Figure 5c

```
library(readxl)
library(lubridate)
##
## Attaching package: 'lubridate'
## The following objects are masked from 'package:base':
##
##
       date, intersect, setdiff, union
library(mgcv)
## Loading required package: nlme
## This is mgcv 1.8-34. For overview type 'help("mgcv-package")'.
library(ggplot2)
library(ggsignif)
library(readr)
library(emmeans)
library(tidyr)
library(plyr)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:plyr':
##
##
       arrange, count, desc, failwith, id, mutate, rename, summarise,
##
       summarize
## The following object is masked from 'package:nlme':
##
##
       collapse
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
#load in raw data
dat <- data.frame(read_csv("5c_data.csv"))</pre>
##
## -- Column specification -----
## cols(
     `Repeat(Date)` = col_datetime(format = ""),
##
##
     Algae = col_character(),
```

```
##
     Treatment = col_character(),
##
     Well_ID = col_character(),
##
     Larvae = col character(),
     Time_to_first_expulsion = col_double()
##
## )
head(dat)
     Repeat.Date. Algae Treatment Well_ID Larvae Time_to_first_expulsion
## 1
       2019-05-14 SSB01
                                        SC1
                                               1-1
                           control
       2019-05-14 SSB01
                                                                          95
                           control
                                        SC1
                                               1-2
## 3
       2019-05-14 SSB01
                                        SC1
                                               2-1
                                                                          5
                           control
## 4
      2019-05-14 SSB01
                          control
                                        SC1
                                               2-2
                                                                          95
## 5
      2019-05-14 SSB01
                           control
                                        SC1
                                               4-1
                                                                          95
## 6
       2019-05-14 SSB01
                           control
                                        SC1
                                               4-2
                                                                          95
#clean up raw data
dat$Algae <- factor(dat$Algae, levels= c("SSB01", "M. gaditana", "N. oculata", "C. velia"))
dat$Treatment <- factor(dat$Treatment)</pre>
dat$Repeat.Date. <- factor(as.character(dat$Repeat.Date.))</pre>
dat$expulsion <- ifelse(dat$Time_to_first_expulsion<90,1,0)</pre>
dat$Well_ID <- factor(dat$Well_ID)</pre>
head(dat)
##
     Repeat.Date. Algae Treatment Well_ID Larvae Time_to_first_expulsion expulsion
## 1
       2019-05-14 SSB01
                           control
                                        SC1
                                               1-1
## 2
       2019-05-14 SSB01
                                               1-2
                                                                          95
                                                                                     0
                                        SC1
                           control
## 3
       2019-05-14 SSB01
                           control
                                        SC1
                                               2-1
                                                                          5
                                                                                     1
                                                                          95
                                                                                     0
## 4
      2019-05-14 SSB01
                                        SC1
                                               2-2
                          control
## 5
       2019-05-14 SSB01
                           control
                                        SC1
                                               4-1
                                                                          95
                                                                                     0
## 6 2019-05-14 SSB01
                                        SC1
                           control
                                               4-2
                                                                          95
                                                                                     \cap
#model data with generalised linear mixed modes
summary <- data.frame()</pre>
for(i in unique(dat$Algae))
  #i <- unique(dat$Algae)[1]
 Dat <- droplevels(dat[dat$Algae==i,])</pre>
  ##Regression model:
  mod <- gam(expulsion ~ Treatment + s(Well_ID,bs='re'),data=Dat,family=binomial())</pre>
  S <- summary(mod)$p.table
  if(S[2,4]>0.1)
  {
   p <- ""
  if(S[2,4]<0.1)
  {
   p <- ""
  if(S[2,4]<0.05)
  {
       <- "*"
  if(S[2,4]<0.01)
```

```
p <- "**"
 }
  if(S[2,4]<0.001)
   p <- "***"
 newline <- data.frame(Algae=i,outcome=S[1,1],SE=S[1,2],type="control",p=NA,pvalue=NA)</pre>
  summary <- rbind(summary,newline)</pre>
 newline <- data.frame(Algae=i,outcome=S[1,1] + S[2,1],SE=S[2,2],type="LPS",p=p, pvalue=S[2,4])
  summary <- rbind(summary,newline)</pre>
}
summary$up <- plogis(summary$outcome + 1.96*summary$SE)</pre>
summary$down <- plogis(summary$outcome - 1.96*summary$SE)</pre>
summary$outcome <- plogis(summary$outcome)</pre>
summary
##
     Algae
              outcome
                              SF.
                                    type
                                                    pvalue
                                                                             down
                                            р
                                                                   up
## 1 SSB01 0.09999999 0.4714069 control <NA>
                                                         NA 0.2186985 0.04224192
## 2 SSB01 0.45238182 0.5642140
                                     LPS *** 0.0003769807 0.7139829 0.21468533
write_csv(summary, "5c_summary.csv")
summary$Algae <- factor(summary$Algae, levels= c("SSB01", "M. gaditana", "N. oculata", "C. velia"))</pre>
#Plot graph
P1 <- ggplot(data = summary,aes(x = Algae, y = outcome, group = type)) +
geom_bar(stat = "identity", width = 0.75, size=1.5, position = position_dodge(width = 0.8), fill = c(
geom_errorbar(aes(ymin=down, ymax=up), width=.5, position =position_dodge(.75)) +
geom_text(data = summary, aes(x=Algae,y=0.95),label=summary$p,size=6) +
theme_classic() +
labs(x = "", y = "Fraction expelled Algae") +
scale_y_continuous(expand = c(0, 0), limits = c(0,1), labels = scales::number_format(accuracy = 0.01))
plot(P1)
```

Warning: Removed 1 rows containing missing values (geom_text).

```
1.00

0.75

90

0.75

0.00

SSB01

ggsave("Figure_5c.pdf",width=5.75,height=11,units="cm")

## Warning: Removed 1 rows containing missing values (geom_text).
```