# ml<br/>4ec - Machine Learning for Eddy Covariance data

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2021-09-22

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# Set up

### 1.1 Apps

For this workshop, you need R and RStudio. Follow the links for downloading and installing these apps.

### 1.2 Libraries

Install missing packages for this tutorial.

```
list_pkgs <- c("caret", "recipes", "rsample", "tidyverse", "conflicted", "modelr", "forcats", "ya
new_pkgs <- list_pkgs[!(list_pkgs %in% installed.packages()[, "Package"])]
if (length(new_pkgs) > 0) install.packages(new_pkgs)
```

This book was compiled with the *bookdown* library and source files (RMarkdown), available on Github. Navigate there also for working on the exercises (Chapter 7) and using the solutions (Chapter 8).

This book is published under a GNU-GPL v3 license. Please cite this as:

Benjamin Stocker. (2021). ml4ec\_workshop: v1: Initial version (v1.0). Zenodo. https://doi.org/10.5281/zenodo.5521563

# Introduction

### 2.1 Learning objectives

After this workshop unit, you will be able to ...

- Understand how overfitting models can happen and how it can be avoided.
- Implement a typical workflow using a machine learning model for a supervised regression problem.
- Evaluate the power of the model

### 2.2 Some primers

Machine learning (ML) refers to a class of algorithms that generate statistical models of data. There are two main types of machine learning:

Unsupervised machine learning: Detecting patterns without prior specification.

**Supervised machine learning**: Model fitting by optimising predictions for a given target.

**Loss** is a function predicted and observed values derived from the validation set. It is minimised during model training.

# 2.3 Overfitting

This example is based on this example from scikit-learn.

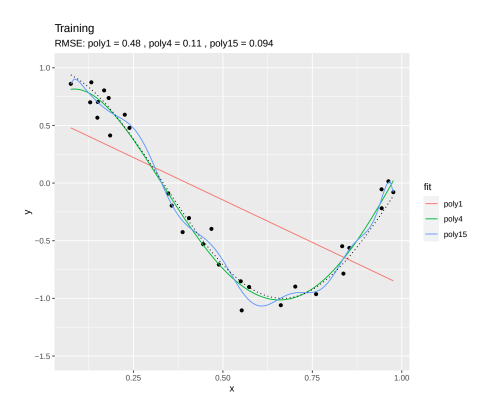
Machine learning (ML) may appear magical. The ability of ML algorithms to detect patterns and make predictions is fascinating. However, several challenges have to be met in the process of formulating, training, and evaluating the models. In this practical we will discuss some basics of supervised ML and how to achieve best predictive results.

In general, the aim of supervised ML is to find a model  $\hat{Y} = f(X)$  that is trained (calibrated) using observed relationships between a set of features (also known as predictors, or labels, or independent variables) X and the target variable Y. Note, that Y is observed. The hat on  $\hat{Y}$  denotes an estimate. Some algorithms can even handle predictions of multiple target variables simultaneously (e.g., neural networks). ML algorithms consist of (more or less) flexible mathematical models with a certain structure and set of parameters. At the simple extreme end of the model spectrum is the univariate linear regression. You may not want to call this a ML algorithm because there is no iterative learning involved. Nevertheless, also univariate linear regression provides a prediction  $\hat{Y} = f(X)$ , just like other (proper) ML algorithms do. The functional form of a linear regression is not particularly flexible (just a straight line for the best fit between predictors and targets) and it has only two parameters (slope and intercept). At the other extreme end are, for example, deep neural networks. They are extremely flexible, can learn highly non-linear relationships and deal with interactions between a large number of predictors. They also contain very large numbers of parameters, typically on the order of thousands. You can imagine that this allows these types of algorithms to very effectively learn from the data, but also bears the risk of overfitting.

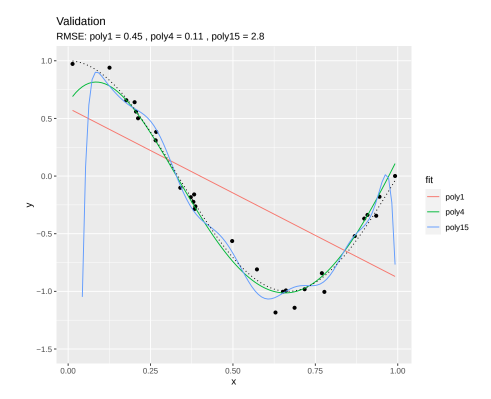
What is overfitting? The following example illustrates it. Let's assume that there is some true underlying relationship between a predictor x and the target variable y. We don't know this relationship (in the code below, this is  $true_fun()$ ) and the observations contain a (normally distributed) error  $(y = true_fun(x) + 0.1 * rnorm(n_samples))$ . Based on our training data  $(df_train)$ , we fit three polynomial models of degree 1, 4, and 15 to the observations. A polynomial of degree N is given by:

$$y = \sum_{n=0}^{N} a_n x^n$$

 $a_n$  are the coefficients, i.e., model parameters. The goal of the training is to get the coefficients  $a_n$ . From the above definition, the polynomial of degree 15 has 16 parameters, while the polynomial of degree 1 has two parameters (and corresponds to a simple linear regression). You can imagine that the polynomial of degree 15 is much more flexible and should thus yield the closest fit to the training data. This is indeed the case.

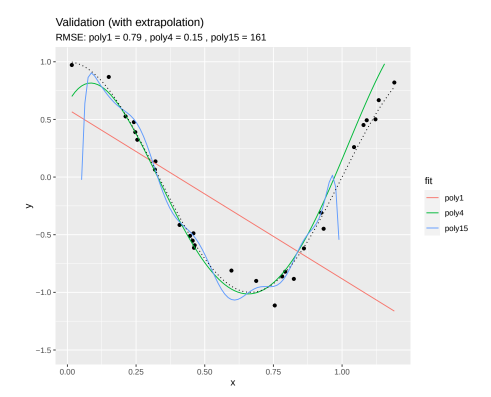


We can use the same fitted models on unseen data - the  $validation\ data$ . This is what's done below. Again, the same true underlying relationship is used, but we sample a new set of data points in x and add a new sample of errors on top of the true relationship.



You see that, using the validation set, we find that "poly4" actually performs the best - it has a much lower RMSE that "poly15". Apparently, "poly15" was overfitted. Apparently, it indeed used its flexibility to fit not only the shape of the true underlying relationship, but also the observation errors on top of it. This has obviously the implication that, when this model is used to make predictions for data that was not used for training (calibration), it will yield misguided predictions that are affected by the errors in the training set. In the above pictures we can also conclude that "poly1" was underfitted.

It gets even worse when applying the fitted polynomial models to data that extends beyond the range in x that was used for model training. Here, we're extending just 20% to the right.



You see that the RMSE for "poly15" literally explodes. The model is hopelessly overfitted and completely useless for prediction, although it looked like it fit the data best when we considered at the training results. This is a fundamental challenge in ML - finding the model with the best *generalisability*. That is, a model that not only fits the training data well, but also performs well on unseen data.

The phenomenon of fitting/overfitting as a function of the model "flexibility" is also referred to as bias vs. variance trade-off. The bias describes how well a model matches the training set (average error). A model with low bias will match the data set closely and vice versa. The variance describes how much a model changes when you train it using different portions of your data set. "poly15" has a high variance, but much of its variance is the result of misled training on observation errors. On the other extreme, "poly1" has a high bias. It's not affected by the noise in observations, but its predictions are also far off the observations. In ML, we are challenged to balance this trade-off. In Figure ?? you can see a schematic illustration of the bias-variance trade-off.

This chapter introduces the methods to achieve the best model generalisability and find the sweet spot between high bias and high variance. The steps to get there include the preprocessing of data, splitting the data into training and testing sets, and model training that "steers" the model towards what is considered a good model fit in terms of its generalisation power.

You have learned in video 6a about the basic setup of supervised ML, with input data containing the features (or predictors) X, predicted  $(\hat{Y})$  and observed target values (Y), also known as labels. In video 6b (title 6c: loss and it's minimization), you learned about the loss function which quantifies the agreement between Y and  $\hat{Y}$  and defines the objective of the model training. Here, you'll learn how all of this can be implemented in R. Depending on your application or research question, it may also be of interest to evaluate the relationships embodied in f(X) or to quantify the importance of different predictors in our model. This is referred to as model interpretation and is introduced in the respectively named subsection. Finally, we'll get into feature selection in the next Application session.

The topic of supervised machine learning methods covers enough material to fill two sessions. Therefore, we split this part in two. Model training, implementing the an entire modelling workflow, model evaluation and interpretation will be covered in the next session's tutorial (Supervised Machine Learning Methods II).

Of course, a plethora of algorithms exist that do the job of Y = f(X). Each of them has its own strengths and limitations. It is beyond the scope of this course to introduce a larger number of ML algorithms. Subsequent sessions will focus primarily on Artificial Neural Networks (ANN) - a type of ML algorithm that has gained popularity for its capacity to efficiently learn patterns in large data sets. For illustration purposes in this and the next chapter, we will briefly introduce two simple alternative "ML" methods, linear regression and K-nearest-neighbors. They have quite different characteristics and are therefore great for illustration purposes in this chapter.

## 2.4 Our modelling challenge

The environment determines ecosystem-atmosphere exchange fluxes of water vapour and CO2. Temporally changing mass exchange fluxes can be continuously measured with the eddy covariance technique, while abiotic variables (meteorological variables, soil moisture) can be measured in parallel. This offers an opportunity for building models that predict mass exchange fluxes from the environment.

In this workshop, we formulate a model for predicting ecosystem gross primary production (photosynthesis) from environmental covariates.

This is to say that GPP\_NT\_VUT\_REF is the target variable, and other available variables available in the dataset can be used as predictors.

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#### 2.5 Data

Data is provided here at daily resolution for a site ('CH-Dav') located in the Swiss alps (Davos). This is one of the longest-running eddy covariance sites globally and measures fluxes in a evergreen coniferous forest with cold winters and temperate and relatively moist summers.

For more information of the variables in the dataset, see FLUXNET 2015 website, and Pastorello et al., 2020 for a comprehensive documentation of variable definitions and methods.

#### 2.5.1 Available variables

- TIMESTAMP: Day of measurement.
- TA\_F: Air temperature. The meaning of suffix \_F is described in Pastorello et al., 2020.
- SW IN F: Shortwave incoming radiation
- LW IN F: Longwave incoming radiation
- VPD\_F: Vapour pressure deficit (relates to the humidity of the air)
- PA\_F: Atmospheric pressure
- P\_F: Precipitation
- WS\_F: Wind speed
- GPP\_NT\_VUT\_REF: Gross primary production the target variable
- NEE\_VUT\_REF\_QC: Quality control information for GPP\_NT\_VUT\_REF. Specifies the fraction of high-quality underlying high-frequency data from which the daily data is derived. 0.8 = 80% underlying high-quality data, remaining 20% of the high-frequency data is gap-filled.

### 2.6 More primers

#### 2.6.1 K-nearest neighbours

As the name suggests, the K-nearest neighbour (KNN) uses the k observations that are "nearest" to the new record for which we want to make a prediction. It then calculates their average (in regression) or most frequent value (in classification) as the prediction. "Nearest" is determined by some distance metric evaluated based on the values of the predictors. In our example (GPP\_NT\_VUT\_REF ~ .), KNN would determine the k days where conditions, given by our set of predictors, were most similar (nearest) to the day for which we seek a prediction. Then, it calculates the prediction as the average (mean) GPP value of these days. Determining "nearest" neighbors is commonly based on either the Euclidean or Manhattan distances between two data points  $x_a$  and  $x_b$ , considering all p predictors j.

Euclidean distance:

$$\sqrt{\sum_{j=1}^{p}(x_{a,j}-x_{b,j})^2}$$
 
$$\sum_{j=1}^{p}|x_{a,j}-x_{b,j}|$$

Manhattan distance:

$$\sum_{j=1}^{p} |x_{a,j} - x_{b,j}|$$

In two-dimensional space, the Euclidean distance measures the length of a straight line between two points (remember Pythagoras!). The Manhattan distance is called this way because it measures the distance you would have to walk to get from point a to point b in Manhattan, New York, where you cannot cut corners but have to follow a rectangular grid of streets. |x| is the positive value of x (|-x| = x).

KNN is a simple algorithm that uses knowledge of the "local" data structure for prediction. A drawback is that the model training has to be done for each prediction step and the computation time of the training increases with  $x \times p$ . KNNs are used, for example, to impute values (fill missing values) and have the advantage that predicted values are always within the range of observed values of the target variable.

#### 2.6.2 Random Forest

Random forest models are based on decision trees, where binary decisions for predicting the target's values are based on thresholds of the predictors' values. The depth of a decision tree refers to the number of such decisions. The deeper a tree, the more likely the model will overfit. Here are some links for more information on decision trees:

#### • Hands On Machine-Learning in R

Just as forests are made up by trees, random Forest models make use of random subsets of the original data and of available predictions and respective decision trees. Predictions are then made by averaging predictions of individual base learners (the decision trees). The number of predictors considered at each decision step is a tunable parameter (a hyperparameter, typically called  $m_{tru}$ ). Introducing this randomness is effective because decision trees tend to overfit and because of the wisdom of the crowd - i.e., the power of aggregating individual predictions with their random error (and without systematic bias) for generating accurate and relatively precise predictions.

Random forest models have gained particular popularity and are widely applied in environmental sciences not only for their power, but also for their ease of use. No preprocessing (centering, scaling) is necessary, they can deal with skewed data, and can effectively learn interactions between predictors.

Here are some links for more information on random forest:

• Hands On Machine-Learning in R

# Data splitting

### 3.1 Reading and wrangling data

Let's start by reading our data and apply few processing steps (wrangling).

There is a difference between data wrangling and pre-processing as part of the modelling workflow, which we will learn about in Chapter 4. Data wrangling can be considered to encompass the steps to prepare the data set prior to modelling, including, combining variables from different sources, removal of bad or missing data, and aggregating to the desired resolution or granularity (e.g., averaging over all time steps in a day, or over all replicates in a sample). See the Quartz Guide to Bad Data for an overview of how to deal with different types of bad data.

In contrast, *pre-processing* refers to the additional steps that are either required by the ML algorithm (e.g. centering and scaling for KNN or neural networks) or the transformation of variables guided by the resulting improvement of the predictive power of the ML model. In other words, pre-processing is part of the modelling workflow and includes all steps that apply transformations that use parameters derived from the data.

We are provided with a data file in the format of comma-separated-values (CSV), obtained through FLUXNET2015. It contains data from one site (CH-Dav) at a daily time step, and includes quality control information for each variable.

Let's read the data, select relevant variables, convert the time stamp column to a time object and interpret missing values (encoded -9999 in the file).

```
library(tidyverse)

ddf <- read_csv("./data/FLX_CH-Dav_FLUXNET2015_FULLSET_DD_1997-2014_1-3.csv") %>%
```

```
## select only the variables we are interested in
select(TIMESTAMP,
       GPP_NT_VUT_REF,
                          # the target
       NEE_VUT_REF_QC,
                          # quality control info
       ends_with("_F"), # includes all all meteorological variables
       -contains("JSB")
                          # weird useless variable
       ) %>%
## convert to a nice date object
mutate(TIMESTAMP = lubridate::ymd(TIMESTAMP)) %>%
## set all -9999 to NA
na if(-9999) %>%
## drop QC variables (no longer needed), except NEE_VUT_REF_QC
select(-ends_with("_QC"), NEE_VUT_REF_QC)
```

If the style of the code above looks unfamiliar - this is the **tidyverse**. The tidyverse is a R syntax "dialect" and a collection of R functions and packages. They share the structure of arguments and function return values than can be combined to a chain by the %>% ("pipe") operator. For this, the output of each function is a data frame which is "piped" to the next function, and each function takes a data frame as input. What is piped into a function takes the place of the first argument, normally provided inside the brackets. This enables ease with typical data wrangling and visualization tasks (**ggplot2** is part of the tidyverse). This tutorial is generally written using tidyverse packages and code syntax.

The column NEE\_VUT\_REF\_QC provides information about the fraction of gap-filled half-hourly data used to calculate daily aggregates. Let's use only GPP\_NT\_VUT\_REF data, where at least 80% of the underlying half-hourly data was good quality measured data, and not gap-filled. Make sure to not actually remove the respective rows, but rather replace values with NA.

```
ddf <- ddf %>%
  mutate(GPP_NT_VUT_REF = ifelse(NEE_VUT_REF_QC < 0.8, NA, GPP_NT_VUT_REF))</pre>
```

At this stage, we won't use NEE\_VUT\_REF\_QC any longer. So we can drop it.

```
ddf <- ddf %>%
select(-NEE_VUT_REF_QC)
```

### 3.2 Splitting into testing and training sets

The introductory example impressively demonstrated the importance of validating the fitted model with data that was *not* used for training. Thus, we can test the model's *generalisability*. The essential step that enables us to assess the model's *generalization error* is to hold out part of the data from training, and set it aside (leaving it absolutely untouched) for *testing*.

There is no fixed rule for how much data are to be used for training and testing, respectively. We have to balance the trade-off between:

- Spending too much data for training will leave us with too little data for testing and the test results may not be robust. In this case, the sample size for getting robust validation statistics is not sufficiently large and we don't know for sure whether we are safe from an over-fit model.
- Spending too much data for validation will leave us with too little data for training. In this case, the ML algorithm may not be successful at finding real relationships due to insufficient amounts of training data.

Typical splits are between 60-80% for training. However, in cases where the number of data points is very large, the gains from having more training data are marginal, but come at the cost of adding to the already high computational burden of model training.

In environmental sciences, the number of predictors is often smaller than the sample size (p < n), because its typically easier to collect repeated observations of a particular variable than to expand the set of variables being observed. Nevertheless, in cases where the number p gets large, it is important, and for some algorithms mandatory, to maintain p < n for model training.

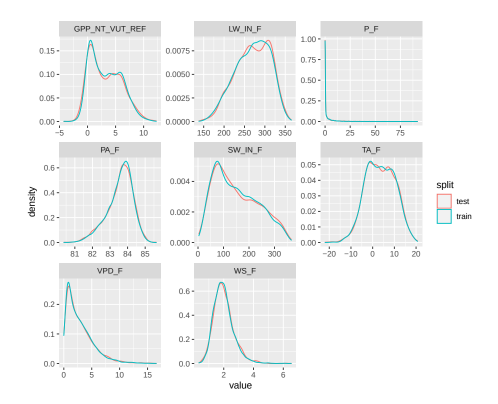
An important aspect to consider when splitting the data is to make sure that all "states" of the system for which we have data are approximately equally represented in training and testing sets. This is to make sure that the algorithm learns relationships f(X) also under rare conditions X, for example meteorological extreme events.

Several alternative functions for the data splitting step are available from different packages in R. We will use the **rsample** package as it allows to additionally make sure that data from the full range of a given variable's values (VPD\_F in the example below) are well covered in both training and testing sets.

```
library(rsample)
set.seed(123) # for reproducibility
split <- initial_split(ddf, prop = 0.7, strata = "VPD_F")
ddf_train <- training(split)
ddf_test <- testing(split)</pre>
```

Plot the distribution of values in the training and testing sets.

```
ddf_train %>%
  mutate(split = "train") %>%
  bind_rows(ddf_test %>%
    mutate(split = "test")) %>%
    pivot_longer(cols = 2:9, names_to = "variable", values_to = "value") %>%
  ggplot(aes(x = value, y = ..density.., color = split)) +
  geom_density() +
  facet_wrap(~variable, scales = "free")
```



# Pre-processing

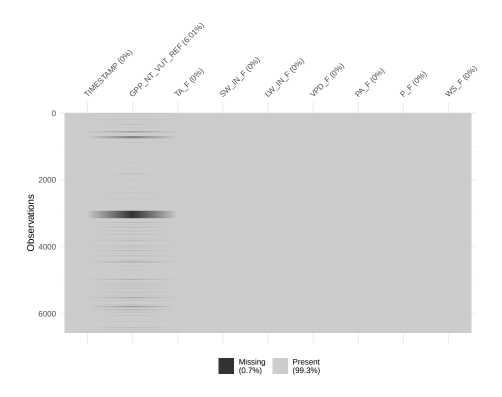
Skewed data, outliers, and values covering multiple orders of magnitude can create difficulties for certain ML algorithms, e.g., or K-nearest neighbours, or neural networks. Other algorithms, like tree-based methods (e.g., Random Forest), are more robust against such issues.

### 4.1 Dealing with missingness and bad data

Several ML algorithms require missing values to be removed. That is, if any of the cells in one row has a missing value, the entire cell gets removed. Data may be missing for several reasons. Some yield random patterns of missing data, others not. In the latter case, we can speak of *informative missingness* (Kuhn & Johnson, 2003) and its information can be used for predictions. For categorical data, we may replace such data with "none" (instead of NA), while randomly missing data may be dropped altogether. Some ML algorithms (mainly tree-based methods, e.g., random forest) can handle missing values. However, when comparing the performance of alternative ML algorithms, they should be tested with the same data and removing missing data should be done beforehand.

Visualising missing data is essential for making decisions about dropping rows with missing data versus removing predictors from the model (which would imply too much data removal). The cells with missing data in a data frame can be early visualised e.g. with vis\_miss() from the *visdat* package.

```
library(visdat)
vis_miss(
    ddf,
    cluster = FALSE,
    warn_large_data = FALSE
)
```



The question about what is "bad data" and whether or when it should be removed is often critical. Such decisions are important to keep track of and should be reported as transparently as possible in publications. In reality, where the data generation process may start in the field with actual human beings writing notes in a lab book, and where the human collecting the data is often not the same as the human writing the paper, it's often more difficult to keep track of such decisions. As a general principle, it is advisable to design data records such that decisions made during its process remain transparent throughout all stages of the workflow and that sufficient information be collected to enable later revisions of particularly critical decisions.

#### 4.2 Standardization

Several algorithms explicitly require data to be standardized. That is, values of all predictors vary within a comparable range. The necessity of this step becomes obvious when considering neural networks, the activation functions of

each node have to deal with standardized inputs. In other words, inputs have to vary over the same range, expecting a mean of zero and standard deviation of one.)

To get a quick overview of the distribution of all variables (columns) in our data frame, we can use the skimr package.

library(skimr)
knitr::kable(skim(ddf))

skim_type	skim_variable	n_missing	complete_rate	Date.min	Date.max	Date.median	D
Date	TIMESTAMP	0	1.0000000	1997-01-01	2014-12-31	2005-12-31	
numeric	GPP_NT_VUT_REF	395	0.9399148	NA	NA	NA	
numeric	TA_F	0	1.0000000	NA	NA	NA	
numeric	SW_IN_F	0	1.0000000	NA	NA	NA	
numeric	LW_IN_F	0	1.0000000	NA	NA	NA	
numeric	VPD_F	0	1.0000000	NA	NA	NA	
numeric	PA_F	0	1.0000000	NA	NA	NA	
numeric	P_F	0	1.0000000	NA	NA	NA	
numeric	WS F	0	1.0000000	NA	NA	NA	

We see for example, that typical values of LW\_IN\_F are by a factor 100 larger than values of VPD\_F. KNN uses the distance from neighbouring points for predictions. Obviously, in this case here, any distance would be dominated by LW\_IN\_F and distances in the "direction" of VPD\_F, even when relatively large, would not be influential, neither for a Euclidean nor a Manhattan distance (see 2). In neural networks, activation functions take values in a given range (0-1). Thus, for both algorithms, data has to be standardized prior to model training.

Standardization is done, for example, by dividing each variable, that is all values in one column, by the standard deviation of that variable, and then subtracting its mean. This way, the resulting standardized values are centered around 0, and scaled such that a value of 1 means that the data point is one standard deviation above the mean of the respective variable (column). When applied to all predictors individually, the absolute values of their variations can be directly compared and only then it can be meaningfully used for determining the distance.

Standardization can be done not only by centering and scaling (as described above), but also by *scaling to within range*, where values are scaled such that the minimum value within each variable (column) is 0 and the maximum is 1.

In order to avoid *data leakage*, centering and scaling has to be done separately for each split into training and validation data (more on that later). In other words, don't center and scale the entire data frame with the mean and standard deviation derived from the entire data frame, but instead center and scale with mean and standard deviation derived from the training portion of the data, and apply that also to the validation portion, when evaluating.

The *caret* package takes care of this. The R package **caret** provides a unified interface for using different ML algorithms implemented in separate packages. The preprocessing steps applied with each resampling fold can be specified using the function preProcess(). More on resampling in Chapter 6.

```
library(caret)
pp <- preProcess(ddf_train, method = c("center", "scale"))</pre>
```

As seen above for the feature engineering example, this does not return a standardized version of the data frame ddf. Rather, it returns the information that allows us to apply the same standardization also to other data sets. In other words, we use the distribution of values in the data set to which we applied the function to determine the centering and scaling (here: mean and standard deviation).

### 4.3 More pre-processing

Depending on the algorithm and the data, additional pre-processing steps may be required. You can find more information about this in the great and freely available online tutorial Hands-On Machine Learning in R.

One such additional pre-processing step is *imputation*, where missing values are imputed (gap-filled), for example by the mean of each variable respectively. Also imputation is prone to cause data leakage and must therefore be implemented as part of the resampling and training workflow. The **recipes** package offers a great way to deal with imputation (and also all other pre-processing steps). Here is a link to learn more about it.

# Model formulation

The aim of supervised ML is to find a model  $\hat{Y} = f(X)$  so that  $\hat{Y}$  agrees well with observations Y. We typically start with a research question where Y is given - naturally - by the problem we are addressing and we have a data set at hand where one or multiple predictors (or features) X are recorded along with Y. From our data, we have information about how GPP (ecosystem-level photosynthesis) depends on set of abiotic factors, mostly meteorological measurements.

#### 5.1 Formula notation

In R, it is common to use the *formula* notation to specify the target and predictor variables. You have probably encountered formulas before, e.g., for a linear regression using the lm() function. To specify a linear regression model for GPP\_NT\_VUT\_REF with three predictors SW\_F\_IN, VPD\_F, and TA\_F, we write:

```
lm(GPP_NT_VUT_REF ~ SW_F_IN + VPD_F + TA_F, data = ddf)
```

### 5.2 The generic train()

Actually, the way we formulate a model is independent of the algorithm, or engine that takes care of fitting f(X). As mentioned in Chapter 4 the R package caret provides a unified interface for using different ML algorithms implemented in separate packages. In other words, it acts as a wrapper for multiple different model fitting, or ML algorithms. This has the advantage that it unifies the interface (the way arguments are provided). caret also provides implementations for a set of commonly used tools for data processing, model training, and

evaluation. We'll use caret for model training with the function train() (more on model training in Chapter 6). Note however, that using a specific algorithm, which is implemented in a specific package outside caret, also requires that the respective package be installed and loaded. Using caret for specifying the same linear regression model as above, the base-R lm() function, can be done with caret in a generalized form as:

```
library(caret)
train(
  form = GPP_NT_VUT_REF ~ SW_IN_F + VPD_F + TA_F,
  data = ddf %>% drop_na(),
  method = "lm"
)
```

```
## Linear Regression
##
## 6179 samples
##
      3 predictor
## No pre-processing
## Resampling: Bootstrapped (25 reps)
## Summary of sample sizes: 6179, 6179, 6179, 6179, 6179, ...
## Resampling results:
##
##
    RMSE
               Rsquared
                          MAE
##
    1.599712 0.6646581 1.241035
##
## Tuning parameter 'intercept' was held constant at a value of TRUE
```

Of course, this is an overkill compared to just writing lm(...). But the advantage of the unified interface is that we can simply replace the method argument to use a different ML algorithm. For example, to use a random forest model implemented by the ranger package, we can write:

```
## do not run
train(
  form = GPP_NT_VUT_REF ~ SW_F_IN + VPD_F + TA_F,
  data = ddf,
  method = "ranger",
  ...
)
```

The ... hints at the fact that there are a few more arguments to be specified for training a random forest model with ranger. More on that in Chapter 6.

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### 5.3 Recipes

The **recipes** package provides another way to specify the *formula* and preprocessing steps in one go and is compatible with caret's **train()** function. For the same formula as above, and an example where the data **ddf\_train** is to be centered and scaled, we can specify the "recipe" using the *tidyverse*-style pipe operator as:

```
library(recipes)
pp <- recipe(GPP_NT_VUT_REF ~ SW_IN_F + VPD_F + TA_F, data = ddf_train) %>%
    step_center(all_numeric(), -all_outcomes()) %>%
    step_scale(all_numeric(), -all_outcomes())
```

The first line assigns *roles* to the different variables. GPP\_NT\_VUT\_REF is an *outcome* (in "recipes speak"). Then, we used selectors to apply the recipe step to several variables at once. The first selector, all\_numeric(), selects all variables that are either integers or real values. The second selector, -all\_outcomes() removes any outcome (target) variables from this recipe step.

The object pp can then be supplied to train() as its first argument:

```
## do not run
train(
   pp,
   data = ddf_train,
   method = "ranger",
   ...
)
```

As seen above for the pre-processing example, this does not return a standardized version of the data frame ddf\_train, but rather the information that allows us to apply the same standardization also to other data sets.

# Model training

Model training in supervised ML is guided by the match (or mismatch) between the predicted and observed target variable(s), that is, between  $\hat{Y}$  and Y. The loss function quantifies this mismatch  $(L(\hat{Y},Y))$ , and the algorithm takes care of progressively reducing the loss during model training. Let's say the ML model contains two parameters and predictions can be considered a function of the two  $(\hat{Y}(w_1,w_2))$ . Y is actually constant. Thus, the loss function is effectively a function  $L(w_1,w_2)$ . Therefore, we can consider the model training as a search of the parameter space of the machine learning model  $(w_1,w_2)$  to find the minimum of the loss. Common loss functions are the root mean square error (RMSE), or the mean square error (MSE), or the mean absolute error (MAE). Loss minimization is a general feature of ML model training.

Model training is implemented in R for different algorithms in different packages. Some algorithms are even implemented by multiple packages (e.g., nnet and neuralnet for artificial neural networks). As described in Chapter 4, the caret package provides "wrappers" that handle a large selection of different ML model implementations in different packages with a unified interface (see here for an overview of available models). The caret function train() is the centre piece. Its argument metric specifies the loss function and defaults to the RMSE for regression models and accuracy for classification (see sub-section on metrics below).

### 6.1 Hyperparameter tuning

Practically all ML algorithms have some "knobs" to turn in order to achieve efficient model training and predictive performance. Such "knobs" are the *hyperparameters*. What these knobs are, depends on the ML algorithm.

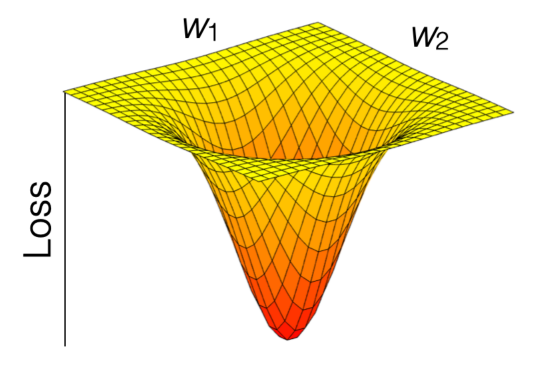


Figure 6.1: Visualization of a loss function as a plane spanned by the two parameters  $w_1$  and  $w_2$ .

For KNN, this is k - the number of neighbours to consider for determining distances. There is always an optimum k. Obviously, if k=n, we consider all observations as neighbours and each prediction is simply the mean of all observed target values Y, irrespective of the predictor values. This cannot be optimal and such a model is likely underfit. On the other extreme, with k=1, the model will be strongly affected by the noise in the single nearest neighbour and its generalisability will suffer. This should be reflected in a poor performance on the validation data.

For random forests from the **ranger** package, hyperparameters are:

- mtry: the number of variables to consider to make decisions, often taken as p/3, where p is the number of predictors.
- min.node.size: the number of data points at the "bottom" of each decision tree
- splitrule: the function applied to data in each branch of a tree, used for determining the goodness of a decision

Hyperparameters usually have to be "tuned". The optimal setting depends on the data and can therefore not be known *a priori*.

In **caret**, hyperparameter tuning is implemented as part of the **train()** function. Values of hyperparameters to consider are to be specified by the argument **tuneGrid**, which takes a data frame with column(s) named according to the name(s) of the hyperparameter(s) and rows for each combination of hyperparameters to consider.

Here, expand.grid() is used to provide a data frame with all combinations of values provided by individual vectors.

### 6.2 Resampling

The goal of model training is to achieve the best possible model generalisability. That is, the best possible model performance when predicting to data that was not used for training - the test data. Resampling mimicks the comparison of predictions to the test data. Instead of using all training data, the training data is resampled into a number further splits into pairs of training and validation data. Model training is then guided by minimising the average loss determined on each resample of the validation data. Having multiple resamples (multiple folds of training-validation splits) avoids the loss minimization from being misguided by random peculiarities in the training and/or validation data.

A common resampling method is k-fold  $cross\ validation$ , where the training data is split into k equally sized subsets (folds). Then, there will be k iterations, where each fold is used for validation once (while the remaining folds are used for training). An extreme case is leave-one-out  $cross\ validation$ , where k corresponds to the number of data points.

Fold 1 Fold 2 **Train** Test Train Test Full set of **Train Train** training data Train **Train** Train **Train** 

To do a k-fold cross validation during model training in R, we don't have to implement the loops around folds ourselves. The resampling procedure can be specified in the **caret** function train() with the argument trControl. The object that this argument takes is the output of a function call to trainControl(). This can be implemented in two steps. For example, to do a 10-fold cross-validation, we can write:

In certain cases, data points stem from different "groups", and generalisability across groups is critical. In such cases, data from a given group must not be used both in the training and validation sets. Instead, splits should be made along group delineations. The *caret* function <code>groupKFold()</code> offers the solution for this case.

# Exercises

Now that you are familiar with the basic steps for supervised machine learning, you can get your hands on the data yourself and implement code for addressing the modelling task outlined in Chapter ??.

### 7.1 Reading and cleaning

Read the CSV file "./data/FLX\_CH-Dav\_FLUXNET2015\_FULLSET\_DD\_1997-2014\_1-3.csv", select all variables with name ending with "\_F", the variables "TIMESTAMP", "GPP\_NT\_VUT\_REF", and "NEE\_VUT\_REF\_QC", and drop all variables that contain "JSB" in their name. Then convert the variable "TIMESTAMP" to a date-time object with the function ymd() from the *lubridate* package, and interpret all values -9999 as missing values. Then, set all values of "GPP\_NT\_VUT\_REF" to missing if the corresponding quality control variable indicates that less than 90% are measured data points. Finally, drop the variable "NEE\_VUT\_REF\_QC" - we won't use it anymore.

## write your code here

### 7.2 Data splitting

Split the data a training and testing set, that contain 70% and 30% of the total available data, respectively.

## write your code here

#### 7.3 Linear model

#### 7.3.1 Training

Fit a linear regression model using the base-R function lm() and the training set. The target variable is "GPP\_NT\_VUT\_REF", and predictor variables are all available meterological variables in the dataset. Answer the following questions:

- What is the  $R^2$  of predicted vs. observed "GPP\_NT\_VUT\_REF"?
- Is the linear regression slope significantly different from zero for all predictors?
- Is a linear regression model with "poor" predictors removed better supported by the data than a model including all predictors?

#### ## write your code here

Use caret and the function train() for fitting the same linear regression model (with all predictors) on the same data. Does it yield identical results as using lm() directly? You will have to set the argument trControl accordingly to avoid resampling, and instead fit the model on the all data in ddf\_train. You can use summary() also on the object returned by the function train().

#### ## write your code here

#### 7.3.2 Prediction

With the model containing all predictors and fitted on  $ddf_{train}$ , make predictions using first  $ddf_{train}$  and then  $ddf_{test}$ . Compute the  $R^2$  and the root-mean-square error, and visualise modelled vs. observed values to evaluate both predictions.

Do you expect the linear regression model trained on ddf\_train to predict substantially better on ddf\_train than on ddf\_test? Why (not)?

#### Hints:

- To calculate predictions, use the generic function predict() with the argument newdata = ....
- The  $R^2$  can be extracted from the model object as summary(model\_object)\$r.squared, or is (as the RMSE) given in the metrics data frame returned by metrics() from the yardstick library.
- For visualisation the model performance, consider a scatterplot, or (better) a plot that reveals the density of overlapping points. (We're plotting information from over 4000 data points here!)

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## write your code here

## 7.4 KNN

#### 7.4.1 Check data

## write your code here

The variable PA\_F looks weird and was not significant in the linear model. Therefore, we won't use it for the models below.

## 7.4.2 Training

Fit two KNN models on  $ddf_{train}$  (excluding "PA\_F"), one with k=2 and one with k=30, both without resampling. Use the RMSE as the loss function. Center and scale data as part of the pre-processing and model formulation specification using the function recipe().

## write your code here

#### 7.4.3 Prediction

With the two models fitted above, predict "GPP\_NT\_VUT\_REF" for both and training and the testing sets, and evaluate them as above (metrics and visualisation).

Which model do you expect to perform better on the training set and which to perform better on the testing set? Why? Do you find evidence for overfitting in any of the models?

## write your code here

## 7.4.4 Sample hyperparameters

Train a KNN model with hyperparameter (k) tuned, and with five-fold cross validation, using the training set. As the loss function, use RMSE. Sample the following values for k: 2, 5, 10, 15, 18, 20, 22, 24, 26, 30, 35, 40, 60, 100. Visualise the RMSE as a function of k.

Hint:

• The visualisation of cross-validation results can be visualised with the plot(model\_object) of ggplot(model\_object).

## write your code here

## 7.5 Random forest

## 7.5.1 Training

Fit a random forest model with ddf\_train and all predictors excluding "PA\_F" and five-fold cross validation. Use RMSE as the loss function.

Hints:

- Use the package ranger which implements the random forest algorithm.
- See here for information about hyperparameters available for tuning with caret.
- Set the argument savePredictions = "final" of function trainControl().

## write your code here

#### 7.5.2 Prediction

Evaluate the trained model on the training and on the test set, giving metrics and a visualisation as above.

How are differences in performance to be interpreted? Compare the performances of linear regression, KNN, and random forest, considering the evaluation on the test set.

## write your code here

Show the model performance (metrics and visualisation) on the validation sets all cross validation folds combined.

Do you expect it to be more similar to the model performance on the training set or the testing set in the evaluation above? Why?

## write your code here

# Chapter 8

# **Solutions**

Now that you are familiar with the basic steps for supervised machine learning, you can get your hands on the data yourself and implement code for addressing the modelling task outlined in Chapter ??.

## 8.1 Reading and cleaning

Read the CSV file "./data/FLX\_CH-Dav\_FLUXNET2015\_FULLSET\_DD\_1997-2014\_1-3.csv", select all variables with name ending with "\_F", the variables "TIMESTAMP", "GPP\_NT\_VUT\_REF", and "NEE\_VUT\_REF\_QC", and drop all variables that contain "JSB" in their name. Then convert the variable "TIMESTAMP" to a date-time object with the function ymd() from the *lubridate* package, and interpret all values -9999 as missing values. Then, set all values of "GPP\_NT\_VUT\_REF" to missing if the corresponding quality control variable indicates that less than 90% are measured data points. Finally, drop the variable "NEE\_VUT\_REF\_QC" - we won't use it anymore.

```
mutate(TIMESTAMP = lubridate::ymd(TIMESTAMP)) %>%

## set all -9999 to NA
na_if(-9999) %>%

## drop QC variables (no longer needed), except NEE_VUT_REF_QC
select(-ends_with("_QC"), NEE_VUT_REF_QC) %>%

mutate(GPP_NT_VUT_REF = ifelse(NEE_VUT_REF_QC < 0.9, NA, GPP_NT_VUT_REF)) %>%
select(-NEE_VUT_REF_QC)
```

## 8.2 Data splitting

Split the data a training and testing set, that contain 70% and 30% of the total available data, respectively.

```
library(rsample)
set.seed(1982) # for reproducibility
split <- initial_split(ddf, prop = 0.7)
ddf_train <- training(split)
ddf_test <- testing(split)</pre>
```

## 8.3 Linear model

#### 8.3.1 Training

Fit a linear regression model using the base-R function 1m() and the training set. The target variable is "GPP\_NT\_VUT\_REF", and predictor variables are all available meterological variables in the dataset. Answer the following questions:

- What is the  $R^2$  of predicted vs. observed "GPP\_NT\_VUT\_REF"?
- Is the linear regression slope significantly different from zero for all predictors?
- Is a linear regression model with "poor" predictors removed better supported by the data than a model including all predictors?

```
## fit linear regression model
linmod_baser <- lm(
  form = GPP_NT_VUT_REF ~ .,
  data = ddf_train %>%
     drop_na() %>%
```

```
select(-TIMESTAMP)
)
## show summary information of the model
summary(linmod_baser)
##
## Call:
## lm(formula = GPP_NT_VUT_REF ~ ., data = ddf_train %>% drop_na() %>%
##
      select(-TIMESTAMP))
##
## Residuals:
             1Q Median
##
     Min
                           3Q
                                 Max
## -5.1941 -1.0393 -0.1299 0.8859 7.7205
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.9570704 3.4840107 -0.275 0.783558
## TA F
             ## SW_IN_F
             0.0137226  0.0003914  35.064  < 2e-16 ***
             0.0207311 0.0012040 17.219 < 2e-16 ***
## LW_IN_F
## VPD F
            ## PA_F
            -0.0402727 0.0404675 -0.995 0.319704
## P F
            ## WS_F
            ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.549 on 4135 degrees of freedom
## Multiple R-squared: 0.6904, Adjusted R-squared: 0.6899
## F-statistic: 1317 on 7 and 4135 DF, p-value: < 2.2e-16
## Models can be compared, for example by the Bayesian Information Criterion (BIC).
## It penalizes complex models and is optimal (lowest) where the trade-off between
## explanatory power on the training set vs. number of predictors is best. The BIC
## tends to be more conservative than the AIC.
BIC(linmod_baser)
```

#### ## [1] 15450.89

The variable PA\_F was not significant in the linear model. Therefore, we won't use it for the models below.

```
## Fit an lm model on the same data, but with PA_F removed.
linmod_baser_nopaf <- lm(</pre>
  form = GPP_NT_VUT_REF ~ .,
 data = ddf_train %>%
    drop_na() %>%
    select(-TIMESTAMP, -PA_F)
)
## The R-squared is slightly lower here (0.6903) than in the model with all predictors
summary(linmod_baser_nopaf)
##
## Call:
## lm(formula = GPP_NT_VUT_REF ~ ., data = ddf_train %>% drop_na() %>%
       select(-TIMESTAMP, -PA_F))
##
##
## Residuals:
      Min
               1Q Median
                                3Q
                                       Max
## -5.1849 -1.0390 -0.1306 0.8867 7.7594
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -4.4102425 0.3135013 -14.068 < 2e-16 ***
## TA_F
               0.1595814 0.0101445
                                     15.731 < 2e-16 ***
## SW_IN_F
               0.0136923 0.0003902
                                     35.093 < 2e-16 ***
## LW_IN_F
               0.0210424 0.0011626
                                     18.099 < 2e-16 ***
                                     -7.203 6.98e-13 ***
## VPD_F
              -0.1400240
                          0.0194405
## P_F
              -0.0212661 0.0046147
                                     -4.608 4.18e-06 ***
## WS_F
              -0.1343263 0.0391409
                                     -3.432 0.000605 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.549 on 4136 degrees of freedom
## Multiple R-squared: 0.6903, Adjusted R-squared: 0.6899
## F-statistic: 1537 on 6 and 4136 DF, p-value: < 2.2e-16
```

## ... but the BIC is clearly lower, indicating that the model with PA\_F removed is be BIC(linmod\_baser\_nopaf)

```
## [1] 15443.55
```

Use caret and the function train() for fitting the same linear regression model (with all predictors) on the same data. Does it yield identical results as using lm() directly? You will have to set the argument trControl accordingly to

library(caret)

avoid resampling, and instead fit the model on the all data in ddf\_train. You can use summary() also on the object returned by the function train().

```
linmod_caret <- train(</pre>
 form = GPP NT VUT REF ~ .,
 data = ddf_train %>%
   drop_na() %>%
   select(-TIMESTAMP),
 method = "lm",
 trControl = trainControl(method = "none")
summary(linmod_caret)
##
## Call:
## lm(formula = .outcome ~ ., data = dat)
##
## Residuals:
      Min
             1Q Median
                           3Q
                                 Max
## -5.1941 -1.0393 -0.1299 0.8859 7.7205
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.9570704 3.4840107 -0.275 0.783558
## TA F
             ## SW IN F
             0.0137226  0.0003914  35.064  < 2e-16 ***
## LW IN F
             0.0207311 0.0012040 17.219 < 2e-16 ***
## VPD F
            -0.1420886   0.0195509   -7.268   4.35e-13 ***
## PA F
            ## P F
            -0.0214314 0.0046177
                                -4.641 3.57e-06 ***
## WS_F
            ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

#### 8.3.2 Prediction

With the model containing all predictors and fitted on  $ddf_{train}$ , make predictions using first  $ddf_{train}$  and then  $ddf_{test}$ . Compute the  $R^2$  and the

## Residual standard error: 1.549 on 4135 degrees of freedom
## Multiple R-squared: 0.6904, Adjusted R-squared: 0.6899
## F-statistic: 1317 on 7 and 4135 DF, p-value: < 2.2e-16</pre>

root-mean-square error, and visualise modelled vs. observed values to evaluate both predictions.

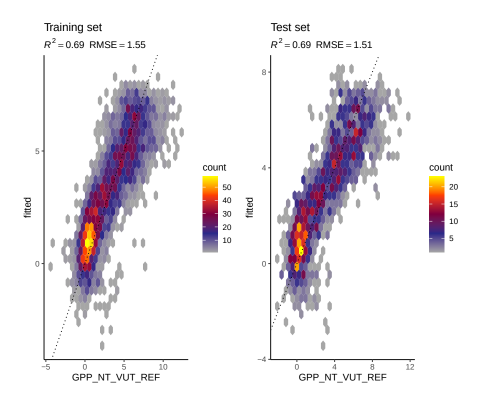
Do you expect the linear regression model trained on ddf\_train to predict substantially better on ddf\_train than on ddf\_test? Why (not)?

#### Hints:

- To calculate predictions, use the generic function predict() with the argument newdata = ....
- The  $R^2$  can be extracted from the model object as summary(model\_object)\$r.squared, or is (as the RMSE) given in the metrics data frame returned by metrics() from the yardstick library.
- For visualisation the model performance, consider a scatterplot, or (better) a plot that reveals the density of overlapping points. (We're plotting information from over 4000 data points here!)

```
library(patchwork)
library(yardstick)
## made into a function to reuse code below
eval_model <- function(mod, df_train, df_test){</pre>
  ## add predictions to the data frames
 df_train <- df_train %>%
   drop na() %>%
   mutate(fitted = predict(mod, newdata = .))
  df_test <- df_test %>%
    drop na() %>%
   mutate(fitted = predict(mod, newdata = .))
  ## get metrics tables
 metrics_train <- df_train %>%
    yardstick::metrics(GPP_NT_VUT_REF, fitted)
 metrics_test <- df_test %>%
   yardstick::metrics(GPP_NT_VUT_REF, fitted)
  ## extract values from metrics tables
 rmse_train <- metrics_train %>%
   filter(.metric == "rmse") %>%
    pull(.estimate)
 rsq_train <- metrics_train %>%
   filter(.metric == "rsq") %>%
   pull(.estimate)
```

```
rmse_test <- metrics_test %>%
   filter(.metric == "rmse") %>%
   pull(.estimate)
  rsq_test <- metrics_test %>%
   filter(.metric == "rsq") %>%
   pull(.estimate)
  ## visualise with a hexagon binning and a customised color scale,
  ## adding information of metrics as sub-titles
  gg1 <- df_train %>%
   ggplot(aes(GPP_NT_VUT_REF, fitted)) +
    geom_hex() +
   scale_fill_gradientn(
      colours = colorRampPalette( c("gray65", "navy", "red", "yellow"))(5)) +
    geom_abline(slope = 1, intercept = 0, linetype = "dotted") +
    labs(subtitle = bquote( italic(R)^2 == .(format(rsq_train, digits = 2)) ~~
                            RMSE == .(format(rmse_train, digits = 3))),
         title = "Training set") +
    theme_classic()
  gg2 <- df_test %>%
   ggplot(aes(GPP_NT_VUT_REF, fitted)) +
   geom_hex() +
   scale_fill_gradientn(
      colours = colorRampPalette( c("gray65", "navy", "red", "yellow"))(5)) +
    geom_abline(slope = 1, intercept = 0, linetype = "dotted") +
    labs(subtitle = bquote( italic(R)^2 == .(format(rsq_test, digits = 2)) ~~
                            RMSE == .(format(rmse_test, digits = 3))),
         title = "Test set") +
    theme_classic()
  return(gg1 + gg2)
eval_model(mod = linmod_baser, df_train = ddf_train, df_test = ddf_test)
```



Here, the function eval\_model() returned an object that is made up of two plots (return(gg1 + gg2) in the function definition). This combination of plots by + is enabled by the patchwork library. The individual plot objects (gg1 and gg2) are returned by the ggplot() functions. The visualisation here is density plot of hexagonal bins. It shows the number of points inside each bin, encoded by the color (see legend "count"). We want the highest density of points along the 1:1 line (the dotted line). Predictions match observations perfectly for points lying on the 1:1 line. Alternatively, we could also use a scatterplot to visualise the model evaluation. However, a large number of points would overlie each other. As typical machine learning applications make use of large number of data, such evaluation plots would typically face the problem of overlying points and density plots are a solution.

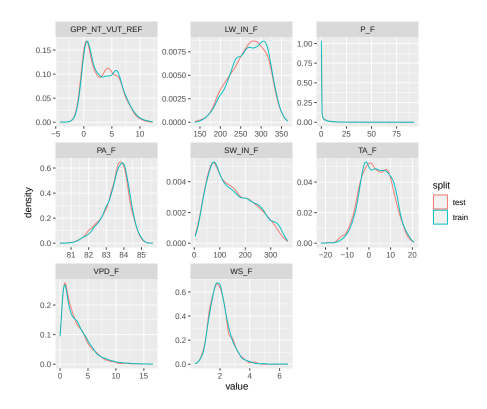
Metrics are given in the subtitle of the plots. Note that the  $\mathbb{R}^2$  and the RMSE measure different aspects of model-data agreement. Here, the measure the correlation (fraction of variation explained), and the average error. We should generally consider multiple metrics measuring multiple aspects of the prediction-observation fit to evaluate models.

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# 8.4 KNN

## 8.4.1 Check data

```
ddf_train %>%
  mutate(split = "train") %>%
  bind_rows(ddf_test %>%
    mutate(split = "test")) %>%
  pivot_longer(cols = 2:9, names_to = "variable", values_to = "value") %>%
  ggplot(aes(x = value, y = ..density.., color = split)) +
  geom_density() +
  facet_wrap(~variable, scales = "free")
```



## 8.4.2 Training

Fit two KNN models on ddf\_train (excluding "PA\_F"), one with k=2 and one with k=30, both without resampling. Use the RMSE as the loss func-

tion. Center and scale data as part of the pre-processing and model formulation specification using the function recipe().

```
library(recipes)
## model formulation and preprocessing specification
myrecipe <- recipe(</pre>
  GPP_NT_VUT_REF ~ TA_F + SW_IN_F + LW_IN_F + VPD_F + P_F + WS_F,
  data = ddf_train %>% drop_na()) %>%
  step_center(all_numeric(), -all_outcomes()) %>%
  step_scale(all_numeric(), -all_outcomes())
## fit model with k=2
mod_knn_k2 <- train(</pre>
  myrecipe,
  data = ddf_train %>%
    drop_na(),
  method = "knn",
  trControl = trainControl("none"),
  tuneGrid = data.frame(k = c(2)),
  metric = "RMSE"
  )
## fit model with k=30
mod_knn_k30 <- train(</pre>
  myrecipe,
  data = ddf_train %>%
    drop_na(),
  method = "knn",
  trControl = trainControl("none"),
  tuneGrid = data.frame(k = c(30)),
  metric = "RMSE"
  )
```

### 8.4.3 Prediction

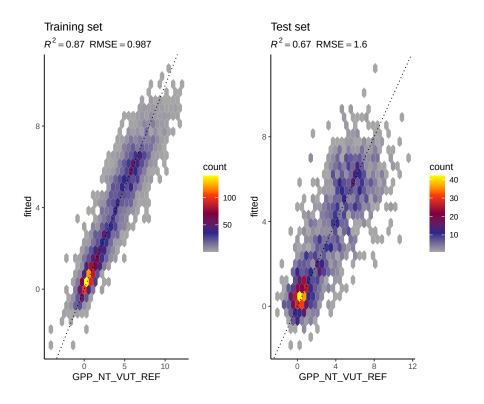
With the two models fitted above, predict "GPP\_NT\_VUT\_REF" for both and training and the testing sets, and evaluate them as above (metrics and visualisation).

Which model do you expect to perform better better on the training set and which to perform better on the testing set? Why?

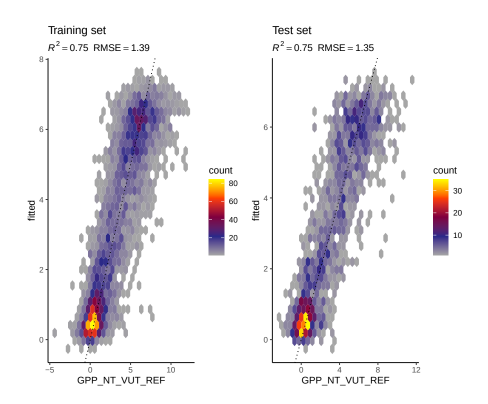
```
## with k = 2
eval_model(
```

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```
mod_knn_k2,
df_train = ddf_train,
df_test = ddf_test
)
```



```
## with k = 30
eval_model(
  mod_knn_k30,
  df_train = ddf_train,
  df_test = ddf_test
)
```



## 8.4.4 Sample hyperparameters

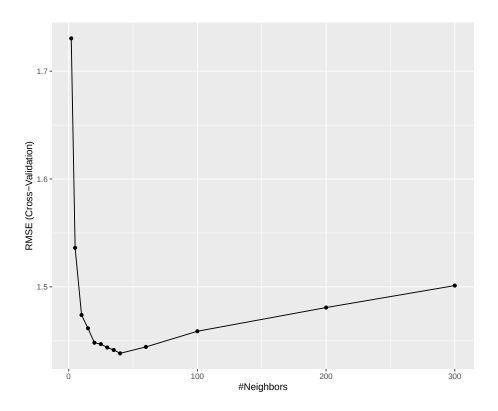
Train a KNN model with hyperparameter (k) tuned, and with five-fold cross validation, using the training set. As the loss function, use RMSE. Sample the following values for k: 2, 5, 10, 15, 18, 20, 22, 24, 26, 30, 35, 40, 60, 100. Visualise the RMSE as a function of k.

Hint:

• The visualisation of cross-validation results can be visualised with the plot(model\_object) of ggplot(model\_object).

```
mod_knn <- train(
  myrecipe,
  data = ddf_train %>%
    drop_na(),
  method = "knn",
  trControl = trainControl(method = "cv", number = 5),
  tuneGrid = data.frame(k = c(2, 5, 10, 15, 20, 25, 30, 35, 40, 60, 100, 200, 300)),
  metric = "RMSE"
```

```
ggplot(mod_knn)
```



# 8.5 Random forest

## 8.5.1 Training

Fit a random forest model with ddf\_train and all predictors excluding "PA\_F" and five-fold cross validation. Use RMSE as the loss function.

#### Hints:

- $\bullet\,$  Use the package ranger which implements the random forest algorithm.
- See here for information about hyperparameters available for tuning with caret.
- Set the argument savePredictions = "final" of function trainControl().

```
library(ranger)
## no pre-processing necessary
myrecipe <- recipe(</pre>
  GPP_NT_VUT_REF ~ TA_F + SW_IN_F + LW_IN_F + VPD_F + P_F + WS_F,
  data = ddf_train %>%
    drop_na())
rf <- train(
  myrecipe,
  data = ddf_train %>%
   drop na(),
 method = "ranger",
  trControl = trainControl(method = "cv", number = 5, savePredictions = "final"),
  tuneGrid = expand.grid( .mtry = floor(6 / 3),
                          .min.node.size = 5,
                          .splitrule = "variance"),
  metric = "RMSE",
 replace = FALSE,
  sample.fraction = 0.5,
 num.trees = 2000,
                              # high number ok since no hperparam tuning
  seed = 1982
                              # for reproducibility
)
rf
```

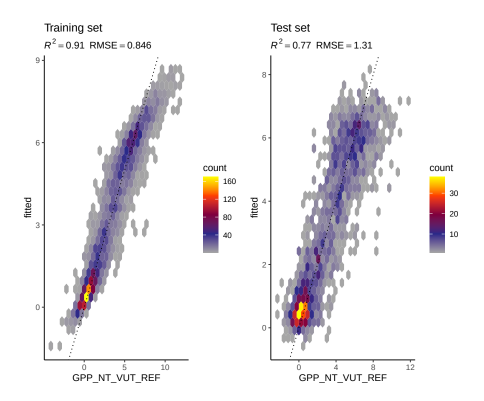
```
## Random Forest
##
## 4143 samples
##
      8 predictor
##
## Recipe steps:
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 3315, 3315, 3315, 3315, 3312
## Resampling results:
##
##
    RMSE
               Rsquared
                          MAE
##
    1.396752 0.7485248 1.055638
##
## Tuning parameter 'mtry' was held constant at a value of 2
## Tuning
## parameter 'splitrule' was held constant at a value of variance
##
## Tuning parameter 'min.node.size' was held constant at a value of 5
```

#### 8.5.2 Prediction

Evaluate the trained model on the training and on the test set, giving metrics and a visualisation as above.

How are differences in performance to be interpreted? Compare the performances of linear regression, KNN, and random forest, considering the evaluation on the test set.

```
eval_model(rf, df_train = ddf_train, df_test = ddf_test)
```



Show the model performance (metrics and visualisation) on the validation sets all cross validation folds combined.

Do you expect it to be more similar to the model performance on the training set or the testing set in the evaluation above? Why?

```
metrics_train <- rf$pred %>%
  yardstick::metrics(obs, pred)
```

```
rmse_train <- metrics_train %>%
  filter(.metric == "rmse") %>%
  pull(.estimate)
rsq_train <- metrics_train %>%
  filter(.metric == "rsq") %>%
  pull(.estimate)
rf$pred %>%
  ggplot(aes(obs, pred)) +
  geom_hex() +
  scale_fill_gradientn(
    colours = colorRampPalette( c("gray65", "navy", "red", "yellow"))(5)) +
  geom_abline(slope = 1, intercept = 0, linetype = "dotted") +
  labs(subtitle = bquote( italic(R)^2 == .(format(rsq_train, digits = 2)) ~~
                          RMSE == .(format(rmse_train, digits = 3))),
       title = "Validation folds in training set") +
  theme_classic()
```

