### Purees data

2024-01-13

#### Load libraries

```
shhh = function(lib_name){ # It's a library, so shhh!
  suppressWarnings(suppressMessages(require(lib_name, character.only = TRUE)))
}
shhh("tidyverse")
shhh("ACutils")
shhh("mvtnorm")
shhh("salso")
shhh("FGM")
shhh("gmp")
shhh("mcclust")
shhh("mcclust.ext")
shhh("logr")
shhh("tidygraph")
shhh("ggraph")
shhh("igraph")
shhh("Rcpp")
shhh("RcppArmadillo")
shhh("RcppEigen")
## Load custom functions
source("functions/utility_functions.R");
source("functions/bulky_functions.R");
source("functions/data_generation.R")
sourceCpp("functions/wade.cpp")
Rcpp::sourceCpp('functions/UpdateParamsGSL.cpp')
library('RcppGSL')
library(fda)
library(tidyverse)
library(coda)
library(lattice)
```

#### Goal

Stima della partizione e grafo con bulky functions 2024.

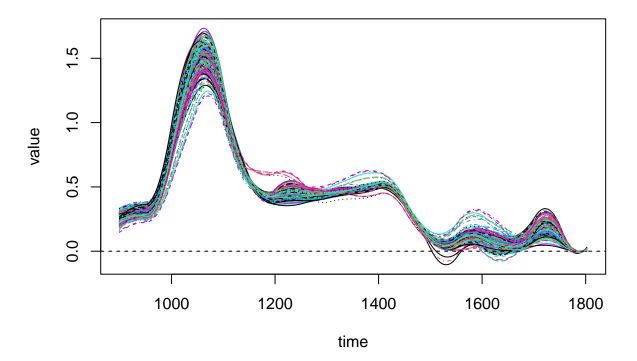
#### Purees data

```
load("purees.Rdat")
curves = purees$data
wavelengths = purees$wavelengths
strawberry <- curves[which(curves$Group == "Strawberry"), ]
data = strawberry[,-1]*100
data = as.matrix(data)  # n x r

#Generate basis
p = 20
n = dim(data)[1]
r = dim(data)[2]
range_x = range(wavelengths)

data_W <- t(as.matrix(data))
basis <- create.bspline.basis(rangeval=range(wavelengths), nbasis=p, norder=3)
data.fd <- Data2fd(y = data_W, argvals = wavelengths, basisobj = basis)
plot.fd(data.fd, main="B-splines")</pre>
```

### **B**-splines



## [1] "done"

#### Initialization

```
# Compute quantities for function UpdateParamGSL
tbase_base = t(BaseMat)%*%BaseMat # p x p (phi_t * phi)
tbase_data = t(BaseMat)%*%t(data) # p x n (phi_t * Y_t)
Sdata = sum(diag(data%*%t(data)))
set_UpdateParamsGSL_list = set_UpdateParamsGSL(
   tbase_base = tbase_base,
   tbase_data = tbase_data,
   Sdata = Sdata,
   a_tau_eps = 2000,
   b_{tau_eps} = 2,
   sigma mu = 100,
             = r,
   Update_Beta = TRUE,
   Update_Mu = TRUE,
   Update_Tau = TRUE
)
# Set the number of iterations and burn-in
niter <- 50000
burn_in <- 1000
# Set the initialization values of the chains
initialization_values = set_initialization(
   Beta = matrix(rnorm(n=p*n), nrow = p, ncol = n),
   mu
               = rnorm(n=p),
   tau_eps = 100,
K = matrix(0,p,p),
                = matrix(0,p,p),
                = rep(1,p),
   z
   rho
                = p,
             = 1,
= 1,
   a_sigma
   b_sigma
   c_sigma
               = 1,
   d_sigma
               = 1,
              = 100,
   c_{theta}
               = 10,
   d_theta
   sigma
               = 0.5,
   theta
              = 1,
   weights_a0 = rep(1,p-1),
weights_d0 = rep(1,p-1),
   total_weights = 0,
   total_K = matrix(0,p,p),
   total_graphs = matrix(0,p,p),
   graph_start = NULL,
   graph density = 0.3,
   beta_sig2 = 0.2,
```

```
d = 3
```

#### Gibbs sampler

```
chains = Gibbs_sampler_update(
 set_UpdateParamsGSL_list,
 niter,
 initialization_values,
 alpha_target
                      = 0.234,
                     = 0.5,
 alpha_add
 adaptation_step
                   = 1/(p*1000),
                      = 22111996,
 seed
 update_sigma_prior = TRUE,
 update_theta_prior = TRUE,
 update_graph

perform
                   = TRUE
 perform_shuffle
```

#### Useful plots: 1. Plot smoothed curves

```
# Compute the mean of Beta in order to have data_post
sum_Beta <- matrix(0, p, n)
for(i in (burn_in+1):niter){
    sum_Beta <- sum_Beta + chains$Beta[[i]]
}
mean_Beta <- sum_Beta/(niter-burn_in)
data_post <- BaseMat %*% mean_Beta

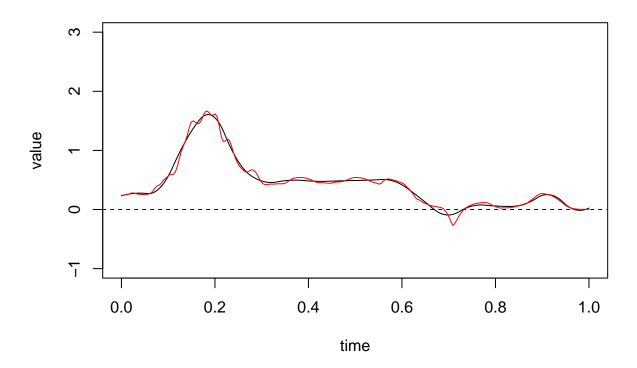
# Compute the x value, create the basis and the functional object
x <- seq(0, 1, length.out=r)
basis <- create.bspline.basis(rangeval=range(x), nbasis=p, norder=3)
data.fd <- Data2fd(y = data_post, argvals = x, basisobj = basis)

# Plot smoothed curves
plot.fd(data.fd[1,], main="smoothed curves", ylim=c(-1,3))

## [1] "done"

# plot(x, data_post[,1], type='l', ylim=c(-2,4))
lines(x,data[1,], main="smoothed curves", col='red')</pre>
```

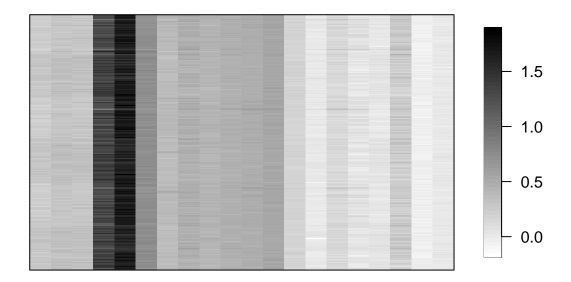
### smoothed curves



## Useful plots: 2. Plot of the final Beta matrix

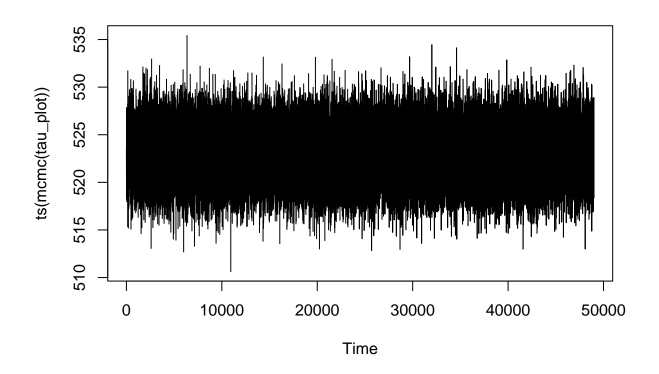
```
ACutils::ACheatmap(
  chains$Beta[[niter]],
  use_x11_device = F,
  horizontal = F,
  main = "Estimated Beta matrix",
  center_value = NULL,
  col.upper = "black",
  col.center = "grey50",
  col.lower = "white"
)
```

### **Estimated Beta matrix**

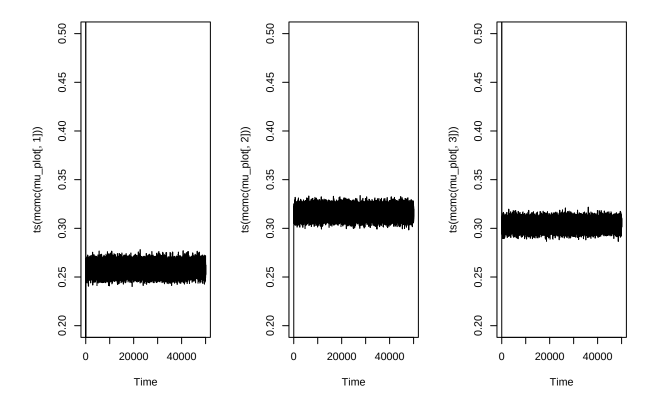


# Useful plots: 2. Traceplots (tau\_eps, mu, beta)

```
# tau_eps
tau_plot <- as.vector(chains$tau_eps)
tau_plot <- tau_plot[(burn_in+1):niter]
plot(ts(mcmc(tau_plot)))</pre>
```

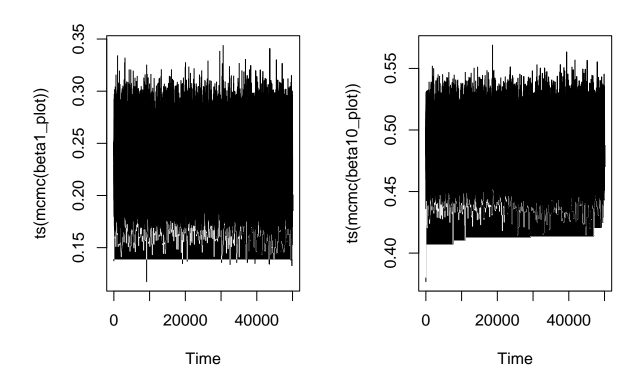


```
# mu
mu_plot <- matrix(0, niter, p)
for(i in 1:niter){
    mu_plot[i, ] <- chains$mu[[i]]
}
par(mfrow=c(1,3))
plot(ts(mcmc(mu_plot[, 1])), ylim=c(0.2,0.5))
plot(ts(mcmc(mu_plot[, 2])), ylim=c(0.2,0.5))
plot(ts(mcmc(mu_plot[, 3])), ylim=c(0.2,0.5))</pre>
```



```
# first element of first beta
beta1_plot <- rep(0, niter)
for(i in 1:niter){
    beta1_plot[i] <- chains$Beta[[i]][1,1]
}
# and 10th element of first beta
beta10_plot <- rep(0, niter)
for(i in 1:niter){
    beta10_plot[i] <- chains$Beta[[i]][10,1]
}

par(mfrow=c(1,2))
plot(ts(mcmc(beta1_plot)))</pre>
```

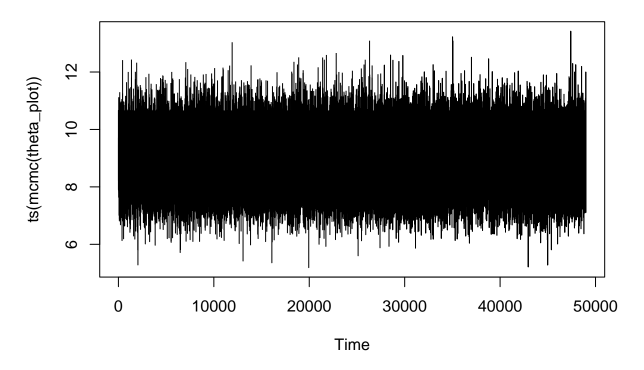


## Posterior analysis

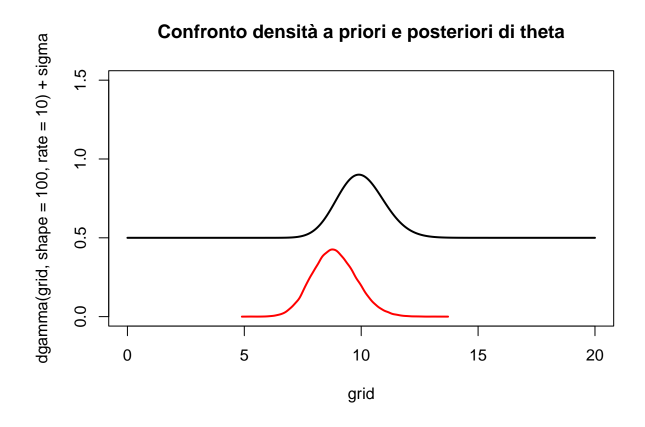
```
#prior and posterior of theta

#traceplot theta
theta_plot <- as.vector(chains$theta)
theta_plot <- theta_plot[(burn_in+1):niter]
plot(ts(mcmc(theta_plot)), main='Trace plot theta')</pre>
```

## Trace plot theta



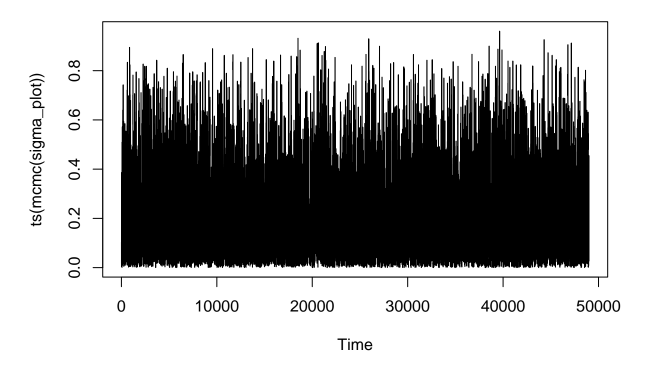
```
# confronto densità a priori e posteriori
min_val <- 0
max_val <- 20
sigma <- 0.5
grid <- seq(min_val, max_val, length.out=10000)
plot(grid, dgamma(grid,shape=100, rate=10)+sigma, type='l', lwd=2, ylim=c(0,1.5), main='Confronto densi
lines(density(mcmc(theta_plot))$x, density(mcmc(theta_plot))$y, type='l', lwd=2, col='red') #posterior</pre>
```



```
#prior and posterior of sigma

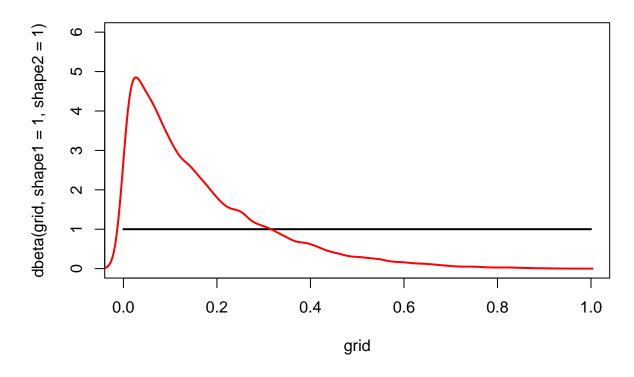
#traceplot sigma
sigma_plot <- as.vector(chains$sigma)
sigma_plot <- sigma_plot[(burn_in+1):niter]
plot(ts(mcmc(sigma_plot)), main='Trace plot sigma')</pre>
```

## Trace plot sigma



```
# confronto densità a priori e posteriori
min_val <- 0
max_val <- 1
grid <- seq(min_val, max_val, length.out=10000)
plot(grid, dbeta(grid,shape1=1, shape2=1), type='l', lwd=2, ylim=c(0,6), main='Confronto densità a prior
lines(density(mcmc(sigma_plot))$x, density(mcmc(sigma_plot))$y, type='l', lwd=2, col='red') #posterior</pre>
```

### Confronto densità a priori e posteriori di sigma



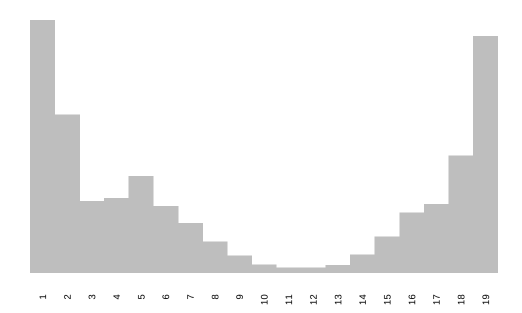
## Recomputing the partition in other forms and the number of groups

cex.names=.6,

```
rho <- chains$rho
r = do.call(rbind, lapply(chains$rho, rho_to_r))
## Warning in (function (..., deparse.level = 1) : number of columns of result is
## not a multiple of vector length (arg 2)
z = do.call(rbind, lapply(chains$rho, rho_to_z))
num_clusters = do.call(rbind, lapply(chains$rho, length))
num_clusters = as.vector(num_clusters)
### Barplot of changepoints
bar_heights = colSums(r)
barplot(
  bar_heights[1:(p-1)],
  names = seq_along(bar_heights[1:(p-1)]),
  border = "NA",
  space = 0,
  yaxt = "n",
  main="Changepoint frequency distribution",
  \#col = color,
```

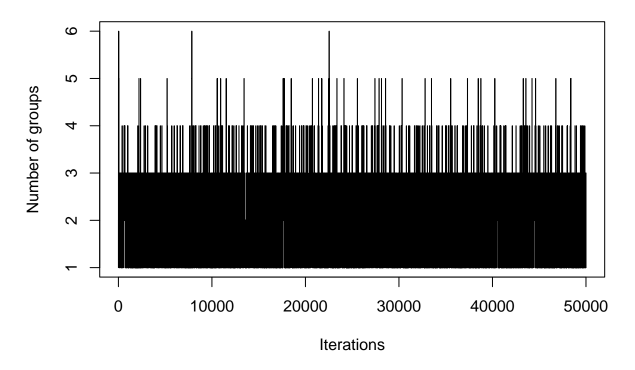
```
las=2
)
```

# **Changepoint frequency distribution**



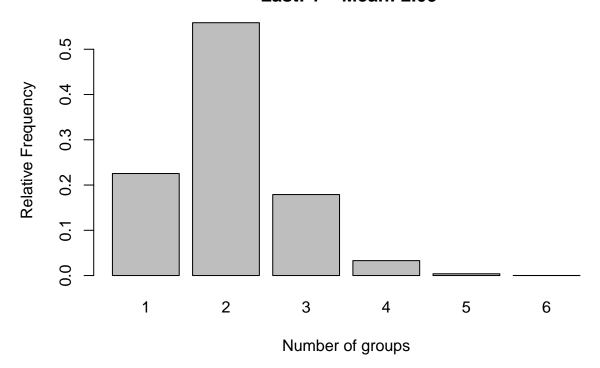
```
### Evolution of the number of clusters
plot(
    x = seq_along(num_clusters),
    y = num_clusters,
    type = "n",
    xlab = "Iterations",
    ylab = "Number of groups",
    main = "Number of groups - Traceplot"
)
lines(x = seq_along(num_clusters), y = num_clusters)
```

# **Number of groups – Traceplot**



```
barplot(
  prop.table(table(num_clusters)),
  xlab = "Number of groups",
  ylab = "Relative Frequency",
  main = paste(
    "Number of groups - Relative Frequency\n",
    "Last:",
    tail(num_clusters, n = 1),
    "- Mean:",
    round(mean(num_clusters), 2)
  )
)
```

### Number of groups – Relative Frequency Last: 1 – Mean: 2.03



```
### Retrieving best partition using VI on visited ones (order is guaranteed here)
# compute VI
sim_matrix <- salso::psm(z)</pre>
dists <- VI_LB(z, psm_mat = sim_matrix)</pre>
# select best partition (among the visited ones)
best_partition_index = which.min(dists)
rho_est = rho[[best_partition_index]]
z_est = z[best_partition_index,]
# VI loss
dists[best_partition_index]
## [1] 0.3413531
# select best partition
unname(z_est)
   ## Graph
# Extract last plinks
last_plinks = tail(chains$G, n=1)[[1]]
```

```
# Criterion 1 to select the threshold (should not work very well) and assign final graph
threshold = 0.5
G_est <- matrix(0,p,p)
G_est[which(last_plinks>threshold)] = 1

#Criterion 2 to select the threshold
bfdr_select = BFDR_selection(last_plinks, tol = seq(0.1, 1, by = 0.001))

# Inspect the threshold and assign final graph
bfdr_select$best_treshold
```

## [1] 0.859

```
G_est = bfdr_select$best_truncated_graph

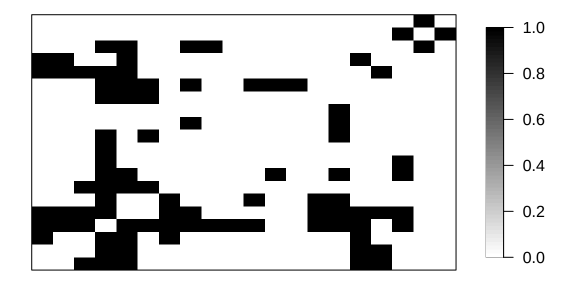
### Plot estimated matrices
ACutils::ACheatmap(
    last_plinks,
    use_x11_device = F,
    horizontal = F,
    main = "Estimated plinks matrix",
    center_value = NULL,
    col.upper = "black",
    col.center = "grey50",
    col.lower = "white"
)
```

# **Estimated plinks matrix**



```
ACutils::ACheatmap(
   G_est,
   use_x11_device = F,
   horizontal = F,
   main = "Estimated Graph",
   center_value = NULL,
   col.upper = "black",
   col.center = "grey50",
   col.lower = "white"
)
```

## **Estimated Graph**



```
ACutils::ACheatmap(
  tail(chains$K,n=1)[[1]],
  use_x11_device = F,
  horizontal = F,
  main = "Estimated Precision matrix",
  center_value = NULL,
  col.upper = "black",
  col.center = "grey50",
  col.lower = "white"
)
```

## **Estimated Precision matrix**

