# Summarizing HTP Temporal Data

singhdj2@ksu.edu August 8, 2019

### Introduction

The objective of today's exercise is to make you familiar with the processing of high-throughput phenotyping data. We will use plant height trait as an example and along the way will learn how to summarize and explore data. Some of the concepts/topics discussed here would be applicable to your research as well.

#### About the Data

The data was collected from two wheat experiments in South Asia in field season 2018 using an unmanned aerial vehicle a.k.a. drone. There were a total of 1200 wheat test plots (size 1.2m x 3m) in each experiment. The two experiments are identical in terms of genotypes and plot layout except these were planted at two different sowing dates (Optimum on 23-Oct, Late on 11 Nov). The main objective of this field experiment was to assess the effect of sowing dates on plant growth and yield. The data was collected at multiple times during the wheat growing season. Overall this dataset captures the dynamic changes in wheat height during its growth cycle.

## **Expected Outcome**

Through this exercise you will learn:

- How to process plot-level height data from multiple dates
- Model the growth dynamics of height through non-linear regression
- Visualize the results and make sense of the trends
- Compare the effects of sowing dates on wheat height and growth

#### First we will load all the required packages

First, let us create a logistic regression function to model wheat growth. This is just the definition of a function, we will call it during the later steps.

Quick question: Why is there any need to create functions?

Non-linear Logistic Regression Function

```
nest()
  res <- data.frame(phi1=NA, phi2=NA, phi3=NA,
                    flag="no-data", stringsAsFactors = F)
 fit.stats <- as.data.frame(matrix(nrow = 1,ncol = 11,
                                    dimnames = list(1,fit.stat.cols))) # empty fit.stats
 pred.res <- bind cols(pred,res,fit.stats)</pre>
                                                 # bind columns
 return(pred.res)
                                       # return the data frame
if(max(phenValue, na.rm=TRUE) < 60){  ## skip plots that never reach 60cm height</pre>
 pred = data.frame(x=NA,y=NA) %>%
   nest()
 res <- data.frame(phi1=NA, phi2=NA, phi3=NA, flag="short.max.ht",
                    stringsAsFactors = F) # flag short plots
 fit.stats <- as.data.frame(matrix(nrow = 1,ncol = 11,</pre>
                                    dimnames = list(1,fit.stat.cols))) # empty fit.stats
 pred.res <- bind_cols(pred,res,fit.stats)</pre>
  return(pred.res)
## get Thermal Time
days = dates
if(sum(!is.na(phenValue)) >= 9) { # run only plots that have at least 9 time-points
  # add some dummy variables for min and max height at 50-100 ThermalDays before/after the phenotypin
  # buffer raw data to force start values to zero
 days = c(days, c(0,50,75,100), max(days,na.rm=T)+c(50,75,100,150))
 height = c(phenValue, c(0,0,0,0), rep(max(phenValue,na.rm=T),4))
                                                                    #c(0,0,0,100,100,100))
 ##find initial starting values for phi2 and phi3, fix phi1 at max pheno value
 phi1 = max(phenValue,na.rm=T)
                                                     # maximum height parameter
 phi2 = coef(lm(logit(height/phi1)~days))[2]
                                                     # growth rate
 phi3 = -coef(lm(logit(height/phi1)~days))[1]/phi2 # time at half-max growth
  # Run logistic regression model
 growth_mod <- try(nls(height~ max(phenValue,na.rm=T)/(1+exp(-phi2*(days - phi3))),</pre>
                        start=list(phi2=phi2,phi3=phi3), trace=FALSE), silent=TRUE)
  if(class(growth_mod)!="try-error"){
                                                 # this condition is to handle errors
   phi2 = coef(growth_mod)[1]
   phi3 = coef(growth_mod)[2]
    ## get predicted value
   pred = getPred(phi1, phi2, phi3) %>%
    res = data.frame(phi1, phi2, phi3,flag="data.complete",
                     stringsAsFactors = F) # combine all growth coeficients
   fit.stats <- accuracy(list(growth_mod),plotit = F)$Fit.criteria # gives pseudo-Rsq and other fit
    colnames(fit.stats) <- fit.stat.cols</pre>
   pred.res = bind_cols(pred,res,fit.stats)
                                                  # bind growth and predicted values
   return(pred.res)
  } else {
                                                   # skip if doesn't converge
   pred = data.frame(x=NA,y=NA) %>%
     nest()
    res <- data.frame(phi1=NA,phi2=NA,phi3=NA,flag='try-error',
                      stringsAsFactors = F)
   fit.stats <- as.data.frame(matrix(nrow = 1,ncol = 11,dimnames = list(1,fit.stat.cols))) # empty f
    pred.res <- bind_cols(pred,res,fit.stats)</pre>
   return(pred.res)
```

Helper function to extract predictions for desired input 'x' values (e.g. days, thermal days).

Now lets get started with our analysis by reading the data file. Enter help(read.csv) in the console to know about the read.csv function.

```
# read curated data UAV height data
dat.ht <- read.csv("C:/Users/singhdj2/Documents/daljit/r_stuff/tutorials/data/18LDH-tutorial-log-regress
head(dat.ht)

## condition plot_id thermal.time DAS phenotype_value</pre>
```

```
## 1
      Optimum
                4120
                            1101 78
                                           48.64035
## 2
      Optimum
                4119
                            1101 78
                                           53.21288
## 3
     Optimum
                4118
                            1101 78
                                           41.98752
     Optimum
## 4
                4117
                            1101 78
                                           47.62004
## 5
     Optimum
                            1101 78
                                           47.49351
                4116
## 6
     Optimum
                4115
                            1101 78
                                           42.04319
```

There are five columns in the data file. Column *condition* refers to the optimum and late sowing conditions; *plot\_id* is unique plot\_id for 1200 plots in each condition; *thermal.time* is the cumulative sum of the mean daily air temperature at each condition; DAS is Days after sowing (in Julian Day units); *phenotype\_value* is the height value estimated from drone based images.

Before moving forward with your analysis, it is always good to take a look at the data structure and column attributes. Here we will take a quick glimpse at our data file.

Let us find out how many unique time-points are there per condition?

In which condition the plants grow taller? Lets get mean height per time-point for each condition to find out.

```
dat.ht %>%
  group_by(condition,thermal.time) %>%
  summarise(ht.mean=mean(phenotype_value,na.rm=T))

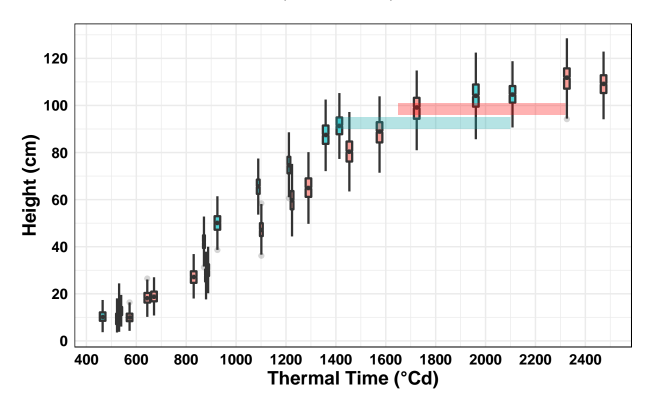
## # A tibble: 26 x 3
## # Groups: condition [2]
```

```
##
     condition thermal.time ht.mean
               <int>
##
     <fct>
                             <dbl>
## 1 Optimum
                            12.7
                       546
## 2 Optimum
                       573
                             9.98
## 3 Optimum
                       644
                            18.2
## 4 Optimum
                       671
                             18.8
## 5 Optimum
                       830
                             27.1
## 6 Optimum
                             27.7
                       881
## 7 Optimum
                       898
                             30.1
## 8 Optimum
                             47.3
                       1101
## 9 Optimum
                       1226
                             59.8
                       1290
                             65.1
## 10 Optimum
## # ... with 16 more rows
```

We can dig a little deeper to look at the raw height trends at both field conditions.

```
x.label = "Thermal Time (°Cd)"
ggplot(data=dat.ht, aes(y = phenotype_value,
                        x = thermal.time,
                        group=interaction(thermal.time,condition),
                        fill=condition)) +
  geom_boxplot(width=25,alpha=0.7,notch = TRUE, size=0.8,
               notchwidth = 0.5, outlier.alpha = 0.2) +
  labs(x=x.label,y="Height (cm)") +
  scale_y_continuous(breaks= seq(0,150,20)) +
  scale_x_continuous(breaks= seq(0,2400,200)) +
  theme_minimal() +
  theme(panel.border = element_rect(colour = "black", fill = NA, size = .5),
        axis.text.x = element_text(colour = "black", size = 11, angle = 0),
        axis.text.y = element_text(colour = "black", size = 11),
       plot.title = element_text(size = 14, face = "bold"),
        text = element_text(size=14,face="bold"),
       legend.position="top") +
  guides(fill=guide_legend(title="")) +
  annotate("rect", xmin=1650, xmax=2320, ymin=96, ymax=101, alpha=0.3, fill="red") +
  annotate("rect", xmin=1400, xmax=2100, ymin=90, ymax=95, alpha=0.3, fill="#0da3a7")
```





```
# We can also save the above plot by using ggsave function.
# create a file name character string
# fName = pasteO('../data/18LDH_tutorial-height-trend_boxplot_',Sys.Date(),'.jpg')
# # save the plot on disk
# ggsave(fName, plot = last_plot(), device = "jpg", path = "output/",
# scale = 1, width = 10, height = 6.5, units = c("in"), dpi=800)
```

Question: By chance, did you notice different height trends at two conditions? Which condition has plants growing faster/slower?

We have data collected at multuple time-points. This rich temporal information can be summarized into a few growth parameters that have a straightforward biological interpretation. To do so, we will deploy our logistic regression function (remember we defined logistic function at the beginning of this script!) for each plot and condition using group\_by and do functions from tidyverse package.

```
dat.ht.growth <- dat.ht %>%
  group_by(condition,plot_id) %>%
  do(runLogReg(dates = .$thermal.time, phenValue = .$phenotype_value)) %>%
  ungroup()
head(dat.ht.growth)
```

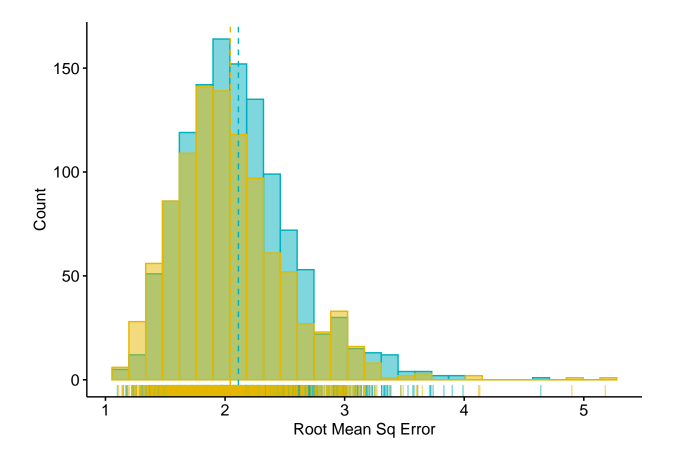
## # A tibble: 6 x 18

```
##
    condition plot_id data    phi1
                                    phi2 phi3 flag Min.max.accuracy
              <int> <lis> <dbl>
##
    <fct>
                                    <dbl> <dbl> <chr>
                                                                 <dbl> <dbl>
                1001 <tib~ 95.4 0.00357 1186. data~
                                                                0.799 1.13
## 1 Optimum
## 2 Optimum
                1002 <tib~ 109. 0.00381 1245. data~
                                                                0.782 1.37
                 1003 <tib~ 107. 0.00333 1135. data~
## 3 Optimum
                                                                0.793 1.81
## 4 Optimum
                 1004 <tib~ 104. 0.00340 1266. data~
                                                                0.782 1.65
                 1005 <tib~ 106. 0.00334 1197. data~
## 5 Optimum
                                                                0.786 1.79
                 1006 <tib~ 112. 0.00309 1204. data~
## 6 Optimum
                                                                0.786 2.35
## # ... with 9 more variables: MAPE <dbl>, MSE <dbl>, RMSE <dbl>,
      NRMSE.mean <dbl>, NRMSE.median <dbl>, NRMSE.mean.accuracy <dbl>,
      NRMSE.median.accuracy <dbl>, Efron.r.squared <dbl>, CV.prcnt <dbl>
```

Note that the parameters phi1, phi2, phi3 correspond to the maximum plant height (cm), growth rate (cm/thermal time), thermal time when plant is halfway its maximum height (measured in degree Cd units), respectively.

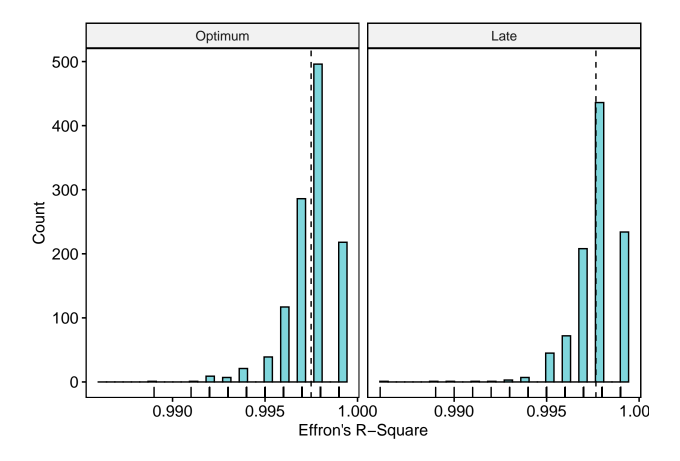
Now that we have succefully summarized our time-points into three biologically interpretable parameters, we can go ahead and take a look at the regression fit summaries to find out how accurately our model was able to fit the original data.

```
gghistogram(dat.ht.growth, x = "RMSE",
  add = "mean", rug = T,
  color = "condition", fill = "condition",
  palette = c("#00AFBB", "#E7B800")) +
  rremove("legend") +
  xlab("Root Mean Sq Error") +
  ylab("Count")
```



Keep in mind that the R-square statistic is a bit tricky for non-linear models. Therefore, Efron.r-sq is used as a workaround approximation for checking the model fits in non-linear models. We will plot Efron.r-sq to check the fit of our regression model.

```
gghistogram(dat.ht.growth, x = "Efron.r.squared",
  add = "mean", rug = T,
  #color = "condition",
  fill = "#00AFBB",
  facet.by = "condition") +
  rremove("legend") +
  xlab("Effron's R-Square") +
  ylab("Count")
```



It looks like our logistic model fit our data really well. Almost 99% of the data had a perfect fit.

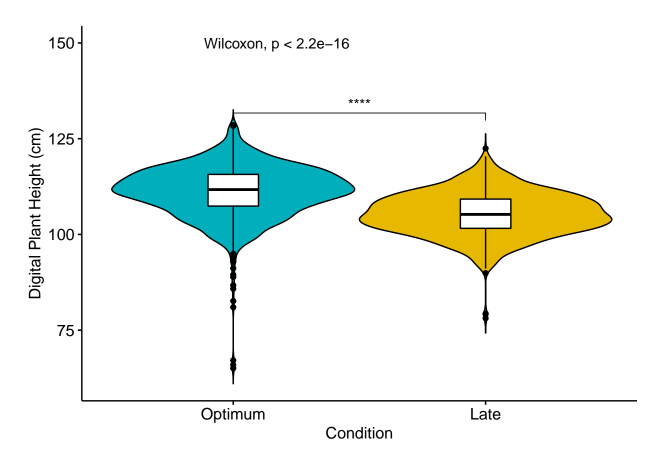
Now we are quite sure that our data fits worked well. Now we can ask some biological questions, for example, does sowing date have any effect on plant growth? Lets dig further...

Let us check mean values for each growth parameter.

Looks like the late sown plants are gowing much faster in short time!

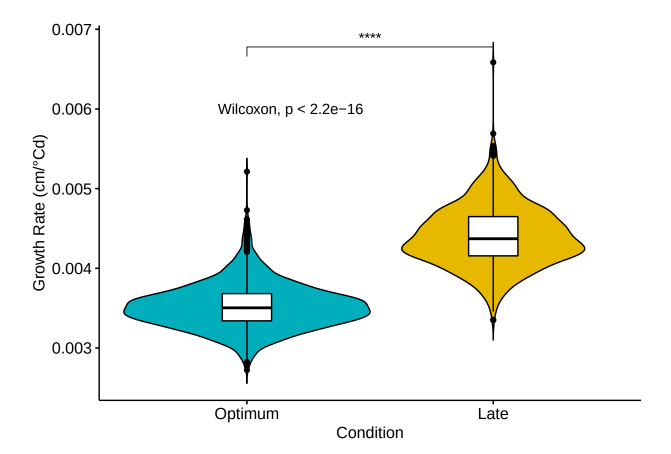
Wait...but, are these mean values statistically significant?

We can create violin plots and also perform mean comparisons for phi1 (maximum plant height) in a single step.



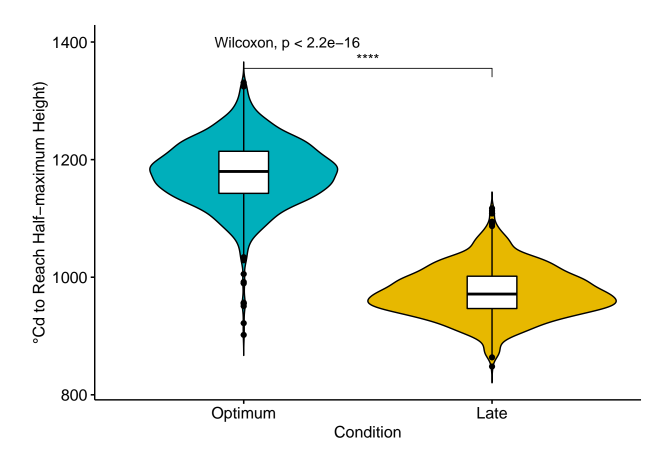
What do you see in these plots? Does this figure answer our question of sowing date effect on plant height?

Violin plots for phi2 (height growth rate)



Does the late planted crop grow faster? If so, any guess why?

Violin plots for phi3 (time to reach half maximum height).



This plot is also consistent with what we see in previous two plots, i.e., the late sown crop grows significantly faster and take less time to reach to its half maximum height.

## Concluding remarks

Through this tutorial you have familiarized yourself with a number of programming and statistical concepts namely: R functions, conditional workflows (if, else), tidy-data, exploratory data analysis, graphics, non-linear regression modeling, mean comparisons etc. However, always remeber that the tutorials such this one can only provide a primer or starting point- the real learning actually comes from a goal-oriented program exercises. Just like any other skill, programming is a game of regular practice. I would suggest you to pick a topic of your choice/interest and start playing with some example datasets; or better yet, if you have your own data try that out first.

As always, if you have any questions or interesting ideas to share, please free to reach out to me or other members of the K-State PBG journal club.

## Happy Coding!!!

#### sessionInfo()

```
## R version 3.5.3 (2019-03-11)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 14393)
## Matrix products: default
##
## locale:
## [1] LC COLLATE=English United States.1252
## [2] LC CTYPE=English United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC NUMERIC=C
## [5] LC_TIME=English_United States.1252
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                   base
##
## other attached packages:
   [1] rcompanion_2.2.2 ggpubr_0.2.1
                                          magrittr_1.5
                                                           car_3.0-3
   [5] carData_3.0-2
                         nlme_3.1-137
                                          forcats_0.4.0
                                                           stringr_1.4.0
  [9] dplyr_0.8.3
                         purrr_0.3.2
                                          readr_1.3.1
                                                           tidyr_0.8.3
##
## [13] tibble_2.1.3
                         ggplot2_3.2.0
                                          tidyverse_1.2.1 pacman_0.5.1
##
## loaded via a namespace (and not attached):
  [1] matrixStats_0.54.0 lubridate_1.7.4
                                              httr_1.4.0
  [4] tools 3.5.3
                          backports_1.1.4
                                              utf8 1.1.4
## [7] R6_2.4.0
                           nortest 1.0-4
                                              lazyeval 0.2.2
```

```
## [10] colorspace_1.4-1
                           withr_2.1.2
                                               tidyselect_0.2.5
## [13] curl_4.0
                           compiler_3.5.3
                                               cli_1.1.0
                                               xml2 1.2.0
## [16] rvest 0.3.4
                           expm 0.999-4
                                               scales_1.0.0
## [19] sandwich_2.5-1
                           labeling_0.3
## [22] lmtest_0.9-37
                           mvtnorm_1.0-11
                                               multcompView_0.1-7
## [25] digest_0.6.20
                           foreign_0.8-71
                                               rmarkdown 1.14
## [28] rio 0.5.16
                           pkgconfig 2.0.2
                                               htmltools 0.3.6
## [31] manipulate_1.0.1
                           rlang_0.4.0
                                               readxl_1.3.1
## [34] rstudioapi_0.10
                           generics_0.0.2
                                               zoo_1.8-6
## [37] jsonlite_1.6
                                               modeltools_0.2-22
                           zip_2.0.3
## [40] Matrix_1.2-17
                           fansi_0.4.0
                                               Rcpp_1.0.1
## [43] DescTools_0.99.28
                           munsell_0.5.0
                                               abind_1.4-5
## [46] stringi_1.4.3
                                               yaml_2.2.0
                           multcomp_1.4-10
## [49] MASS_7.3-51.4
                                               grid_3.5.3
                           plyr_1.8.4
## [52] parallel_3.5.3
                           crayon_1.3.4
                                               lattice_0.20-38
## [55] haven_2.1.1
                           splines_3.5.3
                                               hms_0.5.0
## [58] zeallot_0.1.0
                           knitr_1.23
                                               pillar_1.4.2
## [61] EMT 1.1
                           boot 1.3-23
                                               ggsignif_0.5.0
## [64] codetools_0.2-16
                           stats4_3.5.3
                                               glue_1.3.1
## [67] evaluate 0.14
                           data.table_1.12.2
                                               modelr_0.1.4
## [70] vctrs_0.2.0
                           cellranger_1.1.0
                                               gtable_0.3.0
## [73] assertthat 0.2.1
                           xfun_0.8
                                               openxlsx_4.1.0.1
                                               broom_0.5.2
## [76] coin_1.3-0
                           libcoin_1.0-4
## [79] survival_2.44-1.1 TH.data_1.0-10
```