

Introgression of Novel Traits from a Wild Wheat Relative Improves Drought Adaptation in Wheat^{1[W]}

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Root architecture traits are an important component for improving water stress adaptation. However, selection for aboveground traits under favorable environments in modern cultivars may have led to an inadvertent loss of genes and novel alleles beneficial for adapting to environments with limited water. In this study, we elucidate the physiological and molecular consequences of introgressing an alien chromosome segment (7DL) from a wild wheat relative species (*Agropyron elongatum*) into cultivated wheat (*Triticum aestivum*). The wheat translocation line had improved water stress adaptation and higher root and shoot biomass compared with the control genotypes, which showed significant drops in root and shoot biomass during stress. Enhanced access to water due to higher root biomass enabled the translocation line to maintain more favorable gas-exchange and carbon assimilation levels relative to the wild-type wheat genotypes during water stress. Transcriptome analysis identified candidate genes associated with root development. Two of these candidate genes mapped to the site of translocation on chromosome 7DL based on single-feature polymorphism analysis. A brassinosteroid signaling pathway was predicted to be involved in the novel root responses observed in the *A. elongatum* translocation line, based on the coexpression-based gene network generated by seeding the network with the candidate genes. We present an effective and highly integrated approach that combines root phenotyping, whole-plant physiology, and functional genomics to discover novel root traits and the underlying genes from a wild related species to improve drought adaptation in cultivated wheat.

During the Green Revolution, high-yielding semi-dwarf wheat (*Triticum aestivum*) and rice (*Oryza sativa*) cultivars were bred for improved responsiveness to irrigation and fertilizer (Borlaug, 2008). In wheat, this was made possible by *Rht* dwarfing alleles, which improved harvest index and reduced lodging under favorable conditions (Hedden, 2003). However, selection for the aboveground traits and yield under optimal environments may have led to a reduction in root parameters that may be adaptive to environments with limited water. Root biomass in modern high-yielding cultivars from the International Maize and Wheat Improvement Center is lower than in old tall landraces (Waines and Ehdaie, 2007). This difference was hypothesized to be consistent with an inadvertent selection of smaller, more compact root systems under optimal conditions.

As agriculture is increasingly shifted to marginal lands and drought events become more frequent and intense, specific root morphological traits that can improve drought tolerance and sustain yields in suboptimal

conditions hold immense potential. To effectively incorporate drought-adaptive root traits in crops like wheat and rice, we need to better understand the genetic and physiological basis of adaptive root traits and explore the novel alleles or genes that were likely lost during domestication or through breeding for yield under optimal environments. Considerable genetic variation exists for root-related traits in major crops such as wheat, rice, and maize (*Zea mays*) in both the domesticated species as well as wild relatives (Reynolds et al., 2007; Ribaut et al., 2009; Gowda et al., 2011).

Here, we focus on the potential of wild wheat species to contribute to adaptive root traits under water limitations. Introduction of alien chromosome segments from wild relatives into wheat has resulted in improved pest resistance and yield (Rabinovich, 1998; Singh et al., 1998; Reynolds et al., 2001; Monneveux et al., 2003; Waines and Ehdaie, 2007). Rye-wheat translocations, for example, have been used to introduce traits for tolerance to abiotic and biotic stresses, resulting in improved yields in some cases (Zeller, 1973; Waines and Ehdaie, 2007). Genes providing resistance to biotic stresses such as stem rust (*Puccinia graminis*), powdery mildew (*Blumeria graminis* f. sp. *tritici*), and greenbug have been successfully introgressed from rye (*Secale cereale*) into modern wheat germplasm (Friebe et al., 1996; Graybosch, 2001; Waines and Ehdaie, 2007; Yediay et al., 2010). In addition to pathogen resistance, the rye-wheat translocation (1RS) has been reported to have significant effects on yield, root

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morphology, and water and nitrogen use efficiency (Rabinovich, 1998; Ehdaie et al., 2003, 2012; Sharma et al., 2010). The presence of the 1RS translocation reduced central metaxylem diameter, a trait that has been associated with drought tolerance in wheat (Richards and Passioura, 1989; Sharma et al., 2010).

Introgressions from *Agropyron elongatum* into bread wheat have improved pest resistance, biomass, and yield under irrigated conditions (Monneveux et al., 2003). Specifically, chromosome 7E of *A. elongatum* has been a valuable source of genetic variation for bread wheat, improving yield and providing resistance to barley yellow dwarf virus, stripe rust (*Puccinia striiformis*), and leaf rust (*Puccinia triticina*) resistance (Sharma et al., 1995; Ma et al., 2000; Zu-Jun and Zheng-Long, 2001; Monneveux et al., 2003). In addition to harboring pathogen resistance genes, several *Agropyron* species and wheat \times *A. elongatum* lines have been reported to enhance tolerance to abiotic stresses, including drought, waterlogging, and salinity (Dvorák et al., 1988; Wang and Zhang, 1996; McDonald et al., 2001). In hexaploid wheat, introgressions of 7DL.7EL, originally used to introduce leaf rust resistance, have been reported to increase biomass and yield in various wheat backgrounds (Sears, 1956; Reynolds et al., 2001).

The reported yield increase associated with the 7DL.7EL translocation lines relative to control genotypes and the presence of genes improving tolerance to abiotic stresses on 7EL led us to investigate this specific chromosomal segment for traits associated with yield (Singh et al., 1998; Monneveux et al., 2003; Zhang et al., 2005). We hypothesized that the 7DL.7EL translocation could be improving yields and water use efficiency in some environments due to adaptive root traits introduced through the *A. elongatum* chromosomal segment. It is possible that 7EL translocations could be either introducing novel root-related genes into wheat from the wild relative or incorporating alleles lost during intense selection under optimal environment. The original 7DL.7EL translocation carried locus Y for yellow flour, which limited its use in breeding. For this study, we chose a newly engineered version of the 7DL.7EL translocation with Y removed (Zhang et al., 2005), the recipient parent spring wheat line, and sister lines of the translocation line served as additional controls. Our experiments aim to address whether the 7DL.7EL translocation improves root traits. If it does, then what are the physiological consequences of the altered root traits under water-limiting environments? We next explore the molecular basis of altered root traits to identify underlying genes or alleles that can be used to improve wheat yields in marginal environments. We present an effective approach for addressing these questions by integrating root phenotyping, with whole plant physiology, transcriptomics, and computational biology tools, to discover novel root traits and candidate genes underlying these traits.

RESULTS

A. elongatum Translocation Introduces a Drought-Adaptive Response in Wheat

We hypothesized that the differences in biomass and yield between the *A. elongatum* 7DL lines and the corresponding parental genotypes could be due to increased investment in root production, thus increasing the acquisition of water and nutrients from the soil. We used a parent line, Pavon76 (P76), the 1-96-1 translocation line (TL), and a negative control (NC) for the translocation event for our experiments (Fig. 1). Phenotypic screening at an early seedling stage showed that reduced water availability had negative effects on several root traits in P76 and NC. Primary root length was significantly reduced at 4 and 6 d in P76 and NC ($P \leq 0.05$; Table I), while significant reductions in seminal root length were observed in these genotypes at 2, 4, and 6 d when compared with well-watered conditions ($P \leq 0.05$; Supplemental Table S1). Furthermore, the number and density of lateral roots on the primary and seminal root axis of P76 and NC under limited-water conditions were significantly less than in well-watered conditions at 4 and 6 d ($P \leq 0.01$; Table I; Supplemental Table S1). Water deficit also reduced shoot length in P76 and NC at day 6 ($P \leq 0.05$; Table I). In striking contrast, TL exhibited no significant reductions in lateral root number, density, or the length of primary or seminal roots at any time point in response to water stress. The shoot length in TL did not decrease significantly during water stress. These data suggest that the *A. elongatum* translocation could influence both root and shoot traits when plants are water stressed. Similar results were observed when the experiment was repeated with one-tenth-strength

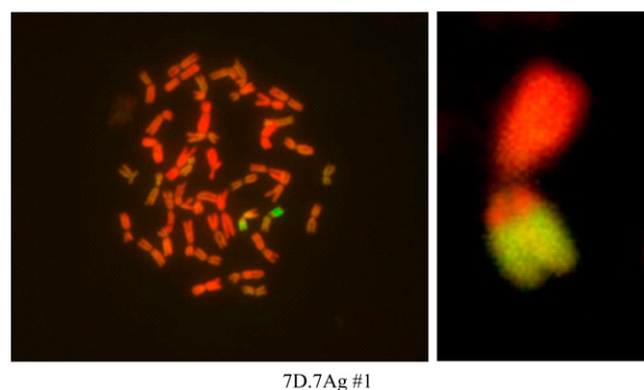


Figure 1. Visualization of wheat-*A. elongatum* translocation. An International Maize and Wheat Improvement Center hexaploid wheat cultivar, P76, carrying a chromosome 7 segment from *A. elongatum* (7Ag.7DL) is visualized by genomic in situ hybridization. In both images, green staining represents the translocated segment from *A. elongatum*. Wheat genomic probe was directly labeled with rhodamine (red). We performed root system architecture analysis on the background parent P76 as well as TL and NC.

Table 1. Phenotypic evaluation of the early seedling root system

Phenotyping of Spring Wheat genotypes P76, TL, and NC was done using the cigar roll method. The total number of lateral roots (more than 1 mm) on the primary root axis was counted. Lateral root density was calculated by dividing the number of lateral roots on the primary root axis by the primary root length. Statistical significance was determined using a one-way ANOVA using Tukey's method. Asterisks indicate significant differences observed between treatments for each genotype as follows: *** $P \leq 0.001$, ** $P \leq 0.01$, * $P \leq 0.05$. SE values are shown. nd, Not detected. This method was repeated three times using 30 wheat seedlings per cultivar per experiment.

Parameter	P76			TL			NC		
	2 d	4 d	6 d	2 d	4 d	6 d	2 d	4 d	6 d
Lateral root number									
Well watered	nd	4.6 ± 0.3	23.0 ± 0.5	nd	5.0 ± 0.3	26.6 ± 0.6	nd	4.8 ± 0.2	23.4 ± 0.9
Limited water	nd	1.3 ± 0.6**	8.8 ± 1.9***	nd	5.1 ± 0.9	27.6 ± 2.3	nd	1.2 ± 0.6**	8.8 ± 1.1***
Primary root length (mm)									
Well watered	7.6 ± 0.2	19.2 ± 0.3	25.0 ± 0.6	8.3 ± 0.2	19.8 ± 0.2	27.1 ± 0.8	8.1 ± 0.2	18.5 ± 0.5	25.1 ± 0.9
Limited water	7.2 ± 0.2	14.1 ± 1.0***	18.7 ± 1.9*	7.2 ± 0.7	18.2 ± 0.9	27.5 ± 0.6	7.4 ± 0.3	14.4 ± 0.4***	19.5 ± 0.6***
Lateral root density (mm ⁻¹)									
Well watered	nd	0.24 ± 0.02	0.92 ± 0.01	nd	0.25 ± 0.02	1.00 ± 0.03	nd	0.26 ± 0.02	0.93 ± 0.01
Limited water	nd	0.08 ± 0.04**	0.47 ± 0.08***	nd	0.28 ± 0.05	1.00 ± 0.08	nd	0.08 ± 0.04**	0.44 ± 0.05***
Shoot length (mm)									
Well watered	2.9 ± 0.2	8.3 ± 0.6	12.5 ± 0.6	3.3 ± 0.2	9.2 ± 0.6	12.7 ± 0.7	3.2 ± 0.2	8.6 ± 0.8	13.1 ± 0.8
Limited water	2.4 ± 0.2	6.8 ± 0.5	9.7 ± 1.0*	2.9 ± 0.1	8.5 ± 0.6	12.3 ± 0.5	3.0 ± 0.2	7.3 ± 0.5	9.8 ± 0.5*

Hoagland solution to ensure any confounding effects of nutrient availability (Supplemental Table S2). Collectively, these observations suggested that the presence of a 7DL.7EL segment in the P76 background could be conferring an adaptive response under water limitation in the translocation line. The results from early-seedling phenotyping prompted us to investigate the differences in root architecture and shoot photosynthetic traits in older plants.

TL Lines Have Higher Shoot and Root Biomass under the Limited-Water Condition

To observe the effect of the 1-96-1 translocation on root and shoot phenotypes at the pretillering stage, we sampled plants at 18 d after germination. The root and shoot biomass of P76 and NC plants under water stress decreased significantly ($P \leq 0.01$; Fig. 2; Table II). However, in TL, water stress did not significantly alter root or shoot biomass. Under water stress, TL had higher root ($P \leq 0.01$) and shoot ($P \leq 0.001$) biomass compared with P76 and NC (Fig. 2; Table II). Both root and shoot length measured slightly higher in TL under water stress relative to well-watered TL plants. Under well-watered conditions, there were no differences in root and shoot lengths and dry weights among genotypes. The phenotypic measurements indicate that the 1-96-1 translocation alters root traits but not at the cost of shoot biomass under water-limiting conditions.

Stomatal Factors Limit Photosynthesis in P76 and NC during Water Deficit

To elucidate the physiological impact of higher root biomass in TL during water stress, the CO₂ assimilation and gas-exchange rates were measured for the three genotypes. Carbon assimilation was different among the three genotypes under limited-water conditions (Fig. 3).

Photosynthetic rates were reduced by 45% and 46% in P76 and NC ($P \leq 0.001$), respectively, but did not change significantly in TL (Fig. 3A). Water deficit also severely impacted stomatal conductance and transpiration, reducing stomatal conductance by 50% and 56% ($P \leq 0.001$) and transpiration rate by 39% and 49% ($P \leq 0.001$) compared with well-watered conditions in P76 and NC, respectively (Fig. 3, B and C). Stomatal conductance was also reduced ($P \leq 0.001$) in TL, but only by 16%. No reductions in transpiration were observed for TL during water deficit. Clearly, TL showed lower sensitivity to water limitation compared with P76 and NC.

Reductions in gas exchange and carbon assimilation are common in drought-stressed plants due to decrease

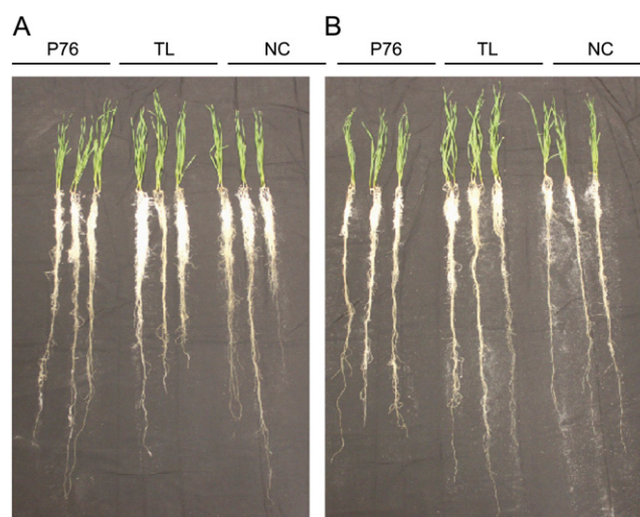


Figure 2. Harvested spring wheat P76 genotypes grown in the greenhouse under well-watered (A) and limited-water (B) environments. The images show 18-d-old wheat plants (pretillering stage) grown in tubes containing fine sand. The genotypes are indicated.

Table II. Summary of shoot and root lengths and biomasses of Spring Wheat genotypes P76, TL, and NC

The lengths were measured at the time of harvest, while the shoot and root dry mass were weighed separately after 5 d of incubation in a 60°C chamber. Statistical significance was determined using a one-way ANOVA using Tukey's method. Asterisks indicate significant differences observed between treatments for each genotype as follows: *** $P \leq 0.001$, ** $P \leq 0.01$. SE values are shown. This method was repeated three times using 35 wheat plants per cultivar per experiment.

Genotype	Shoot		Root	
	Well Watered	Limited Water	Well Watered	Limited Water
Dry weight (mg)				
P76	366.6 ± 57.8	209.5 ± 30.0***	343.6 ± 56.1	192.2 ± 18.2***
TL	327.0 ± 51.1	336.4 ± 53.1	335.7 ± 81.0	290.2 ± 44.4
NC	320.9 ± 16.9	219.5 ± 30.5**	318.9 ± 54.1	206.7 ± 39.4***
Length (cm)				
P76	35.2 ± 13.1	24.8 ± 2.0	76.2 ± 6.8	71.1 ± 3.8
TL	26.1 ± 2.3	27.5 ± 2.8	75.6 ± 4.0	79.6 ± 7.0
NC	27.6 ± 3.0	26.3 ± 1.9	79.8 ± 10.6	78.4 ± 2.9

in stomatal aperture. However, other aspects of photosynthesis, such as light utilization and photosynthetic electron transport, can also be impacted by drought stress. Therefore, we also measured chlorophyll fluorescence at the pretillering stage. Calculation of maximum quantum efficiency of open PSII reaction centers (F_v'/F_m') and the amount of energy used to drive photosynthesis (photochemical quenching) or dissipated as heat (nonphotochemical quenching) from fluorescence data are described in "Materials and Methods." In well-watered conditions, no differences in F_v'/F_m' , photochemical quenching, or nonphotochemical quenching were observed among the genotypes (Fig. 3, D–F). Photochemical quenching increased under limited water by 12% and 9% in P76 and NC, respectively, but not in TL.

In addition to stomatal aperture and photosynthetic electron transport, biochemical components of carbon assimilation, such as the carboxylation of ribulose-1,5-bisphosphate (RuBP) or the rate at which RuBP is regenerated, contribute to the photosynthetic capacity of a plant. To assess whether biochemical factors were contributing to the decline in CO_2 assimilation in P76, NC, and TL during water deficit, CO_2 assimilation was measured at a series of CO_2 concentrations and plotted against intercellular CO_2 , and the maximum rate of carboxylation by Rubisco (V_{cmax}) and the RuBP regeneration rate (J_{max}) were calculated according to the methods outlined by Dubois et al. (2007). Under limited-water conditions, J_{max} declined significantly by 31% and 24% in P76 and NC, respectively ($P \leq 0.05$; Fig. 4; Table III) but did not change in TL. V_{cmax}

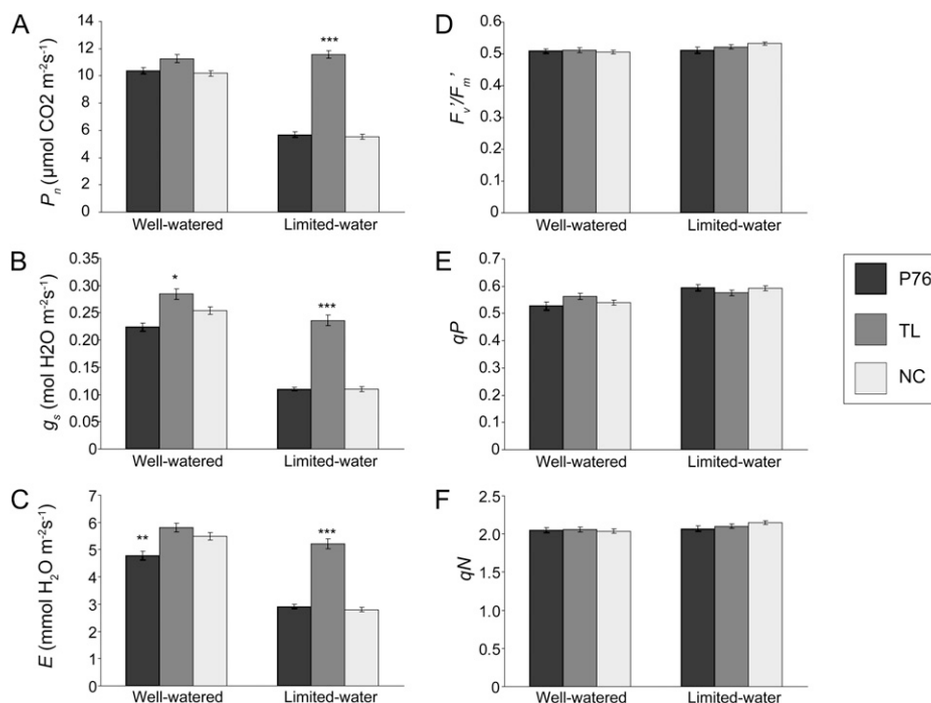


Figure 3. Physiological analyses to assess the response by each genotype under well-watered and limited-water conditions. A to C, Photosynthetic rate (P_n ; A), stomatal conductance (g_s ; B), and transpiration rate (E ; C) measurements using the LI-COR 6400 for 18-d-old posttransplanted spring wheat plants. D to F, F_v'/F_m' (D), photochemical quenching (q_p ; E), and nonphotochemical quenching (q_n ; F). Experiments were performed three times independently. At least nine individual leaves for each genotype in each experiment were measured. Error bars represent SE. Statistical significance was determined using a one-way ANOVA using Tukey's method. Asterisks represent statistically significant differences between genotypes within a treatment: *** $P \leq 0.001$, ** $P \leq 0.01$, * $P \leq 0.05$. The genotypes are indicated in the key at right.

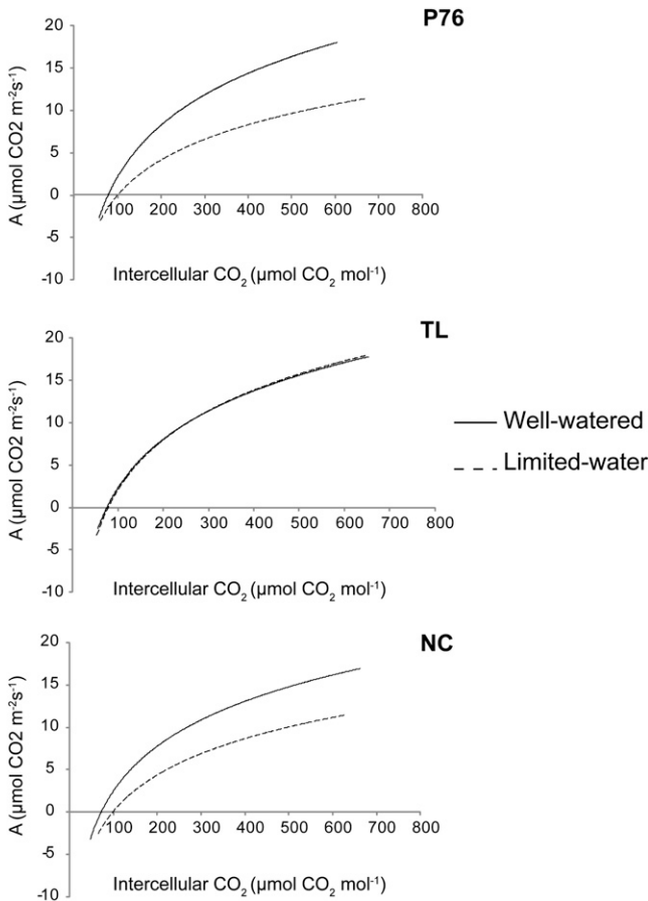


Figure 4. A/C_i curve analysis of P76, TL, and NC under well-watered and limited-water environments. The CO_2 assimilation rate (A ; $\mu\text{mol m}^{-2} \text{s}^{-1}$) was measured at light saturation (photosynthetically active radiation = $1,000 \mu\text{mol m}^{-2} \text{s}^{-1}$) and recorded at different chamber CO_2 concentrations. Experiments were performed three times independently. Six leaf samples per genotype per experiment were used for measurements.

was reduced by 31% in P76 ($P \leq 0.05$). However, it increased in TL by 18%. Although a 7% reduction in V_{cmax} was observed in NC, the difference was not significant. The results suggest that the major cause of photosynthetic decline in P76 and NC during water stress could be RuBP regeneration rate.

TL Differentially Regulates Genes Associated with Root Architecture

To elucidate the molecular mechanism underlying the increased root biomass phenotype observed in TL during water stress, we analyzed the root transcriptome of the three genotypes at the pretillering stage (18 d) under well-watered and water-stressed conditions. Lists of differentially expressed genes from various comparisons are provided as Supplemental Table S3 and Supplemental File S1. Several genes known to play roles in root development were differentially expressed between the TL and P76/NC lines in limited-water samples. One of these genes is a wheat ortholog of the *KNOTTED-LIKE3* (*KNAT3*) homeobox gene in *Arabidopsis thaliana*. *KNAT3* was down-regulated in roots and shoots of TL compared with P76 and NC under limited-water conditions (Fig. 5, D and G). In *Arabidopsis*, *KNAT3* has been proposed to act as a negative regulator of lateral root development (Truernit et al., 2006; Truernit and Haseloff, 2007). *KNAT3* expression level differences among the genotypes are notable in the context of increased lateral root formation in TL under water stress. *KNAT3* transcript abundance in early seedlings was lower ($P \leq 0.001$) at 4 d in TL compared with P76 and NC (Fig. 5A). At 6 d after germination, *KNAT3* was down-regulated in TL compared with P76 at 2 d, while *KNAT3* expression was slightly up-regulated in both P76 and NC when compared with P76 at 2 d. No genotypic differences were observed at 2 d after germination.

A wheat *E2F-RELATED* (*E2F*) transcription factor is down-regulated in roots and shoots of TL under water-limited conditions compared with P76 and NC (Fig. 5, E and H). *E2F* proteins are a family of transcription factors that regulate cell cycle progression in plants and animals. In *Arabidopsis*, *E2F* is known to play an antagonistic role in cell division and is a negative regulator of lateral root formation (del Pozo et al., 2002; Ramirez-Parra et al., 2004). We validated the expression of *E2F* using quantitative PCR (qRT-PCR) in pretillering stage and early seedling stage plants (Fig. 5, B, E, and F). Water stress caused a significant reduction ($P \leq 0.05$) in *E2F* expression in both TL and P76 during the pretillering stage, while no significant reductions were observed in NC when compared with well-watered plants. However, at the pretillering

Table III. V_{cmax} and J_{max}

Summary of CO_2 assimilation (A/C_i) curve estimates of V_{cmax} and J_{max} of P76, TL, NC using the methods described by Dubois et al. (2007). Asterisks denote statistical differences between treatments for each genotype: * $P \leq 0.05$.

Genotype	V_{cmax}		J_{max}		r^2	
	Well Watered	Limited Water	Well Watered	Limited Water	Well Watered	Limited Water
P76	69.6 ± 3.8	50.1 ± 2.1*	93.9 ± 4.7	64.3 ± 2.3*	0.86	0.92
TL	63.1 ± 2.6	74.9 ± 2.7*	90.6 ± 4.1	90.8 ± 2.2	0.90	0.95
NC	64.4 ± 4.1	59.8 ± 4.7	83.2 ± 3.9	63.0 ± 2.5*	0.86	0.92

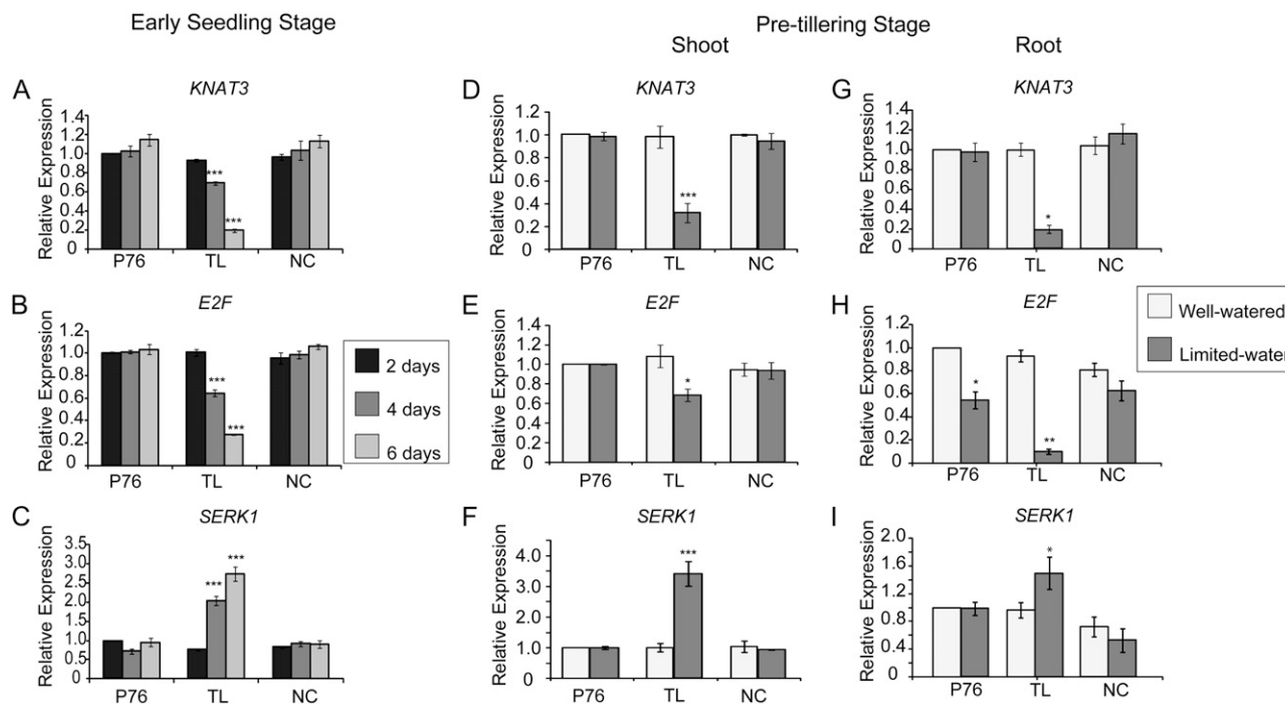


Figure 5. Gene expression analysis of root-related genes by qRT-PCR. A to C, Expression of microarray genes in early seedling stages using root tissues collected at 2, 4, and 6 d. The genotypes are indicated. P76 at 2 d was used as a reference for comparison for each gene expression. D to F, Validation of microarray gene expression using cDNA from 18-d shoot tissues of P76, TL, and NC. G to I, Validation of microarray gene expression using cDNA from 18-d root tissues of P76, TL, and NC. The P76 well-watered value was used as a baseline for comparison for gene expression analysis for both root and shoot assays. The representative genes are as follows: *KNAT3*, *E2F*, and *SERK1*. The error bars represent se. The experiment was performed using three biological replicates with two technical replicates per sample. Statistical significance was determined using a one-way ANOVA using Tukey's method. Asterisks represent statistically significant differences: *** $P \leq 0.001$, ** $P \leq 0.01$, * $P \leq 0.05$.

stage, the transcript abundance of *E2F* was 5-fold less in TL compared with P76 in water-limited conditions. The expression patterns of *E2F* in seedling roots was significantly repressed ($P \leq 0.05$) in TL compared with P76 and NC at 4 and 6 d (Fig. 5B).

The expression of *SOMATIC EMBRYOGENESIS RECEPTOR KINASE1* (*SERK1*), a member of the Leu-rich repeat, receptor-like kinase protein family, was notable among the differentially expressed genes among the genotypes during water stress. *SERK1* was up-regulated in roots and shoots of TL under water limitation compared with P76 in the array experiment (Fig. 5). Expression of *SERK1* in TL was 3-fold higher than in NC under water stress. *SERK1* plays a critical role in root differentiation in response to auxin in addition to being involved in somatic embryogenesis and gamete development (Walker, 1994; Schmidt et al., 1997; Somleva et al., 2000; Hecht et al., 2001). Higher expression of *SERK1* in TL relative to P76 and NC was validated by qRT-PCR in roots of pretiltering plants. *SERK1* expression was significantly induced in 4- and 6-d TL seedling roots but not in P76 and NC (Fig. 5C). In summary, our transcriptome analysis identified several differentially regulated genes that could directly or indirectly be associated with the 1-96-1 translocation. We have focused on a small subset of genes

for further characterization based on their role in growth and development in model species.

Gene Network Analysis Predicts the Differential Regulation of Brassinosteroid-Associated Genes in the *A. elongatum* Translocation Line during Water Stress

To further extract *A. elongatum*-associated genetic components involved in the water stress response from our list of differentially expressed microarray genes, we used the wheat-rice orthologous relationships to generate gene regulatory networks using a computational approach. Gene networks were constructed using the recently developed Web-based tool RiceNet to determine putative relationships among genes and predict root trait-associated genes (Lee et al., 2011). Although RiceNet was designed to predict functional relationships among rice genes, it has been shown to accurately predict functional interactions in other monocotyledonous crop species such as maize (Lee et al., 2011). Coexpression network analysis is likely to uncover some genes that are expected to be missed in transcriptome analysis due to low expression levels or lack of representation on the wheat array. RiceNet combines functional genomics, proteomics, and comparative

genome-scale data sets from diverse organisms to predict functional relationships among genes using a Bayesian log likelihood scoring method (Lee et al., 2011). Comparisons of the root transcriptomes of P76 and TL under limited-water conditions resulted in the identification of 384 differentially expressed wheat transcripts. Of these, 224 were assigned to corresponding rice loci. After removal of duplicate loci, a list of 202 genes was queried in RiceNet, resulting in the construction of nine networks from 58 connected genes. Fifty genes were not found in RiceNet, while an additional 94 genes remained unlinked to other genes in the input set. We selected candidate genes with a log likelihood score of 2.0 or greater, with particular focus on genes associated with the regulation of cell division, growth, and root development.

We found a network populated by genes that are either induced by brassinosteroids (BRs) or directly involved in BR signaling, suggesting that BR-associated genes could be differentially regulated in TL (Fig. 6). This network consists of six genes identified during microarray analysis and five predicted genes. The network includes a component of the BR receptor protein complex, BRASSINOSTEROID-INSENSITIVE1 (*BRI1*). Although *BRI1* was not identified from our microarray analysis, its expression was also up-regulated in TL compared with P76 under water-limited conditions in our real-time PCR assays (Fig. 6; Supplemental Table S4). This is consistent with the role of BR in promoting cell wall loosening, root elongation, and lateral root development (Müssig et al., 2003; Bao et al., 2004; Wolf et al., 2012). During the early seedling stage, the expression of network members *CELLULOSE SYNTHASE3* (*CESA3*) and *SERK1* was higher in TL at 4 and 6 d compared with P76 and NC. Differences in lateral root production were also observed at these time points in TL relative to the other two genotypes. The BR gene regulatory network identified from our analysis yielded other members that are listed in Supplemental Table S4. Several of these genes have been reported previously to

have root-specific phenotypes when their expression is altered in mutants and/or transgenic plants in model species.

Single-Feature Polymorphism Analysis to Map Wheat Orthologs to Brachypodium and Rice Genomes for Syntenic Clusters

We next explored if the root-related differentially expressed genes identified from the microarray analysis between TL, P76, and NC map to the 7DL translocation. We used the wheat array transcriptome data set to mine for polymorphisms between TL, P76, and NC using single-feature polymorphism (SFP) analysis (Cui et al., 2005; Walia et al., 2007; Kim et al., 2009). SFP analysis can utilize the transcriptome data as a proxy for DNA to identify polymorphic probes indicative of underlying sequence differences between the *A. elongatum* genes and cultivated wheat alleles. It is important to mention that the presence of a high SFP score probe does not imply differential regulation of the gene represented by the SFP probe. We hypothesized that if most of the SFP-based polymorphisms are due to *A. elongatum* introgression, then many of their orthologs should cluster together on a reference genome such as Brachypodium (*Brachypodium distachyon*) or rice due to the conservation of synteny. Using 19 as an arbitrary cutoff value, our analysis yielded 29 probe sets. The highest score was 42.3 (scores and BLAST hits to the Brachypodium and rice genomes are provided as Supplemental Table S5 and Supplemental File S2). Eight probe sets either did not have a BLAST match to the Brachypodium genome or the e-score for the alignment was higher than -20 . Of the 20 probe sets that met the threshold criterion, eight mapped to a 7DL syntenic region spanning 2.2 Mb in the Brachypodium genome (Fig. 7). When the same threshold criterion was applied to the rice genome, five of the 18 SFP probe sets mapped to a 7DL syntenic region

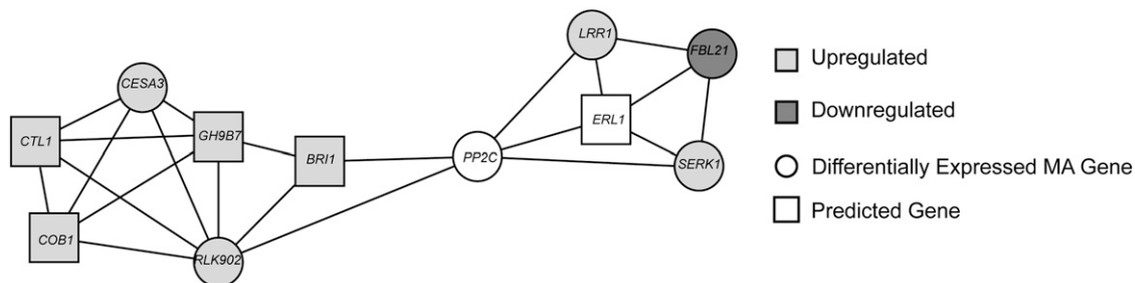


Figure 6. Predicted root-associated gene network based on transcriptome data from the wheat experiment. The network was generated by leveraging the wheat-rice orthologous relationship. Wheat genes differentially expressed between P76 and TL were identified, and their orthologs were used to seed the network using the RiceNet network tool. The displayed network is populated by genes known to be involved in root trait regulation in model species such as Arabidopsis. New genes (square nodes) connected to the input gene set were identified based on RiceNet analysis. The network includes *CESA3*, *CTL1*, *COB1*, *RLK902*, *GH9B7*, *BRI1*, *PROTEIN PHOSPHATASE2C* (*PP2C*), *ERECTA-LIKE1* (*ERL1*), *SERK1*, *F-BOX21* (*FBL21*), and *LEU-RICH REPEAT* (*LRR1*).

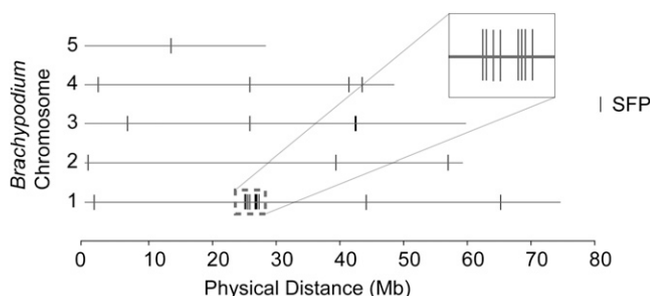


Figure 7. SFP analysis and in silico mapping of wheat orthologs to the Brachypodium genome. High-scoring SFP probe sets from the comparison of P76 samples with TL samples (both well watered and limited water) were BLAST aligned against the Brachypodium genome. Best gene hits to wheat probe sets are mapped to the five chromosomes of Brachypodium. Based on this analysis, we found that eight out of the 20 Brachypodium orthologs (vertical bars) map to the syntenic region on chromosome 1. No other clusters (three or more genes in a 1-Mb window) were observed.

on rice chromosome 6 based on BLAST alignments (Supplemental Fig. S1; Sorrells et al., 2003). Although several high-score SFP calls mapped to nonsyntenic regions, none of them clustered (three or more genes within 1 Mb) in the Brachypodium or rice genome. We selected three predicted SFPs mapping to the syntenic region on the Brachypodium and rice genomes for validation by sequencing the PCR amplicons covering the region around the individual SFP probe in wheat. We were able to confirm the polymorphic probe sequences for these three probe sets (Supplemental Fig. S2).

Mapping of Root-Related Wheat Candidate Genes to Chromosome 7DL

Based on our SFP analysis, we found that two of the previously characterized candidate genes (*KNAT3* and *SERK1*) map to the chromosome 6 region in the SFP cluster in the rice genome (Supplemental Fig. S1). Mapping results suggest that these genes are likely located on the 7DL.7EL translocation. We confirmed the genomic locations of *KNAT3* and *SERK1* in wheat using genomic DNA from a subset of ditelosomic and nullisomic-tetrasomic genetic stocks. The set includes six nullisomic-tetrasomic lines, which lack chromosome 7 for a given genome (A, B, or D) and carry four copies of a corresponding homeologous chromosome of another genome, and two ditelosomic lines, in which one chromosome pair is composed entirely of either short or long arms. Both *SERK1* and *KNAT3* mapped to 7DL (Fig. 8). Two additional differentially expressed genes (*ANNEXIN* and a *Leu-rich repeat extensin-like* gene) also mapped to the wheat 7DL (Fig. 8). Although *E2F* mapped to the distal segment of rice chromosome 6 in the SFP analysis, we were unable to PCR amplify the transcript in wheat for confirmation.

DISCUSSION

A. elongatum 7DL.7EL translocations in wheat are known to improve yield under some rain-fed and water-limiting environments. Our results suggest that the improved yield performance could be due to a more adaptive photosynthesis and root system of the translocation lines. Initial screening at the early seedling stage revealed that TL produced more seminal and lateral roots compared with P76 and NC during water deficit. Lateral root production and primary and seminal root growth were maintained at levels similar to well-watered conditions. Higher root biomass in TL was also observed in older plants at the pretillering stage when grown under moderate water stress. A more resilient photosynthesis is maintained in the pretillering TL plants under water stress, leading to higher carbon fixation compared with control genotypes. Increased availability of carbon under water stress could be partitioned for further root growth and increased water extraction. Collectively, our data suggest that the translocation from *A. elongatum* introduces drought-adaptive traits that involve novel cross talk between shoot and roots that is either absent or regulated differently in P76 and NC. Water stress greatly reduced stomatal conductance and transpiration in P76 and NC during water deficit, while TL

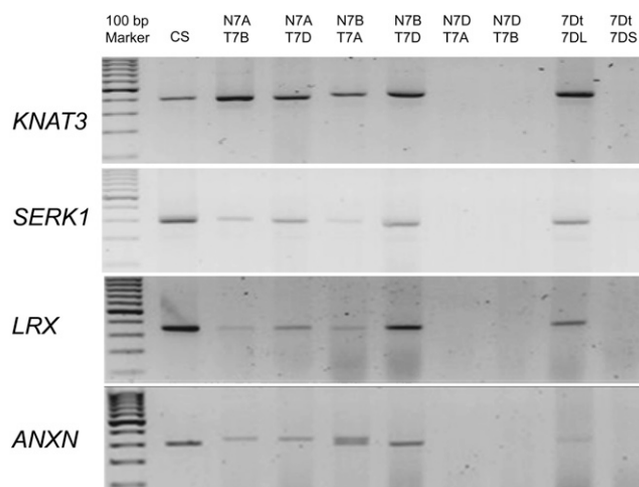


Figure 8. Chromosome mapping of five root trait-associated candidate genes using the wheat genetic stocks. Using cv Chinese Spring and nullisomic-tetrasomic and ditelosomic lines for group 7, PCR-based mapping of candidate genes was performed. The representative genes are as follows: *KNAT3*, *SERK1*, *ANNEXIN* (*ANXN*), and *LEU-RICH REPEAT EXTENSIN-LIKE* (*LRX*). These four genes were selected because they were differentially regulated between TL and control genotypes and their rice orthologs mapped to chromosome 6 with a syntenic relation to the wheat chromosome 7D long arm. CS, cv Chinese Spring; N7AT7B, nullisomic 7A-tetrasomic 7B; N7AT7D, nullisomic 7A-tetrasomic 7D; N7BT7A, nullisomic 7B-tetrasomic 7D; N7BT7D, nullisomic 7B-tetrasomic 7A; N7DT7A, nullisomic 7D-tetrasomic 7B; N7DT7B, nullisomic 7D-tetrasomic 7B; 7Dt7DL, ditelosomic 7DL; 7Dt7DS, ditelosomic 7DS.

maintained stomatal conductance and transpiration rates near control conditions. Chlorophyll fluorescence measurements indicated that the imposed stress was not severe enough to impair PSII photochemistry. This is evidenced by F_v'/F_m' values, which represent the quantum efficiency of PSII photochemistry and remain nearly identical between treatments. Reductions in F_v'/F_m' can be attributed to nonphotochemical quenching or a reduction in the fraction of PSII reaction centers capable of photochemistry (i.e. due to photoinhibition; Papageorgiou and Govindjee, 2004). Therefore, we propose that the reduction in carbon assimilation in P76 and NC under water-limited conditions is due to reduced stomatal aperture and the resulting decline in intercellular CO_2 . This is supported by the observed decline in J_{max} and the increase in photochemical quenching, both of which may be attributed to photorespiration. During drought periods in C3 plants, stomatal closure increases the possibility of Rubisco oxygenation. The reactions of the photorespiratory and glycolate pathways require considerable amounts of ATP and NADPH. Therefore, photorespiration may compete for ATP needed for the phosphorylation of ribulose 5-phosphate and the formation of RuBP. RuBP and ATP contents have been reported to decrease early during drought periods, even as stomatal conductance remains above $100 \text{ mmol water m}^{-2} \text{ s}^{-1}$ (Flexas and Medrano, 2002). Studies by Foyer et al. (2009) reported a significant increase in photosynthetic electron transport caused by photorespiration, which would then drive up photochemical quenching, much like what was observed in P76 and NC during water deprivation.

Due to the adaptive advantage of increased root biomass under water stress, it is important to elucidate the underlying genes and pathways that are differentially regulated in the 7DL.7EL translocation line. The transcriptome study provided candidate genes and a possible link to a BR signaling pathway that could be contributing to root biomass under water stress. Although we highlight root trait-related genes and a BR-related gene network from computational analysis, there are likely to be other important genes and networks involved in root and shoot responses to water stress that were not identified. Based on our array analysis and leveraging a rice computational resource, we have identified differentially regulated genes that could be involved in root differentiation and lateral root production in TL. For instance, a lower transcript level of *KNAT3*, a homeobox transcription factor, was observed during the early seedling and pretiltering stages in TL compared with P76 and NC under water stress. In Arabidopsis, *KNAT3* has been proposed to act as a negative regulator of lateral root development (Truernit et al., 2006; Truernit and Haseloff, 2007). Notably, lower expression of *KNAT3* in TL was correlated with increased lateral root formation during the early seedling stage. Repression of *KNAT3* in wheat in TL, which exhibits increased lateral roots, is consistent with the previously hypothesized role of

KNAT3 in negatively regulating lateral root formation. *KNAT3* maps to the 7DL region in wheat, further supporting its importance as a potential candidate for root architecture.

A second transcript corresponding to the E2F-related protein showed expression patterns similar to *KNAT3* in the early seedling and pretiltering stages. E2F proteins belong to a family of transcription factors that regulate cell cycle progression and cell proliferation (del Pozo et al., 2002; Ramirez-Parra et al., 2004). E2FA, a group I E2F protein, promotes cell division in pericycle tissue, stimulating lateral root development in response to auxin (Berckmans et al., 2011; Magyar et al., 2012). Other E2F proteins assume an opposite role and act as negative regulators of cell cycle progression. Down-regulation of *E2Fc* via RNA interference increased cell proliferation in pericycle cells, resulting in increased lateral root production (del Pozo et al., 2006). Due to the lack of sequence information regarding *E2F*, it remains to be determined whether the identified wheat transcript corresponds to a group I or II E2F protein. Therefore, the contribution of *E2F* to increased lateral root production in TL will require improved access to the wheat genome sequence.

SERK1 was mapped in silico to the distal region of rice chromosome 6, and the location was confirmed to 7DL in wheat. The network analysis revealed a connection between BR signaling and cell growth. *SERK1* forms a protein complex with *BRI1* in Arabidopsis. The Arabidopsis *bri1* mutant exhibits reduced lateral root formation (Clouse et al., 1996; Fukaki and Tasaka, 2009). Endogenous BR application is known to alter root system architecture through synergistic action with auxin (Bao et al., 2004). Arabidopsis *dwarf4* mutants defective in BR biosynthesis fail to form lateral roots when exposed to exogenous indole-3-acetic acid (Yoshimitsu et al., 2011). A number of studies have associated *SERK1* expression with lateral root initiation and development (Hecht et al., 2001; Kwaaitaal et al., 2005; Kwaaitaal and de Vries, 2007; Nolan et al., 2009). This is consistent with the higher expression levels of *SERK1* and increased lateral root production observed in TL during the early seedling stage. Longer primary and seminal root lengths and an increase in the number of seminal roots during the early seedling stage could possibly be due to the influence of the *A. elongatum* translocation on embryogenesis. In this context, it is noteworthy that *SERK1* expression was reported during ovule development in all the cells of the embryo sac in Arabidopsis (Albrecht et al., 2005). Based on the expression patterns of *SERK1* and *E2F* in TL during water stress, their role in lateral root formation, and known synergism between the auxin and BR pathways, it is possible that *A. elongatum* translocation could be altering hormone signaling in the roots.

The coexpression network with *SERK1* identified in our analysis includes several cell wall-related genes, such as *CESA3*, *COBRA1* (*COB1*), *GLYCOSYL HYDROLASE9B7* (*GH9B7*), and *CHITINASE-LIKE1* (*CTL1*), that

are linked to *BRI1* (Fig. 6). BR is known to compensate for the loss of cell wall integrity by inducing the expression of genes involved in cell wall biosynthesis (Wolf et al., 2012). Network analysis also revealed a putative relationship between *CESA3* and *RECEPTOR-LIKE KINASE902* (*RLK902*). Both genes shared similar expression patterns and were up-regulated in TL (Fig. 6; Supplemental Table S4). Repression of *RLK902* has been associated with reduced root growth and root meristem size (ten Hove et al., 2011). Both phenotypes were also observed in other cellulose synthase-defective mutants (Ellis et al., 2002; ten Hove et al., 2011). The expression of several genes involved with cell growth and primary cell wall formation also showed higher expression levels in TL when compared with P76 under limited-water conditions. A transcript corresponding to the *CESA3* gene in Arabidopsis was up-regulated in TL (Fig. 6; Supplemental Table S4). BR is known to induce *CESA* genes through the BR transcription factor *BRI1-EMS-SUPPRESSOR1* (Xie et al., 2011). *CTL1* was up-regulated in TL compared with P76 during water deficit (Fig. 6; Supplemental Table S4). Loss-of-function *ctl1* mutants have reduced root growth and cellulose deposition and increased sensitivity to drought and saline conditions (Kwon et al., 2007). Differential expression of some of the cell wall- and BR-related network members in TL relative to P76 and NC suggests that BR signaling could be a component of root biomass plasticity observed in TL during water deficits.

For the transcriptome analysis and gene discovery, our experiments relied on the representation on the wheat microarray and the orthologous relationship between rice and wheat for input into RiceNet. It is possible that our analysis could have missed some rare genes introduced from *A. elongatum* that confer the observed shoot- and root-related traits under water stress. Two recent reports in wheat (*TaSC*) and rice (*Pstol1*) indicate the value of such genes that are missing from commonly used germplasm and reference genomes (Gamuyao et al., 2012; Huang et al., 2012). As the wheat draft genome becomes more accessible, future experiments using the RNAseq approach can enable the discovery of novel stress tolerance genes.

A combination of whole-plant physiological assays with functional genomics and network biology approaches enabled us to better understand the adaptive advantage introduced by an alien chromatin introgression into wheat. Our results suggest that increased lateral root production displayed by introgression line TL in response to water depletion could be one of the contributing factors to improved plant performance. Future studies will determine whether phenotypic differences observed during these early stages translate to differences in grain production during moderate drought events. Future work will involve functional characterization of the candidate genes identified from the transcriptome analysis. As phenotyping technology becomes automated and allows

larger populations to be screened in a noninvasive manner, we will test if allelic variants for candidate genes identified in this study correlate with phenotypic variation for drought adaptability. Should these candidate genes provide higher yields or greater yield stability, they can be valuable for wheat breeding programs in many environments.

MATERIALS AND METHODS

Plant Material

All plant materials used in this study were obtained from A.J. Lukaszewski (University of California, Riverside). These included a third-generation derivative of the original wheat (*Triticum aestivum*)-*Agropyron elongatum* translocation chromosome produced by E.R. Sears (Sears, 1978) and controls. The starting chromosome transfer 1 (Sears, 1978) was generated by *ph1b*-induced homeologous recombination and consists of normal 7DS, a short proximal segment of 7DL, and a long distal segment of *A. elongatum* chromosome 7EL. A line of wheat 'Chinese Spring' homozygous for transfer 1 was crossed and backcrossed three times to P76. The last two backcrosses were to a *ph1b* line of Pavon, resulting in several plants heterozygous for transfer 1 and homozygous for the *ph1b* mutation. These plants were grown and self-pollinated, and their progeny were screened to identify secondary wheat-*A. elongatum* recombinant chromosomes. Among many such chromosomes isolated was a chromosome labeled 1-96 (Zhang et al., 2005). Chromosome 1-96 is essentially a normal chromosome 7D of wheat with the terminal segment of the long arm, perhaps 20% in length, originating from 7E of *A. elongatum*. The *A. elongatum* segment carries *Lr19* and *Y*. This chromosome was again combined with the *ph1b* mutation in P76, and a tertiary recombinant chromosome was recovered, labeled 1-96-1. This chromosome differs from 1-96 by the absence of the terminal *A. elongatum* segment that normally carries the *Y* locus. The translocated chromosome 1-96-1 has a normal 7DS arm, a long (approximately 75%) proximal segment of 7DL, a homeologous segment from 7EL of *A. elongatum*, and a short terminal segment of 7DL (Fig. 1). Since the chromosome was produced by crossing over, it is assumed to be fully compensating for normal 7D. No indications of reduced compensation have been observed.

After recovery, plants with chromosome 1-96-1 were backcrossed an additional three times to P76, and two sets of sister lines were isolated among progeny from self-pollinated BC₃ heterozygotes: homozygotes for the 1-96-1 translocations and disomics of normal chromosomes 7D. These two are sister lines that differ by the presence/absence of the *A. elongatum* segment; disomics of 7D serve as additional controls for the translocation lines. Overall, the recovered lines have eight backcrosses to P76; however, it needs to be taken into account that the *ph1b* lines carry substantial and extensive genome rearrangements, and new ones are generated in each generation. There were only three backcrosses to normal P76 after the last round of the *ph1b*-induced recombination. For this reason, additional controls in the form of sister lines are highly desirable. For simplicity, the line of P76 homozygous for the 1-96-1 translocation is referred to as TL here, P76 is referred to as P76, while the sister lines of TL without the translocation are called NC. Nullisomic-tetrasomic lines for the homeologous group 7 as well as ditelosomic lines of chromosome 7D in cv Chinese Spring were used to determine and verify the chromosome locations of various candidate genes.

Cigar Roll Root Assays

For the early seedling stage root screening, we used the cigar roll method (Zhu et al., 2006). Similar-sized seeds were germinated in a petri dish with a moist paper towel for 3 d in the dark at 22°C to 25°C. Three seedlings per genotype were placed between two sheets of germination paper and rolled vertically. Three batches of 10 cigar rolls, totaling 30 plants per genotype, were soaked in tap water and placed vertically in a 1-L beaker. For well-watered conditions, the beakers were filled with 100 mL of tap water, while 50 mL of tap water was supplied for water-limited conditions. Seedlings were grown in a controlled environment using a Percival Intellus Control System incubator (model I36VLC8) set at 25°C/27°C, 11.5-h/12.5-h day/night, 50%/70% relative humidity, and illumination of 80 $\mu\text{mol m}^{-2} \text{s}^{-1}$ for the duration of the experiments (Percival Scientific). Water was added to each beaker each day to

maintain the volume of 50 mL. All measurements were carried until 6 d after germination. Measurements after day 6 can potentially be affected by limiting seed resources that drive seedling growth. To ensure that the observed differences in measured traits were not due to nutrient limitations, the experiment was repeated twice with a one-tenth-strength Hoagland solution (Supplemental Table S2). Root length was measured from the tip of the root to the root-shoot junction, while shoot length was measured from the tip of the longest leaf to the root-shoot junction. To calculate lateral root density, the number of lateral roots on a branch from a main axis (primary or seminal root) was divided by the length of the root. For gene expression experiments, seedlings were harvested at 2, 4, and 6 d after germination. Root tissue samples were collected and snap frozen.

Greenhouse Experiments

Seeds were germinated in glass petri dishes in the dark at 25°C. Once the coleoptiles were approximately 1 cm, uniform seedlings were selected and transplanted to polyvinyl chloride pipes (1 m × 15.5 cm [height × diameter]; NDS). Each pipe was lined with clear polyplastic bags (153 cm × 18.5 cm [length × width]; Uline) and filled with approximately 9 kg of fine silica sand (Lane Mountain). A gap was maintained between the plastic bags and the inner wall of the pipes. Small holes were made at the side and bottom of the plastic bags as well as the pipes to provide adequate aeration and drainage. For the first week after transplant, each tube was watered daily with 100 mL. After the first week, the well-watered, control plants were supplied with 50 mL of tap water and 50 mL of one-half-strength Hoagland solution twice weekly (irrigated four times per week). For the limited-water treatment, each tube received 50 mL of one-half-strength Hoagland solution twice weekly. Each treatment consisted of seven replicates per genotype with five plants per tube/replicate (35 plants per treatment). For limited-water conditions, the watering regime of 100 mL per week was based on preliminary experiments with 200, 100, and 50 mL of water twice per week for water stress. At the 200-mL watering level, we did not observe a difference in root and shoot biomass among the genotypes, and the sand column was fairly wet at 3 weeks post germination. At the 50-mL watering level, almost all plants from the three genotypes had senesced and were not suitable for photosynthetic measurements. We observed clear differences in the 100-mL treatment in root and shoot biomass. Furthermore, in the 100-mL water stress treatment, the sand column was fairly dry and all three genotypes showed a decrease in stomatal conductance indicative of successful imposition of a moderate water stress.

The pretillering stage experiment was done from July to October 2011. The greenhouse temperature was set between 26.7°C and 32.2°C during the day and 26.1°C to 29.4°C during the night, humidity ranged from 33% to 73%, and the light period was 7:00 AM to 6:00 PM by Metal Halide Lamp 1,000-W bulbs (Philips Lighting). Using the LI-250A Light Meter, the average light reading was $215.8 \pm 44.5 \mu\text{mol m}^{-2} \text{s}^{-1}$ (LI-COR Biosciences).

Wheat roots were harvested 18 d after germination. The plastic sleeve was carefully removed from each tube, and the plastic lining was cut gently to avoid root damage. Roots were gently removed from the sand column. Excess sand was removed using a paintbrush with soft bristles, and root and shoot lengths were recorded immediately. Shoot length was measured from the tip of the leaf to the root-shoot junction, while root length was measured from the root-shoot junction to the longest root. Root and shoot dry weights were determined after drying for 5 d at 60°C. Statistical analyses were done using the one-way ANOVA function using Tukey's method in the R-Commander package for R (Fox, 2005).

Chlorophyll Fluorescence, Photosynthesis, and Gas-Exchange Measurements

Chlorophyll fluorescence measurements were recorded at day 18 after transplant for each treatment on the leaf below the newest fully expanded leaf using a portable photosynthesis system (LI-COR 6400xt) equipped with a leaf chamber fluorometer (LI-COR 6400-40; LI-COR Biosciences). The area of each leaf in the leaf chamber was determined, and values were entered manually into the system. Measurements were recorded between 10:00 AM and 2:00 PM. Reference CO₂ levels were maintained at $400 \mu\text{mol s}^{-1}$, and chamber flow was set at $500 \mu\text{mol}$. Actinic light intensity was set at $1,000 \mu\text{mol}$ with 10% blue light-emitting diodes. Maximal fluorescence values of light-adapted leaves were obtained by saturating the leaf using a multiphase flash (Loriaux et al., 2006). The first phase consisted of an 8,000- μmol pulse for 300 ms followed by

a 100% ramp for 400 ms and a second saturating flash of 8,000 μmol for 300 ms. The maximum quantum efficiency of PSII ($\frac{F_v'}{F_m'}$) was calculated using the equation:

$$\frac{F_v'}{F_m'} = \frac{F_m' - F_o'}{F_m'}$$

where F_v' refers to variable fluorescence, F_m' is maximal fluorescence, F_o' is minimum fluorescence, and F_s is the steady-state fluorescence. Photochemical quenching was calculated using the equation:

$$\frac{F_m' - F_s'}{F_m' - F_o'}$$

Nonphotochemical quenching was determined using the equation:

$$\frac{F_m - F_m'}{F_m - F_o'}$$

Maximum fluorescence in the dark-adapted state (F_m) was determined by temporarily shading leaf tissue with a black cloth for 30 min and applying a far-red pulse for 3 s to fully oxidize PSII reaction centers. After this, a single saturating flash was applied (8,000- μmol pulse for 300 ms) to obtain F_m .

Carbon Assimilation/Intercellular CO₂ Curve with Chlorophyll Fluorescence

Gas-exchange and fluorescence measurements were recorded for each replicate with the LI-COR 6400xt using polyvinyl chloride-cultured wheat plants on day 18. Instrument settings were identical to those described above. Six CO₂ response curves per genotype were generated for each treatment. Gas-exchange and fluorescence measurements were recorded at 13 CO₂ flow rates (400, 300, 200, 100, 50, 100, 150, 200, 300, 400, 500, 600, and 800 $\mu\text{mol s}^{-1}$). CO₂ assimilation was plotted against intercellular CO₂ (C_i), and the data points were fitted to the equation of Farquhar et al. (1980) using SAS and the methods described by Dubois et al. (2007).

RNA Isolation and RNA Cleanup

Eighteen days after transplanting, leaf and root tissues were collected and snap frozen in liquid nitrogen for RNA extraction. RNA was extracted using the TRIzol method. RNA concentration was quantified (1:1,000) with the DU 730 Life Science UV/Vis Spectrophotometer (Beckman Coulter). RNA cleanup was performed according to the RNeasy MinElute Cleanup kit protocol (Qiagen). The RNA was eluted with 15 μL of RNase-free water along with on-column DNaseI treatment.

First-Strand cDNA Synthesis and Quantitative Reverse Transcription-PCR

We used SuperScript VILO to synthesize complementary DNA (cDNA) for semiquantitative PCR and real-time qRT-PCR (Invitrogen). First-strand cDNA was generated for two-step reverse transcription PCR by following the protocol from the Superscript VILO cDNA synthesis kit. An amount of 2 μg of RNA was used in the 20- μL reaction mixture. For qRT-PCR, 4 μL of the diluted cDNA (1:20) was used in a 15- μL reaction mixture. In the qRT-PCR volume, 7.5 μL of iQ SYBR Green Supermix was used (Bio-Rad Laboratories). qRT-PCR was run using LightCycler 480 II with the following parameter settings (Roche Applied Science): 95°C preincubation for 10 min; amplification was done for 45 cycles at 95°C for 10 s and 60°C for 10 s; the melting curve was set for 95°C, 65°C, and 97°C; cooling was set at 40°C for 30 s. Each qRT-PCR run was performed with three independent tissue samples, each sample having two technical replicates. Two genes that showed stable expression values in a large set of microarray experiments across treatments and tissue samples were used as internal controls. A description of the genes and primer sequences is provided in Supplemental Table S4. Crossing point value, which was the point at which the fluorescence crosses the threshold, and melting curve analyses were obtained using the Roche LightCycler 480 software version 1.5. The melting curve data were collected for all samples and genes to ensure a single peak, indicating amplification of a specific region by a pair of primers.

Microarray Data Analysis

Wheat Array GeneChip data were generated using a standard Affymetrix protocol. Two independent biological replicates were used for each array experiment. CEL files were analyzed using the limma package in the R software program. Briefly, the Bioconductor affy and limma packages were used for reading CEL files and normalizing microarray data (Smyth, 2005). The raw intensity values for the GeneChip arrays were background corrected, \log_2 transformed, and quantile normalized with robust multiarray average within the R package affy (Izarzary et al., 2003). The statistical computation and assessment of differential expression were done with the empirical Bayes function, which moderates the SE values of estimated log fold changes toward a pooled SD value. *P* values were adjusted to the Benjamini and Hochberg method (BH) to control the false discovery rate. The top table function was used to list the genes according to the specified *P* value of 0.05, \log_2 fold change of 1, and *P* value adjust = BH. The probe set annotated was generated from HarvEST: WheatChip (Close et al., 2007). The raw array data are publicly available through the National Center for Biotechnology Information Gene Expression Omnibus (GSE42214). Complete lists of differentially expressed transcripts are given in Supplemental Table S3 and Supplemental File S1.

Construction of the Coexpression Network

The coexpression network was constructed with RiceNet using rice (*Oryza sativa*) orthologous information from HarvEST: Wheat for differentially expressed transcripts between P76 and TL under limited-water conditions (Lee et al., 2011). A BLASTX search against the rice genome (Michigan State University version 6; January 2009) was performed to identify rice orthologs for each differentially expressed transcript. Wheat sequences without a hit or with *e*-values exceeding $1e^{-8}$ were excluded from analysis. Genes corresponding to ribosomal proteins were removed to reduce the occurrence of false relationships between otherwise nonconnected genes. A Bayesian log likelihood score threshold of 2 was used to reduce the number and improve the reliability of predicted connections. Predicted rice loci without a BLASTX hit to wheat Affymetrix transcripts or with *e*-values greater than $1e^{-8}$ were omitted from further analysis. Primers for the genes identified from the network analysis and validated with qRT-PCR are listed in Supplemental Table S6, and their relative expression is summarized in Supplemental Table S4. P76 under limited-water conditions was used as a reference for the validation of expression using qRT-PCR.

SFP Identification and in Silico Mapping of Candidate Genes

Microarray data were preprocessed as described above. To eliminate probe sets with absent transcripts, we adopted the procedure suggested by Schuster et al. (2007), making "present" and "absent" calls for each probe set. A probe set was retained and used for SFP detection if it had present calls in all biological replicates of the two genotypes under comparison. We used the robustified projection pursuit method for SFP detection (Cui et al., 2005) through pairwise comparison among parent genotype P76, TL, and NC for all samples. For each comparison, we used an arbitrary cutoff value of 19 for calling SFP-containing probe sets, within which a probe will be identified as an SFP probe if it accounts for more than 40% of the overall outlying score of its residing probe set. A BLASTX search against the Brachypodium (*Brachypodium distachyon*) genome (Phytozome Bradi 1; May 2009) was performed for each probe set exceeding the SFP threshold of 19. Also, sequences without a BLASTX hit or with an *e*-value less significant than $1e^{-20}$ were excluded from in silico mapping to the Brachypodium genome. A complete list of probe sets identified through SFP analysis is provided as Supplemental File S2.

Primers were designed so that the region flanking each probe was amplified. Sequences were amplified from cDNA using the primers listed in Supplemental Table S6 using PCR. A 25- μ L reaction was prepared using GoTaq Green Master Mix (Promega) consisting of 1 μ L each of forward and reverse primers at 10 μ M and 2 μ L of cDNA. Sequences were amplified using the following program: denaturation at 94°C for 3 min, followed by 30 cycles of amplification (30 s at 94°C, 30 s at 58°C, and 30 s at 72°C), and a final temperature of 72°C for 7 min. Prior to cloning, each reaction was purified using the GeneJET PCR Purification Kit (Thermo Scientific). Amplicons were cloned into pGEM T Easy Vector (Promega), and JM109 competent cells were transformed and cultured according to the manufacturer's protocol. Cultures were plated on Luria-Bertani plates supplemented with 100 μ g of ampicillin and

5-bromo-4-chloro-indolyl- β -D-galactopyranoside-isopropyl- β -D-1-thiogalactopyranoside. Plasmids were retrieved using the GeneJET Plasmid Miniprep Kit (Thermo Scientific) and sequenced.

PCR-Based Mapping of Genes to Wheat Chromosome 7D

A subset of microarray genes that were mapped through SFP analysis to 7DL using rice-wheat synteny maps were confirmed using a series of nullisomic-tetrasomic and ditelosomic wheat 'Chinese Spring' lines (Devos et al., 1999; Sorrells et al., 2003). Primers specific to 7DL were designed using probe set sequences obtained from HarvEST: WheatChip version 1.59, Grain Genes (<http://wheat.pw.usda.gov/>) and the National Center for Biotechnology Information BLAST (Altschul et al., 1997; Close et al., 2007). The selected 7D sequences were used to design primers using Primer3 software (Rozen and Skaletsky, 2000). PCR was performed to map the genes using genomic DNA from cv Chinese Spring, nullisomic-tetrasomic and ditelosomic 7DL, and 7DS wheat 'Chinese Spring' plants. PCR was performed using the following settings: 95°C preincubation for 3 min, amplification for 40 cycles at 95°C for 30 s, 58°C for 30 s, and 72°C for 1 min, and extension at 72°C for 3 min. The PCR products were run on a 2% agarose gel (molecular biology grade; Benchmark Scientific) and imaged using the BioDoc-IT Imaging system UV Transilluminator (UVP).

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number GSE42214.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. In silico mapping of wheat genes on the rice genome using SFP.

Supplemental Figure S2. Validation of polymorphic probes identified via SFP/robustified projection pursuit analysis.

Supplemental Table S1. Phenotypic data for early seedling screens using the cigar roll method with tap water.

Supplemental Table S2. Phenotypic data for early seedling screens using the cigar roll method with a one-tenth-strength Hoagland solution.

Supplemental Table S3. Subset of differentially expressed genes from a comparison of P76 with TL during water deficit.

Supplemental Table S4. Relative expression of root-associated network genes during water deficit during the pretiltling stage.

Supplemental Table S5. Subset of polymorphic probe sets identified through SFP analysis.

Supplemental Table S6. List of primer sequences used for PCR and qRT-PCR.

Supplemental File S1. Complete list of differentially expressed genes identified by transcriptional profiling of wheat roots.

Supplemental File S2. Complete list of probe sets identified through SFP analysis.

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