RCode - MHC class II genotype does not contribute towards the chemical encoding of heterozygosity and relatedness in a wild vertebrate population

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Packages

```
if (!require("magrittr", quietly = TRUE)) {
  install.packages("magrittr")
  library(magrittr)
} else {
  library(magrittr) # pipe operators
if (!require("tidyverse", quietly = TRUE)) {
  install.packages("tidyverse")
  library(tidyverse)
} else {
  library(tidyverse) # package collection for easy and pretty data science with R
if (!require("phyloseq", quietly = TRUE)) {
  if (!require("BiocManager", quietly = TRUE)) {
    install.packages("BiocManager")
  BiocManager::install(pkgs = "phyloseq")
 library(phyloseq) # phyloseq objects
} else {
  library(phyloseq) # phyloseq objects
if (!require("GCalignR", quietly = TRUE)) {
  install.packages("GCalignR")
 library(GCalignR)
} else {
  library(GCalignR) # handling/aligning chromatograms
if (!require("inbreedR", quietly = TRUE)) {
  install.packages("inbreedR")
  library(inbreedR)
} else {
  library(inbreedR) # population genetic analyses
```

```
}
if (!require("vegan", quietly = TRUE)) {
  install.packages("vegan")
  library(vegan)
} else {
  library(vegan) # statistical tools
if (!require("ggpubr", quietly = TRUE)) {
  install.packages("ggpubr")
 library(ggpubr)
} else {
  library(ggpubr) # ggplot grid and plot alignment functions
if (!require("ape", quietly = TRUE)) {
  install.packages("ape")
 library(ape)
} else {
  library(ape) # handling phylogenetic tree data
if (!require("performance", quietly = TRUE)) {
  install.packages("performance")
 library(performance)
} else {
  library(performance) # tools for models
if (!require("MuMIn", quietly = TRUE)) {
  install.packages("MuMIn")
  library(MuMIn)
} else {
  library(MuMIn) # tools for models
if (!require("partR2", quietly = TRUE)) {
  install.packages("partR2")
  library(partR2)
} else {
  library(partR2) # tools for models
}
if (!require("ggbeeswarm", quietly = TRUE)) {
  install.packages("ggbeeswarm")
  library(ggbeeswarm)
} else {
  library(ggbeeswarm)
}
# archived package as is dependent on `fts` package
```

```
# for execution of the code, users need to manually install Rtools to be able
# to install packages `Demerelate` and `fts`
library(fts)
library(Demerelate)
```

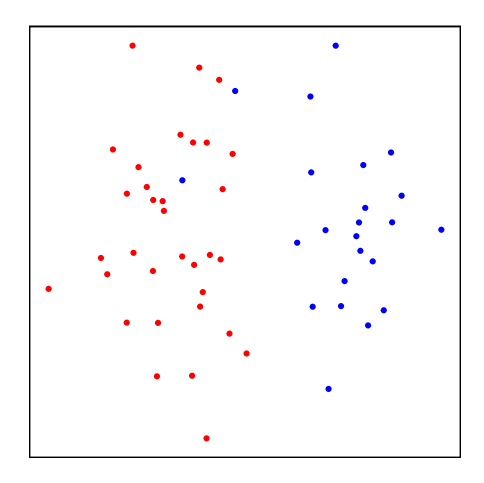
Packages for relatedness calculations

Not supported on newer versions of R, to execute code you must have Rtools installed on your machine in order to load older version of the Demerelate and fts package.

```
if (!require("remotes", quietly = TRUE)) {
  install.packages("remotes")
  library(remotes)
} else {
  library(remotes)
}
if (!require("fts", quietly = TRUE)) {
  install_version("fts", "0.9.9.2")
  library(fts)
} else {
  library(fts)
}
if (!require("Demerelate", quietly = TRUE)) {
  install version("Demerelate", "0.9.9.2")
  library(Demerelate)
} else {
  library(Demerelate)
options(digits = 12)
```

Subset scent data to correlate same individuals

```
scent %<>%
  .[keep_i, ] %>%
  `rownames<-`(meta$real_id)
## NMDS with reduced data
## GCalignR contains factors for the chemical dataset
data("peak factors")
peak_factors <- peak_factors[match(meta$id, rownames(peak_factors)),] %>%
 `rownames<-`(meta$real_id)</pre>
## keep order of rows consistent
scent <- scent[match(rownames(peak_factors),rownames(scent)),]</pre>
## NMDS using Bray-Curtis dissimilarities
scent_nmds.obj <- vegan::metaMDS(comm = scent, distance = "bray")</pre>
## get x and y coordinates
scent_nmds <- as.data.frame(scent_nmds.obj[["points"]])</pre>
## add the colony as a factor to each sample
scent_nmds <- cbind(scent_nmds,colony = peak_factors[["colony"]])</pre>
## quick plotting
scent_plot <- ggplot(data = scent_nmds,aes(MDS1,MDS2,color = colony)) +</pre>
 geom_point() +
 theme void() +
 scale_color_manual(values = c("blue","red")) +
 theme(panel.background = element_rect(colour = "black",
                                         linewidth = 1,
                                         fill = NA),
        aspect.ratio = 1,
        legend.position = "none")
scent_plot
```



Calculate MHC heterozygosity relatedness between individuals

```
## read in mhc genotype data
mhc_het_dat <- read.table("data/clone_mhc_het.txt")</pre>
## restructure `mhc_het_dat`to fit `Demerelate()::inputdata)
## id and colony as factors; alleles as integers or numeric
## otherwise `rxy`cannot handle computations
mhc_het_dat %<>%
  rownames_to_column(., var = "id") %>%
  # mutate(., a1 = str_pad(a1, 2, pad = "0")) %>%
  \# mutate(., a2 = str_pad(a2, 2, pad = "0")) %>%
  mutate(., colony = as.factor(rep("col", 56))) %>%
  mutate(., id = as.factor(id)) %>%
  .[,-4] %>%
  relocate(., colony, .before = a1)
  ## order mhc_het_dat$id after meta$real_id
  ## so data is consistently ordered same in all data.frames
## get matching indeces
id_index <- match(meta$real_id, mhc_het_dat$id)</pre>
## sort correspondingly
mhc_het_dat %<>% .[id_index,]
## calculate relatedness after Queller & Goodnight
mhc_relatedness_res <- Demerelate(inputdata = mhc_het_dat,</pre>
```

```
value = "rxy",
                                  object = T,
                                  NA.rm = F,
                                  Fis = F)
## Warning in Demerelate(inputdata = mhc_het_dat, value = "rxy", object = T, : Careful, bi-allelic mark
     Especially, rxy and ritland estimator are not defined when bi-allelic estimates are used with alle
     You should consider removing bi-allelics which tend to have very evenly distributed alleles or swi
     Be careful even if allele frequencies are not perfectly 0.5, during randomizations problems may oc
mhc_relatedness <- unlist(mhc_relatedness_res$Empirical_Relatedness)</pre>
## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals
relate_mat_mhc <- matrix(nrow = 56, ncol = 56)</pre>
## fill distance matrix row wise, thus fill upper.tri
relate_mat_mhc[upper.tri(relate_mat_mhc)] <- mhc_relatedness</pre>
## transpose to keep consistency with other distance matrices
relate_mat_mhc <- t(relate_mat_mhc)</pre>
relate_mat_mhc %<>% `colnames<-`(meta$real_id) %>% `rownames<-`(meta$real_id)
## vectorize again to identify whether relatedness pairs were consistent in the first place
a <- relate_mat_mhc %>% as.vector() %>% na.omit()
Create vectorized distance measurements for scent data
# bray-curtis distance measurement on scent profiles
scent_dist <- vegdist(scent) %>% as.matrix()
```

Generate UniFrac distances from MHC DQB II individual genotypes

scent_dist[upper.tri(scent_dist, diag = T)] <- NA
b <- scent_dist %>% as.vector() %>% na.omit()

```
# handle genotypes as otu table
phylo_mat <- read.table("data/phyloseq-mat.txt") %>%
    as.matrix()

# make sample names consistent
n <- match(meta$real_id, colnames(phylo_mat))

phylo_mat %<>% .[, n] %>%
    otu_table(., taxa_are_rows = T)

# create phylogenetic tree from file
phylo_tree <- ape::read.tree("data/unifrac_tree_p.nwk")

# merge into Formal class phyloseq
arga_phylseq <- merge_phyloseq(phylo_mat, phylo_tree)

# create UniFrac as genetic diversity measurement for single locus data
mhc_dqb2_ufrac <- UniFrac(arga_phylseq, weighted = F) %>%
    # distances to distance matrix
as.matrix()
```

```
# vectorize distances matrices
mhc_dqb2_ufrac[upper.tri(mhc_dqb2_ufrac, diag = T)] <- NA
c <- mhc_dqb2_ufrac %>% as.vector() %>% na.omit()
```

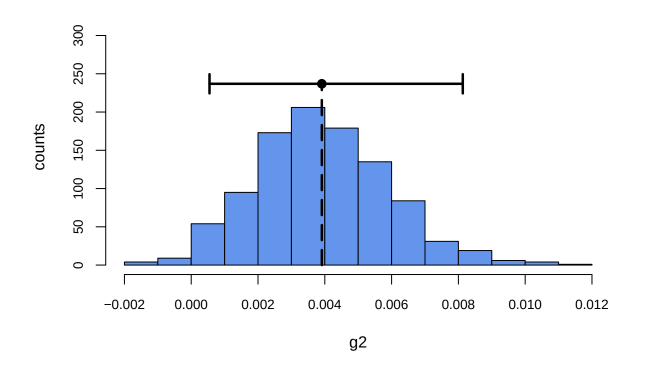
Calculate identity disequilibirum g2

```
msats_g2 <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t") %>%
    convert_raw()

g2 <- g2_microsats(msats_g2, nperm = 1000, nboot = 1000, CI = 0.95)

plot(g2, main = "Microsatellites",
    col = "cornflowerblue", cex.axis=0.85)</pre>
```

Microsatellites



Calculate microsatellite relatedness values

create data.frame in correspondence to Demerelate input format

```
# read in genotype data table
msats_df <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t")

# update data.frame with additional info
# "delete" colony info, otherwise relatedness is only calculated for individuals
# within their own colonies -> no complete pairwise comparison
msats_df <- cbind(id = as.factor(rownames(msats_df)),</pre>
```

Calculate relatedness of individuals based on Queller & Goodnight

Coerce output to a vector

```
relatedness <- unlist(relatedness_results$Empirical_Relatedness)

## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals
relate_mat <- matrix(nrow = 56, ncol = 56)

## fill distance matrix row wise, thus fill upper.tri
relate_mat[upper.tri(relate_mat)] <- relatedness

## transpose to keep consistency with other distance matrices
relate_mat <- t(relate_mat)
relate_mat %<>% `colnames<-`(meta$real_id) %>% `rownames<-`(meta$real_id)

## vectorize again to identify whether relatedness pairs were consistent in the first place
d <- relate_mat %>% as.vector() %>% na.omit()
```

Analyse Odour and genetic association by MHC DQB II and neutral genomic background

Create data.frame to plot in ggplot2

```
## substitute once tested correctly
## scent_mds shall contain similarity values but `b` contains
## dissimilarity values based on Bray-Curtis -> substracting
## dissmilarities from 1 returns similarities

model_rel.df <- cbind(mhc_rel = a, scent_mds = 1-b, ufrac = c, rel = d) %>%
    as.data.frame()
```

Custom theme to make plot aesthetics consistent

Plot odour by mhc similarity

Plot odour by relatedness

Model odour relationship on MHC and neutral genetic background

Pool underlying data dependencies

Create a function that generates pairwise variables in a systematic matter for pairwise comparisons

```
create_pair_vars <-function(row_cross, col_cross, split_vars = F){
   require(stringr)</pre>
```

```
rc <- row cross
  cc <- col_cross
  # create empy matrix
  # keep row and col names from existing distance matrices
  empty_mat <- matrix(nrow = length(rc),</pre>
                      ncol = length(cc)) %>%
    `colnames<-`(cc) %>%
    `rownames<-`(rc)
  # fill each matrix i,j-th cell with the crossing from their corresponding
  \# i-th rowname and j-th colname
  for (i in 1:dim(empty_mat)[1]) {
    for (j in 1:dim(empty_mat)[2]) {
      empty_mat[i,j] <- paste0(rc[i], "/", cc[j])</pre>
    } # end j
  } # end i
  # delete `upper.tri()` of `empty_mat` to resemble structure of the other
  # distance matrices in use
  empty_mat[upper.tri(empty_mat, diag = T)] <- NA</pre>
  pair_vars <- empty_mat %>% as.vector() %>% na.omit()
  # split `pair_vars` if needed
  if (split_vars == T) {
    pair_vars1 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][1]
    pair_vars2 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][2]
                          })
    pair_vars_split <- list(pair_variable1 = pair_vars1,</pre>
                             pair_variable2 = pair_vars2)
    return(pair_vars_split)
  } else {
    return(pair_vars)
} #end create_pair_vars
```

Helper function to combine double entries

```
## for x, overwrite specified replacer with specified value
f <- function(x, replacer, overwrite){
  if (x == replacer) {
    x <- overwrite
  } else {
    x <- x
  }
}</pre>
```

Transform model variables

```
agePaired <- create_pair_vars(row_cross = meta$maturity,</pre>
                                col_cross = meta$maturity) %>%
  sapply(., f, "P/M", "M/P")
colonyPaired <- create_pair_vars(row_cross = meta$colony,</pre>
                                   col_cross = meta$colony) %>%
  sapply(., f, "FWB/SSB", "SSB/FWB")
colonyID1 <- create_pair_vars(row_cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[1] %>%
  unlist() %>%
 paste0("f", .) %>%
  as.vector()
colonyID2 <- create_pair_vars(row_cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[2] %>%
 unlist() %>%
 paste0("f", .) %>%
  as.vector()
colonyBool <- ifelse(colonyID1 == colonyID2, 1, 0)</pre>
familyPaired <- create_pair_vars(row_cross = meta$family,</pre>
                                   col_cross = meta$family)
familyID1 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[1] %>%
  unlist() %>%
 paste0("f", .) %>%
  as.vector()
familyID2 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[2] %>%
  unlist() %>%
 paste0("f", .) %>%
 as.vector()
pairID1 <- create_pair_vars(row_cross = meta$real_id,</pre>
                             col_cross = meta$real_id,
```

Update data.frame with model variables

Color Chemical similarity by same or different beach

'colonyBool' encodes whether individual from same colonies (SSB vs SSB and FWB vs FWB) are compared or from different colonies

Chemical similarity models

```
# mhc
a1 <- lmerTest::lmer(scent_mds ~ ufrac + colonyBool + agePaired + (1|familyBool) +
                       (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# relatedness
a2 <- lmerTest::lmer(scent_mds ~ rel + colonyBool + agePaired + (1|familyBool) + (1|pairID1) +
                       (1|pairID2),
                     data = model_rel.df)
# mhc & relatedness
a3 <- lmerTest::lmer(scent_mds ~ rel + ufrac + colonyBool + agePaired + (1|familyBool) +
                       (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# no genetic effect
a4 <- lmerTest::lmer(scent_mds ~ colonyBool + agePaired + (1|familyBool) + (1|pairID1) +
                       (1|pairID2),
                     data = model_rel.df)
# compare model performance scores
compare_performance(a1, a2, a3, a4, rank = T) %>%
  arrange(Name)
```

```
## # Comparison of Model Performance Indices
##
                   Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc weight
## -----
## a1 | lmerModLmerTest | 0.696 | 0.140 | 0.646 | 0.060 | 0.062 | ## a2 | lmerModLmerTest | 0.690 | 0.143 | 0.638 | 0.060 | 0.062 | ## a3 | lmerModLmerTest | 0.689 | 0.144 | 0.636 | 0.060 | 0.062 | ## a4 | lmerModLmerTest | 0.698 | 0.139 | 0.649 | 0.060 | 0.062 |
                                                                                    0.198 |
                                                                                   0.212
                                                                                    0.082 l
                                                                                0.508 |
summary(a2)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ rel + colonyBool + agePaired + (1 | familyBool) +
##
       (1 | pairID1) + (1 | pairID2)
##
      Data: model_rel.df
##
## REML criterion at convergence: -3928.2
## Scaled residuals:
            Min
                         1Q
                                 Median
                                                     30
## -3.108444259 -0.690869875 -0.068233408 0.583996354 3.762633304
##
## Random effects:
## Groups Name
                           Variance
                                        Std.Dev.
            (Intercept) 0.00136165106 0.0369005564
## pairID2
## pairID1 (Intercept) 0.00132283514 0.0363708006
## familyBool (Intercept) 0.00404648919 0.0636120208
                           0.00381948877 0.0618020126
## Residual
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
##
## Fixed effects:
                      Estimate
                                  Std. Error
                                                           df t value Pr(>|t|)
## (Intercept) 3.15186662e-01 4.73683692e-02 1.14156516e+00 6.65395 0.074931 .
               8.71026170e-03 2.08593513e-02 1.33740535e+03 0.41757 0.676328
## colonyBool1 8.37421175e-02 8.06181663e-03 1.20111837e+02 10.38750 < 2e-16 ***
## agePairedM/P 2.39466928e-03 8.06276850e-03 1.28046621e+02 0.29700 0.766945
## agePairedP/P 1.21988991e-02 1.53157945e-02 1.03077423e+02 0.79649 0.427577
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
              (Intr) rel clnyB1 agPM/P
##
## rel
               -0.090
## colonyBool1 -0.152 0.012
## agePairdM/P -0.145 -0.016 -0.023
## agePairdP/P -0.137 -0.004 -0.026 0.915
summary(a4)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ colonyBool + agePaired + (1 | familyBool) + (1 |
##
       pairID1) + (1 | pairID2)
```

0.19 0.21

0.08

0.51

Data: model_rel.df

##

```
## REML criterion at convergence: -3933.9
## Scaled residuals:
                         1Q
                                  Median
## -3.113665362 -0.687292046 -0.067704903 0.586076727 3.745905490
## Random effects:
## Groups
              Name
                          Variance
                                        Std.Dev.
## pairID2
              (Intercept) 0.00136551721 0.0369529053
## pairID1
              (Intercept) 0.00132698089 0.0364277489
## familyBool (Intercept) 0.00435745709 0.0660110377
## Residual
                          0.00381639653 0.0617769903
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
##
## Fixed effects:
##
                     Estimate
                                  Std. Error
                                                         df t value Pr(>|t|)
## (Intercept) 3.17056257e-01 4.87992888e-02 1.13632796e+00 6.49715 0.077614 .
## colonyBool1 8.36965038e-02 8.07029419e-03 1.20281473e+02 10.37094 < 2e-16 ***
## agePairedM/P 2.44412815e-03 8.07041285e-03 1.28202135e+02 0.30285 0.762495
## agePairedP/P 1.22231134e-02 1.53347232e-02 1.03314754e+02 0.79709 0.427228
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
              (Intr) clnyB1 agPM/P
## colonyBool1 -0.147
## agePairdM/P -0.142 -0.023
## agePairdP/P -0.134 -0.026 0.915
# if interested
# check model performance by
# check_model(a2)
```

Correlations of genetic main effects

```
# correlation of ufrac and relatedness
u_r_model1 <- lmerTest::lmer(ufrac ~ rel + (1|pairID1) + (1|pairID2),
                             data = model_rel.df)
summary(u_r_model1)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: ufrac ~ rel + (1 | pairID1) + (1 | pairID2)
##
      Data: model_rel.df
##
## REML criterion at convergence: -663.8
##
## Scaled residuals:
##
                          1Q
                                   Median
## -4.441714289 -0.385476875 0.272592692 0.681993885 1.501197368
##
## Random effects:
## Groups Name
                                       Std.Dev.
                         Variance
```

```
## pairID1 (Intercept) 0.00216331204 0.0465114183
## pairID2 (Intercept) 0.00101136839 0.0318020186
                       0.03589648773 0.1894636844
## Residual
## Number of obs: 1540, groups: pairID1, 55; pairID2, 55
## Fixed effects:
                    Estimate
                                  Std. Error
                                                         df t value Pr(>|t|)
## (Intercept) 7.97297459e-01 9.69913681e-03 6.44203957e+01 82.20293 < 2.22e-16
## rel
              -2.38911481e-01 5.12089926e-02 1.42412461e+03 -4.66542 3.3692e-06
##
## (Intercept) ***
## rel
              ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
      (Intr)
## rel -0.009
u_r_model2 <- lmerTest::lmer(ufrac ~ rel + (1|pairID1) + (1|pairID2) + (1|familyBool),
                           data = model_rel.df)
summary(u_r_model2)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: ufrac ~ rel + (1 | pairID1) + (1 | pairID2) + (1 | familyBool)
##
     Data: model rel.df
##
## REML criterion at convergence: -674.6
## Scaled residuals:
                        1Q
                                 Median
## -4.473364209 -0.379228923 0.272204191 0.676906270 2.150144511
## Random effects:
                         Variance
                                       Std.Dev.
## Groups
              Name
              (Intercept) 0.00215194602 0.0463890722
## pairID1
              (Intercept) 0.00107608003 0.0328036587
   pairID2
## familyBool (Intercept) 0.01743074553 0.1320255488
                         0.03554491277 0.1885335853
## Residual
## Number of obs: 1540, groups: pairID1, 55; pairID2, 55; familyBool, 2
## Fixed effects:
##
                                Std. Error
                                                      df t value Pr(>|t|)
                   Estimate
                                            0.9921110956 7.34418 0.087397 .
                              0.0967753027
## (Intercept)
               0.7107349721
## rel
               ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
      (Intr)
## rel -0.126
```

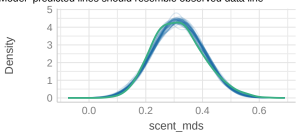
```
compare_performance(u_r_model1, u_r_model2, rank = T)
## Some of the nested models seem to be identical and probably only vary in
## their random effects.
## # Comparison of Model Performance Indices
                         Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc
## Name
## u r model2 | lmerModLmerTest | 0.369 |
                                                 0.003 | 0.368 | 0.185 | 0.189 |
                                                                                        0.874 l
## u r model1 | lmerModLmerTest |
                                    0.095 | 0.015 | 0.081 | 0.186 | 0.189 |
                                                                                        0.126 l
u_r_model3 <- lmerTest::lmer(ufrac ~ rel + colonyBool + (1|pairID1) + (1|pairID2) + (1|familyBool),
                           data = model_rel.df)
summary(u_r_model3)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: ufrac ~ rel + colonyBool + (1 | pairID1) + (1 | pairID2) + (1 |
##
      familyBool)
##
      Data: model_rel.df
##
## REML criterion at convergence: -667.8
##
## Scaled residuals:
##
           Min
                         1Q
                                  Median
                                                   30
                                                               Max
## -4.473094103 -0.378995809 0.272905728 0.676506794 2.148306764
##
## Random effects:
## Groups
                          Variance
                                        Std.Dev.
            (Intercept) 0.00217394338 0.0466255657
## pairID1
## pairID2
              (Intercept) 0.00108902249 0.0330003408
## familyBool (Intercept) 0.01740183944 0.1319160318
                          0.03555503513 0.1885604283
## Number of obs: 1540, groups: pairID1, 55; pairID2, 55; familyBool, 2
## Fixed effects:
                     Estimate
                                   Std. Error
                                                           df t value Pr(>|t|)
## (Intercept) 7.11336570e-01 9.72959228e-02 1.01443951e+00 7.31106 0.084313 .
              -1.26595370e-01 5.94527269e-02 9.16826722e+02 -2.12935 0.033492 *
## colonyBool1 -7.42382735e-04 1.31644657e-02 2.09024905e+02 -0.05639 0.955083
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
              (Intr) rel
## rel
              -0.126
## colonyBool1 -0.110 0.013
compare_performance(u_r_model1, u_r_model2, u_r_model3, rank = T)
## Some of the nested models seem to be identical and probably only vary in
    their random effects.
## # Comparison of Model Performance Indices
##
```

```
| Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc
## -----
## u r model2 | lmerModLmerTest | 0.369 |
                                            0.003 | 0.368 | 0.185 | 0.189 |
                                                                               0.661 |
## u_r_model3 | lmerModLmerTest |
                                0.369 |
                                            0.003 | 0.368 | 0.185 | 0.189 |
                                                                              0.243 |
## u_r_model1 | lmerModLmerTest |
                                  0.095 |
                                             0.015 | 0.081 | 0.186 | 0.189 |
                                                                               0.096 |
summary(u_r_model2)# colony effect unsubstantial but family important!
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: ufrac ~ rel + (1 | pairID1) + (1 | pairID2) + (1 | familyBool)
     Data: model_rel.df
## REML criterion at convergence: -674.6
## Scaled residuals:
                      1Q
                              Median
## -4.473364209 -0.379228923 0.272204191 0.676906270 2.150144511
## Random effects:
## Groups
                       Variance
                                   Std.Dev.
            (Intercept) 0.00215194602 0.0463890722
## pairID1
## pairID2 (Intercept) 0.00107608003 0.0328036587
## familyBool (Intercept) 0.01743074553 0.1320255488
                       0.03554491277 0.1885335853
## Number of obs: 1540, groups: pairID1, 55; pairID2, 55; familyBool, 2
## Fixed effects:
                                                  df t value Pr(>|t|)
                  Estimate
                            Std. Error
## (Intercept) 0.7107349721 0.0967753027 0.9921110956 7.34418 0.087397 .
            ## rel
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
      (Intr)
## rel -0.126
(aov_u_r <- anova(u_r_model2))</pre>
## Type III Analysis of Variance Table with Satterthwaite's method
                     Mean Sq NumDF
                                       DenDF F value Pr(>F)
## rel 0.1616032287 0.1616032287 1 922.8588459 4.54645 0.033251 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Adressing collinearity
# use performance tools to check model goodness
# model with all genetic effects that were
# evaluated in chemical similarity associations:
```

check model(a3)

Posterior Predictive Check

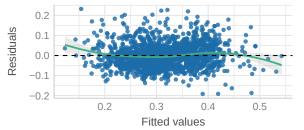
Model-predicted lines should resemble observed data line



— Observed data — Model-predicted data

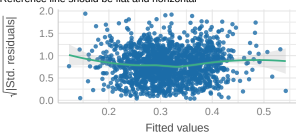
Linearity

Reference line should be flat and horizontal



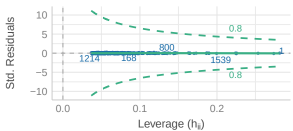
Homogeneity of Variance

Reference line should be flat and horizontal

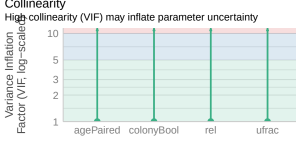


Influential Observations

Points should be inside the contour lines

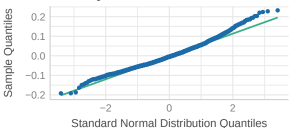


Collinearity



Normality of Residuals

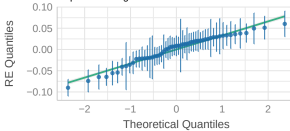
Dots should fall along the line



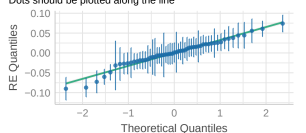
Low (< 5)

Normality of Random Effects (pairID2)

Dots should be plotted along the line

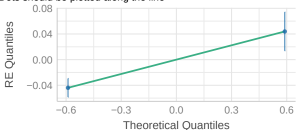


Normality of Random Effects (pairID1) Dots should be plotted along the line



Normality of Random Effects (familyBool)

Dots should be plotted along the line

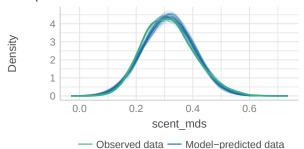


```
# check by re-running a3 model with residuals for relatedness effect
res_u_r <- residuals(u_r_model2)
temp_df <- cbind(model_rel.df, col_residuals = res_u_r)</pre>
a3_residual <- lmerTest::lmer(scent_mds ~ col_residuals + rel + colonyBool + agePaired +
                                (1|familyBool) + (1|pairID1) + (1|pairID2),
                              data = temp_df)
summary(a3_residual)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ col_residuals + rel + colonyBool + agePaired + (1 |
       familyBool) + (1 | pairID1) + (1 | pairID2)
##
      Data: temp_df
##
## REML criterion at convergence: -3920.6
##
## Scaled residuals:
           Min
                          1Q
                                   Median
                                                    3Q
                                                                Max
## -3.103258827 -0.690769473 -0.068686908 0.585558921 3.761156979
##
## Random effects:
                                         Std.Dev.
## Groups
              Name
                           Variance
## pairID2
               (Intercept) 0.00136213257 0.0369070802
               (Intercept) 0.00132201584 0.0363595358
## pairID1
## familyBool (Intercept) 0.00404272164 0.0635824004
                           0.00382206362 0.0618228406
## Residual
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
##
## Fixed effects:
##
                        Estimate
                                      Std. Error
                                                              df t value Pr(>|t|)
## (Intercept)
                 3.15169410e-01 4.73489770e-02 1.14124101e+00 6.65631 0.074941
## col residuals -1.72524676e-03 8.62343567e-03 1.43952773e+03 -0.20006 0.841458
## rel
                 8.77922600e-03 2.08681133e-02 1.33588284e+03 0.42070 0.674041
                 8.37532200e-02 8.06204794e-03 1.20109868e+02 10.38858 < 2e-16
## colonyBool1
## agePairedM/P
                 2.38288843e-03 8.06317648e-03 1.28076263e+02 0.29553 0.768070
## agePairedP/P
                 1.21929643e-02 1.53155887e-02 1.03075963e+02 0.79611 0.427795
##
## (Intercept)
## col_residuals
## rel
## colonyBool1
## agePairedM/P
## agePairedP/P
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
                                    clnyB1 agPM/P
               (Intr) cl rsd rel
## col residls 0.002
## rel
              -0.090 -0.014
## colonyBool1 -0.152 -0.008 0.012
```

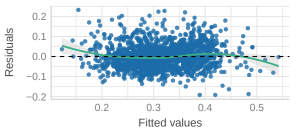
check_model(a3_residual)

Posterior Predictive Check

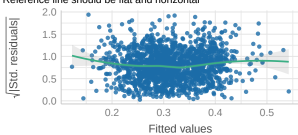
Model-predicted lines should resemble observed data line



Linearity Reference line should be flat and horizontal

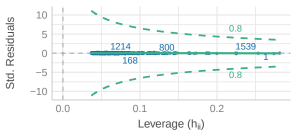


Homogeneity of Variance Reference line should be flat and horizontal

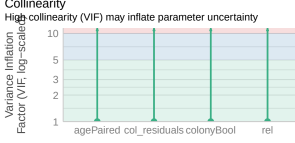


Influential Observations

Points should be inside the contour lines

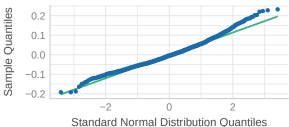


Collinearity



Normality of Residuals

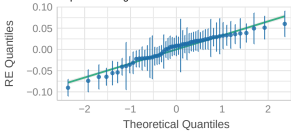
Dots should fall along the line



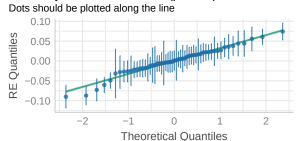
Low (< 5)

Normality of Random Effects (pairID2)

Dots should be plotted along the line

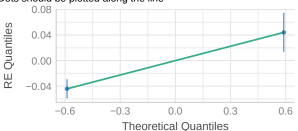


Normality of Random Effects (pairID1)



Normality of Random Effects (familyBool)

Dots should be plotted along the line

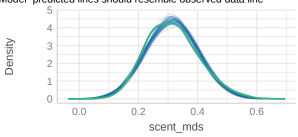


Neither model, performs noticeably better, and by VIC both models (with and without residuals fitted) show no high amount of collinearity. However, 'a4' is within this same goodness and while retaining the same information with less fixed effects, thus being more parsimonious.

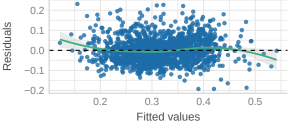
check_model(a4)

Posterior Predictive Check

Model-predicted lines should resemble observed data line



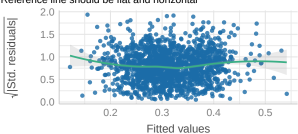
Reference line should be flat and horizontal



Observed data — Model-predicted data

Homogeneity of Variance

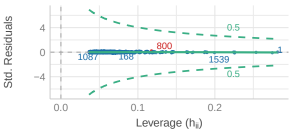
Reference line should be flat and horizontal



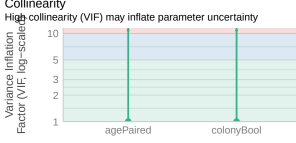
Influential Observations

Linearity

Points should be inside the contour lines

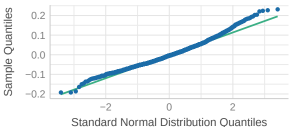


Collinearity



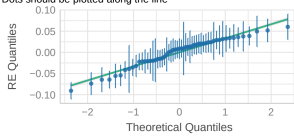
Normality of Residuals

Dots should fall along the line

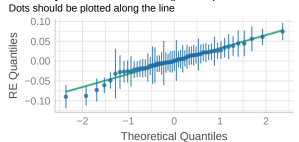


Low (< 5)

Normality of Random Effects (pairID2) Dots should be plotted along the line

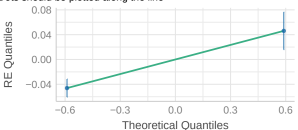


Normality of Random Effects (pairID1)



Normality of Random Effects (familyBool)

Dots should be plotted along the line



Model relationship between chemical diversity and mhc plus msats diversity update data frame with meta data

Include information about MHC heterozygosity, sMLH from microsatellite data and chemical diversity by number of compounds per individual

```
scent.abs <- ifelse(scent != 0, 1, 0)</pre>
compound_n <- apply(scent.abs, 1, sum)</pre>
names(compound_n) == meta$real_id
# read in heterzygosity information
het table <- read.table("data/arga mhc het.txt", sep = "\t")
# keep names consistent
match_het <- match(meta$real_id, rownames(het_table))</pre>
het_table %<>% .[match_het,]
# generate sMLH with microsatellite data
# table is pre-prepped, thus rows correspond to same individuals in meta data
smlh_res <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t") %>%
 # convert to inbreedR format
 convert_raw() %>%
 # generate sMLH
 sMLH()
meta %<>% cbind(., compound_n = compound_n,
           mhc_het = het_table$het,
            smlh = smlh res)
meta %<>% mutate(
 real_id = as.factor(real_id),
 colony = as.factor(colony),
 maturity = as.factor(maturity),
 family = as.factor(family)
```

General meta data information

```
# 32 adult female individuals
meta_mom <- meta %>% filter(maturity == "M")

# check n for each beach colony
n_fwb_mom <- meta_mom %>% filter(colony == "FWB") %>% dim() %>% .[1]
n_ssb_mom <- meta_mom %>% filter(colony == "SSB") %>% dim() %>% .[1]

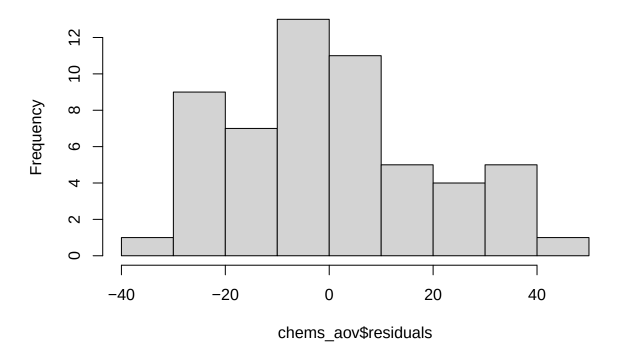
# test for difference in number of chemicals by beach
chems_fwb_mom <- meta_mom %>% filter(., colony == "FWB") %>%
    select(compound_n) %>%
    unlist()
```

```
chems_ssb_mom <- meta_mom %>% filter(., colony == "SSB") %>%
  select(compound_n) %>%
 unlist()
(colony_t_test_mom <- t.test(chems_fwb_mom, chems_ssb_mom))</pre>
##
##
  Welch Two Sample t-test
## data: chems_fwb_mom and chems_ssb_mom
## t = -1.522037389, df = 22.29327201, p-value = 0.142058931
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -29.16483741389
                      4.46483741389
## sample estimates:
## mean of x mean of y
       52.15
                 64.50
##
# 24 pups (mixed sexes)
meta_pup <- meta %>% filter(maturity == "P")
# check n for each beach colony
n_fwb_pup <- meta_pup %>% filter(colony == "FWB") %>% dim() %>% .[1]
n_ssb_pup <- meta_pup %>% filter(colony == "SSB") %>% dim() %>% .[1]
# test for difference in number of chemicals by beach
chems_fwb_pup <- meta_pup %>% filter(., colony == "FWB") %>%
  select(compound_n) %>%
  unlist()
chems_ssb_pup <- meta_pup %>% filter(., colony == "SSB") %>%
  select(compound_n) %>%
  unlist()
(colony_t_test_pup <- t.test(chems_fwb_pup, chems_ssb_pup))</pre>
##
## Welch Two Sample t-test
##
## data: chems_fwb_pup and chems_ssb_pup
## t = 1.175783211, df = 20.29290702, p-value = 0.253292061
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -6.90357099378 24.77769686791
## sample estimates:
##
       mean of x
                     mean of y
## 58.8461538462 49.9090909091
# standard error function
se <- function(x) sd(x)/sqrt(length(x))</pre>
mean_chems_by_colony <- list(chems_fwb_mom, chems_fwb_pup, chems_ssb_mom, chems_ssb_pup)
sapply(mean_chems_by_colony, mean)
```

[1] 52.1500000000 58.8461538462 64.5000000000 49.9090909091

```
sapply(mean_chems_by_colony, se)
## [1] 4.80037005153 6.31324021184 6.54182348779 4.23288971884
mean(meta$compound_n)
## [1] 55.9107142857
se(meta$compound_n)
## [1] 2.80714265061
# neither in pups nor moms compound_n differs for colony, calculate anova
# check whether compound number varies for different ages and different colonies
chems_aov <- aov(compound_n ~ maturity * colony, data = meta)</pre>
summary(chems_aov)
                                     Mean Sq F value Pr(>F)
##
                  Df
                           Sum Sq
                                   56.584821 0.13023 0.71966
## maturity
                   1
                        56.58482
                        115.19930 115.199303 0.26513 0.60880
## colony
                   1
## maturity:colony 1 1504.61805 1504.618048 3.46285 0.06842 .
## Residuals
                  52 22594.15140 434.502912
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
hist(chems_aov$residuals)
```

Histogram of chems_aov\$residuals



```
# post-hoc analysis
TukeyHSD(chems_aov)
##
     Tukey multiple comparisons of means
##
      95% family-wise confidence level
##
## Fit: aov(formula = compound_n ~ maturity * colony, data = meta)
##
## $maturity
##
                          lwr
                                        upr
## P-M -2.03125 -13.326105701 9.26360570104 0.7196568197
##
## $colony
##
                    diff
                                    lwr
                                                  upr
                                                               p adj
## SSB-FWB 2.90513833992 -8.45648556259 14.2667622424 0.610058289387
## $`maturity:colony`
##
                          diff
                                           lwr
                                                                       p adj
## P:FWB-M:FWB 6.69615384615 -13.01371483797 26.40602253028 0.803961243091
## M:SSB-M:FWB 12.35000000000 -7.85146210548 32.55146210548 0.375145035385
## P:SSB-M:FWB -2.24090909091 -23.00834798997 18.52652980816 0.991714931867
## M:SSB-P:FWB 5.65384615385 -16.49347177283 27.80116408052 0.905141795957
## P:SSB-P:FWB -8.93706293706 -31.60181836421 13.72769249008 0.723034927926
## P:SSB-M:SSB -14.59090909091 -37.68444251969 8.50262433787 0.346056221246
#### Permanova
perm_chem_data <- vegan::adonis2(</pre>
  # term
  scent ~ colony + family + smlh + maturity,
  by = "terms",
  # data for grouping
  data = meta)
perm_chem_data
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## vegan::adonis2(formula = scent ~ colony + family + smlh + maturity, data = meta, by = "terms")
           Df
                   SumOfSqs
                                      R2
                                                F Pr(>F)
           1 1.671427973 0.1273987963 11.47634 0.001 ***
## colony
            34 8.546158183 0.6514012466 1.72587
                                                  0.001 ***
## family
            1 0.120713508 0.0092009682 0.82884 0.717
## maturity 1 0.159812162 0.0121811274
                                         1.09730 0.338
## Residual 18 2.621540965 0.1998178615
           55 13.119652790 1.00000000000
## Total
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Compare chemical diversity models

Correlate Chemical diversity per sample with their sMLH and MHC, respectively. Also accounting maturity and family as fixed and random effect.

```
b1 <- lmerTest::lmer(compound_n ~ mhc_het + maturity + colony +(1 family),
                     data = meta)
b2 <- lmerTest::lmer(compound_n ~ smlh + maturity + colony + (1|family),
                     data = meta)
b3 <- lmerTest::lmer(compound_n ~ mhc_het + smlh + maturity + colony + (1 | family),
                    data = meta)
b4 <- lmerTest::lmer(compound_n ~ maturity + colony + (1|family),
                    data = meta)
compare_performance(b1, b2, b3, b4, rank = T) %>% arrange(Name)
## # Comparison of Model Performance Indices
##
                   Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc weigh
## Name |
        | lmerModLmerTest |
                                0.747 |
                                             0.006 | 0.746 | 7.248 | 10.847 |
                                                                                    0.005 |
## b2
       | lmerModLmerTest |
                                0.772 |
                                             0.119 | 0.741 | 6.696 | 9.994 |
                                                                                    0.713 |
       | lmerModLmerTest |
                                0.769 |
                                             0.117 | 0.738 | 6.706 | 10.127 |
                                                                                    0.269 |
## b3
      | lmerModLmerTest |
                                0.752 |
                                             0.006 | 0.751 | 7.202 | 10.670 |
                                                                                    0.014
## b4
summary(b2)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: compound_n ~ smlh + maturity + colony + (1 | family)
      Data: meta
##
## REML criterion at convergence: 450.6
##
## Scaled residuals:
##
                           1Q
                                     Median
## -1.1174803761 -0.5102355060 -0.0397205379 0.2311200062 2.1221413149
##
## Random effects:
## Groups
                        Variance
                                    Std.Dev.
            (Intercept) 285.8932945 16.90837942
## family
                         99.8834741 9.99417201
## Number of obs: 56, groups: family, 36
##
## Fixed effects:
                  Estimate
                            Std. Error
                                                  df t value Pr(>|t|)
## (Intercept) -21.15641099 24.20114640 44.84179770 -0.87419 0.3866725
               76.53192951 24.10779559 43.94521713 3.17457 0.0027408 **
## smlh
## maturityP
               -4.62266359 3.17801210 24.39984244 -1.45458 0.1585308
## colonySSB
                4.10983380 6.36424962 34.10616354 0.64577 0.5227505
```

0.0

0.7

0.2

0.0

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

```
##
## Correlation of Fixed Effects:
##
            (Intr) smlh
            -0.984
## smlh
## maturityP 0.257 -0.303
## colonySSB -0.165 0.055 -0.076
Correlate zygosity effects
smlh_het_m1 <- lmerTest::lmer(smlh ~ mhc_het + (1|family), data = meta)</pre>
summary(smlh het m1)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: smlh ~ mhc_het + (1 | family)
     Data: meta
##
## REML criterion at convergence: -98.4
## Scaled residuals:
##
           Min
                         1Q
                                  Median
                                                  3Q
                                                              Max
## -2.193649022 -0.656470901 0.102448217 0.718635430 1.907640611
## Random effects:
## Groups
                        Variance
                                     Std.Dev.
            (Intercept) 0.00131322013 0.0362383793
## family
                        0.00720048654 0.0848556806
## Residual
## Number of obs: 56, groups: family, 36
## Fixed effects:
                               Std. Error
                                                    df t value Pr(>|t|)
                   Estimate
## (Intercept) 1.0286614945 0.0268377389 51.3350533536 38.32892 < 2e-16 ***
              ## mhc_het
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
          (Intr)
## mhc het -0.876
# check performance for including colony as fixed effect, as well
smlh_het_m2 <- lmerTest::lmer(smlh ~ mhc_het + colony + (1|family), data = meta)</pre>
summary(smlh_het_m2)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: smlh ~ mhc_het + colony + (1 | family)
##
     Data: meta
## REML criterion at convergence: -93.1
## Scaled residuals:
                                  Median
           Min
                         1Q
                                                  3Q
## -2.095136459 -0.653315112 0.113695726 0.687418314 1.847436161
## Random effects:
```

```
## Fixed effects:
                   Estimate
                              Std. Error
                                                   df t value Pr(>|t|)
## (Intercept) 1.03101391140 0.02871644361 45.35909433637 35.90326
                                                                <2e-16 ***
## mhc het
             -0.03679744212  0.03054090182  52.11941131342  -1.20486  0.2337
## colonySSB -0.00838946614 0.02680960458 26.33503941544 -0.31293 0.7568
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
           (Intr) mhc_ht
##
## mhc_het -0.799
## colonySSB -0.329 -0.074
(aov <- anova(smlh_het_m2))</pre>
## Type III Analysis of Variance Table with Satterthwaite's method
                              Mean Sq NumDF DenDF F value Pr(>F)
##
                 Sum Sq
## mhc_het 0.010366750137 0.010366750137 1 52.11941131 1.45168 0.2337
## colony 0.000699292576 0.000699292576
                                        1 26.33503942 0.09792 0.7568
compare_performance(smlh_het_m1, smlh_het_m2, rank = T)
## # Comparison of Model Performance Indices
##
                        Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc
## Name
## ------
## smlh_het_m2 | lmerModLmerTest |
                                    0.202 |
                                               0.029 | 0.178 | 0.076 | 0.085 |
                                                                                   0.277 l
                                            0.028 | 0.154 | 0.077 | 0.085 |
## smlh het m1 | lmerModLmerTest |
                                   0.178 |
                                                                                   0.723 |
```

Plot chemical complexity by mhc heterozygosity

Name Variance Std.Dev.

0.00714119814 0.0845056101

family (Intercept) 0.00154644902 0.0393249160

Number of obs: 56, groups: family, 36

Groups

```
panel2.a <- ggplot(data = meta,</pre>
                   aes(y = compound_n,
                       x = as.factor(mhc_het),
                       fill = as.factor(mhc_het),
                       color = as.factor(mhc_het))) +
  scale fill manual(values = c("darkgrey", "orange")) +
  geom_boxplot(width = 0.4,
               color = "black",
               size = 1) +
  geom_jitter(height = 0.02,
              width = 0.1,
              color = "black",
              size = 3.5,
             alpha = 0.25) +
  scale_x_discrete(name = "MHC heterozygosity",
                     breaks = c(0,1),
                     labels = c("homozygous", "heterozygous")) +
  scale_y_continuous(name = "Chemical diversity") +
  custom_theme
```

Plot chemical complexity by sMLH

```
panel2.b <- ggplot(data = meta,</pre>
                   aes(y = compound n,
                       x = smlh)) +
  geom point(size = 3.5,
             alpha = 0.25) +
  geom_smooth(method = "lm",
              se = T,
              color = "orange") +
  scale_x_continuous(name = "sMLH") +
  scale_y_continuous(name = "Chemical diversity") +
  scale_color_manual(name = "Senescence",
    values = c("#E8B54D", "#000000"),
    labels = c("Mother", "Pup")) +
  scale_fill_manual(name = "Senescence",
    values = c("#E8B54D", "#000000"),
    labels = c("Mother", "Pup")) +
  custom_theme
```

Effect size boostrap of linear mixed effects models

Demanding code and results are prerendered to optimize run-time of this markdown script. Bootstraps take several hours, only run, if you're interested in a different seed as a different randomization should switch up the data points but not overall results.

```
# subsample 30 individuals
model_r2_btrap <- function(x){</pre>
  n <- 1:dim(meta)[1]</pre>
  n_btrap <- sample(n, 30, replace = F)</pre>
  btrap_meta <- meta[n_btrap, ]</pre>
  div_trap <- lmerTest::lmer(compound_n ~ mhc_het + smlh + maturity + colony + (1|family),</pre>
                               data = btrap_meta)
  # bootstrap effect size with partR2
  partR2(div_trap, partvars = c("mhc_het", "smlh", "maturity", "colony"), nboot = 100, CI = 0.95)
}
# amount of times boostrap shall be repeated
nbtrap <- 1000
# repeat function call as often as specified for bootstrap iterations
system.time(
  div_r2_btrap <- lapply(1:nbtrap, model_r2_btrap)</pre>
sim_r2_btrap <- function(x){</pre>
n <- 1:dim(model rel.df)[1]
```

Plot effect size results

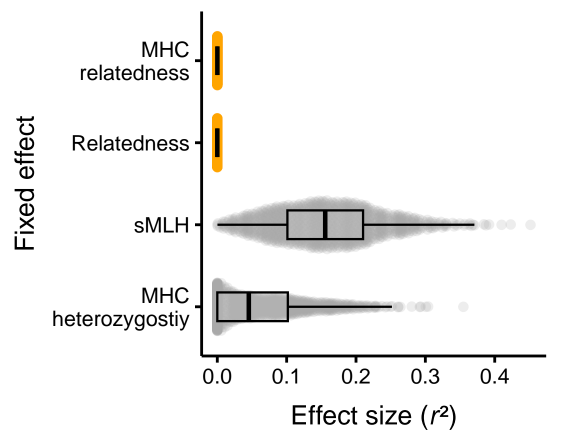
Load in the pre-rendered data.

```
load(file = "RData/div_r2_btrap.RData")
load(file = "RData/sim_r2_btrap_res.RData")
```

Create a ggplot workable data frame

Plot effect sizes

```
alpha = 0.1, lwd=0.8, notch = F) +
labs(y = expression(paste("Effect size (", italic("r"), "²)")),
    x = "Fixed effect") +
scale_x_discrete(labels = c(
    "ufrac" = "MHC\nrelatedness",
    "relatedness" = "Relatedness",
    "r2_smlh" = "sMLH",
    "mhc_het" = "MHC\nheterozygostiy")) +
coord_flip() +
custom_theme
effectsize_best_model_gg
```



```
ggsave(filename = "figures/figure3.png",dpi = 400,
    width = 22, height = 20,
    units = "cm",
    bg = "white")
```

Addition to model selection process by 'dredge' and 'partR2' Similarity model selection and power

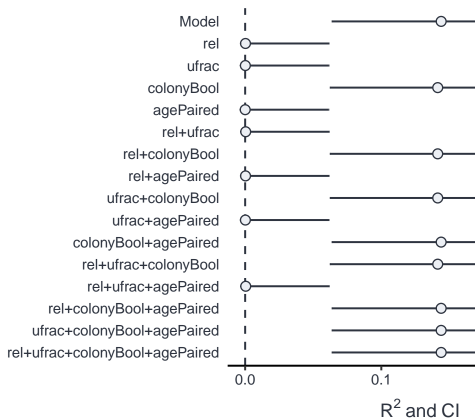
```
# premise for 'dredge' to work correctly
options(na.action = "na.fail")
# model with all effects
```

```
sim_m <- lmerTest::lmer(scent_mds ~ rel + ufrac + colonyBool + agePaired + (1|familyBool) +</pre>
                       (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# calculate different versions
m1 <- dredge(sim_m)</pre>
Find appropiate models
## Warning in dredge(sim_m): comparing models fitted by REML
## Fixed term is "(Intercept)"
subset(m1, delta < 4)</pre>
### Global model call: lmerTest::lmer(formula = scent_mds ~ rel + ufrac + colonyBool +
##
       agePaired + (1 | familyBool) + (1 | pairID1) + (1 | pairID2),
##
       data = model_rel.df)
## ---
## Model selection table
     (Intrc) clnyB df
                        logLik
                                  AICc delta weight
## 3 0.3197
                + 6 1974.155 -3936.3
## Models ranked by AICc(x)
## Random terms (all models):
     1 | familyBool, 1 | pairID1, 1 | pairID2
# chose best
summary(get.models(m1, 1)[[1]])
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ colonyBool + (1 | familyBool) + (1 | pairID1) + (1 |
##
       pairID2)
##
      Data: model_rel.df
##
## REML criterion at convergence: -3948.3
##
## Scaled residuals:
##
                          1Q
                                   Median
## -3.138411025 -0.682850860 -0.075254444 0.581948282 3.762734892
##
## Random effects:
## Groups
               Name
                           Variance
                                         Std.Dev.
               (Intercept) 0.00134929136 0.0367327015
## pairID2
               (Intercept) 0.00132586922 0.0364124872
## familyBool (Intercept) 0.00421404024 0.0649156394
## Residual
                           0.00381737000 0.0617848687
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
##
## Fixed effects:
                     Estimate
                                  Std. Error
                                                          df t value Pr(>|t|)
## (Intercept) 3.19681994e-01 4.75484750e-02 1.09434933e+00 6.72329 0.080084
## colonyBool1 8.39162891e-02 8.04874742e-03 1.22040771e+02 10.42601 < 2e-16 ***
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
               (Intr)
## colonyBool1 -0.154
r2_sim_m <- partR2(sim_m, partvars = c("rel", "ufrac", "colonyBool", "agePaired"),</pre>
                   nboot = 100, CI = 0.95)
```

Calculate effect size by partR2

```
forestplot(r2_sim_m, type = "R2", line_size = 0.7, text_size = 14, point_size = 3)
```



Visualize effect sizes of fixed effects

 $r2_sim_m$

```
##
##
## R2 (marginal) and 95% CI for the full model:
           CI_lower CI_upper nboot ndf
    0.1439 0.0636
                    0.2569
                              100
##
##
##
## Part (semi-partial) R2:
```

```
## Predictor(s)
                                       CI_lower CI_upper nboot ndf
                                0.1439 0.0636
## Model
                                               0.2569
                                                        100
                                                             6
                                0.0002 0.0000
                                               0.0616
## rel
                                                        100
                                                             5
                                0.0001 0.0000
                                               0.0615
## ufrac
                                                        100
                                                             5
## colonyBool
                                0.1414 0.0622
                                               0.2532
                                                        100
                                                             5
## agePaired
                               0.0000 0.0000 0.0614
                                                       100
                                                             4
## rel+ufrac
                               0.0004 0.0000
                                               0.0618 100
## rel+colonyBool
                               0.1414 0.0622
                                               0.2532
                                                       100
## rel+agePaired
                               0.0003 0.0000
                                               0.0617
                                                        100
                                                             3
                                                       100
## ufrac+colonyBool
                              0.1414 0.0622
                                               0.2532
## ufrac+agePaired
                               0.0002 0.0000
                                               0.0616
                                                       100
## colonyBool+agePaired
                               0.1439 0.0636
                                               0.2569
                                                        100
                                                             3
## rel+ufrac+colonyBool
                                0.1414 0.0622
                                               0.2532
                                                       100
                                                             3
                                0.0004 0.0000
## rel+ufrac+agePaired
                                                       100
                                                             2
                                               0.0618
## rel+colonyBool+agePaired
                                0.1439 0.0636
                                               0.2569
                                                        100
                                                             2
## ufrac+colonyBool+agePaired
                                0.1439 0.0636
                                               0.2569
                                                        100
                                                             2
## rel+ufrac+colonyBool+agePaired 0.1439 0.0636
                                               0.2569
                                                        100
                                                             1
```

Diversity model selection

Find appropiate models

```
## Warning in dredge(div_m): comparing models fitted by REML
## Fixed term is "(Intercept)"
subset(m3, delta < 4)</pre>
## Global model call: lmerTest::lmer(formula = compound_n ~ mhc_het + smlh + maturity +
##
      colony + (1 | family), data = meta)
## ---
## Model selection table
       (Int) cln mtr mhc_het
                              sml df
                                       logLik AICc delta weight
## 16 -22.62 + + 1.1860 77.18 7 -222.645 461.6 0.00 0.620
## 12 -21.16
                            76.53 6 -225.280 464.3
                                                     2.65 0.165
## 15 -20.32
                  + 1.2930 76.51 6 -225.621 465.0 3.33 0.117
## 14 -12.22
                    -0.2606 66.23 6 -225.800 465.3 3.69 0.098
## Models ranked by AICc(x)
## Random terms (all models):
##
    1 | family
```

```
select best
```

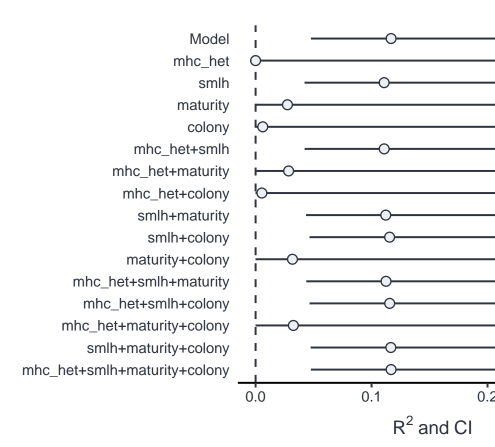
summary(get.models(m3, 1)[[1]])

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: compound_n ~ colony + maturity + mhc_het + smlh + (1 | family)
## Data: meta
##
```

```
## REML criterion at convergence: 445.3
##
## Scaled residuals:
##
            Min
                          1Q
                                    Median
                                                     30
                                                                  Max
## -1.0928313033 -0.4734130794 -0.0457156417 0.2360966398 2.0910996889
##
## Random effects:
## Groups Name
                        Variance
                                  Std.Dev.
## family (Intercept) 289.098507 17.0028970
                       102.550738 10.1267338
## Residual
## Number of obs: 56, groups: family, 36
##
## Fixed effects:
##
                                                df t value Pr(>|t|)
                  Estimate
                           Std. Error
## (Intercept) -22.62052081 25.23757770 41.63258609 -0.89630 0.3752445
                           6.41330434 33.84088527 0.63366 0.5305648
## colonySSB
               4.06383229
## maturityP
               -4.75717833 3.27180878 24.21976233 -1.45399 0.1587859
## mhc het
               1.18588993 5.45889047 42.15362665 0.21724 0.8290693
## smlh
               77.17560246 24.48424644 42.14207673 3.15205 0.0029835 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
            (Intr) clnSSB mtrtyP mhc ht
##
## colonySSB -0.152
## maturityP 0.289 -0.068
## mhc_het -0.247 -0.034 -0.181
## smlh
            -0.973 0.051 -0.313 0.100
r2_div_m <- partR2(div_m,
                  partvars = c("mhc_het", "smlh", "maturity", "colony"),
                  nboot = 100, CI = 0.95)
```

Calculate effect size by partR2

```
forestplot(r2_div_m, type = "R2", line_size = 0.7, text_size = 14, point_size = 3)
```



Visualize effect sizes of fixed effects

```
r2_div_m
##
##
## R2 (marginal) and 95% CI for the full model:
##
           CI_lower CI_upper nboot ndf
    0.1168 0.0477
                     0.3054
                               100
##
##
##
##
  Part (semi-partial) R2:
##
    Predictor(s)
                                   R2
                                          CI_lower CI_upper nboot ndf
##
                                                    0.3054
##
    Model
                                   0.1168 0.0477
                                                              100
                                                                    5
##
    mhc_het
                                   0.0000 0.0000
                                                    0.2155
                                                              100
                                                                    4
##
                                   0.1108 0.0423
                                                    0.3008
                                                              100
                                                                    4
    smlh
                                                    0.2370
                                                                    4
##
    maturity
                                   0.0275 0.0000
                                                              100
##
    colony
                                   0.0062 0.0000
                                                    0.2207
                                                              100
                                                                    4
##
    mhc het+smlh
                                   0.1109 0.0424
                                                    0.3009
                                                              100
                                                                    3
                                                    0.2378
                                                              100
                                                                    3
##
    mhc_het+maturity
                                   0.0285 0.0000
##
    mhc het+colony
                                   0.0054 0.0000
                                                    0.2201
                                                              100
                                                                    3
                                                    0.3019
##
    smlh+maturity
                                   0.1122 0.0436
                                                              100
                                                                    3
    smlh+colony
                                   0.1155 0.0466
                                                    0.3044
                                                              100
                                                                    3
##
    maturity+colony
                                   0.0318 0.0000
                                                    0.2403
                                                              100
                                                                    3
    mhc_het+smlh+maturity
                                   0.1125 0.0438
                                                    0.3021
                                                              100
                                                                    2
##
##
    mhc_het+smlh+colony
                                   0.1156 0.0466
                                                    0.3045
                                                              100
                                                                    2
    mhc_het+maturity+colony
                                   0.0326 0.0000
                                                    0.2409
                                                              100
                                                                    2
```

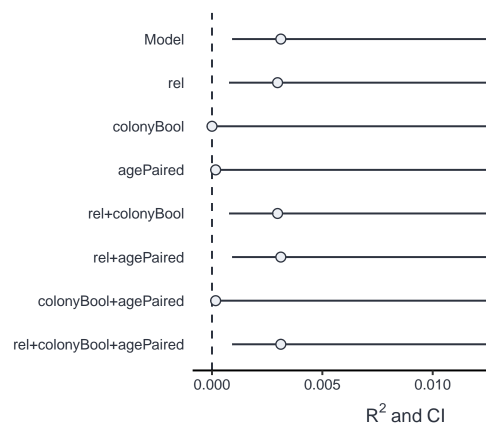
```
## smlh+maturity+colony 0.1166 0.0476 0.3052 100 2 ## mhc_het+smlh+maturity+colony 0.1168 0.0477 0.3054 100 1
```

```
Correlation of genetic effects
gen_m <- lmerTest::lmer(ufrac ~ rel + colonyBool + agePaired + (1|pairID1) + (1|pairID2) + (1|familyBoo
                        data = model rel.df)
m2 <- dredge(gen_m)</pre>
Find appropriate models
## Warning in dredge(gen_m): comparing models fitted by REML
## Fixed term is "(Intercept)"
subset(m2, delta < 4)</pre>
## Global model call: lmerTest::lmer(formula = ufrac ~ rel + colonyBool + agePaired +
       (1 | pairID1) + (1 | pairID2) + (1 | familyBool), data = model_rel.df)
## ---
## Model selection table
   (Intrc)
              rel df logLik AICc delta weight
                     5 337.011 -664.0 0.00 0.675
## 1 0.6825
## 5 0.7107 -0.1267 6 337.288 -662.5 1.46 0.325
## Models ranked by AICc(x)
## Random terms (all models):
    1 | pairID1, 1 | pairID2, 1 | familyBool
summary(get.models(m2, 1)[[1]])
select best
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: ufrac ~ (1 | pairID1) + (1 | pairID2) + (1 | familyBool)
     Data: model_rel.df
##
## REML criterion at convergence: -674
##
## Scaled residuals:
                                   Median
##
                         1Q
                                                    3Q
## -4.485507072 -0.371589025 0.270089938 0.677691079 2.138879974
##
## Random effects:
## Groups
                          Variance
                                        Std.Dev.
## pairID1
             (Intercept) 0.00218749622 0.0467706769
## pairID2 (Intercept) 0.00115356500 0.0339641723
## familyBool (Intercept) 0.02938341728 0.1714159190
## Residual
                           0.03556353542 0.1885829669
## Number of obs: 1540, groups: pairID1, 55; pairID2, 55; familyBool, 2
##
## Fixed effects:
##
                  Estimate Std. Error
                                                df t value Pr(>|t|)
```

(Intercept) 0.682529592 0.123310856 1.004398662 5.53503 0.11302

Calculate effect size by partR2

```
forestplot(r2_gen_m, type = "R2", line_size = 0.7, text_size = 14, point_size = 3)
```



Visualize effect sizes of fixed effects

##

```
r2_gen_m
```

```
## R2 (marginal) and 95% CI for the full model:
          CI_lower CI_upper nboot ndf
   0.0031 9e-04
                   0.0175
                            100
##
##
   _____
##
## Part (semi-partial) R2:
##
  Predictor(s)
                            R2
                                   CI_lower CI_upper nboot ndf
## Model
                            0.0031 9e-04
                                            0.0175
                                                      100
                                                           5
                            0.0030 8e-04
                                            0.0173
                                                     100
## rel
                                                           4
## colonyBool
                            0.0000 0e+00
                                            0.0132
                                                      100
                                                           4
                            0.0002 0e+00
                                                      100
                                                           3
## agePaired
                                            0.0134
## rel+colonyBool
                            0.0030 8e-04
                                            0.0173
                                                      100
                                                           3
## rel+agePaired
                            0.0031 9e-04
                                            0.0175
                                                     100
                                                           2
```

```
## colonyBool+agePaired
                             0.0002 0e+00
                                             0.0134
                                                       100
## rel+colonyBool+agePaired 0.0031 9e-04
                                             0.0175
                                                       100
                                                            1
```

PERMANOVA for individual genotypes and alleles respectively

Create workable dataframe

```
# create data frame containing of:
  # individual substance count for every animal
  # an animals individual genotype, represented by 0 and 1 for a given number
  # of alleles (here ranging from 1 to 19)
idv_allele <- t(phylo_mat) %>%
  # coerce to data.frame
  as.data.frame() %>%
  # combine individual compound number with mhc genotype
  cbind(., compound_n) %>%
  # rename columns
  `colnames<-`(c(paste0("a",1:19), "compound_n"))</pre>
```

Run PERMANOVA on each allele

a13

a14

a15

a16

a17

a18

a19

```
# run permanova to associate individual alleles to compound complexity
allele_permanova <-
  vegan::adonis2(compound n ~ a1 + a2 + a3 + a4 + a5 + a6 + a7 + a8 + a9 + a10 +
                   a11 + a12 + a13 + a14 + a15 + a16 + a17 + a18 + a19
               data = idv_allele)
# View results
allele_permanova
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
##
## vegan::adonis2(formula = compound_n ~ a1 + a2 + a3 + a4 + a5 + a6 + a7 + a8 + a9 + a10 + a11 + a12 +
##
                   SumOfSqs
                                      R2
                                               F Pr(>F)
## a1
             1 0.0151593876 0.0084655024 0.46315
             1 0.0096098366 0.0053664500 0.29360
## a2
## a3
             1 0.0069896633 0.0039032587 0.21355
                                                  0.746
             1 0.0378931425 0.0211607815 1.15771
## a4
                                                  0.279
## a5
             1 0.0026201425 0.0014631741 0.08005 0.882
            1 0.0442421031 0.0247062507 1.35169 0.261
## a6
## a7
            1 0.0221723994 0.0123817997 0.67741 0.418
## a8
            1 0.0564621645 0.0315303363 1.72503 0.184
## a9
            1 0.0022475191 0.0012550888 0.06867 0.891
## a10
            1 0.0239470526 0.0133728246 0.73163 0.423
## a11
            1 0.0201670701 0.0112619576 0.61615 0.483
## a12
            1 0.0751690749 0.0419768926 2.29657
```

0.046

0.887

1 0.0171102090 0.0095549055 0.52275 0.501

1 0.0129836481 0.0072504976 0.39668 0.576

1 0.0591946266 0.0330562334 1.80852 0.206 1 0.0120118860 0.0067078336 0.36699 0.579

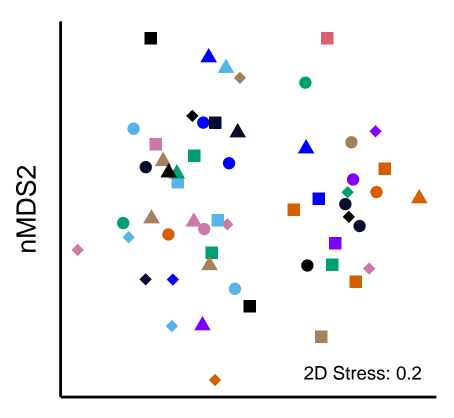
1 0.0578941065 0.0323299800 1.76878 1 0.1338806176 0.0747633560 4.09033

1 0.0026532032 0.0014816363 0.08106

Residual 36 1.1783172402 0.6580112405

```
## Total
                          55 1.7907250935 1.0000000000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# give out p-values for each individual allele
pvals <- allele_permanova[1:19,5]</pre>
# correct p-values by fdr
pvals_corrected <- p.adjust(pvals, method = "fdr") %>% as.data.frame()
pvals corrected
##
## 1 0.785785714286
## 2 0.808133333333
## 3 0.885875000000
## 4 0.757285714286
## 5 0.89100000000
## 6 0.757285714286
## 7 0.785785714286
## 8 0.757285714286
## 9 0.89100000000
## 10 0.785785714286
## 11 0.785785714286
## 12 0.757285714286
## 13 0.785785714286
## 14 0.757285714286
## 15 0.757285714286
## 16 0.89100000000
## 17 0.785785714286
## 18 0.757285714286
## 19 0.785785714286
PERMANOVA for associated odour nmds profiles with genotypes
# combine individuals alleles for each individual to genotype in same dataframe
het_table %<>% mutate(gtype = as.factor(paste0(a1, "/", a2)))
vegan::adonis2(scent ~ het_table$gtype)
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
##
## vegan::adonis2(formula = scent ~ het_table$gtype)
                                           Df
                                                           SumOfSqs
                                                                                                     R2
                                                                                                                          F Pr(>F)
## het_table$gtype 36 8.803613197 0.6710248615 1.07653 0.173
## Residual
                                          19 4.316039594 0.3289751385
                                          55 13.119652790 1.0000000000
## Total
scent_nmds %<>% cbind(., gtype = as.factor(het_table$gtype))
Plot PERMANOVA results
# create color palette for the plot
clr \leftarrow c("\#D55E00", "\#0000ff", "\#56B4E9", "\#009E73", "\#000000", "\#CC79A7", "\#a4805c", "#56B4E9", "
                    "turquoise", "#ed0c2e", "#8000ff", "#ffb700", "#ffff00", "#0a0c2e", "#db5e71")
```

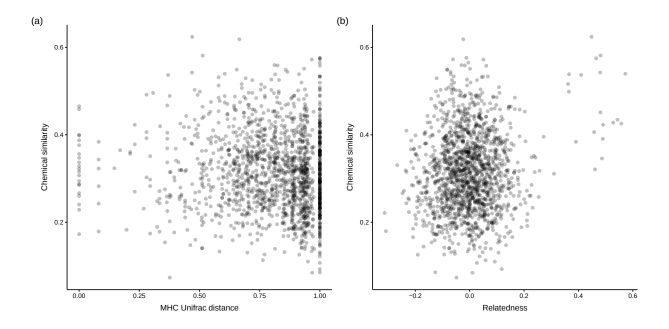
```
# assign pch values for plotting
shp \leftarrow c(17, 15, 16, 18)
color_shape_pairs <- crossing(clr,shp)</pre>
shape_pair_df <- data.frame(fam = levels(scent_nmds$gtype),</pre>
                             color_shape_pairs[1:length(levels(scent_nmds$gtype)),])
cross_ref <- match(scent_nmds$gtype, shape_pair_df$fam)</pre>
shape_pair_df %<>% .[cross_ref,]
scent_nmds %<>% cbind(.,
                    shape_pair_df[,2:3])
scent_nmds %<>% mutate(across(clr:shp, as.factor))
ggplot(data = scent_nmds,aes(MDS1,MDS2, color = clr, shape = shp)) +
  geom_point(size = 4) +
  scale_shape_manual(values = as.numeric(levels(scent_nmds$shp))) +
  theme void() +
  scale_color_manual(values = levels(as.factor(scent_nmds$clr))) +
  annotate("text", x = 0.48, y = -0.75, label = "2D Stress: 0.2", size = 5) +
  scale_x_continuous(name = "nMDS1") +
  scale_y_continuous(name = "nMDS2") +
  custom_theme +
  theme(
   legend.position = "none",
   axis.ticks = element_blank(),
    axis.text = element_blank()
  )
```

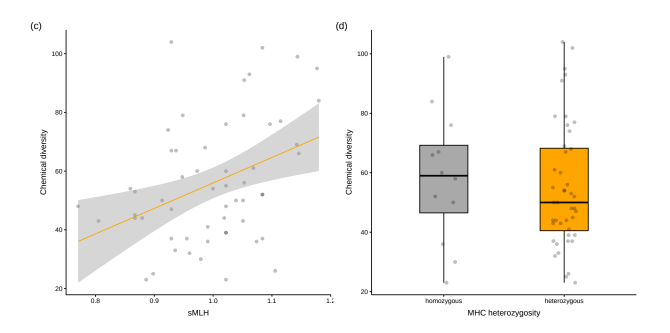


nMDS1

```
# save output
ggsave(filename = "figures/supplementary_figure1.png",
    width = 32, height = 16,
    units = "cm", dpi = 400)
```

Create manuscript panel figure





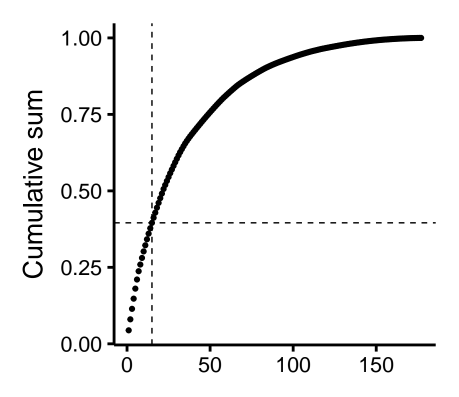
Repeat for best subset by SIMPER analysis

```
# simper analysis
min_range <- 12
max_range <- 42</pre>
scent_range <- scent[,as.numeric(colnames(scent)) >= min_range & as.numeric(colnames(scent)) <= max_ran</pre>
simp_colony <- vegan::simper(scent_range, meta$colony)</pre>
# getting best substances and their contribution to colony dissimilarity
#set numbers of best compounds to retain
keep_compounds <- 15
simp_colony_names <- summary(simp_colony, ordered = T)[[1]] %>%
        rownames(.) %>%
        .[1:keep_compounds]
contribution <- summary(simp_colony, ordered = TRUE)[[1]] %>%
        .$average %>%
        .[1:keep_compounds]
# indices of colony substances (58,62,68,74,86,89,90,98,106,107,110,164,181,189,211)
ind_col <- which(names(scent) %in% simp_colony_names)</pre>
# connect to data frame and compute contribution in percent
col_simp <- data.frame(comp = simp_colony_names, contrib = contribution*100, stringsAsFactors = FALSE)
col_simp
##
                  comp
                             contrib
## 1 15.4498717948718 3.05575894307
## 2 19.7148780487805 2.48839019949
## 3 19.5174358974359 2.37671240496
## 4 16.3895238095238 2.30226998568
## 5 26.7808450704225 2.27321897210
## 6 21.3982051282051 2.06493210540
## 7
                 21.34 1.87869623710
## 8 30.7885294117647 1.51069283280
## 9 37.5734782608696 1.49482594047
## 10 38.4982142857143 1.47661837144
## 11 17.4137142857143 1.38914137040
## 12 20.509347826087 1.35687404025
## 13 15.5008108108108 1.27741681524
               38.5256 1.23455984162
## 15 19.6568571428571 1.21537246870
```

```
print(
   paste0(
     "Subset of ", keep_compounds, " chemicals accounts for ", round(sum(col_simp$contrib), digits = 2),
     "% of colony similarity"))
```

[1] "Subset of 15 chemicals accounts for 27.4% of colony similarity"

cumulative contribution plot



Substances in SIMPER

```
ggsave("figures/supplementary_figure2.png", dpi = 400,
       plot = plot_simper_cusum)
## Saving 6.5 \times 4.5 in image
adjust scent matrix, include only 15 most explanatory compounds
scent_simper <- scent[, ind_col]</pre>
## keep order of rows consistent
scent <- scent[match(rownames(peak_factors),rownames(scent)),]</pre>
## NMDS using Bray-Curtis dissimilarities
scent_nmds_simper.obj <- vegan::metaMDS(comm = scent, distance = "bray")</pre>
## Run 0 stress 0.237312208882
## Run 1 stress 0.260622188214
## Run 2 stress 0.252609955688
## Run 3 stress 0.247186199252
## Run 4 stress 0.260781284585
## Run 5 stress 0.237246533424
## ... New best solution
## ... Procrustes: rmse 0.00328746829711 max resid 0.017717070994
## Run 6 stress 0.247186199283
## Run 7 stress 0.237319083164
## ... Procrustes: rmse 0.00471213195535 max resid 0.019276015911
## Run 8 stress 0.265021569917
```

```
## Run 9 stress 0.237259771147
## ... Procrustes: rmse 0.00329416346959 max resid 0.0184551380275
## Run 10 stress 0.261552436591
## Run 11 stress 0.237312207932
## ... Procrustes: rmse 0.00328679804084 max resid 0.0177205495253
## Run 12 stress 0.267009219007
## Run 13 stress 0.257209600275
## Run 14 stress 0.25905051738
## Run 15 stress 0.24974859163
## Run 16 stress 0.237246535186
## ... Procrustes: rmse 1.73970827882e-05 max resid 0.000115321703975
## ... Similar to previous best
## Run 17 stress 0.248999811536
## Run 18 stress 0.237319083127
## ... Procrustes: rmse 0.00471212433109 max resid 0.0192740471189
## Run 19 stress 0.252609960826
## Run 20 stress 0.23731908334
## ... Procrustes: rmse 0.00471217644571 max resid 0.0192790842828
## *** Best solution repeated 1 times
## get x and y coordinates
scent nmds simper <- as.data.frame(scent nmds.obj[["points"]]) %>%
## add the colony as a factor to each sample
cbind(.,colony = peak_factors[["colony"]])
```

Create vectorized distance measurements for scent data

```
# bray-curtis distance measurement on scent profiles
scent_dist_simper <- vegdist(scent_simper) %>% as.matrix()
scent_dist_simper[upper.tri(scent_dist_simper, diag = T)] <- NA
b_simper <- scent_dist_simper %>% as.vector() %>% na.omit()
```

Repeat chemical similarity models with best chemicals subset

```
model rel.df %<>% cbind(., scent simper mds = b simper)
# mhc
z1 <- lmerTest::lmer(scent_simper_mds ~ ufrac + colonyBool + agePaired + (1 familyBool) +
                       (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# relatedness
z2 <- lmerTest::lmer(scent_simper_mds ~ rel + colonyBool + agePaired + (1|familyBool) + (1|pairID1) +
                       (1|pairID2),
                     data = model_rel.df)
# mhc & relatedness
z3 <- lmerTest::lmer(scent_simper_mds ~ rel + ufrac + colonyBool + agePaired + (1|familyBool) +
                       (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# no genetic effect
z4 <- lmerTest::lmer(scent_simper_mds ~ colonyBool + agePaired + (1|familyBool) + (1|pairID1) +
                       (1|pairID2),
                     data = model_rel.df)
# compare model performance scores
```

```
compare_performance(z1, z2, z3, z4, rank = T) %>%
 arrange(Name)
## # Comparison of Model Performance Indices
##
                Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc weight
## ------
                             0.670 |
                                          0.350 | 0.493 | 0.096 | 0.099 |
       | lmerModLmerTest |
      | lmerModLmerTest |
                            0.658 |
                                         0.365 | 0.461 | 0.096 | 0.099 |
## 72
      | lmerModLmerTest | 0.658 | | lmerModLmerTest | 0.670 |
                                         0.365 | 0.462 | 0.096 | 0.099 |
## z3
## z4
                                         0.351 | 0.492 | 0.096 | 0.099 |
summary(z2)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_simper_mds ~ rel + colonyBool + agePaired + (1 | familyBool) +
##
      (1 | pairID1) + (1 | pairID2)
##
     Data: model_rel.df
##
## REML criterion at convergence: -2489.9
## Scaled residuals:
##
          Min
                       1Q
                                Median
## -3.361727005 -0.601762806 0.044879466 0.656512064 4.481398163
## Random effects:
## Groups
                        Variance
                                     Std.Dev.
## pairID2
            (Intercept) 0.00371962719 0.0609887464
## pairID1 (Intercept) 0.00222537088 0.0471738368
## familyBool (Intercept) 0.00248148577 0.0498145136
## Residual
                        0.00985985813 0.0992968183
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
## Fixed effects:
##
                    Estimate
                              Std. Error
                                                     df
                                                         t value Pr(>|t|)
## (Intercept)
                ## rel
               -0.0339705052 0.0329143620 473.1835094380 -1.03209 0.302558
## colonyBool1
               -0.2041643091 0.0118146357 112.2879453844 -17.28063 < 2e-16
## agePairedM/P -0.0165814738 0.0119597238 128.9256179904 -1.38644 0.168005
## agePairedP/P -0.0317056901 0.0224867649 99.4223955740 -1.40997 0.161669
##
## (Intercept) *
## rel
## colonyBool1 ***
## agePairedM/P
## agePairedP/P
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
                          clnyB1 agPM/P
             (Intr) rel
             -0.149
## rel
## colonyBool1 -0.256 0.016
```

0.157

0.304 |

0.113 |

0.426 l

0.15

0.30

0.11

0.43

```
## agePairdM/P -0.250 -0.024 -0.002
## agePairdP/P -0.237 -0.009 -0.003 0.900
summary(z4)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_simper_mds ~ colonyBool + agePaired + (1 | familyBool) +
       (1 | pairID1) + (1 | pairID2)
##
##
      Data: model_rel.df
##
## REML criterion at convergence: -2493.8
## Scaled residuals:
##
           Min
                          1Q
                                   Median
                                                   3Q
                                                                Max
## -3.369275911 -0.607272157 0.045559137 0.657635847 4.508880002
## Random effects:
## Groups
                          Variance
                                        Std.Dev.
               (Intercept) 0.00373524835 0.0611166781
## pairID2
               (Intercept) 0.00223267976 0.0472512409
## pairID1
## familyBool (Intercept) 0.00356932632 0.0597438392
## Residual
                          0.00985536801 0.0992742062
## Number of obs: 1540, groups: pairID2, 55; pairID1, 55; familyBool, 2
## Fixed effects:
##
                     Estimate
                                  Std. Error
                                                          df
                                                              t value Pr(>|t|)
## (Intercept)
                 0.5579296640 0.0474009636
                                               1.3493551125 11.77043 0.025259
## colonyBool1
                -0.2038667903
                                0.0118300291 112.1681349190 -17.23299 < 2e-16
                                0.0119726612 128.7456942475 -1.40215 0.163278
## agePairedM/P -0.0167874180
## agePairedP/P -0.0318598080
                                0.0225198624 99.4410644101 -1.41474 0.160268
##
## (Intercept) *
## colonyBool1 ***
## agePairedM/P
## agePairedP/P
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
               (Intr) clnyB1 agPM/P
## colonyBool1 -0.224
## agePairdM/P -0.224 -0.001
## agePairdP/P -0.210 -0.003 0.900
# if interested
# check model performance by
# check_model(z2)
#
```

Repeat analyses for moms only

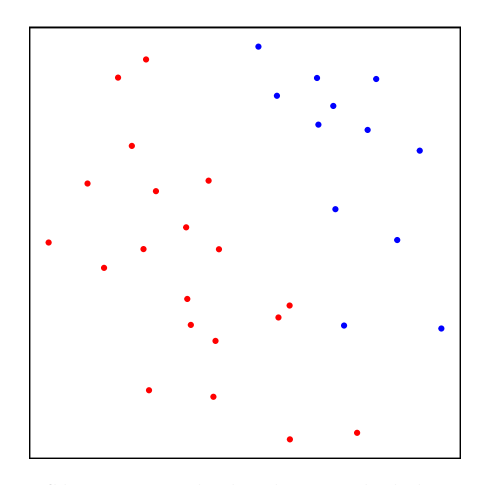
empty workspace to repeat analyses for moms

```
rm(list=ls())
```

Subset scent data to correlate same individuals

```
## read in meta data
meta <- read.table(file = "data/arga_metadata.txt", sep = "\t") %>%
  `colnames<-`(unlist(.[1,])) %>%
  .[-1,] %>%
  # subset for only moms
 filter(., maturity == "M")
## normalise area and return a data frame
scent <- norm_peaks(aligned_peak_data,</pre>
                    conc_col_name = "area",
                    rt col name = "time",
                    out = "data.frame")
## common transformation for abundance data to reduce the extent of mean-variance trends
scent <- log(scent + 1)</pre>
n_scnt <- rownames(scent)</pre>
keep_i <- match(meta$id, n_scnt)</pre>
scent %<>%
  .[keep_i, ] %>%
  `rownames<-`(meta$real_id)
## NMDS with reduced data
## GCalignR contains factors for the chemical dataset
data("peak_factors")
peak_factors <- peak_factors[match(meta$id, rownames(peak_factors)),] %>%
  `rownames<-`(meta$real_id)
## keep order of rows consistent
scent <- scent[match(rownames(peak_factors),rownames(scent)),]</pre>
## NMDS using Bray-Curtis dissimilarities
scent_nmds.obj <- vegan::metaMDS(comm = scent, distance = "bray")</pre>
## Run 0 stress 0.236250449583
## Run 1 stress 0.235605102249
## ... New best solution
## ... Procrustes: rmse 0.06131503612 max resid 0.278985505315
## Run 2 stress 0.254789734088
## Run 3 stress 0.236896130641
## Run 4 stress 0.260508086835
## Run 5 stress 0.233612595113
## ... New best solution
```

```
## ... Procrustes: rmse 0.0691442965345 max resid 0.269369957952
## Run 6 stress 0.233612593985
## ... New best solution
## ... Procrustes: rmse 2.18937130058e-05 max resid 7.89459539875e-05
## ... Similar to previous best
## Run 7 stress 0.233785520575
## ... Procrustes: rmse 0.00876312592908 max resid 0.0371260735997
## Run 8 stress 0.258333360017
## Run 9 stress 0.241730929293
## Run 10 stress 0.237657583017
## Run 11 stress 0.234290238728
## Run 12 stress 0.264573253674
## Run 13 stress 0.234468822273
## Run 14 stress 0.243895058834
## Run 15 stress 0.234374205772
## Run 16 stress 0.260508153374
## Run 17 stress 0.233785517964
## ... Procrustes: rmse 0.00876234172883 max resid 0.0371229389694
## Run 18 stress 0.235283613596
## Run 19 stress 0.254789858386
## Run 20 stress 0.250551297406
## *** Best solution repeated 1 times
## get x and y coordinates
scent_nmds <- as.data.frame(scent_nmds.obj[["points"]])</pre>
## add the colony as a factor to each sample
scent_nmds <- cbind(scent_nmds,colony = peak_factors[["colony"]])</pre>
## quick plotting
ggplot(data = scent_nmds,aes(MDS1,MDS2,color = colony)) +
  geom_point() +
 theme_void() +
  scale_color_manual(values = c("blue", "red")) +
 theme(panel.background = element_rect(colour = "black",
                                        size = 1.
                                        fill = NA),
        aspect.ratio
                        = 1,
        legend.position = "none")
## Warning: The `size` argument of `element_rect()` is deprecated as of ggplot2 3.4.0.
## i Please use the `linewidth` argument instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
```



Calculate MHC heterozygosity relatedness between individuals

```
## read in mhc genotype data
mhc_het_dat <- read.table("data/clone_mhc_het.txt")</pre>
## restructure `mhc_het_dat`to fit `Demerelate()::inputdata)
## id and colony as factors; alleles as integers or numeric
## otherwise `rxy`cannot handle computations
mhc_het_dat %<>%
  rownames_to_column(., var = "id") %>%
  # mutate(., a1 = str_pad(a1, 2, pad = "0")) %>%
  \# mutate(., a2 = str_pad(a2, 2, pad = "0")) %>%
  mutate(., colony = as.factor(rep("col", 56))) %>%
  mutate(., id = as.factor(id)) %>%
  .[,-4] %>%
  relocate(., colony, .before = a1)
  ## order mhc_het_dat$id after meta$real_id
  ## so data is consistently ordered same in all data.frames
## get matching indeces
id_index <- match(meta$real_id, mhc_het_dat$id)</pre>
## sort correspondingly
mhc_het_dat %<>% .[id_index,]
## calculate relatedness after Queller & Goodnight
mhc_relatedness_res <- Demerelate(inputdata = mhc_het_dat,</pre>
```

```
value = "rxy",
                                  object = T,
                                  NA.rm = F,
                                  Fis = F)
## Warning in Demerelate(inputdata = mhc het dat, value = "rxy", object = T, : Careful, bi-allelic mark
     Especially, rxy and ritland estimator are not defined when bi-allelic estimates are used with alle
     You should consider removing bi-allelics which tend to have very evenly distributed alleles or swi
     Be careful even if allele frequencies are not perfectly 0.5, during randomizations problems may oc
##
mhc_relatedness <- unlist(mhc_relatedness_res$Empirical_Relatedness)</pre>
## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals
relate_mat_mhc <- matrix(nrow = dim(mhc_het_dat)[1],</pre>
                         ncol = dim(mhc_het_dat)[1])
## fill distance matrix row wise, thus fill upper.tri
relate_mat_mhc[upper.tri(relate_mat_mhc)] <- mhc_relatedness</pre>
## transpose to keep consistency with other distance matrices
relate_mat_mhc <- t(relate_mat_mhc)</pre>
relate_mat_mhc %<>% `colnames<-`(meta$real_id) %>% `rownames<-`(meta$real_id)
## vectorize again to identify whether relatedness pairs were consistent in the first place
a <- relate_mat_mhc %>% as.vector() %>% na.omit()
Create vectorized distance measurements for scent data
# bray-curtis distance measurement on scent profiles
scent_dist <- vegdist(scent) %>% as.matrix()
```

Generate UniFrac distances from MHC DQB II individual genotypes

scent_dist[upper.tri(scent_dist, diag = T)] <- NA
b <- scent_dist %>% as.vector() %>% na.omit()

```
# handle genotypes as otu table
phylo_mat <- read.table("data/phyloseq-mat.txt") %>%
    as.matrix()

# make sample names consistent
n <- match(meta$real_id, colnames(phylo_mat))

phylo_mat %<>% .[, n] %>%
    otu_table(., taxa_are_rows = T)

# create phylogenetic tree from file
phylo_tree <- ape::read.tree("data/unifrac_tree_p.nwk")

# merge into Formal class phyloseq
arga_phylseq <- merge_phyloseq(phylo_mat, phylo_tree)

# create UniFrac as genetic diversity measurement for single locus data
mhc_dqb2_ufrac <- UniFrac(arga_phylseq, weighted = F) %>%
    # distances to distance matrix
as.matrix()
```

```
# vectorize distances matrices
mhc_dqb2_ufrac[upper.tri(mhc_dqb2_ufrac, diag = T)] <- NA
c <- mhc_dqb2_ufrac %>% as.vector() %>% na.omit()
```

Calculate microsatellite relatedness values

create data.frame in correspondence to Demerelate input format

```
# read in genotype data table
msats_df <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t")</pre>
# update data.frame with additional info
# "delete" colony info, otherwise relatedness is only calculated for individuals
# within their own colonies -> no complete pairwise comparison
msats_df <- cbind(id = as.factor(rownames(msats_df)),</pre>
                   # colony = meta$colony,
                  colony = rep("col", 56),
                  msats_df[1:56,]) %>%
  # clear df from rownames/ only keep colnames/ variable names
  `rownames<-`(NULL)
msats_df[is.na(msats_df)] = 0
str(msats_df)
# write.table(msats df, file = "data/msats genotypes demerelate.txt",
              sep = " \setminus t",
#
              row.names = F)
msats df %<>% . [match(meta$real id, .$id),]
```

Calculate relatedness of individuals based on Queller & Goodnight

sum(table(relate.non.X.mean))), : Chi-Quadrat-Approximation kann inkorrekt sein

```
## Warning in Demerelate(inputdata = msats_df, value = "rxy", object = T, NA.rm = F, : Careful, bi-alle
## Especially, rxy and ritland estimator are not defined when bi-allelic estimates are used with alle
## You should consider removing bi-allelics which tend to have very evenly distributed alleles or swi
## Be careful even if allele frequencies are not perfectly 0.5, during randomizations problems may oc
## Warning in prop.test(c(emp, non), c(sum(table(empirical.list)),
```

Coerce output to a vector

```
relatedness <- unlist(relatedness_results$Empirical_Relatedness)
## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals</pre>
```

Analyse Odour and genetic association by MHC DQB II and neutral genomic background

Create data.frame to plot in ggplot2

```
## substitute once tested correctly
## scent_mds shall contain similarity values but `b` contains
## dissimilarity values based on Bray-Curtis -> substracting
## dissmilarities from 1 returns similarities

model_rel.df <- cbind(mhc_rel = a, scent_mds = 1-b, ufrac = c, rel = d) %>% as.data.frame()
```

Custom theme to make plot aesthetics consistent

Model odour relationship on MHC and neutral genetic background

Pool underlying data dependencies

Create a function that generates pairwise variables in a systematic matter for pairwise comparisons

```
create_pair_vars <-function(row_cross, col_cross, split_vars = F){
   require(stringr)

   rc <- row_cross
   cc <- col_cross

# create empy matrix
# keep row and col names from existing distance matrices

empty_mat <- matrix(nrow = length(rc),</pre>
```

```
ncol = length(cc)) %>%
    `colnames<-`(cc) %>%
    `rownames<-`(rc)
  # fill each matrix i,j-th cell with the crossing from their corresponding
  \# i-th rowname and j-th colname
  for (i in 1:dim(empty_mat)[1]) {
    for (j in 1:dim(empty_mat)[2]) {
      empty_mat[i,j] <- paste0(rc[i], "/", cc[j])</pre>
    } # end j
  } # end i
  # delete `upper.tri()` of `empty_mat` to resemble structure of the other
  # distance matrices in use
  empty_mat[upper.tri(empty_mat, diag = T)] <- NA</pre>
  pair_vars <- empty_mat %>% as.vector() %>% na.omit()
  # split `pair_vars` if needed
  if (split_vars == T) {
    pair_vars1 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][1]
                          })
    pair_vars2 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][2]
    pair_vars_split <- list(pair_variable1 = pair_vars1,</pre>
                             pair_variable2 = pair_vars2)
    return(pair_vars_split)
  } else {
    return(pair_vars)
} #end create_pair_vars
```

Helper function to combine double entries

```
## for x, overwrite specified replacer with specified value
f <- function(x, replacer, overwrite){
  if (x == replacer) {
    x <- overwrite
  } else {
    x <- x
}</pre>
```

}

Transform model variables

```
agePaired <- create_pair_vars(row_cross = meta$maturity,</pre>
                                col_cross = meta$maturity) %>%
  sapply(., f, "P/M", "M/P")
colonyPaired <- create_pair_vars(row_cross = meta$colony,</pre>
                                   col_cross = meta$colony) %>%
  sapply(., f, "FWB/SSB", "SSB/FWB")
colonyID1 <- create pair vars(row cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[1] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
colonyID2 <- create_pair_vars(row_cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[2] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
colonyBool <- ifelse(colonyID1 == colonyID2, 1, 0)</pre>
familyPaired <- create_pair_vars(row_cross = meta$family,</pre>
                                   col cross = meta$family)
familyID1 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[1] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
familyID2 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[2] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
pairID1 <- create_pair_vars(row_cross = meta$real_id,</pre>
                             col_cross = meta$real_id,
                              split_vars = T)[1] %>%
  unlist() %>%
  as.vector()
pairID2 <- create_pair_vars(row_cross = meta$real_id,</pre>
                              col_cross = meta$real_id,
                              split_vars = T)[2] %>%
```

```
unlist() %>%
as.vector()

familyBool <- ifelse(familyID1 == familyID2, 1, 0)</pre>
```

Update data.frame with model variables

update data frame with meta data

Include information about MHC heterozygosity, sMLH from microsatellite data and chemical diversity by number of compounds per individual

```
scent.abs <- ifelse(scent != 0, 1, 0)</pre>
compound_n <- apply(scent.abs, 1, sum)</pre>
names(compound n) == meta$real id
## [31] TRUE TRUE
# read in heterzygosity information
het_table <- read.table("data/arga_mhc_het.txt", sep = "\t")</pre>
# keep names consistent
match_het <- match(meta$real_id, rownames(het_table))</pre>
het_table %<>% .[match_het,]
# generate sMLH with microsatellite data
# table is pre-prepped, thus rows correspond to same individuals in meta data
smlh res <- read.table("data/msats genotypes inbreedR.txt", sep = "\t")</pre>
smlh_res <- smlh_res[match(meta$real_id, rownames(smlh_res)), ] %>%
 # convert to inbreedR format
 convert_raw() %>%
 # generate sMLH
 sMLH()
meta %<>% cbind(., compound_n = compound_n,
             mhc_het = het_table$het,
             smlh = smlh_res)
meta %<>% mutate(
real_id = as.factor(real_id),
```

```
colony = as.factor(colony),
maturity = as.factor(maturity),
family = as.factor(family)
)
```

Repeat models for chemical similarity

```
# without age and indicator for family groupings
# as grouping factors must have >1 sampled level
c1 <- lmerTest::lmer(scent_mds ~ ufrac + colonyBool + (1|pairID1) + (1|pairID2),</pre>
                     data = model_rel.df)
# relatedness
c2 <- lmerTest::lmer(scent_mds ~ rel + colonyBool + (1|pairID1) + (1|pairID2),</pre>
                     data = model_rel.df)
# mhc & relatedness
c3 <- lmerTest::lmer(scent_mds ~ rel + ufrac + colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# no genetic effect
c4 <- lmerTest::lmer(scent_mds ~ colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# compare model performance scores
compare_performance(c1, c2, c3, c4, rank = T) %>%
 arrange(Name)
```

```
## # Comparison of Model Performance Indices
##
## Name |
                    Model | R2 (cond.) | R2 (marg.) |
                                                         ICC | RMSE | Sigma | AIC weights | AICc weight
## ---
        | lmerModLmerTest |
                                 0.524 |
                                               0.245 | 0.370 | 0.060 | 0.064 |
                                                                                      0.200 |
                                                                                                      0.19
## c1
        | lmerModLmerTest |
                                               0.245 | 0.370 | 0.060 | 0.064 |
## c2
                                 0.524 |
                                                                                      0.196 |
                                                                                                      0.19
        | lmerModLmerTest |
                                               0.244 | 0.369 | 0.060 | 0.064 |
                                                                                      0.074 |
                                                                                                      0.07
## c3
                                 0.524 |
        | lmerModLmerTest |
                                 0.525 |
                                               0.245 | 0.370 | 0.060 | 0.064 |
                                                                                      0.531 |
                                                                                                      0.53
## c4
summary(c2)
```

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ rel + colonyBool + (1 | pairID1) + (1 | pairID2)
     Data: model rel.df
##
## REML criterion at convergence: -1211.3
##
## Scaled residuals:
##
           Min
                                   Median
                          1Q
                                                    30
                                                                Max
## -2.434739031 -0.668546769 -0.125072431 0.516608517 3.362588564
##
## Random effects:
                                       Std.Dev.
## Groups
           Name
                        Variance
## pairID1 (Intercept) 0.00129394285 0.0359714171
## pairID2 (Intercept) 0.00108030441 0.0328679845
```

```
0.00404600916 0.0636082475
## Number of obs: 496, groups: pairID1, 31; pairID2, 31
## Fixed effects:
                    Estimate
                                 Std. Error
                                                        df t value
## (Intercept) 2.64239187e-01 1.31288102e-02 4.08268587e+01 20.12667 < 2.22e-16
              1.70656495e-03 3.59170911e-02 4.83363527e+02 0.04751
## colonyBool1 9.11773096e-02 1.13258476e-02 7.04830509e+01 8.05037 1.4125e-11
##
## (Intercept) ***
## rel
## colonyBool1 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) rel
## rel
               0.030
## colonyBool1 -0.666 0.045
summary(c4)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ colonyBool + (1 | pairID1) + (1 | pairID2)
     Data: model_rel.df
## REML criterion at convergence: -1216.1
##
## Scaled residuals:
##
           Min
                                  Median
                         1Q
                                                   3Q
                                                               Max
## -2.438554364 -0.669736096 -0.127401072 0.518947513 3.362266186
##
## Random effects:
## Groups
           Name
                        Variance
                                      Std.Dev.
## pairID1 (Intercept) 0.00129455037 0.0359798606
## pairID2 (Intercept) 0.00108126826 0.0328826437
                        0.00403675432 0.0635354572
## Residual
## Number of obs: 496, groups: pairID1, 31; pairID2, 31
## Fixed effects:
                               Std. Error
                                                     df t value
                                                                   Pr(>|t|)
                   Estimate
## (Intercept) 0.2642215710 0.0131250678 41.6332801738 20.13106 < 2.22e-16 ***
## colonyBool1 0.0911511887 0.0113126069 71.5115810251 8.05749 1.2502e-11 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
              (Intr)
## colonyBool1 -0.668
```

Repeat models for chemical diversity

```
# exclude family as random effects (no accounting of same family in different
# individuals)
# exlude age as only one 'age' group being moms is modelled
d1 <- lm(compound_n ~ mhc_het + colony , data = meta)</pre>
d2 <- lm(compound n ~ smlh + colony, data = meta)
d3 <- lm(compound_n ~ mhc_het + smlh + colony, data = meta)
d4 <- lm(compound_n ~ colony, data = meta)
compare_performance(d1, d2, d3, d4, rank = T) %>% arrange(Name)
## # Comparison of Model Performance Indices
                     R2 | R2 (adj.) |
                                        RMSE | Sigma | AIC weights | AICc weights | BIC weights | Perf
## Name | Model |
## d1
             lm | 0.091 |
                              0.029 | 21.014 | 22.075 |
                                                              0.086 |
                                                                              0.086 |
                                                                                            0.079 |
## d2
             lm | 0.190 |
                              0.134 | 19.841 | 20.842 |
                                                              0.540 l
                                                                             0.543 l
                                                                                            0.500 I
                              0.104 | 19.831 | 21.200 |
## d3
             lm | 0.191 |
                                                              0.202 |
                                                                             0.134 |
                                                                                            0.090 |
## d4
             lm | 0.074 |
                              0.043 | 21.217 | 21.913 |
                                                              0.172 |
                                                                              0.236 |
                                                                                            0.331 |
summary(d2)
## Call:
## lm(formula = compound_n ~ smlh + colony, data = meta)
## Residuals:
##
                                   Median
                                                    3Q
                          1Q
                                                                Max
## -36.56900864 -15.29316134
                             -0.55473784 12.51322130 45.43810011
##
## Coefficients:
                   Estimate
                              Std. Error t value Pr(>|t|)
## (Intercept) -27.91731762 39.51389375 -0.70652 0.485504
                80.82602303
                             39.60992119
                                         2.04055 0.050493 .
## colonySSB
                10.39704337
                              7.67024625 1.35550 0.185721
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 20.8416688 on 29 degrees of freedom
## Multiple R-squared: 0.189883606,
                                        Adjusted R-squared: 0.13401351
## F-statistic: 3.3986626 on 2 and 29 DF, p-value: 0.0471995222
```

Repeat analyses for pups only

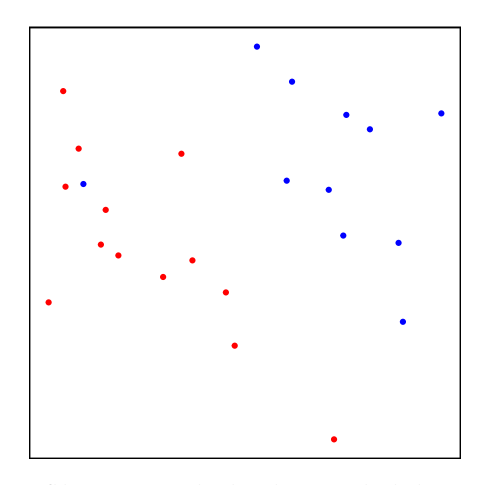
Empty Workspace to analyse pups isolated

```
rm(list=ls())
```

Subset scent data to correlate same individuals

```
## read in meta data
meta <- read.table(file = "data/arga_metadata.txt", sep = "\t") %>%
  `colnames<-`(unlist(.[1,])) %>%
  .[-1,] %>%
  # subset for only pups
 filter(., maturity == "P")
## normalise area and return a data frame
scent <- norm_peaks(aligned_peak_data,</pre>
                    conc_col_name = "area",
                    rt_col_name = "time",
                    out = "data.frame")
## common transformation for abundance data to reduce the extent of mean-variance trends
scent <- log(scent + 1)</pre>
n scnt <- rownames(scent)</pre>
keep_i <- match(meta$id, n_scnt)</pre>
scent %<>%
  .[keep_i, ] %>%
  `rownames<-`(meta$real_id)
## NMDS with reduced data
## GCalignR contains factors for the chemical dataset
data("peak_factors")
peak_factors <- peak_factors[match(meta$id, rownames(peak_factors)),] %>%
  `rownames<-`(meta$real_id)
## keep order of rows consistent
scent <- scent[match(rownames(peak_factors),rownames(scent)),]</pre>
## NMDS using Bray-Curtis dissimilarities
scent_nmds.obj <- vegan::metaMDS(comm = scent, distance = "bray")</pre>
## Run 0 stress 0.212491025469
## Run 1 stress 0.212490961771
## ... New best solution
## ... Procrustes: rmse 0.000124038173527 max resid 0.000445047276052
## ... Similar to previous best
## Run 2 stress 0.222517647992
## Run 3 stress 0.211374911381
## ... New best solution
## ... Procrustes: rmse 0.0908135796951 max resid 0.287857012631
## Run 4 stress 0.243883351843
## Run 5 stress 0.212490950078
## Run 6 stress 0.21286277287
## Run 7 stress 0.211374952444
## ... Procrustes: rmse 0.000132619074264 max resid 0.000392554343063
## ... Similar to previous best
```

```
## Run 8 stress 0.217889011628
## Run 9 stress 0.211374919107
## ... Procrustes: rmse 6.55462389298e-05 max resid 0.000195486026936
## ... Similar to previous best
## Run 10 stress 0.21895680594
## Run 11 stress 0.235679395094
## Run 12 stress 0.258084684067
## Run 13 stress 0.212490930286
## Run 14 stress 0.211374915078
## ... Procrustes: rmse 4.43203524851e-05 max resid 0.000121164575607
## ... Similar to previous best
## Run 15 stress 0.211374954904
## ... Procrustes: rmse 0.000102396870182 max resid 0.000302952901169
## ... Similar to previous best
## Run 16 stress 0.235679385533
## Run 17 stress 0.212491041528
## Run 18 stress 0.216772430909
## Run 19 stress 0.211477658673
## ... Procrustes: rmse 0.00869275672358 max resid 0.0298859601908
## Run 20 stress 0.212862832757
## *** Best solution repeated 4 times
## get x and y coordinates
scent_nmds <- as.data.frame(scent_nmds.obj[["points"]])</pre>
## add the colony as a factor to each sample
scent_nmds <- cbind(scent_nmds,colony = peak_factors[["colony"]])</pre>
## quick plotting
ggplot(data = scent_nmds,aes(MDS1,MDS2,color = colony)) +
 geom_point() +
 theme_void() +
  scale_color_manual(values = c("blue","red")) +
  theme(panel.background = element_rect(colour = "black",
                                        size
                                              = 1,
                                        fill = NA),
        aspect.ratio = 1,
        legend.position = "none")
```



Calculate MHC heterozygosity relatedness between individuals

```
## read in mhc genotype data
mhc_het_dat <- read.table("data/clone_mhc_het.txt")</pre>
## restructure `mhc_het_dat`to fit `Demerelate()::inputdata)
## id and colony as factors; alleles as integers or numeric
## otherwise `rxy`cannot handle computations
mhc_het_dat %<>%
  rownames_to_column(., var = "id") %>%
  # mutate(., a1 = str_pad(a1, 2, pad = "0")) %>%
  \# mutate(., a2 = str_pad(a2, 2, pad = "0")) %>%
  mutate(., colony = as.factor(rep("col", 56))) %>%
  mutate(., id = as.factor(id)) %>%
  .[,-4] %>%
  relocate(., colony, .before = a1)
  ## order mhc_het_dat$id after meta$real_id
  ## so data is consistently ordered same in all data.frames
## get matching indeces
id_index <- match(meta$real_id, mhc_het_dat$id)</pre>
## sort correspondingly
mhc_het_dat %<>% .[id_index,]
## calculate relatedness after Queller & Goodnight
mhc_relatedness_res <- Demerelate(inputdata = mhc_het_dat,</pre>
```

```
value = "rxy",
                                  object = T,
                                  NA.rm = F,
                                  Fis = F)
## Warning in Demerelate(inputdata = mhc het dat, value = "rxy", object = T, : Careful, bi-allelic mark
     Especially, rxy and ritland estimator are not defined when bi-allelic estimates are used with alle
     You should consider removing bi-allelics which tend to have very evenly distributed alleles or swi
     Be careful even if allele frequencies are not perfectly 0.5, during randomizations problems may oc
##
mhc_relatedness <- unlist(mhc_relatedness_res$Empirical_Relatedness)</pre>
## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals
relate_mat_mhc <- matrix(nrow = dim(mhc_het_dat)[1],</pre>
                         ncol = dim(mhc_het_dat)[1])
## fill distance matrix row wise, thus fill upper.tri
relate_mat_mhc[upper.tri(relate_mat_mhc)] <- mhc_relatedness</pre>
## transpose to keep consistency with other distance matrices
relate_mat_mhc <- t(relate_mat_mhc)</pre>
relate_mat_mhc %<>% `colnames<-`(meta$real_id) %>% `rownames<-`(meta$real_id)
## vectorize again to identify whether relatedness pairs were consistent in the first place
a <- relate_mat_mhc %>% as.vector() %>% na.omit()
Create vectorized distance measurements for scent data
# bray-curtis distance measurement on scent profiles
scent_dist <- vegdist(scent) %>% as.matrix()
```

Generate UniFrac distances from MHC DQB II individual genotypes

scent_dist[upper.tri(scent_dist, diag = T)] <- NA
b <- scent_dist %>% as.vector() %>% na.omit()

```
# handle genotypes as otu table
phylo_mat <- read.table("data/phyloseq-mat.txt") %>%
    as.matrix()

# make sample names consistent
n <- match(meta$real_id, colnames(phylo_mat))

phylo_mat %<>% .[, n] %>%
    otu_table(., taxa_are_rows = T)

# create phylogenetic tree from file
phylo_tree <- ape::read.tree("data/unifrac_tree_p.nwk")

# merge into Formal class phyloseq
arga_phylseq <- merge_phyloseq(phylo_mat, phylo_tree)

# create UniFrac as genetic diversity measurement for single locus data
mhc_dqb2_ufrac <- UniFrac(arga_phylseq, weighted = F) %>%
    # distances to distance matrix
as.matrix()
```

```
# vectorize distances matrices
mhc_dqb2_ufrac[upper.tri(mhc_dqb2_ufrac, diag = T)] <- NA
c <- mhc_dqb2_ufrac %>% as.vector() %>% na.omit()
```

Calculate microsatellite relatedness values

create data.frame in correspondence to Demerelate input format

```
# read in genotype data table
msats_df <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t")</pre>
# update data.frame with additional info
# "delete" colony info, otherwise relatedness is only calculated for individuals
# within their own colonies -> no complete pairwise comparison
msats_df <- cbind(id = as.factor(rownames(msats_df)),</pre>
                   # colony = meta$colony,
                  colony = rep("col", 56),
                  msats_df[1:56,]) %>%
  # clear df from rownames/ only keep colnames/ variable names
  `rownames<-`(NULL)
msats_df[is.na(msats_df)] = 0
str(msats_df)
# write.table(msats df, file = "data/msats genotypes demerelate.txt",
              sep = " \setminus t",
#
              row.names = F)
msats df %<>% . [match(meta$real id, .$id),]
```

Calculate relatedness of individuals based on Queller & Goodnight

```
## Warning in Demerelate(inputdata = msats_df, value = "rxy", object = T, NA.rm = F, : Careful, bi-alle
## Especially, rxy and ritland estimator are not defined when bi-allelic estimates are used with alle
## You should consider removing bi-allelics which tend to have very evenly distributed alleles or swi
## Be careful even if allele frequencies are not perfectly 0.5, during randomizations problems may oc
## Warning in prop.test(c(emp, non), c(sum(table(empirical.list)),
## sum(table(relate.non.X.mean))), : Chi-Quadrat-Approximation kann inkorrekt sein
```

Coerce output to a vector

```
relatedness <- unlist(relatedness_results$Empirical_Relatedness)
## fill distant matrix / make sure that it follows same systematics as previous distance matrices
## create empty matrix with equal rows and cols similar to sample size of indidivuals</pre>
```

Analyse Odour and genetic association by MHC DQB II and neutral genomic background

Create data.frame to plot in ggplot2

```
## substitute once tested correctly
## scent_mds shall contain similarity values but `b` contains
## dissimilarity values based on Bray-Curtis -> substracting
## dissmilarities from 1 returns similarities

model_rel.df <- cbind(mhc_rel = a, scent_mds = 1-b, ufrac = c, rel = d) %>% as.data.frame()
```

Custom theme to make plot aesthetics consistent

Model odour relationship on MHC and neutral genetic background

Pool underlying data dependencies

Create a function that generates pairwise variables in a systematic matter for pairwise comparisons

```
create_pair_vars <-function(row_cross, col_cross, split_vars = F){
    require(stringr)

    rc <- row_cross
    cc <- col_cross

# create empy matrix
# keep row and col names from existing distance matrices

empty_mat <- matrix(nrow = length(rc),</pre>
```

```
ncol = length(cc)) %>%
    `colnames<-`(cc) %>%
    `rownames<-`(rc)
  # fill each matrix i,j-th cell with the crossing from their corresponding
  \# i-th rowname and j-th colname
  for (i in 1:dim(empty_mat)[1]) {
    for (j in 1:dim(empty_mat)[2]) {
      empty_mat[i,j] <- paste0(rc[i], "/", cc[j])</pre>
    } # end j
  } # end i
  # delete `upper.tri()` of `empty_mat` to resemble structure of the other
  # distance matrices in use
  empty_mat[upper.tri(empty_mat, diag = T)] <- NA</pre>
  pair_vars <- empty_mat %>% as.vector() %>% na.omit()
  # split `pair_vars` if needed
  if (split_vars == T) {
    pair_vars1 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][1]
                          })
    pair_vars2 <- sapply(pair_vars,</pre>
                          function(x){
                            str_split(x, pattern = "/")[[1]][2]
    pair_vars_split <- list(pair_variable1 = pair_vars1,</pre>
                             pair_variable2 = pair_vars2)
    return(pair_vars_split)
  } else {
    return(pair_vars)
} #end create_pair_vars
```

Helper function to combine double entries

```
## for x, overwrite specified replacer with specified value
f <- function(x, replacer, overwrite){
  if (x == replacer) {
    x <- overwrite
  } else {
    x <- x
}</pre>
```

}

Transform model variables

```
agePaired <- create_pair_vars(row_cross = meta$maturity,</pre>
                                col_cross = meta$maturity) %>%
  sapply(., f, "P/M", "M/P")
colonyPaired <- create_pair_vars(row_cross = meta$colony,</pre>
                                   col_cross = meta$colony) %>%
  sapply(., f, "FWB/SSB", "SSB/FWB")
colonyID1 <- create pair vars(row cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[1] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
colonyID2 <- create_pair_vars(row_cross = meta$colony,</pre>
                                col_cross = meta$colony,
                                split_vars = T)[2] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
colonyBool <- ifelse(colonyID1 == colonyID2, 1, 0)</pre>
familyPaired <- create_pair_vars(row_cross = meta$family,</pre>
                                   col cross = meta$family)
familyID1 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[1] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
familyID2 <- create_pair_vars(row_cross = meta$family,</pre>
                                col_cross = meta$family,
                                split_vars = T)[2] %>%
  unlist() %>%
  paste0("f", .) %>%
  as.vector()
pairID1 <- create_pair_vars(row_cross = meta$real_id,</pre>
                              col_cross = meta$real_id,
                              split_vars = T)[1] %>%
  unlist() %>%
  as.vector()
pairID2 <- create_pair_vars(row_cross = meta$real_id,</pre>
                              col_cross = meta$real_id,
                              split_vars = T)[2] %>%
```

```
unlist() %>%
as.vector()

familyBool <- ifelse(familyID1 == familyID2, 1, 0)</pre>
```

Update data.frame with model variables

update data frame with meta data

colony = as.factor(colony),

Include information about MHC heterozygosity, sMLH from microsatellite data and chemical diversity by number of compounds per individual

```
scent.abs <- ifelse(scent != 0, 1, 0)</pre>
compound_n <- apply(scent.abs, 1, sum)</pre>
names(compound n) == meta$real id
# read in heterzygosity information
het_table <- read.table("data/arga_mhc_het.txt", sep = "\t")</pre>
# keep names consistent
match_het <- match(meta$real_id, rownames(het_table))</pre>
het_table %<>% .[match_het,]
# generate sMLH with microsatellite data
# table is pre-prepped, thus rows correspond to same individuals in meta data
smlh_res <- read.table("data/msats_genotypes_inbreedR.txt", sep = "\t")</pre>
smlh res <- smlh res [match(meta$real id, rownames(smlh res)), ] %>%
 # convert to inbreedR format
 convert_raw() %>%
 # generate sMLH
 sMLH()
meta %<>% cbind(., compound_n = compound_n,
              mhc_het = het_table$het,
              smlh = smlh_res)
meta %<>% mutate(
 real_id = as.factor(real_id),
```

```
maturity = as.factor(maturity),
family = as.factor(family)
)
```

Repeat models for chemical similarity

```
# without age and indicator for family groupings
# as grouping factors must have >1 sampled level
# mhc
e1 <- lmerTest::lmer(scent_mds ~ ufrac + colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# relatedness
e2 <- lmerTest::lmer(scent_mds ~ rel + colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# mhc & relatedness
e3 <- lmerTest::lmer(scent_mds ~ rel + ufrac + colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# no genetic effect
e4 <- lmerTest::lmer(scent_mds ~ colonyBool + (1|pairID1) + (1|pairID2),
                     data = model_rel.df)
# compare model performance scores
compare_performance(e1, e2, e3, e4, rank = T) %>%
 arrange(Name)
```

```
## # Comparison of Model Performance Indices
##
## Name |
                    Model | R2 (cond.) | R2 (marg.) | ICC | RMSE | Sigma | AIC weights | AICc weight
        | lmerModLmerTest |
                                 0.517 |
                                              0.188 | 0.404 | 0.058 | 0.062 |
                                                                                     0.287 |
                                                                                                    0.28
## e1
                                              0.186 | 0.417 | 0.058 | 0.062 |
## e2
        | lmerModLmerTest |
                                 0.525 |
                                                                                     0.207 |
                                                                                                    0.20
       | lmerModLmerTest |
                                 0.520 |
                                              0.191 | 0.407 | 0.058 | 0.062 |
                                                                                     0.228 |
                                                                                                    0.21
## e3
      | lmerModLmerTest |
                                 0.522 |
                                              0.184 | 0.414 | 0.058 | 0.062 |
                                                                                     0.278 |
                                                                                                    0.29
summary(e2)
```

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ rel + colonyBool + (1 | pairID1) + (1 | pairID2)
     Data: model_rel.df
##
##
## REML criterion at convergence: -669.2
## Scaled residuals:
                                     Median
##
                           1Q
## -2.8734024086 -0.6551149100 -0.0127392185 0.5907406140 2.6919018000
## Random effects:
                                      Std.Dev.
## Groups Name
                        Variance
## pairID1 (Intercept) 0.00114373121 0.0338190954
## pairID2 (Intercept) 0.00158812032 0.0398512273
## Residual
                        0.00382618866 0.0618561934
```

```
## Number of obs: 276, groups: pairID1, 23; pairID2, 23
##
## Fixed effects:
                               Std. Error
##
                   Estimate
                                                    df t value
                                                                Pr(>|t|)
## (Intercept)
               0.2930925713 0.0159793884 29.8480662643 18.34191 < 2.22e-16
               ## rel
## colonyBool1
               ##
## (Intercept) ***
## rel
## colonyBool1 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
             (Intr) rel
              0.108
## rel
## colonyBool1 -0.639 -0.031
summary(e4)
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: scent_mds ~ colonyBool + (1 | pairID1) + (1 | pairID2)
     Data: model_rel.df
##
## REML criterion at convergence: -671.8
## Scaled residuals:
##
                                  Median
## -2.8754878831 -0.6724901336 -0.0272566346 0.5651130445 2.6590829271
##
## Random effects:
                      Variance
                                   Std.Dev.
## Groups
## pairID1 (Intercept) 0.00114074594 0.0337749306
## pairID2 (Intercept) 0.00156895325 0.0396100145
                      0.00383649427 0.0619394403
## Residual
## Number of obs: 276, groups: pairID1, 23; pairID2, 23
##
## Fixed effects:
                  Estimate
                             Std. Error
                                                 df t value
                                                              Pr(>|t|)
## (Intercept) 0.2910051000 0.0158359198 29.1599058492 18.37627 < 2.22e-16 ***
## colonyBool1 0.0767098586 0.0135826001 52.0344825538 5.64766 6.8802e-07 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
             (Intr)
## colonyBool1 -0.640
```

Repeat models for chemical diversity

```
# exclude family as random effects (no accounting of same family in different
# individuals)
```

```
# exlude age as only one 'age' group being moms is modelled
f1 <- lm(compound_n ~ mhc_het + colony , data = meta)
f2 <- lm(compound_n ~ smlh + colony, data = meta)
f3 <- lm(compound_n ~ mhc_het + smlh + colony, data = meta)
f4 <- lm(compound n ~ colony, data = meta)
compare_performance(f1, f2, f3, f4, rank = T) %>% arrange(Name)
## # Comparison of Model Performance Indices
##
## Name | Model |
                  R2 | R2 (adj.) | RMSE | Sigma | AIC weights | AICc weights | BIC weights | Perf
## -----
         lm | 0.055 |
                        -0.035 | 18.471 | 19.746 |
                                                       0.071 |
                                                                      0.070 |
## f1
                                                                                   0.067 |
         lm | 0.201 |
                       0.125 | 16.980 | 18.152 |
                                                       0.536 |
                                                                      0.526 |
                                                                                   0.503 |
## f3
      lm | 0.202 |
                         0.082 | 16.973 | 18.593 |
                                                       0.199 |
                                                                      0.106 |
                                                                                   0.104 |
                        0.012 | 18.471 | 19.293 |
           lm | 0.055 |
## f4
      0.193 |
                                                                      0.298 |
                                                                                   0.327 |
summary(f2)
##
## Call:
## lm(formula = compound_n ~ smlh + colony, data = meta)
## Residuals:
##
          Min
                       1Q
                               Median
                                               30
                                                          Max
## -33.84774747 -10.72822530 -2.63667025 13.89024838 32.64888040
## Coefficients:
                           Std. Error t value Pr(>|t|)
##
                 Estimate
## (Intercept) -32.76603326 46.95021186 -0.69789 0.49290
              89.23084048 45.46613021 1.96258 0.06308 .
## smlh
## colonySSB
              -3.59128851
                          7.91958429 -0.45347 0.65486
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 18.1520925 on 21 degrees of freedom
## Multiple R-squared: 0.201400231, Adjusted R-squared: 0.12534311
## F-statistic: 2.64801282 on 2 and 21 DF, p-value: 0.0942879933
Session information
```

```
sessionInfo()
## R version 4.3.1 (2023-06-16 ucrt)
## Platform: x86 64-w64-mingw32/x64 (64-bit)
## Running under: Windows 11 x64 (build 22621)
## Matrix products: default
```

```
##
##
## locale:
## [1] LC_COLLATE=German_Germany.utf8 LC_CTYPE=German_Germany.utf8
  [3] LC_MONETARY=German_Germany.utf8 LC_NUMERIC=C
  [5] LC TIME=German Germany.utf8
## time zone: Europe/Berlin
## tzcode source: internal
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                                datasets methods
                                                                    base
## other attached packages:
   [1] remotes_2.4.2.1
                           Demerelate_0.9-3
                                               fts_0.9.9.2
                                                                  zoo_1.8-12
##
    [5] ggbeeswarm_0.7.2
                           partR2_0.9.1
                                               MuMIn_1.47.5
                                                                  performance_0.10.4
##
  [9] ape_5.7-1
                                                                  lattice_0.21-8
                           ggpubr_0.6.0
                                               vegan_2.6-4
## [13] permute 0.9-7
                            inbreedR 0.3.3
                                               GCalignR_1.0.5
                                                                  phyloseq_1.44.0
## [17] lubridate_1.9.2
                           forcats_1.0.0
                                               stringr_1.5.0
                                                                  dplyr_1.1.2
## [21] purrr_1.0.1
                           readr 2.1.4
                                               tidyr_1.3.0
                                                                  tibble_3.2.1
## [25] ggplot2_3.4.2
                           tidyverse_2.0.0
                                               magrittr_2.0.3
##
## loaded via a namespace (and not attached):
##
     [1] rstudioapi 0.15.0
                                  jsonlite 1.8.7
                                                          datawizard 0.8.0
##
     [4] farver_2.1.1
                                  nloptr 2.0.3
                                                          rmarkdown 2.23
     [7] ragg_1.2.5
                                  zlibbioc_1.46.0
                                                          vctrs 0.6.3
##
    [10] multtest_2.56.0
                                                          RCurl_1.98-1.12
                                  minqa_1.2.5
##
   [13] rstatix_0.7.2
                                  htmltools_0.5.5
                                                          broom_1.0.5
##
                                                          rhdf5_2.44.0
   [16] Rhdf5lib_1.22.0
                                  Formula_1.2-5
   [19] plyr_1.8.8
                                  sfsmisc_1.1-15
                                                          igraph_1.5.0.1
##
   [22] lifecycle_1.0.3
                                  iterators_1.0.14
                                                          pkgconfig_2.0.3
##
   [25] Matrix_1.6-0
                                  R6_2.5.1
                                                          fastmap_1.1.1
  [28] GenomeInfoDbData_1.2.10
                                 rbibutils_2.2.13
                                                          digest_0.6.33
  [31] numDeriv_2016.8-1.1
                                  colorspace_2.1-0
                                                          patchwork_1.1.2
##
   [34] S4Vectors 0.38.1
                                  textshaping 0.3.6
                                                          labeling_0.4.2
##
  [37] fansi_1.0.4
                                  timechange_0.2.0
                                                          abind_1.4-5
  [40] mgcv 1.8-42
                                  compiler 4.3.1
                                                          withr 2.5.0
##
  [43] backports_1.4.1
                                  carData_3.0-5
                                                          mlogit_1.1-1
   [46] highr_0.10
                                  ggsignif_0.6.4
                                                          MASS_7.3-60
##
## [49] biomformat_1.28.0
                                  tools_4.3.1
                                                          vipor_0.4.5
## [52] lmtest 0.9-40
                                  beeswarm 0.4.0
                                                          glue_1.6.2
## [55] nlme 3.1-162
                                 rhdf5filters\_1.12.1
                                                          grid_4.3.1
##
   [58] cluster 2.1.4
                                  reshape2_1.4.4
                                                          ade4_1.7-22
##
  [61] see_0.8.0
                                  generics_0.1.3
                                                          gtable_0.3.3
  [64] tzdb_0.4.0
                                  data.table_1.14.8
                                                          hms_1.1.3
##
   [67] car_3.1-2
                                  utf8_1.2.3
                                                          XVector_0.40.0
##
   [70] BiocGenerics_0.46.0
                                  ggrepel_0.9.3
                                                          foreach_1.5.2
##
  [73] pillar_1.9.0
                                  splines_4.3.1
                                                          survival_3.5-5
  [76] tidyselect_1.2.0
                                  pbapply_1.7-2
                                                          Biostrings_2.68.1
## [79] knitr_1.43
                                  IRanges_2.34.1
                                                          stats4_4.3.1
## [82] xfun_0.39
                                  Biobase_2.60.0
                                                          statmod_1.5.0
## [85] dfidx 0.0-5
                                  stringi_1.7.12
                                                          yaml_2.3.7
## [88] boot_1.3-28.1
                                  evaluate_0.21
                                                          codetools_0.2-19
## [91] cli_3.6.1
                                  systemfonts 1.0.4
                                                          Rdpack 2.4
```

| ## | [94] | munsell_0.5.0 | Rcpp_1.0.11 | <pre>GenomeInfoDb_1.36.1</pre> |
|----|-------|----------------|---------------------------|--------------------------------|
| ## | [97] | parallel_4.3.1 | bayestestR_0.13.1 | bitops_1.0-7 |
| ## | [100] | lme4_1.1-34 | <pre>lmerTest_3.1-3</pre> | scales_1.2.1 |
| ## | [103] | insight_0.19.3 | crayon_1.5.2 | rlang_1.1.1 |
| ## | Γ1067 | cowplot 1.1.1 | | |

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