Late embryogenesis abundant proteins protect human hepatoma cells during acute desiccation

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Expression of late embryogenesis abundant (LEA) proteins is highly correlated with desiccation tolerance in anhydrobiotic animals, selected land plants, and bacteria. Genes encoding two LEA proteins, one localized to the cytoplasm/nucleus (AfrLEA2) and one targeted to mitochondria (AfrLEA3m), were stably transfected into human HepG2 cells. A trehalose transporter was used for intracellular loading of this disaccharide. Cells were rapidly and uniformly desiccated to low water content (≤0.12 g H2O/g dry weight) with a recently developed spin-drying technique. Immediately on rehydration, control cells without LEA proteins or trehalose exhibited 0% membrane integrity, compared with 98% in cells loaded with trehalose and expressing AfrLEA2 or AfrLEA3m; surprisingly, AfrLEA3m without trehalose conferred 94% protection. Cell proliferation across 7 d showed an 18-fold increase for cells dried with AfrLEA3m and trehalose, compared with 27-fold for nondried controls. LEA proteins dramatically enhance desiccation tolerance in mammalian cells and offer the opportunity for engineering biostability in the dried state.

It is well established that water is essential for active life (1, 2), yet animals from four different phyla, selected land plants, and certain fungi and bacteria survive severe desiccation for extended periods (3). The mechanisms by which animals protect cellular structure and function in the state of anhydrobiosis, or “life without water” (4–6), are not only of fundamental interest, but also of potential biomedical importance for cell stabilization (7–9). Multiple molecular components may contribute to the intracellular conditions required for successful desiccation in nature, and are often associated with metabolic preconditioning for embryonic stages (10). Many organisms accumulate compatible osmolytes of low molecular weight for osmotic balance during water stress, biopreservation and intrinsically disordered proteins. Osmolytes such as trehalose, halose. Here we report the profound protection conferred during drying, including action as a “molecular shield” to sterically reduce aggregation of denatured proteins, stabilization of target proteins via chaperone-like activity, protection of cell membranes (particularly for mitochondria-targeted forms; refs. 27, 28), stabilization of vitrified sugar glasses by increasing the glass-transition temperature (Tg), and sequestration of divalent ions (15, 16). Finally, synergistic interactions have been documented for LEA proteins and sugars such as trehalose, related to their ability to protect target proteins during drying (29).

For the purpose of the present study, we chose to focus on two LEA proteins naturally expressed in embryos of the brine shrimp Artemia franciscana, AfrLEA2 and AfrLEA3m. Several LEA proteins are expressed in these embryos, including group 1 and group 3 LEA proteins (25, 26, 30, 31). Bioinformatic analyses have identified AfrLEA2 in the cytoplasm (25) and AfrLEA3m in mitochondria (26). Expression in HepG2/C3A cells of a chimeric protein composed of the leader sequence from AfrLEA3m and GFP has shown targeting of the construct to mitochondria (26). We developed a HepG2 cell line based on the tetracycline (Tet)-inducible expression system and then stably transfected this line with either AfrLEA2 or AfrLEA3m under doxycycline (Dox) control. To introduce trehalose into the cytoplasm, we also stably transfected these cells with trehalose transporter 1 (TRET1) from the African chironomid Polypedilum vanderplanki (32) that is expressed constitutively.

To avoid problems commonly encountered with evaporative drying of sessile droplets containing suspended cells, we have developed a spin-drying technique that yields rapid and uniform drying across the sample and low final water contents (33, 34). Droplet drying is inherently slow, and water removal is spatially nonuniform, because a glassy skin forms at the sample interface when solutions of glass-forming solutes such as trehalose are dried (35, 36). This phenomenon slows water removal and prevents desiccation beyond certain limits. The negative biological impact is that cells can be trapped in a partially desiccated and nonvitrified sample and consequently subjected to prolonged hypoxic stress.

In this work, we merged biological and physicochemical strategies in our attempt to enhance the desiccation tolerance of mammalian cells. Specifically, we applied principles gleaned from animals whose evolutionary history has provided the capacity for severe desiccation tolerance (e.g., by means of LEA proteins and protective glass-forming solutes) and combined these with an improved drying method based on thermodynamic and kinetic studies of sugar solutions and relevant thermophysical properties.


The authors declare no conflict of interest.

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of cells. We report remarkable improvement in the tolerance of mammalian cells to acute desiccation, arguably at the lowest moisture levels yet attained for uniformly dried cells.

Results

HepG2 Tet-On Cell Lines and Inducible Expression of AfrLEA2 and AfrLEA3m. A Tet-inducible gene expression system in HepG2 cells was developed by stable transfection with a vector encoding the Tet transactivator protein. To verify whether these HepG2 Tet-On cells are competent for efficient expression of the target protein under Dox control, the luciferase gene was transiently transfected into the cells in an appropriate expression vector. Western blot analysis showed strong induction of luciferase protein across the range of 10–1,000 ng/mL Dox (Fig. 1A), with undetectable expression in the uninduced control. A similar pattern was observed when luciferase enzyme activity was plotted as a function of increasing Dox concentrations (Fig. 1B). The induced activities at 10–1,000 ng/mL Dox were significantly higher (P < 0.001) compared with the uninduced control, with the maximum induction at 1,000 ng/mL being at least 500-fold above control. Thus, the Tet-On expression system operates efficiently in HepG2 cells.

The original nucleic acid sequences for Afrlea2 and Afrlea3m cloned from A. fransciscana embryos (25, 26) were first optimized for human codon bias, after which each gene was inserted into a Tet-On expression vector (pTRE3G) and stably transfected into HepG2 Tet-On cells. Fig. 1C shows a representative Western blot for AfrLEA2 expression induced with 1,000 ng/mL Dox across a 5-d induction period. Polyclonal antibody raised against recombinant AfrLEA2 exhibited a strong band induced at ~45 kDa that matched migration of the purified recombinant protein. Quantification of data averaged from three individual experiments (Fig. 1D) indicates that AfrLEA2 induction was significant at day 1 and continued to increase across the 5-d induction period, with maximum induction at ~11-fold above the uninduced control by 120 h. Western blot analysis for AfrLEA3m expression during a 5-d time course of induction is shown in Fig. 1E. The polyclonal antibody against the recombinant protein recognized a band for AfrLEA3m at ~34 kDa, consistent with the predicted mass of the mature protein (26) after removal of the mitochondrial targeting sequence (30.9 kDa), allowing for the fact that intrinsically disordered proteins show an increased apparent molecular mass on SDS/PAGE gels owing to reduced binding of SDS (37). Quantification of AfrLEA3m indicates statistically significant induction across the 5-d period (Fig. 1F). The maximum cellular expression (~20-fold over the uninduced control) occurred at 24 h and declined gradually thereafter. It should be noted that migration and incorporation into the mitochondrial compartment may take longer than 24 h, based on localization of a chimeric protein composed of the AfrLEA3m leader sequence plus GFP (26). Thus, the Tet-On system effectively induces expression of AfrLEA2 and AfrLEA3m in human HepG2 cells. Our ability to document acceptable stability of these LEA proteins in the transfected cells is important, because the uncoiled conformation of intrinsically disordered proteins in aqueous solution can render them susceptible to proteolytic degradation, and, accordingly, they may exhibit reduced half-lives (18).

Subcellular Localization of AfrLEA2 and AfrLEA3m. The subcellular localization of both AfrLEA2 and AfrLEA3m proteins in HepG2 Tet-On cells was documented using confocal microscopy. For AfrLEA2, transient transfection was done with a plasmid encoding a chimeric protein composed of AfrLEA2 plus GFP. At 24 h after transfection, AfrLEA2-GFP is localized to the cytoplasm of HepG2 cells (Fig. 2A). Lack of colocalization in the merged image detected by staining with MitoTracker Red (Life Technologies) does not support any mitochondrial targeting for AfrLEA2. This cytoplasmic distribution for AfrLEA2 is consistent with predictions from bioinformatic analysis (25). At 3 d after transfection of HepG2 Tet-On cells, a fraction of the AfrLEA2-GFP was also detected in the nucleus (Fig. S1). Small proteins tagged with GFP are known to readily enter the nucleus without the requirement for a nuclear localization signal (38); apparently this is the case for AfrLEA2-GFP, given that bioinformatic algorithms failed to detect a nuclear localization signal (25).

Immunofluorescence staining with primary AfrLEA3m antibody, followed by secondary antibody conjugated to FITC, was used to colocalize AfrLEA3m. Images for both induced and uninduced HepG2 Tet-On-AfrLEA3m cells were captured with confocal microscopy. Fig. 2B definitively shows that AfrLEA3m is targeted to the mitochondrial network of HepG2 cells, based on colocalization with MitoTracker Red. These results provide direct evidence of mitochondrial localization of AfrLEA3m using the full-length protein. Thus, we should be able to evaluate the impact of differentially targeted LEA proteins on the protection of HepG2 cells during desiccation.

Cellular Transfusion with TRET1 and Kinetics of Sugar Uptake. In some cases (6, 39), but certainly not all (40–42), animals with natural desiccation tolerance accumulate low molecular weight solutes, such as trehalose, along with protective proteins. Trehalose is a nonreducing sugar with a high Tg value, and consequently it vitrifies readily at biologically relevant temperatures. Vitrification of the intracellular compartment is considered one factor contributing to dehydration tolerance during anhydrobiosis (43). Although trehalose is not permeable to biological membranes, its effectiveness as a stabilizer is greatest when present on both sides of a lipid bilayer (44). Thus, we used a naturally occurring membrane channel TRET1 to load the sugar into HepG2 cells. A TRET1 construct designed for constitutive expression was stably transfected into HepG2 Tet-On cells with and without the capacity for inducible expression of AfrLEA2/AfrLEA3m. Evaluation of TRET1 mRNA expression by RT-PCR revealed expression in TRET1-transfected cells, but not in control cells (Fig. 3A). Next, the functionality of the transporter protein was 20% increased by measuring trehalose uptake into TRET1-transfected cells and nontransfected control cells that had been preincubated with 50 mM trehalose for up to 24 h (Fig. 3B). HPLC analyses indicated that HepG2-TRET1 cells contained an average of 0.0217 pmol trehalose per cell after 24 h of loading, approximately sevenfold higher than the average level in WT HepG2 cells under the same conditions. The intracellular concentration of trehalose in the HepG2-TRET1 cells was estimated as ~20 mM. No significant change in cell morphology due to the presence of intracellular trehalose was observed in HepG2 or HepG2-TRET1 cells before spin-drying.

Biotubalization of HepG2 Cells During Severe Desiccation. Fluid movements during spin-drying of cells in a glass-forming solution of trehalose are depicted in Fig. 3C. Control HepG2 cells (without LEA protein or intracellular trehalose) exhibited 0% membrane integrity (n = 9) when spin-dried to a final water content of <0.12 g H2O/g dry weight, as confirmed by FTIR microscopy and bulk gravimetric analysis of the water content (Fig. 4A). Membrane integrity after spin-drying and immediate rehydration was based on staining of live (green) and dead (red) cells with Syto-13 and ethidium bromide (Fig. 4B). The membrane integrity of HepG2-TRET1 cells preloaded with trehalose increased to 44.5 ± 22.2% (mean ± SD; n = 3). In contrast, the membrane integrity of HepG2-AfrLEA2 cells without trehalose was 57.2 ± 13.0% (n = 9), increasing to 98.3 ± 22.2% (n = 9) when these cells were preloaded with trehalose. The most remarkable result is the striking integrity of 93.6 ± 4.6% (n = 9) measured for HepG2-AfrLEA3m cells in the absence of trehalose. Consistent with this finding, evidence indicates that model synthetic peptides composed of several tandem motifs of group 3 LEA proteins (22- to 44-aa residues) vitrify at a high Tg, which suggests that dried LEA proteins themselves may vitrify and act in this way as excellent protectants against desiccation damage (45). It is appropriate to emphasize that although AfrLEA3m is indeed mitochondrial-targeted, synthesis occurs in the cytosol, and importation into mitochondria is time-dependent (26), and thus presumably some...
fraction of the protein is continuously present in the cytoplasm. In the absence of trehalose, membrane integrity was 0% (n = 3) for both uninduced HepG2-AfrLEA2 and uninduced HepG2-AfrLEA3m cells. The integrity of HepG2-AfrLEA3m cells preloaded with trehalose increased slightly, to 97.7 ± 3.8% (n = 9). Thus, both cellular treatments containing LEA–sugar mixtures display ~98% membrane integrity after severe desiccation. In vitro studies have shown that mixing LEA-like peptides into trehalose solutions stabilizes the vitrified sugar glasses on drying, as judged by upward shifts in $T_g$ (45).

Fig. 1. Construction of HepG2 Tet-On cell lines and the induction of transfected proteins. (A) Representative Western blot of luciferase enzyme expression in the constructed HepG2 Tet-On cell line at 24 h after induction with Dox. α-tubulin served as the loading control. (B) Luciferase catalytic activity measurements (mean ± SEM; n = 8) versus Dox addition. "a" indicates significant differences ($P < 0.001$) vs. the uninduced control. Both luciferase expression and activity were strongly induced by Dox in a dose-dependent manner. (C) Induction with Dox (1,000 ng/mL for the 120-h period) of AfrLEA2 in HepG2 Tet-On cells as shown by Western blot analysis. (D) AfrLEA2 expression from three individual experiments (mean ± SEM) normalized to α-tubulin. Densitometric quantification was performed using Bio-Rad Quantity One 1-D software. Data are presented as fold increase relative to the uninduced control (hour 0). "a" indicates significant differences ($P < 0.005$) vs. the uninduced control, and "b" indicates significance ($P < 0.005$) vs. the 24-h time point. (E) Western blot analysis of AfrLEA3m induction in HepG2 Tet-On cells with 1,000 ng/mL Dox present throughout the 120-h period. (F) AfrLEA3m expression (mean ± SEM; n = 3) normalized to α-tubulin. Quantification and statistical tests are as described above.
is a primary contributor to this lyoprotection of SD (www.pnas.org/cgi/doi/10.1073/pnas.1214893109). When 0.21). In contrast, the growth kinetics ≤ | Li et al. |

TRET1 transporter, trehalose uptake kinetics, and the spin-drying 0.05). Plant seeds are known to Subcellular localization of AfrLEA2 and AfrLEA3m HepG2 Tet-On cells. (A) Cells were imaged with confocal laser scanning microscopy performed at 24 h after transient transfection with AfrLEA2-GFP. (Left) Distribution of green fluorescence from AfrLEA2-GFP, with nuclei clearly visible as the absence of green fluorescence. (Center) Distribution of mitochondria based on MitoTracker Red staining. (Right) Merged image, confirming the cytoplasmic localization of AfrLEA2-GFP as predicted from bioinformatic data. (B) AfrLEA3m was stably transfected into HepG2 Tet-On cells. The top row presents images obtained from cells induced with Dox for 2 d, and the bottom row shows images for the uninduced control cells. (Left) Immunofluorescent staining with specific AfrLEA3m primary antibody (chicken IgY), followed by secondary antibody (goat anti-chick IgY) tagged with FITC. (Center) Mitochondrial distribution imaged with MitoTracker Red staining. (Right) Merged images clearly showing the yellow color resulting from overlap of the green AfrLEA3m staining with the MitoTracker Red in induced cells, which confirms the mitochondrial localization of AfrLEA3m in the HepG2 cell line.

Growth studies conducted across ensuing days after rehydration of dried cells revealed the greatest proliferation for those containing AfrLEA3m plus trehalose compared with all other treatment groups (Fig. 4C). By day 7, numbers were increased 18-fold for cells with AfrLEA3m plus trehalose vs. 27-fold for nondried controls during the same growth period (statistically equivalent; P = 0.21). In contrast, the growth kinetics for cells expressing AfrLEA2 (with or without trehalose) and AfrLEA3m (without trehalose) were markedly slower compared with nondried controls (P ≤ 0.05). Plant seeds are known to contain mitochondrial-targeted LEA protein (46). Macherel et al. (47) have argued persuasively that mitochondria play a profound and vital role in desiccation tolerance, and that their protection in the dry state and rapid return to full respiratory activity on rehydration is a matter of life and death in plant cells. The superior protection against desiccation stresses afforded HepG2 cells by AfrLEA3m is consistent with the importance of intact and functional mitochondria during the recovery process.

Discussion

We have successfully applied lessons learned from organisms that are naturally desiccation-tolerant to the protection of a mammalian cell line from very low water activity during acute drying. The significantly improved viability of human HepG2 cells obtained after desiccation below 0.12 g H2O/g dry weight is a very encouraging step toward the ultimate goal of preparing dried cells for storage at ambient temperature. Ectopic expression of LEA proteins cloned from the anhydrobiotic crustacean A. franciscana is a primary contributor to this lyoprotection of human HepG2 cells. LEA proteins are not represented in mammalian genomes, although a short protein motif of 10 amino acids that resembles a LEA protein has been reported in humans (48); however, regions of low sequence complexity characteristic of LEA proteins complicate the assessment of relatedness.

Previous work on air drying of mammalian cells has generally resulted in poor viability at water content below ~0.35 g H2O/g dry weight obtained by sessile droplet drying (49, 50). A nanoliter droplet approach also yielded low viability (<10%) at a residual water content of ~0.1 g H2O/g dry weight (51). Successful air drying to the point of no residual water has been reported for

Fig. 2. Subcellular localization of AfrLEA2 and AfrLEA3m HepG2 Tet-On cells. (A) Cells were imaged with confocal laser scanning microscopy performed at 24 h after transient transfection with AfrLEA2-GFP. (Left) Distribution of green fluorescence from AfrLEA2-GFP, with nuclei clearly visible as the absence of green fluorescence. (Center) Distribution of mitochondria based on MitoTracker Red staining. (Right) Merged image, confirming the cytoplasmic localization of AfrLEA2-GFP as predicted from bioinformatic data. (B) AfrLEA3m was stably transfected into HepG2 Tet-On cells. The top row presents images obtained from cells induced with Dox for 2 d, and the bottom row shows images for the uninduced control cells. (Left) Immunofluorescent staining with specific AfrLEA3m primary antibody (chicken IgY), followed by secondary antibody (goat anti-chick IgY) tagged with FITC. (Center) Mitochondrial distribution imaged with MitoTracker Red staining. (Right) Merged images clearly showing the yellow color resulting from overlap of the green AfrLEA3m staining with the MitoTracker Red in induced cells, which confirms the mitochondrial localization of AfrLEA3m in the HepG2 cell line.

Fig. 3. TRET1 transporter, trehalose uptake kinetics, and the spin-drying approach. TRET1 is expressed constitutively and is not under Dox control. (A) RT-PCR showing the presence of mRNA for TRET1 in HepG2 Tet-On-TRET1 cells and its absence in HepG2 Tet-On control cells. The TRET1 plasmid was used as template for a positive control (P). A negative control with no added template (N) is shown as well. Actin was used for cell samples as a loading control. (B) Trehalose loading kinetics for HepG2 Tet-On-TRET1 cells and control HepG2 Tet-On cells (without TRET1). Cells were incubated in 50 mM trehalose for the indicated times. Trehalose uptake was more pronounced in the presence of TRET1. Values are mean ± SD (n = 3). When error bars are not visible, they are within the size of the symbol. (C) Schematic of the spin-drying process, in three steps: (1) At the beginning of the spin-drying process, the excess liquid is removed from the sample by radial flow (E) due to spinning (ω); (2) after the excess liquid is removed by radial flow, the evaporation (E) into an environment continuously purged with dry nitrogen gas causes further drying of the samples; (3) in the final stage of the drying process, a thin film of dried trehalose with very low moisture content is formed.
human primary fibroblasts provisioned with intracellular trehalose (52), but a subsequent report from the same group indicated that those water content analyses apparently were inaccurate (53). A group 3 LEA protein from the nematode *Aphelenchus avenae* transfected into the human cell line T-REx293 (derived from the embryonic kidney cell line HEK293) improved the tolerance to hyperosmotic shock induced by exposure to 100 mM NaCl or 400 mM sugars, but exposure of cells to air drying over periods of hours at different relative humidity levels resulted in mortality (54). In the present study, the protection of HepG2 cells during drying provided by AfrLEA2 versus AfrLEA3m was statistically identical in the presence of intracellular trehalose, as judged by membrane integrity immediately after rehydration (98.3% vs. 97.7%). Without intracellular trehalose, AfrLEA2 was significantly less effective than AfrLEA3m. This difference was particularly evident in growth studies extended over 7 d, in which proliferation was far lower for cells with AfrLEA2 compared with cells with AfrLEA3m, with or without intracellular trehalose. The reasons for the improved performance of HepG2 cells that expressed AfrLEA3m remain unclear, but this finding points to the critical role in protecting mitochondrial integrity during drying. Insight into this issue has been provided by studies of an LEA protein from pea seed mitochondria (PsLEAm) localized to the matrix (46). Using differential scanning calorimetry and FTIR spectroscopy, Macherel and coworkers (27, 28) found that PsLEAm interacts with liposomes in the dry state. The protein protects liposomes during drying by preventing membrane fusion and lysis. Furthermore, protein modeling studies by that group showed that the amphiphatic α-helices of PsLEAm formed during drying take on an interesting pattern in which negative amino acid residues form a stripe bordered on either side by positive residue stripes. This pattern may facilitate the protein’s integration with the lipid membrane parallel to its plane (28). Evidence also shows that a LEA protein from a bdelloid rotifer (ArLEA1B) interacts with dried liposomes and decreases the gel-to-liquid crystalline phase-transition temperature ($T_{m}$) of dry liposomes (55). Finally, it is noteworthy that mitochondria isolated from embryos of the anhydrobiotic species *A. franciscana* contain at least one LEA protein (AfrLEA3m), and likely more (16, 30), targeted to the mitochondrion. When these mitochondria are subjected to water stress represented by freezing in a trehalose solution, respiratory control ratio (succinate plus rotenone) for mitochondria isolated from frozen/thawed mammalian cells were remarkably well-preserved compared with ratios for control, nonfrozen mitochondria (26). Taken together, these findings add to the mounting evidence that mitochondrial-targeted LEA proteins offer strong protection to organelles against desiccation-induced damage, and this function is important for improving the desiccation tolerance of mammalian cells.

Spin-drying is another key step in the development of an improved dry-processing technique for mammalian cells (34), primarily because of the high degree of spatial uniformity in moisture content obtained in the samples and the rapid drying of cells. To our knowledge, both the short-term viability and the long-term viabilities obtained after immediate rehydration in this study are the highest yet reported for mammalian cells dried uniformly to a water content below 0.12 g H$_2$O/g dry weight. As we have discussed previously in more detail (34), traditional drying procedures result in substantial spatial nonuniformity, such that regions with higher residual water are prone to higher rates of chemical reactions that can foster unwanted metabolic processes, energy depletion, and cell degrada-

Fig. 4. Membrane integrity and long-term growth of HepG2 cells after spin-drying and immediate rehydration. (A) HepG2 cells with TRET1 transporters were incubated in 50 mM trehalose-containing medium for 18 h before spin-drying. Membrane integrity was determined using Syto-13 and ethidium bromide viability dyes. Values are mean ± SD (n = 3–9 independent determinations). "a" indicates statistically significant differences ($P < 0.05$) vs. control (no trehalose or LEA protein), and "b" indicates statistically significant differences ($P < 0.05$) vs. cells with intracellular trehalose alone. (B) Fluorescent micrograph of HepG2 cells stained with Syto-13 and ethidium bromide after spin-drying and rehydration (green, live cells; red, dead cells). (Scale bars: 100 μm.) (C) Cells were rehydrated with fully complemented cell culture medium and incubated in a 5% (vol/vol) CO$_2$ environment at 37 °C. Cell counts on days 1, 3, and 7 were obtained with trypan blue exclusion for parallel samples. The error bars indicate SEM. By day 7, cell numbers were increased by 18-fold for those with AfrLEA3m plus trehalose, vs. 27-fold for nondried controls during the same growth period (statistically equivalent; $P = 0.21$). In contrast, the growth kinetics for cells expressing AfrLEA2 (with or without trehalose) and AfrLEA3m (without trehalose) were markedly different compared with the nondried controls ($P < 0.05$), as indicated by asterisks.

The improved dry-processing technique for mammalian cells (34), traditional drying procedures result in substantial spatial nonuniformity, such that regions with higher residual water are prone to higher rates of chemical reactions that can foster unwanted metabolic processes, energy depletion, and cell degra-

In summary, LEA proteins dramatically improve the surviv-

orship of human cells during acute desiccation to low water conditions.
activity achieved by spin-drying. Spin-drying represents a significant step forward in the processing procedures used for water removal. Adding intracellular trehalose in combination with LEA proteins enhances the long-term growth of cells, as measured across 7 d after rehydration, to proliferation rates approaching those of control cells that were never dried. Successful strategies for long-term storage of dried cells at room temperature may require metabolic preconditioning signals (10), elevated $T_d$ values, and the simultaneous expression of several LEA proteins targeted to various intracellular compartments. In desiccation-tolerant plant tissues, LEA proteins are localized to the nucleus, mitochondrion, chloroplast, endoplasmic reticulum, vacuole, peroxisome, and plasma membrane (16).

Materials and Methods

Detailed information on materials, cell culture, construction of the HepG2 Tet-On cell line, gene transfection, selection of stable cell lines, Western blot analysis, confocal imaging and immunofluorescence staining, trehalose uptake kinetics, spin-drying of mammalian cells, and membrane integrity and long-term viability is provided in SI Materials and Methods.

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Supporting Information

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**SI Materials and Methods**

**Materials.** The plasmids (pCMV-Tet3G, pTRE3G, pTRE3G-Luc, pEGFP-C3, and a linear hygromycin marker), Xfect transfection system, tetracycline (Tet) system-approved FBS, and doxycycline (Dox) were purchased from Clontech. PCR primers were synthesized by Integrated DNA Technologies. BamHI and SalI restriction enzymes and Phusion Hot Start DNA polymerase were purchased from New England BioLabs. The Endofree Plasmid Mini Kit and Maxi Kit were obtained from Qiagen. Tissue culture reagents, including MEM, FBS, glutamine, penicillin, and streptomycin, were obtained from Life Technologies. G 418 sulfate was purchased from Calbiochem. Tubulin antibody was purchased from Cell Signaling Technology. Antibodies against late embryogenesis abundant (LEA) proteins AfrLEA2 and AfrLEA3 were raised against the expressed and purified proteins by Aves Labs. Trehalose was obtained from Ferro Pfanstiehl. All other reagents were purchased from Sigma-Aldrich.

**Cell Culture.** Human hepatocellular carcinoma (HepG2) cells were obtained from American Type Culture Collection (catalog no. HB-8065) and cultured in MEM containing 10% FBS (vol/vol), 2 mM glutamine, 100 U/mL penicillin, and 100 μg/mL streptomycin. Cells were maintained in 25-cm² T flasks at 37 °C and equilibrated in a humidified atmosphere with 5% CO₂ and 95% air (vol/vol). For subculture, cells were detached with 0.25% trypsin–EDTA solution, and 1 × 10⁵ cells were seeded into new flasks. Cells were counted with a hemocytometer when required during experimental procedures.

**Construction of HepG2 Tet-On Cell Line.** The regulatory plasmid pCMV encoding a Tet transactivator with a neomycin expression gene was transfected into HepG2 cells. In brief, 5 × 10⁶ HepG2 cells were seeded into single wells of six-well plates, and Clontech Xfct transfection reagent and 2.0 μg of the regulatory plasmid were added into each well containing HepG2 cells following the manufacturer’s protocol. After a 24-h incubation, the cells were transiently transfected with pTRE3G encoding with the luciferase reporter gene to test the inducibility of HepG2 Tet-On cells. Selection for stable cell lines was initiated by adding 800 μg/mL G418 into Tet-On culture medium (HepG2 culture medium with Tet-On-approved FBS added instead of the normal FBS). After an approximate 4 wk selection period, single clones appeared, which were cultured and resteted for maximum inducibility with the luciferase reporter.

To test the HepG2 Tet-On 3G cells for induction efficiency, Dox, a synthetic Tet derivative, was used to promote the inducible binding of transactivator protein to pTRE3G promoter. The plasmid pTRE3G-Luc, which contains the Tet-responsive element (TRE) and luciferase reporter gene (Luc), was transiently transfected into HepG2 Tet-On 3G cells. Various amounts of Dox (0, 1, 10, 100, and 1,000 ng/mL) were added, after which luciferase activity and enzyme expression were analyzed. Luciferase activity was evaluated using the Promega Bright-Glo Luciferase assay system and a fluorescence microplate reader. Luciferase protein expression was detected by Western blot analysis using the same dosages for Dox induction as above.

**Gene Transfections and Selection of Stable Cell Lines.** To express Afrle2 (−1.1 kb; GenBank accession no. EU477187) and Afrlea3m (−0.9 kb; GenBank accession no. FJ592175) under the control of the Tet-On–inducible expression system, and to obtain maximal expression in HepG2 cells, the full-length sequences (1, 2) were revised to reflect human codon bias and synthesized using Gene Oracle software. SalI and BamHI, used to insert the LEA genes into the Tet-On expression vector pTRE3G, were added at 5' and 3' of each translational sequence. A Kozak sequence was added as well. AfrLea2 and AfrLea3m sequences were then inserted into the precut pTRE3G vector and ligated with T4 ligase. The resulting plasmid was transformed into Escherichia coli cells for amplification. The construct was verified by double digestion of the plasmid with XhoI and EcoR. HepG2 Tet-On cells were cotransfected with the Tet-On expression plasmid and a linear hygromycin marker (20:1) following the procedures described above. After 48 h, Tet-On medium containing 200 μg/mL hygromycin was added to select individual clones that appeared after approximately 3 wk. Single clones were then picked for evaluation of AfrLEA2 and AfrLEA3m protein expression by Western blot analysis.

TRET1 was cloned from anhydrobiotic larvae of *Polypedilum vanderplanki* (3) and the original vector (a gift from Takahiro Kikawada, National Institute of Agrobiological Sciences, Ibaraki, Japan). The TRET1 gene was then inserted into the expression vector (pcDNA 6.2/V5-DEST Gateway Vector; Invitrogen), and its sequence was verified by presequencing PCR with T7 promoter and V5 reverse primers, followed by sequencing of the cleaned product. Note that the Tet-On vector pTRE3G was not used for this gene, so that TRET1 expression in HepG2 Tet-On cells was constitutive. Transfection into HepG2 Tet-On cell lines (with or without the capacity for AfrLEA2/AfrLEA3m expression) was done using GenePORTER transfection reagent (Genlantis) according to the manufacturer’s instructions. Stable cell lines were selected with 5 μg/mL blasticidin for 5–7 d and then maintained in Tet-On medium with 2 μg/mL blasticidin. RT-PCR was used to verify TRET1 expression in the HepG2 cells. Total RNA was extracted using a RNaseasy Mini Kit (Qiagen) and reverse-transcribed to cDNA using a DynaNo cDNA Synthesis Kit (Thermo Scientific). The forward PCR primer was 5'-GGGTTTGGCAACATCGGATCCCTCAT-3' and the reverse primer was 5'-AACTTTGGCTCCACGCGCATCTAT-3'. The PCR used Crimson Taq DNA polymerase (New England BioLabs) and 25 cycles of 95 °C for 30 s, 60 °C for 30 s, and 68 °C for 60 s, followed by a final extension at 72 °C for 5 min. Human β-actin was used as the loading control, and the primers were 5'-GGCCACCCAGCAAATGAGATCAA-3' (forward) and 5'-ACTCTGATACCTCGTTGCTGA-3' (reverse).

**Western Blot Analysis.** Luciferase and AfrLEA2/AfrLEA3m protein expression were evaluated by Western blot analysis. Transfected cells were harvested with SDS sample buffer containing 2% (wt/vol) SDS, 25% (vol/vol) glycerol, 5% (vol/vol) β-mercaptoethanol, and 0.625 M Tris-HCl (pH 6.8). The protein concentration was determined using Pierce 660-nm protein assay reagent with ionic detergent compatibility reagent (Thermo Scientific) using BSA as the standard. Equal amounts of protein from the whole-cell lysates were electrophoresed on 10% (wt/vol) SDS/PAGE gels, and the proteins were electrotransferred to nitrocellulose membranes. Membranes were incubated with blocking solution containing 5% (wt/vol) nonfat milk prepared in Tris-buffered saline [50 mM Tris base, 150 mM NaCl, 0.05% Tween-20 (pH 7.6) with HCl], and then incubated with the primary antibodies for AfrLEA2 and AfrLEA3m (brine shrimp anti-chick IgY, 1:100,000; Aves Labs, Inc.) and for luciferase (firefly antirabbit IgG fraction, 1:1,000 dilution; Sigma-Aldrich) at 4 °C overnight. After three 5-min rinses with Tris-buffered saline,
the membrane was incubated with the secondary antibodies (1:10,000, rabbit anti-IgY for AfrLEA2/AfrLEA5m; 1:10,000, anti-rabbit IgG for luciferase) conjugated with HRP at room temperature for 1 h. The LumiGlo developing system and high-performance chemiluminescence film (GE Healthcare) were used to develop blots.

Confocal Imaging of AfrLEA2-GFP and Immunofluorescent Staining of AfrLEA2m. For transient transfection with AfrLEA2-GFP, HepG2 Tet-On cells were detached from culture flasks using 0.25% trypsin-EDTA solution and replated into six-well glass-bottom plates coated with type 1 rat tail collagen. When cells reached 60% confluency, GenePORTER transfection reagent was used to transfect the cells with the expression vector encoding for the AfrLEA2-GFP protein (Clontech pEGFP-C3). Before confocal visualization, cells were stained with 100 nM MitoTracker Red (Life Technologies) for 20 min and then washed three times with PBS. Cellular localization of the chimeric protein was visualized by confocal imaging performed daily across a 3-d period after transfection. Images were acquired with a Leica Microsystems TCS SP2 microscope equipped with a Plan Apo 63× oil immersion lens. Two lasers, 488 nm at 15% power and 543 nm at 25% power, were used to sequentially excite GFP and MitoTracker Red, respectively. Emission passes of 500–535 nm and 560–650 nm were used to detect the signals.

HepG2 Tet-On cells stably transfected with Afrlea3m were induced with Dox for 3 d, and uninduced cells were parallel. Live cells were stained with MitoTracker Red as above, then fixed in 0.4% paraformaldehyde for 20 min at room temperature, washed with PBS, blocked with 5% (vol/vol) normal goat serum and 0.3% Triton X-100 in PBS for 1 h, and incubated with anti-AfrLEA3m chick IgY at 1:200 dilution in 5% normal goat serum and PBS overnight. Control cells without primary antibody were prepared as well. After another washing in PBS, cells were incubated in secondary antibody (goat anti-chick IgY conjugated with FITC; excitation, 488 nm; emission, 515 nm; Jackson ImmunoResearch) at a 1:200 dilution in 5% normal goat serum in PBS for 1 h. Cells were then washed three times in PBS, treated with ProLong Gold antifade solution (Life Technologies), and imaged with confocal microscopy.

Trehalose Uptake Kinetics. HepG2 cells with or without TRET1 were incubated in cell culture medium containing 50 mM trehalose for specified time periods (0, 8, 18, and 24 h) to load the sugar. The trehalose-containing medium was then completely removed, and the attached cells were rinsed twice with PBS. Cells were harvested by incubation for 5 min in 0.25% trypsin-EDTA, dilution with cell culture medium, and centrifugation. Cells were then resuspended in deionized water, subjected to three freeze-thaw cycles in liquid nitrogen, and then centrifuged at 16,000 × g for 15 min. Trehalose in the supernatant was analyzed by HPLC as described previously (4). The trehalose content per million cells was converted to an approximate intracellular concentration using a measured diameter of 15.1 μm for HepG2 cells and assuming a spherical form and 70% osmotically active water.

Spin-Drying of Cells. Monolayers of HepG2 cells expressing various combinations of AfrLEA2, AfrLEA3m, and TRET1 were allowed to attach to round glass coverslips (22-mm diameter, collagen-coated; BD BioCoat catalog no. 354089; BD Biosciences) during an 18-h incubation in cell culture medium. For all cells expressing TRET1, the medium also contained 50 mM trehalose. Cells were prevented from attaching to the edge of the coverslips using 20-mm-diameter Press-to-Seal silicone isolators (Life Technologies). Immediately before spin-drying, the cell culture media was completely removed using a Pasteur pipette and replaced with the spin-drying solution (1.8 M trehalose, 10 mM KCl, 5 mM glucose, 20 mM Hepes, and 120 mM choline chloride; pH 7.4). This trehalose-containing solution was essential for a successful outcome regardless of the intracellular variables tested. Spin-drying was performed using a commercially available spinning machine (Brewer Science model Cee 200) as described previously (4).

FTIR spectromicroscopy was performed with a Nicolet Continuum FTIR spectromicroscope (Thermo Scientific) equipped with a mercury cadmium telluride detector, to examine the uniformity of water distribution in spin-dried trehalose films prepared with the aforementioned drying solution and determine the average moisture content of these noncellular samples (4). Residual water was estimated by interpolation using the trehalose-water binary phase diagram (5) derived from the glass-transition temperature (Tg) measured by FTIR spectromicroscopy. The FTIR results were then compared with values obtained by bulk gravimetric analysis of cell monolayers overlain with drying solution and spin-dried (4). The two datasets were statistically identical. After drying, the samples were quickly removed from the spin-dryer and placed into six-well (35 mm) culture plates (Corning). Cells were immediately rehydrated in 0.5 mL of cell culture medium, and membrane integrity and long-term viability were evaluated.

Assessment of Membrane Integrity and Long-Term Viability. Membrane integrity after spin-drying was assessed using Syto-13/ethidium bromide membrane integrity assays (Molecular Probes). The stock solution for Syto-13/ethidium bromide membrane staining was prepared by adding 10 mL of 1 mg/mL Syto-13 solution (aq.) and 5 mL of 1.0 mg/mL ethidium bromide solution (aq.) to 8 mL of MEM without phenol red or serum (Life Technologies). After rehydration, 500 μL of Syto-13/ethidium bromide solution was added to the cells attached on coverslips, and the samples were incubated at 37 °C for 5 min. These samples were then imaged using an inverted microscope (Carl Zeiss) with FITC and PI filters. Cell viability was determined immediately after rehydration using this technique by counting the live (green) and dead (red) cells in seven representative images obtained at different locations on the coverslip.

Long-term viability and growth profiles were determined by incubating the rehydrated cell samples in fully complemented medium for 7 d in six-well (35 mm) culture plates. Parallel samples were used for the various experimental treatments, and the viability of cells after dehydration was measured by counting cells on days 1, 3, and 7. To ensure that only viable cells were counted, the membrane integrity of these cells was determined by trypan blue exclusion.


Fig. S1. AfrLEA2-GFP exhibits a predominately cytoplasmic distribution, but at 72 h after the initial transfection event, some of the chimeric protein has entered the nucleus, as demonstrated by the increased green staining of the organelle. The nuclear staining is far more prevalent than that seen after 24 h (Fig. 3). (Left) Distribution of green fluorescence from AfrLEA2-GFP. (Center) Distribution of mitochondria based on MitoTracker Red labeling. (Right) Merged image.