# R-code for "Chemical fingerprints encode mother-offspring similarity, colony membership, relatedness and genetic quality in fur seals"

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This document provides the code for all major analysis in our paper. Duplicate analyses that have not been part of an argument (e.g. most analysis for pups) were strapped out for readability. Both the Rmarkdown file and the data can be assessed directly from the hyperlinks in the methods section of the paper (Dataset S1, Dataset S2) or by downloading this GitHub repository. Just click on the link and then on *Download ZIP* on the right-hand side of the page. Make sure you read the instructions in the xlsx sheet if you downloaded the data from the paper or from the Readme file on GitHub if you downloaded the repository. If you have any questions, don't hesitate contacting me: martin.adam.stoffel[at]gmail.com.

The sequence of code follows roughly the sequence of analysis in the Results section of the paper.

# Prerequisites:

- Some of the functions in the analysis (g2, sMLH) are part of the inbreedR package, that we wrote in parallel to the analyses. The inbreedR package provides a framework for calculating inbreeding and HFCs from genetic markers (SNPs and microsatellites) and will soon be published on CRAN. To download the newest version of the package from the developer platform GitHub, you need to install the devtools package as shown in the next section.
- For running the complete code you need a files subfolder with all the raw data files.

Installing inbreedR from GitHub.

```
# install.packages("devtools")
library(devtools)
# uncomment the next line for downloading and installing inbreedR
# install_github("mastoffel/inbreedR")
library(inbreedR)
```

See ?inbreedR for further information on the functions or type browseVignettes(package = "inbreedR") to have a look at the vignette.

# Loading data, standardisation and transformation

Loading the

- chemical data (scent\_raw, called scent data from now), which is the output of Gas-chromatogramm peak detection done in Xcalibur 2.0.5. (A first preprocessing was done by aligning the raw chemical data and removing substances that have been present in the control sample, see Methods part of the paper)
- identities for colony membership (colony), mother-offspring pairs (family) and mothers and pups, respectively (age)

```
scent_raw <- as.data.frame(t(read.csv(".\\files\\scent_raw.csv", row.names = 1)))</pre>
factors <- read.csv(".\\files\\factors.csv",row.names=1)</pre>
head(factors)
       colony family age
#>
#> M10
             2
                    10
                         1
#> M12
             2
                    12
                         1
             2
                    14
                         1
#> M14
#> M15
             1
                    15
                         1
             2
#> M16
                    16
                         1
#> M17
             2
                    17
```

Standardising observations by total, such that within every observation compounds add up to 100% (Thus averaging out absolute concentration differences between samples)

```
scent_stand <- as.data.frame(t(apply(scent_raw, 1, function(x) (x/sum(x)) * 100)))</pre>
```

Log(x+1) transformation of the standardised scent data.

```
scent <- log(scent_stand + 1)</pre>
```

The scent matrix contains 82 observations and 213 compounds (retention times of chemicals are column names, values are relative concentrations) in total.

```
dim(scent)
#> [1] 82 213
head(scent[1:6])
       8.061111111
                       8.23 8.307142857 8.394 8.47375 8.516153846
#> M10
          0.000000 0.000000
                              0.0000000
                                            0 0.000000
                                                         0.6562090
#> M12
          0.000000 0.000000
                                            0 0.000000
                                                         0.0000000
                              0.4864961
#> M14
         3.222626 1.665421
                              0.0000000
                                            0 0.000000
                                                         0.0000000
#> M15
          0.000000 0.000000
                              0.0000000
                                            0 0.000000
                                                         0.0000000
#> M16
         0.000000 0.000000
                              0.6849915
                                            0 1.008018
                                                         0.5654895
#> M17
         2.330450 0.000000
                              0.0000000
                                            0 0.000000
                                                         0.0000000
```

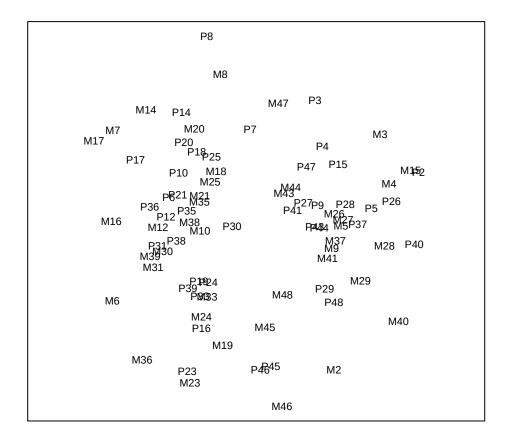
# Colony differences in chemical fingerprints

```
library(vegan)
library(MASS)
```

Non-metric multidimensional scaling (nMDS) visualizes a distance matrix (Bray-Curtis similarity). The nMDS algorithm aims to place each individual in a 2-dimensional space such that the between-individual distances are preserved as well as possible. Axis coordinates are arbitrary and not shown. The plot is better visualized with colours (see paper) and is shown here for the purpose of demonstration. Mother-offspring pairs can be identified by labels (e.g. M14, P14).

```
scent_mds <- MASS::isoMDS(vegdist(scent))
#> initial value 28.002906
#> iter    5 value 21.594484
#> final value 21.345037
#> converged
```

```
vegan::ordiplot(scent_mds, type = "t", ylab = "", xlab = "",axes=FALSE, frame.plot=TRUE)
```



Analysis of Similarities (ANOSIM) is a non-parametric test for group differences based on a Bray-curtis (or any other) similarity matrix. We use the vegan package (Oksanen et al. 2015) for ANOSIM and several other functions. Most analysis are done for the whole sample as well as for mothers and pups seperately to avoid pseudoreplication. ANOSIM is based on a permutation test, which is why results can slightly differ from the paper.

Dissimilarity between the two colonies.

Dissimilarity between mothers from the two colonies.

Dissimilarity between pups from the two colonies.

Genetic differentiation of the two colonies was assessed through bayesian structure analysis, with the software "Structure" (Pritchard, Stephens, and Donnelly 2000)

Mother offspring similarity in chemical fingerprints.

Full sample

### Mother offspring similarity within colony 1 (Special study beach)

1, ]\$fam

2, ]\$fam

# Mother offspring similarity within colony 2 (Freshwater beach)

## Chemical similarity vs. geographic distance on special study beach

• location data in meters is available for this population as the special study beach on Bird Island provides an aerial walkway

Loading X-Y coordinates of each individual.

Converting coordinates to pairwise euclidian distance matrix.

```
dist_mat <- as.matrix(dist(coord, method = "euclidian"))</pre>
```

Constructing a bray curtis similarity matrix (from chemical fingerprints) of all individuals from beach 1 (special study beach). We constantly used spearman rank correlation in mantel tests.

### Geographic distance vs. chemical similarity in mothers

```
geo mum <- dist mat[1:20, 1:20]
scent_mum <- scent_bc[1:20, 1:20]</pre>
vegan::mantel(geo_mum, scent_mum, method = "spearman")
#> Mantel statistic based on Spearman's rank correlation rho
#>
#> Call:
#> vegan::mantel(xdis = geo_mum, ydis = scent_mum, method = "spearman")
#>
#> Mantel statistic r: 0.008091
#>
        Significance: 0.475
#>
#> Upper quantiles of permutations (null model):
   90% 95% 97.5%
                     99%
#> 0.200 0.256 0.306 0.359
#> Permutation: free
#> Number of permutations: 999
```

### Geographic distance vs. chemical similarity in pups

```
geo_pup <- dist_mat[21:40, 21:40]
scent_pup <- scent_bc[21:40, 21:40]
vegan::mantel(geo_pup, scent_pup, method = "spearman")
#>
#> Mantel statistic based on Spearman's rank correlation rho
#>
#> Call:
#> vegan::mantel(xdis = geo_pup, ydis = scent_pup, method = "spearman")
```

```
#> Mantel statistic r: 0.06039

#> Significance: 0.304

#>

#> Upper quantiles of permutations (null model):

#> 90% 95% 97.5% 99%

#> 0.164 0.222 0.268 0.329

#> Permutation: free

#> Number of permutations: 999
```

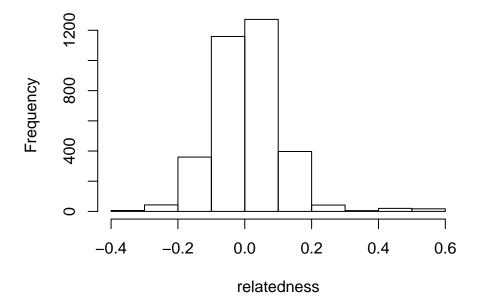
# Correlation between genotype and overall chemical fingerprints.

## Relatedness and overall chemical similarity

Load pairwise relatedness (Queller and Goodnight 1989) based on 41 microsatellite markers.

```
relatedness <- as.matrix(read.csv(".\\files\\relatedness.csv",row.names=1))</pre>
head(relatedness[1:6, 1:6])
#>
               M10
                            M12
                                                     M15
                                                                 M16 M17
                                        M14
#> M10
                NA
                             NA
                                         NA
                                                      NA
                                                                  NA NA
#> M12 -0.09578940
                             NA
                                         NA
                                                      NA
                                                                      NA
                                                                  NA
#> M14 -0.10861601 -0.16464236
                                                      NA
                                                                      NA
                                                                  NA
#> M15 -0.03246021 -0.11981456 -0.12591268
                                                                  NA
                                                                      NA
#> M16 0.07639825 0.13027995 0.01176970 -0.02336469
                                                                  NA
                                                                      NA
#> M17  0.04367833  -0.09591802  -0.06258925  0.03730164  -0.08061711
hist(relatedness)
```

# Histogram of relatedness



Pairwise bray curtis similarity in chemical fingerprints of all individuals.

Mantel test between genetic relatedness and bray curtis similarity in chemical fingerprints of all individuals.

```
vegan::mantel(relatedness, scent bc, method = "spearman", permutation = 1000)
#> Mantel statistic based on Spearman's rank correlation rho
#>
#> Call:
#> vegan::mantel(xdis = relatedness, ydis = scent_bc, method = "spearman",
                                                                                permutations = 1000)
#> Mantel statistic r: 0.07231
#>
        Significance: 0.005994
#>
#> Upper quantiles of permutations (null model):
#>
     90%
           95% 97.5%
                           99%
#> 0.0398 0.0487 0.0573 0.0632
#> Permutation: free
#> Number of permutations: 1000
```

We find a significant relationship between the overall chemical fingerprints and genetic relatedness. However, we are likely to have a problem of pseudoreplication here. For that reason, we are analysing mothers and pups seperately.

Fur seal mothers: mantel test between genetic relatedness and bray curtis similarity of chemical fingerprints.

```
vegan::mantel(relatedness[factors$age == 1, factors$age == 1],
              scent_bc[factors$age == 1, factors$age == 1],
             method = "spearman", permutation = 1000)
#>
#> Mantel statistic based on Spearman's rank correlation rho
#> Call:
#> vegan::mantel(xdis = relatedness[factors$age == 1, factors$age ==
                                                                         1], ydis = scent_bc[factors$a
#>
#> Mantel statistic r: 0.05938
#>
        Significance: 0.10689
#> Upper quantiles of permutations (null model):
     90%
            95% 97.5%
#> 0.0603 0.0757 0.0900 0.1059
```

```
#> Permutation: free
#> Number of permutations: 1000
```

Fur seal pups: mantel test between genetic relatedness and bray curtis similarity of chemical fingerprints.

```
vegan::mantel(relatedness[factors$age == 2, factors$age == 2],
             scent_bc[factors$age == 2, factors$age == 2],
             method = "spearman", permutation = 1000)
#>
#> Mantel statistic based on Spearman's rank correlation rho
#>
#> Call:
#> vegan::mantel(xdis = relatedness[factors$age == 2, factors$age ==
                                                                         2], ydis = scent bc[factors$a
#> Mantel statistic r: 0.02985
#>
        Significance: 0.25475
#>
#> Upper quantiles of permutations (null model):
     90%
            95% 97.5%
                          99%
#> 0.0587 0.0732 0.0894 0.1055
#> Permutation: free
#> Number of permutations: 1000
```

# Correlation between heterozygosity (sMLH) and diversity (number of compounds) of chemical fingerprints

• The function sMLH is part of the inbreedR package, currently available on GitHub.

```
library(inbreedR)
# ?inbreedR
```

Loading raw genotypes and calculating standardised multilocus heterozygosity (sMLH) based on 41 markers.

\* inbreedRpackage requires a special format, see ?convert raw for more information'\*

```
genotypes <- read.table(".\\files\\genotypes.txt", row.names=1)
genotypes[1:6, 1:6]
#> V2 V3 V4 V5 V6 V7
#> M10 168 184 237 241 164 164
#> M12 168 168 233 233 164 164
#> M14 168 170 231 237 164 166
#> M15 168 182 227 229 164 164
#> M16 168 168 231 233 164 164
#> M17 166 172 231 237 164 164
genotypes_formatted <- inbreedR::convert_raw(genotypes, miss = NA)
heterozygosity <- inbreedR::sMLH(genotypes_formatted)</pre>
```

Number of compounds per individual.

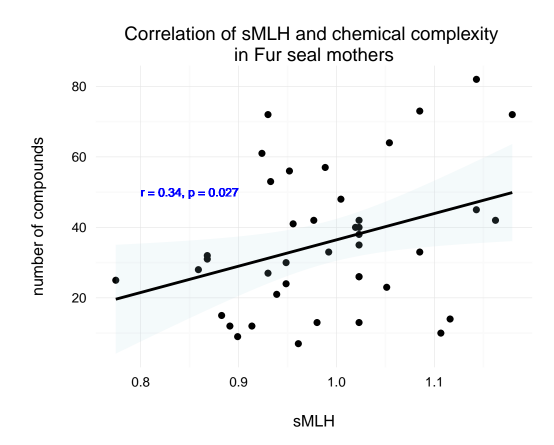
```
num_comp <- as.vector(apply(scent, 1, function(x) length(x[x>0])))
```

### Linear model of heterozygosity on number of compounds in mothers

A clear association between sMLH and chemical complexity in mothers but not pups.

```
het_mum <- heterozygosity[factors$age == 1]</pre>
num_comp_mum <- num_comp[factors$age==1]</pre>
summary(lm(het_mum ~ num_comp_mum))
#>
#> Call:
#> lm(formula = het_mum ~ num_comp_mum)
#>
#> Residuals:
#>
       Min
                 1Q
                      Median
                                   3Q
#>
#> Coefficients:
#>
               Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 0.9339147 0.0282437 33.066
                                         <2e-16 ***
#> num_comp_mum 0.0015914 0.0006936
                                           0.0272 *
                                   2.294
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> Residual standard error: 0.08633 on 39 degrees of freedom
#> Multiple R-squared: 0.1189, Adjusted R-squared: 0.09633
\#> F-statistic: 5.264 on 1 and 39 DF, p-value: 0.02724
```

Plotting with ggplot2, an implementation of the grammar of graphics (Wickham 2009)



## Linear model of heterozygosity on number of compounds in pups

```
het_pup <- heterozygosity[factors$age == 2]</pre>
num_comp_pup <- num_comp[factors$age==2]</pre>
summary(lm(het_pup ~ num_comp_pup))
#>
#> Call:
#> lm(formula = het_pup ~ num_comp_pup)
#>
#> Residuals:
#>
                   1Q
                         Median
                                       3Q
#> -0.199070 -0.067303 -0.001971 0.050500 0.152902
#>
#> Coefficients:
                Estimate Std. Error t value Pr(>|t|)
#>
#> (Intercept) 0.9940863 0.0235812 42.156
                                            <2e-16 ***
#> num_comp_pup 0.0003928 0.0005542
                                     0.709
                                               0.483
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> Residual standard error: 0.08035 on 39 degrees of freedom
#> Multiple R-squared: 0.01272, Adjusted R-squared: -0.0126
\#> F-statistic: 0.5025 on 1 and 39 DF, p-value: 0.4826
```

Strength of correlation between sMLH and number of compounds increases with an increasing number of genetic markers in mothers.

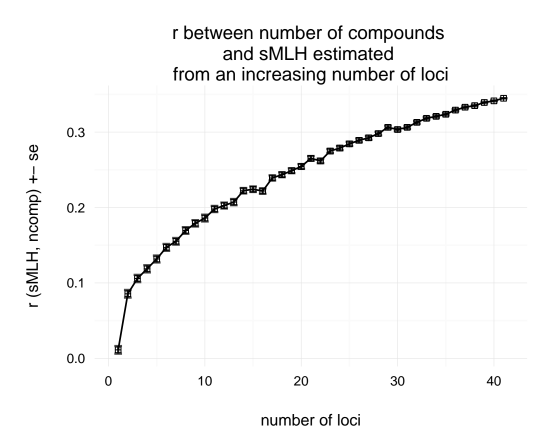
The resample\_loci() function samples an increasing subset of loci, calculates sMLH and correlates with a vector y (here: number of compounds in chemical fingerprints).

```
resample_loci <- function(genotypes, y, num_iter = 1000) {</pre>
# genotypes in inbreedR format. See ?inbreedR
# y is a vector to correlate with sMLH
# num iter is the number of resamplings per added locus
        # calculate number of loci
        num_loci <- ncol(genotypes)</pre>
        results <- data.frame(matrix(nrow = num iter, ncol = num loci))
        for (i in seq_along((1: num_loci))){
                for (k in seq_along(1:num_iter)) {
                loci_ind <- sample(1:num_loci, i, replace = FALSE)</pre>
                het <- inbreedR::sMLH(genotypes[, loci_ind])</pre>
                results[k, i] <- cor(het[1:41],y) # heterozygosity subsetted for mothers
results
}
# Converting genotypes into the right format
genotypes_formatted <- inbreedR::convert_raw(genotypes, miss = NA)</pre>
# Resampling 1 - 40 loci each 1000 times, compute sMLH and correlate with number of compounds
resample_mums <- resample_loci(genotypes_formatted, num_comp_mum, num_iter = 1000)
```

Calculating summary statistics for the resampling output: mean, sd, se of the correlations per subset of markers.

Plotting mean correlation of heterozygosity (estimated by an increasing number of markers) with number of compounds in chemical fingerprints for Fur seal mothers.

Pups are not shown here for simplicity and to avoid code replication. For the full figure see the results section of the paper



# Estimation of identity disequilibrium g2 with the inbreedR package. (can diverge slightly from the RMES program)

Instead to just finding a correlation between heterozygosity and a trait such as chemical complexity, one can ask whether variation in inbreeding (so called-general effects) is a potential cause. This can be measured with a parameter called g2 (David et al. 2007), that assesses identity disequilibrium through quantification of excess double heterozygote loci. We are currently working on the inbreedR package, which provides functions for calculation g2 with both microsatellites and SNPs.

Calculate g2.

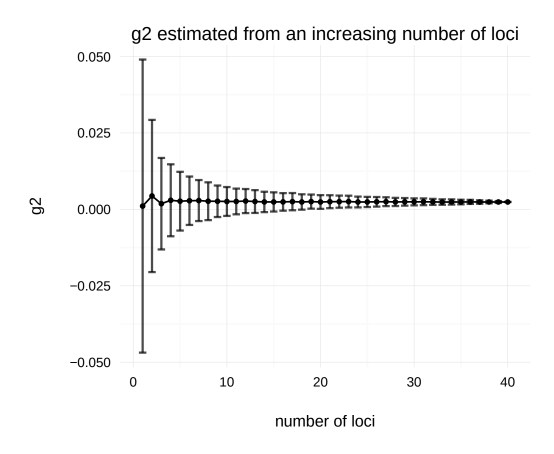
Flexibility of the g2 functions in the inbreedR package allow for further analysis of the parameter. A good start is estimate g2 from an increasing number of subsampled loci.

```
resampling_g2 <- resample_loc_g2(genotypes_formatted, niter = 1000)</pre>
```

```
results_g2 <- sum_results(resampling_g2)
```

Plotting the mean +- sd of g2 estimated from increasing amount of microsats.

```
# plotting
library(grid)
ggplot2::ggplot(results_g2, aes(x = locnum, y = cormean)) +
        geom_line(size = 0.6, colour = "black") +
        geom_errorbar(aes(ymin = cormean-corsd, ymax = cormean+corsd),
                      width=0.8, alpha=0.7, size = 0.8, colour = "black") +
        geom_point(size = 2, shape = 16) +
        theme_minimal(base_size = 12) +
        theme(axis.title.x = element_text(vjust= -2 ,size = 12),
              axis.title.y = element_text(vjust=3,size = 12),
              axis.ticks.x = element_blank(),
              axis.ticks.y = element_blank(),
              plot.margin = (unit(c(.5, .5, 2, 2), "cm"))) +
        #geom_hline(yintercept=0.305) +
        ylab("g2") +
        xlab("number of loci") +
        labs(title = "g2 estimated from an increasing number of loci")
```



# Factor analysis on the chemical compounds data with the package HDMD.

HDMD (McFerrin 2013) allows for doing a Factor analysis with high dimensional data(where the number of variables exceeds the number of observations) by calculating a general inverse matrix.

```
library(HDMD)
library(minmodelr)
```

Factor analysis and extraction of factor scores for the first 4 factors. Promax rotation of the factors allows them to be non-orthogonal and thus correlated. After FA, the factor scores for each individual on all 4 factors are extracted.

```
fa_scores <- as.data.frame(scent_fa$scores)

head(fa_scores)

#> F1 F2 F3 F4

#> M10 0.21501860 -0.8384633 0.02809373 0.41819811

#> M12 0.00144057 -0.4989538 0.19093716 0.63217100

#> M14 -0.32163481 -0.7076035 0.16407820 -0.18508919

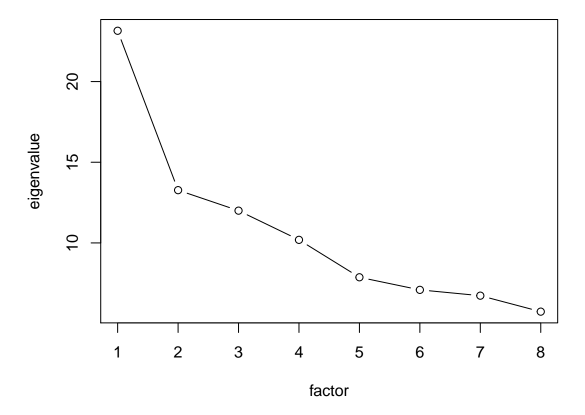
#> M15 -0.34857816 -0.5187462 0.12530472 -0.93750096

#> M16 -0.38997409 -0.1759757 0.53663454 1.18679542

#> M17 -0.10514763 -0.6294724 0.18209318 0.02876237
```

The eigenvalue course seen in the screeplot allows for decisions on the number of factors to retain.

# **Screeplot**



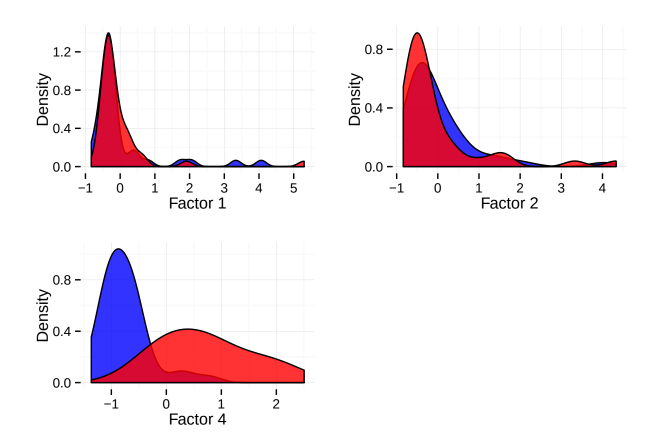
Plotting the distribution of factor scores seperately for each colony. Similar distributions suggest the compounds which are represented by a given factor to be similarly distributed across colonies and could thus be of potential genetic origin, while different distributions as in factor 4 suggest this factor to represent environmentally influenced compounds.

# Multiplot function from cookbook-r.com for plotting multiple ggplots

```
multiplot <- function(..., plotlist=NULL, cols) {</pre>
  require(grid)
  \# Make a list from the ... arguments and plotlist
  plots <- c(list(...), plotlist)</pre>
 numPlots = length(plots)
  # Make the panel
  plotCols = cols
                                            # Number of columns of plots
 plotRows = ceiling(numPlots/plotCols) # Number of rows needed, calculated from # of cols
  # Set up the page
  grid.newpage()
  pushViewport(viewport(layout = grid.layout(plotRows, plotCols)))
  vplayout <- function(x, y)</pre>
    viewport(layout.pos.row = x, layout.pos.col = y)
  # Make each plot, in the correct location
  for (i in 1:numPlots) {
    curRow = ceiling(i/plotCols)
    curCol = (i-1) %% plotCols + 1
    print(plots[[i]], vp = vplayout(curRow, curCol ))
  }
}
```

Plotting all factor distributions.

```
multiplot(f1_plot, f2_plot, f4_plot, cols = 2)
```



Linear model of heterozygosity on factors (factor scores) as explanatory variables in mothers.

```
# bind heterozygosity and the factor scores in one data.frame and subset mothers
het_df <- cbind(heterozygosity, fa_scores)[factors$age == 1, ]</pre>
het_model <- lm(heterozygosity ~., data=het_df)</pre>
summary(het_model)
#>
#> Call:
#> lm(formula = heterozygosity ~ ., data = het_df)
#> Residuals:
#>
        Min
                  1Q
                       Median
#> -0.19439 -0.06271 0.01210 0.04769 0.14674
#>
#> Coefficients:
#>
                Estimate Std. Error t value Pr(>|t|)
                                     75.780
                0.987971
                           0.013037
                                               <2e-16 ***
#> (Intercept)
#> F1
                0.029393
                           0.012288
                                       2.392
                                               0.0221 *
#> F2
                0.027521
                           0.011898
                                       2.313
                                               0.0265 *
#> F3
                0.004033
                           0.012964
                                       0.311
                                               0.7575
                           0.014180
                                               0.5020
#> F4
               -0.009617
                                     -0.678
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> Residual standard error: 0.08315 on 36 degrees of freedom
#> Multiple R-squared: 0.2455, Adjusted R-squared: 0.1616
```

```
#> F-statistic: 2.928 on 4 and 36 DF, p-value: 0.03406
```

While Factor1 and Factor2 seem to represent substances that are accociated with heterozygosity, Factor 3 and Factor 4 clearly don't. To simplify the model we used deletion testing (Crawley, Statistics). The minmodelr package contains some helper functions for this task. See ?MinMod, ?DelTestVar. We don't generally recommend a deletion testing procedure. In our case, results are clear and we use it for simplicity rather than for fishing significant results.

```
library(devtools)
# install_github("mastoffel/minmodelr")
library(minmodelr)
```

```
het_reduced <- minmodelr::MinMod(het_df)
#>
#> Call:
#> glm(formula = depVar ~ ., family = family, data = bestmodeldf)
#> Deviance Residuals:
              1Q
       Min
                           {\it Median}
                                           3Q
                                                     Max
#> -0.191202 -0.060032 0.009789
                                   0.057485 0.155757
#>
#> Coefficients:
#>
              Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 0.98814 0.01278 77.328 <2e-16 ***
#> F1
               0.02783
                          0.01167 2.385
                                             0.0222 *
#> F2
                0.02809
                          0.01164
                                     2.414
                                            0.0207 *
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#> (Dispersion parameter for gaussian family taken to be 0.006649744)
#>
       Null deviance: 0.32986 on 40 degrees of freedom
#> Residual deviance: 0.25269 on 38 degrees of freedom
#> AIC: -84.303
#> Number of Fisher Scoring iterations: 2
# extract data frame
het_reduced_df <- het_reduced[[1]]</pre>
# extract reduced model
het_reduced_mod <- het_reduced[[2]]</pre>
# deletion testing for both variables in the reduced model. See ?DelTestVar
table <- minmodelr::DelTestVar(het reduced df)
                Estimate Deviance Explained
                                                    F P (F-test)
#> (Intercept) 0.98814218
                                                   NA
                                  11.46936 5.689346 0.02215967
#> F1
              0.02783316
#> F2
               0.02808759
                                    11.74642 5.826780 0.02071002
#>
               P (Chisquared-test)
#> (Intercept)
                               NA
#> F1
                        0.01706822
#> F2
                        0.01578399
# deviance explained by the reduced model
dev_expl <- (het_reduced_mod$null.deviance - het_reduced_mod$deviance) / het_reduced_mod$null.deviance
summary(het_reduced_mod)
```

```
#>
#> Call:
#> glm(formula = depVar ~ ., family = family, data = bestmodeldf)
#> Deviance Residuals:
#>
       Min 1Q
                         Median
                                        3Q
#> -0.191202 -0.060032 0.009789 0.057485 0.155757
#>
#> Coefficients:
             Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 0.98814 0.01278 77.328 <2e-16 ***
#> F1
             0.02783
                        0.01167 2.385
                                        0.0222 *
#> F2
              0.02809
                                  2.414 0.0207 *
                         0.01164
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> (Dispersion parameter for gaussian family taken to be 0.006649744)
      Null deviance: 0.32986 on 40 degrees of freedom
#> Residual deviance: 0.25269 on 38 degrees of freedom
#> AIC: -84.303
#>
#> Number of Fisher Scoring iterations: 2
```

Creating a new variable F1F2 which is the sum of the two factor scores and using this variable as predictor in a linear model of heterozygosity.

```
# sum of factors as variable
het_df$F1F2 <- het_df$F1 + het_df$F2</pre>
table <- minmodelr::DelTestVar(as.data.frame(cbind(het_df$heterozygosity, het_df$F1F2)))
#>
               Estimate Deviance Explained F P (F-test)
#> (Intercept) 0.98813169
                                        NA
                                                 NA
#> V2
              0.02796073
                                   23.39391 11.90979 0.001356642
#>
              P (Chisquared-test)
#> (Intercept)
                     0.0005583969
summary(lm(heterozygosity ~ F1F2, data = het_df))
#>
#> Call:
#> lm(formula = heterozygosity ~ F1F2, data = het_df)
#>
#> Residuals:
       Min
                  1Q
                        {\it Median}
                                       3Q
                                                Max
#> -0.191204 -0.060060 0.009766 0.057466 0.155764
#>
#> Coefficients:
              Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 0.988132 0.012596 78.449 < 2e-16 ***
                         0.008102 3.451 0.00136 **
#> F1F2
             0.027961
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Residual standard error: 0.08049 on 39 degrees of freedom
```

```
#> Multiple R-squared: 0.2339, Adjusted R-squared: 0.2143
#> F-statistic: 11.91 on 1 and 39 DF, p-value: 0.001357
```

### Linear model of genetic relatedness on factor scores as explanatory variables for mothers.

Pairwise genetic relatedness is represented as a matrix. To model the relationship between relatedness and factor scores we created a matrix for each factor, whereby each pairwise value represents the difference in factor scores for a pair of seals.

get\_pairdiff() creates these matrices. We based these analysis on mothers and pups seperately.

```
get_pairdiff <- function(relate, scores, df=F) {</pre>
# creates data.frame with
# pairwise differences in factor scores
# input should be: relatedness data frame (lower triangular),
# data frame with factor scores in columns
# if df=TRUE, get_pairdiff will return a list of score-difference
# dataframes (for each component/factor) with pairwise pc-differences.
# make sure to have data.frames
relate <- as.data.frame(relate)</pre>
scores <- as.data.frame(scores)</pre>
# copy similarity matrix and clear
score_mat <- relate</pre>
score_mat[, ] <- NA</pre>
# get vector of pairwise-rownames
allnames <- vector()
for (i in 1:ncol(relate)) {
        for (k in 1:nrow(relate)) {
                 nametemp <- paste(names(relate)[i],</pre>
                                     row.names(relate)[k], sep = "")
                 allnames <- append(allnames, nametemp)</pre>
        }
}
# roll out as vector
relate vec <- unlist(relate)</pre>
# label the rows
names(relate vec) <- allnames</pre>
# delete na´s
relate_vec <- relate_vec[!is.na(relate_vec)]</pre>
# get new row-names vector
pairnamessub <- names(relate_vec)</pre>
# create raw data frame
fac_diff_all <- data.frame("relatedness"= relate_vec)</pre>
# construct similarity matrix out of pairwise differences in factors
names <- rownames(relate)</pre>
row.names(scores) <- names</pre>
fac diff mats <- list()</pre>
for (z in 1:ncol(scores)) {
        for (i in names) {
                 for (k in names) {
                          if (!(is.na(relate[i,k]))) {
                                   diff_fac <- abs(scores[i,z] - scores[k,z])</pre>
                                   score_mat[i,k] <- diff_fac</pre>
```

```
}
        }
        # create list of data frames, containing difference matrices per Factor
        fac_diff_mats <- c(fac_diff_mats, list(score_mat))</pre>
        # turn into vector
        factor_diff <- as.vector(as.matrix(score_mat))</pre>
        factor_diff <- factor_diff[!is.na(factor_diff)]</pre>
        fac_diff_all <- cbind(fac_diff_all, factor_diff)</pre>
## check argument for what to return
if (df == T) {
        names(fac_diff_mats) <- names(scores)</pre>
        return(fac_diff_mats)
} else if (df == F) {
        names(fac_diff_all) <- c("relatedness",names(scores))</pre>
        row.names(fac_diff_all) <- pairnamessub</pre>
        return(fac_diff_all)
}
}
```

Creation of 4 pairwise distance matrices for each factor.

The ecodist package (Goslee and Urban 2007) can handle multiple distance matrices by doing a partial mantel test.

Every partial mantel test just tests for the association with the first response, while the other are permutated

```
rel_dist <- as.dist(relatedness[factors$age == 1, factors$age == 1])</pre>
ecodist::mantel(rel_dist ~ f1_diff + f2_diff + f3_diff + f4_diff, mrank = T, nperm = 1000)
                                 pval3 llim.2.5% ulim.97.5%
    mantelr
               pval1
                        pval2
ecodist::mantel(rel_dist ~ f2_diff + f1_diff + f3_diff + f4_diff, mrank = T)
                             pval2
                                        pval3 llim.2.5%
                  pval1
ulim.97.5\%
#> -0.005076159
ecodist::mantel(rel_dist ~ f3_diff + f2_diff + f1_diff + f4_diff, mrank = T)
    mantelr
                        pval2
                                pval3 llim.2.5% ulim.97.5%
              pval1
#> 0.08900545 0.06100000 0.94000000 0.12200000 0.04792259 0.12570957
ecodist::mantel(rel_dist ~ f4_diff + f3_diff + f2_diff + f1_diff, mrank = T)
                                     pval3 llim.2.5% ulim.97.5%
                 pval1
     mantelr
                           pval2
#> 0.052327208 0.118000000 0.883000000 0.247000000 0.002718941 0.095575727
```

Linear model for associations between genetic relatedness and factor scores as explanatory variables for pups.

```
fa_diff_pups <- get_pairdiff(relatedness[factors$age == 2, factors$age == 2],</pre>
                          fa_scores[factors$age == 2, ], df = F)
for (i in seq along(1:4)) {
       assign(paste("f", i, "_diff", sep=""), fa_diff_pups[, i+1])
}
rel_dist <- as.dist(relatedness[factors$age == 2, factors$age == 2])</pre>
ecodist::mantel(rel_dist ~ f1_diff + f2_diff + f3_diff + f4_diff, mrank = T)
                                    pval3 llim.2.5% ulim.97.5%
      mantelr
                  pval1
                             pval2
#> 0.02435999 0.31700000 0.68400000 0.64300000 -0.02535968 0.05825877
ecodist::mantel(rel_dist ~ f2_diff + f1_diff + f3_diff + f4_diff, mrank = T)
                                               llim.2.5% ulim.97.5%
#>
      mantelr
                  pval1
                             pval2
                                        pval3
#> 0.01340994 0.39600000 0.60500000 0.81500000 -0.02573603 0.05067160
ecodist::mantel(rel_dist ~ f3_diff + f2_diff + f1_diff + f4_diff, mrank = T)
                          pval2
                                   pval3 llim.2.5% ulim.97.5%
     mantelr
                pval1
#> 0.08487309 0.08400000 0.91700000 0.15100000 0.03461989 0.13741796
ecodist::mantel(rel_dist ~ f4_diff + f3_diff + f2_diff + f1_diff, mrank = T)
                  pval1
                           pval2 pval3 llim.2.5% ulim.97.5%
```

Colony differences in factor scores: Just factor 4 shows significant differences.

```
col_df <- cbind(factors["colony"], fa_scores)</pre>
col_reduced <- minmodelr::MinMod(col_df)</pre>
#>
#> Call:
#> glm(formula = depVar ~ ., family = family, data = bestmodeldf)
#>
#> Deviance Residuals:
     Min
           1Q Median
                                 3Q
                                          Max
#> -0.8127 -0.2569 -0.1038 0.2719
                                       0.7243
#>
#> Coefficients:
              Estimate Std. Error t value Pr(>|t|)
#> (Intercept) 1.51220 0.03696 40.91 < 2e-16 ***
              0.38454
                        0.03791 10.14 5.08e-16 ***
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> (Dispersion parameter for gaussian family taken to be 0.1120233)
#>
      Null deviance: 20.4878 on 81 degrees of freedom
#> Residual deviance: 8.9619 on 80 degrees of freedom
#> AIC: 57.179
#> Number of Fisher Scoring iterations: 2
col_reduced_df <- col_reduced[[1]]</pre>
# dev_expl <- (col_reduced_df$null.deviance - col_reduced_df$deviance) / col_reduced_df$null.deviance
table <- minmodelr::DelTestVar(col reduced[[1]])
```

```
#> Estimate Deviance Explained F P (F-test)

#> (Intercept) 1.5121951 NA NA NA

#> F4 0.3845353 56.25756 102.8887 5.082064e-16

#> (Intercept) NA

#> F4 3.545071e-24
```

### Identification of substance subsets.

```
# subsets and identification
library(vegan)
library(ggplot2)
library(dplyr)
library(magrittr)
library(vegan)
library(reshape2)
```

Similarity percentages analysis (simper) identifies the contribution of a specific compound to group similarity / dissimilarity. ANOSIM was used to test whether a small subset of the compounds with the highest contributions shows significant patterns.

### Identification of best substances encoding mother-offspring similarity.

For this analysis we have 41 groups (mother-offspring pairs) and want to look at withing group similarities rather then between group dissimilarities. This was done in Primer-E, as the simper function from the vegan package computes discriminating compounds, rather then compounds that make a mother-pup pair unique (although both sets overlap of course).

```
# results from simper analysis in Primer-E
mp_simp <- read.csv(".\\files\\simper_mp_results.csv", colClasses = c("character", "numeric"))</pre>
mp_simp
#>
             comp contrib
#> 1 19.72268293
                   15.54
#> 2 15.45769231
                    12.25
#> 3 26.78859155
                    11.97
#> 4
     16.3974359
                    11.30
#> 5 19.52538462
                    10.87
#> 6
           21.405
                     8.49
#> 7 37.56363636
                     6.48
#> 8 15.62272727
                     6.48
#> 9 33.63655172
                     6.28
#> 10 30.80365385
                     6.03
#> 11
       20.361875
                     5.34
#> 12 17.40942623
                     4.79
```

Mother offspring similarity based on a Bray-curtis similarity matrix which was computed from just the subset of 12 top compounds from the SIMPER analysis is highly significant, both overall, as well as within colonies.

Full sample

# Within colony 1 (Special study beach)

#### Within colony 2 (Freshwater beach)

### Identification of best substances encoding colony dissimilarity.

Using simper from the vegan package to find the important substances for discriminating between the two colonies. And sorting them subsequently in order of contribution to colony dissimilarity.

```
# simper analysis
simp_colony <- vegan::simper(scent, factors$colony)</pre>
```

```
# getting 15 best substances and their contribution to colony dissimilarity
simp_colony_names <- rownames(summary(simp_colony, ordered = TRUE)[[1]])[1:15]</pre>
contribution <- summary(simp_colony, ordered = TRUE)[[1]]$contr[1:15]</pre>
# indices of colony substances (58,62,68,74,86,89,90,98,106,107,110,164,181,189,211)
ind_col <- paste(which(names(scent)%in%simp_colony_names), collapse = ",")</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 15.45769231 3.006785
#> 2 16.3974359 2.419310
#> 3 26.78859155 2.069715
#> 4 19.52538462 1.965096
#> 5
           21.405 1.894868
#> 6 21.34820513 1.671251
#> 7 19.72268293 1.669923
#> 8 30.80365385 1.482931
#> 9 38.5183871 1.440400
#> 10 17.40942623 1.330158
#> 11 20.51086207 1.288827
#> 12 33.63655172 1.266612
#> 13 21.57529412 1.208037
#> 14 15.74219178 1.180692
#> 15 19.66514286 1.128883
```

### Colony dissimilarity based on 15 compounds.

# Identification of substanced encoding relatedness.

All the following analyses are shown for the subset of mothers.

The core of the idea is to use a bootstrapping procedure on the BIO-ENV function, originally by Clarke (Clarke and Warwick 2001), which was modified (Taylor 2014) to work with a bray curtis similarity matrix. For details see the methods part of the paper. The function is built to run on parallel with snowfall (Knaus 2013) on a server or similar, but still takes a couple of days to finish.

Additional packages used are Hadley Wickham's dplyr (Wickham and Francois 2015) and stringr (Wickham 2015). First, the BIO-ENV function code is presented, followed by the bootstrap function.

```
bio.env <- function(fix.mat, var.mat,</pre>
                     fix.dist.method="bray", var.dist.method="euclidean",
                     scale.fix=FALSE, scale.var=TRUE,
                     output.best=10,
                     var.max=ncol(var.mat)
){
        \# \ if(dim(fix.mat)[1] \ != \ dim(var.mat)[1]) \{ stop("fixed \ and \ variable \ matrices \ must \ have \ the \ same \ n
        if(var.max > dim(var.mat)[2]){stop("var.max cannot be larger than the number of variables (colu
        require(vegan)
        combn.sum <- sum(factorial(ncol(var.mat))/(factorial(1:var.max)*factorial(ncol(var.mat)-1:var.m</pre>
        if(scale.fix){fix.mat<-scale(fix.mat)}else{fix.mat<-fix.mat}</pre>
        if(scale.var){var.mat<-scale(var.mat)}else{var.mat<-var.mat}</pre>
        # fix.dist <- vegdist(fix.mat, method=fix.dist.method)
        fix.dist <- fix.mat</pre>
        RES_TOT <- c()
        best.i.comb <- c()</pre>
        iter <- 0
        for(i in 1:var.max){
                 var.comb <- combn(1:ncol(var.mat), i, simplify=FALSE)</pre>
                 RES <- data.frame(var.incl=rep(NA, length(var.comb)), n.var=i, rho=0)
                 for(f in 1:length(var.comb)){
                          iter <- iter+1</pre>
                          var.dist <- vegdist(as.matrix(var.mat[,var.comb[[f]]]), method=var.dist.method)</pre>
                          temp <- suppressWarnings(cor.test(fix.dist, var.dist, method="spearman"))</pre>
                          RES$var.incl[f] <- paste(var.comb[[f]], collapse=",")</pre>
                          RES$rho[f] <- temp$estimate</pre>
                          if(iter %% 100 == 0){print(paste(round(iter/combn.sum*100, 3), "% finished"))}
                 }
                 order.rho <- order(RES$rho, decreasing=TRUE)</pre>
                 best.i.comb <- c(best.i.comb, RES$var.incl[order.rho[1]])</pre>
                 if(length(order.rho) > output.best){
                          RES_TOT <- rbind(RES_TOT, RES[order.rho[1:output.best],])</pre>
                 } else {
                          RES_TOT <- rbind(RES_TOT, RES)</pre>
                 }
        }
        rownames(RES_TOT) <- NULL
        if(dim(RES_TOT)[1] > output.best){
                 order.by.best <- order(RES_TOT$rho, decreasing=TRUE)[1:output.best]
        } else {
                 order.by.best <- order(RES_TOT$rho, decreasing=TRUE)</pre>
        OBB <- RES_TOT[order.by.best,]
        rownames(OBB) <- NULL</pre>
        order.by.i.comb <- match(best.i.comb, RES_TOT$var.incl)</pre>
```

```
OBC <- RES_TOT[order.by.i.comb,]
rownames(OBC) <- NULL

out <- list(
          order.by.best=OBB,
          order.by.i.comb=OBC,
          best.model.vars=paste(colnames(var.mat)[as.numeric(unlist(strsplit(OBB$var.incl[1], "," best.model.rho=OBB$rho[1]
)
    out
}</pre>
```

```
#### run seperately on multicore server #####
#### aim: resampling test for finding the substances associated with genetic
#### relatedness. Basic assumption: Each variable will be tested in many different
#### environments (individuals, other variables), which will prevent spurious
#### correlations, as the really important substances will occur in best subsets
#### in many different constellations. (see methods section)
# parallel computing using 40 cores, takes some days nevertheless and is just
# shown here.
library(vegan)
library(stringr)
library(dplyr)
library(snow)
library(snowfall)
# number of cores
ncores <- 2
# subset
scent_mum <- filter(scent, factors$age == 1)</pre>
relate_mum <- relatedness[factors$age == 1, factors$age == 1]</pre>
# initialise results vector
all best <- vector()
# initialise cluster
snowfall::sfInit(parallel=TRUE, cpus=ncores, type="SOCK")
# export libraries and main function to all cores
snowfall::sfSource("bio.env.R")
snowfall::sfLibrary(vegan)
snowfall::sfLibrary(stringr)
snowfall::sfLibrary(dplyr)
bootstrap <- function(iter_comp) { # main resampling function</pre>
       for (i in 1:500) {
               # sample 20 out of 41 mothers, indices
               ind_obs <- sort(sample(1:41, size = 20, replace = F))</pre>
               # subset relate_mum and scent_mum
               reltemp <- 1-as.dist(relate_mum[ind_obs, ind_obs])</pre>
```

```
abundtemp <- scent_mum[ind_obs, ]</pre>
                for (i in iter_comp) {
                        # sample 10 compounds
                        index_comps <- sort(sample(1:213, size = 10, replace = F))</pre>
                        abundtemp_sub <- abundtemp[, index_comps]</pre>
                        # get vector with O for null-column and 1 for non-null column
                       nullcomps <- apply(abundtemp_sub, 2, function(x) sum(x>0))
                        abundtemp sub <- subset(abundtemp sub,
                                                subset = c(rep(TRUE, nrow(abundtemp_sub))),
                                                select = (nullcomps >= 2))
                        # new iteration if too less substances left
                       if (ncol(abundtemp_sub) <= 2) next</pre>
                        # main function: bio.env finds subset that mostly correlates
                        # with relatedness
                       results <- bio.env(reltemp, abundtemp_sub,
                                          var.dist.method = "bray",
                                           scale.fix = F, scale.var = F)
                       mods <- results$best.model.vars</pre>
                       best <- unlist(str_split(mods, ","))</pre>
                        all_best <- append(all_best, best)</pre>
                        # write(best, file = "best.txt", append = TRUE, sep = " ")
                }
       return(all_best)
# export objects
snowfall::sfExportAll(except = NULL, debug = FALSE)
snowfall::sfClusterEval(ls())
# create list of 500 iterations for all cores
vals <- list()</pre>
for (i in 1:ncores) {
       vals[[i]] <- 1:500
}
# run analysis
# best is a list of all best subsets
best <- snowfall::sfLapply(vals, bootstrap)</pre>
# stop cluster
sfStop()
# bring all results
results <- unlist(best)
```

### Analysing results from the BIO-ENV bootstrap analysis.

best\_mums is a data frame containing the number of occurences of each variable in the best subset from the BIO-ENV bootstrap analysis. Substances, that were retained more often are therefore likely to be genuinly associated with genetic relatedness.

```
# substance occurences are sorted in the table
best_mums <- read.csv("files/bootstrap_mums.csv",row.names=1)</pre>
```

To analyse how many of these compounds are really important, the idea is to take an increasing number of "best" compounds and compute a mantel test with relatedness for each of the subsets. The subsequent plot

shows a nice peak, which could be seen as the optimal number of chemicals encoding relatedness.

```
# subset mothers
scent_mum <- dplyr::filter(scent, factors$age == 1)
relate_mum <- 1-relatedness[factors$age == 1, factors$age == 1]
sub_names_mums <- row.names(best_mums)

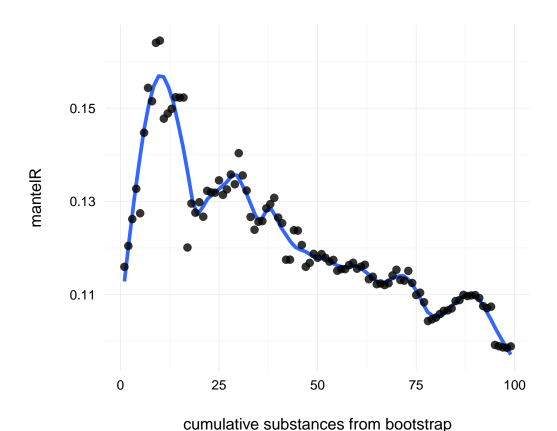
statm <- vector()
sigm <- vector()

# compute mantelR for an increasing set of best substances
for (i in 2:100) {
    bc_dist <- vegan::vegdist(scent_mum[, sub_names_mums[1:i]], method = "bray")
    mod <- vegan::mantel(relate_mum, bc_dist, na.rm = T, method = "spearman")
    statm <- append(statm, mod$statistic)
    sigm <- append(sigm, mod$sig)
}

stat_df <- data.frame(num_comps = 1:length(statm), mantelR = statm)</pre>
```

Plotting mantelR for an increasing number of best substances.

```
library(grid)
# simple plot
ggplot2::ggplot(stat_df, aes(x = num_comps, y = mantelR)) +
        stat_smooth(se = FALSE, span = 0.16, size = 1.3, method = "loess") +
        geom_point(colour = "black", size = 3, alpha = 0.8) +
        theme minimal(base size = 12) +
        theme(strip.text.x = element_text(vjust=1, size = 12),
              axis.title.x = element_text(vjust= -2 ,size =12),
              axis.title.y = element_text(vjust=3,size = 12),
              axis.ticks.x = element_blank(),
              axis.ticks.y = element_blank(),
              plot.margin = (unit(c(.5, .5, 2, 2), "cm"))) +
        \# scale_x_continuous(breaks=c(seq(from = 0.8, to = 1.20, by = 0.1))) +
        \#geom\_text(aes(0.85,80, label="(a) r = 0.34, p = 0.027"), size=4) +
        xlab("cumulative substances from bootstrap") +
        ylab("mantelR")
```



The plot peaks at 10 substances. We now want to do a single mantel test for chemical bray-curtis similarities based on these 10 compounds and genetic relatedness. As already shown in the plot, the mantelR is 0.164 and is highly significant.

```
# indices of the 10 best compounds associated with relatedness -
comp_ind_m \leftarrow c(36,52,86,88,96,103,110,203,206,207)
# bray curtis similarity matrix based on this 10 compounds
scent_bc <- 1-(as.matrix(vegan::vegdist(as.matrix(scent[factors$age == 1, comp_ind_m])),</pre>
                      method = "bray"))
# relatedness matrix
rel_m <- relatedness[1:41, 1:41]
# mantel test for association between both
vegan::mantel(rel_m, scent_bc, method = "spearman", permutation = 1000, na.rm = TRUE)
#> Mantel statistic based on Spearman's rank correlation rho
#>
#> Call:
#> veqan::mantel(xdis = rel_m, ydis = scent_bc, method = "spearman",
                                                                            permutations = 1000, na.rm =
#> Mantel statistic r: 0.164
         Significance: 0.002997
```

```
#>
#> Upper quantiles of permutations (null model):
#> 90% 95% 97.5% 99%
#> 0.0706 0.0911 0.1045 0.1260
#> Permutation: free
#> Number of permutations: 1000
```

R version and platform.

```
sessionInfo()
#> R version 3.2.1 (2015-06-18)
#> Platform: x86 64-w64-mingw32/x64 (64-bit)
#> Running under: Windows 8 x64 (build 9200)
#> locale:
#> [1] LC_COLLATE=English_United Kingdom.1252
#> [2] LC_CTYPE=English_United Kingdom.1252
#> [3] LC_MONETARY=English_United Kingdom.1252
#> [4] LC_NUMERIC=C
#> [5] LC_TIME=English_United Kingdom.1252
#> attached base packages:
#> [1] qrid
               stats qraphics qrDevices utils datasets methods
#> [8] base
#>
#> other attached packages:
#> [1] reshape2_1.4.1 magrittr_1.5 dplyr_0.4.2
                                                          HDMD_1.2
#> [5] psych_1.5.4 ggplot2_1.0.1 minmodelr_0.1 inbreedR_0.1
#> [9] yegan 2 3-0 nermyte 0 8-4 MASS 7 3-40 lattice 0 20
                        permute 0.8-4 MASS 7.3-40
#> [9] vegan 2.3-0
                                                          lattice 0.20-31
#> [13] devtools_1.8.0
#>
#> loaded via a namespace (and not attached):
\#> [1] Rcpp_0.11.6 formatR_1.2 git2r_0.10.1 \#> [5] tools_3.2.1 digest_0.6.8 evaluate_0.7
                                                          plyr_1.8.3
                                                          memoise\_0.2.1
#> [9] nlme_3.1-120 gtable_0.1.2 mgcv_1.8-6
#> [13] DBI 0.3.1 curl 0.9 yaml 2.1.13
                                                          Matrix 1.2-1
                                           yaml_2.1.13
#> [13] DBI_0.3.1
                         curl_0.9
                                                             parallel_3.2.1
#> [17] proto_0.3-10 stringr_1.0.0
                                           xml2_0.1.1
                                                             cluster_2.0.1
\# [21] knitr_1.10.5 rversions_1.0.1 ecodist_1.2.9 R6_2.0.1
\# [25] rmarkdown_0.7 codetools_0.2-11 scales_0.2.5 htmltools_0.2.6
#> [29] assertthat_0.1 mnormt_1.5-3
                                           colorspace_1.2-6 labeling_0.3
#> [33] stringi_0.5-2
                       lazyeval_0.1.10 munsell_0.4.2
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