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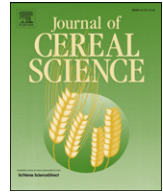
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Variant high-molecular-weight glutenin subunits arising from biolistic transformation of wheat

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ABSTRACT

Genetic transformation *via* the biolistic method has been used to introduce genes encoding natural and novel high-molecular-weight glutenin subunits (HMW-GS) into wheat. The appearance of new seed proteins of sizes not predicted by the transgene coding sequences was noted in some experiments. In this report, the identities of thirteen of these novel proteins were determined by tandem mass spectrometry (MS/MS). Seven different proteins larger than and two proteins smaller than the native protein were shown to contain peptides from 1Dx5. A novel protein found in some progeny of crosses between a transgenic plant and Great Plains winter wheats was larger than but contained several peptides from 1Dy10. In one line, a protein larger than and a protein smaller than HMW-GS each contained peptides from the N- and C-terminus of 1Dx5 and from the repeat region of 1Dy10. In a sixth transgenic line, the native *Bx7* gene was apparently replaced by a gene that encodes a larger version of 1Bx7. The variant proteins accumulate in the polymeric protein fraction, indicating that they can form inter-molecular disulfide bonds. These results show that novel proteins found in some transformants are encoded by altered versions of either the transforming or endogenous HMW-GS genes.

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1. Introduction

Genetic transformation is one of several approaches used to understand the contributions of individual genes and seed proteins to wheat (*Triticum aestivum*) end-use functionality. The most-studied wheat quality genes are those encoding high-molecular-weight glutenin subunits (HMW-GS), which typically constitute 5–10% of seed storage proteins and are important determinants of dough strength and elasticity (reviewed and referenced in Shewry et al., 2003). Addition of native or modified HMW-GS genes to wheat generally results in accumulation of proteins of the expected sizes in addition to those encoded by the native genes (Altpeter et al., 1996; Alvarez et al., 2000; Barro et al., 1997; Blechl and Anderson, 1996; He et al., 1999, 2005; León et al., 2009). Transformants carrying expressed transgenes encoding native HMW-GS produce flours that make doughs with increased mixing times and tolerances compared to their non-transformed parent lines (Alvarez et al., 2001; Barro et al., 1997, 2003; Blechl et al., 2007; Darlington et al., 2003; He et al., 1999; León et al., 2009; Popineau

et al., 2001; Rakszegi et al., 2005; Rooke et al., 1999; Vasil et al., 2001). However, some HMW-GS gene transformation events contain proteins of unpredicted sizes in addition to or instead of those encoded by the introduced transgenes (Altpeter et al., 1996; Alvarez et al., 2000; Barro et al., 1997; Blechl and Lin, 2007; He et al., 2005). In this report, we characterize thirteen of these proteins by tandem mass spectrometry (MS/MS) and show that they are size variants of HMW-GS.

2. Materials and methods

2.1. Derivation of transformed lines

The transformation events characterized here were produced in the same experiments as those described previously (Blechl et al., 2007). Briefly, cultivar 'Bobwhite' was co-transformed with the Ubi::BAR selection plasmid and separate DNA plasmids carrying the wheat genomic *EcoR1* fragments from cultivar 'Cheyenne' that encode either 1Dx5 or 1Dy10 (Anderson et al., 1989). After selection with bialaphos, putative transformants were identified by changes in the levels of 1Dx5 and/or Dy10 in T₁ seeds. Homozygous progeny from each event were identified by SDS-PAGE of seed proteins and used to establish true-breeding lines (Blechl et al., 2007). Lines were named by the introduced HMW-GS gene(s), e.g., Dx5 + Dy10, followed by a unique event letter.

Abbreviations: HMW-GS, high-molecular-weight glutenin subunits; MS/MS, tandem mass spectrometry; RP-HPLC, reverse-phase liquid chromatography.

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An additional event “B52a-6” was characterized after the transgene had been crossed into hard winter wheat lines. The derivation and characterization of those lines are described in Graybosch et al. (2011). The original ‘Bobwhite’ transformant was co-transformed by Ubi::BAR and a plasmid carrying the native gene encoding 1Dy10.

2.2. Protein characterization

2.2.1. Mass spectrometry

To prepare individual protein bands for mass spectrometry, SDS-PAGE was performed as described previously (Blechl and Anderson, 1996 for Fig. 1C; Blechl et al., 2007 for Figs. 1A, B, D and 3). In some experiments, proteins were reduced and then alkylated by incubation for 30 min at room temperature in 4% (w/v) iodoacetamide, 50 mM Tris–HCl (pH 8.8), 2% (w/v) SDS before loading onto the gels (method adapted from Görg et al., 2004). Bands corresponding to novel proteins were excised from the gels and placed into the reaction plate of a DigestPro (Itavis, Koeln, DE), where the protein-containing spots were destained, reduced, alkylated, and digested with trypsin. The resulting tryptic peptides were automatically eluted into a 96-well plate in preparation for LC-MS/MS.

Identification of protein-containing bands was carried out as previously described (Vensel et al., 2005). The 96-well collection plate from the DigestPro was inserted into the autosampler of the Reverse-Phase (RP)-HPLC system that was interfaced to a QSTAR PULSAR *i* quadrupole time-of-flight mass spectrometer (Applied Biosystems/MDS Sciex, Toronto, CA). Conditions for operation of the instrument were as previously described (Vensel et al., 2005). A minimum of four independent samples for each protein variant were excised from gels and analyzed.

2.2.2. Protein identification

Extraction of peak lists from the QSTAR Analyst QS wiff files and their conversion to text files was carried out using Mascot Daemon (<http://www.matrixscience.com/>). MS/MS samples were analyzed using Mascot (Matrix Science, London, UK; version 2.1.04) and X! Tandem (www.thegpm.org; version 2006.04.01.2). Mascot was set up to search a specially constructed database with all Triticace sequences (as of January 2009) and the protein sequences of Phosphinothricin Acetyl Transferase (Accession No. P16426 encoded by the BAR gene), beta lactamase (Accession No. AAB59737.1 encoded by the *bla1* gene for ampicillin resistance), and β -galactosidase

(Accession No. P00722 encoded by the *lacZ* gene). The specified digestion enzyme was trypsin. Using the post-analysis software package Mascot and X! Tandem (Craig and Beavis, 2004), results were searched with a fragment ion mass tolerance of 1.00 Da and a parent ion tolerance of 1.00 Da. Iodoacetamide derivative of cysteine was specified in Mascot and X! Tandem as a fixed modification while oxidation of methionine and deamidation at N and Q were specified as variable modifications.

Validations of tandem MS-based peptide and protein identifications were performed using Scaffold (Scaffold version 2.06.01, Proteome Software Inc., Portland, OR). Protein probabilities were assigned by the Protein Prophet algorithm (Keller et al., 2002; Nesvizhskii et al., 2003). The protein identifications reported in Table 1 were established at 100.0% probability and contain at least 2 peptides identified with greater than 90% probability.

2.2.3. Protein solubility

Whole seeds homozygous for the transgenes were ground and extracted three times with 50% propanol. The residue was dissolved in SDS-PAGE sample buffer containing 2-mercaptoethanol and loaded onto 4–12% gradient gels for electrophoresis (Blechl et al., 2007).

3. Results

Previously we reported the generation of multiple independent transformation events obtained by particle bombardment with the Ubi::BAR selection gene and plasmids containing the native genes encoding HMW-GS 1Dx5 and/or 1Dy10 (Blechl et al., 2007). Analyses by SDS-PAGE of seed proteins showed that 23 of 30 transformants had an increase in the protein band(s) corresponding to HMW-GS 1Dx5 and/or 1Dy10, as expected (open circles in Fig. 1), while the other seven exhibited transgene-mediated co-suppression (Blechl and Lin, 2007; lane 2 of Fig. 1B). However, seven of the events with increased expression of 1Dx5 and/or 1Dy10 also contained one or more novel protein bands (arrowheads in lanes 1, 3–7, 9, 11, and 12 of Fig. 1). Most of the novel proteins were of higher molecular weight than native HMW-GS, although lower molecular-weight bands were also apparent in three transformants (lanes 1, 3 and 4, Fig. 1). Among 13 transformants with the 1Dx5 gene, we observed three lines with variant proteins. Among 10 transformants with both 1Dx5 and 1Dy10 genes, we observed four such lines (Blechl and Lin, 2007). Among 7 lines transformed with

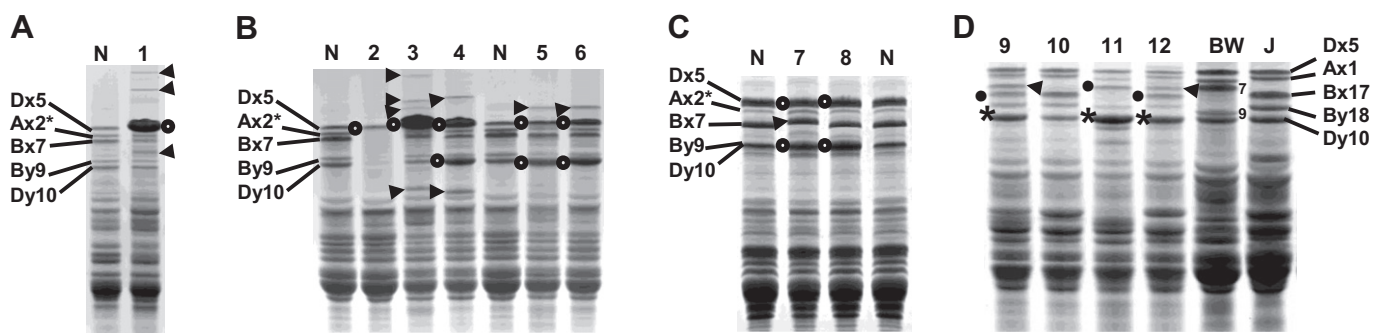


Fig. 1. SDS-PAGE of seed proteins from transgenic (numbered) and non-transformed ‘Bobwhite’ (N, BW) and ‘Jagger’ (J) wheat plants. Arrowheads to the left or right of the lanes indicate the positions of the variant proteins analyzed in this paper. Open circles indicate the subunit(s) with increased accumulation due to transgene expression. The names and positions of HMW-GS native to ‘Bobwhite’ are shown to the left of the first three panels. The names and positions of HMW-GS native to ‘Jagger’ are shown to the right of panel D. A. Transgenic line Dx-M (1) containing three variants. B. Transgenic lines are as follows: a segregant of transgenic event Dx5-F that showed transgene-mediated suppression (2); a different segregant of transgenic event Dx5-F, called Dx5-F* (3), that contained four proteins not found in (2) or in non-transformed Bobwhite extracts; Dx5 + Dy10-H (4) containing two variants; Dx5 + Dy10-I (5); Dx5 + Dy10-C (6). C. Transgenic line Dx5 + Dy10-A: a segregant called Dx5 + Dy10-A* (7), that contained a variant protein that migrated more slowly than native 1Bx7 (arrowhead), and a different segregant (8) that contained unaltered 1Bx7. D. Individual seeds (9–12) from a segregating population derived from crosses between transgenic event B52a-6 and Nebraska hard winter wheat lines (Graybosch et al., 2011). Asterisks indicate over-expressed 1Dy10 encoded by the transgene. Closed circles indicate other HMW-GS identified by tandem MS. The positions of subunits 1Bx7 (7) and 1By9 (9) are indicated to the right of the BW lane.

Table 1
Protein and peptide identities for variants in transgenic wheat lines.

Transgenic event [figure, gel lane] variant band	Number of independent samples with identified proteins	Proteins identified with 100% probability (# samples)	Average % of protein covered by detected peptides	Peptides identified by Scaffold with probabilities of 90% or higher (# samples)
Dx5-M [Fig. 1A, 1] largest	7 alkylated	1Dx5 (7) Sucrose synthase I (2)	4.5 2	2 (2), 5 (7), 6 (5), *10 (3), 12 (2) Two unique
Dx5-M [Fig. 1A, 1] 2nd largest	7 alkylated	1Dx5 (7) Sucrose synthase I (3)	5 3.5	2 (5), 5 (5), 6 (5), *10 (4), 11 (1), 12 (5) Two or three unique
Dx5-M [Fig. 1A, 1] between native 1Bx7 and 1By9	8 alkylated	1Dx5 (8) Sucrose synthase I (4) Sucrose synthase II (4)	7 22 12.5	2 (6), 3 (1), 4 (1), 5 (8), 6 (8), 8 frag (1), *10 (3), 11 (2), 12 (7) Eleven to eighteen unique Two to seven unique
Dx5-F* [Fig. 1B, 3] largest	6 non-alkylated 2 alkylated	1Dx5 (6) 1Dx5 (2)	7.5 3	1 (5), 2 (4), 3 (6), 4 (1), 5 (5), 6 (3), 11 (4), 12 (3) 5 (2), 6 (2)
Dx5-F* [Fig. 1B, 3] 2nd largest	7 non-alkylated 3 alkylated	1Dx5 (7) 1Dx5 (3)	10 4	1 (7), 2 (6), 3 (7), 4 (3), 5 (7), 6 (4), 11 (5), 12 (4) 2 (2), 5 (3), 6 (2), *10 (1)
Dx5-F* [Fig. 1B, 3] 3rd largest	4 non-alkylated 4 alkylated	1Dx5 (4) 1Dx5 (4)	9 4.5	1 (3), 2 (2), 3 (4), 5 (4), 6 (4), 11 (4), 12 (2) 2 (2), 5 (4), 6 (4), *10 (1), 12 (1)
Dx5-F* [Fig. 1B, 3] smallest	8 non-alkylated	1Dx5 (8) Protein disulfide isomerases (5) Globulin 3 (4)	10 28 23	1 (8), 2 (7), 3 (8), 4 (4), 5 (8), 6 (7), 11 (4), 12 (3) Two to eighteen unique Nine or ten unique
	8 alkylated	1Dx5 (7) Globulin 3 (8) Phosphoglucosyltransferase (1) Unknown 60 kDa (1)	3 39.5 4 7	5 (7), 6 (7), 12 (1) Fourteen to twenty-six unique Two unique Two unique
Dx5 + Dy10-H [Fig. 1B, 4] larger	4 non-alkylated	1Dx5 (4) 1Dy10 (4)	7.5 4	1 (4), 2 (2), 3 (3), 5 (4), 6 (4), 11 (1), 12 (1) 26 (1), 38 (3), 39 (4)
	8 alkylated	1Dx5 (8) 1Dy10 (1)	4.5 7	2 (1), 3 (3), 5 (8), 6 (8) *37 (1), 38 (1)
Dx5 + Dy10-H [Fig. 1B, 4] smaller	4 non-alkylated	1Dx5 (4) 1Dy10 (2)	10.5 2	1 (4), 2 (4), 3 (4), 4 (1), 5 (4), 6 (4), 11 (4), 12 (3) Two versions of 38 (2)
	8 alkylated	1Dx5 (8) 1Dy10 (5) Globulin 3 (8)	10.5 4.5 20	2 (8), 3 (8), 4 (5), 5 (8), 6 (7), *10 (6), 11 (1), 12 (8) *37 (2), frag *37 (3), 38 (5) Four to ten unique
Dx5 + Dy10-I [Fig. 1B, 5] large	4 non-alkylated 4 alkylated	1Dx5 (4) 1Dx5 (4)	9 9.5	1 (4), 2 (3), 3 (4), 5 (4), 6 (4), 11 (3), 12 (2) 2 (4), 3 (4), 5 (4), 6 (4), *10 (4), 11 (2), 12 (4)
Dx5 + Dy10-C [Fig. 1B, 6]	7 non-alkylated	1Dx5 (7) 1Dy10 (6)	14.5 8	1 (7), 2 (7), 3 (7), 4 (7), 5 (7), 6 (7), 7 (1), 8 (2), 11 (6), 12 (6) 25 (1), 26 (5), 27 (3), 30 (2), 32 (1), 33 (2), fragment 34 (1), 35 (1), 39 (4)
	7 alkylated	1Bx7 (4) 1Dx5 (7)	6.5 10	14 + 15 (4), 15 (1), 16 (4), 18 (1), 11 (4) 2 (7), 3 (5), 4 (2), 5 (7), 6 (7), frag 8 (1), *10 (7), 11 (3), 12 (7)
Dx5 + Dy10-A* [Fig. 1C, 7]	4 non-alkylated	1Bx7 (3) 1Ax2 (2)	26.5 13.5	13 (3), 14 + 15 (3), 16(2), 17 (3), 18 (2), 19 (3), 20 (2), 21 (1), *23 (1), 24 (2), 11 (3) Two or seven unique including one in N-terminus (2) and six in repeats
	4 alkylated	1Dx5 (4) Sucrose synthase I (1) 1Bx7 (4) 1Ax2* (4)	6.5 4 15 13	1 (1), 2 (1), 3 (3), 4 (1), 5 (2), 6 (4), 11 (4) Two unique 14 + 15 (2), 16 (4), 17 or fragment (4), 19 (2), 20 (1), 21 (4), 23 (3), 24 (4) Three to eight unique including two in N-terminus (2) and six in repeat regions (4)
Dy10 – B52a-6 [Fig. 1D, 9] Larger	4 alkylated	1Bx7 or 17 (4) 1Ax2* or 1Ax1	13.5 8	14 + 15 (1), 16 (4), 17 frag (2), 19 (1), 21 (4), *22 (1), *23 (4), 24 (4) Two to six unique, including one in N-terminus and five in repeat regions
		1Dy10 (4) or 1By9 (2) Unknown 90 kDa protein (1) Ortho dikinase (1)	4 4 6	26 (4), 27 (4), 30 (2) Two unique (1) Two unique (1)
Dy10 – B52a-6 [Fig. 1D, 9] smaller	4 alkylated	1Bx7 or 1Bx17 (4) 1Ax2* or 1Ax1(2) Sucrose synthase II (4) Elongation factor (2)	15.5 4.5 18 16	14 + 15 (4), 16 (4), frag 17 (2), 18 (1), 19 (2), 21 (4), *23 (4), 24 (4) Two unique in repeat region (2) Eight to fourteen unique Two unique
Dy10 – B52a-6 [Fig. 1D, 11]	4 alkylated	1Bx7 or 1Bx17 (4)	7.5	11 (1), 16 (4), 19 (1), 21 (1), *23 (4), 24 (4)
Dy10 – B52a-6 [Fig. 1D, 12] larger	4 alkylated	1Dy10 (4)	10.5	25 (4), 26 (3), 27 (4), 30 (4), 31 (3), 32 frag (3), 38 (3)
Dy10 – B52a-6 [Fig. 1D, 12] smaller	4 alkylated	1Bx7 or 1Bx17 (4) Sucrose synthase II (4)	8 14.5	16 (4), 19 (3), *23 (4), 24 (4) Six to thirteen unique peptides

The GenBank Accession numbers for the identified proteins are 1Dx5 gi|110341804|, 1Bx7 gi|262205164|, 1Ax2* gi|110341791|, 1By9 gi|22090|, 1Dy10 gi|164457873|, Sucrose synthase I gi|3393044|, Sucrose synthase II gi|3393067|, Protein disulfide isomerases gi|222446340|, gi|12056115|, Globulin 3 gi|215398470|, phosphoglucosyltransferase gi|18076790|, unknown 60 kDa protein gi|257693194|, unknown 90 kDa protein gi|257716429|, Pyruvate orthophosphate dikinase gi|32400838|, Elongation factor gi|58500286|.

The "Peptides Identified" in the last column of Table 1 are presented from Figure 2.

only the 1Dy10 gene, we found one with an unexpected protein in the HMW-GS region of gels. This line was crossed with Great Plains winter wheats and further characterized in those backgrounds (Graybosch et al., 2011). Seeds from these derived lines can contain seed storage proteins other than those in the transformed cultivar 'Bobwhite' (BW in Fig. 1D), including HMW-GS 1Ax1, 1Bx17 and 1By18 found in winter wheat 'Jagger' (J in Fig. 1D). Protein extracts from four individual seeds of the derived lines are shown in lanes 9–12 of Fig. 1D.

In two events, we noted segregation of progeny with different seed protein patterns from a single regenerated transformed plant. One such event, Dx5-F, had some progeny seeds that exhibited transgene-mediated co-suppression (lane 2 of Fig. 1B) and other seeds (lane 3 of Fig. 1B) that showed increased accumulation of 1Dx5, some accumulation of the other HMW-GS, and four additional protein bands that were not present in either the suppressed segregant or in non-transformed 'Bobwhite' (lanes N of Fig. 1B). In the first generation progeny, we were able to select segregants with the different transgene expression patterns, which then bred true, producing homozygous progeny with the uniform seed protein compositions seen in lanes 2 and 3 of Fig. 1B. We designated the line with the variant bands Dx5-F*. Another transgenic event, line Dx5 + Dy10-A, also produced two types of progeny with overexpression of 1Dx5 and 1Dy10 subunits. In some T₁ seeds, the native 1Bx7 band was replaced by a protein of slightly slower mobility (Fig. 1C, compare lanes 7 and 8). The difference in migration was apparent in large (15 × 15 cm) 10% SDS-PAGE gels (Blechl and Anderson, 1996), but not in a smaller (10 × 10 cm) 4–12% gradient gel format (Blechl et al., 2007). Homozygous lines with the native 1Bx7 or the variant protein were derived from the primary transformant. The line with the variant was called Dx5 + Dy10-A*.

The variant bands were excised from SDS-PAGE gels, digested with trypsin, and subjected to MS/MS. Fig. 2 shows the protein sequences of 1Dx5 (A), 1Bx7 (B) and 1Dy10 (C) with the locations of arginine (R), lysine (K) and cysteine (C) amino acids highlighted. The peptides that were actually identified by the MS/MS analyses reported here are assigned numbers. As expected for trypsin digestion, all the detected peptides ended in K or R. However, not all the peptides identified were directly preceded by K or R. The starts of those peptides are marked by vertical lines and their numerical designations are preceded by asterisks (Fig. 2). Table 1 lists the proteins identified in each variant band with 100% certainty by the Scaffold™ software, based on the presence of at least 2 peptide sequences identified with certainties of 90% or more. Also shown is the average % of the identified protein that was covered by identified peptides and the frequency of the HMW-GS peptides numbered in Fig. 2 in the samples analyzed for each variant. In early experiments, samples were loaded after reduction onto the gels, as usual for analytical gels. In some of these "non-alkylated" samples, peptides from two or three different HMW-GS were found. To rule out that these were from dimers or other multimers between individual subunits that might migrate at positions larger than native 1Dx5, later experiments were performed with proteins that had been alkylated prior to gel loading. In most cases in which both alkylated and non-alkylated samples were analyzed (the four variants in Dx5-F*, the two variants in Dx5 + Dy10-H, the large variant in Dx5 + Dy10-I), there was no difference in the HMW-GS identified. In the cases of Dx5 + Dy10-C and Dx5 + Dy10-A*, fewer different HMW-GS were identified in the alkylated samples than in non-alkylated ones.

MS/MS unambiguously identified the large variants in Dx5-M, Dx5-F*, and Dx5 + Dy10-I as HMW-GS 1Dx5. Every sample that was assigned any protein identity was assigned this identity. Protein coverage ranged from 4.5 to 10% and both N- and C-terminal

peptides were detected. For the other variants, more than one protein was identified in the majority of samples, but each sample included at least one HMW-GS. Fifteen of 16 samples of the small variant (around 57 kDa) in event Dx5-F* contained peptide 5 from 1Dx5 and many samples contained other N- and C-terminal peptides. Average coverage was 10% in the non-alkylated samples. The other proteins identified in these samples can be accounted for by their co-migration in this region of the gels. In particular, globulin 3 was identified in samples from the same gel region of non-transformed 'Bobwhite' (data not shown).

Three different HMW-GS were identified in non-alkylated samples of Dx5 + Dy10-C. All samples contained 1Dx5. The seven alkylated samples had only 1Dx5. The presence of 1Dy10 and 1Bx7 in some of the non-alkylated samples may be due to the presence of HMW-GS dimers or other multimers in this region of the gel. We occasionally found 1Dy10 in this region in lanes containing non-alkylated samples of non-transformed 'Bobwhite' (data not shown).

The larger and smaller variants from event Dx5 + Dy10-H also contained peptides from two different HMW-GS, but these were found in both alkylated and non-alkylated samples. With the exception of one non-alkylated sample of the larger variant, the only peptides from 1Dy10 were from the repeat region: *37, its fragment, and 38 (double-underlined in Fig. 2). N-terminal peptides of 1Dx5 were found in most of the samples of both variants. C-terminal peptides of 1Dx5 were found in nearly all the samples of the small variant. The simplest explanation for these findings is that the variants in Dx5 + Dy10-H are chimeric proteins consisting of the N- and C-terminus of 1Dx5 and at least part of the repeat region of 1Dy10.

The variant protein in Dx5 + Dy10-A* migrates more slowly than native 1Bx7 from 'Bobwhite' (lanes 7 and 8 in Fig. 1C) and was found instead of the native protein in segregating T₁ seeds. MS/MS identified 1Bx7 peptides in 3 of 4 non-alkylated samples and 4 of 4 alkylated samples. The presence of 1Dx5 and 1Ax2* in most of the samples is probably due to contamination of the gel slice with these proteins that migrate nearby. The presence of a version of 1Bx7 is difficult to explain since the plasmids used to make the transformant encoded 1Dx5 and 1Dy10, not 1Bx7.

The identity of the variant protein in winter wheat lines derived from crosses with event B52a-6 was assigned by comparing results from MS/MS identifications of several protein bands (arrowheads and closed circles in Fig. 1D) in the HMW-GS region of three seeds (lanes 9, 11 and 12 in Fig. 1D). Since we did not have the original event for comparison and since these seeds contain potentially co-migrating HMW-GS from parents besides 'Bobwhite', it was important to eliminate potentially co-migrating native HMW-GS from consideration. The bands labeled with circles in lanes 9, 11 and 12 of Fig. 1D were identified as containing an Ax (lane 9 only) and a Bx HMW-GS. The peptides detected by MS/MS in these samples do not distinguish between Ax1 and Ax2* or between Bx7 and Bx17. However, the positions of the bands in the gels relative to subunits in 'Bobwhite' (BW) and 'Jagger' (J) suggest that the proteins tagged by circles are or include 1Bx17 in lanes 9 and 12 and 1Bx7 in lane 11. The band tagged by an arrowhead in lane 9 contains Ax-, Bx- and y-type HMW-GS. The peptides in two of the samples of this band unambiguously identify 1Dy10, while the peptides in the other two samples do not distinguish between 1By9 and 1Dy10. This band is evidently a mixture of HMW-GS, including Bx7 and the variant protein, which is bigger than but related to 1Dy10. This conclusion is reinforced by the unambiguous identification of the variant band in lane 12 as 1Dy10 in all four samples. Average protein coverage for these samples is 10.5% and peptides were identified from both the N-terminus and repeat regions of the native protein.

Peptides 1, 3, 4, 11, 13 and 14 contain cysteine residues conserved in all x-type subunits. The presence of these cysteines in

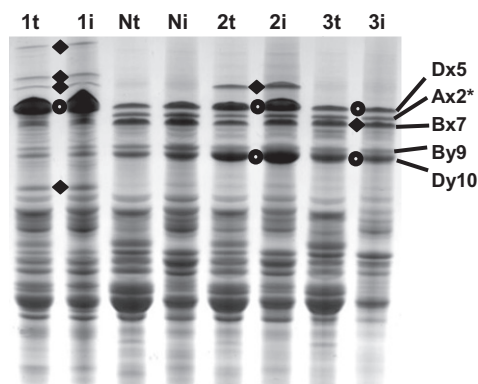


Fig. 3. SDS-PAGE of total seed proteins (t) and those insoluble in 50% propanol (i) from non-transformed 'Bobwhite' (N) and transgenic lines D5-F* (1); D5 + Dy10-C (2); and D5 + Dy10-A* (3). Diamonds show the positions in adjacent lanes of the variant bands analyzed in this report. Open circles between adjacent lanes indicate the subunits with increased accumulation due to transgene expression. The names and positions of HMW-GS native to 'Bobwhite' are shown to the right. In this gel system, the variant 1Bx7 subunit in D5 + Dy10-A* co-migrates with native 1Bx7.

the peptides detected by MS/MS predicted that many of the variant proteins containing them would be capable of forming disulfide bonds. To test whether the variants could form inter-molecular disulfide bonds, whole grain flours from homozygous progeny of lines D5-F*, D5 + Dy10-A* and D5 + Dy10-C were extracted with 50% 1-propanol. SDS-PAGE of the insoluble fractions revealed that both the 1Dx5 and 1Bx7 variants were found along with native HMW-GS in the propanol-insoluble fractions (Fig. 3), indicating that they did form inter-molecular disulfide bonds.

4. Discussion

We have shown that thirteen novel proteins of unexpected sizes in seven different transgenic lines are versions of HMW-GS (summarized in Table 2). These comprise 3 variants smaller and 10 larger than their related native HMW-GS. Most of the variants we characterized were derived from 1Dx5. One was derived from 1Dy10 and another appears to be hybrid between the N- and C-terminus of 1Dx5 and at least some of the repeat region of 1Dy10. The thirteenth is a variant of 1Bx7 that accumulated instead of the native version encoded at the *Glu-B1* locus.

Other researchers who have transformed wheat with HMW-GS genes have also noted exceptions to the expected expression patterns in transgenic wheat seeds (Alvarez et al., 2000; Barro et al., 1997; He et al., 2005; Vasil et al., 2001). Alvarez et al. (2000) were able to segregate two expression types from their event "F", as we did from our events D5-F and D5 + Dy10-A. In the former case, the two different types were a co-suppressed line expressing 1Ax1 and 1Dx5 transgenes and a non-suppressed line expressing only the 1Ax1 transgene (Alvarez et al., 2000). Vasil et al. (2001) found two of their nine 'Bobwhite' lines transformed with the gene encoding 1Ax1 each contained a novel protein larger than the endogenous HMW-GS. Preliminary investigation of this protein by N-terminal sequence analysis revealed similarity to an asparagine synthetase rather than to a HMW-GS.

Table 2

Summary of transgenic events and novel proteins analyzed.

Event name	HMW-GS intended for over-expression	Proteins larger than expected and analyzed	Proteins smaller than expected and analyzed
D5-M	1Dx5	2 Dx5-like	1 Dx5-like
D5-F*	1Dx5	3 Dx5-like	1 Dx5-like
D5 + Dy10-H	1Dx5, 1Dy10	1 Dx5/Dy10-like	1 Dx5/Dy10-like
D5 + Dy10-I	1Dx5, 1Dy10	1 Dx5-like	None
D5 + Dy10-C	1Dx5, 1Dy10	1 Dx5-like	None
D5 + Dy10-A*	1Dx5, 1Dy10	1 Bx7-like	None
Dy10 – B52a-6	1Dy10	1 Dy10-like	None

These results are not in agreement with our findings for the novel proteins in our transgenic lines, but the authors emphasized that their data were preliminary (Vasil et al., 2001). They also noted a variant protein in a third transgenic line that co-migrated with 1Ax2* in SDS-PAGE, but did not share its free zone capillary electrophoresis or RP-HPLC mobility (Vasil et al., 2001). Barro et al. (1997) observed novel protein bands of varying sizes in five of the nine transformants they analyzed for expression of 1Ax1 and 1Dx5 transgenes. In three of the lines, the novel proteins were recognized by a monoclonal antibody specific for the N-terminal domain of 1Ax1 and 1Dx5 subunits. The authors hypothesized that such variants were the products of rearranged HMW-GS genes (Barro et al., 1997).

Despite their large size and the relative scarcity of R and K in their repetitive regions, HMW-GS were readily identified by MS/MS. We were able to assign identities to even faint gel bands. Because of their close relationship, many subunits share some peptides and could not always be distinguished from one another. Some tryptic peptides, particularly those derived from the repetitive regions of the proteins, were not identified because they are smaller or larger than the range detected by MS/MS.

The finding that the variant HMW-GS can form inter-molecular disulfide bonds predicts that their presence will have some influence on the bread-making quality of flours that contain them. However, since they have always been found in transgenic lines that contain additional native subunits, it has not been possible to distinguish the effects of the variants on end-use properties from those due to increases in overall HMW-GS content.

The molecular mechanism for formation of these variants is unknown, but we speculate that the size differences arise mainly from changes in the lengths of the central repetitive coding regions. Such changes are difficult to assess by MS/MS because these regions are large and devoid of the protease cleavage sites commonly used to fragment proteins for MS. The central repetitive domains are known to be subject to length changes during evolution of native HMW-GS genes, perhaps by homologous recombination between misaligned repeated DNA segments (D'Ovidio et al., 1996; Shewry et al., 1989) or by slippage mistakes made by DNA polymerases during replication (Kroutil and Kunkel, 1999). Some of the same mechanisms may be operative during the introduction of DNA into the wheat embryo cells via biolistics. These hypothetical mechanisms predict that larger and/or more regular repeat regions would be more prone to such rearrangements than smaller or less regular repetitive regions. In support of

Fig. 2. Amino acid sequences encoded by HMW-GS genes for 1Dx5 (A) 1Bx7 (B) and 1Dy10 (C). Cysteines (C) are shadowed in gray. Tryptic peptides detected by tandem MS and used to identify variant proteins are underlined and numbered. All end in arginine (R) or lysine (K) (highlighted in reverse black and white), but those whose numbers are preceded by asterisks start after amino acids other than R or K (their starts are indicated by vertical bars). In some cases, smaller fragments of those peptides were also found (Frag *). Some of the commonly identified peptides were fusions of adjacent peptides, e.g., 14 + 15, 28 + 29. Peptide 3 was sometimes found fused to the preceding peptide ELK. Sometimes a version of peptide 24 with the Q after the R was detected. C. Peptides *37 and 38, which were found in the small variant of D5 + Dy10-H that may be a hybrid between 1Dx5 and 1Dy10 subunits, are double-underlined.

this prediction, He et al. (2005) found that 2 of 4 lines transformed with a version of the 1Dx5 gene with a 22.5% longer repeat region contained proteins smaller than would be encoded by the intact coding region in the transforming plasmid. For comparison, we found variants in 3 of 13 transformants bombarded with the natural 1Dx5 gene.

Of particular interest in the work reported here is the 1Bx7 variant that apparently replaced the native subunit in line Dx5 + Dy10-A*. The protein change found in this event suggests that some interaction may have occurred between incoming plasmid DNAs bearing the 1Dx5 or 1Dy10 genes and the native 1Bx7 gene at the *Glu-B1-1* locus. Alvarez et al. (2000) noted the disappearance of the endogenous 1Ax2* subunit in two of their six transformed lines. Interestingly, in their line called “D”, there was a novel protein they called “x” that migrated just below the 1Bx7 band in their gels. Both 1Ax2* and x were present in T₁ seed extracts, but in T₂ and T₃ transgenic segregants, 1Ax2* was missing. We speculate that their protein “x” could be a smaller version of 1Ax2*, generated by a mechanism similar to the one that resulted in a larger version of 1Bx7 in our line Dx5 + Dy10-A*, and replacing the native 1Ax2* gene on the 1A chromosome in some of their segregants.

Some clues to the possible molecular processes at work following biolistic transformation can be found in the structures of the transgene integration sites that result. Because of the complexities of both the insertions and the wheat genome, only a few such transgene integration sites have been analyzed (Abranches et al., 2000; Jackson et al., 2001; Rooke et al., 2003). These studies reveal that genes originally on separate plasmids are co-integrated into the same sites in multiple tandem and non-tandem copies, sometimes interspersed with genomic DNA. Rooke et al. (2003) characterized six transgenic wheat lines transformed by biolistics with HMW-GS and selection genes. They found that between one to about 15 copies were clustered in one to three loci, in some cases separated by genomic DNA. Some transgenes were present as tandem repeats and others were rearranged or truncated (Rooke et al., 2003). The interspersed genomic DNA in transgenes and their truncation and co-integration suggest that incoming DNA is broken and joined together by non-homologous end-joining during the biolistic process. The apparently chimeric Dx5/Dy10 variants found in event Dx5 + Dy10-H could have been formed by such a process. The results of the integration site analyses and the MS/MS data presented here indicate that the incoming plasmid DNA's interact with one another and with genomic DNA before and/or during integration, but the relative roles played by homology-based and non-homologous processes in these interactions is not known.

Regardless of the molecular mechanism, these alterations of HMW-GS genes are quite common during biolistic transformation. As detailed above, they have been documented in several independent laboratories. Three of the events we report here had more than one variant. The proteins we analyzed were clearly visible as individual protein bands in un-crowded regions of 1-dimensional SDS-PAGE gels in seven of 30 transgenic lines. It is possible that other variants could be present in more crowded regions of the gels where they are not readily distinguished from endogenous proteins. Thus, we are probably underestimating the frequency with which such size variants are generated by biolistic transformation. Better separation methods such as RP-HPLC and 2-D electrophoresis might reveal more such variants in our transformants. It is possible that the HMW-GS transgenes are especially prone to such expansion and contraction because there are six closely related genes in each bread wheat cultivar and because they contain long regions of repetitive coding sequence. Whether or not such alterations are common in the coding

regions of other genes used in biolistic transformations remains to be investigated.

5. Conclusions

Biolistic transformation of wheat with genes encoding HMW-GS often results in changes in their coding regions that lead to production of related proteins larger or smaller in size than the native subunits. In one case, transformation resulted in apparent replacement of the native *Glu-B1-1* gene with a version that encoded a larger 1Bx7 subunit. Such variants are detectable by one-dimensional SDS-PAGE in about one-fifth of lines transformed with native genes encoding 1Dx5 and/or 1Dy10. All the variant proteins were HMW-GS and 6 of 6 tested were found in the polymeric protein fraction. Up to four altered HMW-GS were found in the same transgenic event. The mechanism by which these variants arise requires further investigation.

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