

# Genome-Wide ORFeome Cloning and Analysis of Arabidopsis Transcription Factor Genes<sup>1[w]</sup>

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Here, we report our effort in generating an ORFeome collection for the Arabidopsis transcription factor (TF) genes. In total, ORFeome clones representing 1,282 Arabidopsis TF genes have been obtained in the Gateway high throughput cloning pENTR vector, including 411 genes whose annotation lack cDNA support. All the ORFeome inserts have also been mobilized into a yeast expression destination vector, with an estimated 85% rate of expressing the respective proteins. Sequence analysis of these clones revealed that 34 of them did not match with either the reported cDNAs or current predicted open-reading-frame sequences. Among those, novel alternative splicing of TF gene transcripts is responsible for the observed differences in at least five genes. However, those alternative splicing events do not appear to be differentially regulated among distinct Arabidopsis tissues examined. Lastly, expression of those TF genes in 17 distinct Arabidopsis organ types and the cultured cells was profiled using a 70-mer oligo microarray.

Transcription factors (TFs) play critical roles in all aspects of a higher plant's life cycle. It is the programmed and regulated interactions between TFs and genomic DNA that bring a genome to its life and define many of its functional features (Grandori et al., 2000; Dimova et al., 2003; Kohler et al., 2003). An initial analysis indicated that Arabidopsis has at least 1,533 TF genes (approximately 6% of the coding capacity of its genome) that belong to more than 30 different families, each possessing a highly conserved and characteristic region recognized as the DNA-binding domain (Riechmann et al., 2000, 2002). Besides the DNA-binding domain in different TF families, there are conserved sequence motifs that help to further classify the TF genes into subgroups (Hosoda et al.,

2002; Heim et al., 2003; Parenicova et al., 2003; Toledo-Ortiz et al., 2003). Among the Arabidopsis TFs, about 45% are plant-specific, whereas the rest share DNA-binding domains common to other eukaryotes (Riechmann et al., 2000, 2002). Arabidopsis has several large TF families, each with more than 100 members. Those include the MYB, bHLH, MADS, and AP2/EREBP family of TFs (Riechmann and Ratcliffe, 2000; Hosoda et al., 2002; Heim et al., 2003; Parenicova et al., 2003; Toledo-Ortiz et al., 2003).

Although extensive studies have been carried out for functional analysis of individual TFs, the function of only a small fraction of these TFs has been revealed so far (Riechmann, 2002). The complete sequence of the Arabidopsis genome (The Arabidopsis Genome Initiative, 2000) made it possible not only to approach the function of TFs on a genomic scale, but to examine the transcriptional network and cascade involved. Transcriptional networks or cascades are common features in controlling Arabidopsis development and its response to various environmental challenges (Shinozaki and Yamaguchi-Shinozaki, 2000; Riechmann, 2002). In these regards, microarray profiling has become an important approach to analyze TF genes at the genome

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level. For example, profiling of 402 Arabidopsis TF genes under various environmental conditions revealed that 74 of the bacterial pathogen-responsive TF genes were also involved in salicylic acid, jasmonic acid, and ethylene signaling (Chen et al., 2002). Among the 43 genes that were activated during senescence, 28 were also induced by other stress treatments, indicating the possibility of extensive overlaps of cellular events downstream of the same TFs (Chen et al., 2002). Recently, a PCR-amplified fragment-based microarray containing approximately 95% of all TF genes from Arabidopsis was generated and used to reveal genome-wide differential expression of TF genes between white light- and dark-grown seedlings (Jiao et al., 2003; Ma et al., 2003). Although microarray profiling is a power tool in revealing TF gene expression patterns and in some instances the temporal and functional interdependency among TF gene expression, it alone often cannot provide sufficient knowledge of TF function.

Further functional analysis of TF genes requires a careful examination of the encoded proteins and their interaction in the cells. A prerequisite for genome-wide analysis of TF genes at the protein level is a collection of cDNA clones with intact open-reading-frames (ORFs). Unfortunately, the initial identification of TF genes in the Arabidopsis genome sequence was carried out mainly by *ab initio* gene predictions, sequence homology comparisons, motif analysis, and other nonexperimental methods (The Arabidopsis Genome Initiative, 2000). Dramatic progress in Arabidopsis genome annotation was achieved by expressed sequence tag and full-length cDNA analysis (Seki et al., 2002) and tiling-path oligo microarray studies (Yamada et al., 2003). However, transcriptional activity was demonstrated for only about 70% of the Arabidopsis genes by microarray analyses and currently available full-length cDNA clones cover only about 41% of Arabidopsis genes (Yamada et al., 2003). Thus there is an urgent need for higher coverage of ORFome clones (cDNA clones containing full-length ORFs) of TF genes for the Arabidopsis community.

Here we report our genome wide effort in generating Arabidopsis TF ORFome clones, which succeeded in covering 1,282 unique Arabidopsis TF genes. In the process, sequence analysis of our ORFome clones allowed us to correct a number of errors in the annotation of these genes. Further, comprehensive expression profiles of those TF genes in the Arabidopsis life cycle were conducted. This ORFome clone collection has been deposited in the Arabidopsis stock center and is available to the research community for in-depth functional analysis of Arabidopsis TF genes.

#### GENERATION OF ORFome cDNA CLONES FOR 1282 ARABIDOPSIS TF GENES

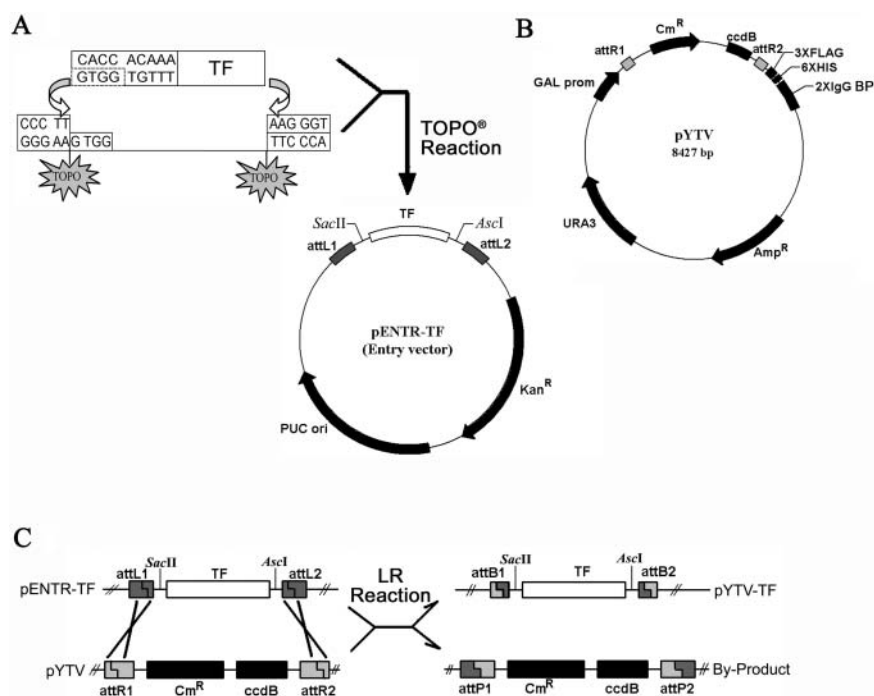
Through searching MIPS database using previously described InterPro and GenBank accessions as family

identifier (Riechmann et al., 2000), we obtained a collection of 1,581 TF genes from the Arabidopsis genome at the start of this study (see "Materials and Methods"). Table I summarizes TF gene numbers in each family as reported by a prior study (Riechmann et al., 2000) and the number of TF genes for which we were able to retrieve sequence information at the time. Note that a total of about 20 TF genes that could not be placed in any known families were reported as "others." Further, about 30 TF genes in a newly

**Table I.** The number of genes encoding putative TFs in the Arabidopsis genome

TF Families	InterPro or GenBank Acc. Access	A	B	C
ABI3/VP1	CAA48241	14	17	4
ALFIN-like	AAA20093	7	7	7
AP2/EREBP	IPR001471	144	118	142
ARF	AAC49751	23	23	11
ARID	IPR001606	4	5	3
AS2		0	0	30
AUX/IAA	AAC39440	26	28	26
bHLH	IPR001092	139	108	81
BZIP	IPR001871	81	82	54
C2C2 (Zn)-co-like	A56133	33	32	24
C2C2 (Zn)-dof	CAA66600	37	35	32
C2C2 (Zn)-gata	IPR000679	28	22	33
C2C2 (Zn)-yabby	AAD30526	6	5	3
C2H2 (Zn)	IPR000822	105	152	92
C3H-type1 (Zn)	IPR000232	17	46	32
C3H-type (Zn)	CAA65242	16	26	10
CCAAT	A26771/P13434/Q02516	36	36	34
	AAB51375			
CPP (Zn)	CAA09028	8	8	5
E2F/DP	O00716/Q64163	8	8	1
EIL	AAC49750	6	6	5
GARP	AAD55941/BAA74528	56	53	24
GRAS	AAB06318	32	32	26
HB	IPR001356	89	83	56
HMG-box	IPR000910	10	13	10
HSF	IPR000232	26	16	12
JUMONJI	T30254	9	8	4
LFY	AAA32826	1	1	0
MADS	IPR002100	82	91	70
MYB	IPR001005/IPR000818	190	258	243
NAC	BAB10725	109	106	93
Nin-like	CAB61243	15	14	2
PCG		4	4	2
SBP	CAB56581	16	16	14
TCP	AAC26786	25	21	19
Trihelix	S39484	28	21	1
TUB	IPR000007	11	9	7
WRKY (Zn)	S72443	72	70	54
Others		20	20	16
Totals		1,533	1,581	1,282

Column A, number of putative TFs reported by Riechmann et al. (2000); Column B, number of TFs whose sequence and annotation can be retrieved from MIPS database at the time we started this effort (December, 2000); Column C, number of TFs whose ORFs were cloned in the current work. All genes are counted only once, even in some genes that belong to more than one family based on their multiple signature motifs.



**Figure 1.** Schematic diagrams showing strategies of high-throughput cloning of blunt-ended PCR amplified TF gene ORFeome products. A, Cloning of ORFeome product into the pENTR-TOPO vector. The PCR products containing individual TF gene ORFs were directionally cloned into pENTR-TOPO vectors using the TOPO DNA recombination reaction facilitated by topoisomerase I attached to one of the vector strands. B, Map of pYTV. This yeast expression vector was modified based on vector pDEST 52 (Invitrogen). The fused tags at the C-terminal of TF ORF include three copies of Flag, six His amino acids, and two copies of IgG-binding motif from protein A. Those tags should enable tandem affinity purification (TAP) of the TF proteins (Rigaut et al., 1999). C, Generation of yeast expression cassette using pENTR clones and pYTV vector. Recombination between different pENTR-TFs and pYTV vectors were carried out in the presence of Gateway LR Clonase enzyme mix. *Ascl* and *SacII* were designed for determination of the insert size. For further details about this gateway system, please refer to the Invitrogen manual ([www.invitrogen.com](http://www.invitrogen.com)).

defined AS family were included as well. Based on the retrieved sequence and annotation information, primer pairs were designed for all TF genes based on a recombinant cloning strategy and used to amplify ORF regions of mRNAs through reverse transcription (RT) and PCR (see Fig. 1 and "Materials and Methods"). A variety of Arabidopsis tissue types and treatments (see "Materials and Methods") were employed to isolate total RNA to serve as template for RT-PCR. In total, we succeeded in obtaining ORFeome clones for 1,282 TF genes in the pENTR TOPO vector system. Of the

1,282 TF genes that had ORFeome clones, 411 had no matches in current cDNA collections (data not shown).

As expected, most of the ORFeome clones matched to the existing cDNA sequences or gene annotation based on single read sequencing from both ends of each clone. Our sequence analysis also revealed differences in 39 clones. Among those, 34 ORFeome clones were completely sequenced and their differences confirmed from either reported cDNA sequences or annotation in public databases (Tables II and III). Table II summarizes those 15 TF genes that have prior

**Table II.** Fifteen of our TF genes showed different in ORF sequences from previously reported cDNA submission

Families	Locus ID	Old Acc.	New Acc.	ORF in bp		Differences (New VS old)
				Old	New	
C2H2 (Zn)	At1g03840	BT006209	AJ630476	1,521	1,515	6 bp deleted at 151 <sup>1</sup>
bHLH	At1g26260	RAFL 16-92-I23	AJ630483	1,173	1,020	153 bp deleted at 178
C2H2 (Zn)	At1g34790	ATH318491	AJ630477	912	909	3 bp deleted at 219
C2H2 (Zn)	At1g72050	RAFL 09-22-M23	AJ630478	975	1,239	9 bp omitted and 273 bp added from codon 1
SBP	At1g76580	RAFL 08-16-H24	AJ630503	1,467	1,510	43 bp added at 1,440
SBP	At2g42200	RAFL 04-13-I11	AJ628864	1,128	1,137	9 bp added at 824
C2H2 (Zn)	At3g13810	RAFL 09-12-L04	AJ630504	1,542	1,551	9 bp added at 175 <sup>2</sup>
MYB	At3g46590	RAFL 07-12-A13	AJ630475	939	1,659	3 bp deleted at 81, 6 bp deleted and 729 bp added at 933
bHLH	At3g61950	RAFL 19-75-M14	At3g61950	924	1,077	153 bp added from codon 1
HMG	At4g11080	AY133687	AJ630485	1,341	1,353	12 bp added at 90
WRKY (Zn)	At4g26640	RAFL 07-10-M14	AJ630479	1,458	1,674	12 bp omitted and 228 bp added from codon 1
bHLH	At4g29930	RAFL 17-41-A21	AJ630482	765	792	89 bp deleted and 116 bp added at 676
WRKY (Zn)	At5g22570	RAFL 21-21-O16	AJ630480	870	867	3 bp deleted at 677
C3H-type2 (Zn)	At5g46730	RAFL 15-50-L08	AJ630502	873	813	60 bp deleted at 586
C2H2 (Zn)	At5g54630	RAFL 09-11-B05	AJ630501	1,419	593	826 bp deleted at 568

<sup>1</sup>Nucleotide positions counted from the A of translation initiation codon.

<sup>2</sup>This 9-bp sequence was not found in the Arabidopsis genome database.

**Table III.** Nineteen clones that were different in their ORF from those predicted annotations in the genome database

Families	Locus ID	New GenBank Accessions	ORF in bp		Differences (Ours VS MIPS) <sup>1</sup>
			MIPS	Ours	
MYB	At1g14600	AJ630486	252	768	33 bp deleted and 549 bp added at 219
ABI3/VP1	At1g28300	AJ630496	1,092	1,089	3 bp deleted at 1,011
WRKY (Zn)	At1g29280	AJ630494	780	762	18 bp omitted from codon 1
AP2/EREBP	At1g79700	AJ580379	927	912	6 bp deleted at 171, 9 bp deleted at 250
C2H2 (Zn)	At3g01030	AJ630491	1,062	1,116	54 bp added at 633
MYB	At3g10000	AJ630487	1,491	1,446	13 bp added and 58 bp deleted at 368
CPP (Zn)	At3g16160	AJ630495	1,083	1,107	3 bp deleted at 137, 6 bp added at 405, 16 bp added and 4 bp deleted at 630, 9 bp added at 716
AP2-EREBP	At3g23230	AJ580377	369	420	51 bp added at 293
ABI3/VP1	At3g26790	AJ630497	942	933	9 bp deleted at 715
C2H2 (Zn)	At3g27970	AJ630492	1,065	1,074	9 bp added at 315
MYB	At4g12670	AJ630488	1,563	1,500	63 bp deleted at 1,232
MYB	At4g39160	AJ630489	1,638	1,806	168 bp added at 49
C2H2 (Zn)	At5g03150	AJ630493	1,506	1,512	6 bp added at 187
MYB	At5g17780	AJ630490	1,260	1,254	6 bp deleted at 418
bHLH	At5g38860	AJ630499	1,062	897	165 bp deleted at 665
AP2/EREBP	At5g50080	AJ580378	663	663	27 bp deleted at 75, 27 bp added at 202
bHLH	At5g53210	AJ630498	885	1,095	210 bp added at 583.
C2H2 (Zn)	At5g54360	AJ630500	813	763	9 bp deleted at 681, 24 bp deleted at 697, 17 bp deleted at 750
bHLH	At5g61270	AJ630484	804	837	19 bp added at 159, 14 bp added at 245

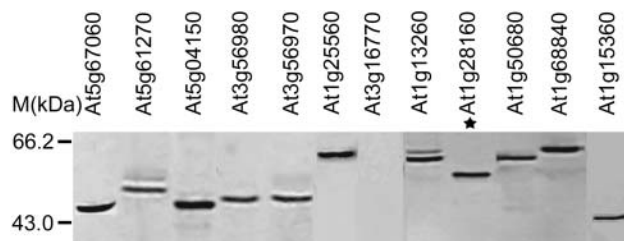
<sup>1</sup>All nucleotide positions started from A of the translation initiation codon.

deposited cDNA sequences but nevertheless show clear sequence discrepancy between our ORFeome clones and the cDNA. Table III summarizes those 19 Arabidopsis TF genes that had predicted annotation without experimental support. In these cases, we were able to correct inaccuracy in the gene annotation using our ORFeome clones. All those 34 ORFeome clone sequences were submitted to GenBank and their corresponding accession numbers are listed in Table II and III.

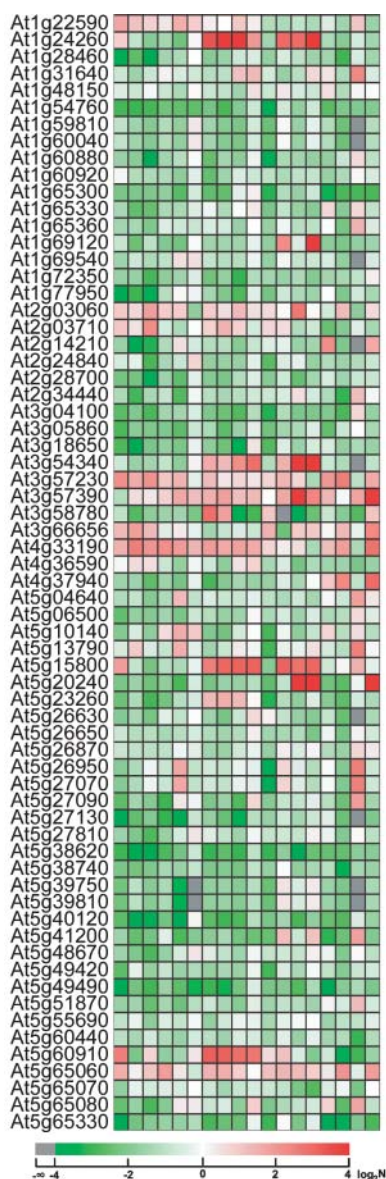
#### EXPRESSION OF REPRESENTATIVE TF ORFeome CLONES IN YEAST

As a way to confirm the intactness of ORFeome clones and to test the feasibility of high-throughput expression of TF proteins, all the ORFeome clone inserts in our collection were transferred into a yeast expression vector (pYTV; see Fig. 1) for expression analysis in yeast. About 300 representative ORFeome clones in pYTV vector were selected from different TF gene families and their expression in yeast was examined via protein-blotting analyses. Using antibodies against a His-tag fused to ORFeome inserts in the pYTV vector, our results indicated that up to 85% of ORFeome clones expressed TF proteins above the detection limit. Examination of the protein size in SDS-PAGE indicated that about 90% of the expressed proteins were of expected  $M_r$  (data not shown). For example, as illustrated in Figure 2, of the yeast protein extracts from 12 distinct TF ORFeomes, 10 of them

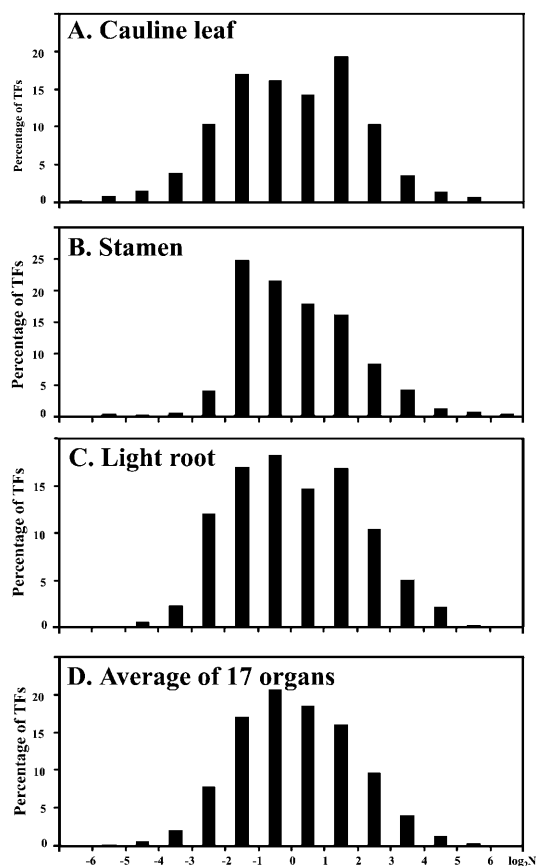
produced strong protein blot signals that match with the calculated protein sizes, while 1 protein migrated significantly slower than predicted (Fig. 2). Protein extract from 1 clone failed to produce a detectable amount of protein (Fig. 2). These results largely confirmed the intactness of our ORFeome clones. The small amount of clones (<10%) that failed to produce proteins with the expected size (with most migrating slower) suggest that the proteins encoded by these ORFeome clones may have unusual conformations or promiscuous interactions with other proteins in yeast such that they migrated in larger size than expected. For those failed to be detected, the proteins might be unstable or expressed at very low levels in yeast.



**Figure 2.** Protein-blot analysis of Arabidopsis TF protein expression in *Saccharomyces cerevisiae*. Total proteins were fractionated by SDS-PAGE, probed with 1  $\mu$ g/mL monoclonal antipolyhistidine antibody, and visualized after incubating with goat anti-mouse AP-conjugated secondary antibody. \* marks the protein that migrated significantly slower than its predicted  $M_r$ .



**Figure 3.** Expression profiles of 66 Arabidopsis MADS family TF genes in 17 different Arabidopsis organs and cultured cells. Locus ID is listed at the left of the column, and only those MADS box genes whose ORFeome clones are presented in our collection were analyzed. All comparisons were done against the absolute median point obtained for the respective organ. The ratio value of the normalized signal intensity in each organ for a given gene (see "Materials and Methods") relative to the median value from that organ was first calculated. Then this ratio was subjected to logarithmic ( $\log_2$ ) transformation, with the resulting value as indicator of relative expression level among organ type. Therefore, a value of zero indicates an expression level equal to the median of all gene expression in that organ, a positive value indicates an expression level higher than the media, and a negative value indicates an expression level lower than the median. The 18 lanes are as follows: a, cauline leaf; b, light cotyledon; c, rosette leaf; d, dark cotyledon; e, dark hypocotyls; f, light hypocotyl; g, pistil 1 d after pollination; h, pistil 1 d before pollination; i, Silique 3 d after pollination; j, silique 8 d after pollination; k, stem; l, sepal; m, stamen; n, petal; o, dark root; p, light root; q, germinating seed; and r, cultured cells.



**Figure 4.** Distribution of expression abundance for the 858 TF genes. A, Distribution of TF gene expression levels in cauline leaf. B, Distribution of TF gene levels in stamen. C, Distribution of TF gene levels in light root. D, An averaged distribution of TF gene expression levels in all 17 organs. A similar method was used to calculate relative expression as described in the legend for Figure 3 and the  $\log_2$  values for the ratio for the normalized expression levels with the median are shown in the x axis.

#### EXPRESSION PROFILING OF ARABIDOPSIS TF GENES

An Arabidopsis 70-mer oligo microarray covering more than 25,000 Arabidopsis genes were used for organ-specific expression analysis (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data). In this array, 1,222 of the 1,282 ORFeome genes were present. The profiling data of those 1,222 TF genes from the 17 representative Arabidopsis organs and suspension cultured cells were extracted from the above-mentioned data set and allow us to estimate the relative expression abundance for each transcript in different organs. As the detailed characterization of the AP2/EREBP and MYB families of TF genes will be reported separately, the expression patterns for the remaining 858 cloned TF genes are summarized in this report. The expression patterns of MADS family of TF genes among the 17 organs and cultured cells were illustrated in Figure 3, while the expression patterns for the entire

858 TF genes are shown in an appendix figure available at [www.plantphysiol.org](http://www.plantphysiol.org). One notable feature of the TF gene expression profile is that a vast majority of the TF genes exhibited organ specific expression patterns. On the other hand, some notable exceptions are present. For example, four genes (At5g65670, At2g22430, At2g18160, and At1g30970) were found to express at quite higher levels in all organs and cultured cells tested in the current work. The relative expression levels of the 858 TF genes follow similar distribution in most organ types (Fig. 4) with a general pattern not much different from the total gene expression level distribution in each organ type (data not shown).

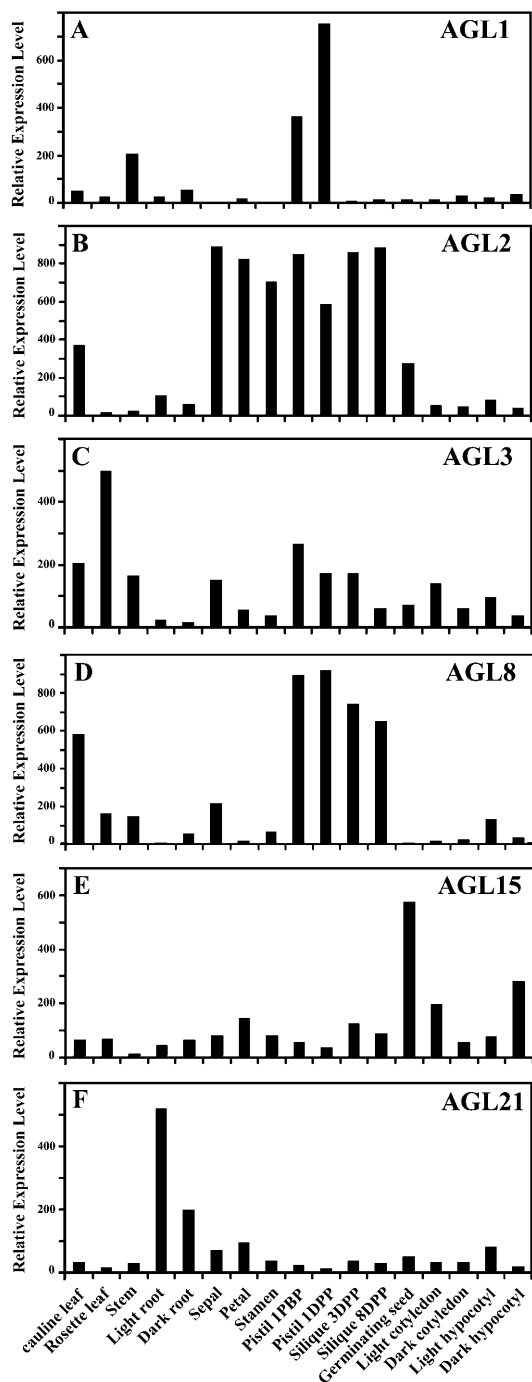
As described in a previous analysis (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data), close comparisons between the known expression patterns of several well-characterized TF genes and the microarray result is a valuable mean to validate our microarray data. The genes examined in that work (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data) included the *PISTILLATA* (Goto and Meyerowitz, 1994; Honma and Goto, 2000), *APETALA1* (Honma and Goto, 2001; Ng and Yanofsky, 2001), and *LATERAL ORGAN BOUNDARIES* genes (Shuai et al., 2002). The data from the microarray analysis all exhibited organ or tissue expression patterns that are consistent with prior studies (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data). In this

report, we extend this comparative analysis to 14 known MADS box TF genes that have expression data derived from in situ or northern-blot analysis. We found that all 14 TF genes exhibited largely similar expression patterns between our microarray analysis and previous reported nonmicroarray data (Table IV). The detailed expression patterns from six of those MADS box genes are illustrated in Figure 5. For example, *AGL1* is specifically expressed in particular regions of the gynoecium and ovule (Fig. 5A). *AGL2* transcript is very abundant in the primordia of all four floral organs: sepals, petals, stamens, and carpels. The *AGL2* transcript remains abundant in each organ during morphological differentiation but diminishes as each organ undergoes the final maturation phase of development (Fig. 5B). *AGL3* is expressed in all aerial organs but roots (Fig. 5C). *AGL8* RNA does not accumulate during vegetative growth; it accumulates to high levels in the inflorescence apical meristem as well as in the inflorescence stem and cauline leaves (Fig. 5D). *AGL15* is preferentially expressed in embryos and accumulates significantly in germinating seedlings (Fig. 5E). *AGL21* is highly expressed in roots and in developing embryos (Fig. 5F). The only possibly minor exception is the *FLC* gene, which exhibited a small difference in relative expression level in inflorescence organs from our microarray analysis compared to prior studies. In a prior northern analysis, it was shown that *FLC* expression was not detectable in inflorescence organs (Michaels and Amasino, 1999),

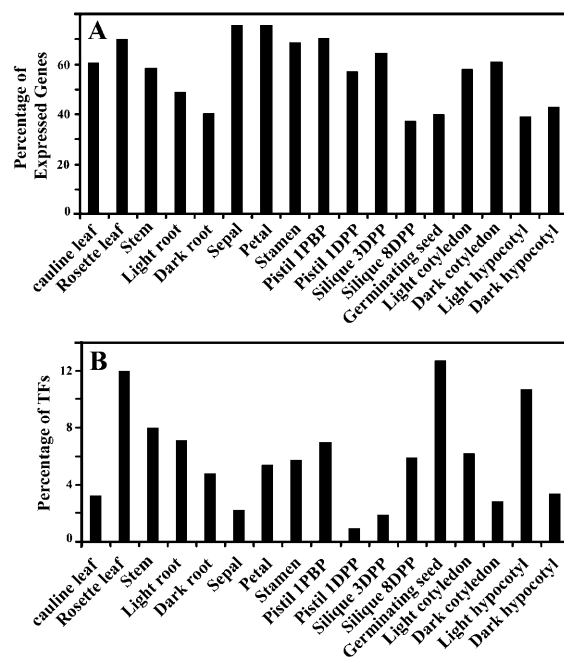
**Table IV.** Comparison of our microarray results with prior studies for representative MADS box genes

Locus ID	Gene name	Reference	Comment
At3g58780	AGL1	Flanagan et al. (1996) Ma et al. (1991) Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g15800	AGL2 SEP1	Flanagan and Ma (1994) Ma et al. (1991) Kofuji et al. (2003)	Microarray data are consistent with reported results
At2g03710	AGL3	Huang et al. (1995) Ma et al. (1991) Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g60910	AGL8 FUL	Mandel and Yanofsky (1995) Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g13790	AGL15	Rounsley et al. (1995) Kofuji et al. (2003)	Microarray data are consistent with reported results
At4g37940	AGL21	Burgeff et al. (2002) Kofuji et al. (2003)	Microarray data are consistent with reported results
At1g24260	AGL9 SEP3	Kofuji et al. (2003)	Microarray data are consistent with reported results
At2g03060	AGL30	Kofuji et al. (2003)	Microarray data are consistent with reported results
At3g57230	AGL16	Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g20240	PI	Goto and Meyerowitz (1994) Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g23260	AGL32	Kofuji et al. (2003)	Microarray data are consistent with reported results
At1g69120	AGL7/AP1	Mandel et al. (1992) Kofuji et al. (2003)	Microarray data are consistent with reported results
At3g54340	AP3	Jack et al. (1992) Kofuji et al. (2003)	Microarray data are consistent with reported results
At5g10140	AGL25 FLC	Michaels et al. (1999) Kofuji et al. (2003)	Possible minor discrepancy in inflorescence or floral organs

while our microarray result indicated that *FLC* expressed at levels slightly below the median expression level in floral organs. This minor discrepancy could be due to the not exactly same organ types used or cross-hybridization in the oligo microarray. Overall,



**Figure 5.** The relative expression levels of the 6 well-characterized MADS box TF genes in 17 organ types as determined by our microarray analysis. A, AGL1 (At3g58780); B, AGL2 (At5g15800); C, AGL3 (At2g03710); D, AGL8 (At5g60910); E, AGL15 (At5g13790); and F, AGL21 (At4g37940). The observed expression patterns from our microarray studies are consistent with the previously reported results (see Table IV).



**Figure 6.** Summary of TF gene expression characteristics among Arabidopsis organ types. A, The percentage of TF genes expressed in different organs. Abbreviations: DPP, day post pollination; DBP, day before pollination. A total 858 TF genes (see text) are analyzed, and 61% (cauline leaf), 70% (rosette leaf), 58% (stem), 49% (light root), 41% (dark root), 76% (sepal), 76% (petal), 79% (stamen), 71% (pistil 1DBP), 57% (pistil 1DPP), 65% (silique 3DPP), 37% (silique 8DPP), 40% (germinating seed), 58% (light cotyledon), 61% (dark cotyledon), 39% (light hypocotyl), 43% (dark hypocotyl) were detected expression respectively. B, Distribution of TF genes in their highest expression level among 17 different tissues. Among the 858 TF genes analyzed, 3% (cauline leaf), 12% (rosette leaf), 8% (stem), 7% (light root), 5% (dark root), 2% (sepal), 5% (petal), 6% (stamen), 7% (pistil 1DBP), 1% (pistil 1DPP), 2% (silique 3DPP), 6% (silique 8DPP), 13% (germinating seed), 6% (light cotyledon), 3% (dark cotyledon), 11% (light hypocotyl), and 3% (dark hypocotyl) are expressed at their highest level in the indicated organ types, respectively.

the results suggest that whole genome oligo microarray is a valid approach to determine specific gene expression patterns. As knowledge of expression patterns of a TF gene often offers the initial clues needed in dissecting its biological function, these results should provide useful information for further functional studies.

We also examined the number (and percentage) of the 858 TF genes whose expression can be detected experimentally in each organ type and in any of the organs examined. This analysis revealed that the expression for 831 (97%) out of the 858 genes can be detected in at least one of the 17 organs or cultured cells examined. This result confirms that the vast majority of known and predicted TFs are expressed during Arabidopsis development; while the percentage of TF genes expressed in each organ types varied from 37.4% (silique 8 d-post-pollination) to 75.8% (petal; see Fig. 6A). We also calculated the numbers of genes exhibiting highest relative expression levels

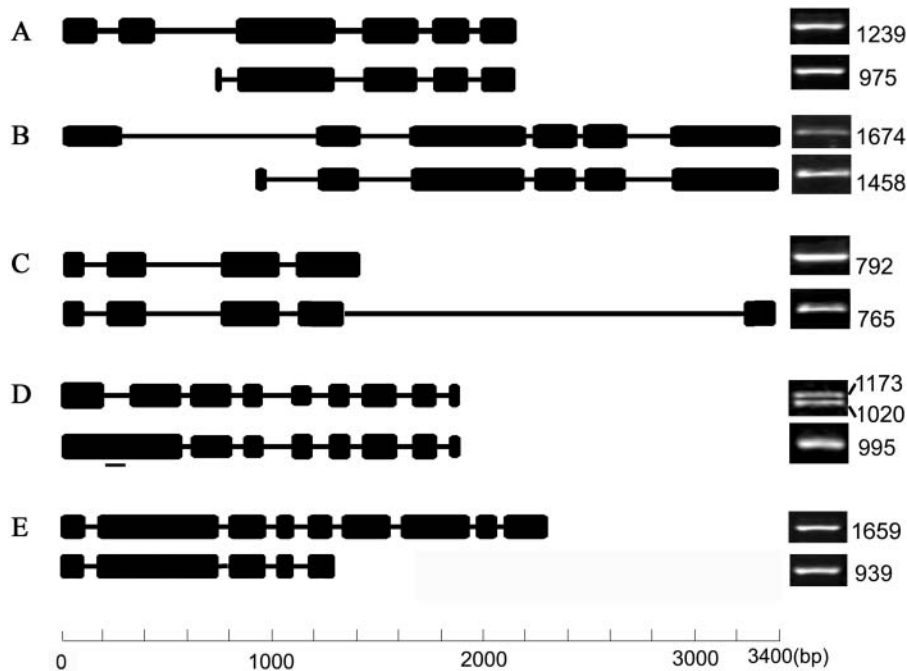


in each organ types. As shown in Figure 6B, the percentage of the highest expressed TF genes in each organ type varies from organ to organ. Vegetative organs have large numbers of TF genes that have the highest expression level. This may be consistent with the fact that vegetative organs are where most of metabolism activities reside. About 20% TF genes exhibit highest expression levels in flower organs, which may hint at their special roles in flower development and reproduction. The germinating seed has the highest number of TFs, with highest expression among all organs examined here. It is interesting to note that the germinating seed has a very low percentage of the total TF genes with detectable expression (Fig. 6A). This result indicated that during seed germination, a relatively large fraction of genes turned on highly are TFs. This is consistent with the fact that those early expressed genes will initiate the developmental and metabolic processes to follow.

#### ALTERNATIVE SPLICING OF FIVE TF GENES

For the 34 TF genes where our ORFeome sequences differ from prior cDNA sequence or predicted gene

annotation (Tables II and III), we examined whether alternative splicing contributes to the observed differences. Indeed, we were able to confirm that alternative splicing variants were present for five TF genes (At1g26260, At1g72050, At3g46590, At4g26640, and At4g29930) by RT-PCR (Fig. 7) and were responsible for the observed differences in their ORFeome clone sequences. Different mRNA forms of At1g72050 and At4g26640 appear to be generated by using extra or different exons located at the very 5' end, while At4g29930 and At3g46590 include alternative exons at the 3' of the mature RNAs. In At1g26260, one form of the mRNAs simply contains the first intron that is spliced out in the other form of transcript. In the case of At1g72050, the previously reported mature transcript contains 5 exons with a 975 bp ORF (GenBank accession nos. AY054225 and AY066042), and its encoded protein possesses one C2H2-type Zinc-finger domain (PSSIM-Id: 20248) plus a partial domain (PSSIM-Id: 21389) that is incomplete in both ends. By using two extra exons at the 5' end of the transcript, the new ORF predicted from our cloned mRNA species is 1,239 bp in length and has two C2H2-type Zinc-finger domains, a scenario resembling a previously reported alternative splicing event



**Figure 7.** Alternative splicing of five Arabidopsis TF genes. A diagram of the exon/intron structures of the alternative spliced transcripts of At1g72050 (A), At4g26640 (B), At4g29930 (C), At1g26260 (D), and At3g46590 (E) is shown on the left. Sequence-specific primers were synthesized for each form of the mature RNA as outlined on the left side. For each gene, our version of gene structure and PCR product is placed on top, while the gene structure and PCR products based on prior information is placed on bottom. The levels of the two mature RNA molecules from Arabidopsis plant samples were analyzed by RT-PCR, and only one representative result is shown as ethidium bromide-stained DNA band, with the length of full-length ORF marked to the right. At the bottom is a scale that indicates the lengths of introns (represented by thin lines) and exons (thick black boxes) in all the genes compared. The small bar to the lower transcript shown in D designates the position of the forward primer used to amplify the RAFL reported version of cDNA since the 5' end of these two mature RNA molecules were identical and hence only the 995 bp band represented a partial ORF. However, the PCR primer pair for the 1,173 bp cDNA (based on the full-length ORF of RAFL cDNA) can amplify both the 1,020 bp cDNA band (our new version) and the 1,173 bp cDNA, so that doublet bands were observed.



for the rice *Myb7* gene (Magaraggia et al., 1997). In that case, one of the two transcripts that encodes a partial DNA-binding domain was known to act as a repressor to switch off the expression of its target genes, while the other served as a transcription activator (Magaraggia et al., 1997). However, in the other four genes, alternative splicing variants do not affect the DNA-binding domains.

In all five cases, the typical GT-AG binucleotide splicing junctions were observed in the alternatively spliced transcripts. To test if alternative splicing of these genes is developmentally regulated, we designed specific PCR primer pairs for the two alternative spliced transcripts of each of those five genes and used semiquantitative RT-PCR to examine the presence and abundance of those alternative mRNAs in selected Arabidopsis tissue samples (Fig. 7). The alternative spliced transcripts for each of those five genes were present at similar abundance in all tissue types tested (data not shown), suggesting that alternative splicing of these genes is constitutive and is not regulated by developmental or environmental conditions tested.

## MATERIALS AND METHODS

### Plant Materials

Arabidopsis (ecotype Columbia) plants were grown in fully automated growth chambers (Conviro, Canada) under 16 h light illumination each 24 h period. Plants were maintained at 23°C during the light period and 21°C during the dark period.

To provide additional RNA samples to cover those TF genes that may not be expressed under normal growth conditions, Arabidopsis plants at 6 to 8 rosette stages were subjected to the following 8 specific treatments and were used for total RNA isolation: (1) NaCl treatment, whole pots were submerged in 300 mM NaCl for 8 h. (2) Heat shock (heat), plants were preconditioned at 37°C for 2 h before being transferred to 45°C for another 2 h. (3) UV treatment, plants were radiated with UV light (100 J m<sup>-2</sup>) for 6 h. (4) Water depletion treatment, entire plants were uprooted, placed on filter papers, and allowed to dry for 6 h. (5) Ethylene treatment, plants were placed in a closed jar containing 100 ppm C<sub>2</sub>H<sub>4</sub> for 24 h. (6) Cold treatment, plants were placed in a 4°C cold room for 8 h. (7) Wound treatment, rosette leaves were cut into approximately 5 mm strips and were left in the growth chamber for 8 h before being used for RNA isolation. (8) Dark adaptation, plants were placed in darkness for 48 h before being harvested for RNA isolation.

### Identification of Arabidopsis Transcription Factor Genes

All known and predicted TF genes were selected from the MIPS Arabidopsis genome database (<http://www.mips.biochem.mpg.de/proj/thal/db/index.html>) as of December 21, 2000. Each gene was identified by its chromosome locus ID (e.g. At5g61270). The MIPS Arabidopsis database August 17, 2003 update was used as our final reference for gene annotation.

### Isolation and Cloning of ORFeome into pENTR and Expression Vectors

Total RNA was isolated from pooled Arabidopsis plant samples harvested from the 6 to 8 rosette leaves before bolting and plants a week after flowering using the RNeasy plant mini kit (QIAGEN, Germany) and was quantified at 260 nm with a spectrophotometer. This RNA sample was used as a generic initial template for RT-PCR. For any TF genes that were not able to be cloned from those generic RNA samples, RNA samples from specific treated Arabidopsis seedlings (see "Plant Materials" section) were used as an alternative template for RT-PCR amplification.

Three μg of total RNA sample was reverse transcribed using SuperScript First-Strand Synthesis System for RT-PCR (Invitrogen, Carlsbad, CA) in a total volume of 20 μL. Primers for ubiquitin amplification (forward: 5'-GGTGCTAAGAAGAGGAAGAAT-3' and reverse: 5'-CTCCTCTTTCTGGTAAACGT-3') were added as the internal control together with gene-specific primers. PCR was performed using *Pfu* polymerases (Sangon, China). For tissue-specific expression analyses, different plant materials harvested at indicated stages were used. PCR products were purified with a gel extraction kit (CLONTECH Laboratories, Palo Alto, CA), cloned into pENTR/D/TOPO vector (Invitrogen), and verified by sequencing using M13 primers. Primers for different TF genes were designed using information obtained from the Arabidopsis genome. The forward primer contained the sequence 5'-CAC-CACAAA-3' at the 5' end. The CACC base paired with the overhang sequence, GTGG, in pENTR TOPO vector (Fig. 1A). The yeast expression vector, pYTV, was a modified version of the pDEST 52 (Invitrogen). The original tag was removed and was replaced by 3XFLAG, 6Xhis, and a 3C cleavage and 2XlgG binding protein added as C-terminal tags to facilitate purification of the fusion protein (Fig. 1B). To clone the gene of interest in frame with C-terminal tags present in the pYTV, the reverse primer was designed in such a way that the stop codon in the target gene was deleted in the final PCR product for ORF amplification for initial cloning into pENTR TOPO vector (Fig. 1C).

### Protein-Blot Analysis of TF Proteins in Yeast

Total proteins extracted from 50 to 75 μL saturated yeast cells expressing target genes were fractionated by SDS-PAGE. Each gel was probed with 1 μg/mL monoclonal antipolyhistidine antibody (R&D Systems, Minneapolis) and visualized after incubating with goat anti-mouse AP-conjugated secondary antibody (Promega, Madison, WI).

### Expression Profile Analysis using Microarray

Gene-specific 70-mer oligos were designed based on Arabidopsis genome annotation data available on February 20, 2002 by Qiagen ([http://omad.qiagen.com/download/genelist/arabidopsis\\_V1\\_384.prn](http://omad.qiagen.com/download/genelist/arabidopsis_V1_384.prn)), and the microarray slide was printed at Yale University as described (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data). The signal was scanned at 532 nm (Cy3) and 635 nm (Cy5) wavelengths with an Axon GenePix 4000B scanner (Axon, Foster City, CA) at 5-nm resolution and quantified with Axon GenePix Pro 3.0 image analysis. The intensity of different organs was normalized by equalizing the median value of all gene intensities from each organ. The normalized intensity value for each gene was considered its relative expression level (L.G. Ma, N. Sun, X.G. Liu, Y.L. Jiao, H.Y. Zhao, and X.W. Deng, unpublished data).

Sequence data from this article have been deposited with the EMBL/GenBank data libraries under accession numbers listed in Tables II and III.

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