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RESEARCH ARTICLE



Comparison of low molecular weight glutenin subunits identified by SDS-PAGE, 2-DE, MALDI-TOF-MS and PCR in common wheat

Li Liu¹, Tatsuya M Ikeda², Gerard Branlard³, Roberto J Peña⁴, William J Rogers⁵, Silvia E Lerner⁶, María A Kolman⁵, Xianchun Xia¹, Linhai Wang¹, Wujun Ma⁷, Rudi Appels⁷, Hisashi Yoshida⁸, Aili Wang⁹, Yueming Yan⁹ and Zhonghu He*1,10

Abstract

Background: Low-molecular-weight glutenin subunits (LMW-GS) play a crucial role in determining end-use guality of common wheat by influencing the viscoelastic properties of dough. Four different methods - sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), two-dimensional gel electrophoresis (2-DE, IEF × SDS-PAGE), matrixassisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) and polymerase chain reaction (PCR), were used to characterize the LMW-GS composition in 103 cultivars from 12 countries.

Results: At the Glu-A3 locus, all seven alleles could be reliably identified by 2-DE and PCR. However, the alleles Glu-A3e and *Glu-A3d* could not be routinely distinguished from *Glu-A3f* and *Glu-A3g*, respectively, based on SDS-PAGE, and the allele Glu-A3a could not be differentiated from Glu-A3c by MALDI-TOF-MS. At the Glu-B3 locus, alleles Glu-B3a, Glu-B3b, Glu-B3c, Glu-B3q, Glu-B3h and Glu-B3j could be clearly identified by all four methods, whereas Glu-B3ab, Glu-B3ac, Glu-B3ad could only be identified by the 2-DE method. At the Glu-D3 locus, allelic identification was problematic for the electrophoresis based methods and PCR. MALDI-TOF-MS has the potential to reliably identify the Glu-D3 alleles.

Conclusions: PCR is the simplest, most accurate, lowest cost, and therefore recommended method for identification of Glu-A3 and Glu-B3 alleles in breeding programs. A combination of methods was required to identify certain alleles, and would be especially useful when characterizing new alleles. A standard set of 30 cultivars for use in future studies was chosen to represent all LMW-GS allelic variants in the collection. Among them, Chinese Spring, Opata 85, Seri 82 and Pavon 76 were recommended as a core set for use in SDS-PAGE gels. *Glu-D3c* and *Glu-D3e* are the same allele. Two new alleles, namely, Glu-D3m in cultivar Darius, and Glu-D3n in Fengmai 27, were identified by 2-DE. Utilization of the suggested standard cultivar set, seed of which is available from the CIMMYT and INRA Clermont-Ferrand germplasm collections, should also promote information sharing in the identification of individual LMW-GS and thus provide useful information for quality improvement in common wheat.

Background

Glutenin proteins are the major factors responsible for the unique viscoelastic characteristics of wheat dough. They determine rheological properties and bread-making performance [1-3]. The polymeric glutenin proteins, with molecular weights ranging from less than 300 to more

than 1,000 kDa, are composed of two groups of subunits. These subunits include the LMW-GS, which are similar in size and structure to the γ - gliadins (30-40 kDa), and the high-molecular-weight glutenin subunits (HMW-GS) which range in molecular mass from ~65 to 90 kDa [4]. The LMW-GS represent about one-third of the total seed protein and ~60% of total glutenins [5], and are essential in determining dough properties, such as dough extensibility [6] and gluten strength [2]. Hence characterization of allelic variation among cultivars and investigation of their relationships with end-use quality has been a key



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area of research on quality improvement during the last 15 years, and is the basis for the success of using specific LMW-GS alleles in breeding programs [7-9].

The genes coding for LMW-GS are located on the short arms of homoeologous group 1 chromosomes at the Glu-A3, Glu-B3 and Glu-D3 loci, and are tightly linked to the *Gli-1* loci [10-12]. The *Glu-A3* locus on chromosome 1A encodes relatively few LMW-GS, with alleles Glu-A3e in hexaploid or common wheat and Glu-A3h in tetraploid wheat being null alleles that do not express any Glu-A3 product [13,14]. In contrast, there is extensive variation for LMW-GS encoded by chromosome 1B in common wheat. The Glu-D3 locus has less variability with five alleles reported originally by Gupta and Shepherd [13], four alleles by Lerner et al. [15] and only three alleles observed by Jackson et al. [16] and Eagles et al. [17]. Nonetheless, recent studies using protein and PCR analyses have identified 11 Glu-D3 alleles [18,19], suggesting that a reexamination should be carried out to clarify the genetic variability at this locus.

Despite the abundance of the LMW-GS, they have received much less attention than the HMW-GS, probably due to their complexity, heterogeneity and co-migration with gliadins in SDS-PAGE [19,20]. In the SDS-PAGE system, utilizing gliadins as indicators provided an indirect way to define LMW-GS alleles [16]. The 2-DE analytical process that could generate much more information than SDS-PAGE [21] was not generally recommended for use in breeding programs, due to its timeconsuming procedure, high costs and skill requirements. MALDI-TOF-MS is currently the most efficient method to analyze proteins and requires only 4-5 minutes per sample. It is a high throughput technology for analyzing wheat gluten proteins [22-25], but being relatively new and expensive, few wheat breeding programs can afford to acquire such equipment. Recently, a simple, rapid and sensitive PCR approach, has proven to be a very useful tool for identifying LMW-GS composition in common wheat [19,26-28].

LMW-GS were first identified by gel filtration and starch gel electrophoresis of extracts of wheat flour [29,30]. They were classically subdivided into B, C, and D groups (no relationship to the A, B and D genomes of wheat), according to their electrophoretic mobilities in SDS-PAGE and their isoelectric points (pI) [31]. Based on the locations of cysteine residues involved in the formation of intermolecular disulfide bridges, Ikeda et al. [32] classified LMW gene sequences into six types, each containing several different groups based upon differences in their N- and C-terminal acid-amino compositions. Altogether, 12 groups were differentiated, but an additional five groups were reported by Juhász and Gianibelli [33].

The allelic nomenclature system for the LMW-GS was defined through the chromosomal location of the DNA

coding regions by Gupta and Shepherd [13] and was reviewed by Jackson et al. [16]. Branlard et al. [34] proposed a schematic presentation of SDS-PAGE relative subunit mobilities to characterize the different alleles encoded at *Glu-A3*, *Glu-B3* and *Glu-D3* loci. Ikeda et al. [35] recently compared *Glu-3* allele identifications from five laboratories, confirming inconsistencies between laboratories in identifying *Glu-3* alleles due to differences between the separation and identification methods. The study also indicated new *Glu-3* alleles in a number of the cultivars analyzed.

The N-terminal sequences of LMW-GS were used to divide the protein subunits into two main groups [32,36]. The first group corresponded to typical LMW-GS, i.e., LMW-i (or i-type, first amino acid isoleucine) and LMWm (or m-type, methionine) types, and the second group, named gliadin-like sequences [37] as these subunits have N-terminal sequences similar to α -, γ - and ω - gliadins. Most gliadins are monomeric, but some have an extra cys that allows them to be incorporated into glutenin polymers. Payne [1] termed the prominent bands observed by SDS-PAGE of reduced glutenin protein as A (HMW-GS), B (many of the LMW-GS) and C (the smaller LMW-GS). Later, other researchers also observed larger gliadin-like subunits, between the A and B bands, and they named them as D- subunits [31]. Most of the B- subunits were shown to possess i-, m- or s (serine) -type N-terminal sequences [38]. C- subunits including α -, and γ - gliadinslike subunits as well as subunits with classic LMW-GS sequences occur in large numbers, although their relative amounts are lower than those of B- subunits. Similarly, D- subunits have N-terminal sequences that correspond to ω - gliadins, another type of gliadin-like sequence [2,39,40].

The use of two distinct nomenclature systems, one based upon the relative mobilities in SDS-PAGE and the other upon N-terminal sequences, make it extremely difficult to compare work from different laboratories. The main ambiguities from these different classification systems can be summarized as follows: 1) at the Glu-A3 locus, both Glu-A3a and Glu-A3c were reported for the same cultivar, and similarly, Glu-A3a, Glu-A3b, Glu-A3c, Glu-A3d were reported to be identical to Glu-A3e; 2) at the Glu-B3 locus, results differed for Glu-B3b and Glu-*B3g*, and for *Glu-B3f* and *Glu-B3g* in the same cultivars; and 3) at the *Glu-D3* locus, there was ambiguity between Glu-D3a and Glu-D3c, and between Glu-D3a and Glu-D3b in the same cultivars [41]. As a consequence of these problems, reports of correlations between certain allelic forms of LMW-GS and quality parameters in common wheat have often been contradictory [7,42-45]. It is, therefore, essential to establish a simple and uniform classification through a set of standard cultivars for each LMW-GS allele.

In 2005, a cooperative program was developed among the following five laboratories to establish such a set of standard cultivars for identifying LMW-GS alleles: Chinese Academy of Agricultural Sciences (CAAS, China), International Maize and Wheat Improvement Center (CIMMYT, Mexico), National Institute for Agricultural Research (INRA, France), National Agriculture and Food Research Organization (NARO, Japan), and National University of the Center of the Province of Buenos Aires (Universidad Nacional, Argentina). A set of 103 cultivars used in various previously studies [35] in 12 countries was assembled and distributed to all laboratories, including Murdoch University as an additional laboratory, for the identification of LMW-GS alleles. Their preliminary *Glu-3* allelic assignments were summarized in a previous paper [35]. The objectives of the current paper are 1) to compare the LMW-GS compositions obtained by SDS-PAGE, 2-DE, MALDI-TOF-MS and PCR in order to clearly identify the protein compositions of cultivars in the collection; and 2) to establish a set of standard cultivars for the identification of LMW-GS alleles, enabling information regarding the effects of individual LMW-GS on gluten properties to be readily and continuously shared between laboratories and applied in breeding programs.

Results and discussion Analysis of LMW-GS by SDS-PAGE

The LMW-GS compositions identified in participating laboratories by SDS-PAGE were combined and listed in Table 1 (details available upon request); discrepancies among different laboratories were discussed by Ikeda et al. [35]. At the *Glu-A3* locus, alleles *Glu-A3a*, *Glu-A3b*, *Glu-A3c* and *Glu-A3f* could be readily identified using SDS-PAGE (Figure 1). Alleles *Glu-A3d* and *Glu-A3g* could be differentiated with the aid of the gliadin SDS-PAGE gel; by the presence or absence of the *Gli-A1o* allele, which we believe is linked to *Glu-A3d*, but not to *Glu-A3g* (Figure 2). It was difficult to distinguish *Glu-A3f* from *Glu-A3e* (null allele). In previous studies [7,46,47] both alleles tended to be detected as *Glu-A3e*.

Figure 3 shows cultivars representing different Glu-B3 alleles. At the Glu-B3 locus, three alleles, Glu-B3d, Glu-B3h and Glu-B3i, each carried the slowest LMW-GS bands in the SDS-PAGE region B among the cultivars studied. The slowest Glu-B3 band, Glu-B3b, almost coincided with Glu-A3a, but the Glu-B3b band was usually lighter and thinner, permitting their discrimination. Allele Glu-B3f could not be reliably discriminated from Glu-B3g since these bands had very similar mobilities, including the presence of a band in the SDS-PAGE region (Figure 3, lanes 8-10) as previously reported [7,34,41]. However, taking advantage of the Glu-B3/Gli-B1 linkage, one can look at the omega-gliadins region in SDS-PAGE, to identify with confidence several of the Glu-B3 alleles (Figure 4). Actually, differentiating between several Glu-B3 alleles is possible only looking at both, gliadin and glutenin SDS-PAGE gels. Using this criteria, Glu-B3 alleles in lanes 15, 16, 17, 18, and 19, seem to correspond to Glu-B3b, Glu-B3g, Glu-B3g, Glu-B3i, and Glu-B3i, respectively (Figures 3 and 4), however, 2-DE analysis indicates that these genotypes correspond to new alleles provisionally designated as Glu-B3ab, Glu-B3ac, Glu-B3ac, Glu-B3ad, and Glu-B3ad, respectively (Figure 3).

Figure 5 shows cultivars representing different Glu-D3 alleles. Although alleles Glu-D3a, Glu-D3b, Glu-D3c and Glu-D3d were frequently identified in germplasm from various origins [35], only alleles Glu-D3a, Glu-D3b and Glu-D3d were consistently differentiated [34]. Glu-D3 alleles had similar mobilities to gliadins and were generally faintly stained due to the rapid diffusion of low molecular mass proteins from the gel. Thus the identification of Glu-D3 alleles was quite difficult using only SDS-PAGE, leading to the reported discrepancies [13,19,41]. Although improvements to the SDS-PAGE protocol now allow differentiating several of the Glu-D3 alleles with more certainty, as it is shown in Figure 5, other methods for definitive identification of these alleles, such as 2-DE, MALDI-TOF-MS and PCR, had to be implemented to facilitate identification of *Glu-D3* alleles.

Characterization of LMW-GS by 2-DE

The identification of the LMW-GS alleles by 2-DE was consistent between the two laboratories. The discrimination between LMW-GS alleles in the collection by high resolution 2-DE is illustrated in Figures 6, 7, 8 and 9 and the results are shown in Table 1. The *Glu-A3* alleles *Glu-A3d* (Figure 6, (4)), *Glu-A3e* (Figure 6, (5)), *Glu-A3f* (Figure 6, (6)) and *Glu-A3g* (Figure 7, (1)), were readily differentiated on the basis of protein spots with clearly different molecular masses and pI. Alleles *Glu-A3a* (Figure 6, (1)), *Glu-A3b* (Figure 6, (2)) and *Glu-A3c* (Figure 6, (3)) had identical pI but different molecular masses, making it possible to discriminate between them.

At the *Glu-B3*, the alleles *Glu-B3ab* (Figure 7, (4)), *Glu-B3ac* (Figure 8, (2)), *Glu-B3h* (Figure 8, (3)), *Glu-B3ad* (Figure 8, (4)) and *Glu-B3j* (Figure 8, (5)) were easily differentiated by protein spots having different molecular masses and pI. Alleles *Glu-B3ab* (Figure 7, (4)), *Glu-B3ac* (Figure 8, (2)) and *Glu-B3ad* (Figure 8, (4)) were each discriminated from *Glu-B3b* (Figure 7, (3)), *Glu-B3g* (Figure 8, (1)) and *Glu-B3i* (image not provided) by two distinct protein spots. Although the majority of the protein spots for alleles *Glu-B3b* and *Glu-B3g* had identical molecular masses and pI, they could be discriminated since allele *Glu-B3g* had one additional spot, at pH6, located between the HMW-GS and gliadins. There were no obvious differences in molecular mass or pI between alleles *Glu-B3d*

Cultivar	Origin	Glu-A3	Glu-B3	Glu-D3
Aca 303	Argentina	f/f/f*	h/h/h/h	c/c/c/-
Aca 601	Argentina	f/f/f	b/b/b/b	c/c/c/-
Aca 801	Argentina	c/c/a or c/c	g/ac/g/g	b/b/-
Buck Brasil	Argentina	f/f/f	g/ac/g/g	d/d/?/-
Buck Mejorpán	Argentina	f/f/f	b/b/b/b	c/c/c/-
Buck Pingo	Argentina	f/f/f	i/ad/d or i/i	c/c/c/-
Klein Capricornio	Argentina	c/c/a or c/c	h/h/h	b/b/-
Klein Chaja	Argentina	c/c/a or c/c	h/h/h	b/b/-
Klein Flecha	Argentina	c/c/a or c/c	h/h/h	b/b/-
Klein Jabal 1	Argentina	d/d/g	g/g/g/g	c/c/-
Klein Martillo	Argentina	e/e/e	j/j/j/j	b/b/-
Klein Proteo	Argentina	g/g/e/g	g/ac/g/g	b/b/-
Nidera Baguette 10	Argentina	d/d/d	g/g/g/g	c/c/c/-
Nidera Baguette 20	Argentina	f/f/f/f	g/g/g/g	c/c/c/-
ProINTA Amanecer	Argentina	f/f/f	j/j/j	a/a/b/-
ProINTA Colibr 1	Argentina	d/d/d	b/b/b/b	a/a/a/-
ProINTA Isla Verde	Argentina	b/b/b/b	b/b/b/b	b/b/-
ProINTA Redomon	Argentina	c/c/a or c/c	h/h/h	c/c/c/-
Thomas Nevado	Argentina	c/c/a or c/c	j/j/j/j	b/b/-
Angas	Australia	c/c/a or c/c	g/g/g/g	c/c/-
Avocet	Australia	c/c/a or c/c	b/b/b/b	b/b/-
Carnamah	Australia	c/c/a or c/c	i/ad/d or i/i	c/c/-
Gabo	Australia	b/b/b/b	b/b/b/b	b/b/-
Grebe	Australia	c/c/a or c/c	j/j/j/j	b/b/-
Halberd	Australia	e/e/e	c/c/c/c	c/c/-
Insignia	Australia	f/f/f/f	c/c/c/c	c/c/-
Millewa	Australia	c/c/a or c/c	g/g/g/g	b/b/-
Spear	Australia	e/e/e/e	h/h/h	c/c/-
Stiletto	Australia	c/c/a or c/c	h/h/h	c/c/c/-
Tasman	Australia	b/b/b/b	i/ad/d or i/i	a/a/a/-
Trident	Australia	e/e/e	h/h/h	c/c/c/-
Westonia	Australia	c/c/a or c/c	h/h/h	c/c/c/-
Wilgoyne	Australia	d/d/d	h/h/h	b/b/-
AC Vista	Canada	e/e/e	i/ad/d or i/i	c/c/c/-
Bluesky	Canada	g/g/e/g	g/g/g/g	c/c/c/-
Glenlea	Canada	g/g/e/g	g/g/g/g	c/c/-
Katepwa	Canada	e/e/e	h/h/h/h	c/c/c/-
Marquis	Canada	e/e/e	b/b/b/b	a/a/a/-
Neepawa	Canada	e/e/e	h/h/h	c/c/-
Pioneer	Canada	e/e/e	i/ad/d or i/i	c/c/-
99G46	China	f/f/f/f	j/j/j/j	c/c/-
CA9641	China	d/d/d/d	h/h/h	c/c/-
CA9722	China	c/c/a or c/c	h/h/h/h	c/l/c/-

Table 1: Compositions of LMW-GS alleles in 103 wheat cultivars identified by SDS-PAGE, 2-DE, MALDI-TOF-MS and allele-specific markers

Chinese Spring	China	a/a/a or c/a	a/a/a	a/a/a/-	
Demai 3	China	c/c/a or c/c	i/d or i/d or i/i	b/b/b/-	
Fengmai 27	China	c/c/a or c/c	f/f/f/f	a/n/a/-	
Guanfeng 2	China	c/c/a or c/c	b/b/b/b	a/a/a/-	
Huaimai 16	China	f/f/f/f	h/h/h	c/c/c/-	
Jing 411	China	c/c/a or c/c	h/h/h	c/l/c/-	
Lumai 23	China	c/c/a or c/c	d/d or i/d or i/d	c/l/c/-	
Neixiang 188	China	a/a/a or c/a	j/j/j	a/a/a/-	
Shan 229	China	c/c/a or c/c	j/j/jj	b/b/b/-	
Wanmai 33	China	d/d/d/d	g/g/g/g	a/a/b/-	
Yan 239	China	c/c/a or c/c	j/j/jj	b/b/b/-	
Yangmai 158	China	c/c/a or c/c	g/g/g/g	c/c/c/-	
Yumai 54	China	c/c/a or c/c	d/d or i/d or i/d	c/c/c/-	
Yumai 63	China	c/c/a or c/c	d/d or i/d or i/d	c/c/c/-	
Yumai 69	China	c/c/a or c/c	d/d or i/d or i/d	a/a/b/-	
Zhongyou 9507	China	d/d/d/d	b/b/b/b	c/c/c/-	
Zhongyou 9701	China	d/d/d/d	d/d or i/d or i/d	c/c/c/-	
Zhongyu 415	China	c/c/a or c/c	d/d or i/d/d	c/c/c/-	
Ruso	Finland	c/c/a or c/c	i/ad/d or i/i	a/a/a/-	
Brimstone	France	c/c/a or c/c	g/g/g/g	d/d/?/-	
Cappelle-Desprez	France	d/d/d/d	g/g/g/g	c/c/c/-	
Chopin	France	c/c/a or c/c	h/h/h	c/c/c/-	
Clément	France	f/f/f/f	j/j/j/j	c/c/c/-	
Courtot	France	c/c/a or c/c	b/b/b/b	c/l/c/-	
Darius	France	d/d/d/d	g/g/g/g	b/m/m/-	
Etoile de Choisy	France	d/d/d/d	i/d or i/d or i/i	c/l/c/-	
Festin	France	f/f/f/f	b/b/b/b	c/l/c/-	
Magali Blondeau	France	e/e/e/e	g/g/g/f	b/b/b/-	
Magdalena	France	d/d/d/d	b/b/b/b	a/a/a/-	
Petrel	France	d/d/d/d	h/h/h	c/c/c/-	
Renan	France	f/f/f/f	b/b/b/b	b/b/b/-	
Soissons	France	c/c/a or c/c	b/b/b/b	c/c/c/-	
Thesee	France	c/c/a or c/c	g/ac/g/g	c/l/c/-	
Apollo	Germany	d/d/d/d	j/j/j/j	c/c/c/-	
Manital	Italy	c/c/a or c/c	b/b/b/b	a/a/a/-	
Salmone	Italy	c/c/a or c/c	c/c/c/g	c/c/c/-	
Aoba-komugi	Japan	e/e/e/e	b/b/b/b	c/c/c/-	
Eshimashinriki	Japan	c/c/a or c/c	d/d or i/d or i/d	a/a/a/-	
Haruyutaka	Japan	c/c/a or c/c	h/h/h	b/b/b/-	
Kanto 107	Japan	c/c/a or c/c	g/g/g/g	a/a/a/-	
Kitanokaori	Japan	f/f/f/f	j/j/j/j	c/c/c/-	
Nanbu-komugi	Japan	d/d/d/d	b/ab/b/b	a/a/a/-	
Norin 61	Japan	d/d/d/d	i/d or i/d or i/-	c/c/c/-	
Norin 67	Japan	c/c/a or c/c	g/g/g/g	b/b/b/-	
Shinchunaga	Japan	c/c/a or c/c	i/ad/d or i/-	a/a/a/-	
Shirane-komugi	Japan	e/e/e/e	i/ad/d or i/i	a/a/a/-	

Table 1: Compositions of LMW-GS alleles in 103 wheat cultivars identified by SDS-PAGE, 2-DE, MALDI-TOF-MS and allele-specific markers (Continued)

Amadina	Mexico	e/e/e/e	j/j/j/j	c/l/c/-
Attila	Mexico	c/c/a or c/c	h/h/h/h	b/b/b/-
Heilo	Mexico	f/f/f/f	i/ad/d or i/i	c/l/c/-
Opata 85	Mexico	b/b/b/b	i/ad/d or i/i	a/a/a/-
Pastor	Mexico	c/c/a or c/c	g/g/g/g	b/b/b/-
Pavon 76	Mexico	b/b/b/b	h/h/h/h	b/b/b/-
Pitic	Mexico	c/c/a or c/c	b/b/b/b	b/b/b/-
Rebeca	Mexico	c/c/a or c/c	g/g/g/g	b/b/b/-
Seri 82	Mexico	c/c/a or c/c	j/j/j/j	b/b/b/-
Orca	Netherlands	d/d/d	d/d or i/d or i/d	c/c/c/-
Pepital	Netherlands	f/f/f/f	d/d or i/d or i/d	c/l/c/-
Ernest	USA	d/d/d	d/d or i/d or i/d	d/?/?-
Splendor	USA	e/e/e/e	g/g/g/g	b/b/b/-
Verde	USA	f/f/f/f	h/h/h/h	c/c/c/-

Table 1: Compositions of LMW-GS alleles in 103 wheat cultivars identified by SDS-PAGE, 2-DE, MALDI-TOF-MS and allelespecific markers (Continued)

*, the first, second, third and fourth symbol in each column are alleles of *Glu-3* loci identified by SDS-PAGE, 2-DE, MALDI-TOF-MS and PCR, respectively.

Data from five laboratories are combined data for SDS-PAGE. - indicates data not available

(Figure 7, (6)) and *Glu-B3i* (image not provided), or between *Glu-B3f* (image not provided) and *Glu-B3g* (Figure 8, (1)), making differentiation by 2-DE impossible.

At the *Glu-D3*, only *Glu-D3c* (Figure 9, (2)), *Glu-D3l* (Figure 9, (3)) and *Glu-D3m* (Figure 9, (4)) could be definitely identified by 2-DE. Allele *Glu-D3l* (Figure 9, (3)) had two more distinctive spots compared to *Glu-D3c* (Figure 9, (2)) in 2-DE separations. As expected, alleles *Glu-D3c* and *Glu-D3e* (image not provided) could not be separated by 2-DE. These alleles appeared to be the same based on SDS-PAGE and MALDI-TOF-MS in the present study as they were in a previous study [16].

2-DE did not distinguish *Glu-D3a* (Figure 8, (6)), *Glu-D3b* (Figure 9, (1)) and *Glu-D3d* (image not provided), hence further investigation should target discrimination of *Glu-D3* alleles by combining 2-DE with other methods such as PCR.

Identification of LMW-GS by MALDI-TOF-MS

The compositions of LMW-GS analyzed by MALDI-TOF-MS are presented in Table 1. As shown in Figures 10, 11, 12 and 13, the spectra of LMW subunits analyzed by this method consist of complex sets of peaks, consistent with the extensive diversity of the subunits. The LMW-GS exhibited molecular masses of 25-43 kDa in MALDI-TOF-MS spectra, considerably lower than the corresponding molecular masses of 42-51 kDa determined by SDS-PAGE and indicative of limitations of the SDS-PAGE method in determining the molecular masses of LMW glutenins [24]. Two major regions with masses from 30 to 35 kDa and from 36 to 43 kDa were separated in spectra of MALDI-TOF-MS (Figures 10, 11, 12 and 13). These regions correspond to the C- LMW-GS and B-LMW-GS classified by SDS-PAGE. The region with molecular masses of 30-35 kDa also corresponds in mass to the major gliadins range [1]. The results were in agreement with previous studies based on SDS-PAGE, where there were extensive overlaps between gliadins and LMW-GS with lower molecular masses [48].

MALDI-TOF-MS-based identification of LMW-GS alleles was established using a set of 19 near-isogenic lines (NIL) of cultivar Aroona (unpublished data, A Wang, W Ma, R Appels, Murdoch University, Australia). Most of the distinct peaks of the *Glu-A3* alleles exhibited higher masses in the ranges of about 41.8-42.1 kDa and 43.5-43.8 kD, whereas the distinct peaks of the *Glu-D3* alleles showed lower masses of 33.2-33.7 kDa. The middle masses in the ranges of about 40.1-40.2 kDa and 42.8-43.3 kDa corresponded to the *Glu-B3* alleles. The distributions of distinct peaks of the *Glu-3* alleles in the MALDI-TOF-MS were in agreement with their position in SDS-PAGE [34].

Compared to the other loci, *Glu-A3* was less diverse and most protein bands had lower mobilities, so discrimination between them using SDS-PAGE is usually feasible. Similarly, most of the distinct peaks of the *Glu-A3* alleles were well separated in MALDI-TOF-MS spectra, and alleles *Glu-A3b* (Figure 10, (2)), *Glu-A3d* (Figure 10, (3)), *Glu-A3e* (Figure 10, (4)) and *Glu-A3f* (Figure 11, (1)) were reliably discriminated.

The *Glu-B3* alleles *Glu-B3a* (Figure 11, (2)), *Glu-B3b* (Figure 11, (3)), *Glu-B3c* (Figure 11, (4)), *Glu-B3h* (Figure

12, (3)), and *Glu-B3j* (Figure 12, (4)), as well as seven other alleles, were readily distinguished by MALDI-TOF-MS.

With regard the *Glu-D3* locus, MALDI-TOF-MS clearly differentiated the *Glu-D3a* (Figure 13, (1)), *Glu-D3b* (Figure 13, (2)), *Glu-D3c* (Figure 13, (3)) and *Glu-D3m* (Figure 13, (4)) alleles. As expected, *Glu-D3e* (image not provided) could not be discriminated from *Glu-D3c* (Figure 13, (3)). Improved discrimination will be achieved as calibration technology improves. In addition, it may be of value to utilize the close linkage between gliadin and LMW glutenin alleles to further improve the power of MALDI-TOF-MS in differentiating LMW glutenin alleles.

Detection of LMW-GS by allele specific PCR markers

Seven primer pairs [27], including *gluA3a*, *gluA3b*, *gluA3ac*, *gluA3d*, *gluA3e*, *gluA3f* and *gluA3g*, were used to identify *Glu-A3* alleles (Table 1). The amplified fragment sizes for each marker were 529 bp for *Glu-A3a*, 894 bp for *Glu-A3b*, 967 bp for *Glu-A3d*, 158 bp for *Glu-A3e*, 552 bp for *Glu-A3f*, and 1345 bp for *Glu-A3g*, indicating that the *Glu-A3* alleles in the collection could be readily distinguished from one another. Since no *Glu-A3c* allele-specific primer has been developed, identification of this allele required the use of the *gluA3ac* with a 573 bp band in combination with the marker *gluA3a* [27].

Ten primer pairs developed by Wang et al. [28] were utilized to test for *Glu-B3* alleles and the results are sum-



Figure 2 SDS-PAGE of gliadins. The gliadins are 50% propanol (v/v) soluble fractions (The same as below). Cultivars in lanes 7, 8, 13, and 14, correspond to the same shown with same number in Figure 1. The linkage between *Gli-A1o* (indicated in lanes 7 and 8 in the omega-gliadin zone) and *Glu-A3d* helps to differentiate the latter from *Glu-A3g*.

a

a

Glu-A3

d

marized in Table 1. Specifically amplified fragments included 1095 bp for *Glu-B3a*, 1549 bp for *Glu-B3b*, 472 bp for *Glu-B3c*, 662 bp for *Glu-B3d*, 669 bp for *Glu-B3e*, 853 bp for *Glu-B3g*, 1022 bp for *Glu-B3h*, and 621 bp for *Glu-B3i*, indicating that the *Glu-B3* alleles could be well differentiated based on corresponding markers. Detection of *Glu-B3f* required the use of the *Glu-B3fg* marker with an 812-bp marker in combination with the *Glu-B3g* marker since no *Glu-B3f* allele-specific marker has been designed. Although *Glu-B3f* could not be clearly distinguished from *Glu-B3g* by protein based methods, these alleles could be definitively differentiated by PCR. In addition, there were obvious differences between genes *GluB3-1* and *GluB3-2* in the gene sequences of *Glu-B3f*

and *Glu-B3g* [28]. The differences were firstly, the sequence length of *Glu-B3f* was 60 bp longer than that of *Glu-B3g* in the *GluB3-1* gene, and secondly, there were single base differences between *Glu-B3f* and *Glu-B3g* in both *GluB3-1* and *GluB3-2*. Therefore, alleles *Glu-B3f* and *Glu-B3g* reported in previous studies were different alleles although they could not be reliably differentiated by SDS-PAGE, 2-DE or MALDI-TOF-MS [13,34].

Glu-D3 appeared to be the most complicated locus. It contains the highest number of genes and expressed subunits compared to the other two loci, and yet most of the subunits across different alleles have similar molecular weights. Electrophoresis based methods and PCR are not efficient in differentiating *Glu-D3* alleles. The MALDI-TOF-MS based method can differentiate *Glu-D3* alleles since it is able to differentiate subtle changes in mass values. High accuracy mass calibration to remove the variations in mass measurement is the key to improve the efficiency of MALDI-TOF in differentiating these alleles.

Comparison of the four methods for identification of LMW-GS composition

The data from all five laboratories and the four methods employed showed that alleles Glu-A3b, Glu-A3d and Glu-A3e were consistently identified by all four methods. Similarly, analyses of alleles Glu-B3a, Glu-B3b, Glu-B3c, Glu-B3h and Glu-B3j were in agreement for all four methods. At the Glu-D3, only the Glu-D3c allele was consistently identified by SDS-PAGE, 2-DE and MALDI-TOF-MS. The discrepancies in allelic identification using the different methods are indicated in Table 2. Alleles Glu-A3a and *Glu-A3c* could not be distinguished by MALDI-TOF-MS due to their nearly identical molecular masses. Similarly, these two alleles could not be reliably identified by SDS-PAGE and 2-DE due to their identical mobilities and pI. However, it was easy to differentiate them by PCR. In SDS-PAGE gels, the higher mobility patterns of alleles Glu-B3d, Glu-B3h, Glu-B3i overlapped with those of alleles Glu-A3a or Glu-A3c, and lower mobility patterns overlapped with those of allele Glu-A3b. These results were in agreement with the reports of Gupta and Shepherd [13], who concluded that ambiguous identification of subunits was possibly caused by differential staining intensity of banding patterns. The difficulty to differentiate Glu-B3b and Glu-B3g based on SDS-PAGE banding patterns arose from their similar mobilities. However, as shown in Figures 3 and 4, several *Glu-B3* alleles could be readily discriminated using gliadins as a marker for glutenin by SDS-PAGE. These alleles had clearly different peaks or spots using MALDI-TOF-MS or 2-DE, respectively. Alleles Glu-D3a and Glu-D3b could not be reliably separated by MALDI-TOF-MS or 2-DE. It is suggested that the Glu-D3 alleles should be differentiated by a combination of primers [49-51].



The 2-DE method is generally considered as the most powerful tool for identifying storage protein polymorphism of proteins in wheat [52]. However, different bands in SDS-PAGE separations were not always distinguishable in 2-DE separations. For example, alleles *Glu-B3d* and *Glu-B3i* could be identified by SDS-PAGE, but not by 2-DE. For LMW-GS identification in wheat breeding programs, PCR and/or SDS-PAGE of both gliadin and glutenin extracts should be used as the basic method, with 2-DE and MALDI-TOF-MS as complementary approaches. A combination of different methods is recommended for differentiating certain LMW-GS alleles, particularly those suspected as being novel.

Comparison of the four methods is presented in Table 3. Utilization of a particular method will depend upon research objectives and the targeted materials. With appropriate classification of glutenin alleles, it is possible to improve wheat quality by selection of alleles and allelic combinations with desired quality performance. If progeny screening and cultivar development is the objective, PCR will likely be adequate for the identification of *Glu-A3* and *Glu-B3* alleles. However, if the aim is to determine the glutenin subunits of potential parents for predicting cross performance and designing crossing schemes, or to identify specific alleles such as *Glu-A3g*, *Glu-B3ab*, *Glu-B3ac*, or distinguish between the *Glu-D3* alleles, a combination of methods should be used, i.e. PCR with 2-DE or PCR with SDS-PAGE and 2-DE, in order to achieve the correct identification of LMW-GS alleles.

A set of standard cultivars for identification of LMW-GS

From this study of 103 wheat cultivars from 12 countries we propose a set of 30 cultivars for determination of LMW-GS (Table 4) irrespective of the method to be used. Figures 1, 3, 5 show glutenin electropherograms of 28 (missing Ernest and Darius) of the 30 genotypes presented in Table 4. They cover all LMW-GS allelic variants identified in the original set. A core set of Chinese Spring, Opata 85, Seri 82 and Pavon 76 is recommended for inclusion in all gels. Most of the common *Glu*-3 alleles





Figure 5 SDS-PAGE of LMW-GS. Cultivars: 1. Chinese Spring, 2. Neixiang 188, 3. Gabo, 4. Avocet, 5. Insignia, 6. Cappelle- Desprez, 7. Amadina, 8. Heilo, 9. Fengmai 27. Arrow heads indicate bands corresponding to different *Glu-D3* alleles.

are represented among this group and their distributions on gels will provide useful landmarks for comparison with other bands. In this classification, it is possible to differentiate alleles *Glu-A3g* from *Glu-A3d*, *Glu-B3ab* from *Glu-B3b*, *Glu-B3ac* from *Glu-B3g*, *Glu-B3ad* from *Glu-B3i*, and *Glu-D3l* from *Glu-D3c*. Alleles *Glu-D3e* and *Glu-D3c* are assumed to be identical. The allele in cultivar Darius, with no distinct spot in 2-DE gels, is a new allele, *Glu-D3m*. The new allele *Glu-D3n* identified in the cultivar Fengmai 27 has a distinct spot in 2-DE and different mobility in SDS-PAGE (Figure 5). However, more work is needed to further characterize these new alleles at the *Glu-D3* locus. The other alleles were the same as those observed by Gupta and Shepherd [13].

Allele *Glu-A3g*, identified in the Canadian cultivars Bluesky and Glenlea by 2-DE in the current collection, is widely distributed in many cultivars from Canada and the U.S.A. [41]. In previous studies, allele *Glu-A3g* was frequently identified as *Glu-A3d* due to their similar SDS-PAGE patterns. The role of *Glu-A3g* in bread making quality therefore requires further study. Similarly, effects on bread making quality of alleles *Glu-B3ab*, *Glu-B3ac*, *Glu-B3ad* and *Glu-D3l*, with two additional distinct spots



Figure 6 Identification of LMW-GS by two-dimensional gel electrophoresis (2-DE). Discrimination of alleles *Glu-A3a*, *Glu-A3b*, *Glu-A3c*, *Glu-A3d*, *Glu-A3c*, *Glu-A3d*, *Glu-A3c*, *Glu-A3d*, *Glu-A3e*, and *Glu-A3f*. Cultivars: 1. Neixiang 188, 2. Gabo, 3. Pitic, 4. Nidera Baguette 10, 5. Amadina, 6. Kitanokaori.











compared to alleles *Glu-B3b*, *Glu-B3g*, *Glu-B3i* and *Glu-D3c*, respectively, also need further investigation.

Conclusions

Four methods, SDS-PAGE, 2-DE, MALDI-TOF-MS and PCR, were used for identifying the LMW-GS composition in wheat cultivars from 12 countries. All seven *Glu-A3* alleles could be identified by 2-DE and PCR, and only four and five of the seven could be differentiated by MALDI-TOF-MS and SDS-PAGE of the glutenin extract,

respectively. The *Glu-B3* alleles *Glu-B3a*, *Glu-B3b*, *Glu-B3c*, *Glu-B3g*, *Glu-B3h* and *Glu-B3j* could be identified by all four methods, but alleles *Glu-B3ab*, *Glu-B3ac*, *Glu-B3ad* could only be identified by the 2-DE method. *Glu-D3* alleles were very difficult to clearly distinguish by SDS-PAGE, 2-DE and PCR. MALDI-TOF-MS was promising in reliably differentiating them. PCR is a simple, accurate, and low cost method for identifying *Glu-A3* and *Glu-B3* alleles that are currently routinely analysed by SDS-PAGE in breeding programs. However, SDS-PAGE

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Locus	Subunit	SDS-PAGE	2-DE	MALDI-TOF-MS	PCR
Glu-A3	Glu-A3a	$\sqrt{*}$		\checkmark	
	Glu-A3b	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-A3c	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-A3d	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-A3e	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-A3f		\checkmark	\checkmark	\checkmark
	Glu-A3g		\checkmark		
Glu-B3	Glu-B3a		\checkmark		
	Glu-B3b	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-B3c	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-B3d	\checkmark		\checkmark	\checkmark
	Glu-B3f			\checkmark	\checkmark
	Glu-B3g	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-B3h	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-B3i	\checkmark			\checkmark
	Glu-B3j	\checkmark	\checkmark	\checkmark	\checkmark
	Glu-B3ab		\checkmark		
	Glu-B3ac		\checkmark		
	Glu-B3ad		\checkmark		
Glu-D3	Glu-D3a	\checkmark		\checkmark	-
	Glu-D3b	\checkmark		\checkmark	-
	Glu-D3c	\checkmark	\checkmark	\checkmark	-
	Glu-D3m		\checkmark	\checkmark	-
	Glu-D3l	\checkmark	\checkmark		-
	Glu-D3n		\checkmark		

Table 2: Allelic variants of LMW-GS identified using different methods

 $\sqrt{}$ - confirmed; - data not available

using a multi-gel buffer chamber, and running both gliadins and glutenin extracts is also a highly reliable method. A combination of all methods will help to identify specific alleles, especially potentially new alleles.

A set of 30 cultivars (Table 4) was recommended for identifying LMW-GS alleles. These standard cultivars cover all variants of LMW-GS in the collection investigated. Among them, Chinese Spring, Opata 85, Seri 82 and Pavon 76, are recommended as a core set to be included in each SDS-PAGE gel when identifying alleles of LMW-GS genes. The 30 cultivars have been placed in CIMMYT's and INRA Clermont Ferrand, France germplasm banks and seed is being multiplied to make them freely available as a set upon request. Accession numbers will be assigned once the *Glu-1/Glu-3* allelic composition is confirmed.

Methods

Plant materials

One hundred and three cultivars of common wheat collected from 12 countries were used to develop a set of standard cultivars for identification of LMW-GS alleles (Table 1). They included 21 cultivars from China, 19 from Argentina, 15 from Australia, 14 from France, 10 from Japan, eight from Mexico, seven from Canada, three from the USA, two from Italy, two from the Netherlands, one from Finland and one from Germany. These cultivars were widely utilized in investigating glutenin subunit

Subject	SDS-PAGE	2-DE	MALDI-TOF-MS	PCR
Required sample amount	40 μg (Protein)	150 μg (Protein)	0.04 μg (Protein)	2 μL (DNA)
Purity required	Low	High	High	Medium
Number of alleles	19	22	21	16
Alleles efficiently resolved	<i>Glu-B3b</i> and <i>Glu-B3g</i> , <i>Glu-B3d</i> and <i>Glu-B3i</i> ,	Glu-A3e, Glu-A3f, Glu-A3g, Glu-B3b, Glu-B3g, Glu-B3ab, Glu-B3ac, Glu-B3ad, Glu-D3l and Glu-D3m	Glu-A3e, and Glu-A3f; Glu-D3a, Glu-D3b, Glu-D3c, and Glu-D3m	<i>Glu-A3e</i> and <i>Glu-A3f,</i> <i>Glu-B3d</i> and <i>Glu-B3i,</i> <i>Glu-B3f</i> and <i>Glu-B3g</i>
Mass accuracy	Inaccurate	Inaccurate	Accurate	Accurate
pl	Unknown	Known	Unknown	Unknown
Cost of equipment	≈\$7,000	≈\$30,000	≈\$20,000-400,000	≈\$5,500
Cost per sample	≈\$1.0	≈\$70.0	≈\$0.3	≈\$0.3
Number of samples analysed per day for skilled technician	30-160*	1	100	100
Automation	Not possible	Not possible	Possible	Possible
Experience required	Considerable	Considerable	Less	Less
Safety	High toxicity	High toxicity	Safe	Toxicity
False positives	No	Yes	No	Yes
Accuracy level	++	+++	++	++

Table 3: Relative efficiencies of methods of gluten analysis for situations where cultivar identification is required

*. Thirty samples/day if running two gels. Up to 160 samples/day if using multi-gel (8 gels) buffer tank.

compositions and their relationships to processing quality [41].

Protein extraction

A similar protocol was adopted for protein extraction in all five laboratories. Proteins were extracted from 100 mg whole meal according to the sequential procedure of Branlard and Bancel [53]. The samples were treated with 1.0 mL of 50% propanol-1-ol (v/v) for 5 min with continuous vortexing, followed by incubation (20 min at 65°C), vortexing (5 min), and centrifugation (5 min at 10, 000 × g). This step was repeated three times to remove most of the gliadins. The glutenin in the pellet was reduced with 50% propanol-1-ol, 50 mM Tris-HCl solution containing 1% w/v dithiothreitol (DTT), after which 1.4% v/v of 4vinylpyridine was added, and alkylation was continued overnight at room temperature. The protein of each cultivar was extracted in three replicates.

SDS-PAGE

SDS-PAGE was performed in all five laboratories. Glutenin and gliadin protein extracts were separated using the method of Singh et al. [46] with some modifications in different laboratories to obtain the best resolution. To summarize, there were differences in three aspects. The concentrations of separation gel were 14.0% concentration (T) with 1.3% cross linker (C), 15.0% T with 1.3% C, 12.5% T with 0.97% C, 15.0% T with 1.4% C, and 13.5% T with 0.8% C in the laboratory of CAAS, CIMMYT, INRA, NARO and Universidad Nacionalm of Argentina, respectively. The pH for separation gel was pH8.8 in all laboratories except in CIMMYT with pH8.5. The currents of running gel were 16, 12.5, 30, 30 and 40 mA in the laboratory of CAAS, CIMMYT, INRA, NARO and Universidad Nacionalm of Argentina laboratory, respectively. Generally, lower current results in better resolution, but we could not find the optimum conditions for maximum resolution of LMW-GS in all laboratories since each laboratory used its own optimum conditions. Details were reported by Ikeda et al. [35].

The LMW-GS compositions were identified according to Singh et al. [46] and Jackson et al. [16] and the gliadins were used as indicators of LMW-GS based on the linkage between LMW-GS and gliadin because the gliadin composition can be screened more readily than specific LMW-GS. The nomenclature system of LMW-GS followed Gupta and Shepherd [13], Jackson et al. [16], Branlard et al. [34], Ikeda et al. [35], Appelbee et al. [19] and the catalogue of gene symbols for wheat <u>http://</u> wheat.pw.usda.gov/ggpages/awn/53/Textfile/WGC.html.

Table 4: Thirty cultivars recommended as standards for the determination of LMW-GS alleles

Locus	Allele	Standard cultivar
Glu-A3	Glu-A3a	Neixiang 188, Chinese Spring
	Glu-A3b	Gabo, Pavon 76
	Glu-A3c	Pitic, Seri 82
	Glu-A3d	Nidera Baguette 10, Cappelle-Desprez
	Glu-A3e	Amadina, Marquis
	Glu-A3f	Kitanokaori, Renan
	Glu-A3g	Bluesky, Glenlea
Glu-B3	Glu-B3a	Chinese Spring
	Glu-B3b	Renan, Gabo
	Glu-B3c	Insignia, Halberd
	Glu-B3d	Pepital, Ernest
	Glu-B3f	Fengmai 27
	Glu-B3g	Splendor, Cappelle-Desprez
	Glu-B3h	Aca 303, Pavon 76
	Glu-B3i	Norin 61
	Glu-B3j	Grebe, Seri 82
	Glu- B3ab	Nanbu-komugi
	Glu- B3ac	Thesee, Aca 801
	Glu- B3ad	Heilo, Opata 85
Glu- D3	Glu-D3a	Chinese Spring, Neixiang 188
	Glu-D3b	Gabo, Avocet
	Glu-D3c	Insignia, Cappelle-Desprez
	Glu- D3m	Darius
	Glu-D3l	Amadina, Heilo
	Glu-D3n	Fengmai 27

The core group is in bold

2-DE procedure

The 2-DE method was only performed at CAAS and NARO. The 2-DE procedure employed to identify LMW-GS was performed with an IPGphor (GE Healthcare, Sweden) for isoelectric focusing (IEF), and an AE-6530 chamber and an AE-8450 power supply (ATTO, Japan) for SDS-PAGE. The glutenin fraction was precipitated with 80% acetone [54], and the resulting pellets containing 150 μ g protein were dissolved in 250 μ L of IEF rehydration solution [7 M urea, 2 M thiourea, 4% w/v CHAPS, 2% v/v IPG buffer pH 6-11 (GE Healthcare) and

20 mM DTT] for very basic proteins [55]. After incubation for 30 min at room temperature, samples were applied to Immobiline Dry-Strip pH 6-11 (13 cm, GE Healthcare). The rehydration step was carried out for 12 h at 20°C. IEF was performed with a step-wise protocol to 45 kVh. After IEF, the strips were stored at -80°C or prepared directly for 2-DE as follows: the gel strips were first equilibrated under gentle shaking for 15 min in equilibration buffer (50 mM Tris-HCl, pH 8.8, 6 M urea, 30% v/v glycerol, 2% w/v SDS) with 2% w/v DTT, and then in equilibration buffer containing 1.4% v/v 4-vinylpyridine. The second dimension separations (SDS-PAGE) were carried out on 13% acrylamide constant gels and ran at 7 mA/gel for 45 min and then 25 mA/gel for approximately 4 h, until the bromophenol blue had run off the bottom of the gel [56]. After the completion of 2-DE, gels were fixed and stained with Coomassie Brilliant Blue-G250 according to Neuhoff et al. [57]. The resulting gels were scanned using an Image Scanner (GE Healthcare) and the images analyzed with ImageMaster 2D Platinum v6.0 software (GE Healthcare). At least three gel images of each sample were taken and compared. The LMW-GS compositions were identified with the distinctive spot on 2-DE gels according to Ikeda et al. [18]. The nomenclature system of LMW-GS was the same as above SDS-PAGE separation.

In some cases the 2-DE was modified where glutenin proteins were not alkylated; 16% isopropanol was added to the IEF buffer, and IEF was performed at 18 kVh [18].

MALDI-TOF-MS protocol

MALDI-TOF-MS was performed at the State Agriculture Biotechnology Center, Murdoch University, Australia. The glutenin fraction was precipitated with 80% acetone [54], and the resulting pellets containing 100 μ g protein were dissolved in 60 μ L acetonitrile (ACN)/H₂O (50:50 v/ v) containing 0.05% v/v trifluoroacetic acid (TFA) for 1 h at room temperature. Sample preparation was carried out according to the dried droplet method [58], using sinapinic acid (SA) as matrix. The matrix solution was prepared by dissolving SA in 50% ACN/0.05% TFA (w/w) at a concentration of 10 mg/mL. A sandwich matrix/sample/matrix 1:1:1 (0.7 μ L) was deposited on to a 96-sample MALDI target, and dried at room temperature.

MALDI-TOF-MS was performed on a Voyager DE-PRO TOF mass spectrometer (Applied Biosystems, Foster City, CA, USA) equipped with a 337 nm nitrogen laser and delayed extraction. Analyses were carried out on a positive linear ion mode at a mass range of 10000-50000 m/z with an accelerating voltage of 25 kV and a delay time of 900 ns. A low mass gate value of 10000 m/z was selected for analysis to avoid saturation of the detector. The identification of LMW-GS alleles based on MALDI-TOF-MS was established using a set of 19 near-isogenic lines (NIL) of cultivar Aroona (unpublished data, A Wang, W Ma, R Appels, Murdoch University, Australia).

DNA extraction and PCR amplification

PCR was performed only at CAAS. Genomic DNA was extracted from seeds using a modified CTAB procedure [59]. PCR was performed using TaKaRa (Dalian, China) *Taq* DNA polymerase (1.0 unit) in 20 μ L reaction volumes containing approximately 50 ng of genomic DNA, 1× PCR buffer (1.5 mM MgCl₂), 100 μ M of each dNTP and 7.5 pmol of each PCR primer. Details of allele-specific markers for the discrimination of *Glu-A3* and *Glu-B3* alleles and PCR conditions were reported previously [27,28].

Abbreviations

2-DE: two-dimensional gel electrophoresis (IEF × SDS-PAGE); ACN: acetonitrile; BIOLAB AZUL: Laboratory of Functional Biology and Biotechnology; CAAS: Chinese Academy of Agricultural Sciences; CEBB-MdP: Biotechnology and Biodiversity Study Center, Mar del Plata, Argentina; CHAPS: 3-[(3-Cholanidopropyl) dimethylammonio]-1-propanesulfonate; CICPBA: Scientific Research Commission of the Province of Buenos Aires; CIMMYT:International Maize and Wheat Improvement Center; CIISAS: Center for Integrated Research in Sustainable Agricultural Systems, COUNTRY; CONICET: National Science and Technology Research Council, Argentina; CRESCAA: Regional Center for Systemic Study of Agro-alimentary Chains, COUNTRY; CTAB: cetyltrimethylammonium bromide; DNA: deoxyribonucleic acid; dNTPs: deoxynucleoside triphosphates; DTT: dithiothreitol; HMW-GS: high-molecular-weight glutenin subunits; HPLC: highperformance liquid chromatography; IEF: isoelectric focusing; INBA: Research Institute of Agricultural and Environmental Biosciences: INRA: National Institute for Agricultural Research; kDa: kilodalton; kVh: kilo-volt-ampere-hour; LMW-GS: low-molecular-weight glutenin subunits; MALDI-TOF-MS: matrix-assisted laser desorption/ionization time-of-flight mass spectrometry; ${\rm MgCl}_{\rm 2}{\rm :}$ magnesium chloride; NARO: National Agriculture and Food Research Organization; NIL: near-isogenic lines; PCR: polymerase chain reaction; pl: isoelectric points; SA: sinapinic acid; SDS: sodium dodecyl sulfate; SDS-PAGE: sodium dodecyl sulphate-polyacrylamide gel electrophoresis; TFA: trifluoroacetic acid; Tris-HCI: tris (hydroxymethyl) aminomethane hydrochloride

Authors' contributions

LL carried out SDS-PAGE, 2-DE and MALDI-TOF-MS analyses and drafted the manuscript. TI participated in the design of the study, and performed SDS-PAGE and 2-DE analyses. GB participated in the design of the study and carried out SDS-PAGE analysis. RP participated in the design of the study and carried out SDS-PAGE analysis. RP participated in the design of the study and carried out study and revised the manuscript. SL carried out SDS-PAGE analysis and revised the manuscript. SL carried out SDS-PAGE analysis and revised the manuscript. LW carried out PCR analysis. WM participated in the identification of LMW-GS by MALDI-TOF-MS. RA participated in the identification of LMW-GS by MALDI-TOF-MS and revised the manuscript. HY participated in the design of the study. AW participated the identification of LMW-GS by MALDI-TOF-MS and revised the manuscript. HY participated in the design of the study. AW participated the identification of LMW-GS by MALDI-TOF-MS. The identification of LMW-GS by 2-DE. ZH conceived of the study and revised the manuscript. All authors read and approved the final manuscript.

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