

## Differentially expressed genes during malting and correlation with malting quality phenotypes in barley (*Hordeum vulgare* L.)

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**Abstract** Breeding for malting quality is an important goal of malting barley breeding programs. Malting quality is a complex phenotype that combines a large number of interrelated components, each of which shows complex inheritance. Currently, only a few genes involved in determining malting quality have been characterized. We combined transcript profiling with phenotypic correlations to identify candidate genes for malting quality. The Barley1 GeneChip<sup>®</sup> array containing 22,792 probe sets was used to conduct transcript profiling of genes expressed in several different stages of malting of four malting cultivars. Genes that were differentially expressed in comparisons between different malting stages relative to ungerminated seed, as well as in comparisons between malting cultivars in the

same malting stage were identified. Correlation analysis of 723 differentially expressed genes with malting quality phenotypes showed that 11–102 of these genes correlated with six malting quality phenotypes. Genes involved in carbohydrate metabolism were among the positively correlated genes. Genes for protein and lipid metabolism, cell wall organization and biogenesis, and genes involved in stress and defense response also correlated with malting quality phenotypes. Expressed sequence tags (ESTs) were generated from a ‘malting-gene enriched’ cDNA library made by suppression subtractive hybridization between malted and ungerminated seeds of ‘Morex’. Eleven percent of the ESTs had no significant homology with sequences in the databases, suggesting that there may be other malting-related genes

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not represented in the barley gene chip array. The results provide candidate genes for malting quality phenotypes that need to be functionally validated.

## Introduction

Barley (*Hordeum vulgare* L.) is the fifth most important cereal crop in terms of total world production (FAOSTAT 2006; <http://faostat.fao.org/site/370/default.aspx>), and it is the major grain used for malting and brewing. Malting is seed germination under strictly controlled environmental conditions. During malting, the stored biopolymers in barley grains are converted into low molecular weight molecules that can be utilized by yeast during brewing (Jones 2005a). The malting process involves three stages: steeping, germination, and kilning (Briggs 1998). During steeping, the barley seed is soaked in water to reach predefined moisture content. During the germination phase, hydrolytic enzymes are synthesized by the aleurone cells (Bamforth and Barclay 1993) and scutellum (Macfadden et al. 1988; Ranki 1990). These hydrolytic enzymes are secreted into the starchy endosperm where they convert proteins and carbohydrates into partially degraded biopolymers (Fincher 1989). In kilning, the malted seed is dried by forcing air at 40–60°C and then the temperature is gradually increased up to 85–95°C (Briggs 1998). Other than the low molecular weight biopolymers, the malt also contains heat-stable hydrolytic enzymes required for further breakdown of carbohydrates in the subsequent mashing process (Potokina et al. 2004).

Understanding the genetic basis of malting quality is important in malt barley breeding. Malting quality is a complex phenotype that combines a large number of inter-related components, each of which shows complex inheritance (Hayes and Jones 2000). Genetic mapping of malting quality phenotypes resulted in more than 150 QTLs associated with 19 traits used to measure malting quality (Gao et al. 2004; Zale et al. 2000). Currently, only a few genes involved in determining malting quality have been characterized. These include genes coding for the starch degrading enzymes  $\alpha$ -amylase (1,4- $\alpha$ -D-glucanohydrolase, EC 3.2.1.1),  $\beta$ -amylase (1,4- $\alpha$ -D-glucan malto-hydrolase, EC 3.2.1.2),  $\alpha$ -glucosidase, and limit dextrinase (LD, EC 3.2.1.41) (for review, see Fincher 1989). However, the large number of QTLs associated with malting quality traits suggest that many other genes play important roles in malting.

Analysis of gene expression profiles using cDNA-based microarrays has proven useful for deciphering biochemical pathways involved in plant responses to stress and pathogens (Caillaud et al. 2007; Caldo et al. 2004; Venu et al. 2007). Differentially expressed genes during germination in

barley were also studied using cDNA arrays (Potokina et al. 2002; Watson and Henry 2005). Using serial analysis of gene expression (SAGE) to conduct transcript profiling during malting in barley, the most abundant transcripts identified by (White et al. 2006) overlapped with most genes identified in cDNA array-based studies (Potokina et al. 2002; Watson and Henry 2005).

The use of cDNA-based arrays for identifying genes that are involved in malting was previously demonstrated (Potokina et al. 2004, 2006). Ten malting barley genotypes showing variation in six malting quality characteristics were compared for gene expression during malting using a cDNA array containing 1400 barley ESTs. The differentially expressed genes were then correlated with malting quality phenotypes (Potokina et al. 2004) resulting in the identification of 17–30 candidate genes for each of the malting parameters. A subsequent study further analyzed a candidate gene, serine carboxypeptidase I (*Cxp1*), by mapping the relative level of *Cxp1* expression in a population made from a cross between the malting barley cultivar Morex and a feed barley cultivar, Steptoe (Potokina et al. 2006). An expression QTL mapped to the same region on chromosome 3H containing a QTL for a malting quality phenotype, diastatic power. A single nucleotide polymorphism (SNP) for *Cxp1* was further correlated with phenotypic variation in diastatic power.

This study was conducted to identify genes that are involved in determining malting quality phenotypes. A barley microarray containing more than 22,792 probe sets representing approximately 21,439 barley genes has been developed (Close et al. 2004). We used the barley gene chip array to conduct transcript profiling of genes that are differentially expressed during malting in barley and then correlated differentially expressed genes with malting quality phenotypes. Furthermore, we generated ESTs from a suppression-subtractive hybridization cDNA library made from malted seeds of the malting cultivar ‘Morex’.

## Materials and methods

Plant material, experimental design, and micromalting conditions

Four North American spring type malting barley cultivars that have different malting quality phenotypes and represent 2-row and 6-row varieties were used. The 2-row varieties were ‘Harrington’ (Klages/7211) and ‘Merit’ (Manley/2B80-350), and the 6-row varieties were ‘Legacy’ (6B86-3517/ Excel), and ‘Morex’ (Cree/Bonanza). All four lines were grown in Idaho Falls during the summer of 2003 in standard plots (1.5 × 4 m) consisting of three replicates in randomized complete block trials optimized for nearest

neighbor analysis. Harrington and Merit were grown in one block with several other two-row spring barleys while Morex and Legacy were grown in an adjacent block with several other six-row barleys. The plots were harvested with a plot combine and a sub-sample of 300 g of grain was retained from each plot. Sub-samples of each replicate of each variety were blended together in equal proportions to form a representative sample of about 900 g. The blended grain samples were subsequently cleaned over an A/S Rationel Kornservice model SN4 cleaner to remove thin kernels (by passing through a 6/64th slotted screen). The cleaned blended samples were used for the replicate experiments in this study. By blending samples from replicate plots, field effects should therefore be minimal. Fifty to 100 g seeds of each variety were placed inside a steel container and arranged at random inside the cabinet of a Joe White micromalter located at Busch Agricultural Resources, LLC, Fort Collins, CO. The micromalting conditions were as follows: steeping at 12°C for 37 h, germination at 12°C for 5 h, 17°C for 65 h, 18°C for 23 h, and kilning stages consisting of 55°C for 10 h, 60°C for 4 h, 68°C for 3 h, 80°C for 2 h and 90°C for 3 h. Twenty gram samples of micromalted seeds were collected at four stages for Morex and two stages for Legacy, Harrington, and Merit (Table 1). The four malting stages for Morex were: steeping, 24 h germination (day 1), 93 h germination (day 4), and finished malt after kilning was completed (Fig. 1). Two 20 g samples of un-germinated dry seeds of Morex from the same batch of seeds were used as control. For the three other cultivars, collection was done at day 1 and day 4. Tissues were immediately frozen in liquid nitrogen for RNA extraction.

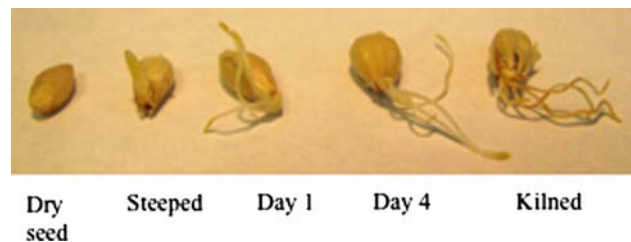
#### RNA extraction, microarray experiments and data analysis

Approximately 100 mg of frozen, ground tissue from each sample was used for RNA extraction. Total RNA was extracted using Qiagen RNeasy Plant Mini Kit with RNase Free/DNase (Qiagen, Valencia, CA) following the manufacturer's instructions. Integrity and quantity of the RNA was tested using Bio-Rad Experion RNA StdSen Chips (Bio-Rad, Hercules, CA). The RNA samples were sent to the University of California at Irvine or the Virginia Bioinformatics Institute Core facility, where additional quality control was performed. These facilities then performed RNA labeling and hybridization to the Affymetrix 22 K Barley1 GeneChip according to the manufacturer's protocols (<http://www.affymetrix.com>).

Following hybridization to the gene chip arrays, different quality control checks were performed including inspection of hybridized images, boxplots and histograms of  $\log_2(\text{PM})$  values, and examination of hybridization and PolyA controls. GeneChips with poor quality were discarded from the analysis. Twenty-two arrays were analyzed with two

**Table 1** Malting stages and cultivars used in microarray experiments with corresponding GEO accession numbers for the microarray data

Malting stage	Replication no.	GEO accession no.
A. Malting stage time course experiment in Morex		
Steeping	1	GSM282034
	2	GSM282036
Day 1	1	GSM282032
	2	GSM282037
Day 4	1	GSM282035
	2	GSM282039
Kilned	1	GSM282038
	2	GSM282040
Dry seed	1	GSM282025
	2	GSM282041
Cultivar	Malting stage	GEO accession no.
B. Cultivar comparisons		
Morex (6-row)	Day 1 (rep 1)	GSM282032
	(rep 2)	GSM282037
	Day 4 (rep 1)	GSM282035
	(rep 2)	GSM282039
Legacy (6-row)	Day 1 (rep 1)	GSM282027
	(rep 2)	GSM282026
	Day 4 (rep 1)	GSM282021
	(rep 2)	GSM282020
Merit (2-row)	Day 1 (rep 1)	GSM282028
	(rep 2)	GSM282029
	Day 4 (rep 1)	GSM282024
	(rep 2)	GSM282023
	(rep 3)	GSM282022
	Harrington (2-row)	Day 1 (rep 1)
Day 4 (rep 1)	GSM282030	
	(rep 2)	GSM282033



**Fig. 1** Barley seeds of cultivar 'Morex' at different malting stages analyzed in this study: dry seed, steeped for 37 h, day 1 (germination for 24 h), day 4 (germination for 93 h), and kilning (see text for details of malting conditions)

biological replications per cultivar per time point, with the exception of Merit day 4 with three replications and Harrington day 1, where one of the replicates was discarded (Table 1). All microarray data have been deposited in NCBI's Gene Expression Omnibus (Edgar et al. 2002) and

are accessible through GEO Series accession number GSE11200 (<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE11200>) (Table 1).

Data analysis was carried out using Bioconductor in R (Gentleman et al. 2004). Data preprocessing and summarization were performed using Robust Multichip Average (RMA) (Irizarry et al. 2003). Statistical tests of differential expression were conducted using the moderated *t* test through the limma (Linear Models for Microarrays) package in Bioconductor. The Benjamini-Hochberg multiple testing adjustment was applied in order to control the comparison-wise false discovery rate (Benjamini and Hochberg 1995).

Two sets of analyses were performed (Table 1). First, expression values in four malting stages in Morex cultivar (steeping, day 1, day 4, and kilned) were compared to the ungerminated, dry seed (non-malting control) stage. Second, expression values were compared during day 1 and day 4 among the four cultivars Morex, Legacy, Harrington, and Merit.

For the Morex time course experiment, 5,579 probe sets that were called “Absent” on every Morex array were removed from further analysis. For the cultivars comparison, a comparison of 2-row cultivars versus 6-row cultivars and all pair-wise cultivar comparisons were performed for day 1 and day 4 samples separately. We note that since there was only a single Harrington replicate for day 1 we did not consider any pair-wise cultivar comparisons involving Harrington for day 1. Five thousand two hundred sixty probe sets that were called “Absent” on every array (in the cultivars experiment) were removed from further analysis. For both analyses, genes were considered to be differentially expressed if the probe sets had an absolute value of  $\log_2$  fold change  $|\log_2(\text{FC})| > 1$  and adjusted *P*-values of  $\leq 0.05$ .

The probe sets identified as differentially expressed were annotated using HarvEST:Barley (Version 1.65) assembly

25 (<http://www.harvest-web.org/>). For gene ontology (GO), we used the *Arabidopsis* gene model for each probe set and used the program PLAN (He et al. 2007) to obtain functional categories. GO annotation was obtained for the top significant hit (using an *e*-value cutoff of  $1e-4$ ) for each probe set.

### Malting quality phenotypes

To mitigate bias of environmental effects from a single location, we used data from a relatively large number of field experiments. Nine malting quality traits routinely used for screening breeding materials at Busch Agricultural Resources were initially considered. The data for malting quality phenotypes included phenotypic measurements of individual malting traits from as many as 49 independent field experiments (Table 2). These trials were conducted from 2001 to 2006 over a wide range of geography and different farming practices, such as irrigated versus dry land. For a given trait, the analysis was restricted to those field experiments for which all four cultivars were represented. A two-way analysis of variance (ANOVA) including a random experiment effect was calculated for each of the nine traits (Table 2). Only wort protein did not have a statistically significant cultivar effect (at  $P \leq 0.05$  level) and it was excluded from further analysis.

Broad-sense heritability ( $H^2$ ) estimates for some of these traits have been reported (Foster et al. 1967; Hockett and Nilan 1985; Therrien 2006) and there is a wide range of values obtained in different studies, as well as different sets of environments within a study. Because we used a relatively large set of paired phenotypic data over a range of locations and years as the best unbiased estimate of the true phenotypic response of the four varieties for the respective malting traits the distorting effects of individual locations and years should have been minimized in our analysis.

**Table 2** Summary of analysis of variance of nine malting quality phenotypes with the least squares means for each cultivar by phenotype

Malting quality phenotype (units)	Number of field locations <sup>a</sup>	Harrington	Legacy	Merit	Morex	Std. error <sup>b</sup>	<i>P</i> -value <sup>b</sup>
$\alpha$ -amylase dextrining unit (DU)	34	73.14	71.86	82.65	61.23	2.483	<0.0001
$\beta$ -glucan (ppm)	34	207.36	225.37	171.07	214.63	22.211	0.01
Barley protein (%)	44	12.64	12.49	12.81	13.12	0.294	0.02
Diastatic power (DU)	34	127.78	165.13	150.50	168.65	6.381	<0.0001
Free amino nitrogen (ppm)	9	186.00	179.53	194.02	158.51	5.291	<0.0001
Fine extract (%)	34	79.90	79.03	79.93	78.62	0.416	<0.0001
Malt protein (%)	27	12.02	12.74	12.13	13.05	0.386	0.0003
Soluble/total protein	34	45.93	44.62	45.87	41.37	1.222	<0.0001
Wort protein (%)	34	5.52	5.54	5.53	5.37	0.099	0.16

<sup>a</sup> Indicates the number of locations out of the total 49 where measurements for the phenotype was available for all four cultivars

<sup>b</sup> Standard error and *P*-value based on the *F*-test for cultivar differences

A short description of the nine malting quality phenotypes follows.  $\alpha$ -Amylase is a measure of the units of enzyme activity to break down a known standardized branched dextrin into smaller linear pieces by internal cleavage in a given amount of time at a constant temperature and pH 9. Beta glucan is a measure of the degree of cell wall modification and indicates the enzyme activity level of beta-glucanase, as measured by the ability of a cold water extract of malt to digest a known standardized beta glucan substrate using a Skalar instrument. Barley protein is a measure of the percent protein found in the grain. It is measured indirectly using near-infrared technology and is corrected to a standard dry moisture (13%) basis. Diastatic power refers to the starch-degrading capacity of malt, and represents the complementary actions of  $\alpha$ -amylase and  $\beta$ -amylase on starch to produce sugars and low-molecular weight dextrans. Diastatic power is measured like  $\alpha$ -amylase by incubating malt extract with a starch substrate, but the malt extract used as enzyme source is not heated leaving the more thermolabile  $\beta$ -amylase active. Percent fine extract refers to the concentration of solids when finely ground malt is mixed with water and the stirred mash is subjected to a time/temperature program that facilitates amylolysis, proteolysis and other enzyme reactions. Free amino nitrogen is a measure of the concentration of individual wort amino acids and small peptides which can be utilized by yeast for cell growth and proliferation. Malt protein is a measurement of total nitrogen on kilned finished malt. Soluble/total protein ratio, also known as Kolbach index, represents the ratio of soluble to total nitrogen. Wort protein, also called soluble protein, is a measure of percent soluble protein in the wort after hot water extraction of the malt. It is measured spectrophotometrically as set by a wet chemistry standard curve. Measurements for these malting quality traits are based on standard procedures described by the American Society of Brewing Chemists (ASBC 1992) and are discussed in previous reviews (Bamforth and Barclay 1993; Burger and LaBerge 1985).

#### Correlation between gene expression and malting quality phenotypes

A subset of genes that were differentially expressed in at least one malting stage versus dry seed and in at least one comparison between cultivars in day 1 and day 4 was correlated with the four malting quality phenotypes. The correlation between gene expression (taking day 1 and day 4 separately) and average phenotype (for each of the nine traits averaged of field experiments) was calculated. Correlation was calculated based on the four pairs of observations representing the expression and phenotypic data across the four cultivars. Calculation of correlation between

gene expression and phenotype average, however, ignores experiment and environmental effects which are known to be present in the data. In order to take the variability due to field experiment into account, we performed an additional analysis where the correlation was calculated by gene and by field experiment. Genes that showed statistically significant correlation to average phenotype ( $P \leq 0.01$ ) and strong correlation (absolute value of correlation  $\geq 0.7$ ) for at least half of the individual field experiments for a phenotype were identified.

#### Quantitative reverse transcription-PCR (qRT-PCR)

Primer pairs for each target gene were based on specific barley sequences from the 22 K Barley1 Affymetrix Gene-Chip probe array (Affymetrix, Inc., Santa Clara, CA) and were designed online using Primer3 (Rozen and Skaletsky 1998). Four genes were selected for these experiments:  $\alpha$ -amylase,  $\alpha$ -glucosidase, limit dextrinase, and  $\beta$ -ketoacyl synthase. Primer sequences were as follows:  $\alpha$ -amylase (contig3953) L: 5'-CGGCAATGACTATGCCGAT-3', R: 5'-GCATGTCCCTCATCCTCACT-3';  $\alpha$ -glucosidase (contig7937) L: 5'-AGCGATTGGTGATGGTAAG-3'; R: 5'-AGTCCTCTGCGTTCTGGTA-3'; limit dextrinase (contig11648) L: 5'-CGGTTTCAACACGAGGATCT-3'; R: 5'-ACTAGCAGCTTGGG CACTA-3'; and  $\beta$ -ketoacyl synthase: (contig6642\_at) L: 5'-GAGGACACAACTCG GTGGTT-3', R: 5'-GCAATGGATCTTGGATCCTC-3'. qRT-PCR was performed using the qPCR MasterMix Plus for SYBR(R) Green I with fluorescein (Eurogentec, Inc., San Diego, CA, USA) using 100 ng of cDNA and 10 pmol of both forward and reverse primers per 25  $\mu$ l reaction. All qPCR reactions were carried out in triplicate. Amplification and fluorescence measurements were carried out on iCycler iQ Real-Time PCR Detection Instrument (Bio-Rad Laboratories, Hercules, CA) with the following thermocycling parameters: 2 min at 50°C and 10 min at 95°C, followed by 60 amplification cycles of 15 s at 95°C and 1 min at 58°C. A melting curve was constructed for the amplification product by increasing the temperature from 55°C by 0.5°C per cycle for 80 cycles. Standard curves were generated using a 10-fold dilution series of the Merit cDNA sample and used to determine the PCR efficiency for each target transcript amplification. The unregulated 18S rRNA transcript was selected as suitable endogenous control and used for normalization during relative quantification of target genes (Pfaffl 2001).

#### Construction of subtracted cDNA library from malted barley seeds and DNA sequencing

A cDNA library was constructed from malted seeds of cv. Morex using suppression subtractive hybridization (SSH)

technique (Diatchenko et al. 1996). Total RNA was extracted from dry seed (control) and seed at day 4 micromalting (96 h at 20°C) as described above. cDNA populations were derived from the two RNA samples using the SMART cDNA synthesis system (BD Biosciences-Clontech, Palo Alto, CA). SSH was performed between these two cDNA populations designating the one derived from dry seed as “Driver” and the population derived from malted seed as “Tester”. Forward PCR-based subtraction was performed (Tester minus Driver) and the amplification products ligated into the pTAdvantage PCR cloning vector (BD Biosciences-Clontech, Palo Alto, CA). *E. coli* was transformed with this ligation reaction by electroporation. Aliquots of the plasmid library were plated on Luria Broth plates containing the appropriate antibiotic and grown overnight at 37°C. Seven hundred sixty-eight white clones were picked onto 96-well plates. DNA sequences for ~650 clones were determined by Sanger sequencing (Macrogen Inc., Maryland, USA and Washington University Genome Sequencing Center, St. Louis, MO).

#### Annotation of ESTs

Vector sequences, polyA and polyT stretches, were trimmed from the DNA sequences. Sequences less than 100 bp were discarded. Annotation of individual ESTs was performed using Basic Local Alignment Search Tool (BlastN and tBlastX) (Altschul et al. 1990) searches against the non-redundant protein sequences at NCBI (<http://www.ncbi.nlm.nih.gov/blast>). Best matches ( $E$ -value <  $10^{-10}$ ) were used for putative gene assignments. Nucleotide BLAST searches against the EST database at NCBI was conducted for ESTs showing no significant hits with protein sequences. Functional categories were assigned on the basis of biological, functional and molecular annotations available from the gene ontology (GO) database (<http://www.geneontology.org/>). The sequences were deposited into the NCBI dbEST resource using NCBI sequence-submission protocols.

## Results

### Transcript profiles of different malting stages

The number of differentially expressed genes (i.e., probe sets with absolute value of  $\log_2$  fold change >1 and adjusted  $P$ -values of  $\leq 0.05$ ) increased as barley seeds went from steeping to 24 h germination (day 1), 93 h germination (day 4), and kilning (Table 3). During steeping, when seeds imbibed water, 966 probe sets were up-regulated in comparison to dry seed. Categories of the genes based on biological processes assigned using the PLAN program

(He et al. 2007) are shown in Table 3 (see Supplemental Tables S1 for expression data and S2 for list of genes under each category). Altogether, genes involved in metabolism of carbohydrates, nucleotides, amino acids, and proteins, made up 15% of the genes induced during this stage. Another large group of genes (~15%) were those involved in response to stimulus, including pathogens, chemicals, and abiotic stress. Following steeping, extensive reprogramming occurred as shown by the dramatic increase in numbers of induced genes in day 1 and day 4 malting (1,432 and 1,997 genes, respectively). Genes involved in metabolism of macromolecules (i.e., proteins, amino acids, nucleotides, carbohydrates, and lipids) continued to increase from steeping to day 4. Genes involved in transport more than doubled from steeping to day 4. The number of up-regulated genes decreased slightly during kilning (1466). At this stage when the seed was subjected to heat, the number of genes involved in metabolism of macromolecules decreased. Genes involved in cell organization and development decreased by more than half. In comparison, the percentage of genes involved in stress response slightly increased.

The number of down-regulated genes increased from steeping to kilning (Table 3). While up-regulated genes decreased from day 4 to kilning, down-regulated genes continued to increase during this transition. The kilning stage contained the largest number of down-regulated genes (2135). Down-regulated genes involved in metabolism of macromolecules increased from day 4 to kilning, reflecting the necessity to decrease or shut down these processes. The exception is carbohydrate metabolism where down-regulated genes did not increase from day 4 to kilning. The number of down regulated genes for stress response decreased from day 4 to kilning, indicating the possible need for induction of these genes. Genes for which there were no significant homology with DNA sequences in GenBank made up approximately 40–50% of the differentially expressed genes during the different stages of malting.

### Differentially expressed genes among four malting barley cultivars

To determine the best stages to use for comparing gene expression during malting in different barley cultivars, we investigated the expression patterns of barley genes that had been previously associated with one or more malting quality phenotypes. The list included genes encoding starch degradation enzymes, cell wall degrading enzymes, protein hydrolytic enzymes, and proteinase inhibitors (Jones 2005a, b; Potokina et al. 2004). Seventy probe sets representing these genes were identified and hierarchical clustering was performed according to expression patterns in dry seed and the four malting stages of Morex (Supplemental

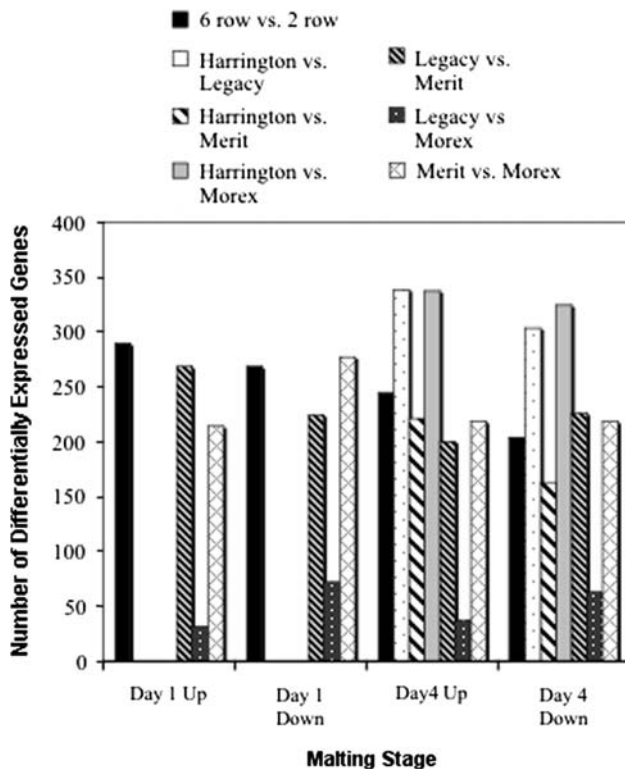
**Table 3** Number and functional categories of differentially expressed genes in four malting stages relative to expression in dry seed

Functional category	GO Category	Up-regulated				Down-regulated			
		Steep	Day 1	Day 4	Kiln	Steep	Day 1	Day 4	Kiln
Regulation of cellular physiological process	GO:0050794	14	22	28	18	6	17	21	48
Carbohydrate metabolism	GO:0005975	25	32	37	25	6	14	13	6
Alcohol metabolic process	GO:0006066	17	19	17	10	4	8	7	5
Generation of precursor metabolites and energy	GO:0006091	7	12	29	18	8	11	13	11
Nucleobase, nucleoside, nucleotide and nucleic acid metabolism	GO:0006139	38	50	63	30	9	24	31	92
Amino acid and derivative metabolism	GO:0006519	21	34	49	34	2	9	10	17
Lipid metabolic process	GO:0006629	12	22	30	20	11	15	15	17
Cell wall organization and biogenesis	GO:0007047	6	6	10	7	0	2	2	3
Cell cycle	GO:0007049	5	7	9	3	0	2	2	7
Signal transduction	GO:0007165	10	16	16	10	4	10	12	19
Sensory perception	GO:0007600	0	1	2	2	0	0	0	0
Photosynthesis	GO:0015979	0	1	7	4	1	1	1	0
Cell organization and biogenesis	GO:0016043	43	54	67	24	12	25	28	90
Cell growth	GO:0016049	4	3	5	1	2	1	1	5
Regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolism	GO:0019219	9	14	19	13	4	14	17	35
Protein metabolic process	GO:0019538	53	85	106	44	15	33	38	129
Development	GO:0032502	38	47	59	27	13	25	36	76
Negative regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolism	GO:0045934	4	5	3	1	0	0	0	8
Cell division	GO:0051301	4	2	3	1	0	3	3	4
Response to stimulus	GO:0050896	57	76	119	116	34	80	80	79
Response to stress	GO:0006950	29	36	60	69	21	53	45	36
Defense response	GO:0006952	10	16	27	23	6	8	15	14
Response to hormone stimulus	GO:0009725	14	15	23	22	8	15	17	18
Response to other organism	GO:0051707	9	16	27	25	6	9	16	13
Response to chemical stimulus	GO:0042221	20	34	55	51	14	36	35	27
Response to abiotic stimulus	GO:0009628	21	25	42	49	21	44	36	36
Transport	GO:0006810	26	49	62	39	11	24	27	40
Other		227	349	509	316	147	326	355	609
Unique targets		444	706	989	604	244	538	590	1,015
No hits		472	608	815	738	289	614	796	1,042
Total unique probe sets		966	1,432	1,997	1,466	546	1,211	1,458	2,135

Fig. S1). A majority of the genes were down-regulated in the dry seed and steeping stage. More than 90% of the probe sets were highly expressed during either or both day 1 and day 4 germination stages. Day 4 contained the largest number of up-regulated genes, consistent with the pattern for all differentially expressed genes in the different malting stages. Based on these results, we decided to use day 1 and day 4 germination stages to investigate expression patterns in three additional cultivars.

Pairwise comparisons of expression patterns between cultivars show that the largest numbers of differentially

expressed genes were in comparisons between 2-row and 6-row cultivars (Fig. 2). In day 1, the largest number of differentially up-regulated genes was in the combined 2-row versus 6-row comparison, followed by Legacy (6-row) versus Merit (2-row), then Merit versus Morex (6-row) (see Supplemental Table S3 for list of genes). A small number of genes (~30) were differentially up-regulated between Legacy and Morex. A similar trend was observed for differentially down-regulated genes in day 1, with the exception that a greater number of differentially expressed genes were found between Merit and Morex compared to Legacy and

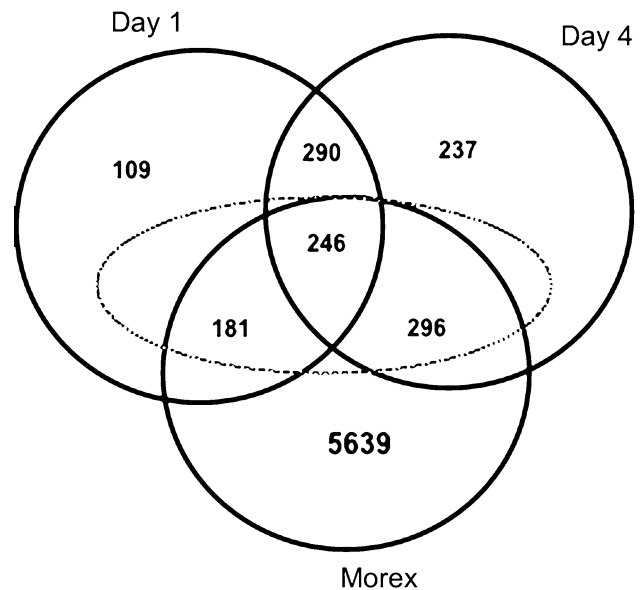


**Fig. 2** Number of differentially expressed genes in all possible pairwise comparisons between the four malting barley cultivars

Merit. Pairwise comparisons involving Harrington were not available in day 1 since there was only one replication. For day 4, the largest numbers of differentially up-regulated genes were found in comparisons between Harrington and the 6-row cultivars, Legacy and Morex (see Supplemental Table S4 for list of genes). This was followed by combined 2 row versus 6 row, and Harrington versus Merit and Merit versus Morex. The Legacy versus Morex comparison showed the least number of differentially expressed genes (~30). The pattern for down-regulated genes for day 4 was similar to that of up-regulated genes. In general, the lowest numbers of differentially expressed genes were found in comparisons between same row-class cultivars (Merit vs. Harrington and Legacy vs. Morex).

#### Correlation of differentially expressed genes with malting quality phenotypes

Genes that were differentially expressed compared to dry seed from the Morex time course experiment are potentially important in determining malting quality traits. However, many genes expressed during these stages are involved with growth and development and may not contribute directly to malting quality parameters. Similarly, many genes that were differentially expressed between the cultivars may potentially underlie malting quality trait differ-



**Fig. 3** Venn diagram showing the overlap between differentially expressed genes in four malting stages of Morex, and differentially expressed genes in the cultivars-comparisons for day 1 and day 4. The broken-lined ellipse shows the 723 genes that overlapped between the Morex malting stages, day 1 cultivars-comparisons, and day 4 cultivars-comparisons. Expression levels of these 723 genes were used to correlate with malting quality phenotypes

ences among the cultivars. However, since the cultivars have different genetic backgrounds, many genes identified in these comparisons may not be directly related to the malting quality differences among the cultivars. In an attempt to identify genes that may be the most relevant to malting quality traits, we set the following criteria: the genes had to be differentially expressed in at least one of the four malting stages in the Morex time course experiment, as well as differentially expressed in the cultivars comparison for day 1 or day 4 (Fig. 3). There were a total of 723 genes that overlapped between Morex malting stages and at least one or both of the two malting stages for all cultivar comparisons (see Supplemental Table S5 for list of genes).

We examined the correlation between the expression values of the 723 genes and the malting quality phenotypes using a cut-off value  $P \leq 0.05$ , and requiring more than 50% of the sites to have correlation values  $\geq 0.7$ . Correlations with six malting quality traits ( $\alpha$ -amylase, diastatic power, free amino nitrogen, fine extract, malting protein, and soluble/total protein ratio) were detected (Table 4). Eleven to 72 genes showed positive correlation while 19–102 genes showed negative correlation with these traits. Representative genes correlated with the malting quality phenotypes are listed in Table 5 (see Supplemental Table S6 for complete list). The list includes genes involved in metabolism of carbohydrates, proteins, lipids, and amino acids, genes involved in transport, cell wall organization and biogenesis, as well as genes for stress and



**Table 4** Summary of genes showing positive (+) or negative (-) correlations with six malting quality phenotypes

Stage	$\alpha$ -amylase		Diastatic power		Free amino nitrogen		Fine extract		Malt protein		S/T protein	
	+	-	+	-	+	-	+	-	+	-	+	-
Day 1	8	10	38	55	9	15	27	72	37	14	8	15
Day 4	6	10	49	54	7	22	30	64	30	14	5	23
Total	12	19	72	85	15	33	47	102	54	26	11	34

defense response. Genes for carbohydrate metabolism include genes that were previously associated with malting quality traits (Clark et al. 2003; Jones 2005a, b; Potokina et al. 2004).  $\beta$ -Glucosidase was positively correlated with diastatic power while *limit dextrinase* was positively correlated with fine extract. Other genes coding for carbohydrate metabolic enzymes, such as *Sucrose synthase 1* and  $\beta$ -fructofuranidase 1, were also positively correlated with diastatic power. On the other hand, many genes coding for enzymes involved in glycolysis negatively correlated with fine extract, diastatic power, soluble/total protein ratio, and free amino nitrogen. Lipid metabolic enzyme genes showing positive correlations with the malting quality traits included *Lipoxygenase 2.1*, several *lipases*, and *oleosin*. Protein inhibitor genes including  $\alpha$ -amylase inhibitor, *protein synthesis inhibitor 1*, and *lipid transfer protein (LTP)* showed negative correlation with many of the malting traits (Table 5). Other genes showing correlation with malting quality phenotypes are those involved in transport of proteins and macromolecules and genes involved in defense and stress response. Genes with unclear functions as well as genes with no significant homology were also identified.

#### Characterization of barley ESTs from malted seeds

To identify other barley genes that may be important in malting, ESTs were generated from a cDNA library made by suppression-subtractive hybridization between cDNA made from day 4 malted Morex seeds and cDNA from Morex dry seeds. Suppression-subtractive hybridization was performed to enrich for genes expressed during day 4 germination. A total of 509 ESTs were produced and grouped according to functional categories (Fig. 4 and Supplemental Table S7). Fifty-six percent of the genes were of unknown function. Many of these did not produce BLASTX hits with proteins. Significant homologies were only found with barley ESTs and may be barley-specific genes. Other genes in this group showed homology to hypothetical proteins or to ESTs from other plant or animal species. Eleven percent of the ESTs showed no significant homology to sequences in the databases. Among the genes that could be classified into functional categories, the largest group (9%) was involved in defense and stress response.

*Jasmonate-induced protein* genes made up 26% of the genes in this group. Genes involved in lipid biosynthesis made up approximately 6% of all ESTs, although 96% of the ESTs in this group consisted of  $\beta$ -ketoacyl ACP synthase. Surprisingly, this gene was not differentially expressed during malting in any of the four cultivars examined. Genes involved in gene regulation, protein metabolism, and carbohydrate metabolism made up 4.3, 3.9, and 3.1% of the ESTs, respectively.  $\alpha$ -Amylase genes were the largest group of genes under carbohydrate metabolism (Supplemental Table S7). Other ESTs identified were involved in metal ion binding (2.2%), membrane biosynthesis (1.4%), amino acid metabolism (1.8%), energy production (1%), cell wall biogenesis (0.6%), and seed storage protein (0.4%).

#### Quantitative RT-PCR of selected genes

To validate gene expression observed in microarray experiments, qRT-PCR of four selected genes ( $\alpha$ -amylase,  $\alpha$ -glucosidase, *limit dextrinase*, and  $\beta$ -ketoacyl synthase) was conducted on the four malting cultivars using cDNA prepared from day 4 malted seeds (Fig. 5a). cDNA from day 4 malted seed of a non-malting cultivar, Steptoe, was included for comparison. For all four genes, there was a significant difference in expression levels among the five cultivars (Fig. 5a).  $\alpha$ -Amylase and  $\alpha$ -glucosidase expression levels were significantly different among the cultivars at  $P \leq 0.0001$ . *Limit dextrinase* and  $\beta$ -ketoacyl synthase expression levels were significantly different among the cultivars at  $P \leq 0.001$  and 0.01, respectively. In comparison, only  $\alpha$ -amylase and *limit dextrinase* showed differential expression in microarrays at day 4 as defined by  $|\log_2(\text{FC})| > 1$  and adjusted  $P \leq 0.05$  (Fig. 5b). There were also differences in expression patterns between qRT-PCR and microarrays for  $\alpha$ -amylase expression (Fig. 5a, b). Merit and Legacy showed the highest levels of  $\alpha$ -amylase expression in qRT-PCR while Harrington showed the highest level of  $\alpha$ -amylase expression in microarrays. For *limit dextrinase*, Merit and Harrington showed the highest levels of expression in both qRT-PCR and microarrays (Fig. 5a, b). Therefore, there was limited correlation between expression patterns observed with qRT-PCR and microarrays.

**Table 5** List of representative genes correlated with malting quality phenotypes

Functional category	Probe set ID	Trait	Correlation coefficient	<i>P</i> Value	No. of positive trials	No. of negative trials
Carbohydrate metabolism						
Sucrose/starch metabolism						
Limit dextrinase	Contig11648_at	FE	0.976	0.024	23	1
		MPRT	−0.961	0.039	3	15
Glucan endo-1,3-beta-glucosidase	Contig5219_s_at	FE	0.996	0.004	20	1
		Beta-glucosidase	Contig2736_s_at	DP	0.975	0.025
Beta-fructofuranosidase 1	Contig3392_at	DP	−0.956	0.044	0	21
Beta-fructofuranosidase 5	Contig4470_s_at	ST	0.952	0.048	18	3
Sucrose synthase 1	Contig361_s_at	DP	0.984	0.016	22	0
Glycolysis						
Pyruvate kinase	Contig3010_s_at	FAN	−0.989	0.011	0	7
		FE	−0.967	0.033	1	20
		ST	−0.970	0.030	3	19
Glyceraldehyde-3-phosphate dehydrogenase B	baak1k18_s_at	DP	−0.975	0.025	0	22
Phosphoenolpyruvate carboxylase kinase	Contig10338_at	FE	−0.995	0.005	1	21
		MPRT	0.982	0.018	16	3
Protein metabolism						
Alpha-amylase/trypsin inhibitor	Contig566_s_at	AA	−0.954	0.046	0	25
		FAN	−0.985	0.015	0	7
		FE	−0.969	0.031	1	20
		ST	−0.955	0.045	3	19
Cysteine proteinase EP-B 1 precursor	Contig5278_at	DP	−0.957	0.043	1	21
Protein synthesis inhibitor I	Contig572_at	DP	−0.963	0.037	0	22
Lipid transfer protein (LTP) family protein	Contig7968_at	FE	0.997	0.003	21	1
		MPRT	−0.983	0.017	3	15
Nonspecific lipid transfer protein 2	Contig9069_at	FE	0.973	0.027	20	1
Protease inhibitor	HVSMEh0099O01f_s_at	FE	0.976	0.024	22	1
Subtilisin-chymotrypsin inhibitor CI-1B	Contig34_s_at	FE	−0.977	0.023	1	23
		FE	−0.965	0.035	1	23
Amino acid metabolism						
Arginine decarboxylase	Contig5994_s_at	FE	−0.984	0.016	1	21
		MPRT	0.964	0.036	15	3
Serine carboxypeptidase 3	Contig600_at	DP	−0.980	0.020	0	22
Carbon fixation						
Transketolase	Contig1618_at	DP	−0.950	0.050	1	20
Cytochrome c oxidase subunit VIa	Contig4843_at	AA	−0.959	0.041	0	25
		FAN	−0.952	0.048	0	6
Lipid metabolism						
Lipoxygenase 2.1	Contig2306_s_at	AA	0.974	0.026	26	1
Lipase	Contig10522_at	DP	−0.959	0.041	0	20
		FE	0.972	0.028	22	1
Cholinephosphate cytidylyltransferase	rbaal36m18_s_at	FE	0.974	0.026	23	1
Oleosin Bn-V	Contig19860_at	FE	1.000	−0.989	0.011265623	1
		MPRT	1.000	0.997	0.003087643	14

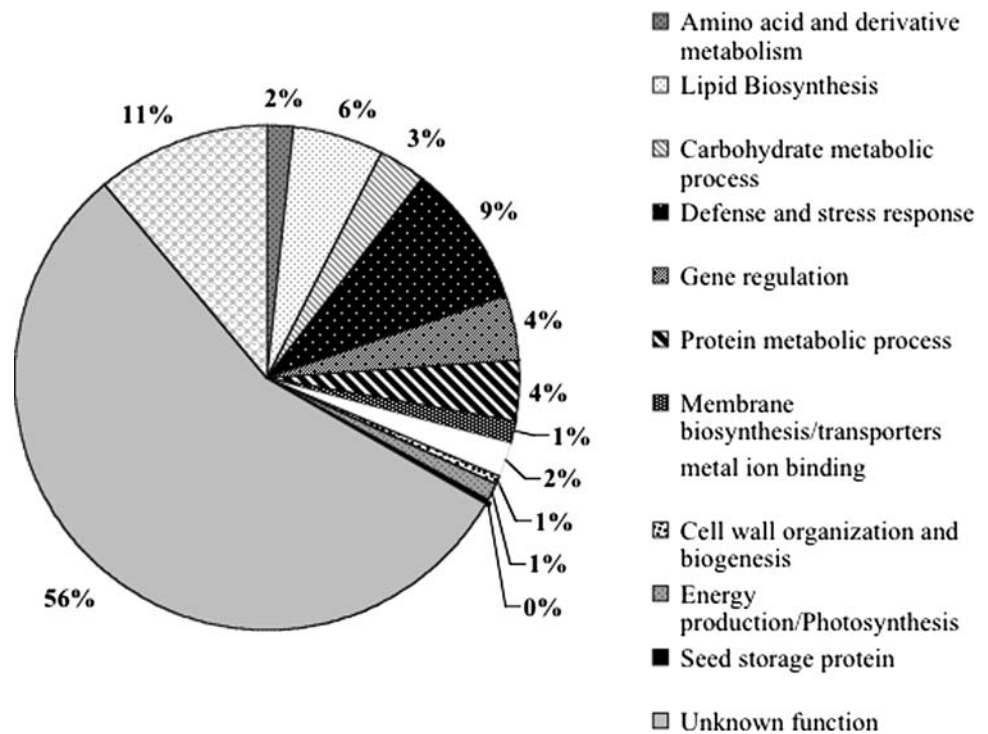
**Table 5** continued

Functional category	Probe set ID	Trait	Correlation coefficient	<i>P</i> Value	No. of positive trials	No. of negative trials
Lipase class 3 family protein	Contig20537_at	DP	1.000	0.955	0.04521111	22
Transport						
Proton-dependent oligopeptide transport (POT) family protein	Contig20130_at	FE	0.963	0.037	22	1
		FE	0.963	0.037	22	1
Protein aquaporin TIP3.1	Contig3772_at	ST	−0.969	0.031	3	18
Cell wall organization and biogenesis						
Caffeic acid <i>O</i> -methyltransferase	Contig2528_x_at	FE	0.964	0.036	23	1
	Contig2539_s_at	FE	−0.976	0.024	1	23
		FE	−0.979	0.021	1	23
Aldehyde oxidase-like protein	Contig8610_at	ST	−0.962	0.038	3	18
Xylanase inhibitor precursor	Contig14679_at	DP	0.995	0.005	22	0
Gene regulation						
Transcription elongation factor 1	Contig7809_at	FE	−0.964	0.036	1	22
WIR1A protein	Contig5974_s_at	DP	0.962	0.038	20	0
		FAN	−0.980	0.020	0	7
		ST	−0.997	0.003	3	18
Type A response regulator 4	Contig5952_at	DP	−0.979	0.021	0	22
Leucine-rich repeat family protein	Contig518_s_at	DP	−0.960	0.040	0	20
ATAPY1 (APYRASE 1); calmodulin binding	HW09M17u_at	FE	−0.984	0.016	1	21
		MPRT	0.989	0.011	14	1
Storage proteins						
B1 hordein— <i>Hordeum vulgare</i> (Barley)	Contig69_x_at	FE	−0.981	0.019	1	21
		MPRT	0.957	0.043	15	3
Embryo globulin	Contig1353_s_at	DP	0.967	0.033	21	0
	Contig1356_s_at	FE	−0.970	0.030	1	23
Defense response						
Heat shock factor protein hsf8-like	Contig9927_s_at	FE	−0.977	0.023	1	23
		MPRT	0.965	0.035	15	3
Jasmonate <i>O</i> -methyltransferase	EBro02_SQ005_I23_at	DP	0.956	0.044	20	0
Jasmonate-induced protein	Contig3499_at	FE	0.958	0.042	22	1
Stress responsive protein	Contig1249_s_at	DP	−0.956	0.044	0	20
		DP	−0.983	0.017	0	24
		FE	0.960	0.040	20	1
		MPRT	−0.979	0.021	1	14
Oxalate oxidase GF-2.8	Contig3017_at	DP	0.987	0.013	23	0
Oxalate oxidase-like protein	Contig3156_s_at	AA	−0.956	0.044	0	26
		FAN	−0.988	0.012	0	7
		FE	−0.985	0.015	1	21
		MPRT	0.997	0.003	14	1
		ST	−0.968	0.032	3	18
Small heat shock protein HSP17.8	Contig3286_s_at	FAN	0.963	0.037	7	0
		ST	0.992	0.008	19	3
No Significant hit						
	Contig9743_at	AA	1.000	0.000	26	0
	HV10I20u_at	AA	0.989	0.011	25	0
	HV10I20u_at	AA	0.977	0.023	27	0

**Table 5** continued

Functional category	Probe set ID	Trait	Correlation coefficient	<i>P</i> Value	No. of positive trials	No. of negative trials
	Contig16067_at	DP	0.999	0.001	23	0
	Contig16067_at	DP	1.000	0.000	24	0
	Contig1852_at	DP	0.995	0.005	23	0
	Contig23146_at	DP	0.998	0.002	23	0
	Contig23396_at	DP	0.964	0.036	21	0
	Contig3233_at	DP	1.000	0.000	24	0
	HT09D15r_s_at	DP	0.994	0.006	23	0
	HVSMEi0006K11r2_at	DP	0.997	0.003	23	0

**Fig. 4** Functional categories of barley ESTs generated from a cDNA library made by suppression-subtractive hybridization between cDNA from ungerminated seed of Morex as ‘driver’ and cDNA from day 4 malted seed of Morex as tester



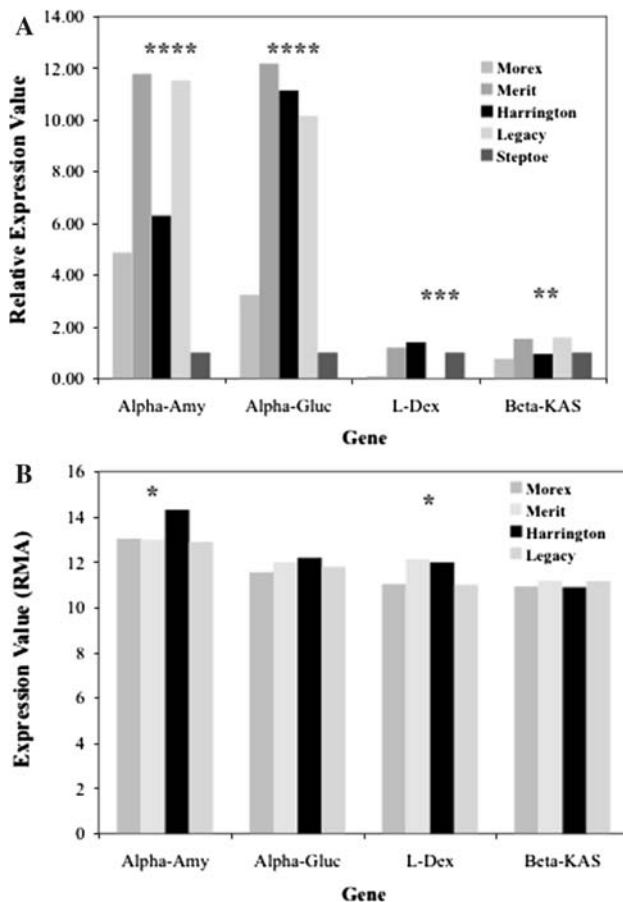
Including Steptoe in qRT-PCR experiments allowed comparison of gene expression between the four malting cultivars and a non-malting cultivar. Steptoe showed at least three times lower expression levels for  $\alpha$ -amylase and  $\alpha$ -glucosidase than the lowest expressing malting cultivar, Morex (Fig. 5a). For *limit dextrinase*, Legacy had lower expression than Steptoe, while for  $\beta$ -ketoacyl synthase, Steptoe, Morex, and Harrington had similar expression levels.

## Discussion

Transcript profiles of germinating barley seeds

The use of the Barley1 GeneChip<sup>®</sup> array in this study provided a comprehensive view of global changes in gene

expression during germination in barley. Up-regulation of genes involved in metabolism of carbohydrates, nucleotides, amino acids, and proteins during steeping indicates that a reactivation of metabolism was occurring. Twenty-four hours following steeping (day 1), extensive reprogramming of gene expression occurred, similar to what others have previously reported (Watson and Henry 2005; White et al. 2006). Every functional category included in the analysis increased in the number of induced genes during the malting stages. Doubling in number of induced genes involved in transport from steeping to day 4 may reflect the importance of protein movement from embryo and scutellum to the endosperm (Fincher 1989). Hydrolytic enzymes synthesized in the embryo are transported to the endosperm, where they break down the carbohydrate and protein reserve (Bamforth and Barclay 1993; Macfadden et al.



**Fig. 5** Bar charts showing expression levels of  $\alpha$ -amylase (Alpha-Amy),  $\alpha$ -glucosidase (Alpha-Gluc), limit dextrinase (L-Dex), and  $\beta$ -ketoacyl synthase (Beta-KAS) based on (a) qRT-PCR, and (b) microarrays during day 4 germination in Morex, Merit, Harrington, and Legacy. For A, a non-malting cultivar, Steptoe, was included in the qRT-PCR experiments. Relative quantification of transcript abundance was performed using Steptoe as calibrator; this sample therefore had a baseline relative fold change of one. Asterisks denote *P*-values based on ANOVA: \*\*\*\*  $\leq 0.0001$ , \*\*\*  $\leq 0.001$ , \*\*  $\leq 0.01$ , \*  $\leq 0.05$ . Differential expression in B was based on absolute  $\log_2$  fold change  $>1$  and  $P \leq 0.05$

1988; Ranki 1990). In addition, transcripts for hydrolytic enzymes are also transcribed or stored in the endosperm as recently reported (Sreenivasulu et al. 2008). In this recent study, the authors investigated the transcriptome of barley during maturation, desiccation, and germination, separating the endosperm plus aleurone from the embryo. They showed that transcriptional activation of storage reserve mobilization events occurs at an early stage during germination, well before protrusion.

Genes involved in stress and defense response made up a major portion of differentially expressed transcripts in all four malting stages studied. Previous studies of germinating barley seeds have reported similar findings (Potokina et al. 2002; Watson and Henry 2005; White et al. 2006). This is not surprising since malting is not performed under aseptic

conditions and barley seeds contain an inherent microbial flora consisting of fungi, bacteria, and yeasts (Flannigan 2003). It has been shown that the molds and bacteria in barley increase during malting, reaching a peak during the germination stage (Flannigan 2003), consistent with the observation that defense and stress response genes increased from steeping to day 4. Analysis of the proteome of malt and beer also showed the predominance of proteins involved in protection against pathogens and insects (Perrocheau et al. 2005). In addition, seed germination may be considered a critical time point in a plant's life cycle and it is not surprising that a large number of defense-related genes are normally upregulated as a protective measure.

While previous studies of gene expression in germinating barley seeds mainly focused on steeping and germination stages, we also looked at kilning. Exposure of the seeds to gradually increasing temperatures resulted in the overall decrease in number of up-regulated genes involved in metabolism of macromolecules. This is to be expected as kilning is done to terminate the germination process. Some genes that remained up-regulated from germination through kilning included those coding for  $\alpha$ -amylase, limit dextrinase,  $\beta$ -glucosidase, and  $\beta$ -glucanase, indicating that the enzymes are thermostable. Other genes that were up-regulated from germination through kilning included those coding for peptidases, lipid transfer proteins, and proteinase inhibitors. These proteins were identified as thermostable components of malt and beer (Perrocheau et al. 2005). Other genes that were up-regulated from germination to kilning were those for heat shock proteins, which function to protect enzymes during heat stress (Jinn et al. 1995; Waters et al. 1996).

#### Genes correlated with malting quality phenotypes

The strategy used to identify barley genes associated with malting quality was to focus on genes that were differentially expressed in both the malting stages in Morex and in the four-cultivar comparisons. Using this approach, we took advantage of differential expression between the malting stages relative to dry seed as well as differential expression among four malting cultivars. There were 723 genes that met the criteria. To address the concern that this approach might be too stringent and might eliminate genes that are important in malting, we determined whether genes that have been previously associated with malting quality phenotypes were represented among the 723 genes. The genes for  $\alpha$ -amylase, limit dextrinase, and  $\alpha$ -glucosidase were in this group of genes. The endosperm-specific  $\beta$ -amylase gene,  $\beta my1$ , has also been associated with diastatic power (Clark et al. 2003) but was not included in the list, since  $\beta my1$  is transcribed during seed development (Daussant and Corvazier 1970). Other genes previously

identified based on differential expression and correlation with malting quality traits including  $\beta$ -1,3 *glucanase*, *Sucrose synthase I*, and *Lipoxygenase* (Potokina et al. 2004), were also among the 723 genes. Therefore, it appears that the approach used was able to identify genes that are potentially important in malting.

Of the 723 genes, 11–102 genes correlated with six malting quality phenotypes. Well-characterized genes for hydrolytic enzymes known to be important in malting (e.g., *Limit dextrinase*,  $\beta$ -*glucanase*) showed the expected association with malting quality phenotypes (Table 5). Many other genes that have not been characterized for their roles in determining specific malting quality traits were identified in this study. For example,  $\alpha$ -amylase measurements were negatively correlated with genes coding for  $\alpha$ -amylase inhibitors and positively correlated with *Lipoxygenase 2.1*. Diastatic power measurement, which is an indication of  $\alpha$ -amylase and  $\beta$ -amylase activities, was correlated with genes for carbohydrate metabolism enzymes, as well as genes for lipid metabolism, several defense and stress response genes, and genes of unknown function.

It is noteworthy that many of the genes showing correlation with malting quality traits were also previously shown to be associated with the same or different malting quality traits in this current report (Potokina et al. 2004). The previous study investigated the expression of 1400 ESTs in ten malting barley varieties different from those used in our study. Examples of common genes that correlated with malting quality traits in the two studies are those coding for  $\beta$ -glucosidase, sucrose synthase I, limit dextrinase, cysteine protease,  $\beta$ -glucanase, lipid transfer proteins, lipoxygenase 2.1, lipases, peroxidase, and heat shock proteins.

The finding that several proteinase inhibitor genes correlated with malting quality traits agrees with the suggestion by Jones (2005b) that it is important to define the hydrolytic enzyme inhibitors in malting barley since these proteins can regulate the activity of the endoproteases. It has been recognized that current measurements of malting quality are not precise since they do not identify all enzymes that contribute to a malting quality trait (Bamforth and Barclay 1993). The results presented here may provide candidate genes that are involved in malting quality traits. A caveat to keep in mind is that these results are exploratory and the role of these candidate genes in determining malting quality will need to be validated.

One way to validate candidate genes is by determining whether the gene or the expression level polymorphism maps to a malting quality QTL, as previously demonstrated (Potokina et al. 2006). A candidate gene (*CxpI*) and the relative expression level polymorphism both mapped to a region containing a QTL for diastatic power on chromosome 3H (Potokina et al. 2006). Another potential approach for validating the function of genes identified in this study

would be to knock down gene expression using RNA interference or TILLING (McCallum et al. 2000; Till et al. 2003). TILLING lines developed in malting barley cultivars are available (Caldwell et al. 2004). The use of TILLING lines provides an advantage in that transformation is not required, since barley transformation is still a technical challenge and requires a particle bombardment facility. Identification of SNPs associated with malting quality traits would provide another confirmation for the function of candidate genes in malting as well as markers that can be used by breeders. SNP analysis of the *CxpI* gene in 90 barley cultivars identified two haplotypes, one of which showed association with diastatic power (Potokina et al. 2006).

ESTs from malted barley seed include novel genes

The majority of ESTs generated from a cDNA library made by suppression-subtractive hybridization between malted and dry seed were in common with the 723 differentially expressed genes identified in the microarray experiments. Furthermore, many of the ESTs corresponded to genes correlated with malting quality phenotypes (Supplemental Table S7). Some examples are *limit dextrinase*, *lipid transfer protein*,  $\alpha$ -*amylase inhibitors*, and *jasmonate-induced proteins*.

Approximately 6% of the ESTs generated showed homology to  $\beta$ -*ketoacyl ACP synthase*. This gene was not differentially expressed in any of the malting stages of the four cultivars investigated using microarrays. However, qRT-PCR results showed significant differences ( $P < 0.01$ ) in the expression level of this gene among the four cultivars, with transcript levels in Morex lower than in the non-malting cultivar Steptoe. Whether this gene plays an important role in malting is not clear.  $\beta$ -ketoacyl ACP synthase catalyzes the chain elongation step in fatty acid synthesis leading to palmitoyl-ACP and stearoyl ACP (Kauppinen 1992). Other genes involved in lipid metabolism were found to be correlated with malting quality traits in this study and in previous studies (Potokina et al. 2004).

## Conclusion

In conclusion, transcript profiling using microarrays and ESTs identified a common set of genes that were differentially expressed during malting in barley. Correlation of expression profiles with malting quality phenotypes resulted in the identification of 11–102 genes showing correlation with six malting quality traits. These candidate genes will need to be further validated for their importance in malting. A large group of genes with unknown function were identified with both methods, emphasizing the fact that the molecular basis of malting quality traits is not well

understood. Finally, ESTs generated in this study with no known homology to sequences in the genome database may be novel genes and these may be important in malting.

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