A Novel Arabidopsis Vacuolar Glucose Exporter Is Involved in Cellular Sugar Homeostasis and Affects the Composition of Seed Storage Compounds\[W][OA]

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Subcellular sugar partitioning in plants is strongly regulated in response to developmental cues and changes in external conditions. Besides transitory starch, the vacuolar sugars represent a highly dynamic pool of instantly accessible metabolites that serve as energy source and osmoprotectant. Here, we present the molecular identification and functional characterization of the vacuolar glucose (Glc) exporter Arabidopsis (Arabidopsis thaliana) Early Responsive to Dehydration-Like6 (AtERDL6). We demonstrate tonoplast localization of AtERDL6 in plants. In Arabidopsis, AtERDL6 expression is induced in response to factors that activate vacuolar Glc pools, like darkness, heat stress, and wounding. On the other hand, AtERDL6 transcript levels drop during conditions that trigger Glc accumulation in the vacuole, like cold stress and external sugar supply. Accordingly, sugar analyses revealed that Aterdl6 mutants have elevated vacuolar Glc levels and that Glc flux across the tonoplast is impaired under stress conditions. Interestingly, overexpressor lines indicated a very similar function for the ERDL6 ortholog Integral Membrane Protein from sugar beet (Beta vulgaris). Aterdl6 mutant plants display increased sensitivity against external Glc, and mutant seeds exhibit a 10% increase in seed weight due to enhanced levels of seed sugars, proteins, and lipids. Our findings underline the importance of vacuolar Glc export during the regulation of cellular Glc homeostasis and the composition of seed reserves.

The transport of sugars across membrane barriers is essential for higher plants, because sugars represent transport and storage units of conserved energy and thus play a fundamental role during developmental processes and stress responses (Williams et al., 2000). The disaccharide Suc and the monosaccharide Glc are the most prominent soluble sugars in many plants and fulfill multiple functions in nutrition, signaling, and osmoregulation. In order to sustain proper development of nongreen sink tissues, plants have to distribute these sugars via long-distance transport, mostly in the form of Suc (Sauer, 2007; Dinant and Lemoine, 2010), which is taken up from the apoplastic into sink cells directly or as its hydrolysis products Glc and Fru (Sherson et al., 2003; Büttner, 2007). In addition, sugars have to be distributed at the cellular level depending on the actual requirements. Within a cell, carbohydrates are stored in chloroplasts in the form of starch and in the vacuole as soluble sugars. While considerable effort has been expended in elucidating the flux of carbon in and out of the starch pool in chloroplasts (Zeeman et al., 2007), our current knowledge of carbon allocation in the vacuole and the corresponding transport steps is rather limited. However, understanding the details of vacuolar sugar partitioning is of major relevance, since several agriculturally important plants, like sugar beet (Beta vulgaris; Doll et al., 1979; Briskin et al., 1985; Getz, 1991; Getz and Klein, 1995) and sugarcane (Saccharum officinarum; Thom et al., 1982), accumulate considerable amounts of sugars in the vacuoles of storage organs.

The large central vacuole of a typical mesophyll cell can represent up to 90% of the total cell volume. It contains water, ions, enzymes, and a variety of primary and secondary metabolites and therefore fulfills vital functions, such as pH regulation, storage of essential ions, structural support, osmoregulation, and cell enlargement. Hence, vacuoles provide additional space for transient and long-term storage of sugars.
Dehydration-Like Early Responsive to osmotic stress has been attributed to one of the sugar transporters characterized biochemically, but not yet demonstrated in Arabidopsis (Thom and Komor, 1984; Martinoia et al., 1987, 2000). Furthermore, determination of subcellular sugar concentrations in a variety of plants via nonaqueous fractionation (Gerhardt and Heldt, 1984) revealed that in leaves, Glc is predominantly found in the vacuole, while Suc is mainly found in the cytosol or is evenly distributed (Wagner, 1979; Heineke et al., 1994; Pollock et al., 2000; Voitsekhovskaja et al., 2006).

In addition to the daily fluctuations due to photosynthetic activity, the vacuolar allocation of soluble sugars is also strongly influenced by stress factors like temperature changes, desiccation, and salt (Ingram and Bartels, 1996). Especially cold stress causes a strong increase of cellular Glc levels (Kaplan et al., 2004), and recently, this cold-induced accumulation of Glc was attributed to the vacuole (Wormit et al., 2006).

The passage of sugars across biomembranes is primarily mediated by transport proteins. The corresponding carriers for the uptake of Suc (Sauer, 2007) and hexoses (Buttner, 2010) across the plasma membrane have been identified and characterized for a variety of plants. In contrast, little is known about sugar transport proteins in the tonoplast. In sugar beet, a putative hexose transporter has been identified, and in vitro colocalization studies suggested vacuolar localization (Chiou and Bush, 1996), but evidence for transport activity was not provided. Only very recently, two previously unknown Glc transporter families, the tonoplast monosaccharide transporters (AtTMTs) and the vacuolar Glc transporters, have been identified in Arabidopsis (Arabidopsis thaliana), and direct or indirect transport assays support an import function for both types of transporters (Wormit et al., 2007), a Ser residue at position 32 or 33 can be phosphorylated. Notably, phylogenetic analysis revealed the existence of ERDL6 orthologs in all plant genomes sequenced to date, including dicots, monocots, gymnosperms, ferns, and mosses (Supplemental Fig. S1).

AtERDL6 is widely expressed, especially in tissues with high metabolic turnover.

First, the Genevestigator Arabidopsis Microarray Database and Analysis Toolbox (Eidgenössische Hochschule Zurich) was queried for plantwide AtERDL6 expression levels. According to these data sets, relatively high AtERDL6 expression was found in many plant tissues and organs, including green tissues as well as nonphotosynthetic sink tissues at various developmental stages (Supplemental Fig. S2). In order to get a detailed picture of the AtERDL6 expression profile during plant development and at the tissue level, we generated transgenic reporter plants expressing the GUS gene under the control of the AtERDL6 promoter. Consistent with the microarray data, GUS reporter plants showed wide-
spread activity of the AtERDL6 promoter in leaves, roots, and flower organs at various stages of development. In addition, these reporter lines allowed precise allocation of AtERDL6 expression in various tissues, such as cotyledons, hypocotyls, and roots (except elongation zone) of seedlings and 3-week-old plants grown on petri dishes (already visible after 1 h of staining; Fig. 1, A–C), in all green leaves (rosettes, cauline leaves, and sepal; Fig. 1, D–H), in mature pollen grains and anther filaments (from flower stage 14; Fig. 1, G and J), in sepal (from flower stage 13; Fig. 1), in pedicels starting at flower stage 15 (Fig. 1G), in the abscission zone of flower leaves from flower stage 16 (Fig. 1G), and in seed pod walls (Fig. 1K).

Taken together, our analyses demonstrated a widespread pattern of AtERDL6 expression especially pronounced in tissue types and developmental phases with high metabolic turnover, like seed germination, seedling establishment, pedicel and filament elongation, abscission, and wound response.

Figure 1. AtERDL6 expression pattern. Histochemical localization of AtERDL6 promoter-driven GUS expression is shown. A, Germinating seed with GUS activity in the emerging radicle. B, Two-day-old seedling with overall GUS expression. C to E, Five-day-old seedling with strong GUS staining in hypocotyl and cotyledons, root (except elongation zone), and root tip. F, Three-week-old plate-grown plant. G, Inflorescence and developing siliques. The inset shows an enlargement of the abscission zone of flower leaves and siliques. H, Mature rosette leaf. I, Cauline leaf. J, Flower with strong GUS staining in sepal, anthers, filaments, and pollen. K, Silique with GUS staining in seed pod walls. L, GUS activity in a mature leaf after wounding. In all experiments, GUS staining was stopped after 2 to 4 h and analyzed by light microscopy. Bars = 100 μm in A and J (inset), 200 μm in B, D, E, G (inset), K (inset), and L, 1 mm in C, I, and J, and 2 mm in F, G, H, and K.

AtERDL6 Is a Transporter of the Vacuolar Membrane

Next, we determined the subcellular localization of AtERDL6 by transient expression of an AtERDL6-GFP fusion construct in Arabidopsis and tobacco (Nicotiana benthamiana). As depicted in Figure 2, Arabidopsis mesophyll protoplasts transiently expressing the AtERDL6-GFP fusion construct showed GFP fluorescence only in the tonoplast, clearly visible after protoplast lysis by mild osmotic shock as a ring with chloroplasts (red autofluorescence) attached at the outside (Fig. 2B). Additionally, tobacco leaves were infiltrated with an Agrobacterium tumefaciens culture harboring an AtERDL6-GFP fusion construct. Confocal sections of transformed epidermal cells showed GFP fluorescence of the vacuolar membrane that does not encompass chloroplasts (Fig. 2C). Furthermore, enzymatic removal of the cell wall and lysis of the protoplasts demonstrated that, also in tobacco, the AtERDL6-GFP fusion protein was clearly confined to the vacuolar membrane (Fig. 2D).
AtERDL6 Activity Affects Vacuolar Glc Levels

In order to examine the precise molecular function of the putative sugar transporter AtERDL6, we obtained two independent mutant lines, SAIL_528_F09 and SALK_106049, carrying T-DNA insertions at positions +2,262 (exon 12) and +3,139 (exon 17), respectively (Supplemental Fig. S3). Genomic PCR analyses verified homozygosity for the T-DNA alleles, and RT-PCR analyses confirmed the absence of full-length AtERDL6 transcripts for both mutant lines (Supplemental Fig. S3).

Due to the demonstrated vacuolar localization of AtERDL6 and its homology to known MSTs, we examined the effect of the erdl6 knockout on cellular sugar homeostasis. To this end, we quantified the content of soluble sugars in leaves from wild-type and AtERDL6 mutant lines. As shown in Figure 3A, levels for Suc, Fru, and myoinositol were similar in the wild type and mutants. In contrast, a drastic increase by a factor of 2 was measured for the Glc content in both AtERDL6 knockout lines (Fig. 3A).

In order to distinguish whether the lack of AtERDL6 affects whole cell Glc levels or results in vacuolar Glc accumulation, we determined sugar concentrations in subcellular compartments. Applying nonaqueous fractionation (Gerhardt and Heldt, 1984), we showed that the observed Glc increase in AtERDL6 mutants was mainly due to an accumulation in the vacuole (398–402 ng mg⁻¹ fresh weight in AtERDL6 mutants and 140 ng mg⁻¹ fresh weight in the wild type), while in the extravacuolar space, the difference in Glc levels was only marginal (32–34 ng mg⁻¹ fresh weight in AtERDL6 mutants and 22 ng mg⁻¹ fresh weight in the wild type). Thus, in wild-type leaves, 86% of the total Glc is allocated to the vacuole, while in the AtERDL6 mutants, the vacuolar Glc content accounts for 92% to 93% (Table I).

This increase of vacuolar Glc in plants lacking AtERDL6 strongly points toward an export function for this transporter. To further support this idea, we...
We then addressed the question of whether *AtERDL6* expression is regulated by soluble sugars, which are a measure of the cellular energy status. Therefore, mature Arabidopsis leaves were incubated in 2% Glc, 2% Suc, or Murashige and Skoog (MS) medium without sugars, and the relative amounts of *AtERDL6* transcripts were determined by RT-PCR. As compared with the mock treatment, both Glc and Suc led to reductions of *AtERDL6* expression after 2 h of about 40% and 30%, respectively, which were still detectable after 6 h (Fig. 4D). Hence, *AtERDL6* expression is down-regulated by high external sugar concentrations, a condition where sugar release from the vacuole is not required.

**Figure 4.** *AtERDL6* expression is significantly induced during darkness, heat stress, and wound response and down-regulated by sugars and cold stress. For all treatments, total RNA from 4-week-old rosette leaves was used to determine *AtERDL6* transcript levels by quantitative real-time PCR. A, Induction of *AtERDL6* expression after 3 and 24 h of darkness. B, Transcriptional induction by heat stress. Plants were incubated at normal (23°C) and elevated (37°C) temperatures for 6 h. C, Strong up-regulation of *AtERDL6* expression by sugars. Leaves were incubated in liquid MS medium containing no sugars (black bars), 2% Glc (light gray bars), or 2% Suc (dark gray bars) for 2 or 6 h. E, Cold response of *AtERDL6*. Leaves were harvested 5 h after the onset of light before (day 0, 22°C) and during (1, 2, 4, and 8 d at 4°C) cold treatment. Shown are *AtERDL6* expression values relative to day 0 (no cold treatment). Data represent mean values ± se of four independent biological replicates using *LDL3* (At4g16310) as a reference gene for A to D and of three independent experiments using *EF-1α* (At5g60390) as a reference gene for E.

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**Table 1.** Subcellular Glc distribution determined by nonaqueous fractionation and HPAEC-PAD

Mature rosette leaves were assayed by nonaqueous fractionation according to Krueger et al. (2009). Values represent means ± se of five biological replicates each for the wild type and mutant lines. Due to the much smaller volumes of chloroplasts and cytosol compared with vacuoles, the so in these compartments is higher (Tilsner et al., 2005); thus, it was not possible to exactly determine whether the slightly elevated extravacuolar Glc levels are caused by changes in chloroplasts, the cytosol, or both.

| Plant Line | Vacular Glc | Extravacular Glc |
|------------|-------------|-----------------|
| Wild type  | 140 ± 20 (86%) | 22 ± 3 (14%) |
| SALK_106049 | 409 ± 79 (92%) | 34 ± 6 (8%) |
| SAIL_528_F09 | 398 ± 45 (93%) | 32 ± 4 (7%) |
It has previously been shown that several soluble sugars accumulate in vacuoles of Arabidopsis leaves upon exposure to low temperature (Wormit et al., 2006). In order to test if the application of cold stress also influences the expression level of the putative vacuolar Glc exporter ATERDL6, we determined ATERDL6 transcript levels in plants that were transferred from normal temperature (22°C; day 0) to 4°C for 8 d. As shown in Figure 4E, already after 1 d, ATERDL6 expression was reduced to about 40%, and it dropped continuously until a level of about 15% was reached at day 4, which remained constant for the rest of the sample period.

In summary, factors that affect vacuolar Glc allocation also influence ATERDL6 transcript levels.

**Aterdl6 Mutants Are Affected in Glc Export from the Vacuole Induced by External Signals**

The impact of low temperature on vacuolar Glc accumulation and on ATERDL6 expression prompted us to investigate the physiological consequences of the lack of ATERDL6 export function during the plant response to cold stress. Therefore, plants were kept at 4°C for 4 d to induce vacuolar Glc accumulation. Then, we transferred the plants to 22°C and compared the changes in Glc levels of wild-type and Aterdl6 mutant plants during the period of rewarming, when Glc is released from the vacuole. In comparison with the wild type, the rate of Glc release from the vacuole was significantly slower in the Aterdl6 mutants (Fig. 5A). While in wild-type plants, the Glc level dropped down to 37% after 7 h and to 16% after 14 h of rewarming, the Aterdl6 mutants still had 57% to 58% and 40% to 41% of the initial Glc level after 7 and 14 h, respectively, indicating that a major route of Glc export from the vacuole is missing in Aterdl6 mutants.

In contrast to the vacuolar Glc accumulation during cold stress, responses to elevated temperatures consume cellular energy and thus require the export of Glc from the vacuole. Accordingly, we found that Aterdl6 mutants have a less marked mobilization of vacuolar Glc upon heat stress than wild-type plants. As seen in Figure 5B, wild-type plants reduced their Glc level to 39% after 6 h of heat stress, while mutants showed a reduction to only 62% to 71%. This difference was even more apparent when plants were kept in the dark during heat treatment. Under these conditions, wild-type plants reduced their Glc level to 27%, while mutants showed a reduction to only 44% to 46% (Fig. 5B).

Taken together, sugar measurements in Aterdl6 mutants demonstrate that during adaptation to ambient temperatures, AtERDL6 is responsible for the export of a significant portion of Glc from the vacuole.

**The ERDL6 Ortholog of Sugar Beet Also Mediates Glc Export When Expressed in Arabidopsis**

Phylogenetic analyses revealed that ERDL6 orthologs are present in all plant genomes (Supplemental Fig. S1). Therefore, we investigated if an ERDL6-type transporter from another plant species also mediates Glc export from the vacuole. To this end, we used the previously identified cDNA clone (accession no. U43629, locus BVU43629) from sugar beet (Chiou and Bush, 1996) coding for Integral Membrane Protein (BvIMP), which is 80% identical and 92% similar to AtERDL6, for overexpression in Arabidopsis. U43629 transcript levels were monitored by northern-blot analysis, and two lines (lines 6 and 8) displaying...
substantial expression of the \textit{BvIMP} gene (Fig. 6A) were selected for further analysis. In order to measure the Glc levels in these overexpressor lines, we used seedlings grown in liquid culture (Wingenter et al., 2010). As shown in Figure 6B, both overexpressor lines showed significant decreases in Glc, representing about 66% to 69% of the wild-type level, similar to the transgenic lines overexpressing \textit{AtERDL6} (Fig. 3B).

Cold treatment induces Glc relocation into the vacuole, as previously demonstrated. Therefore, we wanted to test if lines overexpressing the sugar beet cDNA show differences in Glc release during rewarming after cold incubation. Five- to 6-week-old plants were kept at 4°C for 3 d and then transferred to 22°C at the beginning of the dark phase. As seen in Figure 6C, in wild-type plants, the level of Glc dropped to 65% and 35% after 7 and 14 h, respectively, demonstrating release and metabolization of Glc, which previously accumulated in the vacuole during the cold treatment. However, lines expressing \textit{BvIMP} displayed a more pronounced Glc release, indicated by the lower Glc levels being only 44% and 17% (of the initial value) after 7 and 14 h, respectively (Fig. 6C). This difference strongly suggests a stimulated Glc export capacity at the tonoplast in these lines. No differences between the wild type and overexpressors were observed for the release of Suc, which also accumulates during the cold (data not shown). Consistent with the proposed export function, \textit{BvIMP} overexpression resulted in the opposite effect as observed for the \textit{Aterdl6} mutant lines (Fig. 5A), namely, a faster decrease in cellular Glc levels due to a more rapid release from the vacuole.

\textbf{Lack of \textit{AtERDL6} Export Activity Causes a Delay of Seed Germination in the Presence of Glc}

Since our GUS reporter plants demonstrated a high expression level of \textit{AtERDL6} in germinating seeds (Fig. 1A), we examined the \textit{Aterdl6} mutants for detectable changes during this vital developmental process. While germination rates of mutant and wild-type seeds were nearly identical on 2% sorbitol and 2% Suc (data not shown), \textit{Aterdl6} seed germination was significantly delayed with increasing amounts of Glc in the growth medium (Fig. 7). In the absence of sugars, wild-type seeds started to germinate when incubated for 2 d in light (DIL) and reached 40% germination at 3 DIL (Fig. 7, top panel). It has been shown before that Glc delays seed germination in Arabidopsis (Dekkers et al., 2004). Accordingly, 2% Glc in the growth medium slightly delayed wild-type seed germination, reaching 70% at 4 DIL (Fig. 7, middle panel). This delay was even more pronounced in the presence of 7.2% Glc, where 50% germination was reached only at 7 DIL (more than 5 d later than in the wild type).

\textit{Aterdl6} Mutants Show a Significant Elevation of the Major Seed Storage Compounds

Due to the observed difference in seed germination, we also analyzed seed sugar composition as well as protein and lipid content in \textit{Aterdl6} mutant lines. As shown in Table II, mutant lines contain more than

\begin{figure}[h]
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\caption{Overexpression in Arabidopsis of the \textit{ATERDL6} ortholog \textit{BvIMP} supports the Glc export function from the vacuole. A, Northern-blot analysis of Arabidopsis lines 6 and 8 overexpressing the sugar beet clone U43629, coding for \textit{BvIMP}. Total RNA was isolated from 5-week-old plants; rRNA represents the loading control. B, Glc levels in the wild type (WT) and \textit{BvIMP}-overexpressing lines grown in liquid culture. Wild-type plants (black bars) and \textit{BvIMP}-overexpressing lines 6 (light gray bars) and 8 (dark gray bars) were grown in standard liquid culture medium for 12 d. Each data point represents the mean value of three independent biological experiments ± ss. C, Release of Glc after cold-induced accumulation in the vacuole. Both the wild type (black bars) and \textit{BvIMP}-overexpressing line 6 (light gray bars) were incubated for 3 d at 4°C under long-day conditions and transferred to 22°C for rewarming after the start of the dark phase (0 h) at day 3. Leaf Glc contents were quantified by ion chromatography. Each data point represents the mean value of three independent biological experiments ± ss. FW, Fresh weight.}
\end{figure}
experiments with 80 to 140 seeds per line.

Figure 7. AtERDL6 mutant seeds display enhanced sensitivity to external Glc during seed germination. Wild-type (WT) and AtERDL6 mutant seeds were germinated on MS plates containing no sugar, 2% Glc, and 7.2% Glc. Glc reduces the germination rate of wild-type seeds (black bars). However, a further delay can be seen with seeds of mutant lines SAIL_528_F09 (light gray bars) and SALK_106049 (dark gray bars) germinating in the presence of 2% Glc, which is even more prominent with 7.2% Glc. Data represent mean values ± s of three replicate experiments with 80 to 140 seeds per line.

double the amount of Glc compared with the wild type, whereas the amount of Suc was unchanged. Further analyses of the mutant seeds revealed that the total protein content is increased by more than 50% and also that the lipid content is more than 10% higher than in the wild type. Most interestingly, as a consequence of elevated soluble sugars, proteins, and lipids, the AtERDL6 mutant seed weight is increased significantly, as determined by the “thousand seed weight” being 16.7 ± 0.6 mg for the wild type and 18.4 ± 0.4 and 18.5 ± 0.4 mg for the mutant lines SAIL_528_F09 and SALK_106049, respectively (Table II). Moreover, this increase in single seed weight is not at the expense of total seed quantity, as shown by the nearly unchanged number of seeds per silique (61.2 ± 1.2 in the wild type versus about 63 ± 1.2 in the mutant lines) and the total seed weight per plant (191 ± 16 in the wild type versus 222 ± 16 and 223 ± 14 in the mutant lines). Thus, the total seed yield per plant is increased by more than 10% in the AtERDL6 mutant lines (Table II).

DISCUSSION

Sugars have essential functions in a number of plant cellular processes, including storage units of energy and building of structural components and signal molecules. Therefore, metabolic conversion, storage, and compartmentalization have to be tightly regulated, which often requires the transport of sugars across biomembranes. Consequently, plant genomes contain a great number of genes coding for sugar transport proteins, which sometimes form large families according to their specific functions as determined by substrate specificity, subcellular localization, and/or transport mechanism (Büttner, 2007; Sauer, 2007). While numerous transport proteins that import sugars into cells and cellular compartments have been identified to date, little effort has been devoted to identifying transporters that mediate sugar flux in the opposite direction, out of cells or organelles.

In this paper, we present the molecular identification and functional characterization of a previously unknown vacuolar Glc exporter, AtERDL6. A survey of public plant sequence databases revealed that genes coding for ERD6-like transporters exist in all plant genomes sequenced so far, including higher plants, ferns, and mosses (Supplemental Fig. S1), underlining a fundamental function of this type of sugar transporter.

Analysis of GUS reporter plants revealed a widespread expression pattern of AtERDL6 (Fig. 1) that was particularly prominent in tissues and developmental phases with high metabolic turnover, which included germinating seeds, young seedlings, elongating ped-
icels and filaments, programmed and induced cell death in abscission zones, and wounded tissues. Our results are supported by AtERDL6 expression data found in public microarray databases (Supplemental Fig. S2) and indicate a role of this new sugar transporter in cellular carbon distribution. Localization of a translational GFP fusion clearly assigned AtERDL6 to the tonoplast in Arabidopsis and tobacco (Fig. 2). This finding is in agreement with results from a recent proteome study, where AtERDL6 was identified in the vacuolar fraction (Jaquinod et al., 2007).

**AtERDL6 Regulates Cellular Glc Homeostasis in Response to Several Environmental Factors**

In support of a role in Glc export from the vacuole, we found a strong transcriptional regulation of AtERDL6 expression in response to conditions that alter cellular sugar levels and require the activation of cellular carbohydrate stores. It is long known that exposure to drastic temperature changes induces profound adaptations in carbon metabolism (Ristic and Ashworth, 1993). In Arabidopsis, cold stress at 4°C leads to a drastic increase in soluble sugar levels within the first 12 h (Kaplan et al., 2004). A similar effect on primary metabolism is also seen under heat stress, although cold stress appears to cause a more dramatic metabolic response (Shulaev et al., 2008). However, while Suc (and many other soluble sugars like Fru, maltose, and raffinose) accumulates under both temperature treatments, Glc is accumulating in the cold but is decreasing under heat stress (Kaplan et al., 2004). It is discussed that Glc is released from the vacuole to sustain Suc production in the cytosol, which in turn serves as an osmoprotectant in maintaining cell membrane integrity and cellular function under stress (Ruan et al., 2010). We determined AtERDL6 expression levels under both temperature stress conditions. In line with its Glc export function, cold stress, which causes Glc accumulation in the vacuole, strongly represses AtERDL6 (Fig. 4E). Interestingly, the closely related AtERD6 gene (At1g08930) shows an opposite regulation and is induced during cold stress (Kiyosue et al., 1998). This might indicate different functions within the ERD6-like transporter family with respect to substrates and transport direction. In contrast to its cold repression, we found a strong AtERDL6 induction when plants were exposed to elevated temperatures and consumed Glc (Fig. 4C), coinciding with the decrease in cellular Glc levels under these conditions.

Wounding induces a wide number of cellular responses, including respiration, cell division, and biosynthesis of secondary metabolites. Sealing wound sites and repairing damaged tissues requires metabolic energy provided by pools of soluble sugars (Roberts and de Bruxelles, 2001). We found that wounding triggered a more than 7-fold increase of AtERDL6 expression (Fig. 4C). Interestingly, treatment with methyl jasmonate, a phytohormone regulating the plant’s response to wounding, pathogen attack, and other stress conditions, also elicits a strong increase in AtERDL6 expression (1 h, 2.2-fold; 3 h, 3.5-fold; NASCARRAYS-174).

Another condition that requires access to cellular energy stores is darkness. At the light-to-dark transition and during the night, transitory starch represents the main pool of stored energy. However, vacuolar Glc also can serve the same purpose, especially for immediate compensation of photosynthesis switch off. We measured AtERDL6 transcript abundance after a 3-h dark period interrupting the normal light cycle during the day as well as after a prolonged night. Both conditions of “light starvation” led to a strong induction of AtERDL6 expression (Fig. 4A).

Since we could not distinguish whether the observed dark induction is a direct effect of the absence of light or is indirectly mediated by the drop in photosynthetic products, we tested the effect of sugars on AtERDL6 expression. We could measure a downregulation of AtERDL6 by feeding leaves with the primary metabolites Glc and Suc (Fig. 5), indicating that AtERDL6-mediated Glc export from the vacuole is controlled via sugar signaling. Consistent with this finding, a significant regulation of AtERDL6 by sugars was also found in transgenic plants with mutations related to sugar metabolism. The phy3 mutant, which harbors a defective copy of the Suc transporter SUC2 gene, accumulates large amounts of soluble sugars. In this mutant, AtERDL6 transcript levels are reduced to about 50% (Lloyd and Zakhleniuk, 2004). The Snf1-related protein kinase AKIN10 controls reprogramming of transcription in response to darkness, sugar, and stress conditions. Transgenic AKIN10 overexpression, which confers enhanced starvation tolerance (Baena-González et al., 2007), triggers a 2.5-fold increase of AtERDL6 expression (Gene Expression Omnibus series GSE8257), which in turn enhances the availability of vacuolar Glc. Finally, in independent studies investigating the effect of Glc treatment on the transcriptome level, AtERDL6 expression is markedly affected. Price et al. (2004) found a more than 5-fold reduction by 3% Glc, which was not seen with the nonmetabolizable analog 3-O-methylglucose. Similarly, Li et al. (2006) reported 4- and 5-fold reductions after treatment with 3% Glc for 2 and 6 h, respectively, which could not be observed with 3% mannitol. Interestingly, also the grapevine (Vitis vinifera) monosaccharide transporter VvHT1 was shown to be regulated by Glc at both the transcriptional and posttranscriptional levels (Conde et al., 2006).

In conclusion, our studies on AtERDL6 transcriptional regulation revealed a strong responsiveness to changes in the cellular sugar status that require instant access to vacuolar sugar pools.

**Physiological Role of the Vacuolar Glc Exporter AtERDL6**

To verify the role of AtERDL6 as a vacuolar Glc exporter in planta, we have chosen a reverse genetics
approach and isolated two independent mutant lines, SAIL_528_F09 and SALK_106049, in which AtERDL6 expression was absent (Supplemental Fig. S2). Analysis of cellular sugar levels revealed that while most of the common sugars were not affected in the Aterdl6 mutants, a clear increase in cellular Glc levels was detectable, indicating that in the mutants a transport activity at the tonoplast is missing, which traps Glc within the vacuole. Moreover, nonaqueous fractionation clearly demonstrated that, in contrast to wild-type leaves, which allocate 86% of the total Glc in the vacuole, in Aterdl6 mutant leaves, 92% to 93% of the total Glc is found in the vacuole (Table I). Consistent with AtERDL6-driven Glc export from the vacuole, we found markedly decreased Glc levels in AtERDL6-overexpressing lines (Fig. 3B). However, we also observed lower levels of Fru and Suc in these over-expressor lines (Fig. 3B). It is unlikely that AtERDL6 also directly mediates the export of Fru and Suc, since we could not measure an accumulation of these sugars in Aterdl6 knockout plants. It seems more likely that the excessive export of Glc from the vacuole alters cytosolic Glc signaling/homeostasis and thus indirectly affects Fru and Suc levels. To further substantiate our model of AtERDL6-driven Glc export from the vacuole, we analyzed the sugar levels in mutant lines exposed to cold stress that was shown to affect AtERDL6 expression. Since Glc accumulates in the vacuole during cold treatment, we expected an effect on Glc release from the vacuole during rewarming of the plants. In fact, both Aterdl6 mutant lines displayed significantly slower reductions of cellular Glc levels as compared with the wild type, which can be attributed to less efficient export from the vacuole (Fig. 5A). Similarly, upon heat stress (and even more obviously in combination with darkness), the liberation of the vacuolar Glc pool was significantly slower in the Aterdl6 mutants as compared with the wild type (Fig. 5B). The fact that in Aterdl6 mutants, Glc export from the vacuole is not completely absent is most likely due to one (or more) of the 18 other members of the AtERDL6 gene family, some of which show partial overlap in their expression patterns, according to public microarray data sets (Genevestigator).

Finally, we also addressed the physiological function of the AtERDL6 ortholog BvIMP from sugar beet. BvIMP overexpression in Arabidopsis resulted in a similar decrease of cellular Glc levels as observed for AtERDL6 overexpressors. In addition, when BvIMP overexpressors were subjected to low temperatures in order to induce vacuolar Glc accumulation and subsequently brought back to normal growth temperatures, the cellular Glc levels dropped significantly faster as compared with wild-type plants, indicating a faster Glc release from the vacuole during rewarming due to additional export capacity (Fig. 6).

All our results are in line with an export activity for AtERDL6 mediating Glc transport from the vacuole to the cytosol, and a lack of this transport step results in significant accumulation of Glc in the vacuole. However, Aterdl6 mutant lines developed completely normally and were indistinguishable from the wild type under standard growth conditions as well as after application of stress conditions like cold shock, drought, or salt stress (data not shown). Since AtERDL6 transcript levels are relatively high in mature seeds (NASCARRAYS-69, NASCARRAYS-183, NASCARRAYS-195, NASCARRAYS-497; Gene Expression Omnibus series GSE15700), we examined the impact of different sugars on seed germination. Interestingly, with increasing amounts of Glc in the medium, the seed germination rate was significantly delayed (Fig. 7). This effect could not be detected with comparable amounts of Suc or sorbitol (as osmotica), indicating the involvement of a Glc-specific signaling pathway.

Several recent studies revealed a negative influence of external Glc on seed germination. The production of sugars from starch is inhibited by increasing sugar levels in germinating cereal seeds by the repression of α-amylases (Karrer and Rodriguez, 1992). In germinating Arabidopsis seeds, the presence of exogenous Glc significantly retards the mobilization of seed storage lipid (To et al., 2002). Dekkers et al. (2004) have shown that Glc acts on embryo growth and that the inhibition of seed germination is independent of hexokinase function. Furthermore, exogenous Glc blocks seed germination by delaying the decline in endogenous abscisic acid concentration (Price et al., 2003). Thus, if germinating on high external Glc concentrations, seeds might sequester excess Glc from the cytosol into the vacuole to avoid these negative effects via Glc signaling. However, the vacuolar buffering capacity for Glc might be exhausted in Aterdl6 mutants, which already have a constantly elevated Glc level in the vacuole and, therefore, are more susceptible to external Glc, which then causes the observed delay in seed germination.

Due to the higher Glc sensitivity during seed germination, we have analyzed Aterdl6 mutant seeds in more detail and found that the major seed storage compounds were elevated (i.e. Glc by more than 100%, proteins by 53%, and lipids by 10%), while Suc was unchanged. This rise in storage compound deposition also led to an increase in single seed weight and total seed yield per plant of about 10% (Table II). This is in perfect agreement with the effect that was observed after overexpressing the tonoplast monosaccharide importer TMT1 (Wingenter et al., 2010), which also led to a significant increase of vacuolar Glc levels as well as lipids and proteins, causing a seed weight gain similar to that in Aterdl6 knockouts. Further analyses will be needed to show if the increase in Glc, proteins, and lipids in Aterdl6 mutants is balanced by a compensatory decrease of other components (e.g. secondary metabolites).

In summary, this paper presents the characterization of the vacuolar Glc exporter AtERDL6, which localizes to the tonoplast and is transcriptionally regulated by a number of internal and external factors that entail vacuolar partitioning of Glc. Lack of AtERDL6 causes...
significant increases in vacuolar Glc levels and negatively affects the release of vacuolar Glc during the heat stress response and during a rewarming phase after a cold period. Our findings underpin the important role of vacuolar Glc export during the regulation of cellular Glc homeostasis and the composition of seed reserves.

**MATERIALS AND METHODS**

**Strains and Growth Conditions**

Escherichia coli strain DH5α (Hanahan, 1983) was used for cloning. Arabidopsis (Arabidopsis thaliana) ecotype Columbia was grown in potting soil in the greenhouse or on agar medium in growth chambers at 22°C and 55% relative humidity under short-day (8 h of light/16 h of dark) or long-day (16 h of light/8 h of dark) conditions, as indicated. Arabidopsis transformation was performed with Agrobacterium tumefaciens strain GV3101 (Holsloot et al., 1980) using the floral dip method (Clough and Bent, 1998).

**Cloning of the AtERDL6 cDNA**

The full-length AtERDL6 cDNA (GenBank accession no. AY124845) was obtained using gene-specific primers AtERD67g-10_OBSPI (5’-GAATCAGGATCAGTGGTGGATGATAATGAGGAGGC-3’) and AtERD67c+1199r_OBSPI (5’-GGGATCCATGATATCAACAGAGTTAAGTCTC-3’) and total RNA from leaves to amplify the AtERDL6 open reading frame by RT-PCR. The corresponding 1,472-bp PCR product was cloned into pGEM-T Easy and sequenced, yielding pSR6740u.

**Generation of AtERDL6 Promoter/GUS Lines and Histochemical Localisation of GUS Activity**

In order to generate AtERDL6 promoter/GUS plants, a 2-kb fragment upstream of the AtERDL6 coding region was amplified by PCR using primers ERD67g-2056fs_PstI (5’-GATCTTGGTAGCTGCAGACGATACATTGC-3’) and ERD67g-11er_NcoI (5’-CATCCCTGAAACCCACGCTTGGTGCAT-3’) and genomic DNA, cloned into pGEM-T Easy, and sequenced, yielding pSR6730. From here, the promoter fragment was cloned into PstI and NcoI sites into pSR1, a pUC19-based plasmid harboring the GUS reporter gene, yielding pSR6721. A PstI/NcoI fragment containing the AtERDL6 promoter, the GUS reporter gene, and the Nos terminator was transferred to pGT7-bar (Becker et al., 1992), yielding construct pSR6722. Agrobacterium-mediated transformation of Arabidopsis (floral dip) produced 30 independent transgenic lines that were analyzed for GUS activity in the T1 and T2 generations.

**Subcellular Localization of AtERDL6**

To determine the compartment to which AtERDL6 is targeted within the cell, we cloned the AtERDL6 cDNA from pSR6740u into pC5Os5e, a pUC19-based plasmid harboring the GFP reporter gene (Aluri and Büttner, 2007), and AtERDL6-GFP expression was monitored by fluorescence microscopy. In addition, an AtERDL6-GFP fusion construct was transiently expressed in tobacco leaves and analyzed after 2 d by confocal laser scanning microscopy. To determine the cold stress response of AtERDL6, plants were kept for 1 week at 4°C. RNA was prepared from leaves harvested 5 h after the beginning of the light phase before (day 0, 22°C) and during (days 1, 2, 4, and 8) cold treatment.

**Generation of AtERDL6 and BolIMP Overexpression Mutants**

To generate an AtERDL6 overexpression construct, we used a modified version of the plant vector pCRIPT-bar (Becker et al., 1992). To this end, the EcoRI site was removed by restriction digestion following a nucleotide fill-in reaction and religation, yielding vector pRGl. An 814-bp HindIII/Xmal fragment from pSG35e (Aluri and Büttner, 2007) containing the enhanced CaMV 35S promoter was inserted into pRGl digested with HindIII and Xmal, yielding pRGl3. Restriction digestion of pRGl3 with BstXI and Sall yielded the new vector pRGl5. The AtERDL6 cDNA was PCR amplified using primers AtERD67c+271f_EcoRI (5’-AAATCAAGCCTGCTAAGAAAAGATGTTGCGAGGATGATAATGAGGAGGC-3’) and AtERD67c+1464r_Sall (5’-GGGATCCATGATATCAACAGAGTTAAGTCTC-3’) to generate the construct for subcellular localization, a Gateway LR reaction was performed according to the supplier’s instructions (MBI Fermentas). To this end, the ERDL6 open reading frame by RT-PCR, identifying BvIMP-for_U43629 and BvIMP-rev_HindIII (5’-TTAAGCTTTTATCTTGCAGTGGTGGATGATAATGAGGAGGC-3’) and BolIMP-rev_HindIII (5’-TTAAGCTTTTATCTTGCAGTGGTGGATGATAATGAGGAGGC-3’). The product was inserted into pBSK (digested with EcoRV), excised by using XhoI and HindIII, and further cloned into the correspondingly digested pMyosin vector. Thereafter, the construct was digested with SacI and cloned into the correspondingly treated pART27 vector. Transformation of Arabidopsis by Agrobacterium-mediated floral dip generated eight independent transgenic overexpression lines, of which lines 6 and 8 were identified as the strongest by northern-blot analysis.

**Plant Treatments for AtERDL6 Expression Analyses**

To analyze the effect of darkness and external sugars on AtERDL6 expression, plants were treated as described (Poschet et al., 2010). For heat stress experiments, plants were wrapped in aluminum foil to prevent dehydration and incubated for 6 h at 37°C (stress) or 23°C (control) in an incubator (Binder). To determine the impact of wounding, leaves were pinched with a syringe in a homogenous pattern. Leaves of unwounded plants as well as unwounded leaves from treated plants served as controls. Total RNA was extracted after 6 h. To determine the cold stress response of AtERDL6, plants were kept for 1 week at 4°C. RNA was prepared from leaves harvested 5 h after the beginning of the light phase before (day 0, 22°C) and during (days 1, 2, 4, and 8) cold treatment.

**RNA Isolation and Quantitative Time-PCR Analysis**

Total RNA was isolated as described (Schneideret al., 2003), and RT-PCR was performed according to the supplier’s instructions (MBI Fermentas). Primers AtERD67c+10680f (5’-TGGTCCATCCCAAATAGGAGAAAAG-3’) were used to PCR amplify AtERDL6-specific products from the indicated tissues. In parallel, primers AtLDL3-R (5’-TACTCTCAAGGCCAGAGTCAACAACA-3’) and AtDL3-R (5’-TTTCTGGTCCTCAACACCTTCACTC-3’) were used to amplify the LDL3 transcript as a reference (Czechowski et al., 2005). PCR conditions were 0.2 μL of each primer as indicated, and 2× Maxima SYBR Green/ROX qPCR Master Mix (Fermentas), and PCR amplification was performed in a RotorGene 2000 system (Corbett Research). Relative transcript abundance was determined using the comparative ΔΔCt method (Livak and Schmittgen, 2001) with the RotorGene software version 5.
kept in the dark. Leaves were harvested 0, 7, and 14 h after cold treatment and subjected to sugar analysis by ion chromatography.

Sugar Quantification by Ion Chromatography

Soluble sugars were extracted from plant material with 80% ethanol for 1 h at 80°C and quantified by high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) in an ICS-3000 system (Dionex) with a CarboPac PA1 column and 15 to 300 msi NaOH (Fluka) in HPLC-water (VWR) as eluent. Quantitative calculation of sugars was performed using Chromleon software 6.7 (Dionex).

Identification of T-DNA-Tagged Arabidopsis Knockout Lines for AtERDL6

Line SAIL_528_F09 (insertion at position +2,284, exon 12) was obtained from the Torrey Mesa Research Institute. Line SALK_106049 (insertion at position +3,107, exon 17) was obtained from the T-DNA mutant collection at the Salk Institute and provided by the Nottingham Arabidopsis Stock Centre (http://nasc.nott.ac.uk/). Plants homozygous for both AtERDL6 alleles were identified by PCR with genomic DNA and primers flanking the insertion site. In addition, the absence of ATERDL6 transcript in these knockout lines was verified by RT-PCR analyses with total RNA and ATERDL6-specific primers spanning the insertion site. AtERDL6c+1039f (5'-GAACTCAAACTAGGTTTCAAGGGATGA-3') and ATERDL6c+1462r_BspHI (5'-CGCGTCATGAATCTGAA-CAG-3') for the SAIL_528_F09 line and ATERDL6c+1039f (5'-CAGTCGCTGGCTTCTGCTT-3') and ATERDL6c+1512r (5'-CAGAGGGAGAGAGAGGGTGAC-3') for the SALK_106049 line.

Nonaqueous Fraction

Unshaded mature rosette leaves were harvested in the middle of the light period and used for subcellular fractionation according to the method described by Krueger et al. (2009). Compartment-specific marker enzymes were used to assign the different gradient fractions to the cytosolic, plastidic, or vacuolar compartment. Activity of the plastid-specific NADP-glyceraldehyde-3-phosphate dehydrogenase (EC 1.1.1.94) was measured according to Zrenner et al. (1993), and nitrate was used as a vacuolar marker as described by Winter et al. (1994; Cross et al., 2006). Five biological replicates each were done for wild-type or vacuolar compartment. Activity of the plastid-specific NADP-glyceraldehyde-3-phosphate dehydrogenase (EC 1.1.1.94) was determined as described by Zrenner et al. (1993), and nitrate was used as a vacuolar marker as described by Winter et al. (1994; Cross et al., 2006). Five biological replicates each were done for wild-type and mutant lines. For each biological replicate, data from two technical replicates of marker measurements and from three technical replicates of sugar quantifications via HPAEC-PAD were averaged.

Seed Germination Assay and Quantification of Seed Storage Compounds

For seed germination assays, seeds of both ATERDL6 mutant lines and the wild type were surface sterilized, and germination was carried out on MS plates containing 2% Glc, 7.2% Glc, 2% Suc, 2% sorbitol, or no sugar. Germination was determined by measuring the time of radicle emergence from triplicate assays with 80 to 140 seeds per plate...

Protein and lipid content of seeds was determined as described (Wingenter et al., 1994; Cross et al., 2006). Five biological replicates each were done for wild-type and mutant lines. For each biological replicate, data from two technical replicates of marker measurements and from three technical replicates of sugar quantifications via HPAEC-PAD were averaged.

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Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. Phylogenetic tree of Arabidopsis MSTs and plant ERDL6 orthologs.

Supplemental Figure S2. Tissue- and organ-specific expression patterns of ATERDL6 (At1g75220) according to the Genevestigator microarray database.

Supplemental Figure S3. Genomic organization of the ATERDL6 gene and molecular characterization of two independent T-DNA insertion lines for ATERDL6.

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Supplemental Figure 1: Phylogenetic tree of *Arabidopsis* MSTs and plant ERDL6 orthologs.

Currently available *ERDL6* orthologous sequences as well as all members of the Arabidopsis MST-like gene family were collected to build an un-rooted phylogenetic tree using the neighbour-joining method of the MEGA 4.1 software (Tamura K, Dudley J, Nei M & Kumar S (2007) MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0. Molecular Biology and Evolution 24:1596-1599). A 1000 trial of bootstrap analysis was used to provide confident estimates for phylogenetic tree topology and corresponding bootstrap values are given in red at the basis of each branch. The scale bar corresponds to 0.1 estimated amino acid substitutions per site.
Supplemental Figure 2: \textit{AtERDL6} (At1g75220) expression pattern according to microarray data (www.genevestigator.com). Error bars represent standard deviation of a varying number n of experiments as indicated in brackets. Note the different scale applied for the high expression levels found in protoplasts and stamen/pollen (grey bars).
Supplemental Figure 3: Genomic organization of the *AtERDL6* gene and molecular characterization of two independent T-DNA insertion lines for *AtERDL6*.

Diagram of the *AtERDL6* gene (At1g75220) and its primary transcript. Exons (black boxes) and introns (lines) are indicated. 5’ and 3’ untranslated regions are depicted in red. Additionally, the positions of the two T-DNA insertions in mutant lines SALK_106049 (+3107; exon 17) and SAIL_528_F09 (+2284; exon 12) are shown. As indicated above the insertion sites, the absence of *AtERDL6* specific transcript in both mutant lines was demonstrated by RT-PCR analyses using *AtERDL6*-specific primers along with *Actin2*-specific primers for cDNA quality control (see Material section for primer details).