

Supplementary material 1: The R code for fitting multiple survival curves (data for multiple seed lots) and hypothesis testing.

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This supplementary material demonstrates how to analyze seed longevity in R to fit the viability equation. Original data is from Rezaei *et al.* (2023).

```
# load libraries
library(tidyverse) # usefull for all manor of data manipulation and visualizartion

## -- 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   3.5.1    v tibble    3.2.1
## v lubridate  1.9.3    v tidyr    1.3.1
## v purrr     1.0.2
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()   masks stats::lag()
## i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all conflicts to become errors

library(MASS) # the dose.p() function used to calculate P50 and P85

##
## Attaching package: 'MASS'
##
## The following object is masked from 'package:dplyr':
## 
##     select

library(knitr) # used to make tables in R Markdown

# set working directory
#setwd("my/file/path")

#
# Read in data
# brna short for the study organism, Brassica napus
brna <- read.csv("Rezaei_etal_2023_SS.R.csv", header = TRUE)
glimpse(brna)
```

```

## Rows: 117
## Columns: 10
## $ Treatment      <chr> "Desorption 170", "Desorption 170", "Desorption 170", "De-
## $ MC             <dbl> 9.141008, 9.141008, 9.141008, 9.141008, 8.000368, 8.00036-
## $ storage.RH    <dbl> 66.91, 66.91, 66.91, 66.91, 60.95, 60.95, 60.95, 60.95, 5-
## $ period         <int> 8, 17, 24, 27, 13, 24, 31, 41, 20, 37, 55, 90, 50, 59, 65-
## $ sown           <int> 90, 90, 90, 90, 87, 90, 90, 90, 90, 90, 90, 90, 90, 90, 9-
## $ germinated     <int> 89, 23, 3, 0, 83, 28, 10, 0, 84, 73, 14, 0, 73, 57, 51, 3-
## $ germ.percent   <dbl> 98.888889, 25.555556, 3.333333, 0.000000, 95.402299, 31.1-
## $ sorption        <chr> "Desorption", "Desorption", "Desorption", "Desorption", "~
## $ chamber.RH    <int> 70, 70, 70, 70, 60, 60, 60, 60, 50, 50, 50, 50, 40, 40, 4-
## $ cycle          <int> 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, ~

```

```

# citation for dataset;
# Rezaei, S., Buitink, J. & Hay, F.R. (2023) Contrasting seed moisture sorption
# behaviour between two species and the implication for seed longevity.
# Seed Science Research, 1-9. https://doi.org/10.1017/S0960258523000156

```

```
# Tell R which variables are factors
```

```

brna <- brna |>
  mutate(Treatment = as.factor(Treatment),
         sorption = as.factor(sorptio),
         chamber.RH = as.factor(chamber.RH),
         cycle = as.factor(cycle))
glimpse(brna)

```

```

## Rows: 117
## Columns: 10
## $ Treatment      <fct> Desorption 170, Desorption 170, Desorption 170, Desorptio-
## $ MC             <dbl> 9.141008, 9.141008, 9.141008, 9.141008, 8.000368, 8.00036-
## $ storage.RH    <dbl> 66.91, 66.91, 66.91, 66.91, 60.95, 60.95, 60.95, 60.95, 5-
## $ period         <int> 8, 17, 24, 27, 13, 24, 31, 41, 20, 37, 55, 90, 50, 59, 65-
## $ sown           <int> 90, 90, 90, 90, 87, 90, 90, 90, 90, 90, 90, 90, 90, 90, 9-
## $ germinated     <int> 89, 23, 3, 0, 83, 28, 10, 0, 84, 73, 14, 0, 73, 57, 51, 3-
## $ germ.percent   <dbl> 98.888889, 25.555556, 3.333333, 0.000000, 95.402299, 31.1-
## $ sorption        <fct> Desorption, Desorption, Desorption, Desorption, Desorptio-
## $ chamber.RH    <fct> 70, 70, 70, 70, 60, 60, 60, 60, 50, 50, 50, 50, 40, 40, 4-
## $ cycle          <fct> 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, ~

```

```
### analysis and plotting with three different Treatments ###
```

```

# subset desorption, cycle 1, and 3 RHs (30, 50, 70%)
des<-droplevels(subset(brna, sorption == "Desorption" & cycle == "1"))
des<-droplevels(subset(des, chamber.RH == "30" | chamber.RH == "50" | chamber.RH == "70"))

```

```
### Independent (independent Ki and -sigma^-1)
```

```

independent <- glm(formula = cbind(germinated, sown-germinated) ~ Treatment + Treatment:period -1 , fam-
summary(independent)

```

```

##
## Call:
## glm(formula = cbind(germinated, sown - germinated) ~ Treatment +
##       Treatment:period - 1, family = binomial(link = "probit"),

```

```

##      data = des)
##
## Coefficients:
##                               Estimate Std. Error z value Pr(>|z|)
## TreatmentDesorption 130      4.315944  0.281077 15.355 <2e-16 ***
## TreatmentDesorption 150      3.433957  0.344167  9.978 <2e-16 ***
## TreatmentDesorption 170      3.978369  0.427873  9.298 <2e-16 ***
## TreatmentDesorption 130:period -0.032867  0.002419 -13.585 <2e-16 ***
## TreatmentDesorption 150:period -0.077701  0.007848 -9.900 <2e-16 ***
## TreatmentDesorption 170:period -0.261960  0.025678 -10.202 <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 1086.437 on 14 degrees of freedom
## Residual deviance: 36.411 on 8 degrees of freedom
## AIC: 90.036
##
## Number of Fisher Scoring iterations: 6

```

#generating coefficient and confidence intervals for plotting

```
coefficients.for.plotting1<- cbind(summary(independent)$coef[,c(1)], confint(independent, level = 0.95))
```

Waiting for profiling to be done...

```
plot(des$period, des$germinated/des$sown, xlab="Time (days)", ylab="Proportion germinated", xlim=c(0,200),
curve(pnorm(coefficients.for.plotting1[1] + coefficients.for.plotting1[4]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting1[2] + coefficients.for.plotting1[5]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting1[3] + coefficients.for.plotting1[6]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=1,lwd=1, col =
```

If confidence intervals are to be shown see below code

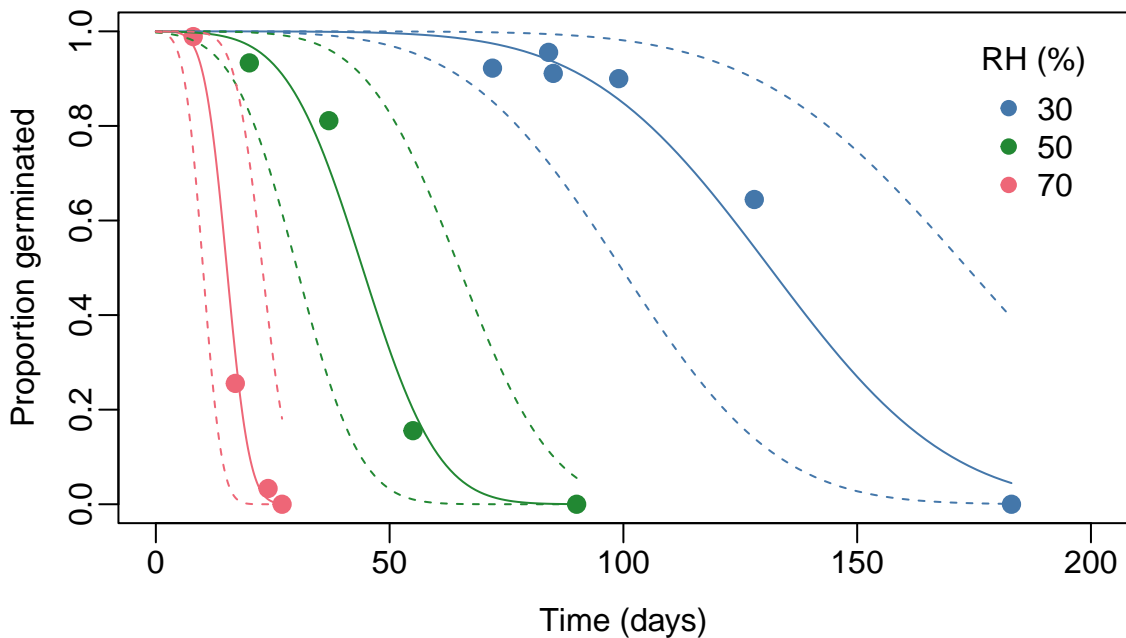
```
curve(pnorm(coefficients.for.plotting1[1,2] + coefficients.for.plotting1[4,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting1[1,3] + coefficients.for.plotting1[4,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =
```

```
curve(pnorm(coefficients.for.plotting1[2,2] + coefficients.for.plotting1[5,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting1[2,3] + coefficients.for.plotting1[5,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=2,lwd=1, col =
```

```
curve(pnorm(coefficients.for.plotting1[3,2] + coefficients.for.plotting1[6,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting1[3,3] + coefficients.for.plotting1[6,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =
```

```
legend(175, 1, title = "RH (%)", rev(unique(des$chamber.RH)),col=c("#4477AA","#228833","#EE6677"), pch=
title("Independent")
```

Independent



```
#dev.off()

###Common intercept (Common Ki)
com.inter <- glm(formula = cbind(germinated, sown-germinated) ~ Treatment:period , family = binomial(link = "probit"))
summary(com.inter)

##
## Call:
## glm(formula = cbind(germinated, sown - germinated) ~ Treatment:period,
##       family = binomial(link = "probit"), data = des)
##
## Coefficients:
##                               Estimate Std. Error z value Pr(>|z|)
## (Intercept)                3.981964   0.193851  20.54  <2e-16 ***
## TreatmentDesorption 130:period -0.030063   0.001709 -17.59  <2e-16 ***
## TreatmentDesorption 150:period -0.089767   0.004793 -18.73  <2e-16 ***
## TreatmentDesorption 170:period -0.262164   0.013179 -19.89  <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 1079.71  on 13  degrees of freedom
## Residual deviance: 40.39  on 10  degrees of freedom
```

```

## AIC: 90.015
##
## Number of Fisher Scoring iterations: 6

# generating coefficient and confidence intervals for plotting
coefficients.for.plotting2<- cbind(summary(com.inter)$coef[,c(1)], confint(com.inter, level = 0.95)) #

## Waiting for profiling to be done...

# plot
plot(des$period, des$germinated/des$sown, xlab="Time (days)", ylab="Porportion germinated", xlim=c(0,200))

curve(pnorm(coefficients.for.plotting2[1] + coefficients.for.plotting2[2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting2[1] + coefficients.for.plotting2[3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting2[1] + coefficients.for.plotting2[4]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=1,lwd=1, col =

### If confidence intervals are to be shown see below code ###

curve(pnorm(coefficients.for.plotting2[1,2] + coefficients.for.plotting2[2,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting2[1,3] + coefficients.for.plotting2[2,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =

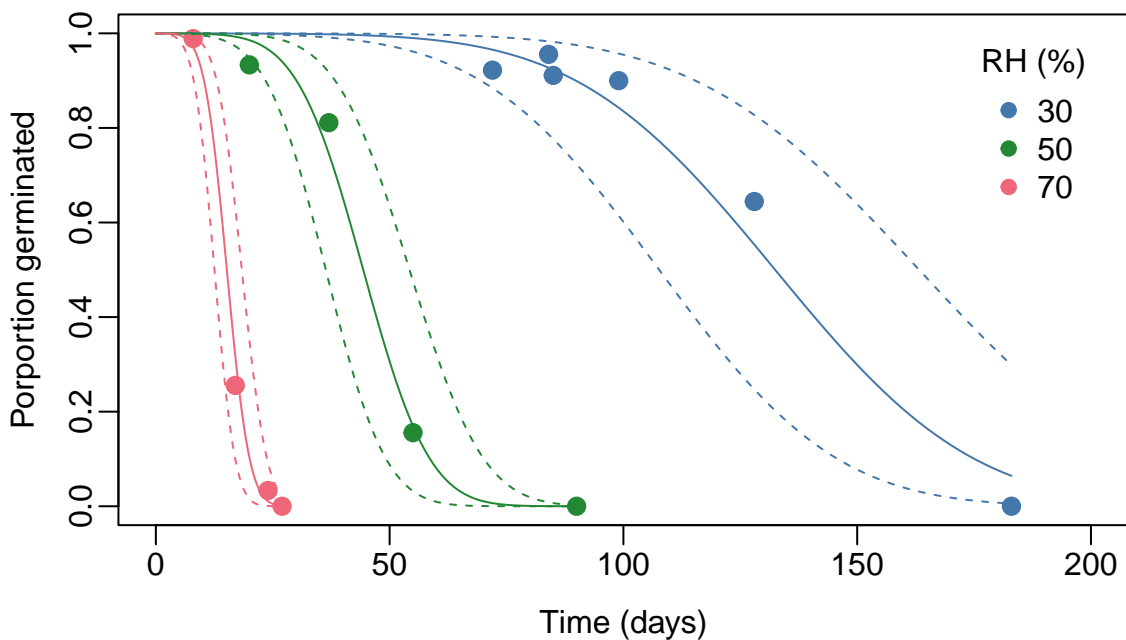
curve(pnorm(coefficients.for.plotting2[1,2] + coefficients.for.plotting2[3,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period),
      add=TRUE, lty=2,lwd=1, col = "#228833")
curve(pnorm(coefficients.for.plotting2[1,3] + coefficients.for.plotting2[3,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=2,lwd=1, col =

curve(pnorm(coefficients.for.plotting2[1,2] + coefficients.for.plotting2[4,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting2[1,3] + coefficients.for.plotting2[4,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =

legend(175, 1, title = "RH (%)", rev(unique(des$chamber.RH)),col=c("#4477AA","#228833","#EE6677"), pch=
title("Common intercept")

```

Common intercept



```
#dev.off()

# Table of coefficients for common intercept

###Common slope (common -sigma^-1)
com.slope <- glm(formula = cbind(germinated, sown-germinated) ~ Treatment + period -1, family = binomial)
summary(com.slope)

## 
## Call:
## glm(formula = cbind(germinated, sown - germinated) ~ Treatment +
##       period - 1, family = binomial(link = "probit"), data = des)
## 
## Coefficients:
##                               Estimate Std. Error z value Pr(>|z|)    
## TreatmentDesorption 130  6.24593   0.35392 17.648 < 2e-16 ***
## TreatmentDesorption 150  2.31196   0.15725 14.702 < 2e-16 ***
## TreatmentDesorption 170  0.43695   0.09078  4.813 1.48e-06 ***
## period                  -0.05077   0.00312 -16.270 < 2e-16 ***
## ---                     
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## 
```

```

## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 1086.44 on 14 degrees of freedom
## Residual deviance: 255.08 on 10 degrees of freedom
## AIC: 304.71
##
## Number of Fisher Scoring iterations: 6

#generating coefficient and confidence intervals for plotting
coefficients.for.plotting3<- cbind(summary(com.slope)$coef[,c(1)], confint(com.slope, level = 0.95)) #

## Waiting for profiling to be done...

# plot
plot(des$period, des$germinated/des$sown, xlab="Time (days)", ylab="Porportion germinated", xlim=c(0,200))

curve(pnorm(coefficients.for.plotting3[1] + coefficients.for.plotting3[4]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[2] + coefficients.for.plotting3[4]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=1,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[3] + coefficients.for.plotting3[4]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=1,lwd=1, col =

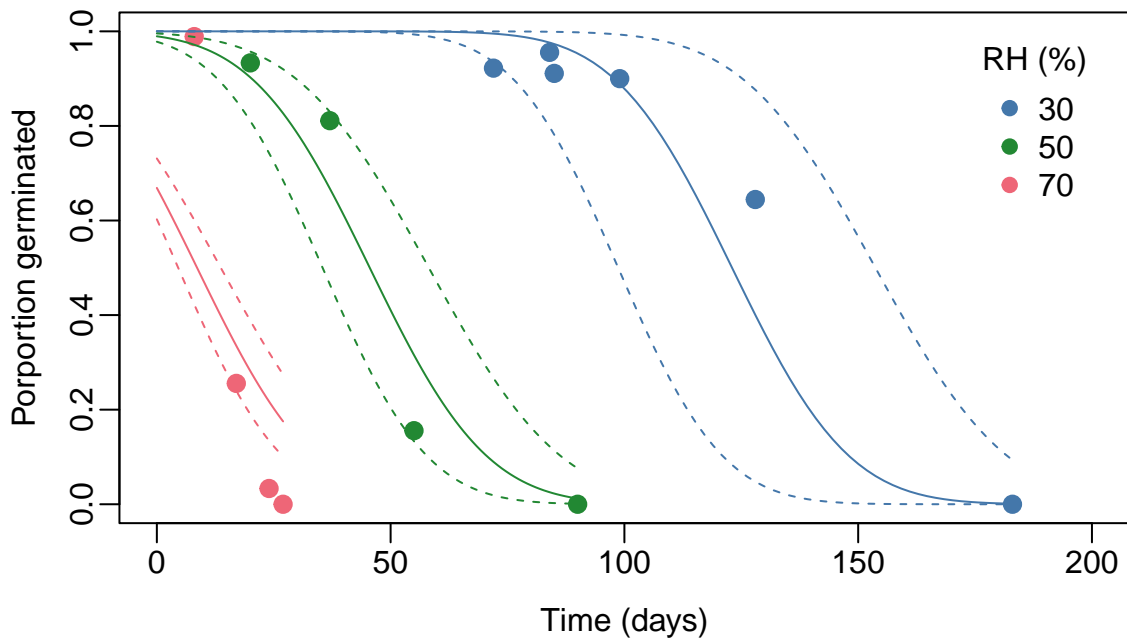
### If confidence intervals are to be shown see below code ###

curve(pnorm(coefficients.for.plotting3[1,2] + coefficients.for.plotting3[4,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[1,3] + coefficients.for.plotting3[4,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 130")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[2,2] + coefficients.for.plotting3[4,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period),
      add=TRUE, lty=2,lwd=1, col = "#228833")
curve(pnorm(coefficients.for.plotting3[2,3] + coefficients.for.plotting3[4,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 150")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[3,2] + coefficients.for.plotting3[4,2]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =
curve(pnorm(coefficients.for.plotting3[3,3] + coefficients.for.plotting3[4,3]*x),
      from = 0, to = max(subset(des, Treatment=="Desorption 170")$period), add=TRUE, lty=2,lwd=1, col =

legend(175, 1, title = "RH (%)", rev(unique(des$chamber.RH)),col=c("#4477AA","#228833","#EE6677"), pch=
title("Common slope"))

```

Common slope



```
#dev.off()
```

```
## one line
one.line <- glm(formula = cbind(germinated, sown-germinated) ~ period , family = binomial(link = "probit"), data = des)
summary(one.line)

##
## Call:
## glm(formula = cbind(germinated, sown - germinated) ~ period,
##      family = binomial(link = "probit"), data = des)
##
## Coefficients:
##             Estimate Std. Error z value Pr(>|z|)
## (Intercept)  0.262001   0.060829  4.307 1.65e-05 ***
## period       -0.002573   0.000743 -3.464 0.000533 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 1079.7 on 13 degrees of freedom
## Residual deviance: 1067.8 on 12 degrees of freedom
## AIC: 1113.4
##
## Number of Fisher Scoring iterations: 5
```

```

#generating coefficient and confidence intervals for plotting
coefficients.for.plotting4<- cbind(summary(one.line)$coef[,c(1)], confint(one.line, level = 0.95)) # le

## Waiting for profiling to be done...

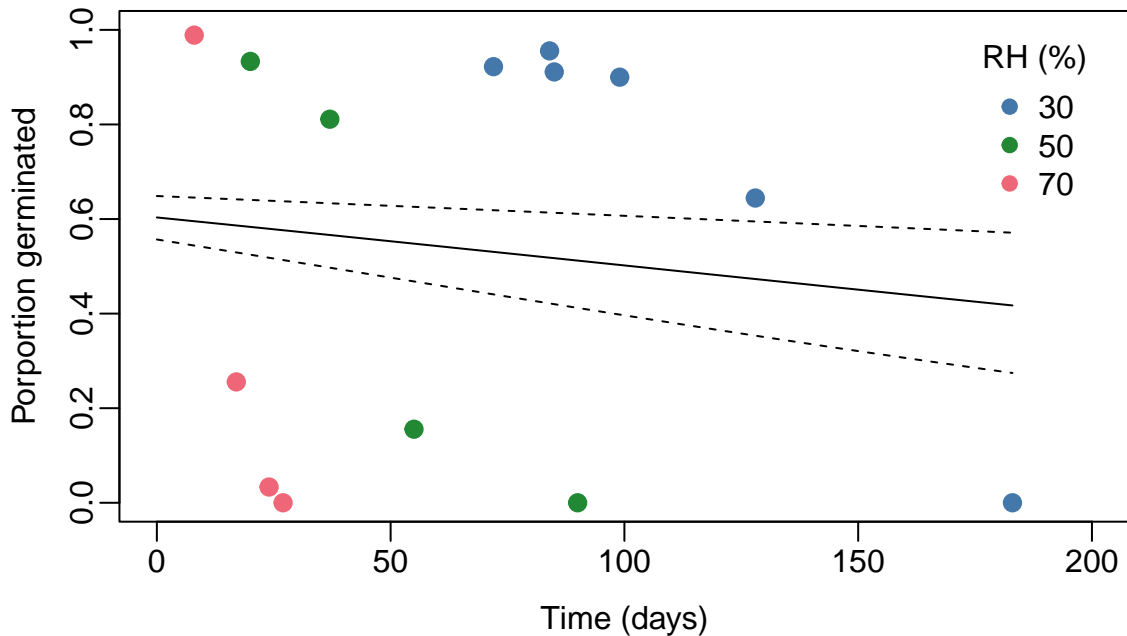
# plot
plot(des$period, des$germinated/des$sown, xlab="Time (days)", ylab="Porportion germinated", xlim=c(0,200),
      curve(pnorm(coefficients.for.plotting4[1] + coefficients.for.plotting4[2]*x),
             from = 0, to = max(des$period), add=TRUE, lty=1,lwd=1)

### If confidence intervals are to be shown see below code ####
curve(pnorm(coefficients.for.plotting4[1,2] + coefficients.for.plotting4[2,2]*x),
       from = 0, to = max(des$period), add=TRUE, lty=2,lwd=1)
curve(pnorm(coefficients.for.plotting4[1,3] + coefficients.for.plotting4[2,3]*x),
       from = 0, to = max(des$period), add=TRUE, lty=2,lwd=1)

legend(175, 1, title = "RH (%)", rev(unique(des$chamber.RH)),col=c("#4477AA","#228833","#EE6677"), pch=16,
      title="One line")

```

One line



```

#dev.off()

# Table of coefficients for one line model
table.oneline<- signif(as.data.frame(summary(one.line)$coefficients[,c(1,2,4)]), 4)

```

```

table.oneline[2,1] <- signif(-1/table.oneline[2,1], 4)

# dose.p
table.oneline[3,1] <- signif(dose.p(one.line, cf = 1:2, p = 0.5), 4)
table.oneline[3,2] <- signif(unname(attributes(dose.p(one.line, cf = 1:2, p = 0.5))$SE[, 1]), 4)
table.oneline[4,1] <- signif(dose.p(one.line, cf = 1:2, p = 0.85), 4)
table.oneline[4,2] <- signif(unname(attributes(dose.p(one.line, cf = 1:2, p = 0.85))$SE[, 1]), 4)

row.names(table.oneline) <- c("Ki", "sigma", "p50", "p85")
table.oneline$`Std. Error` <- round(table.oneline$`Std. Error`, 4)

table.oneline[3,3] <- "-"
table.oneline[4,3] <- "-"
table.oneline$AIC <- "-"
table.oneline$AIC[1] <- round(summary(one.line)$aic, 4)

# Call the table
knitr::kable(table.oneline)

```

	Estimate	Std. Error	Pr(> z)	AIC
Ki	0.262	0.0608	1.653e-05	1113.4349
sigma	388.700	0.0007	0.000533	-
p50	101.800	17.1400	-	-
p85	-301.000	107.0000	-	-

```

### Compare models ###

# Create deviance table
dev.table<-data.frame(
  Model = c("independent", "Common.intercept", "Common.slope", "One.line"),
  Deviance = c(independent$deviance, com.inter$deviance, com.slope$deviance, one.line$deviance),
  df = c(independent$df.residual, com.inter$df.residual, com.slope$df.residual, one.line$df.residual),
  AIC = c(independent$aic, com.inter$aic, com.slope$aic, one.line$aic)
)

# Call the table.
knitr::kable(dev.table, row.names = FALSE)

```

Model	Deviance	df	AIC
independent	36.41096	8	90.03587
Common.intercept	40.39037	10	90.01528
Common.slope	255.08134	10	304.70624
One.line	1067.81000	12	1113.43490

Here we demonstrate pairwise comparisons of common slope and common intercept with independent and one.line.
However, in practice this would only be done with the models that have the lowest and closest in their
(e.g. independent and common intercept in this data presented).

Common intercept vs Independent

```

df.change1<-com.inter$df.residual - independent$df.residual
df.change1

## [1] 2

F1<-((com.inter$deviance-independent$deviance)/df.change1)/(independent$deviance/independent$df.residual)
F1

## [1] 0.437166

F1prob<-pf(F1, df.change1, independent$df.residual, lower.tail = FALSE) # note the lower.tail = False is important
F1prob<-format(signif(F1prob, 4), nsmall = 4)
F1prob

## [1] "0.6604"

F1sum<-c(df.change1, F1, F1prob)
F1sum

## [1] "2"           "0.437166049341884" "0.6604"

# Common slope vs Independent
df.change2<-com.slope$df.residual - independent$df.residual
df.change2

## [1] 2

F2<-((com.slope$deviance-independent$deviance)/df.change2)/(independent$deviance/independent$df.residual)
F2

## [1] 24.02248

F2prob<-pf(F2, df.change2, independent$df.residual, lower.tail = FALSE) # note the lower.tail = False is important
F2prob<-format(signif(F2prob, 4), nsmall = 4)
F2prob

## [1] "0.0004152"

F2sum<-c(df.change2, F2, F2prob)
F2sum

## [1] "2"           "24.0224765318435" "0.0004152"

# Common intercept vs One line
df.change3<-one.line$df.residual - com.inter$df.residual
df.change3

## [1] 2

```

```

F3<-((one.line$deviance-com.inter$deviance)/df.change3)/(com.inter$deviance/com.inter$df.residual)
F3

## [1] 127.1862

F3prob<-pf(F3, df.change3, one.line$df.residual, lower.tail = FALSE) # note the lower.tail = False is n
F3prob<-format(signif(F3prob, 4), nsmall = 4)
F3prob

## [1] "8.359e-09"

F3sum<-c(df.change3, F3, F3prob)
F3sum

## [1] "2"           "127.18620508823" "8.359e-09"

# Common slope vs One line
df.change4<-one.line$df.residual - com.slope$df.residual
df.change4

## [1] 2

F4<-((one.line$deviance-com.slope$deviance)/df.change4)/(com.slope$deviance/com.slope$df.residual)
F4

## [1] 15.93077

F4prob<-pf(F4, df.change4, com.slope$df.residual, lower.tail = FALSE) # note the lower.tail = False is n
F4prob<-format(signif(F4prob, 4), nsmall = 4)
F4prob

## [1] "0.0007779"

F4sum<-c(df.change4, F4, F4prob)
F4sum

## [1] "2"           "15.9307746494971" "0.0007779"

# Combine into one summary table. Report in appendices
# Most important value is F probability
# If F probability > alpha, you can accept the model
# In this case there's no significant difference between common intercept vs independent
F.table1<-data.frame(Model.comparison = c("Common intercept vs. Independent",
                                             "Common slope vs. Independent",
                                             "One line vs. Common intercept",
                                             "One line vs. Common slope"),
                        "DF change" = c(df.change1, df.change2, df.change3, df.change4),
                        "F statistic" = c(F1, F2, F3, F4),
                        "F probability" = c(F1prob, F2prob, F3prob, F4prob))
# call the table
knitr::kable(F.table1)

```

Model.comparison	DF.change	F.statistic	F.probability
Common intercept vs. Independent	2	0.437166	0.6604
Common slope vs. Independent	2	24.022476	0.0004152
One line vs. Common intercept	2	127.186205	8.359e-09
One line vs. Common slope	2	15.930775	0.0007779

```

rm(df.change1, df.change2, df.change3, df.change4, F1, F2, F3, F4, F1prob, F2prob, F3prob, F4prob, F1sum)
# N.B. Common line = one line
# N.B. Independent

# Note: A quick way to view the deviance between two models is with function anova()
# adding, test = test="Chisq", will output a p-value.
anova(independent, com.inter, test="Chisq")

## Analysis of Deviance Table
##
## Model 1: cbind(germinated, sown - germinated) ~ Treatment + Treatment:period -
##           1
## Model 2: cbind(germinated, sown - germinated) ~ Treatment:period
##   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1      8     36.411
## 2     10     40.390 -2    -3.9794   0.1367

#### Table of coefficients
### Presented for only the selected model. In this case the independent model.

table.independent<- round(as.data.frame(summary(independent)$coefficients[,c(1,2,4)]), 3)
table.independent[c(4:6),1]<- signif(-1/table.independent[c(4:6),1], 3)
table.independent[c(4:6),2] <- table.independent[ c(4:6),2]*table.independent[ c(4:6),1]

# dose.p
table.independent[7,1]<- signif(dose.p(independent, cf = c(1,4), p = 0.5), 3)
table.independent[7,2]<- signif(unname(attributes(dose.p(independent, cf = c(1,4), p = 0.5))$SE[, 1]), 3)
table.independent[8,1]<- signif(dose.p(independent, cf = c(2,5), p = 0.5), 3)
table.independent[8,2]<- signif(unname(attributes(dose.p(independent, cf = c(2,5), p = 0.5))$SE[, 1]), 3)
table.independent[9,1]<- signif(dose.p(independent, cf = c(3,6), p = 0.5), 3)
table.independent[9,2]<- signif(unname(attributes(dose.p(independent, cf = c(3,6), p = 0.5))$SE[, 1]), 3)

row.names(table.independent)<-c( "Ki-Desorption 130", "Ki-Desorption 150", "Ki-Desorption 170", "sigma-Desorption")
table.independent$`Std. Error` <- round(table.independent$`Std. Error`, 3)

table.independent[c(7:9),3] <- "-"
table.independent$AIC<- "-"
table.independent$AIC[1]<- round(summary(independent)$aic, 3)
names(table.independent)<- c("Estimate", "SE", "P", "AIC")
table.independent$P <- ifelse(table.independent$P=="0", "<0.001", table.independent$P)

# Call the table.
knitr::kable(table.independent)

```

	Estimate	SE	P	AIC
Ki-Desorption 130	4.316	0.281	<0.001	90.036
Ki-Desorption 150	3.434	0.344	<0.001	-
Ki-Desorption 170	3.978	0.428	<0.001	-
sigma-Desorption 130	30.300	0.061	<0.001	-
sigma-Desorption 150	12.800	0.102	<0.001	-
sigma-Desorption 170	3.820	0.099	<0.001	-
p50-Desorption 130	131.000	2.730	-	-
p50-Desorption 150	44.200	1.250	-	-
p50-Desorption 170	15.200	0.449	-	-