

# Statistical Analysis of Method Comparison Studies

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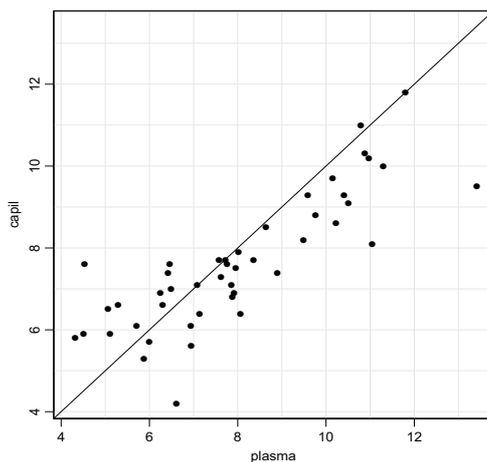
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

## What this is about

- ▶ Two (laboratory) methods for measuring the same clinical quantity.
- ▶ Persons are measured with both methods.
- ▶ Scaled measurements (continuous).
- ▶ Errors in both variables.

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## Glucose measurements



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## Course outlook

- ▶ **Model** based approach
- ▶ Explicit **parametric** models:
  - ▶ Assumptions are made clear
  - ▶ — relaxing assumptions is clear
- ▶ **Comparison** of methods:
  - can one replace the other?
- ▶ **Conversion** between methods:
  - if measurement is  $y_1$  with method 1, what would it be with method 2?
- ▶ Examples from [MethComp](#) package for **R**.
- ▶ Code and output included on the slides
- ▶ — and on the course web-site.

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## Order of topics 19-20 March

- ▶ Wednesday 19th
  - ▶ One measurement by each method
  - ▶ Computing
  - ▶ Linear bias between methods
  - ▶ Variable SD
  - ▶ Practical milk, plvol
  - ▶ Replicate measurements, exchangeable / linked
  - ▶ Practical fat, sbp2
  - ▶ Repeatability, reproducibility
  - ▶ Coefficient of variation
- ▶ Thursday 20th
  - ▶ Replicate measurements and linear bias
  - ▶ Practical ox 1–8
  - ▶ Converting between methods
  - ▶ MCMC methods for estimation of variance components
  - ▶ Practical ox 9–

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## Comparing two methods with one measurement on each

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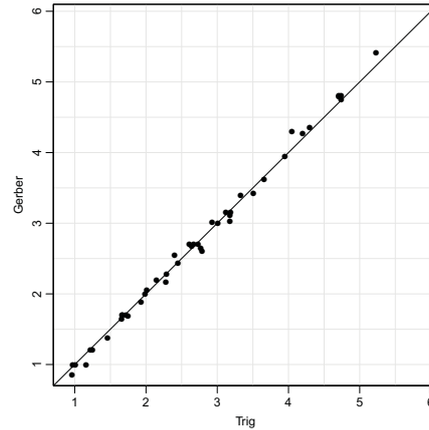
(Comp-simple)

## Comparing measurement methods

General questions:

- ▶ Are results systematically different?
- ▶ Can one method safely be replaced by another?
- ▶ What is the size of measurement errors?
- ▶ Different centres use different methods of measurement: How can we convert from one method to another?
- ▶ How precise is the conversion?

## Fat content in human milk:



The relationship looks like:

$$y_1 = a + by_2$$

```
> library(MethComp)
> # sessionInfo()
> data( milk )
> milk <- Meth( milk )
```

The following variables from the dataframe "milk" are used as the Meth variables:

```
meth: meth
item: item
y: y
```

Method	#Replicates	1	#Items	#Obs:	90	Values:	min	med	max
Gerber	45	45	45	45	0.85	2.67	6.20		
Trig	45	45	45	45	0.96	2.67	6.21		

```
> par( mar=c(3,3,1,1),mgp=c(3,1,0)/1.6 )
> BA.plot( milk, pl.type="comp", col.line="transparent",
+         lwd=c(3,0,0), axlim=c(1,6)-0.1 )
> abline(0,1)
```

## Two methods — one measurement by each

- ▶ How large is the difference between a measurement with method 1 and one with method 2 on a (randomly chosen) person?

$$D_i = y_{2i} - y_{1i}, \quad \bar{D}, \quad \text{s.d.}(D)$$

- ▶ 95% prediction interval for the difference between a measurement by method 1 and one by method 2.  $[1, 2]$
- ▶ **Limits of agreement:**

$$\bar{D} \pm 2 \times \text{s.d.}(D)$$

## Limits of agreement: Interpretation

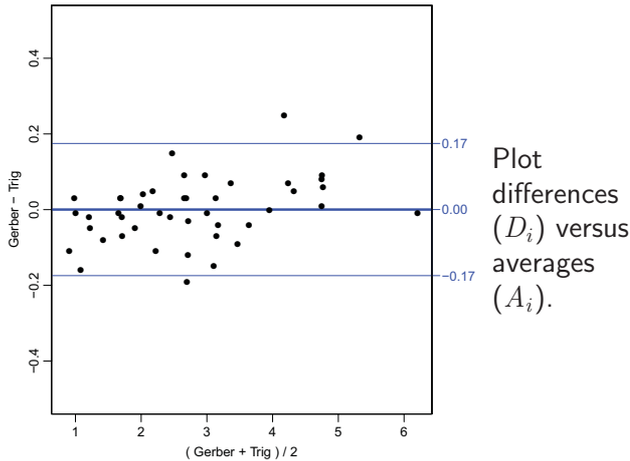
- ▶ If a new patient is measured **once** with each of the two methods, the difference between the two values will with 95% probability be within the limits of agreement.
- ▶ This is a **prediction** interval for a single (future) difference.
- ▶ Interpretation requires a **clinical** input: Are the limits of agreement sufficiently narrow to make the use of either of the methods clinically acceptable?
- ▶ Is it relevant to test if the mean is 0?

## Limits of agreement: Test? **No!**

Testing whether the difference is 0 is a bad idea:

- ▶ Small study: Null accepted even if the difference is important.
- ▶ Large study: Null rejected even if the difference is clinically irrelevant.
- ▶ It is an **equivalence** problem:
  1. How small can we reasonably safely assume the differences to be?
  2. **Testing is irrelevant:** — not interesting if the **mean** difference is significantly different from 0.
  3. **Clinical input is required** to interpret the **prediction** interval.

## Limits of agreement:



Comparing two methods with one measurement on each (Comp-simple)

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```
> par( mar=c(3,3,1,3), mgp=c(3,1,0)/1.6 )
> BA.plot( milk, diflim=c(-0.5,0.5), grid=FALSE )
```

Comparing two methods with one measurement on each (Comp-simple)

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## Model behind "Limits of agreement"

- Methods  $m = 1, \dots, M$ , applied to  $i = 1, \dots, I$  individuals:

$$y_{mi} = \alpha_m + \mu_i + e_{mi}$$

$$e_{mi} \sim \mathcal{N}(0, \sigma_m^2) \quad \text{measurement error}$$

- Two-way analysis of variance model, with **different** variances in columns.
- Different variances are not identifiable without replicate measurements for  $M = 2$ .

The variances  $\sigma_m$  are based on the distance of the obs to the mean across methods, but they are always numerically identical with only 2 methods.

Models

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## Limits of agreement:

- Usually interpreted as the likely difference between two future measurements, one with each method:

$$\widehat{y_2 - y_1} = \hat{D} = \alpha_2 - \alpha_1 \pm 2 \text{s.d.}(D)$$

- Convert to prediction interval for  $y_2$  given  $y_1$ :

$$\hat{y}_{2|1} = \hat{y}_2 | y_1 = \alpha_2 - \alpha_1 + y_1 \pm 2 \text{s.d.}(D)$$

- Formally, we should replace:

$$2 \rightarrow t_{0.975}^{(I-1)} \sqrt{1 + 1/I}$$

which equals 2 for  $I = 85$  and 1.96 for  $I = \infty$

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## Spurious correlation?

Different variances induce correlation between  $D_i$  and  $A_i = (y_{1i} + y_{2i})/2$ , if the variances of  $y_{1i}$  and  $y_{2i}$  are  $\zeta_1^2$  and  $\zeta_2^2$  respectively:

$$\text{cov}(D_i, A_i) = \frac{1}{2}(\zeta_2^2 - \zeta_1^2) \neq 0 \quad \text{if } \zeta_1 \neq \zeta_2$$

In correlation terms:

$$\rho(D, A) = \frac{1}{2} \left( \frac{\zeta_2^2 - \zeta_1^2}{\zeta_1^2 + \zeta_2^2} \right)$$

i.e. the correlation depends on whether the difference between the variances is large relative to the sizes of the two.

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## ... not really...

The variances we were using were the **marginal** variances of  $y_1$  and  $y_2$ :

$$y_{mi} = \alpha_m + \mu_i + e_{mi}$$

$$\text{var}(y_m) = \text{var}(\mu_i) + \sigma_m^2$$

and hence the correlation expression is:

$$\rho(D, A) = \frac{1}{2} \left( \frac{\zeta_2^2 - \zeta_1^2}{\zeta_1^2 + \zeta_2^2} \right) = \frac{1}{2} \left( \frac{\sigma_2^2 - \sigma_1^2}{2\text{var}(\mu_i) + \sigma_1^2 + \sigma_2^2} \right)$$

Hence only relevant if  $\text{var}(\mu_i)$  is small relative to  $\sigma_1^2$  and  $\sigma_2^2$ .

**Not** likely in practise — the  $\mu$ s are normally chosen to be widely spread, so  $\text{var}(\mu_i) \gg \sigma_1^2, \sigma_2^2$

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# Introduction to computing

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<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Intro-comp)

## Course structure

The course is both theoretical and practical, i.e. the aim is to convey a basic understanding of the problems in method comparison studies, but also to convey practical skills in handling the statistical analysis.

- ▶ **R** for data manipulation and graphics.
- ▶ Occasionally BUGS (JAGS) for estimation in non-linear variance component models.

## How it works

Example data sets are included in the MethComp package.

Functions in MethComp are based on a data frame with a particular structure; a **Meth** object:

meth — method (factor)  
item — item, person, individual, sample (factor)  
repl — replicate (if present) (factor)  
y — the actual measurement (numerical)

Once converted to a Meth object, just use `summary`, `plot` etc.

## How it looks I

```
> library( MethComp )  
> data( ox )  
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth  
item: item  
repl: repl  
y: y  
#Replicates  
Method 1 2 3 #Items #Obs: 354 Values: min med max  
CO 1 4 56 61 177 22.2 78.6 93.5  
pulse 1 4 56 61 177 24.0 75.0 94.0
```

```
> ( subset( ox, as.integer(item)<3 ) )
```

## How it looks II

```
meth item repl y  
1 CO 1 1 78.0  
2 CO 1 2 76.4  
3 CO 1 3 77.2  
4 CO 2 1 68.7  
5 CO 2 2 67.6  
6 CO 2 3 68.3  
7 pulse 1 1 71.0  
8 pulse 1 2 72.0  
9 pulse 1 3 73.0  
10 pulse 2 1 68.0  
11 pulse 2 2 67.0  
12 pulse 2 3 68.0
```

```
> subset( to.wide(ox), as.integer(item)<3 )
```

```
item repl CO pulse  
1 1 1 78.0 71  
2 1 2 76.4 72  
3 1 3 77.2 73  
4 2 1 68.7 68  
5 2 2 67.6 67  
6 2 3 68.3 68
```

## Analyses in this course

- ▶ Scatter plots.
- ▶ Bland-Altman plots ( $(y_2 - y_1)$  vs.  $(y_1 + y_2)/2$ )
- ▶ Limits of agreement.
- ▶ Models with constant bias.
- ▶ Models with linear bias.
- ▶ Conversion formulae between methods.
- ▶ Plots of conversion equations.
- ▶ Reporting of variance components.
- ▶ Transformation of response.

## Data objects in MethComp

- ▶ `Meth` Dataframe in the “long” format, with predefined variable names.
- ▶ `MethComp` Results from an analysis with estimated conversions between methods and (if applicable) variance components. Produced by different functions.
- ▶ `MCmcmc` Results from a MCMC analysis of a model. Can be converted to a `MethComp` object.

## Functions in the MethComp package

5 broad categories of functions in MethComp:

- ▶ Data manipulation — reshaping and changing data.
- ▶ Graphical — exploring data.
- ▶ Simulation — generating datasets or replacing variables.
- ▶ Analysis functions — fitting models to data.
- ▶ Reporting functions — displaying the results from analyses.

## Data manipulation functions

- ▶ `Meth` Sets up a `Meth` object — a dataframe in the “long” format, with predefined variable names.
- ▶ `make.repl` Generates a `repl` column in a data frame with columns `meth`, `item` and `y`.
- ▶ `perm.repl` Randomly permutes replicates within (method,item) and assigns new replicate numbers.
- ▶ `to.wide/to.long` Transforms a data frame in the long form to the wide form and vice versa.
- ▶ `Meth.sim` Simulates a dataset (a `Meth` object) from a method comparison experiment.

## Graphical functions (basic)

- ▶ `plot.Meth` Plots all methods against all other, both as a scatter plot and as a Bland-Altman plot.
- ▶ `BA.plot` Makes a Bland-Altman plot of two methods from a data frame with method comparison data, and computes limits of agreement.
- ▶ `bothlines` Adds regression lines of  $y$  on  $x$  and vice versa to a scatter plot.

## Analysis functions (simple)

- ▶ `DA.reg`, regresses the differences on the averages. Also regresses the absolute residuals on the averages to check whether the variance is constant. Returns a `MethComp` object.
- ▶ `BA.est` Estimates in the variance components models underlying the concept of limits of agreement, and returns the bias and the variance components. Assumes constant bias between methods. Returns a `MethComp` object.
- ▶ `VC.est` The workhorse behind `BA.est`.
- ▶ `Deming` Performs Deming regression, i.e. regression with errors in both variables.

## Analysis functions (general)

- ▶ `MCmcmc` Estimates via BUGS (JAGS) in the general model with non-constant bias. Produces an `MCmcmc` object. Which can be converted to a `MethComp` object.
- ▶ `AltReg` Estimates via ad-hoc procedure (alternating regressions) in a model with linear bias between methods. Returns a matrix of estimates with the conversion parameters as well as the variance components. Returns a `MethComp` object.

## Reporting functions

- ▶ `print.MethComp` Prints a table of conversion equations based on an estimated model.
- ▶ `plot.MethComp` Graphs the estimated relationship between methods based on an estimated model.
- ▶ `print.MCmcmc` Table of conversion equations between methods analyzed.
- ▶ `plot.MCmcmc` Conversion lines between methods with prediction limits.
- ▶ `post.MCmcmc` Smoothed posteriors of estimates.
- ▶ `trace.MCmcmc` Simulation traces from an MCmcmc object.

## Does it work? I

You should get something reasonable out of this:

```
> library( MethComp )
> data( ox )
> ox <- Meth( ox )
> summary( ox )
> plot( ox )
> BA.plot( ox )
> BA.est( ox )
> ( AR.ox <- AltReg(ox,linked=TRUE,trace=TRUE) )
> MCmcmc( ox, code.only=TRUE )
> MC.ox <- MCmcmc( ox, n.iter=500 )
> print( MC.ox )
> plot( MC.ox )
> trace.MCmcmc( MC.ox )
> post.MCmcmc( MC.ox )
```

## Non-constant difference

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(Non-const)

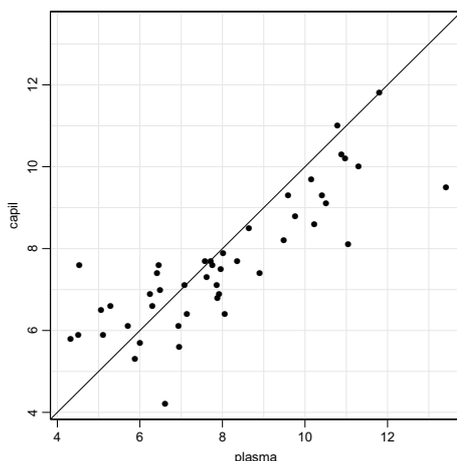
## Limits of agreement — assumptions

- ▶ The difference between methods is constant
- ▶ The variances of the methods (and hence of the difference) is constant
- ▶ “Constant” means constant across the range of measurement values

Check this by:

- ▶ Regress differences on averages.
- ▶ Regress absolute residuals from this on the averages.

## Glucose measurements



```
> options( width=61 )
> library(MethComp)
> data( glucose )
> gluc <-subset( glucose, type %in% levels(type)[c(2,4)] &
+ meth %in% c("h.cap","o.cap","n.plas1"),
+ select=c(2,3,4,6) )
> str( gluc )

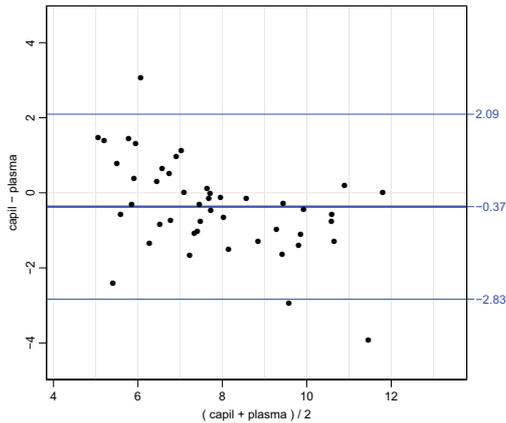
'data.frame': 472 obs. of 4 variables:
 $ type: Factor w/ 4 levels "blood","plasma",...: 2 4 2 4 2 4 2 4 2 4 ...
 $ item: num 1 1 1 1 1 1 1 2 2 ...
 $ time: num 0 0 30 30 60 60 120 120 0 0 ...
 $ y : num 6.36 5.1 10.3 9.8 13.33 ...

> glu120 <- Meth( subset( gluc, time==120 ), meth="type", print=F )
> summary( glu120 )

#Replicates
Method 1 #Items #Obs: 119 Values: min med max
plasma 73 73 73 4.32 7.92 13.42
capil 46 46 46 4.20 7.45 11.80

> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, wh.comp=2:1, pl.type="comp",
+ col.line="transparent" )
> abline( 0, 1 )
```

## Glucose measurements



Non-constant difference (Non-const)

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```
> par( mar=c(3,3,1,3), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, wh.comp=2:1, pl.type="BA" )
```

Non-constant difference (Non-const)

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## Regress differences on averages

$$D_i = a + bA_i + e_i, \quad \text{var}(e_i) = \sigma_D^2$$

If  $b$  is different from 0, we could use this equation to derive LoA:

$$a + bA_i \pm 2\sigma_D$$

or convert to prediction as for LoA:

$$y_{2|1} = y_1 + a + bA_i \approx y_1 + a + by_1 = a + (1 + b)y_1$$

Exchanging methods would give:

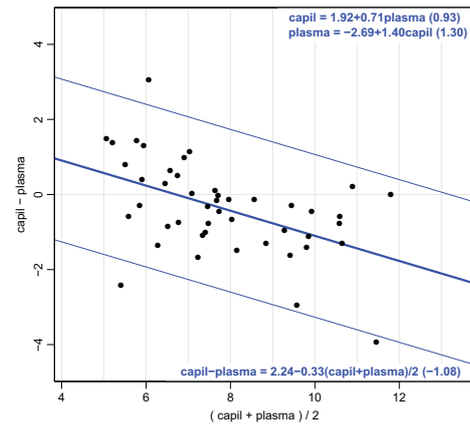
$$y_{1|2} = -a + (1 - b)y_1$$

instead of: 
$$y_{1|2} = \frac{-a}{1 + b} + \frac{1}{1 + b}y_1$$

Non-constant difference (Non-const)

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## Variable limits of agreement



Non-constant difference (Non-const)

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```
> par( mar=c(3,3,1,3), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, dif.type="lin", wh.comp=2:1, pl.type="BA" )
```

```
> par( mar=c(3,3,1,3), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, dif.type="lin", wh.comp=2:1, pl.type="BA",
+         eqn=TRUE )
```

Relationships between methods:  
 capil-plasma = 2.24-0.33(capil+plasma)/2 (-1.08)  
 capil = 1.92+0.71plasma (0.93)  
 plasma = -2.69+1.40capil (1.30)

Non-constant difference (Non-const)

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## Using the regression of $D$ on $A$ properly

$$y_{2i} - y_{1i} = a + b(y_{1i} + y_{2i})/2 + e_i$$

$$y_{2i}(1 - b/2) = a + (1 + b/2)y_{1i} + e_i$$

$$y_{2i} = \frac{a}{1 - b/2} + \frac{1 + b/2}{1 - b/2}y_{1i} + \frac{1}{1 - b/2}e_i$$

$$y_{1i} = \frac{-a}{1 + b/2} + \frac{1 - b/2}{1 + b/2}y_{2i} + \frac{1}{1 + b/2}e_i$$

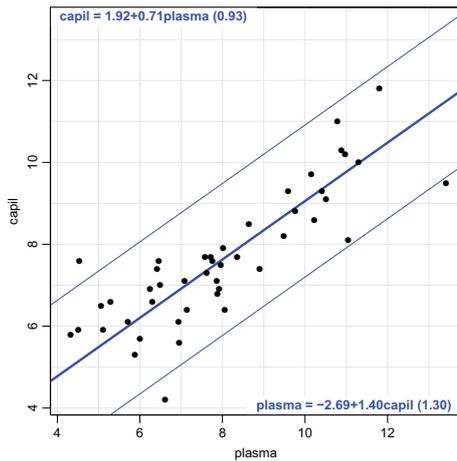
Details found in [5]

This is what comes out of the functions `DA.reg` and `BA.plot`.

Non-constant difference (Non-const)

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## Conversion equation with prediction limits



Non-constant difference (Non-const)

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```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> BA.plot( glui20, dif.type="lin",wh.comp=2:1, pl.type="conv",
+         eqn=TRUE )
```

```
Relationships between methods:
capil-plasma = 2.24-0.33(capil+plasma)/2 (-1.08)
capil = 1.92+0.71plasma (0.93)
plasma = -2.69+1.40capil (1.30)
```

Non-constant difference (Non-const)

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## Why does this work?

The general model for the data is:

$$y_{1i} = \alpha_1 + \beta_1 \mu_i + e_{1i}, \quad e_{1i} \sim \mathcal{N}(0, \sigma_1^2)$$

$$y_{2i} = \alpha_2 + \beta_2 \mu_i + e_{2i}, \quad e_{2i} \sim \mathcal{N}(0, \sigma_2^2)$$

- ▶ Work out the prediction of  $y_{2i}$  given an observation of  $y_{1i}$  in terms of the  $\alpha$ s and  $\beta$ s.
- ▶ Work out how differences relate to averages in terms of  $\alpha$ s and  $\beta$ s.
- ▶ Use til to work out relationship between the  $(\alpha, \beta)$  and  $(a, b)$
- ▶ Then the prediction is as we just derived it.

Non-constant difference (Non-const)

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## So why is it wrong anyway?

### Conceptually:

Once the  $\beta_m$  is introduced:

$$y_{mi} = \alpha_m + \beta_m \mu_i + e_{mi}$$

measurements by different methods are on different scales.

Hence it has formally no meaning to form the differences.

Non-constant difference (Non-const)

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## So why is it wrong anyway?

### Statistically:

Under the correctly specified model, the induced model for the differences on the averages  $A_i$ , these contain the error terms, and so does the residuals.

So the covariate is not independent of the error terms.

Thus the assumptions behind regression are violated.

Non-constant difference (Non-const)

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## Then why use it?

- ▶ With only one observation per (method,item) there is not much else to do.
- ▶ If the slope linking the two methods ( $\beta_1/\beta_2$ ) is not dramatically different from 1, the violations are not that big.
- ▶ Implemented in `BA.plot` and in `DA.reg`, which also checks the residuals.

For further details, see [5].

Non-constant difference (Non-const)

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## Limits of agreement — assumptions

- ▶ The difference between methods is constant
- ▶ The variances of the methods (and hence of the difference) is constant
- ▶ Residuals follow a normal distribution

Check this by:

- ▶ Regress differences on averages
- ▶ Regress absolute residuals from this on the averages
- ▶ ... the central limit theorem?

## Regressing residuals on averages

- ▶ Residuals  $\sim \mathcal{N}(0, \sigma^2)$   
 $\Rightarrow$  absolute residuals half-normal.
- ▶ Mean of standard half normal is:

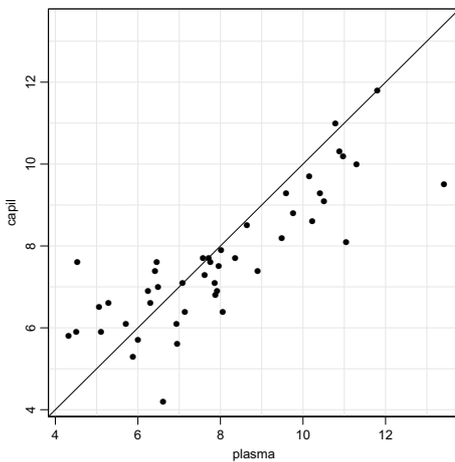
$$\int_0^{\infty} x(2/\sqrt{2\pi})\exp(-x^2/2) dx = \sqrt{2/\pi}$$

- ▶ Mean of absolute residuals is  $\sigma\sqrt{2/\pi}$
- ▶ Linear relationship of absolute residuals ( $R_i$ ) to averages ( $A_i$ ):

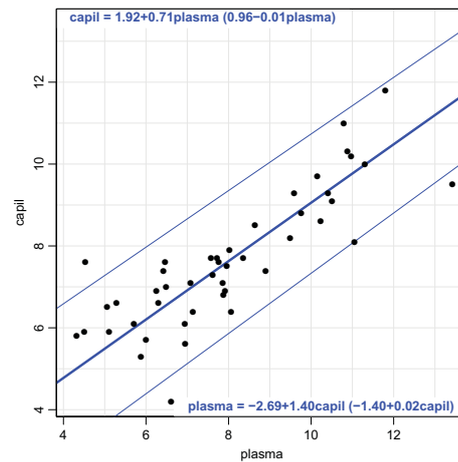
$$R_i = a + bA_i \Leftrightarrow \sigma(A) \approx a\sqrt{\pi/2} + b\sqrt{\pi/2}A$$

- ▶ Implemented in DA.reg.

## Glucose measurements



## Variable standard deviation



```
> ( da <- DA.reg( glu120 ) )

Conversion between methods:
      alpha  beta  sd.pr beta=1 in(t-f) sl(t-f) sd(t-f) in(sd) sl(sd)
To: From:
plasma plasma 0.000 1.000    NA    NA 0.000 0.000    NA    NA    N
capil  plasma -2.695 1.402  1.302 0.000 -2.244 0.335  1.084 1.138 -0.01
capil  plasma  1.922 0.713  0.928 0.000  2.244 -0.335 -1.084 1.138 -0.01
capil  capil   0.000 1.000    NA    NA 0.000 0.000    NA    NA    N

> round( ftable( da$Conv[, -(1:4)] ), 3 )

      in(t-f) sl(t-f) sd(t-f) in(sd) sl(sd)  sd=K LoA-lo LoA-up
To: From:
plasma plasma 0.000 0.000    NA    NA    NA    NA    NA    NA
capil  plasma -2.244 0.335  1.084 1.138 -0.015 0.833 -2.095 2.833
capil  plasma  2.244 -0.335 -1.084 1.138 -0.015 0.833 -2.833 2.095
capil  capil   0.000 0.000    NA    NA    NA    NA    NA    NA

> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, wh.comp=2:1, pl.type="BA", dif.type="const",
+         sd.type="lin", eqn=TRUE )
```

```
Relationships between methods:
capil-plasma = -0.37 (1.70-0.07Avg.)
capil = -0.37+plasma (1.65-0.07plasma)
plasma = 0.37+capil (-1.75+0.07capil)

> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, wh.comp=2:1, pl.type="BA",
+         dif.type="lin", sd.type="lin", eqn=TRUE )

Relationships between methods:
capil-plasma = 2.24-0.33(capil+plasma)/2 (1.14-0.02Avg.)
capil = 1.92+0.71plasma (0.96-0.01plasma)
plasma = -2.69+1.40capil (-1.40+0.02capil)

> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> BA.plot( glu120, wh.comp=2:1, pl.type="comp",
+         dif.type="lin", sd.type="lin", eqn=TRUE )

Relationships between methods:
capil-plasma = 2.24-0.33(capil+plasma)/2 (1.14-0.02Avg.)
capil = 1.92+0.71plasma (0.96-0.01plasma)
plasma = -2.69+1.40capil (-1.40+0.02capil)
```

## Variable mean and standard deviation

- ▶ 2-step procedure:
  - ▶ Regress  $D_i$  on  $A_i$ .
  - ▶ Regress  $R_i$  (absolute residuals) on  $A_i$
- ▶ Can be done using quadratic rather than linear terms, or even splines. (Not in MethComp — yet, any takers?)
- ▶ Allows very flexible form of the relationships between differences and averages
- ▶ —and flexible form of the s.d. to the mean.
- ▶ The relationship  $D \sim A$  is easily back-transformed to a relationship  $y_1 \sim y_2$ , with prediction intervals.
- ▶ Beware: **over-modelling!**

## Comparing two methods with replicate measurements

### Bendix Carstensen

SAoMCS  
19–20 March 2014  
Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Comp-repl)

## Replicate measurements on each item

Fat data; **exchangeable** replicates:

item	repl	KL	SL
1	1	4.5	5.0
1	2	4.7	4.9
1	3	4.4	4.8
3	1	6.4	6.5
3	2	6.2	6.4
3	3	6.5	6.1

Oximetry data; **linked** replicates:

item	repl	CO	pulse
1	1	78.0	71
1	2	76.4	72
1	3	77.2	73
2	1	68.7	68
2	2	67.6	67
2	3	68.3	68

## Replicate measurements on each item

Fat data; exchangeable replicates:

item	repl	KL	SL
1	1	4.5	4.9
1	2	4.4	5.0
1	3	4.7	4.8
3	1	6.4	6.5
3	2	6.2	6.4
3	3	6.5	6.1

Oximetry data; **linked** replicates:

item	repl	CO	pulse
1	1	77.2	73
1	2	78.0	71
1	3	76.4	72
2	1	68.7	68
2	2	67.6	67
2	3	68.3	68

## Extension of the model: exchangeable replicates

$$y_{mir} = \alpha_m + \mu_i + c_{mi} + e_{mir}$$

s.d. ( $c_{mi}$ ) =  $\tau_m$  — “matrix”-effect  
s.d. ( $e_{mir}$ ) =  $\sigma_m$  — measurement error

- ▶ Replicates within  $(m, i)$  is needed to separate  $\tau$  and  $\sigma$ .
- ▶ Even with replicates, the  $\tau$ s are only estimable if  $M > 2$ .
- ▶ Still assumes that the difference between methods is constant.
- ▶ Assumes **exchangeability** of replicates.

## Extension of the model: linked replicates

$$y_{mir} = \alpha_m + \mu_i + a_{ir} + c_{mi} + e_{mir}$$

s.d. ( $a_{ir}$ ) =  $\omega$  — between replicates  
s.d. ( $c_{mi}$ ) =  $\tau_m$  — “matrix”-effect  
s.d. ( $e_{mir}$ ) =  $\sigma_m$  — measurement error

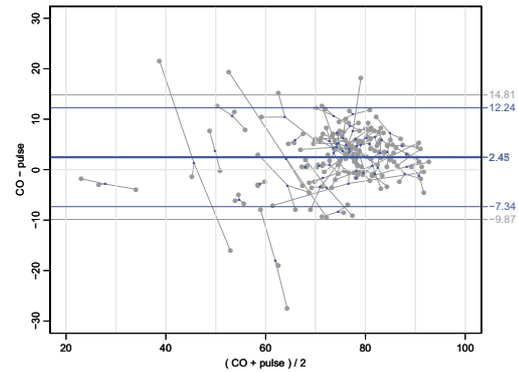
- ▶ Still assumes difference between methods constant.
- ▶ Replicates **linked** between methods:  $a_{ir}$  is common across methods; first replicate on a person is made under similar conditions for all methods, second too etc.

## Replicate measurements

Three approaches to LoA with replicate measurements:

- Means over replicates within each method by item stratum.
- Replicates within item are taken as items.
- Fit the model and use it for the LoA:
  - The model is a standard linear mixed model with separate variances per method.
  - The model is fitted using `BA.est(data,linked=TRUE)` — later.

## Oximetry data



```
> library(MethComp)
> data(ox)
> ox <- Meth(ox, print=FALSE)
> summary(ox)

#Replicates
Method  1  2  3 #Items #Obs: 354 Values: min med max
CO      1  4  56   61   177   22.2 78.6 93.5
pulse   1  4  56   61   177   24.0 75.0 94.0

> par(mar=c(3,3,1,3), mgp=c(3,1,0)/1.6)
> BA.plot(ox, pl.type="BA",
+         xlim=c(20,100), diflim=c(-30,30))

> par(mar=c(3,3,1,3), mgp=c(3,1,0)/1.6)
> BA.plot(ox, pl.type="BA", col.points=gray(0.5), repl.conn=TRUE,
+         xlim=c(20,100), diflim=c(-30,30), col.lines=gray(0.5))

> par(mar=c(3,3,1,3), mgp=c(3,1,0)/1.6)
> BA.plot(ox, pl.type="BA", col.points=gray(0.6), repl.conn=TRUE,
+         xlim=c(20,100), diflim=c(-30,30), col.lines=gray(0.6))
> par(new=TRUE)
> BA.plot(mean(ox), pl.type="BA", col.points="blue", cex=0.5,
+         xlim=c(20,100), diflim=c(-30,30))
```

## Replicate measurements

- The limits of agreement should still be for difference between future **single** measurements.
- Analysis based on the **means** of replicates is therefore **wrong**:
- If the model is:

$$y_{mir} = \alpha_m + \mu_i + a_{ir} + c_{mi} + e_{mir}$$

- ... then the correct limits of agreement are:

$$\alpha_1 - \alpha_2 \pm 2\sqrt{\tau_1^2 + \tau_2^2 + \sigma_1^2 + \sigma_2^2}$$

## Wrong or almost right?

- $\text{var}(y_{1jr} - y_{2jr}) = \tau_1^2 + \tau_2^2 + \sigma_1^2 + \sigma_2^2$   
— note that the term  $a_{ir} - a_{ir}$  cancels because we are referring to the **same** replicate.
- If we are using means of replicates to form the differences we have:

$$\begin{aligned} \bar{d}_i &= \bar{y}_{1i} - \bar{y}_{2i} \\ &= \alpha_1 - \alpha_2 + \sum_r a_{ir}/R_{1i} - \sum_r a_{ir}/R_{2i} \\ &\quad + c_{1i} - c_{2i} + \sum_r e_{1ir}/R_{1i} - \sum_r e_{2ir}/R_{2i} \\ \Rightarrow \\ \text{var}(\bar{d}_i) &= \tau_1^2 + \tau_2^2 + \sigma_1^2/R_{1i} + \sigma_2^2/R_{2i} \\ &< \tau_1^2 + \tau_2^2 + \sigma_1^2 + \sigma_2^2 \end{aligned}$$

## (Linked) replicates as items

- If replicates are taken as items, then the differences are:

$$d_{ir} = y_{1ir} - y_{2ir} = \alpha_1 - \alpha_2 + c_{1i} - c_{2i} + e_{1ir} - e_{2ir}$$

- which has variance  $\tau_1^2 + \tau_2^2 + \sigma_1^2 + \sigma_2^2$ , and so gives the correct limits of agreement.
- But the differences are not independent:

$$\text{cov}(d_{ir}, d_{is}) = \tau_1^2 + \tau_2^2$$

- Negligible if the residual variances are very large compared to the interaction, variance likely to be only slightly downwards biased.

## Exchangeable replicates as items?

- ▶ Exchangeable replicates: not clear how to produce the differences with replicates as items.
- ▶ If replicates are paired at random (see the function `perm.repl`), the variance will still be correct using the model without the  $i \times r$  interaction term ( $a_{ir}$ ):

$$\text{var}(y_{1ir} - y_{2is}) = \tau_1^2 + \sigma_1^2 + \tau_2^2 + \sigma_2^2$$

- ▶ Differences will be positively correlated within item:

$$\text{cov}(y_{1ir} - y_{2is}, y_{1it} - y_{2iu}) = \tau_1^2 + \tau_2^2$$

— slight underestimate of the true variance.

## Recommendations

- ▶ Fit the correct model, and get the estimates from that, e.g. by using `BA.est`.
- ▶ If you must use over-simplified methods:
  - ▶ Use linked replicates as item.
  - ▶ If replicates are not linked; make a random linking.
  - ▶ Note: If this give a substantially different picture than using the original replicate numbering as linking key, there might be something fishy about the data.

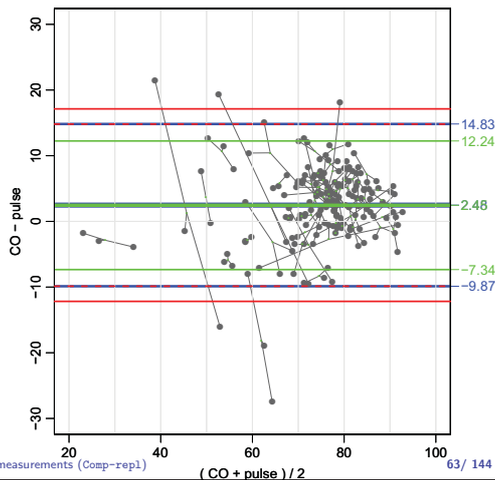
Further details, see [6].

## Oximetry data (linked replicates)

Linked replicates used as items

Mean over replicates as items

Limits based on model — dashed line assuming linked, full exchangeable replicates



```
> ( ox.link <- BA.est( ox, linked=TRUE ) )
```

```
Conversion between methods:
      alpha  beta  sd.pr  LoA-lo  LoA-up
To: From:
CO   CO      0.000  1.000   3.146  -6.293   6.293
     pulse  2.470  1.000   6.169  -9.867  14.808
pulse CO     -2.470  1.000   6.169  -14.808  9.867
     pulse  0.000  1.000   5.649  -11.298  11.298
```

```
Variance components (sd):
      IxR  MxI  res
CO     3.416  2.928  2.225
pulse  3.416  2.928  3.994
```

```
> ( ox.exch <- BA.est( ox, linked=FALSE ) )
```

```
Conversion between methods:
      alpha  beta  sd.pr  LoA-lo  LoA-up
To: From:
CO   CO      0.000  1.000   5.755  -11.509  11.509
     pulse  2.476  1.000   7.326  -12.175  17.127
pulse CO     -2.476  1.000   7.326  -17.127  12.175
     pulse  0.000  1.000   7.417  -14.835  14.835
```

```
Variance components (sd):
      IxR  MxI  res
CO     0  2.191  4.069
pulse  0  2.191  5.245
```

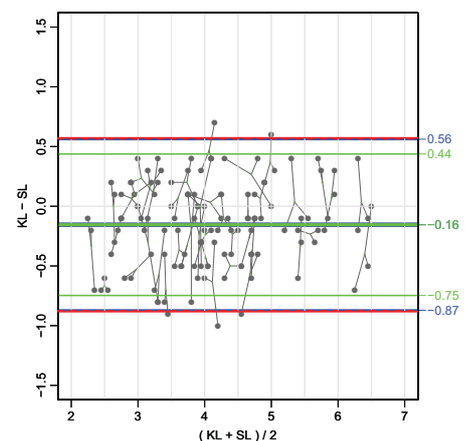
```
> par( mar=c(3,3,1,3), mgp=c(3,1,0)/1.6 )
> BA.plot( ox, pl.type="BA", model=NULL,
+         col.points=gray(0.4), repl.conn=TRUE,
+         xlim=c(20,100), ylim=c(-30,30), col.lines="blue",
+         lwd=c(6,3,3) )
> par( new=TRUE )
> BA.plot( mean(ox), pl.type="BA", col.points="green",
+         cex.points=0.3, xlim=c(20,100), ylim=c(-30,30),
+         col.lines="green", lwd=c(4,2,2) )
> abline( h=ox.link[["LoA"]][2:3], col="red", lwd=2, lty=2 )
> abline( h=ox.exch[["LoA"]][2:3], col="red", lwd=2, lty=1 )
```

## Visceral fat data (exchangeable replicates)

Randomly paired replicates used as items

Mean over replicates as items

Limits based on model — dashed line assuming linked, full exchangeable replicates



```
> data(fat)
> vis <- Meth(fat, 2, 1, 3, 5)
```

The following variables from the dataframe "fat" are used as the Meth variables:

```
meth: Obs
item: Id
repl: Rep
y: Vic
```

```
#Replicates
Method 3 #Items #Obs: 258 Values: min med max
KL 43 43 129 2.0 3.9 6.5
SL 43 43 129 2.3 4.1 6.7
```

```
> (vis.link <- BA.est(vis, linked=TRUE))
```

```
Conversion between methods:
alpha beta sd.pr LoA-lo LoA-up
To: From:
KL KL 0.000 1.000 0.264 -0.528 0.528
SL -0.155 1.000 0.360 -0.874 0.564
SL KL 0.155 1.000 0.360 -0.564 0.874
SL 0.000 1.000 0.235 -0.471 0.471
```

```
Variance components (sd):
IxR MxI res
KL 0.048 0.183 0.187
SL 0.048 0.183 0.166
```

```
> (vis.exch <- BA.est(vis, linked=FALSE))
```

```
Conversion between methods:
alpha beta sd.pr LoA-lo LoA-up
To: From:
KL KL 0.000 1.000 0.273 -0.545 0.545
SL -0.155 1.000 0.364 -0.883 0.573
SL KL 0.155 1.000 0.364 -0.573 0.883
SL 0.000 1.000 0.245 -0.490 0.490
```

```
Variance components (sd):
IxR MxI res
KL 0 0.181 0.193
SL 0 0.181 0.173
```

```
> par(mar=c(3,3,1,3), mgp=c(3,1,0)/1.6)
> BA.plot(vis, pl.type="BA", model=NULL,
+ col.points=gray(0.4), repl.conn=TRUE,
+ xlim=c(2,7), diflim=c(-3,3)/2, col.lines="blue",
+ lwd=c(6,3,3))
> par(new=TRUE)
> BA.plot(mean(vis), pl.type="BA", col.points="green",
+ cex.points=0.3, xlim=c(2,7), diflim=c(-3,3)/2,
+ col.lines="green", lwd=c(4,2,2))
> abline(h=-vis.link[["LoA"]][2:3], col="red", lwd=2, lty=2)
> abline(h=-vis.exch[["LoA"]][2:3], col="red", lwd=2, lty=1)
```

## How the data is generated I

- ▶ A statistical model is a description of a machinery that may have generated data
- ▶ Illustrate how the various components make up the observed data.

```
> source("mc-ill.R")
> mc.ill
```

## How the data is generated II

```
function(prefix, Nm = 2, Ni = 11, Nr = 3, alpha = c(-4, 7),
beta = c(0.95, 1.05), sigma.ir = 5, sigma.mi = c(3, 5), sigma.mir = c(2,
3))
{
meth <- rep(1:Nm, Ni)
item <- rep(1:Ni, each = Nm)
reps <- rep(Nr, length(meth))
dfr <- data.frame(meth = meth, item = item)[rep(1:length(meth),
reps), ]
dfr <- make.repl(dfr)
dfr <- dfr[with(dfr, order(meth, item, repl)), ]
mu <- runif(Ni, 15, 85)
dfr$mu <- mu[dfr$item]
dfr$alpha <- alpha[dfr$meth]
dfr$beta <- beta[dfr$meth]
e.ir <- rnorm(nlevels(IR) <- with(dfr, interaction(item, repl))),
mean = 0, sd = sigma.ir)
dfr$e.ir <- e.ir[as.integer(IR)]
e.mi <- rnorm(nlevels(MI) <- with(dfr, interaction(meth, item))),
mean = 0, sd = sigma.mi)
dfr$e.mi <- e.mi[as.integer(MI)]
dfr$e.mir <- rnorm(nrow(dfr), mean = 0, sd = sigma.mir[meth])
dfr <- transform(dfr, y = alpha + beta * (mu + e.ir + e.mi) +
e.mir, yrm = alpha + beta * (mu + e.ir + e.mi), yr = alpha +
beta * (mu + e.ir), y0 = alpha + beta * mu)
dfr
```

## How the data is generated III

```
d1 <- subset(dfr, meth == 1)
d2 <- subset(dfr, meth == 2)
mu1 <- d1$mu
y10 <- d1$y0
y1r <- d1$yr
y1m <- d1$yrm
y1f <- d1$y
mu2 <- d2$mu
y20 <- d2$y0
y2r <- d2$yr
y2m <- d2$yrm
y2f <- d2$y
x <- 4
xx <- 1.7
clr <- rainbow(Ni)
pdf(paste("../graph/", prefix, "-ill-1.pdf", sep = ""), height = 2 *
x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y10, xlim = c(0, 100), ylim = c(0, 100), xlab = expression(mu),
ylab = "y1", pch = 16, cex = xx, col = clr[d1$item])
abline(0, 1)
plot(mu2, y20, xlim = c(0, 100), ylim = c(0, 100), xlab = expression(mu),
ylab = "y2", pch = 16, cex = xx, col = clr[d1$item])
abline(0, 1)
plot(y10, y20, xlim = c(0, 100), ylim = c(0, 100), xlab = "y1",
```

## How the data is generated IV

```
ylab = "y2", pch = 16, cex = xx, col = clr[d1$item])
abline(0, 1)
dev.off()
pdf(paste("../graph/", prefix, "-ill-2.pdf", sep = ""), height = 2 *
x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y10, xlim = c(0, 100), ylim = c(0, 100), col = clr[d1$item],
xlab = expression(mu), ylab = "y1", pch = 1, lwd = 2,
cex = xx)
segments(mu1, y10, mu1, y1r, col = grey(0.7))
points(mu1, y1r, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(mu2, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
xlab = expression(mu), ylab = "y2", pch = 1, lwd = 2,
cex = xx)
segments(mu2, y20, mu2, y2r, col = grey(0.7))
points(mu2, y2r, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(y10, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
xlab = "y1", ylab = "y2", pch = 1, lwd = 2, cex = xx)
segments(y10, y20, y1r, y2r, col = clr[d1$item])
points(y1r, y2r, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
dev.off()
pdf(paste("../graph/", prefix, "-ill-3.pdf", sep = ""), height = 2 *
```

## How the data is generated V

```
x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y10, xlim = c(0, 100), ylim = c(0, 100), col = clr[d1$item],
      xlab = expression(mu), ylab = "y1", pch = 1, lwd = 2,
      cex = xx)
segments(mu1, y10, mu1, y1r, col = clr[d1$item])
points(mu1, y1r, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(mu1, y1r, mu1, y1m, col = clr[d1$item])
points(mu1, y1m, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(mu2, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = expression(mu), ylab = "y2", pch = 1, lwd = 2,
      cex = xx)
segments(mu2, y20, mu2, y2r, col = clr[d1$item])
points(mu2, y2r, col = clr[d1$item], pch = 16, cex = xx)
segments(mu2, y2r, mu2, y2m, col = clr[d1$item])
points(mu2, y2m, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(y10, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = "y1", ylab = "y2", pch = 1, lwd = 2, cex = xx)
segments(y10, y20, y1r, y2r, col = clr[d1$item])
points(y1r, y2r, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(y1r, y2r, y1m, y2m, col = clr[d1$item])
points(y1m, y2m, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
```

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## How the data is generated VI

```
dev.off()
pdf(paste("../graph/", prefix, "-ill-4.pdf", sep = ""), height = 2 *
    x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y10, xlim = c(0, 100), ylim = c(0, 100), col = clr[d1$item],
      xlab = expression(mu), ylab = "y1", pch = 1, lwd = 2,
      cex = xx)
segments(mu1, y10, mu1, y1r, col = clr[d1$item])
points(mu1, y1r, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(mu1, y1r, mu1, y1m, col = clr[d1$item])
points(mu1, y1m, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(mu1, y1m, mu1, y1f, col = clr[d1$item])
points(mu1, y1f, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(mu2, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = expression(mu), ylab = "y2", pch = 1, lwd = 2,
      cex = xx)
segments(mu2, y20, mu2, y2r, col = clr[d1$item])
points(mu2, y2r, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(mu2, y2r, mu2, y2m, col = clr[d1$item])
points(mu2, y2m, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(mu2, y2m, mu2, y2m, col = clr[d1$item])
points(mu2, y2m, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
plot(y10, y20, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = "y1", ylab = "y2", pch = 1, lwd = 2, cex = xx)
```

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## How the data is generated VII

```
xlab = "y1", ylab = "y2", pch = 1, lwd = 2, cex = xx)
segments(y10, y20, y1r, y2r, col = clr[d1$item])
points(y1r, y2r, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(y1r, y2r, y1m, y2m, col = clr[d1$item])
points(y1m, y2m, col = clr[d1$item], pch = 1, lwd = 2, cex = xx)
segments(y1m, y2m, y1f, y2f, col = clr[d1$item])
points(y1f, y2f, col = clr[d1$item], pch = 16, cex = xx)
abline(0, 1)
dev.off()
pdf(paste("../graph/", prefix, "-ill-5.pdf", sep = ""), height = 2 *
    x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y1f, xlim = c(0, 100), ylim = c(0, 100), col = clr[d1$item],
      xlab = expression(mu), ylab = "y1", pch = 16, lwd = 2,
      cex = xx)
abline(0, 1)
plot(mu2, y2f, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = expression(mu), ylab = "y2", pch = 16, lwd = 2,
      cex = xx)
abline(0, 1)
plot(y1f, y2f, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = "y1", ylab = "y2", pch = 16, lwd = 2, cex = xx)
abline(0, 1)
```

Comparing two methods with replicate measurements (Comp-rep1)

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## How the data is generated VIII

```
dev.off()
pdf(paste("../graph/", prefix, "-ill-6.pdf", sep = ""), height = 2 *
    x + 2, width = 3 * x + 3, pointsize = 21)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
layout(matrix(c(1, 2, 3, 3, 3, 3), 2, 3))
par(mai = c(3, 3, 1, 1)/4, mgp = c(3, 1, 0)/1.6)
plot(mu1, y1f, xlim = c(0, 100), ylim = c(0, 100), col = clr[d1$item],
      xlab = expression(mu), ylab = "y1", pch = 16, lwd = 2,
      cex = xx)
abline(0, 1)
plot(mu2, y2f, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = expression(mu), ylab = "y2", pch = 16, lwd = 2,
      cex = xx)
abline(0, 1)
plot(y1f, y2f, xlim = c(0, 100), ylim = c(0, 100), , col = clr[d1$item],
      xlab = "y1", ylab = "y2", pch = 16, lwd = 2, cex = xx)
abline(alpha[2] - alpha[1] * beta[2]/beta[1], beta[2]/beta[1],
        lwd = 3, col = gray(0.6))
dev.off()

> library( MethComp )
> mc.ill("vcx",beta=c(1,1),sigma.ir=0)
```

Comparing two methods with replicate measurements (Comp-rep1)

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## How the data is generated IX

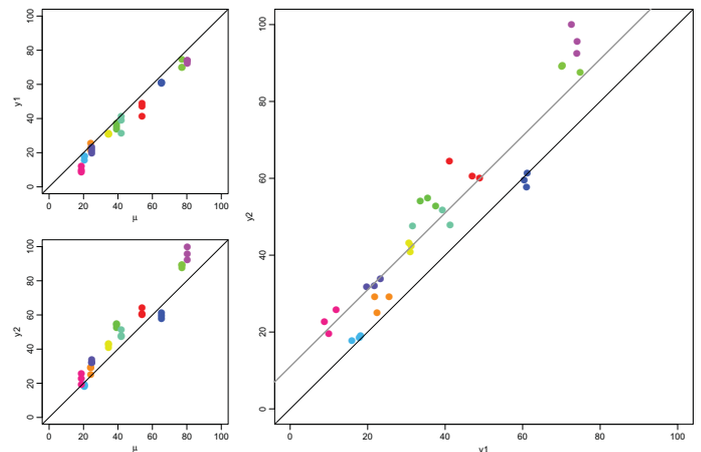
```
null device
1

> mc.ill("vc1",beta=c(1,1),sigma.ir=5)

null device
1
```

Comparing two methods with replicate measurements (Comp-rep1)

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$$y_{mir} = \alpha_m + \mu_i + C_{mi} + e_{mir}$$

Comparing two methods with replicate measurements (Comp-rep1)

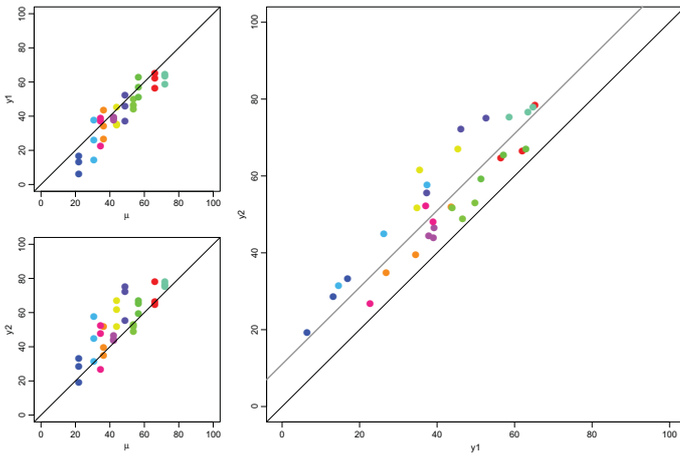
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# Repeatability and reproducibility

Bendix Carstensen

SAoMCS  
19–20 March 2014  
Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Repro)



$$y_{mir} = \alpha_m + \mu_i + a_{ir} + c_{mi} + e_{mir}$$

Comparing two methods with replicate measurements (Comp-rep1)

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## Accuracy of a measurement method

(ISO 5625)

- ▶ Repeatability:  
The accuracy of the method under exactly similar circumstances; i.e. the same lab, the same technician, and the same day.  
(**Repeatability conditions**)
- ▶ Reproducibility:  
The accuracy of the method under comparable circumstances, i.e. the same machinery, the same kit, but possibly different days or laboratories or technicians.  
(**Reproducibility conditions**)

Repeatability and reproducibility

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## Quantification of accuracy

- ▶ Upper limit of a 95% confidence interval for the difference between two measurements.
- ▶ Suppose the variance of the measurement is  $\sigma^2$ :

$$\text{var}(y_{mi1} - y_{mi2}) = 2\sigma^2$$

- standard error of difference:  $\sqrt{2}\sigma$
- ▶ Confidence interval for the difference:

$$0 \pm 1.96 \times \sqrt{2}\sigma = 0 \pm 2.772\sigma \approx \pm 2.8\sigma$$

- ▶ This is called the reproducibility coefficient or simply the **reproducibility**.  
(2.8 is used as a convenient approximation).

Repeatability and reproducibility

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## Quantification of accuracy

- ▶ Where do we get the  $\sigma$ ?
- ▶ Repeat measurements on the same item.
- ▶ The conditions under which the repeat (replicate) measurements are taken determines whether we are estimating repeatability or reproducibility.
- ▶ In larger experiments we must consider the **exchangeability** of the replicates — i.e. which replicates are done under (exactly) similar conditions and which are not.

Repeatability and reproducibility

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## Coefficient of variation

- ▶ Defined as s.d. relative to mean:  $CV = \sigma/\mu$
- ▶ Measurements with varying mean and s.d. may still have constant CV.
- ▶ Assumption of s.d. proportional to  $\mu$  across the range of  $y$ ,  $\text{s.d.}(y) = CV\mu(y)$   
— implies that measurements are positive.
- ▶ LoA could be:

$$\mu \pm 2CV\mu$$

- ▶ But what if  $CV > 0.5$  — lower bound  $< 0$ ?
- ▶ Immaterial — "2" depends on the degree of confidence chosen anyway.

Repeatability and reproducibility

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## Coefficient of variation

- ▶  $\sigma$  proportional to  $\mu$
- ▶  $\Rightarrow$  confidence intervals should be multiplicative:  $\mu \times \text{erf}$  for some error-factor.
- ▶ Specifically:

$$\text{s.d.}(\log(Y)) \approx \sigma \times \left. \frac{d\log(y)}{dy} \right|_{y=\mu} = \sigma/\mu = CV$$

- ▶ ... so using CV is just doing analysis on the log-scale.

## Coefficient of variation

- ▶ CV small:  
CV is the same as the s.d. of the log-transformed data.
- ▶ CV large:  
CV is **not** same as the s.d. of the log-transformed data.
- ▶ ... but it is the log-transformed analysis that is meaningful.
- ▶ Empirical question if this gives a better model.

## A common misconception

*There are other approaches that might also be used (e.g., coefficients of variation, item response theory, or the “signal to noise ratio”). [7]<sup>1</sup>*

- ▶ The authors seem to think that coefficient of variation is another **model**.
- ▶ It is not a different model — just the same model on a transformed **scale**,
- ▶ — focusing on the variance (of the log-transformed data)

<sup>1</sup>Guidelines for Reporting Reliability and Agreement Studies (GRRAS)

## Linear bias between methods

### Bendix Carstensen

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19–20 March 2014  
Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Lin-bias)

## Extension with non-constant bias

$$y_{mir} = \alpha_m + \beta_m \mu_i + \text{random effects}$$

- ▶ There is now a **scaling** between the methods.
- ▶ Methods do not measure on the same scale — the relative scaling is **estimated**, between method 1 and 2 the scale is  $\beta_2/\beta_1$ .
- ▶ Consequence: Multiplication of all measurements on one method by a fixed number does not change results of analysis:
  - ▶ The  $\alpha$ s &  $\beta$ s are multiplied by the same factor
  - ▶ as is the s.d.s of the variance components for this method.

## Variance components

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

- ▶ The random effects  $c_{mi}$  and  $e_{mir}$  have variances specific for each method.
- ▶ Variance of  $a_{ir}$  does not depend on  $m$  — reporting scaled to each of the methods by the corresponding  $\beta_m$ .
- ▶ Implies that  $\omega = \text{s.d.}(a_{ir})$  is irrelevant — the scale is arbitrary.
- ▶ Relevant quantities are  $\beta_m \omega$  — the between replicate variation within item **as measured on the  $m$ th scale**.

## Variance components

Method, Item, Replicate.

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

$$\text{s.d.}(c_{mi}) = \tau_m$$

- ▶ **Matrix-effect:** Each item reacts differently to each method.
- ▶ If only two methods:
  - ▶  $\tau_1$  and  $\tau_2$  cannot be separated.
  - ▶ Variances must be reported on the scale of each method, as  $\beta_m \tau_m$ .

## Variance components

Method, Item, Replicate.

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

$$\text{s.d.}(a_{ir}) = \omega$$

- ▶ Common across methods — must be scaled relative to the methods.
- ▶ Included if replicates are linked across methods, e.g. if there is a sequence in the replicates.
- ▶  $a_{ir}$  nuisance parameters —  $(\mu_i + a_{ir})$  is the “true” value underlying measurements  $y_{mir}$ .

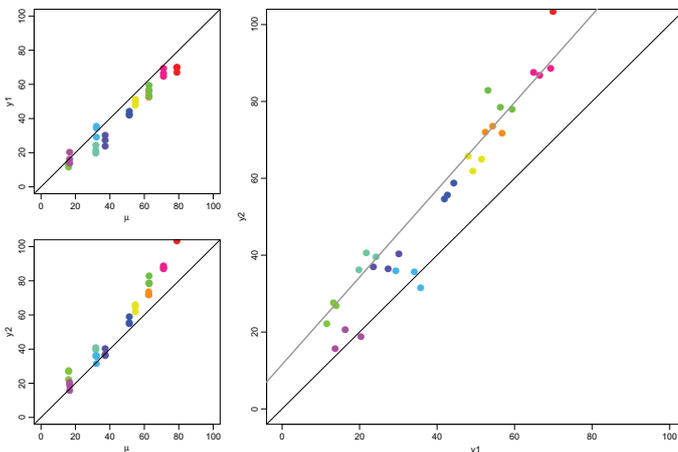
## Estimation in the extended model

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

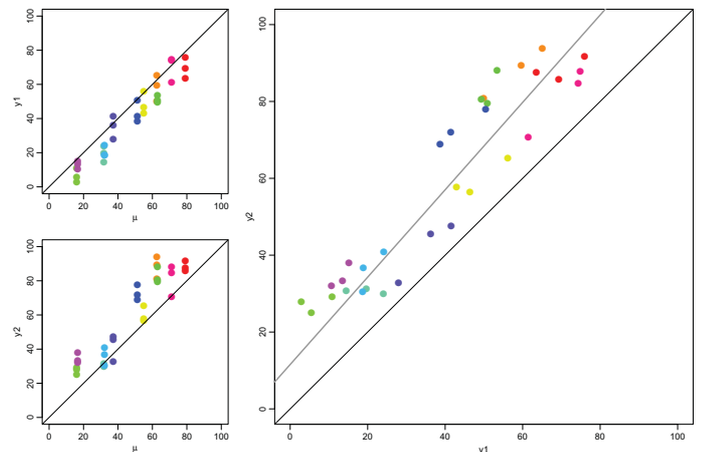
- ▶ Not a standard linear mixed model.
- ▶ Does not fit into usual software.
- ▶ Fitted in BUGS, using JAGS via MCmcmc.
- ▶ ... or AltReg — we shall return to this later

## How the data is generated I

- ▶ A statistical model is a description of a machinery that may have generated data
- ▶ Illustrate how the various components make up the observed data.



$$y_{mir} = \alpha_m + \beta_m(\mu_i + c_{mi}) + e_{mir}$$



$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

# Converting between methods

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Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Convert)

## Predicting method 2 from method 1

$$\begin{aligned} y_{10r} &= \alpha_1 + \beta_1(\mu_0 + a_{0r} + c_{10}) + e_{10r} \\ y_{20r} &= \alpha_2 + \beta_2(\mu_0 + a_{0r} + c_{20}) + e_{20r} \\ &\Downarrow \\ y_{20r} &= \alpha_2 + \frac{\beta_2}{\beta_1}(y_{10r} - \alpha_1 - e_{10r}) \\ &\quad + \beta_2(-c_{10} + c_{20}) + e_{20r} \end{aligned}$$

The random effects have expectation 0, so:

$$E(y_{20}|y_{10}) = \hat{y}_{20} = \alpha_2 + \frac{\beta_2}{\beta_1}(y_{10} - \alpha_1)$$

- ▶ Intercept:  $\alpha_{2|1} = \alpha_2 - \alpha_1 \frac{\beta_2}{\beta_1}$
- ▶ Slope:  $\beta_{2|1} = \frac{\beta_2}{\beta_1}$
- ▶ Invariant under linear transform of  $\mu$ :  
 $a + b\mu_i \rightarrow \tilde{\mu}_i \Rightarrow \alpha_m + \beta_m\mu_i \rightarrow \tilde{\alpha}_m + \tilde{\beta}_m\tilde{\mu}_i$   
 where:  $\tilde{\alpha}_m = \alpha_m - a\beta_m/b$ ,  $\tilde{\beta}_m = \beta_m/b$
- ▶  $\Rightarrow$  the conversion is invariant too:

$$\begin{aligned} \alpha_{2|1} &= \tilde{\alpha}_2 - \tilde{\alpha}_1 \frac{\tilde{\beta}_2}{\tilde{\beta}_1} \\ \beta_{2|1} &= \frac{\tilde{\beta}_2}{\tilde{\beta}_1} \end{aligned}$$

$$y_{20r} = \alpha_2 + \frac{\beta_2}{\beta_1}(y_{10r} - \alpha_1 - e_{10r}) + \beta_2(-c_{10} + c_{20}) + e_{20r}$$

$$\text{var}(\hat{y}_{20}|y_{10}) = \left(\frac{\beta_2}{\beta_1}\right)^2(\beta_1^2\tau_1^2 + \sigma_1^2) + (\beta_2^2\tau_2^2 + \sigma_2^2)$$

The prediction s.d. is:

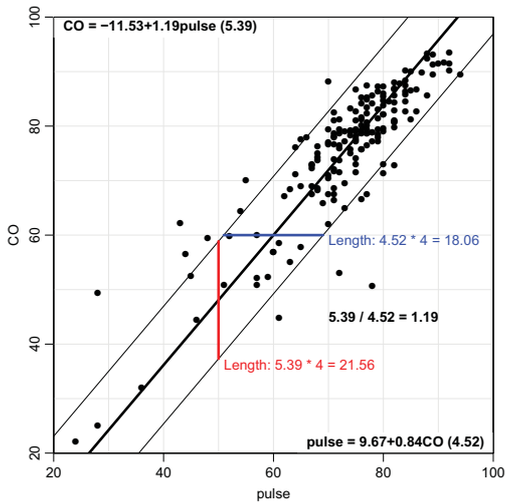
$$\sigma_{2|1} = \sqrt{\left(\frac{\beta_2}{\beta_1}\right)^2(\beta_1^2\tau_1^2 + \sigma_1^2) + (\beta_2^2\tau_2^2 + \sigma_2^2)}$$

If we do the prediction the other way round ( $y_1|y_2$ ) we get the same relationship i.e. a line with the inverse slope,  $\beta_1/\beta_2$ .

The width of the prediction interval in this direction is (by permutation of indices):

$$\begin{aligned} \sigma_{1|2} &= \sqrt{(\beta_1^2\tau_1^2 + \sigma_1^2) + \left(\frac{\beta_1}{\beta_2}\right)^2(\beta_2^2\tau_2^2 + \sigma_2^2)} \\ &= \frac{\beta_1}{\beta_2} \sqrt{\left(\frac{\beta_2}{\beta_1}\right)^2(\beta_1^2\tau_1^2 + \sigma_1^2) + (\beta_2^2\tau_2^2 + \sigma_2^2)} = \frac{\beta_1}{\beta_2}\sigma_{2|1} \end{aligned}$$

i.e. if we draw the prediction limits as straight lines they can be used both ways.



```
> options( width=61 )
> library(MethComp)
> data( ox )
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

```
> system.time( MCox <- MCmcmc( ox, IxR=TRUE ) )
```

Comparison of 2 methods, using 354 measurements on 61 items, with up to 3 replicate measurements, (replicate values are in the set: 1 2 3) (2 \* 61 \* 3 = 366):

```
No. items with measurements on each method:
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

Simulation run of a model with  
 - method by item and item by replicate interaction:  
 - using 4 chains run for 2000 iterations (of which 1000 are burn-in),  
 - monitoring all values of the chain:  
 - giving a posterior sample of 4000 observations.

Initialization and burn-in:  
 Compiling model graph  
 Resolving undeclared variables  
 Allocating nodes  
 Graph Size: 2868

Initializing model

```
Sampling:
user system elapsed
13.94 0.07 14.45
```

```
> ( Mox <- MethComp( MCox ) )
```

```
Conversion between methods:
alpha beta sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 1.963 0.000 0.000 1.963
pulse -11.531 1.192 5.390 -10.519 0.175 4.917
pulse CO 9.671 0.839 4.515 10.519 -0.175 4.911
pulse pulse 0.000 1.000 6.108 0.000 0.000 6.108
```

```
Variance components (sd):
s.d.
Method IxR MxI res
CO 3.861 3.311 1.388
pulse 3.212 2.774 4.319
```

```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Mox, points=TRUE, axlim=c(20,100), xaxs="i", yaxs="i" )
```

```
Relationships between methods:
CO-pulse = -10.52+0.18(CO+pulse)/2 (4.92)
CO = -11.53+1.19pulse (5.39)
pulse = 9.67+0.84CO (4.52)
```

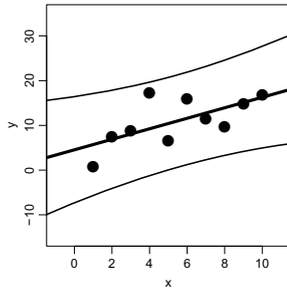
```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Mox, points=TRUE, axlim=c(20,100), xaxs="i", yaxs="i" )
```

```
Relationships between methods:
CO-pulse = -10.52+0.18(CO+pulse)/2 (4.92)
CO = -11.53+1.19pulse (5.39)
pulse = 9.67+0.84CO (4.52)
```

```
> segments( 50, Mox$Conv["CO", "pulse", "alpha"] +
+ Mox$Conv["CO", "pulse", "beta"] * 50 -
+ Mox$Conv["CO", "pulse", "sd.pr"] * 2,
+ 50, Mox$Conv["CO", "pulse", "alpha"] +
+ Mox$Conv["CO", "pulse", "beta"] * 50 +
+ Mox$Conv["CO", "pulse", "sd.pr"] * 2,
+ col="red", lwd=3 )
> text( 51, Mox$Conv["CO", "pulse", "alpha"] +
+ Mox$Conv["CO", "pulse", "beta"] * 50 -
+ Mox$Conv["CO", "pulse", "sd.pr"] * 2.02,
+ paste( "Length:", formatC(Mox$Conv["CO", "pulse", "sd.pr"],
+ format="f", digits=2),
+ "* 4 =", formatC(Mox$Conv["CO", "pulse", "sd.pr"]*4,
+ format="f", digits=2) ),
+ col="red", adj=c(0,1) )
> segments( Mox$Conv["pulse", "CO", "alpha"] +
+ Mox$Conv["pulse", "CO", "beta"] * 60 -
+ Mox$Conv["pulse", "CO", "sd.pr"] * 2, 60,
+ Mox$Conv["pulse", "CO", "alpha"] +
+ Mox$Conv["pulse", "CO", "beta"] * 60 +
+ Mox$Conv["pulse", "CO", "sd.pr"] * 2, 60,
+ col="blue", lwd=3 )
> text( Mox$Conv["pulse", "CO", "alpha"] +
+ Mox$Conv["pulse", "CO", "beta"] * 60 +
```

```
+ Mox$Conv["pulse", "CO", "sd.pr"] * 2 + 1, 60,
+ paste( "Length:", formatC(Mox$Conv["pulse", "CO", "sd.pr"],
+ format="f", digits=2),
+ "* 4 =", formatC(Mox$Conv["pulse", "CO", "sd.pr"]*4,
+ format="f", digits=2) ),
+ col="blue", adj=c(0,1) )
> text( 70, 45, paste( formatC( Mox$Conv["CO", "pulse", "sd.pr"],
+ format="f", digits=2 ), "/",
+ formatC( Mox$Conv["pulse", "CO", "sd.pr"],
+ format="f", digits=2 ), "=",
+ formatC( Mox$Conv["CO", "pulse", "beta"],
+ format="f", digits=2 ) ),
+ adj=0, font=2 )
```

## What happened to the curvature?



Usually the prediction limits are curved:

$$\hat{y}|x \pm 1.96 \times \hat{\sigma} \sqrt{1 + x'x}$$

In our prediction we have ignored the last term ( $x'x$ ), i.e. effectively assuming that there is no estimation error on  $\alpha_{2|1}$  and  $\beta_{2|1}$ .

```
> set.seed(17676)
> par(mar=c(3,3,1,1),mgp=c(3,1,0)/1.6)
> x <- 1:10
> y <- 3 + 1.6*x + rnorm(x,,6)
> m0 <- lm(y~x)
> plot(y~x,pch=16,ylim=c(-15,35),xlim=c(-1,11),cex=2)
> nx <- seq(-3,13,,200)
> matlines( nx, predict( m0, interval="pred", newdata=data.frame(x=nx)),
+          lwd=c(4,2,2), col="black", lty=1 )

> # The same but now with 100 points
> set.seed(17676)
> par(mar=c(3,3,1,1),mgp=c(3,1,0)/1.6)
> x <- seq(1,10,,100)
> y <- 3 + 1.6*x + rnorm(x,,6)
> m0 <- lm(y~x)
> plot(y~x,pch=16,ylim=c(-15,35),xlim=c(-1,11),cex=0.7)
> nx <- seq(-3,13,,200)
> matlines( nx, predict( m0, interval="pred", newdata=data.frame(x=nx)),
+          lwd=c(4,2,2), col="black", lty=1 )
```

## Comparing to a gold standard

- ▶ The prediction s.d. is:

$$\sigma_{2|1} = \sqrt{\left(\frac{\beta_2}{\beta_1}\right)^2 (\beta_1^2 \tau_1^2 + \sigma_1^2) + (\beta_2^2 \tau_2^2 + \sigma_2^2)}$$

- ▶ If method 1 is the gold standard (no error), i.e. **assumed**:  $\tau_1 = \sigma_1 = 0$
- ▶ Estimate relationship by regressing  $y_2$  on  $y_1$ , deriving  $\tau_2$  and  $\sigma_2$  — standard linear regression.
- ▶ Prediction of  $y_1$  (what would the gold standard give?):
- ▶ Limits for  $y_2|y_1$ , but used the other way.

## Implementation in BUGS/JAGS

### Bendix Carstensen

SAoMCS  
19–20 March 2014  
Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(BUGS-impl)

## Implementation in BUGS

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

Non-linear hierarchical model:

- ▶ The model is *symmetrical* in methods.
- ▶ Mean is overparametrized.
- ▶ Choose a prior (and hence posterior!) for the  $\mu_s$  with finite support.
- ▶ Keeps the chains nicely in place.

This is the philosophy in the function `MCmcmc`.

## Results from fitting the model

The posterior dist'n of  $(\alpha_m, \beta_m, \mu_i)$  is singular.

But the relevant translation quantities **are** identifiable:

$$\alpha_{2|1} = \alpha_2 - \alpha_1 \beta_2 / \beta_1$$

$$\beta_{2|1} = \beta_2 / \beta_1$$

— so are the variance components.

Posterior medians used to devise prediction equations with limits.

## Implemented model:

$$y_{mir} = \alpha_m + \beta_m(\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

- ▶ Replicates required in data.
- ▶ **JAGS** (or R2WinBUGS or BRUGS) is required.
- ▶ Dataframe with variables meth, item, repl and y (a Meth object)
- ▶ The function MCmcmc writes a BUGS-program, initial values and data to files.
- ▶ Runs **JAGS** and sucks results back in to **R**, and gives a nice overview of the conversion equations.

```
> options( width=61 )
> library(MethComp)
> data( ox )
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

```
> system.time( MCox <- MCmcmc( ox, IxR=TRUE, n.iter=10000 ) )
```

Comparison of 2 methods, using 354 measurements on 61 items, with up to 3 replicate measurements, (replicate values are in the set: 1 2 3 ) ( 2 \* 61 \* 3 = 366 ):

```
No. items with measurements on each method:
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

Simulation run of a model with  
- method by item and item by replicate interaction:  
- using 4 chains run for 10000 iterations (of which 5000 are burn-in),  
- monitoring every 5 values of the chain:  
- giving a posterior sample of 4000 observations.

Initialization and burn-in:  
Compiling model graph  
Resolving undeclared variables  
Allocating nodes  
Graph Size: 2868

Initializing model

```
Sampling:
user system elapsed
69.49 0.09 69.93
```

```
> MCox
```

```
Conversion between methods:
alpha beta sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 2.316 0.000 0.000 2.316
pulse -6.933 1.129 5.124 -6.513 0.121 4.813
pulse CO 6.140 0.886 4.544 6.513 -0.121 4.819
pulse pulse 0.000 1.000 6.050 0.000 0.000 6.050
```

```
Variance components (sd):
s.d.
Method IxR MxI res
CO 3.824 3.155 1.638
pulse 3.377 2.798 4.278
```

```
Variance components with 95 % cred.int.:
method CO pulse
qnt 50% 2.5% 97.5% 50% 2.5% 97.5%
SD
IxR 3.824 3.074 4.546 3.377 2.741 4.054
MxI 3.155 2.323 4.150 2.798 2.024 3.760
res 1.638 0.298 2.697 4.278 3.631 5.005
tot 5.260 4.632 6.037 6.169 5.541 6.841
```

```
Mean parameters with 95 % cred.int.:
50% 2.5% 97.5% P(>0/1)
alpha[pulse.CO] 6.144 -2.900 13.632 0.918
alpha[CO.pulse] -6.928 -17.274 2.921 0.082
```

```
beta[pulse.CO] 0.886 0.788 1.003 0.028
beta[CO.pulse] 1.129 0.997 1.270 0.972
```

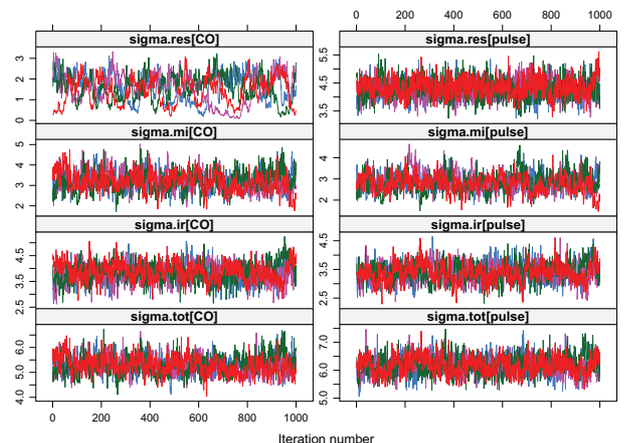
Note that intercepts in conversion formulae are adjusted to get conversion formulae that represent the same line both ways, and hence the median intercepts in the posterior do not agree exactly with those given in the conversion formulae.

```
> MethComp( MCox )
```

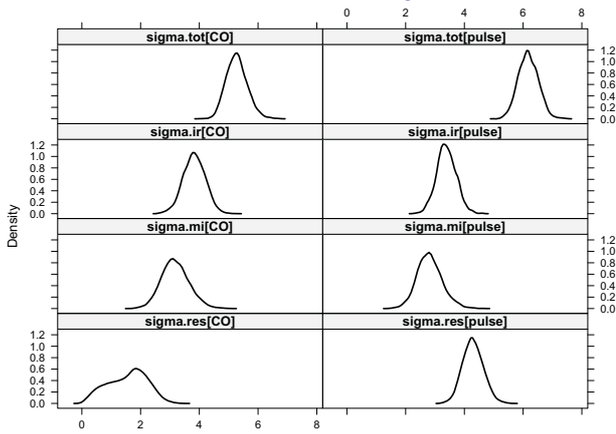
```
Conversion between methods:
alpha beta sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 2.316 0.000 0.000 2.316
pulse -6.933 1.129 5.124 -6.513 0.121 4.813
pulse CO 6.140 0.886 4.544 6.513 -0.121 4.819
pulse pulse 0.000 1.000 6.050 0.000 0.000 6.050
```

```
Variance components (sd):
s.d.
Method IxR MxI res
CO 3.824 3.155 1.638
pulse 3.377 2.798 4.278
```

## Traces of the chains



## Posteriors for variance components



Implementation in BUGS/JAGS (BUGS-impl)

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```
> trace.MCmcmc( MCox )
> post.MCmcmc( MCox )
> post.MCmcmc( MCox, check=FALSE )
```

Implementation in BUGS/JAGS (BUGS-impl)

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```
> data( sbp )
> sbp <- Meth( sbp )
```

The following variables from the dataframe "sbp" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
```

Method	#Replicates	3	#Items	#Obs:	765	Values:	min	med	max
J	85	85	255	74	120	228			
R	85	85	255	76	120	226			
S	85	85	255	77	135	228			

```
> system.time( MCBp <- MCmcmc( sbp, IxR=TRUE, n.iter=10000 ) )
```

Implementation in BUGS/JAGS (BUGS-impl)

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Comparison of 3 methods, using 765 measurements on 85 items, with up to 3 replicate measurements, (replicate values are in the set: 1 2 3) ( 3 \* 85 \* 3 = 765 ):

No. items with measurements on each method:

Method	#Replicates	3	#Items	#Obs:	765	Values:	min	med	max
J	85	85	255	74	120	228			
R	85	85	255	76	120	226			
S	85	85	255	77	135	228			

Simulation run of a model with  
 - method by item and item by replicate interaction:  
 - using 4 chains run for 10000 iterations  
 (of which 5000 are burn-in),  
 - monitoring every 5 values of the chain:  
 - giving a posterior sample of 4000 observations.

Initialization and burn-in:  
 Compiling model graph  
 Resolving undeclared variables  
 Allocating nodes  
 Graph Size: 5982

Initializing model

Sampling:

Implementation in BUGS/JAGS (BUGS-impl)

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```
user system elapsed
179.57 0.22 180.22
```

```
> MCBp
```

Conversion between methods:

To: From:	alpha	beta	sd.pr	in(t-f)	sl(t-f)	sd(t-f)
J J	0.000	1.000	2.173	0.000	0.000	2.173
R J	-1.143	1.010	2.293	-1.137	0.010	2.282
S J	-50.444	1.246	24.899	-44.929	0.219	22.176
R R	1.132	0.990	2.271	1.137	-0.010	2.282
S R	0.000	1.000	2.374	0.000	0.000	2.374
R S	-48.832	1.234	24.689	-43.720	0.209	22.104
J S	40.501	0.803	20.008	44.929	-0.219	22.196
R S	39.577	0.810	20.019	43.720	-0.209	22.114
S S	0.000	1.000	28.242	0.000	0.000	28.242

Variance components (sd):

Method	IxR	MxI	res
J	5.992	0.316	1.482
R	5.935	0.184	1.658
S	4.804	17.860	8.923

Variance components with 95 % cred.int.:

method	J	R	S						
qnt	50%	2.5%	97.5%	50%	2.5%	97.5%	50%	2.5%	97.5%

Implementation in BUGS/JAGS (BUGS-impl)

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SD	IxR	MxI	res	tot
5.992	5.379	6.719	5.935	5.331
6.650	4.804	4.082	5.698	
0.316	0.009	0.840	0.184	0.025
0.635	17.860	15.163	21.274	
1.482	0.765	2.023	1.658	1.016
2.125	8.923	8.011	10.024	
6.191	5.573	6.897	6.176	5.570
6.866	20.551	18.270	23.607	

Mean parameters with 95 % cred.int.:

alpha	beta	sd	in	sl	sd
alpha[R.J]	1.132	-0.209	2.332	0.952	
alpha[S.J]	40.477	27.049	53.826	1.000	
alpha[J.R]	-1.143	-2.375	0.209	0.048	
alpha[S.R]	39.561	25.983	52.959	1.000	
alpha[J.S]	-50.474	-76.857	-30.134	0.000	
alpha[R.S]	-48.852	-74.837	-28.416	0.000	
beta[R.J]	0.990	0.981	1.001	0.033	
beta[S.J]	0.803	0.699	0.905	0.000	
beta[J.R]	1.010	0.999	1.019	0.967	
beta[S.R]	0.810	0.706	0.913	0.000	
beta[J.S]	1.246	1.105	1.430	1.000	
beta[R.S]	1.234	1.095	1.415	1.000	

Note that intercepts in conversion formulae are adjusted to get conversion formulae that represent the same line both ways, and hence the median intercepts in the posterior do not agree exactly with those given in the conversion formulae.

```
> MethComp( MCBp )
```

Implementation in BUGS/JAGS (BUGS-impl)

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```

Conversion between methods:
      alpha      beta      sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
J  J      0.000      1.000      2.173      0.000      0.000      2.173
   R     -1.143      1.010      2.293     -1.137      0.010      2.282
   S    -50.444      1.246     24.899    -44.929      0.219     22.176
R  J      1.132      0.990      2.271      1.137     -0.010      2.282
   R      0.000      1.000      2.374      0.000      0.000      2.374
   S    -48.832      1.234     24.689    -43.720      0.209     22.104
S  J     40.501      0.803     20.008     44.929     -0.219     22.196
   R     39.577      0.810     20.019     43.720     -0.209     22.114
   S      0.000      1.000     28.242      0.000      0.000     28.242

Variance components (sd):
s.d.
Method  IxR    MxI    res
J  5.992  0.316  1.482
R  5.935  0.184  1.658
S  4.804 17.860  8.923

```

# Alternating regressions

Bendix Carstensen

SAoMCS  
 19–20 March 2014  
 Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Alt-reg)

## Alternating random effects regression

Carstensen [3] proposed a ridiculously complicated approach to fit the model

$$y_{mir} = \alpha_m + \beta_m \mu_i + c_{mi} + e_{mir}$$

based in the observation that:

- ▶ For fixed  $\mu$  the model is a linear mixed model.
- ▶ For fixed  $(\alpha, \beta)$  it is a regression through 0.

This has been improved by Carstensen in [4]

## Alternating random effects regression

The correctly formulated version of the slightly more general model:

$$y_{mir} = \alpha_m + \beta_m (\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

- ▶ For fixed  $\zeta_{mir} = \mu_i + a_{ir} + c_{mi}$  the model is a linear model, with residual variances different between methods.
- ▶ For fixed  $(\alpha, \beta)$  scaled responses  $y$  follow a standard mixed model:

$$\frac{y_{mir} - \alpha_m}{\beta_m} = \mu_i + a_{ir} + c_{mi} + e_{mir} / \beta_m$$

## Estimation algorithm

$$y_{mir} = \alpha_m + \beta_m (\mu_i + a_{ir} + c_{mi}) + e_{mir}$$

1. Start with  $\zeta_{mir} = \bar{y}_{mi}$ .
2. Estimate  $(\alpha_m, \beta_m)$ .
3. Compute the scaled responses and fit the random effects model.
4. Use the estimated  $\mu_i$ s, and BLUPs of  $a_{ir}$  and  $c_{mi}$  to update  $\zeta_{mir}$ .
5. Check convergence in terms of identifiable parameters.

## The residual variances

- ▶ The variance components are estimated in the model for the scaled response.
- ▶ The estimation of parameters  $(\alpha_m, \beta_m)$  are not taken into account in the calculation of the residual variance d.f.
- ▶ Hence the residual variances must be corrected *post hoc*.
- ▶ This machinery is implemented in the function AltReg in the MethComp package.

```
> options( width=100 )
> library(MethComp)
> data( ox )
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

```
> system.time( AR.ox <- AltReg( ox, linked=T, trace=T ) )
```

Alternating regressions

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```
iteration 1 criterion: 1
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO 0.911 0.988 1.861 74.419 74.417 1.000 0.974 3.371 3.502 2.292
pulse -1.039 1.014 1.860 74.422 74.419 1.027 1.000 3.460 3.595 3.958

iteration 2 criterion: 0.07508045
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -0.714 1.011 1.255 74.419 74.956 1.00 0.99 3.399 3.311 2.251
pulse -2.006 1.022 3.020 73.878 74.419 1.01 1.00 3.433 3.344 3.981

iteration 3 criterion: 0.0594666
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -2.363 1.035 1.215 74.419 75.433 1.000 1.005 3.425 3.173 2.211
pulse -2.971 1.030 3.082 73.412 74.419 0.995 1.000 3.407 3.156 4.002

iteration 4 criterion: 0.04281372
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -4.019 1.058 1.177 74.419 75.831 1.000 1.019 3.447 3.084 2.175
pulse -3.963 1.039 3.139 73.034 74.419 0.982 1.000 3.384 3.027 4.021

iteration 5 criterion: 0.02856943
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -5.668 1.081 1.143 74.419 76.145 1.000 1.03 3.466 3.031 2.145
pulse -5.009 1.049 3.186 72.744 74.419 0.971 1.00 3.365 2.943 4.036

iteration 6 criterion: 0.01820552
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -7.307 1.103 1.113 74.419 76.382 1.000 1.039 3.482 3.003 2.121
pulse -6.124 1.062 3.223 72.530 74.419 0.962 1.000 3.351 2.890 4.048
```

Alternating regressions

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```
iteration 7 criterion: 0.01140264
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -8.936 1.126 1.09 74.419 76.556 1.000 1.046 3.493 2.989 2.102
pulse -7.314 1.076 3.25 72.377 74.419 0.956 1.000 3.340 2.858 4.057

iteration 8 criterion: 0.007169339
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -10.562 1.148 1.071 74.419 76.680 1.000 1.051 3.502 2.982 2.08
pulse -8.576 1.092 3.269 72.269 74.419 0.951 1.000 3.331 2.837 4.06

iteration 9 criterion: 0.005074459
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -12.190 1.169 1.057 74.419 76.768 1.000 1.055 3.508 2.980 2.07
pulse -9.904 1.109 3.282 72.193 74.419 0.948 1.000 3.325 2.824 4.06

iteration 10 criterion: 0.003705422
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -13.826 1.191 1.047 74.419 76.830 1.000 1.058 3.513 2.978 2.06
pulse -11.290 1.126 3.292 72.140 74.419 0.945 1.000 3.321 2.816 4.07

iteration 11 criterion: 0.002686236
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -15.476 1.213 1.039 74.419 76.873 1.000 1.06 3.516 2.978 2.06
pulse -12.727 1.145 3.298 72.104 74.419 0.944 1.00 3.318 2.810 4.07

iteration 12 criterion: 0.001930191
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -17.144 1.236 1.034 74.419 76.903 1.000 1.061 3.518 2.978 2.06
```

Alternating regressions

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```
pulse -14.211 1.165 3.303 72.079 74.419 0.942 1.000 3.315 2.807 4.07

iteration 13 criterion: 0.001381194
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -18.834 1.258 1.030 74.419 76.924 1.000 1.062 3.520 2.978 2.05
pulse -15.736 1.185 3.306 72.061 74.419 0.941 1.000 3.314 2.804 4.07

iteration 14 criterion: 0.0009863462
alpha beta sigma Intercept: CO pulse Slope: CO pulse IxR MxI res
CO -20.548 1.281 1.027 74.419 76.938 1.000 1.063 3.521 2.978 2.05
pulse -17.301 1.205 3.308 72.049 74.419 0.941 1.000 3.313 2.802 4.07

AltReg converged after 14 iterations
Last convergence criterion was 0.0009863462
user system elapsed
12.71 0.03 12.78

> AR.ox
```

Alternating regressions

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```
Conversion between methods:
alpha beta sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 2.906 0.000 0.000 2.906
pulse -2.159 1.063 6.385 -2.093 0.061 6.190
pulse CO 2.031 0.941 6.007 2.093 -0.061 6.190
pulse pulse 0.000 1.000 5.769 0.000 0.000 5.769
```

```
Variance components (sd):
s.d.
Method IxR MxI res
CO 3.521 2.978 2.055
pulse 3.313 2.802 4.079
```

Alternating regressions

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# Transformation of data

**Bendix Carstensen**

SAoMCS  
19–20 March 2014  
Haukeland University Hospital, Bergen, Norway  
<http://BendixCarstensen.com/MethComp/Courses/Bergen.2014>

(Transform)

## If variances are not constant

A transformation might help:

```
> library( MethComp )
> data( ox )
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
```

```
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

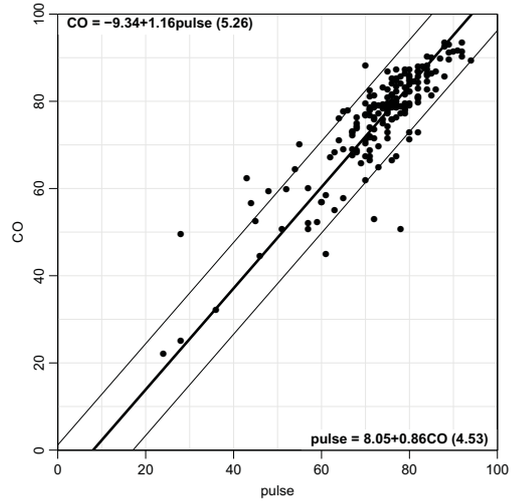
```
> DA.reg(ox)
```

```
Conversion between methods:
      alpha  beta  sd.pr beta=1 in(t-f) sl(t-f) sd(t-f) in(sd) sl(sd)
To: From:
CO CO 0.000 1.000 NA NA 0.000 0.000 NA NA NA
pulse -1.977 1.061 6.342 0.142 -1.919 0.059 6.155 17.602 -0.162
pulse CO 1.864 0.943 5.979 0.142 1.919 -0.059 -6.155 17.602 -0.162
pulse pulse 0.000 1.000 NA NA 0.000 0.000 NA NA NA
```

```
> round( ftable(DA.reg(ox)$Conv[,,-(1:4)]), 3 )
```

Transformation of data (Transform)

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Transformation of data (Transform)

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```
> library(MethComp)
> data( ox )
> ox <- Meth( ox )
```

The following variables from the dataframe "ox" are used as the Meth variables:

```
meth: meth
item: item
repl: repl
y: y
```

```
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

```
> system.time( MCox <- MCmcmc( ox, IxR=TRUE ) )
```

Transformation of data (Transform)

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Comparison of 2 methods, using 354 measurements on 61 items, with up to 3 replicate measurements, (replicate values are in the set: 1 2 3) ( 2 \* 61 \* 3 = 366 ):

No. items with measurements on each method:

```
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 22.2 78.6 93.5
pulse 1 4 56 61 177 24.0 75.0 94.0
```

Simulation run of a model with

- method by item and item by replicate interaction:
- using 4 chains run for 2000 iterations (of which 1000 are burn-in),
- monitoring all values of the chain:
- giving a posterior sample of 4000 observations.

Initialization and burn-in:

```
Compiling model graph
  Resolving undeclared variables
  Allocating nodes
  Graph Size: 2868
```

Initializing model

```
Sampling:
  user system elapsed
16.27 0.05 16.39
```

Transformation of data (Transform)

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```
> ( Mox <- MethComp( MCox ) )
```

```
Conversion between methods:
      alpha  beta  sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 2.553 0.000 0.000 2.553
pulse -9.341 1.161 5.263 -8.645 0.149 4.871
pulse CO 8.045 0.861 4.526 8.645 -0.149 4.864
pulse pulse 0.000 1.000 5.985 0.000 0.000 5.985
```

```
Variance components (sd):
s.d.
```

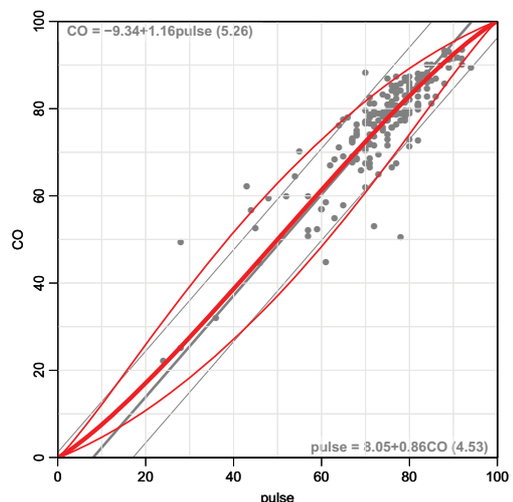
```
Method IxR MxI res
CO 3.706 3.089 1.805
pulse 3.173 2.647 4.232
```

```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Mox, points=TRUE, axlim=c(0,100), xaxs="i", yaxs="i" )
```

```
Relationships between methods:
CO-pulse = -8.64+0.15(CO+pulse)/2 (4.87)
CO = -9.34+1.16pulse (5.26)
pulse = 8.05+0.86CO (4.53)
```

Transformation of data (Transform)

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Transformation of data (Transform)

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## Using the Transform argument I

```
> system.time( MCox <- MCmcmc( ox, IxR=TRUE, Transform="pctlogit" ) )
```

Comparison of 2 methods, using 354 measurements on 61 items, with up to 3 replicate measurements, (replicate values are in the set: 1 2 3) (2 \* 61 \* 3 = 366):

```
No. items with measurements on each method:
#Replicates
Method 1 2 3 #Items #Obs: 354 Values: min med max
CO 1 4 56 61 177 -1.254049 1.300981 2.666159
pulse 1 4 56 61 177 -1.152680 1.098612 2.751535
```

Simulation run of a model with  
 - method by item and item by replicate interaction:  
 - using 4 chains run for 2000 iterations (of which 1000 are burn-in),  
 - monitoring all values of the chain:  
 - giving a posterior sample of 4000 observations.

```
Initialization and burn-in:
Compiling model graph
Resolving undeclared variables
Allocating nodes
Graph Size: 2869
```

Transformation of data (Transform)

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## Using the Transform argument II

Initializing model

```
Sampling:
user system elapsed
16.12 0.00 16.19
```

```
> ( Tox <- MethComp( MCox ) )
```

Note: Response transformed by: function (p) log(p/(100 - p))

```
Conversion between methods:
alpha beta sd.pr in(t-f) sl(t-f) sd(t-f)
To: From:
CO CO 0.000 1.000 0.184 0.000 0.000 0.184
pulse CO 0.000 1.141 0.264 0.000 0.132 0.247
pulse pulse 0.000 0.876 0.232 0.000 -0.132 0.247
pulse CO 0.000 1.000 0.283 0.000 0.000 0.283
```

```
Variance components (sd):
s.d.
Method IxR MxI res
CO 0.257 0.176 0.13
pulse 0.224 0.154 0.20
```

Transformation of data (Transform)

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## Using the Transform argument III

```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Mox, points=TRUE, xlim=c(0,100), xaxs="i", yaxs="i",
+ col.lines=gray(0.5), col.points=gray(0.5) )
```

Relationships between methods:  
 CO-pulse = -8.64+0.15(CO+pulse)/2 (4.87)  
 CO = -9.34+1.16pulse (5.26)  
 pulse = 8.05+0.86CO (4.53)

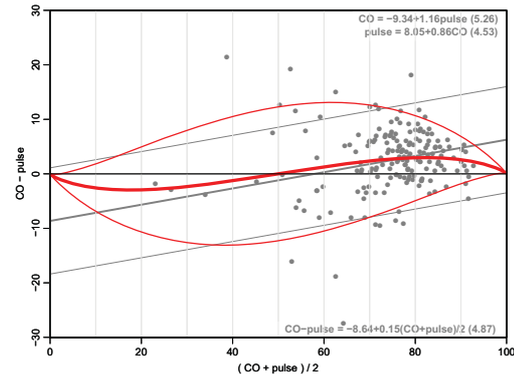
```
> par( new=TRUE )
> plot( Tox, points=FALSE, xlim=c(0,100), xaxs="i", yaxs="i",
+ col.lines="red", lwd=c(5,2,2) )
```

Transformation of data (Transform)

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## Transformation to a Bland-Altman plot

Just convert to the differences versus the averages:



Transformation of data (Transform)

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```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Mox, pl.type="BA", points=TRUE, xlim=c(0,100), diflim=c(-30,30),
+ xaxs="i", yaxs="i" )
```

Relationships between methods:  
 CO-pulse = -8.64+0.15(CO+pulse)/2 (4.87)  
 CO = -9.34+1.16pulse (5.26)  
 pulse = 8.05+0.86CO (4.53)

```
> abline( h=0 )
```

```
> par( mar=c(3,3,1,1), mgp=c(3,1,0)/1.6 )
> plot( Tox, pl.type="BA", points=TRUE, xlim=c(0,100), diflim=c(-30,30),
+ xaxs="i", yaxs="i", col.lines=gray(0.5), col.points=gray(0.5) )
```

Relationships between methods:  
 CO-pulse = -8.64+0.15(CO+pulse)/2 (4.87)  
 CO = -9.34+1.16pulse (5.26)  
 pulse = 8.05+0.86CO (4.53)

```
> abline( h=0 )
> par( new=TRUE )
> plot( Tox, pl.type="BA", points=FALSE, xlim=c(0,100), diflim=c(-30,30),
+ xaxs="i", yaxs="i", col.lines="red", lwd=c(5,2,2) )
> abline( h=0 )
```

Transformation of data (Transform)

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Transformation of data (Transform)

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