Adaptive Sampling simulations

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Motivation for adaptive sampling

In many scenarios we would like to be able to adaptively increase sampling effort when certain observed values are of interest. This would be especially beneficial in the case where we are try to observe rare events that are likely to be clustered together. For example a rare plant species that has very particular growing conditions. This would be the case where it may be very rare to see the plant but if we see one we see 100 and it is desired to be able to sample all of the plants in the cluster.

Simulations

First I'll write a few functions that I think will be helpful in making this adaptive sampling demonstration. I want to recreate the example in Adaptive cluster Sampling (1990) where there are points distributed in clusters in an area. The area is split into a grid, then grid cells are chosen as the primary sampling unit.

Goals:

- choose a number of clusters
 - easy enough, this can just be rpois(1,n), this will choose a number of clusters randomly based on some mean. We could use any discrete distribution to decide this
- generate points in those clusters
 - decide how many points are in each cluster, this it is probably actually most appropriate to use a poisson distribution since it will be a count.
- at this step I also need to choose the locations of the centers, runif(ncenters, 0, 20)
- I can then generate points normally distributed around these centers using rnorm(ncenters*2,mu,sd)
- choose a sample of initial grid cells
 - sample(1:ncells,n)
- expand if the cell next to it is occupied
 - need a list of currently occupied cells and a way to determine adjacent cells. maybe just a vector called occupied
 - function takes vector of neighbours, checks against vector of occupied, returns neighbors in occupied

Creating functions for sampling

Loading required package: ggridges

```
require(tidyverse)
## Loading required package: tidyverse
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
             1.1.4
                       v readr
                                   2.1.5
## v forcats
              1.0.0
                        v stringr
                                   1.5.1
## v ggplot2
              3.5.0
                        v tibble
                                   3.2.1
## v lubridate 1.9.3
                        v tidyr
                                   1.3.1
## v purrr
              1.0.2
                                          ## -- Conflicts -----
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
# install.packages("qqridqes")
require(ggridges)
```

```
# install.packages("colorspace")
require(colorspace)
```

Loading required package: colorspace

1 7.951749 12.67416 Cluster 1 ## 2 6.299387 15.44295 Cluster 1 ## 3 5.416986 13.04660 Cluster 1

make clusters

```
make clusters<-function(grid size=20, nclusters=3, avg size=20, force inclusion=FALSE){
  # create a list of clusters.
  centers<-split(runif(nclusters*2,0,grid_size),seq(nclusters))</pre>
  # get the three cluster sizes
  sizes<-rpois(3,avg_size)</pre>
  # get list of matrices of the centers
  center_dfs<-map2(sizes,centers,</pre>
                   ~ kronecker(matrix(rep(1,.x),ncol=1), matrix(.y,ncol=2)))
                               # I actually used the kronecker product holy crap!
  if(force_inclusion==T){
    # get coordinates of locations
    locations <- sizes %>% # use sizes
      map( ~data.frame(matrix(rnorm(.x*2),ncol=2))) %% # make list of df of changes from centers
      map2(.y=center_dfs, ~.x+.y) %>% # add the changes on to the center
      map2(.y=seq(nclusters), ~ mutate(.x, Group=paste("Cluster",.y)))%>%
      bind rows()
    colnames(locations)<-c("X","Y","Group")</pre>
    # force points to snap inside if they were generated outside. This means some
    # tiles will be more densely clustered at edges when a center is near an edge
    # should help maintain the average number of points though
    locations<-locations%>%
      mutate(X=ifelse(X>grid_size,grid_size,X))%>%
      mutate(X=ifelse(X<0, 0, X))%>%
      mutate(Y=ifelse(Y>grid_size,grid_size,Y))%>%
      mutate(Y=ifelse(Y<0, 0, Y))</pre>
  }else{
    # get coordinates of locations
    locations<-sizes %>% # use sizes
      map(~data.frame(matrix(rnorm(.x*2),ncol=2))) %>% # make list of df of changes from centers
      map2(.y=center_dfs,~.x+.y) %>% # add the changes on to the center
      map2(.y=seq(nclusters), ~ mutate(.x, Group=paste("Cluster",.y)))%>%
      bind rows()
    colnames(locations)<-c("X","Y","Group")</pre>
  }
  locations
}
head(make_clusters(force_inclusion = TRUE))
##
                           Group
```

```
## 4 8.794119 14.05494 Cluster 1
## 5 6.514118 12.90943 Cluster 1
## 6 8.152712 13.70114 Cluster 1
```

plotting

```
plot_clusters<-function(cluster_df, grid_size=20,samp=NULL){</pre>
  # cluster_df is the set of points on the grid
  # samp is the set of tiles that were sampled
  # create a plot of the clusters
 p<-ggplot(cluster_df, aes(x=X, y=Y, color=Group))+</pre>
    geom_point()+
    scale_x_continuous(breaks=seq(grid_size))+
    scale_y_continuous(breaks=seq(grid_size))+
    coord_cartesian(xlim=c(0,grid_size), ylim=c(0,grid_size))
  if(!is.null(samp)){
    # this will highlight tiles that are sampled
    # samp is the dataframe containing the coordinates for tiles
    p<-p+geom_rect(data=samp,
                   aes( xmin=X-1, xmax=X, ymin=Y-1, ymax=Y),
                   color="black",
                   fill=NA)
 }
 plot(p)
```

Check whether a tile is occupied

```
is_occupied<-function(tile=c(1,1,1),df){
  # check if a tile is occupied tile location is given as c(row,column)
  # give a dataframe with coordinates of point
  # check if there are any points in xrange & yrange at the same time
  #sum((df[,1]<tile[2] & df[,1]>tile[2]-1) * (df[,1]<tile[2] & df[,1]>tile[2]-1))>0
  sum((df[,1]<tile[1]&df[,1]>tile[1]-1) & (df[,2]<tile[2] & df[,2]>tile[2]-1))>0
}
```

find the average in a single tile this will get used in the hansen hurwitz estimator as well. Just including it here so we can have y_k in the simulated data.

```
tile_sum<-function(tile=c(X=1,Y=1,K=1), df){
    # This function returns the y_k for tile k
    # tile is the tile to find the response of
    # df is the dataframe of points on the grid
    sum((df[,1]<tile[1]&df[,1]>tile[1]-1) & (df[,2]<tile[2] & df[,2]>tile[2]-1))
}
```

get the neighbours of a tile

choose tiles for sample

```
get_tiles<-function(grid_size=20,n1=10,...){
    # this function returns a sample of n1 tiles using a square grid_size grid
    # get a tile
    tiles<-sample(1:grid_size^2,n1,...)
    # convert tile numbers into X and Y
    samp<-data.frame(X=tiles%,grid_size, Y=(tiles-1)%,/%grid_size+1)
    samp$X[samp$X==0]<-grid_size
    samp$k<-1:n1
    samp
}</pre>
```

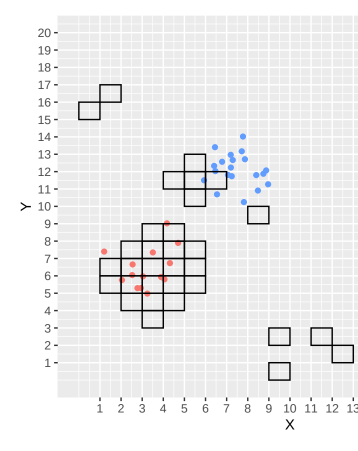
Performing adaptive cluster sampling

One sample from a population of points

```
simulate_one<-function(nclusters=3, grid_size=20,n1=10,force_inclusion=FALSE, hard_border=TRUE,...){
    # generate clusters
    points<-make_clusters(force_inclusion = force_inclusion)
    # choose the starting grid cells
    sample_tiles<-get_tiles(n1=10,...)
    # save a copy to check against for updates
    temp<-sample_tiles
    # check whether or not they are occupied based on the clusters
    occupied<-apply(sample_tiles,1,function(x) is_occupied(x, points))
    # find the neighbours of the occupied points
    neighbours<-apply(sample_tiles[occupied,], 1, function(x) get_neighbours(x,hard_border = TRUE))%>% #
    bind_rows() %>% # turn list of neighbours into tibble
    as.data.frame() # into dataframe
    # update sample tiles to include neighbours
    sample_tiles<-rbind(sample_tiles,neighbours) %>%
```

```
unique()
  # keep looping until sample_tiles does not grow
  while(dim(temp)[1]!=dim(sample_tiles)[1]){
    # save a copy to check against for updates
    temp<-sample_tiles
    # check whether or not they are occupied based on the clusters
    occupied<-apply(sample_tiles,1,function(x) is_occupied(x,points))</pre>
    # find the neighbours of the occupied points
    neighbours<-apply(sample_tiles[occupied,], 1, function(x) get_neighbours(x,hard_border = TRUE))%>%
      bind_rows() %>% # turn list of neighbours into tibble
      as.data.frame() # into dataframe
    # update sample tiles to include neighbours
    sample_tiles<-sample_tiles %>%
      bind_rows(neighbours) %>%
      unique()
  }
  # get the values of the response for each unit in the sample
  sample_tiles$y_k<-apply(sample_tiles,1,function(x) tile_sum(x, points))</pre>
  # get the number of units in each network m_k and add occupied to sample_tiles
  sample_tiles<-sample_tiles%>%
    cbind(occupied)%>%
    group_by(k)%>%
    mutate(m k=sum(occupied))
  # minimum for m_k is 1 not 0, set all 0s to 1
  sample_tiles$m_k[sample_tiles$m_k==0]<-1</pre>
  return(list(sample_tiles=sample_tiles,
              points=points,
              nclusters=nclusters,
              grid_size=grid_size,
              n1=n1)
}
set.seed(pi)
sim_data<-simulate_one()</pre>
```

```
set.seed(pi)
sample_one<-simulate_one(n1=10,nclusters=3, grid_size=20,force_inclusion=TRUE)
plot_clusters(sample_one$points,samp=sample_one$sample_tiles)</pre>
```



Demonstrating the function works to generate a sample

Multiple samples from the same population of points

Goals: - take m samples from a population - return output in the same format as simulate_one but since there are m samples there will be a list of lists

```
# m numer of samples from the same population
simulate_m<-function(m=10,nclusters=3, grid_size=20,n1=10,force_inclusion=FALSE, hard_border=TRUE,...
  results<-list()
  # generate clusters
  points<-make_clusters(force_inclusion = force_inclusion)</pre>
  for(i in 1:m){ # loop for the m samples
    # this is just the code from simulate_one() see that for more comments
    sample_tiles<-get_tiles(n1=10,...)</pre>
    temp<-sample_tiles</pre>
    occupied<-apply(sample_tiles,1,function(x) is_occupied(x,points))</pre>
    neighbours<-apply(sample_tiles[occupied,], 1, function(x) get_neighbours(x,hard_border = hard_border</pre>
      bind_rows() %>%
      as.data.frame()
    sample_tiles<-rbind(sample_tiles,neighbours) %>%
      unique()
     # keep looping until sample_tiles does not grow
    while(dim(temp)[1]!=dim(sample_tiles)[1]){
      # save a copy to check against for updates
```

```
temp<-sample_tiles
      # check whether or not they are occupied based on the clusters
      occupied<-apply(sample_tiles,1,function(x) is_occupied(x,points))</pre>
      # find the neighbours of the occupied points
      neighbours <- apply (sample_tiles [occupied,], 1, function(x)get_neighbours(x, hard_border = hard_bord
        bind_rows() %>% # turn list of neighbours into tibble
        as.data.frame() # into dataframe
      # update sample tiles to include neighbours
      sample_tiles<-sample_tiles %>%
        bind_rows(neighbours) %>%
        unique()
    }
    # get the values of the response for each unit in the sample
    sample_tiles$y_k<-apply(sample_tiles,1,function(x) tile_sum(x, points))</pre>
    # get the number of units in each network m_k and add occupied to sample_tiles
    sample_tiles<-sample_tiles%>%
      cbind(occupied)%>%
      group_by(k)%>%
      mutate(m_k=sum(occupied))
    # minimum for m k is 1 not 0, set all 0s to 1
    sample_tiles$m_k[sample_tiles$m_k==0]<-1</pre>
    results[[i]]<-list(sample_tiles=sample_tiles,</pre>
                                   points=points,
                                   nclusters=nclusters,
                                   grid_size=grid_size,
                                   n1=n1)
 }
  return(results)
set.seed(pi)
sim_data_m<-simulate_m()</pre>
```

When trying to write this simulation the largest issue I ran in to was determining how to treat points that lie outside of the grid. Should they be included in the sample? should they be ignored? should I change the generating mechanism to force points to be bounded by the edges? There are lots of things you need to specify in regards to how the points are generated. This seems like it could also be an issue that comes up in practical situations as well, for example the case where you have a defined area you are allowed to collect samples from but the thing you are measuring can occur up to and outside of that area. It seems like we would be underestimating the average number in the greater population if those are excluded from the cluster since it means we are underestimating the cluster size.

Finding the HT and HH Estimators

Vocabulary

- neighborhood: the collection of units that are immediately included in the sample if a given unit is included. This relationship is symmetric and is typically (but not necessarily) geographic.
- Cluster: the collection of all the units that are observed under the design as a result of initial selection of unit i

- Network: selection of any unit within the network would lead to the inclusion in the sample of every other unit in the network.
- edge unit: any unit not satisfying the condition but in the neighborhood of one that does

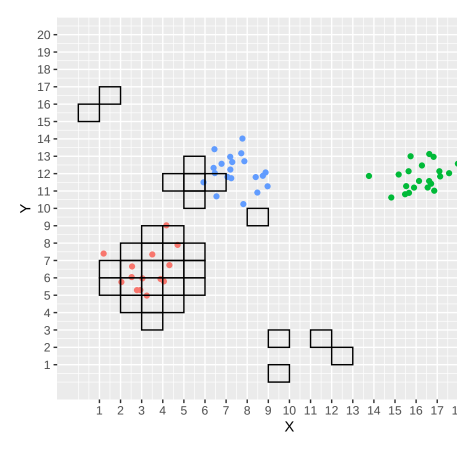
Hansen-Hurwitz

- Let Ψ_k be the network that includes unit k and m_k be the number of units in that network. Any unit not satisfying the criterion is size 1.
- Let $\bar{y}_k^* = (m_k)^{-1} \sum_{j \in \Psi_k} y_j$ represent the average number of observations in the network that includes the kth unit of the initial sample
- The modified Hansen-Hurwitz estimators given as $t_{HH^*} = n_1^{-1} \sum_{k=1}^{n_1} \bar{y}_k^*$

Steps

- Find average number of response in each network
 - file the average number of the response in a single tile.
- compute estimator using the averages

```
modified_HH<-function(sim_data,plot=F,n1=10,...){</pre>
  # Find the hansen hurwitz estimator of a sample called sim data
  if(plot==T){ # plot sample
    plot_clusters(sim_data$points,samp=sim_data$sample_tiles)
  }
  # sample_one$points and sample_one$sample_tiles
  # get the response values for each tile sampled
  y_k<-sim_data$sample_tiles$y_k</pre>
  # find the sum of each network that a unit belongs to this returns
  # k and the mean
  temp<-data.frame(y_k,group=sim_data$sample_tiles$k) %>%
    group_by(group)%>%
    filter(y_k>0) %>%
    summarize(network means=mean(y k))
  means<-data.frame(init_sample=1:n1,network_means=rep(0,n1))</pre>
  means[temp$group,2]<-temp$network_means</pre>
  mean (means$network_means)
}
modified_HH(sim_data,plot=T)
```



Applying the function to a single sample

[1] 0.25

Hovritz-Thompson

The classic Horvitz-Thompson estimator is given by dividing each y-value by the associated inclusion probability and is an unbiased estimator of the population mean. This is not viable in adaptive cluster sampling as the inclusion probabilities for all units are not known. None-the-less we can still create an unbiased estimator by modifying the Horvitz-Thompson estimator. First we define

$$\alpha_k^* = 1 - \binom{N - m_k}{n_1} / \binom{N}{n_1}$$

Where m_k is the number of units in the network that includes unit k, N is the number of units in the population, and n_1 is the number of units in the initial sample. Next let

$$J_k = \begin{cases} 0 & \text{If the condition is not satisfied} \\ 1 & \text{Otherwise} \end{cases}$$

Then the modified estimator is given by

$$t_{HT^*} = N^{-1} \sum_{k=1}^{v} y_k J_k / \alpha_k^*$$

where v is the number of distinct units in the sample

Steps

- Find α_k^* for each unit in the sample
- Find $y_k J_k / \alpha_k^*$ for each unit
- Find t_{HT^*}

 α_k^*

$$\alpha_k^* = 1 - \binom{N - m_k}{n_1} / \binom{N}{n_1}$$

```
alpha_k<-function(sim_data){
    # get the response values for each tile sampled
    y_k<-sim_data$sample_tiles$y_k
    n1<-sim_data$n1
    N<-sim_data$grid_size^2
    1-choose(N-sim_data$sample_tiles$m_k, sim_data$n1)/choose(N,sim_data$n1)
}
alpha_k(sim_data)

## [1] 0.0250000 0.0250000 0.0250000 0.0250000 0.0250000 0.1848315 0.0250000
## [8] 0.0250000 0.0250000 0.0250000 0.0250000 0.0250000 0.0250000 0.0250000
## [15] 0.1848315 0.1848315 0.1848315 0.1848315 0.1848315 0.1848315
## [22] 0.1848315 0.1848315 0.1848315 0.1848315 0.1848315 0.1848315
## [29] 0.1848315 0.1848315 0.1848315 0.1848315 0.1848315
## [29] 0.1848315 0.1848315 0.1848315 0.1848315</pre>
```

Finding the estimator

$$t_{HT^*} = N^{-1} \sum_{k=1}^{v} y_k J_k / \alpha_k^*$$

```
modified_TH<-function(sim_data,plot=F){
   if(plot==T){
      plot_clusters(sim_data$points,samp=sim_data$sample_tiles)
   }
   a_k<-alpha_k(sim_data)
   N<-sim_data$grid_size^2
   y_k<-sim_data$sample_tiles$y_k
   J_k<-sim_data$sample_tiles$occupied
   1/N*sum(y_k*J_k/a_k)
}
modified_TH(sim_data)</pre>
```

[1] 0.26231

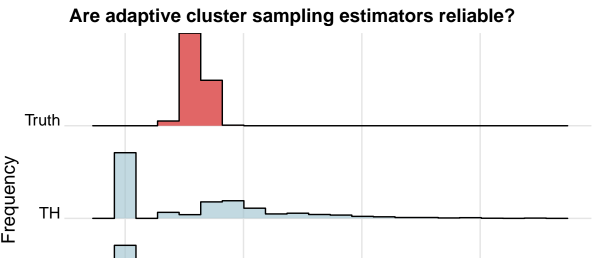
Running simulations

Initial results, Hard border, inclusion not forced

This was the first simulation performed where 1000 populations were sampled once, this limits what we are able to see though as the variation of the sampling method for a given population may also be of interest.

Moving forward with simulations rather than sampling thousands of populations we will only generate 9 populations but sample from them 1000 times each.

```
n_sim < -1000
results < - data.frame(HH=rep(NA,n_sim),
                     TH=rep(NA,n_sim),
                     Truth=rep(NA,n_sim))
for(i in 1:n_sim){
  sim_data<-simulate_one()</pre>
  results$HH[i] <-modified_HH(sim_data)</pre>
  results$TH[i] <-modified_TH(sim_data)</pre>
  results$Truth[i] <-length(sim_data$points$X)/(sim_data$grid_size^2)</pre>
}
write.csv(results,file="sampling 1000 populations.csv")
results <- read_csv("sampling 1000 populations.csv", show_col_types = FALSE)
## New names:
## * '' -> '...1'
results_long<-results %>%
  pivot_longer(cols = HH:Truth,
               names_to = "Estimator",
               values_to = "Value")
ggplot(results_long,aes(x=Value, y=Estimator, fill=Estimator))+
  geom_density_ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
  theme ridges()+
  theme(legend.position="none",
        axis.title.y = element_text( hjust = 0.5),
        axis.title.x = element_text( hjust = 0.5))+
  xlab("Estimate") +
  ylab("Frequency")+
  ggtitle("Are adaptive cluster sampling estimators reliable?")+
  scale_fill_manual(values=c("lightblue3","lightblue3","red3"))
```



exploring combinations of inclusion and border type Below are simulations for all four combinations of border type and inclusion type for 9 populations. There are 4 loops for simulations, one for inclusion type, one for border type, one for the 9 populations, and one to compute the estimators of the 1000 samples for the 9 populations.

0.50

Estimate

0.75

0.25

HH

0.00

```
m_samples<-1000 #
n_pops<-9
border_type<-c(FALSE,TRUE) # h</pre>
border_label<-c("hard", "soft")</pre>
inclusion_type<-c(FALSE,TRUE) # k</pre>
inclusion_label<-c("not_","")</pre>
for(h in 1:length(border_type)){
  for(k in 1:length(inclusion_type)){
    # Create blank dataframes for the estimates and the populations of points
    results <- data.frame(HH=rep(NA,n_pops*m_samples),
                         TH=rep(NA,n_pops*m_samples),
                         Truth=rep(NA,n_pops*m_samples),
                         Population=rep(NA,n_pops*m_samples))
    population_points<-data.frame(X=c(),Y=c(),Group=c(),Population=c())</pre>
    # set seed to ensure each set of populations are the same
    set.seed(pi)
    for(i in 1:n_pops){
      sim_data<-simulate_m(m=m_samples,</pre>
                             hard_border = border_type[h],
```

```
force_inclusion = inclusion_type[k])
      # save the population points to plot later
      population_points<-population_points%>%
        rbind( cbind(sim_data[[i]] $points, Population=paste("pop",i)) )
      for(j in 1:m_samples){
        # for each sample from the population in sim_data we compute our estimator
        # saving estimators into results
        results$HH[(i-1)*m samples+j]<-modified HH(sim data[[j]])
        results$TH[(i-1)*m_samples+j]<-modified_TH(sim_data[[j]])</pre>
        results$Truth[(i-1)*m_samples+j]<-length(sim_data[[j]]$points$X)/(sim_data[[j]]$grid_size^2)
        results$Population[(i-1)*m_samples+j]<-paste("Population",i)</pre>
     }
    }
    # save end results of simulation
    write.csv(results,file=paste("estimates-inclusion_",
                                  inclusion_label[k],
                                  "forced-",
                                  border_label[h], "_border.csv",
                                  sep=""))
    write.csv(population_points,file=paste("pop_points-inclusion_",
                                            inclusion_label[k],
                                            "forced-",
                                            border_label[h],
                                            " border.csv",sep=""))
 }
}
```

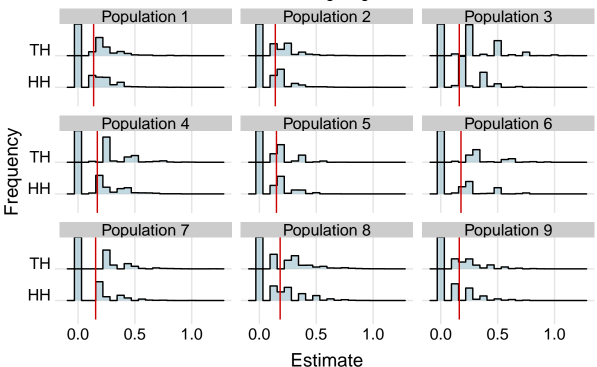
plotting simulation results and populations

```
for(i in c("soft", "hard")){
  for(j in c("_not","")){
    # read in results from simulation above
   results <-read_csv(paste("estimates-inclusion",j,"_forced-",i,"_border.csv",sep=""),
                      show_col_types = FALSE,
                      col_select = HH:Population)
   results long<-results%>%
      pivot_longer(cols = HH:TH,
                   names_to = "Estimator",
                   values_to = "Value")
        # plot the distributions of the estimators across the 9 populations
   p<-ggplot(results_long,aes(x=Value, y=Estimator, fill=Estimator))+</pre>
      geom density ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
      theme_ridges()+
      theme(legend.position="none",
            axis.title.y = element_text( hjust = 0.5),
            axis.title.x = element_text( hjust = 0.5))+
      xlab("Estimate") +
      ylab("Frequency")+
      ggtitle("Are adaptive cluster sampling estimators reliable?",
```

```
subtitle=paste("Inclusion",gsub("_","",j),"forced,", i,"border finding neighbours"))+
scale_fill_manual(values=c("lightblue3","lightblue3"))+
facet_wrap(~Population)+
geom_vline(aes(xintercept=Truth),color="red3")
plot(p)
}
```

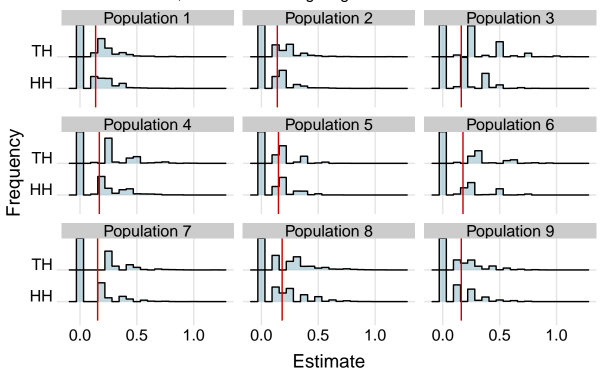
```
## New names:
## New names:
## * '' -> '...1'
```

Inclusion not forced, soft border finding neighbours



```
## New names:
## * '' -> '...1'
```

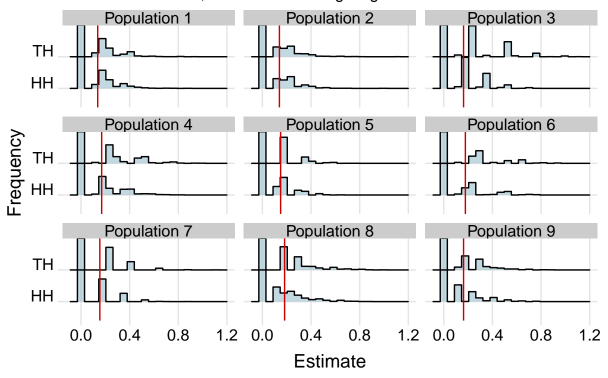
Inclusion forced, soft border finding neighbours



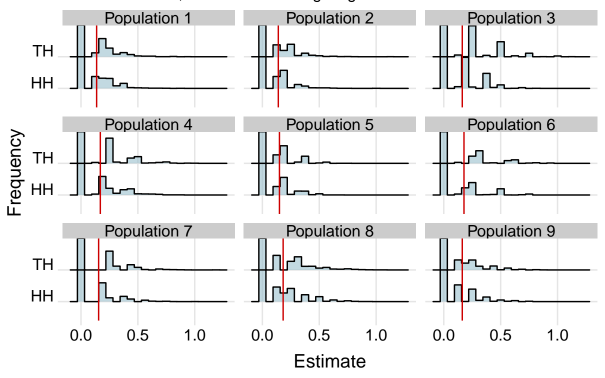
New names:

* '' -> '...1'

Inclusion not forced, hard border finding neighbours



Inclusion forced, hard border finding neighbours



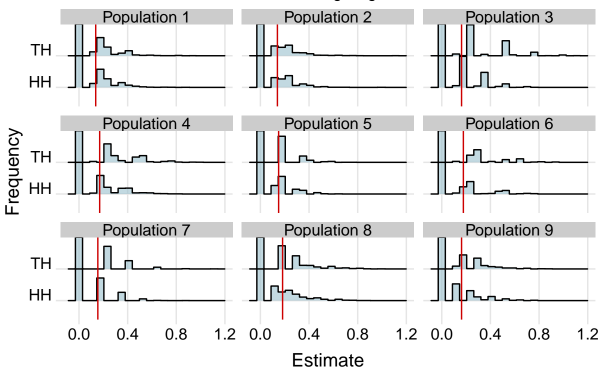
```
for(i in c("_not","")){
}
```

Inclusion not forced

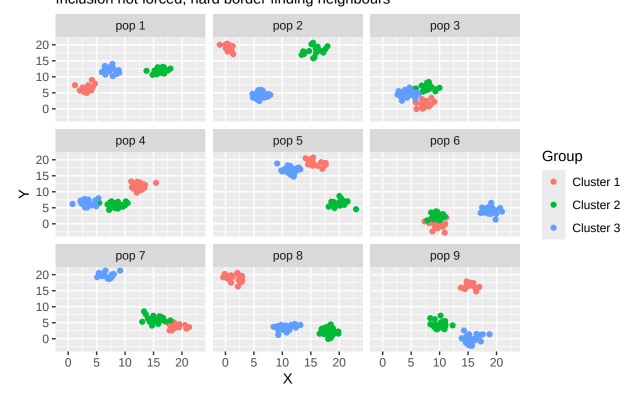
Hard border when finding neighbours

```
for(j in 1:n_samples){
    # saving estimators into results
   results$HH[(i-1)*n_samples+j]<-modified_HH(sim_data[[j]])
   results$TH[(i-1)*n_samples+j]<-modified_TH(sim_data[[j]])
   results$Truth[(i-1)*n_samples+j]<-length(sim_data[[j]]$points$X)/(sim_data[[j]]$grid_size^2)
   results $Population [(i-1)*n_samples+j] <-paste("Population",i)
 }
}
# save end results of simulation
write.csv(results,file="estimates-inclusion_not_forced-hard_border.csv")
write.csv(population_points,file="pop_points-inclusion_not_forced-hard_border.csv")
# read in results from simulation above
results <- read_csv("estimates-inclusion_not_forced-hard_border.csv", show_col_types = FALSE)
## New names:
## * '' -> '...1'
results_long<-results%>%
  pivot_longer(cols = HH:TH,
               names_to = "Estimator",
               values_to = "Value")
population_points<-read_csv("pop_points-inclusion_not_forced-hard_border.csv",show_col_types = FALSE)
## New names:
## * '' -> '...1'
# plot the distributions of the estimators across the 9 populations
ggplot(results_long,aes(x=Value, y=Estimator, fill=Estimator))+
  geom_density_ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
  theme_ridges()+
  theme(legend.position="none",
        axis.title.y = element_text( hjust = 0.5),
        axis.title.x = element_text( hjust = 0.5))+
  xlab("Estimate") +
  ylab("Frequency")+
  ggtitle("Are adaptive cluster sampling estimators reliable?",
          subtitle="Inclusion not forced, hard border finding neighbours")+
  scale_fill_manual(values=c("lightblue3","lightblue3"))+
  facet_wrap(~Population)+
  geom_vline(aes(xintercept=Truth),color="red3")
```

Inclusion not forced, hard border finding neighbours



How do these estimators vary across populations? Inclusion not forced, hard border finding neighbours

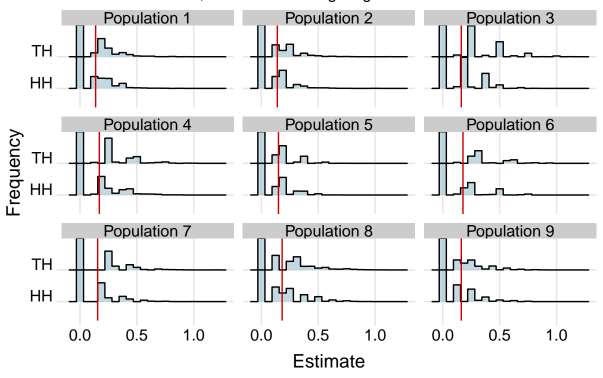


Soft border when finding neighbours

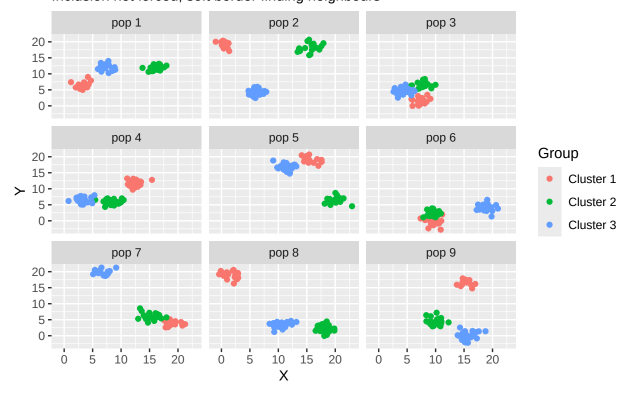
```
set.seed(pi)
n_samples < -1000
n_pops<-9
# estimator results
results <- data.frame(HH=rep(NA,n_pops*n_samples),
                    TH=rep(NA, n pops*n samples),
                    Truth=rep(NA,n_pops*n_samples),
                    Population=rep(NA,n_pops*n_samples))
# saving the population sampled from
population_points<-data.frame(X=c(),Y=c(),Group=c(),Population=c())</pre>
for(i in 1:n_pops){ # generating the populations and samples
  sim_data<-simulate_m(m=n_samples,hard_border = FALSE, force_inclusion = FALSE)</pre>
  # save the population points to plot later
  population_points<-population_points%>%
    rbind( cbind(sim_data[[i]]$points,Population=paste("pop",i)) )
  for(j in 1:n_samples){
    # save estimates for each sample
    results$HH[(i-1)*n_samples+j]<-modified_HH(sim_data[[j]])
    results$TH[(i-1)*n_samples+j]<-modified_TH(sim_data[[j]])
    results$Truth[(i-1)*n samples+j]<-length(sim data[[j]]$points$X)/(sim data[[j]]$grid size^2)
    results$Population[(i-1)*n_samples+j]<-paste("Population",i)</pre>
```

```
}
}
# save estimates of the samples and the populations sampled from
write.csv(results,file="estimates-inclusion_not_forced-soft_border.csv")
write.csv(population_points,file="pop_points-inclusion_not_forced-soft_border.csv")
results <- read_csv("estimates-inclusion_not_forced-soft_border.csv", show_col_types = FALSE)
## New names:
## * '' -> '...1'
results_long<-results%>%
  pivot_longer(cols = HH:TH,
               names_to = "Estimator",
               values_to = "Value")
population_points<-read_csv("pop_points-inclusion_not_forced-soft_border.csv",show_col_types = FALSE)
## New names:
## * '' -> '...1'
ggplot(results_long,aes(x=Value, y=Estimator, fill=Estimator))+
  geom_density_ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
  theme_ridges()+
  theme(legend.position="none",
        axis.title.y = element_text( hjust = 0.5),
        axis.title.x = element_text( hjust = 0.5))+
  xlab("Estimate") +
  ylab("Frequency")+
  ggtitle("Are adaptive cluster sampling estimators reliable?",
          subtitle="Inclusion not forced, soft border finding neighbours")+
  scale_fill_manual(values=c("lightblue3","lightblue3"))+
  facet_wrap(~Population)+
  geom_vline(aes(xintercept=Truth),color="red3")
```

Inclusion not forced, soft border finding neighbours



How do these estimators vary across populations? Inclusion not forced, soft border finding neighbours



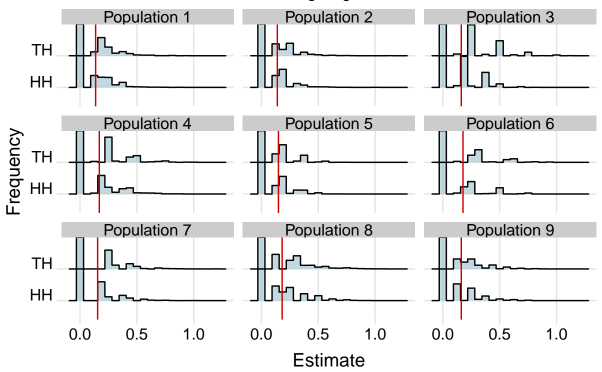
Inclusion forced

Hard border when finding neighbours

```
set.seed(pi)
n \text{ samples} < -1000
n_pops<-9
# Create blank dataframes for the estimates and the populations of points
results <- data.frame(HH=rep(NA,n_pops*n_samples),
                    TH=rep(NA,n_pops*n_samples),
                    Truth=rep(NA,n_pops*n_samples),
                    Population=rep(NA,n_pops*n_samples))
population_points<-data.frame(X=c(),Y=c(),Group=c(),Population=c())</pre>
for(i in 1:n_pops){
  sim_data<-simulate_m(m=n_samples,hard_border = TRUE, force_inclusion = TRUE)</pre>
  # save the population points to plot later
  population_points<-population_points%>%
    rbind( cbind(sim_data[[i]]$points,Population=paste("pop",i)) )
  for(j in 1:n_samples){
    # saving estimators into results
    results$HH[(i-1)*n samples+j]<-modified HH(sim data[[j]])
    results$TH[(i-1)*n_samples+j]<-modified_TH(sim_data[[j]])
```

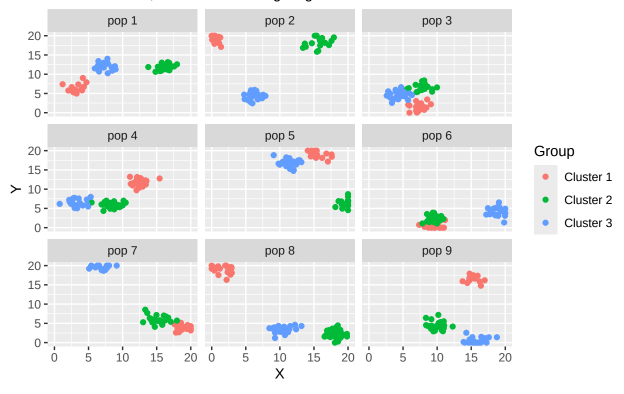
```
results$Truth[(i-1)*n_samples+j]<-length(sim_data[[j]]$points$X)/(sim_data[[j]]$grid_size^2)
    results$Population[(i-1)*n_samples+j]<-paste("Population",i)
  }
}
# save end results of simulation
write.csv(results,file="estimates-inclusion_forced-hard_border.csv")
write.csv(population points,file="pop points-inclusion forced-hard border.csv")
# read in results from simulation above
results<-read_csv("estimates-inclusion_forced-hard_border.csv",show_col_types = FALSE)</pre>
## New names:
## * ' ' -> ' . . . 1 '
results_long<-results%>%
  pivot_longer(cols = HH:TH,
               names_to = "Estimator",
               values_to = "Value")
population_points<-read_csv("pop_points-inclusion_forced-hard_border.csv",show_col_types = FALSE)
## New names:
## * '' -> '...1'
# plot the distributions of the estimators across the 9 populations
ggplot(results long,aes(x=Value, y=Estimator, fill=Estimator))+
  geom_density_ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
  theme_ridges()+
  theme(legend.position="none",
        axis.title.y = element_text( hjust = 0.5),
        axis.title.x = element_text( hjust = 0.5))+
  xlab("Estimate") +
  ylab("Frequency")+
  ggtitle("Are adaptive cluster sampling estimators reliable?",
          subtitle="Inclusion forced, hard border finding neighbours")+
  scale_fill_manual(values=c("lightblue3","lightblue3"))+
  facet_wrap(~Population)+
  geom vline(aes(xintercept=Truth),color="red3")
```

Inclusion forced, hard border finding neighbours



How do these estimators vary across populations?

Inclusion forced, hard border finding neighbours

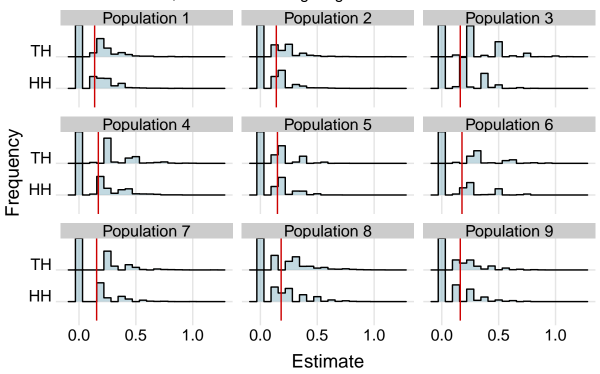


Soft border when finding neighbours

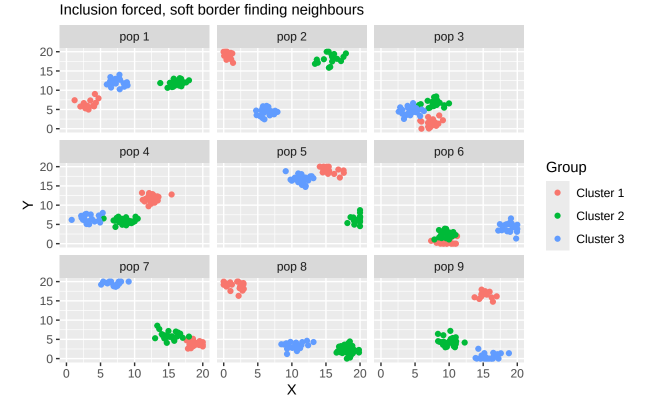
```
set.seed(pi)
n_samples < -1000
n_pops<-9
# Create blank dataframes for the estimates and the populations of points
results <- data.frame(HH=rep(NA,n_pops*n_samples),
                    TH=rep(NA, n pops*n samples),
                    Truth=rep(NA,n_pops*n_samples),
                    Population=rep(NA,n_pops*n_samples))
population points<-data.frame(X=c(),Y=c(),Group=c(),Population=c())
for(i in 1:n_pops){
  sim_data<-simulate_m(m=n_samples,hard_border = FALSE, force_inclusion = TRUE)</pre>
  # save the population points to plot later
  population_points<-population_points%>%
    rbind( cbind(sim_data[[i]] points, Population=paste("pop",i)) )
  for(j in 1:n_samples){
    # saving estimators into results
    results$HH[(i-1)*n_samples+j]<-modified_HH(sim_data[[j]])
    results$TH[(i-1)*n_samples+j]<-modified_TH(sim_data[[j]])
    results$Truth[(i-1)*n samples+j]<-length(sim data[[j]]$points$X)/(sim data[[j]]$grid size^2)
    results$Population[(i-1)*n_samples+j]<-paste("Population",i)</pre>
```

```
}
}
# save end results of simulation
write.csv(results,file="estimates-inclusion_forced-soft_border.csv")
write.csv(population_points,file="pop_points-inclusion_forced-soft_border.csv")
# read in results from simulation above
results <- read_csv("estimates-inclusion_forced-soft_border.csv", show_col_types = FALSE)
## New names:
## * '' -> '...1'
results_long<-results%>%
  pivot_longer(cols = HH:TH,
               names_to = "Estimator",
               values_to = "Value")
population_points<-read_csv("pop_points-inclusion_forced-soft_border.csv",show_col_types = FALSE)</pre>
## New names:
## * '' -> '...1'
# plot the distributions of the estimators across the 9 populations
ggplot(results_long,aes(x=Value, y=Estimator, fill=Estimator))+
  geom_density_ridges(alpha=0.6, stat="binline",bins=20,scale=1)+
  theme ridges()+
  theme(legend.position="none",
        axis.title.y = element_text( hjust = 0.5),
        axis.title.x = element_text( hjust = 0.5))+
  xlab("Estimate") +
  ylab("Frequency")+
  ggtitle("Are adaptive cluster sampling estimators reliable?",
          subtitle="Inclusion forced, soft border finding neighbours")+
  scale_fill_manual(values=c("lightblue3","lightblue3"))+
  facet_wrap(~Population)+
  geom_vline(aes(xintercept=Truth),color="red3")
```

Inclusion forced, soft border finding neighbours



How do these estimators vary across populations?



exploring population 7

[1] 12