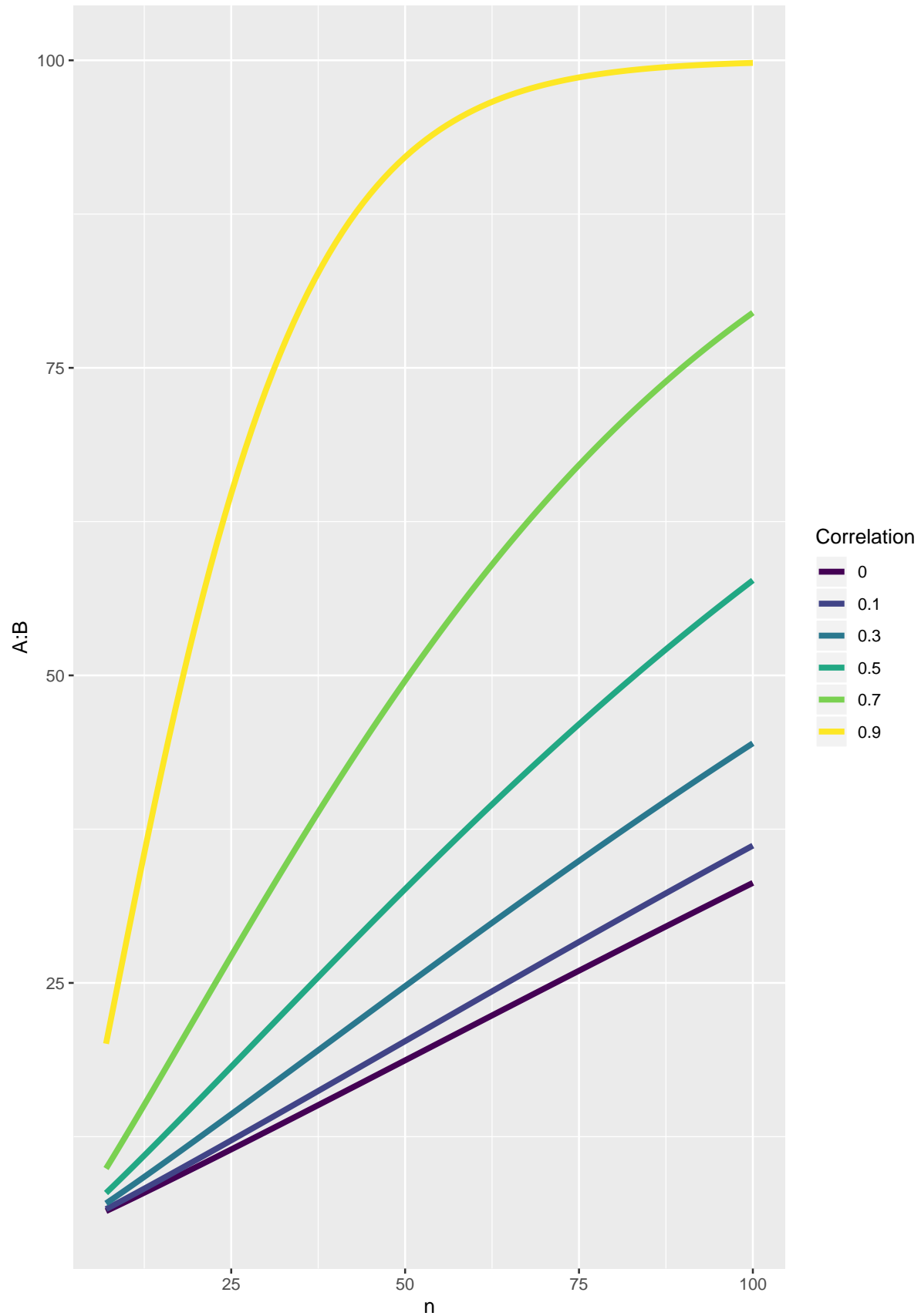
12.1 Explore increase in effect size for cross-over interactions.

The design has means 0, 0, 0, 0, with two cells increasing by 0.1, up to 0.5, 0, 0, 0.5. The standard deviation is set to 1. The correlation between all variables is 0.5.



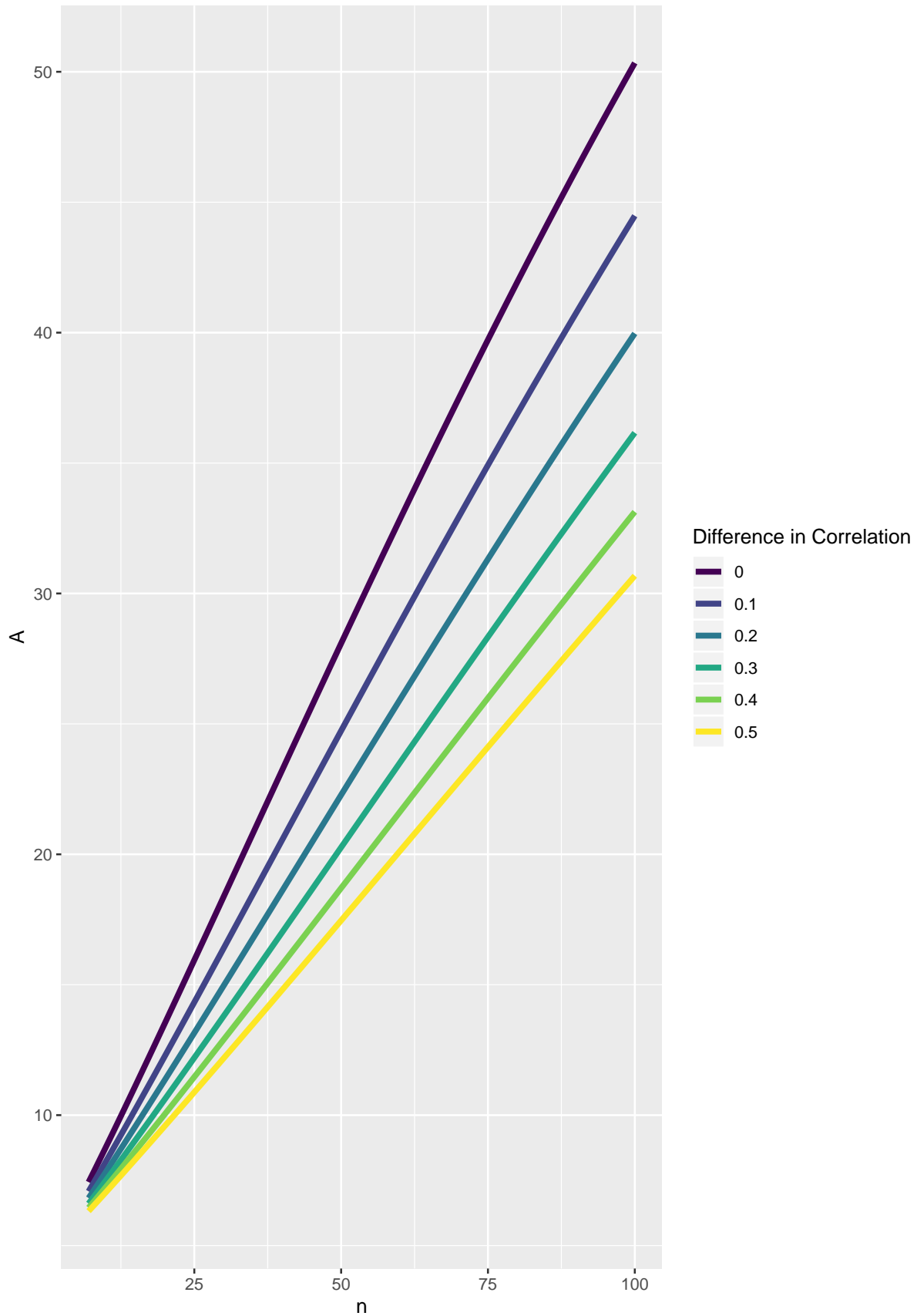
12.2 Explore increase in correlation in moderated interactions.

The design has means 0, 0, 0, 0.3. The standard deviation is set to 1. The correlation between all variables increases from 0 to 0.9.



12.3 Increasing correlation in on factor decreases power in second factor

As Potvin and Schutz (2000) write: “The more important finding with respect to the effect of r on power relates to the effect of the correlations associated with one factor on the power of the test of the main effect of the other factor. Specifically, if the correlations among the levels of B are larger than those within the AB matrix (i.e., $r_B - r_{AB} > 0.0$), there is a reduction in the power for the test of the A effect (and the test on B is similarly affected by the A correlations).” We see this in the plots below. As the correlation of the A factor increases from 0.4 to 0.9, we see the power for the main effect of factor B decreases.



Chapter 13

Appendix 1: Direct Comparison to pwr2ppl

In this appendix we have included the performance of **Superpower** to the **pwr2ppl** package using Chris Aberson's examples in *Applied Power Analysis for the Behavioral Sciences* (2nd edition).

13.1 Examples from Chapter 5

13.1.1 Example 5.1/5.2

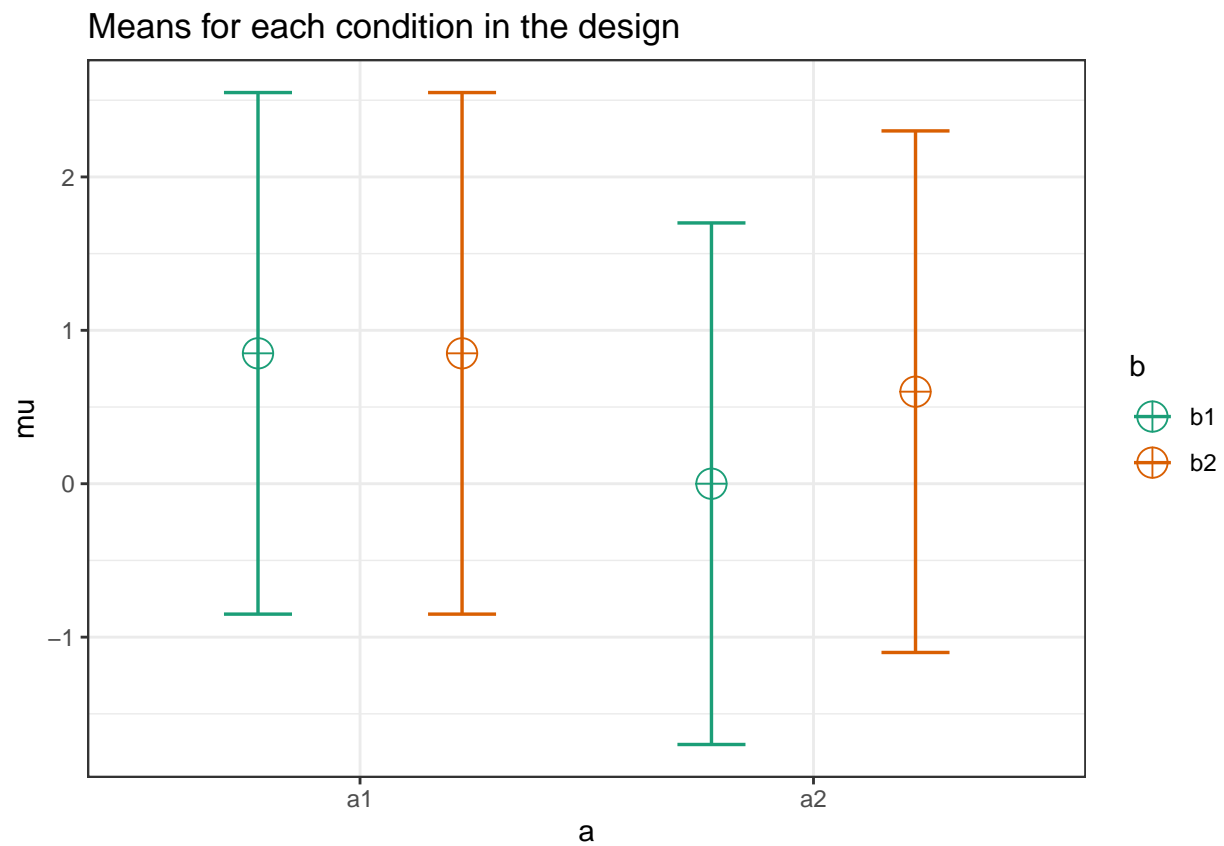
In this example, Aberson proposes a study expecting an average “score” of 80, 82, 82, and 86 for the control and three treatment groups respectively. The common standard deviation is 10 and the sample size per cell is 60.

```
design_result <- ANOVA_design(design = "4b",
  n = 60,
  sd = 10,
  mu = c(80, 82, 82, 86),
  labelnames = c("DORM",
    "Control",
    "T1",
    "T2",
    "T3"),
  plot = TRUE)
```


13.1.2 Example 5.3

Now a 2 x 2 between-subject ANOVA.

```
design_result <- ANOVA_design(design = "2b*2b",
                             n = 100,
                             sd = 1.7,
                             mu = c(.85, .85,
                                     0, .6),
                             plot = TRUE)
```



Now we calculate the analytical result from Superpower.

```
analytical_result <- power_twoway_between(design_result)
```

```
analytical_result$power_A
```

```
## [1] 89.75072
```

```
analytical_result$power_B
```

```
## [1] 42.10204
```

```
analytical_result$power_AB
```

```
## [1] 42.10204
```

The ANOVA_exact result.

```
exact_result <- ANOVA_exact(design_result, verbose = FALSE)
exact_result$main_results
```

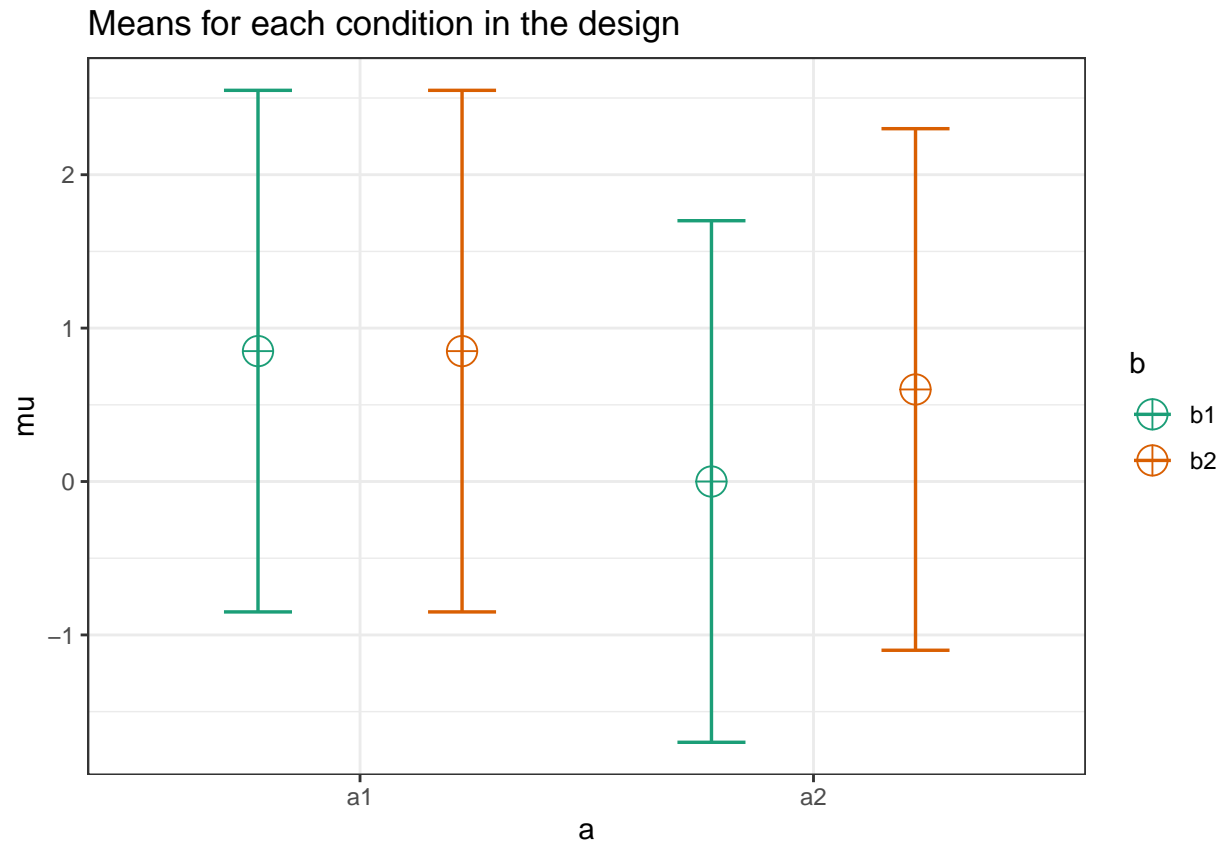
```
##      power partial_eta_squared cohen_f non centrality
## a   89.75           0.0258  0.1626      10.4671
## b   42.10           0.0078  0.0887       3.1142
## a:b 42.10           0.0078  0.0887       3.1142
```

And these match pwr2ppl. From Table 5.12.

```
pwr2ppl_result <- anova2x2(m1.1 = 0.85, m1.2 = 0.85, m2.1 = 0.00, m2.2 = 0.60,
                           s1.1 = 1.7, s1.2 = 1.7, s2.1 = 1.7, s2.2 = 1.7,
                           n1.1 = 100, n1.2 = 100, n2.1 = 100, n2.2 = 100,
                           alpha = .05)
```

Now we can increase the sample size to 250 per cell

```
design_result <- ANOVA_design(design = "2b*2b",
                             n = 250,
                             sd = 1.7,
                             mu = c(.85, .85,
                                     0, .6),
                             plot = TRUE)
```

Now we calculate the analytical result from Superpower.

```
analytical_result <- power_tway_between(design_result)
```

```
analytical_result$power_A
```

```
## [1] 99.91852
```

```
analytical_result$power_B
```

```
## [1] 79.60496
```

```
analytical_result$power_AB
```

```
## [1] 79.60496
```

The ANOVA_exact result.

```
exact_result <- ANOVA_exact(design_result, verbose = FALSE)
```

```
exact_result$main_results
```

```
##      power partial_eta_squared cohen_f non centrality
## a    99.92                0.0256  0.1621      26.1678
## b    79.61                0.0078  0.0884       7.7855
## a:b  79.61                0.0078  0.0884       7.7855
```

And these match `pwr2ppl`.

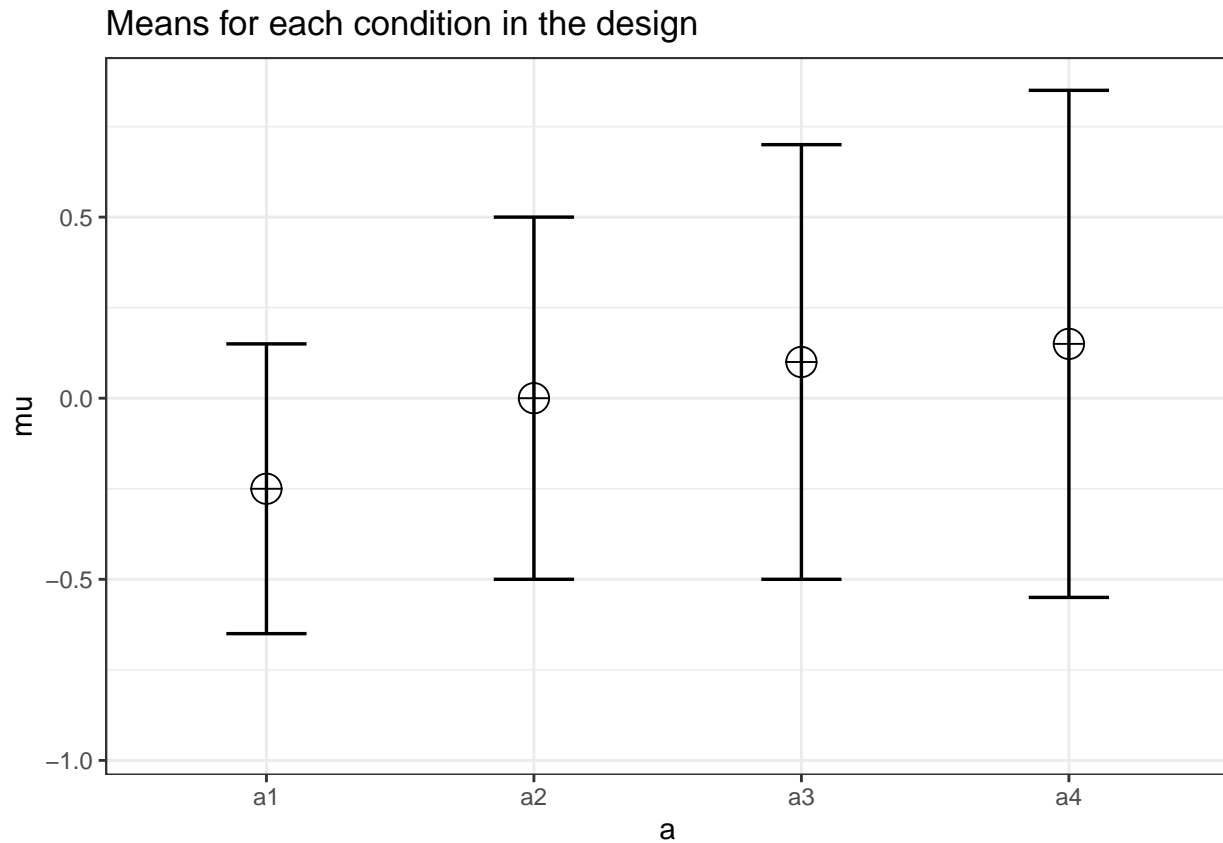
```
pwr2ppl_result <- anova2x2(m1.1 = 0.85, m1.2 = 0.85, m2.1 = 0.00, m2.2 = 0.60,
  s1.1 = 1.7, s1.2 = 1.7, s2.1 = 1.7, s2.2 = 1.7,
  n1.1 = 250, n1.2 = 250, n2.1 = 250, n2.2 = 250,
  alpha = .05)
```

13.2 Examples from Chapter 6

Repeated measures ANOVAs

13.2.1 Example from Table 6.2

```
design_result <- ANOVA_design(design = "4w",
  n = 25,
  sd = c(.4, .5, .6, .7),
  mu = c(-.25, .00, .10, .15),
  r = c(.50,
    .30,
    .15,
    .5,
    .30,
    .50),
  plot = TRUE)
```



```
design_result$cor_mat
```

```
##      a1 a2 a3 a4
## a1 1.00 0.5 0.3 0.15
## a2 0.50 1.0 0.5 0.30
## a3 0.30 0.5 1.0 0.50
## a4 0.15 0.3 0.5 1.00
```

There is no analytical result from **Superpower** when the correlations vary.

Now we prouce 3 **ANOVA_exact** results representing no sphericity correction, Greenhouse-Geisser, and Huynh-Feldt corrected results.

```
exact_result <- ANOVA_exact(design_result, verbose = FALSE)
```

```
exact_result$main_results
```

```
##  power partial_eta_squared cohen_f non centrality
## a 80.95                0.1405  0.4043          11.7671
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "GG",
                             verbose = FALSE)
```

```
exact_result$main_results
```

```
## power partial_eta_squared cohen_f non centrality
## a 74.46          0.1405  0.4043          9.5852
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "HF",
                             verbose = FALSE)

exact_result$main_results
```

```
## power partial_eta_squared cohen_f non centrality
## a 78.14          0.1405  0.4043          10.7526
```

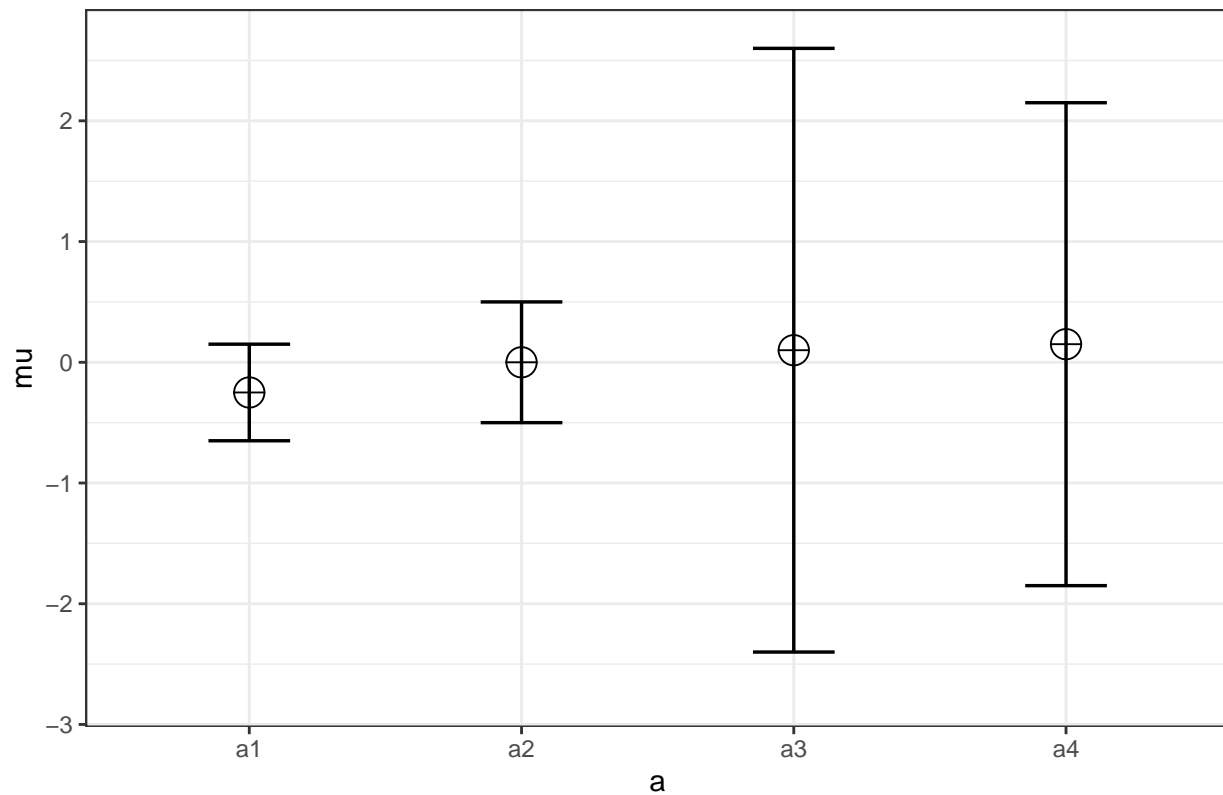
And these match pwr2ppl.

```
pwr2ppl_result <- win1F(m1 = -.25, m2 = .00, m3 = .10, m4 = .15,
                        s1 = .4, s2 = .5, s3 = .6, s4 = .7,
                        r12 = .50, r13 = .30,
                        r14 = .15, r23 = .5,
                        r24 = .30, r34 = .50,
                        n = 25)
```

13.2.2 Example from Table 6.6

```
design_result <- ANOVA_design(design = "4w",
                             n = 100,
                             sd = c(.4, .5, 2.5, 2),
                             mu = c(-.25, .00, .10, .15),
                             r = c(.50,
                                    .30,
                                    .1,
                                    .5,
                                    .30,
                                    .40),
                             plot = TRUE)
```

Means for each condition in the design



```
design_result$cor_mat
```

```
##      a1  a2  a3  a4
## a1  1.0  0.5  0.3  0.1
## a2  0.5  1.0  0.5  0.3
## a3  0.3  0.5  1.0  0.4
## a4  0.1  0.3  0.4  1.0
```

There is no analytical result from **Superpower** when the correlations vary.

Now we prouce 3 ANOVA_exact results representing no sphericity correction, Greenhouse-Geisser, and Huynh-Feldt corrected results.

```
exact_result <- ANOVA_exact(design_result, verbose = FALSE)
```

```
exact_result$main_results
```

```
##   power partial_eta_squared cohen_f non centrality
## a 39.75                0.015  0.1235          4.5292
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "GG",
                             verbose = FALSE)
```

```
exact_result$main_results
```

```
## power partial_eta_squared cohen_f non centrality
## a 31.79          0.015  0.1235          2.998
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "HF",
                             verbose = FALSE)

exact_result$main_results
```

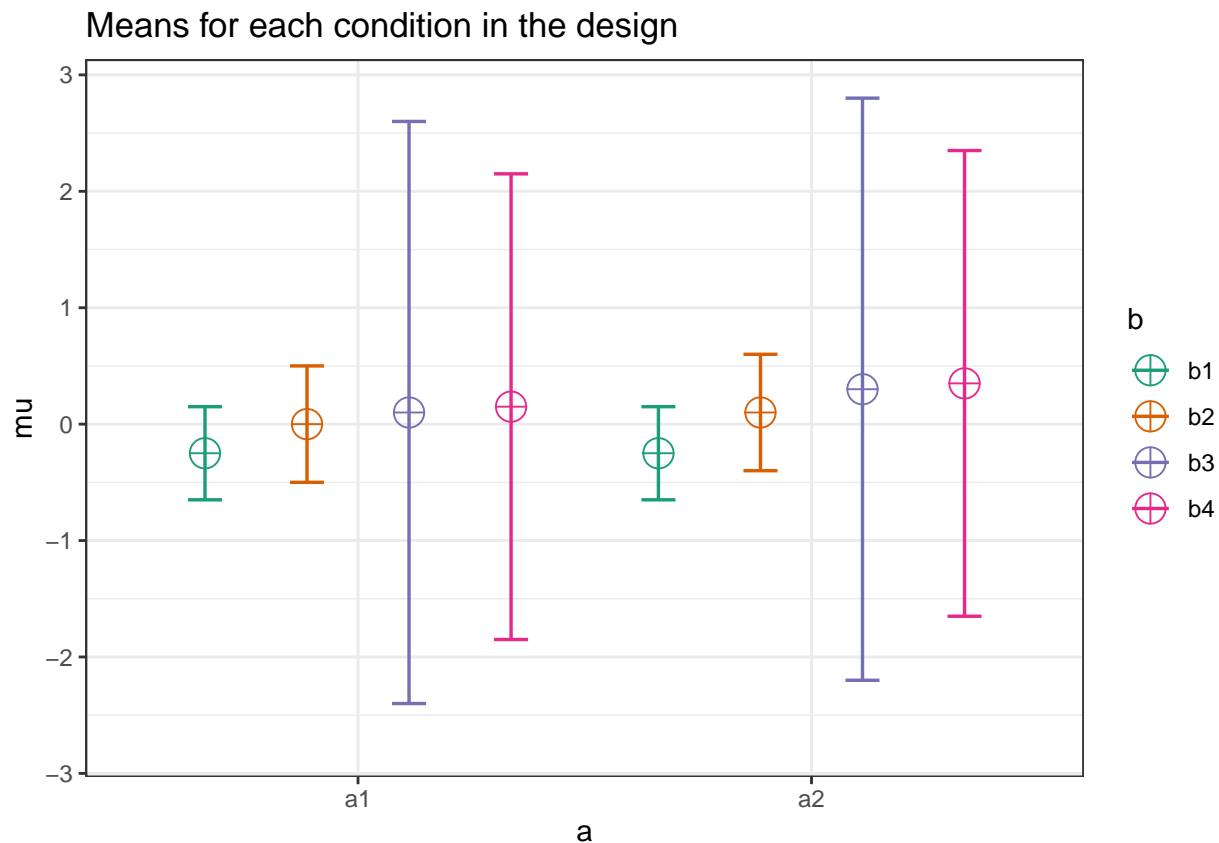
```
## power partial_eta_squared cohen_f non centrality
## a 32.12          0.015  0.1235          3.0591
```

And these match pwr2ppl.

```
pwr2ppl_result <- win1F(m1 = -.25, m2 = .00, m3 = .10, m4 = .15,
                        s1 = .4, s2 = .5, s3 = 2.5, s4 = 2.0,
                        r12 = .50, r13 = .30, r14 = .10,
                        r23 = .5, r24 = .30, r34 = .40,
                        n = 100)
```

13.2.3 Example from Table 6.8

```
design_result <- ANOVA_design(design = "2w*4w",
                              n = 80,
                              sd = c(.4, 0.5,
                                      2.5, 2.0,
                                      0.4, 0.5,
                                      2.5, 2.0),
                              mu = c(-0.25, 0.0,
                                      0.10, 0.15,
                                      -0.25, 0.10,
                                      0.30, 0.35),
                              r = c(.5),
                              plot = TRUE)
```



```
design_result$cor_mat
```

```
##      a1_b1 a1_b2 a1_b3 a1_b4 a2_b1 a2_b2 a2_b3 a2_b4
## a1_b1  1.0  0.5  0.5  0.5  0.5  0.5  0.5  0.5
## a1_b2  0.5  1.0  0.5  0.5  0.5  0.5  0.5  0.5
## a1_b3  0.5  0.5  1.0  0.5  0.5  0.5  0.5  0.5
## a1_b4  0.5  0.5  0.5  1.0  0.5  0.5  0.5  0.5
## a2_b1  0.5  0.5  0.5  0.5  1.0  0.5  0.5  0.5
## a2_b2  0.5  0.5  0.5  0.5  0.5  1.0  0.5  0.5
## a2_b3  0.5  0.5  0.5  0.5  0.5  0.5  1.0  0.5
## a2_b4  0.5  0.5  0.5  0.5  0.5  0.5  0.5  1.0
```

There is no analytical result from **Superpower** for two-way within subjects designs.

Now we prouce 3 ANOVA_exact results representing no sphericity correction, Greenhouse-Geisser, and Huynh-Feldt corrected results.

```
#In comparision to pwr2ppl the main effects are "flipped"
# e.g. Superpower a = pwr2ppl "B"
exact_result <- ANOVA_exact(design_result, verbose = FALSE)

exact_result$main_results
```

```
##      power partial_eta_squared cohen_f non centrality
## a      27.24                0.0232  0.1541          1.8762
```

```
## b    74.85          0.0401  0.2043          9.8948
## a:b  10.23          0.0035  0.0590          0.8255
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "GG",
                             verbose = FALSE)
```

```
exact_result$main_results
```

```
##      power partial_eta_squared cohen_f non centrality
## a    27.24          0.0232  0.1541          1.8762
## b    58.54          0.0401  0.2043          5.9072
## a:b   9.13          0.0035  0.0590          0.5072
```

```
exact_result <- ANOVA_exact(design_result,
                             correction = "HF",
                             verbose = FALSE)
```

```
exact_result$main_results
```

```
##      power partial_eta_squared cohen_f non centrality
## a    27.24          0.0232  0.1541          1.8762
## b    59.18          0.0401  0.2043          6.0353
## a:b   9.17          0.0035  0.0590          0.5188
```

And these match pwr2ppl.

```
pwr2ppl_result <- win2F(m1.1 = -.25, m2.1 = 0,
                        m3.1 = .10, m4.1 = .15,
                        m1.2 = -.25, m2.2 = .10,
                        m3.2 = .30, m4.2 = .35,
                        s1.1 = .4, s2.1 = .5,
                        s3.1 = 2.5, s4.1 = 2.0,
                        s1.2 = .4, s2.2 = .5,
                        s3.2 = 2.5, s4.2 = 2.0,
                        r = .5, n = 80)
```

13.3 Example from Chapter 7

Mixed effect ANOVA

13.3.1 From Table 7.2

In this case we must write out an entire correlation matrix. This means the diagonal element is equal to 1 and the off-diagonal elements corresponding to between-subjects factors are equal to zero.

```
design_result <- ANOVA_design("2b*4w",
                              n = 50,
                              sd = c(.4, .5, 0.6, .7,
```



```

        .4, .5, .6, .7),
r = c(1.0,0.5,0.3,0.15,0.0,0.0,0.0,0.0,
      0.5,1.0,0.5,0.3,0.0,0.0,0.0,0.0,
      0.3,0.5,1.0,0.5,0.0,0.0,0.0,0.0,
      0.15,0.3,0.5,1.0,0.0,0.0,0.0,0.0,
      0.0,0.0,0.0,0.0,1.0,0.5,0.3,0.15,
      0.0,0.0,0.0,0.0,0.5,1.0,0.5,0.3,
      0.0,0.0,0.0,0.0,0.3,0.5,1.0,0.5,
      0.0,0.0,0.0,0.0,0.15,0.3,0.5,1.0),
mu = c(-.25, 0.0, 0.10, 0.15,
      -.25,-.25,-.25,-.25))

design_result$cor_mat

```

```

##      a1_b1 a1_b2 a1_b3 a1_b4 a2_b1 a2_b2 a2_b3 a2_b4
## a1_b1 1.00  0.5  0.3  0.15 0.00  0.0  0.0  0.00
## a1_b2 0.50  1.0  0.5  0.30 0.00  0.0  0.0  0.00
## a1_b3 0.30  0.5  1.0  0.50 0.00  0.0  0.0  0.00
## a1_b4 0.15  0.3  0.5  1.00 0.00  0.0  0.0  0.00
## a2_b1 0.00  0.0  0.0  0.00 1.00  0.5  0.3  0.15
## a2_b2 0.00  0.0  0.0  0.00 0.50  1.0  0.5  0.30
## a2_b3 0.00  0.0  0.0  0.00 0.30  0.5  1.0  0.50
## a2_b4 0.00  0.0  0.0  0.00 0.15  0.3  0.5  1.00

```

Now the results from ANOVA_exact.

```

exact_result <- ANOVA_exact(design_result,
                           correction = "none",
                           verbose = FALSE)

exact_result$main_results

##      power partial_eta_squared cohen_f non centrality
## a   86.43                0.0888 0.3122          9.5493
## b   82.68                0.0385 0.2001          11.7671
## a:b 82.68                0.0385 0.2001          11.7671

```

```

exact_result <- ANOVA_exact(design_result,
                           correction = "GG",
                           verbose = FALSE)

exact_result$main_results

##      power partial_eta_squared cohen_f non centrality
## a   86.43                0.0888 0.3122          9.5493
## b   76.47                0.0385 0.2001          9.5852
## a:b 76.47                0.0385 0.2001          9.5852

```

```

exact_result <- ANOVA_exact(design_result,
                           correction = "HF",
                           verbose = FALSE)

exact_result$main_results

```

```
##      power partial_eta_squared cohen_f non centrality
## a   86.43                0.0888 0.3122          9.5493
## b   77.32                0.0385 0.2001          9.8486
## a:b 77.32                0.0385 0.2001          9.8486
```

And the results from `pwr2ppl`.

```
win1bg1(m1.1 = -.25, m2.1 = 0, m3.1 = 0.10, m4.1 = .15,
        m1.2 = -.25, m2.2 = -.25, m3.2 = -.25, m4.2 = -.25,
        s1.1 = .4, s2.1 = .5, s3.1 = 0.6, s4.1 = .7, s1.2 = .4,
        s2.2 = .5, s3.2 = .6, s4.2 = .7,
        n = 50,
        r1.2_1 = .5, r1.3_1 = .3, r1.4_1 = .15,
        r2.3_1 = .5, r2.4_1 = .3, r3.4_1 = .5,
        r1.2_2 = .5, r1.3_2 = .3, r1.4_2 = .15,
        r2.3_2 = .5, r2.4_2 = .3, r3.4_2 = .5)
```