

# Predicting Cherry Blossom Bloom Dates

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## Introduction

In this document, we demonstrate how we went about predicting the peak bloom dates in the coming decade for all four locations required by the competition. We use different modeling approaches due to different available data.

```
# Loading in packages and data
library(tidyverse)
library(forecast)
library(lubridate)
library(astsa)
library(here)
library(rnoaa)
library(gridExtra)

cherry <- read_csv(here('peak-bloom-prediction', 'data', 'washingtondc.csv')) %>%
  bind_rows(read_csv(here('peak-bloom-prediction', 'data', 'liestal.csv'))) %>%
  bind_rows(read_csv(here('peak-bloom-prediction', 'data', 'kyoto.csv'))) %>%
  select(location, year, bloom_date, bloom_doy)
```

## Predicting peak bloom dates for DC

In the field of phenology, plant growth is often modeled by using growing degree-day (GDD) as a predictor. Since the [National Park Service \(NPS\)](#) publishes dates of various growth stages for 2004 through 2021, we use this to build a growth model that helps us estimate the threshold GDD for which we can expect the plant to bloom. Details can be found in [Phenological Models of Flower Bud Stages and Fruit Growth of ‘Montmorency’ Sour Cherry Based on Growing Degree-day Accumulation by C. Zavalloni, J. Adresen, and J. Flore](#). The authors use 4 degrees Celsius as the base for calculating GDD and fit a logistic function to the phenological stages from side green as a function of accumulated GDD.

$$y = \frac{k}{1 + \left[ \frac{k-n_0}{n_0} \cdot \exp(-r \cdot \text{GDD}) \right]}$$

where  $y$  is the phenological stage and  $k$  and  $r$  are empirical factors related to growth rate. Now, with just enough knowledge to be dangerous, we attempt to use this model to fit the growth stages published by the NPS as a function of cumulative GDD. Note that the authors use  $n_0 = 2$  because they were interested in phenological stages after side green (Stage 2). To be consistent with the paper’s authors, we label the phenological stages published by the NPS in a similar fashion.

## Obtaining temperature and phenological data for DC

Since the species in the paper is different from what we’re interested in, we must first identify the typical day of year when the cherry blossom trees in DC reach the green buds stage (Stage 3, as defined by the authors).

```
# Import historical weather data from DayMet
dc_tmin <- read_csv(here('peak-bloom-prediction', 'data', 'statistics_tmin.csv'))
dc_tmax <- read_csv(here('peak-bloom-prediction', 'data', 'statistics_tmax.csv'))
```

```

# DayMet data stops at 2020. Use rnoaa for 2021.
# Original code from demo analysis
# https://competition.statistics.gmu.edu/wp-content/uploads/2022/02/demo_analysis.html
getTemperature <- function (stationid,date_min,date_max) {
  tbls <- ghcnv2_search(stationid = stationid, var = c("tmax","tmin"),
    date_min = date_min, date_max = date_max)
  tmax <- tbls[[1]] %>% select(id,tmax,date)
  tmin <- tbls[[2]] %>% select(tmin,date)
  temp <- inner_join(tmax,tmin,by="date") %>%
    group_by(date) %>%
    # Calculate average and convert unit from tenth degree Celsius to regular Celsius
    summarize(id,tavg=0.5*sum(tmax,tmin,na.rm=TRUE)/10) %>%
    ungroup() %>%
    mutate(year = year(date))
  return(temp)
}

dc_noaa <- getTemperature("USC00186350","2021-01-01","2022-01-31") %>%
  select(-id)

# Temperature data
dc_temp <- left_join(dc_tmin,dc_tmax, by = 'dt') %>%
  select(dt,value_mean.x,value_mean.y) %>%
  rename(tmin=value_mean.x,tmax=value_mean.y) %>%
  summarize(date=dt,tavg=0.5*(tmin+tmax), year=year(date)) %>%
  bind_rows(dc_noaa)

# Phenological stage data
dc_bloomdata <- cherry %>% filter(location=='washingtondc') %>%
  summarize(date=bloom_date,bloom_doy) %>%
  summarize(year=year(date),FullBloom=bloom_doy)

phenostage <- read_csv(here('peak-bloom-prediction','data',
  'Phenological_Stage.csv')) %>%
  inner_join(dc_bloomdata,by='year')

summary(phenostage$GreenBuds)

```

```

##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
##    50.00   59.00   64.00   63.94   69.50   77.00

```

This result suggests that Stage 3 occurs as early as 50 days from the start of the year. Thus, we shift the dates by 49 days to start calculating GDD to ensure that we start calculating GDD as soon as Stage 3 occurs. We also set  $n_0 = 3$  because we started from Stage 3 instead of 2.

## Building the growth model

As mentioned above, we calculate the GDD using a base of 4 degrees Celsius and starting from 49 days from the beginning of the year. Then, we fit the known phenological stages for the cherry blossoms in DC to the calculated GDDs.

```

thresh <- 4
GDD_final <- dc_temp %>%
  mutate(NewYear=year(date %m-% period("49 day"))) %>%
  mutate(DD = ifelse(tavg - thresh > 0, tavg - thresh, 0)) %>%
  group_by(NewYear) %>%
  mutate(GDD = cumsum(DD),doy = yday(date)) %>%
  ungroup()

```

```

# obtain GDD for each phenological stage for DC
stages <- tibble(stage=names(phenostage)[2:7], stagenum=seq(3,8))
pheno_GDD <- phenostage %>%
  select(-GreenBuds) %>% #This stage is the starting point for our model.
  pivot_longer(cols=FloretsVisible:FullBloom, names_to = "stage", values_to="doy") %>%
  left_join(GDD_final, by=c("year", "doy")) %>%
  summarize(year, stage, doy, GDD) %>%
  left_join(stages, by='stage')

# Fit growth model
n0 <- 3 #Starting phenological stage for our model
growth_model <- nls(stagenum ~ k/(1+(k-n0)/n0*exp(-r*GDD)),
  data=pheno_GDD, start=list(k=1, r=0.01))
summary(growth_model)

```

```

##
## Formula: stagenum ~ k/(1 + (k - n0)/n0 * exp(-r * GDD))
##
## Parameters:
##      Estimate Std. Error t value Pr(>|t|)
## k 10.223870    1.362473    7.504 4.72e-11 ***
## r  0.011973    0.001598    7.495 4.93e-11 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.9374 on 88 degrees of freedom
##
## Number of iterations to convergence: 8
## Achieved convergence tolerance: 7.797e-06

```

The model results show that both  $k$  and  $r$  are significant. However, inference is not of interest to us. We only care about prediction performance.

## Checking performance on historical data

Now that we have the growth model, we can test its prediction performance out on historical data that we did not use.

```

testdata <- GDD_final %>% filter(year<=2003) %>% #Used data from 2004 onward
  group_by(year) %>%
  filter(doy>=50 & doy <= 200) %>%
  ungroup() %>%
  select(date, GDD)

preds <- tibble(date=testdata$date, GDD=testdata$GDD,
  pred=predict(growth_model, newdata=testdata)) %>%
  mutate(year=year(date))

# Predicted date is the earliest date when predicted stage rounds to Stage 8
# Compare predicted date to actual historical dates
comp <- preds %>% group_by(year) %>%
  filter(round(pred) >= 8) %>%
  filter(row_number()==1) %>%
  ungroup() %>%
  mutate(pred.doy=yday(date)) %>%
  inner_join(dc_bloomdata, by='year') %>%
  summarize(year, GDD, pred.doy, act.doy=FullBloom, err_GDD=abs(pred.doy-act.doy)) %>%
  # compare to using simply using a lag predictor

```

```
mutate(lag_bloom = lag(act.doy),err_lag=abs(act.doy - lag_bloom))

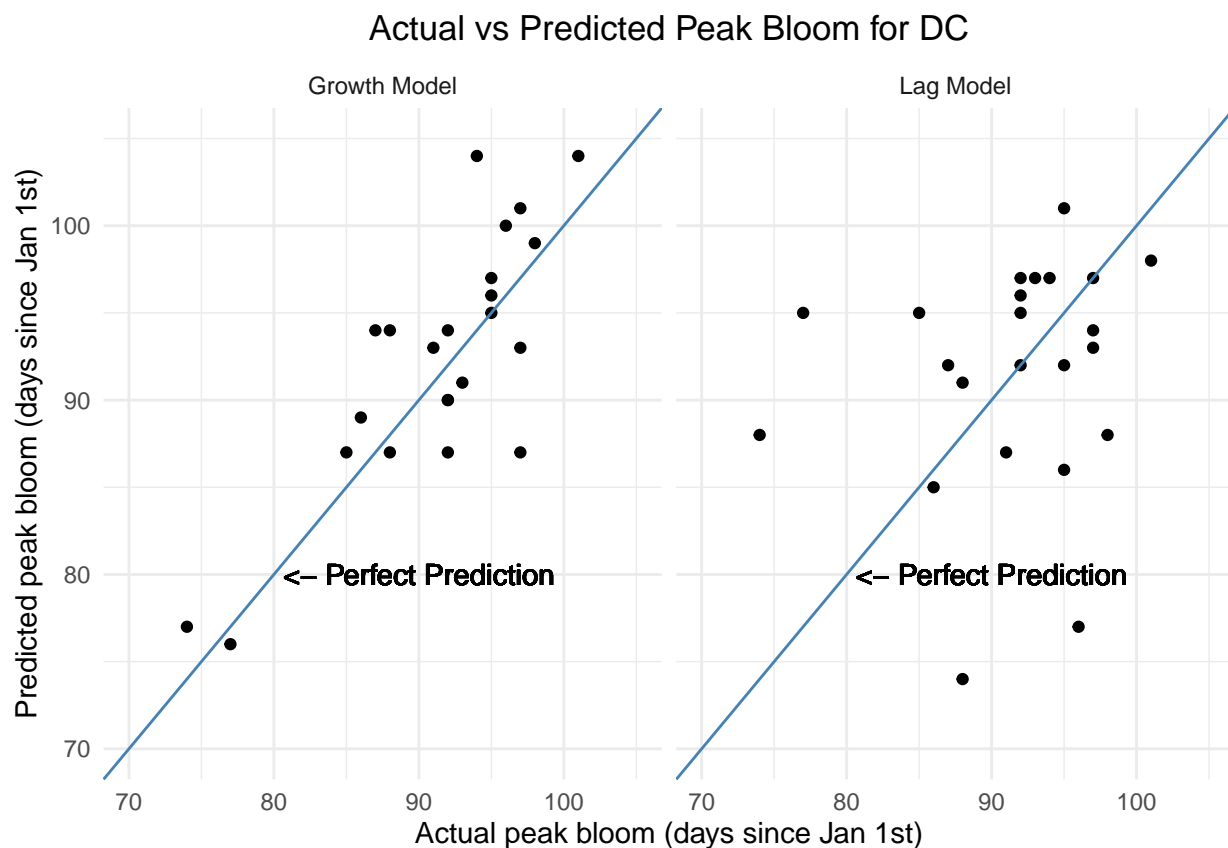
data.frame(MAE_gdd=mean(comp$err_GDD[which(!is.na(comp$err_lag))]),
  MAE_lag=mean(comp$err_lag,na.rm=TRUE))

##      MAE_gdd MAE_lag
## 1 3.347826 6.304348
```

Our growth model has a much lower mean absolute error (MAE) compared to simply using the previous year's bloom date as a predictor. This is good because this is the bare minimum a useful model should be able to do. This is more apparent when we plot the two models against the actual peak bloom days since January 1st.

```
plt_data <- comp %>% rename(Actual=act.doy,GrowthModel=pred.doy,LagModel=lag_bloom) %>%
  pivot_longer(cols=c("GrowthModel","LagModel"),names_to="type",values_to="doy") %>%
  filter(!is.na(err_lag))
plt_data$type <- factor(plt_data$type,levels=c("GrowthModel","LagModel"),
  labels=c("Growth Model","Lag Model"))

# Plot Actual vs Predicted for Growth Model
ggplot(plt_data,aes(x=Actual)) + geom_point(aes(y=doy)) +
  facet_grid(~type) +
  geom_abline(slope=1,col="steelblue") +
  geom_text(aes(x=90,y=80,label="← Perfect Prediction")) +
  theme_minimal() +
  ggtitle("Actual vs Predicted Peak Bloom for DC") +
  xlab("Actual peak bloom (days since Jan 1st)") +
  ylab("Predicted peak bloom (days since Jan 1st)") +
  xlim(c(70,105)) + ylim(c(70,105)) +
  theme(plot.title=element_text(hjust=0.5))
```

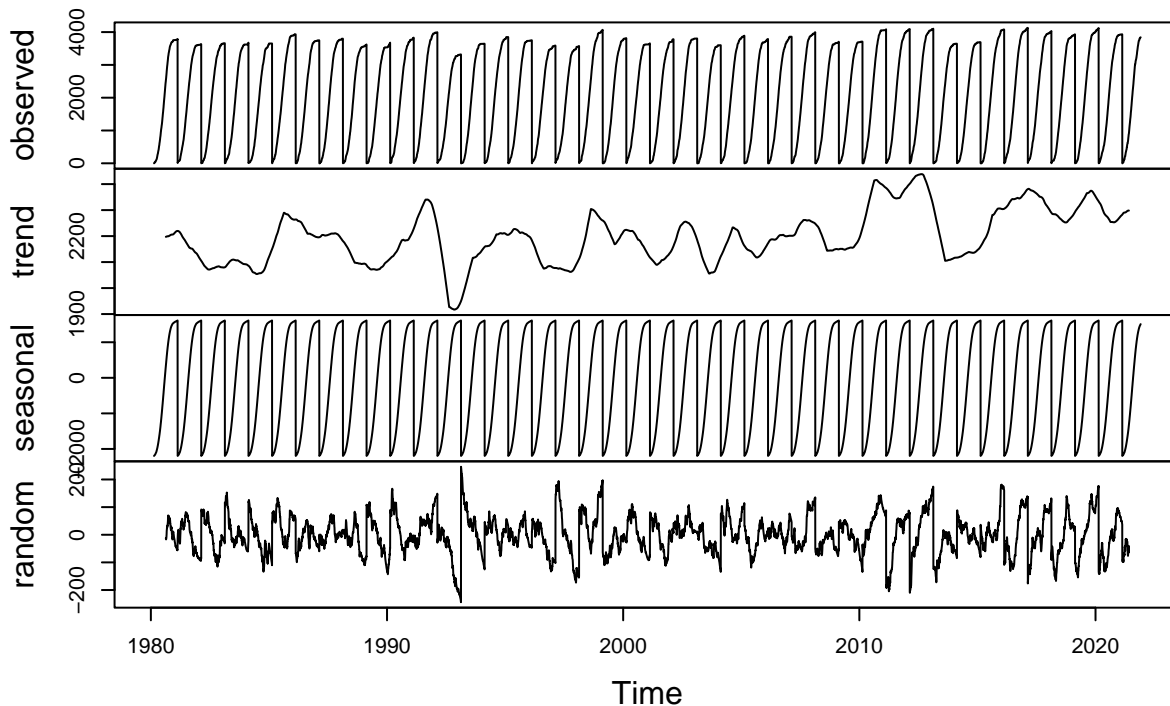


## Predicting future GDD and future peak bloom dates

For us to predict future peak bloom dates for DC, we must first predict future GDDs. We could predict future temperature and calculate GDD from that; however, that would be unnecessary because we always shift the dates by 49 days and use the base of 4 degrees Celsius to calculate GDD. We can directly predict GDD to avoid that step.

```
# Create time series
ts_gdd <- ts(GDD_final$GDD[50:nrow(GDD_final)],frequency = 365,
            #Only start counting GDD from day 50 & 2021 DC data only goes to November
            start=c(1980,50),end=c(2021,334))
plot(decompose(ts_gdd))
```

### Decomposition of additive time series



```
if (file.exists('pred_dc.rds')){
  pred_dc <- readRDS(here("pred_dc.rds"))
} else{
  # Fit time series to GDD
  ts_fit <- auto.arima(ts_gdd, ic = 'aicc')
  ar <- ts_fit$arma[1:2]
  ma <- ts_fit$arma[3:4]
  dd <- ts_fit$arma[5:7]

  # Use result from fitted time series to predict GDD until May 2031
  preddays <- difftime("2031-05-31", "2021-12-01", units="days")
  pred_dc <- sarima.for(ts_gdd, as.numeric(preddays), ar[1], dd[2], ar[2], ma[1], dd[3], ma[2], dd[1])
  saveRDS(pred_dc, here("pred_dc.rds"))
}
```

We can now predict the peak bloom day for DC using the predicted GDD as an input for the growth model.

```
# Forecasted GDD and growth stage
newdata <- tibble(date=seq.Date(as.Date("2021-12-01"),length.out=length(pred_dc$pred),by="day"),
                  GDD=pred_dc$pred,doy=yday(date),year=year(date))
future_growthstage <- predict(growth_model,newdata=newdata)
forecasts_dc <- newdata %>% mutate(pred.gs=future_growthstage) %>%
```

```
group_by(year) %>%
filter(doy >= 50 & doy <= 200) %>%
filter(round(pred.gs) >= 8) %>%
filter(row_number()==1) %>%
select(year,doy)

forecasts_dc
```

```
## # A tibble: 10 x 2
## # Groups:   year [10]
##   year    doy
##   <dbl> <dbl>
## 1  2022     86
## 2  2023     86
## 3  2024     86
## 4  2025     85
## 5  2026     85
## 6  2027     85
## 7  2028     84
## 8  2029     83
## 9  2030     83
## 10 2031     83
```

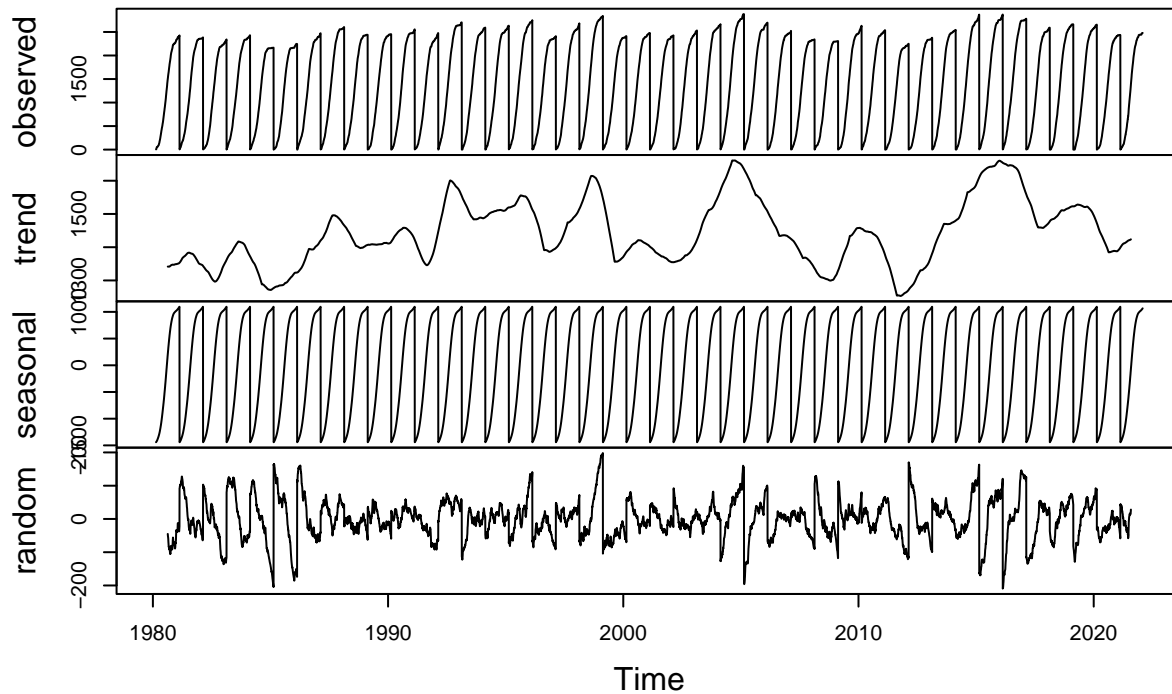
## Predicting peak bloom dates for Vancouver

Vancouver doesn't provide historical peak bloom dates. However, we can use the growth model from DC to predict future peak bloom dates because the tree species are identical (or close to identical) and the peak bloom definition is the same. We simply need to calculate historical GDD using the same definitions and predict future GDD for Vancouver to be able to predict the peak bloom dates.

### Obtaining historical weather data

Similar to DC, we obtained the historical weather data for Vancouver via DayMet. However, this dataset only goes to December 2019, so we use `rnoaa` to obtain the remaining data up to Jan 2022.

## Decomposition of additive time series



```
## # A tibble: 10 x 2
## # Groups:   year [10]
##   year    doy
##   <dbl> <dbl>
## 1  2022    109
## 2  2023    108
## 3  2024    108
## 4  2025    107
## 5  2026    106
## 6  2027    106
## 7  2028    106
## 8  2029    105
## 9  2030    104
## 10 2031    104
```

## Predicting peak bloom dates for Liestal

The cherry tree in Liestal is a different species from the ones in DC and Vancouver. Ideally, we would want to fit a new growth model using historical phenological stages. Unfortunately, such a dataset is not readily available. So we use the same growth model from DC to calculate the mean growth stage for historical peak bloom dates. We use the average growth stage as the cutoff for our prediction.

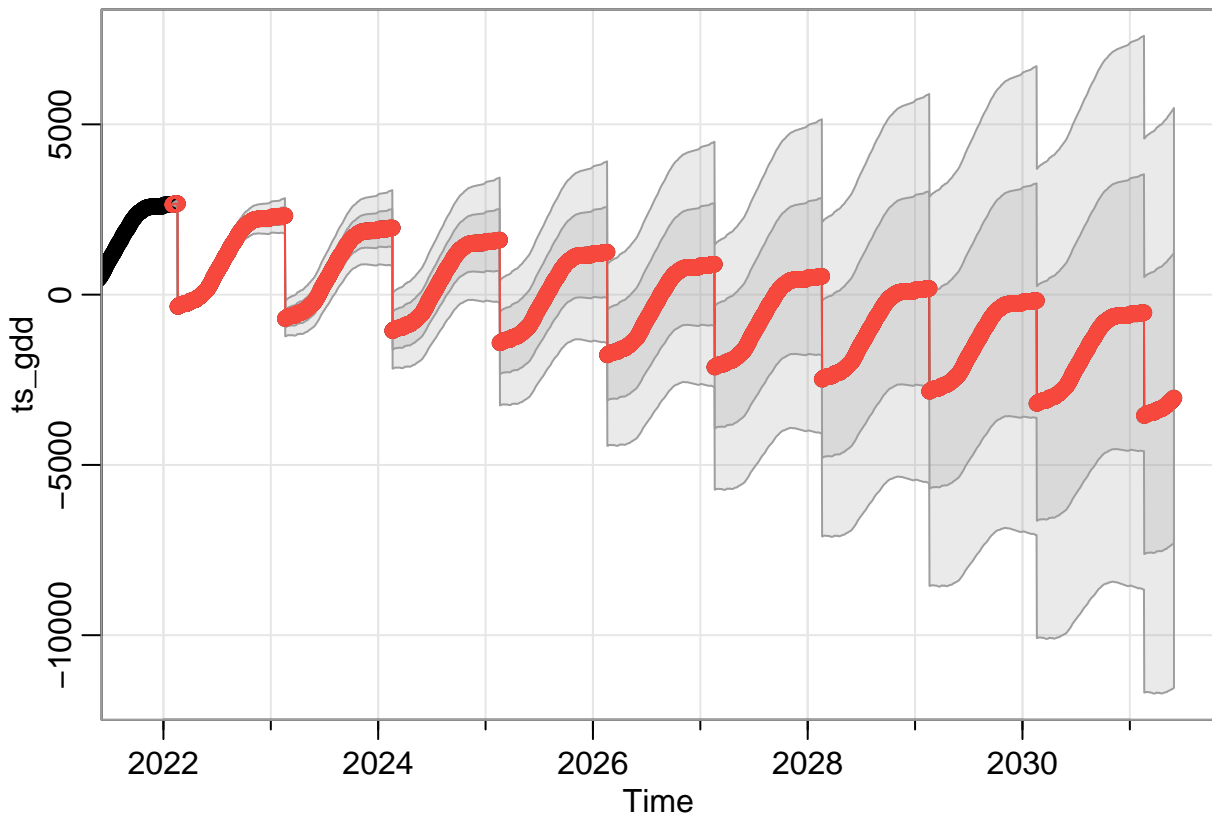
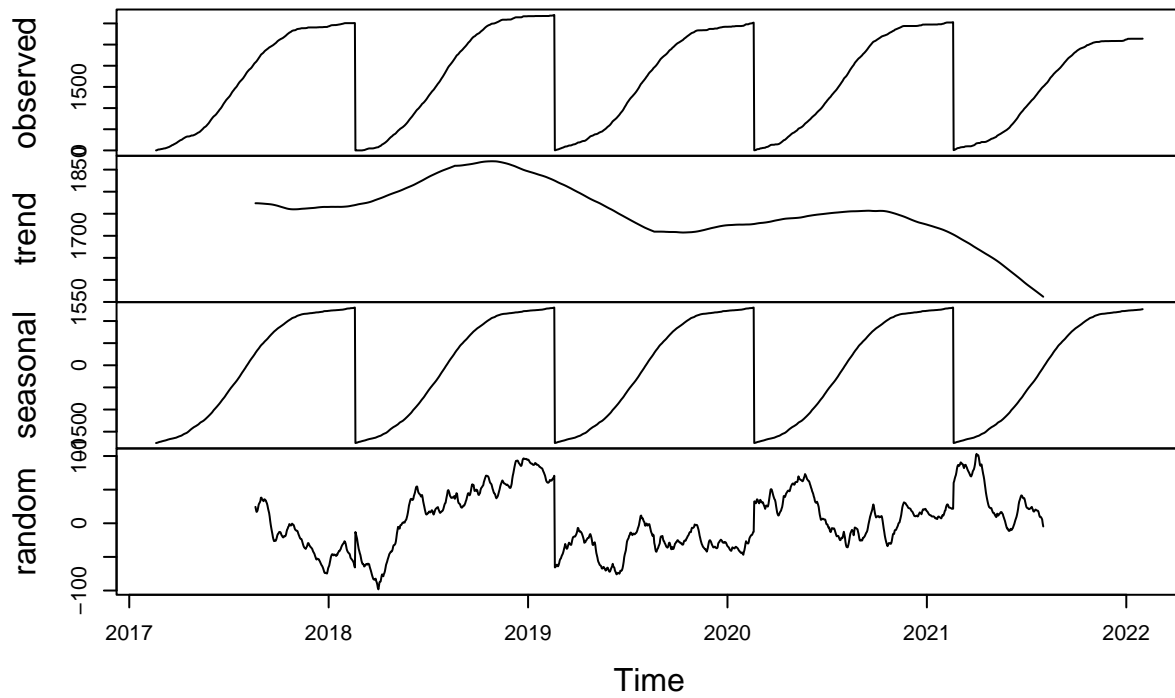
```
# Check historical GDD and corresponding predicted growth stages for actual bloom dates in Liestal
testdata <- cherry %>%
  left_join(GDD_final %>% select(date,GDD), by=c('bloom_date'='date')) %>%
  filter(year >= 2017 & location=='liestal') %>%
  select(GDD)

mean(predict(growth_model,newdata=testdata))
```

```
## [1] 6.798815
```

We can approximate the peak bloom for the cherry trees in Liestal (as defined by MeteoSwiss) using the growth model for the trees in DC if we use 6.8 as a threshold for the predicted growth stage. In other words, we estimate the peak bloom date by finding the earliest date for which the predicted growth stage rounds to 6.8.

## Decomposition of additive time series



```
## # A tibble: 3 x 2
## # Groups:   year [3]
##   year   doy
##   <dbl> <dbl>
```



```
## 1 2022 148
## 2 2023 171
## 3 2024 195
```

## Predicting peak bloom dates for Kyoto

Similar to predicting peak bloom dates for Liestal, the peak bloom dates for Kyoto (as defined by a local newspaper in Arashiyama) need to be approximated using the growth model for the DC cherry trees.

```
jp_temp <- temps %>% filter(city == 'Kyoto') %>%
  mutate(doy = yday(date)) %>%
  filter(doy != 366) #Leap year mess

# GDD calculations
GDD_final <- jp_temp %>%
  mutate(NewYear=year(date %m-% period("49 day"))) %>%
  mutate(DD = ifelse(tavg - thresh > 0, tavg - thresh, 0)) %>%
  group_by(NewYear) %>%
  mutate(GDD = cumsum(DD),doy = yday(date)) %>%
  ungroup()

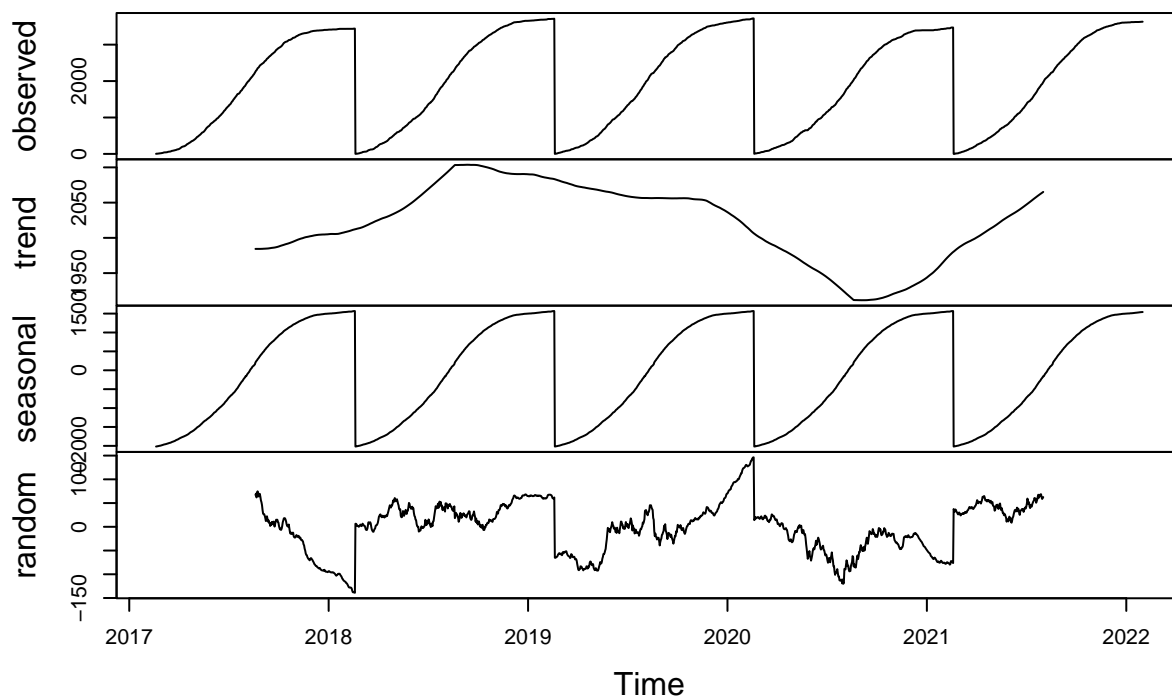
# Check historical GDD and corresponding predicted growth stages for actual bloom dates in Liestal
testdata <- cherry %>%
  left_join(GDD_final %>% select(date,GDD), by=c('bloom_date'='date')) %>%
  filter(year >= 2017 & location=='kyoto') %>%
  select(GDD)

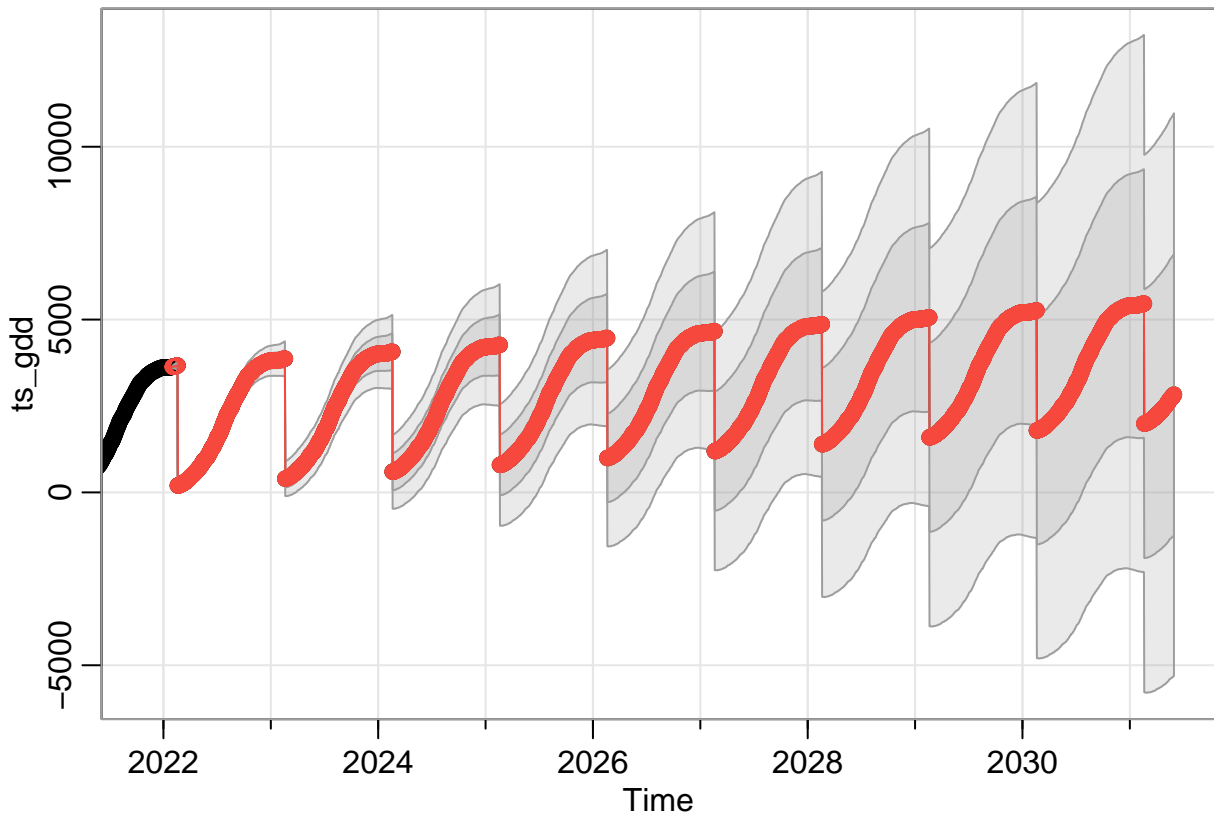
mean(predict(growth_model,newdata=testdata))

## [1] 8.165088
```

Based on the above result, we estimate the peak bloom date by finding the earliest date for which the predicted growth stage rounds to 8.2.

## Decomposition of additive time series





```
## # A tibble: 10 x 2
## # Groups:   year [10]
##   year    doy
##   <dbl> <dbl>
## 1  2022     50
## 2  2023     50
## 3  2024     50
## 4  2025     50
## 5  2026     50
## 6  2027     50
## 7  2028     50
## 8  2029     50
## 9  2030     50
## 10 2031     50
```

## Final predictions

The summary table below shows the final predictions for the four locations from 2022 to 2031.

## Closing thoughts

If historical phenological stages were recorded for all locations in this competition, we would have been able to fit a growth model for each species and location. This would have allowed us to create more accurate models. Additionally, weather data was extremely limited or unreliable for locations outside the United States. If historical weather data was free and readily available to the public, we would have been able to create a better time series model for GDD.