Ramírez-Mejía et al. (2021) Activity and habitat use by understory birds in a native Andean forest and a eucalypt plantation. The Wilson Journal of Ornithology.

Actividad y uso de hábitat de aves de sotobosque en un bosque andino nativo y una reforestación de eucalipto

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null

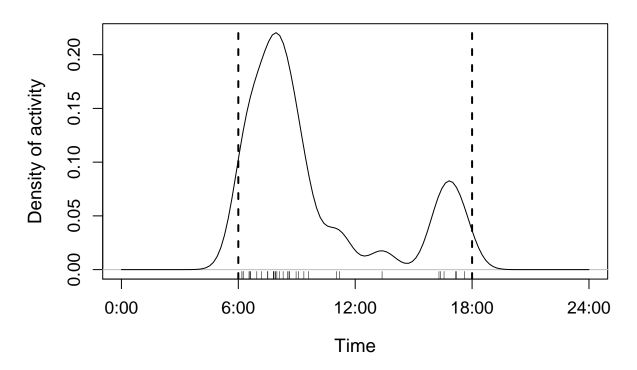
This document's primary purpose is to facilitate access to the code from different devices, so I did not include a detailed explanation. I hope it is clear enough to understand the way I conducted the activity analysis. If you have any concerns, questions, or suggestions, feel free to contact me.

```
#### aves bosques de la chec
library(tidyverse)
chec <- read.delim("bird_activity_data.txt")</pre>
str(chec)
                    299 obs. of 3 variables:
## 'data.frame':
    $ species: chr "brun" "brun" "brun" "brun" ...
  $ forest : chr "andean" "andean" "andean" "andean" ...
                    0.304 0.347 0.656 0.442 0.281 ...
for (i in 1:ncol(chec)) {
  if (is.character(chec[[i]])) {
    chec[[i]] <- as.factor(chec[[i]])</pre>
 } else {
    next
}
head(chec)
```

```
## species forest time
## 1 brun andean 0.3041667
## 2 brun andean 0.3472083
## 3 brun andean 0.6562500
## 4 brun andean 0.4423333
## 5 brun andean 0.2805417
## 6 brun andean 0.3770833
```

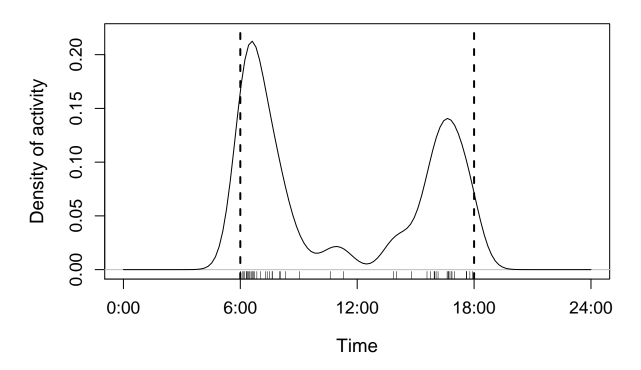
```
summary(chec)
##
     species
                         forest
                                        time
## brun :49
                            :224 Min.
                                         :0.2389
               andean
               reforestation: 75 1st Qu.:0.3079
## frena:97
## goud :53
                                   Median :0.4014
## mille:28
                                   Mean :0.4540
## rufi :39
                                   3rd Qu.:0.6235
## torq :33
                                   Max. :0.7583
range(chec$time)
## [1] 0.2388750 0.7583333
timeRad <- chec$time * 2 * pi</pre>
table(chec$species,chec$forest)
##
##
           andean reforestation
##
     brun
              15
                             34
               82
##
                             15
     frena
               44
                             9
##
    goud
                             2
               26
##
    mille
##
    rufi
               35
                             4
##
    torq
               22
                             11
library(boot)
library(circular)
library(overlap)
library(cowplot)
library(gridExtra)
library(png)
#patrón de actividad por especie----
\#Arremon\ torquatus
torquatus <- timeRad[chec$species == "torq"]</pre>
densityPlot(torquatus, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1.2,
abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)
```

torquatus



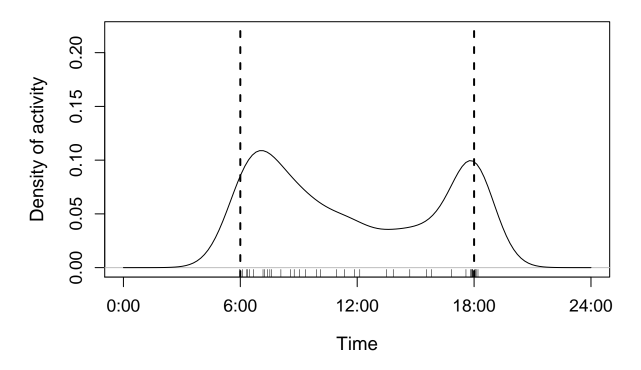
```
#Arremon brunneinucha
brunneinucha <- timeRad[chec$species == "brun"]
densityPlot(brunneinucha, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1
abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)</pre>
```

brunneinucha



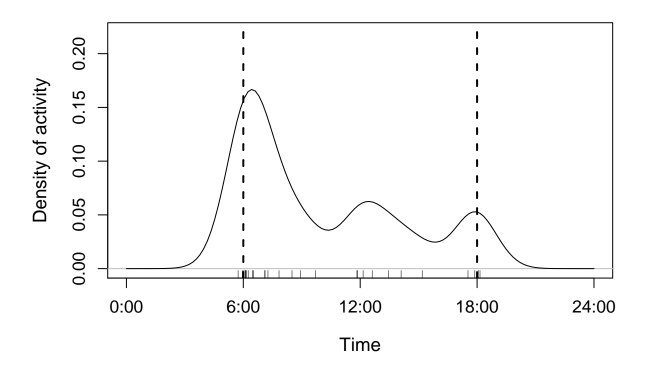
```
#Grallaria ruficapilla
ruficapilla <- timeRad[chec$species == "rufi"]
densityPlot(ruficapilla, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1.
abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)</pre>
```

ruficapilla



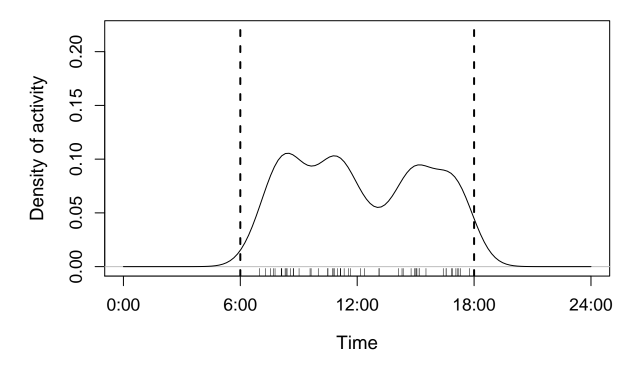
```
#Grallaria milleri
milleri <- timeRad[chec$species == "mille"]
densityPlot(milleri, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1.2, e
abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)</pre>
```

milleri



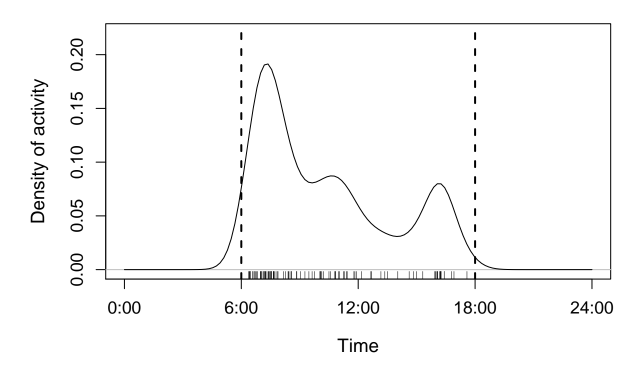
```
#Chamaepetes goudotii
goudotii <- timeRad[chec$species == "goud"]
densityPlot(goudotii, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1.2, abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)</pre>
```

goudotii



```
#Zentrigon frenata
frenata <- timeRad[chec$species == "frena"]
densityPlot(frenata, rug=TRUE, ylim=c(0,0.22), ylab="Density of activity", cex.axis=1.1, cex.lab=1.2, e
abline(v=6, lty="dashed", col="black", lwd=2); abline(v=18, lty="dashed", col="black", lwd=2)</pre>
```

frenata



```
levels(chec$species)
```

```
## [1] "brun" "frena" "goud" "mille" "rufi" "torq"
```

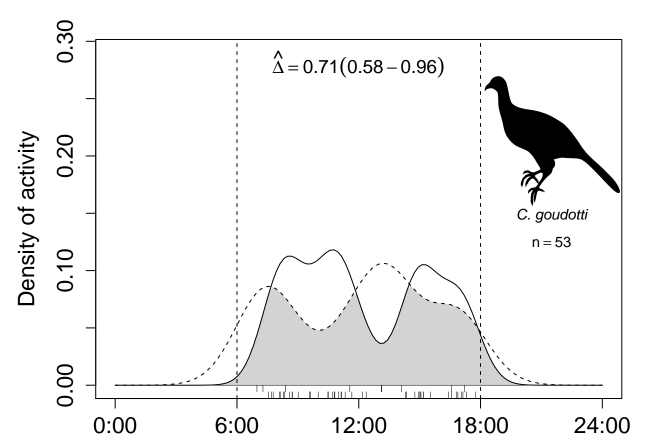
table(chec\$species, chec\$forest)

```
##
##
           andean reforestation
##
                15
                              34
     brun
##
     frena
               82
                               15
                               9
                44
##
     goud
##
     mille
               26
                                2
     rufi
                35
                               4
##
                22
                               11
##
     torq
```

```
#
#
#coeficiente de solapamiento----
#C. goudotti ----
#overlap calculation
bosq <- timeRad[chec$species == "goud" & chec$forest == "andean"]
plant <- timeRad[chec$species == "goud" & chec$forest == "reforestation"]
min(length(bosq), length(plant))</pre>
```

```
BosqPlantMILLE <- overlapEst(bosq, plant)
BosqPlantMILLE</pre>
```

```
## Dhat1 Dhat4 Dhat5
## 0.7271702 0.7294513 0.6666667
```



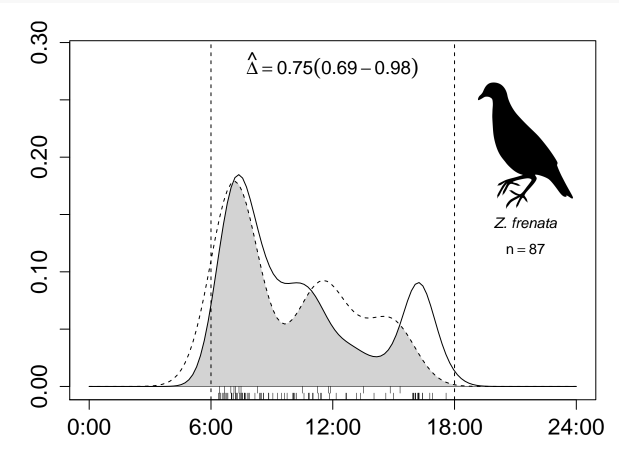
```
#dev.off()
#legend("topleft", c("Andean ", "G. ruficapilla"), lty=c(1,2), col=c(1,4), bty="n")
#bootstrap
bosqboot <- resample(bosq, 10000)
dim(bosqboot)</pre>
```

```
## [1]
          44 10000
plantboot <- resample(plant, 10000)</pre>
dim(plantboot)
## [1]
           9 10000
#overlap estimates by boostrap
BosqPlant <- bootEst(bosqboot, plantboot, adjust = c(1, NA, NA))</pre>
dim(BosqPlant)
## [1] 10000
BSmean <- colMeans(BosqPlant)</pre>
BSmean
##
                 Dhat4
                            Dhat5
       Dhat1
## 0.7098421
                    NA
                               NA
#calculation of CI
tmp <- BosqPlant[, 1]</pre>
bootCI(BosqPlantMILLE[1], tmp)
##
              lower
                         upper
## norm 0.5517511 0.9372454
## norm0 0.5344230 0.9199173
## basic 0.5807783 0.9659897
## basic0 0.5056787 0.8908901
## perc 0.4883506 0.8735620
tmp <- BosqPlant[, 1] # Extract the required column of the matrix</pre>
bootCIlogit(BosqPlantMILLE[1], tmp)
##
              lower
                         upper
## norm 0.5107887 0.8798348
## norm0 0.5016149 0.8759001
## basic 0.5069488 0.8815541
## basic0 0.4975240 0.8775605
## perc 0.4883506 0.8735620
#Z. frenata ----
#overlap calculation
bosq <- timeRad[chec$species == "frena" & chec$forest == "andean"]</pre>
plant <- timeRad[chec$species == "frena" & chec$forest == "reforestation"]</pre>
min(length(bosq), length(plant))
```

[1] 15

```
BosqPlantMILLE <- overlapEst(bosq, plant)
BosqPlantMILLE</pre>
```

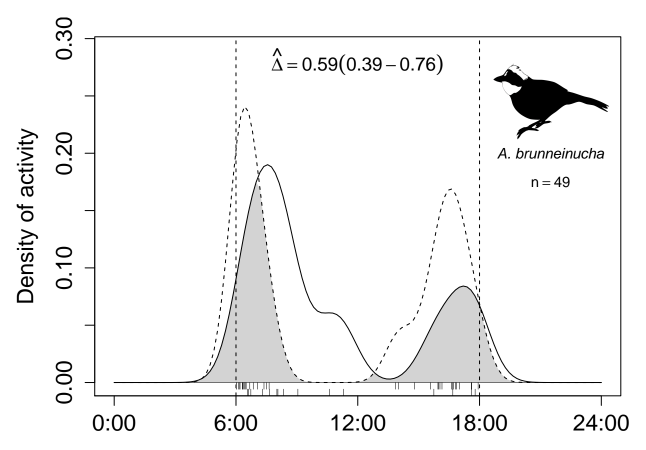
```
## Dhat1 Dhat4 Dhat5
## 0.7904093 0.8246503 0.8617886
```



```
#dev.off()
#legend("topleft", c("Andean ", "G. ruficapilla"), lty=c(1,2), col=c(1,4), bty="n")
#bootstrap
bosqboot <- resample(bosq, 10000)
dim(bosqboot)</pre>
```

[1] 82 10000

```
plantboot <- resample(plant, 10000)</pre>
dim(plantboot)
## [1]
          15 10000
#overlap estimates by boostrap
BosqPlant <- bootEst(bosqboot, plantboot, adjust = c(1, NA, NA))</pre>
dim(BosqPlant)
## [1] 10000
                 3
BSmean <- colMeans(BosqPlant)</pre>
BSmean
      Dhat1
                         Dhat5
##
               Dhat4
## 0.757651
                            NA
                  NA
#calculation of CI
tmp <- BosqPlant[, 1]</pre>
bootCI(BosqPlantMILLE[1], tmp)
##
              lower
                         upper
## norm 0.6812005 0.9651347
## norm0 0.6484422 0.9323764
## basic 0.6998351 0.9832851
## basic0 0.6302918 0.9137418
## perc 0.5975335 0.8809835
tmp <- BosqPlant[, 1] # Extract the required column of the matrix</pre>
bootCIlogit(BosqPlantMILLE[1], tmp)
##
              lower
                         upper
## norm 0.6631668 0.9063621
## norm0 0.6297424 0.8931832
## basic 0.6576889 0.9054746
## basic0 0.6321671 0.8954914
## perc 0.5975335 0.8809835
#A. brunneinucha ----
#overlap calculation
bosq <- timeRad[chec$species == "brun" & chec$forest == "andean"]</pre>
plant <- timeRad[chec$species == "brun" & chec$forest == "reforestation"]</pre>
min(length(bosq), length(plant))
## [1] 15
BosqPlantMILLE <- overlapEst(bosq, plant)</pre>
{\tt BosqPlantMILLE}
       Dhat1
                 Dhat4
                            Dhat5
## 0.5838830 0.5863419 0.5549020
```



```
#dev.off()
#legend("topleft", c("Andean ", "G. ruficapilla"), lty=c(1,2), col=c(1,4), bty="n")
#bootstrap
bosqboot <- resample(bosq, 10000)
dim(bosqboot)</pre>
```

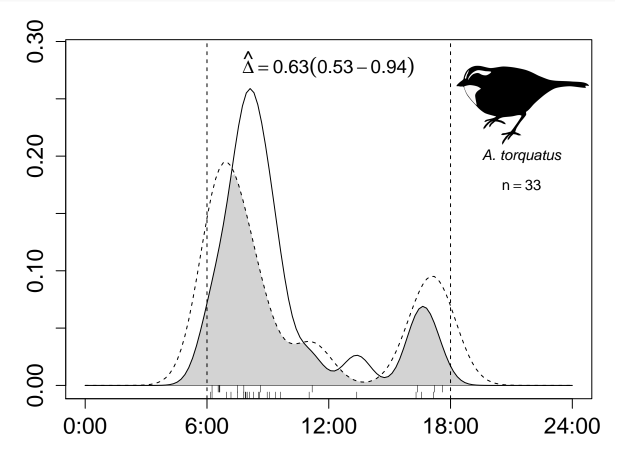
[1] 15 10000

```
plantboot <- resample(plant, 10000)
dim(plantboot)</pre>
```

[1] 34 10000

```
#overlap estimates by boostrap
BosqPlant <- bootEst(bosqboot, plantboot, adjust = c(1, NA, NA))</pre>
dim(BosqPlant)
## [1] 10000
                 3
BSmean <- colMeans(BosqPlant)</pre>
BSmean
                            Dhat5
##
       Dhat1
                 Dhat4
## 0.5905579
                    NA
                               NA
#calculation of CI
tmp <- BosqPlant[, 1]</pre>
bootCI(BosqPlantMILLE[1], tmp)
##
              lower
                         upper
## norm 0.3947913 0.7596248
## norm0 0.4014662 0.7662997
## basic 0.3987255 0.7650094
## basic0 0.3960816 0.7623656
## perc 0.4027565 0.7690404
tmp <- BosqPlant[, 1] # Extract the required column of the matrix</pre>
bootCIlogit(BosqPlantMILLE[1], tmp)
##
              lower
                        upper
## norm 0.3797421 0.7472527
## norm0 0.3896951 0.7551115
## basic 0.3715827 0.7448746
## basic0 0.3926846 0.7614875
## perc 0.4027565 0.7690404
#A. torquatus ----
#overlap calculation
bosq <- timeRad[chec$species == "torg" & chec$forest == "andean"]</pre>
plant <- timeRad[chec$species == "torq" & chec$forest == "reforestation"]</pre>
min(length(bosq), length(plant))
## [1] 11
BosqPlantMILLE <- overlapEst(bosq, plant)</pre>
BosqPlantMILLE
       Dhat1
                 Dhat4
                            Dhat5
## 0.6849370 0.6993380 0.9090909
```

```
#plot activity overlap
#png('a_torquatus1.png', width = 15, height = 10, units = 'cm', res = 300)
par(mar=c(2.8,4.1,1,1))
torq <- overlapPlot(bosq, plant, ylim=c(0,0.29), ylab="", cex.axis=1.4, cex.lab=1.4, cex.main= 1.4, xlarug = T, linecol = c("black", "black"), linewidth = c(1,1), main='', extend = NULL)
a <- readPNG(paste(getwd(), "/a_torquatus.png", sep = ""))
rasterImage(a, xleft=18.3, xright=24.8, ybottom=0.21, ytop=0.28)
abline(v=6, lty="dashed", col="black", lwd=1); abline(v=18, lty="dashed", col="black", lwd=1)
text(21.5,0.175,expression(n == 33), cex = 0.95);text(21.5,0.20,expression(paste(italic("A. torquatus"))
text(12,0.28,expression(hat(Delta) == 0.63 (0.53 - 0.94)), cex = 1.2)</pre>
```



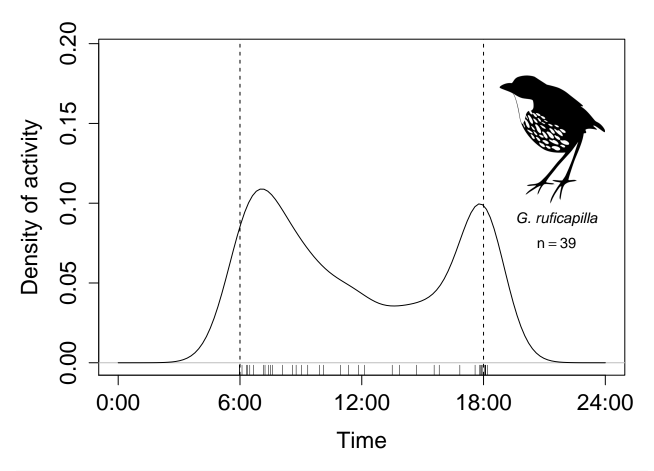
```
#dev.off()
#legend("topleft", c("Andean ", "G. ruficapilla"), lty=c(1,2), col=c(1,4), bty="n")
#bootstrap
bosqboot <- resample(bosq, 10000)
dim(bosqboot)</pre>
```

[1] 22 10000

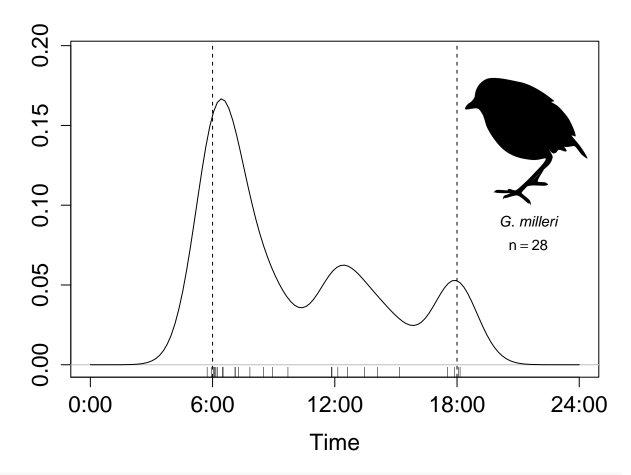
```
plantboot <- resample(plant, 10000)
dim(plantboot)</pre>
```

[1] 11 10000

```
#overlap estimates by boostrap
BosqPlant <- bootEst(bosqboot, plantboot, adjust = c(1, NA, NA))</pre>
dim(BosqPlant)
## [1] 10000
                 3
BSmean <- colMeans(BosqPlant)</pre>
##
      Dhat1
               Dhat4
                        Dhat5
## 0.645683
                NA
                           NA
#calculation of CI
tmp <- BosqPlant[, 1]</pre>
bootCI(BosqPlantMILLE[1], tmp)
##
              lower
                        upper
## norm 0.5132849 0.9350971
## norm0 0.4740309 0.8958431
## basic 0.5366065 0.9489145
## basic0 0.4602135 0.8725214
## perc 0.4209595 0.8332675
tmp <- BosqPlant[, 1] # Extract the required column of the matrix</pre>
bootCIlogit(BosqPlantMILLE[1], tmp)
##
              lower
                        upper
## norm 0.4869442 0.8687309
## norm0 0.4515388 0.8516453
## basic 0.4860398 0.8666831
## basic0 0.4559605 0.8521021
## perc 0.4209595 0.8332675
#G. ruficapilla ----
#overlap calculation
bosq <- timeRad[chec$species == "rufi"]</pre>
#plant <- timeRad[chec$species == "mille" & chec$forest == 2]</pre>
#min(length(bosq), length(plant))
#BosqPlantMILLE <- overlapEst(bosq, plant)</pre>
\#BosqPlantMILLE
#plot activity overlap
#png('g_ruficapilla1.png', height = 10, width = 15, units = 'cm', res = 300)
par(mar=c(4,4.1,1,1))
rufi <- densityPlot(bosq, ylim=c(0,0.195), ylab="Density of activity", cex.axis=1.4, cex.lab=1.4, cex.m
                     rug = T, linecol = c("black", "black"), linewidth = c(1,1), main='', extend = NULL
abline(v=6, lty="dashed", col="black", lwd=1); abline(v=18, lty="dashed", col="black", lwd=1)
a <- readPNG(paste(getwd(), "/g_ruficapilla.png", sep = ""))</pre>
rasterImage(a, xleft=18.8, xright=24, ybottom=0.10, ytop=0.18)
text(21.6,0.075,expression(n == 39), cex = 0.95);text(21.6,0.09,expression(paste(italic("G. ruficapilla
```



```
#dev.off()
#G. milleri ----
#overlap calculation
bosq <- timeRad[chec$species == "mille"]</pre>
#plant <- timeRad[chec$species == "mille" & chec$forest == 2]</pre>
#min(length(bosq), length(plant))
#BosqPlantMILLE <- overlapEst(bosq, plant)</pre>
#BosqPlantMILLE
#plot activity overlap
#png('g_milleri1.png', width = 15, height = 10, units = 'cm', res = 300)
par(mar=c(4,4,1,1))
mille <- densityPlot(bosq, ylim=c(0,0.195), ylab="", cex.axis=1.4, cex.lab=1.4, cex.main= 1.4, rug = T, linecol = c("black", "black"), linewidth = c(1,1), main='', extend = NULL)
abline(v=6, lty="dashed", col="black", lwd=1); abline(v=18, lty="dashed", col="black", lwd=1)
a <- readPNG(paste(getwd(), "/g_milleri.png", sep = ""))</pre>
rasterImage(a, xleft=18.4, xright=24.4, ybottom=0.10, ytop=0.18)
text(21.5,0.075,expression(n == 28), cex = 0.95);text(21.5,0.09,expression(paste(italic("G. milleri"))))
```



#dev.off()