Interchromosomal Transfer of Immune Regulation During Infection of Barley with the Powdery Mildew Pathogen

Priyanka Surana,† Ruo Xu,‡ Gregory Fuerst,§ Antony V. E. Chapman,*,**, Dan Nettleton,*,† and Roger P. Wise*,†,**

*Interdepartmental Bioinformatics and Computational Biology Program, †Department of Plant Pathology and Microbiology, ‡Department of Statistics, §Com Insects and Crop Genetics Research, United States Department of Agriculture-Agricultural Research Service, and **Interdepartmental Genetics and Genomics, Iowa State University, Ames, Iowa 50011

ORCID IDs: 0000-0002-7167-0875 (P.S.); 0000-0001-7786-1528 (R.P.W.)

ABSTRACT Powdery mildew pathogens colonize over 9500 plant species, causing critical yield loss. The Ascomycete fungus, Blumeria graminis f. sp. hordei (Bgh), causes powdery mildew disease in barley (Hordeum vulgare L.). Successful infection begins with penetration of host epidermal cells, culminating in haustorial feeding structures, facilitating delivery of fungal effectors to the plant and exchange of nutrients from host to pathogen. We used expression Quantitative Trait Locus (eQTL) analysis to dissect the temporal control of immunity-associated gene expression in a doubled haploid barley population challenged with Bgh. Two highly significant regions possessing trans eQTL were identified near the telomeric ends of chromosomes (Chr) 2HL and 1HS. Within these regions reside diverse resistance loci derived from barley landrace H. laevigatum (MLa) and H. vulgare cv. Algerian (Mla1), which associate with the altered expression of 961 and 3296 genes during fungal penetration of the host and haustorial development, respectively. Regulatory control of transcript levels for 299 of the 961 genes is reprioritized from MLa on 2HL to Mla1 on 1HS as infection progresses, with 292 of the 299 alternating the allele responsible for higher expression, including Adaptin Protein-2 subunit μ AP2M and Vesicle Associated Membrane Protein VAMP72 subfamily members VAMP721/722. AP2M mediates effector-triggered immunity (ETI) via endocytosis of plasma membrane receptor components. VAMP721/722 and SNAP33 form a Soluble N-ethylmaleimide-sensitive factor Attachment Protein REceptor (SNARE) complex with SY121 (PEN1), which is engaged in pathogen associated molecular pattern (PAMP)-triggered immunity via exocytosis. We postulate that genes regulated by alternate chromosomal positions are repurposed as part of a conserved immune complex to respond to different pathogen attack scenarios.

KEYWORDS immune regulation expression quantitative trait locus (eQTL) barley (Hordeum vulgare) Blumeria graminis host–pathogen interactions
resistance protein interacts, directly or indirectly, with corresponding pathogen-secreted effector proteins, which alter host processes to promote nutrient acquisition and colonization (Bent and Mackey 2007; Jacob et al. 2013; Cesari et al. 2014). This interaction triggers defense signaling in the cytoplasm or the nucleus, resulting in programmed cell death (Li et al. 2015).

Reaction to Bgh is controlled by several loci distributed across the barley genome, termed Mi (for mildew resistance locus). Positioned on the short arm of Chr 1H (Wei et al. 2002), the Mla locus harbors over 30 alleles that confer varying levels of resistance when corresponding avirulence (AVR) effectors are present in the pathogen (Shen et al. 2003; Halterman and Wise 2004; Seeholzer et al. 2010). Mla alleles encode coiled-coil, nucleotide binding, leucine-rich repeat (NLR) proteins that accumulate in the nucleus after recognition of corresponding AVR effector proteins from Bgh (Li et al. 2015). After recognition, MLA dissociates the MYB6 transcriptional activator from the WRKY1 repressor and promotes its binding to corresponding cis-regulators, initiating disease resistance signaling (Chang et al. 2013).

Despite the observation that plant resistance to disease is often associated with single segregating loci, the underlying genetic architecture that contributes to the ultimate phenotype can be comprised of hundreds to thousands of genes (Kliebenstein 2009a). OnceeQTL analysis is used to associate gene expression measurements with polymorphisms in a segregating population (Farrall 2004; Gilad et al. 2008). eQTLs are classified as cis- or trans-acting by comparing their genomic location to the position of the target genes. A cis-regulatory change is inferred when the eQTL coincides with the target gene, whereas a trans-regulatory change is assumed when the eQTL is not near the target gene (Wittkopp 2005). Regions of the genome that associate with the expression of many genes are referred to as eQTL hotspots (Chesler et al. 2005; Hansen et al. 2008), which tend to act in trans, regulating 1000 genes in some cases (West et al. 2007; Hansen et al. 2008; Li et al. 2013). Thus, eQTL hotspots can be used to characterize chromosomal positions of major regulators (de Koning and Haley 2005), including, for example, those involved in disease resistance (Mozhui et al. 2008; Kliebenstein 2009b). Alleles at the Mla locus were completely randomized within each of the five trays. Seedlings were grown in a 20°C controlled glasshouse to 10 cm prior to the application of treatment. Seedlings were inoculated with fresh Bgh conidiospores and trays were transferred to a growth chamber at 18°C (16 hr light and 8 hr darkness) where the trays were stored side-by-side in randomly assigned positions. Six days after inoculation, all leaves were photographed and scored for infection type (Table S2). The entire procedure was repeated three times in successive weeks.

Expression QTL mapping

eQTL analysis was carried out separately for the 16 and 32 HAI time points. Each of 22,840 probe sets was scanned for association with each of 377 nonredundant markers (Moscou et al. 2011b) using data from the 75 QSM lines. For probe set $i = 1, \ldots, 22,840$ and marker $j = 1, \ldots, 377$, a simple linear regression model was fitted to 75 data points with robust multi-array average (RMA)-normalized expression level as the response and an indicator of marker genotype as the explanatory variable. For missing genotypes, observed flanking marker genotypes were used with linear interpolation to impute a continuous genotype value in the interval [0,1]. A t-statistic ($t_i$) for testing the significance of the slope coefficient in the simple linear regression model was computed to quantify the strength of association between probe set $i$ and marker $j$. For each probe set $i$, $T_i$ was the maximum value of $|t_i|$ over markers $j = 1, \ldots, 377$ was computed, and the identity of the marker yielding the maximum value was recorded.

To identify statistically significant associations of expression level with marker genotype, the statistics $T_1, \ldots, T_{22,840}$ were first converted to p-values ($p_1, \ldots, p_{22,840}$) using a standard permutation approach that accounts for multiple testing across the 377 markers (Churchill and Doerge 1994). Specifically, the values of $T_1, \ldots, T_{22,840}$ were recomputed

MATERIALS AND METHODS

Plant growth, RNA extraction, and GeneChip hybridization

The 75 Q21861 × SM89010 (QSM) doubled haploid barley lines (Steffenson et al. 1995) were derived from a single F1 plant, and have been maintained via single seed descent (Moscou et al. 2011b). Seedlings were grown in two 98-cone-tray experiments in a climate-controlled greenhouse using a randomized block design. Each tray contained five seedlings/cone of the 75 QSM lines plus four replicates of each parent. Seven days after sowing, both trays were inoculated with Bgh isolate 5874 (AVRla, AVRao, visla, visao, and AVRwl) at an average density of 200 conidiospores per cm². The five seedlings were harvested, pooled, and placed in liquid nitrogen for each progeny and parent line at 16 and 32 hr after inoculation (HAI). Total RNA was isolated using a hot (60°C) phenol/guanidine thiocyanate method (Caldol et al. 2004). Labeling, hybridization, washing, and scanning were performed according to standard Affymetrix protocols at the Iowa State University GeneChip Core facility using 22K barley1 GeneChip expression arrays (Affymetrix part number 900515) (Close et al. 2004).

Phenotyping

Bgh isolate 5874 was propagated on Hordeum vulgare cv. Morex at 18°C while in 16 hr of light and 8 hr of darkness. H. laevigatum-derived lines were obtained from the USDA-Germplasm Resources Information Network (GRIN). Alleles at the Mla locus were amplified via polymerase chain reaction (PCR) using primers Mla LRR F and R (Supplemental Material, Table S1), which amplified the LRR region, spanning ~850 nucleotides between positions 1897 and 2725 bp relative to the Mla6 genomic sequence (AJ302293.1). Sequence verification was done by alignment against known Mla alleles (Seeholzer et al. 2010), and the alleles in Q21861 and SM89010 were assigned as Mla1 (AY009939.1) and Mla8 (GU245940.1) respectively.

To phenotype the QSM population, five 98-cone trays of Q21861, SM89010, and 75 QSM progeny were planted in sterilized potting soil. Each of the five experimental trays (7 rows × 14 columns) consisted of five seedlings for each parent (Q21861 and SM89010) and one seedling for each QSM progeny. One seedling was planted per cone and genotypes were completely randomized within each of the five trays. Seedlings were grown in a 20°C controlled glasshouse to 10 cm prior to the application of treatment. Seedlings were inoculated with fresh Bgh conidiospores and trays were transferred to a growth chamber at 18°C (16 hr light and 8 hr darkness) where the trays were stored side-by-side in randomly assigned positions. Six days after inoculation, all leaves were photographed and scored for infection type (Table S2). The entire procedure was repeated three times in successive weeks.

Expression QTL mapping

eQTL analysis was carried out separately for the 16 and 32 HAI time points. Each of 22,840 probe sets was scanned for association with each of 377 nonredundant markers (Moscou et al. 2011b) using data from the 75 QSM lines. For probe set $i = 1, \ldots, 22,840$ and marker $j = 1, \ldots, 377$, a simple linear regression model was fitted to 75 data points with robust multi-array average (RMA)-normalized expression level as the response and an indicator of marker genotype as the explanatory variable. For missing genotypes, observed flanking marker genotypes were used with linear interpolation to impute a continuous genotype value in the interval [0,1]. A t-statistic ($t_i$) for testing the significance of the slope coefficient in the simple linear regression model was computed to quantify the strength of association between probe set $i$ and marker $j$. For each probe set $i$, $T_i$ was the maximum value of $|t_i|$ over markers $j = 1, \ldots, 377$ was computed, and the identity of the marker yielding the maximum value was recorded.

To identify statistically significant associations of expression level with marker genotype, the statistics $T_1, \ldots, T_{22,840}$ were first converted to p-values ($p_1, \ldots, p_{22,840}$) using a standard permutation approach that accounts for multiple testing across the 377 markers (Churchill and Doerge 1994). Specifically, the values of $T_1, \ldots, T_{22,840}$ were recomputed
for each value of 100 permutations, where for each permutation, the 75 expression vectors associated with the 75 QSM lines were randomly assigned to the 75 genotype vectors associated with the 75 QSM lines. This yielded a null distribution of 22,840 × 100 permutation test statistics. The p-value for the original statistic T was given by the proportion of the 22,840 × 100 permutation test statistics ≥ T. To account for multiple testing across 22,840 probe sets, the 22,840 permutation p-values were converted to q-values (q1, … , q22,840) (Benjamini and Hochberg 1995). To control false discovery rate (FDR) at level 0.001, the association between probe set i and its most strongly associated marker was declared significant if and only if its q-value (q) was ≤ 0.001.

For this experiment, we interrogated the data set for only the most associated markers to identify the primary eQTL that had the greatest impact on gene expression. While other secondary and tertiary eQTL can also exert influence, we focused on changes in the most associated regulatory eQTL, with the hypothesis that it represents the most significant modification in the underlying biology of the gene in question.

**Differential expression analyses for QSM progeny groups**

The 75 QSM progeny were grouped by the four QSM progeny groups Mla1, MlLa, Mla2, and MlLa—analyzed using R-package limma (Ritchie et al. 2015). A model was fitted with expression value as response and treatment (genotype × time point) as explanatory variables. p-values were adjusted for multiple testing using R-package q-value (Storey et al. 2003). Q-values were estimated using the smoother method described in the BLAST+ command line suite (Camacho et al. 2009). This alignment gave an average of 86% coverage and 60% identity among the orthologous sequence pairs.

Next, protein sequences for *Arabidopsis* assembly Arabo11 were obtained from TAIR (www.arabidopsis.org) and compared against the Harvest 21 assembly that was used to derive barley1 GeneChip probe sets (Close et al. 2004) using blastx in the BLAST+ command line suite (Camacho et al. 2009). Coverage was calculated using the query as well as the subject and the maximum of the two was used. Sequences that aligned with 86% coverage and 60% identity were selected as candidates for homologs.

**Computational prediction of secretion-associated genes/proteins**

To reliably map the *Arabidopsis* proteome to the barley1 GeneChip probe sets, we obtained empirical scores for coverage and identity. Rice (*Oryza sativa* subsp. *japonica*) was selected as the monocot representative as it had the highest number of protein sequences (3716) in Swiss-Prot (UniProt Consortium 2015) as compared to barley (350 sequences).

**Transcript pattern analyses of published data sets**

We conducted linear model analyses of the normalized signal intensities for each of the 22,840 barley1 probe sets (Close et al. 2004) using packages in BioConductor/R (Gentleman et al. 2004; R Core Team 2016). RNA normalization and data transformation was done using package affy (Gautier et al. 2004). Linear model analysis and multiple testing adjustments were conducted using package limma (Ritchie et al. 2015) and q-value (Storey et al. 2015). The model used RMA normalized expression values as the response, and treatment (genotype × time point) as the input variable. Contrasts were used to compare transcript levels between treatments after infection with *Bgh* 5874. Q-values were estimated using the smoother method described in Storey and Tibshirani (2003).

Custom R program scripts were used to parse for highly similar patterns of expression among incompatible and compatible interactions up to 16 HAI, followed by significant (q-value ≤ 0.01) divergence from 16 to 32 HAI. This was used to distinguish two opposing scenarios: (i) downregulation of the transcript in the compatible interaction after 16 hr, as compared to paired time points in the incompatible interaction (designated "iPacMan"), or (ii) upregulation of the transcript in the compatible interaction after 16 hr, as compared to paired time points in the incompatible interaction (designated "PacMan").

**MlLa marker design and testing**

The two markers, MLOC_20004 and MLOC_25435 on Chr 2H, represent the outermost boundaries of a region flanked by recombination breakpoints and implicated to contain *MlLa*, at 621 and 624.5 Mb, respectively. Using the available barley Morex genome on Ensembl (Assembly 082214v1, INSDC Assembly GCA_000326085.1) (International Barley Genome Sequencing Consortium et al. 2012), 29 high-confidence candidates were selected. To further delineate the region that contains *MlLa*, markers were generated to identify genes within this region that cosegregated with the *MlLa* phenotype. Multiple primer sets were used to amplify candidate genes using standard *Taq* polymerase (Invitrogen, Carlsbad, CA) from cultivars SM89010 (*Mla8* and *MlLa*) and Q21861 (*Mla1* and *mlLa*). Any primer sets that showed potential as PCR-based markers (presence/absence or band-shift) were utilized on the QSM DH population (Table S3) and *H. laevigatum*-derived lines (Table S4). Those that showed no discernible difference between the two parents were sequenced to design cleaved amplified polymorphic sequence markers.

**Connecting alternatively regulated genes to biological function**

Of the 344 membrane trafficking encoding genes associated with 2HL at 16 HAI and/or 1HS at 32 HAI (Table S5), we focused on the 37 barley1 probe sets that exhibit alternate regulation between the two developmental stages. *Arabidopsis* proteins corresponding to these probe sets were interrogated for previously established protein–protein interactions in BioGRID (Stark et al. 2006). This resulted in a network of 129 nodes partitioned into 19 communities developed using the igraph package (Csardi and Nepusz 2006) in R (R Core Team 2016). These communities had a high modularity of 0.842, which reflects dense connections within communities and sparse connections across communities.

**Data availability**

All MIAME-compliant GeneChip profiling data from the QSM eQTL experiment are available as accession BB96 at the PLEXdb expression resource for plants and plant pathogens, as well as accession GSE68963 at NCBI-GEO. The previously published PLEXdb data set, BB4, is also available as NCBI-GEO accession GSE33396.

Table S1 details the PCR primers used in this study. Table S2 shows infection type data for three replications. Table S3 contains the complete QSM genetic map. Table S4 details the genotyping and
Figure 1 illustrates that Q21861 confers an IT of 0 in response to Bgh isolate 5874 and photographed 6 d later. The three leaves shown represent an average infected plant for each genotypic group. An average infection phenotype of <2.5 across three replications was considered resistant (incompatible interaction), whereas a value of ≥2.5 was considered susceptible (compatible interaction). The values are rounded to the nearest 0.5. QSM, Q21861 × SM89010.

| Bgh Isolate | Q21861 | SM89010 | QSM Progeny | QSM Progeny | QSM Progeny | QSM Progeny |
|-------------|--------|---------|-------------|-------------|-------------|-------------|
| Mla1        | mla1   | Mla1   | mla1        | Mla1        | mla1        | mla1        |
| MlLa        | MlLa   | MlLa   | MlLa        | MlLa        | MlLa        | MlLa        |

| Average Infection Type | 0 | 2 | 0 | 0 | 2 | 3 |
|------------------------|---|---|---|---|---|---|
| Number of Progeny Lines| – | – | 20| 18| 14| 23|

Table S5 features genes associated with membrane trafficking. Table S6 lists genes associated with each marker on the QSM map. Table S7 provides differential expression results for comparison between CI 16151 and CI 16155 (BB4), and between QSM progeny groups (Mla1 and mla1) and (mla1 and mla1). Table S8 shows the modes of alternate regulation for the genes transferred between 2HL at 16 HAI and 1HS at 32 HAI. Figure S1 shows a subnetwork connected via PEN1 (SYP121) in community A and VAMP721, SNAP33, and KEULE in community B. Figure S2 illustrates a community with the alternatively regulated AP2M as the hub that interacts with the remaining five proteins, including AP2A1.

**RESULTS**

**Mla1 is epistatic to MlLa**

In order to determine if interactions between Mla and MlLa had an effect on infection type (IT), the 75 QSM barley lines (Steffenson et al. 1995) were inoculated with Bgh isolate 5874. This isolate harbors both the AVR<i>a</i> and AVR<i>la</i> AVR effectors, and thus, was used to monitor the various infection types segregating in the QSM progeny. Figure 1 illustrates that Q21861 confers an IT of 0 in response to Bgh 5874, whereas SM89010 imparts an average IT of 2 with the same isolate. A 1259 cM transcript-derived marker map of the QSM population (Moscou et al. 2011b) was used to position Mla1 and MlLa. This map is divided into 377 bins across seven Chr and harbors 1494 Barley1 GeneChip-derived markers (Table S3).

Infection phenotyping (Wei et al. 1999) positioned a Bgh resistance specificity on the Mla locus on Chr 1HS (1H.I05), and another on the distal end of Chr 2HL. The second resistance that mapped to 2H1.67 colocalized with the position associated with MlLa (Schweizer and Stein 2011). ITs for three replications of each QSM recombinant line are detailed in Table S2.

MlLa is derived from the barley landrace <i>H. laevigatum</i>, but its precise position and sequence are currently unknown. Therefore, to determine whether we were indeed tracking the MlLa locus, we compared infection types and MlLa flanking marker sequences of SM89010 (the QSM parent line putatively harboring MlLa) to <i>H. laevigatum</i> and 10 other <i>H. laevigatum</i>-derived accessions (Dros 1957). SM89010 displayed identical infection type and DNA marker sequences to <i>H. laevigatum</i>, as well as the <i>H. laevigatum</i>-derived Minerva and Vada. This suggests that the locus mapped is likely MlLa (Table S4). Mla1-derived sequences were amplified and aligned against known Mla alleles (Seelholzer et al. 2010), confirming that Q21861 possessed Mla1, which was consistent with the observed IT of 0 associated with this allele.

Taking epistatic interactions into account, it was conceivable that some of the recombinant progeny in the QSM population would display an IT of 0 in response to Mla1, but still contain MlLa. Indeed, all 38 lines that contained Mla1 displayed an IT of 0, whereas the 14 lines that contained MlLa only displayed an average IT of 2, and lines that contained neither Mla1 nor MlLa exhibited an average IT of 3. Genotypes for the double haploid progeny fit a ratio of 1:1:1:1 (χ² 1:1:1:1 = 2.28; p-value = 0.3216) and a phenotypic ratio of 2 (IT = 0):1 (IT = 2):1 (IT = 3) (χ² 2:1:1 = 2.17; p-value = 0.3373) as expected for an epistatic interaction.

**Trans eQTL hotspots are associated with MlLa during Bgh penetration and Mla duringhaustorial development**

The QSM population is ideal for identifying regulatory components that reprogram the defense transcriptome of barley in response to inoculation with Bgh. Parental and QSM progeny lines were inoculated with Bgh isolate 5874 and harvested at 16 and 32 HAI; time points that are associated with appressorial penetration and early formation of haustorial development, respectively (Caldo et al. 2004, 2006; Moscou et al. 2011a). Gene expression for each of the 166 experimental units was estimated using the 22K barley1 GeneChip (Close et al. 2004) (BB96 in PLEXdb and GSE68963 in NCBI-GEO).

A simple linear regression model was fit to 75 data points with normalized expression level as the response and an indicator of marker genotype as the explanatory variable to identify eQTLs. Significance thresholds were determined by using q-values to control the FDR (Benjamini and Hochberg 1995). Table S6 lists all 22,840 probe sets along with their most associated eQTL marker and its level of significance. The values are rounded to the nearest 0.5. QSM, Q21861 × SM89010.
were of highest significance to at least 100 probe sets each. For the same threshold, 371 positions were classified as the top association for at least one probe set at 32 HAI. Likewise, 6 of these 371 were found to be most associated with at least 100 probe sets. In simulations performed with a multinomial distribution, the probability for an eQTL marker position to be randomly associated with 100 probe sets by chance is essentially 0 with eQTL marker associations controlled at q-value < 0.001. Thus, for the purposes of this manuscript, only loci associated with >100 probe sets were considered eQTL hotspots.

The five eQTL hotspots significant at 16 HAI, corresponding to penetration of epidermal cells by Bgh, map to linked positions on Chr 2HL Bins 63 and 65–68 (MlLa region) and associate with a total of 3103 genes (Table S6). Likewise, the six hotspots significant at 32 HAI, corresponding to formation of Bgh haustorial feeding structures and delivery of secreted effectors into the host, localized to Chr 1HS Bins 3–8 (including Mla1) and associate with 5070 genes. In this report, 2HL will be the designation for Chr 2H Bins 63 and 65–68 and 1HS refers to Chr 1H Bins 3–8. Figure 2 illustrates off-diagonal probe sets, indicative of trans eQTL, including those associated with 2HL at 16 HAI and 1HS at 32 HAI.

We then wished to examine any changes within each of the four genotypic progeny groups and identify the alleles at 2HL and 1HS eQTL that suppress or upregulate associated genes. To do this, we compared the gene expression for each progeny line against the median of the genotypic progeny groups and identify the alleles at 2HL and 1HS eQTL. For the same purposes of this manuscript, only loci associated with >100 probe sets were considered eQTL hotspots.

Transcript pattern analysis associates eQTL hotspots on Chr 2HL and 1HS with compatibility or incompatibility

Next, we integrated the current eQTL results with previous barley1 GeneChip expression data to examine changes in transcript accumulation across developmental time. Accession BB4 (PLEXdb; GSE33396 in NCBI-GEO) facilitated comparison with independent incompatible and compatible interactions mediated by the Mla1 and Mla13 alleles, respectively, at six different time points (0, 8, 16, 20, 24, and 32 HAI) with Bgh isolate 5874 (AVRr13, AVRg13, and avr13) (Caldo et al. 2004).

Of particular interest were two classes of transcript accumulation; both were highly similar among incompatible and compatible interactions up to 16 HAI, coinciding with germination of Bgh conidiospores and formation of appressorium, followed by significant divergence from 16 to 32 HAI, during development of the perihaaustorial membrane between fungal haustoria and host epidermal cells (Caldo et al. 2004). Computational pattern analysis was used to distinguish the two opposing scenarios: The first represents genes downregulated during compatible interactions after 16 HAI, as compared to corresponding time points in incompatible interactions; this pattern was designated PacMan (Figure 4, A–C and Table 1). In contrast, the second represents genes upregulated from 16 to 32 HAI during compatible interactions, as compared to the equivalent time points in incompatible interactions; this pattern was designated iPacMan (Figure 4, D–F and Table 1). The first scenario may represent positive regulators of host immune signaling that are suppressed by the pathogen during compatible interactions, whereas the second may represent a situation where the pathogen is inducing host factors for disease susceptibility. Of the 3103 genes associated with the eQTL hotspot cluster on Chr 2HL, 321 and 258 exhibited PacMan and iPacMan expression patterns, respectively. Similarly, of the 5070 genes associated with 1HS, 404 are associated with a PacMan pattern, whereas 606 displayed an iPacMan pattern (Table 1 and Table S6).

The genotypes represented by PLEXdb accession BB4 lack MlLa, but do possess Mla1. Therefore, when inoculated with Bgh isolate 5874 (AVRr13, AVRg13, and avr13), a comparison of differential gene expression can be made between QSM progeny groups (Mla1 and mla1; resistant) and (mla1 and MlLa; susceptible) in BB96, and between CI 16151 (Mla1 and mla1; resistant) and CI 16155 (Mla13 and mLa; susceptible) from BB4. At 32 HAI, of the 1712 genes that are differentially expressed between CI 16151 and CI 16155, 1356 (80%) are also differentially expressed between (Mla1 and MlLa) and (mla1 and MlLa) in BB96. Of these 1356 genes, 559 are upregulated and 797 are downregulated in the Mla1 and mLa QSM progeny group (Table S7).

Temporal regulation of immunity

Up to now, a handful of genes (and their encoded proteins) have been functionally connected to NLR-mediated pathways (Shirasu 2009; Jacob
et al. 2013), but these interactions have not been associated with specific stages of pathogen development. Previous studies have shown that MLA1 interacts with SGT1 (suppressor of the G2 allele of skp1) (Bieri et al. 2004), which binds SKP1, a component of the Skp1-Cullin-F-box ubiquitin ligase complex, and facilitates regulation of defense in both yeast and plants (Kitagawa et al. 1999; Azevedo et al.

Figure 3  Number of genes up- and downregulated in each QSM progeny for 16 and 32 HAI as compared to the gene-wise median for the Q and SM parent lines. At 16 HAI, (A) all genes, (B) genes associated with 2HL, and (C) genes associated with 2H.67 (Mla1) are plotted for each QSM progeny. At 32 HAI, (D) all genes, (E) genes associated with 1HS, and (F) genes associated with 1H.05 (Mla1) are represented. The purple, red, blue, and brown lines refer to QSM progeny genotype (Mla1 and Mla1), (Mla1 and mla1), (mla1 and Mla1), and (mla1 and mla1), respectively. Within each group, the barley lines are in numerical order (Table S2). Note that the graphs are plotted on different scales. HAI, hr after inoculation; QSM, Q21861 × SM89010.
Along with RAR1 (required for Mla12 resistance) and HSP90 (Heat Shock Protein 90), SGT1 positively controls the steady state levels of resistance proteins recognizing viral, bacterial, oomycete, or fungal pathogens (Azevedo et al. 2002; Takahashi et al. 2003; Schulze-Lefert 2004; Shirasu 2009). Our findings show that SGT1 is associated with 2H.67 (MlLa locus) at 16 HAI, whereas HSP90 is associated with 1H.05 (Mla1) at 32 HAI. However, RAR1 does not associate significantly with either 2HL or 1HS, or any other eQTL marker.

Formerly, HvWRKY10, HvWRKY19, and HvWRKY28 were shown to positively regulate Mla-triggered immunity to Bgh; however, current evidence suggests that these three nuclear-localized WRKY transcription factors do not directly bind MLA (Meng and Wise 2012). Of these, we show that WRKY19 is regulated (either directly or indirectly) by 2H.67 (MlLa) at 16 HAI, whereas WRKY10 is associated with 1H.05 (Mla1) at 32 HAI. Additional WRKYs associated with 2HL at 16 HAI include WRKY20, WRKY33, WRKY34, WRKY41, WRKY46, and WRKY61. From this group, WRKY33, WRKY41, and WRKY46 change association from 2H.65, 2H.66, and 2H.67 (MlLa) at 16 HAI, respectively, to 1H.05 (Mla1) at 32 HAI (Table S6).

Lastly, the previously known defense-related proteins, syntaxin (VAMP727), thaumatin, and SOD1 display no significant associations at 16 HAI, but were associated with 1H.08, 1H.05 (Mla1), and 1H.06 at 32 HAI, respectively. BLN1, a negative regulator of basal defense (Meng et al. 2009; Xu et al. 2015), was associated with 2H.68 during penetration, but 1H.07 during haustorial growth, implying control by unlinked host loci during pathogen development.

To further extend the temporal analysis of the known NLR-associated proteins above, we also queried the 2HL and 1HS associations for encoded proteins annotated to membrane trafficking, including the secretory pathway, nuclear and vesicle transport, signaling G-proteins, and their known interactors (Stark et al. 2006; Paul et al. 2014; Rutter and Innes 2017). This identified 356 distinctive probe sets encoding several SNAREs, Rab GTPases (part of the Ras superfamily of small GTPases), tethering factors, COat Protein complex (COPs), and Clathrin-Coated Vesicle proteins (CCVs). Of these 356 probe sets, 80 were associated with 2HL at 16 HAI, 126 associated with 1HS at 32 HAI, and 38 were associated with both 2HL at 16 HAI and then 1HS at 32 HAI (Table 1 and Table S6). Taken together, the current eQTL analysis
Table 1 Number of genes associated with each hotspot in the Chr 2HL and 1HS clusters and the number represented by PacMan and iPacMan expression patterns, respectively

| Position (Chr.Bin) | Associations | Expression Pattern | PacMan | iPacMan |
|--------------------|--------------|--------------------|--------|---------|
| 2H.63              | 217          | 17                 | 10     |         |
| 2H.64              | 26           | 0                  | 0      |         |
| 2H.65              | 261          | 33                 | 24     |         |
| 2H.66              | 1331         | 163                | 167    |         |
| 2H.67 (M1la)       | 961          | 100                | 26     |         |
| 2H.68              | 333          | 8                  | 31     |         |
| 2HL total          | 3103         | 321                | 258    |         |
| 1H.03              | 261          | 13                 | 12     |         |
| 1H.04              | 276          | 14                 | 6      |         |
| 1H.05 (Mla1)       | 3296         | 341                | 431    |         |
| 1H.06              | 783          | 21                 | 121    |         |
| 1H.07              | 114          | 7                  | 13     |         |
| 1H.08              | 340          | 8                  | 23     |         |
| 1HS total          | 5070         | 404                | 606    |         |

Chr, chromosome; PacMan, genes downregulated during compatible interactions after 16 HAI, as compared to corresponding time points in incompatible interactions; iPacMan, genes upregulated from 16 to 32 HAI during compatible interactions, as compared to the equivalent time points in incompatible interactions.

a Associated genes are represented at 16 HAI for Chr 2H and 32 HAI for Chr 1H hotspots at a q-value ≤ 0.001. The annotations for these genes are presented in Table S6. Expression pattern changes across developmental time in the comparison of incompatible (Mla1-AVRr) and compatible (Mla13-Avr1) barley-Bgh interactions (Accession BB4 at PLEXdb.org). Described further in Pattern Analyses of Published Data Sets under Materials and Methods.

b Marker bin not included in analyses as the number of associations is below 100 but shown here for informational purposes.

identifies proteins coupled to NLR-mediated signaling, but also provides new associations that appear to play important temporal roles during penetration, haustorial growth, or both.

Alternating temporal control: prioritizing immune regulation from 2HL (associated with penetration) to 1HS (associated with haustorial development)

Plant transcriptomes undergo significant reprogramming in response to environmental stimuli, such as cold, salinity, drought, or pathogen stress. A marked example of this reprogramming is the transfer of association with Chr 2HL (Bins 63, 65–68) at 16 HAI, and then to Chr 1HS (Bins 3–8) at 32 HAI, where a total of 1470 genes come under alternate regulation postpenetration (Figure 5). Of these 1470 genes, 260 exhibited PacMan patterns of transcript accumulation, whereas 192 showed patterns similar to iPacMan.

Alternate transcriptional control suggests that postpenetration regulators at 1HS supplant those at 2HL. There are four modes of regulation between 2HL at 16 HAI (penetration) and 1HS at 32 HAI (haustoria), as shown in Table 2 and Table S8. We designated these SM2HL,16 → SM1HS,32, SM2HL,16 → SM1HS,32, Q2HL,16 → Q1HS,32, and Q2HL,16 → SM1HS,32. Represented by the quadrants delineated at point (0,0) in Figure 6. Genes depicted in the upper-left-hand quadrant (SM2HL,16 → Q1HS,32) display higher expression in recombinant lines where the SM allele is present on 2HL at 16 HAI, and in lines possessing the Q allele at 32 HAI. This scenario also represents the presence of an active M1la during penetration, in addition to a functioning M1la during haustorial development. Thirty-one percent of the genes in this quadrant are associated with PacMan patterns of expression (vs. 0% iPacMan), and include, among other annotations, BLN1, Rab GTPase, and Syntaxin.

Genes depicted in the lower-left-hand quadrant (SM2HL,16 → SM1HS,32) display higher expression at 16 HAI in recombinant lines where the SM allele is present on 2HL, and at 32 HAI in lines possessing the SM allele on 1HS. This scenario is indicative of the presence of a functional M1la, but is accompanied by the absence of M1la. Genes depicted in the upper-right-hand quadrant (Q2HL,16 → Q1HS,32) display higher expression at 16 HAI in recombinant lines where the Q allele is present on 2HL, and at 32 HAI in lines possessing the Q allele on 1HS. This scenario is indicative of the absence of a functional M1la, but the presence of a functional M1la. Genes in these two quadrants do not have a distinct expression pattern or associated pathway.

Lastly, genes depicted in the lower-right-hand quadrant (Q2HL,16 → SM1HS,32) display higher expression at 16 HAI in recombinant lines where the Q allele is present on 2HL, and at 32 HAI in lines possessing the SM allele on 1HS. Similar to the second quadrant, this scenario is indicative of the absence of a functional M1la, but also the absence of M1la. Thirty-two percent of these genes are associated with transcript accumulation designated as iPacMan (vs. 0% PacMan). Among other annotations, these genes include nuclear transport factor 2, aquaporin PIP, and a calcium-binding EF hand family protein. Of the 1470 genes that show alternate regulation between 2HL at 16 HAI and 1HS at 32 HAI, 1420 switch the allele responsible for higher expression (SM2HL,16 to Q1HS,32 or Q2HL,16 to SM1HS,32) when association is reallocated from 2HL to 1HS.

DISCUSSION

We used eQTL analysis of the QSM doubled haploid barley population after infection with the fungal Ascomycete pathogen, B. graminis f. sp. hordei, to interrogate temporal control of plant immunity. Near-synchronous penetration by thousands of Bgh conidiospores delivers an ideal stimulus to induce dynamic reprogramming of the leaf transcriptome to response to pathogen attack. Two major clusters of trans eQTL hotspots on Chr 2HL and 1HS were identified along with immune response genes associated with them, corresponding to appressorial penetration and development of haustoria, respectively.

Dynamic regulation of immunity

Trans-regulatory regions typically influence transcript levels via a protein intermediate and have the potential to interact with many proteins, depending on the milieu present in the cell in a given condition. A single variant may be able to activate transcription in one condition and repress it in another, resulting in a change in direction of the effect, as observed when the two carbon sources, glucose and ethanol, are compared during yeast fermentation and aerobic respiration, respectively (Smith and Kruglyak 2008). In the current eQTL study, 1470 genes associated with 2HL during penetration were repurposed by 1HS at later stages of infection, but more specifically, 299 genes reallocated control from the M1la locus (2H.67) to 1H.05 (M1la1). Finally, an additional 138 genes were identified that change association from other non-Chr 2HL positions at 16 HAI to 1H.05 (M1la1) at 32 HAI. Of the total 1608 (1470 + 138) genes that undergo alternate regulation, 271 exhibited PacMan patterns of expression, which is indicative of host immune signaling that is typically suppressed by pathogen effectors (Caldo et al. 2004). Another 197 genes displayed a contrasting pattern of transcript accumulation designated iPacMan, which we postulate represent a scenario where the pathogen induces defense, possibly by recognition of PAMPs by pattern recognition receptors, but then coopts these genes for its own purposes (Figure 4). While we focused on transcript accumulation signified by PacMan and iPacMan patterns, undoubtedly there are additional changes influenced post-transcriptionally or post-translationally, for example by small RNAs (Liu et al. 2014;
Xu et al. 2014; Fei et al. 2016; Zhang et al. 2016). Nonetheless, the population-based eQTL analysis reported here supports the conclusion that these genes are part of an immune regulon that can be activated or repressed by disparate resistance loci in response to multiple infection stages.

Alternate control of gene expression suggests that one or more regulators on 1HS supersede that exerted by eQTLs on 2HL. There are four modes of alternate transcriptional control, specifically SM2HL,16–Q1HS,32, SM2HL,16–SM1HS,32, Q2HL,16–Q1HS,32, and Q2HL,16–SM1HS,32 (Table 2). For example, SM2HL,16 denotes that at 16 HAI recombinant lines with the SM allele at 2HL (including MlLa at 2H.67) had higher gene expression compared to the lines with the Q allele at 2HL. Similarly, Q1HS,32 implies that at 32 HAI the lines with the Q allele at 1HS (including Mla at 1H.05) had higher gene expression compared to the lines with the SM allele at 1HS. As illustrated in Figure 6A, allelic effects were interchanged (Q2HL,16 to SM1HS,32 or SM2HL,16 to Q1HS,32) for 1420 of the 1470 genes that reallocated control from 2HL at penetration to 1HS during haustorial development.

One explanation for these correlated phenomena may be multiple linked polymorphisms. Trans-regulatory regions are much larger targets for variation than cis-regulatory regions (Brem et al. 2002; Smith and Kruglyak 2008). Multiple mutations could accumulate at loci, and, depending on the condition, could compensate differentially. The mean phenotype would be stable over conditions, yet the direction of the effect within a condition could vary across the population, as observed in the phenotypic range for lines lacking a functional Mla1. Another example, which is observed across taxa rather than the population, is suggested by the hotspots on Chr 1HS, including those in fl influenced by Mla1 (1H.05). In this case, the emerging narrative is consistent with multiple polymorphisms that influence disease resistance to diverse pathogens (Seeholzer et al. 2010; Periyannan et al. 2013; Mago et al. 2015; Cesari et al. 2016).

### Table 2 Modes of alternate regulation between 2HL at penetration and 1HS in haustoria and associated expression patterns

| Modes of Alternate Regulation | Number of Genes | Expression Pattern |
|-------------------------------|-----------------|--------------------|
| SM2HL,16–Q1HS,32              | 836             | PacMan 260         |
| SM2HL,16–SM1HS,32             | 12              | iPacMan 0          |
| Q2HL,16–Q1HS,32               | 38              | PacMan 0           |
| Q2HL,16–SM1HS,32              | 584             | iPacMan 191        |

PacMan, genes downregulated during compatible interactions after 16 HAI, as compared to corresponding time points in incompatible interactions; iPacMan, genes upregulated from 16 to 32 HAI during compatible interactions, as compared to the equivalent time points in incompatible interactions.

---

**Membrane trafficking regulated by immune-associated loci**

To further elucidate the underlying mechanisms behind these phenomena, we queried the 2HL and 1HS associations annotated to secretory pathway, vesicle transport, and signaling G-proteins. This identified 356 probe sets including several SNAREs, Rab GTPases (part of the Ras superfamily of small GTPases), tethering factors, COPs, and CCVs (Table S5). Rab GTPases regulate vesicle formation, vesicle movement, and membrane fusion (Stenmark and Olkkonen 2001), while SNARE proteins mediate fusion of vesicles with their target membrane-bound compartments (Burri and Lithgow 2004). These processes make up the route through which cell surface proteins are trafficked from the Golgi to the plasma membrane and are recycled. In plants, membrane vesicles targeted to the cell division plane fuse with one another to form the partitioning membrane, progressing from the center to the periphery of the cell. This is seen in the two types of SNARE complexes formed by KNOLLE that are jointly needed to mediate membrane fusion in...
cytokinesis. One of these complexes, a KNOLLE–SNAP33–VAMP721/722 trimer (El Kasmi et al. 2013) (Figure S1), is localized on the plasma membrane and drives membrane fusion of vesicles destined for exocytosis (Collins et al. 2003). PEN1 is engaged in a similar SNARE complex mediating extracellular immunity via exocytosis also with SNAP33 and VAMP721/722 (Kwon et al. 2008) (Figure S1). Intriguingly, the alternate regulation exhibited by VAMP721/722, first by the MlLa locus at penetration and then by Mla1 during haustorial growth, could provide clues as to how its secretion apparatus is coopted for immune responses to pathogens.

Tethering factors have emerged as key regulators of membrane traffic and organelar architecture (Dubuke and Munson 2016). The restricted subcellular localization of tethering factors and their ability to interact with RABs and SNAREs suggests that tethers participate in

Figure 6 Summary of allelic effects between Chr 2HL and Mla1 (1H.05) show modes of alternate regulation. (A) Representation of the allelic effects between 2HL at 16 HAI and 1HS at 32 HAI. The allelic effects between 1H.05 (Mla1) and (B) 2H.63, (C) 2H.65, (D) 2H.66, (E) 2H.67 (MlLa), and (F) 2H.68, are shown in respective panels. The quadrants delineated by (0,0) in each panel illustrate the four modes of regulatory transfer between hotspots and are labeled along with their number of associated genes. Chr, chromosome; HAI, hr after inoculation. Panel (G) summarizes the gene expression effects for each of the four quadrants.
determining the specificity of membrane fusion. An accepted model of tether function considers them molecular “bridges” that link opposing membranes before SNARE pairing. Tethers are also implicated to function as integration switches that simultaneously transmit information to coordinate distinct processes required for membrane traffic. Additionally, CCVs selectively sort cargo at the cell membrane, trans-Golgi network, and endosomal compartments, whereas COPs transport proteins between the Golgi complex and the rough endoplasmic reticulum. Adaptor protein complexes are vesicle coat components and appear to be involved in cargo selection and vesicle formation. AP-2 is involved in clathrin-dependent endocytosis in which cargo proteins are incorporated into vesicles surrounded by clathrin (CCVs) that are destined for fusion with the early endosome (Happel et al. 2004). The μ-adaptin of AP-2 (AP2M) localizes to the plasma membrane in plants for CCV formation, similar to AP-2 in animals. AP2M is involved in ETI mediated by plasma membrane-localized disease resistance proteins, possibly by mediating endocytosis of immune receptor components from the plasma membrane (Hatsugai et al. 2016) (Figure S2). However, it is not the coat proteins that determine the target of a transport vesicle but the SNAREs (Sanderfoot and Raikhel 1999). Together, the coat proteins and SNAREs coordinate the trafficking of cargo between various organelles of the endomembrane system.

Membrane trafficking pathways are involved in immune receptor activation, signal transduction, and execution of multiple defense responses including programmed cell death (Teh and Hofius 2014). Although mechanisms of pathogenic modulation of endocytic, secretory, and vacuolar trafficking and their roles in plant–microbe interactions need additional investigation, these pathways appear to function in the rapid responses to environmental stimuli (Inada and Ueda 2014).

**Shared components in immunity**

In previous studies, we localized a trans eQTL hotspot to 2H.16 that associates with an enhancer for adult plant resistance to the obligate Basidiomycete fungus *Puccinia graminis tritici* race TTKSK, more commonly known as Ug99 stem rust (Moscou et al. 2011b). This hotspot, unlinked to the *Bgh*-responsive eQTL described here, overtook regulatory control of 368 genes from several unlinked loci when plants are challenged with Ug99 (Moscou et al. 2011b). Of these 368 genes, 97 are associated with the current *Bgh*-induced 2HL at 16 HAI, 139 with 1HS at 32 HAI, and 57 undergo alternate regulation between 2HL and 1HS (Table S6), indicating substantial conservation of defense components to these two obligate biotrophic fungi.

The *Bgh*-susceptible *Arabidopsis pen2 pad4 sag101* triple mutant transformed with *Mla1* recognizes *Mla1*-incompatible isolates of *Bgh* (Maekawa et al. 2012). In order to compare MLA1-dependent signaling in *Arabidopsis* and barley, the top 100 differentially expressed genes from Maekawa et al. (2012) were mapped onto Barley1 probe sets; 14 of these returned high-confidence alignments (Table S6). Seven of these 14 were associated with 2HL at 16 HAI, six with 1HS at 32 HAI, and four associated with alternate regulation, first by 2HL (including *Mla1*) at penetration, and then 1HS (including *Mla1*) during development of haustoria.

Only one MLA1-dependent protein from *Arabidopsis* also aligned with a probe set associated with both *Mla1* and *Mla1* in barley. This was annotated as a putative serine/threonine-protein kinase (PBL2). PBL2 contributes to PTI signaling and defense responses downstream of LRR receptor-like serine/threonine-protein kinase (FLS2) (Zhang et al. 2010). Ligand-activated FLS2 is internalized in a clathrin-dependent manner that converges at ARA7 (Rab GTPase Homolog F2B) endosomes facilitating the responses required for full plant immunity (Mbengue et al. 2016).

**Future prospects**

Plant immune systems exemplify multi-tiered signaling networks comprised of biological molecules interacting in space and time. For example, we show that of the HSP90-RAR1-SGT1 chaperone complex, SGT1 is associated with 2H.67 (*Mla1* locus) at 16 HAI, whereas HSP90 is associated with 1H.05 (*Mla1*) at 32 HAI. We also show that the transcript encoding BLN1, a negative regulator of basal defense, is associated with 2H.68 at penetration, but 1H.07 during haustorial development, inferring specific control by unlinked chromosomal regions in response to different phases of pathogen infection. The regulatory mechanisms of the many examples in this report need further study, and questions also remain whether there is an epigenetic component to these phenomena. It would be intriguing to quantify methylation sites across the QSM population at different infection stages and integrate that with the current eQTL data.

One general implication of our results is that many genetic effects on most traits are likely to be detected without testing for gene–environment interactions, provided that the relevant environmental factors are known and controlled either experimentally or statistically. However, analyses that ignore temporal gene-by-environment interactions introduce strong biases regarding the types of loci that are detected. Our focus on transcript levels as quantitative traits allowed us to study a very large number of traits simultaneously and to delineate general patterns, as well as to provide detailed examples of loci that show gene-by-environment interactions, specifically infection of a plant host by an obligate biotrophic pathogen. The quantitative details would undoubtedly differ if different species, environments, and phenotypes were studied. However, some environmental differences (for example, exposure to pathogens) can have a dramatic effect on health. Our detailed studies in a model grain crop provide examples of the types of effects that may be expected in host–parasite interactions, and thereby inform practical study design.

**ACKNOWLEDGMENTS**

We thank Matt Moscou for initial phenotyping of the QSM (Q21861 × SM89010) population and for development of the QSM genetic map (Moscou et al. 2011b). We are grateful to Divya Mistry and Julie Dickerson for extensive discussions on eQTL gene coexpression networks. This article is a joint contribution of the Iowa Agriculture and Home Economics Experiment Station and the Corn Insects and Crop Genetics Research Unit, United States Department of Agriculture (USDA)-Agricultural Research Service. Mention of trade names or commercial products in this publication is solely for the purpose of providing scientific information and does not imply recommendation or endorsement by the USDA or the National Science Foundation. USDA is an equal opportunity provider and employer. Research supported in part by the National Science Foundation Plant Genome Research Program grants 09-22746 and 13-39348 to R.P.W. and D.N., and USDA-Agricultural Research Service project 3625-21000-060-00D to R.P.W. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Author contributions: R.P.W conceived and designed the experiments. G.F., R.P.W., and A.Y.E.C. performed the experiments. P.S., R.X.,
LITERATURE CITED

Azavedo, C., A. Sadanandom, K. Kitagawa, A. Freialdenhoven, K. Shirasu et al., 2002 The RAR1 interactor SGT1, an essential component of R gene-triggered disease resistance. Science 295: 2073–2076.

Benjamini, Y., and Y. Hochberg, 1995 Controlling the false positive rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. B 57: 289–300.

Bent, A. F., and D. Mackey, 2007 Elicitors, effectors, and R genes: the new paradigm and a lifetime supply of questions. Annu. Rev. Phytopathol. 45: 399–436.

Bieri, S., S. Mauch, Q. H. Shen, J. Peart, A. Devoto et al., 2004 RAR1 positively controls steady state levels of barley MLA resistance proteins and enables sufficient MLA6 accumulation for effective resistance. Plant Cell 16: 3480–3495.

Brem, R. B., G. Yvert, R. Clinton, and L. Kruglyak, 2002 Genetic dissection of transcriptional regulation in budding yeast. Science 296: 752–755.

Burris, L., and T. Lithgow, 2004 A complete set of SNAREs in yeast. Traffic 5: 45–52.

Caldo, R. A., D. Nettleton, and R. P. Wise, 2004 Interaction-dependent gene expression in Mla-specified response to barley powdery mildew. Plant Cell 16: 2514–2528.

Caldo, R. A., D. Nettleton, J. Peng, and R. P. Wise, 2006 Stage-specific suppression of basal defense discrimates barley plants containing fast- and delayed-acting Mla powdery mildew resistance alleles. Mol. Plant Microbe Interact. 19: 939–947.

Camacho, C., G. Coulouris, V. Avagyan, N. Ma, J. Papadopoulos et al., 2009 BLAST+: architecture and applications. BMC Bioinformatics 10: 421.

Cesari, S., M. Bernoux, P. Moncuquet, T. Kroj, and P. N. Dodds, 2014 A novel conserved mechanism for plant NLR protein pairs: the “integrated decay” hypothesis. Front. Plant Sci. 5: 606.

Cesari, S., J. Moore, C. Chen, D. Webb, S. Periyannan et al., 2016 Cytosolic activation of cell death and stem rust resistance by cereal MLA-family CC-NLR proteins. Proc. Natl. Acad. Sci. USA 113: 10204–10209.

Chang, C., D. Yu, J. Jiao, S. Jing, P. Schulze-Lefert et al., 2013 Barley MLA immune receptors directly interfere with antagonistically acting transcription factors to initiate disease resistance signaling. Plant Cell 25: 1158–1173.

Chen, X., C. A. Hackett, R. E. Níks, P. E. Hedley, C. Booth et al., 2010 An eQTL analysis of partial resistance to Puccinia hordei in barley. PLoS One 5: e8598.

Chesler, E. J., L. Lu, S. Shou, Y. Qu, J. Gu et al., 2005 Complex trait analysis of gene expression uncovers polygenic and pleiotropic networks that modulate nervous system function. Nat. Genet. 37: 233–242.

Churchill, G. A., and R. W. Doerge, 1994 Empirical threshold values for quantitative trait mapping. Genetics 138: 963–971.

Close, T. J., S. I. Wananmaker, R. A. Caldo, S. M. Turner, D. A. Ashlock et al., 2004 A new resource for cereal genomics: 22K barley GeneChip comes of age. Plant Physiol. 134: 960–968.

Collins, N. C., H. Thordal-Christensen, V. Lipka, S. Bau, E. Kombrink et al., 2003 SNARE-protein-mediated disease resistance at the plant cell wall. Nature 425: 973–977.

Csardi, G., and T. Nepusz, 2006 The igraph software package for complex network research. Inter. J. Comp. Sys. 1695: 1–9.

Cui, H., K. Tsuda, and J. E. Parker, 2015 Effector-triggered immunity: from pathogen perception to robust defense. Annu. Rev. Plant Biol. 66: 487–511.

de Koning, D. J., and C. S. Haley, 2005 Genetical genomics in humans and model organisms. Trends Genet. 21: 377–381.

Dros, J., 1957 The creation and maintenance of two spring barley varieties. Euphytica 6: 45–48.

Dubuque, M. L., and M. Munson, 2016 The secret life of tethers: the role of tethering factors in SNARE complex regulation. Front. Cell Dev. Biol. 4: 42.

El Kasmi, F., C. Krause, U. Hiller, Y. D. Steinhof, U. Mayer et al., 2013 SNARE complexes of different composition jointly mediate membrane fusion in Arabidopsis cytokinesis. Mol. Biol. Cell 24: 1593–1601.

Farrall, M., 2004 Quantitative genetic variation: a post-modern view. Hum. Mol. Genet. 13: R1–R7.

Fei, Q., Y. Zhang, R. Xia, and B. C. Meyers, 2016 Small RNAs add zing to the zig-zag-zig model of plant defenses. Mol. Plant Microbe Interact. 29: 165–169.

Gautier, L., L. Cope, B. Bolstad, and R. Irizarry, 2004 affy--analysis of Affymetrix GeneChips at the data probe level. Bioinformatics 20: 307–315.

Gentleman, R. V., C. Carey, D. Bates, B. Bolstad, M. Dettling et al., 2004 Bioconductor: open software development for computational biology and bioinformatics. Genome Biol. 5: R80.

Glad, Y., S. A. Rifkin, and J. K. Pritchard, 2008 Revealing the architecture of gene regulation: the promise of eQTL studies. Trends Genet. 24: 408–415.

Halterman, D. A., and R. P. Wise, 2004 A single-amino acid substitution in the sixth leucine-rich repeat of barley MLA6 and MLA13 alleviates dependence on RAR1 for disease resistance signaling. Plant J. 38: 215–226.

Hansen, B. G., B. A. Halkier, and D. J. Kliewenstein, 2014 Identifying the molecular basis of QTLs: eQTLs add a new dimension. Trends Plant Sci. 19: 72–77.

Happel, N., S. Honing, J. M. Neuhaus, N. Paris, D. G. Robinson et al., 2004 Arabidopsis mu A-adaptin interacts with the tyrosine motif of the vacuolar sorting receptor VSR-PS1. Plant J. 37: 678–693.

Hatsugi, N., R. Hillmer, S. Yamaoka, I. Hara-Nishimura, and F. Katagiri, 2016 The μ subunit of Arabidopsis adaptor protein-2 is involved in effector-triggered immunity mediated by membrane-localized resistance proteins. Mol. Plant Microbe Interact. 29: 345–351.

Inada, N., and T. Ueda, 2014 Membrane trafficking pathways and their roles in plant-microbe interactions. Plant Cell Physiol. 55: 672–686.

International Barley Genome Sequencing Consortium, Mayer, K. F., R. Waugh, J. W. Brown, A. Schulman, P. Langridge et al., 2012 A physical, genetic and functional sequence assembly of the barley genome. Nature 491: 711–716.

Inumaa, T., S. A. Khodaparast, and S. Takamatsu, 2007 Multilocus phylogenetic analyses within Blumeria graminis, a powdery mildew fungus of cereals. Mol. Phylogenet. Evol. 44: 741–751.

Jacob, F., S. Vernaldi, and T. Maekawa, 2013 Evolution and conservation of plant NLR functions. Front. Immunol. 4: 297.

Kitagawa, K., D. Skowrya, S. J. Elledge, J. W. Harper, and P. Hieter, 1999 SGT1 encodes an essential component of the yeast kinetochore assembly pathway and a novel subunit of the SCF ubiquitin ligase complex. Mol. Cell 4: 21–33.

Kliewenstein, D., 2009a Quantitative genomics: analyzing intraspecific variation using global gene expression polymorphisms or eQTLs. Plant Biol. 60: 93–114.

Kliewenstein, D. J., 2009b Quantification of variation in expression networks, pp. 227–245 in Plant Systems Biology, edited by Belostotsky, D., Humana Press, Totowa, NJ.

Kwon, C., C. Neu, S. Pajonk, H. S. Yun, U. Lipka et al., 2008 Co-option of a default secretary pathway for plant immune responses. Nature 451: 835–840.

Li, L., K. Petsy, R. Shimizu, S. Liu, W. W. Xu et al., 2013 Mendelian and non-Mendelian regulation of gene expression in maize. PLoS Genet. 9: e1003202.

Li, X., P. Kapos, and Y. Zhang, 2015 NLRs in plants.Curr. Opin. Immunol. 32: 114–121.

Liu, J., X. Cheng, D. Liu, W. Xu, R. Wise et al., 2014 The miR9863 family regulates distinct Mla alleles in barley to attenuate NLR receptor-triggered disease resistance and cell-death signaling. PLoS Genet. 10: e1004755.
Macho, A., and C. Zipfel, 2014 Plant PRRs and the activation of innate immune signaling. Mol. Cell 54: 263–272.

Maekawa, T., B. Kracher, S. Vernaldi, E. Ver Loren van Themaat, and P. Schulze-Lefert, 2012 Conservation of NLR-triggered immunity across plant lineages. Proc. Natl. Acad. Sci. USA 109: 20119–20123.

Mago, R., P. Zhang, S. Vaurtin, H. Simkova, U. Bansal et al., 2015 The wheat Sr30 gene reveals rich diversity at a cereal disease resistance locus. Nat Plants 1: 15186.

Mbengue, M., G. Bourdais, F. Gervasi, M. Beck, J. Zhou et al., 2016 Clathrin-dependent endocytosis is required for immunity mediated by pattern recognition receptor kinases. Proc. Natl. Acad. Sci. USA 113: 11034–11039.

Meng, Y., and R. P. Wise, 2012 HvWRKY10, HvWRKY19, and Mbengue, M., G. Bourdais, F. Gervasi, M. Beck, J. Zhou et al., 2016 Clathrin-dependent endocytosis is required for immunity mediated by pattern recognition receptor kinases. Proc. Natl. Acad. Sci. USA 113: 11034–11039.

Meng, Y., and R. P. Wise, 2012 HvWRKY10, HvWRKY19, and HvWRKY28 regulate Mla-triggered immunity and basal defense to barley powdery mildew. Mol. Plant Microbe Interact. 25: 1492–1505.

Meng, Y., M. J. Moscow, and R. P. Wise, 2009 Blufensin1 negatively impacts basal defense in response to barley powdery mildew. Plant Physiol. 149: 271–283.

Moscou, M. J., N. Lauter, R. A. Caldo, D. Nettleton, and R. P. Wise, 2011a Quantitative and temporal definition of the Mla transcriptional regulon during barley-powdery mildew interactions. Mol. Plant Microbe Interact. 24: 694–705.

Moscou, M. J., N. Lauter, B. Steffenson, and R. P. Wise, 2011b Quantitative and qualitative stem rust resistance factors in barley are associated with transcriptional suppression of defense regulons. PLoS Genet. 7: e1002208.

Mozhui, K., D. C. Giobanu, T. Schikorski, X. Wang, L. Lu et al., 2008 Dissection of a QTL hotspot on mouse distal chromosome 1 that modulates neuro-behavioral phenotypes and gene expression. PLoS Genet. 4: e100260.

Paul, P., S. Simm, O. Mirus, K. D. Scharf, S. Fragkostefanakis et al., 2014 The complexity of vesicle transport factors in plants examined by orthology search. PLoS One 9: e97745.

Periyanayan, S., J. Moore, M. Ayliffe, U. Bansal, X. Wang et al., 2013 The gene Sr33, an ortholog of barley Mla genes, encodes resistance to wheat stem rust race Ug99. Science 16: 786–788.

R Core Team, 2016 R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria.

Ritchie, M. E., B. Phipson, D. Wu, Y. Hu, C. W. Law et al., 2013 The complex of vesicle transport factors in plants examined by orthology search. PLoS One 9: e97745.

R Core Team, 2016 R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria.

Ritchie, M. E., B. Phipson, D. Wu, Y. Hu, C. W. Law et al., 2015 Limma powers differential expression analyses for RNA-sequencing and microarray studies. Nucleic Acids Res. 43: e47.

Rutter, B. D., and R. W. Innes, 2017 Extracellular vesicles isolated from the leaf apoplast carry stress-response proteins. Plant Physiol. 173: 728–741.

Samad-Zamini, M., W. Schweiger, T. Nussbaumer, K. F. X. Mayer, and H. Buerstmayr, 2017 Time-course expression QTL-atlas of the global transcriptional response of wheat to Fusarium graminearum. Plant Biotechnol. J. DOI: 10.1111/pbi.12729.

Sanderfoot, A. A., and N. V. Raikhel, 1999 The specificity of vesicle trafficking: coat proteins and SNAREs. Plant Cell 11: 629–642.

Schulze-Lefert, P., 2004 Knocking on the heaven behavior of phenotypes and gene expression. PLoS Genet. 4: e1000260. e1002208.

Shen, Q. H., F. Zhou, S. Bieri, T. Haizel, K. Shirasu et al., 2003 Recognition specificity and RAR1/SGT1 dependence in barley Mla disease resistance genes to the powdery mildew fungus. Plant Cell 15: 732–744.

Shirasu, K., 2009 The HSP90–SGT1 chaperone complex for NLR immune sensors. Annu. Rev. Plant Biol. 60: 139–164.

Smith, E., and L. Kruglyak, 2008 Gene–environment interaction in yeast gene expression. PLoS Biol. 6: e83.

Stark, C., B. J. Breitkreutz, T. Reguly, L. Boucher, A. Breitkreutz et al., 2006 BioGRID: a general repository for interaction datasets. Nucleic Acids Res. 34: D353–D359.

Steffenson, B. J., Y. Jin, B. G. Rossnagel, J. B. Rasmussen, and K. Kao, 1995 Genetics of multiple disease resistance in a doubled-haploid population of barley. Plant Breed. 114: 50–54.

Stennmark, H., and V. M. Olikkonen, 2001 The Rab GTPase family. Genome Biol. 2: REVIEWS0007.

Storey, J., and R. Tibshirani, 2003 Statistical significance for genomewide studies. Proc. Natl. Acad. Sci. USA 100: 9440–9445.

Storey, J. D., A. J. Bass, A. Dabney, and D. Robinson, 2015 qvalue: Q-value estimation for false discovery rate control pp. R package version 2.6.0. Available at: https://github.com/StoreyLab/qvalue.

Takahashi, N., M. Yanagida, S. Fujiyama, T. Hayano, and T. Isobe, 2003 Proteomic snapshot analyses of preribosomal ribonucleoprotein complexes formed at various stages of ribosome biogenesis in yeast and mammalian cells. Mass Spectrom. Rev. 22: 287–317.

Teh, O. K., and D. Hofius, 2014 Membrane trafficking and autophagy in pathogen-triggered cell death and immunity. J. Exp. Bot. 65: 1297–1312.

Uniprot Consortium, 2015 Uniprot: a hub for protein information. Nucleic Acids Res. 43: D204–D212.

Wei, F., K. Gobelman-Werner, S. M. Morroll, J. Kurth, L. Mao et al., 1999 The Mla (powdery mildew) resistance cluster is associated with three NBS-LRR gene families and suppressed recombination within a 240-kb DNA interval on chromosome 5H (1HS) of barley. Genetics 153: 1929–1948.

Wei, F., R. A. Wing, and R. P. Wise, 2002 Genome dynamics and evolution of the Mla (powdery mildew) resistance locus in barley. Plant Cell 14: 1903–1917.

West, M. A., K. Kim, D. J. Klebeinstein, H. van Leeuwen, R. W. Michelmore et al., 2007 Global eQTL mapping reveals the complex genetic architecture of transcript-level variation in Arabidopsis. Genetics 175: 1441–1450.

Wittkop, P. J., 2005 Genomic sources of regulatory variation in cis and trans. Cell. Mol. Life Sci. 62: 1779–1783.

Xu, W., Y. Meng, and R. P. Wise, 2014 Mla- and Rom1-mediated control of microRNA398 and chloroplast copper/zinc superoxide dismutase regulates cell death in response to the barley powdery mildew fungus. New Phytol. 201: 1396–1412.

Xu, W., Y. Meng, P. Surana, G. Fuerst, D. Nettleton et al., 2015 The knottin-like Blufensin family regulates genes involved in nuclear import and the secretory pathway in barley powdery mildew interactions. Front. Plant Sci. 6: 409.

Zhang, J., W. Li, T. Xiang, Z. Liu, K. Laluk et al., 2010 Receptor-like cytoplasmic kinases integrate signaling from multiple plant immune receptors and are targeted by a Pseudomonas syringae effector. Cell Host Microbe 7: 290–301.

Zhang, Y., R. Xia, H. Kuang, and B. C. Meyers, 2016 The diversification of plant NBS-LRR defense genes directs the evolution of microRNAs that target them. Mol. Biol. Evol. 33: 2692–2705.

Communicating editor: J. Faris