Down-Regulation of the CSLF6 Gene Results in Decreased (1,3;1,4)-β-D-Glucan in Endosperm of Wheat1[C][W]

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(1,3;1,4)-β-D-Glucan (β-glucan) accounts for 20% of the total cell walls in the starchy endosperm of wheat (Triticum aestivum) and is an important source of dietary fiber for human nutrition with potential health benefits. Bioinformatic and array analyses of gene expression profiles in developing caryopses identified the CELLULOSE SYNTHASE-LIKE F6 (CSLF6) gene as encoding a putative β-glucan synthase. RNA interference constructs were therefore designed to down-regulate CSLF6 gene expression and expressed in transgenic wheat under the control of a starchy endosperm-specific HMW subunit gene promoter. Analysis of wholemeal flours using an enzyme-based kit and by high-performance anion-exchange chromatography after digestion with lichenase showed decreases in total β-glucan of between 30% and 52% and between 36% and 53%, respectively, in five transgenic lines compared to three control lines. The content of water-extractable β-glucan was also reduced by about 50% in the transgenic lines, and the M₆₆ distribution of the fraction was decreased from an average of 79 to 85 × 10⁴ g/mol in the controls and 36 to 57 × 10⁴ g/mol in the transgenics. Immunolocalization of β-glucan in semithin sections of mature and developing grains confirmed that the impact of the transgene was confined to the starchy endosperm with little or no effect on the aleurone or outer layers of the grain. The results confirm that the CSLF6 gene of wheat encodes a β-glucan synthase and indicate that transgenic manipulation can be used to enhance the health benefits of wheat products.

Wheat endosperm cell walls comprise two major components, arabino-(1,4)-β-D-xylan (arabinoxylan [AX]) and (1,3;1,4)-β-D-glucan (β-glucan), which account for about 70% and 20% of the total, respectively (Bacic and Stone, 1980). In addition about 4% cellulose [(1,4)-β-D-glucan] and 7% (1,4)-β-D-glucomannans are present (Bacic and Stone, 1980). This contrasts with starchy endosperm tissues of oats (Avena sativa) and barley (Hordeum vulgare), in which the proportions of AX and β-glucan are reversed (Henry, 1987; Stone, 1996). This is of particular interest as soluble β-glucans from barley and oats have benefits in reducing coronary heart disease that have been accepted by the U.S. Food and Drug Administration for health claims on food products (Anonymous, 2008). It is not known whether these benefits are shared by β-glucan from wheat, which differs from barley and oat β-glucans in its distribution of 1,3 and 1,4 linkages (Lazaridou and Biliaderis, 2007). Differences in M₆₆ of β-glucans have also been reported, although it is difficult to conclude whether these result from true genetic differences, the effects of growth conditions, or the methods used for extraction and characterization (Lazaridou and Biliaderis, 2007). However, differences in linkage distribution are likely to affect β-glucan solubility and

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viscosity (Cui et al., 2000; Lazaridou and Biliaderis, 2007), which are considered to be key parameters determining health benefits (Wood, 2007).

Despite the biological importance of plant cell walls, we know relatively little about their biosynthesis. However, it is becoming clear that members of the cellulose synthase super family are responsible for the synthesis of the backbone of hexose polysaccharides, with cellulose itself being synthesized by enzymes encoded by the CELLULOSE SYNTHASE A (CESA) family (Richmond and Somerville, 2000) and other polysaccharides by enzymes encoded by genes in CELLULOSE SYNTHASE-LIKE (CSL) families designated A to H and J. The latter include genes encoding (1,4)-β-D-mannans (CSLA; Dhugga et al., 2004; Liepman et al., 2007) and the (1,4)-β-D-glucan backbone of xyloglucan (CSLC; Cocuron et al., 2007).

Burton et al. (2006) provided convincing evidence that the rice (Oryza sativa) CSLF genes encode β-glucan synthases, by expression in transgenic Arabidopsis (Arabidopsis thaliana) and detection of β-glucan (which is not normally synthesized in Arabidopsis) with a highly specific antibody. Furthermore, the corresponding CSLF genes of barley were mapped to a locus on chromosome 2H corresponding to a major quantitative trait locus (QTL) for grain β-glucan content, and two other CSLF genes to loci on chromosomes 1H and 7H that were also close to β-glucan QTLs. The seventh barley CSLF gene mapped to a locus on chromosome 5H and showed no linkage to β-glucan QTLs (Burton et al., 2008). However, more recently Doblin et al. (2009) have shown that the barley CSLH gene also leads to the synthesis of β-glucan when expressed in transgenic Arabidopsis. The authors concluded that both the CSLF and CSLH gene families contribute to β-glucan synthesis in this species, and suggested that they act independently, based on the fact that there was no significant correlation between their transcripts.

We have identified the CSLF6 gene of wheat as a putative β-glucan synthase and confirmed this function by RNA interference (RNAi) suppression in transgenic wheat. This demonstrates that transgenic manipulation can be used to modify the amounts and properties of β-glucan in wheat, to enhance the health benefits and improve the properties for processing and livestock feed.

RESULTS

Bioinformatic Identification of Candidate Genes

We hypothesized that candidate genes for β-glucan synthesis should be expressed in the developing caryopses during the early (cell division and expansion) and mid (grain filling) stages of development, when cell wall synthesis is actively occurring. Furthermore, we proposed that such genes were likely to be members of the CSL gene family. Figure 1 therefore shows the abundances of transcripts for CSL genes represented on the wheat Affymetrix GeneChip, hybridized to cDNA isolated from whole caryopses of wheat throughout development (Wan et al., 2008; Fig. 1A) and from fractions enriched in starchy endosperm tissue taken at 14 to 21 DPA (Fig. 1B). Transcripts are labeled according to the most similar rice sequences following the naming convention for CSL genes (Hazen et al., 2002). Data are only shown for the six most abundant CSL transcripts in the endosperm-enriched samples; the complete list of CSL genes represented on the chip and their average expression levels are given in Supplemental Table S1. The most highly expressed transcripts in the endosperm-enriched fraction (Fig. 1B) are members of the CSLD family (D2) and the CSLA family (A7-1, A7-2, A12), the latter being implicated in mannan synthesis (Dhugga et al., 2004; Liepman et al., 2007). The molecular function of CSLD genes, which are the most similar to CESA, are unknown, but Arabidopsis knockout mutants for CSLD genes have defective root tip or pollen tube development (Bernal et al., 2008). The next most abundant transcripts in the endosperm-enriched fraction (Fig. 1B) are CSLF6 and CSLC1, implicated in β-glucan and xyloglucan synthesis, respectively (Burton et al., 2006; Cocuron et al., 2007), with both being particularly abundant early in grain development (Fig. 1A). These expression patterns are consistent with the report that β-glucan synthesis is initiated earlier and proceeds more actively during early grain development than the synthesis of AX in wheat (Philippe et al., 2006).

The relative abundances of CSL transcripts from the array data are consistent with counts of wheat ESTs from grain libraries; the five transcripts identified as most abundant in Figure 1 being in the top six ranked by EST counts (Supplemental Table S2). These ESTs indicate the presence of at least seven CSLF genes in
wheat, corresponding to the rice CSLF1, CSLF2, CSLF3, CSLF4, CSLF6, CSLF8, and CSLF9 genes. Of the CSLF transcripts in barley, the HvCSLF6 gene, an ortholog of TaCSLF6, is also by far the most abundant during the cell expansion phase in developing barley endosperm (Burton et al., 2008). Another barley transcript, HvCSLF9, was shown to be reasonably abundant early in endosperm development, but the wheat ortholog has only two associated wheat ESTs (Supplemental Table S2), compared with 30 barley ESTs, and is not represented on the wheat Affymetrix array.

Since the CSLA genes are implicated in the synthesis of mannans (Dhugga et al., 2004; Liepman et al., 2007) that account for only 7% of wheat endosperm cell walls (Stone and Morell, 2009), CSLD2 and CSLF6 transcripts were selected as targets for RNAi suppression to generate changes in endosperm cell wall composition. During the progress of this work, the rice CSLF genes were also implicated in the synthesis of β-glucan by expression in transgenic Arabidopsis (Burton et al., 2006), making the CSLF6 gene the stronger candidate. Similarly, a recent study of the β-D-glucan less (bgl) mutant of barley showed cosegregation of the mutation with polymorphism in the CSLF6 gene, suggesting that CSLF6 encodes a β-glucan synthase (Tonooka et al., 2009).

Construction of RNAi Cassettes

Supplemental Figure S1 shows the protein sequence corresponding to the TaCSLF6 gene cloned from cv Cadenza endosperm cDNA, compared with those of the rice OsCSLF2 gene that was shown by Burton et al. (2006) to confer the capacity to synthesize β-glucan by expression in transgenic Arabidopsis and the orthologous barley HvCSLF6 gene that is highly expressed in the endosperm (Burton et al., 2008). Supplemental Figure S2 compares the nucleotide sequence of the 3′ end of this gene with single nucleotide polymorphisms present in two other forms (determined by sequencing of 15 cDNA clones covering the whole region shown and a further 21 covering the region used for the RNAi construct and that are almost certainly homoeologs) present in hexaploid cv Cadenza cDNA. The fragments used for the RNAi constructs against CSLF6 (Supplemental Fig. S2) and against CSLD2/CSLD4 (Supplemental Fig. S3) are also shown; an RNAi construct is expected to repress transcripts that contain regions of 20 bp identity or more, so homoeologs and other closely related isoforms will be affected. Partial CSLF6 and CSLD4 cDNAs cloned from cv Cadenza were used as templates to PCR amplify fragments for RNAi. Two RNAi plasmids were constructed (Supplemental Fig. S4), one designed to silence only CSLF6 (pHmw-Adh-Nosf6ri/ri) and the other designed to simultaneously silence both CSLD2 and CSLD4 (pHmw-Adh-Nosd4ri/ri) by using regions of CSLD4 cDNA possessing high similarity to CSLD2. The RNAi cassettes were constructed with inverted repeats flanking the maize (Zea mays) Adh2 intron and under the control of the endosperm-specific high M₆ glutenin subunit 1Dx5 promoter (Lamacchia et al., 2001).

Production and Initial Screening of RNAi Lines

Two RNAi cassettes were used, both together and separately, to generate transgenic lines of wheat (cv Cadenza). A total of 29 independent transgenic T0 lines were made, 13 containing the CSLF6 RNAi cassette, eight containing the CSLD2/4 RNAi cassette, and a further eight containing both constructs. Presence of the selectable marker gene and RNAi cassette in genomic DNA from each T0 plant was confirmed by PCR. Additional evidence for the presence of transgene cassettes in these plants was provided using TaqMan real-time PCR analysis performed on selected T1 plants by iDNA Genetics.

T1 seed from 16 lines (10 containing the CSLF6 RNAi cassette only, four containing the CSLD2/4 RNAi cassette only, and two containing both constructs) were germinated and the T1 progeny were screened by PCR of genomic DNA to determine the segregation ratios of the transgenes. The χ² goodness of fit test indicated that the transgenes were segregating with a 3:1 ratio in 12 of these lines, including line 15T that possessed both CSLF6 and CSLD2/4 cassettes, indicative of a single transgene locus containing both cassettes (Supplemental Table S5). In two of the lines, the χ² goodness of fit test predicted a 15:1 ratio, indicating that a CSLF6 cassette had inserted in two unlinked loci (Supplemental Table S5). Line 22T that also contained both CSLF6 and CSLD2/4 cassettes possessed a classic 9:3:3:1 ratio of two unlinked loci, with one containing at least one copy of CSLF6 and the other, at least one copy of CSLD2/4. The remaining line (10T) segregated in a non-Mendelian ratio.

Enzyme fingerprinting of T1 seed of transgenic wheat plants carrying only RNAi designed against the CSLD genes showed no detectable differences in grain cell wall composition from the controls. We therefore concluded that the CSLD genes do not control the synthesis of the major groups of cell wall polymers in wheat endosperm (β-glucan and AX). In contrast, enzyme fingerprinting of T1 seeds of 10 heterozygous transgenic lines carrying RNAi designed against CSLF6 showed a mean decrease in β-glucan content of over 20% (data not shown). It was therefore decided to focus attention on the lines expressing the CSLF6 RNAi constructs.

Five independent lines that displayed both 3:1 segregation of the transgene and significantly reduced β-glucan content in T1 seeds were subjected to transgene copy number analysis and zygosity estimation via quantitative PCR. From each of these lines, homozygous T1 sublines were propagated to collect T2 seed that were then sown to provide T3 seed for further analyses. Four of these lines possessed the CSLF6 RNAi cassette (lines 4T, 7T, 9T, and 21T) and one possessed both CSLF6 and CSLD2/4 cassettes (line...
Transgene copy number ranged from one to 14 copies per line (Supplemental Table S5). These five lines were used for β-glucan analysis. The 100 grain weights of the five transgenic lines ranged from 2.96 g to 4.21 g (mean 3.75 g), which did not differ significantly (P = 0.14) from the 100 grain weights of three control lines (range 2.91 g to 3.45 g, mean 3.22 g). This confirms that the expression of the transgene did not have any significant effect on grain filling.

### Transgene and Endogenous CSLF6 Expression

The expression of the CSLF6 RNAi transgene and its effect on the abundance of the endogenous CSLF6 transcript were determined in line 15T and controls by quantitative reverse transcription (qRT)-PCR (Fig. 2). In RNA samples isolated from dissected endosperm at 14 DPA, primers designed against the transgene as well as the native gene showed abundance 11-fold greater in 15T compared to null controls (P < 0.001), whereas primers designed exclusively to the endogenous CSLF6 transcript showed abundance decreased by 4-fold on average (P = 0.025).

### β-Glucan Contents of Mature Grain

The mean content of total β-glucan in wholemeal fractions of the five transgenic lines was reduced by 42.2% compared to that in the three control lines, with the greatest reduction of 52.2% occurring in line 21T (Table I).

This decrease in total β-glucan was confirmed by enzyme fingerprinting, digesting with lichenase and endoxylanase to determine AX and β-glucan in a single analysis (Ordaz-Ortiz et al., 2005; Saulnier et al., 2009). Typical high-performance anion-exchange chromatography (HPAEC) separations of the digestion products from grain of control (null) and transgenic lines are shown in Figure 3.

Oligosaccharides released from AX by endoxylanase were Xyl (X), xylobiose (XX), and oligosaccharides of DP4 to 9 (XA²X², XA³XX, XA²+³XX, XA³A²XX, XA²A²+³XX, XA³XX, and XA²³XX; Ordaz-Ortiz et al., 2005; Saulnier et al., 2009; see Fauré et al., 2009 for nomenclature). In addition, lichenase mainly released glucooligosaccharides of DP3 (3-O-β-cellobiosyl-D-Glc, G3) and DP4 (3-O-β-cellobiose-β-glucan, G4). These two fragments have been reported to account for 90% of all wheat β-glucan (Lazaridou and Biliaderis, 2007). Further small peaks corresponding to larger glucooligosaccharides (DP5, DP6) were also observed, but these accounted for small proportions of the total glucooligosaccharides and were not quantified in this study. The full analyses of the transgenic and control lines are reported in Supplemental Table S3.

Principal component analysis (PCA) was applied to HPAEC peaks obtained from transgenic and control samples (five transgenic and three control lines, three biological replicates). This revealed variation in AX and β-glucan composition of endosperm cell walls and showed clear separation between the transgenic and control lines. Figure 4 shows the similarity map for the two first principal components that together accounted for 95.2% of the total variance. The loading plot (Supplemental Fig. S5) indicated that the first principal component that accounted for 77.7% of the total variance related to G3 and G4 and thus indicated variation in the proportions of β-glucan in the samples. Principal component 2 mainly related to xylobiose (peak XX) and accounted for 17.5% of the total variation. Principal component 2 revealed small differences between the lines irrespective of the presence or absence of the transgene, showing that AX structure was not affected by the presence of the transgene. Since β-glucan and AX are both synthesized from UDP, Glc compensatory increases in AX could occur if β-glucan synthesis is down-regulated. However, this was not the case, the HPAEC analysis (Fig. 2; Supplemental Table S3) showed that the amount of AXOS released by the enzyme was similar in transgenic and in control lines.

The combined proportions of G3 + G4 decreased by a mean of 45.9% (range from 35.5% to 53.2%) in the transgenic lines (Fig. 5A), showing excellent agreement with the levels of suppression determined by total β-glucan analysis. The ratio of G3 to G4 fragments was also slightly higher (P < 0.001), with the mean in the control lines being 2.33 and that in the transgenic lines being 2.58. These G3 to G4 ratios are lower than those reported for β-glucan prepared from wheat bran (3.7–4.5; Cui et al., 2000; Li et al., 2006; Lazaridou and Biliaderis, 2007) and flour (3.04–3.84; Wood et al., 1991), but similar to those determined using the same method for developing and mature caryopses of wheat cv Hereward (ranging from 2.31–2.55 during the period from 10 DPA to maturity; authors’ unpublished results on the developmental series described by Shewry et al., 2009). The values are also within the range reported for β-glucan from a range of barley samples (1.8–3.5; for reviews, see Lazaridou and Biliaderis, 2007).
Table 1. Contents and properties of β-glucans in wholegrain samples of the transgenic and control lines

<table>
<thead>
<tr>
<th>Line (n = 3)</th>
<th>Total β-Glucan (g/100 g Dry Wt; Mean ± sd)*</th>
<th>WE β-Glucan (% Dry Wt; Mean ± sd)*</th>
<th>WE β-Glucan (% Total β-Glucan)</th>
<th>Average β-glucan (g/mol × 10^6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>9T (transgenic)</td>
<td>0.437 ± 0.09</td>
<td>0.05 ± 0.01</td>
<td>12.58</td>
<td>36.0 ± 4.5</td>
</tr>
<tr>
<td>7T (transgenic)</td>
<td>0.577 ± 0.09</td>
<td>0.06 ± 0.01</td>
<td>9.71</td>
<td>43 ± 6.4</td>
</tr>
<tr>
<td>21T (transgenic)</td>
<td>0.410 ± 0.10</td>
<td>0.06 ± 0.01</td>
<td>14.63</td>
<td>45.3 ± 4.2</td>
</tr>
<tr>
<td>4T (transgenic)</td>
<td>0.461 ± 0.09</td>
<td>0.06 ± 0.01</td>
<td>13.67</td>
<td>47.5 ± 6.1</td>
</tr>
<tr>
<td>15T (transgenic)</td>
<td>0.607 ± 0.09</td>
<td>0.07 ± 0.01</td>
<td>11.86</td>
<td>56.6 ± 1.7</td>
</tr>
<tr>
<td>20N (control)</td>
<td>0.872 ± 0.04</td>
<td>0.16 ± 0.02</td>
<td>18.24</td>
<td>84.8 ± 9.5</td>
</tr>
<tr>
<td>23C (control)</td>
<td>0.888 ± 0.04</td>
<td>0.14 ± 0.02</td>
<td>15.54</td>
<td>78 ± 5.1</td>
</tr>
<tr>
<td>15N (control)</td>
<td>0.852 ± 0.04</td>
<td>0.11 ± 0.02</td>
<td>13.57</td>
<td>83.9 ± 2.5</td>
</tr>
</tbody>
</table>

*Biological replicates.  Figures with different superscripts within a column are significantly different from each other (P < 0.01).  15N and 20N are null segregants from transformed lines.  23C is a nontransformed control line that had been subjected to the same transformation and regeneration conditions as the transformed lines.

These analyses were carried out on whole-grain samples, whereas the RNAi constructs were expressed under the control of the HMW subunit 1Dx5 gene promoter that is expressed only in the starchy endosperm and starting at about 10 to 12 DPA (Lamacchia et al., 2001), with significant levels of the encoded protein being detected from 14 DPA (Shewry et al., 2009). Hence, the true impact of the down-regulation on β-glucan content and properties would have been diluted by the presence of the β-glucan-rich aleurone but also by β-glucan synthesized in the young developing starchy endosperm.

β-Glucan in Developing Grain

The HMW subunit 1Dx5 promoter is only expressed from about 10 to 12 DPA (Lamacchia et al., 2001), whereas CSLF6 expression begins earlier (Fig. 1). Enzyme fingerprinting was therefore also carried out on developing caryopses of transgenic line 15 (chosen as the only line with a single copy of the transgene), the corresponding null segregant control line (15N), and the wild-type control line 23 at 10 DPA, 14 DPA, and 21 DPA (Fig. 5B). The results were consistent with the expression pattern of the HMW subunit promoter, with no differences being observed between the G3 + G4 contents of the three lines at 10 DPA but with significant (P = 0.002; P = 0.005) decreases in the proportions of G3 + G4 fragments in the transgenic line at 14 DPA (24.06% decrease) and 21 DPA (16.79% decrease). The difference in G3 + G4 between the 15T and 15N lines was therefore less than half as great at 21 DPA (Fig. 5B) as in mature seed (Fig. 5A). A possible explanation for this observation is that although the abundance of the CSLF6 transcript was low after 21 DPA the encoded enzyme was still active, leading to greater cumulative differences between β-glucan amounts in the mature 15T and 15N grain.

No statistically significant differences were observed in the ratios of G3 to G4 fragments in the three lines at 10 DPA and 14 DPA (P = 0.210 and 0.135, respectively) while at 21 DPA the transgenic line was not significantly different from the wild-type control line 23 (lowest value) but the null segregant control line 15N (highest value) was significantly different from the other two lines (P = 0.015). The ratios in the three lines at 10 DPA were consistent with those in the mature grain samples (2.01–2.65 compared with 2.25–2.62) whereas the ratios at 14 DPA and 21 DPA were lower (1.62–1.74 and 2.13–2.08, respectively). However, it is clear that these differences were not related to the presence of the transgene.

![Figure 3. HPAEC analysis of oligosaccharides produced by enzyme mapping of transgenic and control lines. A trace from transgenic line 9 (green) is overlaid on a trace from the null control line 15 (red). Detector response is measured in nC, retention time in minutes. G3 and G4 are glucan fragments released from β-glucan by digestion with lichenase; X and XX are Xyl and xylobiose released from AX by digestion with endoxylanase. Peaks 1 to 7 are AXOS released from AX by digestion with endoxylanase: 1 = XA3X, 2 = XA3XX, 3 = XA2+3XX, 4 = XA1XX, 5 = XA2XX, 6 = XA1XX, and 7 = XA2XX. All identified peaks were integrated (Supplemental Table S4) and used for PCA analysis (Fig. 4; Supplemental Fig. S5).](Image 311x181 to 551x370)
Effect on β-Glucan Molecular Weight

The content of hot-water-extractable β-glucan in wholemeal flour was 0.11% to 0.16% of dry matter in the control lines, with the highest content in line 20N. The content was significantly lower (P = 0.001) in the transgenic samples (0.05%–0.07% of dry matter), with the highest content being in sample 15T. These values correspond to between 9.7% and 14.6% of the total β-glucan in the transgenic lines and between 13.6% and 18.2% of the total β-glucan in the control lines. However, the differences between the transgenic and control lines were not significantly different (P = 0.069).

The Mr distribution of the hot-water-extractable β-glucan was determined by high-performance size-exclusion chromatography (Fig. 6). The Mr distribution in the control samples (lines 20N, 23C, and 15N) was either bimodal or polymodal (i.e. with two or three populations of β-glucan being present). The transgenic samples showed clear differences in Mr distribution, with the high Mr population of β-glucan being reduced to a small shoulder (lines 9T, 15T, and 21T are shown in Fig. 6; lines 4T and 7T had similar patterns and are therefore not shown).

The Calcofluor average Mr (Mr cf) of the hot-water-extractable β-glucan was 79 to 85 × 10^4 g/mol in the control samples, but was significantly lower in the transgenic samples (36–57 × 10^4 g/mol; Table I). Within the transgenic lines the highest Mr cf was observed in line 15T that had the smallest decrease in β-glucan compared to the control plants, and the Mr cf decreased with increasing reduction in β-glucan. The Mr percentiles at 10%, 50%, and 90% were also significantly lower for the transgenic lines (2.0–3.1 × 10^4 g/mol, 16.3–21.3 × 10^4 g/mol, and 89.7–160.3 × 10^4 g/mol, respectively) than for the control samples (4.6–5.2 × 10^4 g/mol, 40.4–44.5 × 10^4 g/mol, and 213–232 × 10^4 g/mol, respectively), with 15T again having the highest value of the transgenic lines. The Mr percentile at 90% also decreased with increasing down-regulation for the transgenic samples.
The decrease in β-glucan content determined by enzyme fingerprinting of the transgenic lines was significantly correlated with the decrease in the average $M_r$ ($r = 0.902$, $P = 0.036$) and with p90 ($r = 0.925$, $P = 0.024$).

**Immunolocalization of β-Glucan**

Immunolocalization of β-glucan was carried out in semithin sections of mature grain (Fig. 7, A–D) and developing caryopses (Fig. 7, E–J). Strong labeling of the aleurone layer was observed in all samples (which is consistent with the presence of thick cell walls comprising about 29% β-glucan; Bacic and Stone, 1981) and of the pericarp and nucellar epidermis in the developing caryopses. No differences were observed between the control and transgenic lines in labeling of the starchy endosperm cell walls at 10 DPA (Fig. 7, I and J) and little difference at 14 DPA (Fig. 7, G and H). However, clear differences were observed at 21 DPA (Fig. 7, E and F) and maturity (Fig. 7, A–D) with line 9T (Fig. 7, A and E) showing reduced labeling throughout the starchy endosperm when compared with the control lines and line T15 showing a less uniform pattern of reduction in labeling (Fig. 7, C and F; being greater in the outer cells of the tissue). These outer endosperm cells correspond to those that are added most recently during grain development by division of the aleurone cells (Bechtel and Wilson, 2003). The labeling differences are consistent with the control of RNAi expression by the glutenin promoter and with the amounts of β-glucan determined using the Megazyme kit and enzyme fingerprinting.

**DISCUSSION**

The amount and properties of β-glucan have important impacts on the utilization of both wheat and barley. Both cereals are widely used as feed for mono-
gastric livestock, notably pigs and poultry. In this case the high viscosity conferred by β-glucan has negative effects on feed intake, feed conversion rate, and weight gain, and may result in sticky feces when used to feed chickens (Hesselman et al., 1981). Although this problem may be alleviated by adding β-glucanase to degrade β-glucan in the feed and thus lower gut viscosity (Hesselman and Åman, 1986), reducing the amount and viscosity of the β-glucan in the grain is a more attractive option. β-Glucan content and M₄ are also important in malt and beer production, where high viscosity causes problems with haze formation and wort filtration (Bamforth, 1982), and in starch production where a high content of β-glucan may reduce the yield of starch. In contrast, high contents of soluble β-glucan are favored for food products as they may reduce serum cholesterol levels (McIntosh et al., 1991; Bourdon et al., 1999) as well as regulate blood Glc and insulin (Cavallero et al., 2002; Wood, 2007).

We have shown that down-regulation of the TaCSLF6 gene is sufficient to reduce the amount of β-glucan in the starchy endosperm of wheat to less than half of that in control lines. Expression of the transgene did not result in any consistent effects on the ratio of G3 to G4 fragments released on digestion with lichenase, suggesting that the synthesis of both 1,3 and 1,4 linkages was affected. This does not necessarily imply the encoded enzyme is capable of forming both linkages, as its removal could affect a multienzyme complex or stop the production of a substrate. However, the M₄ of the polymer was reduced, which could result from the deposition of incompletely synthesized, and therefore shorter, polysaccharide chains into the wall due to limiting activity of the glucan synthase. Only one form of CSLF was expressed in the developing grain at significant levels, which is consistent with the same enzyme catalyzing β-glucan synthesis in the developing starchy endosperm (the tissue studied here) and other grain tissues.

The TaCSLF6 gene reported here is a close ortholog of the barley HvCSLF6 gene (Supplemental Fig. S1) that was shown by Burton et al. (2008) to be similarly highly expressed in barley endosperm. A role of this gene in β-glucan synthesis in barley grain is also implied by analysis of the bg/l mutant line (Tonooka et al., 2009).

Rice CSLF genes were shown to mediate the synthesis of β-glucan in transgenic Arabidopsis (Burton et al., 2006). However, subsequent work from the same group showed that similar activity was displayed by transformation with a barley CSLH gene (Doblin et al., 2009). An ortholog of this gene is present in wheat but, as was the case in barley, the transcript levels in developing grain were very low (too low to show in Fig. 1A), which implies that it has little or no role in the synthesis of wheat grain β-glucan. Doblin et al. (2009) also suggested that the major role of the HvCSLH1 gene is the synthesis of β-glucan in secondary cell walls, based on in situ PCR localization in leaf tissues.

We therefore conclude that TaCSLF6 plays a major role in mediating β-glucan synthesis in developing grain of wheat and hence a key target for manipulation to increase or decrease total β-glucan. Furthermore, the effects of RNAi suppression on β-glucan M₄ indicate that indirect effects on M₄ and viscosity may occur by altering the relationship between enzyme activity and substrate availability.

MATERIALS AND METHODS

Array Analysis

Array analysis procedures were as previously reported (van et al., 2008). Wheat (Triticum aestivum) ESTs were assigned to rice (Oryza sativa) genes and assembled into contigs using the WinETS tool (Mitchell et al., 2007). PCR primers were designed to these sequences as the first step in cloning the CSL transcripts from grain cDNA.

cDNA Cloning and Construction of RNAi Expression Cassettes

Partial (CSLF6 and CSLD4) and full-length (CSLF6) cDNA was amplified by RT-PCR from RNA isolated from 14-DPA endosperm of cv Cadenza using PCR primers SP for CK201603, ASPI for RACE/FeDonger, SP_CSLF6 and ASP_CSLF6 (CSLF6), and SP_CSLD4 and ASP_CSLD4 (CSLD4) (Supplemental Table S4). cDNA was A tailed and cloned in pGEM-T Easy (Promega) and sequenced. For construction of RNAi expression cassettes DNA fragments corresponding to CSLF6 and CSLD4 were amplified using the PCR primers indicated in Supplemental Table S4 using partial cDNA clones as the templates. The RNAi fragment sequences were used to search all public domain wheat sequences and the only sequences that had stretches of identity of 20 bp or more were the target genes; namely CSLF6 for the CSLF6 fragment and CSLD2 and CSLD4 for the CSLD4 fragment. The reaction products were cloned, excised using BamHII and BglII, and ligated as inverted repeats on either side of the maize (Zea mays) Adh2 intron in the phasmid pHMW-Adh-Nos, an RNAi expression cassette constructed in pGEM-3Zf(+), under the control of the high M₄ glutelin subunit 1DX5 promoter (Supplemental Fig. S4; Lamacchia et al., 2001). The desired orientation of the inserts was confirmed by BamHI and BglII restriction digestion, PCR (using primers DoSPwfd with Adh5’rev and Adh3’fwd with Nos5’rev; Supplemental Table S4), and sequencing.

Production of Transgenic Wheat Lines and Analysis of Transgene Segregation and Copy Number

Transformation was carried out by particle bombardment (PDS1000, Bio-Rad) of immature scutella of cv Cadenza (Sparks and Jones, 2009) with RNAi constructs being co-bombarded with plasmid pCalNeo containing NptII for plant selection. PCR of genomic DNA was used to confirm the presence of the transgenes and the segregation in T1 plants: using primers Neo1 with Neo2 for the NptII gene and SP_CSLF6_BglII or SP_CSLD4_BglII with Nos5’rev for the CSLF6 and CSLD4 genes, respectively (Supplemental Table S4). The chi² goodness of fit test (with 1 degree of freedom for 3:1 ratios and 3 degrees of freedom for 9:3:3:1 ratios) was used to separate plants into genetic classes. Genomic DNA was subjected to quantitative PCR using TaqMan probes to determine transgene copy number and zygosity of individual plants (Supplemental Table S5; analysis performed by iDNA Genetics).

Plant Material and Growth Conditions

Plant material for transformation was produced according to Sparks and Jones (2009). All other plant material was grown in the glasshouse with 18°C day and 14°C night temperatures with a 16-h photoperiod provided by natural light supplemented with banks of Sun-T 400 W sodium lamps (Osram, Ltd.) giving 400 to 1,000 µmol m⁻² s⁻¹ photosynthetically active radiation. T2 grain from five homozgyous transgenic wheat lines carrying RNAi designed against CSLF6 (lines 9T, 7T, 21T, 4T, and 15T) and three controls (lines 20N, 15N, and 23C) were sown to provide mature T3 grain for analyses. For each line three 25-cm pots, with four plants per pot, were randomized and grown in the same glasshouse compartment at the same time. Grain weight
was determined using four replicate samples (except for line 7T for which only two replicate samples were available) of grain from the main tillers of individual plants for each of five transgenic and three control lines.

**qRT-PCR**

Seeds of line 1ST and segregating nulls, 14 DPA, were frozen in liquid nitrogen and stored at ~80°C. For RNA extraction seeds were freeze dried and the endosperm extracted by peeling off the outer layers. After homogenization in a Tissuelyser (Qiagen) total RNA was extracted according to Wan et al. (2008). RT and qRT-PCR were performed as described in Pelinly et al. (2008). The expression of the native gene as well as the transgene was determined with the primers SP_CS56F, RNe_7593 and ASP_CS56F, RNe_2828 that give an amplicon within the RNAi construct. To test for the down-regulation of the native gene the sense primer was chosen to be just outside of the construct (SP_CS56F, RNe_2669) with the same antisense primer amplifying an extra 30 bp at the 5’ end of the RNAi construct and all but 3 bp at the 3’ end. The expression of these amplicons was normalized against Ta2652 (pfTYW19, Ta2652, gPCR_F and pfTYW20, Ta2652, gPCR_R), an EST that was shown to be stably expressed in wheat grain development (Wan et al., 2008). Results are displayed as relative expression of four transgenics compared to between four and seven nulls and significant differences were calculated by ANOVA after data transformations as proposed by Rieu and Powers (2009).

**β-glucan Analysis**

For total β-glucan analysis, enzyme fingerprinting and determination of β-glucan M, distribution finely ground freeze-dried developing caryopses (enzyme fingerprinting only) or wholegrain flour (all methods) was used. Three biological replicates consisting of 30 seeds each from the main tillers of three individual plants (a total of 90 seeds per replicate) were analyzed.

**Total β-glucan was determined using a Megazyme kit. Endoxylanase (EC 3.2.1.8) from *Trichoderma viride* was purchased from Megazyme (Xylanase M1).** The activity of the enzyme preparation determined by the supplier on WE-AX (40°C, pH 6.5) was 1,670 units/mL, pH optimum is 4.5 to 5.0. Lichenase (endo-1,3(4)-β-D-glucanase, EC. 3.2.1.73) from *Bacillus subtilis* was purchased from Megazyme. The activity of the enzyme preparation on barley (*Hordeum vulgare*) β-glucan (40°C, pH 6.5) determined by the supplier was 1,000 units/mL, pH optimum is 6.5 to 7.0.

Enzyme fingerprinting samples (100 mg) weighed in an Eppendorf tube (2 mL) were treated with 1.5 mL ethanol solution (80%, v/v) in a boiling water bath for 5 min. The supernatant was removed and the residue successively washed with 1.5 mL ethanol (80%, v/v) and 1.5 mL ethanol (95%, v/v) solutions. The supernatant was then discarded and the residue dried for 2 h in an oven at 40°C.

Samples were then digested overnight (16 h) with 2 units lichenase and 16 units endoxylanase (Ordz-Ortiz et al., 2005; Saulnier et al., 2009). The degradation was carried out in water at a pH 6.0 compatible with the use of both enzymes in one run. After centrifugation, supernatant of the digested samples was diluted with water (1/40) and 10-μl aliquots were injected on a Carbospec PC-A (5 × 250 mm) analytical column (Dionex) run at 25°C with a flow rate of 1 mL/min. AXOS and GOS were separated using the following elution conditions with Ultrapure water (A), 1 mM NaOAc (B), and 0.5 mM NaOH (C): 0 min (A: 80%, C: 20%), 35 min (A: 60%, B: 20%, C: 20%); 36 min (A: 80%, C: 20%), hold up to 60 min. Detection was realized with a TSP EC2000 pulse amperometric detector (Thermo Separation Products) using the pulse potentials E1 = +0.05 V, E2 = +0.6 V, and E3 = −0.6 V. Typical chromatograms are shown on Figure 2. Peak identification was based on the retention times of reference compounds previously isolated (Ordz-Ortiz et al., 2004, 2005; Saulnier et al., 2009). It was checked that in the conditions used, GOS and especially G3 and G4 were not coeluted with peaks generated by the endoxylanase (Saulnier et al., 2009). Overlay of chromatograms was realized using raw data and adjustment of the baseline to the same level without peak normalization.

For PCA (Fig. 3; Supplemental Fig. S5) peak areas (Supplemental Table S3) were normalized for each chromatogram as follows: $S_{\text{Peak}} = (S_{\text{Peak}} / \sum_{i=1}^{100}$ $S_{\text{Peak}})$, with $S_{\text{Peak}}$ being the normalized area of peak $i$ and $S_{\text{Peak}}$, the area of peak $i$ (Ordz-Ortiz et al., 2005; Saulnier et al., 2009).

For determination of M, distribution, samples were extracted with boiling water containing thermostable α-amylase and analyzed by high-performance size-exclusion chromatography with fluorescence detection as described by Rimsten et al. (2003), except that 0.0025% Calcofluor was used and that the pH of the Calcofluor solution was not adjusted. This detection method is selective for β-glucan in extracts and the fluorescence response is independent of M, it will, however, exclude M; below 10^(4) (Munck, 1989). The M is therefore given as Calcofluor average M, (M), which is the same as weight average M, for samples that only contain high M, β-glucan. (M) and percentiles (p10, p50, and p90) describing the M, at which 10%, 50%, and 90% of the distribution fall below were calculated by using a calibration curve established with narrow M, β-glucan standards (Rimsten et al., 2003). The content of β-glucan in the extracts (content of water-extractable β-glucan) was also calculated by using the area of the peak as described by Rimsten et al. (2003). The repeatability was <15% for the three biological replicates analyzed. The results were evaluated statistically with ANOVA, generalized linear model procedure, and Tukey’s pairwise comparison (Minitab 15).

**Immunocytochemistry**

One-millimeter sections were fixed in 3% paraformaldehyde and 2% glutaraldehyde in phosphate buffer (0.1 M, pH 7.2), dehydrated in a graded aqueous ethanol series, and infiltrated with London Resin White acrylic resin. One-micrometer sections were mounted onto multwell glass sides and preincubated in phosphate-buffered saline (PBS) containing 3% (w/v) bovine serum albumin (BSA) for 30 min and then incubated for 1 h with PBS containing 1% (w/v) BSA, 0.01% Tween 20 (PBS-BSA), and 1,000 dilution of mAb anti-β-glucan (Mekile et al., 1994; Biosupplies Pty Ltd.). After extensive washing in PBS they were incubated for 1 h in the dark with marker-conjugated goat anti-mouse antibody labeled with Alexa Fluor 546 (Molecular Probes, distributed by Interchim) diluted 100-fold in PBS 0.3% BSA. Nonspecific staining was identified using controls omitting the primary antibody. Sections were examined with a LEICA DM2500 microscope equipped with epifluorescence using a band-pass filter at 515 to 560 nm for excitation filters and fluorescence detected at >570 nm.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number AM743080.

**Supplemental Data**

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

**Supplemental Figure S1.** Alignment of wheat, barley, and rice CSLF6 protein sequences, with that encoded by the *OsCSLF2*.

**Supplemental Figure S2.** Sequence of 3’ end of *TaCSLF6* gene coding region (accession AM743080.2).

**Supplemental Figure S3.** Sequence of 3’ end of *TaCSLD4* gene coding region and *CSLD2/4* RNAI fragment.

**Supplemental Figure S4.** Schematic representation of construction of RNAi expression vectors.

**Supplemental Figure S5.** Loading plot for PCA of the enzyme fingerprints of mature seeds of five control and three transgenic lines.

**Supplemental Table S1.** Transcript abundance for all CSL genes represented on wheat Affymetrix GeneChip.

**Supplemental Table S2.** Wheat EST counts for sequences mapped to rice CSL genes.

**Supplemental Table S3.** Enzyme mapping of cell wall polysaccharides in mature grain, determined by HPAGE.

**Supplemental Table S4.** DNA sequences and targets for PCR primers used in this work.

**Supplemental Table S5.** Segregation and copy number analysis of 16 transgenic wheat lines.

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