

Development of an R Toolbox for Near-Infrared Spectroscopy Data Processing and Analysis of Plant Metabolic Phenotypes

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Abstract

The abstract of this thesis is.....

Contents

Abbreviations

Abbreviation	Full Form
NIRS	Near Infrared Spectroscopy
LC-MS	Liquid Chromatography Mass Spectroscopy
CNN	Convolutional Neural Network
DL	Deep Learning
PLSR	Partial Least Squares Regression
PCA	Principal Component Analysis
ML	Machine Learning
NIRS	Near-Infrared Spectroscopy
SLA	Specific Leaf Area
RF	Random Forest
SLA	Specific Leaf Area
LDMC	Lead Dry Matter Content
R ²	Coefficient of Determination
ADAM	Adaptive Moment Estimation
SGD	Stochastic-Gradient Descent
AdaGrad	Adaptive Gradient Algorithm
RMSProp	Root Mean Square Propagation
MSC	Multiplicative Scatter Correction
SNV	Standard Normal Variate
GMO	Genetically Modified Organisms
HCA	Hierarchical Cluster Analysis
PLSDA	Partial Least Squares Discriminant Analysis
ANA	Artificial Neural Network
LDA	Linear Discriminant Analysis
RNN	Recurrent Neural Network
CRAN	Comprehensive R Archive Network
LC	Liquid Chromatography
RT	Retention Time

Chapter 1

Introduction

The understanding of interplay between plant physiology and its hidden biochemical process is crucial for the improvement of basic plant science and addressing global challenges such as food security, crop resilience and combating climate change [1]. In recent years, advanced High-throughput analytical techniques such as Near-Infrared Spectroscopy (NIRS) and Liquid Chromatography-Mass Spectrometry (LC-MS) has instigated a paradigm shift in plant biology [2,3]. These High-throughput techniques are mostly used in areas like genomics, imaging and spectroscopy and are known for their ability to collect and analyse the data faster than traditional techniques[3]. High-throughput techniques are widely used since they enable the efficient collection of vast amount of data at various scales, from molecular to field level over significant time periods[4]. The big data generated by these high throughput procedures present both opportunities and challenges at the same time. It requires efficient processing to extract the maximum useful results and this is where Machine Learning (ML) or Deep Learning (DL) becomes indispensable [4,5]. ML as part of Artificial Intelligence (AI) refers to the ability of computers to find patterns and learn from existing data, which can be employed in processing high-dimensional data [6,4]. The ML algorithms are powerful enough to analyse complex, high dimensional datasets, enabling accurate predictions of plant traits or other features based on the input data. Additionally, integrating these big data with ML could help the researchers to optimize data processing pipelines, enhance predictive accuracy and thereby enter into a new era of data-driven decision-making [4,5]. This project employs linear model, non-linear model and neural networks to predict various plant features and compare their predictive accuracy, error rate and training time.

A significant shift in the realm of the biomedical community has brought new guidelines to ensure readability, modularity, transparency and extensibility of computational toolboxes. A toolbox, which stores multiple functions, parameters and results in a central location should be maintainable and uncomplicated for the developers and members of the open-source community [7]. R is a powerful and widely used programming language in the analysis and processing of high throughput data. Additionally, R contains a multitude of statistical and high quality visualization packages such as ggplot2 which are capable of processing and integrating big data to different ML methods [8]. Bioconductor is an open source software for bioinformatics, which contains more than 3691 packages (according to the last update in 2024) for statistical computing. This offers an object oriented framework for the high dimensional data, cutting edge visualization capabilities and interoperability [9]. Existing tools in Near-Infrared Spectroscopy (NIRS) data processing lack functionalities that could simplify and standardize data workflows when integrated with the SummarizedExperiment framework from the Bioconductor package. To address these gaps, the R toolbox, “nearspectRa” was developed for processing NIRS data. This package has a modular structure which creates a SummarizedExperiment object from

NIRS data.

Metabolomics, the study of small molecular compounds in biological systems, is a rapidly advancing field of science with applications in biotechnology, medicine, synthetic biology and environmental science [6]. Metabolomics has emerged as a transformative tool in plant biology, enabling cost-efficient and high throughput molecular characterization. The integration of metabolomics with different omics approaches has proven invaluable for functional genes identification and developing trait specific markers [10]. Metabolomics, which is built on the advancement of phenomics and genomics, provides high throughput and precise profiling of metabolites, revealing the physiological state of cells [6,10]. Metabolites play a crucial role in plant metabolism, influencing its biomass and architecture therefore study of these small molecules will aid in uncovering plant regulatory mechanisms and pathway interactions [10]. The coupling of liquid or gas chromatography with mass spectrometry or nuclear magnetic resonance spectroscopy (NMR) facilitates measurement of thousands of metabolites, thereby providing a comprehensive view of biochemical and biological mechanisms [11]. Therefore, Mass spectrometry (MS) remains the most widely used analytical approach among others due to its versatility and sensitivity [6]. Mass spectrometry based metabolomics generate data of high sensitivity and throughput requiring advanced computational methods. Machine learning not only offers a powerful solution to analyse such data, but also helps in resolving the challenges like noise, batch effects and missing values [12]. Integrating ML with Liquid Chromatography-Mass Spectroscopy (LC-MS) data helps us to analyse this complex heterogeneous data rapidly, enabling deeper insights.

Near-Infrared Spectroscopy (NIRS) is an advanced high throughput and non-destructive analytical technique that uses light in the near-infrared region (700-2500 nm) to assess the chemical composition of samples [13]. The light is either absorbed or reflected by the sample at different wavelengths and thereby creating a spectrum [13]. The NIRS is widely used in plant research due to its ability in predicting sample structure and traits by analysing the spectral patterns. NIRS can also be used in the quantitative analysis of key plant features such as protein and carbohydrate content, secondary metabolites and physiological traits such as Specific Leaf Area (SLA) by developing calibration models between spectra and reflectance trait data [14,13]. NIRS is not only used in plant biology but also in various fields such as food science, agriculture and pharmaceuticals. When compared to other analytical techniques, NIRS is rapid, requires minimal sample preparation and less expensive, which makes it more attractive and interesting to the scientific communities [13]. However, on the flip side it requires complex statistical methods to extract different complex features due to the highly-correlated nature of NIRS data [13]. To tackle this problem, the conventional methods such as Partial Least Square Regression (PLSR) and Principal Component Analysis (PCA) imply dimension reduction which result in loss of information and often struggles to extract important features from the spectral data [13]. To address the challenge of data complexity and generalizability, different ML methods can be used to predict the traits from the NIRS data [13,15]. In this project different ML and Deep Learning (DL) has been employed to predict different plant leaf traits with use of NIRS data.

In recent years, studies using NIRS data coupled with PLSR have been used as an alternative for traditional methods such as high-performance liquid chromatography (HPLC) and mass spectrometry which are both labor-intensive and expensive to predict different plant traits. A notable example is the prediction of glucobrassicin (GBS) concentrations from NIRS data. This has shown that GBS concentrations could be reliably predicted from NIRS data [16]. Another prominent example is the tree and mycorrhizal fungal diversity experiment and trait variation

in temperate forests conducted by Pablo Castro Sanchez-Bermejo, where he combined Deep Learning (DL) approaches with leaf-level spectral data to predict 5 different leaf traits [15]. Another good example is a project involving the development of white-box workflow for regression tasks [17]. The project marks the potential of Regression (Sensitive) Neural Gas (RSNG) for generating interpretable results while maintaining high accuracy [17]. From the above studies, it is evident that NIRS data has a wide range of applications in plant research. This can also be expanded further to predict complex metabolites which are usually assessed via techniques like LC-MS. Moreover, integrating NIRS with advanced ML could further enhance the prediction accuracy and unlock new possibilities in plant science.

The past decade has witnessed the increasing popularity of Artificial Intelligence (AI) in different fields. However, this idea of AI has been under development since 1956, starting from the concept of “programming computers to think and reason” [18]. In other words, AI can be described as “automating intellectual tasks normally done by humans” [18]. Machine learning (ML) and Deep learning (DL) are the methods that fall under the realm of AI [18,19]. Nowadays, there are different ML algorithms in use, in which the most popular ones include Partial Least Square Regression (PLSR), Random Forest (RF) and Convolutional Neural Network (CNN) [20,18]. PLSR is a linear and one of the most simple ML approach. It uses the advantages of PCA and linear regression to solve the regression problem in the high dimensional data [18]. On the other hand, Random forest is a non-linear approach in ML that is primarily used for classification. It can also be used for regression tasks and can be represented as a decision tree with a series of nodes starting from a root node. The terminal node will predict the class of data in classification tasks [18]. For regression tasks, the random forest works differently compared to classification but the core principles remains the same. The terminal node of RF in regression takes the average of predictions from the individual trees [44]. The Convolutional neural networks are a specialized type of neural network which are mainly used in the field of image processing [21]. A recent study of mycorrhizal fungal diversity experiment and trait variation in temperate forests conducted by Pablo Castro Sanchez-Bermejo, demonstrated the application of CNN in predicting the leaf trait values from NIRS data, achieving superior results [15]. This outcome strongly suggests the potential of CNN not only for classification tasks such as image processing but also for regression tasks. In another project, CNN was used to predict gene expression status on the basis of sequence of gene transcription start regions. The CNN model had achieved roughly 80% accuracy [22]. These studies highlight the growing versatility of CNN models.

Omics is a term associated with the field of large scale biological data, including genomics, epigenomics, proteomics, transcriptomics and metabolomics [23]. Combination of data from these techniques along with advanced microscopy techniques helps in the study of biomolecules in cellular and subcellular levels [23]. However, the high throughput data from these omics instruments poses challenges in processing and analysing it without the loss of information [23,10]. The complexity and scale of this data make ML essential for effective integration and analysis, raising the critical question: which ML model is best suited to handle this data? How much programming expertise is necessary to implement these models? And, which models are most suitable for regression tasks?. Each ML model handles the data differently. For instance, PLSR uses latent variables to capture the covariance between predictor and response variables. Moreover, PLSR uses the combination of Principal Component Analysis (PCA) and linear regression [20]. In the case of RF, it follows the concept of “a forest made of many trees” which uses the combination of predictions from many trees [18]. Among these ML techniques, CNN is gaining attention on its ability in handling high throughput data and predicting with remarkable accuracy [15,22]. These ML models also require different levels of programming

proficiency and computational resources, depending on the scale of data.

In the light of the findings, it is clear that ML can significantly improve the analysis and processing of high throughput data from analytical techniques such as LC-MS and NIRS [15,22,17]. Among these, NIRS stands out as a non-destructive, cost-effective and rapid method, offering valuable insights into the chemical composition of the biological samples [12,13,14]. These qualities make NIRS a promising technique to optimize and integrate with ML and DL models for predictive accuracy.

Given the popularity of R programming within the ecological and bioinformatic community, it was chosen as the foundation for this project [7,9]. Recognizing the need for specialized tools to process the NIRS data, an R package, `nearspectRa`, was developed to handle data from two widely used NIRS instruments namely “ASD Fieldspec 4” and Spectra Vista Corporation (SVC) HR-1024i. Leveraging supporting packages like “R-FieldSpectra”, the high dimensional data was structured into a “SummarizedExperiment” object, aligning with Bioconductor standards for interoperability and integration.

Apart from developing a Good Scientific Practice (GSP) compliant package, this project involved two key analyses: first, predicting plant leaf traits from NIRS data using three popular ML methods, PLSR, RF and CNN and second predicting LC-MS features from NIRS data using the same models. To evaluate these approaches, performance was compared using metrics such as the coefficient of determination (R^2), Root Mean Squared Error (RMSE) and training time of each model. Additionally, extrapolation studies were conducted on PLSR and RF to assess the robustness and performance of those beyond training data. This project not only exemplifies good scientific practice in developing an R toolbox but also provides a comprehensive comparison of linear, non-linear and neural network based NL approaches in predicting plant traits and LC-MS features from NIRS data. By achieving this, the project makes a significant milestone, paving the way for a new era of cost-effective, rapid biochemical analysis in metabolomics.

Chapter 2

Background

2.1 Near Infrared Spectroscopy (NIRS)

Near infrared spectroscopy (NIRS) is a non-invasive measurement technique that uses light in the near infrared region to analyse a sample [35,13]. Infrared (IR) is a form of electromagnetic radiation that interacts with samples through absorption or reflection. The NIRS is widely used in plant research due to its ability in predicting sample structure and traits by analysing the spectral patterns. The analysis relies on how the sample absorbs or reflects light at various wavelengths. Infrared radiation is classified into three categories based on wavelength: (1) Near Infrared (NIR), ranging from 780 to 2500 nanometers (nm), (2) Mid Infrared (MIR), ranging from 2500 to 25,000 nm, and (3) Far Infrared (FIR), ranging from 25,000 to 1,000,000 nm [35]. When a substance such as plant leaf is exposed to NIR light, the molecular bonds in the infrared range will interact with the light thereby causing absorption or reflection of light by the sample. The light transmitted or reflected is then measured to generate the NIR spectrum. This spectrum provides a detailed representation of the molecular composition of the substance and the peaks in the spectrum corresponding to different vibrational modes of different chemical bonds. This occurs due to the change in the vibrational or rotational state of molecules or transition between their energy levels [35,13]. The group which is the most dominant in absorbing the NIR are hydrogen-containing groups such as C-H, O-H and N-H while other groups like C=C and C=O absorbs light in weaker intensities. These groups are key components in organic substances and their absorption wavelengths and intensities differ depending on the chemical composition of the substance [35].

This figure (Figure 2.1) illustrates the relationship between reflectance and wavelength for near-infrared (NIR) spectra where the variations in reflectance provide valuable information about the chemical and physical properties of the sample. Compared to other analytical methods, NIRS is faster, requires little sample preparation, and is more cost-effective, making it a highly appealing choice for researchers and scientists [13]. NIR spectrometers typically consist of a light source, a beam splitter, optical detectors and optionally a processing system or a monitor. The components vary depending on the purpose of the instrument to ensure accuracy and consistency. These systems can operate in different modes such as transmission, reflection, diffuse reflectance or transfectance depending on the type of analysis [36]. The collected spectra will be later subjected to chemometric analysis to develop a calibration model using key NIR bands. However, NIRS require reference data from traditional chemical analysis for accurate quantitative analysis [36]. The spectra is usually subjected to preprocessing to remove irrelevant information to enhance the analysis accuracy. The most common preprocessing techniques include, baseline correction, scatter correction, noise removal and wavelength selection. The scatter correction method and spectral derivatives are the most used preprocessing approaches



Figure 2.1: Example near-infrared spectra illustrating reflectance across wavelengths. Each line represents a unique sample, showing how NIRS captures molecular absorption/reflectance patterns.

in NIRS. The scatter correction method includes, Multiplicative scatter correction (MSC) and Standard normal variate (SNV). Additionally a wide range of normalization methods such as mean centering, auto scaling, vector normalization and area normalization are also used in pre-processing of NIRS data. The preprocessing methods are aimed to highlight important spectral features but must be carefully selected to avoid losing critical information [36].

NIRS consist of overlapping weak bands that require multivariate calibration for quantitative analysis. These methods analyze the data in a way that can identify patterns or make predictions [36]. Techniques like Principal component analysis (PCA), Partial least squares discriminant analysis (PLSDA), Hierarchical cluster analysis (HCA) and Linear discriminant analysis (LDA) are used for classification, while others like Artificial neural networks (ANNs) handle more complex relationships. Deep learning (DL) is a subset of ML that has a wide range of applications including image and audio recognition [36]. ML can also be used in the handling of “Big Data” emerging from the growing spectral libraries. DL models, unlike the traditional neural networks, consist of multiple layers to extract different features from the data resulting in better predictions [36]. In recent years techniques like Convolutional neural network (CNN), recurrent neural network (RNN) and DeepSpectra model have gained popularity. Combining DL with spectroscopy shows a great promise in applications like food quality assessment and genetically modified organisms (GMO) detection [36].

The NIRS has diverse biological applications, particularly in agricultural and soil sciences. In recent years, it has been employed to analyse the crops, food properties and plant traits, including water content, pH, oil, protein, and fatty acids. Another example would be the use of NIRS to predict anthocyanin levels in grapes and black rice seeds. Researchers have also used NIRS to identify pest attacks and pesticide residues in crops making it invaluable for quality assessment in agriculture [36]. Even Though, much of the NIRS applications focused on agriculture and food quality, its applications in food safety are also growing. For instance, NIRS can effectively evaluate the lamb meat tenderness, pH, fat and water content. The portable NIRS devices allow real time industrial monitoring making it suitable for large scale monitoring of meat, fruits, vegetables and beverages for quality and safety [36]. Genetically modified organisms (GMO) pose a risk of gene transfer, which could threaten environmental biodiversity. NIRS has shown potential for rapid and cost effective GMO detection in labs and fields, making

it a favored tool among researchers for monitoring GMOs [36].

NIRS Instruments The ASD FieldSpec4 and SpectraVista Corporation (SVC) HR-1024i are two widely used NIRS instruments that are highly relevant to this project.

ASD FieldSpec4

ASD FieldSpec4 is a portable NIRS instrument produced by Malvern panalytical and comes in different variants. It has a wide range of applications including, plant physiology, remote sensing and geology, atmospheric remote sensing research and biomass analysis among others. It provides data across the wavelength of 350 to 2500 nm. There are multiple optional accessories, contact probe (contact measurements like solid raw materials, grains, other granular materials), reflectance panels, pistol grips and leaf clip version 2 are few out of many [40].

SVC HR-1024i

This is a high resolution portable spectroradiometer which provides data across a full spectral region from 350 to 2500 nm. Compared to ASD FieldSpec4, SVC HR-1024i comes with a built-in touch screen to set parameters and review data in real time, which makes it more advanced. It is also light weight and has an internal GPS and a camera which adds critical information to the spectral signature. The output from this instrument is human readable and easy to interpret without the use of any package [41].

2.2 Liquid Chromatography Mass Spectrometry

Liquid Chromatography-Mass Spectrometry is the combination of two powerful analytical techniques for the separation, identification and quantification of biological compounds from complex mixtures. This technique has a wide range of applications in Metabolomics, Pharmaceuticals, environmental science and food chemistry [43]. Liquid Chromatography (LC) is a technique used to separate components in a mixture based on the rates at which they move through a stationary phase under the influence of mobile phase gradient. The varying affinities of the components to the stationary and mobile phases leads to their separation, with some attached to the mobile phase and eluting faster, while others attach to the stationary phase and thereby elute more slowly, leading to different retention times (RT) [43]. Mass spectrometry (MS) is one of the several detectors which can be coupled with LC to identify compounds. MS detects the mass to charge ratio (m/z) and the abundance of ions generated during ionization of the sample extracts [43]. Ions are charged molecules and are easier to analyze than neutral molecules therefore ionization is a crucial step in MS [43]. The MS is made of an ionization source, a mass analyzer and a detector, all these should be kept under vacuum to optimize the ion transmission. A mass spectrum is then generated with the help of a computer [43].

The diversity of the metabolic profile analyzed in Liquid Chromatography-Mass Spectrometry (LC-MS) makes it one of the essential tools in plant metabolomics. LC-MS has a great number of uses within the field of plant biology for plant metabolite identification and analysis. Unlike other techniques such as gas chromatography coupled to mass spectrometry, the steps involving derivatization are avoided in LC-MS [43]. This makes LC-MS particularly fit for the detection of varied classes, including semi-polar compounds and secondary metabolites common in plants [43]. The flexibility of LC-MS in gradient elution further allows effective separation of complex mixtures. Using different polarities of solvents during the elution process allows

for the separation of polar and nonpolar compounds. Reverse-phase LC-MS, a widely used approach, often employs a gradient starting with an aqueous solvent and progressively increasing the proportion of an organic solvent, ensuring the efficient separation of a broad spectrum of metabolites [43].

2.3 R Programming

R is an open-source and free software developed by professors Robert Gentleman from the University of Waterloo and Ross Ihaka from University of Auckland as a programming language to teach statistics [37]. This free programming language is widely used for statistical analysis, data visualization and graphical output [38]. Quoting the words of Ross Ihaka, “R is a computer language and run-time environment which can be used to carry out statistical (or other quantitative) computations” [37]. The name “R” itself comes from the shared first letters of both authors and being the successor of “S” language [37]. The strength of R is its extensive add-on packages which helps to analyse and visualize complex data. Even though R primarily has a command line interface, multiple third party graphical user interfaces such as Rstudio, Jupyter are available for better user friendly experience [37,38]. R is a versatile software suited for data manipulation, computational and graphical representation [39]. The GNU General Public License of R offers users significant freedom, including the ability to use, modify and distribute the software making it a favourite programming language for researchers across all the fields [37,39].

R is often associated with statistics creating a versatile environment that incorporates many traditional and modern statistical techniques [39]. Some of these techniques are part of base R setup while others are available through additional packages. R offers a greater flexibility and control over analytical processes by step-by-step analysis with intermediate results saved as objects. This allows users to interact with and refine results using additional R functions [39]. R is an expression-based programming language with a straightforward syntax. It is also case-sensitive, meaning variables like “A” and “a” are distinct and names must start with a letter or a period not followed by a digit [39]. The commands in R can either be expressions or assignments. Expressions are evaluated, displayed and then discarded. Assignments, on the other hand, evaluate expressions and assign their results to variables without displaying it, allowing more flexible programming [39].

Initially, R gained popularity for its syntax, closely resembling the S programming language, which made it easy for the S-PLUS users to transition to R. The versatility of R made it popular since it could run on various platforms, such as tablets and smartphones due to its open-source nature [8]. The frequent updates, including annual major release and timely bug fixes, highlight active development in R. Its advanced graphical capabilities is another excellent quality among many, supporting detailed and publication-quality visuals with help of packages such as “ggplot2” surpass many competitors [8]. These visualization packages enable high-dimensional visualizations and are more user friendly [8]. R remains true to its original philosophy, offering an interactive and programmable environment for creating tools. Its strong community contributes a wide range of packages, share knowledge and support on the forums like Stack Overflow, allowing innovation and collaboration among users worldwide [8]. The main R system can be downloaded from the Comprehensive R Archive Network (CRAN), which also hosts many additional packages that extend R’s capabilities [8]. The R system is divided into two parts

- The core R system, which can be downloaded from CRAN for different platforms like Linux, Windows and MacOS.
- Additional functionality through various packages.

R is a powerful tool for statistical computing, this along with high quality visualization capabilities provides a robust environment for machine learning (ML) and deep learning (DL) [42,8]. The availability of a wide range of supporting packages and libraries makes R a powerful tool for both traditional ML methods and modern DL techniques [42]. There are various packages available to perform ML and DL in R, some of them include caret, randomForest, keras and tensorflow. These packages contain the algorithm for different ML and DL models. This project involves the implementation of various ML and DL models using the R programming language.

2.4 Machine Learning

Machine learning (ML) as the name indicates is the field of computer science which applies mathematical models and algorithms to enable the system to learn and make predictions without being programmed explicitly [18,19]. In contrast to classical programming where someone explicitly programmes an algorithm to execute predefined tasks, ML uses a subset of (training) data to learn patterns and relationships within the data to create an algorithm which can generalize to unseen data[18]. This versatility and flexibility enables ML methods to improve performance over time, leading to advanced data driven decision making [23,18]. As a subgroup of Artificial intelligence (AI) is often simply represented as a 3 layer model which includes an inputlayer that receives the data, a hidden second layer which processes the data according to the mathematical backend of the model and finally the third output layer that outputs the prediction [21]. The hidden layer which does the linear regression or classification differs according to the ML model in use and these are often compared to a single human neuron, where dendrite represents input layer, cell body corresponds to hidden layer, and axon functions as output layer [21]. ML employs four primary learning methods namely, supervised, unsupervised, semi-supervised and reinforcement learning [18].

Supervised Learning Supervised learning is a ML approach that aims to predict a known output based on the input data. It excels in tasks where the patterns in data can augment human decision making [18]. For instance, handwriting recognition or object classification (example, distinguishing an elephant and tiger). These tasks are easily done by humans and supervised learning strives to replicate or enhance this performance [18]. Another example would be in medicine where supervised ML identifies patterns in the electrocardiogram (ECG) which is an easy job for a trained cardiologist. These classification tasks from supervised learning are achieved through training an algorithm on labeled datasets containing ECG features (heart rate, rhythm and waveform shape) and their corresponding diagnosis. By mapping these features (X) to diagnostic outcomes (Y), the algorithm learns the function $f(X)$ to accurately predict for new unseen ECG data [18,24]. Another common application of supervised learning is in regression and classification tasks [18]. Regression focuses on predicting continuous numerical values such as LC-MS values from NIRS data and test scores. In contrast, classification predicts which category does the given instance belong such as elephant or tiger as seen in the previous example [23,24].

Unsupervised Learning Unsupervised learning is considered to be more challenging compared to supervised learning since the former focuses on discovering patterns or groupings within data without predefined targets [18]. Common unsupervised learning tasks include clustering, association and anomaly detection, where the algorithm independently identifies underlying structures in the data [23,18]. For instance, clustering data points into separate groups based on the shared features (Figure 2.2). This approach has already proven successful in genomics, where identifying an eosinophilic subtype of asthma led to a novel therapy targeting interleukin-13, a cytokine secreted by eosinophils. Unlike supervised learning, there were no predicted outcomes, in fact there was a greater interest in identifying the patterns within the data [24].

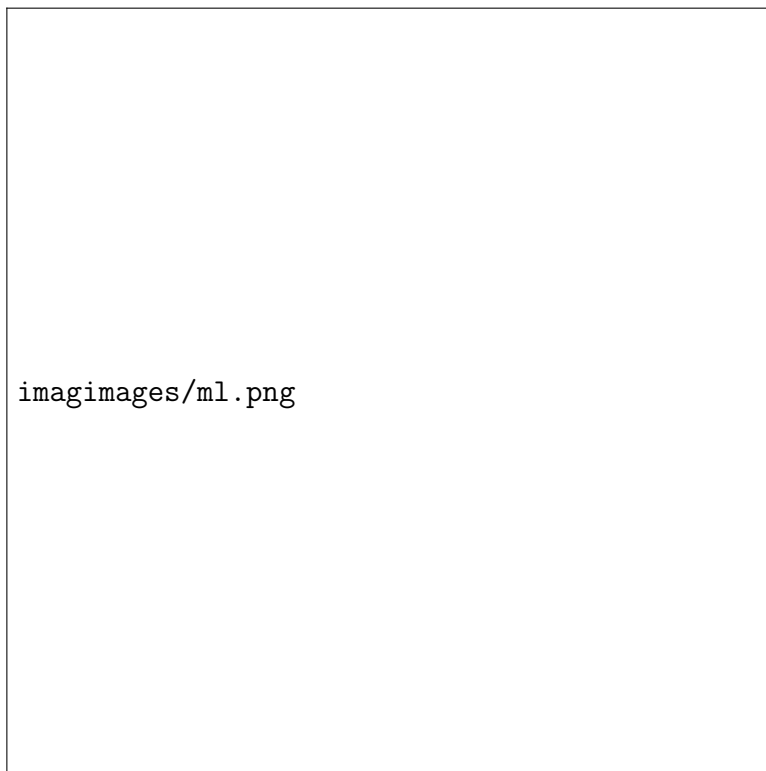


Figure 2.2: Different machine learning models illustrating the four primary learning methods: supervised, unsupervised, semi-supervised, and reinforcement learning.

Semi-supervised and Reinforcement Learning Semi-supervised learning bridges the gap between supervised and unsupervised learning by utilizing datasets that have both labelled and unlabelled data. For instance, the labelling of medical images is time-consuming and expensive. A physician might label a few medical images and this is then used to train a preliminary model which then aids in classification of the unlabelled images. This newly created labelled dataset will then be used to train a more robust final model[24]. On the other hand, reinforcement learning mimics the human learning process, where it relies on trial and error then solely on data [24].

In the era of modern molecular plant breeding, integration of ML with the large, noisy and heterogeneous data is important to uncover complex patterns and enable accurate predictions of plant features [23]. The “big data” resulting from high throughput techniques in plant sciences can be leveraged to drive discoveries, enhanced precision and accelerate advancement in plant research [23]. A plant genetic makeup (genotype) has a significant influence in its growth,

development and biochemical composition. This results in the expression of plant traits such as yield, stress tolerance and pest resistance. Understanding how genotype and environment influences on phenotypes is crucial for insights into regulatory mechanisms, and development of plants [23]. This knowledge enables the prediction of yield and other plant traits based on the genotypes under different environmental conditions, which in turn paves the way for modern molecular plant breeding [23]. Different ML approaches such as Partial Least Square Regression (PLSR), Random Forest (RF) and Convolutional Neural Networks (CNN) can be employed to make predictions by leveraging patterns in the data.

2.4.1 Partial Least Square Regression (PLSR)

In machine learning, Partial Least Square Regression (PLSR) is a statistical method which combines the benefits of Principal Component Analysis (PCA) and linear regression to predict the outcomes [26]. It is a linear regression model which is arguably the simplest machine learning algorithm that uses a straight line to solve a regression problem [18]. PLSR uses the advantage of PCA for dimensionality reduction and the regression for prediction [27]. This fitting of linear regression between two data matrices has a wide range of application in plant biology, especially in crop breeding, ecosystem monitoring and predicting plant traits from its spectral data [25]. In PLSR, the predictor variable (often denoted as X) refers to a set of independent variables or features that are used to predict response variable (y). The predictor variables are typically high dimensional and often include multiple correlated features. The response variable (y) represents the outcome or dependent variable. PLSR works by identifying the latent variables, which summarizes the covariance between predictor and response variables. This latent variable captures the most relevant information from the predictors (X) in relation to response (y) variables, allowing the model to predict y more effectively [25,27,28].

Principal Component Analysis (PCA) uses the principal components as explanatory variables to capture most of the variance in predictor variables (X), ensuring dimensionality reduction while retaining most of the data's variability. Partial Least Squares (PLS) prioritize relevance to both X (predictor) and y (response) variables, rather than maximising the variance in X alone (Figure 2.3) [27]. By combining the PCA with linear regression, PLSR constructs the latent variable that summarizes predictor variables and maximises their relevance to the response variables. This makes PLSR particularly suitable for high dimensional datasets such as spectral data from NIRS instruments.

2.4.2 Random Forest (RF)

Random Forest (RF) is a non-linear ensemble technique used in machine learning which is also depicted as a forest of decision trees [18][29]. Random forest is commonly used for classification tasks, though it can also be applied to regression such as predicting the leaf trait values from the NIRS data [18]. It is a supervised learning method consisting of decision trees and the root node serves as the initial point for dividing the dataset. Recursive partition in which the data is separated into two binary part begins with the root node [18]. RF combines multiple trees to improve accuracy, robustness and reduce overfitting [18,29]. In the classic tree based model, the dataset is divided into two groups based on the certain criteria until it encounters a predetermined stopping condition. The endpoints of a decision tree, known as leaf nodes, represents the final data division. A random forest model consisting of an ensemble of decision trees, can be utilized for both regression and classification tasks, depending on the partitioning

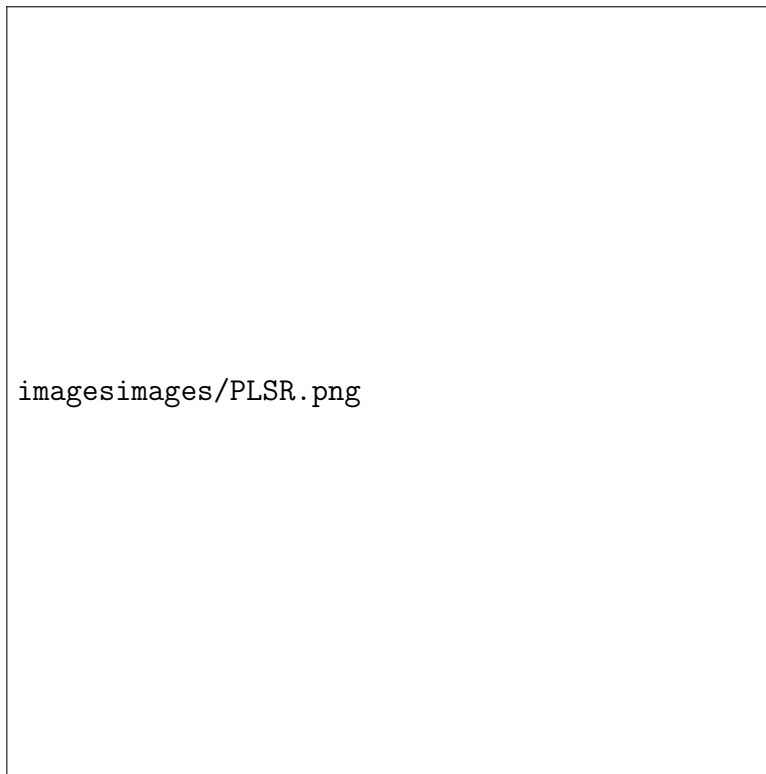


Figure 2.3: A comparison of PCA (A) and PLS (B). In the PCA plot, the x-axis represents a combination of variables (e.g., three metabolites) that captures the greatest variation in the dataset, independent of group classification. In contrast, PLS focuses on explaining the relationship with an explanatory variable, such as "Treatment" in this example

strategy and stopping criteria [30]. RF has proven its importance in many scientific domains and one of which was in environmental science, in a study employed learning algorithms such as Least Absolute Shrinkage and Selection Operator (LASSO) Regression, RF and neural network to predict ragweed pollen concentrations, with RF delivering the most accurate predictions [30,31]. A graphical representation of the RF model with decision trees can be found in the figure (Figure 2.4) .

Random Forest often outperforms linear regression models since, linear regression assumes a linear relationship between the variables. Even though this makes linear regression easier to interpret, it limits their flexibility in capturing complex patterns in the dataset. On the flip side, RF can easily adapt to non linear relationships, making them better flexible and suited for such tasks [30]. The RF algorithm estimates the error rate by out-of-bag (OOB) during training time. Each tree in the model is built using a subset of data, called a bootstrap sample and about one third of the data is left out during this process. These excluded data points are the OOB. Hence, minimizing the OOB is crucial for better model performance and robustness [30].

2.4.2.1 Mathematical Explanation of Random Forest Regression

For regression tasks, the random forest works differently compared to classification but the core principles remain the same. The terminal node of RF in regression takes the average of predictions from the individual trees [44]

- **Bootstrapping and Tree Construction**

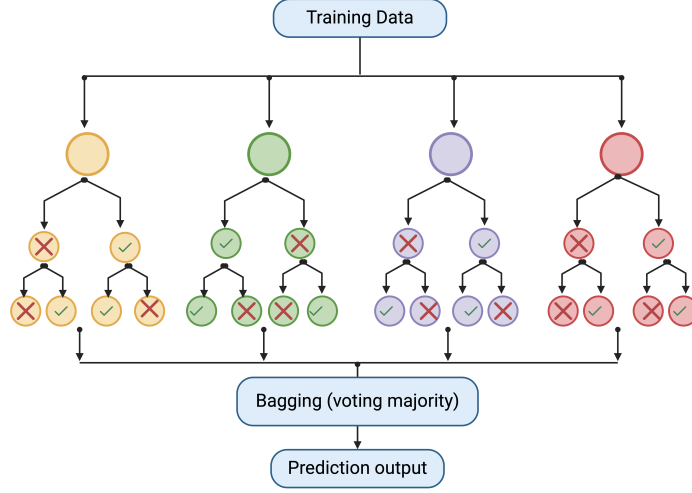


Figure 2.4: Schematic representation of a Random Forest model (classification).

Each tree in a RF model is trained using a bootstrap sample, which is a subset of the sample from the training data of the original dataset. Approximately one third of the samples (data points) are left out during bootstrap sampling for each tree and these left out samples are referred to as Out-Of-Bag (OOB) samples. Given a dataset $D = \{(x_1, y_1), (x_2, y_2), \dots, (x_n, y_n)\}$, where:

- x_i represents the feature (e.g., spectral data),
- y_i represents the response variable (e.g., SLA, LDMC, or other traits),

For each tree (t), a bootstrap sample D_t is taken out of dataset D , and the OOB for that tree is calculated as:

$$D_t^{OOB} = D \setminus D_t$$

• Model training and Prediction

Each tree in a RF model is trained with a separate bootstrap sample (D_t) and later exposed to test data for predictions to be made. For the test data X_{test} , the prediction from each tree (t) is denoted as $\hat{y}_t(x_{test})$. The final prediction from the RF is obtained by taking the average value of predictions from all the individual trees:

$$\hat{y}_{RF}(x_{test}) = \frac{1}{T} \sum_{t=1}^T \hat{y}_t(x_{test})$$

where:

- $\hat{y}_t(x_{test})$ is the prediction from the t -th tree,
- T is the total number of trees in the Random Forest.

In this project, we will be using RF models to do regression tasks on NIRS data and compare the coefficient of determination R^2 , RMSE and training time with that of linear model and neural network, we will also discuss developing the RF model and associated functionalities.

2.4.3 Convolutional Neural Network (CNN)

Convolutional Neural Network (CNN) is a type of deep learning (DL) model inspired by the way neurons in the brain process visual information [32,33]. It is primarily composed of three core components: A convolutional layer that extracts features, a pooling layer to reduce the dimensionality of data and a fully connected layer that produces the final output [32]. Out of these three main layers, the convolutional layer is considered as a fundamental component of a CNN, consisting of a series of mathematical operations, including convolution, which is a distinct form of linear regression [32]. It involves the application of kernels, which is a small array of numbers across the input (tensor) to compute elementwise products. These results are summed to create an output called feature map. Each kernel extracts a different feature of the input data and thereby different feature maps with different characteristics of the input data. The number of kernels determines the depth of the data and is also selected based on the scale of input data. A stride is known as the step size for moving the kernel across the tensor, commonly a stride of 1 is used. To capture the outermost element in the tensor, zero padding technique, which involves adding rows and columns of zero on the sides of input tensor is used to prevent downsizing in the convolutional layer [32]. After the convolutional layer the feature map is then processed through a nonlinear activation function and then to the pooling layer for downsampling [32]. In CNN the widely used pooling method is max pooling, which divides the feature map into small patches and then keeps only the highest values from each patch and ignores the rest. The output from the last pooling layer is flattened to 1 dimensional array and passed to fully connected layers, where each input is connected to every output with a learnable weight. These layers map the extracted features to the final output [32].

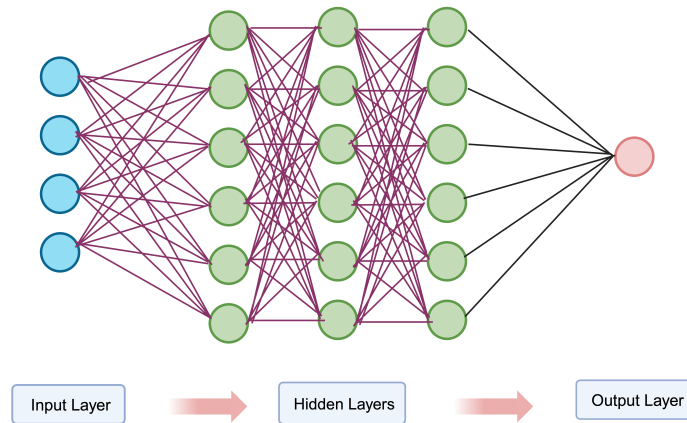


Figure 2.5: Schematic representation of Convolutional Neural Network

A loss function evaluates how well the predicted values match the actual ones. For classification cross-entropy loss is commonly used whereas mean squared error (MSE) is preferred for regression tasks. Choosing the right loss function is essential and is determined according to the given task [32]. In deep neural network training, the weights of each neuron is estimated to establish an accurate relationship between inputs and outputs with a desired level of precision [34]. It is categorized into supervised learning, for classification and regression tasks, and unsupervised learning, for clustering with input data only. Several methods are employed for training deep neural networks, including Backpropagation, Gradient Descent, Stochastic Gradient Descent (SGD), and others. Among these, SGD stands out as one of the simplest

and most widely used optimization algorithms in machine learning [34]. Adaptive Moment Estimation (ADAM) is an advanced optimization algorithm which combines the benefits of two SGD variants: Adaptive Gradient Algorithm (AdaGrad) and Root Mean Square Propagation (RMSProp) [34]. It requires minimal memory and dynamically adjusts the learning rate for each parameter, making it more computationally efficient [34]. In this project CNN is employed to predict the output values from NIRS data.

Chapter 3

Implementation

3.1 R Toolbox Development

The development of a new toolbox was prompted by the realization that existing R packages lacked functions to compartmentalize spectral data in a manner that could be seamlessly integrated into the Bioconductor ecosystem. Bioconductor offers significant advantages in terms of integration and interoperability, which are highly valuable for advanced data analysis. Existing packages commonly used in ecological studies, such as **FieldSpectra**, **plantspec**, and **spectacles**, were reviewed. However, none provided functionality to convert high-dimensional spectral data into a `SummarizedExperiment` object, a data structure widely used in Bioconductor for organizing and analyzing complex datasets.

Recognizing the potential applications of the `SummarizedExperiment` framework in plant biology, along with its ability to streamline workflows and facilitate future analyses, a decision was made to develop a toolbox to address this gap. The primary objective during development was to ensure vendor independence, making the toolbox broadly applicable. Vendors such as Malvern Panalytical offer software for processing data from instruments such as the ASD FieldSpec 4, while the SVC HR-1024i from Spectra Vista Corporation requires additional computational effort to structure the data for analysis. In addition, the ASD FieldSpec 4 generates binary output files. To process these files, functions from the `FieldSpectra` package were utilized as a foundation, with additional custom functions developed to convert the data into `SummarizedExperiment` objects. The entire workflow, including data reading, processing, and conversion, is detailed in this chapter.

3.1.1 Toolbox: `nearspectRa`

The toolbox, named **`nearspectRa`**, is specifically designed to handle Near-Infrared Spectroscopy (NIRS) data. This name was chosen to align with its functionality and field of usage, focusing on plant metabolic phenotype analysis. The toolbox is publicly available on GitHub and can be accessed at the following link: [GitHub Repository](#)

3.1.2 Functions of `nearspectRa`

Function 1: `read.summarizedexperiment_asd`

Key Features:

- Input: A path to either a single .asd file or a directory containing multiple .asd files.
- Output: A SummarizedExperiment object with:
 - Assays: Reflectance data as a matrix where rows are wavelength and columns are samples.
 - colData: Sample metadata.
 - rowData: Wavelength metadata.
- Dependencies: Utilizes `read.asd` and `extract.metadata` from the FieldSpectra package for reading and extracting data, and the `SummarizedExperiment` library from Bioconductor.

Pseudocode:

Function: `read_summarizedexperiment_asd(path):`

Input: path to a single .asd file or a directory containing multiple .asd files.

Output: A SummarizedExperiment object.

1. Check if the path is a file or directory.
2. If it is a directory, list all .asd files in the directory.
3. For each .asd file:
 - (a) Use `read.asd` to extract spectral data.
 - (b) Use `extract.metadata` to obtain sample metadata.
4. Combine spectral data and metadata into matrices.
5. Create a `SummarizedExperiment` object with:
 - Assays: Spectral data matrix (rows: wavelengths, columns: samples).
 - ColData: `DataFrame` of sample metadata.
 - RowData: `DataFrame` of wavelength information.
6. Return the `SummarizedExperiment` object.

Example Usage:

```
path <- "path/to/your/asd/files"
se <- read_summarizedexperiment_asd(path)
print(se)
```

Function 2: `read_summarizedexperiment_sig`

Key Features:

- Input: A path to a single .sig file or a directory containing multiple .sig files.
- Output: A SummarizedExperiment object with:
 - Assays: Reflectance data as a matrix where rows are wavelengths and columns are samples.



Figure 3.1: Example output of the `read_summarizedexperiment_asd` function

- colData: Sample metadata.
- rowData: Wavelength metadata.
- Dependencies: Utilizes `readLines` for file reading, `grep` for pattern matching, and the `SummarizedExperiment` library from Bioconductor.

Pseudocode:

Function: `read_summarizedexperiment_sig(path):`

Input: path to a single `.sig` file or a directory containing multiple `.sig` files.

Output: A `SummarizedExperiment` object.

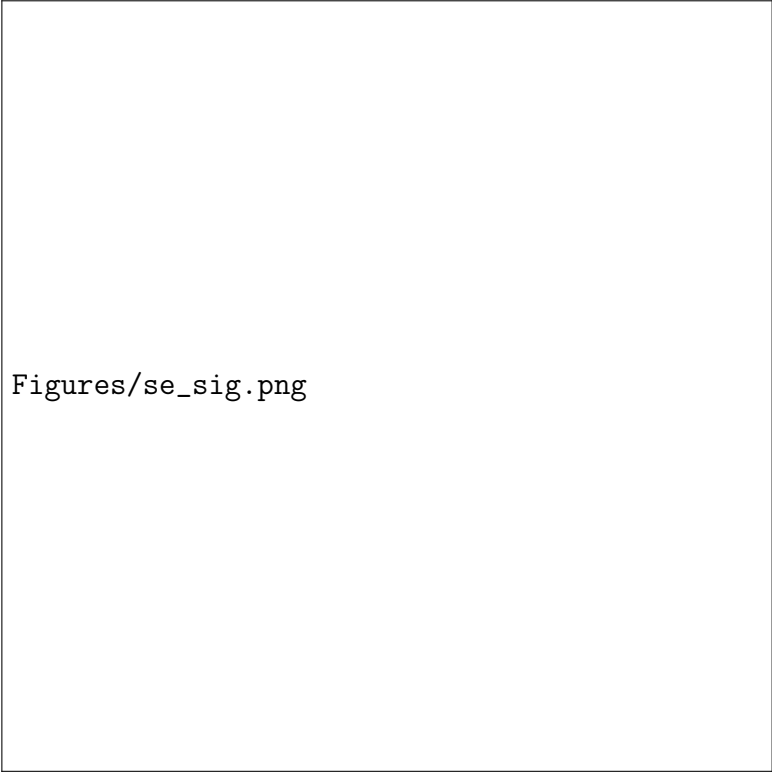
1. Check if the path is a file or directory.
2. If it is a directory, list all `.sig` files in the directory.
3. For each `.sig` file:
 - (a) Use `readLines` to read the file content.
 - (b) Extract the sample name from the line starting with `name=`, or use the file name if not present.
 - (c) Extract spectral data starting from the line that begins with `data=`.
 - (d) Organize the data by wavelengths (first column) and reflectance (fourth column).
4. Combine all data frames by wavelengths, filling missing values with `NA`.
5. Create a `SummarizedExperiment` object with:

- Assays: Reflectance data matrix (rows: wavelengths, columns: samples).
- colData: `DataFrame` containing sample names.
- rowData: `DataFrame` containing wavelengths.

6. Return the `SummarizedExperiment` object.

Example Usage:

```
path <- "/path/to/sig/files"
se <- read_summarizedexperiment_sig(path)
print(se)
```



Figures/se_sig.png

Figure 3.2: Example output of the `read_summarizedexperiment_sig` function

3.1.3 Package Development Overview

3.1.3.1 Purpose and Objectives

This package is aim to ease the NIRS data processing by reading the raw data and outputs a `SummarizEdexperiment` object. The programming language and tools used for this development process is mentioned below.

3.1.3.2 Language

R: is the core programming language used for the development. It is choosen because of its powerful data analytical capabilities and wide usage in bioinformatics. The R version 4.4.0 (eleased on April 24, 2024) has been used for this project.

3.1.3.3 Libraries and Dependencies

Base R Functions: These functions come pre-installed and do not require additional libraries or packages. For instance, `readLines`, `list.files`, `files.info`

FieldSpectra: is Used for reading the ".asd" files and to extract the metadata. The version used is FieldSpectra 0.9.7

SummarizedExperiment: is central to this package and it enable structured storage of spectral data and integrate to the bioconductor world. The version used is 1.34.0

roxygen2: is used for generating documentation directly from inline comments. The version used is 7.3.2

3.1.3.4 Version Control

One of the main challenges during software development is tracking and documenting the code during the development process. These challenges include, managing multiple script versions or integrating collaborator edits. Manual merging and making multiple copies of each version is time consuming and error-prone [46]. To overcome this challenge, Git is used for version control and collaborative development. A git version of 2.32.0 is used.

Examples of few git commands used in this project are given below:

```
# Navigate to the project directory
cd/path/to/the/project

# Initialize a Git repository
git init

# Add a remote repository
git remote add origin https://github.com/username/project.git

# Check the status of the repository
git status

# Stage all files for commit
git add .

# Commit the changes with a message
git commit -m "First commit"

# Push the changes to the remote repository
git branch -M main # Rename branch to main
git push -u origin main
```

3.1.3.5 Testing and Validation

Unit tests are implemented under *test* directory to ensure a robust function performance across different datasets. **testthat** package is used for the unit test development.

3.1.3.6 Documentation

Documentation is a fundamental aspect of software development, regardless of the programming language or platform. It serves as a vital tool for communication between developers and end users, providing the necessary information for effectively utilizing the software to its

full potential [45]. In developing `nearspectRa`, we prioritized clear and comprehensive documentation to ensure the package is user-friendly, while adhering to FAIR principles—Findable, Accessible, Interoperable, and Reusable. This documentation was created with the help of the `roxygen2` package version 7.3.2.

3.1.4 Usage and Integration

3.1.5 Challenges and Future Work

3.2 Contributions elsewhere

3.3 HPC runs

Chapter 4

Results and Discussion

Table 4.1: Comparison of 3 ML models across data

Partial Least Square Regression (PLSR)			
Trait	R^2	RMSE	Training time
SLA	0.89	0.40	13.76 sec
LDMC	0.85	0.41	11.40 sec
C:N	0.72	0.50	15.71 sec
C	0.68	0.55	13.63 sec
P	0.66	0.64	15.19 sec
PLSR Extrapolation			
SLA (H 30%)	0.14	1.06	21.07 sec
SLA (L 30%)	0.70	0.93	52.38 sec
Random Forest (RF)			
Trait	R^2	RMSE	Training time
SLA	0.87	0.42	12.85 min
LDMC	0.54	0.72	12.11 min
C:N	0.57	0.63	12.60 min
C	0.37	0.82	13.17 min
P	0.50	0.66	15.25 min
RF Extrapolation			
SLA (H 30%)	0.035	1.29	11.64 min
SLA (L 30%)	0.17	1.53	3.44 min
Convolutional Neural Network (CNN)			
Trait	R^2	RMSE	Training time
SLA	0.89	0	28.11 min
LDMC	0.88	0	30.20 min
C:N	0.74	0	4.11 min
C	0.36	0	4.83 min
P	0.20	0	31.30 sec

4.1 Data charecterestics

histogram, spectra

4.2 Baseline Machine Learning Models Pablo

PLS, RF, CNN

4.2.1 Variable importance

4.3 Variations in Baseline systems

4.3.1 modifying the Test and Training split

4.3.2 input data length

4.4 Sues

Chapter 5

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