Isolation and characterization of a new mannose-binding lectin gene from *Taxus media*

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In this paper, we report the cloning and characterization of the first mannose-binding lectin gene from a gymnosperm plant species, *Taxus media*. The full-length cDNA of *T. media* agglutinin (TMA) consisted of 676 bp and contained a 432 bp open reading frame (ORF) encoding a 144 amino acid protein. Comparative analysis showed that TMA had high homology with many previously reported plant mannose-binding lectins and that *tma* encoded a precursor lectin with a 26-aa signal peptide. Molecular modelling revealed that TMA was a new mannose-binding lectin with three typical mannose-binding boxes like lectins from species of angiosperms. Tissue expression pattern analyses revealed that *tma* is expressed in a tissue-specific manner in leaves and stems, but not in fruits and roots. Phylogenetic tree analyses showed that TMA belonged to the structurally and evolutionarily closely related monocot mannose-binding lectin superfamily. This study provides useful information to understand the molecular evolution of plant lectins.

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1. Introduction

Plant lectins or agglutinins, widespread throughout the plant kingdom, are considered as an extended group of proteins which, according to a recently updated definition, comprise all plant proteins possessing at least one noncatalytic domain that binds reversibly to specific monoor oligosaccharides (Van Damme *et al* 1998). On the basis of recent advances in biochemistry and molecular cloning and structural analysis, virtually all currently known plant

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Abbreviations used: AAA, Allium ascalonicum agglutinin; ACA, Allium cepa agglutinin; ACPA, Ananas comosus (pineapple) agglutinin; AKA, Amorphophallus konjac agglutinin; APA, Allium porrum agglutinin; ASA, Allium sativum agglutinin; AUA, Allium ursinum agglutinin; AVA, Amaryllis vittata agglutinin; CAA, Crinum asiaticum agglutinin; CTAB, cetyltrimethylammonium bromide; EHA, Epipactis helleborine agglutinin; GEAFP, Gastrodia elata antifungal protein; GNA, Galanthus nivalis agglutinin; LOA, Listera ovata agglutinin; ORF, open reading frame; PCR, polymerase chain reaction; pI, isoelectric point; RACE, rapid amplification of cDNA ends; RT-PCR, reverse transcriptase-polymerase chain reaction; TDA, Typhonium divaricatum agglutinin; TMA, Taxus media agglutinin; UTR, untrslated region; ZCA, Zephyrathes candida agglutinin; ZGA, Zephyrathes grandiflora agglutinin.

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lectins can be classified into seven families of structurally and evolutionarily-related proteins (Van Damme et al 1998), and that most of these belong to four groups of evolutionarily related proteins: legume lectins, chitin-binding lectins, monocot mannose-binding lectins, and type 2 ribosome-inactivating proteins, among which monocot mannose-binding lectins have attracted greater attention because of their great application values in biological and biomedical research; such as in the isolation of mannosecontaining glyconjugates, and in their potent inhibitory effect on human and animal retroviruses, including HIV (Barre et al 1996). In addition, most monocot mannosebinding lectins play an important role in the plant's defense against different kinds of plant-eating organisms; which is due to their recognition of high-mannose type glycans of plant predators (Powell et al 1993; Peumans and Van Damme 1995; Tang et al 2001; Sun et al 2002; Lin et al 2003; Yao et al 2003a). Up to now, monocot mannosebinding lectins have been cloned from seven families of angiosperms including Amaryllidaceae, Araceae, Alliaceae, Orchidaceae, Liliaceae, Iridaceae and Bromeliaceae (Van Damme et al 1991, 1994, 1996, 2000; Smeets et al 1997; Neuteboom et al 2002; Chai et al 2003; Fei et al 2003; Kai et al 2003; Yao et al 2003b; Zhao et al 2003a,b), among which lectins from Amaryllidaceae species have been extensively studied and well documented (Van Damme et al 1992, 1998; Kai et al 2003; Pang et al 2003; Zhao et al 2003a).

However, little is known about the possible origin and molecular evolution of the carbohydrate-binding domains of plant lectins from modern flowering plants with little gene sequence information about plant lectins outside higher flowering plants (Peumans et al 2002). Until now, there have been no reports on molecular cloning of lectin genes from gymnosperms, including family Taxaceae, therefore the molecular evolution relationship of the plant lectins between flowering plants and gymnosperms is unclear. To investigate whether similar mannose-binding lectins also exist in gymnosperms such as Taxaceae, we employed a relatively simple cloning strategy (rapid amplification of cDNA ends, RACE) to successfully isolate a novel lectin gene from Taxus media, which did not belong to any previously reported families containing mannosebinding lectins in the classical taxonomical range above. T. media contains high content of anti-cancer drug Taxol and is a kind of valuable medicinal plant of family Taxaceae in gymnosperms. Up to now, there is no report on the molecular cloning of lectin gene from T. media.

In this paper, we report for the first time the molecular cloning and characterization of a novel mannose-binding lectin gene from *T. media*. The cloned gene is also the first mannose-binding lectin gene isolated from gymnosperm species. The evolutionary relationship among lectins from gymnosperm and angiosperm species was also studied.

2. Materials and methods

2.1 Plant material

T. media plants, provided by Professor Feng Tan at Southwest Normal University in China, were grown in pots in the greenhouse under standard conditions. All tissue materials (1 g) including leaves, fruits, stems and roots from *T. media* plants were frozen separately in liquid nitrogen for RNA extraction.

2.2 Total RNA isolation

Because of the high concentration of secondary products such as phenolic components and polysaccharides in *T. media* tissues, the cetyltrimethylammonium bromide (CTAB) based RNA isolation method (Jaakola *et al* 2001) was adopted in the total RNA extraction. The quality and concentration of the extracted RNA were checked by agarose gel electrophoresis and spectrophotometer (DU-640, Beckman, USA) analysis, respectively. The RNA samples were stored at - 80°C prior to RACE and reverse transcriptasepolymerase chain reaction (RT-PCR) analysis.

2.3 Molecular cloning of tma full-length cDNA

Total RNA (5 µg) was used to synthesize the first strand cDNA (3'-ready cDNA) using SMARTTM RACE cDNA Amplification Kit (Clontech Laboratories Inc., USA) with the 3' RACE CDS Primer A (5'-AAGCAGTGGTATCAA-CGCAGAGTAC $(T)_{30}N_{-1}N-3'$ provided by the kit. The 3' RACE was performed with the Universal Primer A Mix (UPM, long: 5'-CTAATACGACTCACTATAGGG-CAAGCAGTGGTATCAACGCAGAGT; short: 5'-CTAA-TACGACTCACTATAGGGC-3') as the reverse primer and degenerate primer (F2) (5'-ATGCAG(C/G/T)A(G/C)GAC (T/G)GCAACCTG-3') as the forward primer that was designed and synthesized corresponding to the conserved regions [MQ(G/E/Y/D)D(C/G)NL] of previously reported plant lectins (Kai et al 2003; Pang et al 2003; Zhao et al 2003a). PCR was conducted in a total volume of 50 μ l (containing 2 µl cDNA 20 µmol of F2 20 µmol of UPM, 10 µmol dNTPs, 1 × Ex PCR buffer and 5U Ex Taq Polymerase) following the protocol: cDNA was denatured at 94°C for 3 min followed by 35 cycles of amplification (94°C for 30 s, 58°C for 30 s, 72°C for 90 s) and by 10 min at 72°C. The amplified product was purified and cloned into pGEM-T Easy cloning vector (Promega, WI, USA), and transformed into Escherichria coli DH5a. Based on the colour reaction using Xgal-IPTG System and PCR identification, the positive clones were picked out and sequenced by ABI 377 Sequencer (Perkin-Elmer, USA).

The first strand cDNA (5'-ready cDNA) synthesis in 5' RACE was performed according to the manual of the SMARTTM RACE cDNA Amplification Kit (Clontech Laboratories Inc., USA) using the 5'-RACE CDS Primer [5'- $(T)_{25}N_{-1}N$ -3'] provided by the kit. Based on the sequence of the 3' RACE product, the complementary reverse gene specific primer R2 (5'-GCCGTCATTCTGCATACTCAAC-3') was designed and synthesized. The 5' RACE-PCR was carried out using primers R2 and UPM under the following condition: the template (the 5'-ready cDNA) was denatured at 94°C for 2 min followed by 35 cycles of amplification (94°C for 50 s, 56°C for 50 s and 72°C for 90 s) and by 7 min at 72°C. The PCR product was purified and cloned into the pGEM-T Easy cloning vector (Promega, WI, USA) followed by sequencing.

By assembling the sequences of the 3' RACE and 5' RACE products, the full-length cDNA sequence of *tma* was deduced, and was subsequently amplified via PCR using a pair of primers F1 (5'-GGCAATTCTCAATCG-CCATTATA-3') and UPM, which was repeated for three times. The amplified full-length cDNA of *tma* was used for molecular characterization such as sequence homology, signal peptide prediction, presentation of conserved motifs, molecular modelling and phylogenetic tree reconstruction.

2.4 Computer analyses

Bioinformatic-associated analyses of TMA were performed using programs of PSI-Blast, SOPMA, CLUSTAL W1·82 and Swiss-Model. TMA and other plant lectins retrieved from GenBank were aligned with CLUSTAL W (1·82) using default parameters and the phylogenetic tree was constructed using neighbour-joining method (Thompson *et al* 1994).

2.5 Tissue expression analyses

In order to investigate the expression pattern of *tma* in different tissues including leaves, fruits, stems and roots of T. media, semi-quantitative One-step RT-PCR was carried out according to the manufacturer' instruction (Takara, Japan). Aliquots of total RNA (0.5 µg) extracted from leaves, fruits, stems and roots of T. media, individually were used as templates in one-step RT-PCR reaction with the forward primer KF1 (5'-ATGGGAGAATCATCTG-TAAC-3') and reverse primer KR1 (5'-TCAGAGAAG-using one-step RNA PCR kit. Meanwhile, the RT-PCR reaction for the housekeeping gene (actin gene) using specific primers actF (5'-GTGACAATGGAACTGGAA-TGG-3') and actR (5'-AGACGGAGGATAGCGTGAGG-3') designed according to the conserved regions of plant actin genes was performed as an internal control to estimate whether equal amounts of RNA among samples were used in

semi-quantitative RT-PCR (Kai *et al* 2004). Amplifications were performed under the following condition: 50°C for 30 min and 94°C for 2 min followed by 25 cycles of amplification (94°C for 30 s, 60°C for 30 s and 72°C for 2 min). The amplified products were separated on a 1% agarose gel and analysed with Gene analysis software package (Gene company, USA).

3. Results

3.1 Cloning of the full-length cDNA of tma

Based on sequences of the conserved regions of mannosebinding lectins from other plant species, a degenerate primer (F2) was designed and used for the amplification of 3'-end of T. media agglutinin cDNA (tma). A single fragment of about 480 bp was amplified using primers F2 and UPM, and a 3' untranslated region (UTR) of 109 bp was found downstream from the stop codon in the amplified sequence. Complementary reverse-specific primer R2 was designed and used for the cloning of the 5'-end sequence of tma by 5' RACE based on the sequence of 3' RACE fragment. A 400 bp fragment was amplified in which a 5' UTR of 62 bp was found upstream of the first ATG codon. Based on the sequences of the 3' and 5' RACE products, the full-length cDNA fragment was deduced and amplified by PCR using a pair of primers F1 and UPM. The cloned full-length cDNA of tma was 676 bp and contained a 432 bp ORF encoding a protein of 144 amino acids, with an isoelectric point of 9.48 and calculated molecular weight of 15,166 Da (figure 1). The richest amino acid in the deduced TMA was Ala (13% by frequency), followed by Leu (11%), Ser (10%), Val (8%), Gly (8%), and Asn (7%). Acidic and basic amino acids constituted 4% and 9% of the polypeptide, respectively. Nineteen per cent of the total amino acids were charged and the per centages of polar and hydrophobic amino acids were 34% and 40%, respectively.

3.2 Sequence analyses of TMA

Sequence comparison by performing Blast Search in GenBank database (http://www.ncbi.nih.gov) revealed that TMA had high homology with many other mannosebinding lectins such as Zephyrathes grandiflora agglutinin (ZGA), Zephyrathes candida agglutinin (ZCA), Epipactis helleborine agglutinin (EHA), Listera ovata agglutinin (LOA), Amaryllis vittata agglutinin (AVA), Galanthus nivalis agglutinin (GNA) and Crinum asiaticum agglutinin (CAA), suggesting that TMA belonged to the monocot mannose-binding lectin superfamily. Furthermore, TMA also showed high homology to Gastrodia elata antifungal protein (GEAFP). On the amino acid level, TMA was 50%, 51%, 50%, 50%, 49%, 46%, 47% and 49% identical to ZGA, ZCA, EHA, LOA, AVA, GNA, GEAFP and CAA, respectively, and was 61%, 60%, 63%, 66%, 61%, 61%, 59% and 60% similar to ZGA, ZCA, EHA, LOA, AVA, GNA, GEAFP and CAA respectively (figure 2). Signal peptide of TMA was predicted by the signal peptide prediction software (http://www.cbs.dtu.dk/services/signalP)

1										g	gca	att	ctc	aat	cgcc	
18	at	tat	atc	aga	tca	gca	gct	tag	cct	cgc	taa	cag	atc	aaa	cata	
63	at	<i>e</i> gg	aga	atc	atc	tgt	aac	tac	agc	cgc	cat	tgt	agc	gct	ggta	
	М	G	Ε	S	S	۷	T	T	A	A	I	۷	A	L	۷	
108	at	att	act	tac	gtt	tgc	aaa	tcc	ttg	cto	ato	aaa	atc	tgt	tctc	
	Ι	L	L	T	F	A	N	Ρ	C	S	S	Κ	S	۷	L	
153	aa	aag	cgg	gga	atc	act	ggc	ggc	tgg	aga	gto	tct	gca	ata	cgct	
	K	S	G	Ε	S	L	A	A	G	Ε	S	L	Q	Y	A	
198	${\tt caatatatattggtaatgcaaggcgactgcaacctggttttgtat}$															
	Q	Y	Ι	L	۷	М	<u>Q</u>	G	D	С	<u>N</u>	L	۷	L	<u>Y</u>	
243	gc	aaa	caa	ggt	gaa	agt	gtt	gtg	ggc	gtc	aag	gac	aaa	tgg	aaaa	
	A	Ν	Κ	۷	Κ	۷	L	W	A	S	R	Т	N	G	K	
288	ggcggcgccgcctcctgcaagttgagtatgcagaatgacggcaat															
	G	G	A	A	S	С	Κ	L	S	М	<u>Q</u>	N	D	G	<u>N</u>	
333	tt	ggt	gat	ata	tgc	ggc	gac	cac	acc	tgt	atg	ggc	tag	tag	aacc	
	Ł	۷	Ι	<u>Y</u>	A	A	T	Т	Ρ	۷	W	A	S	R	T	
378	tc	cag	agc	ctt	tgc	ttc	cta	caa	gct	caa	tct	tca	ggg	cga	tggt	
	S	R	A	F	A	S	Y	Κ	L	N	L	<u>Q</u>	G	D	G	
423	aa	tgt	tgt	tat	tta	tgg	gcc	gtc	tgg	agc	tat	ttg	ggc	tac	taat	
	N	۷	۷	Ι	<u>Y</u> ,	G	Ρ	S	G	A	I	W	A	Т	N	
468	aca	agc	tca	gaa	caa	gaa	aaa	act	tct	c <u><i>tg</i></u>	<u>a</u> ac	ttt	ctg	gtg	gggg	
	Т	A	Q	Ν	Κ	К	Κ	L	L	*						
513	aa	ott	gtt	ttt	tag	ata	cta	cag	gag	tta	acg	atc	cag	taa	cttg	
558	gc	aca	ctt	ctt	gac	gtg	tta	ata	ata	gga	aga	tac	aat	ctt	cttc	
603	cg	tta	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaa	aaaa	
~ * ~																

Figure 1. The full-length cDNA sequence and deduced amino acid sequence of *T. media* agglutinin (TMA). The start codon (atg) is in italics and the stop codon (tga) is in italics and underlined. The conserved mannose-binding sites (QDNY) are underlined. Asterisk mark (*) represents that stop codon can not be translated into any amino acid.

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and the result showed that *tma* encoded a precursor lectin with a signal peptide with the cleavage site being between Ser^{26} (S) and Lys^{27} (K) (before KSVL), which was very similar to some mannose-binding lectins from angiosperm species such as *G. nivalis*, and *Z. grandiflora* (Van Damme *et al* 1992; Kai *et al* 2003).

NCBI conserved domain search results suggested that TMA belonged to mannose-binding lectin superfamily and contained consensus sequence motif (QDNY) which was involved in alpha-D-mannose recognition (figure 2). The conserved motifs were often conservative during evolution, while some variations on un-conserved domain can form the molecular foundation for the diversity of the structures and functions.

The secondary structure of TMA was analysed by SOPMA (Geourjon and Deléage 1995) and the result showed that the putative TMA peptide contained 21% of alpha helices, 32% of extended strands, 9% of beta turns and 38% of random coils (figure 3A). The random coil and extended strands constituted interlaced domain of the main part of the secondary structure.

3.3 Molecular modelling of TMA

Swiss-Model structure prediction for the three-dimensional structure (Guex and Peitisch 1997) revealed that TMA had similar structure to those of many other mannosebinding lectins and also had three specific conservative mannose-binding sites with the same amino acid sequences (QDNY) symmetrically arranged on TMA molecular (figure 3B), in consistent with many other mannose-binding lectins, such as ZGA, ZCA and GNA (Van Damme et al 1991; Kai et al 2003; Pang et al 2003; Zhao et al 2003a,b), indicating TMA is a mannose-specific lectin. The three conserved mannose-binding sites may play an important role in sugar-binding activity and thus are preserved in evolution. As shown in figure 3B, it was very noteworthy that **b**-sheets occurred predominantly and the overall folding was typically built from **b**-sheets connected by turns and loops, which created very tight structural scaffolds. Molecular modelling provides new evidence for an evolutionary relationship among the plant lectins. Purification of TMA protein from T. media and X-ray crystal structure analysis will further elucidate the detailed structure of TMA in the future.

3.4 Molecular evolution analyses

To investigate the evolutionary relationship among different plant lectins, a phylogenetic tree was constructed based on the deduced amino acid sequences of TMA and other plant lectins. As shown in dendrogram (figure 4), all the monocot mannose-binding lectins were grouped into four

families such as Taxaceae, Araceae, Amaryllidaceae, Orchidaceae and Bromeliaceae. All the lectins from family Alliaceae including *Allium cepa* agglutinin (ACA), *Allium ascalonicum* agglutinin (AAA), *Allium sativum* agglutinin (ASA), *Allium porrum* agglutinin (APA) and *Allium*

big clusters. The first cluster comprised TMA, ZGA, CAA, LOA, EHA, *Typhonium divaricatum* agglutinin (TDA), GEAFP, *Amorphophallus konjac* agglutinin (AKA) and *Ananas comosus* (pineapple) agglutinin (ACPA), which seemed to be made up of some taxonomically unrelated

ZGA	MTTVSINVLLLATTLGLLMTTGARAANVLQSGQQLGTGGSLDYAAYNFRMQADCN 55
CAA	MARNYILLPAAIFGLLLTS-CMAANVLYSGQALSTGASLSFAAYQFVMQADCN 52
LOA	-VSFLSMSTILLVSTMALMSLLATPVSASDRLNAGQSLGTGGSLALGGYIFIIQGDCN 57
EHA	MASFLSLRTILLLSTAALMSLLATPVSAADRLNAGQSLGTGGSLALGGYIFKIQADCN 58
GEAFP	MAASASTAVILFFAVTTVMSLSAIPAFASDRLNSGHQLDTGGSLAQGGYLFIIQNDCN 58
TMA	MGESSVTTAAIVALVILLTFANPCSSKSVLKSGESLAAGESLQYAQYILVMQGDCN 56
ZCA	MTRITFL-VVAAIFLGVITPSCLSVCDSILYSGETLSAGQSLNNGPYTFIMQEDCN 55
AVA	MAKTSFLIVVAAIVLGVITPSCLSDNILYSGETLSAGESLNYGPYTFIMQQDCN 54
GNA	TSLLILATIFLGVITPSCLSDNIMYSGETLSTGEFLNYGSYVFIMQEDCN 50
Consensus	GLGLY QDCN
ZGA	LVLYDNG-NAVWASGTNGRGSGCTCRMQYDGNLVVYTNA-NRAVWASNTNRGTG-SYQ 110
CAA	LVLYDNG-NPVWASGTNGRGSGCQCRMQYDGNLVVYTSG-NRAVWASNTNRGTG-RYQ 107
LOA	LVLYDNN-RAVWASGTNGRGSNCILSMORDGNLVIYS-S-GRAIWASNTNRONG-NYY 111
EHA	LVLYDNN-RAVWASGTNGRATDCILSMQRDGNLVIYS-G-SRVIWASNTNRQSG-NYY 112
GEAFP	LVLYDNN-RAVWASGTNGKASGCMLKMQNDGNLVIYS-G-SRAIWASNTNRQNG-NYY 112
TMA	LVLYANKVKVLWASRTNGKGGAASCKLSMQNDGNLVIYA-A-TTPVWASRTSRAFA-SYK 113
ZCA	LVLYNVD-KPIWASNTGGLARGCHLSMQRDGNLVVYSQR-NRPIWASNTGGHNAANYV 111
AVA	LVLYDVD-KPIWASNTGGLARGCHLSMQSDGNLVVYSPSGNRPIWASNTQGENG-NYV 110
GNA	LVLYDVD-KPIWATNTGGLSRSCYLNMQTDGNLVVYNPS-NKPIWASNTGGONG-NYV 105
Consensus	LVLY WATG C MQ DGNLVY WAST
7GA	LILQRDRNVVIYDPSGRAIWATGTNVAAAN-VTLSAARPTNLTITAAKLTNNNVPAA 167
CAA	CILQRDRNVVVYDPQGRPIWATGTNVAAASNVTLAAAAAAHGPTNVTVAALNPAA 162
LOA	LILORDRNVVIYDNSNNAIWATGTNVGNAAIAVIPHNNGTAAASGA 157
FHA	LILORDRNVVIYDNSNNAIWATGTNVGNAAIAVIPHNNGTAAASGA 158
GEAFP	LILORDRNVVIYDNSNNAIWATHTNVGNAEITVIPHSNGTAAASGA 158
TMA	LNLQGDGNVVIYGPSG-AIWATNT 136
7CA	I ILOKDRNVVIYGPAKWATGTYTGAVGISGFETGTAGIIKP 152
AVA	CIVOKDRNVVIYGTARWATGTYTGTVGISGFATGMAGIIKS 151
GNA	CILOKDRNVVIYGPARWATGTNIHGAGIVGVPGSAPONSTAEMIKL 151
Consensus	Q D NVV Y WAT T
00113011303	
764	ATEKASTSNGSNDTOVPVTTAGKN 101
	AKTTAKTTAAGGN 175
104	
GENED	
1 m/A 7 C A	
	VTDKSAK 159
AVA CNA	VDKVI I TK
UNA	AUVITIN198
Consensus	

Figure 2. Multiple alignment of TMA with other mannose-binding lectins. The completely identical amino acids were indicated with single capital letter. ZGA (AAQ18904); ZCA (AAM94381); EHA (AAC48927); LOA (AAC37423); AVA (AAP57409); GNA (AAA33349); CAA (AAO59506); AVA (AAP57409) and GEAFP (CAB94239). Mannose-binding sites (QDNY) are underlined.

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ursinum agglutinin (AUA) were grouped into the second cluster as expected. The third cluster grouped most lectins of Amaryllidaceae family such as Zephyranthes candida, Amaryllis vittata, Hippeastrum rutilum, Narcissus hybrid, Galanthus nivalis, Clivia miniata and Lycoris radiata. As expected, the fourth cluster contained most lectins from Araceae species including Polygonatum cyrtonema, Polygonatum multiflorum, Arisaema heterophyllum, Pinellia ternata and Crocus sativus. All the above four groups of lectins were derived from a common ancestor in evolution, suggesting that these lectins shared a common evolutionary ancestor in plants. It is very surprising and interesting that some non-Orchidaceae lectins such as ZGA and CAA from family Amaryllidaceae are very homologous to the typical Orchidaceae lectins such as LOA and EHA, while TMA from gymnosperm species also has much higher similarity to LOA and EHA than those in other families. The finding that mannose-binding lectin exists in gymnosperms suggests that the mannosebinding lectins are much more widely distributed than previously believed.

3.5 *Tissue expression pattern analyses*

To investigate *tma* expression pattern in various tissues of *T. media*, total RNA was isolated from different tissues including leaves, fruits, stems and roots, and subjected to one-step RT-PCR analysis using the primers KF1 and KR1. The result showed that *tma* is expressed in a tissue-specific manner. *Tma* expression could be detected in leaves and stems, but not in fruits and roots (figure 5). Therefore, the *tma* was considered to be a non-constitutively expressing gene.

4. Discussion

The full-length cDNA of a lectin was successfully isolated from *T. media* and Blast search result showed that *tma* had very low identity to all the other known lectin genes in GenBank at nucleotide level (data not shown). However amino acid sequence comparison revealed that TMA shared comparatively higher similarities to other lectins (figure 2). This result suggested that lectin genes



Figure 3. Secondary and three-dimensional structures of TMA. (**A**) The secondary structure of TMA. The helices, sheets, turns and coils are indicated respectively as blue, red, green and purple vertical lines. (**B**) The three-dimensional structure of TMA. **b**-sheets are indicated as patches. Turns and loops are indicated as lines, and the amino acids constituting mannose-binding sites are indicated as balls.



Figure 4. Phylogenetic tree analysis of TMA and other mannose-binding lectins. ZGA (AAQ18904); ZCA (AAM94381); EHA (AAC48927); LOA (AAC37423); AVA (AAP57409); GNA (AAA33349); CAA (AAO59506); GEAFP (CAB94239); *L. radiata* agglutinin (LRA, AAP20877); *Clivia miniata* agglutinin (CMA, AAA19910); *Allium sativum* (garlic) agglutinin (ASA1, AAB64238); *Allium sativum* (garlic) agglutinin (ASA2, AAB64237); AKA (AAP04617); *Tulipa hybrid* agglutinin (THA, AAC49386); APA (AAC37361); TDA (AAQ55289); AAA (S39488); ACA (S39487); *Cymbidium hybrid* agglutinin (CHA, S43463); AUA (AAA16280); *Narcissus hybrid* agglutinin (NHA, AAA33546); *Polygonatum cyrtonema* agglutinin (PCA, AAM28644); ACPA (AAM28277); *Polygonatum multiflorum* agglutinin (PMA, AAC49412); *Hippeastrum rutilum* agglutinin (HRA, AAN73326); *Arisaema heterophyllum* agglutinin (AHA, AAP50524); *Arum maculatum* (cuckoopint) agglutinin (AMA, AAC48998); *Pinellia ternata* agglutinin (PTA, AAP20876); *Crocus sativus* agglutinin (CSA, AAK29077).



Figure 5. Expression analyses of *tma* in different *T. media* tissues. Total RNA ($0.5 \mu g$ /sample) was isolated from leaf, fruit, stem and root respectively, and subjected to one-step RT-PCR amplification (upper panel). *Actin* gene was used as the control to show the normalization of the amount of templates in PCR reactions (lower panel).

in different species are very diverse at nucleotide level but their deduced proteins are fairly conservative, especially the conserved motifs, which may play an important role in biological functions and thus are preserved in evolution. Molecular characterization of TMA such as mannose-binding site analysis, signal cleavage site prediction, secondary and three-dimensional structures analysis revealed that TMA had many typical characters possessed by monocot mannose-binding lectins, implying that TMA might have similar functions with many other mannose-binding lectins such as inhibition of fungal growth or insect feeding. The cloning of *T. media* agglutinin gene (*tma*) will also enable us to test its potential functions for pest resistance by transferring the gene into tobacco in the future.

To our knowledge this is the first report of a mannosebinding lectin gene isolated from gymnosperms. Interestingly, molecular evolution analyses result reveal that some non-Orchidaceae lectins from higher plants and even TMA from gymnosperms show more closer evolutionary relationship to typical Orchidaceae lectins than those in other families; implying that Orchidaceae lectins may be much lower in higher flowering plant species which is consistent with previous study (Peuman *et al* 2002). Considering the relatively low similarity between *tma* and other mannose-binding lectin genes, the divergence of their respective genes from a common ancestor is not a recent event. This unexpected finding extends the existence of the monocot mannose-binding lectin superfamily to family Taxaceae, a taxonomically unrelated family with those of angiosperms. The identification of TMA for the first time as a new member of the family of monocot mannose-binding lectins not only suggests that these lectins are more widespread among plants than previously believed but also provides new insights into further understanding of the molecular evolution of monocot mannose-binding lectins.

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