

Comparison of statistical methods and designs for a high throughput phenotyping experiment

Alexandre BOHYN

Supervisor: Prof. P. Goos
KULeuven

Mentor: Pr. X. Draye
UCLouvain

Thesis presented in
fulfillment of the requirements
for the degree of Master of Science
in Statistics

Academic year 2018-2019

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Chapter 1

Introduction

In the recent years, food security has grown to become a worldwide concern due to the challenges imposed on farmers by climate change. In this respect, genetic improvement to increase crop resistance to abiotic stresses is seen as a solution to generate novel traits in plants and help fight those threats (Tester & Langridge 2010). In the last decades, genetic editing and sequencing techniques have greatly improved, leading to a high-throughput of genetic data and associated data analysis techniques (Schiml & Puchta 2016). However, to capitalize on those discoveries, similar advancements need to be made regarding plant phenotyping. Even though many breakthrough regarding sensors and imaging techniques are made, plant phenotyping still remains a challenge (Furbank & Tester 2011). Due to the extreme sensitivity of plants to their growing environment, a large amount of the research is dedicated to study plants' variations that are unrelated to their genetic traits. Researchers are setting up high-throughput platforms, where thousands of plants are sequentially analysed, to be able to quantify the influence of environmental conditions on plant growth (Tardieu *et al.* 2017).

In parallel, plant breeding trials often involve a large number of genotypes and large areas where spatial variation is likely to be an obstacle to reliable genetic prediction. To account for complex spatial variations, researchers often use spatial analysis methods that model the correlation between neighbouring plots (Velazco *et al.* 2017). There is a large array of different methods such as nearest neighbours analysis (Wilkinson *et al.* 1983), spatial covariance structures (Gilmour *et al.* 1997; Piepho & E. Williams 2010), polynomials models (Federer 1998) or smoothing splines (Durban *et al.* 2001). However, for these approaches to be efficient, they require a complementing experimental design. Over the years, several types of complex experimental designs have been created and tested in field trials (Cullis, Smith, *et al.* 2006; Patterson & Hunter 1983; Yates 1939). However, even though those designs allow good correction for field trends, they are rarely used in practice because of their complexity and the specific cases to which they apply. This is even more relevant for phenotyping platforms, where the experimental set-up is rarely like the one of a field.

With this background in mind, the motivation behind this thesis was to design and conduct an experiment in a phenotyping platform and then analyse the results using different spatial models. We created an optimal design, fitted for the platform, and applied in experiment with different genotypes of maize. We then harvested the plants

and analysed the weights using different spatial models, to compare their efficiency on this given platform. The fact that we over-viewed each step of the experiment from the seeds germination to the analysis of the results took a large amount of time but allowed more control over the different phases of the process. On this platform, plants are constantly moving to be characterized, which renders the spatial trend quantification complicated. Even though this feature is often available in platforms, it is poorly evaluated. Therefore, to make this thesis relevant we wanted to evaluate the impact of constant movements on the plants growth.

To summarize, this master thesis can be divided in two sections. The first one is about creating an experiment adapted to the platform's characteristics to evaluate the impact of movements on plant growth, as well as the effect of each genotype. The second part is about fitting different spatial models to see which one is better able to capture, and explain, the spatial trends in the data. We do not aim at creating new spatial models or revolutionizing experimental design, but rather to apply those concepts on a practical case, to see how well they allow the analysis and interpretation of phenotypic data. The following sections are a literature review of the topic; then a development of the material and methods used (that is the experimental procedure and the spatial models); a presentation and discussion of the results; and finally a conclusion to discuss if and how the goals, presented in this introduction, were met in the thesis.

Chapter 2

Literature review

2.1 Plant phenotyping

The terms phenotype and phenotyping are often interpreted in diverse ways between authors and between studies. In order to avoid any confusion, it is important to define these concepts clearly. Plant phenotyping is defined as the identification of effects on the phenotype (i.e., the plant appearance and performance) as a result of genotype differences (i.e., differences in the genetic code) and the environmental conditions to which a plant has been exposed (Fiorani & Schurr 2013; Houle *et al.* 2010). In this thesis, we refer to phenotyping more precisely as the set of methodologies and protocols used to measure plant growth, architecture, and composition with a certain accuracy and precision at different scales of organization (Fiorani & Schurr 2013).

Plant phenotyping is an important tool to address and understand plant environment interaction and its translation into application in crop management practices, effects of biostimulants, microbial communities, etc... (Pieruschka, Schurr, *et al.* 2019). In our current society, food security is a rising issue and genetic crop improvement is seen as a solution to deal with this issue. While genetic editing techniques and genome mapping technologies are blooming, they depend on a similar improvement in phenotyping, since they are key to analyse plant responses to environmental characterization. In recent years, high-throughput and high-resolution phenotyping tools have made impressive progress and can now help relieving the current phenotyping bottleneck (Fiorani & Schurr 2013; Furbank & Tester 2011; Tardieu *et al.* 2017). Different phenotyping platforms are emerging around the world. They range from high-precision platforms for cell and organ characterization (Vargas *et al.* 2006) to multi-environment networks of fields, exploiting remote sensing (Virlet *et al.* 2017). An experiment generates a large amount of raw data that provides a condensed set of multi-dimensional information (2D usually, but 3D scanning platforms are developping (Mooney *et al.* 2012)). A lot of tools are available for data analysis in phenotyping platforms (Lobet, Draye & Périlleux 2013). This makes the choice complicated for an external user, especially since most of these softwares are designed for a single specific purpose. Another challenge in root system architecture (RSA) characterisation is the inherent complexity of the system. Different techniques have been developed to best characterize the RSA in a cost-efficient way (Lobet & Draye 2013; Pound *et al.* 2013). However, at all scales, phenotyping facilities display spatial heterogeneity that needs to be separated from the genetic signal. For example, the spatial variability of incident light raises up to 30% between pots within

a glasshouse or a growth chamber (Cabrera-Bosquet *et al.* 2016). There are also evidences of microclimate variations in greenhouses experiments (Brien *et al.* 2013). Therefore, correcting for spatial trends and using appropriate experimental designs is crucial for a precise estimation of genetic effects. Hence, the existing design and modelling theory for field experiments needs to be adapted for the phenotyping platforms.

2.2 Experimental design in field trials

Experimental field trials in agriculture have always been affected by soil heterogeneity. As Van Es (2002) explains, soil is a continuum with variability on multiple scales. The heterogeneity is as much affected by microscopic interactions as by field-sized effects. Therefore, agricultural trials have always heavily relied on randomisation, blocking and replication to account for spatial variability and remove bias from the estimation of the treatment effects (Atkinson & Bailey 2001). For randomisation to be truly effective, stationarity of the mean and spatial independence assumptions need to be verified. Several studies have proven that it is rare that both these assumptions hold in field trials (Davidoff *et al.* 1986; Iqbal *et al.* 2005; Nielsen *et al.* 1973). Moreover, Van Es (1993) showed that even randomized designs can still be problematic for experiments with large numbers of treatments and low numbers of replications in the presence of spatial autocorrelation. A new class of design has been proposed involving the use of replicated plots for a percentage of the test lines: the “p-rep” designs (Cullis, Smith, *et al.* 2006; Velazco *et al.* 2017). Local field trends can influence groups of treatments in specific blocks. As a solution, several authors (Fagroud & Van Meirvenne 2002; Watson 2000) have suggested considering the spatial trends and autocorrelation structures when creating the design, by using prior soil information, but taking into consideration spatial variability in the design of a trial not only require previous information on the plot but is often costly and cumbersome. Furthermore, in practice, most experimenters have neither the capacity to implement advanced designs (in terms of computation power and statistical training), nor the capacity to analyse them. Finally, Van Es *et al.* (2007) showed that completely randomized (43 % in greenhouse trials) and random block designs (70 % in field trials) are still widely used. Considering this global issue, finding and using an appropriate design is complex task.

2.3 Spatial modelling for field trials

In order to increase the precision of the estimation of genetic effects, experimental designs need to be complemented with appropriate models of analysis. Mixed model analyses using the autoregressive (AR_1) functions (Cullis & Gleeson 1991) have become a standard strategy in field trials. However, Piepho, Möhring, *et al.* (2015) recently discussed several issues with this model and have therefore proposed the use of the linear variance (LV) model (E. R. Williams & Luckett 1988) instead. More specifically, Piepho & E. Williams (2010) have proposed a revised version of this model, augmenting it into two dimensions ($AR_1 \times AR_1$). The main novelty resides in the addition of spatial components to a classic rows-columns model. Recently, Rodriguez-Alvarez

et al. (2018) introduced a novel spatial model that adjusts for both global and local trends simultaneously: the SpATS model (Spatial Analysis of field Trials with Splines). The new spatial method makes use of penalized splines (P. H. C. Eilers & Marx 1996) to estimate a bivariate smooth function over the rows and columns of a plot. Using the work of Lee Hwang (2010), Lee & Durbán (2011), and Lee, Durbán & P. Eilers (2013) the spatial variability is characterized using tensor products of two-dimensional P-splines (Dierckx 1995) and decomposed in a PS-ANOVA system. By exploiting the similarities between P-splines and mixed models (Currie & Durban 2002; Durban *et al.* 2001; Wand 2003), the P-splines are expressed as a mixed model, which allows the use of classical mixed-model software but also the use of additional random and fixed effects to the model to better capture the variation along the 2-dimensional field. It has already been tested on simulated data (Rodríguez-Alvarez *et al.* 2018) and previous field trials data (Lado *et al.* 2013) and showed promising results.

As Wilkinson *et al.* (1983) and Gilmour *et al.* (1997) highlight, in field trials data modelling, three main sources of spatial variations need to be accounted for:

Stationary¹ variations: Large scale trends across the field (e.g. fertility trend, depth of soil, moisture)

Non-stationary variations: Also natural variations but localized on part of the field (e.g. patch of soil moisture)

Extraneous variations: Variations unrelated to a natural process, often due to the way the field is prepared (e.g. tillage, sowing practices, etc. . .)

A part of these variations can be attributed to systematic effects, e.g. sowing or planting, other to random effects such as fertility trends. While systematic effects can easily be modelled using factors and row-columns attributes, it is not case the case for random spatial variation. They are harder to model because there are no covariates to relate it to. Since the spatial variation has both random and systematic components, it makes sense to use the mixed model framework.

There are two main approaches to model spatial trends: one based on spatial variance-covariance structures; and the other based on smoothing techniques. The SpATS model uses a smooth bivariate surface to model both the global and local trends, while accounting for the extraneous variations by using extra random and/or fixed coefficients. Models using spatial covariance structure (such as the $AR \times AR$) model the global trends using functions of the spatial coordinates (both linear trends and smoothing splines), while the local trends are estimated with the use of spatially dependant error term (thus the reason why these models use spatial covariance structure) and the extraneous trends are managed similarly to the SpATS model. In this thesis, the data extracted from the phenotyping platform are modelled using these 2 different models.

¹Risser (2016) defines a stationary process as follows:

Let C be a spatial covariance function, it is said to be stationary if the features of C do not depend on spatial location. More formally, a process $\{Y(s) : s \in G\}$ is said to be second-order stationary (or weakly stationary) if the following two properties hold:

1. $E[Y(s)] = E[Y(s + h)] = c$ for some constant c and
2. $C(s, s + h) = C(0, h)$ for some spatial lag $h \in \mathcal{R}^d$.

2.4 Thesis objectives

This master thesis falls within the scope of the second research activity of the European project EPPN²⁰²⁰². It is a research infrastructure project funded by Horizon 2020, that will provide access to 31 key plant phenotyping installations. It defines three research activities: (1) novel technologies and methods for environmental and plant measurements, (2) innovative design and analysis of phenotyping experiments across multiple platforms and (3) a European plant phenotyping information system. The project revolves around data acquisition, data analysis and data networking, so that every platform uses common, standardized practices and analysis protocols, that have been tested for robustness and quality.

The main goal was to assess the utility of statistical designs and mixed models to identify and correct for spatial trends (heterogeneity) in an aeroponic root installation at UCLouvain (Louvain-la-Neuve). The idea is to set up an experiment in this installation using different genotypes (plant varieties) and a custom experimental design to account for possible complex environmental variations. It was created using JMP®(Version 14.3, SAS Institute Inc., Cary, NC, 1989-2019.), taking into account the specificities of the platform and the number of genotypes used. This approach allows the design to fit the experiment properly and avoids having to use a pre-made design, not optimal for the experiment. After data collection and image analysis, two different models will be used to model the spatial variability and to assess the quality of spatial prediction. The first one is a two-dimensional version of the linear variance model, revised by Piepho & E. Williams (2010). The second one is the SpATS model, recently created by Rodriguez-Alvarez *et al.* (2018). The two models will be compared in term of their ability to estimate genotypic effects and to quantify spatial variability. These comparisons will be made using classical indicators (RMSE, ...) and other indicators, specific to spatial models for field trials (Oakey *et al.* 2006).

The experiment took place in February 2019 in the UCLouvain greenhouses. The installation consists of two aeroponic tanks of 495 plants located in a 64 m² greenhouse. Plants are held on strips, 5 plants per strip, 99 strips per tank. The specificity of the platform is that the plant rotate constantly so that their root system can be photographed every two hours. The experiment lasted 3 weeks, after which the plants became too large for the platform. The experiment included two tanks. In the first one, plants constantly moved to be pictured every 2 hours (usual set-up on this platform). In the second one, plants moved twice or three times a day to be pictured. This allowed comparing the effect of moving versus non-moving plants, which is a feature often available in the phenotyping platforms but poorly evaluated so far.

Since the UCLouvain platform focuses on the analysis of the root system, the main variable of interest in the experiment is the overall growth of the root system of each plant. Scientists of the UCLouvain platform have developed pipelines³ that allow easy processing of the images captured in the platform to extract quantitative root architec-

²European Plant Phenotyping Network 2020 <https://eppn2020.plant-phenotyping.eu/>

³Here, pipelines are defined as computer programs designed to analyse raw data from phenotyping platforms.

ture information for the spatial models (Lobet & Draye 2013; Lobet, Pagès, *et al.* 2011).

This thesis was divided in four main parts: create an appropriate experimental design for a phenotyping experiment, analyse data from a high-throughput platform, comparing the efficiency of various spatial models to correct for heterogeneous and non-linear spatial trends and developing the appropriate R scripts.

Chapter 3

Material and methods

3.1 Optimal experimental designs

In the context of design of experiments (DOE), optimal designs are the holy grail that every experimenter want. However, researcher often use pre-made designs that fit a large amount of experiments instead of creating an optimal one. Since experimentations exist in all sizes and forms, not all pre-made designs are ideal. Therefore, creating an optimal design is a great choice to ensure that the design fits the experiment and not the other way around. In this section, we explain how and why we created a custom, optimal, design to fit our experiments.

3.1.1 Orthogonal designs

Orthogonal designs are interesting because they guarantees that each main effect and interaction can be estimated independently. Meaning that the effect of one factor or interaction can be estimated separately from the effect of other factors and that the addition or subtraction of terms in the model does not affect the estimates. In a regression or ANOVA-type model, the best linear unbiased predictor (or BLUE) of the regression coefficients is obtained by using the ordinary least squares (OLS) method, because it minimizes the variance of the estimators. These variances are often represented in variance-covariance (VCOV) matrix, where the diagonal is the variance of the estimators and the non-diagonal elements are the pairwise covariances between estimators. The inverse of this matrix is the information matrix, because it summarizes the available information on the models coefficients (a low variance means a lot of information, and inversely). As detailed in Goos & Jones (2011), when the information matrix is diagonal, then the design is said to be orthogonal.

3.1.2 Optimality criteria

To compare different designs, the two main criteria are the D-optimality and the I-optimality. The first one aims at minimizing the variance of the factors effects estimates and is more useful for significance testing. D-optimal designs are therefore more ap-

propriate for screening experiments¹. The second one aims at minimizing the average relative prediction variance over the experimental region. I-optimal designs are focused on prediction and thus are more suited to response surface experiments. There also exists a G-optimality criterion that is similar to the I-optimality criterion but minimizes the maximum prediction variance. Recent work (Rodríguez *et al.* 2010) has shown that I-optimal designs are often better choices than the G-optimal ones. Since this is a screening experiment, only the D-optimality is detailed here. For more information about I-optimal and G-optimal design, refer to Atkinson (2014) and Goos & Jones (2011).

D-optimality

As said previously, for an orthogonal design all the non-diagonal elements of the VCOV matrix are null, and thus, the determinant is simply the product of all the diagonal elements, i.e. the estimators variances. Since the goal is to have the smallest variance of the estimates, the VCOV matrix with the smallest determinant will have the estimates with the smallest variance. Minimizing the determinant of the VCOV matrix is similar to maximizing the determinant of the information matrix. Therefore, the design with factor settings that maximize the determinant of information matrix, will maximize the available information about the models parameters. This design is called the "D-optimal design", where the "D" stands for determinant and the value of the determinant itself is called the "D-optimality criterion".

For any model with two-levels factors and two-factor interaction effects, orthogonal designs will always be D-optimal. However if the number of runs is not a multiple of 4 then there are no orthogonal designs available for two-level factors. This condition offers little flexibility for experimenters and is not always feasible. In contrast, the optimal experimental design approach allows for any number of runs. However, in non-orthogonal designs the variance of the estimates is inflated and the estimates are correlated. Nevertheless this inflation is usually small and the correlation is too small to cause any concerns. Therefore there exist non-orthogonal designs that still maximize the information of the model being estimated. The D-optimal designs may not be unique. For a specified number of runs, several designs might have the maximal value for the determinant of the information matrix.

3.1.3 Generating optimal designs

In order to generate an optimal design, the determinant of the variance-covariance matrix needs to be computed multiple times. Therefore algorithms are used to gain time and avoid errors. Several algorithms exist but the most common one is the coordinate exchange algorithm, created by Meyer & Nachtsheim (1995). It has the advantage to run in polynomial time, which means that the time needed to find an optimal design

¹Screening experiments are experiments designed to evaluate the significance of factors and factor-interactions in a model. The factor are usually two-level factor because they are either present in the model or not. They are opposed to response surface (RS) experiments that are designed to find the optimal settings for the factors. In RS experiments, the significant factors of the model are already determined.

does not explode when the size of the design increases. Another similar algorithm is the point-exchange algorithm, created by Fedorov (1972) and modified several times to speed it up (Atkinson & Donev 1989; Johnson & Nachtsheim 1983). The main drawback of this algorithm is that it needs a list of possible design points as input, which can be quite tedious to do for large designs. In recent years, other types of algorithm such as genetic algorithms (Heredia-Langner, Carlyle, *et al.* 2003; Heredia-Langner, Montgomery, *et al.* 2004), simulating annealing algorithms (Bohachevsky *et al.* 1986; Meyer & Nachtsheim 1988) and tabu search algorithms (Jung & Yum 1996) have been used in experimental designs. While these algorithms maintain a level of performance comparable to more traditional design construction techniques, they are not as popular because they are either far more complex, only feasible in some specific cases or better for some specific models and do not lead to designs that make a significant difference in practice.

Coordinate-exchange algorithm

The coordinate-exchange algorithm proceeds by iterating through the rows of the matrix of factors settings

$$\mathbf{D} = \begin{bmatrix} x_{11} & x_{21} & \dots & x_{k1} \\ x_{12} & x_{22} & \dots & x_{k2} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1n} & x_{2n} & \dots & x_{kn} \end{bmatrix}, \quad (3.1.31)$$

called the design matrix, of an experiment with n runs and k factors. The lines of this matrix essentially represent the coordinates of the runs in the experimental space, where each factor of the experiment is a dimension. This algorithm is called the coordinate-exchange algorithm, because in each iteration of the algorithm, possible changes for every element of the design matrix are considered.

It is straightforward to see that the design matrix \mathbf{D} is a submatrix of the model matrix \mathbf{X} . It can happen the D-optimality criterion is zero. In those cases, the design is called singular and the inverse of the $\mathbf{X}'\mathbf{X}$ matrix does not exist. To avoid singularity, the number of design points (different rows in the design matrix \mathbf{D}) must be greater than or equal to the number of model parameters.

The algorithm starts by generating a random design. For all continuous factors, the algorithm generates random values on the interval $[-1, +1]$. For all factors that are categorical, the algorithm randomly chooses a value in a discrete set of levels. This random starting design is almost always non-singular. If the design happens to be singular, then another new random starting design is computed.

In the next step, the algorithm improves the design on an element-by-element basis. For each element of the starting design, x_{ij} , a change to either -1 or $+1$ is considered, and its impact on the D-optimality criterion is evaluated. The change that increases the value of the D-optimality criterion the most, is kept. After investigating changes in each element of the design, the process is repeated until no element changes within an entire iteration through the factor settings or until a prespecified maximum number of iterations is reached. The obtained design is the best among a set of neighbouring designs but it is often a locally optimal design that is different for each random starting

design. To select the best among all locally optimal designs, the algorithm is repeated a large number of times. The globally optimal design is then selected among all the locally optimal ones, as the one that yields the highest D-optimality criterion.

3.1.4 Generating the design

A custom design was created for the experimental set-up of the phenotyping platform, where the goal was to quantify the genotype and tank effect. Four factors were considered:

Tank In which tank was the plant situated (moving or still).

Strip Which of the 99 strip was used (1 to 99).

Position What was the position on the strip (1 to 5).

Genotype Which one of the 30 genotypes was used (1 to 30).

To fit the design, the design of experiment (DOE) tool of JMP was used. The four categorical factors were specified and *Tank* and *Strip* were set to "very hard to change" and "hard to change", respectively. Two whole plots of 99 sub-plots each were specified to match the tanks and the strips. With 99 strips of five positions inside two tanks, 990 experimental units were available.

Initially the design was supposed to take into account the 99 different strips individually but the program couldn't converge to an optimal design because of its complexity. Instead only 33 strips were considered and the design was replicated 3 times to match the number of runs. Figures 3.1 and 3.2, display a schematic view of the design for the moving and still tank respectively.

The seeds were provided by researchers from the french national institute of agro-nomic research (INRA). A subset of 30 maize hybrids was chosen among an historical series² of maize hybrids that were characterised by their commercial success at the time of their release. Claude Wlecker (LEPSE, INRA) and Carine Palaffre (SMH Maïs, INRA) prepared and sent a total of 30 seeds per genotype plus an extra 150 seeds of genotype called "border genotype" hereafter, used to fill the gaps in the platform left by non germinated seeds and the 90 empty spots left. This genotype is not part of the historical series and was therefore not considered in the design of the experiment.

3.1.5 Design characteristics

²Historical series correspond to varieties that have been cultivated and bred for some time, mainly due to their physiological specificities.

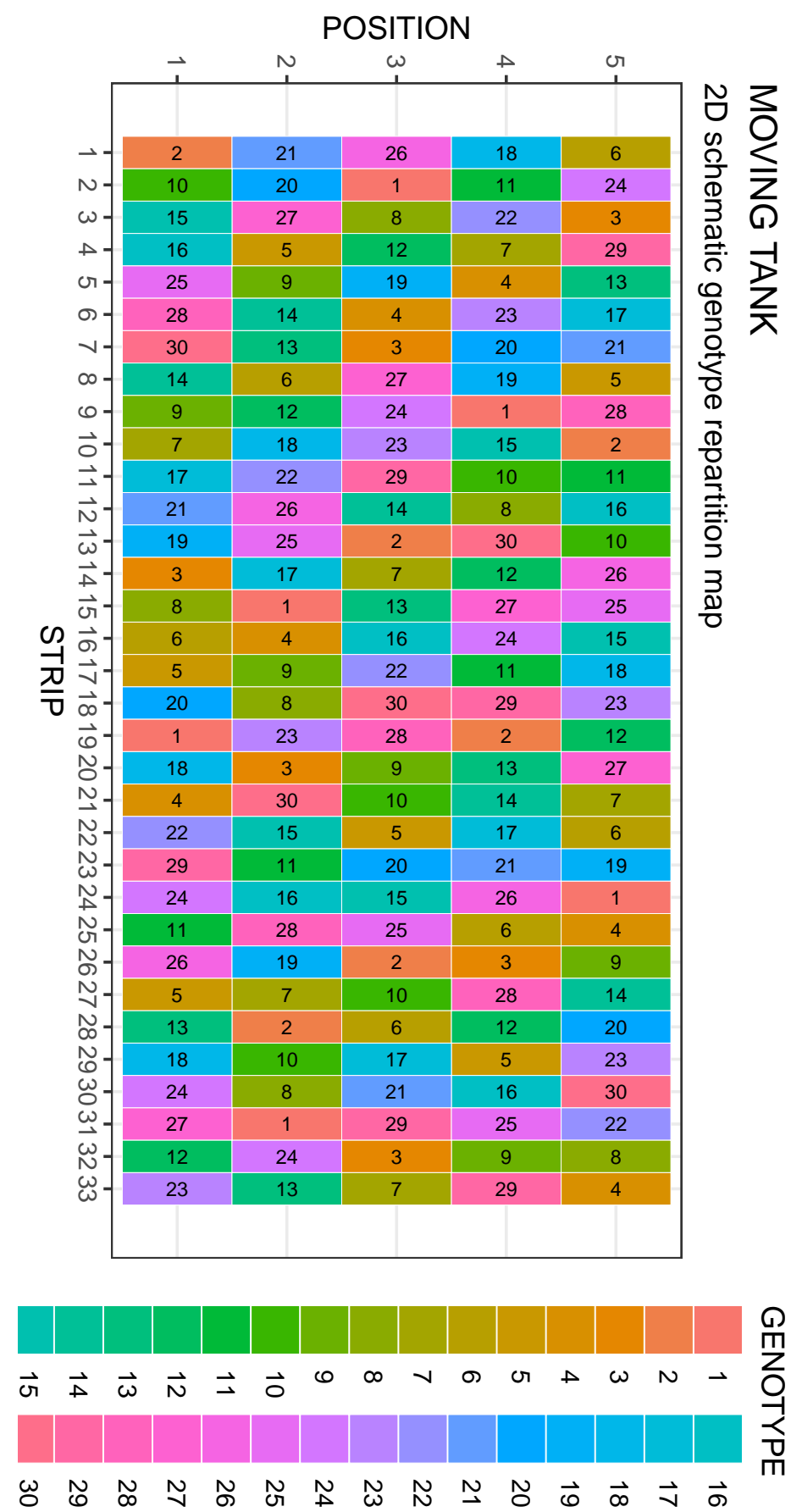


Figure 3.1: 2D schematic representation of the experimental design for 33 individual strips and 30 genotypes in the moving tank.

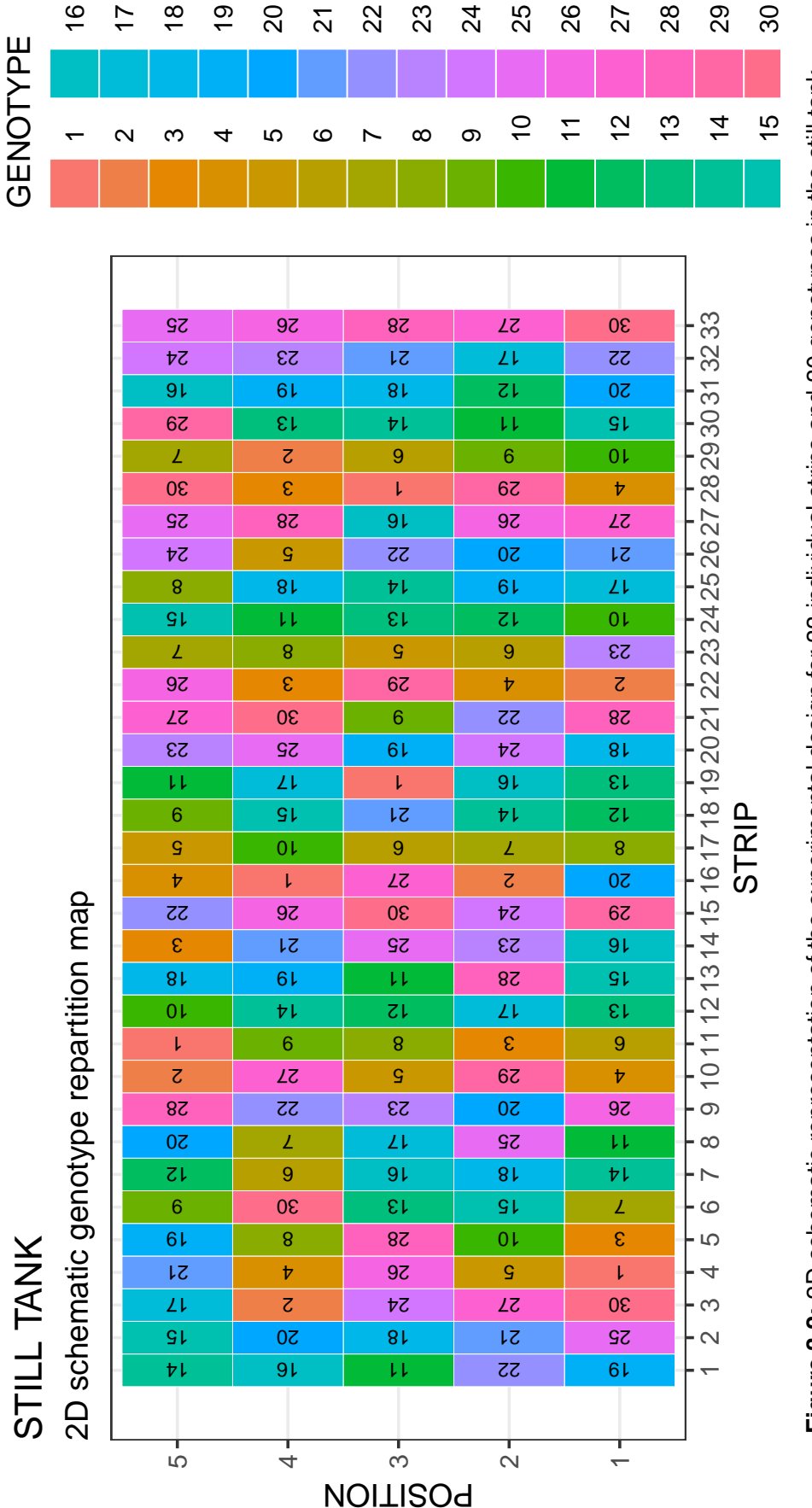


Figure 3.2: 2D schematic representation of the experimental design for 33 individual strips and 30 genotypes in the still tank.

3.2 Phenotyping experiment

The phenotyping experiment took place between February 25th and March 13th 2019. The seeds were first germinated and then transferred onto the platform. After the end of the experiment the plants were weighted, dried and weighted again to obtain dry and fresh weight.

3.2.1 Germination

Previous experiments in the greenhouses showed that germination of maize seeds on the platform often lead to asphyxiation of the seeds. Because of this, the seeds were germinated in an outside germination chamber and were only transferred onto the platform once germinated.

The seeds were placed in a temperature-controlled room at 20°C for 3 days, inside a germination chamber. The chamber consisted of 2 PVC trays to which an air-fog machine was connected, to keep the seeds moist. Inside each tray, PVC plates were disposed diagonally and evenly spaced (fig. 3.3a). On those plates, the seeds were arranged on a filtering paper sheet with ledges to support the weight of the seeds (figure 3.3b and figure 3.3c). The bottom of the trays were filled with water to keep the filtering paper moist. There were 17 plates in total, 15 for the 30 genotypes and 2 for the border genotype (150 seeds dispatched on 2 plates).

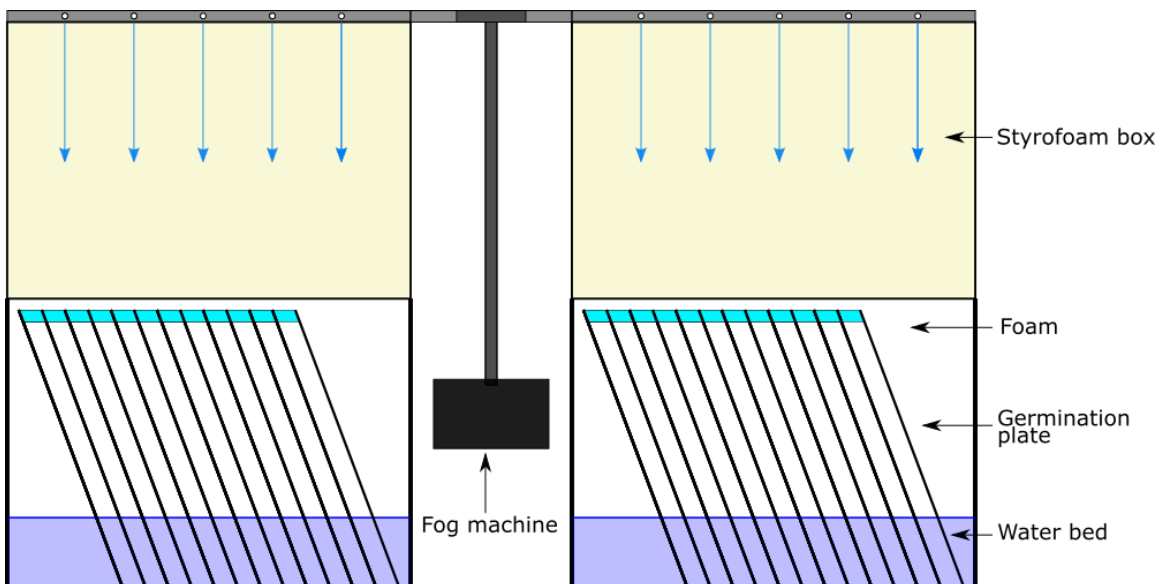
After 3 days into the chamber, not all seeds were germinated, table 3.1 presents the germination rates and mean seed weights for all the genotypes used (including the border genotype). Here the germination rate is defined as the number of germinated seeds over the total number of seeds used in the chamber. However, the actual germination rate (updated rate where the number of fallen seeds, that did not get the chance germinate, is removed from the total) was not recorded in this experiment. It can be an interesting information since germination rates can be genotypes-related too. The non-germination was mainly due to the fact that seeds fell into the water bed and because mold grew on some filtering paper.

3.2.2 Phenotyping platform

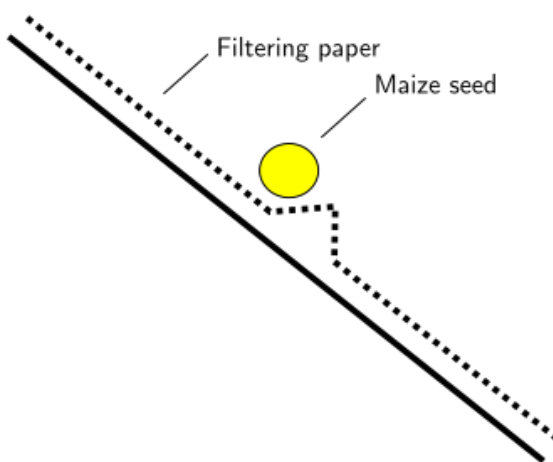
The phenotyping platform is located inside a greenhouse in the facilities of the UCLouvain (Louvain-la-Neuve, Belgium). It consists of two aeroponic tanks on which are arranged 99 styrofoam strips, each with five holes. At the end of each tank is a high definition camera that scans the root system of each plant individually, when it passes in front of it (fig. 3.4a). The strips rotate in a clockwise fashion in the tank and a full rotation is completed in 2 hours. Three sprinklers are placed regularly at the bottom of each side of the tank (fig. 3.4b). The sprinklers spray nutrient solution³ at regular intervals, set by the operator. The spraying pattern (interval and duration) can be differentiated between day and night and can be modified at any moment of the experiment. In this case the patterns were 5 seconds of spraying every 295 seconds all the time. During the experiment, the temperature of the greenhouse was set to 20°C at day and 18°C at night and the lights were on from 6 AM to 10 PM. At the start of the

³The precise concentration of the Hoagland solution is presented in appendix C

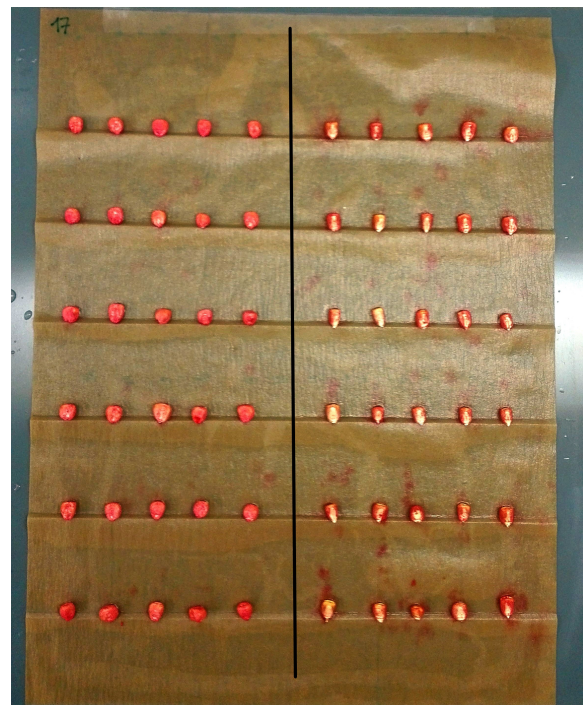
3.2. PHENOTYPING EXPERIMENT



(a) Global schematic view of the germination chamber: a fog machine assure constant humidity in the germination chambers by creating fog at regular intervals (the blue arrows represent the path of the fog). The plates are placed at a 60° angle and 5 cm apart



(b) Schematic view of a germination ledge on a PVC plate: each seed is fixed in position on the ledge by an additional drop of agar solution to avoid any fall-off



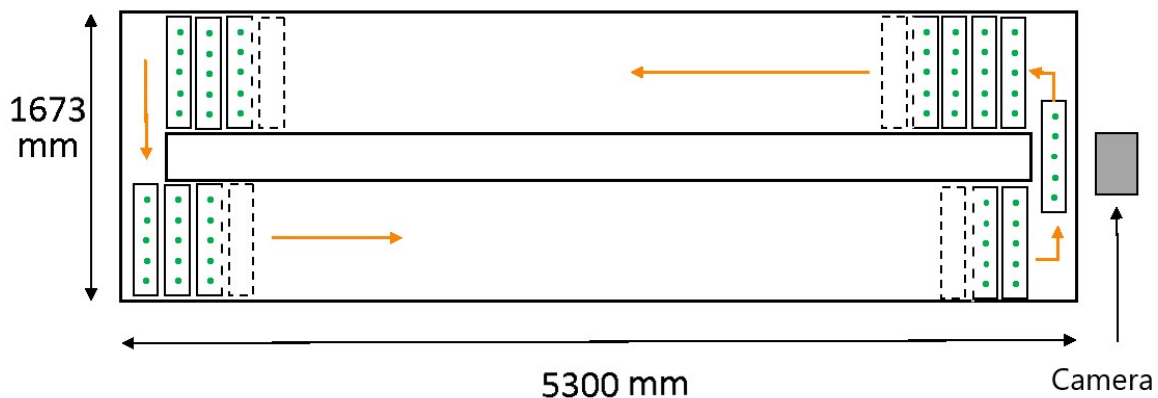
(c) 30 cm by 40 cm PVC plate with seeds on filtering paper (the black line represents the separation between the two genotypes on the plate). Each sheet had 6 rows of 10 seeds with one genotype on the left and one genotype on the right.

Figure 3.3: Germination chamber diagram with detailed view and pictures

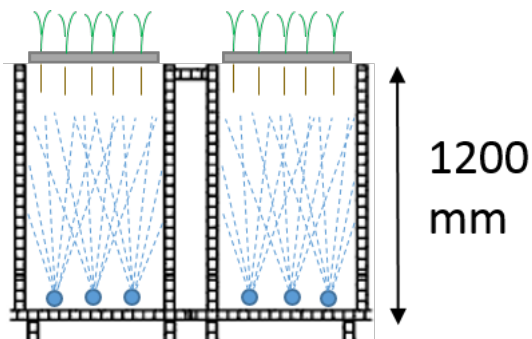
CHAPTER 3. MATERIAL AND METHODS

experiment seeds were placed inside a foam cork and then placed inside a hole on a strip (fig 3.4c). They were placed at the bottom to allow the root system to grow freely. The corks are drilled vertically to let the leaf system develop with less resistance and allow a direct access to sunlight. More information about the platform is available in appendix D.

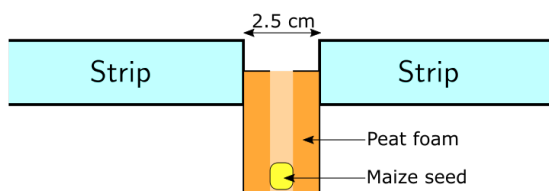
After 3 days in the germination chamber, the germinated seeds were transferred onto the platform following the created design, and the non-germinated seeds were discarded. The first tank was constantly moving but only scanning the root systems once a day, while the second tank was only moving once a day to scan the root system of the plants and stayed still the rest of the day.



(a) Schematic view of an aeroponic tank: plants are hold on strips, 5 plants per strip (green dots on layout). There are 99 strips in the tank for a total of 495 plants/tank. Strips move in the direction indicated by orange arrows



(b) Transversal schematic view of an aeroponic tank of the platform: at the bottom of each tank, sprinklers (represented in blue in the layout) are disposed at regular interval and spray nutritive solution



(c) Close up schematic view of a strip: inside each hole, seeds are placed inside a pierced peatfom cork to allow the root system to develop frelly

Figure 3.4: Detailed diagrams about the phenotyping platform

Table 3.1: Germination rate and mean seed weight for each genotype used. (there is no data concerning the germination rate of the border genotype because it was not measured).

Genotype	Germination rate (%)	Mean seed weight (g)
1	80.00	0.28
2	86.67	0.36
3	96.67	0.36
4	73.33	0.28
5	100.00	0.32
6	96.67	0.24
7	96.67	0.19
8	70.00	0.31
9	96.67	0.33
10	96.67	0.25
11	60.00	0.33
12	93.33	0.27
13	90.00	0.24
14	86.67	0.31
15	56.67	0.35
16	90.00	0.28
17	90.00	0.30
18	86.67	0.26
19	100.00	0.26
20	86.67	0.28
21	86.67	0.32
22	53.33	0.30
23	73.33	0.28
24	100.00	0.16
25	96.67	0.19
26	96.67	0.25
27	96.67	0.28
28	83.33	0.36
29	93.33	0.30
30	73.33	0.35
31	/	0.38

3.3 Data processing

After 15 days, the plants were considered fully grown and the experiment was stopped. The leaf and root systems were separated and weighted individually on scales precise to 0.001 g. They were then dried for 3 days in a 70°C oven. After the drying process, they were weighted again. For each plant, the remaining of the seed was consistently kept on the root system. From the weighting, four variables were kept for the spatial analysis:

- *FRESH_RS*: fresh weight of the root system
- *FRESH_LS*: fresh weight of the leaf system
- *DRY_RS*: dry weight of the root system
- *DRY_LS*: dry weight of the leaf system

For some plants, the germinated seeds placed on the platform did not fully grow or had an abnormal growth, but all the plants were still weighted to avoid leaving out any data. Therefore some data points need to be handled more carefully, as they do not represent the genotype's growth correctly. However, a growth representative of the genotype is hard to define because the influence of the conditions on each genotype is unknown. Therefore, instead of choosing which plants are outliers in a binary way, we attributed weights to each plant to express the quality of the data. Those weights were established by reviewing the final root scan of each plant and checking the different factors that could have an influence. The factors chosen are the following:

- *NO_RS*: no additional root to the primary root
- *NO_LS*: no visible leaf system
- *BAD_LS*: leaf system grew under the strip (or abnormally in general)
- *NO_SEED*: no seed (or an empty cork) present on this position
- *NOT_FG*: plant not fully grown
- *OVERLAP*: leaf (or root) system of another plant overlaps on the root scan
- *OK*: no influential factors

A visualization of those factors with example pictures is presented in figure 3.5. Some plants were attributed several influential factors, but *OK*, *NO_SEED* and *NOT_FG* were considered as exclusive. The factor attribution was ambiguous for some pictures, in those cases the plant were considered *OK* to avoid losing any data points. Following the determination of the factors, weights were attributed to each variable according to the weight matrix, presented in table 3.2.

Table 3.2: Weight attribution matrix for the different factors and variables. LS weight is both the fresh and dry weight for leaf system and RS weight is the same for the root system

Code	Weight
NO_RS	1
NO_LS	2
BAD_LS	3
NOT_FG	4
NO_SEED	0
OK	5
OVERLAP	5

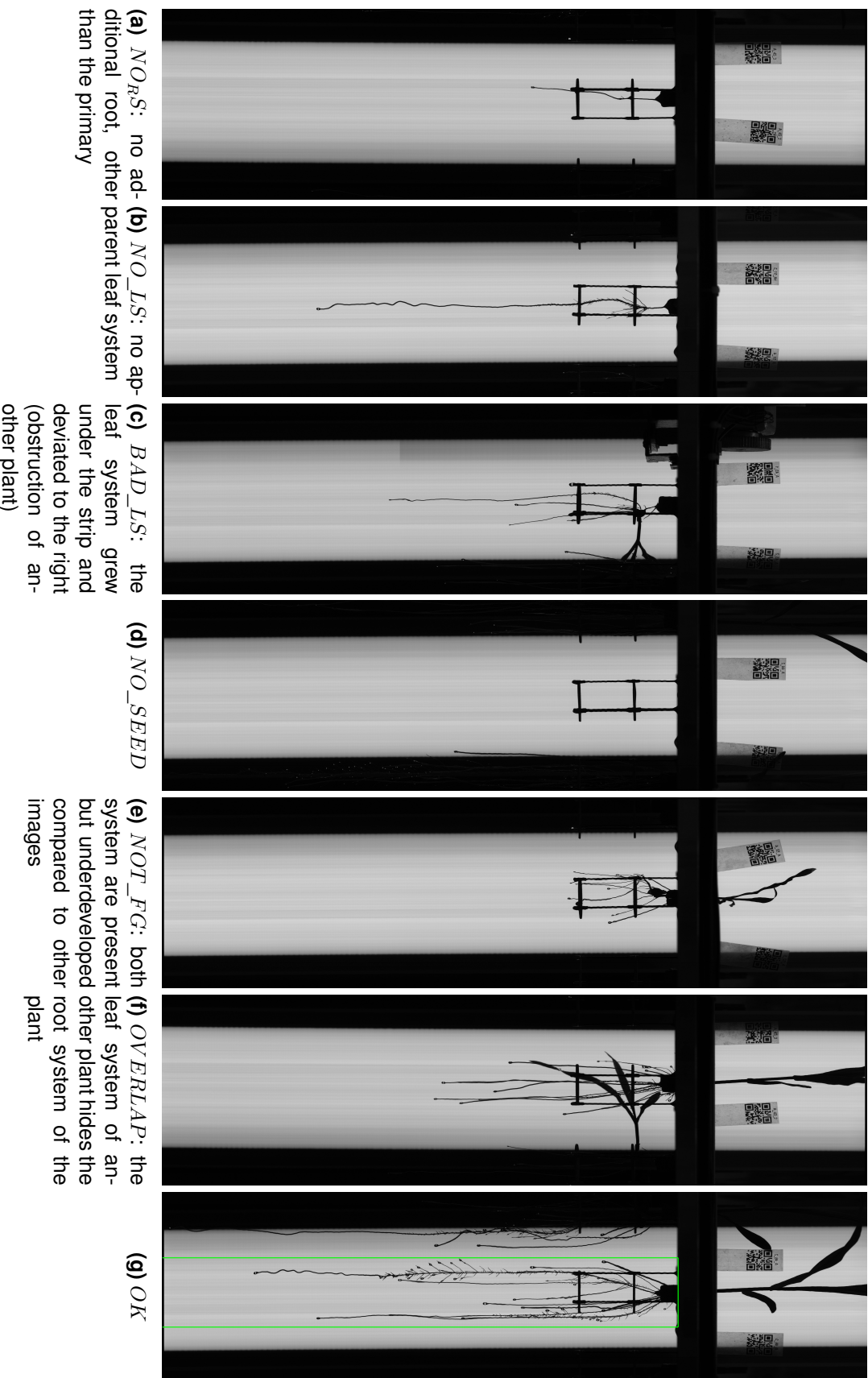


Figure 3.5: Example pictures of the influential factors chosen for the weight attribution.

3.4 SpATS model

In this section, the SpATS model is introduced. For a more thorough treatment of the model and all its components, see M. X. Rodríguez-Álvarez, Boer, *et al.* (2016).

Consider a field trial of n plots arranged in a rectangular grid, where the plot positions are collected in vectors of row (\mathbf{r}) and column (\mathbf{c}) coordinates. If \mathbf{y} is the vector of plot data in field order, a common model for \mathbf{y} , to use as a starting point is

$$\mathbf{y} = \mathbf{1}_n \beta_0 + \mathbf{Z}_r \mathbf{c}_r + \mathbf{Z}_c \mathbf{c}_c + \varepsilon \quad (3.4.01)$$

where $\mathbf{1}_n$ is a column-vector of ones of length n , \mathbf{c}_r and \mathbf{c}_c are, respectively, the random effect coefficients for the rows and columns and associated matrix \mathbf{Z}_r and \mathbf{Z}_c . To fully capture complex spatial patterns, a smooth bivariate surface jointly defined over the row and column positions is added to the model, which becomes

$$\mathbf{y} = f(\mathbf{u}, \mathbf{v}) + \mathbf{Z}_r \mathbf{c}_r + \mathbf{Z}_c \mathbf{c}_c + \varepsilon \quad (3.4.02)$$

where \mathbf{u} and \mathbf{v} are, respectively, the vector of row and columns positions and where $f(\cdot, \cdot)$ represents the smooth bivariate function. Note that the intercept term, β_0 is embedded into $f(u, v)$. To better understand this function, let us decompose it in a nested-ANOVA structure

$$\begin{aligned} f(\mathbf{u}, \mathbf{v}) &= \underbrace{\mathbf{1}_n \beta_0 + \mathbf{u} \beta_1 + \mathbf{v} \beta_2 + \mathbf{u} \odot \mathbf{v} \beta_3}_{\text{Bilinear polynomial}} \\ &\quad + \underbrace{f_u(\mathbf{u}) + f_v(\mathbf{v}) + \mathbf{u} \odot h_v(\mathbf{v}) + \mathbf{v} \odot h_u(\mathbf{u}) + f_{u,v}(\mathbf{u}, \mathbf{v})}_{\text{Smooth part}} \end{aligned} \quad (3.4.03)$$

where \odot denotes the element-wise matrix product⁴. There are now two components to the model: a bilinear polynomial part(parametric) and a smooth part (non-parametric). The parametric part includes the linear trends along rows (β_1) and columns (β_2) as well as a linear interaction trend (β_3). The smooth part models the deviation from the compound linear trend, and can be decomposed in the following elements:

- $f_u(\mathbf{u})$ is a smooth trend along the rows, identical for all columns (i.e., a main smooth effect).
- $f_v(\mathbf{v})$ is a smooth trend along the columns, identical for all rows.
- $\mathbf{v} \odot h_u(\mathbf{u})$ and $\mathbf{u} \odot h_v(\mathbf{v})$ are linear-by- smooth interaction trends. For instance, $\mathbf{u} \odot h_v(\mathbf{v})$ is a varying coefficient surface trend, consisting of functions, linear in the rows, for each column, but with slopes that change smoothly along the columns, h_v .
- $f_{u,v}(\mathbf{u}, \mathbf{v})$ is a smooth-by-smooth interaction trend jointly defined over the row and column directions.

⁴See appendix A for details about the element-wise matrix product.

In total, six components are used to model the surface f . This may seem like a lot but this allows the translation of model 3.4.02 into a standard mixed model. An interesting property of this proposal is that since u and v are row and column position, it allows depicting the spatial trend in a grid finer than the number of rows and columns. Figure 3.6 shows an example of those six components in the context of a barley uniformity performed by K. A. Williams (1988). It shows clearly how the additional components, help to capture small variations in the spatial data.

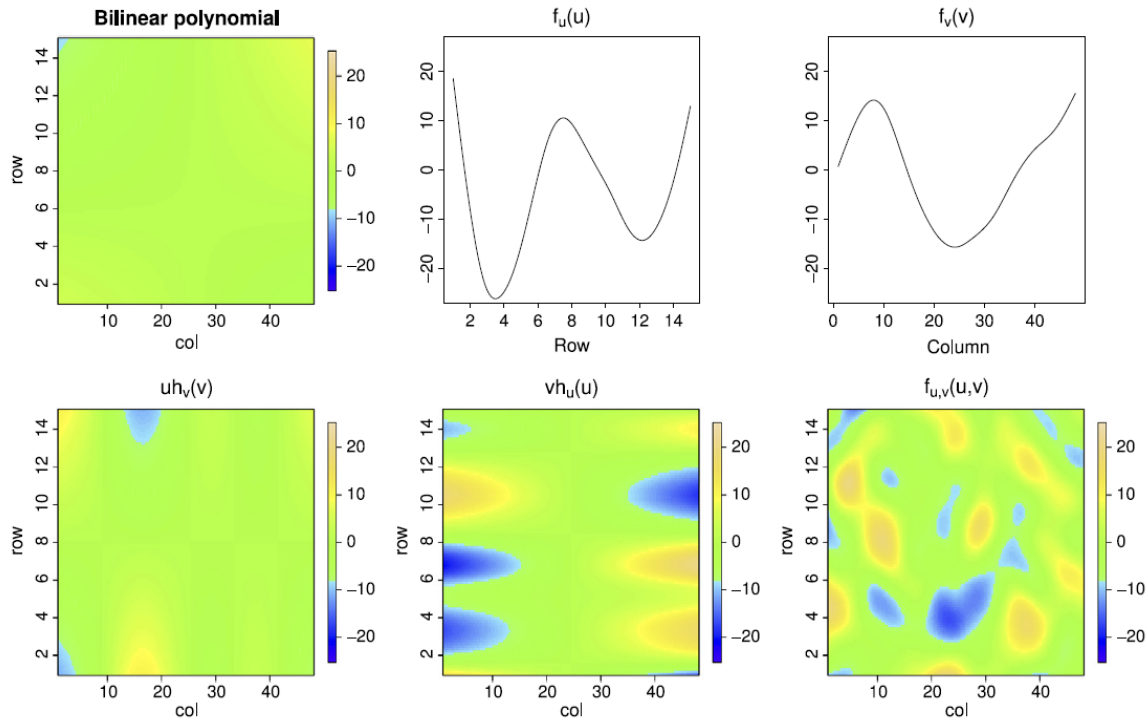


Figure 3.6: Bilinear and smooth components of the PS-ANOVA decomposition of the estimated spatial trend for the barley uniformity data from Rodriguez-Alvarez *et al.* (2018).

3.4.1 Modelling using P-splines

The functions f_u , f_v , h_u and h_v are constructed with variations on one-dimensional P-splines, while $f_{u,v}$ is based on tensor products P-splines.

For clarity's sake, let us consider a model only containing a smooth bivariate surface and an error term

$$y_i = f(u_i, v_i) + \varepsilon_i, \text{ with } \varepsilon_i \sim N(0, \sigma^2). \quad (3.4.11)$$

Lee, Durbán & P. Eilers (2013) show that it can be represented using B-splines⁵. Let us form two B-splines basis:

1. one for the columns, \hat{B} with $b_{il} = \hat{B}_l(u_i)$, where $\hat{B}_l(u_i)$ is the l th B-spline of the basis, evaluated at u_i

⁵See appendix A for details about B-splines and P-splines.

2. and one for the rows, $\check{\mathbf{B}}$ with $b_{ip} = \check{B}_p(v_i)$, where $\check{B}_l(v_i)$ is the p th B-spline of the basis, evaluated at v_i .

Then, the smooth-by-smooth interaction can be written using those basis

$$f(u_i, v_i) = \sum_{l=1}^L \sum_{p=1}^P \hat{B}_l(u_i) \check{B}_p(v_i) \alpha_{lp}, \quad (3.4.12)$$

where $\boldsymbol{\alpha} = (\alpha_{11}, \dots, \alpha_{1P}, \dots, \alpha_{LP})^t$ is a vector of unknown regression coefficients of dimension $(LP \times 1)$. Note that $\hat{\mathbf{B}}$ and $\check{\mathbf{B}}$ are matrices of order $n \times L$ and $n \times P$ respectively, where L and P are the number of the B-spline basis functions. Dierckx (1995) shows that, in the P-spline framework, the smooth-by-smooth interaction $f(u_i, v_i)$ is modelled by the tensor product of B-splines bases. Then, we can write, in matrix notation,

$$\mathbf{B} = \hat{\mathbf{B}} \square \check{\mathbf{B}} = (\hat{\mathbf{B}} \otimes \mathbf{1}_L^t) \odot (\mathbf{1}_P^t \otimes \check{\mathbf{B}}), \quad (3.4.13)$$

where the operation \square is defined in terms of the Kronecker product of two matrices (denoted by \otimes) and the element-by-element multiplication of two matrices (denoted by \odot)⁶. Therefore model (3.4.11) can be written in matrix notation:

$$\mathbf{y} = \mathbf{B}\boldsymbol{\alpha} + \boldsymbol{\epsilon}. \quad (3.4.14)$$

The coefficients of this parametric model can be estimated by minimizing the sum of squares. The explicit solution is then:

$$\hat{\boldsymbol{\alpha}} = (\mathbf{B}^t \mathbf{B})^{-1} \mathbf{B}^t \mathbf{y} \quad (3.4.15)$$

To prevent over-fitting, P. H. C. Eilers & Marx (1996) propose to incorporate a discrete penalty on the coefficient associated to adjacent B-splines. As described in details in appendix A.3, this penalty also determines the smoothness of the splines. The solution of equation 3.4.14 then becomes

$$\hat{\boldsymbol{\alpha}} = (\mathbf{B}^t \mathbf{B} + \mathbf{P})^{-1} \mathbf{B}^t \mathbf{y}, \quad (3.4.16)$$

where \mathbf{P} is the penalty matrix. The details of this solution are presented in appendix A.4, but the important point to remember here, is that the smoothness of the bivariate surface is defined by the penalty matrix, which only depend on two tuning parameters λ (smoothing along the columns) and $\check{\lambda}$ (smoothing along the rows).

3.4.2 Mixed model based smoothing parameter selection

As explained in M. X. Rodríguez-Álvarez, Boer, *et al.* (2016), \mathbf{P} is rank-deficient, meaning that the rank⁷ is smaller than the number of rows and/or columns, and this causes numerical instability when applying mixed model estimation techniques. To obtain a full-rank penalty matrix, the key is to write model 3.4.14 as

$$\mathbf{B}\boldsymbol{\alpha} = \mathbf{X}_s \boldsymbol{\beta}_s + \mathbf{Z}_s \boldsymbol{c}_s. \quad (3.4.21)$$

⁶See appendix A for details about the Kronecker, and the element-wise products

⁷The rank of a matrix is the maximum number of linearly independent row-vectors.

There are now two bases: \mathbf{X}_s , with coefficients that are not penalized at all, and \mathbf{Z}_s , with a size penalty on its coefficients. This decomposition follows the proposal by Lee & Durbán (2011), based on eigenvalue decomposition which gives rise to a diagonal penalty matrix.

The two bases have the following structures:

$$\mathbf{X}_s = [\mathbf{1}_n, \mathbf{u}, \mathbf{v}, \mathbf{u} \odot \mathbf{v}] \quad \text{and} \quad \mathbf{Z}_s = [\mathbf{Z}_v, \mathbf{Z}_u, \mathbf{Z}_v \square \mathbf{u}, \mathbf{v} \square \mathbf{Z}_u, \mathbf{Z}_v \square \mathbf{Z}_u], \quad (3.4.22)$$

where \mathbf{u} and \mathbf{v} are still, respectively, the vectors of row and column positions. Here \mathbf{Z}_u and \mathbf{Z}_v are penalized version of the B-splines basis \mathbf{B} (rows) and $\hat{\mathbf{B}}$ (columns). This new way of writing the problem leads to another penalty matrix $\tilde{\mathbf{P}}$, which is a block diagonal matrix. Each block of $\tilde{\mathbf{P}}$ corresponds to a block in \mathbf{Z}_s . Similarly to \mathbf{P} , the penalty matrix of the previous section, this new penalty matrix only depends on the two tuning parameters λ (smoothing along the columns) and $\hat{\lambda}$ (smoothing along the rows). Figure 3.7 presents a diagram clarifying the structures and relations of the different matrices presented throughout this section.

This reformulation provides the ANOVA type decomposition discussed in the previous section (3.4.03), and explains how the bilinear smooth surface can be modelled using P-splines and tensor products of P-splines. The block structure of \mathbf{X}_s and \mathbf{Z}_s implies

$$\begin{aligned} f(\mathbf{u}, \mathbf{v}) &= \mathbf{X}_s \boldsymbol{\beta}_s + \mathbf{Z}_s \mathbf{c}_s \\ &= \mathbf{1}_n \beta_0 + \mathbf{u} \beta_1 + \mathbf{v} \beta_2 + \mathbf{u} \odot \mathbf{v} \beta_3 \\ &\quad + \underbrace{f_v(\mathbf{v})}_{\mathbf{Z}_v \mathbf{c}_{s1}} + \underbrace{f_u(\mathbf{u})}_{\mathbf{Z}_u \mathbf{c}_{s2}} + \underbrace{\mathbf{u} \odot h_v(\mathbf{v})}_{[\mathbf{Z}_v \square \mathbf{u}] \mathbf{c}_{s3}} + \underbrace{\mathbf{v} \odot h_u(\mathbf{u})}_{[\mathbf{v} \square \mathbf{Z}_u] \mathbf{c}_{s4}} + \underbrace{f_{u,v}(\mathbf{u}, \mathbf{v})}_{[\mathbf{Z}_v \square \mathbf{Z}_u] \mathbf{c}_{s5}}, \end{aligned} \quad (3.4.23)$$

where \mathbf{c}_{sk} ($k = 1, \dots, 5$) contains the elements of \mathbf{c}_s that correspond to the k th block of \mathbf{Z}_s , i.e. $\mathbf{c}_s = (c_{s1}^t, \dots, c_{s5}^t)^t$. The details about the specific block component of \mathbf{Z}_s and the computation of the new penalty matrix are available in the paper of Rodriguez-Alvarez *et al.* (2018) and the appendices therein.

Therefore, using this new notation, model 3.4.11 that only contains a smooth bivariate surface and an error term can be rewritten in the following way:

$$\mathbf{y} = \mathbf{X}_s \boldsymbol{\beta}_s + \mathbf{Z}_s \mathbf{c}_s + \boldsymbol{\varepsilon}, \quad \text{with } \boldsymbol{\varepsilon} \sim N(\mathbf{0}, \sigma^2 \mathbf{I}_n) \quad \text{and } \mathbf{c}_s \sim N(\mathbf{0}, \mathbf{G}_s), \quad (3.4.24)$$

where $\mathbf{G}_s = \sigma^2 \tilde{\mathbf{P}}^{-1}$. It is straightforward to see that \mathbf{G}_s also has a block diagonal structure, similar to that of $\tilde{\mathbf{P}}$ (this structure is also represented on figure 3.7). However, \mathbf{G}_s depends on two different parameters, $\sigma^2 = \sigma/\lambda$ and $\hat{\sigma}^2 = \sigma/\hat{\lambda}$, which are variances parameters. As shown in the diagram in figure 3.7, the same variance parameters control the smoothness of the both the main effects and interactions terms. This prevents the use of standard mixed models software for estimation since \mathbf{G}_s has its last block depending on both σ^2 and $\hat{\sigma}^2$, but in a non-linear way. Even though M. X. Rodríguez-Alvarez, Lee, *et al.* (2015) presented a specialized algorithm to deal with this issue, here the PS-ANOVA decomposition approach (Lee, Durbán & P. Eilers 2013) is used to allow the use of standard mixed model estimation procedures. Lee, Durbán & P. Eilers (2013) therefore propose to use a different variance component for each smooth

component in \mathbf{G}_s , thus redefining this matrix as a linear function of variance parameters:

$$\mathbf{G}_s = \bigoplus_{k=1}^5 \mathbf{G}_{sk} = \text{blockdiag } (\mathbf{G}_{s1}, \mathbf{G}_{s2}, \mathbf{G}_{s3}, \mathbf{G}_{s4}, \mathbf{G}_{s5}), \quad (3.4.25)$$

where \mathbf{G}_{sk} is the k th block of the \mathbf{G}_s matrix, depending on the specific variance component σ_{sk}^2 . In other words, here the tensor product P-splines mixed model is represented as the sum of 5 sets of mutually independent Gaussian random components \mathbf{c}_{sk} , each depending on one variance σ_{sk}^2 ($k = 1, \dots, 5$).

Within this mixed model framework, the smoothing parameters, defined earlier as the ratio between the residual variance and the corresponding variance effect $\lambda_{sk} = \sigma_e^2 / \sigma_{sk}^2$, are determined by restricted maximum likelihood (REML). Therefore the smoothness of the spatial surface is tuned by five distinct parameters, applying anisotropic (direction-dependant) smoothing. This parametrization provides flexibility to account for both global and local variations in the field. Furthermore, the decomposition of $f(\mathbf{u}, \mathbf{v})$ enables a more explicit interpretation of the main patterns of spatial variation (Rodriguez-Alvarez *et al.* 2018).

3.4.3 Spatial models for field trials

The tensor product P-spline, presented in the previous section, constitutes the base for the analysis of agricultural field trials because it allows the modeling of the random spatial variation typically presented in a field. On top of this spatial field, we need to build up a more complex models in order to account for the genetic variation, the different tanks, strips and positions. From now on, we therefore consider the following linear mixed model

$$\mathbf{y} = \underbrace{\mathbf{X}_s \boldsymbol{\beta}_s + \mathbf{Z}_s \mathbf{c}_s}_{f(u,v)} + \mathbf{X}_d \boldsymbol{\beta}_d + \mathbf{Z}_d \mathbf{c}_d + \boldsymbol{\varepsilon}, \text{ with } \mathbf{c}_s \sim N(\mathbf{0}, \mathbf{G}_s) \text{ and } \mathbf{c}_d \sim N(\mathbf{0}, \mathbf{G}_d), \quad (3.4.31)$$

where \mathbf{X}_s , \mathbf{Z}_s and \mathbf{G}_s are defined in the previous section and form the mixed model expression of the smooth spatial surface, and \mathbf{X}_d and \mathbf{Z}_d represent column-partitioned matrices, associated respectively with fixed and random components. Since we do not have any check genotypic varieties or resolvable block effect, the only extra fixed effects are: an intercept (1_n) and the tank effect (t). We assume that the \mathbf{X}_d matrix has full-rank. The position on the strip (p) and strip (s) variables are added as random effects in \mathbf{Z}_d . Therefore $\mathbf{X}_d = [\mathbf{X}_{1_n}, \mathbf{X}_t]$, $\mathbf{Z}_d = [\mathbf{Z}_{dp}, \mathbf{Z}_{ds}]$ and $\mathbf{G}_d = \text{blockdiag } (\mathbf{G}_{dp}, \mathbf{G}_{ds})$. We further assume that \mathbf{c}_s and \mathbf{c}_d are independent. To keep the notation simple, we rewrite model (3.4.31) as

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\mathbf{c} + \boldsymbol{\varepsilon}, \text{ with } \mathbf{c} \sim N(\mathbf{0}, \mathbf{G}) \text{ and } \boldsymbol{\varepsilon} \sim N(\mathbf{0}, \sigma^2 \mathbf{I}_n) \quad (3.4.32)$$

where $\mathbf{X} = [\mathbf{X}_s, \mathbf{X}_{d_1}, \mathbf{X}_{d_t}]$, $\mathbf{Z} = [\mathbf{Z}_s, \mathbf{Z}_{dp}, \mathbf{Z}_{ds}]$, and

$$\mathbf{G} = \text{blockdiag } (\mathbf{G}_s, \mathbf{G}_{dp}, \mathbf{G}_{ds}) \quad (3.4.33)$$

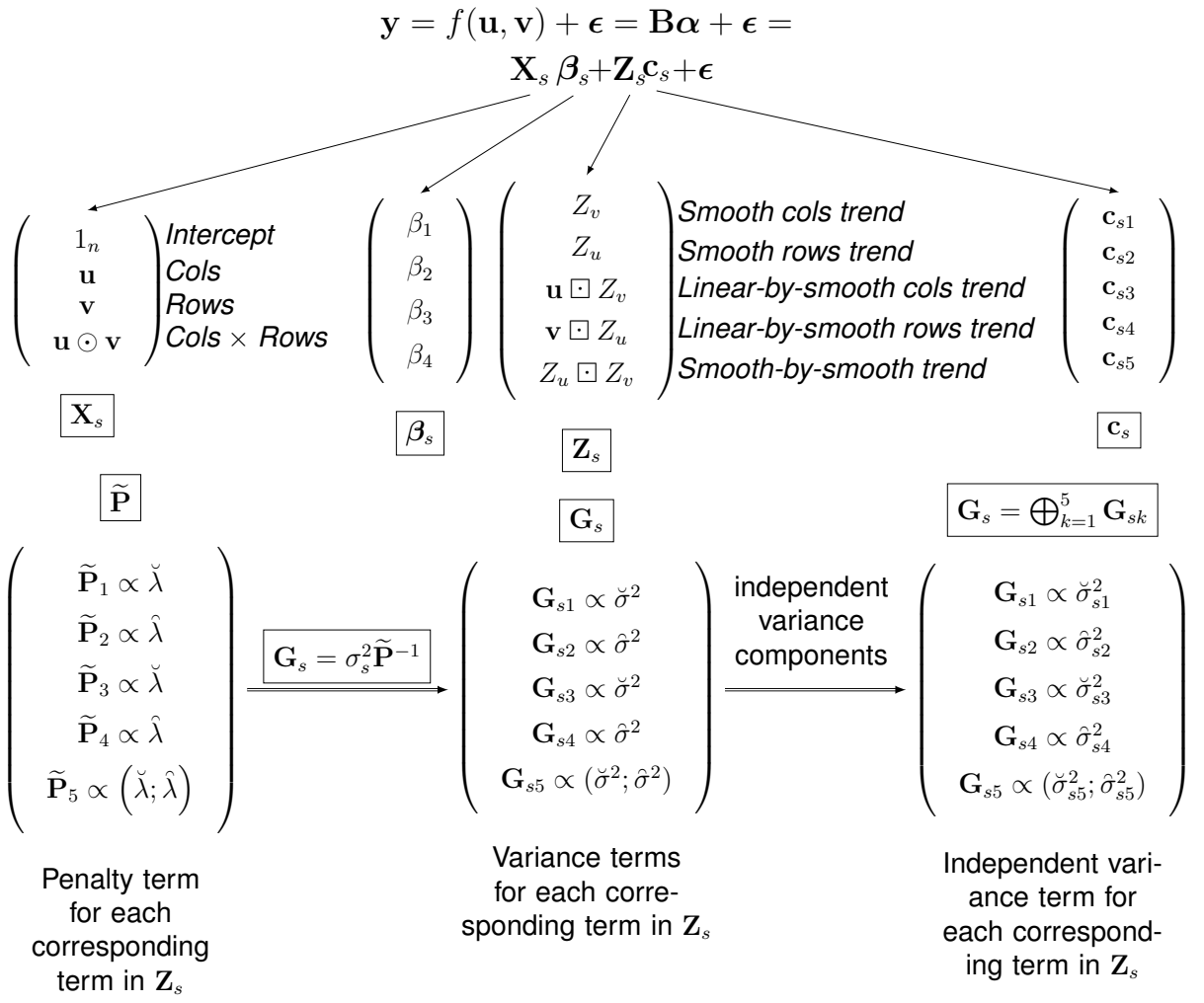


Figure 3.7: Diagram detailing the structure of the matrices used in this section. All matrices are block diagonal matrix with each element represented on the diagram, being an individual block. The symbol \propto shows how each block of the $\widetilde{\mathbf{P}}/\mathbf{G}_s$ matrix relates to the tuning/variance parameters. The last block of both the $\widetilde{\mathbf{P}}$ and \mathbf{G}_s matrices depends on both parameters but in a non-linear way.

3.4.4 Model estimation

With all these specifications in mind, the model was fitted using cubic B-splines and second-order penalties. These settings are commonly used and allow flexibility of the model (M. X. Rodríguez-Álvarez, Boer, *et al.* 2016; Rodriguez-Alvarez *et al.* 2018; M. X. Rodríguez-Álvarez, Lee, *et al.* 2015). We used 99 and 5 equally spaced knots for the P-splines, corresponding to the strips and positions, respectively. In this way there was approximately one knot for every plot. Then there was a total of 362 model parameters to be estimated for the smooth surface (and 501 for the whole model). As Rodriguez-Alvarez *et al.* (2018) explains, the number of knot is not critical since the optimization of the fit to the data is essentially dependant on the smoothing parameters. The estimation procedure was performed using the R-package SpATS (M. Rodríguez-Álvarez *et al.* 2016). This package provides a REML-based estimation of the variances components and computes the best linear unbiased estimators (BLUEs) of the fixed effects and the empirical best linear unbiased predictors (BLUPs) of the random effects. A useful by-product of this computation is the effective dimension associated to each random effect.

Effective dimensions

In P-splines methodology, the effective dimension (ED) measures the complexity of the model components (P. H. Eilers *et al.* 2015), it is similar to the more common concept of effective degree of freedom (Buja *et al.* 1989). It is computed as the trace of the hat matrix \mathbf{H} . If we only take the spatial part of our model (equation 3.4.24), we can write:

$$\begin{aligned}\tilde{\mathbf{y}} &= \mathbf{Hy} \\ \tilde{f}(\mathbf{u}, \mathbf{v}) &= \mathbf{X}_s \hat{\boldsymbol{\beta}}_s + \mathbf{Z}_s \tilde{\boldsymbol{c}}_s, \\ &= \mathbf{H}_{\beta} \mathbf{y} + \mathbf{H}_s \mathbf{y}\end{aligned}\tag{3.4.41}$$

where \mathbf{H}_{β} is the hat matrix of the fixed components and \mathbf{H}_s is the hat matrix of the random components, also known as the smoother matrix. In this context, the sum of the diagonal elements of \mathbf{H}_s expresses the number of parameters effectively involved in the modelling of the spatial surface. This decomposition is allowed from the PS-ANOVA structure of the spatial model. Following this, the smoother matrix can be further decomposed according to the five additive and interaction smooth components of the smooth bivariate surface, giving $\mathbf{H}_s = \sum_{k=1}^5 \mathbf{H}_{s_k}$, and we can compute the individual effective dimension for each component (ED_{s_k}). As explained in M. X. Rodríguez-Álvarez, Boer, *et al.* (2016), the effective dimension varies with the smoothing parameter:

$$\begin{aligned}\lambda_{s_k} = \frac{\sigma_e^2}{\sigma_{s_k}^2} \rightarrow \infty &\text{ then } ED_{s_k} \rightarrow 0 \\ \lambda_{s_k} = \frac{\sigma_e^2}{\sigma_{s_k}^2} \rightarrow 0 &\text{ then } ED_{s_k} \rightarrow \text{upper bound },\end{aligned}\tag{3.4.42}$$

where the upper bound is determined by the number of knots.

Consequently, the total effective dimension ED_s can be interpreted as a measure of the magnitude of field variations with larger values indicating more intense spatial

patterns. In addition, the partial effective dimensions ED_{s_k} are indicative of the contribution of each component to the fitted surface, and reflect the complexity of the spatial pattern.

Generalized heritability

As we just said, the effective dimension is a useful tool to measure the relative importance of each spatial component, and to compare them. However, the effective dimension of the genetic component can't be compared to the rest easily. This is where the concept of heritability comes in handy. In classical genetic models, the standard heritability is defined as the proportion of the total (phenotypic) variation that is attributable to the genetic component. In their paper, Rodriguez-Alvarez *et al.* (2018) show the link between this definition and the genetic effective dimension. They go further by saying that this definition can be extended to the concept of generalized heritability (Oakey *et al.* 2006), which is similar to the standard one, except that the genotype are considered independent. This is useful when there are no pedigree/marker information available about the genotypes. They establish the following relationship:

$$H_g^2 = \frac{ED_g}{n_g} = 1 - \frac{\overline{PEV}}{\sigma_g^2}, \quad (3.4.43)$$

where n_g is the number of genotypes and \overline{PEV} is the prediction error variance for the genotype BLUPs. Since n_g is also the upper bound of the effective genetic dimension, H_g^2 can be seen as a shrinkage factor. Given that our experiment does not incorporate a genetic relationship matrix, we'll use this definition to properly interpret the genetic effect in our model. The right-hand term of the equation, is equivalent to the concept of generalized heritability developed by Welham *et al.* (2010). We can profit from that to compare the heritability between the spats model and the $AR \times AR$ model.

3.5 Standard spatial models

In this section the $AR \times AR$ model, and its extension to the linear variance (LV) model, are presented. For more detailed information about the original $AR \times AR$ model, consult Gilmour *et al.* (1997). For information about the extensions of the model, see Piepho & E. Williams (2010) and E. R. Williams (1986).

As explained in the literature review chapter, the standard spatial models (SS models), such as the $AR \times AR$ model, use a spatially dependant error term to adjust for the local trends, whereas the SpATS model encompasses them in the smooth surface. To estimate the spatial covariance structure of this error process, Gilmour *et al.* (1997) advocate the use of variograms and more precisely, the use of auto-regressive processes ($AR(1)$) to model those variograms. The general representation of the SS model is very similar to the SpATS model:

$$y = \mathbf{X}\beta + \mathbf{X}_s\beta_s + \mathbf{Z}\mathbf{c} + \mathbf{Z}_s\mathbf{c}_s + \boldsymbol{\xi} + \boldsymbol{\varepsilon}, \quad (3.5.01)$$

where $\mathbf{X}\beta$ contains the fixed terms of the model, $\mathbf{Z}\mathbf{c}$ contains the random terms, $\boldsymbol{\varepsilon}$ is the vector of the residuals and $\boldsymbol{\xi}$ is a vector of spatially dependant residuals modelling

the local trends. However, unlike the SpATS model, $\mathbf{X}_s \boldsymbol{\beta}_s$ and $\mathbf{Z}_s \mathbf{c}_s$ do not model a smooth bivariate surface, but contain linear and one-dimensional cubic splines terms that account for the global variations in the data (see Verbyla *et al.* (1999) for details).

3.5.1 Variogram and $AR(1)$ process

As Gilmour *et al.* (1997) explain in their paper, they model an error process using a variogram. The error process is defined as the sum of the independent and spatially-dependent residuals: $e = \xi + \varepsilon$, where the term of interest is the spatially dependant error term ξ .

The variogram is a function that computes the evolution of the covariance of a spatially-dependent variable over the distance. It enables the visualization of the covariance structure of a variable as a function of spatial displacement. They exist both in one and two dimensions. An example of variograms is presented in figure 3.8. In our case, let us consider the error process e , being a stationary (direction-dependent) process over the rows (strips) u and the columns (positions) v . For two distinct points in space x and y , the covariance structure between these two points is modelled by the theoretical variogram (also called the semi-variogram) as:

$$\omega(\mathbf{x}, \mathbf{y}) = \frac{1}{2} \text{var}[e(\mathbf{x}) - e(\mathbf{y})] = \frac{1}{2}[V(\mathbf{x}, \mathbf{x}) + V(\mathbf{y}, \mathbf{y}) - 2V(\mathbf{x}, \mathbf{y})], \quad (3.5.11)$$

where V is the covariance function of our error process e . We can define the distance between the pair of points (x, y) as:

$$\mathbf{l} = \begin{bmatrix} l_1 \\ l_2 \end{bmatrix} = \begin{bmatrix} |x_1 - y_1| \\ |x_2 - y_2| \end{bmatrix},$$

then Zimmerman & Harville (1991) show that the covariance function of the error process can be rewritten a directional exponential covariance (DEC) structure:

$$V(\mathbf{x}, \mathbf{y}) = \exp(-\alpha_1 l_1 - \alpha_2 l_2). \quad (3.5.12)$$

For most field experiments, plots are arranged in regular grids and thus separated by equivalent distances. Therefore the distance between plots is expressed using the displacement vector, which takes values for l_1 of $0, d_1, 2d_1, \dots, (r-1)d_1$ and for l_2 of $0, d_2, 2d_2, \dots, (c-1)d_2$, where d_1 and d_2 are the plot dimensions. From that we can create an indexed displacement vector \mathbf{l}^* with values for l_1^* of $0, 1, 2, \dots, (r-1)$ and values for l_2^* of $0, 1, 2, \dots, (c-1)$. The semi-variogram can then be rewritten as:

$$\begin{aligned} \omega(\mathbf{l}^*) &= \sigma_\varepsilon^2 + \sigma_\xi^2 [1 - \exp(-\alpha_1 d_1 l_1^* - \alpha_2 d_2 l_2^*)] & \mathbf{l}^* \neq 0 \\ &= 0 & \mathbf{l}^* = 0 \end{aligned} \quad (3.5.13)$$

A first order auto-regressive process, is a process in which the term at time t only depends on the value of the process at time $(t-1)$ and on a stochastic term (white noise): $X_t = c + \rho X_{t-1} + \varepsilon_t$. The only parameter in the model (if the intercept is ignored)

is ρ . Therefore Cullis & Gleeson (1991) show that by setting $\rho_1 = \exp(-\alpha_1 d_1)$ and $\rho_2 = \exp(-\alpha_2 d_2)$, the variogram (equation 3.5.13) becomes

$$\begin{aligned}\omega(\mathbf{l}^*) &= \sigma_\varepsilon^2 + \sigma^2 \left(1 - \rho_1^{l_1^*} \rho_2^{l_2^*}\right) & \mathbf{l}^* \neq 0 \\ &= 0 & \mathbf{l}^* = 0\end{aligned}. \quad (3.5.14)$$

This formulation demonstrates the equivalence between the semi-variogram, the DEC model and the $AR(1) \times AR(1)$ process for field experiments. Figure 3.8 shows an example of a simulated variogram using an $AR(1) \times AR(1)$ process and actual variogram fitted using the same process.

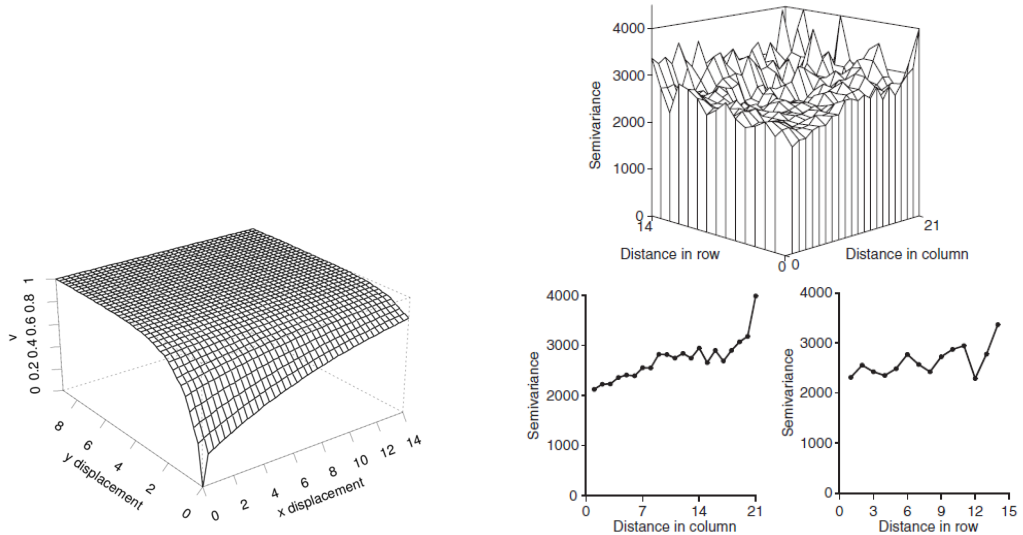


Figure 3.8: Examples of variogram using $AR(1) \times AR(1)$ processes. (Leftmost panel) Variogram for a standardized $AR(1) \times AR(1)$ process, with $\rho_x = 0.9$, $\rho_y = 0.4$ and $\sigma^2 = 0.3$ from Gilmour *et al.* (1997). (Rightmost panel) Variogram fitted to wheat data of Gilmour *et al.* (1997) with the one-dimensional associated variograms in the rows and columns direction (Piepho & E. Williams 2010).

3.5.2 Linear variance structure

The modification of the $AR(1) \times AR(1)$ model to include the linear variance (LV) structure, is based on the one-dimensional LV structure proposed by E. R. Williams (1986) and its elevation to two dimensions by Piepho & E. Williams (2010). Let us first consider the case of LV in one dimension (a single position over several strips in our case) for $n=99$ plots. The baseline covariance structure is written as:

$$V_0 = \sigma^2 I_n + \eta J_n, \quad (3.5.21)$$

where I_n is the n -dimensional identity matrix and J_n is a n by n matrix of ones everywhere, σ^2 is the residual variance and η is the block variance. E. R. Williams (1986) propose to elevate this baseline covariance model by a spatial component V_s such that plots within the same block obey a correlation structure that decays linearly with distance. For two plots x and y , they define the matrix L_n as a $n \times n$ matrix where

the (x, y) -th element is equal to $|x - y|$. From that statement, the decaying correlation structure can be expressed in matrix form as $V_S = \kappa \left(J_n - \tilde{\phi} L_n \right)$, where $\tilde{\phi}$ is a slope parameter determining the decaying process and κ is a spatial variance. To avoid confounding effects, E. R. Williams (1986) rewrite the complete, elevated model as a linear structure:

$$V = V_0 + V_S = \sigma^2 I_n + \eta J_n + \phi M_n, \quad (3.5.22)$$

where $M_n = (n - 1)J_n - L_n$ and $\phi = \kappa\tilde{\phi}$. The components of the model have the following interpretation: (i) $\sigma^2 I_n$ is the residual error (or nugget effect in variogram-specific terms); (ii) the block effect, ηJ_n , captures extraneous variations on the field; and (iii) ϕM_n models a spatial correlation within a block by a LV–covariance structure where correlation decays linearly with distance.

Piepho & E. Williams (2010) present two main ways to extend the model to two dimensions. The first ones is superimposition of two structure: a LV structure along the columns but independent between the rows and a LV structure along the columns but independent between the rows ($LV \otimes I$). The second one is a product of the two LV structure: $LV \otimes LV$. In a desire of clarity, the full notation of these structure is not presented here, but are detailed in the paper of Piepho & E. Williams (2010).

The similarity between the LV structure and the $AR(1)$ process comes from the fact that when the spatial correlation $\rho^l \approx 1$ then $\rho^{|x-y|} = \exp[\log(\rho)|x - y|] \approx 1 - \tilde{\phi}|x - y|$ with $\tilde{\phi} = -\log(\rho)$. Thus, when the correlation is close to one (which is quite common in practice (Pilarczyk 2007)) and when block sizes, hence distances $|x - y|$, are not large $AR(1)$ and LV models are expected to yield similar results. The main advantages of the LV model is that it is more robust to convergence issues when correlation is close to unity and also that it has one parameter instead of two, for the spatial component. To visualize more clearly the different spatial covariance structures of the different models, figure 3.9 shows examples of those structures.

3.5.3 Best standard spatial model

As explained in the previous section, several spatial covariance structures can be used in the standard spatial (SS) model to account for the spatial variations. Comparing the fit to the data of all the different models, would be a tedious and unnecessary process. Instead, all the SS models will be compared in term of AIC and deviance. Those two goodness-of-fit statistics are commonly used to compare statistical models (Gilmour *et al.* 1997; Piepho & E. Williams 2010; Velazco *et al.* 2017). Those two statistics are based on the value of the likelihood function, \mathcal{L} :

$$AIC = 2k - 2\ln(\mathcal{L}) \quad \text{and} \quad \text{Deviance} = -2\log(\mathcal{L}) + C. \quad (3.5.31)$$

The main advantage of the AIC is that it takes into account the number of parameters of the model k , to avoid overfitting. The best standard spatial model (BSS) was retained as the one with the lowest AIC and Deviance. This model was used with the data from the platform and its results were compared to the one from the SpATS model.

$$\sigma^2 \begin{pmatrix} 1 & \rho & \rho^2 & \rho^3 \\ \rho & 1 & \rho & \rho^2 \\ \rho^2 & \rho & 1 & \rho \\ \rho^3 & \rho^2 & \rho & 1 \end{pmatrix}$$

AR(1)

$$\sigma^2 \begin{pmatrix} 1 & \rho_r^{d(1,2,r)} \rho_c^{d(1,2,c)} & \rho_r^{d(1,3,r)} \rho_c^{d(1,3,c)} & \rho_r^{d(1,4,r)} \rho_c^{d(1,4,c)} \\ \rho_r^{d(2,1,r)} \rho_c^{d(2,1,c)} & 1 & \rho_r^{d(2,3,r)} \rho_c^{d(2,3,c)} & \rho_r^{d(2,4,r)} \rho_c^{d(2,4,c)} \\ \rho_r^{d(3,1,r)} \rho_c^{d(3,1,c)} & \rho_r^{d(3,2,r)} \rho_c^{d(3,2,c)} & 1 & \rho_r^{d(3,4,r)} \rho_c^{d(3,4,c)} \\ \rho_r^{d(4,1,r)} \rho_c^{d(4,1,c)} & \rho_r^{d(4,2,r)} \rho_c^{d(4,2,c)} & \rho_r^{d(4,3,r)} \rho_c^{d(4,3,c)} & 1 \end{pmatrix}$$

AR(1) × AR(1)

$$\sigma^2 \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} + \eta \begin{pmatrix} 1 & 1 & 1 & 1 \\ 1 & 1 & 1 & 1 \\ 1 & 1 & 1 & 1 \\ 1 & 1 & 1 & 1 \end{pmatrix} +$$

$$(n-1)\phi \begin{pmatrix} 1 & 1-d(1,2) & 1-d(1,3) & 1-d(1,4) \\ 1-d(2,1) & 1 & 1-d(2,3) & 1-d(2,4) \\ 1-d(3,1) & 1-d(3,2) & 1 & 1-d(3,4) \\ 1-d(4,1) & 1-d(4,2) & 1-d(4,3) & 1 \end{pmatrix}$$

LV

Figure 3.9: Spatial covariance structures of the standard spatial models. The $AR(1) \times AR(1)$ model is presented as anisotropic. In isotropic conditions, the correlation value would be the same along the rows and the columns ($\rho_c = \rho_r$). The distance is represented in units, as the indexed displacement.

3.5.4 Model estimation

The models were implemented using the `proc mixed` of the SAS software (SAS Institute, Cary, NC, USA). The spatial covariance structure was specified in the `type` option of the `random` statement of the procedure. The $AR(1)$ models were fitted using `type = AR(1)`, the $AR(1) \times AR(1)$ were fitted using `type = sp(POWA)` and the LV models were fitted using `type = lin(q)`.

3.6 Model comparison

The SpATS model was compared with the BSS model in terms of quality of fit to the data and ability to detect genotype differences. Similar parameters to those used by Velazco *et al.* (2017), were considered for the comparison:

- Genetic variance (σ_g^2): precision of the genotype effect estimation. The generalized heritability will also be used to assess the same parameter.
- Spatially independent residual variance (σ_e^2): precision of the overall fit to the data and ability to capture the spatial trends.
- BLUPs of the genotypes: similarity between the estimates of the effect of each genotype for the two models.

It is interesting to note that Rodriguez-Alvarez *et al.* (2018) also use the Pearson correlations of predicted genotypes values between environments (i.e. field trials) as a way of comparing models. Since only one trial was studied in this thesis, this correlation cannot be used.

Chapter 4

Results and discussion

4.1 Descriptive statistics

Before the experiment, the seeds were weighted in groups of ten, to see if there was a baseline difference between certain genotypes. Table 3.1, in the previous section, displays the measured weights. After the completion of the experiment, the outliers were identified, and each plant was attributed a specific weight, following the protocol described in material and methods section. Those weights were used to compute the weighted mean and weighted standard deviation of the fresh and dry weight of the root system and the leaf system for each plant.

Because of germination problems on the platform and inside the germination chamber, not all genotypes were similarly represented in the experiment. Table ?? presents the effective germination rates for each genotype, i.e. the number of seed actually kept for the spatial analysis over the number of seeds placed on the platform. This table is interesting because germination rate is a genotypic feature. Since the genotypes are sorted by their effective germination rate, we see clear discrepancies between genotypes, indicating that some may not be well suited to aeroponic growth. Only 6 genotypes have a germination rate lower than 50% (below the dashed line on table ??, even though more than 15 seeds where placed on the platform. Thus, we expect to see a lower yield in all variables for these genotypes.

To verify those assumptions, we created boxplots representing those descriptive statistics as well as the data point are presented in figure 4.1. The numerical values of these results are presented in table B.1, in appendix B.2. Even though no clear conclusions can be made from these figures, we can see large variation of mean values between genotypes. Also, for some genotypes, the difference between tanks seem significant (e.g. genotype 15 on figure ??) while it's clearly not the case for some other (e.g. genotype 7 on figure ??). This implies that the tank and genotypes effects are significant. Overall the values of dry weights seem to have less variations than the fresh weights.

Table 4.1: Effective on-platform germination rates (GR) with the number of seeds kept for data analysis (NS kept) and the number of seeds actually placed on the platform (NS placed) for each genotype.

Genotype	NS placed	NS kept	GR
25	29	27	93.1
23	22	20	90.9
16	27	23	85.2
18	26	22	84.6
3	29	24	82.8
17	27	21	77.8
19	30	23	76.7
1	24	18	75.0
12	28	21	75.0
14	26	19	73.1
28	26	19	73.1
9	29	20	69.0
6	29	19	65.5
7	29	19	65.5
4	23	15	65.2
27	29	18	62.1
21	26	16	61.5
5	30	18	60.0
24	30	18	60.0
10	29	17	58.6
20	26	15	57.7
22	18	10	55.6
29	28	15	53.6
13	27	14	51.9
15	18	9	50.0
2	26	12	46.2
11	19	8	42.1
26	29	12	41.4
8	21	8	38.1
30	23	3	13.0
31	127	0	0.0
	80	0	0.0

CHAPTER 4. RESULTS AND DISCUSSION

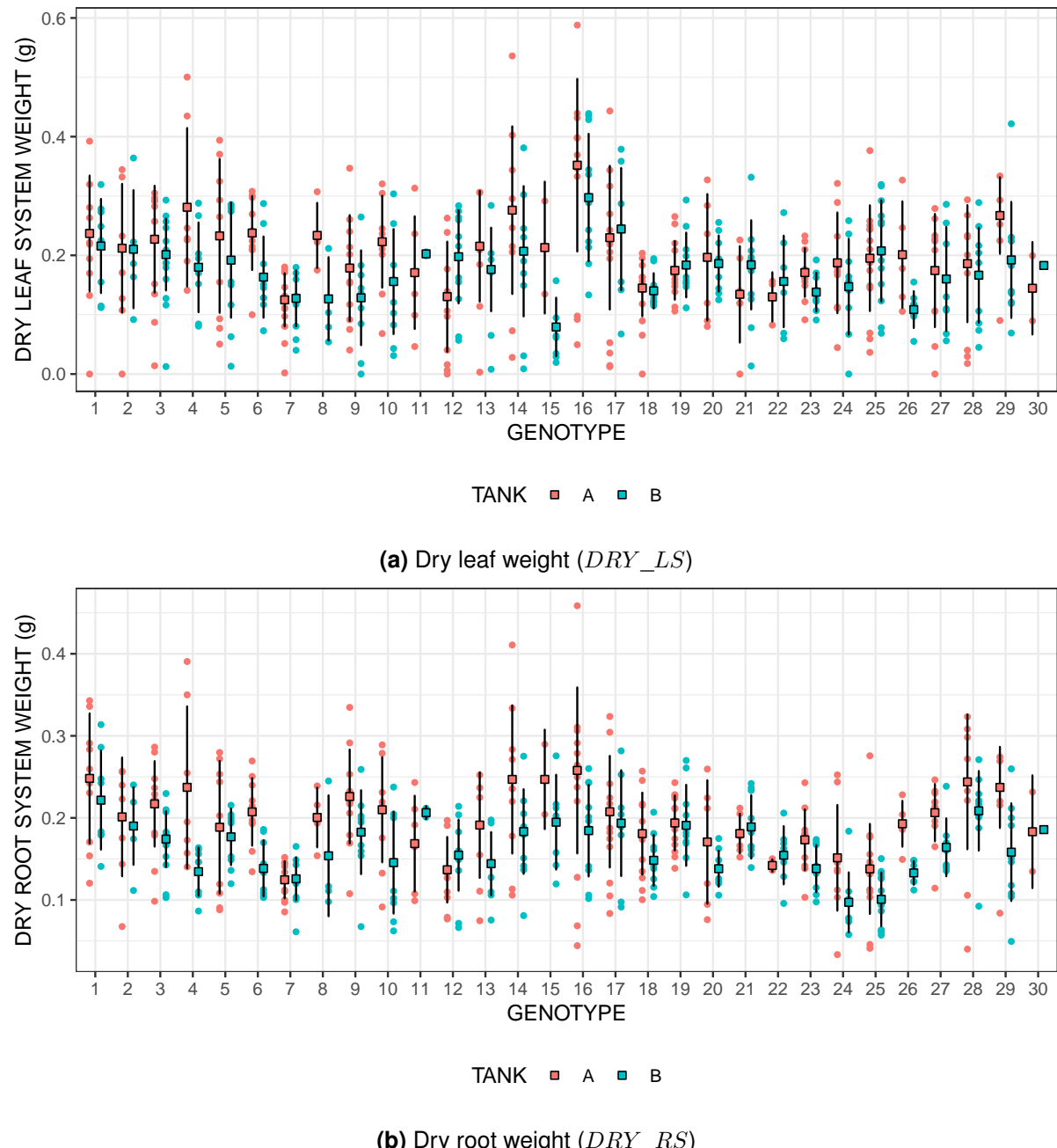


Figure 4.1: Dotplot displaying mean weight (□) and associated standard deviation (—), grouped by tanks for each variable.

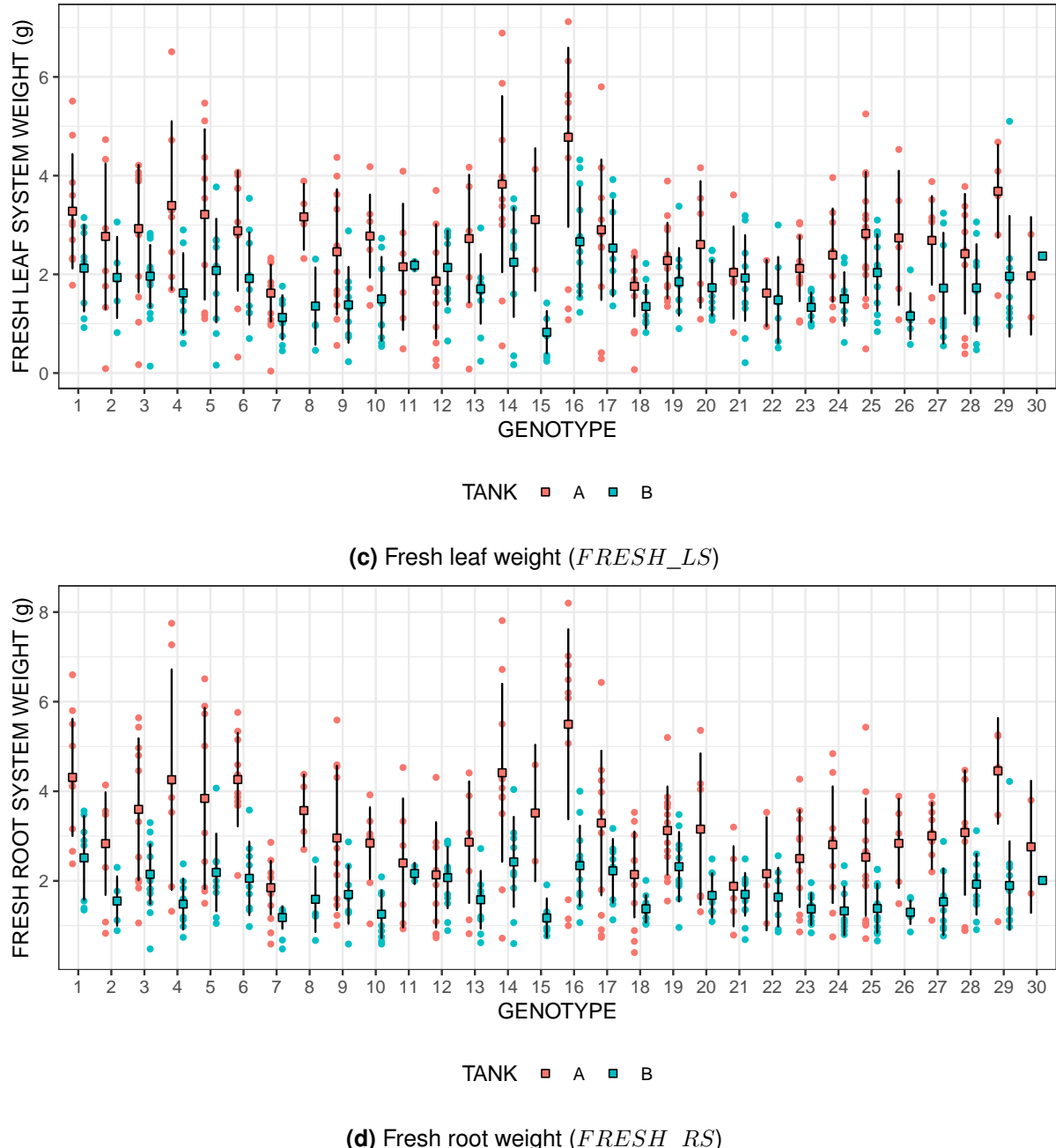


Figure 4.1: Dotplot displaying mean weight (□) and associated standard deviation (—), grouped by tanks for each variable.

	DRY_LS	DRY_RS	FRESH_RS	FRESH_LS
DRY_LS	1.00	0.70	0.84	0.93
DRY_RS	0.70	1.00	0.76	0.68
FRESH_RS	0.84	0.76	1.00	0.93
FRESH_LS	0.93	0.68	0.93	1.00

4.2 SpATS analysis

The SpATS model usually takes rows and columns coordinates as inputs for spatial position. Given that we have tanks (A and B), strips (from 1 to 99) and positions (from 1 to 5), we reshaped the data to give have the tank side by side and the 99 strips divided in two columns (to have a similar disposition to the one in the greenhouse). Figure 4.2 shows the reshaping of the positions. This new display of the data allows us to see the difference between tanks more clearly and to visualize the variables' values as they were in the greenhouse.

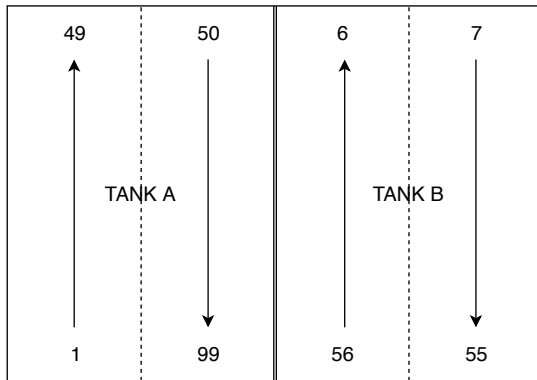


Figure 4.2: Reshaping of the data table to fit the disposition of the greenhouse. The number indicates the original strip number.

The model was then fitted for the four weight variables, using the settings specified in the previous chapter. Table 4.1 presents the effective dimensions associated with the bivariate smooth surface components (see equation 3.4.03) and their relative contribution to the fitted surface for each variable. We see that the fresh weights exhibited a higher complexity in the structure of the spatial surface. This is reflected by the higher value of contribution for the smooth-by-smooth term ($f_{u,v}(u, v)$), that accounts for almost 70% in both variables. Besides this term, the main sources of variation are the linear (for the strips) by smooth (for the positions) term and the smooth trend along the strips term. It is not surprising to have more variation along the strips than along the positions given that there were 99 strips but only 5 positions. Concerning the dry weights, the variation is more spread between all the components for both variables. This means that the variation in the data can be more easily attributed to the strips and positions of the plants. However, all the variables have components with zero value of ED_s ¹, indicating that these terms were not necessary to model the spatial surface.

¹the actual values were not zero but it is denoted as such since they were inferior to 1×10^{-15}

Table 4.2: Model dimensions and effective dimensions (and percentage of the total of the spatial components) of each spatial components for all variables. ED_ϵ represents the effective dimensions for the residuals; ED_g , is the effective dimensions for the genotype and H_g^2 is the heritability. Here \mathbf{v} represents the columns, i.e. the position on the strip; and \mathbf{u} represents the rows, i.e. the strip itself.

Model components	Model	FRESH_LS	FRESH_RS	DRY_LS	DRY_RS
$f_v(\mathbf{v})$	6	0 (0,00%)	0 (0,00%)	0 (0,00%)	0 (0,00%)
$f_u(\mathbf{u})$	100	0,8 (8,73%)	2,26 (14,10%)	1,02 (17,26%)	1,8 (17,26%)
$\mathbf{u} \odot h_v(\mathbf{v})$	6	1,84 (19,95%)	2,63 (16,40%)	1,47 (24,77%)	2,54 (24,77%)
$\mathbf{v} \odot h_u(\mathbf{u})$	100	0,24 (2,56%)	0 (0,00%)	1,09 (18,36%)	0,49 (18,36%)
$f_{u,v}(\mathbf{u}, \mathbf{v})$	150	6,33 (68,76%)	11,16 (69,50%)	2,35 (39,62%)	0,08 (39,62%)
Total	362	9,21 (100%)	16,15 (100%)	5,93 (100%)	4,91 (100%)
ED_ϵ		466,6	459,3	470,2	470,1
ED_g	30	21,02	22,58	21,38	22,98
H_g^2		0,72	0,78	0,74	0,79

Table 4.1 also presents the effective dimension of the genetic component (ED_g) which is quite similar across variables. This is reflected in the heritability values which are all around 75%. However, it should be noted that the SpATS model tends to overestimate the heritability (M. X. Rodríguez-Álvarez, Boer, *et al.* 2016). This means that a good part of the phenotypic variation can be attributed to genotypes.

Figure 4.3 shows the raw data (a), a graphical representation of the fitted spatial trend ($f(\mathbf{u}, \mathbf{v})$) (b) and the spatially independent residuals ϵ (c) obtained from the SpATS package. While there are a lot of missing data, some spatial trends still stand out. Just as predicted with the dotplot in the descriptive statistics section, weights in the B tank are lower than in the A tank. This is especially visible for the fresh weight, where the total weight range is greater than for the dry weights.

The inspection of the fitted spatial surfaces shows us that the trends have been captured by the SpATS model. An additional analysis of the residuals suggests that the spatial patterns have effectively been removed in all four variables by the two-dimensional spline surface. However, some high data points still persist in the residuals, this is mainly due to the high variability of the weights in the raw data. In order to fully analyse the residuals, two additional diagnosis plots have been created: a lagplot to test for spatial independence and a normal distribution to test for the normality assumption. These plots are presented in appendix B.1. Overall, the residuals seem to be independent and normally distributed, which confirms again that the spatial pattern has been well-captured by the model. Another interesting tool to evaluate the spatial independence is the variogram, presented in the previous chapter (section 3.5). As said previously, if the model for spatial trend fits well, the variogram should be a horizontal plane (Piepho & E. Williams 2010).

Table 4.3: Individual variances of all the components of the SpATS model.

	FRESH_LS	FRESH_RS	DRY_LS	DRY_RS
c_g	0.171	0.262	1.27×10^{-3}	6.9×10^{-4}
c_v	3.24×10^{-3}	5.08×10^{-3}	1.95×10^{-7}	8.47×10^{-17}
c_u	1.86×10^{-4}	2.25×10^{-5}	2.42×10^{-7}	3.93×10^{-8}
$f_v(\mathbf{v})$	2.2	11	1.06×10^{-4}	0.265
$f_u(\mathbf{u})$	2.75×10^{-5}	5.36×10^{-5}	5.4×10^{-8}	1.55×10^{-8}
$\mathbf{u} \odot h_v(\mathbf{v})$	5.60×10^{-52}	1.76×10^{-38}	1.51×10^{-52}	1.44×10^{-13}
$\mathbf{v} \odot h_u(\mathbf{u})$	1.81×10^{-9}	4.83×10^{-4}	6.75×10^{-10}	1.43×10^{-6}
$f_{u,v}(\mathbf{u}, \mathbf{v})$	0.398	0.252	6.13×10^{-5}	7.13×10^{-4}
ϵ	4.707	5.014	0.03364	0.01219

Finally, another interesting result from figure 4.3 is the comparison between the scales of spatial variations and residual variations, because they provide an idea of the relative importance of field trends for each variable. We see that for all variables, the scales of the residuals are about ten-fold the scales of the fitted surfaces. This is explained by the effective dimension of the residuals presented in table 4.1. We see that, for all variables, those dimensions are much higher than the others. This means that the spatial variation was lower than the random variation, and even though spatial patterns were captured, much of the variation is still present in the residuals of the models. Given that the raw data present a high variability and a lot of missing values, we expected to have residuals with high variance (see table 4.2 for the variances linked to each component of the model).

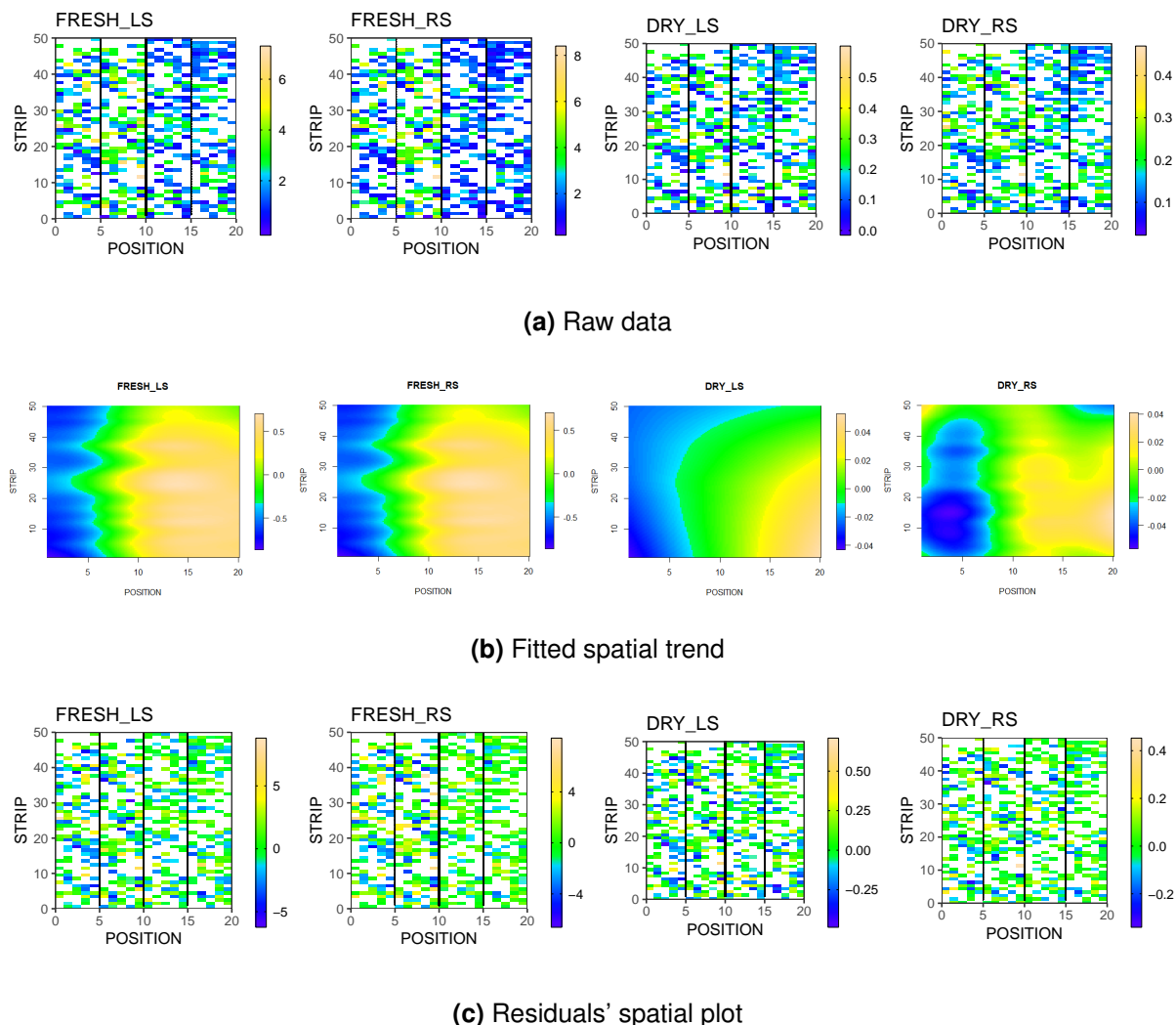


Figure 4.3: Raw data, fitted spatial trend and residuals' plot for each variable.

4.3 Standard spatial model analysis

For the standard spatial model analysis, a baseline model, only considering a fixed effect for the tanks and random effects the rows, columns and the genotypes, was fitted for each of the four variables considered. Each model was then augmented by adding linear regression terms on the rows and columns and one of the following covariance structure:

- $AR(1)$ process along the rows or columns
- $AR(1) \times AR(1)$ process
- LV process along the rows or columns
- Superimposed row and column structure $LV + LV$
- Separable process along the rows or columns $LV \otimes J$
- Separable process along the rows and columns $LV \times LV$

At each step, the AIC and the Deviance were computed and the best model was selected using this criterion (a lower value is preferred). Table 4.3 gives the structure of the final selected model for each variable. ...INTERPRETATION OF THOSE MODELS (similar or not ?).

Then the variograms of the residuals were analysed to see if any leftover spatial pattern was still there. Figure ?? presents those variograms, ..INTERPRETATION.. + ANALYSIS of the residuals in the same way as the SpATS model.

Table 4.4: Best standard spatial (BSS) model selected for each of the four variables. All the models contain an intercept and a fixed effect for the tank and a random effect for the genotypes. P represents a random effect for the positions (columns), S a random effect for the strips (rows) and n represent the spatially independent residuals.

Variable	BSS
FRESH_LS	$S + P + AR(1) \times AR(1) + n$
FRESH_RS	$S + P + AR(1) \times AR(1) + n$
DRY_LS	$S + LV \times LV + n$
DRY_RS	$S + LV + LV + n$

Table 4.4 gives the value of the variance and correlation components for each spatial model.

Table 4.5: Add caption

Variable	ρ_s	ρ_p	σ_g^2	σ_ξ^2	σ_ε^2
FRESH_LS					
FRESH_RS					
DRY_LS					
DRY_RS					

4.4 Model comparison

Table 4.6: Comparison of the estimated TANK fixed effect for both models.

	FRESH_LS	FRESH_RS	DRY_LS	DRY_RS
SpATS	2,9527	3,7282	0,2164	0,2223
BSS	2,7016	3,2975	0,2075	0,1959
Δ	0,2511	0,4307	0,0089	0,0264
SpATS	1,3343	1,1445	0,1622	0,1523
BSS	1,3056	1,1185	0,1660	0,1606
Δ	0,0287	0,0260	0,0038	0,0083

Table 4.7: Comparison of both models in term of genetic variance and residual variance. Δ represents the absolute difference between the two variances.

	FRESH_LS	FRESH_RS	DRY_LS	DRY_RS
SpATS	0,1709	0,2618	1,267E-03	6,902E-04
BSS	0,1829	0,2644	1,339E-03	7,350E-04
Δ	0,0120	0,0026	7,212E-05	4,484E-05
SpATS	4,7073	5,0137	3,364E-02	1,219E-02
BSS	4,5602	4,9245	3,321E-02	1,280E-02
Δ	0,1470	0,0892	4,297E-04	6,147E-04

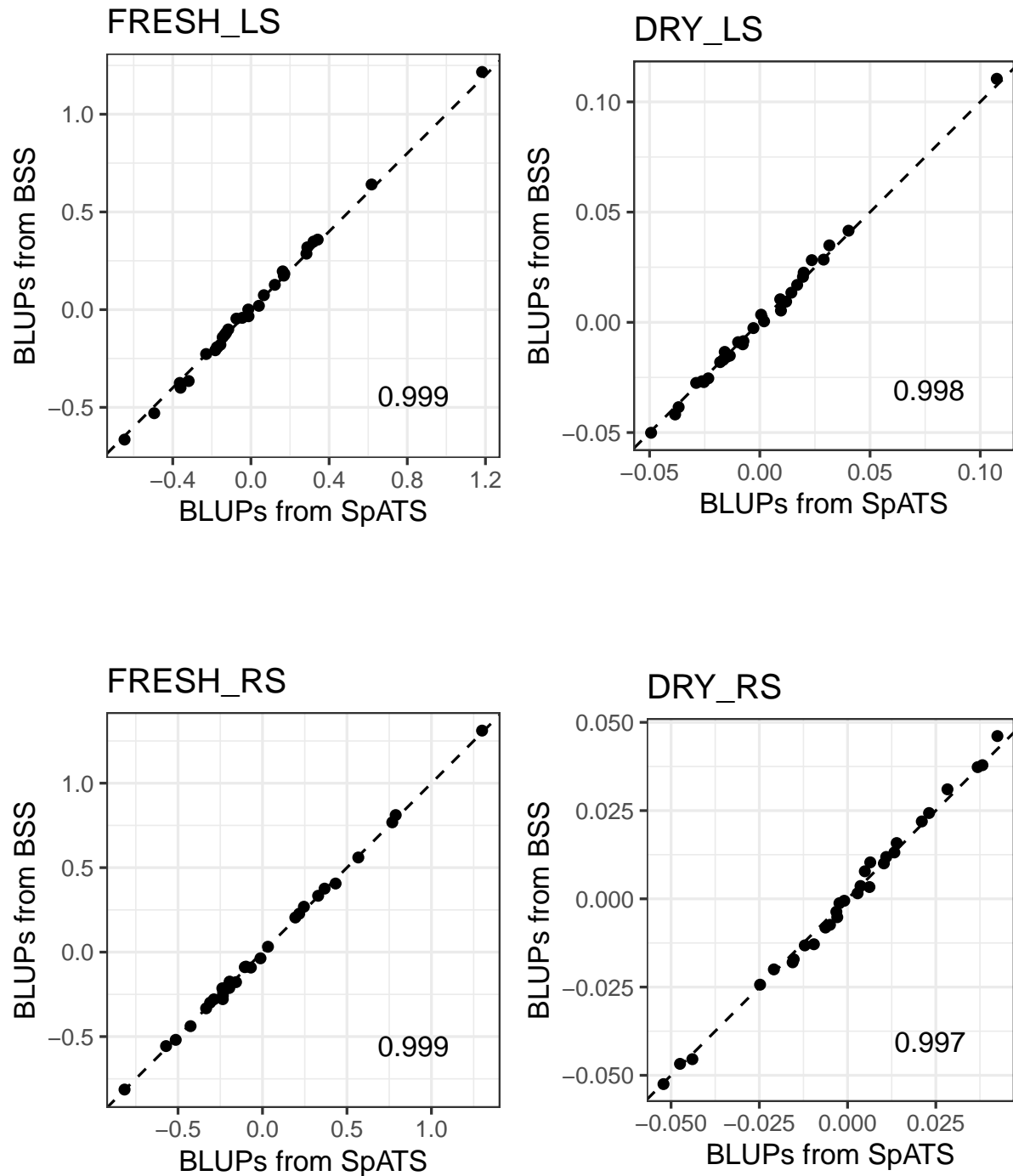


Figure 4.4: Comparison of the genotype BLUPs from the SpATS model and the BSS model, with the spearman's rank correlation (bottom right corner of each panel).

Chapter 5

Conclusion

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Appendices

Appendix A

Additional informations on computation

A.1 Element-wise product

The element-wise product between two matrix \mathbf{A} and \mathbf{B} is noted $\mathbf{A} \odot \mathbf{B}$ and is defined in the following way:

For two matrices \mathbf{A}, \mathbf{B} of same dimensions $n \times m$, the element-wise product is a $n \times m$ matrix where the elements are defined by:

$$(\mathbf{A} \odot \mathbf{B})_{i,j} = (\mathbf{A})_{i,j} \cdot (\mathbf{B})_{i,j}$$

The product is undefined for matrices of different dimensions

A.2 Kronecker product

The Kronecker product of two matrix \mathbf{A} and \mathbf{B} of respective dimensions $n \times m$ and $p \times q$ is a $np \times mq$ block matrix where the elements are defined by:

$$\mathbf{A} \otimes \mathbf{B} = \begin{bmatrix} a_{11}\mathbf{B} & \cdots & a_{1n}\mathbf{B} \\ \vdots & \ddots & \vdots \\ a_{m1}\mathbf{B} & \cdots & a_{mn}\mathbf{B} \end{bmatrix}$$

A.3 Polynomials splines

Fahrmeir *et al.* (2013) state that a function $f : [a, b] \rightarrow \mathbb{R}$ is called a polynomial spline of degree $l \geq 0$ with knots $a = \kappa_1 < \dots < \kappa_m = b$, if it fulfills the following conditions:

1. $f(z)$ is $(l - 1)$ times continuously differentiable. The special case of $l = 1$ corresponds to $f(z)$ being continuous (but not differentiable). We do not state any smoothness requirements for $f(z)$ when $l = 0$.
2. $f(z)$ is a polynomial of degree l on intervals $[\kappa_j, \kappa_{j+1}]$ defined by the knots.

Moreover, it can be shown that each polynomial spline of degree l with knots $\kappa_1 < \dots < \kappa_m$ can be uniquely determined as a linear combination of the $d = l + m - 1$ functions B_1, \dots, B_d , called the *basis functions*, since we can uniquely represent all polynomials splines by using these functions.

A.3.1 B-splines

B-splines are polynomial splines with specific basis functions. B-spline basis functions are constructed from piecewise polynomials that are fused smoothly at the knots to achieve the desired smoothness constraints. More specifically, a B-spline basis function consists of $(l + 1)$ polynomial pieces of degree l , which are joined in an $(l - 1)$ continuously differentiable way. All B-spline basis functions are set up based on a given knot configuration. Using the complete basis, the function $f(z)$ can again be represented through a linear combination of $d = m + l - 1$ basis functions, i.e.,

$$f(z) = \sum_{j=1}^d \gamma_j B_j(z).$$

The B-splines of order $l = 0$ can be written as

$$B_j^0(z) = \begin{cases} 1 & \kappa_j \leq z < \kappa_{j+1} \\ 0 & \text{otherwise} \end{cases} \quad j = 1, \dots, d - 1$$

and the B-splines for higher order l can be written as

$$B_j^l(z) = \frac{z - \kappa_{j-l}}{\kappa_j - \kappa_{j-l}} B_{j-1}^{l-1}(z) + \frac{\kappa_{j+1} - z}{\kappa_{j+1} - \kappa_{j+1-l}} B_j^{l-1}(z).$$

The estimation of a polynomial spline in B-spline representation can be traced back to the estimation of a linear model with a large number of parameters and design matrix

$$\mathbf{Z} = \begin{pmatrix} B_1^l(z_1) & \dots & B_d^l(z_1) \\ \vdots & & \vdots \\ B_1^l(z_n) & \dots & B_d^l(z_n) \end{pmatrix}.$$

The linear combination of basis functions can then be written in matrix form

$$\mathbf{y} = \mathbf{Z}\boldsymbol{\gamma}$$

where the coefficient matrix, $\boldsymbol{\gamma}$ can be estimated using least squares.

The estimation of a B-spline fit can be summarized in three steps:

1. We calculate a complete B-spline basis for a given number of knots.
2. The least squares estimate $\hat{\boldsymbol{\gamma}}$ yields an amplitude $\hat{\gamma}_j$ for the scaling of every basis function.
3. We obtain the final estimate by summing the scaled basis function.

A.3.2 Penalized splines

We clearly see that the quality of the estimation by polynomials splines highly depends on the number of knots and that this can easily lead to an over-fitting issue. To overcome this problem, *penalized splines (P-splines)* introduce a roughness penalty term that prevents over-fitting and minimize a *penalized least squares (PLS) criterion* instead of the usual least squares criterion.

To characterize the smoothness of any type of function, the use of (squared) derivatives is appropriate, since these represent measures for the variability of a function. Therefore penalties based on the second derivative, such as

$$\lambda \int (f''(z))^2 dz,$$

are particularly attractive since they measure the curvature of a function. Since we know that the first derivative of a B-spline can be written as a function of the first differences of the corresponding coefficient vector, we can use differences of a higher order r if we aim at a smooth function in terms of r th-order derivatives. This leads to the penalized residual sum of squares

$$PLS(\lambda) = \sum_{i=1}^n \left(y_i - \sum_{j=1}^d \gamma_j B_j(z_i) \right)^2 + \lambda \sum_{j=r+1}^d (\Delta^r \gamma_j)^2,$$

where Δ^r denotes the r th-order differences. The smoothing parameter $\lambda \geq 0$ controls the compromise between fidelity to the data and smoothness of the resulting function estimate. The PLS criterion can be rewritten using matrix notation

$$PLS(\lambda) = (\mathbf{y} - \mathbf{Z}\boldsymbol{\gamma})'(\mathbf{y} - \mathbf{Z}\boldsymbol{\gamma}) + \lambda \boldsymbol{\gamma}' \mathbf{K}_r \boldsymbol{\gamma}$$

where \mathbf{K}_r is the r th-order difference penalty matrix, and can be decomposed as $\mathbf{D}_r/\mathbf{D}_r'$ with \mathbf{D}_r the r th-order difference matrix. The smoothing parameter $\lambda \geq 0$ controls the compromise between fidelity to the data and smoothness of the resulting function estimate. The PLS estimate of the coefficient matrix is then

$$\hat{\boldsymbol{\gamma}} = (\mathbf{Z}'\mathbf{Z} + \lambda \mathbf{K})^{-1} \mathbf{Z}'\mathbf{y}.$$

For more detailed information about polynomials splines, please refer to Fahrmeir *et al.* (2013) and P. H. C. Eilers & Marx (1996)

A.4 Penalized form of the solution

Let us consider the following model, representing the bivariate surface:

$$\hat{\boldsymbol{\alpha}} = (\mathbf{B}'\mathbf{B})^{-1} \mathbf{B}'\mathbf{y} \tag{A.4.01}$$

Since the model is purely parametric, it can be estimated by minimizing the residual sum of squares (with explicit solution $\hat{\boldsymbol{\alpha}} = (\mathbf{B}'\mathbf{B})^{-1} \mathbf{B}'\mathbf{y}$). To prevent over-fitting, P. H. C. Eilers & Marx (1996) propose to incorporate a discrete penalty on the coefficient associated to adjacent B-splines. For the two-dimensional case, the vector $\boldsymbol{\alpha}$ can be seen

as an $(L \times P)$ matrix of coefficients, $\mathbf{A} = [\alpha_{lp}]$. Now the rows and columns of \mathbf{A} correspond to the regression coefficients in the v and u direction, respectively. In anisotropic (direction-dependant) P-splines, a different amount of smoothing is assumed along the u and v directions. It leads to two penalties: one on all rows of \mathbf{A} , the other on all of its columns; and the penalized least squares objective function becomes (P. H. C. Eilers & Marx 2003)

$$\begin{aligned}
S^* = & \underbrace{\|\mathbf{y} - \mathbf{B}\boldsymbol{\alpha}\|^2}_{\text{Original objective function}} \\
& + \underbrace{\hat{\lambda} \|\hat{\mathbf{D}}\mathbf{A}\|_F^2}_{\text{Penalty along the columns}} \\
& + \underbrace{\check{\lambda} \|\mathbf{A}\check{\mathbf{D}}^t\|_F^2}_{\text{Penalty along the rows}} \\
= & \|\mathbf{y} - \mathbf{B}\boldsymbol{\alpha}\|^2 + \boldsymbol{\alpha}^t \mathbf{P} \boldsymbol{\alpha},
\end{aligned} \tag{A.4.02}$$

where $\mathbf{P} = \hat{\lambda} (\mathbf{I}_P \otimes \hat{\mathbf{D}}^t \hat{\mathbf{D}}) + \check{\lambda} (\check{\mathbf{D}}^t \check{\mathbf{D}} \otimes \mathbf{I}_L)$ is the penalty matrix, $\hat{\lambda}$ and $\check{\lambda}$ are the smoothing parameters acting, respectively, on the columns and rows of \mathbf{A} , and $\hat{\mathbf{D}}$ and $\check{\mathbf{D}}$ are the matrices that form differences of order d_u and d_v respectively. The minimizer of A.4.02 then becomes

$$\hat{\boldsymbol{\alpha}} = (\mathbf{B}^t \mathbf{B} + \mathbf{P})^{-1} \mathbf{B}^t \mathbf{y}. \tag{A.4.03}$$

Appendix B

Additional figures and tables

B.1 Graphical analysis of the residuals

B.1. GRAPHICAL ANALYSIS OF THE RESIDUALS

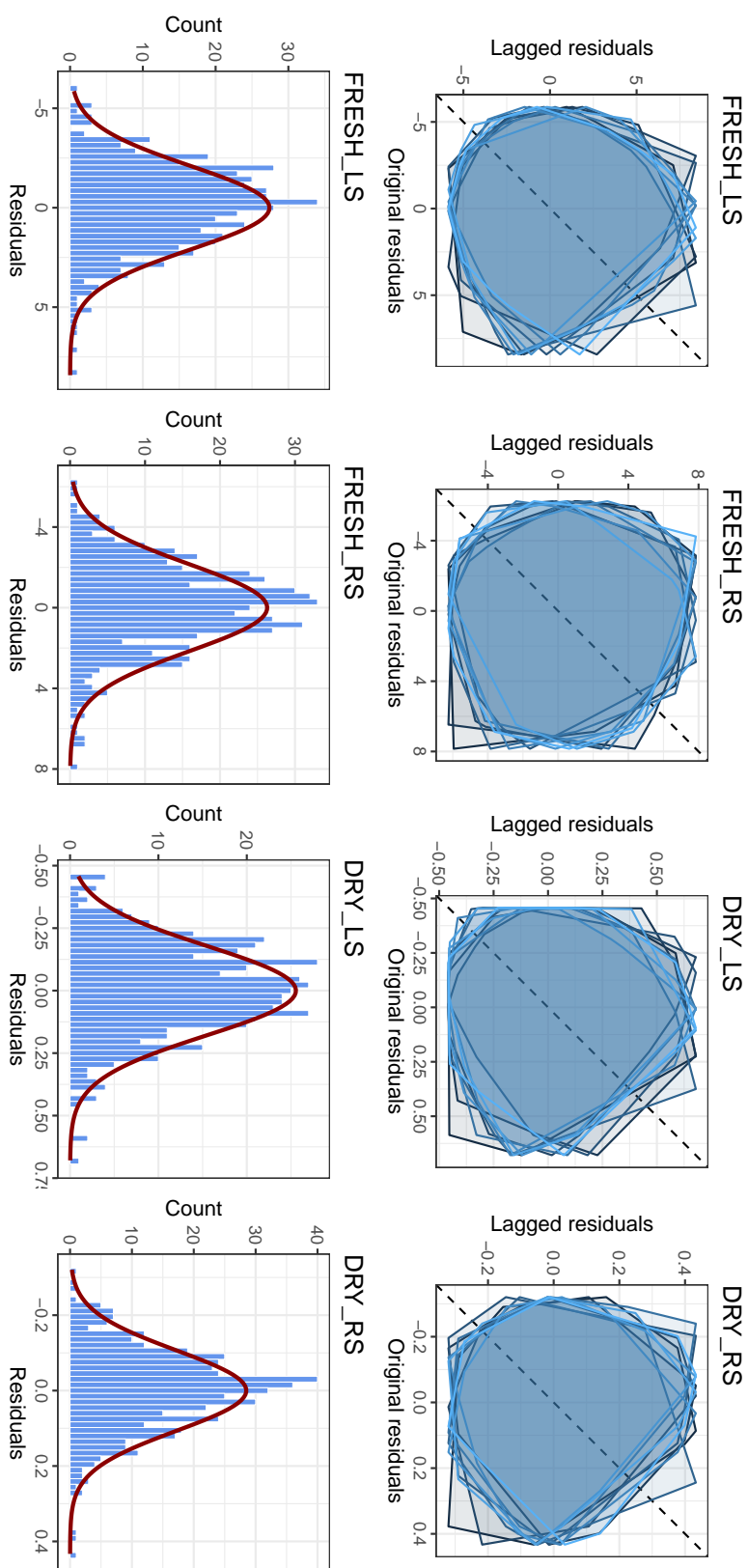


Figure B.1: Graphical analysis of the residuals of the SpATS model for each variable: a convexhull of the lagplots of the residuals with different levels of lag (first row) and an histogram with a zero-mean, and same variance, normal distribution (second row).

B.2 Descriptive statistics

Table B.1: Weighted mean and standard deviation for each genotype. DRY_{LS} represents the dry weight of the leaf system; DRY_{RS} , the dry weight for the root system; $FRESH_{LS}$, the fresh weight for the leaf system and $FRESH_{RS}$, the fresh weight for the root system. All the results are presented as mean \pm standard deviation (g)

Genotype	DRY_{LS}	DRY_{RS}	$FRESH_{LS}$	$FRESH_{RS}$
1	0.2267 ± 0.0869	0.2354 ± 0.0698	2.7231 ± 1.1612	3.4447 ± 1.4431
2	0.2113 ± 0.0993	0.1964 ± 0.06	2.4058 ± 1.254	2.2725 ± 1.1119
3	0.2132 ± 0.0747	0.1939 ± 0.0474	2.406 ± 1.0814	2.8146 ± 1.3663
4	0.227 ± 0.1148	0.1824 ± 0.0861	2.45 ± 1.5508	2.7773 ± 2.1926
5	0.2126 ± 0.113	0.1829 ± 0.0606	2.6521 ± 1.5044	3.0241 ± 1.7336
6	0.2024 ± 0.0739	0.1747 ± 0.051	2.4244 ± 1.1691	3.2168 ± 1.4545
7	0.126 ± 0.0441	0.1251 ± 0.0232	1.4118 ± 0.5677	1.5669 ± 0.5783
8	0.186 ± 0.0805	0.1798 ± 0.0567	2.3614 ± 1.1696	2.6894 ± 1.273
9	0.1559 ± 0.0865	0.2064 ± 0.0574	1.9704 ± 1.1782	2.3862 ± 1.3949
10	0.1885 ± 0.0875	0.1769 ± 0.0693	2.1228 ± 1.046	2.029 ± 1.0451
11	0.1789 ± 0.0811	0.1783 ± 0.0521	2.1608 ± 1.0747	2.339 ± 1.2172
12	0.1684 ± 0.0893	0.1469 ± 0.0417	2.0166 ± 0.9281	2.0985 ± 0.9015
13	0.1927 ± 0.0794	0.1639 ± 0.054	2.1326 ± 1.077	2.1176 ± 1.1582
14	0.2438 ± 0.1288	0.2173 ± 0.0796	3.0901 ± 1.6735	3.486 ± 1.8646
15	0.1175 ± 0.0885	0.2097 ± 0.0591	1.48 ± 1.2962	1.8417 ± 1.3372
16	0.3244 ± 0.1276	0.2208 ± 0.0879	3.7094 ± 1.8196	3.9028 ± 2.2598
17	0.2357 ± 0.1111	0.2019 ± 0.065	2.7519 ± 1.2418	2.8555 ± 1.4
18	0.1427 ± 0.039	0.1653 ± 0.0442	1.5621 ± 0.5627	1.7781 ± 0.8075
19	0.1785 ± 0.0505	0.1926 ± 0.04	2.0917 ± 0.7454	2.7726 ± 0.9668
20	0.1901 ± 0.0721	0.1506 ± 0.0497	2.0641 ± 0.9693	2.2429 ± 1.2908
21	0.1649 ± 0.0786	0.1859 ± 0.0329	1.9682 ± 0.8601	1.7689 ± 0.6407
22	0.1482 ± 0.0669	0.1508 ± 0.0296	1.522 ± 0.7799	1.793 ± 0.838
23	0.156 ± 0.04	0.1574 ± 0.0372	1.7621 ± 0.6537	1.9874 ± 0.9962
24	0.1668 ± 0.0827	0.1236 ± 0.0575	1.9357 ± 0.8693	2.0506 ± 1.2218
25	0.2013 ± 0.0856	0.1195 ± 0.0481	2.4338 ± 1.0937	1.9605 ± 1.1428
26	0.1463 ± 0.075	0.1573 ± 0.0365	1.7995 ± 1.1967	1.9256 ± 1.0053
27	0.1682 ± 0.0894	0.188 ± 0.0401	2.2662 ± 1.0855	2.3647 ± 1.0382
28	0.1753 ± 0.0865	0.2244 ± 0.066	2.0357 ± 1.0699	2.441 ± 1.1785
29	0.215 ± 0.0934	0.1823 ± 0.0663	2.4861 ± 1.3776	2.6743 ± 1.5787
30	0.1573 ± 0.0592	0.184 ± 0.0485	2.1033 ± 0.8712	2.51 ± 1.1265

Appendix C

Hoagland solution

Table C.1: Composition of the *Hoagland* nutritive solution. The pH must be adjusted to 5.0 using HCl 1% before using.

Components	Concentration (g/L)	ml for 25L of solution ¹
2M KNO ₃	202	62.5
2M Ca(NO ₃) ₂ x 4H ₂ O	472	62.5
2M MgSO ₄ x 7H ₂ O	493	25
1M NH ₄ NO ₃	80	25
Minors:		
H ₃ BO ₃	2.86	
MnCl ₂ x 4H ₂ O	1.81	
ZnSO ₄ x 7H ₂ O	0.22	25 ²
CuSO ₄	0.051	
H ₃ MoO ₄ x H ₂ O or	0.09	
Na ₂ MoO ₄ x 2H ₂ O	0.12	
1M KH ₂ PO ₄ (ph to 6.0 with 3M KOH)	136	12.5
Iron (Sprint 138 iron chelate)	15	75

¹ For a 1:1 solution to use with 25L of water.

² All the minors elements are grouped, in the right proportions, in a "minor" solution.

Appendix D

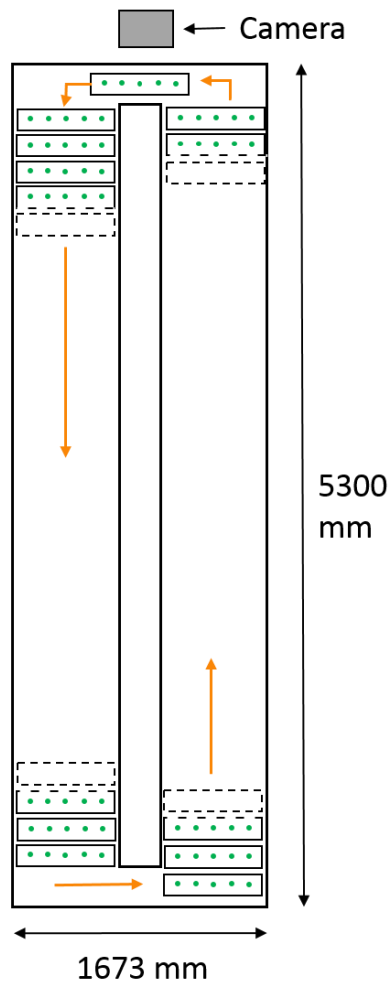
Phenotyping platform information file

JRA2 - Jan. 2018

Platform name

Partner site	UCL
Site and installation	Site: Louvain-la-Neuve, Installation: Aeroponics
Contact person(s)	Xavier Draye xavier.draye@uclouvain.be

Description of the platform structure

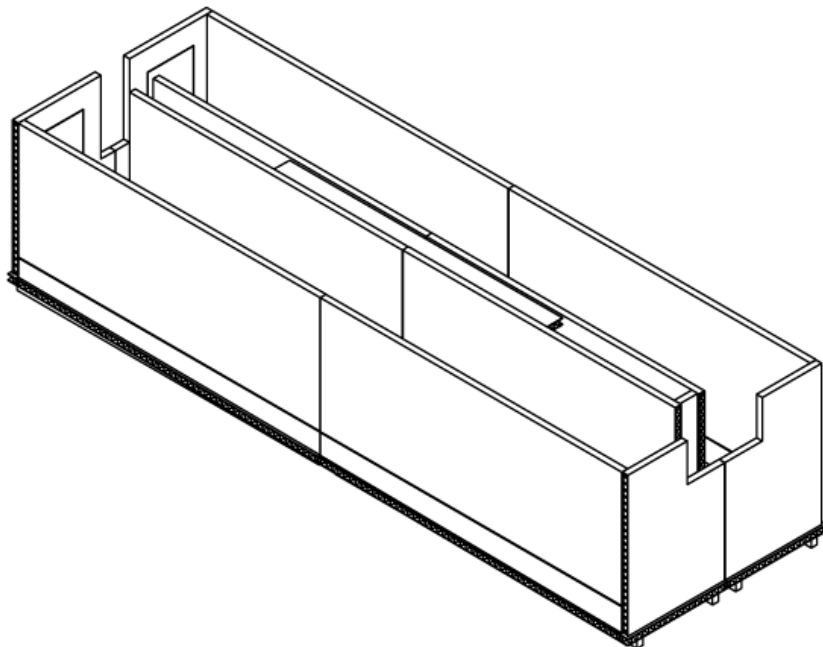


APPENDIX D. PHENOTYPING PLATFORM INFORMATION FILE

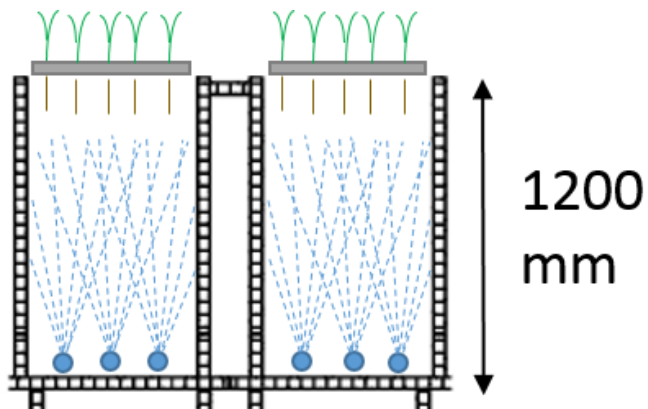


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Aeroponic tank: plants are hold on strips, 5 plants per strip (green dots on layout). There are 99 strips in the tank for a total of 495 plants/tank. Strips move in the direction indicated by orange arrows. A full revolution takes 2 hours. When strips pass in front of the camera, at the top of the layout, plants are imaged individually.



3D view of one tank, without the strips.

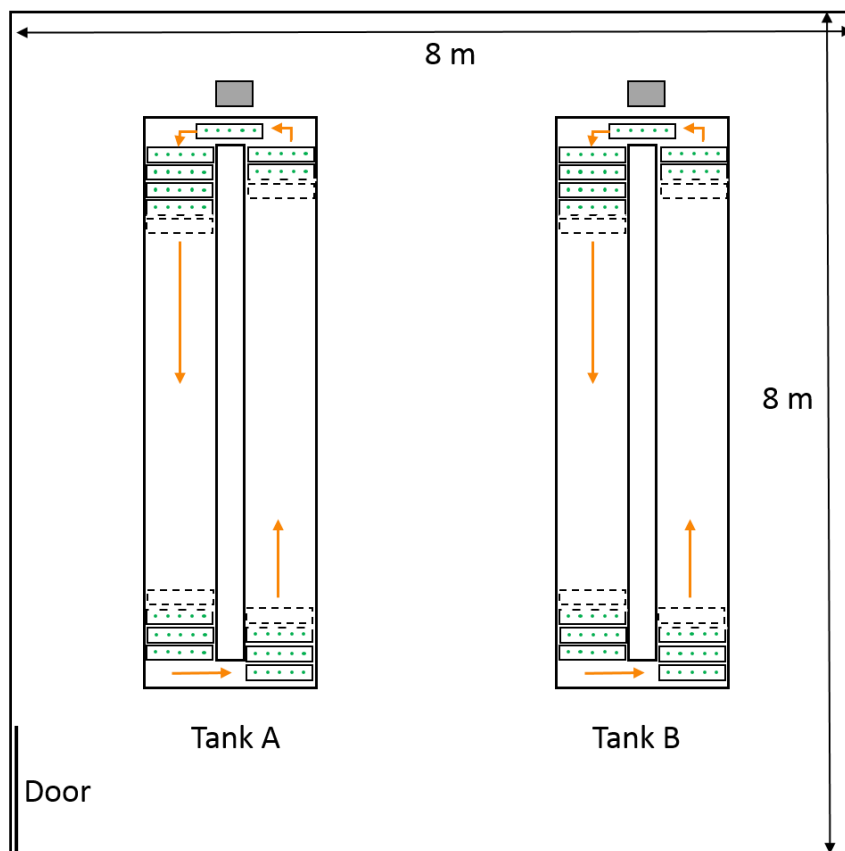


Transversal view of the aeroponic tank: 3 sprinklers are placed regularly in the bottom of each side of the tank. The sprinklers spray nutrient solution at regular interval, set by the operator. The spraying

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pattern (interval and duration) can be differentiated between day and night and can be modified at any moment of the experiment.

2 identical tanks are available in the installation, located next to each other in the same greenhouse.



Sources and directions (if known) of environmental variations in the installation

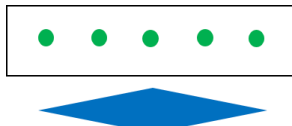
- 1) Between the 2 tanks.
- 2) The side of the tank placed along the greenhouse wall may be warmer than the side near the centre of the greenhouse because of the presence of heating pipes along the walls.

APPENDIX D. PHENOTYPING PLATFORM INFORMATION FILE



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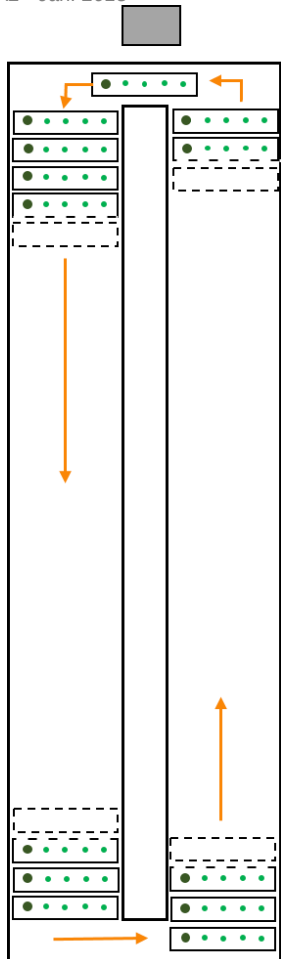
- 3) Inside each tank, between plants that grow in the middle of the strip as compared to plants growing at the border of the strip. We suppose that the plants at the extremity of the strip may receive a bit less water than the others.



Layout of a strip with supposed variation of water availability: more water in the middle and less in the border

- 4) Last year, we observed that the plants growing on the left side of the strips were growing faster than the ones growing on the right side. We understood that the lamps were not exactly centred in the middle of the tank. We moved the lamps to put them exactly at the centre of each tank but we haven't done any new experiment yet.

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Layout representing the plants that grow faster on the left side of the strips. The plants keep moving inside the tank but the left/right distinction is maintained during the whole experiment.

As strips keep moving within each tank, we don't expect to observe environmental variation between the different strips of each tank.

Description of experimental design and randomization and a motivation for the design and the randomization

APPENDIX D. PHENOTYPING PLATFORM INFORMATION FILE

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- Design

Completely randomized design: individual plants are located in a strip and at a position randomly with Excel.

+ 2 treatments (eg: shadow, change of nutrient solution properties...) corresponding to the 2 tanks
OR 2 blocks corresponding to the 2 tanks

- Design specifications

- Motivation

How plant positions are defined and recorded in the experiment

How are the pot positions defined according to the design, i.e. how are the spatial coordinates defined (see example 6)?

QR code associated to each plant



Number of the QR code:

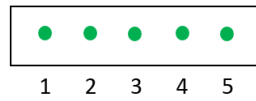
Ex: B_76_5

B: tank id (A or B)

76: strip id (from 1 to 99)

5: position in the strip (from 1 to 5)

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If pots are rearranged during the experiment, how is the change in spatial position recorded?

All strips move at the same pace. Each plant passes every 2 hour in front of the camera, where a picture is taken. The time of the picture enables to record the moment at which each plant passes in front of the camera. It would be possible to compute the pathway the plant had in the tank between two pictures.

No changes between the two tanks or within each strip (position 1 to 5)

If repeated measurements are taken, at what times are these taken?

Every 2 hours, 24h a day

Leuven Statistics Research Centre (LStat)

Celestijnenlaan 200 B

3001 EVERLEE, BELGIË

tel. +32 16 32 88 75

<https://lstat.kuleuven.be/contact>



