

The Rice example: illustrating the first five steps for smoothing and extracting traits (SET) using growthPheno

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29 February, 2024

This example is based on the data whose analysis has been published by Al-Tamimi et al. (2016). The five steps of the method for smoothing and extracting traits (SET) described in detail in Brien et al. (2020) is illustrated for this data.

Initialize

Step 1: Import, select and derive longitudinal data

Step 1(a): Import the data

```
data(RiceRaw.dat)
```

Step 1(b): Organize the data

Here the imaging variables are selected and covariates and factors added to produce `longi.dat`.

```
longi.dat <- prepImageData(data=RiceRaw.dat, smarthouse.lev=c("NE","NW"))

longi.dat <- designFactors(data = longi.dat, insertName = "Reps",
                           nzones = 3, designfactorMethod="StandardOrder")

## Particular edits to longi.dat - add Days after treatment (xDAT)
longi.dat$xDAT <- longi.dat$xDAP - 29
longi.dat <- with(longi.dat, longi.dat[order(Snapshot.ID.Tag,DAP), ])
```

Step 1(c): Derive longitudinal traits that result in a value for each observation

```
# Set responses
responses.image <- c("PSA")
responses.smooth <- paste0("s", responses.image)

# Form growth rates for each observation of a subset of responses by differencing
```

```

longi.dat <- byIndv4Times_GRsDiff(longi.dat, responses = responses.image,
                                    times = "DAP",
                                    which.rates = c("AGR", "RGR"))

# Form PSA.WUI
longi.dat <- within(longi.dat,
                      PSA.WUI <- WUI(PSA.AGR*DAP.diffs, WU))

# Add cumulative responses
longi.dat <- within(longi.dat,
{
  WU.cum <- unlist(by(WU, Snapshot.ID.Tag,
                        cumulate, exclude.1st=TRUE))
  WUI.cum <- PSA / WU.cum
})
# Check longi.dat
head(longi.dat)

```

```

##   Snapshot.ID.Tag DAP Smarthouse Lane Position xDAP Snapshot.Time.Stamp
## 1      045727-C  28        NE   13       2    28 2015-02-18 05:31:00
## 2      045727-C  30        NE   13       2    30 2015-02-20 05:23:00
## 3      045727-C  31        NE   13       2    31 2015-02-21 05:23:00
## 4      045727-C  32        NE   13       2    32 2015-02-22 05:23:00
## 5      045727-C  33        NE   13       2    33 2015-02-23 05:24:00
## 6      045727-C  34        NE   13       2    34 2015-02-24 10:15:00
##   Hour Reps Zone cZone SHZone ZLane ZMainunit Subunit cMainPosn cPosn
## 1  5.516667    1   1   -1     1     1       1     1   -10.5   -11
## 2  5.383333    1   1   -1     1     1       1     1   -10.5   -11
## 3  5.383333    1   1   -1     1     1       1     1   -10.5   -11
## 4  5.383333    1   1   -1     1     1       1     1   -10.5   -11
## 5  5.400000    1   1   -1     1     1       1     1   -10.5   -11
## 6 10.250000    1   1   -1     1     1       1     1   -10.5   -11
##   Genotype.ID Treatment.1 Weight.Before Weight.After Water.Amount WU          PSA
## 1      121146     Control       4013       4032       22  NA 55.311
## 2      121146     Control       4062       4085       26 -30 80.130
## 3      121146     Control       4040       4085       48  45 94.788
## 4      121146     Control       4032       4086       56  53 108.613
## 5      121146     Control       4027       4086       61  59 133.677
## 6      121146     Control       4012       4086       76  74 157.847
##   PSA.SV1 PSA.SV2 PSA.TV Boundary.Points.To.PSA.Ratio.SV1
## 1  11.307  15.456 28.548                  0.563633
## 2  25.816  21.768 32.546                  0.342539
## 3  31.627  23.604 39.557                  0.346413
## 4  37.702  30.704 40.207                  0.342316
## 5  39.861  40.317 53.499                  0.376308
## 6  48.086  46.873 62.888                  0.320904
##   Boundary.Points.To.PSA.Ratio.SV2 Boundary.Points.To.PSA.Ratio.TV
## 1                           0.371442                  0.233571
## 2                           0.439085                  0.220304
## 3                           0.411922                  0.230477
## 4                           0.365262                  0.228343
## 5                           0.346876                  0.222883
## 6                           0.367504                  0.219342

```

```

##   Caliper.Length.SV1 Caliper.Length.SV2 Caliper.Length.TV Compactness.SV1
## 1      736.872        792.324       888.821      0.0491248
## 2      728.754        785.611       797.924      0.0994986
## 3      779.808        889.427       797.332      0.1061870
## 4      956.613        896.909       861.304      0.1008410
## 5     1076.500        1123.540      1202.150      0.0809829
## 6     1016.490        1235.360      1277.480      0.1033860
##   Compactness.SV2 Compactness.TV Convex.Hull.PSA.SV1 Convex.Hull.PSA.SV2
## 1      0.0815964      0.1069410      230.169      189.420
## 2      0.0683533      0.1051310      259.461      318.463
## 3      0.0658089      0.1030830      297.843      358.675
## 4      0.0733596      0.0916347      373.877      418.541
## 5      0.0796217      0.0908369      492.215      506.357
## 6      0.0621912      0.0986004      465.109      753.692
##   Convex.Hull.PSA.TV Center.Of.Mass.Y.SV1 Center.Of.Mass.Y.SV2
## 1      266.952        1822.21       1785.29
## 2      309.575        1809.31       1808.03
## 3      383.738        1815.14       1826.45
## 4      438.775        1827.60       1874.88
## 5      588.957        1823.38       1861.62
## 6      637.807        1830.35       1843.54
##   Max.Dist.Above.Horizon.Line.SV1 Max.Dist.Above.Horizon.Line.SV2 xDAT
## 1                      612                  626    -1
## 2                      637                  636    1
## 3                      591                  628    2
## 4                      650                  691    3
## 5                      599                  618    4
## 6                      695                  707    5
##   DAP.diffs PSA.AGR   PSA.RGR   PSA.WUI   WUI.cum WU.cum
## 1      NA     NA      NA      NA      NA     NA
## 2      2 12.4095 0.1853393 -0.8273000 -2.6710000  -30
## 3      1 14.6580 0.1679925  0.3257333  6.3192000   15
## 4      1 13.8250 0.1361483  0.2608491  1.5972500   68
## 5      1 25.0640 0.2076353  0.4248136  1.0525748  127
## 6      1 24.1700 0.1661998  0.3266216  0.7853085  201

```

Step 2: Exploratory analysis

Step 2(a): Fit splines to smooth the longitudinal trends in the primary traits and calculate their growth rates

The `smoothing.method` used is `direct` and `df` is set to 4. The growth rates are calculated by difference, rather than from the spline derivatives.

```

# Smooth responses and form growth rates by differences
for (response in c(responses.image, "WU"))
  longi.dat <- byIndv4Times_SplinesGRs(data = longi.dat, response = response,
                                         response.smoothed = paste0("s", response),
                                         individuals = "Snapshot.ID.Tag", times="DAP",
                                         df = 4)

## Warning in FUN(X[[i]], ...): Need at least 4 distinct x values to fit a spline

```

```

## - all fitted values set to NA

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## - all fitted values set to NA

## Warning in FUN(X[[i]], ...): Need at least 4 distinct x values to fit a spline
## - all fitted values set to NA

## Warning in log(PGR(x, time.diffs, lag = lag)): NaNs produced

# Finalize longi.dat
longi.dat <- with(longi.dat, longi.dat[order(Snapshot.ID.Tag, xDAP), ])

```

Step 2(b): Compare plots of unsmoothed and smoothed longitudinal data

```

responses.longi <- c("PSA", "PSA.AGR", "PSA.RGR", "PSA.WUI")
responses.smooth.plot <- c("sPSA", "sPSA.AGR", "sPSA.RGR")
titles <- c("PSA (kpixels)",
           "PSA AGR (kpixels per day)", "PSA RGR (per day)",
           "PSA WUI (kpixels per mL)")
titles.smooth<-paste0("s", titles)
nresp <- length(responses.longi)

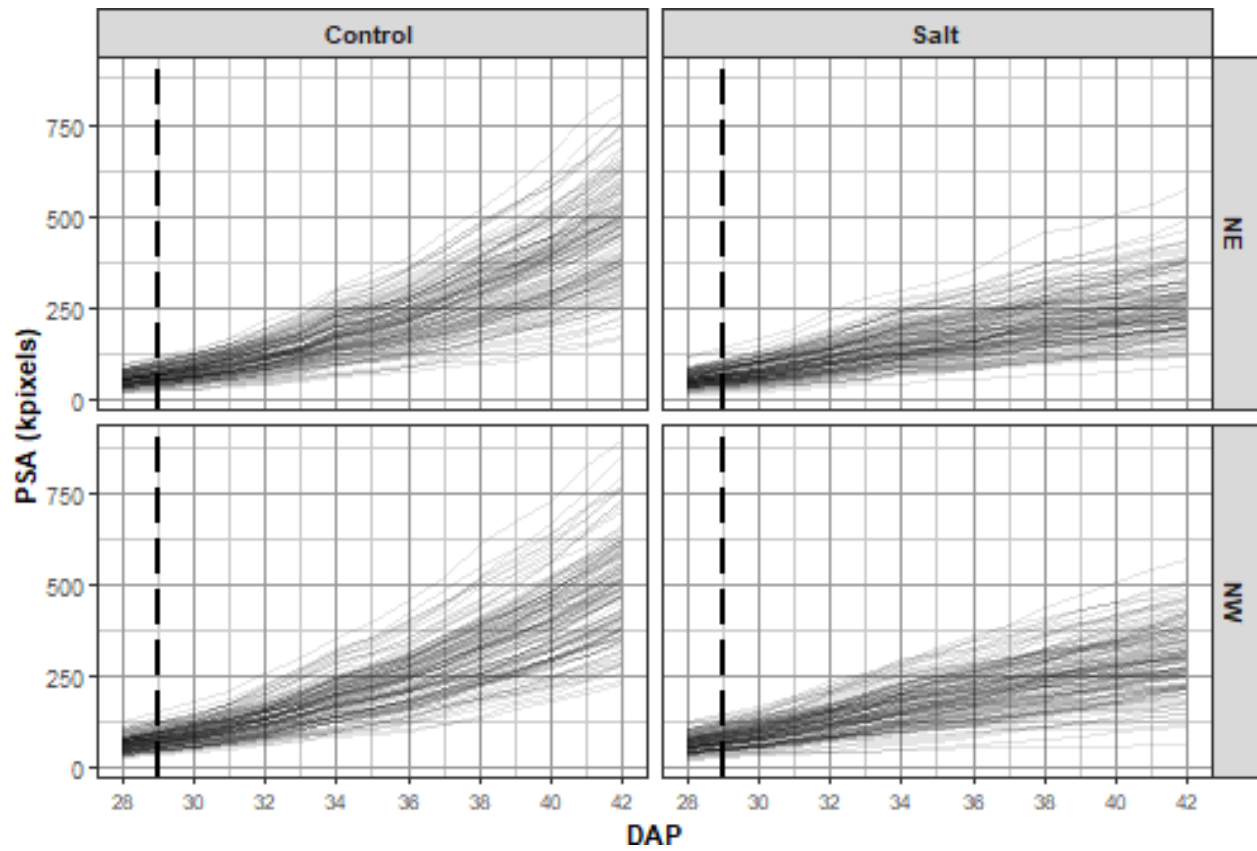
```

Plot unsmoothed profiles for all longitudinal responses

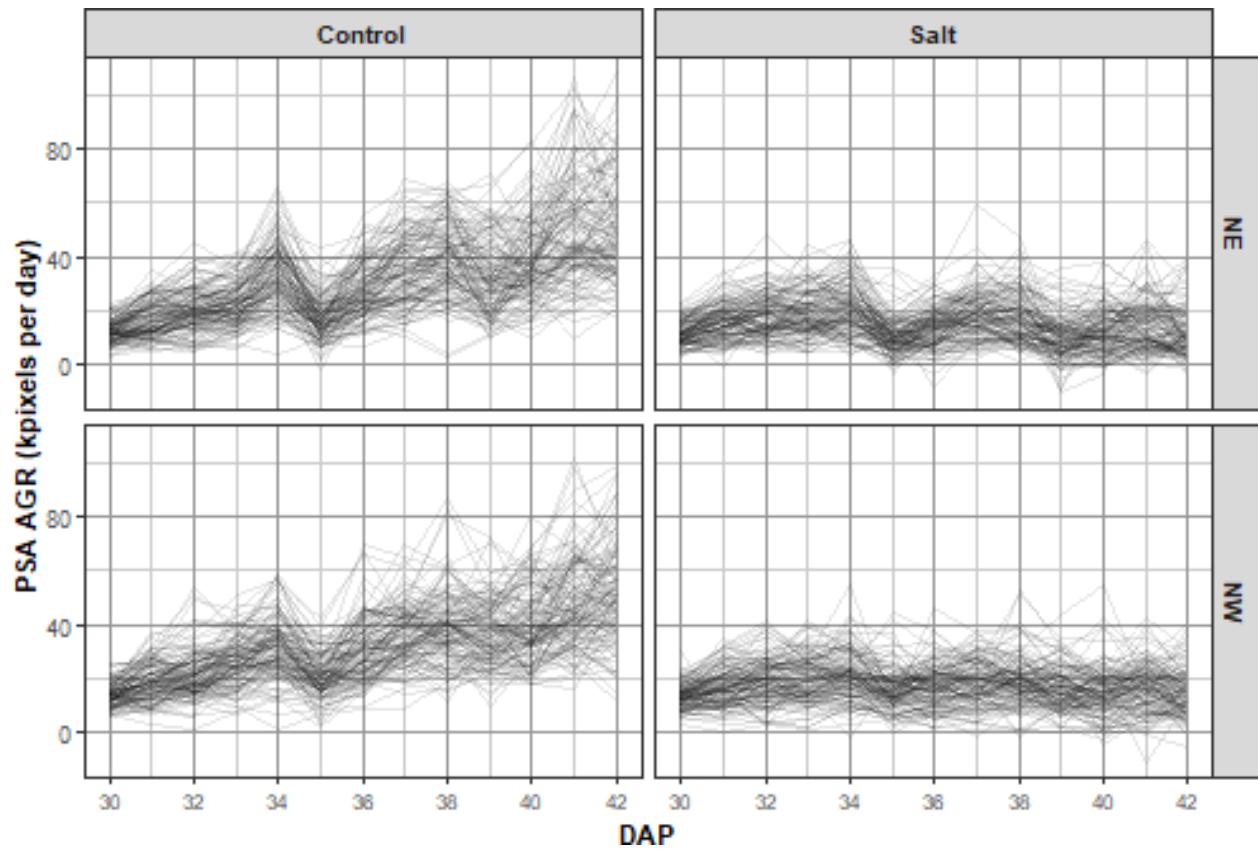
```

for (k in 1:nresp)
{
  plt <- plotProfiles(data = longi.dat, response = responses.longi[k],
                       y.title = titles[k], times = "DAP",
                       facet.x = "Treatment.1", facet.y = "Smarthouse",
                       breaks.spacing.x = 2,
                       printPlot=FALSE)
  plt <- plt + geom_vline(xintercept=29, linetype="longdash", linewidth=1)
  print(plt)
}

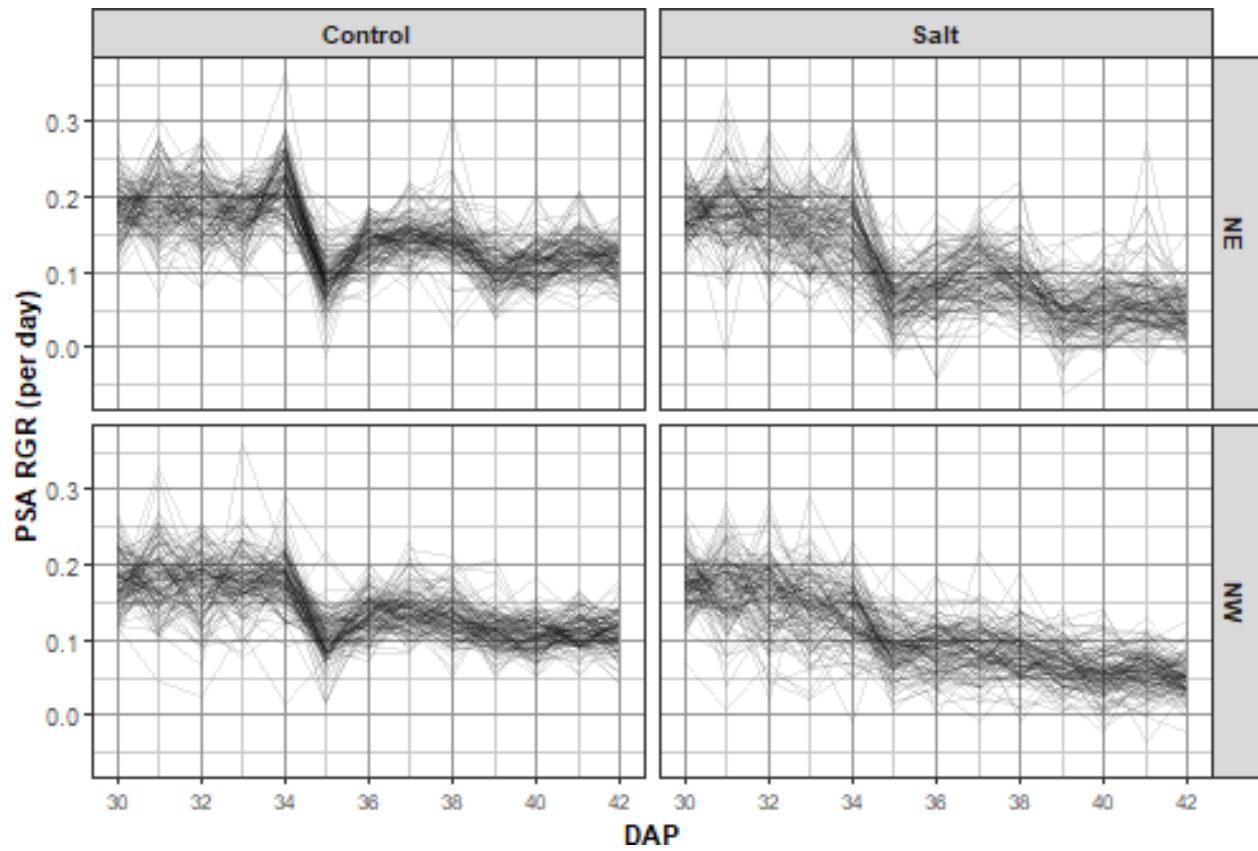
```



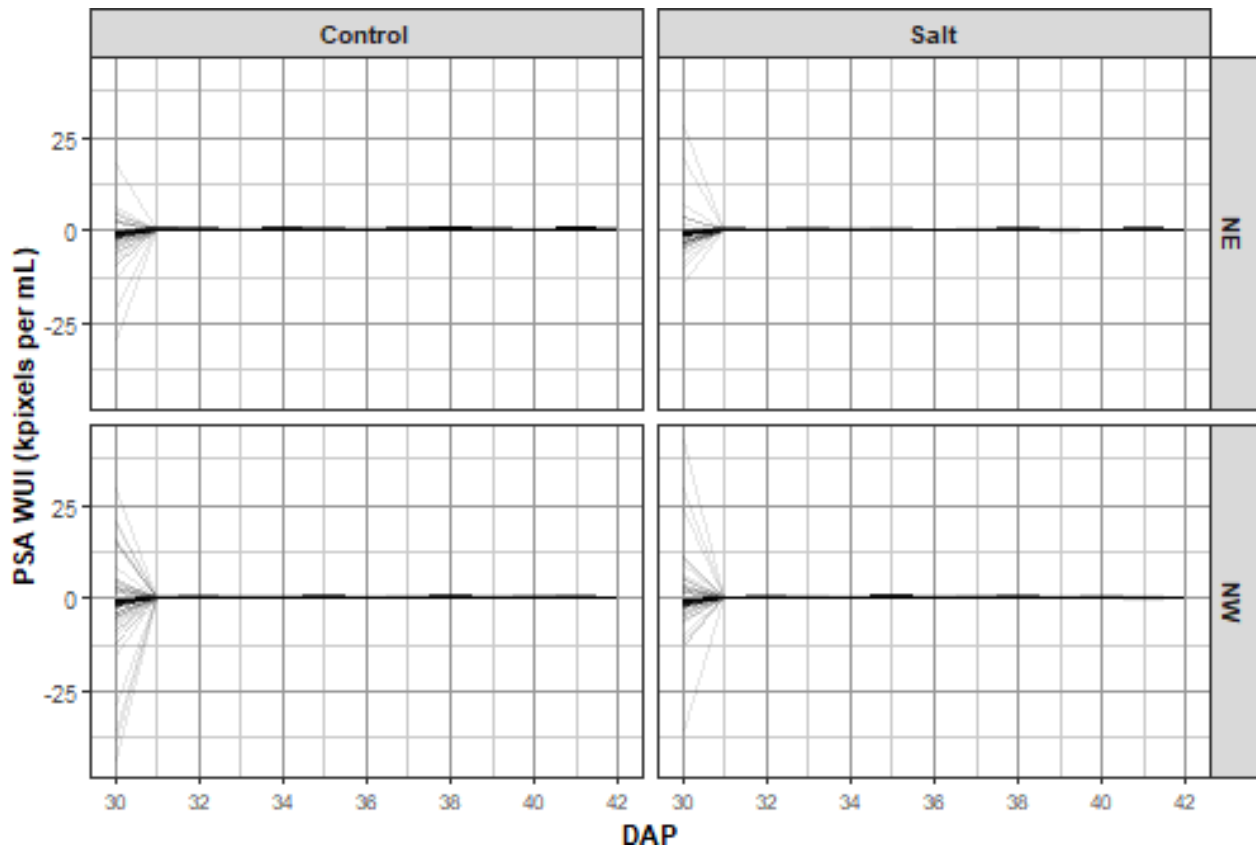
```
## Warning: Removed 4 rows containing missing values ('geom_vline()').
```



```
## Warning: Removed 4 rows containing missing values ('geom_vline()').
```



```
## Warning: Removed 4 rows containing missing values ('geom_vline()').
```

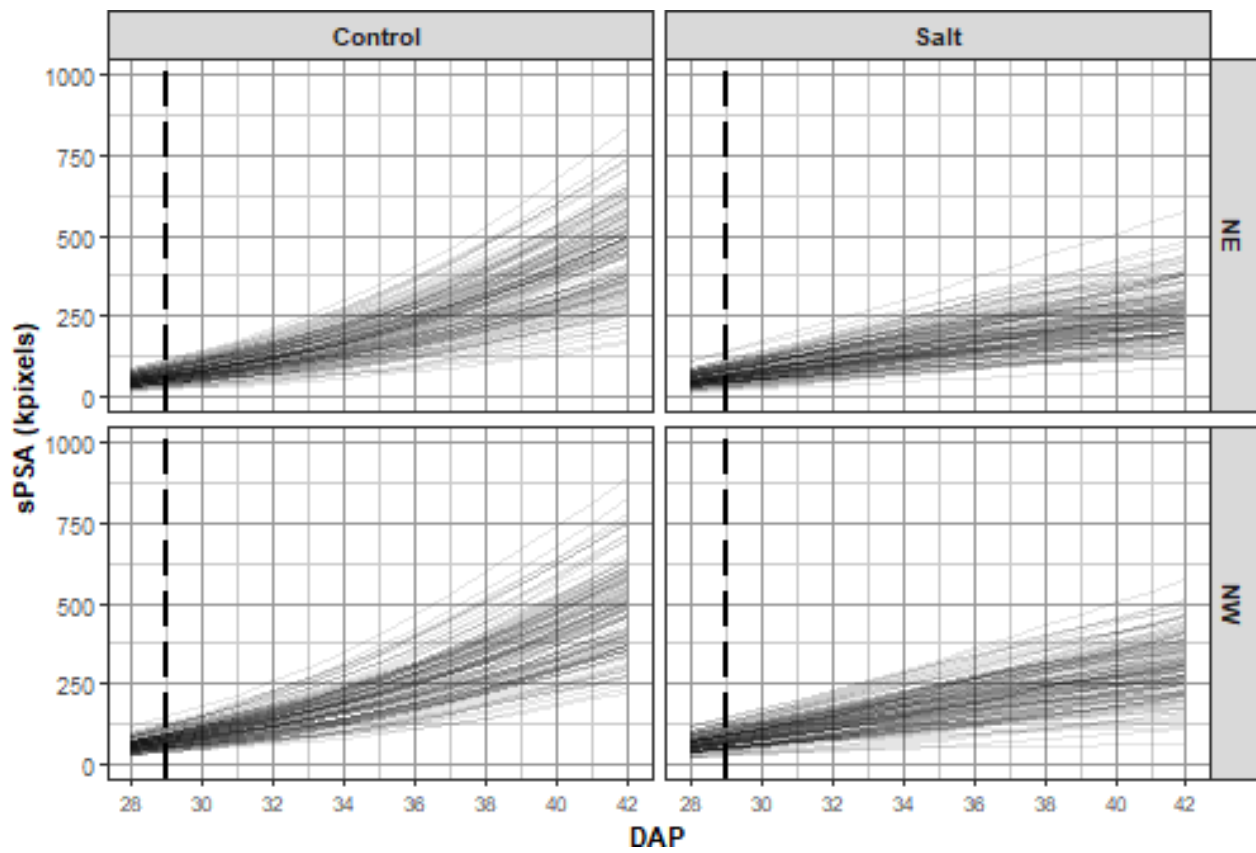


Plot smoothed profiles for all longitudinal responses

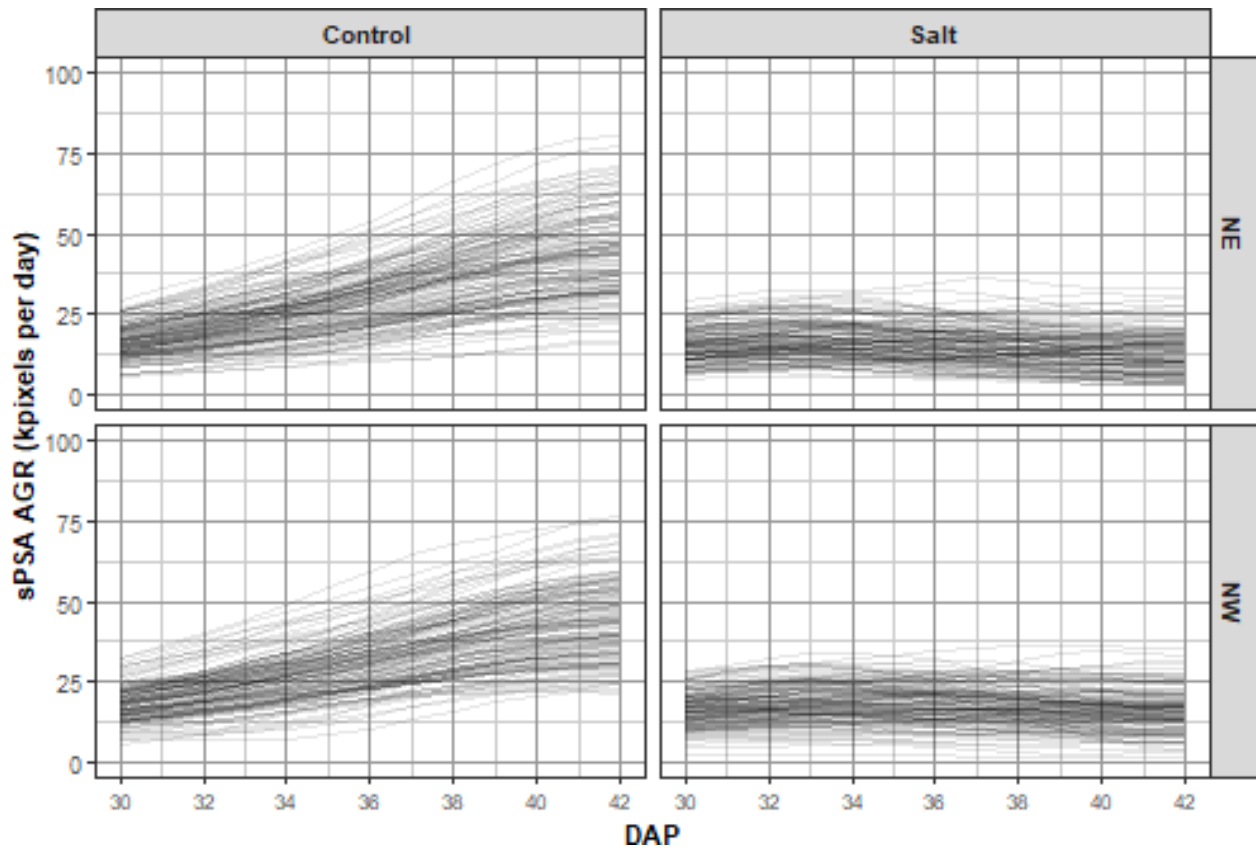
```

nresp.smooth <- length(responses.smooth.plot)
limits <- list(c(0,1000), c(0,100), c(0.0,0.40))
for (k in 1:nresp.smooth)
{
  plt <- plotProfiles(data = longi.dat, response = responses.smooth.plot[k],
    y.title = titles.smooth[k], times = "DAP",
    facet.x = "Treatment.1", facet.y = "Smarthouse",
    breaks.spacing.x = 2,
    printPlot=FALSE)
  plt <- plt + geom_vline(xintercept=29, linetype="longdash", linewidth=1) +
    scale_y_continuous(limits=limits[[k]])
  print(plt)
}

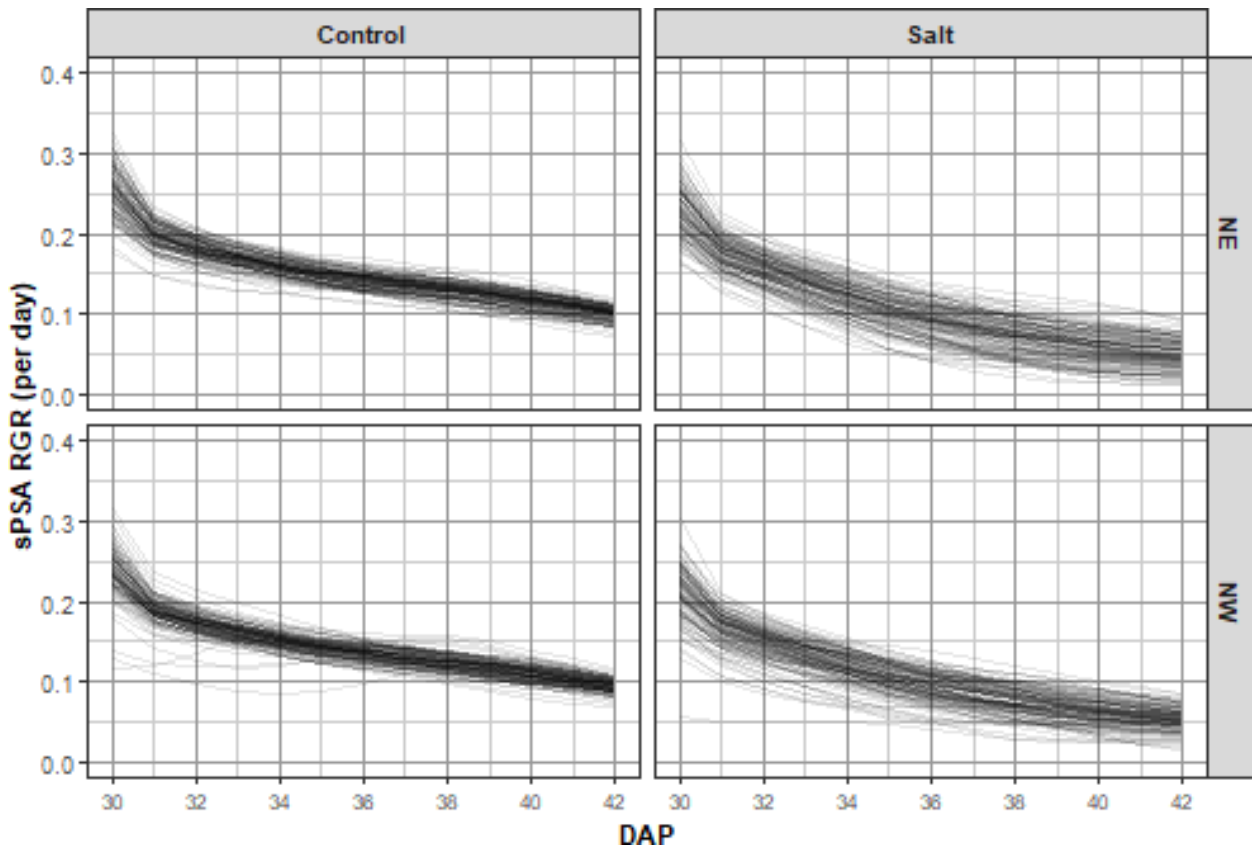
```



```
## Warning: Removed 4 rows containing missing values ('geom_vline()').
```



```
## Warning: Removed 4 rows containing missing values ('geom_vline()').
```



Step 3: Choose the smoothing method and DF

This step has been omitted.

Step 4: Identify potential outliers and clean the data

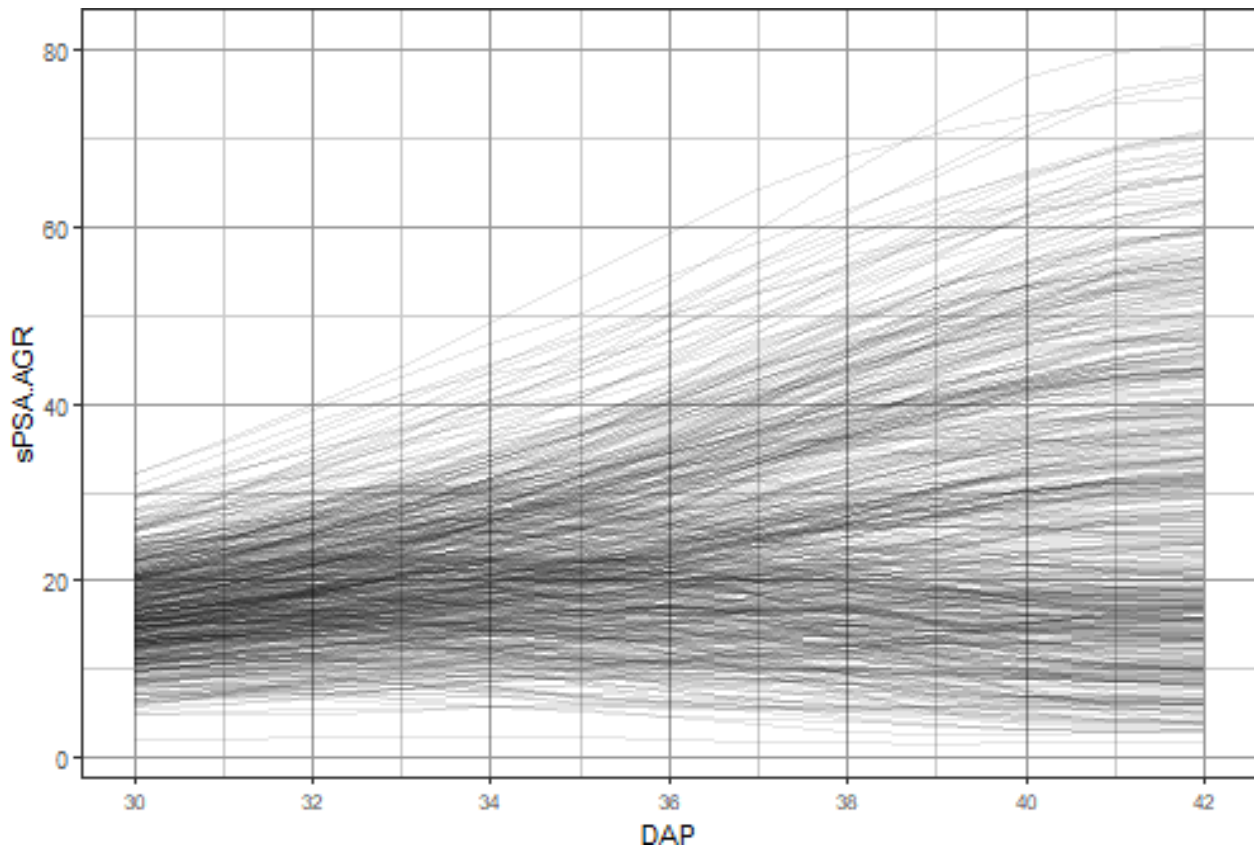
It has been decided that plants whose smoothed AGR are less than 2.5 after Day 40 are growing so slowly as to be considered anomalous. These plants are identified using `plotAnom`. Their values on Day 42 are printed. The plants are plotted without the anomalous plants followed by a plot of just the anomalous plants. The images of these anomalous plants were examined and no particular problems were identified with them. They were retained in the data.

```

anom.ID <- vector(mode = "character", length = 0L)
response <- "sPSA.AGR"
cols.output <- c("Snapshot.ID.Tag", "Smarthouse", "Lane", "Position",
                 "Treatment.1", "Genotype.ID", "DAP")
anomalous <- plotAnom(longi.dat, response=response, lower=2.5, start.time=40,
                      times = "DAP", vertical.line=29, breaks.spacing.x = 2,
                      whichPrint=c("innerPlot"), y.title=response)

## Warning: Removed 1 rows containing missing values ('geom_vline()').

```



```

subs <- subset(anomalous$data, sPSA.AGR.anom & DAP==42)
if (nrow(subs) == 0)
{ cat("\n#### No anomalous data here\n\n")
} else
{
  subs <- subs[order(subs[["Smarthouse"]], subs[["Treatment.1"]], subs[[response]]),]
  print(subs[c(cols.output, response)])
  anom.ID <- unique(c(anom.ID, subs$Snapshot.ID.Tag))
  outerPlot <- anomalous$outerPlot + geom_text(data=subs,
                                                aes_string(x = "DAP",
                                                           y = response,
                                                           label="Snapshot.ID.Tag"),
                                                size=3, hjust=0.7, vjust=0.5)
  print(outerPlot)
}

```

```

##           Snapshot.ID.Tag Smarthouse Lane Position Treatment.1 Genotype.ID DAP
## 6608          046495-S        NW     22      10       Salt    120952  42
##   sPSA.AGR
## 6608 1.809133

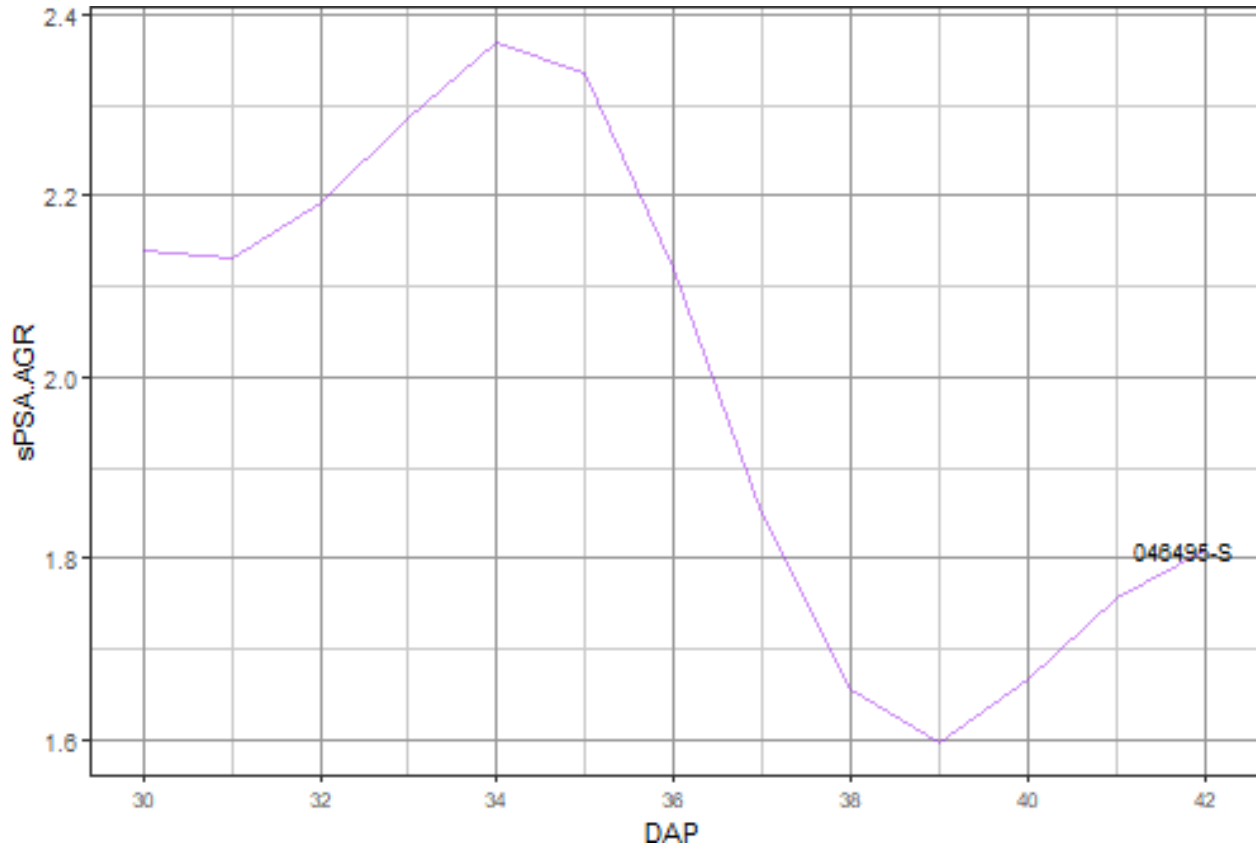
```

```

## Warning: `aes_string()` was deprecated in ggplot2 3.0.0.
## i Please use tidy evaluation idioms with `aes()`'.
## i See also `vignette("ggplot2-in-packages")` for more information.
## Removed 1 rows containing missing values (`geom_vline()`').
## This warning is displayed once every 8 hours.

```

```
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```



Step 5: Extract per-cart traits

A range of single-value plant responses are formed in Snapshot.ID.Tag order.

```
##### Set up intervals
```

```
DAP.endpts <- c(31,35,38,42)
DAP.starts <- c(31,35,31,38)
DAP.stops <- c(35,38,38,42)
DAP.mids <- (DAP.starts + DAP.stops)/2
suffices <- paste(DAP.starts, DAP.stops, sep = "to")
```

Step 5(a): Set up a data frame with factors only

```
cart.dat <- longi.dat[longi.dat$DAP == DAP.endpts[1],
                     c("Smarthouse", "Lane", "Position", "Snapshot.ID.Tag",
                       "cPosn", "cMainPosn",
                       "Zone", "cZone", "SHZone", "ZLane", "ZMainunit", "Subunit",
                       "Genotype.ID", "Treatment.1")]
cart.dat <- cart.dat[do.call(order, cart.dat), ]
```

Step 5(b): Get responses based on first and last date.

```

# Observation for first and last date
cart.dat <- cbind(cart.dat, getTimesSubset(data = longi.dat, responses = responses.image,
                                             times = "DAP", which.times = DAP.endpts[1],
                                             suffix = "first"))
cart.dat <- cbind(cart.dat, getTimesSubset(data = longi.dat, responses = responses.image,
                                             times = "DAP",
                                             which.times = DAP.endpts[length(DAP.endpts)],
                                             suffix = "last"))
cart.dat <- cbind(cart.dat, getTimesSubset(data = longi.dat, responses = "WUI.cum",
                                             times = "DAP",
                                             which.times = DAP.endpts[length(DAP.endpts)],
                                             suffix = "last"))

responses.smooth <- paste0("s", responses.image)
cart.dat <- cbind(cart.dat, getTimesSubset(data = longi.dat, responses = responses.smooth,
                                             times = "DAP", which.times = DAP.endpts[1],
                                             suffix = "first"))
cart.dat <- cbind(cart.dat, getTimesSubset(data = longi.dat, responses = responses.smooth,
                                             times = "DAP",
                                             which.times = DAP.endpts[length(DAP.endpts)],
                                             suffix = "last"))

# Growth rates over whole period.
(tottime <- DAP.endpts[length(DAP.endpts)] - DAP.endpts[1]) #= 11

```

```
## [1] 11
```

```

cart.dat <- within(cart.dat,
{
  PSA.AGR.full <- (PSA.last - PSA.first)/tottime
  PSA.RGR.full <- log(PSA.last / PSA.first)/tottime
})

# Calculate water index over whole period
cart.dat <- merge(cart.dat,
  byIndv4Intvl_WaterUse(data = longi.dat,
    water.use = "WU", response = "PSA",
    trait.types = c("WUI", "WUR", "WU"),
    times = "DAP",
    start.time = DAP.endpts[1],
    end.time = DAP.endpts[length(DAP.endpts)]),
  by = c("Snapshot.ID.Tag"))

```

Step 5(c): Add growth rates and water indices for intervals

```

# Growth rates for specific intervals from the smoothed data by differencing
for (r in responses.smooth)
{
  for (k in 1:length(suffices))

```

```

    {
      cart.dat <- merge(cart.dat,
                          byIndv4Intvl_GRsDiff(data = longi.dat, responses = r,
                                                times = "DAP",
                                                which.rates = c("AGR", "RGR"),
                                                start.time = DAP.starts[k],
                                                end.time = DAP.stops[k],
                                                suffix.interval = suffices[k]),
                          by = "Snapshot.ID.Tag")
    }
}

# Water indices for specific intervals from the unsmoothed and smoothed data
for (k in 1:length(suffices))
{
  cart.dat <- merge(cart.dat,
                     byIndv4Intvl_WaterUse(data = longi.dat,
                                            water.use = "WU", responses = "PSA",
                                            times = "DAP",
                                            trait.types = c("WU", "WUR", "WUI"),
                                            start.time = DAP.starts[k],
                                            end.time = DAP.stops[k],
                                            suffix.interval = suffices[k]),
                     by = "Snapshot.ID.Tag")
}

cart.dat <- with(cart.dat, cart.dat[order(Snapshot.ID.Tag), ])

```

Form continuous and interval SIITs

This experiment involved the extra step of calculating a measure of shoot ion-independent tolerance (SIIT) of pairs of plants, control and a salt-treated co-located plants.

Calculate continuous values

```

cols.retained <- c("Snapshot.ID.Tag", "Smarthouse", "Lane", "Position",
                    "DAP", "Snapshot.Time.Stamp", "Hour", "xDAP",
                    "Zone", "cZone", "SHZone", "ZLane", "ZMainunit",
                    "cMainPosn", "Genotype.ID")
responses.GR <- c("sPSA.AGR", "sPSA.AGR", "sPSA.RGR")
suffices.results <- c("diff", "SIIT", "SIIT")
responses.SIIT <- unlist(Map(paste, responses.GR, suffices.results, sep=". "))

longi.SIIT.dat <-
  twoLevelOpcreate(data = longi.dat, responses = responses.GR, suffices.treatment=c("C", "S"),
                    operations = c("-", "/", "/"), suffices.results = suffices.results,
                    columns.retained = cols.retained,
                    by = c("Smarthouse", "Zone", "ZMainunit", "DAP"))
longi.SIIT.dat <- with(longi.SIIT.dat,
                        longi.SIIT.dat[order(Smarthouse, Zone, ZMainunit, DAP), ])

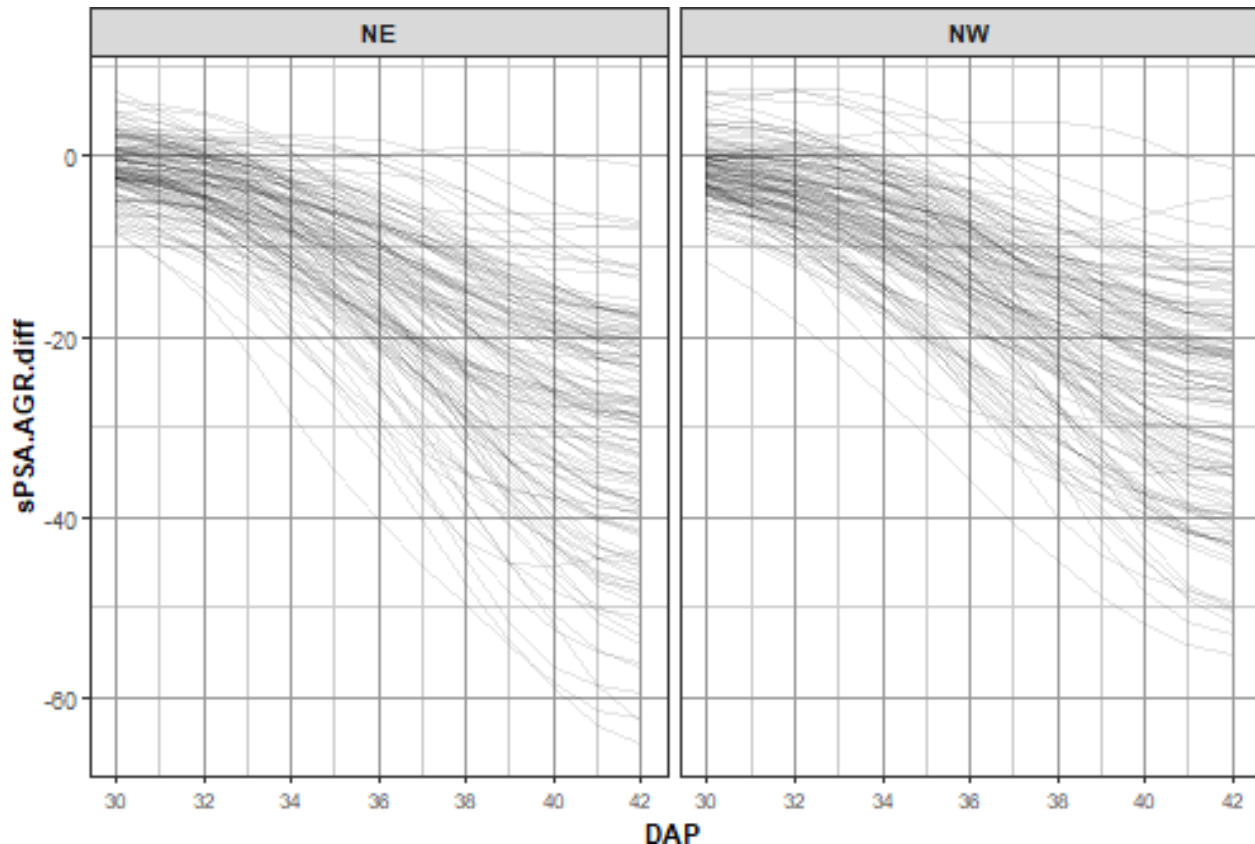
```

```

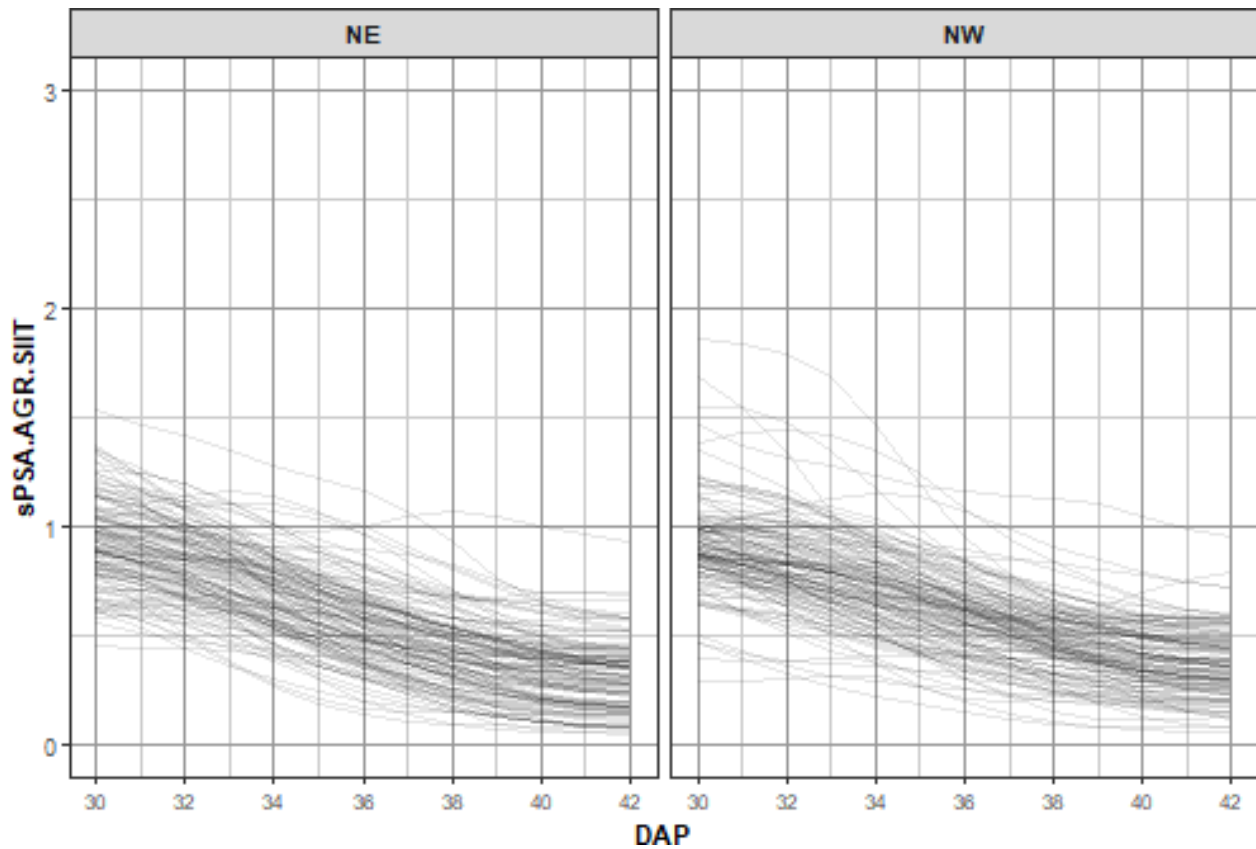
# Plot SIIT profiles
k <- 2
nresp <- length(responses.SIIT)
limits <- with(longi.SIIT.dat, list(c(min(sPSA.AGR.diff, na.rm=TRUE),
                                      max(sPSA.AGR.diff, na.rm=TRUE)),
                                      c(0,3),
                                      c(0,1.5)))
#Plots
for (k in 1:nresp)
{
  plt <- plotProfiles(data = longi.SIIT.dat, times = "DAP",
                        response = responses.SIIT[k],
                        y.title=responses.SIIT[k],
                        facet.x="Smarthouse", facet.y=".",
                        breaks.spacing.x = 2, printPlot=FALSE, )
  plt <- plt + geom_vline(xintercept=29, linetype="longdash", linewidth=1) +
    scale_y_continuous(limits=limits[[k]])
  print=plt)
}

```

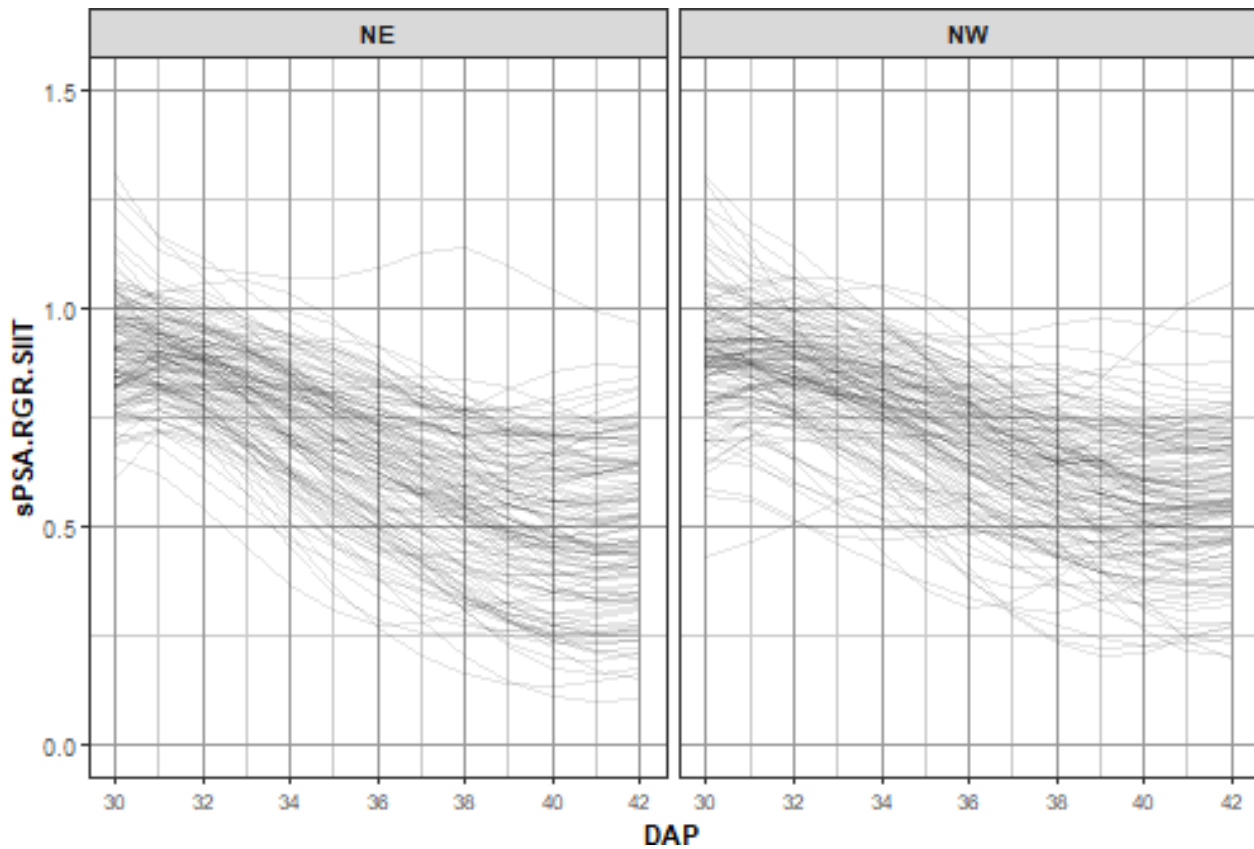
Warning: Removed 2 rows containing missing values ('geom_vline()').



Warning: Removed 2 rows containing missing values ('geom_vline()').



```
## Warning: Removed 2 rows containing missing values ('geom_vline()').
```



Calculate interval SIITs and check for large values for SIIT for Days 31to35

```

response <- "sPSA.RGR.31to35"
SIIT <- paste(response, "SIIT", sep=".") 
responses.SIITinterval <- as.vector(outer("sPSA.RGR", suffices, paste, sep="."))
 
cart.SIIT.dat <- twoLevelOpcreate(data = cart.dat, responses = responses.SIITinterval,
                                     suffices.treatment=c("C","S"),
                                     suffices.results="SIIT",
                                     columns.suffixed="Snapshot.ID.Tag")

tmp<-na.omit(cart.SIIT.dat)
print(summary(tmp[SIIT])) 

##  sPSA.RGR.31to35.SIIT
##  Min.   :0.4240
##  1st Qu.:0.7240
##  Median :0.8033
##  Mean   :0.7940
##  3rd Qu.:0.8720
##  Max.   :1.0789

big.SIIT <- with(tmp, tmp[tmp[SIIT] > 1.15, c("Snapshot.ID.Tag.C","Genotype.ID",
                                               paste(response,"C",sep="."),

```

```

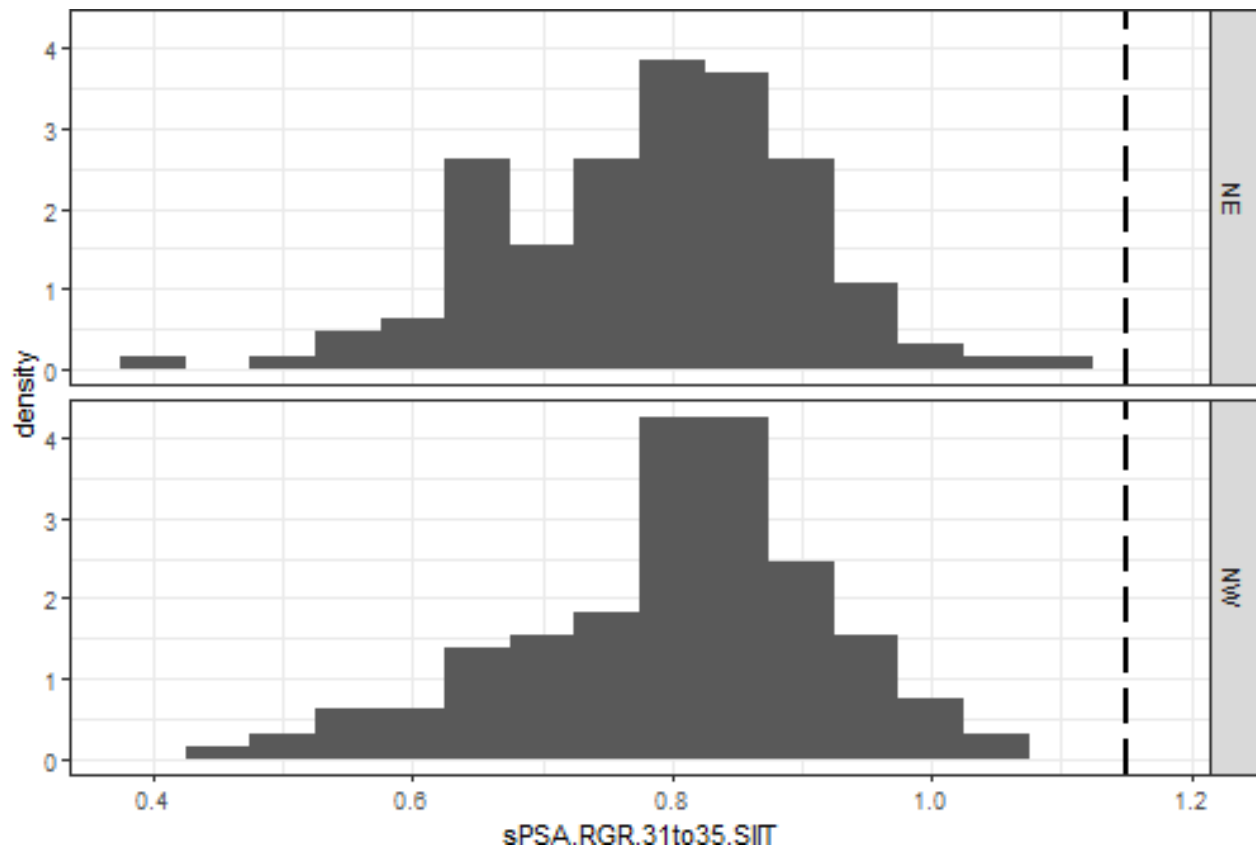
                paste(response, "S", sep=". "), SIIT)])
if (nrow(big.SIIT) > 1)
  big.SIIT <- big.SIIT[order(big.SIIT[SIIT]),]
print(big.SIIT)

## [1] Snapshot.ID.Tag.C      Genotype.ID           sPSA.RGR.31to35.C
## [4] sPSA.RGR.31to35.S     sPSA.RGR.31to35.SIIT
## <0 rows> (or 0-length row.names)

plt <- ggplot(tmp, aes_string(SIIT)) +
  geom_histogram(aes(y = ..density..), binwidth=0.05) +
  geom_vline(xintercept=1.15, linetype="longdash", linewidth=1) +
  theme_bw() + facet_grid(Smarthouse ~.)
print(plt)

## Warning: The dot-dot notation ('..density..') was deprecated in ggplot2 3.4.0.
## i Please use 'after_stat(density)' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.

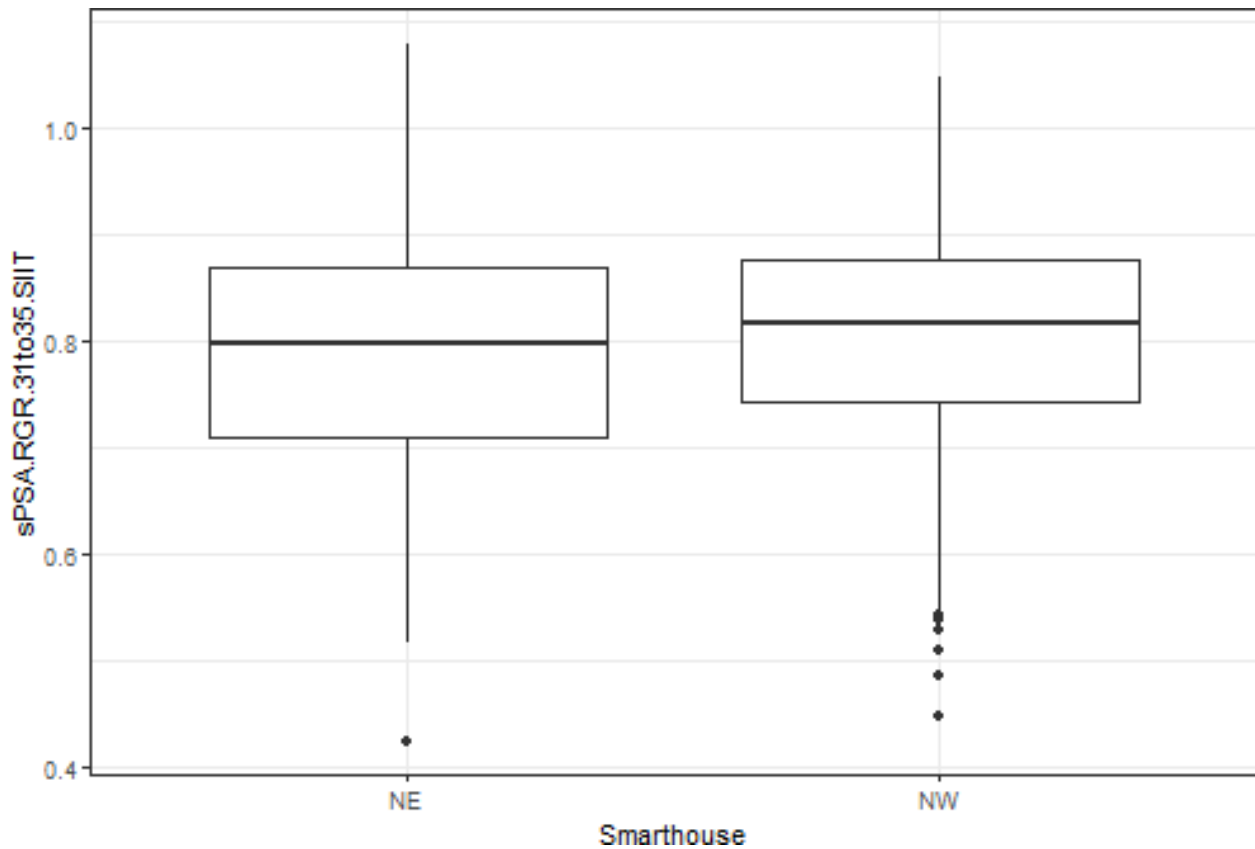
```



```

plt <- ggplot(tmp, aes_string(x="Smarthouse", y=SIIT)) +
  geom_boxplot() + theme_bw()
print(plt)

```



```
remove(tmp)
```

Save image

```
save.image("Rice.RData")
```

References

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