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Gene expression analysis of malting barley

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Gene expression analysis of malting barley

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B.App.Sc (Biotechnology)

A thesis submitted in fulfilment of the degree of Doctor of Philosophy

Centre for Plant Conservation Genetics

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September 2008
Thesis declaration

I certify that the work presented in this thesis is, to the best of my knowledge and belief, original, except as acknowledged in the text, and that the material has not been submitted, either in whole or in part, for a degree at this or any other university. I acknowledge that I have read and understood the University's rules, requirements, procedures and policy relating to my higher degree research award and to my thesis. I certify that I have complied with the rules, requirements, procedures and policy of the University (as they may be from time to time).

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Date
Acknowledgements

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Abstract

Malting barley (*Hordeum vulgare*) is of significant importance and economic value to the malting and brewing industries. The biochemical and physiological processes of barley seed germination (malting) have subsequently been the focus of substantial research; however the underlying molecular processes require further investigation. The introduction of large-scale gene expression technologies has provided an opportunity to contribute significant genetic data to the understanding of the malting process. Long Serial Analysis of Gene Expression (LongSAGE) was applied to malting barley to characterise the transcriptional profile exhibited during grain germination. Eight LongSAGE tag libraries were constructed, one from whole dry seed and one from each of the time points 0, 12, 24, 48, 72, 96 and 120 hours post-steeping. Approximately 20 000 tags were sequenced from each LongSAGE library, generating 41 909 unique tag sequences. Annotations of the 100 most abundant LongSAGE tags from each library were analysed to reveal the major functional groups expressed in the malting transcriptome. Statistical analysis confidently identified 57 LongSAGE tags with significant changes in tag abundance across time. The genes that correspond to eight of these tags were targeted for validation by real-time RT-PCR. Real-time RT-PCR experiments were conducted using RNA extracted from the same grain sample from four of the time points examined by LongSAGE; whole dry seed, 12 hours post-steeping, 48 hours post-steeping and 96 hours post-steeping. Three of the transcripts show co-ordinated expression demonstrating significant up-regulation at 48 hours post-steeping and remain significantly elevated during malting. These transcripts α-amylase type B, (1-3,1-4)-β-D-glucanase and cysteine proteinase EP-B represent key starch, cell wall and protein modifying
enzymes known to play significant roles in the malting process. mRNA abundance levels observed by real-time RT-PCR correlate well with the data obtained from LongSAGE.

Microarray is another large scale gene expression technology with the potential to generate large amounts of genetic data quickly and efficiently. A CombiMatrix 12K ElectraSense™ Custom Array chip was designed from the unique LongSAGE tags from 24 hours post-steeping. RNA from the malting barley 12 hours post-steeping and 48 hours post-steeping samples was hybridised to the array chips. The CombiMatrix ElectraSense™ system incorporates electrochemical detection of a redox reaction proximal to probe specific electrodes. 113 probes show a greater than 10 fold difference between 12 and 48 hours post-steeping with 64 probes preferentially expressed at 12 hours post-steeping and 49 probes preferentially expressed at 48 hours post-steeping. The eight genes of interest analysed by real-time RT-PCR were represented by one or more probes on the malting barley CombiMatrix ElectraSense™ array. Fold changes observed between 12 and 48 hours post-steeping by the CombiMatrix ElectraSense™ microarray support the fold changes observed by LongSAGE and real-time RT-PCR.
List of publications included as part of the thesis

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Statement of contribution of others

The input of co-authors to the published chapters of this thesis is acknowledged by the author. The author would like to recognize the contribution of co-authors to Chapter Two “Abundant transcripts of malting barley identified by serial analysis of gene expression (SAGE)”, published in Plant Biotechnology Journal. T Pacey-Miller assisted with extraction of mRNA, A Crawford prepared the SAGE libraries, excluding HvTaM4 which was prepared by J White, G Cordeiro and D Barbary assisted with databases and annotations, and P Bundock prepared Table 2.5 and comments about antisense transcripts. Comments and suggestions for improving the manuscript were provided by T Pacey-Miller, P Bundock and R Henry, all other work was completed by J White.

The author would like to acknowledge the contribution of co-authors to Chapter Three “Differential LongSAGE tag abundance analysis in a barley seed germination time course and validation with relative real-time RT-PCR” accepted for publication in Plant Science. All laboratory work and analysis was carried out by J White. Comments and suggestions for improving the manuscript were provided by T Pacey-Miller, P Bundock and R Henry.

The author would like to recognize the contribution of co-authors to Chapter Four “Transcriptome analysis of malting barley; comparison of Microarray, LongSAGE and real-time RT-PCR” submitted for publication in Functional and Integrative Genomics. Slide preparation and chip hybridisation was carried out by G Good and P Bundock. All other work was carried out by J White and comments and suggestions
for improving the manuscript were provided by T Pacey-Miller, P Bundock and R Henry.

Signature of the candidate

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Signature of the supervisor

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List of additional publications by the candidate

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Chapter One

Literature Review and General Introduction – Analysis of Gene Expression in Germinating Barley

1.1 Barley

1.1.1 The Importance of Barley

Cultivated barley (*Hordeum vulgare* L.) is an important grain crop, the world’s fourth major cereal crop after maize, wheat and rice (Langridge and Barr, 2003). Barley provides a major contribution to the world food supply in the areas of human food, stock feed and malt production, while also playing an important role as an experimental and model plant species for study (Nilan and Ullrich, 1993). Barley malt is used to add flavour to a variety of different foods; however the most extensive use of barley malt is for brewing (Bamforth and Barclay, 1993).

Breeding strategies and farming practices in the last few decades have resulted in profitable yield increases in barley production (Langridge and Barr, 2003). In Australia, barley production and area have both increased over time. Between the 1996/97 and 2003/04 winter crop seasons, Australian barley production increased 1.3 fold although the sown area was only 1.1 times greater (Figure 1.1) (ABARE, 2003). For Australia the growing seasons for 2002/03, 2004/05 and 2006/07 resulted in poor production due to extremely dry conditions in the main growing states. The forecast for 2007/08, while higher than 2006/07, is still relatively low as most cropping regions of Australia have recorded below average winter rainfall. These extremely dry
conditions have been confounded by above average daytime temperatures which have placed additional pressure on the moisture stressed crops (Brown et al., 2007). It is becoming increasingly obvious that research in breeding and farming practices needs to focus on barley production in dryer, hotter conditions if improvements in yield over area are to continue.

Figure 1.1 Barley production in Australia. Based on data from ABARE Australian Crop Reports (Brown et al., 2007; Hogan et al., 2004) and Australian Commodity Statistics (ABARE, 2003; ABARE, 2007). f  ABARE forecast.

The amount of barley used for domestic malting and malting barley exports have also increased over time. Australia is a large exporter of barley malt, according to the United Nations Statistics Division, Australia was the 4th largest exporter of barley in 2008 (UNdata, 2009). The five year average to 2006/07 for total Australian barley production used for export was approximately 66% (Brown et al., 2007). The average price per tonne of Australian barley is consistently increasing for each of domestic
barley, export feed barley and export malt barley. The increases in the value of
Australian barley and barley malt are likely to be a result of increased crop quality,
increased likelihood of the harvested grain achieving malt quality, and improved
barley varieties and farming practices in the last few decades (Langridge and Barr,
2003).

1.1.2 Barley Taxonomy and Origin

Barley is a member of the grass family, the Poaceae, the barley genus Hordeum
belongs to the Triticeae tribe (Nilan and Ullrich, 1993). Archaeological remains of
barley grains found in the Israel-Jordan region of the Fertile Crescent indicate that
barley was domesticated there about 10 000 years ago when the cultivated form,
*Hordeum vulgare*, is believed to have split from its wild relative *Hordeum
spontaneum* (Badr et al., 2000). Modern nomenclature recognises the two forms as
belonging to different subspecies of *Hordeum vulgare* (H.v.) (Nilan and Ullrich,
1993). They are closely related and are inter-fertile, though *H.v.vulgare* can be both
six- and two-rowed with non-brittle rachis, whereas the wild *H.v.spontaneum* is two-
rowed with a brittle rachis (Nilan and Ullrich, 1993). Morphological seed differences
such as groove depth and endosperm width are some of the characters used to
discriminate wild from cultivated barley lines (Figure 1.2) (Badr et al., 2000). One of
31 Hordeum species, cultivated barley is a diploid with 2n=14 chromosomes. Diploid,
tetraploid and hexaploid varieties exist amongst the wild Hordeum species (Nilan and
Ullrich, 1993).
1.1.3 Morphology of barley inflorescence

The inflorescence of *H. v. vulgare* (Figure 1.3), which as the focus of this review will herein be referred to as barley, is a spike with an axis known as a rachis (Duffus and Cochrane, 1993). The spike has three spikelets alternating on opposite sides of each node of the flat rachis of the head. There is one central and two lateral spikelets (Nilan and Ullrich, 1993) each with a ventral glume or pala, and a dorsal glume, or lemma, which make up the husk, or hull (Duffus and Cochrane, 1993). When all three spikelets are fertile, it is referred to as a six-rowed spike. If only the central spikelet is fertile, the spike is known as two-rowed (Nilan and Ullrich, 1993). Each spikelet is a single floret composed of a lemma, pala, a single ovary and style, and three stamens and anthers. Cultivated barley is effectively a self-pollinator with an extremely low occurrence of cross pollination (Nilan and Ullrich, 1993).

Figure 1.2 Morphological seed differences of wild and cultivated barley

Reproduced from (Badr *et al.*, 2000)
1.2 Seed Germination

Germination begins with uptake of water by the dry seed, and ends with the elongation of the radicle, or embryonic axis (Bewley, 1997). Most of the water
penetrates the seed near the embryo and spreads at different rates through various regions of the seed (Fincher and Stone, 1993). Initially imbibition is the dominant process of water uptake, as it is not dependent on metabolic activity, however once the protoplasm is hydrated and metabolic activity resumes, osmotic forces become more important in controlling the water content (Street and Opik, 1970). Substrates and enzymes for some metabolic processes appear to be present within the dry seed (Bewley, 1997; Street and Opik, 1970), and respiratory activity can be detected within minutes of imbibition (Bewley, 1997). The synthesis of ribosomal proteins under conditions of transcription inhibition suggests that pre-formed transcripts are stored in the embryonic axis of the dry seed (Beltran-Pena et al., 1995). These ready to use mRNA enable the synthesis of proteins and assist in the production of new mRNAs during germination (Bewley, 1997).

1.3 Barley Malting

1.3.1 Malting

Malting is the industrial process applied to cereal grains to generate malt. Malt is used to add flavour, colour, sweetness and even nutrition to many foods. The most common use for malt is to provide a source of fermentable sugars to produce beer and whiskey through alcoholic fermentation (Bamforth and Barclay, 1993). Barley varieties differ widely in their potential as malting barleys. The production of malt for whichever purpose is dependent on the viability of the barley grain. Suitable malt barley will have low dormancy, low water sensitivity and high vigour (Bamforth and Barclay, 1993). Husked barleys only are used for malting (Briggs, 1998) with the
husks providing an efficient filter bed in the brewing process, through which wort is separated from residual solid particles after mashing (Bamforth and Barclay, 1993; Kasha et al., 1993).

Irrespective of the intended use of malt, malting processes are similar. The malting process begins with steeping the barley in water to achieve a sufficient moisture level to encourage germination (Bamforth and Barclay, 1993). Steeping, or imbibition, activates metabolism in the aleurone and embryonic tissues which synthesise hydrolytic enzymes that catalyse the degradation of storage proteins in the starchy endosperm (Fincher and Stone, 1993). Once the starch granules are exposed and the grain is evenly modified or germinated, the grain is ready for kilning. Kilning arrests germination and stabilizes the malt by lowering the moisture levels ready for milling (Bamforth and Barclay, 1993).

1.3.2 Steeping

Steeping, the soaking of barley grains in water, is performed to increase the moisture content of the grain, to ensure rapid and uniform germination while minimising embryo growth and respiration (Bamforth and Barclay, 1993). The moisture content must be taken from 11-12% up to 42-46% in 48-52 hours (Bamforth and Barclay, 1993). To induce germination requires the thorough wetting of the grain, interrupted by air rests that provide the grain access to oxygen for respiration and helps to overcome the water sensitivity of some grain (Bamforth and Barclay, 1993). Grain temperatures are controlled during air rests, as respiration produces a considerable output of heat energy (Briggs, 1998).
During imbibition, the grain swells, increasing its volume by approximately one quarter, the width and depth of the grain increase but the length does not (Briggs, 1998). The acrospire, or sprout, and rootlet begin to grow, and these are traditionally used as a guide to the malting progress. The growing acrospire presses a visible groove into the outer tissues and its length is noted as a fraction of the grain’s (Briggs, 1998). When the average length of the acrospire is approximately $\frac{3}{4}$ the length of the grain, and the endosperm is easily rubbed out between the finger and thumb, the malt is ready to be kilned (Bamforth and Barclay, 1993).

When imbibition begins, the embryo and husk absorb water more rapidly than the endosperm, and it has been thought that the embryo regulates water uptake (Bamforth and Barclay, 1993). More recently it has been suggested that structural differences in the starchy endosperm also affect water uptake during germination (Chandra et al., 1999). Structural differences in barley grains have been described as mealy or steely. Mealy endosperms contain air spaces between their loosely packed cells and starch granules, and appear to display faster distribution of water during steeping, whereas steely endosperms have a more densely packed protein matrix and smaller starch granules (Chandra et al., 1999).

1.3.3 Germination

During the malting process germination is targeted to maximise the production of extractable material for malt, through the process of endosperm modification. Modification describes the physical and chemical changes that occur during malting,
including the degradation of the starchy endosperm and the associated biochemical changes (Briggs, 1998). The rate of modification depends on several factors; the rate of moisture distribution through the endosperm, the rate of enzyme synthesis, the extent of enzyme release and the structure of the endosperm which will determine its digestibility (Bamforth and Barclay, 1993). Hydrolytic enzymes, initially released from the scutellum and later the aleurone layer, degrade cell walls, allowing the matrix proteins to be broken down, freeing the starch granules and exposing them to degradation (Briggs, 1998).

The hormone activity of gibberellic acid (GA) and abscisic acid (ABA) plays a large part in endogenous regulation of germination by inducing and repressing the synthesis and secretion of hydrolytic enzymes respectively (Fincher and Stone, 1993). GA is released from the scutellum and embryo after imbibition; it diffuses through the endosperm to the aleurone layer where it stimulates the production of hydrolytic enzymes, including α-amylase and β-glucanase (Briggs, 1998; Fincher and Stone, 1993; Russell, 1998). α-Amylase acts to break down starch to generate soluble sugars and is the most extensively studied aleurone enzyme (Gibson, 2004; Jones and Jacobsen, 1991). Several α-amylase isoenzymes have been identified and divided into two groups based on their isoelectric points (Fincher, 1989; MacGregor et al., 1988). The group of enzymes with low-pI values degrade starch more rapidly than the high-pI group (Fincher, 1989).

The cell walls within the barley endosperm contain a great quantity of the polysaccharide β-glucan (Nuutila et al., 2002). Cell wall (1-3,1-4)-β-glucan is broken down by (1-3,1-4)-β-glucanases during germination (Fincher, 1989). There are two
closely related isoenzymes of (1-3,1-4)-β-glucanase in barley, EI and EII (Slakeski and Fincher, 1992; Woodward and Fincher, 1982). Beta-glucan concentration is relatively low compared to starch, however it can cause difficulties with malting and brewing, and affect malt quality (Nuutila et al., 2002; Wang et al., 2004). The rate of modification is controlled by water uptake, the production of hydrolytic enzymes and by the content of cell wall β-glucan and more significantly, the ability to synthesise β-glucanase (Bamforth and Barclay, 1993). The native barley β-glucanase is susceptible to inactivation at temperatures above 55°C (Nuutila et al., 2002). The development of a genetically modified, thermotolerant β-glucanase expressed during germination, has increased degradation activity during germination and mashing, which in turn decreases β-glucan concentration in wort and improves the filtration of beer (Nuutila et al., 2002; Wang et al., 1998).

During malting germination is controlled by maintaining moisture, oxygen supply, carbon dioxide removal and elimination of excess heat. Germination normally occurs between 16 and 20°C (Bamforth and Barclay, 1993). Higher temperatures can increase growth rate, however this cannot be sustained over time as the rate of enzyme formation decreases. Malting time is limited; therefore the grain must germinate quickly and evenly, in the dark, and in modern malteries is expected to reach its maximum germination within three days (Briggs, 1998).

In higher plants, germination is usually preceded by a period of dormancy (Kuhlemeier et al., 1987). Seed dormancy in grain crops is not usually considered an advantageous characteristic. Bewley (1997) and Bove et al. (2001) review several studies that identify proteins, mRNAs and hormones involved with dormancy and
seed germination. The mechanisms by which the embryo emerges from the seed after
germination and how this process is blocked in a dormant seed remain to be fully
understood (Bewley, 1997; Bove et al., 2001).

1.3.4 Kilning

When the grain is completely and evenly modified it should be friable and thus ready
for kilning (Bamforth and Barclay, 1993). Kilning gradually dries the grain, reducing
the high moisture levels required for germination down to less than 5%. This must be
regulated carefully to ensure the survival of enzymes that are crucial to brewing and
distilling processes (Bamforth and Barclay, 1993). Some enzymes are more heat
tolerant at lower moisture contents so drying begins at a relatively low temperature
and increases progressively affecting flavour and colour changes over an approximate
24 hour kilning period (Bamforth and Barclay, 1993). On completion of kilning the
malt grain is milled, ready to begin the brewing process.

1.4 Gene Expression

The germination rate of barley is a major contributor to malt quality. The mechanisms
that drive biological events and developmental processes, such as germination, can be
revealed by monitoring the pattern of gene expression under various physiological and
pathological conditions (Schnable et al., 2004; Snustad and Simmons, 2000). Gene
expression can be controlled at several levels in eukaryotes including transcription,
RNA processing, RNA transport, translation, RNA degradation and protein
degradation (Russell, 1998). The ability of a cell to control these processes allows the
production of a variety of diverse proteins in different cells at separate times (Kuhlemeier et al., 1987; Russell, 1998). Molecular biology techniques including RNA extraction, reverse transcription and northern blot analysis, have allowed us to examine the expression of single genes in different tissues or at different developmental time points. These types of analyses are beneficial; however they can be difficult and allow characterisation of only one gene at a time. Modern molecular biology techniques allow researchers to analyse the entire transcriptome of a cell or tissue by characterising differential expression of mRNAs (Donson et al., 2002; Dunwell et al., 2001).

1.5 Molecular Biology Techniques

1.5.1 Genomics

The genomes of eukaryotes are generally much larger than prokaryotes, however in most eukaryote genomes only a small proportion of the DNA encodes amino acid sequences. To focus on the protein-coding content of a genome, it is common to only analyse expressed genes or DNA sequences that are transcribed into messenger RNA (mRNA) copies (Snustad and Simmons, 2000). A specialized enzyme, reverse transcriptase, is used to catalyse the synthesis of DNA molecules complementary to mRNA templates called cDNAs (complementary DNAs) (Snustad and Simmons, 2000). Molecular biologists create cDNA libraries which contain the coding regions of the expressed genes. To construct a library, cDNA sequences are inserted into cloning vectors to form recombinant DNA molecules (Snustad and Simmons, 2000). These are introduced into host bacterial cells which are grown into colonies that will
contain just one cDNA insert that can be isolated and sequenced. Expressed genes can be identified from their DNA sequence. Single pass sequencing of cDNA clones produces Expressed Sequence Tags (ESTs) that are used to characterise the genes expressed in a tissue (Snustad and Simmons, 2000).

The EST approach to functional genomics can be expensive and time consuming and may miss genes expressed at low levels or in only a few specialized cells unless a normalization step is carried out and the library is extensively sequenced. A method to identify patterns of gene expression across different tissues, throughout development, or in response to external stimuli, is northern blot hybridization (Snustad and Simmons, 2000). In this method, RNAs from specific starting material are separated by gel electrophoresis and transferred to membranes for hybridization with a labelled probe. The specific sequence probe will bind with complementary transcripts on the membrane and display a level of expression for that transcript. The presence of certain gene transcripts, in different cells at various or specific times, can indicate functions and interactions of genes during development (Schnable et al., 2004).

1.5.2 Microarray

Northern blot hybridisation is designed to analyse the expression of one transcript at a time. To analyse the expression of large sets of genes, dot blot hybridisation or microarray technology was developed (Schena et al., 1995; Snustad and Simmons, 2000). A microarray is a reverse northern, in which gene specific nucleotide sequences or probes are bound in a specific array pattern to a solid surface, and these are hybridised with fluorescent cDNA or cRNA preparations (Snustad and Simmons,
The amount of cDNA that hybridises to each probe/dot can be measured by scanning the array with an imaging system and analysing the results with computer programs (Snustad and Simmons, 2000). The microarray can monitor many thousands of genes by using high speed robotic printing onto microscope slides (Schena et al., 1995). Microarrays contain replicated features to test the reproducibility of the process (Schena et al., 1995) and allow differential gene expression to be measured by simultaneous two-colour hybridisation (Aharoni and Vorst, 2002; Schena et al., 1995). Two mRNA sources are prepared by reverse transcription to create two cDNA probes with different fluorophores (Schena et al., 1995). Samples of known concentration are used to calibrate the fluorescence of the probes so that expression can be quantified and compared (Schena et al., 1995). Different types of microarray platforms are available for different applications. The major systems include two-colour or one-colour platforms designed to detect the hybridisation of probes to oligonucleotide or cDNA features.

A recent improvement to the development of microarrays is the ElectraSense™ electrochemical detection system developed by CombiMatrix (Ghindilis et al., 2007; Roth et al., 2006). Microarrays have previously been manufactured by spotting pre-synthesized molecules, using inkjet or light-directed synthesis processes and the signal intensities of subsequently hybridised molecules measured with fluorescent detection. CombiMatrix uses computer software to guide the simultaneous synthesis of different oligonucleotides at thousands of platinum microelectrodes using a modified semiconductor (Ghindilis et al., 2007). This electronic system can subsequently be used for electrochemical detection (ElectraSense™) of target molecules. ElectraSense™ technology is based on detection of a redox reaction
proximal to the specific electrode. Target molecules are labelled with biotin, hybridised to the array, the biotin is bound to a horseradish peroxidase (HRP) streptavidin marker and electrodes are exposed to tetramethylbenzidine (TMB) and hydrogen peroxide (Ghindilis et al., 2007; Roth et al., 2006). The oxidized TMB is reduced at the electrode surface which generates an electrochemical signal (Figure 1.4) (Ghindilis et al., 2007; Roth et al., 2006).

**Figure 1.4 Electrochemical detection scheme of CombiMatrix ElectraSense™ microarray.** Figure taken from (Ghindilis et al., 2007).

Microarray analysis is a valuable tool for plant functional genomics and has been thoroughly reviewed as an effective technique for identification and characterisation of differentially expressed genes (Aharoni and Vorst, 2002; Desprez et al., 1998; Donson et al., 2002; Dunwell et al., 2001; Schnable et al., 2004; Wullschleger, 2003; Zhu, 2003). Differentially expressed genes are identified by comparing two or more biological samples. A typical experiment will compare samples of different varieties, different times in development or treated versus untreated samples. It is common to keep almost all attributes of the experiment the same except for one feature of interest. Two samples can then be compared to determine the level of gene expression and
how it is affected in each sample. A differentially expressed gene would present high levels of expression in one sample and low levels of expression in the other. Microarray technology is an important tool for differential expression analysis however it is limited to the analysis of genes with known sequence. Serial Analysis of Gene Expression (SAGE) allows all transcripts in a sample to be profiled without prior knowledge of what the transcripts encode (Dunwell et al., 2001; Velculescu et al., 1995).

1.6 SAGE

1.6.1 SAGE

The SAGE method developed by Velculescu et al. (1995) enables analysis of gene expression of a large number of transcripts in a quantitative and unbiased manner. SAGE is based on the notion that a short nucleotide sequence tag, isolated from a defined position within a transcript, contains enough information to uniquely identify it (Velculescu et al., 1995). Efficient analysis of transcripts is achieved by concatenation of the short sequence tags into a single clone to be sequenced (Velculescu et al., 1995). SAGE has an advantage over EST based methods, in being able to detect low-abundance transcripts (Sun et al., 2004) as it overcomes the obstacle of redundancy present in EST based methods, making it more likely to uncover novel transcripts (Chen et al., 2002b).

In SAGE, as in cDNA library construction, double-stranded cDNA is synthesised from mRNA, using OligodT primers, from the tissue/material of interest. In SAGE the
cDNA is cut with a 4bp anchoring restriction enzyme NlaIII, which would be expected to cleave most transcripts at least once (Figure 1.5) (Velculescu et al., 1995). The cDNA is divided into two pools and each ligated to a different adapter containing a type IIS restriction enzyme priming site for MmeI. Type IIS restriction enzymes (tagging enzymes) cleave at a defined distance downstream of the recognition site (Tucholski et al., 1995). This digestion leaves the adapter attached to a small sequence tag of the original transcript. The two pools are combined at this stage and ligated to form ditags (Velculescu et al., 1995). Production of uniform ditags allows amplification of the sample without introducing PCR bias. Cleavage of the PCR product with the original anchoring enzyme allows the ditags to be isolated and adapter molecules removed (Velculescu et al., 1995). The resulting ditags are ligated to form concatemers to be cloned into the vector pZErO-1 and sequenced. During sequence analysis the use of the anchoring enzyme site for ligation acts as a punctuation boundary for each tag (Velculescu et al., 1995).

Since it was first described, several modifications to the SAGE method have been suggested to improve yield and transcript identification in SAGE libraries (Powell, 1998; Velculescu et al., 2000; Zhang and Gilles, 2003). A major theme among these modifications has been to improve the accuracy of tag identification. A different restriction endonuclease has been used to produce longer tags of 21bp in LongSAGE (Saha et al., 2002; Zhang and Gilles, 2003) and 26bp in the SuperSAGE methods (Matsumura et al., 2003). The most recent and comprehensive review of the SAGE method describes Robust-LongSAGE (RL-SAGE), in which major improvements to the protocol overcome significant technical difficulties including low cloning efficiency and small insert sizes (Gowda et al., 2004).
Figure 1.5 Serial Analysis of Gene Expression

Modified protocols have been described to reduce the amount of mRNA required for cDNA synthesis (Velculescu et al., 2000). To produce sufficient ditags for library construction, these modifications have relied on additional PCR amplifications, which
may introduce bias in gene expression patterns (Gowda et al., 2004). In addition, a large quantity of mRNA can lead to incomplete digestion of cDNA which may generate multiple tags from the same transcript making it difficult to calculate gene expression from tag abundance (Gowda et al., 2004). Modifications present in the RL-SAGE protocol allow efficient libraries to be constructed with limited starting material, as little as 50ng compared with 2-5μg used in conventional and LongSAGE protocols, by extending the incubation times for cDNA digestion, cDNA adapter ligation and ditag ligation (Gowda et al., 2004). Improvements in ditag amplification and gel purification, including steps to decrease linker contamination as recommended by Powell (1998), have increased the cloning efficiency of subsequent steps. Partial digestion of concatemers, suspected to circularise during ligation, significantly improves the efficiency of cloning and increases insert sizes of clones (Gowda et al., 2004). Critical improvement of SAGE has reduced the time required for library construction by avoiding colony PCR screening to remove empty clones (Gowda et al., 2004). Crawford et al. (2005) provide a method for ensuring consistent production of high-molecular weight SAGE concatemers. They demonstrated that the concentration of PCR amplified ditag molecules present in concatemerisation and partial digestion reactions affects the proportion of low-molecular weight contaminants present in the resulting library (Crawford et al., 2005).

SuperSAGE, which uses a type III restriction endonuclease EcoP15l to isolate 26bp tags, has been used to compare two interacting organisms in a host-pathogen relationship by unambiguous gene-to-tag assignment (Matsumura et al., 2003). Unlike previous pathogen transcript profiles that rely on the use of artificial membranes to prepare suitably uncontaminated experimental material (Thomas et al., 2002),
SuperSAGE will be useful for understanding the real molecular mechanisms of both interacting species especially with the advantage of completely sequenced host and pathogen genomes (Matsumura et al., 2003).

SAGE studies have produced vast numbers of tags that cannot be matched to known genes, especially when there is a lack of genomic sequence data for the organism of interest (Dean and Lorenz, 2004; Saha et al., 2002). RT-PCR techniques have been developed to further analyse unknown tags (Chen et al., 2002a; Chen et al., 2000; Lee et al., 2002; Matsumura et al., 2003; van den Berg et al., 1999). The GLGI technique (Generation of Longer complementary DNA [cDNA] fragments from SAGE tags for Gene Identification), uses a SAGE tag sequence as the sense primer, and an anchored oligo(dT) primer as the antisense primer to amplify the original cDNA template from which the SAGE tag was derived (Chen et al., 2002a; Chen et al., 2000; Lee et al., 2002). Unlike the short tags produced by conventional SAGE which prove inadequate in efforts to clone unknown genes (Lorenz and Dean, 2002), the longer tags from SuperSAGE have been established as useful primers for the 3’RACE technique (Rapid Amplification of cDNA Ends) to produce longer cDNA fragments from unknown tags (Matsumura et al., 2003).

1.6.2 SAGE in plants

SAGE was first described in plants by Matsumura et al. (1999). In this study of rice seedlings the 10 122 tags analysed included 5 921 unique tag sequences expressed in rice seedlings (Oryza sativa L.). While only 23.1% of these tags matched to rice cDNA or EST sequence databases, most of the highly expressed tags could be
identified, and matched primarily to housekeeping genes. This quantitative study of
gene expression in rice, further applied the SAGE method to identify genes that were
differentially expressed between anaerobically treated and untreated seedlings.
Twenty-four genes, including eighteen anaerobically induced genes and six that were
repressed, showed expression at significantly different levels in each tag pool
(Matsumura et al., 1999).

Lorenz and Dean (2002) produced the first large scale SAGE libraries in a higher
plant. They produced a combined data set of more than 150,000 tags with
approximately 42,000 different tag sequences using libraries constructed from
lignifying xylem in the crown and base portions of loblolly pine (Pinus taeda). In this
study, it was decided that the omission of singleton tags would not affect the analysis
of differentially expressed tags between the SAGE data sets. The possibility that a tag
seen only once could represent a PCR artefact or sequencing error makes it difficult to
determine true unique genes (Lorenz and Dean, 2002). Zhang et al. (1997) and
Velculescu et al. (1997) however, estimate the sequence error rate for SAGE tag
concatemers as approximately 0.7% or a tag error of 6.8% which would equate to at
least 93% of unique tags representing true transcripts.

The SAGE method was used to analyse gene expression patterns in rice leaf and seed,
and a significant number of tags coding for antisense transcripts were found (Gibbings
et al., 2003). SAGE technology allows the position and orientation of a tag to be
determined with respect to an mRNA molecule, thus allowing the detection of
antisense transcription (Gibbings et al., 2003; Patankar et al., 2001). In Gibbings et al.
(2003), two of the 100 most abundant tags were derived from the non-coding strand in
the 5’ direction, and antisense tags were even more frequent at lower abundances. Many of the tags of low abundance were found to be derived from antisense transcripts of more abundant genes, indicating that the gene number generated by this SAGE analysis is an overestimation. However, there was no association between the frequencies of sense and corresponding antisense tags, indicating that the presence of antisense tags is not an artifact of the SAGE method (Gibbings et al., 2003).

Antisense transcription in plants is more widespread than previously thought and may represent an important mechanism for gene regulation (Gibbings et al., 2003; Terryn and Rouze, 2000). Antisense transcripts have a role in RNA silencing. Small antisense RNAs were first discovered in plants as a mechanism of defence against invading nucleic acids such as transgenes and viruses; however endogenous silencing pathways and microRNA molecules have been found to have widespread roles including gene regulation at transcriptional, RNA stability and translational levels (Brodersen et al., 2006; Bartel, 2004).

SAGE has been used to analyse the biological response of the transcriptome to abiotic stimuli (Jung et al., 2003). Comparison of the expression level of tags identified from cold-treated Arabidopsis leaves to normal leaves identified 272 differentially expressed genes (Jung et al., 2003). The total number of expressed genes in this study was notably reduced after cold stress, but the average copy number per gene was increased (Jung et al., 2003). Of the genes showing more than a ten fold difference in expression, 87 were newly identified as cold-stress related genes (Jung et al., 2003). To confirm the validity of the SAGE data, five putative cold-responsive genes were further analysed by northern blots. The expression data obtained by northern blot analysis were consistent with, and thus confirmed, the differential expression patterns
derived from SAGE (Jung et al., 2003). Several other SAGE studies have used northern blot analyses to validate the expression data produced by SAGE (Patankar et al., 2001; Trendelenburg et al., 2002).

The model plant Arabidopsis has been analysed in many SAGE studies (Fizames et al., 2004; Jung et al., 2003; Lee and Lee, 2003). Lee and Lee (2003) were able to reveal the groups of proteins and metabolic pathways likely to be active during Arabidopsis pollen development. A SAGE library was also constructed from cold stressed pollen of Arabidopsis (Lee and Lee, 2003). The expression of known cold-inducible genes, all of which are highly induced in the Arabidopsis leaf under cold stress (Jung et al., 2003), showed no significant increases in the pollen library (Lee and Lee, 2003). Microarray based analysis was used to support this observation, and the study suggests that the inability to induce expression of genes important for cold acclimation is linked to the cold sensitivity of pollen (Lee and Lee, 2003).

Fizames et al. (2004) conducted a large investigation of the Arabidopsis root transcriptome by SAGE. Taking advantage of the complete sequence of the genome, the study used a computer-based approach to characterise expression of more than 3000 genes that did not exist previously in EST or cDNA databases (Fizames et al., 2004). EST clusters or cDNA sequences had previously been used to identify the gene corresponding to the SAGE tag transcript (Gibbings et al., 2003; Matsumura et al., 1999). This has been a major limitation of SAGE tag to gene assignment as not all transcripts exist in cDNA and EST databases and many tags are left unmatched and unidentified (Fizames et al., 2004; Lash et al., 2000; Pleasance et al., 2003). Despite having the full genome sequence a large proportion of tags in the Arabidopsis root
transcriptome study could not be assigned to any gene (Fizames et al., 2004). The unassigned tags could be artifactual, however erroneous tags are discarded by the exclusion of single tags and duplicate ditags, which suggests that unassigned tags originate from transcripts with incorrect or absent virtual tags in the computer generated tag reference library (Fizames et al., 2004).

SAGE has been employed to investigate barley in whole seed and through a 120 hour malting time course (White et al., 2006) and in a direct comparison to the barley Affymetrix GeneChip (Ibrahim et al., 2005). The most abundant and most significantly up-regulated LongSAGE tag in the study by White et al. (2006), matched to two previously reported genes, B22EL8 and pZE40, which have been described as metallothionein-like proteins with proposed roles in synthesis and transport of metabolites (Klemsdal et al., 1991; Smith et al., 1992). Ibrahim et al. (2005) also reported pZE40 as the most abundant SAGE tag identified in their study. They generated 82,122 tags from two libraries representing whole developing barley caryopsis and dissected embryos, at 12 days post-anthesis (Ibrahim et al., 2005). Difficulties associated with multiple tags matching to one transcript were reported. Sources of multiple tags per transcript evolve from alternative splicing, incomplete digestion with the anchoring enzyme during library construction, alternative polyadenylation sites and possibly non-specific poly dT anchoring of reverse transcription during cDNA synthesis (Ibrahim et al., 2005; Pauws et al., 2001; Welle et al., 1999).

1.7 Real-time reverse transcription PCR
Real-time PCR is an efficient method for determining DNA quantity as it combines DNA amplification and detections steps in one assay. This eliminates the need for post PCR processing steps such as gel electrophoresis, Southern blotting and DNA sequencing. Reverse transcription PCR (RT-PCR) involves the use of reverse transcriptase to synthesise complementary DNA (cDNA) to an mRNA template (Snustad and Simmons, 2000). Monitoring the amplification reaction in real time allows the accurate determination of template quantity in the starting material. Amplification is monitored using fluorescent molecules. There are effectively two types of fluorescent chemistry, template-specific fluorescent probes and non-specific DNA binding fluorophores (e.g. SYBR Green) (Bustin, 2005b). DNA binding fluorophores fluoresce when incorporated into double stranded DNA. The fluorescent signal produced by either chemistry is directly proportional to the amount of amplified product present. The linear relationship between the fluorescence output and sample concentration allows the amount of template present at the start of the reaction to be calculated (Bustin, 2005a). Real-time RT-PCR results are often referred to in terms of the cycle threshold (Ct). This refers to the cycle number at which a PCR product is first detected. Ct is determined in the exponential phase of the reaction and is inversely proportional to target copy number. Therefore the higher the starting copy number, the sooner the PCR product is detected and the lower the Ct value (Bustin, 2005a).

Real-time RT-PCR reactions are a simple, specific, sensitive and highly reproducible method for the detection of RNA. The most important determining factor in the reproducibility of RT-PCR results is the RNA template quality (Bustin, 2005b; Fleige
and Pfaffl, 2006). The introduction of lab-on-chip technologies such as the Bioanalyser allows the accurate assessment of RNA quality and quantity and provides a statistic for this assessment, the RNA Integrity Number (RIN) (Fleige and Pfaffl, 2006).

Real-time RT-PCR results can be obtained by relative or absolute quantification. Relative quantification represents the change in mRNA levels of a gene of interest relative to the levels of a control mRNA, for example, a housekeeping gene such as GAPDH (Bustin, 2005b). Target Ct values are compared to the internal control and expressed as a ratio (Bustin, 2005b). Absolute quantification is an attempt to determine absolute copy number in terms of amount of total RNA or weight of tissue in extraction based on a standard curve. A standard dilution series of known concentrations is used to generate a standard curve by plotting Ct versus known copy number (Bustin, 2005b). The copy numbers of unknown samples can be determined by the position of corresponding Ct values on the curve.

Real-time RT-PCR has been used extensively in the biomedical field and the utility of the technology in the development of diagnostic assays is rapidly increasing (Mackay et al., 2002; Valasek and Repa, 2005). Real-time RT-PCR has also proven to be a valuable research tool to measure gene expression levels for a wide variety of genes in various plant species; for example, Arabidopsis (Klok et al., 2002), canola (Yang et al., 2007), pea (Ayele et al., 2006), peanut (Luo et al., 2005) and barley (Burton et al., 2004; Dean et al., 2002; Liu et al., 2005; Potokina et al., 2006; Wolbang et al., 2004).
1.8 Gene Expression in Barley

In recent times, microarray technology has been exploited to examine gene expression in barley (Close et al., 2004; Potokina et al., 2002; Sreenivasulu et al., 2002; Sreenivasulu et al., 2004; Watson et al., 2001). Most notably a 22K Barley1 GeneChip probe array has become publicly available (Close et al., 2004). This cereal genomics resource, a model for plants without a fully sequenced genome, allows parallel expression analysis and gene discovery in disease defence, abiotic stress response, evolutionary diversity and development (Close et al., 2004). Microarray content was derived from a worldwide contribution of 350 000 ESTs from 84 cDNA libraries, together with 1 145 non-redundant gene sequences from NCBI (Close et al., 2004). The parameters used in designing the array allow a high degree of specificity to enable very similar genes, such as gene family members, to be differentiated (Close et al., 2004). Other features include the use of multiple probe pairs per sequence to reduce false positives and enable expression quantification, and the use of probes representing housekeeping or conserved genes to facilitate comparative studies among grasses (Close et al., 2004). A public database allowing access to the data makes possible the integration of expression profiling experiments from around the world, links annotations and alignments with data from other plant databases, and the option to query a gene of interest to discover the conditions under which it showed significant change (Close et al., 2004). The 22K Barley1 GeneChip is a useful tool for the analysis of barley and other cereal genomes. The GeneChip has been used to profile barley-stem rust interactions (Zhang et al., 2008), maturation, desiccation and germination (Sreenivasilu et al., 2008), genotyping (Walia et al., 2007), Giberellin
and abscisic acid responses (Chen et al., 2006), and most recently to analyse malting barley phenotypes (Lapitan et al., 2009).

To investigate the regulation of gene expression during germination of barley grains, Potokina et al. (2002) developed a cDNA macroarray with 1 440 spotted clones derived from cDNA libraries of developing caryopsis, roots and etiolated leaves. It was their goal to analyse the spatial and temporal patterns of gene expression in embryo, scutellum and endosperm tissue, including the aleurone, during different stages of germination (4, 12, 36, 52 hours after the start of imbibition) (Potokina et al., 2002). The scutellum was considered to be a transitional tissue between embryo and endosperm from a spatial and functional point of view and was used as a comparator for each (Potokina et al., 2002). Genes expressed in the endosperm were clustered according to three distinct behaviours, up-regulated, down-regulated and constant expression. Significant down-regulation of genes appears to be specific to the endosperm, as it was not observed in the embryo or scutellum (Potokina et al., 2002).

A considerable portion of genes are up-regulated in each of the three tissues during the late stages of germination (36 and 52 hours after imbibition), while a few genes identified only in the scutellum were up-regulated only in the final stage 52 hours after imbibition (Potokina et al., 2002). Most of the genes preferentially expressed in the scutellum correspond with expected functions of the tissue, including production of hydrolytic enzymes for storage protein and starch catabolism (Potokina et al., 2002). This study provides some information about the metabolic processes of germination. The array was derived mainly from a cDNA library of developing barley seeds however, and to perform a more detailed analysis of germination would require the incorporation of transcripts from germinating barley grains in the array design.
Sreenivasulu *et al.* (2002; 2004) has produced cDNA macroarrays to infer molecular physiology from maternal pericarp and filial embryo/endosperm tissues of developing barley grains. The gene expression profiles in developing barley caryopses were studied from fertilization to storage phase and it was discovered that expression in the pericarp mainly encodes enzymes involved in carbohydrate and lipid metabolism while expression in the embryo is related to degradation and processing of proteins or the accumulation of starch (Sreenivasulu *et al.*, 2002). An intermediate phase separating the cell division and accumulation processes associated with grain development was identified, during which a considerable rise in gene expression related to photosynthesis and energy production was observed (Sreenivasulu *et al.*, 2004). Like the array by Potokina *et al.* (2002), this study was very useful; however it provides no expression data specific to the germination of barley grains.

In a comparative study of SAGE and Affymetrix array technologies, Ibrahim *et al.* (2005) investigated gene expression in barley caryopsis and dissected embryos at 12 days post-anthesis. The Barley 1 Affymetrix GeneChip contains 22 791 probe sets, approximately half of which were given a present call in both tissues \( p \leq 0.05 \) (Ibrahim *et al.*, 2005). Similar trends were observed between the SAGE and GeneChip scores with only a small proportion of transcripts accounting for the majority of signal. The 100 most abundant transcripts determined by each method were identified and compared, and strong agreement was found between the SAGE and GeneChip technologies (Ibrahim *et al.*, 2005).
Germination rate plays an important role in determining malt quality. Watson et al. (2004; 2001) used cDNA microarrays to examine gene expression in the embryos of germinating barley grain. The expression profiles of barley embryos (including scutellum) from 4, 8, 12, 24, 48, 72 and 96 hours post-imbibition were compared to a reference expression profile of the 24 hour post-imbibition time-point in a series of microarray experiments (Watson and Henry, 2004). Genes with similar gene expression patterns were separated into groups of potentially similar function. Only 514 of the total data set (4 945 clones) could be classified by distinctive patterns of expression (Watson and Henry, 2004). Of this subset, 16.9% of clones showed ten fold or greater differences in expression prior to 24 hours. More than 50% of the subset however, showed a ten fold or greater difference in expression in the period 24-96 hours post-imbibition and a high proportion of these were down-regulated (Watson and Henry, 2004). One of the few clones showing sustained up-regulation in the period after 24 hours post-imbibition was the 23kDa stress-response Jasmonate-Induced Protein (JIP-23). JIP-23 most likely plays a role in mediating osmotic stress in germinating barley embryos (Watson and Henry, 2004). This work provides a useful beginning for the analysis of gene expression in germinating barley embryos by determining temporal differences in gene expression patterns over the first four days post-imbibition. However, a comprehensive analysis of genetic components and mechanisms for developmental, regulatory and response processes in whole germinating barley has yet to be investigated in a full genome examination.

Malting is a highly controlled process that ensures the efficient release of fermentable sugars from the barley grain. The biochemical basis of the malting process has been studied in considerable depth. This knowledge has been used to optimise the quality
and quantity of output from malt barley. The genetic basis of the malting process however has not yet been examined in much detail. Modern molecular biology techniques enable the analysis of thousands of transcripts simultaneously. The high-throughput techniques of SAGE and Microarray will be applied to a time course of malting barley samples. This study aims to provide large data sets for the basis of accurate gene expression analysis with widespread coverage of the malting barley transcriptome.

1.9 Aims and Objectives

Malting is a highly controlled process that ensures the efficient release of fermentable sugars from the barley grain. The biochemical basis of the malting process has been studied in considerable depth. This knowledge has been used to optimise the quality and quantity of output from malt barley. However the genetic basis of the malting process has not been studied in detail. Modern molecular biology techniques enable the analysis of thousands of gene transcripts simultaneously, providing an opportunity to improve knowledge of the malting process at the gene level. This study aims to determine what genes are expressed during the malting of barley and the temporal patterns in which they are expressed and to explore different gene expression analysis technologies.

The high-throughput techniques of SAGE and Microarray will be applied to a time course of malting barley samples to investigate the genes that control and regulate this process. This time course includes samples of dry dormant seeds, post-steeped grain, and grain germinating at time points up to 120 hours post-steeping when the shoots
and roots are emerging from the grain and the endosperm is completely modified. This study aims to provide primary resources of transcription data in the form of a large malting barley LongSAGE tag database and a CombiMatrix 12K Custom ElectraSense™ barley microarray chip.

Chapter one of this thesis is a review of the relevant literature on malting and gene expression analysis. Chapter two describes the generation of LongSAGE libraries from eight time points along a time-course of malting barley. SAGE is an open system capable of capturing global transcript expression data that does not rely on known sequence information, a disadvantage of most other gene expression technologies. SAGE tags are isolated from a defined position within a transcript which enables tags to be annotated by similarity matching to databases of known sequences. Analysis of the most abundant LongSAGE tags highlights the functional groups of dominant mRNA transcripts. This is the largest study to date of global gene expression during grain germination.

Chapter three examines the identification and real-time RT-PCR validation of LongSAGE tags showing differential expression across the malting time course. Transcripts identified with roles in metabolism, particularly in carbohydrate and protein modification, were selected as having a direct relationship to barley seed germination and malting. Real-time RT-PCR was conducted using RNA extracted from the same malting grain samples used in the LongSAGE study. Real-time RT-PCR experiments were performed at four time points corresponding to the LongSAGE study and the mRNA abundance levels observed using real-time RT-PCR compared with the LongSAGE data.
Chapter four describes a microarray study in which the unique tags of the 24 hours post-steeping LongSAGE barley library were used to design a CombiMatrix 12K Custom ElectraSense™ array. Chips were hybridised with cRNA from two important malting time points. The biological sample used to conduct this experiment was the same as that used for the LongSAGE and Real-time RT-PCR experiments. The probes identified as differentially expressed between time points in the malting barley time course are compared between the three gene expression technologies; ElectraSense™ microarray, LongSAGE and real-time RT-PCR.

Chapter five is a general discussion that summarises the findings of the thesis. It also indicates further directions of study and the potential applications of the resources developed from this work.

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Chapter Two

Abundant transcripts of malting barley identified by serial analysis of gene expression (SAGE)

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Chapter 3

Differential LongSAGE tag abundance analysis in a barley seed germination time course and validation with relative real-time RT-PCR

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Chapter 4

Transcriptome analysis of malting barley; comparison of Microarray, LongSAGE and real-time RT-PCR

4.1 Abstract

A 12K CombiMatrix Electrochemical CustomArray was employed to compare gene expression of the Australian malting barley variety Tallon at 12 and 48 hours post-steeping. One hundred and thirteen probes show a greater than ten fold difference in signal intensity, with 64 preferentially expressed at 12 hours post-steeping and 49 at 48 hours post-steeping. The probes with up-regulated expression at 48 hours post-steeping include transcripts associated with stress response, pathological defence and metabolism. The 23kDa Jasmonate-induced protein, JIP-23, is noted as the probe with the highest and fifth highest fold up-regulation from 12 to 48 hours post-steeping. The CombiMatrix ElectraSense™ microarray signal intensity values were compared to expression data from LongSAGE and real-time RT-PCR experiments conducted on the same malting barley samples. The fold change expression data correlated well between the three gene expression technologies. The CombiMatrix ElectraSense™ microarray was the most cost effective and reliable methods for measuring thousands of transcripts simultaneously with the ability to compare many different samples with multiple replicates and relatively lower cost.
4.2 Introduction

Malt barley is the major raw material apart from water used by the brewing industry. The malting of barley involves the controlled germination of the barley grain to promote expression of genes altering the grain composition and generating enzymes for the brewing process. Here we compare the use of different tools to establish the underlying patterns of gene expression during the malting process. High throughput gene expression technologies such as SAGE (Velculescu et al., 1995) and microarrays (Schena et al., 1995) allow us to examine large numbers of gene transcripts in specific samples of malt barley. This provides the opportunity to study changes in gene expression in particular varieties, under various conditions and at specific time points. A time course of malting barley variety Tallon was examined by LongSAGE (White et al., 2006). Eight LongSAGE libraries were constructed from seed sampled at 0, 12, 24, 48, 72, 96 and 120 hours post-steeping and also from dry seed (White et al., 2006). Statistical analysis revealed a set of 57 tags that show significant differential expression across the time course (White et al., 2006). Eight of these were identified as candidates for real-time RT-PCR validation. Real-time RT-PCR was carried out for each of the genes of interest on four of the seed samples analysed in the LongSAGE study. Real-time RT-PCR analysis successfully confirmed the patterns of expression seen in the LongSAGE study (White et al., 2006).

Microarray technology allows the simultaneous expression analysis of large sets of genes of known sequence (Schena et al., 1995). Differential gene expression has most commonly been measured using two-colour hybridisation. Two mRNA sources are prepared to create two cDNA probes with different fluorophores. Samples of known
concentration are used to calibrate the fluorescence, which is captured by scanning the array with an imaging system (Schena et al., 1995). The amount of fluorescence represents the amount of transcript expressed in each sample.

CombiMatrix have developed an electrochemical detection system for oligonucleotide arrays (Ghindilis et al., 2007; Roth et al., 2006). The CombiMatrix system involves a semiconductor matrix of 12,544 individually addressable platinum microelectrodes on which different oligonucleotides can be simultaneously synthesised via digital control (Ghindilis et al., 2007; Roth et al., 2006). The electronics used for the oligonucleotide synthesis are subsequently utilized for the detection of redox active chemistries associated with hybridised target molecules (Ghindilis et al., 2007; Roth et al., 2006). Biotin bound target molecules are labelled with a streptavidin horse radish peroxidase (HRP), the array is exposed to the substrate tetramethylbenzidine (TMB) and hydrogen peroxide, oxidised TMB is reduced at the electrode surface which generates an electrochemical signal that is read with the ElectraSense™ microarray reader (Ghindilis et al., 2007; Roth et al., 2006).

CombiMatrix have demonstrated that the ElectraSense™ system is a sensitive and reproducible detection platform with effective reusability and performance at a level that is at least competitive with the performance of fluorescence based techniques (Ghindilis et al., 2007; Roth et al., 2006). The CombiMatrix ElectraSense™ system also offers advantages over existing technologies including lower reagent and equipment costs, a digital data format that does not require image adjustments, the absence of negative effects related to fluorescent dye quenching, ozone-mediated
signal degradation and autofluorescence, and the potential for a high degree of automation (Ghindilis et al., 2007; Roth et al., 2006).

Microarray analysis is an effective tool for plant functional genomics and has already been successfully used to explore different aspects of the barley transcriptome (Close et al., 2004; Potokina et al., 2004; Potokina et al., 2002; Sreenivasulu et al., 2002; Sreenivasulu et al., 2004; Watson and Henry, 2005). The recent development and public availability of the Affymetrix 22K Barley1 GeneChip is a major contribution to cereal genomics resources (Close et al., 2004). With content derived from a worldwide contribution of 350 000 ESTs from 84 cDNA libraries the array design parameters included high specificity for very similar genes, such as gene family members, so that they might be differentiated (Close et al., 2004). In a comparative study of SAGE and Affymetrix array technologies, Ibrahim et al. (2005) investigated gene expression in barley caryopses and dissected embryos at 12 days post-anthesis using the 22K Barley1 GeneChip. The Affymetrix GeneChip contains 22 791 probe sets, approximately half of which were given a present call in both the caryopsis and dissected embryo tissues ($p \leq 0.05$) (Ibrahim et al., 2005). The 100 most abundant transcripts determined by each method were identified and compared, and strong agreement was found between the SAGE and GeneChip technologies (Ibrahim et al., 2005).

This study employs a 12K CombiMatrix Electrochemical CustomArray to compare gene expression in the Australian malting barley variety Tallon at 12 and 48 hours post-steeping. The array was designed using unigene and contig sequences that match to LongSAGE tags from Tallon at 24 hours post-steeping.
4.3 Materials and Methods

4.3.1 Array design

CombiMatrix 12K Custom ElectraSense™ arrays were designed based on data from a LongSAGE library constructed from malt barley at 24 hours post-steeping (White et al., 2006). LongSAGE tags were blasted to the *Hordeum vulgare* Gene Index database (Version 9.0, release September 15, 2004 The Gene Index Databases, Dana Farber Cancer Institute, Boston, MA USA http://compbio.dfci.harvard.edu/tgi/cgi-bin/tgi/gimain.pl?gudb=barley). Probes were designed using CombiMatrix CustomArray® Software based on the Tentative Consensus sequences (TC’s) and singletons that were returned as perfect match hits to the LongSAGE tags. The optimal oligonucleotide length for the CombiMatrix 12k CustomArray is 35-40 base pairs in length. The array includes 6,390 probes in the sense orientation, 1,417 antisense probes and 3,030 features representing 1,515 probes in both orientations. This equates to 10,837 different probes on the array. There are 12,544 total features on a CombiMatrix 12k CustomArray® with 460 features on an ElectraSense™ array used for quality control, 12,000 user defined probes and the remainder being blank features. Eight bacterial QC probes were used for the hybridisation of complementary spike-in oligonucleotides labelled at the 5’ end with biotin to enable determination of the linearity of concentration versus signal.
4.3.2 Target preparation, array hybridisation, scanning and stripping

Tissue was collected, RNA extracted and samples from 12 and 48 hours post-steeping used from studies previously described (White et al., 2006; White et al., 2008). Target RNA was amplified and labelled with a Kreatech RNA ampULSe: Amplification and Labelling Kit for CombiMatrix arrays with Biotin ULS (Cat. no. EA-026; Kreatech Biotechnology, Amsterdam, The Netherlands). All steps were carried out as per the protocol with aRNA fragmentation carried out using reagents from Ambion (Cat. no. AM8740; Ambion, Austin, TX, USA). Hybridization and electrochemical detection was carried out as per the CombiMatrix Protocol for ElectraSense™ 12K Microarray Hybridization and Electrochemical Detection PTL007 and the CombiMatrix ElectraSense™ Detection Kit (Cat. no. 610027; CombiMatrix, Mukilteo, WA, USA). Arrays were scanned using an ElectraSense™ reader from CombiMatrix. Arrays were stripped and rehybridised two to three times each using the CombiMatrix ElectraSense™ Stripping Kit for 12K (Cat. no. 610029; CombiMatrix) as per the CombiMatrix Stripping and Preparation of ElectraSense™ 12K Microarrays for Re-hybridisation protocol PTL003.

Two microarray slides were used to conduct all of the hybridisations for this series of experiments. The RNA samples were hybridised to the arrays in an alternating pattern. Slide one was hybridised with RNA from 12 hours post-steeping and slide two was hybridised with RNA from 48 hours post-steeping, the slides were stripped and prepared for rehybridisation. The slides were then swapped so that slide one was hybridised with RNA from 48 hours post-steeping and slide two was hybridised with RNA from 12 hours post-steeping. This pattern was repeated and in total, slide one
was hybridised twice with RNA from 12 hours post-steeping and twice with RNA from 48 hours post-steeping. Slide two was hybridised twice with RNA from 48 hours post-steeping but only once with RNA from 12 hours post-steeping. Each hybridisation was scanned so that at least two visually acceptable scans were produced.

4.3.3 Quality control, background correction and data normalisation

The image synthesised from signal data for each hybridisation by the ElectraSense™ software was analysed by visual inspection (Figure 4.1). As an initial step in quality control of hybridisations, the concentration of the spike in controls was plotted against the log of the signal of those controls to determine if an approximate linear relationship resulted (Data not shown). Data for the final analysis did not include the scans from the first hybridisation of slide one to RNA from 48 hours post-steeping or the spike-in controls from any of the hybridisations. Data files for replicate scans and hybridisations were merged and average probe signal intensities used to reduce experimental bias. The data were processed using an RMA (Robust Multi-Array Average) background correction, normalised using quantile normalisation and the data from features of replicate probes were averaged.
Figure 4.1 Representative ElectraSense™ image of slide one hybridised with RNA from 12 hours post-steeping. The spike-in controls are present in five rows spread evenly across the chip and visible due to their relatively higher signal intensity.
4.4 Results

4.4.1 Microarray analysis of malting barley

Two samples of malting barley from important and physiologically contrasting time points, 12 and 48 hours post-steeping, were chosen for a microarray gene expression analysis study. Microarrays were designed based on SAGE tag results from the 24 hours post-steeping LongSAGE library described by White et al. (2006) and malting barley RNA samples from related experiments were used to enable direct comparisons with the relevant results. This analysis assumes that the re-use of slides by stripping and re-hybridising, effects all probes evenly. Scatter plots comparing the probe intensities between replicate scans and between hybridisations on different chips produced very consistent results (Figure 4.2). The median correlation co-efficient for the probe intensities between hybridisations at the M2 time point was 0.931 and for the M4 time point was 0.968 with a very high level of reproducibility observed between replicate scans (median 0.989).
Figure 4.2 Reproducibility of CombiMatrix ElectraSense™ Custom microarrays. Intensity data for each probe is plotted from (a) replicate scans of the same hybridisation, and (b) arrays of the same sample hybridised to different chips. The correlation coefficient (r) for plot (a) is 0.998 and for plot (b) 0.978.

The data from both the LongSAGE and CombiMatrix ElectraSense™ microarray experiments produce a very similar signal range from highest to lowest signal intensity for the samples, 12 and 48 hours post-steeping (Table 4.1). The patterns between the technologies are also very similar. For example at each time point, analysed by each technology, the lowest signal intensity value represents approximately 0.002% of the highest intensity value. However the microarray results appear to provide a better range of signal intensity at low levels. The signal intensity of microarray probes observed at probe numbers 1000 and 2000, in a rank of intensity, represent approximately 0.025% and 0.01% of the total signal on the array respectively. The LongSAGE tag frequency of tag number 1000 is only 0.015% of total tags. The SAGE technology appears more limited in its ability to differentiate expression intensity at the lower end of the frequency range. This limitation may be somewhat overcome by more extensive sequencing of each SAGE library.
Table 4.1 Signal range observed in normalised array data at the probe level and LongSAGE tag count data

<table>
<thead>
<tr>
<th></th>
<th>12 hours post-steeping</th>
<th>48 hours post-steeping</th>
<th>LongSAGE Signal range</th>
<th>12 hours post-steeping</th>
<th>48 hours post-steeping</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total signal intensity</td>
<td>7,303,233</td>
<td>7,318,885</td>
<td>20,042</td>
<td>20,251</td>
<td></td>
</tr>
<tr>
<td>of all probes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Highest intensity value</td>
<td>22,704 (0.31%)</td>
<td>24,641 (0.34%)</td>
<td>499 (2.49%)</td>
<td>203 (1.00%)</td>
<td></td>
</tr>
<tr>
<td>Lowest intensity value</td>
<td>61.4 (0.0008%)</td>
<td>58 (0.0008%)</td>
<td>1 (0.005%)</td>
<td>1 (0.005%)</td>
<td></td>
</tr>
<tr>
<td>Signal intensity</td>
<td>1868.8 (0.0250%)</td>
<td>1854.3 (0.0253%)</td>
<td>3 (0.015%)</td>
<td>3 (0.015%)</td>
<td></td>
</tr>
<tr>
<td>of probe #1000 in rank</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>of intensity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Signal intensity</td>
<td>797.6 (0.0109%)</td>
<td>758.6 (0.0104%)</td>
<td>2 (0.01%)</td>
<td>2 (0.01%)</td>
<td></td>
</tr>
<tr>
<td>of probe #2000 in rank</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>of intensity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

While looking at the LongSAGE results alone, some differences between the time points can be observed. Similar total numbers of LongSAGE tags were analysed in each library, however 12 hours post-steeping has a much higher top ranking frequency count than 48 hours post-steeping. The highest count tag in the 12 hours post-steeping library accounts for almost 2.5% of the total tags analysed in that library. While the highest tag count in the 48 hours post-steeping library accounts for only 1% of total tags analysed. This may indicate that 12 hours post-steeping has a greater diversity of tags at higher frequencies while M4 has a greater diversity of tags at lower frequencies.

4.4.2 Probe identification for Microarray

Probe sequences were compared with complete coding sequences at the NCBI database for *H. vulgare*, and the NCBI database for all barley sequences using BLASTN, to assist in assigning a putative function. Annotations with sequences with perfect matches to probe sequences were assigned along with the original annotations from the TIGR gene indices. The ten microarray probes with the highest signal
intensity in each of the two samples can be compared (Tables 4.2 and 4.3). Five of these are common to the top ten of both malting barley time points. These include nucleic acid binding proteins, metallothionein and lipid transfer proteins, all of which are known to be associated with germination. It appears that those transcripts expressed highly early on remain high throughout germination. This pattern was also observed in the LongSAGE data (White et al., 2006)

Table 4.2 Ten probes with the highest Microarray signal intensity in malting barley at 12 hours post-steeping

<table>
<thead>
<tr>
<th>Probe sequence</th>
<th>TC number</th>
<th>Putative function</th>
<th>12 hours post-steeping signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCCCATATGCTATAGATGCTACCAGTTATGCAAGTTATGCCGAGTTTATGC</td>
<td>X60292.1 a</td>
<td>Lipid transfer protein</td>
<td>22704.75</td>
</tr>
<tr>
<td>TGGAACTATCGCTGTCGTTCACTCTGGAATCAGTTTGTGTG</td>
<td>TC130715</td>
<td>Gamma hordothionin</td>
<td>17054.01</td>
</tr>
<tr>
<td>CCCATGAGACATGCTGTTTCACTCTTCTACAGTTAATGCAATC</td>
<td>TC130715</td>
<td>Gamma hordothionin</td>
<td>16871.29</td>
</tr>
<tr>
<td>CAGATCGTCTTCCCTATCCATCCCTCTCTATGCTCTTACC</td>
<td>Z48624.1 a</td>
<td>Glycine-rich RNA binding protein</td>
<td>16685.38</td>
</tr>
<tr>
<td>CTTAGTTGCTGTCCTCTCGTGCCTCCCCTCAATCTACTAC</td>
<td>BM816569</td>
<td>Metallothionein</td>
<td>15581.15</td>
</tr>
<tr>
<td>TACCGTCTCTACGCCAGCAAGTTATATCTACCTCTCTGTGTGT</td>
<td>TC138719</td>
<td>Nucleic acid binding protein</td>
<td>15277.74</td>
</tr>
<tr>
<td>TTTGCCCCTACTATGCTGTCCTCGATGATGATTCCCAC</td>
<td>TC139387</td>
<td>Oleosin</td>
<td>14919.17</td>
</tr>
<tr>
<td>GTGGAGTTGAGTTGCTAGTTAATCGCTGCGGACTAACCC</td>
<td>X60292.1 a</td>
<td>Lipid transfer protein</td>
<td>14408.44</td>
</tr>
<tr>
<td>TACCGTCTCTACGCCAGCAAGTTATATCTACCTCTATGCTG</td>
<td>U22951.1 a</td>
<td>Gamma-thionin</td>
<td>13491.41</td>
</tr>
<tr>
<td>CATGTGCTGGTGCTGGTCTGCGGTCGCTCTAATGAG</td>
<td>BQ760074</td>
<td>Unclassified</td>
<td>13373.96</td>
</tr>
</tbody>
</table>

* NCBI accession number.

Table 4.3 Ten probes with the highest Microarray signal intensity in malting barley at 48 hours post-steeping

<table>
<thead>
<tr>
<th>Probe sequence</th>
<th>TC number</th>
<th>Putative function</th>
<th>48 hours post-steeping signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTTAGTTGCTGTCCTCTCGTGCCTCCCCTCAATCTACTAC</td>
<td>BM816569</td>
<td>Metallothionein</td>
<td>24640.66</td>
</tr>
<tr>
<td>TCTCCTCGAGCAACCGACAGGAGATGTC</td>
<td>Z34269.1 a</td>
<td>Beta-ketoacyl ACP synthase</td>
<td>18072.14</td>
</tr>
<tr>
<td>TACGGCTGACCAAGTTATCTGACTTTCGCTGTGTG</td>
<td>TC138719</td>
<td>Nucleic acid binding protein</td>
<td>17633.50</td>
</tr>
<tr>
<td>CAGATAGTTGCTCCACCATGCTACGCTCTACCTAATGCC</td>
<td>Z48624.1 a</td>
<td>Glycine-rich RNA binding protein</td>
<td>17429.87</td>
</tr>
<tr>
<td>CTTAGTCTCTAGCTATGCGCTACAGTTATGCGCTCTATG</td>
<td>X60292.1 a</td>
<td>Lipid transfer protein</td>
<td>17341.20</td>
</tr>
<tr>
<td>GCTACGTTAGATGCGCTACGCTCTGCTCTA</td>
<td>U49482.1 a</td>
<td>Low temperature responsive</td>
<td>15364.16</td>
</tr>
</tbody>
</table>
Figure 4.3 shows a comparison between the two samples of all probes following normalisation. There is no strong bias towards expression at either time point with this array design. Most expression signals show a less than two fold deviation from the diagonal. 1 822 probes show a greater than two fold difference, 470 probes show a greater than four fold difference and 113 probes show a greater than ten fold difference between the samples. To focus on genes with the highest fold changes in expression from 12 to 48 hours post-steeping, those probes with signal intensities differing by a factor of ten or more were examined (Tables 4.4 and 4.5). Analysis of the probes with a greater than ten fold change in signal intensity revealed 64 probes preferentially expressed at 12 hours post-steeping (Table 4.4) and 49 probes preferentially expressed at 48 hours post-steeping (Table 4.5). The probes which appear to decrease in expression from 12 to 48 hours post-steeping include several transcription factors and stress-related genes. This pattern of expression could reflect several things; that the transcription factor is no longer required after 12 hours post-steeping, that the encoded protein is relatively stable or that these are transcripts remaining from grain fill and maturation. RNA transcripts vary greatly in their stability and time taken to degrade, with mRNA half-lives in higher plants of up to 24 hours (Aalen et al., 2001; Narsai et al., 2007).
Figure 4.3 Comparison of the normalized signal intensities obtained from barley CombiMatrix ElectraSense\textsuperscript{TM} 12K Custom microarrays. X axis represents normalised signal intensity at 12 hours post-steeping. Y axis represents normalised signal intensity at 48 hours post-steeping. Signals outside the diagonal lines differ by greater than a factor of two between the experiments.

In this study, the probes which show a preference for expression at 48 hours post-steeping are of the greatest interest. It can be assumed that genes with a significant increase in expression at this early stage of malting are significant to the underlying physiology of the malting process. Among the 49 probes with a ten fold or greater increase in expression from 12 to 48 hours post-steeping there are transcripts
associated with stress response, pathogen defence and metabolism. Stress response and pathogen defence transcripts are present as both a preventative and reactive measure to combat pathogen, osmotic, heat and other stresses imposed upon the germinating grain.

The probes with the highest and fifth highest fold change in signal from 12 to 48 hours post-steeping represent the stress response protein JIP-23 (23kDa Jasmonate-induced Protein). JIP-23 is one of several abundant jasmonate-induced proteins of different molecular masses and differing biological functions, the function of several of which is still unknown. The specific expression of JIP-23 mRNA has been noted in the coleoptile and phloem cells of the scutellum and is thought to be expressed in response to osmotic stress brought about by the transport of solutes from the endosperm to the germinating embryo (Hause et al., 1996). Watson et al. (2005) found that JIP-23 had a greater than ten fold increase in expression after 24 hours post-steeping using cDNA arrays. Glutathione-S-transferase, a protein also recognised as responsive to jasmonic acid, is up-regulated by approximately 13 fold at 48 hours post-steeping and has also been found to exhibit high expression in other studies (Sasaki et al., 2000; Potokina et al., 2002; Watson et al., 2005).

Other probes of interest with a ten fold or greater increase in expression at 48 hours post-steeping include Metallothionein, Serine/Threonine protein kinase, β-d-xylosidase, Glycolate oxidase, pZE40, Dehydroascorbate reductase, Xylose isomerase, α-methyltransferase and Aldose-1-epimerase. Dehydroascorbate reductase has been identified by statistical analysis as associated with the malt quality trait of mash viscosity (Potokina et al., 2004). Several of the genes found in this list match to
the 57 tags identified as significantly differentially expressed in the LongSAGE study of malting barley across time (White et al., 2006).

Table 4.4 Microarray probes with signal down-regulated more than 10 fold from 12 to 48 hours post-steeping.

<table>
<thead>
<tr>
<th>Probe sequence</th>
<th>TC number</th>
<th>Putative function</th>
<th>M2 fluorescence signal</th>
<th>M4 fluorescence signal</th>
<th>Fold change</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCCAGAGTGAAGGACCTGAGCAGCTAGTGACTCCGGATTTTGTC</td>
<td>TC140264</td>
<td>Calcium ion binding</td>
<td>5732.62</td>
<td>100.07</td>
<td>57.28</td>
</tr>
<tr>
<td>GCTACACCCGCTCGATGGTTACGCTGCTTCTTC</td>
<td>TC149834</td>
<td>Fruit weight QTL</td>
<td>5602.90</td>
<td>102.40</td>
<td>54.71</td>
</tr>
<tr>
<td>AGAGGGCATGCGAGGATAGGATGGCTGCTTCAAATT</td>
<td>TC134605</td>
<td>Unclassified</td>
<td>8072.10</td>
<td>237.88</td>
<td>33.93</td>
</tr>
<tr>
<td>AGCTCGAGCGAATCAGTACGCGATTTGCTAAAGGCC</td>
<td>TC149899</td>
<td>Unclassified</td>
<td>2493.08</td>
<td>77.09</td>
<td>32.34</td>
</tr>
<tr>
<td>CCGATGCTACTGCTCGTACTGCTAGTCCAGC</td>
<td>TC151637</td>
<td>Viral protein</td>
<td>6682.69</td>
<td>226.34</td>
<td>29.53</td>
</tr>
<tr>
<td>TACGGAGTTGCTGCCCAGTACAGAGAGAT</td>
<td>U01228.1</td>
<td>Oxygen binding</td>
<td>5344.84</td>
<td>182.46</td>
<td>29.29</td>
</tr>
<tr>
<td>CCTTAGTAGTACGATCCACGACAGTACGATGC</td>
<td>TC136288</td>
<td>Unclassified</td>
<td>2438.20</td>
<td>83.40</td>
<td>29.23</td>
</tr>
<tr>
<td>AGCTGCGCGTCTATAGCGACATCAGACTATCTAC</td>
<td>TC140981</td>
<td>Calcium ion binding</td>
<td>5386.63</td>
<td>190.36</td>
<td>28.30</td>
</tr>
<tr>
<td>TGAGACCATCGTGCTGACCACATAGAACCGGAA</td>
<td>TC141920</td>
<td>Unclassified</td>
<td>3223.38</td>
<td>126.35</td>
<td>25.51</td>
</tr>
<tr>
<td>CCAGAATCGTGAATGCTGATGCTGATGAAAGCA</td>
<td>TC146947</td>
<td>Cytochrome P450</td>
<td>6140.34</td>
<td>241.61</td>
<td>25.41</td>
</tr>
<tr>
<td>TCCTGATGGTTGTTACGCTTACCTGATTTGCTCAGC</td>
<td>TC143944</td>
<td>Putative esterase pir7b</td>
<td>3151.45</td>
<td>129.04</td>
<td>24.42</td>
</tr>
<tr>
<td>TTAAGCCGGATCTTGGCTGCTGATTTGCTGATGGA</td>
<td>TC147886</td>
<td>Serine/threonine protein kinase</td>
<td>2885.95</td>
<td>118.85</td>
<td>24.28</td>
</tr>
<tr>
<td>AGCTTAGCTGTTGCTGAGGATCGACATCAGACGAC</td>
<td>TC143368</td>
<td>Wound induced protein</td>
<td>2705.95</td>
<td>114.38</td>
<td>23.66</td>
</tr>
<tr>
<td>TCATTGCCCAGTCCGATGTTACGCTGCTGATGGA</td>
<td>TC139745</td>
<td>AP2 transcription factor</td>
<td>2780.00</td>
<td>121.21</td>
<td>22.94</td>
</tr>
<tr>
<td>CATCAGGCAATCATTGCTGCAGCTGCTGCTGATGGA</td>
<td>TC147201</td>
<td>Fiber protein</td>
<td>8299.14</td>
<td>366.75</td>
<td>22.63</td>
</tr>
<tr>
<td>CGGAATGACTTCCGATGCTGACTCTCAGGGA</td>
<td>TC136788</td>
<td>Unclassified</td>
<td>2636.99</td>
<td>117.00</td>
<td>22.54</td>
</tr>
<tr>
<td>CAGCTCTGTGGTATCGGTTACCTCAGGCTGCTGCTGTT</td>
<td>TC141008</td>
<td>Zinc finger protein</td>
<td>2047.63</td>
<td>92.84</td>
<td>22.06</td>
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<td>CAGGAGCACGCAGTACGCTGCTGCTGATGGA</td>
<td>TC152041</td>
<td>AP2 transcription factor</td>
<td>1545.31</td>
<td>75.17</td>
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<td>CCTGCTTGAATGCAGGATCGTACGCTGCTGATGGA</td>
<td>TC147315</td>
<td>Oxygen binding</td>
<td>7079.50</td>
<td>354.31</td>
<td>19.98</td>
</tr>
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<td>TCCAGATGATAATGCTGCTGCTGCTGCTGATGGA</td>
<td>TC140302</td>
<td>Unclassified</td>
<td>2920.16</td>
<td>148.46</td>
<td>19.67</td>
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<td>TGAGAATGAGCAGATGCTGCTGAGGATCGTACGAT</td>
<td>TC152456</td>
<td>Transcription factor</td>
<td>1830.35</td>
<td>93.14</td>
<td>19.65</td>
</tr>
<tr>
<td>TCAGGAGCAGAGACCTCAGCAGCTGCTGCTGATGGA</td>
<td>TC136749</td>
<td>Calcium ion binding</td>
<td>1808.35</td>
<td>92.06</td>
<td>19.64</td>
</tr>
<tr>
<td>CCTCTGTGTTACGCTGCTGCTGCTGCTGATGGA</td>
<td>TC133815</td>
<td>EBV nuclear antigen 1</td>
<td>4801.65</td>
<td>245.78</td>
<td>19.54</td>
</tr>
<tr>
<td>GAATCTCTGCTGCTGCTGCTGCTGCTGATGGA</td>
<td>TC150059</td>
<td>AP2 transcription factor</td>
<td>1734.88</td>
<td>89.57</td>
<td>19.37</td>
</tr>
<tr>
<td>AGACTACATTAGCAGCAGCATGAAATGCTAAGCTA</td>
<td>TC139585</td>
<td>Pyrophosphate-dependent 6-phosphofructose-1-kinase</td>
<td>2684.25</td>
<td>143.70</td>
<td>18.68</td>
</tr>
<tr>
<td>ATCGTACGGCTGCTGCTGCTGCTGCTGATGGA</td>
<td>TC147875</td>
<td>CG1343-PA</td>
<td>1751.25</td>
<td>97.22</td>
<td>18.01</td>
</tr>
<tr>
<td>CCTACATGGTGCAGCAGCATGAAATGCTAAGCTA</td>
<td>TC140023</td>
<td>Pyrophosphate-dependent 6-phosphofructose-1-kinase</td>
<td>2680.86</td>
<td>148.91</td>
<td>18.00</td>
</tr>
<tr>
<td>CTGTCGACAGCAATTAGGCAGATGCAGACTGACTGACT</td>
<td>TC140265</td>
<td>Calcium ion binding</td>
<td>1551.44</td>
<td>87.72</td>
<td>17.69</td>
</tr>
<tr>
<td>TCTCTGTGCTGCTGCTGAGCATTGCTGCTGCTGATGGA</td>
<td>TC146486</td>
<td>Extensin</td>
<td>2444.57</td>
<td>143.34</td>
<td>17.05</td>
</tr>
<tr>
<td>TAGAATGAGCAGATGCTGCTGCTGCTGATGGA</td>
<td>TC148895</td>
<td>BHLH transcription factor</td>
<td>2170.63</td>
<td>128.69</td>
<td>16.87</td>
</tr>
<tr>
<td>TACCTGCTGCTGCTGCTGCTGCTGCTGATGGA</td>
<td>TC139118</td>
<td>Putative early nodulin</td>
<td>1613.61</td>
<td>96.57</td>
<td>16.71</td>
</tr>
<tr>
<td>AGACGGTGACGGCAAGCAGCAGCATGAAATGCTAAGCTA</td>
<td>TC134228</td>
<td>Unclassified</td>
<td>7840.21</td>
<td>500.13</td>
<td>15.68</td>
</tr>
<tr>
<td>Probe sequence</td>
<td>TC number</td>
<td>Putative function</td>
<td>M2 fluorescence signal</td>
<td>M4 fluorescence signal</td>
<td>Fold change</td>
</tr>
<tr>
<td>----------------</td>
<td>-----------</td>
<td>----------------------------------------</td>
<td>------------------------</td>
<td>------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>GGAAGCTGGGTTAATATCTATGTTGAGACGGTGCC</td>
<td>TC138639</td>
<td>Jasmonate induced protein</td>
<td>135.00</td>
<td>11690.96</td>
<td>86.60</td>
</tr>
<tr>
<td>AGTCGACTACATCTCATCAAGTCACTCACGATTACGGTACACGTA</td>
<td>TC146793</td>
<td>Metacaspase</td>
<td>117.26</td>
<td>5610.08</td>
<td>47.84</td>
</tr>
<tr>
<td>CACATGCCGAGCAAGGCTTCACTCAAGGAGGCTTACAGTGAGCC</td>
<td>TC146793</td>
<td>Metacaspase</td>
<td>115.54</td>
<td>3871.89</td>
<td>33.51</td>
</tr>
<tr>
<td>CGGATGAGTGTTATGAGAGAGGCTTCACTGACTACCTGAGAATTACGGA</td>
<td>TC146793</td>
<td>Wound induced protein</td>
<td>1424.28</td>
<td>137.38</td>
<td>10.37</td>
</tr>
<tr>
<td>CATCCGACGATCAGGATCGTTAGGAGGCTTACAGTGAGCC</td>
<td>TC146793</td>
<td>Gamma-thionin</td>
<td>4753.73</td>
<td>462.53</td>
<td>10.28</td>
</tr>
<tr>
<td>TAGTTGAGGATCACTGTTGATTGATGAGGCTTACAGGAGAATGAGAACG</td>
<td>TC146793</td>
<td>Lipid-transfer protein</td>
<td>1511.10</td>
<td>147.46</td>
<td>10.25</td>
</tr>
<tr>
<td>BQ663776</td>
<td>TC146793</td>
<td>Probable esterase pir7a</td>
<td>1255.43</td>
<td>123.07</td>
<td>10.20</td>
</tr>
<tr>
<td>ACCTAATCCTCTACATTATGCTGTTGAGGCTTACAGGAGAATGAGAACG</td>
<td>TC146793</td>
<td>Hypoxia-responsive family protein-like</td>
<td>1070.75</td>
<td>95.74</td>
<td>11.18</td>
</tr>
</tbody>
</table>

Table 4.5 Microarray probes with signal up-regulated more than 10 fold from 12 to 48 hours post-steeping.
CCTCCATGATCGATGGCTTAGTGCTTTAACGATCTGCC

TC147022 Unclassified 354.86 9768.19 27.53

TGTAAGGATTAAAGGTGGCCGCCTGCTCAGTACCAAC

AF181457.1a Dehydrin 179.91 4722.58 26.25

TCCTGCCGAGGCTTTACACATGCACTCTCACTCC

TC150722 Proline-rich APG-like protein 129.37 3324.17 25.69

GGAACGAGCAGCAGGAAAACATGGATACGTAC

BQ763715 Unclassified 87.97 2236.94 25.43

AGTACGATCGTACGTTACACGAACTATGATGTC

BG947407 Metallothionein 106.46 2686.88 25.24

ATGAAAGTACCAGTGGCCCGCTGCTACCTCTGC

TC133121 Dehydrin 216.98 5336.53 24.59

CATGATCGATGTTGCTGCTCTACGAAATGAAAC

TC140393 Thaumatin-like protein 142.89 3319.96 23.23

TGAAGCAGGAGAGGAGCTGTTAACGATACCTAT

TC132146 Nucleolar protein family A 104.70 2184.80 20.87

GCCATGTCGTTACCTACGAGCAATCGATCTGTT

TC145699 Beta-1,3-glucanase 89.71 1795.47 20.01

ATGCACCCGATGATCGTACGTTACACGAAATG

X74940.1a Pathogenesis-related protein 293.19 5831.04 19.89

TGTCGTGTTACCTATGCTACGAAATGAAAC

BU968132 Unclassified 82.56 1611.67 19.52

AGATCGTACGTTATCTCGGACTGACCAAC

AJ250662.1a Putative protein 126.90 2462.45 19.40

ACCTGACGTTACCTATGCTACGAAATGAAAC

X98326.1a Dehydrin 474.16 9025.92 19.04

CACTGACGTTACCTATGCTACGAAATGAAAC

TC147142 Bundle sheath protein 155.66 2960.13 19.02

TGCACTGACGTTACCTATGCTACGAAATGAAAC

TC147020 Glycerol-rich protein 105.41 1952.73 18.52

GATATCCGTTACCGTACGAAATGAAAC

BM817394 Unclassified 236.54 4309.85 18.22

CACTGACGTTACCTATGCTACGAAATGAAAC

TC130737 Unclassified 174.58 3025.99 17.33

TTGGTCGTTACCGTACGAAATGAAAC

U0963.1a Germin 86.50 1424.72 16.47

CACTGACGTTACCTATGCTACGAAATGAAAC

M19048.1a Leaf specific thionin 85.66 1378.30 16.09

CACTGACGTTACCTATGCTACGAAATGAAAC

TC139061 Glycerol oxide 119.42 1705.17 14.28

CACTGACGTTACCTATGCTACGAAATGAAAC

TC136137 Serine/threonine-protein kinase 106.03 1475.20 13.91

CACTGACGTTACCTATGCTACGAAATGAAAC

AF181452.1a Dehydrin 532.59 7305.82 13.72

CACTGACGTTACCTATGCTACGAAATGAAAC

TC139406 Glutathione transferase 247.46 3329.47 13.45

CACTGACGTTACCTATGCTACGAAATGAAAC

TC134034 Arabinoxylol arabinofuranosylhydrolase isoenzyme 127.26 1705.03 13.40

CACTGACGTTACCTATGCTACGAAATGAAAC

TC141237 O-methyltransferase 128.12 1691.89 13.21

CACTGACGTTACCTATGCTACGAAATGAAAC

TC139613 Subtilisin-chymotrypsin inhibitor 397.41 5075.95 12.77

CACTGACGTTACCTATGCTACGAAATGAAAC

TC146251 Lipid transfer protein 470.56 5967.69 12.68

CACTGACGTTACCTATGCTACGAAATGAAAC

BR966323 pZEA 288.20 3624.92 12.58

CACTGACGTTACCTATGCTACGAAATGAAAC

AJ7626.1a Chitinase 167.80 2103.03 12.53

CACTGACGTTACCTATGCTACGAAATGAAAC

AB331641.1a HvMATE gene for aluminum activated citrate transporter 289.70 3603.44 12.44

CACTGACGTTACCTATGCTACGAAATGAAAC

TC139245 Dehydroascorbate reductase 202.55 2482.96 12.26

CACTGACGTTACCTATGCTACGAAATGAAAC

X95257.1a Xylose isomerase 140.01 1713.99 12.24

CACTGACGTTACCTATGCTACGAAATGAAAC

AY220734.1a Pathogenesis-related protein 287.27 3471.74 12.09

CACTGACGTTACCTATGCTACGAAATGAAAC

TC146761 Phosphoethanolamine methyltransferase 475.75 5532.68 11.63

CACTGACGTTACCTATGCTACGAAATGAAAC

TC132846 Lipid transfer protein 106.06 1223.33 11.53

CACTGACGTTACCTATGCTACGAAATGAAAC

TC139613 Subtilisin-chymotrypsin inhibitor 532.27 5906.97 11.10

CACTGACGTTACCTATGCTACGAAATGAAAC

BF259687 Unclassified 165.99 1834.82 11.05

CACTGACGTTACCTATGCTACGAAATGAAAC

TC147171 Aldose 1-epimerase 133.65 1431.91 10.71

CACTGACGTTACCTATGCTACGAAATGAAAC

TC149510 Cytochrome P450 147.24 1536.69 10.44

CACTGACGTTACCTATGCTACGAAATGAAAC

TC140027 Aldose 1-epimerase 226.80 2360.45 10.41

CACTGACGTTACCTATGCTACGAAATGAAAC

TC146794 Metacaspase 473.27 4818.68 10.18

CACTGACGTTACCTATGCTACGAAATGAAAC

CB866425 Metallothionein 107.54 1080.94 10.05

a NCBI accession number.
4.4.3 Microarray results compared to LongSAGE and real-time PCR results.

The LongSAGE study identified several genes as candidates of interest to the underlying physiology of malting barley (White et al., 2006). Several of these genes were analysed by real-time RT-PCR which validated the expression patterns identified by LongSAGE. These genes were represented by one or more different probes on the malting barley CombiMatrix 12K Custom array. For those genes represented by more than one probe the signal intensity from each of the samples was averaged. Fold changes observed between 12 and 48 hours post-steeping for the genes of interest in the microarray experiment, reinforce the fold changes observed in the LongSAGE and real-time RT-PCR experiments (Table 4.6). The cell wall hydrolysing enzyme (1-3,1-4)-β-D-glucanase is significantly up-regulated between 12 and 48 hours post-steeping and is observed to increase approximately seven fold by microarray, almost thirteen fold by LongSAGE and over eight fold by real-time RT-PCR. Alpha-amylase, a key enzyme responsible for degrading the starchy endosperm of the germinating barley grain, shows significant increases in gene expression at 48 hours post-steeping. It represents the highest fold change observed in the real-time RT-PCR experiment with an increase of almost 28 fold, and at nearly 50 fold shows one of the highest increases in expression in the LongSAGE experiment. However, the microarray results show just a three and a half fold increase in α-amylase expression between 12 and 48 hours post-steeping. The carbohydrate metabolising enzyme β-1,3-glucanase, the protein hydrolysing enzyme Cysteine Proteinase EPB, and the key regulator of the glyoxylate cycle, Isocitrate lyase, produced microarray probe signal intensity fold changes consistent with those obtained for the same genes using LongSAGE and real-time RT-PCR. These genes were observed to show
significant up-regulation of expression at 48 hours post-steeping in comparison to 12 hours post-steeping. Pyruvate kinase, which controls the rate of glycolysis, and Xyloglucan endotransglycosylase, an enzyme that cleaves and regulates xyloglucan polymers, show a decrease in expression between 12 and 48 hours post-steeping by approximately three to six fold as analysed by microarray, real-time RT-PCR and LongSAGE.

Table 4.6 Malting barley microarray signal intensities of genes of interest and comparison to gene expression values of the same genes of interest using LongSAGE and Real-time RT-PCR.

<table>
<thead>
<tr>
<th>Gene of interest</th>
<th>Microarray signal intensity</th>
<th>LongSAGE expression value</th>
<th>Real-time RT-PCR expression value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>12 hours post-steeping</td>
<td>48 hours post-steeping</td>
<td>12 hours post-steeping</td>
</tr>
<tr>
<td>(1-3,1-4)-beta-D-glucanase</td>
<td>1584.91</td>
<td>11056.67</td>
<td>0.065</td>
</tr>
<tr>
<td>Fold change</td>
<td>6.98</td>
<td>12.76</td>
<td>8.16</td>
</tr>
<tr>
<td>Beta-1,3-glucanase</td>
<td>1912.23</td>
<td>11798.74</td>
<td>0.010</td>
</tr>
<tr>
<td>Fold change</td>
<td>6.17</td>
<td>3.95</td>
<td>7.31</td>
</tr>
<tr>
<td>Alpha amylase</td>
<td>229.13</td>
<td>798.91</td>
<td>0.010</td>
</tr>
<tr>
<td>Fold change</td>
<td>3.49</td>
<td>49.38</td>
<td>27.66</td>
</tr>
<tr>
<td>Cysteine proteinase EPB</td>
<td>3340.83</td>
<td>9403.65</td>
<td>0.070</td>
</tr>
<tr>
<td>Fold change</td>
<td>2.81</td>
<td>7.07</td>
<td>5.85</td>
</tr>
<tr>
<td>Isocitrate lyase</td>
<td>3012.36</td>
<td>7797.07</td>
<td>0.005</td>
</tr>
<tr>
<td>Fold change</td>
<td>2.59</td>
<td>75.00</td>
<td>6.89</td>
</tr>
<tr>
<td>Pyruvate kinase</td>
<td>9500.15</td>
<td>3289.57</td>
<td>0.130</td>
</tr>
<tr>
<td>Fold change</td>
<td>2.70*</td>
<td>13.00*</td>
<td>2.89*</td>
</tr>
<tr>
<td>Xyloglucan endotransglycosylase</td>
<td>1962.88</td>
<td>364.67</td>
<td>0.035</td>
</tr>
<tr>
<td>Fold change</td>
<td>4.53*</td>
<td>3.54*</td>
<td>5.38*</td>
</tr>
</tbody>
</table>

*NB. In these fold change ratios the signal intensity is higher at 12 hours post-steeping.

The majority of the microarray signal intensities produce relatively lower fold change ratios between 12 and 48 hours post-steeping than the LongSAGE and real-time RT-PCR technologies. This may be because of a number of contributing factors.
Microarray signal intensity is subject to background noise caused by cross hybridisation and the misreading of high signals adjacent to low signals. This generally leads to overestimation of the signal associated with very lowly expressing transcripts. This may be the case with α-amylase where a signal intensity of only 229 is present at 12 hours post-steeping which is within the range of background noise associated with the array hybridisation. Another point to note is that the fold change ratio of α-amylase as analysed by LongSAGE is considerably higher. This may be a result of more than one gene family member being identified by the one LongSAGE tag, ultimately overestimating the gene expression at 48 hours post-steeping. The oligonucleotide probe on the CombiMatrix array is approximately fifty percent longer than the SAGE tag, allowing for more specific hybridisation and differentiation of gene family members.

The microarray results for α-amylase are different to the LongSAGE and real-time RT-PCR results not only in terms of the fold change value but in the level of total signal intensity compared to the other types of gene transcripts analysed. In the LongSAGE experiment α-amylase has one of the highest total tag counts at 48 hours post-steeping and the relative expression value of α-amylase identified at the same time point by real-time RT-PCR is also quite high compared to the other gene transcripts studied. However the signal intensity of α-amylase detected in the microarray experiment is quite low at 799 compared to the other gene transcripts which reach over 11 000 units. This may be explained in part by the presence of other gene family members in the case of the LongSAGE experiment but not in the real-time RT-PCR experiment where the PCR primers are specific for one gene family member only.
The binding specificities of particular probe sequences have an effect on signal intensity. For example, if the probe sequence representing α-amylase did not have strong binding efficiency it may result in lower signal intensity. Probe sequence specificity may contribute to a general underestimation of microarray signal intensity due to the cross hybridisation of similar probe sequences especially if the binding efficiency of those sequences was not high. This may be the case for Cysteine Proteinase EPB which has a relatively low fold change value in the microarray experiment compared to the other methodologies, apparently due to a comparatively higher signal intensity detected at 12 hours post-steeping than would be expected (Table 4.5). This could have resulted from the cross hybridisation of other proteinase transcripts to the probe.

It is possible that the fold change values calculated from the microarray results for the genes of interest are underestimated due to the saturation of signal intensity at high levels. Some probes on the array are detected at signal intensity levels above 20 000 units (Table 4.2) compared to a maximum of 12 000 units amongst the genes of interest so this seems unlikely (Table 4.5). Conversely the fold change values detected by LongSAGE tag counts may be overestimated if the SAGE library has not been sequenced to a reasonable depth. For example, the fold change ratios for α-amylase and pyruvate kinase are considerably higher when analysed by LongSAGE than by microarray or real-time RT-PCR (Table 4.5). This is because a false value had to be used to estimate the relative expression of transcripts not detected by even one tag count. Those tags with a count of 0 were given a false count of 0.01 to enable comparisons with other libraries in which the tag was present (White et al., 2006). It
cannot be determined whether increasing the number of SAGE tags sequenced would reveal a count for those particular transcripts, and so the fold change is overestimated because the lower level of relative expression is unknown or undetected by the depth of sequencing undertaken in the experiment. The depth of sequencing chosen for SAGE experiments is a critical factor in the effectiveness of the technology for quantitative analysis of gene expression. A comparison of Affymetrix chips with SAGE libraries predicted the sensitivity of an Affymetrix chip to be approximately comparable to a SAGE library of 120 000 total tags (Lu et al., 2004).

SAGE is a relatively expensive method of gene expression analysis. To compare different samples; for example, different developmental time points, tissues or treated versus untreated samples; an entire SAGE library must be created for each sample of interest. The process of making a SAGE library takes approximately two days after which the clones have to be sequenced. Advancements in the SAGE protocol, to analyse longer tag concatemers per sequencing run (Crawford et al., 2005), and increased throughput and efficiency of modern sequencing platforms has reduced both the cost of sequencing and the time taken to generate large amounts of tag sequence data. The effectiveness of SAGE technology has been shown to rely on the depth of tag sequencing undertaken and in a recent extensive study it was found that approximately 650 000 tags were required to sample a transcriptome of approximately 56 000 (Velculescu et al., 1999).

SAGE can be limited in its ability to detect lowly expressed transcripts. The method was developed with the intention of reducing the amplification bias caused by transcripts of significantly high copy number that is observed in some other methods.
By reducing this bias SAGE claims to present a balanced global picture of gene expression (Margulies et al., 2001; Velculescu et al., 1995). However if the SAGE library is not sequenced to a significant depth, transcripts with low levels of expression may not be detected.

Other factors inherent to the SAGE technology which could result in under or overestimation of gene expression include, antisense tags, gene family members and alternative splice variants. LongSAGE tags are 21bp long; theoretically long enough to differentiate approximately $4 \times 10^{21}$ different transcripts. However very similar or exact sequences are likely to arise particularly from gene family members. If more than one gene family member shares the same tag sequence, their counts would be combined resulting in an overestimate of the frequency for that gene transcript. This may be the case for $\alpha$-amylase as mentioned above.

SAGE results can also be underestimated when a transcript is represented by multiple tags. Alternative splicing, incomplete digestion with the anchoring enzyme, alternative polyadenylation and antisense tags are reported as sources of multiple tags (Pauws et al., 2001; Gibbings et al., 2003; Ibrahim et al., 2005; Wahl et al., 2005; White et al., 2006). These events are a common problem in SAGE studies. White et al. (2006) report an occurrence of alternative polyadenylation in a Gamma-thionin transcript. When the tag count of each of the two tags was combined the total relative abundance of the transcript was significantly higher than either of the individual tags indicated. Unless these events are identified and the count of the two or more corresponding tags added together, the transcript expression level of certain SAGE tags can potentially be significantly underestimated.
Despite its downfalls, SAGE is still a useful tool for gene expression analysis. It has the ability to capture expression data for all transcripts in a particular tissue, at a particular time or under specific conditions. This includes transcripts about which no sequence data has previously been obtained. This is a unique advantage of this high throughput gene expression technology.

CombiMatrix ElectraSense™ Microarray technology also has its pros and cons. As seen above, microarray technology provides a more accurate picture of gene expression over a wider range of expression values than SAGE. The CombiMatrix ElectraSense™ CustomArray is also a cost effective method. Once the initial chip has been produced, it is relatively inexpensive to produce replicate chips. The chips can also be stripped and reused up to three times, for a total of four hybridisations per chip. However gene expression values can be subject to cross hybridisation, probe specificity and saturation effects. If these things were corrected, particularly for particular genes of interest, for example for genes directly related to malting quality, then the CombiMatrix ElectraSense™ CustomArray would make an excellent tool for analysis of the transcriptome of barley. A chip such as this could be used to check and QC samples of barley for qualities such as shelf life and malt performance. Such assays could be performed with relatively small amounts of grains, in a short amount of time and at a reasonable cost. This assay would be a valuable tool to provide the malt barley industry with information about the impact of grain storage on shelf life and malt quality, the malting performance of different barley varieties, growing environments and much more.
4.5 Acknowledgements

The authors would like to thank Dana Pascovici and Mervyn Thomas from Emphron Informatics for advice and assistance on background correction and normalisation of microarray data.

4.6 References


Chapter Five

General Conclusions

5.1 Conclusions and summary

The genetic processes involved in barley seed germination, or malting, are poorly understood. Physiological and biochemical changes are occurring within the grain at all times and this study aimed to determine the genetic processes that occur during malting that are associated with these changes.

This study used two large scale gene expression analysis technologies to examine gene expression changes in malting barley over time. As technology has developed over the past ten years, LongSAGE and microarray have become two of the most commonly used methods for generating large amounts of gene expression data. In this study 155 000 LongSAGE tags, representing 41 909 unique tag sequences, were generated from eight tag libraries of malting barley. Analysis of the most abundant LongSAGE tags highlighted the functional groups of dominant mRNA transcripts which include transcripts coding for stress response and cell defence, ribosomal proteins, storage proteins, cell structure and plant growth, and metabolism related transcripts. Annotation of the LongSAGE tags revealed that multiple tags matched to the same transcript. The presence of alternative transcripts has been explored in several SAGE studies and appears to be a result of several variables including alternative splicing, incomplete digestion, alternative polyadenylation and anti-sense transcripts (Pauws et al., 2001; Gibbings et al., 2003; Robinson et al., 2004; Ibrahim et al., 2005; Wahl et al., 2005). SAGE libraries can therefore also be used to examine
aspects of mRNA processing such as the presence of anti-sense transcripts and enzyme sites that enable alternative tags to be created.

Statistical analysis confidently identified 57 LongSAGE tags with significant changes in abundance across the eight malting time points. Three general patterns of expression were observed; tags with higher expression during the first 12 to 24 hours post-steeping, tags showing up-regulation from 24 hours post-steeping and tags with specific up-regulation at 24 hours post-steeping and 120 hours post-steeping. A sample of LongSAGE tags showing significant changes in gene expression across time were chosen for validation by real-time RT-PCR. Metabolism related transcripts, particularly carbohydrate and protein modifying transcripts, were selected for real-time RT-PCR analysis because of a more direct relationship to seed germination and malting.

Real-time RT-PCR experiments were performed on samples obtained at four time points corresponding to the LongSAGE study; pre-steeping, 12 hours post-steeping, 48 hours post-steeping and 96 hours post-steeping. The mRNA abundance levels observed by real-time RT-PCR were compared with the abundance levels from the LongSAGE experiment with similar patterns observed for the two methods. The RT-PCR results provide support for the precision of detection achievable with SAGE and provide greater confidence in the levels of expression detected in the study. Three transcripts, α-amylase, (1-3,1-4)-β-D-glucanase and cysteine proteinase EP-B, produced very similar patterns of expression. The transcripts for these enzymes were significantly up-regulated at 48 hours post-steeping and remained significantly up-regulated throughout malting. The co-ordinated expression of these enzymes provides
substantial support for the view that these enzymes have key roles as the major contributing enzymes in the depolymerisation of starch, the breakdown of cell walls and degradation of protein in the germinating grain.

Chapter four describes a microarray study of malting barley samples. A CombiMatrix 12K ElectraSense™ Custom array was designed on the basis of the unique LongSAGE tags from 24 hours post-steeping. Chips were hybridised with cRNA from 12 and 48 hours post-steeping. The reproducibility of results was examined and very high correlation co-efficients were seen for replicate scans and between chip comparisons. Re-use of the chips of up to three times did not appear to have adverse effects on the signal intensity results and this contributes to the economic advantages of the CombiMatrix ElectraSense™ array system.

There is always the risk in any study for experimental error. Several elements were incorporated into the experimental design to minimize error in this study. The same biological samples were used for each of the three experiments, the LongSAGE libraries, the real-time RT-PCR experiments and the ElectraSense™ microarray hybridisations. Multiple cDNA synthesis reactions and duplicate PCRs were performed in the real-time RT-PCR experiment. A reference gene was amplified in each experiment to normalise all of the real-time RT-PCR experiments and genes of interest were compared to multiple internal control genes. The microarray experiment included several chip swaps, stripping and re-hybridisations to test the reproducibility of results with very high correlation co-efficients. Spike-in controls were also included to control for bias across an individual chip.
Gene expression levels for eight genes of interest were compared from the three different technologies. In most cases fold change values between 12 and 48 hours post-steeping were not remarkably dissimilar, however differences were observed. Several variables contribute to these differences, some which can be readily controlled and others which cannot. Transcript abundances can be underestimated if a transcript is represented by more than one SAGE tag, however if these occurrences are identified the tag count of the individual tags could be added together to generate the frequency by which the transcript is actually represented in the library. The opposite problem can occur with gene family members, whereby more than one transcript can be represented by a unique tag. However with LongSAGE this is unlikely since a tag of 21 bp is likely to provide a unique tag signature for each transcript. The depth to which a library is sequenced will determine how many different transcripts are identified and genes with low expression may not be detected. Despite these limitations SAGE is a very useful tool for gene expression analysis. It provides the opportunity to capture expression data for all transcripts in a particular tissue, at a particular time or under particular conditions, including transcripts about which there is no known sequence data.

The CombiMatrix ElectraSense™ microarray is a very cost effective and accurate tool for large scale gene expression studies. With the ability to re-use chips and the sensitive electrodetection system it has several advantages over other oligonucleotide chip systems. This experiment examined a large number of transcripts and compared the expression at 12 and 48 hours post-steeping. The CombiMatrix ElectraSense™ array has the potential to be used in further studies to compare many different time
points, varieties and treatments of barley grains to generate valuable information for the malting and brewing communities.

This study has highlighted many areas of interest for potential further work. The data collected in this work could be used in further experiments to determine the distinction between transcript levels in the embryo versus the endosperm. The level of various proteins could be examined using assays or antibodies to examine if protein level correlates with transcript level. siRNA or microRNA experiments could be used to determine the level of effect of post-transcriptional silencing. There are also several new technologies available for whole transcriptome analysis including Next Generation (NG) sequencing or ‘deep sequencing’ techniques which include 454 sequencing, Illumina sequencing and the SOLiD system (Wall et al., 2009). These technologies allow the analysis of the whole transcriptome or in many cases the whole genome of an organism in one run. The potential application for this technology is enormous. The same biological samples were used for these experiments which enabled the comparison of three technology platforms however it would also be useful to perform some of these experiments with different biological samples.

This study has provided valuable insight into the types of genes expressed during malting and the temporal patterns in which they are expressed. This information provides a useful understanding of the genetic control of the biological and physiological changes which occur in the embryo and endosperm of the germinating barley grain. Other important outcomes of this study include an extensive LongSAGE database of quantitative transcript data which covers the malting barley transcriptome of a common Australian malting variety. This resource has been validated by real-
time RT-PCR and microarray analyses. The LongSAGE database and the CombiMatrix ElectraSense™ microarray chip are valuable tools for further research of malting barley qualities.

5.2 References


generation sequencing technologies for transcriptome characterization. *BMC Genom.*, 10:347
Appendix One

Acceptance letter

“Differential gene expression analysis in a barley seed germination time course and confirmation with relative real-time RT-PCR” in the journal, Plant Science

Date: Fri, 15 Aug 2008 09:49:24 +0100
From: german.spangenberg@dpi.vic.gov.au
Subject: Your Submission
To: robert.henry@scu.edu.au
X-IronPort-Anti-Spam-Filtered: true
X-IronPort-Anti-Spam-Result: AqMBAKvpEIbJbACCAcQEBCAUIBxGaRIZZgVY
X-IronPort-AV: E=Sophos;i="4.32,214,1217772000"; d="scan'208";a="12893230"
X-SMTP-Vilter-Version: 1.2.4
X-SMTP-Vilter-Virus-Backend: clamd
X-SMTP-Vilter-Status: clean
X-SMTP-Vilter-clamd-Virus-Status: clean
Original-recipient: rfc822;robert.henry@muon.lis.scu.edu.au
X-OriginalArrivalTime: 15 Aug 2008 08:49:24.0992 (UTC) FILETIME=[D1A52000:01C8FEB3]

Ms. Ref. No.: PSL-D-07-00648R2
Title: Differential LongSAGE tag abundance analysis in a barley seed germination time course and validation with relative real-time RT-PCR
Plant Science

Dear Prof. Robert Henry,

I am pleased to confirm that your paper "Differential LongSAGE tag abundance analysis in a barley seed germination time course and validation with relative real-time RT-PCR" has been accepted for publication in Plant Science.

Thank you for submitting your work to this journal.

With kind regards,

German Carlos Spangenberg, Ph.D DSc
Receiving Editor
Plant Science
Appendix Two

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Wiley-Blackwell publishing - Plant Biotechnology Journal

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