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# Salinity tolerance in barley during germination homologs and potential genes<sup>#</sup>

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**Abstract:** Salinity affects more than 6% of the world's total land area, causing massive losses in crop yield. Salinity inhibits plant growth and development through osmotic and ionic stresses; however, some plants exhibit adaptations through osmotic regulation, exclusion, and translocation of accumulated Na<sup>+</sup> or Cl<sup>-</sup>. Currently, there are no practical, economically viable methods for managing salinity, so the best practice is to grow crops with improved tolerance. Germination is the stage in a plant's life cycle most adversely affected by salinity. Barley, the fourth most important cereal crop in the world, has outstanding salinity tolerance, relative to other cereal crops. Here, we review the genetics of salinity tolerance in barley during germination by summarizing reported quantitative trait loci (QTLs) and functional genes. The homologs of candidate genes for salinity tolerance in *Arabidopsis*, soybean, maize, wheat, and rice have been blasted and mapped on the barley reference genome. The genetic diversity of three reported functional gene families for salt tolerance during barley germination, namely dehydration-responsive element-binding (DREB) protein, somatic embryogenesis receptor-like kinase and aquaporin genes, is discussed. While all three gene families show great diversity in most plant species, the *DREB* gene family is more diverse in barley than in wheat and rice. Further to this review, a convenient method for screening for salinity tolerance at germination is needed, and the mechanisms of action of the genes involved in salt tolerance need to be identified, validated, and transferred to commercial cultivars for field production in saline soil.

Key words: Genetics; Barley; Quantitative trait locus (QTL); Germination; Salinity tolerance; Homologous gene; Diversity https://doi.org/10.1631/jzus.B1900400 CLC number: S512.3; Q943.2

# 1 Introduction

Salinity affects about 6% of the world's total land area, including 20% of arable land and 33% of irrigated land (Shrivastava and Kumar, 2015; Machado and Serralheiro, 2017; Kuang et al., 2019; Safdar et al., 2019), causing estimated yield losses of 20% (Ashraf and Harris, 2005; Pirasteh-Anosheh et al., 2016). Furthermore, land salinisation is increasing, with 10 million ha of agricultural land destroyed annually by salt accumulation (Pimentel et al., 2004) due to several factors including the use of contaminated irrigation water, intensive farming and poor drainage, and climate change (Machado and Serralheiro, 2017; Isayenkov, 2019). Without proper and sustainable control, salinity-affected areas will increase to more than 50% of the world's total arable land by 2050 (Ashraf, 2009; Anosheh et al., 2011; Jamil et al., 2011; Emam et al., 2013).

According to Pirasteh-Anosheh et al. (2016), plants experience four types of stress under saline conditions: (1) salinity reduces water uptake due to

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the low water potential of the soil which interferes with the osmotic gradient (Munns and Tester, 2008); (2) the absorbed salt reaches a level that causes severe cellular toxicity due to low sequestration of Na<sup>+</sup> into vacuoles (Nawaz, 2007); (3) the salt interacts with minerals causing nutrient imbalance and deficiency (Nawaz, 2007); and (4) salinity accelerates the production of active oxygen radicles, such as H<sub>2</sub>O<sub>2</sub> (hydrogen peroxide),  $\bullet O_2^-$  (superoxide),  $^1O_2$  (singlet oxygen), and •OH<sup>-</sup> (hydroxyl radicle), which can damage or even kill plants (Hernández et al., 2001). Greenway and Munns (1980) classified plants into two major categories based on their salinity tolerance, namely (1) halophytes and (2) glycophytes. Halophytes have an exceptional ability to produce heteromorphic seeds that have diverse dormancy and germination capacity under saline conditions (Liu RR et al., 2018). Seed germination in glycophytes is severely inhibited under salinity due to both osmotic stress and ionic toxicity stress, unlike in halophytes that are less affected by osmotic pressure (Romo and Haferkamp, 1987; Dodd and Donovan, 1999; Zhang et al., 2010). Halophytes can better regulate the ion-gate-controlled NaCl influx into seed cells (Glenn et al., 1999; Huang et al., 2018). However, salinity-tolerant glycophytes have a lower osmotic potential than sensitive glycophytes, enabling them to absorb more water from the soil during germination (Zhang et al., 2010).

Barley, the fourth most important cereal in the world (Schulte et al., 2009; Visioni et al., 2019), is a glycophyte, but its salinity tolerance varies among genotypes (Mano and Takeda, 1997; Flowers and Hajibagheri, 2001; Xue et al., 2009; Debez et al., 2019). Some barley genotypes are able to thrive in saline conditions (Harlan, 1995; Shen et al., 2018). Salinity-tolerant barley genotypes exhibit halophytic features such as excluding Na<sup>+</sup> from uptake (Chen ZH et al., 2007) and accumulating Na<sup>+</sup> in tissues (Munns et al., 1988; Munns and Tester, 2008). Tolerant genotypes sequester Na<sup>+</sup> in their intracellular vacuoles, thereby maintaining high K<sup>+</sup>/Na<sup>+</sup> levels in the cytosol while reducing damage from Na<sup>+</sup> toxicity (Shabala et al., 2010; Mian et al., 2011; Fu et al., 2018; Han et al., 2018; Ishikawa and Shabala, 2019). They can also synthesize compatible solutes in the cytoplasm to balance the osmotic potential of vacuolar Na<sup>+</sup> (Widodo et al., 2009). Salinity tolerance is controlled by multiple genes that are expressed differently during different growth phases (Qiu et al., 2011; Ahmed et al., 2013a). Germination, which determines seedling vigour and the plant population, is the most important growth stage, but is sensitive to salinity stress (Zhang et al., 2010; Bewley et al., 2013).

Several genes have association with enhanced salinity tolerance in barley (Wu et al., 2011) and are grouped into four classes based on their function (Walia et al., 2006; Wu et al., 2011; Yin et al., 2018). (1) Genes that enhance osmotic protection, such as HvPIP2;5 (Alavilli et al., 2016), HVA1 (Lal et al., 2008), HvDREB1, HvCBF4, HvWRKY38 (Gürel et al., 2016), and reactive oxygen species (ROS)-scavenging genes that include osmoregulatory trehalose synthesis, mannitol-1-phosphate dehydrogenase (M1PD), and pyrroline-5-carboxylase synthetase (P5CS). (2) Genes controlling Na<sup>+</sup> and K<sup>+</sup> transport, such as the highaffinity potassium transporter (HKT) family (e.g. HvHKT1;5 (Hazzouri et al., 2018; Huang et al., 2019), HvHKT1;1 (Han et al., 2018), HvHKT2;1 (Mian et al., 2011; Assaha et al., 2017), HvHAK1 (Mangano et al., 2008), HvHKT1, HvHKT2 (Qiu et al., 2011)), the  $Na^{+}/H^{+}$  exchanger (*NHX*) family (*HvNax4* (Rivandi et al., 2011), and salt overly sensitive (SOS) engaged  $Na^+/H^+$  antiporters (HvSOS1 (HvNHX7), HvSOS2 (HvCIPK24), HvSOS3 (HvCBL4), HvNHX1, HVA) (Yousefirad et al., 2018; Wu et al., 2019). (3) Genes that produce regulatory proteins, such as the CBF/ DREB (C-repeat-binding protein/dehydration-responsive element-binding protein) family (e.g. HvRAF (Jung et al., 2007), HvAP2/ERF (ethylene response factor) (Guo et al., 2016), HvDREB1 (Xu et al., 2009), HvCBF4, HvWRKY38 (Gürel et al., 2016), HvDRF1 (Xue and Loveridge, 2004)) in the signalling pathways of long distance and downstream gene expression. Salinity and drought stresses induce the expression of root abundant factor (RAF), CBF3, and CBF4 from the CBF/DREB gene family in most plants. Twenty CBF genes have been identified in barley, which enhance tolerance to drought, salinity, and low temperature (Wu et al., 2011). (4) Genes that induce jasmonate (JA) biosynthesis, such as late embryogenesis abundant (LEA) protein genes (e.g. HVA1 expressed in response to water and salinity stresses, HVA22 expressed in response to dehydration, extreme temperatures, abscisic acid (ABA) secretion, and salinity stress (al-Yassin and Khademian, 2015)).

# 2 Seed germination process and roles of hormones

Seed germination requires optimum environmental factors, including water, oxygen, and temperature. Other environmental factors, such as light and nitrates, can also affect seed germination (Finch-Savage and Leubner-Metzger, 2006; Rajjou et al., 2012). The process of germination occurs in three main phases regulated by hormones, reactive nitrogen species, and ROS (Ma et al., 2017). The first phase involves the expression of genes that play a major role in cell wall metabolism. The second phase encompasses significant stimulation of hormonal and enzyme activity by genes involved in amino acid synthesis, starch metabolism, nucleic acid synthesis, protein synthesis and transport (Weitbrecht et al., 2011). The third phase involves the induction of genes for photosynthetic metabolism after radicle protrusion (Ma et al., 2017). Hormonal imbalance, caused by factors such as low temperature, drought, and salts (saline and sodic) that increase ground osmotic pressure (Bartels and Nelson, 1994), can affect the seed germination process (Lopez-Molina et al., 2001; Belin and Lopez-Molina, 2008).

Several hormones produced by plant and soil microorganisms such as bacteria play a role in inducing or breaking seed dormancy and thus contribute to the germination process (Bewley, 1997; Baskin and Baskin, 2001; Koornneef et al., 2002; Hoyle et al., 2015). Interestingly, as much as gene expression controls plant hormone activity, the reverse can be true with some hormones regulating gene expression (Miransari and Smith, 2014). For example, ABA usually induces dormancy at seed maturation and gibberellins (GAs) break dormancy during germination (Fig. 1) (Bentsink and Koornneef, 2008; Hauvermale et al., 2012). When an environmental stress such as salinity occurs during germination, ABA is produced in the seeds (Weyers and Paterson, 2001) to upregulate transcription factors (TFs) such as AB13 and AB15 that stimulate genes encoding the osmotolerance protein and block the germination process (Fedoroff, 2002; Lopez-Molina et al., 2002; Graeber et al., 2010; Miransari and Smith, 2014). However, ABA is negatively regulated by ABA-INSENSITIVE1 (ABI1) and ABI2 and the expression of the HvABA8'OH-1 gene (Ma et al., 2017). GAs release seeds from

dormancy by stimulating the production of hydrolases for the germination process (Miransari and Smith, 2014; Abido et al., 2019). The embryo synthesizes GA after imbibing water (Diaz-Mendoza et al., 2019). The GA binds to receptors to promote the breakdown of repressor of GA-like2 (RGL2), a DELL factor suppressing germination, the expression of several genes (*GAMYB*, *HvPTR*, *WRKY*, *PP2C*, *GATA*, and *HvKAO1*), and the production of essential proteins during germination in stress environments (Marrs, 1996; Sun and Gubler, 2004; Ma et al., 2017). The effect of GA is terminated by the expression of GA 2-oxidase 1 (*HvGA2ox*), which is responsible for its inactivation (Fig. 1) (Ma et al., 2017).

During germination, the concentration of ethylene increases (Yang and Hoffman, 1984; Pennazio and Roggero, 1991; Petruzzelli et al., 2000), which reduces plant growth under saline conditions in the presence of ABA (Matilla, 2000; Rinaldi, 2000; Jalili et al., 2009). Auxins (indole-3-acetic acids (IAAs)) are essential for cell elongation, and radicle and embryo growth during germination (Popko et al., 2010; Hauvermale et al., 2012). Brassinosteroids (BRs) and IAA on the other hand stimulate the secretion of ethylene which works in conjunction with GAs to induce germination (Arora, 2005; Miransari and Smith, 2014). Auxins reduce seed sensitivity to ABA by overexpressing microRNAs (Liu et al., 2007) and interacting with GAs to counteract ABA suppression during germination (Chiwocha et al., 2005; Hentrich et al., 2013). Cytokinins interacting with ethylene enhance cell division and alleviate abiotic stresses like salinity during germination (Chiwocha et al., 2005; Subbiah and Reddy, 2010; Peleg and Blumwald, 2011; Miransari and Smith, 2014). BRs, in conjunction with GAs and ethylene, improve seed resistance to abiotic stress, enhance embryo growth out of the seed, and reduce the effects of ABA (Bajguz and Hayat, 2009; Miransari and Smith, 2014; Procházka et al., 2015) and salt stress (Vázquez et al., 2019) during germination. JAs are signalling molecules for plant defence against osmotic stress caused by salt (Kazan and Manners, 2012; Nguyen et al., 2019). They obstruct the production of two primary ABA biosynthesis genes (TaNCED1 and TaNCED2) with acetylsalicylic acid (ASA) in stress germination in wheat (Xu et al., 2016).



**Fig. 1** Interactions among the hormones and gene regulatory pathways in barley during germination under salinity stress Blue lines indicate hormonal regulation, while black lines are for genes and transcription factor expression. Red double-arrowed dotted line shows the negative interaction between ABA and GA during germination, while blue double-arrowed line shows the positive interaction among the hormones. Blue dotted lines show the stimulation outcome of the hormonal interactions, while black dotted lines show the regulatory interactions of genes. *GAMYB*: GA-induced Myb (myeloblastosis)-like protein; *HvPTR*: barley scutellar peptide transporter; *HvKAO1*: barley kaurenoic acid oxidase 1; *HvGA2ox*: barley GA 2-oxidase 1; *HvNCED*: barley nine-*cis*-epoxycarotenoid dioxygenase; *PP2C*: protein phosphatase type 2C; *ABI1*: ABA-insensitive 1; *HvABA8'OH-1*: barley ABA 8' hydroxylase; *HvCBL*: barley calcineurin B-like protein; *HvSERK*: barley somatic embryogenesis receptor-like kinase; *HvCBF*: barley C-repeat-binding protein; *HvDREB*: barley dehydration-responsive element-binding protein; *HvRAF*: barley root abundant factor; *HvPIP2;5*: barley aquaporin 2;5

# 3 Salinity versus sodicity

Saline soils predominantly contain sodium (Na<sup>+</sup>) cations and chloride (Cl<sup>-</sup>) anions and, in some cases, Ca<sup>2+</sup>, Mg<sup>2+</sup>, SO<sub>4</sub><sup>2-</sup>, but not HCO<sub>3</sub><sup>-</sup> or CO<sub>3</sub><sup>2-</sup> (Abrol et al., 1988). Saline soils have good structure, an advantage for proper tillage and crop cultivation, and usually an electric conductivity extract (ECE) higher than 4 dS/m (40 mmol/L), an exchangeable watersoluble sodium of >15%, and a pH of <8.5 but >7. However, when the surface of the ground appears black, hard, and dry, the soil is sodic. In this case, the predominant cation is Na<sup>+</sup>, but the primary anions are Cl<sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, HCO<sub>3</sub><sup>-</sup> and a small amount of CO<sub>3</sub><sup>2-</sup>. Sodic soils have an exchangeable Na<sup>+</sup> of >15%, an ECE of >4 dS/m, and a pH of >8.5 (Abrol et al., 1988).

# 3.1 Causes and types of salinity

Soil salinity is caused by (1) natural or primary, and (2) secondary or human-induced events (Manchanda and Garg, 2008; Parihar et al., 2015) and is either (1) groundwater-associated (dryland salinity), (2) non-groundwater-associated (transient salinity), or (3) irrigation-associated (Ghassemi et al., 1995; Rengasamy, 2006; Majeed and Muhammad, 2019). Transient

salinity fluctuates with soil depth and its effect on plant growth is dependent on rainfall. It occurs in semi-arid environments where there is insufficient rain to leach the soil. Clay layers below the topsoil hinder the movement of water and salts, leading to ion accumulation on the soil surface (Rengasamy, 2002, 2006). Most of Western Australia is characterized by a temperate climate. Ion accumulation in the topsoil following dry, hot summers can result in high salinity at sowing, which affects seed germination. These ions can be leached only by rainfall or irrigation. Dryseeding or seeding with the first rain increases the likelihood that germinating seeds will be affected by salinity stress.

# 3.2 Effect of salinity on germination

Salinity affects the seed germination process by altering water imbibition, changing enzymatic activities causing ionic toxicity, interfering with protein metabolism, causing hormonal imbalances, and reducing the possibility of seeds using their reserves, all of which delay and reduce the number of sprouting seeds (Läuchli and Grattan, 2007; Bordi, 2010; Munns et al., 2012; Parihar et al., 2015; Debez et al., 2019). Saline conditions create an external osmotic stress that reduces water availability and uptake (Bernstein, 1963; Bliss et al., 1986; Sayar et al., 2010; Sabagh et al., 2019) and increases the toxic absorption of ions (Hampson and Simpson, 1990), which decreases seed germination (Dodd and Donovan, 1999; Luan et al., 2014; Narsing Rao et al., 2019; Polash et al., 2019).

# 3.3 Salinity tolerance mechanisms

Plants overcome salinity stress through various mechanisms that involve osmotic adjustment, Na<sup>+</sup> exclusion, and tissue tolerance (Munns and Tester, 2008; Liang et al., 2018; Miransari and Smith, 2019). These encompass biochemical, physiological, and molecular mechanisms of variable complexity (Han et al., 2015; Pirasteh-Anosheh et al., 2016; Kumari et al., 2019). During germination, plants become saline-tolerant by (1) excluding salts from seed cells while maintaining high osmotic potential using organic solutes, or (2) accumulating salt ions in seed cells to increase osmotic potential while putting mechanisms in place to mitigate toxicity (Zhang et al., 2010). Seeds of salt-tolerant barley genotypes take up sodium to increase osmotic potential to absorb water during germination under salinity stress (Zhang et al., 2010), while minimising  $K^+$  losses (al-Karaki, 2001). The vacuole is a vital cell organelle for compound deposition during osmotic stress regulation, and its size is a measure of salinity tolerance (Lauchli and Epstein, 1990; Volkmar et al., 1998; Yarra, 2019; Yarra and Kirti, 2019).

#### 3.3.1 Indicators of salinity tolerance

Salinity tolerance indicators in plants can be divided into three main groups: (1) agronomic/ morphological, (2) physiological, and (3) biochemical. 3.3.1.1 Agronomic/morphological indicators

Salinity tolerance is expressed externally through visible morphological/agronomic traits as a result of several physio-chemical processes that take place inside the plant. They are a reflection of genetic and physiological mechanisms influenced by the environmental effect on the plant that confer salinity tolerance (Ashraf and Harris, 2004). They occur at the whole plant or organ level and are mostly physically visible. These indicators are easy to measure and include germination percentage, yield, survival rate/ percentage, plant height, leaf area, leaf injury, relative growth rate, and relative growth reduction. Distinguishing between tolerant and non-tolerant plants using agronomic indicators can be subjective.

3.3.1.2 Physiological indicators

Like the other indicator groups, physiological indicators are the result of processes that take place inside plant tissues, organs, cells, and organelles when exposed to salinity stress. They include the transportation of excess ions to the vacuole or sequestering them in older tissues. This group of indicators is measured by traits that include relative growth rate, germination speed, ion homeostasis, photosynthesis, transpiration, and senescence (Negrão et al., 2017). They provide more objective information than morphological indicators when combined with knowledge of the genetic model of salinity tolerance (Ashraf and Harris, 2004). Sodium ions are transported and compartmentalized in the vacuoles using two types of  $H^+$  pumps (V-ATPase and  $H^+$ -ATPase) and vacuolar pyrophosphatase (V-PPase). The pumps are facilitated by SOS pathways with three types of proteins (SOS<sub>1</sub>, SOS<sub>2</sub>, and SOS<sub>3</sub>). SOS<sub>1</sub> is essential for regulating Na<sup>+</sup> efflux at the cellular level and enhancing Na<sup>+</sup> transportation in the organelles and tissues. The SOS<sub>2</sub> can not only interact with SOS<sub>3</sub> and subsequently activate SOS1, but also increases transport activity of proteins such as NHX (Gupta and Huang, 2014). Electron transport chains in mitochondria can overflow, deregulate, or become disrupted by salinity stress leading to the accumulation of toxic compounds. Antioxidant enzymes and nonenzymatic compounds are essential for detoxifying ROS, which include helicase proteins, catalase (CAT), peroxidase, polyphenol, flavonoid, ascorbate, and glutathione (Gupta and Huang, 2014).

# 3.3.1.3 Biochemical indicators

Biochemical indicators are chemical solutes that build up inside plants in response to salt stress (Ashraf and Harris, 2004). The accumulation of low molecular weight organic solutes, inorganic ions, compatible osmolytes, soluble sugars, soluble proteins, amino acids and amides, quaternary ammonium compounds, polyamines (PAs), polyols, antioxidants, and ATPases is associated with tolerance to salinity stress. Secretion of biochemical compounds occurs during healthy growth and germination; hence it is not easy to tell whether a chemical is a reaction or tolerance to salinity stress, or an adaptive plant mechanism (Ashraf and Harris, 2004). Plant compatible osmolytes are organic compounds synthesized by various plant species which accumulate in variable amounts and are soluble in cells and do not disturb cellular activities at high or low concentrations (Hanin et al., 2016). These compounds, including proline, glycine betaine, sugars, and polyols, increase within the cell in response to salinity stress, and are sustained by permanent synthesis and/or degradation. Accumulation of these compounds is proportional to the surrounding osmolarity, and they function to shield the cell structure and maintain osmotic balance through constant water entry (Hasegawa et al., 2000; Mansour and Ali, 2017; Alhasnawi, 2019).

The secretion of PAs has a diverse function in healthy plant growth, including seed germination, and is critical for abiotic stress adaptation, including salinity. Diamine putrescine (PUT), tetra-amine spermine (SPM), and triamine spermidine (SPD) are some typical PAs found in plants (Shu et al., 2012) and whose increase has been associated with salinity stress (Gupta and Huang, 2014). Nitric oxide (NO) regulates several plant growth and developmental activities, stress signalling molecules and stress responses, including salinity and the activation of antioxidant enzymes (superoxide dismutase (SOD), CAT, guaiacol peroxidase (GPX), ascorbate peroxidase (APX), and glutathione reductase (GR)), and triggers the expression of various redox-regulated genes that enhance germination and root growth under salinity stress (Gupta and Huang, 2014). Salinity stress mitigation by NO is due to its antioxidant functions, modulation of the ROS detoxification system (Mishra et al., 2011) associated with an increase in antioxidant enzymes, such as SOD, CAT, GPX, APX, and GR (Zhao et al., 2004; Ali et al., 2018), and suppression of malondialdehyde (MDA) production during lipid peroxidation (Nalousi et al., 2012). NO offers a force for  $Na^+/H^+$  exchange, providing the H<sup>+</sup> gradient to stimulate H<sup>+</sup>-ATPase (H<sup>+</sup>-PPase) and contributing to K<sup>+</sup> and Na<sup>+</sup> homeostasis (Zhang et al., 2006; Gupta and Huang, 2014).

Other biochemical indicators include the production of several hormones in response to salinity stress, including ABA in roots and shoots, which mitigates the antagonistic effect of salinity stress on physiological processes (Popova et al., 1995; Jaschke et al., 1997). ABA accumulation triggers the expression of salinity stress tolerance enhancing genes in cultivated barley (*HVP1*, *HVP10*, and *HvVHA-A*) and wheat (*MAPK4-like*, *TIP 1*, and *GLP 1*) (Keskin et al., 2010). Salicylic acid (SA) concentration, along with the SA biosynthetic enzyme, increases under salinity stress in rice seedlings (Jayakannan et al., 2013). Application of BR promotes antioxidant enzyme (SOD, POX, APX, and GPX) activity and the accumulation of salinity stress mitigating nonenzymatic antioxidant compounds, such as tocopherol, ascorbate, and reduced glutathione (Ashraf et al., 2010; el-Mashad and Mohamed, 2012; Gupta and Huang, 2014).

# 3.3.2 Salinity stress signalling pathways and molecules

The response to salinity stress in plants involves three types of signalling pathway: (1) ionic and osmotic stress signalling (homeostasis) essential for the re-establishment of cellular homeostasis, (2) detoxification signalling that regulates and repairs damage, and (3) cell division and expansion signalling (Zhu, 2001). Homeostatic signalling governs detoxification leading to tolerance by reducing the response of growth inhibitors (Zhu, 2002). The signalling information flows in a way that ionic (Na<sup>+</sup>), osmotic (turgor pressure change), and detoxification are stress input, while salinity-induced damage, control, and repair are output (response) (Xiong and Zhu, 2001; Zhu, 2002). The outcomes of osmotic signalling include gene expression or activation of osmolyte biosynthesis enzymes and the water and osmolyte transport system (Zhu, 2001). Detoxification signalling changes induced by salt include phospholipid hydrolysis, changes in the expression of LEA/dehydrintype genes, molecular chaperones, and proteinase to remove denatured proteins, and the activation of enzymes for the generation and removal of ROS and other detoxification proteins (Zhu, 2001, 2002).

Signalling molecules are crucial components for salinity stress tolerance in plants, and they determine downstream actions involving protein phosphorylation, dephosphorylation, phospholipid metabolism, and calcium ion  $(Ca^{2+})$  sensing (Agarwal et al., 2013).  $Ca^{2+}$  signalling is one of the first responses to salt stress and is essential for ion homeostasis (Zhu, 2003; Reddy and Reddy, 2004). It is sensed by calcineurin B-like proteins (CBLs) that are involved in the salt stress indication transduction pathway and govern the influx and efflux of Na<sup>+</sup>. CBLs increase germination

under salinity stress in tobacco (Pardo et al., 1998), Arabidopsis (Liu and Zhu, 1998), and maize (Wang MY et al., 2007). Other molecules include mitogenactivated protein kinase (SAPK) (Diédhiou et al., 2008), CBL-interacting protein kinase (CIPK), nucleoside diphosphate kinase (NDPK), and mitogenactivated protein (MAP) kinase (MPK) (Moon et al., 2003), whose overexpression enhances salinity tolerance capacity by increasing seed germination. Initiation of the MPK cascade is the second response to decode external salinity stress into cellular responses (Teige et al., 2004). This includes signalling modules that phosphorylate particular serine/threonine residues on amide protein substrates that regulate several cellular processes (MAPK Group et al., 2002). Overexpression of MAP in rice (Xiong and Yang, 2003; Jeong et al., 2006) and maize MAP in Arabidopsis (Kong et al., 2011) enhances salinity tolerance and activates transcriptional factors controlling downstream genes such as COR47, RD29A, P5CS2 (Wurzinger et al., 2011), and CDPK (Capiati et al., 2006).

# 3.3.3 Salinity stress tolerance: transcriptional regulation and gene expression

Salt stress prompts the induction of various genes that fall into two broad groups: (1) single function genes that facilitate the production of protective metabolites, and (2) regulatory genes that control the expression of downstream genes (Agarwal et al., 2013). Single function genes promote the release of compounds such as transporters/channel proteins, osmolytes, lipid biosynthesis genes, antioxidative enzymes, and PAs, while DREB, bZIP, NAC, and MYC/ MYB (myelocytomatosis/myeloblastosis) are regulatory genes (Shinozaki and Yamaguchi-Shinozaki, 2007; Agarwal and Jha, 2010). Many of these genes from both groups play different roles in enhancing the adaptation process, and are further categorized into functional groups that include: (1) senescenceassociated genes (SAGs), (2) ion transport or homeostasis genes (SOS, HKT, AtNHX1, and  $H^+$ -ATPase), (3) molecular chaperones (HSP genes), and (4) dehydration-related TFs (DREB) (Hasegawa et al., 2000; Liu et al., 2000; Shi et al., 2000; Yen et al., 2000). Under osmotic pressure from salinity stress, plants synthesize novel proteins that increase the

expression of genes, such as osmotic regulation (OR) genes, at the cell level to aid their adaptation. The OR genes are categorized into nine groups, namely LEA genes, osmolyte biosynthesis genes, transporter genes, OR genes that encode regulatory proteins, photosynthetic genes, OR genes encoding proteins involved in protein synthesis processes and degradation, heat shock protein genes, osmotins, and other protein genes (Zhu et al., 1997; Turan et al., 2012). Other OR genes include those encoding RNA-binding proteins, putative lipid transfer, *RD29A*, *RD29B*, *Kin1*, and *Kin2*. Considerable variation among factors such as plant tissue type, age, and developmental stage also affects the expression of OR genes (Zhu et al., 1997).

Transcription factors (TFs), including bZIP, WRKY, AP2, NAC, C<sub>2</sub>H<sub>2</sub> zinc finger gene, and DREB, are the most important regulators of gene expression under salinity stress (Gupta and Huang, 2014). They interact with diverse cis-elements in the promoter regions of many downstream genes and modify their expression. There are many different types of TFs in the plant kingdom (Shiu et al., 2005) with more than 50 families distinguished by their DNA-binding domains (Riechmann et al., 2000). Several TFs have been identified in Arabidopsis, including a basicleucine zipper, AP2/ERF (APETALA2/ethyleneresponsive factor), MYC, HD-ZIP (homeodomainleucine zipper), MYB, and different classes of zinc finger domains (Shinozaki and Yamaguchi-Shinozaki, 2000). The expression of a given number of genes occurs through TF interactions with non-DNA-binding proteins and/or cis-regulatory elements (Grotewold, 2008). On the same transduction pathways, ABA, SA, ethylene, BR, and JA regulate TF expression (Agarwal et al., 2006; Seo et al., 2008; Agarwal and Jha, 2010; Gürel et al., 2016). The four classes of TFs are: (1) ABA-dependent (CBF/DREB proteins), (2) ABAindependent (MYC, MYB, ABA-responsive elementbinding protein (AREB), ABA-binding factor protein (ABF)) (Agarwal and Jha, 2010; Liu SW et al., 2018), (3) ABA-dependent and -independent (no apical meristem, ATAF1,2 and cup-shaped cotyledon (NAC)) (Agarwal et al., 2013; Joshi et al., 2016), and (4) TFs that do not belong to any of the first three groups, including homeodomain TFs osmotically responsive gene 9 (HOS9) and an R2R3-type MYB protein (HOS10) (Zhu et al., 2004, 2005).

# 4 Genetics of salinity tolerance at germination in barley

# 4.1 QTLs for salinity tolerance in barley and the homologous genes from *Arabidopsis*, soybean, maize, wheat, and rice at the germination stage

At the germination stage, salinity tolerance is controlled by various loci (Mano and Takeda, 1997), indicating that it is polygenic. Angessa et al. (2017) reported transgressive phenotypic segregation for germination percentage in a doubled haploid (DH) population developed from salinity-tolerant genotype CM72 and the sensitive commercial Australian barley cultivar Gairdner. Using 150 and 300 mmol/L NaCl, Angessa et al. (2017) mapped two stable quantitative trait loci (QTLs) to chromosome 2H, close to the sodium concentration QTL reported by Xue et al. (2009). A third QTL reported by Angessa et al. (2017) at 300 mmol/L NaCl was mapped to chromosome 5H, but slightly distant from the potassium concentrationlinked QTL of Xue et al. (2009). Mano and Takeda (1997) reported QTLs controlling ABA response on chromosomes 2H, 3H, 1H, and 5H in Steptoe/Morex DH lines, and 2H and 5H in Harrington/TR306 DH lines. In both crosses, chromosome 5H was very close to the position of the salinity tolerance QTL. A QTL analysis of the Oregon Wolf Barley mapping population (DOM×REC) by Witzel et al. (2010) at the

germination stage identified two chromosomal regions on 5H, one on 7H, and one on 2H associated with the salt stress response. A single chromosomal region on 5H, responsible for 42% of the variation in the phenotype, was constant across all NaCl concentrations (Table 1).

Protein sequences of functional salinity tolerance genes during germination in Arabidopsis, soybean, maize, wheat, and rice were extracted from the National Center for Biotechnology Information (NCBI) website (https://www.ncbi.nlm.nih.gov/protein) (Benson et al., 2013). The protein sequences in FAST formats were blasted on BARLEX (https://apex.ipkgatersleben.de/apex/f?p=284:10) to identify homologous genes from the barley genome with the highest percentage match depending on length and expression levels in tissues at different growth stages (Colmsee et al., 2015). There were 63 functionally characterized genes (Arabidopsis (13), maize (4), rice (12), soybean (7), and wheat (27)) for salinity tolerance at the germination stage (Table 2). Overall, 65 homolog genes were identified in barley, with the percentage match ranging from 100% (between wheat gene TaPLDa(phospholipase D) and barley homolog gene HORVU 1Hr1G048970.4 (phospholipase D P2)) to 30% (between rice gene OsOPT10 (oligopeptide transporter family homolog) and barley's HORVU6Hr1G067430.2 (oligopeptide transporter 4)) (Table 2). Generally, the

| Trait at          | Solt level      | Manning population             | Analysis         | QTL number                    | Percentage | Deference                |
|-------------------|-----------------|--------------------------------|------------------|-------------------------------|------------|--------------------------|
| germination       | Salt level      | Mapping population             | type             | and location                  | explained  | Kelelelice               |
| Germination       | 150 mmol/L NaCl | 103-DH of CM72/Gairdner        | <b>Bi-Parent</b> | 1 (2H)                        | 21.8       | Angessa et al., 2017     |
| percentage        | 300 mmol/L NaCl | 103-DH of CM72/Gairdner        | <b>Bi-Parent</b> | 1 (2H)                        | 16.1       | Angessa et al., 2017     |
|                   | 300 mmol/L NaCl | 103-DH of CM72/Gairdner        | <b>Bi-Parent</b> | 1 (5H)                        | 10.0       | Angessa et al., 2017     |
| Germination speed | 250 mmol/L NaCl | 149-DH of Steptoe/Morex        | Bi-Parent        | 2 (2H), 3 (3H),<br>and 7 (5H) | 7.6–20.3   | Mano and Takeda,<br>1997 |
|                   | 300 mmol/L NaCl | 146-DH of Harrington/<br>TR306 | Bi-Parent        | 1 (7H), 6 (6H),<br>and 7 (5H) | 8.7–15.3   | Mano and Takeda,<br>1997 |
| Salt tolerance    | 250 mmol/L NaCl | 149-DH of Steptoe/Morex        | Bi-Parent        | 4 (4H), 6 (6H),<br>and 7 (5H) | 6.8–46.7   | Mano and Takeda,<br>1997 |
|                   | 300 mmol/L NaCl | 146-DH of Harrington/<br>TR306 | Bi-Parent        | 5 (1H) and 7 (5H)             | 17.4–41.1  | Mano and Takeda,<br>1997 |
|                   | 1.5%*           | 94-DH of DOM/REC               | Bi-Parent        | 1 (5H) and 1 (7H)             | 42         | Witzel et al., 2010      |
|                   | 2.0%            | 94-DH of DOM/REC               | Bi-Parent        | 1 (5H) and 1 (7H)             | 42         | Witzel et al., 2010      |
|                   | 2.5%            | 94-DH of DOM/REC               | <b>Bi-Parent</b> | 1 (5H) and 1 (7H)             |            | Witzel et al., 2010      |
|                   | 1.5%            | 94-DH of DOM/REC               | <b>Bi-Parent</b> | 1 (2H)                        |            | Witzel et al., 2010      |
|                   | 1.5%-2.5%       | 94-DH of DOM/REC               | Bi-Parent        | 2 (5H), 1 (7H),<br>and 1 (2H) |            | Witzel et al., 2010      |

Table 1 Reported quantitative trait loci (QTLs) for salinity tolerance in barley during germination

\* 1%=0.01 g/mL

wheat genes had the highest match with barley homologs, averaging 90.93%, followed by maize (83.00%), rice (67.58%), soybean (59.43%), and Arabidopsis (57.94%). Apart from homolog connections, gene expression patterns in tissues can suggest the function of particular genes. For instance, the expression levels of the 65 barley homologs for the first three development stages (4-d embryo (EMB), root from seedlings (ROO1; 10 cm shoot stage), and shoot from seedling stage (LEA; 10 cm shoot stage)) were more than zero for all except HORVU1Hr 1G080820.6 (CBL protein 8). The aquaporin-like superfamily protein (HORVU2Hr1G096360.13) had the highest expression levels in the EMB, ROO1, and LEA stages at 1305.90, 1094.39, and 1133.08 fragments per million kb, respectively (Table S1) (The International Barley Genome Sequencing Consortium, 2012; Colmsee et al., 2015; Mascher et al., 2017).

# 4.2 Barley salinity tolerance characterized transcriptional factors and genes during germination

A highly positive correlation has been reported between salinity stress and polyethylene glycol treatments, indicating that salt stress in germinating seed is mostly osmotic (Mano et al., 1996). Therefore, osmoprotection is the likely early response in barley geminating seeds (Walia et al., 2006).

# 4.2.1 6PGDH and Glc/RibDH

Salinity tolerant lines at germination in a mapping population containing 94 DH lines of DOM/ REC expressed a higher level of 6-phosphogluconate dehydrogenase (6PGDH) and glucose/ribitol dehydrogenase (Glc/RibDH) in mature seeds (Witzel et al., 2010). Overexpression of the two enzymes in yeast enhanced the growth of transformed cells in saline media (Witzel et al., 2010). Cytosolic 6PGDH is an enzyme participating in the pentose phosphate pathway, which provides reduced nicotinamide adenine dinucleotide phosphate (NADPH) as an electron donor in the reductive biosynthetic process, and is important for the ascorbate-glutathione synthesis phase in the plant antioxidant protection system (Corpas et al., 1998). This protein concentration is increased under salinity stress in olive (Valderrama et al., 2006) and rice (Huang et al., 2003; Hou et al., 2007). Transcripts and protein Glc/RibDH are short alcohol

dehydrogenases (Jornvall et al., 1984) found in developing barley embryos, whose level of transcription reduces with germination (Alexander et al., 1994), and which catalyse the oxidation of D-glucose to synthesize sugars for osmoprotective functions (Witzel et al., 2010).

# 4.2.2 Dehydrins

Dehydrins (DHNs), a subfamily of LEA proteins from Hordeum vulgare (aba2), enhanced seed germination in transgenic lines of Arabidopsis under salinity stress, relative to the wild type (Calestani et al., 2015). Similar protein-coding genes have been identified in wheat (Dhn5) and maize (Rab17) with comparable results when overexpressed in transgenic Arabidopsis plants (Figueras et al., 2004; Brini et al., 2011). DHNs are thought to be involved in protecting the embryo through seed desiccation and rehydration (Skriver and Mundy, 1990; Blackman et al., 1991; Tunnacliffe and Wise, 2007) by increasing their contents during the last phase of seed development, and in dormant embryos. DHN content decreases when imbibition and germination begin as seeds lose their capacity to withstand desiccation (Blackman et al., 1991; Han et al., 1997). DHNs also facilitate water uptake during seed germination on media with low osmotic potential (Hara, 2010), and may act as a hydration buffer inside cells in the presence of sugars (Walters et al., 1997; Hara, 2010). Moreover, they are likely to play a role in cellular detoxification because they contain lipids and metal-binding capacity that can prevent lipid peroxidation (Cheng et al., 2002; Krüger et al., 2002; Alsheikh et al., 2003; Koag et al., 2003, 2009).

#### 4.2.3 CBLs

Rice transgenic lines with CBL protein 8 (*HsCBL8*) transferred from wild barley homolog to HvCBL8 of cultivated barley improved seed germination, reduced Na<sup>+</sup> uptake, adjusted K<sup>+</sup> concentration in tissues, provided in vivo water protection of the plasma membrane, and accumulated more proline under salt stress (Guo et al., 2016). The build-up of compatible osmolytes, like proline, is linked to stress tolerance in plants (Ahmed et al., 2013b; Mekawy et al., 2015). Phylogenetic analysis of *HsCBL8* showed that it encodes proteins of the CBL group modified with only N-myristoylation or S-acylation (Batistič et al.,

| Ta                        | ible 2 Homold | ogs of candida     | te functional salinity gene | s in barle        | $\gamma$ during the germination of $Ara$     | <i>bidopsis</i> , maiz | æ, rice, soybean, and wh  | neat                                  |
|---------------------------|---------------|--------------------|-----------------------------|-------------------|--|------------------------|---------------------------|---------------------------------------|
| Gene source               | Gene ID       | Transgenic<br>host | Homolog ID in barley        | Similarity<br>(%) | Gene name in barley                          | Chromosome             | Location                  | Reference                             |
| Arabidopsis<br>thaliana L | AtRZFP        | Arabidopsis        | HORVU5Hr1G001400.5          | 52                | RING finger protein 5                        | 5                      | 4181876-4184642           | Zang et al., 2016                     |
| A. thaliana L.            |               |                    | HORVU4Hr1G003600.1          | 51                | RING finger protein 5                        | 4                      | 7870080 - 7872190         |                                       |
| A. thaliana L.            | AtNHXI        | Maize/wheat        | HORVU2Hr1G021020.4          | 69                | Sodium hydrogen exchanger 2                  | 7                      | 59025629-59030226         | Yin et al., 2004;<br>Xue et al., 2004 |
| A. thaliana L.            |               |                    | HORVU4Hr1G033760.1          | 69                | Sodium hydrogen exchanger 2                  | 4                      | 238472323-238482634       |                                       |
| A. thaliana L.            |               |                    | HORVU7Hr1G046030.1          | 62                | Sodium hydrogen exchanger 3                  | 7                      | $148709554 {-} 148713307$ |                                       |
| A. thaliana L.            | AtSOSI        | Arabidopsis        | HORVU3Hr1G003150.1          | 58                | Sodium hydrogen exchanger 7                  | ŝ                      | 8165119-8185196           | Wang X et al., 2007                   |
| A. thaliana L.            | AtMYBL        | Arabidopsis        | HORVU5Hr1G077100.1          | 56                | MYB domain protein 3R-3                      | 5                      | 552718664-552726337       | Zhang et al., 2011                    |
| A. thaliana L.            | RAP2.6L       | Arabidopsis        | HORVU2Hr1G071270.4          | 60                | Ethylene-responsive                          | 7                      | 507785971-507787466       | Krishnaswamy                          |
|                           |               |                    |                             |                   | transcription factor 1                       |                        |                           | et al., 2011                          |
| A. thaliana L.            | AtERF96       | Arabidopsis        | HORVU4Hr1G000960.1          | 59                | Ethylene-responsive                          | 4                      | 1356682-1357123           | Wang et al., 2017                     |
|                           |               |                    |                             |                   | transcription factor 14                      |                        |                           |                                       |
| A. thaliana L.            | AtCYSa        | Arabidopsis        | HORVU1Hr1G067870.2          | 56                | Cysteine proteinase inhibitor                | 1                      | 479563716-479564368       | Zhang et al., 2008                    |
| A. thaliana L.            | AtCYSb        | Arabidopsis        | HORVU3Hr1G038190.1          | 99                | Cysteine proteinase inhibitor 12             | ю                      | 219953557-219956556       | Zhang et al., 2008                    |
| A. thaliana L.            | AtNHXI        | Cotton             | HORVU2Hr1G021020.4          | 69                | Sodium hydrogen exchanger 2                  | 7                      | 59025629-59030226         | Cheng C et al., 2018                  |
| A. thaliana L.            | AtPP2-B11     | Arabidopsis        | HORVU6Hr1G090000.1          | 42                | F-box protein PP2-B1                         | 9                      | 571135647-571137859       | Jia et al., 2015                      |
| A. thaliana L.            | AnnAt8        | Arabidopsis        | HORVU6Hr1G074440.1          | 50                | Annexin 7                                    | 9                      | 514097241-514103959       | Yadav et al., 2016                    |
| A. thaliana L.            | AtDIF1        | Arabidopsis        | HORVU5Hr1G079500.1          | 42                | Sister chromatid cohesion 1                  | 1                      | 557145143-557147352       | Gao et al., 2017                      |
|                           |               |                    |                             |                   | protein 4                                    |                        |                           |                                       |
| A. thaliana L.            | NF-YA1-9      | Arabidopsis        | HORVU5Hr1G007890.11         | 99                | Nuclear transcription factor Y               | 5                      | 16694492-16699091         | Mu et al., 2013                       |
| Maize (Zad mane)          | 7mCBL A       | Arahidansis        | HOPVIIIHr160808206          | <u>7</u> 7        | subunit A-10<br>Calcinaurin B-1iba nrotain 8 |                        | 576318770-576370885       | Wang MV at al                         |
| INTAIZE (ZEU MUYS)        |               | Aruvuupsis         |                             | 7                 |  | -                      | C00075075-01/015075       | walig M I Clal.,<br>2007              |
| Maize (Z. mays)           | ZmMKK4        | Arabidopsis        | HORVU0Hr1G038850.2          | 88                | Protein kinase superfamily                   | 0                      | 247582088–247583184       | Kong et al., 2011                     |
|                           |               |                    |                             |                   | protein                                      |                        |                           |                                       |
| Maize (Z. <i>mays</i> )   | ZmMPK5        | Tobacco            | HORVU7Hr1G023760.3          | 89                | Mitogen-activated protein<br>kinase 3        | L                      | 37300015-37306217         | Zhang et al., 2014                    |
| Rice (Oryza sativa        | ) OSISAP I    | Tobacco            | HORVU5Hr1G072920.1          | 70                | Zinc finger A20 and AN1                      | 5                      | 536933399-536936238       | Mukhopadhyay                          |
|                           |               |                    |                             |                   | domain-containing                            |                        |                           | et al., 2004                          |
|                           |               |                    |                             |                   | stress-associated protein 1                  |                        |                           |                                       |
| Rice (O. sativa)          | OsDREB2A      | Soybean            | HORVU6Hr1G050520.2          | 78                | Ethylene-responsive                          | 9                      | 306973236-306973949       | Zhang et al., 2013                    |
| () (d                     | 0475:0        | T-1-7              |                             | ç                 | transcription factor 4                       | ć                      | 0021120CC 170CL           | r ,7                                  |
| Kice (U. sativa)          | USiJAF8       | 1 obacco/rice      | HUKV U2HFIGU336/0.4         | 45                | Zinc linger AZU and ANI                      | 7                      | 328/13904-328/14/08       | Kanneganti and                        |
|                           |               |                    |                             |                   | domain-containing                            |                        |                           | Gupta, 2008                           |
|                           |               |                    |                             |                   | stress-associated protein 9                  |                        |                           |                                       |
|                           |               |                    |                             |                   |  |                        |                           | To be continued                       |

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| Gene source        | Gene ID   | Transgenic<br>host | Homolog ID in barley | Similarit<br>(%) | y Gene name in barley                          | Chromosome | Location Referen                        | nce       |
|--------------------|-----------|--------------------|----------------------|------------------|--|------------|---|-----------|
| Rice (O. sativa)   | OsMYB48-1 | Rice               | HORVU1Hr1G008810.3   | 82               | MYB domain protein 59                          | -          | 19369578-19370912 Xiong et al.,         | , 2014    |
| Rice (O. sativa)   | OsSAP11   | Arabidopsis        | HORVU7Hr1G050270.3   | 99               | Zinc finger A20 and AN1                        | 7          | 180473211-180476075 Giri et al., 20     | 011       |
|                    |           |                    |                      |                  | domain-containing                              |            |   |           |
|                    |           |                    |                      |                  | stress-associated protein 11                   |            |   |           |
| Rice (O. sativa)   | OsRLCK253 | Arabidopsis        | HORVU6Hr1G061280.1   | 69               | Protein kinase superfamily                     | 9          | 410507581-410509354 Giri et al., 20     | 011       |
|                    |           |                    |                      |                  | protein  |            |   |           |
| Rice (O. sativa)   | OsGSTU4   | Arabidopsis        | HORVU6Hr1G026810.3   | 71               | Glutathione S-transferase                      | 9          | 103574244-103575543 Sharma et al.       | l., 2014  |
|                    |           |                    |                      |                  | family protein                                 |            |   |           |
| Rice (O. sativa)   | OsDREBID  | Arabidopsis        | HORVU5Hr1G080420.1   | 71               | C-repeat-binding factor 4                      | 5          | 560569797-560571193 Zhang et al.,       | ,2009     |
| Rice (O. sativa)   | OsNAC9    | Rice               | HORVU7Hr1G060310.1   | 75               | NAC domain protein                             | 7          | 276430981-276436993 Redillas et al      | al., 2012 |
| Rice (O. sativa)   | OsZFP179  | Rice               | HORVU5Hr1G055970.1   | 85               | RING/U-box superfamily                         | 5          | 438292337-438301123 Sun et al., 20      | 010       |
|                    |           |                    |                      |                  | protein  |            |   |           |
| Rice (O. sativa)   | OsOPT10   | Arabidopsis/       | HORVU6Hr1G067430.2   | 30               | Oligopeptide transporter 4                     | 9          | 467362064-467373338 Jung et al., 20     | 2010      |
|                    |           | rice               |                      | ì                |  |            |   | 1         |
| Rice (O. sativa)   | OsMYB3R-2 | Arabidopsis        | HORVU3Hr1G086270.3   | 71               | MYB domain protein 3R-5                        | m          | 616465321–616471572 Dai et al., 20      | 007       |
| Soybean (Glycine   | GmbZIP44  | Arabidopsis        | HORVU5Hr1G023000.1   | 43               | Basic-leucine zipper transcription             | 5          | 119817528-119818004 Liao et al., 20     | 2008      |
| max)               |           |                    |                      |                  | factor family protein                          |            |   |           |
| Soybean (G. max)   | GmbZIP62  | Arabidopsis        | HORVU6Hr1G031330.2   | 49               | Basic-leucine zipper transcription             | 9          | 132098841-132101167 Liao et al., 20     | 2008      |
|                    |           |                    |                      |                  | factor family protein                          |            |   |           |
| Soybean (G max)    | GmbZIP78  | Arabidopsis        | HORVU2Hr1G055230.4   | 61               | Basic-leucine zipper 8                         | 2          | 342114231–342120751 Liao et al., 20     | 2008      |
| Soybean (G max)    | GmDREB2   | Arabidopsis/       | HORVU4Hr1G015350.1   | 99               | Ethylene-responsive transcription              | 4          | 59331360–59332535 Chen M et al.         | ıl., 2007 |
|                    |           | tobacco            |                      |                  | factor 2                                       |            |   |           |
| Soybean (G. max)   | GmDREB1   | Wheat              | HORVU7Hr1G026940.1   | 67               | Ethylene-responsive transcription              | 2          | 47357790-47359063 Jiang et al., 2       | 2014      |
|                    |           |                    |                      |                  | factor 5                                       |            |   |           |
| Soybean (G. max)   | GmNARK    | Arabidopsis        | HORVU5Hr1G098840.2   | 54               | Leucine-rich receptor-like                     | S          | 608904210-608907733 Cheng CH et         | it al.,   |
|                    |           |                    |                      |                  | protein kinase family protein                  |            | 2018                                    |           |
| Soybean (G. max)   | GmFDL19   | Soybean            | HORVU2Hr1G110500.1   | 76               | ABSCISIC ACID-                                 | 2          | 720057018–720058296 Li et al., 2017     | []        |
|                    |           |                    |                      |                  | INSENSITIVE 5-like protein 5                   | 10         |   |           |
| Wheat (Triticum    | TaNAC29   | Arabidopsis        | HORVU2Hr1G017470.1   | 93               | NAC domain protein                             | 2          | 42672748-42675686 Huang et al.,         | , 2015    |
| aestivum)          |           |                    |                      |                  |  |            |   |           |
| Wheat (T. aestivum | ) TaSC    | Arabidopsis        | HORVU5Hr1G067740.1   | 66               | Transmembrane protein 50A                      | 5          | 514305842-514309764 Huang et al.,       | , 2012    |
| Wheat (T. aestivum | ) TaNAC2D | Arabidopsis        | HORVU5Hr1G111590.2   | 95               | NAC domain protein                             | 5          | 636772274–636774317 Huang and W<br>2016 | Wang,     |
| Wheat (T. aestivum | ) TaARGOS | Arabidopsis        | HORVU4Hr1G043960.1   | 88               | Eukaryotic aspartyl protease<br>family protein | 4          | 351442597–351444197 Zhao et al., 2      | 2017      |

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To be continued

Table 2

| Location Reference      | 30039636-630043684 Xiang et al., 2015 | 19528221–519529717 He et al., 2011 |          | 31361349–531362720 Kavas et al., 2016 | 16115076-616116086 Brini et al., 2007 | 74156225-674158128 Hu et al., 2012 | 16404110-716405772 Wang et al 2013   |        | 69768135-469771916 Li et al., 2010 |                      | 88366914–88373289 Kong et al., 2015 |                     | 23061366–623067662 Xu et al., 2008a |                          | 46669169–46672779 Deng et al., 2013 | 82511166–682511937 Rong et al., 2014 |                        | 27004064–627010259 Zhang et al., 2017 | 90705028–390708928 Wang et al., 2015 | 38472323–238482634 Wang et al., 2011 | 89779039–89780525 Hou et al., 2013 | 59025629-59030226 Gouiaa et al., 2012 | 63936492-63936858 Gouiaa et al., 2012      |                       | 39161697–639164555 Gouiaa et al., 2012 |                           | 63556495–363563036 Wang et al., 2014 | 4131150-4132543 Dong et al., 2013 | 44207949–544211747 Ma et al., 2015 |             | 42201795–742204374 Liu et al., 2013 |                         |   |
|-------------------------|---------------------------------------|------------------------------------|----------|---------------------------------------|---------------------------------------|------------------------------------|--------------------------------------|--------|------------------------------------|----------------------|-------------------------------------|---------------------|-------------------------------------|--------------------------|-------------------------------------|--------------------------------------|------------------------|---------------------------------------|--------------------------------------|--------------------------------------|------------------------------------|---------------------------------------|--|-----------------------|--|---------------------------|--------------------------------------|-----------------------------------|------------------------------------|-------------|-------------------------------------|-------------------------|---|
| omosome.                | 3 6                                   | 2                                  |          | 4 5                                   | 5 6                                   | 2 6                                | C                                    | I      | 3 4                                |                      | 0                                   |                     | 5 6                                 |                          | 7                                   | 2 6                                  |                        | 3 6                                   | 4                                    | 4                                    | 7                                  | 2                                     | 7  |                       | 2 6                                    |                           | 1 3                                  | 1                                 | 6 5                                |             | 2 7                                 |                         | : |
| Gene name in barley Chr | Calreticulin 3                        | Family of unknown function         | (DUF662) | Salt-tolerance-related protein        | Dehydrin Rab15                        | Aquaporin-like superfamily         | protein<br>WRKY family transcrintion | factor | Protein DEHYDRATION-               | INDUCED 19 homolog 3 | Transducin/WD40 repeat-like         | superfamily protein | Histone-lysine                      | N-methyltransferase NSD3 | Protein kinase family protein       | Ethylene-responsive                  | transcription factor 1 | Pre-mRNA-processing factor 19         | WRKY DNA-binding protein 3           | Sodium hydrogen exchanger 2          | Protein of unknown function        | Sodium hydrogen exchanger 2           | K <sup>+</sup> -insensitive pyrophosphate- | energized proton pump | Pyrophosphate-energized vacu-          | olar membrane proton pump | Phospholipase D P2                   | 12-Oxophytodienoate reductase 2   | Nuclear transcription factor Y     | subunit A-5 | Phosphatidylinositol 3- and         | 4-kinase family protein |   |
| Similarity<br>(%)       | 98                                    | 97 I                               |          | 60                                    | 79 I                                  | 61                                 | 87                                   | I      | 97 ]                               |                      | 66                                  |                     | 72 1                                |                          | 92 ]                                | 85 1                                 |                        | 95 ]                                  | 92                                   | 67                                   | 90                                 | 70                                    | 97 I                                       |                       | 88                                     |                           | 100                                  | 82                                | 89 ]                               |             | 94 I                                |                         | 0 |
| Homolog ID in barley    | HORVU3Hr1G089830.2                    | HORVU2Hr1G072420.11                |          | HORVU4Hr1G063430.1                    | HORVU5Hr1G103460.2                    | HORVU2Hr1G096360.13                | HORVI DHr1G109330 1                  |        | HORVU3Hr1G061690.1                 |                      | HORVU0Hr1G016450.1                  |                     | HORVU5Hr1G105980.4                  |                          | HORVU2Hr1G018340.3                  | HORVU2Hr1G098330.1                   |                        | HORVU4Hr1G083960.3                    | HORVU4Hr1G048400.12                  | HORVU4Hr1G033760.1                   | HORVU7Hr1G037410.4                 | HORVU2Hr1G021020.4                    | HORVU7Hr1G031480.1                         |                       | HORVU7Hr1G114250.2                     |                           | HORVU1Hr1G048970.4                   | HORVU1Hr1G001850.3                | HORVU6Hr1G081080.12                |             | HORVU2Hr1G118320.7                  |                         |   |
| Transgenic<br>host      | Tobacco                               | Arabidopsis/                       | rice     | Tobacco                               | Arabidopsis                           | Tobacco                            | Tohacco                              |        | Arabidopsis                        |                      | Arabidopsis                         |                     | Arabidopsis                         |                          | Tobacco                             | Wheat                                |                        | Tobacco                               | Tobacco                              | Arabidopsis                          | Arabidopsis                        | Tobacco                               | Tobacco                                    |                       |  |                           | Arabidopsis                          | Arabidopsis                       | Arabidopsis                        |             | Arabidopsis                         |                         | - |
| Gene ID                 | TaCRTI                                | TaSRG                              |          | TaSTRG                                | DHN-5                                 | TaAQP8                             | TaWRKY10                             |        | TaDi19A                            |                      | TaWD40D                             |                     | TaAIDFa                             |                          | TaCIPK29                            | TaERF3                               |                        | TaPUBI                                | TaWRKY44                             | TaVB                                 | TaSRHP                             | TaNHXSI                               | TaVPI                                      |                       |  |                           | $TaPLD\alpha$                        | TaOPR1                            | TaNF-YA10-1                        |             | TaP14KII <sub>Y</sub>               |                         |   |
| Gene source             | Wheat (T. aestivum)                   | Wheat (T. aestivum)                |          | Wheat (T. aestivum)                   | Wheat (T. aestivum) 1                 | Wheat (T. aestivum)                | Wheat (T aestivum)                   |        | Wheat (T. aestivum)                |                      | Wheat (T. aestivum)                 |                     | Wheat (T. aestivum)                 |                          | Wheat (T. aestivum)                 | Wheat (T. aestivum)                  |                        | Wheat (T. aestivum)                   | Wheat (T. aestivum)                  | Wheat (T. aestivum)                  | Wheat (T. aestivum)                | Wheat (T. aestivum)                   | Wheat (T. aestivum)                        |                       | Wheat (T. aestivum)                    |                           | Wheat (T. aestivum)                  | Wheat (T. aestivum)               | Wheat (T. aestivum)                |             | Wheat (T. aestivum)                 |                         |   |

RING, really interesting new gene; MYB, myeloblastosis, NAC, no apical meristem, ATAF1,2 and cup-shaped cotyledon; NSD, nuclear receptor-binding SET domain

.

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Table 2

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2012). Such proteins are involved in relationships with the transmembrane (TM) helix (Kleist et al., 2014) including *HvCBL8*. Reports about other CBLs from different plant species reflected similar trends to that of *HsCBL8*, such as transgenic *Arabidopsis* having 35S-AtCBL5 (Cheong et al., 2010) and poplar harbouring 35S-PeCBL10 (Li DD et al., 2013). OsCBL2 is likely to be involved in GA-signalling that facilitates the vacuolation of aleurone cells (Hwang et al., 2005), and AtCBL1 is involved in upregulation of GA (Li ZY et al., 2013) and downregulation.

# 4.2.4 SERKs

Three orthologs of somatic embryogenesis receptor-like kinase (SERK) genes (HvSERK1/2/3) isolated from barley were induced in microsporederived embryogenic callus under salt stress (Li et al., 2016), indicating their protective role for developing embryos during salinity stress. SERKs are a subfamily of the leucine-rich repeat receptor-like kinase II group (LRR-RLKII) with proline-rich SPP motifs between the LRRs and the TM domain (Hecht et al., 2001). These genes are present during somatic embryogenesis in plants before declining in later stages (Schmidt et al., 1997; Somleva et al., 2000; Nolan et al., 2003; Singla et al., 2008; Li, 2010). They have been characterized in many plants, including maize (Baudino et al., 2001), wheat (Singla et al., 2008), soybean (Yang et al., 2011), rice (Singla et al., 2009), cotton (Pandey and Chaudhary, 2014), and Arabidopsis (Hecht et al., 2001). In rice (Oryza sativa L.), BRs reverse the inhibitory effect of salinity during germination (Anuradha and Rao, 2001), and in Arabidopsis are reportedly signalled by SERK1, SERK3, and SERK4 (Albrecht et al., 2008), while ABA stimulates OsSERK1 during stress (Hu et al., 2005).

# 4.2.5 DREBs

A gene that encodes DREB proteins in *H. vul*gare (*HvDREB1*) is induced by exogenous ABA to enhance germination and early root growth in *Ara*bidopsis plants under salinity stress (Xue et al., 2004). TF *HvDREB1* is a member of the AP2 group of the *DREB* subfamily that is vital for regulating responses to various stresses (Agarwal et al., 2006; Xu et al., 2008b) and reportedly improves salt, drought, and cold responses in transgenic plants (Oh et al., 2007). Overexpressing rice OsDREB1A in Arabidopsis enhanced tolerance to drought and freezing stresses (Dubouzet et al., 2003), and barley's HvCBF4 in rice transgenes increased drought, salt, and cold stress tolerance (Oh et al., 2007). DREB proteins isolated from several plants including rice, maize, soybean, and wheat are involved in several signal transduction pathways during abiotic stress (Dubouzet et al., 2003; Agarwal et al., 2006). Of particular interest are those belonging to the A-2 group that respond to dehydration and salinity stress (Liu et al., 1998; Dubouzet et al., 2003) and regulate ABA-independent gene expression in target plants under stress (Chinnusamy et al., 2006; Nakashima and Yamaguchi-Shinozaki, 2006). It is possible that *HvDREB1* may take part in both ABA-independent and -dependent pathways concurrently (Xue et al., 2004).

# 4.2.6 ERFs

Barley ERF-type TF HvRAF improved seed germination and root growth under salinity stress but was not induced by ABA treatment in transgenic Arabidopsis plants (Jung et al., 2007). TF HvRAF is a member of the AP2 group in the ERF family (Sakuma et al., 2002; Gutterson and Reuber, 2004), which is involved in regulating both biotic and abiotic stress-responsive genes in plants (Lee et al., 2004; Yi et al., 2004). However, the response of Arabidopsis transgenic lines with HvRAF was specific to salinity stress, where it acted as a regulator for ABAindependent signalling in root growth and seed germination (Jung et al., 2007). Similar ABA regulation responses to salinity have been documented in Arabidopsis for DREB2A and DREB2B (Chinnusamy et al., 2005). A tolerance response in terms of root growth and seed germination was observed when CaERFLP1 and JERF1 from pepper and tomato, respectively, were overexpressed in transgenic tobacco plants in response to salt stress (Lee et al., 2004; Zhang et al., 2004).

# 4.2.7 Aquaporin genes

Overexpression of a barley aquaporin gene *HvPIP2;5* in yeast enhanced salt and osmotic stress tolerance, and transgenic *Arabidopsis* with the gene showed better seed germination and root growth than the wild type under salinity stress (Alavilli et al., 2016). Aquaporins form part of the major intrinsic

proteins (MIPs) found in living organisms that enable the transportation of water and small nonpolar molecules across living membranes (Zardoya et al., 2002; Maurel et al., 2015). In plants, all but the GlpF-like intrinsic proteins show water-specific channel movement. Hence, the collective name for MIPs in plants is aquaporins (Maurel et al., 2008), even though they can transport additional small molecules such as CO2 and ammonia inside cells (Uehlein et al., 2003; Jahn et al., 2004). There are five groups of aquaporins in plant genomes, reflective of their diverse physiological roles. Of importance are the plasma membrane intrinsic proteins (PIPs) that are further subdivided into PIP1 and PIP2, each with several isoforms (Javot et al., 2003; Postaire et al., 2010). Overexpression of PIP genes from various plants (O. sativa, Nicotiana tabacum, Vicia faba, Arabidopsis, and Triticum aestivum) enhanced dehydration stress tolerance in their respective transgenic lines (Alavilli et al., 2016).

# 4.3 Diversity of barley salinity tolerance genes at germination

In this review, we have compared studies to determine whether any of the eight genes (Table 3) that have been reported and functionally characterized for salinity tolerance in barley at germination are unique or similar to their homologs from wheat and rice. All the sequences producing a significant match, based on length and total relationship hits, were downloaded and recorded. The following total significant homolog hits from barley, wheat, and rice were recorded: dehydrins (17), dehydrogenase/reductase SDR family member 4 (20), 6PGDH, (29), ERFs (31), CBL protein (72), DREB protein (164), SERKs (215), and aquaporin gene (227).

Three genes families—DREB protein, SERKs and aquaporin gene—were considered for further analysis based on their high numbers of hits. The proteins from the three genes were blasted on the Phytozome (https://phytozome.jgi.doe.gov/pz/portal. html) to download matched homologs (>30%) in 40 different plant species. Mega software (https://www. megasoftware.net) was used to align the 40 species' sequences, and a phylogenetic tree was constructed to estimate the evolutionary distances between the genes using MEGA-X software (https://www.megasoftware. net) (Kumar et al., 2018). Gene alignments showed three distinct regions across the sequences of the gene families. The start and end sections of the sequences were more divergent than the middle segments. The phylogenetic tree of the DREB proteins divided the genes into two major groups that were further divided into two subgroups (Fig. 2a). The groups were distinguished by the presence or absence of genes from wheat, barley, and rice. Among the four sub-groups, barley genes appeared three times in two categories while genes from the three cereals were missing in two sub-groups (Fig. 2a). The phylogenetic tree shows that DREB proteins are no more diverse in barley than in wheat or rice. The SERK phylogenetic tree (Fig. 2b) was similar to that of the DREB proteins, but with more hits. However, the four subgroups contained genes from the three cereals with five hits for barley and wheat, and three hits for rice. The aquaporin gene phylogenetic tree differed from the other two trees in which the genes were first divided into three major groups, two small and one major, containing representative genes from the three cereals (Fig. 3).

#### 5 Conclusions and future prospects

Among the most destructive abiotic stresses, salinity causes massive yield losses in crops in arid, semi-arid, coastal regions, and humid and sub-humid landscapes. Thus, more effort is needed to increase crop yields in these areas to produce enough food for the increasing global population. Growing salt-adapted cultivars requires knowledge of the donating characters at different growth stages. It involves the use of many disciplines to identify and functionally characterize the genes contributing to tolerance, and then to transfer them to commercially acceptable cultivars. Barley is one of the hardiest crops that can grow in saline environments, but its germination is severely affected by salinity stress. Development of cultivars that can acclimatise to salinity at this stage is essential in regions like Western Australia that experience hot and dry summers and increasing salt levels in the topsoil before sowing in autumn. The identification and characterization of salinity tolerance genes, enzymes, and compounds during germination in barley have been ongoing, and some transgenic "salt-tolerant" plants have been developed. However, the seeds of these genotypes have had little success in



Fig. 2 Phylogenetic analyses of barley homolog genes

(a) Dehydration-responsive element binding proteins. (b) Somatic embryogenesis receptor-like kinases. The unrooted phylogenetic trees of 37 (a) and 92 (b) domains comprising four (a) and five (b) domains, respectively, from barley were constructed using MEGA-X

| Reported functional barley gene  | Gene ID in bar   | ley  | Gene name in  | barley  |
|--|--|--|---|---|
| 6-Phosphogluconate dehydrogenase   | HORVU7Hr1G00   | 6160.4 6-Pho                                       | sphogluconate dehydrogen  | ase, decarboxylating 1  |
| Glucose/ribitol dehydrogenase  | HORVU1Hr1G01   | 8140.5 Dehyd                                       | lrogenase/reductase SDR f   | amily member 4  |
| Dehydrin   | HORVU7Hr1G09   | 9800.5 Dihyd                                       | roneopterin aldolase  |   |
| Calcineurin B-like protein 8   | HORVU1Hr1G064  | 4470.4 Calcin                                      | eurin B-like protein 1  |   |
| Somatic embryogenesis receptor-like kinase   | HORVU7Hr1G06   | 8990.2 Recep                                       | tor-like protein kinase 4   |   |
| Dehydration-responsive element binding protein   | HORVU3Hr1G01   | 7950.4 Chron                                       | nodomain-helicase-DNA-b   | inding protein 5  |
| Ethylene response factor   | HORVU4Hr1G07   | 7310.1 Ethyle                                      | ene-responsive transcription  | n factor 1  |
| Aquaporin gene   | HORVU2Hr1G08   | 9940.1 Aquap                                       | oorin-like superfamily prote  | ein   |
|  |  |  |   |   |
| Reported functional barley gene  | Reported gene ID   | Chromosome   | Location  | Reference   |
| Reported functional barley gene   6-Phosphogluconate dehydrogenase   | Reported gene ID<br>6PGDH  | Chromosome<br>7                                    | Location<br>8000958-8002650   | Reference<br>Witzel et al., 2010  |
| Reported functional barley gene6-Phosphogluconate dehydrogenaseGlucose/ribitol dehydrogenase   | Reported gene ID<br>6PGDH<br>Glc/RibDH   | Chromosome<br>7<br>1                               | Location<br>8000958-8002650<br>65592292-65593858  | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010   |
| Reported functional barley gene<br>6-Phosphogluconate dehydrogenase<br>Glucose/ribitol dehydrogenase<br>Dehydrin   | Reported gene ID<br>6PGDH<br>Glc/RibDH<br>aba2                                     | Chromosome<br>7<br>1<br>7                          | Location<br>8000958-8002650<br>65592292-65593858<br>602554874-602555971   | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010<br>Calestani et al., 2015   |
| Reported functional barley gene<br>6-Phosphogluconate dehydrogenase<br>Glucose/ribitol dehydrogenase<br>Dehydrin<br>Calcineurin B-like protein 8   | Reported gene ID<br>6PGDH<br>Glc/RibDH<br>aba2<br>HvCBL8                           | Chromosome<br>7<br>1<br>7<br>1                     | Location<br>8000958-8002650<br>65592292-65593858<br>602554874-602555971<br>461521906-461524442  | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010<br>Calestani et al., 2015<br>Guo et al., 2016   |
| Reported functional barley gene<br>6-Phosphogluconate dehydrogenase<br>Glucose/ribitol dehydrogenase<br>Dehydrin<br>Calcineurin B-like protein 8<br>Somatic embryogenesis receptor-like<br>kinase  | Reported gene ID<br>6PGDH<br>Glc/RibDH<br>aba2<br>HvCBL8<br>HvSERK1/2/3            | Chromosome<br>7<br>1<br>7<br>1<br>7<br>1<br>7      | Location<br>8000958-8002650<br>65592292-65593858<br>602554874-602555971<br>461521906-461524442<br>366099333-366114129   | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010<br>Calestani et al., 2015<br>Guo et al., 2016<br>Li et al., 2016  |
| Reported functional barley gene<br>6-Phosphogluconate dehydrogenase<br>Glucose/ribitol dehydrogenase<br>Dehydrin<br>Calcineurin B-like protein 8<br>Somatic embryogenesis receptor-like<br>kinase<br>Dehydration-responsive element binding<br>protein                             | Reported gene ID<br>6PGDH<br>Glc/RibDH<br>aba2<br>HvCBL8<br>HvSERK1/2/3<br>HvDREB1 | Chromosome<br>7<br>1<br>7<br>1<br>7<br>1<br>7<br>3 | Location<br>8000958-8002650<br>65592292-65593858<br>602554874-602555971<br>461521906-461524442<br>366099333-366114129<br>46482481-46494788                        | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010<br>Calestani et al., 2015<br>Guo et al., 2016<br>Li et al., 2016<br>Xue et al., 2004                      |
| Reported functional barley gene<br>6-Phosphogluconate dehydrogenase<br>Glucose/ribitol dehydrogenase<br>Dehydrin<br>Calcineurin B-like protein 8<br>Somatic embryogenesis receptor-like<br>kinase<br>Dehydration-responsive element binding<br>protein<br>Ethylene response factor | Reported gene ID<br>6PGDH<br>aba2<br>HvCBL8<br>HvSERK1/2/3<br>HvDREB1<br>HvRAF     | Chromosome<br>7<br>1<br>7<br>1<br>7<br>3<br>3<br>4 | Location<br>8000958-8002650<br>65592292-65593858<br>602554874-602555971<br>461521906-461524442<br>366099333-366114129<br>46482481-46494788<br>603804858-603809470 | Reference<br>Witzel et al., 2010<br>Witzel et al., 2010<br>Calestani et al., 2015<br>Guo et al., 2016<br>Li et al., 2016<br>Xue et al., 2004<br>Jung et al., 2007 |

Table 3 Barley salinity tolerance functional transcriptional factors and genes at germination

The protein sequences of identified functional barley genes retrieved from BARLEX (https://apex.ipk-gatersleben.de/apex/f?p=284:10) and blasted on barley (https://webblast.ipk-gatersleben.de/barley\_ibsc/viroblast.php), wheat (https://urgi.versailles.inra.fr/blast\_iwgsc/?dbgroup= wheat\_iwgsc\_refseq\_v1\_chromosomes&program=blastn), and rice (https://www.plantgdb.org/OsGDB) genome explorers



Fig. 3 Phylogenetic analysis of barley homolog aquaporin genes The unrooted phylogenetic tree of 74 domains comprising five domains from barley was constructed using MEGA-X

commercial production because they are not equipped with holistic genes or the mechanisms required for successful germination and growth under saline field conditions.

This review summarized the agronomic/ morphological, physiological, and biochemical traits related to salinity stress. The best trait or combination of traits needs to be identified which can be used to accurately screen for salinity stress tolerance at germination to identify and characterize novel genes. Homolog salinity tolerance genes in barley during germination have been reported, but they need to be validated in barley and other transgenic plants that carry them. Functional characterization by blending genetic, agronomic, biochemical, and physiological indicators can facilitate proof of identity of the genes, leading to the development of barley cultivars with improved salt tolerance at germination and better performance in the field.

#### Contributors

Edward MWANDO performed literature search, data analysis, interpretation of information, and drafting the manuscript. Tefera Tolera ANGESSA and Yong HAN gave guidance on relevant literature search, information and data interpretation. Chengdao LI conceived the projects idea. All authors revised the paper and approved the final version to be published.

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#### **Compliance with ethics guidelines**

Edward MWANDO, Tefera Tolera ANGESSA, Yong HAN, and Chengdao LI declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by any of the authors.

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List of electronic supplementary materials

Table S1 Expression levels in different tissues and growth stages of candidate genes for barley salinity tolerance at germination

# <u>中文概要</u>

#### 题 目: 大麦芽期耐盐相关的同源和候选基因

- 概 要: 土壤盐害影响了全球 6%以上的陆地面积, 并导 致了大量的农作物减产。盐害主要通过渗透和离 子胁迫抑制植物的生长和发育, 而植物相应地通 过渗透调节、转移或外排积累的钠和氯离子以增 强适应性。目前,生产上尚未有实用、经济的方 法治理盐害,因而最为可行的途径是增强植物自 身的耐盐性。盐胁迫严重抑制种子萌发,而作为 全球第四大禾谷类作物的大麦与其他谷物相比 耐盐性更强。本文综述了大麦芽期耐盐性的遗传 机制,总结了已报道的相关数量性状位点和功能 基因,比对了拟南芥、大豆、玉米、小麦和水稻 中耐盐候选基因在大麦中的同源基因并映射到 参考基因组。此外,本文还讨论了三个耐盐功能 基因家族的遗传多样性,包括脱水应答元件结合 蛋白(DREB)、类体细胞胚胎发生受体激酶和 水通道蛋白。上述三个基因家族在植物中都存在 丰富的多样性,但 DREB 家族在大麦中的多样性 高于水稻和小麦。后续研究中, 芽期耐盐性的简 便筛选方法仍有待开发, 耐盐基因及相关机理机 制仍需鉴定、验证,并整合到栽培品种中,以实 现盐土上作物的生产。
- 关键词:遗传;大麦;数量性状位点;发芽;耐盐性;同源基因;多样性