

# BumbleboxExampleAnalysis

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Code available at: [www.github.com/Crall-Lab/BumbleBox\\_BOMBUSS](https://www.github.com/Crall-Lab/BumbleBox_BOMBUSS)

## Set up

```
#Load packages
library(ggplot2)
library(png)
library(lubridate)

##
## Attaching package: 'lubridate'

## The following objects are masked from 'package:base':
##   date, intersect, setdiff, union

library(stringr)
library(grid)
library(lme4)

## Loading required package: Matrix

library(lmerTest)

##
## Attaching package: 'lmerTest'

## The following object is masked from 'package:lme4':
##   lmer

## The following object is masked from 'package:stats':
##   step
```

```

library(gridExtra)
library(igraph)

## 
## Attaching package: 'igraph'

## The following objects are masked from 'package:lubridate':
## 
##     %--%, union

## The following objects are masked from 'package:stats':
## 
##     decompose, spectrum

## The following object is masked from 'package:base':
## 
##     union

#Define parent directory containing data you want to analyze
#This example uses data from a single Bombus impatiens microcolony (data will be provided to workshop p
pdir <- '/Users/jamescrall/Downloads/2023-10-28'
brood_metadata <- read.csv('/Users/jamescrall/Downloads/2023-10-28/bumblebox-01-10-28-nest_image.csv')

#Define image dimensions for plotting
x1 <- 4056
y1 <- 3040

```

## Load and plot data tracking data from single video

```

#Get list of unique trials (using recursive search in case there are subfolders)
trials <- list.files(pdir, pattern = '*averages.csv', full.names = TRUE, recursive = TRUE)

#Select a random trial - trial 10 in this list
i <- 10

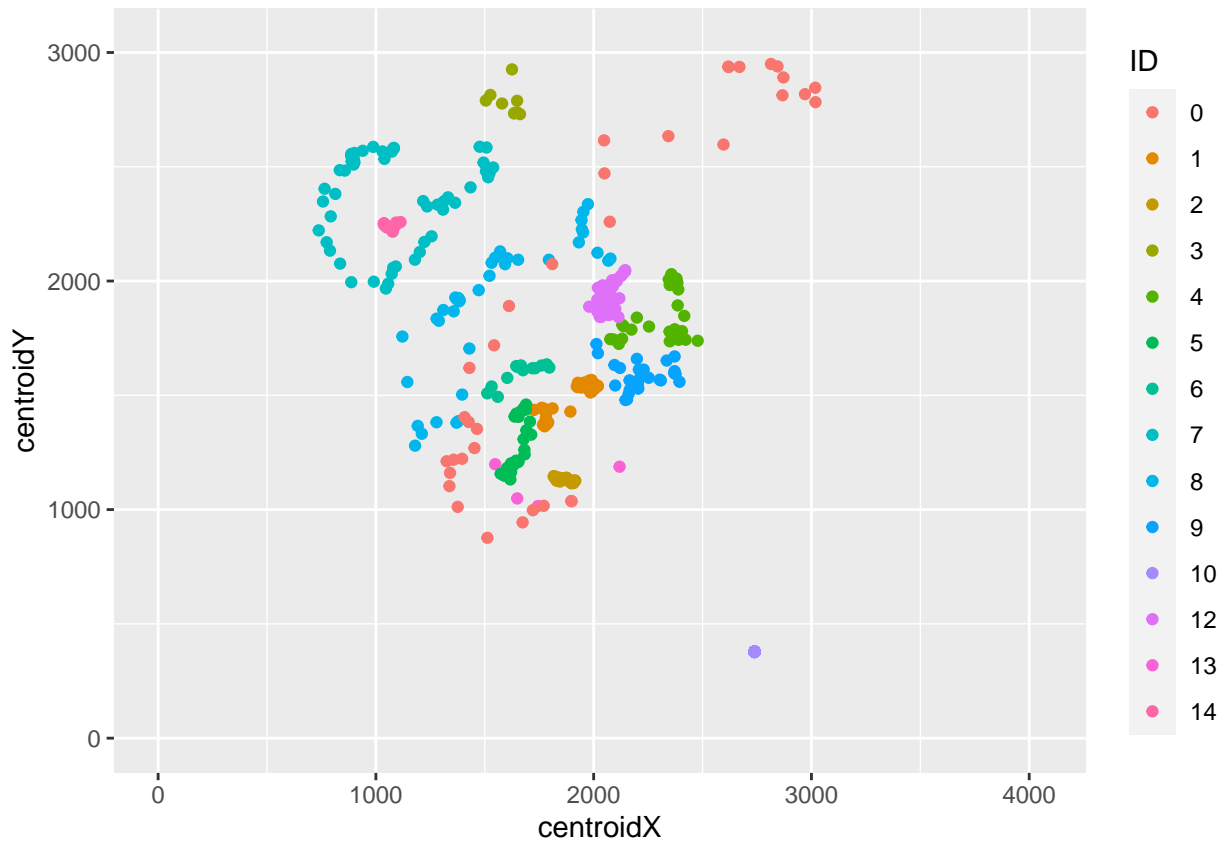
#Load background photo
background_im <- readPNG(str_replace(trials[i], '_averages.csv', '.png'))

#Clip image excess
background_im <- background_im[1:3040,]

#Optional: plot image (this can be slow)
#image(background_im, col = gray.colors(33))

#load and plot tag tracking data
tracking_data <- read.csv(str_replace(trials[i], '_averages.csv', '_updated.csv'))
tracking_data$ID <- as.factor(tracking_data$ID)
ggplot(tracking_data, aes(x = centroidX, y = centroidY, colour = ID)) + geom_point() + xlim(c(0,x1)) + ylim(c(0,y1))

```



```
#average data
summary_data <- read.csv(trials[i])
head(summary_data)
```

```
##                                     filename ID average.distance.from.center
## 1 bumblebox-01_2023-10-28_00-55-03  0                      783
## 2 bumblebox-01_2023-10-28_00-55-03  1                      194
## 3 bumblebox-01_2023-10-28_00-55-03  2                      492
## 4 bumblebox-01_2023-10-28_00-55-03  3                     1060
## 5 bumblebox-01_2023-10-28_00-55-03  4                      497
## 6 bumblebox-01_2023-10-28_00-55-03  5                      433
##   average.speed frames.tracked.in.video
## 1            130                  35
## 2             20                  48
## 3              6                  45
## 4             52                   9
## 5             49                  28
## 6             20                  48
```

## Explore data across all trials

First loop and compile across individual tracking files to compile into a single data frame

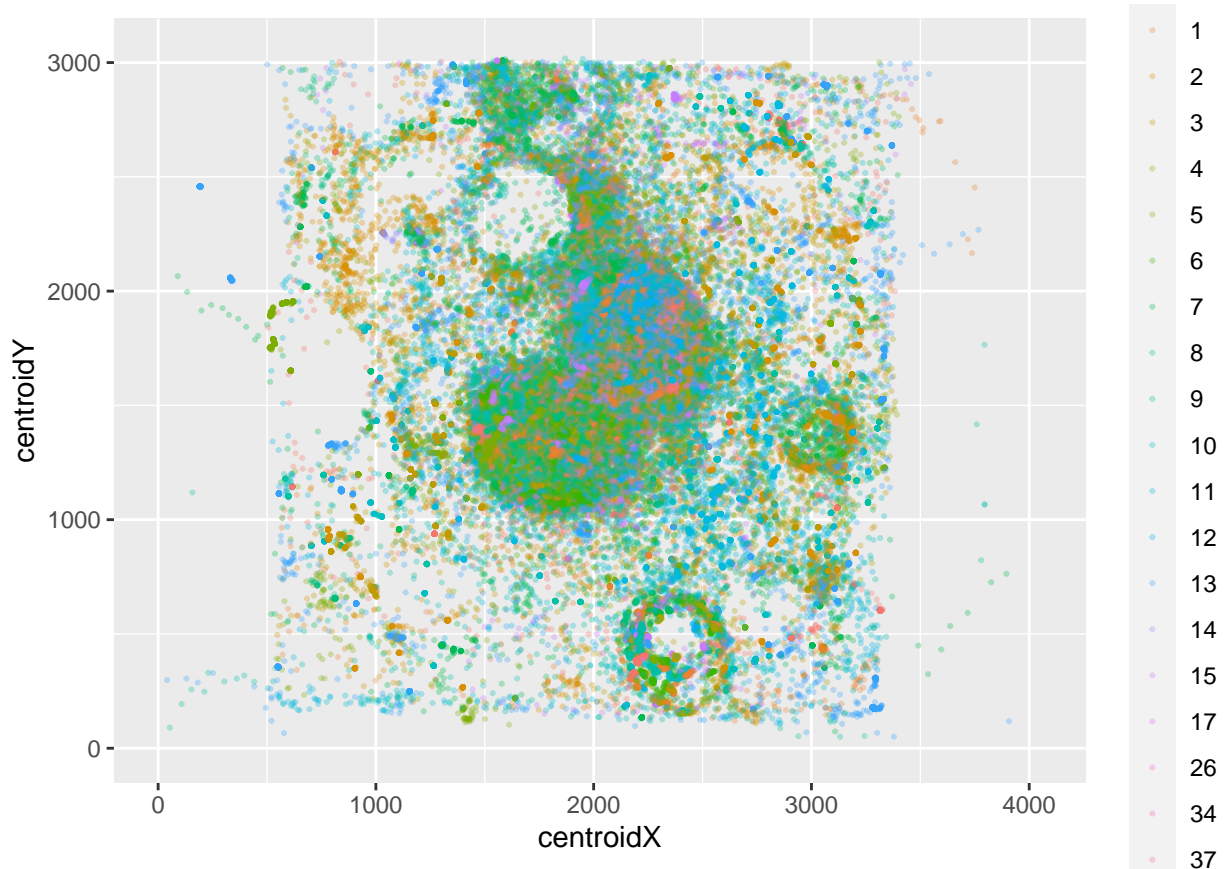
```
#Make sure you don't have a 'comb_data' object in memory
rm('tracking_data')

for(file in trials){

  data <- read.csv(str_replace(file, '_averages.csv', '_updated.csv'))
  ## Compile average data
  #Check to see if output data frame exists
  if(!exists('tracking_data')){
    tracking_data <- data #If not, create it from the first loop
  } else {
    tracking_data <- rbind(tracking_data, data) #Otherwise append new data to the end
  }

}

tracking_data$ID <- as.factor(tracking_data$ID)
ggplot(tracking_data, aes(x = centroidX, y = centroidY, colour = ID)) + geom_point(alpha = 0.3, cex = 0)
```



```
## Next, compile data on within-trial average data (generated directly from BumbleBox scripts)
```

```

#Make sure you don't have a 'comb_data' object in memory
rm('comb_data')

## Warning in rm("comb_data"): object 'comb_data' not found

for(file in trials){

  data <- read.csv(file)
  ## Compile average data

  #Check to see if output data frame exists
  if(!exists('comb_data')){
    comb_data <- data #If not, create it from the first loop
  } else {
    comb_data <- rbind(comb_data, data) #Otherwise append new data to the end
  }

}

#Convert ID to factor instead of numeric
comb_data$ID <- as.factor(comb_data$ID)

```

Plot variation in speed and distance from center across individuals

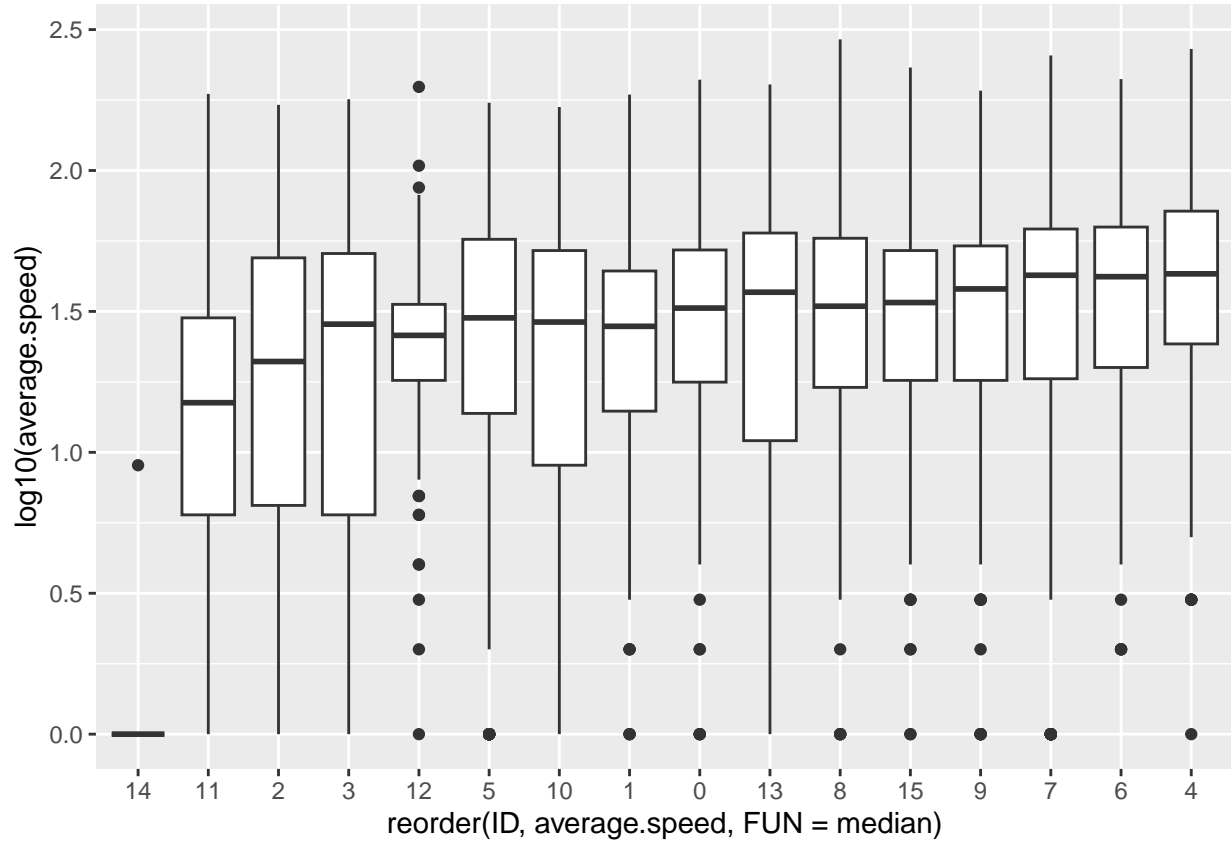
```

#Clean out missing data
comb_data <- comb_data[complete.cases(comb_data),]

#Boxplot of speed vs. individual
ggplot(comb_data, aes(x = reorder(ID, average.speed, FUN = median), y = log10(average.speed)))+geom_boxp

## Warning: Removed 103 rows containing non-finite values ('stat_boxplot()').

```

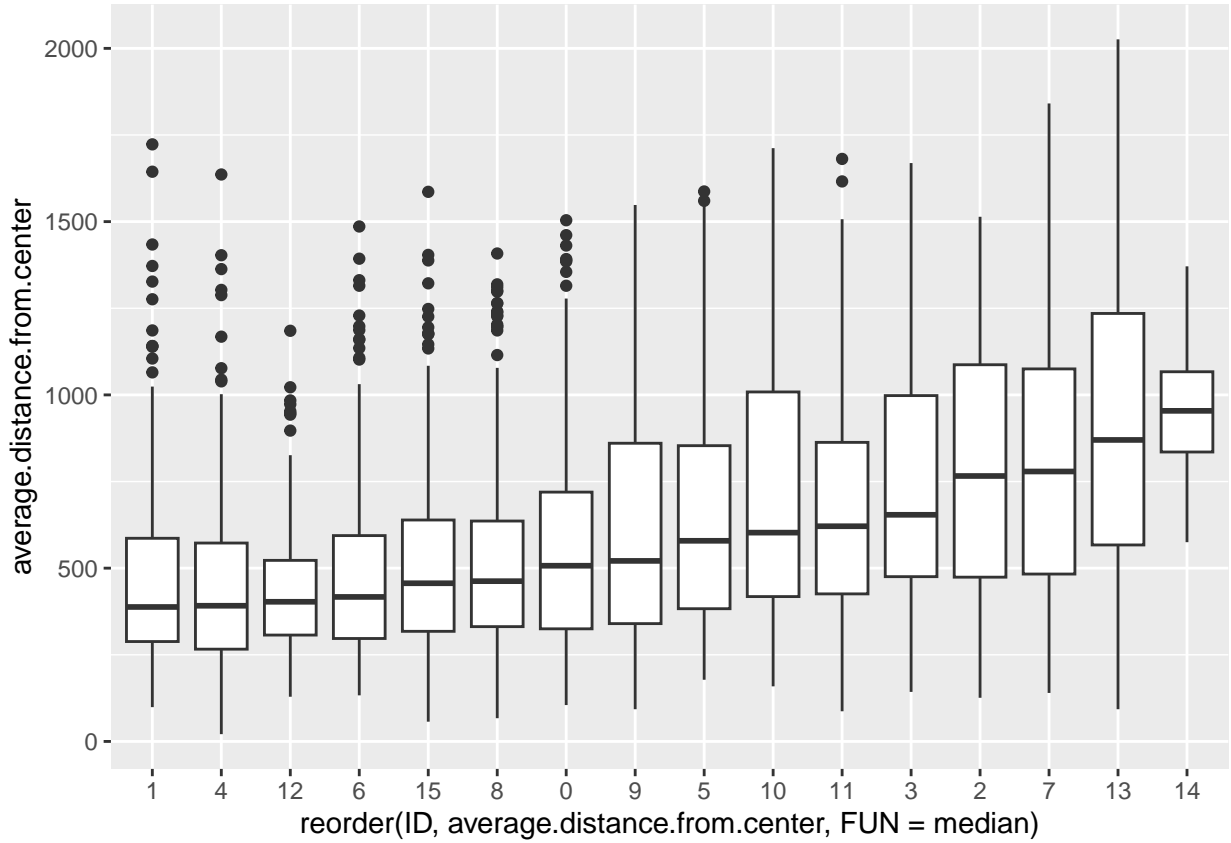


```
#Check for evidence of individual variation
model <- lmer(log10(1+average.speed)~ID + (1|filename), data = comb_data)
model.1 <- lmer(log10(1+average.speed)~1 + (1|filename), data = comb_data)
anova(model, model.1)

## refitting model(s) with ML (instead of REML)

## Data: comb_data
## Models:
## model.1: log10(1 + average.speed) ~ 1 + (1 | filename)
## model: log10(1 + average.speed) ~ ID + (1 | filename)
##          npar   AIC   BIC logLik deviance Chisq Df Pr(>Chisq)
## model.1     3 4660.9 4678.9 -2327.5    4654.9
## model      18 4199.4 4307.3 -2081.7    4163.4 491.56 15 < 2.2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

#Show distance from center by individual
ggplot(comb_data, aes(x = reorder(ID, average.distance.from.center, FUN = median), y = average.distance
```



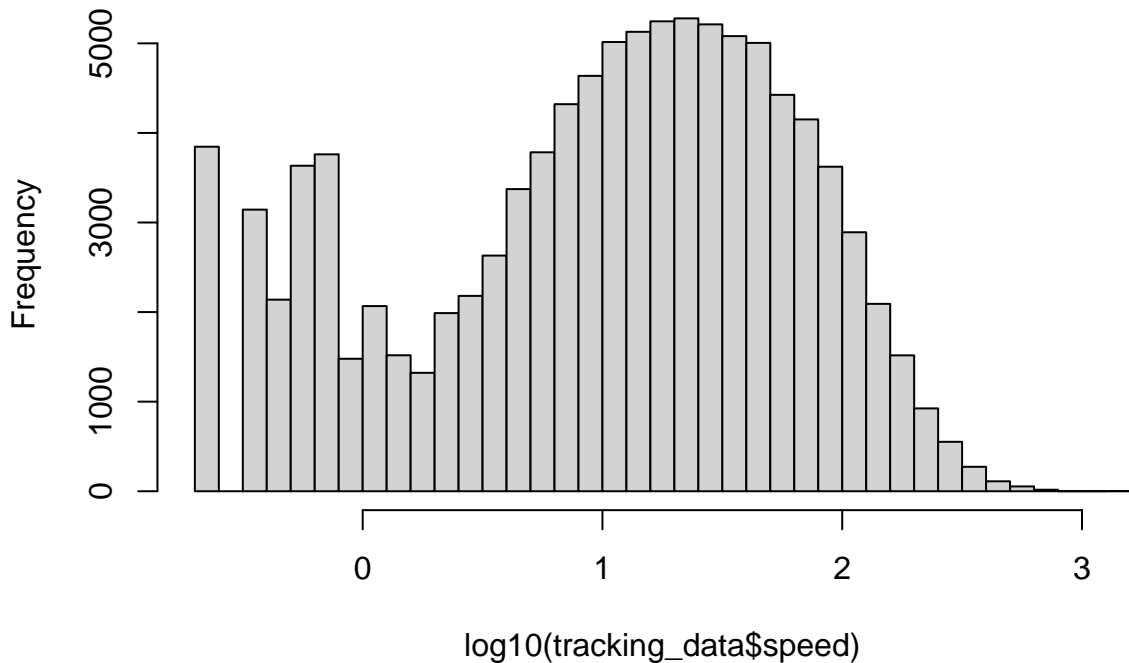
```
#Get individual averages
individual_averages <- aggregate(cbind(average.distance.from.center, average.speed, frames.tracked.in.v),
colnames(individual_averages) <- c('ID', 'dist', 'speed', 'frames_tracked')

# Remove outlier with zero velocity
individual_averages <- subset(individual_averages, speed > 1)
```

Determine activity (moving vs. not moving) by thresholding speed

```
#Plot histogram of framewise speeds to determine threshold for 'real' movement
hist(log10(tracking_data$speed), breaks = 30)
```

## Histogram of log10(tracking\_data\$speed)



```
#Looks like ~3.1 pixels/frame (log10(3.1)~= 0.5)
speed_thresh <- 3.1
tracking_data$moving[tracking_data$speed > speed_thresh] <- 1
tracking_data$moving[tracking_data$speed < speed_thresh] <- 0
```

Visualize changes in individual behavior over time

```
#Visualize individual speed/distance over time

#Set up timestamps
#Reference start time
start.date <- parse_date_time('2023-10-28 00:00:00', "%Y-%m-%d %H:%M:%S")

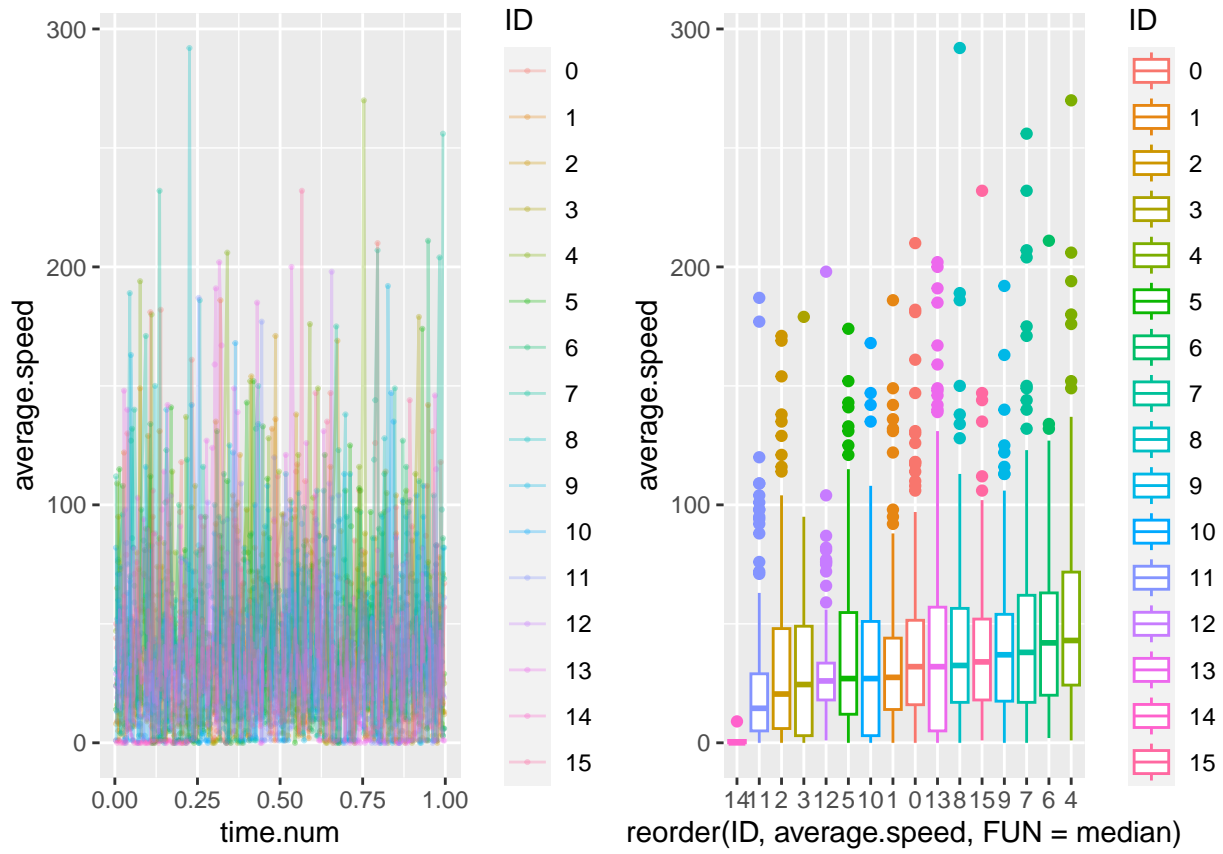
#create timestamp variable
comb_data$datetime <- parse_date_time(substr(comb_data$filename, 14, 32), "%Y-%m-%d_%H-%M-%S")

#Convert into 'days since the start time'
comb_data$time.num <- as.numeric(difftime(comb_data$datetime, start.date, units = 'days'))

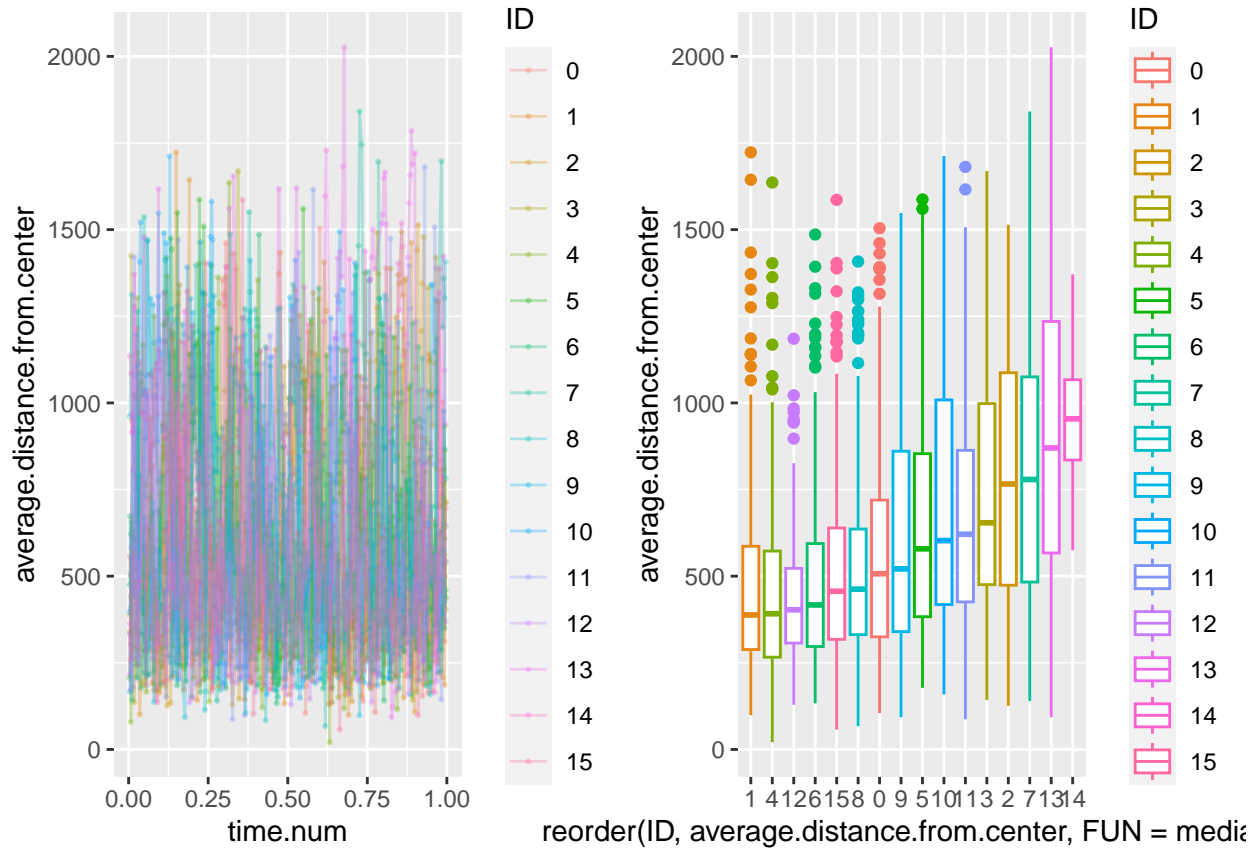
#Reorder
comb_data <- comb_data[order(comb_data$ID, comb_data$time.num),]

#Plots over time and individual variation for speed
p1 <- ggplot(comb_data, aes(x = time.num, y = average.speed, colour = ID)) + geom_point(alpha = 0.3, ce
```

```
p2 <- ggplot(comb_data, aes(x = reorder(ID, average.speed, FUN = median), y = average.speed, colour = ID))
grid.arrange(p1, p2, ncol=2)
```



```
#Plots over time and individual variation for distance from center
p1 <- ggplot(comb_data, aes(x = time.num, y = average.distance.from.center, colour = ID)) + geom_point()
p2 <- ggplot(comb_data, aes(x = reorder(ID, average.distance.from.center, FUN = median), y = average.distance.from.center, colour = ID))
grid.arrange(p1, p2, ncol=2)
```



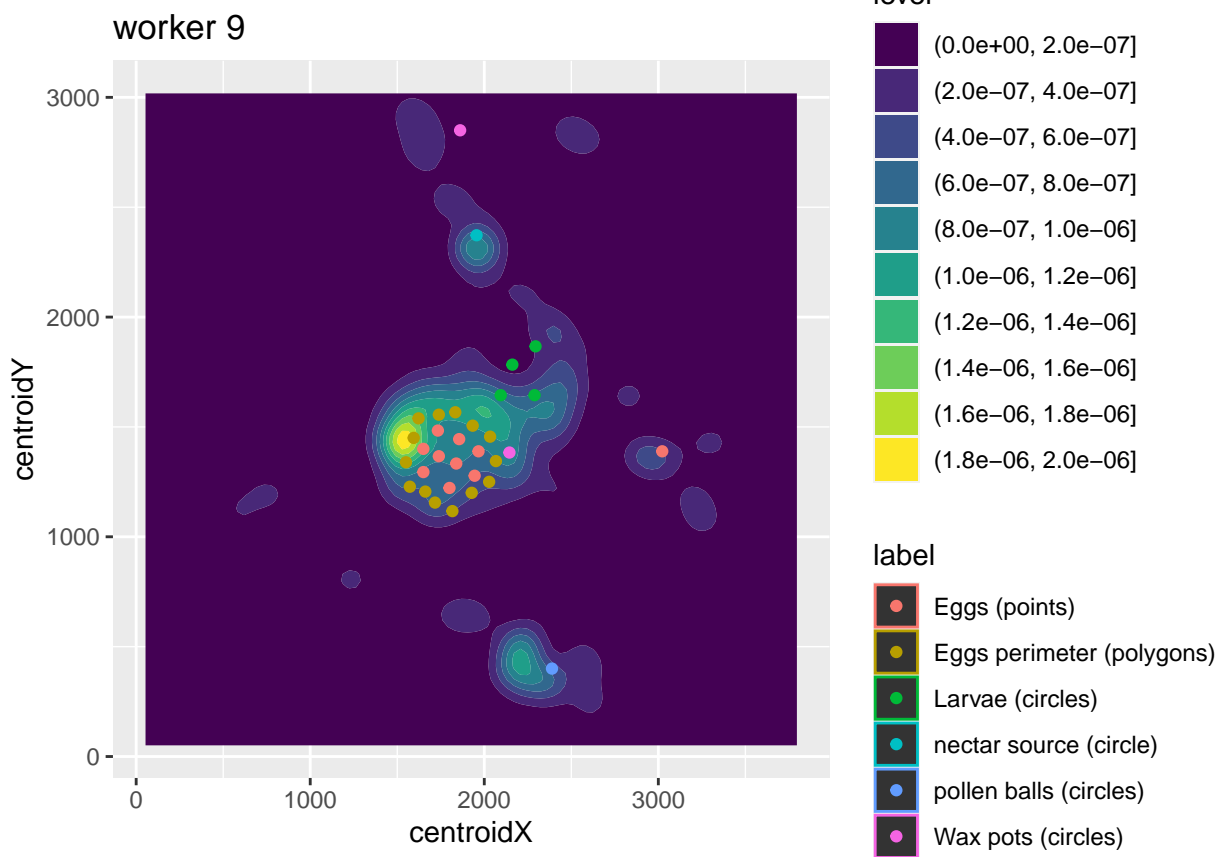
```
#Get average time spent moving for each bee
aggregate(moving~ID, data = tracking_data, FUN = mean)
```

```
##      ID      moving
## 1    0 0.820146223
## 2    1 0.841329966
## 3    2 0.648259994
## 4    3 0.561511140
## 5    4 0.914539920
## 6    5 0.727098956
## 7    6 0.889504950
## 8    7 0.764985260
## 9    8 0.824413818
## 10   9 0.765464770
## 11  10 0.591617345
## 12  11 0.579634465
## 13  12 0.892954723
## 14  13 0.608617095
## 15  14 0.005407654
## 16  15 0.876669635
```

Plot individual spatial occupancy patterns within the nest

```
#Plot for single representative individual, with brood locations overlaid
```

```
ggplot(subset(tracking_data, ID == 9), aes(x = centroidX, y = centroidY)) + geom_density_2d_filled() + gg
```



```
#Plot spatial distribution for select individuals (representing centrality range)
```

```
#NB: if you are working on your own data set, select the IDs here from your own taglist
```

```
worker_list <- c('1', '13', '15', '3')
```

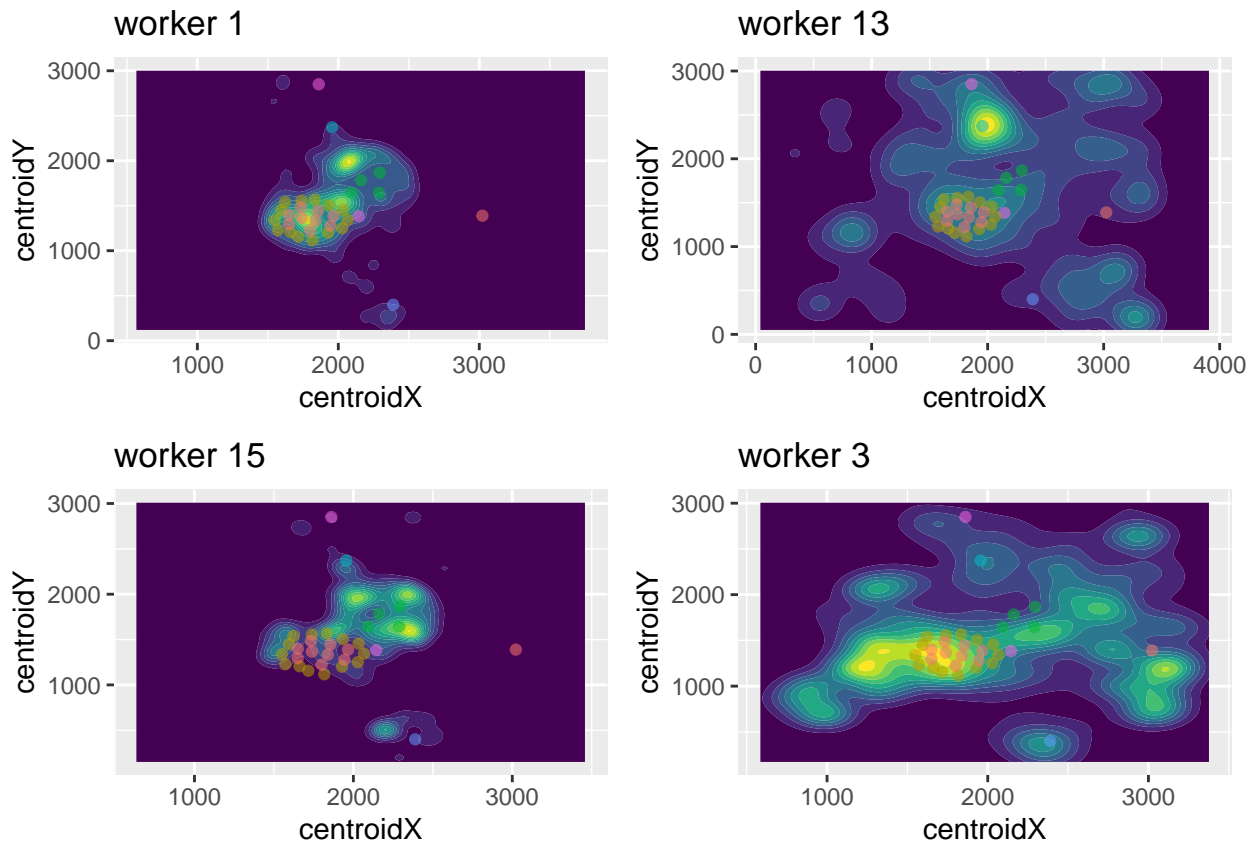
```
p1 <- ggplot(subset(tracking_data, ID == worker_list[1]), aes(x = centroidX, y = centroidY)) + geom_density_2d()
```

```
p2 <- ggplot(subset(tracking_data, ID == worker_list[2]), aes(x = centroidX, y = centroidY)) + geom_density_2d()
```

```
p3 <- ggplot(subset(tracking_data, ID == worker_list[3]), aes(x = centroidX, y = centroidY)) + geom_density_2d()
```

```
p4 <- ggplot(subset(tracking_data, ID == worker_list[4]), aes(x = centroidX, y = centroidY)) + geom_density_2d()
```

```
grid.arrange(p1,p2,p3,p4, ncol = 2)
```



Analyze contact-based social networks. \*NB: this code set up for a data set with sixteen workers, and would need to be updated for different colony sizes

```
#Start by loading contact data by looping over source files

#Create empty output data structure
out_data <- as.data.frame(matrix(ncol = 17, nrow = 0))
colnames(out_data) <- as.character(c('ID', as.character(seq(0,15)))))

#loop over trials
for(file in trials){

  #Get filename specifically for 'contacts.csv' data
  data <- read.csv(str_replace(file, '_averages.csv', '_contacts.csv'))

  #Get rid of 'X' in filenames
  names(data) <- gsub("X", "", names(data))

  #Remove columns that aren't in output data
  data <- data[, colnames(data) %in% colnames(out_data)] 

  #Create and populate dummy intermediate data structure to align columns with output
  empty_data <- as.data.frame(matrix(ncol = 17, nrow = length(data[,1])))
  colnames(empty_data) <- colnames(out_data)
}
```

```

empty_data[,match(colnames(data), colnames(out_data))] <- data

#Aggregate data
out_data <- rbind(out_data, empty_data)

}

#Now, get average interaction rates across all pairwise individuals
interactions <- aggregate(out_data[,2:17], by = list(out_data$ID), FUN = mean, na.rm = TRUE)

#Remove data from errant tags and restructure into adjacency matrix
interactions <- interactions[1:16,2:17]
interactions <- as.matrix(interactions)
interactions[is.na(interactions)] <- 0
interactions[interactions == 1] <- 0

#Make fruchtermann-reingold graph using adjaceny matrix
net <- graph.adjacency(interactions, mode = 'undirected', weighted=TRUE, diag=FALSE)
plot.igraph(net,layout=layout.fruchterman.reingold, edge.width=E(net)$weight*50, vertex.size = 20)

```

