GENE EXPRESSION VARIATION UNDER DYNAMIC GROWTH CONDITIONS IN Arabidopsis thaliana

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Word Counts:

Introduction: x words

Results: y words

Discussion: z words

Abstract

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Thank you all,

Kevin

Questions to people reading this

Does all code have to go in the appendix? This could make it
 VERY long. Or is it ok to give links to github or similar (somewhere publically available).

general notes and TODOs

 $\bullet\,$ Ensure "static growth conditions" is use throughout, not standard GC

- standardise emulate/mimic/simulate references when talking about diff between reality and dynamic cond
- get really anal about stats being reported: use d.f., stat value, and P.
- Standardise the terminology around excess dynamic growth condition vs hot high light.
- ensure all latex cross references are pointing somewhere logical.
- backslash space after all macros
- remove all references to pseudo-natural

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Chapter 1

Introduction

Nearly all terrestrial biomass, including human life, depends on the primary productivity of higher plants (photosynthesis) (Johnston et al. 2009). This primary productivity is reduced when plants are stressed by interaction with their environment (Mittler 2006; Mittler and Blumwald 2010). Abiotic stresses alone account for over 100 billion of dollars of crop losses per annum in the United States alone (Mittler 2006). Therefore, the study of plant-environment interactions is crucial to improving agricultural yield and understanding functional plant ecology in a changing climate.

Central to increasing the understanding of plant-environment intearctions is the development of laboratory study systems that allow classical mechanisite studies of abiote stress response to be placed in the context of the conditions plants experience in natural or cultivated environments outside laboritories. Several techniques and approaches that have been developed over the past decade aide the study of plant stress response. In this thesis, I discuss, develop and apply novel techniques that enable the global study of gene expression in laboratory growth conditions which mimic field-like combinations of light, temperature and humidity. Specifically, I examine the effect of light intensity on global gene expression, within the framework of growth conditions whose temperature, humidity and light follow trends that approximate those observed

in regional climates.

1.1 Abiotic Stress: A limit to Plant Productivity

Abiotic stresses are the non-living stresses imposed upon plants by their growth environment. This includes deletatious extremes in or combinations of environmental variables such as temperature, humidity, osmote potential, water availability, light quality and quantity. Specifically, osmotic and drought stresses cause a decrease in the ability of plants to transpire and obtain inorganic carbon vital to photosynthesis (Mittler 2006; Seki et al. 2003). Excess light creates harmful reactive oxygen species (ROS) that damage the delicate photosynthetic aparatus (Apel and Hirt 2004; Asada 2006; Foyer and Noctor 2009; Li et al. 2009; Mubarakshina et al. 2010; Niyogi 1999). Extremes of temperature impair and damage to many enzymes, particularly those involved in metabolism (Atkin and Tjoelker 2003). Together, abiotic stresses limit plant productivity through their detrimental effects upon the ability of plants to assimilate biomass, and coping mechanisms that plants have evolved generally come at a yield penalty (Mittler 2006). As a result, enormous effort has been directed towards the elucidation and improvement of these coping mechanisms. In particular, the mechanisms by which excess light impact on plants, and how plants respond to this stress, have been major foci of research in the past decade.

blurry comment from barry

Plants respond to abiotic stress via a myriad of mechanisms (Demmig-Adams and Adams 1992), therefore responses to abiotic stress must be studied via a variety of techniques. Studies of photosynthetic characteristics by chlorophyll fluorescence (reviewed in Baker 2008) have illustrated the negative effects of abiotic stress on photosynthesis (Alter et al.

2012; Külheim, Ågren, and Jansson 2002; Mishra et al. 2012; Tikkanen et al. 2012). Phenotypic analysis, particularly of plant morphology, has given insight into the effects of abiotic stress (Armstrong, Wardlaw, and Atkin 2007; Wituszyńska et al. 2013). Studies of the transcriptome, or global cellular pool of expressed genes, have provided insight into plant responses to many abiotic stresses and combinations thereof (Atkinson, Lilley, and Urwin 2013; Kimura et al. 2003; Rossel, Wilson, and Pogson 2002; Seki et al. 2001).

1.1.1 Coping with Detrimental Effects of Excess Light GENOTYPES) ARE Photo-oxidative damage and Photoprotection MORE SUITED FOR DIF

The ability of plants to utilise energy from light to fix inorganic carbon into organic biomass is reduced when the quality or quantity of light a plant receives is not optimal. The quantity of light a plant receives is in excess when plants are unable to utilise energy obained from absorbed photos to fix carbon (Li et al. 2009). Excess light is particularly damaging, causing both reductions in photosynthetic ability, termed photoinhibition, , and cellular or tissue damage and death (Asada 2006; Li et al. 2009; Niyogi 1999; Suzuki et al. 2012). The indispensability of photosynthesis has caused the evolution of mechanisms by which the detrimental effects of excess light can be minimised. These mechanisms, collectively termed photoprotection, work to dissipate excess energy, reduce the amount of light absorbed, and prevent and repair any damage caused (Niyogi 1999; Takahashi and Badger 2011)

1.1.2 Physiological Responses to Excess Light

Upon exposure to excess light, plants immediately begin to mitigate its detrimental effects (Demmig-Adams and Adams 1992; Niyogi 1999). These include mechanisms of respose and damage mitigation within the

better papers JB ADD SOMETHING ABOUT HOW DIFFERENT GENOTYPES) ARE MORE SUITED FOR DIF-FERENT AMOUNTS OF LIGHT EG SHADE VS SUN PLANTS, FOREST, WOODLAND, SAVANA TYPES, TROPIC VS TEM-PERATE, WHAT IS THE GENETIC BASIS OF THIS DIFFERENTIATION IN LIGHT SENSITIVITY? SOME PATHWAYS ARE HYPOTHESIZED, KEY GENES IDENTFIED, PHYA CRY2

genetic variation

13

chloroplast (discussed below), as well as responses on a whole-cell scale. Chloroplast avoidance movement, the movement of chloroplasts parallel to the incident angle of light, decreases the amount of absorbed light (Kasahara et al. 2002). Transcriptional induction of heat shock proteins, antioxidant scavenging and photo-oxidative damage repair occur following excess light, and help to minimise oxidative damage (discussed in detail in subsection 1.1.3) (Jung et al. 2013; Niyogi 1999; Rossel, Wilson, and Pogson 2002). Production of anthocyanin, a protective class of pigments, is induced by the production of reactive oxygen species due to excess light (Vanderauwera et al. 2005). Together, these responses serve to reduce the photo-oxidative damage of cells and tissues due to excess light.

Several mechanisms of photoprotection occur within the chloroplast. Non-photochemical quenching (NPQ) dissipates excess energy from excited state chlorophyll molecules as heat (Müller, Li, and Niyogi 2001). It occurs in photosystem II (PS II), and is particularly important during rapid changes in light intensity (Külheim, Ågren, and Jansson 2002). Cyclic electron flow occurs in photosystem I (PS I) and acts by decreasing the pH of the thylakoid lumen, which in turn is thought to stabilise the oxygen evolving complex and aid NPQ (Takahashi et al. 2009). State transitions act via phosphorylation of PS II and light harvesting complex II (LHC II) proteins. State transitions reversibly alter the balance of excitation energy between PS I and PS II (Tikkanen et al. 2006). State transitions lower the light harvesting ability of PS II and prevent absorption of excess light by PS II, via reversible dissociation of the LHC II and PS II (Johnson et al. 2011). The Arabidopsis mutants non-photochemical quenching $\underline{1}$ (npq1) and \underline{n} on-photochemical quenching 4 (npq4), proton gradient <u>regulator</u> $\underline{5}$ (pgr5), and $\underline{s}tate$ <u>transition</u> $\underline{7}$ (stn7) and $\underline{s}tate$ <u>transition</u> g (stn8) are defective in these three mechanisms respectively, and enable the study of the mechanisms underlying chloroplastic response to excess light and fluctutions in light intensity.

ps 1 and 2 fig

1.1.3 Transcriptional Responses to Light Stress

Transcriptomics, or the global study of gene expression (discussed further below), has been used to study the response of plants to excess light. The excess light induced expression of a number of genes including ASCOR-BATE PEROXIDASE 2 (APX2) (Mühlenbock et al. 2008; Rossel, Wilson, and Pogson 2002), EARLY LIGHT INDUCED PROTEINS 1 and 2 (ELIP1 and ELIP2) (Adamska 1997; Rossel, Wilson, and Pogson 2002), and various heat shock proteins (HSPs) (Rossel, Wilson, and Pogson 2002) is well established. These proteins are induced by alternative putative mechanisms, allowing for differentiation between oxidative induction of retrograde chloroplastic stress signals (APX2) (Karpiński et al. 1999), ROS-mediated but photoreceptor-dependent induction of ELIP1 (Kleine et al. 2007) and ROS-induced, heat shock transcription factor (HSF) mediated induction of HSPs (MILLER and MITTLER 2006). The light induced down-regulation of LIGHT HARVESTING CHLOROPHYLL a/b BINDING LHCB family genes is also well established (Mishra et al. 2012; Rossel, Wilson, and Pogson 2002). Therefore, these genes are used in this thesis as markers of transcritpional response to excess light, allowing comparison of transcriptional responses observed in this thesis to be compared to relevant previous works.

The rapid response of the wider *Arabidopsis* transcriptome to changes in light intensity has been demonstrated (Rossel, Wilson, and Pogson 2002). Acclimation to excess light conditions affects the steady-state level of expression of several classes of genes. In the cyanobacterium *Synechocysitis* sp. PCC 6803, genes involved in light capture are down-regulated and homologues of heat-shock proteins up-regulated after 15 hours of excess light treatment (Hihara et al. 2001). In rice, similar

transcriptional down-regulation of light harvesting and up-regulation of photoprotection after 24 or 72 hours of excess light treatment has been observed (Murchie et al. 2005). Using quantitative real-time PCR, a method to assay relative expression of single genes, Gordon et al. (2013) have show that repeated high light treatments lead to acclimation and reduced induction of high light responsive transcripts. These transcriptomic responses to light overlap with response to drought, pathogen or oxidative stresses as well as hormone response, reiterating the requirement of stresses to be considered in combination. No genome-wide study of transcriptional response to altered light intensity in combination with other abiotic stresses has, to my knowledge, been published.

kimura, other Pogson microarrays

In addition to light quantity, plants perceive alterations in light quality, or the spectral composition of light. Gordon et al. (2013) demonstrate that excess light induced expression of RRTF1 and ZAT10 was dependent on wavelength of excess light in Arabidopsis. Light quality is particularly important in studies of photo-oxidative damage, as the extent of photosystem II damage is not consistent across the visible spectrum (Takahashi et al. 2010), and photo-oxidative damage is relatively more severe under light of wavelengths between 580-620nm than in the remainder of the visible spectrum, which overlaps with a density peak in the spectral power density of fluorescent lamps (see Figure 2.7). The integration of light quality into assays which test plant responses to altered light intensity is lacking in the vast majority of works on the topic.

1.1.4 Plant Growth Under Field Conditions

Plants grown in the field show markedly different phenotypes and responses to those grown in the lab. Under field conditions, plants respond to the combination of stresses they encounter in complex, natural environments. However, in the laboratory plants are exposed to predetermined stresses (or combinations thereof) hypothesised to provoke

perhaps spruik the SPC here also. LED lights are arriving, studies emerging, and you with SPC are in the lead a mechanism or response. This includes metabolic profiles (Jänkänpää et al. 2012), transcript abundances (Mishra et al. 2012; Wituszyńska et al. 2013), photosynthetic differences (Mishra et al. 2012), and survival and fecundity (Külheim, Ågren, and Jansson 2002). Several authors report overlap between genes expressed upon treatment with excess light and those constitutively expressed in field conditions (Mishra et al. 2012; Wituszyńska et al. 2013).

more field constitutive expression papers

1.2 Abiotic Stresses: Studied Individually, Encountered Combinatorially

Despite the large body of published works elucidating responses to abiotic stresses, the majority of works focus on an abiotic stress in isolation (Atkinson and Urwin 2012; Mittler 2006). While there is much merit to this reductionist approach, in natural or agricultural situations, plants rarely experience an abiotic stress in isolation. Unfortunately, this reductionist approach contributes to the difficulty in translating stress-tolerant lines of plants developed in labs to agrigculaural crops that are more field hardy (Mittler 2006; Mittler and Blumwald 2010).

Recent laboratory studies have demonstrated temperature interacts with light stress in a detrimental fashion. At cold temperatures, unacclimated plants are less able to dissipate excess energy to avoid photo-oxidative damage, leading to altered PS I/PS II redox poise (Armstrong, Wardlaw, and Atkin 2007). Many authors demonstrate temperature dependence on the transcriptional response to excess light (jung_subsets_2013; Rossel, Wilson, and Pogson 2002). Interactions with temperature have been observed in the transcriptional response to drought (Rizhsky, Liang, and Mittler 2002; Rizhsky et al. 2004; Seki et al. 2003; Seki et al. 2001). Additionally, interactions between biotic and abiotic stresses exist (Atkinson, Lilley, and Urwin 2013; Atkinson and Urwin 2012; Mit-

tler 2006). The data these authors present warrants a paradigm shift towards the study of abiotic stresses in combination, as advocated by Mittler (2006).

find something more convincing than that paper of Owen's

Field-grown plants tend to demonstrate reduced survival, reproduction and altered transcriptional responses compared to those grown in supposedly comparable lab conditions. For example, in Arabidopsis thaliana the reduction in reproductive success caused by the npq1 and npq4 mutations was severe in the field but had no impact under static sufficient light in the lab (Külheim, Ågren, and Jansson 2002). Similarly, photoinhibition, physiological symptoms of abiotic stress and expression of high light-induced transcripts were more severe in field-grown Solidago altissima than those grown in comparable lab conditions (Barua and Heckathorn 2006). Furthermore, Wituszyńska et al. (2013) observe increased steady-state expression of genes involved in response to excess light and photo-oxidative damage under field conditions compared to labgrown plants, and concomitant physiological acclimation to variable light intensity. Similar findings are presented by Mishra et al. (2012), who found increased NPQ and expression of ELIP proteins and decreased LHC-associated protein abundances in field-grown Arabidopsis. These field studies reiterate interactions noted in laboratory studies, and provide impetus for the study of field-like combinations of abiotic stresses under controlled laboratory conditions, to facilitate the mapping of QTLs for combinatorial stress tolerance (Li et al. 2010; Li et al. 2006).

smoother segue into
GWAS/QTL OUR OLD
PAPERS GIVE YOU THIS
IF YOU REVIEW THEM.

1.3 Transcriptomics: Assessing Global Expression

By studying how, when and to what extent each gene in the genome is expressed, we can gain insight into the response to any perturbation to a plant's environment. RNA sequencing (RNAseq) is a modern method of whole-transcriptome quantification. By creating cDNA from the cellular pool of mRNA, and sequencing libraries of this cDNA using highthroughput sequencing (HTS), quantification is possible (Lister, Gregory, and Ecker 2009; Wang, Gerstein, and Snyder 2009). The process of RNAseq library creation involves the extraction and purification of intact mRNAs, their conversion to complimentary DNA (cDNA), fragmentation, end-repair and A-tailing to allow sequencing adaptor ligation, ligation of sequencing adaptors and library amplification (Kumar et al. 2012; Wang, Gerstein, and Snyder 2009). When sequenced using HTS, raw sequice data is obtained with sequences derived from mRNA molecules proportional to their abundance in the sample mRNA pool. Computational analysis is required to provide quantification from this raw sequence data (Nookaew et al. 2012; Van Verk et al. 2013). This process involves raw sequence quality control, alignment of raw sequence reads to the genome, and quantification of gene-wise expression by aggregating the number of sequences that align to each gene. Quantitative data undergoes statistically rigorous normalisation before hypothesis testing occurs (Robinson, McCarthy, and Smyth 2010; Robinson and Oshlack 2010). Post-hoc analyses of expression data can be performed to glean summarised biological meaning from genome-wide differential expression patterns, including gene ontology (GO) term enrichment analysis (Avraham et al. 2008; Berardini et al. 2004) and gene set enrichment analysis (Kim 2012; Subramanian et al. 2005; Väremo, Nielsen, and Nookaew 2013; Yi, Du, and Su 2013). The RNAseq analysis process takes raw sequence data from high-throughput sequencing of RNAseq libraries and generates data that can be interpreted in the context of the biological basis underlying the experiment, and is presented graphically in Figure 1.1

maybe too methodsy

1.4 Thesis Aims

The overall aim of this project is to examine the effect of light intensity on *Arabidopsis*, within the framework of realistic combinations of abiotic stresses. The remainder of this thesis is devoted to the examination of the following aims:

- Design and implement "Dynamic Growth Conditions" which mimic regional climates
- 2. Select optimal software for the high-throughput study of transcriptome dynamics using High-throughput Sequencing, and implement a framework for generation of analysis pipelines to do so
- 3. Determine the transcriptional response of *Arabidopsis thaliana* to combinatorial application of abiotic stresses, using dynamic growth conditions

Due to technical obstacles and delays, the aims of this project have evolved over the year. As it became apparent that technical difficulties would prevent high-throughput RNAseq experiments required to map expression QTLs in the time available, the aims "Examine the extent of genetic variation in gene expression under these dynamic light conditions, and elucidate gene regulation networks controlling gene expression." and "Explore the effect of genotype-environment interactions on gene expression under dynamic light conditions." were discontinued.

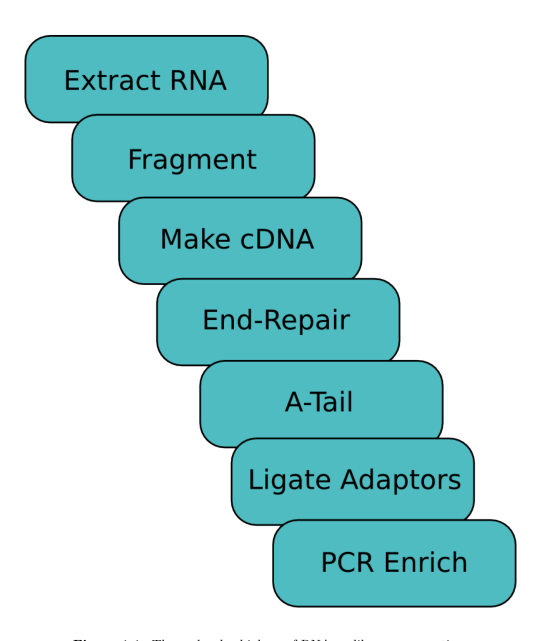


Figure 1.1: The molecular biology of RNAseq library preparation.

Chapter 2

Design and Implementation of Dynamic Growth Conditions

2.1 Background, Aims and Hypotheses

If one studies organisms or mechanisms which have evolved in a given environment, as nearly all biologists do, it follows logically that the study environment should be as similar as possible to the environment in which the subject has evolved. This underpins the enormous wealth of literature derived from field studies. In areas of molecular plant science however, our study organisms are often placed in environments highly dissimilar to those in which our subjects are hypothesised to have evolved (Mittler 2006; Mittler and Blumwald 2010). This has a particular impact on areas of plant science which study the interaction of plants with their environment, and may alter the conclusions of such studies considerably.

In this chapter I describe artificial growth conditions that I have created which vary on diurnal and circannual cycles in an analogous manner to the regional climates cultivated or naturally growing plants experience. This class of laboratory growth condition are termed "dynamic growth conditions". The term dynamic is in contrast with the often unnaturally benign, highly "static growth conditions" typically used in the propagation, growth and experimentation of plants in laboratory settings, in

which temperature, humidity, photoperiod, light intensity and light quality are constant across the period of plant growth. I hypothesise that plants grown under dynamic growth conditions will exhibit phenotypes more similar to those grown outdoors under natural environments, as these dynamic growth conditions are more similar to natural environments when compared to static, benign laboratory growth conditions. This hypothesis is tested in later chapters of this thesis.

A key driver behind the development of dynamic growth conditions is the ability to simulate regional climates in as reliable and reproducible manner as required for scientific study. The second aim for this chapter of my thesis is to create software to allow dynamic growth conditions to be implemented with existing hardware at the ANU. This software should be able to use the model outputs of software previously written to create regional climate models (Spokas and Forcella 2006) that govern dynamic growth conditions, to control multi-spectral LED lamps and computerised reach-in growth chambers. Successful completion of this aim has allowed the implementation of dynamic growth conditions, and their use in research into plant-environment interactions in *Arabidopsis*.

2.2 Materials and Methods

2.2.1 The SpectralPhenoClimatron and Implementation of spcControl

The SpectralPhenoClimatron is a new facility within the Research School of Biology, consisting of computer controllable plant growth cabinets featuring multi-spectral LED lamps, and real-time imaging hardware. Conviron PGC20 reach-in growth chambers (Conviron, Winnipeg, Canada) have been retro-fitted with four Heliospectra Model L4A Series 10 multi-spectral LED growth lamps (Heliospectra AB, Sweden) per chamber, and an image-based phenomics systems (Canon EOS DSLR cameras and

other consumer hardware). The Conviron PGC20 cabinets have a capacity of 320 5cm by 5cm plant growth containers, or 16 250x200mm standard nursery seed trays (e.g. Garden City Plastics part TRSR00). The Heliospectra L4A Series 10 LED lamps contain 7 LED wavelength channels: 400nm (sub-blue), 420nm (blue), 450nm (blue), 530nm (green), 630nm (red), 660nm (red) and 735nm (far red).

Both the Heliospectra LED lamps and Conviron growth cabinets can be controlled via the Telnet protocol, and custom control software was created to utilise this feature. The spcControl program is invoked with a regional climate model in comma-separated value (CSV) format, describing the temperature, humidity, and intensity of each LED wavelength for climate models per SolarCalc calculations (Spokas and Forcella 2006). This software simultaneously sends telnet commands that control growth chamber temperature and humidity to each growth cabinet, and commands that control light quantity and quality to each of four LED arrays per cabinet. The success of each set of control commands is reported to an off-site database.

2.2.2 Design of dynamic growth conditions

SolarCalc was used to create climate models underlying dynamic growth conditions (Spokas and Forcella 2006). Model settings that I used in the creation of the dynamic conditions are described in Table 2.1 if they deviate from program defaults. As SolarCalc by default simulates the climate of a location without variable weather, post-processing work was required to create conditions that mimic cloudy and intermittently cloudy days. A SolarCalc model with a neutral density shade such that model sunlight intensity was 45% that of an unshaded model was created, and the result of both the shaded and unshaded models were spliced together to form a third condition whose light intensity fluctuated on a two hour sufficient light, one-hour excess light rotation, using the

spliceSolarCalc.py script described in subsection 6.2.1. The temperature, humidity and light quality was preserved across these conditions.

2.2.3 Measurement of Spectral Power Density

define spectral power density

Spectral power density, or distribution of light intensity across the visible spectrum, was quantified using a spectrophotometer to record spectral power density across the spectrum between light of wavelengths 400nm to 800nm, with 2nm wavelength resolution. Sun and shade spectra were obtained on July 18 at the Acton campus of the ANU, in a clear, open space and under heavy shade from mature trees of various species, in the courtyard between buildings 46 and 48. Spectra of laboratory growth conditions were obtained from a Conviron PGC20 by placing the spectrophotometer on the lowest shelf level while fluorescent lamps or Heliospectra L4A series 10 lamps were illuminated at their highest intensities. In the case of the Heliospectra L4A series 10 LED lamps, measurements from directly under a single unit were recorded. Intensity-normalised spectral power density was calculated by normalising the intensity recorded for each wavelength by total light source intensity.

2.3 Results

2.3.1 Computer Control of the SpectralPhenoClimatron

To create dynamic growth conditions which change on diurnal and circannual cycles, high temporal resolution is required. Whilst SpectralPheno-Climatron hardware can produce static growth conditions, external software is required enable the creation of such dynamic conditions. Thus, I have created software, spcControl, to do so. This software can im-

Parameter	Temora Setting
Simulation End Date	31/12/12
Simulation Start Date	01/09/12
Chamber Max RH	85
Chamber Min Temp	$5\mathrm{C}$
Chamber Date	01/03/13
Site Elevation	$300 \mathrm{m}$
Site Latitude	-34.446556
Site Longitude	147.53334
Timezone	Sydney
Update Frequency	$1 \min$
Weather Year	2010
Lighting	LED
Set to Threshold	Yes
LED1 Power Threshold	5.00%
LED1 Wavelength	$400\mathrm{nm}$
LED1 Weight Multiplier	5.2
LED2 Power Threshold	5.00%
LED2 Wavelength	$420\mathrm{nm}$
LED2 WeightMultiplier	4.12
LED3 Power Threshold	5.00%
LED3 Wavelength	$450\mathrm{nm}$
LED3 Weight Multiplier	4
LED4 Power Threshold	5.00%
LED4 Wavelength	$530\mathrm{nm}$
LED4 Weight Multiplier	3.92
LED5 Power Threshold	5.00%
LED5 Wavelength	$630\mathrm{nm}$
LED5 Weight Multiplier	4.88
LED6 Power Threshold	5.00%
LED6 Wavelength	$660\mathrm{nm}$
LED6 Weight Multiplier	0.68
LED7 Power Threshold	5.00%
LED7 Wavelength	$740\mathrm{nm}$
LED7 Weight Multiplier	3.64

Table 2.1: Parameter settings of SolarCalc used in the creation of dynamic conditions. LEDs 1-7 correspond to wavelengths 400nm (sub-blue), 420nm (blue), 450nm (blue), 530nm (green), 630nm (red), 660nm (red) and 735nm (far red).

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plement dynamic growth conditions given a climate model generated by SolarCalc (Spokas and Forcella 2006). spcControl will, at time intervals specified in the climate model, send control commands to both the LED arrays and plant growth cabinet, updating LED intensity, temperature and humidity. This process takes around 30-45 seconds, and thus can occur up to every minute, giving extreme temporal resolution in growth condition control. Additionally, as the commands are sent out synchronously, lighting, temperature and humidity will never go "out of sync" if a power outage or device failure occurs. Furthermore, to ensure reliable operation and detection of faults, at every time-point specified in the SolarCalc model, the success or failure is communicated to an off site database, and any error message is emailed to an administrator. An additional program, spcControl.monitor, polls this database and guards against failure of hardware, control computers and software, informing an administrator upon any failure. Together, these features allow highly reliable control of growth cabinets with temporal resolution suitable for the implementation of dynamic growth conditions, while mitigating the risk posed by inconsistencies and failures in equipment, ensuring the validity and reproducibility of scientific experimentation.

The spcControl python module runs with python version 3.2 or later. It is modular in design, with a main program loop which sends a control line to each sub-module per the schedule given by the SolarCalc model. Sub-modules then parse this line, and a configuration file, to formulate commands sent to the relevant device(s); sub-modules for Heliospectra L4S LED lamps and Conviron PCG20 chambers have been implemented. This modular design means that, given hardware specifications, creating new sub modules to control other hardware configurations would be relatively trivial. Status reports are sent to an external PostgreSQL database, and email error messages are generated from within python if an error occurs. In all, this consists of over 740 lines of code and configurations.

ration. This software is actively maintained; as software bugs, hardware limitations and feature requests are discovered solutions are provided. The codebase has expand from a simple script to a fully-fledged python module with 16 versions released thus far (Appendix subsection 6.3.1).

2.3.2 Sufficient, Excess and Fluctuating Dynamic Growth Conditions

To investigate the effect of altered light intensity on the Arabidopsis transcriptome, three novel growth conditions specifically designed to mimic the dynamic nature of temperature, humidity and light intensity and quality that occurs outdoors. The sufficient light dynamic growth condition corresponds to approximately the same daily integral of light as "standard static growth conditions" of 120-150 μ mol photons $m^{-2} s^{-1}$ with a 12-hour photoperiod. The excess light condition is approximately 250% brighter than the sufficient condition. The fluctuating light condition varies between sufficient and excess growth conditions on a 2 hour-1 hour basis, and is designed to simulate the pattern of light intensity variation caused by partial cloud. These conditions simulate the spring season, and display circannual or seasonal variation in temperature, light and humidity. As spring progresses, daily minimal and maximal temperature and peak light intensity increase, while minimal relative humidity decreases.

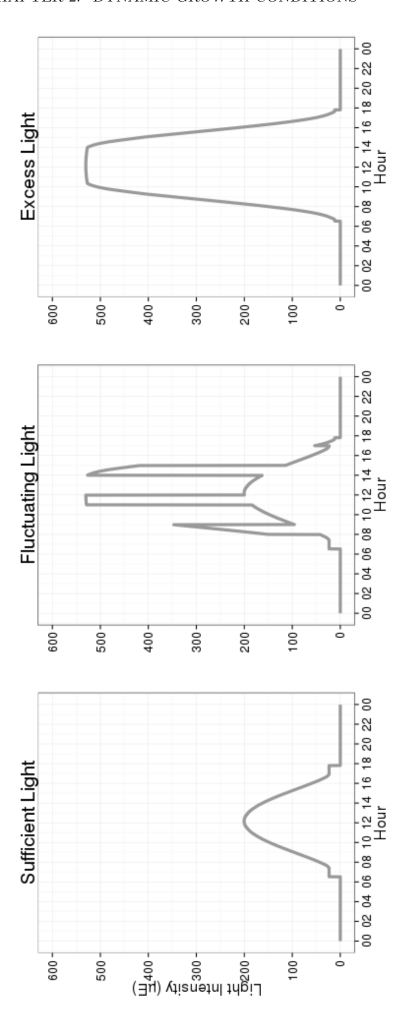


Figure 2.1: Diurnal variation in approximate light intensity of sufficient, fluctuating and excess light dynamic growth conditions (for model date 1 March). Note the altered light intensity between dynamic growth conditions, and identical photoperiod between conditions.

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The light quality of all dynamic growth conditions are markedly different to other light sources. Compared to the fluorescent lamps typically used in laboratory growth chambers the spectral power density, or distribution of light intensity across the visible spectrum, of LED arrays is less variable across the visible and adjacent spectrum, with intensitynormalised spectral power density of fluorescent and LED array light sources of 1.00 \pm 1.74 and 1.00 \pm 0.92 μ mol photons $m^{-2}s^{-1}nm^{-1}$ per total μ mol photons $m^{-2}s^{-1}$ respectively (means \pm SD; see Figure 2.7). The intensity-normalised spectral power density of sunlight on a clear day is remarkably even $(1.00 \pm 0.19 \ \mu \text{ mol photons } m^{-2} s^{-1} nm^{-1} \text{ per total}$ μ mol photons $m^{-2}s^{-1}$; Figure 2.7). The intensity normalised spectral power density (spectral power density per unit total light intensity) of canopy shaded light is similar to sunlight at wavelengths lower than approx 700nm, above which sunlight is not filtered by vegetation and thus is over-represented. Overall, the spectral power density of LED lamps is more even than that of fluorescent lamps, however it still deviates notably from that of sunlight.

The overall light intensity of these natural and laboratory light sources varies drastically. Representative measurements of intensity reveal open sunlight to have an intensity of 2480 μ mol photons $m^{-2} \, s^{-1}$ in the photosynthetically active spectrum on the day of measurement (July 18, 2013 in Canberra, ACT, Australia). The intensity of tree canopy shaded sunlight measured close by on the same day is much lower, at approximately 88 μ mol photons $m^{-2} \, s^{-1}$. This is in contrast with the light intensity under a single Heliospectra L4A series 10 LED lamp of 370 μ mol photons $m^{-2} \, s^{-1}$, and the intensity of light from fluorescent lamps was 140 μ mol photons $m^{-2} \, s^{-1}$ (data shown graphically in Figure 2.7).

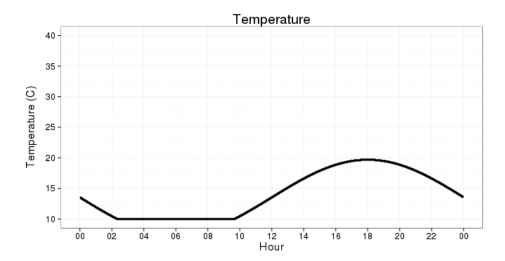


Figure 2.2: Diurnal variation in dynamic growth condition model temperature (for model date 1 March). Temperatures follow an approximation of those observed in temperate climates, reaching a minimum before sunrise (06:00), steadily increasing after sunrise to a peak immediately prior to sunset (18:00). Growth chamber hardware limitations prevent temperatures falling below 10 $^{\circ}$ C for extended periods, thus the model "bottoms out" where temperatures below 10 $^{\circ}$ C would have occurred (02:00 - 10:00).

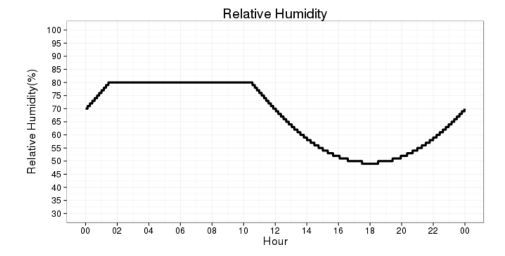


Figure 2.3: Diurnal variation in Dynamic Growth Condition model relative humidity (for model date 1 March). Humidity follows an inverse trend to temperature, peaking before sunrise (06:00) and reaching its minimum at approximately sunset (18:00). Similarly to temperature, growth chamber hardware limitations prevent relative humidities greater than 80% for long periods, and therefore humidity is capped at 80% between the hours of 02:00 and 10:00.

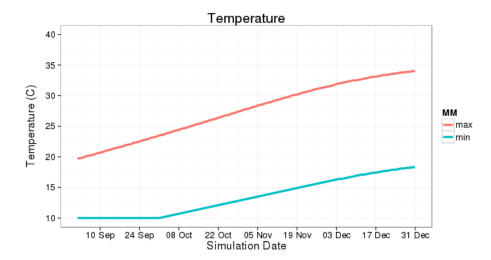


Figure 2.4: Circannual variation in daily minimal (blue) and maximal (red) Dynamic Growth Condition model temperature. Starting at model date September 1, both minimal and maximal daily temperature gradually increase, throughout spring and the first month of summer. Note also the hardware limitation of minimal daily temperature is alleviated after daily minimal temperature exceeds 10 °C (on approximately October 1st).

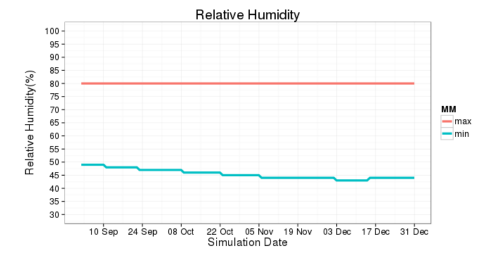


Figure 2.5: Circannual variation in daily minimal (blue) and maximal (red) Dynamic Growth Condition model relative humidity. Unlike the pattern observed in temperature, daily maxima do not increase across the modelled period, due to hardware limitations. However, daily minima in humidity does decrease over the modelled period, concomitant with observed daily maxima, partially preserving the inverse relationship between temperature and humidity.

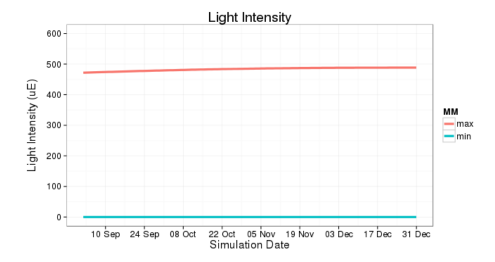


Figure 2.6: Circannual variation in daily minimal (blue) and maximal (red) Dynamic Growth Condition model light. Little change in daily maximal light intensity occurs, due to hardware limitations on the brightness of LED arrays.

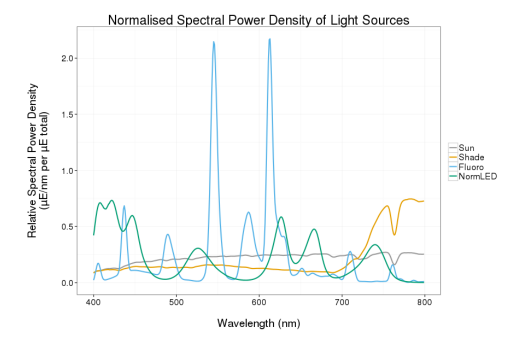


Figure 2.7: Intensity-normalised spectral power density of sunlight, shaded sunlight, fluorescent lamps and Heliospectra L4A series 10 LED lamps. Note the almost flat spectral density of sunlight, compare to the broad peaks of intensity of LED arrays, and large spikes of intensity (Mercury emission peaks) in fluorescent lamp spectra.

2.3.3 Novel Dynamic Growth Conditions That Simulate a Continental Gradient

The spcControl software I have developed can be applied to many experimental designs. For example, it was further used to create conditions which examine hypotheses beyond the scope this thesis. This includes two conditions to test the overall effect of environments with higher light and more diurnal variation in temperature, such as may be experienced in inland regional climates, compared to conditions with lower light intensity and lower diurnal variation in temperature, such as those often encountered in coastal climates. These conditions, named "NSW inland" and "NSW coastal" respectively, generally are more harsh than the sufficient, fluctuating and excess light dynamic growth conditions created for my experiments. Additionally, conditions required to conduct a virtual reciprocal transplant of *Pelargonium* species from Australia and South Africa were created, by simulating regional climates in two locations, in coastal New South Wales, Australia and coastal Western Cape, South Africa. The specifics of these dynamic growth conditions are beyond the scope of this thesis.

2.4 Summary and Technical Discussion

In this chapter, I present software enabling the implementation of dynamic growth conditions that mimic diurnal and circannual trends in temperature, humidity, photoperiod, and light quality and quantity observed in regional climates. This software enables the use of regional climate models to govern growth conditions in laboratory growth chambers, allowing reproducible and reliable implementation of dynamic growth conditions. Practically, there are shortcomings in the SpectralPheno-Climatron. These are largely limitations inherent to the hardware from which it is constructed, and include the limited light intensity, temper-

ature and humidity of the SpectralPhenoClimatron. The limited light intensity and the evenness of the spectral power density of Heliospectra L4A Series 10 lamps will be improved in an upcoming upgrade (pers. comm., Justin Borevitz). Despite these minor shortcomings, the SpectralPhenoClimatron is a phenomenal tool with which to study plant-environment interactions. It has been designed with large scale studies that elucidate underlying genetic mechanisms and examine genetic variation in these interactions in mind.

Three dynamic growth conditions to examine the effect of light intensity in field-like interaction with temperature, humidity and light quality were implemented. These conditions allow for light stress to be studied in the framework of combinatorial application of stresses, as recent literature has advocated (Mittler 2006; Wituszyńska et al. 2013). Examination of genetic variation in transcriptional, physiological or phenomic responses to altered light intensity may provide insight into mechanisms underlying response to light stress in field-like combinations with other abiotic stresses (Li et al. 2010; Li et al. 2006).

summary needs a reword to link it better

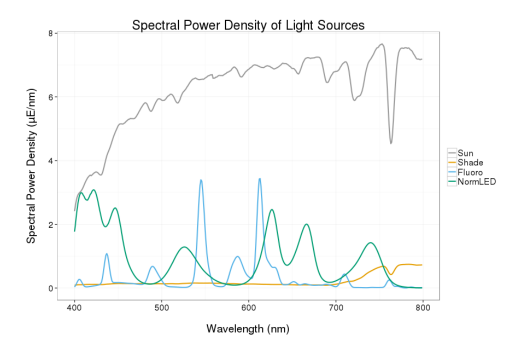


Figure 2.8: Spectral power density of sunlight, shaded sunlight, fluorescent lamps and Heliospectra L4A series 10 LED lamps. Note the higher intensity and less even spectral density of sunlight compared with other light sources.

Chapter 3

Improved Methodology for High-throughput RNAseq Experiments

3.1 Background, Aims and Hypotheses

RNA sequencing (RNAseq) is a modern method for genome-wide expression (transcriptome) quantification. RNAseq works by sequencing the mRNA pool of a cell, tissue or organisms (Martin et al. 2013; Wang, Gerstein, and Snyder 2009). As high-throughput, large-scale experiments such as QTL mapping and Genome Wide Association Studies (GWAS) become more prevalent, increasingly subtle biological contrasts are being examined. RNAseq is an incredibly sensitive tool to assay subtle changes in, for example, plant growth environment or subtle genetic variation (Martin et al. 2013). Typically, RNAseq analytic methods have been developed to investigate stark biological comparisons, such as comparison of healthy and diseased or mutant and wild-type tissues or individuals (Wang, Gerstein, and Snyder 2009). In order to analyse these large scale datasets, updated molecular and analysis methodology must be used.

To gain biological meaning from raw RNAseq data, an analysis pipeline must be employed. In this context, a pipeline is a series of software components that, in succession, manipulate a dataset to obtain a biologically relevant result. Best practice pipelines for RNAseq analysis exist (e.g. Van Verk et al. 2013), but often must be manipulated to suit the idiosyncrasies of each experiment. Thus, an aim in this chapter of my thesis is to create a framework allowing easy creation of pipelines by non-expert bioinformaticians with limited programming experience, and to use this framework to create high-performance pipelines suited to analysis of high-throughput RNAseq experiments.

Additionally, I have conducted an in silico experiment to test the effect of sequencing depth on statistical power of RNAseq experiments. Despite the rapid and continuing reduction in the cost of high throughput sequencing, it is still a very large component of the overall cost of RNAseq experiments (Wang, Gerstein, and Snyder 2009). This is particularly evident with regards to high throughput experiments, and is often combated by increasing multiplexing, i.e. reducing the amount of raw sequence data each sample yields, at a cost to statistical power (Kumar et al. 2012). Therefore, I am to determine the optimal trade-off between sequencing cost and statistical power. Similar experiments suggest an optimal depth of 10 million reads per sample for Chicken tissue samples (Wang et al. 2011). However due to its smaller transcriptome size, I hypothesise that the optimal sequencing depth for Arabidopsis will be smaller, specifically between 2 and 5 million reads, or between 48 and 96 libraries per Illumina HiSeq 2500 sequencing lane (that yield 200 million reads apiece, (glenn_field_2011; illumina_brochure)).

WHY: a pgf here explaining this is a framework, and that it is generalisable, and that it aims to detect subtle differences in high throughput experiments

3.2 Methods

3.2.1 External RNAseq Datasets

RNAseq datasets created by Peter Crisp and Barry Pogson were used both as trial datasets and external references in this thesis. The Rapid Recovery Gene Silencing excess light time-course experiment (hereafter referred to as the RRGS time-course) consists of samples taken in triplicate from an eleven-point excess light stress and recovery time-course. A. thaliana reference accession Col-0 were grown for 3 weeks under standard laboratory growth conditions ($\approx 150 \, \mu \, \text{mol photons} \, m^{-2} \, s^{-1} \text{light inten-}$ sity, 12 hour photoperiod, 21 °C daytime temperature, 21 °C night-time temperature). Whole rosette samples were taken before any treatment, after 30, 60 and 120 minutes of 8x excess light (1000 μ mol photons $m^{-2}s^{-1}$, unfiltered light from a sodium vapour lamp, hereafter EL), after 60 minutes of EL followed by 7.5, 15, 30 and 60 minutes of recovery under standard growth conditions, after 60 minutes of EL, followed by 60minutes of recovery, followed by another 60 minutes of EL, and before and after 60 minutes of EL 24 hours after the original 60 minutes of EL. This complex time-course is illustrated in Figure 3.1. RNA extracted from five plants per replicate was pooled, before Illumina libraries were created using the TruSeq V2 library preparation kit (part number 15026495) per manufacturer's instructions. These libraries were sequenced across two Illumina HiSeq 2500 sequencing lanes, yielding the RRGS timecourse RNAseq dataset. This dataset studies a timecourse over a treatment highly similar to that conducted in the dynamic growth condition experiment, allowing development and validation of bioinformatic protocols for experiments of a similar nature.

3.2.2 Development of an Improved Analysis Pipeline

Bioinformatic experiments were used to validate pipelines against the "gold standard" RNAseq analysis pipeline. In these experiments, programs selected through both literature review and searches of pre-publication software releases (e.g. software on github.com) were tested against a published best-practice pipeline (Van Verk et al. 2013). Specifically, the computational speed and efficiency, and the results obtained with these newer programs were compared to the analysis pipeline of Van Verk et al. (2013). This enables the development of higher-performance analysis pipelines suitable to high throughput experiments, with no cost to the quality of results obtained. A summary of programs selections is described in Table 3.1.

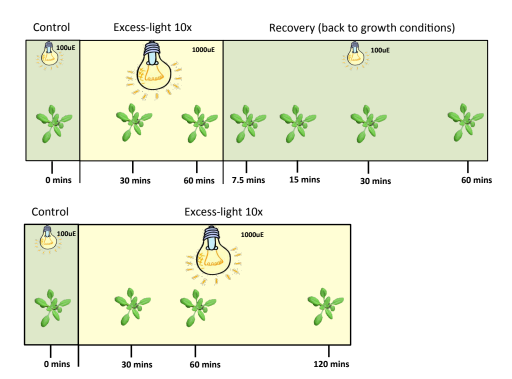


Figure 3.1: Illustration of the RRGS timecourse. Entire rosettes were harvested in triplicate at each indicated time-point along the excess light stress and recovery time-course. Two timepoints after a 24 hour recovery period are not show. This figure was created by Peter Crisp, and is reproduced with his permission

Program	Program's Role	Reasons for Selection	References
fastqc	Determine raw sequence quality of datasets	Easy of use and detailed reporting	(Andrews 2012)
scythe	Remove Illumina adaptor sequence from 3' ends of reads, allowing more accurate mapping	Author's claims of increased accuracy and speed	(Buffalo 2013)
seqtk	Remove sequences with low base-level quality from analysis	Mott trimming algorithm; fast	(Li 2013)
subread	Align short reads to genome	Fast RNAseq compatible	(Liao, Smyth, and Shi 2013b)
tophat2	Align short reads to genome while detecting $de novo$	Capable of de-novo splicing detection	(Kim et al. 2013)
subjunc	Align short reads to genome while detecting mRNA splicing de novo		(Liao, Smyth, and Shi 2013b)
featureCounts	Aggregate gene-wise counts of aligned reads to quantify expression	Fast and well supported	(Liao, Smyth, and Shi 2013a)
edgeR	Perform statistical normalisation and hypothesis testing	Statistically rigorous; supports multi-factor experiments	(McCarthy, Chen, and Smyth 2012; Robinson, McCarthy, and Smyth 2010)
goseq	Perform gene ontology term enrichment	Improved RNAseq- compatible statistical basis	(Young et al. 2010)

Table 3.1: A caption

Comparisons between the computational cost of four pipelines were conducted using a sub-sampled dataset. To demonstrate the improved performance of the aln_subread pipeline, it was compared to the aln_tophat, aln_tophat_htseq and aln_subread_htseq pipelines. The time UNIX command was used to summarise the computational cost of these four pipelines across five identical, independent, non-simultaneous runs. Four time-points of the RRGS timecourse dataset were sub-sampled to 500,000 reads by running seqtk sample -s 10 500000 on both forward and reverse read files, which extracts 500000 random read pairs preserving read pairing. An ANOVA analysis was performed to find significant differences in runtime and CPU utilisation between analysis pipelines (see ??).

To ensure that the subread aligner and featureCounts produced comparable results to the analysis pipeline of Van Verk et al. (2013), several diagnostic measures were used. Firstly, the percentage of reads mapped to the genome, and to protein coding loci within the genome was computed and compared. Then, sample-wise correlations between genewise counts calculated by each pipeline were calculated. Finally, genes called differentially expressed by each pipeline were compared. These measures allow verification of pipeline performance at three major stages in an analysis pipeline: alignment of short reads to a genome, gene-wise count summarisation, and statiscial testing for differential expression.

3.2.3 Measuring the Effect of Sequencing Depth on Analysis of Differential Expression

Six samples from the RRGS-Timecourse experiment (see subsection 3.2.1) were sub-sampled to allow investigation of the effect of sequencing depth on statistical power. To do so, the command seqtk sample \neg s 10 X was run on each pair of read files for these six samples, with X (number of reads to sample) set to 1000, 10000, 20000, 50000, 100000, 200000, 500000, 1000000, 2000000, 5000000 and 10000000. This sub-sampled

dataset allows for titration of the optimal sequencing depth (or multiplexing level) for high throughput experiments, balancing sequencing cost with statistical power.

For each subsampled dataset, the km_subread pipeline followed by the de_pairwise pipeline were applied to find differential expression between the control and 30 minute excess light timepoints (these pipelines are described in subsection 3.3.2). Several metrics were then used to summarise the effect of sequencing depth on the statiscial power of differential expression analysis. The number of genes called as differentially expressed at each sequencing depth was calculated, as was the common biological coefficient of variation. A third measure, the log-transformed mean expression level of the least-expressed differentially expressed gene and the overall least-expressed gene were calculated. These metrics were plotted against sequencing depth to give a graphical overview of the effect of reduction of sequencing depth.

3.3 Results

3.3.1 A Framework for the Creation of RNAseq Analysis Pipelines

In order to allow easy creation of diverse analysis pipelines for the multitude of possible experimental designs and methodologies, I have implemented a generic framework for the creation of RNAseq data analysis pipelines. This framework takes the form of "wrapper scripts", which act as wrappers around programs which other authors have created, and "pipeline" scripts, which combine these wrapped programs to perform an analysis. Wrapper scripts are the workhorses of a pipeline created with this framework, accepting three arguments: an input directory, and output directory, and arguments to be passed to the underlying program. Given these, the wrapper script will run the underlying program,

automatically detecting input files from the input folder and automatically accounting for experimental features such as single or paired-end sequence data. Pipeline scripts describe processes of analysis of RNAseq data. They combine wrapped programs together to perform an analysis specific to a dataset. Each pipeline is run in parallel to utilise multiprocessor computers, and every command is comprehensively logged, ensuring reproducibility. By removing the complexity of command syntax and increasing readability and reproducibility of pipeline workflows, I have enabled their use by a larger community of biologists without detailed training in bioinformatics, and ensured the reproducibility of results obtained.

3.3.2 An Improved Analysis Pipeline for Large Plant RNAseq Datasets

A series of two-step pipelines to analyse RNAseq datasets have been developed. Step one (aln_subread, aln_subjunc and aln_tophat) taking raw sequence reads and produces summarised gene-wise counts. Step two (de_pairwise and de_pairwise) applies statistical normalisation techniques and tests for differential expression. When applied combinatorially, these pipelines allow for different experimental designs to be analysed. Table 3.1 describes each software element of these pipelines, reasons for their selection, and references to literature describing each component (which for brevity are not included below).

The aln_subread pipeline

This pipeline is built around the subread aligner, a very fast RNAseq-compatible short read aligner of expressed RNA sequence to a reference genome. Firstly, quality of sequencing data is checked using the fastqc program (andrews_fastqc_2013), sequencing adaptors are removed with scythe (Buffalo 2013), and seqtk remove low quality se-

quences, before the quality is again checked using fastqc. The subread aligner then aligns reads to the current TAIR10 A. thaliana reference genome, accounting for alternative exon splicing. The resulting SAM file is converted to the BAM format, sorted and indexed, as required by downstream programs. Gene expression is then summarised gene-wise by counting the number of reads which align to genic loci with feature—Counts, completing this section of the analysis and the aln_subread pipeline.

programs citations

The aln_subjunc and aln_tophat pipelines

For studies examining alternative splicing of mRNA transcripts, an aligner able to detect splicing de novo is required (kim_tophat2_2013). Tophat2, one of the most popular RNAseq aligners, is able to align short reads while detecting slicing isoforms (kim_tophat2_2013). Subjunc is an extension to the subread aligner which it allows it to do so (Liao, Smyth, and Shi 2013b). The aln_subjunc and aln_tophat are identical to the aln_subread pipeline, aside from their use of the subjunc and Tophat2 aligners respectively, in place of the subread aligner, allowing study of alternative splicing. However, this de novo detection of splicing comes at a performance cost, and is not necessary for simple quantitation of gene expression. Splicing was not further analysed.

de_pairwise

For paired experimental designs, statistical tests can be performed pairwise between samples. To this end, the de_pairwise pipeline implements these tests using the edgeR R package. This pipeline first reads count files into a DGEList object, removes loci not detected above statistical noise and non-protein coding loci, then normalises counts using the Trimmed Mean of M values normalisation method of Robinson and Oshlack (2010). Common and gene-wise (i.e. gene-wise) dispersion are then calculated using the calcNormFactors, estimateCommonDisp and estimateTag-

wiseDisp functions respectively, yielding a DGElist object containing normalised counts and estimates of expression variability. Tests are then conducted pairwise between groups described in the keyfile, from data in this DGElist, using exactTest. This creates a list() of DGEExact objects, from which tables of differential expression and diagnostic plots can be created. A plot showing the relationship between the gene-wise Biological Coefficient of Variation (BCV) and gene expression level aims to show that variation is largely independent of expression level.

reword to stop it soundling like M+M

de_glm

If the experimental design is multi-factorial, for example if experimental factors such as starting growth condition, treatment, genotype, and replicated block exist, pairwise analysis is inadequate. Thus, the more statistically complex Generalised Linear Model based hypothesis testing functions of edgeR are required. This pipeline takes a keyfile describing the experimental design as the above pipelines do, however it takes and additional R script which describes the statistical model to be fitted, and contrasts within this model to be tested for differential expression. Tag-wise read counts are normalised and dispersions calculated with the GLM-based analogous of the functions used to do so in the de_pairwise pipeline. Then, a generalised linear model is fitted with glmFit, creating a glm object. Then, glmLRT is used to test each constant specified in the model script for differential expression. Analogous plots and tables to those produced in the de_pairwise pipeline are then produced.

3.3.3 Comparison of Differential Expression Pipelines

There was a large difference between the computational cost of four pipelines (aln_subread, aln_tophat, aln_tophat_htseq and aln_subread_htseq). Three measures of computational cost were measured: "real" time, "user" time and "sys" time. The "real" time describes the

time each pipeline took to complete. The "user" and "sys" time describe the CPU time spent running user code and performing operating system calls on behalf of user code respectively. Using an ANOVA model with Tukey's HSD post-hoc testing, significant differences in "real" time, "user" time and "sys" time were observed. As is shown graphically in Figure 3.2, the aln_subread is the fastest, taking an average of 3.48 \pm 0.10 minutes to complete, followed by the aln_subread_htseq pipeline, which took 4.88 \pm 0.07 minutes. The aln_tophat and aln_tophat_htseq were almost 400% slower, taking 12.9 \pm 0.12 and 13.77 \pm 0.07 minutes of real time respectively. The computational cost of user code and kernel processes in CPU-minutes followed similar patterns, as detailed in Figure 3.2. The computational time can become a limitations in very large and heavily replicated data sets needed to identify important effects of environment on common genetic variation.

Quantification of gene expression by the aln_subread pipeline is comparable to that obtained by the aln_tophat pipeline. As shown in Figure 3.3, there is a very tight relationship between counts produced by the Tophat2 and subread aligners. The slope of log(n+1) transformed raw count data when the model $tophatcounts \sim subreadcounts$ is fitted is 0.994, with p < 2e-16 and R^2 of 0.993. This indicates that there is approximately 0.7% statistical variation between these aligners. Table 3.2 illustrates the increased percentage of reads the aln_subread pipeline is able to align to both the entire genome and to the protein-coding transcriptome, when compared to the aln_tophat pipeline.

Add paragraph on edgeR analysis looking for DE between subread and tophat

3.3.4 Substantial reduction of RNAseq coverage is possible

While high-throughput sequencing has reduced sequencing costs dramatically, costs have stabilised in recent years (citation needed). Multi-

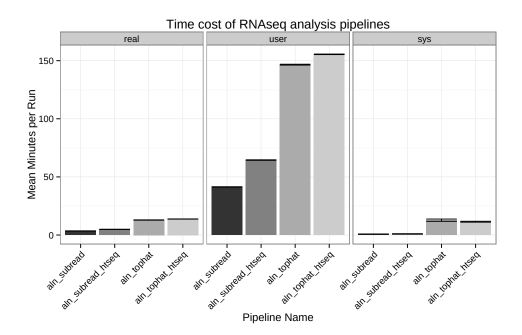


Figure 3.2: Computational cost of the aln_subread, aln_tophat, aln_tophat_ht-seq and aln_subread_htseq RNAseq analysis pipelines differs significantly. The "real" computational cost describes the number of minutes each pipeline took to complete. The "user" and "sys" metrics describe the number of CPU-minutes spent running user code (i.e. the pipeline components) and performing kernel operations on behalf of user code (e.g. input/output, memory (de)allocation and other system calls) for each pipeline execution.

Pipeline Name	Percentage	e Reads Mapped
r ipeime ivame	Entire Genome	Protein-coding Genes
aln_subread	99.43	98.99
aln_tophat	97.02	96.30

Table 3.2: Percentage of mapped reads to genome and transcriptome. The aln_subread pipeline is able to map a slightly higher percentage of reads to both the genome and to the transcriptome when compared to the aln_tophat pipeline.

Correlation of Count Data from Subread and Tophat2 Aligners

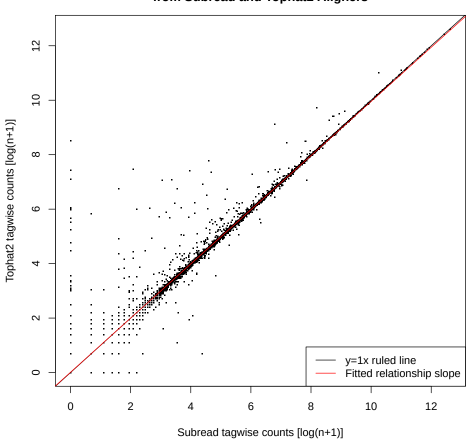


Figure 3.3: Comparison of gene-wise counts derived from the Tophat2 and subread short read aligners. A tight relationship is observed between count data from these aligners. If aligner had no effect, all points (genes) would fall exactly on the y=x line. There are off-diagonal points, however these occur mostly at low expression levels and are likely due to the required log transformation of count data.

plexing many samples per lane is important for high-throughput transcriptomics to increase replication to improve estimates of gene expression, and to increase sample breadth to better estimate transcriptome variation due to genetic or treatment effects. However, improvement to statistical power from increased sequencing depth diminishes as sequencing depth increases as the number of genes called as differentially expressed increases in a non-linear fashion with sequencing depth (Figure 3.4). Below a lower limit — 5 million reads — the number of genes called as differentially expressed reduces rapidly. This is concomitant with a decrease in number of genes whose expression is considered, and increases in biological coefficient of variation (data not shown). For the experimental system used in this experiment, I would recommend an optimal sequencing depth of approximately 5 million reads (or read pairs) to balance statistical power against additional sequencing cost. Budget permitting, more biological replicates or treatments should be included before sequencing is performed. These results have important implications for experimental design of current and future experiments underway.

may need a polish

better caption for Figure 3.4

3.4 Summary and Technical Discussion

summary of br chapter

In this chapter, I have implemented a framework allowing easier creation and optimisation of RNAseq analysis pipelines. Via literature review and brief experimental validation, I have selected software optimised for accuracy and speed in analysis of RNAseq data. Generic analysis pipelines that utilise these software have been created to allow investigation of common RNAseq experimental designs. Subtle transcriptional variation requires software to be optimised for accuracy and statistical

Depth vs Number of Differentially Expressed Genes

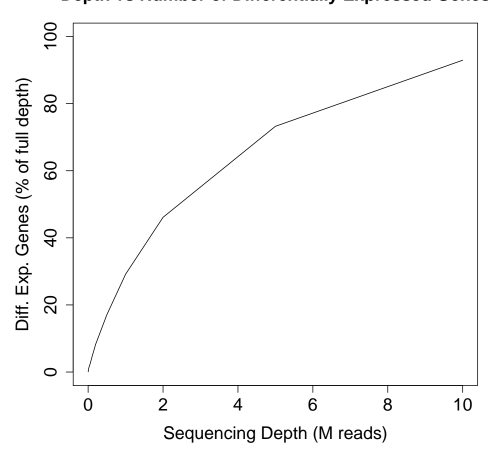


Figure 3.4: Decreasing sequencing depth per sample decreases the number of genes called as differentially expressed. This occurs because the total number of genes which can be examined for differential expression decreases in a similar fashion.

rigour, while optimising software for performance allows end users to conduct their own analyses, ensuring and end-to-end understanding of their experiments. Emphasis has been placed on reproducible analysis of RNAseq, ensuring the validity and reproducibility of results obtained. These advances allow for the examination of subtle transcriptome variation, such as between subtly different growth conditions.

CRUCIAL: needs linking sentence

Chapter 4

Transcriptome Variation Under Dynamic Growth Conditions

4.1 Background, Aims and Hypotheses

Transcriptional responses to abiotic stress have been studied by many authors (drought; pogson). However, theses stresses have rarely been studied in combination (Mittler 2006). This is despite evidence supporting the interactions between stresses (Atkinson and Urwin 2012; Mittler 2006). Additionally, many authors have observed responses to abiotic stresses encountered under field conditions that are not simple additions of the responses to individual stresses as elucidated under laboratory study (Atkinson, Lilley, and Urwin 2013; Atkinson and Urwin 2012; Mittler 2006; Seki et al. 2001; Wituszyńska et al. 2013). Together, these shortcomings warrant the investigation, under controlled laboratory conditions, of abiotic stresses in combinations which mimic those observed under field conditions.

To examine the effect of altered long-term light intensity in field-like combinations of stressful climatic variables such as temperature and humidity, dynamic growth conditions have been designed (see chapter 2).

In this chapter, I present preliminary transcriptomic study of plants grown under these dynamic conditions. Specifically, I aim to test if plants exposed to modest excesses of light either continuously (excess light dynamic growth conditions) or intermittently (fluctuating light) exhibit increased hardening to excess light when compared with sufficient light dynamic growth conditions. I hypothesise that this increased hardening would manifest as increased steady-state transcription of stressresponsive genes (Gordon et al. 2013; Wituszyńska et al. 2013), and a reduced induction of stress responsive genes upon exposure of plants to an 8x hot excess light treatment. More specifically, I expect that the excess and fluctuating light conditions will exhibit increased steady-state expression of stress-responsive transcripts compared with the sufficient growth condition, and with steady-stated expression observed in previous studies of plants grown under static growth conditions. Furthermore, I hypothesise that the fluctuating light condition will exhibit higher steadystate expression of stress-responsive transcripts, as plants are generally less able to acclimate to fluctuations in light intensity than to constitutive excess light (Alter et al. 2012; Gordon et al. 2013; Külheim, Ågren, and Jansson 2002).

clean this up per dad's com-

4.2 Methods

4.2.1 Growth and Harvesting of *Arabidopsis*

Elaborate on RIX lines

Initially, I had aimed to conduct a experiment which mapped expression QTLs under dynamic growth conditions. Thus, 80 A. thaliana Recombinant Inbred Intercross (RIX) lines (RIX line citation) were planted (see Appendix ??). Additionally, plants of reference accessions Cape Verde Islands (Cvi), Columbia (Col) and Landsberg erecta (Ler),

the photoprotection mutants stn8, pqr5, npq1 and npq4 were planted. All lines were planted in triplicate for each dynamic growth condition. Plants were grown in a carefully controlled manner to minimise variation in germination time and the developmental state of plants at a given time. Seeds were sown directly onto Debco Seed Raising Mix (Debco Pty. Ltd.), mixed with 3 grams per litre Osmocote® slow release fertiliser. Following sowing, plants were lightly watered from above and vernalised in a 4 °C cool room for three days. Plants were germinated under static growth conditions, of approximately 120 μ mol photons $m^{-2}s^{-1}$, 21 °C, with a 12 hour photoperiod. Following thinning, the remaining plants were established under these conditions for two weeks, before being distributed between three SpectralPhenoClimatron growth cabinets, with temperature, humidity and lighting controlled according to the Sufficient, 2x Excess and Fluctuating light dynamic growth conditions described in subsection 2.3.2. Plants continued to grow under these conditions until and beyond harvesting. Throught plant growth in the SpectralPhenoClimatron, high resolution images were captured at 20 minutes intervals.

> this really needs some cleaning, per dad's comments also

To assay plant response to hot excess light, plants were exposed to one hour of 8x hot excess light from sodium vapour and tungsten filament lamps in a Conviron growth cabinet. To enable study of transcriptomic responses to this assay, one fully expanded leaf was taken from each of the 576 surviving plants between 0 and 10 mintues before and after this excess light treatment. To do so, petioles were cut with clean scissors, and leaves gently rolled without crushing to facilitate placement in 96 well 1.2 mL deep well plates. Plates were kept in dry ice while harvesting occurred, and were transferred to -80 °C freezers for storage.

check this number

4.2.2 Unsuccessful generation of an eQTL mapping dataset

To increase the throughput and the decrease cost of Illumina RNAseq sequencing library preparation, I attempted to implement the RNAseq library preparation protocol of Kumar et al. (2012), a published protocol enabling the preparation of RNAseq libraries in high throughput using 96 well plates. Specifically, I used a slightly modified version of the High-Throughput RNAseq protocol described in Supplementary Methods 1 of Kumar et al. (2012) (hereafter referenced as the HTR protocol). To test and optimise this protocol, leaf tissue collected from surplus 5-week old A. thaliana Col-0 from a colleague's experiment was used. This tissue had been collected into Qiagen 1.2mL collection tubes containing a single steel ball bearing, and snap frozen in liquid N₂ before grinding in a TissueLyser (PN <+get details+>) for two one minute pulses at 25Hz at a later date. In collection tubes, 750 µLDynabeads Lysis/Binding Buffer was added, before sample was ground for a further 30s in a TissueLyser as before. Then, lysates were prepared per Steps 1.1.6-1.1.8 of the HTR protocol described in Supplementary Methods 1 of Kumar et al. (2012). Isolated mRNA was obtained and cDNA synthesised and fragmented according to steps 1.2 to 3 of the HTR protocol of Kumar et al. (2012). Working with Dr. Norman Warthmann, the remaining steps of the HTR protocol were validated using four cDNA samples, following the HTR protocol with modifications as described below.

To test all bell adaptors, sonicated genomic DNA was used as input material, due to it's similar size and fragmentation properties to fragmented cDNA, and due to the scarcity of large quantities of cDNA of little value. This DNA was obtained from *Oryza sativa* seedlings, diluted to a concentration of approximately $7\,ng\,\mu\,L^{-1}$ (500 ng in 70 $\mu\,L$) before sonication in a Diagenode Bioruptor DNA sonicator. This soni-

part numbers and small details like that are missing throughout this methods section

define bell adaptors

cated DNA was cleaned up per steps 3.5-3.8 of the HTR protocol, with the modification that $30 \,\mu L$ bead binding buffer and $40 \,\mu L$ Ampure XP SPRI beads were used. Then, a modified step 4 of the HTR protocol was used to create unamplified sequencing libraries. In step 4.1 and 4.2, double reactions were performed, however the same quantity of SPRI cleanup reagents were used. Before adaptor ligation, the A-tailed libraries were eluted using a mixture of 5 μL diluted adaptor oligonucleotide, 1 μL 10x ligation buffer and 2 μL water. The DNA ligase was diluted in the remaining $1.5 \mu L$ water and added to each reaction, before proceeding with protocol steps 4.3.3 onwards. The adaptors used were not those specified by Kumar et al. (2012), instead custom bell adaptors were used. These adaptors were designed by Dr Norman Warthmann, and are compatible with the T/A overhang ligation method Kumar et al. (2012) utilise. These adaptors are described in Table 4.1. The protocol described in step 5 of the HTR protocol was used, with the forward and index 1 reverse primers (see Table 4.1), to amplify the libraries.

need this data from Norman

RNA quality and quantity was assayed using the Agilent BioAnalyser digital electrophoresis system. The RNA samples were loaded into a Plant RNA Pico analysis chip and an analysis run per manufacturer's protocol. The effectiveness of various steps in this protocol was assayed by digital electrophoresis with the Shimadzu MultiNA instrument, using the DNA1000 kit. The pre-mix protocol was used: $2\,\mu L$ sample was added to $4\,\mu L$ DNA1000 marker solution, the solution mixed, and loaded into the instrument, which was run according to manufacturer's DNA1000-PreMix protocol. Quantitative PCR was performed to test the ligation and PCR efficiency of each adaptor. To do so, $2\,\mu L$ of each preamplification library was combined with $5\,\mu L$ Sybr Green qPCR master mix, $1\,\mu L$ each of the forward and index 1 reverse primers (see Table 4.1), and $1\,\mu L$ Uracil-Specific Excision Reagent (USER) enzyme mix.

4.2.3 RNAseq Library Preparation and Sequencing

Due to the failure of attempts to implement the high-throughput RNAseq protocol of Kumar et al. (2012) (briefly discussed in ??), a subset of all samples were selected for RNAseq analysis. Specifically, the samples of all Col-0 reference accession plants were used, as described in Table 4.2. Tissue of selected samples was extricated from 96 well plates into pre liquid N₂ cooled 1.5ml micro-tubes, ensuring plates remained frozen. Samples were ground using a Qiagen TissueLyser II for two one minute pulses at 25Hz, cooling racks in liquid N₂ between each pulse.

better sample table caption

Total nucleic acids (TNA) were extracted from samples using a commercial reagent (Trizol, Life Technologies,) immediately after grind- details ing. Total nucleic acids were extracted by adding 1mL of Trizol to each well-ground sample and shaking vigorously by hand, before adding 200 \mu Lchloroform and shaking by hand again. Samples were incubated for 3 minutes at room temperature, before centrifugation at 14000 rcf for 10 min in a chilled centrifuge. The TNA contained in the aqueous phase was re-extracted with chloroform and precipitated with 500 μ Lice cold isopropanol before incubation at -20 °C overnight. Total nucleic acids were precipitated by centrifugation for 20 minutes at 20000 rcf and 4°C , washed with 1mL 75% Ethanol, and resuspended in $50 \,\mu L$ RNase free 10 mM Tris-HCl. The quality of RNA in extracted TNA was assayed using the Agilent Bioanalyser digital electrophoresis platform. Samples were loaded into a Plant Nano analysis chip, and analysis run according to manufacturer's protocol.

RNAseq libraries were then prepared using the Illumina TruSeq V2 RNAseq Sample Preparation kit (Part number RS-122-2002). As previous studies in the Pogson Lab indicated DNAse treatment of TNA samples before RNAseq library preparation was not necessary (pers. comm. Peter Crisp, 2013; Supplementary Figure ??), TNA was diluted with 10mM Tris-HCl to $80\,ng\,\mu\,L^{-1}$ for use as input material. The manu-

Adaptor Name	Sequence
Test	CATCGATGC

 Table 4.1: Bell adaptors and index primers obtained from Dr. Norman Warthmann.

Sample Number	Accession	Growth Condition	High Light Treatment
1	Col-0	Sufficient	0h
2	Col-0	Sufficient	$0\mathrm{h}$
3	Col-0	Sufficient	0h
4	Col-0	Sufficient	$1\mathrm{hHL}$
5	Col-0	Sufficient	$1\mathrm{hHL}$
6	Col-0	Sufficient	$1\mathrm{hHL}$
7	Col-0	Fluctuating	0h
8	Col-0	Fluctuating	0h
9	Col-0	Fluctuating	0h
10	Col-0	Fluctuating	$1\mathrm{hHL}$
11	Col-0	Fluctuating	$1\mathrm{hHL}$
12	Col-0	Fluctuating	$1\mathrm{hHL}$
13	Col-0	Excess	0h
14	Col-0	Excess	$0\mathrm{h}$
15	Col-0	Excess	$0\mathrm{h}$
16	Col-0	Excess	$1\mathrm{hHL}$
17	Col-0	Excess	$1\mathrm{hHL}$
18	Col-0	Excess	$1\mathrm{hHL}$

Table 4.2: Samples analysed by RNAseq and qPCR. These samples are referred to by their sample numbers for the remainder of this thesis

facturer's protocol was then followed to produce RNAseq libraries, with modifications. RNA was fragmented by heating samples at the "Elution 2 - Frag - Prime" stage to 94 °C for 7 minutes, in place of the 8 minutes recommended by manufacturer guidelines to increase median insert size. To create cDNA, the SuperScript III reverse transcriptase (Life Technologies, part number 18080044) was used, and thus the incubation temperature was increased from 42 °C to 50 °C , per SuperScript III guidelines. During every SPRI clean-up step throughout the protocol, DNA bound to SPRI beads was washed with 180 μL ethanol rather than the recommended 200 μL , allowing all liquid to be effectively removed with a P200 multi-channel pipettors.

A pilot enrichment PCR was conducted with a subset of samples, enabling estimation of optimal cycle number for final enrichment PCR. To do so, quarter-volume PCRs ($12.5\,\mu L$ master-mix, $2.5\,\mu L$ sample) were run: samples 1 and 13 were run for 10 cycles, and 2 and 19 were run for 14 cycles, with the $60\,^{\circ}$ C annealing time extended to 45 seconds. Then, half-volume PCRs were used to amplify libraries for 12 cycles. Two libraries (samples 6 and 17) whose amplification failed with 12 PCR cycles were amplified using quarter-volume PCRs with 17 amplification cycles. These libraries were then cleaned up with SPRI beads, per TruSeq kit protocol. The success of these PCRs were assayed by digital electrophoresis, using the MultiNA instrument, as per ??, both before and after SPRI cleanup.

Final sequencing libraries were created by diluting and pooling RNAseq libraries. Libraries were diluted to 10nM, as calculated from MultiNA quantification. They were then quantitated fluorometrically using the Qubit 2.0 instrument (Life Technologies) with the dsDNA BR assay kit per manufacturer's protocol, and diluted to 5nM accordingly. These 5nM libraries were re-quantitated fluorometrically as above. Samples 1-12 and 13-18 plus 4 additional samples from a colleague were pooled to

equimolarity, forming two final sequencing libraries. Raw 100bp paired end sequence data was obtained by sequencing final sequencing libraries on two Illumina HiSeq 2500 sequencing lanes, performed at the Biomolecular Resource Facility, John Curtain School of Medical Research, ANU.

4.2.4 Computational Analysis of RNAseq Data

Raw Illumina paired-end 100bp sequence data was obtained as gzipped FASTQ files from the BRF. To gauge the quality of the obtained sequence data, several analyses were conducted. Firstly, the number of reads obtained from each library were calculated using the code shown in listing 1 below. Then, PHRED scores (defined as $-10 \log_{10}(P)$, where P is the probability of error at a given position) for each sequence base in each library were analysed using analysis pipelines described in subsection 3.3.2.

Analysis pipelines described in subsection 3.3.2 were applied to this dataset. Firstly, the aln_subread pipeline computed read counts genewise from raw short reads, using the kevin-hons-pairwise.key keyfile (see listing ??). Then, both de_pairwise and de_glm pipelines were used independently to test for differential expression, using the kevin-hons-pairwise.key keyfile and the kevin-hons-glm.key keyfile and edgeR-glm-kmhons.R GLM setting script respectively (see listings ??, 2, and ??). A general linear model (GLM) of the form ~ Group was fitted, according to the keyfile shown in listing ??. Statistical tests for differential expression between contrasts shown in listing ??.

stuff on including Pete's dataset

Rewrite analysis methods - lower priority. remove pair-

```
for fqfile in `find -name *.fastq.gz`
do
echo "$fqfile $(($(zcat $fqfile | wc -1) / 4))"
done
```

 ${\bf Listing~1:~Count~the~number~of~reads~in~raw~sequence~files}$

Group	Sufficient.Oh	Sufficient.Oh	Sufficient.Oh	Sufficient.1hHL	Sufficient.1hHL	Sufficient.1hHL	Fluctuating.Oh	Fluctuating.Oh	Fluctuating.Oh	Fluctuating.1hHL	Fluctuating.1hHL	Fluctuating.1hHL	Excess.Oh	Excess.Oh	Excess.Oh	Excess.1hHL	Excess.1hHL	Excess.1hHL
Treatment	Oh Su	Oh Su	Oh Su	1hHL	$1\mathrm{hHL}$	$1\mathrm{hHL}$	q0	QP	q0		1hHL	1hHL	Oh Exce	Oh Exce	Oh Exce	1hHL Ex	1hHL E	1hHL E
	Sufficient	Sufficient	Sufficient	Sufficient	Sufficient	Sufficient	Fluctuating	Fluctuating	Fluctuating	Fluctuating	Fluctuating	Fluctuating	Excess 0	Excess 0	Excess 0	Excess 1	Excess	Excess
GrowthCondition	1_index1	2_index3	3_index8	4_index9	5_index10	6_index11	7_index20	8_index21	9_index22	_10_index23	_11_index25	_12_index27	_13_index1	_14_index3	_15_index8	_16_index9	_17_index10	_18_index11
l Sample	Sample_BJP_K1_1_index1	Sample_BJP_K1_2_index3	Sample_BJP_K1_3_index8	Sample_BJP_K1_4_index9	Sample_BJP_K1_5_index10	Sample_BJP_K1_6_index11	Sample_BJP_K1_7_index20	Sample_BJP_K1_8_index21	Sample_BJP_K1_9_index22	Sample_BJP_K1_10_index23	Sample_BJP_K1_11_index25	Sample_BJP_K1_12_index27	Sample_BJP_K2_13_index1	Sample_BJP_K2_14_index3	Sample_BJP_K2_15_index8	Sample_BJP_K2_16_index9	Sample_BJP_K2_17_index10	Sample_BJP_K2_18_index11
Ordinal	-	O1	σ <u>.</u>	↔	10	62	2	ω 	O.	10	11	12	13	14	15	16	17	18
1	61	m	4	75	9		· ∞	6	10	11	12	13	14	15	16	17	18	19

Listing 2: The kevin-hons-glm.key keyfile

4.2.5 Quantitative Real-time PCR Quantification of Gene Expression

To assay expression of genes shown to respond to hot excess light in previous studies, quantitative PCR (qPCR) was used. RNA extracted for RNAseg analysis (see subsection 4.2.3) was treated with DNAse, to remove genomic DNA contaminants, using Turbo DNAse (Life Technologies). Approximately 15-20 μ gof RNA in 90 μ L was combined with 10 μL Turbo DNAse buffer and 1 μL Turbo DNAse, before incubation at $37\,^{\circ}\text{C}$ for 30 minutes, at which point an additional $1\,\mu\,L$ Turbo DNAse was added to sample. Samples were mixed, and incubated for a further 30 minutes at $37\,^{\circ}\mathrm{C}$. RNA was recovered by phenol-chloroform extraction, adding $100 \,\mu L1:1$ phenol-chloroform mixture to the DNAse reaction solution, mixing, separating phases by centrifugation for 10 minutes at 14000 rcf and 4 °C , and precipitating RNA by adding 200 $\mu\,L{\rm ice}$ cold isopropanol, incubating at -20 °C overnight, centrifuging to pellet RNA, washing RNA pellet with 70% ethanol and re-suspending RNA in 20 \mu LDEPC-treated MilliQ water. RNA quality was assayed by visualising denatured RNA on a 1% Agarose gel, prepared using buffers made with DEPC-treated water to prevent RNA degradation.

Complimentary DNA (cDNA) was synthesised using Invitrogen SuperScript III First-strand cDNA synthesis kit. RNA was diluted to $100\,ng\,\mu\,L^{-1}$, denatured at $65\,^{\circ}$ C for 5 minutes, and cDNA synthesis reactions consisting of $10\,\mu\,L$ denatured RNA sample, $1\,\mu\,L50\,nmol\,L^{-1}$ dT(18)VN primer, $1\,\mu\,L10\,mmol\,L^{-1}$ dNTPs, $2.5\,\mu\,L$ nuclease free water, $4\,\mu\,L5$ x reaction buffer, $1\,\mu\,L100\,mmol\,L^{-1}$ DTT, and $0.5\,\mu\,L$ SuperScript III enzyme. This reaction solution was mixed, centrifuged briefly, and incubated at $50\,^{\circ}$ C for 60 minutes, before enzyme inactivation at $70\,^{\circ}$ C for 15 minutes. Samples were then stored at $-20\,^{\circ}$ C until use.

Expression was quantified with qPCR using the Roche Sybr Gold master mix kit (part number 04-707-516-001) in a Roche LightCycler 480

tukey ref

thermocycler. Reactions of $10\,\mu\,L$ consisting of $1\,\mu\,L$ cDNA, $3.6\,\mu\,L$ nuclease free water, $0.2\,\mu\,L$ of each $20\,\mu\,mol\,L^{-1}$ primer, and $5\,\mu\,L$ Sybr Gold 2x master mix. Reactions were conducted in technical triplicate in 384 well plates sealed with qPCR-compatible plate seals. Expression of eight genes (APX2, ELIP1, ELIP2, LHCB1.4 and HSP90-like, as well as reference genes PP2AA3 and GAP) was quantified using primer sets described in Table 4.3 in 24 samples. These cDNA samples were samples 1-18 of the dynamic growth condition dataset described in Table 4.2, as well as cDNA samples prepared using identical methods as described above from plants grown under static conditions ($100\,\mu$ mol photons $m^{-2}\,s^{-1}$ light intensity, 12 hour photoperiod, 21 °C) for five weeks, before and after exposure to 10x hot excess light for one hour. In addition, template-less and reverse transcriptase free controls were conducted for some primer sets (HSP90-like and PP2AA3) to ensure absence of genomic DNA contamination.

Raw quantification curves were obtained by thermocycling qPCR reactions in a LightCycler 480 thermocycler. Reactions were heated to 95 °C for 10 minutes, before cycling between 95 °C for 30 seconds, 60 °C for 45 seconds, and 72 °C for 60 seconds for 45 cycles. A final incubation at 72 °C for 5 minutes was followed by a slow ramping of temperature from 45 to 95 °C with continuous quantitative analysis, to obtain a melting curve. Raw quantification data was analysed with LinRegPCR (Ruijter et al. 2009) to create N0 values, a statistically rigorous arbitrary unit of quantification suitable for relative quantification. Quantification relative to PP2AA3 was then calculated sample-wise for all primer sets analysed, using custom analysis code in R (see subsection 6.2.3). An ANOVA model was fitted to relative quantification, with Tukey's honest significant differences post-hoc testing to determine specific effects.

GC % Amplicon Size	146	105	~	161	_	136	134
A ½			73		87		
% DD	60.00	52.38 47.62	55.56 43.48	60.00	60.00	60.00	60.00 52.17
$\operatorname{Tm}_{\circ C}$	58.9 58.9	58.5 55.0	54.5 55.9	56.2	59.4 60.4	59.5 59.4	57.8 58.0
Length	20 23	21	18	20	$\frac{20}{20}$	20	20 23
Sequence	GCCGTTAGGCTTCTTGACCC GGCTCAACTTTGTCCAGTCTACC	TCGGAAGAATCGGTCGTTTGG TGTATGTCATGTACTCGGTGG	CCACCACAAATGCCACAG GCAAATCTCCAAACTTCGTACTC	CGACCAAGCGGTTGTGGAGA CACAATTCGTTGCTGTCTTT	TCCTCTCCTGCTTTGACCGG TTTGCGCATGGTGATTCGGC	AGATGCATGGCTGAGGGAGG AGTCGCTAAACTTTGTGCTCACC	CGTGTGGAGAGATCGAGTGG GCCTTCTTAGTCTCAGCCTTAGG
NCBI Acc.	NM_001035587.2	$_{ m NM_101214.3}$	$_{ m NM_117551.2}$	$NM_001035958.1$	$_{ m NM_128995.2}$	NM_113183.3	$_{ m NM}$ _128504.3
AGI	AT3G09640	AT1G13440	AT4G14690	AT1G13320	AT2G34430.1	AT3G22840.1	AT2G29500.1
Name	$\begin{array}{c} APX2_ej4_F \\ APX2_ej4_R \end{array}$	$rac{ ext{GAPC2} - 5 ext{T}}{ ext{GAPC2}}$	ELIP2_F_UPL101 ELIP2_R_UPL101	$\mathrm{PP2AA3}_{-3}$ 3'F $\mathrm{PP2AA3}_{-3}$ 3'B	$\begin{array}{cccc} \text{LHCB1.4} \\ \text{LHCB1.4} \\ \text{LHCB1.4} \end{array} \text{R}$	$ m KM_{ELIP1}_{EM}$ $ m KM_{ELIP1}_{R}$	${ m HSP20Fam_F}$ ${ m HSP20Fam_R}$

Table 4.3: QPCR primer sequences and characteristics.

4.3 Results

4.3.1 QTL mapping datasets

Many RIX lines exhibited very poor germination or survival rates, Because of this, only 40 to 60 RIX lines had at least one plant survive to harvesting, a reduction in statistical power for mapping.

paragraph about RIX mapping dataset. keep it brief.

4.3.2 Quantification of Transcriptome-wide Responses to Altered Light Intensity Under Novel Growth Conditions

Successful Preparation of RNAseq Libraries

RNA of suitable yield and quality for RNAseq library preparation was obtained. Figure 4.1 and Table 4.4 illustrate the high quality and yield of RNA samples. Illumina RNAseq libraries of the expected size and concentration were successfully prepared from all 18 RNA samples. Figures 4.2 and 4.3 demonstrate the successful creation of RNAseq libraries. Two library amplifications (Samples 6 and 17) failed to reach a suitable yield after 12 PCR cycles, so the PCR was re-run with 17 amplification cycles, which created libraries of a suitable concentration (Figure 4.3).

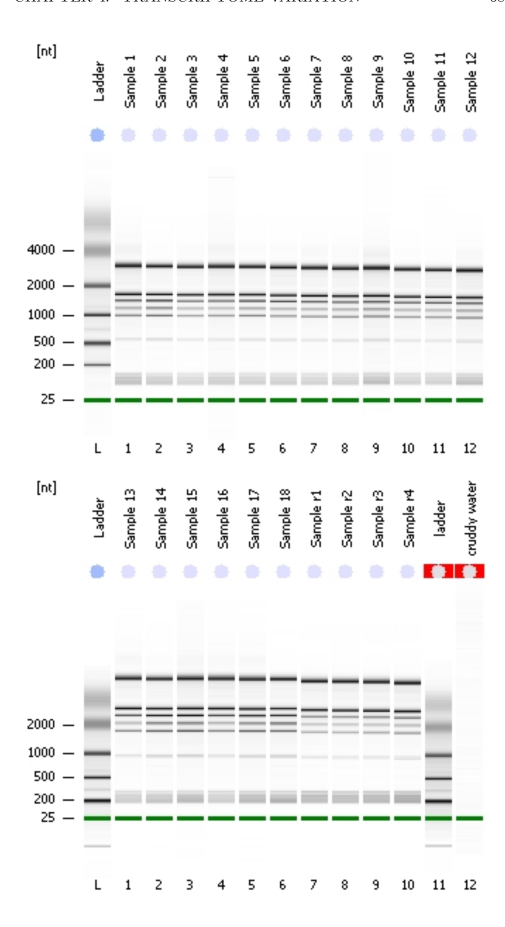


Figure 4.1: Bioanalyser 2100 digital electrophoretograms show expected rRNA peaks and mRNA smear, with no evidence of large-scale sample degradation. Samples 1-18 are numbered according to Table 4.2. The strong, clear bands at approximately 1900 and 3700nt are derived from the 18s and 25s nuclear rRNA species respecively (Babu and Gassmann 2011). The absense of a broad smear indicates minimal degradation of RNA has occured (Babu and Gassmann 2011)

Sample	Conc. $(ng \mu L^{-1})$	RIN
1	451	6.9
2	470	6.5
3	532	7
4	455	7.1
5	453	6.7
6	461	6.8
7	473	6.9
8	528	6.8
9	217	6.9
10	524	6.8
11	505	6.7
12	548	6.7
13	164	6.9
14	205	6.4
15	93	6.6
16	198	6.8
17	149	6.6
18	239	6.3

Table 4.4: RNA sample yield and RNA Integrity Number (RIN). RINs greater than 7 indicate high quality RNA, and RINs greater than 6 are acceptable (Babu and Gassmann 2011). Overall yield is sufficient, and all samples have a RIN of at least 6, indicating accepable quality.

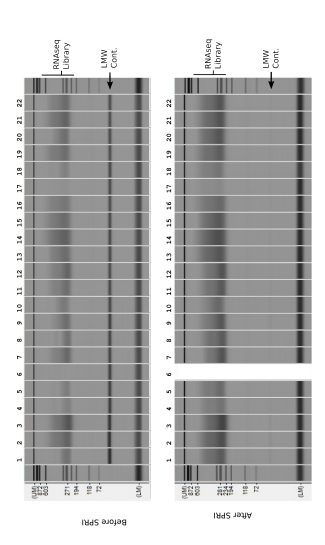


Figure 4.2: RNAseq libraries before (top) and after (bottom) final solid-phase revserible immobilisation (SPRI) cleanup. A broad smear of nucleic acid between approximately 200 and 600bp long is expected, ideally with a peak around 200-300b (illuminamanual). Additionally, after PCR, a band of low molecular weight (approximately 60bp) contaminant is expected, and observed. Clean-up with SPRI is expeceted, and obserbed, to remove this contaminant efficienty.

Obtaining a High Quality RNAseq Dataset

better word for obtaining

To quantify global gene expression accurately, it is important to ensure the quality of raw sequencing data is acceptable. Ensuring approximately even sequencing depth across libraries ensures sequencing depth does not confound differential expression analysis. As detailed in Table 4.5, library depth variation was within an order of magnitude across both lanes sequenced. The two lanes of sequencing yielded 153 and 143 million reads respectively, averaging 15.3 \pm 2.7 million and 11.9 \pm 1.0 million reads per sample respectively (means \pm standard deviation). Coverage in lane 1 is expected to be lower, as its overall coverage is lower and it contained 12 samples compared with 10 in lane 2. Low-level sequence quality as gauged by fastqc was high (Andrews 2012). The 25th percentile PHRED quality score exceeded 28 at every base in all sequence libraries before any quality control (see Figure 4.4). Following sequence quality control, the minimal 25th percentile PHRED score increased to 30. These basic statistics show the sequencing of RNAseq libraries was successful and the resulting data suitable for further analysis.

The very high proportion of sequence reads which aligned to the genome, and moreover to protein coding loci, is a further indicator of dataset integrity. Table 4.6 describes the sample-wise percentages of reads which map to the genome and transcriptome. Over 99 percent of reads map to the genome in all samples, the vast majority of which align to protein-coding genes (97-99%). Ribosomal RNA depletion during library preparation was successful, with between 0.08% and 1.6% rRNA contamination across all libraries.

4.3.3 RNAseq Reveals a Noisy Transcriptome

Once reads were aligned to the genome and gene-wise counts obtained, statistical assessment of differential expression was conducted. After

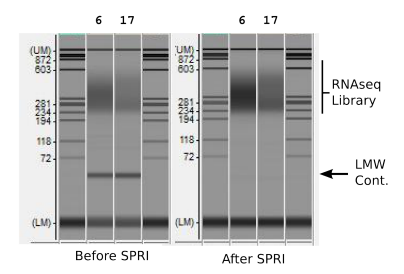
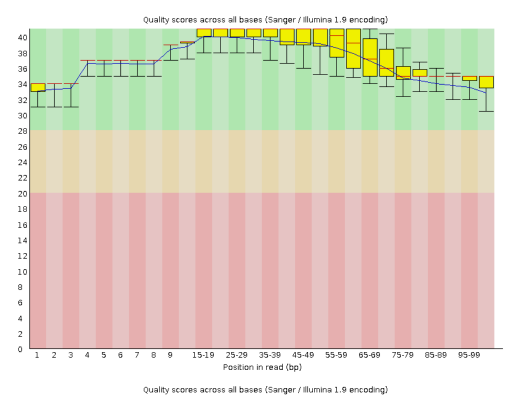


Figure 4.3: PCR amplification of samples that were not amplified after 12 PCR cycles. Note the presence of a band of between 200 and 600bp is expected, with a peak around 200-300b as expected, and as observed in Figure 4.2. Note also the efficient removal of low molecular weight contaminant after SPRI clean-up (right).

Library	Reads per Pre QC	Library (Millions) Post QC
1	11.43	10.78
2	10.66	10.12
3	10.82	10.37
4	10.79	10.41
5	12.39	11.86
6	14.08	13.44
7	12.78	12.28
8	11.58	11.20
9	12.83	12.31
10	12.47	12.10
11	11.25	10.89
12	11.92	11.51
13	17.98	17.31
14	19.35	18.49
15	17.94	17.16
16	14.31	13.84
17	14.11	13.56
18	11.58	11.03
19	12.72	NA
20	16.86	NA
21	15.87	NA
22	12.36	NA

Table 4.5: RNAseq library sequencing depth. Reads per library before and after quality control refer to the length of raw sequence data, and to the size of the libraries immediately before statistical analysis of differential expression, after sparse tags and non-protein coding loci were removed. Libraries 19-22 were sequenced on behalf of a colleague, and do not form part of this thesis, thus have not been analysed for differential expression.



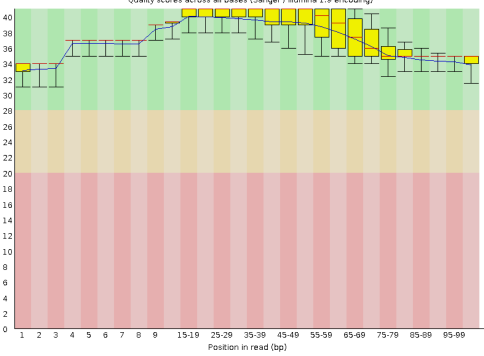


Figure 4.4: Per-base quality before (top) and after (bottom) quality control. These box-plots describe, per-position in each read (x-axis), quartiles and medians of PHRED quality score, which is related to probability of sequencing error at each position. PHRED scores of greater than 30 are considered good, and scores are expected to be lower towards the 3' end of read sequences, due to sequencing technology (andews_fastqc_2013)

C 1. N 1 .	Percentage Reads Mapped to:		
Sample Number	Entire Genome	Protein-coding Genes	
1	99.38%	97.59%	
2	99.15%	98.10%	
3	99.64%	98.60%	
4	99.59%	98.92%	
5	99.10%	98.48%	
6	99.16%	98.26%	
7	99.61%	98.82%	
8	99.53%	99.07%	
9	99.62%	98.84%	
10	99.59%	99.24%	
11	99.62%	99.14%	
12	99.60%	98.94%	
13	99.28%	98.88%	
14	99.16%	98.88%	
15	99.29%	98.70%	
16	99.31%	99.09%	
17	99.10%	98.86%	
18	99.23%	98.59%	

Table 4.6: Percentage of mapped reads to genome and transcriptome. These figures reiterate the successful creation of the RNAseq dataset, with very low (<4%) rates of contamination by non-protein coding RNAs or genomic DNA. This imparts confidence that any short read counted towards quantification of the expression of a protein coding gene is derrived from mRNA transcribed from that gene.

dataset filtering to remove lowly expressed or non-protein coding loci, 17948 loci remained. Figure 4.5 illustrates the high biological coefficient of variation across the majority of genes. This is evidenced by the spread of points upwards in samples from the dynamic growth condition dataset when compared to those from the static growth condition (RRGS) dataset described in subsection 3.2.1. The overall biological coefficient of variation was also much higher in the dynamic growth condition dataset (common BCV = 0.493), when compared to the RRGS dataset (common BCV = 0.128). It is crucial to note that the RRGS experiment is not a control for the dynamic growth condition experiment, but is still useful an external comparison. The high variation between replicates causes reduced statistical power to detect differential expression (Robinson et al. 2013).

The high biological variance in expression is also demonstrated via multiple-dimensional scaling, an unsupervised clustering algorithm that describes the transcriptome-wide similarity of samples. Samples grown under dynamic growth conditions have a higher scatter about both axes of the multiple-dimensional scaling plot when compared to the RRGS dataset (Figure 4.6). Replicates often cluster less closely than treatments, however, upon treatment with one hour of hot excess light, plants grown under both static and dynamic growth conditions exhibit similar patterns (in Figure 4.6, pre and post-treatment samples separate along a axis 1). It is also important to note that replicates within the RRGS dataset cluster together tightly compared with replicates from the dynamic growth condition dataset, a hallmark of the RRGS dataset's lower biological variance. Whist the variability observed in the dyanmic growth condition dataste is concerning, promising qualitative trends transcriptome-wide patterns of differential expression warrant further — albeit cautious investigation of this dataset.

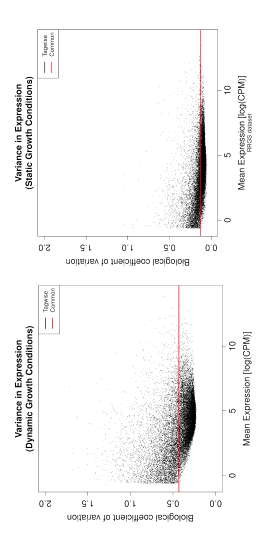


Figure 4.5: Biological coefficient of variation in dynamic (left) and static(right) growth conditions. Here, each point represnts a gene, and the relationship between expression level (x-axis) and gene-wise biological coefficients of variation is plotted. Additionally, a common (analysis-wide) measure of the biological coefficient of variation is ploted (red line), for samples from the dynamic and static growth condition datasets, this equates to 0.493 and 0.128 respectively. The higher biological variance of the dynamic growth condition dataset when compared to the static growth condition (RRGS) dataset is evidenced by a more positive spread of gene-wise biological coefficients of variation, and by a higher common biological coefficient of variation.

mention distance between samples in mds

fix fig sizes

Differential expression between dynamic growth conditions

Eight group contrasts were tested for differential expression. These contrasts test the effect of light intensity within the framework of dynamic growth conditions on both the steady-state transcriptome, and on the transcriptional response to exposure to hot excess light for one hour. A summary of statistically significant differential expression is shown in Table 4.7 and ??. Small numbers of differentially expressed genes were observed between steady-state transcription in the sufficient growth condition and the excess and fluctuating growth conditions, but no differential expresison was detected between the fluctuating and excess growth conditions. A transcriptional response to one hour of hot excess light was observed in plants grown under all growth conditions, however plants grown under excess light dynamic growth conditions showed the greatest number of differentially expressed genes in response to this treatment, followed by fluctuating and sufficient light conditions, in direct confilict with hypotheses. Tests for interaction between growth condition light intensity and treatment effect showed little or no significant differential expression Table 4.7. The differential expression observed between dynamic growth contiitions is in contrast to the 3195 up-regulated and 3146 down-regulated genes differentially expressed after one hour of hot high light treatment in plants grown under static growth conditions (the RRGS dataset described in subsection 3.2.1).

Multi-dimensional Scaling of Static Growth Condition Samples

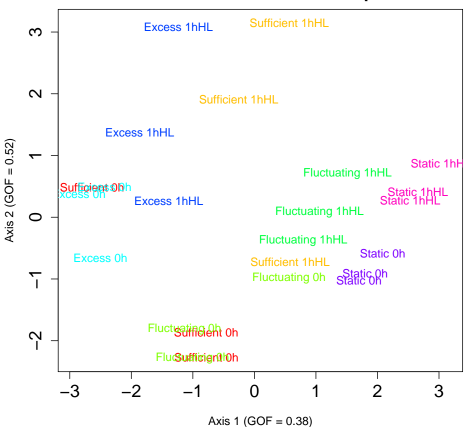


Figure 4.6: Multi-dimensional scaling of dynamic and static growth conditions. Replicate samples are represented in the same colour. Pre-hot excess light samples (0h) tend towards negative values on axis 2, while post-hot high light samples (1hHL) tend towards positive values on axis 2. This trend is preserved across samples from plants grown under both dynamic and static growth conditions. Replicates within the rapid recovery gene silencing dataset (Static 0h and Static 1hHL) cluster together tightly compared with replicates from the dynamic growth condition dataset (Excess, Fluctuating and Sufficient 0h and 1hHL). Note the meaning of axes is arbirary; they are pseudovariables that selected to seperate samples based upon log-fold-changes between samples. Goodness of Fit (GOF) indicates that together, these two axes account for 90% of variation between samples, indicating that the majority of difference across entire transcriptomes between samples is described within this plot.

Contrast Name	Contrast Name Contrast Description	Genes Downregulated Genes Up-regulated	Genes Up-regulated
Exc0-Suf0	Excess Light vs Sufficient Light	4	80
Flu0-Suf0	Fluctuating Light vs Sufficient Light	154	217
Exc0- $Flu0$	Excess Light vs Fluctuating Light	0	0
Suf1-Suf0	Sufficient Light pre- vs post-hot excess light	11	140
Exc1-Exc0	Excess Light pre- vs post-hot excess light	109	426
Flu1-Flu0	Fluctuating Light pre- vs post-hot excess	94	201
	light		
Exc01h-Suf01h	Interaction between Fluctuating light and	0	0
	hot excess light treatment		
Flu01h-Suf01h	Interaction between Sufficient light and hot		
	excess light treatment		

 Table 4.7: Summary of Differential expression

table captions

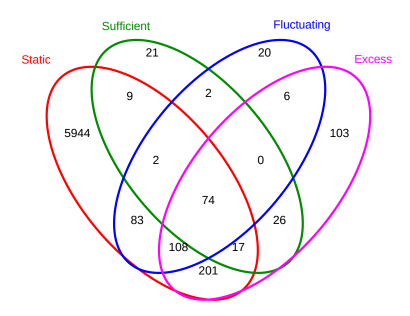
To gain biological insight from patterns of differential expression, gene ontology (GO) term enrichment analysis was used. Statistically significant enrichment of GO terms in genes up- and down-regulated in comparisons of steady-state expression and in transcriptional response to hot excess light. The GO terms enriched in genes up-regulated on exposure to hot high light were highly conserved across all dynamic growth conditions and the RRGS dataset grown under static growth conditions (Figure 4.8). Specifically, terms including 'response to heat', 'response to high light intensity', 'response to hydrogen peroxide' and 'response to jasmonic acid stimulus' are among the most statistically over-represented genes induced by one hour of hot excess light in plants from all dynamic conditions. Moreover, these terms are also amongst the most statistically over-represented genes induced by one hour of hot excess light in plants grown under static growth conditions. Table Table 4.8 describes GO terms that are over-represented in genes induced by hot excess light treatment across all condtions; terms involved in biotic and abiotic stress represent the majority of the 37 such terms. Full details of the 30 most significantly enriched GO terms in all differential expression tests are described in appendix ??.

another table catption

4.3.4 Expression Patterns of Excess Light Marker Genes

Quantitative RT-PCR (qPCR) was used to examine expression patterns of excess light marker genes. Comparisons of both steady-state expression, and induction of expression upon treatemnt with hot excess light have been examined. APX2, a gene induced by oxidative stress and excess light, was upregulated in sufficient, excesss and fluctuating light dynamic light conditions (ANOVA, F=5.63, p=0.0072, 3 degrees of free-

Similarity in response to excess light between static and dynamic growth conditions



Genes differentially expressed in resonse to treatment with hot excess light for 1 hour, by dynamic growth condition (FDR<0.05)

Figure 4.7: Extent of transcriptional response to a one hour treatment with hot excess light. Similarity between between sufficient, excess and fluctuating light dynamic growth conditions, and a similarly treated dataset grown under static growth conditions (from the RRGS dataset discussed in subsection 3.2.1).

Similarity in GO term enrichment in transcriptional response to excess light

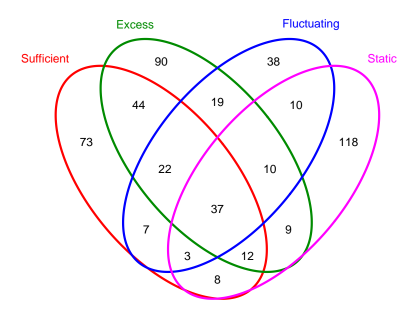


Figure 4.8: Similarly in GO terms between genes whose expression is induced by one hour of hot excess light in plants growth in Sufficient, Excess or Fluctuating dynamic growth conditions, as well as plants grown under static growth conditions (from the Rapid Recovery Gene Silencing dataset. A core set of GO terms is commonly over-represented in genes differentially expressed in all conditions (see Table 4.8).

Gene Ontology Term

abscisic acid mediated signaling pathway anthocyanin-containing compound biosynthetic process cellular response to heat

endoplasmic reticulum

endoplasmic reticulum lumen

gibberellic acid mediated signaling pathway

heat acclimation

hyperosmotic salinity response

jasmonic acid biosynthetic process

jasmonic acid mediated signaling pathway

jasmonic acid metabolic process

membrane

oxygen binding

protein disulfide isomerase activity

protein folding

response to abscisic acid stimulus

response to auxin stimulus

response to bacterium

response to cold

response to desiccation

response to endoplasmic reticulum stress

response to ethylene stimulus

response to fungus

response to gibberellin stimulus

response to heat

response to high light intensity

response to hydrogen peroxide

response to jasmonic acid stimulus

response to karrikin

response to osmotic stress

response to salt stress

response to symbiotic fungus

response to water deprivation

response to wounding

sequence-specific DNA binding transcription factor activity

signal transduction

transport

Table 4.8: Gene Ontology terms signficantly enriched in gene differentially expressed after treatment with one hour of hot excess light in all dynamic growth conditions and static growth conditions

dom, with Tukey's honest significant differences demonstrating significant pairwise differences between expression in sufficient, excesss and fluctuating light dynamic light conditions and in static grwoth conditions with p<0.05). Steady-state expression of LHCB1.4, a photosynthic gene known to be down-regulated by exceess light, was downreulated in plants grown under dynamic growth conditions, with statistically significant down-regulation observed between excess and fluctuating gdynamic light conditions and static growth conditions (ANOVA); no statistically significant effect of hot excess light treatment was found. Expression of ELIP1 is significantly upregulated in excess and fluctuating dynamic light conditions compared to sufficient and static growth conditions. Similar patterns can be qualitatibely observed in ELIP2, however high variance prevents statistical significiance. These patterns of differential expression are summarised in Figure 4.9.

anova stats for LHCB and other genes

4.4 Summary of Findings

In this chapter I present findings obtained from a preliminary dataset examining the transcritptional repsponse to altered light intesnity within the framework of dynamic growth conditions. An RNAseq dataset characterised by high levels of biologogical noise was created; within this dataset differential expression was observed. Upon treatment of plants grown under sufficient, excess or fluctating light dynamic growth conditions with one hour of hot excess light, differential regulation of gene classes involved in abiotic stress response was observed. Comparisons between sufficient, excess and fluctating light dynamic growth conditions elucidated limited differential expression. Gene classes including genes involved in translation, biotic and abiotic stress response and metabolism were significantly enriched in genes differentially expressed between these dynamic growth conditions.

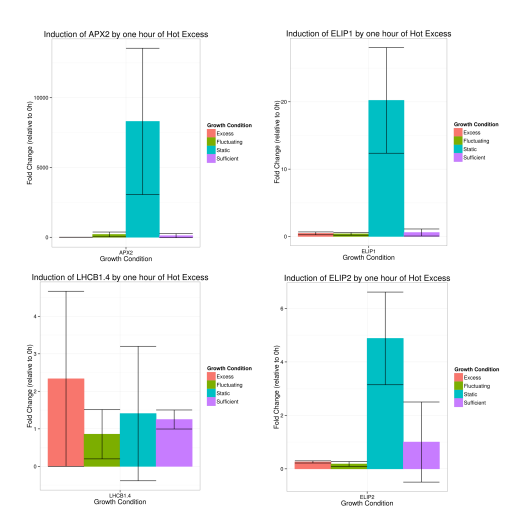


Figure 4.9: Differential expression of by

Chapter 5

Discussion

The study of plant-environment interactions is crucial to improving agricultural yield and understanding functional plant ecology in a changing climate. The study of mechansisms by which plants tolerate or respond to abiotic stresses is of particular importance, as abiotic stresses cause many hundreds of billions of dollars of crop losses and incalculable environmental damage (Mittler 2006). Several recent reviews of abiotic stress research have suggested a changed focus for research, away from studying stresses individually towards more holistic combined stress studies (Mittler 2006; Mittler and Blumwald 2010). Plants in ecological or agricultural settings commonly experience environments unfavourable to optimal growth, composed of multiple abiotic (and biotic) stresses (Boyer 1982; Mittler 2006). Moreover, natural environments are dynamic, and challenge plants to rapidly adapt to heterogenious environmental conditions on a daily basis. Laboratory study of these stresses should therefore account for recent findings suggesting that interactions between abiotic stresses in field conditions are not simply additive, rather are influenced by the dynamic combinations of stresses their environments impose upon them (Atkinson and Urwin 2012; Mittler 2006).

This thesis has developed methods to reproducibly induce abiotic stresses in laboratories in combinations analogous to those in which they can appear naturally and assay plant growth under such conditions, analyse plant response to responses, and presents a preliminary proof-ofconcept transcriptomics dataset analysing light response in the context of combinatorial abiotic stresses.

5.1 Novel Dynamic Growth Conditions

Plant biologists often conduct laboratory experiments in controlled growth facilities. These facilities may lack the ability to mimic characteristics of their natural environment, as they are rarely built from hardware enabling an investigator to do so. While static growth conditions used in labotorires are sufficient to uncover core mecchanisms of plant resposnes to stress, this is not always the case. In fact, evidence from field studies has uncovered phenotypes that are cryptic under static laboratory conditions, and that may be observed and studied closely under dynamic laboratory condtions (Külheim, Ågren, and Jansson 2002; Mishra et al. 2012; Mittler 2006; Wituszyńska et al. 2013). By creating software which connects existing but disparate technologies, I have enabled the implementation of growth conditions which can mimic regional climates or weather. The combination of computer controllable growth cabinets fitted with multi-spectral LED arrays allow for experimentation to consider important parameters that vary in natural regional climates, including temperature, humidity, and light quality and quantity.

In creating dynamic growth conditions, I aimed to combine some elements of the climates experienced by plants in nature with the reliability, reproducibility and convenience of laboratory growth conditions. Diurnal trends in light intensity, temperature and humidity follow those observed in the recent past at the model location (Bureau of Meteorology 2013). However, daily minima in temperature and maxima in humidity are tempered by hardware limitations. Similarly, daily integrated light intensity is lower due to limitations in the brightness of LED arrays used in the SpectralPhenoClimatron. The light produced by LED arrays

does not cover the photosynthetically active portion of the visible and adjacent spectrum with an even intensity of light per wavelength as sunlight does, although LED arrays do so with broader peaks of increased spectral intensity than fluorescent or incandescent lamps. These are important limitations to the SpectralPhenoClimatron, and addressing them in future work is a priority. Circannual variation in temperature and humidity follow similar climactic trends as the historical mean observations of weather at the trial location, with temperature slowly increasing, humidity decreasing and photoperiod lengthening, however light intensity does not gradually increase in the same manner as the observed climate.

While the dynamic growth conditions implemented in the SpectralPhenoClimatron mimic elements of natural environments, they are not designed to be accurate representations of conditions encountered in nature. In nature, weather provides a layer of stochasticity upon the broad trends in climate (citation needed). However, the developments I have described in this thesis facilitate the emulation of stressful growth environments which approximate weather-induced abiotic stresses plants experience in the natural environments. The system I have developed is not only applicable to studies discussed in this thesis, and has been used in experiments beyond the scope of this thesis including virtual recircoial transplants and gene-by-environment interaction QTL mapping. The study of standing genetic variation within regional climates allows links between genotype and reaction to some environmental parameter, and can shed light on mechanistic links via methods including GWAS (Brachi, Morris, and Borevitz 2011; Li et al. 2010; Li et al. 2006). However, when modelling plant growth in, or reaction to, environmental conditions, the system I have developed allows for non-stochastic weather to be imposed over the climatic trends (e.g. the fluctuations in light modelling intermittent cloud cover used in this thesis).

mention ecology, "virtual reciprocal transplants" etc

reiterate reliability, and make it first person

CRUCIAL- need a link

5.2 Improved analytic methods for RNAseq

RNAseq is a precise method to quantify genome-wide expression (Wang, Gerstein, and Snyder 2009). RNAseq can reveal hidden phenotypes and subtle environmental effects of a plant's growth environment (Martin et al. 2013), giving insights into development, regulatory mechanisms, signaling pathways, acclimation and many other aspects of plant biology. It is sufficiently sensitive to measure the subtle differences in expression between both closely genetically related organisms (à la expression QTL mapping, Sun and Hu (2013)), and subtle envronmental effects (e.g. differential response to dynamic growth conditions).

Reproducibility, accuracy, and performance of the computational analysis of any dataset is crucial. Inaccurate and poorly reproducible analyses have lead to embarrassing errors and retractions in many fields (Herndon, Ash, and Pollin 2013; Peng 2009). Computational performance of analysis software is central to interactive analysis of datasets; enabling fast analysis of datasets allows researchers to explore their data without the requirement for expensive clusters or supercomputers. The design and implementation of the RNAseq analysis framework I have created specifically address reproducibility, accuracy, and performance, by selecting the latest advances in high-performance, accurate analysis software, and providing a simple structure to an analysis that allows exact reproduction of the entire analysis.

The specific algorithms used in analysis of RNAseq datasets is a field of active technical research. Validation of pipelines designed around the Subread aligner (Liao, Smyth, and Shi 2013b) reiterate the software authors' claims of increased speed and accuracy. A recent publication suggests superior performance of the trimmed mean of M values normalisation method proposed by Robinson and Oshlack (2010) and implemented by Robinson, McCarthy, and Smyth (2010) compared to its competitors

(Rapaport et al. 2013). However, this review did not consider recently published statistical techniques such as QLspline (Lund et al. 2012). Further investigation of the state of the art in RNAseq statistical and computational analysis software is of great importance to studies of differential expression, and the framework presented in this thesis is specifically designed for modularity, enabling substitution of components for improved versions or alternatives with minimal effort.

mention txome subtlety

When transcriptome variation within a study system is subtle, sensitive and accurate methods are required to glean information. includes accurate analysis software (see above) and optimal experimental design. I examined the effect of sequencing depth on power to detect differential expression, and found that below 5 million reads per sample, power to detect differential expression with three replicates diminishes rapidly (see Figure 3.4). Several recent studies and reviews suggest that the current informal standard of RNAseq experimental design, which emphasises sample sequencing coverage over replication, is unwise. Rapaport et al. (2013) find increasing the level of replication over sequencing depth yields more differentially expressed genes, while Kliebenstein (2012) demonstrate that, even when sequence coverage was very low, nearly all expression QTLs could be mapped. In light of these data, my analysis of sequencing depth is conservative, as it does not consider the increases in replication made possible by increasing the number of samples sequenced per unit cost by a factor of two or more.

> Biologial variation control and multi environment treaments are needed to provide specificity of changes. Gene Atlas idea.

5.3 Elucidating Response to Light Intensity Under Dynamic Growth Conditions

The transcriptomes of *Arabidopsis* grown under dynamic growth conditions have been observed. A preliminary RNAseq dataset characterised by high levels of biological noise indicated differential expression of hundreds of transcripts between growth conditions and in response to treatment with hot excess light. QPCR analysis showed increased steady-state expression of know stress responsive genes under dynamic growth conditions, and reduced fold-change induction of stress genes upon application of hot excess light. Together, these data indicate patterns of differential expression observed under dynamic growth conditions, and provide limited and tentative support for the hypothesised "hardening" of plants to excess light.

The large amount of biological noise and resulting lack of power to detect differential expression is likely caused by a combination of factors. A rapid (within minutes) reduction in transcript abundance of hot excess light induced genes upon removal from hot excess light treatment has been observed (pers. comm. Peter Crisp). In light of these data, the high temporal error in sampling which exists in the dataset that I created may explain some of the biological variability. Additionally, as plants were of an advanced age (5 weeks) when samples were taken rapid estimation of the particular leaf that was samples was infeasible, thus the largest expanded leaf was taken. This variation in leaf number may be an additional reason for the high variability in this dataset, as the transcriptome is dependent on leaf developmental stage (Carmody 2013; Gordon et al. 2013). A final possible explanation for the high biological noise is variation in harvesting time. While plants were harvested as quickly as possible, each replicate took 3 hours to harvest. Previous studies have found circadian effects to both the general transcriptome

this paragraph needs a lot of thought, maybe move this one to summary of ch4, and rewrite the one that ends up here (Covington et al. 2008; Ptitsyn 2008) and in transcriptional response to biotic stress (Wilkins, Bräutigam, and Campbell 2010). The harvesting techniques and experimental design utilised in this study were suited to experiments that mapped eQTLs for stress-reponsive genes. For simple detection of differential expression however, they were not optimal and likely contributed to the high level of biological noise in the obtained dataset. Ideally, true internal controls grown under static growth conditions, and higher levels of biological replication would be more appropriate. These shortcomings result in reduced statistical power, and are a possible cause for the lower than expected number of genes whose expression was induced or repressed upon treatment with hot excess light.

Given this high level of biological noise, all further analyses of differential expression should be approached with caution, even if statistical techniques are sufficiently advanced to allow detection of differential expression. Despite this, patterns of expression similar to those observed by other authors have been noted in this study. Hundreds of genes were differentially expressed in response to one hour of excess light in each dynamic light condition, a similar magnitude to previous examinations of similar stresses (Gordon et al. 2013; Kimura et al. 2003; Rossel, Wilson, and Pogson 2002; Rossel et al. 2007). Specifically, the induction of heat shock family proteins observed by Rossel, Wilson, and Pogson (2002) was also observed here. Additionally, gene ontology (GO) term analysis indicates commonality of the transcriptional response to hot excess light with heat and oxidative stresses, as observed in previous studies (Rossel, Wilson, and Pogson 2002). Together, these data suggest that the response of plants grown under dynamic light conditions to hot excess light is similar to that of plants growth under static light conditions, albeit with somewhat tempered induction of hot excess light-resposive marker genes. An understanding of the power needed and variation present in a typical experiment has been identified.

Limited transcriptomic evidence of the effect of light intensity within the framework of dynamic light conditions was observed. Gene ontology (GO) term enrichment provides evidence that plant response to hot high light is preserved in plants grown under dynamic growth conditions hypothesised to induce acclimation to high light or fluctuations in light intensity (Figure 4.8). Additionally, GO term enrichment analysis details the overlap and interaction with other abiotic stresses. GO terms enriched in genes with differential steady-state expression in plants grown under sufficient light dynamic growth condititons and excess or flucating light dynamic grwoth condtiions exhibit limited overlap with those observed in previous studies of acute changes in light intensity, as hypothesised. However, this evidence of hardening is contrasted by the number of genes differentially expressed in response to hot high light under dynamic growth conditions. If acclimation to modest excesses in light intensity result in reduced fold-change induction of stress-inducible genes, as qPCR quanitification of APX2 expression suggest, then either the relative order of hardening hypothesied is incorrect, or the level of biological noise in this dataset prevented observation of such phenonmena.

fix up go paragrahph

add model of mop vs bucket

A more detailed analysis of this dataset may be warranted by these findings. In particular, examinaiton of possible harvesting or treatment block effects may, if any effect exists, reduce the biological coefficient of variation (Robinson et al. 2013). A detailed analysis of all genes called as differentially expressed may be of use, as high biological noise may not affect the estimates of differential expression in all genes. I hypothesise that upon doing so, genes previously found to be differentially expressed between lab and field growth conditions would have differential steady-state expression between sufficient, fluctuating and excess light conditions, with expression being highest in the excess and lowest in sufficient

light condititions.

ensure expt. methods stuff here makes it into ch4 meth-

Study of the transcriptome is an established method to assess the responses of plants to their environment, but it is far from the only method of doing so. Analysis of chlorophyll fluorescence of plants acclimating to fluctuating light demonstrated up-regulation of NPQ (Alter et al. 2012; Gordon et al. 2013), and study of such characteristics under dynamic light conditions is ongoing, but beyond the scope of this thesis. Analysis of accumulation of stress metabolites has elucidated metabolites involved in acclimation to combined stresses in field conditions (recent jansson paper), warranting similar analysis of metabolites in plants grown under dynamic growth conditions. However, transcriptomics remains a sensitive, accurate and useful tool to measure plants' responses to their environment.

5.4 Discussion of data in broader context of abiotic stresses

JB:This show the extension of the environment and the experiemntal design and the ability to stack layers of phenotype data. This is called a 'systems biology' approach.

better title for this section

While the effect of altered light intensity on plants and their transcriptomes has been studied in depth, few authors have studied under laboratory conditions analogous to those to which plants are adapted. _

intro to section interpretation discussion

Mutants in pathways critical to survival or fecundity of plants in ecological or agricultural settings may show little or no detrimental phenotype under benign laboratory conditions (Külheim, Ågren, and Jansson 2002). This further underlies the importance of examining the physiological, metabolomic and transcriptional responses of plants to field-like combinations of stresses. By examining the responses of plants to their environment in conditions similar to those which they have evolved, and studying stress response to field-like combinations of stresses, it may be possible to obtain a more direct picture of the role of parts of the genome

with little function in the artificially benign conditions under which many laboratory experiments on plants are conducted.

this could do with more refs

Studying the effect of "hardening" on the transcriptional response of plants to stress may reveal mechanistic insights of response genes. As plants acclimate to altered growth conditions, alteration of steady-state transcription often occurs (Alter et al. 2012; Heinrich et al. 2012; Hihara et al. 2001). Transcriptome profiling during acclimation to cold stress, (Chawade et al. 2007; Fowler and Thomashow 2002), excess light (Gordon et al. 2013; Page et al. 2012), and drought (Ding, Fromm, and Avramova 2012) reveals a transcriptional response to long-term stress exposure. The elucidation of genes underlying such acclimatory responses would yield insights into mechanisms by which plant tolerance of chronic abiotic stress could be improved.

need more refs and less fluff for this paragraph

Scientists have repeatedly developed crop lines tolerant to stress assays "in the lab" that, when trialled under agricultural condition, either do not show stress tolerance or have increased susceptibility to other stresses or combinations of stresses (Atkinson and Urwin 2012; Mittler 2006; Mittler and Blumwald 2010; Wituszyńska et al. 2013). Some of the difficulty in translation of stress tolerance from lab to field can be explained by detrimental interaction of stresses (Mittler and Blumwald 2010). By performing genetic screens or genome wide association studies for stress tolerance under dynamic growth conditions, which mimic field-like combinations of stresses, it may be possible to elucidate mechanisms of tolerance to specific stresses which do not conflict with mechanisms of tolerance to other abiotic stresses, as tolerance is assayed in a mildly stressful environment that mimics that encountered under agricultural conditions.

first sentence needs rewording

5.5 Future Directions

The emergence of transcriptional patterns despite a dataset with high noise warrants further investigation. Brief examination of fold-change values indicates that biological noise is the primary factor for reduced differential expression. Utilising recent research on the underlying biology of transcriptional response (Peter Crisp, unpublished data) and RNAseq molecular methodology (Kumar et al. 2012) and experimental design (Rapaport et al. 2013), a repeated experiment which incorporates faster sampling and more accurate application of hot excess light stress, along with many more biological replicates (5-8 replicates) per condition, may further elucidate the patterns of differential expression under dynamic growth conditions, and determine the effect of light intensity on the transcriptome under dynamic light conditions.

rework this a bit

Expression QTL mapping presents an opportunity to simultaneously discover novel transcriptional patterns, and their underlying regulatory mechanisms (Keurentjes et al. 2007). Samples to perform a limited eQTL mapping experiment were collected but not processed due to constraints on time and statistical power. Once genetic material suitable for such analyses has been obtained, successfully conducting an analysis to find gene-by-environment interactions in and eQTLs controlling differential expression of transcripts in response to altered light intensity within the framework of dynamic growth conditions would elucidate not only response mechanisms, but their underlying regulators. This information could be used by crop improvement programs to select for lines with improved ability to respond to or tolerate field-like combinations of stresses.

Application of real-time phenomic analysis to plants grown under dynamic growth conditions, combined with QTL mapping or GWAS analyses, may provide novel insights into physiological responses to combinations of abiotic stresses. Phenomic traits that correlate with transcriptional or physiological response to or tolerance of combinations of abiotic stresses experienced under dynamic growth conditions should be selected. Such traits may include chlorophyll fluorescence measurements of redox state or NPQ (Alter et al. 2012; Külheim, Ågren, and Jansson 2002), or visual estimation of pigmentation caused by anthocyanin accumulation in response to photooxidative stress. GWAS and QTL mapping for these non-destructive measures could be conducted concomitantly with mapping of eQTLs discussed above. Such experiments would present integrative results which give holistic insight to mechanisms of tolerance of or response to combinations of abiotic stresses.

5.6 Conclusions

This thesis has developed several methods to study abiotic stresses in the framework of dynamic conditions, and presents a preliminary dataset which examines plant response to such conditions. Aiming to design and implement dynamic growth conditions that mimic regional climates, within this thesis I have presented both software and protocols to implement such conditions, and have created conditions to test the effect of light intensity in combination with abiotic stresses. Aiming to select optimal software for the high-throughput study of transcriptome dynamics using High-throughput Sequencing, and implement a framework for generation of analysis pipelines to do so, I have presented a comprehensive framework for the creation of pipelines applicable to experiments beyond those discussed in this thesis. Additionally, I have, using this framework, implemented pipelines to analyse generic RNAseq datasets. Finally, aiming to determine the transcriptional response of Arabidopsis thaliana to the combinatorial application of abiotic stresses using dynamic growth conditions, I created a preliminary RNAseq dataset, and samples for preliminary eQTL and phenomic QTL mapping datasets. This RNAseq dataset exhibited high biological noise, preventing reliable detection of differential expression on similar scales to similar published experiments, however yielded insight into patterns of differential expression that warrant further investigation. This work lays the foundation for further work elucidating the response to and tolerance of field-like combinations of abiotic stresses in a manner compatible with that suggested by recent reviews (Atkinson and Urwin 2012; Mittler 2006), and for the further discovery of the genetic architecture underlying any tolerance or response discovered.

add juice to final sentence

Bibliography

- Adamska, I (1997). ELIPs Light-induced stress proteins. *Physiologia Plantarum* 100.4, pp. 794–805. DOI: 10.1111/j.1399-3054.1997. tb00006.x (cit. on p. 14).
- Alter, P, Dreissen, A, Luo, FL, and Matsubara, S (Sept. 2012). Acclimatory responses of Arabidopsis to fluctuating light environment: comparison of different sunfleck regimes and accessions. *Photosynthesis Research* 113.1-3. PMID: 22729524 PMCID: PMC3430843, pp. 221–237. DOI: 10.1007/s11120-012-9757-2 (cit. on pp. 11, 54, 94, 95, 97).
- Andrews, S (2012). FastQC A Quality Control tool for High Throughput Sequence Data (cit. on pp. 41, 71).
- Apel, K and Hirt, H (2004). Reactive oxygen species: Metabolism, oxidative stress, and signal transduction. *Annual Review of Plant Biology* 55, pp. 373–399 (cit. on p. 11).
- Armstrong, AF, Wardlaw, KD, and Atkin, OK (2007). Assessing the relationship between respiratory acclimation to the cold and photosystem II redox poise in Arabidopsis thaliana. *Plant, Cell & Environment* 30.12, pp. 1513–1522. DOI: 10.1111/j.1365-3040.2007.01738.x (cit. on pp. 12, 16).
- Asada, K (June 2006). Production and Scavenging of Reactive Oxygen Species in Chloroplasts and Their Functions. *Plant Physiology* 141.2, pp. 391–396. DOI: 10.1104/pp.106.082040 (cit. on pp. 11, 12).
- Atkin, OK and Tjoelker, MG (July 2003). Thermal acclimation and the dynamic response of plant respiration to temperature. *Trends in Plant*

Science 8.7, pp. 343–351. DOI: 10.1016/S1360-1385(03)00136-5 (cit. on p. 11).

- Atkinson, NJ, Lilley, CJ, and Urwin, PE (Aug. 2013). Identification of Genes Involved in the Response of Arabidopsis to Simultaneous Biotic and Abiotic Stresses. *Plant Physiology* 162.4. PMID: 23800991, pp. 2028–2041. DOI: 10.1104/pp.113.222372 (cit. on pp. 12, 16, 53).
- Atkinson, NJ and Urwin, PE (June 2012). The interaction of plant biotic and abiotic stresses: from genes to the field. *Journal of Experimental Botany* 63.10. PMID: 22467407, pp. 3523–3543. DOI: 10.1093/jxb/ers100 (cit. on pp. 16, 53, 86, 95, 98).
- Avraham, S, Tung, CW, Ilic, K, Jaiswal, P, Kellogg, EA, McCouch, S, Pujar, A, Reiser, L, Rhee, SY, Sachs, MM, Schaeffer, M, Stein, L, Stevens, P, Vincent, L, Zapata, F, and Ware, D (Jan. 2008). The Plant Ontology Database: a community resource for plant structure and developmental stages controlled vocabulary and annotations. *Nucleic Acids Research* 36.suppl 1. PMID: 18194960, pp. D449–D454. DOI: 10.1093/nar/gkm908 (cit. on p. 18).
- Babu, S and Gassmann, M (2011). Assessing integrity of plant RNA with the Agilent 2100 Bioanalyzer (cit. on pp. 68, 69).
- Baker, NR (2008). Chlorophyll Fluorescence: A Probe of Photosynthesis In Vivo. *Annual Review of Plant Biology* 59.1. PMID: 18444897, pp. 89–113. DOI: 10.1146/annurev.arplant.59.032607.092759 (cit. on p. 11).
- Barua, D and Heckathorn, SA (Jan. 2006). The interactive effects of light and temperature on heat-shock protein accumulation in Solidago altissima (Asteraceae) in the field and laboratory. *American Journal of Botany* 93.1, pp. 102–109. DOI: 10.3732/ajb.93.1.102 (cit. on p. 17).

Berardini, TZ, Mundodi, S, Reiser, L, Huala, E, Garcia-Hernandez, M, Zhang, P, Mueller, LA, Yoon, J, Doyle, A, Lander, G, Moseyko, N, Yoo, D, Xu, I, Zoeckler, B, Montoya, M, Miller, N, Weems, D, and Rhee, SY (June 2004). Functional Annotation of the Arabidopsis Genome Using Controlled Vocabularies. *Plant Physiology* 135.2. PMID: 15173566, pp. 745–755. DOI: 10.1104/pp.104.040071 (cit. on p. 18).

- Boyer, JS (Oct. 1982). Plant Productivity and Environment. *Science* 218.4571. PMID: 17808529, pp. 443–448. DOI: 10.1126/science. 218.4571.443 (cit. on p. 86).
- Brachi, B, Morris, GP, and Borevitz, JO (2011). Genome-wide association studies in plants: the missing heritability is in the field. *Genome biology* 12.10. PMID: 22035733, p. 232. DOI: 10.1186/gb-2011-12-10-232 (cit. on p. 88).
- Buffalo, V (2013). Scythe A Bayesian adapter trimmer (cit. on pp. 41, 44).
- Bureau of Meteorology (Oct. 2013). Climate statistics for Australian locations (cit. on p. 87).
- Carmody, ME (May 2013). Rapid leaf-to-leaf communication of high light stress in Arabidiosis. PhD thesis (cit. on p. 91).
- Chawade, A, Bräutigam, M, Lindlöf, A, Olsson, O, and Olsson, B (Sept. 2007). Putative cold acclimation pathways in Arabidopsis thaliana identified by a combined analysis of mRNA co-expression patterns, promoter motifs and transcription factors. *BMC Genomics* 8.1. PMID: 17764576, p. 304. DOI: 10.1186/1471-2164-8-304 (cit. on p. 95).
- Covington, MF, Maloof, JN, Straume, M, Kay, SA, and Harmer, SL (Aug. 2008). Global transcriptome analysis reveals circadian regulation of key pathways in plant growth and development. *Genome Biology* 9.8. PMID: 18710561, R130. DOI: 10.1186/gb-2008-9-8-r130 (cit. on p. 92).

Demmig-Adams, B and Adams, WW (1992). Photoprotection and Other Responses of Plants to High Light Stress. *Annual Review of Plant Physiology and Plant Molecular Biology* 43.1, pp. 599–626. DOI: 10. 1146/annurev.pp.43.060192.003123 (cit. on pp. 11, 12).

- Ding, Y, Fromm, M, and Avramova, Z (Mar. 2012). Multiple exposures to drought 'train' transcriptional responses in Arabidopsis. *Nature Communications* 3, p. 740. DOI: 10.1038/ncomms1732 (cit. on p. 95).
- Fowler, S and Thomashow, MF (Aug. 2002). Arabidopsis Transcriptome Profiling Indicates That Multiple Regulatory Pathways Are Activated during Cold Acclimation in Addition to the CBF Cold Response Pathway. The Plant Cell Online 14.8. PMID: 12172015, pp. 1675–1690. DOI: 10.1105/tpc.003483 (cit. on p. 95).
- Foyer, CH and Noctor, G (Apr. 2009). Redox Regulation in Photosynthetic Organisms: Signaling, Acclimation, and Practical Implications.

 Antioxidants & Redox Signaling 11. 4, pp. 861–905 (cit. on p. 11).
- Gordon, MJ, Carmody, M, Albrecht, V, and Pogson, B (2013). Systemic and local responses to repeated HL stress-induced retrograde signaling in Arabidopsis. *Frontiers in Plant Physiology* 3, p. 303. DOI: 10.3389/fpls.2012.00303 (cit. on pp. 15, 54, 91, 92, 94, 95).
- Heinrich, S, Valentin, K, Frickenhaus, S, John, U, and Wiencke, C (Aug. 2012). Transcriptomic Analysis of Acclimation to Temperature and Light Stress in Saccharina latissima (Phaeophyceae). *PLoS ONE* 7.8, e44342. DOI: 10.1371/journal.pone.0044342 (cit. on p. 95).
- Herndon, T, Ash, M, and Pollin, R (2013). Does High Public Debt Consistently Stifle Economic Growth? A Critique of Reinhart and Rogoff (cit. on p. 89).
- Hihara, Y, Kamei, A, Kanehisa, M, Kaplan, A, and Ikeuchi, M (Apr. 2001). DNA Microarray Analysis of Cyanobacterial Gene Expression during Acclimation to High Light. *The Plant Cell Online* 13.4, pp. 793–806. DOI: 10.1105/tpc.13.4.793 (cit. on pp. 14, 95).

Jänkänpää, HJ, Mishra, Y, Schröder, WP, and Jansson, S (2012). Metabolic profiling reveals metabolic shifts in Arabidopsis plants grown under different light conditions. *Plant, Cell & Environment* 35.10, pp. 1824–1836. DOI: 10.1111/j.1365-3040.2012.02519.x (cit. on p. 16).

- Johnson, MP, Goral, TK, Duffy, CDP, Brain, APR, Mullineaux, CW, and Ruban, AV (Apr. 2011). Photoprotective Energy Dissipation Involves the Reorganization of Photosystem II Light-Harvesting Complexes in the Grana Membranes of Spinach Chloroplasts. *The Plant Cell Online* 23.4, pp. 1468–1479. DOI: 10.1105/tpc.110.081646 (cit. on p. 13).
- Johnston, DT, Wolfe-Simon, F, Pearson, A, and Knoll, AH (Oct. 2009). Anoxygenic photosynthesis modulated Proterozoic oxygen and sustained Earth's middle age. *Proceedings of the National Academy of Sciences* 106.40. PMID: 19805080, pp. 16925–16929. DOI: 10.1073/pnas.0909248106 (cit. on p. 10).
- Jung, HS, Crisp, PA, Estavillo, GM, Cole, B, Hong, F, Mockler, TC, Pogson, BJ, and Chory, J (Aug. 2013). Subset of heat-shock transcription factors required for the early response of Arabidopsis to excess light. *Proceedings of the National Academy of Sciences* 110.35. PMID: 23918368, pp. 14474–14479. DOI: 10.1073/pnas.1311632110 (cit. on p. 13).
- Karpiński, S, Reynolds, H, Karpińska, B, Wingsle, G, Creissen, G, and Mullineaux, P (Apr. 1999). Systemic Signaling and Acclimation in Response to Excess Excitation Energy in Arabidopsis. *Science* 284.5414.
 PMID: 10213690, pp. 654–657. DOI: 10.1126/science.284.5414.
 654 (cit. on p. 14).
- Kasahara, M, Kagawa, T, Oikawa, K, Suetsugu, N, Miyao, M, and Wada, M (2002). Chloroplast avoidance movement reduces photodamage in plants. *Nature* 420.6917, pp. 829–832. DOI: 10.1038/nature01213 (cit. on p. 13).

Keurentjes, JJB, Fu, J, Terpstra, IR, Garcia, JM, Ackerveken, Gvd, Snoek, LB, Peeters, AJM, Vreugdenhil, D, Koornneef, M, and Jansen, RC (Jan. 2007). Regulatory network construction in Arabidopsis by using genome-wide gene expression quantitative trait loci. *Proceedings of the National Academy of Sciences* 104.5, pp. 1708–1713. DOI: 10.1073/pnas.0610429104 (cit. on p. 96).

- Kim, D, Pertea, G, Trapnell, C, Pimentel, H, Kelley, R, and Salzberg, SL (Apr. 2013). TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions. *Genome Biology* 14.4. PMID: 23618408, R36. DOI: 10.1186/gb-2013-14-4-r36 (cit. on p. 41).
- Kim, JH (Dec. 2012). Biological Knowledge Assembly and Interpretation.
 PLoS Comput Biol 8.12, e1002858. DOI: 10.1371/journal.pcbi.
 1002858 (cit. on p. 18).
- Kimura, M, Yamamoto, YY, Seki, M, Sakurai, T, Sato, M, Abe, T, Yoshida, S, Manabe, K, Shinozaki, K, and Matsui, M (2003). Identification of Arabidopsis Genes Regulated by High Light-Stress Using cDNA Microarray. *Photochemistry and Photobiology* 77.2, pp. 226–233. DOI: 10.1562/0031-8655(2003)0770226IOAGRB2.0.CO2 (cit. on pp. 12, 92).
- Kleine, T, Kindgren, P, Benedict, C, Hendrickson, L, and Strand, Å (July 2007). Genome-Wide Gene Expression Analysis Reveals a Critical Role for CRYPTOCHROME1 in the Response of Arabidopsis to High Irradiance. *Plant Physiology* 144.3, pp. 1391–1406. DOI: 10.1104/pp. 107.098293 (cit. on p. 14).
- Kliebenstein, DJ (Sept. 2012). Exploring the Shallow End; Estimating Information Content in Transcriptomics Studies. *Frontiers in Plant Science* 3. PMID: 22973290 PMCID: PMC3437520. DOI: 10.3389/fpls.2012.00213 (cit. on p. 90).

Külheim, C, Ågren, J, and Jansson, S (July 2002). Rapid Regulation of Light Harvesting and Plant Fitness in the Field. *Science* 297.5578, pp. 91–93. DOI: 10.1126/science.1072359 (cit. on pp. 12, 13, 16, 17, 54, 87, 94, 97).

- Kumar, R, Ichihashi, Y, Kimura, S, Chitwood, DH, Headland, LR, Peng, J, Maloof, JN, and Sinha, NR (2012). A high-throughput method for Illumina RNA-Seq library preparation. Frontiers in Plant Genetics and Genomics 3, p. 202. DOI: 10.3389/fpls.2012.00202 (cit. on pp. 18, 37, 56–58, 96).
- Li, H (2013). seqtk Toolkit for processing sequences in FASTA/Q formats (cit. on p. 41).
- Li, Y, Huang, Y, Bergelson, J, Nordborg, M, and Borevitz, JO (Dec. 2010). Association mapping of local climate-sensitive quantitative trait loci in Arabidopsis thaliana. *Proceedings of the National Academy of Sciences* 107.49. PMID: 21078970, pp. 21199–21204. DOI: 10.1073/pnas.1007431107 (cit. on pp. 17, 34, 88).
- Li, Y, Roycewicz, P, Smith, E, and Borevitz, JO (Dec. 2006). Genetics of Local Adaptation in the Laboratory: Flowering Time Quantitative Trait Loci under Geographic and Seasonal Conditions in Arabidopsis. *PLoS ONE* 1.1, e105. DOI: 10.1371/journal.pone.0000105 (cit. on pp. 17, 34, 88).
- Li, Z, Wakao, S, Fischer, BB, and Niyogi, KK (2009). Sensing and Responding to Excess Light. *Annual Review of Plant Biology* 60.1. PMID: 19575582, pp. 239–260. DOI: 10.1146/annurev.arplant.58.032806. 103844 (cit. on pp. 11, 12).
- Liao, Y, Smyth, GK, and Shi, W (May 2013a). featureCounts: an efficient general-purpose read summarization program. arXiv e-print 1305.3347 (cit. on p. 41).

— (May 2013b). The Subread aligner: fast, accurate and scalable read mapping by seed-and-vote. *Nucleic Acids Research* 41.10. PMID: 23558742, e108–e108. DOI: 10.1093/nar/gkt214 (cit. on pp. 41, 45, 89).

- Lister, R, Gregory, BD, and Ecker, JR (Apr. 2009). Next is now: new technologies for sequencing of genomes, transcriptomes, and beyond. *Current Opinion in Plant Biology* 12.2, pp. 107–118. DOI: 10.1016/j.pbi.2008.11.004 (cit. on p. 18).
- Lund, SP, Dan, N, McCarthy, DJ, and Smyth, GK (2012). Detecting Differential Expression in RNA-sequence Data Using Quasi-likelihood with Shrunken Dispersion Estimates. *Statistical Applications in Genetics and Molecular Biology* 11.5, pp. 1–44 (cit. on p. 90).
- Martin, LBB, Fei, Z, Giovannoni, JJ, and Rose, JKC (2013). Catalyzing plant science research with RNA-seq. Frontiers in Plant Systems Biology 4, p. 66. DOI: 10.3389/fpls.2013.00066. (cit. on pp. 36, 89).
- McCarthy, DJ, Chen, Y, and Smyth, GK (May 2012). Differential expression analysis of multifactor RNA-Seq experiments with respect to biological variation. *Nucleic Acids Research* 40.10, pp. 4288–4297. DOI: 10.1093/nar/gks042 (cit. on p. 41).
- MILLER, G and MITTLER, R (Aug. 2006). Could Heat Shock Transcription Factors Function as Hydrogen Peroxide Sensors in Plants?

 Annals of Botany 98.2. PMID: 16740587 PMCID: PMC2803459, pp. 279–288. DOI: 10.1093/aob/mcl107 (cit. on p. 14).
- Mishra, Y, Jänkänpää, HJ, Kiss, AZ, Funk, C, Schröder, WP, and Jansson, S (Jan. 2012). Arabidopsis plants grown in the field and climate chambers significantly differ in leaf morphology and photosystem components. *BMC Plant Biology* 12.1. PMID: 22236032, p. 6. DOI: 10.1186/1471-2229-12-6 (cit. on pp. 12, 14, 16, 17, 87).
- Mittler, R (Jan. 2006). Abiotic stress, the field environment and stress combination. *Trends in Plant Science* 11.1, pp. 15–19. DOI: 10.1016/

```
j.tplants.2005.11.002 (cit. on pp. 10, 11, 16, 17, 21, 34, 53, 86, 87, 95, 98).
```

- Mittler, R and Blumwald, E (2010). Genetic Engineering for Modern Agriculture: Challenges and Perspectives. *Annual Review of Plant Biology* 61.1. PMID: 20192746, pp. 443–462. DOI: 10.1146/annurev-arplant-042809-112116 (cit. on pp. 10, 16, 21, 86, 95).
- Mubarakshina, MM, Ivanov, BN, Naydov, IA, Hillier, W, Badger, MR, and Krieger-Liszkay, A (Aug. 2010). Production and diffusion of chloroplastic H2O2 and its implication to signalling. *Journal of Experimental Botany* 61.13. PMID: 20595239, pp. 3577–3587. DOI: 10.1093/jxb/erq171 (cit. on p. 11).
- Mühlenbock, P, Szechyńska-Hebda, M, Płaszczyca, M, Baudo, M, Mateo, A, Mullineaux, PM, Parker, JE, Karpińska, B, and Karpiński, S (Sept. 2008). Chloroplast Signaling and LESION SIMULATING DISEASE1 Regulate Crosstalk between Light Acclimation and Immunity in Arabidopsis. The Plant Cell Online 20.9. PMID: 18790826, pp. 2339–2356. DOI: 10.1105/tpc.108.059618 (cit. on p. 14).
- Müller, P, Li, XP, and Niyogi, KK (Apr. 2001). Non-Photochemical Quenching. A Response to Excess Light Energy. *Plant Physiology* 125.4, pp. 1558–1566. DOI: 10.1104/pp.125.4.1558 (cit. on p. 13).
- Murchie, EH, Hubbart, S, Peng, S, and Horton, P (Jan. 2005). Acclimation of photosynthesis to high irradiance in rice: gene expression and interactions with leaf development. *Journal of Experimental Botany* 56.411, pp. 449–460. DOI: 10.1093/jxb/eri100 (cit. on p. 15).
- Niyogi, KK (1999). PHOTOPROTECTION REVISITED: Genetic and Molecular Approaches. Annual Review of Plant Physiology and Plant Molecular Biology 50.1. PMID: 15012213, pp. 333-359. DOI: 10.1146/annurev.arplant.50.1.333 (cit. on pp. 11-13).
- Nookaew, I, Papini, M, Pornputtapong, N, Scalcinati, G, Fagerberg, L, Uhlén, M, and Nielsen, J (Nov. 2012). A comprehensive comparison

of RNA-Seq-based transcriptome analysis from reads to differential gene expression and cross-comparison with microarrays: a case study in Saccharomyces cerevisiae. *Nucleic Acids Research* 40.20, pp. 10084–10097. DOI: 10.1093/nar/gks804 (cit. on p. 18).

- Page, M, Sultana, N, Paszkiewicz, K, Florance, H, and Smirnoff, N (2012). The influence of ascorbate on anthocyanin accumulation during high light acclimation in Arabidopsis thaliana: further evidence for redox control of anthocyanin synthesis. *Plant, Cell & Environment* 35.2, pp. 388–404. DOI: 10.1111/j.1365-3040.2011.02369.x (cit. on p. 95).
- Peng, RD (July 2009). Reproducible research and Biostatistics. *Biostatistics* 10.3. PMID: 19535325, pp. 405–408. DOI: 10.1093/biostatistics/kxp014 (cit. on p. 89).
- Ptitsyn, A (Aug. 2008). Comprehensive analysis of circadian periodic pattern in plant transcriptome. *BMC Bioinformatics* 9.Suppl 9. PMID: 18793463, S18. DOI: 10.1186/1471-2105-9-S9-S18 (cit. on p. 92).
- Rapaport, F, Khanin, R, Liang, Y, Pirun, M, Krek, A, Zumbo, P, Mason, CE, Socci, ND, and Betel, D (Sept. 2013). Comprehensive evaluation of differential gene expression analysis methods for RNA-seq data.

 *Genome Biology 14.9. PMID: 24020486, R95. DOI: 10.1186/gb-2013-14-9-r95 (cit. on pp. 90, 96).
- Rizhsky, L, Liang, H, and Mittler, R (Nov. 2002). The Combined Effect of Drought Stress and Heat Shock on Gene Expression in Tobacco. *Plant Physiology* 130.3. PMID: 12427981, pp. 1143–1151. DOI: 10. 1104/pp.006858 (cit. on p. 16).
- Rizhsky, L, Liang, H, Shuman, J, Shulaev, V, Davletova, S, and Mittler, R (Apr. 2004). When Defense Pathways Collide. The Response of Arabidopsis to a Combination of Drought and Heat Stress. *Plant Physiology* 134.4. PMID: 15047901, pp. 1683–1696. DOI: 10.1104/pp.103.033431 (cit. on p. 16).

Robinson, MD, McCarthy, DJ, and Smyth, GK (Jan. 2010). edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. *Bioinformatics* 26.1. PMID: 19910308 PMCID: PMC2796818, pp. 139–140. DOI: 10.1093/bioinformatics/btp616 (cit. on pp. 18, 41, 89).

- Robinson, MD and Oshlack, A (Mar. 2010). A scaling normalization method for differential expression analysis of RNA-seq data. *Genome Biology* 11.3. PMID: 20196867, R25. DOI: 10.1186/gb-2010-11-3-r25 (cit. on pp. 18, 45, 89).
- Robinson, M, McCarthy, D, Chen, Y, and Smyth, GK (2013). edgeR: differential expression analysis of digital gene expression data User's Guide (cit. on pp. 75, 93).
- Rossel, JB, Wilson, IW, and Pogson, BJ (Nov. 2002). Global Changes in Gene Expression in Response to High Light in Arabidopsis. *Plant Physiology* 130.3, pp. 1109–1120. DOI: 10.1104/pp.005595 (cit. on pp. 12–14, 16, 92).
- Rossel, JB, Wilson, PB, Hussain, D, Woo, NS, Gordon, MJ, Mewett, OP, Howell, KA, Whelan, J, Kazan, K, and Pogson, BJ (Dec. 2007). Systemic and Intracellular Responses to Photooxidative Stress in Arabidopsis. *The Plant Cell Online* 19.12, pp. 4091–4110. DOI: 10.1105/tpc.106.045898 (cit. on p. 92).
- Ruijter, JM, Ramakers, C, Hoogaars, WMH, Karlen, Y, Bakker, O, Hoff, MJB van den, and Moorman, AFM (Apr. 2009). Amplification efficiency: linking baseline and bias in the analysis of quantitative PCR data. Nucleic acids research 37.6. PMID: 19237396, e45. DOI: 10.1093/nar/gkp045 (cit. on p. 65).
- Seki, M, Kamei, A, Yamaguchi-Shinozaki, K, and Shinozaki, K (Apr. 2003). Molecular responses to drought, salinity and frost: common and different paths for plant protection. *Current Opinion in Biotechnology*

14.2, pp. 194–199. DOI: 10.1016/S0958-1669(03)00030-2 (cit. on pp. 11, 16).

- Seki, M, Narusaka, M, Abe, H, Kasuga, M, Yamaguchi-Shinozaki, K, Carninci, P, Hayashizaki, Y, and Shinozaki, K (Jan. 2001). Monitoring the Expression Pattern of 1300 Arabidopsis Genes under Drought and Cold Stresses by Using a Full-Length cDNA Microarray. *The Plant Cell Online* 13.1. PMID: 11158529, pp. 61–72. DOI: 10.1105/tpc.13.1.61 (cit. on pp. 12, 16, 53).
- Spokas, K and Forcella, F (Jan. 2006). Estimating hourly incoming solar radiation from limited meteorological data. Weed Science 54.1, pp. 182–189. DOI: 10.1614/WS-05-098R.1 (cit. on pp. 22, 23, 26).
- Subramanian, A, Tamayo, P, Mootha, VK, Mukherjee, S, Ebert, BL, Gillette, MA, Paulovich, A, Pomeroy, SL, Golub, TR, Lander, ES, and Mesirov, JP (Oct. 2005). Gene set enrichment analysis: A knowledge-based approach for interpreting genome-wide expression profiles. *Proceedings of the National Academy of Sciences of the United States of America* 102.43. PMID: 16199517, pp. 15545–15550. DOI: 10.1073/pnas.0506580102 (cit. on p. 18).
- Sun, W and Hu, Y (May 2013). eQTL Mapping Using RNA-seq Data.
 Statistics in Biosciences 5.1, pp. 198–219. DOI: 10.1007/s12561-012-9068-3 (cit. on p. 89).
- Suzuki, N, Koussevitzky, S, Mittler, R, and Miller, G (2012). ROS and redox signalling in the response of plants to abiotic stress. *Plant, Cell & Environment* 35.2, pp. 259–270. DOI: 10.1111/j.1365-3040.2011.02336.x (cit. on p. 12).
- Takahashi, S and Badger, MR (Jan. 2011). Photoprotection in plants: a new light on photosystem II damage. *Trends in Plant Science* 16.1, pp. 53–60. DOI: 10.1016/j.tplants.2010.10.001 (cit. on p. 12).
- Takahashi, S, Milward, SE, Fan, DY, Chow, WS, and Badger, MR (Mar. 2009). How Does Cyclic Electron Flow Alleviate Photoinhibition in

Arabidopsis? *Plant Physiology* 149.3, pp. 1560–1567. DOI: 10.1104/pp.108.134122 (cit. on p. 13).

- Takahashi, S, Milward, SE, Yamori, W, Evans, JR, Hillier, W, and Badger, MR (July 2010). The Solar Action Spectrum of Photosystem II Damage. *Plant Physiology* 153.3, pp. 988–993. DOI: 10.1104/pp. 110.155747 (cit. on p. 15).
- Tikkanen, M, Grieco, M, Nurmi, M, Rantala, M, Suorsa, M, and Aro, EM (Dec. 2012). Regulation of the photosynthetic apparatus under fluctuating growth light. *Philosophical Transactions of the Royal Society B: Biological Sciences* 367.1608, pp. 3486–3493. DOI: 10.1098/rstb.2012.0067 (cit. on p. 12).
- Tikkanen, M, Piippo, M, Suorsa, M, Sirpiö, S, Mulo, P, Vainonen, J, Vener, AV, Allahverdiyeva, Y, and Aro, EM (Nov. 2006). State transitions revisited—a buffering system for dynamic low light acclimation of Arabidopsis. *Plant Molecular Biology* 62.4-5, pp. 779–793. DOI: 10.1007/s11103-006-9044-8 (cit. on p. 13).
- Van Verk, MC, Hickman, R, Pieterse, CM, and Van Wees, SC (Apr. 2013). RNA-Seq: revelation of the messengers. Trends in Plant Science 18.4, pp. 175–179. DOI: 10.1016/j.tplants.2013.02.001 (cit. on pp. 18, 37, 39, 42).
- Vanderauwera, S, Zimmermann, P, Rombauts, S, Vandenabeele, S, Langebartels, C, Gruissem, W, Inzé, D, and Van Breusegem, F (Oct. 2005). Genome-Wide Analysis of Hydrogen Peroxide-Regulated Gene Expression in Arabidopsis Reveals a High Light-Induced Transcriptional Cluster Involved in Anthocyanin Biosynthesis. *Plant Physiology* 139.2, pp. 806–821. DOI: 10.1104/pp.105.065896 (cit. on p. 13).
- Väremo, L, Nielsen, J, and Nookaew, I (Feb. 2013). Enriching the gene set analysis of genome-wide data by incorporating directionality of gene expression and combining statistical hypotheses and methods.

 Nucleic Acids Research. DOI: 10.1093/nar/gkt111 (cit. on p. 18).

Wang, Y, Ghaffari, N, Johnson, CD, Braga-Neto, UM, Wang, H, Chen, R, and Zhou, H (Oct. 2011). Evaluation of the coverage and depth of transcriptome by RNA-Seq in chickens. *BMC Bioinformatics* 12.Suppl 10. PMID: 22165852, S5. DOI: 10.1186/1471-2105-12-S10-S5 (cit. on p. 37).

- Wang, Z, Gerstein, M, and Snyder, M (Jan. 2009). RNA-Seq: a revolutionary tool for transcriptomics. *Nature Reviews Genetics* 10.1, pp. 57–63. DOI: 10.1038/nrg2484 (cit. on pp. 18, 36, 37, 89).
- Wilkins, O, Bräutigam, K, and Campbell, MM (2010). Time of day shapes Arabidopsis drought transcriptomes. *The Plant Journal* 63.5, pp. 715–727. DOI: 10.1111/j.1365-313X.2010.04274.x (cit. on p. 92).
- Wituszyńska, W, Gałązka, K, Rusaczonek, A, Vanderauwera, S, Van Breusegem, F, and Karpiński, S (Apr. 2013). Multivariable environmental conditions promote photosynthetic adaptation potential in Arabidopsis thaliana. *Journal of Plant Physiology* 170.6, pp. 548–559. DOI: 10.1016/j.jplph.2012.11.016 (cit. on pp. 12, 16, 17, 34, 53, 54, 87, 95).
- Yi, X, Du, Z, and Su, Z (Apr. 2013). PlantGSEA: a gene set enrichment analysis toolkit for plant community. Nucleic Acids Research 41.W1, W98–W103. DOI: 10.1093/nar/gkt281 (cit. on p. 18).
- Young, MD, Wakefield, MJ, Smyth, GK, and Oshlack, A (Feb. 2010). Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biology 11.2. PMID: 20132535, R14. DOI: 10.1186/gb-2010-11-2-r14 (cit. on p. 41).

Chapter 6

Appendix

Notes:

Code listings, where included, are illustrative. Full source code of all software developed is large (over 5000 lines of code), and will be distrubuted as a gzipped tar archive. The latest code for all pipleines, scripts, is available online. See Appendix section 6.1 and Table 6.1

6.1 Source Code Repositories

The following source code repositiories have been created as part of this thesis.

Repository	URL
This thesis	http://github.com/kdmurray91/hons-thesis
spcControl module	http://github.com/borevitzlab/spcControl

Table 6.1: Source code repositories

6.2 Miscelanelous Software

The following pieces of software are not part of any software package, however are used within this thesis.

Most scritps are available within the bioscripts reposistory on my github site, available at https://github.com/kdmurray91/bioscripts.

6.2.1 spliceSolarCalc.py

Create solarcalc model files which fluctuate between two model files on a regular cycle. Available at https://github.com/kdmurray91/bioscripts/blob/master/solarcalc/spliceSolarCalc.py.

6.2.2 Analysis of RNAseq analysis pipeline computation cost

Statistical analysis of RNAseq pipeline computation cost.

https://github.com/kdmurray91/hons-thesis-stats/blob/master/pltimes/pltimes.Rmd

6.2.3 qPCR analysis Code

Statistical calculation of relative quantification and differential exression in qPCR datasets.

 $A vailable\ at\ \texttt{https://github.com/kdmurray91/hons-thesis-stats/}$

blob/master/qpcr/qpcr.Rnw

6.3 spcControl Module Implementation Details

6.3.1 Evolution of the spcControl codebase

Initially	SPC evolution
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6.3.2 Invocation of spcControl

Some important notes about the invocation of spcControl:

• SolarCalc is a java program. The implementation of strftime function, which formats date objects into character strings (text) in either SolarCalc or java does not comply with the standards which the datetime module of python does. Thus, each time in the hour after midnight is recorded as, for example, 07/01/12 00:15:00AM, for 15 minutes after midnight, July 1, 2012. As a result, we cannot directly use solarcalc output. There is a very simple fix however, finding all occurances of 00: in the solarcalc output file and replacing them with 12: solves this issue. As these files are very large, this is best accomplished with sed, the Stream EDitor, as show below in listing 3.

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sed -ibak -e 's/00:/12:/g' <file>

 $\textbf{Listing 3:} \ \ \textbf{Code to ensure SolarCalc makes dates which are compatible with python}$