

# Differential Effects of Day/Night Cues and the Circadian Clock on the Barley Transcriptome<sup>1</sup>[OPEN]

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The circadian clock is a complex transcriptional network that regulates gene expression in anticipation of the day/night cycle and controls agronomic traits in plants. However, in crops, how the internal clock and day/night cues affect the transcriptome remains poorly understood. We analyzed the *diel* and circadian leaf transcriptomes in the barley (*Hordeum vulgare*) cultivar 'Bowman' and derived introgression lines harboring mutations in *EARLY FLOWERING3* (*ELF3*), *LUX ARRHYTHMO1* (*LUX1*), and *EARLY MATURITY7* (*EAM7*). The *elf3* and *lux1* mutants exhibited abolished circadian transcriptome oscillations under constant conditions, whereas *eam7* maintained oscillations of  $\approx 30\%$  of the circadian transcriptome. However, day/night cues fully restored transcript oscillations in all three mutants and thus compensated for a disrupted oscillator in the arrhythmic barley clock mutants *elf3* and *lux1*. Nevertheless, *elf3*, but not *lux1*, affected the phase of the *diel* oscillating transcriptome and thus the integration of external cues into the clock. Using dynamical modeling, we predicted a structure of the barley circadian oscillator and interactions of its individual components with day/night cues. Our findings provide a valuable resource for exploring the function and output targets of the circadian clock and for further investigations into the *diel* and circadian control of the barley transcriptome.

The circadian clock is a time-keeping mechanism that reflects the day/night cycle through an endogenous transcriptional rhythm to anticipate dawn and dusk

(McClung, 2006). This clock synchronizes internal rhythms with external light and temperature cycles (Harmer, 2009; Greenham and McClung, 2015). The prevalence of circadian rhythms in all domains of life suggests that circadian clocks provide an adaptive advantage for organisms (Edgar et al., 2012). The Arabidopsis (*Arabidopsis thaliana*) oscillator contains an interconnected regulatory network of transcriptional repressors and activators (Hsu et al., 2013; Fogelmark and Troein, 2014). These components are expressed sequentially to regulate output genes through regulatory elements present in target promoters (Harmer et al., 2000; Covington et al., 2008; Michael et al., 2008b).

In the Arabidopsis circadian oscillator, two morning-expressed MYB transcription factors, CIRCADIAN CLOCK ASSOCIATED1 (*CCA1*) and LATE ELONGATED HYPOCOTYL (*LHY*), inhibit the expression of *TIMING OF CAB EXPRESSION1/PSEUDO-RESPONSE REGULATOR1* (*TOC1/PRR1*) that in turn represses the transcription of *CCA1* and *LHY* in the night (Alabadí et al., 2001; Gendron et al., 2012; Huang et al., 2012). During the day, sequentially expressed *PRR9*, *PRR7*, and *PRR5* repress the transcription of *CCA1* and *LHY*

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M.v.K. and S.J.D. conceived the original research project; L.M.M. and M.v.K. designed the experiments; L.M.M. carried out the experiments and analyzed the RNA-sequencing data; L.M. calculated LTI models with the help of J.G. and A.A.R.W.; A.P. contributed to the bioinformatics analyses; L.M.M., L.M., A.A.R.W., A.P., and M.v.K. wrote the article.

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(Nakamichi et al., 2010, 2012; Liu et al., 2016). CCA1 and LHY in turn promote the expression of PRR9 and PRR7 (Farré et al., 2005). However, this effect might be indirect because transient induction analysis demonstrated that CCA1 can directly repress the expression of both genes PRR9 and PRR7 (Kamioka et al., 2016). Similarly, REV-EILLE8 (RVE8), a close homolog of CCA1/LHY directly activates transcription of PRR5 and TOC1, and likely other evening-phased genes (Farinas and Mas, 2011; Rawat et al., 2011; Nakamichi et al., 2012; Hsu et al., 2013). The downregulation of CCA1 and LHY by PRR proteins allows the induction of *EARLY FLOWERING3* (*ELF3*), *ELF4*, and *LUX ARRHYTHMO* (*LUX*), which encode the Evening Complex (EC) of proteins (Hazen et al., 2005; Kikis et al., 2005). The EC acts at dusk as a transcriptional repressor of PRR9 expression (Helfer et al., 2011; Nusinow et al., 2011; Herrero et al., 2012). Furthermore, the evening-expressed GIGANTEA (GI) protein was modeled as a negative regulator of the EC, which in turn inhibits TOC1 expression (Huang et al., 2012; Pokhilko et al., 2012).

The complex network of transcriptional regulators at the core of the clock underscores the role of transcriptional regulation as a central regulatory mechanism for circadian oscillation. Consequently, the Arabidopsis clock is a master regulator of transcription and controls approximately 30% to 40% of the global gene expression in a time-of-day-specific cycling pattern where transcription of functionally related genes often peaks in clusters (Harmer et al., 2000; Covington et al., 2008; Michael et al., 2008a; Staiger et al., 2013). Expression of such functional clusters often precedes or coincides with the underlying physiological event (Covington et al., 2008; Michael et al., 2008a), suggesting that circadian control anticipates *diel* regulation to improve physiological performance (Greenham and McClung, 2015).

In Arabidopsis, the circadian system controls many agronomically important processes, such as metabolism, growth, photosynthesis, and flowering time (Greenham and McClung, 2015). Consequently, it has been suggested that the circadian clock is key to improving adaptation and performance of crop plants (Hsu and Harmer, 2014; Bendix et al., 2015). Putative circadian oscillator genes have been identified in the monocot crop barley (*Hordeum vulgare*) based on their homology with the Arabidopsis clock genes (Campoli et al., 2012; Calixto et al., 2015). Although the circadian oscillator genes diversified via duplication independently between the monocot and eudicot clades, their structure and expression patterns remained highly similar (Campoli et al., 2012; Hsu and Harmer, 2014; Bendix et al., 2015). For example, in monocots, the morning-expressed MYB-like transcription factor LHY is the only ortholog of the Arabidopsis paralogs CCA1 and LHY (Takata et al., 2009; Campoli et al., 2012). *HvLHY* overexpression in Arabidopsis causes arrhythmia, suggesting circadian functionality (Kusakina et al., 2015). The PRRs duplicated independently from three ancient PRR genes after the divergence of monocots and eudicots such that the orthologous relationship

within the PRR3/7 and PRR5/9 clades of Arabidopsis and monocot plants cannot be immediately resolved (Takata et al., 2010). Partial complementation of Arabidopsis *prp7-11* by *HvPRR37* suggests that the barley gene might retain some functionality of the Arabidopsis ortholog (Kusakina et al., 2015). However, PRR37 orthologs in monocots—*PPD1* in barley and wheat (*Triticum aestivum*; Turner et al., 2005; Beales et al., 2007), and *SbPRR37* in sorghum (*Sorghum bicolor*; Murphy et al., 2011)—are major determinants of photoperiod sensitivity and flowering time. In Arabidopsis, PRR proteins stabilize the CONSTANS protein, a floral promoter, and repress CYCLING DOF FACTOR genes, thereby promoting flowering under long days (Nakamichi et al., 2007; Hayama et al., 2017). However, natural variation in PRR genes in Arabidopsis do not have any notable effect on flowering time (Ehrenreich et al., 2009). Furthermore, the genes underlying the two *early maturity* mutants, *early maturity8* (*eam8*) and *eam10*, have been identified as barley homologs of the Arabidopsis clock genes *ELF3* and *LUX1*, respectively (Faure et al., 2012; Zakhrebekova et al., 2012; Campoli et al., 2013). Mutations in both genes cause photoperiod insensitivity and early flowering under long- and short-day conditions in barley (Faure et al., 2012; Zakhrebekova et al., 2012; Campoli et al., 2013). In addition, several *ELF4*-like homologs that exist in barley, including *HvELF4-like 4*, can complement an Arabidopsis *Atelf4* null mutant (Hicks et al., 2001; Kolmos et al., 2009). Whereas a number of putative clock components in barley have been identified, there is little information on the contribution of the clock versus day/night cues on the global transcriptome in barley.

We generated *diel* and circadian RNA-sequencing (RNA-seq) datasets of four barley genotypes—the spring barley ‘Bowman’ and three derived introgression lines with mutations in *HvELF3* (BW290), *HvLUX1* (BW284), and *EAM7* (BW287; Faure et al., 2012; Campoli et al., 2013). The candidate gene for *EAM7* has not yet been identified, but loss of *EAM7* function accelerates flowering by abolishing sensitivity to the photoperiod (Gallagher et al., 1991). We used the RNA-seq time-course data to analyze the effects of barley clock genes on *diel* and circadian transcriptome oscillations, including changes in phase and period under constant conditions and light and dark cycles. Dynamical modeling allowed us to predict a molecular structure of the barley circadian oscillator and to uncover how circadian oscillator components interact with day/night cues to regulate the global transcriptome in barley.

## RESULTS

### *Diel* and Circadian Oscillations of the Barley Transcriptome

We analyzed the *diel* and circadian global leaf transcriptome of the barley ‘Bowman’ and the derived introgression lines carrying mutations in *HvELF3* (BW290),

*HvLUX1* (BW284), and *HvEAM7* (BW287). Plants were grown under cycles of 12-h light and 12-h night (LD) and the second leaf of replicate plants was harvested every 4 h over 24 h. Additional samples were taken in at 2-h intervals at dusk in all genotypes and additionally at dawn in cv Bowman (Supplemental Fig. S1). Thereafter, plants were transferred to constant light and temperature conditions (LL) and leaf samples were taken every 4 h for 36 h starting from the first subjective night. The sampling strategy was optimized for modeling the circadian oscillator using systems identification, an approach we have used previously to model the circadian *Arabidopsis* (Dalchau et al., 2010; Herrero et al., 2012; Mombaerts et al., 2019). Systems identification models the network based on the dynamical information in time series data sets. The dynamical information is greatest when the system is perturbed by environmental or genetic stimulation. We therefore optimized our RNA-seq strategy to capture the data with a time series that included light-dark transitions to perturb the oscillator and sampled several genotypes with alterations in putative oscillator genes. Whereas it is typical to observe circadian dynamics in prolonged constant conditions to distinguish free-running from entrained behaviors, circadian dynamics occur in both LD and LL and the LD conditions have greater perturbation to inform the modeling. We sampled for 36 h in LL because barley circadian transcript oscillations rapidly dampen upon transfer to constant conditions, which results in a high signal-to-noise ratio of the rhythms that would have compromised the systems identification (Campoli et al., 2012; Hughes et al., 2017). Our strategy of sampling before the onset of dampening of the signal reduces the false-negative discovery rate by increasing the signal-to-noise. This is at the expense of possibly increasing the false-negative rate by having a shorter time series in constant light (Hughes et al., 2017). Consequently, by taking a conservative approach to network identification, we can be more confident of the connections we have identified.

Individual libraries were single-end-sequenced on a HiSeq 2500 System (Illumina) with 10 million reads per library and reads were mapped against a custom reference sequence consisting of 68,739 transcripts (Digel et al., 2015). The nomenclature of the gene models used in this study (Digel et al., 2015) was cross referenced with the identifiers of the HORVU gene models annotated on the barley pseudochromosomes (Mascher et al., 2017). Raw read counts normalized to counts per million (CPM) were used for the downstream rhythmic analysis and modeling. We determined the oscillating patterns of gene expression, including period as the duration of one complete oscillation, and phase as the time point of transcript peak expression (Yang and Su, 2010; Wu et al., 2016). To increase the analytical power for the rhythmic analysis, the 24-h *diel* dataset, but not the 36-h LL data, was duplicated to imitate 48 h of sampling data.

We identified 18,500 transcripts with expression levels >5 CPM in at least two libraries. Among 18,500 transcripts expressed across all the investigated lines,

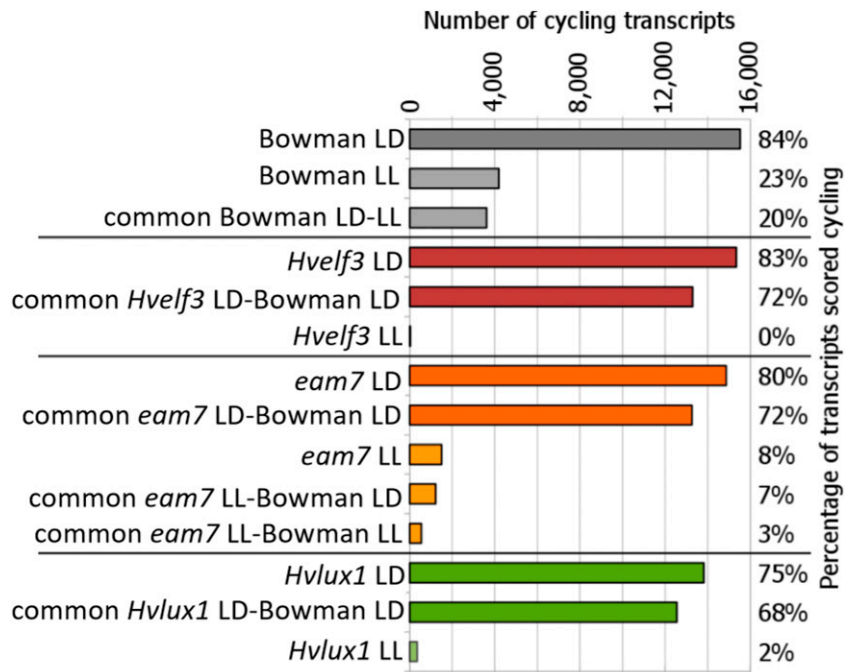
84% were scored as rhythmic under LD in cv Bowman (Fig. 1; Supplemental Dataset S1). The duplicated LD data sets may have resulted in elevated false-positive rates of *diel*-regulated genes estimated per genotype. However, we found that out of all expressed genes ~70% were scored as rhythmic in cv Bowman and at least one other mutant genotype, which represent independent biological data (Fig. 1).

Under LL, ~23% of the 18,500 transcripts were rhythmic, which is a feature of clock-regulated genes (Fig. 1; Supplemental Dataset S1). The gene ontology (GO) analyses revealed that, in cv Bowman under LL, the circadian-controlled transcripts were primarily related to the processes of regulation of DNA-dependent transcription, translation, electron transport, signal transduction, responses to salt stress and cold, and metabolic processes, including amino acid, Suc and starch metabolism (Fig. 2E; Supplemental Dataset S1). The molecular functions of the circadian-controlled transcripts in cv Bowman in LL were primarily represented by protein-, zinc ion-, and ATP binding, DNA- and nucleotide binding, and sequence-specific DNA-binding transcription factor activity GO terms (Fig. 2E).

We found that the majority of the transcripts expressed rhythmically under LL were also rhythmic under LD (20% of all the transcripts, 87% of LL transcripts). This demonstrated that approximately one-quarter of the cv Bowman transcriptome is modulated by the circadian clock, whereas the largest proportion of the rhythmic transcripts in LD required day/night cues for their rhythmic expression.

The large impact of external transitions on transcriptome oscillations independent of the clock was further supported by analysis of *Hvelf3* plants. In *Hvelf3*, no transcript rhythms were detected under LL, demonstrating that HvELF3 function is required for self-sustained transcriptome oscillations in barley (Fig. 1; Supplemental Dataset S1). Environmental cues under LD restored oscillatory dynamics in the *Hvelf3* loss-of-function line with 83% of the global transcriptome being rhythmic in the *Hvelf3* plants (Fig. 1). The number and the identity of oscillating transcripts were similar between *Hvelf3* and cv Bowman plants under *diel* cycles (Fig. 1). In *Hvlux1* plants, only 2% of the expressed transcripts were rhythmic under LL suggesting that, like HvELF3, HvLUX1 is required for free-running oscillations under LL (Fig. 1). Once again, LD cycles were sufficient to restore transcriptional rhythms in the *Hvlux1* mutant, i.e. 75% of the transcriptome oscillated in *Hvlux1* plants under LD (Fig. 1). Mutation of the *EAM7* locus in BW287 reduced the pervasiveness of circadian transcriptional oscillations, but did not completely abolish them because 8% of the expressed transcripts cycled under LL in *eam7*, which was approximately one-third of the number of the oscillating transcripts in cv Bowman (Fig. 1; Supplemental Dataset S1). Under LD, 80% of the global transcriptome was rhythmic in *eam7* and 72% of the rhythmic transcripts were common between *eam7* and the cv Bowman plants (Fig. 1).

**Figure 1.** Fraction of transcripts with oscillating transcription pattern in cv Bowman and cv Bowman-derived introgression lines with mutations in *Hvelf3*, *eam7*, and *Hvlux1* under LD and LL. Fractions refer to a total of 18,500 transcripts expressed in all genotypes.



Our data demonstrate that cycles of light and temperature and the circadian oscillator drive rhythmic expression in barley. HvELF3, HvLUX1, and EAM7 contribute to free-running oscillations under constant conditions whereas environmental rhythms are sufficient to drive rhythmic expression in the absence of a free-running oscillator.

#### EAM7 Is a Modulator of a Bimodal Phase Distribution under LL and Shortens the Free-Running Period

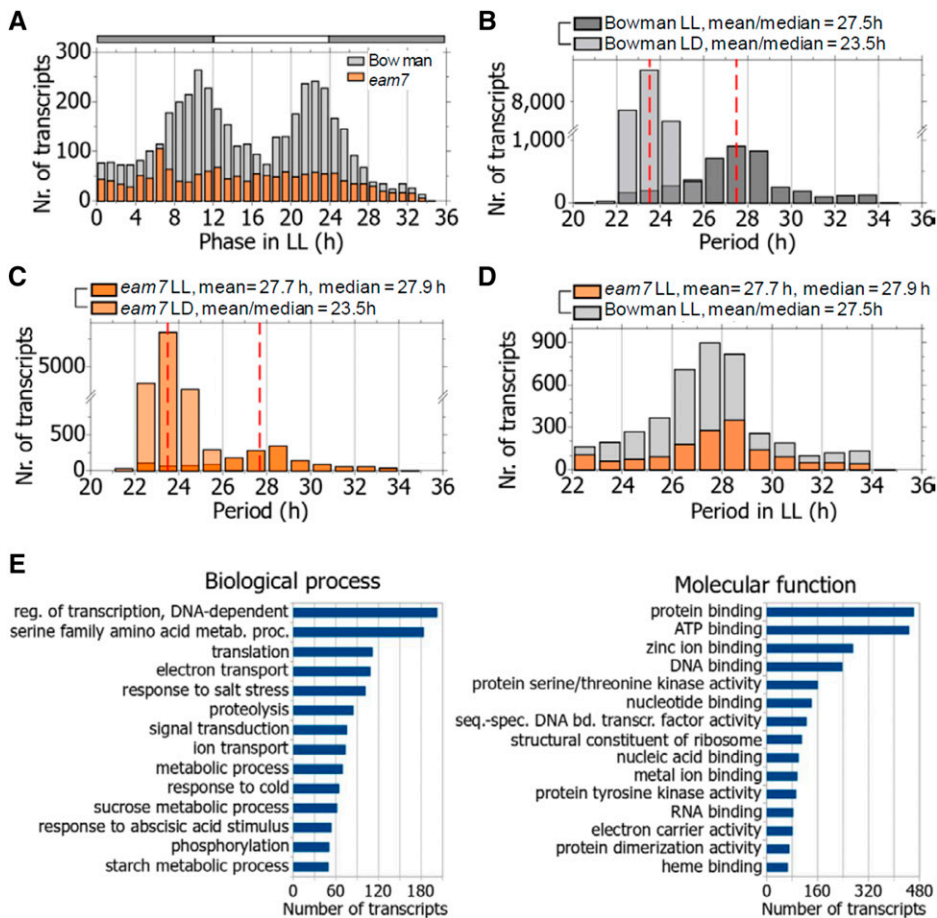
To investigate temporal expression patterns of the circadian-regulated transcripts under free-running conditions, we estimated the phase and the period of every circadian-regulated transcript in cv Bowman and *eam7*, the two genotypes that sustained free-running circadian rhythms. In cv Bowman, the distribution of the circadian transcriptome expression phase followed a bimodal pattern with the highest number of transcripts peaking shortly before the transitions to subjective days and nights (Fig. 2A; Supplemental Dataset S1). By contrast, in *eam7*, this phase pattern of the cumulative circadian transcriptome was not evident (Fig. 2A). These findings indicate that EAM7 is required to modulate the characteristic bimodal pattern of the circadian transcriptome expression in barley. The period estimates of the oscillating transcripts under LL ranged between 22 h and 34 h in cv Bowman and *eam7* and followed a bell-shaped distribution with mean periods of 27.5 h and 27.7 h in cv Bowman and *eam7*, respectively (Fig. 2, B–D). The transcript periods were not statistically different between cv Bowman and *eam7*. In both cv Bowman and *eam7*, the SD of the period distribution was higher under LL (6 h) than under LD

(2.5 h; Fig. 2, B and C). This could arise from either the uncoupled nature of cellular oscillations in free-running conditions or as a consequence from the period estimation as the signal amplitude was lower in LL than in LD.

#### Regulation of the Transcriptome-Wide Phase in Day/Night Cycles

Next, we investigated the transcriptome oscillations under the *diel* LD conditions. In all genotypes, including those that were arrhythmic in LL, the mean of the period distribution was consistent with the enforced 24-h *diel* cycle and ranged between 23.5 h and 23.6 h (Supplemental Fig. S2). The phase was bimodally distributed over the day/night cycle in cv Bowman so that for the highest number of transcripts, the peak of expression occurred before dawn and dusk, and the number of transcripts with their peak expression during the night and day was the lowest (Fig. 3A). This pattern was similar to the phase distribution under LL (Fig. 2A). The transcripts that oscillated in both LL and LD were also bimodally distributed under the *diel* cycles, although the bimodal pattern of these genes was less pronounced under LD (Fig. 3A). This suggested that the bimodal distribution of transcriptome-wide gene expression is, at least partly, under the control of the circadian clock.

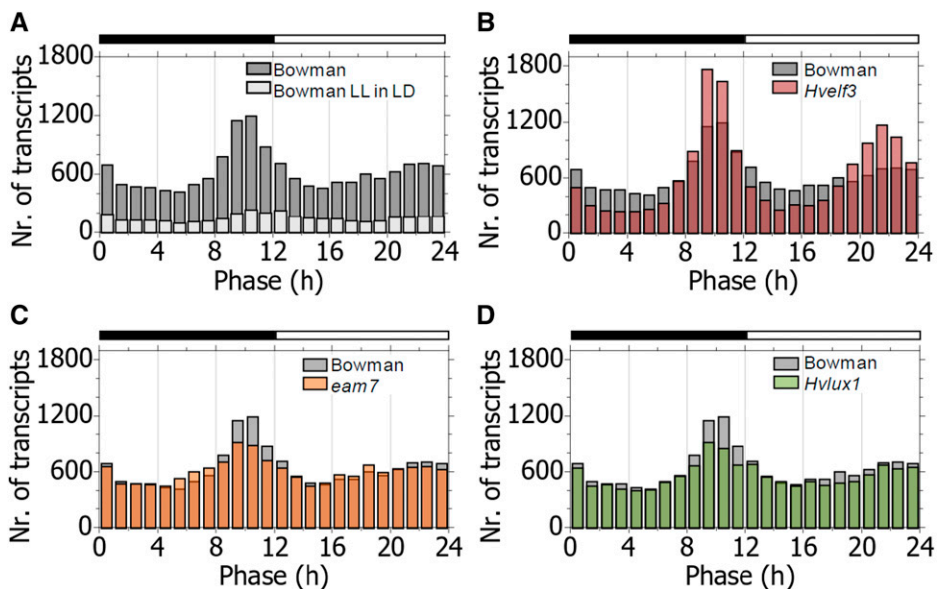
The analysis of the clock mutants, however, suggested that the bimodal phase distribution under LD is controlled by both the circadian clock and day/night cues. In *Hvelf3*, the phase was bimodally distributed under *diel* cycles similar to cv Bowman; however, the quantitative characteristics of the phase distribution



**Figure 2.** Distribution of the period and the phase of the oscillating transcriptome in constant light and their involvement in biological processes and molecular functions. A, Phase distribution in LL of cv Bowman and *eam7*. Gray-orange bars indicate the subjective night (gray) and subjective day (orange) in LL. B and C, Period distribution of the oscillating transcriptome under LL in comparison with LD in (B) cv Bowman and (C) *eam7*. D, Comparison of the period distribution in LL between cv Bowman and *eam7*. E, Top-15 categories of the GO terms for biological processes and molecular function of the transcripts oscillating in cv Bowman under LL.

differed. Specifically, in *Hvelf3*, the phase distribution showed higher peaks at dawn and dusk and deeper troughs during the night or the day than in cv Bowman (Fig. 3B). A large number of the transcripts that peaked around the night-to-day and day-to-night transition in

*Hvelf3* (Fig. 3B) peaked during the day or the night in cv Bowman. This demonstrated that HvELF3 modulates timing of peak expression of multiple transcripts in day/night cycles. This effect was apparently completely or partially independent of the oscillator defect



**Figure 3.** Distribution of the phase of the oscillating transcriptome in night/day cycles. A, Phase distribution in cv Bowman in LD cycles for the global oscillating transcriptome and those transcripts detected oscillating in both night/day cycles and constant light (cv Bowman LL in LD). B to D, Phase distribution in diel cycles in *Hvelf3* mutant (B), *eam7* mutant (C), and *Hvlux1* mutant (D) in comparison with cv Bowman.

that causes arrhythmia in the *Hvelf3* plants under LL, because the phase distribution in *Hvlux1* mutants under LD was similar to that in cv Bowman (Fig. 3, C and D), even though self-sustained circadian oscillations were also absent in this genotype under LL conditions (Fig. 1). This was also evident from the transcriptome-wide comparison of the phase between the barley clock mutants with cv Bowman under LD (Supplemental Fig. S3). Here, the phase distributions strongly correlated between *Hvlux1* and cv Bowman (Pearson correlation  $\rho = 0.97$ ,  $R^2 = 0.94$ ), whereas the phase distributions in *Hvelf3* and cv Bowman were correlated to a lower degree (Pearson correlation  $\rho = 0.93$ ,  $R^2 = 0.86$ ), even though both mutant genotypes harbor an arrested oscillator under LL conditions (Fig. 1).

Day/night cycles had strong effects on the phase distribution of the transcriptome as demonstrated by the analysis of the *eam7* transcriptome. Whereas the phase distribution was not bimodal in *eam7* under LL (Fig. 2A), under LD, the phase distribution was bimodal, similar to that in cv Bowman (Fig. 3C). Consistently, the phase distributions under LD were highly correlated between *eam7* and cv Bowman (Pearson correlation  $\rho = 0.96$ ,  $R^2 = 0.92$ , Supplemental Fig. S3). Consequently, external cues under LD controlled the phase of the global transcriptome in *eam7* to peak at the night/day transitions despite the circadian defects observed in *eam7* under LL. Together, these results demonstrate that the bimodal distribution of the phase in *diel* cycles is controlled by both day/night cues and the clock component HvELF3. The genetic defects and their underlying circadian phenotypes in *Hvlux1* and *eam7* have limited effects on the phase of the global oscillating transcriptome in *diel* cycles despite their strong transcriptional phenotypes under LL.

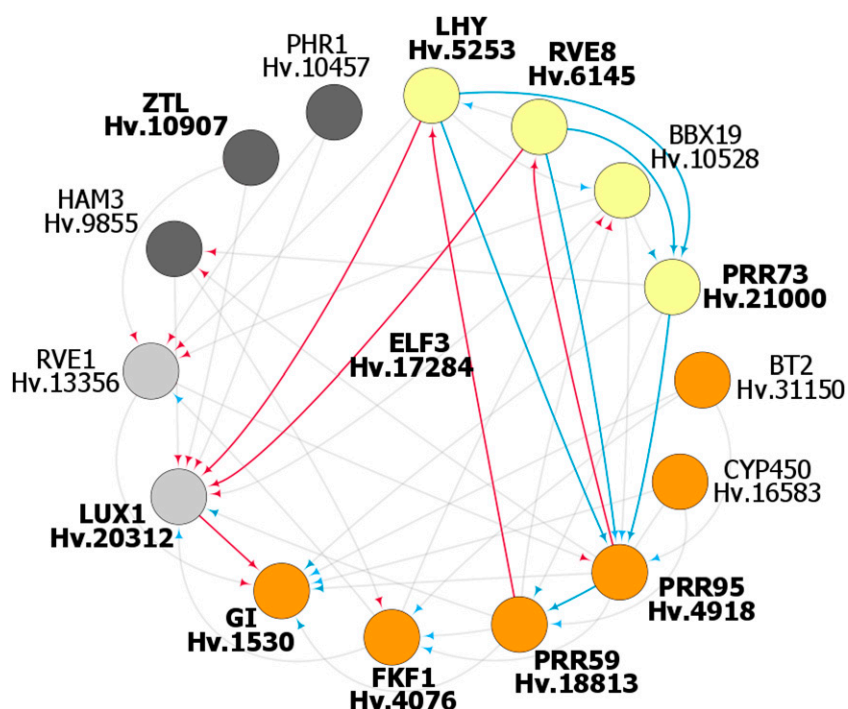
### Dynamical Models Predict Components and Regulatory Interactions of the Barley Clock

We then sought to infer the regulatory relationships between components of the barley circadian clock. To this end, we modeled a transcriptional network based on the RNA-seq time series data. Our data suggested that HvELF3 and HvLUX1 are integral components of the barley oscillator as they were necessary to sustain transcriptome oscillations under LL (Fig. 1). Therefore, we hypothesized that modeling a transcriptional network around HvELF3 and HvLUX1 could identify the regulatory relationships that shape the circadian clock in barley. We followed an approach that searches the dynamic dependencies of HvELF3 and HvLUX1 expression on other transcripts. We used linear time invariant (LTI) models for interpreting expression data without relying on a priori knowledge of the transcriptional network (Dalchau et al., 2010; Herrero et al., 2012; Mombaerts et al., 2019; Supplemental Information). LTI models require transcriptional data sets that display robust changes in expression over time under free-running conditions. Therefore, only the expression

datasets from cv Bowman and *eam7* could be used in modeling, because their transcriptomes oscillated under LL conditions. In both cv Bowman and *eam7*, the transcripts encoding HvELF3 had a very low signal-to-noise ratio due to low rhythmicity under LL and could not be used for modeling. We therefore rooted the network around *HvLUX1*, which displayed robust oscillatory dynamics (Supplemental Fig. S4).

To reduce the identification of erroneous interactions, we filtered all circadian transcripts for those that were homologous to Arabidopsis genes representing transcription factors that were labeled circadian, and thus show circadian expression but are not necessarily components of the core clock (www.geneontology.org). Indeed, whereas our modeling methodology is computationally inexpensive, the uncertainty about the structure of the network is increasing exponentially with the number of genes considered. Additionally, we filtered the resulting 131 transcripts (Supplemental Dataset S2) for those that exhibited unambiguous dynamics and a high signal-to-noise ratio of expression in both cv Bowman and *eam7*. This filter was applied because of the transitional nature of constant light data, which typically shows a large decrease of amplitude after a few hours in barley (Campoli et al., 2012), and the dependency of noise on gene expression levels. *Hvlux1* and *Hvelf3* datasets were not considered in the following network analysis because these mutations led to the arrhythmic transcriptomes. This resulted in 42 transcripts in cv Bowman and 41 in *eam7*, of which 35 transcripts were in common between cv Bowman and *eam7* and used for modeling (Supplemental Dataset S3). For the 35 shared gene transcripts, we predicted all possible pairwise (or single source-target) dynamic dependencies based on the transcript abundance over time for both the cv Bowman and the *eam7* datasets. For each of these 35 pairwise comparisons, we fitted a model that captures the expression pattern of *HvLUX1* in each of the two genotypes.

We then investigated the consistency between the models obtained for cv Bowman and *eam7* using the v-gap metric (Supplemental Fig. S5; Supplemental Dataset S4). This approach estimates differences between models and allowed us to identify regulatory interactions that were maintained or abolished in the *eam7* mutant (Mombaerts et al., 2019). Following this approach, we identified 20 transcripts and 79 regulatory links in cv Bowman, of which 15 transcripts and 49 regulatory links could be cross validated in *eam7* (Fig. 4; Supplemental Figs. S5 and S6; Supplemental Dataset S4). Five transcripts could not be confirmed in *eam7* because they were arrhythmic in *eam7*, or the regulatory link could not be modeled with high confidence (i.e. model fitness), or the models for cv Bowman and *eam7* displayed a large difference (i.e. v-gap). Among the five genes that could not be confirmed in *eam7* transcripts, those with homology to RVE8, RVE6, and RVE4 (Hv.12868, HORVU7Hr1G001830.3), and RVE1 (Hv.25709, HORVU6Hr1G066450.5), were the most prominent, with eight and seven links to putative clock genes in cv



**Figure 4.** The putative circadian network of the barley oscillator as predicted from time series expression data. Genetic evidence but no model prediction allowed placing *HvELF3* as a core clock component. The figure displays the inferred components and interactions that constitute the barley circadian transcriptional network (also see Supplemental Information). Circadian clock components are represented by circles and sorted in clockwise direction for the time point of peak expression starting with *HvLHY* at dawn (yellow, morning; orange, evening; gray, night). The regulatory interactions are represented by directional arrows, where activation is marked in blue and inhibition in red. The components printed in bold and the links highlighted in color are consistent with key components and key regulatory principles present in circadian clock models from *Arabidopsis*.

Bowman, respectively. Both transcripts displayed rhythmic oscillations under LL in cv Bowman, but were arrhythmic in *eam7*. Among the 15 cross-validated candidates, nine transcripts were encoded by barley genes homologous to known *Arabidopsis* core oscillator genes. In addition to *HvLUX1* as a core of the model (Hv.20312, HORVU3Hr1G114970), the predicted components of barley circadian clock were barley homologs of *LHY* (Hv.5253, HORVU7Hr1G070870; Mizoguchi et al., 2002), *RVE8* (Hv.6145, HORVU6Hr1G066000; Hsu et al., 2013), *PRR95* (Hv.4918, HORVU5Hr1G081620; Farré et al., 2005), *PRR59* (Hv.18813, HORVU4Hr1G021000; Nakamichi et al., 2010), *FLAVIN-BINDING*, *KELCH REPEAT*, *F-BOX 1* (Hv.4076, HORVU7Hr1G099010; Baudry et al., 2010), *GI* (Hv.1530, HORVU3Hr1G021140; Dalchau et al., 2011), and *ZEITLUPE* (*ZTL*; Hv.10907, HORVU6Hr1G022330; Más et al., 2003). Such a result supports that putative circadian-clock genes are themselves strongly driven by circadian processes.

On the resulting conjunction network, we noted that *HvPRR95* appeared as a hub with eight connections, whereas *HvLUX1*, as the origin of the graph, had 11 connections. This indicated a significant role of *HvPRR95* in the regulation of the core circadian genes. Therefore, we repeated the search for regulators of *HvPRR95*, computed their interactions in both datasets, and tested for their consistency. Consequently, four genes were added to the final network, namely *REVEILLE-like7* (*HvRVE7*, Hv.13356, HORVU2Hr1G104580), *Cytochrome P450* (Hv.16583), *HvPRR73* (HORVU4Hr1G057550; Farré et al., 2005), and *BTB/POZ and TAZ domain-containing protein2* (*HvBT2*, Hv.31150, HORVU3Hr1G092090; Fig. 4; Supplemental Dataset S4). Our modeling did not

place barley homologs of other known *Arabidopsis* clock genes *TOC1*, *ELF4*, *HvELF3*, and *PRR37* in the barley clock model. In the case of *HvTOC1* and barley homologs of *ELF4*, the inferred regulatory interactions to other putative clock components were weak and did not pass the cutoff filter. The weak transcript oscillations of *HvELF3* precluded its modeling as part of the barley clock. However, we placed *HvELF3* in the core clock model based on the genetic evidence that *HvELF3* is required for clock function in barley (arrhythmic phenotype of *Hvelf3* mutant). Furthermore, *PRR37*, the photoperiod response gene *Ppd-H1*, did not display rhythmic oscillations under LL and was therefore not placed into the circadian oscillator. Based on the timing of the peak expression starting with *HvLHY* expression at subjective dawn, we arranged the predicted components into a model of the barley circadian clockwork (Fig. 4).

In addition to the barley homologs of known *Arabidopsis* oscillator genes, our analysis suggested several previously uncharacterized components of barley circadian clock. These included the B-Box Zinc Finger Protein19 (*HvBBX19*, Hv.10528, HORVU5Hr1G081190) and *HvRVE7* (Hv.13356, HORVU2Hr1G104580; Fig. 4). In our model, both *HvBBX19* and *HvRVE7* regulate *HvPRR95* and are regulated by *HvLHY* (Fig. 4). The modeling predicted that *HvRVE7* represses *HvPRR95*, and *HvBBX19* activates *HvPRR73* and *HvPRR95*. Other predicted components of the barley circadian clock were a homolog of *HAIRY MERISTEM3* (*HAM3*; Hv.9855, HORVU6Hr1G063650) of *BT2*, *CYTCHROME 450* (Hv.16583, HORVU2Hr1G025160), and *PHOSPHATE STARVATION RESPONSE1* (Hv.10457, HORVU4Hr1G051080). However, all of these players

were predicted to regulate clock components, but were not themselves regulated by the clock genes (Fig. 4). To summarize, our analysis was able to predict components of the barley clock, which are close homologs of the Arabidopsis clock components (Campoli et al., 2012; Calixto et al., 2015) and additionally identified HvBBX19, HvRVE7, and HvHAM3 as putative components of the barley clock.

#### Relationship between Internal and External Cues to Regulate the Global Transcriptome in Barley

To quantify the relationship between the circadian oscillator and light signaling in regulating the rhythmicity of barley transcripts, LTI models that integrate both inputs explicitly were computed for each transcript. As a morning clock gene, the expression pattern of *HvLHY* accounted for the contribution of the circadian oscillator, whereas the light/dark cycle was integrated as a rectangular input (1 = light ON, 0 = light OFF; Supplemental Fig. S7). The expression pattern of the output transcript was approximated by finding the combination of inputs that fit the data best. Then, the contribution of each input was formally compared using a Bode analysis (Dalchau et al., 2010). The analysis estimated that 43% of the transcripts that oscillate in both day/night cycles and constant light were predominantly controlled by the circadian clock in light/dark cycles and that 48% were co-regulated by the circadian clock and light/dark cues (Fig. 5A; Supplemental Dataset S1). Only 9% of the transcripts were primarily controlled by light/dark cues (Fig. 5A; Supplemental Dataset S1). This is consistent with the expected under-representation of light/dark-controlled transcripts in a set of genes that oscillate in the absence of environmental cues.

We next investigated the phase relationship between driven and free-running conditions for transcripts predicted to be under clock control, light control, and co-regulated by light and the clock by the Bode analysis (Fig. 5, B and C; Supplemental Dataset S1). The clock-dominated transcripts revealed the highest correlation ( $R^2 = 0.64$ ; Fig. 5B) and the light-dominated transcripts the lowest correlation ( $R^2 = 0.27$ ) of the phase between day/night cycles and constant light (Fig. 5B). The correlation of the phase of co-regulated transcripts under LD and LL was intermediate ( $R^2 = 0.45$ ; Fig. 5B). This suggests that transcripts dominated by the circadian clock maintained a similar expression phase under changing light conditions, whereas the phase of transcripts dominated by light cues reflected the changes in light. These findings suggest that the Bode analysis predicted the main regulatory principles that determine the phase of oscillating transcription in day/night cycles. Namely, it suggests that ~40% of the transcripts with clock-maintained oscillations reveal a phase dominated by the circadian clock in *diel* cycles. For the remaining 60% of the transcripts with clock-maintained oscillations, the peak of their expression is under the control of light signaling pathways or co-regulated by

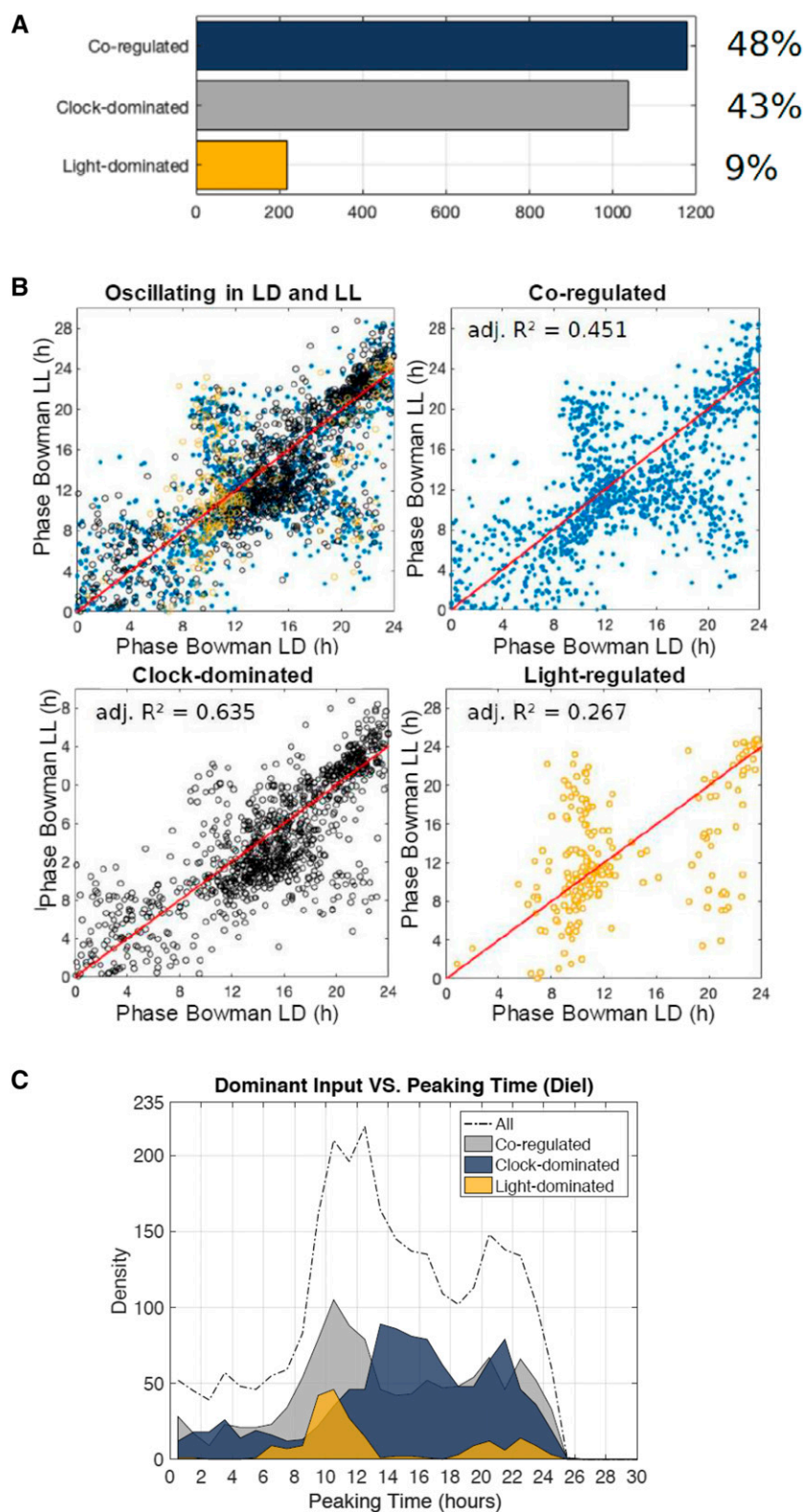
light signaling and clock. This finding highlights the importance of light signaling pathways to regulate the phase of oscillating transcription even for the transcripts, the rhythmicity of which is maintained by the circadian clock.

#### DISCUSSION

The circadian clock was estimated to control a large proportion (~25%) of the barley transcriptome under constant conditions, which is similar to estimates for the proportion of circadian transcripts in Arabidopsis, rice (*Oryza sativa*), and poplar (*Populus trichocarpa*; Covington et al., 2008; Michael et al., 2008b; Filichkin et al., 2011; Gehan et al., 2015). Despite the strong control of the clock on transcript oscillations under LL, day/night cues had a major influence on shaping expression patterns of circadian transcripts under *diel* conditions. First, the expression phase under LL conditions was generally not a strong predictor of the transcript phase under LD conditions. The expression phase of transcripts was therefore a plastic trait where LD conditions delayed or advanced expression phase as compared to LL depending on the transcript. Second, the Bode analysis demonstrated that the majority of circadian transcripts were regulated by light/temperature or a combination of the clock and light/temperature cues under LD conditions. It is well known that the circadian clock is dynamically plastic and constantly entrained by metabolic and environmental cues for synchronization with the cycles of the environment (Webb et al., 2019).

Here, however, we demonstrate that day/night cues entrain the clock and can also largely compensate for the lack of a functioning oscillator. The *Hvelf3* and *Hvlux1* mutants, with no cycling transcriptome under LL conditions, were characterized by transcriptome oscillations under LD comparable to wild-type cv Bowman. In this context, it is interesting to note that *Hvelf3* and *Hvlux1* mutants, which have a disrupted circadian clock, have been used to breed for barley cultivars adapted to northern European environments with strong daily and seasonal changes in light and temperatures (Faure et al., 2012; Campoli et al., 2013; Pankin et al., 2014). Neither of these two arrhythmic mutants has been reported to display any obvious impairment in photosynthesis and growth under conditions of pronounced photo- and thermocycles, in contrast to the corresponding Arabidopsis mutants (Faure et al., 2012; Campoli et al., 2013; Habte et al., 2014). Similarly, Izawa et al. (2011) reported that an *osgi* mutant in the field is not affected in photosynthesis and yield. Only under atypical growing conditions with late transplanting dates in the field was fertility significantly reduced in *osgi* plants, indicating a loss of seasonal adaptability. Our data suggest that *diel* cycles can compensate for circadian defects in the barley clock mutants by increasing the number of oscillating transcripts compared to free-running conditions and





**Figure 5.** Relationship between external and internal cues to regulate the phase of the barley transcriptome. A, Fractions of transcripts identified as clock-dominated, co-dominated by the clock and light, and light-dominated by the Bode analysis. B, Phase relationships among diel cycles (LD) and constant light (LL) for all transcripts oscillating in LD and LL, those co-regulated by the circadian clock, those dominated by the circadian clock, and those that are dominated by light. C, Phase distribution of co-regulated, clock-dominated, and light-dominated transcripts in LD.

strongly influencing the time point of transcript peak expression. These findings suggest that the circadian oscillator has limited control over expression dynamics of circadian transcripts under conditions of pronounced *diel* oscillations in barley.

Whereas the number of cycling transcripts was not different among the *Hvelf3* and *Hvlux1* mutants and cv Bowman, we observed quantitative variation in the phase distribution under *diel* conditions between the three genotypes. HvELF3 altered the timing of transcript oscillations in day/night cycles by suppressing expression at the light and dark interfaces. This effect was apparently completely or partially independent of the oscillator defect that causes arrhythmia in the *Hvelf3* plants under LL. Loss of *HvELF3*, but not of *HvLUX1*, altered the expression phase under *diel* cycles, although both mutants had a disrupted circadian clock. Therefore, HvELF3 modifies the light- and temperature-controlled *diel* transcriptome oscillations in barley. The role of HvELF3 in mediating light and temperature cues is supported by the loss of photoperiod sensitivity in the *Hvelf3* mutant (Faure et al., 2012) and resembles the role of ELF3 in antagonizing light input to the clock during the night in *Arabidopsis* (McWatters et al., 2000; Covington et al., 2001; Thines and Harmon, 2010). However, our experimental setup with light and temperature covarying between day and night did not allow the detection of light- or temperature-specific effects on the *diel* transcriptome. It has already been shown that photic versus thermal entrainment results in a different behavior of the *Arabidopsis* circadian clock (Boikoglou et al., 2011). Future studies on the barley circadian clock should therefore test for the effects of clock genes on the circadian and *diel* transcriptomes under thermal versus photic entrainment.

It is interesting that only HvELF3 but not HvLUX1 had strong effects on the *diel* transcriptome, because out of the *Arabidopsis* core components of the EC ELF3–ELF4–LUX, only LUX has been identified as a transcription factor with direct DNA binding activity (Helfer et al., 2011). The different effects observed in *Hvelf3* and *Hvlux1* regarding the *diel* transcriptome may also be caused by the different nature of the underlying mutations. Whereas the *Hvelf3* mutant line carries a premature stop codon leading to a truncated HvELF3 protein, the *Hvlux1* mutant is characterized by a single amino acid exchange in the Myb-domain, which is important for the binding to cognate DNA sequences and regulation of their target genes (Faure et al., 2012; Campoli et al., 2013). In addition, a paralogue of *HvLUX1* termed *HvLUX2* may have partly redundant function and might have compensated for the mutation in *HvLUX1* (Pankin et al., 2014). On the other hand, chromatin immunoprecipitation experiments demonstrated that ELF3 had many more significant binding sites than LUX, suggesting that ELF3 also binds independently of LUX (Ezer et al., 2017). Therefore, HvELF3 and HvLUX1 might have independent targets in the barley genome.

Based on RNA time series data, we modeled a possible barley clock as a basis for understanding its effects

on physiology, metabolism, and agronomic performance. It is important to emphasize that the resulting interactions between the individual components of the clock represent one of the possible solutions of the barley circadian clock circuit, which may serve as a null model in future studies aimed to experimentally resolve composition and regulation of this clock. We used simple dynamical models to capture gene regulatory dynamics without making a priori assumptions on the structure of the network. These dynamical models have been successfully used in the past to describe circadian processes of *Arabidopsis* under conditions that are similar to those of our dataset (Dalchau et al., 2010; Herrero et al., 2012; Trejo Banos et al., 2015; Mombaerts et al., 2016, 2019). It is also important to stress that our approach could only model genes with circadian expression oscillations, whereas it is well known that post-transcriptional regulation and the rate of protein degradation and activity is an essential constituent of the clock mechanism in *Arabidopsis* (Kim et al., 2003; Más et al., 2003). For example, the expression of the important circadian oscillator component ZTL is not rhythmic (Somers et al., 2000), but ZTL protein is post-translationally regulated by light and its function is modified by GI at specific times of the day (Kim et al., 2007).

Our modeling strategy used *HvLUX1* to reveal the circadian circuitry, which therefore appeared as a major hub in the barley clock. Nevertheless, this predicted central role of *HvLUX1* is consistent with the loss of self-sustained rhythms in the *Hvlux1* mutant. Unlike *HvELF3* and *HvELF4*, *HvLUX1* encodes a protein with a known DNA-binding domain, suggesting that the transcriptional regulation of the EC converges on *HvLUX1* (Nusinow et al., 2011). Our model predicted that *HvLUX1* represses *HvGI* and is itself repressed by HvLHY, consistent with the suggested repression of *HvGI* by the EC and CCA1/LHY repressing the EC in *Arabidopsis* (Hsu et al., 2013; Fogelmark and Troein, 2014).

Further, the regulatory predictions suggested that HvLHY and HVRVE8 are activators of *HvPRR73* and *HvPRR95* in the morning and, at the same time, repress *HvLUX1*. The morning activation of the *HvPRRs* through HvLHY and HVRVE8, together with the repression of *HvLHY* and *HvREV8* through the *HvPRRs* later in the day, are also key regulatory relationships within the *Arabidopsis* clock (Hsu et al., 2013; Fogelmark and Troein, 2014). This suggests that the regulatory links among HvLHY, HVRVE8, and the *HvPRRs* are conserved between barley and *Arabidopsis*, despite the independent evolutionary history of LHY-like and PRR-like genes in the barley and *Arabidopsis* clades (Takata et al., 2009, 2010; Campoli et al., 2012).

Our model suggested that *HvPRR73*, the first PRR expressed in barley in the morning, activates *HvPRR95*, which, in turn, activates *HvPRR59*, such that *HvPRR73*, *HvPRR95*, and *HvPRR59* are expressed in a sequential cascade. This resembles predictions by Pokhilko et al. (2012), who described the PRRs as a series of activators

in the Arabidopsis clock, whereas other models have predicted that direct interactions between the PRRs are negative and directed from the later PRRs in the sequence to the earlier ones (Huang et al., 2012; Carré and Veflingstad 2013; Fogelmark and Troein, 2014). However, the sequential regulation of the PRRs during the day appears to be a common feature of the circadian clock in both barley and Arabidopsis, whereas the sequence of expression of *PRR* genes is altered between Arabidopsis and barley. In Arabidopsis, the sequence of *PRR* expression starts with *PRR9*, followed by *PRR7*, *PRR5*, and *PRR3*, and ends with *PRR1* (Matsushika et al., 2000; Hsu et al., 2013). However, in our data the sequential *PRR* expression wave started with *PRR73* and ended with *PRR59* (*PRR1* was not scored rhythmic in our data). Interestingly, *PRR37*, corresponding to the major photoperiod response gene *PPD1* in wheat and barley, showed no circadian oscillations and is therefore probably not part of the circadian clock in barley. This is consistent with the finding that mutations in *PPD1* do not affect the circadian clock in barley and wheat (Campoli et al., 2012; Shaw et al., 2012).

Our modeling placed several members of the RVE family, including barley homologs of the principal clock activators RVE8, RVE6, and RVE4, into the barley clock (Kuno et al., 2003; Zhang et al., 2007; Rawat et al., 2009). The mutation in *eam7* had a limited effect on the central oscillator, but several RVE homologs lost rhythmicity or were strongly downregulated and this correlated with a lengthening of the period. The suggested that *EAM7* is a component of the slave oscillator that only regulates a subset of clock-controlled transcripts. *RVE*-like genes have been implicated in such slave oscillators (Kuno et al., 2003) and *roe* mutants are characterized by a period lengthening (Hsu et al., 2013); however, none of the expressed *RVE* genes carried mutations that would alter the protein sequence.

BBX19, RVE7, and HAM3 were identified as three new candidate oscillator components in barley. Whereas the three proteins have already been proposed to have connections to the Arabidopsis oscillator, they have not been modeled as an integral part of the circadian clock but rather as clock outputs in Arabidopsis (Kuno et al., 2003; Wang et al., 2014). BBX19 acts as a gatekeeper of EC formation by mediating degradation of ELF3 and is part of a regulatory loop with CCA1 and/or LHY (Wang et al., 2015; Tripathi et al., 2017), supporting the link between BBX19 and LHY in our model. The model also predicted that barley homologs of BT2, CYTOCHROME 450, and PHOSPHATE STARVATION RESPONSE1 are part of the core circadian oscillator in barley. However, these proteins were only predicted to regulate other clock components and were not regulated themselves by the clock. They therefore displayed a low connectivity within the circadian network, consistent with their known functions outside the central clock (Bari et al., 2006; Ren et al., 2007; Bak et al., 2011). Therefore, these components might provide input into the circadian network but are probably not components of the barley oscillator. However, the function of the predicted barley

clock components, their role in the barley circadian clock, and interactions generated by the network modeling need to be experimentally verified using natural and induced mutants and transgenic lines.

## CONCLUSION

Our comparison of *diel* and circadian transcriptomes in the different barley clock mutants revealed that fluctuations of light and temperature have a major effect on the *diel* oscillating transcriptome and can compensate for circadian defects in arrhythmic barley clock mutants. HvELF3, but not HvLUX1, controlled the expression phase of a large number of transcripts under *diel* conditions and this effect was independent from the oscillator arrest under LL. Dynamical modeling suggested novel putative clock genes and connections between clock genes as a basis for experimental explorations into the nature and functions of the barley circadian clock. Finally, our findings and the dataset provide a valuable resource for mining for the output targets of the barley clock components HvELF3, HvLUX1, and EAM7 and to understand the role of the *diel* cues and clock in controlling the barley transcriptome and plant performance.

## MATERIALS AND METHODS

### Genetic Material and Growth Conditions

Four spring barley (*Hordeum vulgare*) genotypes were used in this study, the wild-type spring barley 'Bowman' (wild type) and three derived introgression lines with mutations in *HvELF3* (BW290), *HvLUX1* (BW284), and *EAM7* (BW287; Faure et al., 2012; Campoli et al., 2013). BW290 carries an introgression of the *eam8* allele (*eam8.k*) that is characterized by a bp mutation leading to a premature stop codon in *HvELF3*, which is orthologous to *ELF3* in Arabidopsis (*Arabidopsis thaliana*; Faure et al., 2012; Hicks et al., 2001). BW284 carries the *eam10* locus characterized by a single nonsynonymous nucleotide polymorphism in the conserved Myb-domain of the barley LUX/ARRHYTHMO homolog (Campoli et al., 2013). The candidate gene for *EAM7* has not yet been identified, but *eam7* is characterized by accelerated flowering and reduced sensitivity to the photoperiod (Stracke and Börner, 1998). The lines were sown in soil (Einheitserde) in a 96-well format. Seeds were maintained at 4°C for 3 d, followed by germination in 12-h light/12-h dark cycle at 20°C with a photon flux density 300  $\mu\text{mol m}^{-2} \text{s}^{-1}$  during the day and 18°C during the night, and grown for three weeks.

### Sampling, Extraction of Total RNA, and Sequencing

After three weeks of growth, the second expanded leaf after the cotyledon leaf was harvested from two plants, resulting in two biological replicates per genotype and time point every 4 h for 24 h starting with lights off, except for dusk and dawn, when extra samples were taken for cv Bowman every 2 h (Supplemental Fig. S1). After completion of a 12-h dark/12-h light *diel* cycle, the growth chamber was switched to constant light and 20°C and sampling continued for an additional 36 h starting from the first subjective dusk. Total RNA was extracted from ground tissue using a hybrid protocol of TRIZOL (Invitrogen) and purification columns from a RNeasy RNA extraction kit (Qiagen). Extracted total RNA was DNase-treated (Ambion). The concentration and integrity of the extracted RNA was determined on a BioAnalyzer (Agilent) before library preparation. The library preparation was carried out following the TruSeq protocol and single-end sequenced on a HiSeq 2500 System (Illumina) with 10 million reads per library.

## Mapping of Reads and Rhythmicity Analysis

The quality of the sequencing data were verified using the software FastQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>). The reads were mapped against a custom barley reference transcriptome (Digel et al., 2015) and raw read counts were obtained using the software implementing the full pipeline for RNA-seq analysis RobiNA (v1.2.3) with default settings (Lohse et al., 2012). Raw read counts were normalized to CPM with the R/Bioconductor package edgeR (<http://bioinf.wehi.edu.au/edgeR/>) and used for the downstream rhythmic analysis and modeling. To cross reference a nomenclature of the gene models used in this study (Digel et al., 2015) with the identifiers of the HORVU gene models annotated on the barley pseudochromosomes (Mascher et al., 2017), we used reciprocal BLASTn v2.9.0+ (e value cutoff 10e-05). The reciprocal best BLAST (<https://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/>) hit pairs were extracted as matching gene model identifiers.

For the analysis of the day/night data, the sequence of samples was inverted to start with the night followed by the day samples (12, 14, 16, 20, 0, 2, 4, and 8 h), and this data set was therefore termed "LD." The oscillating patterns of gene expression and period (as a duration of one complete cycle) and phase (as the location of time of the peak of the curve) of the curves were determined using the ARSER algorithm in the R package, MetaCycle (Yang and Su, 2010; Wu et al., 2016). To increase the analytical power for the rhythmic analysis, the 24-h *diel* dataset, but not the 36-h LL data, was duplicated to imitate 48 h of sampling data. The settings for the ARSER algorithm were adjusted for each genotype individually so that the period was normally distributed with two symmetric tails and the number of transcripts passing the cutoff (Benjamini-Hochberg corrected false discovery rate of 0.1; Benjamini and Hochberg, 1995) was maximal with the given range of period estimation being minimal. For cv Bowman and BW287 the settings are shown in Figure 2, B and C, and comprised mean = 28, upper/lower limit = 22/34 for the constant light data. In *diel* cycles, the mean and upper/lower limits were set to 24 and 21/27, respectively. Significant differences between the period lengths of transcripts in cv Bowman and *eam7* were tested with a one-factorial ANOVA ( $P < 0.05$ ).

## Modeling the Barley Circadian Clock

To identify putative barley clock genes, we blasted all expressed barley transcripts against the Arabidopsis transcriptome (geneontology.org). For each barley transcript, the best Arabidopsis BLAST hit was retained, and we then selected only those barley transcripts for which the Arabidopsis homolog carried the GO annotation "circadian" and "transcription" (Supplemental Dataset S1). The resulting 138 barley transcripts were then further filtered for those that exhibited unambiguous dynamics and a high signal-to-noise ratio of expression. This filtering step was necessary to ensure that we did not identify dynamics out of noise. Hence, genes for which the amplitude of oscillation was <20 CPM in the last 24 h were removed. The choice of such a filter is motivated by both the transitional nature of constant light data, which typically shows a large decrease of amplitude after few hours in barley, and the dependency of noise on gene expression levels. Furthermore, genes that were constantly up-/downregulated without exhibiting further significant dynamics were also discarded. This was performed by detrending the final 24 h of constant light data before applying the same filtering criterion. After filtering, out of a total of 138, 49 and 48 genes passed the filtering criteria in the wild-type and BW287 datasets, respectively. BW284 (*Hvlux1*) and BW290 (*Hvelf3*) datasets were not considered in the following network analysis because these mutations led to the arrhythmic transcriptomes. Finally, seven genes (*Hv.21080*, *Hv.22191*, *Hv.23289*, *Hv.32914*, *Hv.33010*, *Hv.6793*, and *MLOC\_7084.3*) were manually discarded from both subset lists of candidates as they were not DNA-binding transcription factors but rather enzymes in a metabolic process, leaving 42 in cv Bowman and 41 in BW287, of which 35 transcripts were in common between the cv Bowman and BW287 final datasets used for modeling. The *HvELF3* transcript did not pass the filtering and, therefore, could not be used to infer dynamical interactions.

To model the barley circadian clock based on the time-course gene transcription data, we adopted an approach based on LTI models. LTI models do not rely on prior knowledge of the transcriptional network to provide accurate predictions and have been shown to provide reliable predictions of the dynamical processes involved in the Arabidopsis circadian network (Dalchau, 2012; Herrero et al., 2012; Mombaerts et al., 2016, 2019). To provide a comprehensive evaluation of the LTI model, the performance of the modeling strategy was evaluated and compared under conditions that replicated those of

the experiments using widely used benchmark models (Supplemental Information).

We used first-order models to represent the system dynamics between two genes at a time using an LTI model with the following equation:

$$\frac{dy(t)}{dt} = au(t) - by(t) + c \quad (1)$$

where  $by(t)$  corresponds to the degradation rate of  $y$  and  $au(t)$  represents the influence of another transcription factor through the synthesis rate of  $y$ . The model, therefore, evaluates whether the rate of change of a particular gene  $y$  depends on another gene  $u$ . Estimating a model means finding (a), (b), and (c) that produce a vector  $y(t)$  as close as possible to the real data. The estimation of parameters was performed using the function "pem" implemented in the software MATLAB (Mathworks; <https://www.mathworks.com/>) that minimizes the prediction error of the data. LL data were used for the estimation, as they represent the autonomous behavior of the oscillator.

The goodness-of-fit of the model, with the data, was calculated as follows:

$$fitness = 100 * \left( 1 - \frac{\sum_{k=1}^N \sqrt{(y_k - \hat{y}_k)^2}}{\sum_{k=1}^N \sqrt{(y_k - \bar{y})^2}} \right) \quad (2)$$

where  $y_k$  is the data (output),  $\bar{y}$  is the average value of the data, and  $\hat{y}_k$  is the estimated output. The MATLAB function *compare* was used to compute the fitness of the model. Each potential link between two genes was validated if the associated model reproduced the dynamics involved with a sufficient degree of precision, which corresponds to a fitness threshold estimated at 60% (Supplemental Information).

To investigate the potential regulators of *HvLUX1*, a collection of independent first-order LTI models was estimated separately between each of the transcripts and *HvLUX1* in the cv Bowman background. In each case, the parameters were estimated so that they together provide the best possible fit to the *HvLUX1* time-course data. This step takes the following form:

$$\begin{aligned} \frac{d[LUX]_t}{dt} &= a_1 u_1(t) - b_1 [LUX]_t + c_1 \\ &\dots \\ \frac{d[LUX]_t}{dt} &= a_n u_n(t) - b_n [LUX]_t + c_n \end{aligned} \quad (3)$$

where  $n$  corresponds to the number of candidates (42 models in total). Each model was characterized by a fitness metric that ranges from 0% to 100%, representing its capability to describe the regulatory dynamics between genes. A gene, therefore, would be further considered as a regulator for *HvLUX1* if the model is capable of reproducing the shape of *HvLUX1* with a sufficient degree of precision. A fitness threshold, evaluated from in silico benchmarks systems (Supplemental Information), was used to validate the models. In this case, the fitness threshold was set to 60% to limit false-positive predictions of regulatory interactions while accounting for sufficient gene regulatory models to describe the system of interest. Finally, 20 models passed the validation step (Supplemental Dataset S5). The methodology is summarized in Supplemental Figure S4.

To further narrow down the predicted regulatory interactions, we estimated the consistency of the candidate models using the filtered *eam7* (BW287) dataset. For this purpose, we evaluated first-order LTI models for each of the previously identified regulations and retained those with the goodness-of-fit > 60% in the *eam7* (BW287) experimental condition (Supplemental Fig. S4). To keep links with the highest confidence only, the dynamical consistency of the LTI models based on these two independent datasets (cv Bowman and BW287) was evaluated using the nu ( $\nu$ ) gap metric (*gapmetric*, MATLAB; Vinnicombe, 1993; Mombaerts et al., 2019). We further considered models that had a  $\nu$ -gap < 0.2 following Carignano et al. (2015). As a result, six regulatory interactions were filtered out (Hv.10528 to Hv.27754, Hv.1530 [GI] to Hv.19411, Hv.19411 to Hv.20312 [LUX], Hv.19759 [TOC1] to Hv.20312 [LUX], Hv.5253 [LHY] to Hv.27754, and Hv.9855 to Hv.18813 [PRR59]). HvPRR95 (Hv.4918) appeared as a hub with eight connections, so we repeated the search for regulators of HvPRR95 (Eq. 3), computed their interactions in both datasets, and checked their consistency.

The relative contribution of light-signaling and circadian-clock pathways in generating oscillating transcriptome was evaluated using a Bode analysis (*bode* function in MATLAB) with the threshold of 7 dB to discern between the two alternative regulatory inputs (Dalchau et al., 2010; Supplemental Fig. S5). Here,

we use the magnitude response of the signal to assess the relative contribution of the inputs  $u_{\text{light}}(t)$  and  $u_{\text{LHY}}(t)$  in each of the validated models, at a frequency of 24 h (or  $0.262 \text{ rad h}^{-1}$ ). Following Dalchau et al. (2010), if the magnitude of the response of the light input was 7-dB higher than the contribution of the clock (represented by *HvLHY* potentially delayed), the circadian-regulated gene (the output of the model) was considered driven mostly by light; in the opposite case, the transcript expression was considered as driven by the clock. If the magnitude difference was <7 dB, then the circadian-regulated gene was considered regulated by both inputs equally. The methodology is summarized in Supplemental Figure S7B.

To this end, we used the 2,759 transcripts that were identified as oscillating in both *diel* and free-running conditions in the wild-type cv Bowman background to calculate another set of LTI models as described earlier. As a reference, we selected a formerly identified clock gene peaking in the morning, *HvLHY* (Hv.5253), with a range of delays integrated into the model to implicitly represent the clock input following Dalchau et al. (2010). The structure of such models is schematically represented in Supplemental Figure S7A. This way, the light input is incorporated on two levels: explicitly through the light input and implicitly through the clock pathway using the following equation:

$$\frac{dy(t)}{dt} = a_1 u_{\text{light}}(t - \mu_{\text{light}}) + a_2 u_{\text{LHY}}(t - \mu_{\text{LHY}}) - by(t) \quad (4)$$

where  $u_{\text{light}}$  was assumed to be binary (1 = light; 0 = dark). We fixed the light delay  $\mu_{\text{light}}$  to 0 h to represent the effect of rapid light signaling on the transcripts, and computed delays ranging from 0 to 8 h, every 0.2 h, for *HvLHY*. The delay that provided the best fit to the data was selected independently for each transcript. Ultimately, models were validated if they succeeded in capturing the regulatory dynamics involved with a goodness-of-fit > 60%.

We assessed the accuracy of our LTI-based network reconstruction algorithm on the circadian model from Pokhilko et al. (2010) as detailed in the Supplemental Information. The performance of the modeling strategy was evaluated based on the Area Under the Curve of the Receiver Operating Characteristic and Area Under the Precision-Recall Curve criteria (Supplemental Information; Supplemental Fig. S8).

The scripts for the modeling of the barley clock are available under: <https://github.com/Lmombaerts/CircadianBarley>.

## Accession Number

The ArrayExpress accession no. E-MTAB-8372 was deposited in the European Molecular Biology Laboratory data library (<https://www.ebi.ac.uk/arrayexpress/>). Accession numbers of genes are given in Supplemental Dataset S1.

## Supplemental Data

The following supplemental materials are available.

**Supplemental Figure S1.** Sampling scheme for the LD and LL samples in Bowman, BW284 (*Hvlux1*), BW287 (*eam7*), and BW290 (*Hvelf3*) grown under 12-h light/12-h dark cycles (20°C day/16°C night).

**Supplemental Figure S2.** Transcriptome-wide period estimates under diel day/night conditions.

**Supplemental Figure S3.** Phase correlation of transcripts of cv Bowman with the clock mutants *Hvelf3*, *Hvlux1*, and *eam7* grown under LD conditions.

**Supplemental Figure S4.** Schematic of the strategy used in this article to investigate the potential dependencies of HvLUX1 through identification of dynamics.

**Supplemental Figure S5.** Predicted regulatory interactions between the regulators of HvLUX1 that have been identified in cv Bowman and further validated using the *eam7* dataset.

**Supplemental Figure S6.** Expression profiles of predicted core clock components of the barley oscillator in free-running conditions of constant light.

**Supplemental Figure S7.** Methodology and examples of the Bode analysis.

**Supplemental Figure S8.** Performance of the modeling strategy compared to other clock models.

**Supplemental Dataset S1.** CPM for transcripts and information about rhythmicity, period, amplitude, and phase, as well as gene annotation.

**Supplemental Dataset S2.** Oscillating barley transcripts homologous to Arabidopsis gene models with the annotations “circadian” and “transcription.”

**Supplemental Dataset S3.** Transcripts with unambiguous dynamics and a high signal-to-noise ratio of expression in cv Bowman and BW287 (*eam7*).

**Supplemental Dataset S4.** Predicted 49 regulatory links between 15 genes in the circadian oscillator network in barley (Fig. 4).

**Supplemental Dataset S5.** LTI model parameters of the 20 models that passed the validation step (>60% fitness).

**Supplemental Information.** Benchmarking models.

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