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# A Putative Proline-rich Protein of B. napus

# Parameswari Namasivayam<sup>1,\*</sup> and David Hanke<sup>2</sup>

<sup>1</sup>Department of Cell and Molecular Biology, Faculty of Biotechnology and Biomolecular Sciences, Universiti Putra Malaysia, 43400 UPM, Serdang, Selangor, Malaysia <sup>2</sup>Department of Plant Sciences, University of Cambridge, Downing Street, Cambridge CB2 3EA, UK \*E-mail: parameswari@biotech.upm.edu.my

#### ABSTRACT

Proline-rich proteins are among the major protein components of plant cell walls. So far, two different proline-rich cell wall proteins have been described in *Brassica napus*. This paper reports a study on expression and sequence analysis of a novel class of a proline-rich putative protein, tentatively designated Ae4. The largest ORF of Ae4 encodes 166 amino acid residues without the start and stop codons. Ae4 is a partial length cDNA. The Ae4 gene expression was investigated and the results demonstrate that it accumulates in all vegetative tissues tested as well as in the embryogenic culture of *Brassica napus*. However, expression of Ae4 was undetectable in the non-embryogenic and cytokinin-treated embryogenic tissues. These results indicate that the Ae4 gene might play a role in somatic embryo formation.

Keywords: Proline-rich protein, PRP, Brassica napus, oilseed rape, somatic embryogenesis

## INTRODUCTION

Proline-rich proteins (PRPs) are one class of structural cell wall protein members (Showalter, 1993). All the PRPs are characterised by the repeating occurrence of Pro-Pro repeats contained within a variety of larger repeat units. These proteins lack the SerPro4 repetitive element defined for extensins (Jose-Estanyol and Puigdomenech, 2000). The most extensively studied PRPs are from soybean that contains the PPVYK motif or variations (Hong et al., 1987, 1989; Datta et al., 1989; Wyatt et al., 1992). Members of the PRP gene family were shown to be developmentally regulated and their expression tissue/organ specific (Hong et al., 1989; Lindstrom and Vodkin 1991; Jose-Estanyol et al., 1992). In general, PRPs are thought to have a structural role in the cell wall (Cassab and Varner, 1988). They have been also implicated in plant defence reactions (Chen and Varner, 1985; Ebener et al., 1993), nodule morphogenesis (Franssen et al., 1987; Wilson et al., 1994), and are expressed during somatic embryogenesis (Aleith and Reichter, 1990; Gyorgyey et al., 1997; Yasuda et al., 2001).

To date, two different proline-rich cell wall proteins have been described in *Brassica napus* (*B. napus*). In this paper, we report the sequence and expression analysis of an additional proline-rich putative protein, tentatively designated as *Ae*4. The amino acid sequence of *Ae*4 has been deduced from the nucleic acid sequence of a copy DNA (cDNA), isolated previously from a subtracted library of *B. napus* embryogenic culture (Namasivayam *et al.*, 2006b).

## MATERIALS AND METHODS

Sources of Plant Materials

Plants of *Brassica napus* ssp. *oleifera* cv. Primor were grown from seeds, in pots with soil in the Botanic Garden, Cambridge. Sources and preparation of plant materials for the preembryogenic, mature embryogenic and non-embryogenic *Brassica napus* ssp. *oleifera* cv. Primor culture was as described in Namasivayam *et al.* (2006a,b). The cytokinin-treated embryogenic tissue was generated from hypocotyls of embryoids grown for 20 days on MS media containing  $10^4$  M kinetin, 2% (w/v) sucrose and

<sup>\*</sup> Corresponding Author

0.8% (w/v) agar. Various organs/tissues such as young leaves, stem, buds, flowers, siliques, roots, stamens, carpels, petals and sepals from mature *B. napus* plants were harvested, immediately frozen in liquid nitrogen and stored at–80°C until isolation of total RNA.

## Sequence Analysis

The Ae4 cDNA sequence has been submitted to the GenBank under the accession number AY570239. Sequence analysis was carried out using BLAST 2.0 (Basic Local Alignment Search Tool; Altschul et al., 1997), accessible from the internet (http://www.ncbi.nlm.nih.gov/BLAST). Alignments of the protein sequence with several closely related genes was carried out using the CLUSTAL W program from the Biology Workbench version 3.2, accessible from the internet (http://biowb.sdsc.edu/CGI/BW.cgi). Other sequence analyses were performed using Biology Workbench version 3.2 to compute molecular weight (MW), hydrophobicity and isoelectric point (pI) determination.

#### Total RNA Isolation

Total RNA from various frozen tissues/organs of the mature plant and tissue culture materials were extracted using the acid guanidium thiocyanate-phenol-chloroform extraction method described by Chomczynski & Sacchi (1987). The concentration of RNA in each sample was determined spectrophotometrically (Sambrook *et al.*, 1989).

# Northern Blot Analysis

Equal amounts of total RNA (10 (g per lane) were resolved on 1.3% (w/v) agaroseformaldehyde denaturing gel and blotted onto Hybond<sup>TM</sup>-XL nylon membrane (Amersham Biosciences). Hybridisation was carried out at 65°C using standard techniques (Sambrook et al., 1989). The entire Ae4 sequence was used as probe labelled with [ $^{32}$ P- $\alpha$ ]-dCTP using the Prime-IT<sup>®</sup> II Random Primer Labeling Kit (Stratagene). Washes were carried out at room temperature in the first wash buffer (40 mM sodium phosphate pH 7.2, 1% (w/v) SDS and 1 mM EDTA) for 10 min and followed by second wash in 40 mM sodium phosphate pH 7.2, 5% (w/v) sodium dodecyl sulphate (SDS) and 1 mM ethylenediaminetetraacetic acid (EDTA) at 65°C for 15 min. The hybridisation signals were captured by a Phosphorimager Typhoon 8600 (Amersham Pharmacia Biotech). After removal of probe, the same blot was hybridised with radiolabelled Arabidopsis *Actin*2/7 cDNA probe as a loading control.

#### RT-PCR

Equal amounts of DNase-treated total RNA (200 ng) from each tissue sample was added individually to a sterile 0.2 ml polymerase chain reaction (PCR) tube and the volume adjusted to 13.5 µl with DEPC-treated sterile deionised water (SDW). Oligo ( $dT_{18}$ ) (1 µl of 20 pmoles/µl) was added to the tube and the reaction mix incubated for 10 min at 70°C. Following brief centrifugation, the following reagents were added: 4 µl 5 x first strand buffer (Promega), 0.5 µl 'RNase Out' ribonuclease inhibitor (40 U/μl) (Invitrogen), 0.5 μl 10 mM Bioline dNTPs mix and 0.5 µl MMLV-RT RNase H minus (200  $U/\mu l$ ) (Promega) and incubated at 37°C for 1 h. Later, the reaction mix was heat deactivated before using for PCR reactions. PCR reactions were performed in 12.5 µl reactions with the following components: 2 µl of the RT product, 1 x Bioline PCR buffer (Mg<sup>2+</sup>- free), 1.5 mM MgCl<sub>2</sub>, 0.4 mM dNTP mix, 2.5 pmoles of forward primer (5' GGACTATAAATTGGTGTTGGAGGTTTCA 3') and reverse primer (5'- TATTTATAGT CCTCCCGTAATGCCA - 3() respectively, and 1.5 U BioTaq DNA polymerase. An internal control was prepared using actin2 primers (forward primer: 5'-CCATTCTTGCTTCCCTCAG-3' and reverse primer: 5(-GACGTAAGTAAAAACCCAG-3') and containing all the components as above to test for equal loading of the template. Also, a negative control without template was included. Amplification was performed as follows: 95°C for 3 min; followed by 35 cycles at 94°C for 30 s, 65°C for 1 min, 72°C for 1 min; and a final extension at 72°C for 3 min. The annealing temperature used for Actin2 was 60°C. Reverse transcription polymerase chain reaction (RT-PCR) products were separated on a 2% (w/v) agarose gel and the gel was photographed. The agarose gel containing the PCR amplified products was blotted (Southern, 1975) and hybridised with a labelled specific probe (Ae4 cDNA).

#### RESULTS

## Sequence Analysis

The BNPE AE4 clone contains a cDNA insert of 678 bp, excluding a poly A tail. The longest open reading frame (ORF) encodes 166 amino

acids starting from nucleotide no. 3 and there is no start and stop codons, as shown in Fig. 1. Therefore, it is unlikely to be a full length clone. The predicted amino acid sequence is rich in proline (44.89 %), lysine (10.2 %), threonine (10.22 %), valine (8.89 %), serine (6.67 %), tyrosine (5.78 %) and glutamine (5.78 %). The predicted protein fragment represents a calculated molecular weight of 17.7 kDa and has a predicted pI value of 10.5. Hydropathy analysis indicated that 13 residues at the N terminal end of the predicted protein fragment are hydrophobic and the other regions are highly hydrophilic. The polypeptide is primarily composed of two repeat units: a 10-mer repeat unit (PPI/VK/MPPPVQK/Q) and a 7-mer repeat unit (PPTPI/S/TYS). Protein database search revealed that there is no significant similarity between the predicted amino acid sequence and protein sequences deposited in

the GenBank. However, comparison of the Ae4 nucleotide sequence with nucleotide sequences in the GenBank showed that it is homologous (82% identical) to a genomic fragment of Arabidopsis in chromosome 2 (At2g27380) which encodes a putative proline-rich protein. Also, a few hits to B. napus seed EST sequences with a homology of 80% to 95% were found in the database. A comparison of the amino acid sequence of this clone with the Arabidopsis putative proline-rich protein and translated EST sequences is depicted in Fig. 2.

## Expression Analysis of Ae4

The pattern of expression of Ae4 in organs and tissues was investigated using Northern analysis and RT-PCR. The Northern analysis on the organ/tissue specific-blot failed to detect a distinct band for Ae4 transcripts except for a very faint smear observed in each sample lane

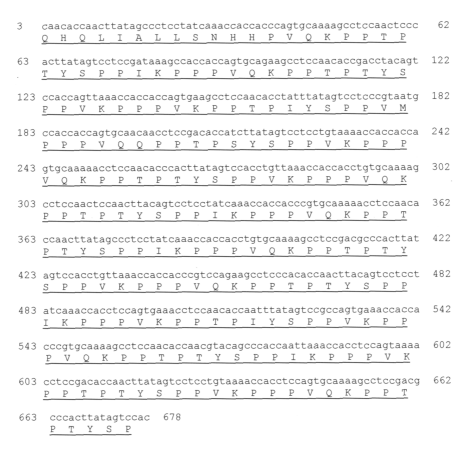


Fig. 1: Nucleotide and deduced amino acid sequence of clone Ae4 (Genbank accession no. AY570239).

The ORF is underlined. This is a truncated clone without the start and stop codons.

# Parameswari Namasivayam and David Hanke

CD825663 Translated CD830533 Translated NP_180307 Ae4	1	MRVPLIDFLRFLVLILSLSGASVAADATVKQNFNKYETDSGHAHPPPIYG	50
CD825663 Translated CD830533 Translated NP_180307	51	APPSYTTPPPPIYSPPIYPPPIQKPPTYSPPIYPPPIQKPPTPTYSPPIY	100
CD825663 Translated CD830533 Translated NP_180307	101	PPPIQKPPTPTYSPPIYPPPIQKPPTPTYSPPIYPPPIQKPPTPSYSPPV	150
CD825663 Translated CD830533 Translated NP_180307	151	KPPPVQMPPTPTYSPPIKPPPVHKPPTPTYSPPIKPPVHKPPTPIYSPPI	200
CD825663 Translated CD830533 Translated NP_180307 Ae4	201	KPPPVHKPPTPIYSPPIKPPPVHKPPTPTYSPPVKPPPVHKPPTPIYSPP	250
CD825663 Translated CD830533_Translated NP_180307	251 1	IKPPPVHKPPTPIYSPPVKPPPVQTPPTPIYSPPVKPPPVHKPPTPTYSP	300 9
CD825663 Translated CD830533_Translated NP_180307	1 1 301 10	PVKPPPVQKPPTPTYSPPVKPPPVQKPPTPTYSPPIKPPPVQKPPTPTYSMPPPVQQPPTPSYSPPVKPPPVQKPPTPTYSPPVKPPPVQKPPTPTYS PVKSPPVQKPPTPTYSPPIKPPPVQKPPTPTYSPPIKPPPV-KPPTPIYSNHHPVQKPPTPTYSPPIKPPPVQKPPTPTYSPPVKPPPV-KPPTPIYS	50 48 350 57
CD825663 Translated CD830533 Translated NP_180307	51 100 351 58	PPIKPPPVQKPPTPTYSPPIKPPPVQKPPTPTYSPPVKPPPV-NPPAPIY PPIKPPPVQKPPTPTYSPPIKPPPVQKPPTPTYSPPVKPP PPVKPPPVHKPPTPIYSPPVKPPPVHKPPTPIYSPPVKPPPIQKPPTPTY PPVMPPPVQQPPTPSYSPPVKPPPVQKPPTPTYSPPVKPPPVQKPPTPTY	139 400
CD825663 Translated	100 140	SPPVKPPPVQQPPTPSYSPPVQKPPPPVQKPPTFTY	133 149

## A Putative Proline-rich Protein of B. napus

NP_180307	401	SPPIKPPPLQKPPTPTYSPPIKLPPVKPPTPIYSPPVKPPPVHKPPTPIY	450
Ae4	108	SPPIKPPPVQKPPTPTYSPPIKPPPVQKPPTPTY	141
CD825663 Translated	134	SPPVKPPPVQKPPTPTYSPPIKPPPVQKPPTPTY	167
CD830533 Translated	150	SPPIKPPPV-KPPTPTYSPPIKPPPVQKPPTPTY	183
NP_180307	451	SPPVKPPPVHKPPTPTYSPPIKPPPVKPPTPTYSPPVQPPPVQKPPTPTY	500
Ae4	142	SPPVKPPPVQKPPTPTYSPPIKPPPVKPPTPIY	174
CD825663 Translated	168	SPPIKPPPVKKPPTPTYSPPVKPPPVQKPPTPTYSPPIKPPPVKTSKTNL	218
CD830533 Translated	184	SPEV	187
BWB4372	501	SPPVKPPPIQKPPTPTYSPPIKPPPV-KPPTPTYSPPIKPPPVHKPPT	550
Ae4	175	SPPVKPPPVQKPPTPTYSPPIKPPPV-KPPTPTYSPPVKPPPVQKPPT	222
CD825663 Translated	219	PTYKATTQCNNL	230
CD830533_Translated NP_180307	551	PTYSPPIKPPPIHKPPTPTYSPPIKPPPVHKPPTPTYSPPIKPPPVHK	
Ae4	223	PTYSP	227
CD825663 Translated CD830533 Translated NP_180307	601	TPTYSPPIKPPPVHKPPTPTYSPPIKPPPVHKPPTPTYSPPIKPPPVHK	
Ae4			
CD825663 Translated CD830533 Translated			
NP_180307	651	PTPTYSPPIKPPPVQKPPTPTYSPPVKPPPVQLPPTPTYSPPVKPPPVQ	QV 700
Ae4			
CD825663 Translated CD830533 Translated NP_180307	701	PPTPTYSPPVKPPPVQVPPTPTYSPPIKPPPVQVPPTPTTPSPPQGGY	
Ae4			
CD825663 Translated CD830533 Translated NP_180307	751	  PPPYAYLSHPIDIRN 761	

Fig. 2: Alignment of predicted amino acid sequence of Ae4 with the Arabidopsis putative proline rich protein and translated EST sequences

Shaded sequences denote identical amino acids and gaps introduced in the alignment are marked with dashes. The amino acid sequences were obtained from GenBank: Arabidopsis At2g27380 (GenBank accession no. NP\_180307), Brassica seed EST clone BN25061G23 (GenBank accession no. CD825663) and Brassica napus seed EST clone BN40045N17 (GenBank accession no. CD830533)

(*Fig. 3A*, top panel). However, hybridisation with the *Actin*2/7 cDNA probe (*Fig. 3A*, bottom panel) showed clear signals, suggesting that the RNA was not degraded. To verify these results, a RT-PCR approach was employed using *Ae4* genespecific primers and equal amount of cDNA from leaves, stems, buds, flowers, siliques, roots and carpels. After 30 amplification cycles, no product was visible on the ethidium bromide stained gel (*Fig. 3C*, top panel). Therefore, the gel was blotted and the RT-PCR gel blot was hybridised with a <sup>32</sup>P-labelled *Ae4* cDNA. An autoradiograph of this blot showed the presence of an approximately 370 bp band (as expected)

in all lanes with various intensities of hybridisation (*Fig. 3C*, bottom panel), suggesting differential expression of *Ae*4 in different organs/tissues of the mature *B. napus* plant. Also, there was an additional faint band at approximately 600 bp in lane 2, suggesting the presence of another isoform of the *Ae*4 gene or possibly an unspliced *Ae*4 transcript.

Northern analysis of the tissue culture blot (Fig. 3B, top panel) detected expression of Ae4 transcripts in the embryogenic culture, both in the pre-embryogenic (lane PEC) and mature embryogenic tissue (lane MEC). Relative to the actin control, expression in both these tissues

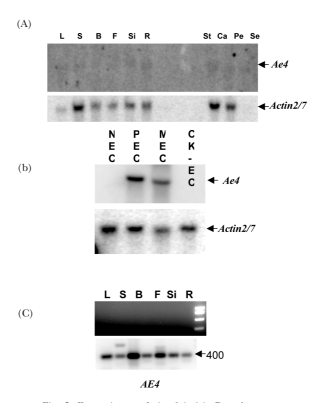


Fig. 3: Expression analysis of Ae4 in Brassica napus

- (A) Tissue/organ-specific and (B) tissue culture RNA gel blots containing 10 µg of total RNA per lane were first hybridised to <sup>32</sup>P-labelled Ae4 and then to an Arabidopsis Actin2/7 cDNA (control). Lanes: L, leaves; S, stem; B, buds; F, flowers; S, siliques; R, roots; St, stamens; Ca, carpels; Pe, petals; Se, sepals; NEC, non-embryogenic tissue; PEC, preembryogenic tissue; MEC, mature embryogenic tissue; CK-EC, cytokinin-treated tissue.
- (C) Top panel shows results of RT-PCR analysis of Ae4 gene expression. (C) Bottom panel is the RT-PCR gel blot probed with <sup>32</sup>P-labelled Ae4 cDNA. Lanes: M, 1 kb Bioline DNA ladder; L, leaves; S, stems; B, buds; F, flowers; Si, siliques; R, roots; Ca, carpels.

seems to be approximately at the same level. By contrast, Ae4 transcripts were not expressed at detectable levels in the non-embryogenic tissue (lane NEC) and the cytokinin-treated embryogenic tissue (lane CK-EC).

#### DISCUSSION

Ae4 Encodes a Partial Length Protein with Prolinerich Domain

The Ae4 cDNA encodes a partial length protein having a proline-rich domain. Both the amino acid composition and the presence of repeating motifs of proline are characteristics of prolinerich cell wall proteins (Jose-Estanyol and Puigdomenech, 2000). The 10-mer motif of the putative AE4 protein is unique in its sequence and belongs to a group of long repeat elements (Gyorgyey et al., 1997). The repeating motifs do not correspond to any of the common motifs previously identified in proline-rich proteins and extensins (Showalter, 1993, Gyorgey et al., 1997; Jose-Estanyol and Puigdomenech, 2000). Therefore, Ae4 may encode a novel class of proline-rich proteins (PRPs). Two PRP genes have already been isolated from Brassica napus, one that accumulated during pod development (Coupe et al., 1993) and the other one induced by cold treatment (Goodwin et al., 1996). However, the Ae4 sequence is not similar to either of them.

## Expression Analysis of Ae4

The expression of Ae4 transcripts in all vegetative and floral tissues examined corresponds to the observations by Fowler et al. (1999) in Arabidopsis. They reported that AtPRP2 and AtPRP4 transcripts were most abundant in the aerial parts of the plant, namely in leaves, stems, flowers and siliques. Also, AtPRP4 expression was detected in the early stages of lateral root formation. Since most PRPs are members of a multigene family, it is likely that the same will be true for those of B. napus.

Based on Northern analysis, Ae4 was detected in the pre-embryogenic but not non-embryogenic tissue and this suggests that the encoded protein may be associated with somatic embryogenesis. There have been a few PRP transcripts that have been shown to accumulate during somatic embryogenesis, such as in carrot (Aleith and Richter, 1990; Holk et al., 1996; Yasuda et al., 2001) and Medicago sativa (Gyorgey et al., 1997).

A proline-rich protein encoded by the DC 2.15 gene was identified as one of the genes that is differentially expressed during induction of somatic embryogenesis in carrot cell suspension culture (Aleith and Richter, 1990). The expression of this gene during somatic embryogenesis was detectable from 3 days after induction, and transcript abundance increased until the heart-shape stage (Aleith and Richter, 1990). This observation was further supported by promoter studies of the DC 2.15 gene (Holk et al., 1996). Also, Gyorgyey et al. (1997) reported that MsPRP5, a cDNA clone encoding a small proline-rich protein is preferentially expressed in alfalfa dedifferentiated callus cells. They proposed that the proline-rich protein may cause structural changes of the cell wall required for certain switches in function by plant cells.

He et al. (2002) reported that expression of a soybean PRP gene was inhibited by treatment with kinetin. This is consistent with our observation from the Northern analysis of the tissue culture blot that Ae4 transcripts could not be detected in cytokinin-treated tissue. This suggests that cytokinin treatment has suppressed Ae4 transcription to undetectable levels, which correlates with the suppression of secondary embryogenesis (Loh et al., 1983). Alternatively, Ae4 transcripts could be down regulated due to the low rate of secondary embryogenesis in the cytokinin-treated tissue.

## Potential Roles of Ae4 in Brassica napus

Proline-rich proteins have been thought to provide strength and rigidity in the cell wall by forming covalently cross-linked networks with cell wall components (Showalter, 1993). PRPs have a relatively high content of tyrosine and lysine residues which have been implicated as the substrate for the peroxidase-mediated insolubilisation of PRPs in soybean via isodityrosine crosslinks (Kleis-San Francisco and Tierney, 1990; Bradley et al., 1992; Brisson et al., 1994). Insolubilisation of PRPs in the cell wall occurs as a rapid response to wounding and treatment with fungal elicitors. PRPs are rapidly insolubilized within the cell wall in response to physical damage, treatment with fungal elicitors, and pathogen infection (Kleis-San Francisco and Tierney, 1990; Bradley et al., 1992; Brisson et al., 1994), indicating an active role in plant defence reactions. Thus, it was proposed that the PRPs

contribute to the cell wall structure of specific cell types based on their patterns of gene expression during plant development and induction by biotic and abiotic stresses. The enhanced expression of Ae4 transcripts in the pre-embryogenic and mature embryogenic tissues suggests that Ae4 may have a role during embryo formation. In carrot cultures, it was suggested that two PRPs encoded by No.93 and DC 2.15 might act as extracellular signal factors during the development of somatic embryos (Yasuda et al., 2001). However, there is no clear evidence for PRPs as signalling molecules that induce somatic embryogenesis.

Assuming that Ae4 encodes a proline-rich protein, we propose that the expression of Ae4 transcripts preferentially in embryogenic tissue could be possibly to provide mechanical strength to the embryonic cells that will protect the cell during later phase of embryo development in vitro. More experiments such as isolation of the full-length sequence of Ae4, immunolocalization and transgenic studies, especially promoter analysis, are required to provide more information on possible biological functions of the Ae4 gene. Also, the inducibility of the Ae4 gene in response to abiotic and biotic stresses in B. napus should be tested to explore its regulation.

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