



Polyamines control of cation transport across plant membranes: implications for ion homeostasis and abiotic stress signaling

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Polyamines are unique polycationic metabolites, controlling a variety of vital functions in plants, including growth and stress responses. Over the last two decades a bulk of data was accumulated providing explicit evidence that polyamines play an essential role in regulating plant membrane transport. The most straightforward example is a blockage of the two major vacuolar cation channels, namely slow (SV) and fast (FV) activating ones, by the micromolar concentrations of polyamines. This effect is direct and fully reversible, with a potency descending in a sequence $\text{Spm}^{4+} > \text{Spd}^{3+} > \text{Put}^{2+}$. On the contrary, effects of polyamines on the plasma membrane (PM) cation and K^+ -selective channels are hardly dependent on polyamine species, display a relatively low affinity, and are likely to be indirect. Polyamines also affect vacuolar and PM H^+ pumps and Ca^{2+} pump of the PM. On the other hand, catabolization of polyamines generates H_2O_2 and other reactive oxygen species (ROS), including hydroxyl radicals. Export of polyamines to the apoplast and their oxidation there by available amine oxidases results in the induction of a novel ion conductance and confers Ca^{2+} influx across the PM. This mechanism, initially established for plant responses to pathogen attack (including a hypersensitive response), has been recently shown to mediate plant responses to a variety of abiotic stresses. In this review we summarize the effects of polyamines and their catabolites on cation transport in plants and discuss the implications of these effects for ion homeostasis, signaling, and plant adaptive responses to environment.

Keywords: cytosolic calcium, ion channels, ion pumps, plasma membrane, polyamines, reactive oxygen species, stress, vacuole

INTRODUCTION

Polyamines (PAs) are acknowledged regulators of plant growth, development, and stress responses. In the model plant *Arabidopsis*, changes in the expression of different enzymes of the PAs biosynthesis and respective levels of individual PAs are stress-specific, and these changes mediate stress tolerance (Alcázar et al., 2010); same may be true also for other plants (see below). Polyamines at physiological pH are polycations, bearing from 2 (putrescine, Put) to 4 (spermine, Spm or thersmospermine, tSpm) positive charges. Thus, PAs can stabilize membranes or nucleic acids, binding to their negative surfaces (Galston and Sawhney, 1990; Kusano et al., 2008). They can also act as a source of reactive oxygen species (ROS) but also as ROS scavengers and activators of key of antioxidant enzymes (Kusano et al., 2008; Moschou et al., 2008; Pottosin et al., 2014a). In animal cells PAs affect a variety of plasma membrane (PM) cation channels, acting primarily as pore blockers but in some cases also affecting the channel gating and/or regulation by extra- and intracellular ligands (Drouin and Hermann, 1994; Lopatin et al., 1994; Williams, 1997; Lu and Ding, 1999; Huang and Moczydlowski, 2001; Xie et al., 2005; Ahern et al., 2006). PAs effects on vacuolar channels in plants were revealed and partly reviewed (Pottosin

and Muñiz, 2002), but the accumulated experimental evidence for the PAs effects on the plant PM ion channels and pumps was not properly discussed until now. This review is aimed to fill this gap and provide a comprehensive overview on our current knowledge of PA control over cation transport across plant membranes, and its implications for ion homeostasis, signaling, and plant adaptive responses to environment.

STRESS-INDUCED CHANGES IN POLYAMINE CONTENT AND THEIR ROLE IN PLANT ADAPTIVE RESPONSES TO ENVIRONMENT

Polyamine levels are strongly modulated by literally every known abiotic factor (see Table 1, for selected examples), often reaching the millimolar level (such as for putrescine; Galston and Sawhney, 1990; Alcázar et al., 2006) under stress conditions. The current consensus is that, rather than being merely collateral effect of stress-induced metabolic changes, these changes are beneficial to plant performance upon stress (Alcázar et al., 2010; Gupta et al., 2013) and therefore represent an important component of plant adaptive mechanisms. Three major lines of evidence support this claim, which can be illustrated for salinity and drought stresses—two key abiotic stresses affecting agricultural crop production

Table 1 | Stress-induced changes in the level of free polyamines in plants.

Species	Stress	Put	Spd	Spm	References
Pine	Drought	Up	Up	Up	De Diego et al., 2013
Pepper	Drought	Up ^a	Up ^a	Up ^a	Sziderics et al., 2010
Rice	Drought	Down	Up	Up	Yang et al., 2007
Wheat	Osmotic	Up	Up	Up	Grzesiak et al., 2013
Potato	Osmotic	Down	Down	Down	Li et al., 2005
Bromus	Osmotic	Down	Down	Down	Gicquiaud et al., 2002
Cucumber	Flooding	Up	Up	Up	Shi et al., 2008
Citrus	Flooding	Up	Up ^b	Up ^b	Arbona et al., 2008
Tobacco	Flooding	Up	Up	Steady	Hurng et al., 1994
Rice	Anoxia	Up	Steady or up	Steady or up	Reggiani et al., 1989
Arabidopsis	Heat	Up	Up	Up	Sagor et al., 2013
Tobacco	Heat	Up	Up	Up	Cvikrova et al., 2012
Wheat	Heat	Down	Up	Up	Goyal and Asthir, 2010
Wheat	Cold	Up	Up	Up	Kovacs et al., 2010
Maize	Cold	Up	Steady	Down	Nemeth et al., 2002
Wheat	CO ₂	Down	Up	Up	Högy et al., 2010
Common sage	UV	Up	Up	Up	Radyukina et al., 2010
Scots pine	K ⁺ deficit	Up	Steady	Steady	Sarjala, 1996
Arabidopsis	K ⁺ deficit	Up	Steady	Steady	Watson and Malmberg, 1996

^aIn leaves but not root tissues; ^bin sensitive genotype only.

around the world. First, externally applied PAs ameliorate stress symptoms. At salt stress, the beneficial effects are due to improved ionic (K⁺/Na⁺) relations (Zhu et al., 2006; Ndayiragije and Lutts, 2007; Roychoudhury et al., 2011; Sharma et al., 2013b) and increased antioxidant activity, both enzymatic (Ozturk and Demir, 2003; Tang and Newton, 2005; Chai et al., 2010) and non-enzymatic, e.g., via proline accumulation (Su and Bai, 2008; Sharma et al., 2013b). Similarly to salinity, PAs improved drought tolerance due to the activation of antioxidant systems, both enzymatic (Kubiś, 2008) and non-enzymatic (such as proline, anthocyanins, and soluble phenolics; Farooq et al., 2009). This reduces the amount of ROS produced (Farooq et al., 2010) and stabilizes membrane structures (Kubiś, 2006). Under natural conditions, PA may also play a beneficial role in mycorrhizal development, contributing to improved plant water status and water use efficiency (Wu et al., 2010). Second, in many cases accumulation of PAs positively correlates with salt (Chattopadhyay et al., 2002; Liu et al., 2006; Mutlu and Bozcuk, 2007) and drought (van der Mescht et al., 1998; Liu et al., 2005), resistance. Third, and maybe the strongest line of evidence came from the experiments with the gain- or loss-of-the-function mutants. Whereas overexpression of enzymes of the PA biosynthesis like arginine decarboxylase, ADC (Roy and Wu, 2001), S-adenosylmethionine synthetase, SAMDC (Waie and Rajam, 2003; Qi et al., 2010), and spermidine synthase, SPDS (Neily et al., 2011) resulted in improved salt tolerance, the loss-of-function mutant of PA biosynthesis genes resulted in reduced stress tolerance in several species (Fariduddin et al., 2013). In Arabidopsis, loss-of-the-function mutants in the synthesis of Spm and thermospermine accumulated more Na⁺ and performed worse than wild type in survival experiments (Alet et al., 2012). Similarly, the introduction of SAMDC gene led to increased polyamine biosynthesis and increased drought

tolerance in tobacco (Waie and Rajam, 2003) and rice (Li and Chen, 2000; Peremarti et al., 2009). Over-expression of ADC in Arabidopsis resulted in a transgenic line with enhanced resistance to high osmoticum, dehydration, and long-term drought (Wang et al., 2011). Transgenic Arabidopsis plants displayed a reduced transpiration rate and stomata conductance, hence, a lesser water loss (Alcázar et al., 2010). Conversely, Arabidopsis mutant plants not capable of producing Spm were hypersensitive to drought (Yamaguchi et al., 2007).

CONTROVERSIES AND INCONSISTENCIES

While a large body of evidence exists suggesting that changes in PA level and plant adaptive responses to environment are positively correlated, negative, or no correlation were reported as well. No correlation between PA content and drought tolerance was found among contrasting rice cultivars (Do et al., 2013) or even higher PAs levels were reported for drought-sensitive chick-pea and beans varieties compared with their tolerant counterparts (Juhász et al., 1997; Nayyar et al., 2005). Although PAs treated rice plants showed improved K⁺/Na⁺ ratio in shoots, PAs did not protect them against salt. Exogenously applied putrescine (Put) induced a decrease in the shoot water content in the presence of NaCl, while spermidine (Spd) and spermine (Spm) were detrimental for cell membrane stability (Ndayiragije and Lutts, 2006). Low-PA lines of maize appeared to be tolerant to salt stress while high-PA lines were more sensitive (Zacchini et al., 1997). Negative correlation between changes in Put content and salinity stress tolerance was reported in experiments comparing different plants (Zapata et al., 2007). Within six selected species, the most tolerant ones accumulate less Put compared with salt-sensitive ones (Zapata et al., 2008). Thus, it appear the concept “the more PA the better for stress tolerance” does not always held.

To support the above statement, we have tabulated some selected evidence for NaCl-induced changes in the level of free PA in root and leaf tissues of various species (Table 2). Of 23 papers cited, eight reported an increase in the level of all three PA; seven have reported a significant reduction in all PA levels; and eight showed mixed results (e.g., an increase in the level of one specific PA accompanied by the concurrent decrease in the level of another PA). Obviously, aspects such as time- and concentration-dependence of PA synthesis and metabolism, its tissue- and organelle-specificity, and inter-conversion between various types and forms of PA, all should be not ignored.

Plant adaptive responses to environment are closely and ultimately related to their ability to control intracellular ion homeostasis and regulate ion transport across cellular membrane (Shabala, 2012). Different tissues show different patterns of ion accumulation, with dicots and monocots sometimes displaying contrasting patterns for a distribution of a certain ion (e.g., Na^+) between different tissues. Understanding of the relative ion accumulation and tissue-specific expression of ion channels and transporters has just started to emerge (Karley et al., 2000; Volkov et al., 2003; Conn and Gilliham, 2010; Gilliham et al., 2011). Plant membranes host hundreds of transport proteins that comprises of ~5% of the entire Arabidopsis genome (Mäser et al., 2001). Some of them are known to be strongly affected by PAs, and PAs can also exert contrasting effects on the same individual ion transporter via diverse mechanisms of action (see below). Thus, the causal role of PA in plant adaptive responses to environment may be established only in the strict context of the tissue- and organelle-specificity.

Table 2 | Selected examples of the changes in the level of free polyamines induced by salt stress in plant root and leaf tissues.

Species	Put	Spd	Spm	References
Cucumber	Up	Up	Up	Fan et al., 2013
Bean	Down	Down	Down	Shevyakova et al., 2013
Pea	Up	Up	Up	Piterkova et al., 2012
Plantago	Down	Down	Down	Radyukina et al., 2009
Mesembryanthemum	Up	Up	Up	Shevyakova et al., 2006
Mesembryanthemum	Down	Down	Up	Stetsenko et al., 2009
Chickpea	Down	Down	Down	Nayyar et al., 2005
Maize	Up	Up	Up	Rodríguez et al., 2009
Apple	Down	Steady	Down	Liu et al., 2008
Grape	Up	Up	Up	Upreti and Murti, 2010
Bromus	Down	Down	Down	Gicquiaud et al., 2002
Tomato	Up	Up	Up	Botella et al., 2000
Tomato	Down	Down	Down	Aziz et al., 1999
Barley	Up	Up	Up	Zhao et al., 2003
Jojoba	Down	Down	Up	Roussos and Pontikis, 2007
Ginseng	Down	Up	Up	Parvin et al., 2012
Wheat	Down	Up	Up	Reggiani et al., 1994
Lupin	Up	Up	Steady	Legocka and Kluk, 2005
Sunflower	Down	Down	Up	Mutlu and Bozcuk, 2007
Soybean	Down	Down	Steady	Xing et al., 2007
Quinoa	Down	Steady	Up	Ruiz-Carrasco et al., 2011
Sunflower	Down	Down	Up	Mutlu and Bozcuk, 2005
Lettuce	Down	Up	Up	Zapata et al., 2003

POLYAMINES EFFECTS ON THE VACUOLAR CATION TRANSPORT

Slow (SV) and fast (FV) vacuolar channels are non-selective cation channels that are ubiquitously and abundantly expressed in higher plant vacuoles (Hedrich et al., 1988; Pottosin and Muñoz, 2002; Hedrich and Marten, 2011; Pottosin and Dobrovinskaya, 2014). SV channels are encoded by the two-pore cation (TPC1) gene (Peiter et al., 2005), whereas the molecular identity of FV channels is still elusive. Both channels conduct a variety of small monovalent cations with a little preference, but SV channels also conduct alkali earth cations like Ca^{2+} and Mg^{2+} (Amodeo et al., 1994; Brüggemann et al., 1999a; Pottosin et al., 2001; Pottosin and Dobrovinskaya, 2014). SV and FV channels only weakly differentiate between K^+ and Na^+ ; this also holds for the case of halophyte plants (Bonales-Alatorre et al., 2013). SV channels are activated by the increase in the cytosolic Ca^{2+} , and, with a lower affinity, by Mg^{2+} (Hedrich and Neher, 1987; Ward and Schroeder, 1994; Pottosin et al., 1997; Carpaneto et al., 2001). At the same time, FV channels are inhibited by the micromolar cytosolic Ca^{2+} and Mg^{2+} (Tikhonova et al., 1997; Brüggemann et al., 1999b; Pei et al., 1999). Therefore, one may propose that the contribution of FV and SV currents into the overall tonoplast cation conductance, among other factors, maybe regulated by the cytosolic Ca^{2+} . In this model, FV channels are more active at the resting Ca^{2+} levels, whereas SV channels require a very substantial cytosolic Ca^{2+} increase for their activation.

In animal cells, several K^+ and cation channels, sensitive to Mg^{2+} , are also sensitive to PAs (Williams, 1997). Similarly, Mg^{2+} -sensitive FV channels were efficiently blocked by micromolar concentrations of Spm and Spd, and by millimolar concentrations of Put (Brüggemann et al., 1998; Dobrovinskaya et al., 1999a). The blockage occurred instantaneously, was dose- but not voltage-dependent, and fully reversible (see Figure 1 and Table 3, for a quantitative description).

SV channels possess a wide pore, with about 0.7 nm diameter in the narrowest constriction (Pottosin and Schönknecht, 2007). Therefore, it is not surprising that PAs act as permeable pore blockers (Dobrovinskaya et al., 1999a,b). This blockage of SV channels is qualitatively very similar to PA blockage of sodium- (Huang and Moczydlowski, 2001), cyclic nucleotide-gated- (Guo and Lu, 2000), and ryanodine- or acetylcholine-receptor channels (Uehara et al., 1996; Haghighi and Cooper, 1998) in animal cells. The block of SV channels was voltage-dependent, with the affinity increasing at increased positive potential at the side of the PAs application. At higher potentials, however, a relief of the block due to a “punchthrough” phenomenon was observed, when PAs were forced to cross the entire channel pore by a strong electric field. Comparison of the voltage-dependence of the block from cytosolic and vacuolar sides displayed several interesting features. First, it appears that PAs reached a common docking point from either membrane side. Second, electric and physical distance from the pore entrance to this stop point was approximately three times longer from the cytosolic side as compared to that from the vacuolar side. Consequently, the cytosolic part of the pore can adopt single Spm or two Put molecules at a time, whereas the vacuolar part can contain a single blocking molecule at a time, regardless of whether it is Spm, Put or a quaternary ammonium ion

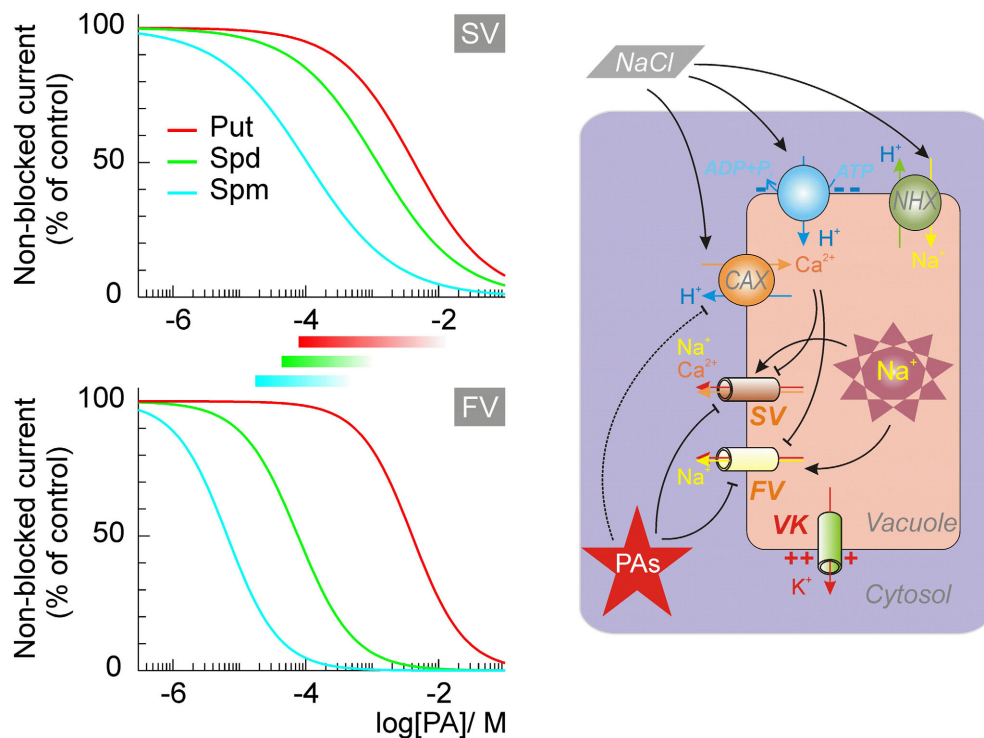


FIGURE 1 | The dose-dependence of the vacuolar cation channels' block by intracellular polyamines and its implications for the salt stress resistance. Dose dependence for SV and FV channels at physiologically attainable (zero) tonoplast potential are drawn, using the values of blocking parameters from Brüggemann et al. (1998) and Dobrovinskaya et al. (1999a,b). Approximate ranges for intracellular PAs in plant cells are indicated by bars. At high salinity, efficient vacuolar Na⁺ sequestration is critical for the salt tolerance. This requires the increased Na⁺/H⁺ antiporter activity and a decrease of Na⁺ leaks through non-selective FV and SV cation channels. The

block by PAs would abolish the FV-mediated current, and strongly suppress the SV current. Continuous operation of the VK, weakly sensitive to PAs, acts as a shunt conductance for the electrogenic H⁺-pump, which fuels the active Na⁺ uptake, and contributes to the recuperation of the salt-induced cytosolic K⁺ loss. Salt stress stimulates expression of the cation-H⁺ antiporters, which may reduce the FV and SV activity via the increase of the luminal Ca²⁺. Over-expression of CAXs is also caused by the inhibition of the Spm⁴⁺ synthesis. Thus, PAs and vacuolar Ca²⁺ may act as alternative regulators of vacuolar cation channels.

(Dobrovinskaya et al., 1999b). At physiological (~ 0 mV) tonoplast potential Spm and Spd block was approximately 10-fold weaker than that of the FV channel, but the affinity of both channels for Put was fairly comparable (Figure 1, Table 3).

Tonoplast also possesses highly K⁺-selective channels (VK), belonging to so called two-pore K⁺ channels family, TPK (Gobert et al., 2007; Isayenkov et al., 2010). These channels are very abundant in guard cells, where they play an important role in mediation of the vacuolar K⁺ release (Ward and Schroeder, 1994). TPK channels may be also found in other tissues (Pottosin et al., 2003; Gobert et al., 2007). TPK is only weakly sensitive to Spm and Spd ($K_d \sim 1$ mM) and practically insensitive to the Put (Hamamoto et al., 2008). Thus, accumulation of PAs during salt stress would primarily inhibit the activity of non-selective cation channels, increasing the overall tonoplast K⁺/Na⁺ selectivity and assisting the efficient vacuolar Na⁺ sequestration (Figure 1).

Effects of PAs on other vacuolar ion transporters are less explored. *Arabidopsis* mutant, lacking the Spm- and tSpm-synthases, consequently has zero Spm and tSpm levels. This mutant shows generally non-altered phenotype under normal growing conditions, except a reduced stem growth (dwarfism) due to the lack of tSpm (Imai et al., 2004). Yet it was hypersensitive

to high NaCl and KCl, but not to the equivalent osmotic stress or high MgCl₂ (Yamaguchi et al., 2006). These mutants also have shown symptoms of the Ca²⁺-deficiency, similar to plants overexpressing vacuolar cation/H⁺ exchangers (CAX). Indeed, transgenic Spm-deficient plants have shown overexpression of several vacuolar CAXs (Figure 1) but same levels of expression of components of the SOS signaling cascade, responsible for the vacuolar Na⁺ sequestration. Causal relations between Spm, CAX-expression, and Ca²⁺ signaling during salt stress remain to be elucidated. Interestingly, whereas Put and Spd but not Spm were essential for the normal growth of *Arabidopsis*, in the case of growth under stress conditions it was just the other way around (Kusano et al., 2008). High ratio of polyamines to diamines positively correlated with a higher activity of vacuolar H⁺-ATPase and PPase as well as with a higher level of phospholipids and lower level of galactolipids in the tonoplast under salt stress (Sun et al., 2002; Liu et al., 2006). In addition to the interaction of PAs with tonoplast phospholipids, binding of PAs to the tonoplast correlated with a higher activity of the V-type H⁺-ATPase and vacuolar Na⁺/H⁺ exchanger, conferring salt tolerance (Zhao and Qin, 2004). Already mentioned tSpm appears to have specific roles in the stem elongation, preventing premature cell death of

Table 3 | Summary of polyamine effects on plant ion channels and pumps.

Channel or pump	Mechanism of the PA action	References
VACUOLAR CHANNELS		
SV (TPC1): slow vacuolar (two-pore cation) Ca^{2+} -permeable channel	Direct, reversible. Voltage-dependent block from either membrane side Spm (50 μM) > Spd (500 μM) > Put (3 mM)*	Dobrovinskaya et al., 1999a,b
FV: fast vacuolar monovalent cation channel	Direct, reversible. Voltage-independent block from the cytosolic side Spm (6 μM) > Spd (80 μM) >> Put (4 mM)*	Brüggemann et al., 1998; Dobrovinskaya et al., 1999a
VK (TPK1): vacuolar K^+ (two-pore K^+)	Direct (?) Voltage-independent, cytosolic side Spm $\sim$ Spd (~ 1 mM) > > Put **	Hamamoto et al., 2008
PLASMA MEMBRANE CHANNELS		
KIRC: inward rectifying K^+ channel -Guard cells (KAT1) -Roots (AKT1)	Indirect, cytosolic side, V-independent Spm $\sim$ Spd $\sim$ Put (0.5–1 mM)** Indirect, extracellular side, V-independent Spm $\sim$ Spd (~ 1.5 mM) > Put**	Liu et al., 2000 Zhao et al., 2007; Zepeda Jazo, 2010
KORC (GORK): outward rectifying K^+ channel	Indirect, extracellular side, V-independent Spm $\sim$ Put (~ 1 mM)**	Zepeda Jazo, 2010
VI-NSCC: voltage-independent cation channel -roots -leaves	Extracellular side, V-independent Spm $\sim$ Spd (~ 0.4 mM) > Put** Extracellular side (indirect?) Spm $\sim$ Put (~ 0.4 mM)**	Zhao et al., 2007 Shabala et al., 2007a,b
ROSIC: weakly voltage-dependent, $\text{OH}^\bullet$ -induced non-selective conductance	Extracellular PAs act as cofactors for ROSIC activation by $\text{OH}^\bullet$ Spm $\sim$ Spd $\sim$ Put (1 mM)	Zepeda-Jazo et al., 2011; Pottosin et al., 2012; Velarde-Buendía et al., 2012
PLASMA MEMBRANE P-TYPE ATPASES		
ACA: autoinhibited Ca^{2+} -ATPase	Rapid activation of Ca^{2+} -pumping Spm $\sim$ Put (0.1–1 mM)	Bose et al., 2011; Zepeda-Jazo et al., 2011; Pottosin et al., 2012; Velarde-Buendía et al., 2012
AHA: autoinhibited H^+ -ATPase	Long-term potentiation Rapid activation (coupled to Ca^{2+} pump) Put (1 mM) Rapid activation and/or inhibition (0.1 or mM Spm) Inhibition of the H^+ pumping; Spm > Spd $\sim$ Put (~ 1 mM) Activation Spm $\sim$ Spd $\sim$ Put (~ 1 mM)** Activation via 14-3-3 proteins binding (Spm only, ~ 0.1 mM)** Long-term suppression, lower expression	Sudha and Ravishankar, 2003 Velarde-Buendía, 2013 Pandolfi et al., 2010 Reggiani et al., 1992; Garufi et al., 2007 Janicka-Russak et al., 2010

*Numbers in brackets are apparent K_d from the data, presented in **Figure 1**.

**Numbers in brackets are approximated concentrations of PAs, producing the half-effect.

When not marked, simply tested experimental concentrations are given.

developing xylem elements (Takehi et al., 2008; Takahashi and Takehi, 2010). On the contrary, the maturation of the xylem elements is achieved via PAs exodus to the apoplast and catabolization therein; released H_2O_2 caused coordinated stiffening of cell walls as well as programmed cell death (PCD) of xylem elements, due to the induction of a Ca^{2+} -permeable conductance in their PM (Tisi et al., 2011; see also the section devoted to PAs and ROS cross-talks below).

At the same time, inhibitory effects of PAs on the vacuolar H^+ -pump activity were also reported (Tang and Newton, 2005; Janicka-Russak et al., 2010). Interestingly, the steady state H^+ pumping was decreased by PAs, but the V- H^+ -ATPase activity was not significantly affected (Janicka-Russak et al., 2010). This may imply that PAs act as uncouplers. In addition, transport of uncharged PAs across the membrane and their

protonation–deprotonation reactions may affect the pH buffering capacity in an acidic compartment and eventually affect the H^+ -ATPase or ATP-synthase activity, as it was shown for the F-type H^+ -ATPase in thylakoids (Ioannidis et al., 2006).

MODULATION OF PLASMA MEMBRANE POTASSIUM AND NON-SELECTIVE CHANNELS BY POLYAMINES

In animal cells, PAs cause a strongly voltage-dependent block of the inward-rectifying (Kir) K^+ channels from the intracellular side; in fact, blockage by PAs is the main cause of the channel inward rectification (Lopatin et al., 1994; Kurata et al., 2010). Inward-rectifying K^+ channels (KIRC) in plants are not related to Kir animal channels and belong to the *Shaker* family, which in animals encode only depolarization-activated K^+ channels (see Sharma et al., 2013a, for a review). It is not surprising, therefore,

that the mechanisms of action of PAs on KIRC may differ from those on Kir. Liu et al. (2000) found that Spm, Spd, and Put, with a little preference, have inhibited KIRC in the guard cell membrane of *Vicia faba* (Table 3). The same work also reported that these PA were also efficient in inhibiting the major component of inward K^+ current, encoded by KAT1 channel, in a heterologic system. The effect of PAs was voltage-independent and showed the same dose-dependence as inhibition of stomata movements. *In planta* measurements revealed that under drought conditions Spd level increased to levels above 1 mM, whereas Put and Spm levels were lower and practically unchanged. This data was interpreted as the evidence for Spd-induced stomata closure to reduce water loss under stress conditions. Importantly, Spd was only efficient from the interior of the guard cell. Yet, when Spd was added at the cytosolic side of small excised membrane patches, no effect on the single channel activity was observed (Liu et al., 2000). Thus, Spd effect on the KIRC was most likely *indirect* and mediated by some unknown intracellular factor or signaling pathway. On contrary, KIRC in barley roots was only affected by PAs from the *extracellular* side (Zhao et al., 2007). In addition to KIRC, the outward-rectifying K^+ channel (KORC) was inhibited indiscriminately by Put or Spm (Table 3). These channels are widely present in root cortex and epidermis and encoded (in *Arabidopsis*) by the GORK gene (Mäser et al., 2001). It should be noted that GORK channel in *Vicia faba* guard cells was unaffected by PAs (Liu et al., 2000). Taken together with a great variability of the PA effects on KORC (e.g., an order of magnitude difference between samples; an occasional but not compulsory reversibility of inhibition) observed in our experiments, it is plausible to suggest that PAs effects on plant *Shaker* K^+ channels are indirect and can be mediated by different factors, present in the apoplast and/or in the cytosol.

In addition to K^+ channels, plants express a variety of non-selective cation currents in the PM (see Demidchik and Maathuis, 2007, for a review). The most common voltage-independent non-selective cation current (VI-NSCC) is almost equally permeable for K^+ and Na^+ , as well as to divalent cations (Ca^{2+}). This current is instantaneous and only weakly voltage-dependent (Demidchik and Tester, 2002). In addition to instantaneous currents, mixed non-selective currents with instantaneous and time-dependent outward-rectifying components can be frequently recorded. The time-dependent but not the instantaneous components were sensitive to (inhibited by) external Na^+ . Thus, it was proposed that VI-NSCC in roots and leaves are major mediators of the toxic Na^+ influx (Shabala et al., 2006). Na^+ -permeable VI-NSCC in roots (Zhao et al., 2007) and leaves (Shabala et al., 2007a,b) were inhibited by externally applied PAs (Table 3). Effect of PAs on the VI-NSCC, albeit reversible, developed slowly (several minutes). There are two possible explanations: (1) PAs acted from the cytosolic side, and their uptake into the cell required a substantial time or (2) PAs effect on the VI-NSCC was indirect. As Zhao et al. (2007) did not found any significant effect of PAs from the cytosolic side on the VI-NSCC, the second possibility seems to be more plausible. Relatively high active concentrations of external PAs raise the question of the physiological significance of their effects on the PM ion channels. However, available data imply that more than half of tissue PAs is associated with the apoplast

(Pistocchi et al., 1988 and references therein). Together with high (up to millimolar) levels of PAs, reached at stress conditions, is justifies a relatively high threshold for the PAs effect on the PM channels.

One of the important determinants of the salt sensitivity mechanism is K^+ loss from plant tissues, caused by the PM depolarization due to the influx of Na^+ (Shabala and Cuin, 2008). In barley the Na^+ -induced K^+ efflux is a main cause of the salt sensitivity and a better control of the membrane potential against the depolarization challenge under salinity is crucial for the tolerance (Chen et al., 2007). In pea mesophyll, externally applied PAs not only inhibited the VI-NSCC, but reduced the salt-induced membrane depolarization and associated loss of K^+ (Shabala et al., 2007a,b). Generalizing this idea, Zepeda-Jazo et al. (2008) proposed a simple model, where PAs inhibition of any NSCC active at depolarized potentials will reduce the membrane depolarization and the loss of K^+ via GORK and NSCC. GORK inhibition by PAs can further reduce the K^+ efflux. A prediction of this hypothesis for the NaCl-induced K^+ efflux was tested on maize and *Arabidopsis* roots. Indeed, PAs could ameliorate NaCl-induced K^+ efflux in some cases. But, depending on the root zone, growing conditions and PA species, the effect of PAs could be null or even resulted in a strong potentiation of K^+ efflux (Pandolfi et al., 2010). Obviously, PAs or their catabolites, can cause not only inhibition but also an activation of some cation currents in the PM (see below).

Extracellular application of PAs *per se* induced the membrane depolarization (Di Tomaso et al., 1989; Fromm et al., 1997; Ozawa et al., 2010; Pottosin et al., 2014b), thus, potentially affecting *any* electrogenic transport across the PM and generating driving force for the K^+ efflux. Our recent pharmacological analysis of the Spm-induced depolarization in barley roots suggested that it was mainly caused by the uptake of PAs via a specific route, not shared with inorganic ions (Pottosin et al., 2014b). Recent advances in the characterization of PA-uptake transporters in plants (Fujita et al., 2012; Mulangi et al., 2012a,b) can provide important clues for the mechanisms of PAs uptake and its impact on the electrogenesis at the plant PM.

INTERPLAY BETWEEN POLYAMINES AND ROS IN THE CONTROL OF PASSIVE CONDUCTANCE AND PUMPING ATPases OF THE PLASMA MEMBRANE

PAs are well-known ROS scavengers and activators of the antioxidant enzymes (Ha et al., 1998; Das and Misra, 2004; Tang and Newton, 2005; Kubiś, 2008; Gill and Tuteja, 2010). At the same time, PAs catabolization generates H_2O_2 , which can be further converted to different ROS, including the most aggressive one, the hydroxyl radical ($\bullet OH$). Export of intracellular PAs to the apoplast and their oxidation therein by available diamine (DAO) and/or polyamine (PAO) oxidase to generate H_2O_2 (Figure 2) is a common signaling pathway segment, which can be found in a variety of plant adaptive and developmental responses. Depending on the strength of the ROS signal generated by the PAs oxidation, very opposite scenarios—e.g., survival vs. PCD—may be realized (see Moschou and Roubelakis-Angelakis, 2014; Pottosin et al., 2014a, for a recent review).

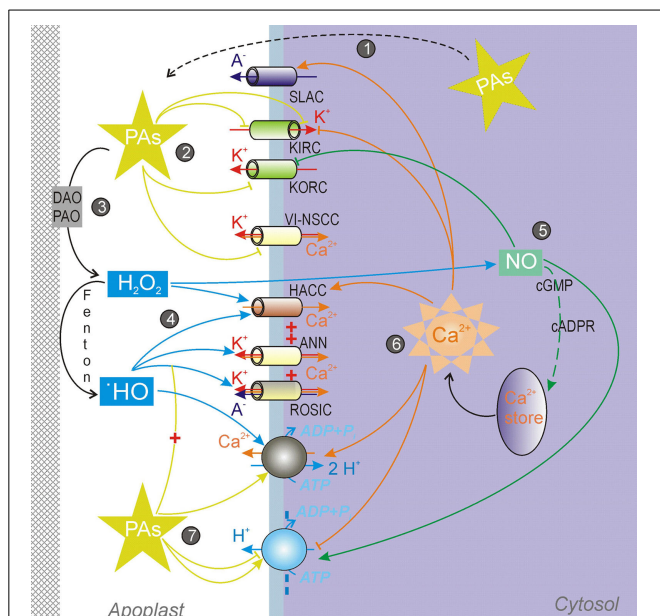


FIGURE 2 | Regulation of ion transport across the plasma membrane by polyamines and their catabolites.

(1) PAs are exported from the cytosol to the apoplast against the electrochemical gradient. (2) PAs inhibit K^+ (inward-rectifying, KIRC and outward rectifying, KORC) and non-selective voltage-independent cation (VI-NSCC) channels. (3) PAs oxidation by diamine (DAO) and/or polyamine (PAO) oxidases generates H_2O_2 in the apoplastic space. There H_2O_2 can be converted to $\bullet OH$ by the single electron reduction catalyzed by transient valency metal ions. (4) H_2O_2 and $\bullet OH$ activate a variety of non-selective Ca^{2+} -permeable channels, including hyperpolarization-activated Ca^{2+} influx channel (HACC), annexin-formed channel (ANN), and non-selective voltage-independent conductance (ROSIC). (5) H_2O_2 , released during PAs catabolization, causes a rapid NO generation. In its turn, NO inhibits KORC by a direct nitrosylation and induces the intracellular Ca^{2+} release via a pathway involving cGMP and cyclic adenosine ribose (cADPR). (6) Ca^{2+} -sensitive network. Several PM channels, including slow anion channel (SLAC) and pumps, are regulated by cytosolic Ca^{2+} . (7) PAs potentiate the ROSIC activation, activate the PM Ca^{2+} -ATPase and alter the activity of the PM H^+ -ATPase.

Studies on transgenic *Arabidopsis* plants, overexpressing enzymes of the Put and Spm biosynthesis, revealed cross-talks between PAs and expression of Ca^{2+} -signaling genes, implying a role of PAs in the Ca^{2+} homeostasis and signaling (Marco et al., 2011). A possible link may be via PAs catabolization and ROS-induced Ca^{2+} fluxes. ROS regulate a variety of ion conductances in the PM. Both H_2O_2 and $\bullet OH$ activate hyperpolarization-activated Ca^{2+} influx currents (HACC) in roots and leaves (Pei et al., 2000; Demidchik et al., 2003, 2007). It appears that properties of HACC, activated by H_2O_2 and $\bullet OH$, are slightly different, despite sharing some characteristics with constitutive HACC. These channels are usually active at non-physiologically large negative potentials (Demidchik and Maathuis, 2007). The presence of the distinct HACC populations, differently responding to H_2O_2 and $\bullet OH$, is manifested by the fact that in the root mature zone HACC are responding only to $\bullet OH$, whereas in the distal elongation zone both H_2O_2 and $\bullet OH$ could induce the Ca^{2+} influx (Demidchik et al., 2007). $\bullet OH$ can activate a variety of conductances, which not only mediate Ca^{2+} influx, but

also cation (K^+) efflux (Figure 2). One of such conductances is mediated by annexin1. It displays both outward and inward rectifying time-dependent components, mediating K^+ efflux and Ca^{2+} influx, respectively (Laohavisit et al., 2012). On the other hand, Demidchik et al. (2010) provided the evidence for the $\bullet OH$ -activation of GORK channels, mediating TEA-sensitive K^+ efflux in *Arabidopsis* roots. However, in pea roots $\bullet OH$ generation caused a rapid cessation of the constitutive GORK (Zepeda-Jazo et al., 2011). Instead, a weakly voltage-dependent instantaneous current, permeable to Ca^{2+} , TEA^+ , K^+ , and even small anions (Cl^-) developed and reached a steady state after 30 min from the initiation of the $\bullet OH$ treatment. Time-dependent depolarization-activated currents developed much later (after 1 h), and were not studied in detail in this plant model. The instantaneous current was termed ROSIC (for ROS-induced conductance) and mediated by tiny (~ 1 pS conductance) channels (Pottosin et al., 2014a). Both $\bullet OH$ -induced K^+ efflux in intact pea roots and ROSIC were sensitive to a variety of cation and anion channels blockers (Zepeda-Jazo et al., 2011). ROSIC activation induced membrane depolarization but caused a massive K^+ efflux so that an efflux of anions likely occurred in parallel (Pottosin et al., 2014a,b). A very similar $\bullet OH$ -induced conductance was reported for barley roots (Velarde-Buendía et al., 2012). PAs unexpectedly stimulated ROSIC and $\bullet OH$ -induced K^+ efflux in intact roots, although by themselves they were incapable to induce any K^+ efflux (Zepeda-Jazo et al., 2011; Velarde-Buendía et al., 2012). In pea such potentiation by PAs was restricted to the mature root zone and not observed in the elongation zone (Pottosin et al., 2012). This is consistent with the idea of the presence of distinct ROS-activated channels populations in these zones. Even more strikingly, ROSIC potentiation by PAs was much larger in salt-sensitive barley than in a salt tolerant variety (Velarde-Buendía et al., 2012), again corroborating the idea on the crucial role of the K^+ retention for the salt tolerance (Chen et al., 2007). Importantly, potentiation of the ROSIC by PAs could be demonstrated in isolated root protoplasts, perfused by an artificial intracellular solution, and containing no amine oxidases. Thus, one has to presume that PAs but not their catabolites acted in this case, and that PAs effects on the ROSIC were likely membrane-delimited.

Externally applied PAs also caused a rapid NO generation, which is likely mediated by PAO and DAO with a generation of H_2O_2 upstream to the NO (Tun et al., 2006; Wimalasekera et al., 2011). NO caused the inhibition of GORK due to a direct protein nitrosylation (Sokolovski and Blatt, 2004). NO also caused stomata closure, via a pathway mediated by cGMP and cADPR, and leading to a Ca^{2+} release from an intracellular store (Figure 2; Neill et al., 2002). This Ca^{2+} release causes the inhibition of KIRC and activation of slow anion channels (SLAC), reducing K^+ uptake and stimulating the anion efflux, respectively; NO did not have any direct effect on these channels (García-Mata et al., 2003). Prolonged (4 days) exposure to NO caused also up to three-fold activation of the PM H^+ -ATPase (Zandonadi et al., 2010).

Activation of the PM Ca^{2+} permeability by ROS and resulting increases of the intracellular free Ca^{2+} could modulate a variety of ion transport processes across the PM (Figure 2). It also exerts a positive feedback regulation on the ROS production by the PM

membrane NADPH-oxidase (Takeda et al., 2008). However, ROS and PAs also affected the Ca^{2+} efflux system, namely the PM Ca^{2+} -ATPase. It was shown that both $\bullet\text{OH}$ and PAs activated eosine-sensitive Ca^{2+} pumping in intact roots (Bose et al., 2011; Zepeda-Jazo et al., 2011; Velarde-Buendía et al., 2012; Velarde-Buendía, 2013). In pea roots the threshold for $\bullet\text{OH}$ activation of the Ca^{2+} pump was at least by an order of magnitude lower as compared for that for the ROSIC (Zepeda-Jazo et al., 2011). Ca^{2+} efflux induced by $\bullet\text{OH}$ in pea roots was transient but could be potentiated by PAs $\text{Spm} > \text{Spd} > \text{Put}$ (Zepeda-Jazo et al., 2011; Pottosin et al., 2012). In the case of Spm, a long-lasting steady state Ca^{2+} efflux was observed, arguing for non-additive effects of $\bullet\text{OH}$ and PAs. No such behavior was found in barley roots, where $\bullet\text{OH}$ - and PA-effects on Ca^{2+} efflux were roughly additive (Velarde-Buendía et al., 2012). Velarde-Buendía (2013) demonstrated that in pea roots PAs affected H^+ fluxes in a differential manner. Whereas Spm caused a net H^+ influx, Put caused a prolonged vanadate-sensitive H^+ efflux, caused by the H^+ -ATPase activation. Spm-induced H^+ influx was consistent with the 1 Ca^{2+} : 2 H^+ exchange mechanism, demonstrated previously (Beffagna et al., 2000). The activation of the H^+ -ATPase by Put appeared to be indirect and coupled to the Ca^{2+} -pump activation (Velarde-Buendía, 2013; Pottosin et al., 2014b). As the PM H^+ -ATPase is inhibited by cytosolic Ca^{2+} (Kinoshita et al., 1995; Brault et al., 2004), it is logical to propose that a cross-talk between the two pumps is mediated by the intracellular Ca^{2+} changes. Yet a supposed decrease of the intracellular Ca^{2+} by PAs needs to be demonstrated directly in future experiments.

H^+ -ATPase operates as a powerhouse, controlling the electric potential difference and the active ion exchange across the PM (Palmgren, 2001). Existing data on immediate effects of PAs on the H^+ -ATPase pumping activity are controversial, with both activation (in rice, Reggiani et al., 1992 and wheat, Liu et al., 2005) and inhibition (in maize, Pandolfi et al., 2010) effects reported. Garufi et al. (2007) described a very specific mechanism of the H^+ -ATPase activation by intracellular Spm, but not by Spd or Put. Spm promoted the binding of 14-3-3 proteins to the unphosphorylated H^+ -ATPase, thus increasing its hydrolytic activity. Long-term treatment with PAs appears to increase the activity of the PM Ca^{2+} -ATPase (Sudha and Ravishankar, 2003) and reduced the magnitude of changes in the H^+ -ATPase activity, induced by the salt stress (Sun et al., 2002; Roy et al., 2005; Tang and Newton, 2005). One day incubation with PAs caused a decrease in transcripts of one of the H^+ -ATPase isoforms, resulting in substantially decreased H^+ pumping across the PM (Janicka-Russak et al., 2010).

IMPLICATIONS FOR STRESS RESPONSES: CURRENT STAND AND PERSPECTIVE

Global climate change is expected to increase the frequency and severity of drought and flooding events in many regions world-wide (Setter and Waters, 2003; Voisenek and Sasidharan, 2013), severely affecting the crop production. On a global scale, the overall loss in food and fiber production due to abiotic stresses is estimated at US\$120 billion p.a. and predicted to increase (<http://www.fao.org/docrep/008/y5800e/Y5800E06.htm>). Also, global availability of good quality water is also

becoming a limiting and increasingly expensive resource, and the cost of irrigation-induced salinity is estimated to exceed US\$11 billion p.a. (Shabala, 2013). Thus, understanding the role of PAs in plant adaptive responses to drought, salinity and flooding may be instrumental in breeding crops with improved tolerance to these stresses to overcome the above losses.

DROUGHT STRESS

Multiple mechanisms contribute to plant adaptive responses to drought. The major ones include (Hu and Schmidhalter, 2005; Shabala and Pottosin, 2014): better stomata control and reduced transpiration under drought conditions; efficient osmotic adjustment, and maintenance of the turgor pressure; prevention of the drought-induced accumulation of ROS; improved water use efficiency and control of a long-distance water transport in plants; maintaining optimal energy status; and improved leaf photochemistry via maintenance of the intracellular ionic homeostasis and charge balance. For obvious reasons, each of these mechanisms is intrinsically dependent on membrane transport activity and implies efficient regulation of ion channels and transporters under stress conditions. Indeed, stomatal opening and closure are strongly dependent on the rapid movement of K^+ into and out of the guard cell (Blatt, 2000). As commented above, the ability of PAs to inhibit KIRC in the guard cell membrane of *Vicia faba* (Liu et al., 2000) may be essential to prevent stomatal opening under drought conditions, thus reducing transpirational water losses. Consistent with this notion, Arabidopsis *gork-1* mutant plants lacking functional GORK channels showed much poorer performance under drought stress, due to their inability to close stomata (Hosy et al., 2003). These findings are in a good agreement with the fact that PAs inhibit opening and induce closure of stomata (Liu et al., 2000; Shi et al., 2010). Importantly, KIRC inhibition by the Spd in *Vicia* guard cells occurred only from the cytosolic side (Liu et al., 2000) highlighting the importance of PA compartmentation.

Drought stress also results in a rapid increase in the level of ROS species in plant tissues (Miller et al., 2010). Polyamines may play a dual role in the process. First, PA may play a critical role in drought stress signaling to confer adaptive responses. According to suggested models, drought-induced increase in ABA content may promote PAs accumulation and exodus into the apoplast, where they were oxidized by the apoplastic amine oxidases, producing H_2O_2 to be used in the signaling cascade (Toumi et al., 2010). On the other hand, PAs are known to significantly enhance activity of both enzymatic (Shi et al., 2010; Wang et al., 2011; Radhakrishnan and Lee, 2013) and non-enzymatic (Högy et al., 2010; Asthir et al., 2012) antioxidants. Therefore, the PA control over the balance between ROS production and scavenging may “shape” H_2O_2 signal, conferring differential stress responses between species and genotypes. Again, tissue- and organelle-specificity of PA accumulation is absolutely essential for this process. Also, given the fact that different PAs may have a different potency for H_2O_2 production and ROS scavenging, it is not the absolute quantity but a balance between various PAs that may be critical to determine the cell fate. Consistent with this notion, An et al. (2012) recently showed that the extent of membrane damage

by drought in maize was dependent on Spd+Spm/Put ratio in plant tissues.

Another important signaling component potentially related to changes in PAs levels is the stress-induced Ca^{2+} signature. Stress- or stimuli-induced elevations in cytosolic free Ca^{2+} , $[\text{Ca}^{2+}]_{\text{cyt}}$, vary in their magnitude, frequency, and shape. These depend on the severity of the stress as well the type of stress experienced, thus creating a unique stress-specific calcium “signature” that is then decoded by signal transduction networks (Bose et al., 2011). The drought stress is not an exception, and transient elevations in cytosolic free Ca^{2+} were reported in response to both hyperosmotic (Ng et al., 2001) and ABA (McAinsh et al., 1997) treatments. As prolonged $[\text{Ca}^{2+}]_{\text{cyt}}$ elevation is detrimental to normal cell metabolism, the basal $[\text{Ca}^{2+}]_{\text{cyt}}$ levels must be restored after the signaling process has been completed. This implies involvement of active Ca^{2+} efflux systems, such as PM and endomembrane Ca^{2+} -ATPase pumps and exchangers (Bose et al., 2011). Meanwhile, Ca^{2+} efflux was shown to be induced by $\bullet\text{OH}$ and potentiated by PAs in a sequence $\text{Spm} > \text{Spd} > \text{Put}$ (Zepeda-Jazo et al., 2011; Pottosin et al., 2012). Thus, the interplay between tissue-specific ROS and PA production and accumulation may confer the shape of stress-specific Ca^{2+} signatures via the modulation of the Ca^{2+} -ATPase activity.

SALINITY STRESS

Plant salinity stress tolerance is a polygenic trait conferred by a large number of sub-traits; each of these is ultimately related to the regulation of membrane-transport activity and ionic homeostasis. The major traits include (Shabala and Munns, 2012): osmotic adjustment; Na^+ exclusion from uptake by roots; intracellular Na^+ sequestration; K^+ retention in the cytosol; tissue-specific Na^+ sequestration; control of the xylem ion loading; Na^+ removal from the shoot; and oxidative stress tolerance. Many if not all these traits may be causally related to, and controlled by, salinity-induced changes in PA levels in various cells compartments.

Rapid osmotic adjustment is absolutely critical to maintain cell turgor and support expansion growth of roots and shoots under saline conditions. Accumulation of K^+ plays a pivotal role in this process, contributing on average to between 35 and 50% of the cell osmotic potential in crops (Shabala and Pottosin, 2014). At the same time, high intracellular K^+ concentrations are required to determine the cell fate and its transition to the PCD. The loss of potassium has been shown to play a primary role in cell shrinkage, caspase activation, and nuclease activity during apoptosis (one of the forms of PCD) in both mammalian (Hughes and Cidlowski, 1999) and plant (Shabala et al., 2007a,b) systems. In this context, the observation that outward-rectifying K^+ channels in root epidermis were inhibited by Put or Spm (Table 3) may be considered as an essential trait enabling K^+ retention in the root and thus contributing to both osmotic adjustment and cell fate determination under saline conditions.

Both changes in the expression levels and activation of existing proteins involved in K^+ transport and sequestration are essential for maintenance in cytosolic K^+ homeostasis under saline conditions. For example, AtCHX17, a member of the CPA2 family of transporters was found to be strongly induced by salinity

(Kreps et al., 2002; Cellier et al., 2004), to compensate for NaCl-induced K^+ exodus from the cytosol resulting from salt-induced depolarization of the PM. However, as the transport capacity of high-affinity K^+ transporters is about 3 orders of magnitude lower compared with channels (Shabala and Pottosin, 2014), cytosolic K^+ homeostasis is seriously compromised. Thus, post-translational regulation and modulation of activities of existing channels or transporters by various factors and second messengers (including PAs) is more significant. More details on factors controlling K^+ transport under stress conditions are available in Shabala and Pottosin (2014).

Efficient vacuolar sequestration of the cytotoxic Na^+ is another prominent mechanism conferring salinity tolerance in plants. This sequestration is achieved by mean of the tonoplast Na^+/H^+ antiporters fueled by the vacuolar H^+ -ATPase and H^+ -PPase pumps (Hasegawa, 2013; Shabala, 2013). In addition, toxic Na^+ ions must be prevented from leaking back into cytosol. Thus, to avoid energy consuming futile Na^+ cycling between cytosol and vacuole and to achieve efficient vacuolar sequestration of toxic Na^+ , passive tonoplast Na^+ conductance has to be kept at absolute minimum. This implies a strict and efficient control over Na^+ -permeable tonoplast SV and FV channels (Bonales-Alatorre et al., 2013). Each of these channels is PA-sensitive and can be blocked by the physiologically relevant concentrations of PAs (see Figure 1). Therefore, salinity stress-induced elevation in PA levels may be essential to enable efficient vacuolar Na^+ sequestration. Consistent with this notion, salt stress inhibited the activity of polyamine synthesizing enzymes (L-arginine decarboxylase and L-ornithine decarboxylase) in glycophyte species of *Vigna radiata* but not in halophyte *P. undulate* (Friedman et al., 1989), thus potentially enabling efficient control over vacuolar Na^+ sequestration in the latter (naturally salt tolerant) species.

Reducing net Na^+ accumulation in the cytosol by controlling the rate of its transport across the PM may be another way of improving plant performance under saline conditions. Non-selective cation channels (NSCC) are considered to be a major pathway of Na^+ uptake into the cell (Demidchik and Maathuis, 2007), and physiologically relevant concentrations of PAs were efficient in inhibiting NSCC-mediated Na^+ currents in leaf (Shabala et al., 2007a,b) and root (Zhao et al., 2007; Zepeda-Jazo et al., 2008) tissues.

FLOODING STRESS

Two major factors affect plant growth and performance in flooded soils: reduced oxygen availability leading to a sharp decline in ATP production, and elemental toxicity originating from the changes in the soil redox potential (Shabala, 2011). Acclimation to flooded conditions requires significant metabolic alterations in living cells. This includes reduced energy consumption, activation of pathways that generate ATP without oxidative phosphorylation, and increased chaperone activity to deal with increased ROS production (Bailey-Serres and Voesenek, 2010; Voesenek and Sasidharan, 2013). Polyamines seem to be instrumental in this metabolic adaptation. Accumulation of Put in flooded roots was shown to be able to stimulate the PM ATPase activity (Bertani et al., 1997), conferring beneficial effects to cell ionic homeostasis and nutrient acquisition. Superoxide radical

and H₂O₂ contents were also reduced in flooding-stressed onion plants after Put pre-treatment (Yiu et al., 2009). Jia et al. (2010) showed that application of exogenous Spd to hypoxic cucumber roots or conversion of Put to Spd and Spm enhanced the aerobic respiration but inhibited the fermentation metabolism in roots, leading to an increase in ATP content and alleviation of the stress symptoms.

A massive increase in the amount of available Mn and Fe in the soil solution is observed within a few days of onset of water-logging, often to above toxic levels (Marschner, 1995; Zeng et al., 2013), due to the changes in the soil redox potential. Being a transition metal, Fe is highly redox active and, in the presence of H₂O₂, can mediate production of the hydroxyl radical through the Fenton reaction (Rodrigo-Moreno et al., 2013). This may cause lipid peroxidation and damage to key cellular structures as well as result in a massive K⁺ exodus from the cytosol mediated by •OH-activated K⁺-selective outward rectifying (Demidchik et al., 2010) and non-selective K⁺ permeable (Zepeda-Jazo et al., 2011) channels. Both these processes are detrimental to cell metabolism and plant performance under stress conditions. Importantly, a regulatory role of PAs in plant adaptation to flooding seems to be closely related to intracellular K⁺ homeostasis. In the absence of K⁺, anoxia led to a decrease in Put, Spd, and Spm levels. The presence of K⁺ ions during the anaerobic treatment abolished the negative effect of anoxia on polyamine titers and slightly increased them (Reggiani et al., 1993).

OUTLOOK

A lesson learned from studies of the PAs effects on plant ion channels is that, contrary to their animal counterparts, a direct pore blockage mechanism is uncommon. A notable exception to that observation is a high affinity block of vacuolar non-selective cation channels of FV and SV types. Because these channels are ubiquitously expressed in plant tissues the model shown in **Figure 1** describing the impact of PAs on vacuolar cation transport, can be considered as a general one. Validation of this working model can be demonstrated by genetic manipulation of SV, FV and VK channels by silencing, or site directed mutations, affecting channels' sensitivity to PAs. While the molecular identities of SV and VK channels are known, that of the FV channel remains cryptic. Completing of this lacking information will allow the manipulation of the overall tonoplast cation transport and eventually will help to understand the role of PAs in its control in plant responses to abiotic stresses.

In most cases, the action spectrum of PAs depends on the side of the membrane/ compartment, whether it is vacuolar lumen, cytosol or apoplast. Unfortunately, PAs compartmentation and membrane transport in plants are poorly explored. In particular, mechanisms of PAs uptake or active efflux across the PM are unknown (Igarashi and Kashiwagi, 2010). This knowledge is very important, however, to understand the exodus of PAs to the apoplast, where they are normally absent. In many cases, it can be proved that rather than PAs themselves, their catabolites (and especially ROS), exert the effect on plant membrane transporters. It is important to mention that the apparent specificity of diamine (Put) vs. PAs (Spd, Spm) effects in this case may be caused by a higher activity of the apoplastic DAO in dicots like Fabaceae or PAO in monocots like Poaceae (Moschou

et al., 2008). Overall, the relation between PAs biosynthesis and catabolism (or, in other words, respective levels of PAs and their catabolites) may determine whether survival or PCD responses would be initiated (Moschou and Roubelakis-Angelakis, 2014). This is also applicable to the balance between PAs actions as ROS scavengers and antioxidant system activators, and PAs as a ROS source. In this regard, ROS speciation also becomes crucial. Whereas H₂O₂ is a relatively long-living and easy membrane-permeable molecule, •OH is short-lived and acts in the closest vicinity of the transient valency metal, which catalyzed its generation. A substantial evidence was obtained for differential effects of •OH and H₂O₂ on plant membrane transporters. In particular, a newly described dual cation and anion conductance, ROSIC, is activated only by •OH and PAs further modulate it in species- and tissue-dependent mode.

Apart of recently revealed cross-talks between PAs and ROS, an important link between PAs catabolism and stress response may be the PAs-induced generation of the NO (Wimalasekera et al., 2011), which in turn affects a variety of PM transporters, either directly or via the intracellular Ca²⁺ signal (**Figure 2**). In addition to ROS-activated Ca²⁺ influx channels, both ROS and PAs are capable to activate PM Ca²⁺ pumps. Thus, the fine tuning of Ca²⁺ signal may be achieved, which is worth of further experimental exploration.

It is conceivable that stress-induced changes of PAs and ROS metabolism were adapted for the stress resistance in a rather opportunistic way. Whereas few direct sensors for PAs and ROS evolved, other targets may be indirect and the net effect, e.g., on the K⁺ transport across the PM, may be rather variable (Pandolfi et al., 2010). While searching for a solution of the equation with many parameters, one needs to take into the account PAs synthesis, transport, and catabolization. In addition, tissue-, species-, and physiological status-dependent expression of different ion channels and transporters as well as the modes of action of PAs and their catabolites should be also always kept in mind. Despite its complexity, this task is the only possible alternative, as the “spray and pray” strategy seems to be not applicable in the case of polyamines.

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REFERENCES

- Ahern, G. P., Wang, X., and Miyares, R. L. (2006). Polyamines are potent ligands for the capsaicin receptor TRPV1. *J. Biol. Chem.* 281, 8991–8995. doi: 10.1074/jbc.M513429200
- Alcázar, R., Marco, F., Cuevas, J. C., Patron, M., Ferrando, A., Carrasco, P., et al. (2006). Involvement of polyamines in plant response to abiotic stress. *Biotech. Lett.* 28, 1867–1876. doi: 10.1007/s10529-006-9179-3
- Alcázar, R., Planas, J., Saxena, T., Zarza, X., Bortolotti, C., Cuevas, J., et al. (2010). Putrescine accumulation confers drought tolerance in transgenic Arabidopsis plants over-expressing the homologous Arginine decarboxylase 2 gene. *Plant Physiol. Biochem.* 48, 547–552. doi: 10.1016/j.plaphy.2010.02.002
- Alet, A. I., Sanchez, D. H., Cuevas, J. C., Marina, M., Carrasco, P., Altabella, T., et al. (2012). New insights into the role of spermine in *Arabidopsis thaliana* under long-term salt stress. *Plant Sci.* 182, 94–100. doi: 10.1016/j.plantsci.2011.03.013
- Amodeo, G., Escobar, A., and Zeiger, E. (1994). A cationic channel in the guard cell tonoplast of *Allium cepa*. *Plant Physiol.* 105, 999–1006.

- An, Z. F., Li, C. Y., Zhang, L. X., and Alva, A. K. (2012). Role of polyamines and phospholipase D in maize (*Zea mays* L.) response to drought stress. *S. Afr. J. Bot.* 83, 145–150. doi: 10.1016/j.sajb.2012.08.009
- Arbona, V., Hossain, Z., Lopez-Clemente, M. F., Perez-Clemente, R. M., and Gomez-Cadenas, A. (2008). Antioxidant enzymatic activity is linked to waterlogging stress tolerance in citrus. *Physiol. Plant* 132, 452–466. doi: 10.1111/j.1399-3054.2007.01029.x
- Asthir, B., Koundal, A., and Bains, N. S. (2012). Putrescine modulates antioxidant defense response in wheat under high temperature stress. *Biol. Plant* 56, 757–761. doi: 10.1007/s10535-012-0209-1
- Aziz, A., Martin-Tanguy, J., and Larher, F. (1999). Salt stress-induced proline accumulation and changes in tyramine and polyamine levels are linked to ionic adjustment in tomato leaf discs. *Plant Sci.* 145, 83–91. doi: 10.1016/S0168-9452(99)00071-0
- Bailey-Serres, J., and Voesenek, L. (2010). Life in the balance: a signaling network controlling survival of flooding. *Curr. Opin. Plant Biol.* 13, 489–494. doi: 10.1016/j.pbi.2010.08.002
- Beffagna, N., Romani, G., and Sforza, M. C. (2000). H^+ fluxes at plasmalemma level: *in vivo* evidence for a significant contribution of the Ca^{2+} -ATPase and for the involvements of its activity in the abscisic acid-induced changes in *Egeria densa* leaves. *Plant Biol.* 2, 168–175. doi: 10.1055/s-2000-9158
- Bertani, A., Brambilla, I., Mapelli, S., and Reggiani, R. (1997). Elongation growth in the absence of oxygen: the rice coleoptile. *Russ. J. Plant Physiol.* 44, 543–547.
- Blatt, M. R. (2000). Cellular signaling and volume control in stomatal movements in plants. *Annu. Rev. Cell Dev. Biol.* 16, 221–241. doi: 10.1146/annurev.cellbio.16.1.221
- Bonales-Alatorre, E., Shabala, S., Chen, Z. H., and Pottosin, I. (2013). Reduced tonoplast fast-activating and slow-activating channel activity is essential for conferring salinity tolerance in a facultative halophyte, quinoa. *Plant Physiol.* 162, 940–952. doi: 10.1104/pp.113.216572
- Bose, J., Pottosin, I., Shabala, S. S., Palmgren, M. G., and Shabala, S. (2011). Calcium efflux systems in stress signalling and adaptation in plants. *Front. Plant Sci.* 2:85. doi: 10.3389/fpls.2011.00085
- Botella, M. A., del Amor, F. M., Amoros, A., Serrano, M., Martinez, V., and Cerda, A. (2000). Polyamine, ethylene and other physico-chemical parameters in tomato (*Lycopersicon esculentum*) fruits as affected by salinity. *Physiol. Plant* 109, 428–434. doi: 10.1034/j.1399-3054.2000.100409.x
- Braut, M., Amiar, Z., Pennarun, A. M., Monestiez, M., Zhang, Z., Cornel, D., et al. (2004). Plasma membrane depolarization induced by ABA in *Arabidopsis thaliana* suspension cells involves reduction of proton pumping in addition to anion channel activation which are both Ca^{2+} dependent. *Plant Physiol.* 135, 231–243. doi: 10.1104/pp.104.039255
- Brügemann, L. I., Pottosin, I. I., and Schönknecht, G. (1998). Cytoplasmic polyamines block the fast activating vacuolar cation channel. *Plant J.* 16, 101–105. doi: 10.1046/j.1365-313X.1998.00274.x
- Brügemann, L. I., Pottosin, I. I., and Schönknecht, G. (1999a). Selectivity of the fast activating vacuolar cation channel. *J. Exp. Bot.* 50, 873–876.
- Brügemann, L. I., Pottosin, I. I., and Schönknecht, G. (1999b). Cytoplasmic magnesium regulates the fast activating vacuolar cation channel. *J. Exp. Bot.* 50, 1547–1552. doi: 10.1093/jxb/50.339.1547
- Carpaneto, A., Cantu, A. M., and Gambale, F. (2001). Effects of cytoplasmic Mg^{2+} on slowly activating channels in isolated vacuoles of *Beta vulgaris*. *Planta* 213, 457–468. doi: 10.1007/s004250100519
- Cellier, F., Conejero, G., Ricaud, L., Luu, D. T., Lepetit, M., and Gosti, F., et al. (2004). Characterization of AtCHX17, a member of the cation/ H^+ exchangers, CHX family, from *Arabidopsis thaliana* suggests a role in K^+ homeostasis. *Plant J.* 39, 834–846. doi: 10.1111/j.1365-313X.2004.02177.x
- Chai, Y. Y., Jiang, C. D., Shi, L., Shi, T. S., and Gu, W. B. (2010). Effects of exogenous spermine on sweet sorghum during germination under salinity. *Biol. Plant* 54, 145–148. doi: 10.1007/s10535-010-0023-1
- Chattopadhyay, M. K., Tiwari, B. S., Chattopadhyay, G., Bose, A., Sengupta, D. N., and Ghosh, B. (2002). Protective role of exogenous polyamines on salinity-stressed rice (*Oryza sativa*) plants. *Physiol. Plant* 116, 192–199. doi: 10.1034/j.1399-3054.2002.1160208.x
- Chen, Z., Pottosin, I. I., Cuin, T. A., Fuglsang, A. T., Tester, M., Jha, D., et al. (2007). Root plasma membrane transporters controlling K^+/Na^+ homeostasis in salt stressed barley. *Plant Physiol.* 145, 1714–1725. doi: 10.1104/pp.107.110262
- Conn, S., and Gilliam, M. (2010). Comparative physiology of elemental distributions in plants. *Ann. Bot.* 105, 1081–1102. doi: 10.1093/aob/mcq027
- Cvikrova, M., Gemperlova, L., Dobra, J., Martincova, O., Prasil, I. T., Gubis, J., et al. (2012). Effect of heat stress on polyamine metabolism in proline-over-producing tobacco plants. *Plant Sci.* 182, 49–58. doi: 10.1016/j.plantsci.2011.01.016
- Das, K. C., and Misra, H. P. (2004). Hydroxyl radical scavenging and singlet oxygen quenching properties of polyamines. *Mol. Cell. Biochem.* 262, 127–133. doi: 10.1023/B:MCBI.0000038227.91813.79
- De Diego, N., Sampedro, M. C., Barrio, R. J., Saiz-Fernandez, I., Moncalean, P., and Lacuesta, M. (2013). Solute accumulation and elastic modulus changes in six radiata pine breeds exposed to drought. *Tree Physiol.* 33, 69–80. doi: 10.1093/treephys/tps125
- Demidchik, V., Cuin, T. A., Svistunenko, D., Smith, S. J., Miller, A. J., Shabala, S., et al. (2010). Arabidopsis root K^+ -efflux conductance activated by hydroxyl radicals: single-channel properties, genetic basis and involvement in stress-induced cell death. *J. Cell Sci.* 123, 1468–1479. doi: 10.1242/jcs.064352
- Demidchik, V., and Maathuis, F. J. M. (2007). Physiological roles of nonselective cation channels in plants: from salt stress to signalling and development. *New Phytol.* 175, 387–404. doi: 10.1111/j.1469-8137.2007.02128.x
- Demidchik, V., Shabala, S. N., Coutts, K. B., Tester, M. A., and Davies, J. M. (2003). Free oxygen radicals regulate plasma membrane Ca^{2+} - and K^+ - permeable channels in plant root cells. *J. Cell Sci.* 116, 81–88. doi: 10.1242/jcs.00201
- Demidchik, V., Shabala, S. N., and Davies, J. M. (2007). Spatial variation in H_2O_2 response of *Arabidopsis thaliana* root epidermal Ca^{2+} flux and plasma membrane Ca^{2+} channels. *Plant J.* 49, 377–386. doi: 10.1111/j.1365-313X.2006.02971.x
- Demidchik, V., and Tester, M. (2002). Sodium fluxes through nonselective cation channels in the plasma membrane of protoplasts from Arabidopsis roots. *Plant Physiol.* 128, 379–387. doi: 10.1104/pp.010524
- Di Tomaso, J. M., Shaff, J. E., and Kochian, L. V. (1989). Putrescine-induced wounding and its effects on membrane integrity and ion transport processes in roots of intact corn seedlings. *Plant Physiol.* 90, 988–995. doi: 10.1104/pp.90.3.988
- Do, P. T., Degenkolbe, T., Erban, A., Heyer, A. G., Kopka, J., Kohl, K. I., et al. (2013). Dissecting rice polyamine metabolism under controlled long-term drought stress. *PLoS ONE* 8:e60325. doi: 10.1371/journal.pone.0060325
- Dobrovinskaya, O. R., Muñoz, J., and Pottosin, I. I. (1999a). Inhibition of vacuolar ion channels by polyamines. *J. Membr. Biol.* 167, 127–140. doi: 10.1007/s002329900477
- Dobrovinskaya, O. R., Muñoz, J., and Pottosin, I. I. (1999b). Asymmetric block of the plant vacuolar Ca^{2+} permeable channel by organic cations. *Eur. Biophys. J.* 28, 552–563.
- Drouin, H., and Hermann, H. (1994). Intracellular action of spermine on neuronal Ca^{2+} and K^+ currents. *Eur. J. Neurosci.* 6, 412–419. doi: 10.1111/j.1460-9568.1994.tb00284.x
- Fan, H. F., Du, C. X., and Guo, S. R. (2013). Nitric oxide enhances salt tolerance in cucumber seedlings by regulating free polyamine content. *Env. Exp. Bot.* 86, 52–59. doi: 10.1016/j.envexpbot.2010.09.007
- Fariduddin, Q., Varshney, P., Yusuf, M., and Ahmad, A. (2013). Polyamines: potent modulators of plant responses to stress. *J. Plant Int.* 8, 1–16. doi: 10.1080/17429145.2012.716455
- Farooq, M., Wahid, A., and Lee, D. J. (2009). Exogenously applied polyamines increase drought tolerance of rice by improving leaf water status, photosynthesis and membrane properties. *Acta Physiol. Plant* 31, 937–945. doi: 10.1007/s11738-009-0307-2
- Farooq, M., Wahid, A., Lee, D. J., Cheema, S. A., and Aziz, T. (2010). Comparative time course action of the foliar applied glycinebetaine, salicylic acid, nitrous oxide, brassinosteroids and spermine in improving drought resistance of rice. *J. Agron. Crop Sci. Z. Acker Pflanzenbau.* 196, 336–345. doi: 10.1111/j.1439-037X.2010.00422.x
- Friedman, R., Altman, A., and Levin, N. (1989). The effect of salt stress on polyamine biosynthesis and content in mung bean plants and in halophytes. *Physiol. Plant* 76, 295–302. doi: 10.1111/j.1399-3054.1989.tb06194.x
- Fromm, J., Meyer, A. J., and Weissenel, M. H. (1997). Growth, membrane potential and endogenous ion currents of willow (*Salix viminalis*) roots are all affected by abscisic acid and spermine. *Physiol. Plant* 99, 529–537. doi: 10.1111/j.1399-3054.1997.tb05353.x
- Fujita, M., Fujita, Y., Iuchi, S., Yamada, K., Kobayashi, Y., Urano, K., et al. (2012). Natural variation in a polyamine transporter determines paraquat tolerance in *Arabidopsis*. *Proc. Natl. Acad. Sci. U.S.A.* 109, 6343–6347. doi: 10.1073/pnas.1121406109

- Galston, A. W., and Sawhney, R. K. (1990). Polyamines in plant physiology. *Plant Physiol.* 94, 406–410. doi: 10.1104/pp.94.2.406
- García-Mata, C., Gay, R., Sokolovski, S., Hills, A., Lamattina, L., and Blatt, M. R. (2003). Nitric oxide regulates K^+ and Cl^- channels in guard cells through a subset of abscisic acid-evoked signaling pathways. *Proc. Natl. Acad. Sci. U.S.A.* 100, 11116–11121. doi: 10.1073/pnas.1434381100
- Garufi, A., Visconti, S., Camoni, L., and Aducci, P. (2007). Polyamines as physiological regulators of 14-3-3 interaction with the plant plasma membrane H^+ -ATPase. *Plant Cell Physiol.* 48, 434–440. doi: 10.1093/pcp/pcm010
- Gicquiaud, L., Hennion, F., and Esnault, M. A. (2002). Physiological comparisons among four related *Bromus* species with varying ecological amplitude: polyamine and aromatic amine composition in response to salt spray and drought. *Plant Biol.* 4, 746–753. doi: 10.1055/s-2002-37401
- Gill, S. S., and Tuteja, N. (2010). Polyamines and abiotic stress tolerance in plants. *Plant Signal. Behav.* 5, 26–33. doi: 10.4161/psb.5.1.10291
- Gillilham, M., Athman, A., Tyerman, S. D., and Conn, S. J. (2011). Cell-specific compartmentation of mineral nutrients is an essential mechanism for optimal plant productivity—another role for TPC1? *Plant Signal. Behav.* 6, 1656–1661. doi: 10.4161/psb.6.11.17797
- Gobert, A., Isayenkov, S., Voelker, C., Czempinski, K., and Maathuis, F. J. M. (2007). The two-pore channel TPK1 gene encodes the vacuolar K^+ conductance and plays a role in K^+ homeostasis. *Proc. Natl. Acad. Sci. U.S.A.* 104, 10726–10731. doi: 10.1073/pnas.0702595104
- Goyal, M., and Asthir, B. (2010). Polyamine catabolism influences antioxidative defense mechanism in shoots and roots of five wheat genotypes under high temperature stress. *Plant Growth Regul.* 60, 13–25. doi: 10.1007/s10725-009-9414-8
- Grzesiak, M., Filek, M., Barbasz, A., Kreczmer, B., and Hartikainen, H. (2013). Relationships between polyamines, ethylene, osmoprotectants and antioxidant enzymes activities in wheat seedlings after short-term PEG- and NaCl-induced stresses. *Plant Growth Regul.* 69, 177–189. doi: 10.1007/s10725-012-9760-9
- Guo, D., and Lu, Z. (2000). Mechanism of cGMP-gated channel block by intracellular polyamines. *J. Gen. Physiol.* 115, 783–797. doi: 10.1085/jgp.115.6.783
- Gupta, K., Dey, A., and Gupta, B. (2013). Plant polyamines in abiotic stress responses. *Acta Physiol. Plant* 35, 2015–2036. doi: 10.1007/s11738-013-1239-4
- Ha, H. C., Sirisoma, N. S., Kuppusamy, P., Zweier, J. L., Woster, P. M., and Casero, R. A. Jr. (1998). The natural polyamine spermine functions directly as a free radical scavenger. *Proc. Natl. Acad. Sci. U.S.A.* 95, 11140–11145. doi: 10.1073/pnas.95.19.11140
- Haghighi, A. P., and Cooper, E. (1998). Neuronal nicotinic acetylcholine receptors are blocked by intracellular spermine in a voltage-dependent manner. *J. Neurosci.* 18, 4050–4062.
- Hamamoto, S., Marui, J., Matsuoaka, K., Higashi, K., Igarashi, K., Nakagawa, T., et al. (2008). Characterization of a tobacco TPK-type K^+ channel as a novel tonoplast K^+ channel using yeast tonoplasts. *J. Biol. Chem.* 283, 1911–1920. doi: 10.1074/jbc.M708213200
- Hasegawa, P. M. (2013). Sodium homeostasis and salt tolerance of plants. *Env. Exp. Bot.* 92, 19–31. doi: 10.1016/j.envexpbot.2013.03.001
- Hedrich, R., Barbier-Brygoo, H., Felle, H., Flüge, U. I., Lüttge, U., Maathuis, F. J. M., et al. (1988). General mechanisms for solute transport across the tonoplast of plant vacuoles: a patch-clamp survey of ion channels and proton pumps. *Bot. Acta* 101, 7–13. doi: 10.1111/j.1438-8677.1988.tb00003.x
- Hedrich, R., and Marten, I. (2011). TPC1-SV channels gain shape. *Mol. Plant* 4, 428–441. doi: 10.1093/mp/ssr017
- Hedrich, R., and Neher, E. (1987). Cytoplasmic calcium regulates voltage-dependent ion channels in plant vacuoles. *Nature* 329, 833–836. doi: 10.1038/329833a0
- Högy, P., Keck, M., Niehaus, K., Franzaring, J., and Fangmeier, A. (2010). Effects of atmospheric CO_2 enrichment on biomass, yield and low molecular weight metabolites in wheat grain. *J. Cereal Sci.* 52, 215–220. doi: 10.1016/j.jcs.2010.05.009
- Hosy, E., Vavasseur, A., Mouline, K., Dreyer, I., Gaymard, F., Poree, F., et al. (2003). The Arabidopsis outward K^+ channel GORK is involved in regulation of stomatal movements and plant transpiration. *Proc. Natl. Acad. Sci. U.S.A.* 100, 5549–5554. doi: 10.1073/pnas.0733970100
- Huang, C. J., and Moczydlowski, E. (2001). Cytoplasmic polyamines as permeant blockers and modulators of the voltage-gated sodium channel. *Biophys. J.* 80, 1262–1279. doi: 10.1016/S0006-3495(01)76102-4
- Hughes, F. M., and Cidlowski, J. A. (1999). Potassium is a critical regulator of apoptotic enzymes *in vitro* and *in vivo*. *Adv. Enzyme Regul.* 39, 157–171. doi: 10.1016/S0065-2571(98)00010-7
- Hurng, W. P., Lur, H. S., Liao, C. K., and Kao, C. H. (1994). Role of abscisic acid, ethylene and polyamines in flooding-promoted senescence of tobacco-leaves. *J. Plant Physiol.* 143, 102–105. doi: 10.1016/S0176-1617(11)82104-8
- Hu, Y., and Schmidhalter, U. (2005). Drought and salinity: a comparison of their effects on the mineral nutrition in plants. *J. Plant Nutr. Soil. Sci.* 168, 541–549. doi: 10.1002/jpln.200420516
- Igarashi, K., and Kashiwagi, K. (2010). Characteristics of cellular polyamine transport in prokaryotes and eukaryotes. *Plant Physiol. Biochem.* 48, 506–512. doi: 10.1016/j.plaphy.2010.01.017
- Imai, A., Takahashi, A., Kato, T., Sato, S., Tabata, S., Yamamoto, K. T., et al. (2004). Spermine is not essential for survival of Arabidopsis. *FEBS Lett.* 556, 148–152. doi: 10.1016/S0014-5793(03)01395-4
- Ioannidis, N. E., Sfichi, L., and Kotzabasis, K. (2006). Putrescine stimulates chemiosmotic ATP synthesis. *Biochim. Biophys. Acta* 1757, 821–828. doi: 10.1016/j.bbabi.2006.05.034
- Isayenkov, S., Isner, J. C., and Maathuis, F. J. M. (2010). Vacuolar ion channels: role in plant nutrition and signaling. *FEBS Lett.* 584, 1982–1988. doi: 10.1016/j.febslet.2010.02.050
- Janicka-Russak, M., Kabała, K., Młodzińska, E., and Kłobus, G. (2010). The role of polyamines in the regulation of the plasma membrane and the tonoplast proton pumps under salt stress. *J. Plant Physiol.* 167, 261–269. doi: 10.1016/j.jplph.2009.09.010
- Jia, Y. X., Sun, J., Guo, S. R., Li, J., Hu, X. H., and Wang, S. P. (2010). Effect of root-applied spermidine on growth and respiratory metabolism in roots of cucumber (*Cucumis sativus*) seedlings under hypoxia. *Russ. J. Plant Physiol.* 57, 648–655. doi: 10.1134/S1021443710050079
- Juhász, A. G., Simon-Sarkadi, L., Velich, I., and Varro, P. (1997). Studies of non-ionic osmotic stress on bean (*Phaseolus vulgaris* L.) callus and seedlings cultures. *Acta Hort.* 447, 455–456.
- Kakehi, J., Kuwashiro, Y., Niitsu, M., and Takahashi, T. (2008). Thermospermine is required for stem elongation in *Arabidopsis thaliana*. *Plant Cell Physiol.* 49, 1342–1349. doi: 10.1093/pcp/pcn109
- Karley, A. J., Leigh, R. A., and Sanders, D. (2000). Differential ion accumulation and ion fluxes in the mesophyll and epidermis of barley. *Plant Physiol.* 122, 835–844. doi: 10.1104/pp.122.3.835
- Kinoshita, T., Nishimura, M., and Shimazaki, K. (1995). Cytosolic concentration of Ca^{2+} regulates the plasma membrane H^+ -ATPase in guard cells of fava bean. *Plant Cell* 7, 1333–1342. doi: 10.1105/tpc.7.8.1333
- Kovacs, Z., Simon-Sarkadi, L., Szucs, A., and Kocsy, G. (2010). Differential effects of cold, osmotic stress and abscisic acid on polyamine accumulation in wheat. *Amino Acids* 38, 623–631. doi: 10.1007/s00726-009-0423-8
- Kreps, J. A., Wu, Y., Chang, H. S., Zhu, T., Wang, X., and Harper, J. F. (2002). Transcriptome changes for Arabidopsis in response to salt, osmotic, and cold stress. *Plant Physiol.* 130, 2129–2141. doi: 10.1104/pp.008532
- Kubiš, J. (2003). Polyamines and scavenging system: influence of exogenous spermidine on catalase and guaiacol peroxidase activities, and free polyamine level in barley leaves under water deficit. *Acta Physiol. Plant.* 25, 337–343. doi: 10.1007/s11738-003-0014-3
- Kubiš, J. (2006). Exogenous spermidine alters in different way membrane permeability and lipid peroxidation in water stressed barley leaves. *Acta Physiol. Plant* 28, 27–33. doi: 10.1007/s11738-006-0065-3
- Kubiš, J. (2008). Exogenous spermidine differentially alters activities of some scavenging system enzymes, H_2O_2 and superoxide radical levels in water-stressed cucumber leaves. *J. Plant Physiol.* 165, 397–406. doi: 10.1016/j.jplph.2007.02.005
- Kurata, H. T., Zhu, E. A., and Nichols, C. G. (2010). Locale and chemistry of spermine binding in the archetypal inward rectifier Kir2.1. *J. Gen. Physiol.* 135, 495–508. doi: 10.1085/jgp.200910253
- Kusano, T., Berberich, T., Tateda, C., and Takahashi, Y. (2008). Polyamines: essential factors for growth and survival. *Planta* 228, 367–381. doi: 10.1007/s00425-008-0772-7
- Laohavisit, A., Shang, Z., Rubio, L., Cuin, T. A., Véry, A. A., Wang, A., et al. (2012). Arabidopsis annexin1 mediates the radical-activated plasma membrane Ca^{2+} - and K^+ -permeable conductance in root cells. *Plant Cell* 24, 1522–1533. doi: 10.1105/tpc.112.097881

- Legocka, J., and Kluk, A. (2005). Effect of salt and osmotic stress on changes in polyamine content and arginine decarboxylase activity in *Lupinus luteus* seedlings. *J. Plant Physiol.* 162, 662–668. doi: 10.1016/j.jplph.2004.08.009
- Li, C. Z., Wang, D., and Wang, G. X. (2005). The protective effects of cobalt on potato seedling leaves during osmotic stress. *Bot. Bull. Acad. Sinica* 46, 119–125.
- Li, Z. Y., and Chen, S. Y. (2000). Differential accumulation of the S-adenosylmethionine decarboxylase transcript in rice seedlings in response to salt and drought stresses. *Theor. Appl. Genet.* 100, 782–788. doi: 10.1007/s001220051352
- Liu, H. P., Yu, B. J., Zhang, W. H., and Liu, Y. L. (2005). Effect of osmotic stress on the activity of H^+ -ATPase and the levels of covalently and non-covalently conjugated polyamines in plasma membrane preparation from wheat seedling roots. *Plant Sci.* 168, 1599–1607. doi: 10.1016/j.plantsci.2005.01.024
- Liu, J. H., Inoue, H., and Moriguchi, T. (2008). Salt stress-mediated changes in free polyamine titers and expression of genes responsible for polyamine biosynthesis of apple *in vitro* shoots. *Environ. Exp. Bot.* 62, 28–35. doi: 10.1016/j.envexpbot.2007.07.002
- Liu, J., Yu, B. J., and Liu, Y. L. (2006). Effects of spermidine and spermine levels on salt tolerance associated with tonoplast H^+ -ATPase and H^+ -PPase activities in barley roots. *Plant Growth Regul.* 49, 119–126. doi: 10.1007/s10725-006-9001-1
- Liu, K., Fu, H. H., Bei, Q. X., and Luan, S. (2000). Inward potassium channel in guard cells as a target for polyamine regulation of stomatal movements. *Plant Physiol.* 124, 1315–1325. doi: 10.1104/pp.124.3.1315
- Lopatin, A. N., Makhina, E. N., and Nichols, C. G. (1994). Potassium channel block by cytoplasmic polyamines as the mechanism of intrinsic rectification. *Nature* 372, 366–369. doi: 10.1038/372366a0
- Lu, Z., and Ding, L. (1999). Blockade of a retinal cGMP-gated channel by polyamines. *J. Gen. Physiol.* 113, 35–43. doi: 10.1085/jgp.113.1.35
- Marco, F., Alcázar, R., Tiburcio, A. F., and Carrasco, P. (2011). Interactions between polyamines and abiotic stress pathway responses unraveled by transcriptome analysis of polyamine overproducers. *OMICS* 15, 775–781. doi: 10.1089/omi.2011.0084
- Marschner, H. (1995). *Mineral Nutrition of Higher Plants*. London: Academic Press.
- Mäser, P., Thomine, S., Schroeder, J. I., Ward, J. M., Hirschi, K., Sze, H., et al. (2001). Phylogenetic relationships within cation transporter families of Arabidopsis. *Plant Physiol.* 126, 1646–1667. doi: 10.1104/pp.126.4.1646
- McAinsh, M. R., Brownlee, C., and Hetherington, A. M. (1997). Calcium ions as second messengers in guard cell signal transduction. *Physiol. Plant* 100, 16–29. doi: 10.1111/j.1399-3054.1997.tb03451.x
- Miller, G., Suzuki, N., Ciftci-Yilmaz, S., and Mittler, R. (2010). Reactive oxygen species homeostasis and signalling during drought and salinity stresses. *Plant Cell Environ.* 33, 453–467. doi: 10.1111/j.1365-3040.2009.02041.x
- Moschou, P. N., Paschalidis, K. A., and Roubelakis-Angelakis, K. A. (2008). Plant polyamine catabolism: the state of the art. *Plant Signal. Behav.* 3, 1061–1066. doi: 10.4161/psb.3.12.7172
- Moschou, P. N., and Roubelakis-Angelakis, K. A. (2014). Polyamines and programmed cell death. *J. Exp. Bot.* 65, 1285–1296. doi: 10.1093/jxb/ert373
- Mulangi, V., Cibucos, M. C., Phuntumart, V., and Morris, P. F. (2012a). Kinetic and phylogenetic analysis of plant polyamine uptake transporters. *Planta* 236, 1261–1273. doi: 10.1007/s00425-012-1668-0
- Mulangi, V., Phuntumart, V., Aoudia, M., Ramotar, D., and Morris, P. F. (2012b). Functional analysis of *OsPUT1*, a rice polyamine uptake transporter. *Planta* 235, 1–11. doi: 10.1007/s00425-011-1486-9
- Mutlu, F., and Bozcuk, S. (2005). Effects of salinity on the contents of polyamines and some other compounds in sunflower plants differing in salt tolerance. *Russ. J. Plant Physiol.* 52, 29–34. doi: 10.1007/s11183-005-0005-x
- Mutlu, F., and Bozcuk, S. (2007). Salinity-induced changes of free and bound polyamine levels in sunflower (*Helianthus annuus* L.) roots differing in salt tolerance. *Pakistan J. Bot.* 39, 1097–1102.
- Nayyar, H., Kaur, S., Smita, Singh, K. J., Dhir, K. K., and Bains, T. (2005). Water stress-induced injury to reproductive phase in chickpea: Evaluation of stress sensitivity in wild and cultivated species in relation to abscisic acid and polyamines. *J. Agron. Crop Sci. Z. Acker Pflanzenbau*. 191, 450–457. doi: 10.1111/j.1439-037X.2005.00184.x
- Ndayiragije, A., and Lutts, S. (2006). Do exogenous polyamines have an impact on the response of a salt-sensitive rice cultivar to NaCl? *J. Plant Physiol.* 163, 506–516. doi: 10.1016/j.jplph.2005.04.034
- Ndayiragije, A., and Lutts, S. (2007). Long term exogenous putrescine application improves grain yield of a salt-sensitive rice cultivar exposed to NaCl. *Plant Soil*. 291, 225–238. doi: 10.1007/s11104-006-9188-y
- Neill, S. J., Desikan, R., Clarke, A., Hurst, R. D., and Hancock, J. T. (2002). Hydrogen peroxide and nitric oxide as signaling molecules in plants. *J. Exp. Bot.* 53, 1237–1247. doi: 10.1093/jexbot/53.7.1237
- Neily, M. H., Baldet, P., Arfaoui, I., Saito, T., Li, Q. L., Asamizu, E., et al. (2011). Overexpression of apple spermidine synthase 1 (MdSPDS1) leads to significant salt tolerance in tomato plants. *Plant Biotech.* 28, 33–42. doi: 10.5511/plant-biotechnology.10.1013a
- Nemeth, M., Janda, T., Horvath, E., Paldi, E., and Szalai, G. (2002). Exogenous salicylic acid increases polyamine content but may decrease drought tolerance in maize. *Plant Sci.* 162, 569–574. doi: 10.1016/S0168-9452(01)00593-3
- Ng, C. K. Y., McAinsh, M. R., Gray, J. E., Hunt, L., Leckie, C. P., Mills, L., et al. (2001). Calcium-based signalling systems in guard cells. *New Phytol.* 151, 109–120. doi: 10.1046/j.1469-8137.2001.00152.x
- Ozawa, R., Berteau, C. M., Foti, M., Narayana, R., Arimura, G., Muroi, A., et al. (2010). Polyamines and jasmonic acid induce plasma membrane potential variations in Lima bean. *Plant Signal. Behav.* 5, 308–310. doi: 10.4161/psb.5.3.10848
- Ozturk, L., and Demir, Y. (2003). Effects of putrescine and ethephon on some oxidative stress enzyme activities and proline content in salt stressed spinach leaves. *Plant Growth Regul.* 40, 89–95. doi: 10.1023/A:1023078819935
- Palmgren, M. G. (2001). Plant plasma membrane H^+ -ATPases: powerhouses for nutrient uptake. *Annu. Rev. Plant Physiol. Plant Mol. Biol.* 52, 817–845. doi: 10.1146/annurev.arplant.52.1.817
- Pandolfi, C., Pottosin, I., Cuin, T., Mancuso, S., and Shabala, S. (2010). Specificity of polyamine effects on NaCl-induced ion flux kinetics and salt stress amelioration in plants. *Plant Cell Physiol.* 51, 422–434. doi: 10.1093/pcp/pcq007
- Parvin, S., Lee, O. R., Sathiyaraj, G., Khorolragchaa, A., Kim, Y. J., Miah, M. G., et al. (2012). Modulation of polyamine levels in ginseng hairy root cultures subjected to salt stress. *Russ. J. Plant Physiol.* 59, 757–765. doi: 10.1134/S10244371206012X
- Peiter, E., Maathuis, F. J. M., Mills, L. N., Knight, H., Pelloux, J., Hetherington, A. M., et al. (2005). The vacuolar Ca^{2+} -activated channel TPC1 regulates germination and stomatal movement. *Nature* 434, 404–408. doi: 10.1038/nature03381
- Pei, Z., Murata, Y., Benning, G., Thomine, S., Klüsener, B., Allen, G., et al. (2000). Calcium channels activated by hydrogen peroxide mediate abscisic acid signalling in guard cells. *Nature* 406, 731–734. doi: 10.1038/35021067
- Pei, Z. M., Ward, J. M., and Schroeder, J. I. (1999). Magnesium sensitizes slow vacuolar channels to physiological cytosolic calcium and inhibits fast vacuolar channels in fava bean guard cell vacuoles. *Plant Physiol.* 121, 977–986. doi: 10.1104/pp.121.3.977
- Peremarti, A., Bassie, L., Christou, P., and Capell, T. (2009). Spermine facilitates recovery from drought but does not confer drought tolerance in transgenic rice plants expressing *Datura stramonium* S-adenosylmethionine decarboxylase. *Plant Mol. Biol.* 70, 253–264. doi: 10.1007/s11103-009-9470-5
- Pistocchi, R., Keller, F., Bagni, N., and Matile, P. (1988). Transport and subcellular localization of polyamines in carrot protoplasts and vacuoles. *Plant Physiol.* 87, 514–518. doi: 10.1104/pp.87.2.514
- Piterkova, J., Luhova, L., Zajoncova, L., Sebela, M., and Petrivalsky, M. (2012). Modulation of polyamine catabolism in pea seedlings by calcium during salinity stress. *Plant Protect. Sci.* 48, 53–64.
- Pottosin, I., and Dobrovinskaya, O. (2014). Non-selective cation channels in plasma and vacuolar membranes and their contribution to K^+ transport. *J. Plant Physiol.* doi: 10.1016/j.jplph.2013.11.013. [Epub ahead of print].
- Pottosin, I., Velarde-Buendía, A. M., Bose, J., Zepeda-Jazo, I., Shabala, S., and Dobrovinskaya, O. (2014a). Cross-talk between ROS and polyamines in regulation of ion transport across plasma membrane: implications for plant adaptive responses. *J. Exp. Bot.* 65, 1271–1283. doi: 10.1093/jxb/ert423
- Pottosin, I., Velarde-Buendía, A. M., Bose, J., Fuglsang, A. T., and Shabala, S. (2014b). Polyamines cause plasma membrane depolarization, activate Ca^{2+} -, and modulate H^+ -ATPase pump activity in pea roots. *J. Exp. Bot.* doi: 10.1093/jxb/eru133. [Epub ahead of print].
- Pottosin, I., Velarde-Buendía, A. M., Zepeda-Jazo, I., Dobrovinskaya, O., and Shabala, S. (2012). Synergism between polyamines and ROS in the induction of Ca^{2+} and K^+ fluxes in roots. *Plant Signal. Behav.* 7, 1084–1087. doi: 10.4161/psb.21185

- Pottosin, I. I., Dobrovinskaya, O. R., and Muñiz, J. (2001). Conduction of mono-valent and divalent cations in the slow vacuolar channel. *J. Membr. Biol.* 181, 55–65. doi: 10.1007/s002320010007
- Pottosin, I. I., Martínez-Estévez, M., Dobrovinskaya, O. R., and Muñiz, J. (2003). Potassium-selective channel in the red beet vacuolar membrane. *J. Exp. Bot.* 54, 663–667. doi: 10.1093/jxb/erg067
- Pottosin, I. I., and Muñiz, J. (2002). Higher plant vacuolar ionic transport in the cellular context. *Acta Bot. Mex.* 60, 37–77.
- Pottosin, I. I., and Schönknecht, G. (2007). Vacuolar calcium channels. *J. Exp. Bot.* 58, 1559–1569. doi: 10.1093/jxb/erm035
- Pottosin, I. I., Tikhonova, L. I., Hedrich, R., and Schönknecht, G. (1997). Slowly activating vacuolar channels can not mediate Ca^{2+} -induced Ca^{2+} release. *Plant J.* 12, 1387–1398. doi: 10.1046/j.1365-313x.1997.12061387.x
- Qi, Y. C., Wang, F. F., Zhang, H., and Liu, W. Q. (2010). Overexpression of suadea salsa S-adenosylmethionine synthetase gene promotes salt tolerance in transgenic tobacco. *Acta Physiol. Plant.* 32, 263–269. doi: 10.1007/s11738-009-0403-3
- Radhakrishnan, R., and Lee, I. J. (2013). Spermine promotes acclimation to osmotic stress by modifying antioxidant, abscisic acid, and jasmonic acid signals in soybean. *J. Plant Growth Regul.* 32, 22–30. doi: 10.1007/s00344-012-9274-8
- Radyukina, N. L., Mapelli, S., Ivanov, Y. V., Kartashov, A. V., Brambilla, I., and Kuznetsov, V. V. (2009). Homeostasis of polyamines and antioxidant systems in roots and leaves of *Plantago major* under salt stress. *Russ. J. Plant Physiol.* 56, 323–331. doi: 10.1134/S1021443709030042
- Radyukina, N. L., Shashukova, A. V., Mapelli, S., Shevyakova, N. I., and Kuznetsov, V. V. (2010). Proline controls the level of polyamines in common sage plants under normal conditions and at UV-B irradiation. *Russ. J. Plant Physiol.* 57, 422–429. doi: 10.1134/S1021443710030155
- Reggiani, R., Aurisano, N., Mattana, M., and Bertani, A. (1993). Effect of K^+ ions on polyamine levels in wheat seedlings under anoxia. *J. Plant Physiol.* 142, 94–98. doi: 10.1016/S0176-1617(11)80113-6
- Reggiani, R., Bozo, S., and Bertani, A. (1994). Changes in polyamine metabolism in seedlings of 3 wheat (*Triticum aestivum* L.) cultivars differing in salt sensitivity. *Plant Sci.* 102, 121–126. doi: 10.1016/0168-9452(94)90028-0
- Reggiani, R., Hochkoepller, A., and Bertani, A. (1989). Polyamines in rice seedlings under oxygen-deficit stress. *Plant Physiol.* 91, 1197–1201. doi: 10.1104/pp.91.3.1197
- Reggiani, R., Zaina, S., and Bertani, A. (1992). Plasmalemma ATPase in rice coleoptiles: stimulation by putrescine and polyamines. *Phytochemistry* 31, 417–419. doi: 10.1016/0031-9422(92)90009-F
- Rodrigo-Moreno, A., Poschenrieder, C., and Shabala, S. (2013). Transition metals: a double edge sword in ROS generation and signaling. *Plant Signal. Behav.* 8, e23425. doi: 10.4161/psb.23425
- Rodríguez, A. A., Maiale, S. J., Menéndez, A. B., and Ruiz, O. A. (2009). Polyamine oxidase activity contributes to sustain maize leaf elongation under saline stress. *J. Exp. Bot.* 60, 4249–4262. doi: 10.1093/jxb/erp256
- Roussos, P. A., and Pontikis, C. A. (2007). Changes of free, soluble conjugated and bound polyamine titers of jojoba explants under sodium chloride salinity *in vitro*. *J. Plant Physiol.* 164, 895–903. doi: 10.1016/j.jplph.2006.05.003
- Roy, M., and Wu, R. (2001). Arginine decarboxylase transgene expression and analysis of environmental stress tolerance in transgenic rice. *Plant Sci.* 160, 869–875. doi: 10.1016/S0168-9452(01)00337-5
- Roy, P., Niyogi, K., SenGupta, D. N., and Ghosh, B. (2005). Spermidine treatment to rice seedlings recovers salinity stress-induced damage of plasma membrane and PM-bound H^+ -ATPase in salt-tolerant and salt-sensitive rice cultivars. *Plant Sci.* 168, 583–591. doi: 10.1016/j.plantsci.2004.08.014
- Roychoudhury, A., Basu, S., and Sengupta, D. N. (2011). Amelioration of salinity stress by exogenously applied spermidine or spermine in three varieties of indica rice differing in their level of salt tolerance. *J. Plant Physiol.* 168, 317–328. doi: 10.1016/j.jplph.2010.07.009
- Ruiz-Carrasco, K., Antognoni, F., Coulibaly, A. K., Lizardi, S., Covarrubias, A., Martínez, E. A., et al. (2011). Variation in salinity tolerance of four lowland genotypes of quinoa (*Chenopodium quinoa* Willd.) as assessed by growth, physiological traits, and sodium transporter gene expression. *Plant Physiol. Biochem.* 49, 1333–1341. doi: 10.1016/j.plaphy.2011.08.005
- Sagor, G. H. M., Berberich, T., Takahashi, Y., Niitsu, M., and Kusano, T. (2013). The polyamine spermine protects Arabidopsis from heat stress-induced damage by increasing expression of heat shock-related genes. *Transgenic Res.* 22, 595–605. doi: 10.1007/s11248-012-9666-3
- Sarjala, T. (1996). Growth, potassium and polyamine concentrations of Scots pine seedlings in relation to potassium availability under controlled growth conditions. *J. Plant Physiol.* 147, 593–598. doi: 10.1016/S0176-1617(96)80051-4
- Setter, T., and Waters, I. (2003). Review of prospects for germplasm improvement for waterlogging tolerance in wheat, barley and oats. *Plant Soil* 253, 1–34. doi: 10.1023/A:1024573305997
- Shabala, S. (2011). Physiological and cellular aspects of phytotoxicity tolerance in plants: the role of membrane transporters and implications for crop breeding for waterlogging tolerance. *New Phytol.* 190, 289–298. doi: 10.1111/j.1469-8137.2010.03575.x
- Shabala, S. (2012). *Plant Stress Physiology*. Oxford: CAB International. doi: 10.1079/9781845939953.0000
- Shabala, S. (2013). Learning from halophytes: physiological basis and strategies to improve abiotic stress tolerance in crops. *Ann. Bot.* 112, 1209–1221. doi: 10.1093/aob/mct205
- Shabala, S., and Cuin, T. A. (2008). Potassium transport and plant salt tolerance. *Physiol. Plantar.* 133, 651–669. doi: 10.1111/j.1399-3054.2007.01008.x
- Shabala, S., Cuin, T. A., and Pottosin, I. I. (2007a). Polyamines prevent NaCl-induced K^+ efflux from pea mesophyll by blocking non-selective cation channels. *FEBS Lett.* 581, 1993–1999. doi: 10.1016/j.febslet.2007.04.032
- Shabala, S., Cuin, T. A., Prismall, L., and Nemchinov, L. G. (2007b). Expression of animal CED-9 anti-apoptotic gene in tobacco modifies plasma membrane ion fluxes in response to salinity and oxidative stress. *Planta* 227, 189–197. doi: 10.1007/s00425-007-0606-z
- Shabala, S., Demidchik, V., Shabala, L., Cuin, T. A., Smith, S. J., Miller, A. J., et al. (2006). Extracellular Ca^{2+} ameliorates NaCl-induced K^+ loss from Arabidopsis root and leaf cells by controlling plasma membrane K^+ -permeable channels. *Plant Physiol.* 141, 1653–1665. doi: 10.1104/pp.106.082388
- Shabala, S., and Munns, R. (2012). “Salinity stress: physiological constraints and adaptive mechanisms,” in *Plant Stress Physiology*, ed S. Shabala (Oxford: CAB International), 59–93. doi: 10.1079/9781845939953.0059
- Shabala, S., and Pottosin, I. (2014). Regulation of potassium transport in plants under hostile conditions: implications for abiotic and biotic stress tolerance. *Physiol. Plantar.* doi: 10.1111/ppl.12165. [Epub ahead of print].
- Sharma, D. K., Dubey, A. K., Srivastav, M., Singh, A. K., Pandey, R. N., and Dahuja, A. (2013b). Effect of paclobutrazol and putrescine on antioxidant enzymes activity and nutrients content in salt tolerant citrus rootstock sour orange under sodium chloride stress. *J. Plant Nutr.* 36, 1765–1779. doi: 10.1080/01904167.2013.807823
- Sharma, T., Dreyer, I., and Riedelsberger, J. (2013a). The role of K^+ channels in uptake and redistribution of potassium in the model plant *Arabidopsis thaliana*. *Front. Plant Sci.* 4:224. doi: 10.3389/fpls.2013.00224
- Shevyakova, N. I., Musatenko, L. I., Stetsenko, L. A., Vedenicheva, N. P., Voitenko, L. P., Sytnik, K. M., et al. (2013). Effects of abscisic acid on the contents of polyamines and proline in common bean plants under salt stress. *Russ. J. Plant Physiol.* 60, 200–211. doi: 10.1134/S102144371301007X
- Shevyakova, N. I., Shorina, M. V., Rakitin, V. Y., and Kuznetsov, V. V. (2006). Stress-dependent accumulation of spermidine and spermine in the halophyte *Mesembryanthemum crystallinum* under salinity conditions. *Russ. J. Plant Physiol.* 53, 739–745. doi: 10.1134/S1021443706060021
- Shi, J., Fu, X. Z., Peng, T., Huang, X. S., Fan, Q. J., and Liu, J. H. (2010). Spermine pretreatment confers dehydration tolerance of citrus *in vitro* plants via modulation of antioxidative capacity and stomatal response. *Tree Physiol.* 30, 914–922. doi: 10.1093/treephys/tpq030
- Shi, K., Dine, X. T., Dong, D. K., Zhou, Y. H., and Yu, J. Q. (2008). Putrescine enhancement of tolerance to root-zone hypoxia in *Cucumis sativus*: a role for increased nitrate reduction. *Funct. Plant Biol.* 35, 337–345. doi: 10.1071/FP08029
- Sokolovski, S., and Blatt, M. R. (2004). Nitric oxide block of outward-rectifying K^+ channels indicates direct control by protein nitrosylation in guard cells. *Plant Physiol.* 136, 4275–4284. doi: 10.1104/pp.104.050344
- Stetsenko, L. A., Rakitin, V. Y., Shevyakova, N. I., and Kuznetsov, V. V. (2009). Organ-specific changes in the content of free and conjugated polyamines in *Mesembryanthemum crystallinum* plants under salinity. *Russ. J. Plant Physiol.* 56, 808–813. doi: 10.1134/S1021443709060107
- Su, G. X., and Bai, X. (2008). Contribution of putrescine degradation to proline accumulation in soybean leaves under salinity. *Biol. Plant* 52, 796–799. doi: 10.1007/s10535-008-0156-7

- Sudha, G., and Ravishankar, G. A. (2003). Influence of putrescine on anthocyanin production in callus cultures of *Daucus carota* mediated through calcium ATPase. *Acta Physiol. Plantar.* 25, 69–75. doi: 10.1007/s11738-003-0038-8
- Sun, C., Liu, Y. L., and Zhang, W. H. (2002). Mechanism of the effect of polyamines on the activity of tonoplasts of barley roots under salt stress. *Acta Bot. Sin.* 44, 1167–1172.
- Sziderics, A. H., Oufir, M., Trognitz, F., Kopecky, D., Matusikova, I., Hausman, J. F., et al. (2010). Organ-specific defence strategies of pepper (*Capsicum annuum* L.) during early phase of water deficit. *Plant Cell Rep.* 29, 295–305. doi: 10.1007/s00299-010-0822-z
- Takahashi, T., and Kakehi, J. (2010). Polyamines: ubiquitous polycations with unique roles in growth and stress responses. *Ann. Bot.* 105, 1–6. doi: 10.1093/aob/mcp259
- Takeda, S., Gapper, C., Kaya, H., Bell, E., Kuchitsu, K., and Dolan, L. (2008). Local positive feedback regulation determines cell shape in root hair cells. *Science* 319, 1241–1244. doi: 10.1126/science.1152505
- Tang, W., and Newton, R. J. (2005). Polyamines reduce salt-induced oxidative damage by increasing the activities of antioxidant enzymes and decreasing lipid peroxidation in Virginia pine. *Plant Growth Regul.* 46, 31–43. doi: 10.1007/s10725-005-6395-0
- Tikhonova, L. I., Pottosin, I. I., Dietz, K. J., and Schönknecht, G. (1997). Fast-activating cation channel in barley mesophyll vacuoles: inhibition by calcium. *Plant J.* 11, 1059–1070. doi: 10.1046/j.1365-313X.1997.11051059.x
- Tisi, A., Angelini, R., and Cona, A. (2011). Does polyamine catabolism influence root development and xylem differentiation under stress conditions? *Plant Signal. Behav.* 6, 1844–1487. doi: 10.4161/psb.6.11.17640
- Toumi, I., Moschou, P. N., Paschalidis, K. A., Bouamama, B., Ben Salem-Fnayou, A., Ghorbel, A. W., et al. (2010). Absciscic acid signals reorientation of polyamine metabolism to orchestrate stress responses via the polyamine exodus pathway in grapevine. *J. Plant Physiol.* 167, 519–525. doi: 10.1016/j.jplph.2009.10.022
- Tun, N. N., Santa-Catarina, C., Begum, T., Silveria, V., Handro, W., Floh, E. I. S., et al. (2006). Polyamines induce rapid biosynthesis of nitric oxide (NO) in *Arabidopsis thaliana* seedlings. *Plant Cell Physiol.* 47, 346–354. doi: 10.1093/pcp/pci252
- Uehara, A., Fill, M., Velez, P., Yasukochi, M., and Imanaga, I. (1996). Rectification of rabbit cardiac ryanodine receptor current by endogenous polyamines. *Biophys. J.* 71, 769–777. doi: 10.1016/S0006-3495(96)79276-7
- Upreti, K. K., and Murti, G. S. R. (2010). Response of grape rootstocks to salinity: changes in root growth, polyamines and abscisic acid. *Biol. Plant* 54, 730–734. doi: 10.1007/s10535-010-0130-z
- van der Mescht, A., de Ronde, J. A., van der Merwe, T., and Rossouw, F. T. (1998). Changes in free proline concentrations and polyamine levels in potato leaves during drought stress. *S. Afr. J. Sci.* 94, 347–350.
- Velarde-Buendía, A. M. (2013). *Remodelación del Transporte Membranal por Especies Reactivas de Oxígeno y Poliaminas en Tejido Radicular*. Dissertation, Universidad de Colima, Mexico.
- Velarde-Buendía, A. M., Shabala, S., Cvikrova, M., Dobrovinskaya, O., and Pottosin, I. (2012). Salt-sensitive and salt-tolerant barley varieties differ in the extent of potentiation of the ROS-induced K⁺ efflux by polyamines. *Plant Physiol. Biochem.* 61, 18–23. doi: 10.1016/j.plaphy.2012.09.002
- Voisenek, L., and Sasidharan, R. (2013). Ethylene - and oxygen signalling - drive plant survival during flooding. *Plant Biol.* 15, 426–435. doi: 10.1111/plb.12014
- Volkov, V., Wang, B., Dominy, P. J., Fricke, W., and Amtmann, A. (2003). *Thellungiella halophila*, a salt tolerant relative of *Arabidopsis thaliana*, possesses effective mechanisms to discriminate between potassium and sodium. *Plant Cell Environ.* 27, 1–14. doi: 10.1046/j.0016-8025.2003.01116.x
- Waie, B., and Rajam, M. V. (2003). Effect of increased polyamine biosynthesis on stress responses in transgenic tobacco by introduction of human S-adenosylmethionine gene. *Plant Sci.* 164, 727–734. doi: 10.1016/S0168-9452(03)00030-X
- Wang, B. Q., Zhang, Q. F., Liu, J. H., and Li, G. H. (2011). Overexpression of PtADC confers enhanced dehydration and drought tolerance in transgenic tobacco and tomato: effect on ROS elimination. *Biochem. Biophys. Res. Commun.* 413, 10–16. doi: 10.1016/j.bbrc.2011.08.015
- Ward, J. M., and Schroeder, J. I. (1994). Calcium-activated K⁺ channels and calcium-induced calcium-release by slow vacuolar ion channels in guard-cell vacuoles implicated in the control of stomatal closure. *Plant Cell* 6, 669–683. doi: 10.2307/3869871
- Watson, M. B., and Malmberg, R. L. (1996). Regulation of *Arabidopsis thaliana* (L.) Heynh arginine decarboxylase by potassium deficiency stress. *Plant Physiol.* 111, 1077–1083. doi: 10.1104/pp.111.4.1077
- Williams, K. (1997). Interactions of polyamines with ion channels. *Biochem. J.* 385, 289–297.
- Wimalasekera, R., Tebart, F., and Scherer, G. F. E. (2011). Polyamines, polyamine oxidases and nitric oxide in development, abiotic and biotic stresses. *Plant Sci.* 181, 593–603. doi: 10.1016/j.plantsci.2011.04.002
- Wu, Q. S., Peng, Y. H., Zou, Y. N., and Liu, C. Y. (2010). Exogenous polyamines affect mycorrhizal development of *Glomus mosseae*-colonized citrus (*Citrus tangerine*) seedlings. *Science Asia* 36, 254–258. doi: 10.2306/scienceasia1513-1874.2010.36.254
- Xie, L. H., John, S. A., Ribalet, B., and Weiss, J. N. (2005). Long polyamines act as cofactors in PIP2 activation of inward rectifier potassium (Kir2.1) channels. *J. Gen. Physiol.* 126, 541–549. doi: 10.1085/jgp.200509380
- Xing, S. G., Jun, Y. B., Hau, Z. W., and Liang, L. Y. (2007). Higher accumulation of gamma-aminobutyric acid induced by salt stress through stimulating the activity of diarnine oxidases in *Glycine max* (L.) Merr. roots. *Plant Physiol. Biochem.* 45, 560–566. doi: 10.1016/j.plaphy.2007.05.007
- Yamaguchi, K., Takahashi, Y., Berberich, T., Imai, A., Miyazaki, A., Takahashi, T., et al. (2006). The polyamine spermine protects against high salt stress in *Arabidopsis thaliana*. *FEBS Lett.* 580, 6783–6788. doi: 10.1016/j.febslet.2006.10.078
- Yamaguchi, K., Takahashi, Y., Berberich, T., Imai, A., Takahashi, T., Michael, A. J., et al. (2007). A protective role for the polyamine spermine against drought stress in *Arabidopsis*. *Biochem. Biophys. Res. Commun.* 352, 486–490. doi: 10.1016/j.bbrc.2006.11.041
- Yang, J. C., Zhang, J. H., Liu, K., Wang, Z. Q., and Liu, L. J. (2007). Involvement of polyamines in the drought resistance of rice. *J. Exp. Bot.* 58, 1545–1555. doi: 10.1093/jxb/erm032
- Yiu, J. C., Liu, C. W., Fang, D. Y. T., and Lai, Y. S. (2009). Waterlogging tolerance of Welsh onion (*Allium fistulosum* L.) enhanced by exogenous spermidine and spermine. *Plant Physiol. Biochem.* 47, 710–716. doi: 10.1016/j.plaphy.2009.03.007
- Zacchini, M., Marotta, A., and de Agazio, M. (1997). Tolerance to salt stress in maize callus lines with different polyamine content. *Plant Cell Rep.* 17, 119–122. doi: 10.1007/s002990050363
- Zandonadi, D. B., Santos, M. P., Dobbss, L. B., Olivares, F. L., Canellas, L. P., Binzel, M. L., et al. (2010). Nitric oxide mediates humic acids-induced root development and plasma membrane H⁺-ATPase activation. *Planta* 231, 1025–1036. doi: 10.1007/s00425-010-1106-0
- Zapata, P. J., Botella, M. A., Pretel, M. T., and Serrano, M. (2007). Responses of ethylene biosynthesis to saline stress in seedlings of eight plant species. *Plant Growth Regul.* 53, 97–106. doi: 10.1007/s10725-007-9207-x
- Zapata, P. J., Serrano, M., Pretel, M. T., Amoros, A., and Botella, A. (2003). Changes in ethylene evolution and polyamine profiles of seedlings of nine cultivars of *Lactuca sativa* L. in response to salt stress during germination. *Plant Sci.* 164, 557–563. doi: 10.1016/S0168-9452(03)00005-0
- Zapata, P. J., Serrano, M., Pretel, M. T., and Botella, M. A. (2008). Changes in free polyamine concentration induced by salt stress in seedlings of different species. *Plant Growth Regul.* 56, 167–177. doi: 10.1007/s10725-008-9298-z
- Zeng, F., Shabala, L., Zhou, M., Zhang, G. P., and Shabala, S. (2013). Barley responses to combined waterlogging and salinity stress: separating effects of oxygen deprivation and elemental toxicity. *Front. Plant Sci.* 4:313. doi: 10.3389/fpls.2013.00313
- Zepeda Jazo, A. I. (2010). *Mecanismos De Intercambio De K⁺ y Ca²⁺ Em Epidermis De Raíces Bajo Estrés Salino*. dissertation, Universidad de Colima, Mexico.
- Zepeda-Jazo, I., Shabala, S., Chen, Z., and Pottosin, I. I. (2008). Na⁺-K⁺ transport in roots under salt stress. *Plant Signal. Behav.* 3, 401–403. doi: 10.4161/psb.3.6.5429
- Zepeda-Jazo, I., Velarde-Buendia, A. M., Enriquez-Figueroa, R., Bose, J., Shabala, S., Muniz-Murguía, J., et al. (2011). Polyamines interact with hydroxyl radicals in activating Ca²⁺ and K⁺ transport across the root epidermal plasma membranes. *Plant Physiol.* 157, 2167–2180. doi: 10.1104/pp.111.179671
- Zhao, F., Song, C. P., He, J., and Zhu, H. (2007). Polyamines improve K⁺/Na⁺ homeostasis in barley seedlings by regulating root ion channel activities. *Plant Physiol.* 145, 1061–1072. doi: 10.1104/pp.107.105882

- Zhao, F. G., and Qin, P. (2004). Protective effect of exogenous polyamines on root tonoplast function against salt stress in barley seedlings. *Plant Growth Regul.* 42, 97–103. doi: 10.1023/B:GROW.0000017478.40445.bc
- Zhao, F. G., Sun, C., Liu, Y. L., and Zhang, W. H. (2003). Relationship between polyamine metabolism in roots and salt tolerance of barley seedlings. *Acta Bot. Sin.* 45, 295–300.
- Zhu, H., Ding, G. H., Fang, K., Zhao, F. G., and Qin, P. (2006). New perspective on the mechanism of alleviating salt stress by spermidine in barley seedlings. *Plant Growth Regul.* 49, 147–156. doi: 10.1007/s10725-006-9004-y

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