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Research Article

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Identification and characterization of a natural polymorphism in *FT-A2* associated with increased number of grains per spike in wheat

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Short Title: *FT-A2* polymorphism increases grain number per spike

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Abstract

Increases in wheat grain yield are necessary to meet future global food demands. A previous study showed that loss-of-function mutations in FLOWERING LOCUS T2 (FT2) increase spikelet number per spike (SNS), an important grain yield component. Unfortunately, associated reductions in fertility offset potential increases in grain number. Here, we report a natural mutation resulting in an aspartic acid to alanine change at position 10 (D10A) associated with significant increases in SNS and no negative effects on fertility. Using a high-density genetic map, we delimited the SNS candidate region to a 5.2 Mb region on chromosome 3AS including 28 genes. Among them, only FT-A2 showed a non-synonymous polymorphism (D10A) present in two different populations segregating for the SNS QTL on chromosome arm 3AS. These results, together with the known effect of the ft-A2 mutations on SNS, suggest that variation in FT-A2 is the most likely cause of the observed differences in SNS. We validated the positive effects of the A10 allele on SNS, grain number, and grain yield per spike in near-isogenic tetraploid wheat lines and in an hexaploid winter wheat population. The A10 allele is present at very low frequency in durum wheat and at much higher frequency in hexaploid wheat, particularly in winter and fall-planted spring varieties. These results suggest that the FT-A2 A10 allele may be particularly useful for improving grain yield in durum wheat and fall planted common wheat varieties.

Key message

We discovered a natural FT-A2 allele that increases grain number per spike in both pasta and bread wheat with limited effect on heading time.
Introduction

Wheat is a global crop of major economic value and nutritional importance as it provides around 20% of the calories and protein consumed by the human population (http://www.fao.org/faostat/en/#data/FBS). However, with ever changing environmental conditions and the rising human population, it is critical to increase wheat grain yield to meet future demands. Yield is a multifaceted trait that can be partitioned into several yield components, including spikes per unit of area, spikelet number per spike (SNS), grains per spikelet, and grain weight. Several genes have been identified that affect these grain yield components (Kuzay et al. 2019; Li et al. 2019; Poursarebani et al. 2015; Sakuma et al. 2019; Shaw et al. 2013; Simmonds et al. 2016; Wang et al. 2019).

Unfortunately, many of the genes affecting SNS also have strong effects on heading date that limit their use in variety development. Significant yield penalties are usually observed for varieties heading before (e.g. incomplete grain filling) or after (e.g. increased risk of heat impacting seed filling) the optimum heading interval to maximize grain yield. For example, the vrnl-null mutant significantly increases SNS by delaying the transition of the inflorescence meristem to a terminal spikelet, but also delays the transition of the vegetative meristem to inflorescence meristem, resulting in a very late heading time (Li et al. 2019). Another good example is the main wheat photoperiod gene PHOTOPERIOD1 (PPD1), which shows a strong correlation between heading date and SNS in lines carrying different dosages of PPD1 loss-of-function mutations ($R^2 = 0.74$) (Shaw et al. 2013). A correlation between heading date and SNS has also been observed in genes regulated by PPD1 such as the FLOWERING LOCUS T1 gene (FT1) (Brassac et al. 2021; Finnegan et al. 2018; Isham et al. 2021; Lv et al. 2014).

*FT1* encodes a mobile protein that travels through the phloem and carries environmental signals from the leaves to the shoot apical meristem (SAM), where it forms a complex with 14-3-3 and FD-like proteins (Florigen Activation Complex) (Taoka et al. 2011). This complex binds to the promoter of the meristem identity gene VERNALIZATION1 (VRN1), promoting its expression and the transition from the vegetative to the reproductive phase in wheat (Li et al. 2015).

Induction of *FT1* also results in the upregulation of SUPPRESSOR OF OVEREXPRESSION OF CONSTANS1-1 (SOC1), LEAFY (LFY) and genes in the gibberellin (GA) pathway that are essential for spike development and stem elongation (Pearce et al. 2013). A deletion of *FT-B1* in
hexaploid wheat delays the transition to reproductive growth and increases SNS (Finnegan et al. 2018).

In addition to \textit{FT1}, wheat has at least five \textit{FT-like} paralogs designated as \textit{FT2} to \textit{FT6} (Lv et al. 2014), which have some overlapping functions but also varying degrees of sub-functionalization (Halliwell et al. 2016; Lv et al. 2014). \textit{FT2} is the most similar paralog to \textit{FT1} (78\% protein identity), but the two genes still exhibit marked differences in transcription and protein interaction profiles. Whereas the \textit{FT1} protein interacts with five out of the six wheat 14-3-3 proteins tested so far, \textit{FT2} failed to interact with any of these members of the Florigen Activation Complex (Li et al. 2015). The two genes also differ in their temporal and spatial transcription profiles. \textit{FT1} transcript levels in the leaves are upregulated earlier than \textit{FT2} when plants are grown at room temperature, but only \textit{FT2} is induced when plants are grown for a long period at 4 °C (vernalization) (Shaw et al. 2019). Interestingly, \textit{FT2} is the only member of the wheat \textit{FT-like} gene family that is expressed directly in the shoot apical meristem (SAM) and in the developing spike (Lv et al. 2014), in addition to leaves and elongating stems (Fig. S1).

Loss-of-function mutations in \textit{FT2}, identified in a sequenced mutant population of the tetraploid wheat variety Kronos (Krasileva et al. 2017), resulted in limited differences in heading time but significantly increased SNS (Shaw et al. 2019). Similar increases in SNS were observed in \textit{ft-B2} natural mutants detected in hexaploid wheat (Gauley et al. 2021). The loss-of-function mutation in the A-genome copy of \textit{FT2} (\textit{FT-A2}) in Kronos was associated with significantly larger increases in SNS (10-15\%) than the mutation in the B-genome copy (\textit{FT-B2}, 2-5\%). This difference in SNS was associated with much higher transcript levels of \textit{FT-A2} relative to \textit{FT-B2} in all tissues and developmental stages (Fig. S1). The increases in spikelet number in the double \textit{ft-A2 ft-B2} mutant (henceforth \textit{ft2-null}) were significantly larger than in the single \textit{ft-A2} mutant confirming that the \textit{FT-B2} gene still has a residual effect on SNS in spite of its lower transcript levels.

Unfortunately, the increase in SNS in the \textit{ft-A2} mutant was associated with reduced fertility, offsetting the potential positive effects of the increase in SNS on total grain yield (Shaw et al. 2019). We hypothesized that strong selection in cultivated wheat for grain yield might have selected an \textit{FT-A2} variant with a positive effect on SNS, but without the associated negative effect on fertility. Analysis of natural variation in \textit{FT-A2} revealed an aspartic acid to alanine
change at position 10 (D10A) that was rare in tetraploid wheat but frequent in modern common wheat varieties, suggesting positive selection for the new allele. In this study, we characterized the effect of the D10A polymorphism on wheat heading time, SNS, grain number, and spike yield in different wheat classes and performed a high-density genetic map of the SNS QTL that identified $FT-A2$ as the most likely candidate gene.

**Material and Methods**

Analysis of the exome capture data generated by the WheatCAP project using the assay developed by NimbleGen (Krasileva et al. 2017) and deposited in the Wheat T3 database (https://wheat.triticeaetoolbox.org/) revealed the existence of an A to C SNP within the $FT-A2$ coding region that resulted in the D10A polymorphism. We studied the effect of this SNP on heading time, SNS, grain number, and spike yield in two segregating populations in tetraploid and hexaploid wheat.

**Biparental mapping population in tetraploid wheat** (*Triticum turgidum* ssp. *durum*)

The tetraploid mapping population included 163 BC$_1$F$_2$ lines from the cross Kronos *2/Gredho (designated KxG hereafter). Kronos (PI 576168, $FT-A2$ D10 allele) is a semi-dwarf ($Rht-B1b$), with reduced photoperiod sensitivity ($Ppd-A1a$) spring wheat, whereas Gredho (PI 532239, $FT-A2$ A10 allele) is a tall ($Rht-B1a$), photoperiod sensitive ($Ppd-A1b$) spring landrace from Oman.

We planted the KxG population as headrows in 2015-2016 season at the UC Experimental Field Station in Davis, CA with each row including on average five individual plants.

**Near isogenic lines of the $FT-A2$ A10 allele from Gredho into Kronos**

We also evaluated the effect of the $FT-A2$ alleles in two sets of near isogenic lines (NILs). For the first set, we selected $FT-A2$ heterozygous BC$_1$F$_2$ and BC$_1$F$_3$ lines from the cross Kronos *2/Gredho using the $FT-A2$ marker, and selected two sets of homozygous BC$_1$F$_{3-4}$ homozygous A10 and D10 sister lines (H2-14 and H2-23). The Kronos isogenic line with the A10 allele was deposited in the National Small Grain Collection (PI 699107). We used the BC$_1$F$_{3-5}$ grains produced by these plants for two field experiments, one at the University of California, Davis (UCD) and the other one at Tulelake (California northern intermountain region). Both field experiments were organized in a complete randomized design with plants as experimental units.
Three to five spikes were measured per plant and averaged for 10 plants per genotype at the UC Davis experiment. In the Tulelake experiment, 23-27 spikes per genotype were randomly collected and used as experimental units in the statistical analyses.

In parallel, we backcrossed the A10 allele into Kronos for three additional generations (Kronos *5/Gredho), and then selected BC$_4$F$_2$ NILs homozygous for the A10 and D10 alleles using the FT-A2 molecular marker. The BC$_4$F$_3$ seed was increased in the greenhouse in 2020 and the BC$_4$F$_4$ grains were used for a second field experiment at UCD in 2021 that used small plots (four 1-m rows, 1.1 m$^2$) as experimental units, organized in a randomized complete block design with 12 blocks.

**Biparental mapping population in hexaploid winter wheat**

The hexaploid population included 358 F$_5$-derived recombinant inbred lines (RILs) derived from the cross between soft-red winter wheat lines LA95135 (CL-850643/PIONEER-2548//COKER9877/3/FL-302/COKER-762) x SS-MVP57 (FFR555W/3/VA89-22-52/TYLER//REDCOAT*2/GAINES). LA95135 is semidwarf (Rht-D1b) and photoperiod sensitive (Ppd-D1b), whereas SS-MVP57 is tall (Rht-D1a) and has reduced photoperiod sensitivity (Ppd-D1a) (DeWitt et al. 2021). This winter wheat population was previously genotyped and phenotyped as 1 m rows in the field at Raleigh, NC and Kinston, NC during the 2017-2018 season, and in Raleigh, Kinston, and Plains, GA in the 2018-2019 season (DeWitt et al. 2021). These locations will be referred to as Raleigh (Ral), Kinston (Kin), and Plains (Pla) followed by the harvest year (18 or 19).

**FT-A2 marker development and allelic frequencies**

We targeted the FT-A2, D10A SNP at position 124,172,909 bp (RefSeq v1.0) on chromosome 3A with a Cleaved Amplified Polymorphic Sequence (CAPS) marker. Primers FT-A2-D10A forward and reverse (Table S1) amplify a fragment of 705 bp. After digestion with the restriction enzyme ApaI, the fragment amplified from the D10 allele remained undigested, whereas the fragment amplified from the A10 allele was digested into two fragments of 448 and 257 bp.

We used this marker to determine the frequency of the D10A mutation in 89 *T. urartu*, 82 *T. turgidum* ssp. *dicoccoides*, 32 *T. turgidum* ssp. *dicoccon*, 417 *T. turgidum* ssp. *durum* and 705 *T. aestivum* accessions summarized in Supplementary Appendix S1. Among the hexaploid lines, we
included a collection of 238 landraces and varieties (He et al. 2019) and a set of 126 winter wheats (T3/Wheat) genotyped by exome capture and with data for the \textit{FT-A2 D10A} polymorphism. We also used the \textit{FT-A2} marker to genotype a panel of 242 spring wheats with reduced photoperiod sensitivity (Zhang et al. 2018) and a panel of 99 varieties and modern breeding lines from the Montana State University wheat breeding program (Supplementary Appendix S1). The spring lines were further classified based on the planting season used in the area where they were developed into those developed under spring planting (hereafter "DuS") or under fall planting (hereafter “DuF”). A previous study has previously shown that DuS and DuF groups are genetically differentiated using the 90K SNP array (Zhang et al. 2018) (Supplementary Appendix S1).

\textbf{High resolution genetic map}

We developed a high-resolution map of the KxG population in two phases. In the first phase, we identified two BC\textsubscript{1}F\textsubscript{3} plants from the KxG BC\textsubscript{1}F\textsubscript{2} head rows, H2 and D12, which were heterozygous for \textit{FT-A2} candidate region. From these heterozygous lines we generated large segregating Heterogeneous Inbred Families (HIF) populations to identify recombination events within the \textit{FT-A2} candidate region. Phenotype screens of these recombinants were space-planted at least three inches apart in a completely randomized design. To generate additional markers in the candidate gene region, we developed markers for 11 genes on both sides of \textit{FT-A2} covering a region of \textasciitilde10 Mb using the exome capture sequence data from Kronos and Gredho (Table S1).

\textbf{Statistical analysis}

In the tetraploid biparental population, we analyzed the effect of the \textit{FT-A2} alleles with a 3 x 2 factorial ANOVA that included the genotypic variation at \textit{PPD-A1} and \textit{RHT-B1} as additional factors, since both genes are known to have pleiotropic effects on heading time and yield components. In the hexaploid winter wheat population, we analyzed the effect of the \textit{FT-A2} in a 4 x 2 factorial ANOVA including the segregating genes \textit{PPD-D1}, \textit{RHT-D1} and \textit{WHEAT ORTHOLOG OF APO1 (WAPO-A1)}, which was previously shown to affect SNS (Kuzay et al. 2019). Analysis of Variance was conducted with the “Anova” function in R package “car” (Fox et al. 2019) with type 3 sum of squares.

\textbf{Yeast two-hybrid assays}
Modified Gateway (Invitrogen) bait/prey vectors pLAW10 and pLAW11 (Cantu et al. 2013) and yeast strain Y2HGold (Clontech, Mountain View, CA, USA) were used in the yeast two-hybrid assays. pLAW10 is the Gateway version of pGBKT7 (GAL4 DNA-binding domain, BD) and pLAW11 is the Gateway version of pGADT7 (GAL4 activation domain, AD). For all Gateway compatible cloning, pDONR/Zeo (Life Technologies, Grand Island, NY, USA) was used to generate the entry vectors. All constructs were verified by sequencing. Yeast two-hybrid assays were performed according to the manufacturer’s instructions (Clontech). Transformants were selected on SD medium lacking leucine (−L) and tryptophan (−W) plates and re-plated on SD medium lacking -L, -W, histidine (−H) and adenine (−A) to test the interactions.

Results

Natural variation in FT-A2

We used exome capture data deposited in the T3 database (https://triticeaetoolbox.org/wheat/) to explore the natural polymorphisms in FT-A2. We identified an A to C SNP at position 124,172,909 in chromosome arm 3AS of the Chinese Spring (CS) RefSeq v1.0, which resulted in an amino acid change of aspartic acid (D) to alanine (A) at position 10 of the FT-A2 protein (henceforth, D10A). In the analyzed accessions of T. urartu, T. turgidum ssp. dicoccoides and T. turgidum ssp. dicoccon, we detected only the D10 allele (Table 1). D10 was also the only allele detected in all the other grass species we analyzed including Lolium perenne (AMB21802), Oryza sativa (XP_021310907), Zea mays (NP_001106251), and Panicum virgatum (APP89655), indicating that D10 is the ancestral grass allele. The Chinese Spring reference genome carries the derived A10 allele, but in this study, we describe the change from the ancestral to the derived allele rather than relative to the reference genome.

We also screened a collection of 417 T. turgidum ssp. durum accessions with a CAPS marker for the D10A polymorphism (see Material and Methods) and found that only 0.7% carried the A10 allele (Table 1). Two of the three accessions with the A10 allele were from Oman (PI 532239 = ‘Gredho’ and PI 532242, ‘Musane and Byaza’) and the other one was from Turkey (PI 167718), suggesting that the A10 allele is almost absent from modern Western durum germplasm.

We detected a higher frequency of the A10 allele (56.5 %) among 705 T. aestivum ssp. aestivum lines (Table 1). This overall frequency was similar to that detected in a worldwide collection of
landraces and varieties combining winter and spring lines (59.7\%) (He et al. 2019). We also analyzed the frequency of the D10A polymorphisms in two collections with known growth habit, and found a higher frequency of the A10 allele among the winter lines (81.7\%) than among the spring lines (44.9\%, Table 1). Among the 341 spring wheat lines genotyped with the \textit{FT-A2} marker, we found that varieties developed under fall-planting (DuF or long cycle) had a significantly higher frequency of the A10 allele (58.4\%) than those developed under spring-planting (DuS or short cycle, 34.4\%, Table 1). A complete list of the accessions used in these calculations is available in Supplementary Appendix 1, and a summary of the frequencies is presented in Table 1.

Effect of the D10A polymorphism in tetraploid wheat

To test the effect of the D10A polymorphism on SNS, we used the diagnostic CAPS marker to screen 163 BC\textsubscript{1}F\textsubscript{2} plants from the KxG population segregating for this polymorphism. We also genotyped this population with markers for the segregating \textit{RHT-B1} (Guedira et al. 2010) and \textit{PPD-A1} (Wilhelm et al. 2009) genes, which can also affect SNS. Plants were grown in the field in the 2015-2016 season in Davis, CA and were phenotyped for individual plant height (HT), days to heading (DTH), and spikelet number per spike (SNS, Table 2).

The three-way factorial ANOVAs including \textit{FT-A2}, \textit{RHT-B1}, and \textit{PPD-A1} as factors showed significant effects for SNS, HT, and DTH and no significant interactions for any of the traits. As expected, \textit{RHT-B1} showed the strongest effect on plant height and \textit{PPD-A1} on heading time, although both genes affected both traits (Table 2). The strongest effect on SNS was detected for \textit{PPD-A1}, but a significant effect was also detected for \textit{FT-A2} (Table 2), with plants homozygous for A10 showing 6.4 \% higher SNS than those homozygous for D10 allele (Table 2). The differences in SNS between the \textit{FT-A2} alleles were larger in the late flowering plants homozygous for the photoperiod sensitive allele from Gredho (2.3 spikelets/spike) than in the early flowering plants homozygous for the Kronos allele for reduced photoperiod sensitivity (1.0 spikelets per spike), but the interaction was not significant.

Effect of the \textit{FT-A2} alleles in Kronos near isogenic lines
To analyze the effect of the D10A polymorphism independently of the variability generated by other major genes, we evaluated two sets of near isogenic lines in field experiments in 2020 at UCD and Tulelake (BC$_{1}$F$_{3-5}$ sister lines), and in 2021 at UCD (BC$_{4}$F$_{2-4}$ sister lines, see Material and Methods). In the 2020 experiment at UCD, lines with the A10 allele (PI 699107) showed large and significant increases in SNS (13.8%), grain number per spike (GNS, 31.7%), grains per spikelet (16.1%, also referred to as fertility) and grain yield per spike (33.0%) relative to the sister lines homozygous for the D10 allele (Table 3). The results from this experiment were consistent between two independent pairs of BC$_{1}$F$_{3-5}$ sister lines (H2-14 and H2-23, Table 3).

The experiments in Tulelake (Northern California, spring planting) using BC$_{1}$F$_{3-5}$ sister lines from family H2-14, also showed increases in SNS (4.0%), GNS (5.4%), grains per spikelet (1.7%), and grain yield per spike (10.5%) associated with the A10 allele. However, the magnitude of the differences between the $FT$-$A_2$ alleles was smaller than those observed at the 2020 UCD experiment under fall planting. Only the differences in SNS were statistically significant in Tulelake (Table 3).

For the 2021 UCD experiment using sister BC$_{4}$F$_{2-4}$ lines, we had more grains available and we were able to use small plots (1.1 m$^2$) as experimental units, with 12 replications per genotype. Lines with the $FT$-$A_2$ allele headed on average 0.8 d later than those with the D10 allele ($P = 0.0252$) and showed significant increases in SNS (5.7%, $P = 0.0011$) and GNS (6.3%, $P = 0.0168$, Table 3). In this experiment we did not detect significant differences in grains per spikelet ($P = 0.7919$). We observed a negative correlation between average GNS and grain weight across the 24 plots ($R = -0.61$) and a significant negative effect of the A10 allele on kernel weight (-7.8 %, $P = 0.0002$). The negative effect on grain weight offset the positive effect of the A10 allele on grain number resulting in non-significant differences in grain weight per spike (Table 3). We harvested the complete plots and measured grain yield per plot and the average yields of the two genotypes were almost identical: D10 = 1,254 ± 26 and A10 = 1,251 ± 32 g ($P = 0.9103$).

The A10 allele has a positive effect on SNS and spike yield in winter wheat

To analyze the effect of the D10A $FT$-$A_2$ alleles in winter wheat, we used phenotypic data available from 358 F$_5$-derived RILs from the cross between soft-red winter wheat lines LA95135...
and SS-MVP57 (DeWitt et al. 2021) and genotypic data for the FT-A2 marker developed in this study. This population was also segregating for PPD-D1, RHT-D1, and WAPO-A1, which were included as factors together with FT-A2 in a 4 x 2 factorial ANOVA.

Plants carrying the FT-A2 allele A10 (SS-MVP57) headed on average 1.7 days later (P < 0.001, Fig. 1a) and had 0.6 more spikelet per spike (5.1 % increase, P < 0.001, Fig. 1b) than plants carrying the D10 allele (LA95135). The differences in SNS were significant in all tested locations. The A10 allele was also associated with an average 5.8% increase in GNS in the two locations where this trait was measured, but the differences were significant only for the Pla19 location (2.7 more grains per spike, P < 0.001, Fig. 1c). The A10 allele was also associated with a 1.2% increase in the number of grains per spikelet but the differences were not significant (Fig. 1d). The differences in SNS were associated with a significant 4.6% increase in average spike yield associated with the A10 allele in two out of the three tested locations (P < 0.001, Fig. 1e).

To delimit the QTL for SNS in this population, we performed ANOVAs for markers flanking FT-A2 (Table S2). Marker S3A_116,149,133 located 1.6 cM (8.0 Mb) distal to FT-A2 and marker S3A_194,830,543 located 3.5 cM (70.7 Mb) proximal to FT-A2 still showed highly significant differences in SNS, but both markers exhibited a decrease in the ANOVA F values relative to FT-A2 (14 % and 16 %, respectively). Based on these results, we delimited a 5.1 cM (78.7 Mb) confidence interval for the SNS QTL in this population including FT-A2.

**High resolution mapping of the SNS QTL on chromosome 3AS**

The previous results showed that the haplotypes associated with the FT-A2 D10 and A10 alleles have a significant effect on SNS. To narrow down the candidate gene region and explore the linkage between the FT-A2 D10A polymorphism and the differences in SNS, we generated a high-density map of the 3AS chromosome region in tetraploid wheat using a total of 3,161 BC1F3, BC1F4, and BC1F5 plants derived from the KxG population. These plants were screened in separate batches over three years using flanking markers 3A-117.83 and 3A-127.82 (numbers indicate coordinates in RefSeq v1.0 in Mb). Within this 9.9 Mb region including FT-A2 (124.17 Mb), we identified 76 recombination events corresponding to a genetic distance of 1.58 cM (6.26 Mb per cM). One of these recombination events (H2-6-#14-5) was detected in the progeny test of
primary recombinant H2-#6, which explains the presence of two close recombination events in this line (Table 4).

In addition to the molecular marker for the $FT-A2$ D10A SNP and the two flanking markers, we developed eight more KASP and CAPS markers in the candidate region (Table S1) and used them to genotype plants carrying recombination events in the region. The lines with the 10 closest recombination events to $FT-A2$ are presented in Table 4 together with the results of the field progeny tests for SNS. Progenies of the lines H2-#6 and H2-14#17-2 heterozygous for $FT-A2$ showed significant differences in SNS ($P < 0.01$) between lines homozygous for the two parental alleles, whereas progeny tests for the eight lines homozygous for $FT-A2$ did not show significant difference in SNS between parental alleles in the heterozygous flanking regions (Table 4). Average SNS were as expected, with the lines homozygous for the A10 allele having 1.3 more spikelets on average than the lines homozygous for the D10 allele.

The phenotype of the critical recombinant line #18-5 with the closest distal recombination event to $FT-A2$ was validated in a separate experiment in Davis in 2021 (Table S3). In this experiment, control lines showed highly significant differences in SNS ($P < 0.0001$) confirming that the differences in SNS were detectable in this experiment. By contrast, there was no significant difference between the sister lines with and without the recombination event #18-5, with both lines showing SNS values similar to the control line with the Gredho allele (Table S3). Taken together, these results confirmed that the causal gene for the 3AS QTL for SNS was proximal to the marker located at CS RefSeq v1.0 coordinate 120,227,651 (Table 4).

We identified an additional line (BC1F4 H2-18 #28-4) with a closer recombination event to $FT-A2$ in the proximal region between $FT-A2-R1$ and 3A-125.4, but we did not have enough grains to evaluate it with the other lines listed in Table 4. We planted a separate field experiment at Tulelake in the spring of 2020, in which we included homozygous sister lines #28-4-1 and #28-4-3 that were fixed for either the Kronos or Gredho alleles in the segregating proximal region (Table 5). As an additional control, we included sister lines derived from plant #17-2 (Table 4) that were either homozygous for the $FT-A2$ D10 (#17-2-18) or A10 allele (#17-2-22, Table 5). These two lines showed highly significant differences in SNS ($P < 0.0001$, Table 5) confirming that it was possible to detect differences between the two $FT-A2$ alleles in this experiment. By contrast, there was no significant difference between the H2-18 #28-4 recombinant sister lines,
confirming that the candidate gene was still linked to *FT-A2* (Table 5). Based on this result, we established a closer proximal flanking marker (3A-125.4), and reduced the candidate region for the 3AS QTL to a 5.2 Mb interval between coordinates 120,227,651 and 125,402,254 (Table 5).

**Genes in the candidate gene region for the 3AS QTL for SNS**

The annotated Chinese Spring reference genome region (RefSeq v1.1) between the two flanking markers defined in the previous section encompasses 28 high-confidence genes (including flanking genes *TraesCS3002G141000* and *TraesCS3002G143700*). The exome capture data revealed non-synonymous SNPs between Kronos and Gredho in only three out of the 28 genes, including the D10A polymorphism in *FT-A2*. The other two genes are described briefly below.

*TraesCS3A02G142200* encodes a leucine-rich repeat receptor-like protein kinase, so it is difficult to predict its potential effects. The predicted R872H amino acid change in Kronos (RefSeq v1.1 3AS 121,646,195) is in a conserved region close to the end of the protein (893 amino acids) and has a BLOSUM62 score of 0, predictive of a low probability of changes in protein structure or function. The R872H polymorphism was not detected in the parental lines LA95135 and SS-MVP57 of the hexaploid winter wheat populations segregating for the 3AS SNS QTL (Table S2), so we ruled out R872H as the causal polymorphism for the SNS phenotype.

*TraesCS3A02G143600* encodes a short peptide (104 amino acids) with a polymorphism in Kronos that generates a premature stop codon (S59*, RefSeq v1.1 3AS 125,094,949 C to A). However, the predicted protein in Gredho also seems to be truncated since it is much shorter (104 amino acids) than the orthologous protein in wild emmer (XP_037404892.1, 483 amino acids) or *T. urartu* (EMS53367.1, 348 amino acids). In addition, the 104 amino acids in Gredho showed no similarity to other plant proteins in the GenBank nr database in species outside the genus *Triticum*, suggesting that *TraesCS3A02G143600* encodes a non-functional protein in both Kronos and Gredho. Similar to R872H, the S59* polymorphism was not detected in winter lines LA95135 and SS-MVP57, providing additional evidence that this polymorphism is not critical for the SNS QTL on chromosome arm 3AS.
In summary, the D10A polymorphism in \textit{TraesCS3A02G143100} (FT-A2) was the only non-
synonymous SNP identified in the candidate gene region that co-segregated with the differences
in SNS in both the LA95135 x SS-MVP57 and Kronos x Gredho populations.

\textbf{Effect of the D10A polymorphism on FT-A2 interactions with 14-3-3 proteins}

Previous results have shown positive interactions between FT1 and six of the seven 14-3-3
proteins tested whereas FT-A2 did not interact with any of the 14-3-3 proteins (Li et al. 2015).
This was a puzzling result because all other four FT-like genes showed positive interactions with
at least one 14-3-3 protein. Since the original study was done using only the FT-A2 D10 allele,
we decided to explore the effect of the A10 allele. In this study, both FT-A2 proteins encoded by
the D10 and A10 allele failed to interact with any of the six tested 14-3-3 proteins, whereas the
FT1 positive control showed a strong interaction signal (Fig. S2). No autoactivation was
observed in the negative controls. Given the lack of interactions between both FT-A2 alleles and
any of the tested 14-3-3 protein, we have initiated Y2H screens to test if there are other protein
partners of FT-A2.

\textbf{Discussion}

\textbf{Candidate gene and causal polymorphism}

Spikelet number per spike is determined early after the transition from the vegetative to the
reproductive phase, when the spike meristem transitions into a terminal spikelet (Li et al. 2019).
This limits the influence of later environmental variability on SNS relative to GNS or grain
weight, which are affected by fertility, grain abortions, and conditions affecting grain filling until
the end of the season. As a result, SNS has a higher heritability ($h > 0.8$) than other yield
component traits (Kuzay et al. 2019; Zhang et al. 2018). This high heritability helped us to
Mendelize this trait and to develop a high-resolution map for the differences in SNS.

Using this high-density map, we delimited a 5.2 Mb candidate gene region on chromosome arm
3AS including 28 annotated high-confidence genes in CS, including three with non-synonymous
polymorphisms between Kronos (D10) and Gredho (A10): \textit{TraesCS3A02G142200} (R872H),
\textit{TraesCS3A02G143100} (D10A) and \textit{TraesCS3A02G143600} (S59*). To test if the S59* and
R872H polymorphisms were present in hexaploid varieties with the D10 allele, we compared the available sequences for this region in the wheat pangenome (Walkowiak et al. 2020). The 124,172,909-A allele (D10) was detected in CDC Landmark, Lancer, and Spelt, whereas the 124,172,909-C (A10) SNP was present in CS, Julius, Jagger, CDC Stanley, ArinaLRFor, Mace, Norin 61, and SY Mattis. The S59* and R872H polymorphisms were not detected in any of these hexaploid wheats, suggesting that these two SNPs originated in durum wheat, and that the A10 mutation occurred in a haplotype different from the one present in modern durum wheat varieties (henceforth S59*-R872H haplotype).

Based on the previous result, it was not surprising that the S59* and R872H polymorphisms were not detected in LA95135 (D10) and SS-MVP57 (A10), the parental lines of the hexaploid winter wheat population segregating for the 3AS SNS QTL. Gene TraesCS3A02G143600 showed no polymorphisms between LA95135 (D10) and SS-MVP57, whereas TraesCS3A02G142200 had only one synonymous polymorphism, suggesting that TraesCS3A02G143600 and TraesCS3A02G142200 are unlikely candidate genes for the SNS QTL. After the elimination of these two genes, FT-A2 is the only other gene in the candidate region that has a non-synonymous polymorphism (D10A) linked to the differences in SNS in both mapping populations. Since we only explored the coding regions, we cannot rule out the possibility of polymorphisms in regulatory regions within the candidate gene region affecting the number of spikelets per spike. However, the genetic data presented here, together with previously published results showing that loss-of-function mutations in FT-A2 affect SNS in wheat (Shaw et al. 2019), point to FT-A2 as the most likely candidate gene for the SNS QTL.

The D10A amino acid change in FT-A2 has a BLOSUM 62 score of -2 and is located in a conserved region of the protein, suggesting a high probability of an effect on either protein structure or function. To test if any other polymorphisms in FT-A2 were associated with the D10A polymorphism, we compared the available exons, introns, 5’ upstream region (5,000 bp) and 3’ downstream region (2,000 bp) of FT-A2 in genomic sequences of hexaploid wheat (Walkowiak et al. 2020). We did not find any additional SNPs to differentiate the varieties with the D10 allele (CDC Landmark, Lancer and Spelta) from those carrying the A10 allele (CS, Julius, Jagger, CDC Stanley, ArinaLRFor, Mace, Norin 61, and SY Mattis) in the analyzed region. Although we cannot completely rule out the possibility of polymorphisms located in regulatory regions outside the investigated region, the available evidence points to D10A as the
most likely causal polymorphism. A conclusive test of this hypothesis will require the editing of
the A124,172,909C, but this is not simple because this is a transversion, and currently available
plant gene editors are not efficient to edit transversions. New prime editing technologies
(Anzalone et al. 2019) may solve this problem once they become more efficient in plants (Lin et
al. 2020).

**Differential recombination rates within the candidate gene region**

The distribution of recombination events (RE) in the 10 Mb region between the flanking markers
used in this study was not uniform. In the 2.4 Mb distal to the candidate gene region (117.8 to
120.2 Mb, 14 genes), we detected 56 RE resulting in an average of 23.3 RE/Mb or 4.0 RE/gene.
In the 2.4 Mb proximal to the candidate region (125.4 to 127.8 Mb, 13 genes), we detected 20
RE resulting in a frequency of 8.3 RE/Mb or 1.5 RE/gene. Surprisingly, not a single RE was
detected in the 5.2 Mb central candidate region (120.2 to 125.4 Mb, 28 genes), despite being
twice as large and including twice the number of genes as the flanking regions. Recombination
events occur mainly in gene regions (Darrier et al. 2017), so we would have expected to find 39
of the 76 RE within the candidate region if RE were distributed proportionally to the number of
genes. The same number would be expected if RE were distributed proportionally to the length
of the interval.

To explore if this lack of recombination in the central region was caused by a structural
rearrangement, we used the sequenced genome of the tetraploid variety Svevo (Maccaferri et al.
2019) that showed the same SNPs as the Kronos exome capture across the candidate gene region.
Since Gredho showed very few polymorphisms with CS across the candidate gene region, we
compared the genomes of CS (A10) and Svevo (D10) in this region. In Svevo, we found
orthologs to the 28 high confidence genes present in CS, with the exception of
*TraesCS3A02G142500* that was present in the correct position and strand in Svevo (100%
identical over all its length) but was not annotated. All the genes were in the same orientation in
CS and Svevo, and the total length of the region was similar in both species (5.2 Mb), suggesting
that no major structural rearrangements occurred in the candidate gene region.

Finally, we did a BLAST comparison of all the Svevo genes to a Kronos scaffold assembly from
the Earlham Institute, U.K. and were able to detect 27 of the 28 genes with 100% identity. The
only exception was *TRITD3Av1G056250* (ortholog of CS *TraesCS3A02G142600*), for which we only detected the B-genome homeolog in Kronos. These results suggest the Kronos genome is not very different from Svevo in this region. We currently do not know the cause of the reduced recombination frequency between 121.5 and 125.1 Mb in the Kronos x Gredho population, but since no pseudomolecule assembly of Kronos or Gredho are available, we cannot rule out the possibility of structural rearrangements in this region in these two varieties.

**Effect of *FT-A2* D10A polymorphism on heading time and fertility**

Wheat varieties are selected to flower within a narrow time window to maximize grain productivity. This limits the introgression of loss-of-function alleles that have beneficial effects on SNS but generate large delays in heading time, such as those in *VRN1* (Li et al. 2019) or *PPD1* (Shaw et al. 2013). By contrast, the *FT-A2* A10 allele has a positive effect on SNS and limited effect on heading time. Even when loss-of-function mutations in *ft-A2* and *ft-B2* were combined in Kronos, the delay in heading time was only 2-4 days (Shaw et al. 2019). In this study, the D10A polymorphism showed small effects on DTH in the different genetic backgrounds, ranging from a non-significant difference in the initial Kronos x Gredho population (Table 2), a marginally non-significant difference of 0.8 d (*P* = 0.053) in the 2021 field experiment comparing Kronos isogenic lines, and an average difference of 1.7 d in the winter wheat population (Fig. 1A).

An important limitation for the utilization of the *ft-A2* loss-of-function mutation for wheat improvement was its negative effect on fertility (Shaw et al. 2019), which offset its positive effect on SNS. This motivated our initial search for *FT-A2* natural variants that separated the positive effects on SNS from the negative effects on fertility. Results presented in this study show that the positive effect of the A10 polymorphism on SNS were translated into positive effects on GNS in both the winter wheat population (Fig. 1e) and in the spring NILs (Table 3). These results suggest that the A10 allele is not associated with negative effect on fertility. This hypothesis was further supported by the higher number of grains per spike observed in the lines carrying the A10 allele in the different field experiments, although the differences were significant only in the two Kronos NILs evaluated in the field in 2020 (Table 3). These results provide a good example of the value of using natural variants selected by breeders to identify mutations that optimize specific traits with limited negative pleiotropic effects.
**FT-A2 effects on SNS, GNS, grain weight and spike yield**

It was encouraging to see that the positive effect of the A10 allele on SNS and GNS was expressed in both winter (Fig. 1) and spring wheats (Table 3), and among the latter in both spring and fall planted spring wheats. However, the magnitude of the increases in SNS, GNS and spike yield associated with the A10 allele varied among experiments, suggesting that the effects of this *FT-A2* polymorphisms on these traits are modulated by the environment. We also observed variable effects of the A10 polymorphisms on grain weight. Whereas no significant effects were detected for this trait in the experiments performed in UCD and Tulelake in 2020, we detected a significant reduction in grain weight in field experiment performed at UCD in 2021, which offset the gains in GNS (Table 3).

Similar observations have been reported for *WAPO-A1*, the causal gene of a wheat SNS QTL on the long arm of chromosome 7AL (Kuzay et al. 2019). Increases in SNS associated with the favorable *Wapo-A1b* allele were translated into significant increases in grain yield only when the favorable *WAPO-A1* allele was present in productive genetic backgrounds and the plants were grown in a favorable environment. When the *Wapo-A1b* allele was present in poorly adapted varieties or when the lines were grown under limiting watering conditions, the plants did not have enough resources to fill the extra grains, resulting in a negative correlation between grain number and grain weight that limited the gains in grain yield (Kuzay et al. 2019). A study with elite CIMYT lines also highlighted the importance of genetic-by-environment interactions on the trade-offs between grain number and grain weight (Quintero et al. 2018). We hypothesize that environmental differences between our 2020 and 2021 field trials may have contributed to the observed differences in grain weight, in spite of the positive effects of the A10 allele on SNS and GNS detected in both years (Table 3).

**FT-A2 as a candidate gene for previously published SNS QTLs on chromosome arm 3AS**

A QTL for DTH (*Qncb.HD-3A*) was previously mapped on chromosome 3A within a 400 Mb interval including *FT-A2* (DeWitt et al. 2021) in the LA95135 x SS-MVP47 population. We found in this study that LA95135 carries the D10 allele and SS-MVP47 the A10 allele, and after genotyping the population with the *FT-A2* marker, we found that the A10 allele was associated not only with a slight delay in heading time but also with higher SNS, GN, and grain yield per
spike (Fig. 1). The similar pleiotropic effects of the SNS QTL in the winter wheat population and
the Kronos x Gredho population, together with the overlapping mapping regions, suggest that the
\textit{FT-A2} D10A polymorphism may have contributed to the \textit{Qncb.HD-3A} identified in the
LA95135 x SS-MVP47 population.

An additional QTL for DTH was identified in the Avalon x Cadenza population (U.K.) on
chromosome arm 3AS around the peak marker BS00021976 (169 Mb RefSeq v1.0) (Martinez et
al. 2021). This QTL interval (60 Mb at each side of BS00021976) includes 536 annotated genes,
among which the authors proposed \textit{FT-A2} as a candidate of particular interest. Using our \textit{FT-A2}
marker, we established that both Avalon and Cadenza carry the A10 allele, so we conclude that
the D10A polymorphism is not the cause for the observed differences in DTH. Martinez et al.
(2021) suggested that differences in \textit{FT-A2} transcript levels may contribute to the differences in
DTH, but more precise mapping of the QTL will be necessary to support this hypothesis.

Several QTLs for grain yield components have been reported in different regions of chromosome
3AS in a recombinant inbred chromosome line from the cross between cultivar Cheyenne and a
substitution of chromosome 3A of Wichita in Cheyenne (CNN(Wichita-3A)) (Ali et al. 2011;
Campbell et al. 2003; Dilbirligi et al. 2006). QTLs for grain yield and grain number per square
meter were mapped in a region between markers \textit{barc86} and \textit{barc67} (54.4 to 464.3 Mb RefSeq
v1.0, “Region 2”) which encompasses the \textit{FT-A2} locus. However, both Cheyenne and
CNN(Wichita-3A) have the A10 allele of \textit{FT-A2} (Supplementary Appendix S1), suggesting that
a different gene (or a different polymorphism in \textit{FT-A2}) was the cause of this QTL.

\textbf{\textit{FT-A2} allele frequencies and breeding applications}

The \textit{FT-A2} alleles show contrasting frequencies in durum and common wheat, with the A10
allele present in less than 1% of the durum accessions and in 56% of the common wheat varieties
analyzed in this study (Table 1). We currently do not know if the A10 allele originated in the few
durum accessions carrying this allele in Oman and Turkey, or if these represent later
introgressions from hexaploid to tetraploid wheat. Either way, since the appearance or transfer of
the A10 allele to common wheat, its frequency increased rapidly suggesting that it was favored
by breeders in common wheat breeding programs.
The low frequency of the A10 allele in durum wheat could be a result of an hexaploid wheat origin combined with lack or infrequent transfers of genes from hexaploid to tetraploid wheat. However, it can also be the result of selection for larger grains and indirect selection for reduced GNS in environments showing a negative correlation between these two traits. Similar to FT-A2, the Wapo-A1a allele for low SNS is almost fixed in durum wheat, whereas the Wapo-A1b allele for high-SNS is found at high frequencies in hexaploid wheat (Kuzay et al. 2019). We interpret this similar asymmetric distribution of WAPO-A1 and FT-A2 alleles for SNS in common and durum wheat as indirect support to the hypothesis that selection for larger grains may result in indirect selection for reduced SNS.

Among hexaploid spring wheats, we also observed significant differences in the distribution of the FT-A2 alleles, with a larger frequency of the A10 allele among spring varieties developed under a long growing cycle (DuF, 58.4%) than among those developed under a short growing cycle (DuS, 34.4%). We speculate that longer cycles may provide more resources to fill the extra grains associated with the A10 allele, facilitating the translation of the difference in SNS into differences in grain yield. This in turn, may result in a stronger selection pressure for the A10 allele in the fall-planted programs. This idea is indirectly supported by the high frequency of the A10 allele among the US winter wheat varieties (Table 1, 81.7%). Additional experiments with D10 and A10 NILs in different genetic backgrounds tested in different spring-planted and fall-planted locations will be necessary to test this hypothesis.

The high frequency of the A10 allele in the winter wheats and fall-planted spring wheats provides additional evidence that this allele has positive effects in those regions. However, as the frequency of the A10 allele increases, the number of varieties that can benefit from its introgression decreases. By contrast, the A10 allele is almost absent from modern durum wheat breeding programs, and may represent a good opportunity to benefit a large proportion of the germplasm in the durum wheat programs. To facilitate the testing and introgression of the A10 allele into durum wheat breeding programs, we deposited the Kronos NIL with the A10 allele in the NSGC (PI 699107). Kronos, is a modern durum wheat variety with excellent pasta quality, which makes it a better donor parent than Gredho.

Our preliminary results suggest that the A10 allele may be more beneficial in fall planted than in the spring planted durum wheat programs, but additional experiments are necessary to test this
hypothesis. It will be also interesting to investigate the combined effect of the A10 allele with alleles from other genes that also result in increases in SNS such as Wapo-A1b (Kuzay et al. 2019) and the Elf3 allele from T. monococcum (Alvarez et al. 2016).

In summary, the genetic information provided in this study, together with the previous mutant information, provides strong evidence that FT-A2 is the causal gene for the differences in SNS, GNS, and spike yield associated with this region on chromosome arm 3AS. The identification of the likely causal polymorphism (D10A) and the development of a perfect marker for this polymorphism can accelerate the deployment of this favorable allele in wheat breeding programs worldwide.

Declarations

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Conflicts of interest/Competing interests

The authors declare no conflict of interests or competing interests.

Author contribution statement
PG conducted most of the experimental work and wrote the first version of the manuscript. JZ contributed experimental work and many of the statistical analysis. KL contributed the Y2H experiments. GBG and ND contributed the LA95135 x SS-MVP57 population and the corresponding genotypic and phenotypic data. JC contributed the frequency of the D10A polymorphism in Montana breeding lines. JD initiated and coordinated the project, contributed to data analyses, and supervised PG. All authors reviewed the manuscript and provided suggestions.

Availability of data and materials

All data and materials described in this paper are available from the corresponding author upon request. The FT-A2 introgression in Kronos is being deposited in the National Small Grain Collection (PI 699107). PI accession numbers are provided for all germplasm used when available. The datasets retrieved and analyzed during the current study are available in the T3/Wheat exome capture database (https://wheat.triticeaetoolbox.org/).

Code availability

Not applicable.
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### Table 1. Frequency of the FT-A2 alleles in different germplasm collections

| Species                        | Ploidy | No. acc. | A10 % | D10 % | A10 | D10 |
|-------------------------------|--------|----------|-------|-------|-----|-----|
| *T. urartu*                   | 2x     | 89       | 0.0%  | 100.0%| 0   | 89  |
| *T. turgidum* ssp. *dicocoides* | 4x     | 82       | 0.0%  | 100.0%| 0   | 82  |
| *T. turgidum* ssp. *dicoccon*  | 4x     | 32       | 0.0%  | 100.0%| 0   | 32  |
| *T. turgidum* ssp. *durum*    | 4x     | 417      | 0.7%  | 99.3% | 3   | 414 |
| *T. aestivum* Exome capture<sup>a</sup> | 6x     | 238      | 59.7% | 40.3% | 142 | 96  |
| *T. aestivum* US winter wheats<sup>b</sup> | 6x     | 126      | 81.7% | 18.3% | 103 | 23  |
| *T. aestivum* Spring DUF<sup>c</sup> | 6x     | 149      | 58.4% | 41.6% | 87  | 62  |
| *T. aestivum* Spring DUS<sup>d</sup> | 6x     | 192      | 34.4% | 65.6% | 66  | 126 |

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<sup>a</sup> He et al. 2019
<sup>b</sup> T3/Wheat
<sup>c</sup> Zhang et al. 2018
<sup>d</sup> Zhang et al. 2018 + 99 breeding lines from MT
Table 2. Effects of \textit{FT-A2}, \textit{PPD-A1} and \textit{RHT-B1} on plant height (HT), days to heading (DTH) and spikelet number per spike (SNS). Three-way ANOVA with \textit{P} values of the main effects and least-square means (LSmeans). Error bars are s.e.m. ns = not significant, * = \textit{P} < 0.05, ** = \textit{P} < 0.01, *** = \textit{P} < 0.001. All the interactions were non-significant.

|           | Kronos (D10) | Gredho (A10) | P value | Kronos | Gredho | P value | Kronos | Gredho | P value |
|-----------|--------------|--------------|---------|--------|--------|---------|--------|--------|---------|
| \textit{FT-A2} | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) |
| LSmear ± s.e.m. | 113.4 ± 3.2 | 130.5 ± 0.9 | 25.1 ± 0.5 | 118.3 ± 2.2 | 130.6 ± 0.6 | 26.7 ± 0.3 |
| Three-way ANOVA | ns | ns | * |
| \textit{PPD-A1} | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) |
| LSmear ± s.e.m. | 108.1 ± 2.3 | 120.8 ± 0.6 | 22.6 ± 0.4 | 121.6 ± 2.6 | 141.0 ± 0.7 | 29.6 ± 0.4 |
| Three-way ANOVA | *** | *** | *** |
| \textit{Rht-B1} | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) | Plant height (HT, cm) | Days to heading (DTH) | Spikelet No./spike (SNS) |
| LSmear ± s.e.m. | 97.1 ± 2.5 | 131.5 ± 0.7 | 26.2 ± 0.5 | 131.4 ± 2.8 | 130.2 ± 0.8 | 25.8 ± 0.4 |
| Three-way ANOVA | *** | * | ns |
Table 3. Comparisons of Near isogenic lines with the \textit{FT-A2} A10 and D10 alleles in field experiments at UC Davis in 2020 and 2021.

| Davis 2020 | Allele | N  | SNS  | GN  | Grains/spikelet | GW mg | Yield / spike g |
|------------|--------|----|------|-----|-----------------|-------|-----------------|
| H2-14 D10 | 10 \(^a\) (54 spikes) | 20.27 | 59.22 | 2.92 | 55.81 | 3.31 |
| H2-14 A10 | 10 \(^a\) (38 spikes) | 21.92 | 70.77 | 3.23 | 56.78 | 4.07 |
| A10 % increase | 7.9 % | 19.6 % | 10.6 % | 1.8 % | 22.9 % |
| \(t\)-TEST | 0.0016 | 0.0004 | 0.0016 | 0.55 | 0.0018 |
| H2-23 D10 | 10 \(^a\) (39 spikes) | 19.36 | 56.31 | 2.89 | 54.47 | 3.07 |
| H2-23 A10 | 10 \(^a\) (38 spikes) | 23.11 | 81.06 | 3.51 | 54.21 | 4.41 |
| A10 % increase | 19.1 % | 44.0 % | 21.5 % | -0.6 % | 43.2 % |
| ANOVA \(P\) | <0.0001 | <0.0001 | 0.0003 | 0.88 | 0.0002 |

| Tulelake 2020 |
|----------------|
| H2-14 D10 | 27 spikes | 17.15 | 44.11 | 2.57 | 38.26 | 1.69 |
| H2-14 A10 | 23 spikes | 17.83 | 46.48 | 2.61 | 40.49 | 1.88 |
| % increase | 4.0 % | 5.4 % | 1.4 % | 5.8 % | 10.5 % |
| ANOVA \(P\) | 0.0004 | 0.2788 | 0.7665 | 0.2487 | 0.1348 |

| Davis 2021 |
|-------------|
| BC\(_4\)F\(_2\)A | 12 \(^b\) (96 spikes) | 18.52 | 67.85 | 3.67 | 60.34 | 4.09 |
| BC\(_4\)F\(_2\)D | 12 \(^b\) (96 spikes) | 19.58 | 72.15 | 3.69 | 55.61 | 4.01 |
| % difference | 5.7 % | 6.3 % | 0.5 % | -7.8 % | -2.0 % |
| ANOVA \(P\) | 0.0011 | 0.0168 | 0.7919 | 0.0002 | 0.2883 |

\(^a\) Experimental units were 1 m rows, with 3-5 spikes measured per row.
\(^b\) Experimental units were 4 row plots (1.86 m\(^2\)), with 8 spikes measured per plot.
**Table 4** Critical recombinant BC$_1$F$_5$ from Davis 2019-2020 field seasons. All lines except recombinant H2 #6 were evaluated in the 2019 field season. Comparisons of SNS for statistical significance are only between sister lines segregating for the heterozygous region.

| Marker     | Chr. 3AS CS  | H2  | H2-6 | H2-14 | H2-23 | D12  | 11-1 |
|------------|--------------|-----|------|-------|-------|------|------|
|            |              | #6  | #14-5| #17-2 | #1-3  | #18-5| #47-1|
| 3A-117.83  | 117,828,272  | H   | H    | H     | H     | K    | H    |
| 3A-120.23  | 120,227,651  | H   | G    | H     | K     | H    | K    |
| 3A-121.48  | 121,482,459  | H   | G    | H     | K     | G    | K    |
| 3A-121.65  | 121,646,195  | H   | G    | H     | K     | G    | K    |
| 3A-122.54  | 122,540,617  | H   | G    | H     | K     | G    | K    |
| FT-A2-L4   | 122,542,102  | H   | G    | H     | K     | G    | K    |
| SNS PHENO. | 124,172,909  | H   | G    | H     | K     | G    | K    |
| FT-A2-R1   | 125,094,949  | H   | G    | H     | K     | G    | K    |
| 3A-125.40  | 125,402,254  | H   | G    | H     | K     | G    | K    |
| 3A-126.57  | 126,567,437  | K   | K    | H     | K     | G    | K    |
| 3A-127.82  | 127,821,835  | K   | K    | K     | K     | G    | H    |

| Number of plants in PT | 34 | 83 | 71 | 72 | 79 | 70 | 74 | 75 | 80 | 81 |
| SNS Avg. Gredho allele (G) | 22.1 | 23.9 | 23.5 | 22.4 | 24.2 | 22.4 | 22.7 | 24.1 | 22.3 | 23.1 |
| SNS Avg. Kronos allele (K) | 21.6 | 23.2 | 21.7 | 22.5 | 23.0 | 21.9 | 22.4 | 24.3 | 21.7 | 22.4 |
| P values K vs G | 3e-05 | NS | 0.004 | NS | NS | NS | NS | NS | NS | NS |
Table 5: Spikelet number per spike (SNS) evaluation of BC1F6 homozygous sister lines from recombinant line H2-18 #28-4 in Tulelake 2020. Sister line #28-4-#1 carried a proximal Kronos chromosome segment and sister line #28-4-#3 a proximal Gredho chromosome segment. Lines #17-2-18 (FT-A2 D10) and #17-2-22 (FT-A2 A10) were included as controls.

| Marker   | Chr.3AS CS | H2-18  | H2-18  | H2-14  | H2-17 |
|----------|------------|--------|--------|--------|-------|
|          |            | #28-4-1 | #28-4-3 | #17-2-18 | #17-2-22 |
| 3A-117.82| 117,828,272| G      | G      | K      | G     |
| 3A-120.2 | 120,227,651| G      | G      | K      | G     |
| 3A-121.4 | 121,482,459| G      | G      | K      | G     |
| 3A-121.64| 121,646,195| G      | G      | K      | G     |
| 3A-122.540| 122,540,617| G     | G      | K      | G     |
| FT-A2-L4 | 122,542,102| G      | G      | K      | G     |
| FT-A2   | 124,172,909| G      | G      | K      | G     |
| **SNS PHENO.** | | G      | G      | K      | G     |
| FT-A2-R1| 125,094,949| G      | G      | K      | G     |
| 3A-125.4| 125,402,254| K      | G      | K      | G     |
| 3A-126.5| 126,567,437| K      | G      | K      | G     |
| 3A-127.8| 127,821,835| K      | G      | K      | K     |

| Number of plants | 40 | 42 | 43 | 40 |
| SNS Avg          | 17.68 | 17.87 | 16.94 | 17.94 |

$P$ values D10 (K) vs A10 (G) | 0.287 | 1.78E-09 |
**Figure legends**

**Fig. 1** Comparison between *FT-A2* A10 (SS-MVP57) and D10 (LA95135) alleles in winter wheat. 

- **a** Days to heading. 
- **b** Spikelet number per spike. 
- **c** Grain number per spike. 
- **d** Grain number per spikelet (fertility). 
- **e** Average spike yield. Bars are least square means from a factorial ANOVA including *PPD*-D1, *RHT*-D1 and *WAPO*-A1 as factors. Error bars are s.e.m. 

*ns* = not significant, *P* = 0.05, **P** = 0.01, ***P** = 0.001.
Figure 1

Comparison between FT-A2 A10 (SS-MVP57) and D10 (LA95135) alleles in winter wheat. a Days to heading. b Spikelet number per spike. c Grain number per spike. d Grain number per spikelet (fertility). e
Average spike yield. Bars are least square means from a factorial ANOVA including PPD-D1, RHT-D1 and WAPO-A1 as factors. Error bars are s.e.m. ns= not significant, * P = 0.05, ** P = 0.01, *** P = 0.001.

**Supplementary Files**

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- SupplementaryAppendix1GlennDubcovsky.xlsx
- SupplementaryTablesandFiguresGlennDubcovsky.pdf