

Invertases involved in the development of the parasitic plant *Phelipanche ramosa*: characterization of the dominant soluble acid isoform, PrSAI1

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SUMMARY

Phelipanche ramosa L. parasitizes major crops, acting as a competitive sink for host photoassimilates, especially sucrose. An understanding of the mechanisms of sucrose utilization in parasites is an important step in the development of new control methods. Therefore, in this study, we characterized the invertase gene family in *P. ramosa* and analysed its involvement in plant development. Invertase-encoded cDNAs were isolated using degenerate primers corresponding to highly conserved regions of invertases. In addition to enzyme assays, gene expression was analysed using real-time quantitative reverse transcriptase-polymerase chain reaction during overall plant development. The dominant isoform was purified and sequenced using electrospray ionization-liquid chromatography-tandem mass spectrometry (ESI-LC-MS/MS). Five invertase-encoded cDNAs were thus characterized, including *PrSai1* which encodes a soluble acid invertase (SAI). Of the five invertases, *PrSai1* transcripts and SAI activity were dominant in growing organs. The most active invertase corresponded to the *PrSai1* gene product. The purified PrSAI1 displayed low *pI* and optimal pH values, specificity for β -fructofuranosides and inhibition by metallic ions and competitive inhibition by fructose. PrSAI1 is a typical vacuolar SAI that is actively involved in growth following both germination and attachment to host roots. In addition, germinated seeds displayed enhanced cell wall invertase activity (PrCWI) in comparison with preconditioned seeds, suggesting the contribution of this activity in the sink strength of infected roots during the subsequent step of root penetration. Our results show that PrSAI1 and, possibly, PrCWI constitute good targets for the development of new transgenic resistance in host plants using proteinaceous inhibitors or silencing strategies.

INTRODUCTION

Broomrapes consist of almost 170 species devoid of chlorophyll and are thereby obligate parasitic plants. Seven species, including *Phelipanche ramosa* L. Pomel (syn. *Orobanche ramosa* L.), are serious parasitic weeds that cause great losses in many economically important crops around the Mediterranean Sea and in Asia Minor (Parker, 2009). *Phelipanche ramosa* attacks Solanaceae crops most severely and is well established in long winter crops (e.g. rapeseed) as well as in shorter spring and summer crops (hemp, melon, tobacco, tomato). Success in controlling broomrape is rare (Rispaill *et al.*, 2007) and there is an urgent need to continue to screen genetic resources and to characterize resistance mechanisms (Pérez-de-Luque *et al.*, 2009). An understanding of the physiological and metabolic features of the parasite is particularly pertinent, because new control methods that target parasite physiology or metabolism are highly desirable (Aly *et al.*, 2009).

Broomrape development occurs in five stages (Labrousse *et al.*, 2001): germination (stage I), which is elicited by stimulants secreted by the roots of host plants; attachment to the host vascular system (stage II) through a haustorium containing both xylem and phloem that serves as both an attaching organ and a bridge for water and nutrient transfer from the host; establishment and growth of a tubercle (stage III), which gives rise to a subterranean shoot (stage IV) and then to a flowering spike after emergence from the soil (stage V). Via the haustorium, the parasite connects to the phloem sieve elements of the host plant (Hibberd and Jeschke, 2001), and thereby becomes a major competitive sink for photoassimilates. Sucrose has long been known to be the main host-derived compound (Aber *et al.*, 1983). However, there is evidence, notably in *P. ramosa*, that sucrose is not accumulated in the parasite, but is utilized for growth and for the accumulation of hexoses and, to a lesser extent, mannitol and starch (Abbes *et al.*, 2009; Delavault *et al.*, 2002; Singh *et al.*, 1968). Broomrape displays both low transpiration rates and low water flux in xylem elements (Hibberd *et al.*, 1999); osmotic

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adjustment through the strong accumulation of potassium and soluble sugars is thus considered to be an essential process in water movement from the host plant, and thus growth of the parasite (Abbes *et al.*, 2009).

Considering the major role of soluble carbohydrates in osmotic adjustment and the sink strength of the parasite, there is a need to better understand the primary steps of sucrose allocation, which consist of the degradation of host-derived sucrose. Sucrose degradation is catalysed by two types of enzyme in plants: invertases (EC 3.2.1.26) and sucrose synthases (SuSys) (EC 2.4.1.13). SuSys are glycosyl transferases which convert sucrose into fructose and UDP-glucose in the presence of uridine diphosphate (UDP), whereas invertases hydrolyse sucrose into glucose and fructose. Sucrose-degrading enzymes play multiple pivotal roles in plants: sugar transport and partitioning, cell turgor and expansion, response to insect wounding and pathogen infection, sugar signalling and plant development (Roitsch and Gonzalez, 2004; Sturm, 1999; Sturm and Tang, 1999). On the basis of their solubility, subcellular localization, pH optima and isoelectric point, three types of invertase isoenzyme have been identified: acid invertases (AIs), including soluble/vacuolar isoforms (soluble acid invertase, SAI) and insoluble/cell wall-bound isoforms (cell wall invertase, CWI), and neutral/cytosolic invertases (soluble neutral invertase, SNI). Both vacuolar and cell wall isoenzymes are β -fructofuranosidases which share a high degree of overall sequence homology. They correspond to acid and glycosylated invertases, showing an acidic pH optimum, and accept other fructofuranosides, such as stachyose and raffinose, as substrates. Vacuolar isoforms control sugar composition in the vacuole and the remobilization of sucrose for metabolic processes, and CWI isoforms are ionically bound to the cell wall and coordinate phloem unloading in sink cells with hexose transporters. Moreover, the sucrose gradient mediated by CWI at sites of phloem unloading is crucial for the normal development of sink organs (Zanor *et al.*, 2009). SNI proteins are unglycosylated proteins and have neutral/alkaline pH optima and high specificity for sucrose. For a long time, SNI isoenzymes had only been described in the cytosol, where they play a housekeeping role and maintain hexose concentrations. Recent work has demonstrated that cytosolic SNIs are essential for normal growth and that mitochondrial and chloroplastic isoforms contribute to intracellular carbon trafficking (Barratt *et al.*, 2009; Vargas *et al.*, 2008). Similarly, SuSys are located in the cytosol and in mitochondria and possibly in plastids as well (Hennen-Bierwagen *et al.*, 2009). Since SuSy is commonly associated with glucan synthesis, it is a key regulator of starch accumulation and thickening of secondary cell walls in the xylem (Baroja-Fernández *et al.*, 2009; Coleman *et al.*, 2009).

Information on the metabolic and molecular bases of parasitic plant–plant interactions is rare. Glucose and fructose are

strongly accumulated in broomrape from host-derived sucrose, suggesting that invertases are the dominant sucrose-degrading enzyme in the parasite (Abbes *et al.*, 2009; Delavault *et al.*, 2002). The present study characterizes the invertase family of the parasite *P. ramosa* at both molecular and biochemical levels, and thereby its involvement in the development of the parasitic plant. Patterns of expression of the different invertase-encoding genes and of SAI, SNI and CWI activities were analysed during germination and post-attachment stages of broomrape development. In addition, the most active invertase isoform was purified and partially sequenced. We characterized the gene encoding this dominant invertase and its associated isoform, which are closely involved in broomrape development before and after attachment to the host plant. This work provides a better understanding of the interaction and opens up new perspectives in host resistance engineering.

RESULTS

Cloning and characterization of *P. ramosa* invertase cDNAs

Five partial *P. ramosa* cDNAs were obtained by reverse transcriptase-polymerase chain reaction (RT-PCR) using sets of primers designed from conserved regions of plant invertase sequences. Using rapid amplification of cDNA ends (RACE) strategies, four full-length cDNAs, *PrSai1*, *PrCwi*, *PrSni1* and *PrSni2* (accession numbers: GU997130, GU99132, GU99133 and GU99134, respectively), and one partial cDNA, *PrSai2* (accession number: GU997131), were isolated. The characteristics of *P. ramosa* invertase cDNAs and the corresponding protein are presented in Table 1.

Three distinct cDNAs encoding AIs were identified. The amino acid sequence encoded by *PrSai1* has 65.6% identity to a *Nicotiana tabacum* vacuolar invertase (CAC83577). The partial *PrSai2*-encoded sequence shares 71% identity with a *Daucus carota* SAI (CAA77266) (Sturm, 1996). The *PrCwi*-encoded sequence shares 66.6% identity with a *Solanum lycopersicum* CWI (AAM28823) (Fridman and Zamir, 2003). All three deduced proteins contain the conserved motifs of plant AIs, including a β -fructosidase motif 'DPNG' and a putative catalytic domain 'WEC(I/V/P)D' (Roitsch *et al.*, 1995; Sturm and Chrispeels, 1990). The plant fructan-metabolizing fructosyltransferases share motifs that are conserved in AIs (Vijn and Smeekens, 1999). Nevertheless, in contrast with plant fructosyltransferases, AIs contain the amino acid triplet WIN or WMN (Gallagher *et al.*, 2004). Thus, the presence of the WIN triplet in the *PrSai1*- and *PrCwi*-encoded sequences (residues 139–141 and 57–59, respectively) and WMN in the *PrSai2*-encoded sequence confirmed that the three genes encoded AIs. As

Table 1 Characteristics of *Phelipanche ramosa* invertase cDNAs and their corresponding proteins.

Gene	cDNA (bp)	5'-UTR (bp)	ORF (bp)	3'-UTR (bp)	Protein (kDa)	pI (theor.)	Identity (%)		Accession number
<i>PrSai1</i>	2160	62	1986	112	73.1	5.42	65.6	<i>Nicotiana tabacum</i>	CAC83577
							64.6	<i>Daucus carota</i>	CAA77267
<i>PrSai2*</i>	1711	—	—	74	—	—	71	<i>D. carota</i>	CAA77266
							65.4	<i>Camellia sinensis</i>	BAF34362
<i>PrCwi</i>	2187	40	1761	386	66.2	8.68	66.6	<i>Lycopersicon esculentum</i>	AAM28823
							63.8	<i>D. carota</i>	CAA55188
<i>PrSni1</i>	2032	115	1710	207	64.8	6.72	80.9	<i>L. esculentum</i>	ABQ28669
							78.6	<i>Vitis vinifera</i>	CAP59642
<i>PrSni2</i>	2445	67	2001	377	76.4	7.16	69	<i>V. vinifera</i>	ABS52644
							67.9	<i>D. carota</i>	CAA76145

The percentage of identity is that with the closest annotated homologue.

*Not full-length.

ORF, open reading frame; Pr, *Phelipanche ramosa*; UTR, untranslated region.

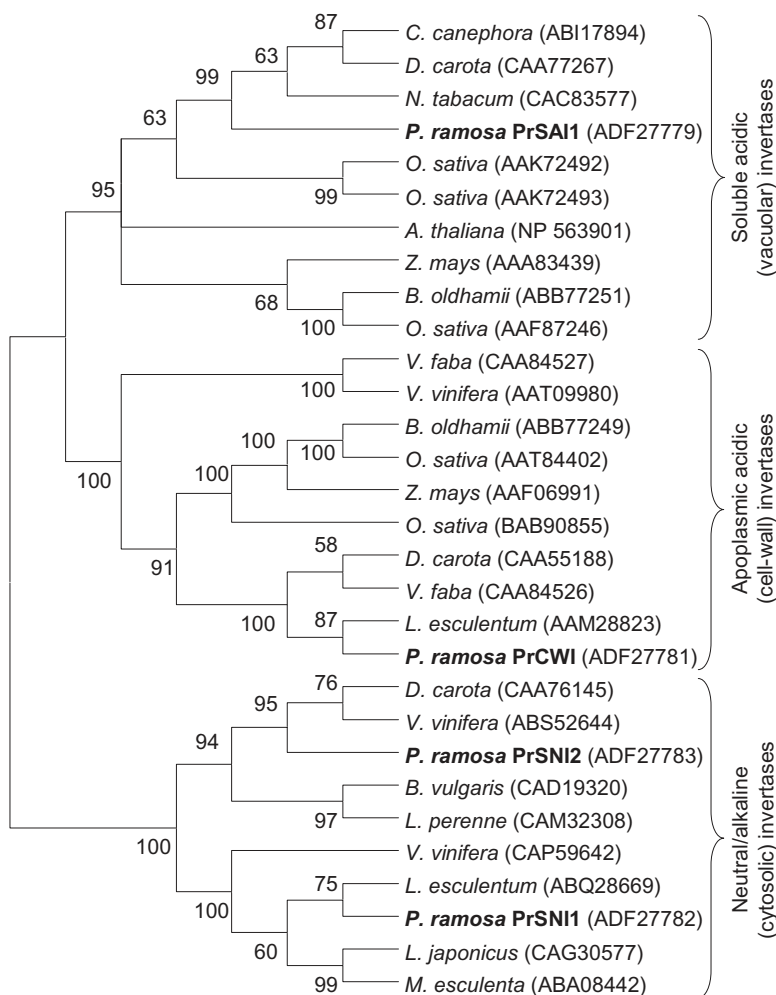


Fig. 1 Dendrogram of invertase amino acid sequences from various plants including sequences deduced from *Phelipanche ramosa* full-length invertase genes. The neighbour-joining consensus tree was inferred from 500 bootstrap replicates. Branches occurring in less than 50% bootstrap replicates were collapsed.

shown in Fig. 1, the *PrSai1*-encoded protein has a closer evolutionary relationship to vacuolar invertases (SAIs) than to apoplasmic isoforms, whereas the protein encoded by *PrCwi* belongs to the CWI family. The partial *PrSai2*-encoded

sequence was also closely related to the SAI family. Three features in the three *P. ramosa* invertase-encoded sequences also supported this classification: (i) the fourth residues of the conserved catalytic domain WEC(VII/P)D in *PrSai1*- and

PrSai2-encoded proteins were isoleucine and valine, respectively, present in vacuolar invertases; the corresponding amino acid residue in the *PrCwi*-encoded protein was proline, a characteristic feature of CWI (Roitsch *et al.*, 1995); (ii) the calculated *pI* values of the *PrSai1*- and *PrCwi*-encoded proteins were 5.42 and 8.68, respectively (Table 1), in accordance with previous observations indicating that vacuolar invertases display acidic *pI* values and CWIs show basic *pI* values (Roitsch and Gonzalez, 2004; Sturm, 1999); (iii) the TargetP 1.1 program strongly predicted that the *PrCwi*-encoded protein is exported to the apoplast, in accordance with the assumed apoplastic location of *PrCwi*.

In addition, two distinct cDNAs encoding putative neutral/alkaline invertase isoforms were isolated. *PrSni1*- and *PrSni2*-encoded proteins can be classified as neutral invertases. The deduced amino acid sequences were indeed highly homologous to neutral/alkaline invertases from other plants (Table 1). The full-length sequence encoded by *PrSni1* shows 80.9% identity with a *Solanum lycopersicum* invertase (ABQ28669) and 78.6% identity with a *Vitis vinifera* putative neutral invertase (CAP59642). The full-length sequence encoded by *PrSni2* shares 69% identity with a *V. vinifera* neutral invertase (ABS52644) and 67.9% identity with a *D. carota* neutral invertase (CAA76145) (Sturm *et al.*, 1999). As shown in Fig. 1, *PrSni1*- and *PrSni2*-encoded proteins have a closer evolutionary relationship to neutral/alkaline (cytoplasmic) invertases than to acidic (apoplastic or vacuolar) invertases. In addition, the *pI* values of *PrSni1*- and *PrSni2*-encoded proteins were calculated to be 6.72 and 7.16, respectively (Table 1), in agreement with previous observations that neutral invertases have neutral *pI* values (Roitsch and Gonzalez, 2004).

Development-related changes in the transcript levels of invertase-encoding genes

The accumulation of transcripts encoding different putative invertases was expressed relative to the level of transcripts of the constitutive elongation factor 1- α gene (*EF1 α*) (Fig. 2a,c). Regardless of the invertase gene assayed, preconditioned seeds displayed low transcript accumulation (Fig. 2a). Transcript accumulation remained low in germinated seeds for *PrSai2*, *PrSni1*, *PrSni2* and *PrCwi* genes. In contrast, large amounts of *PrSai1* transcripts were found in germinating seeds developing a radicle. Similarly, whatever the organ or stage of development after attachment to host roots, transcript accumulation of *PrSai2*, *PrSni1*, *PrSni2* and *PrCwi* was low in comparison with the high *PrSai1* transcript levels (Fig. 2c). *PrSni1* showed the lowest transcript levels. Moreover, the results showed that *PrSai1* transcripts accumulated particularly in growing subterranean shoots and emerged flowering shoots (apices, FS.V). The base of flowering shoots (Bp.V) displayed significantly lower levels of *PrSai1* tran-

scripts. *PrSai1* transcripts also accumulated substantially in fruits containing developing seeds (F.V).

Development-related changes in invertase activities and *in situ* localization of SAI activity

Activities were measured in preconditioned and germinated seeds and in different organs of the subterranean and emerged broomrape (Fig. 2b,d). SAI, SNI and CWI activities were low in preconditioned seeds, and increased following germination by factors of 14-, 1.5- and five-fold, respectively (Fig. 2b). Once the parasite had attached to host roots, levels of CWI, SNI and SAI activities in tubercles were similar and did not change significantly during overall plant development (Fig. 2d). In contrast, enzyme activity increased substantially in both subterranean shoots (S.IV) and at the apices of emerged flowering shoots (FS.V), primarily as a result of a large increase in SAI activity (Fig. 2d). Fruits containing developing seeds (F.V) displayed the highest SAI activity, whereas the base of emerged flowering shoots (Bp.V) showed low SAI activity. *In situ* AI activities were visualized in cross-sections of basal and apical parts of emerged flowering shoots (Fig. 3). Activity was confined to the medullar parenchyma in basal parts, but extended to the cortical parenchyma in the apices (Fig. 3a2,b2). Staining on fixed sections was attributed to SAI (Sergeeva and Vreugdenhil, 2002), because no CWI activity was revealed on unfixed sections (data not shown). The higher SAI activity observed in the apices of emerged shoots may be related to the more specific localization of SAI activity in these young, growing tissues (Figs 2d and 3b2). SAI staining from germinated seeds also showed high SAI activity in the radicle apex (Fig. 3d).

Figure 2 highlights the strong similarity between the patterns of *PrSai1* expression and SAI activity. Figure S1 (see Supporting Information) illustrates the strong relationship between SAI activity (X) and *PrSai1* transcript levels ($Y = 168.14X$; $R^2 = 0.78$).

SAI purification, characterization and sequencing

SAI proteins were purified from growing subterranean shoots (S.IV) and partially sequenced following trypsin digestion to verify that the *PrSai1* gene encodes the most active SAI isoform. The latter was highly purified by a factor of 700-fold with 2% of activity recovery (Table S3, see Supporting Information). The purified SAI displayed an apparent molecular mass of 84 kDa in native conditions and 35 kDa in denaturing conditions (Table 2). The identity of purified SAI was demonstrated on native gels by both activity staining and immunoblotting (Fig. S2, see Supporting Information). As shown by liquid isoelectric focusing experiments and enzyme assays in the pH range 2.0–8.0, purified SAI displayed the expected *pI* and optimal acidic pH (4.63 and 4.5, respectively) (Table 2). When assayed at pH 4.5 and at temperatures ranging

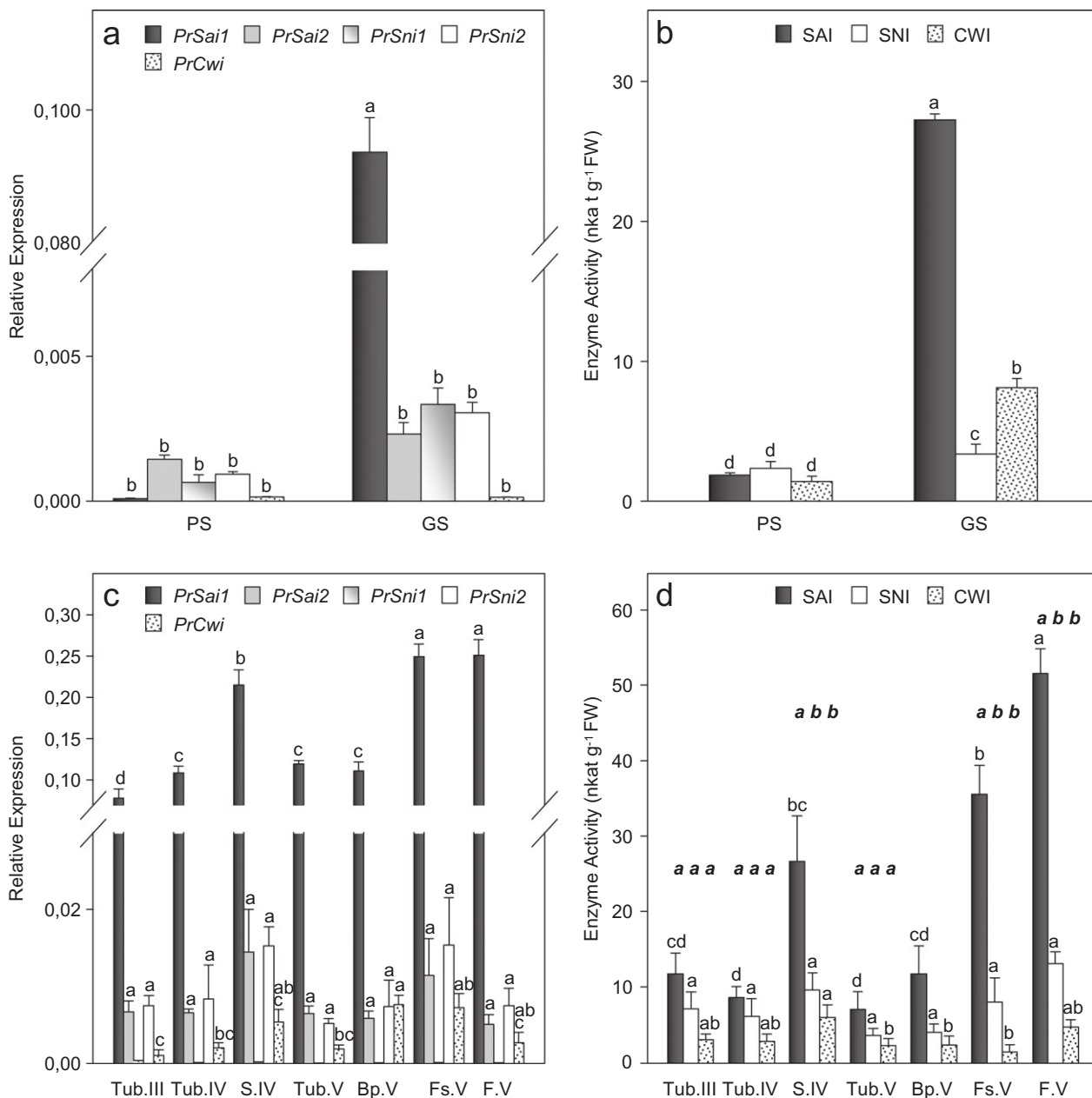


Fig. 2 Development-related changes in the levels of invertase-encoded gene transcripts and invertase activities in *Phelipanche ramosa*: (a, b) independent development following seed germination; PS, preconditioned seeds; GS, germinated seeds; (c, d) parasitic stages following attachment to tomato roots. See Fig. 5 for abbreviations of the various plant organs. (a, c) Transcript accumulation expressed relative to elongation factor 1- α gene (*EF1 α*) transcript levels. Data are means \pm standard error ($n = 3$). For each gene, values with the same letter are not significantly different (ANOVA, SNK test, $P < 0.05$). (b, d) Invertase activities. Data are means \pm standard error ($n = 6$). For each isoform, activities with the same letter (plain typeface) are not significantly different (ANOVA, SNK test, $P < 0.05$). For each stage of development and organ, activities with the same letter (italic and bold typeface) are not significantly different (ANOVA, SNK test, $P < 0.05$).

from 20 to 80 °C, activity was optimal at 50 °C. Purified SAI was active on stachyose, raffinose and, especially, sucrose, where it showed very high activity and affinity. No activity was detected with any other carbohydrates tested as substrate. Fructose inhibited SAI activity in a linear competitive manner ($K_i = 1.63$ mM),

whereas glucose acted as a linear noncompetitive inhibitor ($K_i = 12$ mM). SAI characterization was carried out by testing mineral ions as potential inhibitors (Table 2). Partial inhibition was shown with Ag^+ , Cu^{2+} , K^+ , Na^+ , Mg^{2+} and Pb^{2+} ions, and SAI activity was affected strongly by Hg^{2+} .

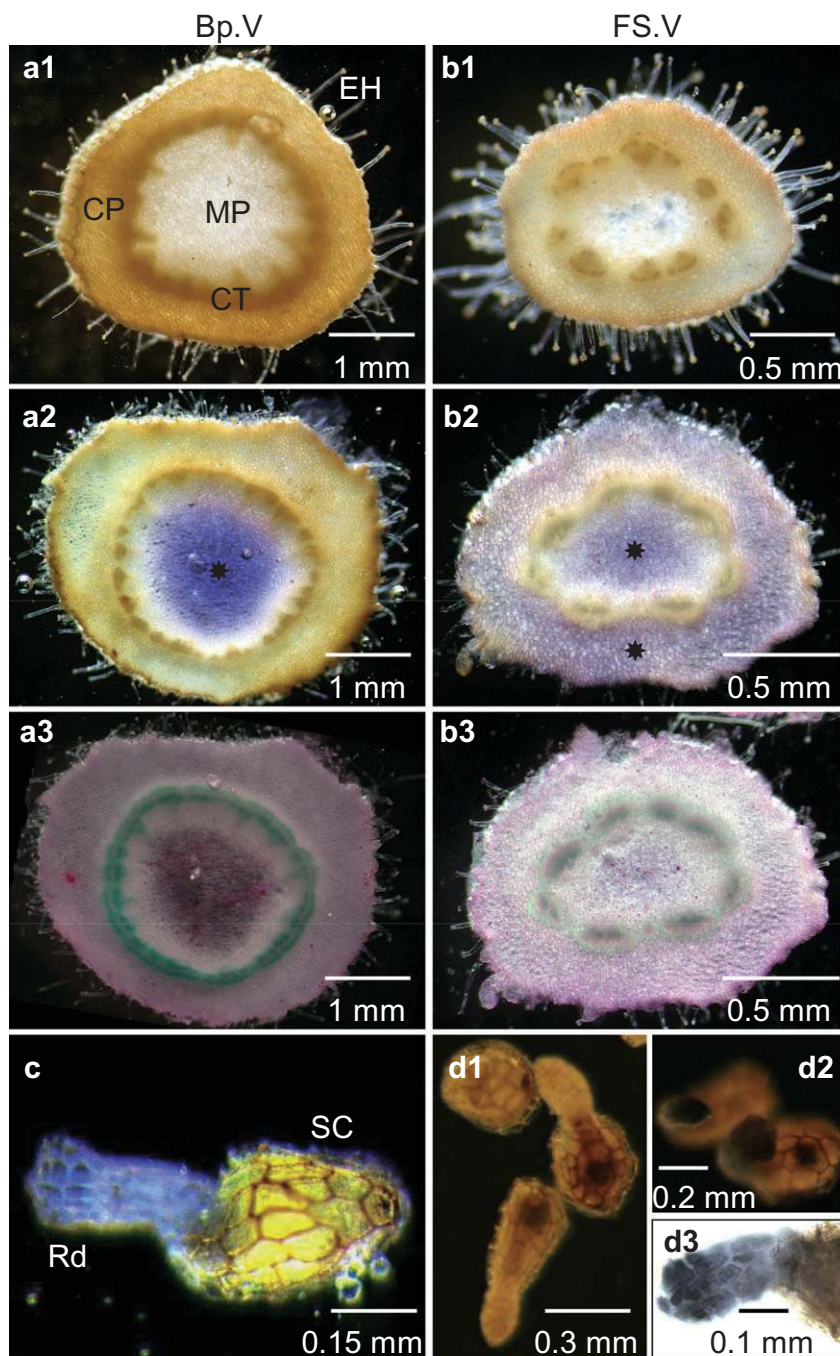


Fig. 3 Tissue localization of soluble acid invertase (SAI) activity in seeds and flowering shoots of *Phelipanche ramosa* parasitizing tomato plants. (a1, b1) Control cross-sections of basal (Fp.V) and apical (FS.V) shoots incubated after fixation in reaction medium without sucrose. (a2, b2) Assay sections incubated after fixation in a reaction mixture containing sucrose. SAI activity (★) is shown in blue in medullar and cortical parenchyma. No cell wall invertase (CWI) activity was detected on unfixed sections (data not shown). Staining on fixed sections was thus attributed to SAI. (a3, b3) Post-activity staining of cellulosic (purple) and lignified (green) tissues using Mirande's reagent. (c) Germinated seeds incubated in the reaction mixture containing sucrose. (d1) NTB staining of seeds without sucrose in reaction medium. (d2) NTB staining of seeds with sucrose in reaction medium. (d3) Zoom of the radicle region stained with NTB: accumulation of reducing sugars produced from SAI-mediated hydrolysis of sucrose is shown in black in the apex of the radicle. CP, cortical parenchyma; CT, conductive tissues; EH, epidermal hairs; MP, medullar parenchyma; NTB, nitroterazolium blue; Rd, radicle; SC, seed coat.

The purified SAI protein was extracted from the band showing both SAI staining and immunoblotting on native gels, and then sequenced using electrospray ionization-liquid chromatography-tandem mass spectrometry (ESI-LC-MS/MS). Trypsin digestion generated 14 peptides, which were aligned on the deduced peptide sequence of the *PrSai1* gene (Fig. 4). These peptides cover 26% of the deduced sequence, attesting that *PrSai1* encodes for the SAI purified in the present study.

DISCUSSION

The present study was designed to both identify and characterize the invertase genes and isoforms that determine sucrose utilization during broomrape development, using *P. ramosa* as a model. We covered the major steps of parasite development, starting from the independent germinating seedling to the ripening parasitic plant after attachment to host roots and emergence from the soil.

Table 2 Characteristics of the soluble acid invertase PrSAI1 purified from growing subterranean shoots of *Phelipanche ramosa* parasitizing tomato plants.

Kinetic properties	
<i>pI</i>	4.6 ± 0.1
Optimal pH	4.5 ± 0.1
Optimal temperature (°C)	50 ± 3
Molecular mass (kDa)	
Native form	86 ± 4
Denatured form	35 ± 2
Substrate specificity (mM)	
<i>K_m</i> (sucrose)	2.7 ± 0.3
<i>K_m</i> (stachyose)	16.9 ± 2.4
<i>K_m</i> (raffinose)	75.6 ± 5.5
<i>K_m</i> (cellobiose, lactose, maltose, melezitose, melibiose, methyl-glucose, trehalose, verbascose)	—
Inhibition	
<i>K_i</i> (fructose) (mM)	1.6 ± 0.1
<i>K_i</i> (glucose) (mM)	12.0 ± 0.4
MnCl ₂ , CoCl ₂ (1 mM, % control)	110 ± 3.1
CaCl ₂ (1 mM, % control)	86 ± 6.1
AgNO ₃ , KCl, NaCl, MgCl ₂ (1 mM, % control)	70.5 ± 2.4
CuSO ₄ , ZnSO ₄ , PbNO ₃ (1 mM, % control)	48.7 ± 7.3
HgCl ₂ (1 mM, % control)	9 ± 2.0

Values are means ± standard error (*n* = 3).

The invertase gene family in *P. ramosa*

Because information on the broomrape genome was scarce when the present study started, we set out to isolate cDNAs encoding invertases from *P. ramosa* using traditional degenerate primers corresponding to highly conserved regions of plant SAI, CWI and SNI genes (Tables S1 and S2, see Supporting Information). The first group of sequences characterized encoded AIs, including three cDNAs, *PrSai1*, *PrSai2* and *PrCwi*. Based on their close evolutionary relationship to vacuolar invertases or CWIs (Fig. 1), the identity of the fourth residue of the conserved catalytic domain (WEC(VII/PD) of plant invertases (Roitsch *et al.*, 1995), the *pI* value (Roitsch and Gonzalez, 2004; Sturm, 1999) and the predicted apoplastic location, the cDNAs *PrSai1* and *PrSai2* encode vacuolar isoforms, whereas the *PrCwi*-encoded protein belongs to the CWI isoform. The second group of sequences isolated encoded soluble neutral/alkaline invertases, including two cDNAs, *PrSni1* and *PrSni2*. In parallel with our study, the Parasitic Plant Genome Project (PPGP) sequenced cDNAs representing transcripts from key life stages of three members of the Orobanchaceae, including *P. aegyptiaca* (syn. *Orobanche aegyptiaca*), a close relative of *P. ramosa* (<http://ppgp.huck.psu.edu>). The *PrSai1*, *PrCwi*, *PrSni1* and *PrSni2* genes share 96.7%, 86.7%, 97.9% and 95.5% identity with four partial sequences encoding putative invertases in *P. aegyptiaca*: *OrAeBC3-2697*, *OrAeBC3-21925*, *OrAeBC3-35083* and *OrAeBC3-36350*, respectively. Other CWI- and SNI-encoding cDNAs exhibiting lower identity could be identified in *P. aegyptiaca*, suggesting the occurrence of other invertase genes in *P.*

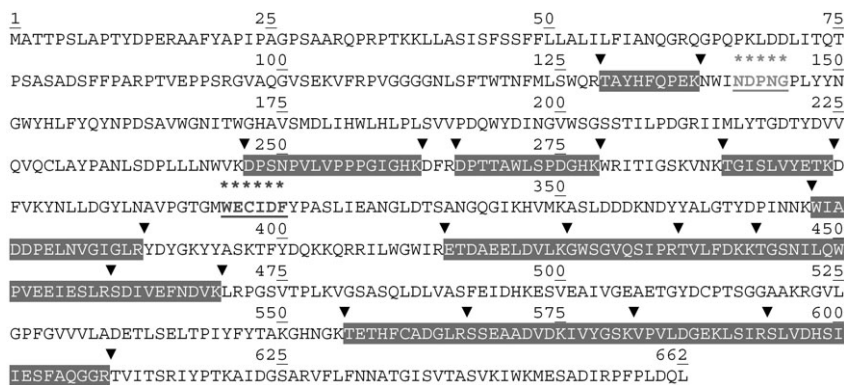
ramosa. Furthermore, the sequence *OrAeBC3-2783* encoding the key enzyme in mannitol production, mannose 6-phosphate reductase (EC 1.1.1.224), shares 99.3% identity with the *PrM6PR* sequence from *P. ramosa* (Aly *et al.*, 2009; Delavault *et al.*, 2002). These data highlight the strong identity of nucleic acid sequences from both species, and the PGP database will clearly facilitate future studies on *P. ramosa* and probably other broomrape species as well.

SAI isoform expression in *P. ramosa* is simple, with a typical dominant *PrSai1*-encoded isoform

Multiple SAI isoforms have been reported in many species. For example, three of five SAI isoforms suspected in mature leaves of *Arabidopsis thaliana* have been characterized and their properties compared (Tang *et al.*, 1996). They exhibited similar *pI*, pH and temperature optima and molecular mass values. Further, sucrose is their preferred substrate with a *K_m* for sucrose varying from 5 to 12 mM, according to the isoform. To date, only two genes, *Atβfruct3* and *Atβfruct4*, encoding vacuolar invertases have been identified in *A. thaliana* (Haouazine-Takvorian *et al.*, 1997). Heterogeneity in SAI isoforms is better understood in other species and involves the expression of different *Sai* genes (e.g. carrot; Unger *et al.*, 1994) or post-translational modification of a single gene product (e.g. radish, tomato; Elliott *et al.*, 1993; Faye *et al.*, 1986). As mentioned above, *P. ramosa* exhibits two genes, *PrSai1* and *PrSai2*, that encode putative vacuolar isoforms (Table 1). Nevertheless, the pattern of expression of SAI isoforms is simple in growing shoots: only one SAI isoform was detected. This finding suggests that the second SAI isoform is not active in shoots, or only has a low basal activity that could not be detected on purification. In addition, patterns of *PrSai1* and *PrSai2* gene expression indicate that *PrSai1* transcripts accumulate much more strongly than do *PrSai2* transcripts (Fig. 2a,c). The purified SAI isoform shares high identity with the deduced peptide sequence of the highly expressed *PrSai1* gene (Fig. 4). We thus assume that it corresponds to the *PrSai1*-encoded protein, therefore called 'PrSAI1'. Furthermore, *PrSai1* transcript levels and SAI activity correlated significantly with overall parasite development (Fig. S1), indicating that SAI activity can be attributed to PrSAI1 in all organs.

PrSAI1 was visualized in native conditions by activity staining on polyacrylamide gels or by immunoblotting using an antibody raised against grape vacuolar invertase (Fig. S1). The molecular mass of the *PrSai1*-deduced protein (73 kDa) is lower than that of the native PrSAI1 (86 kDa, Table 2), which is in agreement with the fact that the native form is glycosylated. This finding also demonstrates that PrSAI1 is monomeric. Accordingly, multiple SAI forms varying from 30 to 220 kDa have been detected in plants, but monomeric or dimeric forms of 50–70 kDa predominate (Sturm, 1999).

Fig. 4 Amino acid sequence as deduced from the *PrSai1* gene and amino acid sequences of the peptides generated by trypsin digestion of the soluble acid invertase (SAI) isoform purified from the subterranean shoots of *Phelipanche ramosa* parasitizing tomato plants. Peptides matching the putative *P. ramosa* acid invertase sequence are shaded, and tryptic cleavage sites are indicated by arrows. The β -fructofuranosidase motif and a highly conserved peptide domain with a catalytic cysteine residue are underlined (Sturm, 1999).



PrSAI1 also exhibited kinetic features typical of many other SAIs. Of all the carbohydrates tested as potential substrates, PrSAI1 was active only on β -fructofuranosides, including sucrose, stachyose and raffinose (Table 2), and was competitively inhibited by fructose, confirming that PrSAI1 is a β -fructofuranosidase, as generally reported for invertases (Sturm, 1999). PrSAI1 exhibited a K_m value for sucrose in a lower millimolar range than for raffinose and stachyose, demonstrating that sucrose is the preferred substrate. The optimal temperature of PrSAI1 was high (50 °C), as demonstrated for other plant SAIs (Marouf and Zeki, 1982; Miller and Ranwala, 1994). PrSAI1 activity was inhibited more or less strongly by metallic ions (Table 2). As generally observed in plant SAIs, PrSAI1 exhibited high sensitivity to Hg^{2+} , suggesting the presence of a sulphhydryl group at the catalytic site of the enzyme (Sturm, 1999).

Sucrose utilization is mediated mainly by PrSAI1 during both host-independent and post-attachment development

In the work presented here, the expression of the five genes encoding invertases and the activities of SAI, SNI and CWI isoforms were followed in both preconditioned and germinated seeds of *P. ramosa* (Fig. 2a,b). Although the transcript levels and activities were low in preconditioned seeds, both *PrSai1* transcript levels and putative PrSAI1 activities increased greatly in seedlings after germination. The gene *PrSai1* was by far the most expressed invertase-encoding gene in germinated seeds, where PrSAI1 activity also predominated, suggesting that sucrose degradation for radicle growth during the mobilization of seed reserves is controlled mainly by PrSAI1. This was confirmed by the high PrSAI1 activity detected at the radicle apex (Fig. 3d3). Considering that broomrape seeds store essentially lipids as reserves, starch being very uncommon (Bar-Nun and Mayer, 2002; Velasco *et al.*, 2000), additional investigations on the mobilization of the lipid reserves in broomrape seeds are recommended. Most of the germination stimulants identified so far are

strigolactones (Yoneyama *et al.*, 2010). Recent studies on arbuscular mycorrhizal fungi have shown that strigolactones stimulate mitochondrial metabolism and catabolic processes, such as β -oxidation of fatty acid degradation (Besserer *et al.*, 2008; Bücking *et al.*, 2008). Accordingly, we speculate that lipids are converted into sucrose, followed by PrSAI1-mediated sucrose utilization, in young *P. ramosa* seedlings following germination. However, this does not preclude PrSAI1 involvement in gentianose hydrolysis, which is stimulated in germinating broomrape seeds (e.g. A. Okazawa, Graduate School of Engineering, Osaka University, 2-1 Yamadaoka, Suita, Osaka, Japan, unpublished data).

Voegelé *et al.* (2006) have reported that, on penetration into the host faba bean leaf, the rust fungus *Uromyces fabae* establishes new sinks which compete with other occurring sink organs through strong expression of the *Uf-INV1* gene encoding an apoplastic invertase in haustoria. Thus, the fungal invertase activity might limit sucrose export from the infected leaf and might condition the tissue for a conversion from source to sink. Such alterations in translocation patterns are well documented (Whipps and Lewis, 1981). Interestingly, the PrCWI activity increased by five-fold in broomrape germinated seeds in comparison with preconditioned seeds, suggesting that the parasite CWI could influence the sink strength of the infected roots during the subsequent step of host root penetration. Additional studies should be performed to assess the role of PrCWI in host root phloem unloading during broomrape attachment. In contrast, the expression of the studied *PrCwi* gene was low in both preconditioned and germinated seeds. At least three cDNAs encoding putative CWI proteins could be detected in the close relative *P. aegyptiaca* (*OrAe41G2B1-99085*, *OrAe3GB1-12647* and *OrAe3GB1-11068* from PGPP), sharing 56%, 54% and 60% identity, respectively, with the *PrCwi* sequence. We can thus hypothesize that at least another cell wall-encoding gene in addition to *PrCwi* exists in *P. ramosa*. Nevertheless, the existence of regulatory mechanisms other than transcriptional regulation cannot be excluded to explain the gain in CWI activity in germinated seeds.

Once the radicle reaches the host roots and establishes a functional haustorium, the attached broomrape first develops a tubercle and then an achlorophyllous shoot. Sucrose uptake from the host leads to hexose accumulation, mainly in the growing shoot, during both subterranean and aerial development (Abbes *et al.*, 2009; Delavault *et al.*, 2002), suggesting that invertases also play a central role during overall post-attachment development. In the present study, we followed the expression of the five genes encoding invertases and the activities of SAI, SNI and CWI in tubercles, shoots and fruits containing developing seeds of *P. ramosa* parasitizing tomato plants (Fig. 2c,d). Of the five transcripts, *PrSai1* transcript levels predominated in all organs regardless of the developmental stage. Nonetheless, the highest levels of *PrSai1* transcripts and of putative PrSAI1 activities occurred in growing organs, including the subterranean shoot, the apex of emerged flowering shoots and fruits containing developing seeds. In old tissues, such as the basal parts of emerged flowering shoots, levels of *PrSai1* transcripts and PrSAI1 activity were significantly lower. Moreover, in these older plant parts, PrSAI1 activity was confined to the medullar parenchyma. In contrast, PrSAI1 was located in both cortical and medullar parenchyma in apices of flowering shoots (Fig. 3b2), as well as in the whole subterranean shoot (data not shown). These findings indicate that PrSAI1 controls major processes in parasite development, including the sink strength for sucrose in these young tissues by reducing water potential through the massive accumulation of hexoses in the vacuole, cell expansion in elongating zones by maintaining turgor through hexose accumulation and adjustment of the hexose to sucrose ratio. As sucrose cleavage by vacuolar PrSAI1 should be the primary sucrose utilization pathway in these cells, PrSAI1 is assumed to actively drive phloem unloading of sucrose in the growing sinks. Sucrose hydrolysis mediated by an SAI isoform has also been demonstrated to be efficient in phloem unloading, hexose accumulation and the regulation of cell turgor and expansion in carrot roots, bean stems and tomato fruits (Klann *et al.*, 1996; Morris and Arthur, 1985; Ohyama *et al.*, 1995; Ricardo and Ap Rees, 1970).

In contrast, *PrSni1*- and *PrSni2*-encoding isoforms display low activity (Fig. 2), as do SNI isoforms in carrot roots (Ricardo and Ap Rees, 1970) and sugar beet (Masuda *et al.*, 1987). Nevertheless, a role of SNI in sucrose utilization cannot be excluded in *P. ramosa*. Indeed, it is commonly assumed that SNI substitutes for SuSy and provides starch and cell wall precursors when SuSy is deficient (Barratt *et al.*, 2009). The study of these activities at the cellular or tissue level is needed to understand their respective roles in sucrose utilization in *P. ramosa*.

Control of steady-state PrSAI1 activity

High sucrose levels and invertase can co-occur in plant vacuoles. Post-translational inhibition of vacuolar invertases can result

from inhibition by hexose products, proteinaceous inhibitors, structural modification and hexose-mediated modulation of expression of some invertase inhibition-related genes (Privat *et al.*, 2008; Rausch and Greiner, 2004). Searching the *O. aegyptiaca* sequences from PPGP with known proteinaceous invertase inhibitor sequences leads to hits for putative invertase inhibitors in the Orobanchaceae. Nevertheless, inhibition of PrSAI1 is unlikely in a parasite which contains only negligible amounts of sucrose (Delavault *et al.*, 2002). Furthermore, the extractable PrSAI1 activity was shown to be typically modulated by glucose as a noncompetitive inhibitor and fructose as a competitive inhibitor (Table 2), indicating that PrSAI1 inhibition by hexoses was overcome in cells, resulting in massive accumulation of hexoses in the vacuole, as observed in growing shoots (Delavault *et al.*, 2002).

Steady-state levels of plant invertases are commonly controlled by a dominant transcriptional regulation mechanism. Likewise, patterns of PrSAI1 transcript levels and activity were highly correlated (Fig. S1). As suggested in germinating legume seeds, which show increased CWI activity in response to cytokinins, as well as the induction of D-type cyclin-encoding genes in response to hexose products (Roitsch and Gonzalez, 2004), cytokinins and hexoses may enhance, in a cooperative manner, cell proliferation and sink metabolism in growing organs of *P. ramosa*. Interestingly, high cytokinin levels have been reported in xylem-feeding parasitic plants (Jiang *et al.*, 2005; Lechowski and Bialczyk, 1996); nevertheless, no data are available for phloem-feeding parasitic plants such as *P. ramosa*.

In addition, PrSAI1 may also be controlled by a post-translational regulatory mechanism involving proteolytic cleavage. We suggest this mechanism as a result of differences in the molecular mass predicted from the *Prsai1* gene (73-kDa polypeptide) and that estimated from PrSAI1 (86 kDa and 35 kDa in native and denaturing conditions, respectively) (Table 2). PrSAI1 in subterranean shoots probably corresponds to two peptides of about 35 kDa produced by proteolytic cleavage. This type of post-translational mechanism has been described for the monomeric 68-kDa and 70-kDa vacuolar forms in mung bean and carrot, respectively (Arai *et al.*, 1991; Unger *et al.*, 1994), in which the fragments are tightly associated under native conditions, and the complex is active. Disassociation appears to be under developmental control in both species, as the ratio between fragments and full-length protein increases with plant age. Developmental control of PrSAI1 in *P. ramosa* was not studied here. In addition, the physiological function of fragmentation remains unclear in plants.

The findings presented in this work indicate that, among the studied invertases, vacuolar PrSAI1 dominates in conditioning the sink strength and growth during both host-independent and post-attachment phases. However, additional studies are needed to confirm that PrCWI acts on the sink strength of infected roots

during root penetration. Consequently, inhibition of PrSA11 and possibly PrCWI activities may be a good strategy for curbing parasite development. Given that the phloem mobility of proteins and nucleic acids between host and parasite has been demonstrated (Aly *et al.*, 2006; Davis and Wurdack, 2004), our results call for additional studies to evaluate the efficiency of new transgenic resistance in host plants based on either the production of proteinaceous inhibitors specifically targeted against PrSA11 (or PrCWI) or the *PrSai1* (or *PrCwi*) silencing technology described by Aly *et al.* (2009) and Tomilov *et al.* (2008). The efficiency of the silencing strategy has already been demonstrated: a silencing mechanism targeted against the parasite's mannose 6-phosphate reductase-encoding gene causes *P. aegyptiaca* to lose accumulated solutes and, as a result, a significant increase in parasite mortality (Aly *et al.*, 2009). Previous studies have shown that the mannose 6-phosphate reductase protein is indeed a key enzyme in mannitol production in broomrape species (Delavault *et al.*, 2002; Harloff and Wegmann, 1993). Moreover, because no transformation method is available for *P. ramosa*, the silencing strategy through host genetic modification could be useful to confirm the key role of PrSA11 in parasite sink strength and growth, and to analyse the relative contribution of the slightly expressed genes *PrCwi* and *PrSni*. To obtain a complete picture of the sink strategy in broomrape species, more investigations are needed on the mechanisms of sucrose unloading and sucrose allocation between mannitol, hexoses and starch in sink cells, and the involvement of other classes of sucrose-degrading enzymes, e.g. SuSy.

EXPERIMENTAL PROCEDURES

Plant materials, cultivation and sampling

Seeds were collected from flowering spikes of *P. ramosa* L. Pomel in France (at Saint Jean d'Angely, 2005) and stored in darkness at 25 °C until use. They were mixed homogeneously with a 1:1:1 peat–sand–clay mixture in 3-L pots (10 mg/L of soil); the pots were watered and protected from light with a black plastic film for 1 week at 20–25 °C (day–night temperature). Then, one tomato seed (*Solanum lycopersicum* L., var. Robusta, Seminis) was sown directly into each pot. The plants were sprinkled weekly with a sterile solution of 50% Coïc nutrient solution (Labrousse *et al.*, 2001) and grown under a photoperiod of 16 h (photosynthetic fluence rate of 300 $\mu\text{mol}/\text{m}^2/\text{s}$ over a waveband of 400–700 nm) at a temperature of 20–25 °C. Twelve weeks after sowing, tomato plants were gently uprooted from the soil and the different developmental stages of *P. ramosa* seedlings were harvested (Fig. 5). Fruits containing developing seeds were also harvested from flowering spikes (stage V).

Phelipanche ramosa seeds were surface sterilized as described previously by Labrousse *et al.* (2001). They were pre-

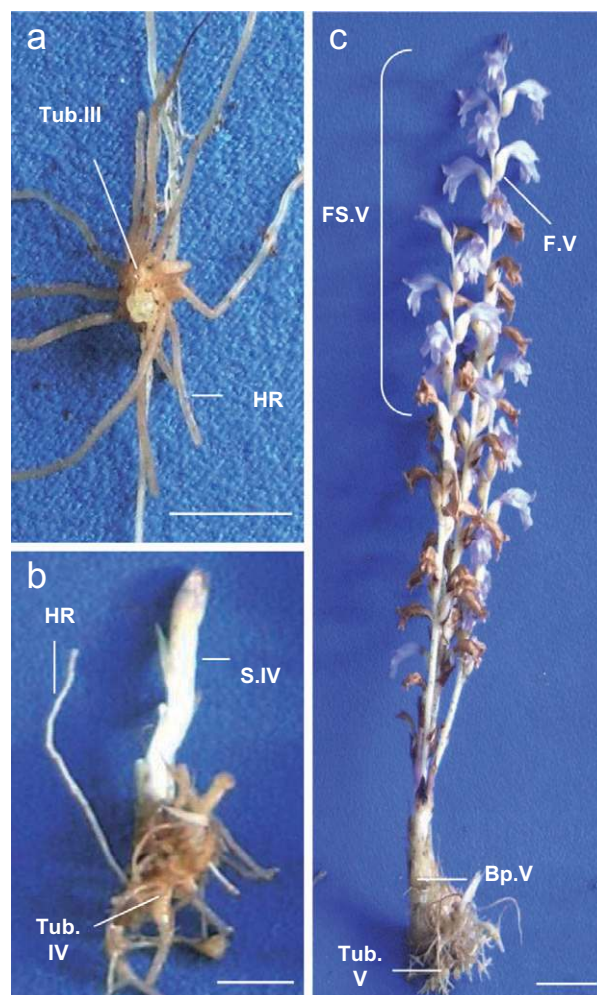


Fig. 5 Developmental stages in *Phelipanche ramosa* parasitizing tomato plants: (a) stage III, growing tubercle (Tub.III); (b) stage IV, tubercle (Tub.IV) bearing the growing subtterranean shoot (S.IV); (c) stage V, tubercle (Tub.V) bearing the emerged flowering shoot. Apical part of the flowering shoot is growing (FS.V) and bears fruits containing developing seeds (F.V). Basal part (Bp.V) does not bear flowers and is larger and more fibrous than the flowering shoot. HR, host roots (tomato). Bar, 1 cm.

conditioned on sterile, watered, glass-fibre filters placed in 9-cm-diameter Petri dishes after a 6-day period in darkness at 21 °C. Germination of a preconditioned seed set was induced following an additional 6-day period in darkness in the presence of GR24 (1 $\mu\text{g}/\text{L}$; Johnson *et al.*, 1976).

Seeds and plant organs were immediately frozen in liquid nitrogen and stored at -80 °C prior to RNA and enzyme extraction.

Total RNA extraction and cDNA preparation

Frozen tissues were ground in liquid nitrogen and total RNA was extracted with the RNeasy Plant Mini Kit (Qiagen, Courtaboeuf,

France) according to the manufacturer's instructions. Extracts were treated with DNase I enzyme (0.02 U/ μ L; New England Biolabs, Ipswich, MA, USA) to eliminate residual genomic DNA. The integrity of total RNA was determined by electrophoresis on 1% (w/v) agarose gel. Using oligo(dT20) as a primer, cDNA was prepared from samples (0.5 μ g) of total RNA using the Super-script II reverse transcriptase kit (Invitrogen, Carlsbad, CA, USA).

Isolation of cDNAs from the invertase family in *P. ramosa*

The cDNAs isolated from an emerged parasite (stage V) were used for PCR amplification. Degenerate primers corresponding to highly conserved regions of AIs (SAI and CWI) and neutral invertases (cytosolic, SNI) were designed (Table S1). After denaturation at 94 °C for 5 min, the amplification consisted of 35 cycles of 45 s at 94 °C, 45 s at 52, 54 or 57 °C for *PrSai* (1 and 2), *PrCwi* and *PrSni* (1 and 2), respectively, and 1 min 30 s at 72 °C. An additional final step of elongation was performed at 72 °C for 5 min. The amplified DNA fragment was purified and cloned into a pGEM-T Easy vector (Promega, Charbonnières-Les-Bains, France). Recombinant plasmid DNA was prepared and then sequenced by GATC Biotech (Konstanz, Germany). Based on the initial sequence information, new primers were generated for RACE of each fragment using the Generacer kit (Invitrogen, Eragny Sur Joise, France). RACE products corresponding to different invertase genes were cloned and sequenced. An incomplete 5' RACE product was amplified for *PrSai2*. To amplify full-length *PrSai1*, *PrCwi*, *PrSni1* and *PrSni2* cDNAs, specific primer pairs were designed. After denaturation at 94 °C for 5 min, the amplification consisted of 35 cycles of 1 min at 94 °C, 1 min at 58, 60 or 54 °C for *PrSai1*, *PrCwi* and *PrSni* (1 and 2), respectively, and 2 min 30 s at 72 °C. An additional final step of elongation was performed at 72 °C for 7 min. The amplified DNA fragments were purified and cloned as mentioned above.

Real-time quantitative PCR

The determined sequences were used to design gene-specific primers for real-time quantitative PCR. Six primer pairs were designed (maximum length, 150 bp; optimal melting temperature T_m at 60 °C; GC percentage between 30% and 80%) with Primer Express 3.0 software (Applied Biosystems, Foster City, CA, USA). The selected primers underwent an extensive search using the BLAST tool to avoid any significant homology with other known nucleotide sequences.

Quantitative PCR was carried out on an Applied Biosystems 7300 real-time PCR system using SYBR Green technology. Each reaction contained 300 nM of each primer, 5 μ L of cDNA sample (~5 ng of input RNA) and 2 \times Power SYBR Green PCR master mix (Applied Biosystems) in a total volume of 25 μ L. Thermal cycling

was performed using the following cycling conditions: 95 °C for 10 min, 40 cycles at 95 °C for 15 s and 60 °C for 1 min. All quantitative PCR runs contained negative controls with no cDNA template, so as to exclude or detect possible contamination. The specificity of the PCR amplification was checked with a heat dissociation protocol (from 60 to 95 °C) following the final cycle of PCR. The efficiency of the primer sets was calculated by performing quantitative PCR on several dilutions of the strands generated in the first cycle. The efficiency of the different primer sets used was checked and found to be similar.

Results were standardized to *PrEF1 α 1* (accession number: HM219554) gene expression levels; *PrEF1 α 1* displays no development-related changes. Experiments were carried out on three biological replicates.

Sequence analysis

Bioinformatics analyses were performed using Vector NTI 9.1.0 software (Invitrogen, Invitrogen Corporation, Carlsbad, CA, USA). Sequence homologies were verified against GenBank databases using BLAST programs (<http://www.ncbi.nlm.nih.gov/blast>). The subcellular location of the deduced proteins was predicted using the TargetP 1.1 Server (<http://www.cbs.dtu.dk/services/TargetP>). Phylogenetic analyses were conducted using MEGA4 software (Tamura *et al.*, 2007). Amino acid sequences of invertase isoenzymes were aligned using CLUSTALW.

Invertase extraction and enzyme assays

Frozen tissue (1 g fresh weight) was homogenized at 4 °C (Ultra-Turrax T25, IKA®-Labortechnik, Staufen, Germany) in 6 mL of buffer A (Table 3) and 0.5 g polyvinylpyrrolidone (PVPP). Following centrifugation (10 000 g, 15 min, 4 °C), the supernatant underwent a soluble invertase assay. The residue was rinsed with 50 mL of buffer A and the supernatant was discarded after centrifugation. CWI was extracted from the washed pellet in 3 mL of buffer A containing 1 M NaCl with gentle shaking for 24 h at 4 °C. After centrifugation, the supernatant was used for the CWI assay. Proteins were measured according to Bradford (1976).

Prior to assay, the enzyme extract (2.5 mL) was desalted on a PD-10 Sephadex column (GE Healthcare, Orsay, France), previously equilibrated with 10 mM sodium phosphate buffer (pH 5 for AIs and pH 7.5 for the neutral/alkaline invertase, SNI) containing 1 mM benzamidine and 0.1 mM phenylmethylsulphonyl fluoride (PMSF). The enzymatic reaction (1 mL) was initiated by adding sucrose to a final concentration of 100 mM in the medium containing 100 mM sodium phosphate buffer (pH 5.0 for AIs and pH 7.5 for SNI) and 1–50 μ g of proteins. The reaction mixture was incubated at 30 °C for 30 min and the reaction was stopped by the addition of 0.5 mL of 100 mM sodium phosphate

Table 3 Chromatographic steps for PrSAI1 purification from growing subterranean shoots of *Phelipanche ramosa* parasitizing tomato plants.

	Buffer	Flow rate (mL/min)	Gradient	Collected fractions (mL)
Extraction	A: 100 mM Tris-HCl (pH 7.0) containing 5 mM EDTA, 10 mM ascorbic acid, 1 mM β -mercaptoethanol, 0.1 mM PMSF, 1 mM benzamidine	—	—	—
Concanavalin-A Sepharose	B: 10 mM sodium phosphate (pH 6.5) containing 1 mM EDTA, 10 mM ascorbic acid, 1 mM DTT, 0.1 mM PMSF, 1 mM benzamidine, 1 mM CaCl_2 , 1 mM MgCl_2 , 1 mM MnCl_2 and 0.5 mM NaCl	3	0–25 M β -methyl-mannoside (60 mL)	3
High Q anion exchanger	C: 50 mM Tris-HCl (pH 8.0) containing 1 mM DTT, 0.1 mM PMSF and 1 mM benzamidine	0.5	0–0.5 M NaCl (50 mL)	2
HR Sephacryl	D: 10 mM sodium phosphate (pH 6.5) containing 5 mM EDTA, 0.1 mM PMSF, 1 mM benzamidine, 0.05% (w/v) NaN_3	0.5	—	1
Liquid isoelectric focusing	50-mL solution containing an aliquot of HR-Sephacryl fraction, 2.5% (v/v) Bio-Lyte® pH 3/10 ampholyte or Bio-Lyte® pH 3/5 ampholyte (Bio-Rad, France)	—	—	2.5

DTT, dithiothreitol; EDTA, ethylenediaminetetraacetic acid; PMSF, phenylmethylsulphonyl.

buffer (pH 7.5) and heating at 95 °C for 3 min. After cooling, the reaction mixture was centrifuged and 0.5 mL of reagent, consisting of 1.5 mM NADP, 1.5 mM ATP, 2 mM MgCl_2 and 1 U of both hexokinase (EC 2.7.1.1) and glucose 6-phosphate dehydrogenase (EC 1.1.1.49) (Sigma, Saint-Quentin Fallavier, France), was added to the supernatant. Glucose conversion was complete after 30 min at 30 °C. Enzyme activity was determined spectrophotometrically (A_{340}) and calculated from the amount of NADPH (nmol) produced per second (nkat).

***In situ* staining of soluble AI activity**

All procedures were carried out at 4 °C. SAI activity was determined in both seeds and flowering spikes using histochemical staining according to Sergeeva and Vreugdenhil (2002). The samples, including seeds and handmade transverse sections of basal and apical parts of flowering shoots, were fixed immediately in 50 mM sodium phosphate buffer (pH 5.0) containing 2% (v/v) paraformaldehyde, 2% (w/v) polyvinylpyrrolidone 40, 1 μM dithiothreitol (DTT), 5 mM ethylenediaminetetraacetic acid (EDTA), 2 mM ethyleneglycol-bis(β -aminoethylether)-*N,N'*-tetraacetic acid (EGTA), 1 mM benzamidine and 2 mM ascorbic acid for 1 h. The fixation step was omitted for control samples intended for CWI staining. The endogenous soluble carbohydrates were removed by washing the samples three times in fixation buffer without paraformaldehyde, and then three times in water over a period of 5 h. Then, the samples were stained in 50 mM sodium phosphate buffer (pH 5.0) containing 294 μM nitroterazolium blue chloride, 460 μM phenazine methosulphate, 25 U/mL glucose oxidase (EC 1.1.3.4) and 100 mM sucrose for 1 h at 37 °C. Controls corresponded to samples incubated in the staining medium without sucrose. In parallel, sections were immersed in sodium hypochlorite (26% active chlorine) for 30 min, rinsed twice in water, incubated for 10 min in 10% (v/v) acetic acid, and then immersed in 0.4% (w/v)

Mirande's reagent for 30 min for better visualization of cellulosic and lignified tissues (Mondolot *et al.*, 2001). After three washes in water, photographs were taken with a Olympus SZX10 stereo microscope using an E330 camera (Olympus Europa GMBH, Hamburg, Germany). Fixed samples revealed both CWI and SAI activity, and unfixed samples revealed only CWI activity.

Purification of the SAI isoform

All procedures were carried out at 4 °C. Subterranean shoots (100 g fresh weight, stage IV) were homogenized in 0.6 L of buffer A (Table 3) in the presence of 50 g PVPP (Ultra-Turrax T25, IKA®-Labortechnik) and centrifuged (10 000 *g*, 20 min). Solid ammonium sulphate was added to 80% saturation of the supernatant (516 g/L). The precipitate formed after 2 h of gentle stirring and was collected following centrifugation, and dissolved in a minimal volume of buffer B (Table 3). Following centrifugation, the supernatant was desalted on PD-10 Sephadex columns (Amersham Pharmacia), previously equilibrated with buffer B.

SAI was purified sequentially on a concanavalin-A Sepharose affinity gel (Amersham Pharmacia, 10 cm \times 1 cm), anion exchanger (Econo-Pac High Q, 1 mL cartridge, Bio-Rad, Marnes-La-Coquette, France), HR Sephacryl column (30 cm \times 1 cm, size exclusion chromatography, GE Healthcare, Orsay, France) with liquid isoelectric focusing on a Rotofor preparative cell (Bio-Rad, Hercules, CA, USA) (Table 3). Protein fractions were desalted on PD-10 Sephadex columns prior to SAI assay on 96-well microplates using the 3,5-dinitrosalicylic acid method (Miller, 1959; iEMS 96 well Microplate Reader, MTX Lab Systems, Vienna, VA, USA). The active fractions were pooled and concentrated in Vivaspin 5-kDa microcentrifuge tubes (Sartorius Stedim France SAS, Aubagne, France). The molecular mass of the purified SAI was determined using an HR Sephacryl column following calibration using molecular mass standards (dextran, 2000 kDa;

thyroglobulin, 669 kDa; ferritin, 440 kDa; catalase, 232 kDa; aldolase, 158 kDa; ovalbumin, 43 kDa; ribonuclease A, 13.5 kDa; GE Healthcare).

Native and sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE), SAI activity staining on native gels and Western blotting

Nondenaturing PAGE was performed on a 12% native gel in a Tris-glycine buffer (pH 8.3) using the Mini Protean® II gel system (Bio-Rad, USA). The gels were stained with Coomassie Brilliant Blue G250 (Neuhoff *et al.*, 1988), or SAI activity was revealed using the method described by Gabriel and Wang (1969), or the proteins were transferred to a nitrocellulose membrane (Hybond-ECL™ Nitrocellulose membrane, GE Healthcare) for 1 h at 200 mA using the Mini Trans-Blot Transfer Cell (Bio-Rad, USA). SAI immunolabelling was performed using polyclonal antibodies raised against grape (*Vitis vinifera*) vacuolar invertase (Dambrouck *et al.*, 2005; 1:2500 dilution) and peroxidase-labelled goat anti-rabbit IgG antibodies (AffiniPure F(ab')₂ fragment, 1:12 500 dilution; Jackson Immuno-Research Europe Ltd., Newmarket, Suffolk, UK), according to the manufacturer's instructions (ECL Plus Western Blotting Detection System, GE Healthcare). Immunolabelled SAI was imaged on Kodak BioMax XAR Film (Sigma). In addition, SDS-PAGE of purified SAI was performed on a 12% gel using Precision Plus Protein™ unstained standards (Bio-Rad, Marnes-La-Coquette, France) as molecular mass standards (Laemmli, 1970).

Analysis by ESI-LC-MS/MS

The band showing SAI activity and immunolabelling was excised from native Coomassie Brilliant Blue-stained gels. In-gel enzymatic digestion of SAI, MS analysis and protein identification-databank searching and validation were performed according to Mary *et al.* (2010). Mass data collected during LC-MS/MS analysis were processed and searched for protein identification against the Uni-Prot/Swiss-Prot, Uni-Prot/TrEMBL and Institute of Genomic Research (TIGR) expressed sequence tag (EST) protein databanks.

Statistical analysis

An analysis of variance (ANOVA) was performed on the results from enzyme assays and quantitative PCR analyses with the isoform (or gene) or the development stage as the tested factor using SigmaPlot® ver. 10.0. Means of three independent RNA isolations or six independent enzyme extractions were tested at $P < 0.05$ (SNK test).

ACKNOWLEDGEMENTS

We gratefully acknowledge Dominique Bozec and Johannes Schmidt for their help in the glasshouse. MS analyses

were conducted at the platform 'Biopolymers-Interaction-Structural Biology' located at the INRA Centre of Angers-Nantes (http://www.angers-nantes.inra.fr/plateformes_et-plateaux_techniques/plate-forme_bibs). This research was supported by grants from the Syrian and French Ministries of Education and Research (grants to RD and TP, respectively).

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Fig. S1 Correlation between levels of *PrSAI1* transcripts and soluble acid invertase (SAI) activities in seeds and different

organs of *Phelipanche ramosa* parasitizing tomato plants. GS, germinated seed; PS, preconditioned seed; Tub.III, growing tubercle; Tub.IV, tubercle bearing a growing subterranean shoot (S.IV); Tub.V, tubercle bearing an emerged flowering shoot. Apical part of the flowering shoot is growing (FS.V) and bears fruits containing developing seeds (F.V). Basal part (Bp.V) does not bear flowers and is larger and more fibrous than the flowering shoot. The accumulation of *PrSAI1* transcripts was expressed relative to elongation factor 1- α gene (*EF1 α*) transcript levels. Data are means \pm standard error ($n = 3$ for quantitative polymerase chain reaction analysis, $n = 6$ for enzyme assay).

Fig. S2 Polyacrylamide gel electrophoresis (PAGE) of PrSAI1 purified from growing subterranean shoots of *Phelipanche ramosa* parasitizing tomato plants. (a) Colloidal blue staining. (b) Soluble acid invertase (SAI) staining. (c) Immunoblotting using polyclonal antibodies targeted against grape (*Vitis vinifera*) vacuolar invertase. CE, 50 μ g crude extract; Con-A, 10 μ g concanavalin-A fraction; MM, molecular markers; Mono Q, 5 μ g Mono Q fraction; 80% SA, 25 μ g 80% ammonium sulphate fraction.

Table S1 Degenerate primer pairs designed for cloning invertase polymerase chain reaction (PCR) fragments.

Table S2 Primer sequences for cloning full-length invertase cDNAs.

Table S3 PrSAI1 purification from subterranean shoots of *Phelipanche ramosa* parasitizing tomato plants.

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