Glucose Analysis

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Meta-regression Glucose

##

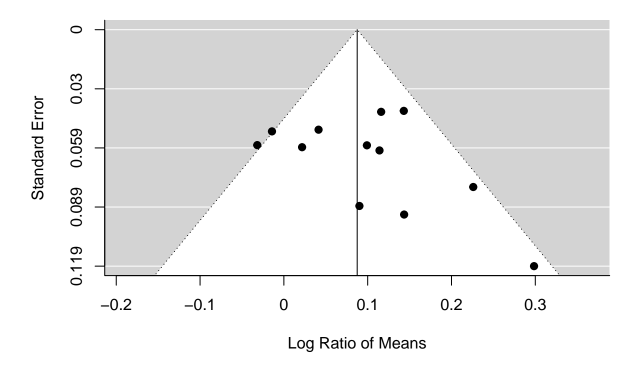
```
load(file="Effect_Size_All_Traits.RData")
load(file="Effect_Size_All_Traits_lnCVR.RData")
#lnRR
Glucose_FBG_lnRR <- subset(Effect_Size_All_Traits, Effect_Size_All_Traits$Trait == "Glucose_FBG")
Glucose_TT_lnRR<- subset(Effect_Size_All_Traits, Effect_Size_All_Traits$Trait == "Glucose_FBG")
Glucose_FBG_lnRR_MG <- subset(Glucose_FBG_lnRR, Glucose_FBG_lnRR$Exposure_Type == "Multigenerational")
Glucose_TT_lnRR_MG <- subset(Glucose_TT_lnRR, Glucose_TT_lnRR$Exposure_Type == "Multigenerational")
Glucose_FBG_lnRR_OF <- subset(Glucose_FBG_lnRR, Glucose_FBG_lnRR$Exposure_Type == "One off")
Glucose_TT_lnRR_OF <- subset(Glucose_TT_lnRR, Glucose_TT_lnRR$Exposure_Type == "One off")
#lnCVR
Glucose_FBG_lnCVR <- subset(Effect_Size_All_Traits_lnCVR, Effect_Size_All_Traits_lnCVR$Trait == "Glucose_TT_lnCVR<- subset(Effect_Size_All_Traits_lnCVR, Effect_Size_All_Traits_lnCVR$Trait == "Glucose_TT_lnCVR, Glucose_FBG_lnCVR$Exposure_Type == "Multigenerational Glucose_TT_lnCVR_MG <- subset(Glucose_TT_lnCVR, Glucose_TT_lnCVR$Exposure_Type == "Multigenerational")
Glucose_FBG_lnCVR_MG <- subset(Glucose_FBG_lnCVR, Glucose_FBG_lnCVR$Exposure_Type == "Multigenerational")
Glucose_FBG_lnCVR_OF <- subset(Glucose_FBG_lnCVR, Glucose_FBG_lnCVR$Exposure_Type == "One off")
Glucose_TT_lnCVR_OF <- subset(Glucose_TT_lnCVR, Glucose_TT_lnCVR$Exposure_Type == "One off")</pre>
```

FBG Analysis (Overall and then split by exposure type)

```
Glucose_overall_lnRR_FBG <- rma.mv(yi, vi, random = list(~1 ES_ID,~1 Cohort_ID), data=Glucose_FBG_lnRR,
summary(Glucose_overall_lnRR_FBG)
##
## Multivariate Meta-Analysis Model (k = 12; method: REML)
##
                           AIC
                                              AICc
##
    logLik Deviance
                                     BIC
   12.1477 -24.2955 -18.2955 -17.1018 -14.8669
##
##
## Variance Components:
##
##
                                              factor
                       sqrt nlvls fixed
              estim
## sigma^2.1 0.0000 0.0000
                                12
                                               ES ID
## sigma^2.2 0.0029 0.0537
                                 9
                                       no Cohort ID
```

```
## Test for Heterogeneity:
## Q(df = 11) = 19.4840, p-val = 0.0529
##
## Model Results:
##
## estimate se zval pval ci.lb ci.ub
## 0.0875 0.0262 3.3447 0.0008 0.0362 0.1388 ***
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

funnel(Glucose_overall_lnRR_FBG)

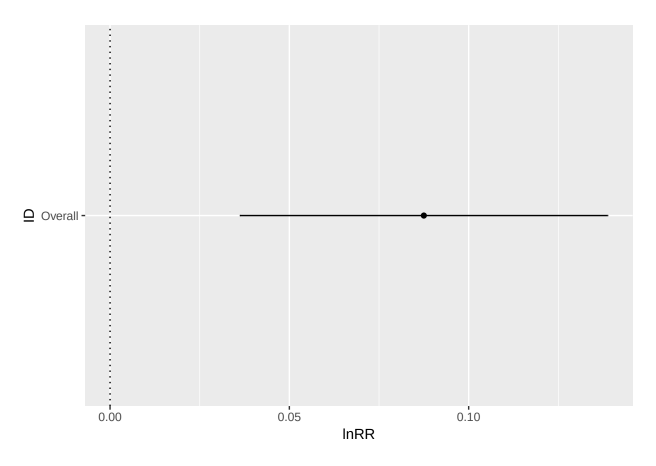


```
#Tibble of overall results

Glucose_overall_lnRR_FBG <- tibble(
   ID = "Overall",
   lnRR = 0.0875,
   ci.lb = 0.0362,
   ci.ub = 0.1388
)</pre>
```

```
## Warning: `list_len()` is deprecated as of rlang 0.2.0.
## Please use `new_list()` instead.
## This warning is displayed once per session.
```

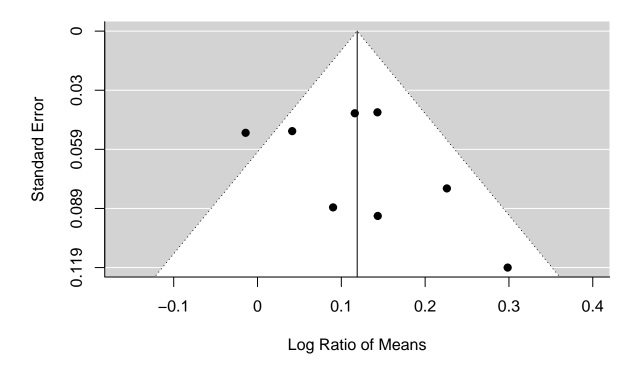
```
plot_Glucose_overall_FBG <- ggplot(Glucose_overall_lnRR_FBG, aes(x=ID, y=lnRR)) +
    geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
    geom_point(aes(x = ID, y = lnRR), position = position_dodge(0.3))+
    geom_hline(yintercept = 0, lty = "dotted") +
    labs(x = "ID", y = "lnRR") +
    coord_flip()
plot_Glucose_overall_FBG</pre>
```



```
#Split by exposure type
Glucose_overall_lnRR_FBG_MG <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_FBG_ln
summary(Glucose_overall_lnRR_FBG_MG)</pre>
```

```
##
## Multivariate Meta-Analysis Model (k = 8; method: REML)
##
##
     logLik Deviance
                            AIC
                                      BIC
                                               AICc
     7.7052 -15.4104
                       -9.4104
                                 -9.5727
                                            -1.4104
##
## Variance Components:
##
##
                        sqrt nlvls fixed
                                               factor
              estim
## sigma^2.1 0.0000 0.0000
                                 8
                                               ES_ID
                                       no
## sigma^2.2 0.0041 0.0641
                                 5
                                       no Cohort_ID
```

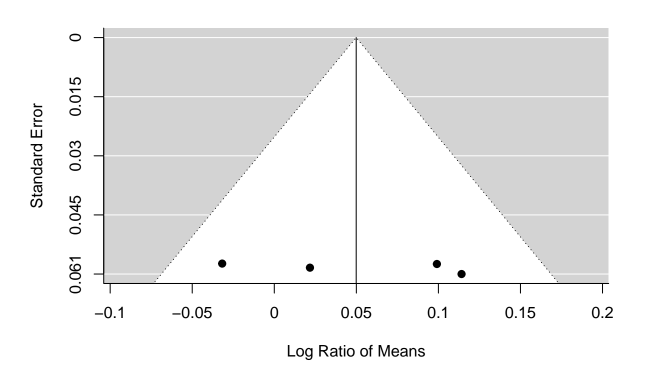
```
##
## Test for Heterogeneity:
## Q(df = 7) = 13.2152, p-val = 0.0670
## Model Results:
##
## estimate
                          pval
                                ci.lb
              se
                   zval
                                      ci.ub
    ##
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
funnel(Glucose_overall_lnRR_FBG_MG)
```



```
Glucose_overall_lnRR_FBG_OF <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_FBG_lnsummary(Glucose_overall_lnRR_FBG_OF)
```

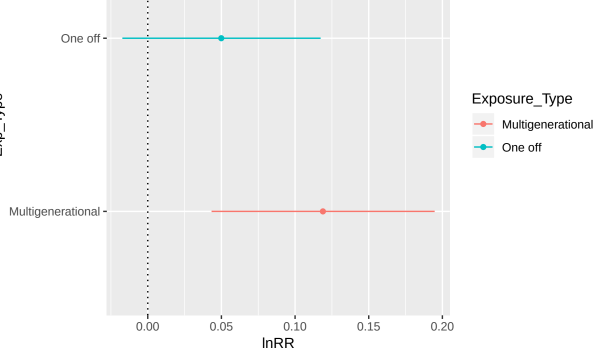
```
##
## Multivariate Meta-Analysis Model (k = 4; method: REML)
##
## logLik Deviance AIC BIC AICc
## 3.7895 -7.5791 -1.5791 -4.2832 22.4209
##
## Wariance Components:
```

```
##
##
                       sqrt nlvls fixed
                                             factor
              estim
## sigma^2.1 0.0006 0.0248
                                 4
                                              ES ID
## sigma^2.2 0.0006 0.0248
                                 4
                                          Cohort_ID
                                      no
## Test for Heterogeneity:
## Q(df = 3) = 4.0610, p-val = 0.2550
##
## Model Results:
##
                      zval
## estimate
                se
                              pval
                                      ci.lb
                                             ci.ub
    0.0499 0.0343 1.4568 0.1452 -0.0172 0.1171
##
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
funnel(Glucose_overall_lnRR_FBG_OF)
```



```
Glucose_FBG_Exp_lnRR <- tibble(
    Exposure_Type = c("Multigenerational", "One off"),
    lnRR = c(0.1189,0.0499),
    ci.lb = c(0.0434,-0.0172),
    ci.ub = c(0.1945,0.1171)
)</pre>
Glucose_FBG_Exp_lnRR
```

```
## # A tibble: 2 x 4
     Exposure_Type
                         lnRR
                                ci.lb ci.ub
     <chr>
                        <dbl>
                                <dbl> <dbl>
##
## 1 Multigenerational 0.119 0.0434 0.194
## 2 One off
                       0.0499 -0.0172 0.117
#Plotting when split by exp type
plot_Glucose_FBG_exp_type <- ggplot(Glucose_FBG_Exp_lnRR, aes(x=Exposure_Type, y=lnRR, colour = Exposur</pre>
  geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
  geom_point(aes(x = Exposure_Type, y = lnRR), position = position_dodge(0.3))+
  geom_hline(yintercept = 0, lty = "dotted") +
  labs(x = "Exp_Type", y = "lnRR") +
  coord_flip()
plot_Glucose_FBG_exp_type
           One off -
```

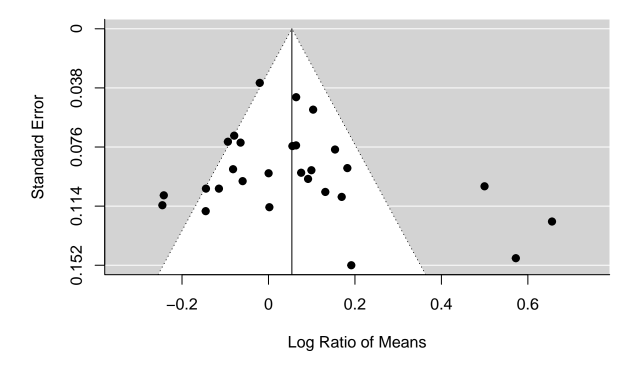


Tolerance test analysis (Overall and then split by exposure type)

```
Glucose_overall_lnRR_TT <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnRR, m
summary(Glucose_overall_lnRR_TT)
```

```
##
## Multivariate Meta-Analysis Model (k = 28; method: REML)
##
##
     logLik Deviance
                            AIC
                                      {\tt BIC}
                                               AICc
     4.7126
              -9.4253
                        -3.4253
                                   0.4623
                                            -2.3818
##
##
## Variance Components:
##
##
               {\tt estim}
                        sqrt nlvls fixed
                                               factor
## sigma^2.1 0.0000 0.0000
                                 28
                                                ES_ID
## sigma^2.2 0.0300 0.1731
                                 27
                                        no
                                            Cohort_ID
## Test for Heterogeneity:
## Q(df = 27) = 105.0308, p-val < .0001
## Model Results:
##
## estimate
                 se
                       zval
                               pval
                                       ci.lb
                                               ci.ub
    0.0541 0.0378 1.4301 0.1527
                                    -0.0200 0.1283
##
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

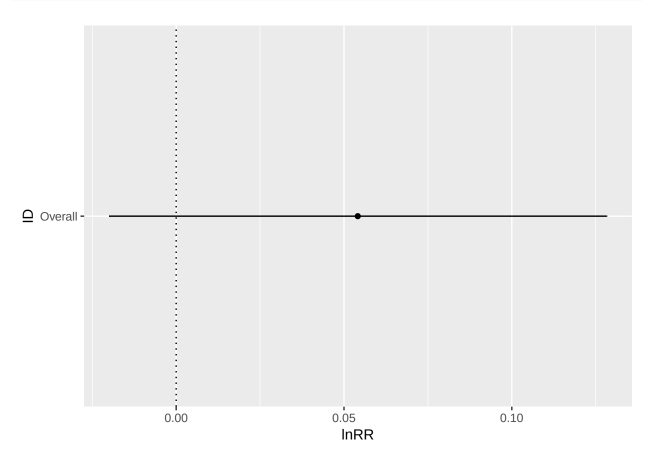
funnel(Glucose_overall_lnRR_TT)



```
#Tibble of overall results

Glucose_overall_lnRR_TT <- tibble(
    ID = "Overall",
    lnRR = 0.0541,
    ci.lb = -0.0200,
    ci.ub = 0.1283
)

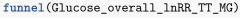
plot_Glucose_overall_TT <- ggplot(Glucose_overall_lnRR_TT, aes(x=ID, y=lnRR)) +
    geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
    geom_point(aes(x = ID, y = lnRR), position = position_dodge(0.3)) +
    geom_hline(yintercept = 0, lty = "dotted") +
    labs(x = "ID", y = "lnRR") +
    coord_flip()
plot_Glucose_overall_TT</pre>
```

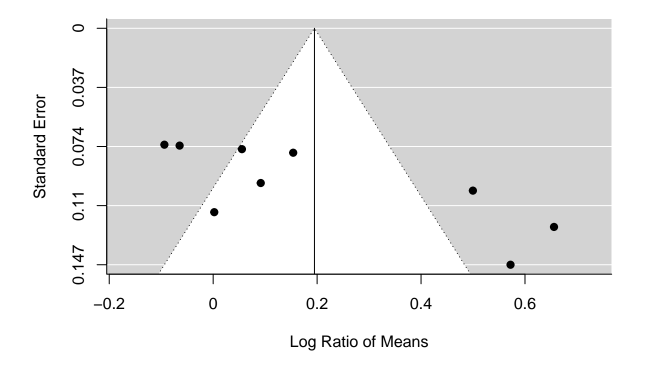


```
#Split by exposure type
Glucose_overall_lnRR_TT_MG <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnRR
summary(Glucose_overall_lnRR_TT_MG)</pre>
```

Multivariate Meta-Analysis Model (k = 9; method: REML)

```
##
##
    logLik Deviance
                         AIC
                                  BIC
                                          AICc
   -1.2829
             2.5658
                      8.5658
                               8.8041
##
                                       14.5658
##
## Variance Components:
##
##
                     sqrt nlvls
                                          factor
             estim
                                fixed
                                           ES_ID
## sigma^2.1 0.0341
                   0.1846
                              9
## sigma^2.2 0.0341 0.1846
                              9
                                   no
                                       Cohort_ID
##
## Test for Heterogeneity:
## Q(df = 8) = 59.3958, p-val < .0001
## Model Results:
##
## estimate
               se
                    zval
                           pval
                                  ci.lb
                                         ci.ub
##
    ##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

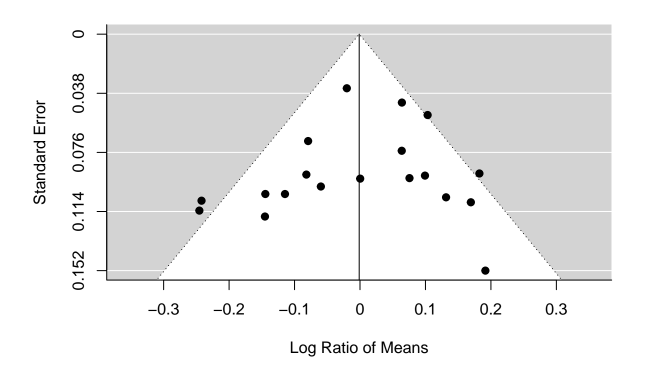




Glucose_overall_lnRR_TT_OF <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnRR
summary(Glucose_overall_lnRR_TT_OF)</pre>

```
##
## Multivariate Meta-Analysis Model (k = 19; method: REML)
##
##
    logLik Deviance
                           AIC
                                    BIC
                                             AICc
  11.8580 -23.7160 -17.7160 -15.0448 -16.0017
##
## Variance Components:
##
##
                       sqrt nlvls fixed
                                             factor
              estim
## sigma^2.1 0.0000 0.0000
                                19
                                              ES ID
## sigma^2.2 0.0064 0.0801
                                18
                                          Cohort_ID
                                      no
## Test for Heterogeneity:
## Q(df = 18) = 35.5344, p-val = 0.0081
##
## Model Results:
##
                               pval
                                      ci.lb
## estimate
                se
                       zval
                                              ci.ub
##
  -0.0012 0.0276 -0.0446 0.9645 -0.0552 0.0528
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

funnel(Glucose_overall_lnRR_TT_OF)



<dbl> <dbl>

0.0124 0.377

-0.00120 -0.0552 0.0528

<dbl>

##

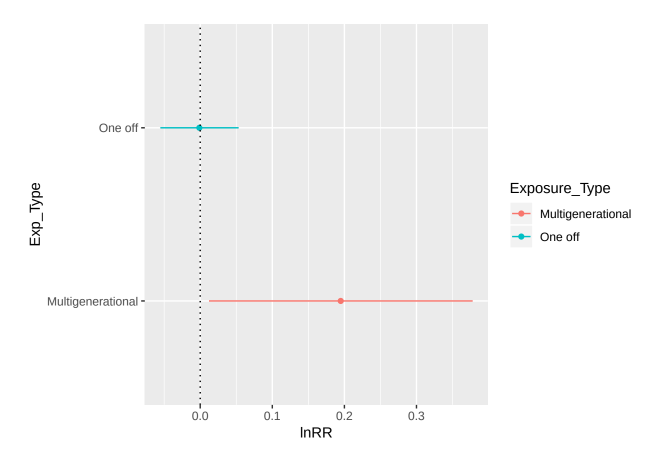
<chr>

2 One off

1 Multigenerational 0.195

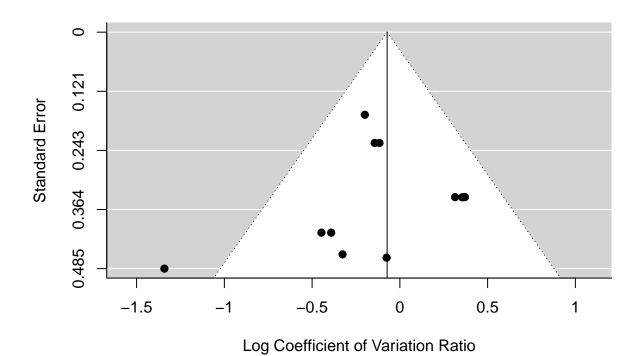
```
#Plotting when split by exp type

plot_Glucose_TT_exp_type <- ggplot(Glucose_TT_Exp_lnRR, aes(x=Exposure_Type, y=lnRR, colour = Exposure_geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
    geom_point(aes(x = Exposure_Type, y = lnRR), position = position_dodge(0.3))+
    geom_hline(yintercept = 0, lty = "dotted") +
    labs(x = "Exp_Type", y = "lnRR") +
    coord_flip()
plot_Glucose_TT_exp_type</pre>
```



FBG Analysis (lnCVR)

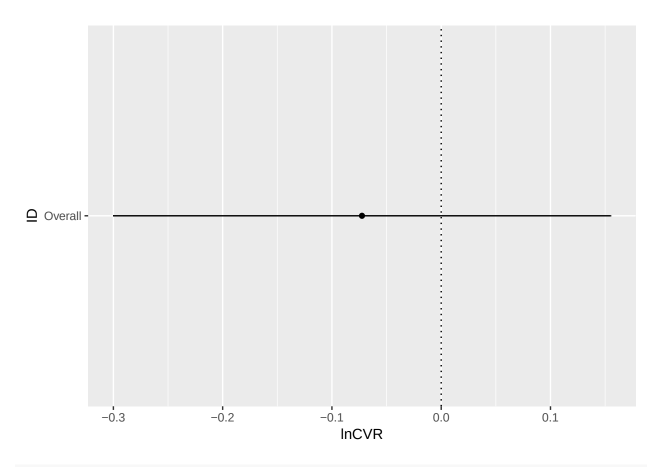
```
Glucose_overall_lnCVR_FBG <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_FBG_lnCV
summary(Glucose_overall_lnCVR_FBG)
##
## Multivariate Meta-Analysis Model (k = 12; method: REML)
##
   logLik Deviance
                          AIC
                                    BIC
                                             AICc
## -5.8607
            11.7215 17.7215
                                18.9152
                                          21.1501
##
## Variance Components:
##
                       sqrt nlvls fixed
                                             factor
              estim
                                              ES_ID
## sigma^2.1 0.0000 0.0000
                               12
                                      no
## sigma^2.2 0.0401 0.2003
                                9
                                      no Cohort_ID
##
## Test for Heterogeneity:
## Q(df = 11) = 15.5523, p-val = 0.1586
##
## Model Results:
##
## estimate
                                      ci.lb
                se
                       zval
                              pval
## -0.0725 0.1162 -0.6243 0.5324 -0.3003 0.1552
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
funnel(Glucose_overall_lnCVR_FBG)
```



#Tibble of overall results

Glucose_overall_lnCVR_FBG <- tibble(
 ID = "Overall",
 lnCVR = -0.0725,
 ci.lb = -0.3003,
 ci.ub = 0.1552
)

plot_Glucose_overall_FBG_lnCVR <- ggplot(Glucose_overall_lnCVR_FBG, aes(x=ID, y=lnCVR)) +
 geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
 geom_point(aes(x = ID, y = lnCVR), position = position_dodge(0.3))+
 geom_hline(yintercept = 0, lty = "dotted") +
 labs(x = "ID", y = "lnCVR") +
 coord_flip()
plot_Glucose_overall_FBG_lnCVR</pre>



```
#Split by exposure type
Glucose_overall_lnCVR_FBG_MG <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_FBG_landom = list(~1|ES_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohort_ID,~1|Cohor
summary(Glucose_overall_lnCVR_FBG_MG)
##
## Multivariate Meta-Analysis Model (k = 8; method: REML)
##
##
                      logLik Deviance
                                                                                                                                       AIC
                                                                                                                                                                                      BIC
                                                                                                                                                                                                                                   AICc
##
              -2.0156
                                                                       4.0313
                                                                                                                   10.0313
                                                                                                                                                                         9.8690
                                                                                                                                                                                                                    18.0313
##
## Variance Components:
##
                                                                                                                                                                                                                                   factor
                                                                       estim
                                                                                                                   sqrt nlvls fixed
## sigma^2.1 0.0000 0.0000
                                                                                                                                                                   8
                                                                                                                                                                                                                                       ES_ID
                                                                                                                                                                                               no
## sigma^2.2 0.0000 0.0000
                                                                                                                                                                   5
                                                                                                                                                                                               no Cohort_ID
##
## Test for Heterogeneity:
## Q(df = 7) = 6.2373, p-val = 0.5123
```

ci.ub

ci.lb

Model Results:

zval

-0.2477 0.1005 -2.4652 0.0137 -0.4446 -0.0508 *

se

pval

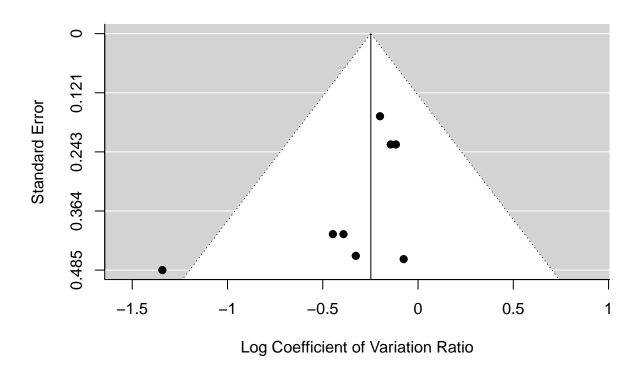
estimate

##

##

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

funnel(Glucose_overall_lnCVR_FBG_MG)
```

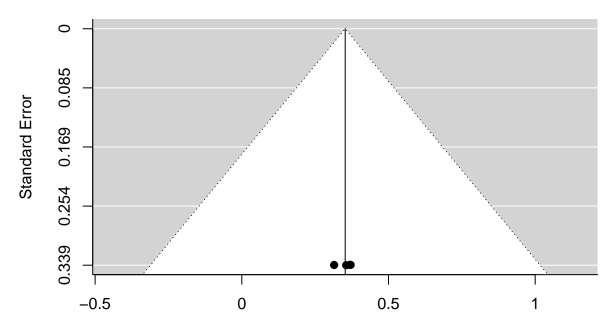


Glucose_overall_lnCVR_FBG_OF <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_FBG_lnsummary(Glucose_overall_lnCVR_FBG_OF)

```
##
## Multivariate Meta-Analysis Model (k = 4; method: REML)
##
##
     logLik Deviance
                            AIC
                                      BIC
                                                AICc
     0.4841
              -0.9683
                         5.0317
                                   2.3276
                                             29.0317
##
##
## Variance Components:
##
               estim
                        sqrt
                              nlvls fixed
                                                factor
## sigma^2.1 0.0000 0.0000
                                  4
                                                 ES_ID
## sigma^2.2 0.0000 0.0000
                                            Cohort_ID
                                        no
## Test for Heterogeneity:
## Q(df = 3) = 0.0177, p-val = 0.9994
## Model Results:
```

```
##
## estimate se zval pval ci.lb ci.ub
## 0.3524 0.1692 2.0825 0.0373 0.0207 0.6842 *
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

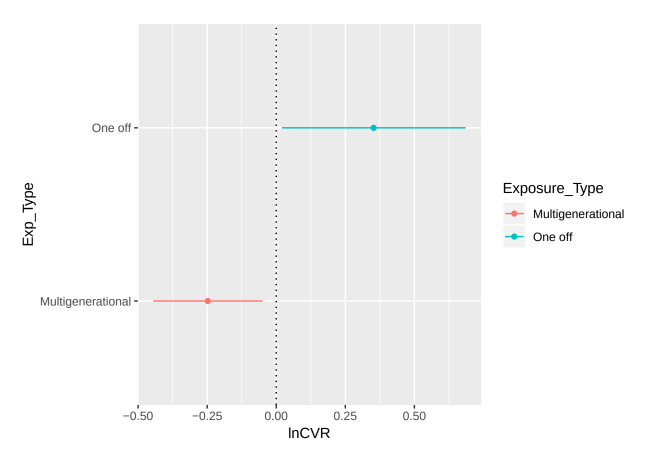
funnel(Glucose_overall_lnCVR_FBG_0F)
```



Log Coefficient of Variation Ratio

```
Glucose_FBG_Exp_lnCVR <- tibble(</pre>
  Exposure_Type = c("Multigenerational", "One off"),
  lnCVR = c(-0.2477, 0.3524),
  ci.lb = c(-0.4446, 0.0207),
  ci.ub = c(-0.0508, 0.6842)
Glucose_FBG_Exp_lnCVR
## # A tibble: 2 x 4
##
    Exposure_Type
                        lnCVR
                                 ci.lb
                                         ci.ub
     <chr>
                                 <dbl>
                                         <dbl>
                        <dbl>
## 1 Multigenerational -0.248 -0.445 -0.0508
## 2 One off
                        0.352 0.0207 0.684
```

```
#Plotting when split by exp type
plot_Glucose_FBG_exp_type_lnCVR <- ggplot(Glucose_FBG_Exp_lnCVR, aes(x=Exposure_Type, y=lnCVR, colour =
    geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
    geom_point(aes(x = Exposure_Type, y = lnCVR), position = position_dodge(0.3))+
    geom_hline(yintercept = 0, lty = "dotted") +
    labs(x = "Exp_Type", y = "lnCVR") +
    coord_flip()
plot_Glucose_FBG_exp_type_lnCVR</pre>
```



Tolerance test analysis (lnCVR)

##

##

-10.5642

logLik Deviance

21.1283

AIC

27.1283

```
Glucose_overall_lnCVR_TT <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnCVR,
summary(Glucose_overall_lnCVR_TT)

##
## Multivariate Meta-Analysis Model (k = 28; method: REML)
##</pre>
```

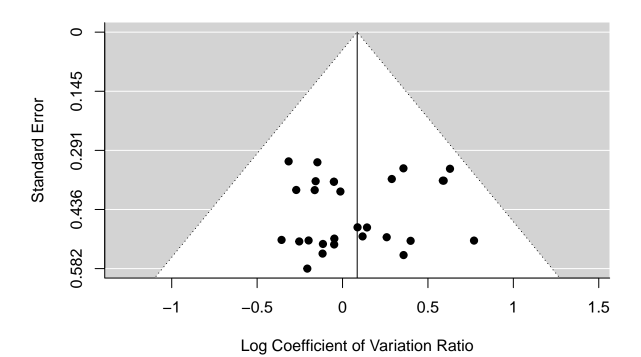
AICc

28.1718

BIC

31.0158

```
## Variance Components:
##
##
              estim
                       sqrt nlvls fixed
                                              factor
## sigma^2.1 0.0000 0.0000
                                28
                                               ES_ID
                                       no
## sigma^2.2 0.0000 0.0000
                                27
                                       no Cohort_ID
##
## Test for Heterogeneity:
## Q(df = 27) = 16.2263, p-val = 0.9485
##
## Model Results:
## estimate
                              pval
                                      ci.lb
                                              ci.ub
                      zval
    0.0859 0.0795 1.0805 0.2799 -0.0699 0.2416
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
funnel(Glucose_overall_lnCVR_TT)
```

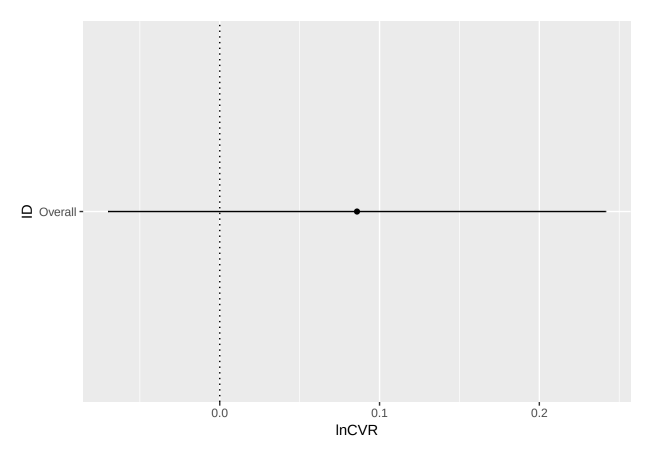


```
#Tibble of overall results

Glucose_overall_lnCVR_TT <- tibble(
   ID = "Overall",
   lnCVR = 0.0859,
   ci.lb =-0.0699 ,</pre>
```

```
ci.ub =0.2416
)

plot_Glucose_overall_TT_lnCVR <- ggplot(Glucose_overall_lnCVR_TT, aes(x=ID, y=lnCVR)) +
    geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
    geom_point(aes(x = ID, y = lnCVR), position = position_dodge(0.3))+
    geom_hline(yintercept = 0, lty = "dotted") +
    labs(x = "ID", y = "lnCVR") +
    coord_flip()
plot_Glucose_overall_TT_lnCVR</pre>
```



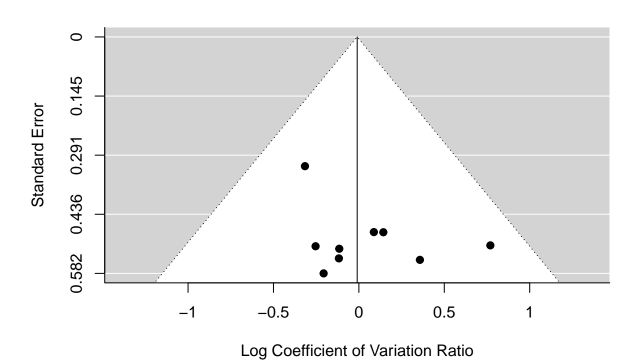
```
#Split by exposure type
Glucose_overall_lnCVR_TT_MG <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnC
summary(Glucose_overall_lnCVR_TT_MG)
## Multivariate Meta-Analysis Model (k = 9; method: REML)
##
##
                            AIC
                                               AICc
    logLik Deviance
                                      BIC
##
   -3.8680
               7.7360
                        13.7360
                                  13.9744
                                            19.7360
```

##

##

Variance Components:

```
estim
                       sqrt nlvls fixed
                                              factor
## sigma^2.1 0.0000 0.0000
                                 9
                                               ES_ID
                                       no
## sigma^2.2 0.0000 0.0000
                                       no
                                           Cohort_ID
##
## Test for Heterogeneity:
## Q(df = 8) = 4.2413, p-val = 0.8347
## Model Results:
##
## estimate
                        zval
                               pval
                                       ci.lb
                                               ci.ub
   -0.0098 0.1592 -0.0615 0.9509 -0.3219 0.3023
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
funnel(Glucose_overall_lnCVR_TT_MG)
```

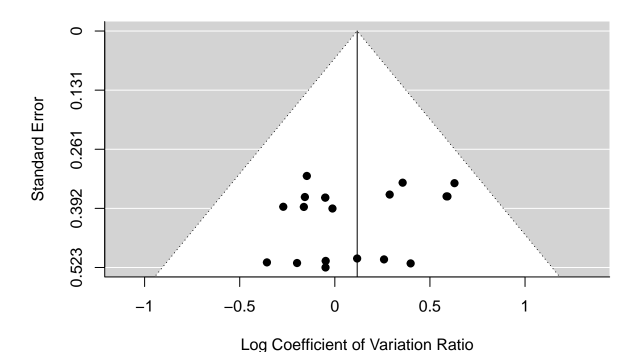


Glucose_overall_lnCVR_TT_OF <- rma.mv(yi, vi, random = list(~1|ES_ID,~1|Cohort_ID), data=Glucose_TT_lnC
summary(Glucose_overall_lnCVR_TT_OF)
##</pre>

```
## Multivariate Meta-Analysis Model (k = 19; method: REML)
##
## logLik Deviance AIC BIC AICc
```

```
-6.3263
            12.6526
                     18.6526
                             21.3237
                                      20.3669
##
## Variance Components:
##
##
             estim
                     sqrt nlvls fixed
                                         factor
## sigma^2.1 0.0000 0.0000
                             19
                                          ES_ID
                                   no
## sigma^2.2 0.0000 0.0000
                            18
                                   no
                                      Cohort_ID
## Test for Heterogeneity:
## Q(df = 18) = 11.5042, p-val = 0.8717
## Model Results:
## estimate
                    zval
                                  ci.lb
                                         ci.ub
              se
                           pval
##
    ##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

funnel(Glucose_overall_lnCVR_TT_OF)



Glucose_TT_Exp_lnCVR <- tibble(
 Exposure_Type = c("Multigenerational", "One off"),
 lnCVR = c(-0.0098,0.1176),
 ci.lb = c(-0.3219,-0.0621),</pre>

```
ci.ub = c(0.3023, 0.2973)
Glucose_TT_Exp_lnCVR
## # A tibble: 2 x 4
##
     Exposure_Type
                         lnCVR
                                 ci.lb ci.ub
     <chr>
                                 <dbl> <dbl>
##
                         <dbl>
## 1 Multigenerational -0.0098 -0.322 0.302
## 2 One off
                        0.118 -0.0621 0.297
#Plotting when split by exp type
plot_Glucose_TT_exp_type_lnCVR <- ggplot(Glucose_TT_Exp_lnCVR, aes(x=Exposure_Type, y=lnCVR, colour = E</pre>
  geom_errorbar(aes(ymin=ci.lb, ymax=ci.ub), width = 0, position = position_dodge(0.3)) +
  geom_point(aes(x = Exposure_Type, y = lnCVR), position = position_dodge(0.3))+
  geom_hline(yintercept = 0, lty = "dotted") +
  labs(x = "Exp_Type", y = "lnCVR") +
  coord_flip()
plot_Glucose_TT_exp_type_lnCVR
```

