

1 Original article

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3 **Scots pine aminopropyltransferases shed new light on evolution of polyamine**  
4 **biosynthesis pathway in seed plants**

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6 Jaana Vuosku<sup>1,\*</sup>, Katja Karppinen<sup>1</sup>, Riina Muilu-Mäkelä<sup>2</sup>, Tomonobu Kusano<sup>3</sup>, G.H.M. Sagor<sup>3</sup>,  
7 Komlan Avia<sup>1</sup>, Emmi Alakärppä<sup>1</sup>, Johanna Kestilä<sup>1</sup>, Marko Suokas<sup>1</sup>, Kaloian Nickolov<sup>1</sup>, Leena  
8 Hamberg<sup>4</sup>, Outi Savolainen<sup>1</sup>, Hely Häggman<sup>1</sup> and Tytti Sarjala<sup>2</sup>

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10 <sup>1</sup>*University of Oulu, Department of Ecology and Genetics, Oulu, Finland*

11 <sup>2</sup>*Natural Resources Institute Finland, Bio-based Business and Industry, Parkano, Finland*

12 <sup>3</sup>*Tohoku University, Graduate School of Life Sciences, Sendai, Miyagi, Japan*

13 <sup>4</sup>*Natural Resources Institute Finland, Management and Production of Renewable Resources,*  
14 *Vantaa, Finland*

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16 **Running title:** Evolution and function of Scots pine aminopropyltransferases

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18 **\*For correspondence.** E-mail [jaana.vuosku@oulu.fi](mailto:jaana.vuosku@oulu.fi)

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## 1 **ABSTRACT**

- 2 • **Background and Aims** Polyamines are small metabolites present in all living cells and play  
3 fundamental roles in numerous physiological events in plants. The aminopropyltransferases  
4 (APTs), spermidine synthase (SPDS), spermine synthase (SPMS) and thermospermine  
5 synthase (ACL5), are essential enzymes in the polyamine biosynthesis pathway. In  
6 angiosperms, SPMS has evolved from SPDS via gene duplication, whereas in gymnosperms  
7 APTs are mostly unexplored and no *SPMS* gene has been reported. The present study aimed to  
8 investigate the functional properties of the SPDS and ACL5 proteins of Scots pine (*Pinus*  
9 *sylvestris* L.) in order to elucidate the role and evolution of APTs in higher plants.
- 10 • **Methods** Germinating Scots pine seeds and seedlings were analyzed for polyamines by HPLC  
11 and the expression of *PsSPDS* and *PsACL5* genes by *in situ* hybridization. Recombinant  
12 proteins of PsSPDS and PsACL5 were produced and investigated for functional properties.  
13 Also gene structures, promoter regions and phylogenetic relationships of *PsSPDS* and *PsACL5*  
14 genes were analyzed.
- 15 • **Key Results** Scots pine tissues were found to contain spermidine, spermine and  
16 thermospermine. PsSPDS enzyme catalyzed synthesis of both spermidine and spermine.  
17 PsACL5 was found to produce thermospermine and *PsACL5* gene expression was localized in  
18 the developing procambium in embryos and tracheary elements in seedlings.
- 19 • **Conclusions** Contrary to previous view, our results demonstrate that SPMS activity is not a  
20 novel feature developed solely in the angiosperm lineage of seed plants but also exist as a  
21 secondary property in the Scots pine SPDS enzyme. The discovery of bifunctional SPDS from  
22 an evolutionary old conifer reveals the missing link in the evolution of the polyamine  
23 biosynthesis pathway. The finding emphasizes the importance of pre-existing secondary

1 functions in the evolution of new enzyme activities via gene duplication. Our results also  
2 associate PsACL5 with the development of vascular structures in Scots pine.

3  
4 **Key words:** aminopropyltransferase, gene duplication, gymnosperm, pathway evolution,  
5 *Pinus sylvestris*, polyamine biosynthesis, Scots pine, spermidine synthase, spermine synthase,  
6 thermospermine synthase

## 8 INTRODUCTION

9 Polyamines (PAs) are ancient low molecular weight polycations which are found in all living  
10 organisms. In addition to the universal occurrence, the fundamental role of PAs in cell  
11 metabolism is emphasized by the conservation of PAs across evolution as well as by the  
12 complexity of compensatory mechanisms that are invoked to maintain PA homeostasis in cells  
13 (Wallace *et al.*, 2003). The cellular functions of PAs may result from their unique chemical  
14 characteristics, from the structural changes of RNA when PA-RNA complexes are formed or  
15 from the PA-dependent alterations in the structure and function of DNA and further in gene  
16 expression (Childs *et al.*, 2003; Igarashi and Kashiwagi, 2010). In plants, PAs have been  
17 considered as growth regulators or as secondary messengers (Kakkar and Sawhney, 2002) and  
18 they are involved in several physiological events, such as organogenesis, embryogenesis, floral  
19 initiation and development, leaf senescence, fruit development and ripening as well as in abiotic  
20 and biotic stress responses (Galston and Sawhney, 1990; Kumar *et al.*, 1997; Malmberg *et al.*,  
21 1998; Bouchereau *et al.*, 1999; Bagni and Tassoni, 2001; Alcázar *et al.*, 2006; Kusano *et al.*,  
22 2008).

23 The major naturally occurring PAs are diamine putrescine (Put), triamine spermidine  
24 (Spd) and tetra-amine spermine (Spm) (Tiburcio *et al.*, 1997). In addition, thermospermine

1 (tSpm), a less abundant structural isomer of Spm, is widely detected in plant kingdom (Takano *et*  
2 *al.*, 2012). In mammalian cells, Put is synthesized from ornithine by ornithine decarboxylase  
3 (ODC; EC 4.1.1.17) and also in many fungi ODC is the sole pathway for Put production  
4 (Coleman *et al.*, 2004). Plants are able to produce Put also from arginine by arginine  
5 decarboxylase (ADC; EC 4.1.1.19) through two intermediate stages (Tiburcio *et al.*, 1997) and  
6 there is also evidence of ADC activity in certain fungal species (Khan and Minocha, 1989;  
7 Biondi *et al.*, 1993; Sannazzaro *et al.*, 2004). Put can be metabolized further to the higher PAs by  
8 the aminopropyltransferases (APTs), which transfer aminopropyl moiety (derived from  
9 decarboxylated S-adenosylmethionine) to an amine acceptor on another PA (Shao *et al.*, 2012)  
10 (Supplementary Data Fig. S1). Spd is formed from Put by spermidine synthase (SPDS; EC  
11 2.5.1.16). It has been suggested that the extant core of the eukaryotic PA biosynthetic pathway in  
12 the last eukaryotic common ancestor consisted of the ODC and SPDS enzymes, because Put and  
13 Spd are the only PAs produced in all PA synthesizing eukaryotes (Michael, 2016). Spd is an  
14 unsymmetrical molecule, which can be further aminopropylated at either end forming Spm or  
15 tSpm. Spermine synthase (SPMS; EC 2.5.1.22) transfers an aminopropyl group to the N<sup>8</sup>-  
16 (aminobutyl) end of Spd to make a symmetrical Spm molecule, whereas thermospermine  
17 synthase (ACL5; EC 2.5.1.B4) transfers an aminopropyl group to the N<sup>1</sup>-(aminopropyl) end of  
18 Spd which results in an unsymmetrical tSpm molecule (Pegg and Michael, 2010) (Supplementary  
19 Data Fig. S1). To distinguish Spm and tSpm, benzylation of the PAs is needed followed by  
20 separation with HPLC (Naka *et al.*, 2010).

21 Several reports have indicated the necessity of Spd for the viability in eukaryotes, whereas  
22 eukaryotes deficient in SPMS and thus lacking Spm are mostly viable, even if they may suffer  
23 different degrees of dysfunctions (Pegg and Michael, 2010). One of the essential functions of Spd  
24 is its role as a substrate for the hypusine modification of the eukaryotic translation initiation

1 factor eIF5A (Chattopadhyay *et al.*, 2003, 2008; reviewed by Wolff and Park, 2015). In  
2 Arabidopsis (*Arabidopsis thaliana* L.), the *spds1-1 spds2-1* double-mutant had abnormally  
3 shrunken seeds and embryos that were arrested morphologically at the heart-torpedo transition  
4 stage (Imai *et al.*, 2004b). In contrast, Spm was not essential for survival of Arabidopsis under  
5 normal growth conditions (Imai *et al.*, 2004a), but Spm-deficient mutants were hypersensitive to  
6 drought and salt stress (Yamaguchi *et al.*, 2006, 2007). The loss-of-function mutations of the  
7 *AtACL5* gene resulted in a severely dwarfed phenotype in Arabidopsis due to a specific defect in  
8 the growth of stem internodes (Hanzawa *et al.*, 2000). These observations suggest that Spd may  
9 act in the basic metabolism of eukaryotic cells and is essential for cell viability, whereas tSpm  
10 may play a role in the development of the structures of multicellular eukaryotes and Spm may  
11 improve the tolerance to different kinds of stresses. Thus, the slight structural difference between  
12 tSpm and Spm seems to be important for their function.

13 APTs share a high degree of sequence similarity at the putative active sites even if they  
14 display different substrate specificities (Rodríguez-Kessler *et al.*, 2010). In plants, SPMS seems  
15 to have evolved via gene duplication and change of the function of an ancestral SPDS in  
16 angiosperms, whereas the *ACL5* gene may have been acquired by an algal ancestor of plants  
17 through a horizontal gene transfer from bacteria or archaea (Minguet *et al.*, 2008). All plant  
18 species, including gymnosperms and *Physcomitrella patens* (Hedw.) Bruch & Schimp, have been  
19 found to possess sequences identifiable as *SPDS* or *ACL5*, but no *SPMS* genes have been reported  
20 from gymnosperms so far (Minguet *et al.*, 2008).

21 The APTs in angiosperms are well known (reviewed by Shao *et al.*, 2012), whereas APTs  
22 in gymnosperms have mostly been an unexplored area. *Pinus* species of the gymnosperms form  
23 an evolutionarily old group of vascular plants that shared a common ancestor with angiosperms  
24 about 285 million years ago (Bowe *et al.*, 2000). In the present study, we hypothesized that at

1 some point of seed plant evolution both SPDS and SPMS activities might coexist in a single  
2 protein and, therefore our focus was specifically on the *SPDS* and *ACL5* genes of Scots pine  
3 (*Pinus sylvestris* L.). We produced recombinant proteins of PsSPDS and PsACL5 to investigate  
4 their functional properties. We also studied the expression of *PsSPDS* and *PsACL5* genes in  
5 Scots pine seeds and seedlings by mRNA *in situ* hybridization. Furthermore, gene structures,  
6 promoter regions and phylogenetic relationships of *PsSPDS* and *PsACL5* genes were analyzed in  
7 order to elucidate the evolution of APTs in higher plants.

8

## 9 **MATERIALS AND METHODS**

### 10 *Plant material*

11 Mature Scots pine seeds were sterilized overnight in 3 % Plant Preservative Mixture TM (Plant  
12 Cell Technology, Washington, DC, USA). Seeds were germinated on moist filter papers in petri  
13 dishes in a growth chamber at +20 °C, in 100 % moisture and under continuous light. For the PA  
14 analyses, the embryos and megagametophytes of the seeds were excised after 2 days imbibition,  
15 and the cotyledons, hypocotyls and roots of the seedlings were excised after 16 days of  
16 germination. The embryos for RNA extraction were frozen in liquid nitrogen and stored at -80 °C  
17 until use. For the *in situ* hybridization assays of *PsSPDS* and *PsACL5* transcripts, seeds (without  
18 seed coat) and hypocotyls were fixed immediately after sampling as described previously in  
19 Vuosku *et al.* (2015).

20 Three populations from different latitudes were used to sequence the *PsSPDS* and *PsACL5*  
21 genes: Kolari (northern Finland, latitude 67°11'N, 24°03'E), Punkaharju (southern Finland,  
22 latitude 61°48'N, 29°19'E) and Radom (Poland, latitude 50°41'N, 20°05'E). Seeds collected  
23 from eight open-pollinated (mostly half-sibs) families of each population were used to sequence  
24 the genes as well as their promoter regions.

1

## 2 *PA analyses in pine tissues with benzylation method*

3       The presence of PAs (Put, Spd, Spm and tSpm) in Scots pine tissues was discovered by  
4 using benzylation method described by Naka *et al.* (2010). The cotyledon, hypocotyl, root,  
5 embryo and megagametophyte samples which were pooled from many seedlings or seeds were  
6 extracted in 5 % (w/v) HClO<sub>4</sub> and the concentration of the benzylated PAs in the tissue samples  
7 and in the reaction mixture was separated by HPLC and detected at 254 nm according to Naka *et*  
8 *al.* (2010). The standard for tSpm was kindly provided by Prof. Masaru Niitsu.

9

## 10 *Production and functional studies of PsSPDS and PsACL5 recombinant proteins*

11       Total RNA was extracted from embryos using the automatic magnetic-based KingFisher™  
12 mL method (Thermo LabSystems, Helsinki, Finland) with the MagExtractor® total RNA  
13 purification kit (Toyobo, Osaka, Japan) according to the manufacturer's instructions. The RNA  
14 samples were treated with RNase-free DNase (Invitrogen, Carlsbad, CA, USA) and purified with  
15 the NucleoSpin® RNA Clean-Up kit (Macherey-Nagel, Düren, Germany). RNA yields were  
16 measured by NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE,  
17 USA). cDNA was reverse-transcribed from 1 µg of total RNA by using the SuperScript® VILO™  
18 cDNA Synthesis kit (Invitrogen) with the SuperScript® III Reverse Transcriptase in a reaction  
19 volume of 20 µl.

20       The coding region of *PsACL5* was amplified by PCR, using upstream primer 5'-  
21 TATGCCATGGGGGAAGTAGCAGC-3' (the translation start codon is in bold and the *NcoI* site  
22 is underlined) and downstream primer 5'-TATGCGGCCGCTTATGATTATGGCAC-3' (the  
23 translation stop codon is in bold and the *NotI* site is underlined). The amplified PCR product was  
24 restricted with *NcoI* and *NotI* and ligated into the *NcoI/NotI* site of the expression vector pET-32a

1 (Novagen, Madison, WI, USA). The coding region of *PsSPDS* was amplified by using upstream  
2 primer 5'-TAGATATCGGATCCATGGCCGAGAAC-3' (the translation start codon is in bold  
3 and the *Bam*HI site is underlined) and downstream primer 5'-  
4 TATGCGGCCGCTTAAGATAATGGTGG-3' (the translation stop codon is in bold and the *Not*I  
5 site is underlined). The amplified PCR product was restricted with *Bam*HI and *Not*I and ligated  
6 into the *Bam*HI/*Not*I site of the expression vector pET-32a. The resulting recombinant plasmids  
7 were confirmed by sequencing using an ABI 3730 DNA sequencer (Applied Biosystems, Foster  
8 City, CA, USA) with a BigDye Terminator Cycle Sequencing Kit (Applied Biosystems). The  
9 *PsSPDS* and *PsACL5* sequence has been deposited in GenBank under accession numbers  
10 KX761190 and HM236828, respectively.

11 The recombinant plasmids were transferred into *Escherichia coli* BL21 (DE3) cells  
12 (Novagen). The cells were grown in Luria-Bertani liquid medium in the presence of ampicillin  
13 (100 µg/ml) at 30 °C until the  $D_{600}$  of the culture reached 0.6. Protein expression was induced by  
14 0.5 mM IPTG and the cultures were further incubated at 16 °C for 18 hours. The *E. coli* cells  
15 were harvested by centrifugation (4000 g for 20 min) and resuspended in buffer (50 mM sodium  
16 phosphate buffer, pH 7.0, containing 600 mM NaCl, 15 mM  $\beta$ -mercaptoethanol, 10 % glycerol, 1  
17 % Tween 20 and 20 mM imidazole). The cells were disrupted using lysozyme (1 mg ml<sup>-1</sup>) and  
18 sonication (Type UP50H; Dr. Hielscher GmbH, Teltow, Germany). The lysate was centrifuged at  
19 17 000 g for 30 min and the supernatant was collected for purification of recombinant protein  
20 using Ni-NTA agarose (Qiagen, Hilden, Germany). Unbound proteins were washed away with  
21 the buffer described above after which the recombinant proteins were eluted with an elution  
22 buffer (50 mM sodium phosphate buffer, pH 7.0, containing 600 mM NaCl, 15 mM  $\beta$ -  
23 mercaptoethanol, 10 % glycerol and 250 mM imidazole). The purified enzyme solution was



1 buffer-exchanged to assay buffer (0.1 M Tris-HCl, pH 7.5, containing 15 % glycerol) using a PD-  
2 10 column (GE Healthcare Bio-Sciences, Little Chalfont, UK). The purity of the recombinant  
3 proteins was verified by SDS-PAGE using a Mini-Protean II electrophoresis system (Bio-Rad,  
4 Hercules, CA, USA). Proteins were visualized with Coomassie Brilliant Blue R-250 (Merck,  
5 Darmstadt, Germany).

6 PA analyses were performed for confirming the SPDS/SPMS activity of the purified  
7 PsSPDS protein. The assay mixture (200  $\mu$ l) contained 0.1 M Tris-HCl (pH 7.5), 5 mM  
8 decarboxylated S-adenosyl-L-methionine (dcSAM, kindly provided by Prof. Keiji Samenima),  
9 substrate (5 mM Put or Spd) and 34  $\mu$ g purified PsSDPD protein. The reaction mixtures were  
10 incubated for two hours at 30 °C, 40 °C, or 50 °C. After two hours incubation, 16  $\mu$ l of 60 %  
11 (w/v) HClO<sub>4</sub> was added into 200  $\mu$ l of the reaction mixture and analysed immediately for PAs or  
12 stored at -80 °C until analysed. For PA analyses, the samples were dansylated (Flores and  
13 Galston, 1982), stored at -20 °C and analysed within one week with HPLC (Merck Hitachi,  
14 Darmstadt, Germany) and fluorescence spectrophotometry (Merck Hitachi F-1050) at excitation  
15 and emission wavelengths of 365 nm and 510 nm, respectively. The PAs were separated by using  
16 LiChrospher 100 RP-18 5  $\mu$ m column and methanol-water gradient as described by Sarjala and  
17 Kaunisto (1993). Putrescine dihydrochloride (Sigma-Aldrich, St. Louis, MO, USA), spermidine  
18 trihydrochloride (Sigma-Aldrich) and spermine tetrahydrochloride (Sigma-Aldrich) were used as  
19 standards.

20 The assay of the tSpm activity of the purified PsACL5 protein was performed in the  
21 reaction mixture (200  $\mu$ l) containing 0.1 M Tris-HCl (pH 7.5), 100 nM Spd, 100 nM dcSAM  
22 (kindly provided by Prof. Yoshihiko Ikeguchi) and 14  $\mu$ g purified enzyme. The reaction mixture  
23 was incubated for one hour at 30 °C, 40 °C, and 50 °C. The presence of PAs in the reaction  
24 mixture was discovered by using benzylation method as described by Naka *et al.* (2010).

1  
2 *mRNA in situ hybridization assay of PsSPDS and PsACL5 transcripts*  
3       The localization of *PsSPDS* and *PsACL5* transcripts in Scots pine seeds and hypocotyls was  
4 performed as described in Vuosku *et al.* (2015). The primers used for the synthesis of a 275 bp  
5 probe for *PsSPDS* transcripts were 5'-CACATGCCCATCATTGAAGA-3' (upstream) and 5'-  
6 CCTATCGCCCTTCTAGCAAA-3' (downstream). The 332-bp long *ACL5* probe was amplified  
7 with the primers 5'-GCCGAGCTCGAGAGTAGAGA-3' (upstream) and 5'-  
8 TCGATTTCTTCGGCGTCTAT-3' (downstream). In seed sections, the DIG-labelled probes  
9 were detected by Anti-DIG-AP Fab fragments and NBT/BCIP substrate (Dig Nucleic acid  
10 detection Kit, Roche Molecular Biochemicals, Mannheim, Germany) as previously described in  
11 Vuosku *et al.* (2015). In hypocotyl sections the probes were visualized by incubating the slides  
12 for 30 min at 37 °C with 1:200 anti-DIG-rhodamine Fab fragments (Roche Molecular  
13 Biochemicals) in the dark. Thereafter, the slides were washed four times in Tris/NaCl buffer for  
14 10 min, dehydrated in a graded series of ethanol, air-dried and mounted with coverslips using  
15 Vectashield mounting medium (Vector Laboratories, Burlingame, CA, USA). The hypocotyl  
16 sections were examined in a confocal laser scanning microscope (LSM 5 Pascal, Carl Zeiss) with  
17 an HBO 103 mercury lamp. Adobe Photoshop CS was used to adjust contrast, brightness and  
18 color uniformly to entire images.

19  
20 *Analysis of PsSPDS and PsACL5 gene structures and copy numbers*

21       Primers (Supplementary Data Table S1) were designed to sequence full *PsSPDS* and  
22 *PsACL5* genes and their promoters. Total genomic DNA was extracted from the haploid  
23 megagametophyte tissue of the seeds (see Plant material) with the Nucleospin Plant II  
24 (Macherey-Nagel) with buffer PL1. DNA quality and quantity were checked with agarose gel

1 electrophoresis and a NanoDrop ND1000 spectrophotometer. All of the fragments belonging to  
2 the same gene were amplified from a single megagametophyte. The MinElute PCR purification  
3 kit (Qiagen, Chatsworth, CA, USA) or the FastAP Thermosensitive Alkaline Phosphatase and  
4 Exonuclease I (Fermentas, Vilnius, Lithuania) were used to purify the PCR products. The  
5 Genome Walker Universal kit (Clontech, Mountain View, CA, USA) was used according to the  
6 manufacturer's instructions to increase the length of the promoter regions. Sequencing reactions  
7 were carried out with an ABI Prism 3730 DNA Analyzer (Applied Biosystems) with Big Dye  
8 Terminator kit v3.1 (Applied Biosystems). Following sequencing, manual checking and edition  
9 were done in Sequencher 4.7 (Gene Codes Corporation). Sequences were aligned using  
10 MUSCLE (Edgar, 2004) in Geneious version 6.1 (created by Biomatters, available from  
11 <http://www.geneious.com>). The genomic sequences of *PsSPDS* and *PsACL5* have been deposited  
12 in GenBank under accession numbers KX788070 and KX788071, respectively.

13 No fully sequenced genome of Scots pine is available to date. Thus, in order to predict the  
14 copy number of the *SPDS* and *ACL5* genes in the Scots pine genome, the current assembly  
15 (version 2.01 in March 2017) of the loblolly pine (*Pinus taeda* L.) genome was downloaded from  
16 the UCDavis Dendrome website  
17 ([http://dendrome.ucdavis.edu/ftp/Genome\\_Data/genome/pinerefseq/Pita/](http://dendrome.ucdavis.edu/ftp/Genome_Data/genome/pinerefseq/Pita/)) and used for full  
18 BLAST search on local server. The results of the BLAST search were analyzed for the  
19 occurrences of complete gene sequences of *SPDS* and *ACL5* using the Geneious software. To  
20 confirm that the best-matching sequences were putative *SPDS* and *ACL5* proteins, BLAST  
21 searches were performed against the NCBI nr or UniProtKB/Swiss-Prot databases. In addition,  
22 the deduced *PsSPDS* and *PsACL5* proteins were used for BLAST search against loblolly pine  
23 transcripts and protein sequences.

24

1 *Functional analysis of SPDS and SPMS promoter regions*

2       The genomic sequences of 3 kb upstream of the SPDS and SPMS genes in Arabidopsis  
3 (*Arabidopsis thaliana* L.) and black cottonwood (*Populus trichocarpa*) were extracted from The  
4 Arabidopsis Information Resource (TAIR; <http://www.arabidopsis.org/>) and The Populus  
5 Genome Integrative Explorer (PopGenIE; <http://popgenie.org/>), respectively. The promoter  
6 sequence of *PsSPDS* (KX788070, 1201 bp) was extended to 3 kb with the 1810 bp promoter  
7 sequence of loblolly pine from conifer Genome Integrative Explorer (ConGenIE;  
8 <http://congenie.org/>). Transcription factor binding sites (TFBS) and the corresponding  
9 transcription factors (TF) within the promoter regions of *SPMS* and *SPDS* genes as well as their  
10 functions were searched using PlantPAN2.0 database (<http://plantpan2.itps.ncku.edu.tw>) (Chang  
11 *et al.*, 2008; Chow *et al.*, 2016). The comparison of the presence and absence of the identified  
12 *cis*-regulatory elements in the promoter regions of the *SPDS* and *SPMS* genes was carried out  
13 manually by generating presence-absence data (Supplementary Data Table S2). The functions of  
14 TFBS/TF were determined in the cases where a TFBS was present in *PsSPDS* but was absent in  
15 at least one of the Arabidopsis or black cottonwood *SPDS/SPMS* genes as well as in the cases  
16 where a TFBS was absent in *PsSPDS* (Supplementary Data Table S3). The similarity of the  
17 TFBS content among the promoter regions of the *SPDS* and *SPMS* genes in Scots pine,  
18 Arabidopsis and black cottonwood, was investigated with global non-metric multidimensional  
19 scaling (GNMDS) in the library vegan (Oksanen *et al.*, 2016) of a statistical program R (R Core  
20 Team, 2015) using presence-absence data. The analysis was performed with 50 separate random  
21 starts with function metaMDS using dissimilarity index Raup-Crick.

22

23 *Phylogenetic analysis of plant SPDS, SPMS and ACL5 genes*

1 For the phylogenetic analysis of plant APT genes, BLAST searches against NCBI  
2 databases (<http://www.ncbi.nlm.nih.gov>) were carried out against the coding sequences of the  
3 *PsSPDS* and *PsACL5* genes. In the case of other conifers, EST information was used to  
4 reconstruct a contig containing the complete coding sequence, when no unigene sequences were  
5 available. In addition, sequences were chosen on the basis of previous literature to the  
6 phylogenetic analysis. The nucleotide sequence alignments of APT genes were performed with  
7 ClustalX (Thompson *et al.*, 1997) (Supplementary Data Fig. S2). The phylogenetic trees were  
8 inferred using the maximum likelihood (ML) method based on the Tamura-Nei model (Tamura  
9 and Nei, 1993). Initial trees for the heuristic search were obtained by applying the neighbor-  
10 joining method to a matrix of pairwise distances estimated using the maximum composite  
11 likelihood (MCL) approach. All codon positions were included. The positions containing gaps  
12 and missing data were eliminated from the dataset. The bootstrap method (Felsenstein, 1985)  
13 with 1000 replicates was used to evaluate the confidence of the reconstructed trees. Bootstrap  
14 values between 70 % and 100 % have been suggested to indicate significant support for a branch  
15 (Soltis and Soltis, 2003). Phylogenetic analyses were conducted in MEGA 5.05 (Tamura *et al.*,  
16 2011).

17

## 18 **RESULTS**

### 19 *Scots pine tissues contain both Spm and tSpm*

20 The PA content was measured in several tissues of Scots pine seedlings and seeds. Especially, we  
21 were interested in the copresence of Spm and tSpm. Both Spm and tSpm were detected in the  
22 cotyledons, hypocotyls and roots of seedlings as well as in the embryos and megagametophytes  
23 of mature seeds. tSpm concentration of the tissues varied from 0.3 to 7.7 nmol g<sup>-1</sup> (Table 1). The

1 content of tSpm was 2-31 % of the content of Spm. In embryos and megagametophytes the  
2 proportion of tSpm was higher than in other Scots pine tissues.

3

#### 4 *Identified Scots pine SPDS and ACL5 proteins*

5 The identified *P. sylvestris* SPDS gene (*PsSPDS*) had an open reading frame (ORF) of 1038  
6 bp coding a protein of 345 amino acids with a calculated molecular weight of 37.90 kDa and a  
7 theoretical *pI* of 4.94. The predicted amino acid sequence of *PsSPDS* showed high similarity with  
8 the SPDS proteins of angiosperms. *PsSPDS* shared 72 % identity with SPDSs identified from rice  
9 (*Oryza sativa* L.) and clementine (*Citrus clementina* Hort. ex Tan.), 71 % with SPDS of black  
10 cottonwood (*Populus trichocarpa*), 70 % with SPDS1 of apple (*Malus x domestica*), and 67 %  
11 identity with Arabidopsis SPDS1 (Supplementary Data Fig. S3). The *PsSPDS* sequence  
12 contained conserved motifs shown to be binding sites for SAM and dcSAM in all members of the  
13 APT family.

14 The *P. sylvestris* ACL5 gene (*PsACL5*) had an ORF of 1125 bp that is predicted to encode  
15 protein of 374 amino acids with a calculated molecular weight of 41.78 kDa and a *pI* of 5.98. The  
16 predicted amino acid sequences of *PsACL5* showed 66 % identity with ACL5 identified earlier in  
17 black cottonwood, 61 % with ACL5 of clementine, 60 % with ACL5 proteins of rice and apple  
18 (*M. sylvestris*), and 59 % identity with the Arabidopsis ACL5 protein (Supplementary Data Fig.  
19 S4). Moreover, the *PsACL5* sequence contained the conserved motifs of the APT family.

20

#### 21 *Bifunctional PsSPDS enzyme produces both Spd and Spm*

22 To study the enzymatic functions of the proteins encoded by *PsSPDS* and *PsACL5*, the  
23 coding regions of the genes were expressed in *Escherichia coli* strain BL21 with pET-32a  
24 expression vector. The SDS-PAGE analysis of the purified recombinant proteins resulted in

1 bands with a molecular mass of ~55 kDa for protein encoded by *PsSPDS* and ~60 kDa for protein  
2 encoded by *PsACL5* (Fig. 1A) which coincides with the calculated molecular masses of  
3 recombinants with N-terminal His-tag (SPDS protein, 55.5 kDa; ACL5 protein, 58.9 kDa).

4 We found that purified PsSPDS catalyzes the synthesis of both Spd and Spm while the  
5 enzyme encoded by PsACL5 produced tSpm but not its isomer Spm (Fig. 1B). The PsSPDS  
6 enzyme reaction mixture containing Put as a substrate produced Spd, whereas the PsSPDS  
7 enzyme reaction mixture containing Spd as a substrate produced Spm (Fig. 1C). Raising the  
8 incubation temperature from 30 °C to 50 °C significantly increased Spm formation (p-value  
9 0,00015). Spd or Spm were not produced in the control reactions lacking PsSPDS enzyme  
10 (Fig.1B). Incubation of reaction mixture with the PsACL5 protein confirmed that tSpm was  
11 produced in all incubation temperatures at 30 °C, 40 °C and 50 °C (Supplementary Data Fig. S5).

12

### 13 *PsSPDS and PsACL5 expression in developing vascular structures*

14 In the Scots pine seed, the embryo lies within the corrosion cavity of the megagametophyte,  
15 which houses the majority of the storage reserves of the seed. In the germinating embryo, intense  
16 *PsSPDS* expression was localized in the procambial cells continuing from cotyledons to the  
17 hypocotyl, but slight *PsSPDS* expression was found in all cells of the embryo (Fig. 2A).  
18 Procambium consists of a few layers of tightly packed cells, where *PsSPDS* also expressed  
19 during cell division (Fig. 2B-E). Unlike the *PsSPDS* gene, *PsACL5* was found to express  
20 specifically only in procambial cells in the embryo (Fig. 2F, G). The specificity of the antisense  
21 *PsSPDS* and *PsACL5* probes was confirmed by the absence of signals in the sections hybridized  
22 with the sense *PsSPDS* and *PsACL5* probes, respectively (Fig. 2H, I). Non-specific hybridization  
23 signal generated by fragmented nucleic acids was observed in the embryo surrounding region  
24 (ESR) of the megagametophyte as previously described in Vuosku *et al.* (2010).

1 In the hypocotyls of Scots pine seedlings, both *PsSPDS* and *PsACL5* were expressed  
2 specifically in developing tracheary elements (Fig. 3A, B, D, E). No signals were detected in the  
3 hypocotyl sections hybridized with the sense *PsSPDS* (Fig. 3C) and *PsACL5* (Fig. 3F) probes.

4

#### 5 *Conserved exon-intron structure of PsSPDS and PsACL5 genes*

6 The full genomic sequence of *PsSPDS* contained nine exons (Fig. 4A) versus ten for  
7 *PsACL5* (Fig 4B). *PsSPDS* harbored overall larger introns leading to gene size of between 5995  
8 bp and 6135 bp for *PsSPDS* versus between 2641 bp and 2653 bp for *PsACL5* from start codon to  
9 stop codon in the 24 samples we sequenced. The intron insertion phases were also very different  
10 between the two genes.

11 The comparison of the exon - intron structure between *PsSPDS* and the *SPDS* genes of  
12 angiosperms revealed that the gene structure has been conserved during the evolution in seed  
13 plants, suggesting an orthologous origin for the genes. The orthologs accumulate much fewer  
14 structural differences unlike the paralogs (Xu *et al.*, 2012). All *SPDS* orthologs harbored nine  
15 exons with same sizes except for exons I and IX, but the size of the large introns was highly  
16 variable (Fig. 4A). For the *ACL5* orthologs, with the exception of *AtACL5*, the situation was  
17 similar, with ten exons of identical sizes except for exons I and X (Fig. 4B). Considering the very  
18 long divergence time between gymnosperms and angiosperms, the conserved gene structure  
19 among seed plants seems remarkable suggesting the important role of these APTs.

20 BLAST search against the assembly version v2.01 of the loblolly pine (*Pinus taeda* L.)  
21 genome resulted in one full-length copy of *ACL5* (in scaffold 80063) and one full-length copy of  
22 *SPDS* (in scaffold 114828). The full-length *PtACL5* showed 95 % sequence identity with the full-  
23 length *PsACL5* while CDS regions showed 98.9 % sequence identity. Likewise, there were 97 %  
24 and 98.9 % sequence identities between the full-length genes and the CDSs of loblolly pine and



1 Scots pine *SPDS*, respectively. BLAST search against the loblolly pine transcript and protein  
2 sequences also resulted in one sequence for *PsSPDS* (PITA\_000022967-RA). No result was  
3 obtained for *PsACL5*; however, the transcript and protein sequences were available only for the  
4 genome assembly version v1.01 and this situation may evolve with subsequent updates. Together,  
5 these results suggest that there are only single copies of the *PsSPDS* and *PsACL5* genes.

6

### 7 *Transcription factor binding site composition of PsSPDS promoter region*

8 In order to get an overview about the roles of bifunctional *PsSPDS* in Scots pine and the  
9 separate *SPDS* and *SPMS* genes in angiosperms, putative transcription factor binding sites  
10 (TFBSs) were searched from their promoter regions. The function of TFBSs, when possible, was  
11 found out by literature search. Promoter regions of the Scots pine *PsSPDS* gene and the *SPDS*  
12 and *SPMS* genes of black cottonwood (*PtSPDS* and *PtSPMS*) and Arabidopsis (*AtSPDS* and  
13 *AtSPMS*) were used for the analyses. In the presence-absence data of the total 81 TFBSs detected  
14 in the promoter regions of the *SPDS* and *SPMS* genes, 16 TFBSs were not present in the *PsSPDS*  
15 promoter (Supplementary Data Table S2). Instead, all TFBSs found from the *PsSPDS* promoter  
16 exist also at least in one gene in angiosperms. None of the detected TFBSs were present only in  
17 the *SPMS* promoters. The functions of the angiosperm-specific TFBSs were mostly related to  
18 embryogenesis, development and abiotic stress responses (Supplementary Data Table S3). The  
19 global non-metric multidimensional scaling (GNMDS) analysis revealed that the TFBS  
20 composition of *PsSPDS* resembles more the TFBS composition of the *SPMS* genes of black  
21 cottonwood and Arabidopsis than the *SPDS* genes of those species (Fig. 4C). The results suggest  
22 that *PsSPDS* carries on many functions, which have been delegated to *SPMS* after the evolution  
23 of separate *SPDS* and *SPMS* genes in angiosperms.

24

## 1 *Phylogenetic relationship of plant SPDS, SPMS and ACL5 genes*

2       The phylogenetic analysis of plant *SPDS*, *SPMS* and *ACL5* sequences confirmed the close  
3 relationship between the *SPDS* and *SPMS* sequences, which is in accordance with the view that in  
4 angiosperms *SPMS* genes have evolved from *SPDS* genes via gene duplication, whereas the  
5 *ACL5* gene seems to have a different evolutionary origin. *SPDS* and *SPMS* sequences divided  
6 into separated phylogenetic groups with high bootstrap support (100 %). The *PsSPDS* sequence  
7 formed the *SPDS* group together with the *SPDS* sequences of other conifers, angiosperms and the  
8 moss *Physcomitrella*. Furthermore, the phylogenetic analysis confirmed that *PsACL5* belonged to  
9 the same main branch with the *ACL5* sequences from other seed plants and *Physcomitrella* (Fig.  
10 5).

11

## 12 **DISCUSSION**

13 In the present study, we revealed that Scots pine possesses a bifunctional *SPDS* enzyme, which is  
14 able to produce both Spd and Spm, and that Scots pine tissues contain Spd, Spm and tSpm. To  
15 our knowledge, Spm and tSpm have been previously detected in gymnosperms only in ginkgo  
16 (*Ginkgo biloba*) nuts (Takano *et al.* 2012) and, so far, the *SPMS* gene has not been found in  
17 gymnosperms (Minguet *et al.*, 2008). Thus, differing from the current view, our results  
18 demonstrate that *SPMS* activity is not a novel characteristic evolved only in the angiosperm line  
19 of the seed plants but *SPMS* activity also exists in the evolutionary old gymnosperms. The Scots  
20 pine *PsSPDS* gene and the *SPDS* genes of the angiosperm species showed highly similar gene  
21 structure, high sequence similarity at amino acid level as well as close phylogenetic relationship  
22 providing evidence for a common evolutionary history of the genes.

23       Only one *SPDS* gene was found from the loblolly pine genome, whereas angiosperm plant  
24 species tend to have several copies of *SPDS* genes. The duplication of *SPDS* genes has been

1 suggested to be the origin of a variety of new activities such as SPMS. During evolution, *SPMS*  
2 genes have arisen from *SPDS* independently at least three times, in animals, fungi and  
3 angiosperm plants (Minguet *et al.*, 2008). Currently, the existence of a separate SPMS gene that  
4 also may contribute to Spm synthesis in gymnosperms in addition to the bifunctional SPDS  
5 revealed in the present study cannot be definitely excluded due to the imperfect annotation of  
6 conifer genomes. Generally, gene duplication has been considered as a major source of new  
7 genes and functional innovations, which can originate in different ways, including mutations that  
8 directly impart new functions (neofunctionalization), subdivision of ancestral functions  
9 (subfunctionalization) and selection for changes in gene dosage (Ohno,1970; reviewed e.g. by  
10 Conant and Wolfe, 2008; Hahn, 2009). However, several studies have suggested that most new  
11 genes have no novel functions (Prince and Pickett, 2002) and, actually, the predominant fate of  
12 most duplicates is loss (Li, 1983; Maere *et al.*, 2005; Hanada *et al.*, 2008).

13 The mechanisms such as subfunctionalization that do not require the evolution of new  
14 functions may play an important role in the initial retention of duplicate genes (Panchy *et al.*,  
15 2016). The patchwork model of enzyme pathway evolution suggests that ancestral enzymes were  
16 unspecific and therefore capable to catalyze chemically similar reactions (Fani and Fondi, 2009).  
17 After duplication, two genes may be preserved if both members of the pair experience  
18 degenerative mutations that result in complementary loss of subfunctions and thus in the  
19 portioning of the functions of an ancestral gene among daughter copies (Force *et al.*, 1999; Lynch  
20 and Force, 2000). Against this background, it seems likely that the broad substrate specificity of  
21 the ancient *SPDS* genes may have favored the repeated evolution of *SPMS* genes from *SPDS*  
22 genes via gene duplications detected in several eukaryotic taxonomic groups by Minguet *et al.*  
23 (2008).

1           There is only little experimental evidence for the various theoretical models of evolution  
2 after gene duplication mostly due to incomplete information on functional properties of the  
3 progenitor gene, because these ancient genes and the proteins they encode usually no longer  
4 exist. Therefore, the bifunctional *PsSPDS* gene found in the present study from an evolutionary  
5 old gymnosperm species together with the separate *SPDS* and *SPMS* genes in angiosperms  
6 provide a valuable example of the preservation and evolution of duplicated genes. Our findings  
7 suggest that SPMS activity already existed as a secondary property in the *SPDS* progenitor gene.  
8 After the *SPDS* duplication in the angiosperm lineage, the functions of the progenitor gene  
9 became divided to the daughter copies and SPMS activity co-opted a primary role. In  
10 gymnosperm lineage, the *SPDS* gene remained bifunctional during evolution (Fig. 6).

11           Subfunctionalization may have occurred simply as a result of accumulation of neutral  
12 degenerative mutations that have removed either SPDS or SPMS activity from each gene copy  
13 after the duplication of the bifunctional progenitor gene in angiosperms. However, both SPDS  
14 and SPMS activities may not have been able to be optimized simultaneously in the bifunctional  
15 enzyme due to the different roles of Spd and Spm in plant cells. Thus, separate SPDS and SPMS  
16 enzymes might have been a solution to the conflict and provided a clear adaptive advantage. In  
17 eukaryotic organisms, hypusine synthesis defines an absolute requirement for Spd  
18 (Chattopadhyay *et al.*, 2003 and 2008; reviewed by Wolff and Park, 2015). *Arabidopsis spds1-1*  
19 *spds2-1* double-mutant lacking Spd shows embryo lethal phenotype (Imai *et al.*, 2004b). Our  
20 findings in the present study supported the essential role Spd in basic cell functions since  
21 *PsSPDS* expression was detected in all cells of the Scots pine embryo. Moreover, *PsSPDS*  
22 expression was localized in dividing cells, where also *ADC*, the enzyme catalyzing the preceding  
23 step in the PA biosynthesis pathway, has been shown to be expressed (Vuosku *et al.*, 2006).  
24 Unlike Spd, Spm is not essential for life but has generally been considered as a stress protective

1 molecule in plants (Yamaguchi *et al.*, 2007; Do *et al.*, 2013). Arabidopsis plants with blocked  
2 activity of *spms* and without Spm show similar phenotype as the wild type under optimal growth  
3 conditions, but the mutants are very sensitive to stresses (Imai *et al.*, 2004a). In pine tissues, Spm  
4 also accumulates under abiotic stresses (Islam *et al.*, 2003; Muilu-Mäkelä *et al.*, 2015). Taken  
5 together, our results propose that the evolution of separate *SPDS* and *SPMS* genes in angiosperms  
6 has provided an adaptive advantage by making not only the Spm production more efficient but  
7 also the regulation of the *SPDS* and *SPMS* activities more flexible, whereas the loss of *SPMS*  
8 activity after gene duplication would have led to fitness cost. The TFBS profile of the *PsSPDS*  
9 promoter resembles more the promoter TFBS profiles in *SPMS* genes than *SPDS* genes in  
10 angiosperm. The result suggests that bifunctional *PsSPDS* due to the production of Spm as a  
11 secondary function also possesses a wide variety of other functions, which have been acquired by  
12 *SPMS* genes in angiosperms. Furthermore, the presence of separate *SPDS* and *SPMS* genes may  
13 have released the regulation of the *SPDS* genes to evolve more freely compared to bifunctional  
14 *PsSPDS*.

15 Both *PsSPDS* and *PsACL5* retained catalytic activity also at temperatures, which are  
16 remarkable high for plant enzymes. Furthermore, *PsSPDS* seemed to have increasing affinity for  
17 Spd as a substrate at high temperatures. Raising the incubation temperature for *PsSPDS* from 30  
18 °C to 50 °C tripled Spm formation, whereas there was no significant change in the amount of  
19 produced Spd. The result suggest the importance of Spd and especially Spm production in  
20 elevated temperatures and heat stress in Scots pine, which is in agreement with the previous  
21 reports on angiosperm species (Kasukabe *et al.*, 2006; Cheng *et al.*, 2009; Sagor *et al.*, 2013;  
22 Sang *et al.*, 2017). During the seasonal fluctuation of free PAs in Scots pine needles, low  
23 Spm/tSpm concentrations coincided with low temperatures in winter (Sarjala and Savonen, 1994;

1 Sarjala and Kaunisto, 1996), which also support the connection between environmental  
2 temperatures and Spm/tSpm production.

3 The *ACL5* in eukaryotes seems to be an ancient plant-specific gene, which has been  
4 proposed to originate from a horizontal gene transfer from Archaea or Bacteria to plants  
5 (Minguet *et al.*, 2008). In *Arabidopsis*, tSpm has been identified as a novel plant growth regulator  
6 that represses xylem differentiation and promotes stem elongation by preventing premature death  
7 of developing xylem vessel elements (Kakehi *et al.*, 2008; Muniz *et al.*, 2008; Vera-Sirera *et al.*,  
8 2010). Our findings in the present study enhance the role of *ACL5* and tSpm in development of  
9 vascular structures in conifers. *PsACL5* and also *SPDS* transcripts were localized in the  
10 procambial cells in Scots pine germinating embryos and, moreover, in the developing tracheary  
11 elements of hypocotyls in young Scots pine seedlings. All vascular tissues are derived from  
12 undifferentiated, meristematic cells, and the body plan for the vasculature in the adult plant is  
13 already established in the embryo (Ye, 2002). Xylem is a specialized vascular tissue that serves  
14 as a conduit of water and nutrients and provides mechanical strength for upright growth (Růžička  
15 *et al.*, 2015). The important part of mature xylem is composed of tracheary elements, which fall  
16 into two broad categories: tracheids, typically found in lycophytes, ferns, and gymnosperms and  
17 vessel elements, which reach their peak of diversity in the angiosperms (Wilson, 2013).  
18 Altogether, the role of *ACL5* in the development of TEs in angiosperm and gymnosperm species,  
19 but also the presence of three *ACL5* copies in non-vascular moss *Physcomitrella* (Rodríguez-  
20 Kessler *et al.*, 2010), suggest that *ACL5* and further tSpm perform multiple tasks in the  
21 development of land plants.

22 In conclusion, our results revealed that an evolutionary old gymnosperm, Scots pine, has a  
23 bifunctional *SPDS* able to produce both spd and spm, when angiosperms seem to lean on separate  
24 enzymes in spd and spm biosynthesis. The *PsACL5* gene was found to code a protein with *ACL5*

1 activity catalyzing the production of tSpm, and the expression of *PsACL5* was associated with the  
2 development of vascular structures during early development of Scots pine. Altogether, the  
3 bifunctional SPDS and vascular-associated ACL5 enzymes of Scots pine shed new light on the  
4 evolution and function of aminopropyltransferases in seed plants.

5

## 6 **SUPPLEMENTARY DATA**

7 Supplementary data are available in online and consist of the following. Fig. S1: Polyamine  
8 biosynthesis pathway in plants. Fig S2: The alignment of the *SPDS*, *SPMS* and *ACL5* sequences  
9 for phylogenetic analysis. Fig. S3: Comparison of the predicted amino acid sequence of the *P.*  
10 *sylvestris* PsSPDS protein with previously characterized homologues from angiosperm. Fig. S4:  
11 Comparison of the predicted amino acid sequence of the *P. sylvestris* PsACL5 protein with  
12 previously characterized homologues from angiosperm. Fig. S5: Area of the tSpm HPLC peaks  
13 in 100 nM reaction mixture with PsACL5 enzyme and Spd as a substrate after 1h incubation  
14 under three different temperatures (30, 40 and 50°C). Table S1: Primers for PCR and sequencing  
15 of the *PsSPDS* and *PsACL5* genes from Scots pine. Table S2: Presence-absence data generated  
16 from the transcription factor binding sites (TFBSs) found in the 3kb promoter regions of *SPDS*  
17 and *SPMS* genes of *Populus trichocarpa* (*PtSPDS* and *PtSPMS*), *Arabidopsis thaliana* (*AtSPDS*  
18 and *AtSPMS*) and *Pinus sylvestris* (*PsSPDS*) using PlantPAN 2.0. database. Table S3:  
19 Transcription factor binding sites (TFBSs) and their functions found in the 3kb promoter regions  
20 of *SPDS* and *SPMS* genes of *Populus trichocarpa* (*PtSPDS* and *PtSPMS*), *Arabidopsis thaliana*  
21 (*AtSPDS* and *AtSPMS*) and *Pinus sylvestris* (*PsSPDS*) using PlantPAN 2.0. database.

22

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8

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## 15 **FIGURE LEGENDS**

16 **FIG 1.** Size and enzymatic functions of Scots pine spermidine synthase (PsSPDS) and  
17 thermospermine synthase (PsACL5) proteins. (A) SDS-PAGE analysis of recombinant PsSPDS  
18 and PsACL5 proteins expressed in *E. coli*; purified PsSPDS and PsACL5 proteins with protein  
19 molecular mass marker. (B) Identification of the reaction products after conversion of putrescine  
20 (Put) and spermidine (Spd) by PsSPDS and Spd by PsACL5. The HPLC chromatograms show  
21 dansylated polyamine standard of Put, Spd and spermine (Spm) (upper chart); reactions without  
22 enzyme as a control (middle chart); reactions with PsSPDS enzyme after 1h incubation under  
23 50°C degrees with Put and Spd as a substrate, respectively, and a reaction with PsACL5 enzyme  
24 after 1h incubation under 50°C degrees with Spd (lower chart). (C) Spd and Spm formation by

1 PsSPDS at 30°C, 40°C and 50°C incubation temperatures with Put and Spd as a substrate,  
2 respectively. Columns labeled with different letters indicate statistically significant differences  
3 (n=3, p<0.05, pairwise t-test, R Software Package 3.3.2).

4  
5 **FIG 2.** Localization of spermidine synthase (*PsSPDS*) and thermospermine synthase (*PsACL5*)  
6 expression in germinating Scots pine seeds. (A) The localization of *PsSPDS* transcripts in the  
7 embryo by mRNA *in situ* hybridization assay using DIG-labelled RNA-probes with NBT/BCIP  
8 detection (blue signal). (B) Intense *PsSPDS* expression in the procambial cells of the embryo. (C,  
9 D and E) *PsSPDS* expression in the dividing procambial cells. (F and G) *PsACL5* expression in  
10 the procambial cells of the embryo. (H) A seed section hybridized with the sense *PsSPDS* probe  
11 as a negative control. (I) A section hybridized with the sense *PsACL5* probe as a negative control.  
12 E=embryo, M=megagametophyte.

13  
14 **FIG 3.** Localization of spermidine synthase (*PsSPDS*) and thermoseprmine synthase (*PsACL5*)  
15 expression in hypocotyls of Scots pine seedlings. (A and B) The localization of *PsSPDS*  
16 transcripts in developing tracheary elements by mRNA *in situ* hybridization assay using DIG-  
17 labelled RNA-probes with rhodamine detection (red signal). (C) A hypocotyl section hybridized  
18 with the sense *PsSPDS* probe as a negative control. (D and E) *PsACL5* expression in developing  
19 tracheary elements. (F) A hypocotyl section hybridized with the sense *PsACL5* probe as a  
20 negative control.

21  
22 **FIG 4.** Structure of Scots pine spermidine synthase (*PsSPDS*) and thermospermine synthase  
23 (*PsACL5*) genes and functional analysis of *PsSPDS* promoter region. (A and B) The exon-intron  
24 structures of *PsSPDS* and *PsACL5* were compared with previously characterized plant



1 homologues. Introns are shown as solid lines while boxes representing exons are numbered from  
2 I to X. (C) Global non-metric multidimensional scaling (GNMDS) analysis of the *SPDS* and  
3 spermine synthase (*SPMS*) promoter regions. The GNMDS is based on the presence-absence data  
4 of the putative transcription factor binding sites (TFBSs) in the promoter regions of the *SPDS* and  
5 *SPMS* genes of *Pinus sylvestris*, *Arabidopsis thaliana* and *Populus trichocarpa*.

6

7 **FIG 5.** Phylogenetic analysis of spermidine synthase (*SPDS*), spermine synthase (*SPMS*) and  
8 thermospermine synthase (*ACL5*) genes in plants.

9

10 **FIG 6.** Proposed model for evolution of aminopropyltransferases in seed plants. In ancient seed  
11 plants, the spermidine synthase (*SPDS*) enzyme possessed a broad substrate specificity and  
12 showed spermine synthase (*SPMS*) activity as a secondary property. After the duplication of the  
13 *SPDS* gene in the angiosperm lineage, the functions of the progenitor enzyme were divided to the  
14 daughter copies and *SPMS* activity co-opted a primary role in one copy, whereas the bifunctional  
15 form was preserved in gymnosperm lineage. The thermospermine synthase (*ACL5*) gene seems to  
16 have a different evolutionary origin and may have been acquired by plants via horizontal gene  
17 transfer.

**TABLE 1.** Polyamine content ( $\text{nmol g FW}^{-1}$ ) in Scots pine seedlings and seeds

Scots pine tissue	Put	Spd	Spm*	tSpm*	tSpm/Spm (%)
<b>Roots</b>	22.7	11.8	6.0	0.4	6.7
<b>Hypocotyls</b>	31.4	21.5	8.3	0.3	3.5
<b>Cotyledons</b>	125.3	68.3	22.5	0.5	2.0
<b>Embryos</b>	47.3	134.4	24.4	7.7	31.5
<b>Megagametophytes</b>	23.3	53.7	10.9	2.7	24.7

Put, putrescine; Spd, spermidine; Spm, spermine; tSpm, thermospermine.

\*Spm and tSPM concentrations in different Scots pine tissues were measured as benzylated polyamines according to Naka *et al.* (2010).