Interindividual differences in response to treatment: fact, fiction and erroneous analyses

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1 Introduction

An example is presented demonstrating why separately analysing treatment and control arms from a randomised controlled trial (RCT) and testing for an association of a covariate (biomarker) with observed 'responders' and 'non responders' in each arm is erroneous, wasteful and a seriously misleading analysis. This analysis scenario is never advisable and demonstrates a misunderstanding of the basics of randomised clinical trials.

Analysts working in the personalised medicine field need to be knowledgeable of variance components analysis, understand the counterfactual premise that underpins RCTs, that is, what would have happened to the same patients in the treatment arm had they been in the control arm and be knowledgeable of appropriate statistical analyses for RCT data. This basic counterfactual tenet indicates that responders and non responders cannot be identified by examining trial arms separately.

It is advisable to first estimate the true individual response using parallel information from both arms, and if present, judge if it is clinically relevant. If so, covariates that potentially modify or mediate response can be included in the statistical model of the two trial arms.

2 Simulate a RCT

Simulate a randomised controlled trial with a baseline and follow up measurement and a constant treatment effect. A constant treatment effect means **everybody** in the treatment arm responds by a **constant** amount. **There are NO interindividual differences!** In the control arm the true baseline and true follow up are the same.

```
n <- 5000
noise <- 5
                     # add noise (within person var & meas. error) to the baseline & foll. up
beta.treatment <- -5 # all trt'd subjects exp same trt effect, so no resp - non responders!!
# beta.treatment <- runif(n,-20,-5) # subjects vary in response to treatment
pop_mu <- 79
                     # population mean
pop_sd <- 10
                     # between person SD
ur.eligible <- 89
                     # eliqibility criteria for trial
y.Otrue <- rnorm(n, pop_mu, pop_sd)
                                                      # true baseline
y.Oobserved <- y.Otrue + rnorm(n, 0, 1*noise)
                                                      # observed baseline
eligible <- ifelse(y.Oobserved > ur.eligible, 1, 0) # 1sd above norm eliqible for trial
treat \leftarrow 1*(runif(n)<.5)
                                                      # random treatment allocation
y.1true <- y.0true + (treat*beta.treatment)</pre>
                                                      # true follow up, treated only respond
y.1observed <- y.1true + rnorm(n, 0, 1*noise)
                                                      # observed follow up, noise added
delta.observed <- y.1observed - y.0observed
d <- data.frame(y.0true, y.0observed, eligible, treat , beta.treatment,</pre>
```

```
y.1true, y.1observed, delta.observed)

# prob that a member of pop observed baseline is eligible
# pnorm(ur.eligible, mean= pop_mu, sd=sqrt(pop_sd^2 + noise^2))
# 1- pnorm( (pop_mu - ur.eligible) / sqrt(pop_sd^2+noise^2) ) # z score calc.

trial <- d[d$eligible==1,] # select the trial subjects</pre>
```

3 First rows of trial data

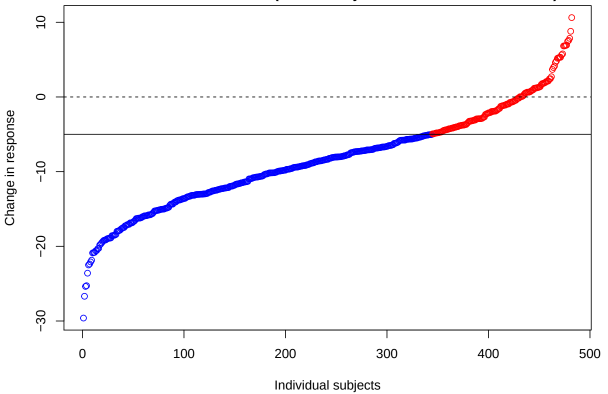
'y.0true' is the true baseline for each subject. Yet it is not observed, 'y.0observed' is that which is observed and includes measurement error. 'y.1true' is 'y.0true'- 'beta.treatment', the treatment effect , for treated subjects only, otherwise equal to the baseline. But again this is not observed as it is measured with error and the estimate is recorded in variable 'y.1observed'.

y.0true	y.0observed	treat	beta.treatment	y.1true	y.1observed	delta.observed
97.67	96.34	0	-5	97.67	99.55	3.22
92.51	105.96	0	-5	92.51	102.02	-3.94
88.59	93.59	0	-5	88.59	90.94	-2.65
96.09	95.78	0	-5	96.09	93.64	-2.13
98.58	95.71	1	-5	93.58	90.78	-4.93
103.26	103.87	0	-5	103.26	104.80	0.93
89.63	96.24	1	-5	84.63	82.93	-13.31
90.62	89.09	0	-5	90.62	88.82	-0.27
90.23	91.75	1	-5	85.23	77.83	-13.92
92.72	89.93	0	-5	92.72	90.30	0.36
91.15	89.26	0	-5	91.15	90.86	1.60
92.29	97.79	1	-5	87.29	78.65	-19.14
92.19	96.44	1	-5	87.19	94.22	-2.21
96.90	89.54	1	-5	91.90	89.87	0.33
96.26	100.01	0	-5	96.26	94.98	-5.03
87.53	89.29	1	-5	82.53	79.13	-10.15
91.16	91.60	1	-5	86.16	85.98	-5.63
89.50	89.68	0	-5	89.50	79.76	-9.92
92.51	89.76	1	-5	87.51	81.66	-8.10
88.34	90.38	0	-5	88.34	79.35	-11.03

4 Focus on the intervention arm only - not recommended!

The subjects in blue were observed to respond only, those in red observed not to respond.

Individual changes in response in treated arm Suggested individual differences due entirely to regression to the mean and random error (within subject and measurement error)



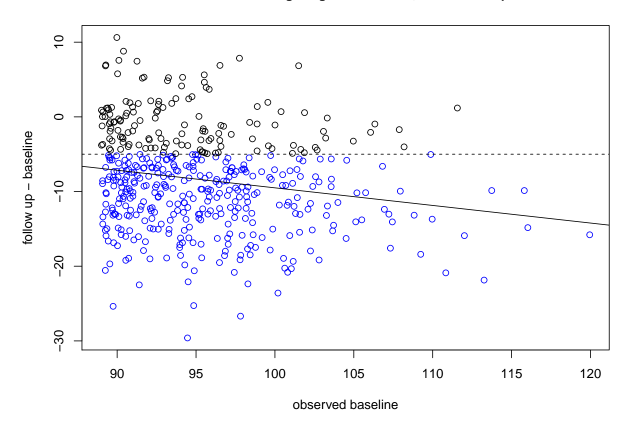
```
# this many were not observed to have reduced response by more than 5
# wrongly labelled as 'non responders'
mean(foo > -5)*length(foo) # shown in red
```

[1] 138

5 Treatment arm only

Observed responders in blue. But **EVERYBODY** responded to the drug **EQUALLY**! Apparent individual difference is due **ENTIRELY** to random within subject error, measurement error and regression to the mean.

Treatment arm: Individual changes against baseline, observed responders in blue



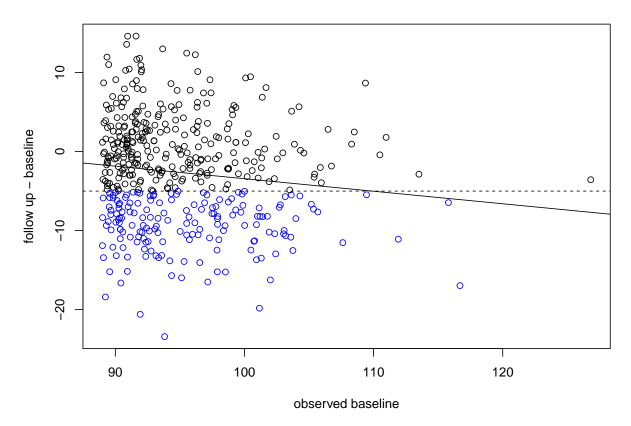
Pearson's product-moment correlation

```
data: diff and y.Oobserved
t = -4.2222, df = 480, p-value = 2.895e-05
alternative hypothesis: true correlation is not equal to 0
95 percent confidence interval:
   -0.2739193 -0.1016379
sample estimates:
        cor
   -0.1892345
```

6 Control arm only

Observed responders in blue. But in truth **NO ONE** responded, apparent individual difference is due **ENTIRELY** to random within subject error, measurement error and regression to the mean.

Control arm: Individual changes against baseline, observed responders in blue



Pearson's product-moment correlation

```
data: diff and y.Oobserved
t = -2.6945, df = 450, p-value = 0.007314
alternative hypothesis: true correlation is not equal to 0
95 percent confidence interval:
   -0.21573253 -0.03416983
sample estimates:
        cor
   -0.1260062
```

7 Analyse the trial correctly. Estimate the treatment effect adjusting for baseline

```
Call:
lm(formula = y.1observed ~ y.0observed + treat, data = trial)
Residuals:
    Min
              1Q Median
                               3Q
-21.3772 -4.4896 0.2056 4.3897 17.9677
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 16.21241 3.85932 4.201 2.91e-05 ***
                      0.04052 19.782 < 2e-16 ***
y.0observed 0.80156
          -5.69619
                      0.42672 -13.349 < 2e-16 ***
treat
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 6.514 on 931 degrees of freedom
Multiple R-squared: 0.3726,
                            Adjusted R-squared: 0.3713
F-statistic: 276.5 on 2 and 931 DF, p-value: < 2.2e-16
               2.5 %
                        97.5 %
(Intercept) 8.638430 23.7863917
y.Oobserved 0.722040 0.8810777
treat -6.533634 -4.8587525
```

8 Look before leaping

Calculate the difference in SDs of the changes between the intervention and control arms, confidence interval for the sd for changes in each arm

```
alpha <- 0.05

x <- trial[trial$treat %in% 0,"delta.observed"]

lstar <- qchisq(alpha/2, df= length(x)-1)
 rstar <- qchisq(1-alpha/2, df= length(x)-1)

up <- sqrt((length(x)-1)*var(x)/(lstar))
 lo <- sqrt((length(x)-1)*var(x)/(rstar))
 pe <- sqrt(var(x))

# ctrl arm estimate with 95% CI
 print(c(pe, lo, up), digits=3)</pre>
```

[1] 6.63 6.22 7.09

```
x1 <- trial[trial$treat %in% 1,"delta.observed"]

lstar <- qchisq(alpha/2, df= length(x1)-1)
  rstar <- qchisq(1-alpha/2, df= length(x1)-1)

up <- sqrt((length(x1)-1)*var(x1)/(lstar))
  lo <- sqrt((length(x1)-1)*var(x1)/(rstar))
  pe <- sqrt(var(x1))

# trt arm estimate with 95% CI
  print(c(pe, lo, up), digits=3)</pre>
```

[1] 6.56 6.17 7.00

9 Typical true interindividual variation in response. Adjust for the influence of biological variation and measurement error (removal of noise).

The linear mixed model p-value provides evidence the SD for true interindividual variation is consistent with zero, as it should be, given that the true magnitude of response in the simulation is constant for all subjects randomised to the treated arm. This result provides information that true individual response differences are negligible and analysis of interindividual response is unwarranted.

```
# True individual response to the intervention sqrt(sd(x1)^2-sd(x)^2) # can be -ve if more var in control group
```

[1] NaN

```
Linear mixed-effects model fit by REML
  Data: trial
  Log-restricted-likelihood: -3076.776
  Fixed: delta.observed ~ treat + y.Oobserved
                  treat y.Oobserved
(Intercept)
 16.2870135 -5.6959245 -0.1992269
Random effects:
Formula: ~1 | treat
        (Intercept) Residual
          0.8122445 6.582047
StdDev:
Variance function:
Structure: Different standard deviations per stratum
Formula: ~1 | treat
Parameter estimates:
1.0000000 0.9797944
Number of Observations: 934
Number of Groups: 2
```

anova(m1,m0) # are the trt ctr interindividual variation in response different?

```
Model df AIC BIC logLik Test L.Ratio p-value m1 1 6 6165.551 6194.569 -3076.776 m0 2 5 6163.745 6187.926 -3076.872 1 vs 2 0.1936813 0.6599
```

```
c.grp <- m1$sigma
t.grp <- coef(m1$modelStruct$varStruct, uncons = FALSE)[[1]]*m1$sigma

# true individual response to the intervention estimate
sqrt(t.grp^2 - c.grp^2)</pre>
```

[1] NaN

```
# truth
sd(beta.treatment )
```

[1] NA

10 References

11 Computing Environment

```
R version 3.2.2 (2015-08-14)
Platform: x86_64-w64-mingw32/x64 (64-bit)
Running under: Windows 8 x64 (build 9200)
locale:
[1] LC_COLLATE=English_United Kingdom.1252
[2] LC_CTYPE=English_United Kingdom.1252
[3] LC_MONETARY=English_United Kingdom.1252
[4] LC_NUMERIC=C
[5] LC_TIME=English_United Kingdom.1252
attached base packages:
[1] stats
              graphics grDevices utils
                                            datasets methods
[7] base
other attached packages:
[1] nlme_3.1-128 knitr_1.13
loaded via a namespace (and not attached):
 [1] magrittr_1.5
                     formatR_1.4
                                       tools_3.2.2
 [4] htmltools_0.3.5 yaml_2.1.13
                                       Rcpp_0.12.6
 [7] codetools_0.2-14 stringi_1.1.1
                                       rmarkdown_1.0
[10] highr 0.6
                     grid_3.2.2
                                       stringr 1.0.0
[13] digest_0.6.10
                      lattice_0.20-33 evaluate_0.9
```

This took 1.22 seconds to execute.