MIKC type genes of the MADS-box family in wheat: molecular and phylogenetic analysis

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INTRODUCTION

The MADS-box family of transcription factors play key roles in the control of development and signal transduction in eukaryotes (1). The MADS family contains a DNA-binding domain (MADS box) and includes two main lineages, type I and type II, both of which are present in plants, animals and fungi (2). Type II MADS-box proteins of plants, also named as MIKCtype proteins, possess three more domains than Type I MADS-box proteins: intervening (I) domain, keratinlike (K) domain and C-terminal (C) domain. In plants MIKC-type genes are involved in several important developmental processes, such as flower morphogenesis, ovule development, vegetative growth, embryogenesis and fruit formation, and through phylogenetic analysis based on sequence comparison, they have been classified into 13 subfamilies (3). The MIKC transcription factors controlling floral organ identity are the best characterized. Analysis of homeotic floral mutants of Arabidopsis resulted in the formulation of the ABCDE genetic model, which explains how the combined functions and organ-specific expression of five gene classes (A, B, C, D and E) specify the identity of the floral organs (sepals, petals, stamens and carpels), forming the four concentric whorls, and ovules (4,5). The most comprehensive cloning and characterization of MIKC genes of grasses have been carried out in maize and rice (6,7). In this study we extend a detailed characterization of the diversity and complexity of MIKC-type genes to hexaploid wheat.

CLONING AND CHARACTERIZATION OF MIKC-TYPE MADS-BOX GENES IN WHEAT

The 34 available sequences of MIKC-type genes of rice (7) and 37 of *Arabidopsis* (3) were exploited to BLAST search the public databases of wheat ESTs: TIGR wheat gene index database (TaGI, version 10), HarvEST wheat (version 1.13) and NCBI. The 29 identified non-redundant MIKC-type consensus sequences were used as templates for 5' and 3' RACE extensions. Full-length cDNAs of the 29 putative wheat MIKC-type genes from various plant tissues were obtained using primer pairs specific for the very 5' and 3' ends, The products of five independent RT-PCR reactions for each of the 29 primer pairs the five full-length cDNAs exhibited the same sequence, whereas the five clones obtained by each of the remaining 14 primer

Clone		Sequence length (nt)			Protein		Domains (aa)			
	Tot.	5'UTR	3'UTR	ORF	(aa)	5'ext.	М	1	К	С
TaSOC1-1A	1000	84	226	690	230		60	12	101	57
TaSOC1-1B	999	84	228	687	229		60	12	100	57
TaAG-1	1101	18	276	807	269	36	60	13	100	60
TaAG-3A	1071	148	167	756	252	-	60	14	101	77
TaAG-3B	1077	148	167	762	254	-	60	14	101	79
TaSEP-1	1076	15	317	744	248		60	14	97	77
TaSEP-2A	976	75	187	714	238		60	14	98	66
TaSEP-2B	976	75	226	675	225		60	14	98	53
TaAP1-1	1163	144	287	732	244		60	14	100	70
TaAP1-2	IIII	116	194	801	267		60	15	100	92
TaAP1-3	1255	140	293	822	274		60	14	100	100
TaAGL6-A	1144	126	241	777	259		60	13	99	87
TaAGL6-B	1141	126	241	774	258		60	13	99	86
TaAGL6-C	1158	134	250	774	258		60	13	99	86
TaSEP-3A	1226	113	357	756	252		60	14	102	76
TaSEP-3B	1233	113	364	756	252		60	14	102	76
TaSEP-4	1033	67	228	738	246		60	14	102	70
TaAGL12	890	124	94	672	240		60	9	102	50
TaAP3	970	104	179	687	229		60	10	100	59
TaPI-1	1035	217	194	624	208		60	10	100	38
TaPI-2	953	24	302	627	208		60	10	100	39
TaWM16	986	118	280	588	196		60	10	100	25
TaSOC1-2	980	32	158	288	259	33	60	17	100	49
		214	280	708	236		60 60	17		49
TaSEP-5A	1202	214 197	280 281	708	236	•			100	
TaSEP-5B	1186					-	60	14	100	62
TaSEP-6	1028	91	256	681	227		60	14	98	55
TaSOC1-3A	937	144	127	666	222		60	13	99	50
TaSOC1-3B	931	138	127	666	222		60	10	102	50
TaSVP-1A	1282	183	415	684	228		60	9	103	56
TaSVP-1B	1297	198	415	684	228		60	9	103	56
TaSVP-2A	1041	86	277	678	226	-	60	18	95	53
TaSVP-2B	1043	89	276	678	226	-	60	18	95	53
TaGGM13	1155	134	265	756	252	-	60	10	101	81
TaAG-4A	1073	49	259	765	255	-	60	12	102	81
TaAG-4B	1081	49	279	753	251	-	60	12	102	77
TaSVP-3A	1163	187	298	678	226	-	60	20	94	52
TaSVP-3B	1171	191	302	678	226	-	60	20	94	52
TaAG-2A	1141	107	215	819	273	37	60	13	101	62
TaAG-2B	1135	107	200	828	276	37	60	13	101	65
TaAGL17-1	1119	123	276	720	240		60	12	100	68
TaAGL17-2A	1155	117	342	696	232		60	13	100	59
TaAGL17-2B	1164	126	342	696	232		60	13	100	59
TaAGL17-2C	1122	91	341	690	230		60	13	100	57
TaAGL17-3A	1012	40	249	723	241		60	12	99	70
TaAGL17-3B	1013	40	250	723	241		60	12	99	70

Table 1. Characteristics of nucleotide and deduced amino acid sequences of 45 full-length cDNAs of wheat MIKC-type genes.

pairs showed either two or three similar but not identical sequences. A code of Ta, the first two letters of Triticum aestivum was assigned to each sequence, followed by the name of its orthologous gene of Arabidopsis identifying the corresponding MIKC phylogenetic subfamily. Multiple sequences clustering into the same subfamily were designed by an additional number (1-6) and the different clones obtained by the same primer pair were distinguished by an additional letter (A-C) (Table 1). Search for conserved motifs using the Pfam HMMs, InterPro and SMART databases revealed that proteins encoded by the 45 full-length wheat cDNAs showed the modular structure typical of plant MIKC-type MADSbox proteins (Table 1). Four wheat cDNAs showed Nterminal extensions upstream their MADS-box domain: TaAG-1 (36 aa), TaSOC1-2 (33 aa), TaAG-2A (37 aa) and TaAG-2B (37 aa). On the basis of sequence homology three of them (TaAG-1, TaAG-2A and TaAG-2B) were assigned to the subfamily including the gene Agamous of Arabidospsis, whose members are characterized by N-terminal extensions. The presence of the N-terminal extension is less obvious for TaSOC1-2,

which belongs to the SOC1/AGL20 subfamily, whose other members do not possess it; therefore its origin and functional meaning would require further studies. Multiple cDNAs cloned from independent amplifications with the same primer pair showed high identity (over 90%) at both nucleotide and amino acid levels, most probably because they derived from transcripts of MADS box genes located in homoeologous chromosomes. Most mutations detected in the coding region of the putative homoeologous cDNA sequences consisted of nucleotide substitutions, with prevalence of synonymous substitutions, and of short in-frame insertions/deletions (indels) of three. six and nine nucleotides, whereas the 5' and 3' untranslated regions showed higher rates of base substitutions and of indels of variable length (data not shown). The most remarkable differences observed among putative homoeologous sequences consisted of two frameshift mutations, in the C-terminal regions of the clone pairs TaSEP-2A/TaSEP-2B and TaAG-4A/TaAG-4B. Previous studies in other plant species have shown that frameshift mutations in the C-terminal domain of specific ancestral MADS-box MIKC-type genes may have contributed to their structural and functional divergence (8).

Southern analyses showed the presence in hexaploid wheat of three homoeologous copies for each of the 29 identified MIKC-type sequence (23 of them in Fig. 1), indicating that the genome of *T. aestivum* contains at least 87 (29x3) type II MIKC MADS-box genes. This assumption was further supported by aneuploid analysis of six sequences assigned by phylogenetic analysis to the *SEP/AGL2* subfamily. The three homoeologous sequences of both TaSEP-1 and TaSEP-2 were located in chromosome arms 4AS, 4BL and 4DL (Fig. 2a and 2d), those of TaSEP3 and TaSEP-5 in the long arms of group 5 chromosomes (Fig. 2b and 2e) and those of TaSEP-4 and TaSEP-6 in the short arms of group 7 chromosomes (Fig. 2c and 2f).

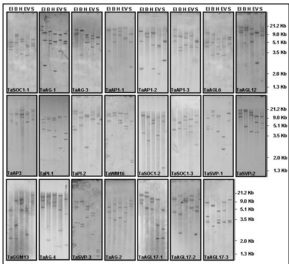
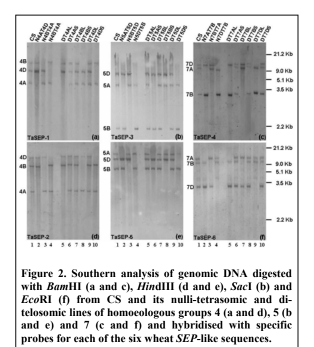


Figure 1. Southern analysis of the hexaploid wheat cv Chinese Spring (CS) genomic DNA digested with *Eco*RI (EI), *Bam*HI (B), *Hin*dIII (H), *Eco*RV (EV) and *SacI* (S) and hybridised with specific probes of 23 MIKC-type sequences of wheat cloned in this study.



The chromosome arm locations of the six SEP-like genes of wheat were compatible with the map locations of their orthologous genes of rice and maize. In fact syntenic relationships, and sometimes high collinearities, have been shown between the chromosomal regions where the SEP-like genes of rice and maize are located and the 4, 5 and 7 chromosome homoeologous groups wherein the wheat genes were located (Table 2).

 Table 2. Chromosome arm locations and syntenic relationships between the SEP-like genes of wheat, rice and maize.

SEP-like	Chromosome arm location					
wheat gene	Wheat	Rice orthologous*	Maize orthologous*			
TaSEP-1 TaSEP-2 TaSEP-3 TaSEP-4 TaSEP-5 TaSEP-6	4AS, 4BL, 4DL 4AS, 4BL, 4DL 5AL, 5BL, 5DL 7AS, 7BS, 7DS 5AL, 5BL, 5DL 7AS, 7BS, 7DS	3S (OsMADSI) 3S (OsMADSI) 9L (OsMADS8) 8L (OsMADS7) 3L (OsMADS34) 6S (OsMADS5)	1S (ZMM14), 9L (ZMM8) 1S (ZMM14), 9L (ZMM8) 7L (ZMM7) 1L (ZMM6) 1L (ZMM27) 9S (ZMM3)			

*The chromosomal locations of *SEP*-like genes from rice and maize have been reported by Lee et al. (7) and Munster et al. (6).

PHYLOGENETIC ANALYSIS

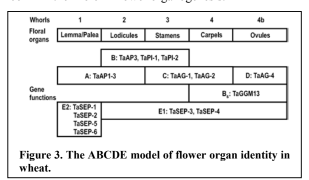
The evolutionary relationships between cloned wheat cDNAs and other known MIKC-type genes of plants were studied by the phylogeny reconstruction of 125 amino acid sequences deduced from 33 genes from *Arabidopsis*, 31 from rice, 32 from maize and 29 from wheat. The phylogenetic analysis revealed that 28 of the 29 non-homoeologous cDNA sequences of wheat were assigned to 11 of the 13 known plant subclasses of the MIKC-type gene family, whereas TA-WM16 and its putative ortholog *OsMADS32* of rice formed a separate clade. More extensive analyses would be needed to verify whether the *Arabidopsis* orthologs of these genes have been lost or are present only in monocot or grass genomes. The absence in wheat of sequences

homologous to FLC- and AGL15-like genes is consistent with previous studies, which found these genes only in Brassicaceae species, but not in other plants, even not in the sequenced genome of rice (3,7). These data indicate that at least 11 subfamilies of MIKC genes are common to both monocot and dicot species and that at least many different MIKC genes were carried by the common ancestor of monocots and dicots. Moreover, in agreement with previous phylogenetic studies, our results demonstrate that several gene duplications occurred in the lineage that originated grass species, preceding the separation of the lineages that led to extant maize, rice and wheat. Our phylogenetic analysis identified 27 minimal clades containing putative orthologs from both wheat and rice, 22 of them included also maize genes. The lack of maize orthologs in five minimal clades (belonging to the subfamilies AGL12, AGL17, SOC1/AGL20 and OsMADS32) could be due either to incomplete MIKC gene sampling in that species or to their deletion after maize speciation. On the basis of these lines of evidence at least 22 (probably 27) MIKC genes were carried by the common ancestor of maize, rice and wheat, which diverged about 50-70 MYA. This represents a significant increase in comparison to the 11 MIKC-type genes that would have been present about 200 MYA in the common ancestor of monocots and eudicots. Most likely the increase of MIKC genes in the progenitor of grass species was caused by multiple duplications followed by gene diversification and could have been a key factor in the evolution of the complex inflorescences common to the species belonging to the Poaceae family.

THE ABCDE MODEL OF FLOWER ORGAN IDENTITY IN WHEAT

The expression of the wheat cDNAs corresponding to the genes of the homeotic classes, A, B, Bs, C, D, and E was analysed by RT-PCR, quantitative real time RT-PCR and northern blot hybridisation in 18 different plant tissues and floral organs of Chinese Spring (data not shown). The expression patterns of these genes were compared with those of functionally characterised MADS-box genes of Arabidopsis and monocot species, such as rice and maize, in the same phylogenetic clades. Potential functions of the wheat genes corresponding to the cloned MADS-box sequences were assumed on the basis of sequence similarities and comparable expression patterns. The morphology of the grass flowers is quite different from that of dicots. However, molecular genetic studies of maize and rice suggest that lemma and palea are flower organs homologous to sepals of dicots and lodicules are modified petals, therefore the ABC model can effectively be extended to grass species (9). On the basis of these considerations and of the phylogenetic and expression analyses of the cloned cDNAs, we propose an adaptation of the "ABCDE model" of Arabidopsis to wheat. The wheat MIKC genes encoding the products, which most likely provide the A, B, C, D and E functions, are shown in Fig. 3. The additional function B sister (Bs) provided by TaGGM13

was introduced and might be required together with the C and D functions for specifying the identities of carpels and ovules. The wheat class E (SEP-like) genes were split into the functional groups E1 and E2. The E1 activity would be provided by TaSEP-3 and TaSEP-4, whose redundant functions would be required for the development of lodicules, stamens and carpel. The E2 activity would be accomplished by TaSEP-1 and TaSEP-2. which are orthologs of the rice gene LHS1, and by TaSEP-5 and TaSEP-6. Theaction of these genes would be required specifically, together with that of the class A genes, for determining the lemma/palea identity. The expression patterns of TaSEP-1, TaSEP-2 and TaSEP-5 indicate that the corresponding genes would be involved also in the development of the reproductive organs of wheat flowers. Although the existence of functional A genes in grass species remains controversial, based on our expression analyses TaAP1-3 would likely be the putative wheat functional equivalent of AP1 in Arabidopsis. The model presented here is mainly intended as a working hypothesis. Functional characterisation of the isolated wheat MIKC-type genes using loss- and gain-of-function mutants, analyses of transcript expression by in situ hybridisation and identification of protein-protein interaction patterns by yeast two-hybrid experiments will be required to confirm their role in flower organogenesis.



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