



## Members of the aquaporin family in the developing pea seed coat include representatives of the PIP, TIP, and NIP subfamilies

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### Abstract

Water and nutrients required by developing seeds are mainly supplied by the phloem and have to be released from a maternal parenchyma tissue before being utilized by the filial tissues of embryo and endosperm. To identify aquaporins that could be involved in this process four full-length cDNAs were cloned and sequenced from a cDNA library of developing seed coats of pea (*Pisum sativum* L.). The cDNA of *PsPIP1-1* appeared to be identical to that of clone 7a/TRG-31, a turgor-responsive gene cloned previously from pea roots. *PsPIP1-1*, *PsPIP2-1*, and *PsTIP1-1*, or their possible close homologues, were also expressed in cotyledons of developing and germinating seeds, and in roots and shoots of seedlings, but transcripts of *PsNIP-1* were only detected in the seed coat. In mature dry seeds, high hybridization signals were observed with the probe for *PsPIP1-1*, but transcripts of *PsPIP2-1*, *PsTIP1-1*, and *PsNIP-1* were not detected. Functional characterization after heterologous expression in *Xenopus* oocytes showed that *PsPIP2-1* and *PsTIP1-1* are aquaporins whereas *PsNIP-1* is an aquaglyceroporin. *PsNIP-1*, like several other NIPs, contains a tryptophan residue corresponding with Trp-48 in GlpF (the glycerol facilitator of *Escherichia coli*) that borders the selectivity filter in the permeation channel. It is suggested that *PsPIP1-1* and/or its possible close homologues could play a role in water absorption during seed imbibition, and that *PsPIP2-1*, possibly together with *PsPIP1-1*, could be involved in the release of phloem water from the seed coat symplast, which is intimately connected with the release of nutrients for the embryo.

**Abbreviations:** MIPs, major intrinsic proteins; NIPs, nodulin 26-like intrinsic proteins; PIPs, plasma membrane intrinsic proteins; SIPs, small, basic intrinsic proteins; TIPs, tonoplast intrinsic proteins

### Introduction

The phloem is the food-conducting tissue of vascular plants. Its sieve elements constitute a symplasmic route by which organic nutrients (mainly sucrose and amino acids) along with inorganic nutrients (mainly K<sup>+</sup>) are translocated hydraulically from source to sink (Münch, 1930; Fisher, 2000). Developing seeds are strong sinks, particularly during the storage phase. In developing legume seeds the nutrients arriving by

the phloem are transferred, via plasmodesmata, from the sieve elements to adjacent parenchyma cells of the seed coat (phloem unloading). Because the embryonic tissues are symplasmically isolated from the maternal seed coat, the nutrients have to be released from seed coat parenchyma cells into the apoplast (seed coat unloading). The molecular basis for the efflux of these solutes is still unknown. It has been suggested that in seed coats of common bean (*Phaseolus vulgaris*) and broad bean (*Vicia faba*) sucrose is

released by an H<sup>+</sup>-antiport mechanism (Patrick and Offler, 2001), whereas work in our group on pea has provided evidence for poorly selective pores in the plasma membrane of seed coat parenchyma cells that allow the passage of sugars, amino acids and possibly inorganic ions as well (de Jong *et al.*, 1996, 1997; van Dongen *et al.*, 2001).

Seed coat unloading implies the transport of a range of nutrients as well as of phloem water (Münch water) across the cell membranes of seed coat parenchyma cells. It may be anticipated that aquaporins are involved in the efflux of water from the parenchyma cells. Aquaporins belong to the family of major intrinsic proteins (MIPs), an ancient family with members in all life forms (Agre *et al.*, 1998; van Dongen and Borstlap, 2003). Based on sequence homology, members of the MIP family in higher plants fall into four major subgroups: plasma membrane intrinsic proteins (PIPs), tonoplast intrinsic proteins (TIPs), Nodulin 26-like MIPs (NLMs or NIPs), and small, basic intrinsic proteins (SIPs). Representatives of the PIPs, TIPs, NIPs, and SIPs have been identified in the moss *Physcomitrella patens*, indicating that the four subfamilies were already established when land plant evolution began (Borstlap, 2002). The PIP subfamily can be further divided into the highly homologous but clearly distinct subgroups PIP1 and PIP2, whereas the TIP subfamily consists of five subgroups, including the  $\alpha$ -,  $\gamma$ -, and  $\delta$ -TIPs which may be markers of different types of vacuoles (Jauh *et al.*, 1999). According to a new nomenclature the subgroups of  $\alpha$ -,  $\gamma$ -, and  $\delta$ -TIPs are now designated TIP3, TIP1, and TIP2, respectively (Johanson *et al.*, 2001; Chaumont *et al.*, 2001).

Functionally, MIPs fall into two main categories: aquaporins and glycerol facilitators (Heymann and Engel, 1999; Agre *et al.*, 1998). High-resolution atomic structures of prototypes of each category, aquaporin-1 (AQP1) from mammals and the glycerol facilitator (GlpF) from *Escherichia coli*, have elucidated aspects of the selectivity of these transporters (Fu *et al.*, 2000; Sui *et al.*, 2001). To some extent the glycerol-transporting MIPs, including GlpF, also allow the passage of water (Borgnia and Agre, 2001), hence their designation as aquaglyceroporins. Some mammalian MIPs transport larger molecules or ions. AQP9 from the liver allows the passage of water and a variety of neutral molecules including carbamides, polyols, purines and pyrimidines (Tsukaguchi *et al.*, 1998), whereas AQP6, an endomembrane aquaporin

in renal epithelia, is permeable to anions, particularly nitrate (Ikeda *et al.*, 2002).

Seed coat parenchyma cells are committed to release a range of solutes that serve as nutrients for the developing seed tissues and as raw materials for the synthesis of storage compounds. Münch's mass flow concept of phloem transport dictates that this solute release must be intimately connected with the release of water. The question may be raised whether MIPs are involved in the flux of water and solutes across the membrane of seed coat parenchyma cells. Here we report the cloning, sequencing and functional characterization of four MIP genes from a cDNA library of developing pea seed coats, which turned out to encode members of the PIP1, PIP2, TIP1, and NIP subgroup, respectively.

## Materials and methods

### *Plant material*

Pea plants (*Pisum sativum* L. cv. Marzia) were grown in a growth chamber as described (de Jong and Wolswinkel, 1995). Developing seeds used for RNA gel blot analysis were taken from pods tagged at the time of flowering. To obtain seedling tissues for RNA gel blot analysis, pea seeds were imbibed in aerated tap water for 24 h and then placed on moist filter paper in petri dishes in the dark at room temperature.

### *Construction of a seed coat cDNA library*

Coats from developing pea seeds in the filling stage were collected and immediately frozen in liquid nitrogen. Total RNA was isolated from 10 g fresh weight of seed coats by the guanidine hydrochloride extraction method (Logemann *et al.*, 1987). Poly(A)<sup>+</sup> RNA was prepared by oligo(dT)-cellulose chromatography (Sambrook *et al.*, 1989). The library was constructed in the  $\lambda$  Uni-ZAP XR vector and packaged into phage particles with the ZAP-cDNA Gigapack II Gold Cloning kit from Stratagene. The resulting library consisted of  $9.7 \times 10^6$  plaque forming units with an average insert length of 1.1–1.3 kb.

### *Reverse transcriptase polymerase chain reaction (RT-PCR)*

PCR was performed with two degenerate primers corresponding to the conserved NPA boxes of the MIP family. The sense primer 5'-(T/C)T(A/C/G/T)AA(C/T)

CC(A/C/G/T)GC(A/C/G/T)GT(A/C/G/T)AC-3' corresponds to the amino acid sequence Leu-Asn-Pro-Ala-Val-Thr in the B-loop, and the antisense primer 5'-AA(A/C/G/T)(C/G)(A/T)(A/C/G/T)C(G/T)(A/C/G/T)GC(A/C/G/T)GG(A/G)TT-3' corresponds to the sequence Asn-Pro-Ala-Arg-Ser-Phe in the E-loop. First-strand cDNA was synthesized from 5 µg of poly(A)<sup>+</sup> RNA from pea seed coats with M-MuLV reverse transcriptase (Boehringer) and the antisense primer. The PCR-mixture consisted of 10 mM Tris-HCl pH 9.0, 50 mM KCl, 1.5 mM MgCl<sub>2</sub>, 200 µM of each deoxynucleotide triphosphate, 3.8 µM of each primer, cDNA corresponding to 300 ng of poly(A)<sup>+</sup> RNA, and 1.5 units of *Taq* DNA polymerase (Boehringer) in a total volume of 50 µl. After 4 min of denaturation at 92 °C the amplification was performed in 35 cycles (1 min 92 °C, 1 min 50 °C, and 1 min 72 °C). PCR products were subjected to phenol/chloroform extractions and precipitated with ethanol. Primers were removed on a SephacrylS-400HR column (Pharmacia). After electrophoresis on an agarose gel a band corresponding to a ca. 400 bp PCR product was isolated, purified with QiaexII (Qiagen) and cloned into the pGEM-3Z plasmid vector (Promega).

#### *Labeling of cDNA probes and screening of the cDNA library*

The PCR fragments of *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1* were radiolabeled with [ $\alpha$ -<sup>32</sup>P]dCTP (3000 Ci/mmol; ICN) by random primer labeling (Feinberg and Vogelstein, 1983) with 40–50 ng of cDNA as a template (Ready-to-Go-DNA-labeling kit, Pharmacia). Excess nucleotides were removed with Microspin S400-HR columns (Pharmacia). The cDNA library was plated on the *Escherichia coli* host strain XL1-Blue MRF<sup>r</sup>. Plaques were blotted onto Hybond-N<sup>+</sup> membranes (Amersham) and hybridization was performed for 16–18 h at 63 °C in 0.5 M sodium phosphate pH 7.2, 1 mM EDTA and 7% w/v SDS, with the [<sup>32</sup>P]-labeled probes of the four MIP gene fragments. Blots were washed three times for 30 min at 65 °C in 2× SSC, 0.5% w/v SDS and autoradiographed (Kodak X-Omat AR) at –80 °C for 2–5 days. Positive clones were isolated from the agar plates and stored in phage buffer (20 mM Tris-HCl pH 7.4, 100 mM NaCl, 10 mM MgSO<sub>4</sub>, 0.01% gelatin). A second screening was performed similarly except that the phage suspension was plated at lower density, allowing the isolation of single phage clones. Following Stratagene's protocol of the ZAP-cDNA Gigapack II

Gold Cloning kit, a helper phage (Exassist) was added to the clones selected from the second screening to excise *in vivo* the pBluescript phagemid from the  $\lambda$  vector.

#### *RNA gel blot analysis*

Collected plant material was frozen immediately in liquid nitrogen and ground to a fine powder using a mortar and pestle. Aliquots of the homogenized tissue (0.5 g) were stored at –80 °C until use for RNA extraction. Total RNA was extracted in an equal volume of extraction buffer (0.35 M glycine, 0.048 M NaOH, 0.34 M NaCl, 0.04 M EDTA, 4% w/v SDS). The homogenates were extracted with phenol and chloroform and the RNA was precipitated with LiCl as described by Sambrook *et al.* (1989). For RNA analysis, 15 µg of RNA was denatured with glyoxal and DMSO (Sambrook *et al.*, 1989), separated electrophoretically on 1.5% w/v agarose gels and blotted onto Hybond-N<sup>+</sup> membranes by capillary transfer. The electrophoresis buffer and blotting buffer consisted of 10 mM and 25 mM sodium phosphate pH 7, respectively. Prior to hybridization RNA blots were washed in 20 mM Tris-HCl pH 8.0 at 65 °C for 1 h. RNA blots were hybridized overnight at 65 °C in 0.5 M sodium phosphate pH 7.2, 7% w/v SDS, 1 mM EDTA with the [<sup>32</sup>P]-labeled, ca. 400 bp PCR fragments of *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1* as probes. Blots were washed three times for 30 min at 65 °C with 2× SSC, 0.5% w/v SDS and hybridization signals were detected by autoradiography (–80 °C; Kodak X-Omat-AR) or by phosphoimaging (BioRad Molecular Imager FX; program Quantity One). Film scans and phosphoimager files were processed in Adobe PhotoShop 5.5. Hybridization with an 851 bp RT-PCR fragment (sense primer 5'-GGCTCATTGAGTCTGGTA-3', antisense primer 5'-GTCGAGGTCTCGTTCGTT-3') from *Pisum sativum* 18S rRNA (accession number U43011) was used as loading control.

#### *Sequencing and sequence analysis*

Sequencing was performed with the DyeDeoxy Terminator Cycle Sequencing kit. Similarity searches were performed by the Advanced BLAST2 Search Service of Bork group at EMBL, Heidelberg (Altschul *et al.*, 1997). Multiple alignments were performed by MultAlin, University of Toulouse, France (Corpet, 1988).

### Subcloning into an oocyte expression vector

cDNA clones encoding PsPIP1-1, PsPIP2-1, PsTIP1-1 and PsNIP-1 were inserted in pT7TS, a vector for transient expression in *Xenopus* oocytes (Johnson and Krieg, UT, Austin, TX). This vector, which is closely related to pSP64T (Krieg and Melton, 1984), contains a 5'- and 3'-untranslated region that originates from *Xenopus*  $\beta$ -globin flanking the polylinker site and a 3' poly(A) tail of 30 nucleotides to increase cRNA stability after injection in *Xenopus* oocytes. Sticky ends of the cDNA clones were made blunt using the Klenow fragment of DNA polymerase I (USB, Cleveland, USA) and ligated into the *EcoRV* site of pT7TS. Constructs were sequenced to ensure that no errors were introduced in the coding sequence. The pT7TS oocyte expression vectors with the different inserts of pea MIP clones were used as template for *in vitro* cRNA preparation.

### Preparation of cRNA

Enzymes and supplements were obtained from Roche Diagnostics (Mannheim, Germany) unless stated otherwise. Plasmids were isolated with a Plasmid DNA isolation kit (Qiagen, Hilden, Germany) and linearized with *SalI*. For *in vitro* transcription, ca. 2  $\mu$ g DNA was incubated for 2 h at 37 °C in 50  $\mu$ l transcription mixture (15 U T7 RNA polymerase; Promega, Madison, WI), 1 mM of each rNTP, 0.75 mM of the methylated cap analogue (m<sup>7</sup>G(5')ppp(5')G), 40 U RNasin (Promega), 10 mM DTT, 2 mM spermidine, 10 mM NaCl, 6 mM MgCl<sub>2</sub>, 40 mM Tris, pH 7.5 in ultrapure water). To increase the yield, another 15 U of RNA polymerase was added after 1 h. The reaction was stopped by a 15 min incubation with 2 U of RQ1 RNase-free DNase (Promega) at 37 °C. Proteins were extracted from the reaction mixture by a phenol/chloroform extraction and the free nucleotides were separated from the cRNA using a Sephadex column. G50-Sephadex (fine, DNA grade, Pharmacia, Uppsala, Sweden) in TNE (20 mM Tris pH 7.5, 50 mM NaCl, 1 mM EDTA) was autoclaved and brought into a Pasteur pipette with a small piece of glass wool in the tip (1.5 ml settled bed volume). The cRNA samples were diluted with sterile ultrapure water to 100  $\mu$ l, loaded onto the column, and eluted with TNE. The eluate was collected in fractions of 100  $\mu$ l in which the cRNA concentration was determined spectrophotometrically (GeneQuant, Pharmacia). Intact cRNA was eluted as a distinct peak in fractions 8–11. The cRNA of these fractions was pre-

cipitated and dissolved in water to a concentration of 0.5  $\mu$ g/ $\mu$ l.

### Injection of cRNA into oocytes

Ovaries were surgically isolated from *Xenopus laevis* frogs anaesthetized with 0.1% w/v 3-aminobenzoic acid ethyl ester methanesulfonate salt in water (Sigma Aldrich, Saint Louis, MO) and placed in modified Barth's solution (MBS: 0.33 mM Ca(NO<sub>3</sub>)<sub>2</sub>, 0.41 mM CaCl<sub>2</sub>, 88 mM NaCl, 1 mM KCl, 2.4 mM NaHCO<sub>3</sub>, 10 mM Hepes pH 7.4, 0.82 mM MgSO<sub>4</sub>). An ovary was torn into small pieces, containing about 10 oocytes each and incubated in MBS with ca. 15 U collagenase A (Roche Diagnostics) in a final volume of 30 ml for 2 h at room temperature. At the end of the reaction, the collagenase was carefully washed away by rinsing the oocytes five times in 30 ml MBS. Stage V and VI oocytes (Dumont, 1972) were collected and stored at 18 °C in MBS supplied with 10  $\mu$ g/ml gentamycin (Sigma Aldrich). The next day, oocytes were injected with 25 ng cRNA in 50 nl ultrapure water for determining water permeability, or 11.5 ng cRNA in 23 nl ultrapure water for solute uptake experiments by using a Nanoject Auto/Oocyte Injector (Drummond Sci. Co., PA). As a control treatment, oocytes were injected with equal volumes of ultrapure water. The day after injection the vitelline membrane around vital oocytes was manually peeled off with two forceps. This horrid job was made easier by incubating oocytes for 5 min in a hypertonic solution (250 mM sucrose, 60 mM KCl, 10 mM EGTA, 8 mM MgCl<sub>2</sub>, 40 mM Hepes, pH 7.0) by which the vitelline layer became detached from the oocyte (Silberberg and Magleby, 1997).

### Water permeability assay

The osmotic water permeability of the oocyte membrane was determined one day after cRNA injection. Single oocytes were transferred to 10% v/v MBS and immediately observed under a microscope (Zeiss Axioskoop equipped with a Nikon digital camera DXM1200). For 2 min, a sequence of digital photographs was made at 5 s intervals. The osmotic water permeability ( $P_f$ ) of the oocyte membrane was calculated from the initial rate of cell volume change with the formula

$$P_f = \frac{V_0(dV_{rel}/dt)}{S \cdot V_w \cdot (osm_{in} - osm_{out})}$$



ment, AJ548795; *PsPIP2-1*, AJ243307; *PsTIP1-1*, AJ243309; *PsNIP-1*, AJ243308.

## Results

### Cloning and sequencing of four MIPs

RT-PCR of mRNA from developing pea seed coats was performed with primers corresponding to the conserved NPA boxes of the MIP family. Electrophoresis of the PCR products on agarose gel yielded a single band of ca. 400 bp (not shown). The PCR products were cloned into the plasmid vector pGEM-3Z. Transformation of *E. coli* cells yielded 37 independent clones. Five distinct PCR fragments could be identified that showed homology to MIP genes. Two fragments belonged to the PIP1 group, three other fragments to the PIP2, TIP1, and NIP subgroup, respectively. Restriction analysis indicated that 12 clones contained the PIP1-1 fragment, 3 clones the PIP1-2 fragment, 4 clones the PIP2 fragment, 1 clone the TIP1 fragment, and 2 clones the NIP fragment.

To obtain full-length sequences the seed coat cDNA library was screened using the [<sup>32</sup>P]-labeled PCR-fragments. From ca. 50 000 plaques, ca. 200 positive clones were obtained with the *PIP1-1* fragment, 80 with the *PIP2* fragment, 80 with the *TIP1* fragment, and 40 with the *NIP* fragment. These abundances of the four transcripts in the cDNA library are in reasonable agreement with their abundances in the mixture of PCR products. Sequencing of the full-length cDNAs confirmed that they encode members of the PIP1, PIP2, TIP1, and NIP subgroup. The nucleotide sequence of the full-length clone of *PsPIP1-1* appeared to be identical to that of *clone 7a / TRG-31*, a turgor-responsive gene cloned previously from pea roots (Guerrero *et al.*, 1990; Guerrero and Crossland, 1993). No attempt was made to obtain a full-length clone of *PIP1-2*. The other full-length clones were named *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1* respectively. An alignment of the deduced amino acid sequences of the four genes together with those of AQP1 and GlpF is shown in Figure 1.

### Expression in developing, mature, and germinating seeds

Expression of aquaporin genes was examined by RNA gel blot analysis with the [<sup>32</sup>P]-labeled PCR fragments (ca. 400 nucleotides) of *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1* as probes. Dot-blots showed

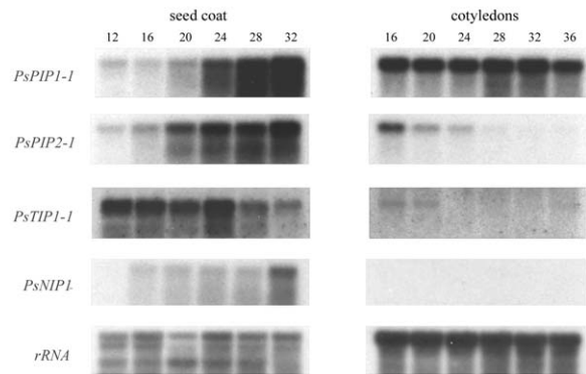


Figure 2. RNA gel blot analysis of seed coat and cotyledons during seed development, with the probes for *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1*. Tissues were harvested 12–36 days after flowering (d.a.f.).

no cross-hybridization, but it is possible that each probe also hybridized with mRNA of its possible close homologues. This is particularly true of the probes of *PsPIP1-1* and *PsPIP2-1* because it is likely that in pea, as in *Arabidopsis* and maize, some members of the PIP1 and PIP2 subfamilies are highly similar isoforms. *PsPIP1-1* and *PsPIP1-2* may be such isoforms since their PCR fragments showed 83% identity.

The expression patterns in seed coat and cotyledons during seed development are shown in Figure 2. In cotyledons a high and remarkably constant hybridization signal was observed with the *PsPIP1-1* probe, whereas the signals with the probes of *PsPIP2-1* and *PsTIP1-1* were confined to the earlier stages of development (<24 days after flowering), and transcripts of *PsNIP-1* could not be detected at all. In seed coats the hybridization signal for the *PsTIP1-1* probe was almost invariable up to 28 days after flowering, and then slightly decreased. By contrast, the signals observed with the probes of *PsPIP1-1*, *PsPIP2-1* and *PsNIP-1* strongly increased in the final stage of seed development (>20 days after flowering).

In the dry, mature seed transcripts of *PsPIP1-1* and/or its close homologues were abundantly present whereas those of *PsPIP2-1*, *PsTIP1-1* and *PsNIP-1* were not detected (Figure 3). After imbibition the hybridization signal of the *PsPIP1-1* probe in the cotyledons first declined, but increased again after four days of imbibition. Hybridization signals of the *PsTIP1-1* probe also reappeared in the cotyledons of the germinating seed, whereas transcripts of *PsNIP-1* and *PsPIP2-1* remained undetectable. In roots of the etiolated pea seedlings high hybridization signals were seen with the probes of *PsPIP1-1*, *PsPIP2-1* and

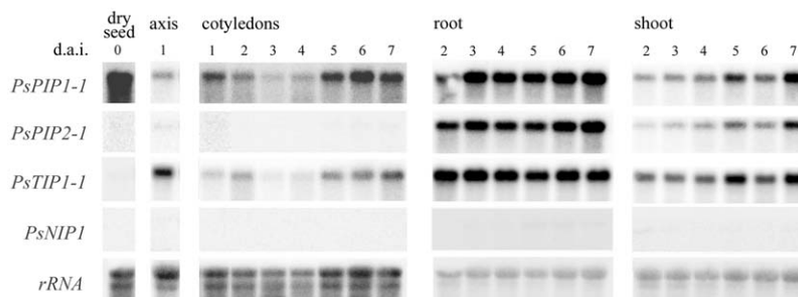


Figure 3. RNA gel blot analysis of dry seeds and of cotyledons, shoots, and roots of etiolated seedlings, using the probes for *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1* and *PsNIP1-1*. Tissues were harvested 0–7 days after imbibition (d.a.i.).

*PsTIP1-1*. The signals were lower in seedling shoots and tended to increase during the 7-day growth period. Transcripts of *PsNIP1* could not be detected in roots and shoots of the seedlings.

#### Functional characterization

The coding sequences of *PsPIP1-1*, *PsPIP2-1*, *PsTIP1-1*, and *PsNIP1-1* were sub-cloned into a vector designed for the preparation of cRNA required for transient expression in *Xenopus* oocytes. The osmotic water permeability ( $P_f$ ) of oocytes was determined from the initial swelling rate of oocytes after transfer to a hypotonic solution (Figure 4). Expression of *PsPIP1-1* in *Xenopus* oocytes had no effect on the swelling rate of the oocytes, whereas *PsPIP2-1* increased  $P_f$  about 5 times, from  $10.2 \pm 0.7 \mu\text{m/s}$  to  $52.3 \pm 4.2 \mu\text{m/s}$ . After expression of *PsTIP1-1* and *PsNIP1-1*, the  $P_f$  of oocytes slightly but significantly increased to  $12.3 \pm 0.7 \mu\text{m/s}$  and  $13.6 \pm 0.9 \mu\text{m/s}$ , respectively.

With the oocyte expression system we also tested the permeability of the four MIPs for urea, glycerol, glycine, and orthophosphate. The permeability of certain aquaporins for glycerol and/or urea is well known, and evidence that GlpF is permeable for glycine has been reported (Heller *et al.*, 1980). We also tested the permeability for phosphate because the cDNAs of two MIP genes from potato have been found to complement a phosphate uptake-deficient yeast mutant (Leggewie *et al.*, 1995). In some experiments a highly significant stimulation of solute uptake was observed. For instance, in one experiment conducted at pH 5.5, the expression of *PsPIP1-1* increased the uptake of glycerol and glycine more than three-fold, whereas no stimulation of the uptake of urea and phosphate was observed (not shown). Unfortunately, these results could not be reproduced in further experiments. In

fact, we were unable to show any reproducible stimulation in the uptake of urea, glycine, or phosphate after the expression of *PsPIP1-1*, *PsPIP2-1*, or *PsTIP1-1*. Uptake of glycerol by oocytes, however, was consistently enhanced after the expression of *PsNIP1-1* but not after expression of the other three seed coat MIPs (Figure 5).

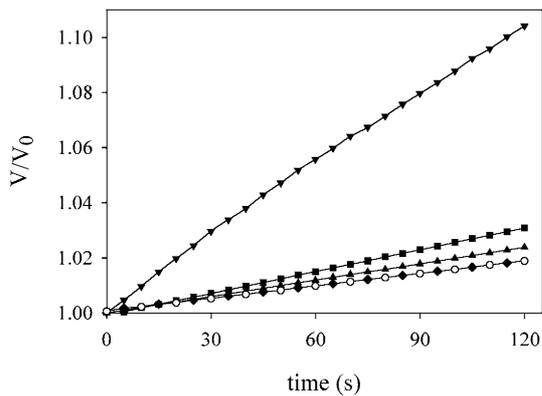
#### Discussion

##### *Aquaporins and the water relations of developing seeds*

During the early stages of development, the dry matter content of the pea seed (ca. 20%) is similar to that of the phloem sap (Lewis and Pate, 1973), which demonstrates that the developing seed can obtain sufficient water from the phloem sap stream. The vascular bundle that runs from the funiculus into the seed coat consists largely of phloem elements but it also contains a central xylem strand (van Dongen *et al.*, 2003). With acid fuchsin as a tracer it can easily be shown, however, that water from the xylem sap stream does not enter into developing legume seeds (P. Wolswinkel and J.T. van Dongen, unpublished observations). It is well known that the water relations of developing seeds are largely independent of those in the rest of the plant (Bradford, 1994; Shackel and Turner, 2000), possibly because water is supplied exclusively by the phloem.

At the end of the filling stage, but well before the final desiccation, the dry matter content of the pea seed had increases to ca. 45%. During seed fill, therefore, the phloem sap stream supplies more water than is needed by the seed. The excessive amount of water is probably cycled back into the xylem, but it is not known which cells are involved in this water flux.

Phloem water entering the developing seed has first to be released from the seed coat symplast by crossing



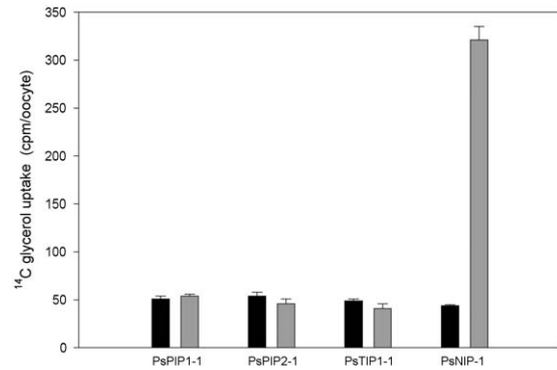
**Figure 4.** Relative volume change ( $V/V_0$ ) of oocytes subjected to a swelling assay. Oocytes of *Xenopus laevis* were injected with water (○) or with 25 ng cRNA coding for PsPIP1-1 (◆), PsPIP2-1 (▼), PsTIP1-1 (▲) and PsNIP-1 (■). The next day, the vitelline membrane was peeled off and oocytes were placed in 10% MBS. Volume changes of individual oocytes from four different frogs were monitored during the first two minutes. For each MIP the result obtained with a single, representative oocyte is shown. Mean values of the osmotic water permeability ( $P_f$  in  $\mu\text{m/s}$ ) were as follows: PsPIP2-1,  $52.3 \pm 4.2$ ; PsNIP-1,  $13.6 \pm 0.9$ ; PsTIP1-1,  $12.3 \pm 0.7$ ; PsPIP1-1,  $9.8 \pm 0.5$ ; control (water injected),  $10.2 \pm 0.7$ .

the plasma membranes of sieve element/companion cell complexes or seed coat parenchyma cells. It seems likely that the aquaporin PsPIP2-1, possibly along with PsPIP1-1 (see below), participates in regulating the transmembrane water movements that are linked to phloem unloading. The possibility that additional aquaporins play a role cannot be excluded because our inventory of aquaporin genes expressed in the pea seed coat is probably not exhaustive.

Translocation of assimilates through the phloem is a hydraulically operated process, but information about the involvement of aquaporins is scanty (Patrick *et al.*, 2001). A study of the distribution of PIP1 and TIP1 homologous proteins in leaves of *Brassica napus* showed that both types of protein were present in phloem elements of the minor veins but at a much lower level than in parenchyma cells of mesophyll and vascular bundles (Frangne *et al.*, 2001). In phloem sap of barley transcripts of the PIP1 member *emip* have been detected (Doering-Saad *et al.*, 2002).

#### *Abundant transcripts of a PIP1 homologue in the dry seed*

Inasmuch as PIP2 members expressed in *Xenopus* oocytes induce a relatively high water permeability they may be regarded as the water channels in plant plasma membranes. The function of PIP1 members is less



**Figure 5.** Glycerol uptake by *Xenopus* oocytes injected with water (black columns) or with 25 ng cRNA coding for PsPIP1-1, PsPIP2-1, PsTIP1-1 and PsNIP-1 (gray columns). The results are from representative experiments, each with 18–24 oocytes. Experiments with PsTIP1-1 and PsNIP-1 were repeated once, that with PsPIP1-1 three times.

clear because, after expression in oocytes, they show a much lower aquaporin activity or no activity at all (Figure 4; Chaumont *et al.*, 2000; Moshelion *et al.*, 2002). Nevertheless, evidence is now accumulating that PIP1 members function in the plant as aquaporins. First, it may be noted that the amino acid residues predicted to border the selectivity filter are the same in PIP1 and PIP2 members (exemplified in Table 1 for PsPIP1-1 and PsPIP2-1). Secondly, recent studies in which the expression of PIP1 members was either impaired or enhanced have provided strong evidence that PIP1 members function in cellular and whole-plant water relations (Siefritz *et al.*, 2002; Martre *et al.*, 2002; Aharon *et al.*, 2003).

Members of *PIP1* and *PIP2* are often co-expressed in plant tissues, but in the dry seed transcripts of *PsPIP2-1* could not be detected (Figure 3). The absence of PIP2 members in dry seeds has been noted before. Neither the RD28 protein from *Arabidopsis* nor the mRNA of *BnPIP1* from *Brassica napus* was detectable in dry seeds (Daniels *et al.*, 1994; Gao *et al.*, 1999). In dry seeds of radish transcripts of both a *PIP2* homologue (*PAQ2*) and a PIP1 homologue (*PAQ1*) were undetectable (Suga *et al.*, 2001). We found the mRNA of *PsPIP1-1* and/or its possible close homologues to be abundantly present in the dry pea seed (Figure 3). To the best of our knowledge, this is the first demonstration of transcripts of a *PIP* homologue in a dry seed, which raises the possibility that PIP1 members play a role in the rehydration of the dry seed during imbibition.

**Table 1.** Amino acid residues lining the selectivity filter and at Froger's positions (P<sub>1</sub>–P<sub>5</sub>) in AQP1 and GlpF and the corresponding residues in several other MIPs. The functional characterization as aquaporin or aquaglyceroporin, and the subgroups to which the plant MIPs belong are indicated. Tryptophan residues corresponding with Trp-48 in GlpF and the residues considered being typical of glycerol transporters (Froger *et al.*, 1998) are printed in bold.

Code	Description	Selectivity filter	P <sub>1</sub>	P <sub>2</sub>	P <sub>3</sub>	P <sub>4</sub>	P <sub>5</sub>				
HsAQP1	human aquaporin	F R C H T S A F W	F	R	C	H	T	S	A	F	W
CvAQPcic	insect aquaporin	F R S H T S A Y W	F	R	S	H	T	S	A	Y	W
EcAqpZ	bacterial aquaporin	F R T H A S A F W	F	R	T	H	A	S	A	F	W
PsPIP1-1	PIP1	F R T H E S A F W	F	R	T	H	E	S	A	F	W
PsPIP2-1	PIP2; aquaporin	F R T H Q S A F W	F	R	T	H	Q	S	A	F	W
NtAQP1	PIP1; aquaglyceroporin	F R T H M S A F W	F	R	T	H	M	S	A	F	W
SsAQP1	PIP1; aquaglyceroporin	F R T H E S A F W	F	R	T	H	E	S	A	F	W
PsTIP1-1	TIP1; aquaporin	F V A I T S A Y W	F	V	A	I	T	S	A	Y	W
NtTIPa	TIP4; aquaglyceroporin	H R A I T S A Y W	H	R	A	I	T	S	A	Y	W
GmNod26	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> S A Y <b>L</b>	<b>W</b>	R	A	V	<b>F</b>	S	A	Y	<b>L</b>
PsNIP-1	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> S A Y <b>M</b>	<b>W</b>	R	A	V	<b>F</b>	S	A	Y	<b>M</b>
LjLIMP2	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> S A Y <b>L</b>	<b>W</b>	R	A	V	<b>F</b>	S	A	Y	<b>L</b>
AtNLM1	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> S A Y <b>L</b>	<b>W</b>	R	A	V	<b>F</b>	S	A	Y	<b>L</b>
AtNLM2	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> S A Y <b>I</b>	<b>W</b>	R	A	V	<b>F</b>	S	A	Y	<b>I</b>
PtNIP1;1	NIP; aquaglyceroporin	<b>W</b> R A V <b>F</b> T A Y <b>M</b>	<b>W</b>	R	A	V	<b>F</b>	T	A	Y	<b>M</b>
EcGlpF	bacterial aquaglyceroporin	<b>W</b> R F G <b>Y</b> <b>D</b> <b>K</b> <b>P</b> <b>L</b>	<b>W</b>	R	F	G	<b>Y</b>	<b>D</b>	<b>K</b>	<b>P</b>	<b>L</b>
ScFPS1	yeast aquaglyceroporin	<b>W</b> R T N <b>W</b> <b>D</b> <b>R</b> <b>P</b> <b>M</b>	<b>W</b>	R	T	N	<b>W</b>	<b>D</b>	<b>R</b>	<b>P</b>	<b>M</b>
HsAQP3	human aquaglyceroporin	F R Y G <b>Y</b> <b>D</b> <b>R</b> <b>P</b> <b>I</b>	F	R	Y	G	<b>Y</b>	<b>D</b>	<b>R</b>	<b>P</b>	<b>I</b>

The analyzed sequences with accession numbers and species names are listed: AtNLM1 (O48595) and AtNLM2 (O49406) from *Arabidopsis thaliana*; CvAQPcic (Q23808) from *Cicadella viridis* (green cicada); EcAqpZ (P48838) and EcGlpF (P11244) from *Escherichia coli*; GmNod26 (P08995) from *Glycine max* (soybean); HsAQP1 (P29972) and HsAQP3 (Q92482) from *Homo sapiens*; LjLIMP2 (Q9LKJ5) from *Lotus japonicus*; NtTIPa (Q9XG70) and NtAQP1 (O24662) from *Nicotiana tabacum* (common tobacco); PsPIP1-1 (P25794), PsPIP2-1 (Q9XGG8), PsTIP1-1 (Q9XGG6) and PsNIP-1 (Q9XGG7) from *Pisum sativum* (garden pea); PtNIP1;1 (Q940D9) from *Pinus taeda* (loblolly pine); ScFPS1 (P23900) from *Saccharomyces cerevisiae* (baker's yeast); SsAQP1 (O65356) from *Samanea saman* (rain tree).

*Some TIP1 members ( $\gamma$ TIPs) are highly expressed in tissues with intensive symplasmic nutrient transfer*

The TIP subfamily can be divided into five subgroups (Johanson *et al.*, 2001; Chaumont *et al.*, 2001), of which the  $\alpha$ -,  $\gamma$ -, and  $\delta$ -TIPs have been studied most extensively. Vacuoles containing seed-type storage proteins are marked by  $\alpha$ -TIPs together with  $\gamma$ - and/or  $\delta$ -TIPs, whereas those marked by  $\gamma$ -TIP alone have characteristics of lytic vacuoles (Jauh *et al.*, 1999). Being the aquaporins of the tonoplast, TIPs are thought to permit a more rapid transcellular water flow by increasing the effective cross-section of the cytoplasm, and to facilitate osmotic adjustment between cytosol and vacuole. It has been proposed that cells with an intensive symplastic transport of assimilates or mineral ions may need an increased capacity for intracellular osmotic equilibration (Barrieu *et al.*, 1998).

The TIP identified in the cDNA library of pea seed coats clearly belongs to the TIP1 subgroup. *PsTIP1-1* and/or its possible close homologues, was expressed in cotyledons, in roots and shoots of seedlings, and at a fairly high level in the seed coat (Figures 2 and 3). The pea seed coat consists of an outer sclerenchyma layer and an inner layer of parenchyma (chlorenchyma, ground parenchyma and branched parenchyma) in which three vascular bundles are embedded. In the stage of seed fill the branched parenchyma has become deteriorated, and the cells of the chlorenchyma/ground parenchyma layer account for some 70–80% of the tissue volume of the seed coat. These parenchyma cells are likely to be involved in nutrient transfer, as they appear to be symplasmically connected with the sieve elements in the vascular bundles (van Dongen *et al.*, 2003).

Three other TIP1 members are known that are expressed in roots, stems, and leaves as well as in developing seeds where they localize to tissues involved in nutrient transfer. Thus,  $\gamma$ TIP1 (= TIP1-1) from *Arabidopsis* is expressed in the seed coat (Ludevid *et al.*, 1992), ZmTIP1 (= ZmTIP1-1) from maize in pedicel parenchyma cells (see Figure 4 in Barrieu *et al.*, 1998), and MIPFG, a TIP1 member from Norway spruce (*Picea abies*) in tapetal-like cells around gametophytes and cells of the future seed coat (Hakman and Oliviusson, 2002).

Transcripts of TIP1-members have never been detected in dry seeds (Ludevid *et al.*, 1992; Gao *et al.*, 1999; Suga *et al.*, 2001). In the cotyledons of the pea seed hybridization signals of the *PsTIP1-1* probe disappeared during seed fill, were undetectable in the dry seed, and reappeared during germination (Figures 2 and 3). Interestingly, the reverse has been found for  $\alpha$ -TIPs (TIP3 members), which accumulate late in seed maturation and disappear during germination and seedling growth (for references, see Maurel *et al.*, 1997).

#### *An aquaglyceroporin is specifically expressed in the seed coat*

We also identified a NIP that seems to be expressed specifically in the seed coat. NIPs are named after Nodulin-26, an abundant intrinsic protein in the peribacteroid membrane of the soybean nodule, and the most extensively studied member of the subfamily (Dean *et al.*, 1999 and references therein). *Nodulin 26* as well as *LIMP2* from the legume *Lotus japonicus* (Guenther and Roberts, 2000) seem to be expressed exclusively in nodules. It may be noted that the peribacteroid membrane of nodule symbiosomes (Udvardi and Day, 1997) and the plasma membrane of the seed coat parenchyma cells have very similar functions, since both membranes have to accommodate the efflux of nutrients from the cytoplasm of the plant cell. We have identified an aquaglyceroporin along with four other MIPs in the seed coat. From a recent study (Wienkoop and Saalbach, 2003) it can be deduced that the peribacteroid membrane of *L. japonicus* harbors an aquaglyceroporin and at least four other MIPs that can be identified as members of the PIP1, PIP2, TIP1 and TIP2 subgroup, respectively.

*PsNIP-1* is another example of a NIP expressed in a reproductive organ. In maize, *ZmNIP1-1* and *ZmNIP3-1* have the highest expression levels in reproductive tissues (Chaumont *et al.*, 2001). Furthermore,

a gene coding for a NIP has been cloned from the pollen of *Nicotiana glauca* (accession number P49173) and another one from rice anthers (Liu *et al.*, 1994), whereas *PtNIP1;1* from loblolly pine (*Pinus taeda*) appeared to be expressed in young, precotyledonary zygotic and somatic embryos, preferentially in the suspensor (Ciavatta *et al.*, 2001).

*PsNIP-1* expressed in *Xenopus* oocytes clearly enhanced the permeability of the oocyte membrane for water and glycerol (Figures 4 and 5). Five other NIPs have been also characterized as channels for water and glycerol (Nodulin-26, Dean *et al.*, 1999; LIMP2, Guenther and Roberts, 2000; AtNLM1 and AtNLM2, Weig and Jakob, 2000; PtNIP1;1, Ciavatta *et al.*, 2001). In GlpF glycerol is transported through an amphipathic channel. The selectivity filter in this channel is lined by four residues, Trp-48, Phe-200, Arg-206, and Gly-199, which are crucial for the selective transport of glycerol (Fu *et al.*, 2000; Sui *et al.*, 2001). Alignment of several glycerol-transporting MIPs showed that Trp-48 in GlpF is conserved in Fsp1, the glycerol facilitator from *Saccharomyces cerevisiae*, whereas it is replaced by phenylalanine in mammalian aquaglyceroporins such as AQP3. The tryptophan side-chain is also present in *PsNIP-1* and all other members of the NIP subfamily that have been experimentally characterized as aquaglyceroporins (Table 1). The importance of the tryptophan residue for glycerol permeation was recently demonstrated for LIMP2 (Wallace *et al.*, 2002). Some NIPs, however, do not contain this tryptophan residue (not shown), and their permeation characteristics have yet to be determined.

Froger *et al.* (1998) pinpointed five positions (P<sub>1</sub>–P<sub>5</sub>) to distinguish between aquaporins and glycerol facilitators. The residues at P<sub>4</sub>P<sub>5</sub> have been found determining factors for the selectivity of water and glycerol transport in the insect aquaporin AQPcic (Lagrée *et al.*, 1999), but not in the aquaporin AqpZ and the glycerol facilitator GlpF from *E. coli* (Borgnia and Agre, 2001). Neither do the amino acid residues at P<sub>2</sub> and P<sub>3</sub> bear general relevance for the selective transport of glycerol and water because all NIPs experimentally characterized as aquaglyceroporins have a serine (or threonine) and alanine at these positions, which has been considered typical of aquaporins. Note that all glycerol-transporting NIPs have amino acid residues at only two out of the five positions (P<sub>1</sub> and P<sub>5</sub>) that Froger *et al.* (1998) regarded as typical of glycerol facilitators (Table 1).

Glycerol transport has been also reported for PIP1 members from tobacco (NtAQP1) and *Samanea saman* (SsAQP1) and for a TIP4 member from tobacco (Biela *et al.*, 1999; Gerbeau *et al.* 1999; Moshelion *et al.*, 2002). This is quite remarkable because in these three MIPs all residues at Froger's positions are typical of aquaporins and, more importantly, the residues predicted to line the selectivity filter in NtAQP1 and SsAQP1 are identical to those in the orthodox PIP2 aquaporins (exemplified in Table 1 for PsPIP2-1).

#### *Solute transport by plant MIPs?*

On the basis of circumstantial evidence it has been suggested repeatedly that plant MIPs may also function as transport channels for solutes. Nodulin 26 has been proposed as a transport channel for malate, which is the major organic nutrient for bacteroids in root nodules (Ouyang *et al.*, 1991). Furthermore, some PIP1 members have been suggested to function as solute transporters. Expression patterns of *TRG-31* promoter::*GUS* constructs in transgenic tobacco lead to the idea that TRG-31 (which is identical to PsPIP1-1) could be involved in the transport of assimilates (Jones and Mullet, 1995), and more recently it was suggested that the ripening-associated membrane protein from tomato, TRAMP, which also belongs to the PIP1 group, might be involved in the transport of sugars and/or organic acids (Chen *et al.*, 2001). Finally, the complementation of yeast mutants deficient in phosphate uptake by PIPs from *Solanum tuberosum* hinted at a possible permeability for phosphate (Leggewie *et al.*, 1995).

From our own experiments and those reported by others (Opperman *et al.*, 1994; Chaumont *et al.*, 2000) it should be concluded, however, that plant MIPs expressed in *Xenopus* oocytes have shown no permeability for any of a great diversity of solutes including sugars, amino acids and ions. The transport of glycerol by certain plant MIPs seems now very well documented (see above), and transport of urea by a TIP4 member (Gerbeau *et al.*, 1999) and of boric acid by a PIP1 homologue (Dordas *et al.*, 2000) has been reported. But, to the best of our knowledge, there is at present no direct evidence of a plant MIP that serves as a transport channel for any other solute.

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