

Visualizing Linear Models: An R Bag of Tricks Session 3: Examples & Extensions

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https://friendly.github.io/VisMLM-course/

Today's topics

- Manipulating & visualizing models
- MANOVA examples
 - Distinguishing among psychiatric groups
 - Robust MLMs: down-weighting outliers
- Multivariate regression
 - PA tests & ability
 - Canonical correlation
 - MANCOVA & homogeneity of regression
- Homogeneity of (co)variance
 - Visualizing Box's M test

Manipulating & visualizing models

- R modeling functions [lm(), glm(), ...] return model objects, but these are "messy"
 - extracting coefficients takes several steps: data.frame(coef(mymod))
 - some info (R², F, p.value) is computed in print() method, not stored
 - can't easily combine models
- Some have associated plotting functions
 - plot(model): diagnostic plots
 - car package: many model plot methods
 - effects package: plot effects for model terms
- But what if you want to:
 - make a table of model summary statistics
 - fit a collection of models, compare, summarize or visualize them?

broom

broom: manipulating models

- The broom package turns model objects into tidy data frames
 - glance(models) extracts model-level summary statistics (R², df, AIC, BIC)
 - tidy(models) extracts coefficients, SE, p-values
 - augment(models) extracts observation-level info (residuals, ...)

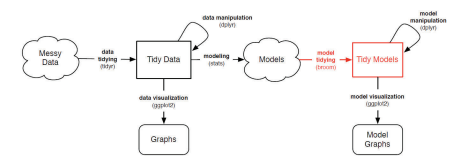


Image from: https://opr.princeton.edu/workshops/Downloads/2016Jan BroomRobinson.pdf



Example: Penguin data

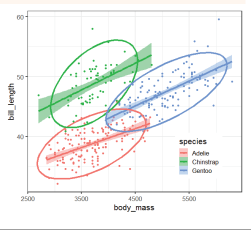
Say we want to model the length of penguin bills

Potential predictors: body mass, species, sex, island, ...

```
ggplot(data=peng, aes(x=body mass, y=bill length, color=species, fill=species)) +
geom point() +
geom smooth(method = "lm", size=1.5) +
stat ellipse(size=1.5)
```

How to extract & plot model statistics?

How to fit multiple models for subsets (by: sex, island)?



Predict bill length from body mass, species, sex, island

```
peng.mod0 <- Im(bill length ~ body mass + species + sex + island, data=peng)
summary(peng.mod0)
```

```
lm(formula = bill_length ~ body_mass + sex + species + island, data = peng)
Residuals:
                                                             observation level
   Min
          1Q Median
                        3Q
-6.938 -1.473 0.002 1.257 10.490
Coefficients:
                                                             component level
                 Estimate Std. Error t value Pr(>|t|)
(Intercept)
                31.237676
                          1.378201 22.67 < 2e-16 ***
                                                             (coefficients)
                 0.001744
                            0.000394
                                       4.43 1.3e-05 ***
body_mass
                 2.532799
                                       7.01 1.4e-11 ***
                            0.361396
speciesChinstrap 10.281510
                            0.410391
                                      25.05 < 2e-16 ***
speciesGentoo
                 6.149100
                            0.672481
                                       9.14
                                             < 2e-16 ***
islandDream
                 -0.463555
                            0.457443
                                       -1.01
                                                0.31
islandTorgersen 0.092246
                            0.474401
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 2.26 on 326 degrees of freedom
                                                             model level
Multiple R-squared: 0.832, Adjusted R-squared: 0.829
F-statistic: 269 on 6 and 326 DF, p-value: <2e-16
```

glance() gives the model level summary statistics

```
> glance(peng.mod0)
 r.squared adj.r.squared sigma statistic p.value df logLik AIC BIC deviance df.residual
            0.829 2.26 269. 4.60e-123 6 -741. 1497. 1528. 1667. 326
```

tidy() gives the model component (term) statistics

```
> tidy(peng.mod0)
 term
                 estimate std.error statistic p.value
1 (Intercept)
                  31.2
                          1.38
                                      22.7 5.90e-69
                                     4.43 1.29e- 5
                  0.00174 0.000394
2 body mass
                  2.53
                                      7.01 1.39e-11
4 speciesChinstrap 10.3
                          0.410
                                      25.1 5.61e-78
5 speciesGentoo
                 6.15
                          0.672
                                      9.14 6.65e-18
6 islandDream
                  -0.464
                          0.457
                                      -1.01 3.12e- 1
7 islandTorgersen 0.0922
```

augment() gives the observation level statistics

```
> augment(peng.mod0) %>% sample n(5)
 bill length body mass sex species
                                     island .fitted .resid .hat .sigma
                 <int> <fct> <fct>
                                     <fct>
                                              <dbl> <dbl> <dbl> <dbl> <dbl>
                                                                           <db1>
                                                                                      <db1>
       <dbl>
        35.7
                 3150 f
                          Adelie
                                     Biscoe
                                               36.7 -1.03 0.0273
                                                                  2.26 0.000858
                                                                                      -0.463
        39.7
                 3550 m
                                               40.0 -0.263 0.0331
                                                                  2.26 0.0000684
                                                                                     -0.118
                            Adelie
                                     Biscoe
                 4050 m
        41.1
                            Adelie
                                     Biscoe
                                               40.8 0.265 0.0257
                                                                  2.26 0.0000532
                                                                                      0.119
        49.3
                  4050 m
                            Chinstrap Dream
                                               50.7 -1.35 0.0177
                                                                   2.26 0.000939
                                                                                     -0.604
        48.4
                 5850 m
                            Gentoo
                                     Biscoe
                                               50.1 -1.72 0.0169
                                                                  2.26 0.00145
                                                                                     -0.769
```

Fitting multiple models

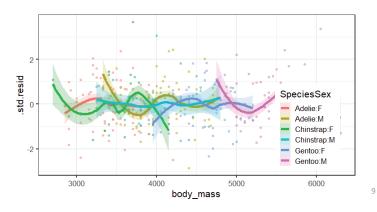
With dplyr::nest_by(), you can fit one model for each subset of the data Feed this to broom::alance() to get model-level statistics

```
models <- peng |>
  nest_by(species, sex) |>
  mutate(model = list(lm(bill_length ~ body_mass, data = data)))
  summarise(broom::glance(model), .groups = "keep") |>
  select(species, sex, r.squared, sigma, statistic, p.value, nobs)
```

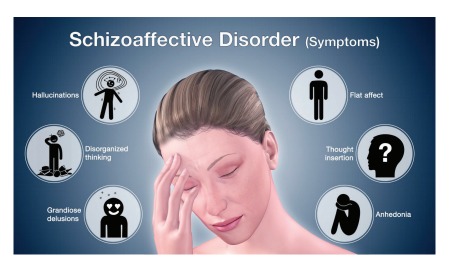
```
# A tibble: 6 x 7
# Groups:
           species, sex [6]
 species
           sex
                 r.squared sigma statistic p.value nobs
  <fct>
                     <db1> <db1>
                                     <dbl> <dbl> <int>
1 Adelie
                    0.0289 2.01
                                      2.12 0.150
                                                      73
2 Adelie
                    0.0486 2.24
                                      3.62 0.0610
                                                      73
                    0.0760 3.03
                                      2.63 0.115
3 Chinstrap f
                                                      34
4 Chinstrap m
                    0.0568
                           1.54
                                      1.93 0.175
                                                      34
5 Gentoo
                    0.0739 1.99
                                      4.47 0.0389
                                                      58
6 Gentoo
                    0.153
                            2.52
                                     10.7 0.00182
```

Plotting via broom::augment

```
models |>
summarise(broom::augment(model), .groups = "keep") |>
mutate(
sex = stringr::str_to_upper(sex),
SpeciesSex = forcats::fct_cross(species, sex)) |> # Combine species and sex
ggplot(aes(x=body_mass, y=.std.res, color=SpeciesSex, fill=SpeciesSex)) +
geom_point(alpha=0.4) +
geom_smooth(method = "Im", size = 2, alpha=0.3)
```



Schizophrenia symptoms: Hallucinations, disorganized thinking, delusions, ... Schizoaffective disorder combines symptoms of schizophrenia with mood disorder (bipolar or depression)



Ex: Neuro- & Social-Cognitive measures in psychiatric groups

- A study by Leah Hartman @York examined whether patients classified as 'schizophrenic' or 'schizoaffective' (on DSM-IV) could be distinguished from a normal, control sample on standardized tests in the following domains:
 - Neuro-Cognitive: processing speed, attention, verbal learning, visual learning, problem solving
 - Social-cognitive: managing emotions, theory of mind, externalizing bias, personalizing bias
- Research questions → MANOVA contrasts
 - Analyze neuro-cog (NC) and social-cog (SC) separately
 - Do the two psychiatric groups differ from the controls?
 - Do the psychiatric groups differ from each other?

See: Friendly & Sigal (2017), Graphical Methods for Multivariate Linear Models in Psychological Research: An R Tutorial *The Quantitative Methods for Psychology, 13*, 20-45, http://dx.doi.org/10.20982/tqmp.13.1.p020

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Neuro-cognitive measures

Data

<pre>> data(NeuroCog, package="heplots")</pre>											
<pre>> car::some(NeuroCog)</pre>											
	Dx	Speed	Attention	Memory	Verbal	Visual	ProbSolv	SocialCog	Age	Sex	
14	Schizophrenia	19	9	19	33	24	39	28	44	Female	
88	Schizoaffective	27	44	24	33	26	31	36	53	Female	
105	Schizoaffective	23	41	42	48	46	29	50	52	Female	
114	Schizoaffective	41	53	47	39	30	50	63	32	Female	
130	Control	44	25	21	37	32	43	29	43	Female	
165	Control	35	35	43	53	57	37	40	30	Male	
194	Control	47	40	53	53	40	45	49	55	Female	

diagnostic group ------ neuro-cognitive measures ------ ignored -----

Questions:

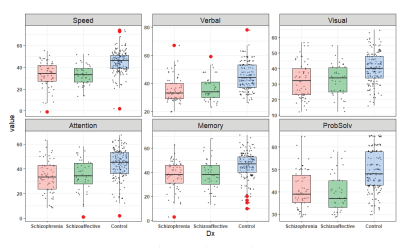
- Do the diagnostic groups differ collectively on the neuro-cognitive measures?
- How do group differences relate to research hypotheses?
- How many dimensions (aspects) are reflected in the differences among means?

NeuroCog: EDA

Always plot the data first!

With boxplots, often useful to show more: jittered points show the data

What can we see here?



Dx = Schizophrenia = Schizoaffective = Control

How did I make this plot?

- 1. Reshape the data wide \rightarrow long
- 2. ggplot() + geom_jitter() + geom_boxplot() + facet_wrap(~response)

```
NC_long <- NeuroCog |>
   select(-SocialCog, -Age, -Sex) |>
   tidyr::gather(key = response, value = "value", Speed:ProbSolv)
```

```
# show a few observations
library(dplyr)
NC_long |>
    group_by(Dx) |>
    sample_n(4) |>
    ungroup()
```

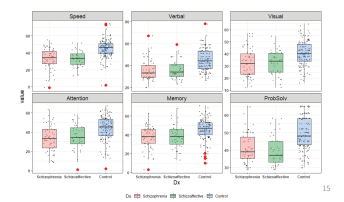
Dx	response	value
1 Schizophrenia 2 Schizophrenia 3 Schizophrenia 4 Schizophrenia 5 Schizoaffective 6 Schizoaffective 7 Schizoaffective 8 Schizoaffective 9 Control 10 Control 11 Control 12 Control	Visual	37 49 61 23 11 39 41 54 45 45 45 61

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How did I make this plot?

- 1. Reshape the data wide \rightarrow long
- 2. ggplot() + geom jitter() + geom boxplot() + facet wrap(~response)

```
ggplot(NC_long, aes(x=Dx, y=value, fill=Dx)) +
  geom_jitter() +
  geom_boxplot() +
  facet_wrap(~response, scales = "free_y") +
  theme_bw()
```



Neuro-cognitive measures: Analyses

Could do a series of univariate ANOVAs

```
car::Anova(NC.mlm1 <- lm (Speed ~ DX, data=NeuroCog))
car::Anova(NC.mlm2 <- lm (Attention ~ DX, data=NeuroCog))
car::Anova(NC.mlm3 <- lm (Memory ~ DX, data=NeuroCog))</pre>
```

Better: MANOVA test for all 6 responses together

Anova(NC.mlm = lm(cbind(Speed, Attention, Memory, ...) ~ Dx, data=NeuroCog)

Research hypotheses: test contrasts

```
    Dx1: Control – (Schiz + SchizAff)/2 = 0 → linearHypotheses (NC.mlm, "Dx1")
    Dx2: Schiz – SchAff = 0 → linearHypotheses (NC.mlm, "Dx2")
```

Dimensions:

- Canonical analysis → NC.can <- candisc(NC.mlm); plot(NC.can)
 Canonical HE plot → heplot(NC.can)
- Tech note: anova() in base R vs. car::Anova
 - anova() uses only Type 1 (sequential) tests, rarely useful; doesn't handle MLM well
 - car::Anova() provides Type 2, 3 (partial) tests; give sensible results for MLMs
 - car::linearHypotheses() gives univariate and multivariate tests of contrasts

Tidy univariate analyses: *F*-tests

Using broom::tidy() we can fit a separate univariate ANOVA model for each response I've used some tidyverse magic to make a nicely formatted table of F values

```
mutate(response = factor(response, levels=unique(response))) |> # keep variable order
group_by(response) |>
do(tidy(anova(lm(value ~ Dx, .)))) |>
                                                           # run anova on each submodel
filter(term != "Residuals") |>
select(-term) |>
rename (F = statistic, df1 = df,
                                                              # use more specific names
      SS = sumsq, MS =meansq) |>
mutate(df2 = 239) |>
                                                     # kludge: extract dfe from object?
relocate(df2, .after = df1) |>
mutate(signif = noquote(gtools::stars.pval(p.value))) |>
                                                                # add significance stars
mutate(p.value = noquote(scales::pvalue(p.value)))
                                                                # format p.values nicely
```

```
df1 df2 SS MS
                                 F p.value
response
 <fct>
          <int> <dbl> <dbl> <dbl> <noquote> <noquote>
       2 239 8360. 4180. 37.1 < 0.001
2 Attention 2 239 5579. 2790. 17.4 <0.001
                                                       All variables
             2 239 3764. 1882. 13.9 < 0.001
3 Memory
                                                       are signif.
4 Verbal
             2 239 4672. 2336. 27.3 < 0.001
             2 239 3692. 1846. 16.6 < 0.001
5 Visual
             2 239 4165. 2083. 25.1 < 0.001
6 ProbSolv
```

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Tidy univariate analyses: *t*-tests

Can do the same to extract the t-tests for coefficients or contrasts in the univariate models for each response

```
NC long |>
 mutate(response = factor(response, levels=unique(response))) |>
 group by (response) |>
  do(tidy(lm(value ~ Dx, .))) |>
 filter(term != "(Intercept)") |>
 mutate(signif = noquote(gtools::stars.pval(p.value))) |>
 mutate(p.value = noquote(scales::pvalue(p.value)))
```

```
response term estimate std.error statistic p.value
  <fct>
         <chr> <dbl>
                        <dbl>
                                <dbl> <noquote> <noquote>
1 Speed
         Dx1
                7.99
                         0.940 8.51 < 0.001
2 Speed Dx2
               -0.0252
                        1.10 -0.0229 0.982
3 Attention Dx1 6.44
                        1.12 5.75 <0.001
4 Attention Dx2 -0.558
                        1.31 -0.426 0.671
5 Memory Dx1 5.31
                        1.03 5.16 < 0.001
         Dx2 -0.375
                        1.20 -0.311 0.756
6 Memory
               5.91
7 Verbal
         Dx1
                         0.818 7.22 < 0.001
                         0.957 -0.461 0.645
8 Verbal
         Dx2
9 Visual
         Dx1
                5.31
                         0.933
                               5.69 < 0.001
10 Visual Dx2
               -0.0371
                         1.09
                              -0.0340 0.973
11 ProbSolv Dx1
                5.70
                         0.807 7.07 < 0.001
12 ProbSolv Dx2
                              0.762 0.447
```

Neuro-cognitive measures: MANOVA

Multivariate tests are more straight-forward & comprehensive

```
# fit the MANOVA model
NC. mlm <- lm(cbind(Speed, Attention, Memory, Verbal, Visual, ProbSoly) ~ Dx,
            dat a=Neur oCog)
Anova (NC. ml m)
Type II MANOVA Tests: Pillai test statistic
  Df test stat approx F num Df den Df Pr(>F)
                         12 470 1.562e-11 *** V
Dx 2 0.2992 6.8902
Signif. codes: 0 å***ç 0.001 å**ç 0.01 å*ç 0.05 å.ç 0.1 å ç 1
```

```
> contrasts (NeuroCog$Dx)
So, the groups differ. But how?
What about the research hypotheses? -
                                              ➤ Schizophrenia -0.5
                                                 Schizoaffective - 0.5
                                                                   1.0
                                                 Cont r ol
                  Dx1: Control – (Schiz + SchizAff)/2 = 0
```

Neuro-cognitive measures: Contrasts

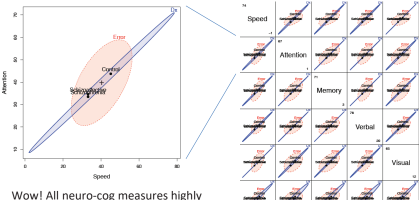
A simple result: Control \neq (Schizophrenia \approx Schizoaffective)

```
> print(car::linearHypothesis(NC.mlm, "Dx1"), SSP=FALSE)
Multivariate Tests:
                Df test stat approx F num Df den Df Pr(>F)
                                                234 2.8e-15 ***
Pillai
                 1
                       0.289
                                 15.9
Wilks
                 1
                       0.711
                                 15.9
                                                234 2.8e-15 ***
                       0.407
                                 15.9
                                                234 2.8e-15 ***
Hotelling-Lawley 1
                       0.407
                                 15.9
                                                234 2.8e-15 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
> print(car::linearHypothesis(NC.mlm, "Dx2"), SSP=FALSE)
Multivariate Tests:
                Df test stat approx F num Df den Df Pr(>F)
Pillai
                       0.006
                                0.249
                                                234
                                                      0.96
Wilks
                       0.994
                                0.249
                                           6
                                                234
                                                      0.96
                       0.006
                                0.249
                                               234
Hotelling-Lawley 1
                                                     0.96
                       0.006
                                0.249
                                                234
                                                      0.96
```

Visualize me: in data space

Bivariate view for any 2 responses: heplot (NC. ml m, var = 1: 2, ...)

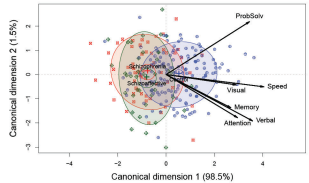
HE plot matrix: for all responses pairs (NC. ml m, ...)



Wow! All neuro-cog measures highly correlated in group means! Only 1 dim. of H variation

Visualize me: in canonical space

- As with biplot, we can visualize MLM hypothesis variation for all responses by projecting **H** and **E** into low-rank space.
- Canonical projection: $\mathbf{Y}_{n \times p} \mapsto \mathbf{Z}_{n \times s} = \mathbf{Y} \mathbf{E}^{-1/2} \mathbf{V}$, where \mathbf{V} = eigenvectors
- This is the view that maximally discriminates among groups, ie max. H



Very simple interpretation

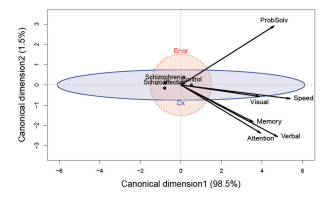
Can1: normal vs. others All vars highly + correlated;

Can2: only 1.5%, NS; but perhaps suggestive (ProbSolv vs. Attention)

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Visualize me: canonical HE plots

- Canonical HE plot is just the HE plot of canonical scores, (z₁, z₂) in 2D,
- or, z₁, z₂, z₃, in 3D.
- As in biplot, we add vectors to show relations of the \mathbf{y}_i response variables to the canonical variates.
- variable vectors here are structure coefficients = correlations of variables with canonical scores.



The multivariate "juicer"

Shows just group means, H ellipse & E ellipse

Variable vectors offer interpretation of Can dimensions.

Social cognitive measures

- These measures deal with the person's perception and cognitive processing of emotions of others
 - Scales: managing emotions, theory of mind, externalizing bias, personalizing bias
- Questions:
 - Do these differentiate normal from patient groups?
 - Can they distinguish between schizophrenic & schizoaffective
 - If so, this could be a major finding.

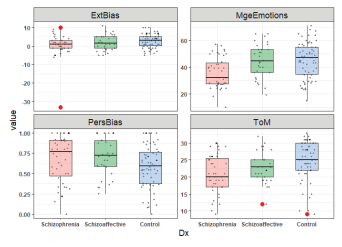
SocialCog: EDA

Again, it is helpful to plot the data!

Same steps: (a) Reshape to long; (b) ggplot() + ... + facet wrap(~response)

What can we see here?

How do groups differ? Unusual features?



Dx = Schizophrenia = Schizoaffective = Control

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Social cognitive measures

Data

```
> car::some(SocialCog)
                Dx MgeEmotions ToM ExtBias PersBias
                            32 18
                                               1.00
      Schizophrenia
58
     Schizophrenia
                            32 17
                                               0.92
      Schizophrenia
                            52 23
                                               0.80
85 Schizoaffective
                            28 12
                                               1.00
99 Schizoaffective
                            40 22
                                               0.36
111 Schizoaffective
                            51 23
                                               0.91
           Control
                            51 24
                                               1.00
155
           Control
                            57 28
                                               0.30
168
           Control
                            47 32
                                               0.77
216
           Control
                            64 31
                                               0.69
```

diagnostic group ------ social-cognitive measures -----

Analyses

Standard MANOVA test:

Anova(SC.mlm = lm(cbind(...) ~ DX, data=SocialCog)

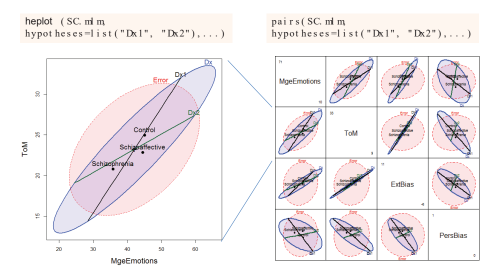
Test contrasts:

linearHypotheses(SC.mlm, "Dx1")
linearHypotheses(SC.mlm, "Dx2")

Social cognitive measures

Test contrasts: Dx1 = Normal vs. Patient; Dx2 = Schizo vs. Schizoaffective

Visualize me: data space



The groups are ordered Schizo < ScAffective < Control on these measures

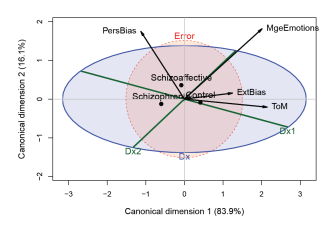
One of these vars is not like the others!

Visualize me: canonical space

Contrasts:

Dx1 : Control vs. patients. Controls > patients on MgeEmotions, ExtBias, ToM

Dx2: Schizo vs. schizoaffective.



Can1: group order

Can2: Schizoaffective vs. others

Dx1 suggests ≈ spacing Schizo < ScAff < Control

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Model checking & remedies

- The MLM assumes residuals are multivariate normal
 - → Squared Mahalanobis distances

$$D_{M}^{2}(\mathbf{y}_{i}) = (\mathbf{y}_{i} - \overline{\mathbf{y}})^{T} \mathbf{S}^{-1}(\mathbf{y}_{i} - \overline{\mathbf{y}}) \sim \chi_{p}^{2} \text{ with p d.f.}$$

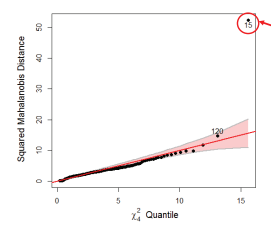
- → a quantile quantile plot of ordered $D_M^2(\mathbf{y}_i)$ vs. quantiles of χ_p^2 should plot as straight line
- Outliers are readily apparent
- plots: heplots::cqplot()
- Influence plots
 - mvinfluence::influence() calculates multivariate analogs of influence measures

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Social cog: cqplot

heplots::cqplot() creates a chi-square QQ plot from a MLM

cqplot(SC.mlm, id.n=2)



One observation appears as an extreme outlier.

This was a case w/ ExtBias = -33, but valid range = (-10, +10)

Refitting w/o case 15: Overall & DX1 tests still OK ☺ Dx2 test: p=0.074, now NS ☺

Social cog: Influence

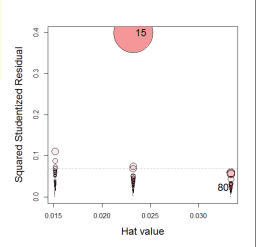
mvinfluence::influencePlot() creates a multivariate analog of an influence plot

>library(mvinfluence)
>influencePlot(SC.mlm, ...)

H Q COOKD L R 15 0.0233 0.3985 0.4202 0.0238 0.4080 80 0.0333 0.0233 0.0352 0.0345 0.0241

Case 15 stands out as hugely influential (a data error)

The 3 columns of circles correspond to the 3 groups



Robust MLMs

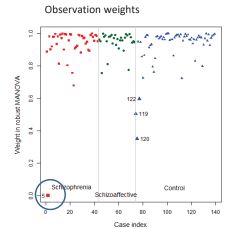
- Robust methods for univariate LMs are now well-developed and implemented
 - → proper SEs, CIs and hypothesis tests
- Analogous methods for multivariate LMs are a current hot research topic
- The heplots package now provides r obml m() for the fully general MLM (MANOVA, MMReg)
 - Uses simple M-estimator via IRLS
 - Weights: calculated from Mahalanobis D^{2,} a robust covariance estimator and weight function, ψ(D²)

$$D^{2} = (\mathbf{Y} - \overline{\mathbf{Y}})^{\mathsf{T}} \mathbf{S}_{\mathsf{robust}}^{-1} (\mathbf{Y} - \overline{\mathbf{Y}}) \sim \chi_{p}^{2}$$

Downside: SEs, p-values only approximate

Robust MLMs

SC.rlm <- robmlm(cbind(MgeEmotions, ToM, ExtBias, PersBias) ~ Dx, data=SocialCog)



Approx test of Dx2 in robust model

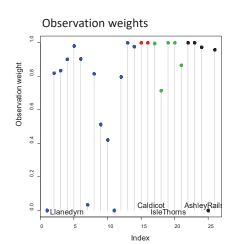
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Robust MLMs: Pottery data

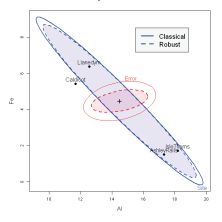
- Chemical composition of ancient pottery found at four sites in Great Britain
 - Sites: AshleyRails, Caldicot, IsleThorns, Llanedyrn
 - Measures: Al Fe Mg Ca Na
- Analyses:
 - standard MANOVA Im(cbind(Al, Fe, Mg, ...) ~ Site, data=Pottery)
 - robust MANOVA robmlm(cbind(Al, Fe, Mg, ...) ~ Site, data=Pottery)
- Data:

Robust MLMs: Pottery data

> pottery.mod < - Im(cbind (Al,Fe,Mg,Ca,Na)~Site, data=Pottery)
> pottery.rmod <- robmlm (cbind (Al,Fe,Mg,Ca,Na)~Site, data=Pottery)



residual E ellipse shrinks a lot overlaid HE plots



3

MMRA example: PA tasks & ability

- Rohwer data from Timm (1975)
- How well do paired associate (PA) tasks predict performance on measures of aptitude & achievement in kindergarten children?
 - Samples: 69 children in two groups (schools): 'Lo' | 'Hi' SES
 - Outcomes (Y):
 - Scholastic aptitude test (SAT)
 - Peabody picture vocabulary test (PPVT)
 - · Raven progressive matrices (Raven)
 - Predictors (X): Scores (0—40) on PA tasks where the stimuli were:
 - named (n), still (s), named-still (ns), named-action (na), sentence-still (ss)

```
> data("Rohwer", package="heplots")
> car::some(Rohwer, n=5)
  group SES SAT PPVT Raven n s ns na ss
                      8 0 0 10 19 14
           8 68
                      11 0 0 7 16 13
      1 Lo 49 74
      1 Lo 19 66
17
                      13 7 12 21 35 27
           38
                66
                      14 0 0 3 16 11
            8
                55
                      16 4 7 19 20 13
```

Having a group factor makes the analysis more complicated (MANCOVA)

Start with analysis of the Hi SES group

> Rohwer2 <- subset(Rohwer, subset=SES=="HI")</pre>

S

Why not univariate models?

```
rohwer.mod1 <- lm(SAT \sim n + s + ns + na + ss, data = Rohwer2)
rohwer.mod2 <- lm(PPVT \sim n + s + ns + na + ss, data = Rohwer2)
rohwer.mod3 <- lm(Raven \sim n + s + ns + na + ss, data = Rohwer2)
library(stargazer)
stargazer(rohwer.mod1, rohwer.mod2, rohwer.mod3, type="text", ...)
```

Univariate regression models for Rohwer data									
	Dependent variable:								
	SAT	PPVT	Raven						
n s ns na ss	3.26* 3.00 -5.86*** 5.67*** -0.62	0.07 0.37 -0.37 1.52* 0.41	0.06 0.49** -0.16 0.12 -0.12						
R2 F Statistic (df = 5; 26)	0.56 6.54***	0.35	0.31 2.32						
Note:	*p<0.05;	**p<0.01;	***p<0.001						

Results are disappointing

- Only model for SAT highly signif.
- Only a few coefs. signif.
 ≠0

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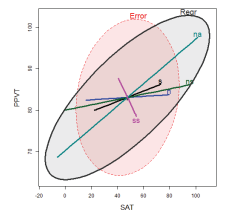
MANOVA tests

```
> rohwer.mlm <- lm(cbind(SAT, PPVT, Raven) ~ n + s + ns + na + ss, data=Rohwer2)
> Anova(rohwer.mlm)
                                                                  Much better!
Type II MANOVA Tests: Pillai test statistic
   Df test stat approx F num Df den Df Pr(>F)
          0.202
                  2.02
                                    24 0.1376
                                                                  Multivariate tests:
                   3.59
                                    24 0.0284 *
          0.310
S
                                                                     pool evidence for all Ys
                                    24 0.0126 *
   1
          0.358
                   4.46
                                                                    take correlations of Ys
                                    24 0.0016 **
   1
          0.465
                   6.96
ss 1
          0.089
                   0.78
                                    24 0.5173
                                                                      into account
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Can also test overall hypothesis, H_0 : **B** = **0** (all coefs = 0)

```
> print(linearHypothesis(rohwer.mlm,
                         c("n", "s", "ns", "na", "ss")), SSP=FALSE)
Multivariate Tests:
                 Df test stat approx F num Df den Df
                                                                       Strongly reject H<sub>0</sub>
Pillai
                     1.0386
                                         15 78.00 0.001912 **
                               2.753
                                                                       by all criteria
Wilks
                       0.2431
                                 2.974
                                          15 66.65 0.001154 **
Hotelling-Lawley 5
                      2.0615
                                 3.115
                                         15 68.00 0.000697 ***
                                           5 26.00 0.000160 ***
                       1.4654
                                7.620
```

Visualize me!



Each predictor gives a 1 df test -> \mathbf{H}_{i} ellipse is a line

E here is a 3D ellipsoid (rank(E) = min(p,q))

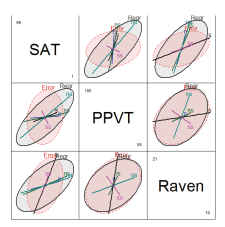
Interpretation:

- Any H ellipse that protrudes outside E ellipse is significant by Roy's test
- Length of each H_i line ≈ strength of association
- Orientation of each H_i line shows relation of X_i to the two Y_i responses shown.

pairs.mlm() plot

View all pairwise HE plots

pairs(rohwer.mlm, hypotheses=hyp, col=cols, fill=TRUE, fill.alpha=0.1, ...)



Can now understand more subtle aspects

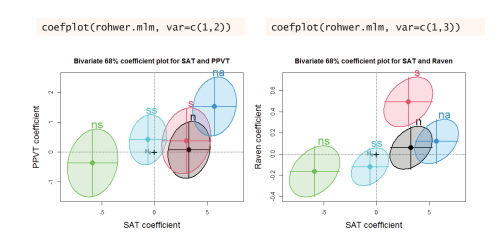
SAT is best predicted overall, but relation with PA tests varies
The na & ns tasks are strongest for SAT

Raven is weakly predicted

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Coefficient plots for MLMs

In multivariate models, confidence intervals for coefficients become ellipsoids Two coefficients are jointly significant if their conf. ellipse $\mathbb{E}(b_1, b_2)$ excludes (0,0) These plots are provided by heplots::coefplot(mlm)



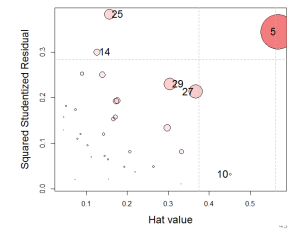
42

Influence plots

As always, we should check for influential observations and multivariate outliers

mvinfluence::influencePlot(Rohwer.mlm, id.n=4, ...)

One point (5) deserves further attention

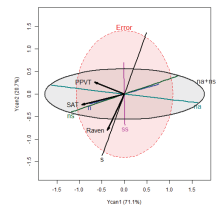


Canonical correlations

For quantitative (X, Y) data, canonical correlation analysis is an alternative to MMRA It finds the weighted sums of the Y variables most highly correlated with the Xs

```
> X <- Rohwer2[, 6:10] # X variables for High SES students
> Y <- Rohwer2[, 3:5] # Y variables for High SES students
> (cc <- cancor(X, Y, set.names=c("PA", "Ability")))</pre>
Canonical correlation analysis of:
           5 PA variables: n, s, ns, na, ss
              Ability variables: SAT, PPVT, Raven
    CanR CanRSQ Eigen percent
                                 cum
                                                                      Two dimensions
1 0.7710 0.5944 1.4654 71.080
2 0.5465 0.2987 0.4259 20.659 91.74
                                                                      acct for 91.7% of
                       8.261 100.00
3 0.3815 0.1455 0.1703
                                                                      (X,Y) association
Test of HO: The canonical correlations in the
current row and all that follow are zero
    CanR LR test stat approx F numDF denDF Pr(> F)
                                                                      Only Can1 is
                       2.9738
                                 15 66.655 0.001154 **
1 0.77096
              0.24307
                                                                      significant
2 0.54652
              0.59926
                       1.8237
                                   8 50.000 0.094538
3 0.38147
              0.85448
                        1.4759
                                   3 26.000 0.244178
```

Visualize CCA in HE plot



Residuals are uncorrelated in canonical space

H ellipses for X terms same as in ordinary HE plots – outside **E** ellipse *iff* signif. by Roy's test

Variable vectors for Ys: correlations with canonical variables Ycan1, Ycan2

- SAT & PPVT: mainly Ycan1
- Raven: more aligned with Ycan2

MANCOVA & homogeneity of regression

- With a group variable (SES) can test differences in means (intercepts)
 - rohwer.mod <- Im(cbind(SAT, PPVT, Raven) ~ SES + n + s + ns + na + ss, data=Rohwer)
 - This assumes that slopes (B) are the same for both groups (homogeneity of regression)
- Can test for equal slopes by adding interactions of SES with Xs
 - rohwer.mod1 <- Im(cbind(SAT, PPVT, Raven) ~ SES * (n + s + ns + na + ss))</p>
- Or, fit separate models for each group

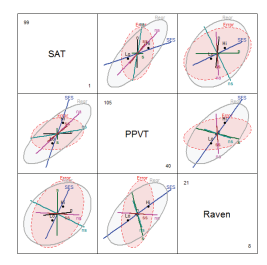
45

MANCOVA

Fit the MANCOVA model & test hypotheses

```
> rohwer.mod <- lm(cbind(SAT, PPVT, Raven) ~ SES + n + s + ns + na + ss,
                   data=Rohwer)
> Anova(rohwer.mod)
Type II MANOVA Tests: Pillai test statistic
    Df test stat approx F num Df den Df Pr(>F)
                                                         SES: diff in means
                                     60 2.5e-06 ***
           0.379
                   12.18
    1
           0.040
                     0.84
                               3
                                     60 0.4773
           0.093
                     2.04
    1
                                     60 0.1173
           0.193
                     4.78
                                     60 0.0047 **
    1
           0.231
                     6.02
                                     60 0.0012 **
                     1.05
                                     60 0.3770
           0.050
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Visualize effects



The SES effect is positive for all Y variables
Hi SES group > Lo SES group

. .

Fit model with interactions

Fit heterogeneous regression model with SES interactions

```
> rohwer.mod1 <- lm(cbind(SAT, PPVT, Raven) \sim SES * (n + s + ns + na + ss),
                     data=Rohwer)
> Anova(rohwer.mod1)
Type II MANOVA Tests: Pillai test statistic
       Df test stat approx F num Df den Df Pr(>F)
              0.391
                       11.78
                                         55 4.5e-06
              0.079
                        1.57
                                         55 0.20638
n
        1
              0.125
                        2.62
                                         55 0.05952 .
                                                              OK, as expected
                                         55 0.00100 ***
              0.254
                        6.25
na
              0.307
                                         55 0.00015 ***
                        1.17
SS
              0.060
                                         55 0.32813
              0.072
                        1.43
                                         55 0.24417
SES:n
SES:s
              0.099
                        2.02
                                         55 0.12117
                        2.44
                                         55 0.07383
                                                              Hmm ???
              0.118
SES:ns 1
SES:na 1
              0.148
                        3.18
                                         55 0.03081
                                         55 0.35094
SES:ss 1
              0.057
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Test interactions

Can test all interactions simultaneously with linearHypothesis() Do I need any interaction terms?

I use a 'grep' trick here to find the names of coefficients like 'SES:' containing a ':'

```
> coefs <- rownames(coef(rohwer.mod1))</pre>
                                        # store coefficient names in a vector
> print(linearHypothesis(rohwer.mod1,
                                        # only test for interaction effects
                         coefs[grep(":", coefs)]), SSP=FALSE)
Multivariate Tests:
                Df test stat approx F num Df den Df Pr(>F)
Pillai
                      0.4179
                               1.845
                                          15 171.0 0.03209 *
Wilks
                                1.894
                                          15 152.2 0.02769 *
                      0.6236
Hotelling-Lawley 5
                      0.5387
                                1.927
                                          15 161.0 0.02396 *
Roy
                      0.3846
                                4.385
                                           5 57.0 0.00191 **
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

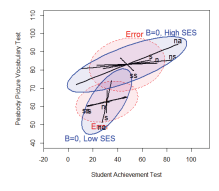
Evidence shows that some slopes differ for Hi/Lo SES

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Fit separate models

Fitting a model for each group allows all slopes to differ Also allows within-group covariances to differ

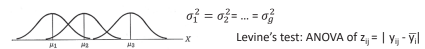


For SAT & PPVT:

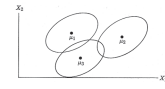
- means higher for Hi SES
- within-group covariance larger for Hi SES
- slopes of predictors smaller for Hi SES
 → SAT more important for this group.

Homogeneity of (co)variances

ANOVA assumes equality of residual variances



MANOVA: assumes equality of covariance matrices



$$\Sigma_1 \texttt{=} \; \Sigma_2 \texttt{=} \; ... \texttt{=} \; \Sigma_g$$

Box's M test: $M = (N-g) \ln |\mathbf{S}_P| - \sum_{i=1}^g (n_i - 1) \ln |\mathbf{S}_i|$

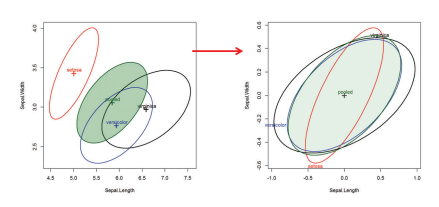
$$-2 \ln(M)^{\sim} \chi^{2}_{df}$$

See: http://www.datavis.ca/papers/EgCov-TAS.pdf

Visualizing covariance matrices

Visualize covariance ellipses in data space Center to see pure differences in size & shape

```
covEllipses(iris[,1:2], iris$Species, ...)
covEllipses(iris[,1:2], iris$Species, center=TRUE, ...)
```

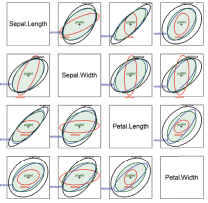


View all pairs

covEllipses(iris[,1:4], iris\$Species, center=TRUE, variables=1:4, ...)

In all cases, setosa stands out as different from the others

- sometimes correlation differs
- sometimes smaller variance(s)



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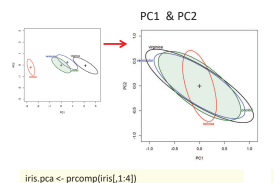
56

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Visualize in PCA space

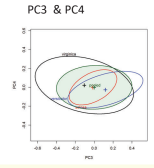
PCA projects the data into an orthogonal space accounting for maximum variance Covariance ellipses show the differences among groups in this space

Surprisingly, the small dimensions contribute largely to Box's M test.



covEllipses(iris.pca\$x, iris\$Species, center=TRUE, ...)

covEllipses(iris.pca\$x, iris\$Species, ...)

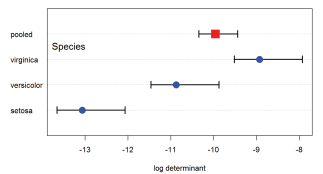


covEllipses(iris.pca\$x, iris\$Species, center=TRUE, variables=3:4, ...)

Visualizing Box's M test

Box's test is based on a comparison of the log $|S_i|$ relative to log $|S_p|$: plot them!

iris.boxm <- boxM(iris[, 1:4], iris[, "Species"])
plot(iris.boxm, gplabel="Species")</pre>

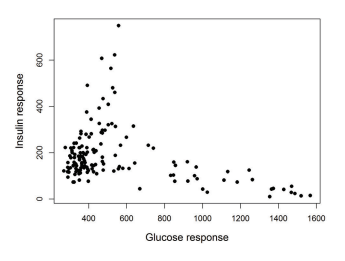


CIs based on an asymptotic CLT ≈ distribution of In |S| (Cai, Liang, and Zhou 2016) (Thx: Augustine Wong)

Unsolved: Bootstrap CI

Diabetes data: 2D mystery

Reaven & Miller (1968) found a peculiar "horse shoe" result in analysis of data on the relationship of blood glucose levels and production of insulin in patients with varying degrees of hyperglycemia



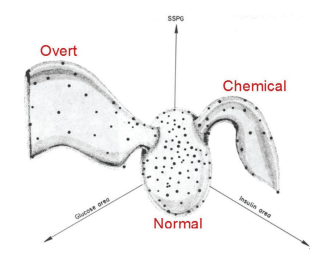
In a 2D plot this was a medical mystery.

What could be the explanation?

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Diabetes data: 3D clarity

Using the first 3D computer graphics system (PRIM-9) they rotated the data in 3-space until a hypothesis was suggested.



Artist's view of the data suggests there were actually three groups in the data.

Two categories of Type 2 diabetes:

- · Overt (advanced)
- Chemical (latent)

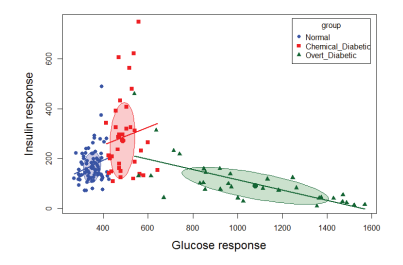
But, these were NOT stages in a progression!

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Diabetes data: scatterplots

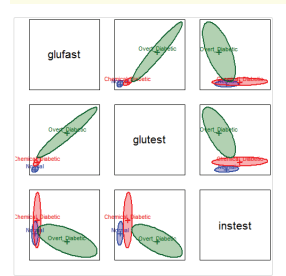
A grouped scatterplot reveals the truer story

scatterplot(instest ~ glutest | group, data=Diabetes, ellipse=TRUE, ...)



Diabetes data: Ellipses

covellipses(Diabetes[,2:5], Diabetes\$group, fill=TRUE, pooled=FALSE, col=c("blue", "red", "darkgreen"), variables=1:3)



The pairwise data ellipses show visual summaries of the data

The differences in correlation and variances are dramatic here

Box's M test

```
diab.boxm <- boxM(Diabetes[,2:5], Diabetes$group)
diab.boxm

##

## Box's M-test for Homogeneity of Covariance Matrices

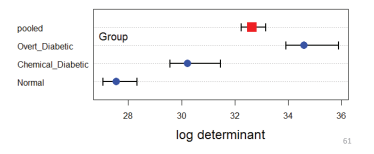
##

## data: Diabetes[, 2:5]

## Chi-Sq (approx.) = 383, df = 20, p-value <2e-16

plot(diab.boxm)
```

All pairs differ!



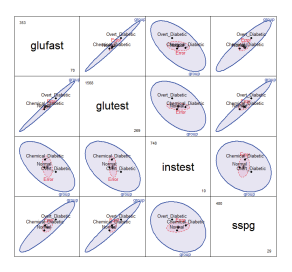
MANOVA & HE plots

diab.mlm <- Im(cbind(glufast, glutest, instest, sspg) ~ group, data=Diabetes) # HE pairs plot pairs(diab.mlm, fill=TRUE, fill.alpha=0.1, var.cex=3, cex=1.25)

The groups are wellseparated in all views

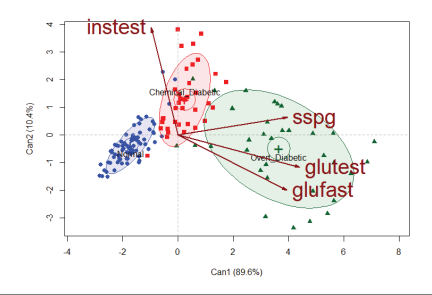
Mostly positive correlation among group means

But: one of these plots is not like the others!



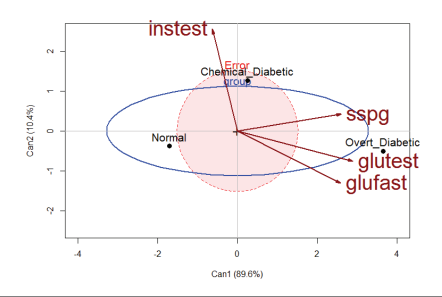
Run the juicer

diab.can <- candisc(diab.mlm)
plot(diab.can, ellipse=TRUE, var.lwd=2, ...)



Filter the juice

heplot(diab.can, fill=c(TRUE, FALSE), fill.alpha=0.1, var.lwd=2, size="effect")



Summary

- MANOVA tests of MLMs are easily visualized in HE plots
 - Contrasts among groups can be easily shown
 - Canonical plots show data in 2D/3D space of max. group differences
 - Robust methods can help guard against outliers
- MMRA models
 - Visualize effects of quant. predictors as lines in data space
 - Test & visualize any linear hypothesis
 - Canonical correlations: visualize in 2D/3D of max. (X, Y) correlations
- Homogeneity of covariances
 - Visualize within-group S_i and pooled S_D by data ellipses
 - Visualize Box's M test by simple dot plot of |S_p| and |S_i|

