Meta-Analysis Reproduction Summary Excluding HUM251

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1. Summary

This document reports the process taken in the model fitting stage of the meta-analysis in thermal reproduction.

2. Setup

We first read in our data and select all of the effect sizes related to reproduction. We do this using the following code.

Next we create new columns in our dataframe which will serve as random factors in our multi-level meta analysis models. The following initialises four new columns, namely "obs", "study_code", "Species.phylo" and "species". Lastly, we create a column name "precision" which is equal to the inverse standard error.

```
### Create random factors into data frame
rdata$obs <- factor(c(1:nrow(rdata))) # Unique observation code
rdata$study_code <- factor(rdata$Paper.code) # Model requires column names study_code
rdata$Species.phylo <- factor(rdata$Species.latin) # Species names for phylo matrix
rdata$species <- factor(rdata$Species.latin) # Another species column for random factor
precision <- sqrt(1/rdata$v) # inverse standard error
rdata[, "precision"] <- precision</pre>
```

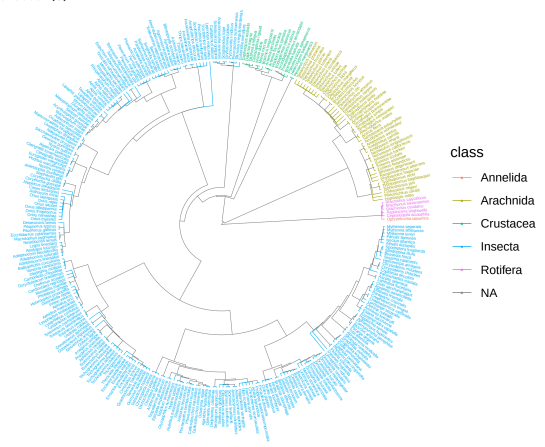
The number of species and total number of studies present in the data are as follows.

```
nlevels(rdata$species) # Check number of species
## [1] 307
nlevels(rdata$study_code) # Check number of studies
## [1] 340
```

The final stage in the setup is to import a phylogentic tree of the data. Below is the code used to produce the tree and a plot of the tree itself.

```
## import tree from map
tree1 <- read.nexus("all_reproduction_excHUM251_tree.nex")
tree_grafen = compute.brlen(tree1, method = "Grafen", power = 1)
phylo_matrix <- vcv(tree_grafen, cor = TRUE, model = "Brownian") # Make phylogenetic matrix</pre>
```

character(0)



3. Random effects models

In this section we determine which random effects to include in our model. For each model I have provided the code used to specify the structure of the model and a summary of the results. We begin with a model that includes all of the random factors we created earlier.

```
# Adding four random factors
meta2 <- rma.mv(es, v, random = list(~1 | Species.phylo, ~1 |</pre>
    species, ~1 | study_code, ~1 | obs), R = list(Species.phylo = phylo_matrix),
    data = rdata, method = "REML")
summary(meta2)
##
  Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
##
                                   AIC
                                                BIC
                                                           AICc
       logLik
                 Deviance
   -4119.6114
                8239.2228
                             8249.2228
                                          8275.5219
                                                      8249.2651
##
##
## Variance Components:
##
##
                                                      factor
                                                                 R
                estim
                                nlvls
                                       fixed
                          sqrt
               0.0000
## sigma^2.1
                        0.0000
                                  307
                                           no
                                               Species.phylo
                                                               yes
## sigma^2.2
               0.0000
                        0.0004
                                  307
                                                     species
                                           nο
                                                                nο
## sigma^2.3
              10.2889
                        3.2076
                                  340
                                           no
                                                  study_code
                                                                no
## sigma^2.4
               5.0483
                        2.2468
                                 1423
                                           nο
                                                          obs
                                                                nο
##
## Test for Heterogeneity:
  Q(df = 1422) = 37705.7968, p-val < .0001
##
## Model Results:
##
## estimate
                                          ci.lb
                                                   ci.ub
                 se
                         zval
                                 pval
##
    -1.5435
             0.1886
                      -8.1844
                              <.0001
                                       -1.9131
                                                -1.1739
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
i2_ml(meta2, method = c("ratio")) # Heterogeneity at each random factor level
##
           I2_Total I2_Species.phylo
                                             I2_species
                                                            I2_study_code
                                                                                     I2_obs
##
       9.945130e+01
                         3.084327e-10
                                           1.214829e-06
                                                             6.671656e+01
                                                                               3.273474e+01
```

Accounting for non-independence of data points from the same experiment

The data has a nested structure. Each study (study_code) may have a number of experiments (effect.size.code) which share a common control temperature. Each effect size has its own unique code, obs. Effect sizes from the same experiment which share a control temperature are thought to be non-independent. The following code create a covariance matrix "VCV_shared" which assumes a correlation of 0.5 between effect sizes from the same experiment. We include this structure in our proceeding models.

```
meta3 <- rma.mv(es, VCV_shared, random = list(~1 | Species.phylo,</pre>
    ~1 | species, ~1 | study_code, ~1 | obs), R = list(Species.phylo = phylo_matrix),
   data = rdata, method = "REML")
summary(meta3)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
                Deviance
##
      logLik
                                 AIC
                                             BIC
                                                        AICc
               7677.0632
## -3838.5316
                           7687.0632
                                       7713.3623
                                                   7687.1055
##
## Variance Components:
##
##
              estim
                       sqrt nlvls fixed
                                                  factor
## sigma^2.1 0.0000 0.0000
                               307
                                       no
                                           Species.phylo yes
## sigma^2.2 0.0000 0.0002
                               307
                                       no
                                                 species
                                                           no
                                              study_code
## sigma^2.3 5.0925 2.2567
                               340
                                       no
                                                           no
## sigma^2.4 4.3215 2.0788
                              1423
                                                     obs
                                       no
                                                           no
## Test for Heterogeneity:
## Q(df = 1422) = 39436.4596, p-val < .0001
## Model Results:
##
## estimate
                se
                       zval
                               pval
                                       ci.lb
## -1.3309 0.1399 -9.5102 <.0001 -1.6052 -1.0566 ***
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
i2_ml(meta3, method = c("ratio")) # Heterogeneity at each random factor level
           I2_Total I2_Species.phylo
                                                        I2_study_code
##
                                          I2 species
                                                                                I2 obs
                                                         5.361290e+01
##
       9.910914e+01
                       3.211305e-09
                                        6.034366e-07
                                                                          4.549624e+01
```

Model without phylogeny

We also originally included a variance-covariance matrix for phylogenetic relatedness of the included species as a random effect in the model (Chamberlain et al., 2012), but this has now been removed as its inclusion did not improve model fit and the phylogenetic signal was very weak.

```
## without phylogeny but with shared control
meta5 <- rma.mv(es, VCV_shared, random = list(~1 | species, ~1 |</pre>
    study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta5)
##
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
       logLik
                 Deviance
                                  AIC
                                              BIC
                                                         AICc
                                        7706.1024
                                                    7685.0914
##
  -3838.5316
                7677.0632
                            7685.0632
##
## Variance Components:
##
##
                                                factor
               estim
                        sqrt nlvls
                                    fixed
## sigma^2.1 0.0000 0.0003
                                307
                                        no
                                               species
## sigma^2.2 5.0925
                      2.2567
                                            study_code
                                340
                                        no
## sigma^2.3 4.3215
                     2.0788
                               1423
                                                   obs
                                        no
##
## Test for Heterogeneity:
## Q(df = 1422) = 39436.4596, p-val < .0001
##
## Model Results:
##
## estimate
                                pval
                                        ci.lb
                 se
                        zval
                                                 ci.ub
   -1.3309 0.1399
                    -9.5102
                             <.0001
                                      -1.6052
                                               -1.0566
##
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
i2_ml(meta5, method = c("ratio")) # Heterogeneity at each random factor level
                    I2_species I2_study_code
##
        I2_Total
                                                    I2_obs
   9.910913e+01 6.890664e-07 5.361289e+01 4.549624e+01
##
```

Model without phylogeny or species

```
## without phylogeny or species
meta4 <- rma.mv(es, VCV_shared, random = list(~1 | study_code,</pre>
   ~1 | obs), data = rdata, method = "REML")
summary(meta4)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
      logLik
                Deviance
                                 AIC
                                             BIC
                                                        AICc
## -3838.5316
               7677.0632
                          7683.0632
                                       7698.8426
                                                   7683.0801
##
## Variance Components:
##
##
                                               factor
              estim
                       sqrt nlvls fixed
## sigma^2.1 5.0925 2.2567
                               340
                                       no study_code
## sigma^2.2 4.3215 2.0788
                              1423
                                                  obs
##
## Test for Heterogeneity:
## Q(df = 1422) = 39436.4596, p-val < .0001
##
## Model Results:
##
## estimate
                                       ci.lb
              se
                       zval
                               pval
                                                ci.ub
## -1.3309 0.1399 -9.5102 <.0001 -1.6052 -1.0566 ***
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
i2_ml(meta4, method = c("ratio")) # Heterogeneity at each random factor level
##
       I2_Total I2_study_code
                                     I2_obs
       99.10914
##
                     53.61290
                                   45.49624
```

Model without phylogeny, species or study_code

```
## without phylogeny, species or study_code
meta7 <- rma.mv(es, VCV_shared, random = list(~1 | obs), data = rdata,</pre>
   method = "REML")
summary(meta7)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
       logLik
                 Deviance
                                  AIC
                                               BIC
                                                          AICc
  -3998.6362
                7997.2725
                            8001.2725
                                        8011.7921
                                                     8001.2810
##
##
## Variance Components:
##
##
               {\tt estim}
                        sqrt nlvls
                                     fixed factor
## sigma^2
              6.8055
                      2.6087
                               1423
                                        no
                                                obs
##
## Test for Heterogeneity:
## Q(df = 1422) = 39436.4596, p-val < .0001
##
## Model Results:
##
                                          ci.lb
## estimate
                 se
                         zval
                                 pval
                                                   ci.ub
   -1.2600 0.0721 -17.4635 <.0001 -1.4014 -1.1185
##
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
i2_ml(meta7, method = c("ratio")) # Heterogeneity at each random factor level
## I2_Total
              I2_obs
## 98.77186 98.77186
```

We can see from the above that the best fitting model according to AIC is "meta4" which includes only the study code and the unique effect size code, obs. There is a AIC difference of 4 between the model meta4 and the next best model meta5. We continue our analysis using meta4 as our base model.

4. Meta-regressions

Starting with the best fitting random-effect model from Section 3, "meta8" we now include single factors as a fixed effect. We initially explore the fixed factors

- reftemp: The experiment's control (reference) temperature.
- treattemp: The treatment temperature
- warm.cool : A categorical variable indicating whether treatment is warmer or cooler than the reference temperature
- diff: The difference between the reference and treatment temperature.

Reference temperature

```
meta_trait_ref <- rma.mv(es, VCV_shared, mod = ~reftemp, random = list(~1 |</pre>
    study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta_trait_ref)
##
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
       logLik
                 Deviance
                                   AIC
                                               BIC
                                                           AICc
##
  -3835.2711
                7670.5423
                             7678.5423
                                         7699.5787
                                                     7678.5705
##
## Variance Components:
##
##
                              nlvls
                                      fixed
                                                 factor
               estim
                         sqrt
## sigma^2.1
              5.0483
                      2.2468
                                 340
                                             study_code
                                         no
## sigma^2.2
              4.3315
                      2.0812
                                1423
                                                     obs
                                         no
##
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39262.5694, p-val < .0001
##
## Test of Moderators (coefficient 2):
## QM(df = 1) = 1.6304, p-val = 0.2016
##
## Model Results:
##
##
            estimate
                           se
                                  zval
                                          pval
                                                  ci.lb
                                                           ci.ub
## intrcpt
              0.1221
                      1.1464
                                0.1065
                                        0.9152
                                                -2.1248
                                                          2.3689
## reftemp
             -0.0588
                      0.0461
                              -1.2769
                                       0.2016
                                                -0.1492
                                                         0.0315
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Treatment temperature

```
meta_trait_treattemp <- rma.mv(es, VCV_shared, mod = ~treattemp,</pre>
   random = list(~1 | study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta_trait_treattemp)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
                               AIC
##
      logLik
               Deviance
                                          BIC
                                                     AICc
## -3828.1823
              7656.3646
                        7664.3646
                                     7685.4011
                                                7664.3929
##
## Variance Components:
##
             estim
                      sqrt nlvls fixed
                                            factor
## sigma^2.1 5.1036 2.2591
                             340
                                     no
                                        study_code
## sigma^2.2 4.2445 2.0602
                             1423
                                               obs
                                     no
##
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39389.3398, p-val < .0001
##
## Test of Moderators (coefficient 2):
## QM(df = 1) = 16.9542, p-val < .0001
##
## Model Results:
##
                                       pval
##
             estimate
                          se
                                zval
                                               ci.lb
                                                       ci.ub
## intrcpt
             -0.4771 0.2496 -1.9113 0.0560 -0.9663
                                                      0.0122
            ## treattemp
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Warm vs Cool

```
meta_trait_warm <- rma.mv(es, VCV_shared, mod = ~warm.cool, random = list(~1 |</pre>
   study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta_trait_warm)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
                Deviance
                                 AIC
                                             BIC
                                                        AICc
      logLik
               7657.9564
## -3828.9782
                           7665.9564
                                       7686.9928
                                                   7665.9846
##
## Variance Components:
##
##
              estim
                       sqrt nlvls fixed
                                               factor
## sigma^2.1 5.0837 2.2547
                               340
                                       no
                                           study_code
## sigma^2.2 4.2511 2.0618
                              1423
                                                  obs
                                       no
##
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39145.1933, p-val < .0001
##
## Test of Moderators (coefficient 2):
## QM(df = 1) = 15.6100, p-val < .0001
##
## Model Results:
##
##
                                                      ci.lb
                 estimate
                               se
                                      zval
                                              pval
                                                               ci.ub
## intrcpt
                  -1.0722 0.1540 -6.9633 <.0001 -1.3739 -0.7704
                 -0.5057 0.1280 -3.9509 <.0001 -0.7565 -0.2548 ***
## warm.coolWarm
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

We model warm versus cool without and intercept so we can visualise the estimates easier.

```
meta_trait_warm_nointer <- rma.mv(es, VCV_shared, mod = ~warm.cool -</pre>
    1, random = list(~1 | study_code, ~1 | obs), data = rdata,
    method = "REML")
summary(meta_trait_warm_nointer)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
       logLik
                 Deviance
                                  AIC
                                              BIC
                                                         AICc
## -3828.9782
                7657.9564
                                        7686.9928
                           7665.9564
                                                    7665.9846
##
## Variance Components:
##
##
                                                factor
               {\tt estim}
                        sqrt nlvls fixed
## sigma^2.1 5.0837
                      2.2547
                                340
                                        no
                                            study_code
## sigma^2.2 4.2511 2.0618
                               1423
                                        no
                                                   obs
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39145.1933, p-val < .0001
##
## Test of Moderators (coefficients 1:2):
## QM(df = 2) = 106.1596, p-val < .0001
##
## Model Results:
##
##
                  estimate
                                se
                                        zval
                                                pval
                                                        ci.lb
                                                                 ci.ub
## warm.coolCool
                  -1.0722 0.1540
                                     -6.9633 <.0001 -1.3739
                                                               -0.7704
                  -1.5778 0.1532 -10.2985 <.0001 -1.8781 -1.2776 ***
## warm.coolWarm
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Difference

```
meta_trait_diff <- rma.mv(es, VCV_shared, mod = ~diff, random = list(~1 |</pre>
   study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta_trait_diff)
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
               Deviance
                                AIC
                                           BIC
                                                     AICc
      logLik
              7657.7529
## -3828.8765
                          7665.7529
                                     7686.7894
                                                 7665.7812
##
## Variance Components:
##
##
              estim
                      sqrt nlvls fixed
                                             factor
## sigma^2.1 5.1566 2.2708
                              340
                                     no
                                         study_code
## sigma^2.2 4.2399 2.0591
                             1423
                                                obs
                                     no
##
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39327.3415, p-val < .0001
##
## Test of Moderators (coefficient 2):
## QM(df = 1) = 15.6189, p-val < .0001
##
## Model Results:
##
           estimate
                        se
                               zval
                                      pval
                                              ci.lb
                                                      ci.ub
## intrcpt
          -1.3368 0.1404 -9.5227 <.0001 -1.6120 -1.0617 ***
## diff
            ##
## ---
```

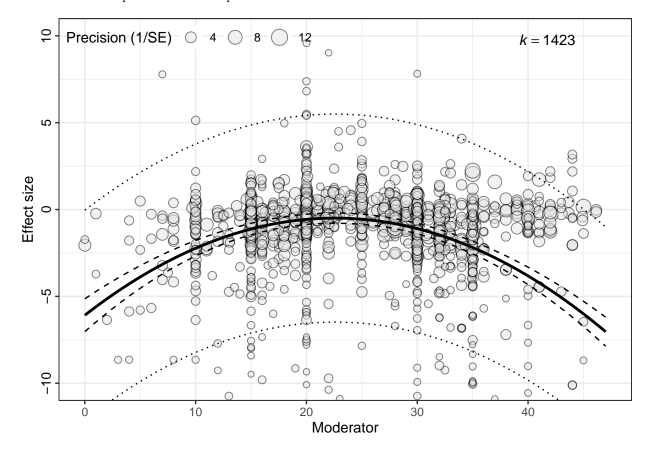
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Treatment temperature as a quadratic effect

We expect that treatment temperature will have a quadratic effect on reproduction. In other words, we expect that reproduction will decrease either side of an optimum temperature for each species. We also assume that this optimum temperature will be close to the control temperature. That said, we investigate **treattemp** as a quadratic fixed effect.

```
meta_trait_treat2 <- rma.mv(es, VCV_shared, mod = ~poly(treattemp,</pre>
    degree = 2, raw = TRUE), random = list(~1 | study_code, ~1 |
   obs), data = rdata, method = "REML")
summary(meta_trait_treat2)
##
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
##
       logLik
                 Deviance
                                   AIC
                                               BIC
                                                          AICc
   -3750.0876
                7500.1753
                            7510.1753
                                         7536.4673
                                                     7510.2177
##
##
## Variance Components:
##
##
                              nlvls
                                     fixed
                                                 factor
               estim
                        sqrt
## sigma^2.1 5.7801
                      2.4042
                                340
                                         no
                                             study_code
## sigma^2.2 3.5306
                     1.8790
                                1423
                                        no
                                                    obs
##
## Test for Residual Heterogeneity:
## QE(df = 1420) = 37352.2717, p-val < .0001
##
## Test of Moderators (coefficients 2:3):
## QM(df = 2) = 186.7480, p-val < .0001
##
## Model Results:
##
##
                                              estimate
                                                            se
                                                                     zval
                                                                             pval
                                                                                     ci.lb
                                                                                              ci.ub
## intrcpt
                                               -6.0828
                                                                -12.2239
                                                                           <.0001
                                                                                  -7.0581
                                                        0.4976
## poly(treattemp, degree = 2, raw = TRUE)1
                                                0.4951
                                                        0.0418
                                                                 11.8515
                                                                           <.0001
                                                                                    0.4133
                                                                                             0.5770
## poly(treattemp, degree = 2, raw = TRUE)2
                                               -0.0110
                                                       0.0008
                                                                -12.9197
                                                                          <.0001
                                                                                  -0.0126
                                                                                            -0.0093
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

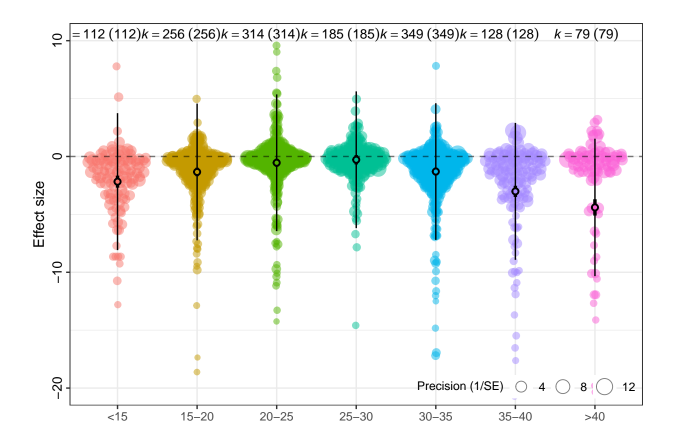
Below is a bubble plot of the fitted quadratic model.



Modelling response with binned treatment temperatures

The last model we will try is one with categorised or 'binned' treatment temperatures. We categories our effect sizes into one seven bins depending on the treatment temperature. The bins and the number of effect sizes in each bin are given below

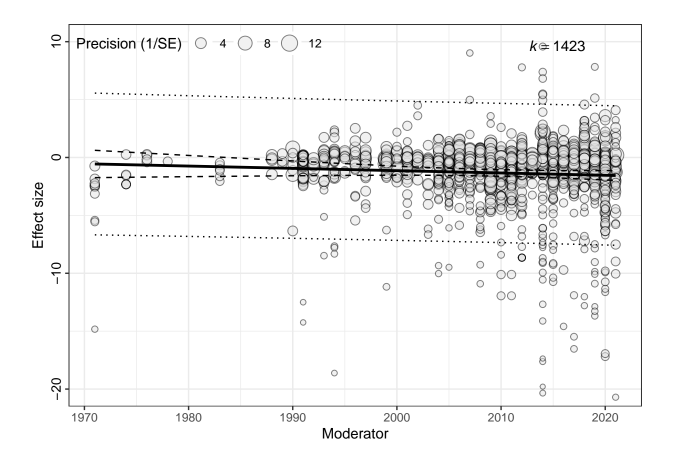
```
## [1] "<15"
               ">40"
                       "15-20" "20-25" "25-30" "30-35" "35-40"
table(rdata$bin.temp)
##
##
     <15
           >40 15-20 20-25 25-30 30-35 35-40
##
     112
            79
                 256
                       314
                             185
                                    349
                                          128
meta_trait_bintemp <- rma.mv(es, VCV_shared, mod = ~bin.temp -</pre>
    1, random = list(~1 | study_code, ~1 | obs), data = rdata,
    method = "REML")
summary(meta_trait_bintemp)
##
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
                                   AIC
                                                           AICc
##
                 Deviance
                                               BIC
       logLik
##
   -3731.3772
                7462.7544
                             7480.7544
                                         7528.0547
                                                     7480.8824
##
## Variance Components:
##
##
               estim
                         sqrt
                              nlvls
                                     fixed
                                                 factor
## sigma^2.1 5.5217
                      2.3498
                                 340
                                         no
                                             study code
## sigma^2.2 3.5140
                      1.8746
                                1423
                                         no
                                                     obs
##
## Test for Residual Heterogeneity:
## QE(df = 1416) = 36953.4464, p-val < .0001
##
## Test of Moderators (coefficients 1:7):
## QM(df = 7) = 295.5739, p-val < .0001
##
## Model Results:
##
##
                                                          ci.lb
                  estimate
                                         zval
                                                 pval
                                                                   ci.ub
                                 se
## bin.temp<15
                   -2.1735 0.2644
                                      -8.2216
                                               <.0001
                                                       -2.6916
                                                                 -1.6553
## bin.temp>40
                   -4.3977
                            0.3593
                                     -12.2402
                                               <.0001
                                                       -5.1018
                                                                 -3.6935
## bin.temp15-20
                   -1.3375 0.1864
                                      -7.1768
                                               <.0001
                                                       -1.7028
                                                                 -0.9722
## bin.temp20-25
                   -0.5414 0.1761
                                      -3.0741
                                               0.0021
                                                       -0.8866
                                                                 -0.1962
                                                                           **
## bin.temp25-30
                   -0.2829 0.2065
                                      -1.3701
                                               0.1707
                                                       -0.6876
                                                                  0.1218
## bin.temp30-35
                   -1.3021 0.1721
                                      -7.5644
                                               <.0001
                                                       -1.6395
                                                                 -0.9647
                                                                          ***
## bin.temp35-40
                   -3.0130 0.2457
                                     -12.2614 <.0001 -3.4946
                                                                 -2.5314
##
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```



Publication Bias.

We fit meta-regression model with year as a moderator to see whether there is an publication bias. In other words, is effect size strongly correlated with publication year. We found now evidence to suggest that year has an effect on the report effect sizes. A summary of the model is given below.

```
meta_year <- rma.mv(es, VCV_shared, mod = ~Publication.year,</pre>
    random = list(~1 | study_code, ~1 | obs), data = rdata, method = "REML")
summary(meta_year)
##
## Multivariate Meta-Analysis Model (k = 1423; method: REML)
##
                 Deviance
                                   AIC
                                               BIC
                                                          AICc
##
       logLik
  -3835.0508
                7670.1016
                            7678.1016
                                         7699.1381
                                                     7678.1299
##
##
## Variance Components:
##
##
               estim
                        sqrt
                              nlvls
                                     fixed
                                                 factor
## sigma^2.1
             5.0693
                      2.2515
                                340
                                             study_code
                                         no
              4.3197
                      2.0784
                                1423
## sigma^2.2
                                         no
                                                    obs
##
## Test for Residual Heterogeneity:
## QE(df = 1421) = 39431.7962, p-val < .0001
##
## Test of Moderators (coefficient 2):
## QM(df = 1) = 1.7366, p-val = 0.1876
##
## Model Results:
##
##
                     estimate
                                                              ci.lb
                                                                       ci.ub
                                     se
                                            zval
                                                    pval
                                                  0.2027
                                                                     98.2813
## intrcpt
                      38.7171
                               30.3905
                                          1.2740
                                                           -20.8471
## Publication.year
                      -0.0199
                                0.0151
                                        -1.3178 0.1876
                                                            -0.0496
                                                                      0.0097
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```



Sensitivty Analysis

Here, we perform a sensitivity analysis by removing the smallest and largest 2.5% of effect sizes.

The 2.5th and 97.5th percentiles are given below.

```
print(minq)

## 2.5%
## -14.17649

print(maxq)

## 97.5%
## 2.347279
```

The summary of the effective sizes is now

```
summary(sdata$es)
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -14.1203 -1.8264 -0.6221 -1.3539 0.0000 2.3381
```

We run the quadratic treatment temperature and the binned temperature models again with our new subsetted data i.e. only with data that is between the 2.5Th and 97.5Th percentile.

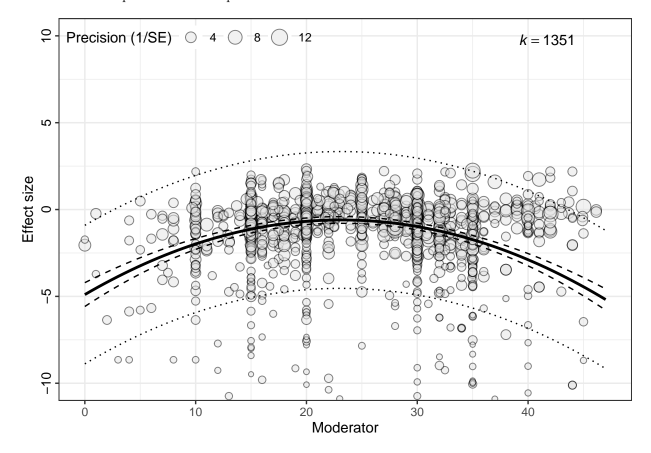
Treatment temperature as a quadratic effect (sessitivity analysis)

We re-create the variance-covariance matrix with our new substited data, which we name sdata. Then we run the meta analysis model again with treatment temperature as quadratic fixed effect,

```
# recreate vcv_shared matrix
VCV_shared_sa <- impute_covariance_matrix(vi = sdata$v, cluster = sdata$shared_control,
meta_sa_treat2 <- rma.mv(es, VCV_shared_sa, mod = ~poly(treattemp,</pre>
    degree = 2, raw = TRUE), random = list(~1 | study_code, ~1 |
    obs), data = sdata, method = "REML")
summary(meta_sa_treat2)
##
## Multivariate Meta-Analysis Model (k = 1351; method: REML)
##
                 Deviance
##
       logLik
                                   AIC
                                               BIC
                                                           AICc
  -2741.7575
                5483.5151
                             5493.5151
##
                                         5519.5470
                                                      5493.5598
##
## Variance Components:
##
##
               estim
                         sqrt
                              nlvls
                                     fixed
                                                  factor
## sigma^2.1
              2.1308
                                 334
                      1.4597
                                             study_code
                                         no
  sigma^2.2
              1.8965
                                                     obs
##
                                         no
##
## Test for Residual Heterogeneity:
## QE(df = 1348) = 29193.2105, p-val < .0001
## Test of Moderators (coefficients 2:3):
## QM(df = 2) = 173.9336, p-val < .0001
##
```

```
## Model Results:
##
##
                                                                  pval ci.lb ci.ub
                                                            zval
                                        estimate
                                                    se
## intrcpt
                                         -4.8959 0.3601 -13.5956 <.0001 -5.6017 -4.1901 ***
                                                        12.1579 <.0001 0.3118 0.4316 ***
## poly(treattemp, degree = 2, raw = TRUE)1
                                        0.3717 0.0306
## poly(treattemp, degree = 2, raw = TRUE)2 -0.0080 0.0006 -12.9093 <.0001 -0.0093 -0.0068 ***
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

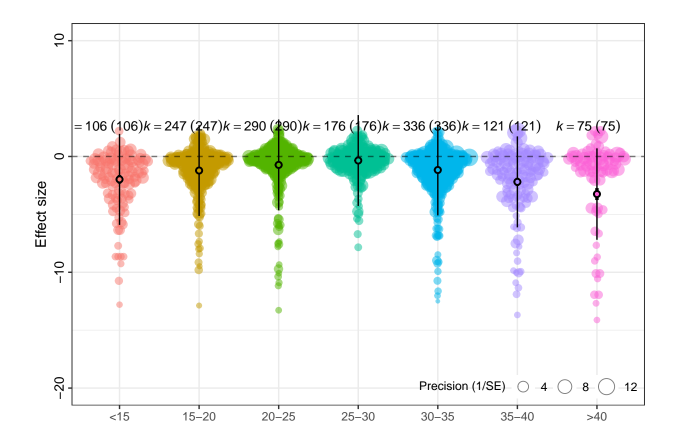
Below is a bubble plot of the fitted quadratic model to the subsetted data.



Modelling response with binned treatment temperatures

Lastly, we rerun the model where our effect sizes are categorised into one seven bins depending on the treatment temperature. The bins and the number of effect sizes in each bin of the new subsetted data are given below

```
table(sdata$bin.temp)
##
           >40 15-20 20-25 25-30 30-35 35-40
##
     <15
##
     106
                                    336
                 247
                        290
                              176
                                          121
meta_sa_bintemp <- rma.mv(es, VCV_shared_sa, mod = ~bin.temp -</pre>
    1, random = list(~1 | study_code, ~1 | obs), data = sdata,
    method = "REML")
summary(meta_sa_bintemp)
## Multivariate Meta-Analysis Model (k = 1351; method: REML)
##
##
       logLik
                 Deviance
                                   AIC
                                               BIC
                                                           AICc
##
  -2735.0546
                5470.1092
                             5488.1092
                                         5534.9399
                                                     5488.2442
##
## Variance Components:
##
##
               estim
                         sqrt
                              nlvls
                                      fixed
                                                 factor
## sigma^2.1
              2.0708
                      1.4390
                                 334
                                         no
                                             study_code
## sigma^2.2 1.9164 1.3843
                                1351
                                                     obs
                                         no
##
## Test for Residual Heterogeneity:
## QE(df = 1344) = 29044.8493, p-val < .0001
##
## Test of Moderators (coefficients 1:7):
## QM(df = 7) = 339.9024, p-val < .0001
##
## Model Results:
##
##
                                                          ci.lb
                  estimate
                                 se
                                         zval
                                                 pval
                                                                   ci.ub
                                               <.0001
                                                                 -1.6050
## bin.temp<15
                   -1.9772 0.1899
                                     -10.4114
                                                       -2.3494
                                                                          ***
## bin.temp>40
                   -3.2455 0.2628
                                     -12.3516
                                               <.0001
                                                       -3.7605
                                                                 -2.7305
## bin.temp15-20
                   -1.2133 0.1286
                                      -9.4342
                                               <.0001
                                                       -1.4654
                                                                 -0.9613
## bin.temp20-25
                   -0.7288 0.1223
                                      -5.9603
                                               <.0001
                                                        -0.9684
                                                                 -0.4891
                                                                 -0.0611
## bin.temp25-30
                   -0.3475 0.1462
                                      -2.3778
                                               0.0174
                                                       -0.6340
                                      -9.9500
                                                                          ***
## bin.temp30-35
                   -1.1681 0.1174
                                               <.0001
                                                       -1.3983
                                                                 -0.9380
## bin.temp35-40
                   -2.1928 0.1777
                                     -12.3429
                                               <.0001
                                                       -2.5410
                                                                 -1.8446
##
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```



Now we completely remove any study that has an effect size in the highest or lowest 2.5%.

```
summary(meta_sub_bintemp)
## Multivariate Meta-Analysis Model (k = 1253; method: REML)
##
##
       logLik
                 Deviance
                                   AIC
                                               BIC
                                                          AICc
   -2377.2698
                            4772.5396
                                         4818.6888
##
                4754.5396
                                                     4772.6852
##
## Variance Components:
##
##
               estim
                        sqrt nlvls fixed
                                                 factor
## sigma^2.1 1.4315
                      1.1965
                                310
                                         no
                                             study_code
## sigma^2.2 1.5062 1.2273
                                1253
                                                    obs
                                         no
##
## Test for Residual Heterogeneity:
## QE(df = 1246) = 25250.1123, p-val < .0001
## Test of Moderators (coefficients 1:7):
## QM(df = 7) = 304.1193, p-val < .0001
##
## Model Results:
```

```
##
                                        pval
               estimate
                           se
                                   zval
                                               ci.lb
                                                         ci.ub
## bin.temp<15
              -1.9033 0.1696 -11.2237 <.0001 -2.2356 -1.5709 ***
## bin.temp>40
                -2.1787 0.2545
                                -8.5598 <.0001 -2.6775 -1.6798 ***
## bin.temp15-20 -1.1631 0.1149 -10.1219 <.0001 -1.3884
                                                       -0.9379 ***
## bin.temp20-25
                                -5.7381 <.0001 -0.8392
               -0.6255 0.1090
                                                       -0.4119
                                -2.6632 0.0077 -0.6036 -0.0918
## bin.temp25-30 -0.3477 0.1306
## bin.temp30-35 -1.1636 0.1050 -11.0774 <.0001 -1.3695 -0.9577 ***
## bin.temp35-40 -1.7249 0.1641 -10.5094 <.0001 -2.0466 -1.4032 ***
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```