




ORIGINAL ARTICLE OPEN ACCESS

# GIGANTEA Is Required for Robust Circadian Rhythms in Wheat

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## ABSTRACT

*GIGANTEA* (*GI*) is a plant-specific protein that functions in many physiological processes and signalling networks. In Arabidopsis, *GI* has a central role in circadian oscillators regulating the abundance of ZEITLUPE and TIMING OF CAB EXPRESSION 1 proteins and is essential for photoperiodic regulation of flowering. We have investigated how orthologues of this component of Arabidopsis circadian oscillators contribute to circadian rhythms and yield traits, including heading (flowering) in wheat. We find that *GI* is a core component of wheat circadian oscillators that is necessary to maintain robust oscillations in chlorophyll fluorescence and circadian oscillator transcript abundance. The predicted lack of functional *GI* results in later flowering of wheat in both long days and short days in controlled environment conditions. Our results support and extend previous work, which suggests that the pathways by which photoperiodism regulates flowering are not fully conserved between Arabidopsis and wheat. Understanding the molecular basis for photoperiodism in wheat is important for breeders looking to manipulate flowering time and develop new elite, high-yielding cultivars.

## 1 | Introduction

Circadian oscillators are endogenous timing mechanisms that act to time internal processes to daily and seasonal cycles in their environment. In the model plant Arabidopsis, well-defined transcription-translation feedback loops result in sequential expression of circadian oscillator genes across the 24-h cycle. At dawn, *CIRCADIAN CLOCK ASSOCIATED 1* (*CCA1*) and *LATE ELONGATED HYPOCOTYL* (*LHY*) are expressed and repress the day phased *PSEUDO RESPONSE REGULATOR* (*PRR*) genes (*PRR7*, *PRR9* and *TIMING OF CAB EXPRESSION 1*, *TOC1*)

(Harmer et al. 2000; Adams et al. 2015). REVILLE (*RVE*)/LIGHT NIGHT INDUCIBLE AND CLOCK REGULATED 1 (*LNK1*) complex promotes *PRR* expression (Rawat et al. 2011; Xie et al. 2014), which feeds back to repress *CCA1/LHY* (Nakamichi et al. 2010). The evening complex (*EARLY FLOWERING 3/EARLY FLOWERING 4/LUX ARRHYTHMO*) represses *PRR* expression at dusk (Dixon et al. 2011; Helfer et al. 2011; Nusinow et al. 2011). *PRR* action is further restricted to the light by ZEITLUPE (*ZTL*)-mediated degradation of *PRR* proteins in the dark (Cha et al. 2017; Lee et al. 2019). Thus, *CCA1/LHY* repression is lifted towards the new dawn and the

**Abbreviations:** *CCA1*, *CIRCADIAN CLOCK ASSOCIATED 1*; CF, chlorophyll fluorescence; CO, CONSTANS; *FKF1*, FLAVIN BINDING KELCH REPEAT 1; FT, FLOWERING LOCUS T; *GI*, *GIGANTEA*; GS55, growth stage 55; LD, light-dark; *LHY*, *LATE ELONGATED HYPOCOTYL*; LL, continuous light; NIAB, National Institute of Agricultural Botany; NPQ, non-photochemical quenching; *PPD-1*, *Photoperiod-1*; *PRR*, *PSEUDO RESPONSE REGULATOR*; RAE, relative amplitude error; *TOC1*, *TIMING OF CAB EXPRESSION 1*; WT segregant, wild type; *ZTL*, ZEITLUPE.

Matthew A. Hannah and Alex A. R. Webb contributed equally to this work and share the last authorship.

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cycle repeats. In Arabidopsis, proteins that might function as scaffolds have profound effects on the circadian oscillator. For example, members of the WD40 repeat family of scaffold proteins are essential for circadian rhythms (Airoldi et al. 2019). GIGANTEA (GI) is another potential scaffold protein that is important in circadian timing. GI regulates ZTL stability and activity through light-dependent binding and affects deubiquitination and also translational activity (Lee et al. 2019). Mutations in *GI* have allele-specific effects on Arabidopsis circadian oscillators but are mostly considered to result in faster running circadian oscillators and therefore shorter circadian periods in constant conditions (Hsu and Harmer 2014). The architecture of circadian oscillators is broadly conserved across the plant kingdom, with alleles of circadian oscillator genes selected during domestication and in breeding programs to enhance agriculturally important crop traits, such as flowering time (McClung 2021; Steed et al. 2021). However, we have found in wheat that transcripts of *ELF3*, which encodes a putative scaffold protein, peak at dawn, rather than dusk as in Arabidopsis (Wittern et al. 2023). Furthermore, wheat *ELF3* proteins are unstable in the light (Alvarez et al. 2023). These data suggest that there are differences in circadian oscillator structure and function in wheat compared to Arabidopsis and warrant further investigation.

In Arabidopsis, mutations in *GI* also have profound effects on the photoperiodic flowering pathway due to both its function in regulating circadian oscillators and more direct roles in regulating the floral induction mechanisms (Fowler 1999). In the external coincidence model of photoperiodism, flowering is induced when external light is coincident with an appropriate phase of the circadian rhythm (Bunning 1960; Pittendrigh and Minis 1964). *GI* connects circadian oscillators and the photoperiodic flowering pathway, and along with *FLAVIN BINDING KELCH REPEAT 1 (FKF1)* and *CONSTANS (CO)* are the molecular components which determine the light-sensitive and -insensitive phase of the circadian cycle for the regulation of flowering (Suárez-López et al. 2001; Valverde et al. 2004; Sawa et al. 2007). In long photoperiods, *GI* and *FKF1* peak expression coincide in the late afternoon, forming a complex which promotes *CO* expression. In Arabidopsis, if light is coincident with the accumulation of *CO* transcripts, *CO* protein is stabilised to pass the threshold necessary to trigger *FLOWERING LOCUS T (FT)* expression (Imaizumi et al. 2003; Sawa et al. 2007; Fornara et al. 2009). Arabidopsis *FT* is a mobile signal which travels from the leaf to the apical meristem to trigger floral development (Corbesier et al. 2007). In short photoperiods, *FKF1* and *GI* maximal expression do not coincide, hence *CO* repression is not lifted during the day and maximal *CO* accumulation occurs at night, outside of the light-sensitive phase of the rhythm (Sawa et al. 2007). Therefore, *GI* mutants such as *gi-1* (premature stop coding resulting in the loss of 171 of the C-terminal domain), *gi-2* (causing the deletion of all but 142 amino acids of the N-terminal domain and addition of the 16 new amino acids in the C-terminal domain) and T-DNA insertion line (*gi-11*, deletion in the 5' half of *GI* caused by the insertion of the T-DNA) are late flowering in long photoperiods but have no, or minor delays to flowering when cultivated in short photoperiods (Fowler 1999; Park et al. 1999).

In wheat, photoperiodic-dependent regulation of flowering is heavily dependent on *Photoperiod 1 (PPD-1)*, an orthologue of

Arabidopsis *PRR3* or *PRR7* (Turner et al. 2005; Beales et al. 2007; Wilhelm et al. 2009). *PPD-1* induces *FT1* expression in leaves activating the transition to the reproductive stage (Alvarez et al. 2023). Analyses of the tetraploid wheat loss of function Target Induced Local Lesions In Genome (TILLING) lines containing stop codon and splice site mutations have demonstrated that *PPD-1* and *CO1/CO2* act in separate but highly connected flowering pathways (Shaw et al. 2020). In tetraploid wheat, light activation of *PPD-1*, mediated by phytochromes *PHYB* and *PHYC*, is facilitated by *ELF3*, which represses the expression of *PPD-1* by directly binding to its promoter (Alvarez et al. 2023). *ELF3* controls the abundance of *PPD-1* to regulate flowering, somewhat independent of its role in circadian oscillators in tetraploid wheat (Wittern et al. 2023). Tetraploid wheat plants carrying a deletion in the promoter of the A homoeologue of *PPD-1 (Ppd-1Aa)* are photoperiod-insensitive (PI) and are early heading under short photoperiods. In contrast, plants that carry the *Ppd-1Ab* allele are photoperiod-sensitive heading later under short photoperiods (Alvarez et al. 2023).

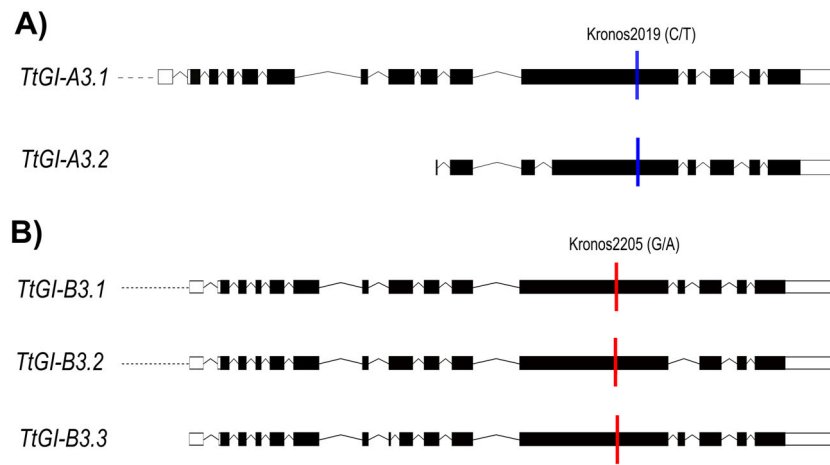
In wheat, *GI* regulates flowering though the mechanisms are not yet resolved. Association studies using wheat lines from diverse geographical regions found that polymorphisms in *GI* are associated with earlier flowering (Rousset et al. 2011). In both wheat and barley, *GI* expression levels peak in the afternoon in long and short photoperiods (Dunford et al. 2005; Zhao et al. 2005; Rees et al. 2022). Wheat *GI* rescues the late flowering phenotype in Arabidopsis *gi-2* with overexpression of the wheat *GI* in Arabidopsis resulting in earlier flowering (Zhao et al. 2005). *GI* loss-of-function mutants in a photoperiod-sensitive wheat background have delayed heading dates in both long and short photoperiods. The stronger effects of *GI* on heading date in long photoperiods required *PPD-1* or *ELF3*, suggesting function in a common pathway (Li et al. 2024).

To investigate the relationship between the function of the circadian oscillator in the regulation of heading, we isolated wheat TILLING lines which carry predicted nonfunctional alleles of the Durum wheat orthologues of Arabidopsis *GI*. We demonstrate that *GI* is required for robust circadian oscillations in Durum wheat. We then investigated whether *GI* functions in the Durum wheat photoperiodic flowering pathway independent of normal *PPD-1* function. Finally, we investigated if lack of functional *GI* affects other aspects related to yield traits.

## 2 | Materials and Methods

### 2.1 | Isolation of *GI* TILLING Mutant Lines in Tetraploid Wheat

To investigate the role of *GI* in wheat, we isolated TILLING mutants in the tetraploid wheat variety *T. turgidum cv. Kronos*. Two TILLING lines with premature stop codons within the A *GI* homoeologues (*GI-A*, TraesCS3A02G116300, Kronos2019) and B *GI* homoeologues (*GI-B*, TraesCS3B02G135400, Kronos2205) genes were obtained from the Germplasm Resource Unit, Norwich UK (Figure 1). BLAST search of the sequenced Kronos TILLING lines for *TtGI* identified pairs of candidate lines with deleterious mutations in each of the subgenomes for *TtGI*, which



**FIGURE 1** | Location of TILLING mutations in the *GI-A3* and *GI-B3* homeologues. Location of premature stop codons (coloured bars) in (A) *GI-A3* and (B) *GI-B3* homeologues (showing their respective splice variants, genetic alterations resulting in an altered coding sequence). Bars represent exons with black and white showing those that are translated and untranslated, respectively. Kronos2019 and Kronos2205 are the original TILLING lines obtained from the GRU. Schematic was adapted from EnsemblPlants database. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

were obtained directly from Dr. Cristobal Uauy (John Innes Centre, Norwich). Further candidates from tetraploid Kronos were identified using a pre-publication version of the <http://www.wheat-tilling.com> website (Krasileva et al. 2017). Kronos2019 contains a C to T SNP 6606 nucleotide base pairs (bp) from the start of the *GI-A3* transcription start site (TSS) (position 3A:84190982 IWGSC RefSeq V1.1). This SNP is situated within the 12th exon of the first splice variant and the fourth exon of the second splice variant and is predicted to result in the deletion of 356 amino acids (Figure 1A). Kronos2205 contains a G to A SNP 6348 bp downstream of the *GI-B3* TSS (position 3B:117929486 IWGSC RefSeq V1.1). This SNP is situated within the 12th exon of the first two splice variants and the 11th exon of the third splice variant and is predicted to result in the deletion of 403 amino acids (Figure 1B). Abundance of *TtGI* at ZT12 and ZT15 is not significantly different between the WT genotypes, single and double mutants, with both homologous expressing at the same level (Figure S1).

Kronos TILLING lines were backcrossed four times to Kronos (Figure S2). The BC4 F1 heterozygous plants were self-crossed to produce BC4 F2 seed. BC4 F2 seeds were genotyped, and seeds were retained from plants that were homozygous for the Kronos2019 or Kronos2205 mutations. The homozygous Kronos2019 (*Ttgi-A3*) and Kronos2205 (*Ttgi-B3*) lines were crossed, self-crossed and plants homozygous for both Kronos2019 and Kronos2205 mutations (*Ttgi-A3/gi-B3*) were selected. To control for possible background mutations, that are present in the TILLING populations, plants homozygous for both WT alleles were selected as mutant sibling WT (segregant). Kronos WT line was selected as non-mutagenised WT control.

## 2.2 | Genotyping

Genotyping was performed by PCR, followed by Sanger sequencing conducted by Source Bioscience Ltd (UK). A Faststart Taq Polymerase Kit (Roche, UK) was used for PCR following the manufacturer's instructions. PCR conditions were as follows: 4 min at 95°C; 40 cycles of 30 s at 95°C, 30 s at 68°C/

69°C (*Ttgi-A3* and *Ttgi-B3* respectively), 60 s at 72°C; 10 min at 72°C. Primers used for the A homologue: forward 5' GCATCATCCCTCTACCATTTGA 3' and reverse 5' GTACAAGCTTCCACCGTCGA 3'. Primers used for the B homologue: forward 5'GTTGGTACCTCCTGGAAGTACA 3' and reverse 5'CATCCCATCTGTAGCACGAAGTA 3'.

## 2.3 | Plant Growth Conditions

Seeds were sown directly into modular trays containing a 5:1 mix of Levington M2 potting compost pre-treated with Intercept 70 W (0.02 gL<sup>-1</sup>, Bayer) and fine vermiculite. Seeds were stratified for 48 h at 4°C and then moved into a growth cabinet (PGR14, Conviron) or room (Conviron) fitted with broad-spectrum LED white light. If plants were to grow to maturity, seedlings were transplanted into 12 by 12 by 13 cm pots containing an identical soil mix after 2 weeks of growth. Plants were grown under 400 μmol m<sup>-2</sup> s<sup>-1</sup> PAR, 22°C white light/18°C dark in either long (16 h light/8 h dark) or short (8 h light/16 h dark) photoperiods.

## 2.4 | Chlorophyll Fluorescence Measurements

Chlorophyll fluorescence (CF) measurements were performed as outlined in Wittern et al. (2023). A 5 mm by 5 mm leaf fragment was placed into a well of a black 96 well plate (Greiner) containing 0.8% (w/v) bactoagar (BD), ½ MS (Duchefa Biochemie) and 0.5 μM 6-benzyl aminopurine (Sigma), adjusted to pH5.7 using 0.5 M KOH (Sigma).

CF imaging and processing were performed using a CFImager and accompanying software (Technologica Ltd. UK). CF images were captured using a Stingray F145B ASG camera (Allied Vision Technologies, UK) through an RG665 long pass filter to exclude blue light from the LEDs. The 'continuous light' protocol included the following steps: 40 min 100 μmol m<sup>-2</sup>s<sup>-1</sup> of blue light, 800 ms 6172 μmol m<sup>-2</sup>s<sup>-1</sup> saturating light pulse and 20 min of darkness. This protocol was repeated 120 times. The parameters non-photochemical quenching

(NPQ) and  $F_v/F_m$  (maximum potential quantum efficiency of Photosystem II, PSII) were used for circadian analysis. Relative amplitude error (RAE) calculation and period estimates were generated using Biodare2 (Zielinski et al. 2014).

## 2.5 | Reverse-Transcription Quantitative PCR

The *Ttgi-A3/gi-B3* and Kronos WT lines were sown and grown in a growth room under long-day conditions (16 h L: 8 h D; 250  $\mu\text{mol m}^{-2} \text{s}^{-1}$ , 20°C). *TtGI* expression was analysed from 14-day old plants at *TtGI* peak expression, ZT12 and ZT15. For analysis of circadian oscillator transcript abundance, samples were collected after 14 days of sowing from the first true leaf every 3 h for 96 h. During the first 24 h, sampling was done under long-day conditions (16 h L: 8 h D; 250  $\mu\text{mol m}^{-2} \text{s}^{-1}$ , 20°C), and then the cabinet was switched to continuous white light (250  $\mu\text{mol m}^{-2} \text{s}^{-1}$ ) and constant temperature (20°C). Flowering gene expression was analysed at the three-leaf growth stage (14 days after sowing) leaves and in GS39 growth stage (flag leaf blade all visible) under long-day conditions (16 h L: 8 h D; 250  $\mu\text{mol m}^{-2} \text{s}^{-1}$ , 20°C). Sampling commenced at time 0 for 24 h every 3 h.

Total RNA was extracted from leaves using the RNeasy Plant Mini Kit (Qiagen, UK) with an on-column DNase digest (Qiagen, UK), concentration and quality was determined using the Nanodrop ND-1000 (ThermoFischer scientific). cDNA was synthesised from 500 ng RNA using the RevertAid First Strand cDNA synthesis kit (Thermo Scientific, UK). Three technical replicates of gene-specific products were amplified in 10  $\mu\text{L}$  reactions using QuantiNova SYBR Green PCR Kit (Qiagen) on a CFX384 Touch Real-Time PCR detection system (Bio-Rad). Transcript levels were determined relative to the expression of two housekeeping genes, *RP15*, *RPT5A* and *Ta22845* as described in Wittern et al. (2023). Primer sequences are shown in Table 1.

## 2.6 | Plant Phenotyping (Laboratory)

Growth stage 55 (GS55), that is, when half of the ear emerged above flag leave ligule (Zadoks et al. 1974) was recorded daily. The first tiller to reach GS55 (primary tiller) per plant was marked. Post senescence, the height of each plant was measured using a metre rule and the total number of tillers and number of productive tillers were counted. The primary head (head from primary tiller) was collected, and the length, spikelet number, seed number and seed weight were recorded. Seed number and weight were then determined for the whole plant. Seed number was counted by hand, and seed weight was measured using a balance.

## 2.7 | Field Trial

Observational (1  $\text{m}^2$ ) plots of Kronos, WT segregant, *Ttgi-A3* and *Ttgi-A3/gi-B3* lines were grown at the National Institute of Agricultural Botany (NIAB, Cambridge, UK) experimental farm during the 2020 field season to provide preliminary data and bulk seed for a larger trial the following year. Environmental

**TABLE 1** | Primers used in this study.

Gene	Sequence 5' - 3'
<i>TtLHY</i>	F: CCTGGAATTGGAGATGGAGA R: TGAGCATGGCTTCTGATTTG
<i>TtELF3</i>	F: TCTCCAGATGATGTTGTGCGGT R: CTCGAACACTTGGACAGCAAA
<i>TtFT1</i>	F: TGAGGACCTTCTACACACTCG R: ACCGGGGATATCTGTCAACAAG
<i>TtGI</i>	F: GGTAGGTGATAGACGGCACTT R: GTGCTACAGATGGGATGCTTG F: ACAAGCGGTTTCGTGGAGG R: CCTGCATCCGCTTGACGTA
<i>TtPpd-1</i>	F: CCTGTGGACTGTGCATCTCAA R: CAAGGGATGGCAGCGATAATG
<i>TtPRR73</i>	F: TCCCGAAGTTCCTCTCTTTCC R: AGCGGTAGTGGAATGACA
<i>TtTOC1</i>	F: GGCATGGCACTTCATTCAAGTT R: GCACATTCATACCAGCAGGAC
<i>TtGID1 Li</i>	F: GGAGGAGGGGATCAAGATACAC R: CGATCTCCTCCATCACCTCG
<i>TtGA20ox2</i>	F: CTACGAGCCAATGGGGAG R: CCAGCAGCTCCATGATCCT
<i>TtRP15</i>	F: GCACACGTGCTTTCAGATAAG R: GCCCTCAAGCTCAACCATAACT
<i>TtRPT5A</i>	F: GCTGGCTCGTTCAACTGATG R: GGACCAAGCGTTCTGATTACTC
<i>Ta22845</i>	F: GCTGGCTCGTTCAACTGATG R: GGACCAAGCGTTCTGATTACTC
<i>TtGI</i>	F: GCTCTGGCATAAGCTTATTGCA R: TTCGCTGGTTGACTCTCCAC

data are included in Supporting Information S1. On 9 April 2021, Kronos, WT segregant, *Ttgi-A3* and *Ttgi-A3/gi-B3* yield plots (3.8 m x 2 m = 7.6  $\text{m}^2$ ) were drilled at NIAB following a randomised block design (Figure S3). The trial was flanked by two rows of Paragon to separate the *GI* field experiment from other trials located at the site. *Ttgi-B3* was not included due to insufficient number of seeds during the 2020 field season.

Five plants per plot were randomly tagged from the middle of each plot. The date each tagged plant reached GS55 was recorded. Post senescence, the height of each tagged plant was measured using a metre rule and the total number of tillers per tagged plant was counted. A representative head was taken from each tagged plant and the length, spikelet number, grain number and grain weight were recorded. Data were not recorded from Paragon control plants as the higher plant density and tillering meant that the tags were no longer visible within the Paragon plots. On 4 September, the plots were harvested, and the weight of grain plus the percentage of grain moisture for each plot was determined by the NIAB field team.

## 2.8 | Statistics

All statistical analysis was performed using R. For CF data sets, significant differences between the groups were tested using an ANOVA followed by a post hoc Tukey test. Statistics for phenotypic data collected from plants grown in the laboratory and tagged plants from the field were as follows. Continuous variables (plant height, primary head length, weight of seed) were tested using an ANOVA followed by a post hoc Tukey. All continuous-discrete variables (days till GS55, tiller number, seed number, spikelet number) were tested using a Kruskal–Wallis test followed by a post hoc Dunn test adjusted for multiple comparisons.

## 2.9 | Sequence Analysis

*TaGI* genomic sequences and CDS were retrieved from the reference genome sequence (IWGSC RefSeqv1.1) and submitted as query sequences to local Basic Local Alignment Search Tool server to identify *GI* in wheat cultivar genomes released by the 10+ genomes project (Walkowiak et al. 2020) (Table S1). A reciprocal BLAST approach was used to confirm ambiguous results. Alignments were created using CLC Genomics between the BLAST results, the genomic BLAST query sequence (IWGSC RefSeqv1.1) and the CDS and mRNA. SNPs and InDels within the CDS were manually annotated and counted.

## 3 | Results

### 3.1 | *TtGI* Is Required for Robust Circadian Rhythms Under Constant Light

To establish if *GI* contributes to the functioning of wheat circadian oscillators, we quantified circadian rhythms in chlorophyll *a* fluorescence from *Triticum turgidum* cv. *Kronos* by measuring the period and the RAE (measurement of the goodness of fit of the data to a cosine curve,  $RAE > 0.5$  usually indicates a lack of circadian rhythms) (Figure 2). Robust circadian oscillations of the chlorophyll *a* fluorescence derived parameter reporting NPQ were measured in the recurrent parent and WT segregant lines ( $22.70 \pm 0.27$  h,  $23.30 \pm 0.28$  h, respectively) (Figures 2A,B, Table S2). The measured periods were shorter than 24 h due to the high blue light intensity illumination used in our CF apparatus, which accelerates circadian oscillators and therefore shortens circadian period in constant conditions. Predicted loss of function of *Ttgi-B3* had little effect on circadian function (period  $22.70 \pm 0.22$  h,  $RAE 0.21 \pm 0.01$ ; Figures 2F,G, Table S2). Whereas plants in which there was a loss of functional *Ttgi-A3* had shorter period NPQ circadian rhythms (period  $21.30 \pm 0.14$  h,  $RAE 0.26 \pm 0.01$  Figures 2F,G, Table S2). The difference in period between single mutants was 1.40 h. Mutation of both copies of *GI* (*Ttgi-A3/gi-B3*) resulted in loss of robust circadian rhythms demonstrated by an  $RAE$  of  $0.59 \pm 0.05$  (Figures 2F,G, Table S2). The double mutant led to higher variation of plants that are rhythmic or arrhythmic. Here, 25% of the leaf fragments tested had an  $RAE < 0.5$ , indicating rhythmicity with a period of approximately 18 h. The high variability of  $RAE$  values and shorter period is also seen in Arabidopsis *gi-3* mutant lines (Mizoguchi et al. 2005). Assessment of another chlorophyll *a*  $RAE$  parameter  $F_v/F_m$  (Figure S4; Table S2) resulted in the same

conclusions that loss of *Ttgi-A3* shortens circadian period (*Ttgi-A3* period  $21.71 \pm 0.11$  h  $RAE = 0.27 \pm 0.02$ , *Ttgi-B3* period  $22.05 \pm 0.11$  h,  $RAE = 0.26 \pm 0.02$ ), with a difference in period of 0.75 h, and loss of both *Ttgi-A3* and *Ttgi-B3* abolishes robust circadian rhythms ( $RAE 0.52 \pm 0.04$ ; Figure S4G; Table S2).

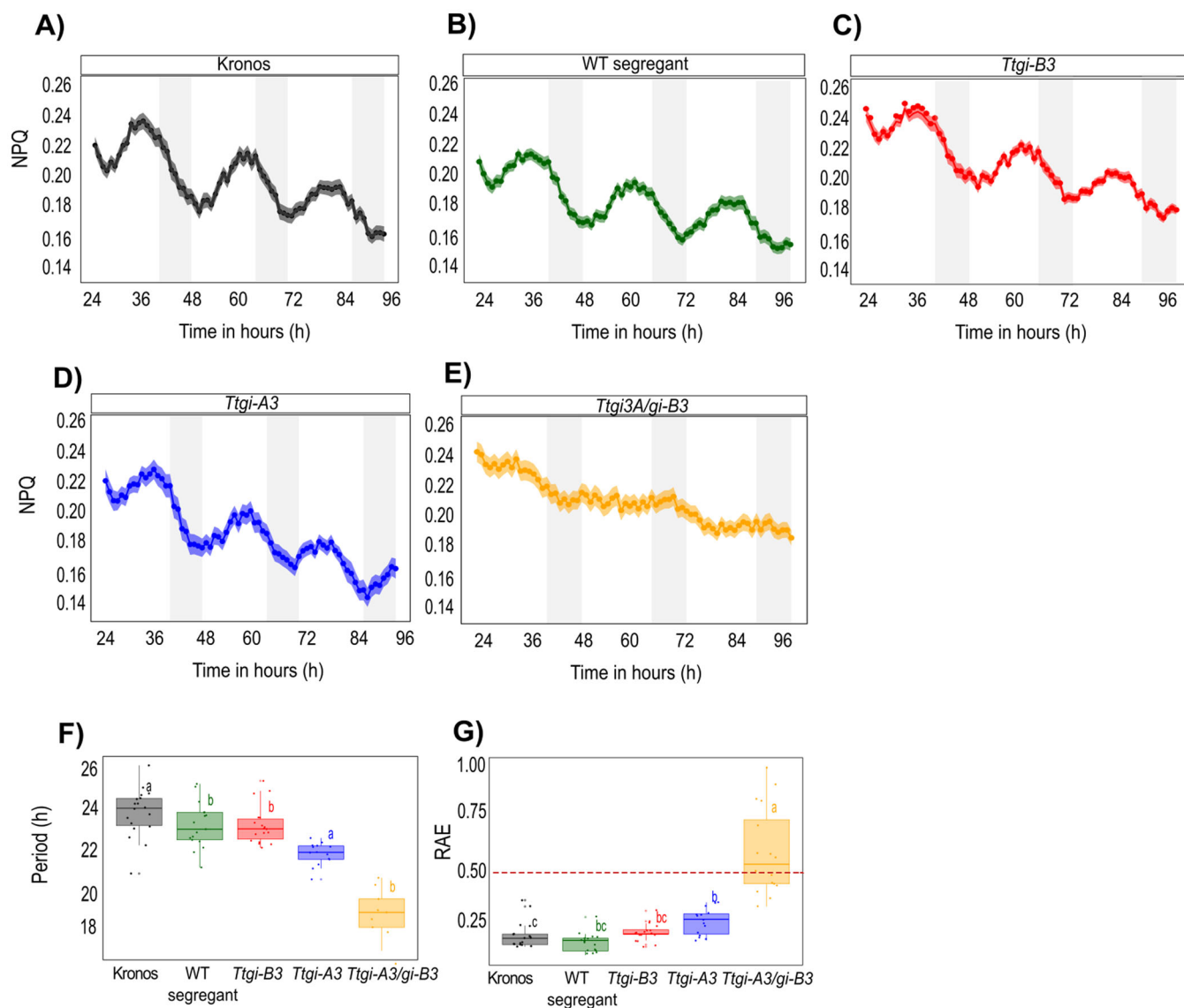
### 3.2 | Circadian Oscillator Transcript Abundance Is Perturbed in *Ttgi-A3/gi-B3* Loss-of-Function Lines

Having found that *GI* double mutants affect circadian rhythms in wheat, we investigated how this might occur by examining regulation of circadian oscillator transcript abundance in *Ttgi-A3/gi-B3* in white light-dark (LD) and continuous white light (LL). Under LD, transcripts of *TtLHY*, *TtTOC1*, *TtPRR73*, *TtLUX* and *TtGI* oscillated robustly in the *Kronos* genotype and *Ttgi-A3/gi-B3* (Figure 3). The loss of functional *GI* resulted in a reduction in amplitude of *TtGI* and *TtELF3*, an increase in amplitude of *TtTOC1* and little effect on the other components (Figure 3). The slightly advanced wave forms in LD in *Ttgi-A3/gi-B3* suggested that loss of *TtGI* advances the entrained phase of circadian oscillators in wheat (Figure 3). This early phase phenotype advanced the timing of maximal expression of *TtLHY* and *TtTOC1* in *Ttgi-A3/gi-B3* compared to *Kronos* (*TtLHY* ZT0 compared to ZT3, *TtTOC1* ZT9-12 compared to ZT12) in LD (Figure 3A,B), while phasing of peak transcript abundance of *TtPRR73* (ZT6), *TtGI* (ZT9) and *TtLUX* (ZT12) was unaffected in *Ttgi-A3/gi-B3* (Figure 3C,E,F). In LD conditions, there were no significant differences in *TtPPD-1* peak transcript abundance between the *Ttgi-A3/gi-B3* and *Kronos* (Figure 3D).

In the first true circadian cycle (beginning 24 h after the transition to LL), peak transcript abundance of *TtLHY*, *TtTOC1*, *TtPRR73*, *TtLUX* and *TtGI* was phased earlier in *Ttgi-A3/gi-B3* than in *Kronos* (denoted by orange stars). Oscillations of transcript abundance were detected in *Kronos* in the subsequent LL cycles, albeit with reducing amplitude, whilst in *Ttgi-A3/gi-B3* transcript abundance tended to be less rhythmic (Figure 3). Both *TtPPD-1* and *TtELF3* expression had undetectable rhythms in prolonged LL in both *Kronos* and *Ttgi-A3/gi-B3* (Figures 3D,G), consistent with previous reports of low amplitude rhythms of wheat *ELF3* transcripts (Wittern et al. 2023). JTK cycle (Figure 3H) analysis reported that *TtGI*, *TtLHY*, *TtPPD-1* and *TtTOC1* expression as rhythmic (BH,  $Q < 0.05$ ) in LL, however visual inspection suggests the rhythmic dynamics were not as robust, this was particularly evident for *TtLHY* (Figure 3A) and *TtPRR73* (Figure 3C).

### 3.3 | *TtGI* Is a Mild Promoter of Flowering in Long and Short-Day Photoperiods

Our data demonstrate that mutation of *GI* affects wheat circadian oscillators. In Arabidopsis, *GI* contributes to photoperiodism through its role in maintaining robust circadian rhythms and through the regulation of FKF1 (Sawa et al. 2007). The TILLING population used in our studies carries the semi-dominant *PPD1a* allele which reduces the sensitivity of heading date to photoperiod, though the mechanism by which this occurs is not currently understood. This gave us the opportunity to investigate whether loss of *GI* function, and the associated effect on circadian oscillators can affect heading date additively to the *PPD-1-*



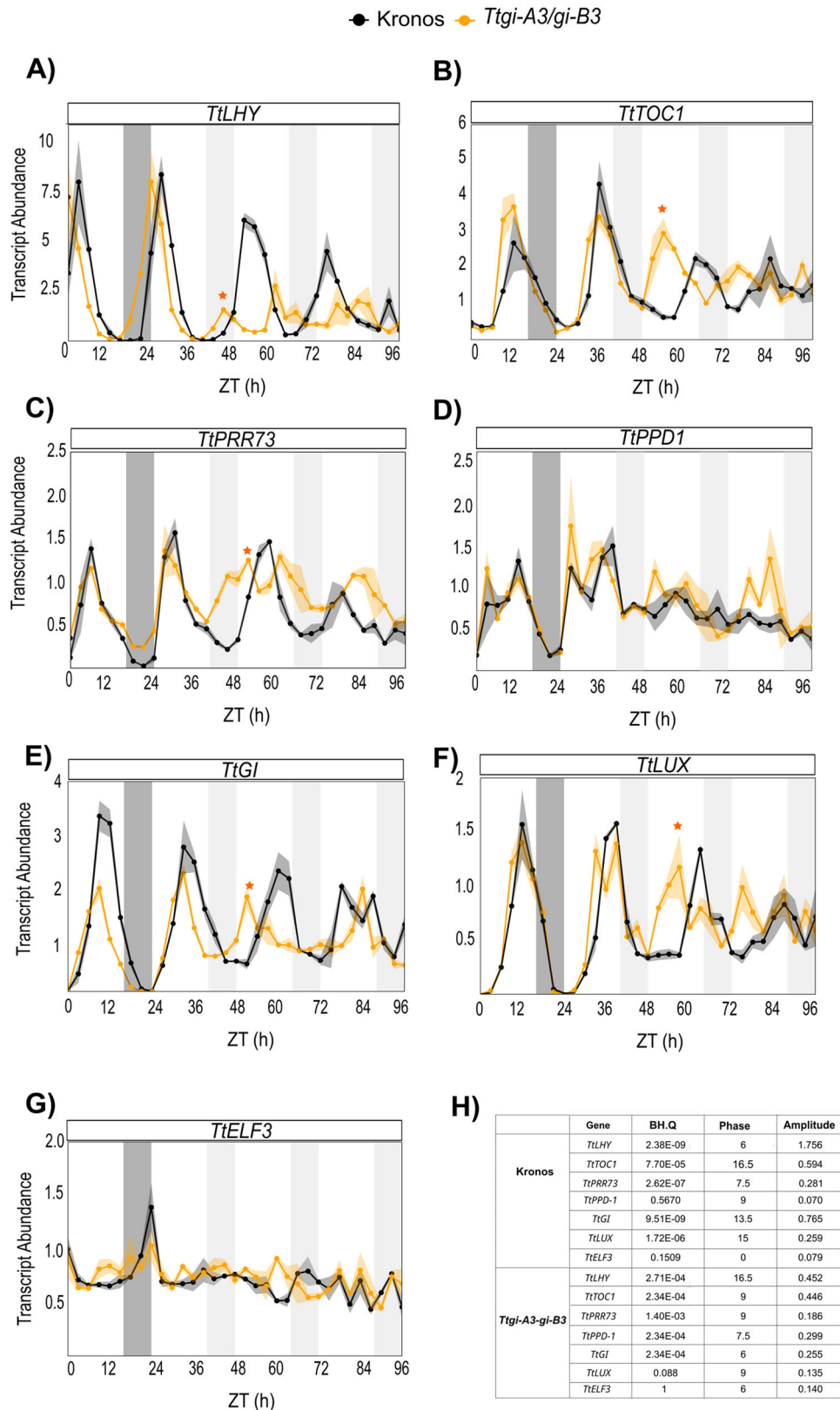
**FIGURE 2** | Functional *TtGI* is necessary for maintaining robust circadian rhythms in chlorophyll fluorescence. Mean of NPQ ( $\pm$  SEM, represented by the shaded ribbon) of Kronos (A), WT segregant (B), *Ttgi-B3* (C), *Ttgi-A3* (D) and *Ttgi-A3/gi-B3* (E) leaf fragments in constant light ( $n > 10$ ). White bars represent the subjective day, and grey bars represent the subjective night. (F) Circadian period (hours). (G) Relative error of amplitude (RAE). An RAE value above 0.5 (indicated by the dashed line) is considered arrhythmic. Period and RAE were calculated using Biodare2 for the CF parameter NPQ. Significant differences ( $p < 0.05$ ) calculated in R using the Kruskal–Wallis test followed by post hoc Dunn’s test. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

mediated photoperiod perception pathway. We grew Kronos, WT segregant, *Ttgi-A3*, *Ttgi-B3* and *Ttgi-A3/gi-B3* lines in long and short photoperiods recording when each plant reached the heading growth stage (GS55, half the ear emerged from the ligule), an easily observed growth stage which is commonly used as a proxy for flowering time in wheat (Zadoks et al. 1974).

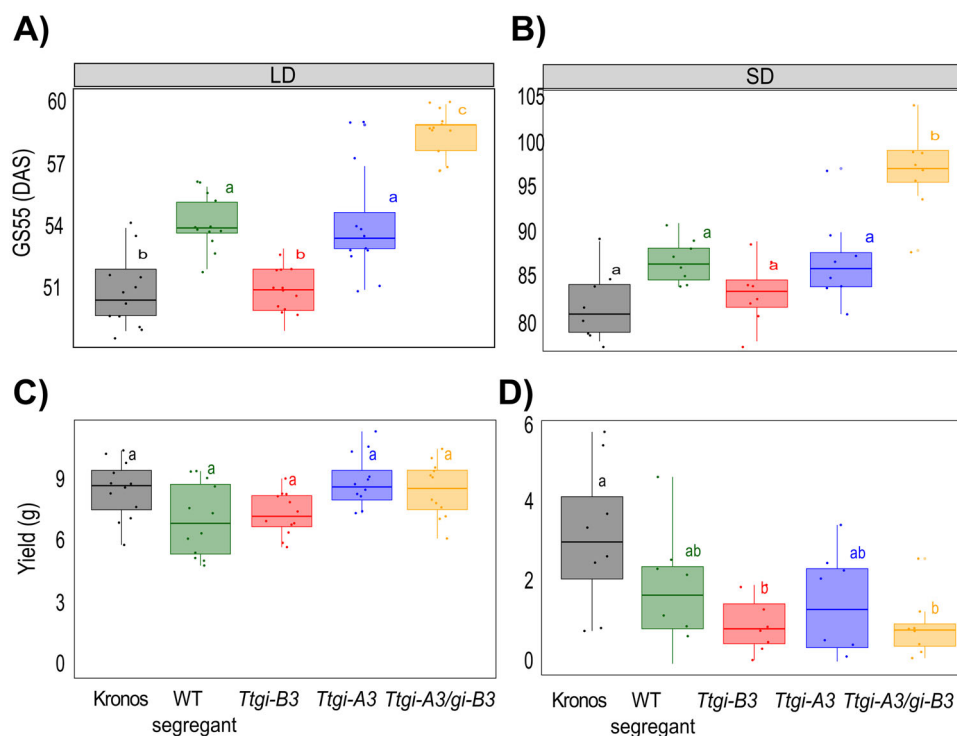
Under long photoperiod, heading time in the double mutant was later ( $59 \pm 0.31$  days) than either single mutant (*Ttgi-A3*  $54 \pm 0.78$  days, *Ttgi-B3*  $51 \pm 0.28$  days) and even later than Kronos ( $51 \pm 0.51$  days) and WT segregant ( $54 \pm 0.37$  days) (Figure 4A) demonstrating that the delayed flowering observed in the double mutant is caused by the absence of a functional *GI* rather than background mutations. In short photoperiods, the *Ttgi-3A/gi-3B* double mutants reached GS55 10 days after *Ttgi-A3* and WT segregant, 13 days after *Ttgi-B3* and 14 days after Kronos

(Figure 4B), demonstrating an effect on heading. These results differ from *Arabidopsis* where *GI* mutants (*gi-1-6*) and T-DNA insertion line (*gi-11*) flower later in long-day and have minor delays in short-day flowering (Fowler 1999). Taken together these data indicate that mutation of *GI* delays flowering in both long and short photoperiods, and that mutation of either the copies of *GI* on the A or B genome alone is without effect, or very weak.

Flowering time is a key determinant of yield in wheat (Hyles et al. 2020). To investigate if *Ttgi-A3/gi-B3* had an effect on yield, we quantified yield traits and measured the yield of each plant (total seed weight) grown in long or short photoperiods (Figures 4C,D). Predicted loss of functional *TtGI* had a greater effect on yield and yield traits under short photoperiods compared to long photoperiods. Under long photoperiods, seed yield was not significantly different between the genotypes, the mean yield per



**FIGURE 3** | Functional *TtGI* is required for persistence of robust oscillations of circadian oscillator transcript abundance in continuous light. (A–G) Mean abundance ( $\pm$  SEM, represented by the shaded ribbon) of circadian oscillator transcripts ( $n = 3$ ) in *Ttgi-A3/gi-B3* (yellow) and Kronos WT (black). Transcript abundance ( $\Delta\Delta Cq$ ) is relative to *TtRP15* and *TtRPT5A*, (A) *TtLHY*, (B) *TtTOC1*, (C) *TtPRR73*, (D) *TtPPD1*, (E) *TtELF3*, (F) *TtLUX*, and (G) *TtGI*. (H) Parameters of the rhythms of transcript abundance in LL. Orange stars indicate the peak of first true circadian cycle in LL in *Ttgi-A3/gi-B3*. White bars represent light, dark grey bars represent darkness in the last cycle before release into constant light. Light grey bars represent subjective night in constant light. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]



**FIGURE 4** | Flowering time is delayed in *Ttgi-A3/gi-B3* in long- and short-photoperiods. *Ttgi-A3/gi-B3*, *Ttgi-A3*, *Ttgi-B3*, WT segregant and Kronos were grown in long photoperiod (LD, 16 h light at  $250 \mu\text{mol m}^{-2} \text{s}^{-1}$ ,  $20^\circ\text{C}$ : 8 h dark  $16^\circ\text{C}$ ) and short photoperiod (LD, 8 h light at  $250 \mu\text{mol m}^{-2} \text{s}^{-1}$ ,  $20^\circ\text{C}$ : 16 h dark  $16^\circ\text{C}$ ). GS55 is used as reference for heading date and defined as days after sowing (DAS). (A) GS55 in long photoperiod ( $n = 12$ ). (B) GS55 in short photoperiod ( $n = 8$ ). (C) Yield (total seed weight) under long photoperiod ( $n = 12$ ). (D) Yield (total seed weight) under short photoperiod ( $n = 8$ ). Upper and lower hinges represent the first and third quartiles (25th and 75th percentiles), the middle hinge represents the median value, whiskers represent the third quartile +  $1.5\times$  interquartile range (IQR) and the first quartile -  $1.5\times$  IQR, each dot represent individual replicates. Significant differences in heading date were tested using a Kruskal–Wallis test followed by a post hoc Dunn test, and yield differences were tested using one-way ANOVA followed by Tukey’s test. The letters within each panel indicate statistical difference, samples that share the same letter in that experiment are not significantly different. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

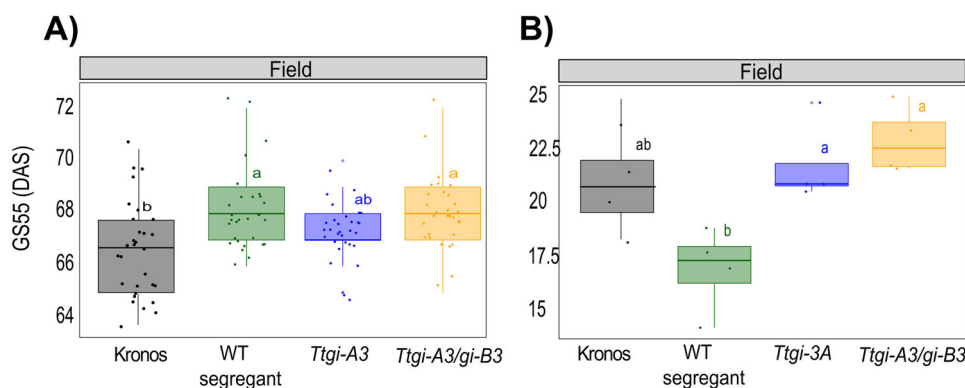
*Ttgi-A3/gi-B3* plant was  $8.55 \pm 0.39$  g compared to  $8.56 \pm 0.41$  g in Kronos. The mean yield of each individual Kronos plant grown in short photoperiods ( $3.13 \pm 0.64$  g) was significantly greater than *Ttgi-A3/gi-B3* ( $0.92 \pm 0.27$  g), and *Ttgi-B3* ( $0.99 \pm 0.24$  g), however there were no differences in seed weight between the WT segregant line ( $1.76 \pm 0.30$  g) and the double mutant (Figure 4C). The mean yield of the WT segregant line was lower than in Kronos, this could be due to the presence of the mutation load in this line and that short photoperiod conditions are less favourable for optimal growth. Under long photoperiods, height, number and length of the primary head, number and weight of seeds did not significantly differ (Figure S5). Under short photoperiod height, number of tillers and length of the primary head, were not statistically different between any of the lines, however, total seed number, seed weight and number of seeds of the primary head were substantially lower in *Ttgi-A3/gi-B3*, *Ttgi-A3*, *Ttgi-B3* compared to Kronos (Figure S6).

### 3.4 | Lack of Functional *TtGI* Did Not Have Large Effects on Heading Date and Yield Traits in the Field

We had established that *Ttgi-A3/gi-B3* had a late heading phenotype in long and short photoperiods in controlled environment conditions. Next, we wished to determine if this effect was

observed under field conditions. Wheat is a commercial crop, and it was therefore important to understand if the heading date phenotype seen in the laboratory was observable in the field, where interactions with the environment are more complex.

We performed a small field trial at the NIAB experimental farm in Cambridge, UK in the 2021 field season (environmental information in Supporting Information S1). Plots of Kronos, WT segregant, double *Ttgi-A3/gi-B3* mutant and single A mutant were grown (*Ttgi-A3*); a single B mutant was not available at the time of the trial. Five plants per plot were tagged (six plots per genotype) and the date each plant reached GS55 was recorded. *Ttgi-A3/gi-B3* reached GS55 only 1 day later than Kronos (*Ttgi-A3/gi-B3* 67 days, Kronos 68 days), while there was no difference in heading date between the double mutant and the WT segregant line (Figure 5A), which could be because of the presence of background mutations within in the WT segregant affecting flowering time. Therefore, whilst Kronos is not optimised for growth in UK field conditions, we found that if *GI* affects heading in the field, the effects are modest and possibly overridden by other regulators of flowering time, particularly in a *PPD-A1a* background. The mean weight of seeds at 15% grain moisture (Kg) per head gathered from *Ttgi-A3/gi-B3* ( $1.45 \pm 0.08$ ) plants was not significantly different to Kronos ( $1.69 \pm 0.08$ ), the single A mutant ( $1.54 \pm 0.09$ ), or WT segregant ( $1.51 \pm 0.07$ ) (Figure 5B). After the plants had fully senesced,



**FIGURE 5** | Mutations of single *TtGI* homologues and double mutant have no effect on flowering time in the field and yield. Kronos, WT segregant, *Ttgi-A3* and *Ttgi-A3/gi-B3* were grown in the field at NIAB experimental farm in the 2021 field season. (A) Heading date as defined as days after sowing to reach GS55. Data were pooled for all tagged plants of each genotype ( $n = 30$ ). Significant differences in heading date were tested using a Kruskal–Wallis test followed by a post hoc Dunn test. (B) Yield at 15% grain moisture in representative heads in each genotype grown in the field ( $n = 6$ ). Each jitter point represents the total yield of an individual plot. Upper and lower hinges represent the first and third quartiles (25th and 75th percentiles), the middle hinge represents the median value, whiskers represent the third quartile  $+1.5\times$  interquartile range (IQR) and the first quartile  $-1.5\times$  IQR, each dot represent individual replicates. Significant differences in heading date were tested using a Kruskal–Wallis test followed by a post hoc Dunn test, and yield differences were tested using one-way ANOVA followed by Tukey's test. Significant differences in yield were tested using one-way ANOVA followed by Tukey's test. The letters within each panel indicate statistical difference, samples that share the same letter in that experiment are not significantly different. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

morphological traits were measured and a representative head was taken from each tagged plant (Figure S7).

### 3.5 | Expression of Flowering Genes in the *Ttgi-A3/gi-B3* Line

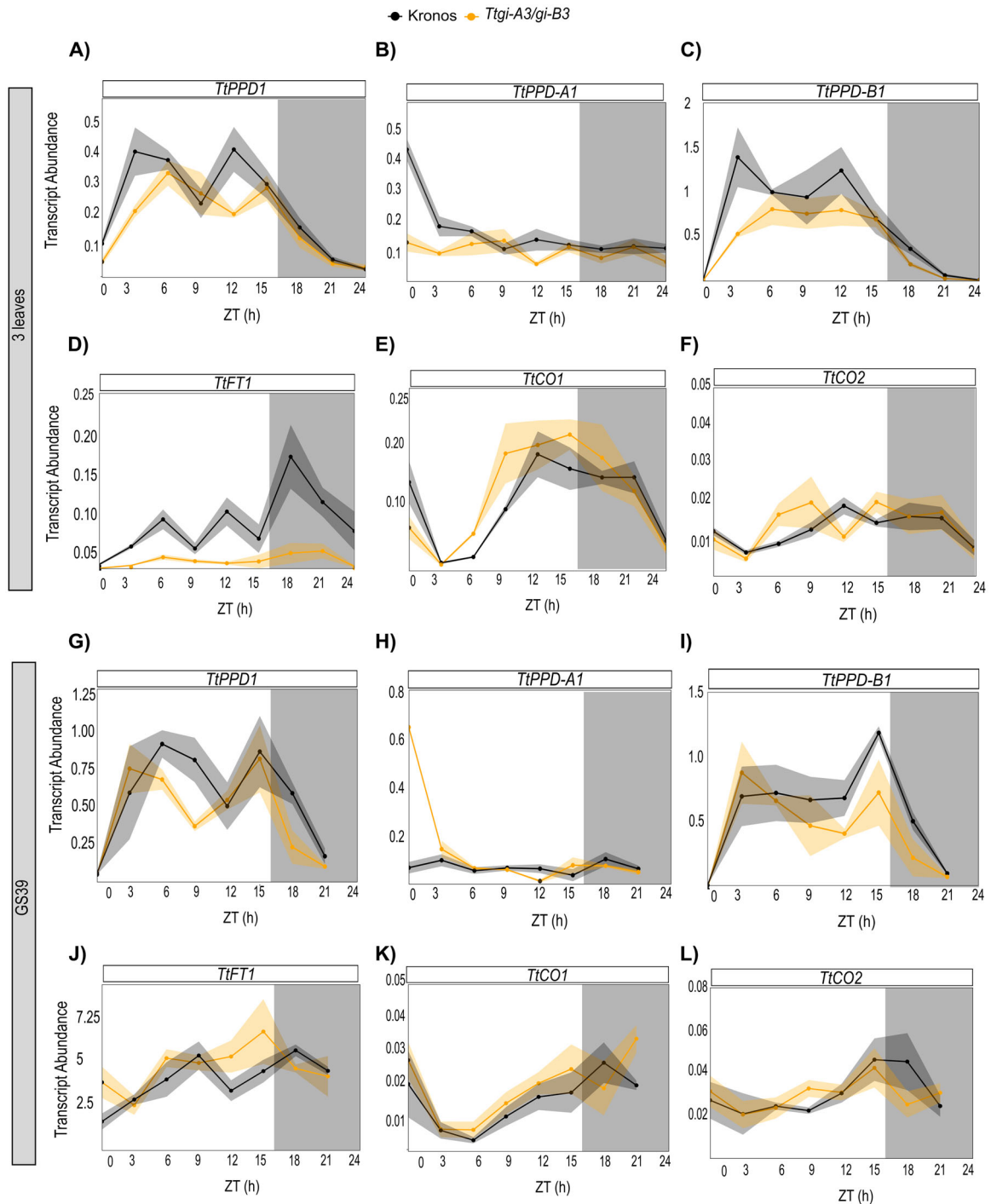
Having shown that *GI* is required for robust circadian rhythms, and that *TtGI* is a mild inducer of flowering in long and short photoperiods in a *PPD-A1a* background, we then characterised the effect of loss of function of *TtGI* on the transcription profile of several flowering regulatory genes. The absence of *TtGI* did not have a major effect on the expression of flowering genes at either the three-leaf stage or GS39 (Figure 6). Expression of *TtPPD-1* was measured using common primers for both sub-genome and sub-genome-specific primers. At the three-leaf growth stage, there were two peaks of *TtPPD1* abundance at ZT3 and 12 h in both genotypes, however, the phase in *Ttgi-A3/gi-B3* was delayed for approximately 3 h and the amplitude of transcript abundance was slightly lower than in Kronos (Figure 6A). *TtFT1* abundance was low in both *Ttgi-A3/gi-B3* mutant and Kronos but there was some evidence for loss of *GI* derepressing *TtFT1* at night (Figure 6D). There were no differences in *TtCO1* and *TtCO2* expression between *Ttgi-A3/gi-B3* and Kronos (Figure 6E,F). Similar results were observed in growth stage GS39, *TtPPD1* amplitude was lower in the double mutant than in the Kronos line, phasing at the same time in both genotypes (Figure 6G). No differences were detected on the expression of *TtFT1*, *TtCO1* and *TtCO2* between *Ttgi-A3/gi-B3* and Kronos (Figure 6F–H). Overall, our results show that mutation of *TtGI* does not affect the expression of photoperiodic flowering genes in wheat, contrary to Arabidopsis, indicating that there is functional divergence between the two species.

In Arabidopsis, the phytohormone gibberellin (GA) is essential for development processes including flowering and its signalling pathway is modulated by *GI* (Nohales et al. 2019). We therefore

investigated whether it was the same in wheat by analysing the expression of two genes involved in GA biosynthesis (*TtGA20ox2* and *TtGID1*). There were no differences in expression in either three-leaf growth stage or GS39, between the WT and the mutant line for any of the genes analysed (Figure S8).

### 3.6 | *TtGI* Sequence Is Conserved Across Modern Wheat Varieties

Our data indicate that whilst *GI* is required for robust circadian function, the effects on flowering in our conditions and the *PPD-A1a* background were modest. Variation within *loci* can have profound effects on the contribution of alleles to flowering, and so we investigated whether there was evidence for variation at *GI* *loci*, that could indicate that *GI* allelic variation might have been selected for its direct or epistatic interaction with other selected *loci* in other backgrounds and varieties. We identified *GI* orthologues in genome-sequenced wheat cultivars important in modern breeding efforts (Avni et al. 2017; Walkowiak et al. 2020) (Table S1) by performing a BLAST search using the IWGSC RefSeq V1.1 *TaGI* homoeologues as a query sequences. We then created alignments between *GI* from the different wheat cultivars and IWGSC RefSeq V1.1 to investigate variation in *GI*. Remarkably low levels of variation were observed in *GI* across the different wheat cultivars (Table 2). No variation existed in the *GI-D* coding sequences. Two SNPs within *GI-A* coding sequences were identified in the wild emmer tetraploid Zavitan, but not in any bread wheat cultivars. Five *GI-B* allele groups were identified within all varieties. The maximum number of non-synonymous SNPs in the *GI B* allele group was two. Thus, we found little variation in *GI* sequence at coding sequence level across modern wheat cultivars which can be exploited by breeders to manipulate flowering time and other circadian output traits. However, further research on noncoding sequences is needed.



**FIGURE 6** | Loss of function of *TtGI* did not affect the abundance of flowering transcripts. Transcript abundance (mean  $\pm$  SEM, represented by the shaded ribbon) of wheat genes involved in flowering in *Ttgi-A3/gi-B3* (yellow,  $n = 4-5$ ) and Kronos WT (black,  $n = 4-5$ ) grown under long-day (16 h light at  $250 \mu\text{mol m}^{-2} \text{s}^{-1}$ ,  $20^\circ\text{C}$ : 8 h dark  $16^\circ\text{C}$ ). Once plants reached the three-leaf stage or GS39, sampling of the first true leaf commenced at time 0 to 24 h, every 3 h. White bars represent light, and grey bars represent darkness. (A–F) Mean abundance of flowering transcript at the three-leaf growth stage. (G–L) Mean abundance of flowering transcripts at the GS39 growth stage. Transcript abundance ( $\Delta\Delta\text{Cq}$ ) is relative to *TtR15* and *TtRPT5A*, (A–G) *TiPPD1*, (B–H) *TiPPD1-A*, (C–I) *TiPPD1-B*, (D–J) *TiCO1*, (E–K) *TiCO2* and (F–L) *TiFT1*. [Color figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

## 4 | Discussion

### 4.1 | *TtGI* Is Required for Robust Circadian Oscillations in Wheat

Wheat circadian oscillators are perturbed in lines with predicted lack function copies of *TtGI*. *Ttgi-A3/gi-B3* lines have no detectable

rhythms of NPQ or  $F_v/F_m$  in constant conditions. Similarly, circadian oscillator transcript abundance lost robust oscillations in LL. Mutation of one *GI* homoeologue (*Ttgi-A3* or *Ttgi-B3*) had no effect on the robustness of circadian rhythms in CF. This demonstrates that similar to our previous findings concerning *TtELF3*, one homoeologue of a circadian oscillator gene is sufficient to maintain robust circadian rhythms (Steed et al. 2021; Wittern

**TABLE 2** | *GI* sequence is conserved across modern wheat varieties. Total number of SNPs within the CDS of *GI* across cultivars important in modern breeding efforts. A SNP was recorded if the base was different to IWGSC RefSeqv1.1. SNPs within the same codon were recorded individually.

Sub-genome	A		B		D	
	ns	s	ns	s	ns	s
Cultivar	ns	s	ns	s	ns	s
Arina	0	0	2	3	0	0
CDC Stanley	0	0	2	2	0	0
Landmark	0	0	2	3	0	0
Cadenza	0	0	2	3	0	0
Claire	0	0	NA	NA	0	0
Jagger	0	0	2	3	0	0
Julius	0	0	2	3	0	0
Lancer	0	0	2	3	0	0
Norin 61	0	0	2	2	0	0
Mace	0	0	2	3	0	0
P190962	0	0	0	1	0	0
Paragon	0	0	1	3	0	0
Robigus	0	0	1	3	0	0
SY Mattis	0	0	2	3	0	0
Zavitan	1	1	1	5	0	0
Total no. SNPs	1	1	23	40	0	0
Total no. alleles	2		6		1	

Note: Allele groups are indicated by colour. Cultivars were considered to share the same allele if the CDS were identical. Zativan is a tetraploid variety and so lacks a D subgenome. Claire *GI* B analysis was not included due to poor sequence quality. Abbreviations: NS, non-synonymous SNP; S, synonymous SNP.

et al. 2023). Due to differences in expression of circadian oscillator gene homoeologues it has been proposed that one dominant homoeologue performs the majority of the biological function in the circadian oscillator (Rees et al. 2022). We found that levels of transcription of both homoeologues was similar, however mutation of the *GI* A homoeologue had a greater effect on circadian period than mutation of the B *GI* homoeologue.

In Arabidopsis, *GI* contributes to the daily rhythms of ZTL activity (Cha et al. 2017; Lee et al. 2019). Wheat has two orthologous genes to Arabidopsis ZTL (Calixto et al. 2015), however it is unknown whether the two wheat ZTL orthologs are functionally redundant in wheat circadian oscillators. In Brachypodium, *GI* and ZTL interact in a yeast two hybrid screen (Hong et al. 2010). Further work is required to confirm the role of *GI* and ZTL in wheat at the molecular level.

#### 4.2 | *TtGI* Is a Promoter of Flowering in Wheat

To investigate the role of *GI* in photoperiodic flowering, we measured GS55 and yield parameters in long, short photoperiod and in field conditions. Our study focuses on flowering time regulation independent of *PPD-1*-dependent pathways because the *Ttgi-A3/gi-B3* mutant was generated in a *PPD-A1a* photoperiod-insensitive background. This allowed us to

investigate the role of *GI* independent of *PPD-1* but does mean we have not been able to study genetic interactions between *GI* and *PPD-1*. In both long and short photoperiods, *Ttgi-A3/gi-B3* headed later than Kronos but few differences were observed in the field indicating that *TtGI* is a mild promoter of flowering in wheat, in a *PPD-A1a* background. In Arabidopsis, *GI* and *FKF1* complex during the light period only during long days, promoting *CO* expression in a photoperiod-dependent manner (Fowler 1999; Sawa et al. 2007). This mechanism, plus targeting of *CO* for degradation by *COP1* in the dark, accelerates flowering in Arabidopsis in long days and forms the molecular basis of the external coincidence model.

A core principle of the external coincidence model is that there are light-sensitive and insensitive phases of the flowering-promoting rhythm (Bunning 1960). This can be demonstrated experimentally by treating plants with skeleton photoperiods (Thomas and Vince-Prue 1997; Roden et al. 2002). Flowering occurred earliest in wheat plants when the light period was applied in the middle of the night (Pearce et al. 2017), thus demonstrating that there are rhythms in light sensitivity in the wheat flowering pathway. The circadian clock in wheat forms a complex network involving the interactions of various proteins and genes that regulate photoperiodic flowering. *ELF3* acts as a mediator between the light signalling pathway and flowering in wheat by binding to *PhyB* and *PhyC*. Moreover, in photoperiodic regulation of flowering, *ELF3* regulates *PPD-1* by binding directly to its promoter and repressing its expression (Alvarez et al. 2023). *TtCO1/CO2* and *TtPPD1* operate in two distinct but highly connected systems, *PPD-1* represses expression of *CO1* affecting the expression of *FT1* (Shaw et al. 2020). According to our gene expression data, *TtGI* does not promote the expression of *TtCO1* and *TtCO2* under long-day photoperiod which indicates that *TtGI* does not directly regulate *CO1/2*. However, a recent study in a photoperiod-sensitive line of Kronos carrying a *Ppd-A1b* allele found that *gi* mutants increase *TtCO2* expression and that *CO1* and *CO2* physically interact with *GI* in Y2H (Li et al. 2024). In a photoperiod-sensitive *Ppd-A1b* allele-carrying line of Kronos *gi* mutants headed early with a strong interaction with photoperiod (Li et al. 2024). Our data support the conclusion that *GI* has little effect on heading date in a *PPD-A1a* background (Li et al. 2024). Taken together these data suggest that *TtGI* regulates *TtCO1/2* through a *PPD-1*-dependent pathway and that *GI* might differentially regulate flowering in wheat and Arabidopsis.

## 5 | Conclusion

We find that reduction of *GI* function, reduces robustness of circadian oscillations but had little effect on flowering time in a *PPD-A1a* photoperiod-insensitive background. This and the recent finding that *gi* mutants affect flowering in *PPD-A1b* photoperiodic backgrounds (Li et al. 2024) suggest that if the circadian oscillator and *GI* do contribute to photoperiodism in wheat, it is likely to be mostly dependent on *PPD1*.

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### Conflicts of Interest

Mathew Hannah is an employee of BASF. The other authors declare no conflicts of interest.

### Data Availability Statement

The data that support the findings of this study are openly available in the University of Cambridge Data Repository at <https://doi.org/10.17863/CAM.107919>.

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### Supporting Information

Additional supporting information can be found online in the Supporting Information section.