

Heteromerization of PIP aquaporins affects their intrinsic permeability

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The plant aquaporin plasma membrane intrinsic proteins (PIP) subfamily represents one of the main gateways for water exchange at the plasma membrane (PM). A fraction of this subfamily, known as PIP1, does not reach the PM unless they are coexpressed with a PIP2 aquaporin. Although ubiquitous and abundantly expressed, the role and properties of PIP1 aquaporins have therefore remained masked. Here, we unravel how FaPIP1;1, a fruit-specific PIP1 aquaporin from *Fragaria x ananassa*, contributes to the modulation of membrane water permeability (P_f) and pH aquaporin regulation. Our approach was to combine an experimental and mathematical model design to test its activity without affecting its trafficking dynamics. We demonstrate that FaPIP1;1 has a high water channel activity when coexpressed as well as how PIP1–PIP2 affects gating sensitivity in terms of cytosolic acidification. PIP1–PIP2 random heterotetramerization not only allows FaPIP1;1 to arrive at the PM but also produces an enhancement of FaPIP2;1 activity. In this context, we propose that FaPIP1;1 is a key participant in the regulation of water movement across the membranes of cells expressing both aquaporins.

The plasma membrane (PM) is the first barrier that limits water exchange in plant cells. The rate of its water transport capacity is mainly associated with aquaporins. Among the seven aquaporin subfamilies described in the plant kingdom, only plasma membrane intrinsic proteins (PIP) and some members of the nodulin-26-like intrinsic proteins (NIP) and X intrinsic proteins (XIP) subfamilies have been shown to be preferentially localized at the PM (1, 2). Of these, PIP aquaporins appear to have a large role in controlling membrane water permeability, whereas NIP and XIP have been mainly described as solute transporters (2–4). Plant PIP aquaporins represent a conserved subfamily that has been historically divided into two subgroups due to their differences in primary structure, PIP1 and PIP2. Interestingly, PIP aquaporins compose ~40% of the total aquaporin set, and the PIP1 and PIP2 ratio among different species is relatively constant (5–12). Fig. S1 shows the distribution of all aquaporin genes present in plants whose genome has been completely sequenced and analyzed. Antisense inhibition experiments on *Arabidopsis thaliana* PIP1 and PIP2 have suggested that the two subgroups of aquaporins contribute to root or leaf hydraulic conductivity in the same way (13). In several plant species, members of the PIP1 and PIP2 subgroups were shown to be coexpressed in the same cell type (14–17).

Although PIP1 are as ubiquitous as PIP2, the functional properties of each type of channel are different. PIP2 are very well described as a homotetramer with high water transport activity (18, 19) and a gating mechanism unequivocally associated with specific and conserved amino acid motifs triggered by cytosolic acidification (20–22), phosphorylation (23, 24), or divalent cation concentration (22). In contrast, PIP1 have shown complex heterogeneity in water and solute transport and post-translational regulation. Many reports show that some PIP1 are nonfunctional in regard to water transport (6, 25), whereas other PIP1 act as low-efficiency water channels (26–28), and a minority

group shows activity comparable to that of PIP2 (20, 29) or, in contrast, serves as solute channels (25, 30).

In addition to their transport properties, many PIP1 show membrane relocation as a regulatory mechanism, a feature that clearly distinguishes them from any PIP2. These PIP1 fail to reach the PM when expressed alone, but they can succeed if they are coexpressed with PIP2. It has been proposed that this process is a consequence of a physical interaction between PIP1 and PIP2, as reported in both homologous (31) and heterologous systems (14, 32). Although there are some PIP1 with the ability to reach the PM on their own (20, 27, 29), this PIP1–PIP2 interaction seems to be present for several pairs of PIP among different species with functional consequences (14, 21, 28, 33–35).

Although the molecular basis of this interaction is still not clear, some data support a model in which the aquaporins of the two subgroups physically interact—very likely by heterooligomerization—to facilitate PIP1 trafficking (31). Recently, it was shown that the first extracellular loop of PIP2 (loop A in BvPIP2;1) could be relevant to the formation of heterotetramers with PIP1 (32). The modification of cytosolic pH sensing, reflected by a shift in the EC₅₀ of oocytes coexpressing BvPIP2;2 and BvPIP1;1 compared with BvPIP2;2 expressed alone, favors the heterooligomerization hypothesis (21).

The aim of this work is to contribute to the understating of the PIP1 and PIP2 interaction and to elucidate the functional properties of PIP1 and the role it plays in defining overall membrane permeability.

Significance

Aquaporins are known for their capacity to increase trans-cellular water exchange. In plants, a highly conserved group known as plasma membrane intrinsic proteins (PIP) affects the adjustment of not only membrane water permeability but also overall plant hydraulic conductivity. An experimental design combined with a mathematical modeling approach allowed us to explore the interplay of channel gating, membrane translocation, and channel stoichiometric arrangement of a pair of PIP1 and PIP2 aquaporins. We dissect the individual contribution of each PIP, showing that (i) PIP1 has a high water transport capacity when coexpressed with PIP2, (ii) PIP2 water permeability is enhanced if it physically interacts with PIP1, and (iii) the PIP1–PIP2 interaction results in the formation of heterotetramers with random stoichiometric arrangement.

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Results

FaPIP1;1 Cannot Reach the PM Unless It Is Coexpressed with FaPIP2;1.

The localization of FaPIP1;1-EYFP and FaPIP2;1-EYFP was studied by means of confocal microscopy in *Xenopus* oocytes expressing the tagged aquaporins. Fig. 1 shows that whereas FaPIP2;1-EYFP is localized mostly at the oocyte PM, FaPIP1;1-EYFP is restricted to internal structures. However, when FaPIP1;1-EYFP is coexpressed with FaPIP2;1, the fluorescence signal is mainly visualized at the limit of the cell, indicating its relocation to the PM.

Water transport assays show that FaPIP1;1-EYFP displayed no significant increase in PM osmotic water permeability in accordance with FaPIP1;1-EYFP localization in the interior of the cell (Fig. S2). Oocytes expressing FaPIP2;1-EYFP showed high water transport activity ($P_f = 112 \pm 8 \cdot 10^{-4} \text{ cm} \cdot \text{s}^{-1}$). Coexpression of FaPIP1;1-EYFP with FaPIP2;1 and coexpression of FaPIP1;1 with FaPIP2;1-EYFP present high P_f values, $250 \pm 8 \cdot 10^{-4} \text{ cm} \cdot \text{s}^{-1}$ and $250 \pm 13 \cdot 10^{-4} \text{ cm} \cdot \text{s}^{-1}$, respectively; in both cases, the values are almost 2.2-fold higher than the P_f with FaPIP2;1-EYFP expressed alone. These results are consistent with previously reported values of P_f for non-EYFP-tagged FaPIP coexpression (36).

FaPIP2;1–FaPIP1;1 Coexpression Has a Different pH Inhibition Response than FaPIP2;1 Expressed Alone.

In a previous study, we showed that cytosolic acidification (pH 6.0) of oocytes expressing FaPIP2;1 or coexpressing FaPIP1;1–FaPIP2;1 triggers a reduction in P_f (36). Here, we analyze the pattern of this inhibition of P_f vs. $[\text{H}^+]$ dose–response curves (Fig. 2), which is sigmoidal for both systems, indicating allosteric behavior. Both maximal P_f and EC_{50} are different in each case. The EC_{50} value shifts to 6.60 ± 0.02 when both aquaporins are coexpressed, showing a different gating sensitivity to proton concentration (Fig. 2, *Inset*).

FaPIP2;1 Mutant Reveals FaPIP1;1 Water Transport Activity. Because FaPIP1;1 is not localized at the PM unless it is coexpressed with FaPIP2;1, we constructed a FaPIP2;1 mutant to characterize

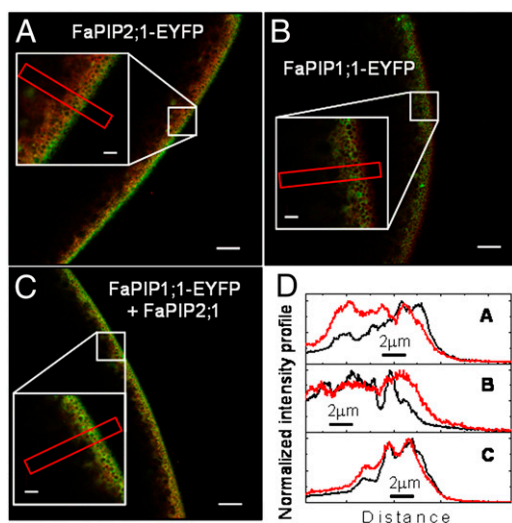


Fig. 1. Subcellular localization of FaPIP2;1-EYFP and FaPIP1;1-EYFP. (A–C) Radial (x – z) confocal images of *Xenopus laevis* oocytes expressing FaPIP2;1-EYFP (A) (green) and FaPIP1;1-EYFP (B and C) (green), previously injected with TMR-Dextran (red). The oocyte surface is near the right of each image frame, and the interior of the oocyte is to the left. *Insets* show an enlargement of the indicated square section. (A) FaPIP2;1-EYFP localizes mostly in the oocyte PM; (B) FaPIP1;1-EYFP expressed alone is restricted to internal structures. (C) When FaPIP1;1-EYFP is coexpressed with FaPIP2;1, the fluorescence is mainly visualized at the limit of the cell, indicating the relocation of FaPIP1;1-EYFP to PM. (D) Normalized intensity profile of selected areas in A–C. The black line represents EYFP intensity levels, and the red line shows the TMR-Dextran intensity levels.

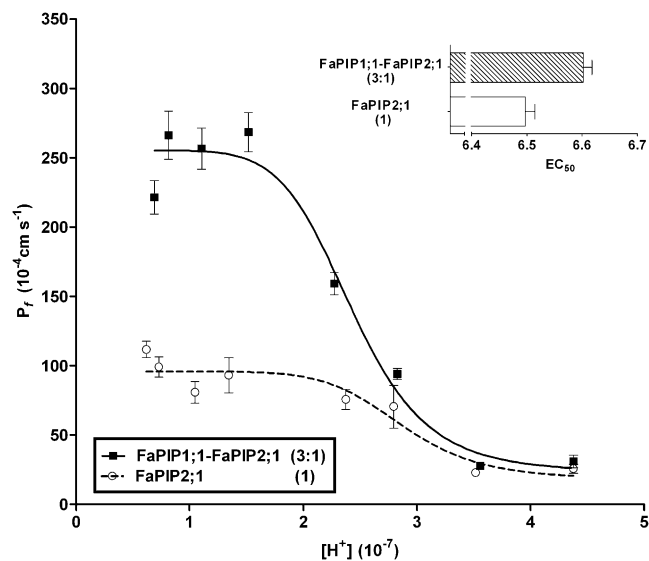


Fig. 2. P_f inhibitory response triggered by cytosolic acidification for FaPIP1;1–FaPIP2;1 (at a 3:1 cRNA mass ratio) and for FaPIP2;1 expressed alone. Shown is P_f vs. $[\text{H}^+]$ inhibitory response profile for oocytes injected with FaPIP2;1 cRNA and oocytes injected with FaPIP1;1 and FaPIP2;1 cRNA in a 3:1 mass ratio. The amount of cRNA is represented in parentheses with an arbitrary unit of measure where 1 is equivalent to 1.25 ng of cRNA. The data points are representative values obtained from the same batch of oocytes (mean $P_f \pm \text{SEM}$). The data were fitted to an allosteric sigmoidal dose–response curve. (*Inset*) For each K_i ($[\text{H}^+]$ at which half of the maximum effect occurs), an EC_{50} value was calculated using $\text{EC}_{50} = -\log K_i$. The mean $\text{EC}_{50} \pm \text{SEM}$, $n = 3$ –5) was calculated for each treatment.

the FaPIP1;1 water transport activity. Our objective was to generate a nonfunctional FaPIP2;1 mutant while preserving its capacity to interact with FaPIP1;1. To do so, we followed a strategy already applied to AQP1 (37), replacing a highly conserved residue (Asn228) with Asp. Fig. S3 shows that the mutant FaPIP2;1N228D-EYFP is unable to reach the PM when expressed alone, but when FaPIP1;1 and FaPIP2;1N228D-EYFP or FaPIP1;1-EYFP and FaPIP2;1N228D are coexpressed, both fluorescently tagged aquaporins reach the PM. These results are consistent with the P_f measurements (Fig. 3 and Fig. S3). When both proteins are coexpressed, the P_f of the oocyte PM increases (Fig. 3). However, this P_f value cannot be a priori assigned to FaPIP1;1 because it has not yet been proved that the mutant is inactive for water transport. Testing the latter hypothesis is not straightforward because FaPIP2;1N228D requires functional interaction with FaPIP1;1 to reach the PM. Therefore, water transport assays were performed in oocytes after the injection of variable complementary RNA (cRNA) mass ratios of FaPIP1;1–FaPIP2;1 or FaPIP1;1–FaPIP2;1N228D. Fig. 4 shows that when FaPIP1;1–FaPIP2;1 are coexpressed, the P_f increase correlates well with the amount of cRNA of FaPIP2;1 injected ($P < 0.05$). On the other hand, when the amount of FaPIP1;1 cRNA injected was increased, there were no significant differences observed between the P_f of oocytes injected with cRNA mass ratios of 1:1, 1:2, and 1:3 ($P < 0.05$). When analyzing FaPIP1;1–FaPIP2;1N228D coexpression, the amount of increase in P_f correlated with the amount of cRNA of FaPIP1;1 injected ($P < 0.05$) but remained constant when the amount of FaPIP2;1N228D cRNA increased ($P < 0.05$). This result suggests that FaPIP2;1N228D is unable to transport water; moreover, the final P_f could reflect the water transport activity of FaPIP1;1.

Mathematical Models Dissect the Intrinsic Contribution of Each Coexpressed Aquaporin to the Total P_f . To determine the real contributions of both FaPIP1;1 and FaPIP2;1N228D in terms of water transport, a set of mathematical models was developed (Fig. 5). The difference between the models lies in the composition of

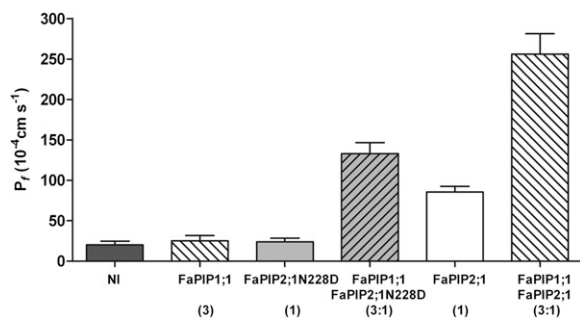


Fig. 3. Osmotic water permeability of FaPIP2;1, FaPIP1;1, and FaPIP2;1N228D and coexpression in *Xenopus laevis* oocytes. Shown are P_f measurements of oocytes injected with FaPIP1;1, FaPIP2;1, or FaPIP2;1N228D cRNA and coinjections. The amount of cRNA is represented in parentheses with an arbitrary unit of measure where 1 is equivalent to 1.25 ng of cRNA. Noninjected (NI) oocytes were used as controls. P_f values are represented as $P_f \pm$ SEM, $n = 9$ –12.

tetramers and the intrinsic permeabilities of each FaPIP (p_f). For instance, models 1A and 1B assumed that a random aggregation of homodimers forms the tetramer, whereas models 2A and 2B assumed that a random aggregation of monomers forms the tetramer. The classification as A or B models depends on the p_f of FaPIP2;1. The A models assume that the FaPIP2;1 p_f is not necessarily equal in a homotetramer and in a heterotetramer configuration, whereas the B models assume the same p_f for FaPIP2;1 independent of the type of tetramer.

The first round of selection was between model 1 and model 2. Because model 2 fits the data better than model 1 for FaPIP1;1–FaPIP2;1N228D coexpression, the latter was discarded. Model 2 was then preselected, and we proceeded to check the Akaike information criterion (AIC) values of models 2A and 2B for FaPIP1;1–FaPIP2;1 coexpression. Model 2A presents a better fit (AIC = -19.99) in comparison with model 2B (AIC = -7.59). Because only model 2A fitted the data well, the hypothesis associated with this model (i.e., random monomer heterotetramerization and a different FaPIP2;1 p_f for homo- and heterotetramer configuration) seems to be the most plausible.

The fitting parameters using model 2A for FaPIP1;1–FaPIP2;1 coexpression and for FaPIP1;1–FaPIP2;1N228D coexpression are shown in Fig. 6. The parameter values of three independent FaPIP1;1 and FaPIP2;1 coexpression experiments and their fittings indicate that the FaPIP1;1 and FaPIP2;1 intrinsic permeabilities within the heterotetramer double the intrinsic permeability of FaPIP2;1 when a homotetramer is formed. The fitting parameter that corresponds to the FaPIP2;1N228D relative intrinsic permeability ($p_{f,rel-FaPIP2;1N228D}$) was zero in all three independent adjustments of FaPIP1;1–FaPIP2;1N228D coexpression experiments, reinforcing the hypothesis that the FaPIP2;1N228D mutant lacks water transport capacity and therefore contributes only to the translocation of FaPIP1;1 to the PM. Additionally, the fitting parameter that corresponded to the FaPIP1;1 relative intrinsic permeability ($p_{f,rel-FaPIP1;1}$) in all three independent adjustments was similar for both coexpression experiments.

FaPIP1;1 Is Important in pH Sensing. According to our model, FaPIP2;1N228D does not transport water; thus, the observed water permeability when it is coexpressed with FaPIP1;1 can be attributed entirely to FaPIP1;1. With these data, it is possible to explore FaPIP1;1 pH sensitivity and its contribution to the sensitivity of the whole heterotetramer configuration. We analyzed the pH inhibition of the oocyte plasma membrane P_f when different mass ratios of cRNA of FaPIP1;1 and FaPIP2;1N228D or FaPIP2;1 were coinjected. Fig. 7 shows that the EC_{50} value remains constant regardless of the cRNA coinjection performed, differing only when FaPIP2;1 is expressed alone ($P < 0.05$).

Discussion

The functional interaction reported for certain PIP1–PIP2 pairs (31, 35) provides an additional regulatory mechanism to adjust water exchanges at the plant PM level. Determining the specific PIP1 contribution to the PM P_f and pH sensitivity when coexpressed with PIP2 was the goal of this work.

We explored FaPIP1;1 as a channel that requires coexpression with FaPIP2;1 to be translocated to PM (Fig. 1). To elucidate its functional activity, we combined coexpression experiments of FaPIP1;1 with FaPIP2;1 or FaPIP2;1N228D with a mathematical approach. The four models developed assumed that the total P_f observed would represent the water transport activity of each aquaporin at the oocyte PM plus the water directly crossing the lipid bilayer. Because it is well established that the MIP quaternary structure is tetrameric (18, 19, 38, 39), all models assumed that all expressed aquaporins form tetrameric structures. Moreover, recent discoveries indicate that PIP interaction occurs via heterotetramerization (25, 32).

To build our models, we had to consider channel arrangement within the tetramer. Studies of a variety of channels and receptors have shown that subunit arrangement in the conformation of heterooligomers varies depending on which membrane protein is being analyzed. It has been proposed that individual proteins might assemble with a random arrangement that is dependent on their abundance (40–43), with a random arrangement of preformed homomeric dimers (44), and some channels even assemble as heteromeric structures with fixed stoichiometry (45, 46). With respect to aquaporins, it was reported that the AQP4 isoforms M1 and M23 seem to assemble as heterotetramers with a random arrangement (47). In regard to PIP aquaporins, it was previously suggested that a disulfide bridge between the cysteines of two monomers could be important in determining the heterotetramer composition, which would assemble through a dimer-dimer association in *Zea mays* PIP1 and PIP2 (48). Here we considered that (i) random heterooligomerization of two homodimers occurs following a binomial distribution (models 1A and 1B) or (ii) random heterooligomerization of monomers occurs following a binomial distribution (models 2A and 2B).

Our results show that model 2A best explained the experimental results. Models 1A and 1B were discarded because they could not explain the P_f obtained within different coexpression ratios of FaPIP1;1 and FaPIP2;1N228D. These models assume that the interaction of FaPIP2;1N228D dimers with FaPIP1;1 dimers would form heterotetramers with a 2:2 stoichiometry capable of reaching the PM and as a consequence predict that increases in the amount of FaPIP1;1 or FaPIP2;1N228D cRNA would have the same effect as increasing the quantity of 2:2 heterotetramers formed, producing a similar P_f enhancement.

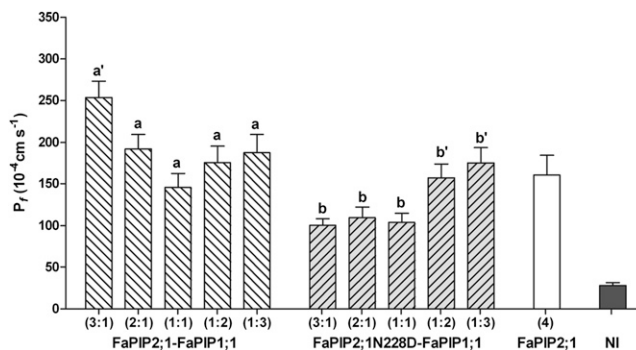


Fig. 4. P_f measurements of oocytes injected with different ratios of the coexpression pairs FaPIP2;1–FaPIP1;1 and FaPIP2;1N228D–FaPIP1;1. Shown are P_f measurements of oocytes injected with different ratios of cRNA. The amount of cRNA is represented in parentheses with an arbitrary unit of measure where 1 is equivalent to 0.25 ng of cRNA. a' and b' are significantly different from a and b, respectively ($P < 0.05$).

	p_f Assumption	$K_m = 1$ (plasma membrane localization)	$K_m = 0$ (internal structures)	Fitting parameters	AIC
1 HOMODIMER + RANDOM TETRAMERIZATION	1A p_f FaPIP2;1 homotetramer is different from p_f FaPIP2;1 heterotetramer			\circ/\circ \bullet/\circ	-18.54
				\bullet/\circ \bullet/\circ	-3.42
	1B p_f FaPIP2;1 homotetramer is equivalent to p_f FaPIP2;1 heterotetramer			\bullet/\circ \bullet/\circ	-20.54
				\bullet/\circ \bullet/\circ	-3.42
2 MONOMER + RANDOM TETRAMERIZATION	2A p_f FaPIP2;1 homotetramer is different from p_f FaPIP2;1 heterotetramer			\circ/\circ \bullet/\circ	-19.99
				\bullet/\circ \bullet/\circ	-17.97
	2B p_f FaPIP2;1 homotetramer is equivalent to p_f FaPIP2;1 heterotetramer			\bullet/\circ \bullet/\circ	-7.59
				\bullet/\circ \bullet/\circ	-17.97

Fig. 5. Schematic representation of the mathematical models including their assumptions and fitting parameters. Four different models are presented, each with distinctive assumptions regarding each parameter. The second and third columns indicate which tetramers may form within each model and their localization coefficient (k_m). The fitting parameters are schematized using circles to represent the intrinsic permeability (p_f) of each aquaporin. In model A, a distinction was made for FaPIP2;1 intrinsic permeability when it is part of a heterotetramer, and open dotted circles represent the FaPIP2;1 intrinsic permeability when it is part of a homotetramer. The Akaike Information Criterion (AIC) value presented in the sixth column was used to analyze the goodness of fit and the complexity (number of fitting parameters) of the different models (56).

In contrast, models 2A and 2B could explain our results satisfactorily. Models 2A and 2B discriminate between FaPIP2;1 with different or the same intrinsic permeability when forming a homotetramer or a heterotetramer, respectively. A better fit was obtained for model 2A, which confirmed that FaPIP2;1N228D does not transport water (Fig. 6). This lack of activity is in accordance with *in silico* studies of the conserved NPA motif (49) and constitutes a unique report of a PIP that lacks the ability to transport water due to a mutation of one of the NPA sites. Finally, because FaPIP2;1N228D is a nonfunctional aquaporin, the coexpression of this mutant with FaPIP1;1 reveals the activity of FaPIP1;1 as a water channel.

According to our results, FaPIP2;1 or FaPIP2;1N228D and FaPIP1;1 are organized as heterotetramers with both types of monomers participating in the tetramer randomly, depending on their relative amount. Moreover, the most plausible interpretation of these results is that FaPIP2;1 has a different p_f depending on its tetramer configuration. If FaPIP1;1 and FaPIP2;1 form a heterotetramer, our results show a p_f that is twice the p_f of FaPIP2;1 when it is in the homotetramer configuration (Fig. 6). These data are important because they could be interpreted as indicating a dual role for FaPIP2;1, not only acting to assist FaPIP1;1 in translocating to the PM but also augmenting its own p_f when coexpressed with PIP1. Moreover, because FaPIP1;1 and FaPIP2;1 have approximately the same p_f in the heterotetramer configuration, each heterotetramer would be expected to have equal water channel activity regardless of its stoichiometry.

PIP gating by cytosolic acidification has been explored exhaustively in PIP2 (20–22, 36) but analyzed only with AtPIP1;2, a PIP1 that is able to reach the PM alone in *Xenopus* oocytes (20). Therefore, it was assumed that PIP2 and PIP1 share the same mechanism of triggering an on-off response under acidification. Here, we show that when FaPIP1;1 and FaPIP2;1 are coexpressed, the interaction promotes a change in pH sensing as shown by a modification of the observed EC_{50} value. Similar results were previously obtained for BvPIPs (21, 32). We demonstrate that coexpressing FaPIP1;1 with FaPIP2;1 or FaPIP2;1N228D produced different maximal P_f . Regardless of the proportion of the channels, the EC_{50} remained the same, supporting the idea of a characteristic pH sensitivity for the heterotetramers different from that corresponding to the FaPIP2;1 homotetramer (Fig. 7). When PIP2 is expressed in the absence of PIP1, a lower EC_{50} implies a lower probability of closed water channels in physiological

conditions. Clearly, FaPIP1;1 has an important impact on pH sensing by shifting the EC_{50} when expressed at the PM.

As proved by recent discoveries of the subcellular dynamics of plant AQPs, cellular trafficking and its effect on AQP function are important regulatory mechanisms defining plant cell water permeability (50, 51). Here, we demonstrated that PIP1 aquaporins retained intracellularly are active water channels that are located at the PM when coexpressed with PIP2. The combination of channel relocalization and enhancement of their p_f broadens not only the landscape of aquaporin plasticity in adjusting cell-to-cell pathways but also the understanding of the role of heteromerization as a regulatory mechanism.

Materials and Methods

Plasmid Construction and Site-Directed Mutagenesis. To generate the EYFP-tagged vectors, FaPIP2;1 and FaPIP1;1 cDNAs were amplified by PCR, using the pT7Ts-FaPIP2;1 and pT7Ts-FaPIP1;1 plasmids as templates, respectively. Using specific primers, a SacII site was incorporated at the 5' end of both constructs and a BglII site at the 3' end. Then, the two PCR-amplified fragments were subcloned into the SacII and BglII sites of a pT7Ts-BvPIP2;2-EYFP

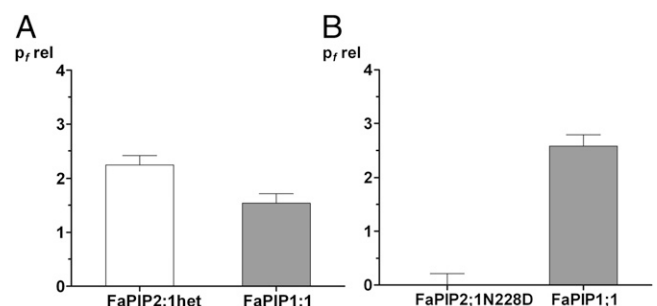


Fig. 6. Model 2A fitting parameters: relative intrinsic permeability values for each aquaporin. Fitting was performed with three independent experiments that yielded similar results. A representative independent experiment is shown. Because the fitting parameters are the intrinsic permeability of each aquaporin relative to FaPIP2;1 intrinsic permeability when it is part of a homotetramer, the FaPIP2;1 relative intrinsic permeability when it is part of a homotetramer would be equal to 1. (A) Fitting parameter values are reported as $p_{f,rel} \pm SEM$ for the FaPIP1;1–FaPIP2;1 coexpression results. (B) Fitting parameter values are reported as $p_{f,rel} \pm SEM$ for the FaPIP1;1–FaPIP2;1N228D coexpression results.

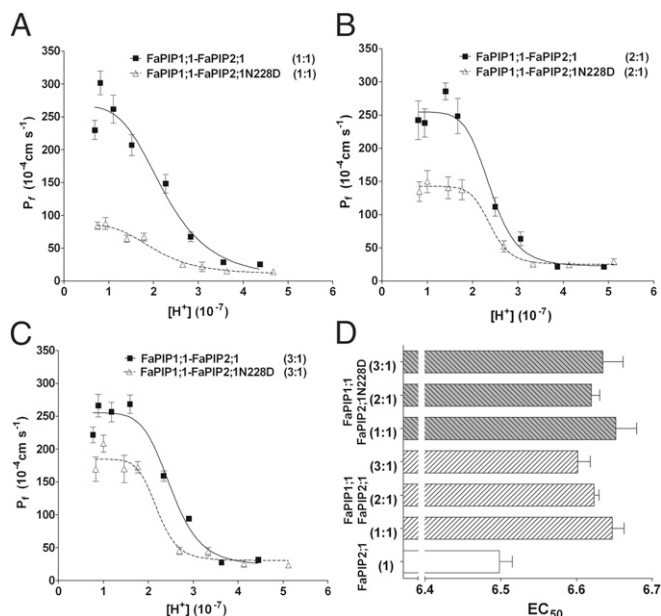


Fig. 7. P_f inhibitory response triggered by cytosolic acidification with FaPIP1;1-FaPIP2;1 and FaPIP1;1-FaPIP2;1N228D coexpression. (A–C) P_f vs. $[H^+]$ inhibitory response profile for oocytes coexpressed with FaPIP1;1 and FaPIP2;1 cRNA and oocytes coexpressed with FaPIP1;1 and FaPIP2;1N228D cRNA at 1:1 (A), 2:1 (B), and 3:1 (C) mass ratios. The amount of cRNA is represented in parentheses with an arbitrary unit of measure where 1 is equivalent to 1.25 ng of cRNA. The data points are representative values obtained from the same batch of oocytes (mean $P_f \pm$ SEM). Data were fitted to an allosteric sigmoidal dose-response curve. (D) EC_{50} values reported as the average of three to five independent experiments ($EC_{50} \pm$ SEM, $n = 3–5$) for each treatment and compared with the FaPIP2;1 EC_{50} shown in Fig. 2. The FaPIP2;1 EC_{50} is significantly different from the EC_{50} values presented in coexpression experiments ($P < 0.05$).

vector, thus exchanging BvPIP2;2 for FaPIP2;1 or FaPIP1;1. The products were pT7Ts-FaPIP2;1-EYFP and pT7Ts-FaPIP1;1-EYFP vectors where the EYFP sequence is 10 codons downstream of the aquaporin sequence.

Mutated cDNA encoding FaPIP2;1N228D and FaPIP2;1N228D-EYFP was obtained by site-directed mutagenesis (QuikChange; Stratagene), following the manufacturer's recommendations, using custom-made oligonucleotide primers (Eurofins MWG Operon).

All of the constructs were confirmed by DNA sequencing (Macrogen).

cRNA in Vitro Synthesis. Capped cRNAs encoding FaPIP1;1, FaPIP2;1, FaPIP2;1N228D, FaPIP1;1-EYFP, FaPIP2;1-EYFP, and FaPIP2;1N228D-EYFP were synthesized in vitro, using the mMESSAGE mMACHINE T7 High Yield Capped RNA Transcription Kit (Ambion). The pT7Ts-derived vector carrying the corresponding sequence as template was linearized by XbaI. The synthesized products were suspended at a final concentration of $0.1 \mu\text{g} \cdot \mu\text{L}^{-1}$ in RNase-free water supplemented with Recombinant RNasin (Ribonuclease inhibitor; Promega). The cRNA was quantified by fluorescence, using a Quant-iT RNA Assay Kit (Invitrogen). At least two independent cRNA syntheses were assayed.

Oocyte Transport Studies. Defolliculated *Xenopus* oocytes were injected with different masses of cRNA in a final volume of 50 nL, using an automatic injector (Drummond Scientific). Injected oocytes were incubated for 72 h at 18 °C in ND96 medium supplemented with $1 \mu\text{g} \cdot \text{mL}^{-1}$ gentamycin sulfate (21). Osmotic water permeability (P_f) was determined by measuring the rate of oocyte swelling induced by a hypo-osmotic shock of $160 \text{ mOsm} \cdot \text{kg}^{-1}$. Changes in cell volume were video monitored, and the P_f was calculated according to previous reports (52, 53). All osmolarities were determined using a vapor pressure osmometer (5520C; Wescor). Noninjected oocytes were used as a negative control (36). The results from experiments performed with different oocyte batches were not pooled; therefore, all of the experiments shown in this report are representative of at least three different experiments.

Confocal Microscopy. Confocal fluorescence microscopy was used to determine the localization of the respective PIP subgroups tagged with EYFP. We used

tetramethylrhodamine (TMR) dextran (molecular weight 10,000; Invitrogen-Molecular Probes), i.e., an unconjugated nonspecific fluorochrome marker (54), as a marker of the interior of the oocyte. Three to 4 d after cRNA injection and 40 min before imaging, oocytes were microinjected with 50 nL of a 33 mM aqueous solution of TMR-dextran. If the tagged aquaporin fluorescence localized in the same area of the marker fluorescence, the aquaporin is retained in the interior of the cell. However, if the tagged aquaporin fluorescence appeared on the limit of the cell supports, they were localized in PM. Intensity profiles were calculated averaging 5 consecutive pixels along the direction specified in each figure. Fluorescence images of EYFP distribution, together with TMR, were obtained with a Fluoview1000 spectral confocal scanning microscope (Spectral FV1000 Olympus, Japan), using a 60X UPLSAPO oil immersion objective lens with a numerical aperture of 1.35. To avoid crosstalk, images were recorded line by line in a sequential order. EYFP and TMR were excited using the 488 nm line of the argon laser and the 543 nm He-Ne, respectively, and the emitted fluorescence was detected in the 500–530 nm and 570–670 nm range. Autofluorescence (monitored in control oocytes) was negligible in comparison with cells expressing fluorescent PIP. We analyzed 3–5 oocytes from each of at least 6 donor frogs.

Aquaporin Inhibition by pH. The oocyte cytosolic pH was modified following a previously described protocol (20). Briefly, oocytes were preincubated in different NaAc solutions with distinct final pH. For the 5.8–6.8 pH interval, the NaAc solutions were as follows: 50 mM NaAc, 20 mM Mes, supplemented with mannitol 1 M to adjust the osmolarity to $\sim 200 \text{ mOsmol} \cdot \text{kg}^{-1}$ and supplemented with NaOH to adjust to the final desired pH. For the 6.8–7.6 pH interval, the NaAc solutions were as follows: 50 mM NaAc, 20 mM HEPES, supplemented with mannitol 1 M to adjust the osmolarity to $\sim 200 \text{ mOsmol} \cdot \text{kg}^{-1}$ and supplemented with NaOH to adjust to the final desired pH. To calculate internal proton concentration ($[H^+]_{\text{int}}$), we used a calibration curve described previously (21). The P_f values for each $[H^+]_{\text{int}}$ were fitted with the sigmoidal dose-response equation presented in *SI Text* (55). The results from experiments performed with different oocyte batches were not pooled; therefore, all of the dose-response curves shown in this work are representative of at least three different curves.

Mathematical Model Analysis. To analyze the oocyte total water permeability in coexpression assays and dissect the contribution of each aquaporin to the total P_f achieved, four mathematical models were proposed.

The equations of each model are shown in *SI Text*. The mass of cRNA of each aquaporin injected was used as independent model variables. The dependent variable was $[(P_{\text{FCO}} - P_{\text{NI}})/(P_{\text{FaPIP2;1}} - P_{\text{NI}})]$, where $P_{\text{FCO}} - P_{\text{NI}}$ is the total permeability provided by the tetramers formed by coexpressed aquaporins relative to $P_{\text{FaPIP2;1}} - P_{\text{NI}}$, which represents the permeability provided by the injection of a fixed amount of cRNA of FaPIP2;1. The fitting parameters are the intrinsic permeability (p_i) of each aquaporin forming a heterotetramer relative to the intrinsic permeability of FaPIP2;1 forming a homotetramer; i.e., $p_i^{\text{rel}}_{\text{FaPIP2;1}}$, $p_i^{\text{rel}}_{\text{FaPIP1;1}}$, and $p_i^{\text{rel}}_{\text{FaPIP2;1N228D}}$.

The general assumptions of all models include the following:

- The water permeability of the lipid bilayer (P_{LB}) is equal to the water permeability shown by noninjected oocytes (P_{NI}).
- All of the synthesized aquaporins formed tetrameric structures.
- The permeability of the oocyte PM is equal to the permeability of the lipid bilayer plus the permeability of the total amount of each type of tetramer expressed in PM.
- FaPIP1;1, FaPIP2;1, and FaPIP2;1N228D are equally expressed. A translation coefficient indicating the amount of protein expression per mass of injected RNA was defined: k_e (i.e., $k_{e\text{FaPIP1;1}} = k_{e\text{FaPIP2;1}} = k_{e\text{N228D}} = k_e$). This coefficient is canceled when the permeability of each coexpression is normalized to the permeability of oocytes injected with 1 ng of FaPIP2;1 cRNA.
- A localization coefficient (k_{m}) is defined for each tetramer on the basis of confocal microscopy data. To simplify the analysis, this coefficient will be 0 or 1 depending on the tetramer localization, with 1 for the PM and 0 for other locations other than the PM.
- All heterotetramers (FaPIP1;1-FaPIP2;1 or FaPIP1;1-FaPIP2;1N228D) are assumed to target completely and equally to the PM ($k_{\text{m}} = 1$).

The particular assumptions of each model are shown in *SI Text*.

Each model was fitted to three independent experiments. Fig. 6 shows one representative experiment with the fitted parameters. The AIC was used to compare the different models (56). The AIC was calculated as

$$\text{AIC} = n \ln \text{SC} + 2p,$$

where n is the number of experimental data, SC is the residual squares sum, and p is the number of parameters. The model with the lowest AIC value was chosen.

Statistical Analysis. The results are reported in the form of means \pm SEM. Significant differences between treatments were calculated using Student's *t* test.

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