

9-30-2004

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Leelapon, Oranuch; Sarath, Gautam; and Staswick, Paul E., "A single amino acid substitution in soybean VSP α increases its acid phosphatase activity nearly 20-fold" (2004). *Publications from USDA-ARS / UNL Faculty*. Paper 46.
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A single amino acid substitution in soybean VSP α increases its acid phosphatase activity nearly 20-fold

Received: 10 February 2004 / Accepted: 17 April 2004 / Published online: 23 July 2004
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Abstract Soybean [*Glycine max* (L.) Merr.] contains two proteins called vegetative storage proteins (VSPs) that function as temporary storage reserves, but are also closely related to plant acid phosphatases of the haloacid dehalogenase (HAD) superfamily. This study examined the biochemical basis for the relatively low catalytic activity previously reported for these VSPs. The specific activity of purified recombinant VSP α on GMP was about 40-fold lower than for a related soybean root nodule acid phosphatase (APase), which had a specific activity of 845 U mg⁻¹ protein. Conversion of Ser¹⁰⁶ to Asp increased VSP α activity about 20-fold. This Asp residue is present in nodule APase and is a highly conserved nucleophile in the HAD superfamily. Related VSPs from cultivated soybean and from three wild perennial soybeans, as well as a pod storage protein (PSP) from *Phaseolus vulgaris* L. all lack the catalytic Asp, suggesting they too are catalytically inefficient. Phylogenetic analysis showed the VSPs and PSP are more closely related to each other than to 21 other VSP-like proteins from several plant species, all of which have the nucleophilic Asp. This study suggests that loss of catalytic activity may be a requirement for the VSPs and PSP to function as storage proteins in legumes.

Keywords Acid phosphatase · Enzyme activity · *Glycine* · Mutation · Vegetative storage protein

Abbreviations APase: Acid phosphatase · GST: Glutathione *S*-transferase · HAD: Haloacid dehalogenase · pNPP: *Para*-nitrophenol phosphate · PSP: Pod storage protein · RIP: Ribosome inactivating protein · VSP: Vegetative storage protein

Introduction

A wide variety of plant species abundantly accumulate proteins that function as storage reserves. The best studied of these are the seed storage proteins synthesized during seed development and then degraded during germination (Derbyshire et al. 1976). Also well documented are numerous vegetative storage proteins (VSPs) that accumulate in other plant tissues. These proteins are often synthesized in sink tissues, such as developing leaves, and then may be degraded within the same growing season to contribute to the needs of developing seeds or other plant sinks (Wittenbach 1983; for review, see Staswick 1994). Other VSPs accumulate in overwintering tissues, such as tree bark (O’Kennedy and Titus 1979), tubers and non-tuberous roots (for review, see Bewely 2002), and can supply amino acids to reinitiate growth in the spring.

Proteins are generally classified as storage reserves based on their abundance and pattern of accumulation and degradation. Some have no other known biological activity, but others are enzymatically active or have other biological properties that can raise questions about what their primary function is. For example, certain lectins of bark tissue have been considered storage proteins (Greenwood et al. 1986), but they may also have other roles, such as plant defense (Peumans and Van Damme 1995). The abundant potato storage reserve patatin has lipid acyl hydrolase activity (Andrews et al. 1988) and the sporamin tuber protein of

Accession numbers for the VSP α sequences reported in this paper are from *G. falcata*, AY523602; *G. tomentella*, AY523603; *G. curvata*, AY523604

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sweet potato is related to trypsin inhibitors (Yeh et al. 1997). An abundant storage protein from the Andean tuber crop oca has antimicrobial activity, which may also be important for protection against pathogens (Flores et al. 2002). A functional β -amylase also appears to serve as a storage reserve in alfalfa taproots and it is not clear that enzymatic activity has an important function (Gana et al. 1998).

There are also storage proteins that are clearly derived from active proteins based on sequence homology, but they have lost some or all of their activity. Among these is an abundant but inactive RNase-like storage protein from rhizomes of *Calystegia sepium* (hedge bindweed; Van Damme et al. 2000) and bark lectin-like proteins from *Cladastris lutea* (yellow wood) and *Sambucus nigra* (black elderberry) that lack sugar-binding capacity (Van Damme et al. 1995; Chen et al. 2002). Loss of these biological properties may have occurred because they were redundant and offered no selective advantage, or because inactivation was required for the protein to function as an abundant storage reserve.

Soybean (*Glycine max*) accumulates a limited number of VSPs to high level in developing vegetative sink tissues and these are later preferentially degraded. The major soybean VSPs are VSP α and VSP β , two glycoproteins of about 27 kDa that are around 80% identical in sequence (for review, see Staswick 1994). Consistent with a storage role, the corresponding VSP genes are regulated developmentally in a source/sink-dependent manner, and are induced by the removal of seed pods and by the availability of excess nitrogen (Mason and Mullet 1990; Staswick et al. 1991; Mason et al. 1992; Sadka et al. 1994). On the other hand, their induction by stresses such as drought and high salt could suggest other roles are also possible.

Soybean VSP α and β are related to tomato acid phosphatase-1 (Aarts et al. 1991; Williamson and Colwell 1991). The VSPs occur as both homo and heterodimers and have acid phosphatase activities on *o*-carboxyphenyl phosphate ranging from 0.3 U mg⁻¹ protein for VSP α homodimer to 10 U mg⁻¹ protein for the heterodimer (DeWald et al. 1992). Compared to several other plant acid phosphatases these values are somewhat low, raising the question of the relevance of VSP catalytic activity. Although VSP levels and total acid phosphatase activity increased dramatically in leaves of depodded soybean plants, VSP α and β accounted for no more than 0.1% of the total acid phosphatase activity in these leaves (Staswick et al. 1994). Rather, an unrelated 51-kDa phosphatase was responsible for most of the activity, having a specific activity of 1,353 U mg⁻¹ protein.

Soybean root nodules also contain an acid phosphatase with 69% sequence identity with the VSPs (Penheiter et al. 1997; Penheiter 1998). Interestingly, the specific activity of recombinant nodule acid phosphatase (APase) on its optimal substrate (monophosphates) was about 30-fold higher than that previously reported for purified soybean VSP α/β , which was most active on polyphosphates (DeWald et al. 1992; Penheiter et al.

1998). The VSPs and nodule APase contain three short sequence motifs that suggest they belong to the bacterial class-B family of acid phosphatases, which are members of the haloacid dehalogenase (HAD) superfamily (Koonin and Tatusov 1994; Penheiter 1998; Morais et al. 2000; Selengut 2001). In plant and bacterial class-B APases the motif-I consensus sequence is FD[I,V]D[D,E]TXL. By analogy with the extensively characterized bacterial L-2-HAD (Liu et al. 1995) it was suggested that the first Asp in motif I is critical for enzyme activity because it makes a nucleophilic attack on the substrate phosphate (Penheiter 1998). However, the role of this Asp has not been experimentally demonstrated for plant APases. In L-2-HAD and in several related acid phosphatases, motif I is at the amino terminus (Selengut 2001), whereas it is near the center of the plant APases and VSPs. This suggests that the structure of the bacterial and plant proteins is somewhat different. Therefore, it is conceivable that one of the other acidic residues of motif I could play the catalytic role.

Interestingly, the putative catalytic Asp is present in nodule APase (Asp¹¹⁶), but it is substituted by Ser¹⁰⁶ and Gly¹⁰⁶ in VSP α and VSP β , respectively. It was suggested that the relatively low enzyme activity of soybean VSPs might result from this substitution of Asp in motif I (Penheiter 1998). Recombinant VSP has not been previously reported so it has not been possible to directly compare its activity with recombinant nodule APase.

The purpose of this study was to investigate the biochemical basis for the apparent low catalytic activity of soybean VSPs. Specifically, we tested whether restoring the putative catalytic Asp in motif I would elevate the activity of VSP α when expressed as a glutathione *S*-transferase (GST) fusion protein in *Escherichia coli*. We also compared the sequence of VSP cDNAs isolated from distant relatives of cultivated soybean and evaluated the phylogenetic relationship among 25 other plant proteins related to soybean VSPs.

Materials and methods

Plant and DNA material used

Perennial soybeans from the subgenus *Glycine* were grown in a temperature-regulated greenhouse and young leaves were collected for RNA extraction as described by Staswick (1997). cDNA libraries were constructed in lambda UNI ZAP-XR vectors (Stratagene) with reverse-transcribed poly(A)-mRNA from *G. falcata* and *G. tomentella* leaves. ³²P-labeled *VSPA* and *VSPB* DNA was used as a probe to screen the cDNA libraries. Low-stringency hybridizations were done overnight at 55°C and blots were washed twice with 1× SSC, 0.1% SDS solution at 53°C. Plaques that produced positive signals were selected and re-screened to homogeneity. After confirming the clones' relationship to the *VSPA/B* sequences by restriction endonuclease digestion, one cDNA from each species was sequenced.

A partial cDNA that included motif I for a *VSP* homologue was obtained from *G. curvata*. Total RNA was used with the 3' RACE System from GIBCO-BRL according to the manufacturer's instructions. The *VSPA*-related fragment was amplified with primers for sites flanking motif I that are relatively conserved in the other *VSPs* (Fwd: 5' GTGGAAGCACACAACATC 3', *VSPA* nucleotides 145–162; Rev: 5' TCTTCCTGACAAGAATA 3', *VSPA* nucleotides 485–502). PCR products were cloned into pGem-T-Easy vector (Promega) and putative *VSP* clones were sequenced, translated and aligned with other *VSP* sequences.

Preparation and expression of recombinant *APase* and *VSPA*

The soybean [*Glycine max* (L.) Merr.] root nodule *APase* and *VSPA* cDNAs were amplified by PCR with primers designed to eliminate the amino terminal signal peptide and incorporate *XhoI* sites compatible with the GST fusion expression vector. The primers used for PCR of *APase* were:

- Fwd, 5' GATCTCGAGATTCCGGAGGTATCATGC 3';
- Rev, 5' TCACTCGAGTCAACTAATGTAGTACATGGGATCAGG 3'.

For *VSPA* the primers were:

- Fwd, 5' CCACTCGAGAACACTGGCTATGGTG 3';
- Rev, 5' GATCTCGAGCTACTGAATGTAGTACAG 3'.

The PCR products were cloned into pGem-T-Easy vector, sequenced to verify their integrity, and then fused into the *XhoI* site of pGEX-4T-1. The *APase*:pGEX-4T-1 and *VSPA*:pGEX-4T-1 were transformed into *E. coli* strain BL21. For expression, cultures in Luria-Bertani (LB) medium containing 100 µg ml⁻¹ ampicillin were grown to OD₆₀₀ = 0.6 and then induced at room temperature with 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) for 4–6 h. Cells were harvested and washed once in Mes–NaOH (pH 6.0), and were stored as pellets at –80°C if not used immediately. Proteins were evaluated by SDS–PAGE and the molecular weights of the GST–*APase* and GST–*VSPα* fusions were approximately 52 kDa, as expected.

PCR was also used to introduce a site-specific mutation in *VSPα* converting Ser¹⁰⁶ to Asp using the wild-type *VSPA* as template. Because the targeted region was in the internal region of the cDNA, four oligonucleotides were used to generate two PCR fragments that could be joined by ligation at a common *Clal* restriction endonuclease site. Primers for the 5'-end cDNA fragment were:

- Fwd: 5' GATCTCGAGAACACTGGCTATGGTG 3';
- Rev: 5' ATCGATATCGAACACAAATGTGTCC-TTGGG 3'.

Primers for the 3'-end cDNA fragment were:

- Fwd: 5' GTGTTTCAGTATCGATGGCACCG 3';
- Rev: 5' GATCTCGAGCTACTGAATGTAGTACAG 3'.

The two PCR products were ligated into pGem-T-Easy, verified by sequencing and the reconstructed insert was then cloned into the *XhoI* site of pGEX4-T-1

Purification of GST fusion proteins

The GST–*APase* and GST–*VSPα* were partially purified from sonicated *E. coli* extracts by ammonium sulfate precipitation and ion-exchange chromatography. Ammonium sulfate was added to 60% saturation and the extract left on ice for 30 min. The precipitate was collected by centrifugation (12,000 rpm, 20 min) and resuspended in 4 ml of 50 mM Mes–NaOH (pH 6.0). The suspension was desalted by dialysis at 4°C against 20 mM Tris–HCl, pH 8.0 (two exchanges, 6–12 h). The dialyzed sample was then centrifuged at 15,000 rpm for 20 min. The supernatant was loaded onto a DEAE-50 cellulose column pre-equilibrated with 50 mM Mes–NaOH (pH 6.0). The eluted sample was concentrated osmotically and assayed for protein and *APase* activity. Fractions were analyzed by SDS–PAGE on 12% minigels (Bio-Rad) following the manufacturer's procedures. Protein concentrations were determined with the Bio-Rad DC protein assay according to the protocol supplied by the company. Activity of acid phosphatase was examined by monitoring phosphate released from *p*-nitrophenol phosphate (pNPP) at 405 nm. Activity determinations were routinely performed in triplicate at room temperature in 0.05 M Mes–NaOH (pH 6.0) containing 1 mM MgCl₂.

Enzymes for the analysis of mutated *VSPα* were affinity-purified with glutathione agarose. Induced cells were pelleted, resuspended in 10 mM Tris–HCl, 5 mM NaCl, and 3 mM MgCl₂ at pH 7.5, and then sonicated. Insoluble material was precipitated by centrifugation at 10,000 rpm for 10 min, and the supernatant was combined with glutathione agarose beads for 30 min at 4°C with continuous rotation. Beads were prepared in the same buffer as used for cell lysis. Recombinant proteins bound to the beads were washed 5 times with the same buffer and then digested with Thrombin Protease (Amersham) at 4°C for 6 h to release each protein from the GST fusion. The protease-digested supernatant was collected after centrifugation at 300 rpm for a few seconds. Protein amount was estimated using the Bio-Rad DC assay according to the manufacturer's instructions; bovine serum albumin was used as a standard. Proteins were analyzed by SDS–PAGE on 12% minigels (Bio-Rad) according to the manufacturer's procedure.

Assays for enzyme activity

Phosphatase activity was determined by the method of Fiske and Subbarow (1925) using a kit from Sigma

following the manufacturer's instructions. A variety of phosphorylated compounds were used as substrates: ADP, GMP, FMN, pNPP, P-tyrosine, and tripolyphosphate, each at 3 mM. Acid phosphatase reactions were performed in triplicate at 37°C in 0.05 M Mes-NaOH containing 1 mM MgCl₂, pH 6.0. Kinetic measurements were determined by activity assays at pH 6.0 with pNPP and GMP substrate concentrations ranging from 0 to 6 mM. The kinetic parameters V_{max} and K_m were evaluated by the Michaelis-Menten method.

Analysis of DNA and protein sequence

DNA sequencing was done by the University of Nebraska Genome Core Research Facility. DNA and protein sequence was analyzed with SeqWeb v. 2 (Accelrys). For phylogenetic analysis VSP-related plant sequences were identified by blastP searches of the non-redundant translated database using soybean VSP α (AAA34020) and tomato APase-1 (AAA34135) as the query sequences. Non-redundant sequences were judged to be full length based on comparison with those known to be complete (e.g. soybean VSPs, soybean nodule APase, tomato APase-1). Phylogenetic relationships were analyzed with Grow Tree using the default settings (Kimura distance, neighbor distance, blossom62 scoring matrix, gap penalty 8). The tree was displayed using the Nexus output with Tree View 1.6.6 (<http://taxonomy.zoology.gla.ac.uk/rod/rod.html>). Sequences for the motif-I region included those used for tree construction, three partial sequences from the blastP search (PPI309082, AY106317, CAB71336) and the partial sequence we derived for *G. curvata*.

Results

The putative nucleophilic Asp is substituted in VSPs from perennial soybeans

We first isolated cDNA clones from wild perennial relatives of cultivated soybean in order to determine whether the absence of the putative catalytic Asp was unique to the two *G. max* VSPs, or whether this was more generally the case in the *Glycine* genus. Full-length clones from *G. falcata* and *G. tomentella* had predicted open reading frames encoding 253 amino acids, compared with 254 for VSP α and VSP β from *G. max*. Analysis with Grow Tree indicated the perennial proteins were more closely related to VSP α than VSP β . The proteins from *G. falcata* and *G. tomentella* shared 85% sequence identity with each other and were each 81 and 76% identical with VSP α and VSP β , respectively. Based on this evidence we have classified these as VSP α proteins. Sequence identity with soybean nodule APase was 64 and 62% for the *G. falcata* and *G. tomentella* proteins, respectively.

The sequence of 22 amino acids surrounding motif I for each of these proteins is shown in Fig. 1, along with

the same region from previously characterized *G. max* VSPs and from the VSP-like pod storage protein (PSP) from *Phaseolus vulgaris* (Zhong et al. 1997). In addition, a translated sequence from a partial cDNA from *G. curvata* is also shown. Like the *G. max* VSPs, those from all three perennial species lacked the putative catalytic Asp, which in each case was substituted by a Ser residue. The Asp is also substituted in PSP by Asn. Together these are denoted as Group-I proteins. The sequence from this region was also compared to that of 25 other plant VSP/APase-like proteins. Included in this second group are two enzymes that are known to have acid phosphatase activity, the soybean nodule APase and tomato Apase-1 (Aarts et al. 1991; Williamson and Colwell 1991; Penheiter et al. 1997). All of the Group-II proteins from eight different plant species contained the Asp that is predicted to have a catalytic role. Two arabidopsis genes have been called *VSP1* and *VSP2* based on predicted amino acid sequence similarity with the soybean VSPs (Utsugi et al. 1998), but the proteins they encode have not been evaluated for their potential enzymatic activity. The remainder are putative proteins

Species	Accession	Name/Function	Motif I region
Group I			
Gm	AAA34020	VSP α	DTFVFSIDGTVLSNIPYYKKHG
Gm	AAA34021	VSP β	DI FIFGIDNTVLSNIPYYEKHG
Gt	AY523603	VSP α	DVFVFSIDATVLSNVPYYSEHG
Gf	AY523602	VSP α	DVFLFSIDGTALS NVPYYSEHG
Gc	AY523604	VSP	DVFLFSIDGTVLSNVPYYSEHG
Pv	BAA23563	PSP (Pod Storage)	DVILEFDIGTALS NIPYYSQHG
Group II			
Gm	CAA11075	nodule Apase	TAWVFDIDETLLSNLPYYADHG
Le	AAA34135	Apase-1	DVWIFDVDDTLLSNLPYYSDHR
Gm	BAB86895	pathogen-associated	DAWILDVDDTCISNIDYYKGRR
Pv	AAL17638	pathogen-associated	DAWILDVDDTCISNVSYKTKR
At	At5g24780	VSP1	NVWIFDLDDTLLSSIPYYAKYG
At	At5g24770	VSP2	NVWIFDLDDTLLSSIPYYAKYG
At	NP194656		DAWVFDIDETLLSNIEYYKANG
At	At4g25150		DIWIFDIDETLLSNLPYYMEHG
At	At5g51260		DIWIFDIDETLLSNLPYYIDHG
At	At4g29260		DVWIFDIDETLLTNIYYKAHG
At	At2g38600		DAWILDVDDTFCFSNVFYYRLKR
At	At1g04040		DAWIFDIDETLLSTIPYKKNHG
At	At5g44020		DAWIFDIDETLLSTIPYKKNHG
At	At4g29270		DAWVFDIDETLLSNIEYYKANG
Os	AK058604		DAWVFDVDETTLLSNLPYYADHG
Os	AK059825		EI WVFDIDETSLSNLPYYAKHG
Os	AK104624		EI WVFDVDDTALS TVPYQANHG
Os	AK064745		DAWVFDVDDTCLS NLFYQAKRQ
Os	AK069355		PAWVFDVDETTLLTNPYYAVNG
Os	AK104267		EI WVFDVDETTALS TL PYYQANHG
Os	AK109335		EI WVFDVDETTALS TL PYYQAKHG
Os	NM189664		ATWVFDVDETTALS HVKFFYKKNHG
Pp	PPI309082		DAWVFDADETLLSNIPYYENYE
Zm	AY106317		EVWVFDIDETSLSNLPYYATHG
Hv	CAB71336		DVWVFDIDETLLSNLPYYATHG

Fig. 1 Comparison of conserved motif I from soybean VSPs and related plant APases. The invariant nucleophilic Asp of Group-II proteins is substituted by the boxed residue indicated for each Group-I protein (Ser¹⁰⁶ in *G. max* VSP α). Invariant or highly conserved Group-II residues are shaded and represented similarly if present in Group I. Species designations are *At*, *Arabidopsis thaliana*; *Gc*, *Glycine curvata*; *Gf*, *Glycine falcata*; *Gt*, *Glycine tomentella*; *Gm*, *Glycine max*; *Hv*, *Hordeum vulgare*; *Le*, *Lycopersicon esculentum*; *Os*, *Oryza sativa*; *Pv*, *Phaseolus vulgaris*; *Pp*, *Pinus pinaster*; *Zm*, *Zea mays*

based on complete or partial database nucleotide sequences. In addition to the putative nucleophilic Asp, two other residues conserved in Group-II proteins are absent in Group I. These are Trp three residues upstream and Asp or Glu three residues downstream of the putative nucleophilic Asp

Nodule APase expressed in *E. coli* is catalytically active

Recovery of soybean VSPs and nodule APase in active form from *E. coli* had not previously been reported. In order to test the feasibility of obtaining these enzymes from bacteria, VSP α and nodule APase were each expressed as GST fusion proteins. The constructs excluded the signal peptide found in each full-length cDNA. GST-APase and GST-VSP α were partially purified by DEAE ion-exchange chromatography following precipitation with 60% ammonium sulfate (Table 1). The specific activity of GST-APase was about 72 $\mu\text{mol min}^{-1} \text{mg}^{-1}$ using 3 mM pNPP as the substrate. This was similar to the activity reported previously for this enzyme expressed in yeast and purified in a similar manner (66.7 $\mu\text{mol min}^{-1} \text{mg}^{-1}$ with 5 mM pNPP; Penheiter 1998). This result indicated that APase could be recovered from *E. coli* in active form.

GST-APase was assayed for its pH optimum with pNPP as substrate. Figure 2 shows that the enzyme had a broad pH activity profile typical of acid phosphatases, with a maximum occurring around pH 6.0. This is similar to the pH optimum found for APase expressed in yeast (Penheiter 1998). Therefore, all further kinetic studies were carried out at this pH. To evaluate the substrate specificity for GST-APase, several phosphorylated substrates were tested (Table 2). The highest activity was observed with the monophosphorylated substrate 5'-GMP, which was in agreement with previous results for native and recombinant APase from yeast (Penheiter 1998). Several other compounds (FMN, ADP, and P-tyrosine) were also dephosphorylated, but at a much lower rate than 5'-GMP. In contrast, we found no evidence that triphosphorylated substrate was hydrolyzed. Based on these data, the kinetics for the two most active substrates was determined and is shown in Table 3. Highest affinity was observed for 5'-GMP ($K_m=0.9$ mM). The observed K_m value for pNPP was somewhat higher at

8.8 mM. The V_{max} value for pNPP was greater than that calculated for 5'-GMP and the V_{max}/K_m values for pNPP and 5'-GMP were 279 and 1,926, respectively.

In contrast to GST-APase the VSP α fusion protein possessed little phosphatase activity. The only significant activity among the substrates tested was with GMP and pNPP (22.6 and 3.2 U mg^{-1} , respectively). This was about 20-fold lower than for GST-APase with the same substrates. Extracts from *E. coli* expressing GST alone exhibited no detectable activity with any of these substrates (data not shown), indicating that the low activity observed for GST-VSP α was in fact due to the recombinant enzyme and not endogenous phosphatases co-purified from *E. coli*. As observed for nodule GST-APase, P-tyrosine, ADP and triphosphorylated substrate, were poor substrates for GST-VSP α .

Previous studies showed that cyclic nucleotides inhibited purified nodule APase (Penheiter 1998). To determine the effect of this inhibitor on GST-APase, enzyme activity was assayed in the presence of cAMP. Substrate pNPP concentrations ranging from 2.5 to 20 mM, and two concentrations of inhibitor, 5 and 50 μM cAMP, were used. The K_i value of GST-APase was about 15 μM (not shown), only slightly higher than that determined for APase from nodule extract (12 μM ; Penheiter 1998). Collectively, these results indicate that in all regards tested nodule GST-APase from *E. coli* behaves similarly to native APase.

Mutation of VSP α increases its acid phosphatase activity

We next examined whether conversion of Ser¹⁰⁶ to Asp in VSP α would alter its phosphatase activity relative to the wild-type protein. For this assay, highly purified enzymes were recovered by isolation on glutathione agarose followed by cleavage with thrombin to isolate the enzyme from GST. Analysis of wild-type and mutant VSP α , as well as nodule APase, by SDS-PAGE indicated the major protein bands in each case had the expected molecular weights of approximately 27 kDa after thrombin cleavage (Fig. 3). For each protein only minor contaminants were evident.

The result of the assay of purified enzyme with five phosphorylated substrates is shown in Table 4. The activity for thrombin-cleaved APase on GMP was 845 U mg^{-1} . This was the most active substrate, as

Table 1 Purification of soybean (*Glycine max*) nodule APase in *Escherichia coli*

Step	Total protein ^a (mg)	Total activity ^{a,b} (U)	Specific activity ^a (U/mg)	Purification (-fold)	Recovery (%)
Crude extract	251.0 ± 19	6200 ± 900	24.5 ± 1	1.0	100
60% (NH ₄) ₂ SO ₄	119.0 ± 9	4413 ± 548	37.2 ± 1	1.5	71
DEAE	15.2 ± 1	1088 ± 58	72.1 ± 4	3.0	18

^aMean ± SD, $n=3$. 0.15 mg protein was used in each assay

^bOne unit = 1 μmol *p*-nitrophenol released from pNPP (3 mM) per minute at room temperature in 0.05 M Mes-NaOH (pH 6.0) + 1 mM MgCl₂

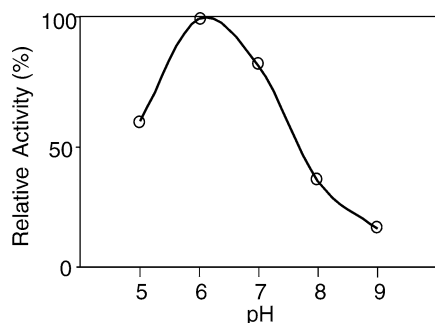


Fig. 2 The effect of pH on soybean (*Glycine max*) nodule APase activity. Enzyme purified from *Escherichia coli* was assayed with pNPP as a substrate at the indicated pH values. Activity is expressed as percent of the value at pH 6. Mes–NaOH was the buffer for pH 5 and 6, and Tris–HCl for pH 7–9

Table 2 Activity of soybean nodule APase, and VSP α fusions against a variety of substrates

Substrates	APase ^{a,b} (U/mg)	VSP α ^a (U/mg)
pNPP	72.1 \pm 4	3.2 \pm 1
ADP	28.0 \pm 2	0
GMP	473.3 \pm 9	22.6 \pm 1
FMN	26.3 \pm 1	0
O-P-Tyrosine	24.3 \pm 1	0.5 \pm 0.1
Tripolyphosphate	0	0

^aOne unit = 1 μ mol *p*-nitrophenol released from pNPP (3 mM) per minute at room temperature in 0.05 M Mes–NaOH (pH 6.0) + 1 mM MgCl₂.

^bMean \pm SD, *n* = 3. 0.15 mg protein was used in each assay

Table 3 Kinetic values of the soybean nodule APase fusion from *E. coli*

Substrates	K_m ^a (mM)	V_{max} ^{a,b} (U/mg)	V_{max}/K_m ^a
5'-GMP	0.9 \pm 0.2	2160.5 \pm 99	1926.5 \pm 92
pNPP	8.8 \pm 0.6	2687.0 \pm 82	279.5 \pm 20

^aMean \pm SD, *n* = 3. 0.15 mg protein was used in each assay

^bOne unit = 1 μ mol phosphate released per minute at room temperature in 0.05 M Mes–NaOH (pH 6.0) + 1 mM MgCl₂

noted earlier for the partially purified GST–APase. Wild-type VSP α cleaved from GST yielded 30- to 40-fold lower activity on both pNPP and GMP. In contrast, the conversion of Ser¹⁰⁶ to Asp dramatically increased the enzyme activity of VSP α on GMP to 439 U mg⁻¹, almost 20-fold higher than for wild-type VSP α and about half that of nodule APase. VSP α Asp¹⁰⁶ also hydrolyzed pNPP at 35 U mg⁻¹ and P-tyrosine at 20 U mg⁻¹. This was about one-third of the level found for nodule APase with pNPP as a substrate and about 10-fold higher than for wild-type VSP α on these substrates. As for nodule APase, the mutant VSP α had no detectable activity towards FMN or tripolyphosphate. These results establish that the major reason for the low catalytic activity of VSP α is the single amino acid sub-

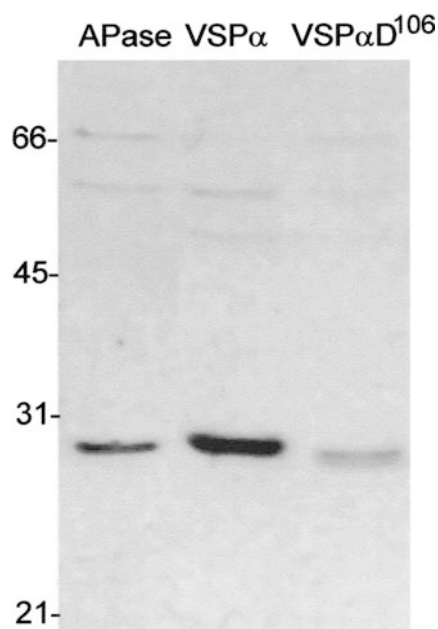


Fig. 3 Analysis of glutathione agarose-purified GST fusion proteins. Major bands at about 27 kDa are wild-type nodule APase, VSP α and mutated VSP α Asp¹⁰⁶, as indicated. Position of molecular markers is indicated to the left in kDa. Purified proteins were loaded on 12% SDS–PAGE gels

Table 4 Phosphatase activity of purified proteins. Phosphatase activity of soybean VSP α , mutant VSP α and nodule APase with a variety of phosphorylated compounds (3 mM). Reactions were assayed in triplicate using 0.05 mM Tris–NaOH (pH 6.0) + 1 mM MgCl₂

Substrate	Activity (U/mg) ^{a,b}		
	APase	VSP α Ser106Asp	VSP α
pNPP	97 \pm 11	35 \pm 7	3.2 \pm 1
GMP	845 \pm 152	439 \pm 90	22.6 \pm 1
FMN	0	0	0
P-tyrosine	31 \pm 4	20 \pm 0.7	0.5 \pm 0.1
Tripolyphosphate	0	0	0

^aOne unit = 1 μ mol of phosphate released per minute at room temperature in 0.05 M Mes–NaOH (pH 6.0) + 1 mM MgCl₂

^bMean \pm SD, *n* = 3. 1 mg protein used in each assay

stitution of Ser¹⁰⁶ for the catalytic Asp found in this family of APases.

Discussion

Penheiter et al. (1998) previously reported that nodule APase expressed in *E. coli* as a His tag fusion was found in inclusion bodies rather than as a soluble protein. Our results show that expression of both VSP α and nodule APase as GST fusions is a viable means to obtain these enzymes in soluble and active form from bacteria. This expression system permitted the rapid affinity purification of the enzymes, which facilitated the study of VSP α

by mutational analysis. The specific activity of the glutathione-purified and thrombin-cleaved APase on GMP was about 2-fold higher than that purified by ion exchange. The substrate specificity was also similar to that found for the native enzyme, confirming that purification by this method yields enzyme that is representative of the natural enzyme. We were also able to directly compare the activity of recombinant nodule APase with that of VSP α , because the two enzymes were produced and purified under identical conditions. Wild-type VSP α had about 40-fold lower activity than nodule APase, confirming previous indications that soybean VSP is a relatively weak acid phosphatase. A recent study showed that a marked down-regulation of the *VSP* genes, resulting in only about 2% of the normal VSP protein, had no adverse effect on plant growth or productivity (Staswick et al. 2001). While this does not necessarily mean that VSPs have no catalytic role, at least the role is not essential and is possibly made redundant by other enzymes. The same is apparently true for the storage role that has been assumed for these VSPs.

Site-specific mutation of the first Asp of motif I in bacterial L-2-HAD and in magnesium-dependent acid phosphatase-1 from mouse has shown this residue is critical for catalytic activity (Liu et al. 1995; Selengut 2001). The functional relevance of this residue had not previously been determined in plant enzymes. This was important to establish because all Group-II plant proteins, including two demonstrated to be acid phosphatases, contain two additional acidic residues just down stream of the proposed catalytic Asp in motif I (DxD[D/E]). It was conceivable that one of these, rather than the first Asp, might be structurally positioned to act as a nucleophile in these enzymes. We found that substitution of Ser¹⁰⁶ with Asp at the position corresponding with the proposed catalytic Asp increased VSP α activity nearly 20-fold, confirming its essential role in catalysis in the plant enzymes. Interestingly, all Group-I proteins also lacked the invariant Trp and the conserved acidic residue found in Group-II proteins corresponding to positions 103 and 109 in VSP α , respectively. It is possible that these residues also enhance acid phosphatase activity, perhaps affecting the substrate-binding site.

Although Asp¹⁰⁶ is clearly of major importance, wild-type recombinant VSP α did have low activity, as was previously reported for native VSPs (DeWald et al. 1992). It is not clear why this is so if Asp¹⁰⁶ is essential for catalysis. All Group-I VSPs retain the Asp corresponding to position 108 in VSP α . It would be of interest to determine whether Asp¹⁰⁸ can also act as the nucleophile for catalysis in VSPs, if only inefficiently. VSP α Asp¹⁰⁶ had only half the activity of nodule APase. It is possible that other residues in VSP α are also sub-optimal for activity. The fact that the VSP β homodimer purified from plants had a specific activity about 10-fold higher than the VSP α homodimer (DeWald et al. 1992) supports this possibility.

In contrast to the earlier finding of highest activity on polyphosphates for native VSP α/β (DeWald et al. 1992),

we found no detectable activity on this substrate. One possible explanation is that we evaluated only VSP α , which presumably forms a homodimer in *E. coli*. The native VSP α homodimer from plant extracts had much lower activity overall than VSP α/β and was relatively less active on polyphosphates than was the heterodimer. We chose to investigate VSP α because it is the more abundant of the two subunits in soybean during normal plant growth. The previous analysis of native VSP (DeWald et al. 1992) also did not include GMP, which was the substrate giving the highest activity in our analysis. It will be important in future studies to determine the level of activity and substrate specificity for the β VSP that has had Asp¹⁰⁶ restored. VSP β Asp¹⁰⁶ could be expressed alone in *E. coli* and along with VSP α Asp¹⁰⁶, in which case it could presumably form the heterodimer as in soybean.

The sequence of VSPs from perennial soybeans had not previously been reported. We characterized two new VSP cDNAs and found they encode proteins with high sequence identity to the *G. max* VSPs. The VSPs from the perennial *Glycine* spp. are nearly identical in size to the *G. max* VSPs and are expected to have signal peptides of 29 amino residues. Previous analysis indicated there was considerable apparent size heterogeneity among proteins from the perennials that cross-reacted with VSP antisera (Staswick 1997). Our results here suggest that part of the heterogeneity on SDS-PAGE is due to variable mobility caused by charge differences or glycosylation, rather than differences in polypeptide length. This is also the basis for the apparent size difference between *G. max* VSP α and VSP β . In the earlier study, leaf extracts from *G. falcata* produced a single VSP immuno cross-reacting band while *G. tomentella* yielded two distinct bands. This suggests there may be a second gene in *G. tomentella*, possibly a VSP β -type protein as in *G. max*. Like the *G. max* VSPs, those from the perennial species also lacked the first Asp residue in motif I, suggesting that they are probably catalytically inefficient as well. It appears that the loss of the catalytic Asp originated before the domestication of *Glycine max*. We would also predict that the enzyme activity of *P. vulgaris* PSP would be low due to the absence of the catalytic Asp in this protein as well.

A phylogenetic analysis of the full-length sequences of 30 VSP and VSP-like plant proteins suggests that catalytic inactivation of VSPs may not be a recent event (Fig. 4). Of the currently known VSP-like plant proteins the one most closely related to the *Glycine* VSPs is PSP from *P. vulgaris*. Together, these constitute the Group-I proteins described in Fig. 1. All of the Group-II proteins have the nucleophilic Asp, including the *G. max* nodule and pathogenesis-associated APases (Acc. No. BAB86895), and are more distantly related to the VSPs than is PSP. These relationships suggest that enzymatic inactivation of an ancestral APase may have occurred after the divergence of VSPs from soybean root nodule APase, but before the divergence of *Phaseolus* from *Glycine*. We cannot, however, rule out the possibility

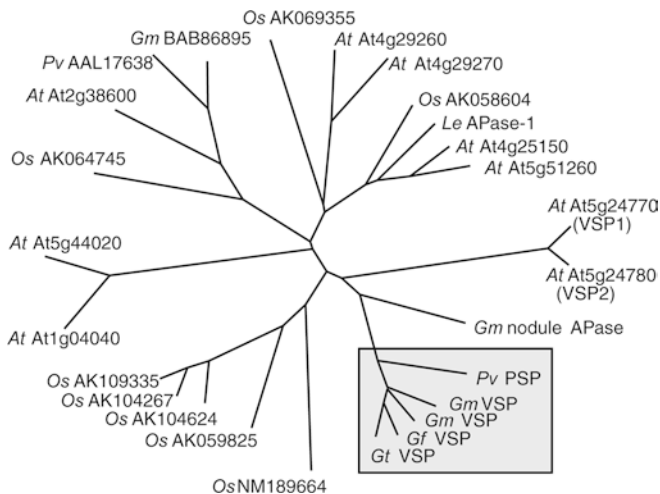


Fig. 4 Phylogenetic analysis of plant VSPs and VSP-like APases. Group-II proteins lacking the nucleophilic Asp residue are boxed. Proteins are designated by a two-letter genus/species designation (see Fig. 1), followed by the database accession number

that PSP lost the nucleophilic Asp independently of the soybean VSPs.

Our results document clearly that inactivity in VSP α is due primarily to a single amino acid substitution in an APase active site, because we have been able to restore activity to this protein by mutation. It is somewhat surprising that other residues necessary for catalysis had not also diverged, since there would not seem to be a reason to retain them in this inactive enzyme. It is possible that some of these must be maintained for proper folding and protein stability. However, the low overall level of sequence conservation among VSPs comprising Group-I proteins, as well as in the Group-II VSP-like acid phosphatases, indicates there is wide room for sequence variation in this family of proteins.

Other storage proteins also appear to be inactivated by simple mutations, although in each case mentioned below it has not been experimentally demonstrated that repairing the proposed defect alone is sufficient to restore activity. For example, sequence comparison suggested the RNase-like rhizome storage protein of *Calystegia sepium* is inactive due to substitution of a conserved His that is involved in catalysis in related plant RNases (Van Damme et al. 2000). Type-2 ribosome-inactivating proteins (RIP) can also function as storage reserves. A critical component of their activity is a B-chain polypeptide with agglutinating activity. *Sambucus nigra* contains an abundant RIP-like protein that is inactive apparently because its B-chain lacks critical residues for agglutinating activity. Substitution of one of these in a closely related active B-chain reduced binding activity 50%, and a second mutation further lowered activity (Chen et al. 2002). Structural modeling of a lectin-like storage protein from *Cladostrius lutea* also suggested its inactivity is due to three extra amino acids in the presumed carbohydrate-binding site (Van Damme et al. 1995). In each of these cases it would be of interest

to correct the apparent defects in the inactive proteins and determine whether this alone would restore activity, or if additional residues must also be changed.

An intriguing question is whether loss of acid phosphatase activity was required for the VSPs to function as storage proteins, or whether they were recruited for storage proteins because they were redundant and not serving another critical role. Storage proteins by definition accumulate to a high level. It may be that over accumulation of an active phosphatase would raise the cellular phosphate pool to toxic levels. It is also possible that in their role as storage proteins, legume VSPs have simply lost a non-essential function, but that loss is not required. One way to address this question would be to express the gene for activated VSP α Asp¹⁰⁶ in transgenic soybean under its native promoter to determine whether VSPs with high catalytic activity would be tolerated.

Because the complete genome is available we should have identified all putative VSP-like proteins from arabidopsis. None of the nine proteins that were found lacked the nucleophilic Asp. This suggests that if any of these are storage proteins their enzymatic inactivation is not required or involves other amino acids not yet identified. It has not been conclusively established that arabidopsis contains proteins that function specifically as storage reserves. Two genes have been called *VSP1* and *VSP2* based on sequence homology and some regulatory similarity with the soybean *VSP* genes. However, their relative abundance has not been well characterized, so it is not clear whether they could have a significant storage role. Retention of the catalytic Asp in these proteins suggests they may be active enzymes. It should also be noted that regulatory similarity with the soybean *VSP* genes does necessarily imply they encode storage proteins. For example, soybean *VSPs* respond to several stress factors that may also trigger the production of APases that function in stress response. Depodding of soybean plants dramatically elevated acid phosphatase activity from a 51-kDa protein, along with the rise in VSP level. But the low abundance of the 51-kDa protein indicates it is not a significant storage reserve (Staswick et al. 1994).

In summary, the five soybean VSPs from four distinct species of *Glycine* sequenced to date, as well as a PSP from *P. vulgaris*, all lack an Asp residue that we have demonstrated is critical for high catalytic function in VSP α . A single amino acid substitution replacing the native Ser with Asp in VSP α increased its acid phosphatase activity about 20-fold, to a level similar to that of nodule APase. This establishes that the major biochemical basis for the low enzymatic activity of VSP α is the substitution of this single amino acid residue, and that other residues necessary for high enzyme activity are retained in VSP α .

Acknowledgements We thank Martha Rowe for her technical assistance in carrying out this study. This is Journal Series Paper No. 14,443. A contribution of the University of Nebraska Agricultural Research Division, Lincoln, NE 68583. This work was

supported in part by NIH Grant Number 1 P20 RR16469 from the BRIN Program of the National Center for Research Resources (G.S.).

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