Identification and characterization of transcription factor IIIA and ribosomal protein L5 from *Arabidopsis thaliana*

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### ABSTRACT

Thus far, no transcription factor IIIA (TFIIIA) from higher plants has been cloned and characterized. We have cloned and characterized TFIIIA and ribosomal protein L5 from *Arabidopsis thaliana*. Primary sequence comparison revealed a high divergence of *At*TFIIIA and a relatively high conservation of *At*L5 when compared with other organisms. The *At*TFIIIA cDNA encodes a protein with nine Cys₂-His₂-type zinc fingers, a 23 amino acid spacer between fingers 1 and 2, a 66 amino acid spacer between fingers 4 and 5, and a 50 amino acid non-finger C-terminal tail. Aside from the amino acids required for proper zinc finger folding, *At*TFIIIA is highly divergent from other known TFIIIA s. *At*TFIIIA can bind SS rRNA, as well as 5S rRNA, and efficiently stimulates the transcription of an *Arabidopsis* SS rRNA gene in vitro. *At*L5 identity was confirmed by demonstrating that this protein binds to SS rRNA but not to 5S rRNA. Protoplast transient expression assays with green fluorescent protein fusion proteins revealed that *At*TFIIIA is absent from the cytoplasm and concentrated at several nuclear foci including the nucleolus. *At*L5 protein accumulates in the nucleus, especially in the nucleolus, and is also present in the cytoplasm.

### INTRODUCTION

Ribosome biogenesis in eukaryotic cells requires the synthesis of RNAs by all three nuclear RNA polymerases. RNA polymerase II (pol II) produces mRNAs that encode ribosomal proteins, while the 5.8S, 18S and 28S rRNAs are co-transcribed in the nucleolus by RNA polymerase I. Transcription factor IIIA (TFIIIA) is pol III transcription factor specifically required for transcription of 5S rRNA genes. It binds to the internal control region of the 5S rRNA genes as the first step in the assembly of a transcription complex, allowing the recruitment of TFIIIC, TFIIIB and pol III (1,2).

TFIIIA has been studied extensively in *Xenopus laevis* where it was first isolated from oocytes (3). Analysis of TFIIIA sequences from several species including *X.laevis* (3) and other frog species (4,5), human (6), catfish (7), mouse, rat (8), *Saccharomyces cerevisiae* (9) and *Schizosaccharomyces pombe* (10) has revealed remarkably poor conservation of primary sequence. All known TFIIIA s, except in *S.cerevisiae* and *S.pombe*, have a similar organization: nine consecutive zinc fingers of the Cys₂-His₂ type, followed by a C-terminal domain of unknown structure required for the support of transcription of 5S rRNA genes in *X.laevis* (11). The *S.cerevisiae* TFIIIA protein bears an 81 amino acid spacer insertion between zinc fingers 8 and 9. In *S.pombe*, TFIIIA contains an unprecedent tenth zinc finger. TFIIIA has been shown to bind to the 5S rRNA in *Acanthamoeba castellanii* (12) and *X.laevis*, where it is involved in a network of interactions that couple 5S rRNA synthesis to accumulation of ribosomal proteins (13). Purified proteins containing TFIIIA activity, isolated from tulip (14) and maize (15), have also been shown to bind 5S rRNA.

Ribosomal protein L5 is also known to bind specifically to 5S rRNA and is involved in its nucleocytoplasmic transport (16–19). After transcription, 5S rRNA binds either to its own transcription factor IIIA or to ribosomal protein L5, forming 7S or 5S ribonucleoprotein particles (RNPs), respectively (reviewed in Pieler and Rudt (20)). It has been suggested that the 5S RNP acts as a precursor to ribosome assembly by delivering 5S rRNA from the nucleoplasm to the nucleolar assembly site of 60S pre-ribosomal subunits (21).

Studies in *X.laevis* oocytes have shown that 5S rRNA can be exported from the nucleus to the cytoplasm, for storage (22). Pre-vitellogenic oocytes of amphibians and fish accumulate two major RNP particles, the 7S RNP and the 42S RNP. The 42S RNP is composed of various tRNAs and oocyte-specific 5S rRNA, along with two proteins, p50 and p43, a 5S rRNA-binding protein (23,24). In fully grown oocytes, 7S and 5S RNPs migrate out of the nucleus and accumulate in the cytoplasm (16,22). 5S rRNA must then re-enter the nucleus to ensure that the ribosome can be fully

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assembled (21,25). The nuclear re-entry of 5S rRNA is mediated exclusively by the ribosomal protein L5 (26–28). The most plausible interpretation for cytosolic export of 5S rRNA in *X. laevis* may lie in control of 5S rRNA synthesis, as suggested by the current view of the feedback regulation mechanism (13). Cyttoplasmic storage sites for 5S rRNA have not been observed in mammalian somatic cells.

To date, no TFIIIA from higher plants has been cloned and characterized. We have identified, cloned and characterized TFIIIA and L5 from *Arabidopsis thaliana*. Primary sequence comparison revealed a high divergence of *AtTFIIIA* and a relatively high conservation of *AtL5* compared with other organisms. As previously shown in *Xenopus*, we have demonstrated that *AtTFIIIA* can bind to 5S rDNA, as well as to 5S rRNA, the gene product. *AtTFIIIA* is able to stimulate the transcription of an *Arabidopsis* 5S rRNA gene in vitro. *AtL5* identity was confirmed by showing that this protein binds to 5S rRNA but not to 5S tRNA. Protoplast transient expression assays with green fluorescent protein (GFP) fusion proteins indicate that *AtL5* protein accumulates in the nucleus and in the nucleolus. *AtL5* is also present in the cytoplasm, probably incorporated in the large ribosomal subunit in association with 5S rRNA. *AtTFIIIA* is only detected in the nucleus, with a strong accumulation in the nucleolus and at additional foci, suggesting that *AtTFIIIA* can be imported efficiently from the nucleoplasm into the nucleolus. We assume that the additional foci found in the nucleoplasm represent accumulation of *AtTFIIIA* on transcribed 5S rDNA loci or into Cajal bodies.

**MATERIALS AND METHODS**

**Isolation of the *Arabidopsis* TFIIIA cDNA and purification of recombinant protein**

A cDNA encoding the putative *Arabidopsis* TFIIIA homolog was amplified by PCR from an *Arabidopsis* cDNA library (29) using primers designed according to the *Arabidopsis* sequence database. The direct primer (5'-ATCATAGGATCTGCG-GGAAGAACTAAAA-3') and reverse primer (5'-ATTACAGGATCTGCG-GGAAGAACTAAAA-3') included an *BamHI* restriction site (underlined). After PCR amplification and *BamHI* digestion, the coding sequence of TFIIIA was cloned into the pGEX-5X-1 expression vector (Amersham Biosciences). In the resulting construct named pGEX-*AtL5*, *AtTFIIIA* is fused to the C-terminal end of GST. Prior to expression in bacteria, sequencing was performed to verify the sequence of the cDNA and the translational fusion. To express the *AtTFIIIA* recombinant protein, pGEX-*AtTFIIIA* was transformed into *Escherichia coli* BLR (DE3) cells. A fresh 2 ml starter culture of BLR (DE3)/pGEX-*AtTFIIIA* was used to inoculate 200 ml of LB containing 100 µg/ml ampicillin and 12.5 µg/ml tetracycline. The culture was grown at 37°C to an OD<sub>600</sub> of 0.3–0.4. Recombinant protein expression was then induced by addition of 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) followed by an incubation of 2–3 h at 30°C. Induction was verified by SDS–PAGE analysis followed by Coomassie blue staining. Induced cells were harvested by centrifugation at 4000 g for 15 min at 4°C, and resuspended in 1 ml of ice-cold buffer A (20 mM Tris–HCl pH 8.0, 150 mM NaCl). Cell lysis was performed by addition of 100 µg/ml lysozyme, followed by a 15 min incubation at 30°C and sonication. Soluble cell extract containing *AtTFIIIA* was recovered by centrifugation at 12 000 g at 4°C for 30 min, and saved at 4°C for purification.

The extracts containing the GST–*AtTFIIIA* recombinant protein or GST were mixed with glutathione–agarose (Sigma) for 30 min at room temperature, and unbound protein was removed by three washes in 1 M NaCl and three washes in 1× phosphate-buffered saline (PBS). Subsequently, bound protein was removed by elution with 10 mM reduced glutathione pH 7.6 (Sigma). Eluates were analyzed by SDS–PAGE.

**Isolation of the *Arabidopsis* L5 cDNA and purification of recombinant protein**

As for *AtTFIIIA*, the cDNA encoding the *Arabidopsis* L5 homolog was amplified by PCR from the *Arabidopsis* cDNA library, using primers designed according to the L5 cDNA sequence present in the database (accession no. AY081701). The direct primer (5'-ATTCTATAGATCTTTGTTGTGGTGGTGT-3') and reverse primer (5'-ATTCTATAGATCTTTAGCTTTCATC-3') included an EcoRI restriction site. After PCR amplification and EcoRI digestion, the L5 cDNA was cloned into the pGEX-5X-1 expression vector. In the resulting construct named pGEX-*AtL5*, *AtL5* is fused to the C-terminal end of GST. Prior to expression in bacteria, sequencing was performed to check the sequence of the cDNA (accession no. AY186611) and the translational fusion.

Expression and purification of the GST–*AtL5* recombinant fusion protein were as described above for *AtTFIIIA*, except that cells were grown at 37°C after induction with 1 mM IPTG.

**DNA gel retardation assays**

The 238 bp *NotI* fragment of pGEMT-5S containing the transcribed region and a part of the spacer sequence of an *A. thaliana* 5S rRNA gene was labeled by a fill-in reaction performed with 25 µCi of [γ-<sup>32</sup>P]dCTP (3000 Ci/mmol), 100 µM each dATP, dGTP and dTTP, 1 mM dithiothreitol (DTT), 1 µM T7 polymerase, 10 mM MgCl<sub>2</sub>, 50 mM NaCl, 1 mM dithiothreitol (DTT), 100 ng of the 5S DNA fragment and 25 U of Klenow fragment (Amersham). After a 30 min incubation at room temperature, the labeled fragment was purified with a PCR purification kit (Qiagen). Recombinant proteins (see legends to figures for concentrations) were incubated with 1 µl (20 000 c.p.m., 1–3 ng, ~1 nM) of the labeled 5S rDNA fragment and variable concentrations of unlabeled DNA in 20 µl reactions containing buffer EMSA (20 mM Tris–HCl pH 7.5, 7 mM MgCl<sub>2</sub>, 10 µM ZnCl<sub>2</sub>, 1 mM DTT, 10% glycerol, 70 mM KCl) supplemented with 10 µg/ml poly(dI–dC) and 100 µg/ml bovine serum albumin (BSA). Unlabeled DNA referred to as ‘specific’ is the 238 bp *NotI* fragment of pGEMT-5S, while ‘non-specific’ refers to a 490 bp EcoRI fragment from the plasmid pGEM-APT1 that contains the cDNA of the adenine phosphoribosyltransferase (APRT) gene (30). The reaction mixtures were incubated at 25°C for 30 min and then quickly cooled on ice before addition of 3 µl of loading buffer [50% (v/v) glycerol, 1 mg/ml bromophenol blue]. The samples were loaded onto an 8% polyacrylamide gel containing 5% glycerol in 25 mM Tris–HCl pH 8, 200 mM glycine. Prior to loading, the gels were pre-run at 50 V for 30 min. After loading, electrophoresis...
was continued for an additional 2 h at 140 V at room temperature. The gels were dried, and visualized on a PhosphorImager.

RNA gel retardation assays

The transcribed sequence of an *A. thaliana* 5S rRNA gene was fused to a T7 promoter by PCR. Labeled 5S rRNA was then synthesized *in vitro* using the SP6/T7 transcription kit (Roche), following the manufacturer’s instructions. Non-specific RNA was generated similarly using the first 120 bp of the *APTJ* cDNA fused to a T7 promoter. Transcripts were separated on an 8% acrylamide gel containing 7 M urea, and RNAs were eluted from the gel by an overnight incubation at 37°C in elution buffer (0.5 M ammonium acetate, 1 mM EDTA, 0.2% SDS). After ethanol precipitation, RNAs were resuspended in diethylpyrocarbonate (DEPC)-treated water. Before use in the gel retardation assays, the RNA probes were incubated for 10 min at 65°C in renaturation buffer (50 mM Tris–HCl pH 8.0, 50 mM KCl, 5 mM MgCl₂) and allowed to cool slowly to room temperature. RNA gel retardation assays were then performed according to the procedure described above for DNA gel retardation.

DNase I footprinting assay

For footprinting of the template strand, 5′-labeled 5S rDNA was generated by PCR using primers OLIGO5 (5′-TATATACGATGGCATTGCATATAC-3′) and 32P-labeled 1037PE-rev (5′-[32P]-GGAAAACGCTATGACCATGGAGGGATGCAACACGAGGAC-3′) from a 5S rDNA transcribed unit (1037). For footprinting of the RNA-like strand, 5′-labeled OLIGO5 primer was used instead of labeled 1037PE-rev. Binding was performed in a 50 µl volume containing 30 mM HEPES-KOH pH 7.9, 3 mM MgSO₄, 80 mM KOAc, 0.1 mM EDTA, 2 mM DTT, 10% glycerol, 0.8 µg of poly(dl–dC)-poly(dl–dC), 0.15 pmol of labeled 168 bp double-stranded DNA including the ~28 to +120 region of 5S rDNA, and ~400 ng of recombinant protein (GST or TFIIA). After incubation for 15 min at room temperature, 5 µl of Ca²⁺/Mg²⁺ solution (5 mM CaCl₂ and 10 mM MgCl₂) was added, and then 0.33 U of DNase I was added to the mixture and incubated for just 1 min at room temperature. After phenol/chloroform extraction and ethanol precipitation, the 32P-labeled fragments were separated on an 8% polyacrylamide gel containing 7 M urea and TBE. Radioactivity was detected and measured using BAS-2000 II.

**Protoplast transient expression assay with GFP fusion proteins**

The NcoI-ArTFIIA cDNA was PCR amplified using the primers (5′-ATTCTATGAATCTTGGTCTTGTG-3′) and (5′-ATTCTATGAATCTTACCCTCTC-3′) and inserted at the NcoI restriction site of the GFP fusion vector pAVA393 (32) containing a cytosolic derivative of the GFP5 cDNA (33) driven by the constitutive 35S cauliflower mosaic virus (CaMV) promoter. In the resulting construct, ArTFIIA is fused to the N-terminal region of the GFP5. Similarly, the ArL5 cDNA was amplified by PCR using the primers (5′-ATTCTATCCATGTTTGTTGGAAGG-3′) and (5′-AATCTATCCAGCTTACATGCATCT-3′) including NcoI restriction sites, and cloned into pAVA393. The resulting constructs, named pAVA-ArTFIIA and pAVA-ArL5, respectively, were verified by sequencing.

Protoplasts were prepared from *Arabidopsis* cultured cells and transformed as described (34) with minor modifications (C.I. White, unpublished). Transformations were performed with 50 µg of plasmids (pAVA393, pAVA-ArTFIIA or pAVA-ArL5) purified using the Plasmid midiprep Kit (Qiagen). Protoplasts were observed 30 h after transformation using a ZEISS Axioplan 2 microscope.

**RESULTS**

**Amplification and cloning of the putative *Arabidopsis* TFIIA and L5 cDNAs**

Searching Cys₂-His₂-type multi-zinc finger proteins in the *Arabidopsis* database resulted in the identification of only one predicted protein containing nine zinc fingers (accession no. AAG51140). As the presence of nine zinc fingers is the only common property between TFIIA proteins characterized to date, we considered this protein to be a likely candidate to be *Arabidopsis* TFIIA (named ArTFIIA). The open reading frame encoding this protein (accession no. AC069273) is carried by the BAC clone F28P5 mapped on chromosome 1. A cDNA encoding the putative ArTFIIA protein was amplified by PCR from a cDNA library using primers designed according to the sequence of the predicted cDNA. The sequence of the obtained cDNA (accession no. AY186610) revealed that the predicted second exon was 66 bp too long in its 3′ end. Furthermore, predicted exons 5 and 6 were 20 and 43 bp too short in their 3′ and 5′ end, respectively. The remainder of the coding sequence was identical to the database prediction.

The 412 amino acid encoded protein contains nine Cys₂-His₂-type zinc fingers as well as 19 amino acid N-terminal and 50 amino acid C-terminal non-zinc finger tails. The nine zinc finger sequences comply with the F/I-X(4)-X(3)-X(2)-3-L-X₂-H-X₂(3)-5-H consensus. Sequence similarity between ArTFIIA and any other known TFIIA is dominated by the seven residues found in each zinc finger that are required for proper folding of the protein. These include the...
Figure 1. Primary sequence alignment of known TFIIAs with Arabidopsis TFIIIA. Sequences were aligned manually to match each of the nine zinc fingers of AtTFIIIA with the corresponding finger of the other TFIIAs. Zinc fingers are framed and numbered. Note that the ninth finger from S.cerevisiae was aligned with finger 10 from S.pombe only for illustration. Non-aligned regions between fingers 1 and 2, and 4 and 5 in the AtTFIIIA sequence are underlined. AtTFIIIA zinc fingers are in red, and zinc fingers of other organisms are shown in blue. Conserved residues (in >50% of the sequences) in the zinc fingers or in non-finger regions are highlighted in yellow or gray, respectively. The oocyte motif MGEK/R and the NES motifs are highlighted in green and blue, respectively. NLS motifs are written in green characters, and TAS motifs are indicated in black bold type. Asterisk marks the end of the sequence. The residue number is shown to the right of each line of sequence.
hydrophobic residues: F or I in *Arabidopsis* (but F or Y in other species), F, L and two Zn$^{2+}$-coordinating cysteines and histidines. Excluding these conserved residues, the overall identity of AtTFIIIA to known TFIIIA is low. For instance, there is only 26% sequence identity between *A. thaliana* and *X. laevis*, and 17% sequence identity between *A. thaliana* and *S. cerevisiae*. Alignment of TFIIIA sequences reveals that AtTFIIIA is organized differently from its homologs (Fig. 1). AtTFIIIA contains two unique 23 and 66 amino acid spacers located between zinc fingers 1 and 2, and 4 and 5, respectively. Amongst known TFIIIA, such a long spacer sequence (81 amino acids) exists between fingers 8 and 9 in *S. cerevisiae* TFIIIA. The *S. pombe* TFIIIA carries a unique tenth zinc finger which has no equivalent in other organisms. Several sequence motifs have been defined in the different TFIIIA sequences, but none of them could be found either in the N- and C-terminus tails, or in the spacers of the AtTFIIIA protein sequence. We searched for the N-terminal MGEK motif characteristic of the smaller oocyte form of *X. laevis* TFIIIA (3–5,35), the nuclear localization signal (NLS) (7,8), the transcription-activating signal (TAS) (7,11,36) and the nuclear export signal (NES) (7,37) found in some of the reported TFIIIA sequences (Fig. 1), but could not identify any of them in the terminal tails nor in the spacer sequences of *AtTFIIIA*.

The cDNA encoding the putative *Arabidopsis* 5S rRNA-binding ribosomal protein L5 (*AtL5*) was PCR amplified from the cDNA library, using primers designed according to the sequence of the cDNA present in the database. Unlike AtTFIIIA, the primary sequence of the putative AtL5 protein revealed a high degree of conservation (~54% identity between *Arabidopsis* and human, see Fig. 2).

**Sequence-specific DNA-binding activity**

To investigate the biochemical properties of the putative AtTFIIIA and AtL5 proteins, the cloned cDNAs were used for expression of the proteins in the E. coli BLR(DE3) strain. Two recombinant GST fusion proteins (GST–AtTFIIIA and GST–AtL5) were purified (Fig. 3A and D).

One property of TFIIIA is to bind specifically to the 5S rRNA gene. Gel retardation assays demonstrate that the recombinant protein indeed possesses 5S rDNA-binding activity, although the GST protein alone does not (Fig. 3B, C and F). Competition experiments revealed that the DNA-binding activity of the AtTFIIIA recombinant protein is 5S sequence specific (Fig. 3B and C). As expected, GST–AtL5 protein does not bind to the 5S rRNA gene (Fig. 3E). The equilibrium binding constant ($K_d$) for the interaction between AtTFIIIA and 5S rDNA, reflecting the affinity of AtTFIIIA for the 5S rRNA gene, was determined using a gel mobility shift assay as described previously (10). We measured a $K_d$ of 0.33 nM (SE ± 0.05) for the interaction between AtTFIIIA and 5S rRNA. This value is comparable with those reported previously for other known TFIIIA (10,38,39).

**DNase I footprinting**

To analyze further the binding of AtTFIIIA to the 5S rRNA gene, DNase I footprinting was performed on both the template and the RNA-like strands (Fig. 4), using recombinant GST–AtTFIIIA and GST as a control. The protection pattern of the AtTFIIIA protein extends from position +43 to +97 on the template strand and from +45 to +103 on the RNA-like strand. Unprotected regions are found from +63 to +76 on the template strand and from +71 to +76 on the RNA-like strand. This protection pattern is similar to those observed for *X. laevis* [+47 to +96; (10)], and recently reported for *A. castellanii* [+44 to +97; (12)] and *S. pombe* [+45 to +95; (10)]. However, we did not observe any DNase I-hypersensitive site around position +63 as described for these known TFIIIA proteins (10,12,40). As expected, GST alone does not produce any protection along the 5S gene.

**5S rRNA-binding activity of putative AtTFIIIA and AtL5**

Another characteristic of TFIIIA is to bind not only to the 5S rRNA gene, but also to 5S rRNA, the gene product. Similarly, the L5 ribosomal protein can bind to 5S rRNA, forming the 5S RNP. To investigate this common property between TFIIIA and L5 on the putative AtTFIIIA and AtL5 proteins, we performed gel retardation assays using an in vitro transcribed 5S rRNA. Incubation of increasing concentrations of

![Figure 2. Primary sequence alignment of AtL5 with some known L5s. Sequences were aligned using the CLUSTALW program. Identical residues (in 100% of the sequences) are highlighted in black, and conserved residues (in >50% of the sequences) are highlighted in gray. The residue number is shown to the right of each line of sequence.](https://academic.oup.com/nar/article-abstract/31/9/2424/1080337/2428-Nucleic-Acids-Research-2003-Vol-31-No-9)
GST–ATFIIIA or GST–AtL5 protein with 5S rRNA resulted in the appearance of a 5S rRNA–protein complex with slower mobility in non-denaturing gel electrophoresis (Fig. 5A and B). Control experiments showed that GST alone does not bind to 5S rRNA (Fig. 5D) and that neither GST–AtL5 nor GST–ATFIIIA bind to a non-specific RNA (Fig. 5C).

Taken together, these results demonstrate that the putative ATFIIIA protein indeed possesses a 5S rRNA-binding activity and that AtL5 is the actual Arabidopsis homolog of the ribosomal L5 protein.

Figure 3. DNA binding assays of GST–ATFIIIA and GST–AtL5 recombinant proteins. (A and D) SDS–PAGE of GST–ATFIIIA and GST–AtL5 recombinant proteins, respectively. Shown are results for uninduced E.coli BLR (DE3) (lane 1), IPTG-induced cells (lane 2) and purified recombinant proteins (lane 3). Proteins were visualized by staining with Coomassie brilliant blue. The sizes (in kDa) of molecular mass markers run in lane M are indicated on the left. Increasing concentrations (1–170 nM) of unlabeled 5S rDNA (B) or non-specific competitor DNA (C) were added to binding reactions including labeled 5S rDNA and GST–ATFIIIA (2.5 ng/μl). Binding reactions were performed with increasing concentrations (2.5–10 ng/μl) of either AtL5 (E) or GST alone (F). Arrowheads indicate free (unbound) probe, and protein–DNA complexes are indicated by an asterisk. n.p., no protein.

Figure 4. DNase I footprint of ATFIIIA. DNase I cleavage of each strand is presented. Numbers to the right of each gel indicate the position of DNase I cleavage relative to the start site of transcription (+1). The positions of the A-box, the intermediate element (IE) and the C-box are shown to the left of each gel.

Figure 5. 5S rRNA binding analysis of GST–ATFIIIA and GST–AtL5 proteins. A constant amount of labeled 5S rRNA probe synthesized in vitro was incubated with increasing concentrations (2.5–10 ng/μl) of GST–AtL5 (A), GST–ATFIIIA (B) or GST (D) proteins and then subjected to gel mobility shift analysis. As a control, GST–ATFIIIA and GST–AtL5 proteins were incubated with non-specific RNA (C). Arrowheads indicate free (unbound) probe, and protein–RNA complexes are indicated by an asterisk. n.p., no protein.
Transcriptional activity of the putative AtTFIIIA protein

To test the ability of putative AtTFIIIA to support transcription of the Arabidopsis 5S rRNA gene, we used an in vitro transcription system from tobacco cells (41,42). This system can transcribe an Arabidopsis 5S rRNA gene without added recombinant AtTFIIIA (GST–AtTFIIIA) or GST. (B) Quantification and graphical representation of the data from several independent experiments similar to that shown in (A). The asterisk and arrowhead indicate endogenous tRNA and 5S rRNA, respectively. Molecular sizes are indicated (in nucleotides) next to the gel.

Figure 6. In vitro transcription assays. (A) Reactions were performed without added recombinant protein (n.p.) or with increasing concentrations (2.5–7.5 ng/μl) of GST–AtTFIIIA or GST. (B) Quantification and graphical representation of the data from several independent experiments similar to that shown in (A). The asterisk and arrowhead indicate endogenous tRNA and 5S rRNA, respectively. Molecular sizes are indicated (in nucleotides) next to the gel.

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Taken together with 5S rDNA- and 5S rRNA-binding activities, the ability of putative AtTFIIIA to stimulate 5S rRNA gene transcription efficiently in vitro clearly demonstrates that the protein we have characterized is indeed A.thaliana TFIIIA.

Cellular localization of AtTFIIIA and AtL5 proteins

To investigate the cellular localization of the AtTFIIIA and AtL5 proteins, translational fusions of AtTFIIIA and AtL5 with an enhanced version of GFP (mGFP5) were made. Each construct was then transiently transfected to Arabidopsis cells protoplasts and monitored for GFP expression 30 h post-transfection.

As previously reported (43,44), and expected due to its small size, expression of GFP alone resulted in cytoplasmic and nuclear signals, the nucleolus being clearly devoid of signal (Fig. 7A and B). AtL5–GFP protein accumulates in the nucleus with significant nuclear enrichment. A faint signal was observed in the cytoplasm, indicating that the protein is also present in this cellular compartment (Fig. 7C and D). This result is in good agreement with previous observations in somatic mammalian cells (21).

In contrast, expression of the AtTFIIIA–GFP protein resulted in complete absence of fluorescence in the cytoplasm (Fig. 7E and F), suggesting the presence of a functional NLS.
in AtTFIIA although not identified by sequence comparison (see Fig. 1). The nuclear localization of AtTFIIA was heterogeneous, with a higher accumulation of the fusion protein in the nucleolus together with additional foci (Fig. 7F–H). As for AtL5, we conclude that AtTFIIA is imported efficiently from the nucleoplasm to the nucleolus.

**DISCUSSION**

We report here the first cloning and characterization of TFIIIA from a higher plant, A. thaliana. We have shown that the protein has a specific 5S rRNA-binding activity together with a 5S rRNA-binding property as previously reported for X. laevis and A. castellanii TFIIAs (12,45). Moreover, AtTFIIA can stimulate the transcription of an Arabidopsis 5S rRNA gene efficiently *in vitro*.

In the Arabidopsis database, only predicted TFIIA cDNA was present. AtTFIIA cDNA escaped sequencing programs probably because of the low content of AtTFIIA in somatic cells. Indeed, TFIIA was first purified from X. laevis oocytes which were shown to contain up to 10^{12} molecules per cell (3). For comparison, somatic A. castellanii and HeLa cells were estimated to contain only 170 and 400 molecules of TFIIA, respectively (12,46). Rat TFIIA was purified from breast tumor, and tumoral cells are known to contain enhanced pol III transcription [for a review see Brown et al. (47)]. The reason for our success in the amplification of AtTFIIA cDNA could lie in the fact that we used a cDNA library from metabolically active Arabidopsis cells.

AtTFIIA bears nine Cys_{2}-His_{2}-type zinc fingers including the conserved residues required for proper folding and specific 5S rRNA binding. Irregular spacing between these residues in the zinc fingers, C-X_{2-4}-C-X_{11-12}-H-X_{3-5}-H, may play a role in the alignment of AtTFIIA zinc fingers along the 5S rRNA gene. The overall sequence identity between AtTFIIA and yeast or vertebrate homologs ranges between 17 and 26%. Thus the divergence is too important to detect any convincing pattern of higher similarity between Arabidopsis and mammals, amphibians or yeasts. Apart from zinc fingers, known TFIIAs contain several sequence motifs. In the non-finger N-terminal region, the mammalian, yeast and Arabidopsis TFIIAs lack the conserved MGEKR/R motif characteristic of the smaller oocyte form of TFIIA from amphibians, also found in catfish TFIIA purified from immature ovarian tissue (7). We assume that Arabidopsis does not synthesize an oocyte form of TFIIA as already proposed for mammals (8).

Vertebrate TFIIAs contain a variable length (49–68 amino acids) non-zinc finger region at the C-terminus, which contains a TAS, approximately 25 amino acids long, in amphibians (5,11) and mammals (8). In S. cerevisiae, the 81 amino acid spacer located between fingers 8 and 9 exhibits a leucine-rich oligopeptide required for transcription (48), which differs in sequence from the amphibian or mammalian TAS motifs. In Arabidopsis, the TAS domain could lie in the 66 amino acid spacer between fingers 4 and 5 or, alternatively, in the 50 amino acid long C-terminal tail. We could not find the NLS consensus sequence (KKKM/LXXKL) present in the C-terminal region of vertebrates TFIIAs (8) nor the RLRRKLKKRR NLS found in S. cerevisiae (48). Nonetheless, AtTFIIA–GFP fusion experiments have clearly revealed a nuclear accumulation and an absence of the protein in the cytoplasm of Arabidopsis protoplasts, indicating the presence of a functional NLS in AtTFIIA which remains to be identified. Finally, a NES (LXXLTI) has been identified and functionally tested in amphibians (37). A similar motif, based on sequence comparison, has been described in catfish (LXXLSL) (7) and humans (VAVLTL) (8). No similar sequence motif could be found in AtTFIIA primary sequence and, accordingly, AtTFIIA–GFP fusion protein does not localize in the cytoplasm of transformed protoplasts.

The nuclear export of 5S rRNA by TFIIA for subsequent accumulation at distinct cytoplasmic storage sites has only been reported in Xenopus oocytes (16,22). Pre-vitellogenic oocytes store 5S rRNA in the cytoplasm as either 7S (5S rRNA–TFIIA complex) or 42S RNPs. In the 42S RNP particle, the p43 protein binds to the 5S rRNA (23,24). Mature oocytes represent a particular cell type where TFIIA accumulates to 10^{12} molecules per cell. TFIIA or L5 protein binds to 5S rRNA, and each of these two RNPs migrates out of the nucleus and accumulates in the cytoplasm, prior to development. Cytoplasmic storage sites for 5S rRNA have not been observed in somatic mammalian cells (19). This cytoplasmic phase of the 5S rRNA biosynthetic pathway is probably unique to oocytes and does not occur in somatic cells (18).

We show here that the Arabidopsis ribosomal protein L5 binds to the 5S rRNA and accumulates in the nucleolus. Immediately after transcription, 5S RNA is transiently associated with the La protein which, amongst other things, functions in transcription termination of pol III transcripts (49,50). After association with La, 5S RNA is bound by ribosomal protein L5 to form a 5S RNP particle. Then the 5S RNP migrates to the nucleolus to participate in large ribosomal subunit assembly. L5 accumulates in the nucleolus at a concentration which greatly exceeds that of assembling ribosomal subunits (18). Rosorius et al. (19) have shown that the binding of L5 protein to 5S rRNA correlates with its ability to accumulate in the nucleolus, as previously demonstrated for the nucleolin protein (51). After nucleolar localization, the 5S RNP becomes incorporated into large ribosomal subunits and is then exported from the nucleus to the cytoplasm. In Arabidopsis protoplasts, the AtL5–GFP fusion protein localizes predominantly to the nucleolus and to a lesser extent to the nucleoplasm, and exhibits a faint homogenous staining pattern in the cytoplasm which probably reveals the protein incorporated into ribosomes. This pattern is in good agreement with previous observations made in monkey and human cells using an anti-5S RNP antibody (21).

In Arabidopsis protoplasts, the AtTFIIA–GFP fusion protein was concentrated in the nucleolus and at several nuclear foci. Since 5S rRNA transcription occurs in the nucleoplasm and AtTFIIA specifically binds to 5S rRNA, these nuclear foci could correspond to the transcribed 5S rDNA loci. In the germline vesicle of Xenopus oocytes, it was reported previously, using polyclonal antibodies, that TFIIA localizes in nuclear organelles, called Cajal bodies, which were assumed to be the primary site for assembly of the transcription machinery of the nucleus (52,53). Hence, we hypothesize that the fluorescent foci observed along with the nucleolus could also correspond to Cajal bodies where AtTFIIA would accumulate and be incorporated in the pol III transcription machinery before delivery to the
chromosomal sites of 5S rDNA transcription. In germinial vesicle of Xenopus oocytes, TFIIIA was only detected in Cajal bodies but not in nucleoli (52,53). To our knowledge, our results describe for the first time the localization of TFIIIA in somatic cells and its presence in the nucleolus. The reason for the nucleolar accumulation of AtTFIIIA remains unclear because nucleoplasm–nucleolus shuttling of 5S rRNA is known to be mediated by L5. However, as AtTFIIIA binds 5S rRNA, this nucleoplasm–nucleolus trafficking could occur as 7S RNP.

A model in which a network of nucleic acid–protein interactions, involving TFIIIA, L5, 5S rRNA and 5S rDNA, regulates 5S rRNA synthesis has been proposed (13). In this model, formation of 7S RNP regulates 5S rRNA synthesis because it lowers the amount of free TFIIIA available for 5S rDNA transcription. In Xenopus oocytes, 7S RNP are sequestered in the cytoplasmic compartment. In the light of our results, we formulate the hypothesis that in somatic cells, 7S RNP are stored in the nucleolus.

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