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AINTEGUMENTA homologue expression in *Gnetum* (gymnosperms) and implications for the evolution of ovulate axes in seed plants

Running title -- ANT homologue expression in Gnetum

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AINTEGUMENTA (ANT) found in the gymnosperm Gnetum parvifolium, was analyzed by RT-PCR and in situ hybridization. GpANTL1 was expressed in the leaf primordia, root tips, and young ovules. In the ovulate axis, expression was detected as four distinct rings around the outer, middle, and inner envelope primordia as well as around the nucellar tip. This pattern of expression is similar to that of ANT in Arabidopsis thaliana. A comparison of the expression of GpANTL1 with that of PtANTL1 in the conifer Pinus thunbergii suggests that the integrated expression of PtANTL1 may have been caused by congenital fusion of the integument, ovuliferous scale, and bract.

INTRODUCTION

Since the origin of the ovule in the Devonian period (Stewart and Rothwell 1993), the morphology of the distal-proximal ovulate axis has diversified in seed plants (Florin 1951; Crane 1985; Doyle and Donoghue 1986). The nearly sessile gymnosperm ovule is composed of a distal nucellus, which harbors the female gametophyte with several archegonia and a proximal integument. The integument surrounds the nucellus completely except for a small opening known as the micropyle. The micropyle is at the distal end of the ovule, opposite the proximal connection to the sporophyll, resulting in an orthotropous configuration (Coulter and Chamberlain 1917; Bouman 1984; Gifford and Foster 1988). In some extant gymnosperm lineages, such as the Gnetales and Coniferales, the ovule is enveloped by a single integument (i.e., a unitegmic ovule), and this structure is incorporated with the shoot and lateral organs (i.e., the sporophylls and bract) into a complicated ovulate axis. In contrast, in the Cycadales, the ovules are simply borne on the

megasporophyll, while in the Ginkgoales, pairs of ovules terminate on a stalk (Coulter and Chamberlain 1917; Florin 1951; Crane 1985; Doyle and Donoghue 1986; Gifford and Foster 1988; Shindo et al. 1999).

The angiosperm ovulate axis is characterized by an acropetal sequence that includes the funiculus, which connects the ovule to the placenta, outer and inner integuments that flank the chalaza, and the nucellus around the embryo sac (Bouman 1984; Robinson-Beers et al. 1992). The micropyle, which is formed either by both integuments or by only the inner integument, sits adjacent to the funiculus in anatropous ovules, or opposite in orthotropous ovules. Developmental similarities suggest homology between the inner integument of the angiosperms and the lone integument of the gymnosperms (Bouman 1984; Umeda et al. 1994), while the outer integument is inferred to be homologous to a leaflike organ (Umeda et al. 1994; Imaichi et al. 1995; Meister et al. 2003; Yamada et al. 2001a, b, 2003). Thus, in angiosperms, the unitegmic ovule and a leaflike organ should be integrated into the ovulate axis.

To determine what evolutionary changes gave rise to the

complicated ovulate axis of seed plants, it is necessary to evaluate the genetic basis of ovulate axis formation among extant seed plants. AINTEGUMENTA (ANT), a member of the AP2 subfamily within the AP2/ERBP family, is involved in ovule development in *Arabidopsis* thaliana and required for proper integument growth (Elliott et al. 1996; Klucher et al. 1996; Mizukami and Fischer 2000; Shigyo and Ito 2004; Kim et al. 2006; Shigyo et al. 2006). ANT is transcribed throughout the ovule primordia during Stage 8, when the primordia emerge from the placenta (Elliot et al. 1996; Klucher et al. 1996; Schneitz 1998; Balasubramanian and Schneitz 2000, 2002; Seiber et al. 2004). During Stages 9 and 10, ANT expression is negatively regulated in the nucellus and funiculus by NOZZLE/SPOROCYTRLESS (NZZ/SPL), resulting in restricted expression in the chalaza (i.e., where the inner and outer integuments will be formed) (Elliot et al. 1996; Klucher et al. 1996; Schneitz 1998; Balasubramanian and Schneitz 2000, 2002; Seiber et al. 2004). Previously, a gymnospermous ANT homolog was isolated from Pinus thunbergii Parl. (Pinaceae, Coniferales) and its expression was analyzed by in situ hybridization and RT-PCR (Shigyo and Ito 2004).

However, due to its highly modified nature (e.g., Florin 1951; Gifford and Foster 1988), the ovulate axis in the Coniferales is difficult to compare with the ovulate axis of the angiosperms. Gnetales is the putative sister of the Pinaceae or Coniferales among the extant gymnosperms (Winter et al. 1999; Soltis et al. 2002; Burleigh and Mathews 2004), but its members have a less complex ovulate axis than the Coniferales (Coulter and Chamberlain 1917; Florin 1951; Crane 1985; Gifford and Foster 1988; Shindo et al. 1999). Thus, the expression of an *ANT* homolog in the Gnetales would not only help explain the genetic basis underlying the differences between the ovules of gymnosperms and angiosperms, but it would also help elucidate the evolutionary process that led to the modified ovulate structure of the Coniferales.

As well as the role that *ANT* played in the ovule evolution, its significant developmental role roused much interests to the evolution of *ANT* homologs (Shigyo and Ito, 2004; Kim et al. 2006; Shigyo et al. 2006; Floyd and Bowman, 2007). *ANT* is believed to regulate cellular proliferation via maintenance of the meristematic competence of other lateral organs (i.e., on the vegetative and reproductive axes), such as

the leaves, floral organs and integuments in *A. thaliana* (Mizukami and Fischer 2000). However, a little data is available on the function and expression of *ANT* homologs in other species than *A. thaliana*, hindering the inference of their evolution.

In this study, we analyzed the expression of *GpANTL1*, an *ANT* homolog in *Gnetum parvifolium* (Warb.) C. Y. Cheng (Shigyo et al. 2006), by RT-PCR and *in situ* hybridization. We then compared our data with that of *ANT* in *A. thaliana* (Elliot et al. 1996; Klucher et al. 1996; Schneitz 1998; Balasubramanian and Schneitz 2000, 2002) and inferred the ancestral function of *ANT* homologs. Furthermore, we compared the expression of *GpANTL1* in ovulate axis to that of *ANT* in ovule of *A. thaliana* as well as that of *PtANTL1* in the reproductive shoots of *P. thunbergii* (Shigyo and Ito 2004). The implications of our data for the evolution of the ovulate axis in seed plants is also discussed.

MATERIALS AND METHODS

Plant materials

Female strobili of *G. parvifolium* were collected from a plant cultivated in the Botanical Gardens, Graduate School of Science, University of Tokyo, from May to June 2006, and the ovules were dissected from the strobili on ice. Leaf primordia, root tips, and vegetative shoots were collected from a single plant of *G. parvifolium* in August and December 2007. This plant was also used for the isolation of *GpAP2L1* and *GpANTL1* by Shigyo et al. (2006).

All tissues were frozen in liquid nitrogen for use in the isolation of *GpANTL1* and RT-PCR analysis. For *in situ* hybridization, female strobili and vegetative shoots were fixed in 4% formaldehyde (w/v) in which the pH had been adjusted to 7.0 with NaP_i buffer for 15 h. In female strobili, the collenchymatous tissue of the annular collar was removed to facilitate fixation.

Isolation of GpANTL1

Total nucleic acids were extracted from the ovules with cethy trimethyl ammonium bromide (CTAB) (Doyle and Doyle 1987) and then applied to an RNeasy mini column (QIAGEN, Hilden, Germany) from an RNeasy Mini Kit (QIAGEN) to obtain total RNA. Single-stranded cDNA was subsequently synthesized using the 3' RACE System for Rapid Amplification of cDNA Ends (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's protocol. Two gene-specific primers, GpANTF1 (5'-CACAAGGCACAGATGGACAG-3') and GpANTF2 (5'-GGCTCATTTGTGGGACAACA-3'), were designed based on partial sequences of GpANTL1 (AB195247) registered in the DDBJ/EMBL/NCBI database, and nested PCR was conducted using the amplified product. GpANTF1 and Universal Amplification Primer (UAP) (Invitrogen) were used in the first round of PCR, while GpANTF2 and UAP were used for the nested PCR. Three internal primers for 5' RACE were subsequently designed based on the sequences obtained from the 3' RACE amplification: GpANT5R1 (5'- GCTTCATCTCATCAATCTGC-3'), GpANTR2 (5'- GCCTTTTCCTCCTGATCGTA-3'), and GpANT5R3 (5'-GCTCTGCCCTTCTTTTTGC-3'). Amplification of the 5' partial cDNA

fragments was performed using the 5' RACE System for Rapid

Amplification of cDNA Ends (Invitrogen). The sequence of *GpANTL1*(2005 bp) is registered as AB297493 in the DDBJ.

Alignment and phylogenetic analysis

Initially, the nucleotide sequences of *ANT* and several (outgroup) *AP2* genes were retrieved from the DDBJ/EMBL/NCBI DNA database (S1). Their amino acid sequences were then deduced and aligned using CLUSTAL X, v. 1.83 (Thompson et al. 1997). The residues within the two AP2 domains and the linker region between them were used for the phylogenetic analyses.

Bayesian phylogenetic analysis was performed using BEAST v. 1.4 (Drummond and Rambaut 2006). The JTT + G model of amino acid substitution, which assumes a gamma distribution to accommodate for the rate of variation among sites, was adopted for the analysis because the Akaike Information Criterion (AIC), the correlated AIC, and the likelihoods calculated by ProtTest v. 1.2.6 (Abascal et al. 2005) all

predicted it as the best model of molecular evolution. The starting tree for the Markov chain Monte Carlo (MCMC) analysis was generated using a coalescent process. The trees were sampled every 1,000 steps for 10,000,000 steps after discarding 1,000,000 burn-in steps.

Convergence of the chain was verified using Tracer v. 1.3 (Rambaut and Drummond 2005).

Maximum parsimony (MP) and bootstrap analyses were implemented by PAUP*4.0b10 (Swofford 2001) using a heuristic search strategy with 100 random addition sequence replicates and the TBR branch-swapping algorithm. All optimal trees were saved in our MP analyses, while ten trees per replicate of 1,000 total replicates were saved in our bootstrap analyses.

Neighbor Joining (NJ) analysis was conducted using PHYLIP v.

3.56 (Felsenstain 1994). The distance matrix was generated using the

JTT matrix model of PROTDIST (Felsenstain 1994). A NJ tree was

generated using NEIGHBOR (Felsenstain 1994). Support for the clade

was evaluated by bootstrap analysis with 1,000 replicates.

RT-PCR

Total RNA was extracted from root tips, vegetative shoots, leaf primordia, young ovules, and developed ovules (i.e., those with three envelopes). RNA extraction and cDNA synthesis were performed as described for the isolation of *GpANTL1*.

Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was

(AB362487 in the DDBJ) was amplified for normalization of the mRNA in the samples by PCR using the primers GpGAPF1

(5'-AAAGGTGTTACCTGCTCTTA-3') and GpGAPR1

(5'-CGTTCAGGGCTATGCCGGCC-3') (260-bp product). Partial

GpANTL1 cDNA was amplified using GpANTPF1 (5'
ACTGCCATCTGGCCACGTCG-3') and GpANTPR1 (5'
TGGGTGGCGTGGCTCAGGTG-3'). The amplified fragment included

410 bp just downstream of the AP2-repeat 2-domain coding region. A

total of 30 cycles was used for the amplification of GAPDH and

GpANTL1. The products were electrophoresed in a 1% agarose gel and stained with ethidium bromide.

In situ hybridization

The partial cDNA of *GpANTL1* was amplified by PCR using GpANTPF1 and GpANTPR1. The product was then cloned into pCR II (Invitrogen) and amplified using M13⁺ (Invitrogen) and GpANTPR1 or M13⁻ (Invitrogen) and GpANTPF1 for probe synthesis. Contaminating RNases were removed with RNAsecure Reagent (Ambion, Austin, TX, USA), and DIG-labeled antisense and sense riboprobes were synthesized as described previously (Yamada et al. 2003). *In situ* detection of *GpANTL1* transcription was performed as per Yamada et al. (2004).

RESULTS

ANT gene phylogeny

By Bayesian phylogenetic analysis, the monophyly of the ANT genes

was supported with a posterior probability of 100%. The ANT group consisted of two clades, euANT and basalANT (sensu Kim et al. 2006), each of which had a posterior probability of 100% (Fig. 1). Within the euANT clade, two major clades, BBM/PLT and ANT (sensu stricto, s.s.), were recognized with 99 and 100% posterior probabilities, respectively, but the relationship between the two clades and *PpANT1* was not fully resolved (Fig. 1). All gymnospermous ANT genes were nested in the ANT (s.s.) clade, and formed a clade with a moderate posterior probability value of 82% (gymnoANT clade in Fig. 1). However, all of the angiosperm genes except AIL1 were included in a single clade with a posterior probability of 99% (angioANT clade in Fig. 1). The relationships among AIL1, the angioANT clade, and the gymnoANT clade were not unambiguously supported (Fig. 1).

By both MP and NJ analyses, the ANT, basal ANT, euANT, and BBM /PLT clades were recovered with more than 70% bootstrap support. NJ analysis indicated monophyly for ANT (s.s.) including the gymnoANT clade with 64% bootstrap support, but the ANT (s.s.) clade was not recovered in a strict consensus tree by MP analysis.

GpANTL1 expression

GpANTL1 expression was detected in the vegetative shoots including leaf primordia, leaf primordia, root tips, and young ovules. *GpANTL1* expression was extremely weak in the developed ovules compared to that in the young ovules (Fig. 2). In vegetative shoots, GpANTL1 was expressed in leaf primordia. The stronger transcription is observed in the abaxial and adaxial tissues of the leaf primordia, where the laminar growth was taken place by active cell divisions (Fig. 3A, B).

Ovule formation was observed in the axil of the annular collar. Of the three envelopes, the outer envelope developed first and covered the ovulate axis long before the middle and inner envelope primordia emerged. At this stage, *GpANTL1* expression was detected in the abaxial epidermis and subdermis of the outer envelope as well as in the inner and middle envelopes (Fig. 3C, D). It was also detected in the nucellar tip (Fig. 3C, D). No signal was detected in the internode (i.e., between the inner and middle envelope anlagen) or in the tissues

between the middle and outer envelopes. Together, the regions where *GpANTL1* was expressed formed four separate rings around the ovulate axis corresponding to the future nucellus and the outer, middle, and inner envelopes (Figs. 3C, D). A little later, the middle envelope emerged slightly, while the inner envelope primordium remained hidden (Fig. 3E, F). As in the previous stage, *GpANTL1* expression was detected in four distinct domains (the future nucellus and three envelopes) (Figs. 3E, F). As the ovule developed further, the inner and middle envelope primordia emerged (Fig. 3G, H). *GpANTL1* expression was greatly diminished at this point in the outer and middle envelopes, while some expression remained in the nucellar tip and inner envelope (Fig. 3G, H).

DISCUSSION

Phylogeny of the ANT genes

The phylogenetic framework of the *ANT* genes included in this study is similar to those of previous studies in two respects: the group is subdivided into the basalANT and euANT clades, and the ANT (s.s.) and BBM/PLT clades are recovered within the euANT clade (Shigyo and Ito 2004; Kim et al. 2006; Shigyo et al. 2006; Floyd and Bowman 2007). As described earlier (Shigyo and Ito 2004; Shigyo et al. 2006; Floyd and Bowman 2007), all of the gymnosperm *ANT* genes reported so far are nested within the ANT (s.s.) clade. This implies that gene loss occurred at least once in the BBM/PLT and basalANT clades, or that genes related to these two lineages have not yet been isolated in gymnosperms.

Putative ancestral function of the ANT (s.s.) clade genes

In *Arabidopsis*, *ANT* is expressed in the developing lateral organs (i.e., on the vegetative and reproductive axes), such as the leaves, floral organs, and the inner and outer integuments (Elliott et al. 1996; Klucher et al. 1996; Krizek 1999; Mizukami and Fischer 2000). In the *ant* mutant,

organ size is reduced due to a decrease in cell number, while constitutive ANT expression causes organ hyperplasia due to an increase in cell number (Mizukami and Fischer 2000). Thus, ANT is believed to regulate cellular proliferation via maintenance of the meristematic competence of the lateral organs (Mizukami and Fischer 2000). Consistent with this idea, ANT expression in the lateral organs decreases as organ development progresses (Elliott et al. 1996; Klucher et al. 1996; Krizek 1999; Mizukami and Fischer 2000; Nole-Wilson et al. 2005). In addition to its expression in the lateral organs, ANT is also expressed in the roots in Arabidopsis, although its function there is unknown (Nole-Wilson et al. 2005). In gymnosperms, similar expression of the ANT homolog PtANTL1 was reported in the lateral organs of the reproductive shoot in P. thunbergii (Pinaceae) (Fig. 4B; Shigyo and Ito 2004). However, no expression data are available for the leaf primordia and roots, and thus the ancestral expression pattern and function of the ANT (s.s.) genes are not well understood.

Developmental studies suggest that the inner envelope of *Gnetum* is homologous to the integument (e.g., Takaso and Bouman 1986), while

the outer and middle envelopes are comparable to the leaves that enclose unitegmic ovule(s) (Crane 1985; Shindo et al. 1999; Hasebe 1999). *GpANTL1* is expressed in the outer, middle, and inner envelopes as well as in the leaf primordia and roots. This pattern is consistent with that of *ANT* in *A. thaliana* (Elliott et al. 1996; Klucher et al. 1996; Krizek 1999; Mizukami and Fischer 2000; Nole-Wilson et al. 2005), suggesting that this is the ancestral pattern of the ANT (s.s.) clade. Together, the enhanced expression of *GpANTL1* in the young envelope primordia versus that in the old primordia was observed in RT-PCR analysis and its transcription is decreased in the differentiated tissues in the leaf primordia. These data imply that regulation of cellular proliferation in the lateral organs is the ancestral function of the *ANT* (s.s.) genes.

A major difference between *GpANTL1* and *ANT* is that *GpANTL1* is expressed in the nucellar tip while *ANT* expression is absent from the nucellus (Fig. 4A, C). Expression in the nucellar tip was previously reported for *PtANTL1* in *P. thunbergii* (Fig. 4B; Shigyo and Ito 2004), implying that it represents an ancestral pattern of gene expression in the gymnospermous ovulate axis. Such divergent gene expression in

gymnosperms and angiosperms may be attributable to differential regulation of nucellar development or a remarkable change in the angiosperm nucellus.

Evolution of the ovulate axis in seed plants

The ANT (s.s.) clade genes are transcribed differently in *G. parvifolium* (Gnetales), *P. thunbergii* (Coniferales), and *A. thaliana* (angiosperms), but their expression is similar with respect to the lateral organs of the ovulate axis. *GpANTL1* is expressed separately in the developing inner, middle, and outer envelope primordia (Fig. 4A). In comparison, *PtANTL1* expression in *P. thunbergii* has been detected throughout the entire primordium of the bract and ovule—ovuliferous scale complex (Fig. 4B: Shigyo and Ito 2004). In contrast, in *A. thaliana*, *ANT* is expressed in a single band of cells including the chalaza and the developing inner and outer integument (Elliott et al. 1996; Klucher et al. 1996; Mizukami and Fischer 2000; Fig. 4C). How can this difference be interpreted from an evolutionary point of view?

It is widely accepted that two pairs of leaves fuse to create radially symmetrical outer and middle envelopes in the ancestral fertile shoot axis, which bears unitegmic ovules at its apical end (Crane 1985; Shindo et al. 1999), although no direct evidence is available. According to this interpretation, the ovulate axis of *Gnetum* is a highly contracted axis with three envelopes, each of which is a distinct developmental unit (Fig. 5A). The ovule-ovuliferous scale complex of the Coniferales is likely another highly condensed, modified fertile shoot subtended by the scaly bract (Florin 1951; Gifford and Foster 1988). Gene expression analyses supported this interpretation (Shindo et al. 1999; Hasebe 1999). Axis condensation may have caused lateral congenital fusion of the ovules and ovuliferous scale on the ovulate axis and subtending bract (Fig. 5B). The integrated expression of PtANTL1 in Pinus may be attributable to fusion between lateral organs, but the distinct expression of GpANTL1 in three domains with subsequent organogenesis is retained in *Gnetum* (Fig. 5A, B).

Recent molecular phylogenetic analyses have suggested that the extant angiosperms are distantly related to the extant gymnosperms

(Winter et al. 1999; Soltis et al. 2002; Burleigh and Mathews 2004). Therefore, the leaves on the fertile shoots of extant gymnosperms may or may not be homologous in a strict sense to those of angiosperms. Nonetheless, the difference in expression between the *Gnetum* and Pinus ANT homologs may provide an important analogy regarding the evolutionary history of the angiosperm ovulate axis. Gymnosperms and angiosperms have similar fertile axes, into which the unitegmic ovules and leaves are incorporated; in gymnosperms, the middle and outer envelopes (in Gnetum) and the ovuliferous scale (in conifers) are modified leaves, while in angiosperms, the outer integument of the bitegmic ovule may also be a modified leaf (Yamada et al. 2001a,b; Doyle 2006). In light of the hypothesis that the two integuments evolved from different organs at different times, they may have distinct origins. ANT expression in the chalaza in Arabidopsis is partly similar to the expression of PtANTL1 in Pinus because no tissue without ANT expression exists between the two integuments (Elliott et al. 1996; Klucher et al. 1996; Mizukami and Fischer 2000; Fig. 4C). This suggests that the internodal tissues between the outer and inner integuments are extremely reduced (or could be congenitally fused into the chalaza) in angiosperms, like the ovuliferous scale and bract in conifers. This hypothesis is supported by the fact that the pattern of *ANT* expression in the chalaza is also observed for *BELL1*, which is involved in integument formation (Reiser et al. 1995; Klucher et al. 1996; Baker et al. 1997; Gasser at al. 1998; Balasubramanian and Schneitz 2000, 2002). The role which the internode played in the ovulate axis evolution could be tested by examining the involvement of genes regulating internode growth, such as BREVIPEDICELLUS/OSH15 homologs (Sato et al. 1999; Douglas et al. 2002; Venglat et al. 2002; Smith and Hake 2003), in seed plants.

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Figure legends

Fig. 1. Bayesian phylogenetic tree for the *ANT* genes. The numbers above and below the branches are the bootstrap values from the MP analysis (>50%), bootstrap values from the NJ analysis (>50%), and posterior probability values from the Bayesian analysis (>80%). Those branches with >90% posterior probability values are indicated by bold black lines and those with 80–90% are indicated by bold gray lines. The branches of the *ANT* and *AP2* genes are contracted. Bar = 0.1 substitution per site.

Fig. 2. Expression of *GpANTL1* in vegetative shoots (VS), leaf primordia (LP), root tips (RT), young ovules with developing inner and middle envelopes (OV1), and ovules with three developed envelopes (OV2). *GpGAPDH* was used to normalize the amount of mRNA.

Fig. 3. Expression of GpANTL1 in the shoot apex and ovulate axis of Gnetum parvifolium. (A, C, E, G) antisense probe. (B, D, F, H) sense probe (negative control). (A, B) Obliquely cross sections through shoot apex. Arrow indicates the boundary between the shoot apex and leaf primordium. Scale bars = 100 µm. (C, D) Longitudinal sections of the ovulate axis with the outer envelope completely covering the ovule and middle envelope. Expression in the inner and middle envelope is indicated by arrowheads. Scale bars = 50 µm. (E, F) Longitudinal sections of the ovulate axis at a slightly later stage than that in A and B. Expression in the inner and middle envelope is indicated by arrowheads. Scale bars = $100 \mu m$. (G, H) Longitudinal sections of the inner and middle envelopes developing on the ovulate axis. Scale bars = $50 \mu m. i$, integument (inner envelope); I, leaf primordium; me, middle envelope; n, nucellus; oe, outer envelope; s, shoot apex.

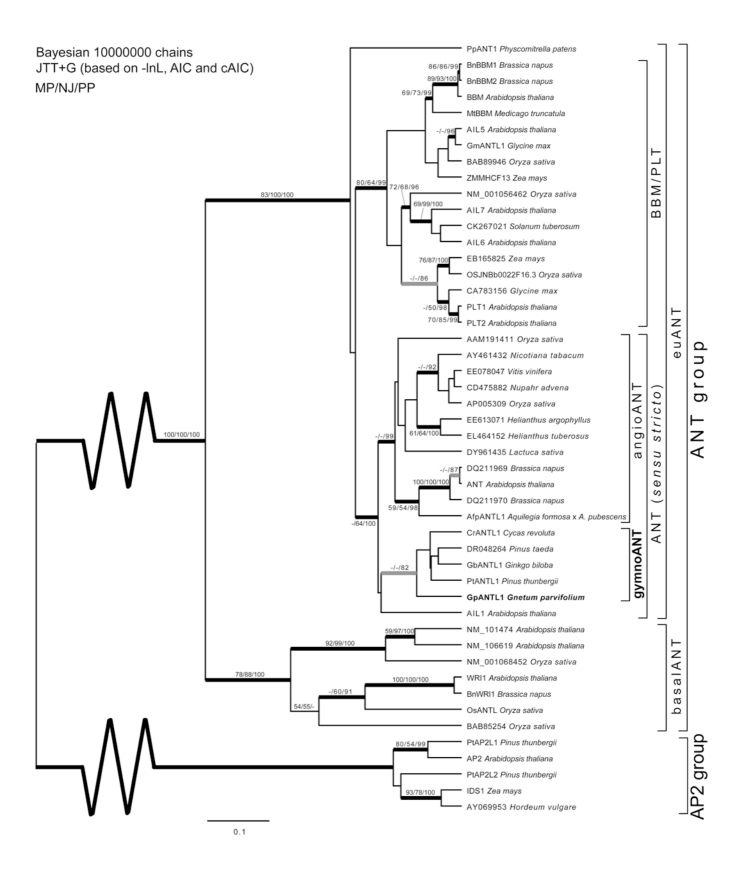
Fig. 4. Schematic drawings of ANT homolog expression. Expression is

orthogonally projected onto the median longitudinal section of the ovulate axis (shaded). (A) *GpANTL1* in *Gnetum parvifolium*. (B) *PtANTL1* in *Pinus thunbergii* (from Shigyo and Ito 2004). (C) *ANT* in *Arabidopsis thaliana* (from Elliot et al. 1996). bs, bract scale; f, funiculus; i, integument (inner envelope); ii, inner integument; me, middle envelope; n, nucellus; oe, outer envelope; oi, outer integument; os, ovuliferous scale.

Fig. 5. *ANT* homolog expression in the ovulate axes of Gnetales and Coniferales and their proposed evolution. *ANT* homolog expression in extant plants (right) and their hypothetical ancestors (left) are indicated in red. Subsumed axes are indicated by dashed lines. The phylogeny is based on Soltis et al. (2002). (A) Gnetales. (B) Coniferales. bs, bract scale; i, integument (inner envelope); I, leaf; me, middle envelope; n, nucellus; oe, outer envelope; os, ovuliferous scale.

Supplementary Material

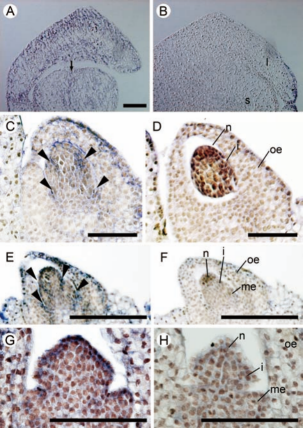
S1. The genes used in the phylogenetic analysis are shown with their accession numbers.

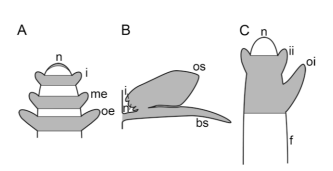


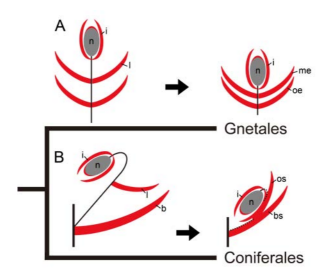
VS LP RT OV1 OV2

GpANTL1

GpGAPDH







Gene	Accession Number	Species
AAM191411	AC103891	Oryza sativa
AfpANTL1	DT750376, DT729383	Aquilegia formosa x A. pubescens
AIL1	DQ446422	Arabidopsis thaliana
AIL5	NM_125122	Arabidopsis thaliana
AIL6	NM_121089	Arabidopsis thaliana
AIL7	NM_125949	Arabidopsis thaliana
ANT	U40256	Arabidopsis thaliana
AP005309	AP005309	Oryza sativa
AP2	U12546	Arabidopsis thaliana
AY069953	AY069953	Hordeum vulgare
AY461432	AY461432	Nicotiana tabacum
BAB85254	AP003247	Oryza sativa
BAB89946	AP003313	Oryza sativa
BBM	AF317907	Arabidopsis thaliana
BnBBM1	AF317906	Brassica napus
BnBBM2	AF317905	Brassica napus
BnWRI1	DQ370141	Brassica napus
CA783156	CA783156	Glycine max
CD475882	CD475882	Nuphar advena
CK267021	CK267021	Solanum tuberosum
CrANTL1	AB195243	Cycas revoluta
DQ211969	DQ211969	Brassica napus
DQ211970	DQ211970	Brassica napus
DR048264	DR048264	Pinus taeda
DY961435	DY961435	Lactuca sativa
EB165825	EB165825	Zea mays
EE078047	EE078047	Vitis vinifera
EE613071	EE613071	Helianthus argophyllus
EL464152	EL464152	Helianthus tuberosus
GbANTL1	AB195245	Ginkgo biloba
GmANTL1	CF807326, AW200688	Glycine max
GpANTL1	AB297493	Gnetum parvifolium
IDS1	AF048900	Zea mays
MtBBM	AY899909	Medicago truncatula
NM_001056462	NM_001056462	Oryza sativa
NM_001068452	NM_001068452	Oryza sativa
NM_101474	NM_101474	Arabidopsis thaliana
NM_106619	NM_106619	Arabidopsis thaliana

Gene	Accession Number	Species
OsANTL	AB247626	Oryza sativa
OSJNBb0022F16.3	AL606446	Oryza sativa
PLT1	AY506549	Arabidopsis thaliana
PLT2	AY506550	Arabidopsis thaliana
PpANT1	BJ178045	Physcomitrella patens
PtANTL1	AB101585	Pinus thunbergii
PtAP2L1	AB101586	Pinus thunbergii
PtAP2L2	AB101587	Pinus thunbergii
WRI1	NM_115292	Arabidopsis thaliana
ZMMHCF13	Z47554	Zea mays