Analysis of Variance

While t-tests allow comparisons of pairs of conditions, it is often more useful to compare all conditions in a single test. I'll talk about Analysis of Variance (ANOVA) in a later lecture. For now, here is the R code and output to run this on the ToothGrowth dataset.

```
> attach(ToothGrowth)
```

>oneway.test(len~supp)

One-way analysis of means (not assuming equal variances)

```
data: len and supp
F = 3.6683, num df = 1.000, denom df = 55.309, p-value = 0.06063
```

You will notice that this only compares supplement, and is thus a pairwise test (and produces the same result as the t-test). What we need to do is include both independent variables.

```
>res.aov2<-aov(len~supp+dose, data=ToothGrowth) >summary(res.aov2)
```

```
Df Sum Sq Mean Sq F value Pr(>F)
supp 1 205.4 205.4 11.45 0.0013 **
dose 1 2224.3 2224.3 123.99 6.31e-16 ***
Residuals 57 1022.6 17.9
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
```

This tells us that we have a significant main effect for 'supp' [F(1,57) = 11.45, p<0.005] and for 'dose' [F(1,57) = 123.99, p<0.0001]. What it doesn't tell us is whether there is a relationship between supplement and dose (as our boxplot suggests). For this, we need to add a further variable which describes the interaction between supplement and dose.

```
>res.aov3<-aov(len~supp+dose+supp:dose, data=ToothGrowth) >summary(res.aov3)
```

```
Df Sum Sq Mean Sq F value Pr(>F)
supp 1 205.4 205.4 12.317 0.000894 ***
dose 1 2224.3 2224.3 133.415 < 2e-16 ***
supp:dose 1 88.9 88.9 5.333 0.024631 *
Residuals 56 933.6 16.7
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Dealing with Non-Normal Data and Repeated Measures ANOVA
```

ii. Repeated Measures ANOVA

Conducting a Repeated Measures ANOVA in R is not the same as the other ANOVA tests we've looked at (and there are several approaches that can be used – in this lab we'll look at two of these). Whereas ANOVA assumes that data are drawn from independent populations (i.e., three separate groups of participants), in Repeated Measures ANOVA we use data from the same population (i.e., the same participants in three different conditions).

The assumptions for Repeated Measures ANOVA are:

- 1. Each participant has a value in each condition (if you are missing a value from a participant in one condition, you need to exclude that participant's data from all conditions);
- 2. Normality of distribution (check with Shapiro Wilke)
- 3. Sphericity (i.e., equality of variance across conditions)
- 4. Data for each condition should not contain outliers (extremes should be checked for and, if necessary, the data winsorized and refer back to 1. If this means that too many participants are removed, then you should consider a non-parametric alternative such as a Friedman test).

Using the transformed data set (b_t) as the Dependent Variable, we next define our Independent Variable (which is the level of Support received in the task) and Subject Variable.

```
> TPT<-c(0.38, 0.54, 0.92, 0.37, 0.26, 0.54, 0.92, 0.57, 0.37, 0.89, 0.21, 0.33, 0.25, 0.5, 0.51, 0.19, 0.642232, 0.288141, 0.197677, 0.232031, 0.256 769, 0.175277, 0.211903, 0.227281, 0.888331, 0.121795, 0.146731, 0.125118, 0.28487, 0.29217, 0.170797, 0.300612, 0.17235, 0.136446, 0.201687, 0.20784 4, 0.227901, 0.569835, 0.272328, 0.172863, 0.122078, 0.130318, 0.132493, 0.2167, 0.220268)
> Support<-as.factor(c("auto", "auto", "auto", "auto", "auto", "auto", "auto", "both", "self", "se
```

Notice that we are defining Part and Support as factors (Support is this is the Independent Variable in our experiment, and Part is the individual participants).

> Part<-as.factor(c(1,2,3,4,5,6,7,8,9,10,11,12,13,14,15))

We then set a data.frame for our analysis.

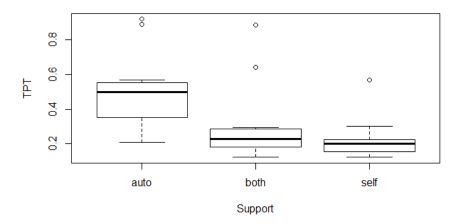
```
> expt<-data.frame(TPT, Support, Part)
```

If we you then type 'expt' you will get a print-out of the table as R sees it. This is useful just to check that the values are in the right columns.

You could also produce a graph of these data:

```
> plot(TPT~Support)
```

This should produce a graph like this:



i. The aov function

We have used an previously and this can also be applied to the Repeated Measures design (once we have indicated the factors in the model).

The line of interest in in the Part: Support table and we can see that there no significant effect of Support [F(2,28) = 12.66, p<0.05]

ii. Using a Linear Mixed Model

We can also run a Repeated Measures ANOVA by implementing a Linear Mixed Model using the library nlme.

> library(nlme)

We construct our linear model, with participant as the random term:

```
> analysis2<-lme(TPT~Support,random=~1|Part,data=expt)</pre>
```

And running the ANOVA to get our results:

> anova(analysis2)

```
numDF denDF F-value p-value
(Intercept) 1 28 86.04928 <.0001
Support 2 28 12.66313 1e-04
```

Notice that the F and p values here are the same as in (i.).

While this looks plausible, we haven't tested for sphericity. This can be done with R, but is not straightforward. One way to do this is to use functions in 'car' library.

```
> library(car)
> myMatrix<-matrix(TPT,nrow=15,ncol=3)</pre>
> myMatrix
               [,2]
      [,1]
 [1,] 0.38 0.190000 0.170797
 [2,] 0.54 0.642232 0.300612
 [3,] 0.92 0.288141 0.172350
 [4,] 0.37 0.197677 0.136446
 [5,] 0.26 0.232031 0.201687
 [6,] 0.54 0.256769 0.207844
 [7,] 0.92 0.175277 0.227901
 [8,] 0.57 0.211903 0.569835
 [9,] 0.37 0.227281 0.272328
[10,] 0.89 0.888331 0.172863
[11,] 0.21 0.121795 0.122078
[12,] 0.33 0.146731 0.130318
[13,] 0.25 0.125118 0.132493
[14,] 0.50 0.284870 0.216700
[15,] 0.51 0.292170 0.220268
> model<-lm(myMatrix ~ 1)</pre>
> design<-factor(c("auto", "both", "self"))</pre>
> options(contrasts = c("contr.sum", "contr.poly"))
> results<-Anova(model, idata=data.frame(design), idesign=~design, type =</pre>
"III")
> summary(results, multivariate = F)
Univariate Type III Repeated-Measures ANOVA Assuming Sphericity
            Sum Sq num Df Error SS den Df F value
                                                      Pr(>F)
                                           86.049 2.349e-07 ***
                           0.82381
(Intercept) 5.0634
                        1
                                        14
                        2
                           0.74554
                                           12.663 0.0001211 ***
design
            0.6743
                                        28
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Mauchly Tests for Sphericity
       Test statistic p-value
              0.98926 0.93223
design
Greenhouse-Geisser and Huynh-Feldt Corrections
 for Departure from Sphericity
        GG eps Pr(>F[GG])
design 0.98938 0.0001296 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
         HF eps
                  Pr(>F[HF])
design 1.151348 0.0001210696
```

```
Warning message:
In summary.Anova.mlm(results, multivariate = F) : HF eps > 1 treated as 1
```

(iii.) Non-Parametric ANOVA

If your data fail the Shapiro-Wilk test (and this means if any single condition produces a value >0.05) then you should apply a non-parametric test (or transform the data so then the distribution is normal). Let's assume that our original TPT data could not be transformed to normality. The Friedman test supports Repeated Measures ANOVA and is called with the command:

```
>friedman.test(TPT ~ Part | Support, data = expt)
```

This gives the following output:

Friedman rank sum test

```
data: TPT and Part and Support
Friedman chi-squared = 17.356, df = 14, p-value = 0.2377
```

MIXED ANOVA

Taken from https://cran.r-project.org/doc/contrib/Baron-rpsych.pdf

Suppose you have 1 between and 2 within Independent Varaiables.

Test of drug treatment effect by one between-subject: group (2 x 8Pp) and two within-subject factors (drug (2) and dose(3)).

We can either enter the data into R by hand and then construct the data frame for our analysis (which is what the next steps illustrate) or we can copy the data in the Ela.uni table (a couple of pages on from here), paste this in a spreadsheet, save it in tab-separated format and then import the dataset to R (which is much easier to do).

```
> Ela.mat <-matrix(c(</pre>
      19,22,28,16,26,22,
      11,19,30,12,18,28,
      20,24,24,24,22,29,
      21,25,25,15,10,26,
      18,24,29,19,26,28,
      17,23,28,15,23,22,
      20,23,23,26,21,28,
      14,20,29,25,29,29,
      16,20,24,30,34,36,
      26, 26, 26, 24, 30, 32,
      22,27,23,33,36,45,
      16, 18, 29, 27, 26, 34,
      19,21,20,22,22,21,
      20,25,25,29,29,33,
      21,22,23,27,26,35,
      17,20,22,23,26,28), nrow = 16, byrow = T)
```

```
Create multivariate format
```

```
> Ela.mul <- cbind.data.frame(subj=1:16, gp=factor(rep(1:2,rep(8,2))), Ela
.mat)
> dimnames(Ela.mul)[[2]] <-c("subj","gp","d11","d12","d13","d21","d22","d2</pre>
> Ela.mul
   subj gp d11 d12 d13 d21 d22 d23
                   22
                                      22
1
              19
                        28
                             16
                                 26
2
       2
           1
                                      28
              11
                   19
                        30
                             12
                                  18
3
       3
           1
                   24
                        24
                             24
                                  22
                                       29
              20
           1
                   25
                        25
                             15
4
       4
                                  10
                                      26
              21
5
       5
           1
                   24
                        29
                             19
              18
                                  26
                                       28
           1
                   23
                        28
                                  23
6
       6
              17
                             15
                                       22
           1
                   23
                        23
7
       7
              20
                             26
                                  21
                                       28
8
       8
           1
                        29
                                  29
                                       29
              14
                   20
                             25
       9
           2
9
              16
                   20
                        24
                             30
                                  34
                                       36
           2
      10
                                  30
10
              26
                   26
                        26
                             24
                                       32
           2
                        23
                             33
                                       45
11
      11
              22
                   27
                                  36
           2
      12
                        29
                             27
12
              16
                   18
                                  26
                                       34
           2
13
      13
              19
                   21
                        20
                             22
                                  22
                                       21
           2
                                  29
14
      14
              20
                   25
                        25
                             29
                                       33
           2
15
      15
              21
                   22
                        23
                             27
                                  26
                                       35
           2
16
      16
              17
                   20
                        22
                             23
                                  26
                                       28
> Ela.uni <- data.frame(effect = as.vector(Ela.mat),subj = factor(paste("s</pre>
", rep(1:16, 6), sep="")), gp = factor(paste("gp", rep(rep(c(1, 2), c(8,8)), 6), sep="")), drug = factor(paste("dr", rep(c(1, 2), c(48, 48)), sep=""))
, dose=factor(paste("do", rep(rep(c(1,2,3), rep(16, 3)), 2), sep="")), row.n
ames = NULL)
> Ela.uni
   effect subj gp drug dose
              s1 gp1
1
        19
                       dr1
                              do1
2
        11
              s2 gp1
                        dr1
                              do1
3
        20
              s3 gp1
                        dr1
                              do1
4
        21
              s4 gp1
                        dr1
                              do1
5
        18
              s5 gp1
                        dr1
                              do1
6
        17
              s6 gp1
                        dr1
                              do1
7
        20
              s7 gp1
                        dr1
                              do1
8
        14
              s8 gp1
                        dr1
                              do1
9
        16
              s9 gp2
                        dr1
                              do1
10
        26
             s10 gp2
                        dr1
                              do1
11
        22
                        dr1
                              do1
             s11 gp2
12
        16
             s12 gp2
                        dr1
                              do1
13
        19
             s13 gp2
                        dr1
                              do1
14
        20
             s14 gp2
                        dr1
                              do1
15
        21
             s15 gp2
                        dr1
                              do1
16
        17
             s16 gp2
                        dr1
                              do1
17
        22
                        dr1
              s1 gp1
                              do2
18
        19
              s2 gp1
                        dr1
                              do2
19
        24
                        dr1
              s3 gp1
                              do2
20
        25
              s4 gp1
                        dr1
                              do2
21
        24
              s5 gp1
                        dr1
                              do2
22
        23
              s6 gp1
                        dr1
                              do2
23
        23
              s7 gp1
                        dr1
                              do2
24
        20
              s8 gp1
                        dr1
                              do2
25
        20
              s9 gp2
                        dr1
                              do2
```

```
26
        26
                              do2
             s10 gp2
                        dr1
27
        27
             s11 gp2
                        dr1
                              do2
28
                        dr1
        18
             s12 gp2
                              do2
29
                        dr1
        21
             s13 gp2
                              do2
30
        25
             s14 gp2
                        dr1
                              do2
31
        22
             s15 gp2
                        dr1
                              do2
32
        20
             s16 gp2
                        dr1
                              do2
33
        28
              s1 gp1
                       dr1
                              do3
34
        30
              s2 gp1
                        dr1
                              do3
        24
35
              s3 gp1
                        dr1
                              do3
36
        25
              s4 gp1
                        dr1
                              do3
        29
37
              s5 gp1
                        dr1
                              do3
38
        28
              s6 gp1
                        dr1
                              do3
39
        23
              s7 gp1
                        dr1
                              do3
        29
40
              s8 gp1
                        dr1
                              do3
41
        24
              s9 gp2
                        dr1
                              do3
42
        26
             s10 gp2
                        dr1
                              do3
43
        23
             s11 gp2
                        dr1
                              do3
        29
44
             s12 gp2
                        dr1
                              do3
45
        20
             s13 gp2
                        dr1
                              do3
46
        25
             s14 gp2
                        dr1
                              do3
        23
47
             s15 gp2
                        dr1
                              do3
48
        22
             s16 gp2
                        dr1
                              do3
49
        16
              s1 gp1
                       dr2
                              do1
50
        12
              s2 gp1
                       dr2
                              do1
51
        24
              s3 gp1
                       dr2
                              do1
52
        15
              s4 gp1
                       dr2
                              do1
53
        19
              s5 gp1
                        dr2
                              do1
54
        15
              s6 gp1
                        dr2
                              do1
55
        26
              s7 gp1
                        dr2
                              do1
56
        25
              s8 gp1
                        dr2
                              do1
57
        30
              s9 gp2
                        dr2
                              do1
58
        24
             s10 gp2
                        dr2
                              do1
59
        33
             s11 gp2
                        dr2
                              do1
60
        27
             s12 gp2
                        dr2
                              do1
61
        22
             s13 gp2
                        dr2
                              do1
62
        29
             s14 gp2
                        dr2
                              do1
             s15 gp2
63
        27
                        dr2
                              do1
64
        23
             s16 gp2
                        dr2
                              do1
65
        26
              s1 gp1
                       dr2
                              do2
66
        18
              s2 gp1
                       dr2
                              do2
              s3 gp1
67
        22
                       dr2
                              do2
              s4 gp1
68
        10
                        dr2
                              do2
69
        26
              s5 gp1
                        dr2
                              do2
              s6 gp1
70
        23
                        dr2
                              do2
71
        21
              s7 gp1
                        dr2
                              do2
72
        29
                        dr2
                              do2
              s8 gp1
73
        34
              s9 gp2
                        dr2
                              do2
74
        30
             s10 gp2
                        dr2
                              do2
75
        36
                        dr2
             s11 gp2
                              do2
76
        26
             s12 gp2
                        dr2
                              do2
77
        22
             s13 gp2
                        dr2
                              do2
78
        29
             s14 gp2
                        dr2
                              do2
79
        26
             s15 gp2
                        dr2
                              do2
80
        26
             s16 gp2
                       dr2
                              do2
        22
81
              s1 gp1
                       dr2
                              do3
        28
              s2 gp1
82
                       dr2
                              do3
        29
83
              s3 gp1
                       dr2
                              do3
        26
84
              s4 gp1
                       dr2
                              do3
        28
85
              s5 gp1
                       dr2
                              do3
        22
86
              s6 gp1
                       dr2
                              do3
        28
87
              s7 gp1
                       dr2
                              do3
```

```
29
88
           s8 gp1
                   dr2
                         do3
89
       36
                   dr2
           s9 gp2
                         do3
90
       32
                   dr2
           s10 gp2
                         do3
       45
91
           s11 gp2
                    dr2
                         do3
92
       34
           s12 gp2
                    dr2
                         do3
93
       21
           s13 gp2
                    dr2
                         do3
94
       33
          s14 gp2
                    dr2
                         do3
95
       35
          s15 gp2
                   dr2
                        do3
                   dr2
96
       28
          s16 gp2
                        do3
> analysis<-aov(effect ~ gp * drug * dose + Error(subj/(dose+drug)), data=</pre>
Ela.uni)
> summary(analysis)
Error: subj
          Df Sum Sq Mean Sq F value Pr(>F)
               270 270.01 7.092 0.0185 *
          1
Residuals 14
                533
                      38.07
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Error: subj:dose
          Df Sum Sq Mean Sq F value
                                      Pr(>F)
                      379.4 36.510 1.58e-08 ***
dose
          2 758.8
              42.3
                       21.1
                              2.034
                                        0.15
gp:dose
Residuals 28 291.0
                       10.4
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Error: subj:drug
          Df Sum Sq Mean Sq F value Pr(>F)
                              13.00 0.00287 **
drug
          1 348.8
                      348.8
                              12.16 0.00362 **
gp:drug
           1
             326.3
                      326.3
             375.6
Residuals 14
                       26.8
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Error: Within
            Df Sum Sq Mean Sq F value Pr(>F)
drug:dose
             2 12.06
                         6.031
                                 0.682 0.514
gp:drug:dose 2 14.81
                                 0.837 0.444
                         7.406
            28 247.79
                         8.850
Residuals
```