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Biotechnology for mechanisms that counteract salt stress in extremophile species: A genome-based view

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Biotechnology for mechanisms that counteract salt stress in extremophile species: a genome-based view

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Abstract Molecular genetics has confirmed older research and generated new insights into the ways how plants deal with adverse conditions. This body of research is now being used to interpret stress behavior of plants in new ways, and to add results from most recent genomics-based studies. The new knowledge now includes genome sequences of species that show extreme abiotic stress tolerances, which enables new strategies for applications through either molecular breeding or transgenic engineering. We will highlight some physiological features of the extremophile lifestyle, outline emerging features about halophytism based on genomics, and discuss conclusions about underlying mechanisms.

Keywords *Thellungiella* · Extremophile species · Genome sequences · Abiotic stress protection · Biotechnology potential

Introduction

Over the last two decades, the search for superior alleles of key genes for plant stress tolerance has often relied on *Arabidopsis thaliana*. This plant provided a model that has significantly contributed to advance our mechanistic understanding of plant responses to environmental stresses, although the species itself is not stress tolerant by any imagination (Sanders 2000; Chen et al. 2004). Even so, the rapid development of powerful tools in molecular biology has opened up unprecedented opportunities to explore the genomes of halophytic species, which could provide better model

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systems in plant salt tolerance research, the stress on which we focus as an example (Inan et al. 2004; Orsini et al. 2010).

There is no straightforward definition of what makes a halophyte. Halophytic lifestyles—in oceans, tidal marshes, salt pans, or in irrigated deteriorated lands—take many shapes. We will focus here on salinity as a causative agent, restricting ourselves to the presence of NaCl, although salinity stress is much more complex in nature (Flowers 2004; Maggio et al. 2006), and similar arguments apply to other types of abiotic stresses.

Most broadly, halophytes on land may be defined as plants that can complete their life cycle in the presence of a salt concentration in the root zone that would kill at least 95 % of higher plant species (Flowers and Colmer 2008). Despite this general and apparently comprehensive definition, an unequivocal classification of halophytes based on both quantifiable growth performance parameters and functional halophytic traits is not available. A partial inventory, compiled by Aronson (1989), classified 1,560 species and 550 genera in 117 families as halophytes. This list refers only to plants with a potential as food, forage, fuel-wood or soil stabilization, and, according to Le Houerou (1993), includes probably only 20–30 % of the terrestrial halophytic flora. Interestingly, approximately 60 % of these species are found in only 13 families. Among those, the Chenopodiaceae contain the largest number of halophytic species (more than 270) followed by Poaceae, Fabaceae, and Asteraceae. All together, these comprise fewer than 5 % of all angiosperm species (Glenn et al. 1999).

Here, we will discuss physiological parameters that are associated with salinity stress tolerance parameters, with particular emphasis on the developmental windows and genetic determinants that appear to mark thresholds in adaptations. In many cases, the salt stress phenotype can be associated with genes that have been revealed by mutagenesis in *Arabidopsis* and increasingly also from rice (Zhu 2000; Oo and Lang 2005). These two species, or, rather most of their lines and ecotypes, are salt sensitive, but mutations typically will increase sensitivity (Zhu 2000).

Technology in the form of improved DNA sequencing, especially so-called next-generation sequencing (NGS), is beginning to provide genome sequences, which, although they are not complete in an all encompassing sense, nevertheless contain all or nearly all (predicted) coding regions. Unlike *Arabidopsis thaliana* and *Oryza sativa*, species with extremophile characteristics include the moss *Physcomitrella patens*, *Populus* sp. (poplar) (Rensing et al. 2008; Tuskan et al. 2006) and two close relatives of *Arabidopsis*: *Thellungiella parvula* and the salt cress *Thellungiella salziginea* (formerly *T. halophila*) (Deng et al. 2009; Oh et al. 2010; Wang et al. 2010; Dassanayake et al. 2011b; Xie Q, personal communication). The analysis of these genomes, which is only just beginning, indicates several avenues that

show a strong correlation with the halophytic lifestyle or, to generalize, identifies gene complexity and structure that seem to reflect different plant habitus and ecological niche. We suggest that the results of comparative genomic studies will initiate breeding and genetic engineering avenues that have not been considered up to now.

The salt tolerance phenotype: how much NaCl is too much?

The fundamental physiological strategies by which halophytes cope with saline conditions such as avoidance, evasion and adaptation are also common to glycophytes (Zhu 2001). Tolerance mechanisms specific to halophytes involve a variety of morphological traits, such as succulence or salt glands (Breckle 2002). A tolerance classification system has been set up based on quantitative analyses of growth performances during and after the adaptation of halophytes to salt stress (Flowers et al. 1986). However, there is no unequivocal physiological basis that would allow a clear line to be drawn between glycophytes and halophytes, with the latter having themselves a large degree of variability in terms of growth in the presence of salt. For instance, moderate salt tolerance is found in crops such as sugar beet (*Beta vulgaris*, Chenopodiaceae), date palm (*Phoenix dactylifera*, Arecaceae) or barley (*Hordeum vulgare*, Poaceae) that can complete their life cycle in concentrations of 5 g l⁻¹ of total dissolved salts (TDS), or equivalent to approximately 80 mM NaCl (Ayers and Westcot 1985). In contrast, extreme halophytic species such as *Salicornia bigelovii* (Chenopodiaceae) generate remarkably high yields even when the soil solution exceeds 70 g l⁻¹ TDS (1,300 mM NaCl) (Glenn et al. 1985, 1991). Most glycophytes relevant to agricultural production have been ranked based on their yield reduction per unit in salinity relative to the maximum yield obtained under non-saline conditions (Maggio et al. 2007). A similar systematic classification does not exist for halophytes, to a large part due to the many degrees of adaptation that in the past had been expressed by distinctions between facultative and obligatory halophytic species. It is also important to point out that none of the definitions of halophytes (Kreeb 1974; Flowers and Colmer 2008) takes into account developmental phases with different salinity tolerance thresholds, or differential organ development in response to salinity. Such information is often also missing for glycophytes, including information on the relative sensitivity of shoot versus root, which can have important adaptive and agronomic implications (Dalton et al. 2000; Maggio et al. 2007). An increased shoot/root ratio at high salinity has been reported for *Atriplex* species, whereas a decrease has been found in Australian salt-marsh plants (as well as in the

glycophytes maize and soybean) (Redondo-Gómez et al. 2007).

The emergence of mechanisms

Even though traits associated with salt tolerance of halophytes have been extensively studied, as documented by the volume of publications, it has been difficult to define salt tolerance in morphological, physiological or molecular terms. The accumulation of Na^+ in a controlled fashion, as a driver for water uptake against low external water potential, is probably the mechanism that has early on, and subsequently most often, been identified as a requirement of the halophytic lifestyle (Flowers et al. 1977; Ungar and Pfeiffer 1991; Bohnert and Cushman 2000). Much attention has been given to the uptake and distribution systems of ions that can be toxic, and this has also been the major feature in attempting the classification of halophyte types. The discrimination of Na^+ versus K^+ , often credited as a major feature of halophytism, is an important though not sufficient determinant. *Salicornia europaea* and *Suaeda maritima* are considered true Na^+ accumulators, whereas the equally tolerant *Puccinellia distans* possesses a strong selective capacity of K^+ versus Na^+ at both low and high salinity (Tarasoft et al. 2007). Degrees of ion selectivity have been reported for halophytic and non-halophytic Chenopodiaceae (Flowers and Colmer 2008). Differences in Na^+ accumulation among dicotyledonous and monocotyledonous halophytes have also been documented (Greenway 1968) and used to distinguish, on a highly inconsistent scale, inclusion versus exclusion strategies to cope with a highly saline environment. In fact, early observations on the level of accumulation of NaCl into the shoots led to the classification of dicotyledonous halophytes as “includers”, since they accumulated more NaCl in shoot tissues compared to monocotyledonous plants, which were then termed “excluders” (Greenway 1968). Subsequently, this distinction was clarified by Greenway and Munns (1980), who recognized that monocots (mostly grasses) still use leaf Na^+ accumulation for osmotic adjustment, yet apparently to a lesser extent compared to halophytic dicots. Further studies with *Atriplex canescens*, however, demonstrated that the growth rate of different accessions was strongly correlated with their capacity for accumulating Na^+ and negatively correlated with their capacity to accumulate K^+ , highlighting remarkable interspecies variability (Glenn et al. 1992; Mata-González et al. 2001).

It is notable that almost no studies have exploited the use of genetics, either through crosses and segregation analyses, or by mutagenesis and selection systems, with very few exceptions (Cushman and Bohnert 1999;

Cushman 2001; Munns 2002). Importantly, the species used were either not halophytic or their other features precluded efficient identification of genetic loci involved in coping with high external Na^+ levels (Munns 2002).

Specific anatomical features may be responsible for salt tolerance in halophytes. Salt glands are typical morphological adaptations in black mangroves (*Avicennia* spp.), which excrete excess salts onto the leaf surface, yet this seems not to be a critical mechanism for the salt-tolerant red mangroves, *Rhizophora* spp. (Popp 1995; Popp and Albert 1995). Salt glands (Lipshitz and Waisel 1982; Balsamo and Thomson 1993, 1996) and salt bladders (Schirmer and Breckle 1982; Freitas and Breckle 1992; Adams et al. 1998; Orsini et al. 2011) may contribute by excreting up to and exceeding 50 % of the salt entering the leaf to cope with transitory imbalances of tissue NaCl (Glenn et al. 1999). In desert halophytes such as *Atriplex*, salt excretion may also have secondary adaptive functions (e.g., light reflection) that may protect the photosynthetic machineries (Osmond et al. 1980).

Ion accumulation and export

Succulence or reduction in leaf water content may function to dilute excess NaCl in leaf tissues (Kramer 1984) or to adjust the amount of NaCl necessary to establish a favorable osmotic flux (Glenn and O’Leary 1984; Glenn 1987). Regulation of water flux through the plant has emerged as an important component of salinity tolerance in halophytes. Developmental regulation of stomata density, size and closure has been functionally associated with stress tolerance (Farnsworth 2004; Yang et al. 2005). This trait has not been systematically analyzed for halophytes. It tends to be linked to excluder species. Halophytes in general exhibit reduced stomatal conductance compared to glycophytes, with transpiration further decreased as the exposure to salinity increases (Flowers and Yeo 1995; Serrano 1996). In fact, several halophytes, including the well-studied model *Mesembryanthemum crystallinum*, are capable of CAM metabolism that is stress induced and developmentally controlled to limit stomatal water loss (Adams et al. 1998). Although there have been several attempts to match morphological and physiological characters to specific halophyte habitats or growth strategies, the classification systems that have been developed still cannot predict degrees of halophytism.

A comparative physiological study

A recent study (Orsini et al. 2010) compared 11 wild relatives of *Arabidopsis thaliana* with respect to differences



Fig. 1 Plants of *A. thaliana* (Col-0) and *T. parvula* at different growth stages. Plants were grown in perlite and pumice, in a growth chamber with a photosynthetic photon flux of $500 \text{ mmol m}^{-2} \text{ s}^{-1}$ from cool-white fluorescent lamps and a 16-h light/8-h dark photoperiod. Plants of *T. parvula* at 42 days were grown on Turface[®] calcined clay (Profile Products, Buffalo Grove, IL, USA), in a growth chamber with a photosynthetic photon flux of $250 \text{ mmol m}^{-2} \text{ s}^{-1}$ from cool-white fluorescent lamps and a 16-h light/8-h dark photoperiod

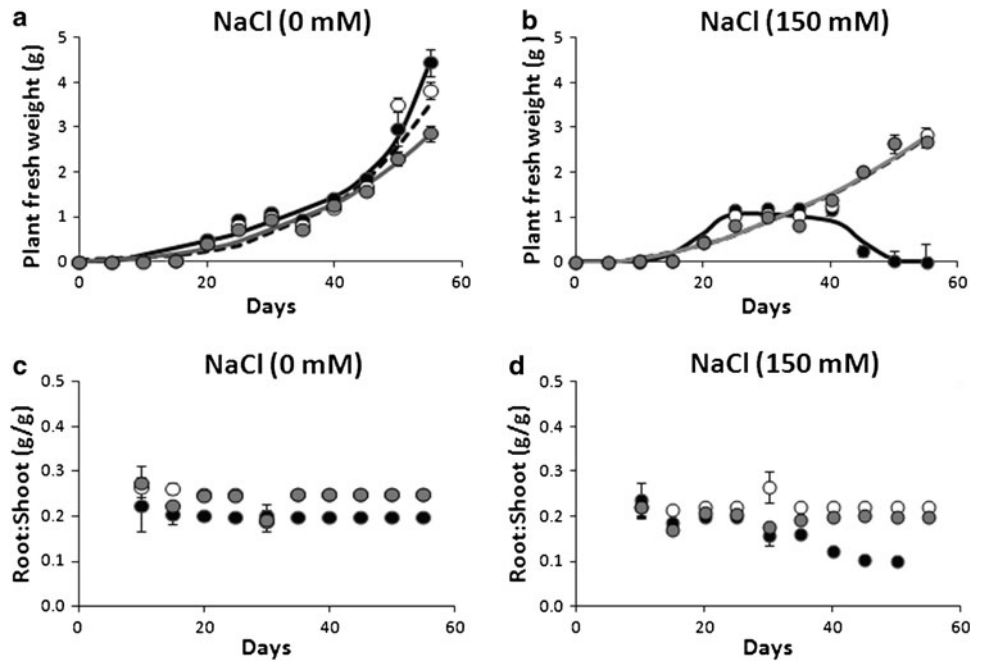
in growth, water transport properties, and ion accumulation to identify *Arabidopsis*-relative model systems (ARMS) that could serve as the genetic basis to advance our understanding of halophyte-specific mechanisms. It has been demonstrated that *T. parvula* (Fig. 1) has a true halophytic habitus comparable to *Thellungiella salsuginea*, a species that has been more extensively studied. The analysis presented by Orsini et al. (2010) added phenotypic

markers that allowed us to place several close *Arabidopsis* relatives with halophytic characters and an extremophile lifestyle into a framework for further studies. The halophytic habitus of these species (Fig. 2a, b) was associated with several morpho-physiological traits that have so far been largely underexplored in salinity tolerance research, including the ability to control transpiration water loss versus growth, reduced daily transpiration rates, and differential control of Na^+ versus K^+ fluxes (Volkov and Amtmann 2006). Among these qualities, the ability of these plants to maintain a close to constant root-to-shoot ratio at increasing salinity appears to be critical (Fig. 2c, d). A significant drop of the root-to-shoot ratio in *Arabidopsis*, due to a relatively higher sensitivity of roots versus shoots to salinity, appeared to coincide with overall salt-induced growth inhibition. We do not know how this and other morphological adaptation traits (see also the higher stomatal density in *T. parvula* and *T. salsuginea* compared to *A. thaliana*; Orsini et al. 2010) are functionally linked to the overall metabolic program that distinguishes halophytes from glycophytes. However, we imagine that the experience gained over the last decade in developing genetic systems based on insertion mutagenesis and the application of recently developed high-throughput DNA sequencing technologies (Oh et al. 2010; Dassanayake et al. 2011b) will rapidly lead to results from which we can begin to understand the evolutionary patterns that are at the basis of halophytic adaptations to extreme environments. Genomics, it seems to us, will become the tool that can bridge the gap that lets us go from correlative to mechanistic understanding.

Genomics takes center stage

The complexity of the halophytic lifestyle is best addressed by exploiting and further developing those genetic resources that can fit the principle of effective ARMS. Together with the ongoing generation of salt cress mutants with disruptions in osmotic regulatory properties, comparative analysis with other ARMS should be started. We argue that, for *T. salsuginea* (salt cress) and *T. parvula*, additional phenotypic and physiological tolerance markers should be established to go along with exhaustive transcriptome profiling to accompany the full genome sequence models that already exist for both *Thellungiella* species (Dassanayake et al. 2011b; Xie Q., personal communication). Comparative analyses and the juxtapositioning of responses to salinity by *Arabidopsis* and species in the category of ARMS will bring about an understanding of the mechanisms and will establish links between diverse tolerance traits and the genetic bases that are responsible for these characteristics (Bressan et al. 2001). These

Fig. 2 Effects of NaCl on shoot fresh weight (**a, b**) and root:shoot ratio (**c, d**) of *T. parvula*, *T. salsuginea* and *A. thaliana* (Col-0). Plants were grown in Perlite and Pumice in a growth chamber with a photosynthetic photon flux of $500 \mu\text{mol m}^{-2} \text{s}^{-1}$ from cool-white fluorescent bulbs and a 16-h light/8-h dark photoperiod. NaCl was added incrementally to the irrigation water every 5 days starting from 15 days after sowing, to a final concentration of 150 mM. Plants were harvested every 5 days until day 55. Gray circles *T. parvula*, white circles *T. salsuginea*, black circles *A. thaliana*. Values are mean \pm SE; $n = 9$



genetic/bioinformatics comparative analyses will be greatly enhanced by the addition of genome sequence information of additional ARMS with established extremophile phenotypes.

Several studies over the past decade have produced enigmatic results that can, with genomic hindsight, be rationalized. For example, a comparison of the transcription programs of two rice lines (salt-tolerant Pokkali and sensitive IR29) showed that both lines induced a very similar set of genes after a salt shock, but Pokkali terminated expressing this set of genes within a few hours and switched on genes in metabolism and development that are not indicative of severe stress responses. IR29 continued to express genes that have been labeled stress-inducible after a 150-mM NaCl treatment and died (Kawasaki et al. 2001). Another study using rice microarrays with $\sim 55\text{-K}$ probes compared a salt tolerant line FL478 with the sensitive IR29, revealing larger numbers of genes regulated in IR29 by salt stress, as well as different regulation of genes in flavonoid biosynthesis pathways and cell wall restructuring between the two lines (Walia et al. 2005). Similarly, a comparison of *A. thaliana* Col-0 and salt cress salt-dependent transcription programs indicated a large overlap in the types of genes induced, repressed or unaltered during salt stress treatments. The significant difference was that salt cress showed responses only at much higher salt levels than *Arabidopsis* and rapidly went back to a new state (Gong et al. 2005). One tentative conclusion in interpretation of these differences in response capacity was that there did not appear to be a battery of magical stress response and protection genes, that these close relatives are

distinguished by degrees of sensitivity by which different outcomes were orchestrated. However, the outcome was still determined by a common set of genes.

Genome and transcriptome sequences

The genome sequences for *T. salsuginea* and *T. parvula* are available in the form of chromosome models (Dassanayake et al. 2011b; Xie Q., personal communication, <http://thellungiella.org/>, <http://www.phytozome.net/thellungiella.php>). In both genomes, the amount of transcripts and deduced amino acid sequences exceeds the number of authentic transcript units by a few hundred putative additional genes. In both species, the numbers of genes and size of gene space are similar; differences exist mainly in the content of repetitive DNAs, which, in *T. parvula*, are less than 15 % of the genetic space, while they amount to more than 50 % in the case of *T. salsuginea*. A detailed analysis and comparison between the *A. thaliana* and *T. parvula* genomes identifies the “movement” of centromeric regions that accompanied the evolution of crucifers to result in a variable number of chromosomes (Mandáková and Lysak 2008; Dassanayake et al. 2011b). Other distinguishing aspects in genome organization include the proliferation of different transposable elements, and the emergence of species-specific putative reading frames for which no function is known in *T. parvula* or *A. thaliana*. In addition, sequences upstream of coding regions, for which functions have been determined in *A. thaliana*, are to a very large percentage different in *T. parvula*. It seems that gene

control elements have evolved in a different and novel way, and those differences appear to be a factor in how *T. parvula* copes with the abiotic stresses to which it is exposed (Oh et al., in preparation). A preliminary analysis of the *T. salsuginea* genome suggests that both *Thellungiella* species share the differences in promoter structure.

The notion about the potential importance of copy number variation (CNV) in shaping ecological adaptation and lifestyle has recently been substantiated by more detailed analyses of differences between *Arabidopsis* and *Thellungiella*; differences first in how the SOS system is engaged during stress (Oh et al. 2009; Dassanayake et al. 2011a) and by monitoring differences in genome structure not only at the macro level but also in the details of the uptake systems of sodium ions (Oh et al. 2010; Dassanayake et al. 2011b; Ali et al. 2012; Hong et al., in preparation).

One outstanding feature emerging from a comparison of the *Arabidopsis* and *Thellungiella* genome sequences concerns gene content, especially the number and type of gene duplications. For example, comparable numbers of tandem duplication events accumulated during the evolution of *Arabidopsis* and *Thellungiella*. What is dramatically different are the type of genes that have become, or have been retained as, duplicated sequences, and the number of genes that have been lost or failed to become duplicated, as exemplified by the different GO profiles of genes in species-specific tandem duplicated arrays between *A. thaliana* and *T. parvula* (Fig. 3a). *Thellungiella* added or retained after (random) duplication genes that are relevant to stress management and survival under abiotic stress, while duplications in the *Arabidopsis* genome focus on biotic stress defenses (Cannon et al. 2004; Dassanayake et al. 2011b). Further differences between *A. thaliana*, and *T. parvula* are observed when GO enrichment for transcript expression abundance is considered between orthologs (Fig. 3b).

A detailed comparison of selected genomic DNA regions in *A. thaliana*, *T. salsuginea* and *T. parvula* provided confirmatory results (Oh et al. 2010). First, an analysis by quantitative PCR revealed that, in a syntenic stretch of DNA in both *A. thaliana* and *T. parvula*, a number of genes showed higher expression under control conditions in *T. parvula* and a delayed but stronger induction of the genes at very high NaCl stress. Seemingly, expression strength and response threshold distinguish the two species. Even more revealing was the strong divergence in promoter structure between *Arabidopsis* and both *Thellungiella* species for the crucial salinity tolerance gene *SOS1* (Oh et al. 2010). In both *Thellungiella* species studied so far, *SOS1* shows very different regulation when compared with *A. thaliana*. *SOS1* and the entire SOS pathway constitute a critical component for salt stress

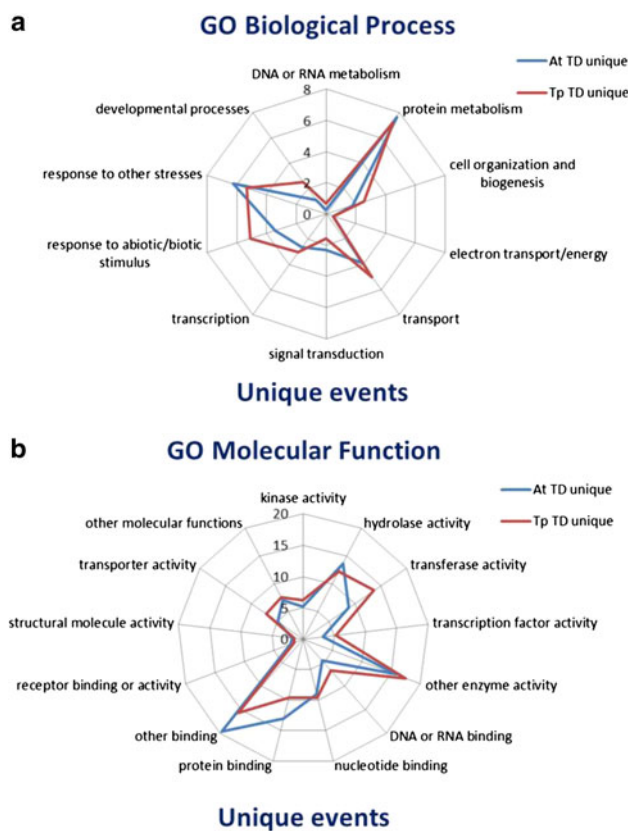


Fig. 3 Comparison of GO profiles in genomes and transcriptomes of *Arabidopsis thaliana* and *Thellungiella parvula*. **a** Distribution of TAIR GO slim-categories of genes in tandem duplicated arrays unique to either *A. thaliana* (At) or *T. parvula* (Tp) genomes. Y axes show the percentage of each categories per total number of genes in unique tandem duplicated gene arrays, which are 1,429 and 1,424 for At and Tp, respectively (modified after Dassanayake et al. 2011b). **b** GO enrichment in basal transcript level for high CNV. *A. thaliana* (3,459) and *T. parvula* (3,601) highly abundant transcripts ($p < 0.5$ with FDR < 0.05) were selected out of 19,935 orthologs for GO enrichment analysis against the total GO assignments for the *A. thaliana* genome as a background. Only significant enrichment for GO 'Molecular Function' is included for each transcriptome ($p < 0.01$ and FDR < 0.05)

adaptation. The activity of the Na^+/H^+ antiporter *SOS1* is regulated through protein phosphorylation by the *SOS2/SOS3* kinase complex which converts inactive *SOS1* to an active form (Bertorello and Zhu 2009). In both *Thellungiella* species, *SOS1* and *SOS2* transcript expression seems to be regulated very differently from the regulation in *A. thaliana*. Both genes are highly expressed in the absence of salinity stress, while stress leads to further increases in steady-state transcript amounts (H. Hong, and D.H. Oh, unpublished).

A second difference appeared, unexpectedly, in comparing the 5'-UTR regions between the three species. Both *Thellungiella* species are characterized by a high C/T-rich stretch within 100 nucleotides of the translation initiation codon, and in both *Thellungiella* species, a stable

stem-loop structure can form immediately upstream of the ATG codon. Further investigations showed that both features are found in other SOS1 genes of other halophytes, and are present only in halophyte SOS1 genes (Oh et al. 2009, 2010; Dassanayake et al. 2011b).

At present, such observations may be viewed as inconclusive or anecdotal unless they are backed by additional data. However, it can be pointed out that the higher transcriptional expression of genes in a species, line, or ecotype must be viewed as akin to a phenomenon that has recently attracted much attention. The term for this phenomenon, copy number variation (CNV), is applied to both gene copy number in a genome as well as to transcript abundance, both of which, on first principles, should increase protein amount. In fact, CNV has been associated with line differences in animal systems, underlying medically relevant symptoms (e.g., Kato et al. 2010; Waszak et al. 2010). CNV has also been invoked as a factor distinguishing maize from its non-domesticated relatives (Swanson-Wagner et al. 2010).

The case for CNV has been strengthened by the recent completion of the genome sequence of *Thellungiella parvula*, for which a first draft sequence has been published (Dassanayake et al. 2011a). A second version of *T. parvula* genome annotation is now available (<http://thellungiella.org/data/>). The first draft genome sequence and annotation of *Thellungiella salsuginea* have been prepared by two groups (Xie Q., personal communication; <http://www.phytozome.net/thellungiella.php>).

Preliminary comparison of *T. parvula* and *T. salsuginea* with their non-halophyte relative *A. thaliana* identified CNVs, in genome and transcriptome levels, of genes that are related to especially stress tolerance (Dassanayake et al. 2011b; Ali et al. 2012; Hong et al., in preparation). In addition to the shared functional classes in highly abundant transcripts, in the GO molecular function, transcription factor activity is enriched in *T. parvula*, while the category phosphatases activity is slightly higher in *A. thaliana* (Fig. 3b).

Higher basal level expressions of stress-related genes often coincide with genomic structural variations, such as tandem duplications and translocation duplications (Hong et al., in preparation). This confirms the hypothesis that assumed long ago the evolution by gene duplication—originally stated by Susumu Ohno (1970)—as a major driver of evolutionary changes. Duplications followed by sub- and neo-functionalizations of one of the duplicated genes appear as major mechanisms of evolutionary niche adaptations. For example, genes encoding HKT1, a Na⁺/K⁺ transporter, are tandemly duplicated in *T. parvula* and *T. salsuginea*. HKT1 is a single gene in *A. thaliana*. The duplicated copies showed both sub- and neo-functionalization in their gene expression pattern and protein functionality. One of the duplicated copies includes

Thellungiella-specific mutation of two amino acids near the second pore domain, which changed the transporter to favor K⁺ over Na⁺ (Ali et al. 2012). The overall expression levels of these copies are dramatically higher in both *Thellungiella* species than in *A. thaliana* (Ali et al. 2012; Hong et al., in preparation). Other examples of known stress-related genes that are duplicated in *Thellungiella* but not in *Arabidopsis* include genes encoding *AVP1*, *NHX8*, *MYB47*, *CBL10* and *KEA1* (Dassanayake et al. 2011a). Many of them show dramatically higher basal-level gene expression in *Thellungiella* than in *Arabidopsis* (Hong et al., in preparation).

It is noteworthy that these CNVs did not originate from whole genome duplication (WGD) events that most crucifers, including *Thellungiella* and *Arabidopsis*, experienced at least three times (Franzke et al. 2011). *Thellungiella* and *Arabidopsis* diverged after the most recent WGD event (Mandáková and Lysak 2008; Oh et al. 2010; Dassanayake et al. 2011a). Our initial observation on *Thellungiella* and *Arabidopsis* gene duplications confirms other studies that WGD resulted in enriched retention of genes involved in regulatory roles and higher numbers of protein–protein interactions, while tandem duplications tended to amplify genes encoding membrane proteins and enriched in responses to environmental stimuli (Freeling 2009; Rodgers-Melnick et al. 2012). In both species, tandem duplications were enriched in categories receptor and transporter activities, and underrepresented with respect to transcription factors, DNA/RNA binding and structural molecule activities (Dassanayake et al. 2011a).

In the analysis of the *Thellungiella* genomes, which is just beginning, one important aspect for future studies will be to find how the expression characteristics may have changed with respect to timing of expression and condition, and tissue or cell specificity. Especially, higher basal level expression of stress-related genes, even without stress, is a recurrently observed pattern distinguishing *Thellungiella* and *Arabidopsis* (Oh et al. 2009, 2010; Ali et al. 2012; Hong et al., in preparation). An exhaustive comparison of the basal-level transcriptomes between the two species will identify genes, of as yet unknown functions, potentially responsible for the phenotypic differences between the two species. Identification of regulatory elements and other genomic structural variations correlated with the observed differences in gene expression strength will reveal the evolutionary mechanism leading to the adaptation of a species to survive in extreme environments.

Towards applications

Until very recently, comparisons of plant genome sequences had to deal with species of widely different

evolutionary histories—*Arabidopsis* to *Oryza*, poplar or *Physcomitrella* (a moss) are examples. Gaps in our knowledge about the evolution of extremophile lifestyles have long existed. Only the availability of the *Thellungiella* genome sequences has made it possible to view different lifestyles in close relatives and to begin to understand the meaning of differences in gene number, gene type, and gene expression. This ability from comparative genomics will also help in understanding the structures of other species' genomes—*Oryza*, *Brachypodium*, *Pennisetum*, *Zea*, *Triticum* and *Hordeum* are examples. While these species in the *Poaceae* are related, it must be taken into account that they have been intensively manipulated over the last 10,000 years of agricultural practice (Rindos 1987; Allaby et al. 2008). At the other extreme are sequences of the many *A. thaliana* accessions (also termed ecotypes) that have been described and the genomes of different *Arabidopsis* species, such as *A. lyrata* (Hu et al. 2011; Cao et al. 2011). The comparison between *A. thaliana* and *T. parvula* and also the comparison with *T. salsauginea* (Q. Xie, personal communication) will provide a very different viewpoint and unprecedented opportunities not simply for comparative genomic studies but—even more importantly—for the future of crop improvement.

It is quite clear that the *Thellungiella* genome sequences signify evolutionary solutions of adaptation to an extremophile lifestyle. We consider this view and statement justified by pointing to the many duplicated genes with relevance to stress defenses. In fact, such a statement could equally be applied to, for example, poplar. Its genome includes, for example, a higher number of aquaporin genes than are seen in other sequenced plant genomes, and the gene complement for cell wall functions is also high.

The question then is about whether having available genome sequences of close relatives with different ways of environmental adaptation (“lifestyle”) means that they are significant messengers. Can genomes tell how, for example, abiotic stress-tolerant crops may be generated? Can we begin to understand the mechanisms of ‘tolerance’ acquisition from gazing at genome sequences?

One starting point can be the recognition that the 25,000–30,000 genes in angiosperm genomes are omnipresent. There exist order-, family- and species-specific genes in addition. Here, we face a problem, because these specific genes are absent from the *Arabidopsis* genome and their putative or possible functions are simply conjecture. However, the vast collection of *Arabidopsis* stress-relevant mutations that exist indicate the existence of conserved mechanisms about how to deal with abiotic stresses, and these genes exist in many or all higher plants. We consider the retained duplication events and expression characteristics in the *Thellungiella* species a fair representation of the genetic makeup of a stress-tolerant, in particular a salt stress-

tolerant, way of life. In a sense, we have known about meaningful tolerance mechanisms for a long time, from many studies, but all glimpses of genes, their functions, and the mechanisms could not be merged into a coherent picture.

Gene duplications and high or low expression alterations in gene control that together signify CNV are at the basis of tolerance (Kvitek et al. 2008; Conrad et al. 2010; Dasanayake et al. 2011a; Waterhouse et al. 2011). That leads to a question about transgenic alterations to, it seems, dozens of genes, mainly transcription factors that respond to different hormonal stimuli, by which hundreds of metabolic genes might be controlled to accomplish stress tolerance. These transcription factors would have to be engaged in a condition-specific way, and one problem could be that multiple inputs, e.g., high temperature and stomatal control, might have to be dealt with. Another level of complexity is provided by the more recent studies showing that stresses can have consequences in chromatin accessibility and activity, and showing the prevalence of small interfering RNAs that generate a different level of complexity (Ito et al. 2011; Luo et al. 2012).

For salinity stress tolerance breeding, or at least increased relative tolerances, another avenue is opening up. For many crop species, wide crosses between significantly different breeding lines or closely related species, e.g., domesticated and wild *Hordeum* sp., exist. In the thousands of lines from these crosses will be individuals that contain alleles for known stress tolerance-enhancing genes—either duplications or genes with altered expression characteristics. Quantitative RT-PCR approaches, or, even better, RNA-seq approaches, may be used to find those individuals. If, out of a population of thousands, 100 plants are identified, crosses can be made that lead to a pyramiding of desired traits in a relatively short time.

Looking beyond the salinity stress problem on which we have focused, the rapidity with which new genome sequences become available foreshadows the immediate future and, we think, a fundamental change in how crop breeding will be approached. The expansion of genome sequencing to non-model species has also opened doors for orphan crop improvement where these are staple food sources in some of the world's poorest regions. These crops are often grown in marginal lands characterized by abiotic stresses. Genome comparisons between extremophile genomes and the recently released orphan crop genomes for cassava and pigeonpea (*Cajanus cajan*) can lead to accelerated identification of genetic markers for molecular plant breeding or underlying conserved stress-tolerant strategies that can be important in crop improvement (Varshney et al. 2011; Prochnik et al. 2012).

Evolution has generated plants that thrive in very hostile environments. These extremophiles are relatives, often close relatives, of traditional crops. Their contrasting

lifestyles provide clues to the ease with which genome sequences and global expression profiles can be obtained to open an entirely new road for crop breeding. Our—still ongoing—comparisons of the *Thellungiella*–*Arabidopsis* pair of species suggest that genome sequence and transcription control can predict the environment in which a species will succeed.

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