

Structural characterization of the *DRF1* gene of *Aegilops speltoides* and comparison of its sequence with those of B and other *Triticeae* genomes

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Abstract The genus Aegilops L. has been intensively investigated due to its close relationship with wheat (Triticum L.) as contributor of B and D subgenomes. Because of their vast genetic diversity, Aegilops species represent a rich source of alleles of agronomic interest, which could be used to widen the wheat gene pool and improve tolerance to diseases, pests, drought, cold and other environmental stresses. We report the isolation and characterization of the Dehydration Responsive Factor 1 (DRF1) gene in three accessions of Ae. speltoides coming from different regions of the Fertile Crescent. The DRF1 gene belongs to the DREB gene family and encodes transcription factors which play a key role in plant response to water stress. As in other cereals, the DRF1 gene in Aegilops speltoides consists of four exons and

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CIMMYT (International Maize and Wheat Improvement Center), Apartado Postal 6-641, 06600 Mexico, DF, Mexico three introns and undergoes alternative splicing. A processed pseudogene was also identified and compared with the sequence of an actual mRNA transcript, breaking new ground in the understanding of the complex regulation mechanism of this gene. The genetic diversity was evaluated by comparison of inter- and intra-species variation among some *Aegilops* and *Triticeae*, by considering both the whole gene and exon 4 sequences. The phylogenetic analyses were able to cluster the sequences in well-supported clades attributable to the genomes analysed. The overall results suggest that there is a high similarity between the B and S genome copies of the *DRF1* gene but also features indicating that the two genomes have evolved independently.

Keywords Aegilops speltoides · Dehydration

Responsive Factor 1 (DRF1) gene · Genetic diversity · Phylogenetic analysis · Pseudogene

Abbreviations

AP2	Apetala 2
AS	Alternative splicing mechanism
DREB	Dehydration responsive element binding
	factor
DRF1	Dehydration responsive factor 1
ERF	Ethylene-responsive element binding factor
GP-2	Secondary gene pool
PTC	Premature termination codon
SSR	Simple sequence repeat

TF	Transcription Factor
Aes	Ae. speltoides
Aesl	Ae. speltoides ssp. ligustica
Aess	Ae. speltoides ssp. speltoides
At	A. thaliana
Та	T. aestivum L.
Td	T. durum (Triticum turgidum ssp. durum L.)

Introduction

The genus Aegilops L. has been intensively investigated due to the close relationship with wheat (Triticum L.), of which it forms the largest part of the so-called secondary gene pool (GP-2) (Harlan and de Wet 1971). Following the commonly accepted system suggested by Van Slageren (1994), the genus comprises 22 species arranged in five sections (Aegilops, Comopyrum, Cylindropyrum, Sitopsis and Vertebrata) and includes diploid, tetraploid and hexaploid genomes. The Sitopsis section is particularly interesting because its five species (Ae.bicornis-S^bS^b, Ae. longissima-S¹S¹, Ae. searsii-S^sS^s, Ae. sharonensis-S^{sh}S^{sh} and Ae. speltoides-SS) share the S genome found to be closely similar to the B genome in T. durum (Kerby and Kuspira 1987; Haider 2013; Ruban and Badaeva 2018), whereas Ae. tauschii (Vertebrata section) has been involved in the evolution of the wheat D genome (Marcussen et al. 2014; Pont et al. 2019). The Aegilops genus thus represents a natural laboratory for the study of the effects of the evolution and domestication of cultivated wheat.

Aegilops species are naturally distributed in an area ranging from the Mediterranean to Western and Central Asia. From the Transcaucasus, thought to be their centre of origin, diploid species migrated west and east, the centre of diversity being in the Fertile Crescent (modern day Iraq and parts of Turkey, Syria and Iran), which contains the majority of Aegilops species (Van Slageren 1994). Such a wide spread implies a great adaptability to different environments and a strong resistance to biotic and abiotic stresses. Indeed, many Aegilops species have a strong resistance to disease and insect pests, high drought and salt tolerance and/or good grain quality, thus representing a significant reservoir of new alleles with valuable traits for broadening the genetic diversity of wheat (Schneider et al. 2008; Brozynska et al. 2016; Zhang et al. 2017; Kumar et al. 2019). A number of studies have analysed and compared the nucleotide diversity of a single gene inside the *Triticeae* tribe; for instance Adderley and Sun (2013) analysed the nuclear phosphoglycerate kinase (pgk1) gene, Liu et al. (2016) studied the Wcor15 gene and Salse et al. (2008) investigated the storage protein activator (SPA) genomic region. In this work, we have analysed the Dehydration Responsive Factor 1 (DRF1) gene in some Aegilops species, with a focus on Aegilops speltoides. DRF1 belongs to the Dehydration Responsive Element Binding (DREB2) gene family and encodes important transcription factors (TFs) involved in plant response to drought. It is characterized by the presence of the APETALA2 (AP2)/Ethylene-Responsive element binding Factor (ERF) motif, through which the TFs bind to the *drought-responsive element* (DRE)/C repeat (CRT) element in the promoters of several stress responsive genes, thus modulating their expression (Agarwal et al. 2006). DREB2 genes were first identified in Arabidopsis thaliana by Yamaguchi-Shinozaki and Shinozaki (1994), and later on in various cereals, such as *Hordeum vulgare* (*HvDRF1*) (Xue and Loveridge 2004), Triticum aestivum (Wdreb2) (Egawa et al. 2006) and Triticum durum (TdDRF1) (Latini et al. 2007). In these cereals, the gene is structurally organized in four exons and three introns, and produces three transcripts through the alternative splicing mechanism (AS). Two of the three transcripts encode putative TF proteins endowed with the AP2 domain, while the other, the DRF1.2 transcript (exon 1, exon 2 and exon 4), encodes an abortive protein, due to the presence of a premature termination codon (PTC), caused by a frameshift mutation.

Here, the whole *DRF1* gene was amplified, cloned and sequenced from three accessions of *Aegilops speltoides*, its nucleotide diversity was analysed and compared with the B subgenome copy of the *DRF1* gene in *T. durum* (Cantale et al. 2018). Furthermore, the nucleotide diversity of the *DRF1* exon 4, containing the AP2 domain, was analysed in several accessions of *Aegilops speltoides* and other diploid and polyploid *Aegilops* species and compared with available corresponding *Triticeae* sequences to investigate the reciprocal relationships.

Materials and methods

Plant material

The *Aegilops* accessions used in the present study are listed in Table 1. Seeds were provided by ICARDA (International Center for Agricultural Research in the Dry Areas), except one accession provided by the Leibniz Institute of Plant Genetics and Crop Plant Research (IPK, DE).

Ten seeds from each accession were germinated in Petri dishes; the plantlets were then transferred into small pots and grown in a greenhouse under controlled standard light and temperature conditions. After 25–30 days, there were sufficient plant leaves (about 100 mg) for harvesting for the molecular analyses.

DNA/RNA extraction and PCR amplification

Genomic DNA was extracted from the leaves using the CTAB method (Murray and Thompson 1980), including a treatment with RNase I to avoid RNA contamination. Total RNA was extracted from the leaves of plants (Ae. speltoides ssp. speltoides, accession IG 47280) grown under normal conditions, using the RNAfast-II Isolation System (Molecular Systems, San Diego, CA, USA). The quality and integrity of the DNA/RNA extracts were checked using agarose gel electrophoresis and their final concentration was measured using a spectrophotometer (OD 260, 280, 260/280). Primer design was based on TdDRF1 gene sequence (NCBI ID: EU089819). The whole gene sequence was isolated by amplifying two gene fragments in optimized specific PCR reactions. The first amplicon was a ~ 2.5 Kb gene fragment obtained using the pair of primers drebfor1 (5'-CATGACGG-TAGATCGGAA-3') and drebrev1 (5'-TGCGCAGG-GAAGTTGGTA-3'). This ~ 2.5 Kb gene fragment mapped exon 1, including the ATG start codon, up to part of exon 4 (experimental conditions: annealing temperature 55 °C for 1 m, extension temperature 72 °C for 5 m). The second amplicon was a ~ 1 Kb gene fragment, obtained using the pair of primers drebfor2 (5'-CATGATCCACAGGGTGCAA-3') and E4rev-down (5'-GGTCCACCATTTGATCTTCATT-3'). This ~ 1 Kb gene fragment overlapped the previous one and completed exon 4, including the 3'-terminal of the gene (experimental conditions: annealing temperature 58 °C for 1 m, extension temperature 72 °C for 2 m). This latter pair of primers was also used to amplify the exon 4 fragment from 42 *Aegilops* accessions (see Table 1). The PCR reactions were performed in an overall volume of 25 µl, containing 12.5 µl REDTaq[®] ReadyMixTM PCR reaction mix (20 mM Tris-HCl, pH 8.3, 100 mM KCl, 3 mM MgCl2, 0.002% gelatin, 0.4 mM dNTP mix, stabilizers and 0.06 unit/ml of Taq DNA polymerase), 20 pmol of each primer and 200 ng of gDNA. The PCR thermal cycle conditions included an initial denaturation at 95 °C for 5 m, then 35 cycles, with melting at 94 °C for 1 m, followed by the annealing and extension steps as reported above for each pair of primers, ending at + 4 °C.

Furthermore, the pair of primers *drebfor1* and *E4rev-down* was used to amplify transcripts from the RNA extract, following the Latini et al. procedure (2007).

In Online Resource 1, the annealing regions of the forward and reverse primers used are represented.

PCR products purification and bacterial transformation

The PCR products were purified using the Gel Band Purification kit (Amersham, USA), followed by Sepharose CL-6B affinity chromatography (Pharmacia Corp., NYC, NY, USA), and then cloned into a $pCR^{\text{B}}\text{II-TOPO}^{\text{B}}$ vector (Invitrogen, USA). After transformation of *E. coli* cells, positive clones were screened by using agar plates containing ampicillin (50 µl/ml), X-Gal (40 mg/ml) and IPTG (40 µl of 100 mM IPTG - Isopropyl β-D-1-thiogalactopyranoside solution). Plasmids were extracted using the SDS plasmid isolation protocol (Sambrook et al. 1990). In order to confirm the presence of the expected cloned fragment, plasmid DNA clones were digested before sequencing by the EcoRI restriction enzyme and visualized on agarose gel.

Sequences from public databases

Various relevant *DRF1* sequences, belonging to the *Aegilops* and *Triticum* genera, were downloaded from NCBI (https://www.ncbi.nlm.nih.gov/) and other data sources. A list of these sequences is provided in Table 2.

sequences derived from different seeds of the same accession) *IPK: accession provided by IPK (Leibniz Institute of Plant Genetics and Crop Plant Research, DE)

Species	Genomic code	IG	Country province	NCBI ID
Ae. speltoides ssp. speltoides	SS	117905	SYR Aleppo	MK803147-48
Ae. speltoides ssp. ligustica	SS	48994	IRQ Ninawa	FJ843102; MK803170
Ae. speltoides ssp. ligustica	SS	47948	TUR Elazig	FJ858187; MK803182
Ae. speltoides ssp. speltoides	SS	110757	SYR Al Hasakah	FJ858188; MK803192
Ae. speltoides ssp. speltoides	SS	126225	SYR Hama	MK803193; MN639698
Ae. speltoides ssp. speltoides	SS	116071	TUR Gaziantep	MK803194-95; MN639699
Ae. speltoides ssp. ligustica	SS	48994	IRQ Ninawa	MK803149-50
Ae. speltoides ssp. speltoides	SS	48993	IRQ Ninawa	MK803151-53
Ae. speltoides ssp. ligustica	SS	48847	IRN Kordestan	MK803154-58
Ae. speltoides ssp. ligustica	SS	48402	SYR Al Hasakah	FJ868038-41; MK803159
Ae. speltoides ssp. speltoides	SS	48764	LBN Chouf	MK803160-63
Ae. speltoides ssp. speltoides	SS	48480	SYR Al Hasakah	MK803164-65
Ae. speltoides ssp. speltoides	SS	47517	SYR Idlib	FJ917411-14
Ae. speltoides ssp. speltoides	SS	131278	IRQ Diyala	MK803166-69
Ae. speltoides ssp. speltoides	SS	131280	IRQ As Sulaymaniyah	MK803171-72
Ae. speltoides ssp. ligustica	SS	47506	SYR Lattakia	MK803174-76
Ae. speltoides ssp. speltoides	SS	47280	SYR Lattakia	FJ917415-18
Ae. speltoides ssp. ligustica	SS	46809	TUR Diyarbakir	FJ917419-22; MK803177
Ae. speltoides ssp. ligustica	SS	47115	TUR Bingol	MK803178
Ae. speltoides ssp. ligustica	SS	47122	TUR Hakkary	FJ930929-32
Ae. speltoides ssp. ligustica	SS	116065	TUR Gaziantep	MK803179-80
Ae. speltoides ssp. speltoides	SS	116079	TUR Gaziantep	MK803181
Ae. speltoides ssp. speltoides	SS	47947	TUR Elazig	MK803183
Ae. speltoides ssp. speltoides	SS	47954	TUR Elazig	MK803184-85
Ae. speltoides ssp. speltoides	SS	46560	SYR Aleppo	MK803186-87
Ae. speltoides ssp. speltoides	SS	47496	SYR Lattakia	MK803188
Ae. speltoides ssp. speltoides	SS	135337	SYR Al Hasakah	MK803191
Ae. speltoides Tausch	SS	*IPK:AE 413/85	ISR	MK803189-90
Aegilops biuncialis	$U^{b}U^{b}M^{b}M^{b}$	46559	SYR Aleppo	MK803133
Aegilops biuncialis	$U^{b}U^{b}M^{b}M^{b}$	46924	TUR Kirsehir	MK803134
Aegilops peregrina	$U^{p}U^{p}S^{p}S^{p}$	132603	AZE Abseron	MK803135
Aegilops peregrina	$U^{p}U^{p}S^{p}S^{p}$	48714	JOR Amman	MK803136
Aegilops umbellulata	UU	46952	TUR Diyarbakir	MK803137
Aegilops umbellulata	UU	49022	IRQ Ninawa	MK803138
Ae. triuncialis ssp. triuncialis	$U^t U^t C^t C^t$	47828	BGR Burgas	MK803139
Ae. triuncialis ssp. triuncialis	$U^t U^t C^t C^t$	48476	SYR Al Hasakah	MK803140
Aegilops neglecta	$U^{n}U^{n}X^{n}X^{n}/$ $U^{n}U^{n}N^{n}N^{n}X^{n}X^{n}$	48837	IRN Lorestan	MK803141
Aegilops geniculata	$U^g U^g M^g M^g$	46576	SYR Aleppo	MK803142
Aegilops geniculata	$U^g U^g M^g M^g$	47311	JOR Irbid	MK803143
Aegilops cylindrica	$D^{c}D^{c}C^{c}C^{c}$	48789	LBN Harmel	MK803144
Aegilops sharonensis	$S^{sh}S^{sh}$	47098	RUS	MK803145
Aegilops sharonensis	$S^{sh}S^{sh}$	47080	PSE	MK803146

Table 2 List of the sequences description	Genus Species		Gene	NCBI ID	Genome
genetic analyses, including	Aegilops	Ae. speltoides	DREB2	KF731665	S
the gene name, the		Ae. tauschii	DRF1	EU197052	D
and the genome type when		Ae. tauschii	-Chr1	BLAST*	D
provided by authors of	Triticum	T. urartu	DREB2	KF731664	А
submission		T. urartu	DRF1	EU197053	А
		T. durum	DRF1	KM520369.2	А
		T. durum	DRF1	KM504244.1	А
BLAST*: sequence retrieved using a BLAST		T. durum	DRF1	KM520370.2	В
search at UC Davis (USA),		T. durum	DRF1	KM504245.1	В
Aegilops tauschii BAC		T. durum	DRF1	FJ560496	na
scaffolds (http://aegilops.		T. durum	DRF1	EU781993	na
wheat.ucdavies.edu/		T. aestivum	-Chr1B	BLAST**	В
BLAST**: sequence		T. aestivum	Wdreb2	AB193608	na
retrieved using a BLAST		T. aestivum	DREB3B	AY781350	na
search at URGI-IWGSC		T. aestivum	DREB4B	AY781355	na
(https://urgi.versailles.inra.		T. aestivum	DREB5B	AY781359	na
no: ganoma information		T. aestivum	DREB	KF737910	na
was not provided		T. aestivum	DREB	JQ004969	na
^a Sequences downloaded		T. dicoccoides	DRF1.3	HM015905	na
from GenBank and used as	Psathyrostachys	Psathyrostachys juncea	DREB2	JF766085	а
outgroup in the phylogenetic analysis	Brachypodium	Brachypodium distachyon	DREB2B	XM003568607	a

Data analyses

Electropherograms were analyzed using FinchTV (Geospiza Inc., USA) and sequences were aligned using Clustal Omega at EMBL-EBI (Sievers et al. 2011) and assembled using CAP3 (Huang and Madan 1999). FGENESH (by Softberry) at http://www.softberry.com/berry.phtml?topic=fgenesh&group=programs&subgroup=gfind (Soloviev et al. 2006) was used to identify the gene structure (parameter settings: organism = triticum; suboptimal exon cutoff = 0.1).

DnaSP v6 (Rozas et al. 2017) was used to assess the diversity and genetic relationships among sequences. Analyses were carried out using alignments manually edited to improve their accuracy and excluded gaps and missing data.

Phylogenetic analyses were carried out using MEGA 7 (Kumar et al. 2016). The Neighbor-Joining (NJ) method was applied and the evolutionary distances were computed using the Maximum Composite Likelihood model. A bootstrap analysis based

on 1000 re-samplings was used to determine the confidence values of the tree branches. All positions containing gaps and missing data were eliminated. For completeness, a separate analysis was also conducted by applying the Maximum Likelihood method and PHYML algorithm at http://www.atgc-montpellier.fr/phyml/ (Guindon and Gascuel 2003; Lefort et al. 2017).

Results

The *DRF1* gene in *Aegilops speltoides* and comparison with the B subgenome copy of *T*. *durum*

The whole *DRF1* gene was isolated and sequenced from three *Ae. speltoides* accessions. Representative PCR amplification and restriction results after cloning of the gene fragments are shown in Fig. 1. The three gene sequences were submitted to GenBank: two

accessions of *Ae. speltoides* ssp. *ligustica* (ICARDA accessions IG 48994 with NCBI ID FJ843102, 3214 bp long and IG 47948 with NCBI ID FJ858187, 3229 bp long) and one accession of *Ae. speltoides* ssp. *speltoides* (ICARDA accession IG 110757 with NCBI ID FJ858188, 3214 bp long).

The FGENESH gene finder tool, used for the prediction of the gene structure of the *Ae. speltoides* sequences, recognized the presence of four exons and three introns, with a high confidence, showing higher scores for exons 1 and 4 than for exons 2 and 3. The same structure had been already observed in wheat and barley. As an example, the prediction obtained using

the *AeslDRF1*-FJ843102 sequence as the template is shown in Fig. 2. An analogous result was obtained when using the other two sequences as templates.

The three *Ae. speltoides* sequences were aligned and the genetic diversity quantification is summarized in Table 3. 3210 sites were analysed and 20 out of 30 identified polymorphisms were located in exon 4, with an overall nucleotide diversity of Pi = 0.006. A total of 4 indels (mutations due to the insertion or deletion of nucleotides) were observed and the lack of a triplet in the simple sequence repeat (SSR) region of exon 1 in *AeslDRF1*-FJ858187 was particularly interesting, as it caused a shorter alanine stretch in the putative



Fig. 1 Electrophoresis gel of products obtained by PCR amplifications of genomic DNA. **a** 2.5 Kb and 500 bp fragments amplified using the pair of primers *drebfor1* and *drebrev1*. Lane 1: *Ae. speltoides* ssp. *speltoides* IG 110757; lane 2: *Ae. speltoides* ssp. *ligustica* IG 47948; lane 3: *Ae. speltoides* ssp. *ligustica* IG 248994. **b** 1 Kb fragment amplified using the pair of primers *drebfor2* and *E4rev-down*. Lane 1: *Ae. speltoides* ssp. *ligustica* IG 47948. **c** EcoRI restriction analysis of clones containing the ~ 500 bp fragments. The samples in lanes 2 and

4 (*Ae. speltoides* ssp. *speltoides* IG 126225 and IG 116071 accessions, respectively) represented two positive clones, whose sequences were deposited (lateral arms of vector were included in the bands, thus justifying 600 bp size), while lanes 1 and 3 were found to be empty plasmids. **d** 1 Kb fragment of exon 4 amplified using the pair of primers *drebfor2 E4rev-down* primers, from various accessions of *Ae. speltoides*. Lane 1: IG 117905, seed 1; lane 2: IG 117905, seed 2; lane 3: IG 116079; lane 4: IG 135337. MM stands for 1 kb molecular marker size

 FGENESH 2.6 Prediction of potential genes in Triticum genomic DNA

 Seq name: Aes/DRF1-FJ843102

 Length of sequence: 3214

 Number of predicted genes 1: in +chain 1, in -chain 0.

 Number of predicted exons 4: in +chain 4, in -chain 0.

 Positions of predicted genes and exons: Variant 1 from 1, Score:185.719043

 ▶ CDSf
 CDSi
 CDSo
 ◆ PolA
 ▼TSS



Fig. 2 The FGENESH prediction of gene structure using *AeslDRF1*-FJ843102 as template. CDSf stands for first exon including start codon, CDSi stands for internal exon, CDSI stands for last coding segment, ending with stop codon, CDSo

stands for genes with only one CDS (starting with start codon and ending with stop codon) TSS stands for the position of transcription start, PolA stands for polyA segment

protein, whereas the other three indels were located in non-coding regions. Worth noting is that the *AessDRF1-FJ858188* sequence showed a nonsynony-mous mutation at nucleotide position 2431, within the AP2 domain coding region, resulting in the replacement of a Thr (<u>A</u>CG) in an Ala (<u>G</u>CG). All the above-reported polymorphisms are highlighted in the alignment shown in Online Resource 1.

Afterwards, the Ae. speltoides sequences were aligned with two different B subgenome copies of T. durum, namely KM520370 and KM504245, recently published (Cantale et al. 2018). A very high similarity (more than 93% identity) was observed among the sequences. The analysis involved 2989 sites and identified 5 haplotypes and 135 polymorphic sites, the overall nucleotide diversity being Pi = 0.025(Table 3). Focusing on the coding regions, the SSR region in exon 1 of the Aegilops group is shorter and characterized by the GCG triplet, while the B genome copies of T. durum showed the distinctive GCC triplet (both triplets coding for alanine). Furthermore, two inframe deletions, each 15 bp long, were present in exon 4 of the B genome copies of T. durum. As regards the non-coding regions, two different insertions at alignment position 325 bp (intron 1) distinguished the Aegilops and Triticum groups, the former with a length of 65 bp and the latter of 25 bp, characterized by completely different nucleotide sequences. There were also few other differences among the indels of the two groups (SSR and indels are highlighted in light grey in Online Resource 1).

Transcription of the *DRF1* gene in *Aegilops* speltoides

The activity of the DRF1 gene in Aegilops speltoides was ascertained with RNA extraction and reverse transcription PCR (RT-PCR) which revealed the presence of the DRF1.2 transcript (Online Resource 2). Its exonic combination (exon 1, exon 2 and exon 4) generated a frameshift causing a premature termination codon (PTC). The sequence of this AesDRF1.2 transcript, isolated from accession IG 47280, was deposited at GenBank (NCBI ID: EU197054) and used as the template in a BLAST search. High identity values were observed between this and DRF1 transcripts of different varieties of T. durum, such as Atil (NCBI ID EU089824, 95.43%), Ciccio (NCBI ID FJ560492, 93.78%) and Karalis (NCBI ID FJ560496, 93.77%), T. aestivum, such as TaDREB3C (NCBI ID AY781351, 95.34%), TaDREB4C (NCBI ID AY781356, 95.16%) and TaDREB5C (NCBI ID AY781357, 95.52%), and also the T. dicoccoides *DRE*-binding transcription factor (NCBI ID HM015904, 95.43%). The DRF1.2 transcript was the only one detectable since plants were grown in normal conditions, as reported in our previous work (Latini et al. 2013).

			Singleton 3 variants	1	Coding region: all in exon 4	Non-synonymous changes	1			Singleton 3 variants	1		Coding region: in exon 4	Non-synonymous changes	1					
nic sites		30	30 Singleton 2 variants			ts	29	all in exon 4	Non-synonymous changes	6	nic sites		ts			n exon 2 and 4	Non-synonymous changes	20		
Polymorph				Singleton 2 variant	Singleton 2 varian	29		29	29	29	Coding region:	Synonymous changes	10	Polymorph	135	Singleton 2 varian		135	Coding region: ir	Synonymous changes
						Non coding region	10							ng region	35					
		Parsimony informative 2 variants -					Parsimony informative 2 variants	102		Non codi	5									
versity Pi		9	Coding region	region 0.01427 (E4)				versity Pi	5	Coding region		0,0338 (E1)	0.0075 (E2)	0.0068 (E3)	(17) 1070.0					
Nucleotide di	900.0 I		Non coding region		0.00250 (11)	0.00349 (I3)		Nucleotide di	0.02	Non coding region			0.0193 (11)	0.0448 (I3)						
Sites	analysed	3210						S ites analysed			0000	6867								
Haplotypes		en						Haplotypes			ı	n								
Gene	seduences							Gene sequences		AesDRF1+ TdDRF1	(FJ843102,	FJ858187,	FJ858188,	KM 520370, KM 504245)						

Table 3 Quantification of genetic variation in the sequences analysed

Deringer

AesDRFI stands for the alignment of the sequences of three accessions of Ae. spelvoides DRFI gene. AesDRFI + TdDRFI stands for the alignment of the sequences of three accessions of Ae. spelvoides and two varieties of T. durum DRFI gene. Parsimony informative stands for sites that have a minimum of two nucleotides that are present at least twice. The analysis was carried out with DNAsp software

Identification of a *DRF1* pseudogene in *Aegilops* speltoides

The genomic DNA amplification using the pair of primers drebfor1 and drebrev1 always amplified shorter fragments, ~ 500 bp, in addition to the expected ~ 2.5 Kb gene fragment (Fig. 1a). These amplicons were cloned (Fig. 1c) and sequenced. For each sample, some sequences turned out to be unspecific genomic fragments, but one corresponded to a DRF1 gene copy lacking introns. The analysis showed that this genomic sequence joined exons 1, 2 and 4, thus looking like the mature DRF1.2 transcript. This structure pointed to the presence in the Ae. speltoides genome of a pseudogene linked to its cognate DRF1 gene. Two sequences, obtained from the accessions IG126225 and IG116071, were deposited at GenBank (NCBI IDs MN639698 and MN639699, respectively). The sequences alignment of these two pseudogenes, the above described mRNA of DRF1.2 of Ae. speltoides and a DRF1.2 transcript of the B genome of *T. durum*, is shown in Fig. 3.

Nucleotide diversity analysis of exon 4 in Aegilops

A genomic DNA fragment of about 1 Kb, mapping in exon 4 (Fig. 1d), was isolated from 42 Aegilops accessions, obtaining a total of 95 cleaned sequences; then, after removing identical sequences obtained from different seeds of the same accession, 81 unique sequences were submitted to GenBank (see Table 1 for Aegilops accessions and the GenBank accession numbers). Three other sequences of *Aegilops* mapping DRF1 exon 4 were downloaded from public databases (see Table 2) and added to the set. The nucleotide diversity of these 84 sequences was analysed and the results are reported in Table 4. Sequences were analysed as a sole group (bulked group) and also split into subgroups, in accordance with their subspecies (Ae. speltoides ssp. speltoides and Ae. speltoides ssp. ligustica, Ae. speltoides, Aegilops). A further analysis was carried out by merging all Ae. speltoides sequences, irrespective of subspecies (Ae speltoides merged group). Interestingly, the nucleotide diversity of diploid Aegilops speltoides sequences was Pi = 0.018, when considering both the merged and individual subgroups, and it was 4.5 times higher than the nucleotide variation observed when analysing the same region in various Triticum durum varieties (nucleotide diversity Pi = 0.0040, data not shown). Most sequences represented a single, non-shared haplotype (71 haplotypes out of 84 sequences) (Table 4), except a few cases shared inside and between taxa, with Haplo 4 being the most common one (Table 5). Furthermore, a comparison between the Ae. speltoides ssp. ligustica and Ae. speltoides ssp. speltoides subspecies highlighted the presence of several SNPs, even if most of them were synonymous and did not cause replacements (data not shown). A phylogenetic analysis was carried out based on these 84 sequences using both the Neighbor-Joining (NJ) and Maximum Likelihood (ML) methods and the inferred NJ tree is shown in Fig. 4. The inferred ML tree showed the same topology (data not shown). Aegilops speltoides and the other Aegilops sequences formed two well-supported clusters and no noticeable separation could be observed between Ae. speltoides ssp. speltoides and Ae. speltoides ssp. ligustica subspecies sequences.

To investigate the relationships among DRF1 exon 4 sequences in Aegilops and Triticum, 16 DRF1 exon 4 sequences from the Triticeae tribe (7 out of 16 were isolated, sequenced and submitted to GenBank by our laboratory) were added to the dataset (see Table 2) and the total 102 sequences, including 2 outgroups, were aligned and analysed. Two further sequences of Psathyrostachys juncea and Brachypodium distachyon mapping DRF1 exon 4 were also added as outgroups. The NJ phylogenetic tree inferred is represented in Fig. 5. The 102 sequences formed distinct clades: all Ae. speltoides sequences clustered in a single clade (S genome), which was graphically collapsed to better highlight the relationships with the other taxa; four sequences grouped together with three B copies of the gene in T. durum, forming a wellsupported clade (96%, B genome), sister to the S one; four other sequences grouped together with two A copies of the gene in T. durum, forming a wellsupported clade (99%, A genome); two sequences grouped together with three Ae. tauschii sequences, forming the D genome clade (99%) and finally all other Aegilops sequences, sharing the U genome copy, formed a single clade (92%, U genome) distinct from the Ae. cylindrica sequence.

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	arebjori	
MN639698	ATGACG GTAGA TCGGA AGGA CGCTG AGGCG GCGG CGGCG GCGGC CGCACCCTT CGAGA TC	60
MN639699	ATGACE GTAGA TCGGA AGGA CGCTG A GCC GCCG CGGCC GCCGCC CCCTT CGAGA TC	60
EU197054	ATGACG GTAGA TCGGA AGCA CGCCG AGGCG GCGG CGGCG GCG CCCTT CGAGA TC	54
KM520370	ATGACG GTAGA TCGGA AGGA CGCTG AGGCG GCCG CCGCCGC CGCA CCCTT CGAGA TC	57

MN629698	RESERVED STORE STATE AND A STREET STORE SCALE AGE A STACCOSS A STCATCTC	120
MIE2GEGG		120
ET107054	CCGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	114
E0197034	CCGGCGCTCCAGCCTGGAAGAACTTGTGGAGCAGAGGAAAGTACCCCGGAGTCATGTTCTC	117
121320870		11/
01600600	CT (3.3.3. (C).3.5.3.C (C).3.3.3.C (C). (C). (C). (C). (C). (C). (C). (C	180
101639690	CTCANCENAL AGRAMANGEOREE COGROUP CALCELOR ALLOSS CELLAR FEASIER CONTRACTOR	180
EN039099		100
20197054	GILAAGULAAI AGGAAAAAGUGAULIUGGAGAILAUGIGAIGGAULIAAIILAGILIUIG	1/4
KM520370	GICAAACCAAIAC GAAAAAGEGAUGHUGGAGAHAAUGUGAHGGGGUGAHGAGUGAHGAGUGHUHG	177
01600600	111001 PO100000 PC (1110 PO1001 001 001 001 PC (0)	0.40
220029090		240
NN639699	AAACGA TCAGG CGATG GAAA GAAG TGAACCAACAACTGG AGCAT GATCCACAG GGTGCAA	240
20197054	AAACGGICAGGCGAIGGAAAGAAGIGAACIGGAGCAIGAICCACAGGGIGCGA	234
KM520370	AAACGATCAGGCGATGGAAAGAAGTGAACCAACAACTGGAGCATGATCCACAGGGTTCAA	237

MIE2GEG8	36366676366336773777673336667773333656776737673	200
MIESSEGG	16165666166116666166676611166677611161166166666776717661666661116616	200
PT107054	10100000000000000000000000000000000000	204
EM520270	ACAGES CONSCREDENCE ACCESSING CARACTERISTICS AND CARACTERISTICS ACCESSING AND CARACTERISTICS ACCESSING CARACTERISTICS ACC	297

MN629698	GACCTGAGAAT ACACA ATGT GGATT CCGTGGTGT AAGGCAACGT ACTT GGGGGAAGTGGG	260
MN639699	GACCTG AGAAT ACACA ATGT GGATT CCGTG GTGT AAGGC AACGT ACTT GGGGG AAGTG GG	360
EU197054	GACCCGAGAATACACAATGT GGTTC CCGTGGTGT AAGGC AACGT ACTT GGGGG AAGTG GG	354
KM520270	GACCTGAGAAT ACACA ATGT GGATT COGTGGTGT AAGGCAACGT ACTT GGGGGAAGTGGG	257

MN629698	TTGCTGAAATT CGGGA GCCA AATCG GGTGA GCAG GCTCT GGTTG GGAA CGTTC CCCACTG	420
MN629699	TTGCTG ANATT CGGGA GCCA ANTCGGGTGA GCAGGCTCT GGTTGGGAA CGTTC CCCACTG	420
EU197054	TTGCTG ANATT CGGGA GCCA ANTCGGGTGA GCAGGCTCT GGTTG GGAA CGTTC CCCACTG	414
KM520270	TTECTER RETECTED SOCIAL TO SECTOR SCALE S	417

MN639698	CTGAGGATGCT GCCCGTGCTTATGA CGAGGCAGCCAGAGCAATGTATGGCGCA CTGGCTC	480
MIE2GEGG	CTG3GG3TGCTGCCCGTGCTT3TG3CG3GGC3GCC3G3GC33TGT3TGGCGC3CTGGCTC	480
EII197054	CTG3GG3TGCTGCCGTGCTT3TG3CG3GGC3GCC3G3GCC3G3GTGT3TGCCGC3CTC	474
E0197034		477
1020010		11/
	deterd	
	arebrevi	
MN639698	GTACCAACTTCCCTGTGCA 499	
MN639699	GTACCAACTTC CCTGT GCA 499	
EU197054	GTACCAACTTC CCTGT GCA 493	
KM520370	GTGCCAACTTCCCTGTGCA 496	

Fig. 3 Alignment of two about 500 bp long fragments with two *DRF1.2* transcripts. MN639698 and MN639699 are the NCBI ID of two sequences, isolated from *Ae. speltoides* accessions IG126225 and IG116071, respectively; EU197054 is the sequence of the *AesDRF1.2* transcript (accession IG 47280); KM520370 is the sequence of a *TdDRF1.2* transcript from the B

Discussion

The study of gene variability in the *Aegilops* species has recently attracted considerable attention as wild relatives represent a valuable source of gene alleles

subgenome copy of *T. durum*. Exon 1 is shown in bold, exon2 using white letters and black background and exon 4 using white letters and grey background. The SSR region, consisting of 15 bp, GCG pattern, plus further 6 bp (all coding for alanine) and the premature stop codon are highlighted in boxes

that can be exploited for wheat improvement. In this context, the *DRF1* gene, involved in the drought response in plants, was studied in *Aegilops speltoides* and in other *Aegilops* species to investigate its various features. The gene was isolated and sequenced in three

Group	Ν	Pi	Н	Hd	θ	S	Tajima's D*
Bulked	84	0.0325	71	0.994	0.0416	169	- 1.11724
Ae. speltoides ssp. speltoides	36	0.0179	31	0.989	0.0241	81	- 1.00961
Ae. speltoides ssp. ligustica	29	0.0168	27	0.995	0.0172	55	- 0.22041
Ae. speltoides	3	0.0189	3	1.000	0.0189	23	nc
Ae. speltoides merged	68	0.0178	57	0.992	0.0257	100	- 1.21173
Aegilops	16	0.0382	14	0.983	0.0312	84	0.30729

Table 4 Estimates of nucleotide diversity in exon 4 sequences evaluated in the various groups analysed

The analysis was carried out with DNAsp software

N number of sequences, Pi nucleotide diversity, H number of haplotypes, Hd haplotype diversity, θ Watterson's θ value, S number of segregating sites; *not significant, P > 0.10; nc = not calculable

Table 5 Eight haplotypes which were found to be shared within or between the groups and their frequencies

Shared Haplotype	36 sequences Ae. speltoides ssp. speltoides	29 sequences Ae. speltoides ssp. ligustica	3 sequences Ae. speltoides	16 sequences Aegilops	84 sequences bulked sequences
Haplo 1	0.083 (3/36)	0	0	0	0.036 (3/84)
Haplo 2	0.055 (2/36)	0.034 (1/29)	0	0	0.036 (3/84)
Haplo 4	0.083 (3/36)	0.069 (2/29)	0	0	0.059 (5/84)
Haplo10	0.027 (1/36)	0.034 (1/29)	0	0	0.024 (2/84)
Haplo 33	0.027 (1/36)	0.034 (1/29)	0	0	0.024 (2/84)
Haplo 45	0	0.069 (2/29)	0	0	0.024 (2/84)
Haplo 57	0	0	0	0.125 (2/16)	0.024 (2/84)
Haplo 68	0	0	0	0.125 (2/16)	0.024 (2/84)

The analysis was carried out with DNAsp software

accessions of *Ae. speltoides* and its structure fully corresponded to that previously observed in barley and wheat (Xue and Loveridge 2004; Egawa et al. 2006; Latini et al. 2007), as deduced by sequence similarity and gene structure prediction (Fig. 2).

The detection of the *DRF1.2* transcript in *Ae. speltoides* suggested the occurrence of transcription involving the alternative splicing mechanism as this transcript arises by joining exons 1, 2 and 4. It was the only transcript isolated, as plants were grown in normal conditions and it is reasonable to speculate that the other two transcripts, *DRF1.1* and *DRF1.3*, would also be expressed and detectable in plants when affected by drought stress, as previously demonstrated in *Triticeae* (Xue and Loveridge 2004; Egawa et al. 2006; Latini et al. 2007). The sequence of this transcript was highly similar to the same transcripts in wheat and barley and does not encode for a transcription factor, but for an abortive short protein whose function is still unknown. Such a kind of transcript variant can often be found in alternatively spliced genes and is reported to play an important role in gene regulation through its modulated expression and degradation by nonsense-mediated mRNA decay (Dubrovina et al. 2013) or other RNA surveillance mechanisms (Reddy et al. 2012; Bedre et al. 2019).

The isolation of a pseudogene suggests an increasingly complex scenario for *DRF1* gene regulation in *Ae. speltoides*. Indeed, during the isolation of the whole gene, an unexpected 500 bp genomic fragment was always amplified and its sequence corresponded to the genomic variant of the mature *DRF1.2* transcript, thus fully satisfying the criteria for representing a processed pseudogene (retropseudogene) arising by reverse transcription and reinsertion into the genome (Zou et al. 2009; Tutar 2012). Because they are derived from mature transcripts, retropseudogenes lack upstream promoters, introns and other regulatory elements and were previously considered transcriptionally silent, dead-on-arrival (Vanin 1985). However, more recently, there has been increasing evidence that they are still active and often involved in the regulation of their parental genes, particularly at the post-transcriptional level, through various mechanisms including RNA interference and RNA antisense (Vinckenbosh et al. 2006; Wen et al. 2012; Sen and Ghosh 2013; Xie et al. 2019). To the best of our knowledge, this is the first reported case of a pseudogene belonging to the DREB gene family in Triticeae.

More targeted research is necessary to investigate the potential roles of both the *DRF1.2* transcript and pseudogene in *DRF1* gene regulation in *Ae. speltoides*.

The analysis of DNA polymorphisms of the nucleotide sequences of the three Ae. speltoides whole DRF1 gene accessions revealed a relatively low total nucleotide diversity (Pi = 0.006), in line with the observation reported that TFs and particularly those containing specific domains, like the AP2 domain, are more conserved than other gene families (Tatarinova et al. 2016). Interestingly, the nucleotide diversity turned out to be lower in introns than in exons, which was somewhat unexpected, as introns are considered freer to accumulate mutations than exons, which are constrained by their function. This genetic feature (exons more variable than introns) has been observed in animals, in specific gene families involved with the immune system or associated with predator-prey responses, and its role in adaptive evolution has been hypothesised (Nakashima et al. 1995; Samonte and Eichler 2002; Caporale 2003). Accordingly, one can speculate that the molecular evolution of the DRF1 gene, involved in response to abiotic stress, has undergone a similar genetic mechanism, even if more investigation is necessary to confirm this result.

The DNA sequence polymorphism analyses highlighted that there were no differences between the two *Ae. speltoides* subspecies, ssp. *ligustica* and ssp. *speltoides*, when considering either the whole gene or the exon 4 sequences (Table 4). Indeed, the three groups, *Ae. speltoides* ssp. *speltoides*, *Ae. speltoides* ssp. *ligustica* and *Ae. speltoides*, showed very similar Pi values and the phylogenetic analysis further confirmed the above observation. In the NJ tree (Fig. 4), clades are well-supported (bootstrap value $\geq 93\%$) and all *Ae. speltoides* sequences clustered in a single clade, in which ssp. *speltoides* and ssp. *ligustica* were scattered and mixed, but definitively separated from other the *Aegilops* species. Our results are in line with the conclusions of Belyayev and Raskina (2013) showing that ssp. *ligustica* and ssp. *speltoides* were a case of population dimorphism in fruit structure and did not in fact represent two different subspecies, as initially proposed by Zhukovsky (1928).

In the phylogenetic tree (Fig. 4) the sequences of Aegilops species were arranged according to their specific known genome even though the sampling performed of the various diploid and polyploid Aegilops species was somewhat limited and a little unbalanced for experimental reasons. It is worth noting that the two sequences of Ae. sharonensis from the Sitopsis section were found to be sisters of the Ae. speltoides clade and the two sequences of Ae. tauschii clustered together in a separated and well-supported branch. Furthermore, Ae. umbellulata, the only other diploid Aegilops species included in this analysis and donor of the U genome to some of the tetraploid species included, was able to form the highly supported (93%) U genome clade. In this regard, it is reported that the U genome occurring in some polyploid species is very well conserved and similar to the parental diploid one, whereas the second genomes evolved rapidly from their ancestors (Badaeva et al. 2004). Interestingly, the Ae. cylindrica sequence, which has the genomic formula $C^{c}C^{c}D^{c}D^{c}$ (Linc et al. 1999), stood apart and was placed in a single branch, leading us to suppose that this sequence proceeded from the C^c genome, since the D genome of Ae. cylindrica is closely related to the D genomes of both Ae. tauschii and T. aestivum (Caldwell et al. 2004).

Due to the close genetic relationship between the S genome of *Aegilops speltoides* and the B subgenome of wheat (Haider 2013; Marcussen et al. 2014), the three *DRF1* gene sequences isolated from *Ae. speltoides* (*AesDRF1* gene) were aligned with the two *DRF1* gene sequences from the B subgenome of *T. durum* (see the alignment in Online Resource 1) and the comparison appeared very interesting, in particular when considering exon 1, containing a SSR region coding for a variable number of alanines (Di Bianco et al. 2011). We recently found that this region consists

Fig. 4 The optimal Neighbor Joining tree based on 84 sequences of DRF1 exon 4 from Aegilops species (sum of branch length = 0.4504). There was a total of 812 positions in the final dataset. The bootstrap support values (> 50%) are shown close to the node. The species, the NCBI IDs and the simplified genome codes are reported on each branch. Genome clusters are indicated in bold to right of tree. Asterisks denote branches shared with the tree inferred by the Maximum-Likelihood method





Fig. 5 The optimal Neighbor Joining tree based on 102 *DRF1* exon 4 sequences from *Aegilops* and *Triticum* species (sum of branch length = 0.9089). For clarity, the *Aegilops speltoides* group of 68 sequences was graphically collapsed. There was a total of 730 positions in the final dataset. The bootstrap support values (> 50%) are shown close to the node. The species, the NCBI IDs and a single capital letter indicating the genome type

of the repetition of two different patterns, both encoding for alanine, namely GCG and GCC, peculiar to the A and B subgenome copies, respectively, thus representing a specific subgenome signature in *T. durum* (Cantale et al. 2018). The length of the SSR region was also variable and we found it was 21 bp long in the A subgenome and 18 bp in the B subgenome copies, respectively. In *Ae. speltoides*, the SSR consisted of 15 nucleotides in *AeslDRF1*-FJ843102 and *AessDRF1*-FJ858188, and 12 nucleotides in *AeslDRF1*-FJ858187, always displaying the GCG triplet. It is worth noting that the SSR regions of the two pseudogene sequences, isolated from different accessions, showed the GCG triplet for a stretch of 15 nucleotides, but also two further downstream triplets

are reported on each branch. When the genome type was unknown, the simplified genome code was indicated: AB for *T. durum* and *T. dicoccoides* or ABD for *T. aestivum*. Genome clusters are indicated in bold on the right of the tree. Asterisks denote branches shared with the tree inferred by Maximum-Likelihood method

(GCC GCA), both encoding for alanine, with sequences identical to those observed in the B subgenome copies (see Fig. 3). The presence of a variable repeat number, the GCG triplet and the GCC GCA motif places the *Ae. speltoides* SSR sequence in an intermediate position between the A and B subgenomes, but the few sequences containing this region currently available do not make it possible to generalize. On the other hand, it is reported that SSRs show an extremely high rate of reversible variation in the repeat number, which can be functionally significant (Gemayel et al. 2010). Recent literature provides evidence of an evolutionary role for SSRs in transcribed regions as important sources of adaptive genetic variation, as they are involved in a number

of effects such as the regulation of transcription and translation of their own genes, organization of chromatin and morphological variations (Kashi and King 2006; Viera et al. 2016). All things considered, further research into the SSR region of the *DRF1* gene in *Ae. speltoides* looks particularly interesting, as it may represent another possible player involved in the gene regulation.

To widen the comparison with the Triticum genus, the exon 4 sequences from Aegilops, T. aestivum and T. durum were aligned and a phylogenetic tree was inferred (Fig. 5). Variability of a single gene has been used to investigate the phylogenetic relationship among species (Adderley and Sun 2013; Liu et al. 2016), although this task appears particularly challenging in this tribe, and sometimes large incongruities among the trees built from different genes could be observed, due to the complex nature of allopolyploidy, possible introgressions and the involvement of mechanisms such as gene hybridization, gene flow or horizontal gene transfer (Degnan and Rosenberg 2006; Liberles and Dittmar 2008; Escobar et al. 2011). In spite of the above, and also considering that the number of accessions analysed among the Aegilops and Triticum genera was not balanced, the phylogenetic tree inferred showed well-supported clades attributable to the current genomes (S, A, B, D and U genomes). Furthermore, those sequences of which the subgenome was initially not known placed themselves correctly with respect to a genomic designation consistent with their species, so that the association enriches information about them. The topology observed among the A, B and D clades could be symbolized by B (A, D) as the A clade is closely related to the D clade and both are distant from the B clade. Marcussen et al. (2014) found that the above topology is the most frequent in bread wheat when analysing both genes in the whole genome and a large number of trees from different genes. Furthermore, with coalescent-based genome divergence analyses they estimated that Ae. speltoides is a sister to the B genome, T. urartu is a sister to the A genome and Ae. sharonensis and Ae. tauschii are successive sisters to the D genome (Marcussen et al. 2014).

In conclusion, we report a study focused on the isolation and characterization of the *DRF1* gene in *Aegilops speltoides* and other *Aegilops* species. The overall results suggest that there is a high similarity between the B and S genome copies of the *DRF1* gene,

but also some features indicating that both genomes have evolved independently, in line with other observations reporting that B is the most variable among the three subgenomes of current polyploid wheats (Wendel 2000; Petersen et al. 2006). As DRF1 is a key gene in drought response, we intend to continue our studies: the possible involvement of the pseudogene, SSR variability and the DRF1.2 transcript in the gene regulation mechanisms is very interesting and deserves further research. Furthermore, the sequencing of the whole gene in a larger taxon sampling would increase knowledge of DRF1 gene diversity in the Triticeae tribe. The advancement of genomics and the growing availability of wild relative genomes, coupled with the development of new statistical methods for comparative genomics, will make it possible for molecular data to be supplemented by other information that is underutilized in phylogenetic analyses, such as protein structural constraints in specific residues, chromosomal information and population genetics, leading in the end to a better understanding of the evolutionary history of Triticeae. Looking further ahead, the increase in knowledge should provide an important contribution to the widening of crop genetic diversity in support of global food security.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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References

- Adderley S, Sun G (2013) Molecular evolution and nucleotide diversity of nuclear plastid phosphoglycerate kinase (PGK) gene in *Triticeae (Poaceae)*. Gene 533(1):142–148
- Agarwal PK, Agarwal P, Reddy MK, Sopory SK (2006) Role of *DREB* transcription factors in abiotic and biotic stress tolerance in plant. Plant Cell Rep 25(12):1263–1274
- Badaeva ED, Amosova AV, Samatadze TE, Zoshchuk SA, Shostak NG, Chikida NN, Zelenin AV, Raupp WJ, Friebe B, Gill BS (2004) Genome differentiation in *Aegilops*. 4. Evolution of the U-genome cluster. Plant Syst Evol 246:45–76
- Bedre R, Irigoyen S, Petrillo E, Mandadi KK (2019) New era in plant alternative splicing analysis enabled by advances in high-throughput sequencing (hts) technologies. Front Plant Sci 10:740
- Belyayev A, Raskina O (2013) Chromosome evolution in marginal populations of *Aegilops speltoides*: causes and consequences. Ann Bot 111(4):531–538
- Brozynska M, Furtado A, Henry RJ (2016) Genomics of crop wild relatives: expanding the gene pool for crop improvement. Plant Biotech J 14(4):1070–1085
- Caldwell KS, Dvorak J, Lagudah ES, Akhunov E, Luo MC, Wolters P, Powell W (2004) Sequence polymorphism in polyploid wheat and their d-genome diploid ancestor. Genetics 167(2):941–947
- Cantale C, Di Bianco D, Thiyagarajan K, Ammar K, Galeffi P (2018) B genome specific polymorphism in the *TdDRF1* gene is in relationship with grain yield. Planta 247(2):459–469
- Caporale LK (2003) Darwin in the genome: molecular strategies in biological evolution. McGrow-Hill, New York
- Degnan JH, Rosenberg NA (2006) Discordance of species trees with their most likely gene trees. PLoS Genet 2(5):e68
- Di Bianco D, Thiyagarajan K, Latini A, Cantale C, Felici F, Galeffi P (2011) Exploring the genetic diversity of the *DRF1* gene in durum wheat and its wild relatives. Plant Genet Res 9(2):247–250
- Dubrovina AS, Kiseklev KV, Zhuravlev YN (2013) The role of canonical and noncanonical pre-mRNA splicing in plant stress responses. Biomed Res Int. https://doi.org/10.1155/ 2013/264314
- Egawa C, Kobayashi F, Ishibashi M, Nakamura T, Nakamura C, Takumi S (2006) Differential regulation of transcript accumulation and alternative splicing of a *DREB2*

homolog under abiotic stress conditions in common wheat. Genes Genet Syst 81(2):77–91

- Escobar JS, Scornavacca C, Cencil A, Guilhaumon C, Santoni S, Douzery EJP, Ranwez V, Glémin S, David J (2011) Multigenic phylogeny and analysis of tree incongruences in Triticeae (Poaceae). BMC Evol Biol 11:181
- Gemayel R, Vinces MD, LegendreM Verstrepen KJ (2010) Variable tandem repeats accelerate evolution of coding and regulatory sequences. Ann Rev Genet 44:445–477
- Guindon S, Gascuel O (2003) A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst Biol 52(5):696–704
- Haider N (2013) The origin of the B genome of bread wheat (*Triticum aestivum* L.). Russ J Genet 49(3):263–274
- Harlan JR, de Wet JMJ (1971) Toward a rational classification of cultivated plants. Taxon 20(4):509–517
- Huang X, Madan A (1999) CAP3: a DNA sequence assembly program. Genome Res 9(9):868–877
- Kashi Y, King DG (2006) Simple sequence repeats as advantageous mutators in evolution. Trends Genet 22(5):253–259
- Kerby K, Kuspira J (1987) The phylogeny of the polyploid wheats *Triticum aestivum* (bread wheat) and Triticum *turgidum* (macaroni wheat). Genome 29(5):722–737
- Kumar S, Stecher G, Tamura K (2016) MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol Biol Evol 33(7):1870–1874
- Kumar A, Kapoor P, Chunduri V, Sharma S, Garg M (2019) Potential of *Aegilops* sp. for improvement of grain processing and nutritional quality in wheat (*Triticum aestivum*). Front Plant Sci 10:308
- Latini A, Rasi C, Sperandei M, Cantale C, Iannetta M, Dettori M, Ammar K, Galeffi P (2007) Identification of a DREBrelated gene in *Triticum durum* and its expression under water stress conditions. Ann Appl Biol 150(2):187–195
- Latini A, Sperandei M, Cantale C, Arcangeli C, Ammar K, Galeffi P (2013) Variability and expression profile of the *DRF1* gene in four cultivars of durum wheat and one triticale under moderate water stress conditions. Planta 237(4):967–978
- Lefort V, Longueville J-E, Gascuel O (2017) SMS: smart model selection in PhyML. Mol Biol Evol 34(9):2422–2424
- Liberles DA, Dittmar K (2008) Characterizing gene family evolution. Biol Proced Online 10(1):66–73
- Linc G, Friebe BR, Kynast RG, Molnar-Lang M, Köszegi B, Sutka J, Gill BS (1999) Molecular cytogenetic analysis of *Aegilops cylindrica* Host. Genome 42:497–503
- Liu F, Si H, Wang C, Sun G, Zhou E, Chen C, Ma C (2016) Molecular evolution of *Wcor15* gene enhanced our understanding of the origin of A, B and D genomes in *Triticum aestivum*. Sci Rep 6:31706
- Marcussen T, Sandve SR, Heier L, Spannagl M, Pfeifer M, The International Wheat Genome Sequencing Consortium, Jakobsen KS, Wulff BBH, Steuernagel B, Mayer KFX, Olsen O (2014) Ancient hybridizations among the ancestral genomes of bread wheat. Science 345(6194):1250092
- Murray MG, Thompson WF (1980) Rapid isolation of high molecular weight plant DNA. Nucleic Acids Res 8:4321–4325
- Nakashima K, Nobuhisa I, Deshimaru M, Nakai M, Ogawa T, Shimohigashi Y, Fukumaki Y, Hattori M, Sakaki Y,

Hattori S, Ohno M (1995) Accelerated evolution in the protein-coding regions is universal in crotalinae snake venom gland phospholipase A_2 isozyme genes. PNAS 92:5605–5609

- Petersen G, Seberg O, Yde M, Berthelsen K (2006) Phylogenetic relationships of *Triticum* and *Aegilops* and evidence for the origin of the A, B, and D genomes of common wheat (*Triticum aestivum*). Mol Phylogenet Evol 39(1):70–82
- Pont C, Leroy T, Seidel M, Tondelli A, Duchemin W, Armisen D, Lang D, Bustos-Korts D, Goué N, Balfourier F, Molnár-Láng M, Lage J, Kilian B, Özkan H, Waite D, Dyer S, Lettellier T, Alaux M, Wheat and Barley Legacy for Breeding Improvement (WHEALBI) Consortium, Russel J, Keller B, van Eeuwijk F, Spannagl M, Mayer KFX, Waugh R, Stein N, Cattivelli L, Haberer G, Charmet G, Salse J (2019) Tracing the ancestry of modern bread wheats. Nat Genet 51(5):905–911
- Reddy AS, Rogers MF, Richardson DN, Hamilton M, Ben-Hur A (2012) Deciphering the plant splicing code: experimental and computational approaches for predicting alternative splicing and splicing regulatory elements. Front Plant Sci 3:18
- Rozas J, Ferrer-Mata A, Sánchez-DelBarrio JC, Guirao-Rico S, Librado P, Ramos-Onsins SE, Sánchez-Gracia A (2017) DnaSP 6: DNA sequence polymorphism analysis of large datasets. Mol Biol Evol 34(12):3299–3302
- Ruban AS, Badaeva ED (2018) Evolution of the S-genomes in *Triticum-Aegilops* alliance: evidences from chromosome analysis. Front Plant Sci 9:1756
- Salse J, Chagué V, Bolot S, Magdelenat G, Huneau C, Pont C, Belcram H, Couloux A, Gardais S, Evrard A, Segurens B, Charles M, Ravel C, Samain S, Charmet G, Boudet N, Chalhoub B (2008) New insights into the origin of the B genome of hexaploid wheat: evolutionary relationships at the SPA genomic region with the S genome of the diploid relative Aegilops speltoides. BMC Genom 9:555
- Sambrook J, Fritsch EF, Maniatis T (1990) Molecular cloning: a laboratory manual, 2nd edn. Cold Spring Harbor Laboratory Press, Cold Spring Harbor
- Samonte RV, Eichler EE (2002) Segmental duplications and the evolution of the primate genome. Nat Rev Genet 3(1):65–72
- Schneider A, Molnár I, Molnár-Láng M (2008) Utilisation of *Aegilops* (goatgrass) species to widen the genetic diversity of cultivated wheat. Euphytica 163:1–19
- Sen K, Ghosh TC (2013) Pseudogenes and their composers: delving in the 'debris' of human genome. Brief Funct Genom 12(6):536–547
- Sievers F, Wilm A, Dineen DG, Gibson TJ, Karplus K, Li W, Lopez R, McWilliam H, Remmert M, Söding J, Thompson JD, Higgins D (2011) Fast, scalable generation of highquality protein multiple sequence alignments using clustal omega. Mol Syst Biol 7:539

- Soloviev V, Kosarev P, Seledsov I, Vorobyev D (2006) Automatic annotation of eukaryotic genes, pseudogenes and promoters. Genome Biol 7(Suppl 1):S10
- Tatarinova TV, Chekalin E, Nikolsky Y, Bruskin S, Chebotarov D, McNally KL, Alexandrov N (2016) Nucleotide diversity analysis highlights functionally important genomic regions. Sci Rep 6:35730

Tutar Y (2012) Pseudogenes. Comp Funct Genomics 424526

- van Slageren (1994) Wild wheats: a monograph of *Aegilops* L. and *Amblyophyrum* (Jaub & Spach) Eig (Poaceae). ICARDA/Wageningen Agricultural University Papers 94–7
- Vanin EF (1985) Processed pseudogenes: characteristics and evolution. Annu Rev Genet 19:253–272
- Viera MLC, Santini L, Diniz AL, de Freitas MC (2016) Microsatellite markers: what they mean and why they are so useful. Genet Mol Biol 39(3):312–328
- Vinckenbosh N, Dupanloup I, Kaessmann H (2006) Evolutionary fate of retroposed gene copies in the human genome. PNAS 103(9):3220–3225
- Wen Y-Z, Zheng L-L, Qu L-H, Ayala FJ, Lun Z-R (2012) Pseudogenes are not pseudo any more. RNA Biol 9(1):27–32
- Wendel JF (2000) Genome evolution in polyploids. Plant Mol Biol 42:225–249
- Xie J, Li Y, Liu X, Zhao Y, Li B, Ingvarsson PK, Zhanga D (2019) Evolutionary origins of pseudogenes and their association with regulatory sequences in plants. Plant Cell 31:563–578
- Xue GP, Loveridge CW (2004) *HvDRF1* is involved in abscisic acid-mediated gene regulation in barley and produces two forms of AP2 transcriptional activators, interacting preferably with a CT-rich element. Plant J 37(3):326–339
- Yamaguchi-Shinozaki K, Shinozaki K (1994) A novel *cis*-acting element in an Arabidopsis gene is involved in responsiveness to drought, low-temperature, or high-salt stress. Plant Cell 6(2):251–264
- Zhang H, Mittal N, Leamy LJ, Barazani O, Song B-H (2017) Back into the wild—apply untapped genetic diversity of wild relatives for crop improvement. Evol Appl 10:5–24
- Zhukovsky M (1928) A critical systematical survey of the species of the genus *Aegilops* L. Bull Appl Bot Genet Plant Breed 18:417–609
- Zou C, Lehti-Shiu MD, Thibaud-Nissen F, Prakash T, Buell CR, Shiu SH (2009) Evolutionary and expression signatures of pseudogenes in *Arabidopsis* and rice. Plant Physiol 151(1):3–15

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